

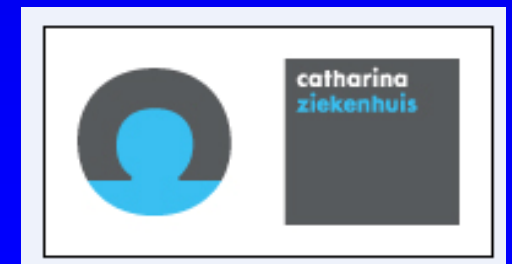
CORONARY PHYSIOLOGY IN THE CATHLAB:

STRUCTURE OF THE CORONARY CIRCULATION

***Educational Training Program ESC
European Heart House
april 24th - 26th 2014***



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Eindhoven, The Netherlands



Disclosures related to this ETP course:

- ***Dr Pijls received institutional research grants from St Jude Medical and Pharma Solutions***
- ***Dr Pijls is consultant to St Jude Medical, and to Heartflow***

ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- relation between vessel size and perfusion area
- endothelium and development of atherosclerosis
- the 2 or 3 compartment model of the coron circulation
- collaterals (*to be discussed tomorrow*)

SUPERIOR VENA CAVAL
BRANCH (NODAL ARTERY)

ANTERIOR R. ATRIAL
BRANCH OF
R. CORONARY
ARTERY

RIGHT
CORONARY
ARTERY

ANTERIOR
CARDIAC
VEINS

SMALL
CARDIAC VEIN

STERNOCOSTAL
ASPECT

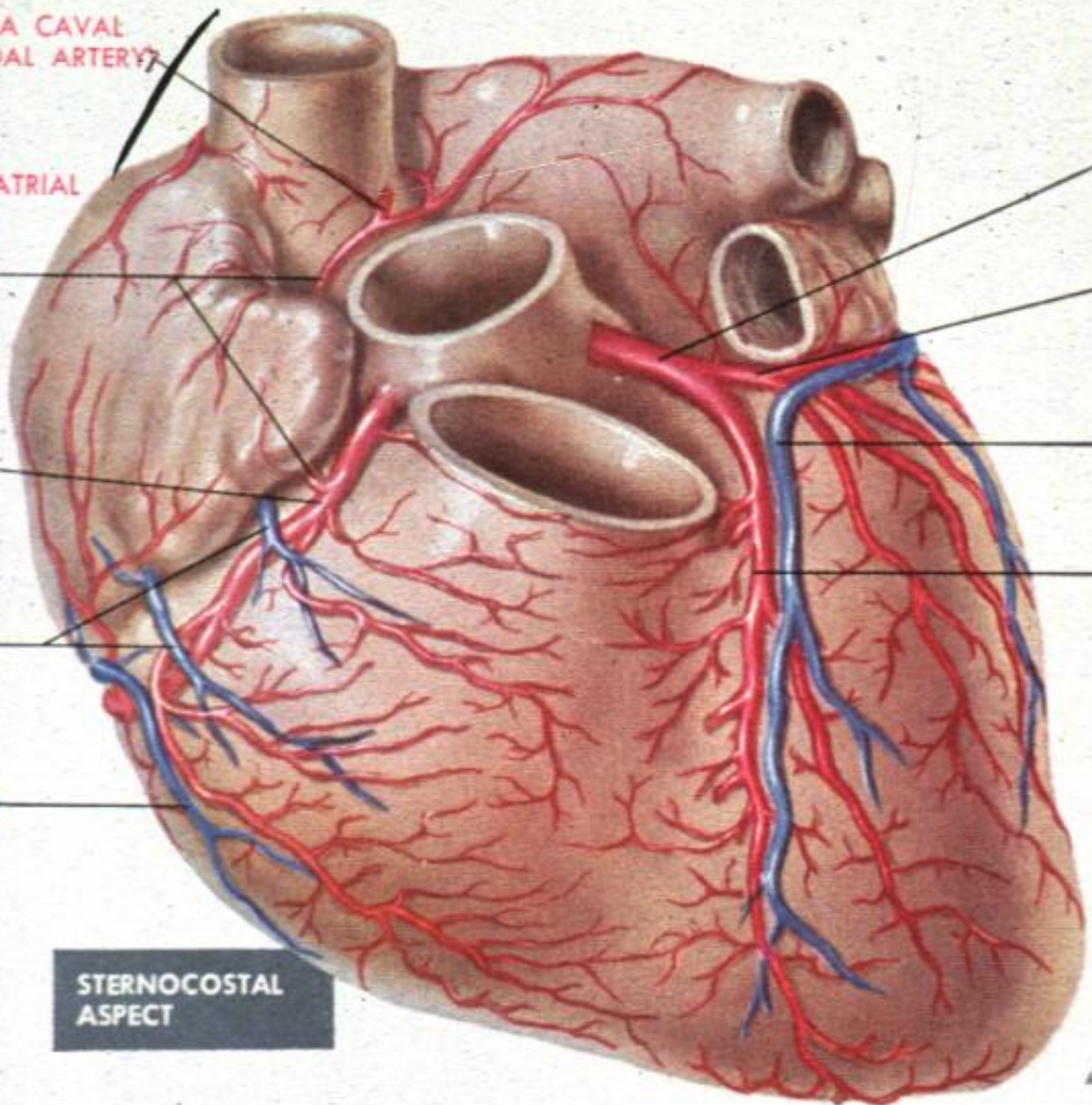
L. CORONARY ARTERY

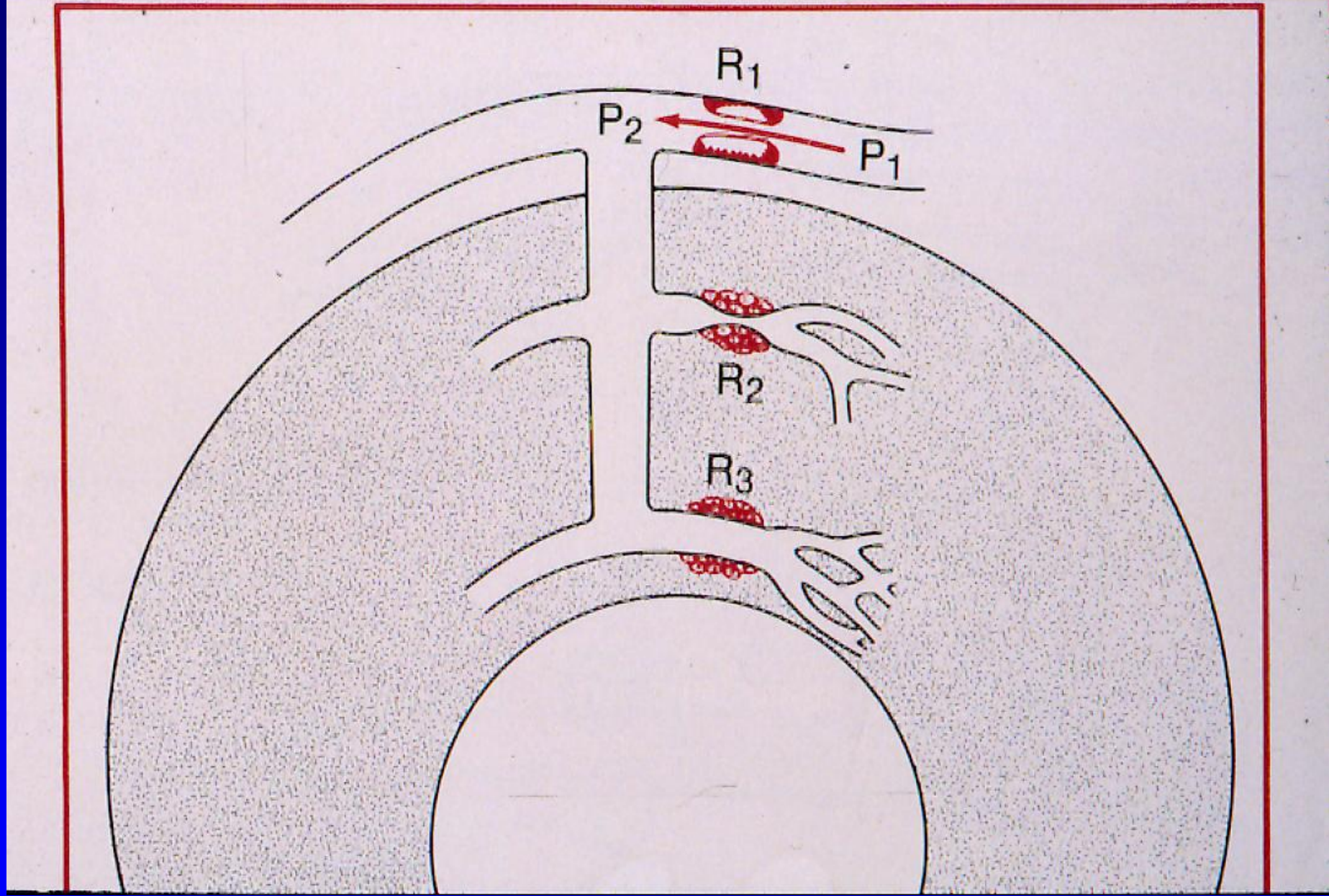
CIRCUMFLEX BRANCH OF
L. CORONARY ARTERY

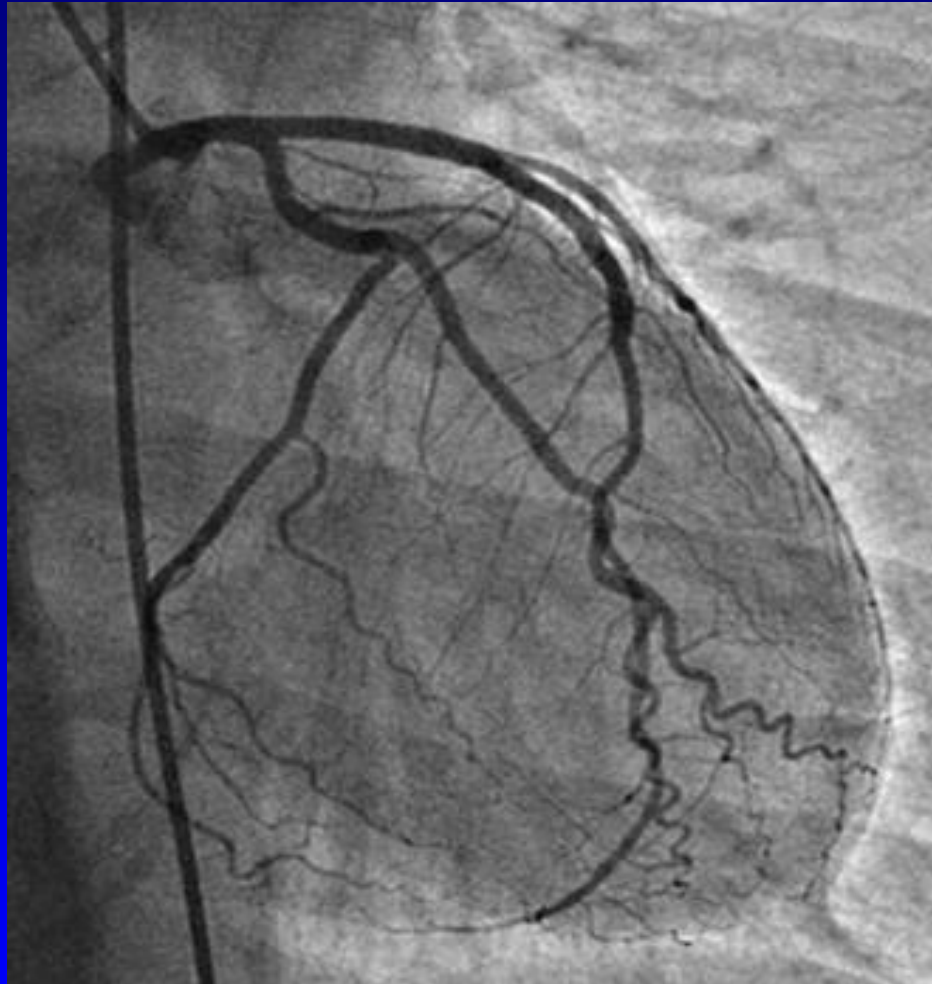
GREAT CARDIAC VEIN

ANTERIOR
INTERVENTRICULAR
(ANTERIOR/DESCENDING)
BRANCH OF L.
CORONARY ARTERY

F. Netter
M.D.
© CIBA

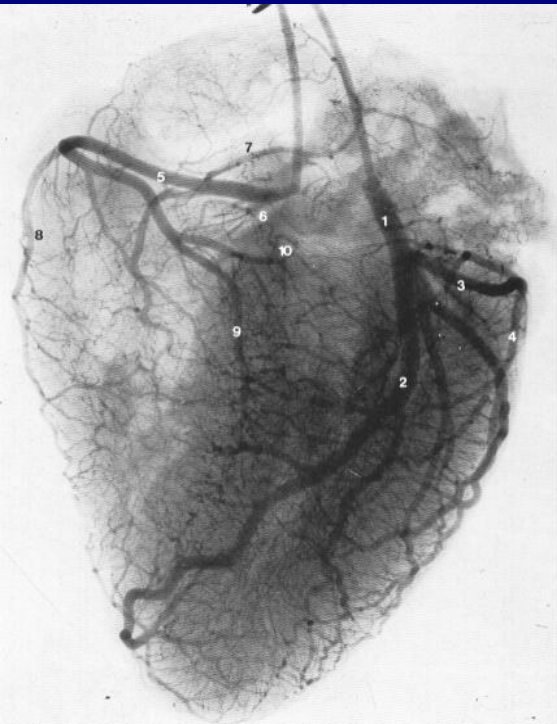






Let's have a closer look at the coronary tree.....

Fractale structure of the coronary circulation (Gould, Finet)



X 1

X 0.75

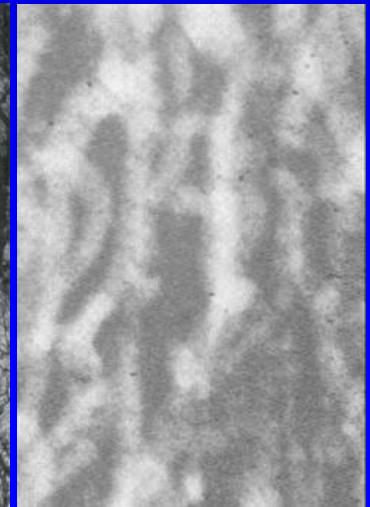


X 4.5

X 10.5

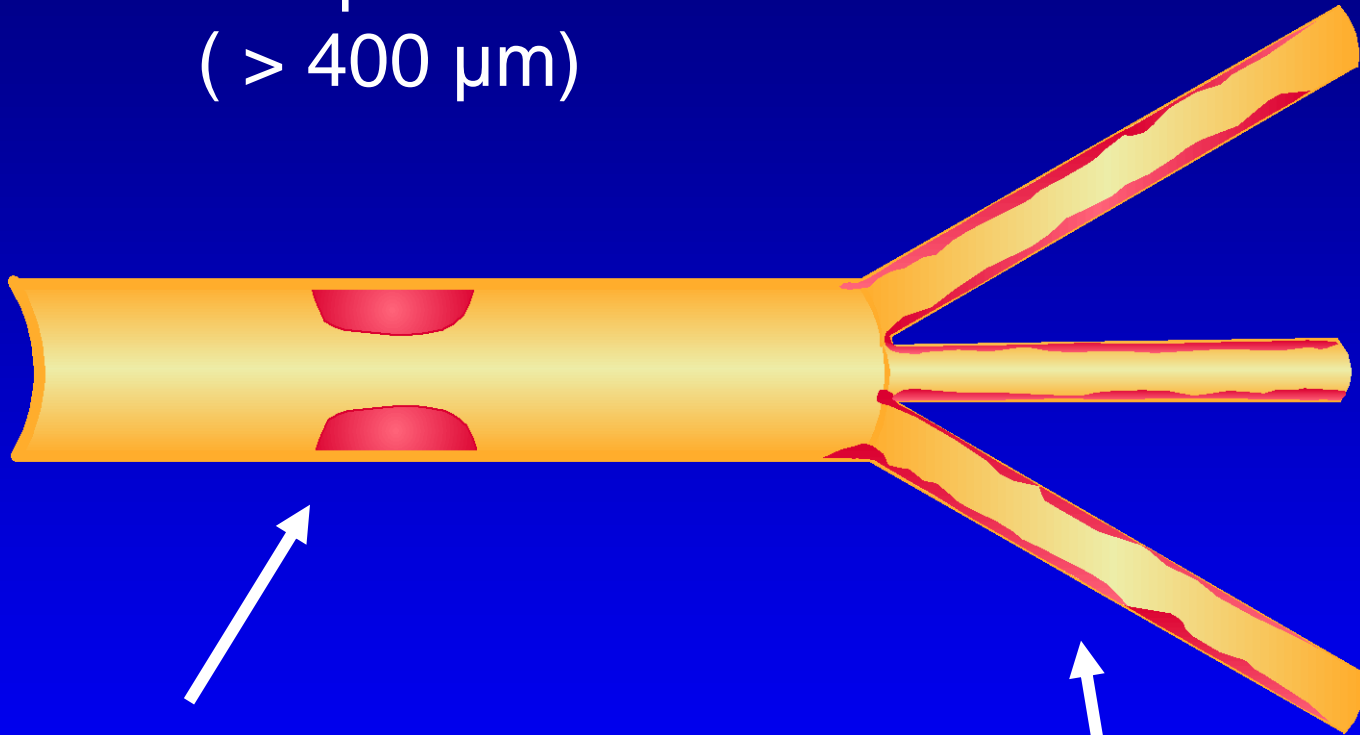
X 90

X 1100



epicardial
compartment
(> 400 μm)

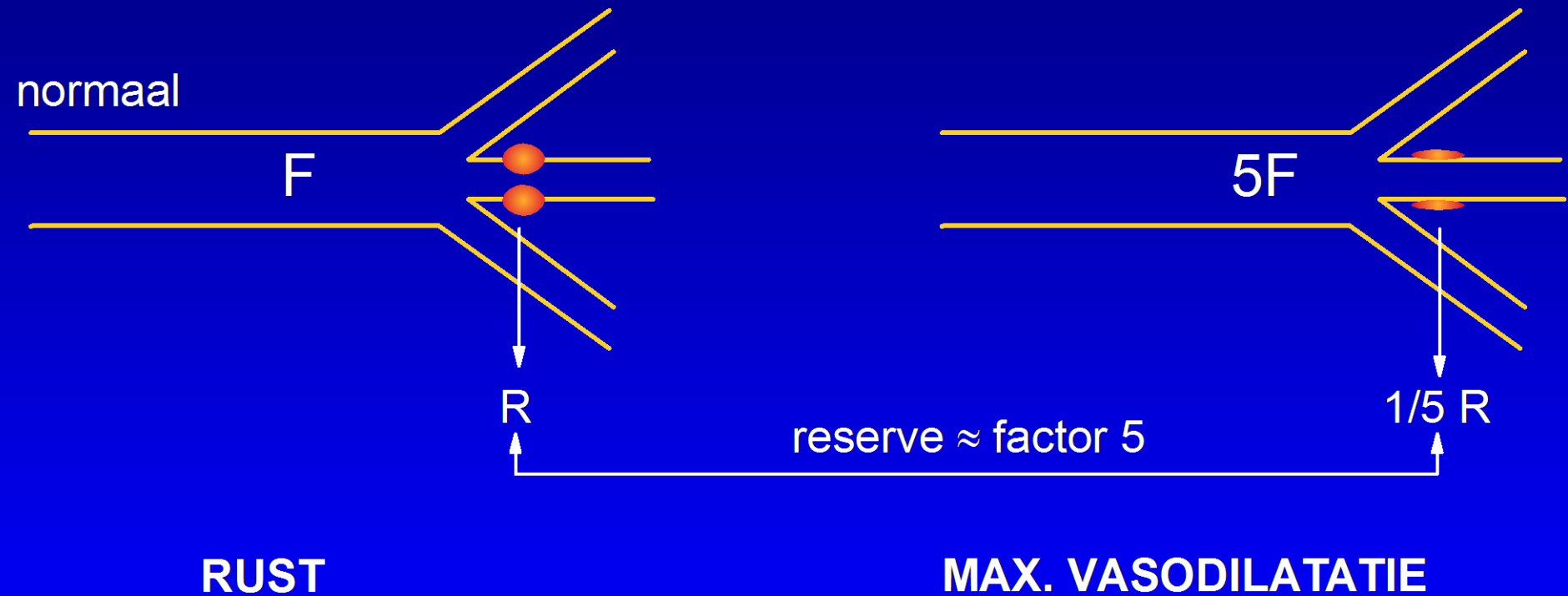
microvascular
compartment



traditionally visible by angiography
and more recently by many invasive
and non-invasive imaging methods

Black box
(until recently)

Regulation of coronary blood flow by arteriolar sphincters



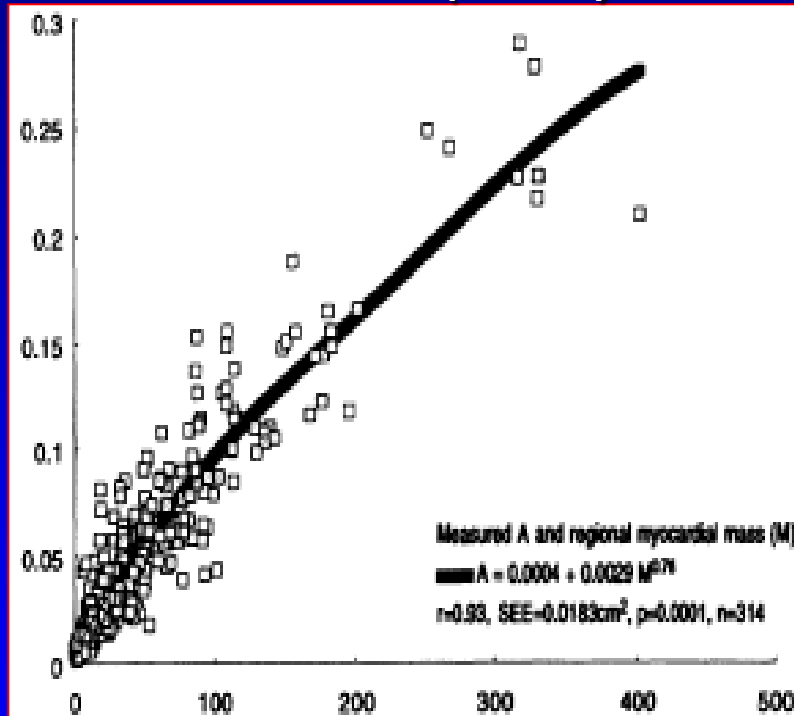
To be further discussed by Dirk Duncker

ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- ***relation between vessel size and perfusion area***
- endothelium and development of atherosclerosis
- the 2 or 3 compartment model of the coron circulation
- collaterals

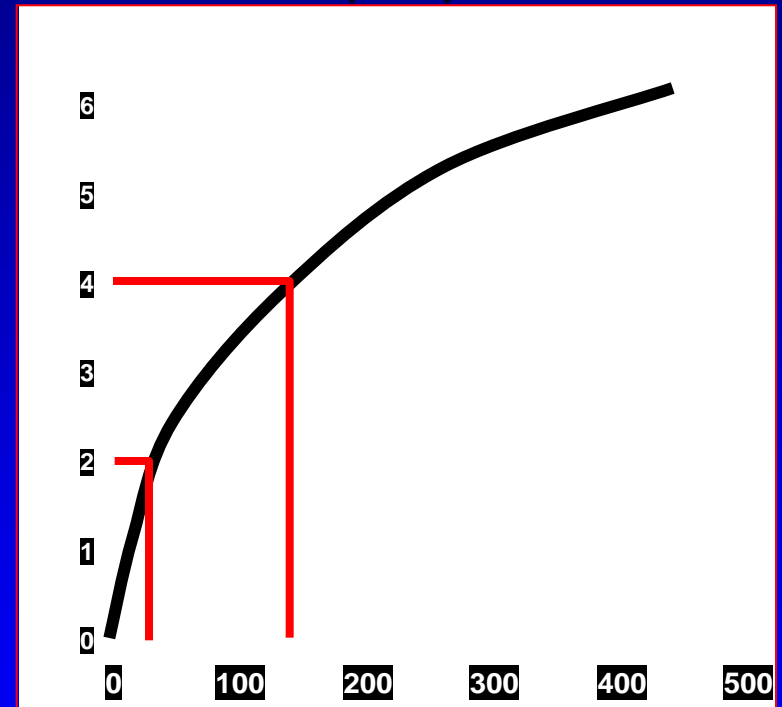
Relationship between vessel size and myocardial mass

Cross Sectional Area (~ flow)



Regional Myocardial Mass

Vessel Diameter (mm)



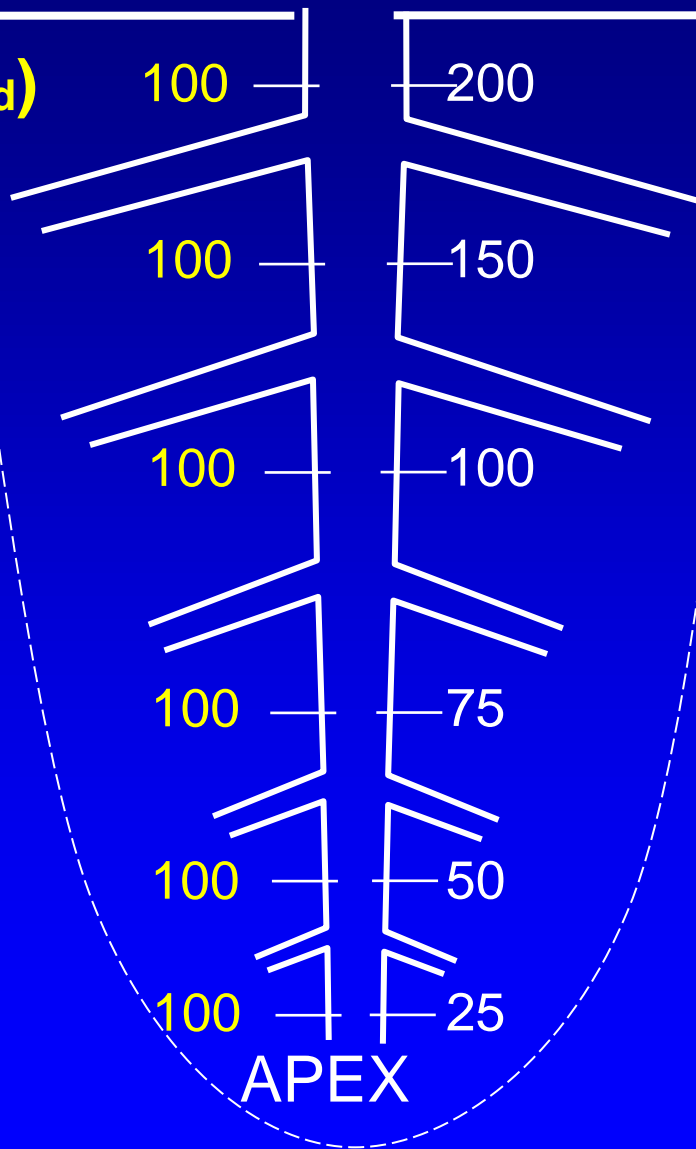
Regional Myocardial Mass (Grams)

AORTA
100 mmHg

IVUS-CSA

pressure (P_d)
(mm Hg)

flow (Q)
(ml/min)



9 mm²

7 mm²

5 mm²

3 mm²

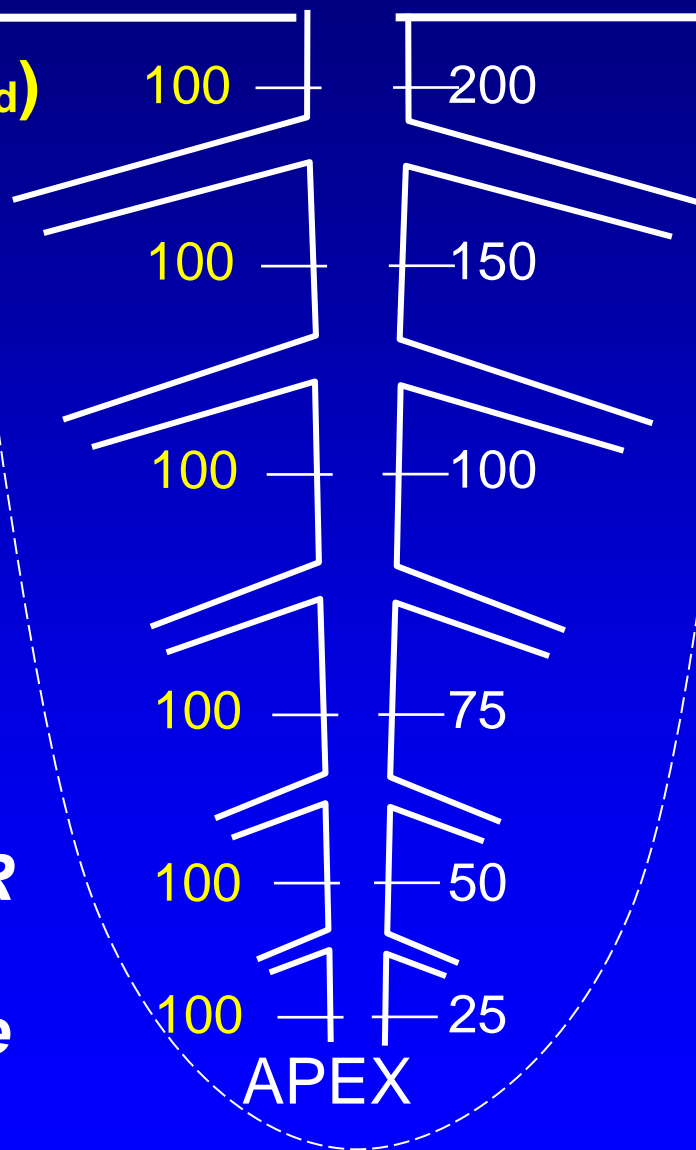
AORTA
100 mmHg

IVUS-CSA

pressure (P_d)
(mm Hg)

flow (Q)
(ml/min)

*Normal FFR
= 1.0
irrespective
where it is
measured*



9 mm²

7 mm²

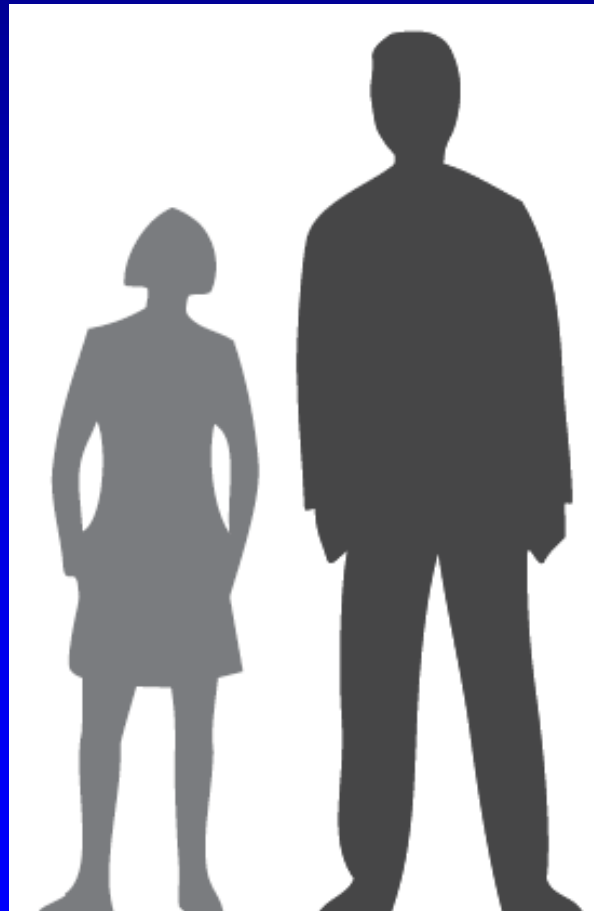
5 mm²

3 mm²

SIZE of the person (importance of perfusion territory)

Suppose both of these 2 persons have a proximal LAD stenosis

*FFR = 0.68
means exactly
the same in
both persons*



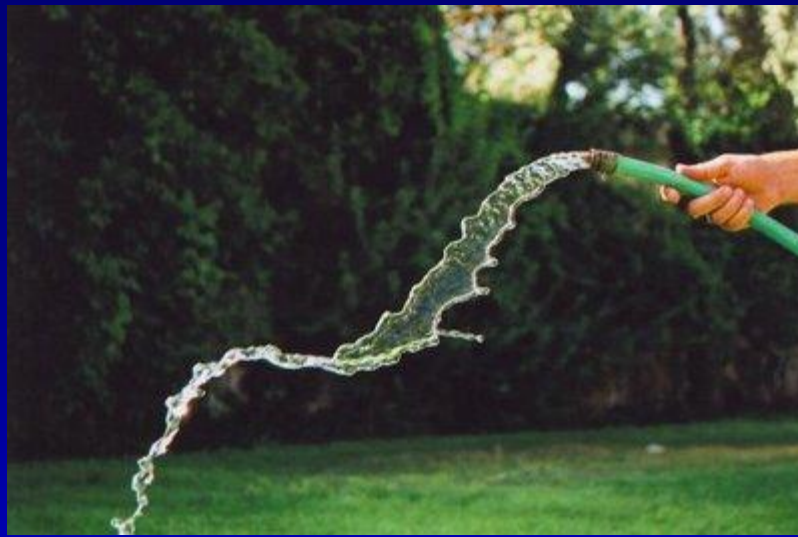
*CSA by IVUS
= 3.3 mm² has
a completely
different meaning
in both persons*

*LAD blood flow
= 100 ml/min has
a completely
different meaning
in both persons*

Value of ANY morphologic methodology (QCA, IVUS, OCT) to assess functional significance of a stenosis is limited by definition because there is simply no normal reference value

! We cannot understand the physiologic significance of a stenosis without taking into account the extent of the distal perfusion territory !

