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Optimal management of renal artery fibromuscular dysplasia

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Abstract: Fibromuscular dysplasia (FMD) is a nonatherosclerotic, noninflammatory angiopathy of unknown cause affecting medium-sized (most commonly renal) arteries and causing renovascular hypertension. The most common medial multifocal type of FMD (with the "string of beads" appearance) is more than four times more prevalent in females than in males. FMD accounts for up to 10% of cases of renovascular hypertension. Compared with patients with atherosclerotic renal artery stenosis, patients with FMD are younger, have fewer risk factors for atherosclerosis, and a lower occurrence of atherosclerosis in other vessels. The etiology is multifactorial, including vessel wall ischemia and smoking, as well as hormonal and genetic factors. Intra-arterial digital subtraction angiography is still the gold standard for exclusion or confirmation of renal artery stenosis caused by FMD, at least in young patients, who more often have lesions in branches of the renal artery. For FMD patients with atherosclerosis and those who are older (>50–55 years), significant renal artery stenosis may be confirmed or excluded with ultrasonography. The FMD lesion is typically truncal or distal, whereas atherosclerotic lesions are more often proximal or ostial. Treatment options are medical, endovascular (percutaneous transluminal renal angioplasty [PTRA]), and surgical. Invasive treatment should be considered when hypertension cannot be controlled with antihypertensive drugs and in patients with impaired renal function or ischemic nephropathy. PTRA has become the treatment of choice and normally yields good results, especially in unifocal disease and young patients. Pressure gradients are normally completely abolished, and there is no indication for stent placement. Surgical revascularization is indicated after PTRA complications; thrombosis, perforation, progressive dissection, repeated PTRA failure or restenosis. Centralization of handling is recommended.

Keywords: fibromuscular dysplasia, renal artery stenosis, arterial hypertension, percutaneous transluminal renal angioplasty, renal artery surgery

Background
One criticism of the ASTRAL (Angioplasty and Stenting for Renal Artery Lesions) and STAR (STent placement and blood pressure and lipid-lowering for the prevention of progression of renal dysfunction caused by Atherosclerotic ostial stenosis of the Renal artery) trials in atherosclerotic renal artery stenosis is that patients with moderate hypertension and a moderate degree of stenosis were included, making the results uncertain. In the recent CORAL (Cardiovascular Outcomes in Renal Atherosclerotic Lesions) trial, the effects of percutaneous renal artery stenting combined with medical therapy were compared with those of medical therapy alone in 950 patients with pronounced hypertension and at least 60% stenosis. This trial replicated earlier findings of no benefit of renal artery stenting with regard to cardiovascular events or renal...
function, establishing beyond reasonable doubt that patients with <70%–80% atherosclerotic renal artery stenosis, stable renal function, and up to two antihypertensive drugs should not undergo renal artery stenting since there is no gain.

These new data might also have relevance in terms of treatment recommendations for patients who have renovascular hypertension with an etiology other than atherosclerosis. This review of fibromuscular dysplasia (FMD), a nonatherosclerotic, noninflammatory angiopathy of unknown cause affecting medium-sized (most commonly renal) arteries and causing renovascular hypertension, is based on information that is as current as possible.

Epidemiology
Symptomatic renal artery stenosis caused by FMD occurs in 0.4% of the population, but the prevalence of asymptomatic FMD in potential renal donors is 2.6%–4%.7,8 and bilateral in about one third of cases. FMD is most often diagnosed in individuals aged 15–70 years, but has been reported to occur from infancy to the age of 89 years.9 The medial multifocal type (Figure 1) of FMD is most common (68%–91%)8,11 and is more than four times more prevalent in females than in males, whereas the less common unifocal intimal type (Figure 2) is more prevalent in men.4,12 The male to female ratio is 1:4.5 according to the studies summarized in Tables 1 and 2. Bilateral lesions occur in about 25%–65% of cases, requiring bilateral treatment in 15%–25%.8,11 In unilateral disease, the right renal artery was found to be affected three times more often than the left.8 FMD accounts for up to 10% of cases of renovascular hypertension, with the remaining being caused mainly by atherosclerosis. Compared with patients who have atherosclerotic renal artery stenosis, FMD patients tend to be female, younger, and have fewer risk factors for atherosclerosis and a lower occurrence of atherosclerosis in other vessels.8,11 FMD is diagnosed more often in whites and less frequently in Hispanic and Asian populations. Few reports have been published from Africa and South America.11 In the US FMD Registry, which contains data for 447 patients, 95% were white, only 2% were African Americans, and 1% were Hispanic or Asian.8 Other vessels, such as the carotid, vertebral, mesenteric, and coronary arteries, were frequently affected. Two vascular beds were found to be affected in 35% of patients and three in 22%, underscoring the need for a good clinical examination and at least carotid artery investigation in FMD patients.8

