



Leading  
European  
Nephrology



Leading European Nephrology



## The 23rd Budapest Nephrology School

Nephrology, Hypertension, Dialysis,  
Transplantation, Nephropathology

Under the Auspices of  
**ISN, ERA-EDTA, RPS, IFKF and ISP**

**26–31 August, 2016**

# Chronic consequences of kidney ischemia

## Prof. Andrzej Wiecek

Department of Nephrology, Transplantation and Internal Medicine  
Medical University of Silesia, Katowice, Poland

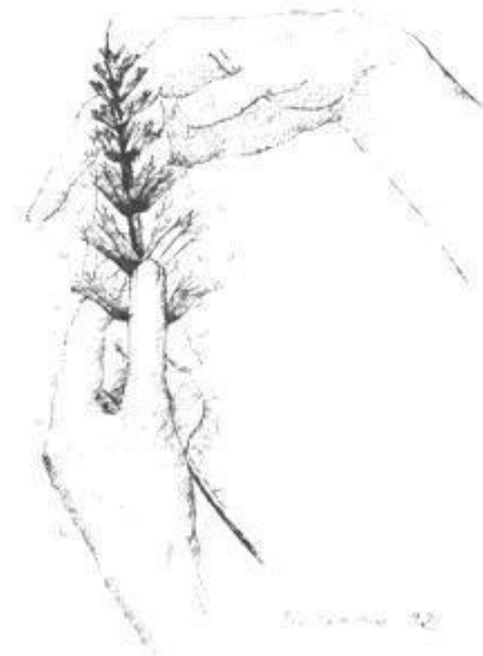
**e-mail: [awiecek@sum.edu.pl](mailto:awiecek@sum.edu.pl)**



Leading  
European  
Nephrology



Leading European Nephrology



# The 23rd Budapest Nephrology School

Nephrology, Hypertension, Dialysis,  
Transplantation, Nephropathology

Under the Auspices of  
**ISN, ERA-EDTA, RPS, IFKF and ISP**

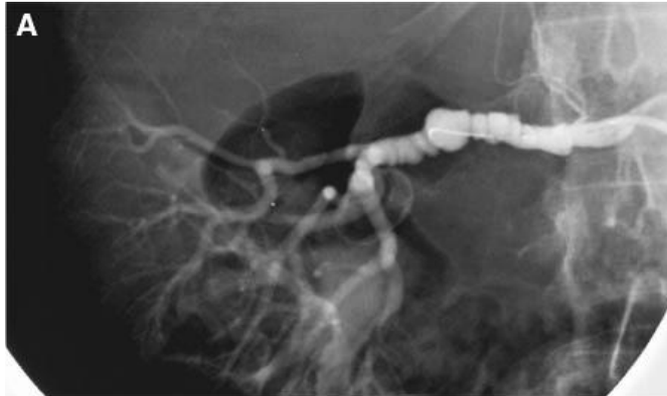
**26–31 August, 2016**

I have no relevant financial relationship to disclose

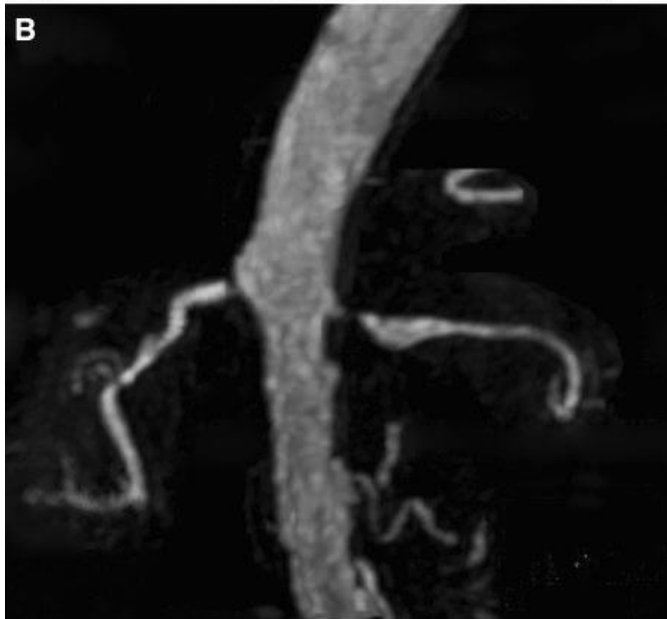
**Andrzej Wiecek**



## Two forms of renal artery lesions



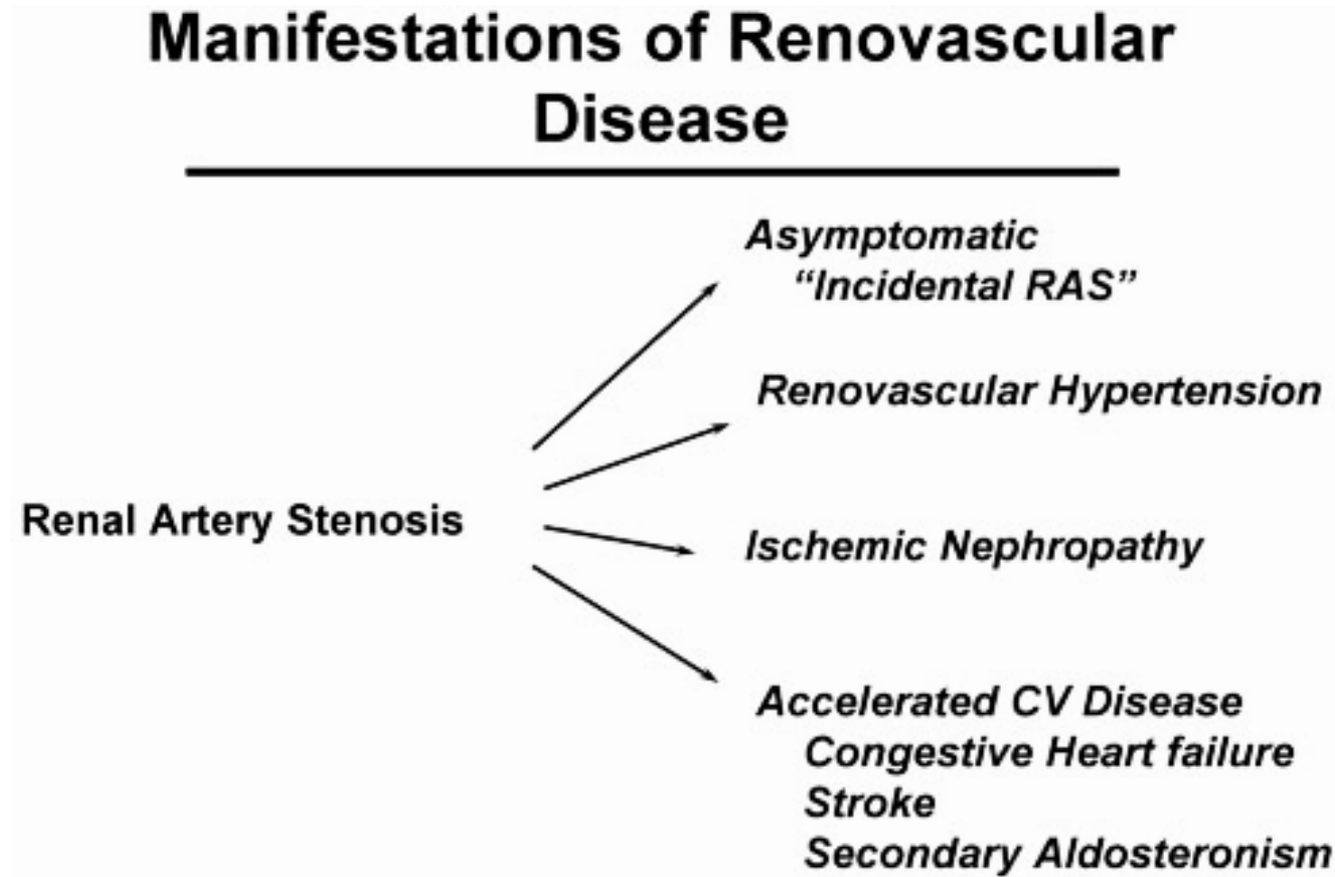
**A**, Angiogram from a patient with FMD with lesions characteristic of medial fibroplasia. The “string-of-beads” appearance typically develops in the mid portion of the vessel from circumferential webs within the vessel. These lesions may progress, particularly in smokers.



**B**, MRA from an individual with atherosclerotic disease affecting the renal arteries. These lesions commonly arise near the ostium of the vessel and may be an extension of aortic plaques.

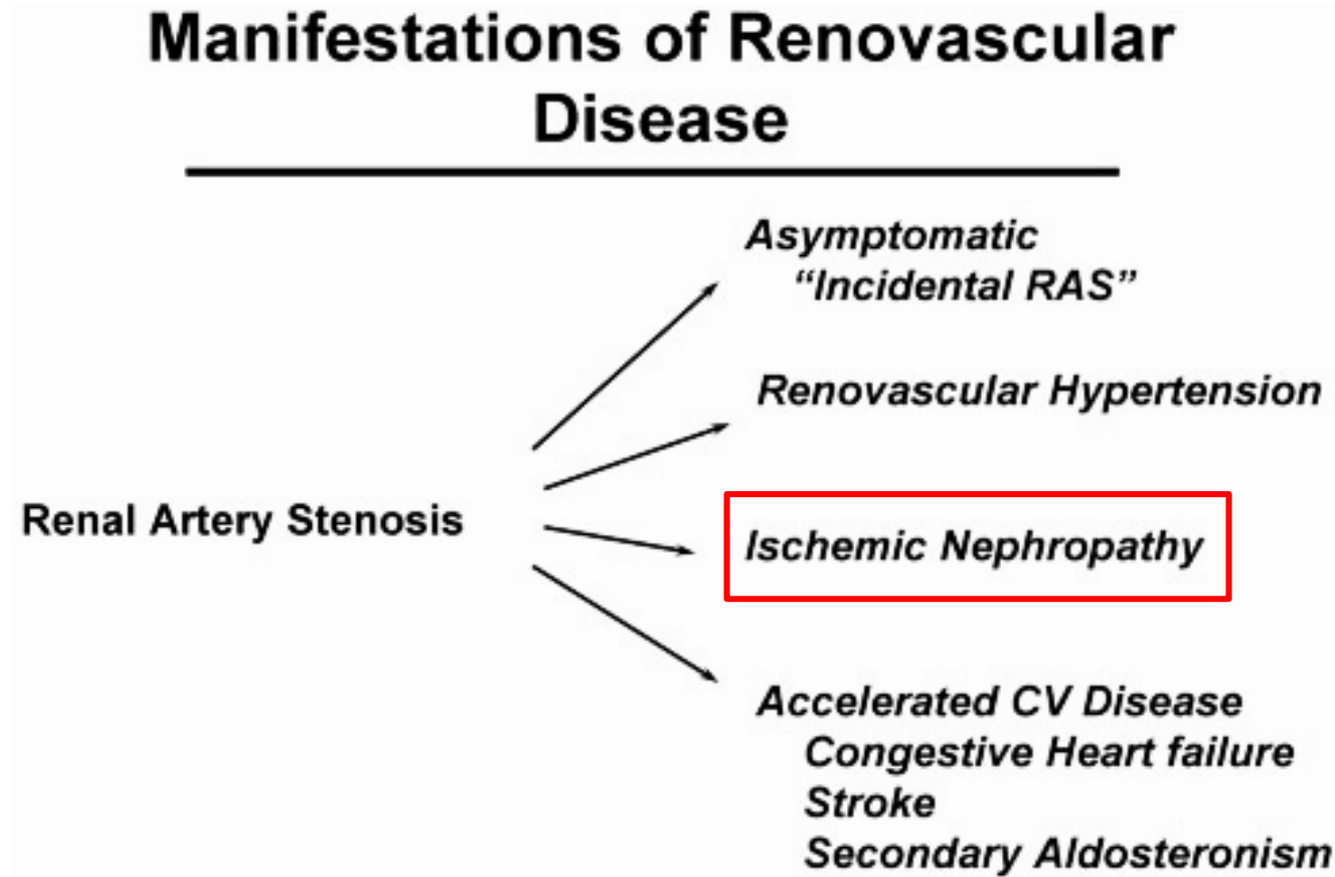


# Schematic summary of the clinical manifestations of renovascular disease

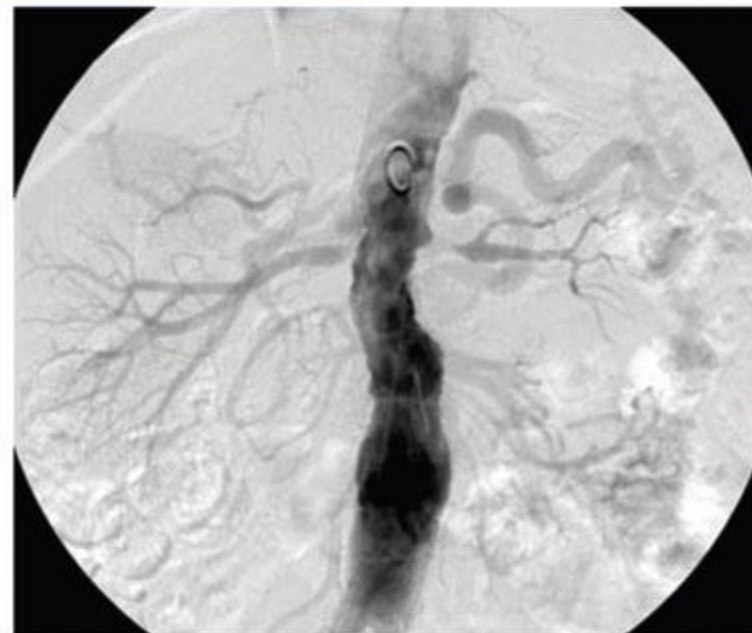
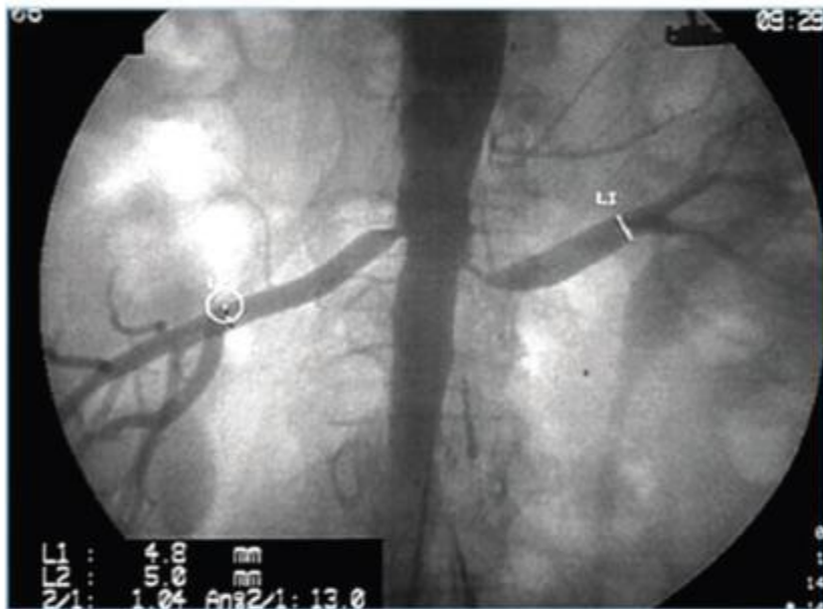




# Schematic summary of the clinical manifestations of renovascular disease



## Spectrum of Renovascular Disease Manifestations

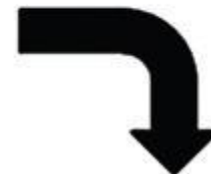


*Asymptomatic  
"Incidental RAS"*

*Renovascular  
Hypertension*

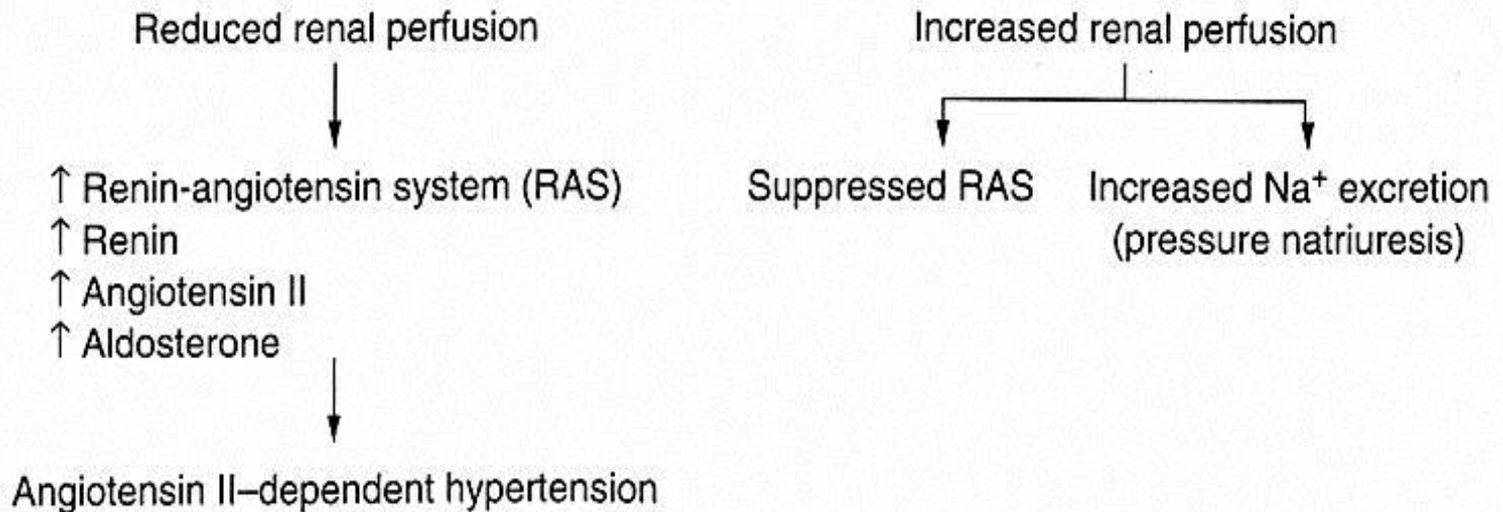
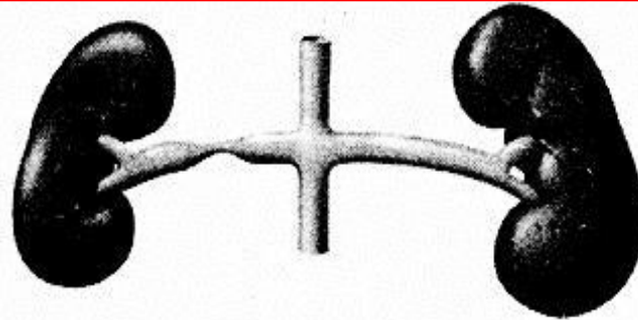


