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The pattern of retinal ganglion cell dysfunction in Leber hereditary optic neuropathy

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\section*{1. Introduction}

Leber hereditary optic neuropathy (LHON) (OMIM 535000) is a primary mitochondrial DNA (mtDNA) disorder that presents with bilateral subacute loss of central vision (Nikoskelainen et al., 1996; Yu-Wai-Man and Chinnery, 2013; Yu-Wai-Man et al., 2014). The majority of patients harbour one of three common mtDNA mutations (m.3460G > A in \textit{MTND1}, m.11778G > A in \textit{MTND4} and m.14484T > C in \textit{MTND6}) that affect complex I subunits of the mitochondrial respiratory chain (Mackey et al., 1996). Despite the universal cellular role of mitochondria, retinal ganglion cells (RGCs) within the papillomacular bundle are particularly severely affected accounting for the characteristic dense central or caecocentral scotoma in this disorder. Although the underlying pathological process is still not fully defined, this tissue specificity has been ascribed to an increased vulnerability of RGCs to both disturbed mitochondrial energy metabolism and the increased formation of reactive oxygen species (Carelli et al., 2004a; Carelli et al., 2004b; Lin et al., 2012; Sadun et al., 2000; Levin, 2015; Sadun et al., 2015). LHON shows maternal inheritance, but there is variable disease penetrance and a marked sex bias with about 50% of male carriers losing vision during their lifetime compared with about 10% of female carriers (Mackey et al., 1996).

The histopathological observation of loss of the small calibre axons that constitute the papillomacular bundle was originally observed on histopathological sections of post mortem optic nerve samples obtained several decades after disease onset (Sadun et al., 1994; Kerrison et al., 1995; Sadun et al., 2000). More recently, in vivo studies involving high-resolution optical coherence tomography (OCT) have revealed a major loss of the temporal peripapillary nerve fibre (RNFL) and macular RGC layers within 3 months of disease onset. Interestingly, pathological thinning within the macular RGC layer was an early sign that was already apparent in the presymptomatic phase (Barboni et al., 2005; Barboni et al., 2010; Akiyama et al., 2013; Zhang et al., 2014; Mizoguchi et al., 2015; Balducci et al., 2015). Following disease conversion, swelling of the peripapillary RNFL spreads circumferentially from the inferotemporal segment of the optic disc to involve the remaining quadrants, before RNFL atrophy becomes established within 3–9 months after the onset of visual loss. Hyperemia and fluctuating
mild swelling of the prepapillary RNFL can also be seen in unaffected LHON carriers without visual loss or progression to full disease conversion (Nikoskelainen et al., 1982).

The function of the papillomacular bundle may be assessed objectively using pattern reversal visual evoked potentials (PR-VEP) and pattern electroretinography (PERG). Reported PR-VEP abnormalities in LHON are consistently severe, but the utility of the PERG in LHON is more controversial and there are conflicting reports in the literature regarding the timing of responses and the sequence of losses (Nikoskelainen et al., 1977; Hrynchak and Spafford, 1994; Mashima et al., 1997; Sharkawi et al., 2012; Ziccardi et al., 2013; Ziccardi et al., 2015; Jarc-Vidmar et al., 2015). The full-field photopic negative response (PhNR) has been used to assess generalized RGC function in glaucoma and other acquired optic neuropathies (Machida, 2012; Morny et al., 2015), but there are no published studies of PhNR in LHON. The applicability of PhNR as a potential objective functional index of RGC function in LHON therefore warrants further investigation.

RGCs are classified into the three major subtypes of RGCs, namely midget, parasol and small bistratified ganglion cells, which are thought to contribute to the parvocellular, midgetcellular and koniocellular pathways, respectively. These distinct RGC populations and their associated pathways can be tested by modifying standard psychophysical measures. In general, the processing of high spatial frequency information has been linked with the parvocellular pathway whereas high temporal frequency information is thought to be integrated by the magnocellular pathway. Red-green processing and blue-yellow processing have been the subject of several recent comprehensive reviews (Lennie and Movshon, 2005; Lee et al., 2010; Dacey et al., 2014).

