

Mechanical weakness of thoracic aorta related to aging or dissection predicted by speed-of-sound with collagenase

メタデータ	言語: English 出版者: 公開日: 2022-01-26 キーワード (Ja): キーワード (En): 作成者: Miura, Katsutoshi, Yamashita, Kanna メールアドレス: 所属:
URL	http://hdl.handle.net/10271/00003939

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1 **Mechanical weakness of thoracic aorta related to aging or dissection predicted by**
2 **speed-of-sound with collagenase**

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14 **Abstract**

15 **Scanning acoustic microscopy reveals information about histology and**
16 **speed-of-sound (SOS) through tissues. Slower SOS corresponds to lower stiffness.**

17 **The present study aimed to investigate whether SOS values reflect the degree of**

18 **degeneration with aging or dissection and whether enzymatic digestion**

19 **susceptibility is distinct.** SOS of media besides the atheromatous areas of normal and

20 surgical dissections was measured and compared using medial degeneration grade

21 (MDG) scores. To evaluate the damage rate, SOS was assessed following collagenase

22 digestion. SOS scores negatively correlated with aging and MDG scores. Dissected

23 aortae showed higher SOS and MDG scores without age correlation. Collagenase

24 digestion was present in all aortae, but older aortae were more injured than younger

25 aortae. Dissected aortae were more vulnerable to collagenase. Older and dissected

26 aortae expressed specific extracellular matrix (ECM) components to compensate for

27 mechanical weakness. The present method can evaluate mechanical weakness

28 corresponding to histology to investigate the cause of rupture.

29

30 **Keywords:** scanning acoustic microscopy, collagenase, extracellular matrix, thoracic

31 aorta, medial degeneration grade, aging, dissected aorta

32

33 **Introduction**

34 Aortic diseases are a major health concern that may result in aneurysm, dissection, and
35 atherosclerotic occlusion (Tsamis et al. 2013). A normally elastic aorta can begin to
36 stiffen with age (Radu-Ionita et al. 2017). Losing elasticity due to conditions such as
37 atherosclerosis or Marfan (MF) syndrome causes serious disorders.

38 The aortic wall media are organized into lamellar units, comprising concentric
39 layers of elastic lamellae, smooth muscle cells, and interlamellar matrix (Brooke et al.
40 2003). Elastic fibers are predominantly composed of elastin, whereas the interlamellar
41 matrix contains structural/supporting proteins, such as type-I collagen, type-III collagen,
42 and fibrillin (FBN). All extracellular matrix (ECM) components provide structural
43 organization and stability to the vessel wall through interactions between the smooth
44 muscle cells and associated ECM elements. Moreover, these elements include lysyl
45 oxidase (LOX), vitronectin (VN), and fibronectin (FN). Collagens bind and signal to
46 smooth muscle cells through specific matrix receptors. Elastic fibers are linked to
47 smooth muscle cells through a microfibril scaffold consisting of FBN and
48 microfibril-associated glycoproteins.

49 All aortic diseases are associated with microstructural changes in the content or
50 architecture of the connective fibers (Tsamis et al. 2013). Connective fibers consist of

51 elastin and collagen (Tsamis et al. 2013), which confer elasticity and strength to a
52 healthy aorta, respectively. Matrix metalloproteinases (MMPs), which include
53 collagenases, are a family of endopeptidases with proteolytic activity toward both
54 elastin and collagen (Choke et al. 2005). The high collagenase activity of the aorta
55 contributes to aneurysms and rupture (Menashi et al. 1987; Busuttil 1980). MMP-1
56 (collagenase-1), -8 (collagenase 2), and -13 (collagenase-3) are enzymes specific for
57 collagens. Collagenase-3 was used to imitate biochemical injury to aortae and detect
58 susceptibility to aneurysm and aortic rupture.

59 **The elasticity of the aortae has been reported in dogs (Hughes et al. 1979)**
60 **and humans (Sutton-Tyrrell et al. 2005; Lehmann et al. 1993) by measuring pulsed**
61 **wave velocity. Other methods to estimate aortic elasticity *in vivo* include**
62 **elastography (Fromageau et al. 2008) or micro-elastography (Schmitt et al. 2010),**
63 **which use shear waves, and 4D ultrasound (Wittek et al. 2013) which apply**
64 **time-resolved three dimensional ultrasound.**

65 These methods can provide macroscopic information about the elasticity of the
66 entire arterial wall, including the intima, media, and adventitia. However, they do not
67 address local tissue elements such as smooth muscle loss, accumulation of ECM
68 components, aneurysm, and dissection.

69 **Scanning acoustic microscopy (SAM) can evaluate both histological and**
70 **viscoelastic properties (Mamou and Rohrbach 2017; Miura 2016; Saijo 2009).**
71 **Since Lemons and Quate (1974) at Stanford University provided the basic design**
72 **of SAM in the biomedical field, many studies have reported on biological objects at**
73 **low frequencies (ranging from 1 to 10 MHz) (Maev 2008). Studies of the acoustic**
74 **properties of tissues at low frequencies suggest that SOS in soft tissues differs only**
75 **slightly from its value in water and is virtually independent of frequency (Duck**
76 **1990). Only solid tissue, such as bones, and tissues rich in fibrillar proteins showed**
77 **significant differences. Many acoustic images were collected and analyzed by**
78 **Quate's team, primarily using fixed unstained tissues. Low-frequency studies (1 to**
79 **7 MHz) of acoustic parameters of tissues (attenuation coefficients, SOS) have**
80 **demonstrated that fixation in 4% formalin changes these parameters only slightly**
81 **(Bamber et al. 1979).**

82 **Studies of acoustic properties of biological tissues at high frequencies (>**
83 **100 MHz) have recently began to obtain high-resolution quantitative images.**
84 **Improvements such as transducers with a very small F-number and high**
85 **sensitivity (Mamou and Rohrbach 2017) and time-frequency analysis method**
86 **(Hozumi et al. 2004) have contributed to this progress. Even single cells can now be**

87 **identified in the subcellular structures (Lemor et al. 2004; Weiss et al. 2007) and**
88 **clinical cytology samples (Miura and Yamamoto 2015).**

89 The speed-of-sound (SOS) of tissues observed using SAM is given as:

90
$$c = \sqrt{K/\rho},$$

91 where c is the SOS, K is the elastic bulk modulus, and ρ is the density.

92 This formula implies that the SOS strongly reflects its elastic parameter. SOS values
93 (m/sec) have been reported for many tissues (Azhari 2010; Saijo et al. 1998). In general,
94 SOS has low values in fluid-filled soft structures and high values in dense solid tissues.
95 SOS values are well-correlated with the palpable tissue stiffness. Therefore, ultrasound,
96 *in vivo* imaging-based methods such as elastography (Mahmood et al. 2016), and *in*
97 *vitro* tests on isolated samples, such as the SAM method, are available for estimating
98 the mechanical properties of biological samples. The current study used a stored
99 collection of ascending aorta (AAo) surgical specimens with detailed clinical history,
100 including dissection episode, to investigate the elasticity of the specimen and observe
101 histology to gain etiological insights.

102 Aortic walls progressively lose their components and increase their content in
103 fibrous tissue, leading to greater stiffness with aging. Aging aortae can lead to two

104 conditions involving greater or thinner wall thickness (Tsamis et al. 2013). The first is
105 atherosclerosis, which causes wall thickening due to development of a fatty plaque. The
106 second is aneurysm, which is produced by balloon-like thinning of the wall due to local
107 weakness. Focal portions of aortae fail to compensate mechanical strength to form
108 aneurysm or second bleb in aneurysm. The specific etiology of dissection due to aging
109 or genetic defects is unknown to date (Wu et al. 2013).

