# One-Month Dual Antiplatelet Therapy Followed by Clopidogrel Monotherapy versus

Standard 12-Month Dual Antiplatelet Therapy with Clopidogrel After Drug-Eluting Stent Implantation:



#### Hirotoshi Watanabe

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# Background

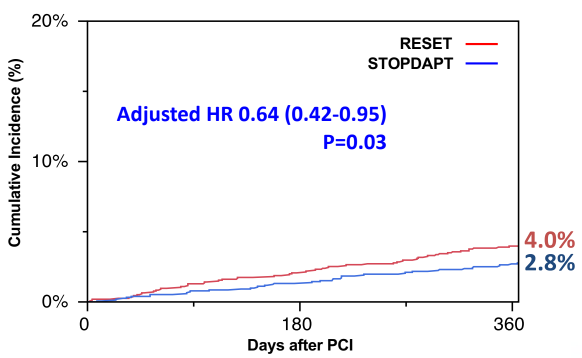
- Mandatory 1-month DAPT had been the standard care after BMS implantation.
- DAPT duration was prolonged after introduction of DES without firm scientific evidence.
- New generation DES has substantially reduced stent thrombosis.
- Prolonged DAPT is inevitably associated with increase in bleeding.
- Bleeding is associated with subsequent mortality risk at least comparable to that of MI.
- Therefore, very short mandatory DAPT duration after DES might be an attractive option, if not associated with increase in ischemic events disproportionate to the reduction in bleeding events.



### STOPDAPT

# Prospective multicenter open-label single arm trial evaluating 3-month DAPT after CoCr-EES implantation

Primary Endpoint
Cardiovascular death, MI, Stroke, Definite ST, and Bleeding



Cardiovasc Interv Ther 2016; 31: 196–209.



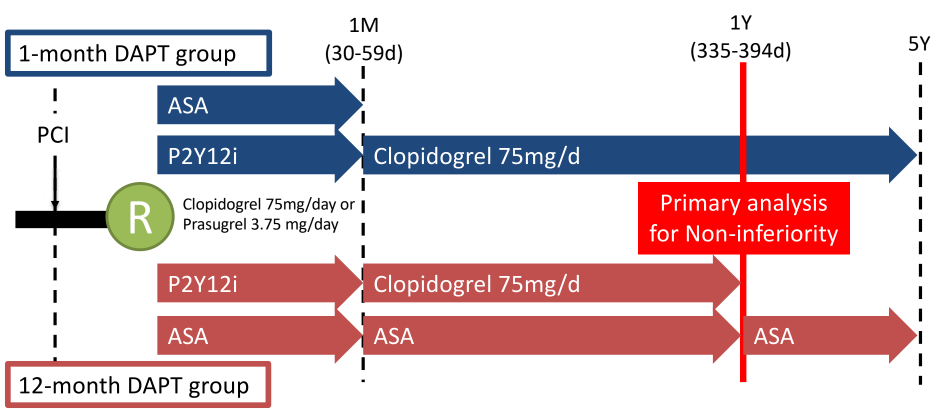
# **Objective**

The objective of the STOPDAPT-2 trial is to explore the safety and efficacy of the experimental regimen of 1-month DAPT followed by clopidogrel monotherapy as compared with the standard 12-month DAPT with aspirin and clopidogrel after implantation of cobalt-chromium everolimus-eluting stents (CoCr-EES).



# **STOPDAPT-2:**

Prospective multicenter open-label randomized trial comparing 1-month versus 12-month DAPT after CoCr-EES implantation with limited exclusion criteria.





# **Study Organization**

#### **Steering Committee**

Takeshi Kimura (PI)
Kazushige Kadota
Ken Kozuma
Yoshihiro Morino
Keiichi Igarashi-Hanaoka
Yuji Ikari
Kengo Tanabe
Kenji Ando
Koichi Nakao
Kazuya Kawai
Mitsuru Abe

#### **Trial Statistician**

Takeshi Morimoto

#### **Clinical Event Committee**

Yoshihisa Nakagawa Yutaka Furukawa Masahiro Natsuaki Hiroki Shiomi Toshiaki Toyota

#### **Safety Evaluation Committee**

Shunichi Miyazaki Ryuji Nohara

#### **Coordinating Center**

Research Institute for Production Development, Kyoto, Japan Saori Tezuka Yumika Fujino

#### **Angiography Core Laboratory**

Cardio Core Japan, Tokyo, Japan

#### Study administrative staff

Masahiro Natsuaki Hirotoshi Watanabe Toshiaki Toyota Toshikazu Jinnai

#### **Funded by**

Abbott Vascular Japan, Co., Ltd.



