Systemic Inflammation Is Associated With Neurologic Involvement in Pediatric Inflammatory Multisystem Syndrome Associated With SARS-CoV-2

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Abstract

Objective
Pediatric inflammatory multisystem syndrome temporally associated with SARS-CoV-2 (PIMS-TS) is a severe immune-mediated disorder. We aim to report the neurologic features of children with PIMS-TS.

Methods
We identified children presenting to a large children’s hospital with PIMS-TS from March to June 2020 and performed a retrospective medical note review, identifying clinical and investigative features alongside short-term outcome of children presenting with neurologic symptoms.

Results
Seventy-five patients with PIMS-TS were identified, 9 (12%) had neurologic involvement: altered conciseness (3), behavioral changes (3), focal neurology deficits (2), persistent headaches (2), hallucinations (2), excessive sleepiness (1), and new-onset focal seizures (1). Four patients had cranial images abnormalities. At 3-month follow-up, 1 child had died, 1 had hemiparesis, 3 had behavioral changes, and 4 completely recovered. Systemic inflammatory and prothrombotic markers were higher in patients with neurologic involvement (mean highest CRP 267 vs 202 mg/L, p = 0.05; procalcitonin 30.65 vs 13.11 μg/L, p = 0.04; fibrinogen 7.04 vs 6.17 g/L, p = 0.07; D-dimers 19.68 vs 7.35 mg/L, p = 0.005). Among patients with neurologic involvement, these markers were higher in those without full recovery at 3 months (ferritin 2284 vs 283 μg/L, p = 0.05; D-dimers 30.34 vs 6.37 mg/L, p = 0.04). Patients with and without neurologic involvement shared similar risk factors for PIMS-TS (Black, Asian and Minority Ethnic ethnicity 78% vs 70%, obese/overweight 56% vs 42%).

Conclusions
Broad neurologic features were found in 12% patients with PIMS-TS. By 3-month follow-up, half of these surviving children had recovered fully without neurologic impairment. Significantly higher systemic inflammatory markers were identified in children with neurologic involvement and in those who had not recovered fully.

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SARS-CoV-2 is a novel and highly pathogenic respiratory coronavirus causing the global COVID-19 pandemic. Neurologic syndromes have been reported within the acute phase of disease, and these may arise from presumed direct viral invasion or result from a para or postinfectious inflammation. Although children present with a milder course of disease and case fatality rates are lower, a severe multisystem hyperinflammatory response has been reported in children during the pandemic, clinically defined as pediatric inflammatory multisystem syndrome temporally associated with SARS-CoV-2 (PIMS-TS). Here, we characterize the spectrum of neurologic involvement was also noted in PIMS-TS. Here, we characterize the spectrum of neurologic features among a cohort of 75 children with PIMS-TS and aim to identify differences to those presenting without neurologic symptoms.

Methods

A retrospective cross-sectional review of patients with PIMS-TS admitted to Evelina London Children’s Hospital was performed to identify the clinical and investigative features alongside short-term outcome of children presenting with neurologic symptoms. Patients younger than 18 years and admitted between March and June 2020 who met the following criteria were included in our cohort:

1. Prospective diagnosis of PIMS-TS according to the Royal College of Paediatrics and Child Health case definition (persistent fever, inflammation, and evidence of single or multiorgan dysfunction and exclusion of other microbial causes) following review by a multidisciplinary team comprising of at least 1 each of infectious diseases, intensive care, cardiology, rheumatology, and general pediatric specialists.

2. New onset of at least one of the following: features of encephalopathy (altered consciousness that persisted for longer than 24 hours, including lethargy, irritability, or a change in personality and behavior), seizures, movement disorder, psychiatric, and/or any neurologic deficit.

All inflammatory and prothrombotic investigations (table e-1, links.lww.com/NXI/A474) were taken before therapy (including aspirin); the highest value was used for analysis. The modified Rankin Scale (mRS) at last clinical review was used to document clinical outcome. Demographic features and inflammatory markers were compared with the cohort of patients with PIMS-TS admitted during the same period with no neurologic symptoms. Statistical analysis was performed using SPSS 24.0 IBM (Chicago, Illinois, US).

