Relapsing polychondritis: state of the art on clinical practice guidelines

Simona Rednic, Laura Damian, Rosaria Talarico, Carlo Alberto Scire, Alexander Tobias, Nathalie Costedoat-Chalumeau, David Launay, Alexis Mathian, Lisa Matthews, Cristina Ponte, Paola Toniati, Stefano Bombardieri, Chariss Frank, Matthias Schneider, Vanessa Smith, Maurizio Cutolo, Marta Mosca, Laurent Arnaud

ABSTRACT

Due to the rarity of relapsing polychondritis (RP), many unmet needs remain in the management of RP. Here, we present a systematic review of clinical practice guidelines (CPGs) published for RP, as well as a list of the most striking unmet needs for this rare disease. We carried out a systematic search in PubMed and Embase based on controlled terms (medical subject headings and Emtree) and keywords of the disease and publication type (CPGs). The systematic literature review identified 20 citations, among which no CPGs could be identified. We identified 11 main areas with unmet needs in the field of RP: the diagnosis strategy for RP; the therapeutic management of RP; the management of pregnancy in RP; the management of the disease in specific age groups (for instance in paediatric-onset RP); the evaluation of adherence to treatment; the follow-up of patients with RP, including the frequency of screening for the potential complications and the optimal imaging tools for each involved region; perioperative and anaesthetic management (due to tracheal involvement); risk of neoplasms in RP, including haematological malignancies; the prevention and management of infections; tools for assessment of disease activity and damage; and patient-reported outcomes and quality of life indicators. Patients and physicians should work together within the frame of the ReCONNET network to derive valuable evidence for obtaining literature-informed CPGs.

INTRODUCTION

Relapsing polychondritis (RP) is a systemic inflammatory disease primarily affecting the cartilaginous structures of the ears, nose and tracheobronchial tree, but also the joints, the inner ear, the eyes and the cardiovascular system. The first case of RP was described in 1923 by Jakusch-Wartenhorst, but little attention has been given to the entity until the 1960s, when Pearson et al introduced the name ‘relapsing polychondritis’. The classification criteria for RP by Michet et al require the presence of proven inflammation in at least two of three of the auricular, nasal or laryngotracheal cartilages, or proven inflammation in one cartilage plus two other signs, including ocular inflammation, vestibular dysfunction, seronegative inflammatory arthritis or hearing loss. RP is a chronic disease with a flaring–remitting course. The exact cause of RP is still unknown, but the disease is mostly regarded as an immune-mediated disease, as there is a well-documented overlap of RP with other rheumatic and autoimmune diseases. Moreover, RP is strongly associated with the Human Leucocyte Antigen (HLA) allele DR4, and various immune responses directed against cartilage components have been demonstrated in patients with RP. Since RP is a very rare disease, with a prevalence estimated to be as low as a few cases per million, it remains an under-researched area. Scores for assessing both disease activity and damage in RP have been developed by our group, with the help of a panel of international experts. Importantly, due to the rarity of the disease, the lack of

Key messages

What is already known about this subject?

► Due to the rarity of the disease and the paucity of available evidence, many unmet needs remain in the field of relapsing polychondritis (RP).

What does this study add?

► We performed a systematic review of clinical practice guidelines published for RP, and we identified the most striking unmet needs in this rare disease.

How might this impact on clinical practice?

► Due to the extreme rarity of the disease, patients and physicians should work together within the frame of the ReCONNET network to derive valuable evidence for informing literature-based clinical practice guidelines for RP.
adequate networks of care for RP in most countries and the paucity of drug efficacy data, many unmet needs remain in the field of RP. Here we present a systematic review of clinical practice guidelines (CPGs) published for RP, as well as a list of the most striking unmet needs in this rare disease. The review has been carried out under the framework of the European Reference Network on Rare and Complex Connective Tissue and Musculoskeletal Diseases (ERN ReCONNET), gathering the experience of several European centres with experience in the diagnosis and follow-up of RP and involving patient representatives.