110 The aim of the present study was to investigate whether SOS values reflect the
111 degree of aortic degeneration and whether resistance to collagenase digestion is
112 different between aged and dissected cases. Moreover, the biochemical changes in ECM
113 components were compared between aging aortae and dissected aortae.

114

115 **Materials and Methods**

116 *Subjects and ethics*

117 All human tissue sections were obtained from samples stored in the tissue archives of
118 the Hamamatsu University Hospital or Shizuoka City Hospital in Japan. The AAO of
119 adult autopsied patients without serious cardiovascular diseases were consecutively
120 selected to investigate the effects of aging on the biomechanical properties of the aortae
121 (n = 36; age, 62.9 ± 20.0 years; 24 men and 12 women). Cases of MF syndrome (n = 7;

122 age, 46.7 ± 16.2 years; three men and four women) and non-MF syndrome ($n = 9$; age,
123 56.3 ± 12.0 years; seven men and two women) were selected from surgical specimens of
124 dissected AAO aneurysm with known clinical history. Formalin-fixed,
125 paraffin-embedded (FFPE) tissue blocks were flat-sectioned into 10- μ m-thick slices and
126 observed using SAM. The research protocol for using stored samples without a link to
127 patient identity was approved by the Ethics Committee of Hamamatsu University
128 School of Medicine (No. 14-135, 19-180). Written consent was waived based on the
129 retrospective design. All procedures were conducted according to approved guidelines
130 and regulations.

131

132 *SAM observations*

133 **The experimental protocol is detailed in Fig. 1. Aortic tissue specimens were**
134 **evaluated using SAM system (AMS-50AI, Honda Electronics, Toyohashi, Aichi,**
135 **Japan) with a central frequency of 320 MHz, lateral resolution of 3.8 μ m, and**
136 **thickness of the focal spot of 13 μ m. A single-pulsed ultrasound with 2-ns pulse**
137 **width was emitted (Hozumi et al. 2004). Distilled water was used for coupling fluid**
138 **between the transducer and the specimen. The transducer was used for both**
139 **transmitting and receiving the signal. Reflected waveforms from the surface and**

140 **the bottom of specimen were compared to measure the SOS and the thickness of**
141 **each point. Waveform from the glass surface without the specimen present was**
142 **used as a reference waveform.**

143 **Scanning the transducer over the specimen formed an acoustic image. The**
144 **mechanical scanner was arranged so that the ultrasonic beam was transmitted at**
145 **every 8, 4 and 2 μm interval over a 2.4, 1.2, and 0.6 mm width, respectively. The**
146 **number of sampling points was 300 in one scanning line, and 300 x 300 points**
147 **made one frame. Four pulse echo sequences were arranged for each scan point in**
148 **order to increase the signal-to-noise-ratio. Each pixel of an image corresponds to**
149 **an echo coming from an x-y position on the specimen.**

150 The observed samples were prepared by cutting FFPE blocks. Although FFPE samples
151 have been demonstrated as having slightly higher SOS than fresh samples, the SOS
152 values were stable (Sasaki et al. 1996) irrespective of periods of formalin fixation from
153 1 day to 3 months (Miura et al. 2015). Therefore, sample bias due to fixation condition
154 was negligible. Areas without calcified deposits and heavy atheroma were selected for
155 comparison. **Calcified areas result in chatter marks on the section, which causes**
156 **irregular reflection and heavy atheromatous portions become translucent due to**
157 **lipid dissolution in organic solvents. In dissected cases, non-dissected portions**

158 away from the separation were used for SOS measurement.

159

160 *SOS difference according to age and disease status (normal, MF syndrome, and*
161 *non-MF syndrome)*

162 To evaluate age-related changes of the aorta, the average SOS values of the aortic media
163 were plotted according to age. To compare the mechanical weakness and structural
164 differences, the samples were divided into two groups: (1) younger adults aged 31–58
165 years (average, 46.7 ± 11.47 years; $n = 6$; four men and two women) and (2) older
166 individuals aged 76–85 years (average, 80.2 ± 3.42 years; $n = 5$; four men and one
167 woman). The SOS values of dissected aortae (including MF syndrome and non-MF
168 syndrome) ($n = 16$; age 52.1 ± 14.3 years; 10 men and 6 women) were also compared
169 with age-matched normal aortae ($n = 12$; age 56.8 ± 15.7 years; 8 men and 4 women) by
170 plotting the average SOS values of the aortic media according to age.

171

172 *Catalytic damage according to collagenase digestion*

173 Paraffin sections were dewaxed using xylene, soaked in distilled water, and submerged
174 into a solution of phosphate-buffered saline containing 0.5 mM CaCl_2 (pH 7.4) plus 250
175 units/mL type-III collagenase (Worthington, Lakewood, NJ, USA) at 37°C for 1.5 h or

176 twice for 1.5 h (Miura and Katoh 2016). According to the manufacturer's instructions,
177 type-III collagenase has typical collagenase activity but lower proteolytic activity than
178 other collagenases. SOS was measured both before (baseline) and after digestion.
179 Digested sections were first washed with distilled water before being analyzed with
180 SAM. The same sections were measured after a repeated digestion (two 1.5-h
181 treatments).

182

183 ***Medial degeneration grade (MDG) evaluation***

184 To compare SAM with light microscopy (LM) images, the same or nearby sections were
185 stained with hematoxylin, eosin, and Elastica Masson trichrome (EMT) to stain collagen
186 and elastic fibers blue and black, respectively. The magnification of each LM image was
187 adjusted to match the corresponding SAM image with scale bars on the bottom and the
188 left side of the screen frame.

189 Consensus criteria was used to evaluate the extent and severity of the aortic
190 MDG scores (Halushka et al. 2016). These criteria included mucoid extracellular matrix
191 accumulation, elastic fiber fragmentation/loss, smooth muscle cell nuclei loss, and
192 laminar medial collapse. The overall MDG was scored as 0 (none), 1 (mild), 2
193 (moderate), or 3 (severe).

194

195 ***Immunohistochemical analysis***

196 Immunostaining was performed using a commercially available Chemmate envision kit
197 (Dako, Glostrup, Denmark). The following primary antibodies used: anti-smooth
198 muscle actin (SMA) (ab5694, Abcam, Tokyo, Japan; 1:400), anti-collagen I (ab88147,
199 Abcam, 1:100) and anti-collagen III (ab7778, Abcam, 1:1000) for collagen types,
200 anti-LOX (ab174316, Abcam, 1:300), anti-VN (ab46808, Abcam, 1:250), anti-FBN
201 (ab53076, Abcam, 1:25), and anti-FN (ab2413, Abcam, 1:250). Heat-mediated antigen
202 (95°C, 20 min) was retrieved with buffer balanced to either pH 6.0 (anti-SMA,
203 anti-collagen III, FN) or pH 9.0 (anti-collagen I, anti-LOX, anti-VN) before staining.

204

205 ***Statistical analyses***

206 The average SOS values of the aortic media were calculated from at least five images
207 per case using the SAM manufacturer and commercial statistics software
208 (Statcel3-Addin forms on Excel, OMS publishing, Tokorozawa, Saitama, Japan), which
209 calculates the average areas-of-interest values. To detect the correlations between age
210 and SOS values and between age and MDG scores, scatter plots were established and
211 subjected to simple linear regression analysis. The correlation strength was quantified

212 using Pearson's correlation coefficients (r). The average SOS (\pm SD) values from
213 younger and older aortae and between dissected aortae and age-matched normal aortae
214 were compared using unpaired Student's t -tests. The average SOS values (\pm SD) among
215 different time points after collagenase were compared using paired t -tests.

216 It was confirmed that each data set followed a normal distribution pattern before
217 statistical analyses were conducted. Significance level for all tests was set at $p < 0.05$.

