Induced Chromosome Deletion in a Williams-Beuren Syndrome Mouse Model Causes Cardiovascular Abnormalities

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Abstract

Aims: The Williams-Beuren syndrome (WBS) is a genetic disorder caused by a heterozygous \textasciitilde{}1.5-Mb deletion. The aim of this study was to determine how the genetic changes in a Wbs mouse model alter \textit{Eln} expression, blood pressure, vessel structure, and abdominal aortic wall dynamics in vivo. Methods: Elastin (ELN) transcript levels were quantified by qRT-PCR and blood pressure was measured with a tail cuff system. M-mode ultrasound was used to track pulsatile abdominal aortic wall motion. Aortas were sectioned and stained to determine medial lamellar structure. Results: ELN transcript levels were reduced by 38–41% in Wbs mice lacking one copy of the ELN gene. These mice also had a 10–20% increase in mean blood pressure and significantly reduced circumferential cyclic strain (p < 0.001). Finally, histological sections showed disorganized and fragmented elastin sheets in Wbs mice, but not the characteristic increase in lamellar units seen in \textit{Eln}^{+/−} mice. Conclusions: The deletion of \textit{Eln} in this Wbs mouse model results in lower gene expression, hypertension, reduced cyclic strain, and fragmented elastin sheets. The observation that the number of medial lamellar units is normal in Wbs deletion mice, which is in contrast to \textit{Eln}^{+/−} mice, suggests other genes may be involved in vascular development.

Introduction

The Williams-Beuren syndrome (WBS), a complex genetic disorder with specific cardiovascular, cognitive, and behavioral manifestations, is caused by a heterozygous \textasciitilde{}1.5-MB pair deletion on chromosome 7q11.23. This deletion is caused by the misalignment during meiosis of large region-specific low-copy repeat elements flanking the WBS critical region (WBSCR), leading to nonallelic homologous recombination [1–3]. Affected individuals have difficulty visualizing complete pictures, are commonly diagnosed with attention deficit disorder, and typically have mean full-scale intelligence quotients of 56 [4]. Other clinical characteristics, about which several thorough review articles have been published [5, 6], include distinctive facial features [7], social disinhibition [8], hyperacusis [9], hypertension [10], and supravalvular aortic stenosis (SVAS) or narrowing of large elastic arteries [11]. Histological characterization of arterial vessel
walls of WBS patients with SVAS showed disorganized lamellar structures, fragmented elastic fibers, and hypertrophy of smooth muscle cells [12]. Previous work has indicated that the increased risk of hypertension is likely due to this large arterial vessel remodeling that occurs during vascular development [13, 14]. Some WBS patients even present smooth segmental narrowing of the abdominal aorta with concomitant stenosis at the origins of the renal arteries [15]. These vascular stenoses have been associated with deletions or disruptions within the elastin (ELN) gene [7, 11]. Because of the distinct set of clinical features and relatively small number of affected genes, WBS has become a common disorder for the study of genotype-phenotype correlations in microdeletion syndromes [16].

Knockout (KO) mouse models are often generated to gain insight into gene function. The entire WBSCR is conserved in mice on chromosome band 5G2 in reverse orientation with respect to the centromere [17, 18]. The Francke laboratory recently developed a mouse model that mimics the WBS heterozygous deletion seen in humans [19]. Two mouse strains were generated, each carrying half of the Wbscr deletion. Mice that contain the entire deletion were produced by crossing mice with the two half deletion genotypes. The two deletions in these mice are in trans and, therefore, do not exactly reproduce the contiguous deletion seen in WBS. Yet, as far as gene dosage is concerned, mice with both half deletions are a model for the human WBS deletion with the single exception that the Limk1 gene is inactivated on both chromosomes [19]. According to their location with respect to the centromere of mouse chromosome 5, the two half deletions were named proximal deletion (PD) and distal deletion (DD), while D/P represents the double deletion. While the exact functions of all genes within the WBS region are not known, the gene that encodes the elastin molecule (Eln) in the DD region has been studied extensively [16, 20]. Elastin is synthesized by smooth muscle cells and provides blood vessels with the ability to reversibly extend during the cyclic loading of the cardiac cycle [21]. Initially secreted as a monomer called tropoelastin, it is eventually organized into a polymer that forms concentric rings of elastic lamellae within the medial layer of the vessel wall. Each elastic lamella surrounds a layer of smooth muscle cells, forming a lamellar unit [22], which provide the resilience arteries need to expand during cardiac systole and recoil during diastole. Since the Eln gene falls within the distal section of the Wbs region, DD and D/P mice have only one copy of the ELN gene.