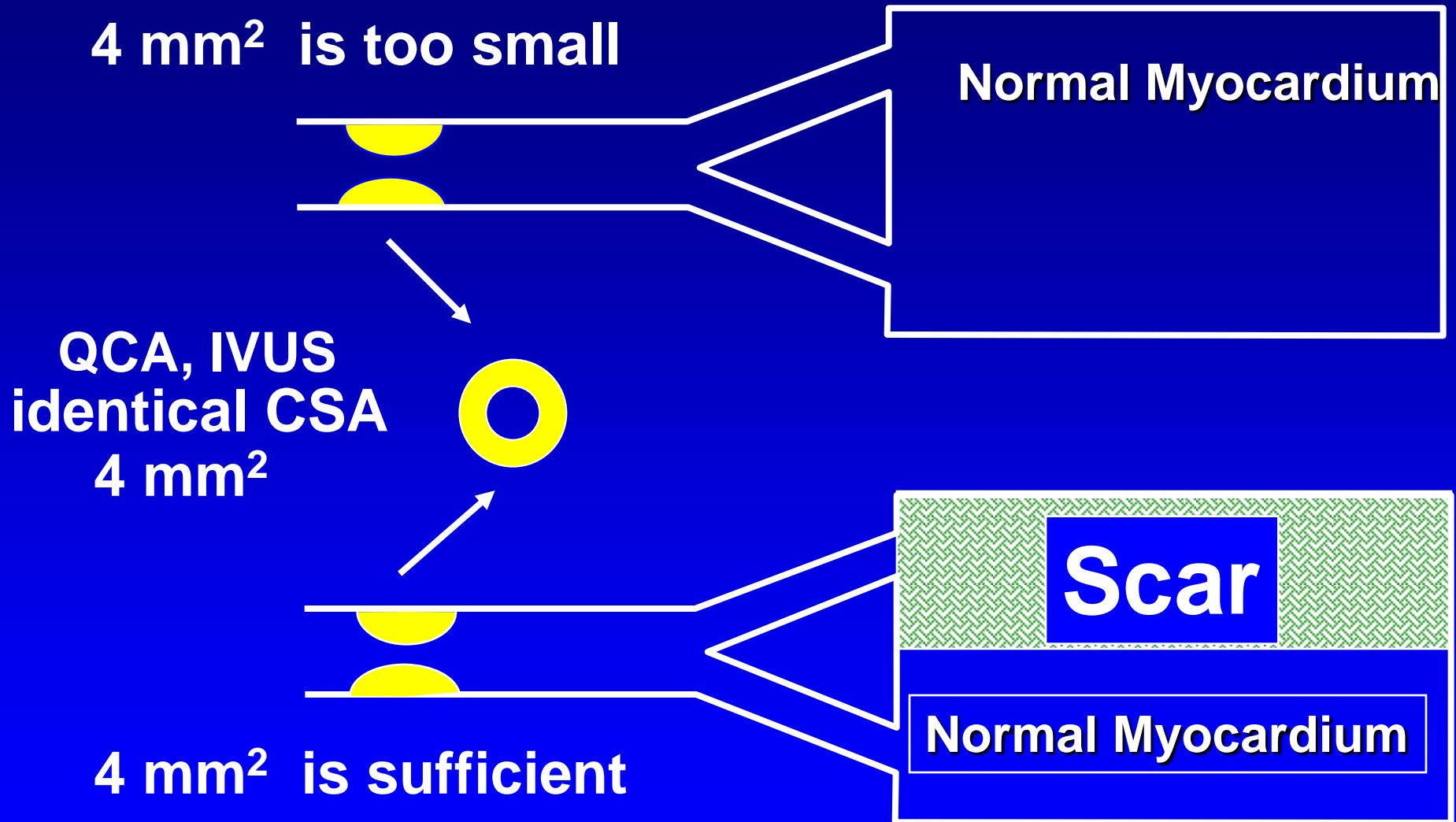
.....especially not under pathologic conditions, when the “physiologic match“ between vessel size and perfusion area has been lost



*With permission of
Dr Haitma Amin,
Bahrain*

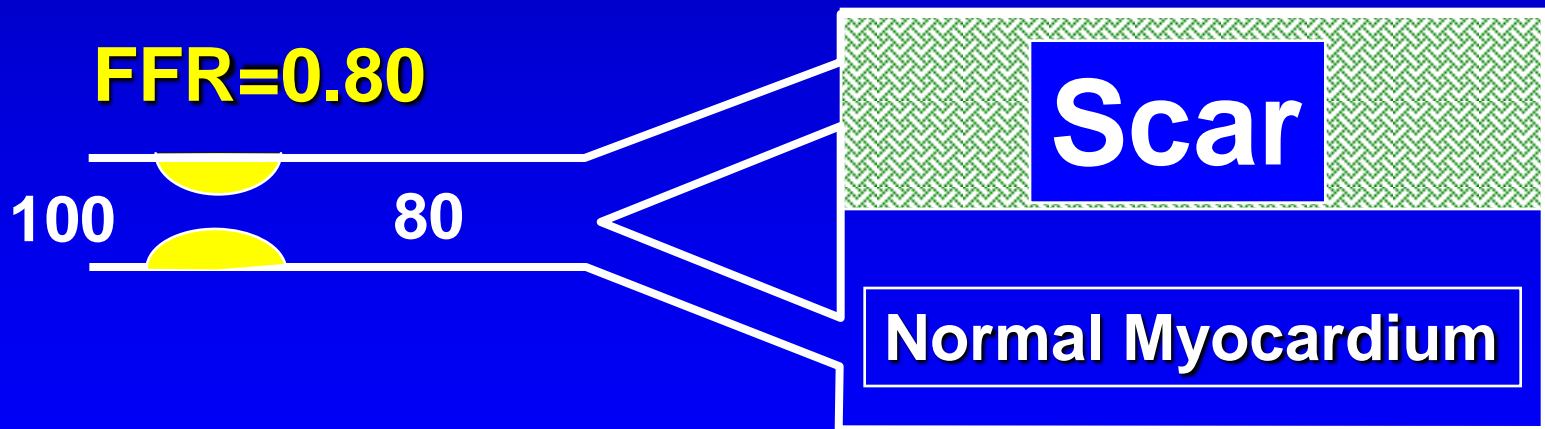
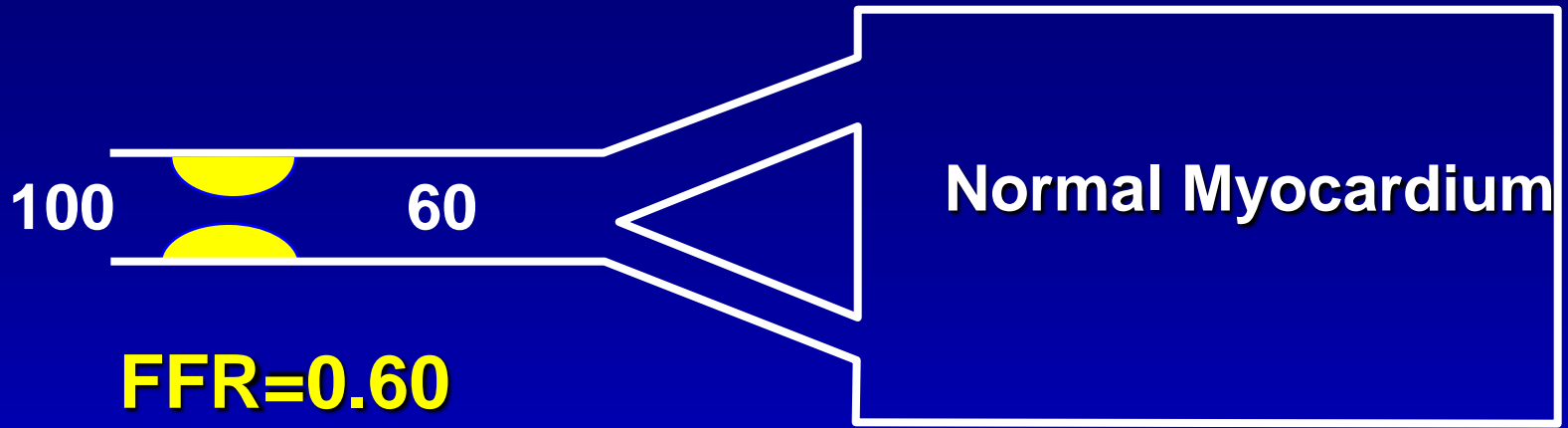


similar stenosis but different extent of perfusion area



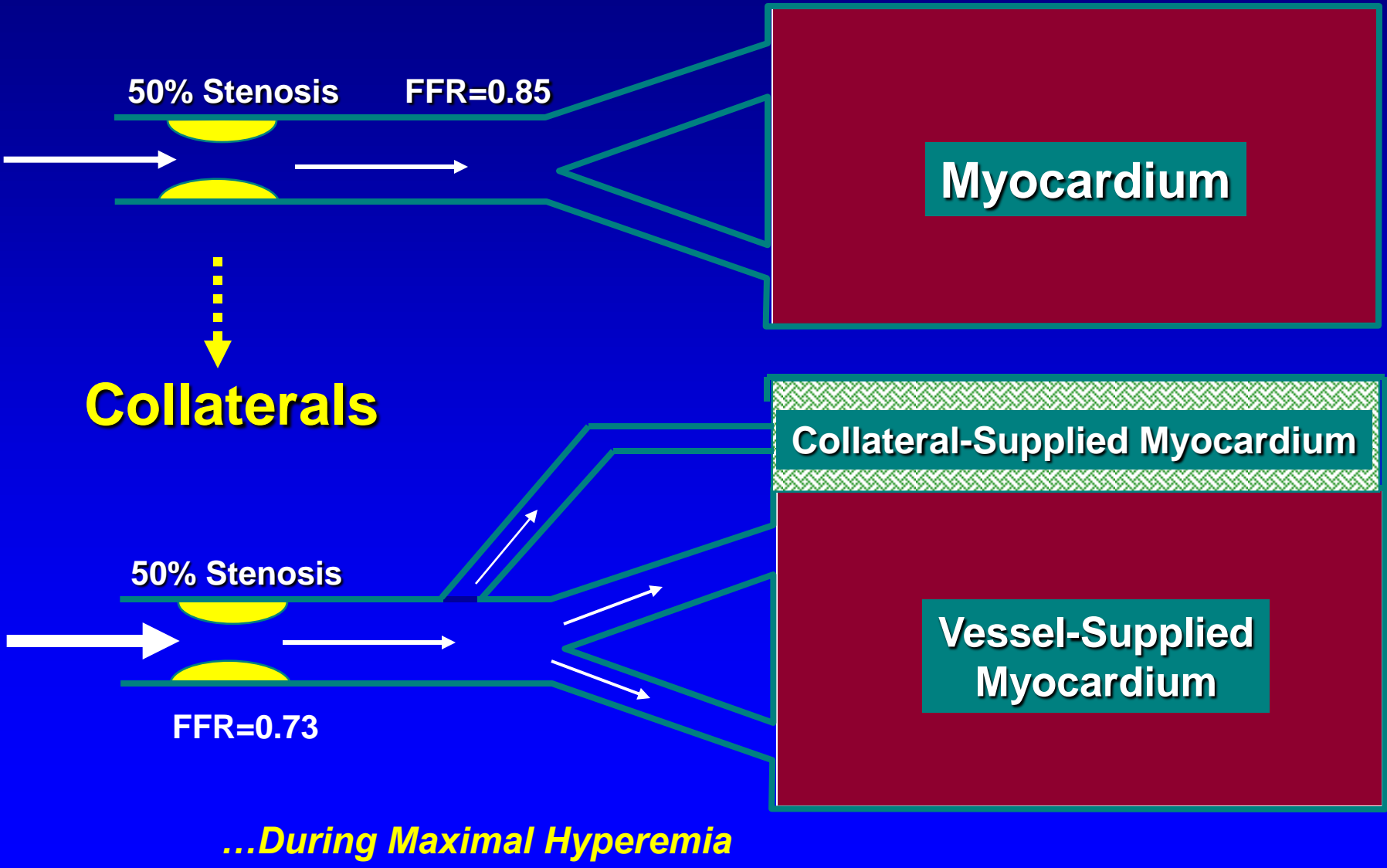
identical CSA, but different significance of stenosis

FFR accounts for the extent of the perfusion area:

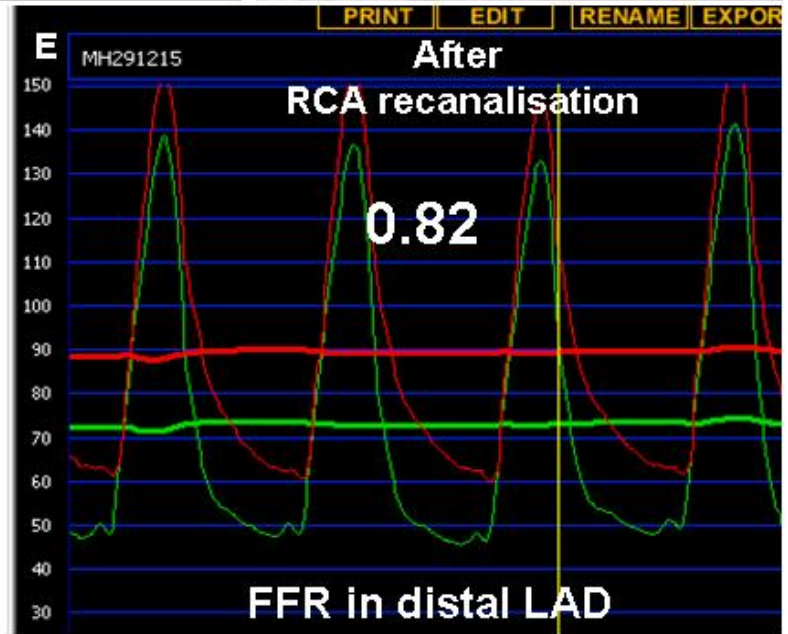
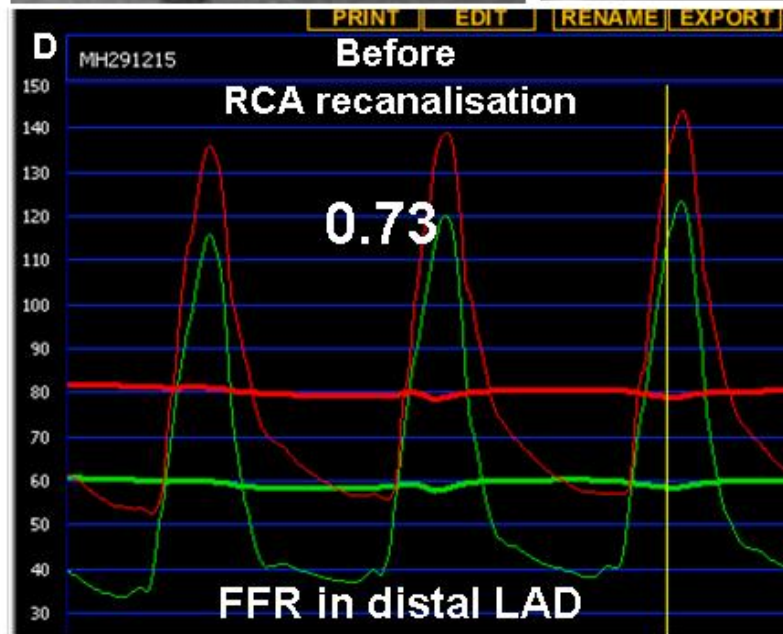
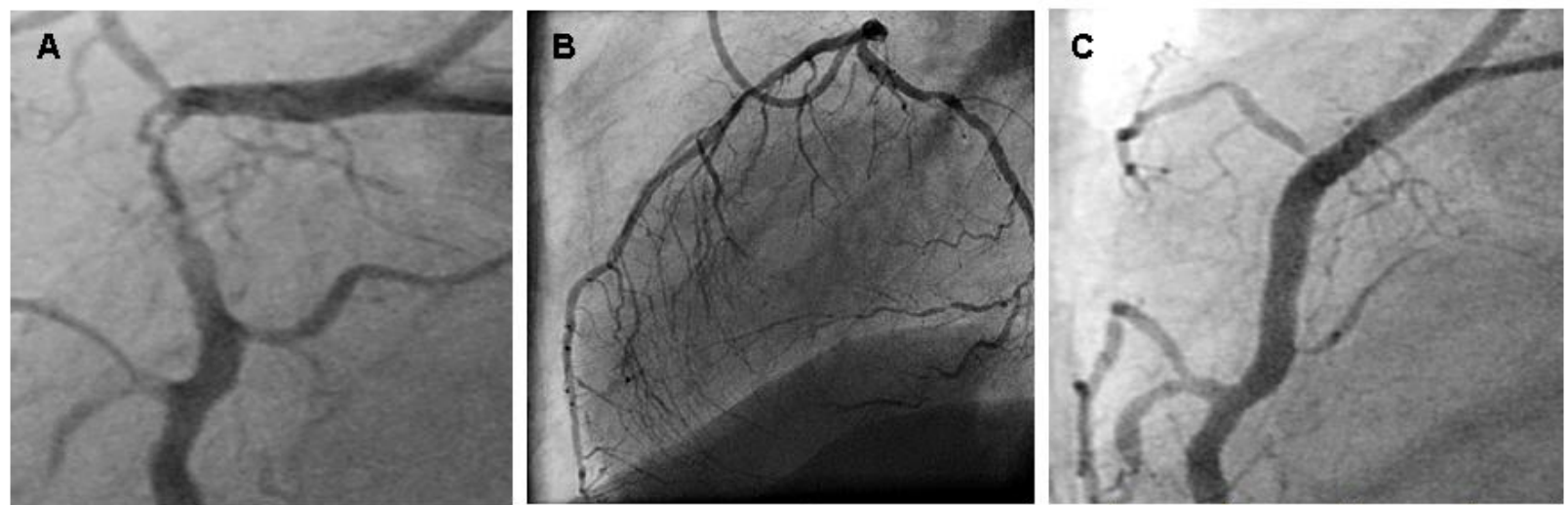


Anatomic stenosis severity by IVUS or QCA is identical but physiologic severity has decreased.
→ **FFR accounts for these changes !!!**

Disconnect between Anatomy and Physiology



FFR in the distal LAD before and After recanalization of the RCA



ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- relation between vessel size and perfusion area
- ***endothelium and development of atherosclerosis***
- the 2 or 3 compartment model of the coron circulation
- collaterals

DEVELOPMENT OF ATHEROSCLEROSIS

Normal



Endothelial dysfunction



First stages of atherosclerosis:



IVUS, OCT, FFR (abnormal pressure decline)

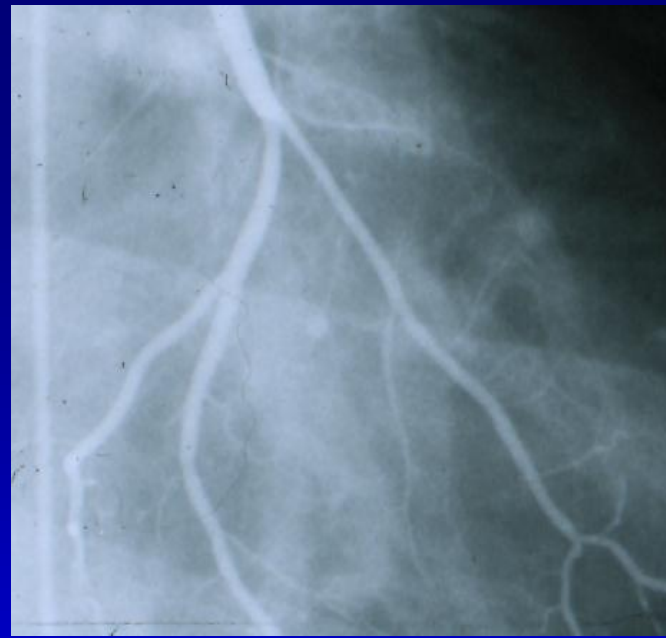
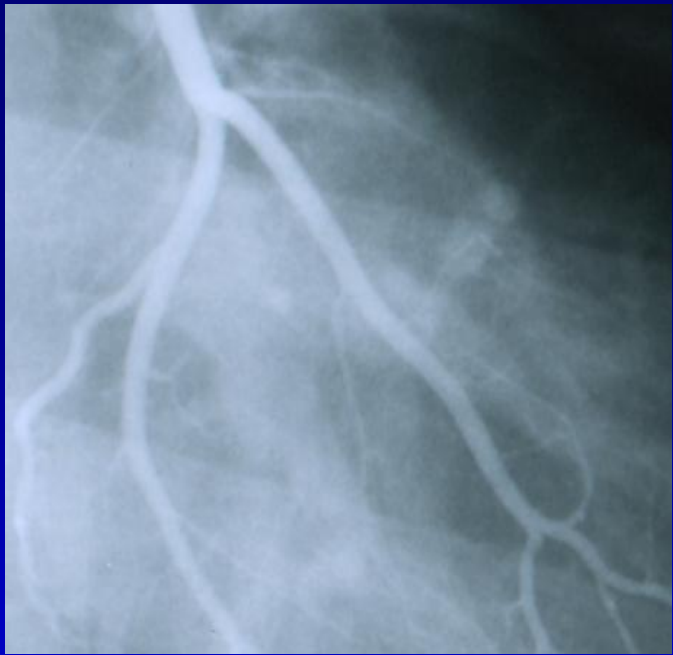
Macroscopic atherosclerotic disease:

angio,

non-invasive imaging (CT, MRI)

*The earliest phase of atherosclerotic coronary disease, is **endothelial dysfunction**.*

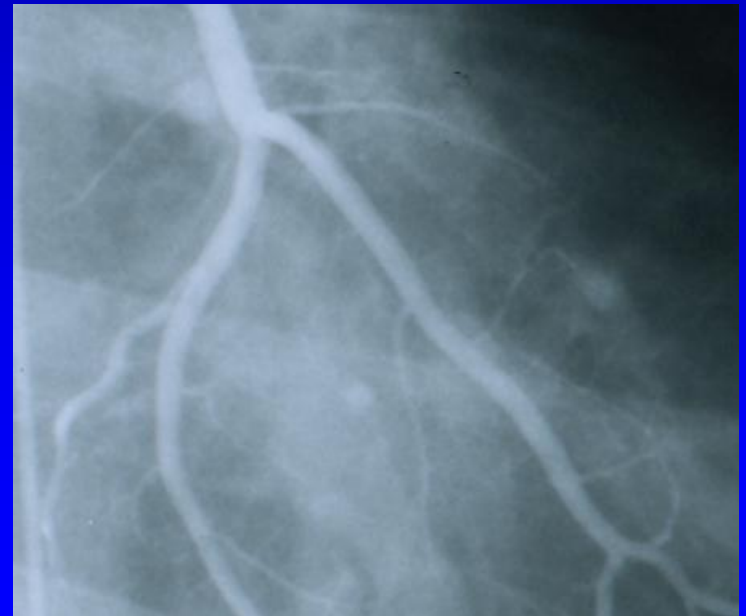
*This is invisible by any imaging method, but can be demonstrated by **functional testing**.*



ACh

baseline

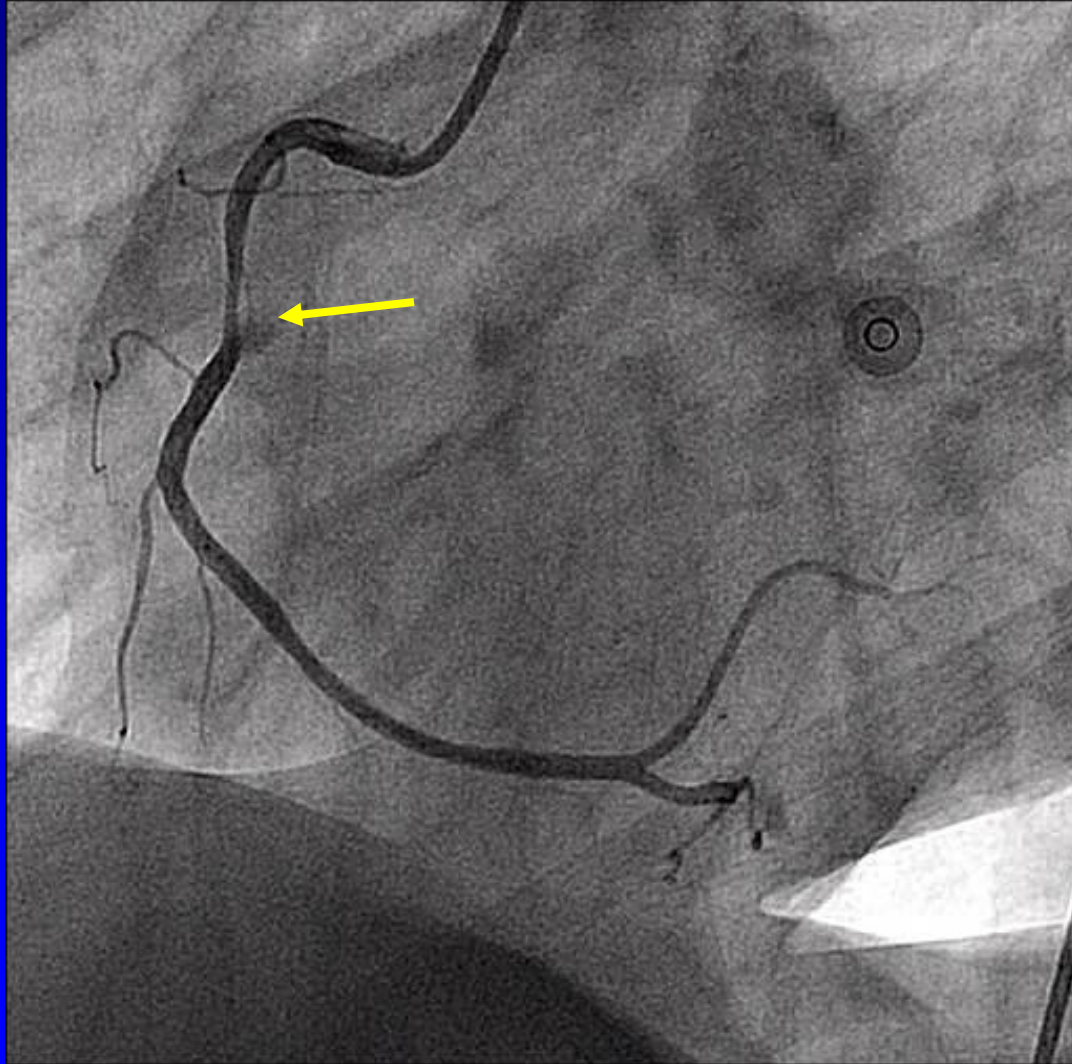
*35-y-old male,
hypertension,
heavy smoker,
chest pain at exercise
and positive ET*

