Angioid streaks, clinical course, complications, and current therapeutic management

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Abstract: Angioid streaks are visible irregular crack-like dehiscences in Bruch’s membrane that are associated with atrophic degeneration of the overlying retinal pigmented epithelium. Angioid streaks may be associated with pseudoxanthoma elasticum, Paget’s disease, sickle-cell anemia, acromegaly, Ehlers–Danlos syndrome, and diabetes mellitus, but also appear in patients without any systemic disease. Patients with angioid streaks are generally asymptomatic, unless the lesions extend towards the foveola or develop complications such as traumatic Bruch’s membrane rupture or macular choroidal neovascularization (CNV). The visual prognosis in patients with CNV secondary to angioid streaks if untreated, is poor and most treatment modalities, until recently, have failed to limit the devastating impact of CNV in central vision. However, it is likely that treatment with antivascular endothelial growth factor, especially in treatment-naive eyes to yield favorable results in the future and this has to be investigated in future studies.

Keywords: angioid streaks, pseudoxanthoma elasticum, choroidal neovascularization

Introduction
Angioid streaks were initially reported in 1889 by Doyne. They were described as “irregular radial lines spreading from the optic nerve head to the retinal periphery” in a patient who had retinal hemorrhages secondary to trauma. Knapp first coined the term “angioid streaks” in 1892 because their appearance suggested a vascular origin. Not until 1917 did Kofler correctly determine that angioid streaks represented changes at the level of Bruch’s membrane. Clinical examination with subsequent histopathological findings by Bock in 1938 in two patients with pseudoxanthoma elasticum confirmed that the underlying abnormality was not vascular in nature but rather a structural alteration in Bruch’s membrane. A few years later similar histopathological results were found in patients suffering pseudoxanthoma elasticum, Paget’s disease, but also from systemic diseases. Despite the fact that many systemic diseases like acromegaly, Ehlers–Danlos syndrome and diabetes mellitus, have been associated with angioid streaks, the most common diseases related to angioid streaks are pseudoxanthoma elasticum, Paget’s disease, bone and sickle-cell anemia.

Histopathological findings
Angioid streaks represent visible irregular crack-like dehiscences in Bruch’s membrane that are associated with atrophic degeneration of the overlying retinal pigmented epithelium (RPE). Histopathology in patients with angioid streaks that suffered from pseudoxanthoma elasticum demonstrated calcium deposition in Bruch’s membrane which has several well demarcated breaks. In pseudoxanthoma elasticum, the primary lesion is the degeneration of elastic fibers of the connective tissue of the organism, while the calcium deposition represents a secondary disorder of unknown origin. In angioid streaks the elastic lamina that occupies the midsegment of Bruch’s membrane is affected, resulting in disintegration and frying of the elastic fibers.
Electron microscopy (EM) studies showed the presence of abundant granulomatous material in this lamina, a fact that supports the theory of pathologic elastic fibers’ production.

In Paget’s disease the bone deformities lead to calcium binding by the elastic fibers.12

For several years the appearance of angioid streaks in sickle-cell hemoglobinopathies was attributed to high level of serum iron.5,13,14 However, other types of anemias with increased iron levels in blood are not associated with angioid streaks. In addition, histochemical and EM studies, which took place in the eyes of patients with homozygotic sickle-cell anemia demonstrated severe tissue calcification;15 this fact favors the hypothesis that angioid streaks in patients with sickle-cell hemoglobinopathies are correlated to calcium deposition at Bruch’s membrane.

Other histopathological findings include the break or absence of choriocapillaris beneath angioid streaks and thinning or decoloration of RPE;16 these changes are identical in angioid streaks despite different underlying systemic diseases.

**Ocular manifestations**

Angioid streaks are mainly asymptomatic. The appearance of symptoms occurs when the angioid streaks involve the foveola or in case of choroidal neovascularization (CNV) in the macular region (Figures 1, 2).

