Pre-Procedural Transesophageal Echocardiography Underestimates Atrial Septal Defect Size Compared to the Balloon Sizing Technique

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ABSTRACT

BACKGROUND: Secundum-type atrial septal defects (ASD) are increasingly being treated with percutaneous placement of occluding devices. Accurate sizing of the ASD is mandatory for subsequent optimal selection of the device. We sought to compare the two most commonly used methods, 2-dimensional transesophageal echocardiography (2D-TEE) and stretched-balloon sizing.

METHODS: Sixteen patients (8 men and 8 women) aged 53.8±9.8 years with ASD were scheduled for implantation of an Amplatzer septal occluder device. The procedure was performed with the use of local only anesthesia via the right femoral vein with fluoroscopic guidance alone and without the intra-procedural use of 2D-TEE. The size of the defect was measured with 2D-TEE prior to the procedure. In the catheterization laboratory the ASD size was measured again with the insertion of a sizing balloon inflated with diluted contrast agent until a waist appeared. Waist dimensions were measured using cineangiographic quantitation (QCA) software.

RESULTS: All ASDs were successfully closed with Amplatzer occluders. A significant correlation was found between echocardiographic and fluoroscopic measurements of the ASD size (r=0.863, p=0.000). However, most of the echocardiographic measurements underestimated ASD diameters compared with conventional QCA balloon sizing by a mean of 3 mm (16.46±5.93 mm vs 19.83±6.3 mm, t= 4.18, p= 0.001). The Bland-Altman analysis evidenced a lack of agreement between echocardiographic and balloon sizing with 95% limits of agreement between 2.9 and -9.7 mm.

CONCLUSIONS: Atrial septal defects can be treated safely and effectively with the percutaneous placement of an Amplatzer septal occluder device via a simple percutaneous technique with use of local anesthesia and fluoroscopy alone without a need for intra-procedural 2D-TEE. Although the use of pre-procedural 2D-TEE has been a common practice for establishing the diagnosis and offering an initial assessment of the size of the defect, it largely underestimates the latter. The single-plane fluoroscopic balloon sizing performed during the procedure seems indispensable for choosing an occluding device of appropriate size.
Surgical closure of atrial septal defects (ASD) has been performed for decades and has been associated with very low morbidity and mortality rates. On the other hand, the placement of closure devices percutaneously via the femoral vein has also been used as an easy alternative approach for several years. Indeed, nowadays, ASDs are increasingly being treated by percutaneous techniques and the placement of a closure device is now considered as the treatment of choice with excellent sealing rates of the defect. Accurate sizing of the ASD is mandatory for subsequent optimal selection of the device and can be done either by echocardiographic measurements (transthoracic, transesophageal or intracardiac) or with the use of sizing balloons in the catheterization laboratory. The stretched balloon measurement of the ASD diameter through fluoroscopy is considered the gold standard among them. Nevertheless, even though balloon sizing is the method preferred by most, an initial assessment of the ASD with echocardiography is usually performed in every candidate for percutaneous closure of the defect prior to the procedure. There have been numerous discussions about the potential of relying on echocardiography alone since this would reduce both the time and the cost of the procedure, but there has been no general consensus on this issue. We sought to evaluate two-dimensional transesophageal echocardiography (2D-TEE) multi-plane measurement in comparison to single-plane fluoroscopic balloon sizing of the ASD in patients undergoing this procedure in our center.

**Patients and Methods**

**Study Population**

Between July 2006 and June 2010, 16 patients with solitary secundum-type ASD had a cardiac catheterization for closure of their defect with an Amplatzer Septal Occluder (AGA Medical Corp., Plymouth, MN) in our department. The patients’ characteristics are shown in Table 1. Most of them underwent the procedure due to a left-to-right shunt with a Qp/Qs ratio of ≥1.5 (n=13). In all subjects, measurements of ASD diameters by TEE before closure were compared with fluoroscopic balloon sizing. For the procedure an informed written consent was obtained from each patient. The study

<table>
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<th>n</th>
<th>Age (years)</th>
<th>Gender</th>
<th>Symptoms</th>
<th>Indication for the procedure</th>
<th>Echo Size (mm)</th>
<th>Balloon Size (mm)</th>
<th>Closure Device</th>
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was approved by the Ethics Committee of our institution.

**TRANSESOPHAGEAL ECHOCARDIOGRAPHY**

Standard 2D-TEE was performed with a Philips iE 33 ultrasound unit (Philips Medical Systems, Bothell, Washington, USA) to diagnose secundum type ASDs, exclude additional congenital malformations and ascertain that the rims were sufficient for a closure device placement. Standard ASD measurements were carried out in the bicaval view at 60-70 degrees and in basal short axis view. The maximum diameter was taken into account.

