Cryopyrin-Associated Periodic Syndromes and Treatment Options

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Abstract

Cryopyrin-associated periodic syndromes (CAPSs) are a growing family of autoinflammatory diseases, also known as periodic fever syndromes. There are three forms of CAPS: (1) Familial Cold autoinflammatory syndrome or familial cold urticaria, (2) Muckle-wells syndrome, and (3) neonatal-onset multisystem inflammatory disease or chronic infantile neurological cutaneous articular syndrome. Genetic mutations in the NLRP3 gene were found to be present in most patients. The foremost common findings between all the CAPS disorders are rash, fever which is sometimes present at birth or in early childhood, joint problems, and conjunctivitis. More extreme forms of CAPS include more persistent inflammation that can cause hearing loss and meningitis and can lead to mental and developmental delays. Drugs for CAPS target the source of inflammation – which is the over-production of interleukin 1β by modified cryopyrin inflammasomes. Three drugs are used to treat CAPS: Rilonacept, canakiumab, and anakinra. With these drugs, the prognosis is greatly improved, with most patients having less frequent episodes, decreased buildup of amyloid in the body, and extended life of severe cases up to adulthood.

Introduction

Cryopyrin-associated periodic syndromes (CAPSs) are a growing family of autoinflammatory diseases, also known as periodic fever syndromes. Autoinflammatory diseases are a group of disorders characterized by repetitive episodes of systemic and organ-specific inflammation. Unlike autoimmune diseases which are caused by activation of the specific immune system, people with autoinflammatory diseases do not produce autoantibodies or antigen-specific lymphocytes, instead, these diseases are caused by genetic mutations in molecules that regulate the innate immune response [1], [2]. There are three forms of CAPS: (1) Familial cold autoinflammatory syndrome (FCAS) or familial cold urticaria, (2) Muckle-Wells Syndrome (MWS), and (3) Neonatal-onset multisystem inflammatory disease (NOMID) or chronic infantile neurological cutaneous articular syndrome (CINCA). Genetic mutations in the NLRP3 gene were found to be present in most patients [3], [4], [5], [6], [7], [8], [9]. These results are connected with the episodes of fever and damage to the body’s cells and tissues. The foremost common findings between all the CAPS disorders are rash, fever which is sometimes present at birth or in early childhood, joint problems, and conjunctivitis. More extreme forms of CAPS include more persistent inflammation that can cause hearing loss and meningitis and can lead to mental and developmental delays.

Epidemiology

CAPS is very rare diseases. The incidence in Germany is 0.34 per 10^6 persons-years [10]. Based on that, research in Republic of Macedonia should be 1–2 new patients every 2 years.
FCAS

This disorder has autosomal dominant inheritance, and genetic mutation is on the NLRP3 gene. Most of the patients with FCAS can have a normal life and have children so they can pass the genetic mutation through generations. Characteristics of the disorder are recurrent episodes of fever and rash that occurs 1–2 h after exposure to cold temperature. The start of the disease is under 6 months of age, or sometimes in early childhood. Some patients have a rash display at birth. Many patients have rash each day, but seriously attacks happened after exposure to cold temperature. Sometimes after the occurrence of the rash, there is elevated temperature and arthralgia. Some patients also have abdominal pain, conjunctivitis, sweating, and headache. The duration of the episodes is less or around 24 h. Treatment incorporates warming of the patient, non-steroid anti-inflammatory drugs, corticosteroids, and IL-1 blocking agents to prevent cellular uptake of IL1 ß [4], [11].

MWS

MWS is a form of CAPS that is caused by is autosomal dominant genetic mutations on the NLRP3 gene. Most families with CAPS have FCAS or MWS and affected individuals of the family frequently share common symptoms. People with MWS have been able to have children which have led to the syndrome being present in some families for many generations. Chronic and recurrent hives on the entire body are present during early childhood. Fevers start in early childhood and are related to flare-ups of symptoms like rashes, joint pains, headaches, and eye inflammation. These symptoms usually last 1–2 days and can be activated sometimes by cold temperatures, stress, or exercise.