LHON is thought to mainly affect midget RGCs, which have the smallest calibre axons and are the predominant subtype within the papillomacular bundle, mediating visual information including high spatial frequencies and red-green chromaticity (Sadun et al., 1994; Hrynchak and Spafford, 1994; Kerrison et al., 1995; Sadun et al., 2000). In contrast, another melanopsin-expressing RGC subtype appears relatively preserved and this peculiarity likely accounts for the frequently reported mild impairment of the magnocellular pathway in unaffected LHON patients (Kawasaki et al., 2010; La Morgia et al., 2010). One previous report indicated mild impairment of the magnocellular pathway in unaffected LHON carriers (Gualtieri et al., 2008), but there are no robust data regarding the involvement of the parasol and small bistratified RGC subtypes in LHON.

In this study, we investigated the pattern of RGC dysfunction in a well-phenotyped cohort of LHON patients in both the acute and chronic phases of the disease by using a comprehensive visual electrophysiological and psychophysical assessment protocol. Our aim was, firstly, to characterise the electrophysiological responses to better define the phenotypic features of LHON and to establish the most appropriate methods for monitoring RGC function and disease progression objectively. Secondly, we used psychophysical tests of temporal, spatial and chromatic vision to investigate the relative involvement of distinct RGC populations in the pathophysiology of LHON.

2. Methods

2.1. Subjects

This was a prospective case study of 12 affected patients (A1–A12) and 9 unaffected carriers (U1–U9) harboring one of the three common mtDNA LHON mutations (Table 1). In addition, retrospective visual electrophysiological data for 5 affected LHON patients (A13-A17) were retrieved from the hospital database of Moorfields Eye Hospital, London, UK. In total, there were 4 affected female and 13 affected male patients. Affected LHON patients and unaffected LHON carriers in our cohort underwent an ophthalmological examination that included the following investigations as indicated in Table 1: best corrected visual acuity (BCVA) assessment using the Early Treatment Diabetic Retinopathy Study (ETDRS) chart; slit lamp examination; automated Humphrey visual field perimetry (Program 30-2, Humphrey Visual Field Analyzer, Model 750, Humphrey Instruments, San Leonardo, CA); and optical coherence tomography (OCT) imaging of the macula and the optic nerve head (see details below).

The normal subjects for psychophysical tests were 15 individuals aged 17 to 78 years old at the time of testing with normal BCVA and normal color vision as assessed by standard color vision tests. Only 12 of the normal subjects had their L-cone temporal contrast sensitivities measured. Written informed consent was obtained from all subjects or their guardians. The study was approved by the local ethics committees at Moorfields Eye Hospital and University College London and it conformed to the standards of the Declaration of Helsinki.

2.2. Optical coherence tomography (OCT) imaging

The Spectralis™ platform (Heidelberg Engineering Ltd., Heidelberg, Germany) was used for SD-OCT imaging of the macula and the optic nerve head. Automated segmentation and thickness analyses were performed for perifoveal volumetric retinal B-scans using the Heidelberg Engineering segmentation tool, included in the Spectralis Glaucoma Module software (version 6.0). Of the 10 retinal layers that were automatically defined and manually confirmed, the following thickness values were recorded from the four sectors of the inner ring (between 1 and 3 mm diameter) of the nine macular ETDRS subfields as described elsewhere (Majander et al., 2016): (i) retina, (ii) retinal nerve fiber layer (RNFL), (iii) combined GCL and inner plexiform layer (IPL), (iv) inner nuclear layer (INL), (v) outer plexiform layer (OPL), (vi) combined OPL and outer nuclear layer (ONL), and (v) inner retina. The thickness of the outer retinal layers was calculated by subtracting the thickness of the inner retinal layers from the total retinal thickness. Normative data was generated from SD-OCT images of 48 healthy eyes of 48 subjects (Majander et al., 2016). For peripapillary RNFL measurement a 3.5-mm-diameter circular scan centered on the optic disc was used and the data for six sectors were collected.

