Very early OCT response to intravenous pulse steroid in Vogt-Koyanagi-Harada disease

Miguel Leitão 💿 , Rita Pinto, Vanda Nogueira

Department of Ophthalmology, Instituto de Oftalmologia Doutor Gama Pinto, Lisboa, Portugal

Correspondence toDr Miguel Leitão;
miguelleitao3@gmail.com

Accepted 3 May 2023

DESCRIPTION

A Caucasian woman in her 40s presented with blurry vision in both eyes for 3 days. Best corrected visual acuity was 20/32 and 20/25 in her right eye and left eye, respectively. Intraocular pressure was 12 mmHg in both eyes, and anterior segment evaluation was unremarkable. Fundus examination revealed multiple neurosensory retinal detachments in the posterior pole and optic disc hyperemia bilaterally. Enhanced depth imaging-optical coherence tomography (EDI-OCT) showed large areas of subretinal fluid (SRF), bacillary layer detachment (BALAD) and markedly thickened choroidal layers in both eyes (figures 1A and 2A). There were no integumentary, neurological or auditory complaints. A diagnosis of presumptive Vogt-Koyanagi-Harada (VKH) was made and the patient was started on intravenous methylprednisolone 1 g/day for 3 days and mycophenolate mofetil. Approximately 20 min after the first pulse of methylprednisolone, there was remarkable improvement of SRF and BALAD already visible on EDI-OCT (figures 1B and 2B), which went on to resolve completely by week 1. In the following 8 months, there was a decrease in choroidal thickness and also progressive and continued reconstitution of the ellipsoid zone on EDI-OCT (figure 3). Mycophenolate was increased to 2 g/day over 8 weeks, and oral steroids were tapered gradually to 7.5 mg/day over 7 months. At 8 months' follow-up, patient remains stable with 20/20 vision in both eyes and sustained remission clinically and on optical coherence tomography (OCT). VKH disease is a primary stromal choroiditis, coursing with exudative retinal detachments and usually mild anterior and vitreous inflammation. Inflammation is thought to originate primarily from the choroid, affecting nearby structures in a

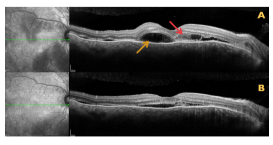


Figure 1 Right eye: (A) Multiple, macular SRF pockets (yellow arrow) and BALAD (red arrow) nasal to the fovea at baseline. (B) Improvement of SRF pockets and BALAD 20 min after the first pulse of intravenous methylprednisolone. BALAD, bacillary layer detachment; SRF, subretinal fluid.

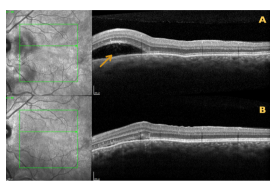


Figure 2 Left eye: (A) Single, peripapillary SRF pocket (yellow arrow). (B) Almost complete reabsortion of peripapillary SRF pocket 20 min after the first pulse of intravenous methylprednisolone. SRF, subretinal fluid.

spill-over fashion.^{1 2} Progression to chronic recurrent granulomatous anterior uveitis and sunset glow fundus with chorioretinal atrophy is likely, unless early and aggressive immunosuppression is implemented.³ It has been previously suggested that there may be a window of therapeutic opportunity, between 2 and 4 weeks of disease onset, where starting aggressive immunosuppression as soon as possible may modify the disease course,^{3 4} allowing for progression to sunset glow fundus to be altogether avoided. To the best of our knowledge, this is the first report wherein objective and remarkable improvement just minutes after a first pulse of intravenous steroid was documented, contributing to the existing body of evidence on the benefit of this approach. Steroid-sparing agents should also be introduced within the first few days to help prevent disease recurrence³ and provide the best chances for true disease course modification.⁵ The ongoing improvement of the outer retinal layers on OCT through to last follow-up at 8 months brings further ultrastructural detail in support of this strategy.

