

DECISION-CTO

**Optimal Medical Therapy With or Without
Stenting For Coronary Chronic Total Occlusion**

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Background

- Benefits of successful CTO-PCI include reduced angina frequency and improvements in quality of life, left ventricular ejection fraction, or survival.
- However, CTO-PCI can lead to procedure-related complications. In addition, the evidence for CTO-PCI was obtained from observational studies, most of which compared successful and failed CTO-PCI without a control group receiving optimal medical treatment.

DECISION CTO Trial

Design

- **DESIGN:** a prospective, open-label, randomized trial
- **OBJECTIVE:** To compare the outcomes of OMT alone with PCI coupled with OMT in patients with CTO.
- **PRINCIPAL INVESTIGATOR**
Seung-Jung Park, MD, PhD,
Asan Medical Center, Seoul, Korea

Participating Centers (N=19)

Country	Site	Investigator
Korea	Asn Medical center	Seung-Jung Park
India	Ruby Hall Clinic	Shirish Hiremath
Korea	Keimyung University Dongsan Medical Center	Seung Ho Hur
Korea	Korea University Guro Hospital	Seung Un Rha
Indonesia	Medistra Hospital	Teguh Santoso
Korea	The Catholic University of Korea, Daejeon ST. Mary's Hospital	Sung-Ho Her
Korea	Chungnam National University Hospital, Daejeon	Si Wan Choi
Korea	Kangwon National University Hospital	Bong-Ki Lee
Korea	Soon Chun Hyang University Hospital Bucheon, Bucheon	Nae-Hee Lee
Korea	Kangbuk Samsung Medical Center, Seoul	Jong-Young Lee
Korea	Gangneung Asan Hospital, Gangneung	Sang-Sig Cheong,
Thailand	King Chulalongkorn Memorial Hospital	Wasan Udayachalerm
Korea	Dong-A University Hospital, Busan	Moo Hyun Kim
Korea	Chonnam National University Hospital, Gwangju	Young-Keun Ahn
Korea	Bundang Cha Medical Center, Bundang	Sang Wook Lim
Korea	Ulsan University Hospital, Ulsan	Sang-Gon Lee
Korea	Hangang Sacred Heart Hospital, Seoul	Min-Kyu Kim
Korea	Sam Anyang Hospital, Anyang	Il-Woo Suh
Taiwan	Shin Kong Hospital	Jun Jack Cheng

Major Inclusion Criteria

- Silent ischemia, stable angina, or ACS
- **De novo** CTO located in a proximal to mid epicardial coronary artery with a reference diameter of ≥ 2.5 mm
- CTO was defined as a coronary artery obstruction with TIMI flow grade 0 of at least three months duration based on patient history.

Major Exclusion Criteria

- CTO located in
 - Distal coronary artery
 - 3 different vessel CTOs in any location
 - 2 proximal CTOs in separate coronary artery
 - left main segment
 - In-stent restenosis
 - Graft vessel
- LVEF < 30%
- Severe comorbidity

Study Procedures (1)

- Patients who were assigned to PCIs underwent CTO-PCI using DES within 30 days after randomization using standard procedures.
- In cases of failed CTO-PCI, additional attempts were allowed within 30 days after the index procedure.
- The use of specialized devices or techniques, and the choice of drug-eluting stent type were left to the operator's discretion.

Study Procedures (2)

- Revascularization for all significant non-CTO lesions within a vessel diameter of ≥ 2.5 mm for patients with multi-vessel coronary artery disease was recommended.
- Patients were prescribed guideline derived optimal medical treatment including aspirin, P2Y12 receptor inhibitors (>12 months in case of PCI), beta-blocker, CCB, nitrate, ACEi/ARB, and statin.
- Blood pressure and diabetic control, smoking cessation, weight control, and regular exercise were recommended.

Primary End Point

At 3 year, a composite of

- Death from any cause
- Myocardial infarction
 - Periprocedural MI: CK-MB > 5 times UNL
 - Spontaneous MI: any cardiac enzyme elevation
- Stroke
- Any repeat revascularization

Original Power Calculation

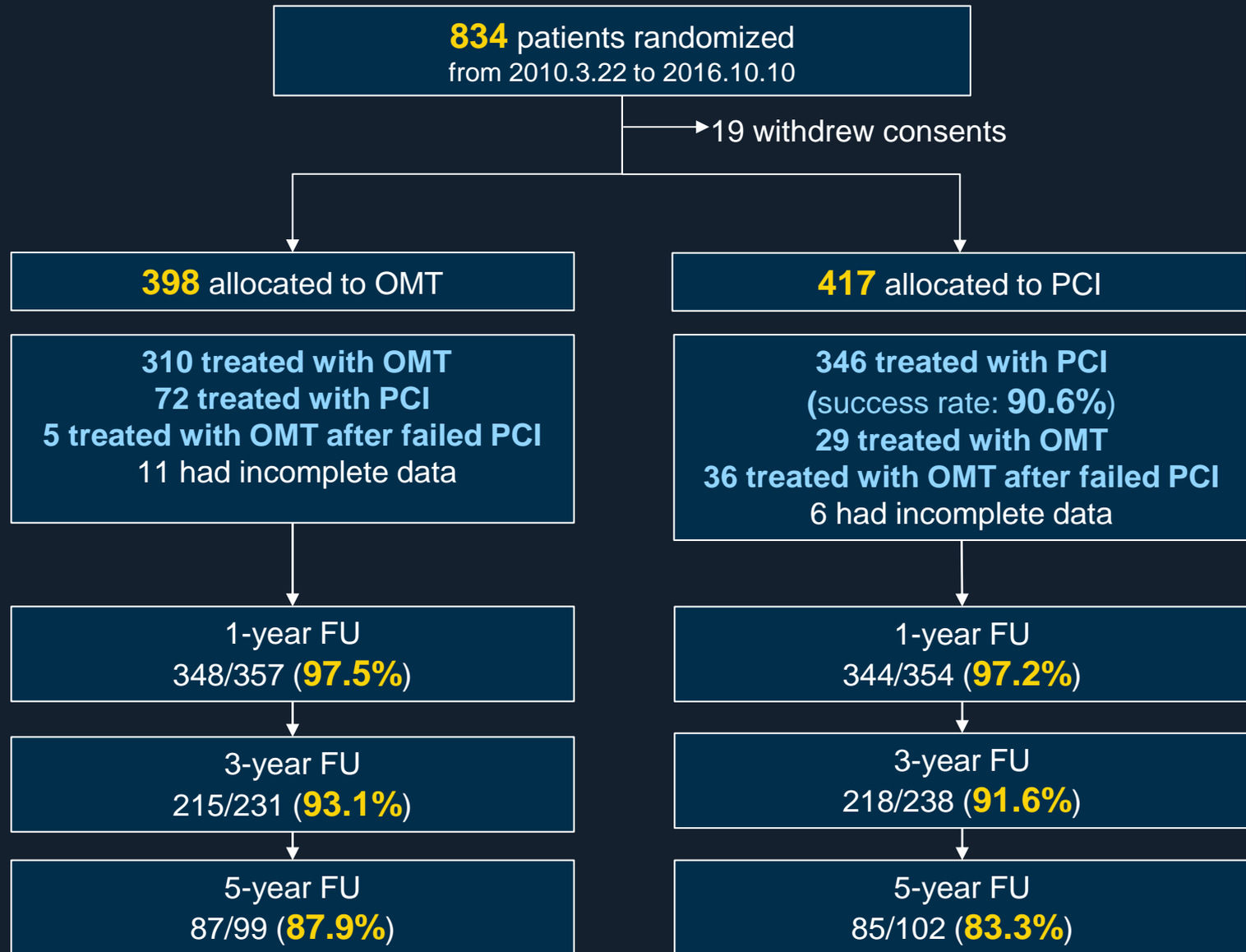
Non-inferiority Design for Primary Endpoint

- Assumed primary event rate: 17% at 3 years
- A noninferiority margin : event rate ratio ≥ 0.7
- A one-sided type I error rate : 0.025
- Power : 80%
- Dropout rate: 5%
- Assumed sample size: 1,284 patients

Premature Termination of Trial

- Because enrollment was slower than anticipated, enrollment was stopped in September 2016 as recommended by the data and safety monitoring board by which time 834 patients had been enrolled.
- The sponsor and study leadership were unaware of study results at the time of this decision.

