Introduction

Idiopathic central serous chorioretinopathy (CSC) is a disease first described by Albrecht von Graefe in 1866 as “recurrent central retinitis”. It is characterized by detachment of the retinal neuroepithelium from the pigment epithelium with the accumulation of clear fluid in the macular region (7). The disease mainly affects middle-aged males (20 to 50 years of age) (7). It is the cause of a sudden, mild decrease of visual acuity, micropsia, metamorphopsia and defects in the centre of the visual field. Generally, the disease has a good prognosis. It depends on the resorption of fluid and thus the normalization of visual functions. The final result is dependant on the eventual damage of the neural part of the retina by fluid accumulation in the acute phase of the disease. Unfavourable prognosis is attributed to recurrence of attacks or less frequently, to the prolonged or chronic course of the disease. This leads to irreversible structural changes of the macula with a permanent decrease of visual acuity. Absorption of the fluid is in many cases spontaneous within a few weeks. The therapy usually used in chronic CSC, comprises of nonsteroidal antiinflammatory drugs, vasoprotectives, resorbients, vitamins, systemic beta-blockers or acupuncture (34). One of the treatment methods of acute idiopathic CSC in strictly indicated cases is direct laser photocoagulation at the source of the fluid leakage from the choroid under the retina. This significantly reduces the duration of ablation of the neuroretina and prevents development of chronic changes due to CSC with irreversible anatomical and functional damage (3, 17, 30, 37).

Many authors describe diminished contrast sensitivity (CS), colour discrimination and focal or multifocal electroretinography (mERG) responses in acute idiopathic CSC not only when best corrected visual acuity (BCVA) decreases, but also when it remains satisfactory (1, 2, 5, 10, 14, 21, 25, 35). Some studies refer co-existing alteration of the retinal function within the central 40° area on the fellow, unaffected eye of the patients (10, 21). Others observe that its abnormalities extend beyond the clinically observed fundus changes mainly in clinically affected eyes (26) or, only in affected eyes limited to the pathological retinal region (35). The persistence of colour vision defects, decreased CS, decreased central retinal sensitivity on microperimetry and subnormal or low-normal signals from central retinal region by means of mERG was described also after spontaneous resolution of unilateral CSC either only on the affected eyes (28, 33), or also in clinically uninvolved, fellow eyes of patients (2, 10, 13, 19, 20, 31). Changes in the foveal pho-
to the receptor layer in CSC may cause the persistent functional deficit and predict visual recovery after macular reattachment in CSC patients (29).

The aim of our study was to evaluate the functional and anatomic retinal changes in patients with the first attack of unilateral CSC within a time period of two years after the treatment with direct laser photocoagulation and compare them with the fellow, unaffected eye of the same patients and with an age-similar normally-sighted control group.

**Material and Methods**

Fourteen patients (14 males) of median age 44 years (range 35 to 51 years) with the first attack of unilateral acute idiopathic CSC were treated at the Department of Ophthalmology of the Charles University in Hradec Králové. Diagnosis was set approximately three weeks from the onset of the disease (range 1 to 8 weeks). In five cases the right eye was affected (36 %) and in nine cases (64 %) the left eye. In all cases, it was the first attack of the disease without any other concurrent pathology of the affected eye and of the fellow eye. The patients had undergone a routine ophthalmologic examination, fundus photography of both eyes at the standard 30° (Zeiss, Germany). Fluorescein angiography (FAG) was performed to localize the leak points at the level of the retinal pigment epithelium (RPE). The patients did not suffer with potential relevant systemic diseases.

All patients underwent direct laser photocoagulation using diode integrated green laser with a wavelength of 532 nm (Integra SP, Laserex, USA). The laser burns were applied to the point where the earliest hyperfluorescent spot was seen in the earliest phase of the FAG. In this way, the laser area could be confined to the source of hyperfluorescence without excess affection of the detached surface of the neuroepithelium that remained unaffected by the laser. The distance between the laser points and the centre of the macula was strictly kept to at least 500 μm, the point size was 200 μm, and the laser was applied for 0.2 seconds with 130–290 mW.

Both eyes of patients were examined at the beginning of the disease and two years after treatment. The affected eyes were examined in addition 2 months and one year after laser photocoagulation. Best corrected visual acuity, contrast sensitivity and optical coherence tomography (OCT) were tested at all visits, colour vision and electroretinography were performed 2 years after the treatment.

