**ABSTRACT.** Aortic rupture is a well recognized cause of sudden death in thoroughbred horses. Some microscopic lesions, such as those caused by cystic medial necrosis and medionecrosis, can lead to aortic rupture. However, these microscopic lesions are also observed in normal horses. On the other hand, a previous study of aortic rupture suggested that underlying elastin and collagen deposition disorders might be associated with aortic rupture. Therefore, the purpose of this study was to compare the structural components of the tunica media of the aortic arch, which is composed of elastin, collagen, smooth muscle cells and mucopolysaccharides (MPS), in fetal to mature thoroughbred horses. The percentage area of elastin was greatest in the young horses and subsequently decreased with aging. The percentage area of collagen increased with aging, and the elderly horses (aged ≥20) exhibited significantly higher percentage areas of collagen than the young horses. The percentage area of smooth muscle cells did not change with age. The percentage area of MPS was inversely proportional to the percentage area of elastin. The fetuses exhibited a markedly larger percentage area of MPS than the mature horses. We concluded that the medial changes seen in the aortic arch, which included a reduction in the amount of elastin and increases in the amounts of collagen and MPS, were age-related variations.

**KEY WORDS:** aorta, aortic arch, equine, histopathology, morphometry

In horses, rupturing of the aorta is a well recognized, but uncommon condition. It mainly affects elderly horses [1, 9] and can lead to sudden death [3]. Histopathologically, aortic lesions, including cystic medial necrosis, elastin fragmentation and medionecrosis of the aortic tunica media, might contribute to aortic rupturing [1, 9]. While such lesions have been evaluated using medial grading systems based on their extent and localization [5, 18, 23], the relationship between these lesions and aortic rupturing remains unclear [14, 19, 21].

On the other hand, recent studies have analyzed the morphometric properties of the aorta in horses, including the proportions of elastin, collagen and smooth muscle cells (SMC) [15, 16]. However, they did not examine horses with a broad age range. In the present study, histopathological and morphometric analyses of the tunica media of the aortic arch were performed in 90 horses ranging in age from 97 days (embryonic horses) to 29 years old.

**MATERIALS AND METHODS**

**Animals**

Ninety thoroughbred horses ranging in age from 97 days to 29 years old were included in the present study (Table 1). Macroscopic and microscopic examinations were performed in 90 and 77 of these animals, respectively. All of the horses underwent postmortem examinations at Rakuno Gakuen University after being euthanized or suffering spontaneous death (mainly due to locomotive or alimentary disorders). Horses with cardiovascular disorders were excluded. The investigated animals were divided into 7 groups (Table 1). Group 1 was comprised of fetuses, group 2 was comprised of newborns-5-month-old horses, group 3 was comprised of 6-11-month-old horses, group 4 was comprised of 1–2-year-old horses, group 5 was comprised of 3–9-year-old horses, group 6 was comprised of 10–19-year-old horses, and group 7 was comprised of horses aged over 20 years.
In all horses, a section of the aortic arch extending from the brachiocephalic trunk to the first cranial artery; i.e., the dorsal intercostal artery or the bronchoesophageal artery, and the surrounding connective tissue were removed. Samples were collected from three sections of the aorta: proximal samples were collected from the region between the brachiocephalic trunk and the ligamentum arteriosum, “middle” samples were obtained from the region starting 2 cm caudal from the ligamentum arteriosum, and distal samples were acquired from the region between the middle sample and the dorsal intercostal artery or bronchoesophageal artery.

Gross examinations
Macroscopically, the luminal diameter (cm) and thickness (mm) of cross-sections of the fixed aorta were measured in 90 horses.

Histological examinations
After the macroscopic examinations, dorsal aortic samples from 77 horses were fixed in 10% formalin and used for histopathological examinations. The samples were processed routinely and embedded in paraffin wax. The resultant sections (4 µm) were stained with hematoxylin and eosin (HE). In addition, Victoria blue (VB), Sirius red (SR), periodic acid Schiff (PAS) and Alcian blue (pH2.5) were used to stain elastic lamellae, collagen fibers and mucopolysaccharides (MPS). Contiguous sections from the same field were used for the histopathological, immunohistochemical and statistical analyses.