*Accelerated CV Disease  
Congestive Heart failure  
Stroke*



*Ischemic Nephropathy*

# UNILATERAL RENAL ARTERY STENOSIS

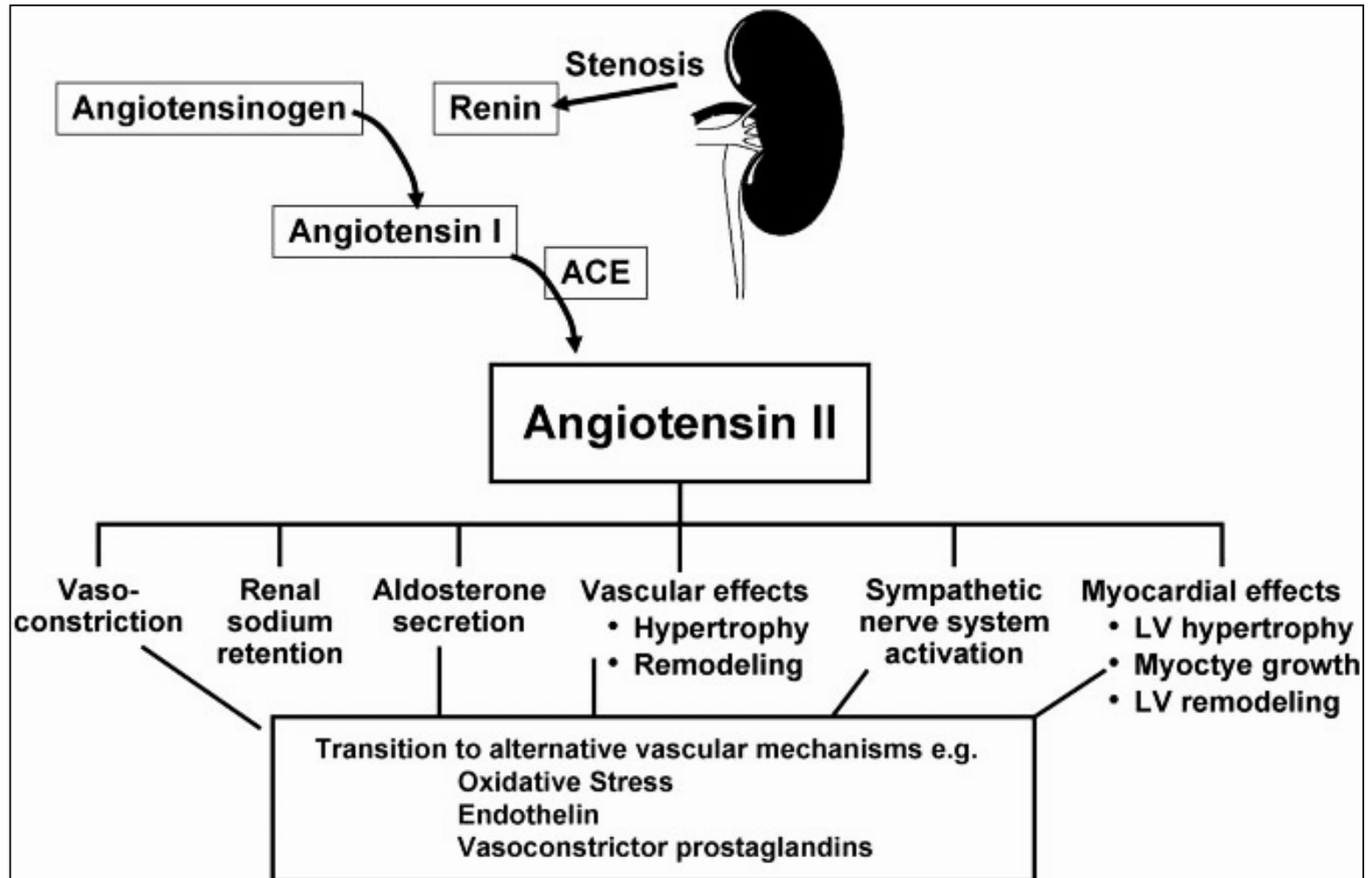


*Effect of blockade of RAS*  
Reduced arterial pressure  
Enhanced lateralization of diagnostic tests  
Glomerular filtration rate (GFR) in stenotic kidney may fall

*Diagnostic tests*  
Plasma renin activity elevated  
Lateralized features, e.g., renin levels in renal veins, captopril-enhanced renography

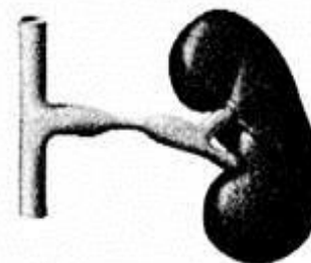
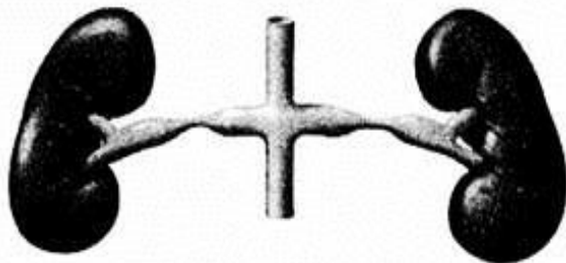


# Pressor mechanisms identified in renovascular hypertension





# BILATERAL RENAL ARTERY STENOSIS



Bilateral

Stenosis of solitary kidney

Reduced renal perfusion

↑ Renin-angiotensin system (RAS)  
 ↑ Renin  
 ↑ Angiotensin II  
 ↑ Aldosterone

Impaired Na<sup>+</sup> and water excretion

Inhibit RAS

Volume expansion

Normal or low angiotensin II

Increased arterial pressure

*Effect of blockade of RAS*

Reduced arterial pressure only after volume depletion  
May lower GFR

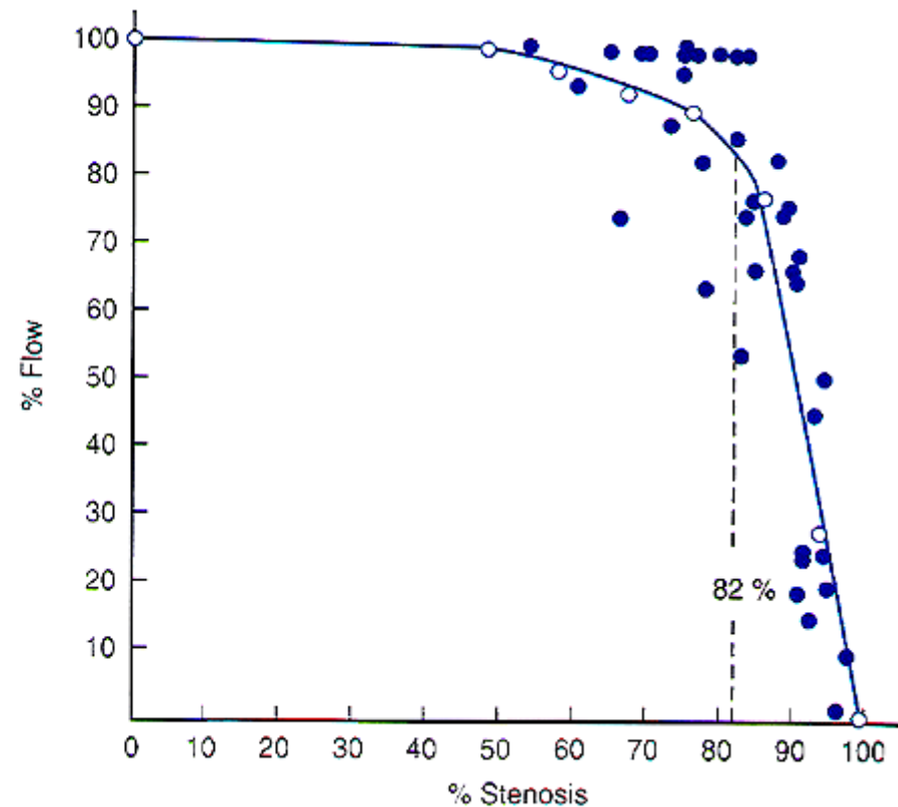
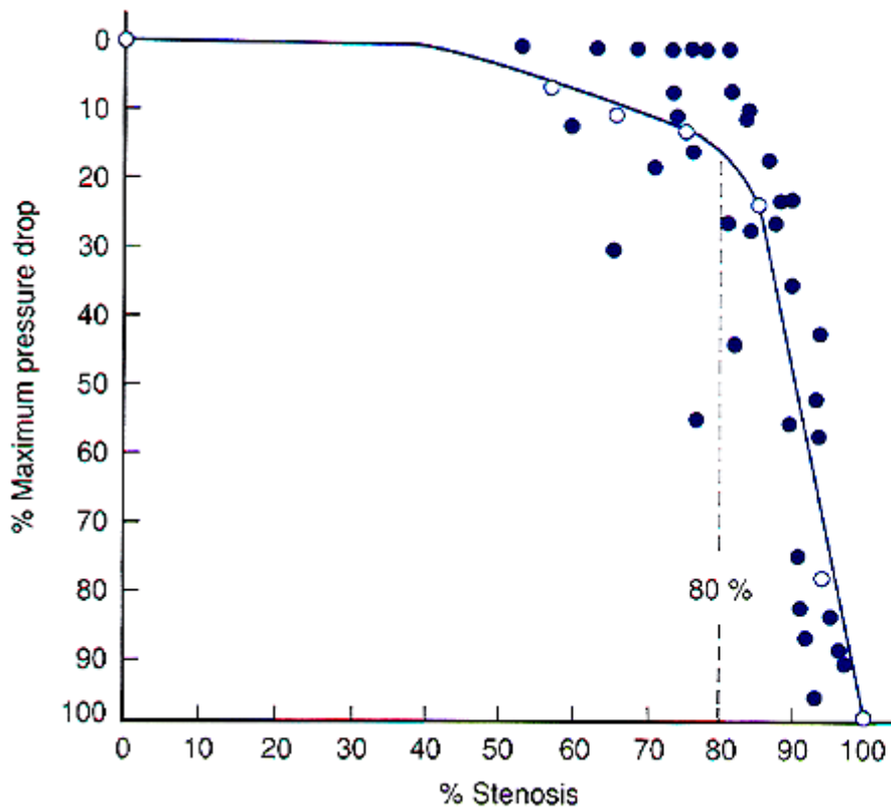
*Diagnostic tests*

Plasma renin activity normal or low  
Lateralized features: none

# Measured fall in arterial pressure and blood flow across stenotic vascular lesion induced in experimental animals



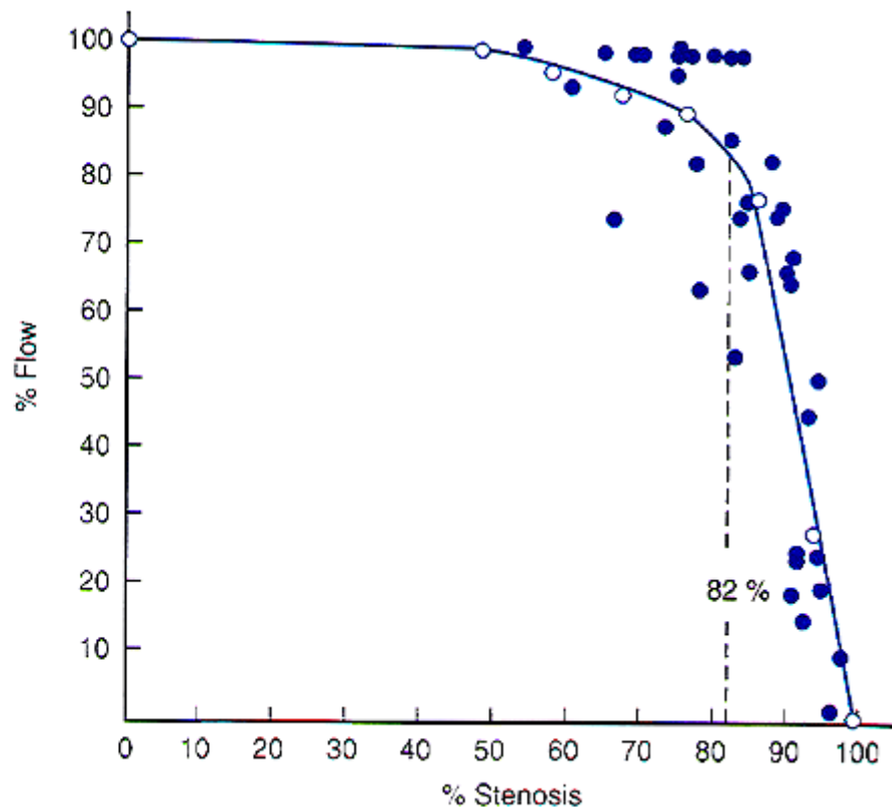
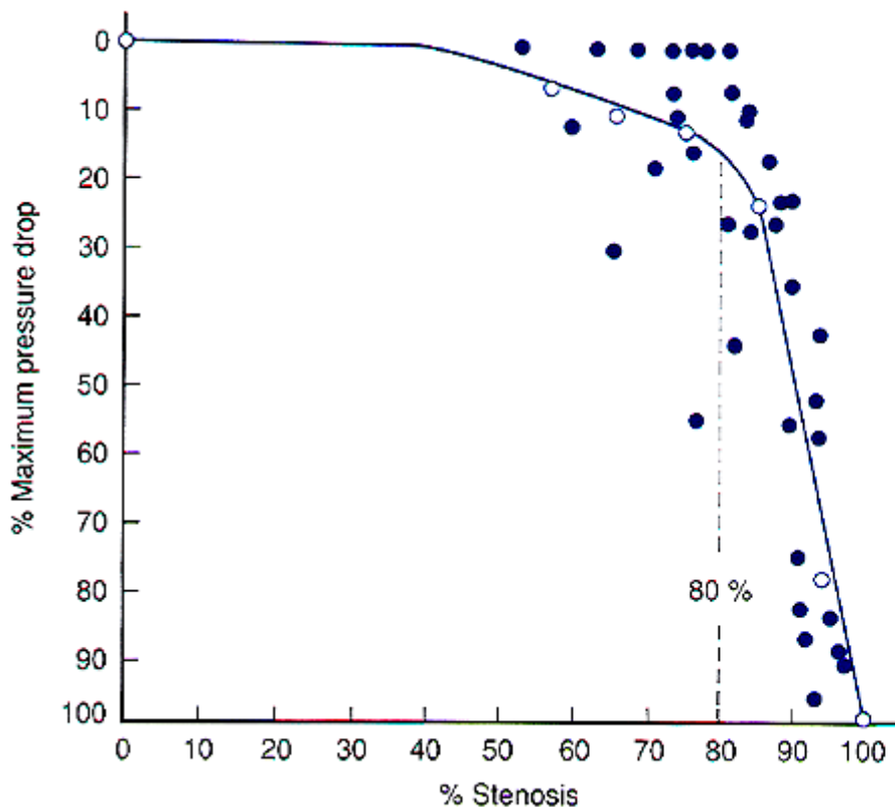
These data indicate that „critical” lesions require 70-80% luminal obstruction before hemodynamic effects can be detected



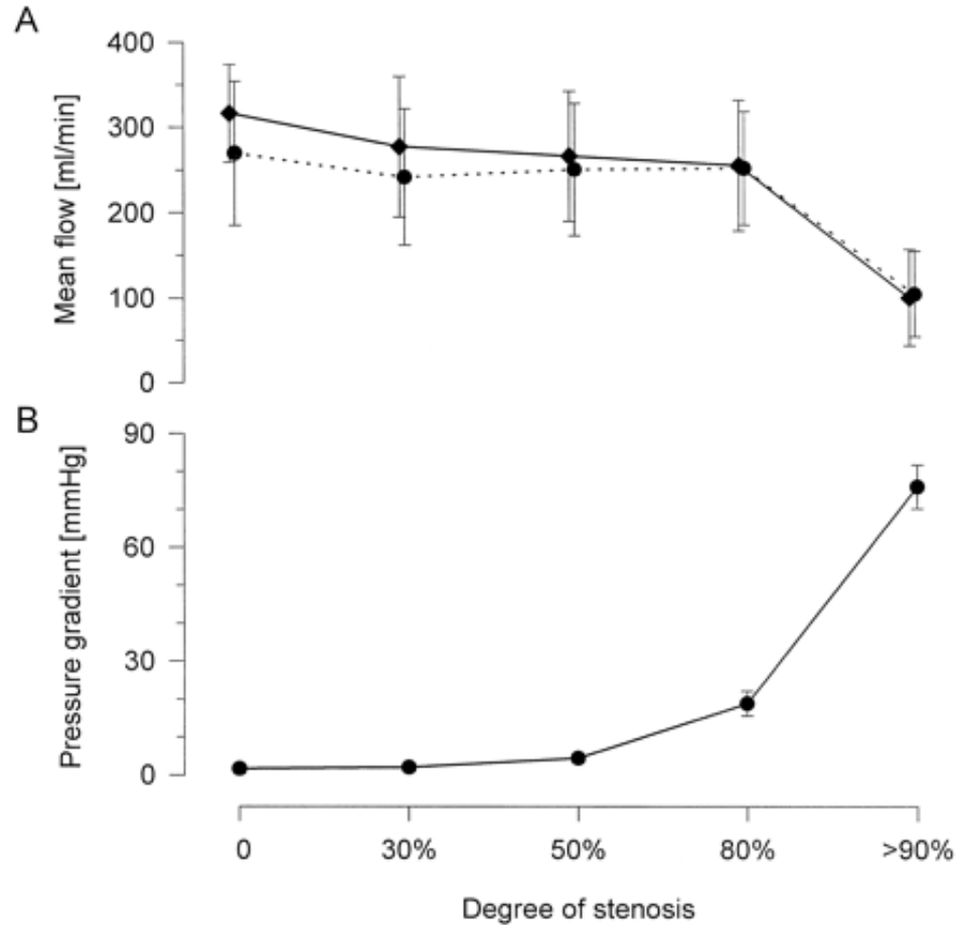
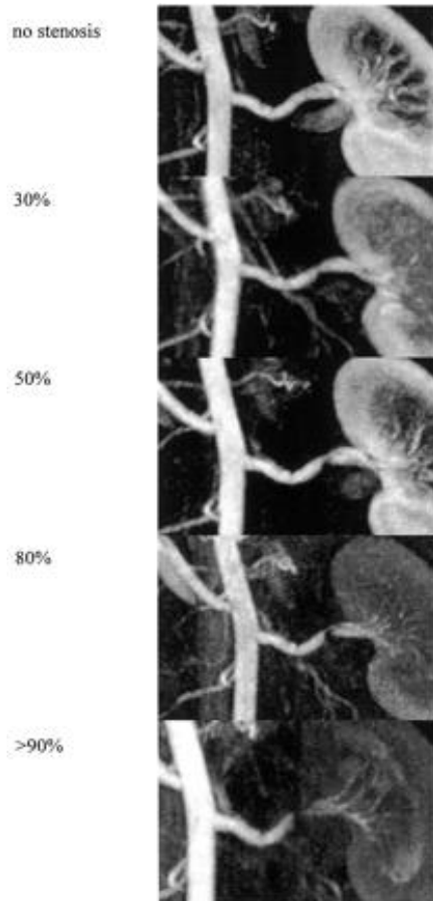


