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# Stent thrombosis in patients treated with drug-eluting stents and bioresorbable vascular scaffolds

Mechanisms, long-term outcomes and sex differences

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Publication date 2021

Link to publication

Citation for published version (APA):

Kerkmeijer, L. S. M. (2021). Stent thrombosis in patients treated with drug-eluting stents and bioresorbable vascular scaffolds: Mechanisms, long-term outcomes and sex differences. [Thesis, fully internal, Universiteit van Amsterdam].

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Download date:11 Feb 2023

# Final five-year results of the AIDA trial: a randomized trial comparing Absorb BVS with Xience EES in daily clinical practice

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Under review Eurointervention

# **Abstract**

**Background:** Absorb bioresorbable vascular scaffold (BVS) related events have been reported between 1 and 3 years – the period of active scaffold bioresorption. Data on the performance of Absorb BVS in daily clinical practice beyond this time point is scarce.

**Aims:** This report provides the final five-year clinical follow-up of the Absorb BVS in comparison with Xience everolimus-eluting stent (EES). In addition, we evaluated the effect of prolonged dual-antiplatelet therapy (DAPT) administration on events in the scaffold group.

Methods and Results: AIDA was a multicentre, investigator-initiated, non-inferiority trial, in which 1,845 unselected patients with coronary artery disease were randomly assigned to either Absorb BVS (n=924) or Xience EES (n=921). Through 5 years follow-up, there was no difference in target vessel failure, composite of cardiac death, target vessel myocardial infarction or target vessel revascularisation, between Absorb BVS (17.7%) and Xience EES (16.1%) (hazard ratio 1.31, 95% CI 0.90-1.41, p=0.302). Definite or probable device thrombosis (DT) occurred in 43 patients (4.8%) of the scaffold group compared to 13 patients (1.5%) of the stent group (hazard ratio 3.32; 95% CI 1.78-6.17; p<0.001). Device thrombosis between 3- and 4-years occurred six times in Absorbarm versus three in Xience-arm. Between 4- and 5-years the incidence was 3 versus 2, respectively. Of those three DT in scaffold group, two occurred in Xience EES treated lesions. When scaffold thrombosis cases matched with controls and tested for effect of DAPT, the odds ratio of scaffold thrombosis in patients on DAPT compared to off DAPT throughout 5-year follow-up was 0.36 (95% CI 0.15-0.86).

**Conclusion:** The excess risk of Absorb BVS on late adverse events, in particular device thrombosis, in routine PCI continues up to 4-years. DAPT appears to mitigate the risk of scaffold thrombosis.

# Introduction

Drug-eluting stents have an ongoing risk of device-related adverse events long after the implantation.(1) The pathogenesis of this ongoing annual hazard is thought to be the permanent presence of a metallic implant. To liberate the coronary artery of its permanent metallic cage and therefore remove the potential cause of restenosis and stent thrombosis, the bioresorbable vascular scaffolds (BVS) were developed. Theoretically, the function of the BVS is to scaffold the arterial wall after balloon dilatation to prevent acute vessel closure and late constrictive remodeling, afterwards it should dissolve over approximately 3 years' time to restore the native structure of the coronary artery. The most widely studied coronary scaffold is the Absorb BVS (Abbott Vascular, Santa Clara, USA), which, in a porcine model, completely resorbs and integrates in approximately 3 years.(2) However, in clinical practice, Absorb BVS was found to be associated with an increased risk of target-vessel myocardial infarction and device thrombosis during the time of reabsorption compared to everolimus-eluting metallic Xience stent (Abbott Vascular, Santa Clara, USA).(3-5) Beyond the 3-year time-point, data on safety and efficacy of the Absorb BVS is scarce.(6) In addition, it is unknown whether prolonged dual antiplatelet therapy (DAPT) benefits patients treated with Absorb BVS. Therefore, long-term outcomes are of interest. The Amsterdam Investigator-initiateD Absorb strategy (AIDA) randomized clinical trial compared the Absorb BVS with the everolimus-eluting metallic Xience stent (Xience EES; Abbott Vascular) in daily clinical practice.(7) Herein we report the final five-year clinical outcomes of Absorb BVS in comparison with Xience EES. In addition, we evaluate whether prolonged DAPT regimes mitigate the occurrence of scaffold thrombosis.

# **Methods**

The study design, endpoint definitions, and results through 3 years have been previously described in detail.(3, 7-9) Briefly, the AIDA trial was an all-comers, multicentre, investigator-initiated, randomized controlled trial. Between August 2013 and December 2015, 1,845 consecutive patients with coronary artery disease undergoing percutaneous coronary intervention (PCI) of one or more target lesions suitable for drug-eluting stent implantation were enrolled. Follow-up was performed at regular intervals through 5 years. Quantitative coronary angiographic analyses were performed at a core laboratory. An independent clinical event committee adjudicated all major adverse cardiac events according to either Third Universal Myocardial Infarction definitions(10), or the Academic Research Consortium definitions.(11) The primary study endpoint was target vessel

failure (TVF), powered for non-inferiority at 2-years. TVF is a composite of cardiac death, target vessel myocardial infarction (TV-MI), or target vessel revascularisation. Secondary endpoints included TVF, its components, and device thrombosis at each follow-up period.

The protocol mandated use of DAPT for at least one-year post-PCI. In January 2017, the data and safety monitoring board (DSMB) noted higher rate of early and late scaffold thrombosis and recommended considering prolonged DAPT in all patients treated with Absorb BVS. Subsequently, this recommendation was implemented and referring cardiologists were advised to prescribe DAPT up to 3 years in all patients treated with Absorb BVS.

The study design was in concordance with the provisions of the Declaration of Helsinki. The research ethics committee of Academic Medical Centre, Amsterdam approved the study protocol for all participating centres. All enrolled patients provided written informed consent.

#### **Effect of DAPT**

To assess the effect of DAPT on occurrence of scaffold thrombosis (ScT), every case with definite ScT was matched with one or two control case(s) based on age, sex, presenting with acute coronary syndrome, total number of stents, total stent length and enrolment date before October 1<sup>st</sup> 2014. At the time of ScT, use of DAPT was scored yes or no for the cases and their controls.

# Statistical analysis

The current paper reports the pre-specified major outcomes at 5-year follow-up. All analyses were performed according to the intention-to-treat principle. Time-to-event curves were constructed by the Kaplan-Meier and compared with log-rank test. Hazard ratios were calculated using cox regression. Landmark analyses were performed at 3 and 4 years after index procedure. All ScT cases were matched fuzzy (1:2). Fuzz of 10 for age, 14 for days, 0.8 for total number of stents and 19 for total stent length were allowed. The effect of DAPT on the occurrence of ScT was assessed by calculating odds ratio, using multivariable logistic regression adjusting for age, total number of stents and total stent length. A p-value of ≤0.05 was considered statistically significant. All statistical analyses were performed with SPSS software, version 26.0 (IBM SPSS Statistics, IBM Chicago, IL, USA).

