Kawasaki disease (KD) is the leading cause of acquired heart disease in children in the developed world (1). Typically, KD presents in children younger than 5 years of age as a febrile illness with mucocutaneous changes (2). A subset of patients will develop permanent damage to the arterial wall, valve leaflets, and myocardium. The acute phase of the illness is self-limited, and the diagnosis may be missed. If untreated, KD can result in coronary aneurysms in 25% of patients (3). Patients who experience coronary artery damage may develop thrombosis or stenotic lesions associated with the aneurysms and are at risk of myocardial infarction, sudden death, and congestive heart failure (4,5). The prognosis for adults who have recovered from KD without coronary aneurysms is postulated to be good, but longitudinal studies have not been performed to test this hypothesis.

In 2004, the American Heart Association (AHA) issued updated guidelines for the care and management of children with KD (6). However, studies of the optimal therapy and management of the sequelae of KD in adults have not been performed, nor have official guidelines been developed that provide direction for cardiologists as they begin to care for the growing number of young adults who experienced KD in childhood. The available data suggest that the pathophysiology of the vascular lesion in KD is unique and distinct from atherosclerosis. Here, we undertake a critical review of the literature describing the cardiovascular manifestations in young adults who experienced KD in childhood.

Epidemiology and Clinical Aspects of Acute KD

It is estimated that >4,000 new cases of KD are diagnosed in the U.S. each year (7). In Japan, where the incidence is approximately 10-fold greater as compared with the U.S., more than 10,000 new cases are diagnosed each year (8). Seasonality of cases, nationwide epidemics, and the self-limited nature of the acute illness suggest an infectious trigger, but no causative agent has been identified. Genetic influences on disease susceptibility and outcome have been identified, and the current paradigm proposes that KD results from exposure to a common agent that triggers the syndrome only in genetically susceptible hosts (9–11).

The self-limited clinical syndrome is recognized through a constellation of clinical signs that include fever for at least 4 days associated with rash, conjunctival injection, erythema of the lips and oropharynx, edema of the hands and feet, erythema of the palms and soles, and, in the convalescent phase, periungual desquamation (6). Up to 25% of untreated children will develop permanent damage to the coronary arteries with inflammatory cell infiltration of the arterial wall, destruction of the internal elastic lamina, necrosis of smooth muscle cells, myointimal proliferation, and subsequent aneurysm formation (3,12). Aneurysms of systemic, extraparenchymal muscular arteries also occur in a subset of patients with coronary aneurysms (13). The proximal coronary arteries can be readily imaged in infants and children by the use of transthoracic echocardiography, which permits a reliable and reproducible measurement of the internal...
diameter of the proximal right and left anterior descending coronary arteries and the expression of these measurements as standard deviation units (Z score) normalized for body surface area (14–16).

Administration of a single dose of intravenous immunoglobulin (IVIG) in conjunction with aspirin within the first 10 days after fever onset reduces the incidence of aneurysms from 25% to 3% to 5% (17,18). The AHA guidelines have categorized these patients into 5 groups on the basis of coronary artery Z scores and morphology of the coronary artery lesions (Table 1) (6). Approximately 30% of IVIG-treated children with KD will develop transient dilation of the coronary arteries (Z score ≥ 2.5 for the right coronary artery or left anterior descending artery, AHA risk level II) (16,19). Another 5% to 10% will develop coronary artery aneurysms, which in some cases can be attributed to delayed diagnosis and treatment. There are no data regarding the number of young adults in the U.S. who experienced KD delay in childhood. If we estimate 4,000 KD patients/year in the U.S. beginning in 1986 when IVIG treatment was first recommended, assume an average age of onset of 2 years, a rate of 1,400 individuals each year. These numbers are clearly underestimated because many KD patients were older than 2 years at the time of disease onset and many clinical cases were never diagnosed. Whatever the real figures are, there is clearly a growing population of young adults with potentially important coronary artery disease after KD in childhood, and cardiologists specializing in adult patients must be prepared to care for them.