# Measured fall in arterial pressure and blood flow across stenotic vascular lesion induced in experimental animals

These data indicate that „critical” lesions require **70-80% luminal obstruction** before hemodynamic effects can be detected

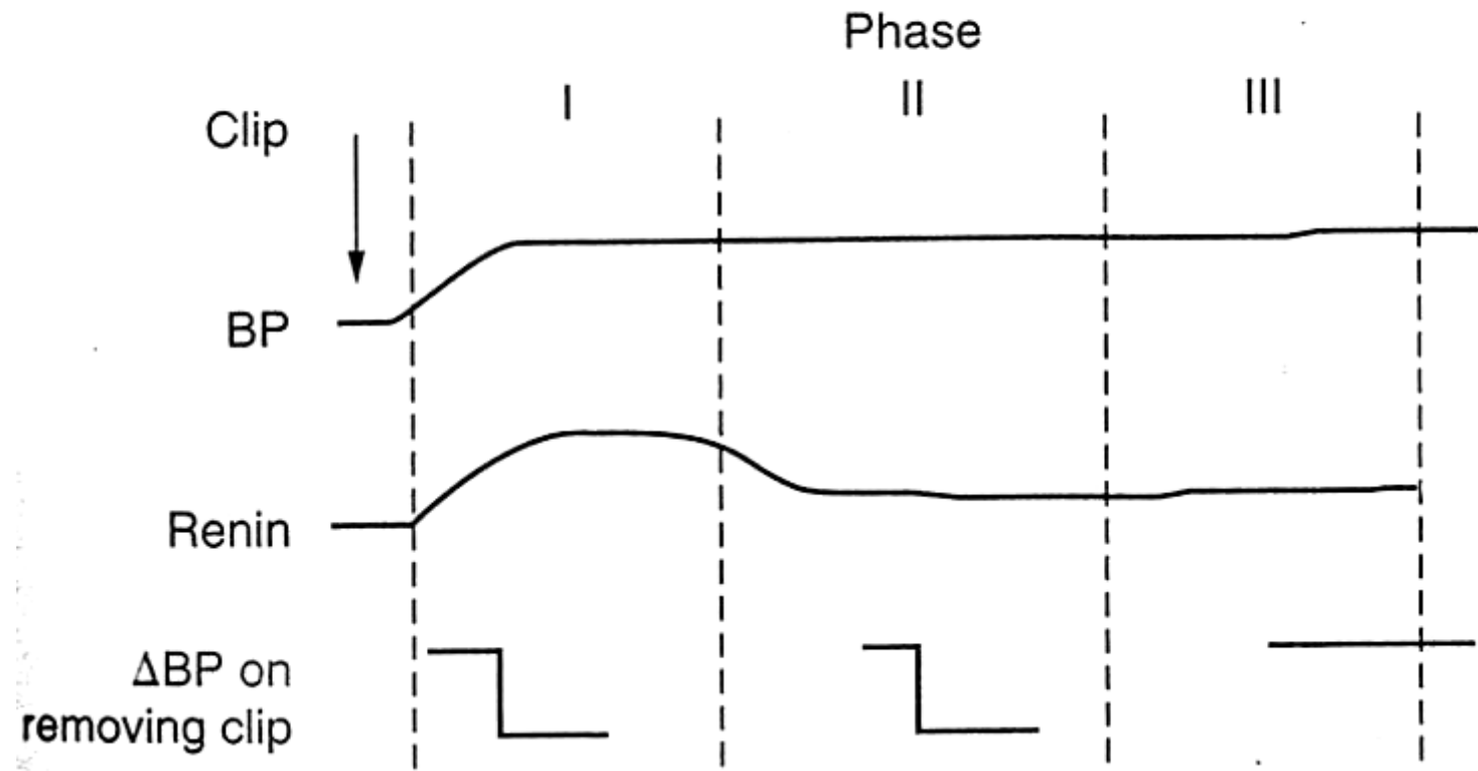


# Hemodynamic consequences of renal artery stenosis (MRI method)



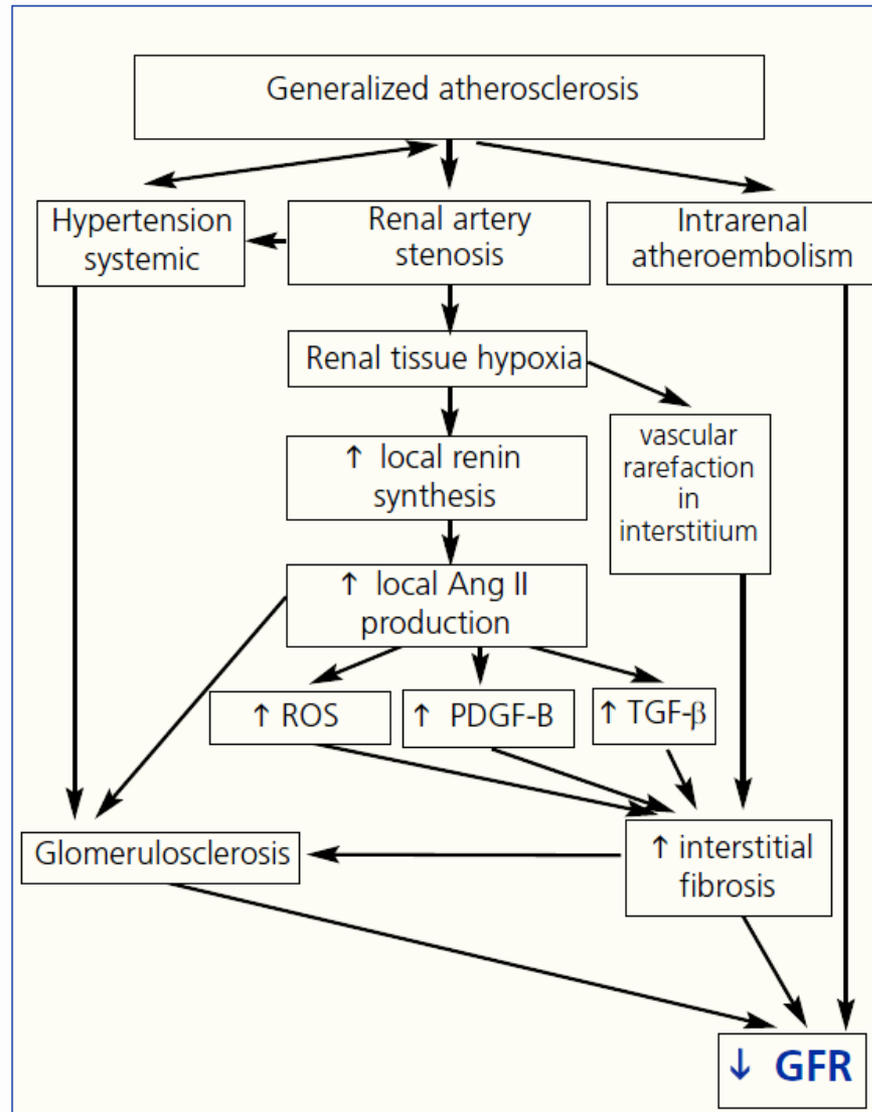


# Depiction of phases observed in experimental renovascular hypertension



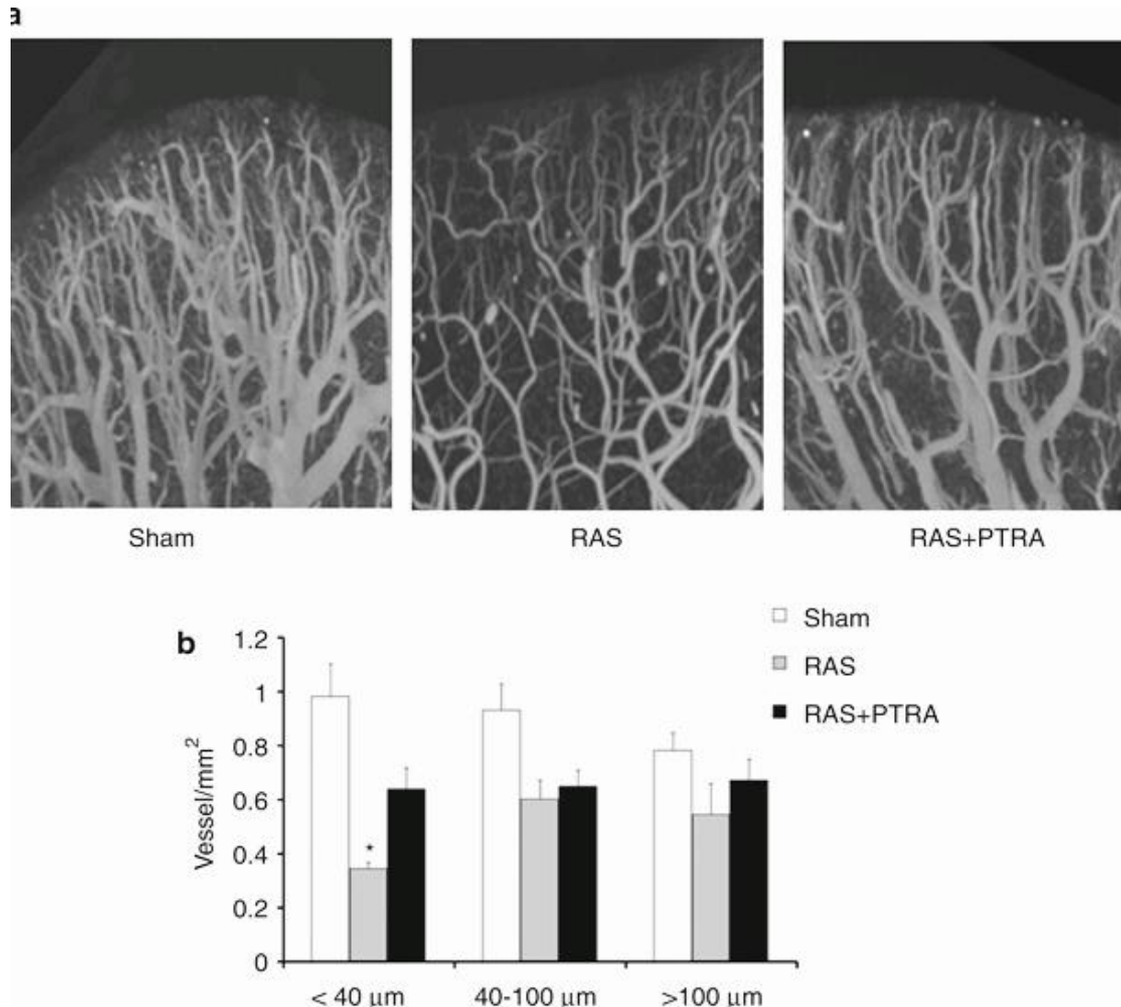


# Pathogenesis of ischemic nephropathy



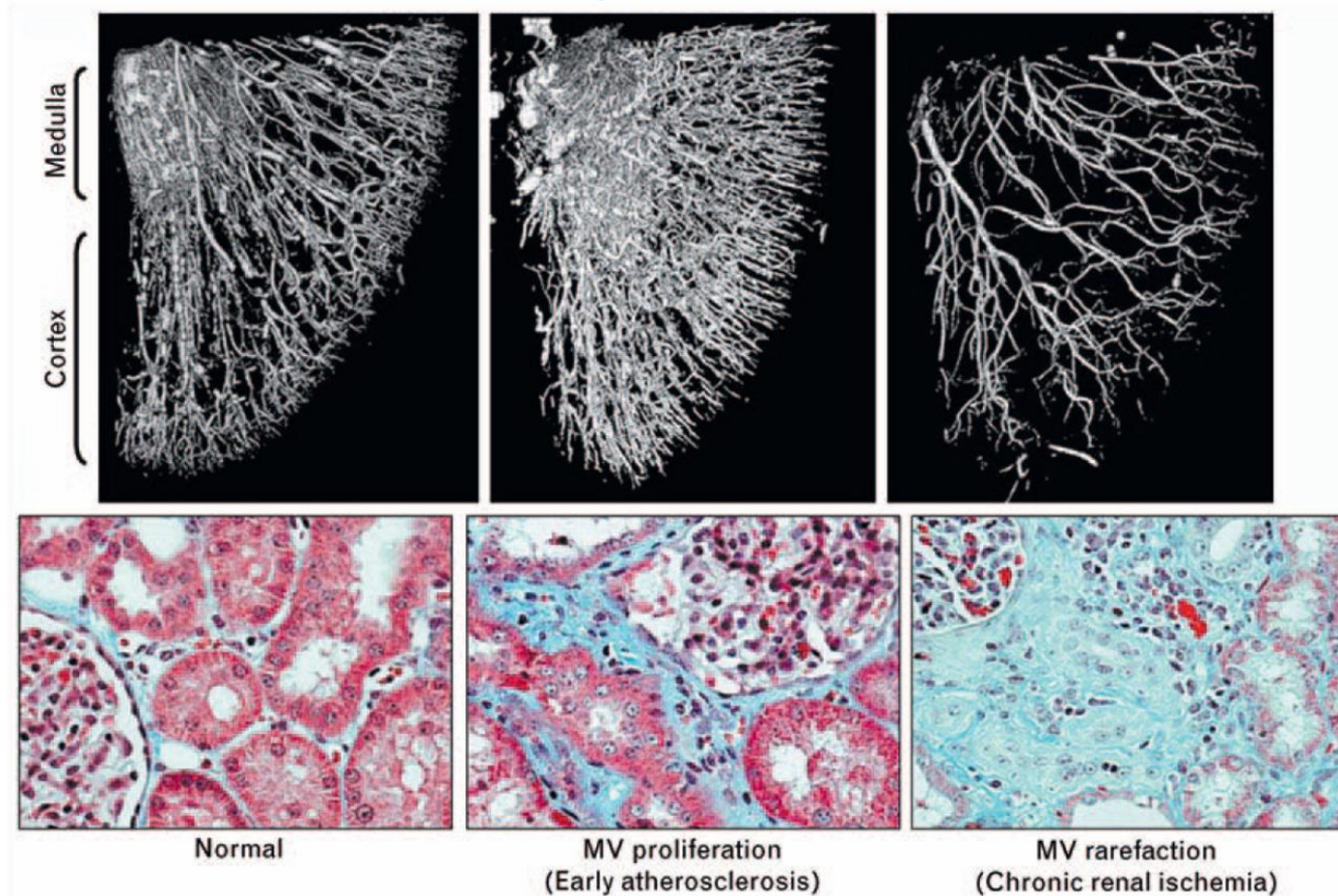


# Three-dimensional tomographic images of the **cortical microcirculation** in sham, RAS, and RAS + PTRA pigs





# Representative three-dimensional reconstruction of the renal microvascular architecture (using microcomputed tomography) and renal morphology (trichrome staining) showing opposing changes in microvascular architecture in early atherosclerosis compared with chronic ischemia

