Acute rheumatic fever: 27 year experience from the Montreal’s Pediatric Tertiary Care Centers

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Objectives
To examine the epidemiology, clinical characteristics and outcomes in a cohort of children with acute rheumatic fever (RF) over the past 27 years in Montreal.

Methods
The medical records of patients younger than 18 years of age hospitalized and diagnosed with RF in Montreal between January 1979 and December 2005 were reviewed.

Results
Among the initial 134 charts selected, 36 children were already followed-up for chronic RF and the remaining 98 patients (51 % females) who fulfilled the Jones criteria for acute RF were included in the analysis. The mean age at diagnosis was 10.1 ± 3.0 years (range: 3-17). Over the 27-year study period, there was a mean incidence of 3.6 patients/year without peaks, but onset occurred in the last 15 years in almost two-thirds of the patients.

Forty-nine percent of the patients were Canadian-born non-aboriginal (CbnA) and the remaining patients were Canadian-born aboriginal (CbnA) or foreign-born (Fb). Carditis was diagnosed in 73 % of the patients and Sydenham’s chorea in 49 %. Of the CbnA children, 59 % had carditis compared with 61 % of children from other ethnic groups (p = 0.003). However, the form of presentation was chorea in 69 % of CbnA children vs. 31 % of children from other ethnic groups (p < 0.001). No deaths were attributable to acute RF although 2 % of the patients relapsed during the study period. Severe cardiac sequelae requiring valve replacements occurred in 6.1 %.

Conclusion
The incidence of acute RF in Montreal was low but consistent over the 27 year study period. Clinical presentation varied depending on ethnicity.

Key words: Acute rheumatic fever. Children. Carditis. Chorea.

FIEBRÉ REUMÁTICA AGUDA: 27 AÑOS DE EXPERIENCIA EN LOS HOSPITALES PEDIÁTRICOS EN MONTREAL

Objetivos
Revisar la epidemiología, las características clínicas y la evolución de una cohorte de niños, durante 27 años, con fiebre reumática aguda en Montreal (Canadá).

Métodos
Las historias clínicas de niños menores de 18 años hospitalizados y diagnosticados de fiebre reumática en Montreal, entre enero de 1979 y diciembre de 2005, fueron revisadas.

Resultados
De las 134 historias seleccionadas, 36 niños eran seguidos por fiebre reumática crónica, los 98 pacientes restantes (51 % de sexo femenino) cumplían los criterios diagnósticos de fiebre reumática aguda y fueron incluidos en el análisis. La edad media en el momento del diagnóstico era de 10,1 ± 3,0 años (intervalo: 3-17). Durante los 27 años del periodo de estudio, hubo una incidencia media de 3,6 pacientes por año sin picos de incidencia, sin embargo dos tercios de los casos se presentaron en los últimos 15 años. El 49 % de los pacientes había nacido en Canadá y no nativos (NCnN), el resto eran nativos nacidos en Canadá (NCN) o nacidos en el extranjero (NE). La carditis fue diagnosticada en el 73 % de los pacientes y la corea de Sydenham en el 49 %. De los niños NCnN, el 59 % presentaban carditis comparativamente al 61 % de los niños de otro origen (p = 0.003). Sin embargo, la corea fue el modo de presentación en el 69 % de los niños NCnN frente al...
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31% en niños de otro origen (p < 0.001). No hubo ninguna mortalidad, a pesar de que el 2% presentaron recidiva durante el periodo de estudio y el 6,1% con secuela cardíaca severa requirieron recambios valvulares.

Conclusion
La fiebre reumática aguda en Montreal, a pesar de ser una enfermedad rara, ha presentado una incidencia constante en los últimos 27 años, la presentación clínica depende del origen étnico.

Palabras clave:

INTRODUCTION
Acute Rheumatic Fever (ARF), a non-suppurative sequela of group A β-hemolytic streptococcal infection (GAS) has become a rare pathology in developed countries. However, every year, pediatric services from large urban centres continue to diagnose patients with ARF. Moreover, it is difficult to establish the diagnosis, as many young physicians have never seen a patient with ARF during their training. Transcontinental migration and changing virulence of certain strains of GAS may influence the epidemiology of ARF.

Regular reassessment of the epidemiology is essential in order to recognize and to diagnose promptly this disease. Recent published literature contains only few studies with large number of pediatric patients. There is the feeling that ARF is more of an issue of the past, however we have found a consistent number of new cases each year over the past 15 years. This paper is meant to highlight the epidemiology and clinical characteristics of the cohort of children diagnosed with ARF in a modern North American/Canadian city.

