

Macular telangiectasia type 2: quantitative analysis of a novel phenotype and implications for the pathobiology of the disease

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Retinal microcystoid spaces in MacTel 2

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Key words

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Summary Statement

Retinal microcystoid spaces are a novel phenotype of Macular Telangiectasia (MacTel) type 2 on optical coherence tomography. It presents a distinct form of cystoid macular oedema with the pathobiology suggesting possible Müller cell involvement in this disorder.

Abstract

Purpose

To investigate retinal microcystoid spaces in macular telangiectasia (MacTel) type 2 with spectral-domain optical coherence tomography.

Methods

Retrospective review of 135 patients enrolled in the MacTel Natural History Observation and Registry Study at Moorfields Eye Hospital, United Kingdom. 172 eyes from 86 patients who had a comparable scan protocol of at least 30 μ m interval were included for analysis. Retinal microcystoid spaces were identified, segmented and metrics analyzed.

Results

From 172 eyes of 86 patients, microcystoid spaces were found in 11 eyes (6.4%) from 8 patients (9.3%). The mean number of microcystoid spaces per eye was 12.9 ± 18.2 . The majority were located in the inner nuclear layer. The inferonasal quadrant of the macula was the least commonly affected region. Microcystoid spaces were distributed entirely within the assumed MacTel area on blue light reflectance imaging in all but two eyes (4 of 142 microcysts). Median diameter of the microcystoid spaces was 31 μ m (range 15 to 80 μ m).

Conclusion

Microcystoid spaces as a phenotype of MacTel should be considered in the differentials for microcystic oedema. Understanding the pathogenesis of these lesions may provide further insight into the role of Müller cell dysfunction in this disorder.

Introduction

Macular telangiectasia type 2 (MacTel) is a rare degenerative disorder of the central retina. The disease causes progressive central vision loss and is typically diagnosed in patients between the age of fifty to sixty, who describe problems with reading vision often for years prior to measurable visual acuity changes.¹⁻³

The exact cause of the disease remains unclear. Although the early description of the disease by Gass and Oyakawa suggested an underlying vascular aetiology, ⁴ it is now thought to be more consistent with a neurodegenerative process. Recent histopathological studies have demonstrated a striking depletion of Müller cells in the central retina in the same distribution as macular pigment reduction, pointing towards neuroglial loss as critical in this disorder, leading to structural and functional impairment in cone and rod photoreceptors. 5–8

In recent years, optical coherence tomography imaging (OCT) has become an increasingly valuable diagnostic tool, providing further phenotyping of the disease. Characteristic findings include hyporeflective spaces of the inner and outer neurosensory retina, ellipsoid zone (EZ) disruption, hyperreflective lesions from pigment migration, as well as foveal atrophy in late stage disease. 9–13 Typically the central macular retinal thickness remains normal or may be reduced despite presence of angiographic leakage.¹⁴

In addition to these imaging findings, optical coherence tomography angiography (OCTA) has also recently identified the presence of small retinal hyporeflective spaces in MacTel patients. Termed 'microcavitations', these spaces were defined as

an optically empty space less than 250 µm in dimension and were frequently distributed around right-angle venules but were also present in areas without obvious vascular changes on OCTA.¹⁵ The microcavitations were mainly located in the inner retina, predominantly in the ganglion cell layer (GCL), inner nuclear layer (INL) as well as Henle's fibre layer. Similar microcavitations, described with varying terminology by others, have also been identified in a case report using light microscopy ¹⁶, as well as on adaptive optics scanning laser ophthalmoscopy (AOSLO) (meeting abstract). 17

Aside from MacTel, microcystoid changes of the macula have been described in a variety of hereditary or acquired optic neuropathies and predominantly occur in the INL on OCT.¹⁸⁻²¹ These lesions are not observed in normal eyes and might correspond to the microcystoid spaces seen in MacTel. The presence of microcystoid spaces in patients with MacTel type 2 may further provide clues to the pathogenesis of the disorder. In this study, we present a quantitative analysis of the metrics and distribution of microcystoid spaces with spectral-domain OCT (SD-OCT).