# 90 Participating Centers

Teine Keijinkai Hospital Hokko Memorial Hospital Hirosaki University Hospital Iwate Medical University Hospital Sendai Kousei Hospital

Sendai Kousei Hospital Sendai Cardiovascular Center

Tohoku Medical and Pharmaceutical University Hospital

Nakadori General Hospital Nihonkai General Hospital Hoshi General Hospital

Jichi Medical University Hospital

Mashiko Hospital

Mitsui Memorial Hospital Juntendo University Hospital The Fraternity Memorial Hospital

Edogawa Hospital

Showa University Koto Toyosu Hospital Tokyo Women's Medical University Hospital

Tokyo General Hospital

Juntendo University Nerima Hospital

Kawakita General Hospital Sakakibara Heart Institute

Tokyo Metropolitan Tama Medical Center

Minamino Cardiovascular Hospital

Higashiyamato Hospital

St. Marianna University School of Medicine Hospital

Yokohama Rosai Hospital

Showa University Fujigaoka Hospital Saiseikai Yokohamashi Tobu Hospital Yokohama City University Medical Center Kitasato University Hospital Hiratsuka Kyosai Hospital

Tokai University Hospital

Kimitsu Chuo Hospital

Kanazawa Cardiovascular Hospital

University of Fukui Hospital

Municipal Tsuruga Hospital

University of Yamanashi Hospital Gifu Prefectural General Medical Center

Ogaki Municipal Hospital

Juntendo University Shizuoka Hospital

Shizuoka General Hospital

Japanese Red Cross Nagoya Daini Hospital

Handa City Hospital
Tosei General Hospital

Ichinomiyanishi Hospital

Yokkaichi Hazu Medical Center

Matsusaka Central General Hospital

Nabari City Hospital

Otsu Red Cross Hospital

Hikone Municipal Hospital

Kyoto University Hospital Kyoto Medical Center

. Mitsubishi Kyoto Hospital

Kitano Hospital

Osaka Red Cross Hospital

National Cerebral and Cardiovascular Center

Kindai University Hospital Mimihara General Hospital

Bell Land General Hospital

Kobe City Medical Center General Hospital

Kindai University Nara Hospital

Tenri Hospital

Japanese Red Cross Wakayama Medical Center

Wakayama Medical University Hospital

Shimane University Hospital

Japanese Red Cross Okayama Hospital

Kurashiki Central Hospital

Hiroshima University Hospital

Iwakuni Medical Center

Tokuyama Central Hospital

Shimonoseki City Hospital Tokushima University Hospital

Tokushima Red Cross Hospital

Kagawa Prefectural Central Hospital

Ehime Prefectural Central Hospital

Matsuyama Red Cross Hospital

Chikamori Hospital

Kokura Memorial Hospital

Hospital of University of Occupational and Environmental Health Japan

Saiseikai Fukuoka General Hospital

Fukuoka Tokushukai Hospital

**Kumamoto University Hospital** 

Saiseikai Kumamoto Hospital

Japanese Red Cross Kumamoto Hospital

Miyazaki Prefectural Nobeoka Hospital

Ibusuki Medical Center

Izumi Regional Medical Center

Urasoe General Hospital

Nakagami Hospital



#### **Inclusion Criteria**

- PCI with exclusive use of CoCr-EES (Xience<sup>™</sup> series)
- No major complications during hospitalization for index PCI
- No plan for staged PCI
- Patients who could take DAPT with aspirin and P2Y<sub>12</sub> inhibitors

# **Key Exclusion Criteria**

- Needs for oral anticoagulants
- History of intracranial hemorrhage



# **Endpoints**

### Primary endpoint:

Net adverse cardiovascular events (NACE: Ischemia and Bleeding)