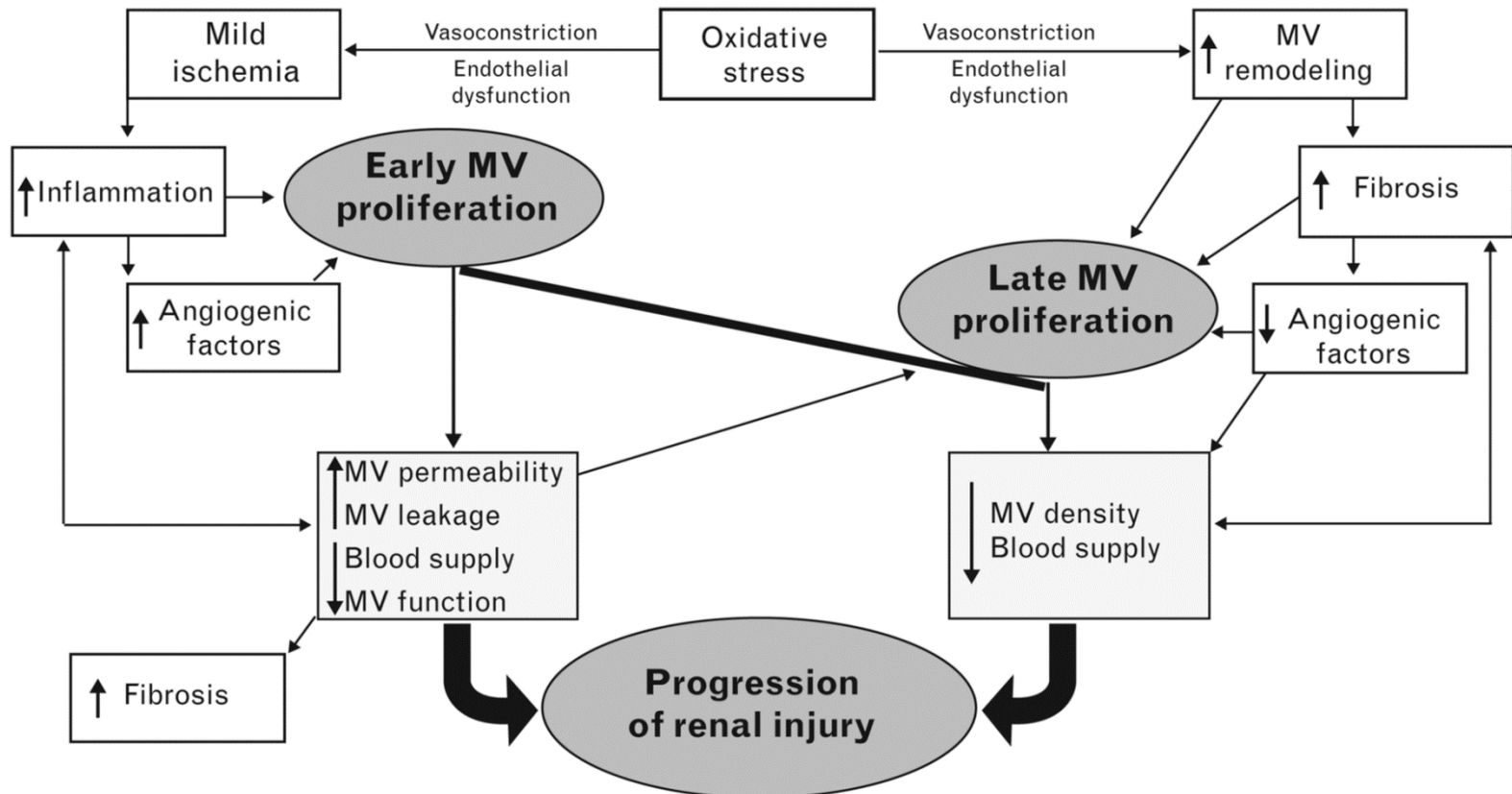


Microvascular rarefaction in particular is accompanied by severe renal fibrosis



At the early stage of renal injury, compensatory mechanisms induce MV proliferation, but abnormal MV structure and function, continuous oxidative stress, and accumulation of extracellular matrix (ECM) subsequently result in MV loss

## Chronic renal ischemia

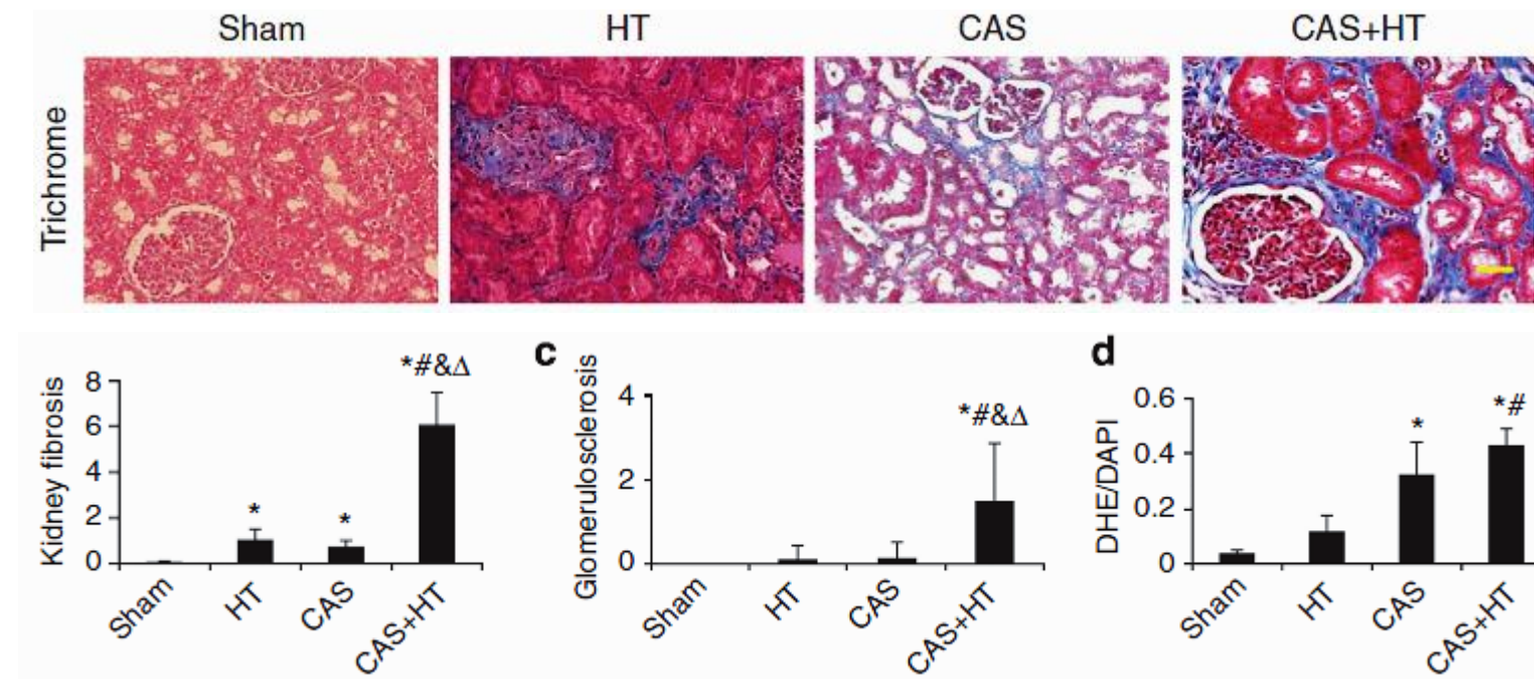




# Experimental coronary artery stenosis accelerates kidney damage in renovascular hypertensive swine

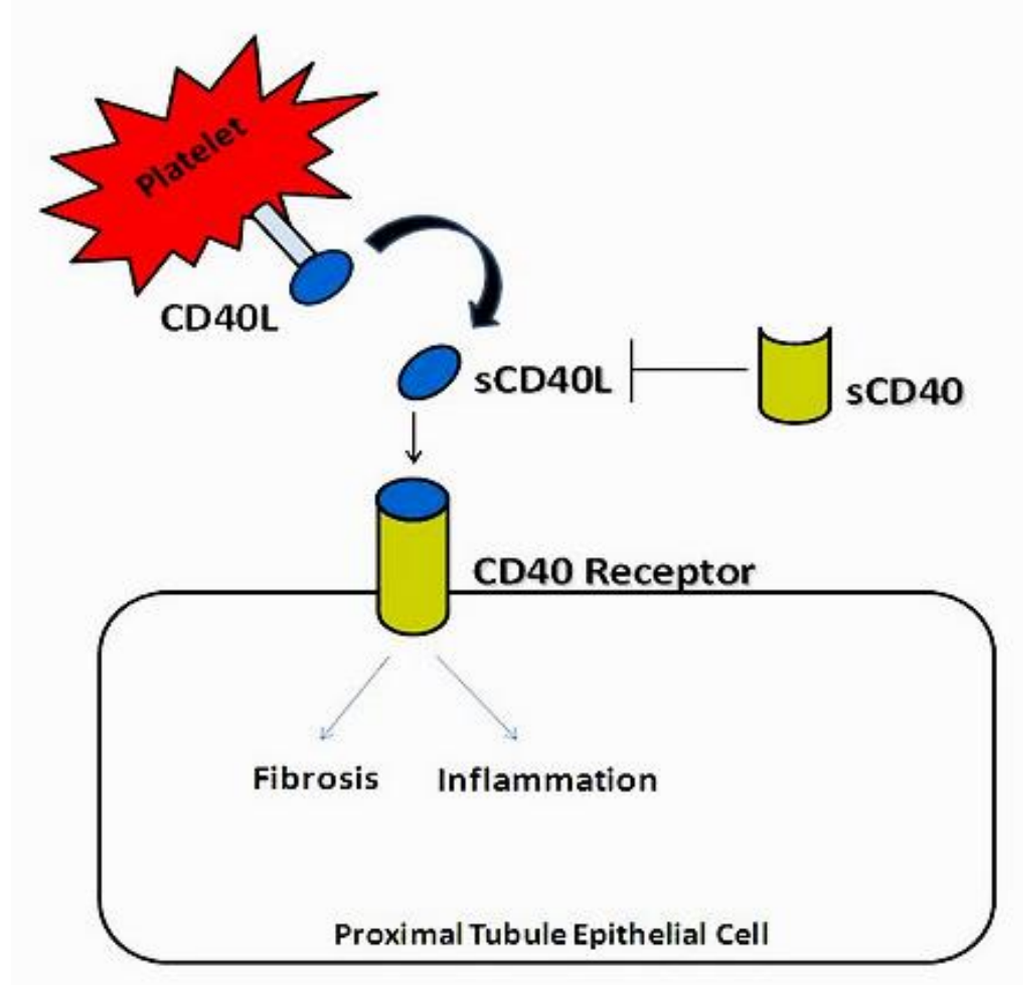
Dong Sun<sup>1,2</sup>, Alfonso Eirin<sup>1</sup>, Xiang-Yang Zhu<sup>1</sup>, Xin Zhang<sup>1</sup>, John A. Crane<sup>1</sup>, John R. Woollard<sup>1</sup>, Amir Lerman<sup>3</sup> and Lilach O. Lerman<sup>1,3</sup>

<sup>1</sup>Division of Nephrology and Hypertension, Mayo Clinic, Rochester, Minnesota, USA; <sup>2</sup>Department of Nephrology, The Affiliated Hospital of Xuzhou Medical College, Xuzhou, China and <sup>3</sup>Division Cardiovascular Disease, Mayo Clinic, Rochester, Minnesota, USA



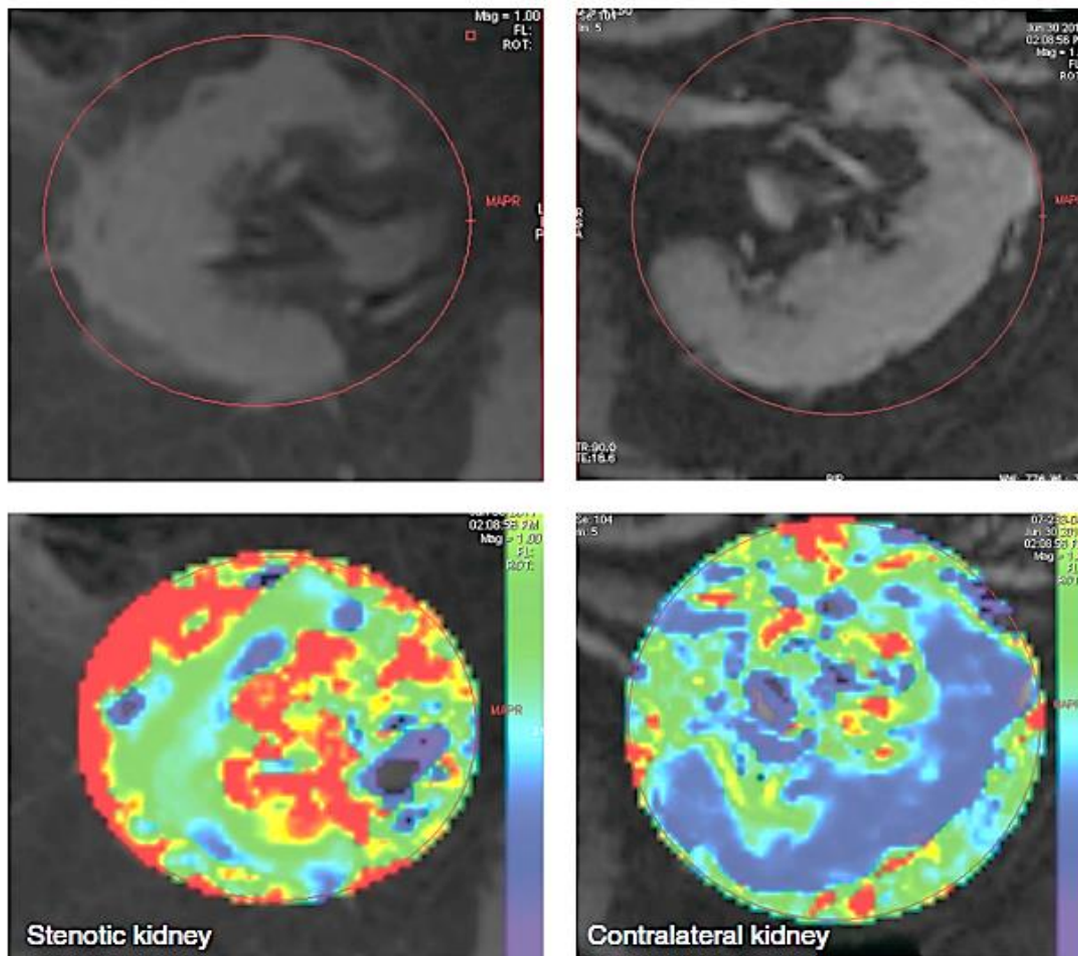


# Proposed mechanisms for **CD40/sCD40L** signaling in the proximal tubule contributing to the development of renal injury in atherosclerotic renal artery stenosis





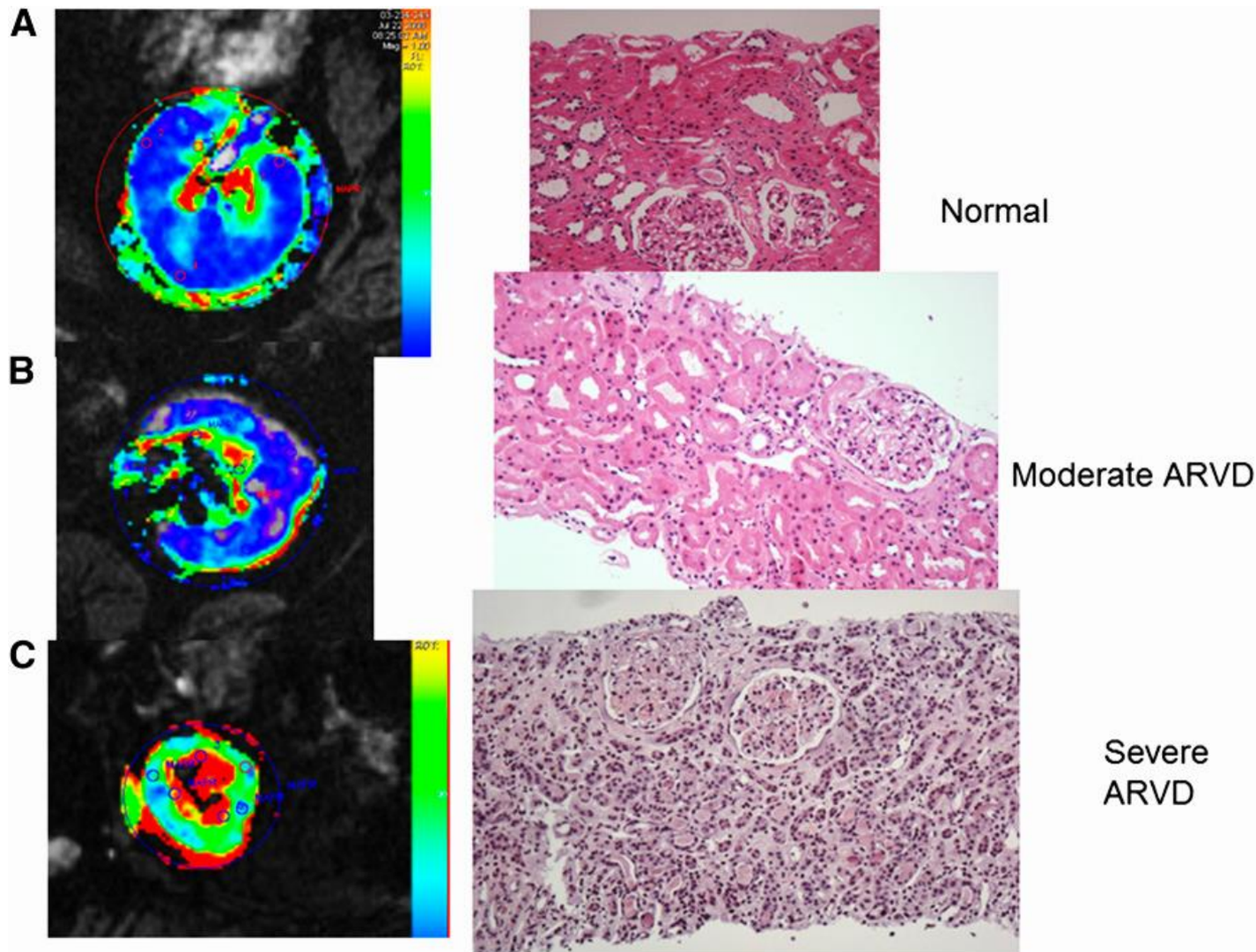
## Blood Oxygen Level-Dependent (BOLD) magnetic resonance imaging technique



**Figure 5 | Blood-oxygen-level-dependent (BOLD) magnetic resonance (MR) images with parametric maps depicting  $R2^*$  levels that correspond to tissue levels of deoxyhemoglobin in axial images of the kidneys. Both of these kidneys had high-grade renal arterial stenosis with velocities  $> 400$  cm/s. Serum creatinine was  $> 3.6$  mg/dl, although the patient was treated with angiotensin receptor blockers and diuretics. The larger kidney (right panel, left kidney) has well-preserved cortical oxygenation (blue zone) and a normal corticomedullary oxygen gradient. The smaller kidney (left panels) is developing overt cortical hypoxia with rising  $R2^*$  levels and expanding zone of medullary hypoxia (inner red zone). These functional imaging tools may assist in defining kidneys that are 'at risk' from critical vascular occlusion, yet remain 'salvageable' from the point of view of restoring renal blood flow (see text).**



