Central serous chorioretinopathy in uveitis patients after corticosteroid therapy: a report of 6 cases

Natalia Skvortsova1, 2 ☞, Ioannis Papasavvas2, Carl P. Herbor2 ☞
1 Posterior Eye Segment Diagnostics and Surgery Center, Moscow, Russian Federation
2 Inflammatory and Retinal Eye Diseases, Centre for Ophthalmic Specialised Care, Lausanne, Switzerland
nat.skvortsova@gmail.com

Purpose. To report central serous chorioretinopathy (CSCR) in uveitis patients. Material and methods. A retrospective chart review of uveitis patients seen in a time frame of 20 years at the Centre for Ophthalmic Specialised Care, Lausanne, Switzerland. The ophthalmic and systemic features are presented. Results. Out of 1793 uveitis patients followed at the Centre for Ophthalmic Specialised Care, 6 patients (0.3%) developed CSCR following corticosteroid therapy due to uveitis. The mean age of patients was 40 ± 13.4 years, disease incidence was not associated with gender. In all 6 patients’ clinical disease was unilateral but subclinical signs were present in all fellow eyes. The mean duration of corticosteroid therapy before CSCR had occurred was 4.95 ± 4.0 months. The mean best-corrected visual acuity at the moment of CSCR was 0.6 ± 0.26 and 0.8 ± 0.17 after discontinuation of corticosteroids. Neurosensory retinal detachment and pigment epithelium detachment were observed in 3 eyes, respectively. During fluorescein angiography (FA), focal dye leakage and areas of alteration of RPE were observed in 6 out of 10 eyes. Diffuse hyperfluorescence of choroidal vessels observed by ICGA was detected in all eyes. Conclusion. Central serous chorioretinopathy should be suspected when functional and morphological deterioration occurs in uveitis patients receiving corticosteroid therapy with no signs of inflammation reactivation. This complication is extremely rare but serious condition which needs a prompt tapering and discontinuing of corticosteroids.

Keywords: uveitis; treatment; central serous chorioretinopathy; corticosteroids

Conflict of interest: The authors declare no conflict of interest.

For citation: Skvortsova N., Papasavvas I., Herbor Jr C.P. Central serous chorioretinopathy in uveitis patients after corticosteroid therapy: a report of 6 cases. Russian ophthalmological journal. 2021; 14 (3): 65-72 (In Russian). https://doi.org/10.21516/2072-0076-2021-14-3-65-72

Центра́льная серозная хориоре́тинопатия у пациен́тов с увеи́том после кортико́стерио́идной терапии: шесть клини́ческих случаев

Н.А. Скворцова1, 2 ☞, И. Папасавvas2, К.П. Хербор2 ☞
1 Центр диагностики и хирургии заднего отдела глаза, ул. 2-я Владимирская, д. 2, стр. 2, Москва, 111123, Россия
2 Центр специализированной офтальмологической помощи, отделение воспалительной и ретинальной патологии, ул. Шарль-Моннар, д. 6, 1003, Лозанна, Швейцария

Цель работы — описать случаи центральной серозной хориоретинопатии (ЦСХР) у пациентов с увеитом. Материал и методы. Представлен ретроспективный анализ, а также офтальмологические и системные особенности пациентов с увеитом, наблюдавшихся в период 20-летней работы Центра специализированной офтальмологической помощи в Лозанне (Швейцария). Результаты. Из 1793 пациентов с увеитом, наблюдавшихся в Центре, у 6 (0,3 %) после кортикостероидной терапии этого заболевания развилась ЦСХР. Средний возраст пациентов составлял 40 ± 13,4 года, возникновение заболевания не имело связи с полом. У всех 6 пациентов выявлялись клинические признаки ЦСХР на одном глазу и субклинические на парном. Средняя продолжительность кортикостероидной терапии перед развитием ЦСХР составляла 4,95 ± 4,0 месяца. Средняя максимально корригированная остра́та зрения на момент ЦСХР составляла 0,6 ± 0,26 и 0,8 ± 0,17 после прекращения кортикостероидной терапии. В 3 глазах наблюдалась соответственно нейросенсорная отслойка сетчатки и отслойка пигментного эпителия. В 6 глазах из 10 при флюоресценцентной ангиографии выявлено фокальное просачивание красителя и зоны повреждения ретинального
Central serous chorioretinopathy (CSCR) was initially described in 1866 by Albrecht von Graefe who called the disease “recurrent central retinitis” [1]. Takashi Masuda examined 192 cases and gave the name of “central serous chorioretinitis” to the disease in 1917 [2]. CSCR is one of the causes of vision loss in middle-aged predominantly male patients. The disease is characterized by the detachment of neurosensory retina (NSRD) with or without pigment epithelial detachment (PED).