Angioid streaks have a typical appearance as narrow, jagged lines deep to the retina, almost always bilaterally. They radiate out in a cruciate pattern from an area of peripapillary pigment alterations. They may circumferentially ring the peripapillary area as well (Figure 1; Figure 2). Angioid streaks are evident in fundoscopy a few millimetres from the optic disc and rarely occur in the periphery of the posterior pole. Clinical diagnosis is usually straightforward. Angioid streaks have a thickness of 50–500 μm and are visible in fundoscopy under the retinal vessels.11,15,17,18

The color of angioid streaks depends on the background coloration of the fundus and the degree of the atrophy of the overlying RPE. Thus, angioid streaks are red in light-colored individuals, while in patients who have darker background pigmentation, they are usually medium to dark brown. Angioid streaks become darker as time passes by and at the same time discoloration of RPE occurs. Sometimes angioid streaks are extremely dark and have several bonds between them giving the appearance of a ‘spider’s web’ in the retina. In other occasions a fibrous connective tissue develops around angioid streaks which appear obscure and light-colored.11,17,18

In cases where angioid streaks are confined to macula and especially if the foveolar avascular zone is affected, patients develop metamorphopsia and reduced visual acuity. In contrary, if the fovea is unaffected, patients remain...
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Table 1 Systemic conditions associated with angioid streaks

- Hemochromatosis
- Acromegaly
- Diabetes mellitus
- Sickle-cell hemoglobinopathies
- Pseudoxanthoma elasticum
- Acquired hemolytic anaemia
- Hereditary spherocytosis
- Myopia
- Neurofibromatosis
- Paget’s disease
- Ehlers–Danlos syndrome
- Sturge–Weber syndrome
- Hyperphosphatemia

asymptomatic and angioid streaks are an accidental finding during a routine ophthalmological investigation.

Fluorescein angiography, indocyanine green angiography

The photographs taken with the filters of fluorescein angiography before the intravenous administration of the contrast material (fluorescein dye) provide the physician with important information because angioid streaks frequently present the phenomenon of auto-illumination. Also, optic disc drusen which are often associated with angioid streaks show auto-illumination in fluoroangiography.11,19

Typically, angioid streaks have a ‘window defect’ in fluorescein angiography due to atrophy of RPE adjacent to them. Leakage of fluorescein is evident when CNV is present. The diagnosis of angioid streaks is usually made on the basis of fundoscopy, but intravenous fluorescein angiography can help to delineate the presence of the disease when the ophthalmoscopic appearance is subtle (Figure 3).20

Indocyanine Green (ICG) angiography is a useful diagnostic tool for angioid streaks only in the rare case that fundoscopy and fluorescein angiography can not confirm the diagnosis. Such occasions involve the severe lesions of RPE which cause hyperfluorescent lines that are obscure or the development of macular CNV; in these cases ICG angiography demonstrates the neovascular membrane more clearly than fluorescein angiography. ICG angiography shows hyperfluorescent lines with ‘pinpoints’ over their whole length that are larger and more numerous than those on fluorescein angiography or red-free photography (Figure 4). On the contrary, recently developed angioid streaks become evident only at the late stage of the examination and have the appearance of hypofluorescent linear distortions around optic nerve head or posterior pole.11,18

Clinical course, complications

Patients with angioid streaks are generally asymptomatic, unless the lesions extend towards the foveola or develop complications such as traumatic Bruch’s membrane rupture or macular CNV (Figures 5, 6).

The increase in length and width of angioid streaks is considered as an expected feature of the disease, but there are no clinical studies determining the rate of their propagation in correlation with time. Increase in length results in lower visual acuity if the angioid streaks spread to the macular region and particularly in the foveolar avascular zone.