Methods

Study Design

This was a retrospective review of the OCT images of 135 patients diagnosed with MacTel type 2 at Moorfields Eye Hospital, London UK and who were enrolled in the MacTel Natural History Observation and Registry study. All patients had fundoscopy, SD-OCT, blue light reflectance (BLR) imaging, dual wavelength autofluorescence for macular pigment optical density (MPOD) mapping and fluorescein angiography as part of the study protocol.²² Only patients who had SD-OCT volume scans with a minimum of 30 μ m interslice distance were included for analysis. Exclusion criteria were any clinical or electrodiagnostic evidence of concomitant optic nerve disease, diabetic retinopathy, and poor imaging quality. Retrospective OCT data from eight eyes of four healthy controls were also included for comparative purposes. Two graders (PM and MO), masked to patient identifying details and disease status, graded all scans meeting eligibility criteria for the presence of visible retinal microcystoid spaces. Macular microcystoid spaces were defined in this study as small hyporeflective spaces that were non-contiguous with typical larger inner or outer retinal cavitations. The terminology 'microcystoid space' is preferred here as it applies to cystoid-like changes seen on imaging. The study was approved by the local institutional review board and conducted according to the tenets of the Declaration of Helsinki. Informed consent was obtained from all study participants.

OCT and BLR Imaging and Image Analysis

OCT imaging was acquired using Heidelberg Spectralis® HRA2 system (Heidelberg Engineering, Heidelberg, Germany). The scan pattern for patients was between 3.8 x 2.5 x 1.9 mm to 4.4 x 2.9 x 1.9 mm. B-scans ranged from 49 to 261 scans per volume, with imaging averaging for $8 - 12$ scans and an interslice distance of 11-30 μ m. The scan pattern for all healthy controls was raster lines $4.5 \times 3.0 \times 1.9$ mm, 261 B-Scans, interslice distance of 11 um, averaging 20 scans using the automatic averaging and tracking feature.

In patients where microcystoid spaces were visually identified, segmentation of all hyporeflective cystoid spaces present in that scan was performed using imageprocessing software AMIRA 6.1 (Materials & Structural Analysis, Merignac, France). Microcystoid spaces were segmented through thresholding of pixel intensity. A threshold from 0 to 50 grey-scale units (scale 0 to 255) showed reasonably wellseparated lesions, further identifying smaller microcystoid spaces that were not visible initially on manual identification. Image artifact from vessel shadowing was excluded. Microcystoid volumes were also measured. Based on a presumed sphere model, data for diameter and surface area were calculated. Average central foveal thickness was recorded using the central area on the Early Treatment Diabetic Retinopathy Study grid area on Heidelberg Eye Explorer viewing module software.

In patients with microcystoid lesions, the location of each lesion in relation to individual retinal layers was recorded on the B-scans according to the International Nomenclature for OCT consensus.²³ The spatial distribution of the microcystoid spaces was also compared to the theoretical MacTel area as seen on BLR imaging, which has been shown to correspond to the area of macular pigment loss as measured by dual wavelength autofluorescence²⁴ and on histology.⁶ For this purpose, open source imaging software ImageJ (v1.467 (ref - Rasband, W.S., ImageJ, U. S. National Institutes of Health, Bethesda, Maryland, USA, https://imagej.nih.gov/ij/, 1997-2016) was used to overlay the patient's scanning laser ophthalmoscopy image (SLO) using

the metric of the SLO image and transferred onto the aligned BLR image with Adobe Photoshop CC 2017 (Adobe Systems, San José, United States). Finally, identified microcystoid spaces were exported and overlaid on the patient's BLR images with AMIRA software to analyse the distribution with respect to the MacTel area and the distribution in quadrants centred on the fovea.

