Williams-Beuren syndrome (WBS) is a microdeletion disorder caused by heterozygous loss of approximately 1.5-Mb pairs of DNA from chromosome 7. Patients with WBS have a characteristic constellation of medical and cognitive findings, with a hallmark feature of generalized arteriopathy presenting as stenoses of elastic arteries and hypertension. Human and mouse studies establish that defects in the elastin gene, leading to elastin haploinsufficiency, underlie the arteriopathy. In this review we describe potential links between elastin expression and arteriopathy, possible explanations for disease variability, and current treatment options and their limitations, and we propose several new directions for the development of nonsurgical preventative therapies based on insights from elastin biology.
Mechanisms and treatment of cardiovascular disease in Williams-Beuren syndrome

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Williams-Beuren syndrome (WBS) is a microdeletion disorder caused by heterozygous loss of approximately 1.5-Mb pairs of DNA from chromosome 7. Patients with WBS have a characteristic constellation of medical and cognitive findings, with a hallmark feature of generalized arteriopathy presenting as stenoses of elastic arteries and hypertension. Human and mouse studies establish that defects in the elastin gene, leading to elastin haploinsufficiency, underlie the arteriopathy. In this review we describe potential links between elastin expression and arteriopathy, possible explanations for disease variability, and current treatment options and their limitations, and we propose several new directions for the development of nonsurgical preventative therapies based on insights from elastin biology.

Williams-Beuren syndrome (WBS, also referred to as Williams syndrome; OMIM 194050) was first recognized as a clinical syndrome, separate from other developmental disability syndromes, because of a unique constellation of cardiovascular (CV) abnormalities. In this review, we describe these characteristic abnormalities, the considerable progress that has been made in understanding their etiology and pathophysiology, and new insights gained into underlying molecular pathways. Finally, we consider currently available therapeutic approaches and opportunities to develop new treatment options.

Overview of WBS

In 1961, J.C.P. Williams described four patients and proposed that “the association of supravalvular stenosis with the physical and mental characteristics here described may constitute a previously unrecognized syndrome” (1). Shortly thereafter A.J. Beuren reported eleven new patients (2), and the condition has fittingly borne the eponym Williams-Beuren syndrome ever since.

WBS affects approximately 1/10,000 individuals, is found worldwide among all racial and ethnic groups, and displays multisystem medical and nonmedical problems (3–5). In 1993 the genetic cause of WBS, a chromosomal microdeletion, was reported (6), and this knowledge permitted development of a laboratory-based diagnostic test, so the diagnosis no longer rests on clinical criteria only. The most widely used method to confirm the diagnosis has been FISH, but DNA dosage techniques such as quantitative PCR, multiplex ligation-dependent probe amplification, and chromosomal microarray, also known as comparative genomic hybridization, may soon become confirmatory tests of choice.

Non-CV clinical features of WBS. Persons with WBS have a subtle but distinctive facial appearance that changes with age (Figure 1) (3). Their linear growth is usually smaller than that of siblings or healthy, age-matched controls. Though infants and young children tend to be thin, or even underweight, many WBS adults are overweight (3, 7, 8).

Endocrine abnormalities are commonly reported and include hypercalcemia, abnormal glucose metabolism, and (subclinical) hypothyroidism. The precise frequencies, etiologies, natural histories, and best treatments of these endocrine problems remain to be determined. Other common findings include dental anomalies (small, abnormally shaped teeth, absent teeth, malocclusion), gastrointestinal dystmotility (reflux, constipation), diverticular disease, musculoskeletal anomalies (joint stiffness, scoliosis), sensorineural hearing loss, genitourinary anomalies (urinary frequency, bladder diverticuli), and neurological problems (abnormal tone, hyperreflexia, and cerebellar findings) (3, 7, 9, 10).