2.3. Electrophysiology investigations

Twelve subjects underwent electrophysiological testing including pattern reversal and flash visual evoked potential (PVEP; FVEP) and pattern electroretinography (PERG), incorporating the standards of the International Society for Clinical Electrophysiology of Vision (ISCEV; Odom et al., 2010, Bach et al., 2013). Pattern ERGs were recorded to a 0.8- degree check size using both a standard checkerboard field (12 × 15°) and additionally to a large field (24 × 30°; LF PERG; Lenassi et al., 2012). The full-field photopic negative response (PhNR) was additionally recorded in 7 cases using diffuse red flash stimulation (640 nm) at 5 flash strengths (0.5, 1.0, 2.0, 5.0 and 10.0 cd·s·m⁻²), superimposed on a blue background (450 nm; 2.25 cd·m⁻²). Gold foil electrodes were used and the results compared to normative data.

2.4. Psychophysical investigations

2.4.1. L- and S-cone critical flicker fusion and L-cone temporal contrast sensitivity

L- and S-cone temporal acuities (critical flicker fusion, CFF) and L-cone temporal contrast sensitivity functions (TCSFs) were measured using a Maxwellian-view optical system described in more detail elsewhere (Stockman et al., 2014a, 2005). Predominantly L-cone or S-cone stimuli were used for the CFF measurements. The L-cone stimulus was produced by flickering a 650-nm circular target of 4° visual angle in diameter superimposed in the center of a steady 481-nm circular background.
TCSF measurements, observers varied the contrast of the sinusoidally-from 6.5 to 10.5 log10 quanta s⁻¹ diameter. The 620-nm background radiance was superimposed in the center of a 620-nm circular background from 6.5 to 10 log10 quanta s⁻¹ background were also used for the L-cone TCSF measurements with the quanta s⁻¹ background and target for at least 2 min. The observers viewed the experiment was repeated 2 and the target radiance was varied in steps 2. Before each run the subjects were light-adapted to 1 deg 2. The 650-nm target and 481-nm S-cone stimulus was measured using a staircase procedure. Observers indicated whether or not they could detect the spatial variation in the Gabor pattern using a 2-button keypad.

### Chromatic discrimination

Chromatic discrimination was tested using the so-called trivector test procedure implemented as part of the Cambridge Color Test (CCT), Technologies Inc., Saint-Bruno, QC, Canada). The full screen subtended a visual angle of 39° × 29° at a test distance of 0.57 m. The experiments were performed at a constant mean luminance of 44.57 cd/m² as measured by a ColorCal calibration device (Cambridge Research Systems Ltd., Rochester, UK). The stimuli were horizontally-orientated Gabor patterns with spatial frequencies ranging from 0.25 to 16 cycles per degree (cpd) and with a spatial Gaussian window with a standard deviation of 6°. The target was presented for 500 ms, preceded and followed by 100 ms cosine-windowed onsets and offsets. The order of presentation was from low to high spatial frequency. Thresholds were measured using a staircase procedure. Observers indicated whether or not they could detect the spatial variation in the Gabor pattern using a 2-button keypad.

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v1.5, (Cambridge Research Systems Ltd., Rochester, UK). The test was performed using a second gamma-corrected Sony FD Trinitron color monitor (Model GDM-F500OR) connected to a VSG 2/5 visual stimulus generator (Cambridge Research Systems, Rochester, UK) with 800 by 600 pixel resolution. The CRT phosphors measured in CIE 1976 \( u' \), \( v' \) chromaticity coordinates (or CIE 1931 x, y coordinates) using ColorCal photometer (Cambridge Research System) were: red phosphor (R) \( u' = 0.416; v' = 0.522 \) (x = 0.610, y = 0.340); green phosphor (G) \( u' = 0.117; v' = 0.559 \) (x = 0.280, y = 0.595); blue phosphor (B) \( u' = 0.159; v' = 0.177 \) (x = 0.142, y = 0.070). The visual stimuli consisted of Landolt “C” targets that varied in orientation presented on a background of neutral chromaticity (CIE 1976 coordinates \( u' = 0.1977, v' = 0.4689 \)). The background and the target were made up of small disks of variable size and luminance (ten equal steps between 8.0 and 18.0 cd·m\(^{-2} \)). The circles making up the Landolt C were up of small disks of variable size and luminance (ten equal steps be-