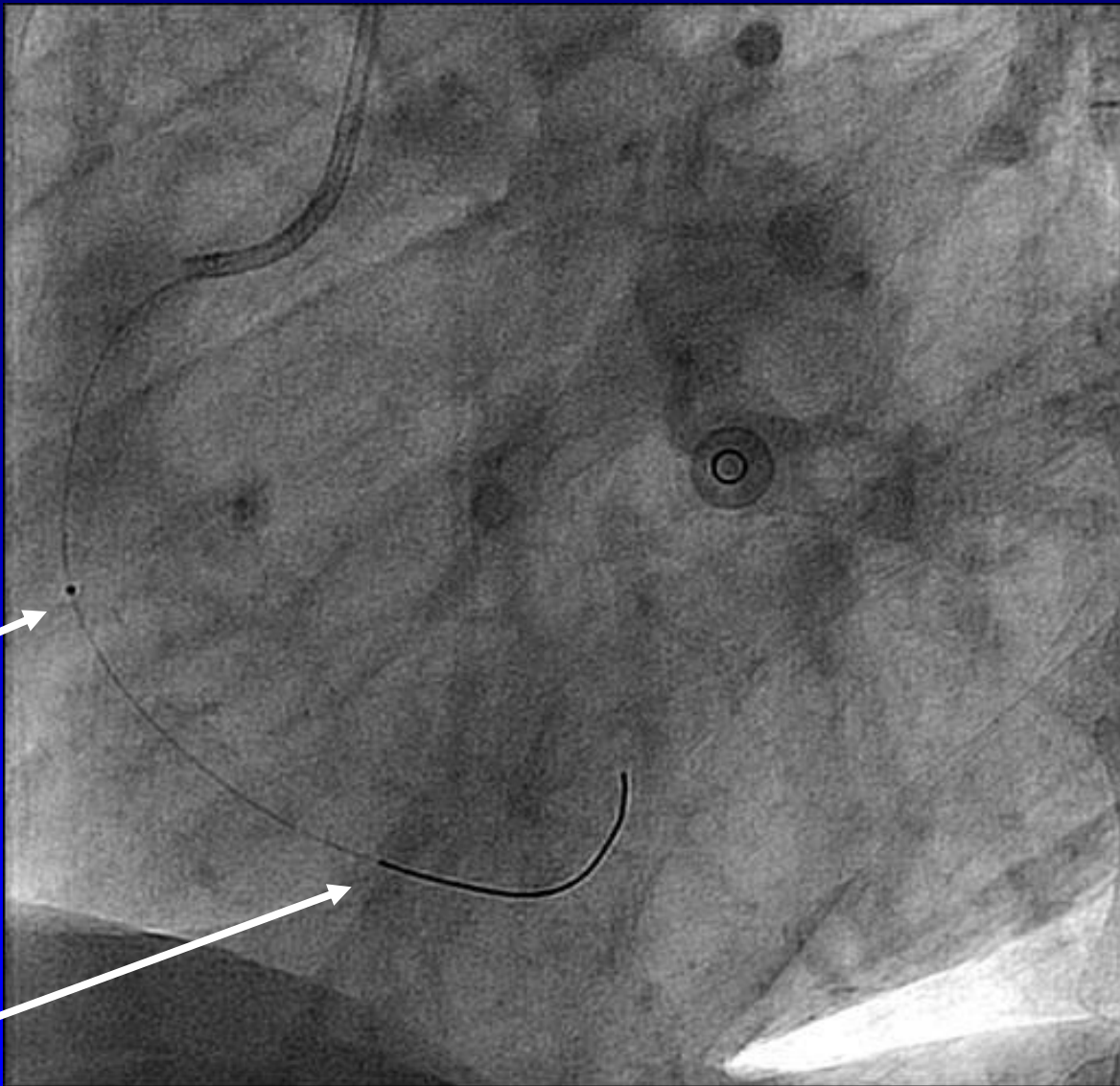


NTG

29 cc/Achol vb (6)

Physiologic **and** pathologic vasomotion in 35-year old male, heavy smoker, and chest pain at exercise

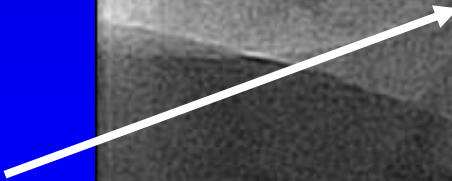


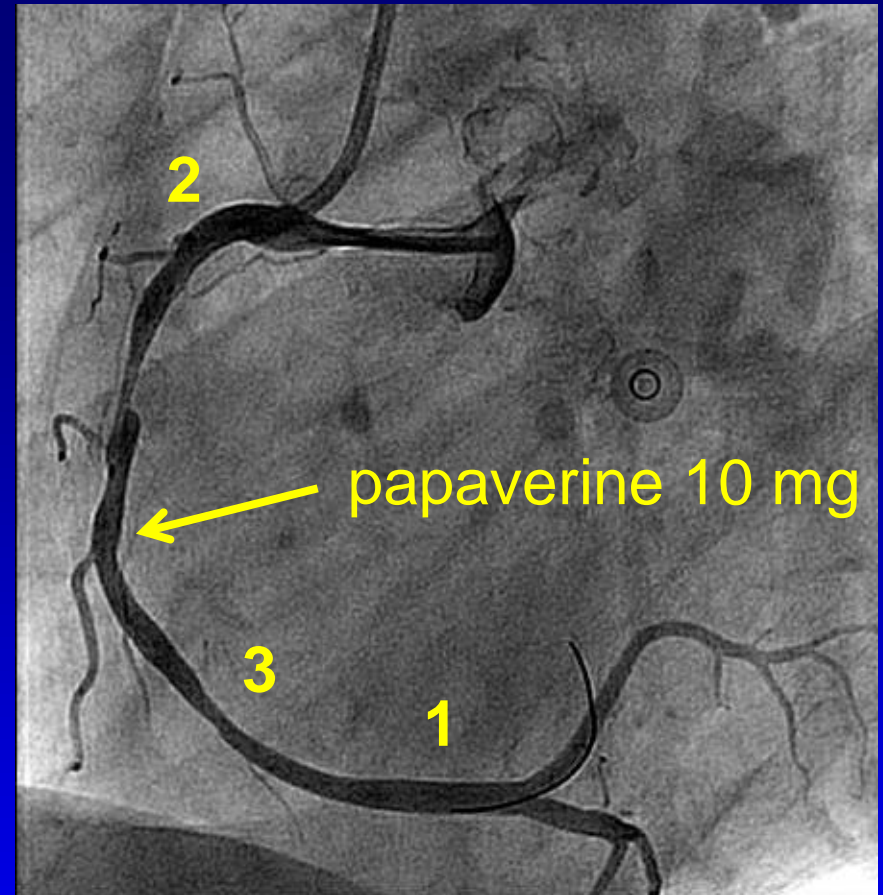
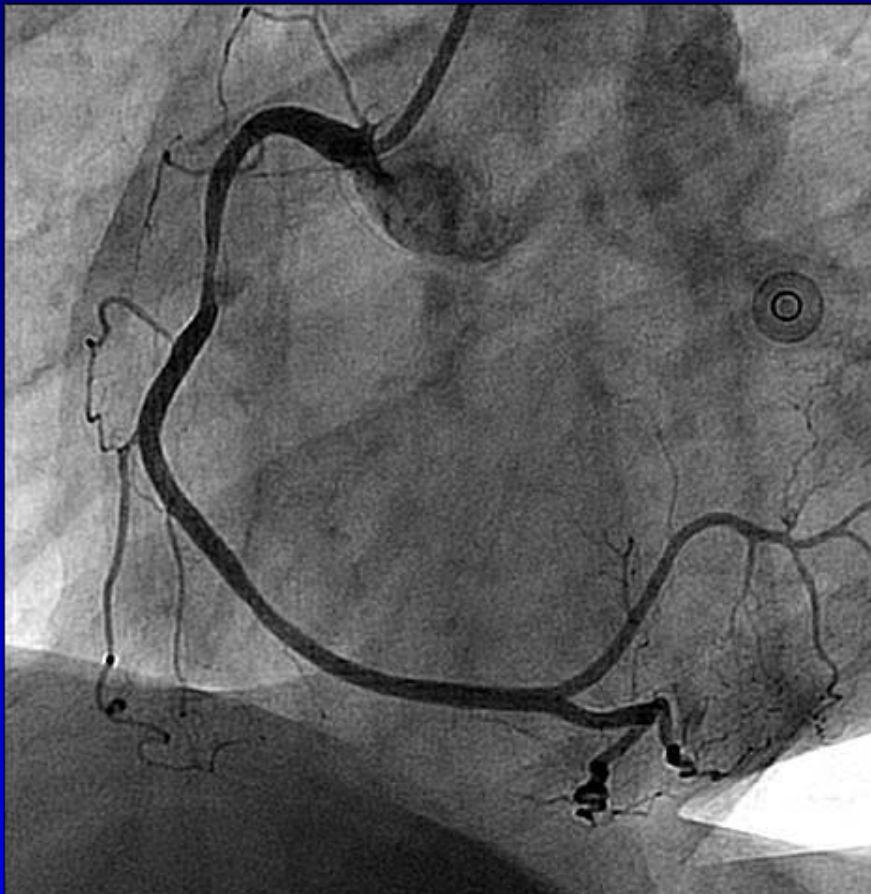


tip of infusion
catheter,
administration
of papaverin



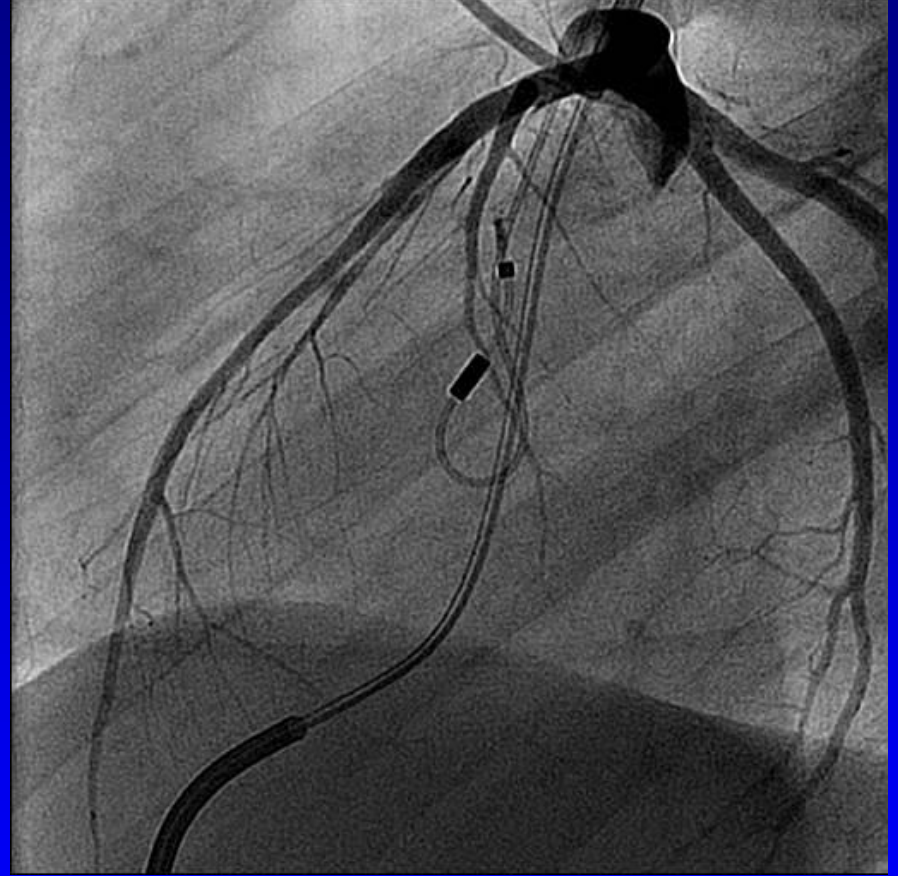
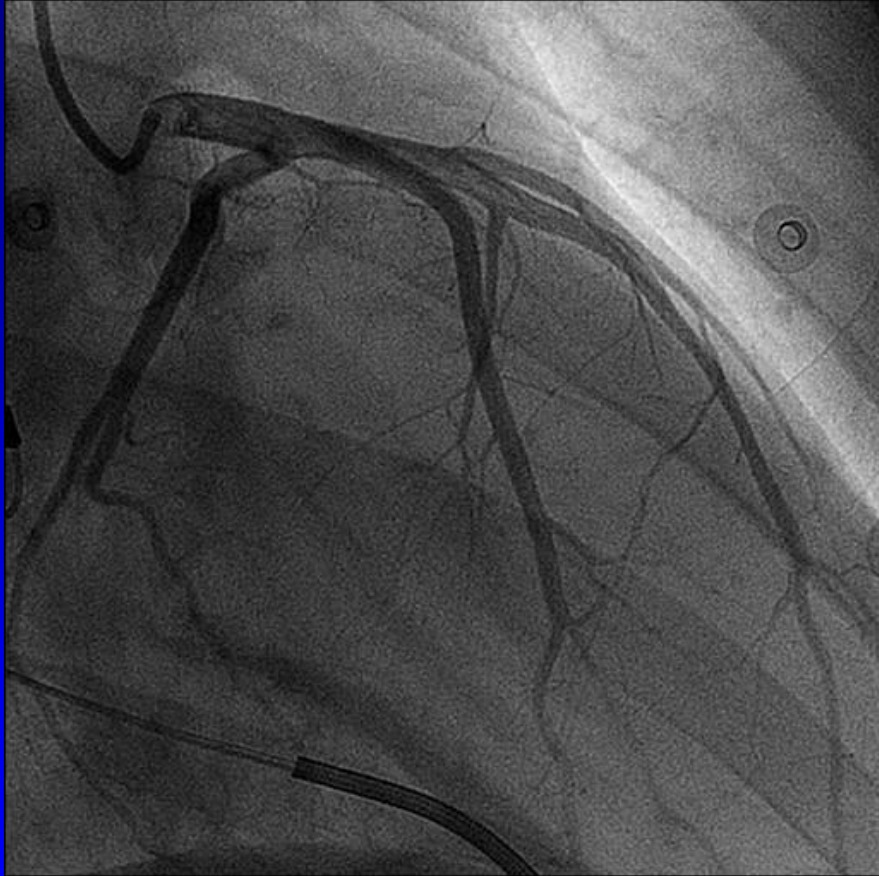
pressure
guidewire



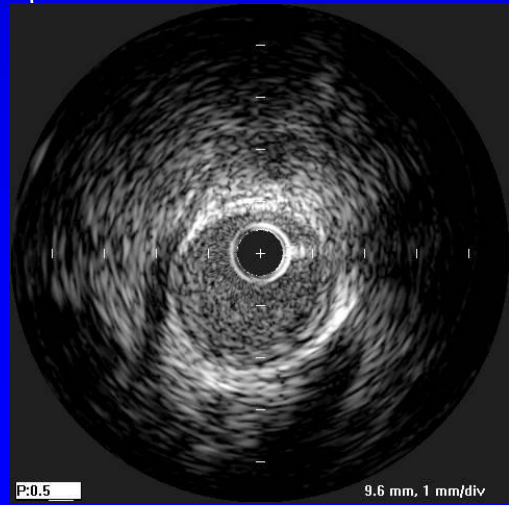
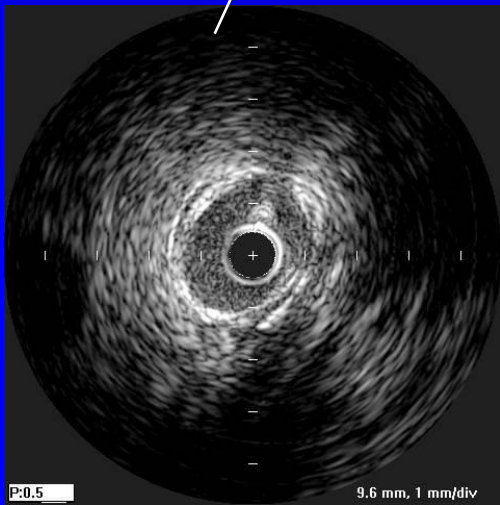
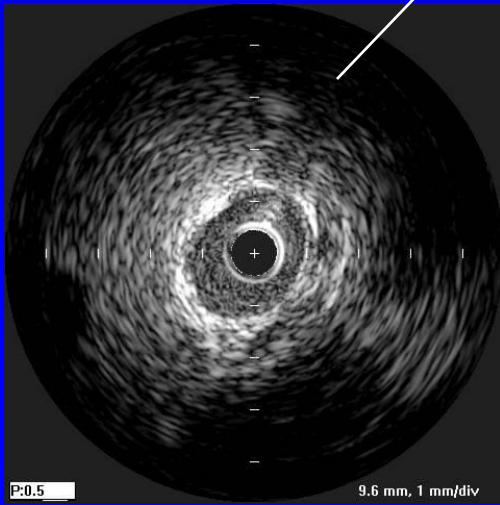
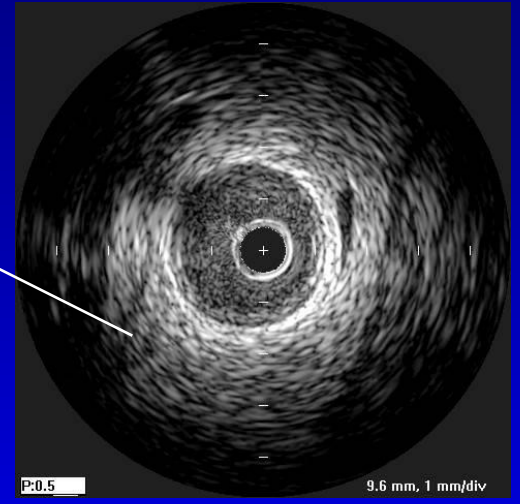
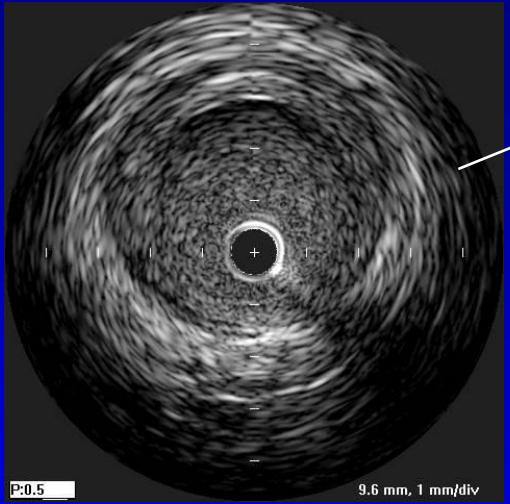
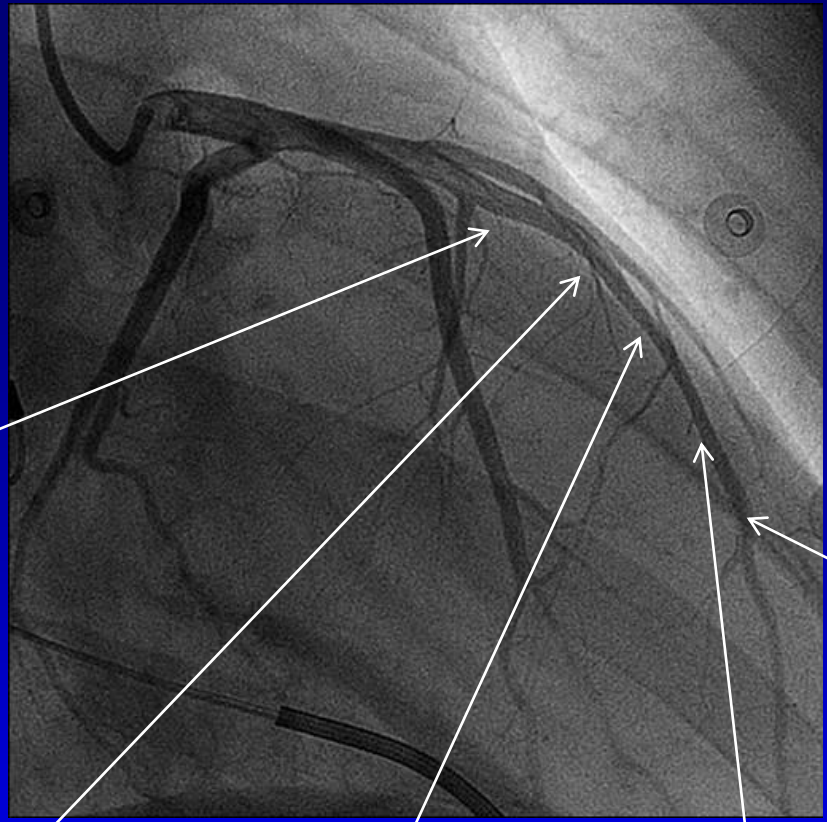


- 1 papaverine induced vasodilation
- 2 flow-induced vasodilation
- 3 flow-induced paradoxical vasoconstriction

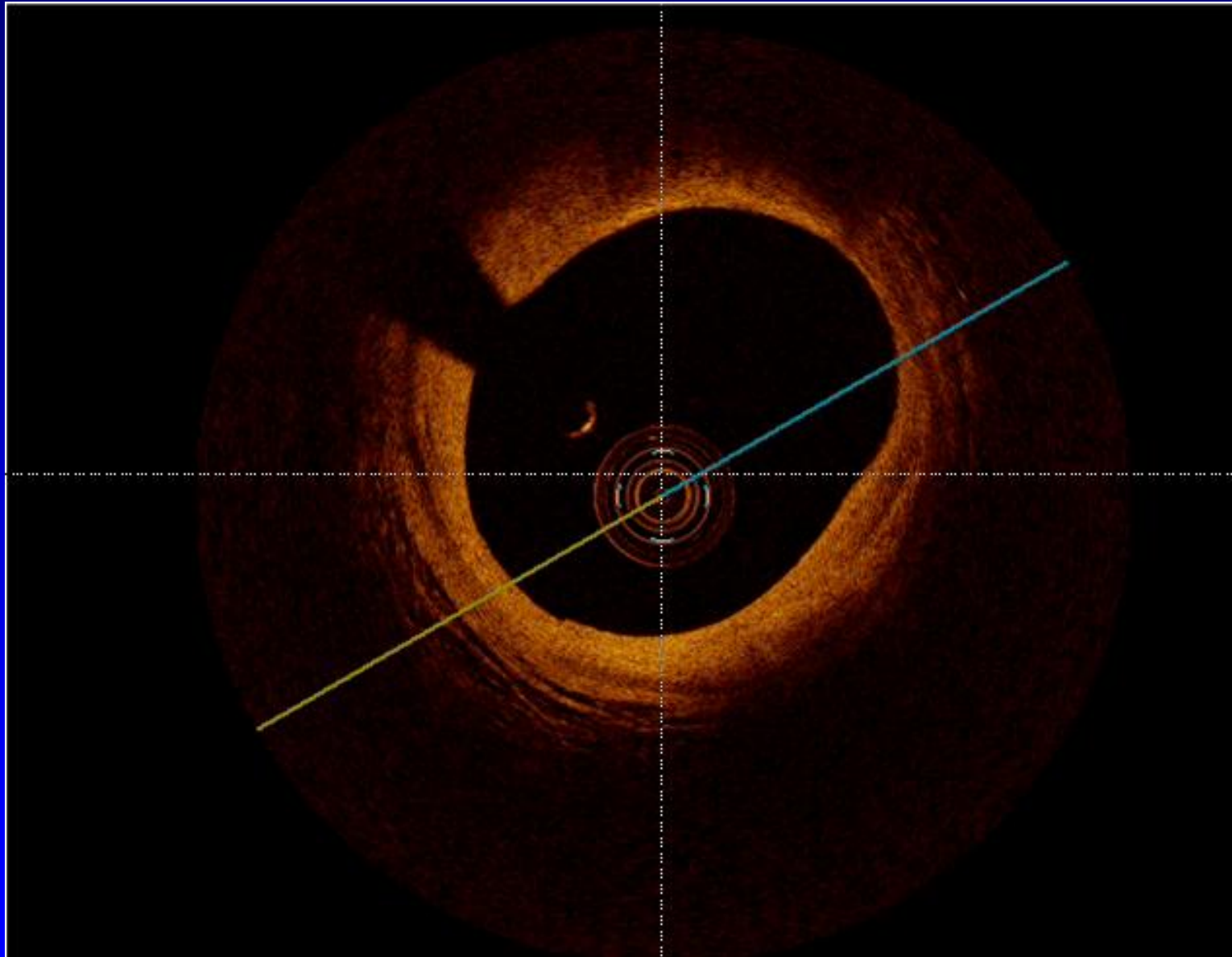
early stage of atherosclerosis



Male, 41-year-old



diffuse atherosclerosis, early stage



Courtesy of Dr Pim Tonino

COM ●

ARCHIVE CUSTOM

C:\RADI\DOWNLOAD\RULO

PATIENT ID	DATE	TIME	VESSEL	PROCEDURE	ACTION	TYPE	SIZE
	2006-03-22	03:37:00				CFR	131Kb
	2006-03-22	03:31:11				FFR	59Kb
	2006-03-22	03:28:44				FFR	8Kb
	2006-03-22	03:10:34				FFR	31Kb
	2006-03-22	03:07:57				FFR	26Kb

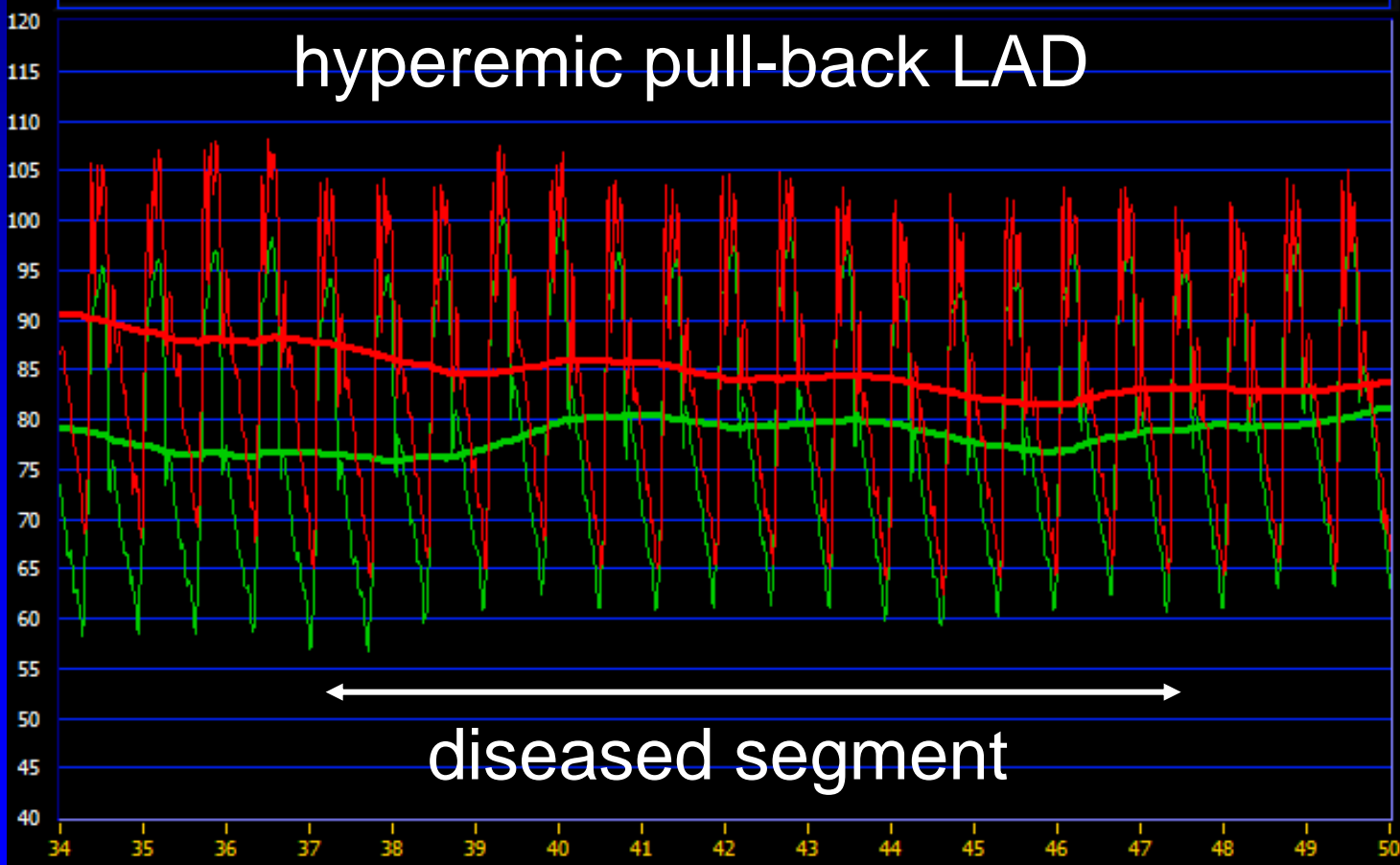


PRINT EDIT RENAME EXPORT ERASE SETUP

[Empty box]

2006-03-22 03:31:11

hyperemic pull-back LAD



82
Pa mean

70
Pd mean

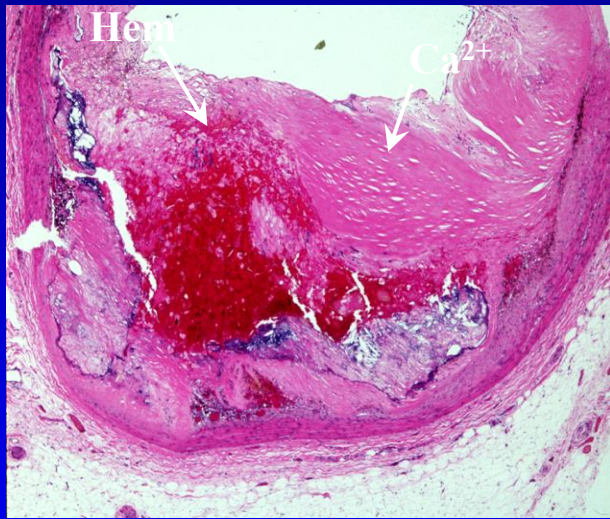
0,85
FFR

21,7
CURSOR

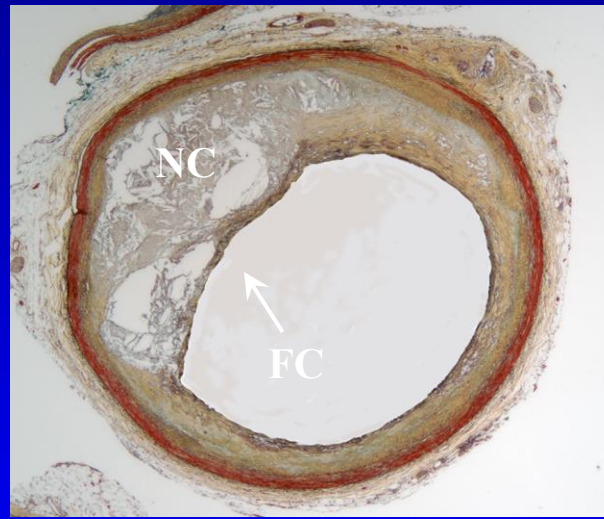


RESET

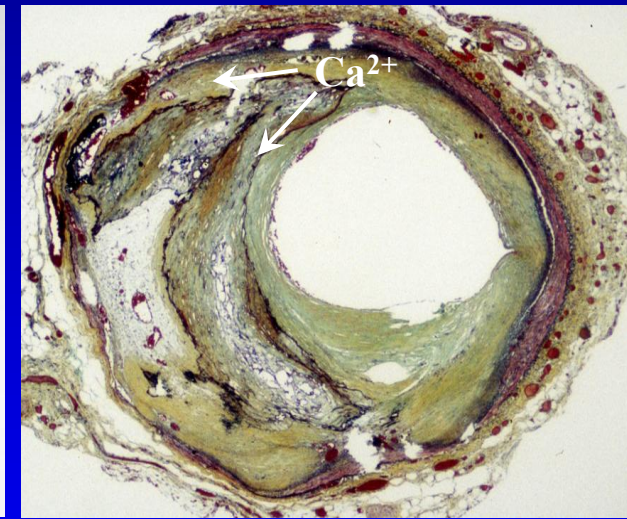
Different stages of gross coronary atherosclerosis, easily visible on angiogram and by several non-invasive methods