Persons with WBS have intellectual handicaps that generally meet the definition of mild to moderate mental retardation on standardized testing. The average full-scale IQ is reported to be 55–60, but a fairly broad range exists, extending from 40–90 (11). Particularly notable is the pattern of intellectual peaks and valleys, referred to as the Williams syndrome cognitive profile, with relative strengths in selected language domains and a prominent weakness in the visuospatial domain (12). Persons with WBS display characteristic personality and emotional traits. Their generally social and friendly demeanor coexists with anxiety (especially anticipatory anxiety), phobias, and perseverative tendencies (i.e., repetitive thoughts or behaviors) (7, 13).

Molecular genetics of WBS. The etiology of WBS, a mystery for more than 30 years after its initial description, is now known to be a contiguous gene deletion or microdeletion syndrome at chromosome 7q11.23. Although the chromosomal location is unique, the mechanism of origin is comparable with that of other microdeletion disorders, namely, a deleted genomic interval resulting from the presence of low-copy repetitive DNA (duplicons) that predispose to nonallelic homologous recombination (NAHR) (14).

The locus for WBS was found through the study of a phenotypically overlapping disorder. Specifically, disruption of the elastin (ELN) gene was identified as the cause of the condition known as familial supravalvular aortic stenosis (familial SVAS) syndrome (OMIM 131400).
185500) by application of linkage analysis and gene sequencing, and by cloning of a chromosome 7q11.23 translocation breakpoint, in selected familial SVAS kindreds (15, 16). Given similar vascular pathologies in familial SVAS and in WBS, study of the ELN gene was undertaken, and deletion of an ELN allele was identified as the cause of WBS (6). Subsequent characterization of the deleted interval, now referred to as the WBS critical region (Figure 2), reveals the following (17–19): (a) 90%–95% of patients clinically diagnosed with WBS have an approximately 1.55-Mb deletion associated with loss of 26–28 genes; (b) 5%–8% of clinically diagnosed WBS patients have a slightly larger, approximately 1.84-Mb pair deletion associated with loss of 28 genes; (c) the deleted intervals are flanked by highly homologous stretches of DNA, organized into a single centromeric duplicon and two telomeric duplicons; (d) each duplicon contains genes, pseudogenes, and clusters of related genes; (e) the duplicons predispose to NAHR through intra- or inter-chromosomal exchange during meiosis; and (f) the deletion arises with equal frequency on either the maternally or the paternally inherited chromosome 7 homolog.

Different breakpoints in the medial block of telomeric repeats determine whether neutrophil cytosolic factor 1 (NCF1) and general transcription factor II I repeat domain–containing 2 (GTF2IRD2) are deleted or not. Several of the deleted genes in patients with WBS are depicted in Figure 2. As this review focuses on CV aspects of WBS, our discussion will emphasize the ELN gene and consequences resulting from its functional loss. For a complete list and description of the genes constituting the WBS critical region, the reader is directed to ref. 20.

Loss of an ELN allele is the single most important genetic change responsible for the CV problems of WBS. The ELN gene encodes a precursor protein, tropoelastin. Multiple isoforms of tropoelastin are generated via alternative splicing, secreted into the extracellular matrix, deposited onto a preformed network of fibrillin microfibrils, and are cross-linked by the lysyl oxidase family of enzymes (21). This process of elastic fiber assembly requires the coordinat ed expression of multiple genes, and peaks during late fetal and perinatal stages of development (22).

The most common pathological consequence of chromosome 7q11.23 duplicon–mediated NAHR is WBS (e.g., deletion of the intervening sequence) (23) has recently been detected by chromosomal microarray; affected individuals have expressive speech delay without evidence of WBS features (24). The most common non-pathological consequence of NAHR at this locus is inversion of the WBS critical region (25). Inversion carriers are asymptomatic, though they may have a slightly greater risk of meiotic NAHR producing a gamete, which, when fertilized, results in an offspring with WBS (26).