Study Flow



Statistical Analysis

- All analyses were performed according to the intention-to-treat principle. Further sensitivity analyses were performed in the per-protocol and as-treated population.
- Hazard ratios (HRs) and 95% confidence intervals (CIs) were estimated using Cox proportional hazard models, with robust standard errors that accounted for clustering effect of stratified randomization.
- Noninferiority test using the Z-test with 95% CI of difference in the 3-year event rate.
- Survival curves were estimated using Cox model and the Kaplan-Meier method
- For quality of life analysis, we assumed the missing values were missing at random, and compared mean values of two groups using Student's t-test at specific time points.
- All P-values and CIs were two-sided. SAS software version 9.3 was used for all statistical analyses.

Baseline Characteristics

ITT Population

	OMT (N=398)	PCI (N=417)	P value
Age (years)	62.9±9.9	62.2±10.2	0.35
Male sex	315 (81.4%)	342 (83.2%)	0.50
BMI, kg/m ²	25.4±3.3	25.6±3.6	0.66
Hypertension	235 (60.7%)	261 (63.5%)	0.50
Diabetes mellitus	133 (34.4%)	132 (32.1%)	
Hypercholesterolemia	215 (55.6%)	248 (60.3%)	0.17
Current smoker	102 (26.4%)	125 (30.4%)	0.20
Previous PCI	74 (19.1%)	62 (15.1%)	0.13
Previous MI	34 (8.8%)	45 (10.9%)	0.31
Previous CABG	5 (1.3%)	4 (1.0%)	0.75
Chronic renal failure	5 (1.3%)	6 (1.5%)	0.84
LVEF, %	57.2±9.4%	57.2±9.8%	0.95

Baseline Characteristics

ITT Population

	OMT (N=398)	PCI (N=417)	P value
Clinical presentation			0.58
Stable angina	290 (74.9%)	297 (72.3%)	
Unstable angina	75 (19.4%)	84 (20.4%)	
AMI	22 (5.7%)	30 (7.3%)	
Location of CTO			0.71
LAD	161 (41.6%)	183 (44.5%)	
LCX	42 (10.9%)	40 (10.2%)	
RCA	184 (47.5%)	186 (45.3%)	
Multivessel disease	286 (73.9%)	301 (73.3%)	0.76
SYNTAX score	21.0±9.5	21.2±9.1	0.79
J-CTO score	2.3±1.2	2.2±1.2	0.23
Number of total stents	2.0±1.4	2.4±1.3	<0.001
Total stent length, mm	53.6±39.4	71.2±40.5	<0.001

CTO PCI Characteristics

Attempted PCI

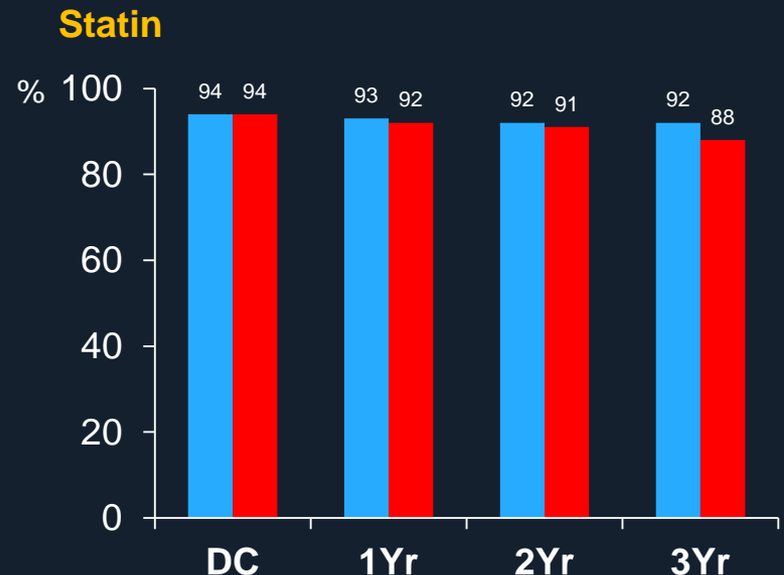
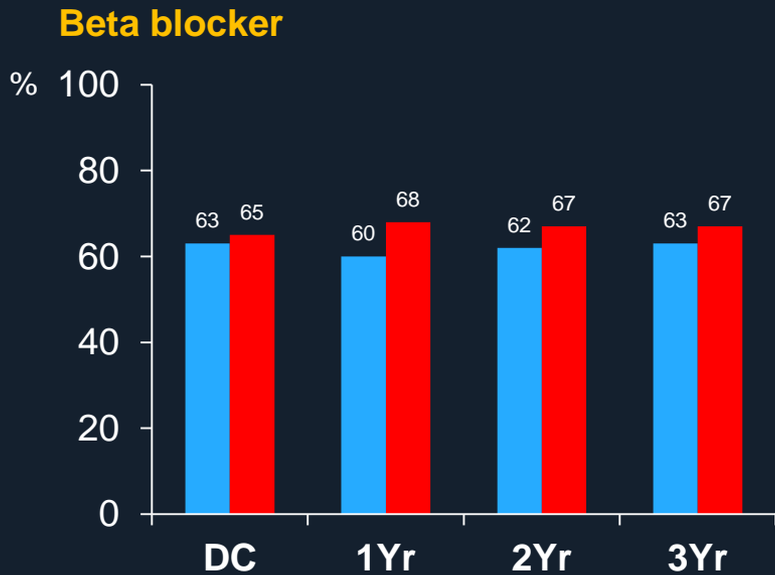
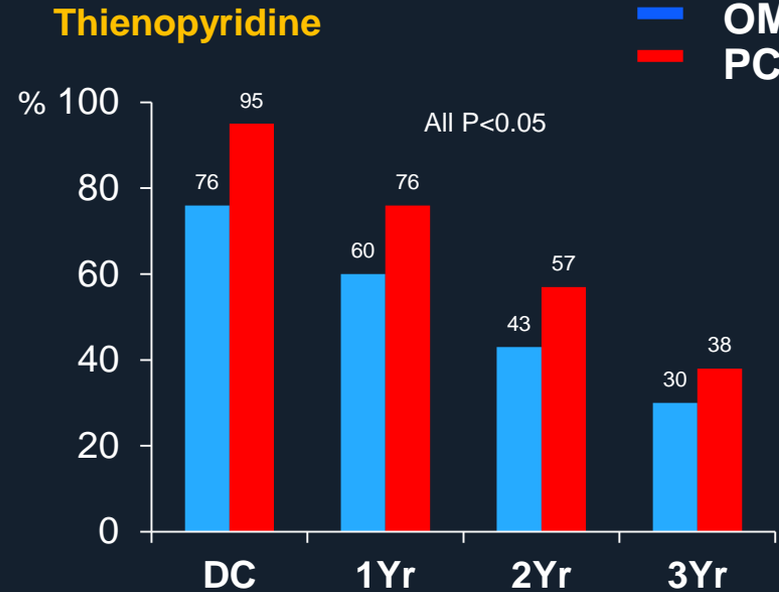
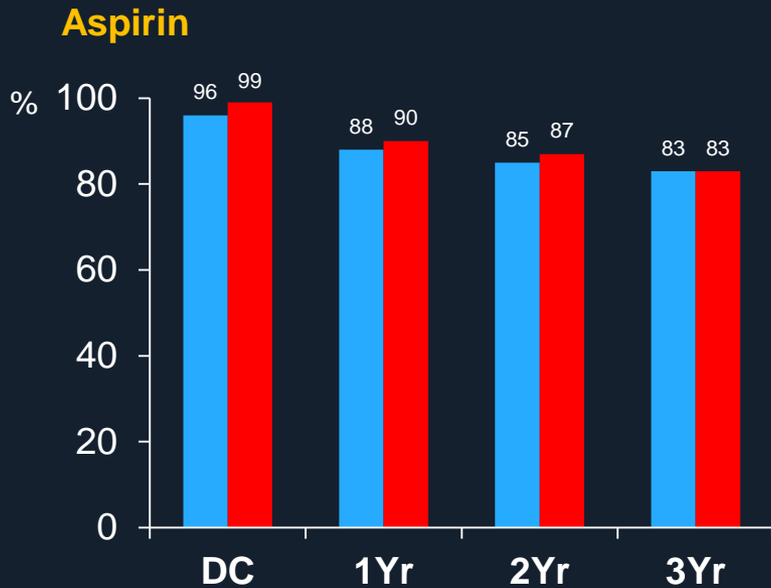
N=459

CTO PCI success	418 (91.1%)
Retrograde approach	113 (24.6%)
Lesion passaged wire	
Low penetration force wire	117/418 (28.0%)
Intermediate to high penetration force wire	301/418 (72.0%)
CTO technique	
Single wire technique only	309/418 (73.9%)
Parallel wire technique	72/418 (17.2%)
IVUS-guided wiring	25/418 (6.0%)
CART technique	55/418 (13.2%)
Additional back-up support	
Corsair	91/418 (21.8%)
Microcatheter other than Corsair	230/418 (55.0%)
Over-the-wire balloon	6/418 (1.4%)

