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**Left atrial appendage occlusion: A better alternative to anticoagulation?**

Akin I *et al*. LAA occlusion

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**Abstract**

Non-valvular atrial fibrillation is associated with a significantly increased risk of embolic stroke due to blood clot forming predominantly in the left atrial appendage (LAA). Preventive measures to avoid embolic events are permanent administration of anticoagulants or surgical closure of the LAA. Various clinical trials provide evidence about safety, effectiveness and therapeutic success of LAA occlusion using various cardiac occluder devices. The use of such implants for interventional closure of the LAA is likely to become a valuable alternative for stroke prevention, especially in patients with contraindication for oral anticoagulantion as safety, clinical benefit and cost-effectiveness of LAA occlusion has recently been demonstrated.

**Key words:** Left atrial appendage; Occlude; Stroke; Thrombus

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**Core tip:** Non-valvular atrial fibrillation is associated with increased risk of embolic stroke. To date, risk-based anticoagulation is the cornerstore to avoid this. However, several patients have got absolute or relative contraindication to this and thus are undertreated. For these patient population the implantation of a local left atrial appendage occluder might be an alternative.

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**INTRODUCTION**

The left atrial appendage (LAA) is an external protrusion of the left atrium located next to the pulmonary trunk[1,2]. Compelling evidence points to the LAA as the primary origin of thrombus formation particular in the presence of non-valvular atrial fibrillation (AF); since the major risk of non-valvular AF to suffer from ischemic stroke, the LAA has drawn much attention in the context of stroke prevention[3-5] considering missing awareness and unrecognized AF prior to strokes[6]. Thus, the current approach for stroke prevention in patients with non-valvular AF of risk-adjusted prevention *via* oral anticoagulants (OAC) or antiplatelet agents[7,8] may be challenged by elective LAA occlusion in selective patients[3,4,9-12]. In the past, physical LAA closure required either surgical excision or exclusion by suture or stapler[10,13]. With the introduction of cardiac occluder devices open surgery is not required any more and redundant for this purpose[14-16]. This review summarizes current knowledge of LAA occlusion as an emerging alternative to chronic OAC therapy for non-valvular AF patients at risk for embolic strokes.

**ANATOMY OF THE LAA**

The function of the LAA is not fully understood but it has been linked to secretion of the hormone “atrial natriuretic factor” (ANF) and, hence, could be involved in the regulation and homoeostatic control of water, salt and fat[17,18]. Regardless, the anatomy of the LAA is highly diverse and was classified into four different morphological types; “chicken wing” was the most frequently identified type (48%) followed by “cactus” (30%), “windsock” (19%), and “cauliflower” (3%)[19]. The “chicken wing” type has a dominant lobe, which may have secondary lobes or twigs, and is bent in the proximal or middle part or even folds back on itself at some distance from the orifice. The “cactus” type has a dominant central lobe with secondary lobes extending in both superior and inferior directions, whereas the primary structure of the “windsock” type is a dominant lobe with variation in the location and number of secondary or even tertiary lobes. Lastly, the “cauliflower” type has a short overall length with more complex internal characteristics, lacks a dominant lobe but has variable number of lobes and a more irregular shape of the LAA orifice. Previous studies indicate that the “chicken wing” type poses the lowest risk for embolism in contrast to the “cauliflower” type, which, notably, exhibits the highest degree of structural complexity[20]. However, due to the complex anatomy of the LAA, it is difficult to correctly assess length, branches and courses, as well as thrombus formation by transesophageal echocardiography (TEE) and it was demonstrated that the outcome of this visualization is dependent on the selection of the imaging plane[21].

**LIMITATIONS OF ORAL ANTICOAGULANT THERAPY**

The current gold standard for stroke prevention in patients with non-valvular AF is the oral administration of anticoagulants to reduce the risk of thrombus formation and prevent any embolic events[7,8]. Chronic anticoagulation is carried out by traditional and novel oral anticoagulants (NOACs) also called directly acting oral anticoagulants (DOACs). Traditional anticoagulants include heparins and coumarins (vitamin K antagonists) of which warfarin is the most common. NOACs are inhibitors of coagulation factors such as factors IIa (*e.g.*,dabigatran) or factor Xa (*e.g.*, rivaroxaban, apixaban and edoxaban)[22-25]. However, chronic OAC therapy is not recommended if contraindications are present or potential interference with other therapies. Moreover, difficulties to adjust treatment, dietary restriction, low compliance or even refusal of the patient to follow treatment protocol are considered contraindication for OAC[10,22-26]. Therefore, alternative strategies for stroke prevention in patients with AF are required.