The chromosome 7q11.23 genomic architecture in the WBS critical region has genetic synteny on mouse chromosome 5 (27). The salient differences are the opposite orientation of the region and the absence of low-copy repeats in the mouse. Such synteny provides a powerful tool for genotype-phenotype analysis through the generation and characterization of genetically modified mice; a complete list of currently published WBS mouse models is provided in Supplemental Table 1 (supplemental material available online with this article; doi:10.1172/JCI35309DS1). The Eln knockout mouse model is discussed in considerable detail below. A CV phenotype has not been appreciated in any other mouse in which a WBS critical region gene has been genetically altered, such as Fzd9−/− (28), Limk1−/− (29), Gtf2ird1−/− (30), or Cyln2−/− mice (31).

A complementary approach to elucidating the role of deleted genes in the WBS phenotype involves detailed characterization of patients with smaller atypical deletions. These patients, in whom less than 26 genes have been deleted from the WBS critical region (Figure 2), are relatively uncommon. Despite their rarity, such cases can provide valuable information. Characterization of patients with atypical deletions that variously encompass LIMK1, GTF2IRD1, and/or GTF2I suggest that loss of one or more of these genes contributes to the WBS cognitive profile (32–34). Given the combination of a relatively small number of genes in the WBS critical region and a remarkably distinct set of features, WBS has become a “model” disorder for the study of genotype-phenotype correlations in microdeletion syndromes.

**CV clinical features of WBS**

CV disease, particularly an arteriopathy consisting of stenoses of medium- and large-sized arteries, is the hallmark of WBS. In early case reports and case series, the diagnosis was primarily established...
by cardiologists evaluating patients for a heart murmur, which usually was diagnosed as SVAS on cardiac imaging or through catheterization and/or surgery (1, 35).

The prevalence of CV abnormalities in 423 patients from nine selected international series published in the last two decades is shown in Table 1 (36–44). Many patients have multiple CV clinical findings. Although the vascular stenoses of WBS predominantly affect the supravalvular aortic (Figure 3) and pulmonary regions, lesions located elsewhere also occur, primarily but not exclusively affecting the vascular branch points. The wide range of published prevalence reflects the age-dependent frequency of clinical features, variable study methods, especially the modality used to ascertain specific clinical features, and biases in case ascertainment. Spontaneous improvement in pulmonary arterial stenosis over time has been well established, whereas SVAS may progress especially in the first five years of life (36, 38–40, 45, 46). Prospective use of echocardiographic dimensions in one study found a 100% frequency of SVAS but only a 3% frequency of pulmonary arterial stenosis (37). Very rare patients have atypical deletions smaller than the common deletion. Schematics of atypical deletions are shown on right and include a very small deletion encompassing ELN and an adjacent gene; a typical centromeric breakpoint but not the common telomeric breakpoint; and a typical telomeric breakpoint but not the typical centromeric breakpoint. Not all genes are shown; see ref. 20 for a complete list of genes. WBSCR, WBS critical region.

A small number of patients with SVAS (as part of either WBS or familial SVAS syndrome) have a diffuse and more severe variant of arteriopathy, sometimes termed middle aortic syndrome (36, 42, 47). These patients have stenosis of the thoracic and abdominal aorta, mesenteric arteries, and renal arteries. Left ventricular hypertrophy and hypertension are common in this severe form of CV disease.

Clinically important coronary artery lesions (Figure 3) are reported infrequently but may put the patient at risk of sudden death. An autopsy series of five patients whose CV disease represented the severe end of the spectrum demonstrated obstructive coronary disease in all individuals, with vessel wall hyperplasia, fibrosis, and disorganization (48). In a catheterization study of 26 patients with WBS, coronary artery dilation or stenosis was found in 27% (39). In mixed cohorts of individuals who had either WBS or mutations in the \textit{ELN} gene in conjunction with severe SVAS requiring surgery, coronary disease was found in 28%–45% (49, 50). Although coronary disease appears to be related to the severity of supravalvular aortic narrowing (39), case reports indicate that severe coronary artery disease leading to death may be the sole vascular feature in some patients with WBS (51, 52). Since the sensitivity of standard non-stress echocardiographic studies for coronary stenosis in the general population is low, perfusion imaging techniques may be needed to investigate the significance of coronary artery disease in WBS (53).
Patients with WBS are at a higher risk of sudden death. Wessel et al. found a risk of 1/1,000 patient years with five cases of sudden death, a 25- to 100-fold increase compared with the normal population (54). Death secondary to myocardial infarction has been detailed in three patients (55, 56). In another report of ten WBS patients with sudden death, pathologic findings suggest that coronary artery stenosis and severe biventricular outflow tract obstruction are mechanisms for myocardial ischemia and arrhythmia (57). Many of the deaths occurred with anesthesia/sedation (often with cardiac catheterization), suggesting that decreased cardiac output from anesthetic agents in concert with coronary artery abnormalities in young patients with WBS in the absence of clinically significant outflow tract obstruction (67) parallels the mouse model findings (68). Cerebral arterial disease with stroke are described infrequently in WBS (69, 70), but the true frequency of intracranial vascular stenoses is not known.