# Results

From August 2013 until December 2015, 1,845 patients were enrolled at five sites throughout the Netherlands. In total, 924 patients were randomized Absorb BVS and 921 patients were randomized to Xience EES. Baseline patient, procedural and lesion characteristics were described in detail in previous reports (1, 5), and are shown in **supplementary table 1 and 2**. Briefly, baseline characteristics were well balanced between the two groups. A total of 54% patients presented with acute coronary syndrome at baseline; 25.2% ST-segment myocardial infarction, 20.4% non-ST-segment myocardial infarction and 8.5% unstable angina. SYNTAX score was available for 1,661 patients (90.0%) with a median of 11 (IQR 7-18). In total, 2,446 lesions were treated.

# **Clinical endpoints**

Complete five-year follow-up was obtained in 95.1% of patients; a study flowchart is displayed in the **supplementary Figure 1**. Clinical outcomes through 5-year follow-up are shown in **Table 1**. Throughout 5 years, no significant difference in TVF-rate was found between patients treated with Absorb BVS (17.7%) versus Xience EES (16.1%) (HR 1.13, 95% CI 0.90-1.41, p =0.302) (**Figure 1**). The rates of TV-MI and target lesions revascularization (TLR) remained significantly increased in the Absorb-arm compared to the Xience-arm, with 5-year follow-up rates of TV-MI 7.7% vs 5.0% (HR 1.57, 95% CI 1.08-2.30; p =0.018) and TLR 10.1% vs 7.3% (HR 1.41, 95% CI 1.02-1.94; p=0.034), respectively.

Landmark analysis of clinical outcomes between 3- and 4-year, and 4- and 5-year follow-up are shown in **Table 2**. Clinical outcomes at 4-years follow-up are shown in **supplementary table 3**. Between 3- and 4-years, the rates of TV-MI were numerically higher in Absorb BVS compared to Xience EES, 1.1% vs. 0.4% HR 3.01, 95% CI 0.82-5.76, p=0.082). The rates of TLR were significantly higher in Absorb BVS compared to Xience EES, 1.6% vs. 0.5% (HR 3.27, 95% CI 1.07-10.02; p=0.028). This difference was mainly driven by TLR due to restenosis, 1.4% vs 0.4%, respectively (HR 3.61, 95% CI 1.01-12.93; p=0.035).

In contrast, between 4- and 5-years, the rates of TV-MI did not differ between Absorb BVS (0.7%) and Xience EES (0.8%) (HR 0.83, 95% CI 0.25-2.73; p=0.763). Also, the incidence of TLR did not differ between Absorb BVS and Xience EES, 0.8% vs. 1.1% (HR 0.75, 95% CI 0.26-9.02; p=0.602).

**TABLE 1.** Clinical Outcomes through 5-year follow-up

		At 5 year	ars	
_	Absorb BVS (n=924)	Xience EES (n=921)	Hazard Ratio (95% CI)	p value <sup>s</sup>
All-cause death	76 (8.4%)	88 (9.8%)	0.85 (0.63-1.16)	0.314
Cardiac death	34 (3.8%)	41 (4.7%)	0.82 (0.52-1.29)	0.396
Cardiovascular death	43 (4.8%)	47 (5.4%)	0.91 (0.60-1.37)	0.641
All myocardial infarction	96 (10.7%)	62 (7.1%)	1.56 (1.13-2.15)	0.006
Target vessel MI	69 (7.7%)	44 (5.0%)	1.57 (1.08-2.30)	0.018
Non-target vessel MI	27 (3.1%)	19 (2.2%)	1.41 (0.79-2.54)	0.246
Any revascularisation	179 (20.1%)	152 (17.3%)	1.18 (0.95-1.47)	0.127
Target vessel revascularisation	119 (13.4%)	94 (10.7%)	1.27 (0.97-1.66)	0.084
Target lesion revascularisation	90 (10.1%)	64 (7.3%)	1.41 (1.02-1.94)	0.034
Device thrombosis related	37 (4.1%)	9 (1.0%)	4.12 (1.99-8.54)	< 0.001
Device stenosis related	58 (6.6%)	56 (6.4%)	1.02 (0.71-1.48)	0.896
Composite endpoints				
Target vessel failure	160 (17.7%)	143 (16.1%)	1.13 (0.90-1.41)	0.302
Target lesion failure†	135 (14.9%)	121 (13.7%)	1.12 (0.88-1.43)	0.356
Patient-oriented composite endpoint‡	259 (28.4%)	241 (26.6%)	1.09 (0.91-1.29)	0.351

p-values were calculated by the log-rank test. p-composite of cardiac death, target vessel myocardial infarction and target lesion revascularisation. p-composite of death, myocardial infarction or any revascularisation. p-bioresorbable vascular scaffold; p-composite of death, myocardial infarction p-composite of death, myocardial infarction p-composite of cardiac infarction p-composite of cardiac death, target vessel myocardial infarction p-composite of death, myocardial infarction p-composite of p-composite of

TABLE 2. Landmark analysis for clinical Outcomes between 3- and 5-years follow-up

		Between 3 and 4 years	ınd 4 years			Between 4 and 5 years	ıd 5 years	
	Absorb BVS (n=924)	Xience EES (n=921)	Hazard Ratio (95% CI)	p value <sup>§</sup>	Absorb BVS (n=924)	Xience EES (n=921)	Hazard Ratio (95% CI)	p value <sup>s</sup>
All-cause death	14 (1.6%)	18 (2.1%)	0.77 (0.38-1.54)	0.453	16 (2.0%)	17 (2.2%)	0.93 (0.47-1.83)	0.828
Cardiac death	4 (0.5%)	7 (0.8%)	0.56 (0.16-1.92)	0.354	6 (0.7%)	8 (1.1%)	0.74 (0.26-2.13)	0.574
Cardiovascular death	5 (0.6%)	9 (1.1%)	0.55 (0.18-1.64)	0.274	9 (1.1%)	10 (1.3%)	0.89 (0.36-2.18)	0.793
All myocardial infarction	13 (1.6%)	(0.8%)	2.19 (0.83-5.76)	0.103	8 (1.1%)	7 (0.9%)	1.15 (0.42-3.18)	0.780
Target vessel MI	9 (1.1%)	3 (0.4%)	3.01 (0.82-11.13)	0.082	5 (0.7%)	(%8.0) 9	0.83 (0.25-2.73)	0.763
Non-target vessel MI	3 (0.4%)	3 (0.4%)	0.99 (0.20-4.91)	0.991	3 (0.4%)	2 (0.3%)	1.49 (0.25-8.90)	0.661
Any revascularisation	26 (3.6%)	14 (1.9%)	1.88 (0.98-3.60)	0.053	13 (2.0%)	18 (2.7%)	0.74 (0.36-1.50)	0.399
Target vessel revascularisation	20 (2.6%)	5 (0.7%)	4.01 (1.50-10.68)	0.003	9 (1.3%)	12 (1.7%)	0.76 (0.32-1.80)	0.526
Target lesion revascularisation	13 (1.6%)	4 (0.5%)	3.27 (1.07-10.02)	0.028	(%8.0) 9	8 (1.1%)	0.75 (0.26-2.18)	0.602
Device thrombosis related	5 (0.6%)	2 (0.2%)	2.51 (0.49-12.96)	0.254	3 (0.4%)	2 (0.3%)	1.51 (0.25-9.02)	0.651
Device stenosis related	11 (1.4%)	3 (0.4%)	3.61 (1.01-12.93)	0.035	4 (0.5%)	(%8.0) 9	0.66 (0.19-2.33)	0.514
Composite endpoints								
Target vessel failure	22 (2.9%)	12 (1.6%)	1.85 (0.91-3.73)	0.082	13 (1.8%)	21 (3.0%)	0.63 (0.31-1.25)	0.181
Target lesion failure†	16 (2.1%)	11 (1.4%)	1.47 (0.68-3.16)	0.324	10 (1.4%)	18 (2.5%)	0.56 (0.26-1.21)	0.136
Patient-oriented composite endpoint;	35 (4.9%)	30 (4.2%)	1.19 (0.73-1.94)	0.481	26 (4.0%)	33 (5.0%)	0.81 (0.48-1.35)	0.415

