Left Main Disease



Comparisons of PCI against CABG 10 years of advances

	Design	N (PCI/ CABG)	Endpoint	FU, yrs	Key findings
MAIN COMPARE (2008, 2010, 2018)	Multicenter registry	1102/1138	Death; death, Q-wave MI, or stroke; TVR	10	Similar rates of mortality and death, Q-wave MI, or stroke; higher rates of TVR with PCI
LE MANS (2008, 2016)	Multicenter RCT	52/53	Change in LVEF	10	Improvement in ejection fraction only with PCI, comparable rates of death, MI, stroke, or TVR
SYNTAX (2010, 2014)	Multicenter RCT	357/348	Death, MI, stroke, or RR	5	Comparable rates of death, MI, stroke, or repeat revascularization
Boudriot et al. (2011)	Multicenter RCT	100/101	Death, MI, or RR	1	PCI with sirolimus-eluting stent inferior to CABG
PRECOMBAT (2011, 2015, 2020)	Multicenter RCT	300/300	Death, MI, stroke, or ischemia-driven TVR	10	PCI non-inferior to CABG at 1, 5, and 10 year, comparable rates of death, MI, stroke, or ischemia-driven TVR
DELTA (2012)	Multicenter registry	1874/901	Death, MI, or stroke	3.5	Comparable rates of death, MI, or stroke. Higher TVR in PCI
NOBLE (2016)	Multicenter RCT	592/592	Death, MI, stroke, or any repeat revascularization	5	CABG superior to PCI (primary end points 28% in PCI group vs in 18% in CABG group)
EXCEL (2016, 2018)	Multicenter RCT	948/957	Death, MI, or stroke	4	Similar rates of primary endpoint of death, stroke, or MI at 4 years

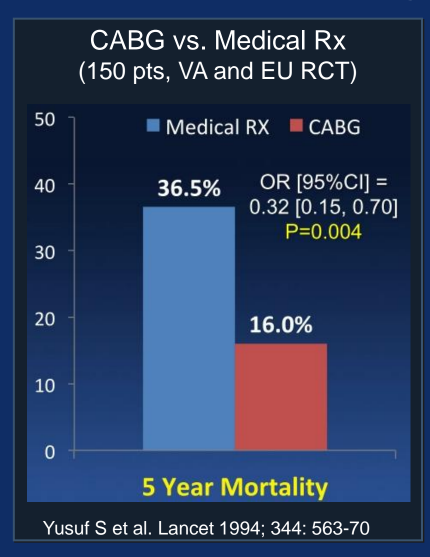


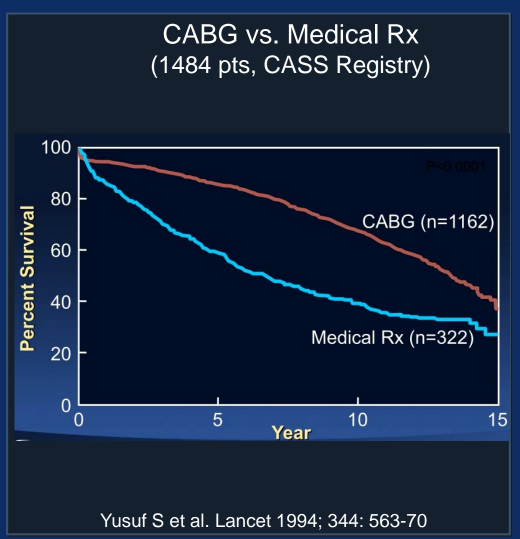


TABLE 1 Summary of Randomized Clinical Trials of PCI With DES Vs CABG for LMCA Disease

	L FARANC30.31	Devidules et =132	CVAITA V 1 1433-35	DDECOMD 4 736 38	EV.CEI 39.40	NOBLE ^{41,42}
	LEMANS ^{30,31}	Boudriot et al ³²	SYNTAX-LM ³³⁻³⁵	PRECOMBAT ^{36,38}	EXCEL ^{39,40}	NORLE
Recruitment period	2001-2004	2003-2009	2005-2007	2004-2009	2010-2014	2008-2015
PCI/CABG, n/n	52/53	100/101	357/348	300/300	948/957	592/592
Follow-up, y	10	1	5 10 (for mortality)	10	5	5
Diabetes, %	18	36	25	32	29	15
Bifurcation, %	58	72	61	64	81	81
SYNTAX score, mean	Not reported	23	30	25	21	22
Stent	BMS and DES (35%)	DP-SES	DP-PES	DP-SES	DP-EES	BP-BES and DP-SES (7.7%)
IVUS	Recommend	Infrequent	Infrequent	At discretion, 91%	Recommended, 77%	Recommended, 74%
FFR guidance	Not reported	Not reported	Infrequent	Not reported	Recommended, 9.0%	Recommended
LIMA, %	72	99	97	94	99	96
Off pump, %	1.9	46	Not reported	64	29	16
Primary trial endpoint	Change in LVEF	Cardiac death, MI, or TVR	Death, MI, stroke, or repeat revascularization 10-y all-cause death	Death, MI, stroke, or TVR	Death, MI, or stroke	Death, nonprocedural MI, stroke, or repeat revascularization
Key finding	There was a trend toward higher LVEF at 10 y with PCI.	PCI was inferior to CABG at 1 y.	PCI was noninferior to CABG at 1 and 5 y in terms of death, MI, stroke, or repeat revascularization. No significant difference in 10-y all- cause death between PCI and CABG.	PCI was noninferior to CABG at 1, 5, and 10 y.		PCI was inferior to CABG at 5 y.

Data for Left Main 30 years ago







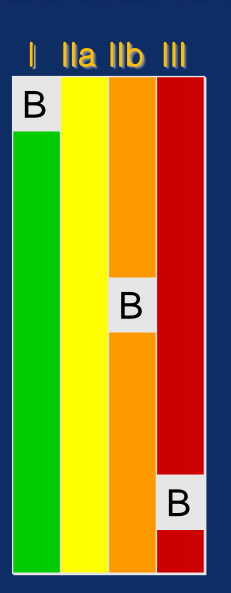


Guideline Changes for LMCA, 10 Years

	Class of recommendation	LOE		
2005 ACC/AHA/SCAI	III—PCI is not recommended in patients with unprotected LMCA disease and eligibility for CABG	С		
2005 ESC/EACTS	IIb —Stenting for unprotected LMCA disease should only be considered in the absence of other revascularization options	С		
:				
	 IIa—For SIHD patients when both of the following are present: Anatomically low risk of PCI procedural complications & high likelihood of good long-term outcomes (e.g., a low SYNTAX score [≤22], ostial or trunk left main stenosis) Clinical characteristics that predict a significantly increased risk of adverse surgical outcomes (e.g., STS-predicted risk of operative mortality ≥5%) 	В		
2011/2014 ACC/AHA/AATS/PCNA /SCAI/STS	 IIb—For SIHD patients when both of the following are present: Anatomically low-to-intermediate risk of PCI procedural complications & intermediate-to-high likelihood of good long-term outcome (e.g., low-intermediate SYNTAX score of <33, bifurcation left main stenosis) Clinically increased risk of adverse surgical outcomes 			
	III: HARM—SIHD patients with unfavorable anatomy for PCI & good candidates for CABG	В		
2014 ESC/EACTS	I—Left main disease with a SYNTAX score ≤ 22. IIb—Left main disease with a SYNTAX score 23–32 III—Left main disease with a SYNTAX score ≥ 33	В		
2018 ESC/EACTS	I—Left main disease with a SYNTAX score ≤ 22. IIa—Left main disease with a SYNTAX score 23–32 III—Left main disease with a SYNTAX score ≥ 33	A B		
	I—In patients with SIHD and significant left main stenosis, CABG is recommended to improve survival.			
2021 ACC/AHA	IIa —In selected patients with SIHD and significant left main stenosis for whom PCI can provide equivalent revascularization to that possible with CABG, PCI is reasonable to improve survival	В		
2024 ESC/EACTS	I—In CCS patients with significant left main coronary stenosis of low complexity (SYNTAX score ≤22), in whom PCI can provide equivalent completeness of revascularization to that of CABG, PCI is recommended as an alternative to CABG, given its lower invasiveness and non-inferior survival. IIa—In CCS patients with significant left main coronary stenosis of intermediate complexity (SYNTAX score 23–32), in whom PCI can provide equivalent completeness of revascularization to that of CABG, PCI should be considered, given its lower invasiveness and non-inferior survival.	A		



Elective PCI for LM Stenosis ESC/EACTS Guidelines 2014



- LM with
 - SYNTAX score ≤ 22

- LM with
 - SYNTAX score 23-32

- LM with
 - SYNTAX score > 32



Elective PCI for LM Stenosis ESC/EACTS Guidelines 2018



- LM with
 - SYNTAX score ≤ 22

- LM with
 - SYNTAX score 23-32

- LM with
 - SYNTAX score > 32



Elective PCI for LM Stenosis ACC/AHA Guidelines 2021



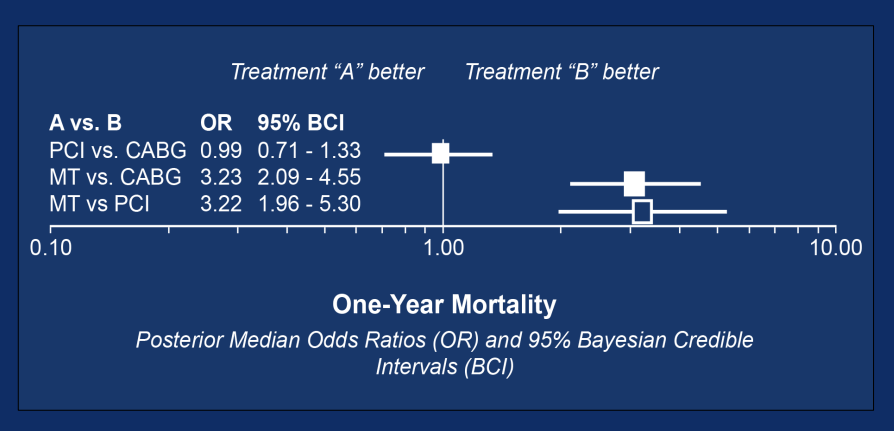
- PCI and provide equivalent revascularization to that possible with CABG
 - PCI is reasonable to improve survival

LM: PCI vs. CABG



PCI vs. Medical Treatment

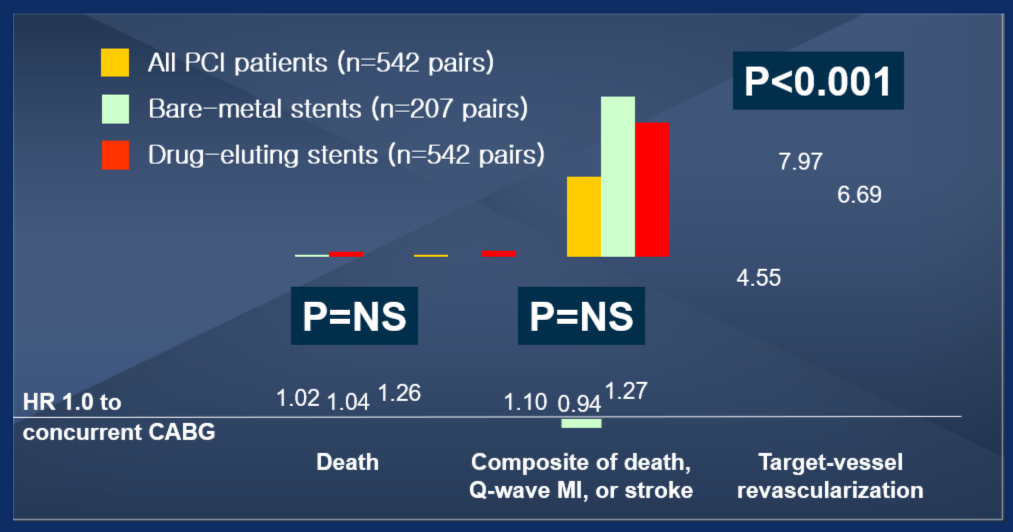
Bayesian network meta-analysis involving 12 (PCI vs. CABG), and 7 (CABG vs. Medication) studies



PCI is superior to medical treatment in the treatment of LM stenosis.



Hazard Ratios for Matched Cohort Outcomes : Median 5-Year Outcomes

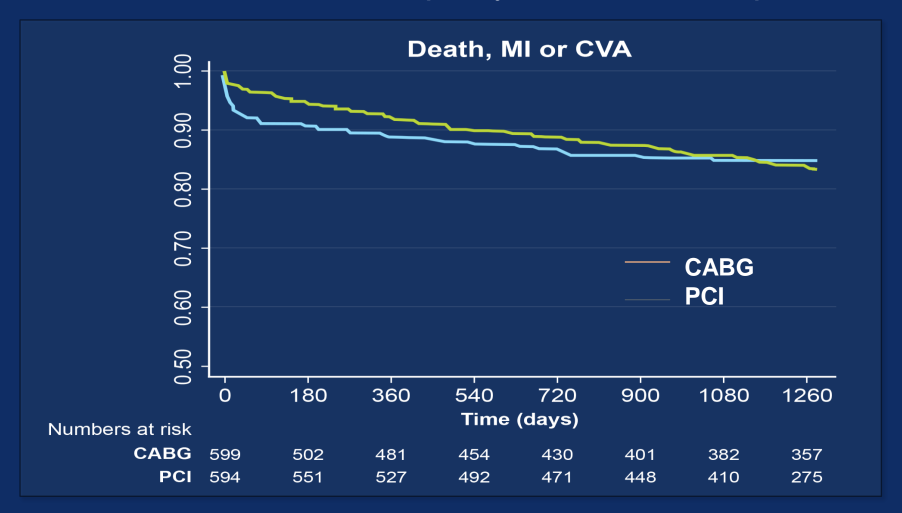




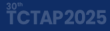


The DELTA Registry LM revascularization: PCI vs. CABG

Death, MI or CVA in Propensity Score-Matched Groups



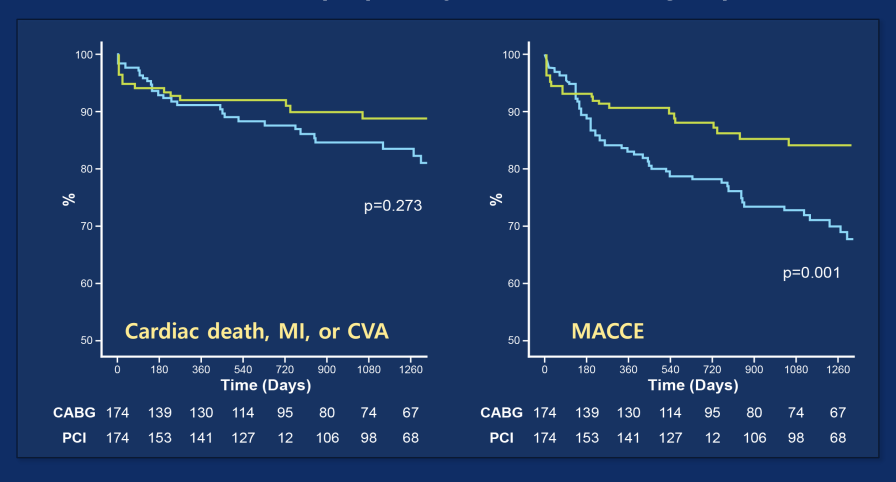




PCI vs. CABG in Females

Female subgroup of <u>DELTA</u> registry (PCI, 489; CABG, 328 patients)

<u>The results of propensity score-matched groups</u>

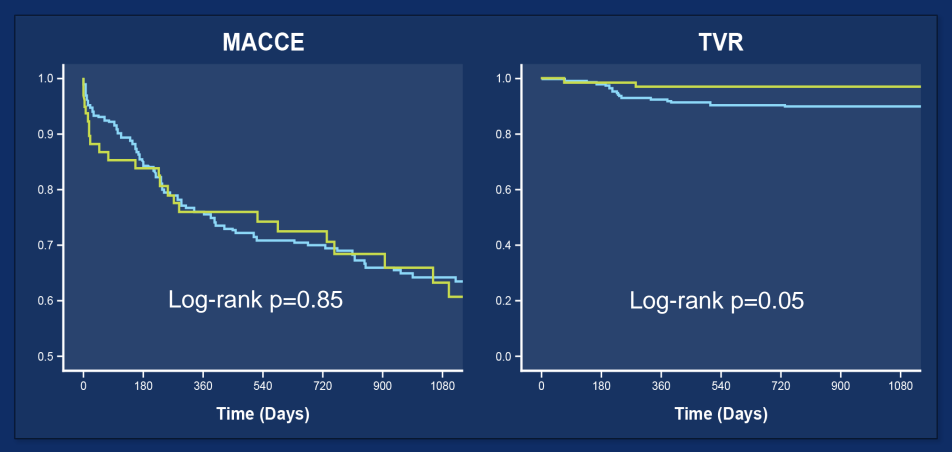


There was no difference in the hard endpoints.



PCI vs. CABG in Octogenarians

Octogenarian subgroup of *DELTA* registry (PCI, 218; CABG, 86)



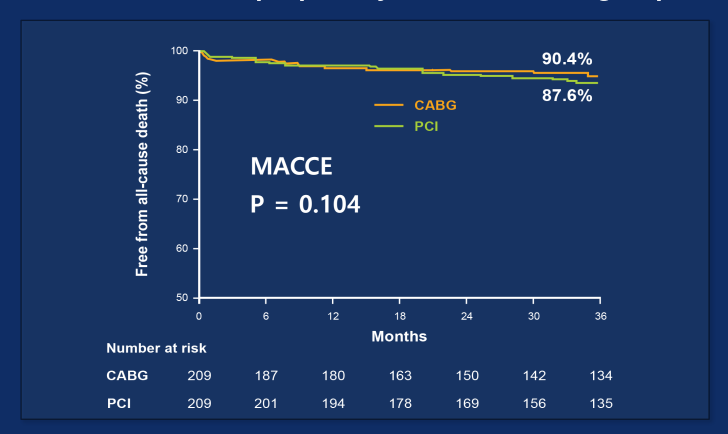
In octogenarians, no difference was observed in the occurrence of the hard endpoint after PCI or CABG.



PCI vs. CABG for Ostial/Midshaft LM stenosis

A subgroup of <u>DELTA</u> registry (PCI, 482; CABG, 374 patients)

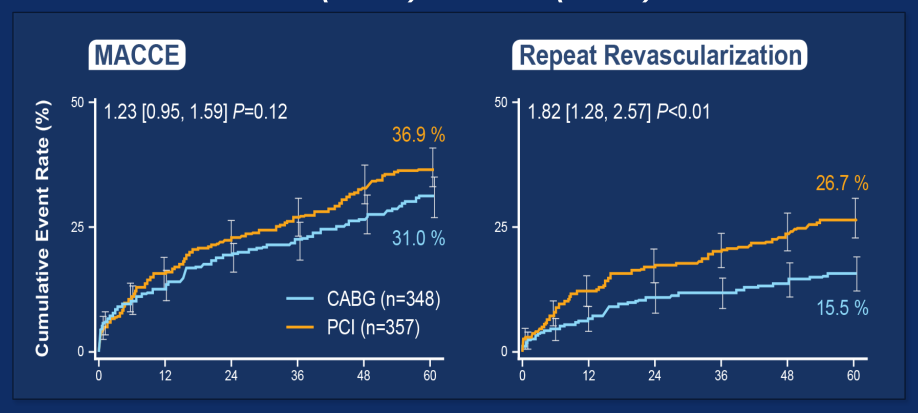
<u>The results of propensity score-matched groups</u>



PCI for ostial/midshaft lesions was associated with clinical outcomes comparable to those observed with CABG



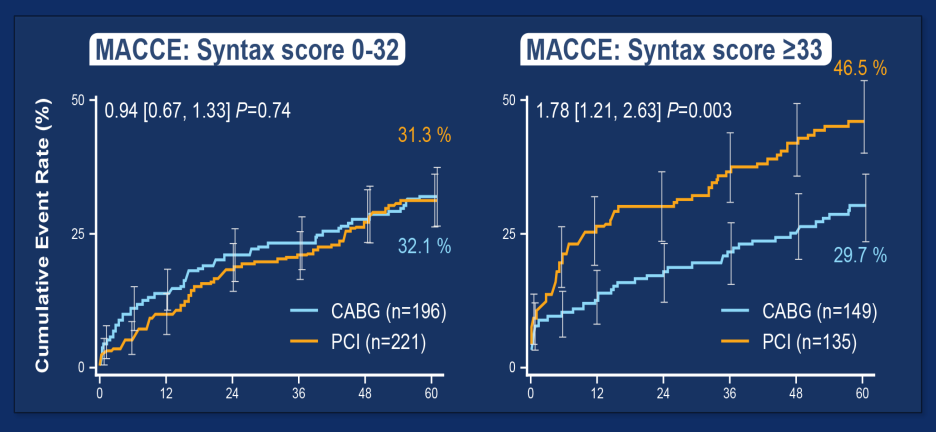
5-year outcomes of the LM subgroup of the <u>SYNTAX</u> trial :PCI (N=357) vs. CABG (N=348)



At 5 years, no difference in MACCE was found between PCI and CABG, but PCI was accompanied by a higher rate of repeat revascularization.



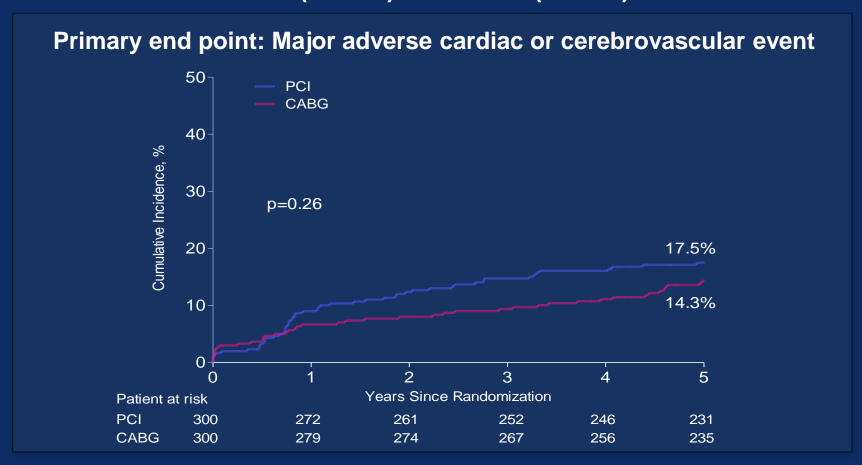
5-year outcomes of the LM subgroup of the <u>SYNTAX</u> trial :PCI (N=357) vs. CABG (N=348)



MACCE were similar between arms in patients with low/intermediate SYNTAX scores but significantly increased in patients with high scores.



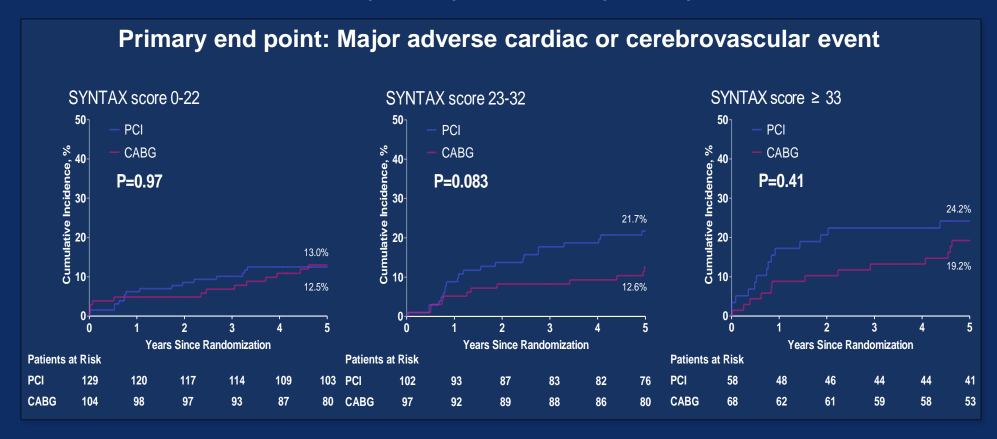
5-year outcomes of the randomized PRECOMBAT trial :PCI (N=300) vs. CABG (N=300)



During 5 year follow-up, no significant difference in the rate of MACCE was observed between the PCI and CABG groups.



5-year outcomes of the randomized **PRECOMBAT** trial :PCI (N=300) vs. CABG (N=300)

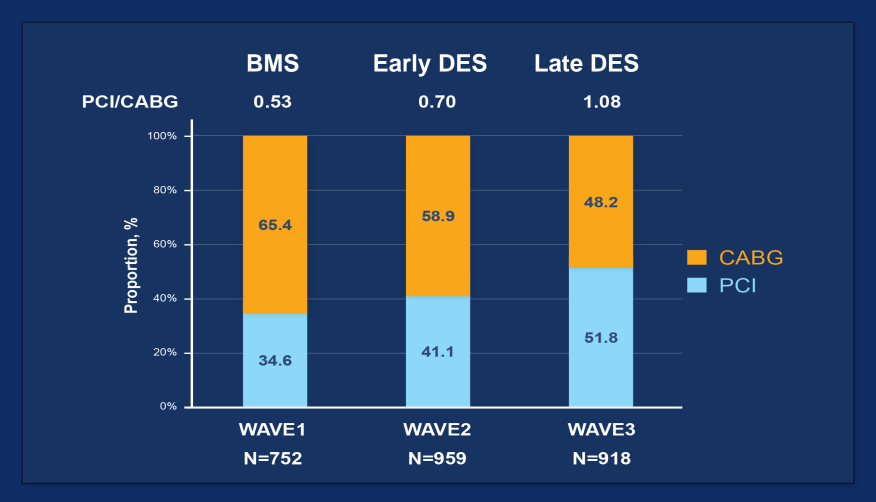


During 5 year follow-up, no significant difference in the rate of MACCE was observed between the PCI and CABG groups.



Temporal Trends

Data From the **Asan Medical Center**-LM Revascularization Registry

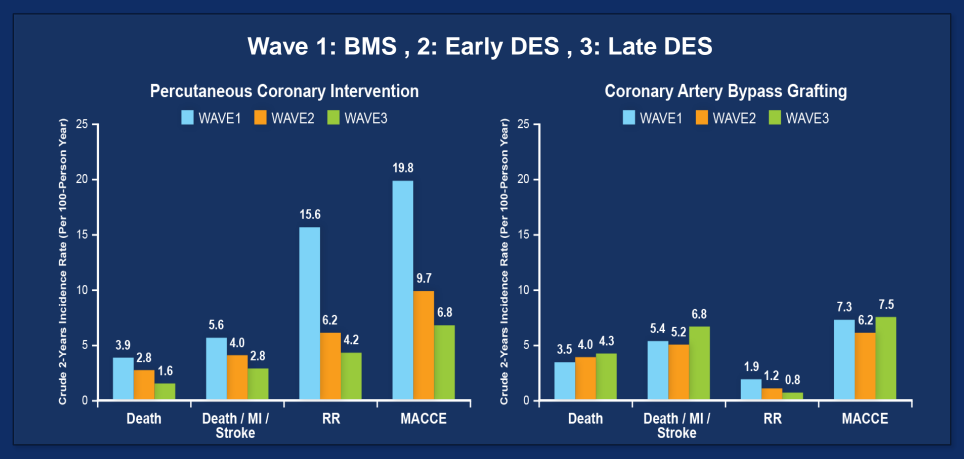


The proportion of PCI is significantly increasing.



Temporal Trends

Data From the Asan Medical Center-LM Revascularization Registry

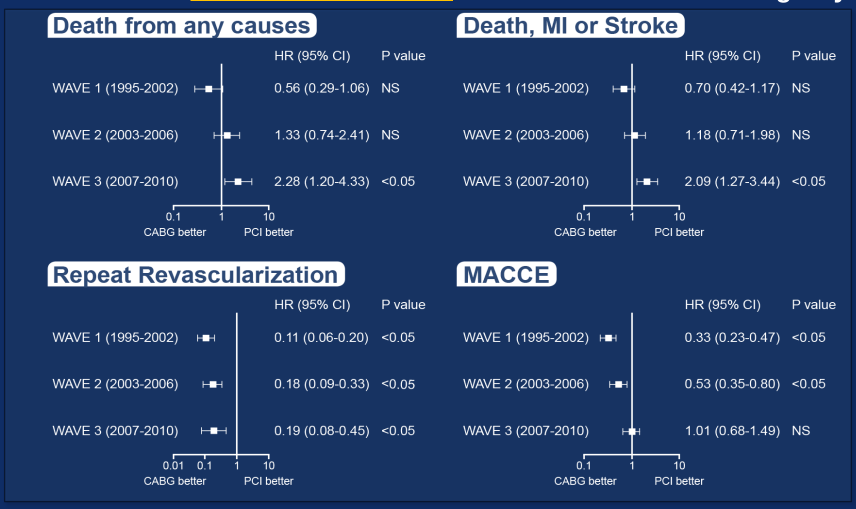


The incidence of adverse events is gradually decreasing with PCI, but the change has been insignificant with CABG.



Temporal Trends

Data From the **Asan Medical Center-LM** Revascularization Registry

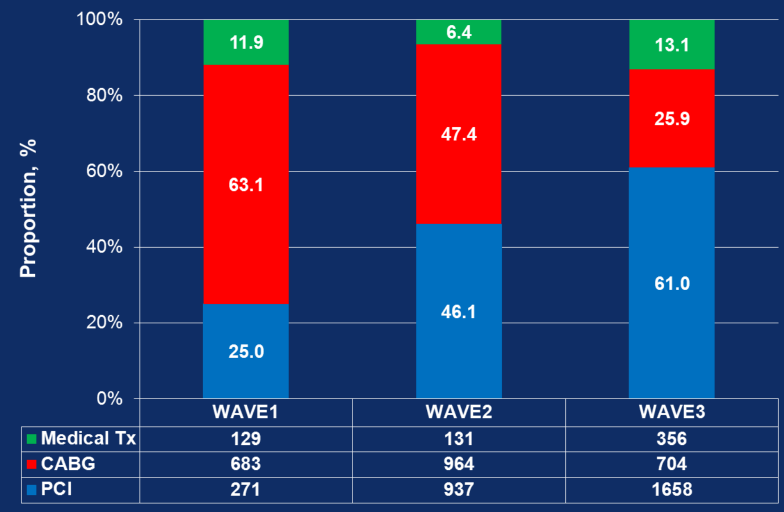


The trend favoring PCI was observed with the coronary stent evolving.



