Clinical Implications of Procalcitonin in Kawasaki Disease: A Useful Candidate For Differentiating From Sepsis and Evaluating IVIG Responsiveness

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Methods

Kawasaki disease (KD), also known as mucocutaneous lymph node syndrome, is an acute, self-limited febrile illness and predominantly affects the cardiovascular system in young children less than 5 years of age. According to a recent epidemiological survey, an increasing trend in the incidence of KD has been noticed in China, with almost a one-fold increase from 50.5 per 100,000 children in 2008 to 94.7 per 100,000 children in 2017 [1, 2]. Depending on the 2017 American Heart Association (AHA) guidelines, the diagnostic criteria of KD include fever ≥ 5 days combined with at least 4 of the 5 principal clinical features. Besides, incomplete KD should be considered in any case with persistent unexplained fever, fewer than 4 of the major clinical features, and compatible laboratory or echocardiographic findings [3]. Coronary artery abnormalities (CAAs) serve as the most important prognostic factor and occur in almost 20% of KD patients based on the database of the 24th nationwide survey in Japan from January 2015 to December 2016 [4]. High-dose intravenous immunoglobulin (IVIG) plus aspirin is the first-line treatment for not only suppressing systemic inflammation in the acute episode of KD but also decreasing the incidence of CAAs to 6.8% [5]. However, approximately 10–20% of KD patients still have recrudescence or persistent fever at least 36 hours after completion of the initial IVIG infusion and are termed IVIG non-responders [2, 4]. To date, the accurate etiology of KD remains unknown, and recent consensus is that an infectious trigger, exaggerated immune response and inflammatory cascades may occur in genetically susceptible children who subsequently develop KD. In a case-control study from Yale University School of Medicine, the detection rate of New Haven coronavirus (HCOV-NH) in respiratory specimens of KD patients was significantly higher than that of the normal control subjects (72.7% versus 4.5%), with an odds ratio (OR) of 16.0 [6]. In addition, Khor et al. [7] conducted a genome-wide association study encompassing 2173 individuals with KD, and revealed that FCGR2A was identified as a susceptibility locus for KD in both Asian and North American populations (OR = 1.32).

Currently, no laboratory test is proved to be specific for the diagnosis of KD apart from the increases in several inflammatory mediators, such as white blood cell (WBC), absolute neutrophil counts (ANC), C-reactive protein (CRP), and so on. In our previous study, 70 children with KD were recruited to observe the alterations in the above 3 inflammatory mediators, and the findings revealed that the mean concentrations of WBC, ANC and CRP were 13.03 ×10⁹/l, 8.43 ×10⁹/l and 54.38 mg/l, respectively [8]. Although the up-regulated syntheses of inflammatory mediators have been universally recognized in the acute episode of KD, their changes are still ubiquitous in all kinds of infectious diseases. According to the report of Reyes et al. [9], the mean concentrations of WBC, ANC and CRP were 11.02 ×10⁹/l, 6.84 ×10⁹/l and 66.80 mg/l in 20 patients with sepsis respectively, appear to be identical to that in KD patients. Given this background, a better mediator is warranted to both reflect the severity of systemic inflammation and differentially diagnose KD from other infectious diseases.

Procalcitonin (PCT), a protein of 116 amino acids with molecular weight of 13 KDa, was primarily isolated 45 years ago as prohormone of calcitonin from C-cell of the thyroid gland and intracellularly degraded by proteolytic enzymes into the active hormone [10]. Although the circulating level of PCT in healthy subjects is far below detection limit, it elevates selectively in the bacterial inflammatory processes and is subjected to a 100-fold increase in the setting of sepsis [11]. Harbarth et al. [12] assessed the diagnostic value of PCT in 78 consecutive patients admitted with acute systemic inflammatory response syndrome (SIRS) and sepsis, and found that median PCT concentrations on admission were 0.6 ng/ml for SIRS, 3.5 ng/ml for sepsis and 21.3 ng/ml for septic shock, respectively; PCT > 1.1 ng/ml yielded a sensitivity of 97% and a specificity of 78% to differentiate SIRS from sepsis, with an area under the receiver operating characteristic curves (AUC) of 0.92. Besides serving as a good discriminative biomarker between sepsis and SIRS, PCT can be also used for monitoring the response to antimicrobial therapy. Nobre et al. [13] conducted a randomized, controlled, open interventional trial involving 79 patients with sepsis, and found that antibiotics for non-localized infections could safely be ceased, once PCT fell to < 0.25 ng/ml or > 90% from its peak concentration. In the last decade, a limited number of studies on serum PCT have been undertaken in KD patients. By the report of Okada et al. [14], a significant increase of PCT was noted in KD patients (2.3 ng/ml) compared with the healthy children (0.2 ng/ml), and moreover, the optimal cut-off value of 3.0 ng/ml enhanced the prediction rate of CAAs. However, further evidence for the association between PCT and KD should also be provided.

