Diagnosis of incomplete Kawasaki disease

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Several authors suggested that the clinical characteristics of incomplete presentation of Kawasaki disease are similar to those of complete presentation and that the 2 forms of presentation are not separate entities. Based on this suggestion, a diagnosis of incomplete Kawasaki disease in analogy to the findings of complete presentation is reasonable. Currently, the diagnosis of incomplete Kawasaki disease might be made in cases with fewer classical diagnostic criteria and with several compatible clinical, laboratory or echocardiographic findings on the exclusion of other febrile illness. Definition of incomplete presentation in which coronary artery abnormalities are included as a necessary condition, is restrictive and specific. The validity of the diagnostic criteria of incomplete presentation by the American Heart Association should be thoroughly tested in the immediate future.

Key words: Mucocutaneous lymph node syndrome, Kawasaki disease, Kawasaki syndrome, Diagnosis

Introduction

Kawasaki disease is an acute, self-limited systemic type of vasculitis that occurs predominantly in young children. After the 1st report by Tomisaku Kawasaki in Japan, it is now recognized as the leading cause of acquired heart disease in children in developed countries. The etiology of Kawasaki disease is still unknown and no single pathognomonic clinical or laboratory finding for the diagnosis has been identified. To establish a diagnosis, we inevitably depend on the diagnostic criteria, which are the typical constellation of signs and symptoms, prepared by the Japanese Kawasaki Disease Research Committee or the American Heart Association (AHA). However, pediatricians sometimes encounter febrile children who do not fulfill the diagnostic criteria but have several findings compatible with those of Kawasaki disease. In this situation, the diagnosis of incomplete Kawasaki disease is a clinical challenge, which cannot be avoided by delaying the diagnosis because of the risk of coronary complications pertaining even to the incomplete presentation of the disease. The diagnosis of incomplete Kawasaki disease is reviewed in this paper.

Use of terms “incomplete” or “atypical”

The phrase “atypical Kawasaki disease” was initially used to describe patients with coronary complications who do not fulfill the classical diagnostic criteria. Recently, the term “atypical Kawasaki disease” has been used to describe patients with incomplete presentation of the disease, regardless of the presence of coronary complications, and is exchangeable for “incomplete Kawasaki disease.” Newburger et al. proposed that “atypical Kawasaki disease” should be reserved for patients with complications, e.g., renal impairment, that are generally not seen in Kawasaki disease. This proposition has been supported by other authors.
Diagnostic criteria of Kawasaki disease

The diagnostic criteria established by the AHA (Table 1) are different from the Japanese criteria (Table 2) with regard to fever prolongation (≥5 days) as a necessary criterion. In the Japanese criteria, fever prolongation is just an optional criterion and there is no description of the size of cervical lymph nodes (Table 2). In spite of rarely reported cases of Kawasaki disease without fever, a diagnosis of complete Kawasaki disease is not difficult in typical cases without regard to a selected criteria. Because of a need for early diagnosis and management, Kawasaki disease is not difficult in typical cases without regard to a selected criteria. Because of a need for early diagnosis and management, Kawasaki disease is not difficult in typical cases without regard to a selected criteria. Because of a need for early diagnosis and management, Kawasaki disease is not difficult in typical cases without regard to a selected criteria. Because of a need for early diagnosis and management

No other demographic features (including gender), except the prevalent age, has been reported to be different between patients with incomplete and complete presentation. The presence of classical clinical findings in patients with incomplete presentation is properly less frequent than in patients with complete presentation. The frequently reported findings which less frequently observed in incomplete presentation are cervical lymphadenopathy (19 to 38.6%) and extremity changes (21 to 44.3%), and this pattern of order of observed frequencies is equal to that noted in patients with complete presentation.

 incomplete Kawasaki disease as a continuous spectrum of complete presentation of the disease

The reported prevalence of incomplete presentation was 15 to 36.2% among patients with Kawasaki disease. Children with incomplete presentation were more likely to be at the extremes of the age spectrum as compared to those with complete presentation. The prevalence of incomplete presentation was relatively higher in the younger-aged patients. One study reported that patients with incomplete presentation were younger than patients with complete presentation. In other reports, there was no difference in the mean/median age between the 2 groups. However, relatively more children with incomplete presentation were in the extremes of the age spectrum (≤1 year old, or ≥5 to 9 years old). The risk of delayed diagnosis and management in young infants with incomplete presentation has been pointed out in the past.