Fixed sections from the middle samples were washed and placed in Holt’s hypertonic gum sucrose to produce frozen sections. The samples were embedded in Tissue Mount (Chiba Medical, Saitama, Japan), before being frozen and stored at −80°C. The frozen sections were stained with Sudan black or oil red O to identify neutral fat. Frozen sections of normal livers collected from 2–21-year-old horses (the same horses from which the aortic arch specimens were obtained) were used as a positive control. Cystic medial necrosis in the tunica media was scored from 0 (no) to 3 (severe) according to the system described by Halushka et al. [2]. Medionecrosis (SMC nuclei loss) in the tunica media was scored from 0 (absent) to 3 (extensive) according to the system described by Halushka et al. [2].

Immunohistochemical examinations
Immunohistochemical examinations of paraffin-embedded sections were performed using the avidin-biotin-peroxidase complex (ABC) method (VECTASTAIN Elite ABC kit; Vector Laboratories, Burlingame, CA, U.S.A.). The primary antibody was a mouse monoclonal antibody against human alpha-smooth muscle actin (SMA) (clone 1A4, 1 in 100 dilution; Dako, Glostrup, Denmark). To remove endogenous peroxidase, the sections were immersed in 0.5% periodic acid solution at room temperature for 20 min. The sections were then incubated in primary antibody solution at 4°C for 14 hr. After being washed with phosphate-buffered saline, the sections were incubated in secondary antibody solution at room temperature for 30 min. After being incubated, the sections were reacted with ABC solution at room temperature for 30 min. Visualization was accomplished using 3′3-diaminobenzidine solution. The sections were counterstained with Mayer’s hematoxylin. Sections without the primary antibody were subjected to the same procedures as a negative control.

Imaging analysis
The thickness of the aortic tunica media was measured in 70 horses by examining sections in which elastin had been stained at magnifications of ×40 and ×100. The percentage thickness of the tunica media was calculated by dividing the thickness of the tunica media by the macroscopic thickness of the whole aorta. The percentage areas of elastin, collagen and SMC were determined based on image analysis. The results were used to calculate the percentage area of MPS. The measurements were obtained with a microscope by examining six randomly selected areas of the tunica media per slide at a magnification of ×400 using a Sony DXC-S500 digital camera (Sony Corporation, Tokyo, Japan). Six areas were selected for each of the three sections of the aorta. Three of the six areas contained the inner third of the tunica media, and the other three areas contained the outer third of the tunica media. The areas of elastin, collagen and SMC were measured as percentages of the total area and were calculated using image analysis software (ImageJ, U. S. National Institutes of Health, Bethesda, MD, U.S.A.) [11]. The percentage area of MPS tissue was calculated in 60 horses.
calculated by subtracting the percentage areas of the three components described above from the total area. A semi-quantitative evaluation of the actual area of each component was conducted based on the percentages of these four components and the thickness of the tunica media.

Statistical analysis

Statistical analyses of the 7 groups and comparisons between the proximal and distal sections were performed using the Tukey-Kramer test and F-test. P-values of <0.05 were considered to indicate statistical significance.

RESULTS

Gross findings

The luminal diameter and thickness of the aortic arch increased with age in all three examined sections (Fig. 1). The luminal diameter and thickness of the aortic arch differed significantly between group 1 and groups 2–7, and between group 2 and groups 3–7. However, the luminal diameter and thickness of the aortic arch did not differ significantly among the three sections (Fig. 1).

Histological findings

Thickness of the tunica intima and media (Fig. 2)

All percentages represent mean values for the relevant group.

In group 1, the tunica intima and media were 32 µm and 3,843 µm thick, respectively. The percentage thickness of the tunica media was 99.8%.