METHODS

We carried out a systematic search in PubMed and Embase based on controlled terms (medical subject headings (MeSH) and Emtree) and keywords of the disease and publication type (CPGs). We reviewed all the published articles in order to identify existing CPGs on diagnosis, monitoring and treatment, according to the definition of the Institute of Medicine 2011 (CPGs are statements that include recommendations intended to optimise patient care that are informed by a systematic review of evidence and an assessment of the benefits and harms of alternative care options). The disease coordinator (DC) of the ERN ReCONNET for RP has assigned the work on CPGs to the healthcare providers involved. Moreover, in order to implement the list of guidelines provided by the Medline and Embase search, the group also performed a hand search. A first screening among papers included in the final list (systematic search + hand search) based on title and abstract selected evidence-based medicine guidelines. A general assessment of the CPGs has been performed following the Appraisal of Guidelines for Research & Evaluation II (AGREE II) tool checklist not for formal appraisal but only to inform discussion. A discussion group was set for the evaluation of the existing CPGs and to identify the unmet needs.

occasionally cartilage biopsies. RP mimickers such as Winkler’s disease) are often suspected first. However, even in the presence of episcleritis and scleritis, which may involve such as intermittent arthritis or eye involvement. The diagnostic delay is associated with recognition early, especially in the absence of typical cartilage involvement. The diagnostic delay is associated with the lack of ear, nose or joint involvement. Early signs of RP in such cases may be intermittent arthritis or eye involvement such as episcleritis and scleritis, which may point a search for RP. Moreover, another immune-mediated disease, such as systemic lupus erythematosus, Sjogren’s syndrome, vasculitis, antiphospholipid syndrome, rheumatoid arthritis, spondyloarthritis, inflammatory bowel disease, thyroiditis and others, is associated with RP in up to 30% of cases and should be searched for systematically. On diagnosis, a baseline assessment is necessary in order to evaluate disease activity, organ involvement and disease damage, and to identify potentially associated diseases. A typical baseline assessment may include otorhinolaryngology and ophthalmology examinations, cardiovascular screening for valvulopathies and aortic involvement, pulmonary assessment, renal function testing, testing for antineutrophil cytoplasmic autoantibodies (ANCA) and possibly haematological assessment to rule out myelodysplastic syndrome (MDS). However, the frequency of these assessments is not standardised and may require further validation. To date, there is a real need for validated diagnostic biomarkers in RP, as well as markers predictive of disease activity, specific organ involvement or prognosis. There is no characteristic laboratory analysis in RP. An inflammatory syndrome is present in more than 60% of patients, but is not constant during flares. Antinuclear antibodies with no particular specificity and ANCA, mostly atypical in immunofluorescence, with no proteinase 3 (PR3) or myeloperoxidase (MPO) specificity, have been reported in several series. The potential candidate biomarkers, the anticollagen type II antibodies and antimitrillin type I antibodies, are neither sensitive nor specific enough. Moreover, cartilage biopsy is positive only in two-thirds of cases and does not show any specific change; its added value is therefore limited. Several potential disease activity markers such as antihyaluronan type II antibodies, antimitrillin-I antibodies, other autoantibodies targeting collagen structures (serum cartilage oligomeric matrix protein, urinary type II collagen neoepitope) or interferon-gamma, interleukin (IL)-12 and IL-2 and serum soluble triggering receptor expressed on myeloid cells 1 (sTREM-1) have been reported to parallel the flares. None of these have entered clinical practice.

Mortality in RP is more than double compared with the general population; the most frequent causes of death are respiratory, cardiac and haematological involvement. Three phenotypes with different presentations have been recently described in a cluster analysis of a cohort: a haematological form (in 10% of cases), a respiratory form (in 25%) and a mild one with good prognosis (about 65% of cases). Some complications such as aortic, vasculitis or anaemia are pejorative prognosis factors. In addition, the male gender is associated with worse prognosis and higher prevalence of uveitis, hearing loss, vestibular disorder, as well as greater necessity for methylprednisolone and cyclophosphamide pulses.

