Microcalcification and Thoracic Aortopathy: A Window Into Disease Severity

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BACKGROUND: Patients with thoracic aortopathy are at increased risk of catastrophic aortic dissection, carrying with it substantial mortality and morbidity. Although granular medial calcinosis (medial microcalcification) has been associated with thoracic aortopathy, its relationship to disease severity has yet to be established.

METHODS: One hundred one thoracic aortic specimens were collected from 57 patients with thoracic aortopathy and 18 control subjects. Standardized histopathologic scores, immunohistochemistry, and nanoindentation (tissue elastic modulus) were compared with the extent of microcalcification on von Kossa histology and 18F-sodium fluoride autoradiography.

RESULTS: Microcalcification content was higher in thoracic aortopathy samples with mild (n=28; 6.17 [2.71–10.39]; P≤0.00010) or moderate histopathologic degeneration (n=30; 3.74 [0.87–11.80]; P<0.042) compared with control samples (n=18; 0.79 [0.36–1.90]). Alkaline phosphatase (n=26; P=0.0019) and OPN (osteopontin; n=26; P=0.0045) staining were increased in tissue with early aortopathy. Increasingly severe histopathologic degeneration was related to reduced microcalcification (n=82; Spearman ρ, −0.51; P<0.0001)—a process closely linked with elastin loss (n=82; Spearman ρ, −0.43; P<0.0001) and lower tissue elastic modulus (n=28; Spearman ρ, 0.43; P=0.026).18F-sodium fluoride autoradiography demonstrated good correlation with histologically quantified microcalcification (n=66; r=0.76; P<0.001) and identified areas of focal weakness in vivo.

CONCLUSIONS: Medial microcalcification is a marker of aortopathy, although progression to severe aortopathy is associated with loss of both elastin fibers and microcalcification.18F-sodium fluoride positron emission tomography quantifies medial microcalcification and is a feasible noninvasive imaging modality for identifying aortic wall disruption with major translational promise.

GRAPHIC ABSTRACT: A graphic abstract is available for this article.

Key Words: aneurysm, dissecting ▪ aortic aneurysm ▪ aorta, thoracic ▪ calcinosis ▪ elastin ▪ sodium fluoride ▪ vascular calcification
identify patients with thoracic aortic aneurysms that are vulnerable to these potentially fatal complications.

Aortic wall microcalcification, or granular medial calcification, is an underappreciated disease process that represents the deposition of hydroxyapatite crystals formed of calcium phosphate within the extracellular matrix. Microcalcification has been described in thoracic aneurysms of various etiologies and is specifically associated with elastin fragmentation, vascular smooth muscle cell phenotypic switching, and increased aortic wall rupture risk on biomechanical testing. We have previously shown that abdominal aortic aneurysm microcalcification can be detected using 18F-sodium fluoride positron emission tomography, and its uptake correlates with aneurysm expansion rate, as well as the risk of subsequent complications. A better understanding of the relationship between thoracic aortic aneurysm microcalcification, 18F-sodium fluoride positron emission tomography, pathological mechanisms, and disease severity has the potential to uncover novel therapeutic targets and act as a clinical tool to diagnose and to monitor thoracic aortopathy.

Using histology and 18F-sodium fluoride autoradiography, we here aimed to assess the relationship between aortic wall microcalcification and thoracic aortopathy and aneurysm disease. We sought to investigate the pathways associated with aneurysm microcalcification and assess the relationship between microcalcification, tissue biomechanical properties, and histological disease severity.

METHODS

Human Ascending Aortic Tissue

Aneurysmal thoracic aortic tissue was obtained from patients with bicuspid (n=28) or tricuspid (n=20) aortic valves undergoing elective repair of aneurysm meeting the American Heart Association or European Society of Cardiology guidelines. Aortic tissue was also retained from the flap and true lumen of patients undergoing surgery for acute or chronic aortic dissection (n=13). Control aortic tissue samples were obtained from patients undergoing proximal vein graft anastomosis during coronary artery bypass grafting (n=9) or donor heart transplant aortic trimmings (n=13). Specimens were obtained from biobanks at 3 separate institutions: the Royal Infirmary of Edinburgh, United Kingdom; the Cardiovascular Tissue Registry at the University of British Columbia and St. Paul’s Hospital Centre for Heart Lung Innovation, Vancouver, Canada; the University of Liverpool, United Kingdom (for overview of methodology, see Figure 1). Samples at each institution were collected from consenting patients or relatives and in accordance with ethical approvals (15/ES/0094, 18/SS/0136, 14/NW/1212, 19-09 Liverpool Bio-Innovation Hub project approval and H21-00987–Providence Health Care/UBC Research Ethics Board). Study data that support the findings of this study are available from the corresponding author upon reasonable request.