Medication at Follow-Up

ITT Population

— OMT
— PCI

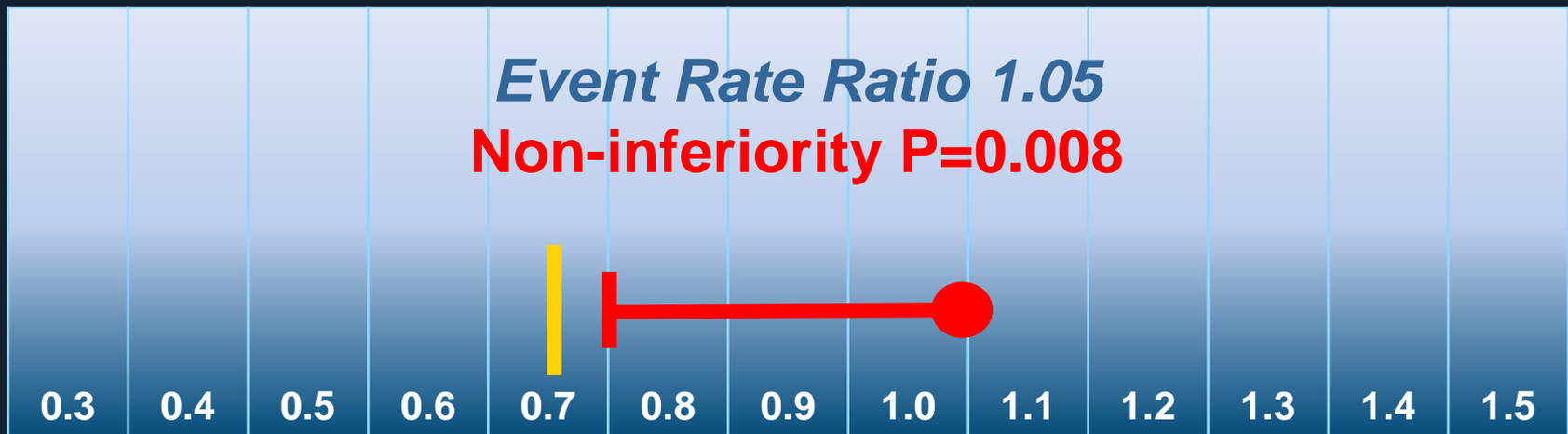


Noninferiority Test for Primary End Point at 3-Year

ITT Population

Estimated 3-year Event Rate OMT: 19.6% PCI: 20.6%

Prespecified non-inferiority margin: 0.7

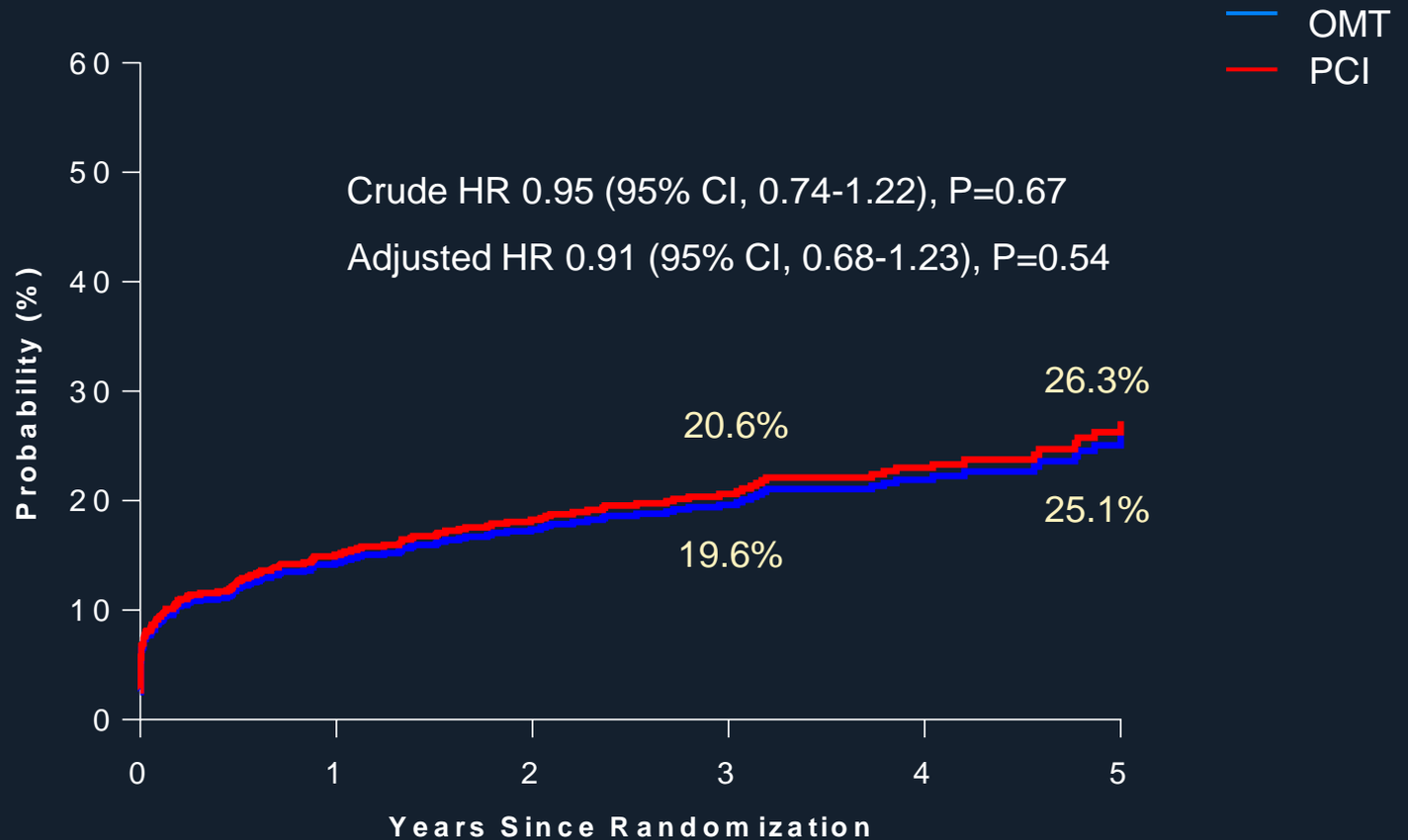


Event Rate Ratio of 3-year MACE rate (PCI/OMT)

 Lower 1-sided 97.5% CI

Primary End Point

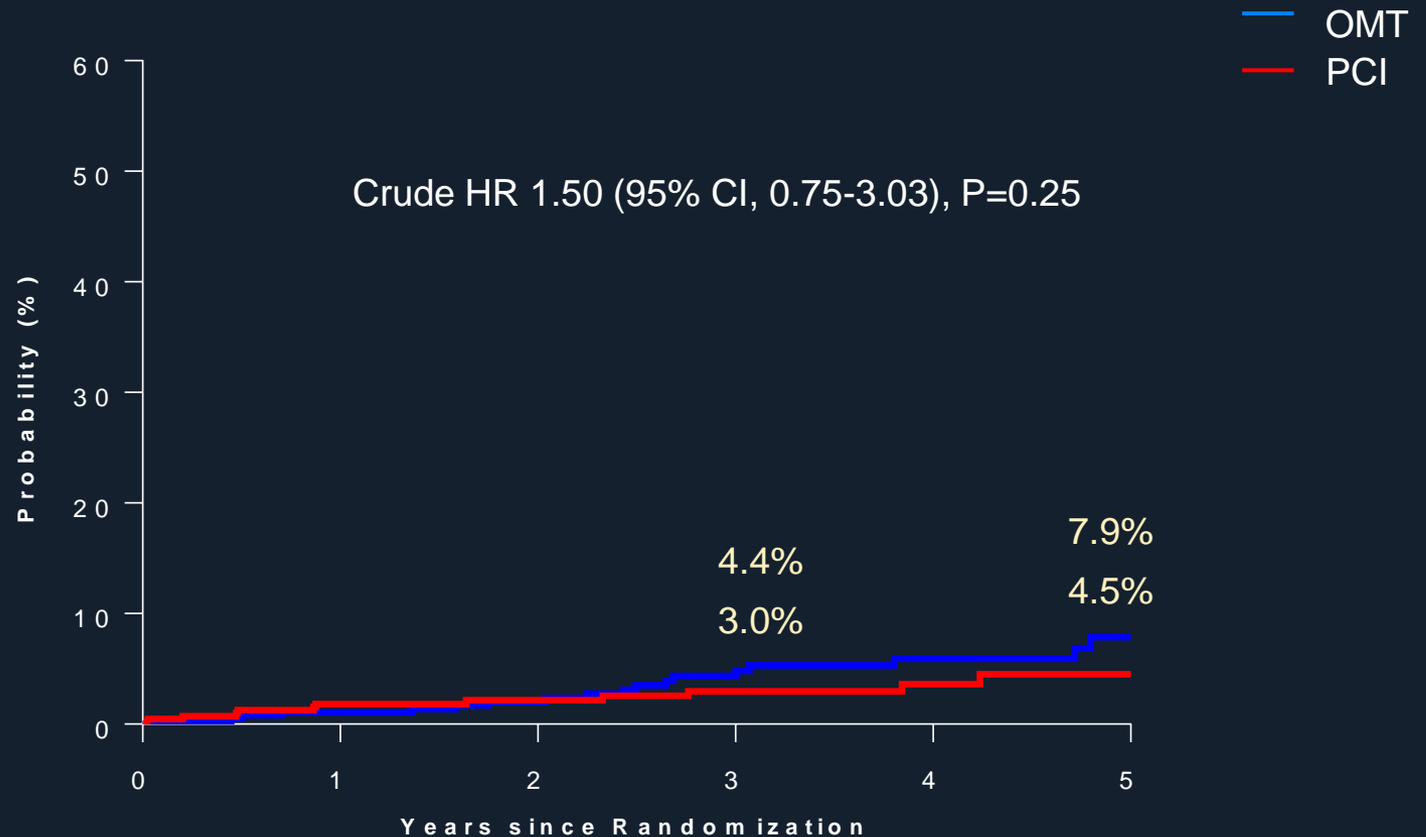
(Death, MI, Stroke, Any Repeat Revascularization)