Results

One hundred and thirty-five patients were enrolled at Moorfields Eye Hospital in the MacTel Natural History Observation and Registry study. Of these, 86 (64%) patients or 172 eyes had at least one volume scan meeting the 30 μ m minimum interslice inclusion criteria. In total, 11 eyes (6.4% of eligible eyes) from 8 patients were identified as having microcystoid spaces as assessed by two masked graders. The demographic and clinical details of the patients are described in Table 1. Mean age of the patients was 53.1 years (range $37 - 71$ years) and mean visual acuity was 0.16 LogMAR units (range 0.0 – 0.78 LogMAR units; equivalent to 20/28 Snellen acuity). Average central foveal thickness using the ETDRS grid was normal in all eyes (mean $249 \pm 22 \mu m$). None of the patients had pigment plaques or evidence of subretinal neovascularisation. One of the patients had highly asymmetric disease with typical BLR and macular pigment changes in the affected eye but no clinical or imaging features of MacTel in the fellow eye.

When present, the mean number of microcystoid spaces per eye was 12.9 ± 18.2 (range 1 to 61 microcystoid spaces) with a total 142 microcystoid spaces across all 11 eyes. None of the control eyes $(n = 8)$ had microcystoid spaces on OCT imaging

either by manual grading with two masked graders or with image processing segmentation using the same threshold criteria. Median volume of the microcystoid spaces was 15,146 μ m³ (range 1,829 to 27,0724 μ m³) (Figure 1), median surface area was 2,960 μ m³ (range 723 to 20,238 μ m²) and median diameter was 31 μ m (range 15 to 80 µm). Eighty-three percent of the assessed lesions had a volume between 16,000 and $44,000 \mu m^3$.

Location of the Microcystoid Spaces within Retinal Layers

Each microcystoid space was assessed according to the predominant retinal layer or zone it was located within on B-scan segmentation. The majority were located within the INL $(42\%, n = 59)$ followed by the outer plexiform layer (OPL) and GCL (Figure 2 and 3). No microcystoid spaces were found posterior to the external limiting membrane.

Distribution of Microcystoid Spaces

Microcystoid spaces were seen in all quadrants of the macula however the inferonasal quadrant was the least common quadrant (inferonasal: 15% versus superonasal: 35%). There was no difference across the vertical meridian with an equal number of microcystoid spaces in the temporal $(50\%, n = 71)$ as compared to the nasal half of the macula. There was no direct correlation between the location of microcystoid spaces and the areas of early hyperfluorescence on fundus fluorescein angiogram.

The microcystoid spaces were distributed within the parafoveal region of the macula (Figures 3 and 4). The spatial location of the microcystoid spaces was analysed and compared to the assumed MacTel area for each patient as defined by the area of

increased reflectance on BLR imaging. In 9 of 11 eyes (82 %) the microcystoid spaces were all entirely contained within the MacTel area. In one eye, the peripheral microcystoid spaces (2 of 61 lesions in that eye) were located temporally, slightly outside the MacTel area (Figure 4). In the second eye, the peripheral lesions (2 of 7 microcystoid spaces in that eye) were in the inferonasal area of the perifoveal region where there was no increased BLR.

Discussion

This paper confirms and extends previously published reports^{15,16} of the presence of macular microcystoid spaces in a separate patient cohort with MacTel type 2 and the findings are using a different imaging modality (SD-OCT rather than OCTA). Furthermore, this study provides a novel quantitative analysis and a detailed examination of the distribution of the microcystoid spaces with respect to the socalled MacTel area, defined here as the area of increased parafoveal BLR. We demonstrate that the microcystoid spaces were concentrated in the INL and OPL, occurring in an oval parafoveal region that largely correlated with the MacTel area. Interestingly, a small number of the lesions extended slightly beyond this hypothetical MacTel area boundary. This might point towards MacTel as a more extensive macular disease than previously thought. It may also suggest that these microcystoid spaces can show up earlier than macular pigment changes as seen in the patient with microcystoid spaces in the inferonasal area where there was sparing on BLR and macular pigment optical density mapping. However, it is also possible that these more peripheral microcystoid spaces are different from the ones within the MacTel area and may even be present in normal aged eyes. We did not see them however in our normal

healthy controls or in the fellow unaffected eye of the patient with asymmetric disease.

