Early atherosclerotic lesions of the cardiac conduction system arteries in infants

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Abstract

Introduction: Although several studies have described initial atherosclerotic lesions of the coronary arteries, already detectable in infancy and even during the intrauterine life, little, if any, attention has been given to the possible involvement of the cardiac conduction system arteries. In particular, to the best of our knowledge, none has considered the lesions of the cardiac conduction arteries as an initial stage of atherosclerosis. Methods: The cardiac conduction system of 70 infants dying suddenly and unexpectedly was removed in two blocks for paraffin embedding and serially cut. Results: The histological study of the cardiac conduction arteries of the 70 cases examined showed a normal structure in 55 cases (78.57%). In 15 cases (21.43%), there was a thickening of the sinoatrial node and/or atrioventricular artery associated with a thickening of varying severity in coronary artery walls. The lesions were marked by thickening and deposits of amorphous material and mainly lipids in the intima, as well as fragmentation of the elastic fiber system. A significant correlation was evident between early atherosclerotic lesions and both formula feeding and parental cigarette smoking ($P < .05$, $\chi^2$ test). Conclusions: The combination of both the considered risk factors seems to increase the early atherogenic effect of each noxa because the coronary lesions were more diffused in formula-fed infants whose parents both smoked. © 2004 Elsevier Inc. All rights reserved.

Keywords: Atrioventricular node artery; Sinoatrial node artery; Early atherosclerotic lesion; Cardiac conduction system; Cigarette smoking; Formula feeding

1. Introduction

Death has climbed in through our windows and has entered our fortresses; it has cut off the children from the streets and the young men from the public squares (Jeremiah 9:21)

Although several studies have described initial atherosclerotic lesions of the coronary arteries, already detectable in infancy [1–4] and even during the intrauterine life [5–7], little, if any, attention has been given to the possible involvement of the cardiac conduction system arteries. In particular, to the best of our knowledge, none has considered the lesions of the cardiac conduction arteries as an initial stage of atherosclerosis.

The finding of early atherosclerotic alterations in infant and fetal coronary arteries [1–4,6,7] prompted us to analyze whether there might be an early atherosclerotic involvement also of the cardiac conduction system arteries.

The aim of the present study is to examine the morphologic structure of the cardiac conduction system arteries in 70 infants dying suddenly and unexpectedly.

2. Material and methods

2.1. Selection of cases and method for examination of the cardiac conduction system

Seventy cases of infants (47 males and 23 females) with unknown medical problems dying suddenly and unexpectedly (SIDS) [8] were selected for this study. Their ages range from 20 to 365 days (mean ± S.E.M. = 91.90 ± 9.55).

The clinical data collected for each case included the type of feeding, as well as any parental smoking habit (maternal cigarette smoking before and/or during pregnancy,
postnatal cigarette smoking by one or both parents and the number of cigarettes smoked daily). The cholesterol level, when available, was within normal range in both infants and mothers.

We have divided the cases into three subgroups: (A) from 0 to 6 weeks; (B) from 7 to 20 weeks; and (C) from 21 to 52 weeks (Table 1).

In each case, a complete autopsy was performed according to the autopsy protocol usually followed by the Institute of Pathology, University of Milan, in case of sudden and unexpected infant death [9], available on the web site: http://users.unimi.it/~pathol/sids/riscontro_diagnostico_e.html.

Cooperating pathological centers received a brief protocol describing the procedures for the removal and preservation of hearts. The well-known procedure of measuring heart size and weight was not omitted, and the values measured were compared with the normal values for infants of that length and age [10]. After the presence of gross cardiac malformations are excluded, the origin of the coronary arteries were carefully inspected. The hearts were regularly examined for pathologic changes in the atria, septa, ventricles, pericardium, endocardium and coronary arteries. Samples of the myocardium were stained with hematoxylin–eosin and trichromic Heidenhain (Azan).

Our cases were examined in blinded fashion, without initial knowledge of the cause of death, age, or other clinicopathological information. Only after the histological assessment of the conduction system had been completed were the pathological findings matched with the corresponding records.