The age of the patients at the time of electrophysiological testing ranged from 7.5 to 59.0 years (mean = 26.6 years; SD = 16.7 years).

The duration of symptoms ranged from 0 to 27 months (mean 12.5 months). Visual acuity was severely impaired in all cases (range = 1.3 logMAR to perception of light). Characteristic electro-physiology waveforms for affected LHON carriers are shown in Fig. 2. Photopic negative responses were attenuated at all stimulus strengths (see corresponding plots in Fig. 2c and d; orange circles). Pattern reversal VEPs were undetectable and flash VEPs were grossly abnormal. Pattern ERG P50 is of short peak time and the waveforms had a low N95:P50 ratio, in keeping with severe optic nerve/retinal ganglion cell dysfunction bilaterally.

Pattern reversal VEPs were undetectable in 20 of 24 eyes and were delayed in 4 eyes of 4 subjects with detectable responses (peak time 121-142 ms; upper limit of normal 115 ms); amplitudes in 2 of the eyes with a delayed VEP were within the normal range (> 4 μV). Flash VEPs were undetectable in 5 eyes of 3 subjects (all with undetectable pattern VEP), were delayed (< 140 ms) in 5 eyes, and were present but subnormal (< 5 μV) in 20 eyes. Flash VEPs were normal in 3 eyes of 2 subjects including one of the youngest individuals (age 7.5 years). The severity of pattern and flash VEP abnormality did not significantly correlate either with age or duration of symptoms.

Pattern ERG P50 to a standard stimulus (field size 12° × 15°) was of abnormally short peak time (< 45.5 ms) in 19 of 21 eyes (Fig. 3a PERG). P50 was of normal amplitude in most, mildly subnormal (< 2.0 μV) in 4 eyes of 4 patients (1.5–1.9 μV) and of borderline amplitude (2.0 μV) in a further 2 eyes (Fig. 3b PERG); P50 was of shortened peak time in all 6 eyes with a borderline or mildly subnormal amplitude. The N95:P50 amplitude ratio was subnormal (< 1.1) in 17 of 19 eyes (0.75–1.0) and at the lower limit of normal (ratio = 1.1) in a further 2 eyes (mean and median ratio of all cases 1.0; SD 0.1; Fig. 3c PERG). PERGs were technically poor in 5 eyes of 4 cases due to variable fixation (N = 1); pupil dilation (N = 1) or physiological noise and those eyes were excluded from analysis. PERG to a doubled stimulus field (24° × 30°) were obtained in 11 patients including 4 eyes in which the standard field PERG was excluded. Data were compared with a normative data set obtained using the same large field stimulus. Sixteen of 22 eyes were of abnormally short peak time (< 46.5 ms; Fig. 3d PERG) and these included 4 patients with additional bilateral P50 reduction (1.9 μV–2.7 μV; normal > 3.9 μV). The N95:P50 ratio was
subnormal in 8, borderline in 10 and normal in 5 eyes (Fig. 3e PERG). There was no significant correlation between pattern ERG parameters and age or duration of symptoms.

International-standard full-field ERGs revealed no clinically significant abnormality in the 7 individuals tested. PhNRs were recorded from 14 eyes of 7 cases, including 5 individuals that did not undergo standard full-field ERGs. The ratios of PhNR/b-wave ratios were within the normal range in 1 eye at all 5 flash intensities; in others the ratio was subnormal to one flash strength (N = 2 eyes), two (N = 3 eyes), three (N = 1 eye), four (N = 4 eyes) or 5 flash strengths (N = 3 eyes) (Fig. 4 PhNR). The mean magnitude of reduction of subnormal responses was 13% (range 1.5–23%) compared with the lower limit of normal. Forty percent of responses showed no abnormality (Fig. 4c and d).