Abstract

Objective Kawasaki disease (KD) is a common childhood vasculitis absent of the specific laboratory definitions, besides a significant elevation in several inflammatory mediators, such as procalcitonin (PCT). However, whether PCT can serve as a useful candidate for differentiating KD from sepsis, and even for predicting incomplete KD, intravenous immunoglobulin (IVIG) nonresponsiveness and coronary artery abnormalities (CAAs) remains unclear.

Methods 254 Chinese KD children were enrolled and divided into 6 subgroups, including complete KD, incomplete KD, IVIG-responsive KD, IVIG-nonresponder KD, KD with CAAs and KD without CAAs. Blood samples were collected from all subjects within 24-h pre- and 48-h post-IVIG infusion, respectively. PCT, C-reactive protein, sedimentation rate and blood cell counts were detected. In addition, both 261 sepsis children and 251 healthy children sex- and age-matched with KD children were enrolled in the same period.

Results (1) PCT experienced the highest increase in sepsis patients before antibiotic therapy, followed by acute KD patients and the healthy controls. (2) The proportion of KD patients with a PCT concentration below 0.25 ng/ml was 11 folds higher than that of sepsis patients. (3) PCT had a sensitivity of 91.7% and a specificity of 30.3% at a cut-off value of >0.15 ng/ml to predict IVIG nonresponsiveness, and the proportion of IVIG-nonresponders with a PCT concentration of 0.25-0.50 ng/ml was 2 folds higher than that of IVIG-responders.

Conclusions The PCT concentrations below 0.25 ng/ml may be useful for discriminating KD from sepsis, and moreover, the PCT concentrations of 0.25-0.50 mg/ml may be helpful in predicting IVIG nonresponsiveness.

Introduction

Kawasaki disease (KD), also known as mucocutaneous lymph node syndrome, is an acute, self-limited febrile illness and predominantly affects the cardiovascular system in young children less than 5 years of age. According to a recent epidemiological survey, an increasing trend in the incidence of KD has been noticed in China, with almost a one-fold increase from 50.5 per 100,000 children in 2008 to 94.7 per 100,000 children in 2017 [1, 2]. Dependent on the 2017 American Heart Association (AHA) guidelines, the diagnostic criteria of KD include fever ≥ 5 days combined with at least 4 of the 5 principal clinical features. Besides, incomplete KD should be considered in any case with persistent unexplained fever, fewer than 4 of the major clinical features, and compatible laboratory or echocardiographic findings [3]. Coronary artery abnormalities (CAAs) serve as the most important prognostic factor and occur in almost 20% of KD patients based on the database of the 24th nationwide survey in Japan from January 2015 to December 2016 [4].

Methods

Subjects
The present study was carried out retrospectively in Department of Pediatrics, the First Affiliated Hospital of Anhui Medical University. A total of 254 children having KD were recruited from July 2015 to July 2020. According to the 2017 AHA guidelines [3], the diagnostic criteria for complete KD include the presence of ≥5 days of fever and ≥4 of the following 5 main features: (1) bilateral conjunctival injection without exudates; (2) changes in the oral mucosa, such as erythema and cracking lips, erythema of the pharynx, strawberry tongue; (3) changes in extremities, such as redness and swelling in the acute phase, periangual desquamation in the subacute phase; (4) polymorphous exanthema; and (5) cervical lymphadenopathy (≥1.5cm in diameter), usually unilateral. However, patients with fever for ≥5 days and at least 2 of the main features were diagnosed as having incomplete KD, if no other febrile illnesses could explain the disease process. All patients received the standard therapy for KD, including a single infusion of high-dose IVIG (2 g/kg) and aspirin (30-50 mg/kg/d), within 10 days since the onset of fever. IVIG-nonresponsive KD was defined as persistent or recrudescent fever ≥36 hours after finishing the initial IVIG infusion. CAAs were defined as a coronary artery having an internal diameter of at least 3 mm in children <5 years or at least 4 mm in children ≥5 years, or a segment with an internal diameter at least 1.5 times larger than that of an adjacent segment by echocardiogram. In addition, 261 sepsis children sex- and age-matched with KD children and 251 healthy children sex- and age-matched with KD children were enrolled in the same period. According to the Third International Consensus Definitions for Sepsis and Septic Shock (Sepsis-3), sepsis was defined as life-threatening organ dysfunction caused by a dysregulated host response to infection [15].

