Clinical Characteristics of Infective Endocarditis in Children

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Background: Infective endocarditis (IE) remains a diagnostic and therapeutic challenge associated with high morbidity and mortality. We evaluated the microbial profile and clinical manifestation of IE in children.

Methods: A retrospective study examining pediatric IE cases treated between 2000 and 2017 at the Department of Pediatric Cardiology, KU Leuven, was conducted. Clinical presentation, treatment, complications, outcome of IE, underlying microorganisms and congenital heart defects were reviewed.

Results: Fifty-three patients were diagnosed with IE. Overall, 19 patients (36%) required cardiac surgery. Seven patients (13%) died. Eighty-seven (19%) cases of IE were detected in this period, which is a substantial decrease compared to the previous period (142 cases). The most frequent causative organism was Staphylococcus aureus (34%), followed by viridans streptococci (17), Staphylococcus epidermidis (15), and coagulase-negative staphylococci (20). Community-acquired (CA) IE increased significantly from 8 (33%) cases in 2000–2007 to 20 (74%) cases in 2008–2017 (P < 0.01). Even with viridans streptococci being significantly more prevalent in the CA group (P < 0.01), we did not observe an increase of streptococcal IE from 2008 to 2017. Seventeen (32%) patients presented with hospital-acquired IE during the first year of life with 14 (82%) children after surgery and a prevalence of coagulase-negative staphylococci (53%).

Conclusions: The incidence of pediatric IE was similar over the investigated time period with a shift toward CA IE. Staphylococcus and staphylococci accounted for the majority of cases in both periods. Awareness of IE and its prevention is crucial in patients after implantation of prosthetic grafts.

Key Words: congenital heart disease, infective endocarditis, artificial valve conduits


Patients with congenital and acquired heart valve disease are at higher risk to develop infective endocarditis (IE). In clinical practice, IE has proven itself a challenging diagnosis with high morbidity and mortality. Novel therapeutic strategies in congenital heart disease (CHD), including catheter- and surgery-derived interventions, have improved patients quality of life but treatment-associated complications, such as IE, are still of major concern. Furthermore, advances in life-saving medical interventions such as critical care and immunosuppressive therapies have increased the population at risk for IE and more frequent use of implanted prosthetic material leads to a higher incidence of device-related infections. In children, patients with underlying CHD are the most important population at risk for IE and because of advanced treatment age at IE diagnoses increases.

Changes in IE epidemiology have become of major interest after the recommended IE prophylaxis has been restricted to patients at high risk from 2007 onwards. Streptococci are still described as the most relevant bacterial pathogen responsible for IE whereas Staphylococcus aureus infections are becoming more frequent.

We conducted a retrospective investigation of pediatric patients with IE treated between 2000 and 2017 at our institution. By evaluating epidemiologic and clinical data, we aimed to improve our understanding of the risk of IE associated with specific CHD, the course of disease and its potential prevention.

PATIENTS AND METHODS

Patients

This retrospective study was conducted using the patient database of the Department of Pediatric Cardiology, University Hospital Leuven, a tertiary referral hospital. All children under the age of 16 years who were diagnosed with definite IE following the modified Duke Criteria between January 1, 2000, and December 31, 2017, were included in the study. Approval by the local ethics committee was asked and informed consent was taken from the parents.

Clinical Data

Patients’ date of birth, gender, underlying CHD and surgical and interventional treatments were noted. Recorded data of the IE episode included the date of IE diagnosis, patient's history, clinical presentation, echocardiographic findings, biochemical and microbiologic data, medical and surgical treatment, length of hospitalization, complications and outcome. Organisms were considered causal if at least 2 blood cultures or a single culture of intraoperative specimens were positive. If no organism met these criteria, the IE episode was considered culture negative. Community-acquired (CA) IE was defined as episodes occurring more than 6 months after cardiac surgery in nonhospitalized patients or within the first 72 hours of hospitalization.

Statistical Analysis

Continuous variables are reported as mean ± standard deviation. Categorical variables are mentioned as frequencies and percentages of the specific group. Statistical analysis was performed by application of the Fisher and χ2 test using GraphPad Prism (7.0d; GraphPad Software, San Diego, CA).

RESULTS

Population Characteristics

Analysis of patient records identified 53 children diagnosed with definite IE according to the Modified Duke Criteria under the
age of 16 between January 1, 2000, and December 31, 2017. None of the patients had recurrent episodes of IE.