The causes of pediatric renal artery stenosis differ in different populations.9,14–16 FMD is the most common cause of pediatric renovascular hypertension in North America and Western Europe, whereas Takayasu’s arteritis dominates in Asia and Africa.

Etiology
Vessel wall ischemia seems to be of relevance for the development of FMD.17 The vasa vasorum of the muscular arteries supplying oxygen and nutrients to the arterial wall originate from branch points of the parent arteries. Experimental occlusion of the vasa vasorum induces formation of dysplastic lesions.19 The vessels most frequently affected by FMD, ie, the renal, internal carotid, and vertebral arteries, have long segments that lack branches.

The female preponderance of FMD suggests the importance of hormonal factors such as estrogen and progesterone.8,19,20 In one study, 16% of women with FMD had previous hypertension in pregnancy compared with 4%–5% of hypertensive pregnancies in the general population.21 FMD has also been diagnosed after ovarian stimulation.22 However, the number of pregnancies and frequency of use of oral contraception or hormone therapy is not higher in FMD patients than in the general population.8,22,23 Young men with FMD often have
gynecomastia, further suggesting hormonal imbalance as part of the etiology of FMD.

In the US FMD Registry, 37% of patients were current or former smokers compared with the rate of 18% reported for US women. In a case-control study of 337 FMD patients, 30% smoked compared with 18% of controls. However, the mechanisms by which smoking contributes to FMD have not been elucidated.

The occurrence of renal FMD in siblings and identical twins suggests autosomal dominant inheritance, with incomplete penetrance and variable clinical symptoms. Further, a US FMD Registry study reported a 7% incidence in relatives. Importantly, stroke (54%), aneurysm (24%), and sudden death (20%) were also common in first-degree and second-degree relatives. FMD can be easily overlooked in asymptomatic relatives. Polymorphisms in the angiotensin-converting enzyme (ACE) insertion allele, ACE-I, and associations with HLA-Drw6 have been reported. Several groups are presently delineating other gene patterns predisposing to development of FMD.

### Differential diagnosis

Important differential diagnoses rely on associated phenotypic traits, including: the characteristic skin lesions in type 1 neurofibromatosis; acrogeric dysmorphism, skin elasticity, and distal joint laxity in vascular Ehlers-Danlos syndrome; and facial dysmorphism, supra-aortic stenosis, and the behavior characteristic of Williams syndrome. Genetic tests can also be used to rule out these conditions. Vasculitis is normally associated with an acute phase reaction.

### Pathology

Histologically, the three main types of dysplasia in FMD are classified according to the arterial wall layer mainly

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### Table 1 Results of percutaneous transluminal renal angioplasty for fibromuscular dysplasia in adults

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<th>Series</th>
<th>Year</th>
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<th>Female (n)</th>
<th>Cure (%)</th>
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**Abbreviation:** NA, data not available.
affected by collagen deposition, ie, intimal, medial, and adventitial. The intimal form accounts for about 10% and occurs without sex difference. Medial dysplasia (accounting for about 80% of cases) occurs as areas with a thinned medial wall layer alternating with thickened fibromuscular ridges containing collagen (Figure 3). Adventitial FMD is uncommon. However, few dysplasias are currently examined histologically, so an angiographic diagnostic classification has been suggested based on differentiation between unifocal and multifocal appearance. Unifocal FMD (Figure 2) has less of a female predominance and is diagnosed more often in younger individuals, with better short-term and long-term results from treatment than multifocal FMD (Figure 1).

