Kidney involvement in multisystem inflammatory syndrome in children: a pediatric nephrologist’s perspective

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ABSTRACT

The initial report of the multisystem inflammatory syndrome in children (MIS-C) was from the UK in April 2020; since then, cases have been reported worldwide. Renal involvement has been seen commonly, ranging from 10% to 46%. Kidney involvement following severe acute respiratory syndrome coronavirus 2 infection in children with MIS-C is more common than initially thought and is associated with higher morbidity and mortality. There are several reports of a direct viral tropism of coronavirus disease 2019 and MIS-C-associated renal damage. This study’s objective was to systematically review the current understanding of kidney involvement in children suffering from MIS-C. Based on our systemic literature search, 19 studies have either partially or fully discussed kidney involvement in MIS-C patients. Furthermore, we discuss the multifactorial pathogenesis contributing to acute kidney injury (AKI) development in MIS-C. The current review gives a pediatric nephrologist’s perspective of the renal involvement in MIS-C, the incidence of AKI, the pathophysiology of AKI in MIS-C and the proposed therapeutic regimens available, including the need for kidney replacement therapy for a child with AKI associated with MIS-C. As the disease is rapidly evolving, more detailed clinical prospective studies are required to understand MIS-C and its role in AKI better.

Keywords: acute kidney injury, COVID-19, MIS-C, nephrology
**INTRODUCTION**

Coronavirus disease 2019 (COVID-19) caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has become a global pandemic [1]. As per the literature [2, 3], with underlying kidney disease, the disease presents with worse outcomes than in those without kidney disease. However, children are mildly affected, and although rare, severe cases have also been documented [4]. There is a lack of strong evidence relating to underlying conditions with severe illness in children [5].

Cases reported from the UK in April 2020 showed a picture similar to incomplete Kawasaki disease (KD) or toxic shock syndrome in children [6]. Since then, reports from different parts of the world describe similarly affected children [7–10]. This clinical disorder has been termed multisystem inflammatory syndrome in children (MIS-C), also known as pediatric inflammatory multisystem syndrome (PIMS) [11, 12], PIMS temporally associated with SARS-CoV-2 (PIMS-TS), pediatric hyperinflammatory syndrome or pediatric hyperinflammatory shock.

Clinicians face a diagnostic dilemma distinguishing KD from MIS-C due to the overlapping clinical features of these two entities and the lack of a definitive diagnostic test for either condition [13]. The signs and symptoms are temporally associated with COVID-19 but presumed to have been developed 2–4 weeks after acute COVID-19, albeit serologic evidence of infection with SARS-CoV-2 [6]. In children, MIS-C occurs as a rare complication of COVID-19 with uncertainty. In a report by Dufort et al., the estimated incidence of laboratory-confirmed SARS-CoV-2 infection in individuals <21 years old was 322/100,000, and the incidence of MIS-C was 2/100,000 [14]. According to the Center of Disease Control and Prevention (CDC), a total of 1659 case of MIS-C have been reported as of 8 January 2021. Acute kidney injury (AKI) is frequently reported in MIS-C as outlined in Table 1, with its incidence ranging from 10% to 60% across studies [15, 16].

**Definition**

The case definitions made by the US CDC in association with the American Academy of Pediatrics, the World Health Organization (WHO), American College of Rheumatology (ACR) and Royal College of Pediatrics and Child Health (RCPCH) are summarized in Tables 2–5. The criteria used for case definition vary slightly between these agencies. The presence of fever remains common along all case definitions except for a much broader duration of fever (>3 days) in WHO guidelines, in contrast to CDC, ACR and RCPCH. The CDC case definition adds several points that are unique to their guideline. Notably, severe illness requiring hospitalization for clinical symptoms and evidence of COVID-19 exposure within 4 weeks are important factors not present in other case definitions. While CDC and WHO case definitions require prior SARS-CoV-2 infection or exposure, RCPCH does not. These definitions are constantly evolving and likely to change as more updated information becomes available (Table 6).