**Cardiovascular Sequelae of KD in Adults**

Only in Japan has an attempt been made to systematically collect mortality data on patients with KD. During a 10-year period from 1982 to 1992, a cohort of 6,576 Japanese patients was created, and the seventh report on the status of the cohort was recently published (20). As of 2004, there were 3,326 adult subjects in the cohort ages 20 to 34 years with an average observation period of approximately 17 years. Standardized mortality ratios (SMRs) were calculated for subsets in the cohort based on sex, coronary artery sequelae, and timing of death after KD. After the acute phase, the SMR was increased only for male patients with known cardiovascular sequelae (expected deaths: 3.9, observed deaths: 10; SMR 2.55, 95% confidence interval: 1.23 to 4.70). The cause of death in these 10 male patients was listed as coronary artery insufficiency (n = 2), acute myocardial infarction (n = 3), sudden death (n = 1), congestive heart failure (n = 1), influenza pneumonia

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**Table 1 American Heart Association Risk Stratification for Kawasaki Disease**

<table>
<thead>
<tr>
<th>Risk Level</th>
<th>Pharmacological Therapy</th>
<th>Follow-Up and Diagnostic Testing</th>
<th>Invasive Testing</th>
</tr>
</thead>
<tbody>
<tr>
<td>I (no coronary artery changes at any stage of illness)</td>
<td>None beyond first 6–8 weeks</td>
<td>Cardiovascular risk assessment, counseling at 5-yr intervals</td>
<td>None recommended</td>
</tr>
<tr>
<td>II (transient coronary artery ectasia disappears within first 6–8 weeks)</td>
<td>None beyond first 6–8 weeks</td>
<td>Cardiovascular risk assessment, counseling at 3- to 5-yr intervals</td>
<td>None recommended</td>
</tr>
<tr>
<td>III (1 small-to-medium coronary artery aneurysm/major coronary artery)</td>
<td>Low-dose aspirin (3–5 mg/kg aspirin/day), at least until aneurysm regression documented</td>
<td>Annual cardiology follow-up with echocardiogram + ECG, combined with cardiovascular risk assessment, counseling; biennial stress test/evaluation of myocardial perfusion scan</td>
<td>Angiography, if noninvasive test suggests ischemia</td>
</tr>
<tr>
<td>IV (≥1 large or giant coronary artery aneurysm, or multiple or complex aneurysms in same coronary artery, without obstruction)</td>
<td>Long-term antiplatelet therapy and warfarin (target international normalized ratio 2.0–2.5) or low-molecular-weight heparin (target: antifactor Xa level 0.5–1.0 U/ml) should be combined in giant aneurysms</td>
<td>Biannual follow-up with echocardiogram + ECG; annual stress test/evaluation of myocardial perfusion scan</td>
<td>First angiography at 6–12 months or sooner if clinically indicated; repeated angiography if noninvasive test, clinical, or laboratory findings suggest ischemia; elective repeat angiography under some circumstances</td>
</tr>
<tr>
<td>V (coronary artery obstruction)</td>
<td>Long-term low-dose aspirin; warfarin or low-molecular-weight heparin if giant aneurysm persists; consider use of beta-blockers to reduce myocardial O2 consumption</td>
<td>Biannual follow-up with echocardiogram and ECG; annual stress test/evaluation of myocardial perfusion scan</td>
<td>Angiography recommended to address therapeutic options</td>
</tr>
</tbody>
</table>

Adapted from Newburger et al. (6). ECG = electrocardiogram.
Late complications of KD may include residual damage from the acute vasculitis as well as the myocarditis (21). In the only longitudinal angiographic study of KD, Kato et al. (3) described the natural history of the vascular lesions in 594 Japanese subjects 10 to 12 years after the onset of KD (Fig. 1). This study was performed before the routine use of IVIG to treat acute KD. Long-term complications included stenosis, myocardial infarction, and death. Other series have reported both symptomatic and asymptomatic coronary artery occlusion in regions of previous aneurysms (22,23), coronary artery stenosis (24), coronary artery calcification (25), diastolic dysfunction (26), and sudden death (27). One study by Tsuda et al. (28) of 562 Japanese KD patients with known coronary artery lesions documented the appearance of new aneurysms 2 to 19 years after disease onset in 15 patients (3%). Patients with KD with aneurysms at least 6 mm in maximal diameter had a greater than 50% chance of developing a clinically significant stenotic lesion during a mean follow-up period of 8 years (24). Most of these series are from Japan, and none report longitudinal outcome in KD patients who were treated with IVIG during the acute illness. On the basis of the accumulating evidence, it is likely that patients with known aneurysms during the acute phase of KD will have some cardiovascular morbidity as young adults. The fate of IVIG-treated children who had no lesions detected by echocardiography as a result of the acute vasculitis is unknown (Figs. 2A and 2B).

