Primary ciliary dyskinesia. From symptoms to diagnosis.
A case report of a 4.5-year-old girl

Abstract
Primary ciliary dyskinesia is a rare genetic disease. Early diagnosis is aimed at delaying pulmonary complications, such as bronchiectasis, reducing lung dysfunction and hearing impairment, as well as providing optimal treatment of rhinitis and sinusitis. The awareness of the disease is still poor, and primary ciliary dyskinesia is often underdiagnosed or confused with other diseases, such as asthma. Chronic productive cough and rhinitis, which are the main symptoms, develop already in infancy. Therefore, differential diagnosis is a key element of the diagnostic and therapeutic process. The area of search depends on patient’s age, the nature of symptoms (type of cough, time of symptom onset, repeatability of symptoms), concomitant symptoms and family history. We present a case report of a 4.5-year-old girl with recurrent lower respiratory tract infections and chronic cough, monitored for asthma and unsuccessfully treated with antiasthmatics.

Keywords: primary ciliary dyskinesia (PCD), diagnosis, chronic cough
INTRODUCTION

Chronic cough and recurrent respiratory infections are currently among the most common reasons of visits to pulmonologists and allergologists. These symptoms may be caused by infectious agents, congenital defects of the airways or psychogenic factors (Tab. 1)(1). In addition to mild conditions, which often resolve spontaneously, chronic disorders frequently leading to irreversible respiratory lesions, respiratory failure and severe complications may also occur. Therefore, differential diagnosis is a key element of the diagnostic and therapeutic process. The area of search depends on patient’s age, the nature of symptoms (type of cough, time of symptom onset, repeatability of symptoms), concomitant symptoms and family history. We present a case report of a child with recurrent lower respiratory tract infections and chronic cough, monitored for asthma and unsuccessfully treated with antiasthmatics.

CASE REPORT

A nearly 4.5-year-old girl was referred to the Department of Pulmonology, Institute of Tuberculosis and Lung Diseases in Rabka-Zdrój for extended pulmonological diagnosis. The child was born at 41 weeks from first pregnancy via vaginal delivery, with a birth weight of 3,340 g and an Apgar score of 7/10/10/10 at 1/3/5/10 minutes, respectively. On the second day of life, the girl was reluctant to eat and developed cough. Physical examination revealed snuffles, nasal discharge as well as dry rales over the lung fields on auscultation. Lung radiography and laboratory testing were unremarkable. A clinical diagnosis of pneumonia was established. Broad-spectrum antibiotic therapy, parenteral nutrition, and physical therapy were included, leading to improved general condition. Screening tests, including immunoreactive trypsinogen test, were performed. An ENT (ear, nose, throat) check was ordered due to abnormal hearing screening.

The girl was re-admitted at 6 weeks of age due to pneumonia. The child was put under the care of a pulmonologist. Sweat test was normal. Further re-admissions due to pneumonia took place at 3 and 5 months of age, the latter one accompanied by airway obstruction. The girl was referred to a pulmonology centre, where (inhaled and food) allergy was excluded, for further treatment and extended diagnosis. An ENT examination showed accumulated discharge at the entrance to the larynx and in the glottis. Systemic glucocorticoids (GCS), bronchodilators, clarithromycin and kinesitherapy were used, leading to resolution of obstruction, auscultatory symptoms and overall clinical improvement. The patient was diagnosed with early-onset asthma and put on chronic inhaled GCS therapy.

At 20 months of age, the girl was re-admitted for pneumonia. IV antibiotic therapy, systemic and inhaled GCS, and bronchodilators were used in the treatment. Patient’s condition improved, but recurrent obstruction and problems with systemic GCS discontinuation were still observed despite the treatment. After 10-day therapy, the girl was discharged with recommendations to continue treatment and to report for allergological and pulmonological consultations.

At the age of four years, the child was re-admitted due to pneumonia. Lung radiography (Fig. 1) showed consolidation of parenchymal atelectatic lesions in the mid-lower parts of the right lung (probably the lower lobe). Furthermore, the description included diffuse parenchymal densities, with the largest lesions located in the lower pulmonary fields and in the perihilar region. Slight amounts of fluid were present in both pleural cavities. Laboratory tests showed increased acute inflammatory markers: C-reactive protein (CRP) 66 mg/L, white blood cells (WBC) 18.09 thousand/µL. Antibiotic treatment (amoxicillin/clavulanic acid, clarithromycin), bronchodilators, inhaled GCSs and mucolytics were administrated. Transient increase in inflammatory markers (CRP 127 mg/L, WBC 24.33 thousand/µL) was observed. The treatment led to improvement in overall condition and partial reduction of auscultatory symptoms.