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**APPROACHES FOR CLOSURE OF THE LAA**

There are two fundamental approaches beyond anticoagulation to avoid emboli in patients with non-valvular AF, *e.g.*, surgical excision of the LAA or exclusion by suture line, stapler or cardiac plug[10,13]. Before introduction of cardiac occluder for LAA closure, surgical excision was the superior method while LAA closure requires either suture line[10,13]. However, a surgical excision is not risk free and may cause bleeding[27]. Nonetheless, surgical excision is still an option mostly in conjunction with other cardiac surgery[10,27]. In recent years, an alternative approach for LAA closure was established by sealing the orifice of the LAA with an occluder. Such a LAA occlusion was performed 2001 for the very first time using the PLAATO system which has been taken off market[28,29]. The current generation of occluders are the Amplatzer™ Cardiac Plug (ACP), Amplatzer Amulet™ from St. Jude Medical and the Watchman™ device from Boston Scientific[29,30]. In addition, a small number of novel devices have been mentioned and applied in the last few years such as the WaveCrest™ device and the Lariat™ device[31,32] (Table 1). The ACP-originally used for closure of atrial septal defects[33]- consists of a self-expanding flexible nitinol mesh with a distal lobe filled with polyester and is equipped with fixation barbs to adhere of the LAA. The distal lobe is connected *via* a small waist to a proximal disc sealing the orifice of the LAA[34]. Similar to the ACP, the Watchman™ device consists of a self-expanding nitinol mesh with fixation barbs and a polyester coating covering the surface facing the left atrium[35].

Implantation of both Watchman™ device and ACP can be performed under local anesthesia and is introduced *via* catheter through the femoral vein by transseptal passage[33,36-39]. TEE guiding or intracardiac echocardiography (ICE) during implantation procedure is used to rule out intracardiac thrombus and to facilitate transseptal puncture. After transseptal puncture heparin is administered to achieve an active clotting time of > 250 s. The LAA is fluoroscopically illustrated in at least 2 standard angulations (RAO 30°, RAO 30°/10° caudal) and sized by TEE measurements and cine angiography. Device size will be chosen according to manufacturer’s recommendations 20% larger than the landing zone (measured from left circumflex coronary artery to the ridge delineating the LAA from the left upper pulmonary vein). Optimally, the device should not protrude more than 5 mm beyond the LAA ostium and should cover the entire ostium with no or minimal (less than 5 mm by colour Doppler) residual flow and a compression grade of 8%-30%. After device releasing and sheath removal the puncture site is dealed with a Z-suture or with a pressure band. Periprocedural anticoagulation is managed by heparine or bivalirudine[14,28,33] (Figure 1). The implanted device becomes initially coated by fibrin and subsequently covered by endothelial cells forming an endocardial lining, which consequently excludes the LAA from circulating blood[40]. In order to allow the process of endothelialization patients have to take warfarin after the intervention for at least 45 d. Warfarin is then replaced by clopidogrel and aspirin for half a year, while aspirin administration is continued life-long[15]. According to newer data a dual antiplatelet therapy is more efficacious than the use of anticoagulants[37,41,42]. In addition to the general risks of catheter-based interventions including air or blood embolism, an incomplete closure of the orifice, pericardial perforation, dislodgement of the implant or the formation of blood clots on the surface of the device leading to prolonged OAC treatment may occur[14,28,33,43]. Finally, since ANF is secreted in the LAA, the LAA closure could interfere with thirst regulation and water retention in the patient, but this theoretical concept has been scarcely investigated so far[17,18].