Box 1 Unmet needs in the field of relapsing polychondritis (RP)

Major unmet needs in the field of RP.
► The diagnosis strategy for RP.
► The therapeutic management of RP.
► The management of pregnancy in RP.
► The management of the disease in specific age groups (for instance in paediatric-onset RP and its relationship with growth, or in the geriatric population).
► The evaluation and management of adherence to treatment.
► The follow-up of patients with RP, including the frequency of screening for the potential complications, mainly of respiratory, cardiac, aortic, ocular and joint involvement, and the optimal imaging tools for each involved region.
► Perioperative and anaesthetic management (due to tracheal involvement).
► Risk of neoplasms in RP, including haematological malignancies, and the need for screening for an occult neoplasm.
► The prevention and management of infections.
► Tools for assessment of disease activity and damage.
► Patient-reported outcomes and quality of life indicators.

doi:10.1136/rmdopen-2018-000788

RESULTS

State of the art on CPGs

The systematic literature review identified 20 citations, among which no CPGs could be identified (figure 1).

Unmet needs

Clinicians’ unmet needs

Following extensive discussions between the DC and other members of the ReCONNeT network, we identified 11 main areas with unmet needs in the field of RP (box 1). We also report detailed list of clinical manifestations that may be seen in RP (table 1).

The diagnostic strategy for RP

The classical classification criteria for RP (such as those by Michet et al, McAdams or Damiani-Levine) have been empirically postulated. The disease is rare and difficult to recognise early, especially in the absence of typical cartilage involvement. The diagnostic delay is associated with the lack of ear, nose or joint involvement. Early signs of RP in such cases may be intermittent arthritis or eye involvement such as episcleritis and scleritis, which may point a search for RP. However, even in the presence of external ear inflammation, an infection, local trauma, insect bite and chondrodermatitis nodularis helicis (Winkler’s disease) are often suspected first.

The diagnosis of RP is mostly clinical, and may be informed by laboratory data, imaging techniques and occasionally cartilage biopsies. RP mimickers such as granulomatosis with polyangiitis (GPA), T-cell lymphoma, sarcoidosis and so on have to be differentiated from RP. Moreover, another immune-mediated disease, such as systemic lupus erythematosus, Sjogren’s syndrome, vasculitis, antiphospholipid syndrome, rheumatoid arthritis, spondyloarthritis, inflammatory bowel disease, thyroiditis and others, is associated with RP in up to 30% of cases and should be searched for systematically. On diagnosis, a baseline assessment is necessary in order to evaluate disease activity, organ involvement and disease damage, and to identify potentially associated diseases. A typical baseline assessment may include otorhinolaryngology and ophthalmology examinations, cardiovascular screening for valvulopathies and aortic involvement, pulmonary assessment, renal function testing, testing for antineutrophil cytoplasmic autoantibodies (ANCA) and possibly haematological assessment to rule out myelodysplastic syndrome (MDS). However, the frequency of these assessments is not standardised and may require further validation.

To date, there is a real need for validated diagnostic biomarkers in RP, as well as markers predictive of disease activity, specific organ involvement or prognosis. There is no characteristic laboratory analysis in RP. An inflammatory syndrome is present in more than 60% of patients, but is not constant during flares. Antinuclear antibodies with no particular specificity and ANCA, mostly atypical in immunofluorescence, with no proteinase 3 (PR3) or myeloperoxidase (MPO) specificity, have been reported in several series. The potential candidate biomarkers, the anticollagen type II antibodies and antimitrillin type I antibodies, are neither sensitive nor specific enough. Moreover, cartilage biopsy is positive only in two-thirds of cases and does not show any specific change; its added value is therefore limited.

Several potential disease activity markers such as antihyaluronan type II antibodies, antimitrillin-I antibodies, other autoantibodies targeting collagen structures (serum cartilage oligomeric matrix protein, urinary type II collagen neoepitope) or interferon-gamma, interleukin (IL)-12 and IL-2 and serum soluble triggering receptor expressed on myeloid cells 1 (sTREM-1) have been reported to parallel the flares. None of these have entered clinical practice.

Mortality in RP is more than double compared with the general population; the most frequent causes of death are respiratory, cardiac and haematological involvement. Three phenotypes with different presentations have been recently described in a cluster analysis of a cohort: a haematological form (in 10% of cases), a respiratory form (in 25%) and a mild one with good prognosis (about 65% of cases). Some complications such as aortic, vasculitis or anaemia are pejorative prognosis factors. In addition, the male gender is associated with worse prognosis and higher prevalence of uveitis, hearing loss, vestibular disorder, as well as greater necessity for methylprednisolone and cyclophosphamide pulses.