Pathophysiology

In the multifocal type, multiple septa in the renal arteries may together induce a significant reduction in renal perfusion in patients with FMD, resulting in renovascular hypertension, but the degree of renal artery stenosis is impossible to evaluate from imaging. Subsequent reduction of arterial perfusion pressure by the unifocally or multifocally stenosed renal artery leads to activation of the renin–angiotensin–aldosterone system, with volume expansion and hypertension. This triggers mechanisms such as increased production of endothelin-1, local activation of the renin–angiotensin–aldosterone system, arterial wall remodeling, and oxidative stress, all helping to sustain the hypertension, which now depends on secondary factors such as central hemodynamics and peripheral resistance.

Table 2 Results of open surgery for fibromuscular dysplasia in adults

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not only on the renin–angiotensin–aldosterone system but also on the local vasoconstrictive and importantly proliferative effects in the arterial/arteriole wall, gradually leading to resistance to therapy.\(^3\)\(^4\) Inflammatory mediators are increased in renovascular hypertension,\(^3\)\(^5\) but neopterin and endothelin-1 levels are lower in renovascular hypertension due to FMD than in atherosclerotic renal artery stenosis.\(^3\)\(^5\)

**Natural history**

Data on the risk of progression of stenosis and deteriorating renal function are scarcer in patients with FMD than in those with atherosclerotic renal artery stenosis, but indicate that progression is less severe in FMD.\(^2\) Serial angiograms have confirmed progression of FMD in all\(^3\)\(^6\) or up to 40% of cases,\(^3\)\(^7\) and about 25% of subjects with asymptomatic FMD develop hypertension within 4 years.\(^6\)\(^7\) Aneurysms\(^8\) and dissection\(^8\) are rather frequent as secondary manifestations of FMD, but complete occlusion,\(^3\)\(^8\) renal infarction,\(^3\)\(^9\) severe renal insufficiency,\(^2\)\(^1\)\(^0\) and regression of stenosis\(^4\)\(^1\) have been reported infrequently.

**Clinical picture and physical examination**

Arterial hypertension of acute onset or high blood pressure that is increasingly difficult to treat suggests the presence of a specific cause of blood pressure elevation. Such secondary hypertension can be identified in about 5% of adult hypertensive patients.\(^1\)\(^3\)\(^2\)\(^4\) Renovascular hypertension caused by one or more stenoses of the extrarenal arteries is the second most common cause of secondary hypertension (after renal parenchymal disease) and occurs in approximately 2% of adult hypertensive patients referred to specialized centers.\(^1\)\(^3\)\(^2\)\(^4\)

**Blood pressure measurement**

Blood pressure should be measured at least twice in the sitting position, and home or 24-hour blood pressure measurements are encouraged to rule out white coat hypertension.\(^1\)\(^3\) Measurement of serum creatinine, calculation of estimated glomerular filtration rate, and testing for microalbuminuria are recommended in all hypertensive patients.\(^1\)\(^3\)\(^2\)\(^4\)

**Diagnostic evaluation**

The recent multicenter studies\(^1\)\(^–\)\(^3\) have implications for optimal selection of patients with hypertension and renal artery stenosis for medical, endovascular, and even open surgery. All diagnostic tools are in some way subjective, but at least in atherosclerotic cases, correct diagnosis of moderate to severe hypertension and renal artery stenosis of at least 60%–70% of the lumen area is important because treatment of stenosis of less than 60-70% leads to no clinical benefit. Appropriate patient selection is crucial to obtain clinical benefit from revascularization of hemodynamically relevant renal artery stenosis. Importantly, there is a difference in work-up according to whether atherosclerotic renal artery stenosis or FMD is suspected.\(^4\)\(^5\)\(^7\)\(^1\)\(^2\)\(^4\)

**Renal ultrasound**

Ultrasonography allows determination of longitudinal kidney size. A difference of >1.5 cm between the sizes of the two kidneys is usually considered diagnostic and is found in 60%–70% of patients with renovascular hypertension.\(^4\)\(^4\)