Fibrous cap atheroma with hemorrhage



Thin fibrous cap atheroma



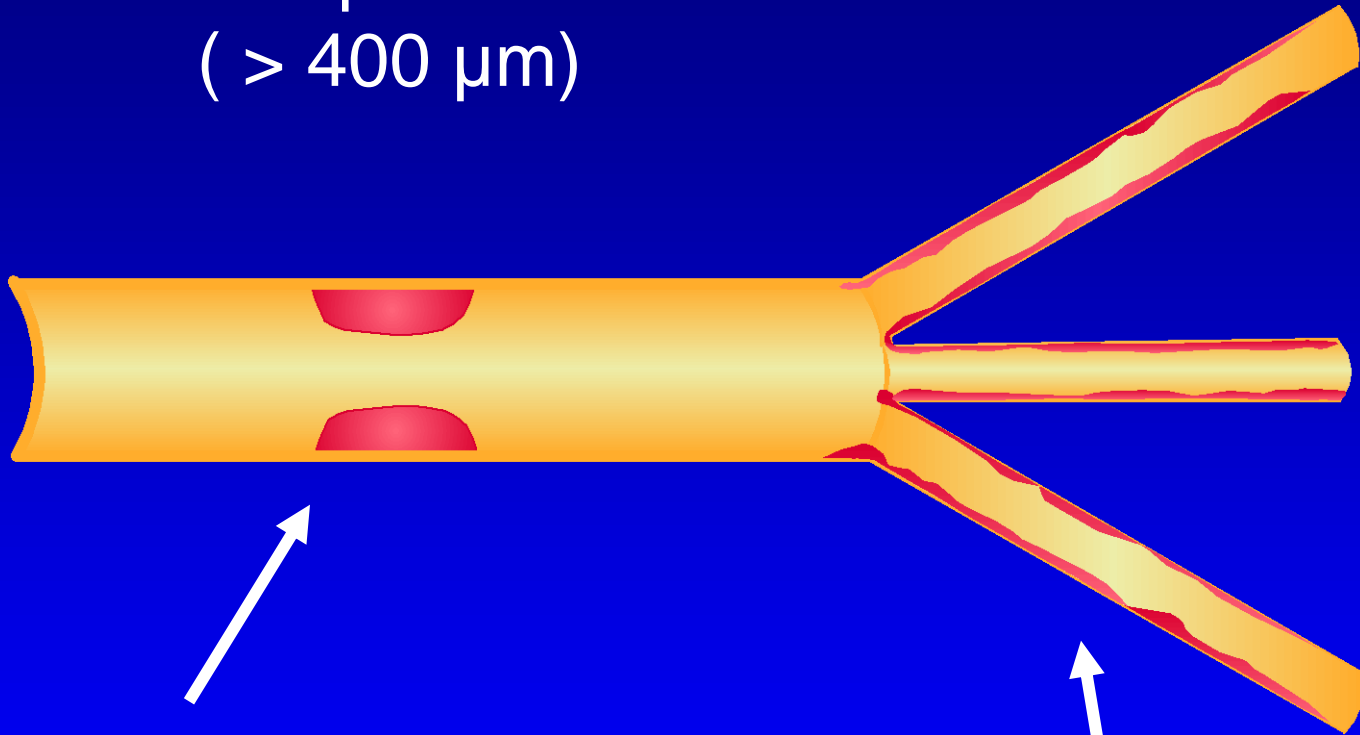
Fibrocalcific plaque

ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- relation between vessel size and perfusion area
- endothelium and development of atherosclerosis
- ***the 2 or 3 compartment model of the coronary circulation***
- collaterals

epicardial
compartment
(> 400 μm)

microvascular
compartment



traditionally visible by angiography
and more recently by many invasive
and non-invasive imaging methods

Black box
(until recently)

IMAGING OF THE EPICARDIAL COMPARTMENT

- non-invasively by CT, MRI
- invasively by angio, IVUS, OCT, and some newer techniques

FUNCTIONAL ASSESSMENT OF THE EPICARDIAL COMPARTMENT

- coronary pressure & FFR

The coronary microcirculation: *Still a black box ??*

IMAGING OF THE MICROVASCULAR COMPARTMENT

- can only be done indirectly (blush, PET, MIBI, MRI)

FUNCTIONAL ASSESSMENT OF THE MICROCIRCULATION:

- IMR (*Bill Fearon, Keith Oldroyd*)
- absolute flow & resistance (*Bernard De Bruyne*)

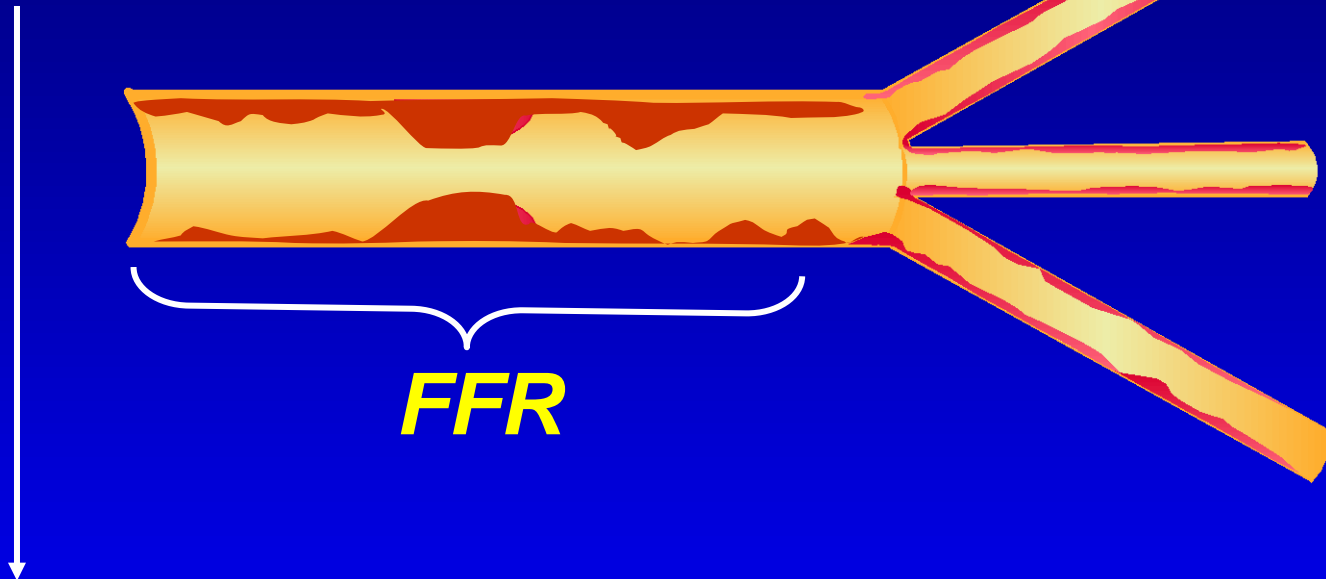
→ Saturday morning session

The third compartment

focal **and diffuse**

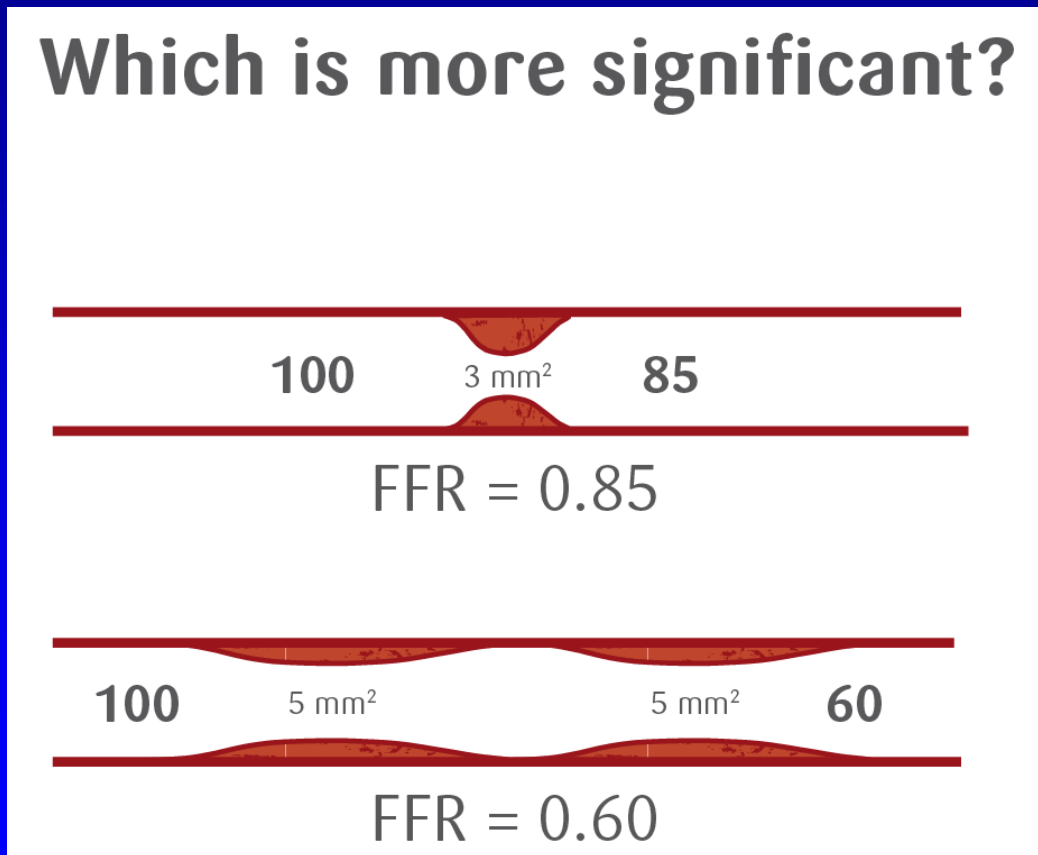
epicardial disease

microvascular
compartment



hard to distinguish by
traditional methods,
but easily assessed
and quantified by FFR
(hyperemic pullback recording)

How to assess the functional significance of diffuse disease, whether or not with super-imposed focal lesions?



CCTA, Angiography, IVUS, or OCT

→ ***Impossible by anatomic methods***

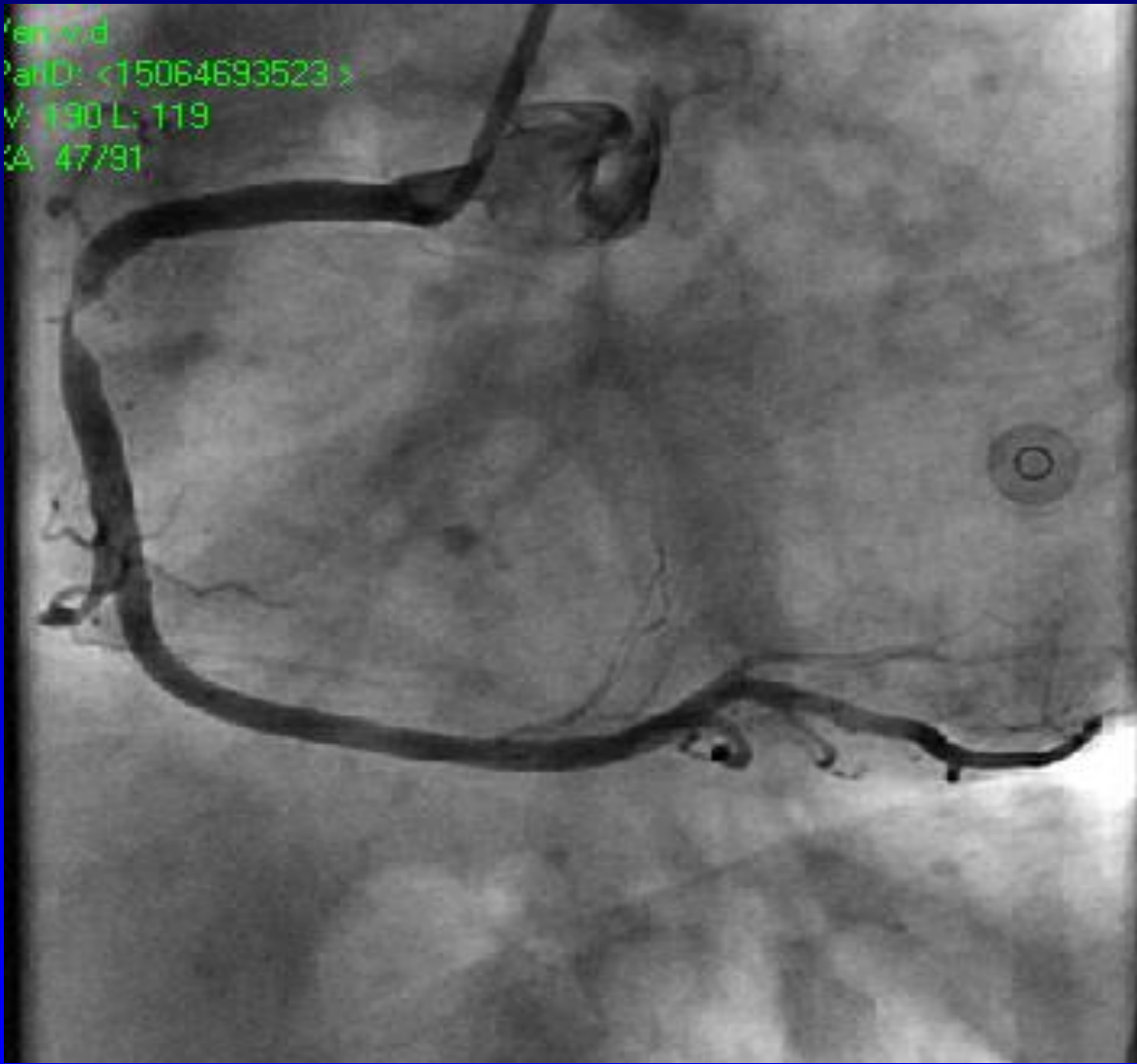
The 3rd compartment:

Diffuse epicardial coronary disease
(whether or not with superimposed focal lesions)
(Bernard De Bruyne, tomorrow morning)

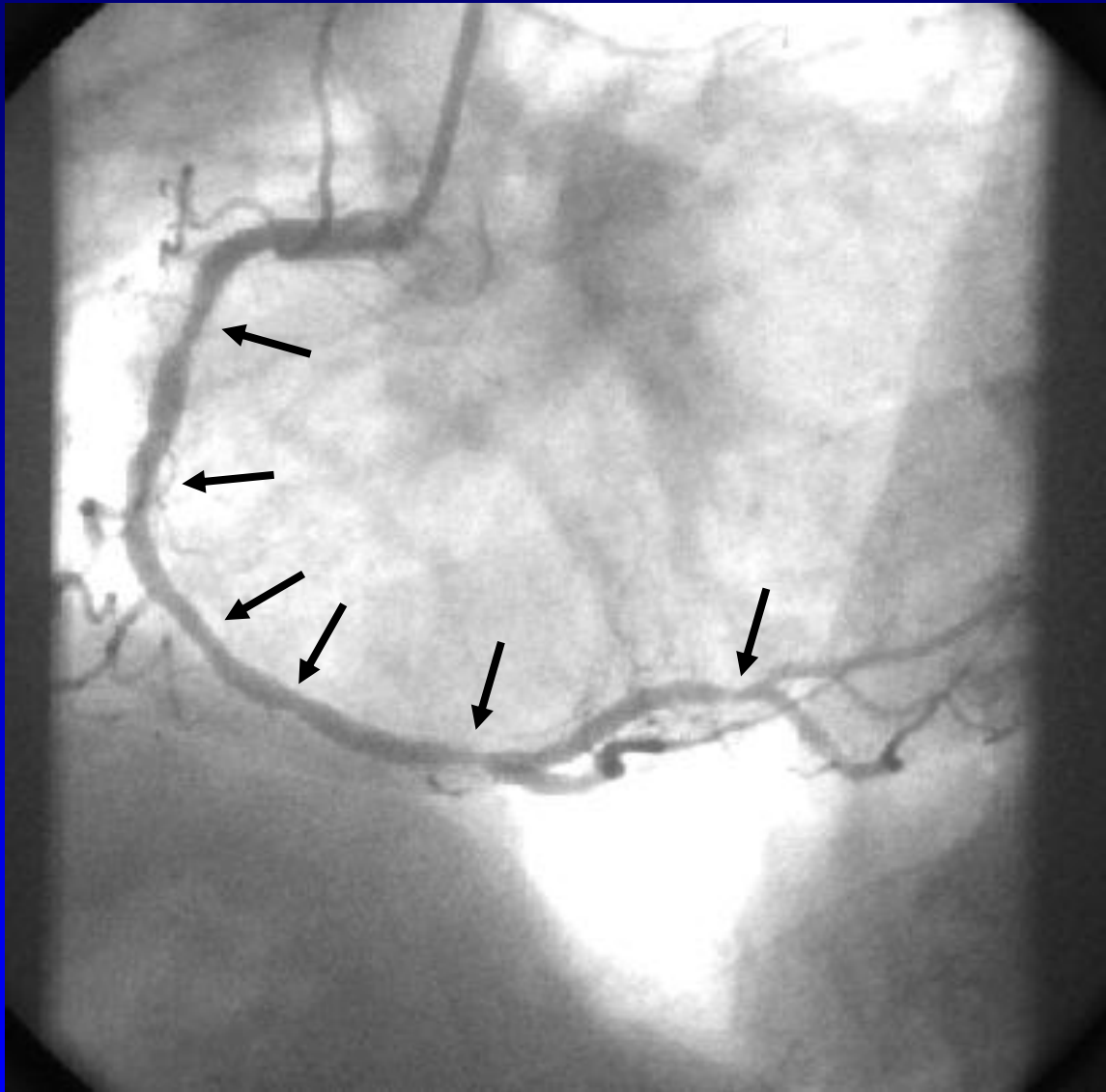
→ **easily evaluable by FFR**
(pressure pull-back recording)



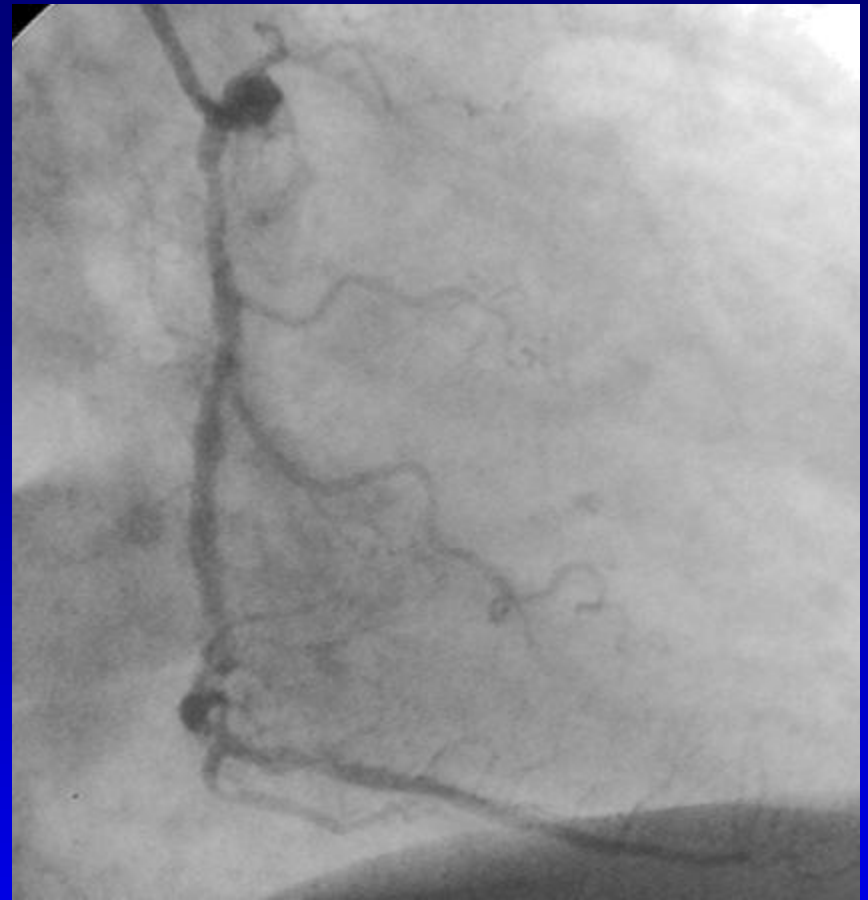
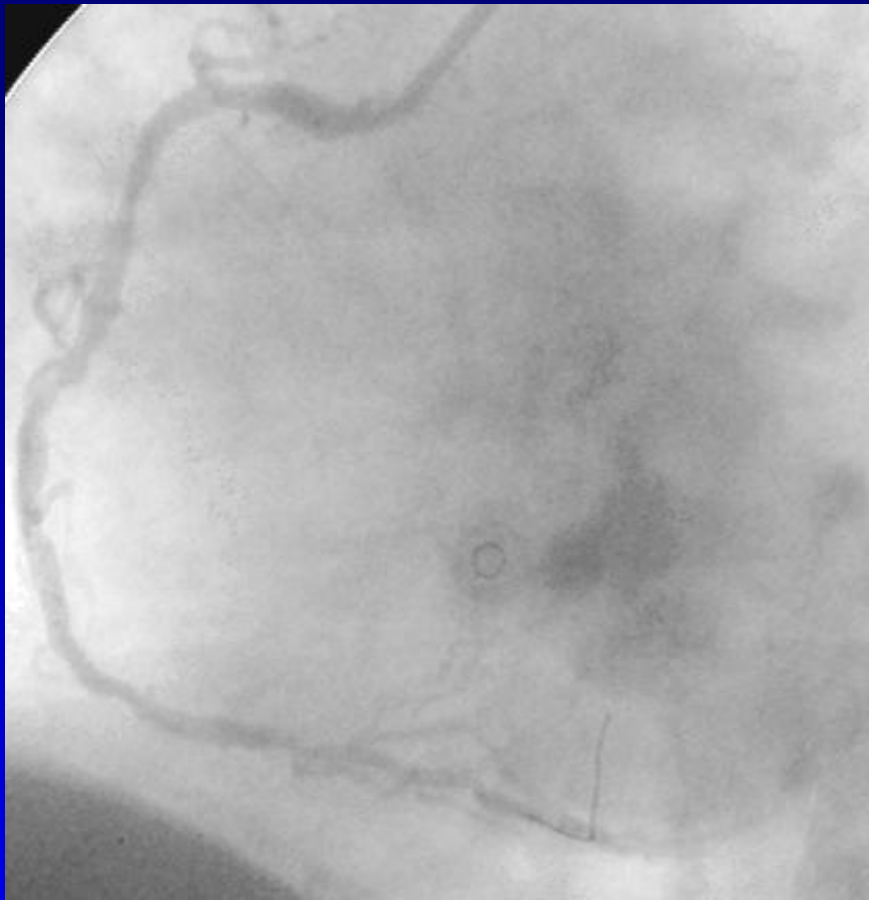
important consequence for treatment
(interventional or medical)



Typical chest pain; positive MIBI-Spect inferior wall

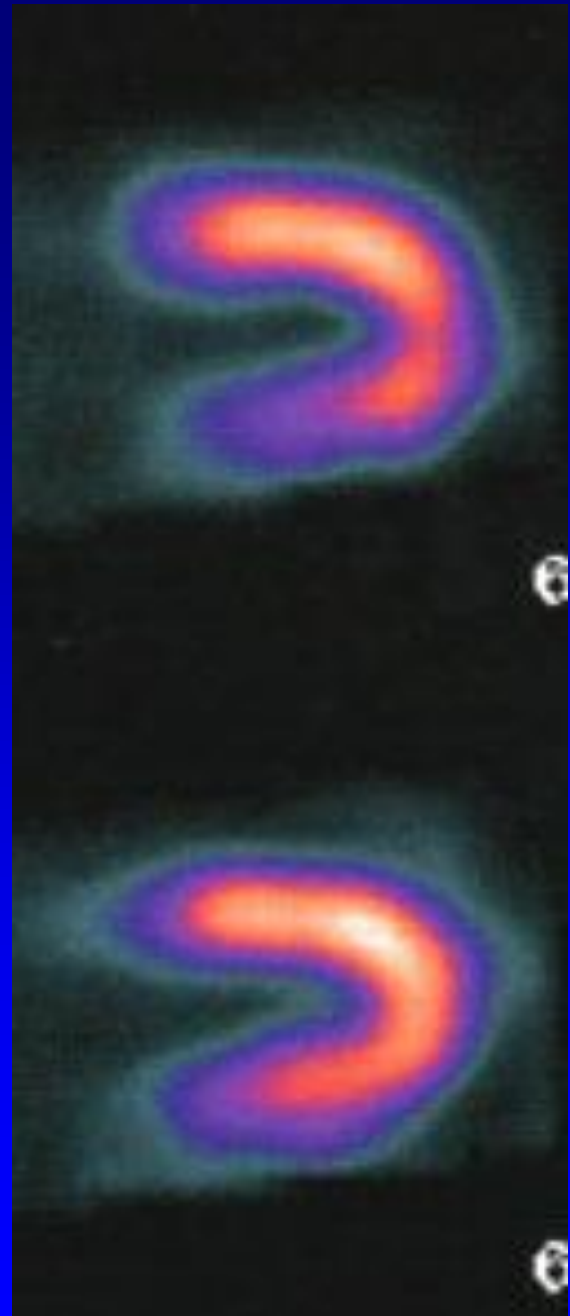
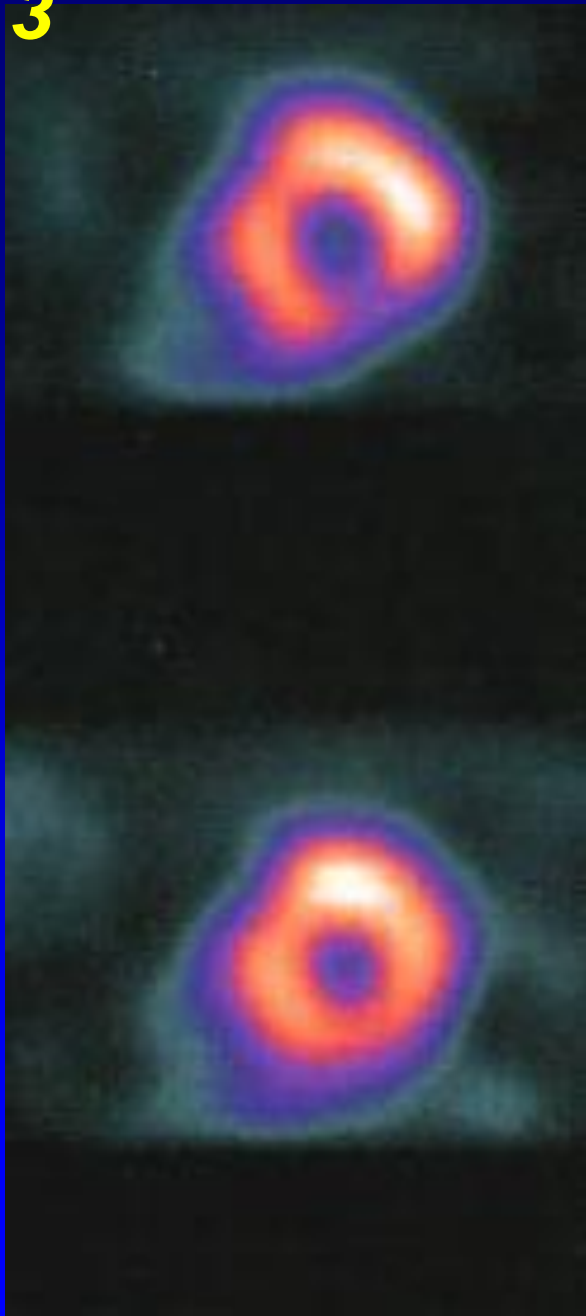