 A composite of cardiovascular death, MI, Definite ST, Stroke, or TIMI major/minor bleeding

### Major secondary endpoints:

Ischemic composite endpoint

- A composite of cardiovascular death, MI, Definite ST, or Stroke
   Bleeding endpoint
- TIMI major/minor bleeding

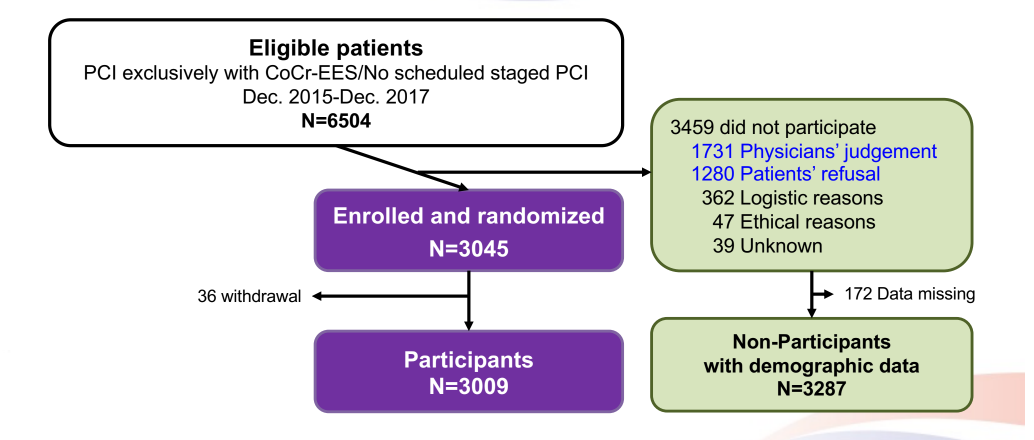


# Sample Size Calculation

- Hypothesis: Non-inferiority of 1-month DAPT to 12-month DAPT for the primary endpoint at 1-year
- Assumption: Event rate at 1-year: 4.6% (Based on RESET study).
- Non-inferiority margin; 50% on the hazard ratio scale
- Randomization ratio: 1:1
- One-sided alpha: 0.025
- Power: 85%
- Sample size: 3000 patients (1500 in each arm)



# Study Flow



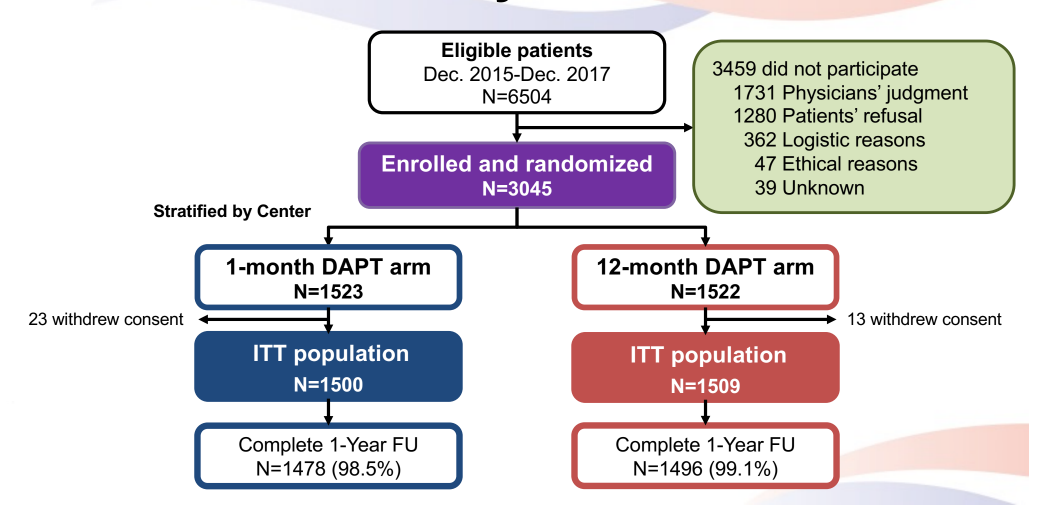
### STOPDAPT-2

# Participants vs Non-participants

	Participants N=3009	Non-participants N=3287	P value
Age, y	68.6±10.7	70.0±11.7	<0.001
ACS	38%	39%	0.61
STEMI	19%	22%	0.003
Prior MI	14%	23%	<0.001
Prior 1st-generation DES implantation	4%	6%	<0.001
Diabetes	39%	39%	0.47
Severe CKD	6%	9%	<0.001
Dialysis	3%	5%	<0.001
Target of LMCA	3%	5%	<0.001
Two or more target vessels	7%	9%	0.003