Mice hemizygous for only the ELN gene (Eln\(^{+/−}\)) have previously been created to determine the effect of elastin loss on vascular development [23]. By comparing vessel mechanics with blood pressure in Eln\(^{+/−}\) mice, it has been shown that cardiac and vascular development is physiologically coupled [24]. Eln hemizygosity in both mice and humans induced a compensatory increase in the number of elastic lamellae created during arterial development [23, 25]. In humans, this results in an increased risk of obstructive vascular disease. Further research with Eln\(^{+/−}\) mice found the animals to be stably hypertensive from birth [26] with slightly smaller and stiffer arteries [27] compared to wild-type littermates (WT-Eln). Hypertension and an increase in the number of lamellar units, both characteristics of human elastin arteriopathy, may be developmental adaptations to normalize vascular perfusion and circumferential vessel strain. There have been no reports of segmental stenoses in Eln\(^{+/−}\) mice, an important difference from the human disease, and previous research reports strongly implicate Eln as a causal gene for SVAS and essential hypertension. Thus, we hypothesize that a vascular phenotype similar to Eln\(^{+/−}\) mice will be seen in Wbs mice, leading to observable changes in large elastic arteries.

The purpose of this study was to determine how the genetic changes in this Wbs mouse model alter Eln expression, blood pressure, vessel structure, and abdominal aortic wall dynamics in vivo. The deletion strategy used to create Wbs mice allowed us to gain a greater understanding of which genes likely produce an observed phenotype, as D/P mice could be compared to littermates that are either wild-type (WT-Wbs) or carry one of the half-deletions (DD or PD). Comparing Eln\(^{+/−}\) to Wbs deletion mice also provides insight into the differences between a single gene KO and a multigene chromosomal deletion. Using a quantitative real-time polymerase chain reaction (qRT-PCR) technique to measure Eln transcript levels, a non-invasive tail cuff blood pressure system, a small animal ultrasound system to track aortic wall motion throughout the cardiac cycle, and histological analysis to quantify differences in medial lamellar structure, a correlation was noted between a portion of the Wbscr deletion and reduced Eln levels, increased blood pressure, stiffer blood vessels, and lamellar unit malformation.

**Materials and Methods**

The investigation conforms with the Guide for the Care and Use of Laboratory Animals published by the US National Institutes of Health (NIH publication No. 85-23, revised 1996). All
experiments were conducted in accordance with the Institution’s ethical guidelines and with local Institutional Animal Care and Use Committee approval.

**Animal Models**

Six different groups of mice were used in this study for a total of n = 105, and all mice were bred on a mixed C57BL/6J and 129/Sv background. Wbs mice were predominantly C57BL/6J (~85% C57BL/6J vs. ~15% 129/Sv) [19], while Eln+/− were mostly 129/Sv. Male and female mice were included in both groups. Mice used for the in vivo portions of this study were between 21 and 42 weeks old with the following means ± SD: WT-Wbs = 32.1 ± 6.3; WT-Eln = 24.8 ± 0.2; PD = 31.0 ± 6.3; DD = 31.9 ± 4.6; D/P = 32.0 ± 7.5, and Eln+/− = 24.8 ± 0.2 weeks. Eln+/− and WT-Eln mouse body weights were significantly lower than all others, likely due to the different genetic background (p < 0.05), and PD, DD, and D/P mice weighed less than WT-Wbs littermates: WT-Wbs = 37.3 ± 6.0; WT-Eln = 26.9 ± 2.6; PD = 36.3 ± 6.6; DD = 34.9 ± 5.3; D/P = 33.5 ± 6.6, and Eln+/− = 26.9 ± 4.1 g. Weight reduction was significant for D/P when compared to WT-Wbs (p < 0.05) and agrees with reported growth delays and short stature in this Wbs model [19]. Wbs mice have been kept alive for >2 years, with no instances of early death or unexpected morbidity later in life [unpubl. data]. Life expectancy is also normal for Eln+/− mice [26]. As described previously, tail clipping was performed within 4 weeks of birth to determine the genotype of each mouse using PCR and appropriate primers [19, 20].