**The most common causes of chronic cough**
- Nonspecific postinfectious cough
- Upper respiratory tract cough syndrome
- Asthma
- Protracted bacterial bronchitis
- Bronchiectasis
- Cystic fibrosis
- Congenital respiratory/cardiovascular defects
- Primary ciliary dyskinesia
- Bronchopulmonary dysplasia
- Foreign body in the airways
- Gastroesophageal reflux disease
- Interstitial lung disease
- Exposure to tobacco smoke and other pollutants
- Habitual cough

**Tab. 1. The most common causes of chronic cough**(1)
Also, the girl’s parents reported the presence of chronic wet cough with expectoration of secretions, nasal obstruction and chronic rhinitis from birth.

The girl was admitted to the Department of Pulmonology at the Institute of Tuberculosis and Lung Diseases a month after the last hospitalization. She was in fairly good overall condition at admission. Physical examination revealed a hypostenic body type, slightly marked signs of dyspnoea, such as tachypnoea, involvement of additional respiratory muscles, retraction at the suprasternal notch, nasal obstruction malocclusion, multiple diffuse crackles, multiple bilateral dry rales, and subscapular bronchial respiratory sound on auscultation; clubbed fingers. Furthermore, there was a delayed speech development – the girl used single indistinct words. Remarkable laboratory findings included increased inflammatory markers: CRP 23 mg/L, WBC 16.6 thousand/L, erythrocyte sedimentation rate (ESR) 44 mm, alpha-1 antitrypsin 2.438 g/L, complement component C3 1.411 g/L. The diagnosis was extended to include sweat test (normal) and upper nasal nitric oxide (nNO) testing (unsuccessful – lack of cooperation). Lung ultrasonography showed extensive atelectasis in the lower right lobe with abolition of bronchial aeration as well as irregular pleura with Z-lines in the basal part of the lungs, and fine consolidation/atelectasis foci in the region of the middle lobe and the lingula. The diagnosis was extended to include high-resolution computed tomography (HRCT) (Fig. 2). Bronchoscopy revealed small amounts of purulent fluid on the tracheal walls; with large amounts of purulent discharge filling the bronchi of both lungs, with right-sided predominance. Samples were collected to investigate the cilia under a light microscope. Moderately numerous conglomerates of ciliary epithelium covered with normal-length cilia were obtained, which showed no movement in most of the fragments (only stiff beat and oscillations around the long axis were occasionally observed). Haemophilus influenzae was present in bronchial culture, with 85% of neutrophils in cytology. An ENT consultation was carried out; otitis media with effusion was diagnosed, but hearing assessment was not possible (uncooperative patient). The girl developed fever and increased dyspnoea requiring temporary use of passive oxygen therapy after bronchoscopy. Inflammatory markers increased. The patient was put on antibiotic therapy, mucolytics, and bronchial drainage, which led to significant improvement of her overall condition, normalisation of inflammatory markers and partial resolution of auscultatory symptoms. Follow-up lung ultrasound showed systematic regression of lesions, reduced atelectasis in the entire lower lobe up to the segment 10 in the right lung. Primary ciliary dyskinesia (PCD) was diagnosed based on the overall clinical picture, basic examinations and laboratory testing. Genetic testing and ciliary evaluation using electron microscopy and immunofluorescence were ordered. No epithelial cilia or microvilli were found in ultrastructural assessment using an electron microscope. No basal bodies were found in the apical epithelial cell layer. The obtained findings and the clinical picture confirmed PCD. Other results were still analysed at the time of publication of the article.

**DISCUSSION**

PCD is a rare genetic disease. The symptoms occur as a result of abnormal structure/function of cilia covering the epithelium of the mucous membranes of sinuses, bronchi, reproductive tract, and the central nervous system. PCD is inherited in an autosomal recessive manner, with a prevalence of about 1:10,000–40,000 live-born children(2). Almost 40 genes are responsible for ciliary defects and, abnormal ciliary movement. A normal cilium is made up of 9 (A+B) symmetrically arranged peripheral microtubule doublets and 1 centrally located doublet. Inner and outer dynein arms are attached to the peripheral microtubule doublets. The peripheral doublets are connected by nexin links, whereas the so called radial spokes attach them to the central doublet. Normal structure determines proper movement (Fig. 3).
Kartagener was first to observe the triad of symptoms: chronic sinusitis, bronchiectasis, and situs inversus. About 45–55% of patients with PCD present with situs inversus (SI)(5). Many patients require repeated medical appointments and hospital admissions before the diagnosis is confirmed(6).

