QED? Not yet!

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Anomalous aortic origin of a coronary artery (AAOCA) from the opposite sinus of Valsalva is a very rare congenital anomaly with a documented potentially devastating risk of sudden cardiac death in young individuals. Starting from the acknowledgement of the entity as a pathology, rather than just a normal anatomic variation, the journey toward an evidence-based treatment has been expedited during the last few years.1,2 Risk stratification of patients continues to remain a challenge, despite the effort to create a detailed description of guidelines.2,3

In this issue of the Journal, Nees and colleagues4 present their retrospective review of 60 patients who underwent a repair exclusively for AAOCA in 2 affiliated institutions. The study, in line with other reports in the literature, offers enough reason to reconsider our surgical techniques to reduce morbidity in these patients.5,6 Even with a short median follow-up period of 1.6 years, Nees and colleagues emphasize 2 unfavorable surgical outcome features. First, a restenosis of the repaired coronary may occur even without positive stress testing findings, resulting in sudden cardiac death years after surgery. In this study, 2 patients required reoperation as a result of coronary stenosis at 3 months and 6 months after surgery. Second, aortic valve regurgitation has been observed to a mild and greater degree in 17% of the patients postoperatively within this short follow-up. Five patients underwent reoperation for aortic regurgitation. In addition to this, supra-aortic stenosis was encountered in 1 patient, and a resolution of the preoperative symptoms could only be documented in 64% of the preoperatively symptomatic patients. Nees and colleagues are to be congratulated for their detailed analysis of a large cohort of the patients and particularly for attracting attention to these extremely important outcome measures. With the inherent limitations of a retrospective, single-center study with a short follow-up but also with a high number of patients, the messages of this evaluation are of great value, highlighting the previously mentioned aspects and leading us to reconsider the impression of apparently “excellent” surgical outcomes in repair of AAOCA. We know that patients after AAOCA repair will remain at risk for lifetime. A clear understanding of the reasons for suboptimal outcomes in some cases is missing. Preoperative evaluations, surgical practices, and follow-up protocols that have significant variations between institutions do not make problem solving easy. As necessary for every surgical patient, each institution will require standardized protocols to allow a smooth multidisciplinary approach to the treatment and follow-up of patients with AAOCA. We are urged to aim strictly for an excellent surgical repair with the least morbidity for these mostly young patients, particularly with current challenges of risk assessment.

There is a consensus among experts that anatomic features should determine the type of repair. For instance, recent reports have suggested that an unroofing of a short intramural course might not suffice to relieve the risk of a recurrent obstruction.7 A reimplantation therefore may be necessary if the coronary ostium cannot be moved beyond the intercoronary pillar to its home sinus. It is likely that similar experiences will be reported in the future, addressing different morphologic features with alternate surgical techniques. Furthermore, once long-term data arrive, we will probably learn more about the fate of the intimal readaptation sutures (suture line) around the coronary ostium with regard to the tendency toward scarring and calcification. Intimal disruption and readaptation, especially in smaller ostial openings, may expose patients to a higher risk for adverse cardiac events and an eventual revascularization requirement in the longer term.

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Central Message

The surgical treatment of an anomalous aortic origin of a coronary artery requires refinement in 3 aspects: indication, techniques, and overall outcome.

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Less evidence exists in cases with an intramural course that is close to or even below the aortic valve commissure with regard to the technique of unroofing. Here, the definition of close remains unknown. Unroofing of an intramural coronary has the potential to distort the support of the aortic valve commissure with immediate proximity. Because the number of patients with new postoperative aortic regurgitation is not negligible in some studies, even without commissural takedown, the question has to be asked whether a resuspension of the aortic valve commissure needs to be performed in a standard manner regardless of the coronary proximity to the commissure. Preliminary data from my own center’s experience indicate that a resuspension can be accomplished with no increased surgical risk but with superior aortic valve function in the medium-term follow-up relative to those patients without resuspension. Another disproportionate complication seems the occurrence of the postcardiotomy syndrome with probable pericardial effusion. Surgical (pleuropericardial window) or medical (anti-inflammatory treatment) may reduce both the burden for our patients and rehospitalization rates.

We are still distant from adequately answering the question of whether today’s surgical indications and repair techniques for AAOCA are effective to prevent sudden cardiac death with acceptable surgical morbidity and sufficient scientific evidence. It thus will not be sufficient to declare the surgery mere to be a safe approach in future to proceed with current practices. Prospective, multicenter studies with longer follow-up periods and standardized, detailed protocols throughout the management of this entity are more essential than ever.

References