**RNA Isolation and qRT-PCR**

To determine whether any of these heterozygous deletions result in a corresponding reduction in Eln transcription, 16 mice (3 WT-Wbs, 4 PD, 4 DD, and 5 D/P) from three different litters were euthanized on postnatal day 7 ([p7] and the visceral organs of the thorax were isolated and homogenized (similar tissue was used in previous research [23]). RNA extraction was performed using TRIzol (Invitrogen) according to the manufacturer’s instructions. Total RNA was quantified with a spectrophotometer, and cDNA from each sample was synthesized according to the manufacturer’s instructions. RT-PCR was performed using SYBR Green Master Mix, consisting of 5 µl SYBR Green Mix (Applied Biosystems). Typically, 25–µl reactions consisting of 5 µl cDNA (1:100), 12.5 µl SYBR Green Master Mix, 2 µl primer (10 nM, forward and reverse), and 5.5 µl water were used. Each sample was analyzed in triplicate in an ABI7500 thermocycler (Applied Biosystems). Threshold cycle values were set manually and analyzed using the comparative method (User Bulletin 2; Applied Biosystems). Amplification of Rps28 transcripts served as RNA control for relative quantification, with primers as described previously [28].

**Blood Pressure Measurement**

We measured systolic, mean, and diastolic blood pressure in conscious mice on three separate instances using a tail cuff system (BP 2000 system; Visitech Systems, Apex, N.C., USA). Mice were held in a black magnetic box on a heated stage. Multiple sessions allowed for each mouse to be trained and improved measurement consistency. At least 10 readings were made for each mouse (n = 7–9 per group).

**Ultrasound**

All mice were imaged in a supine position with the ultrasound transducer placed directly on the abdomen in an anteroposterior projection over the infrarenal aorta (n = 7–10 per group). We removed abdominal hair from each mouse with a depilatory cream. Similar to previous work, M-mode images tracking wall motion were collected at 2–4 discrete locations midway between the renal arteries and iliac bifurcation [29]. The diminutive vessels and rapid heart rates in rodents required a transducer-stabilizing system consisting of a heated platform to maintain body temperature and a transducer mount with an integrated rail system (VisualSonics, Toronto, Ont., Canada). The ultrasound system used (Vevo770; VisualSonics) was specifically designed for the high-resolution needs of small animal imaging with a high transducer frequency (40 MHz, transducer 704). All animals were anesthetized with 1.75–2% isoflurane in 1 l/min of O2 during ultrasound procedures.

**Image Data Analysis and Strain Quantification**

An average of 12 cardiac cycles was calculated from each mouse using the M-mode images. Adobe Photoshop CS (version 8.0; Adobe Systems, San Jose, Calif., USA) was used to measure anterior wall motion (δp), posterior wall motion (δd), systolic diameter (Dp), and diastolic (Dd) diameter. Using these measurements and assuming uniform strain around the vessel, the circumferential component of the Green-Lagrange strain tensor is [30]:

$$F_{sw} = \frac{1}{2} \left( \frac{D_p}{D_d} \right)^2 - 1 = \frac{1}{2} \left( \frac{\delta_p - \delta_d + D_p}{D_d} \right)^2 - 1 \times 100\%$$

(1)

This definition of cyclic circumferential strain assumes the diameter at end diastole is the unloaded reference dimension. It also characterizes deformations independent of rigid body motion and takes into account the nonlinear terms that are ignored when using a small deformation assumption. Aortic circumferential strain for each genotype was estimated and presented as a percentage (x100%).

