KAWSASAKI DISEASE IN SULAYMANIYAH/ KURDISTAN/ IRAQ

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ABSTRACT

Background: Kawasaki disease is the most common cause of acquired heart disease in developed countries. Kawasaki disease has been increasingly reported from developing countries. The aim of this study was to assess the clinical and epidemiological characteristics of Kawasaki disease in children of Sulaimaniyah city.

Subject and Methods: In this study all children diagnosed as Kawasaki disease included during the period between January 2010 and December 2014. The hospital records / pediatric cardiology clinic data were reviewed, and data were abstracted onto standardized forms.

Results: During the study period from January 2010 to December 2014, 36 patients were diagnosed as Kawasaki disease. Their mean age at diagnosis was 2.8± 2 years (range 6 months -9years). Most of the patients (30 patients 83.3%) were under age of 5 years, and the median age was 2.1 year. Male to female ratio was 1.76. Most of cases occurred during winter and spring season. Out of 36 patients, 16 (44.4%) patients had coronary disease, and 4 cases of coronary cases were classified as A4 and A5 class according to Japanese Classification of Severity of Coronary Artery Lesions. There was non-significant difference between coronary and non-coronary group regarding hemoglobin level, platelet count, ESR and WBC count.

Conclusions: Kawasaki disease is not uncommon in Sulaymaniyah, the clinical and epidemiological pattern is not different from other parts of the world. We need to increase the index of suspicion in order not to miss Kawasaki disease cases and avoid its serious cardiac complications.


Keywords: Kawasaki disease, Echocardiography, Coronary disease

A Kawasaki disease (KD) is a multisystem, generalized medium and small vessel vasculitis of unknown etiology, which is the most common cause of acquired heart disease among children living in developed countries 1, 2. It is more common in males than in females, with a male-to-female ratio of 1.5:1 3. The distribution of KD by age in childhood is characteristic; the disease occurs most frequently in young children, 50 percent are younger than 2 years of age, 80 percent are younger than 5 years of age, and cases seldom occur in those older than 12 years of age 4-6.

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The principal symptoms and signs include:
1. Fever persisting 5 days or more (inclusive of those cases in whom the fever has subsided before the 5th day in response to therapy).
2. Bilateral conjunctival congestion without exudates.
4. Polymorphous exanthema.
5. Changes of peripheral extremities:
   a. (Acute phase): Redding of palms and soles, indurative edema
   b. (Convalescent phase): Membranous desquamation from fingertips.
   c. Acute non-purulent cervical lymphadenopathy.

At least five items of 1 to 6 should be satisfied for diagnosis of KD. However, patients with four items of the principal symptoms can be diagnosed as KD when coronary aneurysm or dilatation is recognized by two-dimensional (2D) echocardiography or coronary angiography.

Treatment with intravenous immunoglobulin and aspirin reduces the risk of coronary artery abnormalities when administered within 10 days of fever onset. Coronary artery aneurysms, the most serious consequence of KD, seen in 20% to 25% of untreated patients, and long-term consequences include coronary stenosis, early atherosclerosis, and myocardial infarction.

In the current report, the epidemiology and clinical features of 36 patients with KD, seen over a 5-year period at Sulaymaniya Pediatric Teaching Hospital, Kurdistan Region/Iraq is presented.

**PATIENTS AND METHODS**

A retrospective descriptive study was done at Sulaymaniya Pediatric Teaching Hospital that serve more than 1.5 million population in the Sulaymaniya Governorate; Kurdistan Region/ Iraq. The study included all children diagnosed as KD in the period between January 2010 and December 2014. The hospital records and/or pediatric cardiology clinic charts were reviewed, and data were abstracted onto standardized forms. The data included age at onset, sex, the presence and duration of fever, skin and oral manifestations, LN enlargement, laboratory test (lowest hemoglobin, highest platelet count, highest pretreatment ESR, positive CRP), echocardiography features, treatment, outcomes, and follow-up. All cases were diagnosed according to the study group’s diagnostic guidelines for Kawasaki disease base on the criteria of the Japan Kawasaki Disease Research Committee. Patients were included in the study if they had at least five of the following:

1. Fever persisting more than 5 days
2. Changes of extremities
3. Polymorphous exanthema
4. Bilateral conjunctival congestion
5. Changes of lips and oral cavity
6. Acute non-purulent cervical lymphadenopathy

Echocardiography done with Accuson Cypress Siemens 3MH and 7 MH probes, through subcostal, apical, and parasternal views. Size of coronary arteries plotted on normal value chart against body surface area for each patient accordingly (coronary charts and percentiles).
The coronary lesions were classified according to the Japan Kawasaki Disease Research Committee:\(^7\):
A-1 Patients with no dilatation of coronary arteries, A-2 Patients with slight and transient dilatation of coronary arteries, which subsides within 30 days after the onset of KD, A-3 Patients who have small coronary aneurysms at 30 days after the onset of KD, A-4 Patients who have medium coronary aneurysms at 30 days after the onset of KD, and A-5 Patients who have giant coronary aneurysms at 30 days after the onset of KD.