No. at Risk

OMT	398	305	246	178	129	72
PCI	417	293	241	175	117	65

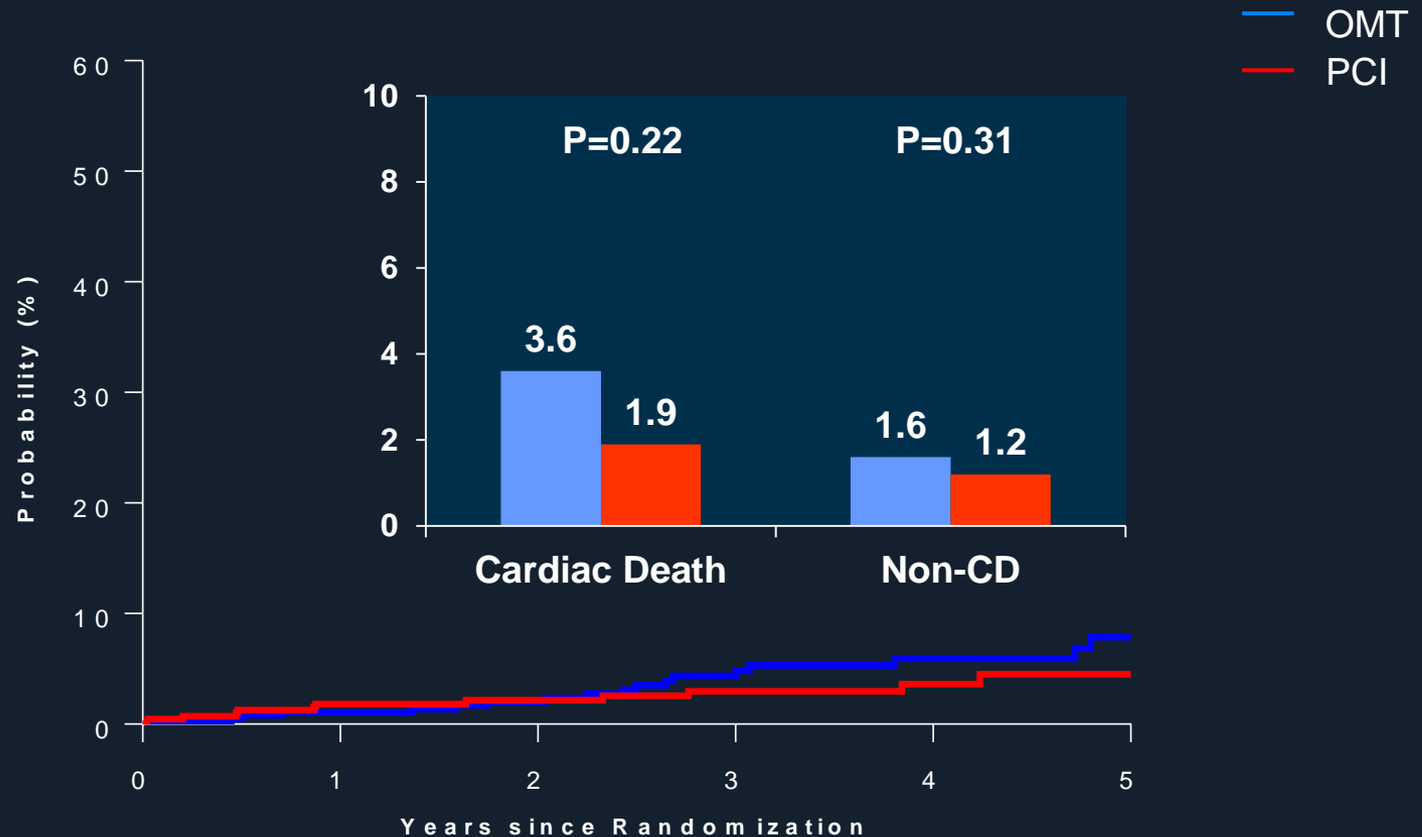
Death from any cause



No. at Risk

OMT	398	344	285	207	140	81
PCI	417	337	285	202	142	74

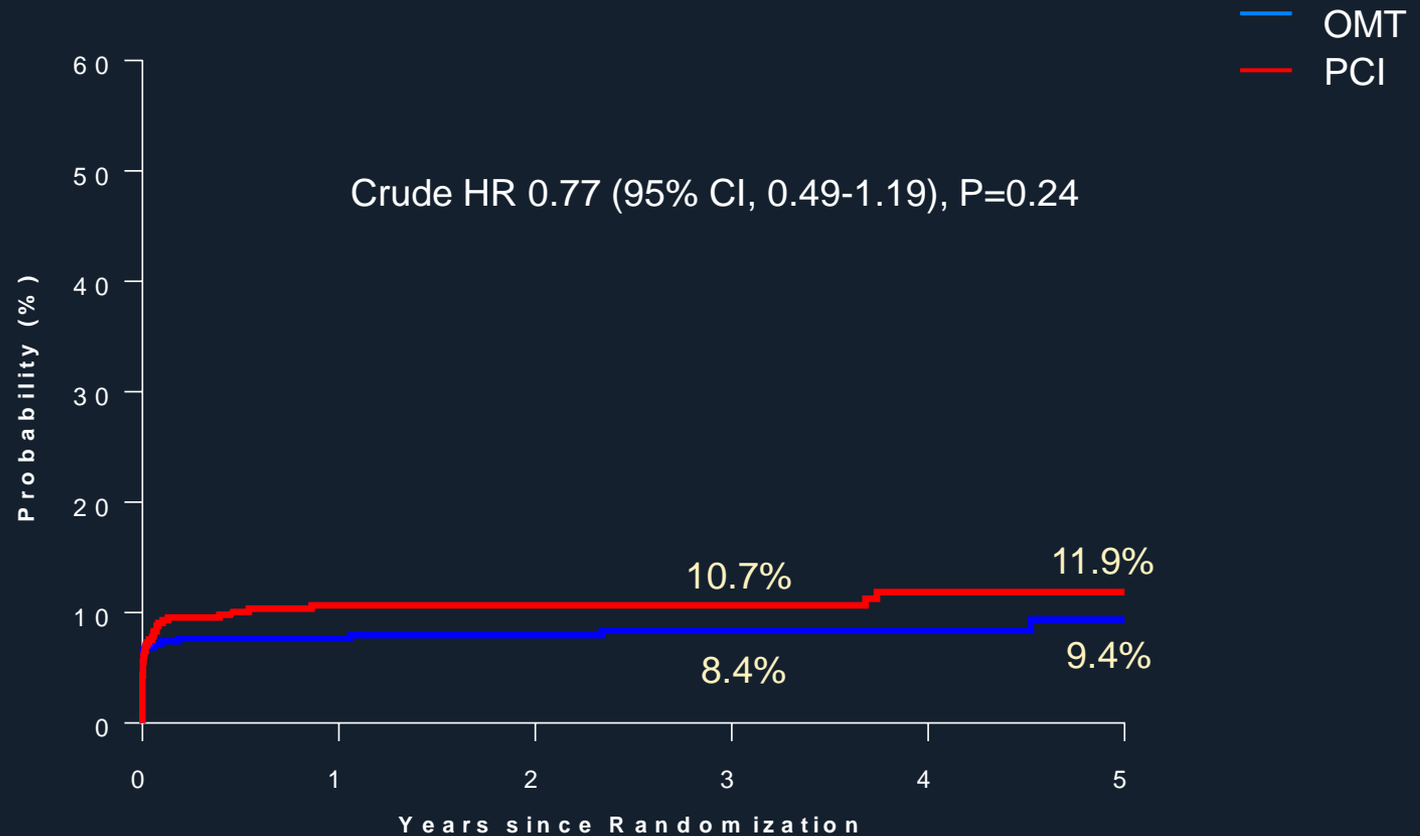
Death from any cause



No. at Risk

OMT	398	344	285	207	140	81
PCI	417	337	285	202	142	74

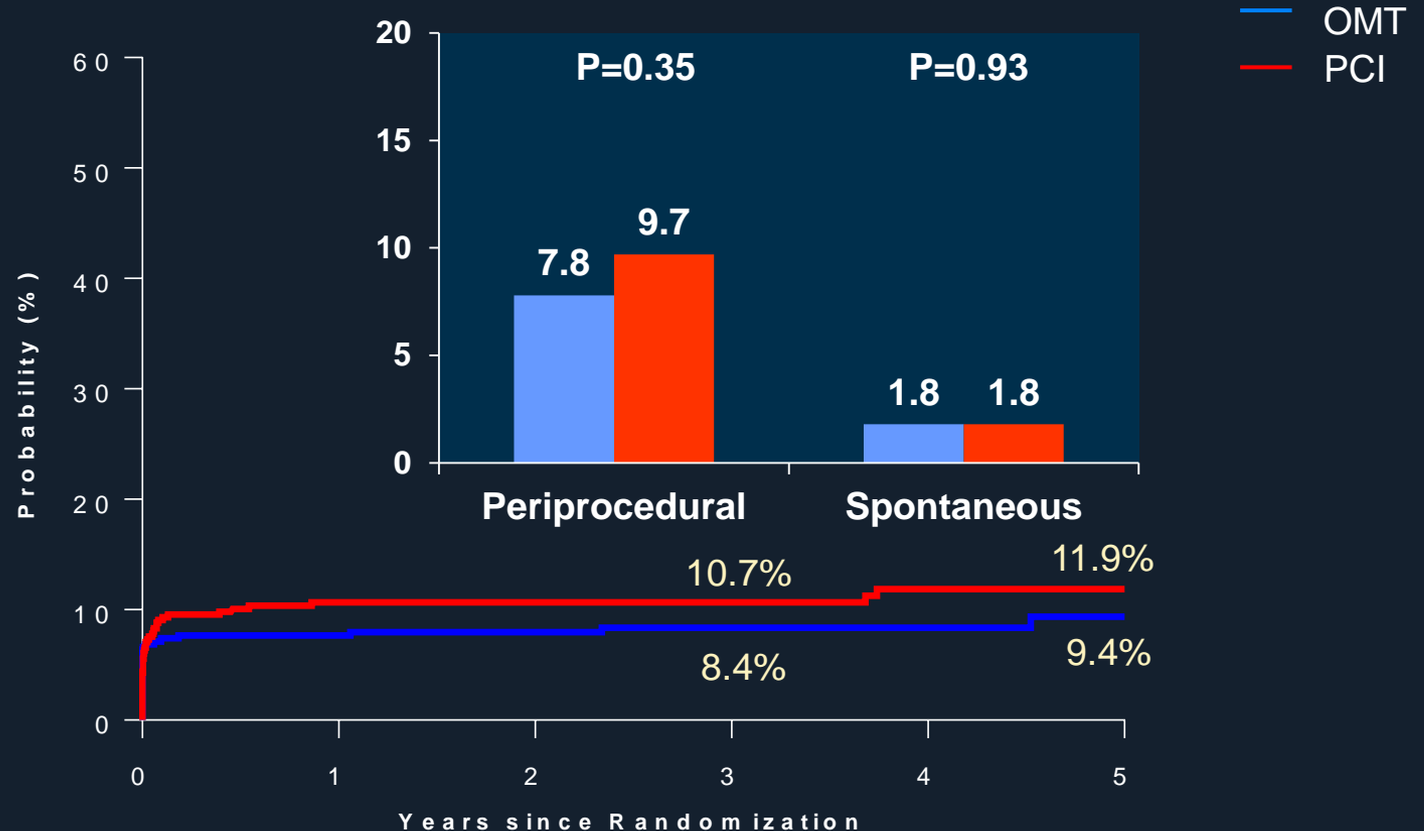
