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One-Year Outcomes After Amulet or Watchman Device for Percutaneous Left Atrial Appendage Closure: a Pre-Specified Analysis of the SWISS-APERO Randomized Clinical Trial

Running title: Galea et al. One-Year Outcomes of SWISS-APERO Trial

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*This article is published in its accepted form, it has not been copyedited and has not appeared in an issue of the journal. Preparation for inclusion in an issue of Circulation involves copyediting, typesetting, proofreading, and author review, which may lead to differences between this accepted version of the manuscript and the final, published version.

**This work was presented as an abstract at TCT, October 23 - 26, 2023



Non-standard Abbreviations and Acronyms

| AF | Atrial Fibrillation |
|-------|---|
| DRT | Device Related Thrombus |
| CCTA | Cardiac Computed Tomography Angiography |
| IDL | IntraDevice Leak |
| LAA | Left Atrial Appendage |
| LAAC | Left Atrial Appendage Closure |
| MIL | Mixed Leak |
| PA | Patent Appendage |
| PANVL | Patent Appendage with Non Visible Leak |
| PDL | PeriDevice Leak |
| TEE | Trans-Esophageal Echocardiography |



Percutaneous left atrial appendage (LAA) closure (LAAC) prevent thromboembolisms in atrial fibrillation (AF) patients. However, residual LAA patency post-LAAC has been associated with higher thromboembolic risk, especially when detected remotely after the procedure (1). Residual leaks are typically assessed by means of transesophageal echocardiography (TEE) or cardiac computed tomography angiography (CCTA), and the latter imaging modality provides a comprehensive operator-independent assessment and higher sensitivity compared with TEE (2). The SWISSAPERO

(https://www.clinicaltrials.gov.-Unique identifier:NCT03399851) trial is the first randomized comparison between Watchman FLX (Boston Scientific, USA) and Amplatzer Amulet (Abbott, USA), which are the two most frequently used devices for LAAC worldwide (3.4). Patients with non-valvular AF and clinically indicated LAAC were eligible if provided written informed consent, had CHA2DS2-VASc score ≥ 2 and either HAS-BLED score ≥ 3 or presence of high bleeding risk features (3). The key exclusion criteria included the presence of LAA thrombus or LAA morphology not suitable for both study devices at TEE. All patients randomized to Watchman before October 2019 received Watchman 2.5, whereas Watchman FLX was exclusively used thereafter. Post-implantation treatment was left at discretion of the treating physician. At 13 months (395 ± 30 days), patients underwent an onsite clinical visit and CCTA. The CCTA images and the clinical events were centrally and blindly assessed. LAA was defined as patent (PA) if LAA density \geq 100 Hounsfield unit (HU) or $\geq 25\%$ of that of the LA (3,4). In patients with PA, the presence of visible leaks was adjudicated as intradevice leak [IDL], peridevice leak [PDL] or mixed leak [MIL] if contrast medium was visible along the entire or part of the length of device respectively; or as PAs with no visible leak (PANVL) if none of the above was detected. DRT was defined as homogenous hypoattenuated thickening (HAT) on the atrial surface of the device and centrally adjudicated as definitive or possible DRT (3). The trial was an investigator initiated

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trial, partially supported by a research grant from Abbott (St. Jude Medical/Abbott, Nathan Lane North Plymouth, MN, USA). The trial protocol was approved by the institutional review boards and all patients provided informed consent before enrolment.

The imaging endpoints were analysed by using chi-squared test and the risk ratios (RR) with 95% confidence intervals (CI) are reported. The clinical endpoints were analysed by using Cox proportional hazards model and the hazard ratios (HR) with 95% CI are reported. Data were analysed according to the intention-to-treat principle. A value of P<0.05 was considered statistically significant.

The data that support the findings of this study are available from the corresponding author upon reasonable request.

A total of 221 patients were assigned to Amulet (111[50.2%]) or Watchman (110[49.8%]; 85 received Watchman FLX and 25 Watchman 2.5). At 13 months (393[378-419] days), 24 patients died, 24 refused CCTA, 2 were lost-to-follow-up and 7 did not undergo CCTA for medical reasons. A total of 164 [74.2%; 84(75.7%) with Amulet and 80 (72.2%) with Watchman 2.5/FLX] patients underwent 13-month CCTA. PAs were observed in 45 (53.6%) patients with Amulet and 39 (48.8%) patients with Watchman 2.5/FLX (RR:1.10; 95%Confidence Interval [CI]:0.81-1.48; p=0.537) (Figure 1). The LAA patency subtypes were similarly distributed between groups. IDL occurred in 20 (23.8%) patients with Amulet and 14 (17.5%) patients with Watchman 2.5/FLX (RR: 1.36; 95%CI: 0.74-2.51; p=0.319). Side gap-leaks (the composite of PDL and MIL) occurred in 24 (28.6%) patients with Amulet and 22 (27.5%) patients with Watchman 2.5/FLX (RR: 1.04; 95%CI: 0.64-1.70; p=0.879). PANVL was detected in a minority of patients in both device groups (7.1% vs. 12.5%; RR: 0.57; 95%CI: 0.22-1.50; p=0.248). Definite or possible DRT at 13-month CCTA did not differ with Amulet or Watchman 2.5/FLX (2.4% vs. 3.8%; RR: 0.63; 95%CI: 0.11-3.70; p=0.610). The composite of cardiovascular death, ischemic stroke and systemic