\$ p-values were calculated by the log-rank test. † Composite of cardiac death, target vessel myocardial infarction and target lesion revascularisation. ‡ Composite of death, myocardial infarction or any revascularisation. BVS = bioresorbable vascular scaffold; EES = everolimus-eluting stent; MI = myocardial infarction

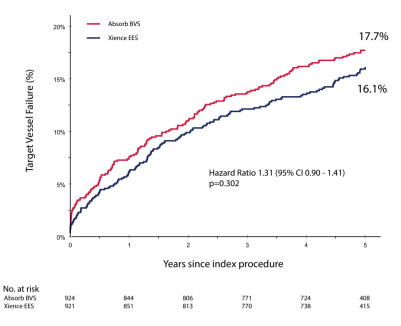


FIGURE 1. Kaplan-Meier curves for target vessel failure up to five-year follow-up per study arm

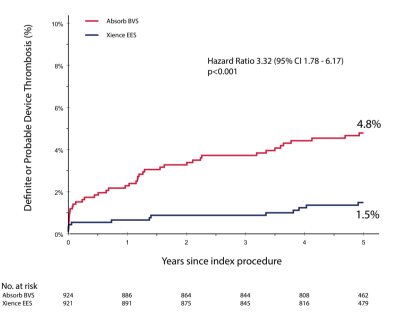


FIGURE 2. Kaplan-Meier curves for definite/probable device thrombosis to five-year follow-up per study arm

#### **Device thrombosis**

Device thrombosis (DT) rates are shown in **Table 3**. At 5-years, 38 Absorb BVS treated patients suffered from definite DT compared to 9 Xience EES treated patients (HR 4.24, 95% CI 2.05-8.77; p<0.001). Descriptive characteristics of the definite DT cases throughout 5-year follow-up are presented in **Supplementary Table 4 and 5**. The rate of definite/probable DT was significantly increased in Absorb BVS arm compared with Xience EES arm, with a 5-year rate of 4.8% (43 cases) versus 1.5% (13 cases) (HR 3.32, 95%CI 1.78-6.17; p<0.001) (**Figure 2**).

Between 3 and 4 years, five definite DT and one probable DT were noted in the Absorb BVS arm compared to three definite DT in the Xience EES arm. Of the five definite scaffold thrombosis cases, one case was treated with two-stent technique in a bifurcation lesion and the DT occurred at 1277 days post index PCI. The second very late scaffold thrombosis (VLST) was described as thrombosis on severe restenosis by the clinical event committee. The other three VLST cases had target lesion revascularization with a DES prior to the occurrence of DT. Between 4 and 5 years, three definite DT in Absorb-arm vs two in Xience-arm were noted. Two of these three DT cases were randomized at baseline to Absorb BVS but treated with Xience EES during index procedure.

**TABLE 3.** Incidence of device thrombosis through 5-year follow-up

	Absorb BVS (n=924)	Xience EES (n=921)	Hazard ratio (95% CI)	p value <sup>s</sup>
Definite	38 (4.3%)	9 (1.0%)	4.24 (2.05-8.77)	<0.001
Probable	5 (0.5%)	4 (0.5%)	1.24 (0.33-4.62)	0.747
Possible	16 (1.8%)	25 (3.0%)	0.63 (0.34-1.18)	0.150
Definite/probable	43 (4.8%)	13 (1.5%)	3.32 (1.78-6.17)	< 0.001
≤ 24 hours (acute)	3	3		
>24 hours to 30 days (subacute)	10	2		
31 days to 1 year (Late)	8	1		
1 – 2 years (Very late)	9	2		
2 – 3 years (Very late)	4	0		
3 – 4 years (Very late)	6	3		
4 – 5 years (Very late)	3	2		
Any device thrombosis	58 (6.5%)	38 (4.4%)	1.53 (1.02-2.31)	0.039

<sup>\$</sup>p\$-values were calculated by the log-rank test. BVS = bioresorbable vascular scaffold; EES = everolimus-eluting stent

#### **Effect of DAPT on Scaffold thrombosis**

During 5-year follow-up, 21 very late definite scaffold thrombosis occurred in the Absorb arm. Only one of these 21 VLST (4.8%) was on DAPT at the time of the event. This is in stark contrast to early DT, where 12 of the 17 patients (70.6%) used DAPT at the time of the event (**Figure 3**). Patients were advised to prolong DAPT up to three years. **Figure 4** shows data on aspirin, P2Y12 inhibitors, DOAC and DAPT use at all follow-up points. All VLST between 3 and 4 years occurred in patients without use of DAPT regimens. These patients discontinued DAPT 331 days (range 119-632) prior to the event. Detailed information on DAPT status at the time of ScT can be found in **supplementary table 4**.

To make the effect of DAPT more transparent, the definite ScT cases were matched with control cases. Four of 38 ScT cases were not eligible; in two ScT cases the SYNTAX score was not available and DAPT-status was unknown in another two ScT cases. DAPT-status was also missing in three matched controls. Therefore, 34 ScT cases with 65 matched controls were included for analysis. Of those who suffered ScT, 13 patients were on DAPT and 21 patients off DAPT. Of those who did not develop ScT, 41 used DAPT and 24 did not. The odds ratio of ScT with the use of DAPT throughout 5-year follow-up was 0.36 (95% CI 0.15-0.86). Within the first year the OR of ScT was 0.14 (95% CI 0.02-0.85) and between 1- and 5-year follow-up the OR was 0.17 (95% CI 0.02-1.63) (**Figure 5**).