# Cortical hypoxia and inflammation develop in severe Atherosclerotic Renovascular Disease (ARVD)



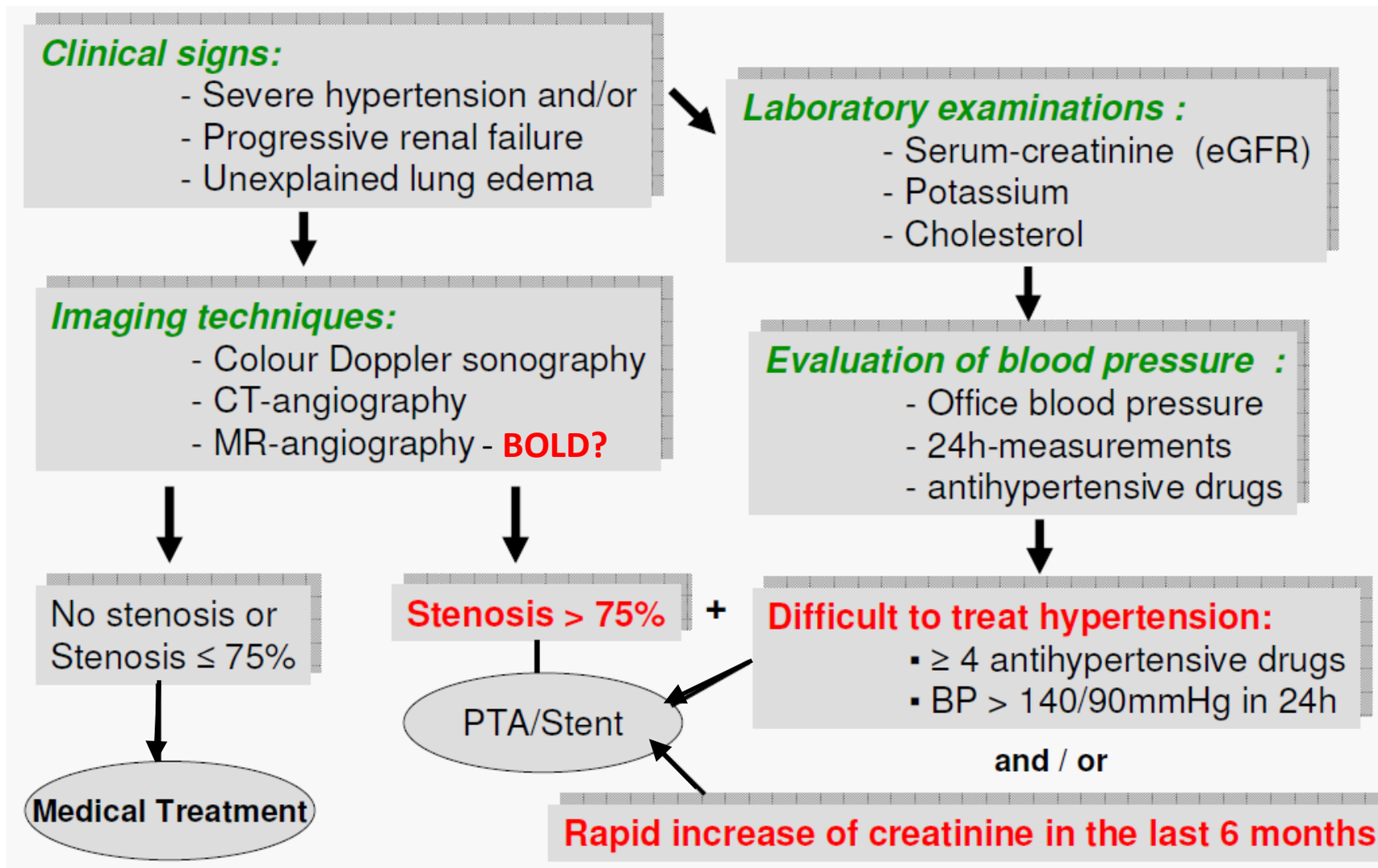


**Table 1. Prevalence of Atherosclerotic Renal Artery Stenosis in Different Subgroups**

| Subgroups                              | Prevalence of<br>Atherosclerotic Renal<br>Artery Stenosis<br>(>60% of renal artery lumen) |
|--|---|
| General population                     | 0.5%  |
| Age > 65 years (Doppler)               | 7%  |
| Healthy kidney donors                  | 3-5%  |
| Chronic kidney disease                 | 5.5%  |
| Suspicion of renovascular hypertension | 14%   |
| Coronary angiography                   | 19%-24% (7% bilateral)  |
| ESRD                                   | 12%-14% (2%-5% as cause of CKD)   |
| Peripheral arterial disease            | 28%-59%   |
| Abdominal aortic aneurysm              | 33%   |
| Eldery with CHF                        | 34%   |
| Refractory CHF                         | 40%-50%   |
| Diffuse arterial disease               | 50%   |

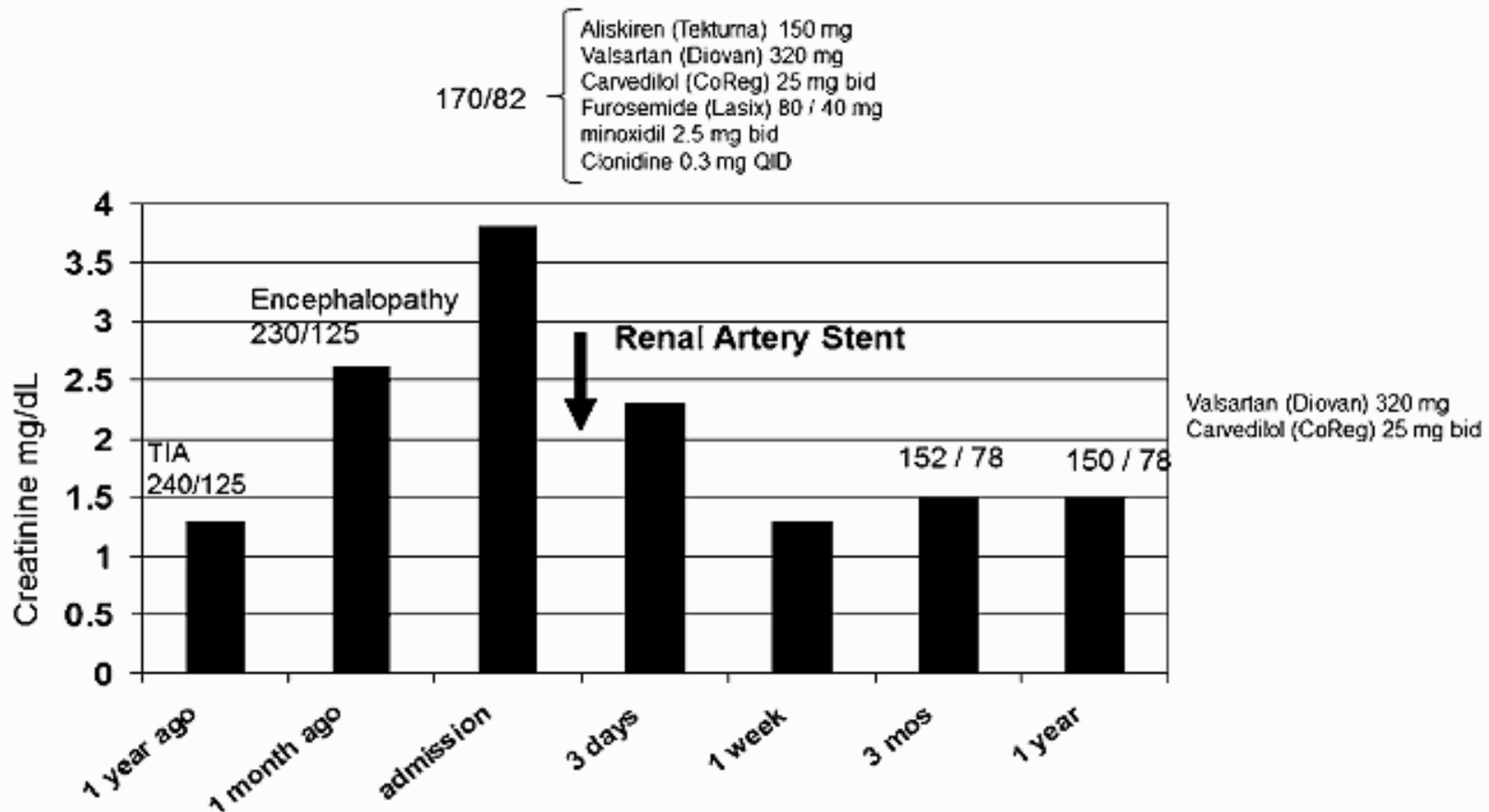


# Diagnostic work up of patients with suspected renal artery stenosis in 2016

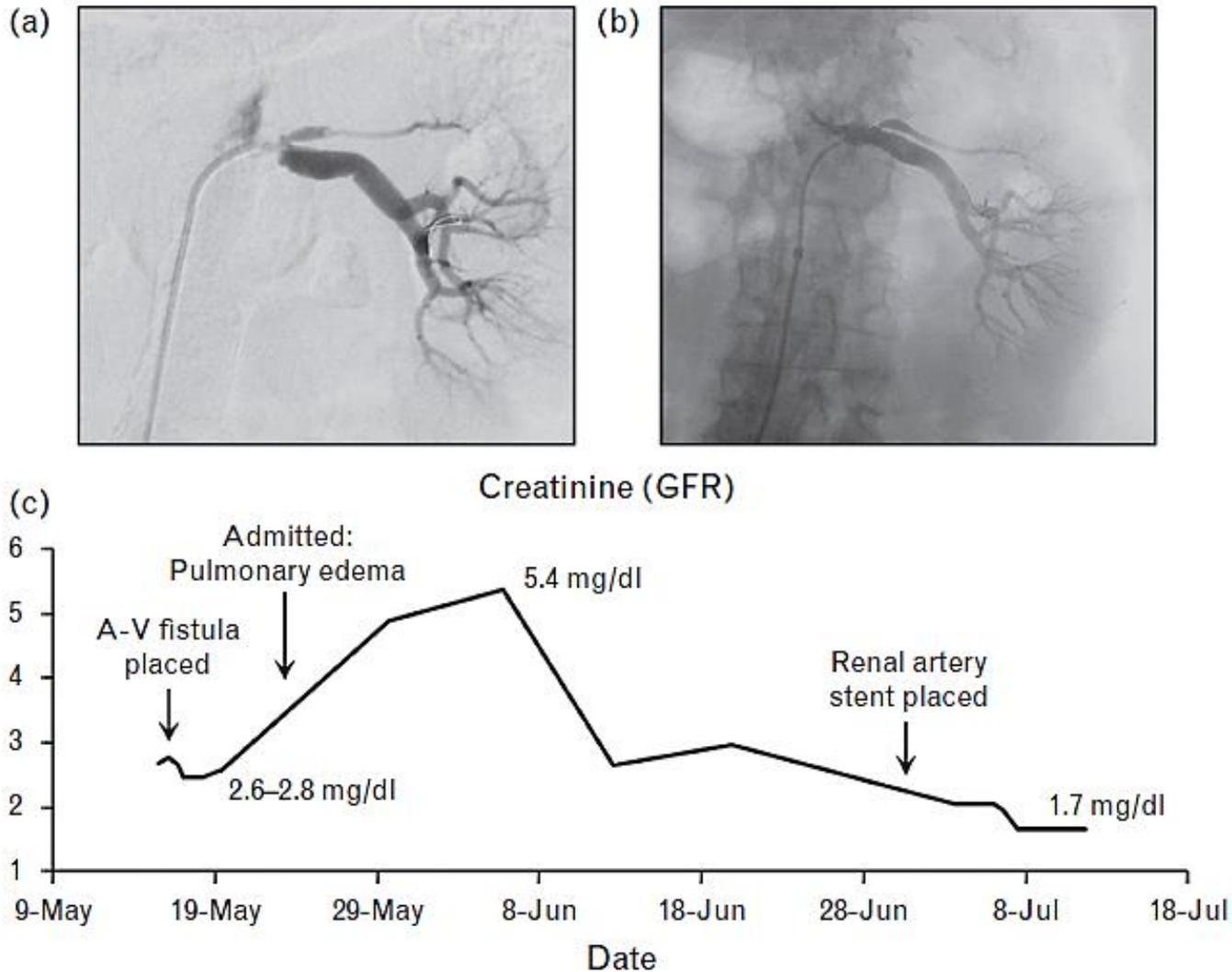


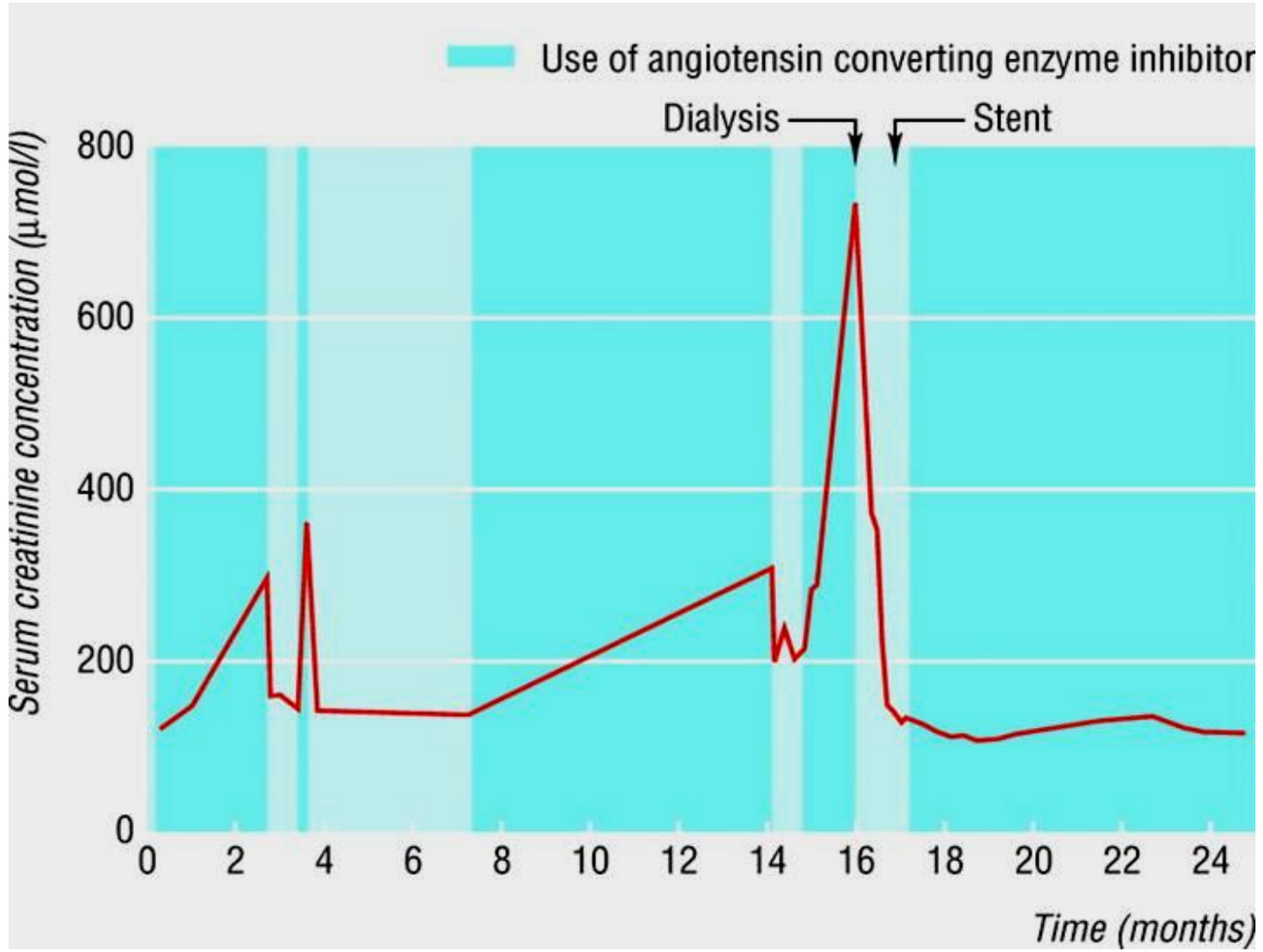


# Serum creatinine levels from 1 year before to 1 year after renal revascularization

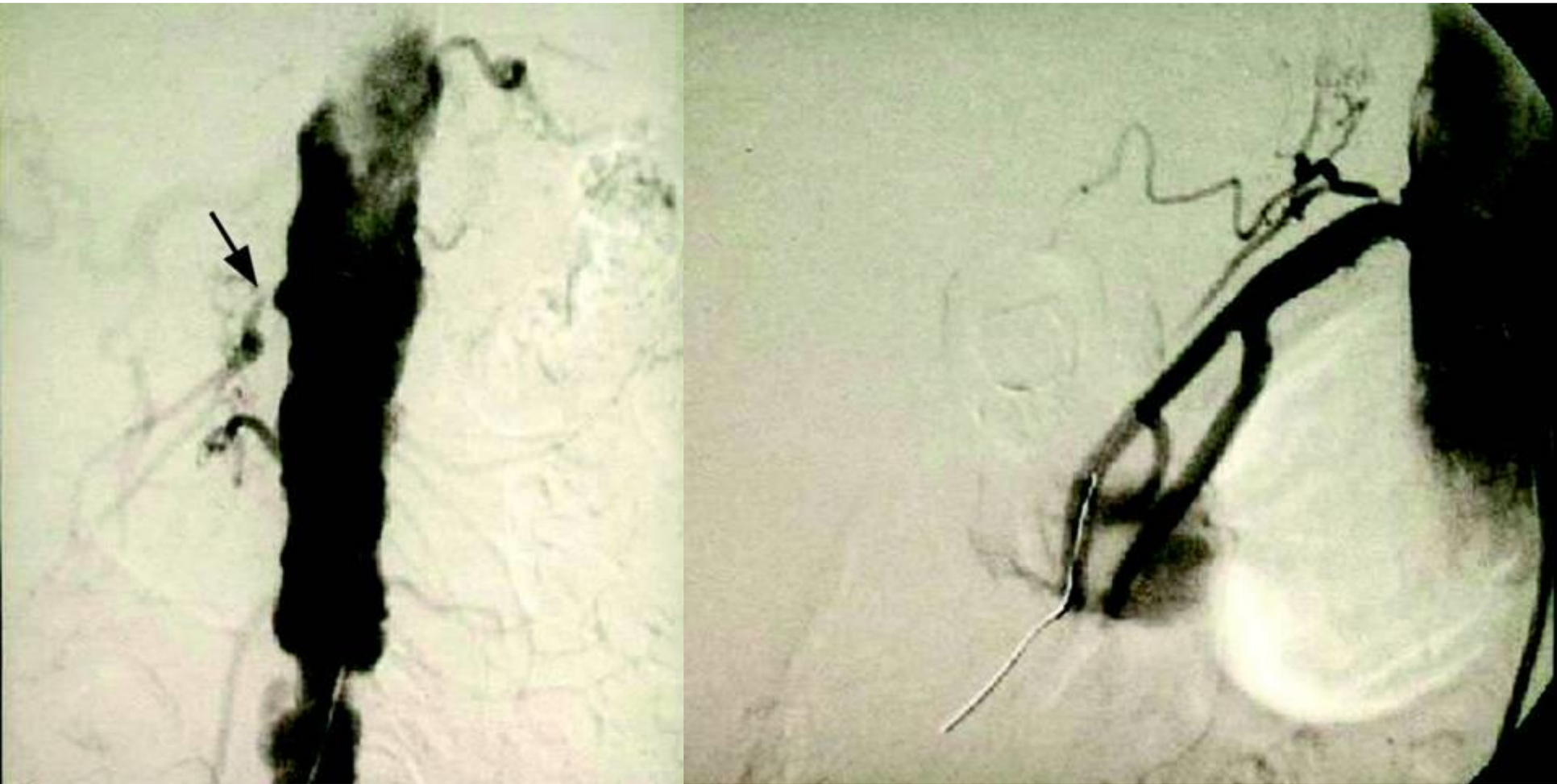


# Renal angiograms and serial serum creatinine values during a 6-week time period obtained for a 62-year-old diabetic patient