Vascular pathology
Arterial abnormalities in WBS include localized or diffuse narrowing of elastic arteries. Diagnostic imaging (64, 71) and pathological studies (72, 73) have both demonstrated generalized arterial wall thickening even in nonstenotic regions of the arterial tree (Figure 4). These nonstenotic areas are characterized by an expansion of the media, caused by up to a 2.5-fold increased number of lamellar units (72), with relatively preserved organization of elastic lamellae and smooth muscle cells.

Three morphological types of SVAS are defined (74). Membranous SVAS is rare and consists of constricting semicircular valve-like membrane or membranes located at the sinotubular junction. The membrane contains small stellate cells in abundant mucopolysaccharide ground substance with few collagen and elastic fibrils but no medial elements (75). The second and third types of SVAS, the hourglass and diffuse types, share histological characteristics of disorganized lamellar architecture in the media, haphazard and fragmented elastic fibers, and focal clumping and hypertrophy of smooth muscle cells (Figure 4D) (75). In addition, enlargement of aorta vasorum has also been observed in the media and adventitia. Focal mural induration is sometimes observed. These regions, in

Other cardiac defects are occasionally observed in WBS. Ventricular septal defects, typically small, were found in 0%–14% of the series described in Table 1. Single cases of more complex defects such as tetralogy of Fallot (malalignment ventricular septal defect with right ventricular outflow obstruction) and atrioventricular canal are reported (40, 66). Mild left ventricular myocardial abnormalities in young patients with WBS in the absence of clinically significant outflow tract obstruction (67) parallels the mouse model findings (68).

<table>
<thead>
<tr>
<th>Clinical finding</th>
<th>Combined prevalence (%)</th>
<th>Range of prevalence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any CV disease</td>
<td>84</td>
<td>53–100</td>
</tr>
<tr>
<td>SVAS</td>
<td>69</td>
<td>28–100</td>
</tr>
<tr>
<td>Pulmonary arterial stenosis</td>
<td>34</td>
<td>0–83</td>
</tr>
<tr>
<td>Hypertension</td>
<td>17</td>
<td>3–30</td>
</tr>
<tr>
<td>Mitral valve disease</td>
<td>15</td>
<td>4–43</td>
</tr>
<tr>
<td>Coarctation of aorta</td>
<td>4</td>
<td>0–19</td>
</tr>
<tr>
<td>Aortic hypoplasia</td>
<td>2</td>
<td>0–14</td>
</tr>
<tr>
<td>Pulmonary valve disease</td>
<td>5</td>
<td>0–47</td>
</tr>
<tr>
<td>Aortic valve disease</td>
<td>3</td>
<td>0–11</td>
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</tbody>
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Aortogram in a 4-year-old individual with WBS. SVAS (arrowhead) and mild narrowing of the proximal left main coronary artery (arrow) are shown. The gradient measured at catheterization was 40 mmHg.
addition to smooth muscle cells, have an extracellular matrix (Figure 4E) that is rich in mucopolysaccharides and contains endothelium-lined lacunae (74, 75).