Myocardial Infarction



No. at Risk

OMT	398	317	260	189	129	73
PCI	417	300	255	181	125	64

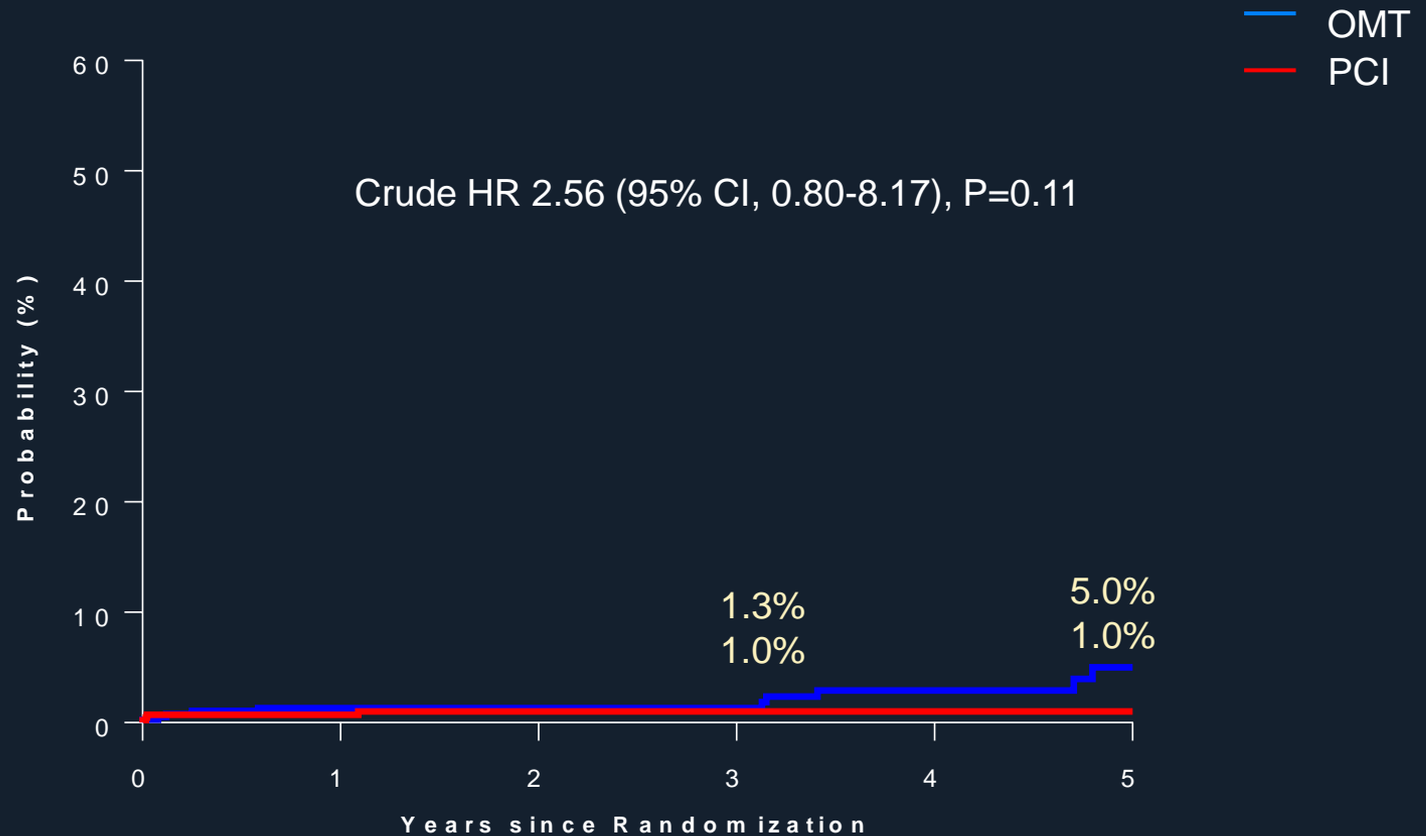
Myocardial Infarction



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OMT	398	317	260	189	129	73
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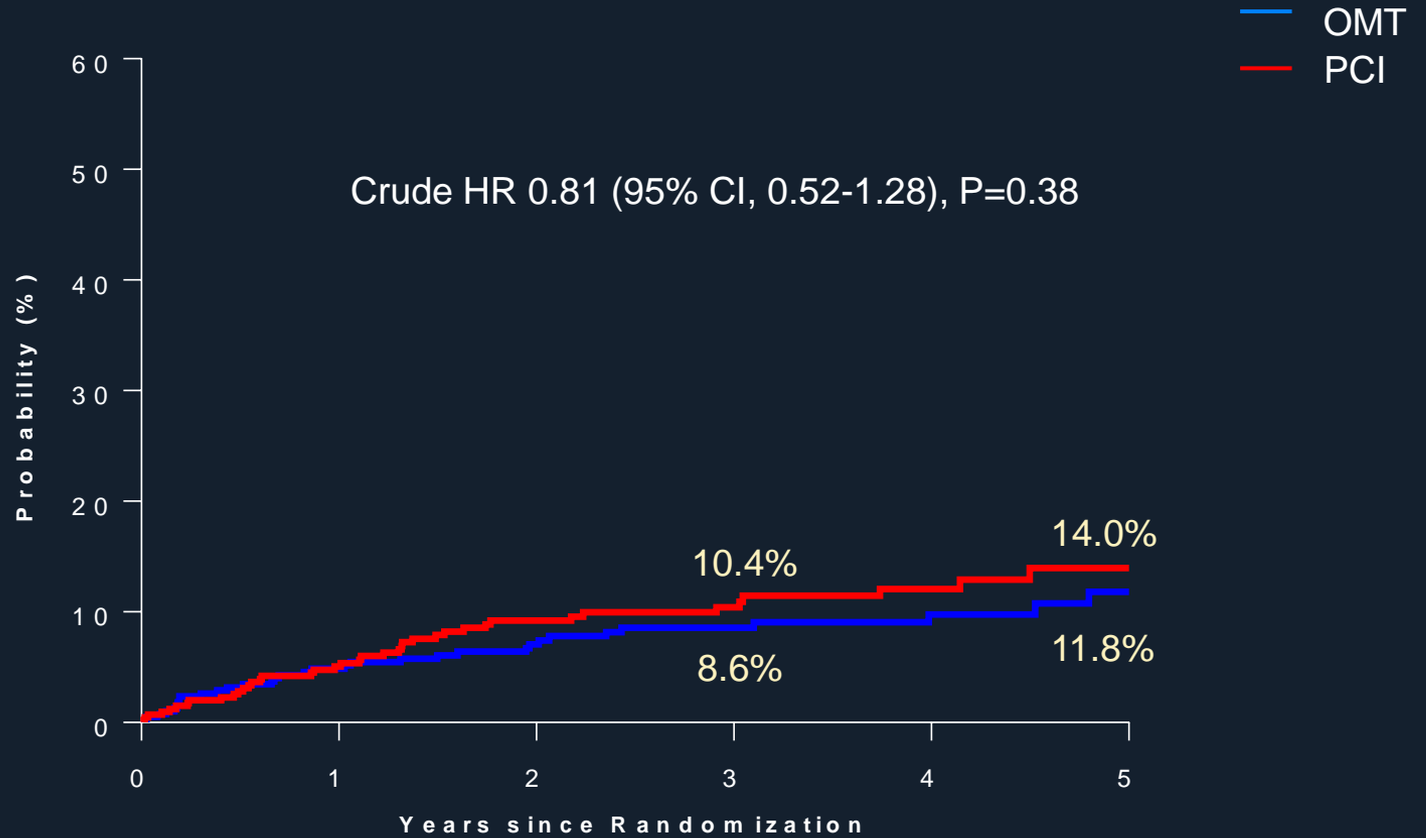
Stroke