**MATERIALS AND METHODS**

**Search**

A systemic literature search was performed in PubMed/Medline from January 2020 to February 2021 to include all the studies related to MIS-C and AKI. Medical subject headings (MeSH) terms including ‘Multisystem inflammatory syndrome’, ‘pediatric inflammatory multisystem syndrome’, ‘Kawasaki-like disease’, ‘Acute kidney injury’, ‘Acute renal injury’ and ‘renal dysfunction’ were used in the search strategy. We excluded foreign language and case reports during our analysis.

**Data selection and extraction**

Two independent reviewers assessed the literature based on the inclusion and exclusion criteria. A third independent reviewer evaluated articles with conflict. The studies were selected based on selecting any incidence of AKI or the use of renal replacement therapy in patients with MIS-C. A standardized data collection form was used to extract the following information from each included article: the last name of the first author, study type, location, number of MIS-C patients in the study, AKI incidence, use of renal replacement therapy.

**RESULTS**

**Search results**

The initial search returned 91 studies; 18 studies met the selection criteria; 7 were multicenter studies; 8 were single-center studies and 1 was a case series. Details of the included studies are provided in Table 1 and Supplementary Figure S1.

**CLINICAL FEATURES AND PRESENTING SYMPTOMS**

MIS-C presents with persistent fever >38.5°C, variable rash, conjunctivitis, peripheral edema, severe abdominal pain and diarrhea. Fever is the most common presenting symptom, lasting for 3–5 days, though fewer days of fever have been reported [10]. Respiratory distress does not seem to be one of the main presenting symptoms of COVID-19, although a significant proportion of patients require respiratory support [8]. More significantly, many of these children present with hypotension and shock, requiring vasopressor support and pediatric intensive care unit (PICU) admission [6]. Echocardiogram findings include ventricular dysfunction and coronary artery abnormalities [29]. Patients with MIS-C also have evidence of severe inflammatory state, including elevation in C-reactive protein (CRP), ferritin, d-dimer, lactate dehydrogenase, pro-brain natriuretic peptide and cardiac enzymes. Additionally, patients showed elevated pro-calcitonin levels, liver transaminases, anemia, along with thrombocytosis or thrombocytopenia (Table 7) [27]. Constitutional gastrointestinal symptoms like abdominal pain, vomiting and diarrhea are particularly prominent, mimicking appendicitis in some children [26, 30], and may be associated with higher mortality [20].

