Procedural Safety and Long-Term Clinical Outcomes in Patients Receiving Ultra-Long Everolimus-Eluting Stent: A Single-Center Real-World Experience

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Abstract

Background: Diffuse long coronary lesions are difficult to treat percutaneously. The aim of the present study was to assess the procedural safety and long-term efficacy of the ultra-long (48-mm) drug-eluting stent Xience Xpedition.

Methods: This was an investigator-initiated, observational, all-comers study. A total of 92 patients with 93 lesions were enrolled in the study from October 2016 to October 2020. The primary outcome of the study was major adverse cardiac events (MACEs). Secondary outcomes were individual components of the primary outcome and procedural success.

Results: The mean (standard deviation (SD)) age of the participants was 58.8 (10.8) years. More than half of the patients had ST-segment elevation myocardial infarction (STEMI) at presentation (55.4%). Ten patients were in cardiogenic shock (CGS; 10.8%). Most of the lesions were located in the left anterior descending artery (48.3%). American College of Cardiology/American Heart Association (ACC/AHA) type C was the most common lesion type amongst the intervened vessels (46.74%), with a mean syntax score (SD) of 16.99 (8.89). The mean stent diameter used was 2.77 mm (0.25). MACE was observed in 7.6% of patients studied at a median follow-up of 24 months. MACE was significantly lower in the population without CGS, occurring in only 2.4% of the patients; a significant difference in MACE was observed in patients with and without CGS (P < 0.001). Procedural success was obtained in 89.2% of total population; however, 96.3% of patients without CGS had procedural success.

Conclusions: The deployment of the ultra-long 48-mm Xience Xpedition stent is feasible, safe, and effective; and it was associated with a good intermediate-term clinical outcome.

Keywords: 48-mm stent; Xience; Percutaneous coronary intervention; Coronary artery disease

Introduction

Diffuse coronary artery lesions account for 20% of all percutaneous coronary intervention (PCI) [1]. Many such complex lesions are addressed percutaneously, leading to an increasing need for techniques to handle them. For example, the use of longer balloons followed by deployment of longer stents or multiple stents with overlapping segments. However, stent overlap results in increased neointimal proliferation, lumen loss due to delayed healing, disparity in drug deposition, and increased inflammation. Furthermore, the overlapping portions make the vessel rigid due to excess metal, which may lead to stent fracture and cause further vascular injury, leading to a higher incidence of restenosis [2]. The length of the stents (drug-eluting stents (DES)) is directly proportionally to the incidence of in-stent restenosis [3]; a study done by Cassese et al showed that lesion length was independently associated with restenosis (odds ratio (OR): 1.27; 95% confidence interval: 1.21 - 1.33, for every 10 mm increase in the length of the stent) [4].

Everolimus-eluting stents (EESs) are more effective than old-generation DES when used in long lesions [5]. Until re-
cently, the longest stent approved by the US Food and Drug Administration (FDA) was the 38-mm DES (stents greater than or equal to 38 mm are considered ultra-long stents). However, a 48-mm EES with platinum-chromium platform with an absorbable polymer has now been approved. The Drugs Controller General of India (DCGI) has approved the 48-mm Xience Xpedition stent (Abbott Vascular, Santa Clara, California). However, there is no long-term data available regarding the clinical outcomes of the ultra-long 48-mm Xience Xpedition stent. In this study, we analyzed the intermediate-term clinical outcomes and immediate procedural success of this device.

Materials and Methods

Study design

This was an investigator-initiated, observational, all-comers study. The present study included all consecutive patients who had undergone PCI using the 48-mm Xience Xpedition stent from October 2016 to October 2020 in the Department of Cardiology, Sri Ramchandra Institute of Higher Education and Research (SRIHER). Xience Xpedition (48 mm) is an L-605 cobalt-chromium alloy stent pre-mounted on a rapid exchange delivery system. It is available in different sizes (2.5 mm, 2.75 mm, 3.00 mm, and 3.5 mm). The abluminal surface of the device is coated with a thin layer of durable fluoropolymer, which is used as the carrier for everolimus, at a concentration of 100 µg/mm² [6].

The details of de-identified patients were collected from the Cardiac Catheterization Registry and patient database software (Health Management Information System - HMIS, SRIHER). Baseline characteristics of the patients, as well as angiographical and interventional details, were recorded. The recruited patients were contacted during the follow-up by the respective physicians or by the second author. There was no interference in the routine care of the included patients.