Typical chest pain; positive MIBI-Spect inferior wall



Typical chest pain; positive MIBI-Spect inferior wall

CASE # 3



15064693523
V: 130 L: 119
SA: 47/81

PRINT EDIT RENAME EXPORT ERASE SETUP

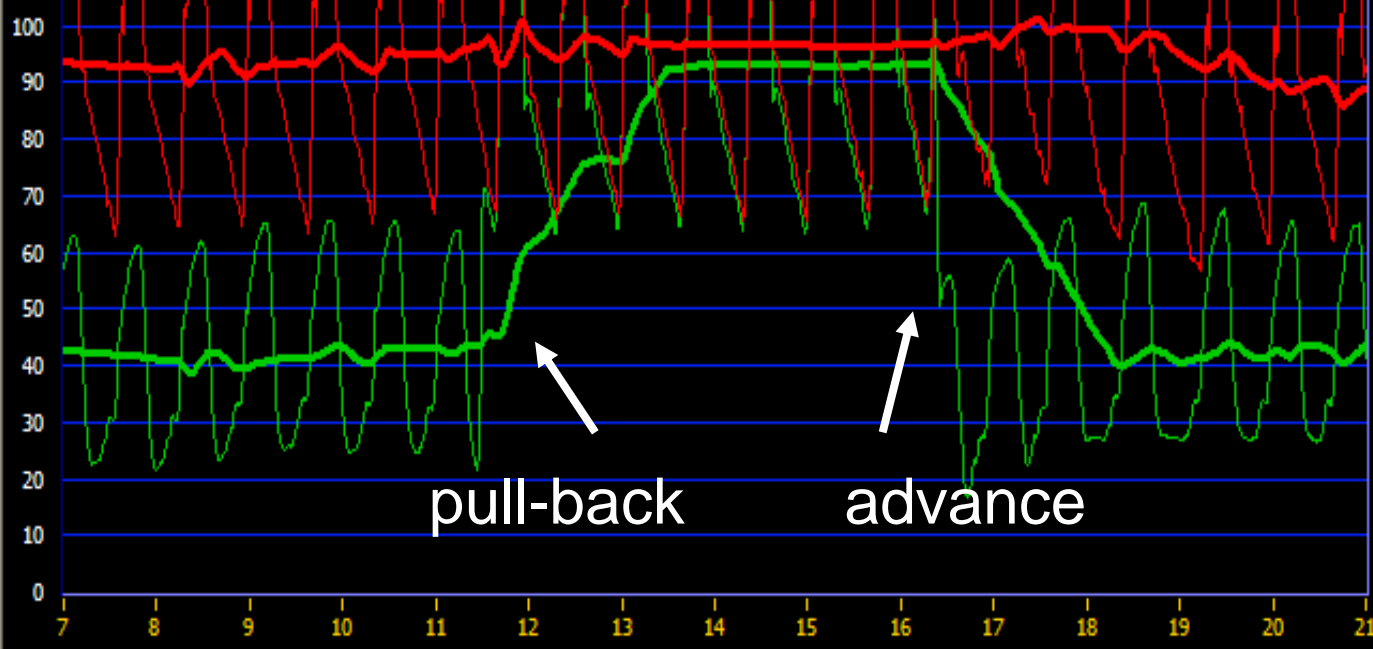
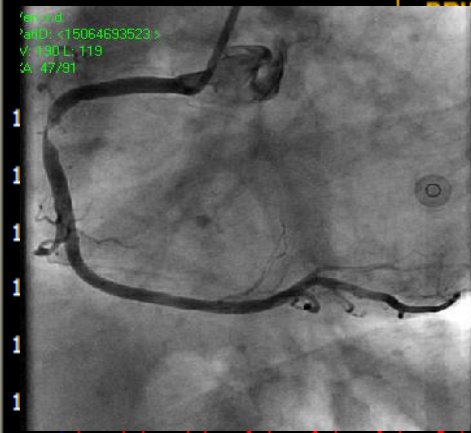
PRE PTCA ADO IV

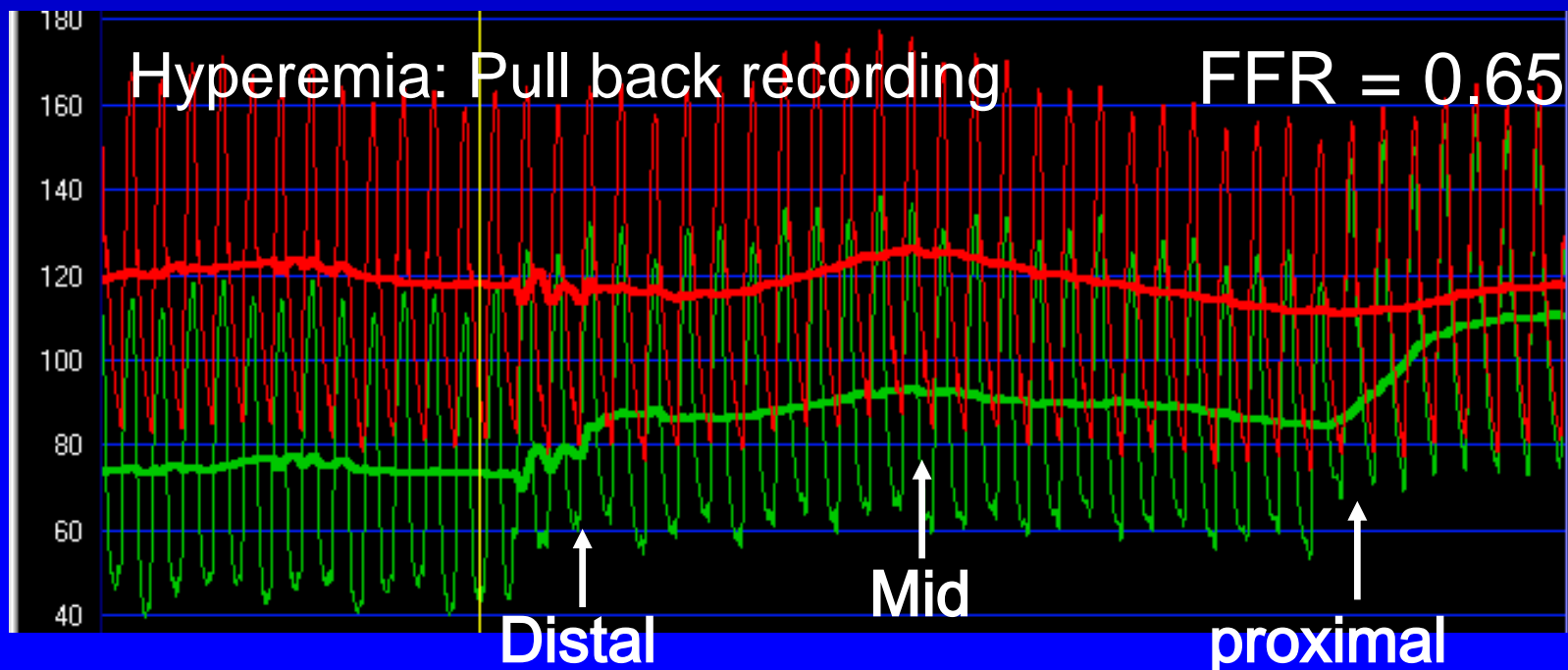
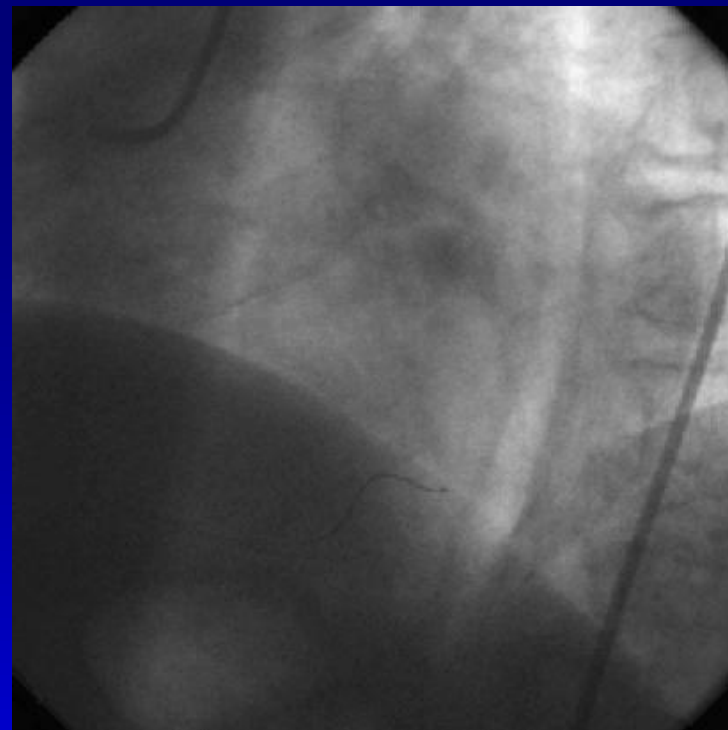
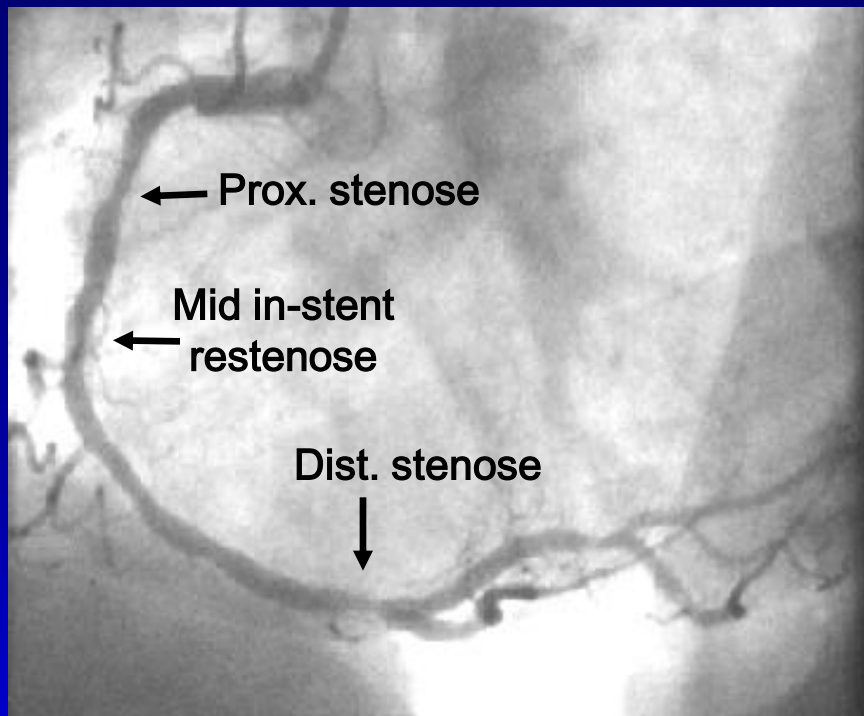
90
Pa mean
40
Pd mean
0,44
FFR

5,6
CURSOR



RESET





EDIT

RENAME

EXPORT

ERASE

SETUP

2004-06-14 15:55:19

109

Pa mean

84

Pd mean

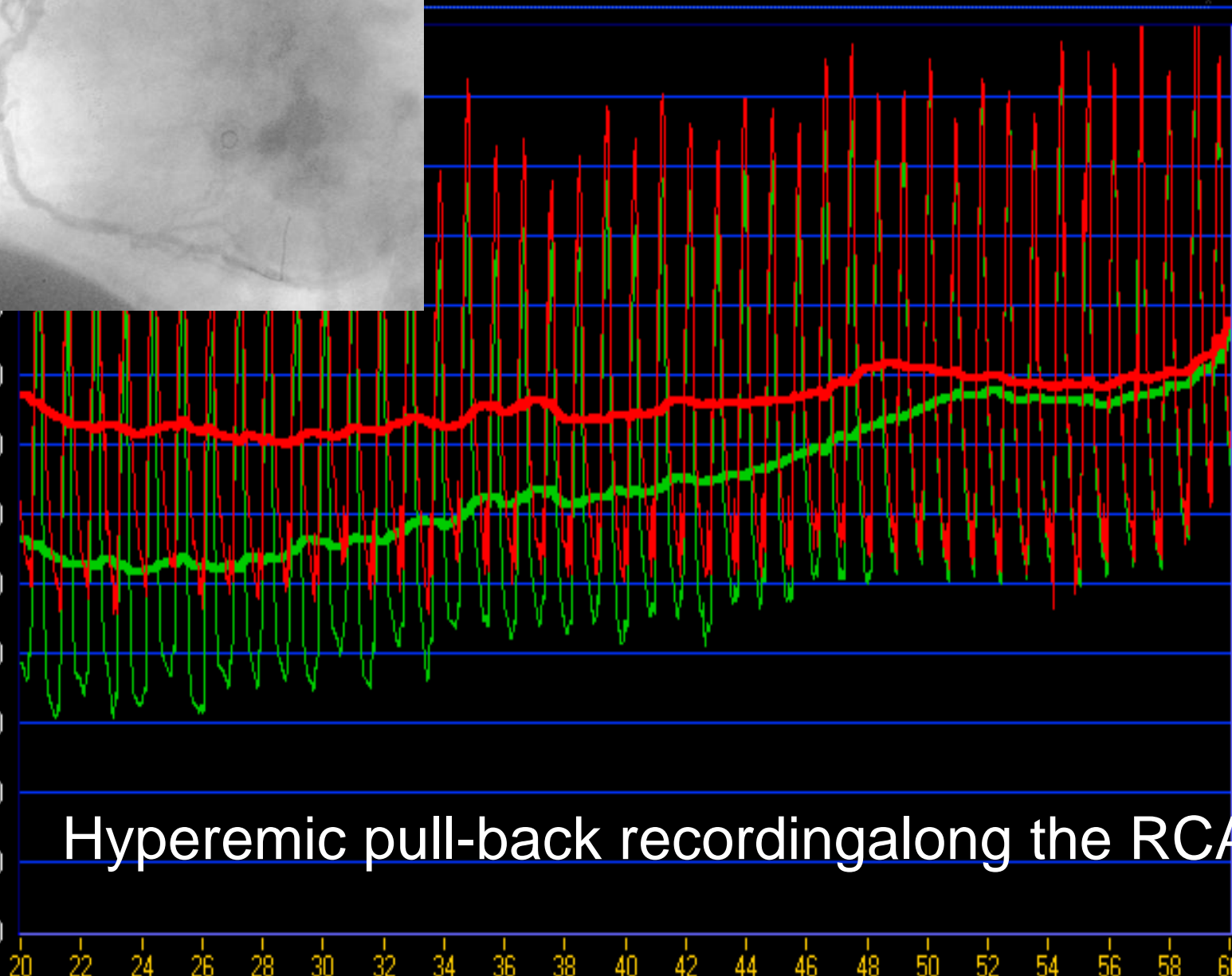
0,77

FFR

7,9

Cursor

15
14
13
12
11
100
90
80
70
60
50
40
30
20



Hyperemic pull-back recording along the RCA

+ 🔍 ↕
RESET

IN SUMMARY:

- Coronary anatomy is just one side of the coin
- There is complex interrelation between the structure and function of the coronary circulation, not only under physiologic circumstances in healthy persons (*vessel size/perfusion area relation, endothelium, regulation of coronary blood flow*), but also under pathologic circumstances (*atherosclerosis, plaques, stenosis, vulnerability, and ischemia*).
- Understanding this relation is paramount to treat our patients in the cathlab in the best possible way.
- Hopefully, this course will contribute both to that *understanding* and to its translation into *practical skills*

EINDE

The 3rd compartment:

Diffuse epicardial coronary disease, whether or not with super-imposed focal disease

(Bernard De Bruyne, tomorrow)

In patients with coronary artery disease, the most important factor with respect to both

- *functional class (symptoms)*
- *and prognosis (outcome)*

Is the presence and extent of inducible ischemia

→ knowledge if and which lesion(s) is / are responsible for inducible ischemia, is paramount for adequate treatment in the cath.lab

→ **FRACTIONAL FLOW RESERVE**

FFR: The Pressure Pull-back Curve

Pressure pull-back curve at maximum hyperemia:

- place sensor in distal coronary artery
- induce sustained maximum hyperemia by i.v. adenosine, or i.c. papaverine
- pull back the sensor slowly under fluoroscopy
- the individual contribution of every segment and spot to the extent of disease can be studied in this way

Coronary pressure is unique in this respect and such detailed spatial information cannot be obtained by any other invasive or non-invasive method

ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- relation between vessel size and perfusion area
- endothelium and development of atherosclerosis
- the 2 or 3 compartment model of the coron circulation
- ***collaterals***
- why functional testing / FFR ?
- which lesions should be treated

ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- relation between vessel size and perfusion area
- endothelium and development of atherosclerosis
- the 2 or 3 compartment model of the coron circulation
- collaterals (*to be discussed tomorrow*)
- ***which lesions should be treated ?*** } next 2 days
- ***why functional testing / FFR ?*** }

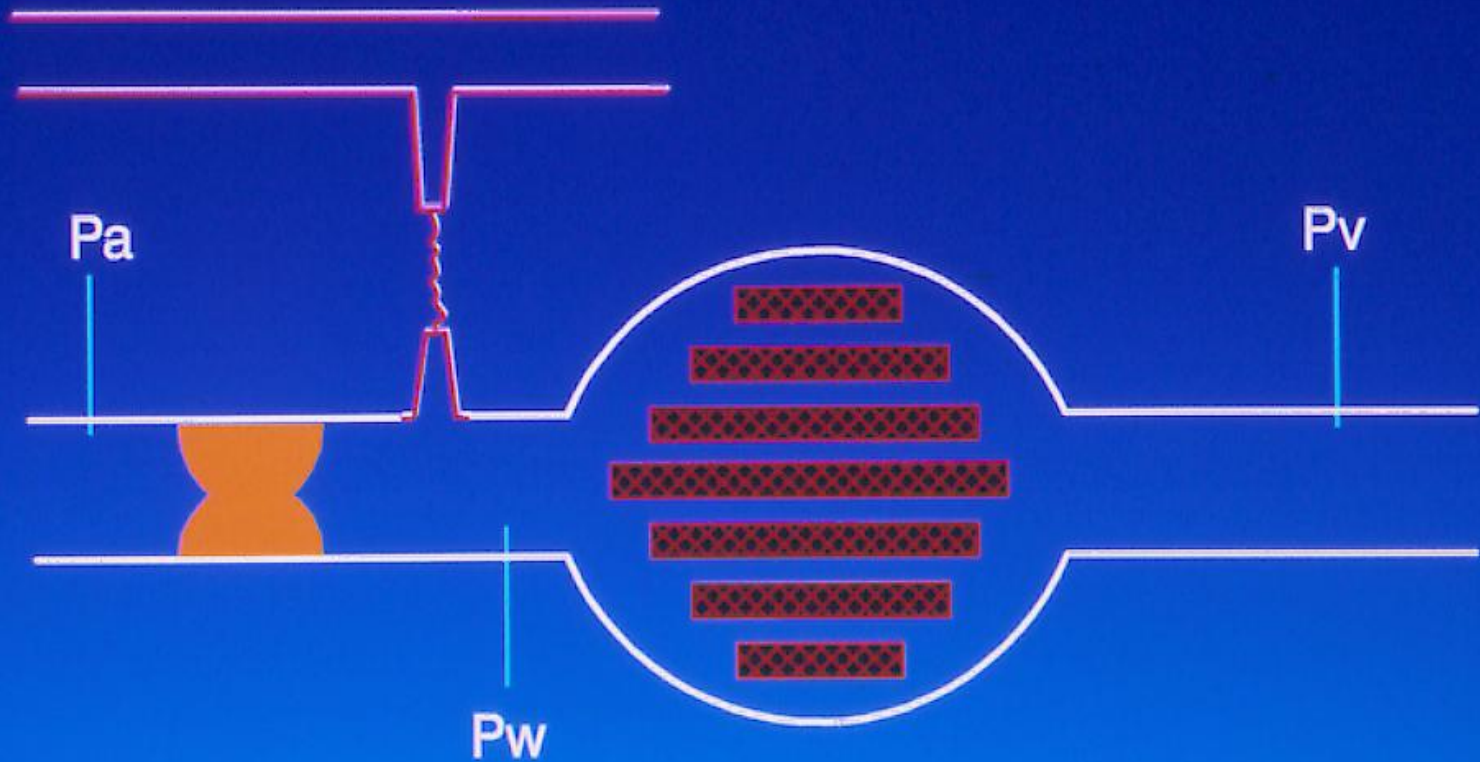
$$Q_{myo} = Q_{cor.artery} + Q_{collateral}$$

Quantitative assessment of the contribution of **coronary arterial** and **collateral flow** to total **myocardial flow** is possible by coronary pressure measurements, but not trivial

Pijls & De Bruyne:

Circulation 1993

Coronary Pressure, sec edition, Kluwer 2000



Fractional collateral flow (also called CFI_p) =

$$FFR_{coll} = \frac{P_w - P_v}{P_a - P_v}$$

Venous pressure not negligible anymore !

EVIDENCE-BASED MEDICINE:

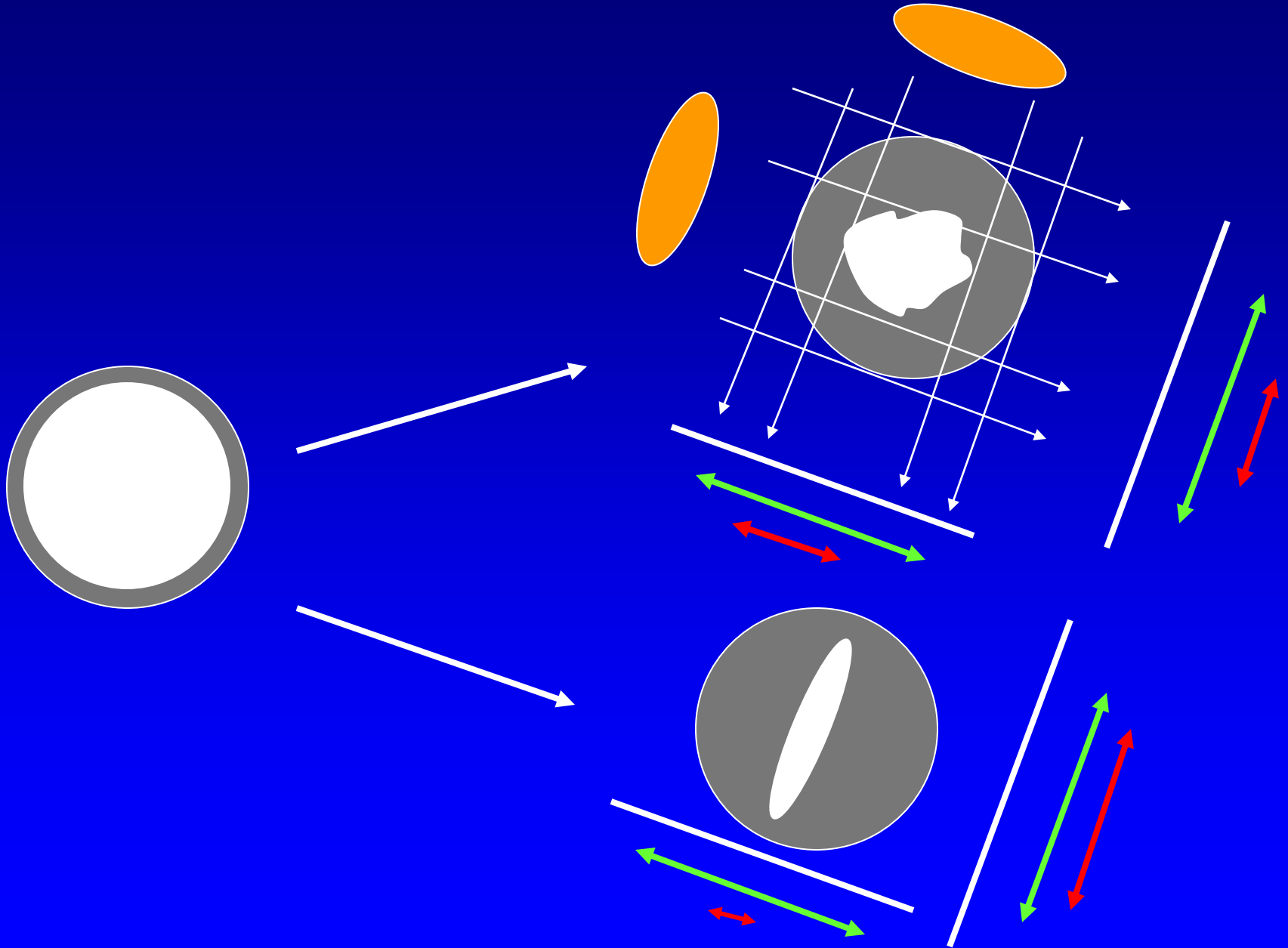
- **PCI of “ischemic” lesions (*associated with reversible ischemia*) makes sense and improves symptoms and sometimes also outcome**
- **PCI of non-ischemic lesions has no benefit, is no evidence-based medicine, is potentially harmful, and unnecessary expensive**

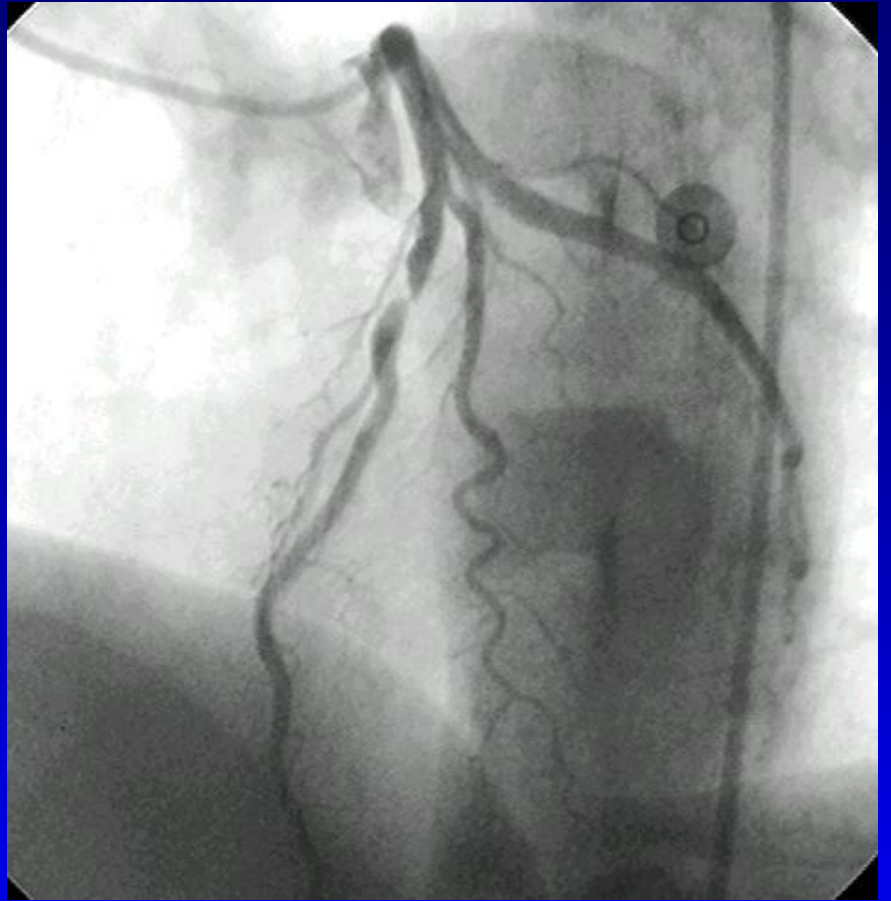
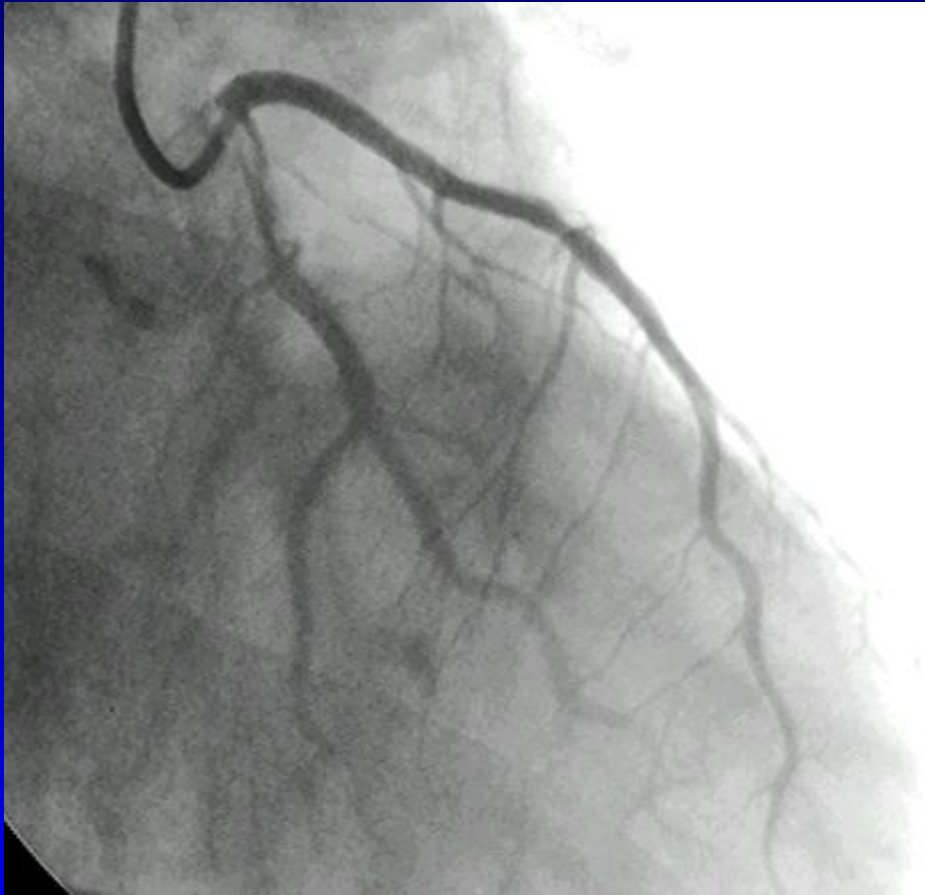
→ **knowledge if and which lesion(s) is / are responsible for inducible ischemia, is paramount for adequate treatment in the cath.lab**