Patients with angioid streaks may develop breaks of the Bruch’s membrane even after relatively mild head injuries (traumatic dehiscences in Bruch’s membrane), since their Bruch’s membrane is brittle.11 A retrospective study on patients with angioid streaks mentioned that 15% of those who suffered a head injury developed significant visual impairment.21 Traumatic breaks of Bruch’s membrane are frequently followed by subretinal hemorrhages which can be easily misinterpreted as CNV. These hemorrhages appear usually next to the angioid streaks and sometimes disseminate into the macula.21,22

Choroidal neovascularization

The commonest and most serious complication of angioid streaks is CNV in the macular region. Such a complication deteriorates dramatically the prognosis of angioid streaks because it is one of the most difficult clinical entities an
ophthalmologist can encounter. Patients with angioid streaks who develop CNV (Figure 6) are symptomatic and their main symptoms are metamorphopsias and reduction of vision.

Fluorescein angiography confirms the presence of classic CNV in most cases and helps define the margins of the neovascular membrane, which is typically above or right next to the angioid streaks (Figures 7, 8). In some occasions it is difficult to determine the borders of the neovascular membrane in fluorescein angiography due to adjacent hemorrhages or RPE lesions. In these cases ICG angiography is an invaluable diagnostic tool.

The incidence of CNV in patients with angioid streaks varies between 72%–86% in numerous studies. Commonly, neovascularization involves both eyes, but does not occur simultaneously; there is an interval of roughly 18 months.2,6,11,21,23

The risk of developing CNV increases with age.24 Other risk factors comprise the width, length and location of the angioid streaks. The wider and longer are the angioid streaks the higher the risk for CNV and especially if the lesions are located in a distance less than one optic disc diameter from the foveola.25,26

Angioid streaks associated with pseudoxanthoma elasticum have a relatively high probability of developing macular CNV;24 the opposite happens in patients suffering sickle-cell anemia.27,28

The standard outcome is poor if CNV in the macular region remains untreated because it leads to further extensive formation of subfoveal scarring of CNV causing severe deterioration of visual acuity.20 More than 50% of such patients eventually become legally blind since their visual acuity is less than 1/20.29

Treatment

Laser photocoagulation

The incidence of CNV in patients with angioid streaks varies between 72%–86% in different studies. Normally, neovascularization involves both eyes, but does not occur simultaneously.

The prophylactic therapy of angioid streaks with laser beams in order to avoid the development of CNV has been used before,30 but it is no longer recommended and many researchers strongly believe that this kind of intervention can actually induce CNV.31

Photocoagulation with thermal laser was regarded as the only possible therapeutic modality for macular CNV and consequently was introduced in angioid streaks. Clarkson and
colleagues\textsuperscript{12} presented a case series report of six patients with
angioid streaks and CNV of the macular region, who were
treated with a thermal laser directly targeting the neovascular
membrane. Their results were devastating and all patients
developed further expansion of the neovascularization
causing loss of central vision. At the same time, other small
case series reports stated that laser photocoagulation had
some encouraging results in neovascularization outside the
foveolar region.\textsuperscript{32,34}

In 1988, Gelisken and colleagues\textsuperscript{35} presented the results of
a study involving 30 eyes with CNV due to angioid streaks.
All patients were treated with green argon or krypton laser
and the follow-up was two months to 16 years. The authors
concluded that the eyes which had extrafoveolar neovascular
membranes benefitted from the application of thermal laser
since they retained a useful vision compared to the eyes that
remained untreated. Additionally, they stressed that in cases
of subfoveal CNV, no treatment should be applied. Finally,
they suggested that no statistically significant difference
was observed in using argon and krypton laser, but they
preferred the latter.

During the last 20 years, several clinicians drew the same
conclusion from these results and thus the dominant con-
temporary theory is that the efficacy of photocoagulation for
macular CNV in angioid streaks is of limited application due
to the high percentage of neovascular membrane relapse.\textsuperscript{36}

Transpupillary thermotherapy
Recently, ophthalmologists’ interest has switched towards laser
treatment with reduced use of energy and such an application
is the diode laser using beams of 810 nm length. Such a laser
beam has better penetration through the transparent media of
the eye, better action and well-controlled thermal effect caus-
ing less absorption by the RPE and deeper penetration in the
choriocapillaris.\textsuperscript{37,38}

Transpupillary thermotherapy uses a diode laser at a
lower threshold that does not cause thermal burn and has been
applied for the treatment of macular CNV of any origin. Aras
and colleagues\textsuperscript{37} tried this method in patients with subfoveal
neovascular membranes in angioid streaks and concluded that
it does not seem to affect the course of the disease and at the
same time they observed a spreading out of the borders and
the leakage of the membrane.