No. at Risk

OMT	398	339	280	203	137	77
PCI	417	337	284	201	142	74

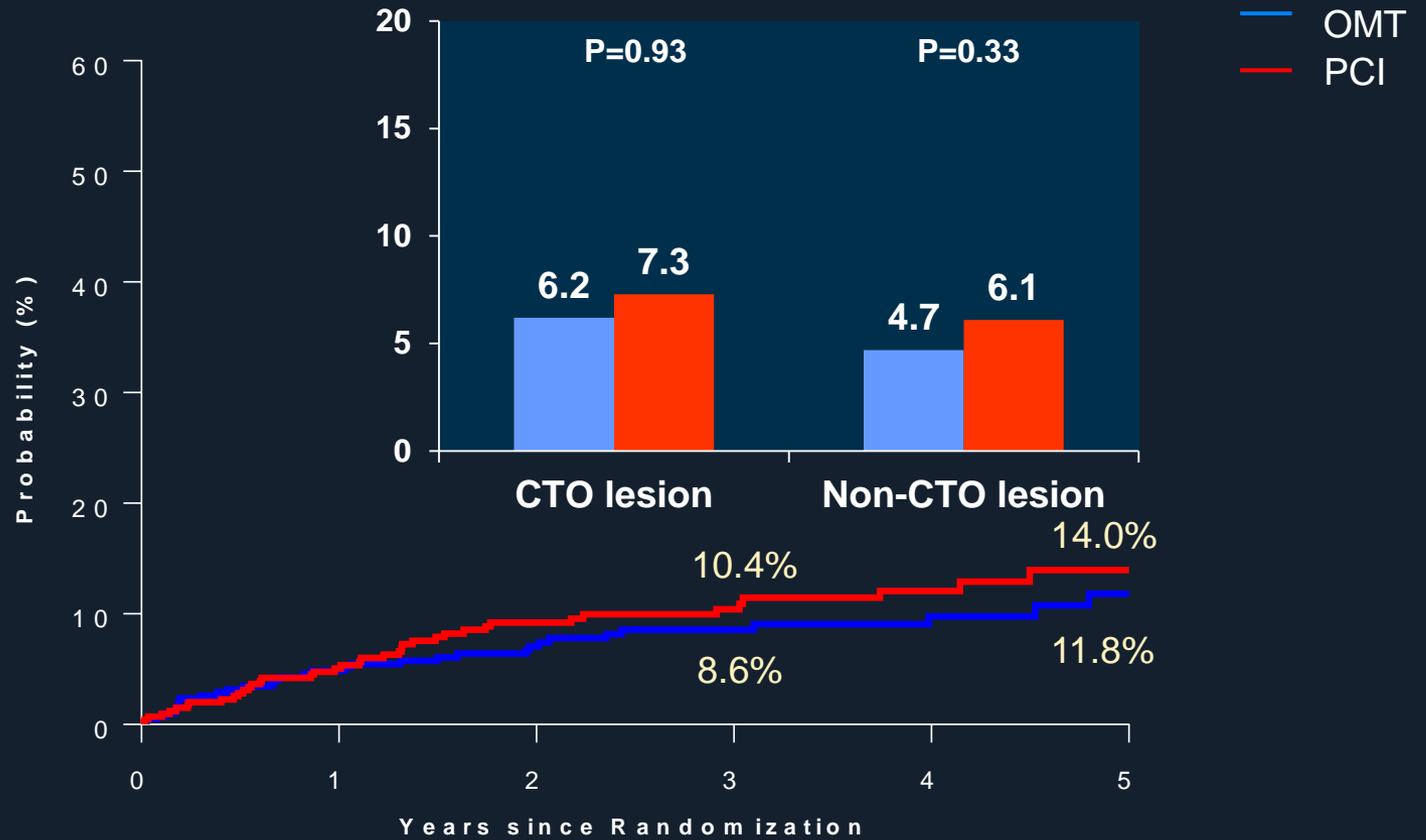
Repeat Revascularization



No. at Risk

OMT	398	330	270	292	129	74
PCI	417	321	259	181	129	65

Repeat Revascularization



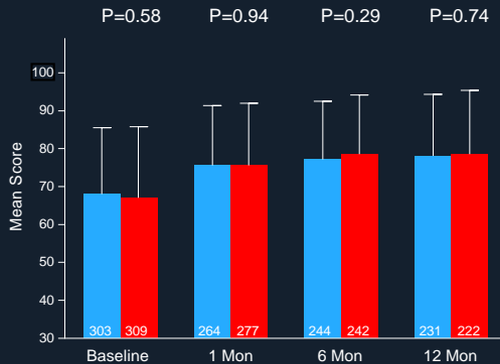
No. at Risk

OMT	398	330	270	292	129	74
PCI	417	321	259	181	129	65

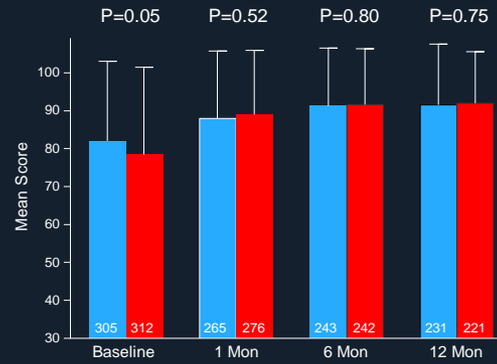
Quality of Life Measures Over Time

ITT Population

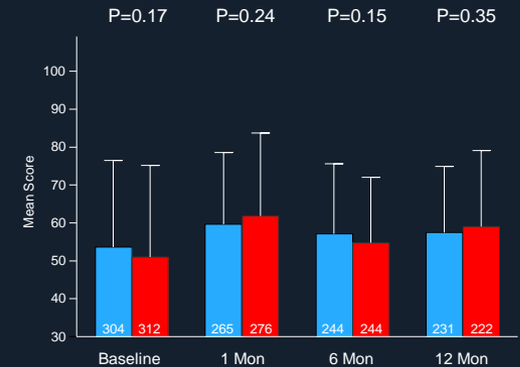
(A) EQ-5D Visual Analogue Scale



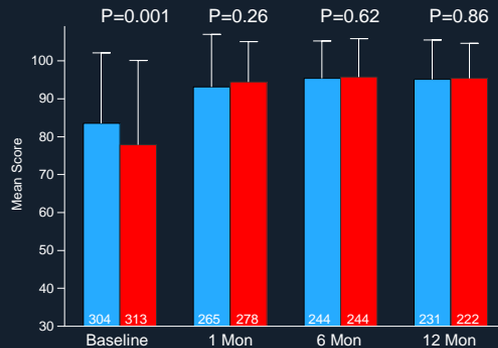
(B) SAQ, Physical Limitation