The control group comprised of 20 healthy eyes (10 males) of median age 44 years (range 35 to 51 years) with the first attack of unilateral acute idiopathic CSC were treated at the Department of Ophthalmology of the Charles University in Hradec Králové. All individuals gave their informed consent prior to their inclusion in the study.

The threshold BCVA was measured under controlled lighting conditions using high-contrast (97 %) optotype logMAR charts (Small medical technique, Czech Republic). The distance for examination was 4 meters. Each of the 14 rows of the chart for testing visual acuity between 0.1 (20/200) and 2.0 (20/10) contained 10 Landolt rings, each with a predefined space in one of eight directions. Their size in the subsequent rows had a logarithmic progression. The number of correct answers was noted and the method of Ferris et al. (4) was used for the calculation of the threshold.

The CS was tested using a computerized Contrast Sensitivity 8010 System (Neuroscientific Corp., Farmingdale USA). The distance for CS measurement was 2.2 m so that a range of spatial frequencies from 0.74 to 29.55 c/deg was achieved. The size of the monitor appeared as 5 deg x 3.5 deg in the centre of the visual field. CS was measured by the method of ascending and descending limits for six spatial frequencies: two low (0.74; 1.97 c/deg), two intermediate (3.69; 7.39 c/deg) and two high frequencies (14.77 and 29.55 c/deg) (15). The background illumination was set to 85 cd m⁻². The contrast sensitivity was evaluated in [dB] = 20*log(CS). The eyes were refracted for CS examination, the median pupil size was 4.0 mm (3.5–4.5 mm).

OCT Stratus (Zeiss, Germany) was used for evaluation of the central retinal thickness of the affected eyes of patients. The macula was examined using “Macular thickness analysis” in 6 radial scans and the central retinal area of 6 mm in diameter was evaluated. As a central retinal thickness we considered the thickness in 1 mm zone.

The colour vision was examined using Farnsworth D-15 and Lanthony D-15 desaturated tests (X-Rite, Grand Rapids USA). The Panel D-15 total error score and colour confusion index were calculated using the calculation proposed by Lanthony (16). The colour confusion index (CCI) was calculated as relationship between the actual and the normal total colour difference score (TCDS_{actual}/TCDS_{normal}) (36).

Electroretinograms (ERG) were recorded from the cornea with DTL fiber electrodes (UniMed Electrode Supplies, England). The reference and ground skin electrodes (gold cup electrodes) were attached to the ipsilateral temples and forehead respectively. The pupils were fully dilated (0.5 % tropicamide and 10 % phenylephrine) and the patients were adapted to light. The eyes were refracted for multifocal ERG.

For conventional photopic full-field electroretinography (phERGs) and mfERG we used the RETI-port plus mfERG (Roland Consult GmbH, Germany). Light adapted 3.0 ERG and 3.0 flicker ERG were registered under recording conditions that conformed to the International Standard for Clinical Electroretinography (ISCEV) (22, 23). The background luminance was set to 30 cd m⁻², the luminance of flash stimulus was 3 cd m⁻². We analysed amplitudes and implicit times of the a- and b-waves of light adapted 3.0 ERG and 3.0 flicker ERG response.
The mfERG stimulus was generated on a 20-inch ViewSonic monitor with a frame rate of 75 Hz positioned 30 cm from the subject’s eye. mfERG traces were recorded from the central 60° of the retina with a resolution of 61-scaled hexagons according to the ISCEV guidelines (9, 24, 32). The luminance of the stimulus was 160 cd s m^{-2} for the white flashes and 1.4 cd s m^{-2} for the black ones and the luminance of the surrounding screen area was set to 80 cd m^{-2}. A red cross was presented for fixation. The recordings were sampled at 1033 Hz, amplified (x 200 000) and band pass filtered (10–100 Hz). We evaluated amplitudes and implicit times of the positive peak component (P1) of the first order kernel analysis for 5 concentric rings centred on the fovea.

For statistical analysis we used a nonparametric Mann-Whitney and Kruskal-Wallis tests and a p≤0.05 was considered statistically significant.