Fig. 2. Photopic negative responses from the right (a) and left (b) eyes of a patient with LHON (A3). For comparison, representative normal recordings over a range of flash strengths (0.5–10.0 unit) have been provided (c). Pattern reversal VEPs, flash VEPs and pattern ERGs are shown for the right (d) and left (e) eyes of Patient A3 compared with representative normal recordings (f). The patient’s recordings have been superimposed for all the parameters tested to demonstrate reproducibility.
3.3. Visual psychophysics

3.3.1. L-cone temporal functions were severely compromised in LHON

The data for L-cone \( \text{cfr} \) (temporal acuity) are presented in Fig. 5 and are summarized in the Supplementary Table S2. In the normal observers (grey triangles), flicker was first seen at a target radiance of 6.6 log 10 quanta s \(^{-1}\) deg \(^{-2}\). The \( \text{cfr} \) then grew linearly with log radiance over about 3 log 10 units with a slope of 9.2 Hz per log 10 unit (see upper blue line in the lower panel) until reaching a plateau near 40 Hz (Stockman et al., 2014a). The linear relationship between \( \text{cfr} \) and log radiance is known as the Ferry-Porter law (Ferry, 1892; Porter, 1902). Eight of the 11 affected LHON carriers were able to detect L-cone flicker, but the mean radiance of 9.6 log 10 quanta s \(^{-1}\) deg \(^{-2}\) at which flicker was first seen was 30 times higher than that for normal observers (Table S2). Three patients could detect flicker only at the highest target radiances. LHON patients thus required higher radiances to detect flicker than normal subjects and their \( \text{cfr} \)s showed severe losses reaching a mean plateau of only 12 Hz, 28 Hz less than normal subjects. The lowest radiance at which unaffected LHON carriers first detected flicker was slightly higher than that for normal subjects (Table S2). Three carriers (U2, U6, U9) had significantly shallower Ferry-Porter slopes than normal and only three carriers (U1, U4, U8) had \( \text{cfr} \)s that reached the normal plateau level of 40 Hz. The majority of the LHON carriers showed some loss of \( \text{cfr} \). Patient A6 experienced spontaneous recovery of BCVA from 1.6 logMAR to 0.1 logMAR between 6 and 18 months after LHON onset. His \( \text{cfr} \) improved, but only marginally (A6 F, yellow squares).

Fig. 6 shows the L-cone temporal contrast sensitivity data. The mean L-cone TCSF for normal subjects peaks in sensitivity near 8 Hz and falls off in sensitivity at both low and high temporal frequencies (Stockman et al., 2014a, 2014b; Ripamonti et al., 2014). Only patients A3, A7 and A9 of the affected carriers were able to perform the test and showed a mean sensitivity loss of 1.0 log 10 unit. Unaffected LHON carriers had normal L-cone temporal modulation sensitivities but were unable to make settings at the highest temporal frequencies, consistent with their lower \( \text{cfr} \)s.

3.3.2. S-cone temporal function was unmeasurable in most LHON patients and reduced in the unaffected m.3460G > A carriers

In the normal subjects, flicker was first seen at a target radiance of about 6.5 log 10 quanta s \(^{-1}\) deg \(^{-2}\). Thereafter, the \( \text{cfr} \) grows linearly with log radiance with a slope of 7.3 Hz per log 10 unit, consistent with the Ferry-Porter law, until reaching a plateau at 9.0 log 10 quanta s \(^{-1}\) deg \(^{-2}\) (see Fig. 7 and the Supplementary Table S3). The rise after 9.9 log 10 quanta s \(^{-1}\) deg \(^{-2}\) is due to M-cone intrusion - the M-cones become more sensitive than S-cones at high radiances and thus mediate flicker detection (Stockman and Plummer, 1998; Stockman et al., 2014a).