Histological Assessment

Histology was performed by 2 institutions (NHS Lothian Clinical Pathology Laboratory and University British Columbia) according to local protocols. One to 4 paraffin-embedded sections were sliced at 5 to 6 µm per sample and slide mounted. Samples containing macrocalcified atheroma requiring decalcification or poor-quality specimens (freezing or imaging artifact) precluding histological grading were excluded. Histological staining included von Kossa, hematoxylin and eosin, elastin van Gieson, and Movat pentachrome performed according to local protocols. One to 4 paraffin-embedded sections were sliced at 5 to 6 µm per sample and slide mounted. Samples containing macrocalcified atheroma requiring decalcification or poor-quality specimens (freezing or imaging artifact) precluding histological grading were excluded. Histological staining included von Kossa, hematoxylin and eosin, elastin van Gieson, and Movat pentachrome performed according to local protocols. All images were captured using the Axioscan slide scanner (Zeiss, Germany) and analyzed using the FIJI software (v 2.0.0, open source) or Aperio Slide Scanner using the ImageScope software (Leica Biosystems, Germany).
Histopathologic severity scoring was performed by an experienced pathologist (WW) blinded to the results of microcalcification quantification and in accordance with international consensus guidelines. Briefly, aortic histology was split into 5 major categories; intimal mucoid extracellular matrix accumulation, transmural mucoid extracellular matrix accumulation, elastin fiber fragmentation or loss, smooth muscle cell nuclei loss, and laminar medial collapse. Each category is
split into absent, mild, moderate, or severe, scoring 0, 1, 2, or 3 points, respectively. The final score represents a sum of each category with possible scores ranging from 0 to 15. The presence or absence of atheroma was also recorded.

Scanning Electron Microscopy and Dispersive Radiograph Energy
Paraffin sections were deparaffinized using Xylene followed by ethanol 100%. Slides were then completely dried out before coating them with carbon. Scanning electron microscopy with secondary electron was used to visualize the tissue. Dispersive radiograph energy coupled with the electron beam of the scanning electron microscopy was used as a chemical analysis method to determine the elemental composition of the mineral deposition on the tissue.

Immunohistochemistry
Immunohistochemistry was performed on 26 samples from 13 patients with aortopathy and 6 control subjects. Scoring was completed from high-resolution slide scans. Immunohistochemistry was completed using OPN (osteopontin; catalog No. 07264; 1:100 dilution; Sigma-Aldrich), alkaline phosphate (AP; ab95462 1:300; Abcam), BMP2 (bone morphogenetic protein 2; AHP960 1:100; Biorad), WNT3A (wingless-type MMTV integration site family member 3a; ab28472; Abcam), cleaved caspase 3 (No. 9664; 1:100 dilution; Cell Signaling), and p-ERK (phospho-extra-cellular signal-related kinase; 9102S, 1:50 dilution; Cell Signaling). Staining was performed via automated staining with a Leica Bond Rx system using Bond Epitope Retrieval Solution 1 (pH=6, catalog No. AR9961) and Bond Polymer Refine Red Detection (catalog No. DS9390). Background staining was assessed via assessment of sections processed with the omission of primary antibody and the consideration of normal human aortic background from normal transplant donors. The omission of the primary antibody served as negative controls. Blinded qualitative categorization of staining was performed by a pathologist (S.S.) using a 4-point classification based on relevant cell positivity as described previously in target tissue:14: 0, no notable staining; 1, 5% to 20% of relevant cells are weakly positive; 2, 21% to 50% of relevant cells are positive; 3, 51% to 100% (see Figure S1 for representative staining of OPN, AP, and cleaved caspase III as examples).