218

219 **Results**

220 *Age-dependent changes in SOS and MDG of normal aortic media*

221 SOS values of the normal aortic media without calcified deposits and heavy atheroma
222 potions decreased with age (Fig. 2a). SOS values were negatively correlated with age (y
223 $= -0.6032x + 1699.3$, $r = -0.39$, $p = 0.015$).

224 The significantly positive relationship between age and MDG scores ($y =$
225 $0.021x + 0.21$, $r = 0.56$, $p = 0.0003$) is shown in Figure 2b. **The significantly negative**
226 **relationship between SOS and MDG scores ($y = -0.012x + 22.47$, $r = -0.505$, $p =$**
227 **0.0016) is shown in Figure 2c.**

228

229 *Relationship between age and SOS or MDG of dissected aortae*

230 **SOS values and MDG scores of dissected aortae were compared with those of**
231 **age-matched normal ones (Fig. 2d).** The dissection group exhibited significantly
232 greater SOS values ($p = 8.50E-10$) and higher MDG scores ($p = 0.043$) than the normal
233 group. No significant correlation between age and SOS values ($y = -0.2334x + 1738$, r
234 $= -0.17$, $p = 0.501$) (Fig. 2a) or age and MDG scores ($y = 0.0025x + 1.83$, $r = 0.075$, p
235 $= 0.772$) (Fig. 2b) was observed in dissected aortae. **There was also no significant**
236 **relationship between SOS value and MDG scores ($y = -0.0054x + 11.45$, $r = -0.21$, p**
237 **$= 0.42$) (Fig. 2c).**

238

239 *Differential changes in SOS values between older and younger aortae after*
240 *collagenase digestion*

241 Before collagenase digestion, the older aortic media ($n = 22$) showed significantly lower
242 SOS values than the younger aortae ($n = 20$) (Figs. 3a and 3b; Table A1). After
243 collagenase digestion, both older and younger aortae showed lower SOS values. At 1.5
244 h after digestion, SOS values significantly decreased in both cases ($p < 0.01$) (Fig. 4).
245 EMT staining revealed wavy, thin, and split elastic fibers in the media from the older
246 aorta, whereas the younger aortae contained thicker and straighter fibers. LM images
247 obtained 3 h after digestion revealed numerous fragmented elastic fibers in the older

248 aortae compared to the younger aortae. Similar to the baseline difference at 0 h, older
249 aortae had significantly lower SOS values than younger aortae after 1.5 h and 3 h of
250 collagenase digestion (p 's < 0.01).

251

252 ***Differential changes in SOS values between dissected aortae (MF and non-MF***
253 ***syndrome) and normal aortae after collagenase digestion***

254 Cases of dissected aortae with MF had cystic mucoid degeneration and elastic fiber
255 fragmentation or loss. Cystic degeneration displayed low SOS values (Fig. 3c). Non-MF
256 cases consisting of thick parallel muscles and wavy elastic fibers with focal splitting
257 showed high SOS values (Fig. 3d). SOS values decreased faster after digestion in cases
258 of dissected aortae compared to age-matched controls (Fig. 4 and Table A2). At baseline,
259 the two groups showed significantly different SOS values ($p = 0.0001$). However, the
260 difference in SOS values disappeared after 1.5 h ($p = 0.100$) and 3 h ($p = 0.346$). In
261 contrast to the SOS results, LM images in EMT staining showed no major differences
262 between MF syndrome and non-MF syndrome cases. There was slight muscle fiber
263 disappearance and loose elastic fibers before and after digestion.

264

265 ***Structure components of SMA, collagen type-I, and collagen type-III in the older,***

266 *younger, MF, and non-MF aortae*

267 Regarding SMA immunostaining (Fig. 5a), old aortae showed focal loss of smooth
268 muscles, whereas younger aortae consisted of continuous parallel muscles. MF cases
269 displayed aggregated muscle fibers with irregular arrangement, whereas non-MF cases
270 showed scattered loss of smooth muscle fibers.

271 Regarding collagen type-I (Fig. 5b) and type-III (Fig. 5c), normal young and
272 old cases displayed parallel collagen fibers. Non-MF cases had rich collagen fibers,
273 although fiber arrangement was irregular. MF cases had poor collagen fibers with
274 irregular arrangement among cells. Table 1 details the immunohistochemical results.

275

276 *ECM components of LOX, VN, FBN, and FN in the younger, older, MF, and non-MF*
277 *aortae*

278 Immunostaining with anti-LOX (Fig. 6a) showed that normal younger and older aortae
279 exhibited no to faint positive staining. However, the MF aortae revealed strong
280 punctate-patterned staining on smooth muscles, whereas the non-MF aortae showed
281 weak positive staining.

282 Regarding the VN (Fig. 6b), all normal young or old aortae and MF and
283 non-MF ones were focally positive, although MF cases showed a rather irregular

284 distribution.

285 About FBN (Fig. 6c), all cases except MF were positive. Concerning FN (Fig.
286 6d), old and non-MF cases were positively associated with muscle fibers, whereas
287 young and MF aortae exhibited negative or focal weak positivity.

288

289

290 **Discussion**

291 Both structural and mechanical alterations in AAo were found with aging and dissection
292 cases using SAM combined with collagenase treatment and immunostaining. Age was
293 negatively correlated with SOS values based on SAM results, suggesting that the
294 non-atheromatous or calcified portions of the aorta become mechanically weaker with
295 age. In general, aging aortic walls show an increased stiffness due to atherosclerosis and
296 calcification. However, non-atheromatous portions of media showed loose and irregular
297 arrangement of smooth muscles and collagen fibers, which contributed to aneurysms
298 and rupture. **The MDG scores that increased with aging were significantly**
299 **negatively correlated with SOS values of non-atheromatous portions. The dissected**
300 **aortae cases, in which SOS values and MDG scores were both higher than those of**
301 **their age-matched controls, showed no significant alteration with aging.** This may

302 indicate that dissected aortic walls have greater stiffness to compensate for structural
303 defects, but are focally fragile and prone to ruptures. The susceptibility to collagenase
304 digestion supported this speculation. Other risk factors for dissection apart from aging
305 must also be present, such as genetic susceptibility. Not only MF syndrome cases but
306 also non-MF young cases may present genetic defects of ECM components.

307 Regarding damage caused by collagenases, both young and older cases
308 exhibited significantly decreased SOS values following 1.5 h and 3 h of collagenase
309 treatment. Young aortae retained rather high SOS values even after 3 h, whereas older
310 aortae reached minimum values close to the risk threshold of rupture. At 3 h, the old
311 aortae were composed of loose filamentous fibers. The dissected aortae group, including
312 MF and non-MF cases, had very high SOS values at baseline and more rapidly
313 decreased values after 1.5 h and 3 h as compared to their age-matched controls.

314 **Dissected aortae may have greater stiffness due to compensation for structural**
315 **defects. Sufficient number of structural components offset fragile quality, as seen**
316 **in smooth muscle bundles of MF cases and rich collagen type-III fibers of non-MF**
317 **cases.**

318 The fibrous portions of the dissected aortae had consistent SOS values,
319 whereas the non-fibrous portions were more sensitive to enzymatic damages. Busuttil et

320 al. (1980) reported that collagenase activity is detected in the aneurysmal wall of
321 abdominal aorta. Therefore, positive results for collagenase testing may predict future
322 rupture.

323 Immunostaining of the structural fibers in young aortae showed a continuous
324 parallel array, while the structural fibers in the old aortae were divided or sparse.
325 Regarding ECM proteins, FN was enriched in the older aortae compared to the younger
326 aortae. In the dissected aortae, MF cases had irregular distribution of SMA, collagen I,
327 and collagen III fibers, whereas most non-MF cases had distribution that was similar to
328 the normal older aortae. MF cases were positive for LOX and VN antigens, although
329 FBN and FN immunostaining was negative or faint. Both non-MF and elderly cases
330 exhibited a similar pattern of positivity for LOX, VN, FBN, and FN.

