	Overnight (reference)	Ambulatory			
	no./total no. (%)	no./total no. (%)	Hazard Ratio (95% CI)	p value	
Low ACEF					
MACE	7/618 (1.1)	18/2214 (0.8)	0.716 (0.299 - 1.714)	0.453	
Readmission	4/618 (0.6)	16/2214 (0.7)			
All-cause Death	1/618 (0.2)	0/2214 (0)			
MI	4/618 (0.6)	3/2214 (0.1)			
High ACEF					
MACE	35/1098 (3.2)	22/1002 (2.2)	0.683 (0.401 - 1.165)	0.162	
Readmission	14/1098 (1.3)	18/1002 (1.8)			
All-cause Death	20/1098 (1.8)	4/1002 (0.4)			
MI	4/1098 (0.4)	1/1002 (0.1)			
ACEF score: age/left ventricular ejection fraction + 1 (if creatinine ≥2 mg/dl)					

Conclusions: In this single-center registry, patients who underwent ambulatory PCI had no worse outcomes than those who stayed at least one night, at high and low ACEF scores.

#### TCT-656

# Low Incidence of Stent Thrombosis in Asian Races: Multicenter Registry in Asia 7 Years Follow-Up Result

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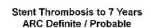
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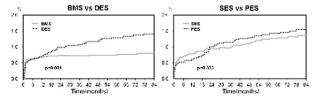
**Background:** The aim of this study was to evaluate the frequency, predictors and the clinical outcome of stent thrombosis after DES implantation and bare metal stent (BMS) implantation in Asian races.

**Methods:** A total of 14,577 consecutive patients who underwent successful DES implantation (8,809 patients, 62% of the lesion with Sirolimus-eluting stent: SES, 38% of the lesion with Paclitaxel-eluting stent: PES) and BMS implantation (5,768 patients) were included in this study. We evaluate the frequency, predictor of stent thrombosis.

**Results:** At a mean follow-up of 78.5±29.9 months in DES and 81.8±26.4 months in BMS. The cumulative incidence of stent thrombosis were subacute stent thrombosis (SAT): 0.5% with DES and 0.6% with BMS, late stent thrombosis (LAST): 0.18% with DES and 0.1% with BMS, very late stent thrombosis (VLAST): 0.18% per year with DES and no BMS. Independent predictors of stent thrombosis are bifurcation lesion (OR=1.90, 95% CI: 1.83 to 24.24, p=0.01) and ejection fraction (OR=0.90, 95% CI: 0.86 to 0.94, p=0.03). Only 0.2 % of the patients were died because of the myocardial infarction after stent thrombosis in both groups.

**Conclusions:** The incidence of stent thrombosis in Asian races is relatively low (0.5 % with DES and 0.6% with BMS of SAT, 0.18% increase per year with DES of late stent thrombosis) at mean follow-up to 7 years. Particular attention will need to be directed to this complication when the patients have bifurcation lesions or low ejection fraction.





### TCT-657

Multi Center, Prospective, Randomized, Single Blind, Consecutive Enrollment Evaluation Of Elixir DESyneTM Novolimus-Eluting Coronary Stent System With Durable Polymer To Endeavor Zotarolimus-Eluting Coronary Stent System: 3-Year Clinical and 9-Month Angiographic And IVUS Results: EXCELLA II Study

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**Background:** Aims: To evaluate safety and effectiveness of the Elixir DESyneTM Novolimus-Eluting Coronary Stent System (CSS) compared to the Endeavor Zotarolimus-Eluting CSS through assessment of clinical, angiographic, and IVUS endpoints.

Methods: 210 patients were randomized 2:1 either to the DESyne CSS loaded with 5mcg per mm of stent length of Novolimus, a sirolimus metabolite, eluted via a durable methacrylate polymer, or to the Endeavor CSS loaded with 10mcg per mm of stent length of Zotarolimus eluted via a durable phosphoryl choline polymer. All patients were analyzed for the primary endpoint of late lumen loss (LLL) assessed by QCA at 9 months. All patients also underwent evaluation for secondary endpoints which included a Device-orientated Composite Endpoint (DoCE) defined as: cardiac death, MI not clearly attributable to a non-intervention vessel, and clinically-indicated target lesion revascularization (TLR); clinically-indicated Target Vessel Revascularization (TVR); and stent thrombosis all evaluated at 1, 6, 9, and 12 months and annually through 5 years. Stents were also assessed for angiographic endpoints at 9 months including: in-stent and in-segment LLL. A subset of patients underwent IVUS evaluation including percent neointimal obstruction at 9 months. The study met the non-inferiority endpoint and also demonstrated superiority of the DESyne CSS as compared to control.

Results: Table 1 summarizes 9-month angiographic and IVUS results and clinical results through 2 years which trend lower for the DESyne stent.

Table 1: 9-month Angiographic, IVUS and Clinical Results

	DESyne	Endeavor	p-value		
Angiographic Results					
Baseline RVD (post- procedure)	2.84 ± 0.43	2.91 ± 0.38	0.2		
9-month angiographic/IVUS					
In-stent LLL	0.11 ± 0.32	0.63 ± 0.42	< 0.001		
% neointimal volume	4.5 ± 5.1	20.9 ± 11.3	<0.001		
Clinical Results					
12-month DoCE (%)	4.3	7.0	0.51		
Clinically-indicated TLR	1.4	5.6	0.18		
24-month DoCE (%)	4.3	9.0	0.14		
Clinically-indicated TLR	1.4	7.0	0.045		

**Conclusions:** The study met the non-inferiority endpoint and also demonstrated superiority of the DESyne CSS as compared to control. Clinical results through 3 years and a review of angiographic and IVUS results will be presented.

### TCT-658

## Do Drug Eluting Stents Improve Survival in All Comers?

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**Background:** Drug eluting stents (DES) have been shown to significantly decrease restenosis with subsequent need for lesion and/or vessel revascularization when compared with bare metal stents (BMS) in selected patient groups in both randomized controlled trials and in observational registries. If their use in all-comers is also associated with a survival benefit over a longer follow-up is controversial.

**Methods:** Retrospective analysis of the MIDAS registry for patients who underwent PCI with BMS between January 1 1997-December 31 1998 (pre DES era, group 1; N=