**Pressure Strain Elastic Modulus and Vessel Stiffness**

As defined previously [31], a vessel’s in vivo pressure strain elastic modulus (or Eps) is defined as:

$$E_p = K \times \frac{P_s - P_d}{(D_s - D_0) / D_0} \times 10^6 Pa$$

(2)

Likewise, a general metric for vessel stiffness (β) is defined as [32]:

$$\beta = \frac{\ln(P_s / P_d)}{(D_s - D_0) / D_0}$$

(3)

In these definitions Ps is the mean systolic pressure, Pd is the mean diastolic pressure (if pressures are measured in mm Hg, K is a constant of 133.3).

**Histology**

Mice were sacrificed with 0.4 mg of tribromoethanol per gram of body mass (n = 3–5 per group). The aortas were exposed and pressure perfused with 10% formalin at 100 mm Hg for 5 min. The ascending thoracic, descending thoracic, and infrarenal abdominal aorta were harvested and imbedded in paraffin. Axial slices 5 µm in thickness were stained with elastic van-Gieson to visualize elastic lamina and Masson’s trichrome for collagen. These ax-

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Fig. 1. Relative Eln transcript levels in p7 mice from each Wbs genotype. The data have been normalized such that the mean of the WT-Wbs group is $1.0 \pm 0.12$. Eln transcripts in DD and D/P mice are significantly reduced compared to combined WT-Wbs and PD levels ($n = 3–5$ per group; * $p < 0.03$). Means $\pm$ SE.

Statistical analysis was performed using ANOVA with a post hoc Tukey-Kramer honestly significant difference comparison between multiple groups (JMP version 7; SAS, Cary, N.C., USA). Values of $p < 0.05$ were considered significant.

Results

Reduced Eln Expression in DD and D/P Mice

The levels of Eln transcript, as determined from the experimental triplicates of each mouse by quantitative RT-PCR, were distinctly different depending on Eln gene dosage. On average, Eln levels in PD mice ($n = 4$) were similar to WT-Wbs ($n = 3$), while Eln levels in DD ($n = 4$) and D/P ($n = 5$) mice showed reductions of 38 and 41%, respectively (fig. 1). This reduction seen in mice with only one copy of the Eln gene was significant for both groups when compared to mice with two copies of the Eln gene (PD and WT-Wbs combined; $p < 0.03$). These data are consistent with previously reported quantitative mRNA levels.

Fig. 2. Systolic (a), mean (b), and diastolic (c) blood pressure in conscious mice from each genotype using a tail cuff measurement system ($n = 7–10$ for each group). DD and Eln$^{+/−}$ have significantly elevated systolic and mean pressures, with DD also showing increased diastolic pressure compared to wild-type littermates (* $p < 0.05$). D/P mice have elevated mean and diastolic pressure, but these increases did not reach significance. Means $\pm$ SD.
Northern analysis, showing a 40–50% reduction of *Eln* expression in *Eln*<sup>+/–</sup> mice [23].

**Increased Blood Pressure in DD and Eln<sup>+/–</sup> Mice**

Blood pressure levels, known to be increased in *Eln*<sup>+/–</sup> mice [26], were measured in conscious mice from all six groups (fig. 2). Both DD and *Eln*<sup>+/–</sup> mice had significantly elevated systolic and mean pressures compared to their wild-type littermates (p < 0.05). D/P mice had slightly elevated mean and diastolic pressure, but these increases did not reach significance. These measurements corresponded to pulse pressures of the following means ± SD:

- WT-Wbs = 36.7 ± 11.2
- WT-Eln = 32.1 ± 10.7
- PD = 30.2 ± 10.8
- DD = 29.1 ± 8.5
- D/P = 26.9 ± 8.7
- *Eln*<sup>+/–</sup> = 34.3 ± 11.4 mm Hg.