During acute KD, approximately 2% of patients will develop valvulitis, followed by scarring of the valve leaflets, most commonly in the mitral valve, leading to valvular incompetence and the need for valve replacement in a subset of patients (29–31). Another manifestation of the cardiovascular damage after acute KD is the progressive dilation of the aortic root (32). In a longitudinal study of 100 children with KD, the body surface area-adjusted aortic root dimension was significantly larger when compared with control subjects. Mild aortic valve regurgitation was noted at the 1-year visit in 4% of KD patients (32,33).
Focus on the coronary aneurysms has diverted attention from other potential cardiovascular complications of KD. Diffuse myocarditis followed by myocardial fibrosis may lead to systolic or diastolic dysfunction in a subset of KD patients (34–36). During acute KD, tissue Doppler interrogation of left ventricular inflow velocities has revealed abnormal relaxation patterns and was associated with increased levels of B-type natriuretic peptide (35). Decreased ventricular contractility and abnormal relaxation may be the consequences of antecedent myocardial inflammation, and late-onset ventricular arrhythmias and congestive heart failure have been observed (21,37).

Missed KD Presenting in Adulthood

In 1992, Kato et al. (38) in Japan published the results of a questionnaire survey sent to adult cardiologists that asked about adult patients with angiographically proven coronary artery disease that could reasonably be attributed to antecedent KD. The survey identified 130 patients ages 20 to 63 years: 21 adults had ischemic heart disease definitely (n = 2) or possibly (n = 19) attributed to antecedent KD, and 109 adults had coronary artery aneurysms with no clear history of clinical KD in childhood. Of the 21 patients with a high likelihood of antecedent KD, 3 had died of ischemic complications, and 18 were alive with serious cardiovascular sequelae, including mitral regurgitation requiring valve replacement, arrhythmias, dilated cardiomyopathy, and congestive heart failure. The authors concluded that all 130 patients likely had KD as the cause of their vascular damage and that cardiologists specializing in adult patients should be aware of these patients in their clinical practice. No subsequent survey from Japan or elsewhere has been performed to estimate the number of adult patients under the care of a cardiologist for the sequelae of KD.

Young adults with no known previous cardiovascular history may present with angina, myocardial infarction, ischemia-induced arrhythmia, or sudden death (39). Characteristics of the coronary arteries that should prompt questioning about antecedent KD include proximal aneurysms with or without calcification followed by an angiographically normal distal segment. Young adults in Japan have presented with ventricular tachycardia in the setting of left heart failure decades after missed KD (37). Investigation of these patients revealed the classic calcified aneurysms of KD. Thus, the acute vasculitis and myocarditis associated with KD may lead to a complex set of cardiovascular problems later in life.