# **Discussion**

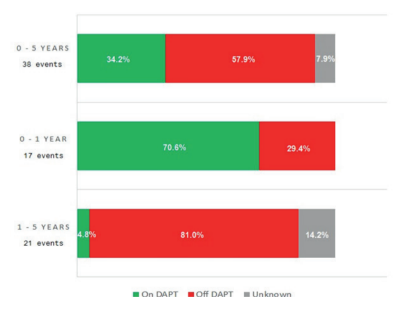
The main findings of this final five-year report on clinical outcomes of Absorb BVS in comparison with Xience EES from the AIDA-trial were as follows: 1) Absorb BVS was associated with a significantly increased risk of target-vessel myocardial infarction and device thrombosis compared to Xience EES tested in daily clinical practice; 2) landmark analysis has shown a plateauing of this excess risk with Absorb BVS starting at four-years; and 3) retrospective analysis indicates a reduced odds ratio of scaffold thrombosis in patients using DAPT regimen.

#### The excess risk of Absorb BVS thrombosis

Randomized clinical trials, comparing Absorb BVS with Xience EES, have identified an increased risk with Absorb BVS on TV-MI and DT up to 3 years after implantation. Stone et al.(12) demonstrated in a pooled analysis of the Absorb trials that this excess risk with Absorb BVS was no longer apparent beyond 3 years. Compared to the first 3 years, the hazard ratios of target lesion failure dropped from 1.42 to 0.92 and the hazard ratio of DT dropped from 3.86 to 0.44 between 3 and 5 years.(12) Our results, however,

show a continued excess risk up to 4-years. Between 3-4 years, the hazard ratio of target lesion failure increased from 1.13(3) to 1.22 at 4 years and the increased risk of device thrombosis diminished but did not disappear (HR dropped from 6.02(3) to 2.52. It was only after four years that the excess risk with Absorb BVS was no longer apparent. The hazard ratio of target lesion failure dropped to 0.56, and for DT it dropped to 1.51. However, two of the three device thrombosis cases between four and five years occurred in Xience EES treated lesions instead of the randomized device, scaffold. Therefore, the hazard ratio is overestimated.

The difference in outcomes between the ABSORB trials and AIDA might be partly explained by the difference in study population; the study population of the ABSORB trials mainly consisted of patients with simple lesions and low risk of restenosis. In comparison, the AIDA trial represented the daily clinical practice and included patients with complex lesions and patients who had presented with acute coronary syndrome including ST-segment elevation myocardial infarction. It might be that the resorption of the Absorb BVS is prolonged in these complex and severe diseased lesions and thereby creates a longer lasting risk of device-related events.(13) A better understanding of this resorption process and the factors that influences it could help us to improve next generation BRS devices.



**FIGURE 3.** Relationship between definite device thrombosis and dual antiplatelet therapy (DAPT) status at the time of the event during 5-year follow-up

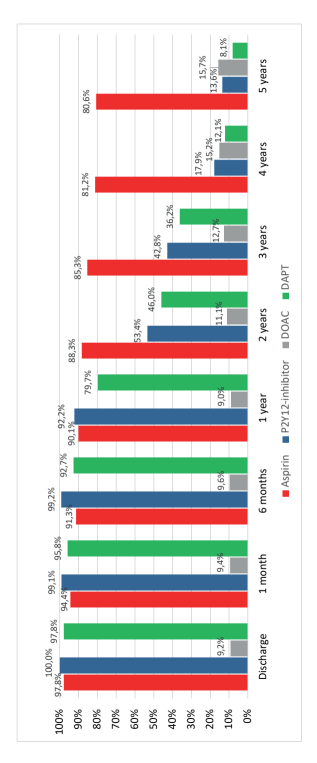


FIGURE 4. Antiplatelet therapy per follow-up in Absorb BVS-group.

#### **Effect of DAPT**

Device thrombosis is a serious complication with high morbidity and mortality. (14) DAPT significantly reduces the risk of stent thrombosis in DES.(15, 16) The introduction of bioresorbable scaffolds led to the question on whether current DAPT recommendations after DES implantation are also applicable to this different technology. A prolonged ischemic risk period could be expected due to its larger footprint (strut thickness 157um) compared with contemporary second-generation DES (60 to 90um), which may lead to greater platelet activation and delayed endothelialisation.(17, 18) In addition, intraluminal dismantling of Absorb BVS at sites without complete endothelialisation during the resorption process has been suggested as a new mechanism of device thrombosis.(19) Indeed, our results demonstrated an increased ischemic risk period of four years with Absorb BVS compared to Xience EES. Especially high complex PCI, as bifurcation stenting, long lesions or double layer stents, led to ischemic events long after the index procedure. Therefore, prolonged DAPT may be justified and outweigh an increased bleeding risk particularly after complex PCI. Although our analyses are retrospective and should be interpreted with caution, the odd ratios of DAPT on ScT are suggestive of possible effect. In addition, there was no temporal relationship between DAPT discontinuation and VLST. For example, all ScT between 3- and 4-years follow-up occurred on average 331 days after DAPT cessation. Well-apposed and embedded struts at baseline can still protrude into the lumen later on during the reabsorption process. (20) It is also plausible that good apposition at baseline would not prevent the occurrence of acquired malapposition, as large plaque burden continues to exert an inner force on the progressively weaker resorbing device. So, a cause of scaffold thrombosis may occur at any time during the reabsorption process, rather than being present continuously and cause thrombosis after DAPT discontinuation.

Nevertheless, lacking data on major bleedings precludes us from commenting on the net clinical benefit of prolonged DAPT. Further research for the recommendation on DAPT duration after implantation of scaffolds, a completely different technology than metallic DES, is warranted.

#### Causes of scaffold thrombosis

The causes of very late scaffold thrombosis are not yet fully understood. It is thought that the underlying mechanism of very late scaffold thrombosis is mostly scaffold dismantling, followed by malappostion and neoatherosclerosis(21). The current data uncovered another possible mechanism of scaffold thrombosis. Three very late scaffold thrombosis occurred in lesions previously treated for restenosis with Xience EES. Lack of Optical Coherence Tomography images precludes us from making a more definitive conclusion about the

mechanisms of these particular cases and allows us only to speculate. It is possible that the DES itself caused device thrombosis. However, it cannot be excluded that resorption of the underlying BVS caused device thrombosis due to protrusion of the thrombogenic material or that it caused acquired malapposition of the DES. As new generations of scaffolds are being developed, it is important to further investigate whether it is safe to implant a metallic stent over the scaffold.

# Overcome very late device-related events

Bioresorbable scaffold were designed to overcome very late device-related events often caused by neoatherosclerosis.(22) However, neoatherosclerosis did also appear in Absorb BVS treated lesions (23) and led to at least one scaffold thrombosis. Neoatherosclerosis will eventually occur within any device if sufficiently potent risk factors remain active and Absorb BVS is not immune to the progression of neoatherosclerosis. In addition, the incidence of patient-oriented and device-related adverse events in the Xience EES group reported in the current article are not negligible. Target lesion failure-rate within the first year was 5.2%, afterwards an annual rate of  $\pm 2.2\%$ , and a total of 16.1% target lesion failure-rate at 5-year follow-up. Patient-oriented composite endpoint in Xience-arm within first year was 10.6%, afterwards an annual rate of  $\pm +/-$  4.0%, and 26.6% at 5-year follow-up. Therefore, regardless of the stent platform, more effort on secondary prevention is needed.