In group 2, the tunica intima and media were 49 µm and 5,216 µm thick, respectively. The percentage thickness of the tunica media was 99.1%.

In group 3, the tunica intima and media were 64 µm and 6,536 µm thick, respectively. The percentage thickness of the tunica media was 99.02%.

In group 4, the tunica intima and media were 61 µm and 7,296 µm thick, respectively. The percentage thickness of the tunica media was 99.2%.

In group 5, the tunica intima and media were 55 µm and 7,409 µm thick, respectively. The percentage thickness of the tunica media was 99.3%.

In group 6, the tunica intima and media were 62 µm and 6,620 µm thick, respectively. The percentage thickness of the tunica media was 99.1%.

In group 7, the tunica intima and media were 62 µm and 7,250 µm thick, respectively.

The mean percentage thickness of the tunica media was 99.14%. The tunica media accounted for most of the aortic wall (at least 98.7%; mean: 99.1%).
Histopathological morphometry

All percentages represent mean values for the relevant group (Figs. 3–8).

Group 1 exhibited the following specific percentage volumetric values: elastin, proximal: 12%, distal: 14%; collagen, proximal: 7%, distal: 12%; SMC, proximal: 21%, distal: 24%; and MPS, proximal: 60%, distal: 51%.

Group 2 displayed the following specific percentage volumetric values: elastin, proximal: 24%, distal: 25%; collagen, proximal: 12%, distal: 12%; SMC, proximal: 24%, distal: 27%; and MPS, proximal: 40%, distal: 36%.

Group 3 demonstrated the following specific percentage volumetric values: elastin, proximal: 29%, distal: 32%; collagen, proximal: 14%, distal: 16%; SMC, proximal: 22%, distal: 27%; and MPS, proximal: 35%, distal: 25%.

Group 4 displayed the following specific percentage volumetric values: elastin, proximal: 23%, distal: 27%; collagen, proximal: 13%, distal: 13%; SMC, proximal: 25%, distal: 30%; and MPS, proximal: 39%, distal: 30%.

Group 5 exhibited the following specific percentage volumetric values: elastin, proximal: 23%, distal: 27%; collagen, proximal: 16%, distal: 14%; SMC, proximal: 26%, distal: 34%; and MPS, proximal: 35%, distal: 30%.

Group 6 demonstrated the following specific percentage volumetric values: elastin, proximal: 17%, distal: 21%; collagen, proximal: 17%, distal: 17%; SMC, proximal: 25%, distal: 28%; and MPS, proximal: 41%, distal: 34%.

Group 7 displayed the following specific percentage volumetric values: elastin, proximal: 15%, distal: 17%; collagen, proximal: 29%, distal: 20%; SMC, proximal: 23%, distal: 32%; and MPS, proximal: 33%, distal: 31%.

Statistical analysis

Statistical comparisons among the 7 groups or between the proximal and distal samples (Figs. 4–6 and Table 2) revealed several significant differences ($P<0.05$). In the proximal region, the percentage area of elastin was significantly lower in group 1 than in groups 2, 3, 4, 5 and 6. In addition, it was significantly higher in group 2 than in groups 6 and 7. Similarly, it was significantly higher in group 3 than in groups 2, 4, 5, 6 and 7, and it was significantly higher in group 4 than in groups 6 and 7. In the distal region, the percentage area of elastin differed significantly between group 1 and groups 2–5, and between groups 2–4 and groups 6–7. The percentage area of collagen in the proximal and distal regions was significantly lower in group 1 than in groups 3 and 6, whereas it was higher in group 7 than in groups 1, 2, 4 and 5. In the proximal region, group 7 exhibited a significantly higher percentage area of collagen than groups 3 and 6. The percentage area of SMC did not differ significantly among the 7 groups. Only the percentage area of elastin in group 3 differed significantly between the proximal and distal regions.