IRIS-MAIN registry

50 academic and community hospitals in Asia (*n*=5883)

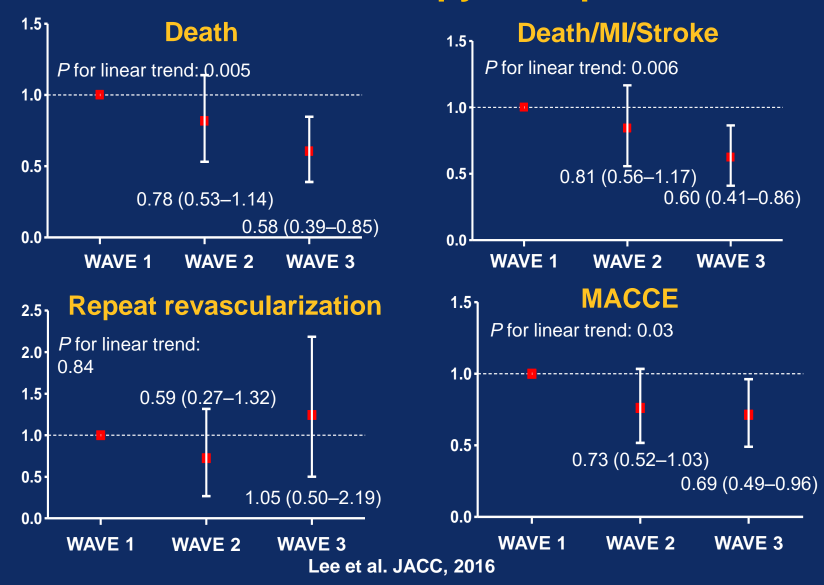


Historical time periods: WAVE1: 1995 – 2002, WAVE2: 2003 – 2006, WAVE3: 2007 – 2013



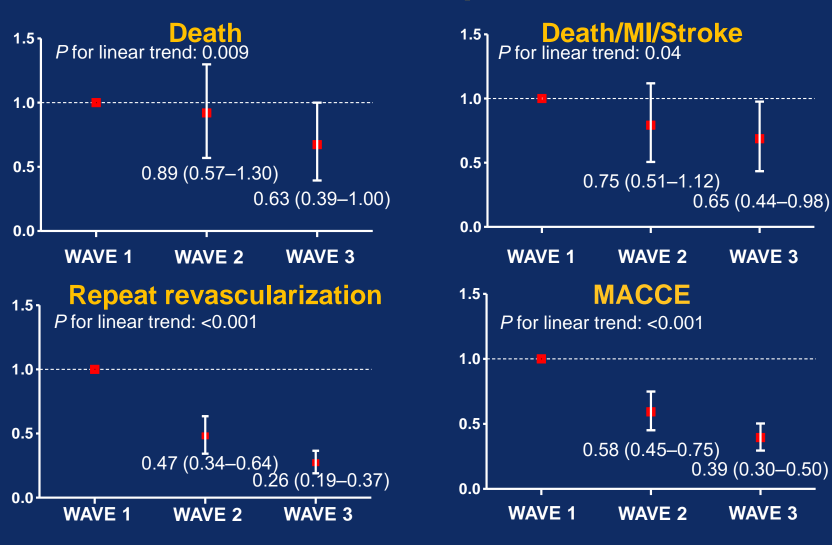
IRIS-MAIN registry

Medical Therapy Group



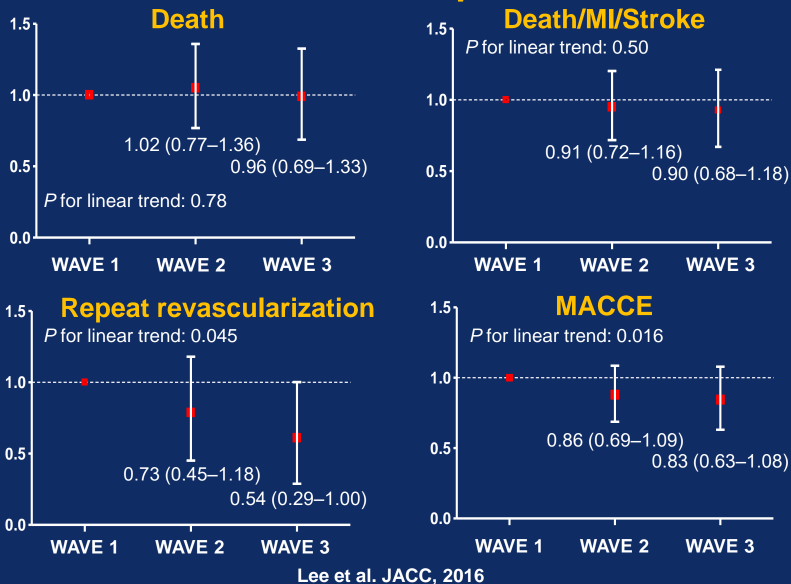
IRIS-MAIN registry

PCI Group

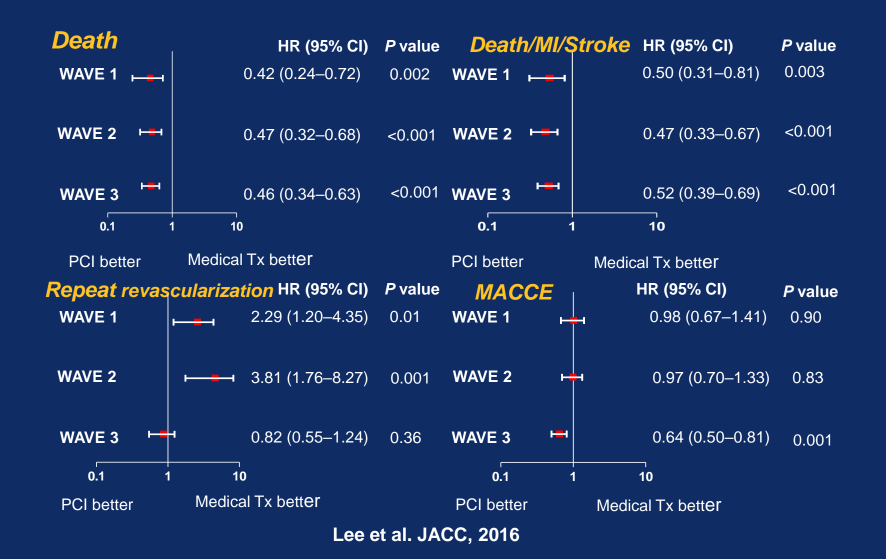


IRIS-MAIN registry

CABG Group

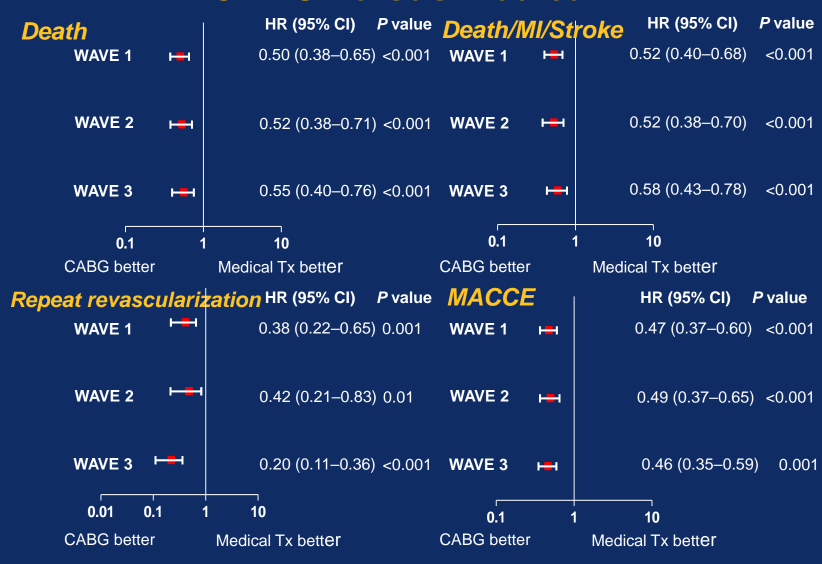


IRIS-MAIN registry PCI versus Medical Tx



IRIS-MAIN registry

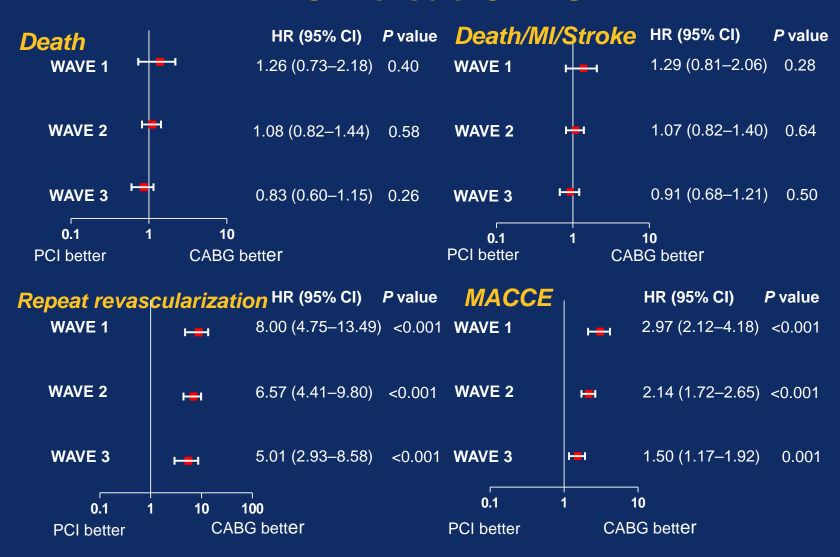
CABG versus Medical Tx





IRIS-MAIN registry

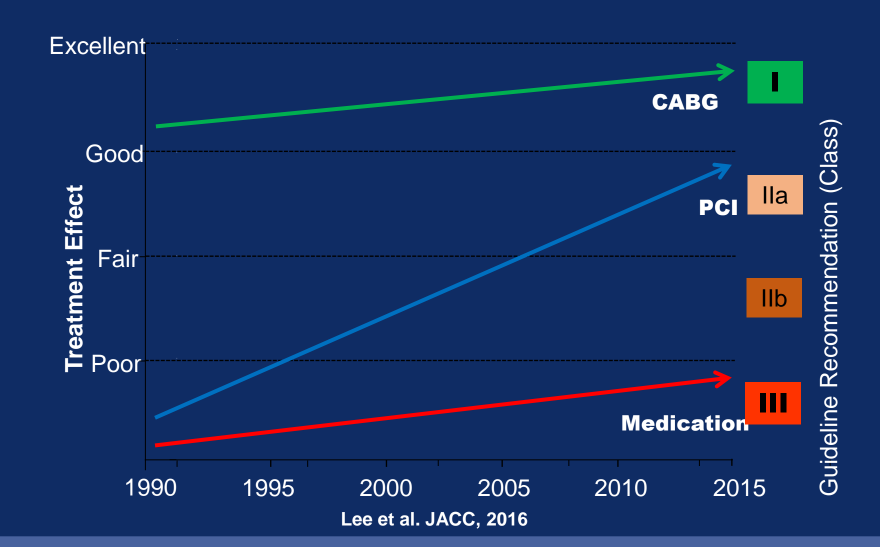
PCI versus CABG





IRIS-MAIN registry

Secular Changes of Treatment Effectof Each Treatment Stratum





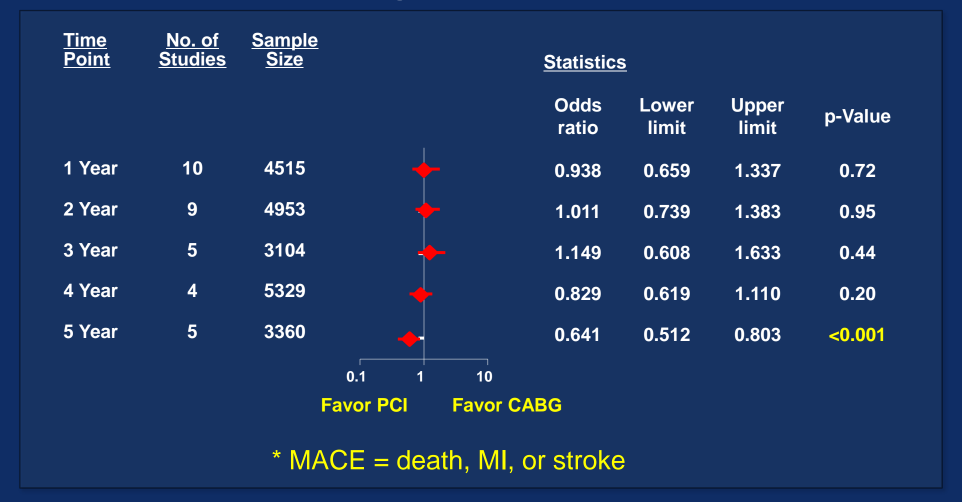
PCI vs. CABG for Left Main Disease Meta-analysis of 4 RCTs, 1,611 Patients

1 Year MACCE						
	PCI	CABG	OR (95%CI)	<i>p</i> -Value	OR (95%CI)	
LEMANS	16/52	13/53	1.37 (0.58-3.23)	0.48		
SYNTAX left main	56/355	46/336	1.18 (0.77-1.80)	0.44		
Boudriot et al.	19/100	14/101	1.46 (0.69-3.10)	0.33		
PRECOMBAT	26/300	20/300	1.33 (0.73-2.44)	0.36		
Fixed effects estiamate	14.5% (117/807)		1.28 (0.95-1.72)	0.11		
Random effects estimat	te		1.28 (0.95-1.72)		0.01 0.1 1 10 100	



PCI vs. CABG for Left Main Disease Meta-analysis of 24 studies, 14,203 patients

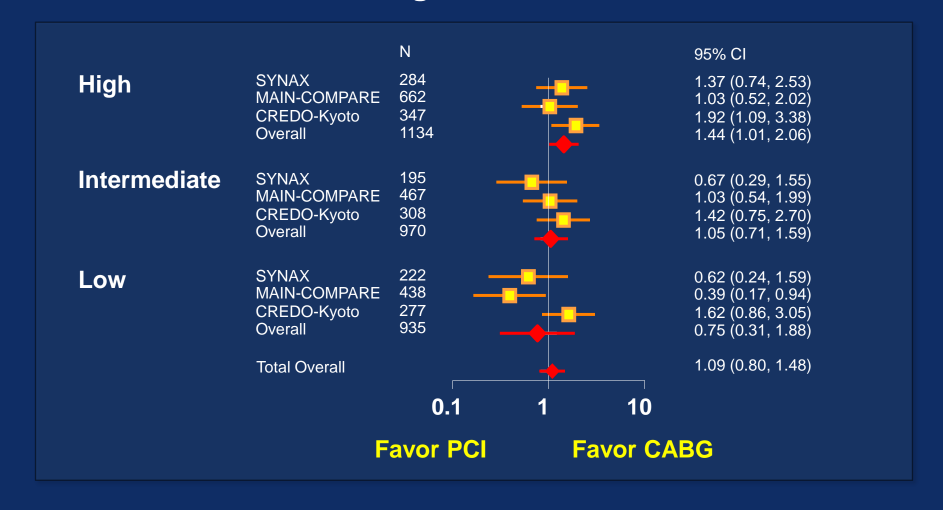
According to follow-up duration







PCI vs. CABG for Left Main Disease Meta-analysis of 24 studies, 14,203 patients According to SYNTAX score

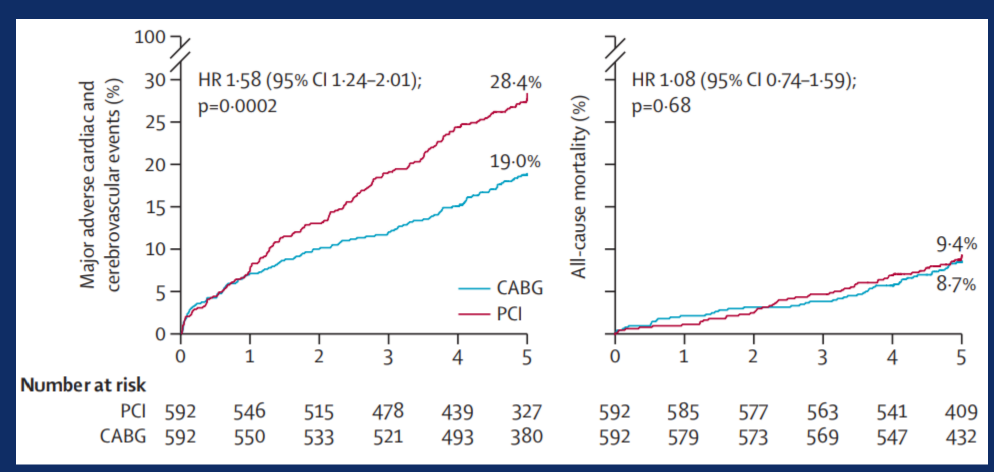




PCI vs. CABG for Left Main Disease

5-year clinical outcomes of the randomized NOBLE trial :PCI (N=592) vs. CABG (N=592)

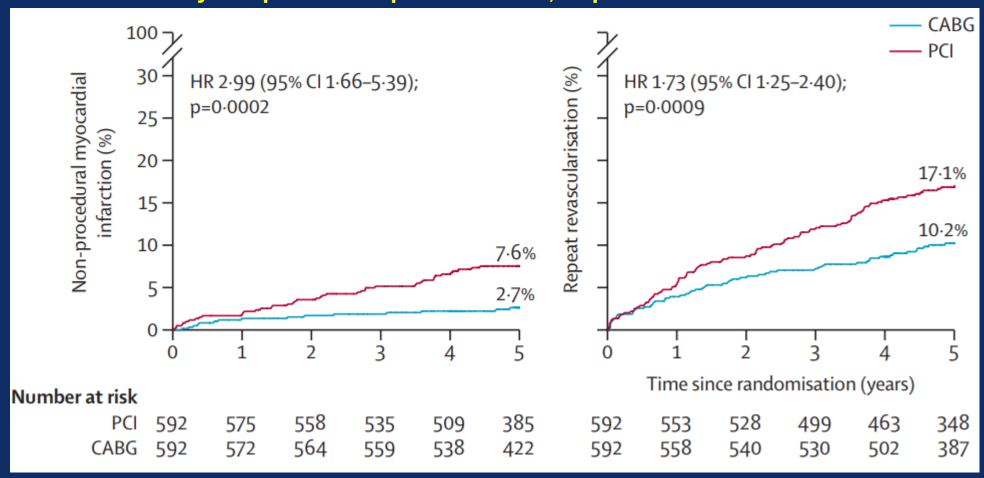
Primary Endpoint: MACCE, All-cause mortality



PCI vs. CABG for Left Main Disease

5-year clinical outcomes of the randomized NOBLE trial :PCI (N=592) vs. CABG (N=592)

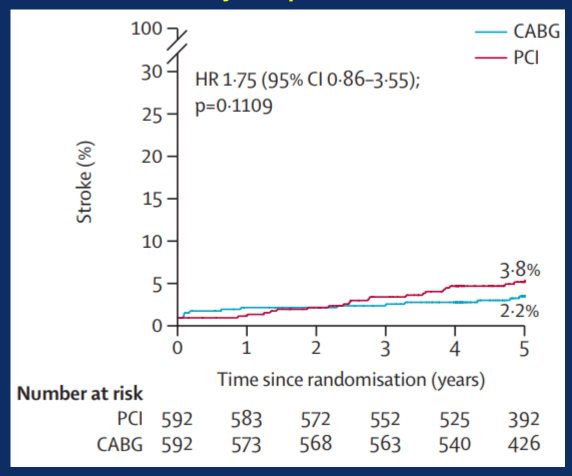
Primary Endpoint: Non-procedural MI, Repeat revascularization



PCI vs. CABG for Left Main Disease

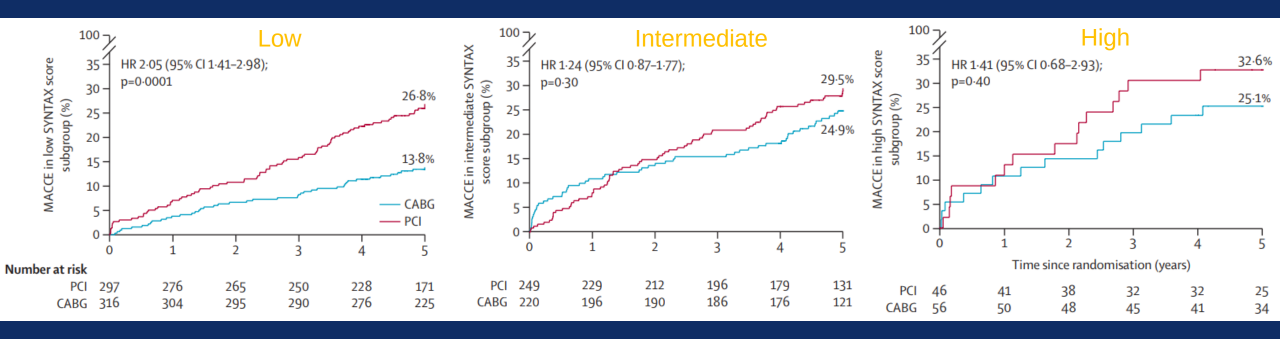
5-year clinical outcomes of the randomized NOBLE trial :PCI (N=592) vs. CABG (N=592)

Primary Endpoint: Stroke



5-year clinical outcomes of the randomized NOBLE trial :PCI (N=592) vs. CABG (N=592)

Primary Endpoint(MACCE) by SYNTAX score subgroups



A low score is defined as 1–22; intermediate is 23–32; high is ≥33.



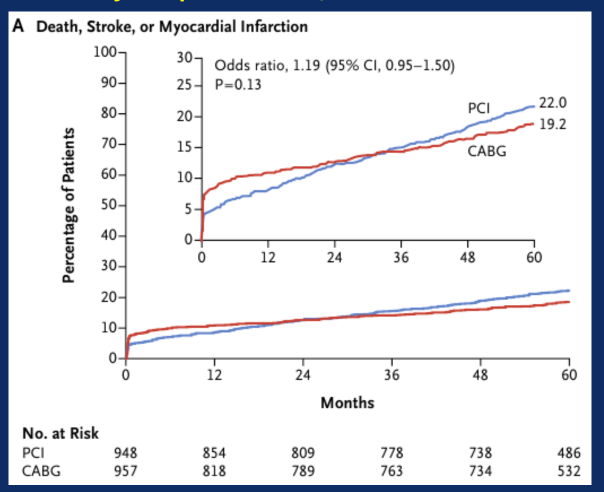
5-year outcomes of the randomized **EXCEL** trial :PCI (N=948) vs. CABG (N=957)

	PCI (n=948)	CABG (n=957)	Diff [95% CI]	OR [95%CI]
Primary endpoint				
Death, stroke or MI at 5 years	22.0%	19.2%	2.8 [-0.9 to 6.0]	1.19 (0.95-1.05)
Secondary endpoints				
Death from any cause	13.0%	9.9%	3.1 [0.2 to 6.1]	1.38 (1.03-1.85)
Death, stroke, MI or ischemia-driven revasc	31.3 %	24.9 %	6.2 [2.4-10.6]	1.39 (1.13-1.71)



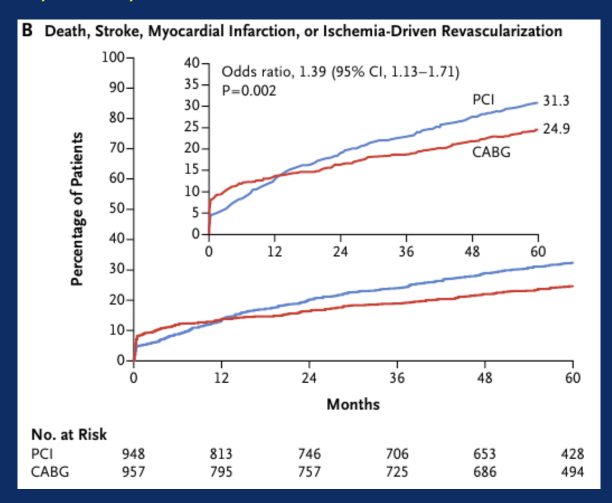
5-year outcomes of the randomized **EXCEL** trial :PCI (N=948) vs. CABG (N=957)

Primary Endpoint Death, Stroke or MI at 5 Years



5-year outcomes of the randomized **EXCEL** trial :PCI (N=948) vs. CABG (N=957)

Death, Stroke, MI or Ischemia-driven Revascularization at 5 Years





5-year outcomes of the randomized **EXCEL** trial :PCI (N=948) vs. CABG (N=957)

Secondary Outcomes Analysis

Outcome	PCI (N = 948)		CABG (N = 957)		Difference in Event Rates (95% CI)	Odds Ratio (95% CI)	
	Events	Events Event Rate		Event Rate			
	no.	%	no.	%	percentage points		
Death from any cause	119	13.0	89	9.9	3.1 (0.2 to 6.1)	1.38 (1.03 to 1.85)	
Cardiovascular	61	6.8	49	5.5	1.3 (-0.9 to 3.6)	1.26 (0.85 to 1.85)	
Definite cardiovascular	45	5.0	40	4.5	0.5 (-1.4 to 2.5)	1.13 (0.73 to 1.74)	
Undetermined cause	16	1.9	9	1.1	0.9 (-0.3 to 2.0)	1.78 (0.78 to 4.06)	
Noncardiovascular	58	6.6	40	4.6	2.0 (-0.2 to 4.2)	1.47 (0.97 to 2.23)	
Stroke	26	2.9	33	3.7	-0.8 (-2.4 to 0.9)	0.78 (0.46 to 1.31)	
Myocardial infarction	95	10.6	84	9.1	1.4 (-1.3 to 4.2)	1.14 (0.84 to 1.55)	
Periprocedural	37	3.9	57	6.1	-2.1 (-4.1 to -0.1)	0.63 (0.41 to 0.96)	
Nonperiprocedural	59	6.8	31	3.5	3.2 (1.2 to 5.3)	1.96 (1.25 to 3.06)	
Ischemia-driven revascularization	150	16.9	88	10.0	6.9 (3.7 to 10.0)	1.84 (1.39 to 2.44)	
PCI	125	14.1	80	9.1	4.9 (1.9 to 7.9)	1.65 (1.22 to 2.22)	
CABG	38	4.3	8	0.9	3.4 (1.9 to 4.9)	4.90 (2.27 to 10.56)	

5-year outcomes of the randomized **EXCEL** trial :PCI (N=948) vs. CABG (N=957)

Additional Outcomes Analysis

Outcome	PCI (N = 948)		CABG (N = 957)		Difference in Event Rates (95% CI)	Odds Ratio (95% CI)	
	Events Event Rate		Events	Event Rate			
	no.	%	no.	%	percentage points		
Additional outcomes							
Any revascularization	153	17.2	92	10.5	6.7 (3.5 to 9.9)	1.79 (1.36 to 2.36)	
Stent thrombosis	16	1.8	0	0	_	_	
Definite	10	1.1	0	0	_	_	
Probable	6	0.7	0	0	_	_	
Symptomatic graft stenosis or occlusion	0	0	58	6.5	_	_	
Therapy failure†	10	1.1	58	6.5	-5.4 (-7.2 to -3.6)	0.16 (0.08 to 0.32)	
Cerebrovascular events‡	29	3.3	46	5.2	-1.9 (-3.8 to 0)	0.61 (0.38 to 0.99)	
Transient ischemic attack	3	0.3	14	1.6	-1.3 (-2.2 to -0.4)	0.21 (0.06 to 0.74)	



5-year outcomes of the randomized **EXCEL** trial: PCI (N=948) vs. CABG (N=957)

Subgroup analysis of Primary outcomes at 5 Years

Subgroup	PCI (N=948)		CABG (N=	957)	Odds Ratio (95%	CI)
	Events/total patients	Event rate	Events/total patients	Event rate		
	no.	%	no.	%	1	
All patients	203/948	22.0	176/957	19.2		1.19 (0.95–1.5
Age (median cutoff)						
≥67 yr	123/466	27.2	98/472	21.8	—	1.39 (1.02–1.8
<67 yr	80/482	16.9	78/485	16.6		1.00 (0.71–1.4
Sex					i	
Male	145/722	20.6	134/742	18.7		1.12 (0.86–1.4
Female	58/226	26.3	42/215	21.1		1.39 (0.88-2.2
Diabetes mellitus, medically treated						
Yes	72/256	29.0	62/249	25.5		1.24 (0.83-1.8
No	131/692	19.4	114/707	16.9	+-	1.17 (0.89–1.5
Chronic kidney disease						
Estimated GFR ≤60 ml/min	54/164	34.0	37/144	27.6		1.44 (0.86-2.3
Estimated GFR > 60 ml/min	147/770	19.5	135/791	17.6	֥-	1.13 (0.87-1.4
Left ventricular ejection fraction					i	
≥50%	158/782	20.6	144/796	18.7		1.14 (0.88-1.4
<50%	33/111	31.5	26/115	24.2	- •	1.35 (0.73-2.4
Geographic region						
North America	89/381	24.2	61/371	17.3	_ -	1.57 (1.09-2.2
Europe	111/534	21.1	102/541	19.6	- -	1.09 (0.81-1.4
Other	3/33	9.6	13/45	29.6	←	0.24 (0.06-0.9
Non-left main diseased coronary arteries (core laboratory assessment)						
0	33/163	20.7	23/167	14.3	-	1.55 (0.86-2.7
1	60/292	21.2	61/292	21.9		0.94 (0.62-1.4
2	79/325	25.0	50/295	17.8	· -	1.58 (1.06-2.3
3	31/162	19.2	37/182	20.7	—	0.93 (0.54-1.5
Left main bifurcation or trifurcation stenosis ≥50% (core laboratory assessment)						
Yes	171/771	22.7	136/741	19.0	֥-	1.24 (0.96-1.6
No	32/171	19.2	35/195	18.9		1.05 (0.62-1.2
SYNTAX score (site reported)	-					
≤22	119/560	21.9	106/588	18.7		1.21 (0.90-1.6
23-32	84/386	22.2	70/366	20.0	- - -	1.16 (0.81-1.6
SYNTAX score (core laboratory assessment)			·			
≤22	49/294	17.2	58/364	16.7	-	0.99 (0.65-1.5
23-32	91/392	23.7	69/346	20.7	+-	1.22 (0.85–1.7
≥33	56/228	25.0	42/216	20.0		1.36 (0.86-2.1
	•		•		0.2 0.5 1.0 1.5 2.0	5.0

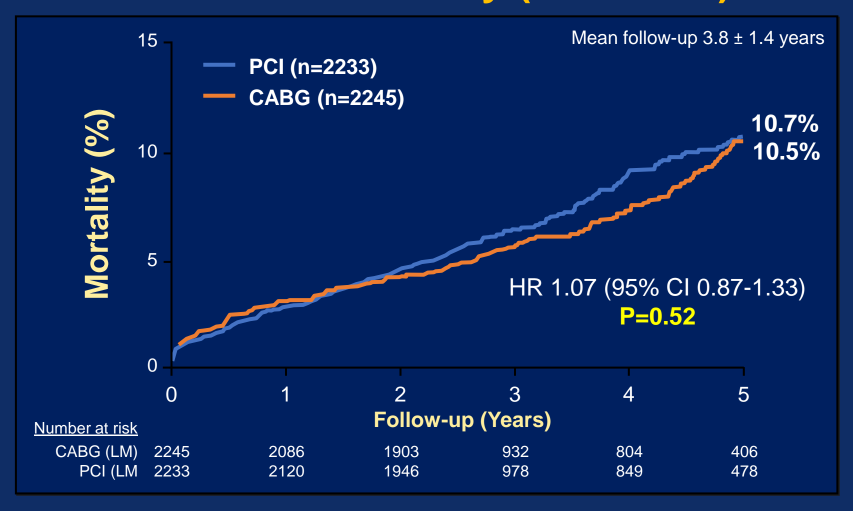
Role of Left Main PCI After EXCEL and NOBLE

Variables	EXCEL	NOBLE
Patients (no.)	1,905	1,201
Median follow-up	5 year	4.9 year
HR (95% CI), CABG/PCI		
Primary endpoint	1.19 (0.95-1.05)	1.58 (1.24-2.01)
All-cause death	1.38 (1.03-1.85)	1.08 (0.74-1.59)
Cardiac death	1.3 (-0.9-3.6)	0.99 (0.57-1.73)
MI	1.4 (-1.3-4.2)	2.99 (1.66-5.39)
Stroke	-0.8 (-2.4-0.9)	1.75 (0.86-3.55)
Revascularization	6.9 (3.7-10.0)	1.73(1.25-2.40)

NOBLE: Stent thrombosis (2% NOBLE vs. 1.8% EXCEL), non-procedural MI excluded (3% CABG vs. 8% PCI)



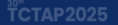
Individual-patient-data Analysis from 11 PCI vs. CABG Trials 11,518 randomized pts; 4,478 (38.9%) with left main ds. All-cause Mortality (Left Main)