Immunofluorescence
To assess the RUNX2 (Runt-related transcription factor 2) expression, we used immunofluorescence on paraffin sections. First, sections were treated with xylene twice 10 minutes and then rehydrated twice in 10 minutes using serial baths of 100% followed by 5-minute baths in 95%, 70%, and 50% each. Slides were then rinsed with water for 10 minutes. Citrate was used to retrieve antigen for 30 minutes. Sections were blocked using goat serum for 1 hour. Primary antibody was diluted 50-fold in dilution buffer; slides were then incubated overnight at 4 ºC. Slides were washed 3x in PBS 1x and then incubated with Alexa red secondary antibody for 1 hour. Sections were then washed with PBS 1x or 5 minutes each. After that, sections were mounted with DAPI (4’, 6-diamidino-2-phenylindole) and imaged using epifluorescence microscope. Image Quantifications: NIH Image J software was used to quantify the mean fluorescence of RUNX2.

Histological Quantification
As multiple samples from the same patient were scored in some instances, the sample with the highest pathology score was used for patient-level analysis, in line with international consensus. Microcalcification quantification was performed by a trained user (A.J.F.) blinded to the histological scoring, immunohistochemistry, and nanoindentation. von Kossa images were transformed to 8-bit gray scale, a threshold value set to 50% of total opacity of the media, and altered by no more than 10% to produce optimal visual coverage of microcalcification. A region of interest was drawn around the intima and media, and the microcalcification concentration reported as the percentage area over threshold values. The percentage area of microcalcification was subdivided into minimal (0%–0.99%), mild (1%–4.99%), moderate (5%–9.99%), or severe (≥10%) categories (for examples, see Figure S2). The intraobserver variability of microcalcification percentage area calculations was determined in 24 randomly selected samples. Analyses were performed twice in random order by a single trained user (A.J.F.) at least 2 months apart to minimize recall bias and blind to the original analysis. To assess the distribution of microcalcification, regions of interest were drawn around the intima, inner media (luminal 50%), and outer media (adventitial 50%). To allow comparison across samples with varying overall microcalcification content, a ratio between each area and the total microcalcification uptake of each sample was reported.

Elastin concentration was determined using a similar method. The elastin van Gieson image was transformed to gray scale 8-bit, with the threshold set using the Otsu formula and altered visually within 20% of the Otsu value to achieve optimal elastin opacity. Wall thickness was measured from the intima to the external elastic lamina. Smooth muscle cell density was measured on the hematoxylin and eosin–stained tissue, and nuclei cell density was calculated as the number of cells in a region of interest divided by the area. In a subset of samples with clear areas of high and low microcalcification, mean wall thickness, elastin concentration, and nuclei cell density were calculated using 3 full thickness regions of interest in both high and low microcalcification concentration areas. Samples with absent microcalcification content, or homogenous distribution (no distinct areas of high or low microcalcification), were excluded from this subanalysis.

Nanoindentation
Oscillatory nanoindentation allows the assessment the elastic and viscous properties of localized regions of the tissue by measuring the shear storage ($G''$) and shear loss modulus ($G'$), respectively. $G''$ is directly related to the tissue elastic modulus ($E$). Small indentations of 0.5 µm at a frequency of 110 Hz were performed using a 100-µm-flat punch indenter tip (Sytont-MDP, Ltd, Nidau, Switzerland) with a G200 nanodenter equipped with a DCM-II actuator (KLA-Tencor, Milpitas) by an experienced investigator (R.A.) as described previously.

Biochemical Assessment
Elastin, collagen, and glycosaminoglycan content were assessed using established protocols as described previously. Briefly, aortic tissue was digested with papain (for
collagen and glycosaminoglycan analysis) or oxalic acid (for elastin analysis). Following digestion, collagen content was determined by measuring hydroxyproline concentration in the tissue using the 1,3-dimethylbutylamine dye, glycosaminoglycan content was measured using the dimethyl methylene blue assay, and elastin was measured using Fastin Elastin Kit (Biocolor, County Antrim, United Kingdom).