**Laboratory analysis**

Approval for the present study was acquired from the Medical Ethic Committee of the First Affiliated Hospital of Anhui Medical University and obtained consent from parents of all subjects. Blood samples were collected from all subjects within 24-h pre- and 48-h post-IVIG infusion, respectively, including WBC, ANC, CRP, sedimentation rate (ESR), red blood cells counts (RBC), hemoglobin (Hb), platelet count (PLT) and PCT.

**Statistical analysis**

Normally distributed continuous data were expressed as mean±SD. Comparisons of the frequencies between groups were analyzed using Chi-square tests and comparisons among groups were made using ANOVA. Serum data were analyzed using the 2 independent samples t test or Mann-Whitney U test, and the paired t test or Wilcoxon test for comparison of pre-IVIG to post-IVIG data. Pearson correlation coefficients were reported for PCT and other variables of interest. Cutoff value, sensitivity and specificity of PCT were identified by receiver operating characteristic (ROC) curve. A value of p<0.05 was considered significant. Statistical analysis was performed using the statistical package for social studies SPSS version 25.0.

**Results**

**Patient characteristics**

254 KD patients (138 males and 116 females) were recruited during the past 5 years, with a mean age of 32.11±27.35 months and a range from 2 months to 13 years. 261 sepsis patients (146 males and 115 females) were recruited in the present study, with a mean age of 36.18±31.54 months and a range from 2 months to 13 years. No significant differences were observed in age (t=2.71, p>0.05) and gender (x²=0.26, p>0.05) among KD patients, sepsis patients and the healthy controls.

Among 254 KD patients, all 5 classic diagnostic criteria for KD were met in 72 cases (28.35%), 4 criteria in 104 (40.94%), 3 criteria in 48 (18.90%), and 2 criteria in 30 (11.81%). Therefore, 176 patients (69.29%) had complete KD, including 101 males and 75 females with a mean age of 32.98±27.56 months; 78 patients (30.71%) had incomplete KD, including 37 males and 41 females with a mean age of 30.17±26.94 months. According to the fever duration after the initial IVIG infusion, 17 KD patients (6.70%) including 8 males and 9 females with a mean age of 29.65±23.54 months had a mean persistent fever of 72.38±23.21 hours and were identified as IVIG-nonresponders. In contrast, 237 patients (93.30%) including 130 males and 107 females with a mean age of 32.29±27.63 months were diagnosed as IVIG-responders and exhibited a noted decrease in fever duration (7.89±6.89 hours; t=11.08, p<0.05). Based on the internal diameter of coronary artery, 20 KD patients (7.87%) including 13 males and 7 females with a mean age of 36.70±32.00 months were defined as having CAAs after IVIG therapy (left coronary artery: 3.09±0.56 mm; right coronary artery: 3.19±1.31 mm). In contrast, 234 patients (92.13%) including 125 males and 109 females with a mean age of 31.72±26.96 months had normal coronary arteries after IVIG therapy (left coronary artery: 2.04±0.34 mm, right coronary artery: 1.96±0.31 mm). The internal diameters of both coronary arteries were significantly larger in KD patients with CAAs (left coronary artery: t=8.30, p<0.05; right coronary artery: t=4.10, p<0.05), compared with KD patients without coronary arteritis. Overall, the mean age and male/female ratio were almost identical between complete KD patients and incomplete KD patients (x²=2.16, p>0.05; t=0.76, p>0.05), between IVIG-nonresponders and IVIG-responders (x²=0.39, p>0.05; t=0.38, p>0.05), and between KD patients with CAAs and KD patients without CAAs (x²=1.00, p>0.05; t=0.78, p>0.05).

