



STREAM

**STRATEGIC REPERFUSION EARLY AFTER
MYOCARDIAL INFARCTION**

Frans Van de Werf: Disclosures



- Study grant from Boehringer Ingelheim to perform the STREAM trial , paid to the University of Leuven ,Belgium
- Honoraria from Boehringer Ingelheim for membership of advisory board related to studies with dabigatran in patients with mechanical heart valves

BACKGROUND



- Large registries have demonstrated delays to primary PCI in STEMI patients first presenting to an EMS or a non-cath capable community hospital, requiring subsequent transfer for primary PCI.
- These delays may exceed guideline recommended times and result in a commensurate increase in morbidity and mortality worse.

AIM OF THE STUDY



To compare a strategy of early fibrinolysis followed by coronary angiography within 6-24 hours or rescue PCI if needed with routine primary PCI in STEMI patients presenting within 3 hours after onset of symptoms with at least 2 mm ST-elevation in 2 contiguous leads and who can not undergo primary PCI within 1 hour of first medical contact.

STUDY PROTOCOL

STEMI <3 hrs from onset symptoms, PPCI <60 min not possible, 2 mm ST-elevation in 2 leads

RANDOMIZATION 1:1 by IVRS, OPEN LABEL

Strategy A: bolus tenecteplase

Strategy B: primary PCI

Ambulance/ER

<75y: full dose

≥75y: ½ dose TNK

no lytic

Aspirin
Clopidogrel:
LD 300 mg + 75 mg QD
Enoxaparin:
30 mg IV + 1 mg/kg SC Q12h

Aspirin
Clopidogrel:
75 mg QD
Enoxaparin:
0.75 mg/kg SC Q12h

Antiplatelet and
antithrombin treatment
according to local standards

PCI Hospital

ECG at 90 min: ST resolution ≥ 50%

YES

NO

cor angio >6 to 24 hrs
PCI/CABG if indicated

immediate cor angio +
rescue PCI if indicated

Standard primary PCI

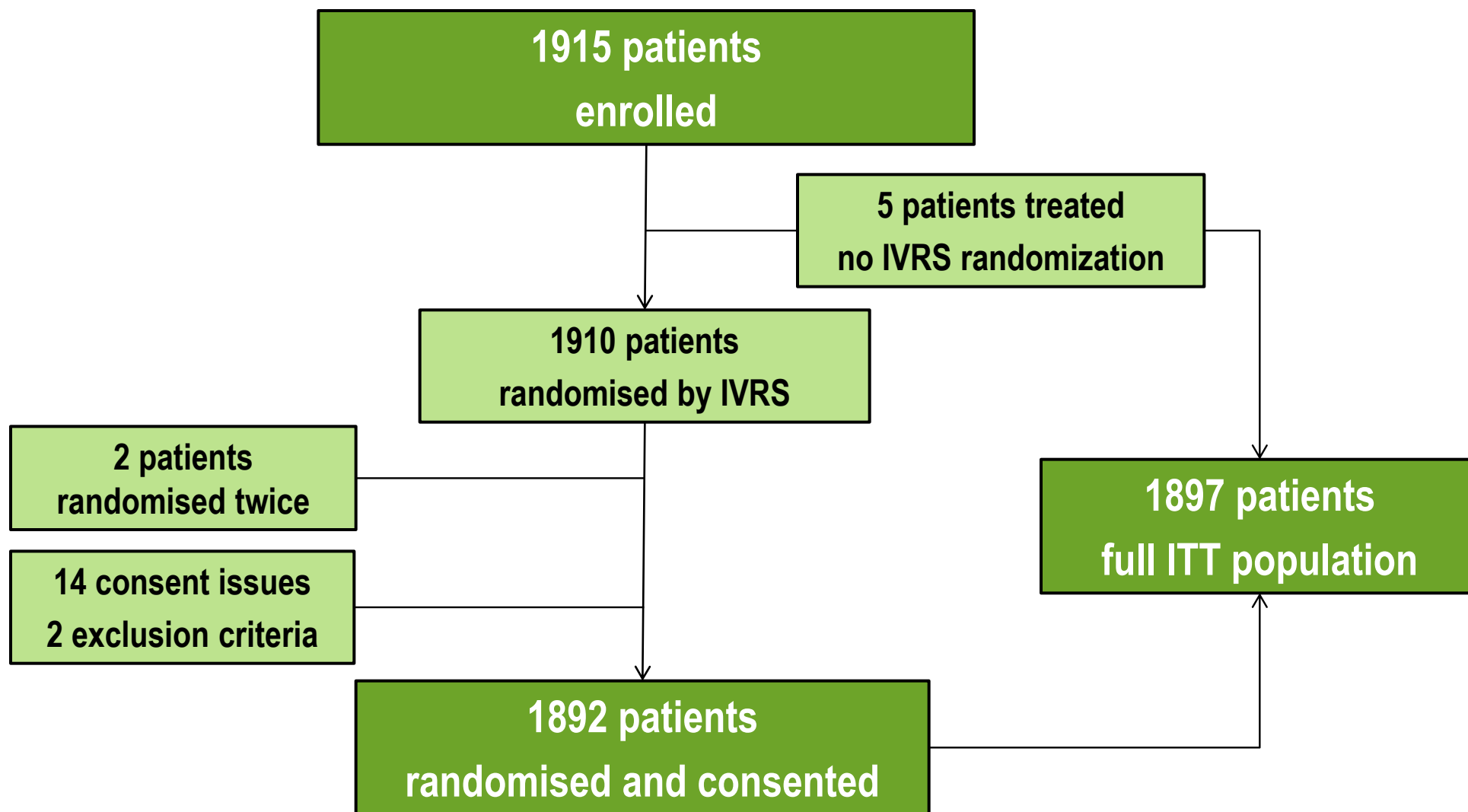
Primary endpoint: composite of all cause death or shock or CHF or reinfarction up to day 30