Our patient population consisted of 34 (64%) boys and 19 (36%) girls with a mean age at diagnosis of 6.5 years. We observed 17 infants under the age of 1 year with 11 of them diagnosed between 2000 and 2007. Overall, 24 patients were diagnosed between 2000 and 2007 while 29 patients were diagnosed from 2008 to 2017. In the entire period, 7 patients (13%) died from IE and 19 patients (36%) required cardiac surgery in the management of IE. All patients were treated for 6 weeks with the respective recommended antibiotic treatment according to the guidelines of the European Society of Cardiology (ESC).21

Characteristics of IE patients are shown in Table 1 and patients’ age distribution is given in Figure 1.

Underlying CHD

Forty-six of the 53 patients (87%) had underlying CHD which included mainly Tetralogy of Fallot (TOF) in 13 patients, a perimembranous ventricular septal defect in 6 patients and a truncus arteriosus in 4 patients. None of the patients had an underlying muscular ventricular septal defect or atrial septal defect and none rheumatic disease. At the time of IE diagnoses, 8 (17%) patients with CHD had not yet undergone surgical repair for their lesion. Detailed diagnoses of CHD are shown in Table 2. In our study population, 7 (13%) children had a structural normal heart with 2 of them having undergone immunosuppressive treatment.

Microbiology

A causative organism was found in 49 cases (92%). In 6 patients, 2 different causative organisms were withheld and included in the analysis. The most frequent causative organisms in the total study period from 2000 to 2017 were viridans group streptococci (17 episodes, 29%), S. aureus (13 episodes, 22%) and coagulase-negative staphylococci (11 episodes, 19%). Other organisms included Candida, enterococci, members of the HACEK group and Abiotrophia species, as well as Mycobacterium avium and Serratia marcescens. Patients were divided into 2 groups depending on the time point of IE diagnosis: 2000–2007 for group 1 and 2008–2017 for group 2. There was no significant difference in the prevalence of any underlying organism, also not for viridans group streptococci, comparing the 2 time periods (P > 0.05). Figure 2 demonstrates the relative importance of the underlying organisms in both groups.

Using the provided criteria, 28 of all cases were classified as CA IE while 25 cases were classified as hospital-acquired (HA) IE. Interestingly, CA IE cases were significantly more frequent between 2008 and 2017 compared with the earlier period (P < 0.01, Table 1 and Fig. 3A).

In general, viridans group streptococci are the major underlying cause for CA IE because they were found in 14 (48%) of CA IE cases while only present in 3 (10%) of HA IE cases (P < 0.01). Conversely, coagulase-negative staphylococci were present in 2 (7%) of CA IE cases and 9 (31%) of HA IE cases (P < 0.05). S. aureus accounts as the second important agent in CA IE 8 (27%) cases compared with 5 (17%) in HA IE. None of the patients was positive for methicillin-resistant S. aureus (MRSA).

Despite increasing CA IE in the later time period, the incidence of viridans streptococcal IE did not increase in CA IE between 2008 and 2017 (P > 0.05). M. avium and S. marcescens caused HA IE.

TABLE 1. Characteristics of IE Patients

<table>
<thead>
<tr>
<th>Definite IE (n = 53)</th>
<th>Number of Patients (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender (female/male)</td>
<td>19 (36%/34/64%)</td>
</tr>
<tr>
<td>Mean age at IE</td>
<td>6.5 ± 5.7 yr</td>
</tr>
<tr>
<td>Surgery for IE</td>
<td>19 (36%)</td>
</tr>
<tr>
<td>Death because of IE</td>
<td>7 (13%)</td>
</tr>
<tr>
<td>Underlying CHD</td>
<td>46 (87%)</td>
</tr>
<tr>
<td>IE between 2000 and 2007/CA</td>
<td>24/8</td>
</tr>
<tr>
<td>IE between 2008 and 2017/CA</td>
<td>29/20</td>
</tr>
<tr>
<td>Right-sided IE</td>
<td>33</td>
</tr>
<tr>
<td>Left-sided IE</td>
<td>15</td>
</tr>
<tr>
<td>Bilateral IE</td>
<td>5</td>
</tr>
</tbody>
</table>