**Reduced Wall Motion and Circumferential Cyclic Strain in DD and D/P Mice**

Ultrasound provided information about the motion and strain experienced by the aorta in vivo. Exemplary B-mode and M-mode ultrasound images of the abdominal aorta are shown in figure 3a and b, respectively. Both anterior and posterior wall motion over several cardiac cycles can be seen in the M-mode tracings (fig. 3b). Across
all groups, it is obvious that the amount of anterior wall motion was greater than posterior wall motion. This may be due to differences in external tissue support and agrees well with previously published data using this technique [29]. Figure 3c shows the average anterior and posterior wall motion for all four genotypes. Anterior wall motion was significantly reduced in DD, D/P, and Eln\textsuperscript{+/−} mice compared to wild-type littermates (p < 0.001): anterior (mm) WT-Wbs = 0.059 ± 0.01; WT-Eln = 0.064 ± 0.004; PD = 0.060 ± 0.01; DD = 0.031 ± 0.007; D/P = 0.028 ± 0.009, and Eln\textsuperscript{+/−} = 0.037 ± 0.004. Likewise, posterior motion was also reduced in the DD and D/P mice compared to WT-Wbs, yet this difference was not significant as the overall amount of posterior motion was small: posterior (mm) WT-Wbs = 0.017 ± 0.006; WT-Eln = 0.016 ± 0.006; PD = 0.017 ± 0.006; DD = 0.012 ± 0.005; D/P = 0.014 ± 0.005, and Eln\textsuperscript{+/−} = 0.022 ± 0.006. Interestingly, posterior motion in Eln\textsuperscript{+/−} mice was higher than in all other groups, with a significant increase versus DD and D/P mice (p < 0.05). Finally, circumferential cyclic Green-Lagrange strain, a mechanical characteristic of vessel loading defined in equation 1, was significantly reduced in DD, D/P, and Eln\textsuperscript{+/−} mice compared to wild-type littermates (fig. 3d, p < 0.001): circumferential strain (%) WT-Wbs = 14.9 ± 2.9; WT-Eln = 15.0 ± 1.4; PD = 15.6 ± 3.1; DD = 9.1 ± 1.8; D/P = 9.2 ± 2.3, and Eln\textsuperscript{+/−} = 10.4 ± 0.9.

**Increased Pressure Strain Elastic Modulus and Vessel Stiffness**

Combining ultrasound and pressure measurements allowed for the estimation of the pressure strain elastic modulus (E\textsubscript{p}) and vessel stiffness (β). DD and Eln\textsuperscript{+/−} mice both had significantly increased E\textsubscript{p} values (equation 2) compared to wild-type littermates (fig. 4a; p < 0.05): E\textsubscript{p} (10\textsuperscript{5} Pa) WT-Wbs = 0.362 ± 0.055; WT-Eln = 0.308 ± 0.027; PD = 0.285 ± 0.050; DD = 0.457 ± 0.082; D/P = 0.431 ± 0.119, and Eln\textsuperscript{+/−} = 0.462 ± 0.038. D/P was also increased, but not significantly compared to WT-Wbs. Finally, vessel stiffness (equation 3) values were larger in Eln heterozygous mice (fig. 4b): vessel stiffness WT-Wbs = 3.00 ± 0.46; WT-Eln = 2.62 ± 0.23; PD = 2.37 ± 0.42; DD = 3.22 ± 0.58; D/P = 3.35 ± 0.93, and Eln\textsuperscript{+/−} = 3.29 ± 0.27. These increases were not significant compared to wild-type littermates, but the increases were larger in DD and D/P mice compared to PD mice (p < 0.05).

**Histological Sections Show Malformed ELN Sheets in DD and D/P Mice**

Vascular histology and morphologic examination provided insight into the structure of the aorta. No segmental stenoses were observed in the thoracic or abdominal aorta in any mouse. Representative pressure-perfused sections of the abdominal aortic wall stained with elastic van-Gieson (fig. 5) as well as color images of elastic van-Gieson and Masson’s trichrome staining (online suppl. figure, www.karger.com/doi/10.1159/000316808) showed disorganized and more fragmented elastin sheets in DD and D/P compared to WT-Wbs and PD. Interestingly, the number of lamellar units differed between mice with only one copy of the Eln gene (table 1). Only Eln\textsuperscript{+/−} mice had a significant increase in the number of lamellar units compared to wild-type littermates (ascending tho-
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Discussion