The cardiac conduction system was removed in two blocks for paraffin embedding [9]. Block 1 consisted of a portion of the right atrial wall including the lateral half of the funnel of the superior venal cava, sulcus and crista terminalis. This first block includes the sinoatrial node, its atrial approaches, the crista terminalis. This second block contains the atrioventricular septum. This second block contains the atrioventricular node, His bundle, bifurcation and bundle branches. For the histological examination of the cardiac conduction system, the two blocks were cut serially at intervals of 40 µm (levels), as devised by one of the present authors [11]. For each level, two sections of 8 µm were saved and stained alternately with hematoxylin–eosin and Azan. All intervening sections were kept and stained as deemed necessary. For each heart, the average number of histologic sections stained and examined is about 200, as described [12].

### 2.2. Coronary and cardiac conduction arteries histologic examination

According to the protocol followed by Roberts and Jones [13], the hearts were fixed in formalin for at least 1 day. The major epicardial coronary arteries (left main, left anterior descending branch, left circumflex, right posterior interventricular descending branch, right marginal branch) were excised transversely to their longitudinal axis in segments approximately 3–4 mm long. Each segment was labeled sequentially from either its aortic ostium or from its origin from the left main coronary artery. They were dehydrated, embedded in paraffin block and serially cut. The sections of each block were stained with hematoxylin–eosin, Azan for histological examination, Alcian blue (at pH 0.5 and 2.5) for acid mucopolysaccharides analysis, Weigert for elastic fibers identification and, in selected cases, submitted to specific immunohistochemical methods for lymphocyte, monocyte and smooth muscle cell typization. The portions of the sinoatrial and atrioventricular nodal arteries were assessed in the prenodal and intraparenchimal branches, following the serial sections of the two cardiac conduction tissue blocks, without restriction by arterial diameter.

### 2.3. Cellular typization of the coronary and cardiac conduction system arteries

In selected cases, immunophenotyping of the cells present in each preatherosclerotic lesion was performed using the immunoperoxidase technique with the following primary antibodies (made from Dako reagents): anti-α-smooth muscle actin to identify smooth muscle cells (SMC),

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**Table 1**

Clinical data and coronary and conduction nodal arteries lesions in infants who died suddenly, divided by postnatal age intervals

<table>
<thead>
<tr>
<th></th>
<th>Subgroup A</th>
<th>Subgroup B</th>
<th>Subgroup C</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of cases</td>
<td>18</td>
<td>39</td>
<td>13</td>
<td>70</td>
</tr>
<tr>
<td>Sex (M/F)</td>
<td>13/5</td>
<td>23/16</td>
<td>11/2</td>
<td>47/23</td>
</tr>
<tr>
<td>Mean age (days) ± S.E.M.</td>
<td>39.13±3.22</td>
<td>86.16±3.94</td>
<td>213.4±25.57</td>
<td>91.90±9.55</td>
</tr>
<tr>
<td>Parents’ smoke</td>
<td>9</td>
<td>19</td>
<td>8</td>
<td>36 (51.42%)</td>
</tr>
<tr>
<td>Bottle feeding</td>
<td>5</td>
<td>15</td>
<td>8</td>
<td>28 (40%)</td>
</tr>
<tr>
<td>Mixed feeding</td>
<td>6</td>
<td>8</td>
<td>3</td>
<td>17 (24.29%)</td>
</tr>
<tr>
<td>Breast feeding</td>
<td>7</td>
<td>16</td>
<td>2</td>
<td>25 (31.71%)</td>
</tr>
<tr>
<td>Coronary artery preatherosclerotic lesions</td>
<td>2</td>
<td>16</td>
<td>5</td>
<td>23 (32.85%)</td>
</tr>
<tr>
<td>Coronary artery juvenile soft atherosclerotic lesions</td>
<td>—</td>
<td>6</td>
<td>5</td>
<td>11 (15.71%)</td>
</tr>
<tr>
<td>Conduction nodal arteries lesions</td>
<td>1</td>
<td>9</td>
<td>5</td>
<td>15 (21.43%)</td>
</tr>
</tbody>
</table>

Subgroups: A: 0–6 weeks; B: 7–20 weeks; and C: 21–52 weeks.
anti-CD68 to identify monocytes and anti-CD20 and anti-UCHL-1 to identify the B- and T-lymphocytes, respectively.