**PATHOGENESIS**

The pathophysiology of MIS-C is still not well established. It is proposed that MIS-C occurs secondary to an abnormal immune reaction to the virus, much like KD and cytokine release syndrome (CRS), causing this disease [22, 21]. The mechanisms through which SARS-CoV-2 triggers the abnormal immune reaction are unknown. Unabated cytokine storm leading to endothelial dysfunction associated with SARS-CoV-2 infection has also been hypothesized [31, 32]. The mechanisms underlying the exaggerated immune reaction in MIS-C are an area of active investigation (Figure 1).
| Study                  | Study type | Location                                      | Number of MIS-C patients in study | Sex (male) | Age mean (SD) (years) | AKI incidence, n (%) | Use of KRT, n (%) | Additional remarks                                                                 |
|-----------------------|------------|-----------------------------------------------|-----------------------------------|------------|-----------------------|----------------------|-------------------|-------------------------------------------------------------------------------------|
| Gonzalez-Dambrauskas et al. [15] | Multicenter | Chile, Colombia, Italy, Spain, and the USA | 17                                | 11         | 6.52 ± 2.99           | 3 (18)               | Not reported         | Different countries patients were recruited; low mortality 3%                        |
| Stewart et al. [17]    | Single center | London, UK                                    | 52                                | NA         | 4 ± 2.67              | 24 (46)              | Not reported         | Dedicated study for AKI                                                            |
| Derespina et al. [18]  | Multicenter | New York, USA                                 | 70                                | 43         | 14.5 ± 1.67           | 9 (12.9)             | 1 (1.4)            | ARDS associated with poor outcome                                                  |
| Riphagen et al. [6]    | Single center | London, UK                                    | 8                                 | 8          | 9.29 ± 3.77           | 1 (12.5)             | 1 (12.5)            | All children tested negative for SARS-CoV-2 on bronchoalveolar lavage or nasopharyngeal aspirates |
| Whittaker et al. [9]   | Multicenter | UK                                            | 58                                | 38         | 9.43 ± 1.38           | 13 (22)              | Not reported         | Uncontrolled case series                                                          |
| Feldstein et al. [10]  | Multicenter | USA                                           | 186                               | 115        | 8.1 ± 1.53            | 17 (9.1)             | Not reported         | High use of IVIG—79% cases, low mortality—2%                                       |
| Dufort et al. [14]     | Multicenter | New York, USA                                 | 99                                | 53         | 9.33 ± 1              | 10 (10)              | Not reported         | High incidence in Blacks and Hispanics                                             |
| Davies et al. [19]     | Multicenter | UK                                            | 78                                | 52         | 11 ± 1               | –                    | 1 (1)               | –                                                                                  |
| Pereira et al. [20]    | Single center | Sao Paulo, Brazil                              | 6                                 | 5          | 8.30 ± 2.94           | –                    | 50                  | 100% PICU admission, 100% cardiac involvement and 67% death                        |
| Lee et al. [21]        | Single center | Sao Paulo, Brazil                              | 28                                | 16         | 8.77 ± 2.82           | 6 (21)               | Not reported         | Cytopenia, level of hyperferritinemia and pattern of cytokine production MIS-C from KD distinguished |
| Pouletty et al. [22]   | Multicenter | Paris                                          | 16                                | 8          | 9.3 ± 1.3             | 9 (56)               | Not reported         | Older age children were more affected compared with younger children in 'classic' KD (10 years versus 2 years, P < 0.0001), respectively |
| Capone et al. [23]     | Single center | New York, USA                                 | 33                                | 20         | 8.83 ± 1.183         | 23 (70)              | Not reported         | –                                                                                  |
| DeBiasi et al. [24]    | Multicenter | Washington, USA                                | 9                                 | NA         | 15.08 ± 4.25         | 1 (11)               | Not reported         | –                                                                                  |
| Dionne et al. [25]     | Single center | Boston, USA                                    | 25                                | 15         | 9.28 ± 2.05          | 2 (8)                | Not reported         | –                                                                                  |
| Godfried-Cato et al. [16] | Multicenter | District of Columbia, New York, USA          | 570                               | 316        | 9.01 ± 3.33          | 105 (18.4)           | 2 (0.4)             | –                                                                                  |
| Mamishi et al. [26]    | Multicenter | Iran                                          | 45                                | 24         | 6.98 ± 0.98          | 13 (29)              | Not reported         | Wide spectrum of sign and symptoms and abnormal inflammatory markers were noted   |
| Shahbaznejad et al. [27] | Case series | Iran                                          | 10                                | 6          | 5.37 ± 3.9           | 2 (20)               | Not reported         | Short-term outcomes of AKI in PIMS-TS appears good, long-term outcomes are unknown |
| Deep et al. [28]       | Multicenter observational study | UK                                          | 116                               | 76         | 10.5 ± 3.5           | 101 (87)             | 3 (2.97)            | –                                                                                  |

ARDS, acute respiratory distress syndrome; NA, not available.
Children tend to be less susceptible to severe respiratory infections than adults. Milder reactions could be attributable to their robust innate immune response, healthier respiratory tracts and few underlying illnesses. In addition to that, children are overprotected by their parents; many are involved in fewer outdoor activities and undertake less international travel. Notably, decreased gene expression of the angiotensin converting enzyme-2 (ACE-2) receptor (the target of SARS-CoV-2) in the pediatric population could result in the age-related difference in the incidence of COVID-19 [33–36].

