Refractory systemic onset juvenile idiopathic arthritis: current challenges and future perspectives

William G. Ambler, Kabita Nanda, Karen Brandt One, and Susan Shenoi

ABSTRACT
Systemic juvenile idiopathic arthritis (SJIA) is a rare disease with distinct features not seen in other categories of juvenile idiopathic arthritis. In recent years, advances in the understanding of disease immunopathogenesis have led to improved targeted therapies with significant improvement in patient outcomes. Despite these advances, there remain subsets of SJIA with refractory disease and severe disease-associated complications. This review highlights existing options for treatment of refractory SJIA and explores potential future therapeutics for refractory disease.

KEY POINTS:
- Despite targeted Interleukin IL-1 and IL-6 inhibitors a subset of SJIA remains refractory to therapy. About 1 in 7 SJIA patients will be refractory to targeted IL-1 or IL-6 therapy.
- There is no current agreed upon definition for refractory SJIA and we propose in this review that refractory SJIA is presence of active systemic or arthritic features despite treatment with anti-IL-1 or anti-IL-6 therapy or disease requiring glucocorticoids for control beyond 6 months.
- SJIA disease associated complications include presence of associated macrophage activation syndrome (MAS), interstitial lung disease (ILD) or amyloidosis and management of each differs.
- Refractory SJIA treatment options currently include additional conventional synthetic disease modifying anti-rheumatic drugs (csDMARDS), biologic (bDMARDS), combination biologic therapy, targeted synthetic (tsDMARDS) or other immunomodulatory therapies.

Background
The International League of Associations for Rheumatology (ILAR) classifies systemic onset juvenile idiopathic arthritis (SJIA) as occurring in children under age 16 with fever for at least 2 weeks, of which there must be 3 consecutive days of quotidian fever, arthritis in one or more joints and with at least one of the following: evanescent erythematous rash, generalised lymphadenopathy, hepatomegaly, splenomegaly, or serositis [1]. Though most forms of JIA are conceptualised as autoimmune diseases, SJIA uniquely has many features of autoinflammatory disease. This has led to a paradigm shift in the treatment of patients, specifically due to the recognition of two critical cytokines in disease pathophysiology. Successful trials using Interleukin (IL)-1 and IL-6 inhibition have led to their widespread use and have dramatically improved patient outcomes [2,3]. IL-1 or IL-6 inhibitors are now recommended as first line therapy [4–7]. Multiple long term outcome studies demonstrate that most patients (60–80%) can achieve disease remission or minimal disease activity [8–12]. Effective treatment of patients earlier in the disease course may change long-term outcomes with fewer patients progressing to chronic synovitis [13]. SJIA disease course varies with approximately 40% having a monocyclic disease course, 10% with a polycyclic course, and 50% with a chronic course [14–16]. Patients with a monocyclic course have a brief duration of active disease (usually less than 6 months) and have excellent outcomes. Prior to the advent of targeted biologic therapies, patients
with a chronic disease course often went on to have severe, erosive, polyarticular arthritis often requiring long term glucocorticoids for arthritis control resulting in associated glucocorticoid toxicity [15,17]. Even in the post targeted biologic era refractory disease in 20% with ongoing systemic symptoms and/or synovitis poses a significant challenge. Additionally, association of SJIA with a rare interstitial lung disease (ILD) in certain high-risk patients can pose significant treatment challenges with morbidity and mortality. Macrophage activation syndrome (MAS), a life-threatening cytokine storm, is also a complication in a subset of patients. This review proposes a definition for refractory SJIA and reviews existing options for management of refractory SJIA. We also discuss SJIA associated complications including MAS, ILD and amyloidosis and discuss their management.

**Pathophysiology**

Understanding the immunologic mechanisms of SJIA has been and will continue to be critical to the development of effective treatments. The innate immune system is heavily implicated in disease pathophysiology. Myeloid cells are increased in number and activation status in the peripheral blood of SJIA [18,19]. Similarly, innate cytokines and alarmins are elevated in patients’ sera [20]. Pascual et al. demonstrated that SJIA sera can increase monocyte transcription of inflammatory cytokines, including IL-1β [21]. Activated monocytes from patients with SJIA release significantly more IL-1β than controls. These findings led to the successful treatment of many SJIA patients with anakinra, an IL-1 receptor antagonist. Subsequently, IL-1 blockade with canakinumab has been further studied and has gained United States Food and Drug Administration (US FDA) approval for the treatment of SJIA. Approximately 60% of patients can achieve remission on IL-1 antagonism as first line therapy [13,22]. IL-6 has also been implicated in SJIA, with increased IL-6 and IL-10 gene expression in SJIA monocytes and B cells compared to controls [23]. IL-6 blockade has similarly been effective, and tocilizumab is US FDA approved for SJIA treatment.

Though innate immune pathways are clearly important, recent evidence points to a role of the adaptive immune system as well. Ombrello et al. conducted GWAS on a large cohort of SJIA and demonstrated that MHC class II alleles portend the largest risk for SJIA development [24,25]. Further, the gene architecture of SJIA is divergent from other JIA categories [25]. These findings are supported by studies demonstrating the importance of T cells in chronic synovitis [26,27]. By analysing patients’ T cells, Henderson et al. demonstrated that different T cell subsets were present in different disease stages [27]. In the early inflammatory stage of disease, genuine IL-17 producing regulatory T cells were expanded. In patients that developed chronic synovitis, IL-17 producing effector CD4 cells were expanded. Patients who achieved early remission with IL-1 blockade failed to expand IL-17 effector CD4 T cells. This provides evidence to the theory of a \"window of opportunity\" where the early inflammatory milieu might favour an adaptive immune response leading to chronic synovitis [28]. Further, the authors hypothesise that medications targeting IL-17 (i.e. secukinumab) may be a logical treatment option for refractory synovitis [27].

MAS develops in 10–30% of patients with SJIA [29,30]. The reader is referred to several in depth reviews on this topic [29,31]. The \"cytokine storm\" of MAS is driven by activated macrophages and cytotoxic T cells [32]. Key findings have demonstrated the importance of IL-18 and interferon gamma (IFN-γ) in MAS. IL-18 is an inflammasome generated cytokine which induces IFN-γ release from T cells [33]. IL-18 is elevated in most patients with SJIA, however IL-18 is highest in those who develop MAS and rises further in active MAS [34]. Notably, IL-18 distinguishes MAS from primary hemophagocytic lymphohistiocytosis (HLH), in which a similar cytokine storm occurs usually in the setting of mutations in cytotoxic T cell killing [35]. IFN-γ is a common downstream cytokine released in both MAS and HLH. IFN-γ is required for HLH pathophysiology and blocking IFN-γ with emapalumab is approved for the treatment of primary HLH [36,37].