Fluorescein angiography (FA) usually reveals leakage sites, demonstrating “ink-plot” or “smokestack” pattern of hyperfluorescent diffusion in acute CSCR. Multiple leakage sites, retinal pigment epithelium (RPE) atrophy and diffuse epitheliopathy are observed in chronic cases. Indocyanine green angiography (ICGA) finds increased choroidal permeability which is manifested by areas of hyperfluorescence from the mid-phase of the angiogram and diffuse late choroidal hyperfluorescence [3]. Optical coherence tomography (OCT) signs include subretinal fluid (neurosensory retinal detachment (NSRD), PED, and disturbances of RPE in chronic cases [4].

The aetiology of CSCR remains uncertain despite the numerous studies which have been published since 1866. The recent systematic review and meta-analysis have revealed several CRCS risk factors including systemic hypertension, steroid usage, sleep disturbance, autoimmune disease, psychopharmacologic medication use, and type-A behavior [5].

One theory considers a dysfunction of the RPE ion pump as a major reason of CSCR development leading to a reverse of fluid movement from the choroid to the retina [6]. However, more recently, CSCR has been included in the newly named group of pachychoroidal diseases [7]. The pathogenesis is thought to be triggered by cortisol and aldosterone which affect the autoregulation of the choroidal vasculature. S. Kuroda, et al. reported a diffusely thickened choroid in CSCR patients [8], while Y. Chung, et al. reported an increased Haller layer both in the affected eye and the unaffacted fellow eye of CSCR patients [9]. Similar changes were noted by R. Agrawal et al., who showed that the choroidal vascularity index was higher in eyes with acute CSCR in comparison to healthy eyes and resolved CSCR eyes [10]. These findings strongly suggest that both choroidal vascular dilatation and hyperpermeability play a crucial role in the development of CSCR [11]. Hyperpermeable choroidal vessels are presumed to increase hydrostatic pressure in the choroid leading to the barrier dysfunction of the RPE and fluid accumulation between the RPE and the retina [3, 11].

High endogenous serum cortisol levels [12] or corticosteroid administration in any form, systemic or topical on skin or on oral, nasal and conjunctival mucosas, are two important promoters of CSCR, as shown in numerous studies [13–19]. A relation was found between high cortisol levels in Cushing’s syndrome and thickening of the choroid and between intraocular dexamethasone and pachychoroid and CSCR development [20, 21], indicating that the role played by corticosteroids in the development of CSCR was through choroidal thickening.

Diagnosis of corticosteroid induced CSCR is relatively straightforward but gets a little trickier when it occurs in uveitis cases receiving systemic corticosteroid therapy.

The AIM of this study was to report a series of uveitis patients that developed CSCR.

**MATERIAL AND METHODS**

Medical charts of patients treated for uveitis and presenting CSCR were retrieved for retrospective analysis. Patients had had a complete work-up applied to patients with uveitis, comprising, in addition to routine features such as Snellen visual acuity, slit-lamp examination, applanation tonometry, and fundoscopy, laser flare photometry (LFP), computerized visual field (VF) testing, OCT, and dual FA and ICGA. The study was performed in accordance with the ethical standards laid down in the Declaration of Helsinki (1964) and in accordance with the IRB of our institution authorizing retrospective, anonymous and non-interventional studies.