No other demographic features (including gender), except the prevalent age, has been reported to be different between patients with incomplete and complete presentation. The presence of classical clinical findings in patients with incomplete presentation is properly less frequent than in patients with complete presentation. The frequently reported findings which less frequently observed in incomplete presentation are cervical lymphadenopathy (19 to 38.6%) and extremity changes (21 to 44.3%), and this pattern of order of observed frequencies is equal to that noted in patients with complete presentation. Comparison of laboratory findings (complete blood count, albumin, hepatic transaminase, acute phase reactants) between these 2 groups also showed significant differences. However, low levels of serum alanine transaminase, low levels of gamma glutamyl transferase, low frequency of hyponatremia, and low frequency of pyuria were reported in children with incomplete presentation. Several authors who had compared clinical characteristics between the 2 forms of Kawasaki disease suggested that they are not separate entities but are probably located along a continuous spectrum, placing more weight on demographic and laboratory similarities between the 2 groups.

In this suggestion, making the diagnosis of incomplete Kawasaki disease in analogy to the findings of complete presentation could be reasonable. Contradicting studies reported incomplete presentation of Kawasaki disease in children to be associated with higher risk, similar risk, or lower risk of coronary artery abnormalities compared to children with complete presentation. Sudo et al. concluded that the higher incidence of coronary artery lesions in patients with incomplete presentation was firstly due to diagnostic bias because of the use of echocardiography in the diagnostic process, and secondly due to delays in the treatment because of difficulties in making the diagnosis. Restricting the diagnosis of incomplete Kawasaki disease within children with echocardiographic coronary artery abnormalities will result in above mentioned diagnostic bias. Manlhiot et al. elucidated equal incidences of coronary artery abnormalities between both patient groups using their definition of incomplete presentation of Kawasaki disease in which the use of echocardiography is excluded. In children with incomplete Kawasaki disease, the time between onset of symptoms and diagnosis has been reported to be longer, and they are less likely to receive IVIG than children with complete presentation. Sudo et al. performed multivariate analysis to delineate risk factors of coronary artery lesions and showed that late IVIG

| Table 1. Classical Diagnostic Clinical Criteria of Kawasaki Disease Prepared by the American Heart Association |
|---|
| Fever persisting ≥5 days |
| Presence of ≥4 principal features |
| Changes in extremities |
| Polymorphous exanthema |
| Bilateral bulbar conjunctival injection without exudates |
| Changes in lips and oral cavity |
| Cervical lymphadenopathy (>1.5 cm diameter) |
| Exclusion of other diseases with similar findings |

| Table 2. Six principal symptoms in the diagnostic guidelines for Kawasaki disease prepared by the Japanese Kawasaki Disease Research Committee |
|---|
| Fever persisting ≥5 days |
| Bilateral conjunctival congestion |
| Changes of lips and oral cavity |
| Polymorphous exanthema |
| Changes of peripheral extremities |
| Acute non-purulent cervical lymphadenopathy |
treatment (≥7 days of illness) was a significant risk factor of coronary artery lesions in both patient groups.

Guidelines for the diagnosis of incomplete Kawasaki disease

The presence of coronary artery abnormalities has been used as criterion for the diagnosis of incomplete Kawasaki disease. According to the Japanese criteria, this criterion can only be used in cases with ≥4 principal symptoms. Many authors believed that this definition is too restrictive and specific. There are at least 3 problems with this restrictive definition of incomplete presentation. First of all, pediatric clinicians sometimes encounter cases in which Kawasaki disease is suspected with less than 4 principal symptoms. In the diagnostic criteria for incomplete presentation prepared by the AHA, a diagnosis of incomplete Kawasaki disease is possible in cases with fever and 2 principal features (according to Japanese criteria, 3 principal symptoms). Secondly, the prevalence of coronary artery abnormalities is in itself an unfavorable outcome of the disease. The duration of fever has been confirmed as an important risk factor of coronary artery abnormalities. Therefore, it may induce a delay of management to postpone diagnosis of incomplete presentation until confirming of coronary artery abnormality. One study in which the definition of incomplete Kawasaki disease included coronary involvement showed higher coronary complications in the group of infants younger than 6 months of age which had more cases of incomplete disease. Finally, the Japanese Ministry of Health criteria (Table 3B), which have been used as a standard for the diagnosis of coronary complications, are too restrictive. De Zorzi et al. showed that the z score of coronary arteries (adjusted for body surface area) was more than 2 in 27% of children with Kawasaki disease whose coronary artery diameters were classified as normal according to the Japanese Ministry of Health criteria. The increase of body surface area-adjusted z score of the coronary diameter was adopted as a standard of coronary artery abnormality by the American Heart Association.