**Refractory SJIA**

For the purpose of this review refractory SJIA is defined as failure to respond to IL-1 and IL-6 inhibitors or need for ongoing treatment with long term glucocorticoids (beyond 6 months) with persistence of systemic and/or arthritic features (Table 1). If a patient does have an adequate response to one of the previously mentioned cytokine inhibitors (IL-1 or IL-6 inhibitor) the other should be used. Refractory SJIA should only be determined after inadequate responses of both IL1 and IL6 blockade (though not necessarily simultaneous use). In the current treatment era with wide availability of effective medications with different targets, inability to taper glucocorticoids should be considered refractory disease due to long term toxicity, regardless of disease activity [7,38]. Another recent
review proposes definitions for different subsets of refractory patients (refractory arthritis, refractory MAS, and ILD) [39]. These could be meaningful partitions of refractory subtypes given the likelihood of different immune drivers in patients with these different refractory phenotypes. For this review we separate out refractory SJIA (as defined above) and SJIA associated complications (including MAS, SJIA associated ILD and amyloidosis). A universally accepted definition will be helpful in order to design future outcome and treatment studies.

In the SJIA literature there exist two different time spans: the pre-biologic era, defined as the period prior to availability of anti-IL-1 and anti-IL-6 agents, and the post-biologic era. Two broad groups are used to identify refractory disease: 1) Lack of efficacy or clinical response including no response to medication, and incomplete response to medications (pertaining to either the systemic or arthritic features of SJIA); 2) Severe adverse reactions or side effects to medication necessitating discontinuation or change of therapy. Also, biologics are still not widely available in all countries. The IL-1 inhibitor canakinumab was approved by the US FDA in 2013 for the treatment of SJIA in children over 2 years of age [3]. The other anti-IL-1 agents, while not US FDA approved, have been used off label in SJIA, including anakinra [40] and rilonacept [41]. Rilonacept is a fusion protein which binds to IL-1 (IL-1 trap) and has been shown to be effective in a phase II trial with 74% of patients meeting the primary outcome of an ACR30 response at 4 weeks [41]. Ultimately the manufacturers did not choose to seek US FDA approval for SJIA and it thus remains largely unavailable. Tocilizumab was US FDA approved for SJIA in children ≥ 2 years of age in 2011 [2]. In the canakinumab trial, the primary end point was met with 83% of patients achieving an ACR30 response at 15 days (compared to 10% in placebo arm). At the end of the withdrawal phase 82% achieved an ACR70 response and 62% had clinical inactive disease (CID). In the tocilizumab trials, 85% of patients achieved the primary outcome of an ACR30 response at 12 weeks (compared to 24% in placebo arm) and 59% achieved an ACR90 response at the end of the open label extension at 52 weeks. The ACR30 is a validated mark of minimal improvement to assess efficacy for drug approval [42]. Clinically, however, an ACR30 response is a minimal response, and the expectation is for most patients is to achieve inactive disease or low disease activity. Despite these dramatic responses, there were still approximately 15–20% of patients with no response to single drug biologic therapy and approximately 40% who continued to have active disease while on single drug biologic therapy. It should be noted that most patients in both trials had long-standing disease. Real world data in the biologic era demonstrates that most patients can achieve inactive disease at long term follow up. In a Canadian inception cohort (ReACCh-Out), 70% of SJIA achieved CID by 2 years and 85% by 5 years [10]. Similar findings were seen in other cohorts [8,43]. A long term follow up (18 year follow up) study in a Nordic cohort showed 3/13 (23%) patients continued to have active disease [11]. This study is somewhat limited by the few patients with SJIA followed. Even in the post-biologic era, approximately 1 in 7 children will continue to have long-term disease activity.

### Treatment options for refractory SJIA

#### Conventional synthetic (cs) DMARDS

While methotrexate is often used as adjunctive treatment for SJIA, other csDMARDS are used with less frequency. While many of these medications are currently used infrequently, it can be reasonable to utilise them in certain situations particularly for refractory arthritic features of SJIA (Table 2). Thalidomide, an immunomodulatory drug has shown benefit in refractory SJIA. While the exact mechanism of action is not entirely understood, thalidomide decreases production of multiple inflammatory cytokines including Tumour necrosis factor alpha (TNF-α), IL-1, and IL-6 [68]. Three small case series and one small trial provide some anecdotal evidence for its use in refractory cases [44–47]. Patients in all three-case series had severe systemic and arthritic disease, refractory to a variety of different medications, and all required high doses of glucocorticoids. After treatment with thalidomide, patients were able to achieve either inactive

| Condition: | Definition: |
|------------|-------------|
| Refractory SJIA | Failure to respond to IL-1 and/or IL-6 inhibitors or need for ongoing treatment with long term glucocorticoids (beyond 6 months) with persistence of systemic and/or arthritic features |
| SJIA associated complications | - MAS |
|  | - SJIA ILD |
|  | - SJIA amyloidosis |
disease or low disease activity while lowering or discontinuing glucocorticoids. The single trial assessed the utility of thalidomide in refractory disease in 13 patients from 4 different centres. Eleven of the 13 patients had improvements in erythrocyte sedimentation rate, anaemia, at least 50% improvement in disease score, and were able to decrease glucocorticoid usage [47]. While thalidomide is a well-known teratogen and can cause neuropathy, it can still be considered, particularly in resource limited settings [68]. Lenalidomide, a thalidomide analogue with significantly less neurotoxicity, may be another option as well. Though there are currently no published reports on efficacy or safety, this author and others have used it successfully in refractory SJIA [16].

Calcineurin inhibitors, particularly cyclosporine, have been used for decades in the treatment of SJIA. Cyclosporine is an alkylating agent, particularly effective at eliminating reactive lymphocytes and inhibiting calcineurin. In a retrospective study from an area where IL-1 and IL-6 inhibitors are prohibitively expensive, cyclosporine led to significant reduction in systemic symptoms and glucocorticoid dosage [53]. Of 13 patients with steroid dependent and refractory SJIA, all had resolution of systemic symptoms within one month of initiation of cyclosporine along with normalisation of serum markers of inflammation. Glucocorticoid dose was able to be decreased by an average 50% at a median of about 2 months. Tacrolimus has not been used as frequently, but in case reports has demonstrated similar efficacy [48,49]. In one recent single centre retrospective case series from Shanghai, of 6 patients with SJIA treated with tacrolimus, all had improvements in laboratory parameters and were able to lower glucocorticoids [48]. Voclosporin, a newer agent has recently gained approval for adult lupus and potentially could be another therapeutic option in the future.