SAMPLE SIZE AND STATISTICAL ANALYSES



- Around 1000 patients per group was planned
- The rate of the primary endpoint in the primary PCI group was projected to be 15.0%
- There was no formal primary hypothesis
- All analyses are therefore explorative

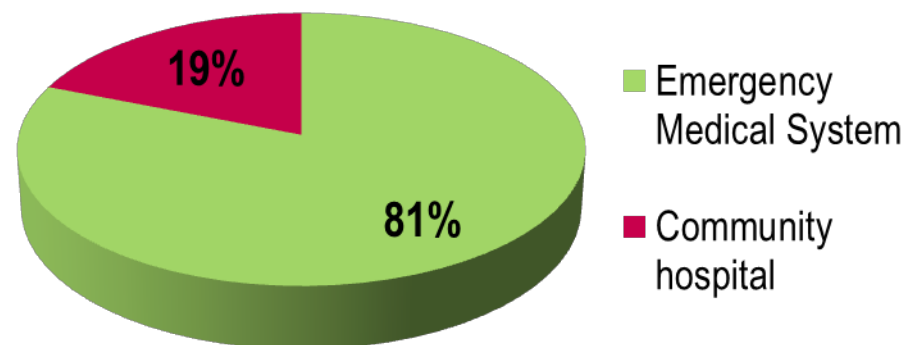
STREAM PATIENTS



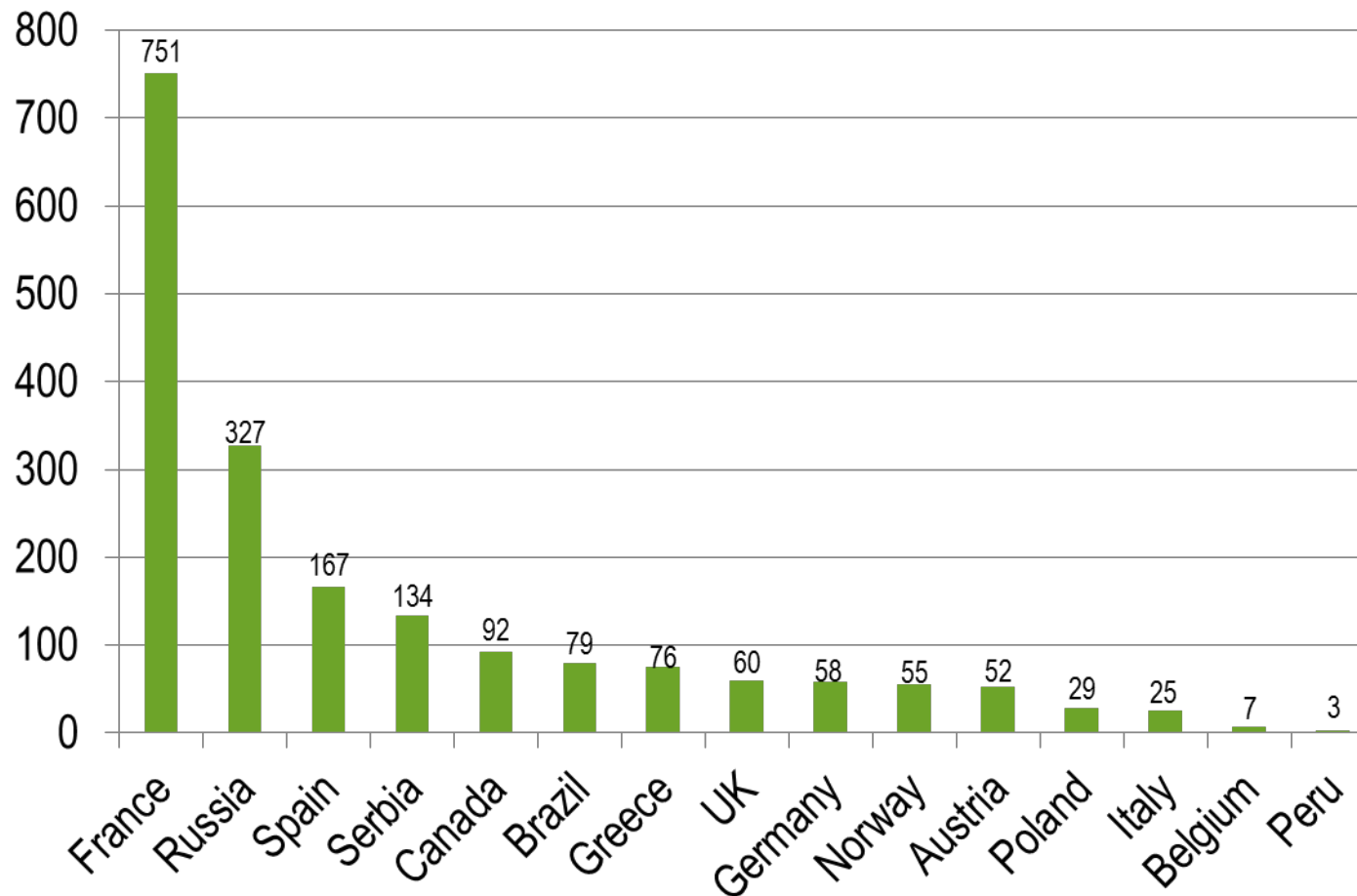
ENROLLMENT AND KEY DATES

- 1892 patients randomized by 99 sites in 15 countries
- First patient in: March 19, 2008
- Last patient in: July 26, 2012
- Last patient out: Sep 7, 2012

Enrolment setting



PATIENTS PER COUNTRY



BASELINE CHARACTERISTICS (1)

		Pharmaco-invasive (N=944)	PPCI (N=948)
Age (yrs)		59.7 (12.4)	59.6 (12.5)
Age >75 y (%)		134/944 (14%)	121/948 (13%)
Women (%)		194/944 (21%)	208/948 (22%)
Weight (kg)		80.5 (14.8)	80.0 (14.9)
Killip class (%)	I	842/895 (94%)	844/894 (94%)
	II/III	52/895 (6%)	47/894 (5%)
	IV	1/895 (<1%)	3/894 (<1%)
Heart rate (bpm)		74.9 (18.4)	75.5 (18.1)
Systolic BP (mmHg)		135.0 (22.7)	135.9 (23.3)
Infarct location	Anterior	453/942 (48%)	431/946 (46%)
	Inferior	468/942 (50%)	497/946 (53%)
	Other	21/942 (2%)	18/946 (2%)

Data are mean (SD) or number (%)

BASELINE CHARACTERISTICS (2)

%	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
Previous MI	81/940 (9%)	98/947 (10%)	0.20
Previous PCI	60/942 (6.37%)	83/944 (8.79%)	0.06
Previous CABG	2/944 (<1%)	3/946 (<1%)	>0.999
Previous congestive heart failure	3/939 (<1%)	16/945 (2%)	0.004
Hypertension	434/930 (47%)	414/932 (44%)	0.33
Diabetes	113/934 (12%)	123/939 (13%)	0.51