Pathology

Although the term “atherosclerosis” has been loosely applied to the progressive vascular lesion after KD, evidence suggests that this is a misnomer. Although many autopsy reports of individuals who died late after KD describe calcified aneurysms, myointimal proliferation, and organizing thrombus in the coronary arteries with recanalization, there is scant mention of lipid-laden macrophages and cholesterol crystals, the hallmarks of established atherosclerosis (40,41). There is ample histological evidence to support the concept that coronary arteries that develop aneurysms as a result of the intense inflammatory process during acute KD have very abnormal architecture despite normalization of the lumen and angiographic evidence of “healing” (42,43). The typical lesions are discrete regions of myointimal proliferation associated with disrupted internal elastic lamina and medial smooth muscle cell necrosis with replacement by fibrosis and calcification. Suzuki et al. (44) examined 7 subjects at autopsy who died 3 to 12 years after acute KD. Active remodeling of the aneurysms with intimal proliferation and neoangiogenesis was evident in all cases. None of the subjects had coronary artery lesions consistent with atherosclerosis. Takahashi et al. (45) described a series.
of 6 subjects with arterial lesions who died from either sudden death (n = 5) or sepsis (n = 1) at least 15 years after acute KD. Only 1 of the 6 subjects had pathology consistent with atherosclerosis. No clinical details were provided in this report, so it is unknown whether this 39-year-old subject with typical atherosclerotic lesions had cardiovascular risk factors superimposed on his antecedent KD. The extent to which KD is a cardiovascular risk factor for the future development of atherosclerosis is unknown.

Another controversial issue is the pathology of arteries in IVIG-treated individuals whose echocardiograms showed no evidence of structural damage to the coronary artery wall. The scant autopsy data available on the histological changes in children and young adults who have died late after KD appear as case reports, many of them in the forensic literature (39). In an autopsy study of a child who died from unrelated causes 13 months after recovery from KD with normal echocardiograms, the intima of the coronary artery was thickened, the internal elastic lamina was disrupted, and smooth muscle cells were observed infiltrating into the intima (46). Further autopsy studies of incidental deaths in subjects with antecedent KD and normal echocardiograms will be necessary to determine how often KD leads to pathologic changes in coronary arteries with no dilation noted during the acute phase of the illness. A prospective registry of KD cases would allow tracking of deaths and correlation of the histopathology with echocardiographic assessment during the acute illness.

Pathologic changes in the myocardium as a result of injury during the acute inflammatory phase also have been reported. Three series from Japan reported endomyocardial biopsies at various time points after the acute illness (47–49). In the study by Yonesaka et al. (48) hypertrophy, degeneration of myocytes, and fibrosis were noted in the majority of biopsies obtained at least 3 years after KD in 38 subjects, and their presence was more frequent in subjects who had experienced coronary artery aneurysms during the acute phase. Yutani et al. (47,50) found myocardial abnormalities, including lymphocyte and plasma-cell infiltration, myocardial fibrosis, and disarray of myocardial fibers in every biopsy from the 201 KD subjects in their study. Changes were most pronounced in patients studied 4 or more years after disease onset and the investigators raised the question of progression of myocardial pathology over time.

Some autopsy reports of adults late after KD describe cardiomyocyte dropout and diffuse fibrosis not in the walled distribution of the epicardial coronary arteries (Fig. 2B) (51–53). Increasing numbers of deaths that are attributed to left ventricular dysfunction and presumed ventricular arrhythmias are being reported in young adults from Japan with antecedent KD (27). Whether the diffuse fibrosis is a consequence of ischemic injury from microinfarcts or inflammatory cardiomyocyte injury or both is unknown.

### Endothelial-Cell Dysfunction and Arterial Wall Injury

In addition to causing coronary artery aneurysms and stenoses, KD also has been shown to have deleterious effects on coronary artery function years after the acute presentation. Arterial function has been assessed in multiple studies in which the authors evaluated myocardial and coronary flow reserve with angiography (54–57), invasive flow measurements (58–60), positron emission tomography (61–63), and transthoracic Doppler ultrasound of the coronary arteries (64). Abnormalities have been observed in endothelial-mediated as well as endothelial-independent coronary artery vasodilation. In patients with persistent coronary artery aneurysms, coronary artery function has repeatedly been shown to be impaired compared with control subjects (56,57,59–61,64). Coronary flow reserve is also abnormal in patients with a history of transiently dilated coronary arteries (54–57,60–62) and in patients with evidence of inducible ischemia but angiographically normal-appearing coronary arteries (58,59). Even in patients with a history of KD without antecedent coronary artery dilation or ischemia, impaired coronary flow reserve has been observed (54,60,61,63,64), although conflicting studies showed no difference from normal control subjects (55–57). Thus, the damage to coronary arteries appears to extend beyond that which is observed angiographically.