Macular translocation surgery
This surgical technique was introduced by Machemer and
Steinhorst\textsuperscript{39} in 1993. Macular translocation involves moving
the neurosensitive retina (macula and varying amounts of
adjacent retina) to a new location, away from the ingrowth of
the new vessels. This may be accomplished by limited trans-
location, in which a limited retinal detachment is made and
the scleral wall is shortened by imbrication or out-pouching,
or there may be a 360-degree retinotomy, with a rotation of
the entire retina. This operation is followed by a strabismus
surgery. Since the first operation was performed there are
several modifications by many vitreoretinal surgeons with
encouraging results.\textsuperscript{40–43} However, we should stress that it is
a complex, difficult, long-lasting operation which has serious
complications (retinal detachment, proliferative vitreoretin-
opathy, endophthalmitis, etc.) threatening the central and
peripheral vision of the patients.

Macular translocation surgery was used in macular CNV
in angioid streaks. Roth and colleagues\textsuperscript{44} described a case
where they performed a successful lower macular translo-
cation followed by laser photocoagulation in the region of
choroidal neovascular membrane in a patient with angioid
streaks. The end result was encouraging and other surgeons
had similarly good results.\textsuperscript{45,46} It should be mentioned that
these encouraging results can not be fully evaluated since
the number of patients and the studies involved are of a very
small number.

Photodynamic therapy
Large randomized clinical trials were performed to evaluate
the efficacy of photodynamic therapy (PDT) with verteporfin
for CNV secondary to age-related macular degeneration
(AMD) and pathologic myopia.\textsuperscript{47–49} Since the results of these
studies were promising, clinician tried to use PDT in other
pathological entities causing CNV.

Figure 7 Fluorescein angiography of left eye with angioid streaks complicated by
choroidal neovascularization.
In 2000, the Archives of Ophthalmology published the first results of the application of PDT for CNV not caused by macular degeneration. In this study group there was one patient with angioid streaks. He was a 55-year-old man who was treated only once with PDT and had a 12-month follow up. During the follow up time there was no further deterioration of his visual acuity, but at the same time the leakage of fluorescein dye from the neovascular membrane remained unaffected.

Two years later, Karacorlu and colleagues from the 'Istanbul Retina Institute' in Turkey published their results from eight patients with angioid streaks who had PDT. None of the patients developed reduction of the visual acuity and the average improvement of visual acuity was 1.37 lines on the Snellen chart. Three patients had fluorescein leak from the neovascular membrane during their last visit. Based on these results the authors of the article concluded that PDT plays an important role in the management of CNV in angioid streaks, but they stressed that more studies with larger number of patients and longer follow up should be done in order to evaluate accurately the efficiency of PDT in the treatment of macular CNV in angioid streaks.

A year later, Shaikh and colleagues presented their results in the same journal. Their study encompassed 11 eyes from nine patients with angioid streaks who had PDT. None of the patients developed reduction of the visual acuity and the average improvement of visual acuity was 1.37 lines on the Snellen chart. Three patients had fluorescein leak from the neovascular membrane during their last visit. Based on these results the authors of the article concluded that PDT plays an important role in the management of CNV in angioid streaks, but they stressed that more studies with larger number of patients and longer follow up should be done in order to evaluate accurately the efficiency of PDT in the treatment of macular CNV in angioid streaks.

A year later, Arias and colleagues reported on PDT in angioid streaks relatively recently. Ten patients (10 eyes) with pseudoxanthoma elasticum had PDT with an average 18-month follow up. The results were appalling since only three patients retained the initial visual acuity, while four patients had dramatic deterioration of visual function (more than six lines on the Snellen chart). Based on their finding, they concluded that...
that PDT is not effective in the treatment of macular CNV with the background of angioid streaks.