Although the cause of WBS, a recurrent deletion of ~1.5-Mb pairs on chromosome 7q11.23, was originally discovered in 1993 [7], the contributions of specific genes within the deletion to the phenotype have not been clearly defined. A recently developed Wbs mouse model has provided new insight into the study of this neurodevelopmental disorder. It has given researchers the ability to correlate phenotypic changes seen in these mice to regions within the WBS deletion. Others have developed several different animal models of genetic vascular disease, providing intriguing insight into the relationship between gene expression and vascular development. Mouse models, specifically, have underscored the prominent role of the elastic matrix in the morphogenesis and homeostasis of the vessel wall [33]. For example, a fibulin-5 KO mouse model has been used to show that defective organization and stabilization of elastic fibers produces stiffer arteries [34]. Other research has focused on the Eln gene because of its known effects on the cardiovascular system and linkage to SVAS [26, 33]. The Eln+/– mice used in this study were developed over a decade ago and have been extensively analyzed by others [20, 23, 27, 35]. The focus of this study was to determine how the different chromosomal deletions in the new Wbs model affect the cardiovascular system by comparing them to wild-type littermates and to single-gene KO Eln+/– mice.

We found reduced Eln transcript levels, determined by qRT-PCR, in mice that carry DD and D/P, suggesting that the amount of elastin produced during development is reduced. This was expected as the Eln gene lies within the distal portion of the Wbs deletion and is consistent with the reported 40–50% reduction of mRNA in Eln+/– mice [23]. We chose to use p7 mice because Eln expression is vital during this early period when elastic arteries are developing and structurally effective elastin deposition occurs [36]. While Eln mRNA levels may differ slightly between aortic tissue and whole thorax homogenate, we would expect the relative expression, with respect to the WT-Wbs group, to be similar for both. The reduction in Eln transcripts during development was likely a factor in the increased blood pressure levels, reduced in vivo cyclic strain, and more disorganized and frayed elastic sheets in the abdominal aorta for each genotype. Dark elastin sheets in the media are easily seen forming lamellar units. The aortic lumen is on the right of each image, and the adventitial layer, mainly composed of collagen, is on the left. More disorganized and jagged elastin sheets in DD and D/P vessels are obvious compared to WT-Wbs and PD. Eln+/– mice have more MLUs that appear to be thinner and more closely spaced, but are not as fragmented as DD and D/P elastin sheets. Bars = 50 μm.

Table 1. Number of MLUs for all six genotypes as means ± SD (number of animals) in the ascending thoracic, descending thoracic, and infrarenal abdominal aorta

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Ascending thoracic aorta</th>
<th>Descending thoracic aorta</th>
<th>Infrenral abdominal aorta</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT-Wbs</td>
<td>7.3 ± 1.1 (4)</td>
<td>4.9 ± 0.4 (4)</td>
<td>3.3 ± 0.3 (8)</td>
</tr>
<tr>
<td>WT-Eln</td>
<td>7.8 ± 0.6 (4)</td>
<td>5.3 ± 0.1 (4)</td>
<td>3.4 ± 0.1 (4)</td>
</tr>
<tr>
<td>PD</td>
<td>7.3 ± 0.8 (4)</td>
<td>4.9 ± 0.4 (4)</td>
<td>3.2 ± 0.5 (8)</td>
</tr>
<tr>
<td>DD</td>
<td>7.6 ± 0.9 (4)</td>
<td>5.5 ± 0.2 (4)</td>
<td>3.4 ± 0.1 (7)</td>
</tr>
<tr>
<td>D/P</td>
<td>7.2 ± 0.8 (5)</td>
<td>5.4 ± 0.2 (5)</td>
<td>3.4 ± 0.3 (9)</td>
</tr>
<tr>
<td>Eln+/–</td>
<td>10.1 ± 0.5 (8)*</td>
<td>6.6 ± 0.4 (8)*</td>
<td>4.7 ± 0.4 (8)*</td>
</tr>
</tbody>
</table>

Eln+/– mice show an increase in the number of MLUs at all three locations when compared to all other groups (* p < 0.01).
seen in these mice. Finally, while differences in strain backgrounds exist between the Wbs and Eln+/– mice, no differences were seen between WT-Eln and WT-WBS mice, suggesting that any changes are most likely caused by the genetic mutations themselves.