**STATUS QUO OF LAA OCCLUSION**

For several years great effort has been devoted to the use of cardiac plugs in the prevention of AF-related strokes. A number of studies have been published about the PLAATO system including a five-year follow-up study[28,44]. In addition to the PLAATO system, the ACP system which has been used for closure of atrial septal defects for more than 20 years and its features, design and applicability are very well studied and has been reported to be successful for LAA closure[33,34]. It is worth mentioning that a large randomized clinical trial to study the ACP for LAA occlusion was recently halted, probably due to the approval of a competitive product (*i.e.*, Watchman device) by the United States Food and Drug Administration (FDA)[45]. The successful use of the Watchman device for LAA occlusion has been shown in two large, prospective randomized clinical trials-the PROTECT-AF and the PREVAIL study-in which this implant was compared to chronic OAC therapy using warfarin[14,15]. Over five years, the PROTECT-AF study examined about 800 patients, of which 463 had received a Watchman™ device and 244 were left on warfarin[14] and demonstrated that the implant is non-inferior to OAC therapy in patients with non-valvular AF being eligible for OAC. In comparison to the OAC treatment group, the incidence of an embolic event was reduced by approximately 30% in the implant group; the overall mortality showed a reduction of the same magnitude. While the per-protocol analysis was in favor of LAA occlusion, the intention-to-treat results were neutral. Furthermore, the safety data also favored warfarin over LAA occlusion, but this was explained by the learning curve phenomenon. The highest risk from LAA occlusion arises from complications associated with the one-time interventional treatment, while the risks from chronic OAC therapy accumulates during time especially with increasing age of the patient. Recently, a 45-months follow-up of the PROTECT-AF study demonstrated that LAA occlusion is not only as efficient as warfarin treatment but even superior in term of stroke and cardiovascular mortality[46]. Further, the safety data showed a considerable reduction in the risk for complications during intervention, likely due to increasing experience of the surgeons. However, the FDA criticized the patient selection and raised questions about the safety of LAA occlusion[47]. To address these limitations, a conﬁrmatory randomized trial (PRE- VAIL) comparing LAA occlusion with the Watchman device to warfarin, which mandated inclusion of new operators, slight modiﬁcations in inclusion criteria, and elimination of clopidogrel 7 d before implant. In PREVAIL more than 400 patients were randomized to either warfarin (*n* = 138) or occluder device (*n* = 269)[15]. This study showed that LAA occlusion achieved non-inferiority in stroke prevention compared to warfarin; the difference between both groups was low and not significant. Most importantly, the number of early safety events (*e.g.*, pericardial effusions) was significantly reduced compared to the PROTECT-AF study and, hence, satisfied the predefined goal. Therefore, the PREVAIL study addressed the concerns rose by the FDA and demonstrated the safety of this intervention for LAA occlusion[15]. As a result of this study the Watchman™ device was approved by the FDA in 2015[48]. Results from the real-world EWOLUTION registry consisting of 1021 patients being implanted with the Watchman device revealed a procedural success rate of 98.5%[49]. During 30-d follow-up 28 subjects experienced serious adverse events with an overall 30-d mortality rate of 0.7%. Serious procedure related complication rates, defined as stroke, pericardial effusion, device embolism and death, were present in 8.7% in PROTECT-AF, 4.1% in CAP registry, 4.2% in PREVAIL and 2.7% in EWOLUTION. However, the average CHADS2 score of 2.8 and CHA2DS2-VASc score of 4.5 in EWOLUTION indicate a relatively higher risk of stroke than either the PROTECT AF (average CHADS2 of 2.2 and CHA2DS2-VASc of 3. 4) or PREVAIL (CHADS2 score of 2. 6 and CHA2DS2-VASc of 4.0) studies. In addition, 40% of EWOLUTION subjects had a HAS-BLED score of ≥ 3, compared with only 20% of PROTECT AF subjects and 30% of PREVAIL subjects (Table 2). Similar results were obtained in a registry with the ACP device[50] and a large meta-analysis[51] including 2406 patients from the PROTECT AF, PREVAIL, CAP I and CAP II registries with a mean follow-up of 2.69 years. Patients receiving LAA occlusion with the Watchman device had signiﬁcantly fewer hemorrhagic strokes [0.15 *vs* 0.96 events/100 patient-years (PY); hazard ratio (HR): 0.22; *P* < 0.004], cardiovascular/unexplained death (1.1 *vs* 2.3 events/100 PY; HR: 0.48; *P* < 0.006), and nonprocedural bleeding (6.0% *vs* 11.3%; HR: 0.51; *P* < 0.006) compared with warfarin. All-cause stroke or systemic embolism was similar between both strategies (1.75 *vs* 1.87 events/100 PY; HR: 1.02; 95%CI: 0.62 to 1.7; *P* = 0.94). There were more ischemic strokes in the device group (1.6 *vs* 0.9 and 0.2 *vs* 1.0 events/100 PY; HR: 1.95 and 0.22, respectively; *P* = 0.05 and 0.004, respectively) [51].