Histopathological lesions

Histopathologically, some aortic lesions, including lesions exhibiting cystic medial necrosis (Fig. 9), elastin fragmentation, medionecrosis, focal calcification or ossification, were incidentally observed. The cystic medial necrosis lesions caused the accumulation of acidic MPS, which were strongly stained with Alcian blue and negatively stained with PAS. Cystic medial necrosis was commonly observed in all groups. In group 2, some small cystic lesions were seen, and the number and size of the lesions increased with age. These lesions were commonly found in the internal media, and exophytic growth was noted in the tunica media. The mean score for cystic medial necrosis increased with advancing age (group 1: 0.4; group 2: 0.64; group 3: 0.7; group 4: 1.1; group 5: 1.43; group 6: 1.38; and group 7: 2.8). The cystic medial necrosis score differed significantly between group 7 and the other groups, as well as between groups 1–3 and groups 5–6. Elastin fragmentation was observed in all horses according
CHANGES IN THE AORTA WITH AGE IN HORSES

Medionecrosis was commonly found in the middle region and was detected at one or two sites per slide. The mean score for medionecrosis did not differ significantly among the 7 groups (group 1: 0; group 2: 0.1; group 3: 0.2; group 4: 0.1; group 5: 0.7; group 6: 0.9; and group 7: 0.6). Focal calcification involving lamellar units was observed in the horses aged ≥17 years. Ossification was seen in a 29-year-old stallion. This lesion was more severe in the proximal region than in the other two regions. In this study, the cystic medial necrosis, elastin fragmentation and calcification lesions did not display any particular predisposition towards any of the three examined regions of the aorta. In addition, the samples containing these lesions were negatively stained with Sudan black and oil red O.

DISCUSSION

Gross examination

The luminal diameter and thickness of all of the examined sections of the aorta increased with age. This suggests that

![Fig. 3. Histological findings obtained with various stains. An 11-month-old horse: the findings obtained with VB (image A), SMA antibody (image C) and SR (image E). A 25-year-old horse: the findings obtained with VB (image B), SMA antibody (image D) and SR (image F). Bar, 100 µm](image-url)
thoroughbred horses exhibit a similar tendency to humans with regard to the age-related changes in their aortas [4, 6].

**Histological findings**

Proportional thickness of the tunica media

Histopathologically, the wall of the aortic arch was predominantly composed of the tunica media. No marked increases in intimal thickness were detected in the present study. This result differs from findings obtained in humans [22] and dogs [13], in which the thickness of the intima increased with age. Therefore, the risk of aortic rupture in thoroughbred horses might be more strongly related to medial lesions than intimal lesions.

**Histopathological morphometry and statistical analysis**

Histopathologically, the aortic media did not exhibit lipid or cholesterol crystal deposition, which have been reported previously [5]. For this reason, we classified the structural components of the aortic media into four categories, elastin, collagen, SMC and MPS. The percentage area of elastin was at least 10% in all age groups. Similarly, the percentage area of SMC was at least 25%
in all age groups. On the other hand, the percentage areas of elastin and SMC were significantly lower and significantly higher, respectively, in the thoracic aortas of Friesian horses that suffered aortic rupture than in the thoracic aortas of control Friesian horses [15]. These findings suggest that specific proportions of elastin and SMC are important for aortic stability. The percentage area of elastin was lowest in group 1. The percentage area of elastin increased from group 1 to group 3 and decreased from groups 3 to group 7. Elastin is only synthesized during the early stages of life and progressively deteriorates with aging [8, 17]. In thoroughbred horses, elastin is synthesized in the first year of life, and the amount of elastin progressively decreases with advancing age. The percentage area of collagen was lowest in group 1. The percentage area of collagen increased with advancing age and peaked in group 7. Increasing collagen density might be associated with hypertension [12] or arterial rupturing/aneurysms

Table 2. The significance of the intergroup differences in the percentage area of elastin

| Group | 1 | 2 | 3 | 4 |
|-------|---|---|---|---|
| 2     |   | ++ |   |   |
| 3     |   | ++ | a |   |
| 4     | **| * | * |   |
| 5     | **| * | * |   |
| 6     | * | * | **| **|
| 7     |   | **| **| **|

a)**: Both the proximal and distal sections displayed significant age-related differences. b)**: Only the proximal section exhibited significant age-related differences (P<0.05).