Color Doppler ultrasonography can often detect stenosis of the renal artery, particularly when localized close to the vessel origin.\(^4\)\(^5\) It also allows determination of the resistance index, which predicts the outcome of angioplasty and stenting in atherosclerotic renal artery stenosis.\(^4\)\(^5\) An increased resistance index suggests structural abnormalities in the small vessels of the kidney, and benefit from renal revascularization has been reported in patients with a resistance index <80.\(^4\)\(^6\) However, this index has not been evaluated in FMD patients. In a study of more than 300 patients using both duplex ultrasonography and angiography, peak systolic velocity and the systolic renal to aortic ratio was accurate for diagnosing a 60% renal artery stenosis. Specific data for 14 FMD patients in this study were not provided.\(^4\)\(^5\) In atherosclerotic renal artery stenosis, ultrasonographic evaluation may therefore identify the majority of patients with less than 60%–70% stenosis, in whom further investigations are not needed.

However, more caution is warranted in patients with suspected secondary hypertension caused by FMD. Limitations of duplex ultrasonography of the renal artery include its dependence on operator skill, the fact that it may be unsuccessful in 20% of patients because of problems visualizing branches or accessory renal arteries, and the difficulty or impossibility of imaging obese patients or those with intervening bowel gas.\(^4\)\(^7\) Because of these limitations and the fact that the distal renal artery and branch lesions are not optimally diagnosed by ultrasound, this technique cannot exclude FMD lesions. Changes in the mid section of the renal artery may well indicate a FMD lesion, but can also be explained by vessel tortuosity.\(^4\)\(^8\) Intravascular ultrasound is useful for distinguishing renal FMD from vasculitis and hypoplasia.\(^4\)\(^9\) However, the method is in limited clinical use.
Captopril renography
The sensitivity and specificity of captopril renography decrease in the presence of renal failure, bilateral disease, branch artery lesions, and a solitary functioning kidney. Although data exist from studies of patients with renal artery stenosis due to FMD, captopril renography is no longer recommended as a first-line method owing to its low sensitivity and specificity, and the same applies to renin measurements.\textsuperscript{13,50}

Magnetic resonance angiography
Magnetic resonance angiography (MRA) is a noninvasive imaging technique. A negative contrast-enhanced MRA probably excludes significant renal artery stenosis, but false-positive results due to turbulence are common, so MRA may overestimate the degree of renal artery stenosis.\textsuperscript{11,12,51} MRA can only identify vessels as small as 2–2.5 mm, so has limited resolution for distal and intrarenal arteries.\textsuperscript{11,21,52} Gadolinium-enhanced MRA has better specificity and sensitivity than ultrasound or scintigraphy, and is as efficient as computed tomographic (CT) angiography for detection of renal artery stenosis.\textsuperscript{59} However, given that the resolution of MRA is inadequate for visualization of branch vessel involvement, its usefulness for exclusion of FMD has been questioned (Figure 4).\textsuperscript{11} Gadolinium toxicity also precludes use of MRA in patients with severe renal impairment. Angeretti et al\textsuperscript{11} reported that breath hold contrast-enhanced MRA is equal in diagnostic accuracy to noncontrast-enhanced electrocardiography gated steady-state free precision and respirator triggered MRA, and even considered maximum intensity projections as being superior.

CT angiography
CT angiography has sensitivity comparable with that of MRA, but requires radiation and infusion of a potentially nephrotoxic contrast substance. CT angiography depicts both the lumen and the vessel wall but has limited resolution for the distal and intrarenal arteries.\textsuperscript{7,12} In some centers, MRA or CT angiography is now used as premapping before invasive treatment after duplex ultrasonography has suggested renal artery stenosis.\textsuperscript{50} Nevertheless, when the index of clinical suspicion is high, neither test can exclude FMD. None of these methods is reliable in the 25% of FMD patients with intrarenal stenosis.