TIME DELAYS

Time difference (min)	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
Onset to first medical contact	62 (40,100)	61 (35,100)	0.36
Onset to randomisation	91 (68,132)	92 (65,132)	0.89
Onset to hospital admission	150 (110,202)	140 (100,185)	<0.001
Onset to start of reperfusion treatment (Tenecteplase or sheath insertion)	100 (75,143)	178 (135,230)	<0.001
Randomisation to arrival at cath lab	483 (135,1140)	67 (45,98)	<0.001
Randomisation to sheath insertion	492 (148,1157)	77 (57,112)	<0.001
Onset to arrival at cath lab	600 (245,1235)	170 (125,220)	<0.001

Time intervals are median (Q1, Q3)

TIME DELAYS

Time difference (hours)	Pharmaco-invasive (N=944)
Randomisation to sheath insertion	
36% required rescue/urgent PCI	2.2 hours (1.8, 2.7)
64% non urgent angiography	17 hours (11, 22)

Time intervals are median (Q1, Q3)

ANGIOGRAPHIC FINDINGS

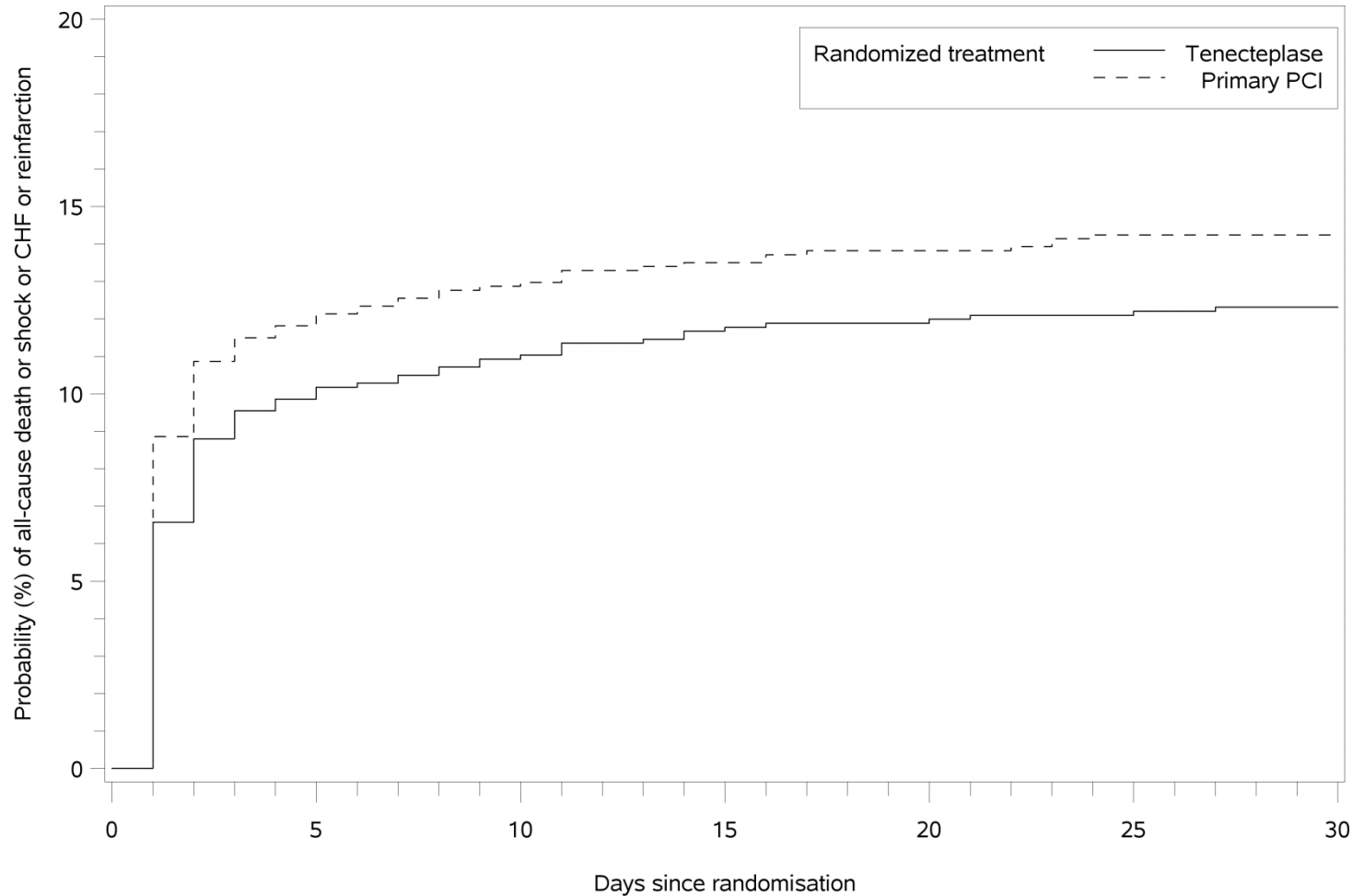
	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
TIMI flow before PCI			<0.001
TIMI 0	141/884 (16%)	534/900 (59%)	
TIMI 1	88/884 (10%)	91/900 (10%)	
TIMI 2	138/884 (16%)	89/900 (10%)	
TIMI 3	517/884 (58%)	186/900 (21%)	
TIMI flow after PCI			0.41
TIMI 0	18/819 (2%)	24/884 (3%)	
TIMI 1	12/819 (1%)	11/884 (1%)	
TIMI 2	43/819 (5%)	33/884 (4%)	
TIMI 3	746/819 (91%)	816/884 (92%)	
Urgent coronary angiography	331/911 (36.3%)		
PCI performed	736/915 (80%)	838/933 (90%)	<0.001
CABG performed	44/943 (4.7%)	20/947 (2.1%)	0.002
Stents deployed	704/736 (96%)	801/838 (96%)	0.95