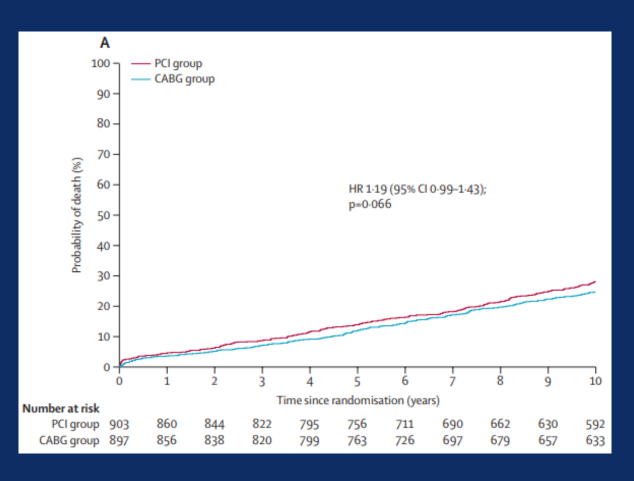


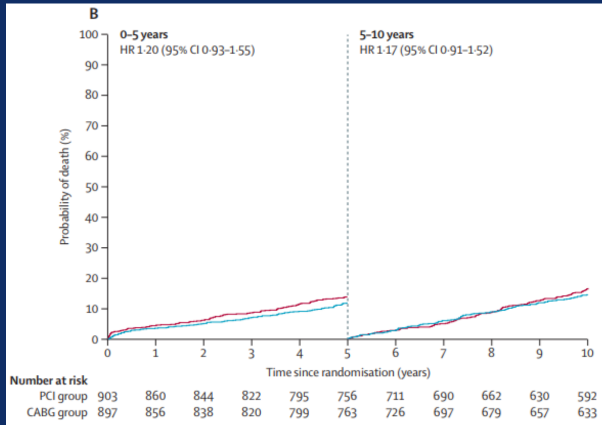
Individual-patient-data Analysis from 11 PCI vs. CABG Trials 11,518 randomized pts; 4,478 (38.9%) with left main ds. All-cause Mortality (LM patients)

	PCI (n=2,233)	CABG (n=2,245)	HR (95%CI]	P value	P _{int}
Overall mortality	10.7% (174)	10.5% (158)	1.07 [0.87, 1.33]	0.52	
Diabetes	16.5% (71)	13.4% (51)	1.34 [0.93, 1.91]	0.11	0.13
No diabetes	8.8% (104)	9.6% (107)	0.94 [0.72, 1.23]	0.65	0.13
SYNTAX score 0-22	8.1% (45)	8.3% (49)	0.91 [0.60, 1.36]	0.64	
SYNTAX score 23-32	10.8% (67)	12.7% (63)	0.92 [0.65, 1.30]	0.65	0.38 (0.06 for trend)
SYNTAX score ≥33	15.0% (56)	12.4% (45)	1.39 [0.94, 2.06]	0.10	



10-year outcomes of the randomized <u>SYNTAX</u> Extended Survival (SYNTAXES) study: *PCI (N=357) vs. CABG (N=348)*

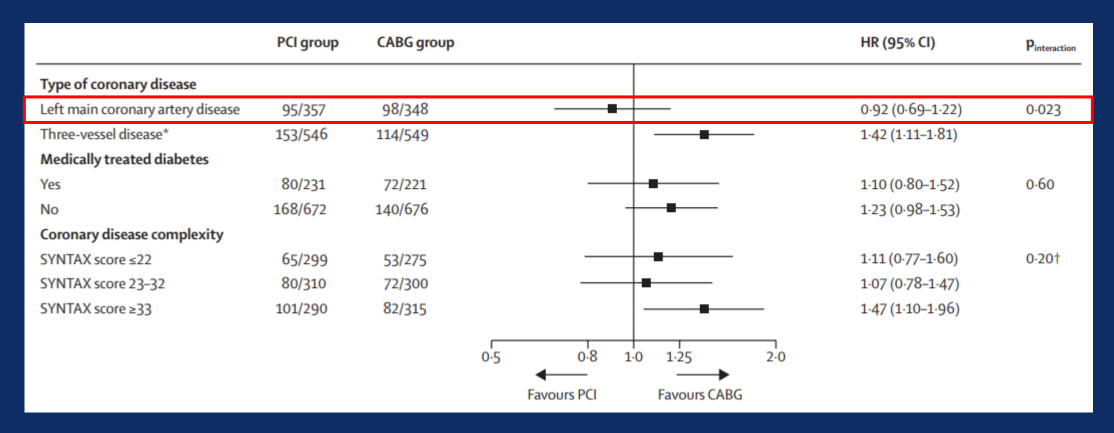






10-year outcomes of the randomized <u>SYNTAX</u> Extended Survival (SYNTAXES) study: *PCI (N=357) vs. CABG (N=348)*

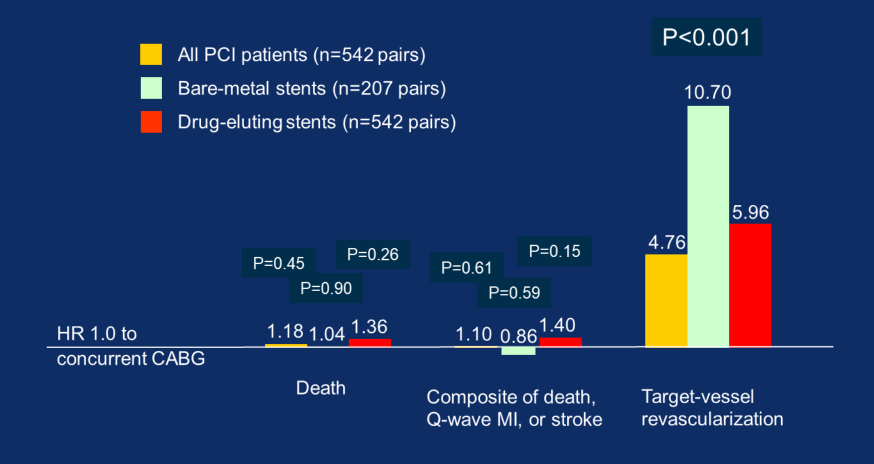
Prespecified Subgroup analysis of 10-year all-cause death





MAIN COMPARE Registry, 3-Year

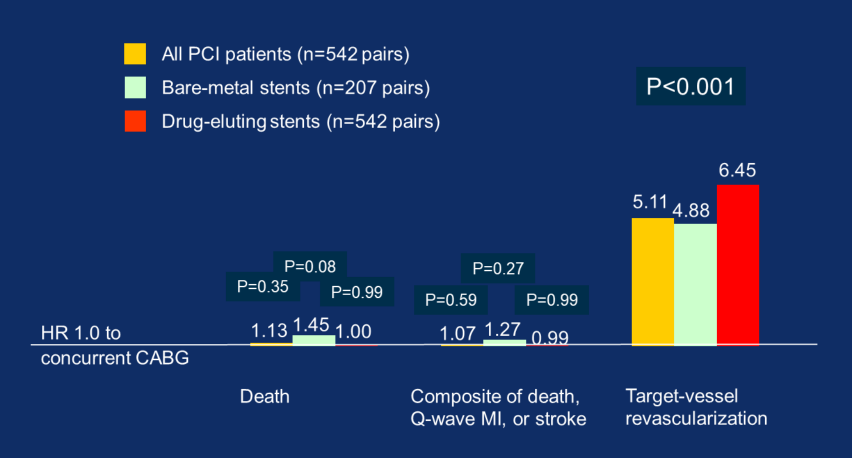
Adjusted HR by Use of PS Matching





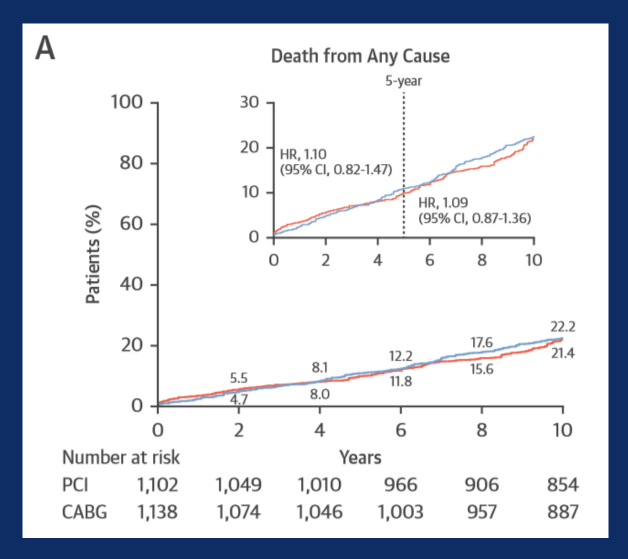
MAIN COMPARE Registry, 5-Year

Adjusted HR by Use of IPTW Method



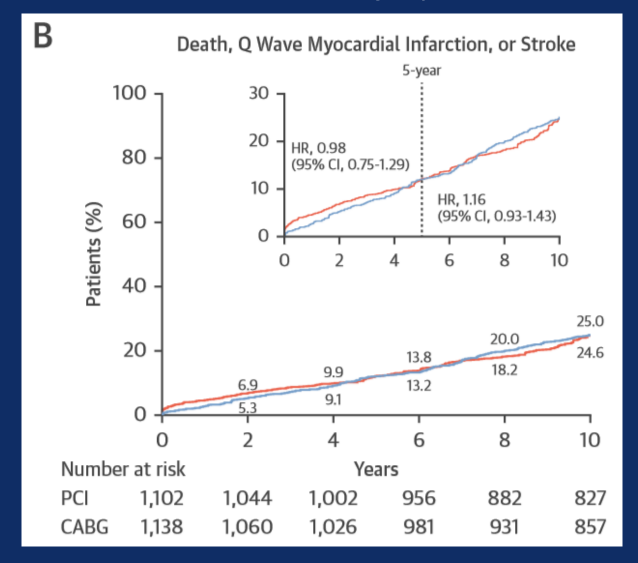


10-year outcomes of the MAIN-COMPARE registry: All-cause death



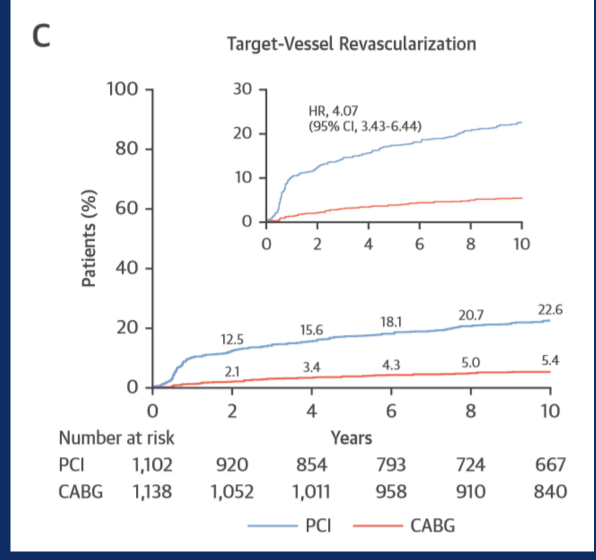


10-year outcomes of the MAIN-COMPARE registry: Death, Q-wave MI, or stroke





10-year outcomes of the MAIN-COMPARE registry: Target-Vessel Revascularization



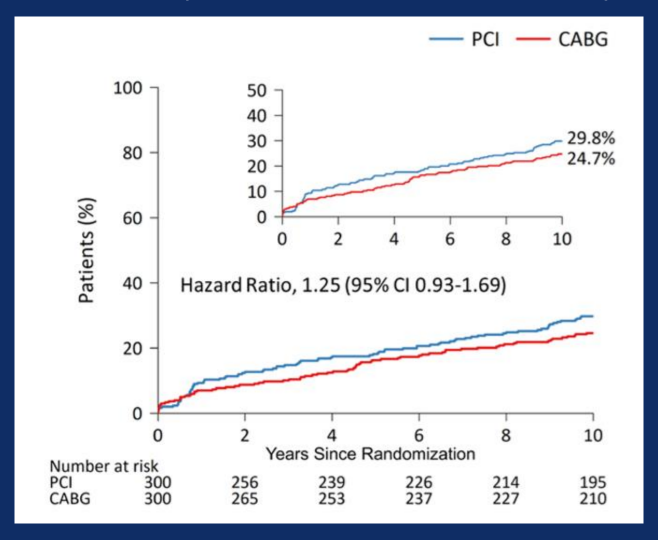


Hazard Ratios for Clinical Outcomes Before and After 5-Year of Follow-up

Outcome	Overall Cohort		Wave 1* (BMS)		Wave 2* (DES)	
	Hazard Ratio [†] (95% CI)	P value	Hazard Ratio [†] (95% CI)	P value	Hazard Ratio [†] (95% CI)	P value
Analyses with IPTW	N = 2240 patients (PCI 1102, CABG 1138)		N = 766 patients (BMS 318, CABG 448)		N = 1474 patients (DES 784, CABG 690)	
Death		0.64		0.05		0.15
0~5 years	1.10 (0.82–1.47)	0.53	1.65 (0.91–2.98)	0.10	1.02 (0.71–1.46)	0.91
>5 years	1.09 (0.87–1.36)	0.48	0.68 (0.46–1.02)	0.06	1.35 (1.00–1.81)	0.05
Composite outcome (death, Q-wave MI or stroke)		0.43		0.06		0.03
0~5 years	0.98 (0.75–1.29)	0.91	1.46 (0.84–2.53)	0.18	0.91 (0.66–1.27)	0.59
>5 years	1.16 (0.93–1.43)	0.19	0.67 (0.46–1.00)	0.05	1.46 (1.10–1.94)	0.009
TVR, All period	4.07 (3.43–6.44)	<0.001	4.45 (2.81–7.05)	<0.001	5.82 (3.77–9.01)	<0.001

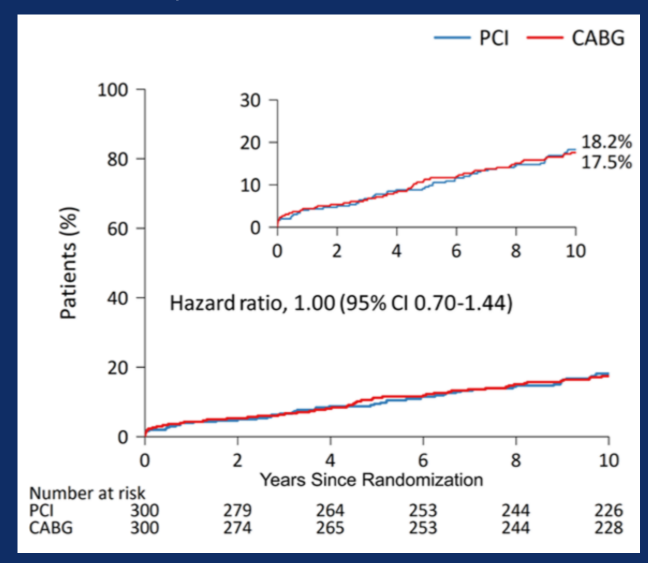


Extended Follow-Up of the PRECOMBAT trial: Primary composite outcome



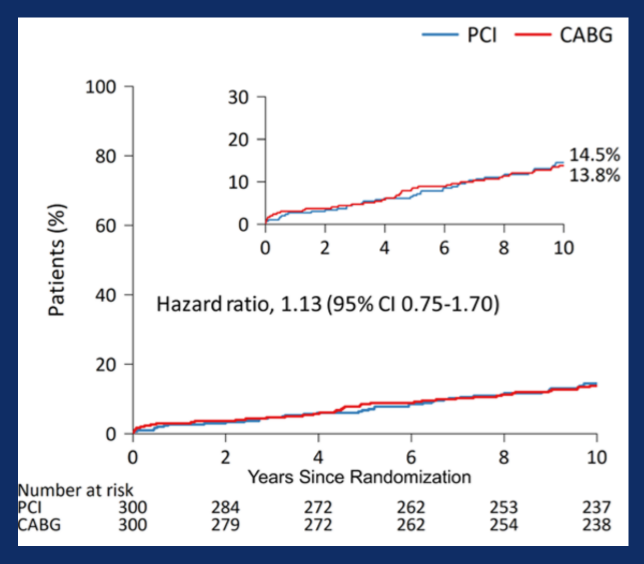


Extended Follow-Up of the PRECOMBAT trial: Death, MI, or Stroke



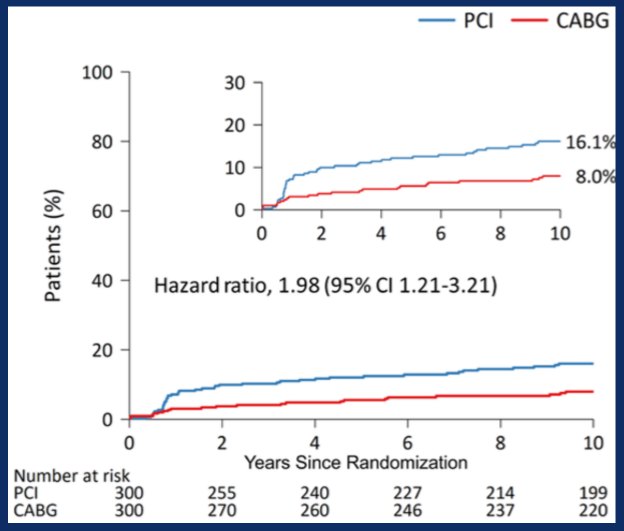


Extended Follow-Up of the PRECOMBAT trial : All-cause Death

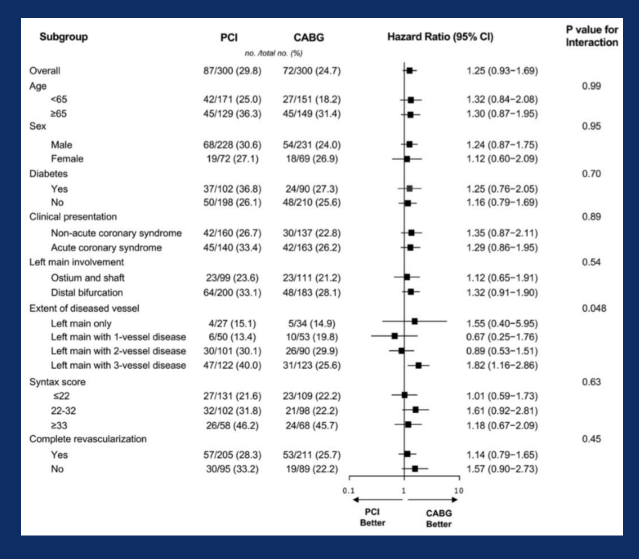




Extended Follow-Up of the PRECOMBAT trial: Target-Vessel revascularization

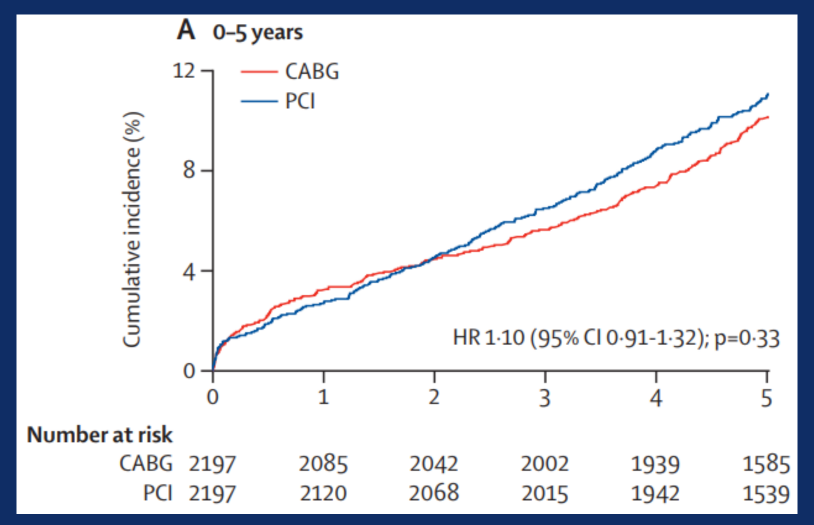


Extended Follow-Up of the PRECOMBAT trial



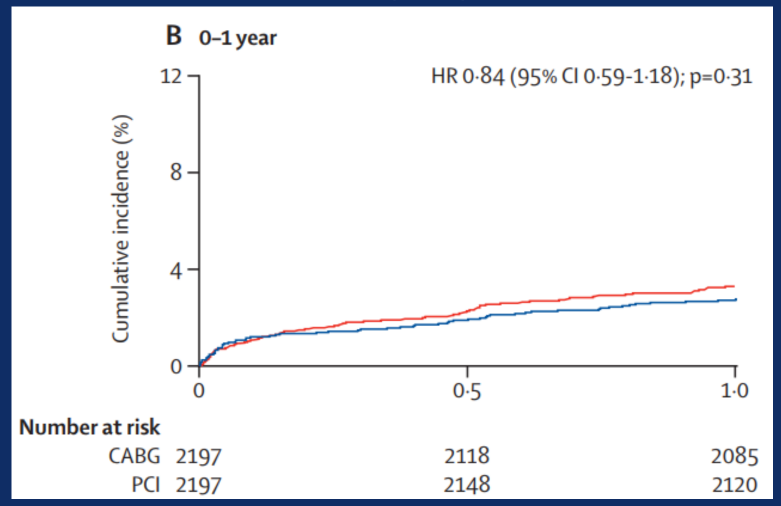


Individual patient data meta-analysis: SYNTAX, PRECOMBAT, NOBLE, EXCEL all-cause death (0-5Yr)



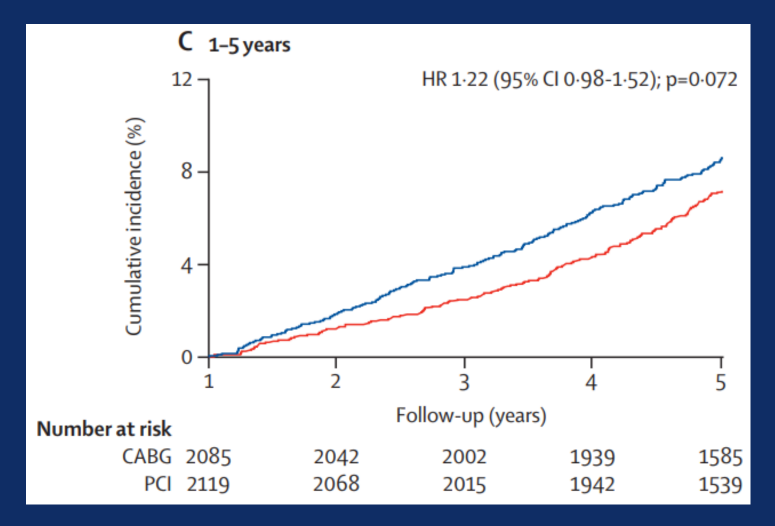


Individual patient data meta-analysis: SYNTAX, PRECOMBAT, NOBLE, EXCEL all-cause death (0-1Yr)



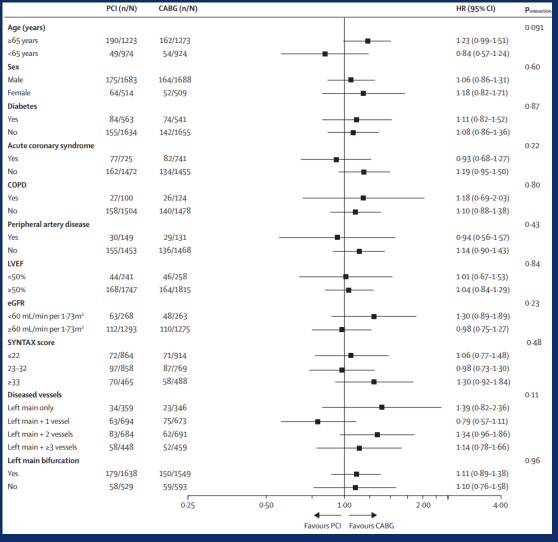


Individual patient data meta-analysis: SYNTAX, PRECOMBAT, NOBLE, EXCEL all-cause death (1-5Yr)





Individual patient data meta-analysis: SYNTAX, PRECOMBAT, NOBLE, EXCEL

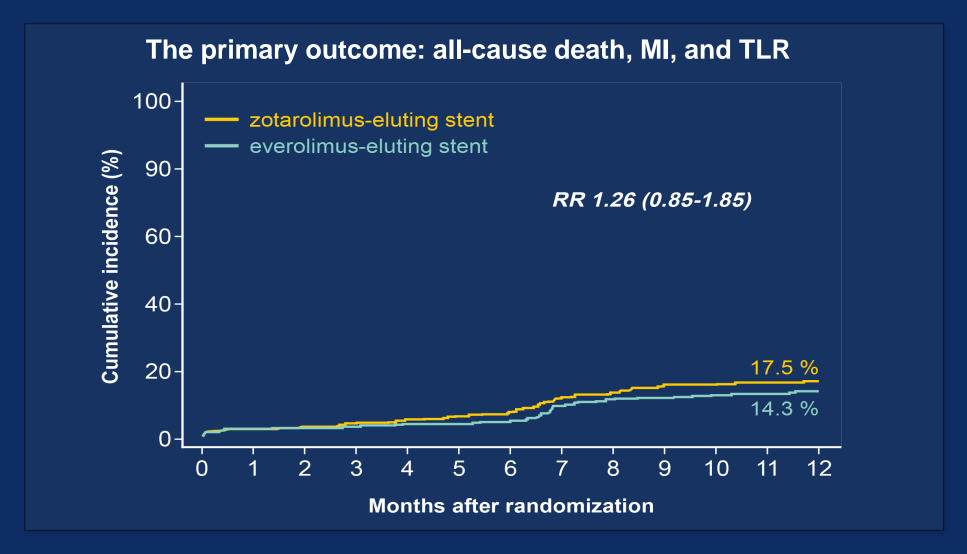




LM: DES vs. DES



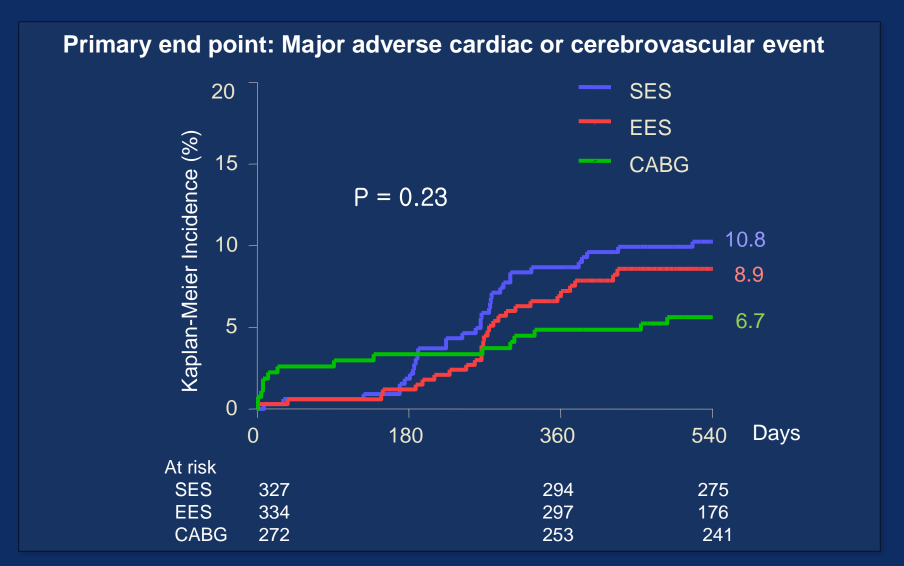
ISAR-LEFT MAIN 2 ZES vs. EES



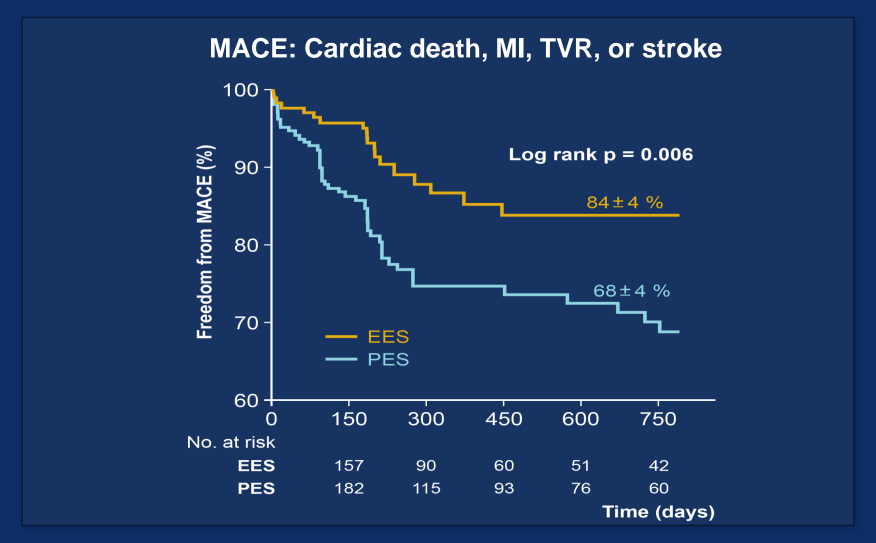


PRECOMBAT-2 Study

EES vs. SES

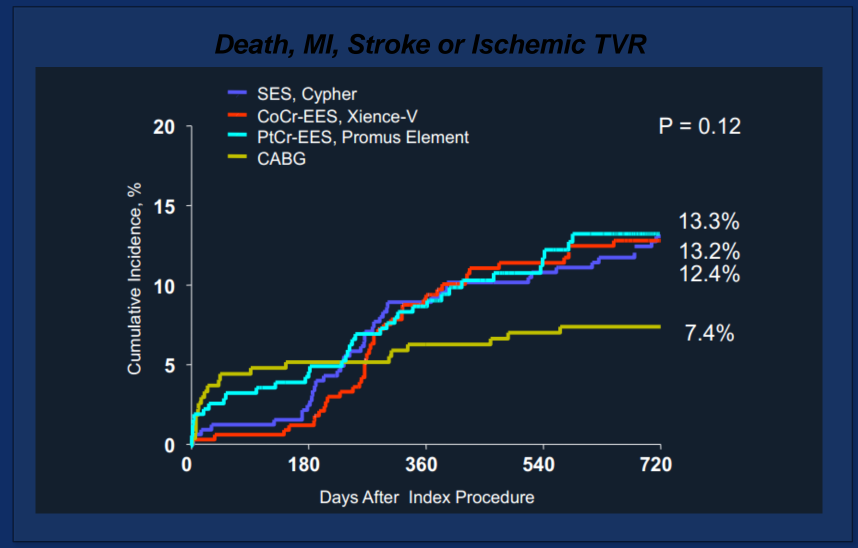


The ULMD Florence registry EES vs. PES





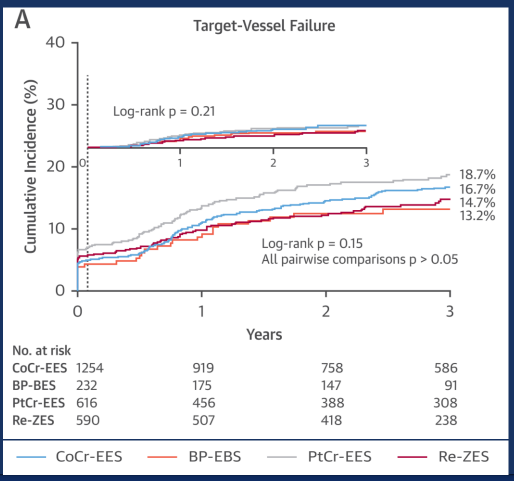
PRECOMBAT-3, 2 Year EES vs. SES





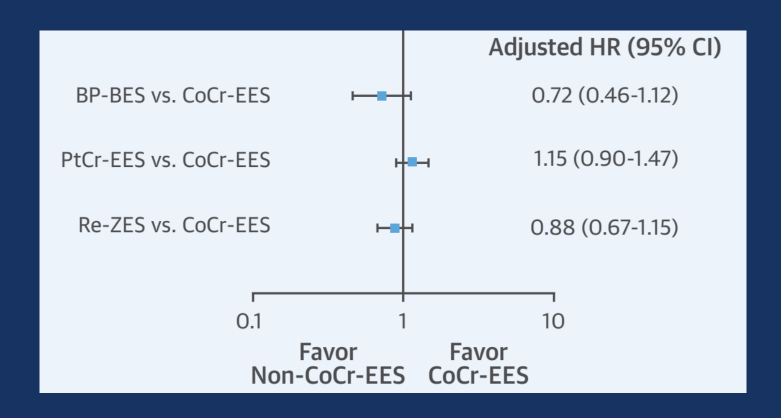
IRIS-MAIN Registry Comparison of 2nd generation DES