Microcystic cavities in MacTel type 2 were first described in a histopathological case report in 1980, when the disease was still described as 'parafoveal retinal telangiectasis'. ¹⁶ Using light and electron microscopy, the authors examined an eye with features of MacTel type 2 which had undergone exenteration, and found 'microcystic changes' in the temporal macula. Microcystic cavities were predominantly noted in the INL and OPL, and were associated with thickening of the temporal macula up to 3mm distance from the fovea. Although histology and OCT are difficult to compare, the retinal layer location and distribution of these 'microcystic cavities' are similar to the microcystoid spaces described on retinal imaging in our study and also reported by Spaide and colleagues.¹⁵

On microscopy the microcystic cavities reported by Green et al. were not empty spaces but rather contained lightly-staining fibrillar material. ¹⁶ Microcystic cavities however were not evident in more recent histological studies.^{5,6} This may be due to differences in disease stage of the patients, which were more advanced in these latter reports. Larger cystoid cavities have been identified but were structurally empty on haematoxylin-eosin staining.

It is also possible that the microcystoid spaces seen here on OCT correspond to the inner retinal 'spherical microcysts' in MacTel type 2 on AOSLO (ARVO abstract).¹⁷ The microcysts described ranged from 30 to 100 μ m in size and were all located within the inner retina, although detail of specific retinal layer involvement was not

provided. Interestingly, the authors found spherical microcysts in 7 of 14 patients, also with early stage disease. Assuming these are similar entities, they found a much higher prevalence. This may be due to differences in the resolution of the imaging method, with our cohort including patients with minimum 30 μ m interval scans, and smaller microcystoid spaces may have been missed. This is also possible given the median diameter of microcystoid spaces in this present study was 31µm size only. Interestingly, the patient with 11 μ m interval scans did not have a higher rate of microcystoid spaces so other factors may be relevant in addition to resolution scan interval.

The microcystoid spaces in MacTel may also be similar in mechanism to the 'macular microcysts' involving the INL in a variety of optic neuropathies. 18,21,25,26 The mechanism for this is still unclear, as conventional teaching would suggest that optic neuropathy should result in GCL thinning alone due to axonal injury. One proposed theory to explain the INL involvement in optic neuropathy entails retrograde synaptic degeneration of optic nerve axons with secondary degeneration of Müller neuroglial cells. 18,20,26 Vitreomacular adhesion with or without traction may also contribute to inward exerting forces on the INL.²⁶⁻²⁸ Given the key finding of central Müller cell depletion in MacTel on histopathology,^{5,6} it is possible that Müller cell dysfunction may also be responsible for the microcystoid disease phenotype presented here. Müller cells span the entire neuroretina and are important not only as a scaffold but also in the osmotic regulation of the retina.²⁹ Further evidence of this is suggested by the presence of INL microcystic changes in 20% of patients with neuromyelitis optica (NMO) who had a history of previous optic neuritis.²¹ Interestingly, retinal Müller

cells are rich in aquaporin-4 water channels, with antibodies against these channels implicated in the immunopathogenesis of NMO-spectrum disorder.