### Ex Vivo 18F-Sodium Fluoride Autoradiography

Formalin-fixed paraffin-processed sections were rehydrated and equilibrated in PBS for 30 minutes. Sections were then incubated with 100 kBq/mL of 18F-sodium fluoride in PBS for 1 hour at room temperature with a blocking control (10 μmol/L sodium fluoride) before two 5-minute washes in PBS and one in deionized water. Dried sections were exposed to a high-resolution autoradiography plate (BAS-IP-SR 2040; Cytiva), which were imaged on an autoradiography imager (Amersham Typhoon IP Biomolecular Imager, Cytiva).

Sample 18F-sodium fluoride content was quantified using the FIJI software (v 2.0.0, open source). Gray scale images were imported and a region of interest drawn around the perimeter of the image to provide mean background activity. Regions of interest were drawn around individual samples providing a mean gray intensity. The result was adjusted by dividing by the background activity, standardizing measurements, and allowing samples across separate experiments to be compared.

### In Vivo 18F-Sodium Fluoride Positron Emission Tomography Example

We include an example 18F-sodium fluoride positron emission tomography scan (Biograph mCT; Siemens Healthcare, Germany) fused with a magnetic resonance imaging angiogram (Biograph mMR; Siemens Healthcare) taken from an ongoing clinical trial; the assessment of risk in thoracic aortic disease using 18F-sodium fluoride study (https://www.clinicaltrials.gov; unique identifier: NCT04083118).

### Statistical Analysis

All statistical analyses were performed in RStudio (V1.3.959, general common license). Categorical variables were presented as number (percentage). Continuous variables with normal distribution are presented as mean±SD, whereas non-normally distributed variables were presented as median (interquartile interval). Analyses of variable influence on patient-level specimen microcalcification content were performed using a univariable linear regression. When comparing between groups, both overall (Kruskal-Wallis) and individual (Wilcoxon) nonparametric tests were used. For correlations between numeric variables, Spearman rank-sum test was used. For comparisons between groups, paired Wilcoxon tests were used. A 2-sided P<0.05 was considered statistically significant.

### RESULTS

#### Study Population

Specimens from 8 participants were excluded from analysis (5 macrocalcified atheromas, 2 freezing or imaging artifacts, and 1 aortitis), leaving specimens from 75 participants (57 patients with thoracic aortopathy and 18 control subjects) for assessment. Participants with bicuspid aortic valve aneurysms tended to be younger although the prevalence of valvular heart disease, cardiovascular risk factors, and maximal aortic diameters were similar between the 3 aortopathy subgroups (Table). Those with dissection had more severe histopathologic degeneration scores than those with bicuspid or tricuspid valve thoracic aortic aneurysms ($\chi^2 P=0.001$; Table S1).

### Aortic Microcalcification

The microcalcification quantification method demonstrated excellent intraobserver repeatability, with a mean absolute percentage area difference of 0.44%, a coefficient of repeatability of 4.5%, and an interclass correlation coefficient of 0.97 (0.93–0.99; Figure S3). When considered as a categorical variable, the repeatability was similarly excellent with 20 of 24 (83%) given the same grade (interclass correlation coefficient, 0.9 [0.78–0.95]; Table S2).

Overall, 43 (57%) of the 75 specimens had at least mild medial microcalcification on von Kossa staining, with only 1 (1.3%) specimen having intimal microcalcification associated with complex atheroma. Microcalcification was clearly localized to the media, with particularly intense microcalcification seen in the outer media (outer media/total sample microcalcification content ratio, 1.36 [1.03–1.54]), but there was very little staining in the intima (intima/total sample microcalcification content ratio, 0.09 [0.04–0.30]; Figure 2).

In 8 specimens with varying degrees of aortic histopathologic disease severity (2 control, 1 mild, 3 moderate, and 2 severe), scanning electron microscopy with dispersive radiograph energy confirmed that the microcalcification was crystalized calcium phosphate. Control samples appeared to have little microcalcification, while those with mild or moderate aortopathy had deposition of microcalcification along intact elastin (Figures S4 and S5). Those with severe histopathologic aortopathy had large areas of mucoid extracellular matrix accumulation devoid of elastin or microcalcification (Figure S5).