Studies of systemic vascular function via flow-mediated dilation of the brachial artery have yielded conflicting results. Dhillon et al. (65) demonstrated abnormal brachial artery reactivity in 20 male subjects in the United Kingdom ages 11 to 19 years who were studied 5 to 17 years after the onset of KD, irrespective of whether they had developed aneurysms during the acute phase. In studies of Japanese subjects of approximately the same age and interval from disease onset, only patients with previous aneurysms had abnormal flow-mediated dilation (66,67). Finally, the authors of a recent study (68) of 52 Canadian KD subjects concluded that there was no evidence of long-term endothelial-cell dysfunction late after KD. It is unclear whether these conflicting results are best explained by methodological issues or genetic differences of the populations studied. Metabolic abnormalities manifested as abnormal lipid profiles with persistently low high-density lipoprotein may also contribute to endothelial cell dysfunction (69,70).

Carotid intimal medial thickness (IMT) measured non-invasively by ultrasound is used as a surrogate marker of coronary atherosclerosis and has been correlated with the risk of stroke and myocardial infarction in adults with atherosclerosis (71). The authors of several small studies have demonstrated increased carotid IMT in patients who have recovered from KD, but the significance of this observation is unclear (66,72–74). Patients with remodeled coronary aneurysms have even greater carotid IMT than those without aneurysms. However, no study to date has
correlated clinical outcomes with IMT findings in KD patients. There is no evidence that increased carotid IMT in KD patients implies atherosclerosis and the prognostic significance of this measurement is unknown.

**Identifying the Adult With Possible Antecedent KD**

The cardiologist specializing in adult patients should be sufficiently familiar with the signs and symptoms of acute KD to allow questioning of the patient or parent about an antecedent KD-compatible illness that was not diagnosed. Features of the illness that are frequently recalled by patients and parents are the prolonged fever, rash, “bloodshot” eyes in the acute phase, and peeling of the fingers and toes in the convalescent phase. Common misdiagnoses for KD include viral syndrome, measles, scarlet fever, allergic reaction to antibiotics, and Stevens-Johnson syndrome (4). Because the etiology remains unknown, there is no specific diagnostic test that can be used to make a retrospective diagnosis.

Imaging studies may be helpful in identifying patients with antecedent KD (Level of Evidence [LOE]: C, Class IIa). Calcification of the arterial wall in regions where former aneurysms have remodeled are hallmarks of KD and may even be apparent on chest radiograph (41,75). In a study of Japanese young adults 20 years after acute KD, 94% of those with aneurysms at least 6 mm in internal diameter during the subacute phase of the illness had calcification observed by electron beam computed tomography (CT) (76). Arterial-wall calcification can be also be imaged by multislice CT angiography (Fig. 3). Coronary calcium scores are elevated in a subset of KD subjects with antecedent vascular injury (77). Suspicion of antecedent KD should be high in the young adult with coronary artery calcification and a low-risk profile for atherosclerosis.

Differentiation of atherosclerotic aneurysms from KD may be aided by angiography (LOE: C, Class IIb). Whereas atherosclerosis tends to involve the arterial wall diffusely, KD results in focal abnormalities only at the site of previous aneurysms. Vessels proximal and distal to the lesion appear healthy and have a smooth luminal surface and normal diameter (Fig. 4). Remodeled aneurysms of KD may be more difficult to recognize, and intravascular ultrasound may be useful to detect the thickened arterial wall of the aneurysm (78,79).
Magnetic resonance (MR) imaging is emerging as the modality of choice for imaging structural damage late after KD (LOE: C, Class IIa). Repeated imaging is possible without concerns for radiation exposure. Myocardial inflammation with the use of T2-weighted images as well as myocardial scarring and fibrosis can be detected as late gadolinium enhancement (80–82). At the present time, gadolinium MR angiography is not as sensitive as CT angiography or cardiac catheterization for the detection of coronary artery stenosis (80).