Echocardiographically all were evaluated every 2 weeks apart for the first 8 weeks of the illness. Patients with no evidence of coronary involvement were evaluated again after 6 months while patients with coronary involvement were evaluated every 2-4 weeks until the echocardiography finding become normal. Diagnostic coronary angio planned to perform in 6-12 months later in giant coronary disease cases.

Statistical analysis done using SPSS software program version 19, applying \( t \)-test to compare means and p-value < 0.05 was considered statistically significant.

### RESULTS

A total of 36 patients with Kawasaki disease were evaluated. The mean age at diagnosis was 2.8 ± 2 years, ranging from 6 months to 9 years. Most of the patients were under age of 5 years (30 patients 83.3%), the median age was 2.1 years. Male to female ratio was 1.76, and most of cases occurred during winter and spring seasons, (Table 1).

### Table 1: Demographic and epidemiologic characteristics of the patients

<table>
<thead>
<tr>
<th>Gender</th>
<th>No (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>23(64)</td>
</tr>
<tr>
<td>Female</td>
<td>13(36)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Seasonal distribution</th>
<th>No (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>11 (31)</td>
</tr>
<tr>
<td>Spring</td>
<td>15 (42)</td>
</tr>
<tr>
<td>Summer</td>
<td>8 (22)</td>
</tr>
<tr>
<td>Autumn</td>
<td>2 (5)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Age distribution</th>
<th>No (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 1yr</td>
<td>5 (14)</td>
</tr>
<tr>
<td>1-2yr</td>
<td>7 (19)</td>
</tr>
<tr>
<td>2-3yr</td>
<td>10 (28)</td>
</tr>
<tr>
<td>3-4yr</td>
<td>5 (14)</td>
</tr>
<tr>
<td>4-5yr</td>
<td>3 (8)</td>
</tr>
<tr>
<td>&gt;5 yr</td>
<td>6 (17)</td>
</tr>
</tbody>
</table>

Presenting signs and symptoms are shown in Table 2. Among the 36 patients; 16(44.4%) had coronary affection, 10 were males, (Figure 1).

### Table 2: Signs and symptoms frequencies in Kawasaki disease cases

<table>
<thead>
<tr>
<th>Clinical feature</th>
<th>Number of cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fever</td>
<td>36 (21)</td>
</tr>
<tr>
<td>Conjunctival changes</td>
<td>32 (19)</td>
</tr>
<tr>
<td>Oral cavity and lip changes</td>
<td>28 (17)</td>
</tr>
<tr>
<td>Changes</td>
<td></td>
</tr>
<tr>
<td>Changes in extremities</td>
<td>30 (18)</td>
</tr>
<tr>
<td>Exanthema</td>
<td>24 (14)</td>
</tr>
<tr>
<td>Cervical lymphadenopathy</td>
<td>18 (11)</td>
</tr>
</tbody>
</table>

Figure 1: Sex distribution among coronary affected Kawasaki disease cases.
The investigations done are shown in Table 3. Among the coronary disease, cases only one patient present with right coronary aneurysm, the rest were left coronary artery affection cases.

### Table 3: Duration of Fever and Investigations in coronary and non-coronary Kawasaki disease

<table>
<thead>
<tr>
<th>Variable</th>
<th>Non coronary Group</th>
<th>Coronary Group</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of cases</td>
<td>20</td>
<td>16</td>
<td></td>
</tr>
<tr>
<td>Duration of fever before starting treatment</td>
<td>8.4+/-.2.6</td>
<td>6.6+/-.2.35</td>
<td>0.004</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>10.5+/-.1.36</td>
<td>11.1+/-.1.3</td>
<td>0.325</td>
</tr>
<tr>
<td>Platelet</td>
<td>487+/-.193</td>
<td>524+/-.96</td>
<td>0.527</td>
</tr>
<tr>
<td>ESR</td>
<td>75+/-.21</td>
<td>81+/-.29</td>
<td>0.497</td>
</tr>
<tr>
<td>WBC count</td>
<td>14457+/-6127</td>
<td>17225+/-7566</td>
<td>0.240</td>
</tr>
</tbody>
</table>