Fig. 8. Areas of the four components of the tunica media in the proximal region. This graph is based on the results shown in Figs. 2 and 7.

Fig. 9. The internal media in the proximal region. Cystic medial necrosis resulted in the accumulation of acidic MPS, which were strongly stained with Alcian blue (arrowheads). The image shows moderate grade (score 2) cystic medial necrosis in an 18-month-old (1.5-year-old) horse. The section was stained with PAS and Alcian blue (pH2.5).
[20]. The blood pressure values of thoroughbred horses are generally higher than those of other breeds of horse [7]. In our study, collagen deposition was considered to be an aging-related physiological reaction. The percentage area of SMC was lowest in group 1 and slowly increased with advancing age. Nevertheless, the change in the percentage area of SMC was the smallest such change seen among the four examined structural components. Furthermore, the percentage area of SMC did not differ significantly among the 7 groups. On the other hand, the proportion of the aortic media accounted for by SMC (Fig. 8) increased with age. This indicates that SMC grow with age. In addition, SMC hypertrophy was detected during histopathological examinations. The percentage area of MPS was largest in group 1. The percentage area of MPS was inversely proportional to the percentage area of elastin. This study is the first to examine the age-related changes in the percentage area of MPS in the equine aortic media. The age-related changes in the percentage area of MPS observed in this study were similar to the changes in the human interstitial ratio described in a study of newborns and children [10].

**Histopathological lesions**

Medionecrosis was only observed in aged horses. Some previous reports have suggested that medionecrosis might predispose the affected vessels to rupture [14, 21]. In Friesian and Dutch halfblood horses, aortic rupturing typically occurs near the ligamentum arteriosum, where medionecrosis is most commonly found [21]. In thoroughbred horses, such rupturing commonly occurs near the junction with the heart [3]. In thoroughbred stallions that had suffered aortic rupture, medionecrosis was found underneath the rupture sites. In addition, the aortic rupturing seen in these cases was considered to have been caused by marked increases in blood pressure during breeding [14, 15]. In our study, medionecrosis was more commonly found in the middle sections near the ligamentum arteriosum than the proximal sections. Therefore, the findings of this study suggest that in thoroughbred horses, rupturing of the aorta is not only caused by medionecrosis. MPS deposition is generally reported to occur in cystic medial necrosis lesions. Cystic medial necrosis has been suggested to be an underlying cause of aortic rupturing [9, 14, 19]; however, other studies have indicated that it is related to physiological activity [13, 18]. In our study, cystic medial necrosis was first found in a 234-day-old (embryonic) horse. These lesions increased in number, size and score with aging. In young horses, the cystic medial necrosis was confined to the internal media, whereas in the elderly horses, cystic lesions were found from the internal to middle media. Our findings indicate that the amount of MPS does not affect the risk of aortic rupturing because the largest amount of MPS was found in group 1. Therefore, it is suggested that the localization of MPS (focal deposition) is more important than the quantity of MPS in terms of the risk of aortic rupture. Thus, cystic medial necrosis might be an underlying cause of aortic rupturing, but such rupturing might be caused by multiple defects.

In conclusion, our study revealed that the medial histological components of the aortic arch undergo changes, including a reduction in the amount of elastin and increases in the amounts of collagen and MPS, with advancing age in thoroughbred horses. These changes are similar to the changes described in a previous report [15], which detected a reduction in the amount of elastin and an increase in the amount of collagen. However, none of the horses in the latter study suffered aortic rupture. Therefore, morphometric changes occur in the aortic media as part of the normal aging process in thoroughbred horses. As the percentage area of each component of the aorta varies according to age, it is necessary to use horses of the same age when performing comparisons of the tunica media in the aortic arch.

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