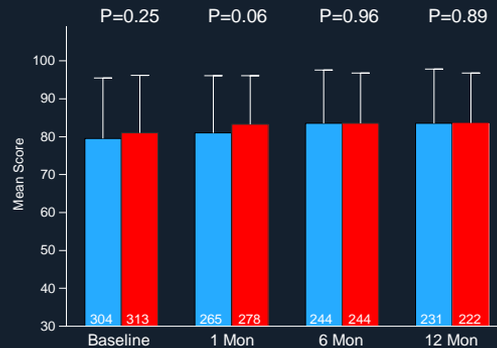
(C) SAQ, Angina Stability



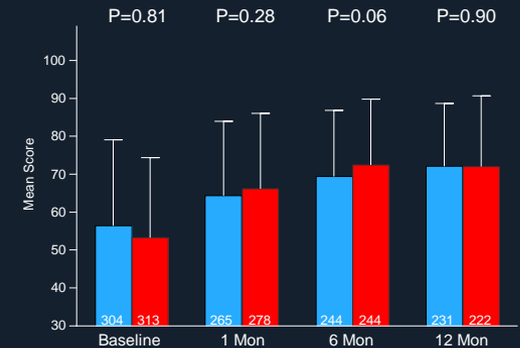
(D) SAQ, Angina Frequency



(E) SAQ, Treatment Satisfaction



(F) SAQ, Quality of Life



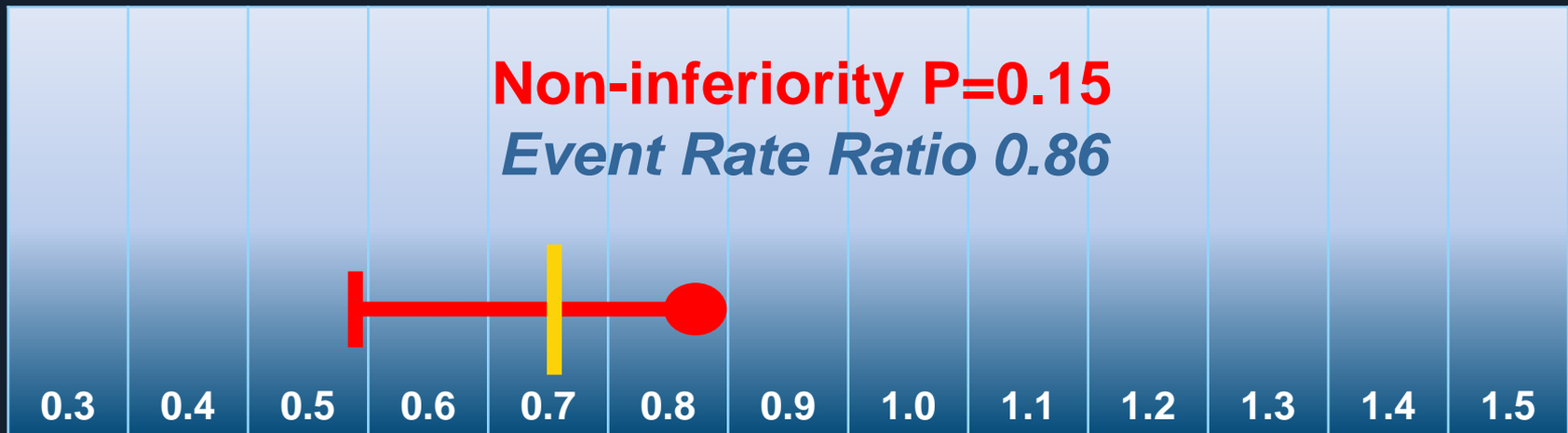
Per Protocol Analysis

Noninferiority Test for Primary End Point at 3-Year

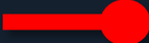
Per-Protocol Population

Estimated 3-year Event Rate OMT: 22.3% PCI: 19.0%

Prespecified non-inferiority margin: 0.7

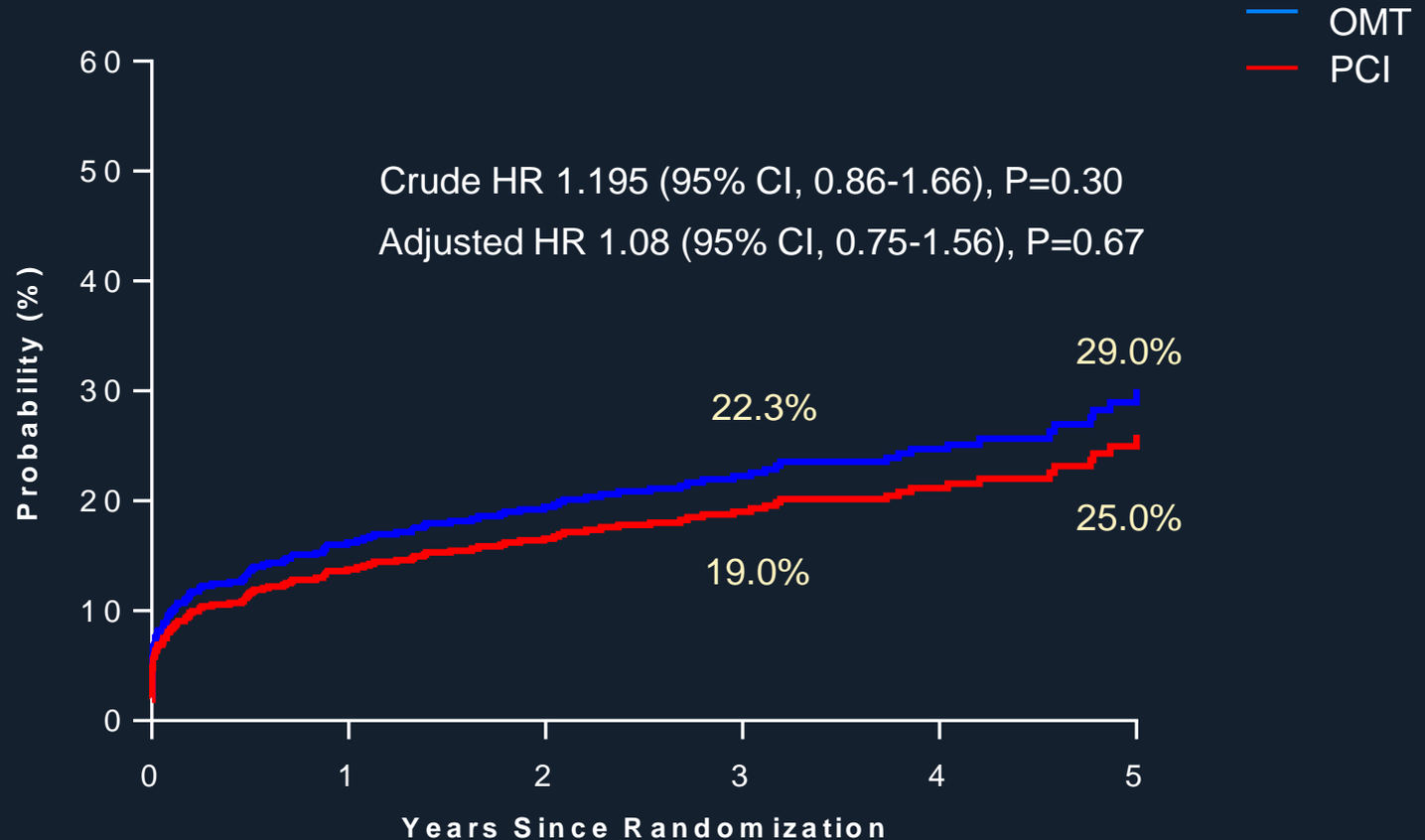


Event Rate Ratio of 3-year MACE rate (PCI/OMT)