Results

Restoration of the detached RPE was reached in 14 eyes of 14 patients (100 %) after laser treatment. One laser burn was enough to restore the neuroepithelium of the retina in seven patients (50 %). For four other patients (28.6 %) 2 or 3 burns were applied. In three cases (21 %) we had to repeat the process once more after 3 to 5 months and thereafter more repetitions were not necessary. Recurrences were not noted in any patient and fellow eyes of all patients remained unaffected within the time period of two years.

Median BCVA of the control group was 1.1 (0.83–1.51). The course of BCVA of the affected eye of patients before and after treatment with the two-year follow-up period in comparison with the normal limits is shown in Fig. 1. The median BCVA of the affected eyes of patients was 0.69 (0.31–0.85) before treatment. After the treatment, BCVA improved significantly (p<0.001) to 0.91 (0.59–0.98) at 2-months follow-up, and further to 0.95 (0.52–1.11) after 1-year in affected eyes, but still remained significantly lower compared to the controls (p=0.03 and p=0.04). The final median BCVA of 0.91 (0.61–1.25) did not differ significantly from the controls. Baseline median BCVA of the fellow eye of patients was 1.02 (0.64–1.33), at 2 years follow-up it increased significantly (p<0.05) to 1.05 (0.94–1.45). Baseline BCVA of both eyes of patients was significantly lower in comparison with that of the controls (p<0.001 for the af-

<table>
<thead>
<tr>
<th>Tab. 1: Contrast sensitivity [dB].</th>
<th>spatial frequencies</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.74</td>
</tr>
<tr>
<td>RCS before therapy</td>
<td>23.25</td>
</tr>
<tr>
<td>RCS after 2 years</td>
<td>28.50</td>
</tr>
<tr>
<td>FE before therapy</td>
<td>28.31</td>
</tr>
<tr>
<td>FE after 2 years</td>
<td>30.18</td>
</tr>
<tr>
<td>controls: P 0.05</td>
<td>26.62</td>
</tr>
<tr>
<td>controls: P 0.95</td>
<td>34.70</td>
</tr>
</tbody>
</table>

The baseline and final median values of contrast sensitivity [dB] of central serous chorioretinopathy (CSC) and the fellow eyes (FE) of patients and normal range of CS in 6 spatial frequencies (90% CL).

Fig. 1: Best corrected visual acuity (BCVA). BCVA of controls and both eyes of patients measured at all examinations. The columns represent their median values.
fected eyes, and \( p < 0.01 \) for fellow eyes) whereas the final BCVA of both eyes of patients was only nonsignificantly lower compared to controls.

The baseline and final CS of both eyes of patients and its normal values are summarized in Table 1. Before treatment, the CS of the affected eyes was significantly lower in comparison with both the clinically unaffected, fellow eye of patients (\( p < 0.01 \) at a frequency of 1.97 c/deg, and \( p < 0.001 \) at all other spatial frequencies) and with the control group (\( p < 0.001 \) at all spatial frequencies). Moreover, baseline CS in the fellow, clinically uninvolved eyes of patients was also significantly lower when compared to the controls (\( p < 0.05 \) to \( p < 0.001 \)). The decrease of CS in fellow eyes of patients was most pronounced at intermediate spatial frequencies and smallest at the highest frequency. After treatment the CS improved significantly and the final CS of both eyes of patients reached normal values, with the exception of significantly lower CS at a spatial frequency of 3.69 c/deg for the affected eyes. The course of recovery of CS in the affected eye within the whole follow-up period is shown in Fig. 2. At 2 months after laser photocoagulation, the CS improved significantly (\( p < 0.001 \)) but still remained significantly lower at all frequencies (\( p < 0.05 \) to \( p < 0.001 \)) in comparison with the controls. Despite a further significant increase after one year at the majority of frequencies (\( p < 0.05 \) to \( p < 0.001 \)), CS remained significantly lower (\( p < 0.05 \)) at intermediate frequencies in comparison with the controls. In the second year CS remained practically stable.

OCT examination confirmed a statistically significant reduction (\( p < 0.001 \)) of central retinal thickness in the affected eyes of patients from an initial median value of 293 μm (range 140 μm – 548 μm) to a median value of 190 μm (range 125 μm – 277 μm) two months after the laser photocoagulation. At the one year follow-up examination, the central retinal thickness further decreased and reached the normal limits with median thickness of 150 μm (range 106 μm – 184 μm) and remained stable at 2-years of follow-up examination.