**COST-EFFECTIVENESS**

An analysis of Panikker *et al*[16] on 110 patients being suitable and unsuitable for long-term OAC and outcome analysis from the PROTECT-AF trial and registry study compared warfarin, dabigatran, rivaroxaban, apxaban, aspirin and no treatment using a network meta-analysis. They revealed that stroke and bleeding rates were significantly lower than PROTECT-AF results. Additionally, LAA occlusion achieved cost parity between 4.9 years *vs* dabigatran 110 mg and 8.4 years versus warfarin and at 10 years, occlusion was cost-saving against all therapies. Similarly, another analysis evaluated the cost effectiveness in patients suffering from AF and absolute contraindication for OAC[52]. For this purpose the ASAP study evaluating the Watchman device, the ACTIVE-A trial evaluating aspirin and clopidogrel and the AVERROES trial evaluating apixaban were compared in a cost-effectiveness analysis. At 5 years, LAA occlusion was cost effective compared with aspirin with an incremental cost-effectiveness ratio of 16971 Euro. As compared with apixaban, it was also cost-effective at 7 years with an incremental cost-effectiveness ratio of 9040 Euro. Apart from the population having an absolute contraindication for OAC similar analysis were performed for patient being eligible for OAC, where LAA occlusion was cost-effective at 7 years and novel oral anticoagulation was cost-effective at 16 years. However, LAA occlusion was superior to novel oral anticoagulation by year 5 and to warfarin by year 10 with respect to cost-effectiveness and cost saving for stroke prevention[53].

**FUTURE CONSIDERATIONS**

In 2012, the European Society of Cardiology (ESC) released a focused update of the guidelines for the management of AF[8]. Interestingly, the ESC also commented on LAA occlusion for prevention of stroke (class IIb recommendation, level of evidence B). Due to insufficient amount of data demonstrating efficacy and safety, the ESC did not recommend an LAA occlusion at this point as a routinely alternative therapy to replace chronic OAC therapy in order to reduce AF-related stroke risk, but recommended to consider this approach for patients with an increased for stroke and contraindications for OAC treatment[8]. However, the references cited as evidence for the recommendation are the PROTECT AF study and the CAP registry. Importantly, neither of these studies included patients who had contraindications to long-term anticoagulation, and both enrolled a majority of patients with relatively low estimated stroke risk. A growing body of evidence suggests that LAA occlusion with the Watchman™ device is an important alternative for OAC therapy using warfarin, yet, a perspective randomized study comparing LAA occlusion to the NOAC is still missing and therefore a final conclusion about the general applicability of LAA occlusion using the Watchman implant cannot be drawn at this point[49,50,54]. Additionally, there are not enough data analyzing different patient cohorts (*e.g.*, sex, age, race, renal insufficiency) as there are data revealing a direct correlation between elevated adiponectin levels and the degree of left atrial blood stasis in men but not in women, and there are more extensive left atrial remodeling and deterioration in LAA function in women than in men[55-57]. There are some subanalyses revealing higher bleeding events in patients older than 75 years after LAA Occluder implanattaionas compared to younger ones (4.4% *vs* 1.4%), as well as in males as compared to females (3.0% *vs* 1.8%)[58]. In accordance with the latest recommendations from the ESC, LAA occlusion should definitely be considered if complications with OAC therapy arise or a high bleeding risk exist, regardless if the patient is treated with traditional or novel anticoagulants. This therapeutic approach is even more justified if the patient undergoing chronic OAC therapy suffers a stroke. Under this circumstance and under the light of recent studies about the safety and efficacy of LAA occlusion, this interventional treatment could be a better choice and advisable for this with a CHA2DS2Vasc Score ≥ 2 (Table 3). Summing up the current data, LAA occlusion is a very promising treatment to prevent AF-related strokes due to its safety, cost-effectiveness and therapeutic success.