**Elastin haploinsufficiency as a cause of CV disease**

*Familial SVAS and WBS.* As introduced earlier, understanding of the molecular basis of CV disease in WBS was aided by studies of a related but genetically distinct disease, familial SVAS (76). The spectrum, natural history, and pathological characteristics of CV and connective tissue lesions in patients with familial SVAS are virtually identical to those found in WBS. However, familial SVAS patients do not have neurobehavioral, metabolic, endocrine, developmental, or some of the craniofacial characteristics of WBS. These differences are due to the fact that familial SVAS is not a microdeletion syndrome but rather is caused by translocations (15), deletions (6, 16), and point mutations (77–79) that disrupt only the *ELN* gene. Despite significant allelic heterogeneity, all *ELN* mutations in familial SVAS studied to date cause loss of function at various levels of elastin biosynthesis, most commonly by eliminating the mutant mRNA via the nonsense-mediated decay pathway (79). Heterozygous loss-of-function mutations in familial SVAS and chromosomal deletion in WBS both cause reduced elastin synthesis and increased proliferation of cultured vascular smooth muscle cells and fibroblasts (73). Thus, both the clinical characteristics of, and the molecular mechanisms underlying, CV disease in WBS and familial SVAS appear to be the same and can be denoted by the term *elastin arteriopathy* (11).

**Animal models.** Animal models provide further evidence for elastin haploinsufficiency as the main cause of CV disease in WBS. Mice homozygous for targeted inactivation of *Eln* die from vascular occlusion (Figure 4M) associated with increased subendothelial smooth muscle cell proliferation in early postnatal life (80). Heterozygous *Eln* knockout mice have hypertension, increased arterial stiffness, and increased number of lamellar units (Figures 4, G and I).
potential genetic modifiers, variants in explain all of the variability in CV disease expression. Among vascular caliber and perfusion, whereas the increased number of vascular compliance in of angiotensin II receptor blockers candesartan and salarasin (68). Hypertension and increased number of lamellae are characteristic of the human and the animal model elastin arteriopathy. However, significantly reduced ELN expression compared with controls even after ing from infantile lethality to no overt or clinically apparent CV involvement. Male sex, a known significant risk factor, is associated with earlier onset and more severe disease (82) but does not ing polymorphisms within ELN pseudogenes, both inactivated by a GT dinucleotide deletion, are located in a telomeric block of low copy number repeats flanking the WBS critical region. In the general population, gene conversion events between the gene and the pseudogenes result in a natural variation of active NCFI gene dose, ranging from 2 to 4 (86). In WBS patients, the deletion breakpoint determines whether the NCFI gene is deleted (14), so that gene dose ranges from 1 to 4. In the WBS patients with more than one copy of NCFI, approximately 56% of the total WBS population, the risk of hypertension is increased 4-fold compared with those with only one functional NCFI allele (18). NCFI encodes the p47phox subunit of the NADPH oxidase, and reduced angiotensin II–mediated oxidative stress in the vasculature is proposed as the mechanism responsible for this protective effect.

Modifying factors
The expression of CV disease in WBS is highly variable, ranging from infantile lethality to no overt or clinically apparent CV involvement. Male sex, a known significant risk factor, is associated with earlier onset and more severe disease (82) but does not explain all of the variability in CV disease expression. Among potential genetic modifiers, variants in ELN, in the size of the WBS critical region, in the “intact” alleles on the normal chromosome 7 homolog, and elsewhere in the genome need to be considered. Given that the developing vasculature is exquisitely sensitive to elastin dose (81), factors that influence elastin biosynthesis, including polymorphisms within ELN, are likely to have important effects on disease severity. Significant differences in elastin synthesis and cross-linking have been observed between different ethnic groups (83) suggesting a genetic basis for natural variation in elastin production. Expression of ELN has been studied in two WBS cohorts. One, enrolling only WBS patients with severe SVAS (73), found significantly reduced ELN expression compared with controls even after accounting for hemizygosity. In contrast, the second study detected normal ELN expression on average, though the authors noted high individual variability (84). Additionally, the second cohort showed a trend for higher ELN expression in patients without SVAS, but the difference did not reach statistical significance. These findings suggest that residual elastin expression protects against the development of CV disease in WBS, but adequately powered and controlled studies are needed to prove this point beyond doubt. Investigation of patients with intracranial aneurysms identified noncoding polymorphisms that affected the expression of ELN, identifying candidate risk alleles for elastin arteriopathy (85).