Autoradiography with 18F-sodium fluoride colocalized precisely with microcalcification on von Kossa staining, demonstrating a good correlation (Spearman $\rho$, 0.76; P<0.0001; Figure 3). Furthermore, to demonstrate the feasibility of using 18F-sodium fluoride to assess aortopathy in vivo, we present the case of a 40-year-old with bicuspid aortic valve and aortopathy with focal aortic wall outpouchings who underwent combined 18F-sodium fluoride positron emission tomography and magnetic resonance imaging angiography (Figure 3). There was high overall uptake in the ascending aorta, with reduced uptake in the areas of focal expansion.
Rise and Fall of Medial Microcalcification With Progressive Aortopathy

In 82 thoracic aortopathy specimens, aortic pathology score was categorized into mild (pathology score, 2–4), moderate (pathology score, 5–7), and severe (pathology score, ≥8) disease (thresholds determined using pathology score tertiles). Compared with control specimens (0.79 [0.36–1.90]), microcalcification content was higher in those with mild (6.17 [2.71–10.39], Wilcoxon with Bonferroni correction $P=0.00010$) or moderate (3.74 [0.87–11.80], Wilcoxon with Bonferroni correction $P<0.042$) aortic pathology scores but was similar in those with severe aortic pathology score (0.40 [0.15–0.87], Wilcoxon with Bonferroni correction, $P=0.42$; Figure 4). In samples from patients with aortopathy, there was an inverse correlation between microcalcification and the overall pathology score (Spearman $r$, $-0.51$; $P<0.0001$; Figure 4).

To understand the processes driving the relationship with pathology scores, we assessed microcalcification content against individual pathological subcategories. Microcalcification content was inversely associated with the severity of elastic fiber fragmentation and loss (Spearman $r$, $-0.43$; $P<0.0001$; Figure 5), mucoid extracellular matrix accumulation (Spearman $r$, $-0.41$; $P=0.0002$), and medial collapse (Spearman $r$, $-0.23$; $P=0.038$) but not cell loss (Spearman $r$, $-0.17$; $P=0.14$). In those with severe disease, any remaining microcalcification colocalized exclusively with areas of residual elastin (Figure 5).

To investigate the mechanisms associated with microcalcification precipitation in early disease, expression of markers for osteogenic smooth muscle cell phenotyping (BMP2, WNT3a, OPN, and RUNX2), pyrophosphate degradation (AP), apoptosis (caspase III), and noncanonical transforming growth factor-$eta$ pathways (p-ERK) were assessed, comparing control (n=6) with aortopathy samples in the early phase of disease (≤mild aortic pathological severity score and ≤mild microcalcification, n=21). Both AP (Wilcoxon $P=0.0019$) and OPN (Wilcoxon $P=0.0045$) were increased in early-phase aortopathic disease. Further, the areas of high positivity colocalized with areas of high microcalcification (Figure S7). In contrast, positivity of WNT3a (Wilcoxon $P=0.51$), RUNX2 (Wilcoxon $P=0.075$), cleaved caspase III (Wilcoxon $P=0.051$), BMP2 (Wilcoxon $P=0.21$), and p-ERK (Wilcoxon $P=0.12$) was similar to control specimens.

To assess the relationship between microcalcification and tissue elastic modulus, both nanoindentation and microcalcification content assessments were performed on sections from the same sample. There was a positive correlation between the elastic modulus and both microcalcification (Spearman $\rho$, 0.43; $P=0.026$) and elastin...
DISCUSSION

Medial microcalcification is an overlooked pathological process related to thoracic aortic aneurysm disease, with the underlying mechanisms and its relationship to disease severity poorly understood. In the current study, we demonstrate a striking association between mild and moderate thoracic aortic aneurysm disease and microcalcification, which is linked to AP and OPN and an increase in tissue elastic modulus. Intriguingly, thoracic aortopathy samples with severe histopathologic degeneration consistently had low levels of microcalcification, a process associated with elastin fiber loss, suggesting a nonlinear pathobiological course of microcalcification in thoracic aortopathy. Finally, we found that uptake of the radiotracer $^{18}$F-sodium fluoride colocalized exquisitely with histological microcalcification, thereby providing support for its use as a noninvasive imaging biomarker of aortopathy severity in similar populations.