Two years after therapy, the colour discrimination was normal in the affected eyes of 12 patients (86 %) and in the fellow eyes of all patients (100 %) with a total mean error score = 0 using the Lanthony Panel D-15 desaturated test. In two patients (14 %), the Farnsworth D-15 test was normal, whereas the Lanthony D-15 desaturated test detected colour vision impairment in the blue region with a total error score of 56 and 136, and CCI of 1.39 and 1.96 respectively.

Two years after treatment, the amplitudes of light adapted 3.0 ERG reached the median value of 35 μV (27–58 μV) for the a-wave, 141 μV (111–207 μV) for the b-wave and 121 μV (90–167 μV) for the 3.0 flicker ERG in the affected eyes of the patients. The amplitudes of both ph-ERG responses were similar in fellow, clinically uninvolved eyes as shown in Fig. 3. The statistical analysis showed only nonsignificant differences in comparison with the control group. Implicit times of both ph-ERG responses were also within the normal range in both eyes of all patients (data not shown).

Figs. 4a and 4b summarize the averaged mfERG amplitudes and implicit times of the P1 component of the first order kernel analysis for all 5 retinal areas for both eyes of all 14 patients at two years follow-up after laser treatment in comparison with the controls. In affected eyes, the alteration of the average median P1 amplitudes was statistically nonsignificant over the entire central retina when compared both to the fellow eyes of patients and to the controls. The average median P1 implicit times were also nonsignificantly prolonged compared to patients fellow eyes and controls, with the exception of a significantly longer P1 implicit time in the retinal region between 1.8° and 7° of eccentricity in affected eyes compared to the controls (\( p = 0.049 \)). However one patient had low average P1 amplitudes in all 5 annular retinal regions (ring 1: 29.8; ring 2: 10.2; ring 3: 4.1; ring 4: 2.78 and ring 5: 1.76 nVdeg⁻²) and in one patient

Fig. 2: Contrast sensitivity (CS). Median values of contrast sensitivity [dB] measured at each examination in the affected eyes of patients and their comparison with normal range of CS (90 % CL).
**Fig. 3:** Photopic ERG (ph-ERG). Amplitudes of both ph-ERG responses. The columns represent their median values [μV].

**Fig. 4 a,b:** Multifocal ERG (mfERG). Box plot with mfERG a) amplitudes [μV] and b) implicit times [msec] of the P1 component of the first order kernel analysis for 5 concentric rings around the fovea. The horizontal lines represent the median values, the boxes display the 25% and 75% percentiles and the error bars show the minimum and maximum data values.
we found low activity in the central 14° area (ring 1: 19.3; ring 2: 29.4; ring 3: 30.1; ring 4: 21.4 and ring 5: 16.5 nVdeg−2) in the affected eyes. In the fellow eyes of all 14 patients, all average mfERG responses were within the normal range.

Discussion

In our prospective study with a 2-year follow-up period we included patients with a first attack of unilateral CSC who had undergone direct laser photocoagulation of the focal leakage sources. We evaluated the macular changes of the affected eyes and compared them with the fellow, uninvolved eyes of the same patients and with an age-similar control group.

Our study shows that the average standardized threshold BCVA of both eyes of the patients was significantly lower than that of the control group at the beginning of the disease. Frisén and Frisén (6) studied BCVA and micropsia in patients with macular oedema and pointed out that a BCVA of 1.0 could be reached even with a loss of 44% of the total amount of fibres in independent pathways joining foveolar cones with visual cortex centre. This suggests that even a BCVA of 1.0 can imply a marked decrease of the central visual functions and thus such a visual acuity should not be accepted as normal without a detailed analysis.

Within two months after laser treatment the BCVA of affected eyes significantly improved and this improvement gradually continued. Two years after the procedure, the average BCVA was similar to that of the unaffected eye of the patient. Additionally, by that time the BCVA of both eyes of the patient was only nonsignificantly lower than that of the control group. Nevertheless in the affected eyes of two patients the BCVA remained low, in the first 0.59 and in a second 0.63, which was consistent with lower levels of CS at high spatial frequencies (13, 31). In the quiescent phase of CSC (the period after the settling of previously detached neuroepithelium), the defects in the photoreceptor layer might have been the cause of the persistently decreased BCVA (6, 29) even though FAG and classical OCT do not show a variation from normal findings.

