The role of devices in the closure of atrial septal defects in the oval fossa

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Increased knowledge, and modern developments in technology, have given birth to several techniques for interventional closure of atrial septal defects. During the last few years, at least five different devices have been the subject of clinical trials. These are various generations of the Sideris' buttoned device; the ASDOS [250]; the self-centring AngelWings [around 325]; CardioSEAL, the modified Clamshell device, [approaching 500]; and the nitinol Amplatzer plug [over 500]. The numbers in brackets refer, according to their manufacturers, to the world-wide implants in atrial septal defects or the oval foramen as of February 1998 for the four less-published devices. These numbers may not only mirror the general acceptance by cardiologists, but also the clinical compliance of the different devices. Only the buttoned device is well documented in the literature, 1-2 although certainly a great part of the literature on the clamshell device 3 might be considered relevant also for CardioSEAL. The other devices are presented in few, smaller series, 4-5 or by their complications, 6-7 representing serious setbacks in the implementation of a new method. Much of the present information is preliminary, and represents personal communications between investigators. It is now increasingly important to publish results with the use of different devices. I welcome, therefore, the articles in this issue of Cardiology in the Young. 8-9

The optimal device should be easy to handle. Implantation should be feasible through a small introducer sheath. The device, when inserted, should be safe, retrievable, compatible with most atrial defects, and produce high rates of closure and low complication rates. Serious complications should not occur, and closure with the device should never induce new lesions within the heart. Unfortunately, several devices do not presently satisfy these criteria. Four of the devices use the same general concept: aiming to close the defect with a membrane on one or both sides of the atrial septum. Stability is achieved through a greater or lesser pressure of arms, constructed in different ways and arranged radially or circumferentially when open against the atrial wall. The Amplatzer is different. It is circular. It stents the hole, and sits like a peg within it, achieving stability by the pressure of the stenting component within the hole against its margins. This very concept may be its main advantage, resulting in high rates of closure and giving few complications. No rigid arms point towards neighbouring structures, avoiding any possibility of injury. The effect of the radial pressure against the atrial septum, and the potential annular distortion, however, is unknown. The somewhat bulky design probably does not matter, but the steel ends which protrude to either side may be a matter of concern regarding complete endothelialization.

In this issue, we publish articles from Melbourne 8 and Toronto 9 respectively, on interventional closure of atrial septal defects. They describe the use of two different devices, one from the stance of the performer, the other from that of the morphologist. Both contribute important knowledge. In their series of patients, Wilkinson and Goh 8 report complete closure of around 90%, 1 month following implantation of the Amplatzer device. Their experience with displacement of one device, obviously the result of underestimation of the size of the defect, emphasizes that correct sizing is particularly critical for this device, demanding a specially designed sizing catheter.
The careful selection of patients for closure will no doubt influence both failures and results. Three-dimensional echocardiography, so beautifully displayed in the article by Maeno et al,9 represents a means to better understanding of the specific anatomy of the septal deficiency, consequently improving the judgement of suitability for interventional closure. The article also discusses results subsequent to implantation, including malpositioning. Following the recommended approach will probably increase the rate of success, but we must realise that there are a number of possible sources of error in the present technology, and the authors themselves accurately describe the difficulty in differentiating between a true septal defect and echo drop-out. Quite a number of paediatric cardiologists will consider it difficult to use transoesophageal echo routinely in children, and some will wonder how the time required can be managed in the daily routine. Needless to say, interventions are preferable to the far more invasive cardiosurgical approach, provided of course that risks and results are comparable. One does not need clairvoyant capabilities to predict that closure of atrial septal defects has a low rate of complications, very little residual shunting, and an almost negligible mortality. Interventions to close atrial septal defects, self-evidently far less invasive than surgery, will still need to provide results comparable to such surgery if they are to compete. Our rates of complication, therefore, must be low even in the 'learning curve'. Serious complications or lesions must not occur but, alas, they have. Some devices certainly will show superior results compared to others with regards to closure, failures, and complications. The evidence from the Melbourne team,8 along with the findings of Masura et al,3 indicate that the Amplatzer might be such a device. It is unlikely that all five devices will survive. Maybe the ultimate device has not yet been made. The devices now leave the era of investigational work and some will become freely available. It will be increasingly important, therefore, to maintain a sound balance between expectations and reality. We must not turn a blind eye to unforeseen errors, complications and poor results. Similarly, we must not perform interventions at any cost. There are certainly moments when ethics should prevail over techniques for the benefit of the patient.

References