1	Sirolimus-eluting Stent Implantation for Ostial Left Anterior Descending
2	Coronary Artery Lesions: Three-Year Outcome from the j-Cypher Registry
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35 Abstract

Background: Ostial left anterior descending coronary artery (LAD) lesion has been regarded as a
 lesion subset unsuitable for coronary stenting. Long-term outcomes of sirolimus-eluting stent (SES)
 implantation for ostial LAD lesions have not been yet adequately evaluated.

39 Methods and Results: Among 12824 patients enrolled in the j-Cypher registry, 3-year outcomes 40 were compared between 481 patients with SES-treated ostial LAD lesions and 5369 patients with 41SES-treated non-ostial proximal LAD lesions. Patients with ostial LAD lesions, as compared with 42patients with non-ostial proximal LAD lesions, had similar incidences of target lesion 43revascularization (TLR) (9.4% vs. 9.7%, p=0.98; adjusted hazard ratio (HR) 0.99 (95% confidence interval (CI): 0.7-1.36), p=0.94) and death/myocardial infarction (MI) (10.7% vs. 11.4%, p=0.82; 44adjusted HR 1.05 (95%CI: 0.76-1.4), p=0.77), respectively. Among 481 patients with ostial LAD 4546lesions, patients undergoing both main- and side-branch stenting (62 patients), as compared with main-branch stenting alone (419 patients), had higher risk for TLR (adjusted HR 4.65 (95%CI: 472.32-9.25), p < 0.0001) but had similar risk for death/MI (adjusted HR 1.15 (95%CI: 0.49-2.41), 4849p=0.73). In patients with main-branch stenting alone, outcomes after crossover-stenting across 50circumflex (225 patients) were not different from those after ostial-stenting (194 patients) (adjusted 51HR 0.77 (95%CI: 0.33-1.82), p=0.55 for TLR, and adjusted HR 1.54 (95%CI: 0.78-3.2), p=0.22 for 52death/MI).

53 Conclusions: In terms of both safety and efficacy, 3-year outcomes of PCI using SES for ostial LAD
54 lesions were comparable to those for non-ostial proximal LAD lesions. Crossover-stenting with

55 one-stent approach might be a reasonable option in treating ostial LAD lesions.

56

57 Key words: Coronary artery disease, Stent, Restenosis, Thrombosis

58 Text

The ostial left anterior descending coronary artery (LAD) lesion is an important target for coronary 59revascularization, since this lesion location subtends a large territory of myocardium. However, the 60 61 ostial LAD lesion has been regarded as a lesion subset unsuitable for percutaneous coronary intervention (PCI) because of frequent atherosclerotic involvement of distal left main coronary 62 63 artery (LMCA) and because of concerns for compromising the circumflex coronary artery (LCX). 64 Furthermore, restenosis rate after implantation of bare-metal stents (BMS) for ostial LAD lesions remained high, ranging from 26% to 33%^{1,2)}. Although randomized controlled trials comparing 6566 drug-eluting stents (DES) with BMS demonstrated significant reduction in the rates of target-lesion 67 revascularization (TLR) with use of DES, ostial LAD lesions have been excluded from most of these randomized controlled trials. Despite increasingly frequent use of DES for the treatment of 68 ostial LAD lesions, its long-term outcome has not been yet adequately evaluated $^{3-5)}$. The current 69 70analysis was conducted to evaluate 3-year clinical outcomes of patients who underwent 71sirolimus-eluting stents (SES) implantation for ostial LAD lesions in a large cohort of patients 72enrolled in the j-Cypher registry.

- 73 Methods
- 74

Study Design and Patient Population

The study design for the j-Cypher registry was previously described ⁶⁾. In brief, the j-Cypher registry is a physician-directed, prospective, multicenter registry in Japan enrolling consecutive patients undergoing SES implantation without any exclusion criteria (Supplemental

Appendix A). While a center actively enrolled patients, technicians in the catheterization laboratory 78registered all the patients undergoing PCI in a screening log. When SES implantation was 7980 undertaken, the patient was invited to participate in the j-Cypher registry. Although data entry was 81 basically left to the individual sites, the experienced clinical research coordinators (Supplemental Appendix B) in the data management center supported data entry when necessary. Logical 82 83 inconsistencies were resolved by inquiries to the site investigators and/or by audits against the 84 original data sources. Follow-up data were obtained from hospital charts or by contacting patients 85and/or referring physicians at 30 days, 6 months, one year and yearly thereafter. When death, 86 myocardial infaction (MI), and stent thrombosis (ST) were reported, the events were adjudicated using the original source documents by a clinical events committee (Supplemental Appendix C). 87 Adjudication of TLR events was left to the decision of the local investigators. The relevant review 88 89 boards in all 37 participating centers approved the study protocol. Written informed consent was 90 obtained from all patients enrolled.