MATERIAL AND METHODS
Study population and data collection
A retrospective chart review of all children diagnosed between 1979 and 2005 with ARF in the pediatric tertiary care referral centres. Montreal (Canada) has two Pediatric hospitals: Sainte-Justine Hospital (a 452 bed pediatric hospital with 25,000 admissions and 300,000 total visits per year, 80,000 emergency room visits) and The Montreal Children’s Hospital (MCH) a 214 bed pediatric hospital with 7,000 admissions and 120,000 total visits per year, 65,000 emergency room visits). Hospitalized children younger than 18 years of age diagnosed with ARF from January 3rd 1979 through December 31st 2005 were included. Patient’s charts were identified through epidemiologic follow-up, hospital database and every year over the past 15 years. This paper is meant to highlight the epidemiology and clinical characteristics of the cohort of children diagnosed with ARF in a modern North American/Canadian city.

INCLUSION CRITERIA
Any patient that fulfilled the Jones Criteria for ARF were retained. Carditis was considered with any new pathologic cardiac murmur, cardiomegaly, pericarditis or congestive heart failure or typical valve involvement at echocardiography. Evidence of recent streptococcal infection was defined by Antistreptolysin-O antibodies (ASO) titres of 400 Todd Units or greater and Anti-deoxyribonuclease B (DNase) B titres of 1/500 or greater.

EXCLUSION CRITERIA
Chorea explained by other causes, uncompleted Jones Criteria and patients with relapses episodes from remote rheumatic fever. Statistics on population in Montreal were obtained from Agence de la Santé et des Services Sociaux de Montréal.

STATISTICAL ANALYSIS
Data from the two hospitals were compiled and combined in the analysis using standard spreadsheet software (Microsoft Excel 2000® and SPSS 11.0 for Windows). Descriptive statistics were calculated on the entire cohort except when mentioned otherwise. Logistic regression was done to establish whether certain covariates were associated with the different manifestations of ARF. Maximum likelihood estimates of regression coefficients were used to estimate odds ratios. Ninety-five percent confidence intervals were calculated for all estimates reported.

RESULTS
Demographic characteristics
During the 27 years study period, 134 charts were selected, of which 86 were excluded because of a past history of ARF. The remaining 48 patients fulfilled the Jones criteria of ARF and were included in the analysis: 47 children from Sainte-Justine Hospital, and 51 from The Montreal Children’s Hospital. Overall, median age was 10.1 ± 3.0 years (range: 3 to 17), 51% were female. An average of 5.6 patients/year (range: 0 to 9) was diagnosed; 67.5% of patients presented in the last 15 years. Forty-nine percent of patients were Canadian-born non-Aboriginal (CbnA: N = 48) and the remaining were either

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Canadian-born Aboriginal (CbA: N = 11; 11.2 %) or Foreign-born (Fb: N = 39; 39.8 %). The median rate of ARF in Montréal was calculated at 1.2/100,000, for the total length of observation. CbnA 0.8/100,000, CbA 10.7/100,000 and Fb 2.1/100,000. A positive family history of ARF was known in only three patients.

Clinical presentation
Carditis was the most frequent clinical manifestation of ARF (72.4 %) followed by Sydenham chorea (49.0 %) (table 1). In our cohort, 51.6 % patients (n = 47) had simultaneously a positive throat culture and elevated ASO at the time of presentation.

CbA or Fb children had more than four-fold increased risk of carditis compared to CbnA children, Crude OR (95 %IC) 4.4 [1.6-11.7] p = 0.003 (fig. 1). CbnA children presented more than five-fold increased risk of chorea compared to children from other cultural backgrounds, Crude OR 95 %IC 5.1 [2.2-12.1] p < 0.001 (fig. 2).

Among 71 patients with carditis diagnosed clinically or by echocardiography, 16 patients (16.5 %) had congestive heart failure and 31 children had involvement of more than one valve. The association of mitral and aortic regurgitation was found in 40.8 % of patients (n = 29), isolated mitral valve or aortic regurgitation were found respectively in 46.5 % and 9.9 % patients (n = 33 and 7), mitral stenosis in 2.8 % (n = 2) and aortic stenosis in 1.0 % (n = 1). Five patients presented with pericarditis and in three children, cardiac nodules were described at the echocardiography.