## Table 2. Medications used in treatment of refractory SJIA.

| Medication used for refractory SJIA | Mechanism of action | Relevant literature | Comments |
|-----------------------------------|---------------------|---------------------|----------|
| csDMARDs                           |                     |                     |          |
| Thalidomide                        | Decreases TNF-α and IL-6 secretion, decreases angiogenesis | [44–47] | Exact mechanism of action is not well described |
| Tacrolimus                         | Inhibits T cell activation and proliferation via inhibiting calcineurin | [48,49] | Volcosporin is newer calcineurin inhibitor with more stable pharmacokinetics |
| Cyclophosphamide                   | Alkylating agent, particularly effective at eliminating reactive lymphocytes | [50,51] |          |
| Cyclosporine                       | Inhibits T cell activation and proliferation via inhibiting calcineurin | [38,52,53] | Also effective add on therapy for MAS and has been used for SJIA ILD |
| bDMARDs                            |                     |                     |          |
| Rituximab                          | Elimination of non-stem B cells | [54–56] | May be effective directly or indirectly through B cell antigen presentation to T cells |
| TNF-α inhibitors                   | Inhibits TNF-α signalling | [57,58] | Pleiotropic pro-inflammatory cytokine |
| Abatacept                          | Inhibits T cell co-stimulation by blocking costimulatory molecules CD80/86 on APCs | [59,60] |          |
| Tadekinig Alfa                     | Blocks IL-18 signalling by binding unbound “free” IL-18 | [34,61] |          |
| tsDMARDs                           |                     |                     |          |
| Jakinib                            | Inhibits signalling through JAK/STAT pathway, inhibiting multiple cytokines | [62,63] | Different jakinibs have reported differing effect on different JAK/STAT pathways *via in vitro* studies [64] |
| Medications used for MAS           |                     |                     |          |
| Emapalumab                         | Inhibits IFN-γ signalling via direct binding | [65] | Dose titrated bases on clinical effectiveness. Can measure effectiveness of blockade by measurement of CXCL9. |
| Anti-thymocyte globulin            | Polyclonal antibodies that eliminate T cells | [66] |          |
| Etoposide                          | Topoisomerase II inhibitor | [67] | Effective at eliminating activated and dividing T cells. Effective at lower doses then in the HLH 94 protocol |
| Medications used for ILD           |                     |                     |          |
| MMF                                | Inhibits inosine monophosphate dehydrogenase leading to decreased lymphocyte activation and proliferation | [38] | Effective in slowing lung fibrosis in scleroderma |
| Cyclosporine                       | See above           |                     |          |
| Jakinibs                           | See above           |                     |          |

**Legend:** APC: Antigen presenting cell, JAK: Janus Kinase, STAT: Signal Transducer and Activator of Transcription; csDMARD: Conventional Synthetic DMARD; bDMARD: Biologic DMARD; tsDMARD: targeted synthetic DMARD; ILD: Interstitial Lung Disease; HLH: Hemophagocytic Lymphohistiocytosis.
refractory SJIA [50,51]. In a small trial prior to the biologic era, 4 patients with severe refractory disease had positive responses to intravenous (IV) CYC given monthly [50]. All patients were able to discontinue glucocorticoids and 3 achieved inactive disease. In another small open label trial of refractory SJIA (pre-biologic era) 18 patients with SJIA had significant improvement in both systemic and articular symptoms after started on IV CYC at 400 mg/m² every 3 months [51]. CYC is broadly immunosuppressive and has toxic effects on myeloid cells and well as lymphocytes, thus its efficacy in SJIA is not surprising [71]. However, CYC has many side effects including increased risk of malignancy and the risk of infertility limiting its usefulness in the long term. The role of CYC in the biologic era is unclear but could be considered in very select cases.

**Biologic (b) DMARDS**

TNF-α inhibitors have been tried in refractory SJIA [57]. In general TNF-α inhibitors are less effective for the systemic features of the disease but may help with arthritic features of SJIA [57]. Kimura et al. highlight this suboptimal response in their paper that surveyed 82 SJIA patients on etanercept. Response was poor in 45%, 33% discontinued etanercept due to lack of response or flare and 2.4% developed MAS while on etanercept [58].

Rituximab, a monoclonal antibody that depletes B cells, has shown some efficacy. Forty-four patients with refractory SJIA (prior to the availability of IL-1 or IL-6 blockade) were treated with rituximab in a prospective, open label, uncontrolled trial [54]. Patients were treated every 24 weeks if there was ongoing disease activity. After 24 weeks, 98% achieved an ACR30 and 25% of patients were able to achieve remission. All but 2 patients had resolution of fever by 24 weeks. In 25 patients that had continued to be followed for 96 months, 24 achieved remission. There are also two case reports that describe significant improvement in both systemic and arthritic features in refractory cases [55,56]. While the role of B cells in the pathogenesis of SJIA is unclear there is some evidence that depleting them can be of benefit. While there are no case reports, we postulate that obinutuzumab another B cell depleting agent might be similarly beneficial in refractory SJIA.

IL-18 is another promising target as IL-18 is elevated in SJIA and correlates with the risk of developing MAS. IL-18 binds to IL-18 receptor protein and it is the unbound fraction of IL-18 which actively signals [72]. Tadekinig alfa is a recombinant IL-18 receptor protein which functionally inhibits IL-18 signalling. A phase II trial showed benefit in patients with Adult Onset Still’s Disease (AOSD) [61], which could potentially translate to patients with SJIA given that AOSD is essentially identical to SJIA both in phenotype and pathophysiology [73,74]. There is single case report showing benefit of IL-18 inhibition in a child with refractory SJIA [75]. In this report IL-18 inhibition was given to 6-year-old with SJIA with recurrent MAS refractory to chronic high doses of steroids as well as numerous bDMARD and csDMARDs including IL-1 and IL-6 inhibition. After starting IL-18 inhibition the patient improved and was able to considerably lower daily steroid usage. Despite having several episodes of MAS on IL-18 therapy, the episodes were significantly less severe than prior episodes. IL-18 binding protein is currently in clinical trials for X-linked inhibitor of apoptosis inhibitor (XIAP) deficiency and NLR Family CARD Domain Containing 4 (NLRC4) gain of function mutations (NCT03113760). These diseases are characterised by elevated IL-18 and are frequently complicated by MAS/HLH. It will be interesting to see if this medication will be beneficial in patients with SJIA, particularly in children with MAS.

Other IL-6 inhibitors which likely are similarly efficacious as tocilizumab exist. Siltuximab, an IL-6 binding chimeric monoclonal antibody, was effective in a 20-year-old patient with SJIA who failed multiple therapies including IL-1 blockade and was unable to tolerate tocilizumab because of infusion reactions [76]. Sarilumab is an alternate IL-6 binding monoclonal antibody currently approved for rheumatoid arthritis that is being studied for its utility in SJIA (NCT02991469). Although these alternative medications have not been rigorously studied in SJIA, and thus currently lack specific approval, it is likely that they are effective. It is unclear if patients that fail tocilizumab might respond to a different IL-6 inhibitor and more data is needed to evaluate this possibility.