The current pre-specified sub-analysis from the j-Cypher registry was intended to evaluate safety and efficacy of SES use in patients with ostial LAD lesions. Among 12824 patients enrolled in the j-Cypher registry from August 2004 to November 2006, 6230 patients underwent PCI for proximal LAD disease. Excluding 380 patients in whom proximal LAD lesions were treated by modalities other than SES, the current study population consisted of 5850 patients whose proximal LAD lesions were treated exclusively with SES. Baseline characteristics and clinical outcomes were compared between 481 patients with ostial LAD lesions, and 5369 patients with non-ostial proximal

98	LAD lesions. Subgroup analysis was also conducted in 481 patients whose ostial LAD lesions were
99	treated exclusively by SES. Baseline characteristics and clinical outcomes were compared between
100	main-branch stenting alone (one-stent approach; 419 patients) and both main- and side-branch
101	stenting (two-stent approach; 62 patients). Furthermore, in patients with one-stent approach, baseline
102	characteristics and clinical outcomes were compared between crossover stenting across LCX
103	(crossover-stenting; 225 patients) and stenting just at the ostium of LAD (ostial-stenting; 194
104	patients) (Figure 1).

105 **Definitions**

106 A "lesion" was defined as the area covered by single or multiple overlapping stents. When 107 two stents were placed without overlap, these two areas were regarded as two separate lesions. Ostial 108 lesion was defined as a narrowing located within 3 mm of the vessel origin in the least foreshortened 109 angiographic projection. Those ostial LAD lesions with concomitant significant LMCA distal bifurcation 110 stenosis were regarded as LMCA lesions and were excluded from the current analysis. Proximal LAD 111 was defined as the segment of LAD proximal to the first major septal branch. Techniques of stenting were pre-specified and recorded in the case report forms during the index stent implantation procedures. 112113Crossover-stenting was defined as stent placement from distal LMCA to LAD across LCX, while 114ostial-stenting as the stenting strategy with an intention not to protrude the stent into LMCA. 115One-stent approach meant stenting of LAD only (including crossover-stenting and ostial-stenting) 116 and two-stent approach denoted stenting of both ostial LAD and ostial LCX. Choice of the stenting 117strategies was left to the discretion of the operators.

118	The primary outcome measure for efficacy in the current analysis was defined as TLR for
119	the index proximal LAD lesions. TLR was defined as either PCI or coronary artery bypass grafting
120	(CABG) surgery due to restenosis or thrombosis of the target lesion that included the proximal and
121	distal edge segments as well as the ostium of the side branches. The composite of death or MI was
122	selected as the primary outcome measure for safety. Death was regarded as cardiac in origin unless
123	obvious non-cardiac causes could be identified. Any death during the index hospitalization was
124	regarded as cardiac death. Sudden death was defined as unexplained death in previously stable
125	patients. MI was adjudicated according to the definition in the Arterial Revascularization Therapy
126	Study ⁷⁾ . ST was defined according to the Academic Research Consortium (ARC) definition ⁸⁾ .

127 Statistical Analysis

128Categorical variables are presented as counts and percentages, and were compared with the 129chi-square test. Continuous variables were expressed as mean value \pm SD unless otherwise indicated. 130 Continuous variables were compared with the Student t test or Wilcoxon rank sum test on the basis 131of their distribution. Cumulative incidences of events were estimated by the Kaplan-Meier method, 132and curves were compared with the log-rank test. A multivariable Cox proportional hazard model 133was developed to adjust the differences in baseline characteristics. Proportional hazard assumptions 134for variables were assessed on the plots of log (time) versus log [-log (survival)] stratified by the variables, and were found justified. For the multivariable analysis, we first selected variables with p 135values < 0.1 in the univariate Cox models among 21 independent variables used in the previous 136 report⁶⁾. In the final multivariable model, we incorporated ostial LAD vs. non-ostial proximal LAD, 137