However, one of the major challenges is to understand the post-viral lag in symptom presentation in MIS-C patients. MIS-C cases predominantly occur 4 weeks post-COVID-19 peak in the population [37]. One-third of reported MIS-C cases test positive for SARS-CoV-2 using reverse transcriptase–polymerase chain reaction (RT-PCR). Most cases test positive for an antibody test immunoglobulin gamma (IgG), indicating immune response development against the virus. This condition’s lag in symptom presentation may be attributed to a higher ratio of antibody positivity with a lower ratio of SARS-CoV-2 positivity by RT-PCR, which suggests a late inflammatory process owing to antibody-and/or immune complex-mediated development to SARS-CoV-2 [38]. Although the pathophysiology of MIS-C remains unclear, the conversion of mild disease to a multisystem inflammatory response in the post-infectious period is thought to emerge from a dysregulated immune response resembling CRS [10, 13].
The timing of the interferon (IFN) response to SARS-CoV-2 infection plays a critical role in the presentation of MIS-C. A hypothesis by Rowley suggests that early IFN response may lead to rapid viral clearance during the early stages of the disease, inducing a mild illness, when the viral burden is low. Conversely, with a high viral burden, the replicating virus may delay the IFN response, resulting in a marked cytokine storm before the body’s adaptive response can clear the virus, driving a severe infection including MIS-C [13].

Furthermore, Martinez et al. theorize that some children with MIS-C could harbor viral RNA in the nasopharynx, which may provide a continuous antigen source for T cells leading to chronic viral exposure in the respiratory system. These theories, however, should be tested in future longitudinal studies [39].

Antibodies against SARS-CoV-2 can play a functional role in MIS-C. Gruber et al. suggest the role of neutralizing antibodies against SARS-CoV-2 in patients with MIS-C and its association with interleukin-18 (IL-18) and IL-16 activation, myeloid chemotaxis and activation of lymphocytes, monocytes and natural killer cells [37].

Table 3. Case definition of MIS-C as per WHO

| WHO case definition |
|---------------------|
| All six criteria must be met: |
| (1) Age 0–19 years |
| (2) Fever for ≥3 days |
| (3) Clinical signs of multisystem involvement (at least two of the following): |
| • Rash, bilateral non-purulent conjunctivitis, or mucocutaneous inflammation signs (oral, hands or feet) |
| • Hypotension or shock |
| • Cardiac dysfunction, pericarditis, valvulitis or coronary abnormalities (including echocardiographic findings or elevated troponin/BNP) |
| • Evidence of coagulopathy (prolonged prothrombin time or partial thrombin time; elevated D-dimer) |
| • Acute gastrointestinal symptoms (diarrhea, vomiting or abdominal pain) |
| (4) Elevated markers of inflammation (e.g. ESR, CRP or procalcitonin) |
| (5) No other obvious microbial cause of inflammation, including bacterial sepsis and staphylococcal/streptococcal TSSs |
| (6) Evidence of SARS-CoV-2 infection any of the following: |
| • Positive SARS-CoV-2 RT-PCR |
| • Positive serology |
| • Positive antigen test |
| • Contact with an individual with COVID-19 |

ESR, erythrocyte sedimentation rate; BNP, brain natriuretic peptide. MIS-C and adolescents with COVID-19: Scientific Brief. 2020. https://www.who.int/publications-detail/multisystem-inflammatory-syndrome-in-children-and-adolescents-with-covid-19 (17 May 2020, date last accessed).