[42x66]informed by laboratory data, imaging techniques and (Winkler’s disease) are often suspected first. However, even in the presence of episcleritis and scleritis, which may involve such as intermittent arthritis or eye involvement. The diagnostic delay is associated with recognition early, especially in the absence of typical cartilage involvement. The diagnostic delay is associated with the lack of ear, nose or joint involvement. Early signs of RP in such cases may be intermittent arthritis or eye involvement such as episcleritis and scleritis, which may point a search for RP. However, even in the presence of external ear inflammation, an infection, local trauma, insect bite and chondrodermatitis nodularis helicis (Winkler’s disease) are often suspected first.

The diagnosis of RP is mostly clinical, and may be informed by laboratory data, imaging techniques and occasionally cartilage biopsies. RP mimickers such as granulomatosis with polyangiitis (GPA), T-cell lymphoma, sarcoidosis and so on have to be differentiated from RP. Moreover, another immune-mediated disease, such as systemic lupus erythematosus, Sjogren’s syndrome, vasculitis, antiphospholipid syndrome, rheumatoid arthritis, spondyloarthritis, inflammatory bowel disease, thyroiditis and others, is associated with RP in up to 30% of cases and should be searched for systematically. On diagnosis, a baseline assessment is necessary in order to evaluate disease activity, organ involvement and disease damage, and to identify potentially associated diseases. A typical baseline assessment may include otorhinolaryngology and ophthalmology examinations, cardiovascular screening for valvulopathies and aortic involvement, pulmonary assessment, renal function testing, testing for antineutrophil cytoplasmic autoantibodies (ANCA) and possibly haematological assessment to rule out myelodysplastic syndrome (MDS). However, the frequency of these assessments is not standardised and may require further validation.

To date, there is a real need for validated diagnostic biomarkers in RP, as well as markers predictive of disease activity, specific organ involvement or prognosis. There is no characteristic laboratory analysis in RP. An inflammatory syndrome is present in more than 60% of patients, but is not constant during flares. Antinuclear antibodies with no particular specificity and ANCA, mostly atypical in immunofluorescence, with no proteinase 3 (PR3) or myeloperoxidase (MPO) specificity, have been reported in several series. The potential candidate biomarkers, the anticollagen type II antibodies and antimitrillin type I antibodies, are neither sensitive nor specific enough. Moreover, cartilage biopsy is positive only in two-thirds of cases and does not show any specific change; its added value is therefore limited.

Several potential disease activity markers such as antihyaluronan type II antibodies, antimitrillin-I antibodies, other autoantibodies targeting collagen structures (serum cartilage oligomeric matrix protein, urinary type II collagen neoepitope) or interferon-gamma, interleukin (IL)-12 and IL-2 and serum soluble triggering receptor expressed on myeloid cells 1 (sTREM-1) have been reported to parallel the flares. None of these have entered clinical practice.

Mortality in RP is more than double compared with the general population; the most frequent causes of death are respiratory, cardiac and haematological involvement. Three phenotypes with different presentations have been recently described in a cluster analysis of a cohort: a haematological form (in 10% of cases), a respiratory form (in 25%) and a mild one with good prognosis (about 65% of cases). Some complications such as aortic, vasculitis or anaemia are pejorative prognosis factors. In addition, the male gender is associated with worse prognosis and higher prevalence of uveitis, hearing loss, vestibular disorder, as well as greater necessity for methylprednisolone and cyclophosphamide pulses.
The therapeutic management of RP

The treatment in RP is mostly based on case reports and case series. The therapy is selected according to the clinical picture and its severity, and depends on the type of organ involvement. Minor nasal or auricular chondritis may respond to non-steroidal anti-inflammatory drugs, glucocorticoids,11 colchicine or dapsone. Conventional disease-modifying antirheumatic drugs (cDMARDs) are employed as glucocorticoid-sparing agents or in cases of more severe disease: methotrexate, azathioprine, ciclosporin, leflunomide, mycophenolate mofetil, cyclophosphamide and so on, sometimes in a step-ladder increment.11 No clinical trials have been published on biologics in RP.11 20 In a recent French series,21 the biologic agents used to treat RP included tumour necrosis factor inhibitors (among which adalimumab was shown to have the best remanence), tocilizumab, anakinra, rituximab and abatacept. Strikingly, only 2/3 of patients responded to treatment at 6 months, with a complete response observed in 19% of cases.