Renal angiography
When one of the aforementioned screening examinations raises strong suspicion of renal artery stenosis, or when a fairly young FMD patient is evaluated and there is a strong clinical indication to proceed to revascularization, intra-arterial digital subtraction angiography should be performed for exclusion or confirmation. This invasive procedure is still the gold standard for detection of renal artery stenosis caused by FMD.\textsuperscript{7,11} FMD lesions are typically truncal or distal, whereas atherosclerotic lesions are more often proximal or ostial.\textsuperscript{4,32} Angiography is especially useful in evaluation of branch vessel disease. Angiography is a two-dimensional analysis and therefore gives only an educated estimate of the degree of stenosis, which might help explain the difficulties in selecting patients who would benefit from treatment in recent trials.\textsuperscript{1–3} Additionally, the potential hemodynamic significance of the multiple stenoses often seen in FMD is almost impossible to evaluate.

Pressure gradients
Intra-arterial measurement of the pressure gradient over the stenosis, such as performed before percutaneous transluminal angioplasty in the iliac arteries,\textsuperscript{54} is recommended in renal artery stenosis.\textsuperscript{55} Different methods for measuring pressure gradients have been used,\textsuperscript{56,57} and there is no consensus regarding the level indicating a hemodynamically significant stenosis. A distal aortic pressure ratio \(<0.90\) has been correlated with increased renin levels in the renal vein,\textsuperscript{58} but pressure gradients could not be used to select patients for successful recanalization of atherosclerotic renal artery stenosis in the recent CORAL trial.\textsuperscript{3} However, prolonged hypertension leads to proliferative effects in the arterial/arteriole wall, resulting in resistance to therapy. Nevertheless, a mean pressure gradient of \(>10\text{ mmHg}\) over the stenosis predicts a favorable response to dilatation, and a persistent pressure gradient after an endovascular procedure indicates a higher risk of restenosis.\textsuperscript{59}

Figure 4 Magnetic resonance angiography, raising suspicion for bilateral fibromuscular dysplasia.

Notes: The right renal artery changes caused significant pressure gradients and were treated with percutaneous transluminal angioplasty. Note the absence of intrarenal branches, that are affected in 25% of patients with fibromuscular dysplasia and are not visualized with magnetic resonance angiography.
Treatment

The treatment options for renal FMD are medical, endovascular, and surgical. Treatment of all renovascular hypertension is controversial due to difficulties in predicting the blood pressure response to renal revascularization in individual patients.\textsuperscript{1–3,7,11} In FMD, invasive endovascular or surgical treatment should be considered when hypertension cannot be controlled with medication, in those who are intolerant of or noncompliant with medication, and in those with impaired renal function or ischemic nephropathy.\textsuperscript{7,11}

Medical therapy

Controlled trials have convincingly shown that lowering blood pressure reduces cardiovascular morbidity and mortality.\textsuperscript{11,60} All patients with renovascular hypertension caused by FMD are therefore candidates for antihypertensive treatment in accordance with current European\textsuperscript{13} and US\textsuperscript{42} guidelines. Blood pressure should be reduced to <140/90 mmHg in hypertensive patients, and according to European recommendations,\textsuperscript{11} further to <140/85 mmHg in patients with diabetes. Current guidelines\textsuperscript{13,42} specify five groups of first-line antihypertensive treatment, ie, ACE inhibitors, angiotensin II receptor blockers, calcium channel blockers, diuretics, and (in the European guidelines) beta blockers.\textsuperscript{13} All of these drugs can be used in the treatment of renovascular hypertension due to FMD.