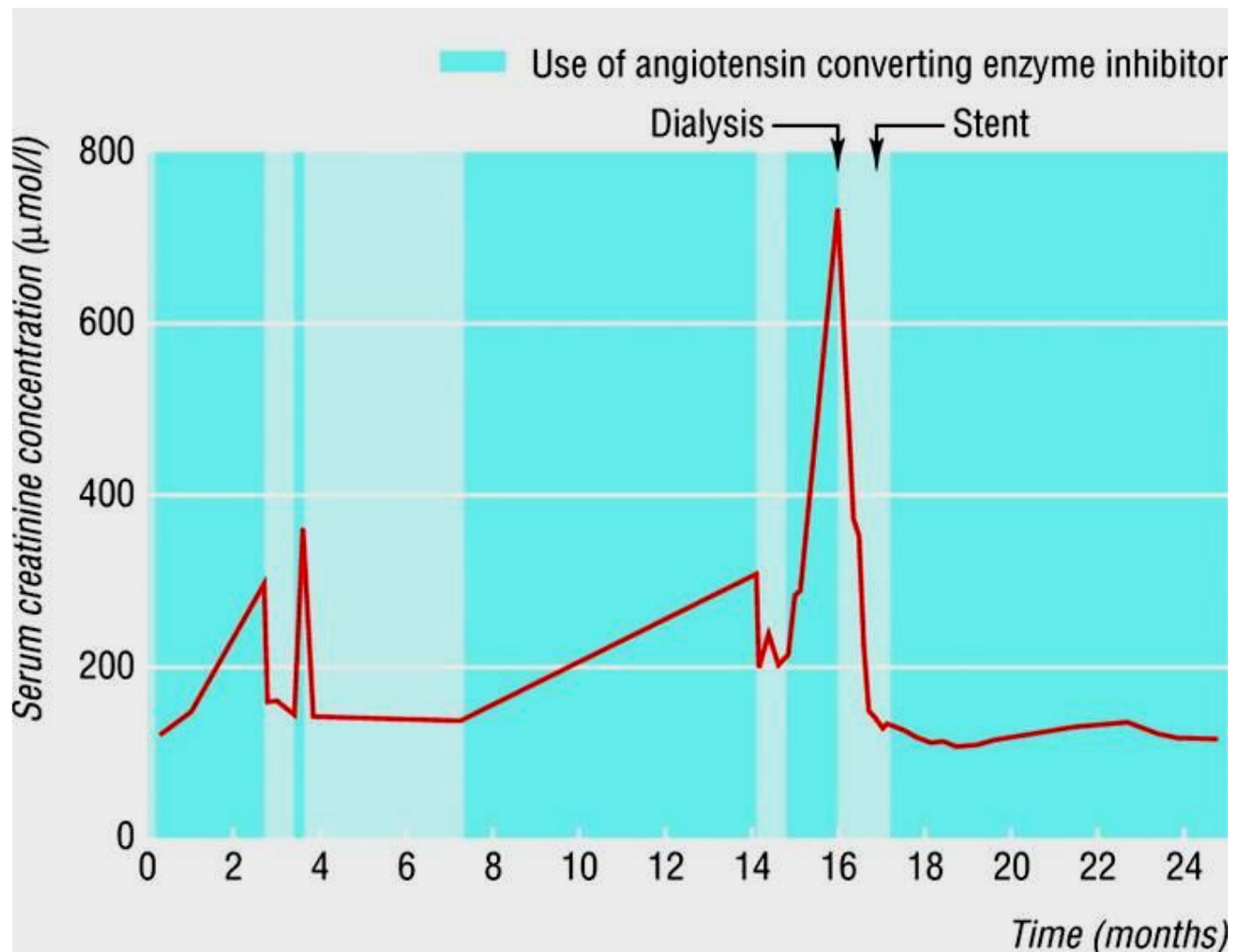




**Serum creatinine concentration increased on four occasions in association with angiotensin converting enzyme inhibition, leading to dialysis, then remained stable despite patient taking angiotensin converting enzyme inhibitor after dilation and stenting of right renal artery.**



**Renal arteriogram showing occlusion of left renal artery (*left*) and tight stenosis of right renal artery before (arrowed) and after stenting (*right*)**

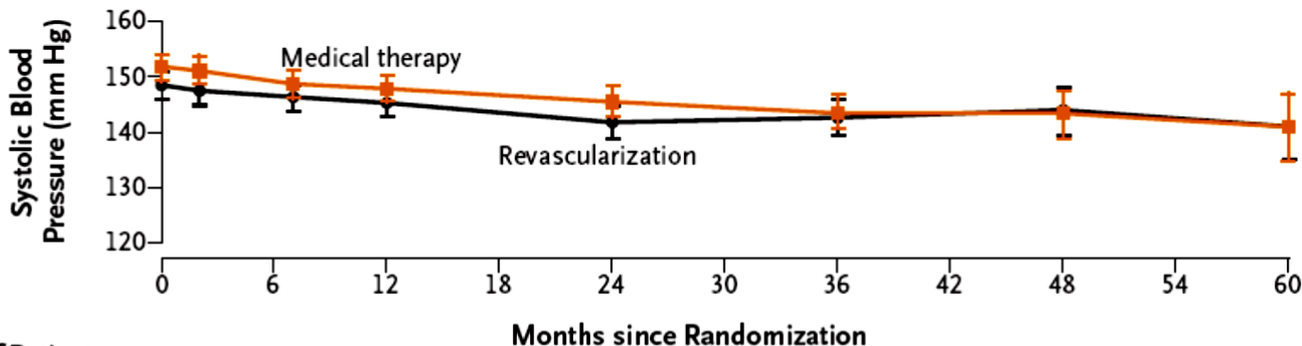


**Serum creatinine concentration increased on four occasions in association with angiotensin converting enzyme inhibition, leading to dialysis, then remained stable despite patient taking angiotensin converting enzyme inhibitor after dilation and stenting of right renal artery.**



# ASTRAL - Systolic and diastolic blood pressure in patients with renal artery stenosis treated with revascularization or medical therapy alone

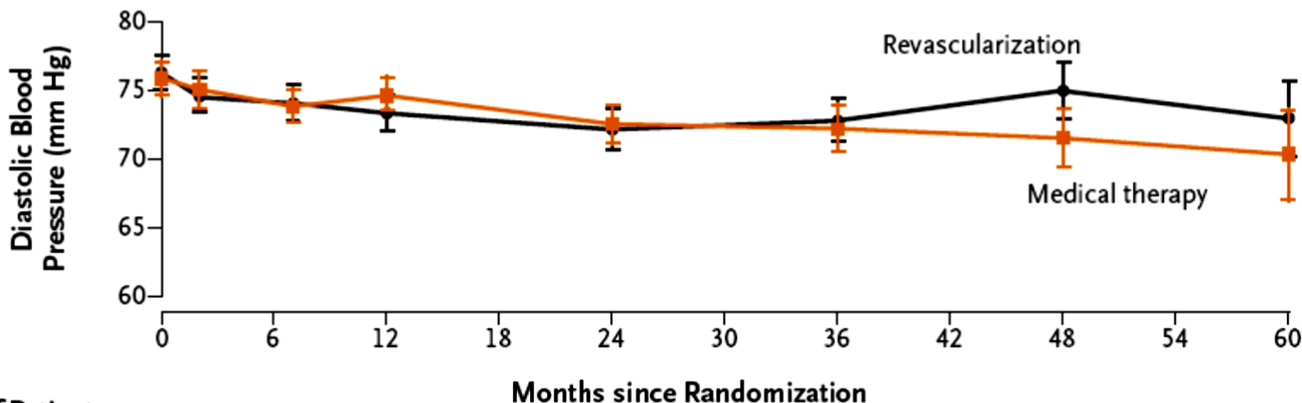
## A Systolic Blood Pressure



### Number of Patients

|                   | 0   | 6   | 12  | 24  | 36  | 48  | 60  |    |
|-------------------|-----|-----|-----|-----|-----|-----|-----|----|
| Revascularization | 385 | 346 | 332 | 321 | 257 | 197 | 125 | 71 |
| Medical therapy   | 388 | 361 | 350 | 336 | 264 | 178 | 124 | 62 |

## B Diastolic Blood Pressure

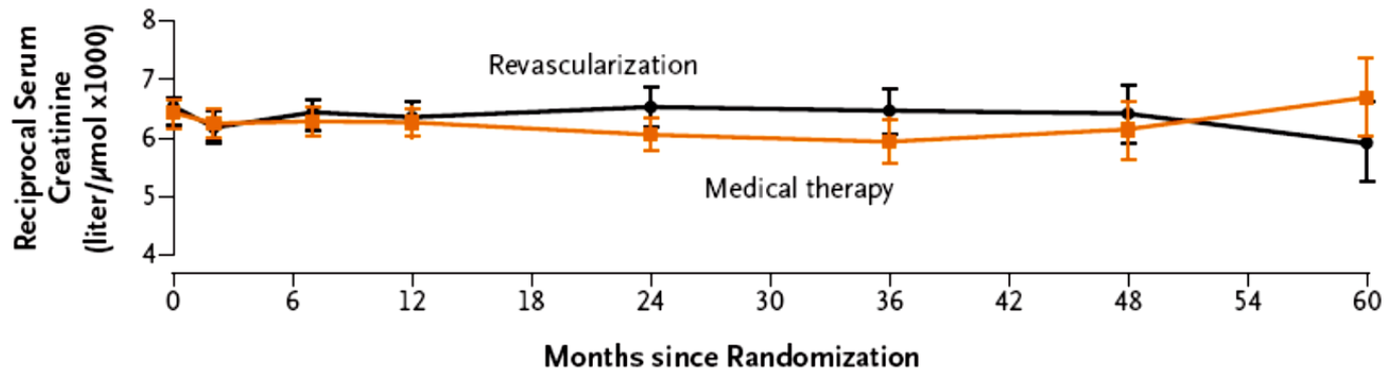


### Number of Patients

|                   | 0   | 6   | 12  | 24  | 36  | 48  | 60  |    |
|-------------------|-----|-----|-----|-----|-----|-----|-----|----|
| Revascularization | 384 | 344 | 330 | 320 | 256 | 197 | 125 | 70 |
| Medical therapy   | 388 | 361 | 349 | 335 | 262 | 178 | 123 | 63 |

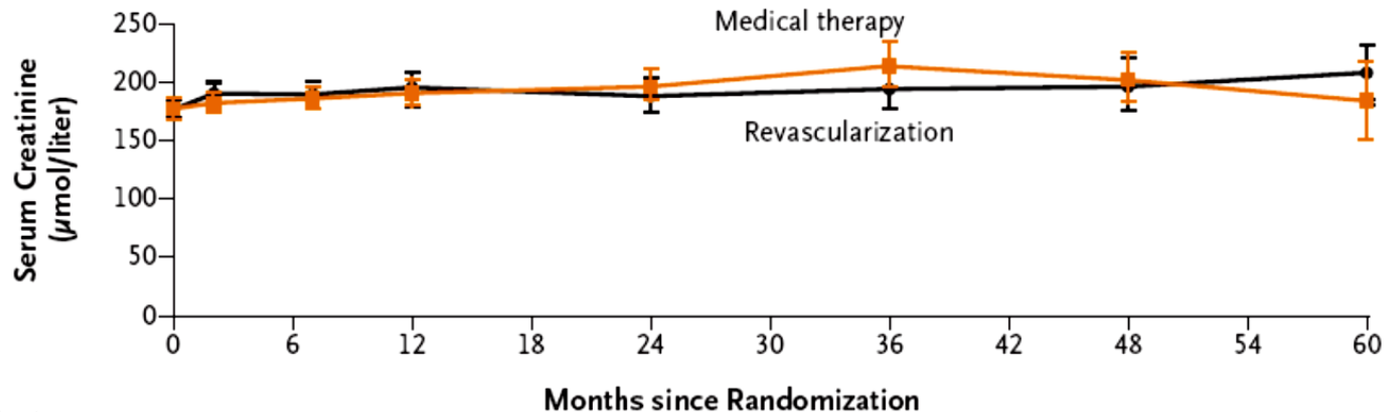
# ASTRAL - Renal function in patients with renal-artery stenosis treated with revascularization or medical therapy alone

## A Reciprocal of Serum Creatinine



| No. of Patients   | 0   |     | 6   |     | 12 |  | 24  |  | 36  |  | 48  |  | 60 |  |
|-------------------|-----|-----|-----|-----|----|--|-----|--|-----|--|-----|--|----|--|
| Revascularization | 403 | 349 | 336 | 329 |    |  | 263 |  | 191 |  | 127 |  | 72 |  |
| Medical therapy   | 403 | 363 | 347 | 343 |    |  | 272 |  | 183 |  | 119 |  | 61 |  |

## B Serum Creatinine

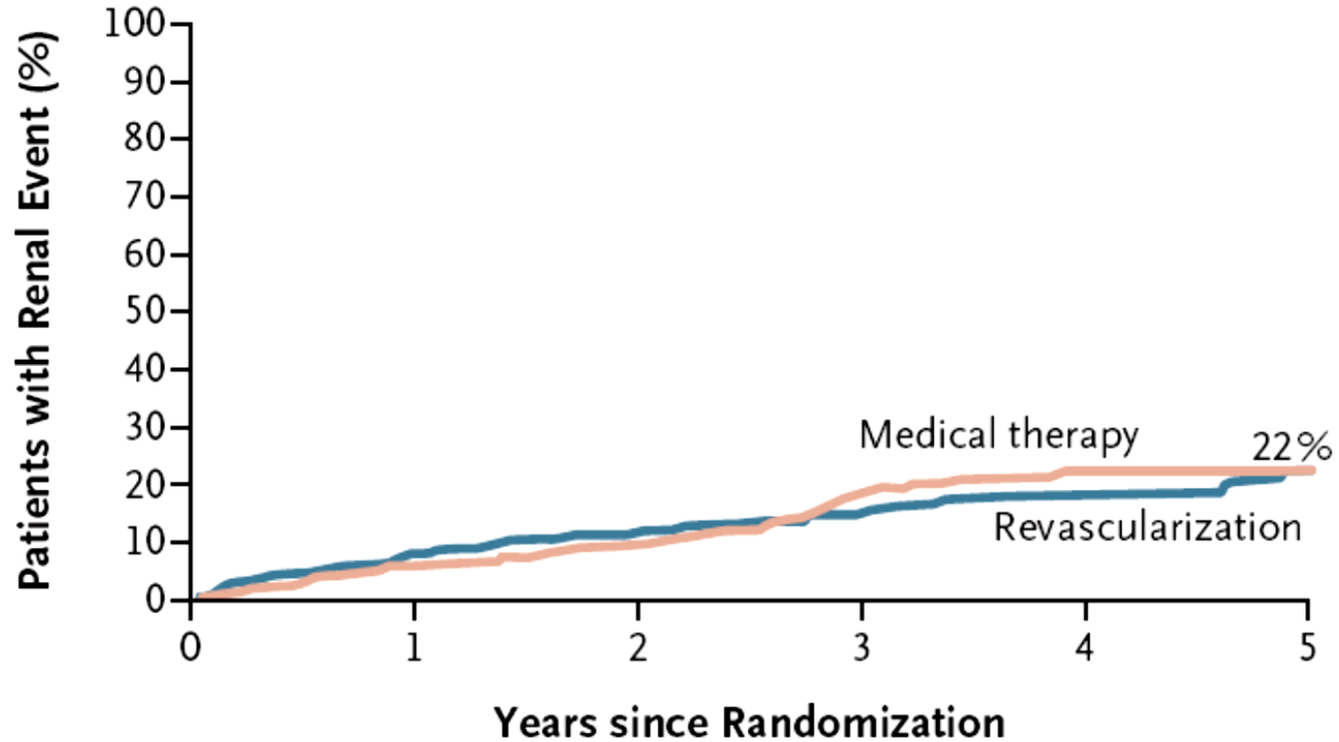


| No. of Patients   | 0   |     | 6   |     | 12 |  | 24  |  | 36  |  | 48  |  | 60 |  |
|-------------------|-----|-----|-----|-----|----|--|-----|--|-----|--|-----|--|----|--|
| Revascularization | 403 | 349 | 336 | 329 |    |  | 263 |  | 191 |  | 127 |  | 72 |  |
| Medical therapy   | 403 | 363 | 347 | 343 |    |  | 272 |  | 183 |  | 119 |  | 61 |  |