Only two affected LHON carriers (A7, A9) were able to detect any 440-nm flicker, and all showed severely impaired temporal acuity. S-cone \( \text{cfr} \) was relatively normal for the unaffected carriers of the m.11778G > A and the m.14484T > C mutations, who first saw flicker at 07.10 ± 0.06 (mean ± 1 SEM) log 10 quanta s \(^{-1}\) deg \(^{-2}\), and exhibited a Ferry-Porter slope of 7.97 ± 0.39 Hz per decade (mean ± 1 SEM) and a \( \text{cfr} \) plateau frequency of 22.94 ± 0.57 Hz (mean ± 1 SEM). In contrast, S-cone \( \text{cfr} \) for the three carriers harboring the m.3460G > A mutation was compromised: those subjects first saw flicker at a radiance of 8.06 ± 0.51 log 10 quanta s \(^{-1}\) deg \(^{-2}\), and exhibited a shallow Ferry-Porter slope of 5.03 ± 1.05 Hz per log 10 unit, the \( \text{cfr} \) reached a plateau frequency of 12.61 ± 3.80 Hz. One carrier of
the three with the m.3460G > A mutation could detect flicker only at the highest radiances, but was within normal limits for the tritan measurements in the CCT (see below).

3.3.3. Achromatic spatial contrast sensitivity function (SCSF) was unmeasurable in most LHON patients and mildly subnormal in the unaffected carriers

The mean achromatic spatial contrast sensitivity function for normal subjects is shown as inverted grey triangles in each of the three panels of Supplementary Fig. S1. The function is band-pass in shape peaking at 2 cpd and falling off in sensitivity at lower and higher spatial frequencies characteristic of other SCSFs measured at moderate and high intensities (Robson, 1966; van Nes and Bouman, 1967). Only two (A7 and A9) of the 11 affected LHON carriers could perform this test. Both patients showed a loss of contrast sensitivity that increased with frequency. In the unaffected LHON carriers, the achromatic SCSF was normal at the lowest spatial frequency but showed increasing loss as spatial frequency increased.

3.3.4. Chromatic discrimination was severely affected in LHON

Fig. 8A shows the vector lengths of the three confusion lines (proton, deutan, and tritan) of the Cambridge Color Test in $10^{-4}$ u\'v\' units (the CIE 1976 u\'v\' color space). The longer the vector length the more saturated the color had to be for the orientation of the gap in the Landolt C to be discriminated. The maximum vector lengths were 1600 $10^{-4}$ u\'v\' units (the CIE 1976 u\'v\' color space). The vector lengths in the normal observers were comparable to those previously reported using standard targets with 1' gaps: 45.1 ± 1.0 for proton, 43.3 ± 0.8 for deutan and 51.5 ± 1.3 for tritan in $10^{-4}$ u\'v\' units (the CIE 1976 u\'v\' color space) [mean ± SEM] (Paramei and Oakley, 2014). Altogether, 8 affected LHON carriers could discriminate the gap in the Landolt C, most viewed the CRT screen from only 5 to 10 cm Their vector thresholds along the proton and the deutan axes on their first tests were 25.8 ± 2.6 (mean ± SEM) and 25.4 ± 2.8 times higher, respectively, than controls whereas along the tritan axes the vectors were only 5.1 ± 0.7 times higher. Fig. 8B shows the vector lengths of the successive tests made on six affected LHON carriers. Three affected carriers (A1, A3, A6), initially studied 6 to 9 months after LHON onset, showed recovery of color discrimination, mainly along proton and deutan axes. Chromatic discrimination in unaffected LHON carriers was only marginally subnormal along all chromatic axes with high normal vector lengths. Affected observers therefore show a general loss along all three axes.

