

Randomized Comparison of Provisional Side Branch Stenting versus a Two-stent Strategy for treatment of True Coronary Bifurcation Lesions Involving a Large Side Branch.

#### **The Nordic-Baltic Bifurcation Study IV**

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#### **Disclosure Statement of Financial Interest**

I, *Indulis Kumsars,* DO NOT have a financial interest/arrangement or affiliation with one or more organizations that could be perceived as a real or apparent conflict of interest in the context of the subject of this presentation.

The Nordic-Baltic Bifurcation IV Study was an academic study primarily funded by participating hospitals.

The participating institutions received unrestricted study grants form Cordis and Abbot.









## **Nordic-Baltic Bifurcation Study IV Participating Centers**

#### Denmark

Aarhus University Hospital Aalborg University Hospital **Odense University Hospital** Rigshospitalet Copenhagen

#### Latvia

P.Stradins University Hospital, Riga (159 pts)

#### Sweden

**Örebro Hospital** (11 pts) Linköping Karolinska University Hospital (1 pts)

112 pts)	
13 pts)	
10 pts)	
(3 pts)	

(3 pts)

Turku University Hospital Kuopio University Hospital

#### Norway

Finland

Tromsø University Hospital (18pts) Arendal Hospital (3 pts)Feiring Heart Clinic (2 pts)

#### Lithuania

Vilnius University Hospital

Oulu University Hospital

Tampere University Hospital

(21 pts)

(75 pts)

(8 pts)

(6 pts)

(2 pt)









# Background

 Provisional (simple) stenting is the preferred strategy in treatment of most bifurcation lesions

 It is unknown if this also applies to true bifurcation lesions involving a large side branch









# Aim

 To compare provisional stenting and twostent techniques for the treatment of true coronary bifurcation lesions involving a large side branch









# Hypothesis

 Two stent techniques are superior to provisional stenting in treatment of true coronary bifurcation lesions involving a large side branch









# Methods

- Open label, randomized, multicenter trial
- 1:1 randomization
- Clinical FU at 0, 1 and 6 months
- Angiographic substudy with 8 months FU
- Study stents:
  - Sirolimus eluting Cordis Cypher Select+ (first 225 patients)
  - Everolimus eluting Abbott Xience V (last 225 patients)









# **Primary endpoint**

Combined endpoint after 6 months:

- cardiac death
- non-index procedure related myocardial infarction
- TLR
- definite stent thrombosis









# Secondary endpoints

- Individual endpoints of:
  - Total death
  - Cardiac death
  - Non-index procedure related MI
  - Target lesion revascularization (TLR)
  - Target vessel revascularization (TVR)
  - Definite stent thrombosis
- Procedure related myocardial infarction
- 8-month angiographic follow-up results









# Methods

## Inclusion criteria

- •Age≥18
- •Stable Angina, UAP, NSTEMI
- •MV≥3.0mm
- •SB ≥2.75mm
- Bifurcation stenosis involving both MV and SB (≥50%DS by eyeballing)

Exclusion criteria

- •STEMI
- •Cardiogenic shock
- •Other critical illness
- •Relevant allergies
- •Cr ≥ 200 µmol/L
- •SB lesion length >15mm









# Implantation techniques

#### Provisional SB stenting

- Two wires
- Predilatation
- MV stenting
- If TIMI flow<III or >75%DS in ostial SB: kissing balloon dilatation
- If SB TIMI flow <III after kissing balloon dilatation, SB stenting using a T- or Culotte technique









# Implantation techniques

- Two-stent techniques
  - Two wires
  - Predilatation of segments to be stented
  - Culotte stenting recommended
    - T-stenting and mini-crush allowed
  - Final kissing balloon dilatation









# Patient flowchart



\*numbers not balanced due to block randomization and sites with less than 4 inclusions









## **Baseline clinical characteristics**

	Provisional (n=221)	Two-stent (n=229)	р
Age (yrs)	64±12	63±11	ns
Diabetes (%)	16.3	15.3	ns
Active smoking (%)	19.1	21.1	ns
Statin treatment (%)	81.8	81.1	ns
Hypertension (%)	70.0	65.6	ns
Family history (%)	50.7	47.4	ns
History of PCI (%)	35.5	33.5	ns
History of CABG (%)	3.6	1.8	ns









## **Lesion characteristics**

	Provisional (n=221)	Two-stent (n=229)	р
LAD/diagonal (%)	74.1	76.7	ns
CX/obtuse marginal (%)	16.8	17.6	ns
RCA PDA/PLA (%)	6.4	4.0	ns
LM/LAD/CX (%)	2.7	1.3	ns
Ref. diameter main vessel (mm)*	3.5	3.4	0.04
Ref. diameter side branch (mm)*	2.9	2.9	ns
Lesion length SB (mm)*	7.4	8.0	<0.0001
Angulation > 60-70° (%)*	50.9	51.1	ns

\*visual estimation









## **Procedural data**

	Provisional (n=221)	Two-stent (n=229)	р
SB dilatation (%)	64.3	78.0	-
SB dilation or final kissing (%)	78.7	-	-
Final kissing balloon dilatation (%)	36.1	91.2	-
SB stented (%)	3.7	96.0	-
Culotte	-	65.6	-
T-stent	-	7.0	-
Other	-	26.4	-
Tx succesful* (%)	97.7	99.1	ns

\* (Residual stenosis <30% of MV + TIMI flow III in SB)









## **Procedural data**

	Provisional (n=221)	Two-stent (n=229)	р
Procedure time (min)	73.9	92.6	<0.0001
Contrast volume (mL)	187	238	<0.0001
Flouroscopy time (min)	14.0	22.8	<0.0001
Tx succesful* (%)	97.7	99.1	ns
Procedural CK-MB>5x UPL** (%)	3.0	3.1	ns
Procedural CK-MB>3x UPL** (%)	6.0	6.1	ns

\* Residual stenosis <30% of MV + TIMI flow III in SB</li>
 \*\* Assessment possible in 327 patients









## **Eventfree survival curve at 6 months**











# Individual endpoints at 6 months

	Provisional (n=220)	Two-stent (n=227)	р
Total death (%)	0	0.4	0.32
Cardiac death (%)	0	0	-
Non-procedural myocardial infarction (%)	1.8	0.9	0.50
Stent thrombosis (%)	0.9	0.4	0.54
Target lesion revascularization (%)	3.2	1.3	0.18
Target vessel revascularization (%)	3.7	1.3	0.11
Angina CCS class ≥ II	2.7	1.3	0.39







# Conclusion

- After 6 months, two-stent techniques for treatment of true bifurcation lesions with a large side branch showed no significant difference in MACE rate compared to provisional side branch stenting
- Longer and more complex procedures in the twostent group did not translate into more procedural myocardial infarctions
- Recommendations on optimal strategy for this lesion subset should await longer term follow-up