Combination biologic therapy may be beneficial in some refractory patients. One case series reports a 17 year old patient with SJIA with persistent systemic and arthritic disease on tocilizumab monotherapy and anakinra monotherapy (with concomitant methotrexate and glucocorticoids) [77]. However, when low doses of tocilizumab (2 mg/kg) and anakinra were used simultaneously, the patient achieved inactive disease. Abatacept has been used with success in combination with anakinra in a case series [59]. Abatacept, a soluble cytotoxic T-lymphocyte associated protein 4 fused the Fc portion of immunoglobulin (CTLA-4-Ig),
inhibits T cell co-stimulation [60]. In this case series, 4 patients with glucocorticoid dependent disease despite use of anakinra, methotrexate, and cyclosporine had significant improvement in disease control and were able to reduce glucocorticoid dosage with the addition of abatacept. Given the importance of T cells in the development of chronic synovitis in SJIA, this may be a promising option as add-on therapy. Despite the more frequent use of combination therapy, there is minimal data on the safety or efficacy of combination therapy.

**Targeted synthetic (ts) DMARDs**

Janus kinase inhibitors (Jakinibs) are a promising option based on the profile of cytokine signalling pathways that are inhibited. A few such cytokines are IFN-γ and IL-6, which as previously mentioned are important in the pathophysiology of SJIA. An in depth review regarding the theoretic benefits of Jakinibs in SJIA is detailed in a recent review by Verweyen and Schulert [78]. There is currently a clinical trial using tofacitinib for treatment of children with SJIA with active systemic features (NCT03000439). While mechanistically Jakinibs are intriguing, there are only few case reports published of their use in the treatment of SJIA. One case report from China describes a 13-year-old girl with recalcitrant systemic and arthritic manifestations of SJIA despite long term glucocorticoids, methotrexate, and TNF-α inhibition [78]. The patient suffered significant glucocorticoid toxicity with vertebral compression fractures and growth retardation. Tofacitinib was started due to family’s refusal of tocilizumab and unavailability of IL-1 antagonism in mainland China. The patient had complete remission by 3 months and discontinued glucocorticoids by 6 months. Though this patient may not have been refractory if given IL-1 or IL-6 inhibitors, this case report does show the potential benefit of Jakinibs in SJIA. Success using Jakinibs in AOSD, may translate to SJIA given their previously mentioned similarities [73,74]. In a case series of 14 patients from China who received tofacitinib, 7 achieved remission and the remainder were partial responders (using the modified Pouchot’s systemic score) [62]. Only 2 patients had received tocilizumab prior to receiving tofacitinib and both were partial responders. Another case report demonstrates success with the use of ruxolitinib in a 4-year-old SJIA complicated by drug reaction to anakinra and canakinumab with development of ILD [63]. After 15 months of follow up the patient had resolution of systemic symptoms, normalisation of C-reactive protein, improvement in oxygen saturation and lung imaging, and was able to lower glucocorticoid usage. Results from the tofacitinib trial for SJIA (NCT03000439) are eagerly awaited as it may expand the existing armamentarium of therapeutic options for SJIA and will likely reduce the number of refractory disease patients.

**Stem cell therapies**

Bone marrow transplantation has been used for refractory disease in both the pre- and post-biologic eras. Earlier work from several groups in Europe studied T cell depleted autologous haematopoietic stem cell transplantation (HSCT) as a strategy to induce remission [79,80]. The Dutch group (Brinkman et al.) used this strategy in 18 patients with refractory SJIA [79]. While 6 achieved remission (per Wallace criteria) and 5 more had a partial response (> ACR 50%) there was significant morbidity and mortality. Two patients died from MAS early post-transplant, which had been reported in prior reports. An additional 2 patients died after one year from viral infection. A group from the United Kingdom had a case series of 7 patients that underwent HSCT and 4 achieved long lasting remission, 2 had disease relapse, and 1 died from viral infection [80]. While autologous HSCT can result in long lasting remission, a large proportion have disease recurrence and there is significant morbidity and mortality with rates of approximately 15–22% [79,80]. Since MAS accounts for a large portion of the mortality post-transplant it is critical to attain minimal disease activity prior to transplant in order to improve transplant related outcomes [81].

Allogeneic HSCT has a higher chance of lifelong cure, however there is also risk of graft versus host disease (~20%) [82]. Silva et al. published the results of performing allogenic HSCT on 16 patients with refractory JIA (11 of whom had SJIA). Many of these patients were refractory to IL-1 and IL-6 blockade. These patients received reduced intensity conditioning with fludarabine based regimens and alemtuzumab. Of the 11 SJIA patients reported, 1 died due to pulmonary haemorrhage, 1 had disease recurrence necessitating reintroduction of medications, and 9 achieved clinical remission. Post-transplant complications included graft versus host disease in 5 patients and viral infections (several severe) in most patients. Early MAS was not observed as it was in autologous HSCT [82]. Allogeneic stem cell transplantation may have a role in refractory disease; however, this procedure comes with known morbidity and mortality risk [82].
Mesenchymal stromal cell (MSC) infusions represent another possible therapeutic option for refractory SJIA cases. MSC are non-embryonic, fibroblast-like stem cells that are isolated from blood or bone marrow and have immunoregulatory properties [83]. They can inhibit T cells, B cells, natural killer (NK) cells, and dendritic cells and can activate Regulatory T cells. They also express very little major histocompatibility complex class I (MHC class I) and thus do not invoke an alloimmune reaction. Swart et al. did a proof-of-concept phase 1 b trial with 6 refractory JIA patients (1 with SJIA). The patient with SJIA was refractory to multiple therapies including IL-1 and IL-6 blockade. This patient was able to wean glucocorticoids, however developed MAS when attempting to discontinue tocilizumab. While clearly warranting additional study, MSC may have a future role in SJIA and other immune mediated disease.

Management of SJIA complications

Macrophage activation syndrome

MAS, a life-threatening manifestation of SJIA, can be difficult to treat. Treatments vary based on severity, but standard treatments include high doses of IV glucocorticoids, cyclosporine, and high doses of anakinra [31,84]. The ACR recently published guidelines for management of SJIA associated with MAS [6]. Intravenous immunoglobulin (IVIG) has been used with mixed success in several case reports and could be considered in refractory cases [85,86]. IFN-γ is a critical cytokine in HLH/MAS pathophysiology [36,87]. Emapalumab, a monoclonal antibody that binds IFN-γ, is effective and has gained approval for the treatment of refractory primary HLH [37]. Preliminary clinical trial data (NCT03311854) suggests that IFN-γ blockade is safe and effective. Nine patients with refractory MAS to high dose IV glucocorticoids that have been treated with emapalumab have been analysed thus far. Of these patients, 4 had prior treatment failure with cyclosporine and 4 with anakinra. All patients had resolution of clinical and laboratory parameters of MAS with emapalumab treatment [65]. IFN-γ blockade represents a promising new drug candidate in the armamentarium for treatment of refractory MAS in SJIA. Anti-thymocyte globulin (ATG) and etoposide, both medications effective at depleting T cells, have been used for refractory cases with success [66,67]. Lower doses of etoposide (when compared with doses per the HLH-94/2004 protocol) are likely efficacious for refractory MAS [67]. In 7 patients with refractory MAS (including 2 with SJIA), weekly etoposide at 50–100mg/m² was used successfully in all patients [67]. In the acute setting there may be a role for leukocytapheresis, particularly for refractory MAS [88–90]. However, application of this is limited due to the temporary nature of action and need for central line access limiting use to tertiary care settings.