The management of eye involvement is critical, because necrotising scleritis is commonly associated with significant morbidity and may lead to ocular perforation. Patients should be referred to an ophthalmologist as early start of local and systemic therapy may avoid further complications. While minor cases such as episcleritis may be managed using topical treatments, the most severe cases usually prompt aggressive systemic treatment such as glucocorticoid and cyclophosphamide infusions. Cases of necrotising scleritis treated with biologics (infliximab or adalimumab) have also been reported.10 11 The use of cyanoacrylate glue repairment has been reported for peripheral corneal perforations.22

The management of respiratory airway involvement depends on its presentation and severity. About 25% of cases will develop a laryngotracheal stricture.23 Early signs of tracheobronchomalacia should be searched for. Tracheal narrowing can be addressed by tracheostomy, tracheal dilation, stenting and reconstructive surgery.11 However, it is not clear whether these procedures do not aggravate RP, as they may result in distal airway inflammation.14 Due to the risk of tracheal perforation, pulmonary fibroscopy should generally be strongly discouraged in RP. When absolutely needed, the procedure should be performed only by very skilled pneumologists, with the backup of an intensive care unit. Nasal reconstructive surgery has controversial results; bone grafts are mostly

---

Table 1 Main clinical features of RP and proposed management (expert opinion)

<table>
<thead>
<tr>
<th>Main clinical manifestations</th>
<th>Typical therapeutic management* (based on expert opinion)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal or auricular chondritis. Peristernal chondritis.</td>
<td>NSAIDs, GCs. In case of relapsing disease colchicine, dapsone, methotrexate or other conventional immunosuppressive agents or biologics.</td>
</tr>
<tr>
<td>Tracheal chondritis.</td>
<td>GCs, methylprednisolone infusion, csDMARDs, conventional immunosuppressive agents (eg, cyclophosphamide) or biologics.</td>
</tr>
<tr>
<td>Articular manifestations. Peripheral and/or axial involvement.</td>
<td>NSAIDs, GCs, csDMARDs, conventional immunosuppressive agents (eg, methotrexate) or biologics.</td>
</tr>
<tr>
<td>Cutaneous involvement.</td>
<td>GCs, colchicine, dapsone (especially in case of neutrophilic dermatitis), methotrexate.</td>
</tr>
<tr>
<td>Cardiac involvement. Valvular involvement.</td>
<td>GCs, csDMARDs, conventional immunosuppressive agents (eg, methotrexate) or biologics.</td>
</tr>
<tr>
<td>Ocular involvement.</td>
<td>Topical GCs, cycloplegic. All patients with ocular involvement should be referred to an ophthalmologist. csDMARDs, conventional immunosuppressive agents or biologics may be necessary.</td>
</tr>
<tr>
<td>Audiovestibular dysfunction.</td>
<td>GCs, methylprednisolone infusion, csDMARDs, conventional immunosuppressive agents or biologics.</td>
</tr>
<tr>
<td>Neurological manifestations.</td>
<td>GCs, methylprednisolone infusion, csDMARDs, conventional immunosuppressive agents (eg, cyclophosphamide) or biologics.</td>
</tr>
<tr>
<td>Renal involvement.</td>
<td>In most cases, renal involvement suggests differential diagnoses such as ANCA-associated vasculitis.</td>
</tr>
</tbody>
</table>
preferred (skull or iliac bone) since common cartilage may become inflamed after trauma; however L-shaped costal cartilage has been employed successfully in disease remission under immunosuppressants.

The management of arthritis is also non-codified. In most cases, the episodes of arthritis are self-limited and non-erosive and its occurrence does not parallel other disease features. The costochondral, sternoclavicular and manubriosternal joints, as well as the peripheral large and small joints, may be involved, sometimes in an asymmetric manner. cDMARDs, sometimes in combination with biologic therapy, may be required.

In a recent large study, aortic involvement, consisting of aneurysms or ectasia, mainly of the thoracic and abdominal aorta, occurred in 6.4% of cases, after a median follow-up of 2 years. An older study showed that a close vascular follow-up is needed in RP and that reintervention may be necessary in some cases. In a recent series, up to 22% of patients had a cardiac valvulopathy, usually not severe. Periodic echocardiography may detect the progression of a valvulopathy despite the apparent quiescence of RP.