Table 1 Demographic and Clinical Information of the 9 Patients Presenting With Neurologic Features

| Patients | Neurologic features (syndrome) | Neurologic investigations | Systemic features | Inflammatory markers | SARS-CoV-2 | Treatment and outcome |
|----------|-------------------------------|--------------------------|-------------------|---------------------|-----------|----------------------|
| 1, female, 2 y, Afro-Caribbean, obesity | Altered consciousness (encephalopathy secondary to systemic inflammation) | Brain CT and MRI; EEG: normal (palecytosis, oligodendral band, and NSABs) | Fever Lymphadenopathy, periorbital and lip edema Abdominal pain Mild left ventricular impairment and coronary aneurysm | Ferritin 275 μg/L; CRP: 189 mg/L; procalcitonin: 4.72 μg/L; fibrinogen: 8.6 g/L; d-dimers: 12.86 mg/L | PCR NP and serology positive, PCR CSF negative | IVIG; NMP (3 d); tocilizumab (1 dose) Complete recovery at 3 mo (mRS 0) |
| 2, male, 4 y, Caucasian | Altered consciousness; behavioral changes (encephalopathy secondary to systemic inflammation) | Brain MRI: normal EEG: normal CSF: mixed oligodendral bands (serum and CSF) | Fever Intense abdominal pain, diarrhea, and vomits Bright coronaries and dilated left anterior descending coronary artery | Ferritin 172 μg/L; CRP: 283 mg/L; procalcitonin: 6.8 g/L; d-dimers: 10.6 mg/L | PCR NP and serology negative | IVIG; VMP (3 d) Behavioral changes at 3 mo follow-up (mRS 3) |
| 3, female, 6 y, Asian | Severe behavioral changes (encephalopathy secondary to systemic inflammation) | Not performed | Fever Rash, face and feet edema Abdominal pain and vomits Coronary aneurysm | Ferritin 74 μg/L; CRP: 80 mg/L; fibrinogen: 4.1 g/L; d-dimers: 1.2 mg/L | PCR NP and serology positive | Oral prednisolone Complete recovery at 4 mo (mRS 0) |

Continued
IL). A p value less than 0.05 was considered statistically significant, and care was given to limiting multiple comparison.

**Standard Protocol Approvals, Registrations, and Patient Consents**

As only data required for standard medical care were collected, a full ethics review under the terms of the Governance Arrangements of Research Ethics Committees in the United Kingdom was deemed not necessary by the institutional PIMS-TS Study Group.

**Data Availability**

Deidentified participant data not included in the article are available on request by any qualified investigator to the corresponding author.