TABLE 2. Underlying Congenital Heart Disease and IE Localization

<table>
<thead>
<tr>
<th>CHD</th>
<th>n = 46</th>
<th>Presurgery (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetralogy of Fallot</td>
<td>13</td>
<td></td>
</tr>
<tr>
<td>Truncus arteriosus</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>pm VSD + coarctation/ TS/ MI/ MS/ AI</td>
<td>5/3</td>
<td></td>
</tr>
<tr>
<td>pm VSD</td>
<td>1/1</td>
<td></td>
</tr>
<tr>
<td>Atrioventricular septal defect</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Univentricular heart</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>AS/ AI</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Complex TGA</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>TGA</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Shone complex/ MI</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Persistent ductus arteriosus</td>
<td>1/1</td>
<td></td>
</tr>
<tr>
<td>Bicuspid aortic valve</td>
<td>2/2</td>
<td></td>
</tr>
<tr>
<td>Ebstein malformation</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

Infective endocarditis distribution with age. A similar distribution of age was found, besides a subpopulation with 17 children <1 year of age.
Infective Endocarditis in Children

IE in Children Under 1 Year of Age

Age distribution among our patients is spread in a similar manner, besides a subpopulation of 17 patients (32%) in which IE was diagnosed during the first year of life. All patients, even younger than 5 months, presented with HA IE. Sixteen of them had underlying CHD and in 14 patients IE was diagnosed 30 days (±42 days) after cardiac surgery. Looking at the causative organisms within this subgroup revealed that coagulase-negative staphylococci were much more prevalent (53% against 9% at older age; *P < 0.05) while viridans group streptococci were not present. In addition, Staphylococcus aureus was found in 3 children and all 4 children with Candida IE belonged to the subgroup of postoperative infants.

Lesion Location

Analyzing cardiac ultrasound records, IE was classified as right-sided IE in 33 (62%) patients and left-sided IE in 15 (28%) patients. Five (10%) patients presented with bilateral lesions (Table 2). Right heart IE was equally caused by the main 3 bacteria (S. aureus 10 episodes, coagulase-negative staphylococci 9 episodes and viridans group streptococci 12 episodes). There was also no prominent causal agent in left or bilateral heart IE. However, culture-negative IE was observed in both left and right heart IE.

All 18 patients with underlying TOF or truncus arteriosus had undergone previous surgery and presented with right-sided IE lesions which were associated with left-sided lesions in 3 patients. The majority of this subgroup (15 patients, 83%) presented with infection of prosthetic material. The primary IE site was determined to be a pulmonary homograft in 6 patients, a stent mounted bovine jugular vein (BJV) graft in 4 patients, a BJV graft in 3 and a bovine pericardial patch (right ventricular outflow tract) in 2 children. The remaining 3 patients (all TOF) had native tricuspid valve IE and 1 patient had both native mitral and pulmonary valve involvement.

In total, 18 of the 53 patients (34%) presented with IE of a prosthetic valve conduit. None of the children had an artificial heart valve in place.

Of the 7 children with a structural normal heart, 4 patients had a right-sided IE (tricuspid valve) and 3 children a left-sided IE (mitral valve). A computed tomography (CT) of the heart/thorax was not required for IE localization but has been performed to visualize associated complications such as long emboli in 8 patients and an aortic aneurysm in 1 patient. Table 2 shows the localization of IE in all patients.

Embolic Risk and Surgical Treatment

Six of 33 (18%) children diagnosed with right heart IE presented with long emboli visualized by CT not leading to respiratory impairment. Left heart IE led to peripheral emboli in 5 patients and 1 patient presented with a central embolization causing hemiplegia. Embolic complications were not observed in patients with bilateral IE lesions. One patient presented with an aortic root and cerebral abscess.

Nineteen of 53 (36%) children required surgical treatment of IE with 6 patients undergoing urgent surgery at time of IE diagnosis and 1 patient presented with a central embolization causing hemiplegia. Embolic complications were not observed in patients with bilateral IE lesions. One patient presented with an aortic root and cerebral abscess.

Nineteen of 53 (36%) children required surgical treatment of IE with 6 patients undergoing urgent surgery at time of IE diagnosis and 6 patients at 7–15 days after IE diagnosis. In follow-up, additional 7 patients underwent late surgery after 1–9 months. Twelve children were operated for right heart IE, 6 children for left heart IE and 1 for bilateral IE. Among these patients, we observed 5 patients with tricuspid valve involvement, 5 patients with mitral valve involvement and 2 with aortic valve involvement. Interestingly, 8 patients with right heart IE needed operative treatment for...
IE of a prosthetic graft or patch (4 stent mounted BJV grafts, 2 BJV grafts, 1 bovine patch, 1 Gore-Tex conduit).