→ **FRACTIONAL FLOW RESERVE**

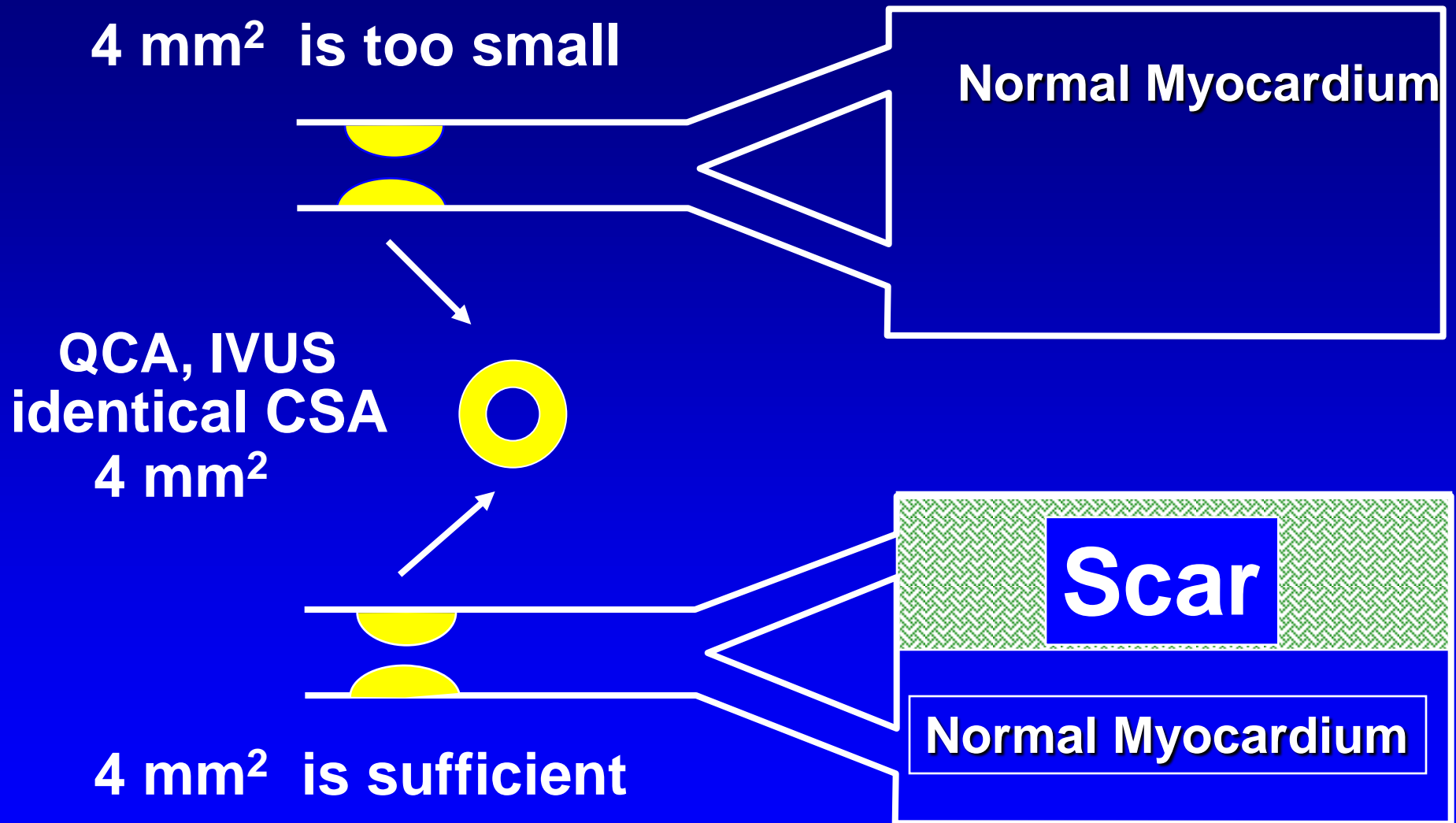
THE CORONARY ANGIOGRAM IS ONLY A CRUDE TOOL TO PREDICT IF A STENOSIS CAUSES ISCHEMIA:

- shortcomings of imaging itself
- discrepancy between structure and function
(especially under pathologic conditions)
- very hard to predict functional severity of disease from structural abnormalities
- complex influence of pathologic structure on blood flow

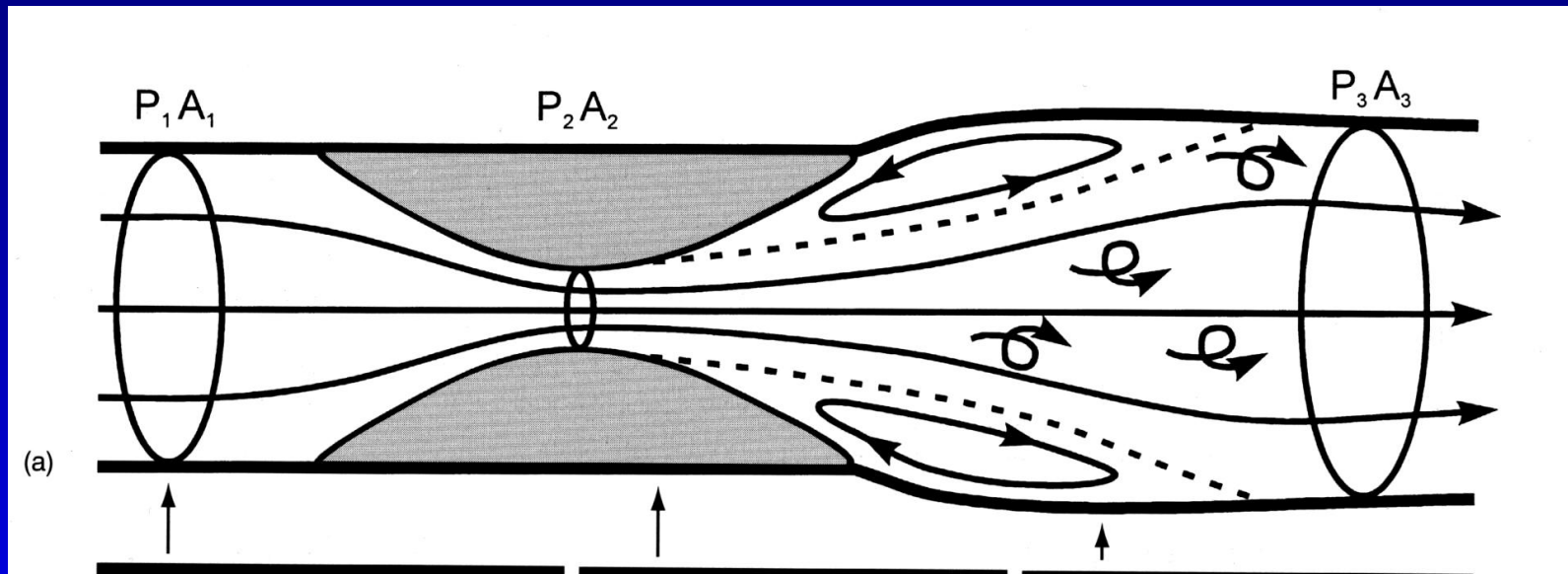


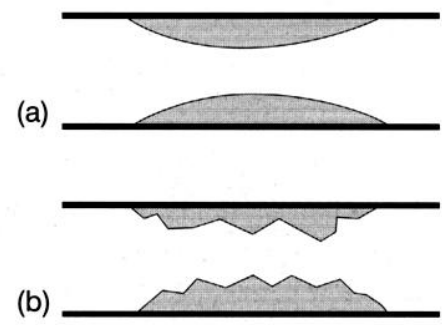
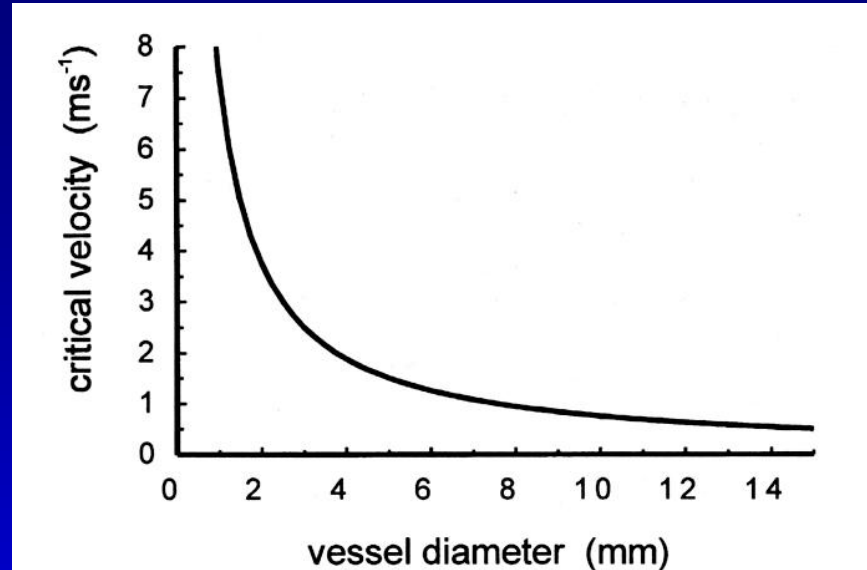
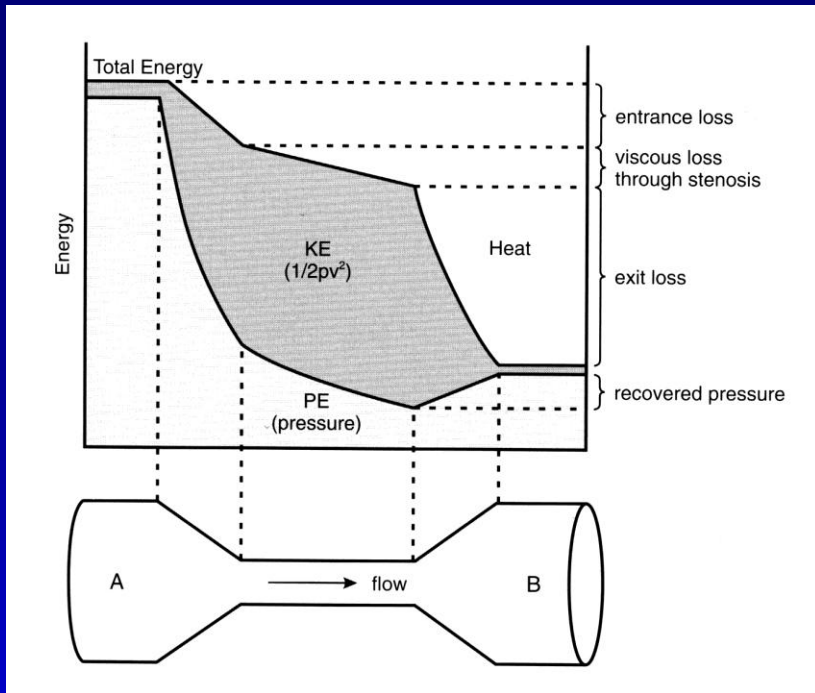


similar stenosis but different extent of perfusion area

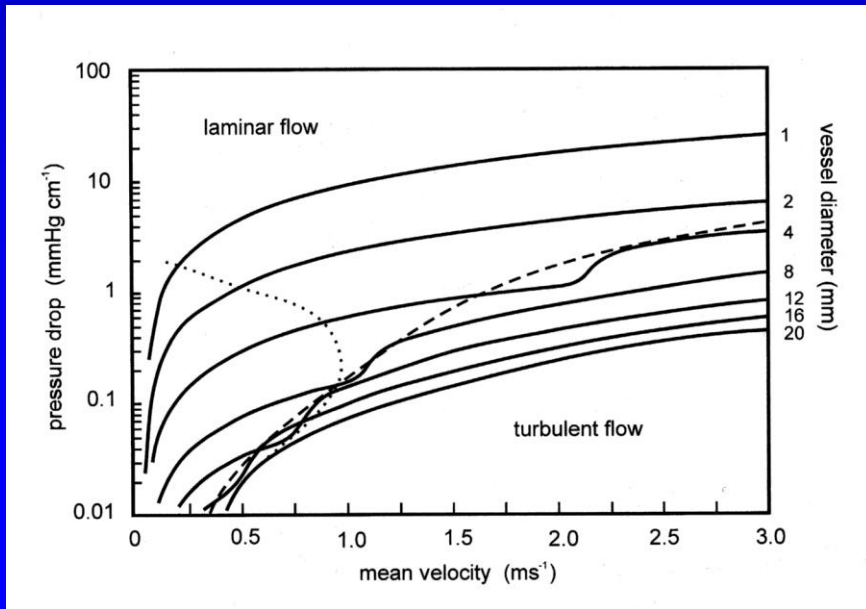


identical CSA, but different significance of stenosis





$$\Delta P = \frac{k\rho(v_s^2 - v_0^2)}{2} = \frac{k\rho Q}{2} \left(\frac{1}{a_s^2} - \frac{1}{a_0^2} \right)$$



Even in the geometrically most “ideal” stenosis, it is impossible to predict the functional severity and influence on blood flow from hydraulic theory

In summary: EVIDENCE-BASED MEDICINE:

- **knowledge if and which lesion(s) is / are responsible for inducible ischemia, is paramount for adequate treatment in the cath.lab**
- *The angiogram (and IVUS!) have fundamental Shortcomings to indicate ischemia correctly*
- *Rationale of Fractional Flow Reserve*

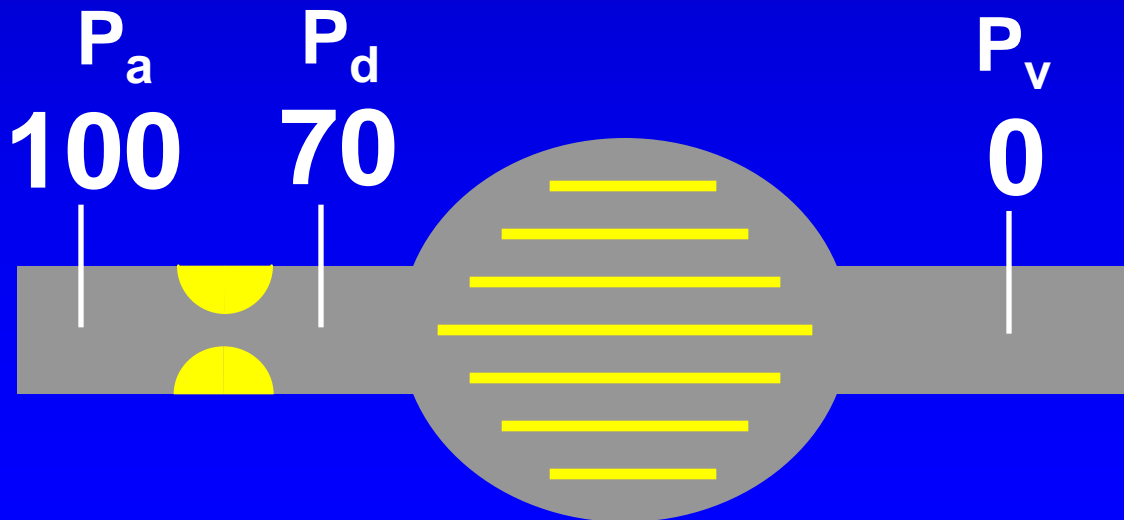
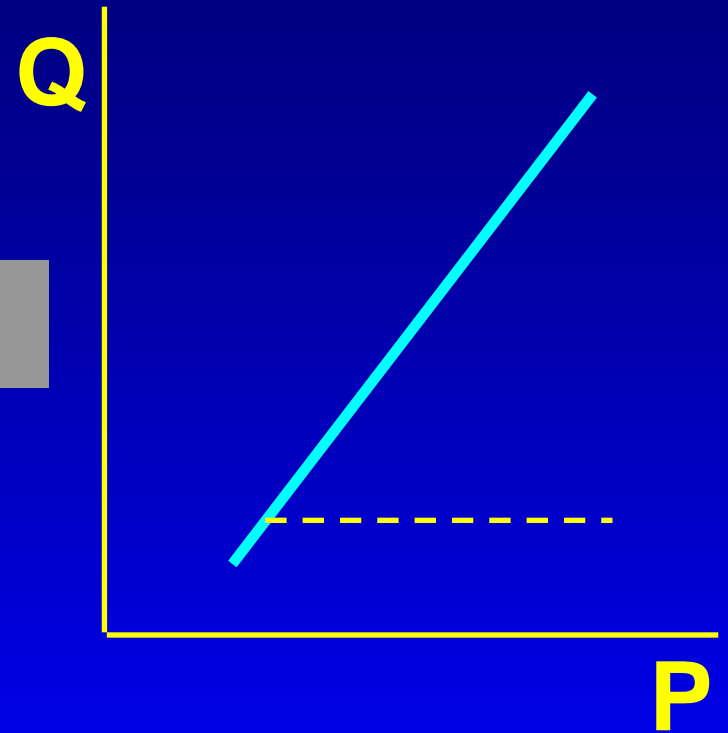
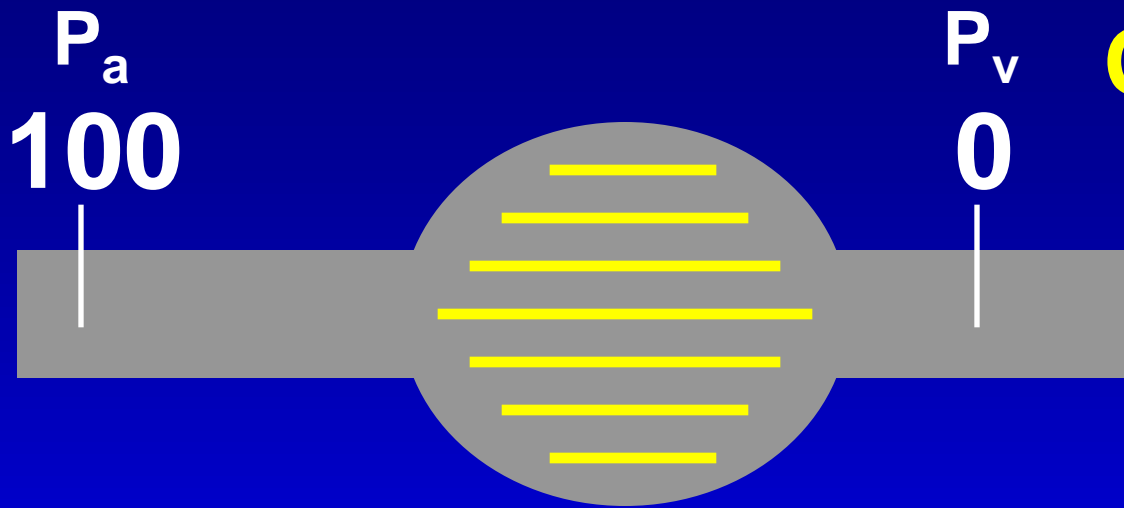
*Whatever the stenosis might look like...,
whatever the pressure/flow relations across
the stenosis might be.....,*

*To understand the meaning of the stenosis for
the patient, the **MOST** important number to know is
the resulting distal perfusion pressure at
hyperemia, as a fraction of normal perfusion
pressure (= aortic pressure)*

*This ratio determines completely the physiologic
significance of the stenosis
and its consequences for the patient !!*

It is called FFR

During Maximal Vasodilatation



$$\text{FFR}_{\text{myo}} = \frac{P_d}{P_a} = 0.70$$

ISSUES TO BE DISCUSSED

- structure of the coronary circulation
- relation between vessel size and perfusion area
- endothelium and development of atherosclerosis
- the 2 or 3 compartment model of the coron circulation
- collaterals
- why functional testing / FFR ?
- **which lesions should be treated**
 - those causing ischemia
- ***ischemia & vulnerability: paradox or antithesis ?***
(Bernard De Bruyne, later today)

ISSUES TO BE DISCUSSED

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 - ***those causing ischemia***
- ischemia & vulnerability: paradox or antithesis ?

Paradox or anthithesis ?

**Excellent outcome of medical treatment in
non-ischemic stenosis
(DEFER study, many non-invasive studies)**

versus

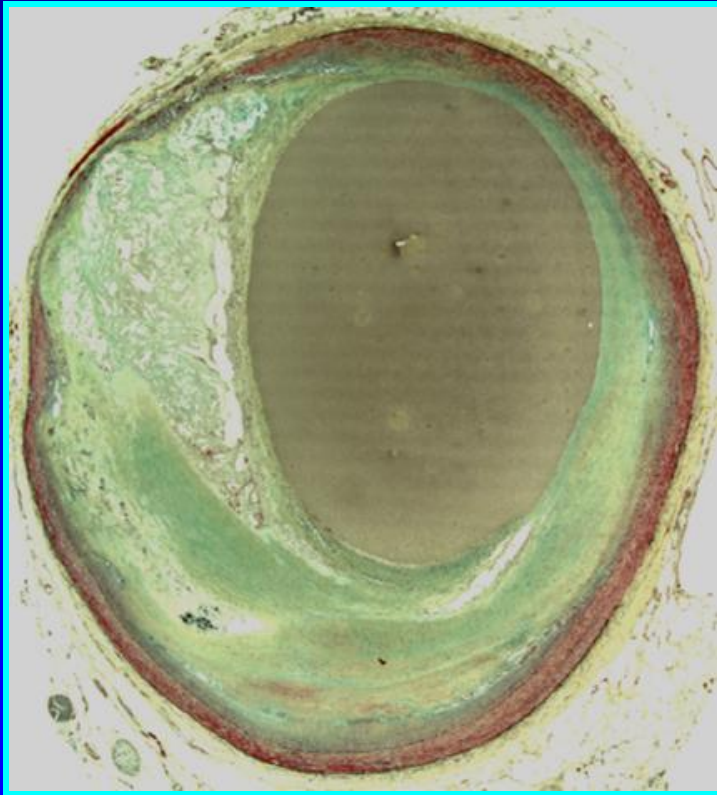
concept of vulnerable plaque

today

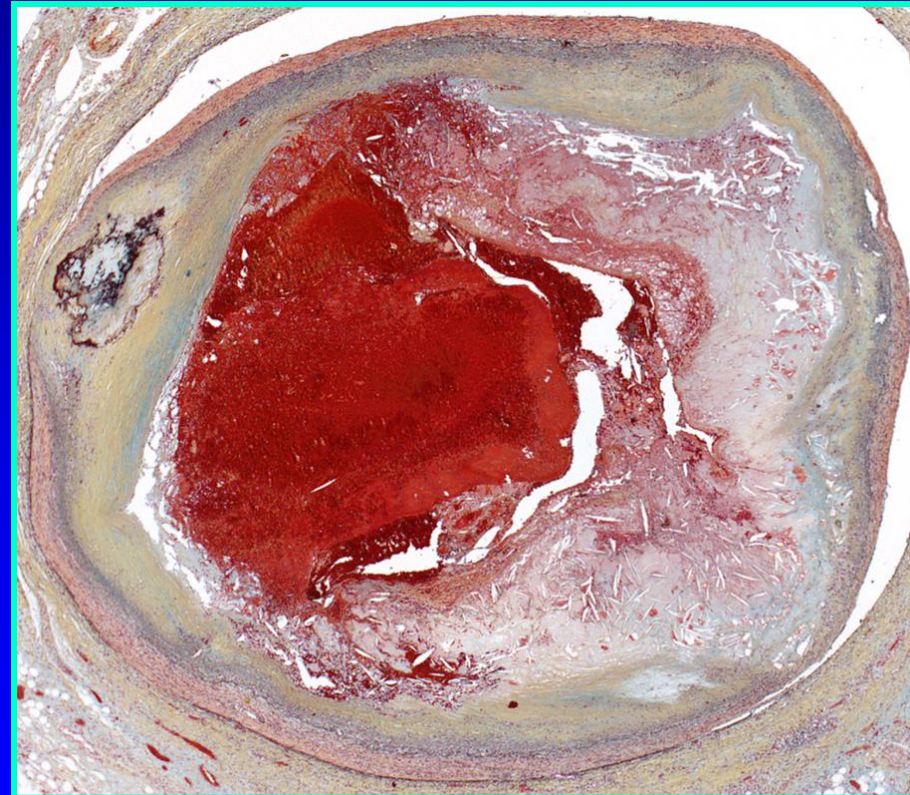


tomorrow

TCFA



Plaque Rupture



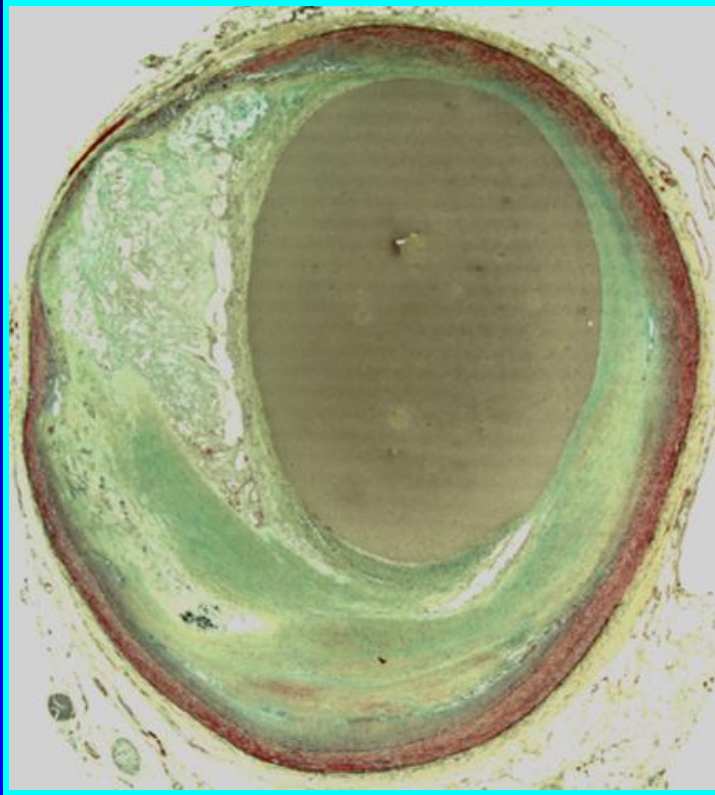
today

?

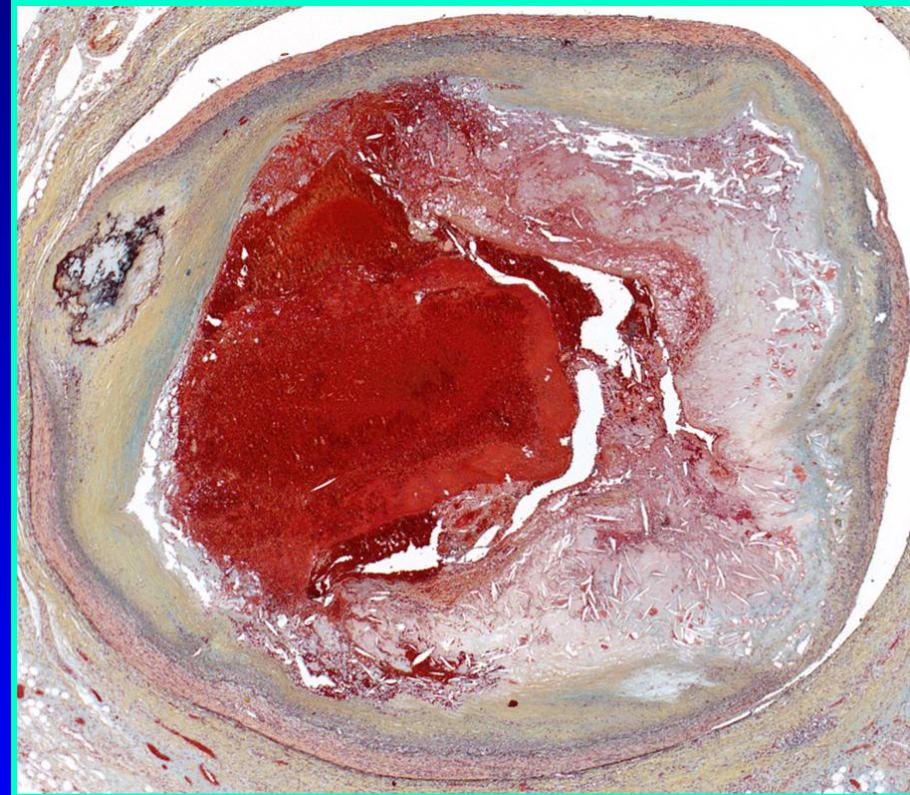
tomorrow

TCFA

Plaque Rupture



?



*Let's look a little bit more critical to such "plaques"....
What are the facts ?? What is the fiction ??*

(Vulnerable) Plaque: Facts and Fiction

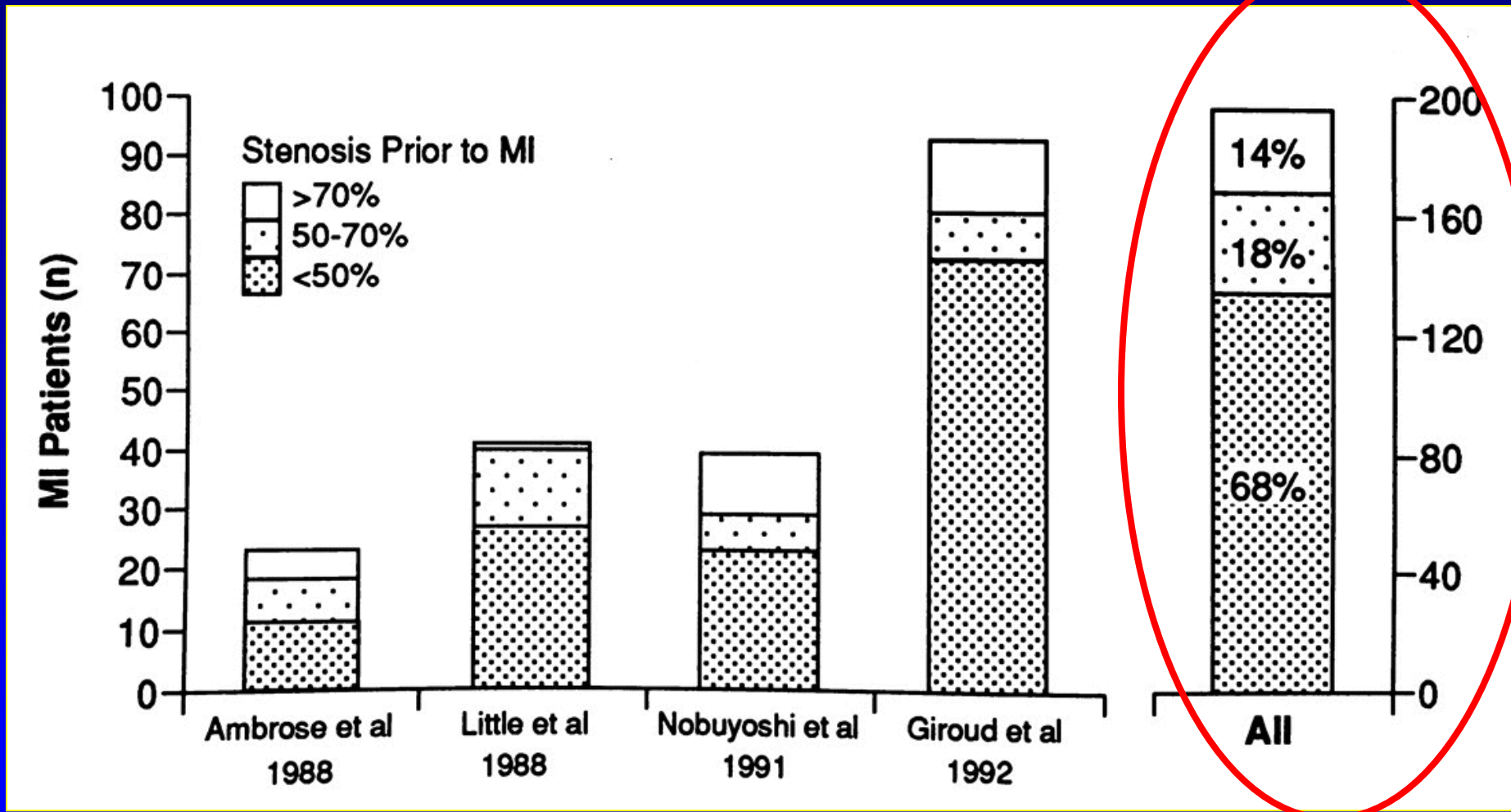
FACTS:

- plaques are very common
- majority of plaques has an excellent prognosis with medical treatment
- only few plaques are vulnerable
- strongest indicator with respect to prognosis is *associated ischemia*

FICTION:

- every plaque is vulnerable
- every vulnerable plaque leads to ACS
- most ACS occurs in mild plaques
- vulnerability can be assessed by imaging

Underlying Stenosis Severity of Abrupt Total Occlusions



Falk, Shah and Fuster, Circulation 1995

“Acute Coronary Syndromes most often occur at the site of mild stenoses”

Do Myocardial Infarctions Evolve from Mild Stenoses ?