**RESULTS**

Out of 1793 uveitis patients followed at the Centre for Ophthalmic Specialised Care during a period of twenty years, 6 patients (0.3%) developed CSCR under treatment. All of them had corticosteroids in their therapeutic regimen. The mean age (range) of patients (3 men/3 women) was 40 ± 13.4 (17–57) years. The mean duration of corticosteroid therapy before CSCR had occurred was 4.95 ± 4.0 months. The mean best-corrected visual acuity (BCVA) at the moment of CSCR diagnosis was 0.6 ± 0.26, whereas it was 0.8 ± 0.17 at last follow-up (p = 0.05, Student’s t test). Patients demographics and clinical data are presented in Table 1.

| Age (years) | Duration of corticosteroid therapy before CSCR | Best-corrected visual acuity (BCVA) |
|------------|-----------------------------------------------|-----------------------------------|
| 57         | 2 years                                       | 0.6 ± 0.26                        |
| 40         | 4.95 ± 4.0 months                             | 0.8 ± 0.17                        |

Mean LFP values were 44.7 ± 91.4 ph/bs showing medium inflammation (normal value of 4–6 ph/bs). OCT which was available in 4 patients (8 eyes) revealed NSRD in 3 eyes and small PEDs in 3 eyes as well. Concomitant NSRD and PED were observed in 1 eye. In 6 out of 10 eyes, focal dye leakage and areas of alteration of RPE were observed on FA. ICGA revealed diffuse hyperfluorescence of choroidal vessels in 10 out of 10 eyes (Table 2).

Three cases are briefly reported here.

*Case 1 (patient 1 on Table 1).* A 43-year-old Caucasian man with the diagnosis of birdshot retinochoroiditis (BRC) was given oral prednisonone (1 mg/kg) and cyclosporine A (5 mg/kg) 2 years after the diagnosis because of worsening of the inflammatory eye condition. The diagnosis was based on the presence of papillitis, diffuse retinal vasculitis, pseudo-delay in retinal FA arteriovenous circulation time, massive fluorescein impregnation of the retina, and cystoid macular oedema. On ICGA multifocal dark dots present in the intermediate and late ICGA phases as well as fuzziness of choroidal vessels seen during the intermediate and late phases of ICGA were typical of BRC. The HLA-A29 antigen was present. For 2 years, the patient’s condition had remained relatively
stable without treatment, BCVA remaining at 1.0 for both eyes, and laser flare photometry not detecting a subclinical anterior chamber inflammation. Both eyes had a slight vitritis and their fundus showed scattered, punched-out creamy lesions. At the moment of
corticosteroid and immunosuppressive therapy administration, the BCVA of the left eye was 0.8 with an anterior chamber inflammation having increased (laser flare photometry for OS = 15.1 ph/ms but normal values for OD (5.1 ph/ms), and visual field deterioration was noted in both eyes. After 10 weeks of systemic treatment, BCVA in both eyes was again 1.0 and LFP decreased to 5.4 ph/ms in the left eye. However, a few months later the patient presented a decrease of vision to 0.4 in the left eye with inflammatory parameters under control on both sides. FA revealed a hyperfluorescent leaking point and ICGA showed a focal brightly hyperfluorescent spot that corresponded to the leakage area on FAG (Fig. 1). All these findings were compatible with CSCR. Therefore, prednisone was progressively tapered with improvement of visual acuity to 0.7 in the left eye. It is interesting to note that hyperpermeable (hyperfluorescent) choroidal vessels were already present on ICGA before corticosteroid administration. As explained hereunder uveitis entities with predominant choroidal inflammation are more prone to secondary CSCR.

**Case 2 (patient 3 on table 1).** A 47-year-old Caucasian man presented with photophobia and vision loss in the right eye. BCVA was 0.2 for the right eye, while BCVA was 0.8 for the left. LFP revealed anterior chamber inflammation (LFP OD: 49.4 ph/ms, OS: 21.2 ph/ms). Vitritis, snowballs, retinal scars,

---

**Table 1.** Demographics and clinical data

| Patient | Gender | Age | Diagnosis       | Duration of corticosteroid treatment | BCVA at the moment of CSCR | BCVA at the last follow-up |
|---------|--------|-----|-----------------|------------------------------------|-----------------------------|-----------------------------|
| 1       | M      | 43  | BRC            | 10 weeks                           | 0.4                         | 0.7                         |
| 2       | F      | 35  | VKH            | 2 months                           | 0.9                         | 0.7*                        |
| 3       | M      | 47  | presumed TB    | 5 weeks                            | 0.6                         | 1                           |
| 4       | M      | 57  | idiopathic     | 6 months                           | 0.7                         | 1                           |
| 5       | F      | 41  | Scleritis, idiop. | 6 months                        | 0.2                         | 0.6**                       |
| 6       | F      | 17  | idiopathic     | 12 months                          | 0.8                         | 0.8                         |

**Note.** BCVA — best corrected visual acuity, BRC — birdshot retinochoroiditis, VKH — Vogt-Koyanagi-Harada disease, TB — tuberculosis. *— choroidal neovascularization successfully treated with intravitreal aflibercept injection, **— after cataract surgery.