# ASTRAL - Kaplan–Meier curves for the time to the first renal events

## A First Renal Event

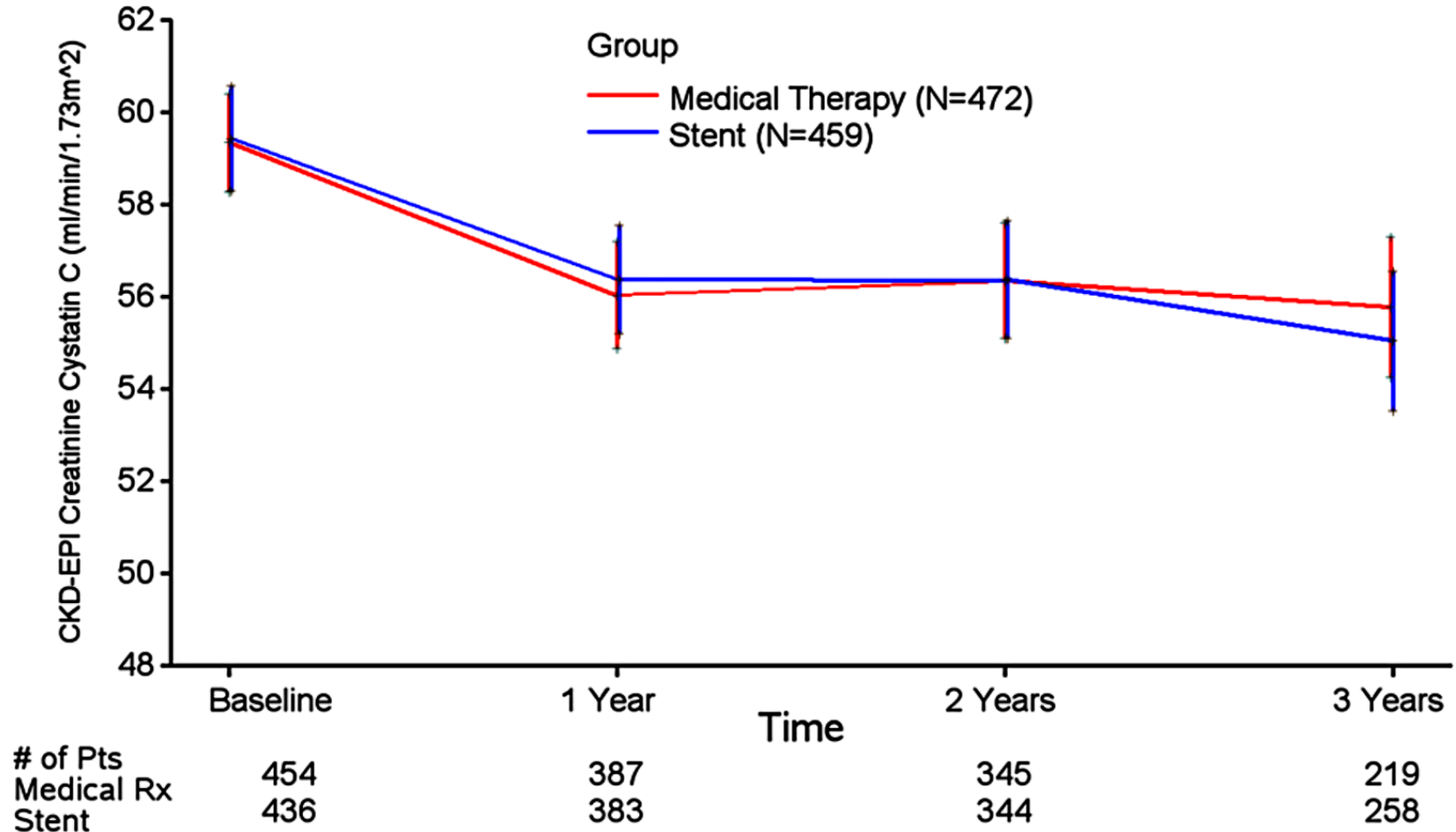


### No. at Risk

|                   |     |     |     |     |    |    |
|-------------------|-----|-----|-----|-----|----|----|
| Revascularization | 403 | 315 | 236 | 157 | 99 | 39 |
| Medical therapy   | 403 | 319 | 233 | 145 | 84 | 37 |



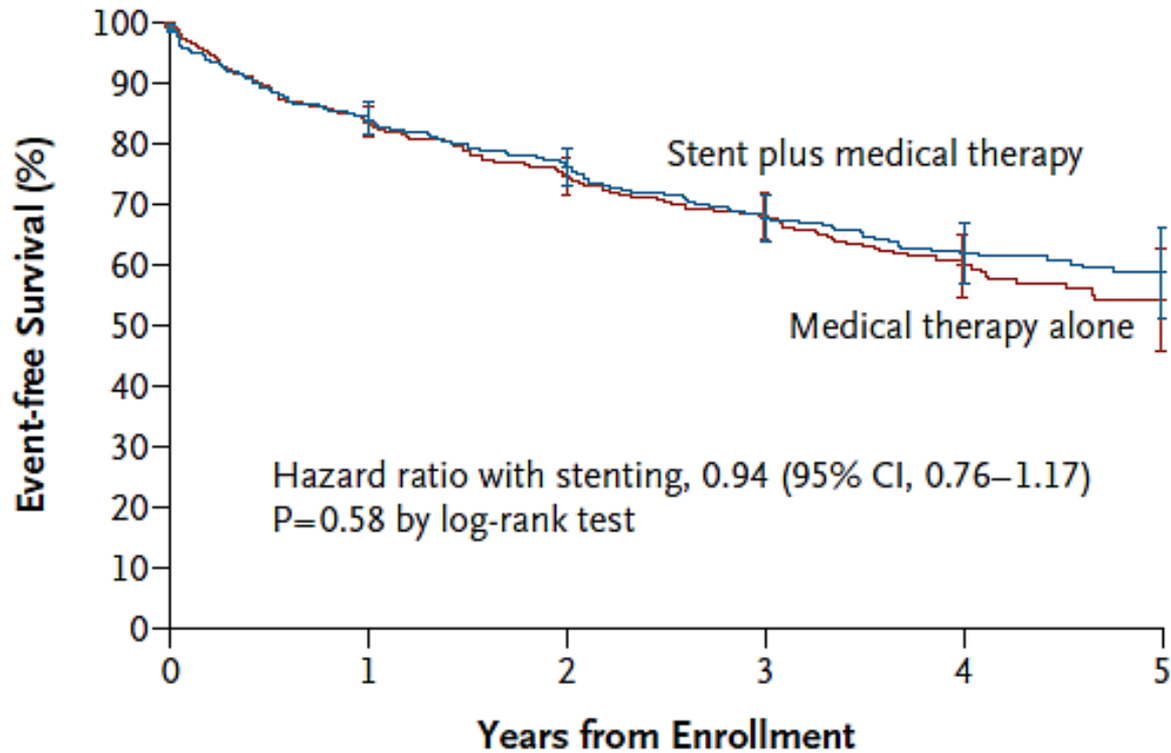
# CKD-EPI eGFR (ml/min./1.73m<sup>2</sup>) in randomized treatment groups of the CORAL Trial





# The CORAL Study

## Kaplan–Meier Curves for the Primary Outcome



### No. at Risk

|                            |     |     |     |     |     |    |
|----------------------------|-----|-----|-----|-----|-----|----|
| Medical therapy alone      | 472 | 371 | 314 | 214 | 115 | 40 |
| Stent plus medical therapy | 459 | 362 | 318 | 224 | 131 | 59 |





# The NEW ENGLAND JOURNAL of MEDICINE

ESTABLISHED IN 1812

JANUARY 2, 2014

VOL. 370 NO. 1

## Stenting and Medical Therapy for Atherosclerotic Renal-Artery Stenosis

Christopher J. Cooper, M.D., Timothy P. Murphy, M.D., Donald E. Cutlip, M.D., Kenneth Jamerson, M.D., William Henrich, M.D., Diane M. Reid, M.D., David J. Cohen, M.D., Alan H. Matsumoto, M.D., Michael Steffes, M.D., Michael R. Jaff, D.O., Martin R. Prince, M.D., Ph.D., Eldrin F. Lewis, M.D., Katherine R. Tuttle, M.D., Joseph I. Shapiro, M.D., M.P.H., John H. Rundback, M.D., Joseph M. Massaro, Ph.D., Ralph B. D'Agostino, Sr., Ph.D., and Lance D. Dworkin, M.D., for the CORAL Investigators\*

### ABSTRACT

#### BACKGROUND

Atherosclerotic renal-artery stenosis is a common problem in the elderly. Despite two randomized trials that did not show a benefit of renal-artery stenting with respect to kidney function, the usefulness of stenting for the prevention of major adverse renal and cardiovascular events is uncertain.

From the University of Toledo, Toledo, OH (C.J.C.); Rhode Island Hospital (T.P.M., L.D.D.) and Alpert Medical School of Brown University (T.P.M., L.D.D.) — both in Providence; Harvard Clinical Research Institute (D.E.C., J.M.M., R.B.D.), Beth Israel Deaconess Medical Center (D.E.C.),

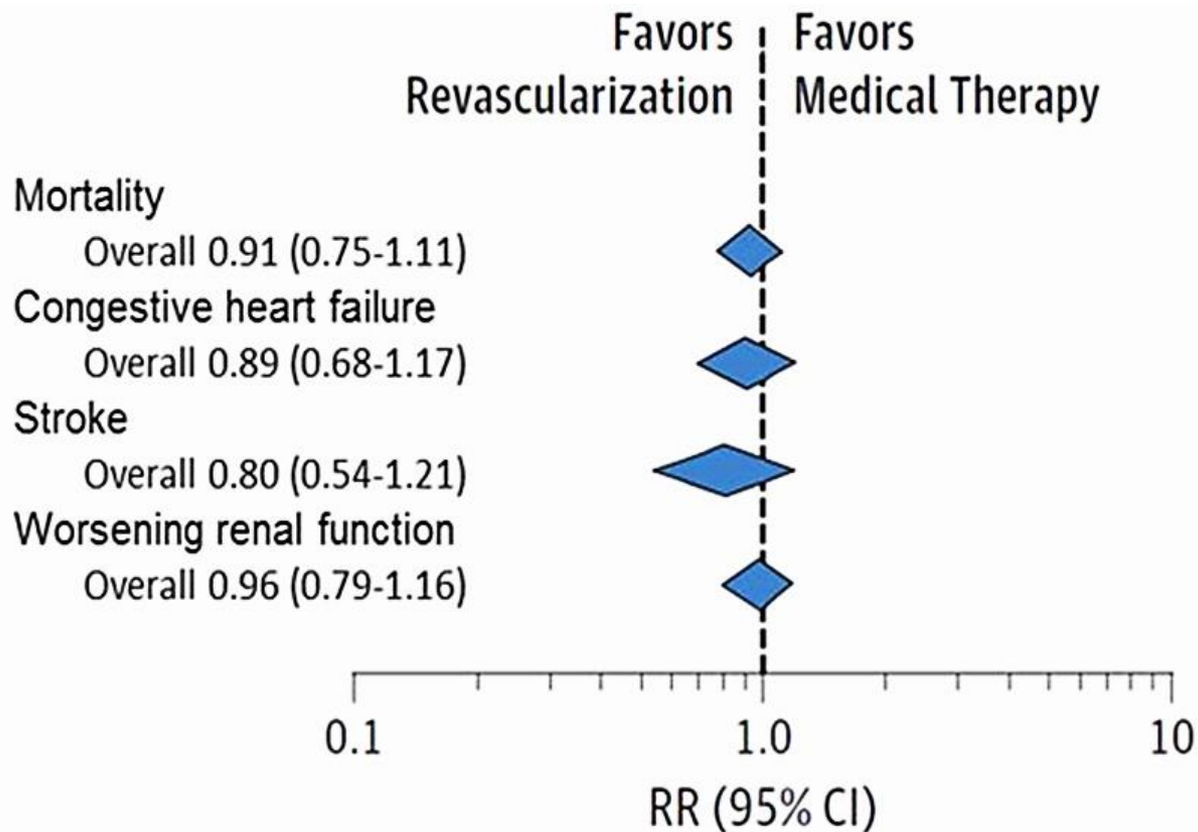
#### CONCLUSIONS

Renal-artery stenting did not confer a significant benefit with respect to the prevention of clinical events when added to comprehensive, multifactorial medical therapy in people with atherosclerotic renal-artery stenosis and hypertension or chronic kidney disease. (Funded by the National Heart, Lung and Blood Institute and others; ClinicalTrials.gov number, NCT00081731.)



# Renal artery revascularization:

updated meta-analysis with the CORAL trial summary estimates of cardiovascular outcomes for revascularization vs medical therapy



Included trials: STAR; ASTRAL; SNARSCG; NITER; CORAL; RASCAD; DRASTIC; EMMA



# Where now in the management of renal artery stenosis? Implications of the ASTRAL and CORAL trials

---

*James Ritchie, Helen V. Alderson, and Philip A. Kalra*

---

## **Purpose of review**

The neutral findings of Angioplasty and Stenting for Renal Artery Lesions and Cardiovascular Outcomes in Renal Artery Lesions trials have shown that unselected revascularization does not improve outcomes in atherosclerotic renovascular disease (ARVD). This review highlights recent translational, clinical and epidemiological studies and suggests directions for future research.

## **Recent findings**

Imaging studies show that the degree of renal artery stenosis is not the most important determinant of outcome and response to therapies in ARVD. Porcine models have established a better understanding of the microvascular and inflammatory changes that occur in ARVD. Biomarkers of inflammation and cardiovascular dysfunction may be informative but do not yet help assess prognosis or response to treatment. Stem cell therapies show promise in animal models but have yet to translate into clinical practice. Analysis of patient subgroups with high-risk presentations of ARVD has provided new insights into treatment response and may guide future studies.

## **Summary**

It is time to reframe thinking and research in ARVD. We need better ways to identify patients likely to benefit from revascularization and to improve response to treatment in these individuals. Many preclinical studies show promise, but these are often small scale and difficult to replicate. Future work should focus on establishing an international disease registry as a foundation for collaborative research.



# Predictor factors for renal outcome in renal artery stenosis

R. CIANCI, P. MARTINA, A. GIGANTE, D. DI DONATO, L. POLIDORI, P. PRESTA\*\*, R. LABBADIA, D. AMOROSO, A. ZACCARIA\*, B. BARBANO, G. FUIANO\*\*

Department of Nephrology, School of Medicine, Sapienza University, Rome, Italy

\*Department of Vascular and Endovascular Surgery, San Pietro-Fatebenefratelli Hospital, Rome, Italy

\*\*Department of Nephrology, School of Medicine, Magna Graecia University of Catanzaro, Catanzaro, Italy

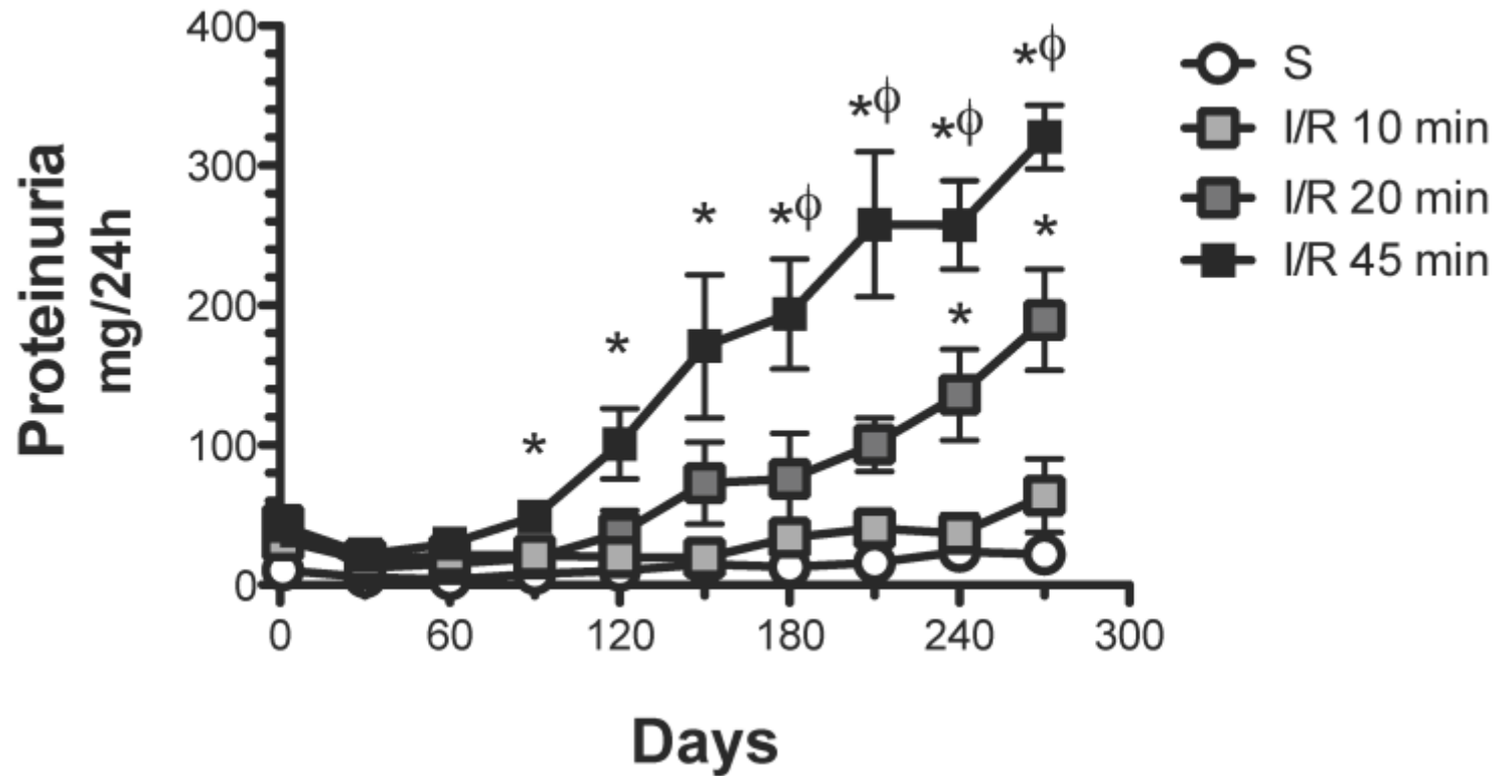
**MATERIALS AND METHODS:** we performed an observational study on a total of 55 patients to find predictive factors of the outcome of renal function after renal percutaneous transluminal angioplasty and stenting (RPTAs).