Serial Angiographic (Retrospective) Studies in Patients with MI and a Prior Coronary Angiogram

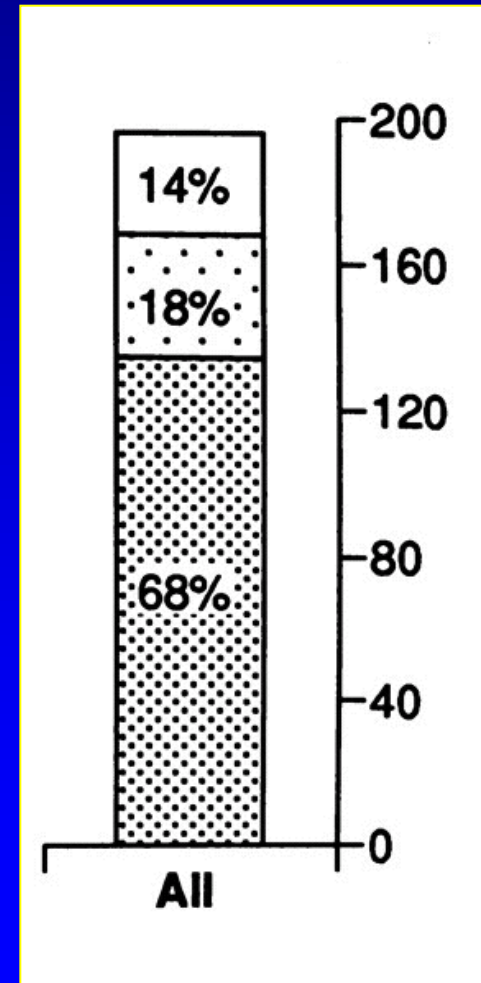
No QCA, No IVUS but unblinded “eyebolling”

	Number of Patients	DelayAngio-MI
Ambrose et al <i>JACC</i> 1988	23	1 month to 7 years
Little et al <i>Circulation</i> 1988	42	4 days to 6.3 years
Giroud et al <i>AJC</i> 1992	92	1 month to 11 years
Moise et al. <i>AJC</i> 1984	116	39 months
Webster et al <i>JACC</i> 1988	30	55 months
Hackett et al <i>AJC</i> 1989	10	21 months

Total

313

**A few days to 11 years
(average 3.9 years !!!)**



THE MYTHE OF THE “DANGEROUS” PLAQUE

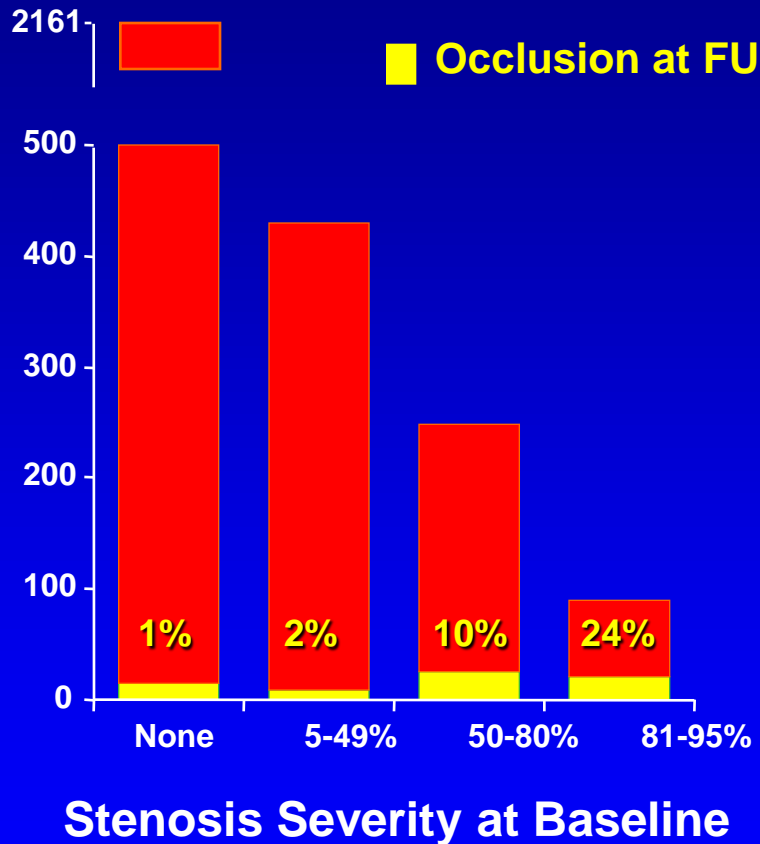
The hypothesis of the occurrence of acute MI on such previously non-significant plaque is based upon

- **6 small retrospective studies**
- **with a total of 313 patients**
- in whom the “index” catheterization was performed an average of **3.9 years** before the acute event

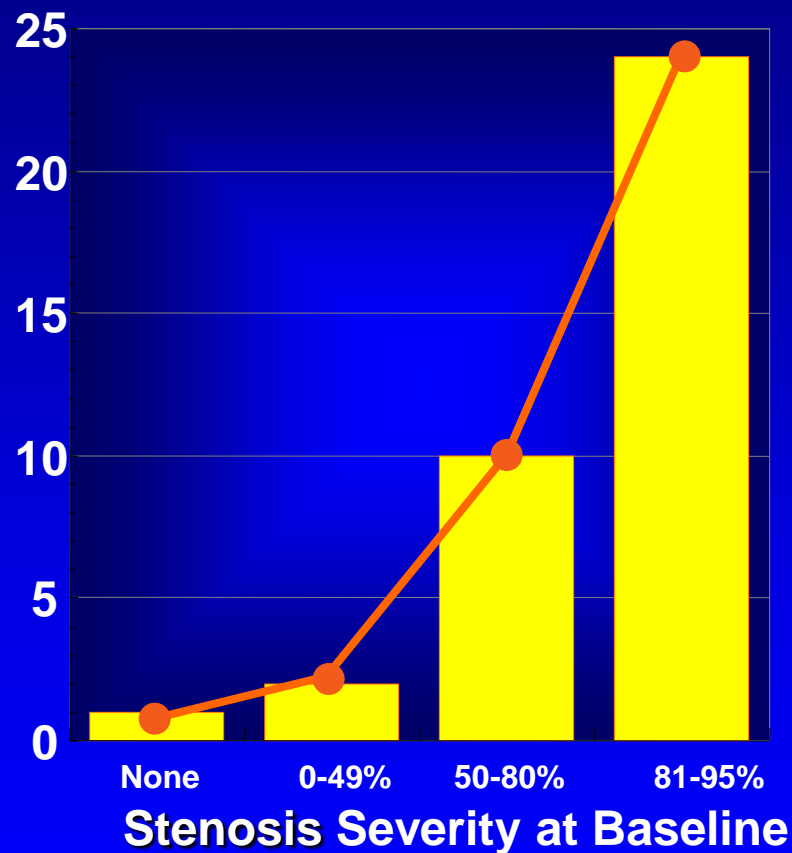
All other literature (21 “meta-analyses” and hundreds of references), refer to these 6 studies !!!

Coronary Occlusion at 5 Years as a Function of Stenosis Severity

Coronary Segments (n)



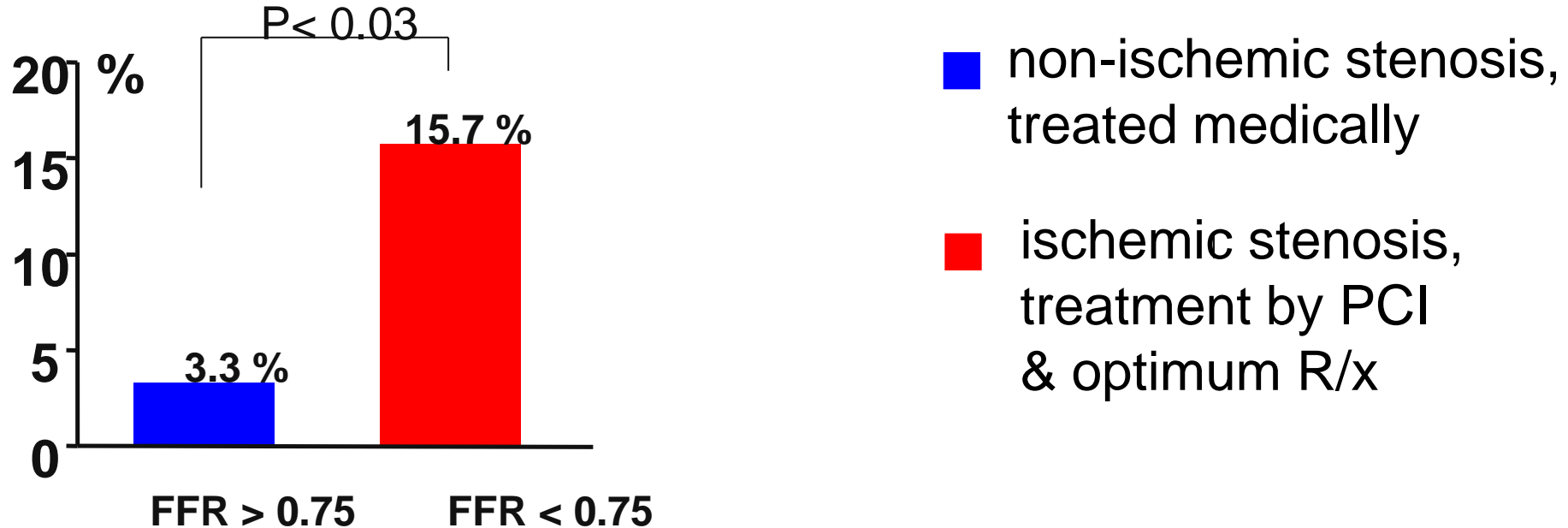
% Occlusion at 5 Year



Adapted from Alderman et al. J Am Coll Cardiol 1993

DEFER study (N=325) :

Cardiac death and Acute MI after 5 years



- ➔ ischemic lesion is much more dangerous than non-ischemic lesion
- ➔ risk of individual non-ischemic lesion to cause death or AMI, is very small and < 1 % per year !!

250 consecutive patients with ST-elevation MI
in the Catharina Hospital:

- underlying stenosis angiographically significant in 92 % of the cases
- *At meticulous anamnesis, 80 % of patients had recurrent chest pain in the year before the acute myocardial infarction occurred !!*

INCIDENCE OF CORONARY STENOSIS IN A GENERAL POPULATION

**Incidence of coronary artery disease in
asymptomatic, apparently healthy persons**

> 50 years old : 25%

> 60 years old : 40%

Sims et al, Am Heart J 1983

Maseri, Ischemic Heart Disease 1995

What about the prognosis of these patients ?

→ Related to inducibility of ischemia

- structure of the coronary circulation
- relation between vessel size and perfusion area
- endothelium and development of atherosclerosis
- the 2 or 3 compartment model of the coron circulation
- collaterals
- why functional testing / FFR ?
- which lesions should be treated
- vulnerable plaques: facts & fiction
- ***ischemia & vulnerability: paradox or antithesis ?***

“The missing link”

Is there a link between vulnerability and ischemia ?

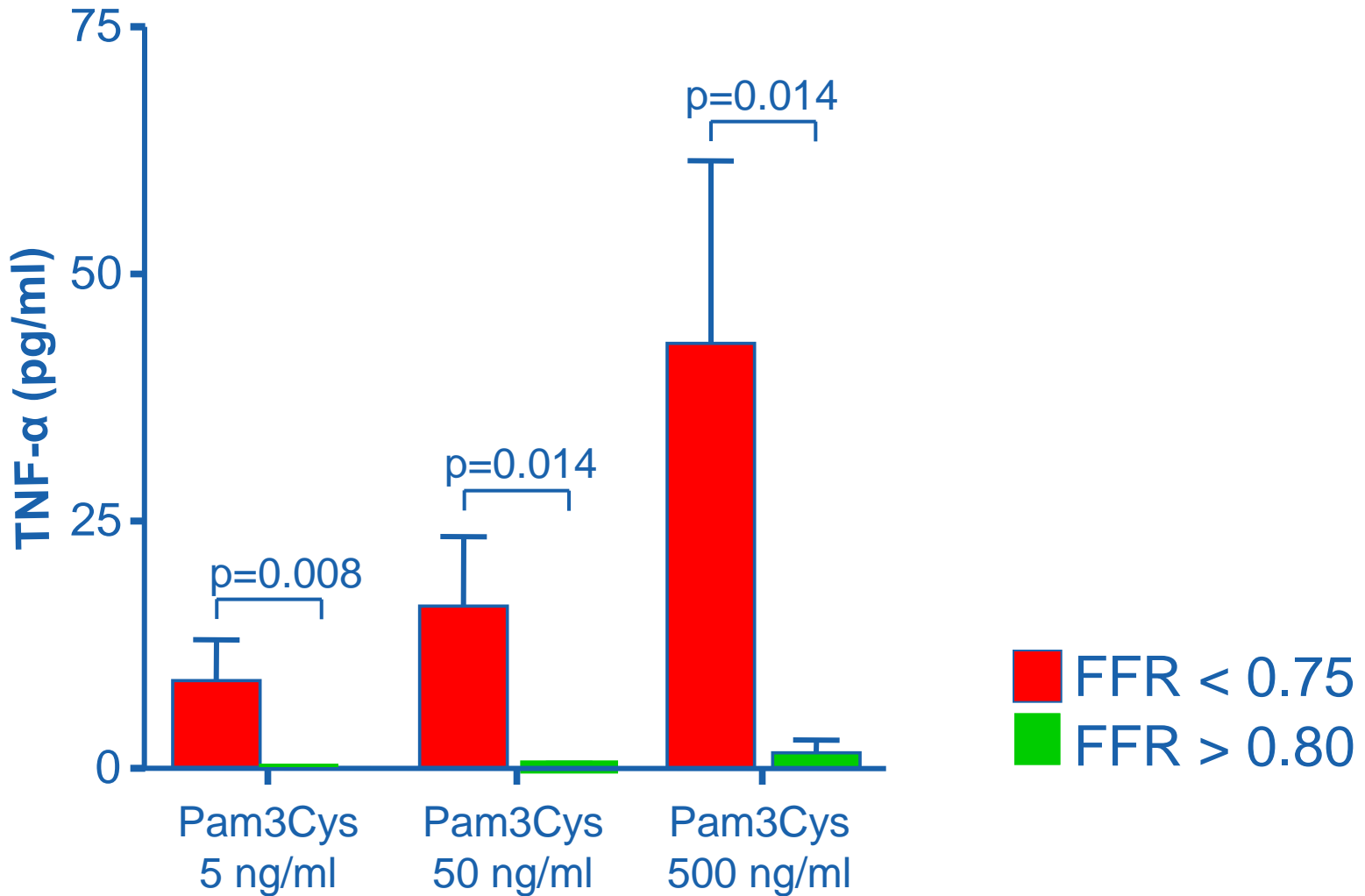
Hypothesis:

- repetitive ischemia *and*
- high shear stress / pressure gradients

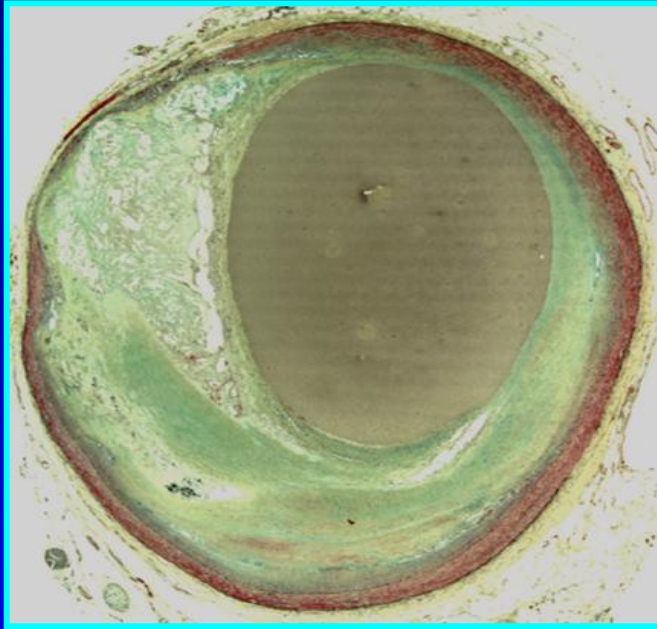
induce vulnerability

→ *Supported by studies on the relation between vulnerability markers and low FFR:
on-going work of Pasterkamp et.al.*

TLR2 stimulation (Pam3Cys)



Concept of Yesterday:

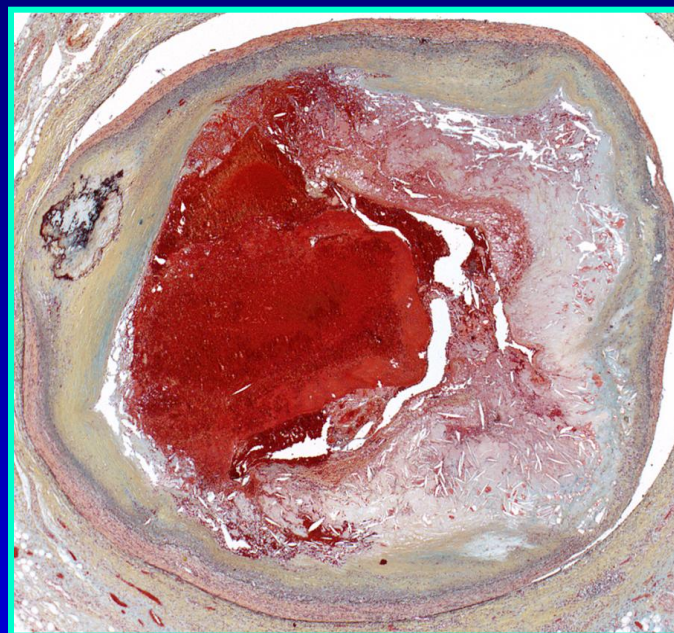
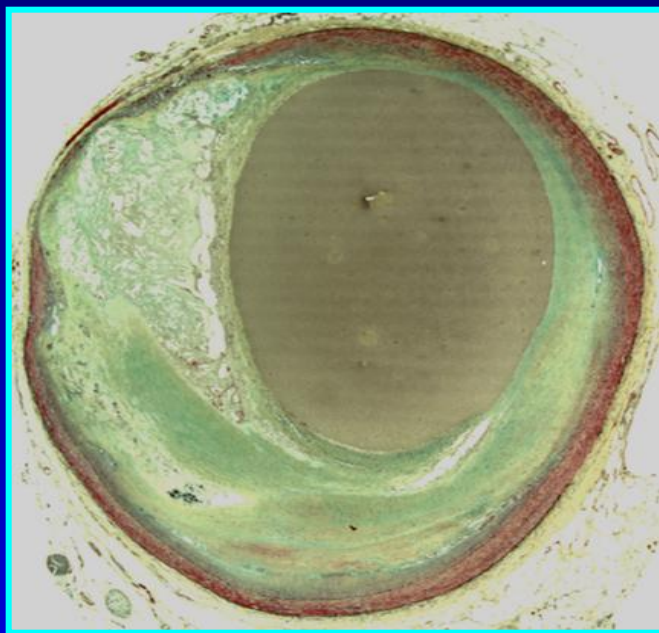


Pro-inflammatory cytokines, activated monocytes, etc



Vulnerability
(“out of the blue”)

Concept of Tomorrow:



ischemic episodes

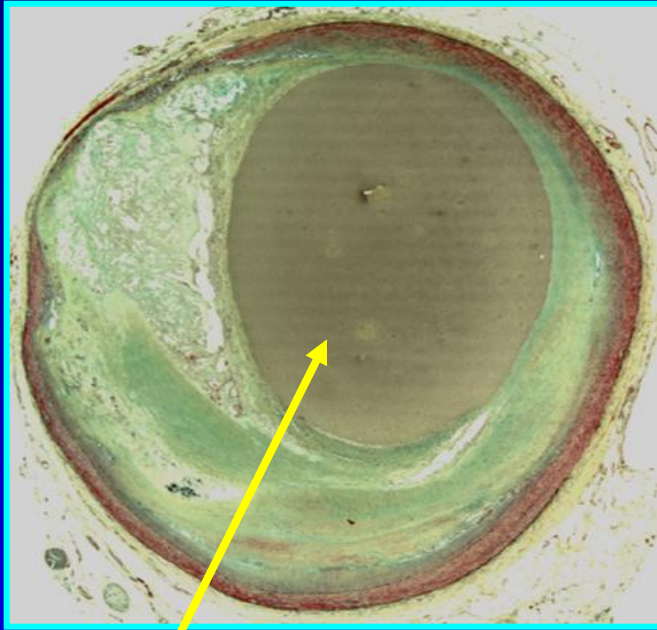


Pro-inflammatory cytokines, activated monocytes, etc



Vulnerability

Concept of today:



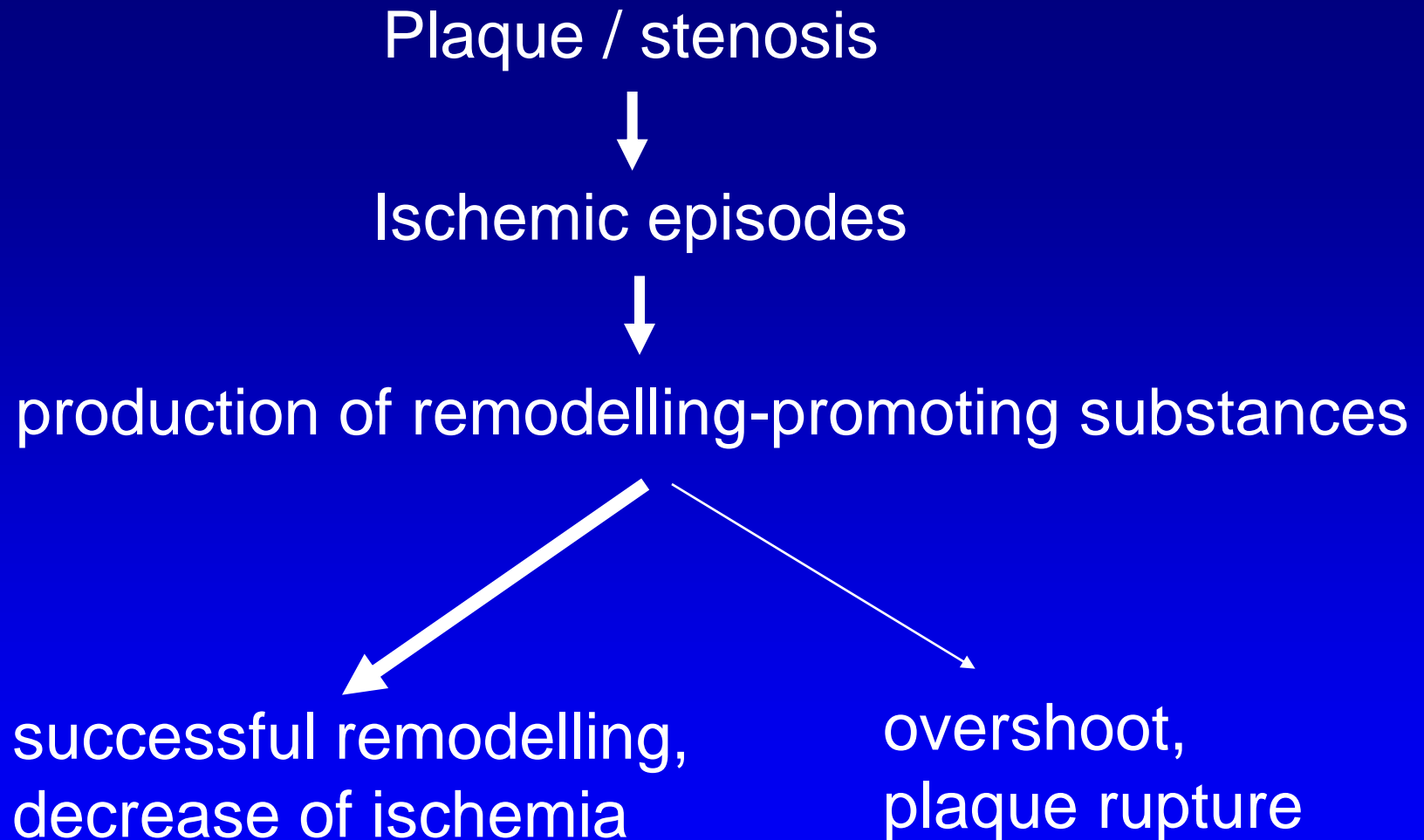
by the way:
70% area
Stenosis !!

ischemic episodes

Pro-inflammatory cytokines etc

Vulnerability

new paradigm:



Searching for vulnerability starts with searching for ischemia

Suppose aliens would visit us and would like to investigate the determinants of a fire.



“Substance X (also called “water”) must be dangerous substance !”

FUNCTIONAL ASSESSMENT OF BOTH COMPARTMENTS TOGETHER:

- non-invasively
(exercise testing, stress echo, Mibi)
- invasively: intracoronary Doppler, absolute flow

FUNCTIONAL ASSESSMENT OF THE MICROCIRCULATION:

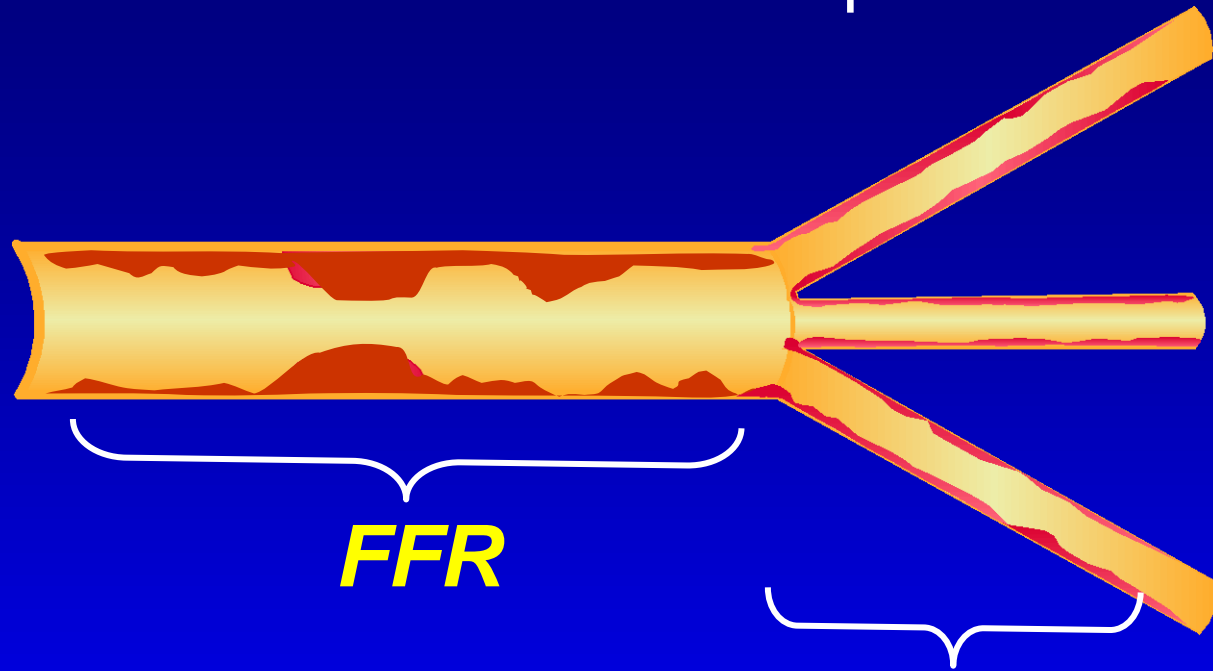
- Index of Microcirculatory Resistance (IMR)

The coronary microcirculation:

Still a black box ??

focal and diffuse
Epicardial disease

microvascular
compartment



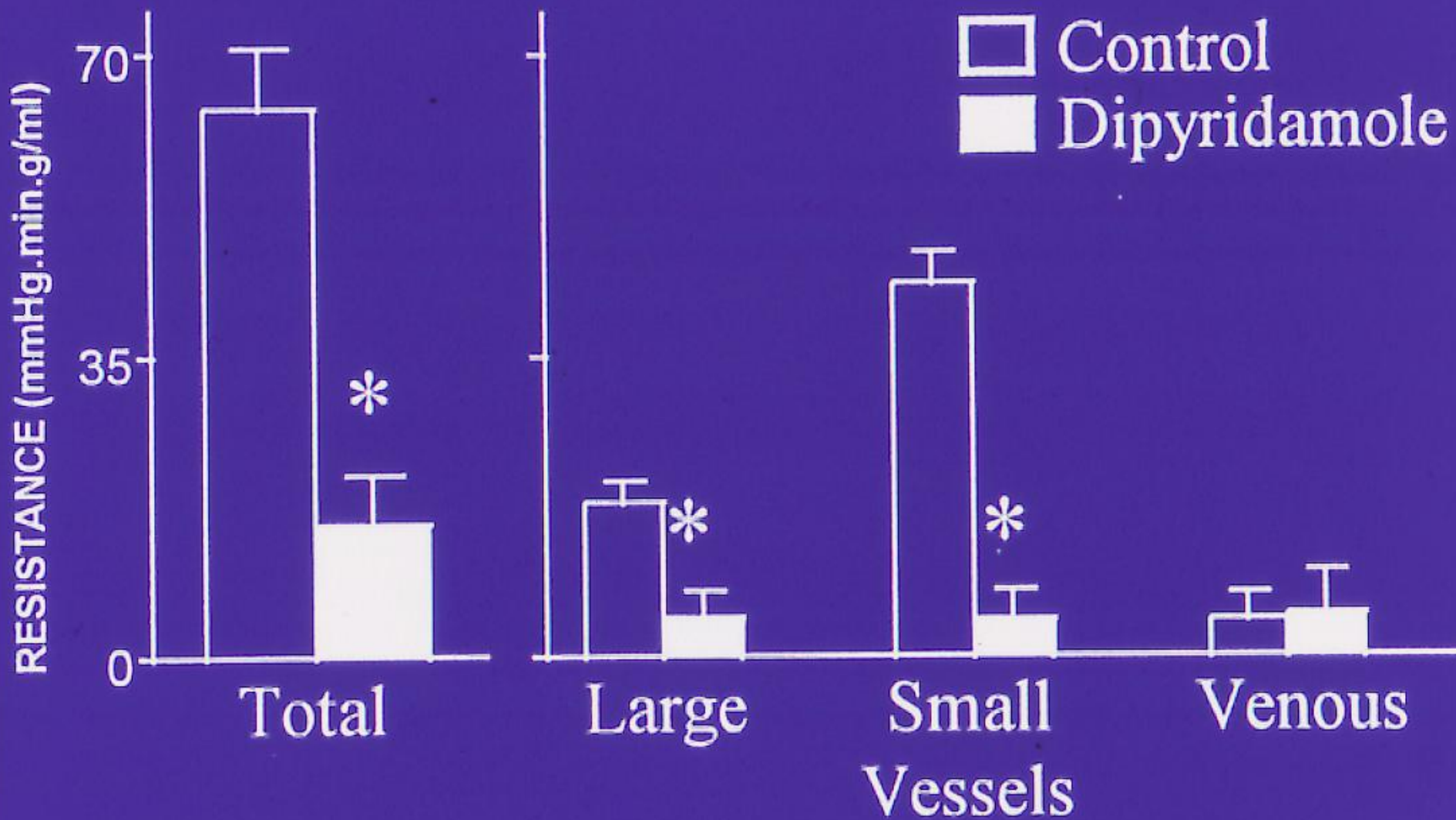
Specific indexes ??