### Table 1 Demographic and Clinical Information of the 9 Patients Presenting With Neurologic Features (continued)

| Patients | Neurologic features (syndrome) | Neurologic investigations | Systemic features | Inflammatory markers | SARS-CoV-2 | Treatment and outcome |
|----------|--------------------------------|--------------------------|-------------------|----------------------|------------|----------------------|
| 6, female, 12 y, Afro-Caribbean, obesity | Behavioral changes; cognitive deterioration (toxo/metabolic encephalopathy) | Brain MRI: subtle cortical changes | Fever | Ferritin: 4,815 μg/L; CRP: 343 mg/L; procalcitonin: >100 μg/L; fibrinogen: 6.8 g/L; d-dimers: 69.76 mg/L | PCR NP negative, serology positive | IVIG (2 doses); IVMP (5 d); infliximab (1 dose) | Mild behavioral changes/low mood at 4 mo follow-up (mRS 1) |
| 7, male, 14 y, Afro-Caribbean, obesity, α-thalassemia trait | Headaches; focal neurology with asymmetric pupils (stroke) | Brain CT: acute infarction (figure 1E) | Fever | Ferritin: 4,220 μg/L; CRP: 556 mg/L; procalcitonin: >100 μg/L; fibrinogen: 10.8 g/L; d-dimers: 13.49 mg/L | PCR NP and serology positive | No immune treatment | Deceased after 8 d of admission |
| 8, female, 15 y, Afro-Caribbean, overweight, Hbc trait | Behavioral changes; visual and auditory hallucinations; new onset of seizures (toxo/metabolic encephalopathy) | Brain MRI: hippocampal and splenium of corpus callosum changes (figure 1, A–D); EEG: focal spike-wave discharges | Fever | Ferritin: 640 μg/L; CRP: 99 mg/L; procalcitonin: 0.22 μg/L; fibrinogen: 6.8 g/L; d-dimers: 28.79 mg/L | PCR NP, PCR CSF, and serology negative | IVIG; IVMP (3 d). | Memory difficulties at 3-mo follow-up. Seizures resolved (mRS 2) |
| 9, male, 10 y, Afro-Caribbean, sickle cell | Encephalopathy; left sided facial weakness. Hypertension and bradycardia (raised ICP) (stroke) | Brain CT/MRI: right frontal intraparenchymal haemorrhage (figure 1F) and infarction | Fever | Ferritin: 1,572 μg/L; CRP: 296 mg/L; procalcitonin: 0.34 μg/L; fibrinogen: 7.3 g/L; d-dimers: 29.08 mg/L | PCR NP negative, serology positive | IVIG; IVMP (3 d); tocilizumab (1 dose) | Left hemiparesis (mRS 4) |

Abbreviations: ARDS = acute respiratory distress syndrome; CRP = C-reactive protein; ICP = intracranial pressure; IVIG = IV immunoglobulin; IVMP = IV methylprednisolone; MRA = magnetic resonance angiogram; mRS = modified Rankin Scale; ND = not done; NSAB = neuronal surface antibody; NP = nasopharyngeal.
Results

Clinical and Investigative Features
A total of 75 children (33% female; median 10 years, interquartile range 7.9 years) were diagnosed with PIMS-TS. At the time of testing availability (June–July 2020), 51 (68%) had detectable immunoglobulin (Ig) G to SARS-CoV-2. Of the 75 children, 9 (12%) had neurologic features (table 1; table e-2, links.lww.com/NXI/A475), altered consciousness (3/9; 33%), acute behavioral changes (3/9; 33%), focal neurologic deficits (2/9, 22%), severe persistent headaches (2/9; 22%), visual/auditory hallucinations (2/9, 22%), excessive sleepiness (1/9, 11%), and new-onset of focal seizures (1/9, 11%). All patients had fever at symptom onset and the range of systemic features include had gastrointestinal symptom (8/9; 89%), cardiovascular instability (5/9; 56%), and respiratory symptoms (4/9; 44%). Four patients (44%) had left ventricular dysfunction and 5 (56%) coronary artery abnormalities.

Four patients had abnormal imaging: focal diffusion restriction involving the splenium of the corpus callosum and mild signal changes in hippocampal regions (patient 8, figure 1, A–D); acute infarction (patient 7, figure 1E); intraparenchymal hemorrhage and infarction (patient 9, figure 1F); and subtle cortical changes as a possible sequelae of hypoxic event (patient 6). EEG showed abnormalities in the 3 of the 4 patients tested. These were reduced amplitude of background rhythms in 2 (patients 1 and 6; features of encephalopathy) and sharpened slow waves in 1 (patient 8; seizure liability). Three children had CSF analysis, and this was normal in 2 and neither had SARS-CoV-2 identified in CSF. One child had matched oligoclonal bands in serum and CSF (patient 2). Mean highest measured inflammatory markers were raised (table 2). Evidence of active infection (positive nasopharyngeal swab PCR) was found in 3 (33%) patients, and a further 3 (33%) had evidence of recent infection (negative PCR and positive IgG serology).