Systemic JIA associated lung disease

In the last several decades there are increasing numbers of a subset of children with SJIA that developed a severe and often fatal lung disease with a mortality rate around 60% [91]. These patients develop a pulmonary alveolar proteinosis-like (PAP) ILD. Compared to SJIA children without ILD, those with ILD have a younger age at disease onset, more frequent MAS, atypical rashes, less prominent arthritis, and frequently develop drug reactions to biologic medications including tocilizumab, anakinra, and/or canakinumab. The exact aetiology of this complication is unclear. Saper and Ombrello et al. showed that there is a strong association of this phenotype with a certain HLA haplotype (HLA-DRB1*15 alleles) [92]. Schulert et al. demonstrated that these patients have elevated IL-18 and the IFN-γ stimulated gene CXCL9, which would be expected in a group that develops more frequent MAS [93]. Lung biopsies demonstrated T cell infiltration with upregulation of gene transcription in both IFN-γ pathways and T cell activation pathways. Based on these observations, a variety of different treatments have been used in attempt to improve the outcomes of these patients. Some clinicians discontinue biologic therapy, and in addition to high doses of glucocorticoids, utilise drugs that target both T cells and/or IFN-γ including cyclosporine, mycophenolate mofetil, and Jakinibs with reported success [38,63]. This is an area of active research to both determine the mechanistic cause of PAP-like ILD and to how to most effectively treat this group of patients.

Amyloidosis

Secondary amyloidosis (AA amyloid), caused by the deposition of serum amyloid A (SAA), is a life-threatening complication of prolonged systemic inflammation [94]. This usually manifests as renal dysfunction and has a high mortality rate. Prior to the biologic era, amyloidosis was seen in increased frequency in SJIA. In one long term follow up study in 2002 from a group in the UK, the rate of amyloidosis in SJIA was approximately 20%. Fortunately, with modern therapies, amyloidosis is now rarely seen in SJIA and recent
rates have not been reported. There are still case reports of AA amyloid, always in the context of chronically active disease [95]. Effective treatment of the underlying disease is critical to the treatment of amyloid, as reduction in systemic inflammation reduces production and deposition of SAA. Tocilizumab or other IL-6 inhibitors may be of particular benefit as IL-6 has been shown to be important in hepatic SAA production [95,96].

**Future directions**

Scientific discovery has led to radical improvements in SJIA outcomes through a deepened understanding of the immunobiology and pathophysiology of disease. Despite this there are still patients that do not respond to contemporary first line treatment with IL-1 or IL-6 inhibition. Patients with ILD also pose significant new challenges. Additional drugs will likely become available for use in our armamentarium to treat SJIA with further scientific advances. Jakinibs, IL-18 receptor blockade, and IFN-γ receptor blockade are agents that will hopefully prove to be safe and effective for patients with SJIA. There may be opportunities for personalised medicine based on individualised patient phenotype, dysregulated immune pathways, and genetics [97,98]. Translational and clinical studies will hopefully continue to lead to effective treatment discoveries to reduce the number of patients with refractory disease and to improve patient outcomes.

**Author contributions**

WA contributed to the literature review, the design of work, and wrote the original manuscript. KN contributed to literature review and critically revised the article. KO contributed to literature review, assisted in the design of the work, and critically revised the article. SS supervised the literature review, conceptualised, and designed the work, and critically revised the article. All authors approved the final manuscript approval.

**Disclosure statement**

Dr. Shenoi serves as a consultant to Pfizer. Dr. Nanda serves as a consultant for Azurity Pharmaceuticals and conducts research funded by Abbvie, Pfizer, and Bristol-Myers Squibb. There are no other disclosures.

**Data availability statement**

Data sharing is not applicable to this article as no new data were created or analysed in this study.

**Funding**

The author(s) reported there is no funding associated with the work featured in this article.

**References**

[1] Petty RE, Southwood TR, Manners P, International League of Associations for Rheumatology, et al. International league of associations for rheumatology classification of juvenile idiopathic arthritis: second revision, edmonton, 2001. J Rheumatol. 2004;31(2):390–392.

[2] De Benedetti F, Brunner HI, Ruperto N, et al. Randomized trial of tocilizumab in systemic juvenile idiopathic arthritis. N Engl J Med. 2012;367(25):2385–2395.

[3] Ruperto N, Brunner HI, Quartier P, et al. Two randomized trials of canakinumab in systemic juvenile idiopathic arthritis. N Engl J Med. 2012;367(25):2396–2406.

[4] DeWitt EM, Kimura Y, Beukelman T, Juvenile Idiopathic Arthritis Disease-Specific Research Committee of Childhood Arthritis Rheumatology and Research Alliance, et al. Consensus treatment plans for new-onset systemic juvenile idiopathic arthritis. Arthritis Care Res. 2012;64(7):1001–1010.

[5] Ringold S, Weiss PF, Beukelman T, American College of Rheumatology, et al. 2013 Update of the 2011 American college of rheumatology recommendations for the treatment of juvenile idiopathic arthritis: recommendations for the medical therapy of children with systemic juvenile idiopathic arthritis and tuberculosis screening among children receiving biologic medications. Arthritis Care Res (Hoboken). 2013;65(10):1551–1563.

[6] Onel KB, Horton DB, Lovell DJ, et al. 2021 American college of rheumatology guideline for the treatment of juvenile idiopathic arthritis: therapeutic approaches for oligoarthritis, temporomandibular joint arthritis, and systemic juvenile idiopathic arthritis. Arthritis Rheumatol. 2022;74(4):521–537.

[7] Onel KB, Horton DB, Lovell DJ, et al. 2021 American college of rheumatology guideline for the treatment of juvenile idiopathic arthritis: therapeutic approaches for oligoarthritis, temporomandibular joint arthritis, and systemic juvenile idiopathic arthritis. Arthritis Care Res. 2022;74(4):521–537.

[8] Yokota S, Itoh Y, Morio T, et al. Tocilizumab in systemic juvenile idiopathic arthritis in a real-world clinical setting: results from 1 year of postmarketing surveillance follow-up of 417 patients in Japan. Ann Rheum Dis. 2016;75(9):1654–1660.