**Table 2.** Features of CSCR revealed by OCT, FAG, and ICGA

| Features                                      | Number of eyes |
|----------------------------------------------|----------------|
| OCT Neurosensory detachment (NSRD)           | 8 (4 patients) |
| Pigment Epith. Detachment (PED)              | 3/8            |
| FAG Focal dye leakage, areas of alteration of RPE | 10 (5 patients) |
| ICGA Hyperfluorescent leaking spot, diffuse hyperfluorescence of choroidal vessels | 10 (5 patients) |

---

**Fig. 1.** Case 1. Birdshot Retinochoroiditis (left eye). (a) — FAG revealed diffuse retinal vasculitis and disc hyperfluorescence at the moment of birdshot retinochoroiditis diagnosis, (b) — ICGA revealed numerous hypofluorescent dark dots (HDDs) at the moment of birdshot retinochoroiditis diagnosis, (c) — FAG showed no signs of more active inflammation with decreased disc fluorescence, a dye leakage point, and signs of detachment of neurosensory retina at the moment of CSCR, (d) — ICGA showed decrease of HDDs, improved aspect of choroidal vessels with decreased fuzziness but hyperpermeable posterior pole choroidal vessels at the moment of CSCR.[already present before corticosteroid administration (b)].

**Рис. 1.** Клинический случай 1. Ретинохориоидит Бирдшота (левый глаз). (а) — FAG выявила диффузный ретинальный васкулит и дисперсию хориоидального диска, (б) — на момент постановки диагноза «ретинохориоидит Бирдшота» ФАГ выявил диффузный ретинальный васкулит и гиперпермутирующую точку диска, (в) — на момент постановки диагноза ЦСХР ФАГ не выявил признаков активного воспаления со снижением флюоресценции диска, точек просачивания красителя и признаков отслойки нейросенсорной сетчатки, (d) — на момент ЦСХР (непосредственно перед кортикостероидной терапией) ангиография с индоксанином зеленым показала снижение HDDs, увеличение вида хориоидальных сосудов в существенно не измененных сосудисто-сетчатых областях (B).
and cystoid macular oedema were observed in the right eye. Mantoux — test showed a hyper-reaction with a skin papule size of 12 × 11 mm. The diagnosis of presumed tuberculous (TB)-related uveitis was posed. After administration of triple anti-TB therapy and oral prednisone (30 mg), BCVA improved to 1.0 in both eyes in 10 days. Five weeks later, the patient presented with a decreased BCVA to 0.6 (OD), while anterior chamber inflammation had substantially decreased (LFP RE: 18.1 ph/ms, LE: 9.3 ph/ms) under 20 mg of prednisone. The condition was interpreted as a recurrence of the inflammatory process and dosage of prednisone was raised up to 50 mg. 5 days later, the patient’s condition did not change but laser flare photometry decreased slightly (16.0 ph/ms on the right eye). NSRD and PED were noted. Central serous chorioretinopathy was diagnosed and steroid treatment was discontinued. 6 months later, on the last follow-up, no signs of active inflammation were seen, BCVA was 1.0 on the right eye, and LFP remained stable at 21.7 ph/ms. LFP was very helpful to precisely follow intraocular inflammation and to avoid incriminating an increase in inflammation for a decrease in function caused by CSCR (Fig. 2, 3).