331 The histopathological alterations in dissection cases include smooth muscle cell
332 disappearance, medial mucoid degeneration, and ECM breakdown (Michel et al. 2018;
333 Wu et al. 2013). Dissection is performed because of genetic causes or degeneration with
334 aging (Michel et al. 2018). Among the 29 thoracic dissected aortae-associated genes
335 identified to date, the majority encode proteins involved in the extracellular matrix,
336 smooth muscle cell contraction or metabolism, or transformation of the growth factor- β
337 signaling pathway (Brownstein et al. 2017). SMA, collagens 1 and 3, elastin, FBN1, and

338 LOX are also included. MF syndrome, an autosomal dominant inherited disorder caused
339 by mutations in FBN1 (Dietz 1991), was reported to have increased expression of
340 matrix metalloproteinase (MMP)-2 and MMP-9 and premature aortic smooth muscle
341 cell differentiation (Dale et al. 2017).

342 Regarding ECM proteins, LOX is an extracellular amine oxidase that primarily
343 functions as a catalyst on the covalent cross-linking of collagen and elastin fibers in
344 ECM (Kagan and Li 2003). VN in ECM is involved in tissue repair and remodeling
345 (Leavesley et al. 2013). FBN-1 plays a crucial role in stabilizing the structure of elastic
346 fiber. FN plays roles in cell adhesion, migration, growth, and differentiation (Sottile
347 2002). FN in ECM controls deposition, organization, and stability of other matrix
348 proteins, including type-I collagen and type-III collagen. Specific types of ECM
349 proteins are expressed differently among MF, non-MF, and normal old cases to maintain
350 the mechanical strength of the aortic wall.

351 Although the resolution and flexibility in SAM imaging is not better than LM
352 imaging, it still has several advantages. First, no staining is required, which allows for
353 imaging of the sample within a few minutes. **Second, SOS values are digital, allowing**
354 **easy quantitative comparison among lesions.** Third, the virtual color images of SOS
355 values are adjustable based on the range of interest, facilitating easy detection of

356 abnormal regional alterations. Fourth, protease digestion alterations can be repeatedly
357 monitored with greater sensitivity compared to LM.

358 This study has several limitations that should be acknowledged. First, the
359 observed tissues were FFPE sections. Compared with fresh tissues, tissues fixed for
360 extended periods, such as those used for autopsies, become harder. However, the current
361 study reported SOS values in normal autopsy cases that were lower than those in
362 surgical specimens. The SOS values of fresh and formalin-fixed aortae have been
363 previously reported. Akhtar et al. (2016) reported that the SOS value of fresh aortae was
364 between 1,538 and 1,709 m/s. Saijo et al. (1998) reported SOS values of $1,614 \pm 30$ m/s
365 in formalin-fixed frozen sections of the aorta without paraffin-embedded treatment. Our
366 cases showed that the overall **mean** SOS value was $1,648.4 \pm 30.2$ m/s (age range, 16–
367 101 years), indicating that SOS values in the FFPE aortae were somewhat greater than
368 those in fresh-frozen aortae, but similar to those previously reported. Sasaki et al.
369 (1996) reported the influence of formalin fixation on the acoustic properties of the
370 normal kidney. No significant differences in acoustic parameters, including SOS values
371 or attenuation were observed. The second limitation is that sex differences were not
372 considered. The consecutive autopsy and surgical tissue samples used were
373 predominantly from men, with a few samples from younger women. Because fewer

374 younger and postmenopausal women present with similar vascular alterations as men
375 (Waddell et al. 2001), gender bias may be low. Third, the non-MF group included
376 different etiologies; therefore, LM images might be distinctly different in non-MF cases.
377 However, these non-MF cases had some common histological findings, such as an
378 intralamellar mucoid matrix and fragmentation or thinning of the elastic fibers, as
379 demonstrated in representative images. Despite these limitations, SAM analysis with
380 protease digestion is a useful method to evaluate mechanical weakness due to histology
381 features and to investigate the cause of rupture.

382 **In conclusion, SOS values appropriately reflected mechanical weakness**
383 **and degeneration with aging. Dissected aortae had higher SOS values irrespective**
384 **of high MDG scores. All normal and dissected aortae were vulnerable to**
385 **collagenase digestion. Old aortae first reached the lowest level of SOS, and**
386 **compared with normal aortae, dissected ones showed more susceptible to**
387 **collagenase digestion. Old and dissected aortae had different structural**
388 **components to protect against mechanical weakness of the thoracic aorta.**

389

390

391 **Acknowledgments**

392 The authors thank T. Moriki, Y. Egawa, Y. Kawabata, and N. Suzuki for their help in
393 preparing the histological samples, Dr K. Kobayashi for his technical supports and
394 advices with SAM, and to Enago (www.enago.jp) for the English language review.

395

396 **Funding**

397 This work was supported by a grant from the Japan Society for the Promotion of
398 Science (KAKENHI), Scientific Research Grant Number (c) 15K08375.

399

400 **Conflict of interest**

401 None.

402

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521

522 **Figure captions**

523

524 **Fig. 1 Flow chart of SAM observation**

525

526 **Fig. 2** Relationship among age, **mean** speed-of-sound (SOS), and grade of medial
527 degeneration (MDG) in normal and dissected aortae

528 a. Relationship between average SOS value and age of normal and dissection aortae. b.

529 Relationship between MDG scores and age of normal and dissected aortae. **c.**

530 **Relationship between SOS value and MDG scores. d. Boxplot showing the SOS and**

531 **MDG difference between normal and dissected cases.** r = Peason's correlation
532 coefficient.

533

534 **Fig. 3** Representative speed-of-sound (SOS) images of the aortae after collagenase

535 type-III digestion (a: 85-year-old woman, b: 31-year-old man, c: Marfan (MF)

536 syndrome, d: Non-MF syndrome)

537 SOS images exhibited gradual decreases in SOS values at 1.5 h and 3 h after digestion.

538 The corresponding light microscopy (LM) images of Elastica Masson trichrome (EMT)

539 staining before and after 3 h of collagenase digestion are shown in the upper rows. The

540 elderly aorta (b) appears with more fragmentation of elastic fibers. The aortae in both

541 MF (c) and non-MF (d) cases exhibit rapidly decreasing SOS values after collagenase
542 digestion. The MF case with cystic mucoid degeneration shows punctate low SOS areas
543 on the right side. The non-MF aorta consisting of parallel muscle fibers and wavy
544 elastic fibers with focal splitting shows high SOS at the baseline. Black bar = 400 μm ,
545 red bar = 10 μm .

546

547 **Fig. 4** Reduction of SOS after collagenase digestion (left: young and old, right:
548 dissection and age-matched normal controls)

549 SOS of all groups decreased from 0 to 1.5 h, and 1.5 h to 3 h after collagenase digestion.
550 SOS values in dissection cases decreased more rapidly than those in normal
551 age-matched aortae. *** $p < 0.001$, ** $p < 0.01$, * $p < 0.05$.

552

553 **Fig. 5** Immunostaining with anti-smooth muscle actin (SMA) (a), anti-collagen type-I
554 (COL 1) (b), and anti-collagen type-III (COL 3) (c) antibodies of young, elderly, MF,
555 and non-MF sections

556 Regarding the SMA, the young aortae showed parallel linear smooth muscles, whereas
557 the old aortae exhibited split smooth muscle fibers. MF cases displayed aggregated
558 SMA bundles with irregular arrangement, whereas non-MF cases showed frayed muscle

559 fibers with irregular arrangement. Regarding COL 1 and COL 3, young and elderly
560 cases displayed parallel fiber components between smooth muscle fibers, although the
561 fiber array was irregular in old cases. MF cases had irregular fiber arrangements among
562 cells, whereas non-MF cases exhibited rich collagen fibers in a focal irregular array.