PRIMARY ENDPOINT

%	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
All cause death or shock or reMI or CHF	116/939 (12.4%)	135/943 (14.3%)	0.21

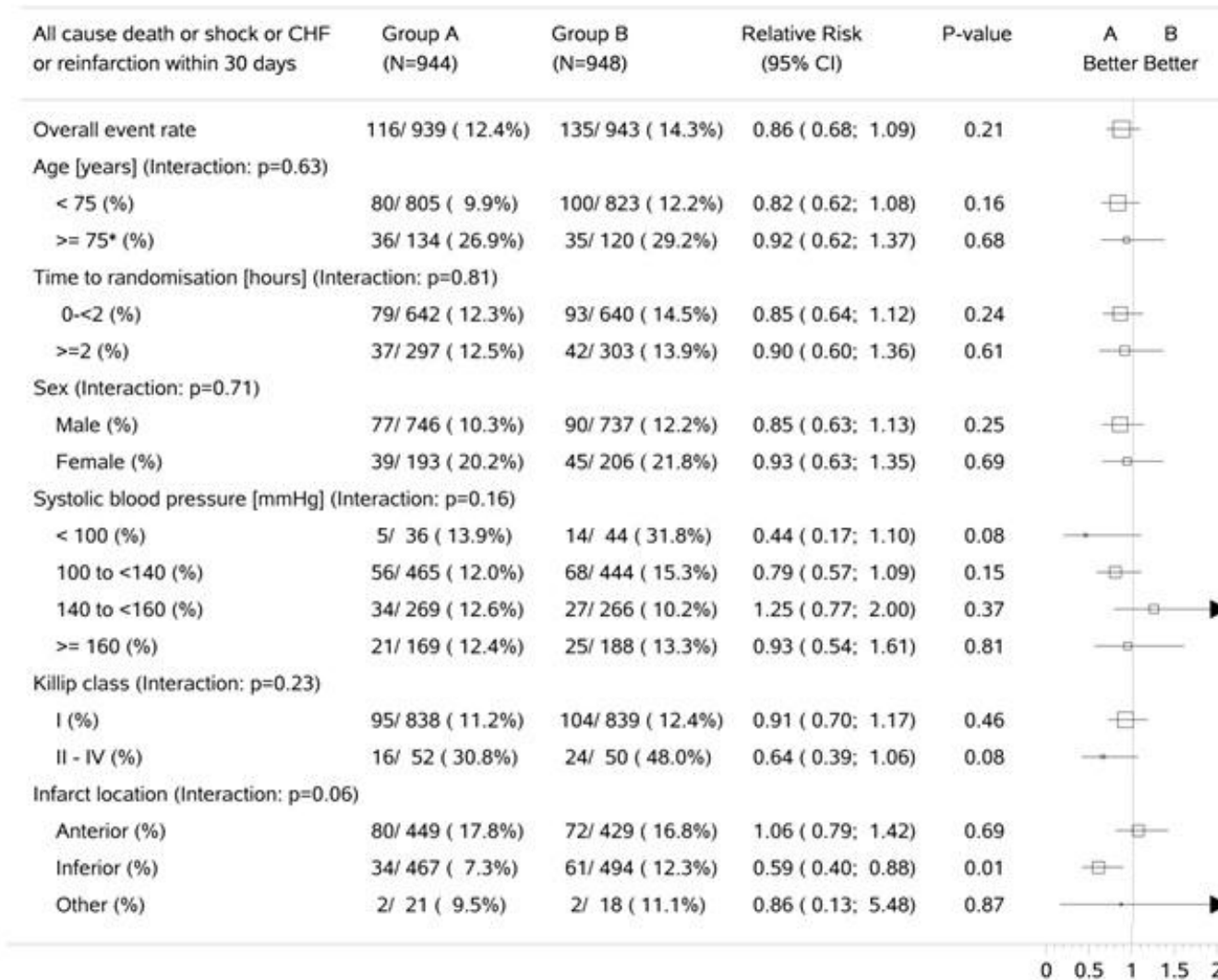
The 95 CI of the observed incidence in the pharmaco-invasive arm would exclude a 1.11% absolute or 9% relative excess compared with PPCI