Reduced expression of a few nondeleted genes mapping to the duplicons flanking the WBS critical region have been documented (84). Variations in the expression of these adjacent genes may be another factor contributing to the phenotypic differences in individuals with WBS.

Among the genes within the WBS critical region, NCFI dose has been associated with the prevalence of hypertension (18). The NCFI gene (Figure 2) and two NCFI pseudogenes, both inactivated by a GT dinucleotide deletion, are located in a telomeric block of low copy number repeats flanking the WBS critical region. In the general population, gene conversion events between the gene and the pseudogenes result in a natural variation of active NCFI gene dose, ranging from 2 to 4 (86). In WBS patients, the deletion breakpoint determines whether the NCFI gene is deleted (14), so that gene dose ranges from 1 to 4. In the WBS patients with more than one copy of NCFI, approximately 56% of the total WBS population, the risk of hypertension is increased 4-fold compared with those with only one functional NCFI allele (18). NCFI encodes the p47phox subunit of the NADPH oxidase, and reduced angiotensin II–mediated oxidative stress in the vasculature is proposed as the mechanism responsible for this protective effect.

Mechanisms of obstructive vascular disease in WBS
Both human and animal studies suggest that elastin is required for the terminal differentiation and quiescence of vascular smooth muscle cells. In elastin-null mice, increased vascular smooth muscle cell proliferation both in vivo and in organ culture occur (80). This hypertrophic phenotype was associated with decreased stress fiber and focal adhesion formation and increased vascular smooth muscle cell migration in vitro and was suppressed by treatment with troponolastin, the soluble precursor protein of elastin (87). Similarly, both dermal fibroblasts and aortic smooth muscle cells isolated from patients with WBS or familial SVAS showed increased proliferation inversely proportional to the amount of elastin produced. Treatment of these cells with insoluble elastin normalized the hyperproliferative phenotype (73).

The formation of segmental obstructive lesions is thought to be a two-step process, consisting of increased number of lamellar units and vessel wall thickening during fetal development, leading to a
uniformly altered vascular tree, followed by postnatal injury-mediated inward remodeling (72). The preferential localization of segmental stenoses to the sinotubular junction and to branch points, areas of high turbulence, supports this notion. Interestingly, Eln−/− mice are protected from vascular remodeling following carotid artery ligation in the ipsilateral vessel while enhanced remodeling occurs in the contralateral artery (88), suggesting that elastin haploinsufficiency may differentially affect ligation-mediated and flow-mediated injury responses. Eln−/− mice show the same CV remodeling response in the renal artery clipping model of adult hypertension as do wild-type animals (89) but are protected from the age-related vessel wall thickening observed in wild-type animals (90). Further clinical studies are needed to determine whether patients with WBS are also protected from age-related vascular changes.

Although it is clear that elastin is a negative regulator of cell proliferation in development, the precise receptors and pathways mediating elastin signaling remain to be identified. Studies in different experimental systems yielded conflicting results. A specific 67-kD elastin-binding protein (EBP) localized to the cell surface has been identified as an enzymatically inactive, alternatively spliced isoform of β-galactosidase (91). EBP has been shown to mediate either increased (92) or decreased (93) cell proliferation depending on whether elastin is present in solution or in solid phase. Elastin peptides were shown to increase the proliferation of coronary artery smooth muscle cells by activating cellular Ca2+ influx through L-type Ca2+ channels and by activating the focal adhesion kinase (FAK), Src, and MAPK pathways (92) (Figure 5A). In contrast, canine coronary vascular cells plated on elastin under cyclic stretch conditions showed reduced serum-induced cell proliferation when compared with cells plated on collagen in an EBP-dependent manner (93). Other studies implicated a GPCR in transmitting elastin signals in elastin-null cells (87). This pathway involves the inhibition of adenylyl cyclase and the activation of RhoA and Rho kinase, leading to increased actin polymerization (Figure 5B).