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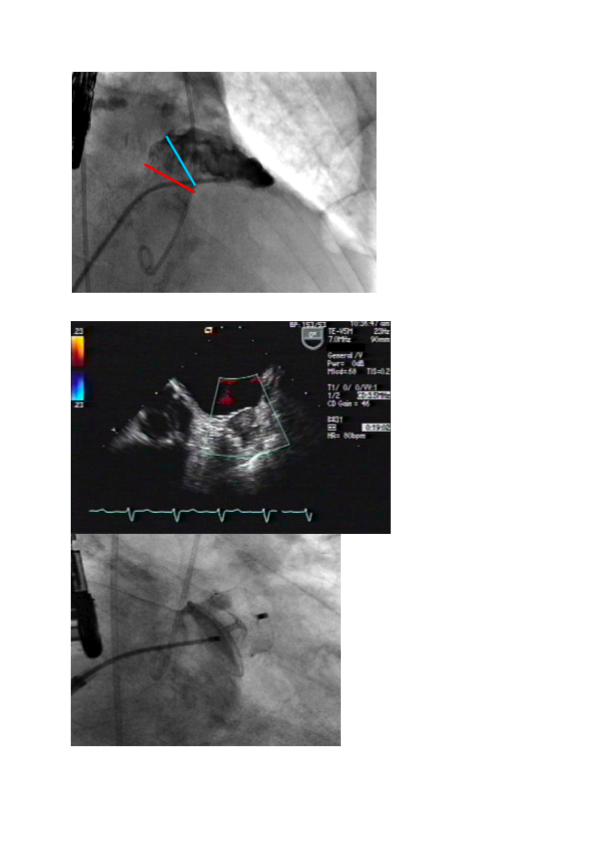
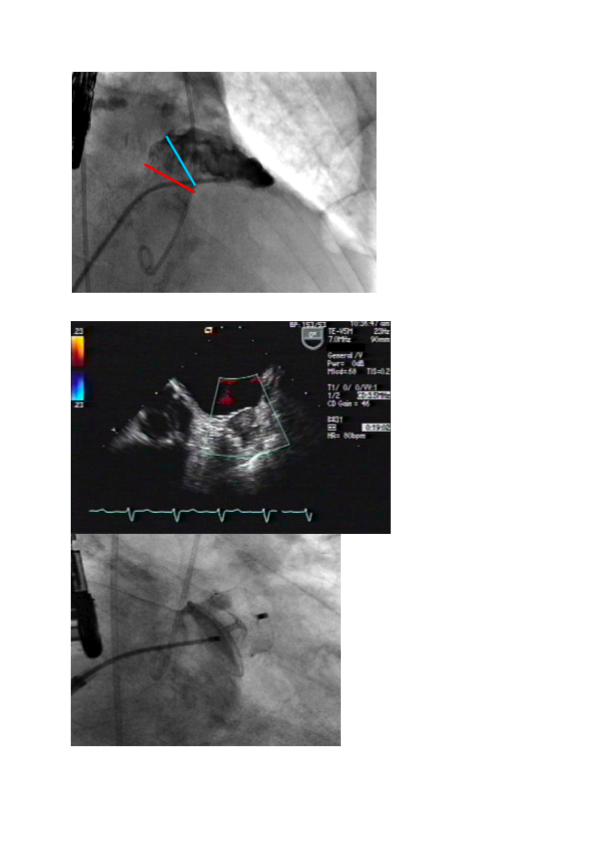
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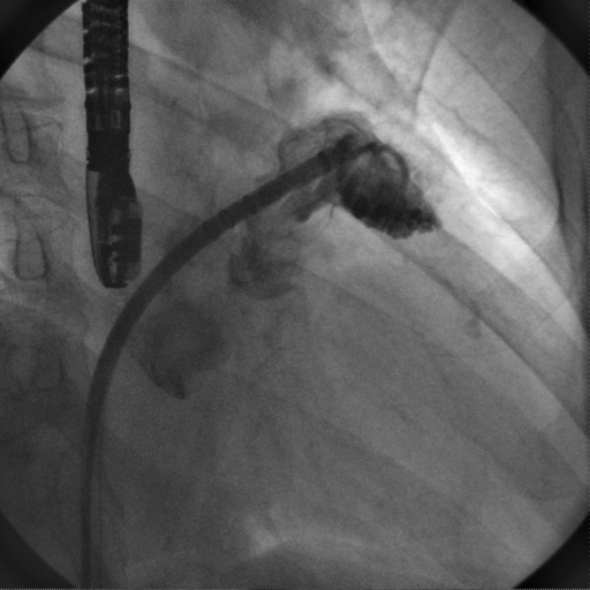
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**Figure 1.** Measurement of left atrial appendage (left), imlantation of an Amulet device (middle) and postinterventional transesophageal echocardiogram revealing good sealing without any leak (right)