Table 4. Case definition of MIS-C as per ACR

| ACR |
|-----|
| (1) All children (age not defined) |
| (2) Unremitting fever ≥38°C |
| (3) At least two suggestive clinical features: |
| • Rash |
| • GI symptoms |
| • Edema of hands/feet |
| • Oral mucosal changes |
| • Conjunctivitis |
| • Lymphadenopathy |
| • Neurologic symptoms |
| (4) Laboratory evidence: 1) CRP ≥5 mg/dL OR 2) ESR ≥40 mm/h |
| 3) At least one suggestive laboratory feature: |
| • ALC <1000/µL |
| • Platelet count <150 000/µL |
| • Na <135 mmol/L |
| • Neutrophilia |
| • Hypoalbuminemia |
| (5) SARS-CoV-2 PCR and/or serologies |
| (6) Exclusion of other infections |

ALC, absolute lymphocyte count; BNP, brain natriuretic peptide; ESR, erythrocyte sedimentation rate; GI, gastrointestinal; LDH, lactate dehydrogenase; Na, sodium.

The timing of the interferon (IFN) response to SARS-CoV-2 infection plays a critical role in the presentation of MIS-C. A hypothesis by Rowley suggests that early IFN response may lead to rapid viral clearance during the early stages of the disease,
The pathogenesis of AKI in MIS-C is a complex, multifaceted entity with overlapping mechanisms. The means through which SARS-CoV-2 triggers the abnormal immune reaction are unknown. Investigators have reported several mechanisms for SARS-CoV-2 triggers the abnormal immune reaction are unknown. Investigators have reported several mechanisms for AKI in COVID-19. They are thought to occur secondary to an abnormal immune response to the virus leading to tubular injury and podocytopathy, inflammatory process, hemodynamic instability and vascular endothelial dysfunction [21]. The underlying exaggerated immune reaction in MIS-C is a place of active investigation.

COVID-19 exhibits viral tropism and can directly affect the kidney. The spike (S) glycoprotein of SARS-CoV-2 binds to the ACE-2 receptor on host cells. Afterward, the active S protein is cleaved by transmembrane serine proteases (TMPRSSs), resulting in membrane fusion facilitated by fusion peptides released by the virus. Finally, the virus penetrates the proximal tubule cells and the podocytes leading to renal impairment (podocytopathy and tubular epithelial cell injury) [43, 44]. This is suggested by a co-expression of ACE2 and TMPRSS2 genes in the podocytes and proximal convoluted tubules, like that in the lung, small intestine and colon [44].

Sub-Saharan African descent with high-risk apolipoprotein-1 (APOL1) genotype (presence of two risk alleles) could be at increased risk of kidney disease in the setting of COVID-19 [45]. Case reports of patients presenting with heavy proteinuria associated with collapsing glomerulopathy with a predisposition in the form of APOL1 polymorphism suggest a crucial role of innate immunity pathways, which are upregulated in viral illness [45].

The development of a hyperinflammatory state as a response to the virus in MIS-C suggests a macrophage activating syndrome (MAS) like cytokine storm or secondary hemophagocytic lymphohistiocytosis (HLH). MAS or secondary HLH is a life-threatening condition triggered by an autoimmune condition or viral illness [46]. The findings indicative of MAS include cytopenias, coagulopathy, tissue damage and hyperferritinemia, leading to a fatal multi-organ failure [21, 47, 48]. Similar findings have been noticed in MIS-C patients, specifically hyperferritinemia on admission, associated with severe AKI contributing to the acute inflammatory state in MIS-C [28]. Furthermore, data from an adult study on secondary HLH demonstrate increased AKI incidence adversely affecting patient survival [49].
Another cardinal finding of MAS bearing similarity with MIS-C is the involvement of the plasma mediators driving the hypercytokinemia state. Cytokine profiling studies show that IFN-$\gamma$-induced response markers (including IFN-$\gamma$, IL-18, IP-10) are the main triggers of inflammation in both MIS-C and MAS [50]. Other important cytokines involved are IL-2, IL-6, IL-7, IL-16, granulocyte-colony stimulating factor, macrophage inflammatory protein-1a, tumor necrosis factor-alpha (TNF-$\alpha$), MCP-1, IL-1$\alpha$ and IL-1 receptor agonist (IL-1Ra) [51].