Joint pains are not associated with tissue and cartilage damage. Recurrent conjunctivitis is often a problem with MWS, and many patients also have a haze on their corneas. Hearing loss occurs in most patients during early adolescence. Due to chronic high levels of inflammation, some patients with MWS may develop amyloidosis, with presence of amyloid within the kidneys and liver [12], [13].

At present, the best medications found to help that the many sufferers of MWS have been the use of different IL-1ß blocking drugs to prevent the cellular uptake of IL-1ß.

NOMID/CINCA

NOMID/CINCA has the highest seriousness of chronic inflammation of all the forms of CAPS. It is also caused by mutations in the same genetic region of the NLRP3 gene. NOMID/CINCA symptoms are present at birth, or shortly after, with the hives rash which can increase during times of flare-ups of inflammation, fevers, pain in the joints, headaches, red eyes, and other symptoms. Swelling of the joints is sometimes with changes to the growth cartilage. Central nervous system symptoms may include headaches, stiffness of the neck and nauscea, elevated spinal fluid pressures, chronic aseptic meningitis with elevated neutrophils, and eosinophils present in the cerebrospinal fluid. The majority of children have significant mental deficits. During inflammatory episodes, many people with NOMID/CINCA and other forms of CAPS can suffer from conjunctivitis which is not caused by infection, they also have chronic papilledema within the eyes which can lead to a serious loss of vision. Progressive deafness can occur in these patients in early childhood. Some patients have dysmorphic facial characteristics, such as a saddleback nose or frontal bossing, smaller teeth, and other dental anomalies. There is a significant risk for amyloidosis with elevated serum amyloid and kidney damage, and some patients also have enlarged liver and spleen [14], [15]. Treatment has been the use of various IL-1ß-blocking drugs.

The main clinical characteristics of CAPS are presented in Table 1.

### Table 1: Clinical characteristics of CAPS

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<th>Triggered by cold</th>
<th>Spontaneous recurrent</th>
<th>Triggers by cold</th>
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<tbody>
<tr>
<td>Familial cold urticaria</td>
<td>Muckle-wells syndrome</td>
<td>Neonatal-Onset multisystem inflammatory disease</td>
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**Key Symptoms**

- **Familial Cold Urticaria**: Recurrent episodes of symptoms such as hives, usually triggered by cold, and sometimes by cold temperatures, stress, or exercise.
- **Muckle-Wells Syndrome**: Recurrent episodes of fever and rash that occur 1–2 h after exposure to cold temperature, followed by joint pains, headaches, and eye inflammation.
- **Neonatal-Onset Multisystem Inflammatory Disease (NOMID/CINCA)**: Symptoms present at birth or shortly after, with a rash, fever, and other systemic symptoms.

**Laboratory Tests**

During acute attacks laboratory tests often reveal signs of non-specific inflammatory markers, such as elevated erythrocyte sedimentation rate (ESR), high C-reactive protein (CRP), anemia, leukocytosis with a high number of polynuclear leukocytes and thrombocytes, and elevated serum amyloid A (SAA) (in MWS and NOMID/CINCA). In the case of NOMID/CINCA, lumbar punction shows cerebrospinal fluid at high pressure with neutrophils and eosinophils and no infection, while blood elements are rarely present in MWS. The following tests may also be performed: Audiogram which shows high-frequency hearing loss, progressing to bilateral deafness, urinary protein to detect amyloidosis, and kidney biopsy to confirm the presence of amyloid deposits. Genetic testing can be performed using a commercial test for an NLRP3 gene.
mutation and should be done for all patients suspected to have CAPS, but should not be the only basis of diagnosis.

**Treatment**

Medications for CAPS target the main source of inflammation which is the over-production of IL-1β by altered cryopyrin inflammasomes. Three drugs that target IL-1 are approved by the US Food and Drug Administration for the treatment of CAPS. The short-acting recombinant IL-1 receptor antagonist named anakinra was approved for the treatment of patients with NOMID in 2012, and the two long-acting IL-1-blocking agents, rilonacept, and canakinumab were approved for the treatment of CAPS in 2008 and 2009.