Target vessel failure: Cardiac death, Target vessel MI, or TVR



IRIS-MAIN Registry Comparison of 2nd generation DES

Target vessel failure: Cardiac death, Target vessel MI, or TVR

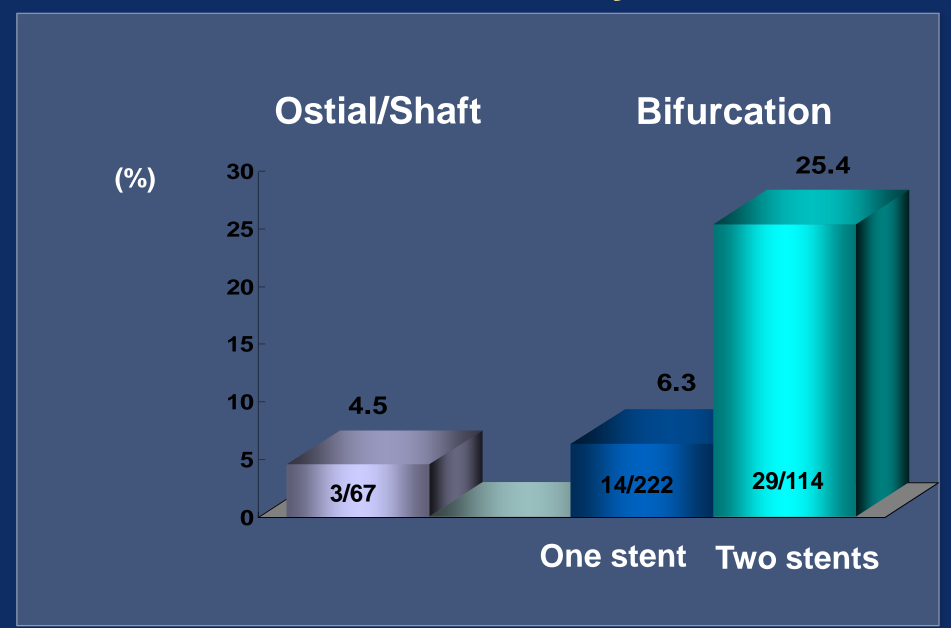




Distal bifurcation vs. Ostial / Shaft lesion

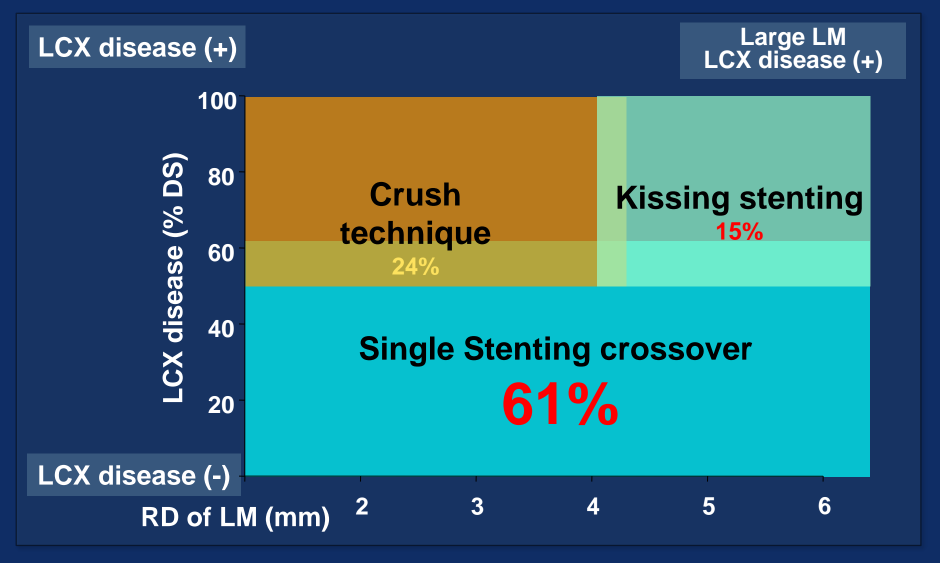


Restenosis at 2 year



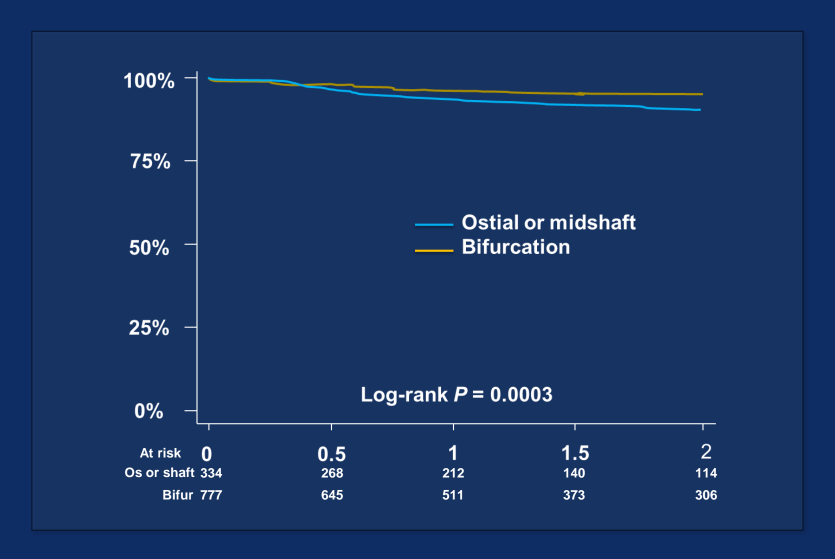


Lesion Specific Approach for LM Bifurcation





Bifurcation vs. Ostial / midshaft lesions TLR: Treated with DES

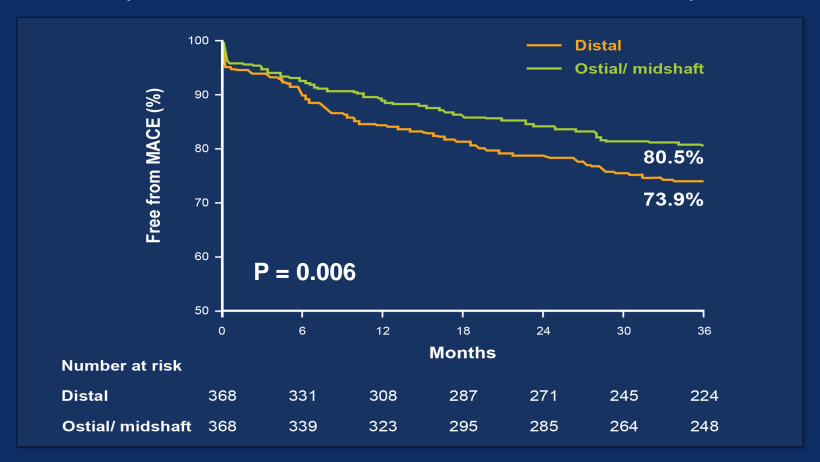






Distal bifurcation vs. Ostial/midshaft

A subgroup of <u>DELTA</u> registry - propensity score-matched groups (Distal bifurcation N=1130, Ostial/mid-shaft N=482)



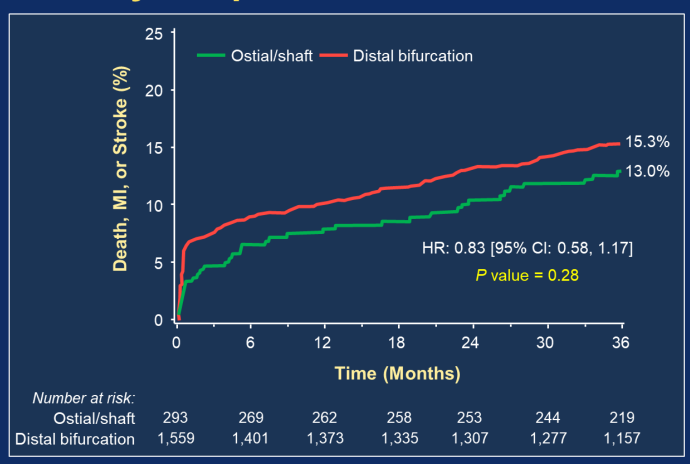
PCI for ostial/mid-shaft lesions was associated with better outcomes than distal bifurcation lesions in LM stenting.



Distal bifurcation vs. Ostial/midshaft

Post-hoc analysis of EXCEL Trial (Distal bifurcation N=1559, Ostial/mid-shaft N=293)

Primary Endpoint: Death, MI or Stroke

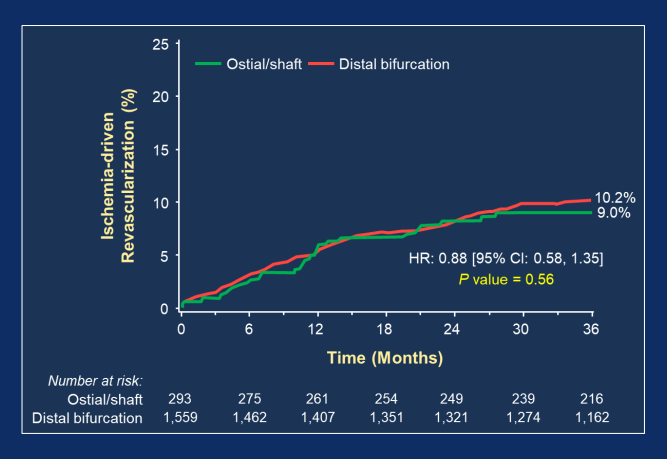




Distal bifurcation vs. Ostial/midshaft

Post-hoc analysis of EXCEL Trial (Distal bifurcation N=1559, Ostial/mid-shaft N=293)

Ischemia-driven Revascularization

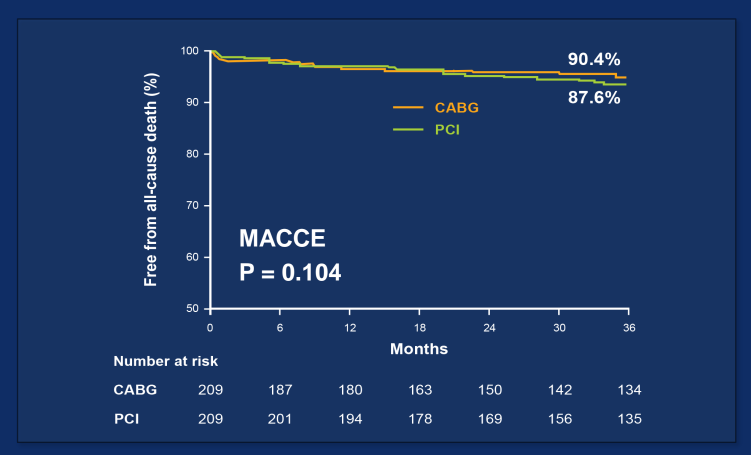




PCI vs. CABG for Ostial/Midshaft LM stenosis

A subgroup of <u>DELTA</u> registry (PCI, 482; CABG, 374 patients)

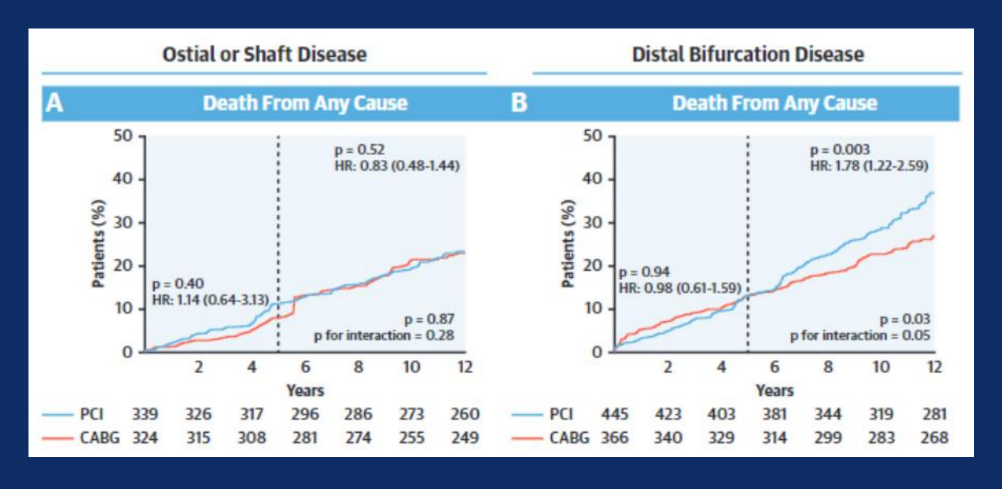
The results of propensity score-matched groups



PCI for ostial/midshaft lesions was associated with clinical outcomes comparable to those observed with CABG

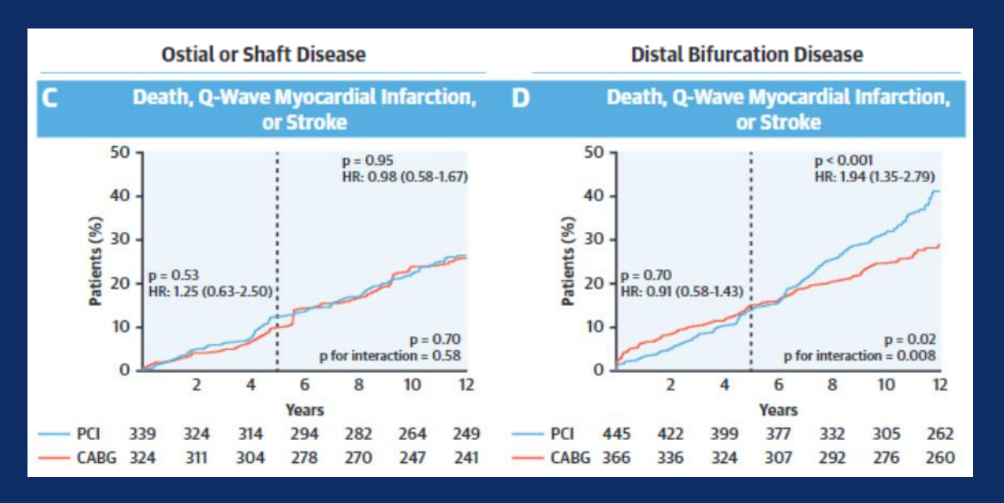


DES vs. CABG for LM Ostial/Shaft & Bifurcation MAIN-COMPARE registry



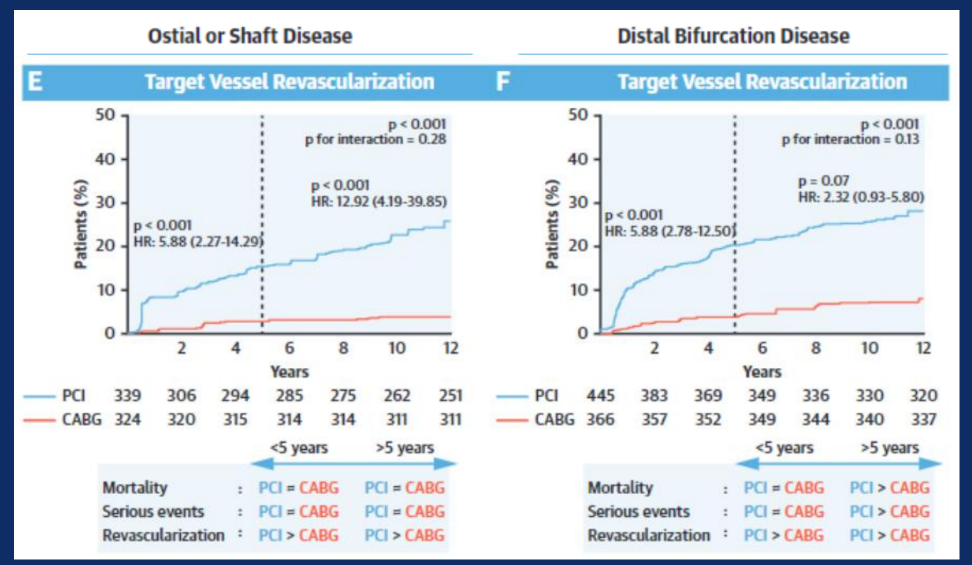


DES vs. CABG for LM Ostial/Shaft & Bifurcation MAIN-COMPARE registry



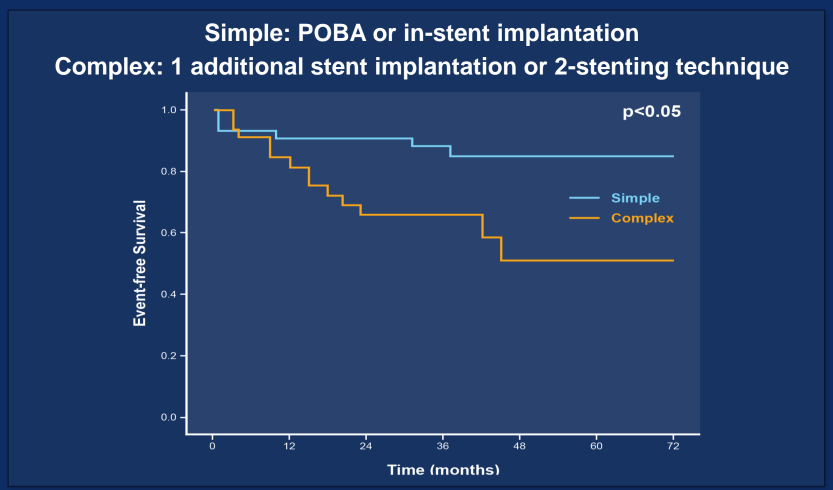


DES vs. CABG for LM Ostial/Shaft & Bifurcation



Distal LM Restenosis

UDLM-ISR subgroup of The <u>CORPAL</u> Registry (N=79)



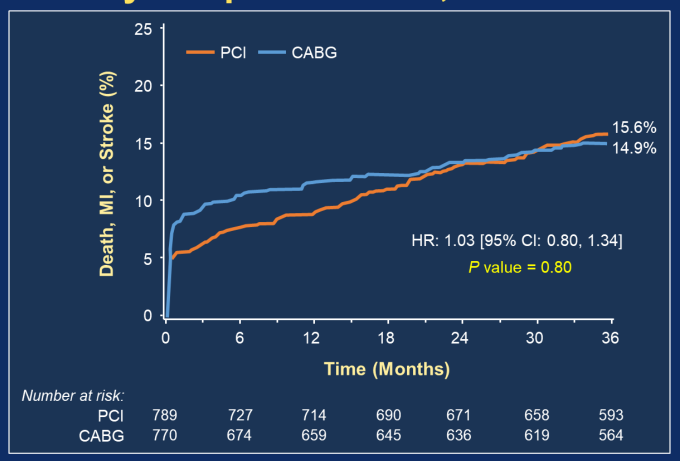
A simple strategy appeared to be a good treatment option, associated with a lower event rate at follow-up.



PCI vs. CABG for Distal Bifurcation LM stenosis

Post-hoc analysis of EXCEL Trial (Distal bifurcation N=1559, Ostial/mid-shaft N=293)

Primary Endpoint: Death, MI or Stroke

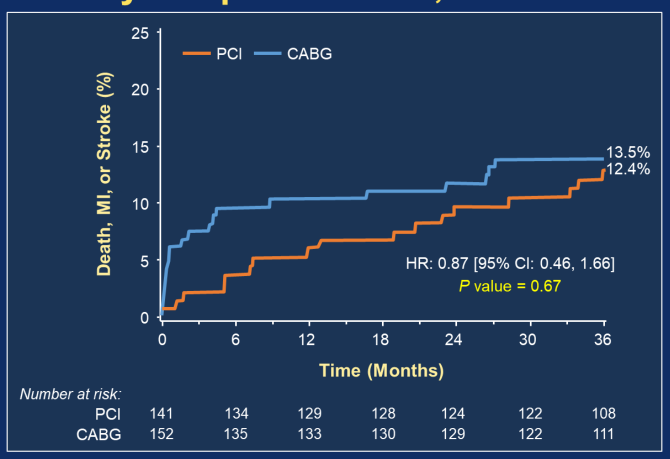




PCI vs. CABG for Ostial/Midshaft LM stenosis

Post-hoc analysis of EXCEL Trial (Distal bifurcation N=1559, Ostial/mid-shaft N=293)

Primary Endpoint: Death, MI or Stroke

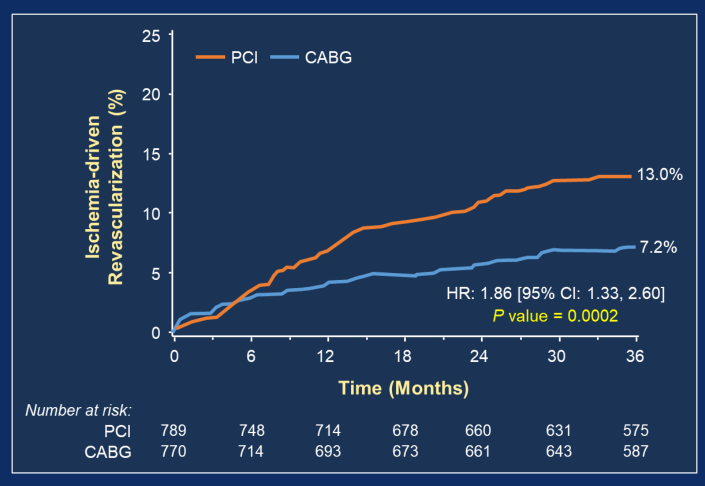




PCI vs. CABG for Distal Bifurcation LM stenosis

Post-hoc analysis of EXCEL Trial (Distal bifurcation N=1559, Ostial/mid-shaft N=293)

Ischemia-driven Revascularization



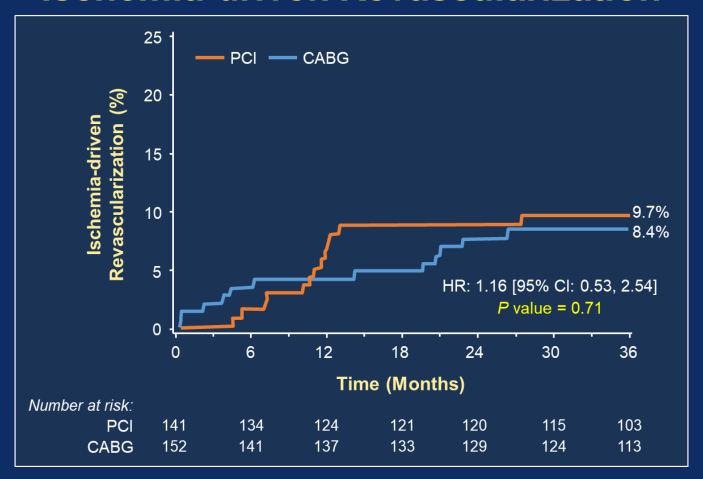




PCI vs. CABG for Ostial/Midshaft LM stenosis

Post-hoc analysis of EXCEL Trial (Distal bifurcation N=1559, Ostial/mid-shaft N=293)

Ischemia-driven Revascularization





Mortality after LM reintervention ISAR-LEFT-MAIN and ISAR-LEFT-MAIN2 registry

TABLE 2 Procedural Findings at the Index Procedure				
	Alive (n = 119)	Dead (n = 47)	p Value	
	(II = 119)	(II = 47)	p value	
Location of left main lesion			0.88	
Ostial	8.4 (10/119)	8.5 (4/47)		
Distal/bifurcation	81.5 (97/119)	78.7 (37/47)		
Body	10.1 (12/119)	12.8 (6/47)		
Occluded right coronary artery	9.2 (11/119)	14.9 (7/47)	0.29	
Trifurcation morphology	14.3 (17/119)	10.6 (5/47)	0.53	
Stenting technique			0.26	
Single	45.4 (54/119)	57.4 (27/47)		
T-stenting	10.1 (12/119)	4.3 (2/47)		
Culotte stenting	44.5 (53/119)	38.3 (18/47)		
Kissing balloon technique	55.5 (66/119)	34.0 (16/47)	0.01	
Coronary artery dominance			0.49	
Left	8.4 (10/119)	10.6 (5/47)		
Right	82.4 (98/119)	74.5 (35/47)		
Balanced	9.2 (11/119)	14.9 (7/47)		
Stent type			0.02	
Sirolimus-eluting stent	29.4 (35/119)	8.5 (4/47)		
Zotarolimus-eluting stent	27.7 (33/119)	44.7 (21/47)		
Paclitaxel-eluting stent	17.6 (21/119)	23.4 (11/47)		
Everolimus-eluting stent	25.2 (30/119)	23.4 (11/47)		

Values are % (n/N).

TABLE 3 Mortality After Target Lesion Revascularization According to Lesion Location and Revascularization Strategy				
	Mortality at 3 Years	Mortality at 5 Years	p Value	
Lesion location Ostial Distal/bifurcation Body	14.3 (2) 20.6 (27) 23.7 (4)	31.8 (4) 29.3 (37) 36.4 (6)	0.90	
Underlying stenting technique Single T-stenting Culotte	20.3 (16) 14.9 (2) 21.5 (15)	35.5 (27) 14.9 (2) 26.9 (18)	0.30	
Revascularization type			0.90	

Values are % by Kaplan-Meier estimate (n).

CABG

PTCA

Stenting



18.1 (3)

24.1 (19)

16.5 (11)

24.4 (4)

31.5 (24)

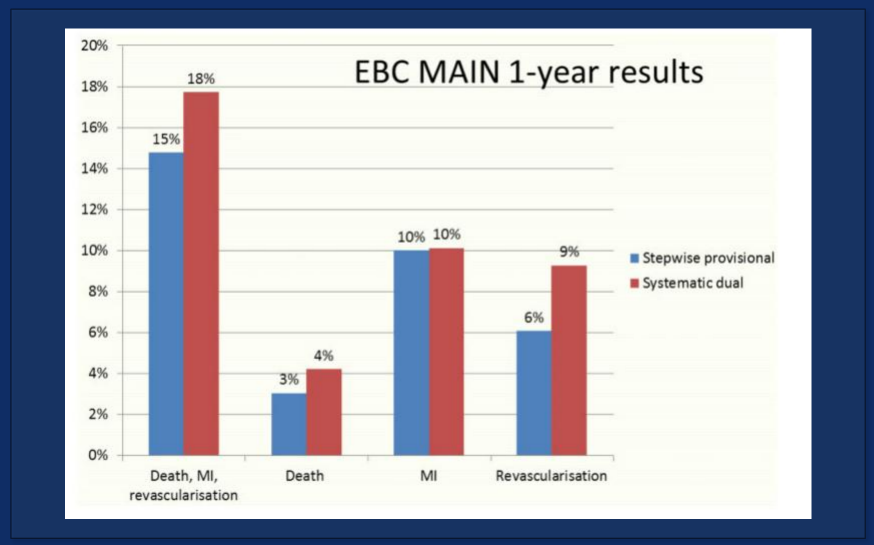
29.9 (19)

Simple cross vs. Two-stent technique



EBC MAIN trial

LM bifurcation: 1 vs. 2 stent tech.

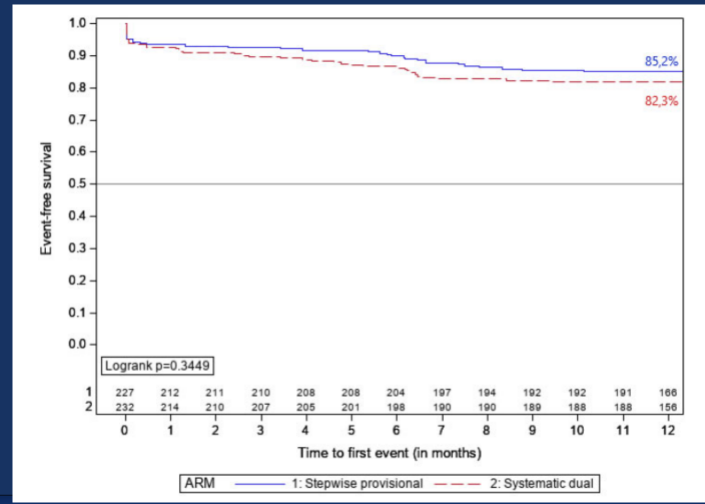




EBC MAIN trial

LM bifurcation: 1 vs. 2 stent tech.

Primary Endpoint: a composite of death, myocardial infarction, and target lesion revascularization at 12month

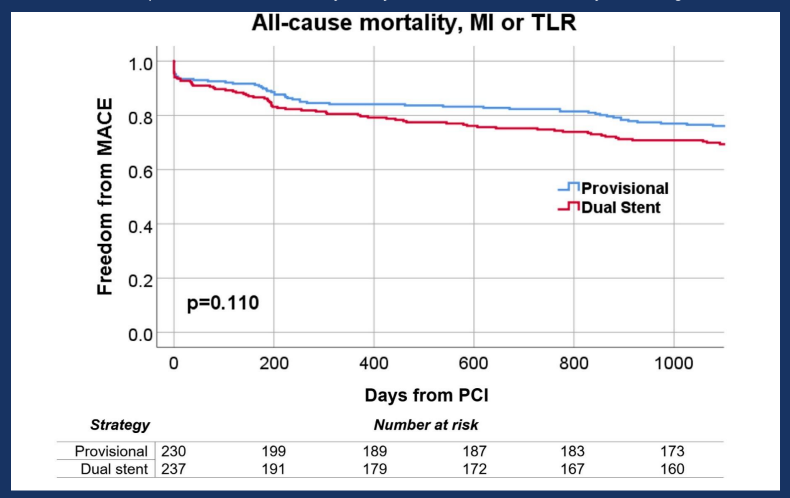




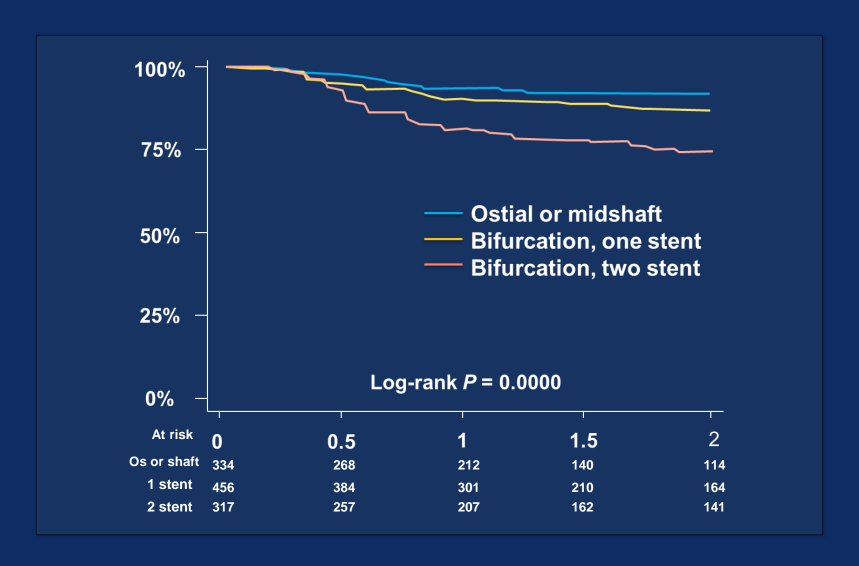
EBC MAIN trial

LM bifurcation: 1 vs. 2 stent tech.