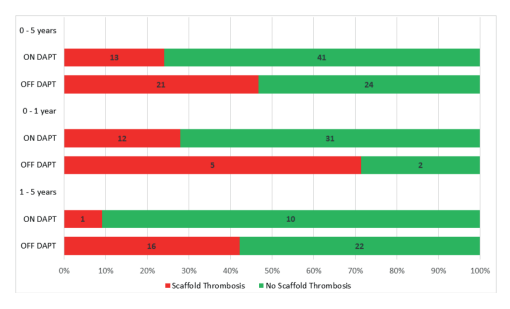


FIGURE 5. Effect of DAPT on occurrence of Scaffold Thrombosis

#### Limitations

The present study has several limitations. First, the AIDA trial was powered for the primary endpoint of TVF at 2 years. All secondary analyses on individual components of the primary endpoint such as scaffold thrombosis should be considered hypothesis generating. Second, the lack of systematic intravascular imaging in patients with clinical events, preclude more definite conclusions about the mechanisms related to BVS failure at different time points. Third, restarting or prolonging DAPT through three years after scaffold implantation was recommended at the request of the DSMB. This recommendation might have influenced the occurrence of thrombosis-related outcomes in patients on prolonged or restarted DAPT compared to patients who were treated according to the applicable guidelines and IFU. Fourth, patients and clinicians were unblinded to treatment assignment after the report of concerns about the safety of Absorb BVS upon the recommendation of the DSMB. Fifth, bleeding events were not monitored or adjudicated by clinical event committee and therefore it precludes us from assessing the net benefit of prolonged DAPT.

### **Conclusions**

In addition to previous reports, the increased risk for device related myocardial infarction and revascularization in patients treated with the Absorb bioresorbable vascular scaffold continues up to 4 years after index PCI and seems to plateau afterwards. Retrospective analyses indicate a reduction of odds with the use of prolonged DAPT on scaffold thrombosis. The later, however, would have to be weighted against the risk of bleeding in individual patients and needs further investigation.

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# **Supplementary material**

**SUPPLEMENTARY TABLE 1.** Characteristics of the Patients at Baseline\*

	Absort (N=9		Xience (N=	e EES 921)
Age - years	64.3	±10.6	64.0	±10.5
Male sex – n (%)	670	(73%)	700	(76%)
Risk factors – n/ total n (%)				
Diabetes mellitus	171/924	(19%)	153/921	(17%)
Requiring oral medication	95/171	(56%)	97/153	(63%)
Requiring insulin	65/171	(38%)	45/153	(37%)
Hypertension	468/920	(51%)	464/919	(51%)
Hypercholesterolemia	344/915	(38%)	350/914	(38%)
Family history of coronary artery disease	451/886	(51%)	469/886	(53%)
Current smoker	248/867	(29%)	273/861	(32%)
History – n/ total n (%)				
Chronic renal failure	70/924	(8%)	91/921	(10%)
Ejection fraction < 30%	22/910	(2%)	17/900	(2%)
Previous stroke or transient ischemic attack	46/923	(5%)	58/921	(6%)
Peripheral vascular disease	65/924	(7%)	56/918	(6%)
Previous myocardial infarction	166/924	(18%)	172/921	(19%)
Previous percutaneous coronary intervention	202/924	(22%)	184/921	(20%)
Previous bypass surgery	38/924	(4%)	26/921	(3%)
Clinical presentation - n (%)				
ST-segment elevation myocardial infarction	240	(26%)	225	(24%)
Non ST-segment elevation myocardial infarction	185	(20%)	192	(21%)
Unstable angina	70	(8%)	87	(9%)
Stable angina and/or documented ischemia	361	(39%)	370	(40%)
Angiographic driven	51	(6%)	36	(4%)
Other	17	(2%)	11	(1%)
SYNTAX score				
Mean	13.2	±8.6	12.6	±8.4
Median	11	(7-18)	11	(7-17)

 $<sup>^*</sup>$  plus-minus values are means  $\pm$  SD. Absorb BVS: Absorb bioresorbable vascular scaffold; n:number; Xience EES: Xience everolimus-eluting stent

**SUPPLEMENTARY TABLE 2.** Procedural Characteristics.\*

Outcome	Absort	BVS	Xience	EES	P Value
Patients					
Total no.	92	4	92	1	
Treated lesions per patient	1.34	±0.63	1.31	±0.59	0.360
Number of devices per patient	1.54	±0.84	1.45	±0.79	0.014
Total device length per patient – mm	31.1	±19.6	29.7	±19.2	0.113
Minimum device diameter per patient - mm	2.73	±0.27	2.88	±0.35	0.050
Device implantation - n (%)					
Any assigned study device	895	(96.9%)	919	(99.8%)	< 0.001
Only assigned study devices	859	(93.0%)	910	(98.8%)	< 0.001
Any unassigned device	65	(7.0%)	11	(1.2%)	< 0.001
Only unassigned devices	29	(3.1%)	2	(0.2%)	< 0.001
After failure assigned device	20		1		
Unassigned device first choice	9		1		
Procedure time – min mean (total n) $\pm$ SD	49 (919)	±26	44 (918)	±23	< 0.001
Contrast use – ml mean (total n) ± SD	160 (902)	±74	151 (897)	±72	0.016
Pre-dilatation first treated lesion – $n/$ total $n$ of target lesions (%)	911	(99%)	892	(97%)	0.012
Procedure success	834	(90%)	889	(97%)	< 0.001
Treated lesions¶					
Total no.	123	37	120	09	
Rotational atherectomy – $n/$ total $n$ of target lesions (%)	24/1232	(1.9%)	26/1208	(2.2%)	0.776
Pre-dilatation performed – n (%)	1199	(97%)	1103	(91%)	< 0.001
Total number of devices implanted	142	25	133	36	
Number of devices per lesion	1.15	±0.40	1.11	±0.34	0.001
Post-dilatation performed – no. (%)	915	(74%)	594	(49%)	< 0.001

<sup>\*</sup> Plus-minus values are means ± SD. # Listed is the diameter of the used pre-dilatation balloon, implanted stent or scaffold, and the used post-dilatation balloon. ¥ Listed is the maximum pressure of the pre-dilatation balloon, the stent or scaffold delivery-system balloon, and the post-dilatation balloon. ¶ All treated lesions at time of randomization and scheduled staged procedures. All abbreviations as in supplementary table 1.