**Laboratory findings**

Blood cell counts and inflammatory mediators in KD patients, sepsis patients and the healthy controls are shown in Table 1. Significant differences in WBC, ANC, RBC, Hb, PLT, CRP, ESR and PCT were noted among KD patients, sepsis patients and the healthy controls (p<0.05). More specifically, KD patients had a significant increase in WBC, PLT, CRP, ESR and PCT, whereas a significant decrease in RBC and Hb compared with the healthy controls regardless of pre-IVIG and post-IVIG therapy (p<0.05). KD patients before IVIG therapy had a significant increase in ANC compared with the healthy controls (p<0.05), whereas no significant difference was found in ANC between KD patients after IVIG therapy and the healthy controls (p=0.05). KD patients had a significant increase in PLT and ESR, whereas a significant decrease in WBC, ANC, RBC, Hb, CRP and PCT compared with sepsis patients regardless of pre-IVIG and post-IVIG therapy (p<0.05). Sepsis patients had a significant increase in WBC, ANC, CRP, ESR and PCT, whereas a significant decrease in RBC, Hb and PLT compared with the healthy controls (p<0.05).
As for KD patients, WBC, ANC, CRP, ESR and PCT showed a significant decrease, whereas PLT experienced a significant increase after IVIG therapy (p<0.05). The associations of PCT with blood cell counts and other inflammatory mediators in both KD patients and sepsis patients before therapy are shown in Fig. 1 and Table 2. PCT was positively correlated with CRP, negatively correlated with RBC and PLT in both KD patients and sepsis patients (p<0.05). Additionally, PCT was positively correlated with ANC, negatively correlated with Hb in KD patients (p<0.05).

The role of PCT in differential diagnosis between KD and sepsis

In order to evaluate the effectiveness of PCT in differential diagnosis between KD and sepsis, the distributions of PCT concentrations are depicted (Fig. 2a). Among 254 KD patients, 101 (39.76%) had a PCT concentration below 0.25 ng/ml, 64 (25.20%) had a PCT concentration of 0.25-0.50 ng/ml, 30 (11.81%) had a PCT concentration of 0.50-1.00 ng/ml, 54 (21.26%) had a PCT concentration of 1.00-10.00 ng/ml, and 5 (1.97%) had a PCT concentration of ≥10.00 ng/ml. In comparison, among 261 sepsis patients, 9 (3.45%) had a PCT concentration below 0.25 ng/ml, 11 (4.21%) had a PCT concentration of 0.25-0.50 ng/ml, 40 (15.33%) had a PCT concentration of 0.50-1.00 ng/ml, 145 (55.55%) had a PCT concentration of 1.00-10.00 ng/ml, and 56 (21.46%) had a PCT concentration of ≥10.00 ng/ml. The proportion of KD patients with a PCT concentration below 0.25 ng/ml was significantly higher than that of sepsis (p<0.05). Therefore, PCT concentrations below 0.25 ng/ml may be candidates for differential diagnosis between KD and sepsis.

Clinical classification

The differences of blood cell counts and inflammatory mediators between complete KD patients and incomplete KD patients are presented in Table 3. No significant differences in WBC, ANC, RBC, Hb, PLT, CRP, ESR and PCT were observed between complete KD patients and incomplete KD patients before IVIG therapy (p>0.05). Furthermore, WBC, ANC, CRP, ESR and PCT showed a significant decrease, whereas PLT experienced a significant increase after IVIG therapy in both complete KD patients and incomplete KD patients (p<0.05).