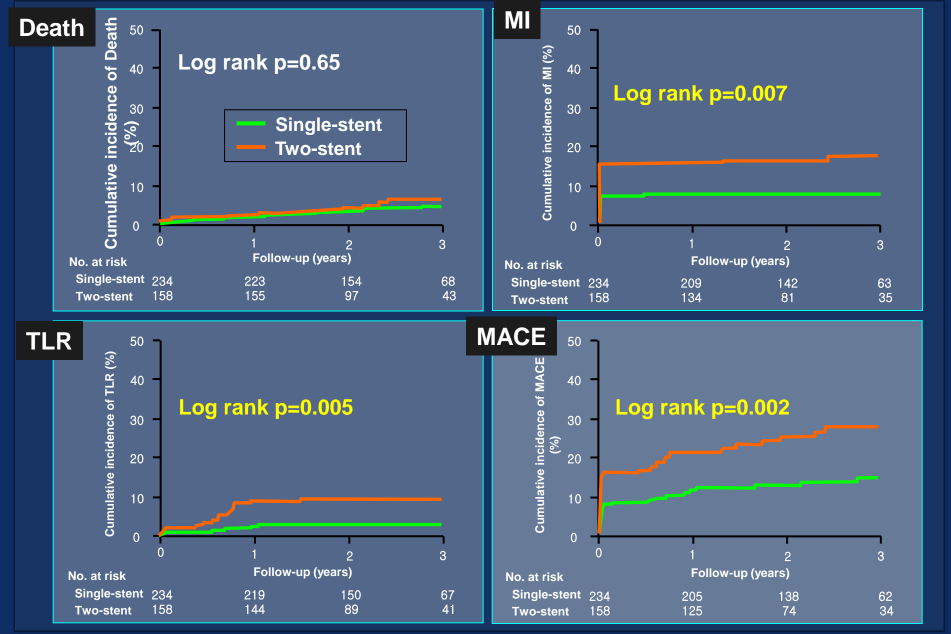
Primary Endpoint: MACE, defined as the composite of all-cause mortality, all myocardial infarction, or clinically driven target lesion revascularization (TLR) at 3 years



Ostial vs. 1 stent vs. 2 stent TLR: Treated with DES

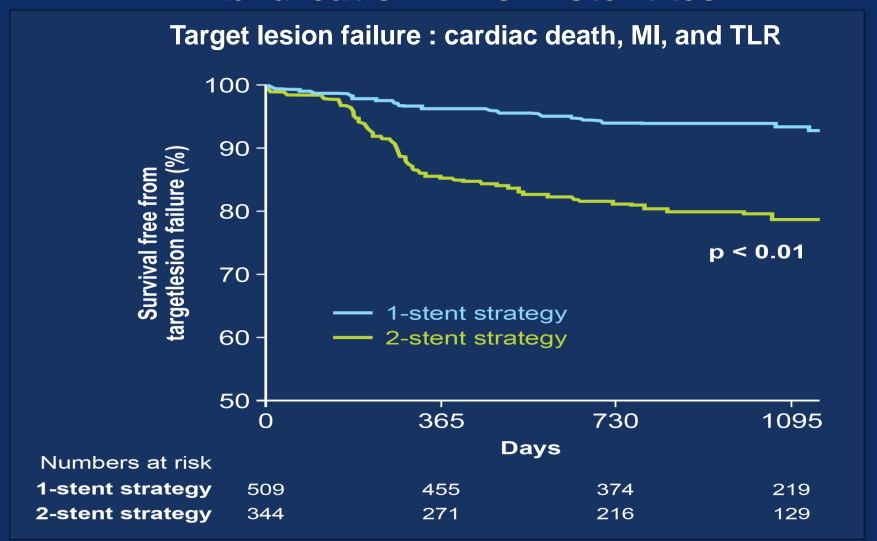


Single- vs. Two-Stent Strategy from MAINCOMPARE



COBIS Registry II

LM bifurcation: 1 vs. 2 stent tech.





IVUS-guided, Lesion-specific

Single stent

- Normal ostial LCX with MEDINA 1.1.0. or 1.0.0.
- Small LCX with < 2.5 mm in diameter
- Diminutive LCX
- Normal or focal disease in distal LCX

Two stent

- Diseased LCX with MEDINA 1.1.1., 1.0.1., or 0.1.1
- Large LCX with ≥ 2.5 mm in diameter
- Diseased left dominant coronary system
- Concomitant diffuse disease in distal LCX



Provisional vs. 2-stent technique for Simple and Complex Bifurcation Lesions - The DEFINITION Study

Adjusted HR with 2-stent technique			
	Simple Complex		
In-hospital			
MI	0.76 (0.45–1.28)	0.58 (0.35–0.94)	
Cardiac death	_	0.53 (0.13–2.12)	
TLR	1.66 (0.41–1.66)	_	
MACE	0.68 (0.40–1.13)	0.58 (0.35–0.94)	
Stent thrombosis	6.68 (1.67–26.80)	_	
At 1 year			
MI	0.68 (0.40–1.13)	0.64 (0.40–1.03)	
Cardiac death	0.95 (0.38–2.34)	0.52 (0.28–0.97)	
TLR	1.78 (1.16–2.74)	1.07 (0.65–1.75)	
MACE	1.03 (0.75–1.42)	0.79 (0.57–1.08)	
Stent thrombosis	1.66 (0.62–4.45)	1.06 (0.42–1.69)	

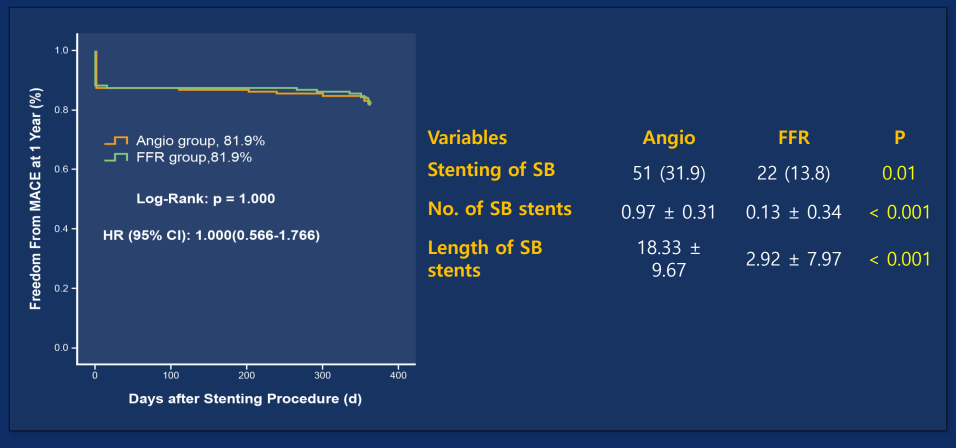
2-stent technique is still needed for complex bifurcation lesions



FFR- vs. Angio-guided Provisional Stenting

The Randomized DKCRUSH-VI Trial

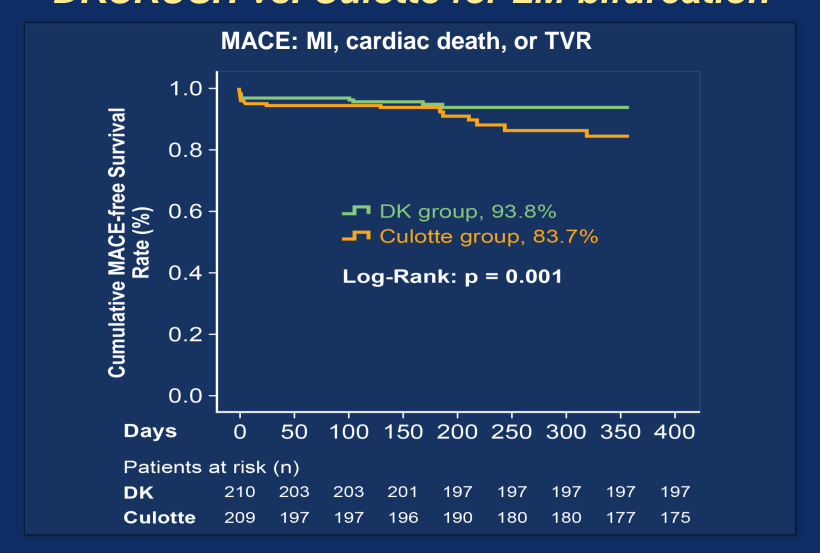
(160 patients with true bifurcation lesion in each group)



FFR-guided provisional stenting showed the similar outcomes with fewer stents



DKCRUSH III DKCRUSH vs. Culotte for LM-bifurcation





Risk Prediction of SB Occlusion

The RESOLVE Score System : a model built from 1545 Chinese patients with bifurcation

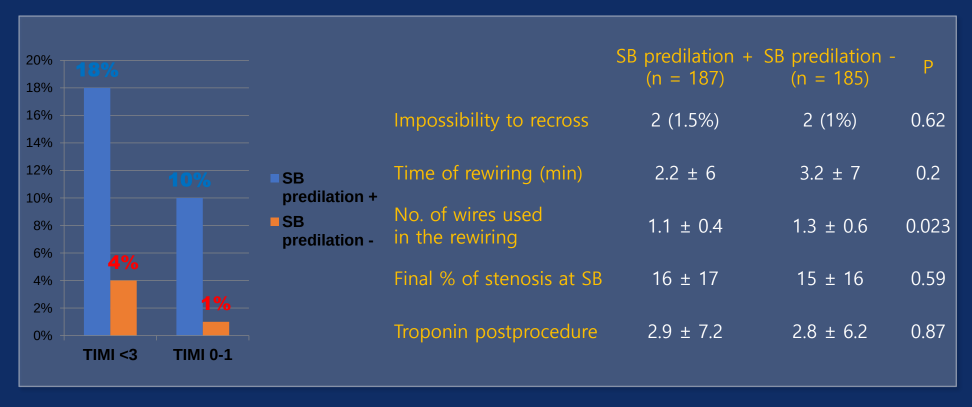
Scores Attributed to Ea			ROC Curve
Risk Factor	Level	Point	
Plaque distribution	At the opposite side of SB At the same side of SB	0	0.8-
MV TIMI flow grade before stenting	TIMI 3 TIMI 2 TIMI 1 TIMI 0	0 6 11 17	
Pre-procedural diameter stenosis of bifurcation core (%)	<50 50–<70 ≥70	0 2 3	Sensitivity 0.6
Bifurcation angle (°)	<70 70–<90 ≥90	0 4 6	AUC 0.77 (95% CI: 0.69 to 0.86)
Diameter ratio between MV/SB	<1.0 1.0–<1.5 1.5–<2.0 ≥2.0	0 2 6 9	Source of the Curve — RESORVE Model
Diameter stenosis of SB before MV stenting (%)	<50 50–<70	0 4	— RESORVE Score — Reference Line
	70–<90 ≥90	6 7	0.0 0.2 0.4 0.6 0.8 1

The RESOLVE score system can help identify patients at risk for SB occlusion during bifurcation stenting.



Effect of SB Predilation Before Provisional Stenting

A randomized study enrolling 372 patients with true bifurcation (SB predilation + vs. SB predilation -)

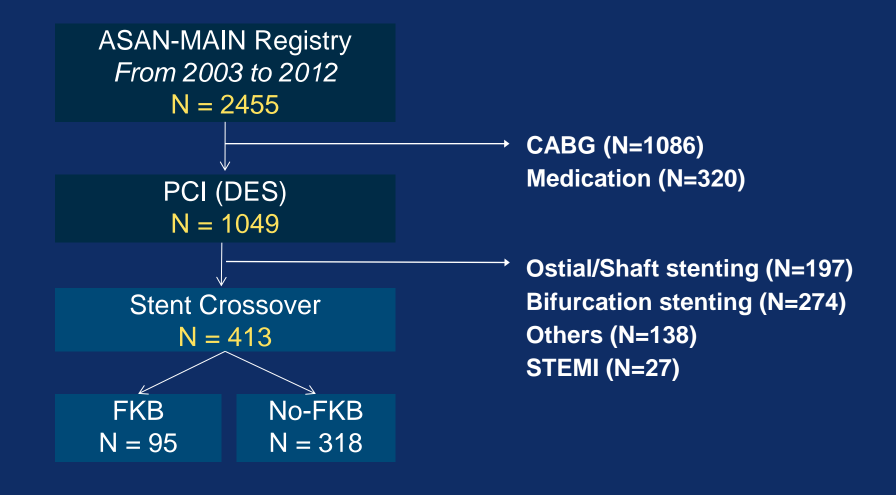


Predilation of the SB resulted in improved TIMI flow after MB stenting, not hindering SB rewiring.





With vs. Without Routine Kissing Balloon Inflation (FKB)



With vs. Without Routine Kissing Balloon Inflation (FKB) 2- year Clinical Outcomes

	FKB (N=95)	Non-FKB (N=318)	Adjusted HR (95% CI)	P value
Death	4 (4.6%)*	12 (3.9%)	1.03 (0.28-3.82)	0.97
Death or MI	4 (4.6%)	13 (4.2%)	0.95 (0.26-3.51)	0.96
TVR	7 (8.1%)	14 (4.8%)	1.12 (0.40-3.11)	0.83
LM-TLR	7 (8.1%)	13 (4.4%)	1.32 (0.46-3.75)	0.60
Definite ST	0	0	NA	NA
MACE#	11 (12.5%)	26 (8.5%)	1.10 (0.49-2.49)	0.82

adjusted for age, DM, clinical presentation, stent No., pre- and post-stenting LCX DS



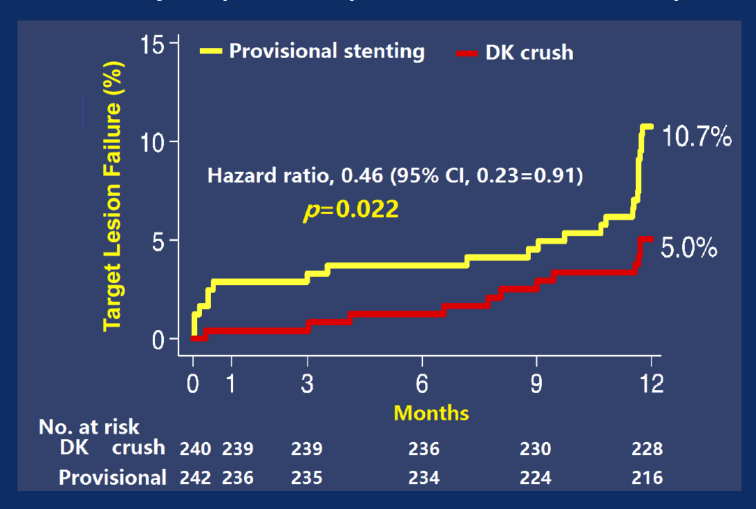
^{*} derived from Kaplan-Meier estimate

[#]composite of death, MI, or LM TLR

DKCRUSH-V Randomized Trial

DKCRUSH vs. Provisional stenting for LM distal bifurcation

Primary Endpoint: TLF (Cardiac death, TVMI, or TLR)

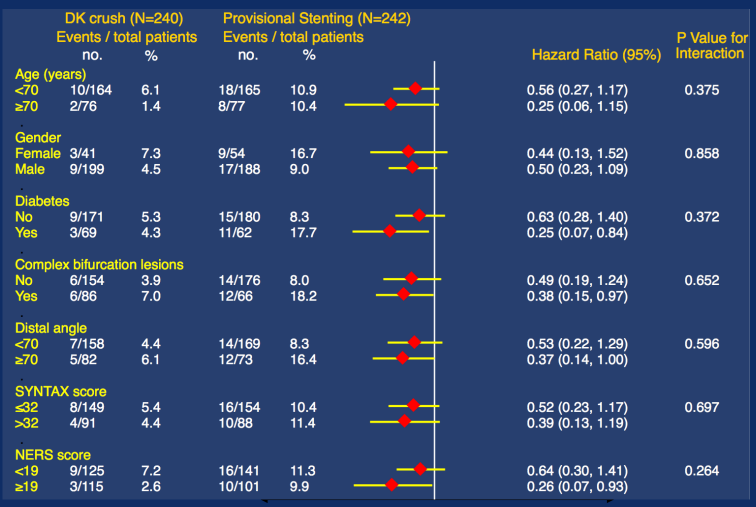




DKCRUSH-V Randomized Trial

DKCRUSH vs. Provisional stenting for LM distal bifurcation

Target Lesion Failure at 1-Year Subgroup analysis





Bifurcation technique



Bifurcation Coronary Disease

- 15~20% of PCI patients
- DES enhanced success rate, but have not resolved completely
- Dependable strategy not established
 - Rare studies evaluating anatomical intricacies
 - Lack of large randomized trials
 - Many anatomical variants
 - → Single technique can't fit all



Difficulties of Bifurcation PCI

- Risk of periprocedural complication
- Relatively high restenosis
- Not all lesions are the same
 - Size of vessels (Meaningful SB size ≥2.25mm)
 - Variable plaque distribution
 - Extent of SB disease
 - Variable angulation
- Higher risk of stent thrombosis

PCI techniques are mainly based on personal experiences from skilled operators



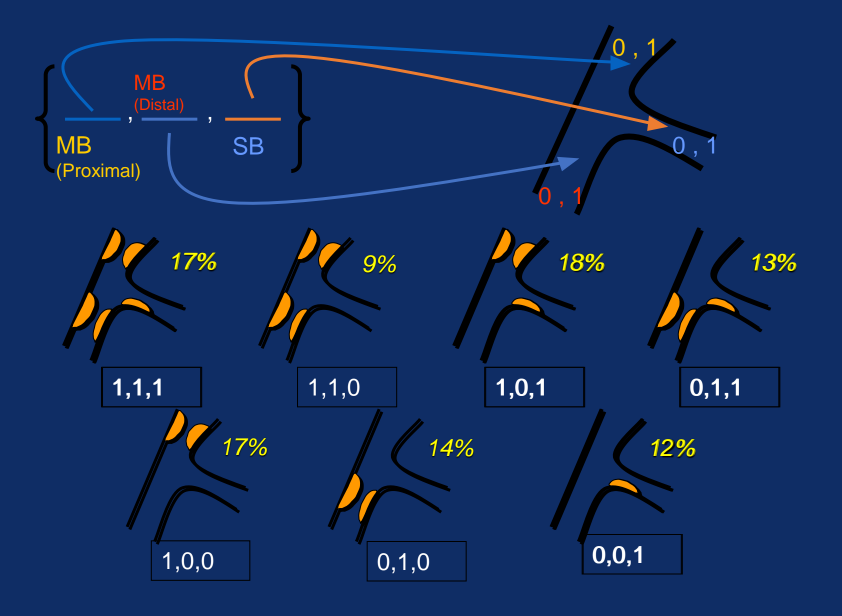
Factors to be considered for PCI strategy

Anatomical factors

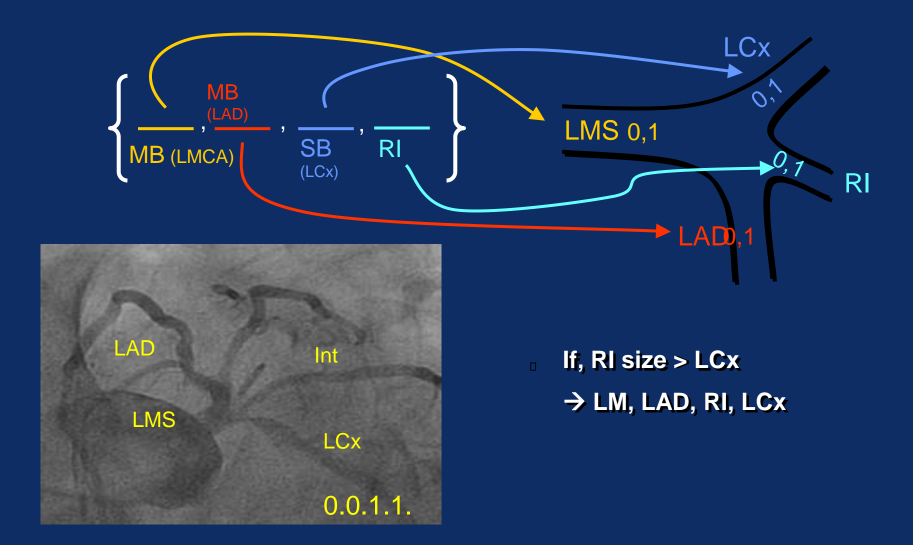
- LMCA bifurcation
- Location of plaque (Anatomical classification)
- Plaque or carina shift
- Angle btw SB and MB
- Dynamic change in bifurcation anatomy
- Modalities for objective anatomical evaluation
 - QCA, IVUS, FFR
- Selection of devices and strategies
 - DES vs. BMS
 - Single vs. Double stent techniques
 - Kissing balloon or not
 - Dedicated bifurcation stents



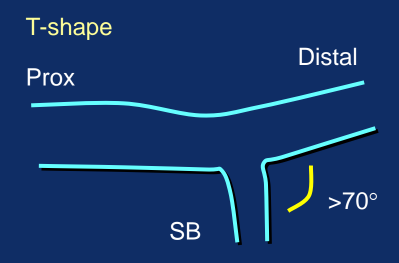
Medina Classification

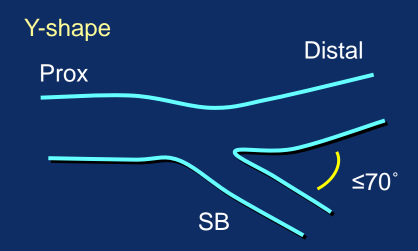


Trifurcation



Angulation



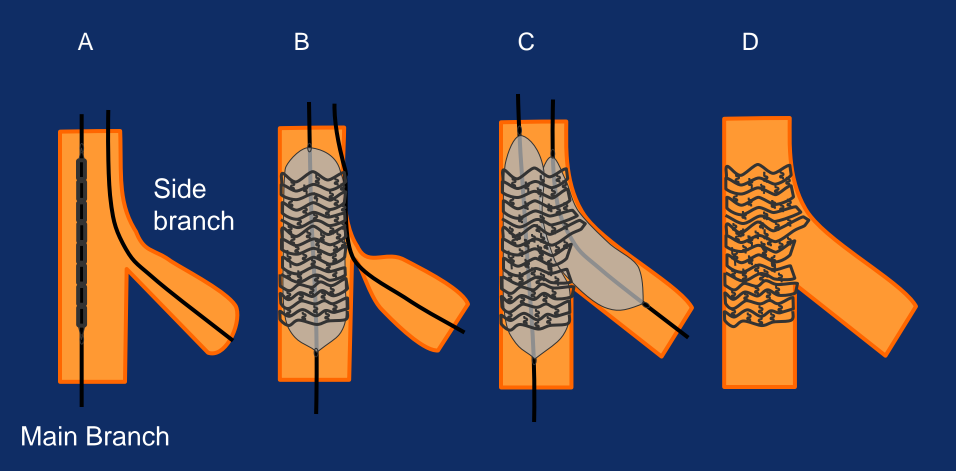


- Difficult SB access
- Less plaque shifting
- T-stenting better

- Easier SB access
- More plaque shifting
- Cullotte or Crush better

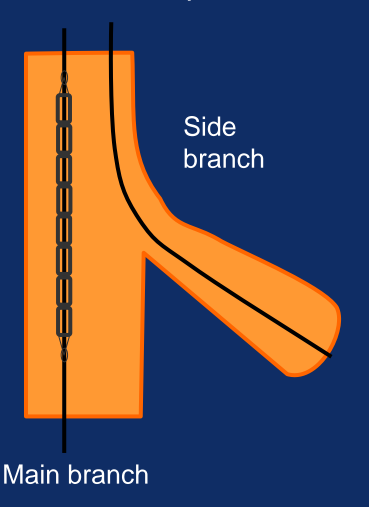


Normal or diminutive side branch ostium

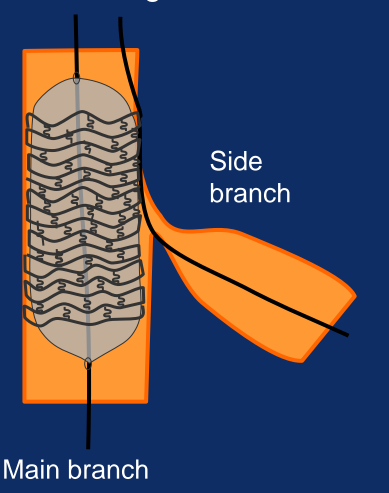




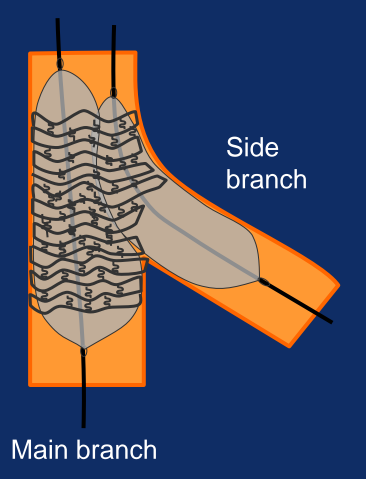
A. Wire both branches and predilate if needed



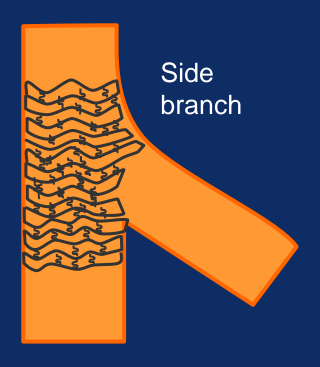
B. Stent the MB leaving a wire in the SB



C. Rewire the SB passing through the strut of the MB stent, remove the jailed wire, dilate toward SB, and perform FKB inflation



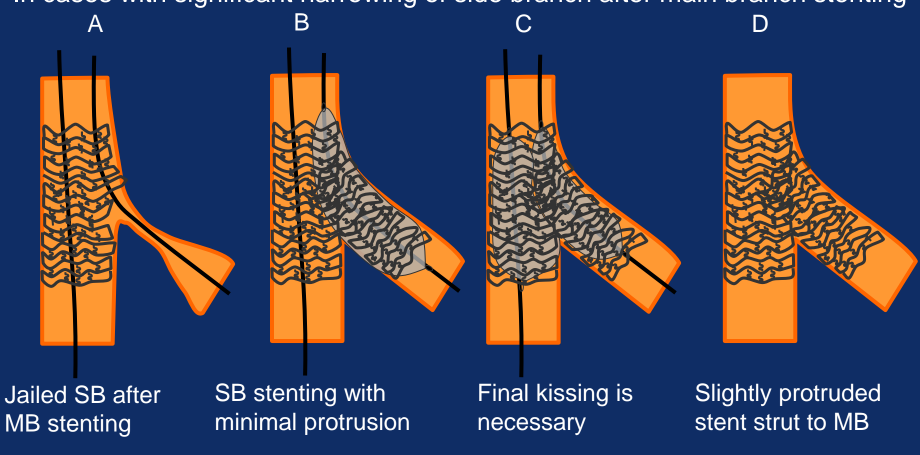
D. Final result



Main vessel



In cases with significant narrowing of side branch after main branch stenting



Advantages

Good SB scaffolding with angles >70°

Disadvantages

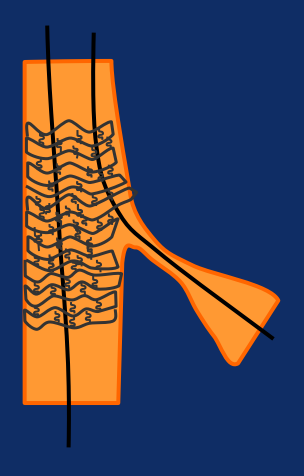
Potential gap at SB ostium

Protrusion of SB stent into the MB



In cases with significant narrowing of side branch after main branch stenting

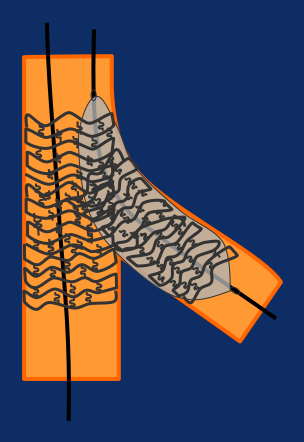
A. Jailed SB after MB stenting





In cases with significant narrowing of side branch after main branch stenting

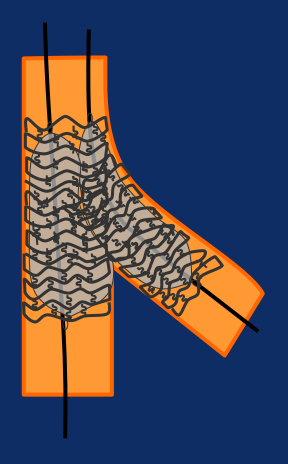
B. SB stenting with minimal protrusion





In cases with significant narrowing of side branch after main branch stenting

C. Final kissing is necessary





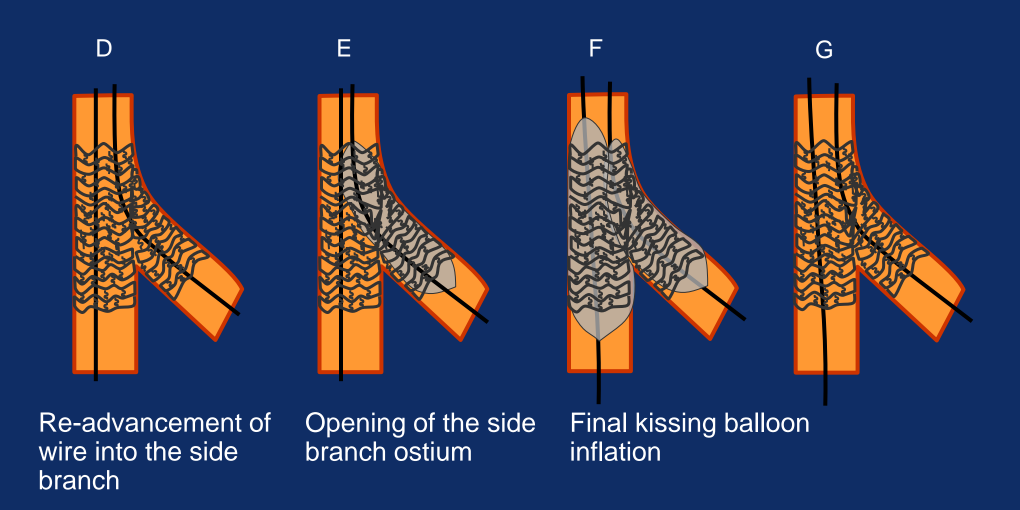
In cases with significant narrowing of side branch after main branch stenting

D. Slightly protruded stent strut to MB





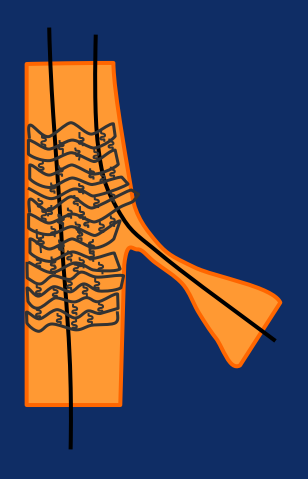
Final kissing balloon dilatation is mandatory





Final kissing balloon dilatation is mandatory

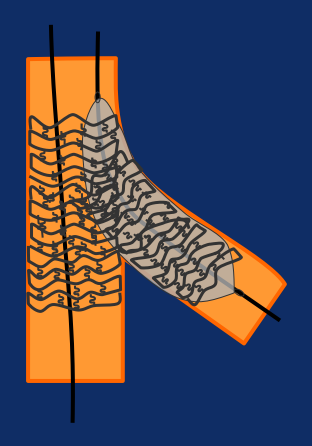
A. Jailed SB after MB stenting





Final kissing balloon dilatation is mandatory

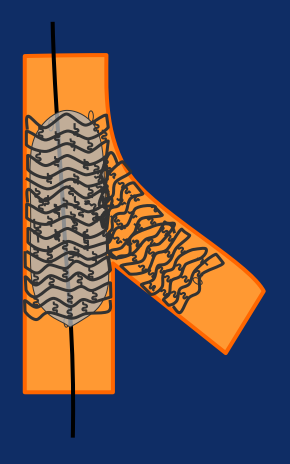
B. SB stenting with minimal protrusion





Final kissing balloon dilatation is mandatory

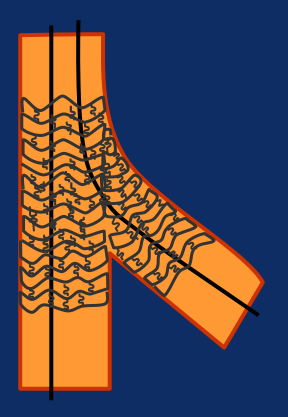
C. Remove SB balloon & wire, and inflate MB at high pressure to crush SB stent