The immunological mechanisms of MAS/secondary HLH overlap with the complex hyperinflammatory pathology of MIS-C in AKI. Extensive clinical and translational studies are required to better understand the cytokine mediators role and their efficacious use and safety as a target for future therapeutics.

IL-6 plays a crucial role in inducing the systemic inflammatory response in COVID-19, and elevated levels have been associated with worsening renal outcomes [52, 53]. A positive correlation was shown between severity of COVID-19 disease and serum levels of IL-6 and IL-2R [54]. Interestingly, experimental studies demonstrate a role of IL-6 activation and secretion by podocytes, endothelial cells, mesangial cells and tubular epithelial cells in renal inflammatory diseases. IL-6 levels would therefore aid in early diagnosis, serve as a target for therapeutics and prevent AKI worsening [55, 56].

Similarly, a proinflammatory cytokine, IL-8, plays a role in recruiting neutrophils and lymphocytes in the tubular interstitium causing inflammation. Investigational studies demonstrate upregulation of IL-8 in tubular epithelial cells due to elevated albumin and proteinuria leading to AKI [57]. A study on adults reported that 34% of COVID-19-infected individuals acquired proteinuria on the first day of admission, with 63% presenting with proteinuria during their hospital stay [58]. Proteinuria of any degree is associated with poor prognosis and increased in-hospital mortality [59]. In conclusion, it is plausible to hypothesize that these cumulative mechanisms could propel renal function impairment in a hypercytokinemia state of MIS-C.

The pathogenesis of post-viral inflammatory response of MIS-C raises the concern for cytokine-mediated hypotension leading to renal hypoperfusion associated with AKI [9]. Deep et al. [28] conducted a multicenter study on 116 PIMS-TS patients, demonstrating 41.4% of children with AKI (any stage) and 27.6% with severe AKI. Nearly half (49%) of patients presented with vasodilatory shock requiring vasopressor support. In the same way, Stewart et al. observed the interaction between hypervolemia and hyperinflammatory shock in majority of their patients with AKI. The renal function improved following fluid resuscitation and use of inotropic support. However, the study was focused on pediatric patients with COVID-19 who suffered renal dysfunction [17]. Other factors, including vomiting and diarrhea, also precipitated dehydration leading to inadequate renal perfusion. These could be contributing to decreasing effective renal perfusion, thereby supporting renal hypoperfusion as a factor causing AKI [60, 61] (Figure 2).

Other etiologies that may contribute to AKI development in MIS-C patients include:

(i) imbalanced renin–angiotensin–aldosterone system activation promoting glomerular dysfunction, fibrosis and vasoconstriction [62];
(ii) endothelial dysfunction, complement and coagulation activation, leading to kidney vascular injuries such as ischemic glomeruli and fibrinoid necrosis [45];
(iii) obstruction of the glomerular capillaries by red blood cells similar to the case in thrombotic microangiopathy [63]; and
(iv) drug toxicity and organ cross-talks are non-specific factors relative to critically ill COVID-19 patients management that may aggravate kidney injury [63–65].

Future studies are needed to further highlight the role of COVID-19 in MIS-C and AKI pathogenesis of hypervolemia and hyperinflammatory state. Additionally, constant renal function surveillance in MIS-C patients can be considered early in the disease course, with the help of collaborative multidisciplinary team, while avoiding AKI exacerbating factors and complications.

TREATMENT

There is no conclusive set of guidelines to manage MIS-C. Still, several organizations have published data based on the locally formulated protocol from previous treatment guidelines of similar diseases, including KD or adult protocols of COVID-19 treatment. Dove et al. conducted a multicenter cross-sectional survey on protocols of early hospital evaluation for treating MIS-C patients. The authors discussed various individual institutional protocols. Of the 40 centers, 21 centers required only 1 day of fever to consider MIS-C as a primary diagnosis. The majority of the centers used intravenous immunoglobulin (IVIG) and corticosteroids as primary therapy (98% and 93%, respectively). Additionally, heparin, anakinra and vasopressor agents were used frequently in children with severe illness [66]. The primary goal of the therapy in MIS-C is to reduce systemic inflammation with organ restoration decreasing mortality and reducing the risk of long-term sequelae. There is a need for multispecialty care guiding management and rehabilitation in patients with MIS-C due to its varied nature and presentation.