IVIG responsiveness

The differences of blood cell counts and inflammatory mediators between IVIG-responders and IVIG-nonresponders are presented in Table 3. No significant differences in WBC, ANC, RBC, Hb, PLT, CRP, ESR and PCT were observed between IVIG-responders and IVIG-nonresponders before IVIG therapy (p>0.05); ANC and CRP declined to normal, whereas PLT remained significantly higher after IVIG therapy in both IVIG-responders and IVIG-nonresponders (p<0.05). The therapy of IVIG significantly inhibited the increase of WBC, ESR and PCT in IVIG-responders (p<0.05). However, the above 3 inflammatory mediators were at a higher level in IVIG-nonresponders persistently (p<0.05). The ROC curve was generated to determine the best cut-off of WBC, ESR and PCT for prediction of IVIG-nonresponders. As shown in Fig. 3, the AUC value for WBC in prediction of IVIG-nonresponders was 0.59, and the estimated sensitivity and specificity were 50.00% and 68.60% with a cutoff of WBC≥15.61×10^9/L, respectively. The AUC value for ESR in prediction of IVIG-nonresponders was 0.54, and the estimated sensitivity and specificity were 66.70% and 60.60% with a cutoff of ESR≥66.50 mm/h, respectively. The AUC value for PCT in prediction of IVIG-nonresponders was 0.55, and the estimated sensitivity and specificity were 91.70% and 30.30% with a cutoff of PCT>0.15 ng/ml, respectively.

To assess the value of PCT for the early recognition of IVIG nonresponsiveness, the distributions of PCT concentrations are depicted in both IVIG-responders and IVIG-nonresponders (Fig. 2b). Among 17 IVIG-nonresponders, 4 (23.53%) had a PCT concentration below 0.25 ng/ml, 7 (41.18%) had a PCT concentration of 0.25-0.50 ng/ml, 4 (23.53%) had a PCT concentration of 0.50-1.00 ng/ml, and 2 (11.76%) had a PCT concentration of 1.00-10.00 ng/ml. In comparison, among 237 IVIG-nonresponders, 95 (40.09%) had a PCT concentration below 0.25 ng/ml, 57 (24.05%) had a PCT concentration of 0.25-0.50 ng/ml, 27 (11.39%) had a PCT concentration of 0.50-1.00 ng/ml, 53 (22.36%) had a PCT concentration of 1.00-10.00 ng/ml, and 5 (2.11%) had a PCT concentration of ≥10.00 ng/ml. The proportion of IVIG-nonresponders with a PCT concentration of 0.25-0.50 ng/ml was significantly higher than that of IVIG-responders (p<0.05). Therefore, PCT concentrations lying in the 0.25-0.50 ng/ml range may be candidates for differential diagnosis between IVIG-responders and IVIG-nonresponders.

Coronary artery involvement

The differences of blood cell counts and inflammatory mediators between KD patients with and without CAAs are presented in Table 3. No significant differences in WBC, ANC, RBC, Hb, PLT, CRP, ESR and PCT were observed between complete KD patients and incomplete KD patients before IVIG therapy (p>0.05); Furthermore, WBC, ANC, CRP, ESR and PCT showed a significant decrease, whereas PLT experienced a significant increase after IVIG therapy in both complete KD patients and incomplete KD patients (p<0.05).

Discussion

PCT is a common serum marker of the inflammatory response and persistently elevated in severe bacterial infection. KD belongs to systemic vasculitis and occurs mainly in children younger than 5 years old. Although the etiology of KD remains unclear, many current researches have indicated that infection is the most prevalent trigger [6,16,17]. In this circumstance, whether PCT can serve as a useful marker for differentiating KD from severe bacterial infection and even guiding antibiotic therapy should be noted by pediatric rheumatologists. In the present study, we compared serum level of PCT among acute KD patients, sepsis patients and the healthy controls, and found that it experienced the highest increase in sepsis patients before antibiotic therapy, followed by acute KD patients and the healthy controls. Consistently, two controlled clinical trials from Chongqing and Guangdong, China [18,19], indicated that serum PCT in acute KD patients had a significant increase compared with the healthy controls (1.4 ng/ml vs 0.2 ng/ml), whereas was significantly lower than that in sepsis patients (1.4 ng/ml vs 4.1 ng/ml). Based on these current evidence, serum PCT in acute KD patients is almost 7-20 folds higher above baseline, whereas 3-5 folds lower than that in sepsis patients. Although the detailed mechanisms of PCT upregulation is still uncertain in acute KD, some studies have demonstrated...
that inflammatory cytokines secreted by activated monocytes/macrophages may be involved. Balog et al. [20] cultured human peripheral monocytes with Staphylococcus aureus (10⁸/ml) to stimulate the release of tumor necrosis factor-α (TNF-α), and found that the mean fluorescence intensities of PCT increased from 74 to 131 after an 18-hour stimulation by S. aureus, which was almost totally abrogated by anti-TNF-α monoclonal antibodies. Oberhoffer et al. [21] assessed the possible expression of PCT in human peripheral monocytes by reverse transcriptase-polymerase chain reaction (RT-PCR), and revealed that the expression of PCT mRNA showed a maximal 90-fold increase after stimulation by TNF-α (50 ng/ml), a maximal 35-fold increase by interleukins (IL) -6 (50 ng/ml), a maximal 18-fold increase by IL-1β (50 ng/ml), a maximal 15-fold increase by IL-2 (50 ng/ml), respectively. Thus, TNF-α had the most pronounced stimulatory effect on the expression of PCT in human peripheral monocytes to date.