2.4. Definition of terms

(1) **Pre-atherosclerotic lesions.** The earliest morphologically recognizable alterations are characterized by fragmentation of the tunica media, with intense infiltration of SMC, often arranged perpendicularly in columns or tangentially to the axis of the tunica itself. These SMC infiltrate the intima, which tends to be thickened, also due to initial mucopolysaccharide deposits, mainly composed of Types A and C chondroitin sulfates and hyaluronic acid. Rare monocytes and B-lymphocytes are also present. The endothelium is morphologically intact. The internal elastic membrane appears to be focally fragmented [5].

(2) **Juvenile soft atherosclerotic plaques.** Such lesions are characterized by wide myointimal thickness with alteration of the elastic fibers and of the tunica media. They are defined as “soft plaques”, as they have abundant SMC, associated with the moderate infiltration of monocytes and rare lymphocytes. Amorphous deposits of acid mucopolysaccharides, mainly composed of B chondroitin sulfates, are present, occasionally also in the internal portion of the tunica media. There is evident fragmentation of the elastic fiber system and of the tunica media. The endothelium is morphologically intact. Reduction of the lumen varies from approximately 10–15% to 30–40% [14–16].

2.5. Statistical analysis

The significance of differences between group parameters were evaluated by $\chi^2$ test. The level of significance chosen was $P < .05$, two-tailed.

3. Results

3.1. Cardiac conduction arteries findings

The histological study of the cardiac conduction arteries of the 70 cases examined showed a normal structure in 55 cases (78.57%). In 15/70 cases (21.43%), there was a thickening of the sinoatrial node and/or atrioventricular artery associated with a thickening of varying severity in coronary artery walls. The lesions were marked by the infiltration in the intima of SMC, monocytes/foam cells, rare B-lymphocytes and deposits of amorphous material, mainly composed of mucopolysaccharides, as well as focal fragmentation of the elastic fiber system (Figs. 1 and 2). The intima was thickened due to the initial mucopolysaccharide deposits, mainly composed of Types A and C chondroitin sulfates and hyaluronic acid. Rare macrophages and B-lymphocytes were also present.

3.2. Coronary arteries findings

In 34/70 cases (48.57%), a thickening of varying severity in the coronary artery walls was detected (Fig. 3). In 23/70 cases, the lesions were preatherosclerotic, mostly located in the anterior descending branch of the left coronary artery. In 11/70 cases, juvenile soft atherosclerotic lesions were observed in one or more coronary branches. Such lesions were characterized by deposits of amorphous material, mainly lipids, even in the innermost portion of the media, and significant fragmentation of the elastic fiber system (Fig. 3).

In 19/34 cases, the thickening of the coronary artery was not associated to lesions of the sinoatrial and atrioventricular nodal arteries.

Fig. 1. Narrowing of the atrioventricular node artery in a 6-month-old child dying suddenly and unexpectedly. A= right atrium; AVN = atrioventricular node; CF= central fibrous body. Trichromic Heidenhain; magnification ×20.

Fig. 2. Fragmentation of the internal elastica lamina in early atherosclerotic lesion. Weigert stain; magnification ×200.
3.3. Feeding type and parents’ cigarette smoking

In 28 cases (40%), the infants were fed from birth, or after a few days of breast feeding, only with formula. In 17 cases (24.29%), feeding was mixed (about 50% formula and 50% human milk) from birth. Twenty-five babies (31.71%) were breast fed. In 36 cases (51.42%), at least one parent was a smoker and generally smoked over five cigarettes a day. In the smoker mothers, the smoking habit had started before pregnancy. In 20 cases (28.57%), formula feeding was combined with cigarette smoking of one or both parents. The lesions in the cardiac conduction arteries appeared to be larger and with higher number of slides, showing changes when the formula-fed babies had smoker parents (9/15 cases, i.e., 60% of cases).