# Study Flow





### **Baseline Clinical Characteristics**

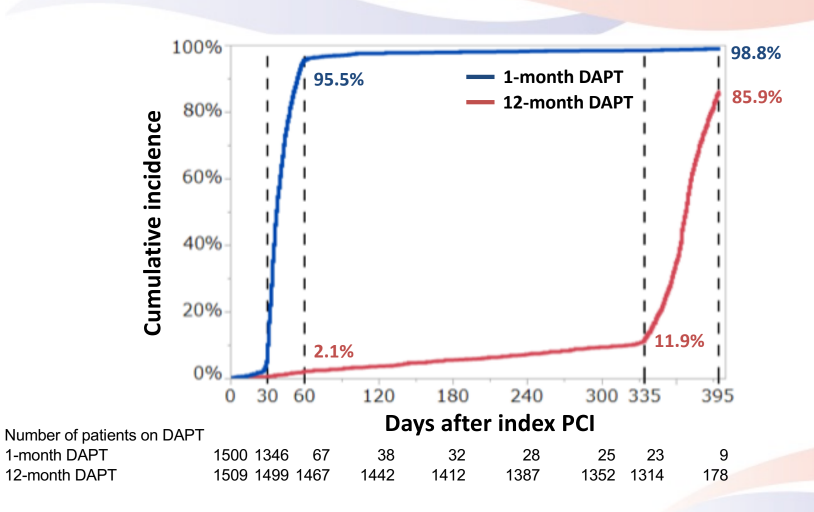
	1-month DAPT N=1500	12-month DAPT N=1509
Age, years	68.1±10.9	69.1±10.4
Men	79%	77%
ACS	38%	39%
STEMI	19%	18%
Stable CAD	62%	61%
Diabetes	39%	38%
Severe CKD (eGFR<30ml/min/m²)	6%	6%
Prior MI	14%	13%
Prior PCI	34%	35%
CREDO-Kyoto thrombotic risk score		
High; Intermediate; Low	8%; 21%; 71%	8%; 24%; 68%
CREDO-Kyoto bleeding risk score		
High; Intermediate; Low	7%; 27%; 66%	7%; 27%; 66%

#### STOPDAPT-2

### **Procedural Characteristics and Medications**

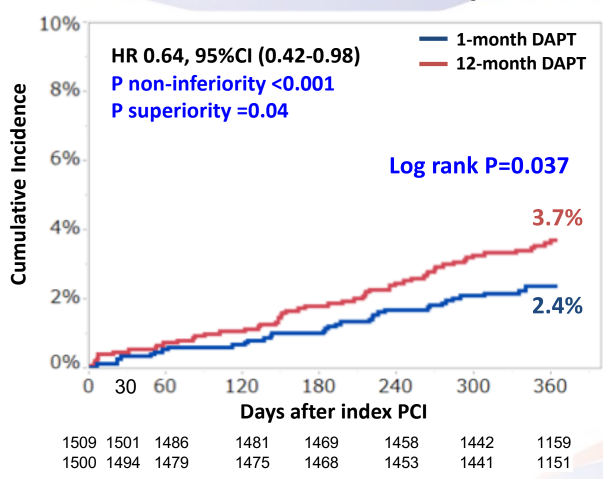
	1-month DAPT N=1500	12-month DAPT N=1509
Transradial approach	82%	84%
N of target lesions	$1.12 \pm 0.35$	$1.14 \pm 0.39$
Minimal stent diameter, mm	$2.98 \pm 0.49$	2.96 ± 0.48
Total stent length, mm	$30.3 \pm 16.7$	$30.5 \pm 16.8$
SYNTAX Score	8 (5-14)	9 (6-15)
Target of LMCA	3%	3%
СТО	4%	4%
IVUS or OCT	97%	98%
ASA	99.8%	100%
Clopidogrel	60%	63%
Prasugrel (3.75mg/day)	40%	37%
Statin	88%	87%
PPI	79%	79%