Table 3. Supplemental Laboratory Criteria (A) and Echocardiographic Criteria (B) for the Diagnosis of Incomplete Kawasaki Disease Prepared by the American Heart Association

| (A)                  |
|---------------------|
| Serum albumin ≤3.0 g/dL |
| Anemia for age       |
| Elevation of alanine aminotransferase |
| Platelets after 7 days ≥450,000/mm³ |
| WBC ≥15,000/mm³      |
| Urine WBC ≥10/HPF   |

| (B)                |
|-------------------|
| Z score of LAD or RCA ≥2.5 |
| Coronary arteries meet Japanese Ministry of Health criteria for aneurysm |
| Internal lumen diameter: |
| >3 mm in children <5 years old, or |
| >4 mm in children >5 years old |
| Of a segment measures ≥1.5 times that of an adjacent segment |
| Clearly irregular coronary lumen |
| Other 6 suggestive features (if ≥3 features, positive) |
| Pervascular brightness of coronary arteries |
| Lack of tapering of coronary arteries |
| Decreased LV function |
| Mitral regurgitation |
| Pericardial effusion |
| Z score in LAD or RCA of 2 to 2.5 |

WBC, white blood cell count in CBC; HPF, high power field; LAD, left anterior descending coronary artery; RCA, right coronary artery; LV, left ventricular.
aneurysms, or ≥3 other suggestive features exist (Table 3B). Several echocardiographic criteria, except the z score of coronary arteries, are qualitative parameters. Recently, quantitation of the perivascular brightness of coronary arteries was performed in children with Kawasaki disease, but its diagnostic validity could not be demonstrated. Decreased myocardial contractility caused by myocarditis, which is histologically universal, is common during the acute phase of Kawasaki disease. Thus, evaluation of myocardial contractility is an important step in the initial diagnosis of Kawasaki disease and was adopted as an echocardiographic criterion. Left ventricular dimensions/volumes, shortening fraction or ejection fraction via the M-mode in parasternal view or via a 2-dimensional examination in apical view are recommended parameters, because they are readily measurable and adequate for routine follow-up.

Left ventricular myocardial deformation was analyzed during the acute phase of Kawasaki disease in 1 study, but the systolic strain rate of the myocardium was not superior to other conventional echocardiographic methods for detecting decreased myocardial contractility. No convincing quantitative threshold values have been proposed for the decision of abnormal pericardial effusion or mitral regurgitation during the acute phase of Kawasaki disease. Newburger et al. acknowledged the diagnostic criteria for incomplete Kawasaki disease prepared by them to be only an informed opinion of a committee of experts (evidence level C). Yellen et al. tested the performance of the diagnostic criteria established by the AHA and reported an applicability of 97%. However, their study subjects were restricted to cases with coronary artery aneurysms. The validity of the diagnostic criteria of incomplete Kawasaki disease established by the AHA should be thoroughly tested in the immediate future because such a systematized guideline is very much needed in pediatric clinical practice.

Additional diagnostically helpful findings are inflammation at the Bacille Calmette-Guérin inoculation site, anterior uveitis, elevated levels of brain natriuretic peptide (BNP) and N-terminal pro-BNP, hyponatremia, elevation of the left ventricular mass, and diastolic dysfunction of the left ventricle. The diagnostic validities of these clinical, laboratory, and echocardiographic findings should also be tested in the future.

Conclusions

Currently, the diagnosis of incomplete Kawasaki disease might be made in cases with fewer classical diagnostic criteria and with several compatible clinical, laboratory, or echocardiographic findings, excluding those of other febrile illnesses. In addition, the validity of the diagnostic criteria of incomplete Kawasaki disease prepared by the AHA should be tested in the future.

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