The severity of coronary artery affection shown in Table 4

### Table 4: Classification of Severity of Cor. Artery Lesions Based on Echocardiographic Findings

<table>
<thead>
<tr>
<th>Coronary disease class</th>
<th>No. of patients (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A 1</td>
<td>2 (13)</td>
</tr>
<tr>
<td>A 2</td>
<td>6 (36)</td>
</tr>
<tr>
<td>A 3</td>
<td>4 (25)</td>
</tr>
<tr>
<td>A 4</td>
<td>2 (13)</td>
</tr>
<tr>
<td>A 5</td>
<td>2 (13)</td>
</tr>
</tbody>
</table>

All patients received IVIG single dose 2gm/kg and Acetylsalicylic acid anti-inflammatory dose. No recurrence noted in our cases but in one case the fever did not subside after the recommended IVIG dose in which case the coronary artery disease was diagnosed on the 2nd echocardiographic examination on 18th day of illness although 1st echo was normal on 8th day of illness. Another patient presented with giant aneurysm (8mm) after 14 days of fever in which the intravenous immunoglobulin treatment was not received within the 10 days of illness, on subsequent echo, the measurement increased to 11mm at left main stem. The third patient developed giant aneurysm from the 2nd weeks and persisted as the same measure till 1 year follow up and then regressed to 7mm on the end of the 2nd year. Subsequently we did coronary catheterization which supported the same diagnosis without other aneurysmal or stenotic lesions.

### DISCUSSION

Most of KD cases (30, 83.4%) were below 5 years of age, this finding was similar to studies done in western Saudi Arabia (83.3%), in Korea (71%) in Iran (77%) and in Taiwan (80%). In addition the average age at diagnosis in our study (2.9 years) was nearly similar to other studies in Korea and Iran (2.8 years in both). Male to female ratio was (1.76:1), which is similar to Saudi Arabia study (1.7:1) and Iranian study (1.5:1). In the current study there was seasonal variation, in which we have two cases recorded during autumn and the peak of our cases was reported in winter, spring and early summer, similar to other studies done in Saudi Arabia, Iran and United States of America. However, in other studies KD had a peak incidence during other seasons.
The high rate of white blood cells, platelet count, ESR and positive C reactive protein was similar to other reports\(^\text{17,19-22}\). There was no significant difference between the presentations of patients with coronary involvement and patients with no coronary involvement except for the duration of fever before commencing treatment. Similar results were found in studies in Iran\(^\text{17}\), and China\(^\text{23}\). In one of our patients, the fever continue in spite of using aspirin and IVIG without explanation, and later she was diagnosed as rheumatoid arthritis.

Delay in diagnosis and treatment, which occur more frequently in older children, are associated with an increased risk of coronary artery aneurysms as we have one case with giant aneurysm diagnosed after 14th day of illness\(^\text{24}\).

If a child clearly meets criteria for Kawasaki disease, the decision to treat with intravenous immunoglobulin is not difficult, especially if surrogate markers support the diagnosis, if the coronary artery abnormalities were detected by echocardiography, continue the aspirin with a dose of 3-5 mg/kg PO qd long term\(^\text{28}\).

Patients with severe coronary affection categories (A4 and A5), like the four patients we have, are at high risk for coronary sequel in the future and if coronary artery abnormalities fail to regress by time, long-term pharmacological therapy and diagnostic follow-up are implicated\(^\text{29}\).

In conclusion, KD is not uncommon in Sulaymaniyah, it seems that the clinical feature is not different everywhere in the world. We need from our pediatrician to be aware of it and not to miss Kawasaki disease cases to avoid its serious cardiac complication.

**REFERENCES**


8- Melish ME, Marchette MJ, Kaplan JC, Kihara S, Ching D, Ho DD. Absence of significant RNA-dependent DNA polymerase activity in lymphocytes


نَهْخُوشی کاواساکی لە سلیمانیه / کوردستان / عراق

تَحَمیل

یَشَکِی: نَهْخُوشی کاواساکی باوْترین نَهْخُوشی وَ وَگنْرَگا دَه لە وَلَائَت بِئتشکووْتکاندَا. مَرَْدَهَا نَهْخُوشی کاواساکی لَوَلائَت گَاشْ-کردووْتکاندَا پَارْهِى سَرَنَدْوَهٔ. نَانَمَّ َهَم تَوْژیزَى وَ وَژَوْیهَا كَه نَاسَرْ-مَوْکَان وَ نَیْشانْ-مَکَان وَمَه نَهْخُوشی وَه دَیْارَی بَکریت لَه مَنْدالَا لَه شَاری سلیمانی بِلْتَانٌ-بیتَیمَهشَی وَوانٌی-کَسْهشَی دَیْیان دَییتَه.