**FLUOROSCOPIC BALLOON SIZING AND PERCUTANEOUS DEVICE CLOSURE TECHNIQUE**

In our laboratory, the procedure of percutaneous ASD closure is routinely performed with use of local only anesthesia via the right femoral vein with fluoroscopy guidance alone and without the intra-procedural use of TEE. All patients received aspirin and clopidogrel for at least 3 days prior to the procedure, and prophylactic antibiotic is administered intravenously one hour prior to the scheduled procedure.

Via a right femoral venous approach, the patients were catheterized and the ASD size was measured with standard sizing balloons with diameters between 20 and 34 mm in the left anterior oblique view with slight cranial angulation (30 degrees left anterior oblique/20 degrees cranial). Sizing balloons were centered in the ASD, aiming to position the distal half of the balloon in the left atrium and the proximal half in the right atrium. The balloon was inflated with diluted contrast agent until a waist appeared. Waist dimensions were measured using cineangiographic quantitation (QCA) software. The procedure was repeated up to two times depending on how well the waist was visualized. After device selection based on balloon measurement of ASD size, a suitable Amplatzer septal occluder (AGA Medical Corp., Plymouth, MN) was selected oversized by 2-4 mm and deployed via the special delivery system. The day following the procedure 2D-TEE was repeated to check for device position and any residual shunt or other peri-procedural complications. All patients were discharged home at 48 hours with aspirin and clopidogrel for 6 moths. This regimen was adopted taking into consideration its safety and efficacy from the wide experience from coronary artery disease. At 6 months all patients underwent a repeat 2D-TEE.

**STATISTICAL ANALYSES**

Data are expressed as mean ±standard deviation. Agreement between methods was evaluated using paired *t* test and Pearson correlation. Calculations were based on a type I error level of 5%. Data were analyzed with SPSS 18 for Windows (SPSS Inc., Chicago, IL). Moreover, since a high correlation does not automatically imply that there is good agreement between two methods we performed also a Bland-Altman analysis to calculate the limits of agreement with the use of the statistical package MedCalc Software (Mariakerke, Belgium).

**RESULTS**

All ASDs were successfully closed with use of Amplatzer occluders. One patient developed a transient paroxysm of atrial fibrillation with no other periprocedural complications noted. At 2 weeks post-procedurally the same patient developed a brief episode of paroxysmal nocturnal dyspnea managed with diuretic without further recurrences. At 6 months after the implantation one patient had a large residual shunt and at attempting to place a new closure device, the device migrated into the pulmonary artery and was successfully recaptured with a percutaneous technique. The patient was subsequently scheduled and underwent surgery 3 months later with successful closure of the ASD. Small residual shunting was noted in another patient at the 6-month follow-up TEE, who was managed conservatively. All other patients reported no symptoms at a mean follow-up of 30.12 ± 4.86 months.

With regards to the measurements of the ASD size obtained with the TEE and the stretched balloon technique, a significant correlation was found between echocardiographic and fluoroscopic measurements (*r*=0.863, *p*=0.000) (Figure 1). However, most of the echocardiographic measurements underestimated ASD diameters compared with conventional QCA balloon sizing (16.46±5.93 mm vs 19.83 ±6.3 mm, *t*= 4.18, *p*= 0.001) (Figure 2).

Moreover, the limits of agreement between the two methods calculated with the Bland-Altman analysis, were found be-

![FIGURE 1. Correlation between fluoroscopic (QCA) and echocardiographic (ECHO) measurements.](image-url)
That means that 2D-TEE measurement may be 2.9 mm above or 9.7 mm below the number obtained from Balloon sizing. This is a fairly wide difference suggesting a lack of agreement between the 2 methods (Figure 3).

**DISCUSSION**

The present study showed that 2D-TEE systematically underestimates stretched ASD size by at least 3 mm. This is reasonable since balloon-sizing technique usually stretches the true anatomical septal shunt measured by TEE. Two-dimensional TEE provides information about the structure of ASD but only one cross-sectional image at a time so it can underestimate the ASD size if the probe is not aligned with the longest diameter of the defect. In addition, ASD changes in shape and size throughout the cardiac cycle and also membranous tissue usually rounds the defect and can easily be stretched if a balloon stretches leading to a significant increase in the true diameter as assessed by the balloon method. Taking this assumption into consideration a study Carcagni et al. introduced a new echocardiographic diameter. Apart from the classic anatomic diameter measured by 2D-TEE, they also measured the procedural ASD diameter, using
as borders of the true defect the points where the thickness of the rims was 2.5 mm. This value was chosen from an in vitro observation of the thickness of the rim at the time the sizing balloon is inflated and the stretched balloon diameter is measured. In this study which included 40 patients a good correlation between procedural ASD diameter and stretched balloon diameter was found ($r=0.99$). The authors suggested that procedural diameter is easily measured, applicable to all echocardiographic techniques without additional cost.