The variable clinical outcome is characterised by recurrent and chronic lower and upper respiratory tract infections, which consequently lead to bronchiectasis and pulmonary dysfunction. Abnormal ciliary function and/or structure leads to ineffective bronchial clearance, causing discharge accumulation.

Symptoms depend on child’s age. In newborns, typically those born full-term, respiratory distress syndrome is observed a few hours after birth. Chronic cough and rhinitis occur already from the first days of life. Congenital heart defects are more common in patients with PCD compared to the general population(7). Impaired ciliary clearance leads to recurrent respiratory infections. Particular attention should be paid to recurrent or chronic otitis media and recurrent bronchitis, which, over time, progress to chronic bronchitis followed by bronchiectasis at a later age. The middle lobe, lingula or lower lobes are a common location. Chronic, productive cough with expectoration of mucopurulent discharge is a typical symptom. These symptoms are accompanied in older children by chronic paranasal sinusitis and, often, hearing impairment. About 50% of adult men are affected by infertility due to the lack of sperm motility; aspermia is diagnosed in some patients. Abnormal ciliary movement in the fallopian tubes may lead to ectopic pregnancy in women(8).

The diagnosis of PCD requires specialist equipment and is performed in specialist centres. Therefore, PICADAR.
with the so-called entry criterion, i.e. a question which re-
concha or bronchus are investigated using a light micro-
scope, which allows for the assessment of ciliary type and
components using florescent markers. A number of anti-ciliary
antibodies are available, including those directed against
outer and inner dynein arms, radial spokes and nexin links.

Immunofluorescence allows the detection of all ultrastruc-
tural abnormalities visible under an electron microscope
and outer dynein arms accompanied by abnormal micro-
tubule structure, without the need for further diagnosis(8).

Normal ciliary ultrastructure may be assessed with trans-
mision electron microscopy (TEM). PCD may be di-
agnosed based on the identification of typical PCD fea-
tures, such as absence of outer dynein arms, absence of
both inner and outer dynein arms, or absence of inner
and outer dynein arms accompanied by abnormal micro-
tubule structure, without the need for further diagnosis(8).

Abnormal ciliary ultrastructure is not visualised in 20% of
patients with typical PCD symptoms(8). Biopsy should
be performed during recovery, i.e. at least 2 weeks after ex-
cacerbations, to exclude secondary causes(11). In the case of
doubts arising from patient's exposure to environmental
or infectious factors, or technical reasons, in vitro epithe-
lial culture may be performed. Epithelial culture reduces
the number of false positive results. It helps evaluate rare
cases, such as ciliary agenesis, and limits the need for fur-
ther biopsies(11). Immunofluorescence is a less available and less commonly
used method. It involves visualizing individual ciliary com-
ponents using florescent markers. A number of anti-ciliary
antibodies are available, including those directed against
outer and inner dynein arms, radial spokes and nexin links.

High-speed video microscopy analysis (HSV A), which re-
ords at 120–500 frames per second with possible assess-
ment of motion at a slow rate (30–60 frames per second –
fps), is an important element of diagnosis in patients with
PCD(3). Fresh, biopsy samples from under the inferior
concha or bronchus or from in vitro epithelial culture may be
investigated using a light micro-
scope, which allows for the assessment of ciliary type and
movement rate. HSV A shows high sensitivity (100%) and
specificity (96%) in the diagnosis of PCD(12). Under nor-
mal conditions, ciliary movement frequency is 8–15 Hz,
with lower values for cilia sampled from more peripheral
airways. Normal beat pattern is synchronised (Fig. 3). It
is described as a sequence of consecutive beats: a strong
beating stroke of an upright cilium followed by a recovery
stroke, which is initiated by bending of the proximal axo-

Tab. 2. PICADAR questionnaire(9)