Early Neurologic and Behavioral Outcome
One patient died after extensive brain infarction (patient 7, mRS 6), and another was transferred to a neurosurgical center for management of intraparenchymal hemorrhage (patient 9, mRS 4). The other 7 patients were discharged home after a median period of 11 (range 6–25) days of admission. Four (57%) patients recovered completely (mRS 0) at 3-month follow-up review. In the other 3 children, the following outcomes were seen: reported persistent significant behavioral and social interaction difficulties (mRS 3), mild behavioral changes with low mood (mRS 1), and memory difficulties (mRS 2).

Comparison With Children Who Did Not Exhibit Neurologic Symptoms
Comparison was made between those with PIMS-TS and associated neurologic involvement (n = 9) and those without neurologic involvement (n = 66) (table 2). Risk factors for PIMS-TS were similar: Black, Asian and Minority Ethnic ethnicity 78% vs 70%, obesity (body mass index >95th centile)/overweight (86th–95th centile) 56% vs 42%. Both groups had comparable number and duration of intensive care unit (ICU) admission and cardiac involvement. Notably, systemic inflammatory markers were higher in patients with neurologic involvement (figure 2). Considering patients with neurologic involvement (n = 9), peak inflammatory and
prothrombotic markers during acute disease were higher in children with sequelae at 3-month follow-up (n = 5) in comparison with children with complete neurologic recovery (n = 4) (table 2, figure 2).

**Table 2** Comparison Between Those With PIMS-TS and Associated Involvement and Those Without Neurologic Involvement

|                          | PIMS-TS and neurology symptoms (n = 9) | No neurology symptoms (n = 66) | p Value |
|--------------------------|----------------------------------------|--------------------------------|---------|
| Female, n (%)            | 5 (56)                                 | 20 (30)                        | 0.15    |
| Median age at symptom onset, y | 10 (IQR 8.4)                          | 10 (IQR 8.1)                   | 0.43    |
| Ethnicity: BAME, n (%)   | 7 (78)                                 | 43 (65)                        | 0.45    |
| Obese/overweight, n (%)a| 5 (56)                                 | 28 (42)                        | 0.49    |
| GI symptoms, n (%)       | 8 (89)                                 | 53 (80)                        | 0.53    |
| Respiratory symptoms, n (%)| 4 (44)                                 | 31 (47)                        | 0.89    |
| Cardiac involvement, n (%)| 7 (78)                                 | 46 (70)                        | 0.62    |
| Evidence of SARS-CoV-2 infection, n (%)b | 6 (67)                                 | 45 (68)                        | 0.93    |
| Median admission duration, d | 11                                     | 8                              | 0.08    |
| ICU admission, n (%)     | 6 (67)                                 | 42 (64)                        | 0.86    |
| Median ICU admission, d  | 6.5                                    | 3                              | 0.69    |
| Immunomodulatory treatment, n (%) | First line (IVIg/IVMP) | 7 (78) | 63 (95) | 0.11 |
|                          | Second line (anakinra/infliximab/    | 2 (22) | 25 (38) | 0.48 |
|                          | tocilizumab)                           |                                |         |
| Inflammatory markersc    |                                        |                                |         |
| CRP, mg/L                | 267 (median: 283; IQR: 244)            | 202 mg/L (median: 188; IQR: 141) | 0.05    |
| Ferritin, µg/L           | 1,395 (median: 640; IQR: 1,400)        | 980 (median: 549; IQR: 871)    | 0.21    |
| Procalcitonin, µg/L      | 30.65 (median: 4.72; IQR: 61.51)       | 13.11 (median: 3.10; IQR: 8.03) | 0.04    |
| Fibrinogen, g/L          | 7.04 (median: 6.8; IQR: 0.5)           | 6.17 (median: 5.9; IQR: 1.7)   | 0.07    |
| ν-dimers, mg/L           | 19.68 (median: 12.86; IQR: 18.43)      | 7.35 (median: 3.36; IQR: 6.87) | 0.005   |
| Residual symptoms (n = 5)| Complete recovery (n = 4)              |                                |         |
| CRP, mg/L                | 315 (median: 296; IQR: 60)             | 207 (median: 142; IQR: 170)    | 0.19    |
| Ferritin, µg/L           | 2,284 (median: 1,572; IQR: 3,580)      | 283 (median: 206; IQR: 245)    | 0.05    |
| Procalcitonin, µg/L      | 36.63 (median: 4.98; IQR: 77.26)       | 20.69 (median: 4.72; IQR: 27.48) | 0.32    |
| Fibrinogen, g/L          | 7.70 (median: 6.8; IQR: 0.5)           | 6.23 (median: 6.1; IQR: 2.18)  | 0.13    |
| ν-dimers, mg/L           | 30.34 (median: 28.79; IQR: 15.59)      | 6.37 (median: 5.78; IQR: 9.82) | 0.04    |