563

564 **Fig. 6** Immunostaining with anti-lysyl oxidase (LOX) (a), vitronectin (VN) (b), fibrillin
565 (FBN) (c), and fibronectin (FN) (d) antibodies of young, elderly, MF, and non-MF
566 sections

567 The normal young aorta was negative for LOX, whereas the old aorta showed focal
568 faint positivity. MF aortae exhibited strong punctate positivity on smooth muscles for
569 LOX, whereas non-MF aortae showed focal weak positivity. In VN, all normal young or
570 old aortae and MF and non-MF aortae, displayed focal positivity, although MF cases
571 show rather irregular distribution. For FBN, only MF cases exhibited negative. For FN,
572 old and non-MF cases show positivity along muscle fibers.

573

Fig. 1

Autopsy or Surgical specimen of aortae

10% buffered formalin fix



Vertical cut aortae



10- μ m flat section

LM sample



Obtain SAM image, use distilled water as coupling medium



Calculate average SOS values of media except calcification and severe atherosclerosis



Collagenase digestion if necessary



Time-lapse observation of the same section

Fig. 2a

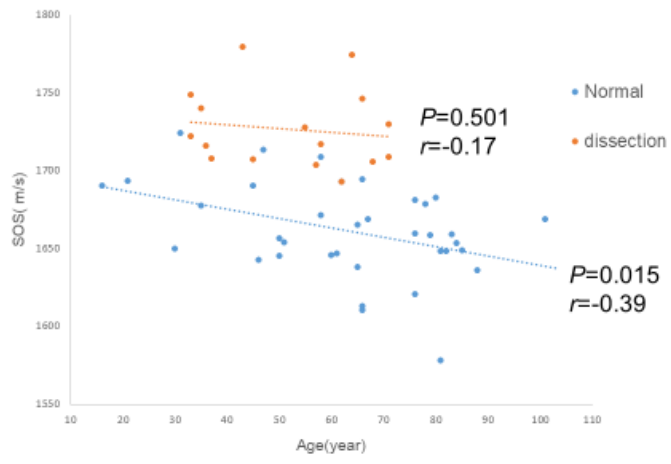


Fig. 2b

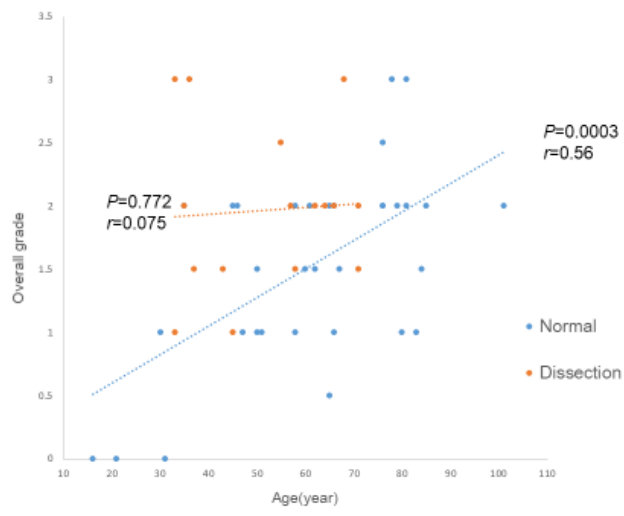


Fig. 2c

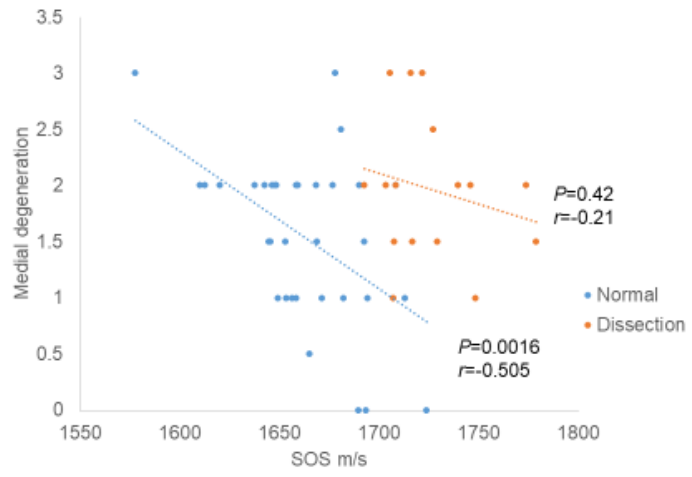


Fig. 2d

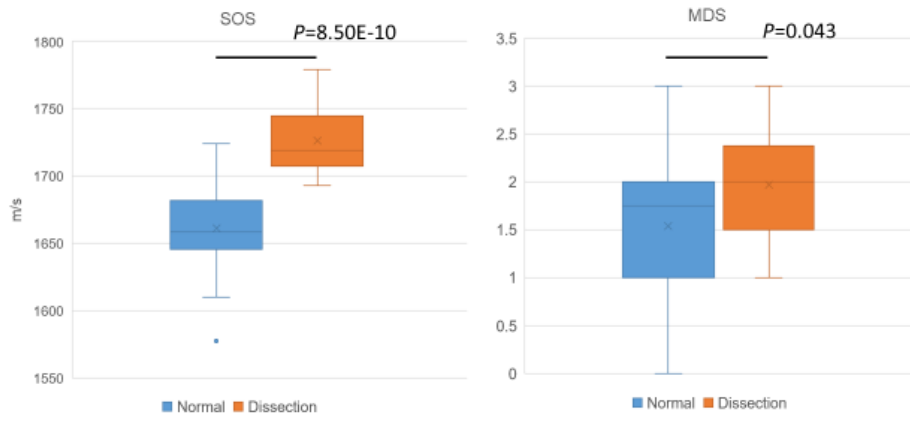


Fig. 3a

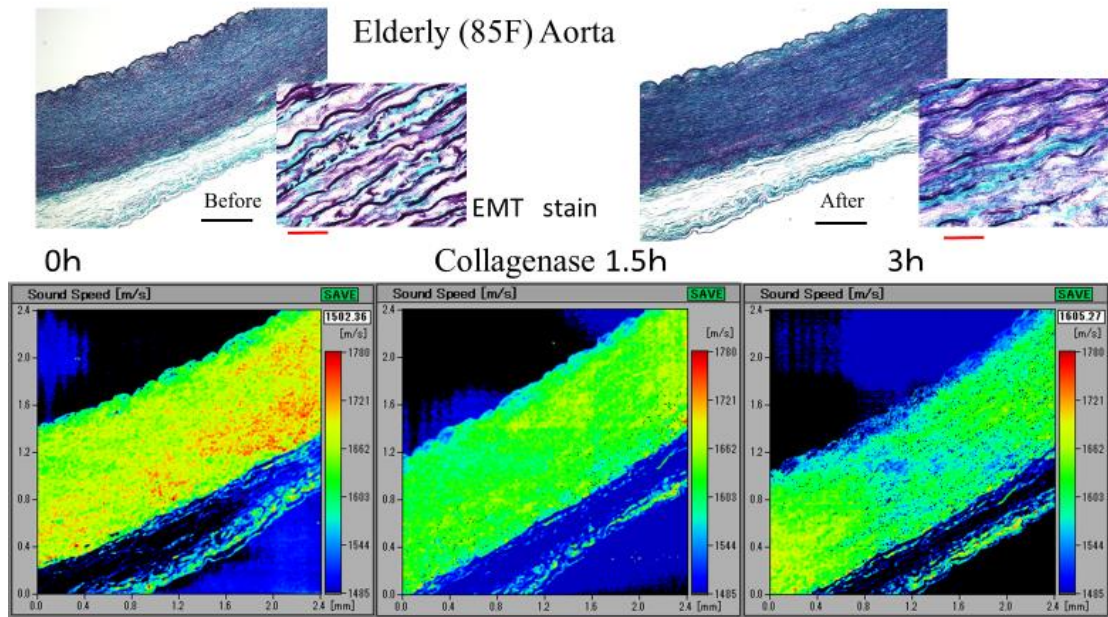


Fig. 3b

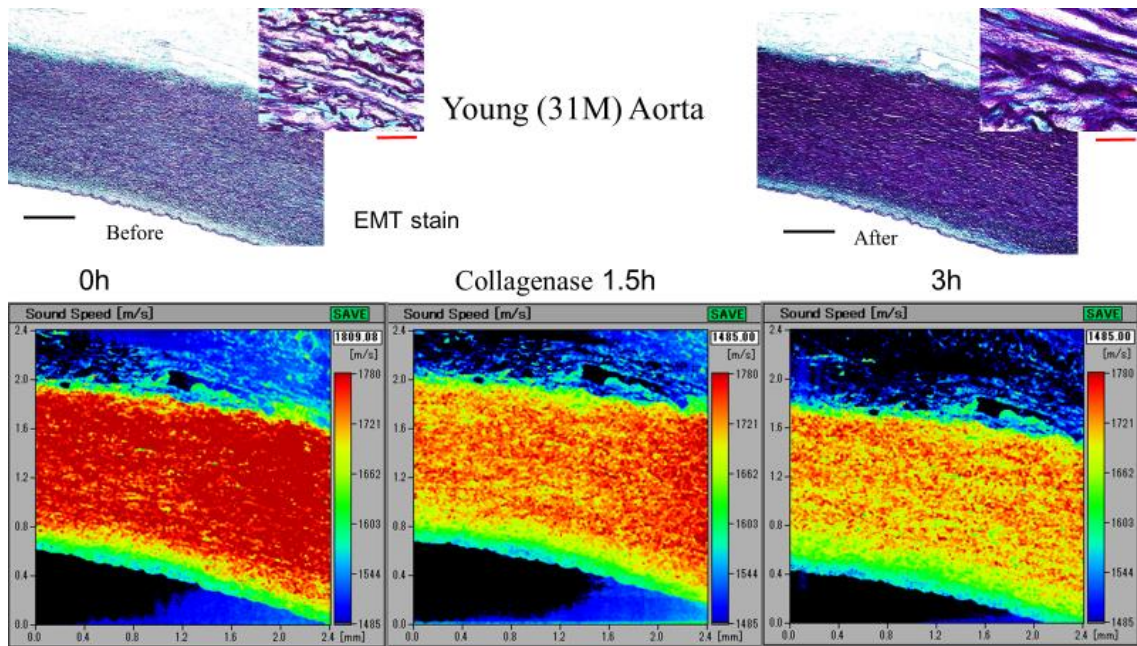


Fig. 3c

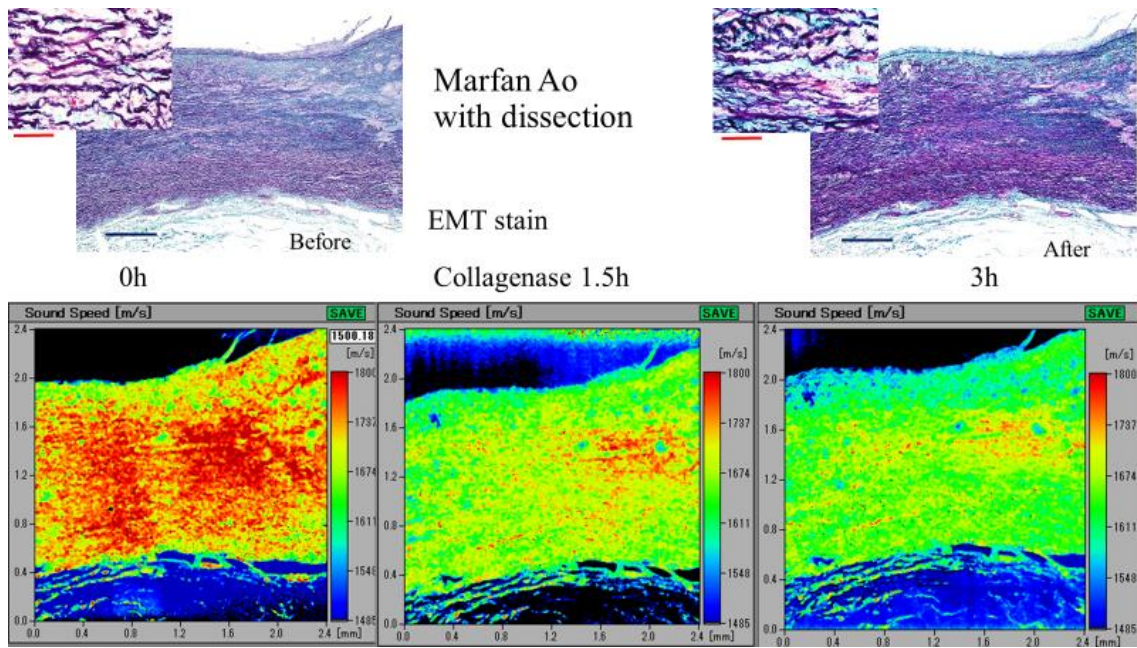


Fig. 3d

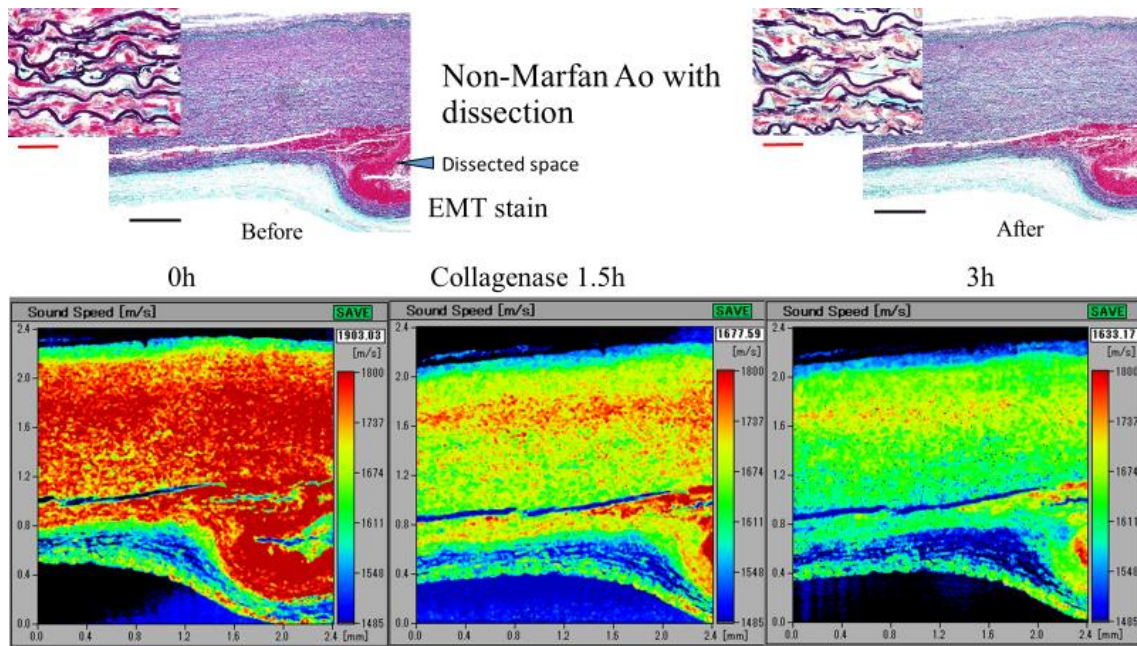


Fig. 4