Forty-eight out of 98 children (49 %) presented with the manifestation of chorea. It was seen significantly more frequently in females (32/48 = 66.7 %) than in males: Crude OR (95 %IC) = 3.6 [1.5-8.2], p = 0.003. Isolated chorea occurred in 29.2 % patients (n = 14), and was unilateral in 43.8 % children (n = 21). Imaging abnormalities at computed tomography or MRI of the brain were found in 44.8 % patients (13 over 29). The mean duration of chorea was 4 months (range: 1-24). Eight patients were treated with Haloperidol, 13 with Valproic Acid. 12 received other drugs and 15 received no treatment. Three patients suffered a prolonged dysarthria with complete resolution after speech therapy.

Isolated polyarthritis, as the only major criteria was observed in 16.3 % patients (n = 16). Erythema marginatum and subcutaneous nodules were found in 23.5 % and 3.1 % of children, respectively. Both carditis and polyarthritis were found in 28.6 % patients (n = 28); carditis and chorea in 30.6 % patients (n = 30), polyarthritis and chorea were presented in two patients (2.0 %) and the three major criteria of polyarthritis, carditis and chorea in two others patients (2.0 %).

Table 1. Clinical features, laboratory and outcome of 98 patients with ARF

<table>
<thead>
<tr>
<th>N</th>
<th>Frequency (%)</th>
<th>N</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carditis</td>
<td>71</td>
<td>72.4</td>
<td></td>
</tr>
<tr>
<td>Chorea</td>
<td>48</td>
<td>49.0</td>
<td></td>
</tr>
<tr>
<td>Polyarthritis</td>
<td>37</td>
<td>37.8</td>
<td></td>
</tr>
<tr>
<td>Erythema marginatum</td>
<td>23</td>
<td>23.5</td>
<td></td>
</tr>
<tr>
<td>Subcutaneous nodules</td>
<td>3</td>
<td>3.1</td>
<td></td>
</tr>
<tr>
<td>Fever</td>
<td>41</td>
<td>41.8</td>
<td></td>
</tr>
<tr>
<td>Arthralgias</td>
<td>52</td>
<td>52.7</td>
<td></td>
</tr>
<tr>
<td>Increased erythrocyte sedimentation rate</td>
<td>56/87</td>
<td>66.7</td>
<td></td>
</tr>
<tr>
<td>Increased PR segment</td>
<td>22/78</td>
<td>28.2</td>
<td></td>
</tr>
<tr>
<td>Positive throat culture</td>
<td>57/91</td>
<td>62.6</td>
<td></td>
</tr>
<tr>
<td>Elevated antistreptolysin-O titer</td>
<td>76/90</td>
<td>84.4</td>
<td></td>
</tr>
<tr>
<td>Elevated anti-DNAse B titer</td>
<td>51/42</td>
<td>73.8</td>
<td></td>
</tr>
<tr>
<td>Relapses</td>
<td>2</td>
<td>2.0</td>
<td></td>
</tr>
<tr>
<td>Valve replacement</td>
<td>6</td>
<td>6.1</td>
<td></td>
</tr>
<tr>
<td>Death</td>
<td>0</td>
<td>0.0</td>
<td></td>
</tr>
</tbody>
</table>

ARF: acute rheumatic fever.

Figure 1. Distribution of cardiac manifestations (N = 71) according to the ethnic origin.

Figure 2. Distribution of chorea (N = 48) according to the ethnic origin.
By the time this review was done: 2 patients suffered clinical relapses; 6 patients (6.1%) required valve replacements, 4 of them in the five years following the initial diagnosis; no deaths were attributed to ARF.

**DISCUSSION**

Despite a high prevalence in the third world, ARF has become a rare disease in industrialized countries, with the overall incidence which has decreased significantly to less than 1/100,000 around the world. During the last decades, ARF has also been on the decline in Canada, the Toronto ARF registry reported in 1961 an incidence of 210/100,000, and in the late 1980s of 0.96/100,000. Our recent experience shows an overall median rate of ARF in Montreal of 1.2/100,000.

Certain ethnic groups may be more susceptible to ARF than others. In 1982, longitudinal studies in Canada, reported an incidence rate of 126/100,000 in native and 29/100,000 in non-native children. In our series in Montreal, we found an incidence for CbA of 0.8/100,000 compared to 10.7/100,000 for Cha and 2.1/100,000 for Fb. The reasons for this observation have not yet been established, since data available were not sufficient to conclude that overcrowding or any other specific factor was responsible. Although, we believe that social and economic factors and in particular housing conditions, are the major reasons for the high rates found in these specific populations. Given the relatively small native population in Quebec (n = 8814 children over total population of 299,150 of children 5-19 years old (statistics from 2001)), it appears that this ethnic group is at higher risk. This may be due to high crude colonization rate of Streptococcus among Inuit, genetic predisposition, poor socio-economic conditions, overcrowding and diminished access to primary health care.