138	or one-stent approach vs. two-stents approach, and crossover-stenting vs. ostial-stenting together
139	with those independent variables with multivariable p values < 0.05. Covariates used in the final
140	model for adjustment were indicated in Supplemental Tables 1-3. The results of the multivariable
141	analysis were expressed as adjusted hazard ratios (HR) and their 95% confidence intervals (CI).
142	Statistical analyses were conducted by two physicians (Kishi K and Kimura T) and a statistician
143	(Morimoto T) with the use of JMP 8.0 (SAS Institute Inc, Cary, NC) software. P values < 0.05 were
144	considered statistically significant.
145	Results
146	Baseline Characteristics: Ostial LAD vs. Non-ostial Proximal LAD
147	The baseline clinical characteristics were generally similar between the ostial LAD group
148	and the non-ostial proximal LAD group, although patients >= 80 years of age, patients with prior MI
149	and statin users were more prevalent in the ostial LAD group (Table 1-A). The baseline angiographic
150	and procedural data were significantly different between the two groups (Table 1-B). The ostial
151	LAD group had larger vessel size, resulting in use of stents and balloons with bigger sizes.
152	Directional coronary atherectomy (DCA) before stenting, intravascular ultrasound (IVUS), and post
153	dilatation were more frequently utilized in the ostial LAD group. Minimal lumen diameter (MLD)
154	post procedure was significantly larger in the ostial LAD group.
155	Clinical Outcomes: Ostial LAD vs. Non-ostial proximal LAD
156	The follow-up interval in surviving patients was significantly longer in patients with ostial
157	LAD lesions (median: 995 days; interquartile range (IQR): 732 to 1095 days) than in patients with

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non-ostial proximal LAD lesions (median: 904 days; IQR: 730 to 1095 days) (P=0.02). Follow-up at

160 Cumulative incidence of TLR in the ostial LAD group was not different from that in the 161 non-ostial proximal LAD group (9.4% vs. 9.7%, p=0.98) (Table 2 and Figure 2-A). Adjusted 162 hazard ratio of ostial LAD vs. non-ostial proximal LAD for TLR was 0.99 (95% CI: 0.7-1.36, 163 p=0.94). Similarly, cumulative incidences of death or MI were not significantly different between 164 the two groups (10.7% vs. 11.4%, p=0.82) (Figure 2-B). Adjusted hazard ratio of ostial LAD vs. 165 non-ostial proximal LAD for death or MI was 1.05 (95% CI: 0.76-1.4, p=0.77).

166 Baseline Characteristics: One-stent vs. Two-stent approach

1 year was completed in 97% of patients.

167 The baseline clinical characteristics were generally similar between the one-stent approach group and the two-stent approach group, although patients >= 80 years of age were more prevalent 168169 in the two-stent approach group (Supplemental Table 4-A). The baseline procedural and 170 angiographic data were significantly different between the two groups. Crossover stenting approach 171and final kissing balloon technique were more frequently utilized in the two-stent approach group. 172The number and length of stents were greater in the two-stent approach group. Obviously, the 173prevalence of significant narrowing at the ostium of LCX was markedly higher in the two-stent 174approach group. Reference diameter (RD) and MLD of LCX before procedure were significantly 175smaller in the two-stent approach group than in the one-stent approach group. Despite these 176 differences in procedural and angiographic characteristics, post-procedural MLD in the main branch 177did not differ between the two groups. Final MLD of LCX was significantly larger in the two-stent approach group than in the one-stent approach group. (Supplemental Table 4-B)

179 Clinical Outcomes: One-stent vs. Two-stent Approach

180 Cumulative incidence of TLR in the two-stent group was significantly higher than that in 181 the one-stent group (28.1% vs. 6.6%, p<0.0001) (Table 3 and Figure 3-A). The adjusted hazard ratio of the two-stent approach vs. one-stent approach for TLR was 4.65 (95% CI: 2.32-9.25, p<0.0001). 182183 Cumulative incidences of stroke, CABG, and any coronary revascularization were also significantly 184 higher in the two-stent group than those in the one-stent group. However, cumulative incidences of 185death or MI were not significantly different between the two groups (16.8% vs. 9.8%, p=0.37) 186 (Table 3 and Figure 3-B). Adjusted hazard ratio of two-stent approach vs. one-stent approach for 187 death or MI was 1.15 (95% CI: 0.49-2.41, p=0.73).