4. Discussion

Our comprehensive electrophysiological and psychophysical study of patients with acute and chronic LHON highlights the marked extent of RGC dysfunction in this classical mitochondrial optic neuropathy. Our study also indicates the utility of the standard transient pattern ERG technique in patients with LHON, with the recordings revealing severe central RGC dysfunction in all the patients tested, irrespective of visual acuity, age or the duration of symptoms. Detailed psychophysical data demonstrated loss of visual function in affected LHON carriers across a range of parameters, including achromatic spatial and L-cone temporal contrast sensitivity, L- and S-cone acuity, and chromatic discrimination. These findings are consistent with substantial losses of the principal RGC subtypes associated with each of the three major retinal pathways. Furthermore, with the exception of L-cone TCSF, unaffected LHON carriers also showed subclinical abnormalities in all the psychophysical measures that were evaluated.

A number of studies have reported abnormal cortical PR-VEPs responses in acute LHON, (Nikoskelainen et al., 1977; Hrynchak and Spafford, 1994; Mashima et al., 1997; Sharkawi et al., 2012; Ziccardi et al., 2013). Our PR-VEP results are consistent with these previous observations and the early involvement of the papillomacular bundle in...
this mitochondrial optic nerve disorder. The PERG reflects central retinal function and has two major components; the positive P50 and negative N95 (Holder, 1987). The N95 component arises in RGCs whereas approximately 30% of the P50 component originates in more anterior retinal structures (Holder, 1987, 2001; Ryan and Arden, 1988; Viswanathan et al., 2000). P50 is widely used to assess macular anterior retinal structures (Holder, 1987, 2004; Ryan and Arden, 1988) (pink, blue and green triangles, respectively) and for the mean of 9 unaffected LHON carriers (yellow triangles). The mean TCSFs for 12 normal observers have been shown as grey triangles. The error bars represent ± 1 SEM either between runs for the individual patients or between subjects for the mean data. The mean difference in log sensitivity between each affected or the mean of unaffected LHON carrier and normal subjects are shown in the lower part of each panel (circles). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

The photopic negative response (PhNR) component of the full-field electrotetrogram is severely reduced in primates treated with tetrodotoxin and in experimental models of glaucoma, in keeping with a possible RGC origin (Viswanathan et al., 1999). The PhNR has also been shown to be attenuated in experimental and clinical studies of glaucoma and in other forms of optic neuropathies (Viswanathan et al., 1999; Gototh et al., 2004; Sustar et al., 2009; North et al., 2010; Preiser et al., 2013; Machida, 2012; Morny et al., 2015). The majority of affected LHON patients in our study had responses near the limits of normal, with a substantial subgroup (40%) being within normal limits. There is no published data describing PhNRs in LHON, but mildly abnormal or borderline reductions in the PhNR have similarly been reported in dominant optic atrophy (Miya et al., 2007). It is notable that both these mitochondrial optic neuropathies predominantly affect the papillomacular bundle with relative sparing of RGC axons in the retinal periphery (Johnston et al., 1979; Kerrison et al., 1995; Kjer et al., 1983; Sadun et al., 1994; Sadun et al., 2000). It should be noted that the full-field PhNR provides a measure of global RGC function and it could therefore be less sensitive in detecting focal RGC loss or dysfunction compared with the pattern ERG, which arises largely in central macular RGCs. Other methods such as focal PhNRs, involving flash stimulation of the central macular area, could potentially offer better sensitivity compared with full-field photopic negative responses, but with the disadvantage of generating comparatively smaller signals. These findings are pertinent to clinical trials of future therapeutic interventions, since an undetectable VEP cannot be used to monitor safety and PERG could prove the objective test of choice to monitor function of the most affected RGCs.