Abbreviations: BAME = Black, Asian and Minority Ethnic; CRP = C-reactive protein; ICU = intensive care unit; IQR = interquartile range; IVIG = IV immunoglobulin; IVMP = IV methylprednisolone.

Peak inflammatory marker comparison between those with residual neurologic symptoms and those with complete neurologic recovery at 3-month follow-up.

a Obesity defined as body mass index >95th centile and overweight as 86th-95th centile for age and sex.
b Positive nasopharyngeal swab PCR or positive serology.
c Mean of highest measured inflammatory marker.

Discussion

In this study, we describe the spectrum of neurologic manifestations in 9 children with PIMS-TS. The frequency of
neurologic symptoms was relatively low (12%), and this was comparable to a previous smaller cohort.5 Two children developed extensive stroke. Stroke in adult patients has been extensively reported and is thought to be due to inflammation driven hypercoagulability with thrombotic microangiopathy playing a major role in COVID-19 syndromes.8 A characteristic radiologic feature noted in our cohort is the splenial lesion. Among other neuroradiologic features, the same splenial findings had been previously reported and also with a favorable clinical outcome, during the acute SARS-CoV-2 infection5 and delayed hyperinflammatory syndrome.9 Of interest, SARS-CoV-2 IgG was undetected in 24 (32%) of children. One plausible explanation for this is that antibody concentrations may have decayed to below the threshold of detection by the time testing was clinically available in our patients.10 Alternatively, antibody specificity and concentrations are known to differ in children with PIMS-TS in comparison to adults with COVID-19,11 explaining an incomplete seroprevalence in this PIMS-TS cohort. Seronegative patients with PIMS-TS were phenotypically similar to seropositive patients with PIMS-TS, presented in the period of the first surge of cases in the UK COVID-19 pandemic, and were negative by PCR and relevant serology to extensive virologic testing.

The key finding of our study is that children with neurologic symptoms as part of their PIMS-TS presentation have significantly higher systemic inflammatory markers than children without neurologic features. These findings support the hypothesis that PIMS-TS–associated neurologic involvement comprises a systemic para or postinfectious immunemediated phenomenon. A study from our institution has demonstrated that there is a distinct immunophenotype with high levels of interleukin activation in children with PIMS-TS.12 The key question of how systemic inflammation might
lead to CNS symptoms often implicates blood-brain barrier integrity or direct transfer via the circumventricular organs and regions devoid of a barrier.\\textsuperscript{13}

Of the surviving 8 children with PIMS-TS and neurologic symptoms, only half have returned to their baseline at 3-month follow-up. Similar difficulties are described short-term complications of ICU admissions in children.\\textsuperscript{14} However, the important observation in our study is that those with residual reported symptoms had higher inflammatory markers at presentation. Systemic hyperinflammation may acutely influence brain structural connections and could lead to long-term neurocognitive sequelae, as seen in some patients following autoimmune encephalitis.\\textsuperscript{15}

This single-center retrospective observational study spans a short period where the case definition and management of a newly evolving disease entity was being established and has inherent limitations. The full spectrum of the neurologic symptoms would have been underestimated in children in critical care and may result in inaccurate cross-sectional comparison. In addition, treatment effects cannot be systematically evaluated, as within this cohort, virtually all have been given first-line immune therapy (70/75, 93%), and that therapy was variably escalated to second line in 28/75 (37%).