One main objective of our study was to evaluate the effectiveness of PCT in differential diagnosis between KD and sepsis. In the present study, we observed that the proportion of KD patients with a PCT concentration below 0.25 ng/ml was 11 folds higher than that of sepsis patients. Based on this finding, PCT concentrations below 0.25 ng/ml may be useful for discriminating KD from sepsis. Recently, Liu et al. [19] developed a novel nomogram model to differentiate KD from sepsis, in which PCT ≤ 0.5 ng/ml was regarded as a strong predictor for KD with 57 points, and associated with a 3.41-fold increase in KD probability. However, in a retrospective study encompassing 49 KD patients and 24 sepsis patients from Korea, Lee et al. [22] divided both KD and sepsis patients into <0.25 ng/ml group, 0.25-1.0 ng/ml group and >1.0 ng/ml group based on PCT concentrations, and found that PCT was not helpful for discriminating KD from sepsis. Because of different identification criteria, analytical methodologies and genetic background, the role of PCT in discriminating KD from sepsis is still controversial. Therefore, further studies of larger sample size and multi-center will be necessary to consolidate our findings.

The other main objective of our study was to assess the value of PCT for the early recognition of IVIG nonresponsiveness. The present study demonstrated that PCT had a sensitivity of 91.7% and a specificity of 30.3% at a cut-off value of >0.15 ng/ml to predict IVIG nonresponsiveness, and the proportion of IVIG-nonresponders with a PCT concentration of 0.25-0.50 ng/ml was 2 folds higher than that of IVIG-responders. In view of the above, PCT concentrations lying in the 0.25-0.50 ng/ml range may be useful for predicting IVIG nonresponsiveness. Nakamura et al. [23] established a multivariate logistic regression model and revealed that PCT had a sensitivity of 46.4% and a specificity of 93.9% at a cut-off value of >2.18 ng/ml to predict IVIG nonresponsiveness and possessed the largest AUC of 0.82 than existing 3 refractory prediction scores: Gunma score [24], Kurume score [25] and Osaka score [26]. Consistently, a clinical trial from Boston Children's Hospital, America, indicated that serum PCT ≥0.5 ng/ml was correlated with nonresponsiveness to IVIG and admission to the pediatric intensive care unit (PICU), and moreover, a PCT concentration of ≥ 4.3 ng/ml showed the most sensitive and specific values for predicting IVIG nonresponsiveness [27]. However, in a prospective study from Sichuan, China, Shao et al. [28] recruited 530 KD patients from January 2015 to March 2019, and discovered that PCT may not be suitable as an independent predictive factor for both initial and repeated IVIG nonresponsiveness. A large controversy exists in these current scattered studies regarding the predictive role of PCT to IVIG nonresponsiveness.

CAAs secondary to KD represent the major contributors to morbidity and mortality in both the acute stage and the long-term. The pathological changes of CAAs compass necrotizing arteritis and inflammatory cell infiltration in the first 2 weeks, and progress to luminal myofibroblastic proliferation thereafter [29]. Furthermore, several inflammatory cytokines including nuclear factor-κB, IL-1β, IL-6, TNF-α and transforming growth factor-β have been reported to participate in CAA onset [30-33]. In acute KD, the overexpression of PCT is associated with elevated inflammatory cytokines. On this background, whether PCT may serve as a predictive factor for CAAs should be elucidated. This study investigated the association between serum PCT and development of CAAs, and found that PCT was not helpful in screening CAAs, which was confirmed with the findings of researches from Europe and North America [34, 27]. On the contrary, a retrospective research encompassing 160 KD patients from Japan revealed that PCT had a sensitivity of 67% and a specificity of 56% at a cut-off value of >0.5 ng/ml to predict CAAs and possessed the highest OR compared with the other inflammatory mediators, such as WBC, ANC, PLT and CRP [35]. Because of different populations and sample sizes, a wide heterogeneity is noted in the predictive role of PCT to CAAs among these current studies, and thus, further studies are warranted to focus on this issue more deeply.