**RESULTS:** We found that uricemia, proteinuria and IR were higher at baseline in patients who worsened renal function after revascularization.

**CONCLUSIONS:** The identification of predictive factors (uricemia, proteinuria and RI) of chronic kidney disease (CKD) progression in patients with RAS undergone revascularization could be useful to predict renal long term outcome and to select patients that really could benefit of this.

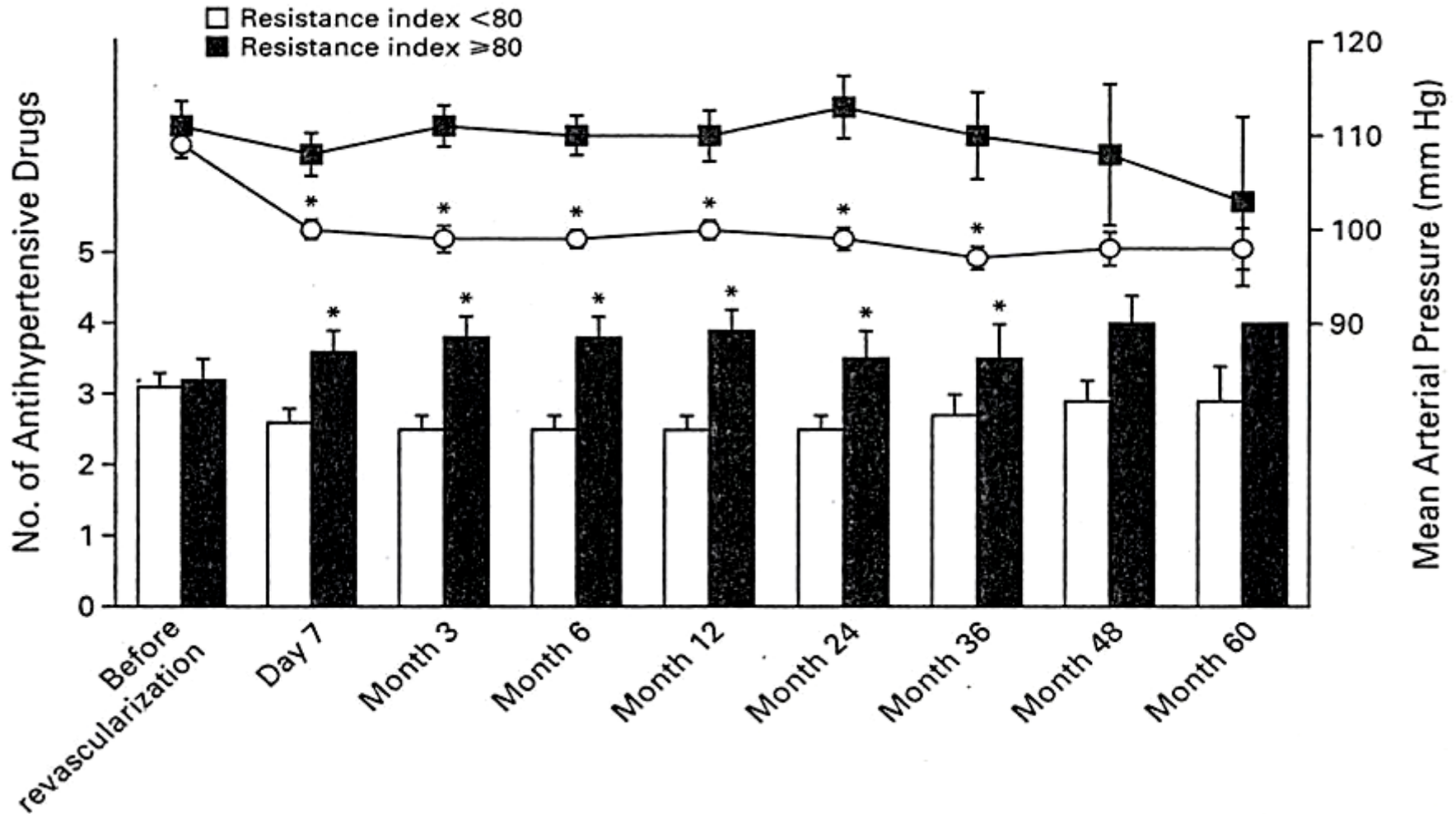


# Effect of various durations of ischemia on proteinuria development

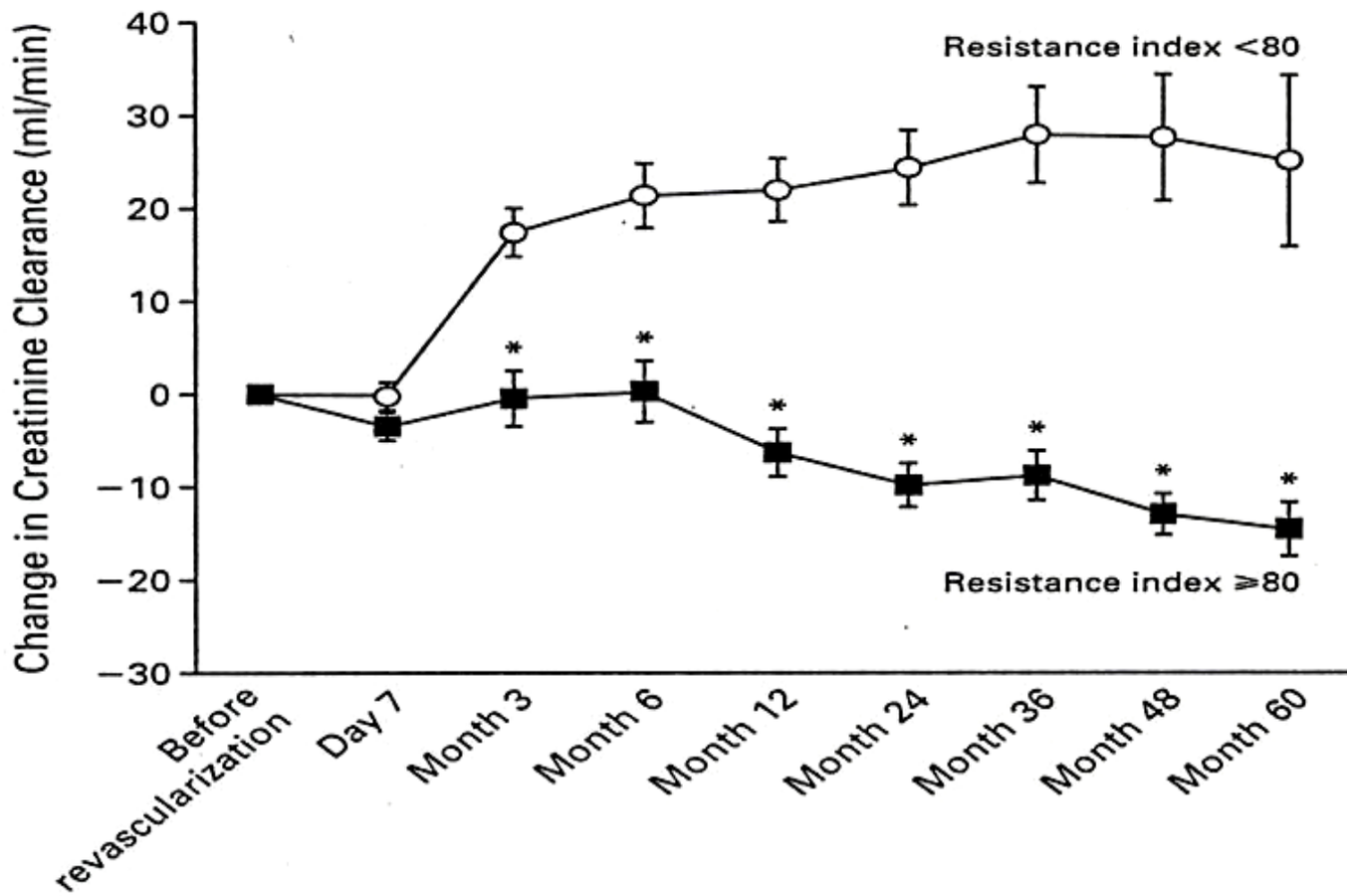




## Mean ( $\pm$ SEM) change in MAP and the number of antihypertensive drugs taken after the correction of RAS, according to resistance index values before revascularization



# Mean changes in creatinine clearance after the correction of RAS, according to resistance index values before revascularization

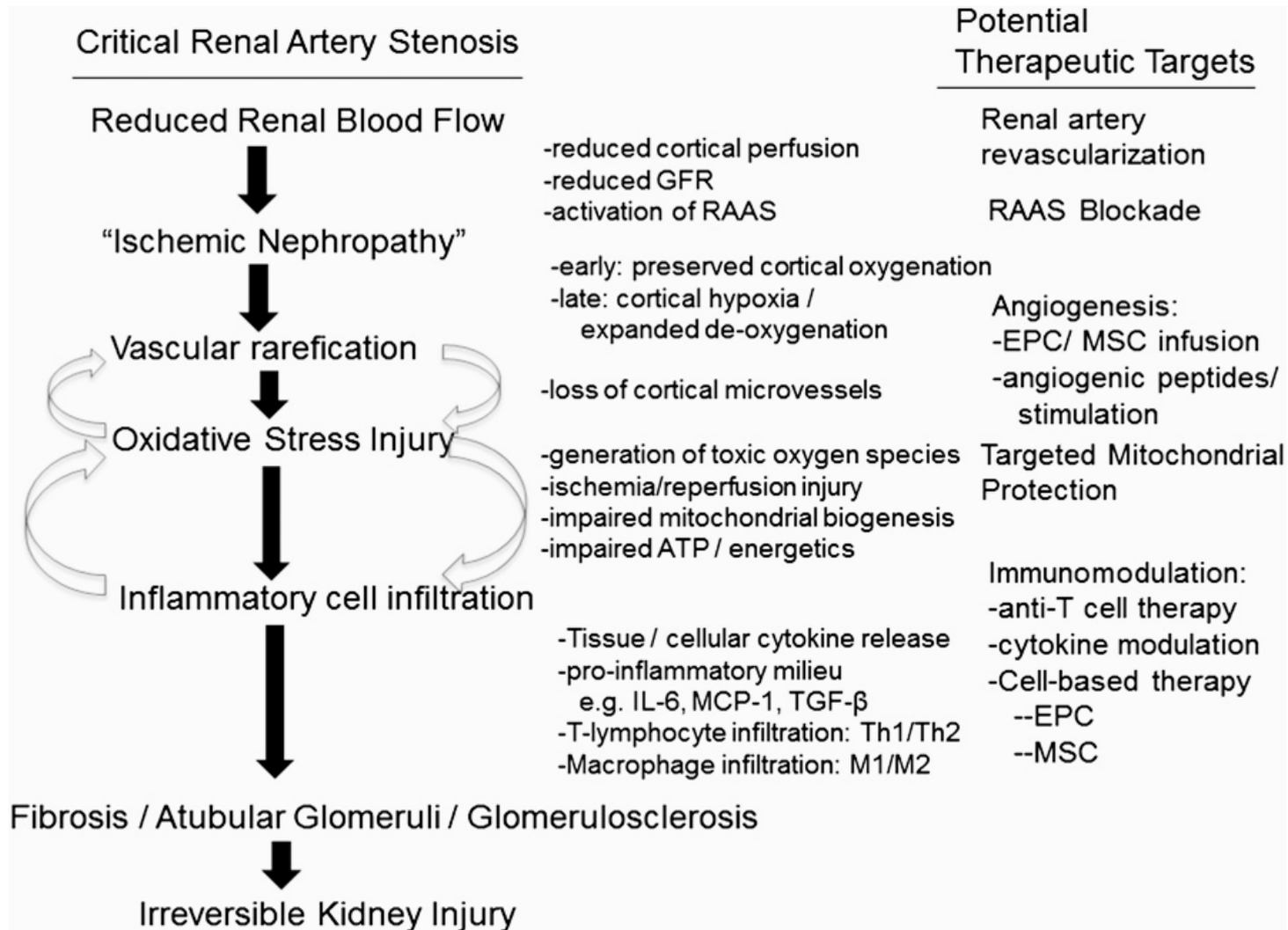


NO. WITH FOLLOW-UP DATA

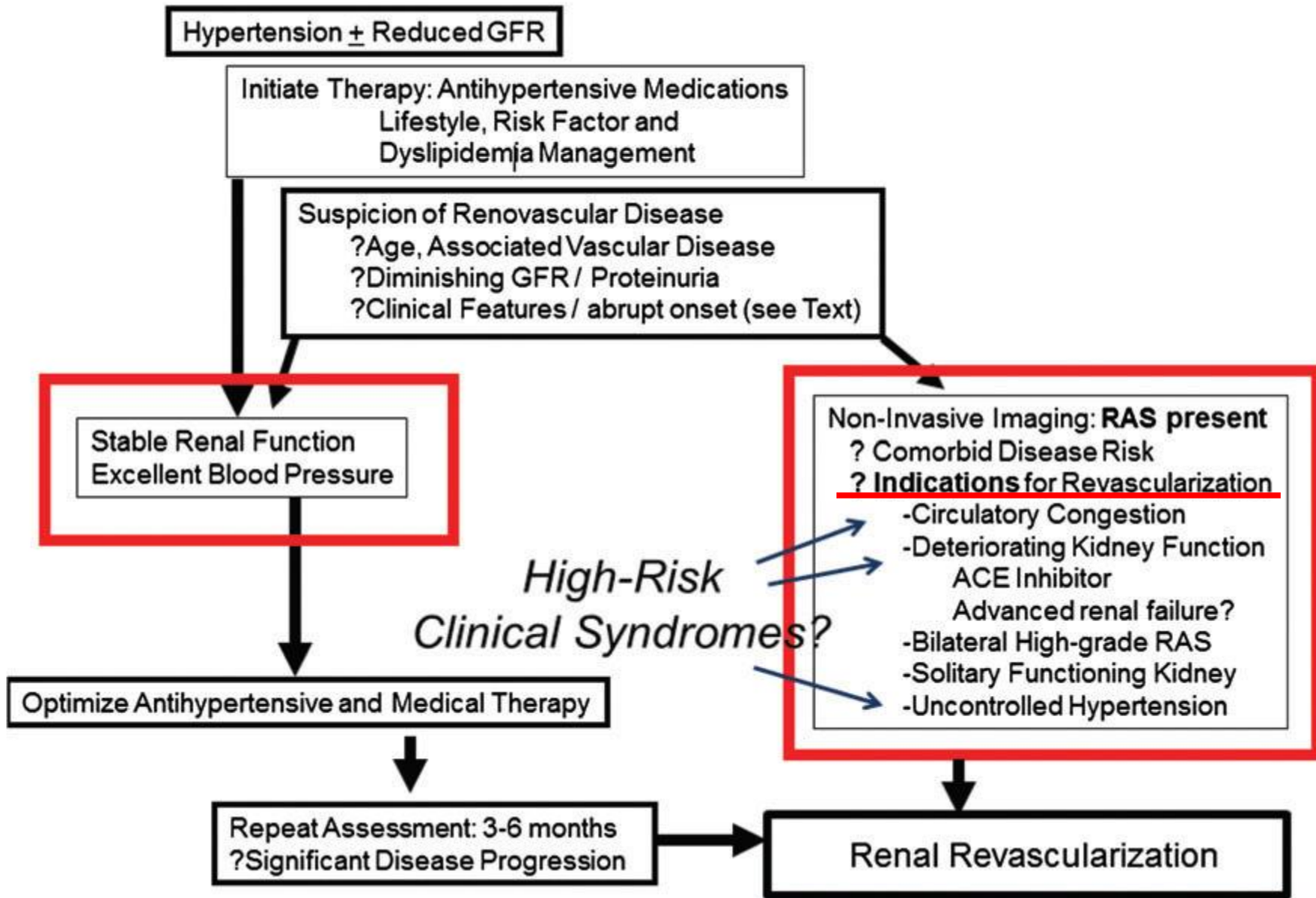
|                      |    |    |    |    |    |    |    |    |    |
|----------------------|----|----|----|----|----|----|----|----|----|
| Resistance index <80 | 96 | 96 | 95 | 83 | 73 | 59 | 43 | 34 | 21 |
| Resistance index ≥80 | 35 | 35 | 33 | 31 | 26 | 21 | 16 | 8  | 5  |



# Injury pathways and targets in Atherosclerotic Renovascular Disease (ARVD)



# Management of Renovascular Hypertension and Ischemic Nephropathy





# Renovascular Hypertension Revisited: To intervene or not?

- Therefore, the assumption that restoring renal artery patency always protects the kidney is false. In addition, sometimes the revascularization procedure causes a worsening of kidney function, in addition to the high risk of major complications and increased economic costs associated with the procedure



## Factors favoring medical therapy plus revascularization for renal artery stenosis:

- Progressive decline in GFR during treatment of hypertension
- Failure to achieve adequate BP control with optimal medical therapy
- Rapid or recurrent decline in GFR in association with a reduction in systemic pressure
- Decline in GFR during therapy with ACE inhibitors or ARBs
- Recurrent congestive heart failure in a patient in whom left ventricular failure does not explain the cause (flash pulmonary edema)



# Factors favoring medical therapy and surveillance of renal artery disease

- Controlled BP with stable renal function
- Stable renal artery stenosis without progression on surveillance studies (e.g., serial duplex ultrasound)
- Advanced age and/or limited life expectancy
- Extensive comorbidities that make revascularization too risky
- High risk for or previous experience with atheroembolic disease
- Other concomitant renal parenchymal diseases that cause progressive renal dysfunction (e.g., diabetic nephropathy) or severely reduced kidney size (< 7.0 cm)





Leading  
European  
Nephrology