We found that patients with mild or moderate thoracic aortic aneurysm disease have over 5× more medial microcalcification content than control subjects, which is in line with previously published reports. The processes mediating the initial precipitation of microcalcification on elastin in thoracic aortic aneurysms are incompletely understood and largely noninflammatory, unlike atherosclerotic aortic aneurysm formation or aortitis. Our experiments confirm hydroxypatite deposition along intact elastin, rather than

(Spearman $\rho$, 0.52; $P=0.0041$) content but not collagen ($P=0.43$) or glycosaminoglycan ($P=0.32$) content.

Figure 2. Distribution of microcalcification in the proximal aortic wall of aneurysms. Assessment of the pattern of microcalcification across the aortic wall. A, Representative image of aortic wall stained with von Kossa, demonstrating significant microcalcification in the media, particularly the outer media, but not the intima. B, Across all samples assessed, as a ratio of total sample uptake, there was relatively little microcalcification of the intima, whereas there was significant deposition in the media, particularly the outer media. C through E, This pattern was consistent across aneurysm and control samples, with bicuspid aortic valve demonstrating a particularly strong preponderance for microcalcification in the outer media.
at fragmentation points, although molecular damage at these sites cannot be excluded. Further, in both within-sample analysis and biochemical analysis, we find that higher elastin content is associated with higher microcalcification content, consistent with reports from a Matrix Gla protein arterial calcification mouse model, where a strong correlation between elastin and microcalcification content was observed.19

We explored potential mechanisms driving microcalcification precipitation in thoracic aneurysm disease focusing on aortopathy samples with mild calcification and mild degeneration where elastin is still intact. By selecting the early stages of disease, we have focused our immunohistochemistry experiments on processes occurring before, but not caused by, the known propagating effects of microcalcification on further calcification.9,20 In these early aortopathy disease samples, AP was increased and is consistent with previous studies using FBN1C1041G/+ murine models of the Marfan syndrome.9 Furthermore, we find higher OPN, which is associated with tissue damage and vascular smooth muscle cell transdifferentiation in vascular disease21—a result consistent with previous reports in bicuspid aortic valve aortopathy.17 Overall, while our results find associations between markers of smooth muscle cell transdifferentiation and AP and early medial microcalcification in aortopathy, further research is required to elucidate the specific pathways involved. For example, Krüppel-like factor 4—an upstream mediator of

Figure 3. 18F-sodium fluoride is an excellent marker of medial microcalcification.
A through D, Visual comparison of samples with minimal (A), mild (B), moderate (C), and severe (D; scale threshold applies to all autoradiography images). There was a good correlation between histologically assessed von Kossa, and 18F-sodium fluoride determined microcalcification in a sample-level analysis (E). In vivo 18F-sodium fluoride positron emission tomography fused with magnetic resonance angiogram in a patient with bicuspid aortic valve and saccular aneurysm of the ascending aorta seen on 3D reconstruction (F). Fused angiogram and 18F-sodium fluoride (G–I). Note reduced 18F-sodium fluoride at sites of aortic wall expansion (green arrows). TBR indicates tissue to background ratio.
Figure 4. The relationship between aortopathy severity and microcalcification. 

A, Patients with mild (Wilcoxon with Bonferroni correction, $P=0.00010$) or moderate (Wilcoxon with Bonferroni correction, $P=0.042$) histopathologic aortopathy severity had more microcalcification than control samples. Those with severe disease had less microcalcification content than either mild (Wilcoxon with Bonferroni correction, $P<0.0001$) or moderate (Wilcoxon with Bonferroni correction, $P=0.0030$).

B, In aortopathy samples, there was an inverse relationship between histopathologic disease severity and microcalcification content (Spearman $\rho$, $-0.51$; $P<0.0001$).

C through K, Representative examples of hematoxylin and eosin (C, F, and I), elastin van Gieson (D, G, and J), and von Kossa (E, H, and K) in mild, moderate, and severe histopathologic disease, with minimal microcalcification coinciding with near total elastin loss in severe disease.
OPN expression in vascular smooth muscle cell osteogenic transdifferentiation—has not been explored in the current work but has been reported in distinct smooth muscle cell subtypes in Marfan syndrome and represents an important research focus for exploring early microcalcification in noninflammatory thoracic aortopathy. Thus, greater characterization is needed to gain a more complete understanding of these processes.