Thiazide diuretics and calcium antagonists can be used with possible addition of a renin–angiotensin–aldosterone system blocker, an ACE inhibitor, or an angiotensin II receptor blocker. However, these should not be used during pregnancy. Such treatment lowers blood pressure in the majority of patients with renovascular hypertension. In patients with bilateral disease or stenosis of a single kidney, blockade of the renin–angiotensin–aldosterone system requires caution because of the risk of compromised renal function.\textsuperscript{61}

Endovascular treatment in adults

Percutaneous transluminal renal angioplasty (PTRA, Figures 2, 3, and 5) has become the treatment of choice for renovascular hypertension caused by FMD.\textsuperscript{11,62} The main reason for treating FMD is unsatisfactorily controlled hypertension. Progressive loss of renal function is less likely in FMD than in atherosclerotic renovascular hypertension.\textsuperscript{63}

Technical considerations

The micropuncture technique should be used to avoid groin hematoma, which is the most common complication (3%–26%)\textsuperscript{11,64–66} but often preventable complication of PTRA. To ensure optimal compression after the procedure, the puncture position should be angiographically controlled before “larger” catheters or introducers are advanced. Low-profile 0.014 inch or 0.018 inch guidewire systems are used to reduce vasospasm. FMD patients are often younger and have more pronounced vasoreactivity than those with atherosclerosis. Therefore, renal vasospasm is common, and premedication with dihydropyridine or repeated doses of nitroglycerin 0.15 mg may be required in hemodynamically stable patients, not only after catheterization. FMD stenoses most often occur in the middle portion of the main renal artery. If the distal renal artery or its branches are affected, use of multiple guide wires may be necessary. The kissing balloon technique can be used in the branches.\textsuperscript{64,67}

Stenting

PTRA alone normally provides good results, in that pressure gradients are completely abolished and there is no indication for stent placement (Table 1).\textsuperscript{14} Indications for stenting of FMD lesions include severe procedural complications, persistent pressure gradients after repeated attempts at angioplasty, and small aneurysms in the renal artery.\textsuperscript{14} No value of drug-eluting stents has been reported.

Procedure-related complications

Groin hematoma is the most common complication, and dissection occurs infrequently (in 1.4%–6.7% of procedures).\textsuperscript{11,21,65} Few cases require repeated or prolonged balloon treatment, and only extensive expanding dissection requires stent deployment and fixation.\textsuperscript{11}

Rupture of the renal artery is uncommon (2%–6%)\textsuperscript{11,21,65} and occurs most often in association with complex FMD lesions. Some of these small ruptures stop leaking after
prolonged balloon treatment, whereas others require stents, covered stents, or emergency open surgery after the balloon has achieved initial hemostasis.

The use of cutting balloons is definitely not a first choice in patients with FMD given that ruptures, especially of hypoplastic arteries, have been reported with their use.\textsuperscript{68–70} CT angiography, MRA, and intravascular ultrasound may add information on hypoplasia and wall thickness.

Branch occlusion is seen in 1\%–5\% of patients with FMD after PTRA,\textsuperscript{1,21,65} but most of these patients have a favorable outcome. Thrombosis of the main renal artery after endovascular reconstruction in adults is very rare. Bonelli et al\textsuperscript{66} reported occurrence of minor nonocclusive thrombotic material in the renal artery after PTRA in five of 105 patients with FMD.

**Early outcome**

Noncontrolled series on endovascular treatment of FMD have included >1,500 patients (Table 1), with two studies reporting on more than 100 patients each.\textsuperscript{65,71} The age and sex distribution differ in these reports, and it is hard to determine which types of FMD were treated, which affects outcome. In addition, recommendations on how to present outcome criteria have changed over time, making interpretation of results even more difficult.\textsuperscript{72} The male to female ratio in the compiled data (Table 1) was 1:4.4, and 25\% of cases had bilateral FMD. Some studies have also reported on the diseased side in unilateral FMD.\textsuperscript{1,21,73} The right renal artery was affected in 81\% of cases, and the left renal artery in 19\%.

PTRA can be performed with good results (Table 1), not only for main artery lesions but also when branch arteries are affected.\textsuperscript{21,66} However, patients with branch artery involvement show less pronounced blood pressure responses and higher restenosis rates.\textsuperscript{21}

Comparisons of different series are limited because of the large variation in selection criteria and length of follow-up. “Cure”, “improvement”, and “benefit” are also defined differently.\textsuperscript{72} In a systematic review of FMD treatment,\textsuperscript{11} cure rates were around 65\% in 30-year-old patients and only about 20\% in 55-year-old patients. This obvious reduction in cure rate with increasing age is also reported elsewhere.\textsuperscript{32,74,75} The systematic review also reported a decrease in cure rate from approximately 60\% in a 1980–1985 series to about 30\% in more recent series, perhaps reflecting changes in the definition of cure.\textsuperscript{11} Treatment is likely to cure hypertension in young individuals, but stricter criteria for invasive treatment should be used in older patients with FMD. The more strict treatment criteria recommended for atherosclerotic renal artery stenosis\textsuperscript{1–3} might perhaps be considered in older FMD patients.