<table>
<thead>
<tr>
<th>Does the patient have a daily productive cough that started in early childhood?</th>
<th>NO (the probability of PCD cannot be estimated based on PICADAR)</th>
<th>YES</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Was the patient born preterm or full-term?</td>
<td>Preterm = 0</td>
<td>Full-term = 2</td>
</tr>
<tr>
<td>2. Did the patient experience chest symptoms in the neonatal period (e.g. tachypnoea, cough, pneumonia)?</td>
<td>NO = 0</td>
<td>YES = 2</td>
</tr>
<tr>
<td>3. Did the patient require prolonged hospitalisation or re-admission during the neonatal period?</td>
<td>NO = 0</td>
<td>YES = 2</td>
</tr>
<tr>
<td>4. Does the patient have a situs abnormality (situs inversus or heterotaxy)?</td>
<td>NO = 0</td>
<td>YES = 4</td>
</tr>
<tr>
<td>5. Does the patient have a congenital heart defect?</td>
<td>NO = 0</td>
<td>YES = 2</td>
</tr>
<tr>
<td>6. Does the patient have persistent renal rinitis?</td>
<td>NO = 0</td>
<td>YES = 1</td>
</tr>
<tr>
<td>7. Does the patient experience chronic ear or hearing symptoms (e.g. chronic otitis media with effusion, hearing loss, tympanic membrane perforation)?</td>
<td>NO = 0</td>
<td>YES = 1</td>
</tr>
</tbody>
</table>

TOTAL:
and cases where the picture is apparently normal or slightly abnormal. The sensitivity and specificity of immunofluorescence are unknown. The results depend on the quality and combination of antibodies used. False negative results are possible if an abnormal protein is present in the cilium. Genetic analysis is another diagnostic method. So far, 39 PCD-associated genes have been identified. A combination of 2 mutations is needed for the diagnosis. Unfortunately, known mutations are detected in only about 65% (50–75%) of patients. There are ongoing studies to identify further mutations.

Differential diagnosis of suspected PCD should include all disease entities characterised by chronic productive cough (Tab. 1) and chronic rhinitis leading to bronchiectasis. It should be emphasised that patients with PCD are at the beginning treated the same way as those with asthma. Obstructive defects in functional tests can be seen with both PCD and asthma. Bronchodilator responsiveness is not exclusive to asthma and does not exclude PCD. Similarities and differences between the symptoms of PCD and asthma are summarised in Tab. 3.

### Treatment

Due to the lack of randomised clinical trials, the recommendations for the treatment of PCD are based on the treatment guidelines for chronic pulmonary diseases, cystic fibrosis (CF) in particular. Although the symptoms seen in patients with PCD and CF are similar, their mechanisms differ: PCD – retention of normal secretions, CF – retention of excessively thick secretions. Treatment aims include prevention of bronchiectasis and maintaining good pulmonary function. Evacuation of secretions using appropriate, age-matched physical therapy, respiratory gymnastics and physical exercise, is also one of the key measurements.

Efficacy of mucoactive agents has been described: recombinant human DNase and hypertonic saline. The benefits of N-acetylcysteine treatment and chronic use of bronchodilators have not been confirmed. Aggressive antibiotic therapy of upper and lower respiratory tract infections should include pathogens that are most common in PCD: *H. influenzae*, *S. aureus* and *S. pneumoniae*.

Due to the large amount of secretion in the respiratory tract, *P. aeruginosa* should be also considered.

ENT care and treatment is also necessary in patients with impaired sinus drainage, chronic paranasal sinusitis and hearing impairment. Avoiding active and passive smoking, elimination of exposure to environmental pollution and minimising exposure to pathogens play a key role in all patients. Vaccinations are also important.

### CONCLUSIONS

The diagnosis of PCD is delayed or missed in the diagnostic process of chronic and recurrent respiratory infections. The median age at diagnosis is 5.3 years in Europe. This age is lower in patients with situs inversus, i.e. 3.5 years vs. 5.8 years. Early diagnosis is aimed at delaying pulmonary complications, such as bronchiectasis, reducing lung dysfunction and hearing impairment, as well as optimal treatment of rhinitis and sinusitis. Patients with PCD require multidisciplinary care with cooperation of a pulmonologist, ENT specialist and physiotherapist. Systematic assessment of respiratory function, bacterial sputum culture, lung imaging and audiogramme are recommended for these patients.

### Conflict of interest

The authors do not report any financial or personal connections with other persons or organizations, which might negatively affect the contents of this publication and/or claim authorship rights to this publication.

### References