Previous research has indeed shown Eln+/– mice to be hypertensive, with a mean arterial pressure 28% higher than their wild-type counterparts [26]. This increase is already present in newborn mice [35] and persists throughout their lifetime without lowering life-expectancy [26]. The lower blood pressure levels reported here (fig. 2) are likely due to differences in methodology. We chose to use a non-invasive tail cuff system, which measures blood pressure in the tail, while others have used an invasive catheter system that can measure pressure directly in the aorta [35]. The cuff system has also been known to be sensitive to body temperature and positioning along the tail. Although the actual blood pressure measurements were slightly lower in our study, we found a similar 18.3% increase in mean pressure of Eln+/– mice, giving us confidence that these single-gene-KO mice are hypertensive. Interestingly, a similar significant increase in mean blood pressure was only seen in DD mice (20.6%) in comparison to the WT-WBS group. Mean pressure in D/P mice increased by only 10.1%, suggesting a modifying effect of gene(s) within or near the PD region on blood pressure in these adult mice. In this regard, the NCF1 gene in humans, which encodes the p47 subunit of the NADPH oxidase and is sometimes included in the WBS gene in humans, which encodes the p47 subunit of the NADPH oxidase and is sometimes included in the WBS model, has been shown to modify the risk of hypertension. More specifically, hypertension was significantly less prevalent in patients with WBS who had a deletion of Eln [37]. While the presence of NCF1 in WBS patients is variable, the mouse homologue (Ncf1) is outside of both deletions in this Wbs mouse model (it is the next gene beyond the breakpoint Gtf2i in PD mice [19]). Therefore, Ncf1 is unlikely to have an impact on blood pressure in this model. The mechanism to explain the hypertension in DD mice is probably related to the loss of one copy of the Eln gene, but the cause of the non-significant increase seen in D/P mice is likely more complex and will be the topic of future investigation.

In vivo ultrasound measurements provide insight into the motion of the vessel with supporting tissue and under physiologic pressure. Reduced aortic distension is consistent with previously published data for the abdominal aorta in Eln+/– mice when ex vivo compliance was compared at physiologic pressure levels [26, 27] and in vivo in newborn mice [35]. We found significantly decreased anterior and posterior wall motion in DD and D/P mice. Interestingly, Eln+/– mice had decreased anterior motion, but larger posterior motion than any other group (0.022 ± 0.006 μm). One possible explanation for this could be the fact that Eln+/– mice have more tortuous vessels, likely due to the offloading of longitudinal tension [38]. Thus, posterior motion may be increased if support of the abdominal aorta from the spine and back muscle is no longer directly posterior to the vessel.

Green-Lagrange circumferential cyclic strain, a metric that takes into account nonlinear terms, was used because relatively large strain values (8–16%) were studied, making linear diameter strain measurements inaccurate. Our circumferential strain results for the WT-Wbs group (14.9 ± 2.9%) compared well to previously published reports for the normal murine abdominal (15.7 ± 2.5%) [29] and thoracic aorta (15.5 ± 3.3%) [35], giving us confidence that our data are accurate. The reduction in strain was similar for DD, D/P and Eln+/– mice, suggesting that aortic compliance is reduced in mice with less elastin in their large arteries and it is the reduced Eln levels that likely caused these dramatic changes in vascular dynamics. Because the vascular phenotype of PD mice is similar to WT-Wbs, this suggests that no other genes within the proximal region influence in vivo aortic motion.

We estimated aortic compliance by calculating the pressure-strain elastic modulus (E_p) and aortic stiffness (β), both of which combine blood pressure and wall motion data. As expected, mice from DD, D/P, and Eln+/– groups all had larger E_p and β values compared to their wild-type littermates. Compared to clinical data from humans of varying age, our results were similar to male volunteers 5–15 years old [39]. These values increase with age, likely due to a combination of changing wall content (i.e. elastin degradation and increased collagen production) and increased atherosclerotic burden. In our study, this increase is likely due to a combination of hypertension and altered vascular development.