Ladas and colleagues\textsuperscript{38} evaluated the effectiveness of conventional PDT in a series of 24 eyes of 22 patients with CNV due to angioid streaks and compared it to the effectiveness of a PDT modification where retreatments were performed earlier (every eight weeks instead of 12).

At the end of the follow-up, final best-corrected visual acuity decreased in 21 of the total 24 eyes and In 19 eyes final best-corrected visual acuity was equal to or less than 20/400. There were not any statistically significant differences in final visual acuity between the two groups and the authors concluded that the functional and the anatomic results of PDT were not satisfactory, even when retreatments were performed earlier than the conventional time of three months.

In conclusion, from all previous studies and case reports it is evident that, despite the initial encouraging results from the application of PDT for the treatment of CNV in the macula, the end results did not fulfill the initial expectations. In addition, several studies present contradictory results and others consider PDT as an adjuvant therapy that does not prevent, but slows down the natural course of CNV.

### Antivascular endothelial growth factor treatment

Anti-vascular endothelial growth factor (VEGF) treatment has resulted in unprecedented visual and anatomic outcomes far outpacing other available treatments for CNV due to AMD.\textsuperscript{59–61} Today physicians and patients can expect visual stabilization in most patients and visual improvement in many, particularly if treatment is begun early in the course of the disease. PDT in combination with anti-VEGF has been also used for the treatment of CNV.\textsuperscript{62,63}

Ranibizumab\textsuperscript{64} (Lucentis) is a recombinant humanized immunoglobulin G1 and isotype monoclonal antibody fragment designed for intraocular use which binds to and inhibits the biologic activity of human VEGF A.

Bevacizumab,\textsuperscript{65} is a recombinant humanized full-length antibody that binds to all isoforms of VEGF, similar to ranibizumab which has been offered as an off-label intravitreal application for the treatment of wet AMD.

Wecke and colleagues\textsuperscript{66} reported favorable result in a patient with CNV due to angioid streaks after intravitreal injection of bevacizumab.

Chang and colleagues\textsuperscript{67} reported their results of intravitreous injection of bevacizumab for CNV from other causes than AMD and among them 11 patients suffered from angioid streaks. The CNV responded well to bevacizumab injections, however, as the authors stated, in between these eyes there was a high proportion of eyes that had previously undergone PDT so the results may have been biased.

Recently Donati and colleagues\textsuperscript{68} reported the use of intravitreal injection of bevacizumab for CNV in six eyes of five patients with angioid streaks. The authors concluded that bevacizumab may be useful in the treatment of CNV due to angioid streaks. However, their results may have been compromised by the fact that all eyes had previously undergone PDT or laser photocoagulation.

Schiano-Lomoriello and colleagues\textsuperscript{69} reported two patients with CNV secondary to angioid streaks who received three intravitreal injections of bevacizumab and followed them for one year. The authors concluded that intravitreal injections of bevacizumab appeared to be an effective and safe treatment for CNV and resulted in a long-term CNV inactivation.

The results from 11 and six patients suffering from CNV associated with angioid streaks were reported recently by Neri and colleagues\textsuperscript{70} and Wiegand and colleagues,\textsuperscript{71} respectively. In both studies, intravitreal bevacizumab was found to mildly reduce central foveal thickness and stabilize visual acuity. Both studies concluded that intravitreal bevacizumab may be a promising treatment. Future studies are required to validate their findings.

In conclusion, the visual prognosis in patients with CNV secondary to angioid streaks, if untreated, is poor and most treatment modalities until recently have failed to limit the devastating impact of CNV in central vision.

However, it is likely that treatment with anti-VEGF, especially in treatment-naive eyes would yield favorable results and this has to be investigated by future studies.

### Disclosure

The authors report no conflicts of interest in this work.

### References