KAPLAN-MEIER CURVES FOR PRIMARY ENDPOINT

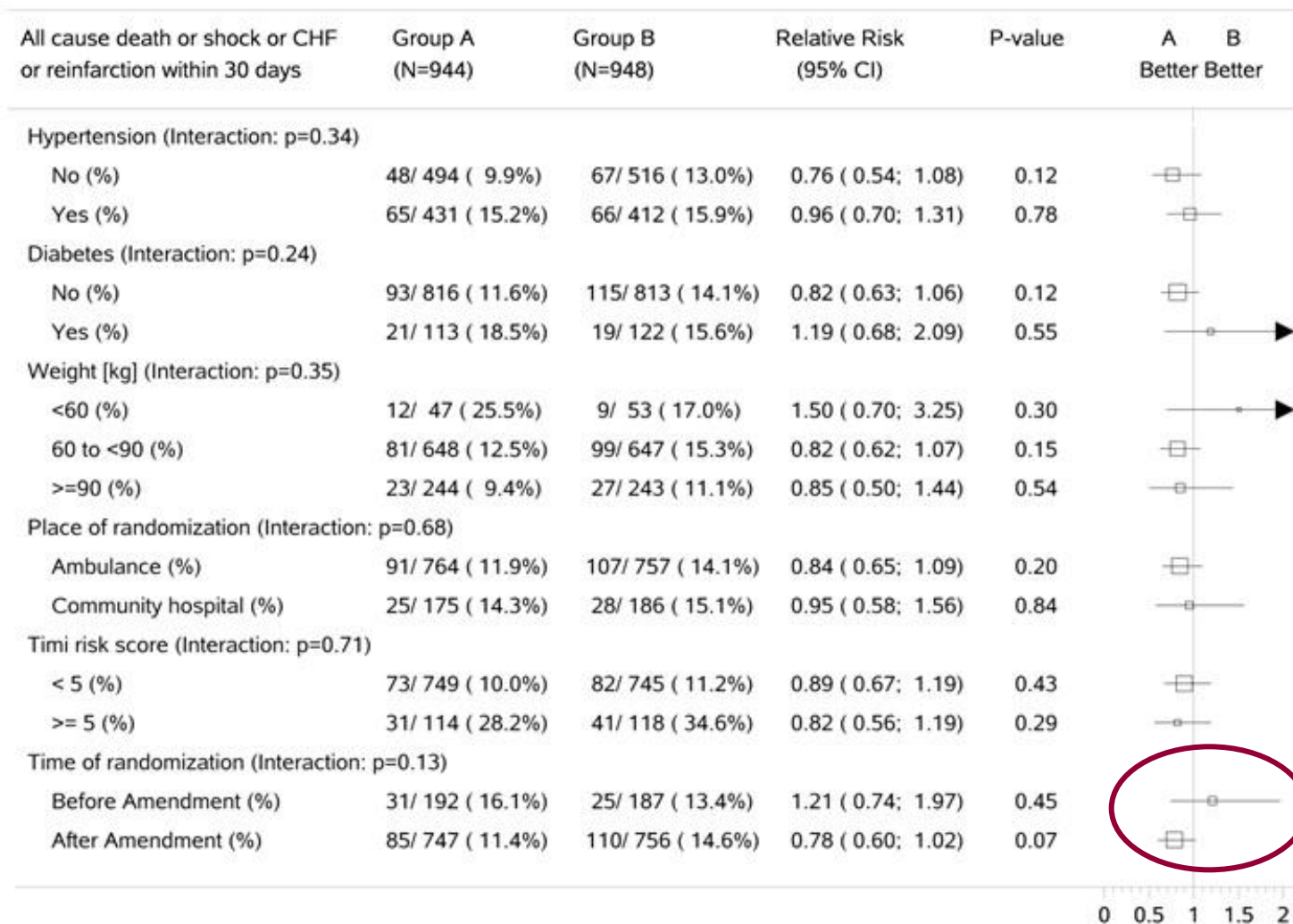


Number at risk							
Tenecteplase	943	848	837	829	827	825	823
Primary PCI	948	836	824	818	815	811	811

Subgroup analyses for primary endpoint within 30 days



Subgroup analyses for primary endpoint up to 30 days



SINGLE ENDPOINTS UP TO 30 DAYS

%	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
All death	43/939 (4.6%)	42/946 (4.4%)	0.88
Cardiac death	31/939 (3.3%)	32/946 (3.4%)	0.92
CHF	57/939 (6.1%)	72/943 (7.6%)	0.18
Cardiogenic shock	41/939 (4.4%)	56/944 (5.9%)	0.13
Reinfarction	23/938 (2.5%)	21/944 (2.2%)	0.74
Rehosp cardiac reason	45/939 (4.8%)	41/943 (4.3%)	0.64

STROKE RATES UP TO DAY 30

%	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
Total stroke (all types)	15/939 (1.6%)	5/946 (0.5%)	0.03
Intracranial haemorrhage	9/939 (1.0%)	2/946 (0.2%)	0.04
after amendment 2*:	4/747 (0.5%)	2/758 (0.3%)	0.45
Primary ischaemic stroke without haemorrhagic conversion	5/939 (0.5%)	3/946 (0.3%)	0.51

*Amendment 2 (Aug 2009): dose reduction of tenecteplase by 50% in patients 75 years of age or older

STROKE RATES UP TO DAY 30



	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
Total population			
Total stroke	15/939 (1.60%)	5/946 (0.53%)	0.03
Fatal stroke	7/939 (0.75%)	4/946 (0.42%)	0.39