**Late outcome**

Renal function was unchanged during follow-up.\textsuperscript{32} Long-term durability and the initial effect after PTRA is good in patients with the “string of beads” appearance in the main artery; however, for more complex and branch lesions, the results are acceptable but redo procedures are needed more frequently (Figure 5).

Bonelli et al reported that 6\% of their 105 patients required redo procedures during 4 years of follow-up,\textsuperscript{65} while Alhadad et al found that although 34\% of their patients required a second and often planned PTRA procedure, only two patients required redo PTRA after more than 4 years of follow-up.\textsuperscript{21} The one-year cure rate is dependent on the angiographic type of FMD; 15/28 (54\%) patients with unifocal FMD versus only 13/50 (26\%) with multifocal FMD were cured in a report by Savard et al.\textsuperscript{32}

**Surgery**

Surgical revascularization for FMD is currently reserved for patients with severe complications of PTRA, such as thrombosis, perforation, or dissection that cannot be handled with endovascular techniques. Open surgery should also be considered after repeated failure or restenosis following PTRA, recoil, and no effect on stenosis following endovascular treatment. In experienced groups, the results of operative treatment after failure of PTRA are favorable.\textsuperscript{75–79}

**Techniques**

Details on reimplantation of a renal artery, aortorenal bypass, and autotransplantation are given elsewhere.\textsuperscript{14} Centers performing endovascular treatment need to have open operative techniques available, demanding further centralization of PTRA. The need for centralization will become more evident as recent published multicenter studies\textsuperscript{1–3} would probably reduce the number of referrals for atherosclerotic renal artery stenosis.

**Results of open surgery**

Published series on open surgery (Table 2) include more than 1,200 patients and report on aortorenal bypass,\textsuperscript{77–80} branch artery reconstructions,\textsuperscript{81,82} and autotransplantation.\textsuperscript{78,83,84} Regardless of the method used, the early outcome is excellent. The more recent series mainly include patients who have undergone unsuccessful PTRA. The proportion of cured patients has been somewhat lower during the last
Complications of open surgery
Renal failure and a fatal outcome are both very uncommon after surgery for FMD. Overall morbidity from surgery is 19%–28%, and is mainly caused by minor complications such as urinary tract infection and postoperative pneumonia. Early postoperative occlusion after FMD reconstruction is seen in 3.8%–13% of cases, and more often with venous grafts than with arterial autografts. Reiher et al detected nine unsuspected thromboses in 90 reconstructed patients on routine postoperative angiography.

Late restenosis and the need for redo procedures have become less common as most anastomoses are now made in an ovoid shape. Restenosis has been reported in 0%–16% of cases, and 1%–15% of patients require a redo procedure. Late vein graft dilatations were reported in 20%–44% in an early series of aortorenal bypass in adults, but not in the last decade, perhaps reflecting that FMD patients are now older and have a longer duration of hypertension. Pseudoaneurysms (Figure 6) are infrequently reported.

FMD and aneurysms
A large proportion of patients have FMD with the “string of beads” appearance, ie, small aneurysmal areas interspaced by webs. “Macroaneurysms” with obvious aneurysmal development are not infrequent; however, 8.5%–26.7% of patients with FMD have aneurysmal changes. Most renal artery aneurysms have a low risk of rupture, and few ruptured FMD lesions have been reported. How FMD-related aneurysms measuring 1–2 cm should be handled is uncertain. Conservative treatment may be successful. Aneurysms larger than 2 cm should probably be excluded. Renal artery aneurysms have occurred in 5%–12% of pediatric patients with FMD undergoing open surgery.