### Persistent DAPT discontinuation rate



# STOPPAPT-2 Primary Endpoint: Net clinical benefit

#### CV death/MI/ST/Stroke/TIMI major/minor bleeding



No. at risk

12-month DAPT

1-month DAPT

#### STOPDAPT-2

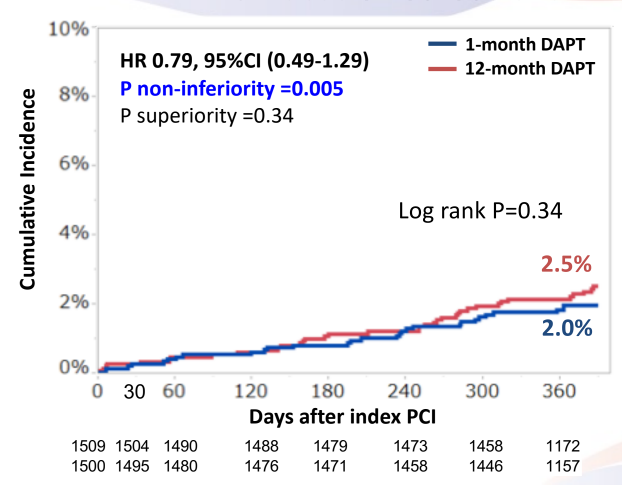
No. at risk

12-month DAPT

1-month DAPT

# Major secondary ischemic endpoint

CV death/MI/ST/Stroke





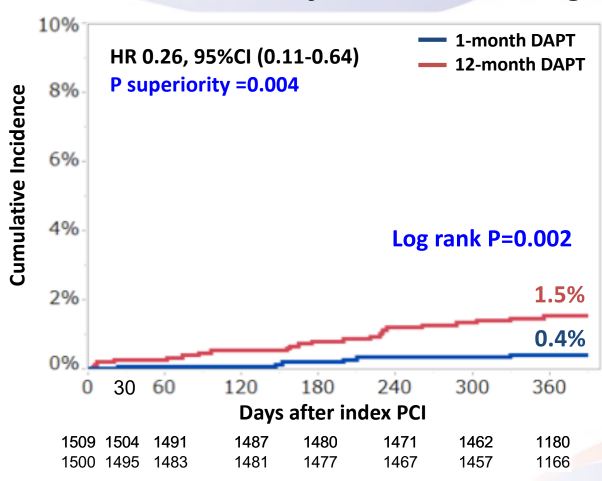
No. at risk

12-month DAPT

1-month DAPT

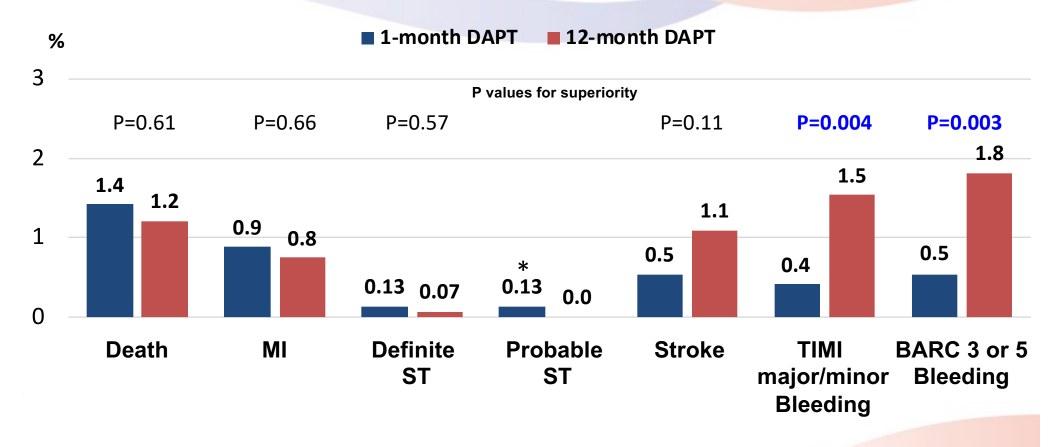
# Major secondary bleeding endpoint

#### TIMI major/minor bleeding



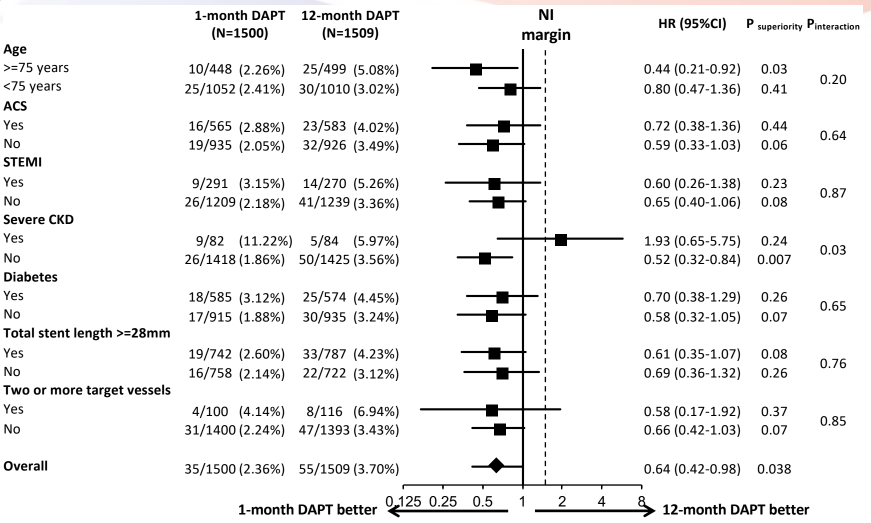
### STOPDAPT-2

# Clinical Outcomes at 1 year

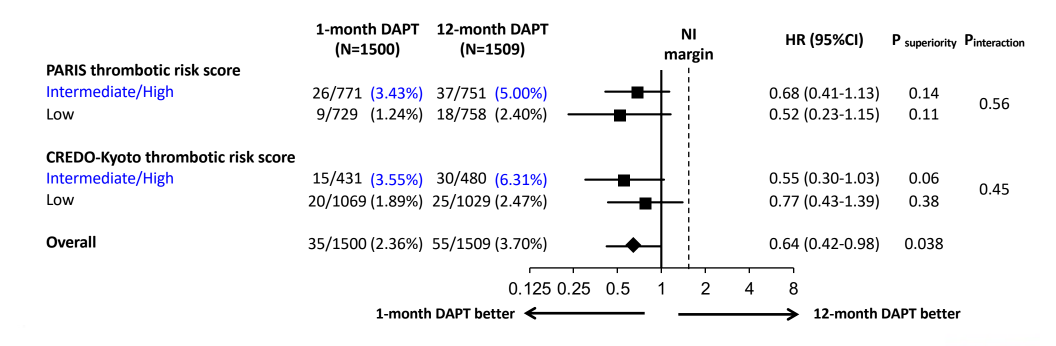


<sup>\* 2</sup> cases of probable ST (undefined death) in the 1-month DAPT group occurred before discontinuing DAPT at 1-month

## STOPDAPT-2 Subgroup analysis for the primary endpoint (1)



# Subgroup analysis for the primary endpoint (2)





## Limitations

- Lack of consensus on the use of the NACE as primary endpoint
- Open label design with its inherent limitations
- Limited enrollment of high ischemic risk patients
- Lower ischemic risk of Japanese versus US/European CAD patients
- Ticagrelor / Prasugrel (standard dose) not available in Japan
- No assessment of aspirin monotherapy after 1-month DAPT



# Conclusions

One-month DAPT followed by clopidogrel monotherapy provided a net clinical benefit for ischemic and bleeding events over 12-month DAPT with aspirin and clopidogrel after CoCr-EES implantation.

The benefit was driven by significant reduction in bleeding events without increase in ischemic events.