ICH	9/939 (0.96%)	2/946 (0.21%)	0.04
Fatal ICH	6/939 (0.64%)	2/946 (0.21%)	0.18
Post amendment population(n=1505)			
Total stroke	9/747 (1.20%)	5/758 (0.66%)	0.30
Fatal stroke	3/747 (0.40%)	4/758 (0.53%)	>0.999
ICH	4/747 (0.54%)	2/758 (0.26%)	0.45
Fatal ICH	2/747 (0.27%)	2/758 (0.26%)	>0.999

IN-HOSPITAL BLEEDING COMPLICATIONS

%	Pharmaco-invasive (N=944)	PPCI (N=948)	P-value
Major non-ICH bleed	61/939 (6.5%)	45/944 (4.8%)	0.105
Minor non-ICH bleed	205/939 (21.8%)	191/944 (20.2%)	0.395
Blood transfusions	27/937 (2.9%)	22/943 (2.3%)	0.473

CONCLUSIONS

Fibrinolysis with bolus tenecteplase and contemporary antithrombotic therapy given before transport to a PCI-capable hospital coupled with timely coronary angiography

- **is as effective as primary PCI in STEMI patients presenting within 3 hours of symptom onset who cannot undergo primary PCI within one hour of first medical contact.**
- **is associated with a small increased risk of intracranial bleeding.**
- **provides the opportunity for a measured approach to invasive coronary interventions, circumventing an urgent procedure in about two thirds of fibrinolytic treated STEMI patients.**

ORIGINAL ARTICLE

Fibrinolysis or Primary PCI in ST-Segment Elevation Myocardial Infarction

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