*S. aureus*, viridans group streptococci and *Staphylococcus epidermidis* were the prominent underlying microorganisms in patients who required surgical treatment (both staphylococci 5 patients, streptococci in 6 patients). All patients but 1 had an underlying CHD. In total, 23 (43%) children required intensive care treatment including 8 (15%) patients under mechanical ventilation.

**Clinical Features at IE Diagnosis**

Patients' record analysis enabled detailed assessment of presenting complaints in 35 (66%) cases. Despite a thorough review, in several patients, it was unclearly documented what initial symptoms led the primary team to consider IE as a possible diagnosis before admission. We therefore did not include these patients in the analysis of clinical features at IE presentation. The majority of patients presented initially with fever of unknown origin [20 of these cases (57%)]. A further 9 (25%) cases presented with sepsis. Three (9%) patients presented with septic embolisms, 2 to the hip and the other to the left foot. An abscess at the sternotomy, dyspnea, toxic shock syndrome and joint complaints in combination with a new cardiac murmur was present each in 1 patient.

**Outcome**

Overall mortality was 13% including one 14-year-old patient with CA IE caused by *S. aureus* that passed away 6 months after medical and surgical treatment of IE because of heart failure. The remaining 6 in-hospital mortalities occurred in HA IE during the first year of life between 2000 and 2004. In these 7 cases, 4 causal organisms were found: coagulase-negative staphylococci present in 4 cases and *S. aureus*, *Candida albicans* and *Enterococcus faecalis* each present in 1 case.

**DISCUSSION**

Analyzing clinical and microbial characteristics of children diagnosed with definite IE, our data clearly indicate the relevance of underlying CHD as a major predisposing risk factor of IE. In addition, a need of surgical intervention in 36% of our population and a mortality rate of 13% indicates that IE is still associated with relevant morbidity and mortality.

Because an underlying CHD was known in 87% of our IE population data strongly support the observation of other reports which state that CHD became one of the major risk factors for IE after the occurrence of rheumatic fever declined. In addition, we observed a high prevalence (62%) of right-sided IE in a CHD heavy population as observed in other CHD cohorts.

Interestingly, IE in unpaired CHD was less common in our patients. Four of 53 patients had an unpaired perimembranous ventricular septal defect, partially associated with valve anomalies or a coarctation. Remarkably, we found no IE in patients with atrial septal defects and only 1 episode because of a patent arterial duct.

Although patients with an univentricular physiology or cyanosis in general are known to be at higher risk to develop IE, we did only recognize 5 of 53 patients. In contrast, patients after TOF or truncus arteriosus repair were very prevalent in our population (18 patients). Because the improvement in survival of patients with complex CHD, more patients appear after surgical treatment with shunts or implanted devices, developing an increasing risk population at pediatric and adult age. Our data verify this evolution by seeing IE in the majority of patients after corrective heart surgery and in 34% after implantation of a prosthetic valve conduit including the cryopreserved homograft and BJV grafts. Sample size of our patient cohort does not allow any suggestion on susceptibility for bacterial adherence to the different valve conduits, but recent clinical observations give special awareness to IE as a complication in patients having undergone transcatheter pulmonary valve replacement using BJV conduits. Larger, multicenter studies are desirable to focus on IE in patients with implanted prosthetic material to gain knowledge on IE pathogenesis in this context in which pathways evoking inflammation and fibrin deposition are not fully understood yet.

It has been stated that the right heart in general is more susceptible for device-related infections, probably because of hemodynamic factors. Cardiac malformations differ in their specific substrate for IE with various disturbances in flow patterns. A non-physiologic flow pattern can favor bacterial adhesion on native or prosthetic valves and, on the other hand, endothelial damage can result of abnormal turbulent blood flow, both mechanisms being considered an IE inciting event.

IE in children with a normal structured heart remains present in a small group of our population (13%). At present, 8%–20% of the IE cases are estimated to occur in children with anatomically normal hearts, in which the course of IE is essentially different if predisposing factors as indwelling central lines leading to healthcare-associated IE are present.

Relatively absent in our patient population were immunocompromised patients (2 children). This supports the hypothesis that endothelial damage associated with transient bacteremia plays a more dominant role in IE pathogenesis than the immunocompromised state.