The clinical data collected disclosed that in 13/15 (86.66%) infants with intimal thickening of the cardiac conduction arteries, the feeding was only with formula (8 cases) or mixed (5 cases). In 12/15 (80%) infants with intimal thickening, at least one parent was a smoker.

Among the infants with normal cardiac conduction arteries, 32/55 (58.18%) were formula (20 cases) or mixed (12 cases) fed, and in 24/55 (43.63%), one or both parents were smokers.

A significant correlation is evident between early atherosclerotic lesions and both formula feeding and parental cigarette smoking ($P < 0.05$, $\chi^2$ test).

Regarding age, the infants’ cases showed a peak between the 5th and the 12th weeks of life. In particular, 21% of cases occurred at the second month, 22% at the third, 13% at the fourth and 10% at the fifth month. Table 1 shows the clinical data and coronary and conduction nodal artery lesions in infants who suddenly died, divided by postnatal age intervals.

4. Discussion

It has been reported that early atherosclerotic or preatherosclerotic lesions are already detectable in infancy [1–4] and even during fetal life [5–7]. Such initial changes are characterized by the thickening of the intima, disruption of the internal elastic lamina, proliferation of the medial SMCs and accumulation of glycosaminoglycans in the fundamental substance of the initial connective tissue [1,5,13–20].

In this study, the histological examination of the serially sectioned cardiac conduction arteries from 70 infants who died suddenly and unexpectedly revealed the frequent presence of early atherosclerotic lesions in 15 of our 70 cases (21.43%). The lesions are marked by infiltration in the intima of SMC, monocytes/foam cells, rare B-lymphocytes and deposits of amorphous material, mainly composed of mucopolysaccharides, as well as fragmentation of the elastic fiber system (Fig. 1).

In a recent study [5] carried out on the coronary arteries of human fetuses, we observed early preatherosclerotic alterations in over 50% of the cases, prevalently on late fetuses of smoker mothers. We interpreted these fetal coronary lesions to have been enhanced by the gaseous products of nicotine.

Fibromuscular hyperplasia or dysplasia [21] of the sinus node and/or AV node arteries has been described as a cause of death in young people and adults [22,23]. Anderson and Hill [24], analyzing 40 victims of crib death, found five cases (12.5%) with increased thickness of the AV artery and a case with stenosing thickness of the sinus node artery. The authors hypothesized that this thickness may explain an ischemia of the conducting tissue, with consequent cardiac arrhythmias and/or heart block. Suárez-Mier and Aguilera [25] reported a fibromuscular hyperplasia in 1.8% of SIDS cases.

The formula feeding and the parents’ cigarette smoking might have a role in the determinism of the herein-described cardiac conduction arteries lesions.

The consequences of baby formula reported in the literature include metabolic disturbances [26–28], delayed
development of the nervous system [29,30] and dyslipidemia [31,32]. In particular, experimental studies on baboons [33,34] and a study on the composition of various types of commercial milk [35] have shown that unlike the fatty acids present in maternal milk, those contained in baby formula cause an increase in the serological cholesterol level, which favors the development of atherosclerosis.

Cigarette smoking has an important atherogenic role and is a major cause of cardiovascular diseases correlated with atherosclerosis in adults [36–39]. Tobacco smoking has been implicated in the initiation and progression of atherosclerotic vascular lesions [31], and recently, we have reported the presence of preatherosclerotic coronary artery lesions in human fetuses of smoker mothers [5]. The combination of both the considered risk factors seems to increase the early atherogenic effect of each noxa because the coronary lesions were more diffused in formula-fed infants whose parents both smoked.

In this study, the relationship between preatherosclerotic lesions and both formula feeding and parental cigarette smoking was statistically significant ($P < .05$, $\chi^2$ test).

According to the present findings, we believe that if the results of our series are confirmed by other further studies of the cardiac conduction system on serial sections, the significance of the lesions of the cardiac conduction system arteries would be better understood. Much meticulous clinical–pathologic correlative work between the early atherosclerotic lesions and the risk factors in larger series remains to be done.

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