Several structural proteins of the elastic fibers including fibrillin-1, fibrillin-2, and fibulin-5 have RGD sequences that serve as attachment sites for cellular integrins and could provide indirect elastin-dependent signals in the arterial wall (21) (Figure SC). Interestingly, patients with recessive mutations in fibrillin-5 develop SVAS (94) and fibrillin-5-null mice show exaggerated neointima formation, as well as increased vascular cell proliferation and migration (88), providing in vivo evidence to support a role for fibrillin-5 in elastin signaling. Better understanding of the pathways that connect elastin deficiency to increased vascular cell proliferation may help identify new targets for the treatment of CV disease in WBS.

**Current treatment**

The series from Table 1 describe operative or catheter-based interventions in 18% of patients for left ventricular outflow tract obstruction and in 4% of patients for right ventricular outflow tract obstruction (36–44). Patients with mild SVAS in infancy (peak catheterization gradient <20 mmHg) often remain stable and do not require intervention (36).

Operative techniques for repair of SVAS have utilized patch aortoplasty that may involve augmentation of 1–3 of the aortic sinuses (95, 96). The symmetric inverted 3-sinus patch plasty has resulted in improved outcome in one large series (49). Early mortality for repair of SVAS is 1%–9% (range of median or mean age at operation, 6–16 years), with 20-year survival of 77%–97% and long-term reduction in peak catheterization gradients in the majority of patients (36, 49, 95, 97). Although gradients maybe relieved, hypoplasia may persist in the majority of the aortic arch (98). Diffuse hypoplasia of the aorta is a risk factor for reoperation (36, 49, 99). In patients with severe left main coronary obstruction, patch enlargement of the coronary ostia, excision of a fused aortic leaflet, and bypass grafting have been utilized (100).

Most patients with pulmonary arterial stenosis without significant SVAS can be observed without need for treatment in view of well-documented spontaneous improvement. For patients with persistent systemic or suprasystemic right ventricular pressure, marked asymmetry in pulmonary blood flow, or symptoms, balloon dilatation angioplasty has been used to improve arterial diameter, especially in distal vessels. After catheter-based therapy, right ventricular pressure often remains elevated due to residual proximal obstruction, and the incidence of aneurysms is higher in comparison with non-WBS subjects (101). Patients with biventricular outflow obstruction with an indication for surgical relief of SVAS may undergo balloon angioplasty of pulmonary arterial stenosis prior to surgery. In a series of 33 patients with median age of operation of 4 years for biventricular obstruction, early mortality was 18% (96). Patients with middle aortic syndrome and long segment narrowing may undergo aorto-aortic bypass or patch plasty with bypass grafting of involved renal and visceral arteries (102). Catheter-based therapy is an option for some middle aortic lesions that are more localized (102).

Many patients require treatment for hypertension, but data are not available to recommend drug selection targeted for WBS. Beta blocker and calcium channel blocker drugs have been utilized frequently in several of the retrospective series (7, 40, 61, 103), and the link between infantile hypercalcemia and hypertension (59) suggests a role for calcium channel blockade. Angiotensin receptor blockade is effective in the Eln−/− mouse model. Medical treatment in WBS can be challenging so that multidrug regimens may be required for adequate control of blood pressure. Patients with hypertension resistant to drug therapy should be studied for a renovascular etiology.

**Future therapies**

Surgical treatment of vascular lesions in WBS frequently relies on the use of vascular grafts, most commonly made of artificial materials such as polyethylene terephthalate (Dacron) or expanded polytetrafluoroethylene (ePTFE). The resilience, long-term stability, and cellular attachment properties of elastin make it an attractive material to explore (104). Elastin-like polymers from recombinant tropoelastin or synthetic elastin peptides can be used to coat or functionalize graft materials or used as a scaffold for vascular cells in tissue engineering approaches. Coating of synthetic materials with elastin peptides was shown to decrease thrombogenicity in a variety of settings (105), and insoluble elastin coating significantly decreased in-stent restenosis (87) in a porcine carotid artery model. Tissue-engineered aortic grafts prepared using autologous endothelial and smooth muscle cells seeded onto biodegradable scaffolds and surrounded with intestinal submucosa showed promising long-term potency in animal models, but achieving appropriate amounts of elastin formation continues to be a limitation (106).