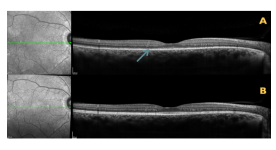


Figure 3 Right eye: (A) at 1 month, disruption of the EZ (blue arrow); (B) at 8 months, complete reconstitution of the EZ without any residual structural abnormalities. EZ, ellipsoid zone.



© BMJ Publishing Group Limited 2023. No commercial re-use. See rights and permissions. Published by BMJ.

To cite: Leitão M, Pinto R, Nogueira V. *BMJ Case Rep* 2023;**16**:e255186. doi:10.1136/bcr-2023-255186



Images in...

Learning points

- ➤ Vogt-Koyanagi-Harada disease is a primary stromal choroiditis that can progress to chronic illness with a poor visual prognosis if not treated promptly.
- On the first 2–4 weeks of presentation, treatment with systemic steroids and early immunosuppression may cure the disease.
- Response to pulsed intravenous methylprednisolone may be extremely fast, starting within minutes.
- Early pulsed intravenous methylprednisolone and/or early immunosuppression may promote continued restoration of outer retinal layers even after 6 months.

Contributors All authors meet the International Committee of Medical Journal Editors criteria for authorship for this article, take responsibility for the integrity of the work as a whole and have given their approval for this version to be published. Conception and design, acquisition of data, analysis and interpretation of data, drafting of the manuscript: revision of the manuscript for intellectual content and final approval of the completed manuscript: ML, RP and VN.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent for publication Consent obtained directly from patient(s).

Ethics approval This publication complied with the World Medical Association Declaration of Helsinki and conformed to the International Committee of Medical Journal Editors Recommendations for the Conduct, Reporting, Editing, and Publication of Scholarly Work in Medical Journals.

Provenance and peer review Not commissioned; externally peer reviewed.

Case reports provide a valuable learning resource for the scientific community and can indicate areas of interest for future research. They should not be used in isolation to guide treatment choices or public health policy.

ORCID ID

Miguel Leitão http://orcid.org/0000-0001-6565-4775

REFERENCES

- 1 Bouchenaki N, Herbort CP. Stromal choroiditis. In: Pleyer U, Mondino B, eds. Essentials in ophthalmology: uveitis and immunological disorders. Berlin, Heidelberg, New York: Springer, 2004: 234–53.
- 2 Bouchenaki N, Cimino L, Auer C, et al. Assessment and classification of choroidal vasculitis in posterior uveitis using indocyanine green angiography. Klin Monbl Augenheilkd 2002;219:243–9.
- 3 Herbort CP, Abu El Asrar AM, Takeuchi M, et al. Catching the therapeutic window of opportunity in early initial-onset Vogt-Koyanagi-Harada uveitis can cure the disease. Int Ophthalmol 2019;39:1419–25.
- 4 Herbort CP Jr, Tugal-Tutkun I, Abu-El-Asrar A, et al. Precise, simplified diagnostic criteria and optimised management of initial-onset Vogt–Koyanagi–Harada disease: an updated review. Eye 2022;36:29–43.
- 5 Papasavvas I, Tugal-Tutkun I, Herbort C. Vogt–Koyanagi–Harada is a curable autoimmune disease: early diagnosis and immediate dual steroidal and non-steroidal immunosuppression are crucial prerequisites. J Curr Ophthalmol 2020;32:310.

Copyright 2023 BMJ Publishing Group. All rights reserved. For permission to reuse any of this content visit https://www.bmj.com/company/products-services/rights-and-licensing/permissions/ BMJ Case Report Fellows may re-use this article for personal use and teaching without any further permission.

Become a Fellow of BMJ Case Reports today and you can:

- Submit as many cases as you like
- ► Enjoy fast sympathetic peer review and rapid publication of accepted articles
- ► Access all the published articles
- Re-use any of the published material for personal use and teaching without further permission

Customer Service

If you have any further queries about your subscription, please contact our customer services team on +44 (0) 207111 1105 or via email at support@bmj.com.

Visit casereports.bmj.com for more articles like this and to become a Fellow