Case 2. Tuberculous chorioretinitis. The crucial role of LFP to distinguish between functional impact of uveitis and CSCR. Parallel evolution of visual acuity (VA) and LFP values in the right eye of the patient in relation to the dose of prednisone (mg) (orange bars). At presentation (sector 1), LFP values are high and VA is low. Following administration of 30 mg of prednisone, VA increases to 1.0 and LFP values decrease substantially (sector 2). In sector 3 VA decreases again under 20 mg of prednisone with persistent low LFP values, due to CSCR lesions. In sector 4, prednisone of azathioprine). 6 months later, the patient presented with vision intensification (32 mg of prednisone with slow tapering plus 200 mg of azathioprine). 6 months later, the patient presented with vision deterioration in the right eye (under 24 mg of prednisone). BCVA was 0.2 for the right eye and 0.5 for the left eye, respectively. PED in the left eye (Fig. 4) was detected by FA and OCT suggestive of CSCR. Corticosteroid therapy was stopped (following tapering during 3 months), anti-TNF-α therapy (Infliximab) was administered. 9 months later, the treatment regimen was azathioprine 200 mg/day. Infliximab every 6 weeks. At the last follow-up after cataract surgery, BCVA was 0.6 for the right eye and 0.5 for the left eye. LFP was 6.5 ph/ms for OD and 6.7 ph/ms for OS, respectively. OCT did not reveal NSRD nor PED in the right eye and FA did not reveal signs of papillitis nor vasculitis.

DISCUSSION

Glucocorticosteroid therapy remains the first-line treatment for the majority of ocular inflammatory diseases including uveitis. Use of systemic and local corticosteroids can be involved in CSCR development. However, reports on CSCR in posterior uveitis patients receiving corticosteroid therapy are limited [22–24, 25]. In our collective of 1793 cases of uveitis patients seen in our centre, 6 uveitis cases (0.3 %) treated with systemic corticosteroids developed CSCR. While CSCR development in autoimmune diseases has been reported often [15, 26], occurrence of CSCR in corticosteroid treated uveitis patients tends to be rare. In three of our six cases, inflammation involved the choroid. Predilection of CSCR for choroiditis entities has been reported previously [25].

Maculopathy associated with corticosteroid use was initially reported in 1966 [27]. I. Jain and K. Singh described a case of 42-year-old male with Reiter syndrome who had been treated with oral prednisone. A serous macular detachment occurred after 1 week of treatment; therefore, Jain and Singh associated this maculopathy with corticosteroid usage. CSCR induced by corticosteroids usage has less male predisposition than the idiopathic disease. The relationship between CSCR and corticosteroids is one of the most interesting aspect of the disease. Two large retrospec-
Central serous chorioretinopathy in uveitis patients after corticosteroid therapy: a report of 6 cases

Case 2. Tuberculous chorioretinitis. FA and ICGA signs of CSCR. Top four frames show the evolution of FA signs. (1) before dual antibiotic and prednisone therapy (cystoid macular oedema and disc hyperfluorescence); (2) good response to treatment; (3) new zone of macular hyperfluorescence (arrow) due to CSCR in a quiet eye; (4) resolution of CSCR after discontinuation of corticosteroids. Bottom two frames show evolution of ICGA signs. The left frame (pre) shows the right eye before therapy. Disc hyperfluorescence on ICGA (usually non fluorescent) indicates severe posterior inflammation. Right frame (post) shows situation after treatment, in particular corticosteroids, have been stopped. Persistent ICGA hyperfluorescence explained by residual pachychoroid

Laser flare photometry is another crucial investigation as it measures exactly intraocular inflammation and can so precisely exclude an inflammatory reactivation [30].

In this study, the strongest positive sign in the diagnosis of CSCR was choroidal hyperfluorescence due to hyperpermeable choroidal vessels revealed by ICGA (seen in 10 eyes out of 10, in one patient (2 eyes) ICGA was not performed). FA findings (focal dye leakage and areas of RPE alteration) and OCT findings (PED...
and NSRD) were helpful in the establishing the diagnosis of CSCR as well. Surprisingly, the simultaneous detachment of neurosensory retina and RPE was observed only in 1 eye.

There may be similarities between acute initial-onset Vogt—Koyanagi—Harada disease and bilateral CSCR. However, the ICGA and OCT patterns differ sufficiently to differentiate the diseases [31, 32].