An alternative approach to the treatment of CV disease in WBS would involve small molecules or biologicals that can either promote elastin biosynthesis or suppress vascular smooth muscle cell proliferation and migration. Minoxidil, a K+–channel blocker and putative NO agonist (107), glucocorticoids (108), and retinoids...
MMP inhibitors may be beneficial in preventing elastin degradation associated with vascular remodeling (110). A number of pulmonary vasodilatory drugs are efficacious in the treatment of primary pulmonary arterial hypertension, including calcium channel blockers, prostacyclin analogs, endothelin receptor antagonists, and NO agonists (111, 112). These agents could be evaluated as potential modulators of vascular lesions in WBS, even though the pathology of ELN arteriopathy is distinct from primary pulmonary arterial hypertension.

Nutritional intervention or nutraceuticals may theoretically be targets affecting elastin arteriopathy. Copper deficiency (113), β-aminopropionitrile (contained in certain legumes), and drugs such as amine oxidase inhibitors and penicillamine (114) all interfere with the activity of lysyl oxidases, a family of enzymes required for elastin cross-linking (115). On the other hand, dill extract has been shown to increase LOXL1 (lysyl oxidase like 1) gene expression and elastin deposition in vitro (116). Finally, ellagic and tannic acid (polyphenols found in berries and nuts) inhibit proteolytic degradation of elastin and increase elastin deposition in skin fibroblast and organ cultures (117). It remains to be shown whether any of these natural products can reverse the effects of elastin deficiency in vivo.

Better understanding of the mechanisms behind the spontaneous improvement of pulmonary vascular lesions in WBS might lead to new treatment options. Developmental, physiological, mechanical, and biochemical differences between the pulmonary and systemic arterial trees need to be considered as potential mechanisms.

Neonatal adaptation to breathing is known to involve a rapid increase in pulmonary flow, with a more gradual growth of the branch pulmonary arteries. One can speculate that slightly elevated flow velocities are usually gone by 6–12 months of life (118, 119). Neonatal adaptation to breathing may account for some of the resolution of 20 older adults with Williams syndrome. The Journal of Clinical Investigation 6. Ewart, A.K., et al. 1993. Hemizygosity at the elastin gene splice variants can be retained (121, 122).

In summary, the insight that the vascular lesions in WBS are linked to hemizygosity of the ELN gene and the creation of an Eln knockout mouse with vascular pathology has provided a focus for connecting elastin biology with vascular disease. In order to translate experimental insights to treatments, newer animal models that better recapitulate the features of WBS may play an important part. For example a second-generation model with regulatable elastin expression in the vessel wall may resolve important questions such as whether re-expression of elastin, or administration of antiproliferative smooth muscle cell pharma-cotherapy, can reverse the disease process, especially in instances where vascular lesions have already formed. These efforts, along with continued research to elucidate pathophysiology and disease modifiers, will hopefully result in therapies that are alternatives to surgery in that they can ameliorate or even prevent the common complications of WBS arteriopathy.

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(109) have been shown to upregulate elastin production in vivo. Although approximately two dozen genes are deleted in WBS patients, their normal chromosome 7 homolog contains an intact copy of each allele. WBS is a particularly compelling model for this treatment approach, especially if transcriptional control of normal ELN gene splice variants can be retained (121, 122). An alternative approach to in situ gene therapy would involve genetic modification of autologous progenitor (or already differentiated) vascular cells ex vivo followed by re-introduction back into the affected individual, though the challenges of appropriate delivery and integration into sites of vascular lesions present a formidable challenge.
Blood.


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