Informed consent was obtained from all participants, and the patients who did not give consent were excluded from the study. The study was approved by the institute’s ethics committee. This study was conducted in compliance with the ethical standards of the responsible institution on human subjects as well as with the Helsinki Declaration.

Outcomes

The primary outcome of the study was major adverse cardiac events (MACEs) which consisted of cardiovascular death, nonfatal myocardial infarction (MI), and repeat intervention in the target vessel.

Cardiovascular death was defined according to the definitions developed by the American College of Cardiology/American Heart Association (ACC/AHA) task force for cardiovascular endpoints in clinical trials [7].

Nonfatal MI was defined by the elevation of cardiac troponin (cTn) values five times the 99th percentile upper reference limit (URL) in patients with normal baseline values as per the fourth universal definition of MI [8]. Similarly, reintervention included target vessel revascularization.

Secondary outcomes included individual components of the composite primary outcome and procedural success, which was defined as the successful deployment of the stent without any procedural complications such as cardiac death, dissection, and stent thrombosis. In addition, we also assessed admission for heart failure, stroke, cardiovascular procedures, and other cardiovascular causes.

Statistical analysis

Descriptive data were expressed in terms of ratio, proportion, or percentage. Mean and median values (interquartile range) were used for discrete quantitative data. Continuous variables were analyzed by the t-test. Categorical variables were analyzed by the Chi-squared test. A Kaplan-Meier curve survival analysis was performed. A P value < 0.05 was considered significant. SPSS Version 20 (IBM) was used for statistical analysis.

Results

A total of 92 patients with 93 lesions were enrolled in the present study, with a median follow-up of 24 months. Baseline characteristics of the patients are described in Table 1. The mean (standard deviation (SD)) age of the participants was 58.8 (10.8) years; 26.1% of the participants were female. More than half of the patients had ST-segment elevation myocardial infarction (STEMI) at presentation (55.4%). The mean ejection fraction (SD) was 51% (0.1%). Ten patients in the study population were in cardiogenic shock (CGS; 10.87%).

The left anterior descending (LAD) artery was the most intervened vessel (48.38%), followed by the right coronary artery (RCA) (45.18%), left circumflex (LCX) (3.22%), and left main (LM) (3.22%). Lesions were classified based on the ACC/AHA guidelines; all lesions were ACC/AHA type C lesions. Approximately 11% (n = 10) of patients had severely calcified lesions, and 20.7% (n = 19) had severe tortuosity in the vessels. Two-thirds of the patients (66.30%) had multivessel involvement. The mean syntax score (SD) of the patients was 16.99 (8.89). The Xience Xpedition 48-mm stent was used for primary PCI in 25% (n = 23) of patients. The mean diameter (SD) of the stent used was 2.77 (0.25) mm. Most of the lesions were predilated (98.91%) with semicompliant or noncompliant balloons. The ultra-long stents were post-dilated in 88.04% of patients with noncompliant balloons. Rotational atherectomy and image guidance using intravascular ultrasound were employed in 2.17% of patients. All the patients received dual antiplatelet therapy (DAPT) for a minimum duration of 12 months with all patients receiving aspirin 75 mg and the majority receiving ticagrelor (82.61%, n = 76).

Outcomes

The primary outcome of the study, MACE, was observed in
7.61% (n = 7) of patients at a median follow-up of 24 months. MACE occurred in five patients who had CGS (50%) as compared with those without CGS (2.4%, n = 2, P ≤ 0.001) The significant difference regarding the occurrence of MACE between the CGS and non-CGS groups was influenced by death at admission.