188 Baseline Characteristics: Crossover-stenting vs. Ostial-stenting

189 Although the baseline clinical characteristics were generally similar between the 190 ostial-stenting group and the crossover-stenting group, the latter included more male patients and 191 patients with prior heart failure (Supplemental Table 5-A). The baseline procedural and angiographic 192 data were significantly different between the two groups. Final kissing balloon technique was more 193 frequently utilized in the crossover-stenting group, reflecting greater prevalence of significant 194 narrowing at the ostium of LCX. Although the crossover-stenting group had larger stent size, larger 195maximum balloon size and longer stent length, post-procedural MLD in the main branch did not 196 differ between the two groups. Final MLD of LCX was significantly smaller in the 197 crossover-stenting group than in the ostial-stenting group. (Supplemental Table 5-B)

198 Clinical Outcomes: Crossover-stenting vs. Ostial-stenting

199	Cumulative incidences of TLR were not significantly different between the crossover-stenting
200	group and the ostial-stenting group (5.4% vs. 7.9%, p=0.81) (Table 4 and Figure 4-A). Adjusted
201	hazard ratio of crossover-stenting vs. ostial-stenting for TLR was 0.77 (95% CI: 0.33-1.82, p=0.55).
202	Similarly, cumulative incidences of death or MI were not significantly different between the two
203	groups (12.2% vs. 7.0%, p=0.07) (Table 4 and Figure 4-B). Adjusted hazard ratio of
204	crossover-stenting vs. ostial-stenting for death or MI was 1.54 (95% CI: 0.78-3.2, p=0.22).
205	Although the crude incidence of all-cause death was significantly higher in the crossover-stenting
206	group (12.2% vs. 4.5%, p=0.01), the difference was no longer significant after adjusting
207	confounders (adjusted HR 2.04 [95% CI: 0.94-4.93, p = 0.07]) (Table 4).

208 Discussion

The main findings of the current analysis in the largest ever reported series of patients undergoing SES implantation for ostial LAD lesions are as follows: (1) In terms of both safety and efficacy, 3-year outcomes of PCI using SES for ostial LAD lesions were comparable to those for non-ostial proximal LAD lesions; (2) The two-stent approach, as compared with the one-stent approach, was associated with significantly higher rate of TLR; and (3) Clinical outcomes after crossover-stenting with one-stent approach for ostial LAD lesions were similar to those after ostial-stenting.

216 Drawbacks of BMS Implantation for Ostial LAD Lesions

217 Ostial LAD lesion has historically been regarded as a lesion subset unsuitable for PCI using

218coronary stents. One of the shortcomings of coronary stenting for ostial LAD lesions was the 219potential for compromising LCX either by plaque shifting or by pinching the LCX ostium. When 220the ostium of LCX had already been significantly narrowed before the procedure, stenting of both 221LAD and LCX might be the only way to optimize the final angiographic result. However, in the era of BMS, stenting both main- and side-branches was considered to be contraindicated in treating 222bifurcation lesions due to unacceptably high restenosis rate ⁹). Also, ostial LAD lesions are often 223224contiguous with the distal LMCA disease, even if the LMCA lesions are not angiographically 225significant. Progression of the LMCA lesions subsequent to the injuries during stent implantation 226 procedure had been another potential concern related to coronary stenting for ostial LAD lesions. 227Furthermore, it is technically demanding to place a stent just at the ostium of LAD without missing 228the adequate coverage of the lesion and without excessive protrusion into the distal LMCA 229bifurcation. Therefore, surgical revascularization could still be considered in patients with ostial 230LAD lesions even if they have single-vessel coronary artery disease.

231 Outcomes of DES Implantation for Ostial LAD Lesions

232 Despite increasingly frequent use of DES for the treatment of ostial LAD lesions, there are only 233 a few small previous studies evaluating the outcome of DES implantation for ostial LAD lesions. 234 Seung et al. compared the clinical outcome of 68 consecutive patients undergoing SES implantation 235 with that of 77 historic control patients undergoing BMS implantation ³⁾. The rate of TLR at 1 year 236 was reported to be less frequent in the SES group than in the BMS group (0% vs. 17%, p < 0.001). 237 Tsagalou et al. compared the clinical outcome of 43 consecutive patients undergoing DES

implantation with that of 43 historic control patients undergoing BMS implantation⁴⁾. The rate of 238239TLR at 9 months was reported to be less frequent in the DES group than in the BMS group (7% vs. 24025.6%, p < 0.001). Our current analysis evaluating larger number of patients clearly demonstrated 241that the rate of TLR at 3 years after SES implantation in patients with ostial LAD lesions was comparable to that in patients with non-ostial proximal LAD lesions, in whom PCI using DES has 242been regarded as the standard of care. The incidences of death or MI were also similar between 243244patients with ostial LAD lesions and patients with non-ostial proximal LAD lesions, suggesting 245safety of PCI using SES for the ostial LAD lesions.