[9] Janow G, Schanberg LE, Setoguchi S, CARRA Legacy Registry Investigators, et al. The systemic juvenile idiopathic arthritis cohort of the childhood arthritis and rheumatology research alliance registry: 2010- 2013. J Rheumatol. 2016;43(9):1755–1762.

[10] Guzman J, Oen K, Tucker LB, et al. The outcomes of juvenile idiopathic arthritis in children managed with contemporary treatments: results from the ReACCh-out cohort. Ann Rheum Dis. 2013;74(10):1854–1860.
[11] Glerup M, Rypdal V, Arnstad ED, the Nordic Study Group of Pediatric Rheumatology, et al. Long-Term outcomes in juvenile idiopathic arthritis: Eighteen years of Follow-Up in the Population-Based nordic juvenile idiopathic arthritis cohort. Arthritis Care Res. 2020;72(4):507–516.

[12] Horneff G, Schulz AC, Klotsche J, et al. Experience with etanercept, tocilizumab and interleukin-1 inhibitors in systemic onset juvenile idiopathic arthritis patients from the BIKER registry. Arthritis Res Ther. 2017;19(1):256.

[13] Nigrovic PA, Mannion M, Prince FHM, et al. Anakinra in first-line disease-modifying therapy in systemic juvenile idiopathic arthritis: report of forty-six patients from an International multicenter series. Arthritis Rheum. 2011;63(2):545–555.

[14] Singh-Grewal D, Schneider R, Bayer N, et al. Predictors of disease course and remission in systemic juvenile idiopathic arthritis: significance of early clinical and laboratory features. Arthritis Rheum. 2006;54(5):1595–1601.

[15] Lomater C, et al. Systemic onset juvenile idiopathic arthritis: a retrospective study of 80 consecutive patients followed for 10 years. J Rheumatol. 2000;27(2):491–496.

[16] Fantini F, Gerlioni V, Gattinara M, et al. Remission in juvenile chronic arthritis: a cohort study of 683 consecutive cases with a mean 10 year followup. J Rheumatol. 2003;30(3):579–584.

[17] Bowyer SL, Roettcher PA, Higgins GC, et al. Health status of patients with juvenile rheumatoid arthritis at 1 and 5 years after diagnosis. J Rheumatol. 2003;30(2):394–400.

[18] Macaubas C, Nguyen K, Deshpande C, et al. Distribution of circulating cells in systemic juvenile idiopathic arthritis across disease activity states. Clin Immunol. 2010;134(2):206–216.

[19] Hinze CH, Fall N, Thornton S, et al. Immature cell populations and an erythropoiesis gene-expression signature in systemic juvenile idiopathic arthritis: implications for pathogenesis. Arthritis Res Ther. 2010;12(3):R123.

[20] Kessel C, Holzinger D, Foell D. Phagocyte-derived S100 proteins in autoinflammation: putative role in pathogenesis and usefulness as biomarkers. Clin Immunol. 2013;147(3):229–241.

[21] Pascual V, Allantaz F, Arce E, et al. Role of interleukin-1 (IL-1) in the pathogenesis of systemic onset juvenile idiopathic arthritis and clinical response to IL-1 blockade. J Exp Med. 2005;202(19):1479–1486.

[22] Vastert SJ, de Jager W, Noordman BJ, et al. Effectiveness of first-line treatment with recombiant interleukin-1 receptor antagonist in steroid-naive patients with new-onset systemic juvenile idiopathic arthritis: results of a prospective cohort study. Arthritis Rheumatol. 2014;66(4):1034–1043.

[23] Ogilvie EM, Khan A, Hubank M, et al. Specific gene expression profiles in systemic juvenile idiopathic arthritis. Arthritis Rheum. 2007;56(6):1954–1965.

[24] Ombrello MJ, Remmers EF, Tachmazidou I, International Childhood Arthritis Genetics (INCHARGE) Consortium, et al. HLA-DRB1*11 and variants of the MHC class II locus are strong risk factors for systemic juvenile idiopathic arthritis. Proc Natl Acad Sci U S A. 2015;112(52):15970–15975.

[25] Ombrello MJ, Arthur VL, Remmers EF, et al; British Society of Pediatric and Adolescent Rheumatology (BSPAR) Study Group, Inception Cohort of Newly Diagnosed Patients with Juvenile Idiopathic Arthritis (ICON-JIA) Study Group, Childhood Arthritis Prospective Study (CAPS) Group, Randomized Placebo Phase Study of Rilonacept in sJIA (RAPPORT) Investigators, Sparks-Childhood Arthritis Response to Medication Study (CHARMS) Group, Biologically Based Outcome Predictors in JIA (BBOP) Group. Genetic architecture distinguishes systemic juvenile idiopathic arthritis from other forms of juvenile idiopathic arthritis: clinical and therapeutic implications. Ann Rheum Dis. 2017;76(5):906–913.

[26] Kessel C, Lippitz K, Weinlage T, et al. Proinflammatory cytokine environments can drive interleukin-17 overexpression by γδ T cells in systemic juvenile idiopathic arthritis. Arthritis Rheumatol. 2017;69(7):1480–1494.

[27] Henderson LA, Hoyt KJ, Lee PY, et al. Th17 reprogramming of T cells in systemic juvenile idiopathic arthritis. JCI Insight. 2020;5(6):e132508.

[28] Nigrovic PA. Review: is there a window of opportunity for treatment of systemic juvenile idiopathic arthritis? Arthritis Rheumatol. 2014;66(6):1405–1413.

[29] Schuilert GS, Grom AA. Pathogenesis of macrophage activation syndrome and potential for cytokine-directed therapies. Annu Rev Med. 2015;66:145–159.

[30] Behrens EM, Beukelman T, Paesler M, et al. Occult macrophage activation syndrome in patients with systemic juvenile idiopathic arthritis. J Rheumatol. 2007;34(5):1133–1138.

[31] Grom AA, Horne A, Benedetti FD. Macrophage activation syndrome in the era of biologic therapy. Nat Rev Rheumatol. 2016;12(5):259–268.

[32] Grom AA, Passo M. Macrophage activation syndrome in systemic juvenile rheumatoid arthritis. J Pediatr. 1996;129(5):630–632.

[33] Novick D, Kim S, Kaplaniski G, et al. Interleukin-18, more than a Th1 cytokine. Semin Immunol. 2013;25(6):439–448.

[34] Yasin S, Fall N, Brown RA, et al. IL-18 as a biomarker linking systemic juvenile idiopathic arthritis and macrophage activation syndrome. Rheumatology (Oxford). 2020;59(2):361–366.