Final kissing balloon dilatation is mandatory

D. Re-advancement of wire into the side branch





Final kissing balloon dilatation is mandatory

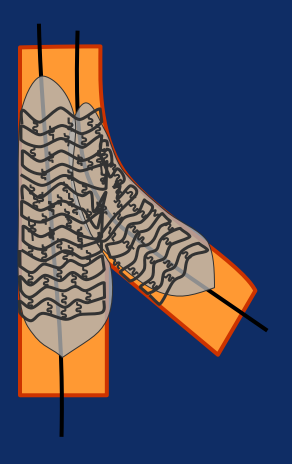
E. Opening of the side branch ostium





Final kissing balloon dilatation is mandatory

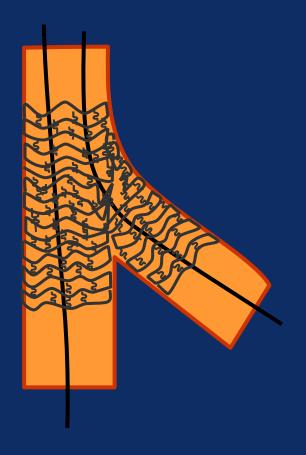
F. Final kissing balloon inflation





Final kissing balloon dilatation is mandatory

G. Final result





В C A D

Advantages

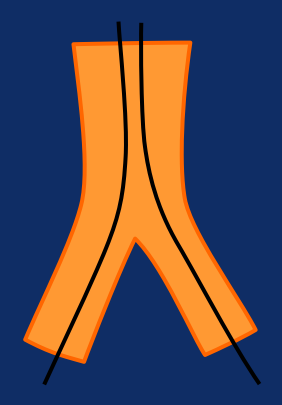
Compatible with 6-Fr guider Independent of bifurcation angle Predictable scaffolding

Disadvantages

Leaves multiple layers of strut Potential acute closure of MB



A. Wire both branches and predilate if needed

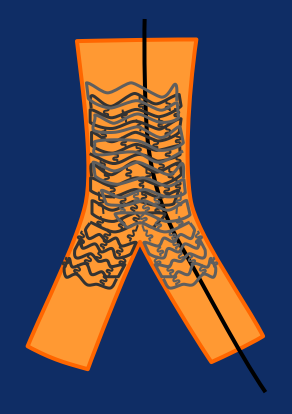




B. Deploy a stent in the more angulated branch (SB)



C. Rewire unstented branch, dilate the stent to unjail the MB, and expand a second stent into the unstented MB

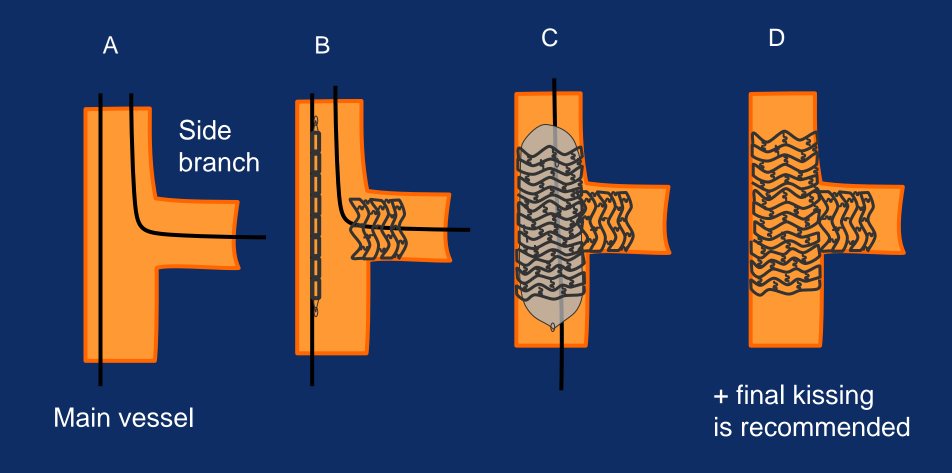




D. Final result after final kissing balloon

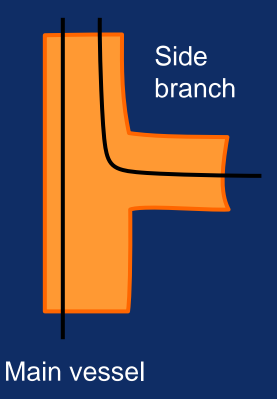






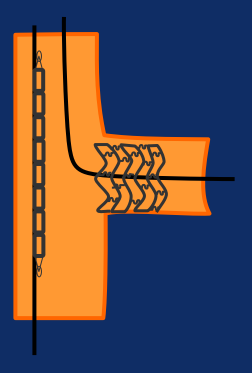


A. Wire both branches and predilate if needed



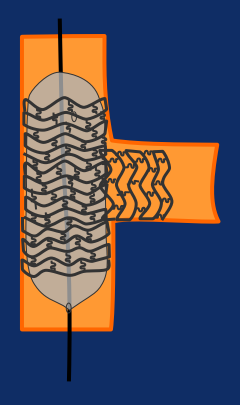


B. SB stent deployed at nominal pressure



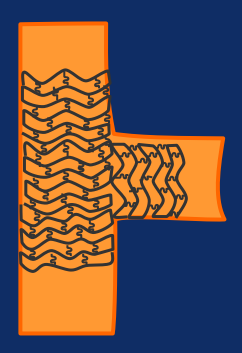


C. Remove balloon and wire from SB, And deploy the MB stent at high pressure





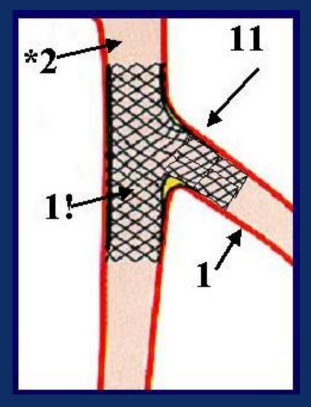
D. Rewire the SB and high-pressure dilatation, then final kissing inflation is recommended

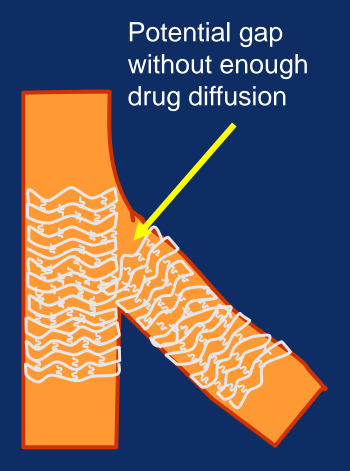




Limitation of Modified T Stenting

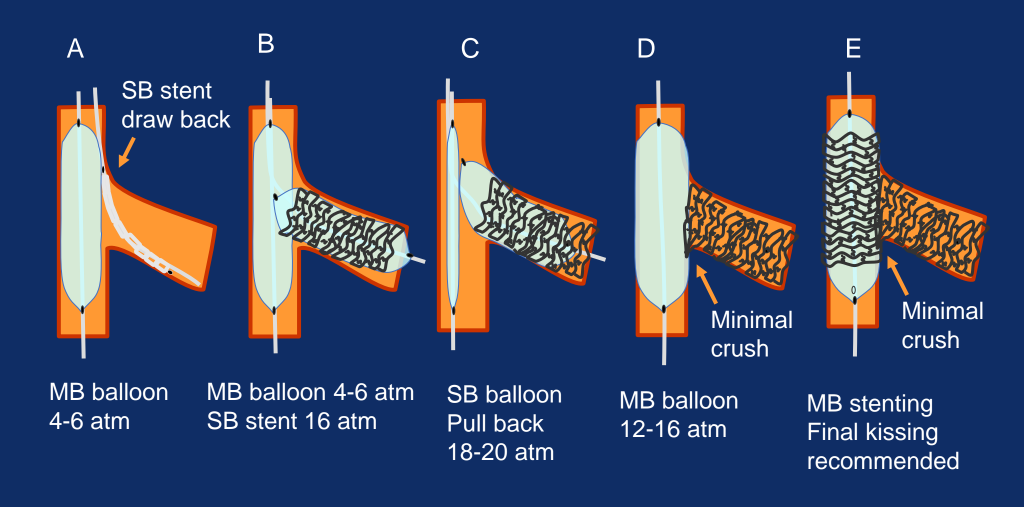
Restenosis site of T stenting in SIRIUS bifurcation



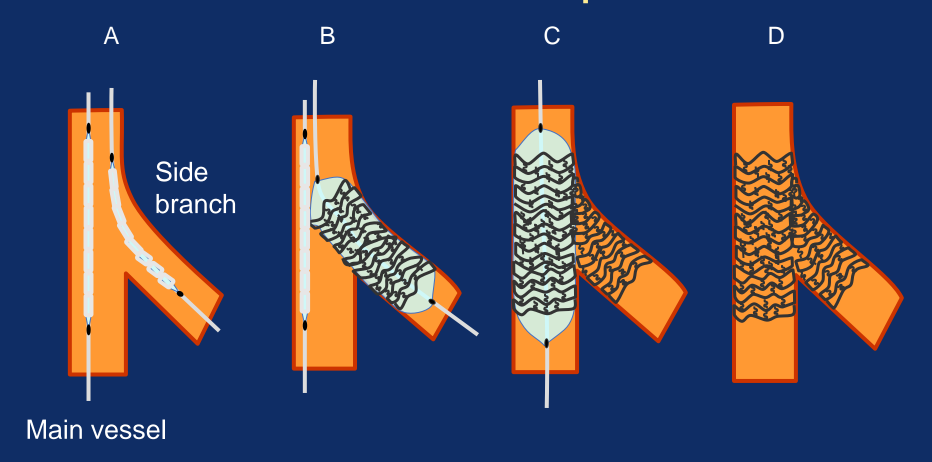


To prevent potential gap at the ostial side branch, the first stent should cover the entire surface of the side branch.

For Proper Ostial positioning



Crush Technique



Advantages

Relatively simple Low risk of SB occlusion Good coverage of SB ostium

Disadvantages

Difficult FKI
Requires 7 or 8-Fr guider
Leaves multiple layers of strut





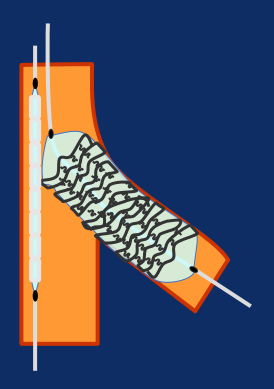
Crush Technique

A. Advance 2 stents



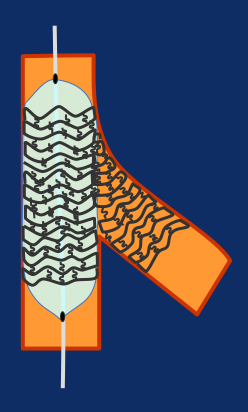
Crush Technique

B. Deploy the SB stent



Crush Technique

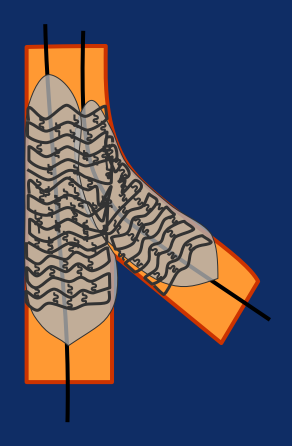
C. Deploy the main stent, then rewire SB and perform high-pressure dilatation





Crush Technique

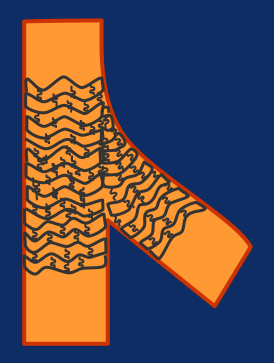
D. Perform final kissing inflation





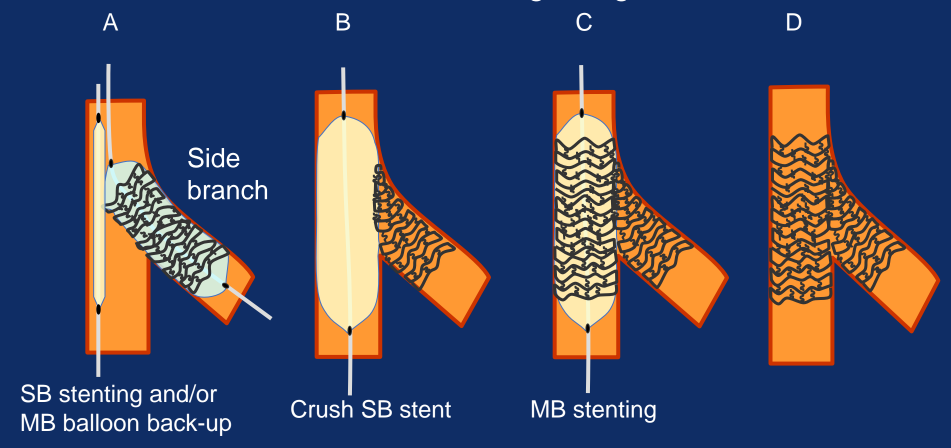
Crush Technique

D. Final result





Performed with 6~7Fr guiding catheter



Advantages

Minimizes multi-layers of struts Good scaffolding at SB ostium Facilitates FKI Compatible with 6-Fr guider

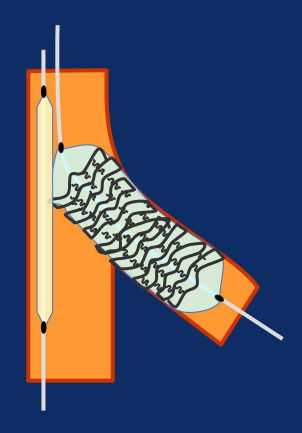
Disadvantages

Still leaves multiple layers of strut



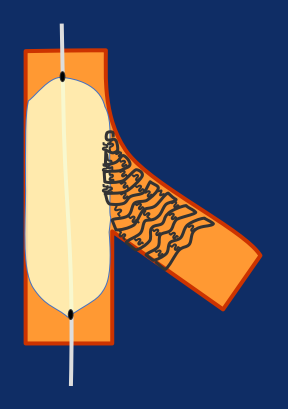
Performed with 6~7Fr guiding catheter

A. Deploy the SB stent \pm MB balloon backup



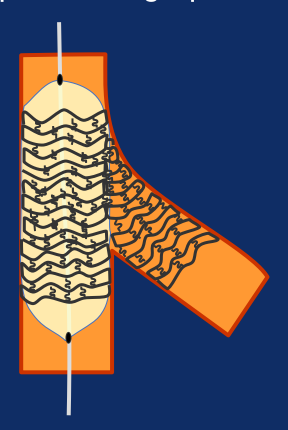
Performed with 6~7Fr guiding catheter

B. Crush SB stent



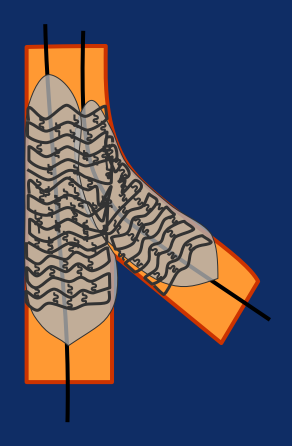
Performed with 6~7Fr guiding catheter

C. Deploy stent in MB, then rewire SB and perform high-pressure dilatation



Performed with 6~7Fr guiding catheter

E. Perform final kissing inflation



Performed with 6~7Fr guiding catheter

F. Final result





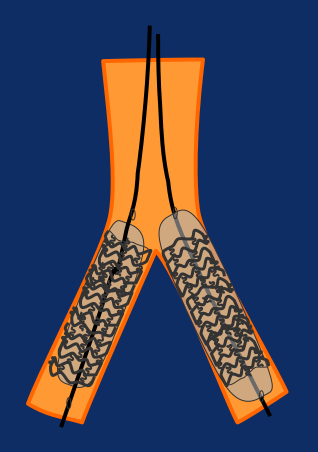
- Bifurcation without stenosis proximal to the bifurcation
- Short LM
- Less angle



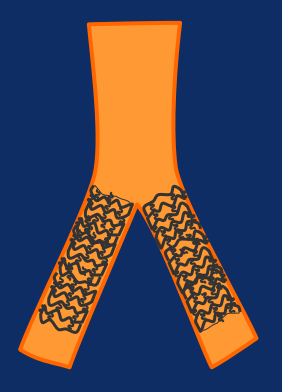
A. Position 2 parallel stents covering both branches with a slight protrusion into the proximal MB



B. Deploy 2 stents individually (or simultaneously)



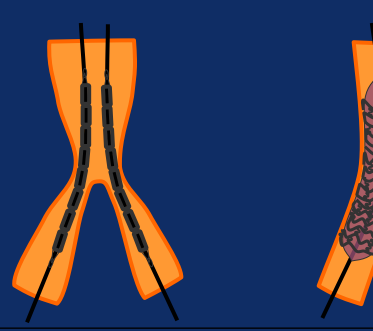
C. Perform high-pressure sequential single stent postdilation, Then medium pressure final kissing inflation





- Large proximal reference
- Bifurcation with stenosis proximal to the bifurcation

A B C





Advantages

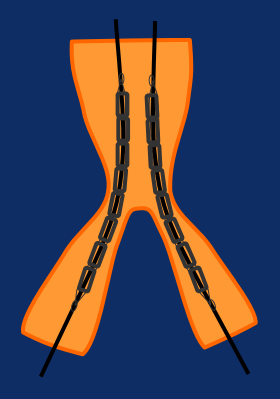
No risk of occlusion for both branches No need to re-cross any stent Technically easy and quick

Disadvantages

Requires 7- or 8-Fr guider
Leaves long metallic carina
Over-dilatation in proximal MB
Diaphragmatic membrane formation
Difficulty in repeat revascularization

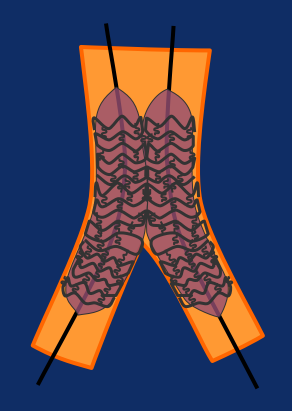


A. Position 2 parallel stents covering both branches with a long double barrel protrusion into the proximal MB



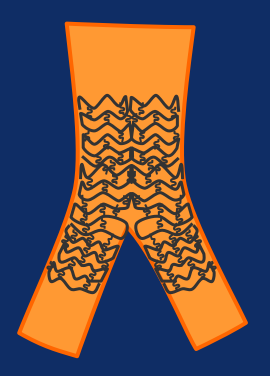


B. Deploy 2 stents





C. Perform final kissing inflation resulting a new metallic carina

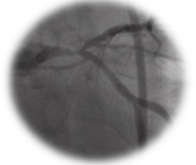


IVUS in LM disease



IVUS Use was Associated with Better 10-yr Outcomes after LM PCI **MAIN-COMPARE** Registry

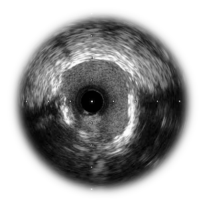
Left Main Disease

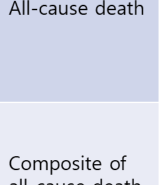


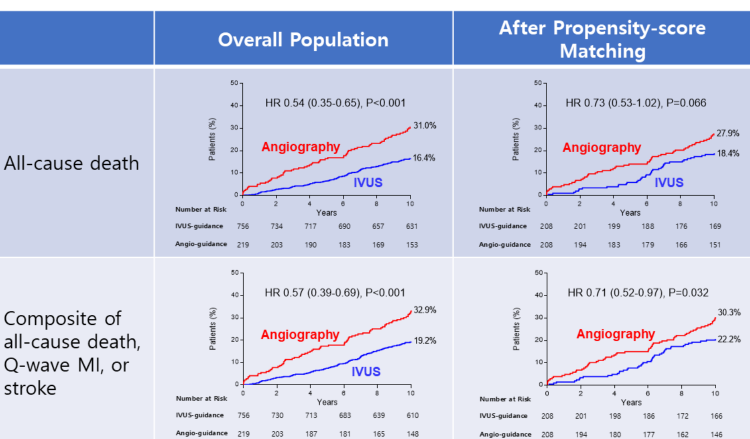
Follow-up

10-Year

IVUS-guided PCI

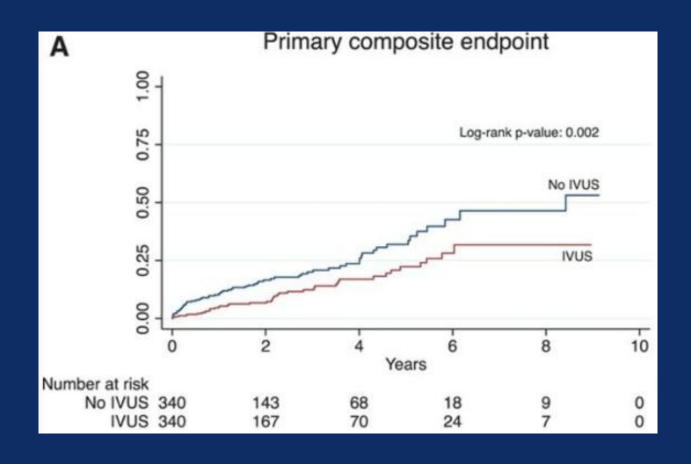






IVUS guidance associated with better outcome in LMCA stenting compared with angiography guidance alone

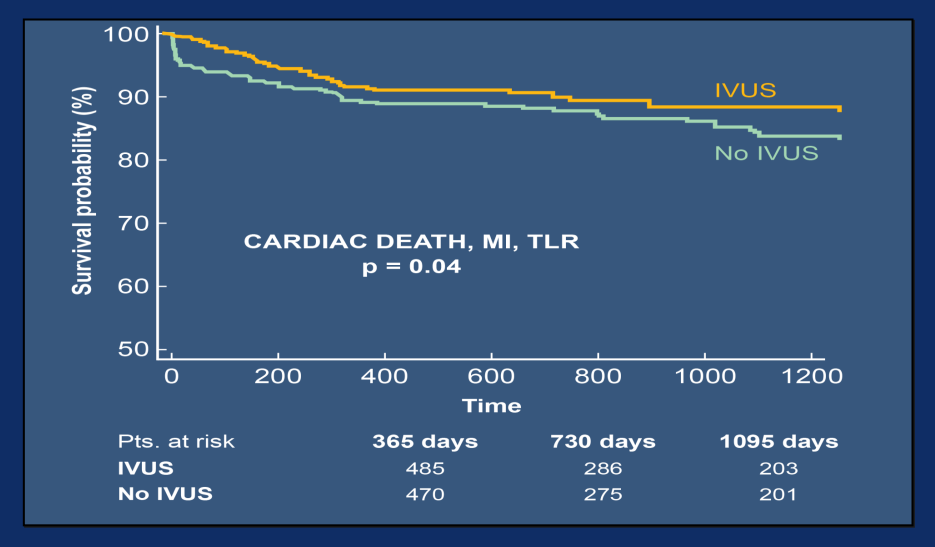
SCAAR Registry





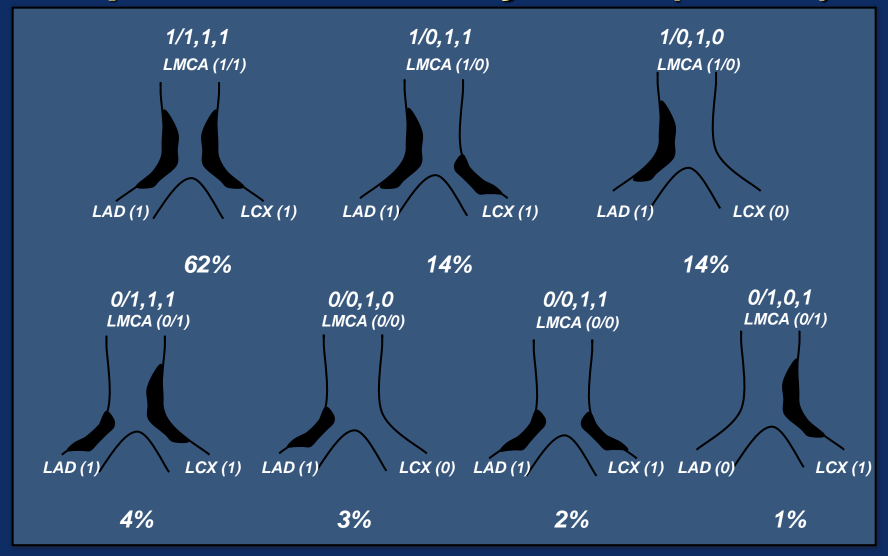
Pooled analysis :ESTROFA-LM, RENACIMIENTO, Bellvitge, Valdecilla

Effectiveness of IVUS on LM PCI





Plaque Distribution by IVUS (n=140)



In 90% plaque extends from LMCA-LAD



TCTAP2025

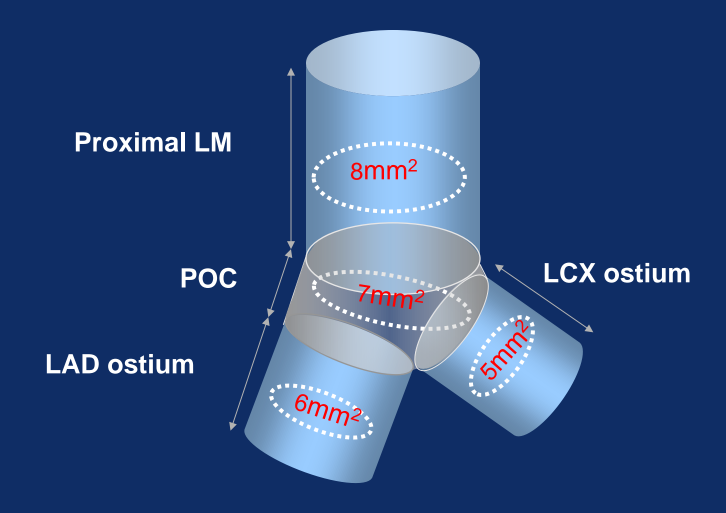
Plaque Distribution by IVUS (n=82)

DLM POC LAD LCX	N. (%)	LAD ostium, MLA (mm²)	POC, MLA (mm²)	DLM, MLA (mm²)	LCX ostium, MLA (mm²)
	5 (6%)	4.4±2.0	9.6±4.4	8.1±4.7	3.4±1.6
	26 (32%)	4.2±2.8	5.3±2.6	4.6±1.5	3.9±2.1
	12 (15%)	2.6±1.3	4.5±1.6	4.5±2.1	3.3±2.0
	9 (11%)	4.3±2.5	5.6±3.3	5.7±3.8	7.6±3.6
	9 (11%)	3.2±1.4	6.1±2.0	4.8±2.5	3.9±1.4
	4 (5%)	3.4±1.9	5.2±1.9	5.8±4.7	3.9±2.0
	4 (5%)	2.8±0.7	5.1±2.1	5.1±2.2	6.6±1.7
	5 (6%)	3.4±1.9	5.2±2.6	5.1±3.8	4.6±2.1

In all cases, the LM disease extended into LAD and LCX continuously.



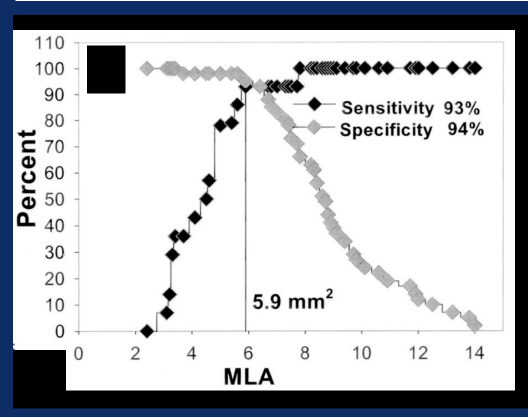
Optimal MSA on a segmental basis

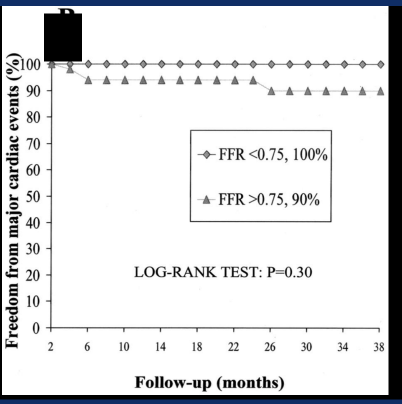




Cut-off for Predicting LM FFR<0.75 : LM MLA 6.0mm²

- Sum of lumen areas of two daughter vessels (Each of LAD and LCx should be 4.0mm²) = 150% of the parent LM
- Murray's Law $(LM r^3 = LAD r^3 + LCx r^3)$





Geometric Abstraction

Old MLA cut-off 6.0mm² was obtained from *Murray's law* considering an MLA 4.0mm² as ischemic threshold of both LAD and LCX

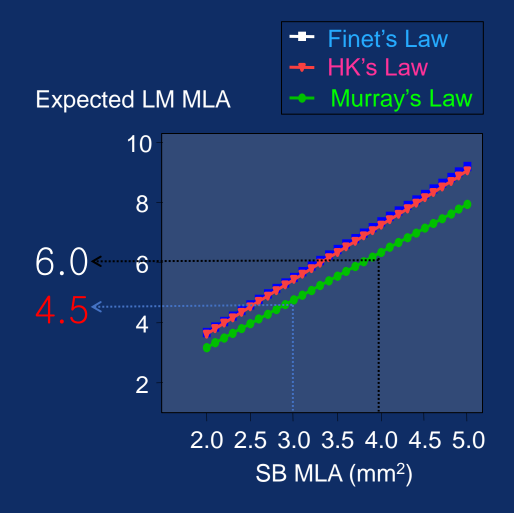


LAD	LCX	LM (Murray's)	
4.0	4.0	6.35	
4.0	3.9	6.27	
4.0	3.8	6.19	
4.0	3.7	6.11	
4.0	3.6	6.04	
4.0	3.5	5.96	



False Assumption... The used cut-off 4.0mm² is too Big!