246 Stent Implantation Techniques for Ostial LAD Lesions

247Relatively high restenosis rate in ostial lesions might be related to incomplete lesion coverage 248due to the technical difficulties in stent positioning in this lesion location. Encouraged by the favorable outcomes after unprotected LMCA stenting with DES, crossover-stenting technique 249emerged as a new stenting strategy for the ostial LAD lesions^{3, 5, 10, 11}. In the current analysis, 250251crossover-stenting was adopted in 56% of patients undergoing SES implantation for ostial LAD 252lesions. Cumulative incidences of TLR and death or MI after crossover-stenting were not different 253from those after ostial-stenting, suggesting safety and efficacy of crossover-stenting in selected 254anatomic situations. The Crossover-stenting technique enabling easier stent positioning and full 255coverage of the lesion seemed to be particularly relevant in treating those ostial LAD lesions with 256concomitant distal LMCA disease.

257 In the current analysis, the rate of TLR in patients who underwent stenting of both main- and

side-branches was unacceptably high, as was reported for unprotected LMCA stenting ¹². Although we could not address the safety issues of the two-stent approach due to the small sample size, it would be too premature to promote PCI using DES in patients in whom the two-stent approach is likely to be required.

262 **Study Limitations**

263There are several important limitations in this study. First, we do not have the control group of patients treated by CABG. However, single digit TLR rate at 3 years after PCI seems to be 264265clinically acceptable even if we do not have the surgical control patients. Second, the choices 266 regarding treatment strategies for the ostial LAD lesions were left to discretion of the operators and 267were not based on randomized assignment. Treatment strategies were chosen according to the 268various anatomic features of the ostial LAD lesions. Therefore, the comparison between the 269crossover-stenting and the ostial stenting may not be clinically relevant. Also, we could not address 270the issue of optimal two-stent technique due to small number of patients treated with two-stent 271approach. Third, angiograms were not analyzed by a core angiographic laboratory and therefore, 272the adjudication of ostial lesion was left to the judgment of the local investigators. Fourth, we could 273not address the issue of lesion progression of LMCA and ostial LCX, since we did not evaluate the 274follow-up angiograms. Fifth, because we could not fully monitor the study patients, there is 275potential for under-reporting adverse events with potential for bias. Finally, although this is the 276largest series of patients undergoing SES implantation for the ostial LAD lesions, the study is 277obviously underpowered to evaluate potential small differences in clinical outcomes. Furthermore,

278	small numbers of events severely limit our ability to make adequate statistical adjustment by
279	multivariable analysis. Therefore, the multivariable findings are exploratory due to the small
280	sample size.
281	Conclusions
282	In terms of both safety and efficacy, 3-year outcomes of PCI using SES for ostial LAD
283	lesions were comparable to those for non-ostial proximal LAD lesions. Crossover-stenting across
284	LCX with one-stent approach might be a reasonable option in treating ostial LAD lesions. The
285	two-stent approach for bifurcation was associated with markedly higher rate of TLR than the
286	one-stent approach.
287	
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351 Figure Legends

- 352 Figure 1. Study flow chart for the current analysis among patients enrolled in the j-Cypher registry.
- 353 LAD = left anterior descending coronary artery, and SES = sirolimus-eluting stent.

354

- Figure 2. Cumulative incidences of target lesion revascularization and death or myocardial
 infarction: ostial LAD lesions vs. non-ostial proximal LAD lesions.
- 357 LAD = left anterior descending coronary artery, and SES = sirolimus-eluting stent.

358

- 359 Figure 3. Cumulative incidences of target lesion revascularization and death or myocardial
- 360 infarction among patients treated for ostial left anterior descending coronary artery lesions:
- 361 one-stent vs. two-stent approach.
- 362 SES = sirolimus-eluting stent.

363

- 364 Figure 4. Cumulative incidences of target lesion revascularization and death or myocardial
- 365 infarction among patients treated for ostial left anterior descending coronary artery lesions with
- 366 one-stent approach: crossover-stenting vs. ostial-stenting.
- 367 SES = sirolimus-eluting stent.