[35] Weiss ES, Girard-Guyonvarc’h C, Holzinger D, et al. Interleukin-18 diagnostically distinguishes and pathogenically promotes human and murine macrophage activation syndrome. Blood. 2018;131(13):1442–1455.

[36] Jordan MB, Hildeman D, Kappler J, et al. An animal model of hemophagocytic lymphohistiocytosis (HLH): CD8+ T cells and interferon gamma are essential for the disorder. Blood. 2004;104(3):735–743.

[37] Locatelli F, Jordan MB, Allen C, et al. Emapalumab in children with primary hemophagocytic lymphohistiocytosis. N Engl J Med. 2020;382(19):1811–1822.

[38] Canna SW, Schuilert GS, de Jesus A, NextGen 2019 Participants, et al. Proceedings from the 2(nd) next gen therapies for systemic juvenile idiopathic arthritis
and macrophage activation syndrome symposium held on October 3–4, 2019. Pediatr Rheumatol Online J. 2020;18(Suppl 1):53.

[39] Erkens R, Esteban Y, Towe C, et al. Pathogenesis and treatment of refractory disease courses in systemic juvenile idiopathic arthritis: Refractory arthritis, current macrophage activation syndrome and chronic lung disease. Rheum Dis Clin North Am. 2021;47(4):585–606.

[40] Quartier P, Allantaz F, Cimaz R, et al. A multicentre, randomised, double-blind, placebo-controlled trial with the interleukin-1 receptor antagonist anakinra in patients with systemic-onset juvenile idiopathic arthritis (ANAJIS trial). Ann Rheum Dis. 2011;70(5):747–754.

[41] Ilowite NT, Prather K, Lokhnygina Y, et al. Randomized, double-blind, placebo-controlled trial of the efficacy and safety of rilonacept in the treatment of systemic juvenile idiopathic arthritis. Arthritis Rheumatol. 2014;66(9):2570–2579.

[42] Lovell DJ, Ruperto N, Giannini EH, et al. Advances from clinical trials in juvenile idiopathic arthritis. Nat Rev Rheumatol. 2013;9(9):557–563.

[43] Sengler C, Klotsche J, Niewerth M, et al. Treatment of systemic juvenile idiopathic arthritis with tacrolimus. Eur J Pediatr. 2007;166(10):1053–1055.

[44] Wang D, Chen X, Li Z. Treatment of patients with systemic-onset juvenile idiopathic arthritis with tacrolimus. Exp Ther Med. 2017;13(3):2305–2309.

[45] Tanaka H, Tsuchiya K, Suzuki K, et al. Treatment of difficult cases of systemic-onset juvenile idiopathic arthritis with tacrolimus. Eur J Pediatr. 2007;166(10):1053–1055.

[46] Wallace CA, Sherry DD. Trial of intravenous pulse cyclophosphamide and methylprednisolone in the treatment of severe systemic-onset juvenile rheumatoid arthritis. Arthritis Rheum. 1997;40(10):1852–1855.

[47] Shawkat AV, et al. Repetitive use of pulse therapy with methylprednisolone and cyclophosphamide in addition to oral methotrexate in children with systemic juvenile rheumatoid arthritis—preliminary results of a longterm study. J Rheumatol. 1992;19(4):612–616.

[48] Ruperto N, Ravelli A, Castelli E, Paediatric Rheumatology International Trials Organisation (PRINTO), et al. Cyclosporine a in juvenile idiopathic arthritis. Results of the PRCSG/PRINTO phase IV post marketing surveillance study. Clin Exp Rheumatol. 2006;24(5):599–605.

[49] Pal P, Giri PP, Sinha R. Cyclosporine in resistant systemic arthritis - A cheaper alternative to biologics. Indian J Pediatr. 2019;86(7):590–594.

[50] Alexeeva EI, Valieva SI, Bzorova TM, et al. Efficacy and safety of repeat courses of rituximab treatment in patients with severe refractory juvenile idiopathic arthritis. Clin Rheumatol. 2011;30(9):1163–1172.

[51] Narváez D, Díaz-Tormé C, Juanola X, et al. Rituximab therapy for refractory systemic-onset juvenile idiopathic arthritis. Ann Rheum Dis. 2009;68(4):607–608.

[52] Kashner-Meron M, Uziel Y, Amital H. Successful treatment with B-cell depleting therapy for refractory systemic onset juvenile idiopathic arthritis: an evidence-based review. Paediatr Drugs. 2010;12(6):367–377.

[53] Kimura Y, et al. Etanercept treatment in patients with refractory systemic onset juvenile rheumatoid arthritis. J Rheumatol. 2005;32(5):935–942.

[54] RECORD JL, BEUKELMAN T, CRON RQ. Combination therapy of abatacept and anakinra in children with refractory systemic juvenile idiopathic arthritis: a retrospective case series. J Rheumatol. 2011;38(1):180–181.

[55] Ruperto N, Lovell DJ, Quartier P, et al. Abatacept in children with juvenile idiopathic arthritis: a randomised, double-blind, placebo-controlled withdrawal trial. The Lancet. 2008;372(9636):383–391.

[56] Gabay C, Fautrel B, Rech J, et al. Open-label, multicentre, dose-escalating phase II clinical trial on the safety and efficacy of tafekini alfa (IL-18BP) in adult-onset still’s disease. Ann Rheum Dis. 2018;77(6):840–847.

[57] Hu Q, Wang M, Jia J, et al. Tofacitinib in refractory adult-onset still’s disease: 14 cases from a single Centre in China. Ann Rheum Dis. 2020;79(6):842–844.

[58] Bader-Meunier B, Hadchouel A, Berteloot L, et al. Effectiveness and safety of ruxolitinib for the treatment of refractory systemic idiopathic juvenile arthritis like associated with interstitial lung disease: a case report. Ann Rheum Dis. 2022;81(2):e20-e20.

[59] Mclnnes IB, Byers NL, Higgs RE, et al. Comparison of baricitinib, upadacitinib, and tofacitinib mediated regulation of cytokine signaling in human leukocyte subpopulations. Arthritis Res Ther. 2019;21(1):183.

[60] De Benedetti F, Brogan P, Bracaglia C, et al. Op0290 emapalumab (anti-interferon-gamma monoclonal antibody) in patients with macrophage activation syndrome (mas) complicating systemic juvenile idiopathic arthritis (sija). Ann Rheum Dis. 2020;79(Suppl 1):180–180.
with anti-thymocyte globulin. Clin Immunol. 2009; 132(1):10–18.

Horne A, von Bahr Greenwood T, Chiang SC, et al. Efficacy of moderately dosed etoposide in macrophage activation Syndrome-Hemophagocytic lymphohistiocytosis. J Rheumatol. 2021;48(10):1596–1602.

Paravar T, Lee DJ. Thalidomide: Mechanisms of action. Int Rev Immunol. 2008;27(3):111–135.