**SUPPLEMENTARY TABLE 3.** Clinical Outcomes per study arm at 4 years follow-up

		, ,		
		At 4 yea	rs	
	Absorb BVS (n=924)	Xience EES (n=921)	Hazard Ratio (95% CI)	p value <sup>s</sup>
All-cause death	60 (6.6%)	71 (7.8%)	0.84 (0.59-1.18)	0.309
Cardiac death	28 (3.1%)	33 (3.7%)	0.84 (0.51-1.39)	0.502
Cardiovascular death	34 (3.7%)	37 (4.1%)	0.91 (0.57-1.45)	0.697
All myocardial infarction	88 (9.8%)	55 (6.2%)	1.61 (1.15-2.26)	0.005
Target vessel MI	64 (7.1%)	38 (4.3%)	1.69 (1.13-2.53)	0.009
Non-target vessel MI	24 (2.7%)	17 (1.9%)	1.40 (0.75-2.61)	0.282
Any revascularisation	166 (18.5%)	134 (15.0%)	1.24 (0.99-1.56)	0.061
Target vessel revascularisation	110 (12.3%)	82 (9.2%)	1.34 (1.01-1.79)	0.042
Target lesion revascularisation	84 (9.4%)	56 (6.3%)	1.50 (1.07-2.11)	0.017
Device thrombosis related	34 (3.8%)	7 (0.8%)	4.87 (2.16-10.99)	< 0.001
Device stenosis related	54 (6.1%)	50 (5.6%)	1.07 (0.73-1.57)	0.734
Composite endpoints				
Target vessel failure	147 (16.2%)	122 (13.5%)	1.21 (0.95-1.54)	0.116
Target lesion failure†	125 (13.7%)	103 (11.4%)	1.22 (0.94-1.58)	0.133
Patient-oriented composite endpoint‡	233 (25.4%)	208 (22.8%)	1.13 (0.94-1.36)	0.197
Device thrombosis				
Definite device thrombosis				
Probable device thrombosis				

Definite/probable device thrombosis

<sup>\$</sup> p-values were calculated by the log-rank test. † Composite of cardiac death, target vessel myocardial infarction and target  $lesion\ revascular is ation.\ \# Composite\ of\ death,\ myocardial\ infarction\ or\ any\ revascular is ation.\ BVS=bioresorbable\ vascular$ scaffold; EES = everolimus-eluting stent; MI = myocardial infarction

SUPPLEMENTARY TABLE 4. Descriptive characteristics of cases of definite device thrombosis

					Ref size	Pre-dilatation	Stent size	Post- dilatation			DAPT therapy	Clinical	
Case	Device	Initial PCI Indication	Treated Vessel	Lesion	(mm)	(atm)	(atm)	(atm)	Initial DAPT therapy	Days to DT	Time of DT	outcome (worst)	Patients note
	Absorb BVS	STEMI	Mid RCA	B2	4.0x15	3.0x15 (12)	3.5x18 (13)	4.0x12 (13)	ASA Ticagrelor	0	ASA Ticagrelor	Myocardial infarction	Dissection distal of stent (OCT)
2	Absorb BVS	STEMI	Prox LAD	B2	3.5x18	3.5x20 (6)	3.5x18 (14)	3.5x12 (20)	ASA Ticagrelor		ASA Ticagrelor	Myocardial	Distal edge dissection (OCT)
e	Absorb BVS	AP	Mid RCA	B2	3.0x15	3.0x15 (10)	3.5x18 (14)	4.0x12 (14)	ASA Clopidogrel	2	ASA Clopidogrel	Myocardial infarction	Malapposition stent (OCT)
4	Absorb BVS	AP	Mid RCA	C	3.0x46	2.5x20 (16)	3.0x28 (12) 3.0x18 (14)	3.0x20 (18)	ASA Clopidogrel	8	ASA Clopidogrel	Myocardial infarction	
5	Absorb BVS	STEMI	Prox LAD	С	3.5x21	2.0x12 (12)	3.0x15 (14) 3.5x12 (16)	3.75x15 (22)	ASA Clopidogrel	4	ASA Clopidogrel	Myocardial infarction	
9	Absorb BVS	AP	Distal RcX	B2	2.5x28	2.5x20 (10)	2.5x28 (10)	2.5x20 (14)	ASA Clopidogrel	~	ASA Clopidogrel	Myocardial infarction	Possible to low therapy compliance
	Absorb BVS	Stabilized	Prox RCA	O	3.0x30	3.5x15 (12) Rotablation	3.5x18 (14) 3.5x18 (14)	3.5x15 (14)	ASA Ticagrelor	9	ASA	Myocardial infarction	Patient forgot to take Ticagrelor
8	Absorb BVS	NSTEMI	Prox LAD	B2	2.5x15	2.5x15(UN)	2.5x18 (10)	3.0x12 (12)	ASA Ticagrelor	11	ASA Ticagrelor	Myocardial infarction	
	Absorb	I Water	Prox LAD Distal RCA	o o	3.0x25 2.7x25	2.5x20 (8) 3.5x20 (12)	3.0x28 (10) 2.5x28 (14)	3.5x15 (10) No	ASA	ć	ASA	Myocardial	DT in both LAD
2	BVS	STEMI	Mid RCA Mid RCA	U U	2.7x25 2.7x25	2.5x20 (10) 3.5x20 (10)	3.0x28 (14) 2.5x28 (14)	°Z °Z	Ticagrelor	67	Ticagrelor	infarction	and RCA