369 Tables

370 Table 1. Baseline Characteristics of Patients Treated for Ostial LAD Lesion as Compared With

(A) Patient characteristics					
	Ostial LAD	Non-ostial Proximal LAD	P value		
Number of patients	481	5369			
Age (years)	68.9±10.8	68.1±10.4	0.14		
Age >= 80 years	84 (17%)	700 (13%)	0.006		
Male	365 (76%)	3933 (73%)	0.21		
Body mass index	23.7±3.2	24.0±3.4	0.046		
Body mass index < 25.0	331 (69%)	3461 (64%)	0.06		
Hypertension	341 (71%)	4023 (75%)	0.051		
Diabetes mellitus	193 (40%)	2150 (40%)	0.97		
Diabetes mellitus on insulin therapy	37 (7.7%)	468 (8.7%)	0.44		
Current smoking	91 (19%)	1121 (21%)	0.31		
Statin use	231 (48%)	2278 (42%)	0.02		
eGFR (mL/min/1.73m2)	59.1±21.8	59.7±22.7	0.56		
eGFR < 30, without hemodialysis	23 (4.8%)	248 (4.6%)	0.87		
Hemodialysis	18 (3.7%)	235 (4.4%)	0.51		

371 Non-ostial Proximal LAD Lesion

Acute coronary syndrome	127 (26%)	1479 (28%)	0.59
STEMI	37 (7.7%)	619(12%)	0.01
NSTEMI	11 (2.3%)	124 (2.3%)	0.97
Prior myocardial infarction	142 (30%)	1252 (23%)	0.002
Prior Stroke	45 (9.4%)	498 (9.3%)	0.95
Peripheral vascular disease	56 (12%)	548 (10%)	0.32
Prior heart failure	62 (13%)	746 (14%)	0.54
Multi-vessel disease	240 (50%)	2806 (52%)	0.32
Ejection fraction <= 40%	52 (12%)	521 (11%)	0.49

(B) Lesion and procedural characteristics

Number of lesions	481	5369	
De novo lesion	343 (71%)	4084 (76%)	0.02
In-stent restenosis	76 (16%)	691 (13%)	0.07
Chronic total occlusion	45 (9.4%)	403 (7.5%)	0.14
Severe calcification	53 (11%)	583 (11%)	0.91
Lesion length >= 30mm	92 (19%)	882 (17%)	0.15
Reference vessel diameter pre < 2.5mm	76 (16%)	1504 (28%)	< 0.0001
Use of intravascular ultrasound	351 (73%)	2579 (48%)	< 0.0001
Direct stenting	94 (20%)	1269 (24%)	0.04

Atherectomy before stenting

Directional coronary atherectomy	41(8.5%)	14 (0.3%)	< 0.0001
Rotational atherectomy	32 (6.7%)	313 (5.8%)	0.46
Post dilatation	296 (62%)	2513 (47%)	< 0.0001
Maximum inflation pressure (atm)	18.4±2.8	18.0±3.2	0.008
Number of stents used	1.6±0.8	1.4±0.7	< 0.0001
Length of stents used (mm)	33.2±19.9	30.4±16.0	0.0003
Maximum stent size (mm)	3.2±0.3	3.0±0.3	< 0.0001
Maximum balloon size (mm)	3.4±0.4	3.0±0.4	< 0.0001
Quantitative coronary angiographic data			
Lesion length (mm)	19.4±13.8	19.9±11.7	0.45
Reference vessel diameter pre (mm)	2.99±0.55	2.73±0.49	< 0.0001
Minimal lumen diameter pre (mm)	0.68±0.50	0.63±0.44	0.009
Diameter stenosis pre (%)	77.4±16.2	76.9±15.9	0.56
Reference vessel diameter post (mm)	3.25±0.48	2.94±0.43	< 0.0001
Minimal lumen diameter post (mm)	2.95±0.55	2.68±0.47	< 0.0001
Diameter stenosis post (%)	9.5±9.7	8.8±9.8	0.13

372 Data was missing for body mass index in 2 patients, for body mass index<25.0 in 2 patients, for

373 statin use in 49 patients, for eGFR in 1 patient, for eGFR < 30, without hemodialysis in 1 patient,

374 for ejection fraction <= 40% in 723 patients, de novo lesion in 1 lesion, in-stent restenosis in 1

375	lesion, chronic total occlusion in 14 lesions, lesion length >= 30mm in 68 lesions, reference vessel
376	diameter pre < 2.5mm in 63 lesions, use of intravascular ultrasound in 17 lesions, direct stenting in
377	7 lesions, post dilatation in 9 lesions, maximum inflation pressure in 43 lesions, lesion length in 68
378	lesions, reference vessel diameter pre in 63 lesions, minimal lumen diameter pre in 63 lesions,
379	diameter stenosis pre in 22 lesions, reference vessel diameter post in 53 lesions, minimal lumen
380	diameter post in 53 lesions, and diameter stenosis post in 23 lesions.
381	eGFR = estimated glomerular filtration rate, LAD = left anterior descending coronary artery,

- 382 NSTEMI = non-ST-segment elevation myocardial infarction, and STEMI = ST-segment elevation
- 383 myocardial infarction.