The management of corticosteroid induced CSCR in posterior uveitis consists in diligent tapering of corticosteroid therapy and discontinuation associated with replacement by other immunosuppressive or immunomodulatory therapies [33]. If this is not sufficient, mineralocorticoid antagonists may be used as for idiopathic CSRC [34]. Indeed, glucocorticosteroid and mineralocorticoid receptors were shown to be co-expressed in the choroid and retina [35]. In our series we did not have to resort to mineralocorticoid antagonists, probably because the trigger of CSCR occurred through the glucocorticoid receptors or through a weaker stimulation of mineralocorticoid receptors by glucocorticoids so that discontinuation of glucocorticosteroids was sufficient [36].

CONCLUSION

Though the rate of CSCR among uveitis patients treated with corticosteroid therapy is rare, misinterpretation of CSCR as worsening of the uveitis is the pitfall to avoid. Indeed, this could lead to the vicious circle of increasing corticosteroids with its harmful consequences. Careful assessment of fundus appearance, laser-flare photometry, FA, ICGA, and OCT enable the clinician not to miss signs of CSCR. This disease should be borne in mind when vision deterioration occurs in uveitis patients under corticosteroid therapy with no signs of reactivating inflammation. It should lead the clinician to tapering and discontinuation of corticosteroids with replacement by other immunosuppressive agents, if necessary.
References/Литература

1. Von Graefe A. Kurzere Abhandlungen. Notizen und casaisitiche Mittheilungen
   vermischten Inhalts: VI. Ueber Zentrale Recidivirende Retinitis. Albrecht
   Von Graefes Arch. Klein. Exp. Ophthalmol. 1866; 12: 211–5.

2. Masuda T. Central serous chorioretinopathy. J. Jpn. Ophthalmol. Soc. (Nihon
   Ganka Gakkai Zasshi) 1917; 21: 777.

3. Spaida R.F., Hall L., Haas A., et al. Indocyanine green videoangiography of older
   patients with central serous chorioretinopathy. Retina. 1996; 16 (3): 203–13.
   doi: 10.1097/00001692-19961603-00004

4. Durach A., Matet A., Draji A., et al. Central serous chorioretinopathy: Recent
   findings and new physiopathology hypothesis. Prog. Retin. Eye Res. 2015; 48:
   12–118. doi: 10.1016/j.preteyeres.2015.05.003.

5. Liu B., Dong T., Zhang J. Risk factors for central serous chorioretinopathy: a
   systematic review and meta-analysis. Retina. 2016; 36 (1): 9–19. doi: 10.1097/
   IARE.0000000000000837

6. Spitznas M. Pathogenesis of central serous retinopathy: A new working hypo-
   thesis. Graefes Arch. Clin. Exp. Ophthalmol. 1986; 224: 321–4. doi: 10.1007/
   BF02150023

7. Yanagi Y. Pachychoroid disease: a new perspective on exudative maculopathy.
   Jpn. J. Ophthalmol. 2020; 64: 323–37. doi: 10.1007/s10384-020-00740-5

8. Kuroda S., Kuno Y., Yasuno Y., et al. Choroidal thickness in central serous chorio-
   retinopathy. Retina. 2013; 33 (2): 302–8. doi: 10.1097/IAE.0b013e318263d1f1

9. Chang Y.R., Kim J.W., Kim S.W., et al. Choroidal thickness in patients with
   central serous retinopathy: assessment of Haller and Sattler layers. Retina.
   2016; 36 (9): 1652–7. doi: 10.1097/IAE.0000000000000998

10. Agrawal R., Chhablani J., Tan K.A., et al. Choroidal vascularity index in central
    serous chorioretinopathy. Retina. 2016; 36 (9): 1646–51. doi: 10.1097/
    IARE.0000000000001040

11. Yannuzzi L.A., Slakter J.S., Gross N.E., et al. Indocyanine green angiography-
    guided photodynamic therapy for treatment of chronic central serous chorio-
    retinopathy: a pilot study. Retina. 2003; 23 (3): 288–98. doi: 10.1097/00001692-
    200306000-00002

12. Lenk J., Sandner D., Schindler L., et al. Hair cortisol concentration in patients
    with active central serous chorioretinopathy is elevated — a pilot study. Acta
    Ophthalmol. 2019; 97 (4): e568–e571. doi: 10.1111/aos.13979

13. Chaine G., Haouat M., Menard-Molcard C., et al. Central serous chorioreto-
    nopathy and systemic corticosteroid therapy. J. Fr. Ophtalmol. 2001; 24 (2): 139–46.