Finally, we saw an interesting difference between Wbs and Eln+/– mice in aortic wall sections stained for elastin and collagen. While the numbers of MLUs in WT-Eln and Eln+/– mice were slightly lower than previously published data [23], our histological images look similar and the percent increase of MLUs in Eln+/– mice was comparable (between 24.5 and 29.5% in the thoracic aorta). Thus, this difference is likely explained by variations in strain background or a slight disparity in what was defined as ascending and descending thoracic aorta. Others have hypothesized that with ELN insufficiency, vascular smooth muscle cells cannot produce enough elastin to withstand the pulsatile circumferential forces, suggesting...
that the increase in MLUs is a developmental adaptation to normalize wall tension [26]. While elastin and collagen fibers did appear more disorganized in DD and D/P mice, with frayed elastin sheets and inhomogeneous collagen distribution, the number of lamellar units was similar for all Wbs mice in the ascending thoracic, descending thoracic, and abdominal aorta. Compared to the DD and D/P mice, we did not observe as much lamellar fragmentation and disorganized elastin in the aortas of Eln$^{+/–}$ mice. These disorganized and dispersed fragmented elastin fibers are similar to the aortic pathology often seen in human cases of WBS [25]. Thus, our results suggest that one or more genes within the DD region may affect vascular development and MLU formation by inhibiting the increase in layers seen in Eln$^{+/–}$ mice. Furthermore, the reduced aortic motion and cyclic circumferential strain cannot be attributed to an increase in lamellar units, as this phenotype was observed in DD, D/P, and Eln$^{+/–}$ mice. More likely, altered vessel dynamics can be attributed to an increased loading of collagen fibers in any vascular tissue with altered elastin structure. All of the above findings are summarized in table 2, with a comparison to SVAS and WBS patients.

Clinical reports have confirmed with intravascular ultrasound that WBS patients have diffusely narrowed and thick-walled stiff arteries, often lacking elastin [40]. These pathologies are hypothesized to be caused by smooth muscle hypertrophy in the medial layer, with abnormal elastic fibers and mild collagenization, as seen in several clinical cases [41–43]. These pathologic changes progress with age in humans, often reducing life expectancy. In animals, Eln expression and organization has been shown to alter the response of vessels to aging in Eln$^{+/–}$ mice [44]. Other clinical case reports have shown no difference in arterial lesions when patients with SVAS and WBS were compared (n = 2 for both) [25], describing elastic fibers as thin, fragmented, and numerous in both diseases. While interesting, further work quantifying elastin structures in a larger number of WBS patients would help improve our understanding of the vascular malformations associated with this disease.

Due to the observed changes in the medial layer, innervation of the abdominal aorta or response to vasoactive hormones may also differ in Wbs mice. First, it has long been known that several different types of neurons innervate all three layers of the abdominal aorta, with abundant nerve endings in the adventitial layer. For example, in humans, cholinergic nerve fibers, known to be parasympathetic, are organized in two plexuses: a superficial layer localized in the adventitia and a deeper layer localized in the adventitial-medial transitional zone. In mice, cholinergic nerve fibers are organized in a single plexus in the adventitia [45]. Other receptor trunks make a stepwise course as they penetrate through the layered elastin sheets in the media [46]. Since vascular elastin does not form typical layered lamellar units in DD and D/P mice, neuronal endings in this layer would likely have a different morphology, possibly affecting their function. Neural control mechanisms could also play a role in the development and maintenance of vascular elastic tissues. Future work will be needed to determine where these innervating structures are located and what role they play in the altered phenotypes observed in Wbs mice. Secondly, previous work has shown that the mouse aorta contracts when exposed to agonists of adrenergic, serotonergic, dopaminergic, and prostaglandin receptors [47]. Thus, future research could determine the ex vivo aortic response of Wbs mice to common vasoactive agonists and antagonists.

<table>
<thead>
<tr>
<th>Model</th>
<th>Clinical Reports</th>
<th>Intravascular Ultrasound</th>
<th>Elastin and Collagen Pathology</th>
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</table>
In summary, mice that carry a deletion in the *Eln* gene (DD, D/P, and *Eln*+/− mice) have a less compliant aorta, likely due to reduced *Eln* transcription and morphological changes in wall structure. We did not observe an increase in the number of elastic lamellae within the medial layer in Wbs mice carrying one copy of the *Eln* gene, a characteristic that is seen in *Eln*+/− mice [23]. This difference may be due to one or more genes within the DD region affecting vascular development and lamellar formation, or uncharacterized variations between inbred strain backgrounds. These data provide further insight into the genetic relationship between *Eln* levels, aortic stiffness, and the cardiovascular abnormalities associated with WBS.

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