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embolism (9.5% vs. 10.2%; HR:0.91; 95%CI: 0.39-2.14; p=0.829), cerebrovascular events (2.7% vs. 3.7%; HR: 0.75; 95%CI: 0.17-3.35; p= 0.706) or bleeding (40.8% vs. 31.4%; HR:1.46; 95%CI: 0.93-2.28; p= 0.098) were comparable between groups. Results remained entirely consistent when only patients with Watchman FLX were compared with Amulet. In the Amulet IDE trial, LAA complete occlusion rate at 12-month TEE was higher with Amulet compared with Watchman 2.5 (63.2% vs. 53.1%; p<0.01)(5). It is unclear if the inconsistent findings between our and Amulet IDE trials are explained by the predominant use of Watchman FLX in the Watchman group in our trial, by the different primary imaging modality used to assess PA, or a combination of both factors. Roughly, a quarter of patients did not undergo 13-month CCTA, which may have impacted study power to assess differences between the two groups. Our trial was not powered to show differences with regard to clinical endpoints. On the other hand, we confirmed in a smaller cohort involving the new iteration Watchman FLX, the similar long-term clinical outcomes after Watchman or Amulet device, which was similarly observed in the 3-year Amulet IDE follow-up(5).

Among patients undergoing clinically indicated LAAC and in whom LAA anatomy was deemed suitable for both Amulet and Watchman 2.5/FLX devices, rates of PA and DRT at 13-month CCTA, as well as clinical outcomes, did not differ between the two groups.

Article Information

Registration: URL: https://www.clinicaltrials.gov; Unique identifier: NCT03399851.

Acknowledgments

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Sources of Funding

This trial is investigator-initiated. The study sponsor, Insel Gruppe AG, Universitätsklinik für Kardiologie, CH-3010 Bern (Switzerland), for the conduction of the study was supported by local available funding and a research grant from Abbott. The funding company was not involved with the study processes, including site selection and management, and data collection and analysis.

Disclosures

NM reports grants and personal fees from Bayer Healthcare, grants and personal fees from BMS Pfizer, personal fees from Astra Zeneca, personal fees from Terumo, grants and personal fees from Abbott, outside the submitted work.. FDM reports consultancies and paid expert testimonies from Abbott and Boston-Scientific. AA is a proctor and consultant for Abbott and Boston-Scientific. KC reports research grant from Janssen Switzerland (Johnson & Johnson). CG reports funding from the Swiss National Science Foundation, InnoSuisse, CAIM foundation, GAMBIT foundation and Novartis foundation, outside of the submitted work. PV reports personal fees from AstraZeneca, Bristol Myers Squibb-Janssen, Bristol

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Myers Squibb-Pfizer, CSL Behring and Daiichi-Sankyo outside the submitted work. UF reports grants from Medtronic, other from Medtronic, Stryker and CSL-Behring, outside the submitted work. FB is proctor for Abbott, Boston-Scientific and Medtronic; he reports consultancies from Terumo and Meril. LR reports research grants to institution by Abbott-Vascular, Boston-Scientific, Biotronik,Heartflow, Sanofi,Regeneron. He reports speaker/consultation fees by Abbott-Vascular, Amgen, AstraZeneca, CSL-Behring, Canon,Occlutech, Sanofi,Vifor. MV has received grants and/or personal fees from AstraZeneca, Terumo, Alvimedica/CID, Abbott-Vascular, Daiichi-Sankyo, Opsens,Bayer, CoreFLOW,Idorsia-Pharmaceuticals-Ltd., Universität Basel Department Klinische Forschung, Vifor, Bristol-Myers-Squibb-SA, iVascular, and Medscape. All other authors have reported that they have no relationships relevant to the contents of this paper to disclose.

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Figure 1. One-Year Outcomes of SWISS-APERO Trial

AF, Atrial Fibrillation; CCTA, Cardiac Computed Tomography Angiography; CI,

Confidence Interval; CVD, Cardiovascular Death; DRT, Device Related Thrombus; HR,

Hazard Ration; LAA, Left Atrial Appendage; LAAC, Left Atrial Appendage Closure; RR,

Relative Risk; SE, Systemic Embolism.



In a European Multicenter Cohort of 221 High Risk AF Patients with Clinical Indication to LAAC