Additionally, the present study also assessed the relationship of serum PCT with clinical classification in the course of KD, and found that no significant difference was determined in serum PCT between complete KD patients and incomplete KD patients. Lee et al. and Shao et al. [22, 28] respectively observed 49 KD Korean and 530 KD Chinese, and both found that PCT could not discriminate incomplete KD patients from complete KD patients.

Conclusions And Perspectives

Thus, The PCT concentrations below 0.25 ng/ml may be useful for discriminating KD from sepsis, and moreover, the PCT concentrations of 0.25-0.50 ng/ml may be helpful in predicting IVIG nonresponsiveness. Recently, PCT was also found to be a predictive factor for multisystem inflammatory syndrome in children (MIS-C) secondary to coronavirus disease 2019 (COVID-19). Based on the latest data from the Coronavirus Resource Center of Johns Hopkins University and Medicine, COVID-19 has caused more than 34 million confirmed cases worldwide and >1,020,000 fatalities up to October 1st, 2020 [36]. Accumulative studies have indicated that all humans including children appear susceptible to COVID-19, in spite the majority of pediatric patients have mild or even no symptoms [37, 38]. However, in early May, 2020, the United Kingdom and several European countries reported that a hyperinflammatory process triggered by COVID-19 could significantly increase the risk of MIS-C in pediatric patients, which was very similar to incomplete KD [39, 40]. By the report of the New York State Department of Health, the incidence of MIS-C was 2 per 100,000 persons younger than 21 years of age, and MIS-C patients had a skyrocketing concentration of serum PCT (6.2 ng/ml) [41].

Declarations

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41. Tables

Table 1 Blood cell counts and inflammatory mediators in KD patients, sepsis patients and the healthy controls

|                      | KD patients  | Sepsis patients | Healthy controls |
|----------------------|--------------|-----------------|-----------------|
|                      | (n=254)      | (n=261)         | (n=251)         |
| WBC (×10^9/l)        | 13.52±4.89   | 19.62±10.19     | 8.26±2.31       |
| ANC (×10^9/l)        | 8.97±4.41    | 14.25±8.84      | 3.73±2.24       |
| RBC (×10^12/l)       | 4.21±0.42    | 4.34±0.61       | 4.60±0.48       |
| Hb (g/l)             | 111.22±12.54 | 115.49±11.73    | 123.70±11.73    |
| PLT (×10^9/l)        | 359.01±125.62| 305.89±121.49   | 326.59±98.21    |
| CPR (mg/l)           | 60.71±45.52  | 60.57±20.94     | 9.32±7.79       |
| ESR (mm/h)           | 61.07±45.52  | 7.47±10.25      | 1.35±0.90       |
| PCT (ng/ml)          | 359.01±125.62| 305.89±121.49   | 9.32±7.79       |

Data are presented as the mean±SD. *p < 0.05, significantly difference between Pre-IVIG and Post-IVIG KD patients; **p < 0.05, significantly difference between KD patients and the healthy controls; ***p < 0.05, significantly difference between KD patients and sepsis patients; #p < 0.05, significantly difference between sepsis patients and the healthy controls

Table 2 The associations of PCT with blood cell counts and other inflammatory mediators in KD patients and sepsis patients (r, p)
| PCT  | WBC  | ANC  | RBC  | Hb   | PLT  | CPR  | ESR  |
|------|------|------|------|------|------|------|------|
| (ng/ml) | (×10⁹/l) | (×10⁹/l) | (×10¹²/l) | (g/l) | (×10⁹/l) | (mg/l) | (mm/h) |
| KD patients | + | − | − | − | − | + |
| 0.128, 0.062 | 0.287, 0.000* | -0.149, 0.038* | -0.148, 0.036* | -0.425, 0.000* | 0.470, 0.000* | 0.125, 0.079 |
| Sepsis patients | − | − | + |
| -0.062, 0.316 | 0.014, 0.822 | -0.133, 0.037* | -0.028, 0.654 | -0.218, 0.000* | 0.232, 0.000* | -0.092, 0.444 |