→ **Invasive indexes** (saturday morning):

IMR (*Bill Fearon*)

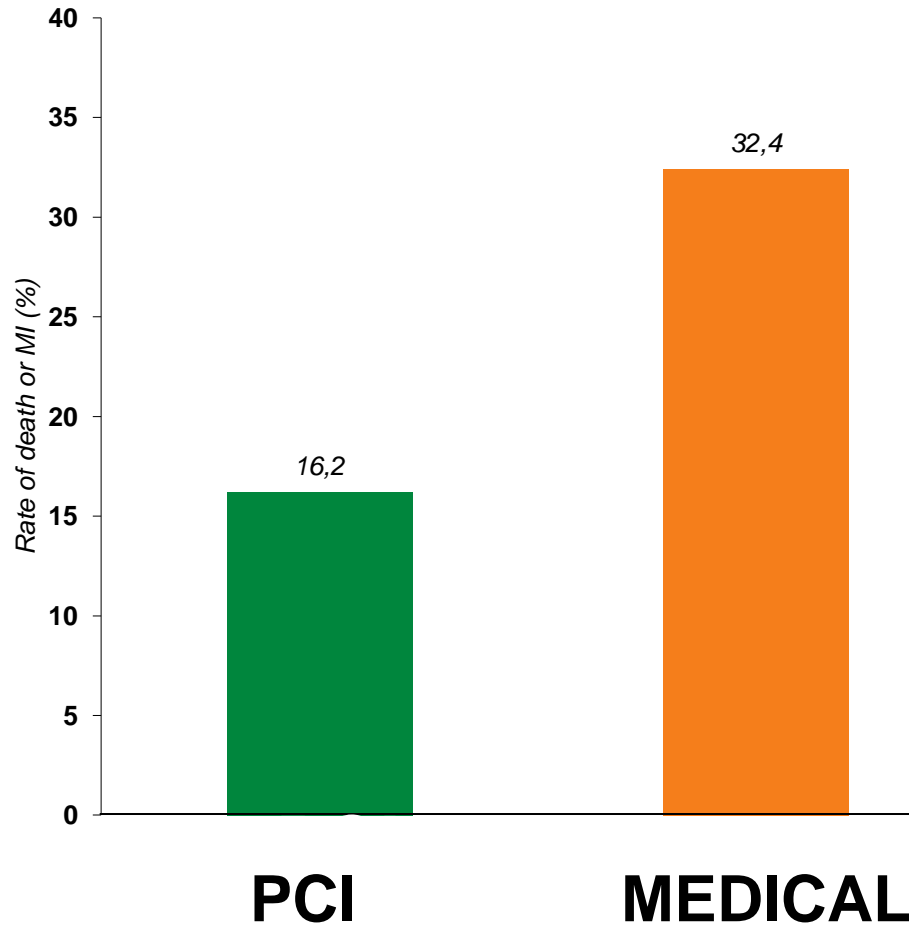
absolute resistance (*Nico Pijls*)

! We cannot understand the physiologic significance of a stenosis without taking into account the distal perfusion territory !

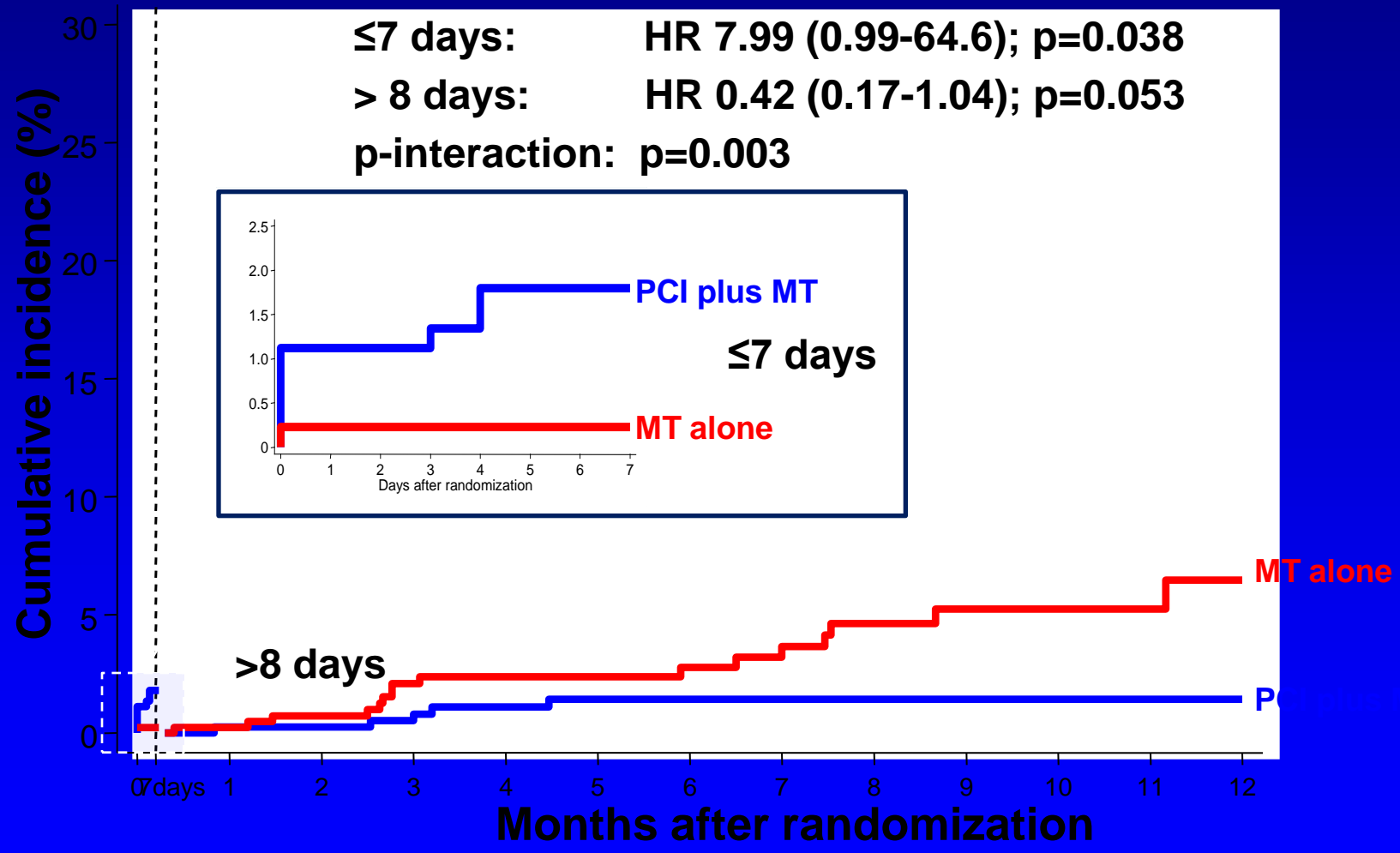


majority of resistance located in arterioles (100-400 μm)

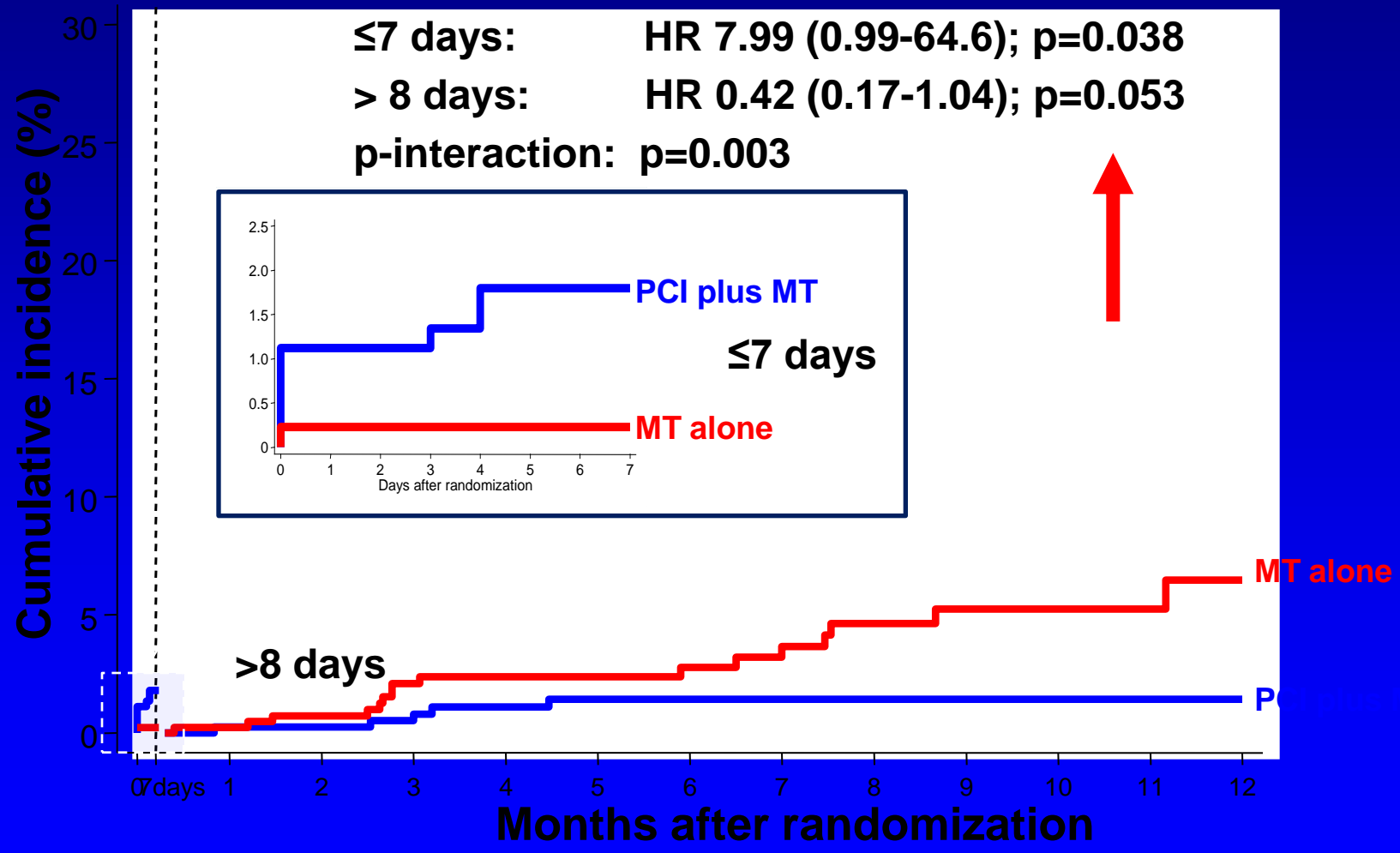
ISCHEMIC



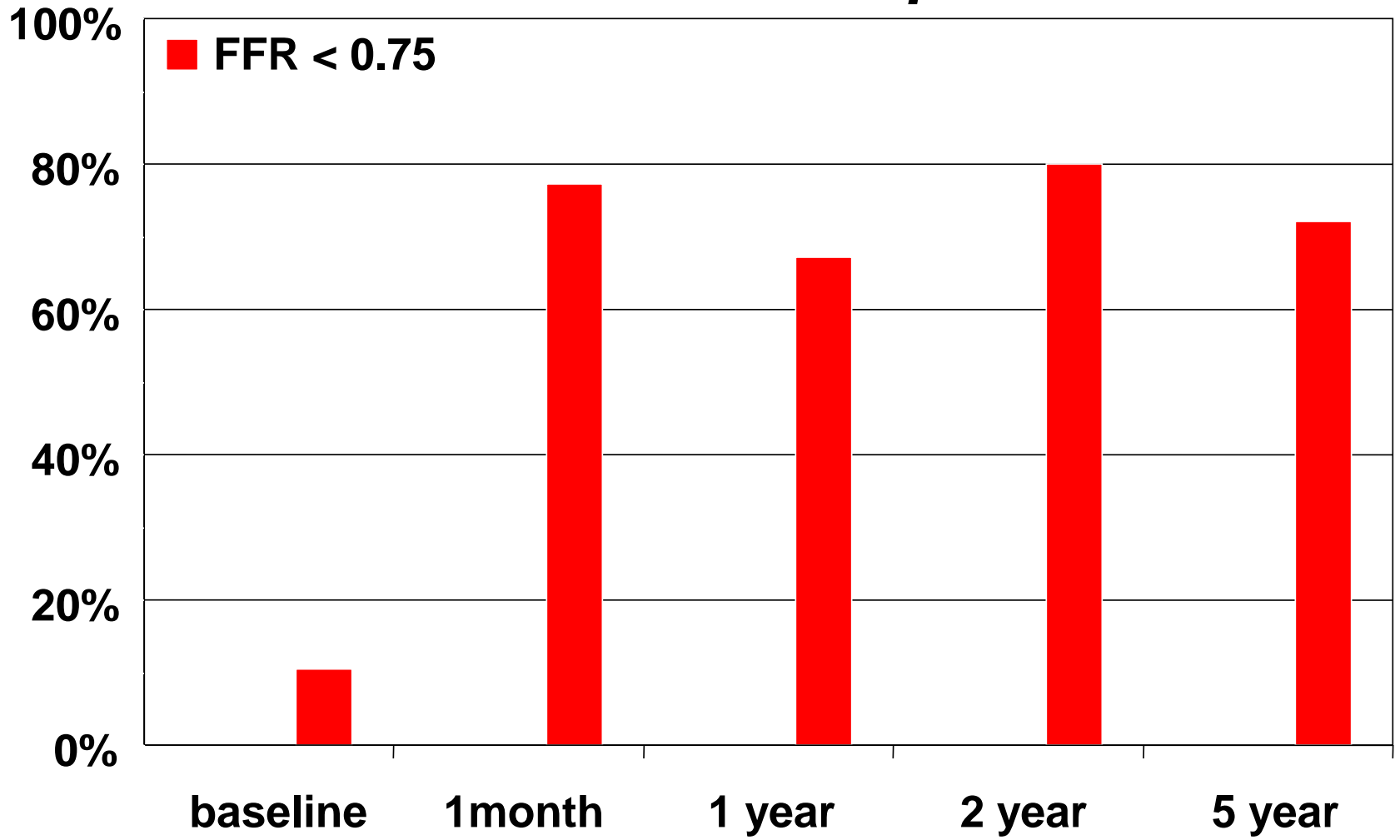
Kaplan-Meier plots of Landmark Analysis of Death or MI



Kaplan-Meier plots of Landmark Analysis of Death or MI

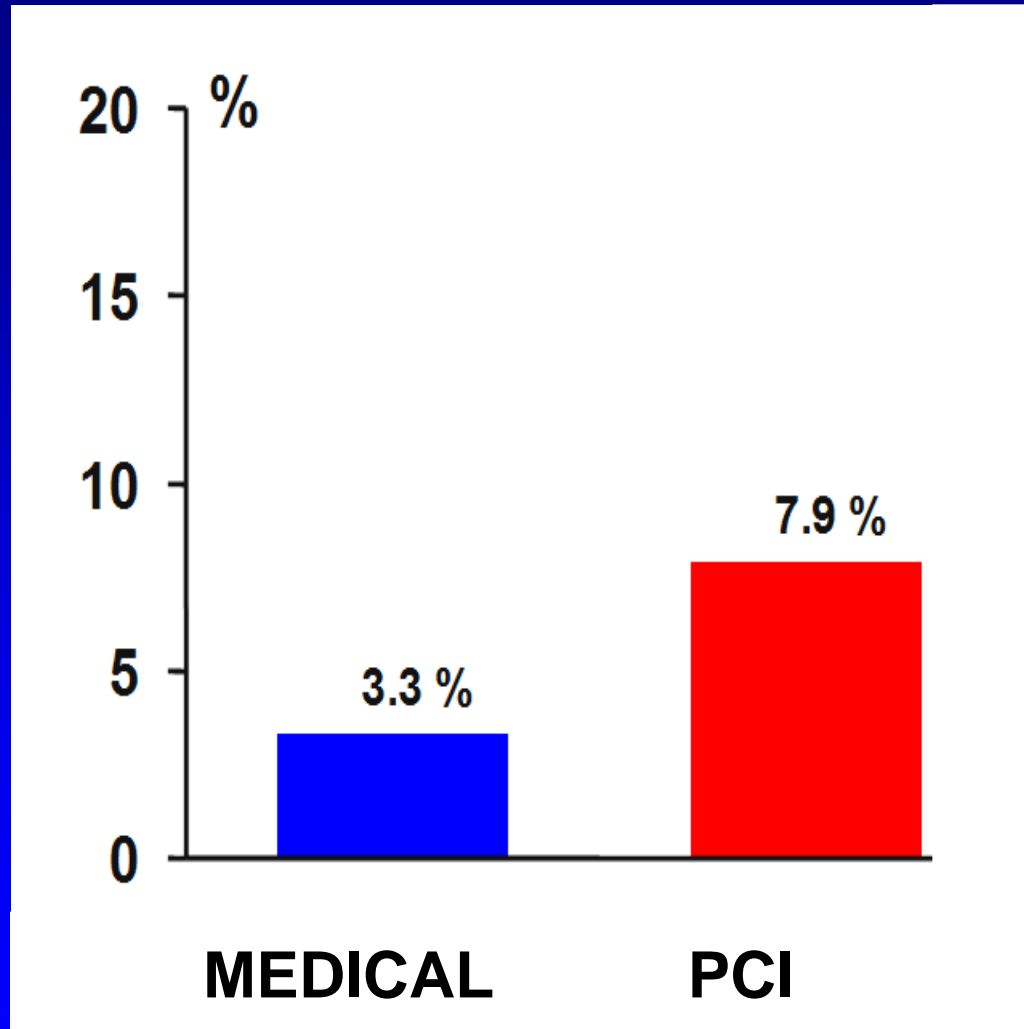


Patients with proven ischemia



freedom from angina after stenting ischemic stenosis

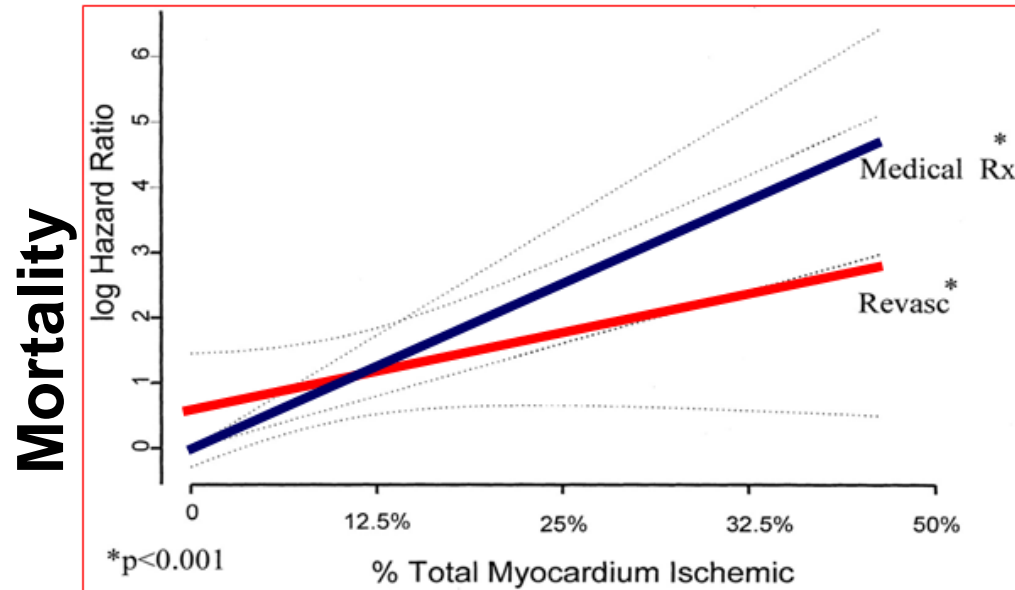
Death & MI 5 during 5 years of follow-up after PCI vs Medical Treatment in NON-ischemic stenosis



*Pijls et al
JACC 2007*

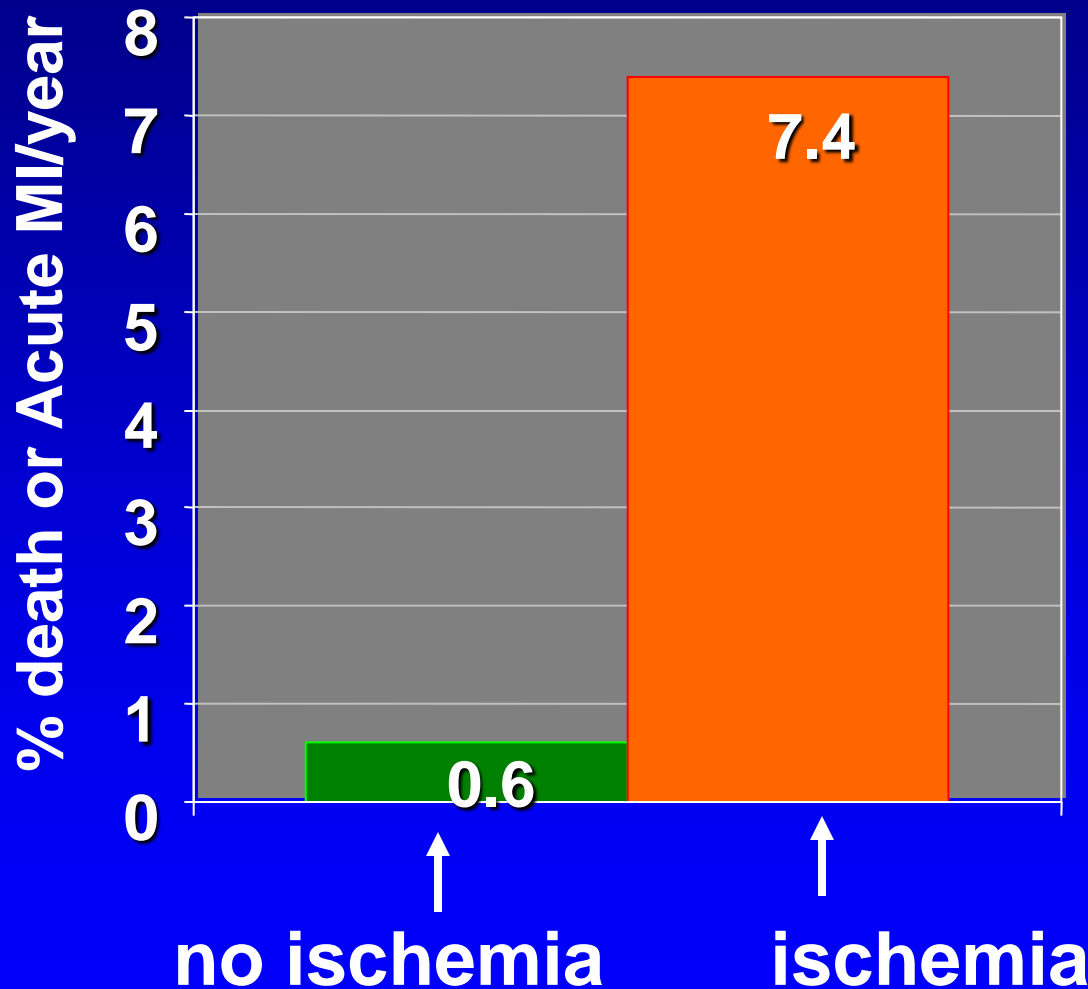
Is it important to detect ischemia ?

Log hazard ratio for revascularization (Revasc) vs medical therapy (Medical Rx) as a function of % myocardium ischemic based on final Cox proportional hazards model



Above 10% ischemic myocardium, the survival benefit from revascularisation increases with the extent of ischemia

The risk for death or acute myocardial infarction in the next five years is 20 times higher for an ischemic lesion compared to a non-ischemic lesion !!!



**12000 Patients
(2 x 6000)**

**similar stenosis
severity by
coronary angio**

***Risk to die or experience myocardial infarction
in the next 5 years related to a coronary stenosis:***

- **non-ischemic stenosis: < 1% per year ***
(NUCLEAR studies, PET, MRI, DEFER, FAME)
- **ischemic stenosis, if left untreated: 5-10% per year**
(Many historical registries, nuclear studies, ACIP, CCTA, MRI, FFR)
- **stented stenosis: 2-3% per year**
(e.g DEFER, FAME, SYNTAX, many large studies and registries)

***THE KEY ISSUE IN INTERVENTIONAL
CARDIOLOGY IS TO DISCRIMINATE
THOSE LESIONS RESPONSIBLE FOR
INDUCIBLE ISCHEMIA***

→ Fractional Flow Reserve

THE EPICARDIAL COMPARTMENT IS RATHER EASY TO ASSESS:

IMAGING OF THE EPICARDIAL COMPARTMENT

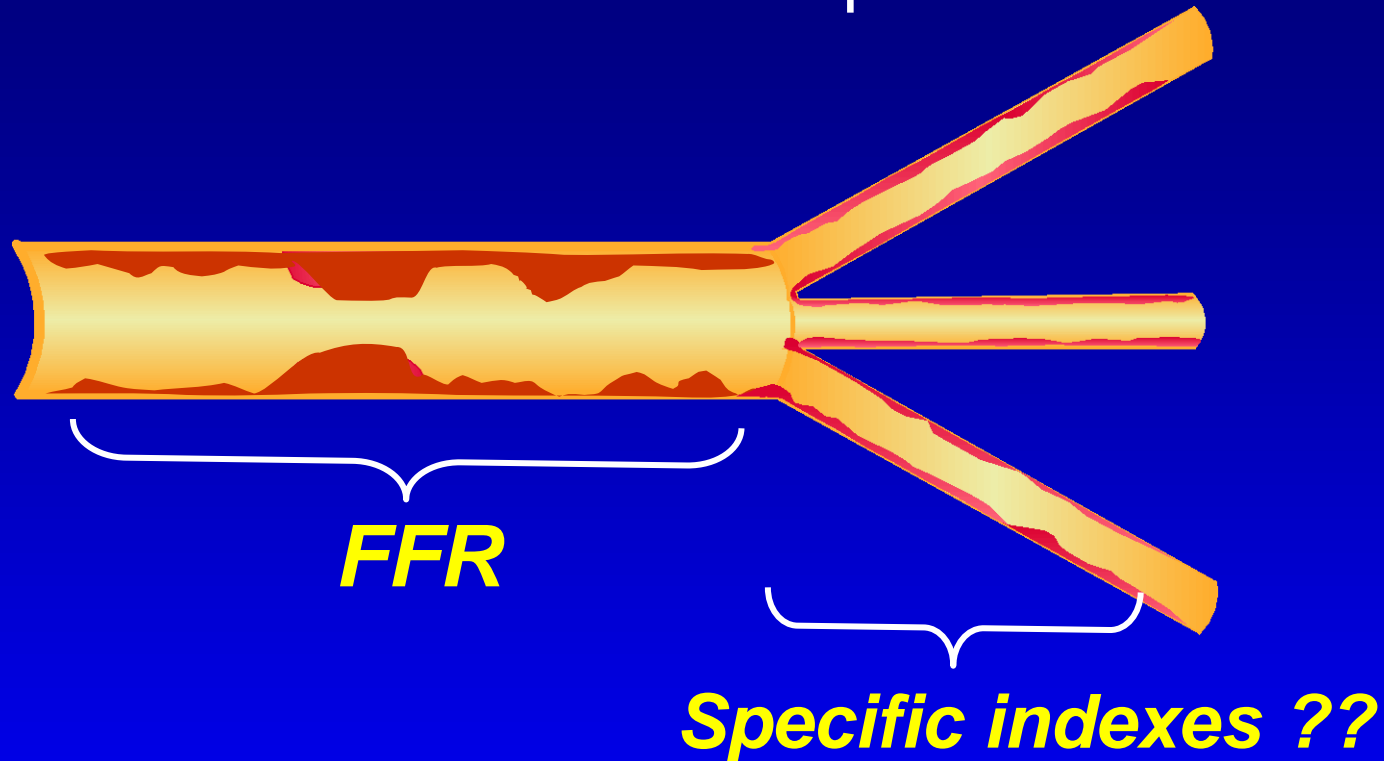
- non-invasively by CT, MRI
- invasively by angio, IVUS, OCT, and some newer techniques

FUNCTIONAL ASSESSMENT OF THE EPICARDIAL COMPARTMENT

- coronary pressure & FFR

focal and diffuse
Epicardial disease

microvascular
compartment



→ **Invasive indexes:**

IMR (*Bill Fearon, Bernard De Bruyne*)

absolute flow & resistance (*Gabor Toth, Inge wijnbergen*)