The existence of a renal involvement is a controversial issue in RP, because in most cases the correct diagnosis is not RP but GPA or microscopic polyangiitis.

The skin involvement is non-specific, including neutrophilic dermatosis, cutaneous vasculitis, nodules, aphthae, superficial venous thrombosis and so on. Chronic skin lesions are more frequent in patients with late-onset RP and are associated with MDS.

**The management of pregnancy in RP**

The management of pregnancy in RP relies on scarce data. In a case series of 25 pregnancies in 11 women, 1 elective medical termination was performed due to cyclophosphamide therapy; flares occurred in 7 of the remaining 24 cases, while the disease was considered stable in 16 cases and asymptomatic in 1. RP onset during pregnancy, at about 20 weeks of gestation, has also been published. No evidence of neonatal RP was observed in neonates, but the risk of fetal loss was increased.

**RP in specific age groups**

A systematic review of paediatric-onset RP identified the most common presenting features to be joint pain, ocular inflammation and chondritis. Paediatric RP shares many features with adult RP; however, children frequently have a familial history of autoimmunity and usually do not have associated autoimmune diseases. The outcome may be fatal, mostly due to cardiac complications; hence, the screening for complications is mandatory. Growth does not seem to be impaired by cartilage involvement, although there are exceptions with regard to the epiphysis plate involvement and destructive arthritis. Increased MRI bone marrow signal of unclear significance has been reported in two children. In very young children, the monogenic autoinflammatory disease chronic atypical neutrophilic dermatosis with lipodystrophy and elevated temperature (CANDLE) proteasome-associated auto-inflammatory syndrome (PRAAS) (CANDLE (PRAAS)) due to proteasome mutations (PSMB8) may evolve with polychondritis features (auricular chondritis, nodular episcleritis and keratitis, along with widespread inflammation and lipodystrophy).

In the elderly, the presence of chondritis may point to an associated vasculitis or to an MDS, more frequent in this age group. Previous age-related cardiac and respiratory involvement may affect the prognosis in the presence of silent valvular or respiratory disease due to RP.

**The evaluation and management of adherence to treatment**

There are no data referring to the adherence to therapy and its evaluation in patients with RP. This issue needs to be addressed, mostly in patients with a refractory disease that fails to respond to conventional therapy.

**The follow-up and the optimal imaging tool**

Follow-up of patients with RP includes screening for potential complications, mainly of respiratory, cardiac, aortic, ocular and joint involvement, as well as a proactive assessment of complications (at least respiratory, cardiac, ocular and otorhinolaryngological). The optimal frequency of the assessment is not known. However, in a recent study, aortic complications were detected at ≈2 years from diagnosis. Arthritis and the ocular features may be inaugural or may appear anytime during the disease. For respiratory involvement, a thorax CT scan may reveal oedematous tracheobronchial wall thickening with or without mural calcifications, deformity of cartilaginous structures, and narrowing of the trachea and bronchi. A typical feature is that the posterior part of the trachea (membranous trachea) is usually spared in case of early tracheal involvement, which is suggestive of RP. Dynamic examinations may reveal air structures collapse. CT scan of the neck may reveal thickening, calcification or destructive lesions of the larynx cartilages (such as cricoid or thyroid cartilage). Thoracoabdominal CT scan identifies the aortic involvement. Also not fully validated, several reports suggest that 18F-fluorodeoxyglucose Positron Emission Tomography (18F-FDG PET/CT) could be interesting for therapeutic response monitoring as the uptake may diminish or disappear after therapy. For articular involvement, Tc-99m bone scintigraphy may reveal active sites of inflammation. Cerebral MRI may be useful if central nervous system involvement is suspected. Otorhinolaryngoscopic examination reveals conductive and sensorineural hearing loss in about 40% of cases, sometimes along with nasal ulcerations, septal perforations, saddle nose deformity, arthrynod swelling and vocal cords deformities. Pulmonary spirometry, including inspiratory volumes, may help assess the tracheal involvement. Ophthalmoscopic examination may reveal any type of structure involvement (conjunctivitis, episcleritis, keratitis, uveitis, retinopathy, glaucoma, dacryocystitis and so on), the most dreaded being necrotising scleritis or...
peripheral ulcerative keratitis, which may rapidly lead to eye loss. 