Nevertheless, we were able to demonstrate that children with neurologic features during acute PIMS-TS illness have a higher burden of inflammation, which may in turn influence their recovery, raising the question as to the utility of these biomarkers in directing inflammation treatments. Presence of neurologic features should alert the clinician to the potential for more systemic inflammation. Further larger scale multicenter collaborative studies are already being initiated\\textsuperscript{16} to map the fuller spectrum of acute and long-term neurologic features of PIMS-TS and to evaluate the treatment effects of attenuating inflammation on neurologic outcome.

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**Disclosure**
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### Appendix 1 Authors

| Name                  | Location                                | Contribution                                                                 |
|-----------------------|-----------------------------------------|------------------------------------------------------------------------------|
| Mario Sa, MD          | Evelina London Children's Hospital (Evelina), United Kingdom | Designed and conceptualized the study; major role in the acquisition of data; analyzed the data; and drafted the manuscript |
| Luwaiza Mirza, BSc    | King's College London, United Kingdom    | Major role in the acquisition of data and analyzed the data                   |
| Michael Carter, MD    | King's College London, United Kingdom    | Major role in the acquisition of data and analyzed the data                   |
| Lalani Carlton Jones, MD | Evelina, United Kingdom                      | Revised the manuscript for intellectual content                             |
| Vasantha Gowda, MD    | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Jennifer Handforth, MD | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Tammy Hedderly, MD    | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Julia Kenny, PhD      | Evelina, United Kingdom                  | Major role in the acquisition of data and revised the manuscript for intellectual content |
| Karine Lascelles, MD  | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Jean-Pierre Lin, PhD  | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Daniel Lumsden, PhD   | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Marilyn McDougall, MD | Evelina, United Kingdom                  | Major role in the acquisition of data and revised the manuscript for intellectual content |
| Owen Miller, MD       | Evelina, United Kingdom                  | Major role in the acquisition of data and revised the manuscript for intellectual content |
| Thomas Rossor, MD     | Evelina, United Kingdom                  | Revised the manuscript for intellectual content                             |
| Vinay Shivamurthy, MD | Evelina, United Kingdom                  | Major role in the acquisition of data and revised the manuscript for intellectual content |

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Appendix 1 (continued)

| Name              | Location                | Contribution                                                                 |
|-------------------|-------------------------|-------------------------------------------------------------------------------|
| Ata Siddiqui, MD  | Evelina, United Kingdom | Revised the manuscript for intellectual content                              |
| Rahul Singh, MD   | Evelina, United Kingdom | Revised the manuscript for intellectual content                              |
| Shan Tang, PhD    | Evelina, United Kingdom | Revised the manuscript for intellectual content                              |
| Marie White, MD   | Evelina, United Kingdom | Major role in the acquisition of data and revised the manuscript for intellectual content |
| Susan Byrne, PhD  | Evelina, United Kingdom | Designed and conceptualized the study; analyzed the data and revised the manuscript for intellectual content |
| Ming Lim, PhD     | Evelina, United Kingdom | Designed and conceptualized the study; analyzed the data and revised the manuscript for intellectual content |

Appendix 2 Coinvestigators

Coinvestigators are listed at links.lww.com/NXI/A473.
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In the article “Systemic Inflammation Is Associated With Neurologic Involvement in Pediatric Inflammatory Multisystem Syndrome Associated With SARS-CoV-2” by Sa et al.,¹ the first sentence under Results in the Abstract should read, “75 patients with PIMS-TS were identified, 9 (12%) had neurologic involvement: altered consciousness (3), behavioral changes (3), focal neurology deficits (2), persistent headaches (2), hallucinations (2), excessive sleepiness (1), and new-onset focal seizures (1).” The authors regret the error.

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1. Sa M, Mirza L, Carter M, et al. Systemic inflammation is associated with neurologic involvement in pediatric inflammatory multisystem syndrome associated with SARS-CoV-2. Neurol Neuroimmunol Neuroinflamm. 2021;8(4):e999. doi:10.1212/NXI.000000000000999.