رَیْکَین ظَفلَکُولِئینی: نَم تَوْژیزَیْوْه نَنْجَمَم دَرَاه لُحَسَر نَهْخُوشی کاواساکی لَوَلائَت ٢٠٠٠-٢٠١٠ لَه بِشَی دَلیً مَنْدالَا لَوَ نَهْخُوشخانَه، مَنْدالاتَی فَیْرکاری شَاری سلیمانی هَمْوَو حَالتَهکان بِهِی، نَاسَرْ-مَوْکَان لِیزَهی نَهْخُوشی کاواساکی زَابْوْیی.

نَفَْتَجَام: زَمرائَه ٣٦ نَهْخُوش لَوَل مَوْبیا بَه کاواساکی تَوْار دَیاری کَراوه کَخاوَندَ تَعَمَهی١٠ غلیسا (٦ مَانگ-٩ سَلَ)، وَوَوَزَیهَا نَهْخُوشکان (٣٠ نَهْخُوش٣٨٪) لَه زَیر تعَمَهی ٥ سَلًا مَوْنَ، رَیْهَی کَر کَرْج١٧٦، وَوَوَزَیهَا حَالتَهکان لَه وَرَزَی زَستَان وَ بِعَمار روَوَوی دَاَوْه. لَه کَری گَنَنی١٦ نَهْخُوش٤٤٪(نهْخُوشی بَوریکتی مَاسْتَکَهی) جَلیان گَمْره بَو، كَه نَهْخُوش لَوَلائَه لَه جَرَویی (٤٨٪) بِولیین نَهْخُوشی کاواساکی جَلی زَابْوْیی دَائَترَون. هَمروُهَا جَیاوازی نِییه لَه رَیْهَی خَستَی خَوْنَ وَ خَرؤْوی سَبیکَهَا وَ خَطای اَیکَی خَوْنَ لَه هَمَرَوو وَو َرَویهَا.

دَرَنْجَاَم: لَه دَرْنَجَامی لیکَلیئَنیمَهه نَوَوْه سَاَغ بَوَوْه كَه نَهْخُوشی کاواساکی نَاسَرْو لَوَلائَتی سَلیمانیا وَ جَیاوازی نِییه لَه وَلائَتی ثَر. هَمروُهَا پَیوْشَیم بَر چَواروووْنیمَهه کَر یَشکَی خَوْنَمَهه لَه کُوردُستَان بَو نَاسَرْبَارو دَیاری کَرِدنی نَم نَهْخُوشی وَه لَه سَرَفْتَانَییا.
الخلاصة

مرض الكوواساكي في السليمانية / كردستان العراق

الخلفية والأهداف: مرض الكوواساكي من الأمراض الشائعة في الدول المتقدمة وفي وقتنا الحاضر المرض أصبح شائعاً في الدول النامية. الغرض من الدراسة متابعة مرضى الكوواساكي في السليمانية و متابعة الاعراض المتلازمة له.

طرق البحث: الدراسة اجريت على الأطفال الذين يشكون من المرض الكوواساكي من فترات 2020 إلى 2014 في قسم قلب الأطفال في مستشفى الأطفال التعليمي في السليمانية، المشخصة حسب الاعراض المتلازمة من مجلس الكوواساكي اليابانية.

النتائج: ش kháchت 36 حالة خلال مدة الدراسة كحالة كواكساكى كاملة ، معدل العمر كان 0±3.0 سنة (3 أشـهر - 9 سنة) ، أكثر من المرضى (82.5%) كانوا دون عمر 5 سنوات . نسبة التذوکر الى الأثنين 1.76 ، و أكثر الحالات شخصت في الشتاء و الريـع . شخصت 16 حالة (44.4%) كتوسع شرايين القلب والقلب و اربعة حالات صنفت بتصنـيف (A4,A5) حسب الاعراض المتلازمة من مجلس الكوواساكي اليابانية للقلب. واستنتجنا أيضا أنه ليس هناك اختلاف في نسبة الدم ، كريات البيضاءو صفائح الدم بين المجموعتين.

الاستنتاجات: استنتجت من الدراسة أن في وقت الحاضر مرض الكوواساكي منتشر في السليمانية و ليس هناك اختلاف في انتشار و اعراض المرض مقارنة بمناطق العالم، و استنتجنا أيضا اننا نحتاج الى توعية الأطباء الأطفال حول اعراض و مضاعفات مرض الكوواساكي لغرض تشخيصه في المراحل الأولى.