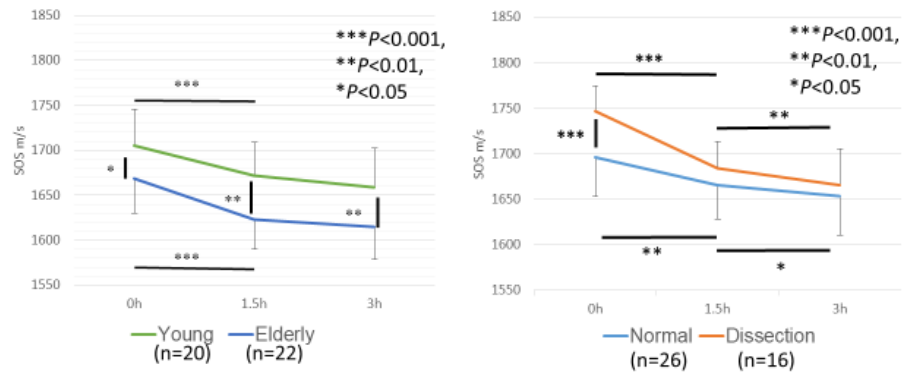


Fig. 5a

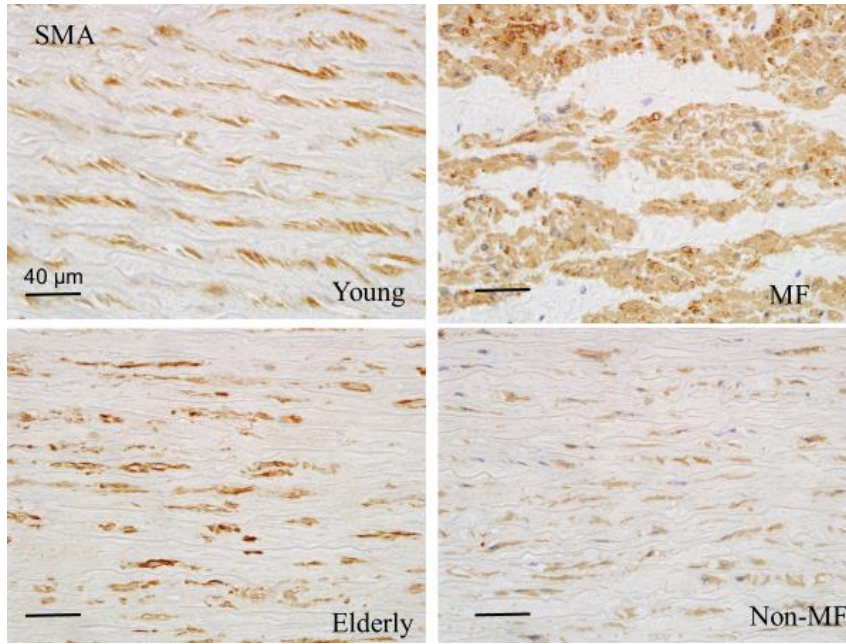


Fig. 5b

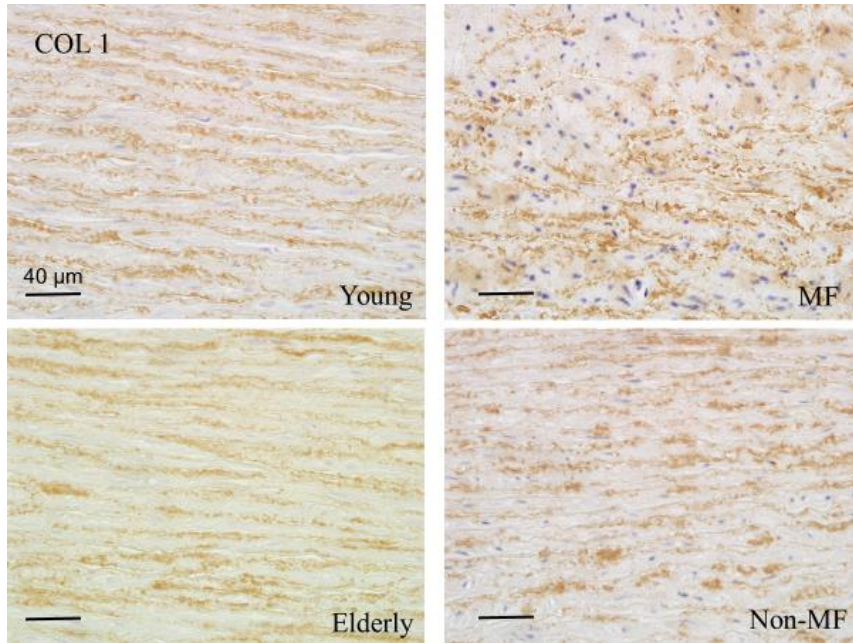


Fig. 5c

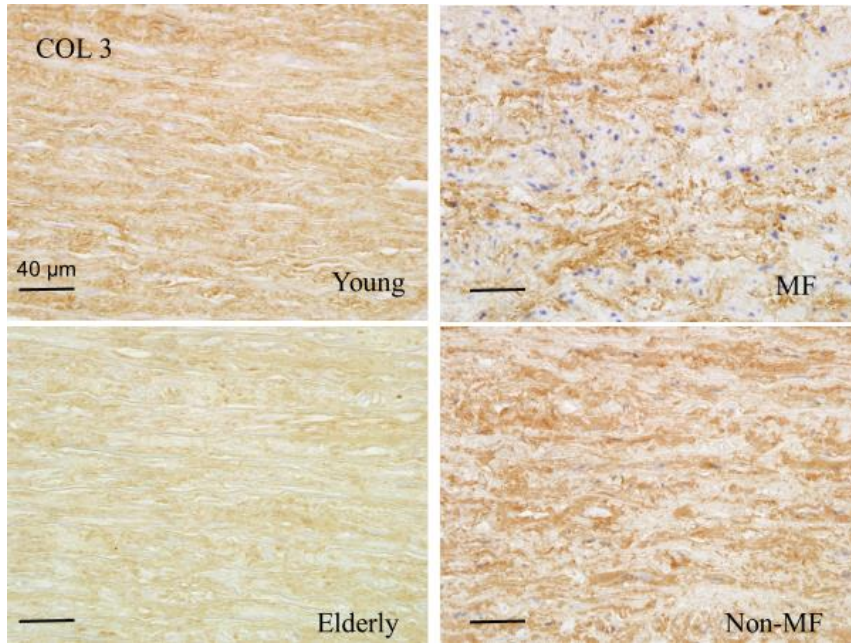


Fig. 6a

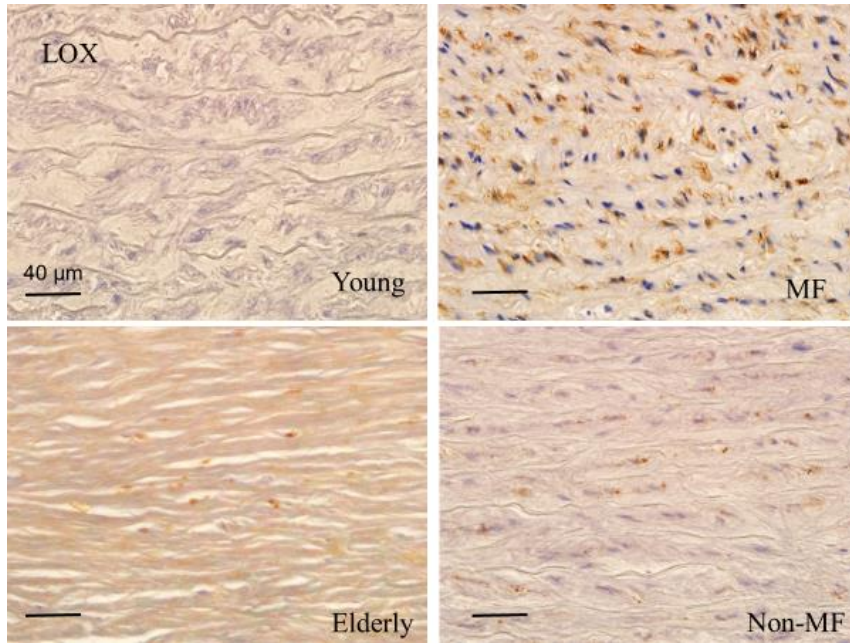


Fig. 6b

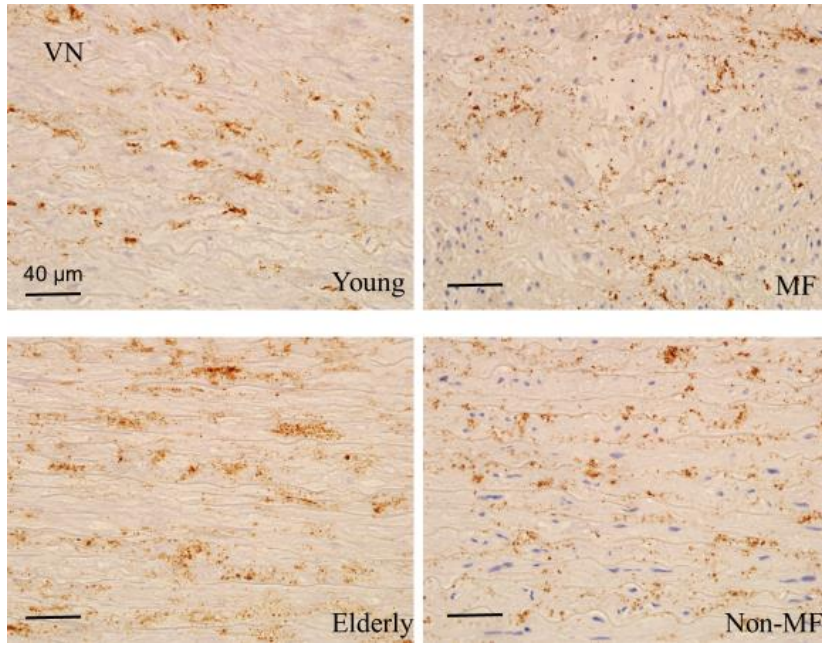


Fig. 6c

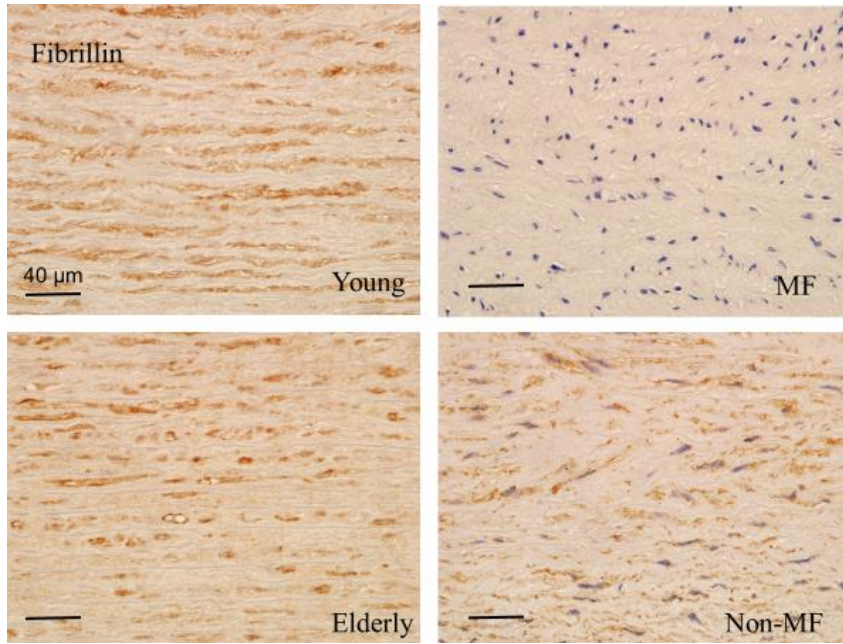
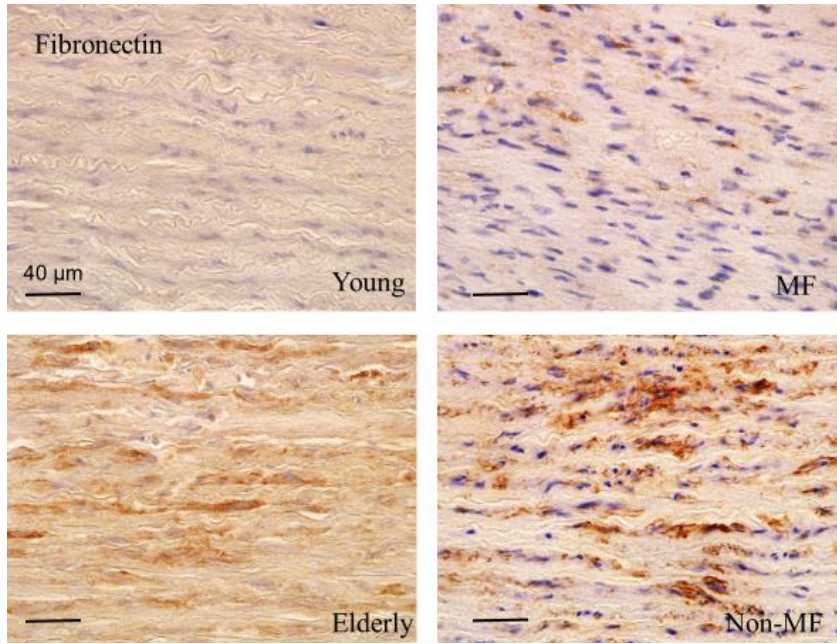


Fig. 6d



Table

Table 1. Immunostaining results of the supporting components

Case	SMA	Col 1	Col 3	LOX	VN	FBN	FN
N-Young	+	+	+	–	+	+	–
N-Old	+F,W	+F,W	+	+W	+F	+	+
MF	+F	+F	+F	++	+	–	+F,W
Non-MF	+F,W	+F,W	+F,W	+F	+	+	+

N-Young, normal young; N-Old, normal old; MF, Marfan syndrome; F, focal; W, weak;

SMA, smooth muscle actin; Col 1, collagen type-I; Col 3, collagen type-III; LOX, lysyl

oxidase; VN, vitronectin; FBN, fibrillin; FN, fibronectin

Table A.1. Average speed-of-sound (SOS) of older (OL) and younger (YG) aortae after collagenase digestion

Collagenase	n	Average	SD	
				m/s
YG0h	20	1705.1	40.6	
YG1.5h	20	1672.3	36.7	
YG3h	20	1658.8	43.5	
OL0h	22	1668.4	38.1	
OL1.5h	22	1622.7	32.9	
OL3h	22	1614.5	36.0	

*** $p < 0.001$. ** $p < 0.01$, * $p < 0.05$

- 1 Table A.2. Average speed-of-sound (SOS) of normal (NOR) and dissection (DIS) aortae
- 2 after collagenase digestion.

Collagenase	n	Average m/s	SD	
0h-NOR	26	1695.8	42.5	**
1.5h-NOR	26	1665.6	37.7	
3h-NOR	26	1653.3	42.5	
0h-DIS	16	1741.1	27.6	***
1.5h-DIS	16	1684.0	29.0	
3h-DIS	16	1665.8	39.3	

3 *** $p < 0.001$, ** $p < 0.01$, * $p < 0.05$

4