Macroscopic and Histological Variations in the Cellular Tapetum in Dogs

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ABSTRACT: We aimed to document macroscopic variations in the cellular tapetum in the dog, to provide a histologic description of the macroscopic results and to evaluate the correlation between the macroscopic appearance and aging. Fifty three dogs including 5 beagles, 1 Chihuahua and 47 mixed breeds of each gender were used. For a macroscopic study, the fresh tapetal fundi were photographed using digital camera. For a histological study, the glutaraldehyde-formalin fixed eyes were embedded in nitrocellulose and stained with hematoxylin-eosin or thionine. The normal tapetum was triangular with the rounded angles and the smooth contour. The atypical tapetum was smaller and more variable in shape, contour and color than the normal one. In severe cases, the fundus was devoid of the tapetum. The atypical tapetum tended to increase in frequency with aging. Retinal pigment epithelial cells on the normal tapetum were unpigmented. In the eye with the atypical tapetum, regardless of tapetal size and shape, unpigmented retinal pigment epithelial cells showed a similar distribution to that on the normal tapetum, even in a dog without a tapetum. Although there is a congenitally hypoplastic tapetum, the atypical tapetum tends to increase in incidence and severity with aging.

KEYWORDS: aging, canine, cellular tapetum, retinal pigment epithelial layer

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Eyeshine in a number of vertebrates including carnivores is a phenomenon of light reflection from the tapetum lucidum of the eyes. The tapetum is thought to reflect light back to the retina, thus enhancing scotopic sensitivity. It is interposed between the choroid capillary layer beneath the retinal pigment epithelial layer (RPEL) and the choroidal proper substance. In the tapetal area, the RPEL lacks pigment so that the tapetum can be clearly seen through the retina [15]. A cellular tapetum is characteristic of carnivores and composed of multiple layers of cells arranged in a brick-like structure. The most striking feature of tapetal cells in the dog as observed by electron microscopy is the presence of numerous slender, electron-dense rods in a regular array in the cytoplasm [13]. The tapetum of the dog eye is triangular in outline with the base of the triangle being ventral. The tapetum occupies the upper half of the ocular fundus. The form and color of the tapetum are variable in dogs [4, 9, 11, 16, 26]. In general, small-sized breeds including the Yorkshire Terrier and Chihuahua have a significantly smaller tapetal area. Some dogs congenitally lack the tapetum entirely, especially in Labrador retrievers and beagles of certain families. Furthermore, the tapetum is small in area in the subalbinotic fundus and completely absent in the albinotic fundus [1, 2, 9–11, 16]. Wyman and Donovan [26] state that beadiness refers to the mosaic pattern observed at the periphery of the tapetum and is most easily seen near the junction of the tapetum and the non-tapetal area. The degree of beadiness increases with the age, a phenomenon that is more apparent in small dog breeds. These previous studies have provided little histological examination of variability in the tapetum.

In this study, we aimed to document macroscopic variations in the canine tapetum, to provide a histologic description of the macroscopic results and to evaluate the correlation between the macroscopic appearance and aging.

MATERIALS AND METHODS

Eyes were obtained from 53 dogs including 5 beagles (2–3 years old), 1 Chihuahua (10 years old) and 47 mixed breeds (1 to 13 years old) of each gender that were euthanized by carbon dioxide overdose (47 mixed breeds) or deep anesthesia using pentobarbital sodium (100 mg/kg, i.p.) (the other 6 dogs) for an unrelated experiment to the eye by other group in our faculty. Both eyes of all animals were removed within 2 hr of death. For a macroscopic study (all cases), the unilateral eyes were cut along the equatorial plane, and the tapetal fundus was photographed by digital camera (EX-Z400, Casio, Tokyo, Japan). For a histological study (20 cases including 5 beagles, 1 Chihuahua and 18 mixed breeds), following macroscopic observations, the unilateral eyes were injected with 3% glutaraldehyde in 10% formalin into the vitreous cavity. After fixation, the posterior cup of each eye was dehydrated, embedded in nitrocellulose (Collodion, Wako, Osaka, Japan), cut at 30–40 µm and then stained with hematoxylin-eosin or thionine. In 5 beagles, 40-µm sections of tapetum with the retina and choroid were cut at 0.5 to 2 mm intervals and reconstructed after measurement of the
tapeptum thickness and observations of the RPEL. These procedures were carried out in accordance with the guidelines of the Ethical Committee of Tottori University, Japan.

RESULTS

Macroscopic observations: The normal tapeptum was triangular with rounded angles and smooth contour. Its base had usually contact with the optic nerve disc (OD), but separated from the OD in rare cases. Its apex located dorsally to the OD. The nasal half was longer than the temporal half in width. Thus, the nasal angle of the tapeptum was more acute than the temporal angle. The color tone of the tapeptum was not even; its center was brightest (Fig. 1). Based on histological observation, the color tone of the tapeptum reflected the thickness of the tapectal tissue.

The atypical tapeptum was smaller and more variable in shape, contour and color than the normal one (Fig. 2). The atypical tapeptum disappeared from the periphery, especially from the nasal and ventral parts. In mild cases, the atypical tapeptum was irregular in contour and faded in the nasal part (Fig. 2a). In severe cases (the Chihuahua), the fundus was devoid of the tapeptum and had a reddish-brown area that was similar in both location and extent to the normal tapeptum (Fig. 2d).

Bilateral differences in appearance of the tapeptum were few through the normal and atypical tapepta.

Regarding the relationship of the atypical tapeptum to age, the occurrence ratio was 31% in the atypical tapeptum and 69% in the normal tapeptum in dogs under age 5 (35 cases) and was 72% in the atypical tapeptum and 28% in the normal tapeptum in dogs over age 6 (18 cases). The tapeptum tended to be atypical with increasing age. Furthermore, the atypical tapeptum tended to increase in severity with aging.