The international standard VEP and PERG stimuli involve high contrast checkerboard reversal stimuli that are suprathreshold, whereas psychophysics enables measurement of detection threshold. Affected LHON carriers in our study had impaired chromatic resolution in the protan and deutan axes. Their achromatic spatial sensitivity was also severely compromised. Only the two least affected patients (A7 and A9) were capable of performing the achromatic spatial CSF measurements, but even then, steep sensitivity losses with increasing spatial frequency were evident. These findings are consistent with the severe loss of the midget RGCs and the small calibre RGC axons found in post mortem histological studies (Sadun et al., 1994; Kerrison et al., 1995; Sadun et al., 2000). The majority of LHON patients in our study were able to detect a flickering long-wavelength light at high luminance levels, but with markedly reduced temporal resolution, in keeping with a severe deficit of magnocellular function. This previously unreported observation is pathologically relevant as it implies that the loss of parasol RGC function is also present in acute LHON, which is consistent with the LGN pathology reported in end-stage disease (Rizzo et al., 2012). Previous studies that have assessed the koniocellular pathway in LHON have been limited to color vision tests (Nikoskelainen et al., 1977). Only the least severely affected patients (A7 and A9) in our study were able to detect flickering short-wavelength light and additional tritan deficits in the CCT tests suggested severe koniocellular involvement. The smaller proportional loss found along the tritan axis compared with the protan and deutan axes in the CCT test of affected LHON carriers is of doubtful clinical significance, since the unusually close viewing distances adopted by 5 of the observers might have selectively reduced the tritan thresholds due to rod intrusion, retinal inhomogeneities or scatter. In addition, melanopsin-expressing RGCs, which are relatively preserved in LHON (Kawasaki et al., 2010; La Morgia et al., 2010), are preferentially sensitive to short-wavelength light (Lucas et al., 2014) and, in theory, may also contribute to blue color discrimination.

Fig. 6. Log10 L-cone temporal contrast sensitivity. Log10 L-cone TCSFs were measured using a sinusoidally-modulated 650-nm target with a time-averaged mean radiance of 10.28 log10 quanta s−1 deg−2 superimposed on a 481-nm background of 8.29 log10 quanta s−1 deg−2 plotted as a function temporal frequency (logarithmic axis) for three affected LHON carriers (A3, A7, A9) (pink, blue and green triangles, respectively) and for the mean of 9 unaffected LHON carriers (yellow triangles). The mean TCSFs for 12 normal observers have been shown as grey triangles. The error bars represent ± 1 SEM either between runs for the individual patients or between subjects for the mean data. The mean difference in log sensitivity between each affected or the mean of unaffected LHON carrier and normal subjects are shown in the lower part of each panel (circles). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Previous studies on asymptomatic LHON carriers harboring the m.11778A > G mutation have reported subclinical visual impairment involving both the parvocellular and the magnocellular pathways as revealed by subtle chromatic and luminance contrast sensitivity deficits and impaired temporal processing (Ventura et al., 2005; Sadun et al., 2006; Ventura et al., 2007; Gualtieri et al., 2008; Mateus et al., 2015).

Our study cohort included unaffected carriers with the m.3460A > G and m.14484T > C LHON mutations, in addition to m.11778A > G. Our data indicate that all three common mtDNA LHON mutations significantly impair achromatic spatial contrast sensitivity worst for higher spatial frequencies. The chromatic thresholds along all three confusion lines showed mild losses compared with normals highlighting dysfunction of the parvocellular pathway and confirming a previous report using similar methods (Mateus et al., 2015). The majority of LHON carriers also showed abnormalities of the long-wavelength temporal visual acuity, but with relatively minimal loss on the L-cone TCSF measurements. These parvocellular and magnocellular related losses were present with all three common mtDNA LHON mutations. Unlike the long-wavelength sensitive temporal visual acuity, impairment of the short-wavelength temporal acuity was limited to the three unrelated unaffected carriers harboring the m.3460A > G mutation.
In conclusion, our study highlights the extent and severity of diffuse and focal electrophysiological measures of RGC dysfunction in LHON. PERG abnormalities in LHON are largely independent of age and can be elicited in patients with severely impaired visual acuity. Furthermore, psychophysical tests of achromatic and chromatic visual function suggest severe involvement or loss of midget, parasol and bistratified RGCs. These findings are highly relevant for the design of future clinical trials aimed at assessing therapeutic interventions and the viability of specific RGC subpopulations in patients affected with LHON.

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