Anakinra is a recombinant and non-glycosylated synthetic form of human IL-1Ra receptor antagonist (Il-1 Ra) and acts like the endogenous human IL-1RA to inhibit IL-1β binding to the IL-1 receptor type1 (IL-1 RI), which helps to inhibit the biologic activity of IL-1β in the body. The drug is administered as a daily subcutaneous injection. Rilonacept is an IL-1β (IL-1β) blocking medication that inhibits IL-1β by attaching and neutralizing IL-1β in the circulating blood. This can prevent the cells from being triggered to activate the increase production of inflammatory mediators and is given as a weekly subcutaneous injection. Canakinumab is a fully-humanized monoclonal antibody against IL-1β with affinity for human soluble IL-1β. It is given every 4–8 weeks by subcutaneous injection.

The goals of therapy in CAPS are (a) to improve the disease symptoms, to reduce the attack frequency and duration, (b) to diminish the systemic inflammatory markers in the blood as CRP, ESR, and SAA, and (c) to prevent progression of organ damage.

**Caps Treatment-clinical Studies**

Clinical studies assessing the efficacy of anakinra and later studies with the other IL-1-blocking agents show significant improvement in the clinical symptoms of CAPS and also improvement in inflammatory markers.

The efficacy of the anakinra has been assessed in case reports [16], as well as several clinical studies. The study on 26 NOMID patients observed for 36–60 months find clinical and laboratory response achieved and sustained in all patients [17]. The study of 12 patients with MWS phenotype up to 14 months on anakinra reported that all patients had a significant improvement at 2 weeks and at the final follow-up. Treatment induced complete and long-lasting resolution of fever, arthralgia/ arthritis, and improvement of conjunctivitis at 2 weeks in all patients. Skin rash resolved in 7 of 8 patients. Amyloidosis improved in 1 of 2 patients with confirmed amyloidosis. Hearing loss resolved in 1 and improved in 1 of 10 individual. Hearing loss worsened in two patients under treatment [18]. Studies in patients with NOMID showed that treatment with anakinra can invert organ inflammation such as aseptic meningitis, papilledema, and cochlear inflammation [19]. Rilonacept was better than to placebo in improving the primary (composite symptom score) and secondary (flare days, single-symptom scores, and disease activity) points in a 24-week study of 47 patients with MWS or FCAS phenotypes [20]. Rilonacept showed improved clinical symptoms induced by cold within days of administration in five patients with observed during 24 months follow-up [11]. Canakinumab achieved complete response in the open-label phase of a 48-week study of 31 patients with MWS and four patients with NOMID. In the randomized phase, 81% of the patients in the placebo group relapsed, whereas all patients in the drug group remained in remission [21].

The doses of IL-1-blocking therapy are needed to suppress systemic and organ inflammation depend on disease severity and the extent of organ involvement. For anakinra, higher doses up to 10 mg/kg/day [19] have been used, and higher dosing has been investigated for patients with more serious forms of CAPS.

**Prognosis**

With suitable medications, most of the patients with FCAS and MWS can have normal life, get married, and have children. NOMID/CINCA patients have more severe form of inflammation through their body which can cause some permanent damage. If early diagnosis and treatment with medications are started, the prognosis is greatly improved, with most patients surviving into adulthood.

**Conclusion**

CAPS is a growing family of autoinflammatory diseases, also known as periodic fever syndromes. CAPS is caused by genetic mutations in the NLRP3 gene that regulates the innate immune response which results in a hyperactive cryopyrin protein and an increased inflammatory response, leading to overproduction of pro-inflammatory cytokine IL-1β.
These changes result in episodes of fever and damage to the body’s cells and tissues. The foremost common findings between all the CAPS disorders are rash, fever which is sometimes present at birth or in early childhood, joint problems, and conjunctivitis. More extreme forms of CAPS include more persistent inflammation that can cause hearing loss, meningitis and can lead to mental and developmental delays.

With medications that target the main source of inflammation which is the over-production of IL 1ß, anakinra, rilonacept, and canakinumab, prognosis is greatly improved, with most patients having less frequent episodes of the disease, preventing amyloidosis, and for severe cases – surviving into adulthood.

References