SUPPLEMENTARY TABLE 4. Continued

					Ref size	Pre-dilatation	Stent size	Post- dilatation			DAPT therapy	Clinical	
Case	Device	Initial PCI Indication	Treated Vessel	Lesion	(mm)	(atm)	(atm)	(atm)	Initial DAPT therapy	Days to DT	Time of DT	outcome (worst)	Patients note
			Mid LAD	B2	3.0x45	2.5x20 (14)	2.5x23 (16)	4.0x15 (18)	ASA		7		
10	Absorb BVS	NSTEMI					3.0x28 (18)		Ticagrelor	46	Clopidogrei	Myocardial infarction	Malapposition stent (OCT)
	) -		Prox LAD	B1	4.0x15	2.5x20 (14)	3.5x18 (18)	4.0x15 (18)	OAC		OAC		
Ξ	Absorb	GATT	G 4 1 1 2 3 4	10	0.13	(01) 21:36	2 0-10 (13)	M	ASA	70	V S V	Myocardial	Interaction
:	BVS	OVE	UNING TWI	DI	2.0x12	7.3XI (10)	3.0x10 (12)	ONI INO	Ticagrelor	00	VCV	infarction	HIV medication
									ASA		Clonidogral	Non-	
12	Absorb RVS	NSTEMI	Prox RCA	B1	3.5x10	3.0x15 (12)	3.5x12 (14)	3.5x8 (22)	Clopidogrel	100	Ciopidogra	followed	
									OAC		OAC	by cardiac death	
-	Absorb	TIAP	Mid I AD	18	3 5 15	2 0×15 (18)	3 5×18 (10)	3 5415 (16)	ASA	191	None	Myocardial	DAPT cessation
Çī	BVS	r con	IMIN EAST	<u>.</u>	0.7417	(01) (1007	3,2010 (10)	7.7417 (10)	Ticagrelor	101	זאסווס	infarction	during surgery
7	Absorb	NSTEMI	Prov B.CV	R2	3 0~78	2 5~15 (12)	3.0~28 (1.6)	3 5215 (14)	ASA	185	None	Myocardial	DAPT cessation
ř.	BVS	HAT TON	I IOX NCA	70	0.0020	(21) (17)	7.0220 (14)	J.JA1 (14)	Ticagrelor	10)	TADIIG	infarction	during surgery
7	Absorb	STFMI	MidIAD	R	2 5x23	2 0×20 (14)	2 5×23 (14)	2 5v15 (18)	ASA	234	ASA	Myocardial	
3	BVS			5	C7VC :-7	(11) 0700.7	(F1) C2V(-7	(01) (10)	Ticagrelor	107	Ticagrelor	infarction	
71	Absorb	ΔD	Dov OM	B1	2 513	2 515 (9)	0) 52.18	Ž	ASA	3/40	ΔSΑ	Myocardial	History of
01	BVS	7.	McA, OIVI	10	4. JA14	(9) (18(:7	7: JAIO (0)	ONT	Ticagrelor	(1.7	Wew	infarction	compliance
17	Absorb	NCTEMI	Drow D.V	ВЭ	2 515	2.5x15 (8)	0 55.18 (14)	0175715	ASA	357	ASA	Myocardial	Dissection after
/1	BVS	HATT I CAT	1108 NCA	70	7. JAI.)	Rotablation	(+1) 010(.7	(10) (10)	Ticagrelor	700	Vev	infarction	(angio)
0	Absorb	ΔĐ	Mid RCA	B2	3.5x25	2.5x20 (12)	3.5x28 (12)	4.0x15 (10)	ASA	376	V S V	Myocardial	Malapposition
01	BVS	117	Distal RCA	B2	3.0x15	2.5x20 (12)	3.0x18 (14)	No	Ticagrelor	0/0	17017	infarction	distal stent (OCT)
9	Absorb	CTTENAT	, C	5	70-0	(01) 00-00	0 0-04	(01) 212 6	ASA	710	۷ ۷	Myocardial	
19	BVS	O I EIVII	Distal RCA	P7	5.0x24	7.0x20 (10)	3.0x2/ (8)	3.5X15 (18)	Ticagrelor	419	ASA	infarction	

SUPPLEMENTARY TABLE 4. Continued

		i de la companya de l	F		Ref size	Pre-dilatation	Stent size	Post- dilatation	Town I want	ć	DAPT therapy	Clinical	
Case	Device	Initial FCI Indication	Vessel	type	(mm)	(atm)	(atm)	(atm)	therapy	Days to DT	Time of DT	(worst)	Patients note
									ASA				
20	Absorb BVS	AP	Dist RcX	B1	3.0x10	3.0x15 (18)	3.0x18 (12)	°Z	Ticagrelor	427	OAC	Myocardial infarction	
									OAC				
									ASA			;	
	-								Prasugrel			Non- fatal MI	OAC cessation
21	Absorb BVS	STEMI	Mid RCA	B2	3.5x23	3.5x20 (10)	3.5x28 (12)	3.5x15 (12)	OAC	430	None	followed	during surgery
									ASA stop after 3 months			death	(2000)
6	Absorb	Angio-	Prox RCA	B1	4.0x16	3.0x20 (16)	3.5x28 (16)	4.0x20 (12)	ASA	107	-	Myocardial	
77	BVS	driven	Prox RcX	Α	3.5x12	3.0x12 (14)	3.0x23 (16)	3.5x40 (16)	Clopidogrel	42/	Unknown	infarction	
;	Absorb	STEMI	Prox RCA	B2	2.5x15	2.5x15 (10)	3.0x18 (12)	No	ASA	5	V 3 V	Myocardial	
C7	BVS	Staged	Prox RcX	B1	$3.0 \times 12$	3.0x15 (10)	3.0x18 (14)	No	Ticagrelor	401	ASA	infarction	
, c	Absorb	ΑĐ	Distal LAD	B1	3.0x28	2.5x28 (14)	3.0x28 (14)	3.0x28 (14)	ASA	471	ASA	Myocardial	
<del>1</del> 7	BVS	7	Prox LAD	А	3.5x12	3.0x12 (14)	3.5x12 (14)	3.5x14 (14)	Ticagrelor	4/1	Ticagrelor	infarction	
35	Absorb	CTEMI	Drow DCA	C	2 5, 10	3.0%15 (12)	3 5,73 (16)	6 0x30 (16)	ASA	295	ASA	Myocardial	
(7	BVS	31 E1VII	FIOX NCA	ر	3. JX 10	3.0x1 C 1x0.C	3.3x23 (10)	4.0xz0 (19)	Prasugrel	/00′	VSV	infarction	
26	Absorb	STEMI	Mid RCA	B2	3.0x25	3.0x15 (12)	3.0x28 (10)	2.25x20 (13)	ASA	593	ASA	Myocardial	
									Licagrelor				
7.0	Absorb	CTEMI	Prov I AD	ر	3 5071	2 5×20 (10)	3 5~23 (18)	3 5×15 (18)	ASA	733	ASA	Myocardial	Patient refused
ì	BVS		T VOI	,	7:7451	(10)	(11) CTV (10)	(10)	Ticagrelor	CC /	T TOTAL	infarction	re-start DAPT
80	Absorb	NSTEMI	AO-MO	R2	3.0v18	2 (V) (2 (12)	3 0×18 (10)	3 0×12 (14)	ASA	692	ASA	Myocardial	
2	BVS		graft		0.000	(71) (17)	0.0410 (10)	0.0012 (1.1)	Clopidogrel	6	YOU	infarction	