Boom V, Anton J, Lahdenne P, et al. Evidence-based diagnosis and treatment of macrophage activation syndrome in systemic juvenile idiopathic arthritis. Pediatr Rheumatol Online J. 2015;13(1):55.

Beukelman T, Patkar NM, Saag KG, et al. 2011 American college of rheumatology recommendations for the treatment of juvenile idiopathic arthritis: Initiation and safety monitoring of therapeutic agents for the treatment of arthritis and systemic features. Arthritis Care Res (Hoboken). 2011;63(4):465–482.

Brummaier T, Pohanka E, Studnicka-Benke A, et al. Using cyclophosphamide in inflammatory rheumatic diseases. Eur J Intern Med. 2013;24(7):590–596.

Novick D, Kim SH, Fantuzzi G, et al. Interleukin-18 binding protein: a novel modulator of the Th1 cytokine response. Immunity. 1999;10(1):127–136.

Nirmala N, Brachat A, Feist E, et al. Gene-expression analysis of adult-onset still's disease and systemic juvenile idiopathic arthritis is consistent with a continuum of a single disease entity. Pediatr Rheumatol Online J. 2015;13(1):50.

Giacomelli R, Ruscitti P, Shoensfeld Y. A comprehensive review on adult onset still's disease. J Autoimmun. 2018;93:24–36.

Yasin S, Solomon K, Canna SW, et al. IL-18 as a therapeutic target in a patient with resistant systemic juvenile idiopathic arthritis and recurrent macrophage activation syndrome. Rheumatology (Oxford). 2020;59(2):442–445.

Leurs A, Launay D, Terriou L, et al. Remission of refractory Systemic-Onset juvenile idiopathic arthritis after treatment with siltuximab. J Clin Rheumatol. 2019;25(4):40–42.

Lee M, Isaacs J. The novel use of combined IL-1 and IL-6 inhibition in a patient with severe, aggressive, erosive, systemic-onset juvenile idiopathic arthritis. Eur J Rheumatol. 2017;4(1):68–69.

Verweyen EL, Schulte GS. Interfering with interferons: targeting the JAK-STAT pathway in complications of systemic juvenile idiopathic arthritis (SJIA). Rheumatology. 2022;61(3):926–935.

Brinkman DMC, de Kleer IM, ten Cate R, et al. Autologous stem cell transplantation in children with severe progressive systemic or polyarticular juvenile idiopathic arthritis: long-term follow-up of a prospective clinical trial. Arthritis Rheum. 2007;56(7):2410–2421.

Abinun M, Flood TJ, Cant AJ, et al. Autologous T cell depleted haematopoietic stem cell transplantation in children with severe juvenile idiopathic arthritis in the UK (2000–2007). Mol Immunol. 2009;47(1):46–51.

Wulfraat NM, van Rooijen EM, Tewarie R, et al. Current perspectives of autologous stem cell transplantation for severe juvenile idiopathic arthritis. Autoimmunity. 2008;41(8):632–638.

M. F. Silva J, Ladomenou F, Carpenter B, et al. Allogeneic hematopoietic stem cell transplantation for severe, refractory juvenile idiopathic arthritis. Blood Adv. 2018;2(7):777–786.

Krampera M, Pasini A, Pizzolo G, et al. Regenerative and immunomodulatory potential of mesenchymal stem cells. Curr Opin Pharmacol. 2006;6(4):435–441.

Griffin G, Shenoi S, Hughes GC. Hemophagocytic lymphohistiocytosis: an update on pathogenesis, diagnosis, and therapy. Best Pract Res Clin Rheumatol. 2020;34(4):1015–15.

Tristan AG, et al. Macrophage activation syndrome in a patient with systemic onset rheumatoid arthritis: rescue with intravenous immunoglobulin therapy. J Clin Rheumatol. 2003;9(4):253–258.

Stéphan JL, Koné-Paut I, Galambrou C, et al. Reactive haemophagocytic syndrome in children with inflammatory disorders: A retrospective study of 24 patients. Rheumatology (Oxford). 2001;40(11):1285–1292.

Lin H, Scull BP, Goldberg BR, et al. IFN-γ signature in the plasma proteome distinguishes pediatric hemophagocytic lymphohistiocytosis from sepsis and SIRS. Blood Adv. 2021;5(17):3457–3467.

Tanaka H, Tsugawa K, Nakahata T, et al. Leukocytapheresis for the treatment of refractory systemic-onset juvenile idiopathic arthritis. Clin Rheumatol. 2007;26(6):1014–1016.

Sato S, Hosokawa T, Kawashima H. Successful treatment of plasma exchange for refractory systemic juvenile idiopathic arthritis complicated with macrophage activation syndrome and severe lung disease. Ann Rheum Dis. 2022;81(4):e61–e61. annrheumdis-2020-217390.

Kinjo N, Hamada K, Hirayama C, et al. Role of plasma exchange, leukocytapheresis, and plasma dialfiltration in management of refractory macrophage activation syndrome. J Clin Apher. 2018;33(1):117–120.

Saper VE, Chen G, Deutsch GH. Childhood Arthritis and Rheumatology Research Alliance Registry Investigators, et al. Emergent high fatality lung disease in systemic juvenile arthritis. Ann Rheum Dis. 2019;78(12):1722–1731.

Saper VE, Ombrello MJ, Tremoulet AH, Drug Hypersensitivity Consortium, et al. Severe delayed hypersensitivity reactions to IL-1 and IL-6 inhibitors link to common HLA-DRB1*15 alleles. Ann Rheum Dis. 2022;81(3):406–415.

Schulte GS, Yasin S, Carey B, et al. Systemic juvenile idiopathic arthritis-associated lung disease: Characterization and risk factors. Arthritis Rheumatol. 2019;71(11):1943–1954.

Pettersson T, Konttinen YT. Amyloidosis-recent developments. Semin Arthritis Rheum. 2010;39(5):356–368.

Chantarog S, Vilaiyuk S, Tim-Aroon T, et al. Clinical improvement of renal amyloidosis in a patient with systemic-onset juvenile idiopathic arthritis who
received tocilizumab treatment: a case report and literature review. BMC Nephrol. 2017;18(1):159.

[96] Okuda Y. AA amyloidosis - Benefits and prospects of IL-6 inhibitors. Mod Rheumatol. 2019;29(2):268–274.

[97] Arthur VL, Shuldiner E, Remmers EF, INCHARGE Consortium, et al. IL1RN variation influences both disease susceptibility and response to recombinant human interleukin-1 receptor antagonist therapy in systemic juvenile idiopathic arthritis. Arthritis Rheumatol. 2018;70(8):1319–1330.

[98] Shimizu M, Nakagishi Y, Yachie A. Distinct subsets of patients with systemic juvenile idiopathic arthritis based on their cytokine profiles. Cytokine. 2013;61(2):345–348.