Another easily evaluated parameter of the eyes’ distinguishing ability is examination of CS. A patient can have normal values of BCVA with high contrast optotypes but significantly abnormal with low contrast. It has been observed that the perception of contrast was disturbed only in high angle critical details and not in low ones. Therefore, introduction of another variable (contrast) into the examination of visual functions can be very useful (31). At the beginning of CSC, the average CS of both eyes of the patient was greatly decreased compared to the control group in our study. The decrease was greater in the affected eyes. The study of Lu (18) also showed that 40% of the fellow eyes of the patient group showed a slight CS decrease at mid-high spatial frequencies, although other visual functional tests showed no abnormality. These fellow eyes may be at risk for CSC. Within two months after the procedure, CS values of the affected eye rapidly increased, and the final CS of both eyes of patients reached normal values, except for a spatial frequency of 3.69 c/deg where the values were significantly lower in the affected eyes. Khosla et al. (12) followed up the changes in BCVA and CS of patients after laser photocoagulation and noted that 3 months after treatment, BCVA rapidly improved while CS in many cases remained decreased even although the BCVA was 1.0. They also noted that a follow-up period of three months is too short for the evaluation of whether the decrease in CS returned to normal values.

Robertson and Ilstrup (30) showed a statistically significant reduction of recurrences within a 1,5 years follow-up period when laser photocoagulation was used. This was also verified by a long-term study by Yap (37). Contrast sensitivity did not seem to recover in all eyes after CSC even if visual acuity had returned to normal. Maaranen (20) therefore points out that contrast sensitivity testing is recommended for patients complaining of visual impairment in spite of good visual acuity.

OCT is an objective means for quantitative assessment of central retinal thickness. The significant decrease and normalization of the retinal thickness of affected eyes at one-year follow-up is consistent with a complete resolution of subretinal fluid in all our patients. Similar results were described by Iida et al. (11). On the other hand, Moschos et al. (26) described that retinal thickness remained abnormal despite of total resolution of subretinal fluid.

Two-years after laser coagulation, we detected colour vision defects in the blue area in only 14.2% of affected eyes; all fellow eyes had normal colour discrimination. In our study, the percentage of eyes with dyschromatopsia was smaller than those published by other authors. Bartos et al. (2) found long-term colour vision deterioration in 20% of patients after a single attack of CSC after conservative treatment or laser coagulation. Maaranen (20) found long-term defects in the blue region in 67% of affected eyes and in 49% of contralateral eyes of patients even if VA had recovered to normal, which may be due to earlier subclinical CSC as the authors speculated. However dyschromatopsia and relative scotomas can occur as a result of defects in the photoreceptor layer even after resolution of subretinal fluid despite normal fundoscopic, FAG and classical OCT findings (27, 29).

Our findings of normal photopic full-field ERG responses in both eyes of all patients ruled out severe peripheral retinal damage in CSC and are thus consistent with the majority of published data (5, 21). The mfERG amplitudes were normal in 86% of affected eyes and in 100% of the fellow eyes of our patients. In 14% of affected eyes (2 eyes) we detected low amplitude with retinal functional impairment in one eye in the central 14° and in the second eye in the whole central 60° region. Fundus changes in the retinal pigment epitheliopathy, however, were apparent in both patients only in the central - 15° area. In these eyes we also found abnormal colour discrimination and lower BCVA.
and CS at high spatial frequencies as mentioned above. There were no fundus abnormalities in the other 12 affected eyes and all 14 fellow eyes of patients. The implicit times of mfERG response were significantly longer in the parafoveal retinal region between 1.8° and 7° eccentricity and within the normal range in the rest of stimulated retina in affected eyes and in the whole posterior pole of fellow eyes of patients. The alteration of the implicit time in the parafoveal area in resolved CSC could be caused by alteration of second-order kernel responses due to inner retinal dysfunction (14).

Conclusions

We conclude that there was an incomplete resolution of functions in central part of the retina in affected eyes with CSC. Even although the fundoscopic findings were normal in 85 % of affected eyes two years after therapy, there remained a mild alteration of CS in the middle spatial frequency (3.69 c/deg) and prolonged implicit times of mfERG remained a mild alteration of CS in the middle spatial frequency (3.69 c/deg) and prolonged implicit times of mfERG in the parafoveal area. On the other hand, we noted a full recovery of all tested parameters in the clinically unaffected fellow eyes.

References