Focusing at age and IE diagnosis, a group that stands out in our dataset is the large group of patients under 1 year of age. An elevated risk of IE in infants with CHD is known and even described in up to 42% of the investigated pediatric IE population. Knowing that operative treatment for CHD is mainly performed in the first year of life, that and other reasons for intensive care treatment as well as central venous lines may contribute to put these infants at high risk for IE. Our study verifies previous cardiac surgery as being a risk factor in this age group as 14 of 17 infants developed IE in the postoperative period. This was also associated with a high risk of adverse outcome and mortality, as previously reported.

IE caused by viridans group streptococci was the most prevalent in our population, closely followed by coagulase-negative staphylococci and *S. aureus*. This is in line with an Italian pediatric IE survey from 2000 to 2015 where streptococci were found as the major underlying cause in CHD patients. In other studies and an international review of more than 30,000 children, however, *S. aureus* is reported as being the most frequent and important bacterial origin.

Interest has been raised on streptococcal IE after restriction of the antibiotic IE prophylaxis to only patients at high risk. Epidemiologic analysis of streptococcal IE showed no change in incidence before and after introduction of the restricted IE prophylaxis in 1157 US children with definite IE while an increase was seen only for the pediatric age group between 10 and 17 years in a more recent study. As well as in our study, unfortunately no conclusive information on antibiotic IE prophylaxis was available. In our complete study population, we saw an unchanged incidence of IE caused by viridans streptococci over the whole time period (29% vs. 35% in the later time interval), nevertheless, statistical power being limited by the relatively small patient number.

Furthermore, our data indicate a shift from HA IE toward CA IE with a significant increase of CA IE in 2008–2017 (69%). As mentioned, even though streptococci were the most frequent organisms in CA IE, overall incidence of streptococcal IE remained the same.

Conversely, a higher prevalence of HA IE was found in pediatric IE cohorts from New Zealand and the United States.
The divergence in evolution could be caused by the different observation periods or geographic region. In contrast to HA IE, the microbiologic pattern of CA IE seems relatively consistent across the globe with streptococcal species being the most prevalent.21-37 In contrast, in patients diagnosed before 1 year of age, no IE caused by viridans streptococci was seen and most cases were caused by coagulase-negative staphylococci, fungi or S. aureus. This microbiologic pattern seems consistent with earlier findings in neonates and young infants43,44 potentially correlating with an increased susceptibility and exposure to these microorganisms related to frequent HA IE. In our population, fungi and enterococci were rare and exclusively found in this specific subpopulation, having chronic indwelling catheters.45

Culture-negative IE remains present in a minority of our children with a decreasing general trend as microbiologic diagnostics have improved.4,15 The classic clinical hallmarks of IE, such as fever and sepsis, remain major clinical signs found at presentation. This highlights the importance of a high index of suspicion, especially in patients with a history of CHD and implanted prosthetic material.3,4,38

Outcome is influenced by many factors as the virulence of the underlying microorganism and the requirement of surgery.37 The need of surgical treatment in 36% of our patient cohort is similar to that of other series and reflects the high morbidity because of IE.4,39 The most frequently performed operation was a valved graft replacement in the right ventricular outflow tract. Surgical treatment was curative in all but 1 patient. We did not observe associated mortality with the need of surgical treatment. Our overall mortality is comparable with results from recent pediatric IE series describing a mortality rate of about 10%.4,9 Mortality was concentrated in those children diagnosed <1 year of age and before 2004. The sharp decline after 2004 highlights the potential impact of vigilance and early treatment, especially in a high-risk population.

Prevalence and prevention of IE gained attention in the context of changed guidelines for IE prophylaxis from 2007 onwards.21-40 As also shown by our data, the vast majority of patients presenting with IE are still included in the classification of high-risk patients and therefore eligible for antibiotic IE prophylaxis. Given the limits of antibiotic prophylaxis and available therapeutic interventions, it is likely that further improvement in IE outcome has to focus on different prevention measures, as also included in the 2015 actualized ESC IE guidelines.4,21,41 Studies evaluating the knowledge on IE in CHD patients or parents show a striking lack of understanding IE, dental hygiene and awareness of symptoms requiring an adequate medical investigation and no self-treatment.42,43 Therefore, a structured education of physicians and patients seems to be a necessary major target in daily practice to improve knowledge and thereby contributing to a decrease in diagnostic delay of IE, which still represents an important burden of the disease.

REFERENCES


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