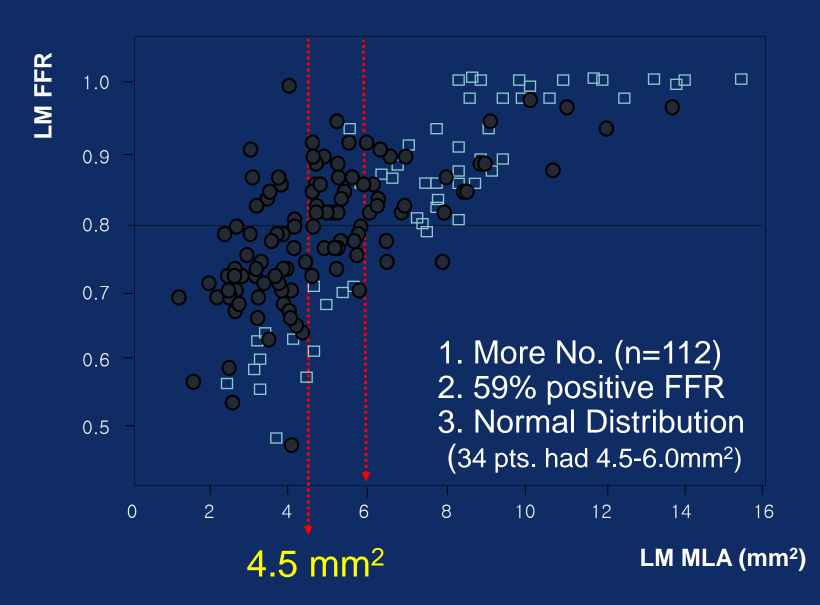
LAD	LCX	LM (Murray's)
3.0	3.0	4.76
3.0	2.9	4.68
3.0	2.8	4.60
3.0	2.7	4.53
3.0	2.6	4.45
3.0	2.5	4.37





TCTAP2025

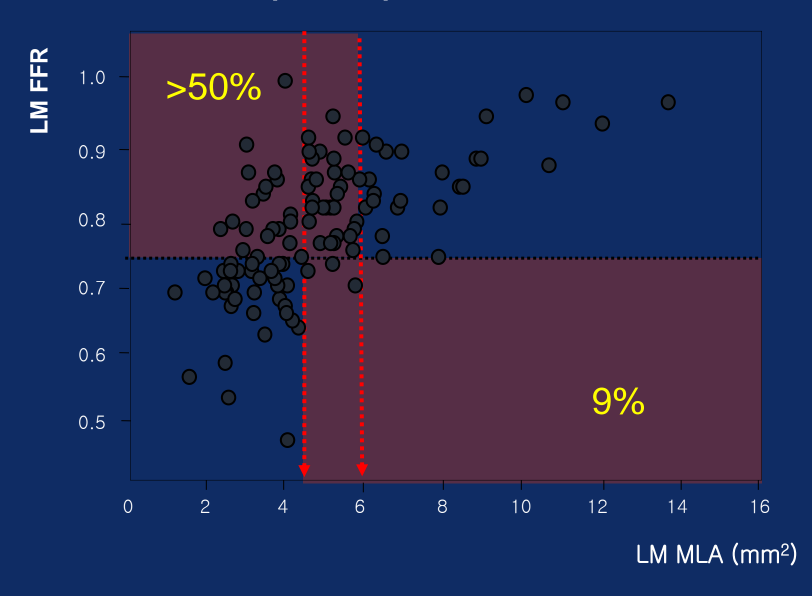
AMC Data (n=112)





TCTAP2025

AMC Data (n=112)





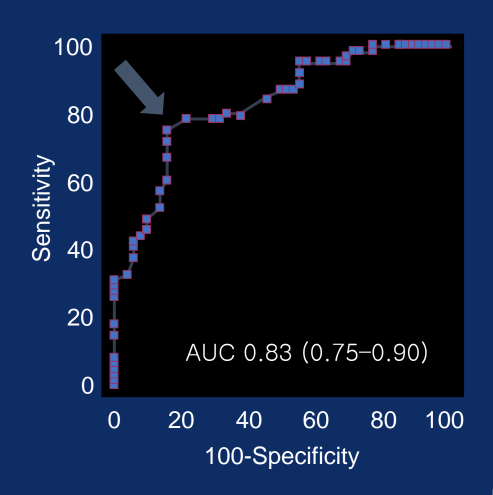
- Old data (MLA 6.0mm²) included downstream SB disease, and 32 of 55 (58%) were distal LM lesions that usually extend to the SB ostia
- Recent data (MLA 4.5mm²) evaluated only pure LM lesions, which more reliably assessed the impact of LM-MLA on functional significance

TABLE 1. Baseline Clinical, Angiographic, and IVUS Characteristics of Patients (n=55)			
Age, y	62±11		
Diabetes mellitus, n	20		
Hypertension, n	50		
Smoking, n	39		
Prior bypass surgery, n	13		
Ostial LM stenosis, n	20		
Mid-I M. stenosis, n	3		
Distal LM stenosis, n	32		



New LM MLA 4.5mm²

Matched with FFR < 0.80 Ostial and Shaft LM Disease (N=112)

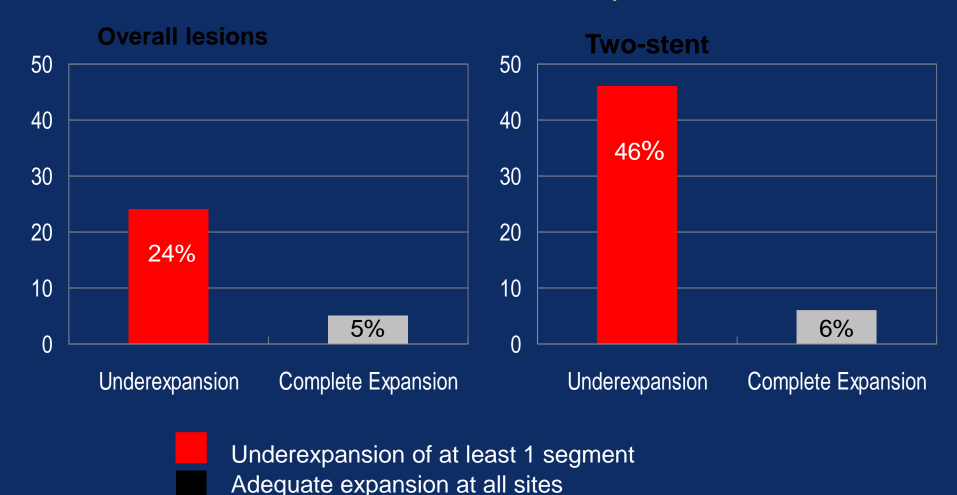


Sensitivity	79%
Specificity	80%
PPV	83%
NPV	76%



Frequency of ISR in LM Lesions

with vs. without Underexpansion



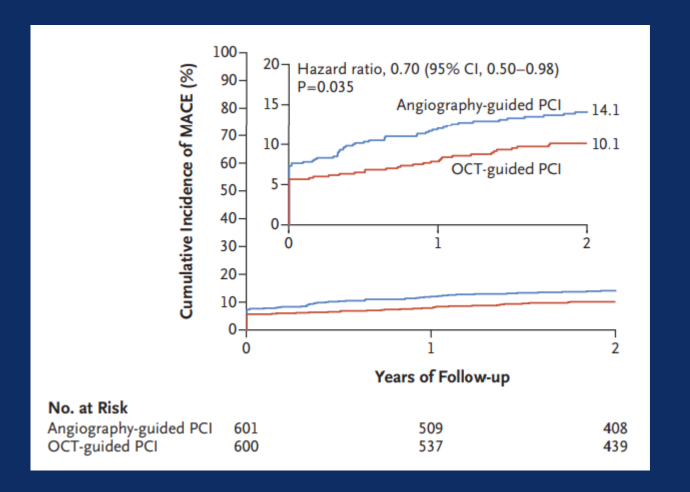




OCT in LM disease



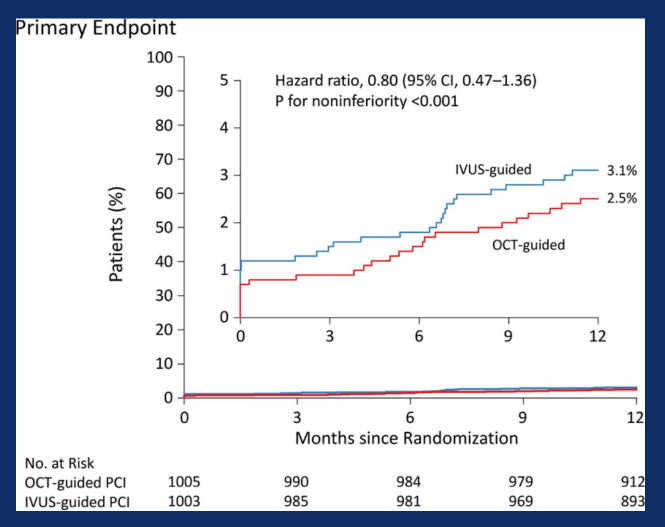
OCT-guidance associated with better outcome in LMCA stenting compared with angiography guidance alone

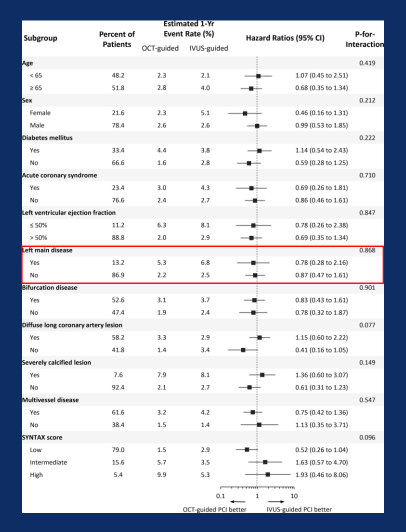


Subgroup	OCT-Guided PCI (N=600)	Angiography-Guided PCI (N=601)	Hazard Ratio	(95% CI)
	no. of events/total no. of patients (%)			
All patients	59/600 (10)	83/601 (14)	-	0.70 (0.50-0.98)
Sex			i	
Female	9/127 (8)	18/126 (15)	→ ; ;	0.49 (0.22-1.08)
Male	50/473 (11)	65/475 (14)		0.76 (0.53-1.10)
Age			i	
<65 yr	29/249 (12)	39/248 (16)		0.73 (0.45-1.18)
≥65 yr	30/351 (9)	44/353 (13)	→	0.67 (0.42-1.07)
Diabetes mellitus				
Yes	10/103 (10)	16/97 (17)		0.55 (0.25-1.20)
No	48/490 (10)	66/497 (14)		0.73 (0.5-1.06)
Left main coronary artery as trial bifu	rcation			
Yes	15/111 (14)	20/116 (19)		0.78 (0.40-1.51)
No	44/489 (9)	63/485 (13)	→	0.68 (0.46-1.00)
Stent technique				
One-stent	12/209 (6)	26/219 (12)	→ ;	0.47 (0.24-0.93)
Two-stent	47/388 (13)	57/382 (15)	-!+ -	0.80 (0.55-1.18)
Multivessel			i	
Yes	12/106 (12)	22/125 (18)		0.63 (0.31-1.28)
No	47/494 (10)	61/476 (13)	→	0.73 (0.50-1.07)
Acute coronary syndrome or staged P from recent AMI	PCI			
Yes	31/270 (12)	39/280 (14)	-i• -	0.81 (0.51-1.30)
No	28/330 (9)	44/321 (14)		0.61 (0.38-0.98)
Calcified lesion			i	
None-to-minor	35/402 (9)	54/405 (14)		0.64 (0.42-0.98)
Moderate-to-severe	24/198 (13)	29/194 (15)	- i+ -	0.81 (0.47-1.39)
SB lesion length >5 mm by QCA				
Yes	40/425 (10)	63/413 (16)	→ i	0.60 (0.40-0.89)
No	19/159 (12)	18/169 (11)	-	1.13 (0.59-2.16)
SYNTAX score				
<17	17/219 (8)	22/221 (10)		0.77 (0.41-1.45)
17–21	15/189 (8)	27/181 (15)		0.52 (0.27-0.97)
>21	27/191 (14)	34/197 (18)	1.0	0.82 (0.49–1.35)
		OCT-G	uided PCI Better Angiography	Guided PCI Better



OCT-guidance is non-inferior in LMCA stenting compared with IVUS-guidance



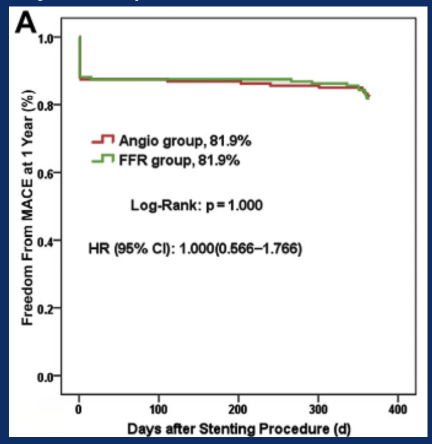


FFR in LM disease



FFR guided and angio guided provisional stenting of LM DKCRUSH-VI trial

primary endpoint : 1 yr composite of MACE

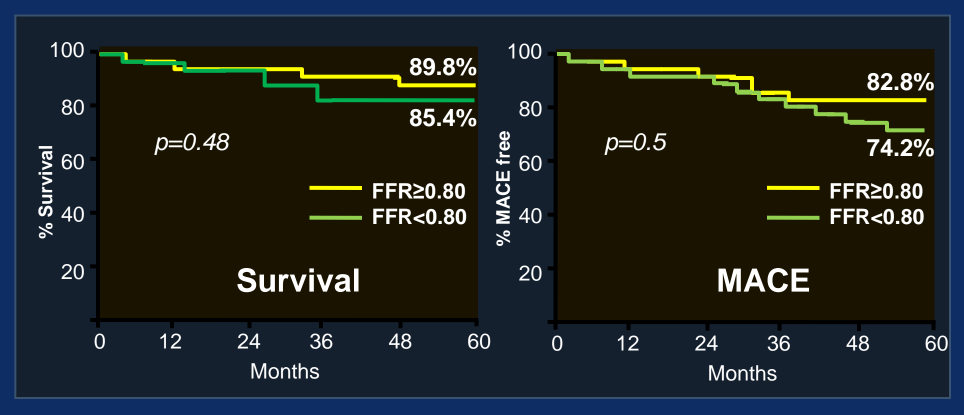


Angiographic and FFR guidance of provisional SB stenting of LM bifurcation lesions provided similar 1-year clinical outcome.

J Am Coll Cardiol Intv 2015:8:536–46

FFR guided PCI in Equivocal LMCA

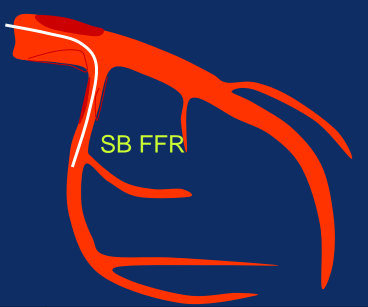
- In 213 patients with an equivocal LMCA stenosis
- FFR ≥0.80: Medication (n=138) vs. FFR<0.80: CABG (n=75)



An FFR-guided strategy showed the favorable outcome.



Use of IVUS vs. FFR in SB Assessment After LM Cross-over

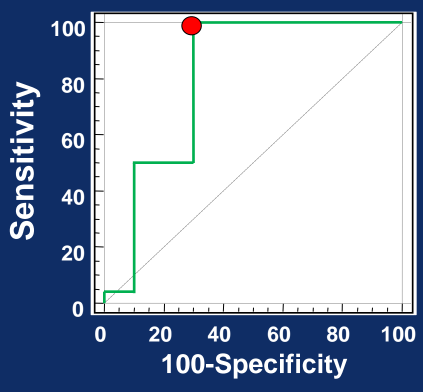


	SB-pullback IVUS	SB FFR
Advantage	 Confirm the anatomical compromise and MLA loss Mechanism of SB jailing 	 Confirm the functional SB compromise
Pitfalls	MLA-FFR mismatchNo MLA criteriaLow feasibility	• Minority - not feasible

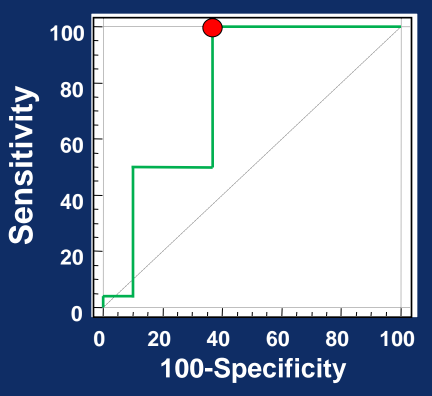


Functional Compromise of LCX after LM Cross-Over Stenting

Preporcedural MLA and plaque burden of poststenting LCX FFR < 0.80



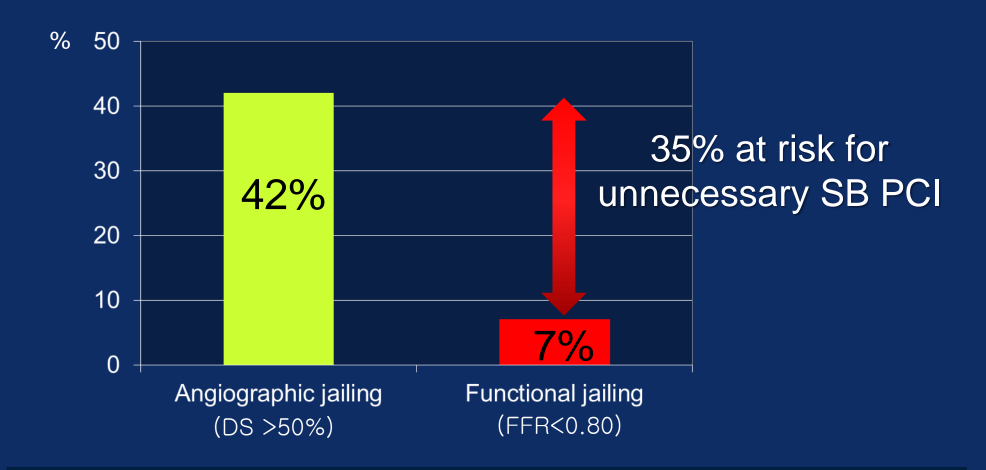
MLA 3.7 mm²



Plaque burden 56%



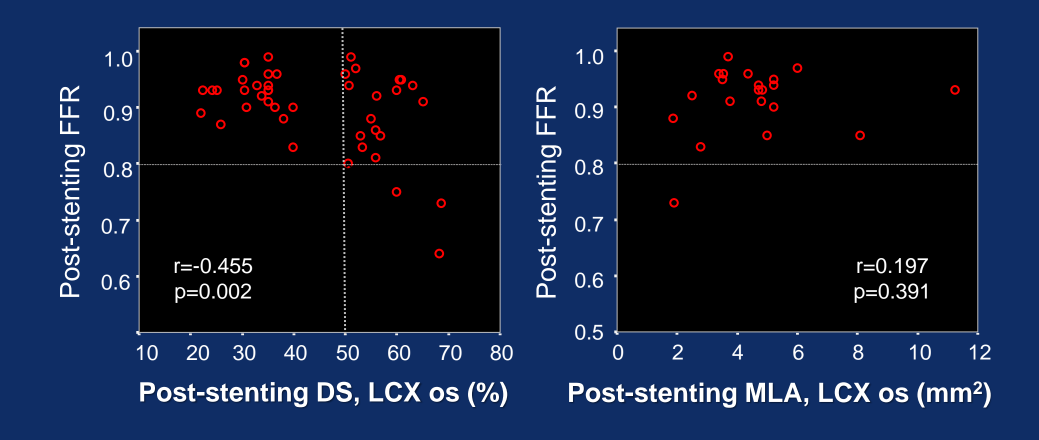
Functional LCX Compromise In LMCA Bifurcations (LCX ostial DS<50%)



When Pre-PCI LCX Ostial DS<50%, Just Do Single Stent!



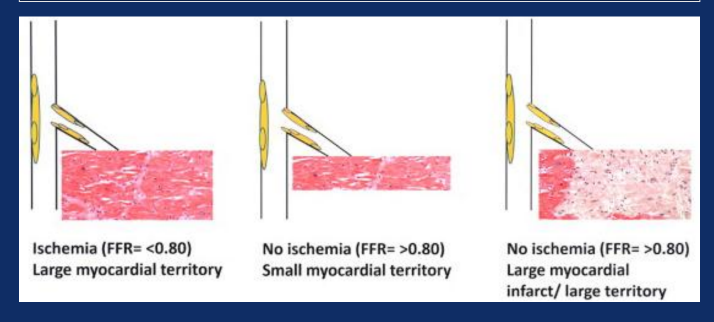
LMCA Bifurcation Post-stenting LCX Stenosis





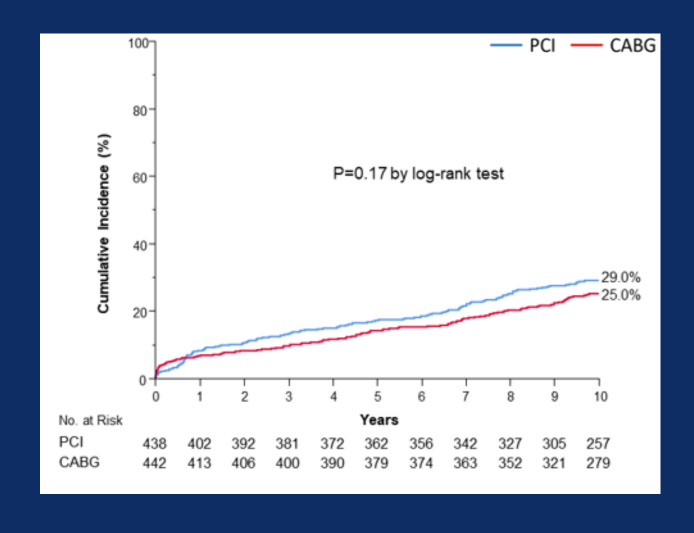
Why Mismatch?

- Lesion eccentricity of SB
- Negative remodeling of ostium
- Various size of myocardium
- Strut artifacts
- Focal carina shift



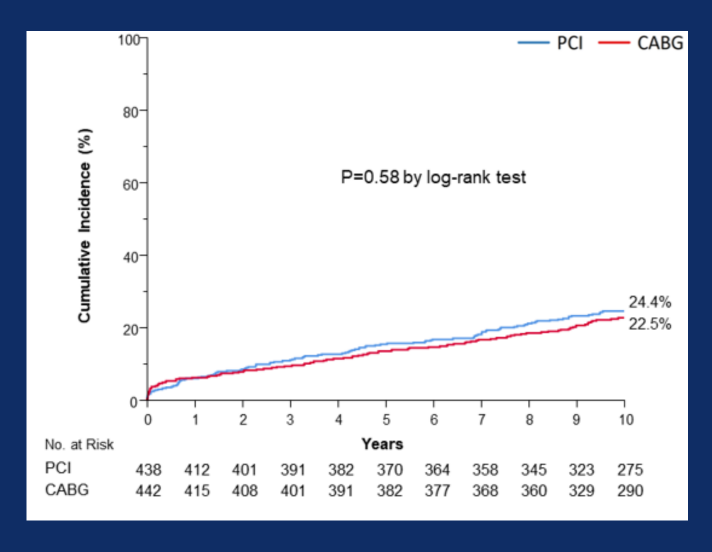


Extended Follow-Up of the **BEST** trial: Primary Composite Endpoint



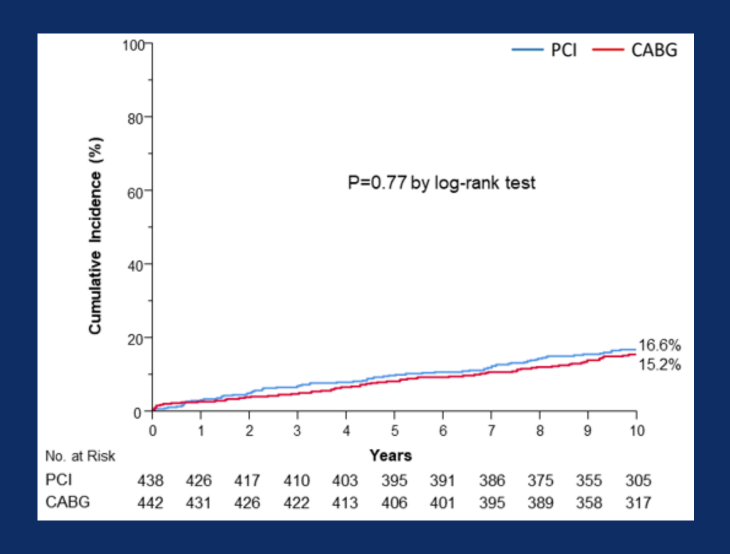


Extended Follow-Up of the **BEST** trial: Death, Stroke, or MI



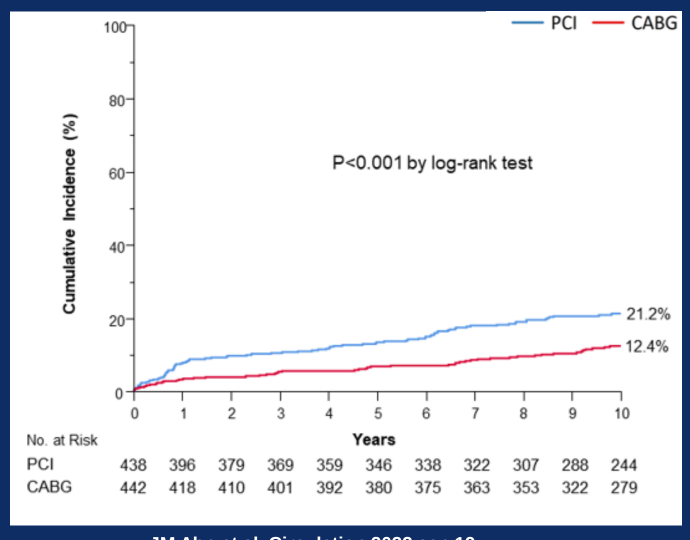


Extended Follow-Up of the **BEST** trial: All-cause death



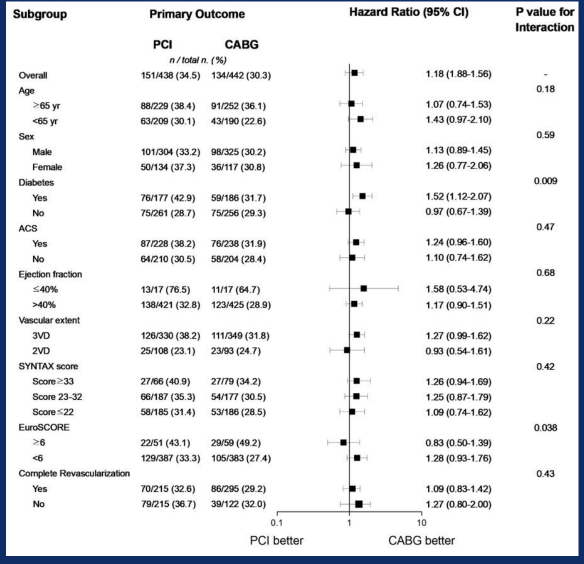


Extended Follow-Up of the **BEST** trial: Repeat Revascularization





Extended Follow-Up of the **BEST** trial: Repeat Revascularization





Fractional Flow Reserve versus Angiography for Treatment-Decision and Evaluation of Significant Left MAIN Coronary Artery Disease

FATE-MAIN Trial

934 Patients with Significant (Angiographic Diameter Stenosis ≥50%) Left Main Coronary Artery Disease Who Were Eligible for PCI

1:1 randomization stratified by (1) participating sites and (2) the presence of concomitant non-left main PCI

FFR-Guided Left Main PCI (N = 467)

Angiography-Guided Left Main PCI (N = 467)

The primary end point was the composite of death from any cause, myocardial infarction, hospitalization for unstable angina, heart failure, or resuscitated cardiac arrest, or repeat revascularization at 2 year.

ISR



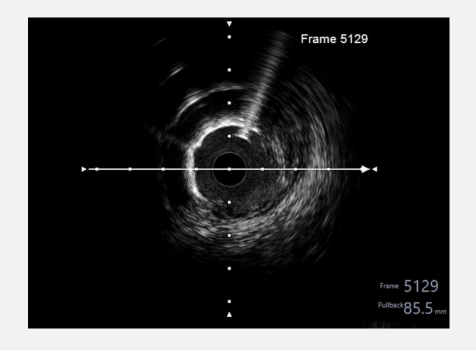


Indications and Intended Use

The WOLVERINE™ Cutting Balloon Device is indicated for use in patients with coronary vessel disease who are acceptable candidates for coronary artery bypass graft surgery, should it be urgently needed, for the purpose of improving myocardial perfusion.

In addition, the target lesion should possess the following characteristics:

- Discrete (< 15 mm in length), or tubular (10 mm to 20 mm in length)
- Reference vessel diameter (RVD) of 2.00 mm to 4.00 mm
- Readily accessible to the device
- Light to moderate tortuosity of proximal vessel segment
- Nonangulated lesion segment (< 45°)
- Smooth angiographic contour
- Absence of angiographically visible thrombus



WOLVERINE™ FDA US IFU Updates November 2021

Scientific

INTENDED USE/INDICATIONS FOR USE

The Wolverine Cutting Balloon Device is indicated for dilatation of stenoses in coronary arteries for the purpose of improving myocardial perfusion in those circumstances where a high pressure balloon resistant lesion is encountered. In addition, the target lesion should possess the following characteristics:

- Discrete (< 15 mm in length), or tubular (10 mm to 20 mm in length)
- . Reference vessel diameter (RVD) of 2.00 mm to 4.00 mm
- Readily accessible to the device
- · Light to moderate tortuosity of proximal vessel segment
- Nonangulated lesion segment (< 45°)
- · Smooth angiographic contour
- Absence of angiographically visible thrombus and/or calcification

Changes

- Removed "and/or calcification" in target lesion characteristics bullet points
- Emergency surgical backup now a clinical consideration
- Additional cleanup and formatting for clarity

Rationale

- Align Instruction for Use with modern product usage
 - Cutting Balloon was first introduced before stents were approved for coronary use
 - · Modern use of cutting balloon has since changed
- Supported by extensive literature, clinical data and real-world experience
- FDA approved changes in Nov 2021

Product Design

Traditional balloon angioplasty can result in complications like:

VESSEL DISSECTION

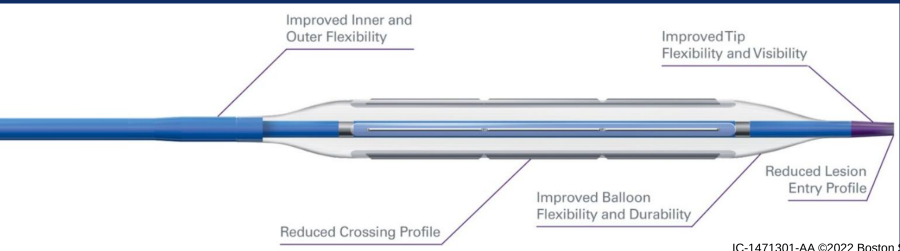
POOR LUMINAL GAIN

LESION RECOIL

BALLOON SLIPPAGE POOR STENT APPOSITION

The WOLVERINE™ Advantage

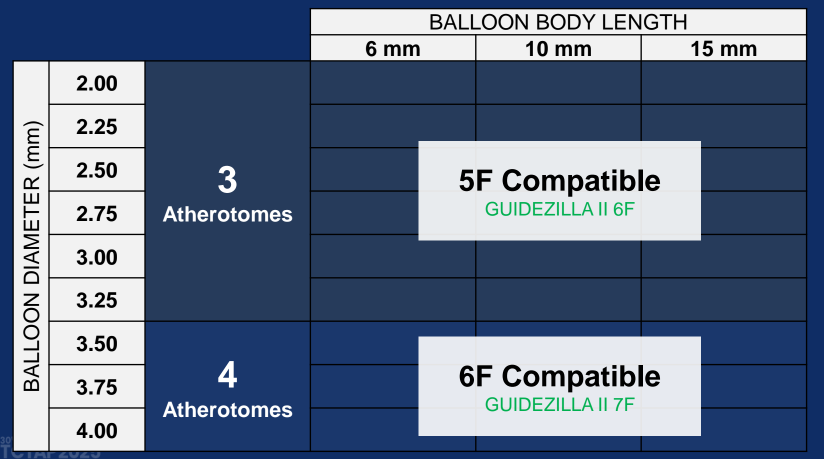
The unique design of the WOLVERINE Cutting Balloon is designed with **proprietary atherotomes** on a **low pressure non-compliant balloon** to directly address each of these complications





Balloon Matrix and Inflation Pressures

Monorail Balloon Catheter with working lengths of 6, 10 and 15 mm For vessels with reference diameter of 2.0 – 4.0 mm



INFLATION PRESSURE RATING

Nominal = 6 ATM

Rated Burst = 12 ATM

Sizing Considerations

WOLVERINE™ utilizes the NC EMERGE™ Catheter Platform, yet the balloon was designed to have a lower nominal pressure resulting in a different compliance



Growth Chart Example (3.0 mm)

Wolverine™ Coronary Cutting Balloon™

MONORAIL™

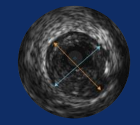
Microsurgical Dilatation Device

atm - kPa		3.00mm
Pressure		Balloon O.D.
3.0 - 304		2.88
4.0 - 405		2.94
5.0 - 507		2.99
6.0 - 608	NOMINAL	3.06
7.0 - 709		3.10
8.0 - 811		3.15
9.0 - 912		3.18
10.0 - 1013		3.22
11.0 - 1115		3.25
12.0 - 1216	RATED*	3.28 -
*Rated Burst	Pressure. DO	NOT EXCEE

Sizing Considerations:

WOLVERINE grows roughly a quarter size when going from nominal (6 ATM) to rated burst pressure (12 ATM)

Physician consensus is to measure the normal distal reference with IVUS and then downsize WOLVERINE a half size from that measurement



Oversizing at nominal pressure will cause atherotomes to be "pillowed" by the balloon and may not provide adequate forces to modify calcium



Oversizing Example Blue = Balloon Red = Vessel

Oversizing at rated burst pressure may lead to vessel stretching and trauma due to balloon growth (not atherotomes)



Device Preparation and Use Instructions

Device Preparation

Important: WOLVERINE™ preparation uses a wet negative prep procedure. Customary balloon preparation methods do not apply!

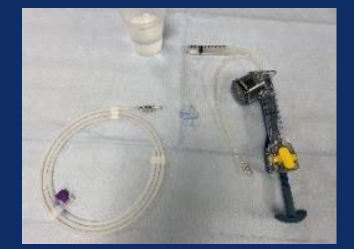
Sizing
• The

 The Wolverine IFU states that the inflated diameter of the device should approximate a ratio of 1.1:1 in relation to the average diameter of the reference vessel. Oversizing increases risk of perforation. As stated earlier, sizing a quarter to half size down may be needed if using higher inflation pressures.

