Autosomal recessive bestrophinopathy

INSERM

Source
INSERM. (1999). Orphanet: an online rare disease and orphan drug data base. **Autosomal recessive bestrophinopathy**. ORPHA:139455

Autosomal recessive bestrophinopathy (ARB) is a retinal dystrophy, characterized by central visual loss in the first 2 decades of life, associated with an absent electrooculogram (EOG) light rise and a reduced electroretinogram (ERG).