We used the static balloon technique in which a sizing balloon catheter is advanced over the wire across the defect, and inflated with dilute contrast medium until the balloon has a definite waist that is visible on fluoroscopy. The balloon indentation can be measured by QCA as the stretched-balloon-diameter by fluoroscopy. The same stretched-diameter can be measured by 2D-TEE only if the examination is performed simultaneously with cardiac catheterization. The usual practice is to select a device 2–4 mm larger than the stretched ASD diameter. Devices smaller than the stretched ASD size cannot be deployed safely. However, issues about accuracy of balloon sizing emerge when the balloon is oblique through the defect or distends the rims.

Others have suggested that three-dimensional trans-esophageal echocardiography (3-D TEE) can offer a better visualization of the anatomy of the ASD, can give accurate assessment of the true anatomical size, can assess tissue surrounding the defect, and can follow its size during the cardiac cycle, providing a far better image of the defect. Even more it can further assist in the measurement during the stretching of the balloon. In the study of Maeno et al both 2-D and the 3-D TEE measurements are lower than the stretching balloon values, with the 3-D TEE measurements having the minimal mean difference between the two. In a previous study no differences were observed between 3-D and stretching balloon measurements. Limitation of the 3-D TEE measurement technique can be considered the time consuming process and the possibility of not capturing the region of interest during data acquisition, especially during the measurement of the stretched ASD when inflation of the balloon occurs. In addition, the need for general anesthesia with or without intubation and the prolonged radiation exposure should always be taken into consideration.

Intracardiac echocardiography (ICE) has been used for guiding the sizing balloon and the closure devices during the procedures. A recent study evaluated the single-plane balloon stretching sizing by ICE compared with the gold standard of fluoroscopy measurement. It was shown that ICE systematically underestimated the stretching balloon diameter by less than 1 mm. Use of ICE may limit the radiation, the time of the procedure and the need for sedation and can always guide the sizing balloon in the center of the defect for offering more accurate values. The major limitations seem to be the single-plane view of the measurement allowing frequent underestimation of the size, the lack of color Doppler capability and the high cost of the catheters. Another study which compared 3-D TEE both with 2-D TEE and ICE suggested that the former is more likely to give larger long-axis dimension and smaller short-axis dimension when compared with both the 2-D TEE and the ICE.

Finally, phase-contrast magnetic resonance imaging offers an alternative way of noninvasive method to ASD sizing, especially in children with inconclusive echocardiographic results. Complications of the balloon sizing technique are rare in general. Oversizing of the balloon, repeated inflation or employment of the pull-through technique instead of the static one may end to tear of the atrium septum and lead to device embolization or entrapment. It has been suggested that inflation of the sizing balloon until flow through the shunt is diminished results in potentially fewer complications. Limitations of the study This study was not a randomized controlled trial. It was rather our experience in our ASD closure program based on a small series of patients. This represents a major limitation in addition to the fact that the operators performed real time calculations and not off-line analysis. Nevertheless, they were blinded to each other’s measurements. Absolute concentricity was not always achievable by balloon inflation and this represents another limitation since a slight underestimation of the balloon sizing might have occurred in some cases. On the other hand, over-stretching of the balloon and “over-sizing” of the defect might have not been avoided in some other patients. However, almost all patients received a device of appropriate diameter since only one distal migration and no aortic erosions occurred. The absence of intra-procedural TEE that might have helped us to reduce the complication
rate (2 residual shunts) should also be noted. Finally, we should emphasize that although a calculation taking into account the thickness of the rim of the ASD might have led to a better correlation with the balloon measurements, applying it to all ASDs was not always feasible probably due to individual anatomical variability.

**CONCLUSIONS**

Atrial septal defects have been treated in our cases safely and effectively with the percutaneous placement of an Amplatzer septal occluder device via a simple percutaneous technique with use of local anesthesia and fluoroscopy alone without a need for intra-procedural 2-D TEE. Although the use of pre-procedural 2D-TEE has been a common practice for establishing the diagnosis and offering an initial assessment of the size of the defect, it largely underestimates the latter. The single-plane fluoroscopic balloon sizing performed during the procedure seems indispensable for choosing an occluding device of appropriate size.

**REFERENCES**