Histological observations: In the normal tapeptum, the RPEL was almost unpigmented on the tapeptum (Fig. 3). The narrow peripheral zone of the tapeptum was covered by the lightly pigmented RPEL. Thus, the distribution of tapectal tissue closely corresponded to the area of the macroscopic tapeptum. The tapeptum was thick in its central part, and its thickest part was located dorso-temporally to the OD (Fig. 4). The thickest part of the typical tapeptum varied in thickness, from 20 to 70 µm (average: 53 µm, standard deviation: 17 µm) and consisted of maximum 20 layers of tapectal cell.

In the atypical cases, in which the tapeptum was markedly small and very irregular in contour, the tapeptum was very thin. In these cases, the choroid was fully pigmented, the same as the normal choroid. Regardless of the shape of the atypical tapeptum, the unpigmented RPEL showed a similar extent to that in the normal tapeptum. Pigmentation of the choroid and pigmented RPEL was normal in all cases (Fig. 5). While the Chihuahua dog macroscopically lacked the tapeptum, unpigmented RPEL showed histologically a similar distribution to that in the normal tapeptum (Fig. 6a). Tapectal tissue was observed in a fragmentary fashion (Fig. 6b).

DISCUSSION

A wide variation in extent and color of the tapeptum has been reported in normal dogs [9, 11, 16]. The tapeptum is poorly developed in the toy breeds including the Chihuahua and in dogs with a merle coat color [11, 16]. Pigmentation of the choroid correlates closely with the development of the tapeptum [10, 11]. Interestingly, the albino ferret and the Siamese cat lack choroidal pigmentation, but have the tapeptum [22, 23]. According to Thibos et al. [22], the choroid in the Siamese cat contains melanocytes, but is defective in pigment production. Thus, the Siamese is completely free from pigment in the choroid but has the tapeptum, because tapectal cells derive from neural crest cells as well as choroidal melanocytes. The blue-eyed white cat has no tapeptum and no choroidal pigmentation, because neural crest cells do not migrate into the choroid. All cases observed in this study were normal in pigmentation of the choroid. In this study, unpigmented RPEL was equally distributed in the dog retina regardless of the extent and shape of the tapectal area. In the Chihuahua, the tapeptum was not found, but the unpigmented RPEL was similar in distribution to that of the normal tapeptum.

The canine tapeptum develops after birth [11, 17]. The developmental relation of the unpigmented RPEL to the tapeptum has not been studied in dog. According to a study of the bovine eye, the adult pattern of pigmented RPEL in nontapetal zones and unpigmented RPEL in the presumptive tapectal zone is well established in the early stage of pregnancy [7]. In other word, the extent of the unpigmented RPEL is decided before the development of the tapeptum. Thus, compartmentalization of unpigmented and pigmented RPEL would not be directly related to the development or
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In this study, the atypical tapetum tended to increase in frequency with aging. Histologically, tapetal layers decreased in number throughout the atypical tapetum. Therefore, thin tapetal parts, namely the periphery of the tapetum, disappeared first, and the contour of the tapetum became irregular with shrinkage of the tapetum. According to Wyman and Donovan [26], the degree of beadiness of the tapetum increases with aging. Beadiness refers to the mosaic pattern observed in ophthalmoscopic examination at the periphery of the tapetum. Beadiness resulted from the irregular contour of the tapetum in this study. Thus, it is suggested that the tapetum in the dog tends to decrease in size macroscopically with aging as the number of tapetal cell layers decreases due to tapetal cell degeneration.

The atypical tapetum was found at high rates, especially in aged dogs. It is speculated that the atypical tapetum results from tapetal cell degeneration in addition to congenital hypoplasia of the tapetum. For example, smaller-sized breeds have significantly smaller tapetal area [9]. In this study, the normal tapetum had a wide variety of thickness (53 ± 17 µm). The thin normal tapetum would be frequently abnormal at a relatively young age. It has been suggested that tapetal cells may be vulnerable. Several studies have shown selective tapetal cell degeneration following administration of drugs, including a beta-adrenergic blocking agent SCH 19927 [20], a macroride antibiotic (rosaramicin) [14], an aziride antibiotics (CP 62,993) [8], an aromatase inhibitor (CGS 14796C) [21] and an antipsychotic agent (1192U90) [5]. Furthermore, tapetal cell degeneration may result from malnutrition. Tapetal cells in the dog are packed with long slender rods [13], which are rich in zinc [12, 24, 25]. Zinc is essential for preservation of the structure of normal tapetal rods, because the congenitally abnormal tapetum which lacks zinc in its tapetal rods is not visible in ophthalmoscopy and shows an absence of the regular arrays of tapetal rods [25]. Drugs that chelate zinc cause a reversible tapetal...
decoloration in dogs [8, 19]. Even in human in developed countries, zinc deficiency is common in the elderly, and zinc supplementation delays the progression of age-related macular degeneration and reduces the risk of loss of vision [18]. Zinc deficiency can affect most organ systems in dogs as well [3]. If zinc deficiency is not uncommon in elderly dogs, tapetal cells may gradually degenerate for zinc deficiency with aging. In addition, the tapetum contains very large amount of cysteine [24]. Cysteine, to which zinc readily binds, is necessary for the biosynthesis of glutathione, which is quantitatively the most important antioxidant and radical scavenger. The age-related decrease in the body’s cysteine and/or glutathione level plays a causative role in various ageing related degenerative processes [6]. As the age-related decrease in zinc and cysteine may synergistically increase tapetal cell degeneration, there may be a trend toward an increase of tapetal cell degeneration with aging.

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