**Management of the Adult With Antecedent KD**

The optimal management of adults with antecedent KD can only be established through systematic study of this patient population with the creation of regional or national registries to record patient histories and outcomes and generate hypotheses regarding their care that could be tested in appropriately powered clinical studies. One challenge of studies designed to collect cardiovascular outcome data will be to differentiate the effects of antecedent KD from the effects of traditional cardiovascular risk factors, which will accumulate as this population ages. For now, the management of these patients must be guided by common sense and an appreciation of the uncertainty about the cardiovascular outcomes in adults who experienced KD in childhood. The Japanese Circulation Society published guidelines (83) based on the opinion of experts in Japan in 2003 for the management and treatment of adults who had KD in childhood (Table 2). The guidelines state that cardiovascular symptoms in KD patients only begin to appear 2 decades after the onset of the acute disease, so only now are patients beginning to present with sequelae.

As with all clinical practice, the admonishment of primum non nocere must be heeded. Subjecting otherwise-healthy, asymptomatic individuals to frequent testing may have undesirable psychological consequences. At the same time, ignoring a past history of KD may lead to missed opportunities for intervention in a subset of these patients. A rational approach to these patients might include standard testing for cardiovascular risk assessment as well as specific testing to detect subclinical ischemia, valvular dysfunction, and myocardial fibrosis caused by antecedent KD. Testing might include a lipid profile, high sensitivity C-reactive protein level, electrocardiogram, 2-dimensional transthoracic echocardiogram, and stress echocardiogram (LOE: C, Class IIb). Because arterial calcification is a feature of KD vascular lesions, a single, baseline coronary artery calcium score by CT may be informative, particularly for patients whose initial evaluation by echocardiogram in childhood was technically inadequate or for whom the results are unknown (77) (LOE: C, Class IIb). Magnetic resonance imaging can be used to detect small myocardial scars and fibrosis that cannot be detected by other modalities, and thus cardiac MR imaging also may be useful in the baseline evaluation of this patient population (LOE: C, Class IIb). Noninvasive assessments of endothelial-cell function such as brachial artery flow–mediated dilation and structural assessments such as carotid IMT remain research tools at this time with uncertain prognostic significance. Follow-up evaluations every 3 to 5 years are currently recommended for KD patients during childhood, and continuing this periodic reassessment into adulthood seems prudent given the lack of information about the long-term consequences of KD (6). No medication is currently recommended for this patient population.

Adults with persistent or regressed aneurysms should be followed at regular intervals with functional and structural cardiovascular studies to determine the need for interventions (LOE: B, Class IIa). When considering CT, cardiac catheterization, or nuclear medicine studies, consideration should be given to the young age of these adults and the need for life-long, repeated assessments when weighing the risks versus the benefits of the radiation exposure. Although repeated MR imaging is without known risk, CT angiography provides a better assessment of coronary artery stenosis and calcification. Future improvements in software and cardiac cycle gaiting techniques will likely reduce the radiation exposure.

Optimal pharmacologic therapy of adult patients with persistent or regressed aneurysms has not been established. The AHA guidelines advocate the use of aspirin as an antiplatelet agent (3 to 5 mg/kg/day) in children with small-to-moderate aneurysms (<8 mm) (Table 1) (6). Ticlopidine or clopidogrel are also added to aspirin therapy at some centers (LOE: C, Class IIb). The current AHA guidelines suggest discontinuation of antiplatelet therapy

### Table 2

**Guidelines for Diagnosis and Management of Cardiovascular Sequelae in Kawasaki Disease as Outlined by the Japanese Circulation Society**

<table>
<thead>
<tr>
<th>Patient Risk Level</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>AHA risk level 1 and 2</td>
<td>Consider noninvasive testing every 3–4 yrs.* No medical therapy.</td>
</tr>
<tr>
<td>AHA risk level 3, 4, 5 without symptoms</td>
<td>Evaluate every 4–6 months with noninvasive testing and angiography every 2–3 years. Treat with low-dose aspirin.</td>
</tr>
<tr>
<td>AHA risk level 3, 4, 5 with symptoms</td>
<td>Evaluate every 3–4 months with noninvasive testing and angiography as needed. Treat with low-dose aspirin; other medications as dictated by the cardiovascular status.</td>
</tr>
</tbody>
</table>

Data from the Japanese Circulation Society (83). *Noninvasive testing: exercise testing, nuclear imaging studies, Holter monitor, transesophageal echocardiogram, magnetic resonance angiography, and computed tomography angiogram.

