PURPOSE: To describe the ocular Behçet’s disease cases followed in our department, to a better understanding of this disease, its manifestations and clinic evolution.

METHODS: Retrospective study, observational, case series, of all patients with a diagnosis of ocular Behçet’s disease followed since 2020 in the eye inflammation consultation, at CHVNG/E. The collected data, namely clinical manifestations, severity of ocular involvement and complications, were analysed.

RESULTS: Since 2020, 10 cases of Behçet’s disease with ocular involvement were followed at CHVNG/E. 60% (n=6) were women and 80% of patients had bilateral ocular involvement. The most common clinical presentation was retinal vasculitis, presented in 60% of cases. Severe complications, as glaucoma (n=3), diffuse chorioretinal atrophy (n=2), and bilateral papillitis (n=1) were registered.

CONCLUSIONS: Behçet’s disease is a chronic multisystem disorder. Still a challenging diagnosis, it is characterized by occlusive vasculitis, which clinical manifestations may include ocular inflammatory involvement. The possible severe presentation and complications may result in severe visual loss. Thus, it is crucial to raise awareness of this disease and respective common clusters of symptoms.

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