**Figure 1 Measurement of left atrial appendage (left), imlantation of an Amulet device (middle) and postinterventional transesophageal echocardiogram revealing good sealing without any leak (right).**

**Table 1** **Different endocardial left atrial appendage occlusion devices[31]**

|  |  |  |
| --- | --- | --- |
| **Device Name** | **Company** | **Design** |
| PLAATO | Appriva Medical Inc. | Single-lobe occluder; nitinol cage; ePTFE membrane hooks |
| WATCHMAN | Boston Scientific | Single-lobe occluder;nitinol frame; PET membrane; hooks |
| ACP | St. Jude Medical | Lobe and disc (polyester mesh); nitinol mesh structure; stabilizing wires |
| Amulet | St. Jude Medical | Lobe and disk (polyester mesh in both); nitinol mesh structure; stabilizing wires |
| WaveCrest | Coherex Medical | Single-lobe occluder; nitinol frame, polyurethane foam and ePTFE membrane; retractable anchors |
| Occlutech LAA | Occlutech | Single-lobe occluder; nitinol wire mesh; stabilizing loops; nanomaterial covering |
| Sideris Patch | Custom Medical Devices | Frameless detachable latex balloon covered with polyurethane |
| Lambre | Lifetech | Lobe and disk; nitinol; PET membrane; distal barbs anchors |
| Pfm | PFM Medical | Dual disk (distal anchor, variable middle connector, proximal disk); nitinol frame |
| Ultrasept | Cardia | Lobe and disk; nitinol frame; Ivalon covering; distal anchors |

**Table 2** **Summary of data for left atrial appendage occlusion**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **PROTECT-AF[46]** | **CAP[43]** | **ASAP[42]** | **EWOLUTION[49]** | **ACP[50]** |
| Patients (*n*) | 463 | 460 | 150 | 1021 | 1047 |
| Follow-up | 4 yr | 16 mo | 14 mo | 30 d | 13 mo |
| CHADS-score | 2.2 | 2.4 | 2.8 | 2.8 | n.a. |
| CHA2DS2-Vasc score | n.a. | n.a. | 4.4 | 4.5 | 4.5 |
| Procedural success | 88.0% | 95.0% | 94.7% | 98.5% | 97.3% |
| Procedural stroke | 1.3% | 0 | 0.7% | n.a. | 0.9% |
| Pericardial effusion | 4.8% | 2.2% | 1.3% | 0.5% | 1.2% |
| Device embolization | 0.6% | 0 | 1.3% | 0.2% | n.a. |
| Major bleeding | 3.5% | n.a. | n.a. | 1.6% | 1.5% |
| Long-term stroke | 2.3% | 1.5% | 0.7% | 0.3% | 2.3% |

AF: Atrial fibrillation; n.a.: Not applicable.

**Table 3 Different aspects of left atrial appendage occlusion**

|  |  |
| --- | --- |
| **Pro** | **Contra** |
| Non-inferiority to oral anticoagulation | Evaluation of other atherothrombotic sources |
| Alternative in patients with contraindication for anticoagulation | Unknown hemodynamic impact |
| Cost-effective | Postprocedural medical treatment not well defined |
| Reduced cumulative bleeding events during follow-up | No coparison between different devices |
| Good results in real-world registries | Undefined impact of residual leaks |