Specific HLA antigens and different allantigen expressed at the surface of lymphocytes and recognized by monoclonal antibodies appear to be a markers of susceptibility to RF and expressed only after stimulation by specific rheumatogenic strains of group A Streptococcus. The high prevalence of Streptococci among Inuit children has been published previously. In our series, information according to family income or persons/household was not available. The decreased incidence of ARF in the world, has been attributed to improved living conditions and medical care, the introduction of antibiotics and the changes in prevalence of rheumatogenic group A Streptococcus.

During the study period, we have seen an average of 3-5 new cases of ARF/year without significant outbreaks such as reported in the United States or in other provinces of Canada. Instead, we calculated 2.7 cases/year in the period 1979-1984, 2.8 cases/year in 1985-1994 and 4.9 cases/year in 1995-2005. It is even possible that the real incidence of ARF in our environment might be higher than the rate observed because we have only included hospitalized patients with diagnosis of ARF, thus the incidence of ARF may have been underestimated if patients were managed without hospitalization. The likelihood of children hospitalized in other secondary care centers surrounding Montreal is not likely, as they reported having not seen patients with ARF for a very long time (personal communication).

Carditis was diagnosed in 72.4% of our patients, a similar incidence was found in US in the 1980s. As known from the literature, carditis is diagnosed in 50% of cases on clinical examination and 70% by cardiac sonography. In our cohort, carditis was statistically significant more commonly found among Cha or Fb than in CbA. In industrialized countries, long-term prognosis of carditis has improved because the disease itself seems to have a modified course and also because antibiotic prophylaxis has prevented subsequent attacks. We outline the good prognosis of our cohort with only two cases of relapse and only 6.1% of patients requiring valve replacements, both statistics comparing favourably to other series. This could have been related to a good compliance to prophylaxis combined with a decreased exposure to other rheumatogenic group A Streptococcal strains.

In our series, chorea was present in 49.0%. In the 1990s, Allen had published an incidence of chorea associated with ARF of only 6% in Toronto, Canada. Our incidence of chorea was also higher than the 6-31% reported in the literature. Patients who present with Sydenham’s chorea often lack evidence for Group A streptococcal infection. Nevertheless, in our patients with chorea, 60% had a positive throat culture and 52% had an elevated ASO titer. During our study, we did not include any case of PANDAS (The Pediatric Autoimmune Neuropsychiatric Disorders Associated with Streptococcal Infection). We found that polyarthritis was present in 57.8% of patients, less frequently than the 50-80% reported in the literature. Isolated polyarthritis, as a single major criterion of ARF, associated with fever and increased acute-phase reactants is the weakest diagnostic criteria of ARF and a problem clinician’s may encounter in practice; in our cohort, it was accepted in 16 patients.

Documentation of an antecedent streptococcal infection is required for diagnosis of ARF. In our cohort,
62.6% of patients tested at the time of diagnosis had positive throat culture for Group A streptococcus. These results have limited usefulness, because it neither distinguishes between recent infection and a more long standing pharyngeal carriage. Nevertheless, this result was confirmed in our cohort, as we found elevated ASO and anti-DNase B titer in 84.4% and 73.8%, respectively. Unfortunately the serotypes of the streptococci were unavailable in our review.

Although our study involves a large series of ARF in children from Montreal, its retrospective nature and the evaluation of only hospitalized cases in tertiary health centers constitute limitation factors. Rheumatic fever may have different clinical presentations in developing countries according to genetic predisposition, prevalence of rheumatogenic strains and socio economical conditions. The results of the present study reflect the clinical picture of an industrialized North American city.

In conclusion, although, it is important to emphasize that ARF has become a rare disease in industrialized countries, it remains prevalent in our environment and clinical presentation varied depending on the ethnicity. The most striking fact in our study was that chorea was significantly more frequently observed in Canadian born-non aboriginal children. Further studies are needed to identify the virulent factors and epitopes of rheumatogenic streptococcal strains as well as the genetic markers of predisposition to explain the differences in this clinical presentation. Physicians ought to know this pathology, in order to recognize and diagnose correctly children with episodes of ARF, a disease which is still with us.

REFERENCES