384 Table 2. Unadjusted and Adjusted Outcomes Through 3 Years in Patients Treated for Ostial LAD

	Ostial LAD	Non-ostial Proximal LAD	Multivariable		
	(N=481)	(N=5369)			
	N of events (Incidence)	N of events (Incidence)	p value	HR (95%CI)	p valu
All-cause death	40 (9.7%)	397 (9.2%)	0.51	1.13 (0.8-1.54)	0.48
Cardiac death	18 (4.5%)	205 (4.7%)	0.92	1.05 (0.62-1.66)	0.84
Sudden death	4 (1.2%)	70 (1.6%)	0.37	0.69 (0.21-1.67)	0.45
Myocardial infarction	11 (2.7%)	171 (4.0%)	0.26	0.73 (0.37-1.28)	0.29
Stroke	23 (5.9%)	178 (4.2%)	0.09	1.38 (0.86-2.09)	0.17
Definite/Probable ST	5 (1.2%)	68 (1.6%)	0.65	0.82 (0.29-1.84)	0.66
Definite ST	4 (1.0%)	60 (1.4%)	0.55	0.77 (0.23-1.86)	0.59
TLR	38 (9.4%)	426 (9.7%)	0.98	0.99 (0.7-1.36)	0.94
CABG	5 (1.2%)	66 (1.4%)	0.73	1.03 (0.36-2.35)	0.94
Any coronary revascularization	110 (27.0%)	1372 (29.5%)	0.21	0.91 (0.74-1.1)	0.33
Death/Myocardial infarction	45 (10.7%)	480 (11.4%)	0.82	1.05 (0.76-1.4)	0.77

385 Lesion as Compared With Non-ostial Proximal LAD Lesion

386 Incidence was estimated by Kaplan-Meier method.

388 anterior descending coronary artery, ST=stent thrombosis, and TLR=target-lesion revascularization

³⁸⁷ CABG=coronary artery bypass grafting, CI=confidence interval, HR=hazard ratio, LAD=left

389 Table 3. Unadjusted and Adjusted Outcomes Through 3 Years in Patients with Ostial LAD Lesions

390 Treated by One-stent Approach as Compared With Those Treated by the Two-stent Approach.

	One-stent approach	Two-stent approach		Multivariable	
	(N=419)	(N=62)			
	N of events (Incidence)	N of events (Incidence)	p value	HR (95%CI)	p value
All-cause death	32 (8.6%)	8 (16.8%)	0.2	1.3 (0.54-2.83)	0.54
Cardiac death	14 (3.6%)	4 (10.5%)	0.24	0.92 (0.25-2.79)	0.89
Sudden death	3 (0.9%)	1 (4.0%)	0.47		
Myocardial infarction	10 (2.8%)	1 (2.1%)	0.69	0.66 (0.04-3.46)	0.68
Stroke	16 (4.7%)	7 (1.4%)	0.01	3.38 (1.3-7.93)	0.01
Definite/Probable ST	4 (1.1%)	1 (2.1%)	0.64		
Definite ST	3 (0.8%)	1 (2.1%)	0.48		
TLR	22 (6.6%)	16 (28.1%)	<0.0001	4.65 (2.32-9.25)	< 0.0001
CABG	1 (0.3%)	4 (7.4%)	<0.0001		
Any coronary revascularization	85 (24.3%)	25 (44.7%)	<0.0001	2.11 (1.3-3.33)	0.003
Death/Myocardial infarction	37 (9.8%)	8 (16.8%)	0.37	1.15 (0.49-2.41)	0.73

391 Incidence was estimated by Kaplan-Meier method.

392 Abbreviations are same as in Table 2.

394 Table 4. Unadjusted and Adjusted Outcomes Through 3 Years in Patients with One-stent Approach

- 395 Treated by Ostial-stenting Technique as Compared With Those Treated by Crossover-stenting
- 396 Technique

	Ostial-Stenting	Crossover-Stenting		Multivariable	
	(N=194)	(N=225)			
	N of events (Incidence)	N of events (Incidence)	p value	HR (95%CI)	p value
All-cause death	8 (4.5%)	24 (12.2%)	0.01	2.04 (0.94-4.93)	0.07
Cardiac death	3 (1.7%)	11 (5.2%)	0.06	1.7 (0.49-7.85)	0.42
Sudden death	0 (0%)	3 (1.6%)	0.1		
Myocardial infarction	6 (3.6%)	4 (2.0%)	0.41	0.59 (0.15-2.07)	0.41
Stroke	8 (4.8%)	8 (4.6%)	0.84	0.87 (0.32-2.38)	0.79
Definite/Probable ST	1 (0.5%)	3 (1.6%)	0.37		
Definite ST	1 (0.5%)	2 (1.1%)	0.62		
TLR	11 (7.9%)	11 (5.4%)	0.81	0.77 (0.33-1.82)	0.55
CABG	0 (0%)	1 (0.5%)	0.35		
Any coronary revascularization	41 (25.1%)	44 (23.5%)	0.8	0.93 (0.61-1.42)	0.73
Death/Myocardial infarction	12 (7.0%)	25 (12.2%)	0.07	1.54 (0.78-3.2)	0.22