SUPPLEMENTARY TABLE 4. Continued

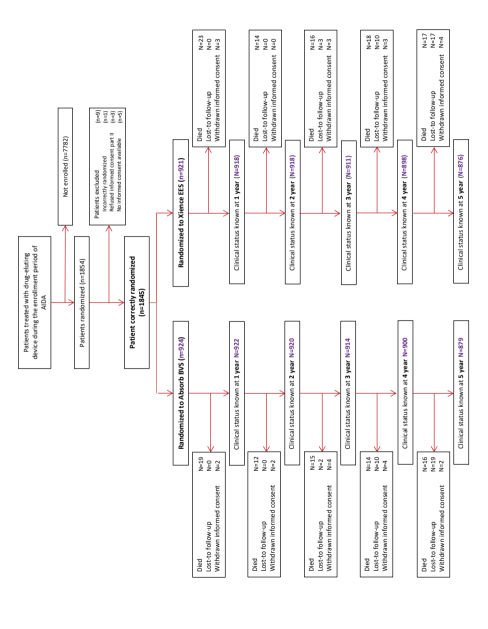
					,								
		Initial PCI	Treated	Lecion	ket size	Pre-dilatation	Stent size	Post- dilatation	Initial DAPT	Days	therapy	Clinical	
Case	Device	Indication	Vessel	type	(mm)	(atm)	(atm)	(atm)	therapy	to DT	Time of DT	(worst)	Patients note
29	Absorb	AP	Prox LAD	∢	3.5x8	3.0x15 (12)	2.5x12 (12)	3.5x8 (20)	ASA	817	ASA	Myocardial	Malapposed non-covered struts
	BVS								Clopidogrel			infarction	distally (OCT)
30	Absorb BVS	NSTEMI	RcX, MO	B1	2.5x10	2.5x15 (20)	2.5x12 (16)	2.75x15 (18)	ASA	825	ASA	Myocardial infarction	
			Mid RcX	B1	3.0x28	2.5x15 (14)	3.0x28 (16)	3.0x28 (18)					RcX prox
31	Absorb BVS	NSTEMI	Prox RCA	B1	3.5x18	2.0x20 (10)	3.5x18 (8)	4.0x12 (20)	ASA	1223	ASA	Post	occluded; TLR Cx Xience at 790
			Distal RCA	B1	2.5x28	2.0x20 (10)	2.5x29 (12)	3.0x12 (10)	Ticagrelor				days
					2.5x45	2.5x30 (12)	2.5x28 (16)	No					
	-	÷	Mid LAD	C	2.5x45	2.5x30 (12)	2.5x28 (16)	No	ASA			-	
32	Absorb BVS	Stabilized			2.5x45	2.5x30 (12)	2.5x18 (16)	°N		1,277	Unknown	Myocardial infarction	DT in LAD
			First Diagonal	B2	3.5x12	3.5x15 (16)	3.5x12 (14)	4.0x9 (14)	Ticagrelor				
23	Absorb	Ę	Distal RcX	B1	3.5x18	3.0x15 (12)	3.5x18 (14)	3.5x12 (16)	ASA		40 <b>4</b>	Myocardial	TLR with Xience
cc	BVS	J.V.	Prox LAD	B1	3.0x18	2.5x15 (12)	2.5x18 (12)	2.5x12 (16)	Ticagrelor	1,312	ASA	infarction	at 2 days
	Abcock		Mid LAD	B1	3.5x15	3.0x15 (12)	3.5x18 (12)	4.0x15 (12)	ASA			Mysocondial	DT in LAD, TLR
34	BVS	UAP	First Diagonal	B2	2.5x10	2.5x10 (10)	2.5x12 (12)	°Z	Clopidogrel	1,330	ASA	infarction	with Xience at 668 days
,	Absorb	4	Prox RCA	C	3.5x23	2.5x15 (12)	3.5x23 (14)	3.5x12 (16)	ASA		•	Myocardial	Acute DT on
ÇÇ	BVS	OAL	Distal RCA	C	2.5x23	2.5x15 (10)	2.5x23 (12)	2.5x12 (16)	Ticagrelor	1,5,1	ASA	infarction	severe scatfold- restenosis
36	Absorb BVS	AP	Prox LAD	B1	3.0x18	2.5x20 (20)	3.0x23 (20)	3.0x15 (20)	ASA	1,506	Unknown	Myocardial	Subtotal lesion RcX proximal of
	Xience EES		Mid RcX	B1	2.5x15	2.0x20 (16)	2.5x23 (16)	°N	Clopidogrel			ınfarction	stent
37	Absorb BVS	STEMI	LAD Mid	CI	3.0x20	2.0x20 (10)	3.0x23 (18)	3.0x15 (16)	ASA Ticagrelor	1,711	ASA	Myocardial infarction	

SUPPLEMENTARY TABLE 4. Continued

Case	Case Device	Initial PCI Indication	Treated Vessel	Lesion	Ref size (mm)	Pre-dilatation (atm)	Stent size (atm)	Post- dilatation (atm)	Initial DAPT therapy	Days to DT	DAPT therapy Time of DT	Clinical outcome (worst)	Patients note
38	Absorb BVS	NSTEMI	LAD Mid	B1	3.3x15	3.0x15 (10)	3.0x18 (12)	3.5x15 (12)	ASA	1,798	ASA		ST in D1
	Xience EES		DI	B2	3.0x20	3.0x15 (6)	3.0x23 (10)	°Z	Ticagrelor				

SUPPLEMENTARY TABLE 5. Descriptive characteristics of cases of definite device thrombosis

		Da Pisi-1	,	1	Refsize	Pre-dilatation	Stent size	Post-dilatation	Initial		DAPT therapy	Clinical	
Case	Device	Indication	Vessel	type	(mm)	(atm)	(atm)	(atm)	therapy	to DT	Time of DT	(worst)	Patients note
1	Xience	Stabilized	Mid RCA	B2	3.5x15	No	3,5x10 (18)	3.5x18 (14)	ASA Ticagrelor	0	ASA Ticagrelor	Myocardial Infarction	
		STEMI	Dist RCA	C	2.5x25	2.5x20 (14)	2,75x28 (14)	2.5x15 (8)					
7	Xienœ	STEMI	Prox LAD	B2	3.0x28	3.0x20 (6)	3.0x38 (14)	3.5x15 (12)	ASA Ticagrelor	0	ASA Ticagrelor	Myocardial Infarction	
8	Xienœ	STEMI	Prox LAD	B2	3.5x15	3.0x15 (16)	3.5x15 (12)	°Z	ASA Ticagrelor		ASA Ticagrelor	Myocardial Infarction	Jailing stent (Angio)
4	Xienœ	AP	Distal RcX	A	3.0x15	2.5x15 (10)	3.0x18 (12)	°Z	ASA Clopidogrel	6	ASA Clopidogrel	Myocardial Infarction	
<b>ν</b>	Xienœ	STEMI	Prox RCA	B2	3.0x15	3.0x15 (10)	3.0x12(16)	°Z	ASA Prasugrel	511	ASA	Myocardial Infarction	Malapposition proximal stent-strut (OCT)
9	Xienœ	STEMI	Mid LAD	B1	3.0x16	3.0x15 (6)	3.0x18 (12)	°Z	ASA Ticagrelor	1,222	ASA	OHCA	
_	Xienœ	UAP	Distal RCA	B1	$3.0 \times 10$	2.5x10 (13)	3.0x15 (14)	No	ASA Clopidogrel	1,391	ASA Clopidogrel	Myocardial Infarction	TLR at 176 days
∞	Xienœ	AP	Mid LAD	B1	3.0x15	2.5x12 (10)	3.0x18 (12)	No	ASA Prasugrel	1,472	ASA	Myocardial infarction	
6	Xienœ	STEMI	RCA mid	A1	3.0x10	2.5x15 (18)	3.0x12 (18)	No	ASA	1,792	none	Myocardial infarction	ST in MO1
			MO1	A1	2.5x12	2.5x15 (16)	2.5x15 (16)	°Z	Ticagrelor				Cessation NOAC 1 month prior



SUPPLEMENTARY FIGURE 1. Study Flowchart