Categorizing patients at risk to an appropriate level of care

The suitable care setting is determined by the severity of the disease, risk of complications and follow-up ability. Children with mild symptoms, normal vital signs and physical examination can be managed in the outpatient setting with appropriate follow-up. Children with moderate to severe symptomatic MIS-C (Table 7) and those at risk for complications should be admitted to the hospital. Admission to a PICU is considered appropriate for children with significant respiratory compromise, hemodynamic instability and other potentially life-threatening complications. Published studies show the requirement of ICU to be as high as 80% [9, 10, 14, 18, 19].

Supportive care

General supportive care is pivotal in management [9]. The focus should be placed on the patient’s vital signs, hydration status, electrolyte and metabolic status. Special care in early detection of signs of hypoxia with prompt intervention should be taken for better outcomes. Children presenting with shock should be resuscitated according to standard protocols, with early goal-directed volume expansion followed by vasoactive agents [23]. In the event of LV dysfunction, epinephrine is preferable, with the addition of milrinone in case of severe LV dysfunction [23].

MIS-C-specific care

Children meeting the criteria for KD should receive standard therapies, including IVIG, aspirin and, if facing persistent signs of inflammation or coronary artery dilation/aneurysm, glucocorticoids. In instances of fulminant disease, mechanical hemodynamic support could also be necessary within the sort of extracorporeal membrane oxygenation (ECMO) or a ventricular assist device [47].

IVIG 2 g/kg and aspirin 20–25 mg/kg/dose every 6 h (80–100 mg/kg/day) should be used for all patients with KD-like illness, evidence of excessive inflammation (ferritin >700 ng/mL, CRP >30 g/dL or multisystem organ failure) or cardiac involvement. Aspirin dosage may also vary at individual centers. IVIG 2 g/kg as a single infusion with 3-day pulse methylprednisolone.
should be reserved for patients in high-risk categories (infants, KD shock syndrome, CRP >130 g/dL, admission echo Z-score >2.5 or aneurysms, Asian race) [47]. Patients with persistent inflammation despite IVIG and steroids can be given anakinra, an IL-1R antagonist, tocilizumab, an IL-6 antagonist, and infliximab, TNF-α blocker. Until now, the data on IL-1 and IL-6 blockade in COVID-19 infection and associated MIS-C are limited, but encouraging with multiple clinical trials currently in progress showing a high short-term survival rate in patients receiving IVIG and steroids [55, 56, 67] (Table 8).

**Treatment of MIS-C with kidney involvement**

Kidney replacement therapy (KRT) in the setting of multiorgan disease syndrome should be instituted with expertise. KRT can be utilized non-selectively to clear inflammatory mediators via convection, adsorption and diffusion. Continuous kidney replacement therapy (CKRT) corrects fluid overload and manages solute levels to provide hemodynamic stability in catabolic pediatric patients [68]. Immediate initiation of preemptive KRT in cases with progressive symptomatic respiratory insufficiency improves overall outcomes.

The pediatric continuous renal replacement therapy group advises the early initiation of KRT in critically ill COVID-19-related immune dysregulation syndrome patients since it has been shown to mitigate fluid overload, enhance cytokine clearance, improve PaO2/FiO2 ratios and establish hemodynamic stability earlier, leading to better overall outcomes [69]. No additional benefit has been reported with hybrid therapies such as prolonged intermittent renal replacement therapy (PIRRT) and sustained low efficiency dialysis. Hence, it is suggested to continue using the local institution’s recommended treatment