A total of eight deaths were reported in the present study population (8.7%), of which five deaths occurred in the CGS group. Out of the five patients who died with CGS, four patients (three of which had multivessel disease and one which had single-vessel disease) presented with acute MI and severe left ventricular (LV) dysfunction (ejection fraction less than 30%), and were taken up for primary PCI. LAD was the culprit lesion in three patients, and RCA was the culprit lesion in one patient. These patients had refractory CGS with multiple organ dysfunction syndrome and succumbed to their illness within 24 h of the procedure. The fifth patient had undergone a procedure with the Xience Xpedition 48-mm stent in 2016 and had

Table 1. Baseline Characteristics and Angiographic and Interventional Details of the Patient Population

| Variable                              | Number (n) | Percentage (%) |
|---------------------------------------|------------|----------------|
| Age (mean ± SD)                       | 58.8 ± 10.8|                |
| Female sex                            | 24         | 26.10%         |
| Diabetes mellitus                     | 63         | 68.48%         |
| Hypertension                          | 40         | 43.48%         |
| Chronic kidney disease                | 3          | 3.26%          |
| Dyslipidemia                          | 6          | 6.52%          |
| Chronic coronary syndrome             | 21         | 22.83%         |
| STEMI                                 | 51         | 55.40%         |
| Mean ejection fraction                | 51%        | Min, 20%; Max, 65% |
| Cardiogenic shock                     | 10         | 10.87%         |
| ACC lesion type                       |            |                |
| B1                                    | 22         | 23.70%         |
| C                                     | 71         | 76.30%         |
| Lesion location                       |            |                |
| Left main                             | 3          | 3.22%          |
| LAD                                   | 45         | 48.38%         |
| LCX                                   | 3          | 3.22%          |
| RCA                                   | 42         | 45.18%         |
| Syntax score (mean ± SD)              | 16.99 ± 8.89|               |
| PCI details                           |            |                |
| Primary PCI with 48-mm stent          | 23         | 25%            |
| Single-vessel lesion addressed with 48-mm stent | 31 | 33.70%         |
| Multivessel disease addressed with 48-mm stent | 61 | 66.30%         |
| IVUS optimizing                       | 2          | 2.17%          |
| Rotational atherectomy                | 2          | 2.17%          |
| Predilation                           | 91         | 98.91%         |
| Stent diameter (mean ± SD), mm        | 2.77 (0.5) |                |
| Stent size: 2.5 mm                    | 29         | 31.52%         |
| Stent size: 2.75 mm                   | 34         | 36.96%         |
| Stent size: 3.0 mm                    | 25         | 27.17%         |
| Stent size: 3.5 mm                    | 4          | 4.35%          |
| Predilation of the lesion             | 92         | 98.91%         |
| Post-dilation of the lesion           | 81         | 88.04%         |

SD: standard deviation; STEMI: ST-segment elevation myocardial infarction; LAD: left anterior descending; RCA: right coronary artery; LCX: left circumflex; PCI: percutaneous coronary intervention; IVUS: intravascular ultrasound; Min: minimum; Max: maximum.
been doing relatively well till 2018 when he developed severe LV dysfunction CGS and succumbed to his illness. Three patients who did not have CGS died during the follow-up. Two of those three patients were doing well for 1 year after stent implantation but had a possible out-of-hospital cardiac death, and one patient had a noncardiac cause of death after colon surgery for carcinoma of the colon. Nonfatal MI of the target vessel and target vessel revascularization were not seen in any of the patients. Non-target vessel revascularization was seen in 4.35% (n = 4) of patients.

Procedural success was seen in 89.2% of the overall population and in 96.3% of patients without CGS. As mentioned in Table 2, a total of nine patients developed complications such as dissection (n = 4), intraprocedural stent thrombosis (definitive stent thrombosis; n = 1), and immediate cardiac death (n = 4). All the patients who died had CGS. Also, the patient who had intraprocedural stent thrombosis was also in CGS. He presented with STEMI with CGS; he had definitive evidence of stent thrombosis (1.08%) as he developed chest pain after the procedure. An immediate angiogram revealed stent thrombosis due to edge dissection. This was addressed by placing an additional stent. A total of four patients had dissections (4.34%) within 3 mm of the stented segment in the LAD. All of them were treated with an additional stent. The procedural success is significantly higher in patients without CGS.

Kaplan-Meier survival curve analysis

We performed a Kaplan-Meier curve analysis to analyze survival in the patient groups. As expected, the survival was lower in patients with CGS compared to those without CGS (Fig. 1). We observed a step-down in the survival in the non-CGS group between 18 and 20 months; however, we did not find any further events till the end of 60 months.