14. Shin W.B., Kim M.K., Lee C.S., et al. Comparison of the clinical manifestations
    between acute Vogt-Koyanagi-Harada disease and acute bilateral central serous
    chorioretinopathy. Korean J. Ophthalmol. 2015; 29 (6): 389–95. doi: 10.3341/
    kjoph.2015.29.6.389

15. Lin D., Chen W., Zhang H., et al. Comparison of the optical coherence tomo-
    graphy characters between acute Vogt-Koyanagi-Harada disease and acute
    central serous chorioretinopathy. BMC Ophthalmol. 2014; 28: 14–87. doi: 10.1186/
    1471-2415-14-87

16. Takayama K., Obata H., Takeuchi M. Efficacy of adalimumab for chronic
    Vogt-Koyanagi-Harada disease refractory to conventional corticoste-
    roids and immunosuppressive therapy and complicated by central serous
    chorioretinopathy. Ocul. Immunol. Inflamm. 2020; 28 (3): 509–12. doi: 10.1080/
    09273948.2019.1603312

17. Bouquet E., Zhao M., Daruich A., et al. Mineralocorticoid antagonists in the
    treatment of central serous chorioretinopathy: review of the pre-
    clinical and clinical evidence. Exp. Eye Res. 2019; 187:107754. doi: 10.1016/j.
    exer.2019.107754

18. van Dijk E.H., Nijhof M.F., de Jong E.K., et al. Central serous chorioretinopathy
    in primary hyperaldosteronism. Graefes Arch. Clin. Exp. Ophthalmol. 2016;
    254 (10): 2033–42. doi: 10.1007/s00417-016-3417-8

19. Daruich A., Matet A., Draji A., et al. Central serous chorioretinopathy: recent
    findings and new physiopathology hypothesis. Prog. Retin. Eye Res. 2015; 48:
    82–118. doi: 10.1016/j.preteyeres.2015.05.003.

Вклад авторов в работе: К.П. Херборт — концепция и дизайн исследования, анализ литературы, написание статьи; Н.А. Скворцова, И. Папасаввас — анализ литературы, написание статьи.

Authors' contribution: Carl P. Herbort Jr — is at the origin of the concept of the study, carried out the bibliographic research and the writing of the study; Natalia Skvortsova, Ioannis Papasavvas — carried out the bibliographic research and the writing of the study.

Поступила: 19.01.2021, Переработана: 14.02.2021. Принята к печати: 15.02.2021

Originally received: 19.01.2021. Final revision: 14.02.2021. Accepted: 15.02.2021.
INFORMATION ABOUT THE AUTHORS/ ИНФОРМАЦИЯ ОБ АВТОРАХ

1 Posterior Eye Segment Diagnostics and Surgery Center, 2 bld., 2, Vladimirskaya str., Moscow, 111123, Russia
2 Inflammatory and Retinal Eye Diseases, Centre for Ophthalmic Specialised care, Rue Charles-Monnard 6, 1003 Lausanne, Switzerland

Natalia A. Skvortsova — MD1, 2
Carl P. Herborg Jr — MD, PD1
Ioannis Papasavvas — FEBOpth2

Contact information: Natalia A. Skvortsova, nat.skvortsova@gmail.com
Carl P. Herborg Jr, cph@herbortuveitis.ch

1 Центр диагностики и хирургии заднего отдела глаза, ул. 2-я Владимировская, 2, стр. 2, Москва, 111123, Россия
2 Центр специализированной офтальмологической помощи, отделение воспалительной и ретинальной патологии, ул. Шарль-Моннар, д. 6, Лозанна, 1003, Швейцария

Наталья Андреевна Скворцова — врач-офтальмолог1, 2
Иоаннис Папасаввас — врач-офтальмолог2
Карл П. Херборт мл. — д-р медицины, профессор2

Для контактов: Наталья Андреевна Скворцова, nat.skvortsova@gmail.com
Карл П. Херборт мл.
cph@herbortuveitis.ch