**Perioperative and anaesthetic management**

In case of surgery, the anaesthesiologist should be informed of the risk of airway damage during intubation, tracheal stenosis or tracheobronchomalacia, structures and scars in the cricoarytenoid area, or dynamic airway obstruction. Anaesthesia may be converted to epidural analgesia whenever possible. A careful preoperative assessment should be planned and the anaesthesiologist may need to prepare various sizes of tracheal tubes and other airway manipulation devices.

**The risk of neoplasms**

Haematological malignancies, mainly MDS, but also lymphoma including mucosa-associated lymphoid tissue, myeloproliferative neoplasms, multiple myeloma and others, and more rarely solid cancers (lung, breast, colon, urothelial carcinoma, sarcoma), have been described in 11%–13% of cases, mainly in male patients with late-onset RP (>60 years) and in those with cutaneous involvement, mostly neutrophilic dermatitis. The reasons for this association are still occult. There are no guidelines regarding the type and frequency of screening for occult neoplasia in RP. Exophthalmos in patients with RP should prompt the search for lymphoma or IgG4-associated disease.

**The prevention and management of infections**

Infections are responsible for the significant morbidity (up to 35%) in the respiratory phenotype patients. Secretion clearance is impaired in patients with airway involvement. In addition, the risk of infections is higher in patients with an underlying MDS or diabetes. Steroids and biologic therapy administration may be complicated by systemic infections in such patients. There are no guidelines regarding infection prophylaxis and therapy in patients with RP, but it is reasonable to believe that the general recommendations used in immunocompromised patients should apply in patients with RP with immunosuppressive treatments.

**Tools for assessment of disease activity and damage**

As RP is a remitting-relapsing disease, disease activity has to be recorded during follow-up. The Relapsing Polychondritis Disease Activity Index (RPDAI) has been designed to assess RP in a standardised manner, taking into account the disease manifestations over a 28-day period. The RPDAI consists of 27 items with individual weights, ranging from 1 to 24, and reaches a maximum theoretical score of 265. More important, as there are no biologic disease activity markers in use, the RPDAI is a useful instrument to assess disease activity in multiple systems, as well as to assess therapeutic response, even in patients with no current inflammatory syndrome (Table 2). A damage index has been recently developed (Arnaud 2018, in press) to assess the disease-induced irreversible changes, with the aim to standardise the assessment and to facilitate studies in this rare disease.

**Patient-reported outcomes and quality of life indicators**

A recent study on more than 300 patients revealed that over 50% have visited the emergency room, had symptoms for more than 5 years and have consulted more than three physicians prior to diagnosis. The diagnostic delay (≥1 year) was attributed to coexisting fibromyalgia and to the lack of auricular and nasal chondritis and arthritis. Some patients may need long-term glucocorticoids and therefore...
Patients’ unmet needs

Most patients are in need of a continuity of care and follow-up, in particular through a multidisciplinary team approach. When there is extensive RP damage, many different appointments are needed for different specialists who do not always communicate with each other or are aware of the disease. There is also a need for a better diagnostic approach, as well as a team that helps patients after the diagnosis with reasonable adjusted advice about employment and disability. It is mainly the care and therapeutic approach that is of most importance for patients. There is a lack of knowledge, not enough studies available, as well as no specialised clinical and research teams available for most patients. RP damage can have a big impact on the well-being of patients. The patients, caregivers and medical caregivers therefore need to be further educated to help in the assistance of patients with RP. The ERN ReCONNET could help play a major role in this matter.

CONCLUSION

This systematic review shows that there are currently no CPGs available for RP. Given the limited data on the pathogenesis of the disease, the association of clinical and laboratory manifestations or imaging strategies, we were able to identify several major unmet needs. To the extreme rarity of the disease, patients and physicians should work together within the frame of the ReCONNET network to derive valuable evidence for the derivation of literature-informed CPGs.

REFERENCES


32. Torrelo A. CANDLE Syndrome as a paradigm of proteasome-related autoinflammation. *Front Immunol* 2017;8:927.