Table 2. Outcomes of Patients Who Received the Ultra-Long 48-mm Xience Xpedition Stent

| Characteristics                                      | Number, n (%) |
|------------------------------------------------------|---------------|
| MACE in all the patients studied                     | 7 (7.61)      |
| MACE in patients without cardiogenic shock           | 2 (2.44)      |
| Total deaths                                         | 8 (8.7)       |
| Noncardiac death                                     | 1 (1.07)      |
| Cardiac death in all the patients studied            | 7 (7.61)      |
| Cardiac death in patients with cardiogenic shock     | 5 (55.5)      |
| Nonfatal MI in the target vessel                     | 0 (0)         |
| Target vessel revascularization                      | 0 (0)         |
| Non-target vessel revascularization                  | 4 (4.35)      |
| Procedural success in all the patients               | 83 (89.2)     |
| Procedural success in patients without cardiogenic shock | 79 (96.3)  |
| Complications                                        | 9 (9.78)      |
| Dissection                                           | 4 (4.34)      |
| Intraprocedural stent thrombosis (definitive stent thrombosis) | 1 (1.07) |

MACE: major adverse cardiac event; MI: myocardial infarction.

Discussion

In our all-comers study, we found that the intermediate-term clinical outcome of Xience Xpedition 48-mm ultra-long stent was very good. The ultra-long 48-mm Xience Xpedition stent was effective even when used during acute coronary events. The strengths of our study are its prospective design, which included all-comers with variable clinical presentation, and the at least intermediate-term follow-up period.
The incidence of long coronary lesions has increased with the increased prevalence of comorbid conditions [9]. Previously, patients with such lesions were excluded from clinical studies. Finn et al performed animal studies to identify the arterial reaction to first-generation DES vs. bare-metal stents (BMS) [10]. First-generation DES demonstrated delayed healing and promoted inflammation at the overlap sites when compared with the BMS. More luminal heterophilis/eosinophilis and fibrin deposition were observed with Taxus stents than Cypher stents [10]. Second-generation DES, such as Xience (Abbott Vascular, Santa Clara, California), Promus (Boston Scientific, MA, USA), and Endeavor Resolute (Medtronic, Minneapolis, USA), have thinner stent struts, and have more biocompatible polymers compared with first-generation DES. Such technological advancements have resulted in a reduction in vascular injury, inflammation and rapid endothelialization [11]. In the porcine studies done by Farooq et al, longer lesions intervened with overlapping everolimus stent platforms have shown to have good overlap endothelialization within 30 days [12].

A meta-analysis involving 13,266 patients undergoing PCI for very long lesions (>35 mm) with overlapping stent treatment with the EES (XIENCE V) from six trials (Spirit II, III, IV, V, Spirit Small Vessel, and XIENCE V USA) demonstrated that there was no difference in outcomes such as target lesion failure (8.9% vs. 10%; P = 0.63) and MACE (9.2% vs. 10%; P = 0.74) compared to the control group comprising lesions > 24 to < 35 mm. The longest stent used in these studies was a 38-mm stent. They concluded that XIENCE V (EES) appeared safe and effective for PCI for long lesions [13]. All the above-described studies used an EES of size less than or equal to 38 mm, and longer lesions were addressed by an overlapping stenting technique.

The only trial available till date for the treatment of very long coronary lesions with a single EES (Xience Xpedition 48-mm stent) was performed by Tan et al [6]. A total of 123 patients were followed up for 12 months. The study included patients presenting with chronic coronary syndrome (CCS) (n = 51) and acute coronary syndrome (ACS) (n = 72). The included patients presented with a wide spectrum of coronary anatomies and clinical presentations. Also, it is a single-center study; whether the results are applicable to other centers is questionable.

Conclusions

On the basis of our all-comers study, which included a small sample size, we conclude that the deployment of the ultra-long Xience Xpedition 48-mm stent is feasible, safe, and effective with a good intermediate-term clinical outcome.

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This study did not receive any funding in any form.

Conflict of Interest

The authors declare that there is no conflict of interest.

Informed Consent

Informed consent was obtained from all participants

Author Contributions

NBS: conceptualization, designing, interpretation supervision, writing of original draft, review and editing. RK: execution,
interpretation, review and editing. Ravi Shankar P: statistics. SS, NVS, SKP, BVK, MI, MR, VS, JVB, PK, SR, PM, TRM and JSM: reviewing and editing. ST: guarantor, reviewing and editing.

**Data Availability**

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

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