The endovascular experience is limited. Small aneurysms may shrink after successful PTRA of stenotic areas, and favorable outcomes have been reported with embolization and stent grafts. Reid et al reported recently on 44 renal artery aneurysms, of which half were openly reconstructed, 19 were embolized with or without stent placement, and four were treated with covered stents. Aneurysmal size (2.5 cm versus 2.2 cm), complications, and renal function did not differ during 21–27 months of follow-up.

Pediatric renal artery stenosis and arterial dysplasia
Renovascular disease is the most common cause of hypertension in children after thoracic aortic coarctation and renal parenchymal disease. Causes of renovascular hypertension are FMD, developmental renal artery stenosis, vascular neurofibromatosis type 1, Moyamoya disease, vasculitis such as Takayasu’s disease, Alagille syndrome, and Williams syndrome. Up to 40% of children with renovascular hypertension have developmental renal artery lesions with mid aortic syndrome and coexisting lesions in both the mesenteric and carotid arteries.

Endovascular treatment in children
Published series on endovascular treatment in children and adolescents have included 237 patients, 150 of whom had FMD. The sex distribution in children is equal. Technical success rates are high in published series, with cure or improvement in about 80%–90% of cases. The majority of children have medial or perimedial dysplasia; however, the “string of beads” type of lesion is not frequent in children. Shroff et al reported long-segment stenosis in 60%, ostial stenosis in 22%, ostial and long-segment stenosis in 12%, aneurysmal dilatation in 6%, and webs in 6% of their series. McTaggart et al reported on 17 children, eight of whom had ostial lesions; the main renal artery was affected in eight cases, and three lesions were seen in branches.
Follow-up of PTRA in children aged 13 months to 12 years demonstrates variable outcome. Benefit is frequently reported, but restenosis occurs in 27%–33% of cases, and branch lesions often require redo PTRA. Treatment of very young children to achieve a moderate improvement may allow open surgery to be delayed until an older age, when the procedure can be performed with better long-term results.

**Open surgery for pediatric renovascular hypertension**

In the University of Michigan Pediatric Renovascular Group’s report, 12 of 97 children undergoing open surgery had multisegmental disease, six had aneurysms, 70 had ostial lesions, 15 had changes in the main artery, and 18 had branch involvement. Stenosis was unilateral in 58% of cases, and

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**Figure 7** Suggested management principles for fibromuscular dysplasia.

**Abbreviations:** FMD, fibromuscular dysplasia; CTA, computed tomography angiography; MRA, magnetic resonance angiography; PTRA, percutaneous transluminal renal angioplasty.
bilateral in 42%. Few girls younger than 10 years were treated, and in that age group, boys were three times more likely to be affected. No sex difference was noted in those aged 10–17 years.

Open surgery series have included 439 children and adolescents, 379 of whom had FMD. The early series included a large proportion (40%–50%) who underwent nephrectomy,97,98 whereas Stanley et al reported on 97 children with irreversible renal disease requiring planned (n=11) or unplanned nephrectomy (n=1) due to failure of a reconstruction.9 Centers of excellence9,99 advocate direct aortorenal anastomosis if feasible, which was possible in 41 of 58 reconstructions.9 In 85 children, Lacombe used direct anastomosis in 17 cases and splenorenal anastomosis in 22.91 Long-term follow-up (1–10 years) shows a benefit of surgery in about 95% of cases and a minimal restenosis rate, in spite of continuing growth. Centralization of endovascular procedures and surgery is required in children.

Conclusion
In summary, the diagnostic work-up in suspected secondary hypertension (Figure 7) should include renal artery ultrasound. When FMD is suspected, further examination with CT angiography or MRA is often warranted but, as with ultrasound, neither technique reliably visualizes FMD in the branch arteries. Renal angiography may be required, and allows for PTRA, which is the treatment of choice for FMD in patients with renovascular hypertension or renal dysfunction. However, indications for invasive treatment in older FMD patients should perhaps be narrowed in the future, as in atherosclerotic renal artery stenosis.

Disclosure
The authors report no conflicts of interest in this work.

References