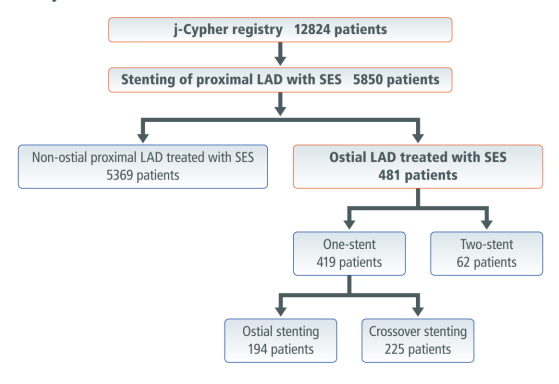
397 Incidence was estimated by Kaplan-Meier method.

398 Abbreviations are same as in Table 2.

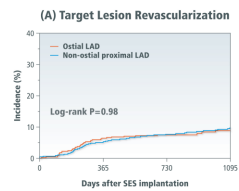
399 Figures

400 Figure 1.

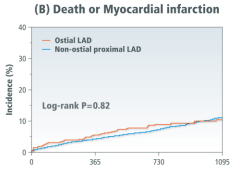
Study Flow Chart



401



Ostial LAD versus Non-ostial Proximal LAD

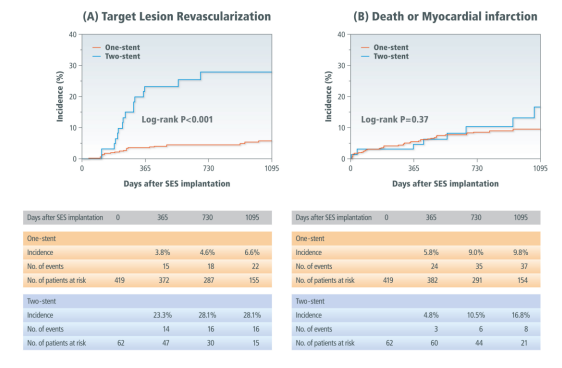


Days after SES implantation

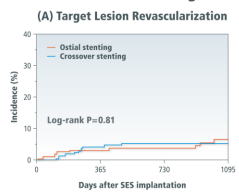
Days after SES implantation	0	365	730	1095					
Ostial LAD									
Incidence		6.3%	7.6%	9.4%					
No. of events		29	34	38					
No. of patients at risk	481	417	316	170					
Non-ostial proximal LAD									
Incidence		5.2%	7.7%	9.7%					
No. of events		268	378	426					
No. of patients at risk	5369	4727	3550	1562					

Days after SES implantation	0	365	730	1095					
Ostial LAD									
Incidence		5.7%	9.2%	10.7%					
No. of events		27	41	45					
No. of patients at risk	481	441	334	175					
Non-ostial proximal LAD									
Incidence		4.5%	7.5%	11.4%					
No. of events		237	374	480					
No. of patients at risk	5369	4942	3797	1661					

404



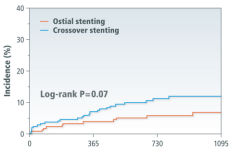
One-stent versus Two-stent



Ostial Stenting versus Crossover stenting

David offer CEC involution

(B) Death or Myocardial infarction



Days after SES implantation

0	365	730	1095
	3.2%	3.8%	7.9%
	6	7	11
194	175	132	79
	4.3%	5.4%	5.4%
	9	11	11
225	197	147	76
	194	3.2% 6 194 175 4.3% 9	3.2% 3.8% 6 7 194 175 132 4.3% 5.4% 9 11

Days after SES Implantation	0	202	/50	1095					
Ostial stenting									
Incidence		4.2%	6.1%	7.0%					
No. of events		8	11	12					
No. of patients at risk	194	178	133	78					
Crossover stenting									
Incidence		7.2%	11.5%	12.2%					
No. of events		16	24	25					
No. of patients at risk	225	204	150	76					