+, a positive correlation; −, a negative correlation; *p < 0.05

Table 3 Blood cell counts and inflammatory mediators in patients with different types of KD

| KD patients (n=254) | WBC  | ANC  | RBC  | Hb   | PLT  | CPR  | ESR  | PCT  |
|---------------------|------|------|------|------|------|------|------|------|
|                     | (×10⁹/l) | (×10⁹/l) | (×10¹²/l) | (g/l) | (×10⁹/l) | (mg/l) | (mm/h) | (ng/ml) |
| Complete KD patients |     |     |     |     |     |     |     |     |
| Pre-IVIG            | 13.35±4.45* | 8.99±3.86* | 4.21±0.40 | 111.40±12.85 | 353.87±128.26* | 62.56±42.83* | 64.87±21.88* | 5.39±27.27* |
| Post-IVIG           | 9.89±4.25* | 3.71±3.70* | 4.19±0.38 | 110.98±10.11 | 499.37±152.14* | 7.47±10.73* | 60.21±20.46* | 0.21±0.38* |
| Incomplete KD patients (n=78) |     |     |     |     |     |     |     |     |
| Pre-IVIG            | 14.22±5.53* | 9.17±4.99* | 4.16±0.43 | 110.45±12.11 | 367.51±125.28* | 69.63±48.87* | 66.56±22.27* | 2.21±4.10* |
| Post-IVIG           | 9.56±3.13* | 3.24±1.88* | 4.20±0.42 | 110.72±12.02 | 546.46±168.74* | 7.76±9.50* | 60.12±22.79* | 0.16±0.15* |
| IVIG-responders (n=237) |     |     |     |     |     |     |     |     |
| Pre-IVIG            | 13.60±4.80* | 8.97±4.25* | 4.21±0.41 | 111.32±12.54 | 358.34±124.46* | 64.93±45.54* | 64.99±20.90* | 0.20±0.32* |
| Post-IVIG           | 9.66±3.67* | 3.43±2.89* | 4.21±0.38 | 111.34±10.54 | 513.77±155.38* | 7.49±10.13* | 59.34±20.90* | 0.16±0.15* |
| IVIG-nonresponders (n=17) |     |     |     |     |     |     |     |     |
| Pre-IVIG            | 13.97±5.26 | 10.09±4.03* | 4.01±0.51 | 107.69±13.55 | 355.00±166.88* | 61.91±32.04* | 70.50±25.71 | 0.40±0.20 |
| Post-IVIG           | 11.60±6.55 | 5.61±6.37* | 3.89±0.42 | 103.92±11.70 | 517.80±205.90* | 8.64±13.78* | 71.83±21.30 | 0.19±0.30 |
| KD patients without CAAs (n=234) |     |     |     |     |     |     |     |     |
| Pre-IVIG            | 13.67±4.85* | 9.10±4.24* | 4.18±0.42 | 110.72±12.64 | 358.52±126.67* | 66.52±45.45* | 66.02±21.59* | 4.69±23.66* |
| Post-IVIG           | 9.78±3.94* | 3.58±3.28* | 4.18±0.39 | 110.61±10.62 | 514.24±153.46* | 7.83±10.72* | 60.35±21.21* | 0.20±0.34* |
| KD patients with CAAs (n=20) |     |     |     |     |     |     |     |     |
| Pre-IVIG            | 13.14±4.50* | 8.40±4.22* | 4.35±0.41 | 115.11±11.64 | 353.74±136.72* | 45.44±31.70* | 57.08±25.45* | 1.31±1.71* |
| Post-IVIG           | 9.88±3.89* | 3.36±2.86* | 4.28±0.41 | 113.95±11.65 | 511.74±212.66* | 4.61±3.70* | 58.00±20.42 | 0.14±0.11* |

Data are presented as the mean±SD. *p < 0.05, significantly difference between Pre-IVIG and Post-IVIG KD patients