AHA — American Heart Association.
when the aneurysm regresses (LOE: C, Class IIa) (Table 1). The use of 3-hydroxy-3-methylglutaryl-coenzyme A reductase inhibitors, angiotensin-converting enzyme inhibitors, and angiotensin receptor blockers may all mobilize endothelial progenitor cells and promote vascular wall remodeling, although there are no data to support their use in the setting of KD (84,85).

For KD patients with persistent aneurysms, treatment regimens should be tailored on the basis of the size of the aneurysm and flow characteristics. In patients with giant aneurysms (≥8 mm) in which blood flow is sluggish and risk for thrombosis is high, systemic anticoagulation with warfarin with an international normalized ratio between 2.0 and 2.5 has been associated with improved survival in small series of KD patients from Canada and Japan (LOE: B, Class IIa) (86,87). There are no data to inform the management of adult patients with large aneurysms, but continuation of systemic anticoagulation seems reasonable. Addition of aspirin for platelet inhibition should also be considered (LOE: C, Class IIa).

On the basis of available data, pregnancy in KD patients with persistent aneurysms is not associated with increased risk (88–90). The authors of a nationwide survey (91) in Japan reported the outcome of 46 deliveries in 30 KD patients. Two-thirds of the deliveries were vaginal. Mothers with persistent giant aneurysms were given subcutaneous low-molecular-weight heparin and low-dose aspirin during the pregnancy. There were no cardiac complications, and the authors concluded that the mode of delivery should be dictated by obstetrical considerations and that coronary artery aneurysms alone were not an indication for Caesarean delivery (LOE: B, Class IIb).

Percutaneous catheter interventions may be necessary if narrowing of the vessel lumen leads to signs or symptoms of ischemia (LOE: B, Class IIa). Ishii et al. (92) have developed consensus guidelines for the use of these procedures in childhood. Percutaneous transluminal angioplasty has been complicated by the need for very high balloon pressures in highly calcified lesions leading to neoaneurysm formation (93). There is limited experience with revascularization procedures that use covered or drug-eluting stents in this patient population (94,95). Rotational atherectomy is the interventional procedure of choice for heavily calcified, stenotic lesions that are not amenable to percutaneous transluminal angioplasty, although current experience is limited (LOE: C, Class IIa) (96).

Surgical approaches that use both venous and arterial grafts have been used in this patient population (LOE: B, Class IIa). In a survey of 156 KD patients older than 12 years treated with internal thoracic artery grafts, the patency rate at 15 years was 91% (97). Use of arterial grafts in patients with large growth potential resulted in a much greater rate of graft patency over time as compared with venous grafts. Surgical intervention in young adults with aneurysms after KD has been reported (98). Surgical risks and graft survival did not differ from those of patients undergoing similar procedures for ischemia due to atherosclerotic disease.

Cardiac transplantation has been successfully performed for KD patients with end-stage cardiomyopathy, severe ventricular arrhythmias, and inoperable multivessel stenotic coronary artery disease (99). There has been no recurrence of KD or coronary artery aneurysms in the transplanted hearts, and all morbidity and mortality in this cohort have been related to rejection and complications of immunosuppression.

Conclusions

In the coming years, growing numbers of patients with cardiovascular complications after KD will present to internists and cardiologists specializing in adult patients for care. The vascular lesions in these patients differ from atherosclerosis and may include coronary artery aneurysms, calcification, and stenosis. These patients may also present with valvular incompetence due to scarring of the leaflets or progressive aortic root dilation and myxoid lesions, including diffuse fibrosis or focal scarring in regions of myocardial ischemia or infarct. The inflammatory insult associated with acute KD has the potential to affect all components of the cardiovascular system. Systematic study of adults with a history of KD in childhood is needed to define the natural history of this enigmatic disease.

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