 Lower 1-sided 97.5% CI

Primary End Point

(Death, MI, Stroke, Any Repeat Revascularization)



No. at Risk

OMT	310	241	190	131	95	54
PCI	346	250	209	150	98	52

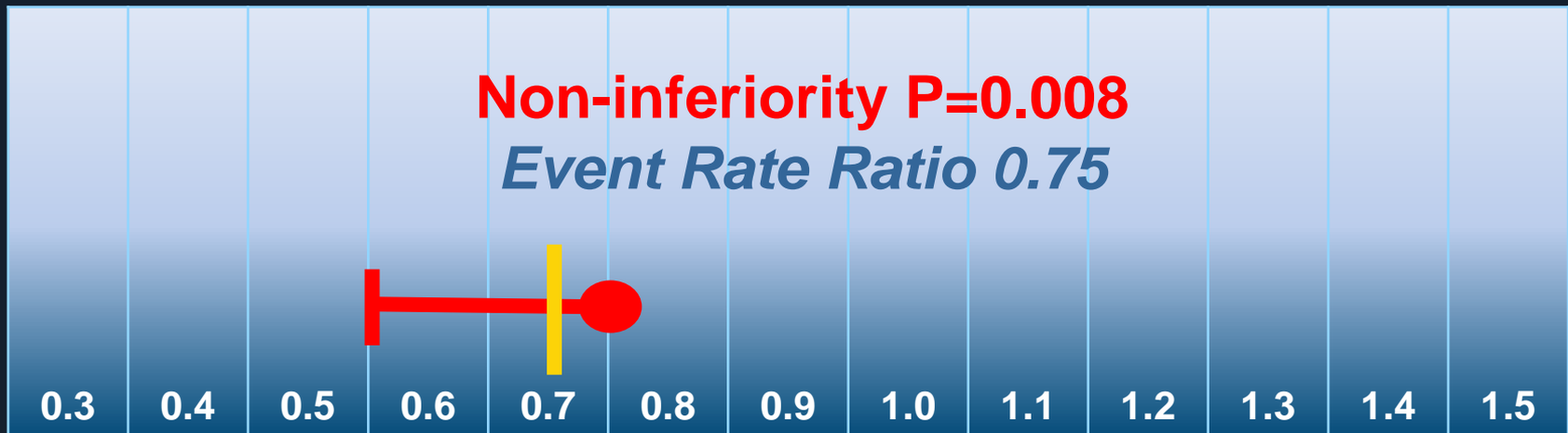
As Treated Analysis

Noninferiority Test for Primary End Point at 3-Year

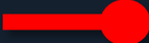
As-Treated Population

Estimated 3-year Event Rate OMT: 19.6% PCI: 20.6%

Prespecified non-inferiority margin: 0.7

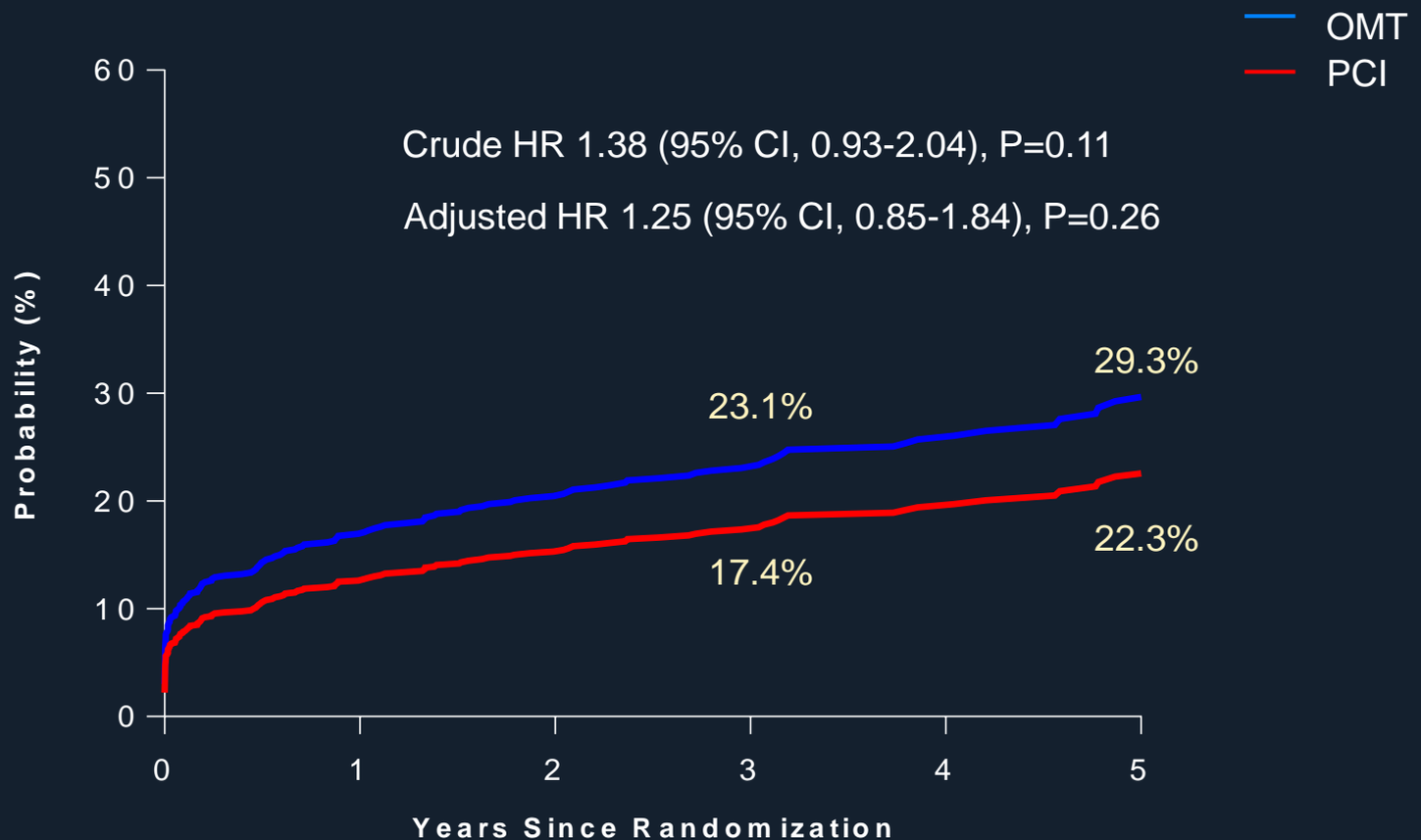


Event Rate Ratio of 3-year MACE rate (PCI/OMT)

 Lower 1-sided 97.5% CI

Primary End Point

(Death, MI, Stroke, Any Repeat Revascularization)



No. at Risk

OMT	380	290	226	160	118	70
PCI	418	309	261	193	128	67

Conclusion

- The DECISION-CTO trial is the first randomized clinical trial to compare the strategy of OMT alone with that of PCI in patients with coronary CTO.
- The ITT analysis showed that OMT as an initial strategy was non-inferior to PCI with respect to the primary endpoint of the composite of death, MI, stroke, or any revascularization at 3 years.
- The measures of health-related quality of life in the OMT and the PCI groups were comparable throughout the follow-up period

Conclusion

- However, OMT did not meet the statistical criteria for noninferiority compared with PCI in the predefined per-protocol and as-treated population-based analyses, although event rates were not significantly different between groups at 3 years.
- This study suggested that OMT could be a reasonable initial treatment strategy for coronary CTO compared with CTO-PCI. Further randomized clinical trials are necessary.



Thank You !!

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