Unpacking

• Using ste

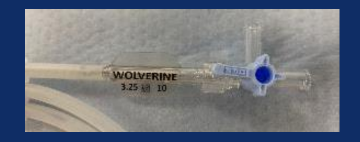
- Using sterile technique, remove the device in its protective hoop from its package and place onto a sterile field.
 - Do not remove the device from its protective hoop.
 - Do not remove the balloon protector from the device tip.



3

Attach Stopcock & Prepare Inflation Device

- Connect a three-way stopcock to the balloon port.
 - Turn stopcock lever OFF to the balloon.
 - Prepare an inflation device with 5 cc of contrast solution (mixture must be at least 50:50 contrast medium and sterile saline).



Device Preparation



Attach Inflation Device & Purge

- Attach the inflation device to stopcock.
 - Assure luer connections are properly aligned to avoid stripping the luer thread causing subsequent leakage and use care when connecting the device to avoid damage (e.g., shaft kink).
 - Purge stopcock by flushing 1-2 cc of contrast medium through the middle port.







Pull Full Negative

• Turn the stopcock lever towards the middle port or open to the balloon and immediately withdraw inflation device plunger to full negative and place the inflation device in a locked position. This will maintain a constant vacuum on the device.







Remove Device from Hoop

• When the device is ready to be inserted into the body, remove the device from its protective hoop. Use care when removing the device to avoid damage (e.g., shaft kink).

Device Preparation

7

Remove Balloon Protector

- Using straight force (not a twisting motion), pull the balloon protector distally from the device tip. For WOLVERINE MR Cutting Balloon Devices, remove the mandrel distally after removing the balloon protector.
 - Caution: If unusual resistance is felt during removal of the balloon protector or mandrel, do not use the device and replace with another.



8

Coiling & Securing with CLIPIT Clip

- The WOLVERINE MR Cutting Balloon Device may be coiled once and secured using the CLIPIT Clip provided in the device package.
 - Only the proximal shaft should be inserted into the CLIPIT Clip; the clip is not intended for the distal end of the device.
 - Remove the CLIPIT Clip prior to inserting the device into the patient's body.

9

Flush Guidewire Lumen

• Flush the guidewire lumen of the device with heparinized saline. For WOLVERINE MR Cutting Balloon Device flush through the distal tip of the device.



Sterility

• Maintain device on a sterile table until ready to use.

Inflation & Removal Instructions

Inflation

1

Go Slow

- Under fluoroscopy, slowly inflate the device (1 ATM/5 sec) to 6 ATM (nominal size).
 - Do not inflate the device above 12 ATM (rated burst pressure).
 - If difficulty is experienced during balloon inflation, do not continue inflation; deflate and remove the device.

2

Treat Distal then Proximal

• When using the device on long lesion segments, treat distal portion first and then proximal lesion segment second. Repeat coronary arteriography after each use to evaluate results.

Tips and Tricks

Prior to advancing the catheter, it may help to increase pressure to 1 atm and then pull negative to aid in loosening the packaged balloon crimp and provide added flexibility

Removal



Deflate & Pull Negative

- Deflate the device by dialing down on the inflation/deflation device, then pull a negative vacuum. Maintain vacuum on the device and verify full deflation under fluoroscopy.
- 2

Confirm Successful Result

· Repeat coronary arteriography to confirm successful result.



Withdraw

• Withdraw the device into the guiding catheter. While withdrawing the deflated device and guidewire from the guide catheter through the hemostasis valve, tighten the hemostasis valve.

Tips and Tricks

Deflating slowly by dialing down pressure methodically to optimize balloon re-wrap



Clinical Use Scenarios

WOLVERINE™ The right tool for vessel preparation device

Proper Solution to Help Prepare Lesions Prior to Stenting

WOLVERINE is right tool at helping treat a wide range of lesions:

- Cuts fibrotic plaque to limit recoil
- Cracks thin concentric and eccentric calcium
- Prepare small vessels prior to Drug Coated Balloon
- Address In-Stent Restenosis
- Limit balloon slippage in coronary ostium and bifurcation lesions
- Cutting balloon angioplasty device designed with improved crossability and deliverability, to deliver precise and controlled cutting action

Clinical Use Scenarios

Small Vessel Lesions

REDUCE RESTENOSIS

• High rates of restenosis

- Tendency to dissect
- Abrupt closure¹

Ostial and Bifurcation

LesionsPLAQUE SHIFT

Recoil

- Plaque Shift
- Side Branch Compromise

Fibrotic Lesions

CHANGE LESION COMPLIANCE

- High concentration of elastin and muscle fibers
- High risk of vessel recoil

Calcified Lesions

CRACK CALCIUM TO ALLOW EXPANSION

- Calcium deposits in plaque that prevent lumen gain
- Varying degrees of burden and arcs

- Use as stand-alone therapy
- DCB or Stent?

- Dilates while reducing elastic recoil²
- More plaque compression
- · Minimal plaque shift
- Less vessel stretching³

- Atherotomes score through fibrotic plaque⁴
- Reduce hoop strain and limit recoil
- Lumen Gain

- Use as stand-alone therapy in eccentric and thin concentric calcium
- Possible additive therapy with atherectomy
- Lumen Gain

CUTTING BALLOON OBJECTIVES

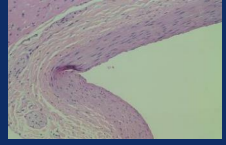
WOLVERINE™ Mechanism of Action

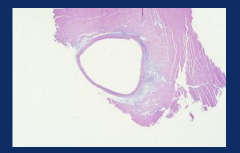
Porcine Artery Models

ACUTE

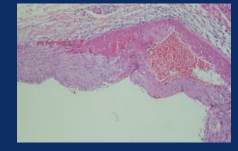
14-DAY





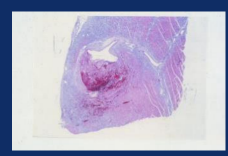






Acutely there is over stretch* and visible trauma to the vessel wall with POBA

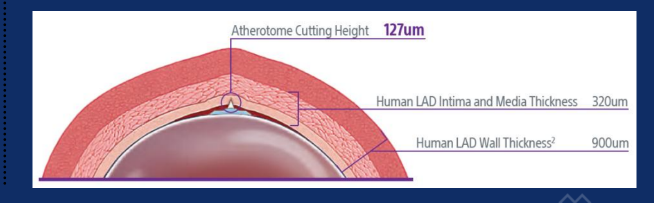
*This level of over-stretch was done for investigational purposes only



At 14-days the vessel has recoiled with POBA and stayed open with cutting balloon

Reliable Option

- 25+ Year Track Record: WOLVERINE has been used for over 25 years, and has a long track record of safety with real-world patients and clinical trials
- Atherotome Height: Approximately the same height as 1st generation stents or a human hair
- Penetration Depth: Even when placed in healthy tissue, WOLVERINE's atherotomes typically only penetrate partially into the media





Calcium Modification

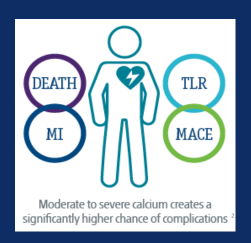
Calcium Needs to be Properly Treated

Calcium is a growing problem that can negatively impact PCIs if left untreated

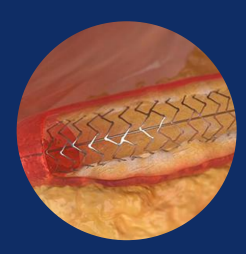
Calcium is prevalent in patients undergoing PCI



Calcium leads to worse clinical outcomes



Calcium can inhibit optimal stenting

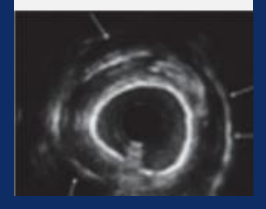


Calcium Morphology

CONCENTRIC



360°Calcium Arc Smooth Surface



ECCENTRIC



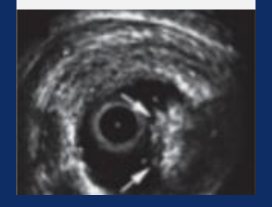
180 – 270° Calcium Arc Irregular Surface



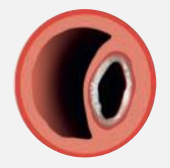
NODULE



90 – 180° Calcium Arc Luminal protrusion and irregular leading edge



PSEUDO-NODULE



Extra-plaque during CTO-PCI



The Right Tools Make a Difference



Controlled Mechanism of Action

Atherotomes anchor to calcium and produce controlled, longitudinal fractures

Strategic Atherotome Placement

Enables up to 4 points of contact with calcium, improving the probability of modification with a single balloon

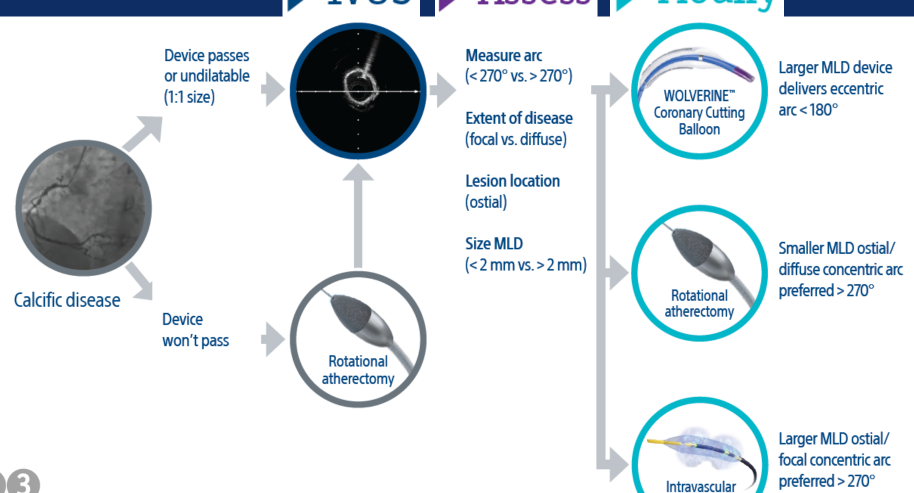
Focused Force to Amplify Impact

Pressure at atherotomes amplified to precisely fracture calcium at lower balloon inflation pressures

Calcific Lesion Modification Strategy



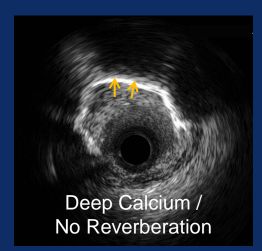
lithotripsy

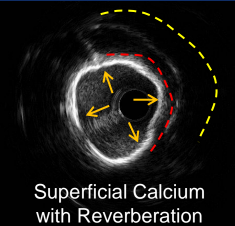




Assess Calcium FIRST with IVUS

Thickness





Angle



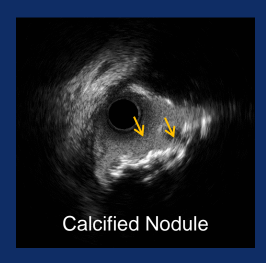
Reverberation
Surface of calcium

Length



LONGITUDINAL VIEW

Nodule





Proven Mechanism of Action

Effective. Safe. Versatile.

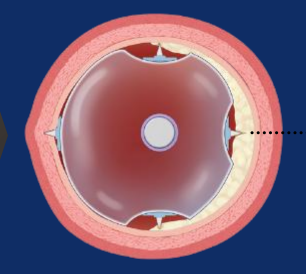
Wolverine's innovative design safely and efficiently cracks calcium³

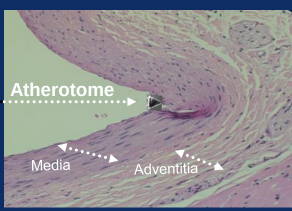
✓ Atherotome Amplified Force.¹

The atherotomes anchor into the plaque and amplify pressures generated by the balloon. This creates controlled, longitudinal cracks in the calcium.¹

Safely Cracks Calcium.

Due to its unique design, Wolverine can modify calcium at lower pressures than POBA.³ Atherotomes penetrate a small distance into the vessel wall, even in healthy tissue.⁴





Pre-clinical Swine Coronary artery post Cutting Balloon

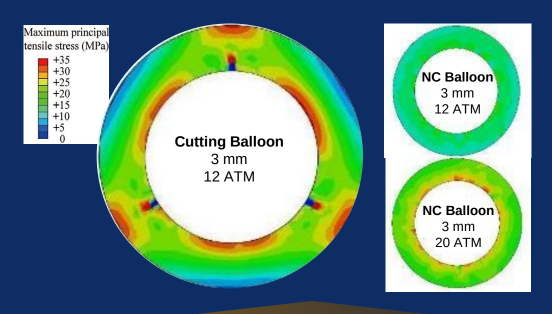
Atherotome Cutting Height	127 µm
Human LAD Media Thickness ²	320 μm
Human LAD Wall Thickness ²	900 μm

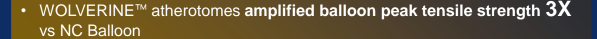
¹ Xiaodong Zhu et al.;Circ Rep 2021; 3: 1 – 8 doi: 10.1253/circrep.CR-20-0070. Results of computer models are not predictive of clinical performance. Clinical results may vary.

³ Mangleri, A. Cutting Balloon to Optimize Predilatation for Stent Implantation: The COPS Randomized Trial, TCT 2022

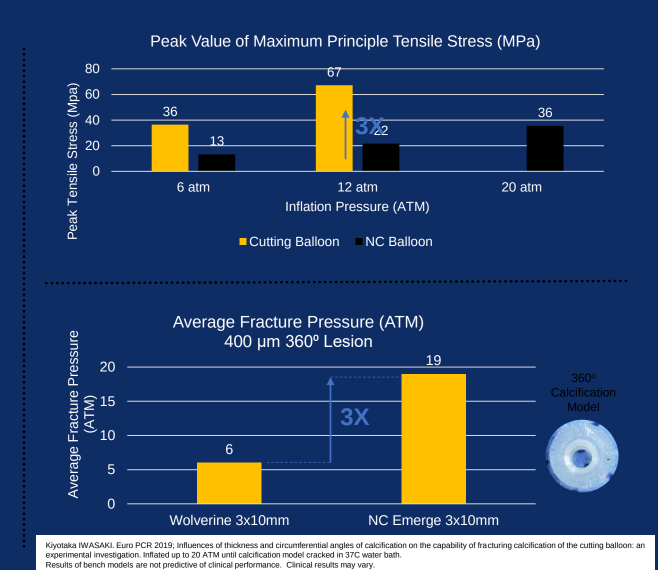
Treating Calcium with WOLVERINE™

Calcification Model Stress Distributions





- Force is focused at atherotomes for controlled even calcium cracking
- Balloon dilation force is enhanced between the anchored atherotomes

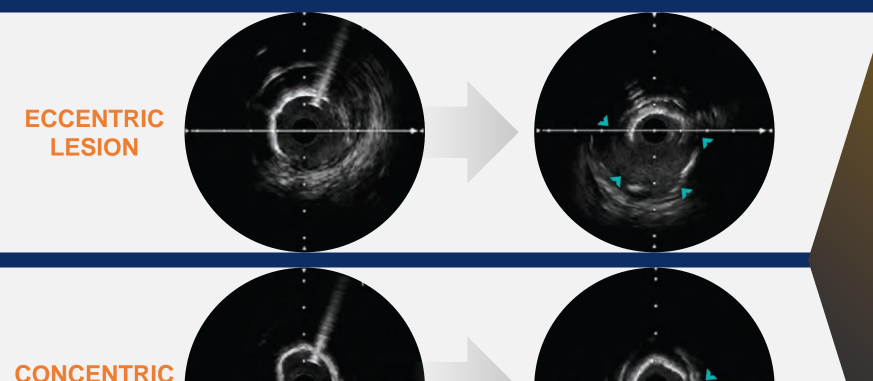


©2022 Boston Scientific Corporation or its affiliates. All rights reserved.

Demonstrated Efficacy in both Concentric and Eccentric Calcium

BEFORE

AFTER



WOLVERINE™ has
clinically demonstrated
effectiveness in calcium
ranging from 0° to 360°
with a proven mechanism
of action.¹

LESION

WOLVERINE™ Cracking Power in Action!



The COPS Trial Cutting balloon to Optimize Predilatation for Stenting



Primary Investigators

Dr. Antonio Mangieri, Dr. Antonio Columbo

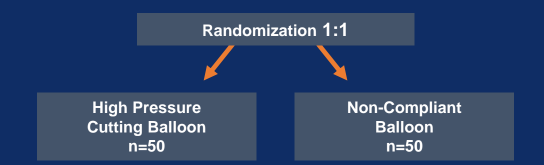
Three hospitals in Italy

Maria Cecilia Hospital, Humanitas Rozzano, Clinica Mediterranea

TAP2025

Study Design

 Prospective, randomized, multicenter open-label trial which enrolled 100 patients with significant calcified lesions evaluated at IVUS



Primary Endpoint

Minimal Stent Area (MSA) at Calcium Site

Secondary Endpoint

- Eccentricity Index : (LD max LD min) / LD max
- MSA
- Device Failure
- Safety: Procedural Complications & One-Year MACE

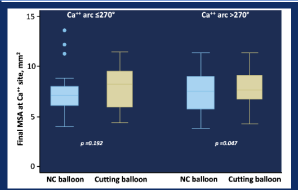
The COPS Trial: Results

Study contained a range calcium 100 – 360° and 29.4% avg of deep calcium

	Overall	CB (n=44)	NCB (n=43)	P value
Lesion Type				
Type B1	25 (28.7)	14 (32.5)	11 (25)	
Type B2/C	62 (71.2)	29 (67.4)	33 (75)	
Calcium distribution				0.482
Mixed Calcium	34 (40)	15 (34.8)	19 (45.2)	
Deep Calcium	25 (29.4)	15 (34.8)	10 (23.8)	
Superficial Calcium	26 (30.5)	13 (30.2)	13 (30.9)	
Arch of calcium (degrees)	266±84	274±84	258±85	0.373
Calcium length (mm)	12±6.6	11.9±7.3	12.5±6	0.667
Lesion length (mm)	24.3±9.7	23.5±9.6	25.1±9.8	0.442
Minimal lumen area (mm²)	3.2±0.9	3.4±1.1	3±0.7	0.02
QCA evaluation				
Reference vessel diameter (mm)	3.4±0.4	3.51±0.3	3.39±0.4	0.112
Percentage of stenosis (%)	81.2±8.1	79.4±7.6	82.7±8.3	0.97

WOLVERINE is clinically proven to provide superior MSA at the calcium site compared to POBA

	CB (n=44)	NCB (n=43)	P value
Final MSA (mm²)	7.1±1.7	6.5±2.1	0.116
Minimal Stent Diameter	2.7±0.4	2.5±0.4	0.064
Maximal Stent Diameter	3.2±0.4	3.1±0.4	0.189
Final MSA at calcium site	8.1±2	7.3±2.1	0.035
Minimal stent diameter at calcium site	2.9±0.7	2.7±0.4	0.016
Maximal stent diameter at calcium site	3.5±0.5	3.3±0.4	0.132
Eccentricity index at calcium site	0.84±0.7	0.8±0.8	0.013



The benefit was magnified in presence of severe calcifications

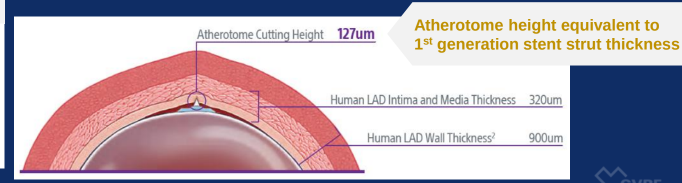
The COPS Trial: Safety

WOLVERINE™ use in calcium is safe, with no significant differences in procedural complications and 1-year MACE

	Overall	CB (n=44)	NCB (n=43)	P value
Device failure	3 (3.4)	3 (6.8)	0 (0)	0.517
Additional use of rotational atherectomy	1 (1.1)	1 (2.2)	0 (0)	0.79
Ellis type 1 vessel rupture	2 (2.2)	2 (4.4)	0 (0)	0.189
Implantation of a covered stent	1 (1.1)	1 (2.2)	0 (0)	0.65
Final TIMI flow >3	87 (100)	44 (100)	43 (100)	0.854
One year Follow-up				
Deaths	3 (3.4)	1 (1.1)	2 (4.6)	0.342
Cardiac deaths	1 (1.1)	0 (0)	1 (2.3)	0.887
Stroke	0 (0)	0 (0)	0 (0)	0.91
MI	0 (0)	0 (0)	0 (0)	0.96
TLR	3 (3.4)	1 (1.1)	2 (4.6)	0.49

WOLVERINE provided excellent procedural success with limited need for atherectomy (n=1) despite a high rate of severe calcium in the study

WOLVERINE is both a safe and effective option for modifying severely calcified lesions



The COPS Trial: Key Learnings



WOLVERINE™ resulted in a **significantly larger minimal stent area** at the calcified segment.



This difference was especially apparent in cases with **severe calcification**.



Stents had significantly more uniform expansion after vessel preparation with WOLVERINE.



WOLVERINE is safe for calcium treatment, even when inflated past rated burst pressure.



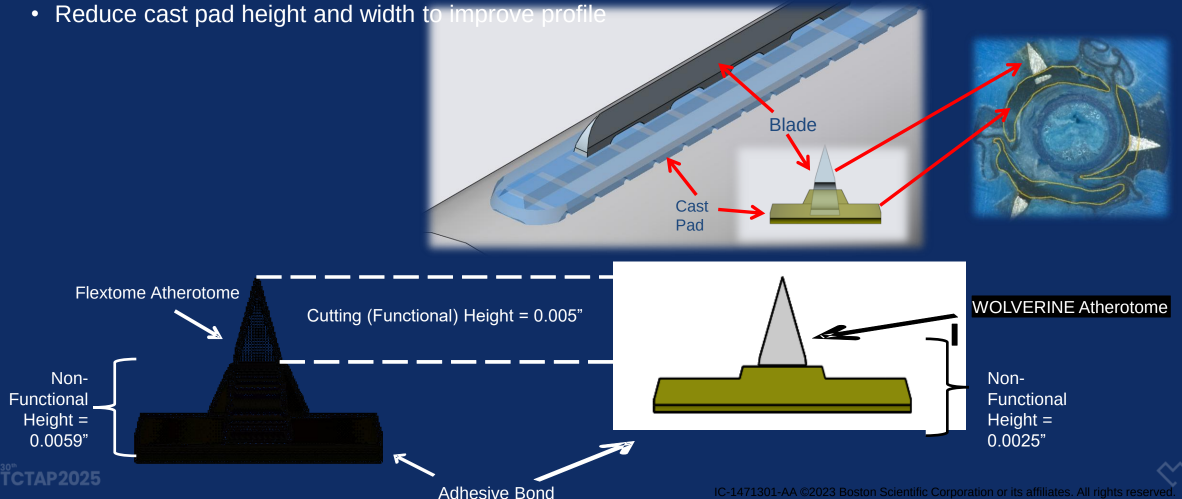
Competitive Product Comparisons

WOLVERINE vs FLEXTOME

	WOLVERINE™	FLEXTOME™	
Manufacturer	Boston Scientific	Boston Scientific	
Guide Cath Compatibility	5F, 6F	5F, 6F	
Size Matrix: Diameter (mm)	2, 2.25, 2.5, 2.75, 3, 3.25, 3.5, 3.75, 4	2, 2.25, 2.5, 2.75, 3, 3.25, 3.5, 3.75, 4	
Size Matrix: Length (mm)	6, 10, 15	6, 10, 15	
Pressures (ATM)	NOM: 6 RBP: 12	NOM: 6 RBP: 12	
Catheter Length (cm)	143	142	
Balloon Compliance	Non-Compliant	Non-Compliant	
Balloon Platform	NC EMERGE	NC Quantum MAVERICK	
Tip Entry Profile	<mark>0.017"</mark>	<mark>0.020"</mark>	
Proximal shaft Distal shaft	1.8Fr / 0.59mm 2.6Fr / 0.86mm	2.0Fr / 0.67mm 2.7Fr / 0.90mm	
Plaque Mod Method	3 or 4 evenly spaced atherotomes	3 or 4 evenly spaced atherotomes	

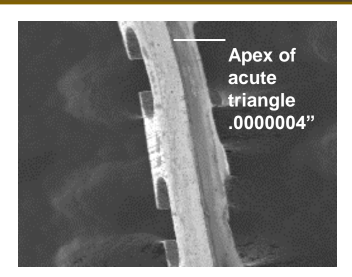
Atherotome Changes

• Reduce non-functional blade height (portion in the cast pad) to improve profile



The Atherotome Advantage

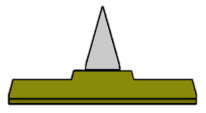
WOLVERINE™ Cutting Balloon™ Device Atherotome



WOLVERINE Atherotome Advantage:

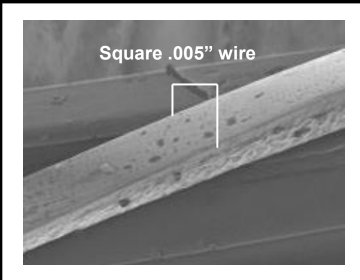
- Amplify balloon inflation pressures in calcium
- Create microsurgical incisions in fibrotic plaque

These two applications help to prepare vessels and limit recoil.



CUTTING BALLOON CROSS SECTION

Product A Nitinol Wire



Scoring Balloon Design:

- Flat scoring design provides a blunt force spread over a greater area.
- May explain why published data shows other scoring balloons to not generate as high of acute gain than cutting balloon.

Matsukawa, et al, Cardiovascular Intervention and Therapeutics (2019) 34:325 - 334



SCORING BALLOON CROSS SECTION

Competitive Specifications

WOLVERINE™ is compatible with smaller guide catheter and offer the broad size matrix to treat according to the type of lesions

	WOLVERINE™	Product A	Product B	Product C
Guide Cath Compatibility	5F, 6F	6F	6F	5F
Size Matrix: Diameter (mm)	2, 2.25, 2.5, 2.75, 3, 3.25, 3.5, 3.75, 4	2, 2.5, 3, 3.5	2, 2.25, 2.5, 2.75, 3, 3.25, 3.5, 4	2, 2.5, 3, 3.5, 4
Size Matrix: Length (mm)	6, 10, 15	6, 10, 15	13	10, 15, 20
Pressures (ATM)	NOM: 6 RBP: 12	NOM: 8 RBP: 16-20	NOM: 6 RBP: 14	NOM: 12 RBP: 20
Catheter Length (cm)	143	137	142	139
Balloon Compliance	Non-Compliant	Semi-Compliant	Semi-Compliant	Non-Compliant
Plaque Mod Method	3 or 4 evenly spaced atherotomes	Wire wrapped balloon	3 scoring elements	Single scoring wire

Clinical Study: Cutting Balloon vs. Scoring Balloon in Severely Calcified Patients

Plaque modification using a cutting balloon is more effective for stenting of heavily calcified lesion than other scoring balloons

Primary Investigator

 Ryuichi Matsukawa, Fukuoka Red Cross Hospital, Fukuoka, Japan

Study Design

 Retrospective analysis of 156 patients treated for calcified coronary artery disease with either Cutting Balloon (n=30), NSE Scoring Balloon (n=39) or Scoreflex Scoring Balloon (n=87) from April 2015 – December 2017

Notable Patient Characteristics

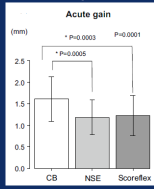
- Patients in all groups had similar characteristics including age, gender, lesion location, Minimum Lumen Diameter, reference vessel diameter and balloon to artery ratio
- However, the cutting balloon patients had a significantly higher rate of severe calcification (83.3%) than NSE (59%) or Scoreflex (44.8%)

Summary of Key Results



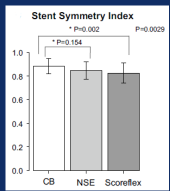
30% HIGHER ACUTE GAIN

Despite a significantly higher percentage of severe calcium, cutting balloon resulted in a statistically significant higher acute gain than scoring balloon.



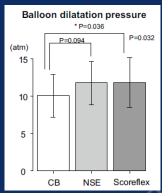


Cutting balloon also had a superior effect on stent symmetry index, meaning that the stent lumen was more symmetrical than with scoring balloon.





This 30% higher acute gain was achieved with cutting balloon despite using a statistically significant lower inflation pressure than scoring balloon.





Brief Summary

WOLVERINE™ Brief Summary

PRECAUTIONS

The device should be used only by physicians trained in the performance of PTCA.

If difficulty is experienced during balloon inflation, do not continue; remove the device and do not attempt to use it.

Infusion of any medium through the guidewire lumen other than heparinized saline may compromise device performance.

Do not attempt to reposition a partially inflated balloon.

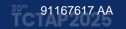
Do not use a guidewire having a diameter greater than 0.014 in (0.36 mm).

Potential ADVERSE EVENTS

Potential adverse events include, but are not limited to, the following:

- Abrupt closure
- Acute myocardial infarction
- · Angina or unstable angina
- Arrhythmias, including ventricular fibrillation
- Arteriovenous fistula
- Cardiac tamponade/pericardial effusion
- Cardiogenic shock
- Cerebrovascular accident/stroke
- Coronary aneurysm
- Coronary artery bypass graft surgery
- · Coronary artery spasm
- Coronary vessel dissection, perforation, rupture, or injury, possibly requiring surgical repair or intervention
- Death
- Drug reactions, including allergic reaction to contrast medium
- Embolism
- · Hemodynamic compromise
- · Hemorrhage or hematoma
- · Hypo/hypertension

- Infection
- Minor vessel trauma
- Myocardial ischemia
- Percutaneous re-intervention
- Pseudoaneurysm (at vascular access site)
- Pyrogenic reaction
- Renal failure
- Respiratory insufficiency
- Restenosis of the dilated vessel
- Side branch occlusion
- Slow flow/no reflow
- Thrombosis
- Total occlusion of the coronary artery or bypass graft
- Transient ischemic attack
- Vasovagal reaction
- Ventricular irritability/dysfunction
- Vessel trauma requiring surgical repair or intervention
- · Volume overload



WOLVERINE™ Brief Summary

CAUTION: Rx only. Prior to use, please see the complete "Directions for Use" for more information on Indications, Contraindications, Warnings, Precautions, Adverse Events, and Operator's Instructions.

INTENDED USE / INDICATIONS FOR USE

The Wolverine Cutting Balloon Device is indicated for use in patients with coronary vessel disease who are acceptable candidates for coronary artery bypass graft surgery, should it be urgently needed, for the purpose of improving myocardial perfusion. In addition, the target lesion should possess the following characteristics:

- Discrete (< 15 mm in length), or tubular (10 mm to 20 mm in length)
- Reference vessel diameter (RVD) of 2.00 mm to 4.00 mm
- Readily accessible to the device
- · Light to moderate tortuosity of proximal vessel segment
- Nonangulated lesion segment (< 45°)
- · Smooth angiographic contour
- Absence of angiographically visible thrombus

CONTRAINDICATIONS

The WOLVERINE Cutting Balloon Device is contraindicated for use in:

Delivery through the side cell of a previously placed stent as the deflated Cutting Balloon could become entangled in the stent. Coronary artery spasm in the absence of a significant stenosis.

WARNINGS

- Exercise extreme care when treating a lesion distal to a stent. When treating lesions at a bifurcation, the device can be used prior to placing a stent, but should not be taken through the side cell of a stent to treat the side branch of a lesion at a bifurcation.
- The atherotomy process, because of its mechanism of action, may pose a greater risk of perforation than that observed with conventional Percutaneous Transluminal Coronary Angioplasty (PTCA). To reduce the potential for vessel damage, the inflated diameter of the device should approximate a 1.1:1 ratio of the diameter of the vessel just proximal and distal to the stenosis.
- The atherotomy process in patients who are not acceptable candidates for coronary artery bypass surgery requires careful consideration, including possible hemodynamic support during the atherotomy process, as treatment of this patient population carries special risk.
- Balloon pressure should not exceed the rated burst pressure.
- When performing percutaneous atherotomy, the availability of on-site surgical backup should be included as a clinical consideration.