**Table 8. Drugs used in the treatment of MISC (relevant to nephrology care)**

| Drug               | Mechanism          | Dose in clinical trial | Dose adjustment | Special consideration                        |
|--------------------|--------------------|------------------------|-----------------|---------------------------------------------|
| Remdesivir         | Nucleotide analog: Inhibits viral replication | 3.5 to <40 kg: 5 mg/kg IV loading dose on Day 1, followed by 2.5 mg/kg IV every 24 h for 5–10 days (5 days for those with a rapid clinical response) <40 kg: 200 mg IV loading dose on day 1, followed by 100 mg IV every 24 h for 5–10 days (5 days for those with a rapid clinical response) | Not recommended GFR <30 mL/min/1.73 m² The excipient, a cyclodextrine accumulates in GFR <30 mL/min/1.73 m² | FDA approved |
| Hydroxychloroquine | Immuno-modulatory effects | 400 mg on day 1, followed by 200 mg twice a day | None | To be avoided in children with underlying QTc abnormalities |
| Lopinavir-ritonavir | Protease inhibitor | 100 mg twice a day | None | Not recommended in children |
| IVIG               | Immunomodulatory effects | 2 g/kg | NA | If IVIG is not available or contraindicated, consider using corticosteroids |
| Glucocorticoids    | Anti-inflammatory  | Low-dose glucocorticoid regimens Dexamethasone 0.15 mg/kg orally, IV or via an NG tube once daily (maximum dose 6 mg); prednisolone 1 mg/kg orally or via an NG tube once daily (maximum dose 40 mg); methylprednisolone 0.8 mg/kg IV once daily (maximum dose 32 mg) | None | Pediatric glucocorticoid arm of the RECOVERY trial is ongoing |
| Tocilizumab        | IL-6 antagonist    | 8 mg/kg maximum 800 mg | None | In clinical trial only |
| Anakinra           | IL-1Ra             | 2–6 mg/kg/day IV/SQ, length of therapy to be decided <30 mL/min GFR—administer every alternate day | Not dialyzable |
| Aspirin            | Anti-inflammatory  | 30–50 mg/kg/day, decrease to 3–5 mg/kg/day once afebrile 48 h GFR <10 mL/min/1.73 m²—caution | Dialyzable—50–100% |

FDA, Food and Drug Administration; GFR, glomerular filtration rate; IV, intravenous; NG, nasogastric tube; QTc, corrected QT interval; SQ, subcutaneous.

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2008 | S.K. Sethi et al.
modality. If CKRT/PIRRT is not available, intermittent hemodialysis is acceptable [69].

In terms of CKRT modality, the use of high flow continuous venovenous hemodiafiltration in critically ill COVID-19 pediatric patients is recommended because of its ability to boost the nonspecific removal of the circulatory cytokine mediators [70]. Increased reports of thrombotic complications are coming up in patients with MIS-C. Care should be taken during dialysis to avoid circuit clotting. As per local protocols, after insertion with heparin/citrare, the catheter’s immediate locking may be considered [71].

Some critically ill patients require mechanical ventilation via ECMO. Integration of the ECMO circuit with the CKRT equipment for these patients can provide additional respiratory support while reducing/preventing fluid overload and cytokine clearance [69]. However, the use of ECMO comes with a risk of potentially accentuating the cytokine activation.

CONCLUSION

Although there has been an increasing number of published data, the overall population-specific incidence of MIS-C remains unknown. The causal relationship and pathogenesis of kidney disease and MIS-C are still under debate. Post-viral immunological reaction to COVID-19 remains the best-implicated theory behind this disease’s pathogenesis, although an overall understanding of the immunopathogenesis induced by SARS-CoV-2 remains elusive. A better understanding of the kidney physiology in COVID-19 and treatment modalities available for children with renal dysfunction and MIS-C will help us to provide better care for this subset of children.

SUPPLEMENTARY DATA

Supplementary data are available at ckJ online.

CONFLICT OF INTEREST STATEMENT

None declared.

DATA AVAILABILITY STATEMENT

The data used in this article will be shared on reasonable request to the corresponding author.

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