

(AMD) is a deterioration of the retina (a thin layer of lightsensitive nerve tissue that lines the back of the eye) and the *choroid* (the layer of blood vessels and connective tissue between the retina and the white of the eye).

AMD may lead to substantial loss in visual sharpness or *acuity*. Yet, many people with AMD maintain reading vision throughout their lives. AMD is the leading cause of significant visual acuity loss in people over age 50 in developed countries.

Causes: The exact cause of AMD is unknown, but the condition develops as the eye ages. There are 2 types of AMD: non-neovascular or dry AMD in 85% of affected individuals; and neovascular or wet AMD in 15%.

Dry AMD can be further divided into early, intermediate, and late stages. In early stages of dry AMD, small drusen—pale-yellow lesions—form beneath the retina (Figure 1A). Drusen are usually harmless, but as they accumulate, dry AMD can progress. Pigmentary changes can occur along with drusen.

Intermediate dry AMD is characterized by larger drusen (*Figure 1B*). Areas of atrophy or wasting also may develop in the retina; if the atrophic area is significant and has sharp borders, it constitutes the late stage of dry AMD and is termed *geographic atrophy* (GA).



Figure 1A. In early stages of dry AMD, small *drusen*—pale-yellow lesions—form beneath the retina. Image by Jerald A. Bovino, MD, FASRS. Retina Image Bank; Image 6374. © American Society of Retina Specialists.

Figure 1B. Intermediate dry AMD is characterized by larger drusen. Photo by Brandon G. Busbee,

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The Foundat

SYMPTOMS

Early stages of dry AMD may not show any symptoms. In intermediate dry-AMD stages, patients may have difficulty reading in dim light or transitioning from light to dark; they may also notice a decrease in the intensity or brightness of colors, and some early distortion (warping) of straight lines.

In the more advanced GA stage, individuals experience more marked distortion of straight lines, difficulty reading and driving at night, gradual loss of central vision, or dark, blurry areas in the center of vision.

In RPD, patients also note decreased vision in dimly lit settings, and may need longer to adjust from bright light to dim light.



THE RETINA is a thin layer of light-sensitive nerve tissue that lines the back of the eye (or vitreous) cavity. When light enters the eye, it passes through the iris to the retina where images are focused and converted to electrical impulses that are carried by the optic nerve to the brain resulting in sight.

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GA most commonly starts in the region next to the center of the retina, called the *fovea* and is called non-central or non-foveal GA (Figure 2A). Progression of GA lesions into the fovea in foveal GA (Figure 2B) leads to loss of central vision over time.

Dry-AMD stages, especially GA, can be also associated with whitishyellow dots connected in a reticular, or net-like pattern. These are termed reticular drusen or reticular pseudodrusen (RPD) or subretinal drusenoid deposits (Figure 3).



Figure 2A. GA most commonly starts in the fovea, the region next to the center of the retina. Image by of Suber Huang, MD, MBA, FASRS. Retina Image Bank; Image 6296. © American Society of Retina Specialists.



Figure 2B. Progression of GA lesions into the fovea leads to loss of central vision. Image by Gerardo Garcia-Aguirre, MD. Retina Image Bank; Image 258. © American Society of Retina Specialists.

Risk Factors:

- Age—the strongest risk factor
- Caucasian race

Possible risk factors:

- Female gender
- Lower education level
- Light iris color
- Far-sightedness
- Higher body mass index (BMI)
- High blood pressure

- Family history of AMD
- Cigarette smoking
- High cholesterol
- Sunlight exposure
- Low dietary fish intake
- Cardiovascular (heart) disease
- Having a diet low in lutein and zeaxanthin/antioxidants

Diagnostic Testing

Disease features related to AMD may be found in the retina and in the layers beneath it.

A diagnosis of dry AMD is made by a clinical examination with a slit lamp and by using several types of imaging, including:

- Optical coherence tomography (OCT)
- Fundus autofluorescence (FAF) imaging

To rule out conversion to the wet form of AMD, the following imaging studies may also be performed:

- Optical coherence tomography angiography (OCT-A)
- Fluorescein angiography (FA) and indocyanine green angiography (ICGA)

Visual function can be measured by using the patient's ability to read letters on eye charts. Also used: a technique named *microperimetry* that evaluates the sharpness of the vision in certain areas of the macula, or center of the retina responsible for central vision. continued next page

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Figure 3. Whitish-yellow dots in a reticular, or net-like pattern—termed *reticular drusen or* reticular pseudo-drusen (RPD) or subretinal drusenoid deposits-are associated with dry-AMD stages, particularly GA. Image by Jeffrey G. Gross, MD, FASRS. Retina Image Bank; Image 1082. © American Society of Retina Specialists.

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Treatment and Prognosis

There are two FDA-approved drugs, Syfovre (pegcetacoplan) and Izervay (avacincaptad pegol) that help to slow geographic atrophy progression. Both drugs are injections into the vitreous cavity of the eye every one to two months.

Also, the Age-Related Eye Disease Studies (AREDS), conducted by the National Eye Institute, have found that a nutritional supplement formula (AREDS2) may delay and prevent intermediate dry AMD from converting to wet AMD.

The AREDS2 supplement formula, which is widely available over the counter, contains specific doses of vitamins and minerals used in the AREDS2 Study: • Vitamin C

- Lutein
- Vitamin E
- Zeaxanthin
- Zinc
- Copper

Although patients with either form of AMD can experience a progressive decrease in visual acuity, it would be very rare to lose all vision.

Following a Mediterranean-type diet (higher whole-fruit consumption, lower red-meat consumption, low to moderate alcohol use, and a higher ratio of monounsaturated fatty acid to saturated fatty acid) is associated with a lower risk of progression to large drusen and to GA, and to slower enlargement of GA.

GA patients can take advantage of low-vision therapy using assistive devices to achieve improved mobility, reading and distance viewing, and facial recognition.

The Implantable Miniature Telescope is a surgical option available to lowvision patients with GA; it enlarges objects in the center of the visual field so they can be seen by intact areas of retina surrounding GA. Importantly, investigational treatments are in late stages of development for intermediate AMD, non-foveal and foveal GA.

Clinical Terms (appearing green within fact sheet text)

Fundus autofluorescence (FAF): An imaging modality that uses naturally occurring fluorescence from the retina to provide an indicator of retinal health.

Fluorescein angiography (FA) and indocyanine green angiography (ICGA): Imaging techniques where a yellow dye called *sodium fluorescein* is injected into a vein in the arm, allowing a special camera to record circulation in and under the retina. These tests can be very useful in diagnosing a number of retinal disorders.

Microperimetry: A test that measures the sensitivity to brightness at various locations in the macula.

Optical coherence tomography (OCT): A non-invasive technique that uses light to create a 3-dimensional image of your eye for physician evaluation.

Optical coherence tomography angiography (OCTA): A non-invasive imaging technique that uses light to image the blood vessels in different layers of the retina and choroid.

Slit lamp: An instrument that combines a high-intensity light source with a microscope to examine the external and internal structures of the eye, including the optic nerve and retina.

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