Figure 5. Elastin fiber loss is associated with reduced microcalcification in severe aortopathy. A, Association between microcalcification content and elastin fragmentation/loss subcategory (Spearman $\rho = -0.43; P < 0.0001$). B and C, Scanning electron microscopy from a sample with moderate histopathologic severity (B) with clear microcalcification precipitation on intact elastin fibers (orange arrows) and severe disease (C) demonstrating similar deposition along intact elastin but areas devoid of elastin with no microcalcification. D through G, Representative von Kossa (E and F) and elastin van Gieson (G and H) image from a patient with bicuspid aortic valve, severe elastin fiber loss, and microcalcification content of 0.4% percentage area (minimal). The microcalcification is clearly colocalized with areas of remaining intact elastin fibers (yellow arrows), whereas there is no microcalcification in areas devoid of elastin fibers (red arrows).
While higher microcalcification was associated with mild and moderate histopathologic thoracic aortic aneurysm severity, we were surprised to find that severe pathological disease was associated with low levels of microcalcification, suggesting a nonlinear pathological course. Through biochemical and both within and between sample histological quantification, we find that this decline in microcalcification is associated with loss of elastin fibers on which the microcalcification has precipitated. While elastin fragmentation is known to have an augmenting effect on microcalcification,25,26 total elastin destruction or loss seen in end-stage disease appears to have the opposite effect, which will need to be accounted for when assessing histological severity by microcalcification. In line with this, we find a linear relationship between both microcalcification content and elastin content with tissue elastic modulus, suggesting biomechanical weakness in those with low microcalcification and elastin content. Supporting this, multiple previous reports have found a close relationship between intact elastin and delamination strength.25,26 If microcalcification could be serially visualized, it has the potential to provide crucial information about the reduction of elastin integrity over time and therefore dissection or rupture potential of thoracic aortic aneurysms.

We found that the radiotracer 18F-sodium fluoride, which binds specifically to hydroxyapatite (calcium phosphate crystals),14,27 had excellent visual colocalization with histological specimens stained with von Kossa. We present the case of a patient with significant aortopathy and focal wall weakness culminating in aortic wall outpouching (Graphic Abstract). While the overall aneurysm 18F-sodium fluoride uptake is high, the areas of weakness have low signal, suggesting focal elastin decay and high biomechanical failure potential. These findings provide proof of concept for 18F-sodium fluoride positron emission tomography imaging in detecting the vulnerable aortic wall in aortopathy. The potential applications of such imaging for those who would most benefit from surgery are 2-fold. First, thoracic aortic aneurysm microcalcification is indicative of mild or moderate disease where the elastin is still largely intact and may represent a lower risk stage in the disease that could be initially managed conservatively and monitored over time. Second, a patient with focal areas of low 18F-sodium fluoride uptake, as in the case presented, or one with decreasing thoracic aneurysm microcalcification over time, would signify advanced disease with poor elastin integrity that might prompt earlier intervention.

It is important to outline some limitations of the current work. All samples were obtained from tissue biobanks, and demographic information and data on comorbidities were incomplete. Specifically, control thoracic aortic tissue was obtained from heart donors and is completely deidentified and anonymized precluding any comparison of demographics to the disease groups. However, tissue from beating heart donors represents the gold standard for control thoracic aortic tissue and represents the most robust control tissue available. While our results demonstrate an important relationship between microcalcification and histopathologic severity in thoracic aortopathy, histological specimens represent a single snapshot biased toward more severe disease, and our results cannot determine the time course of disease progression. Progress in this regard will come from sequential 18F-sodium fluoride imaging of patients with thoracic aortopathy over time, which is currently being explored (NCT04083118).

In conclusion, we found that aortopathy is associated with increased medial microcalcification, AP, and OPN. Microcalcification precipitates on elastin fibers, peaking in mild and moderate thoracic aortic aneurysm disease but falling in severe disease in line with the decay and loss of elastin fibers. The fall in elastin is associated with reduced stiffness and represents an area of biomechanical weakness. Finally, we show that the radiotracer 18F-sodium fluoride colocalizes with medial microcalcification, paving the way for noninvasive assessment of thoracic aortic aneurysm disease severity, with major implications for selecting ideal candidates for elective repair.