Microcystoid spaces in this present study were distributed in the parafoveal region with the inferior-nasal quadrant of the macular being the least common quadrant to be affected. Although most eyes showed increased reflectance infero-nasally on BLR imaging, there is a predilection for sparing of that area in MacTel.³⁰ Analysis of the location of the microcystoid spaces in relation to the theoretical MacTel area revealed most were contained within the area of parafoveal increased reflectance or within the surrounding band of decreased reflectance (corresponding to the outer halo of macular pigment accumulation on MPOD mapping). However in 2 eyes, a few microcystoid spaces extended more peripherally. This is the first demonstration of pathology in MacTel that appears beyond this confined oval area, which is thought to have a boundary of approximately $\leq 5-7^{\circ}$ horizontally and $\leq 5^{\circ}$ vertically on imaging and histologic analysis.^{6,24,31,32} This is in contrast to the findings of Spaide et al. where microcavitations were all reportedly located within the MacTel area.¹⁵ This difference may be due to the imaging modality used to estimate the MacTel area (MPOD versus BLR) or the use of different scan protocols (OCTA versus OCT).

There are several limitations to this study. This was a retrospective study and although we only included images with less than 30 µm interslice distance, scanning protocols varied between patients. It is possible that smaller microcystoid spaces may have been missed and this could explain the difference in frequency of lesions found in this series compared to the Spaide et al cohort. ¹⁵ The findings here are also at a single time point in the disease and microcystoid spaces may not necessarily be seen

at every scan date. Further confirmation of these findings in a larger cohort and analysis of variability with time would be interesting. It is also unclear at this stage why this microcystoid phenotype is seen more commonly in early disease and whether it represents a distinct phenotype. Examination of the relationship between microcystoid spaces and retinal function would be valuable. There are also very few normative datasets available to examine whether microcystoid spaces may be present in normal eyes. We attempted to address this by performing the same thresholding and segmentation in our control eyes. The control group had an even more robust scanning protocol with interslice distance of only 11 μ m and no microcystoid spaces were seen in any normal eyes or in the fellow eye of the patient with asymmetric disease.

In summary, we present quantification and analysis of microcystoid spaces in MacTel type 2 using SD-OCT imaging. Awareness of microcystoid spaces as part of the disease phenotype of MacTel type 2 is important when assessing differentials for microcystoid changes in macular diseases. In this study, microcystoid spaces were present often in early stage disease and were not associated with any retinal thickening as would be expected in traditional vascular causes of microcystic oedema.

We also demonstrate that although the majority of microcystoid spaces are contained within the MacTel area, small numbers can extend slightly beyond this region. Our findings add weight to the evidence suggesting Müller cell involvement in the pathogenesis of this disease. If confirmed, presence and volume of microcystoid spaces may also be a useful biomarker of disease activity. The use of a quantitative imaging method may have value as an endpoint in treatment trials or have prognostic

implications for MacTel and other disorders associated with presumed Muller cell degeneration.

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Figures

Figure 1.

Volume of the microcystoid spaces in MacTel type 2. Eighty-three percent of the assessed lesions had a volume between 16,000 and 44,000 μ m³.

Figure 2.

Location of microcystoid spaces within the retinal layers, showing a predilection for the INL, OPL and GCL. NFL: nerve fibre layer; GCL: ganglion cell layer; IPL: inner plexiform layer; INL: inner nuclear layer; OPL: outer plexiform layer; ONL: outer nuclear layer; HFL: Henle's fibre layer; ELM: external limiting membrane; MZ: myoid zone; EZ: ellipsoid zone; IZ: interdigitation zone; RPE: retinal pigment epithelium

Figure 3. Macular microcystoid spaces on spectral-domain optical coherence tomography imaging in two patients with MacTel 2. Left: overlay of microcystoid spaces (white dots) on the patient's blue light reflectance imaging. Right: Corresponding B-scan with microcystoid spaces in inner nuclear layer (arrows) and ganglion cell layer (arrow head).

Figure 4. Multimodal imaging of the right eye of a 37 year old man with MacTel type 2 (Patient 1 in Table). (A) Macula pigment optical density map showing characterstic central depletion as a darker area surrounded by a brighter ring of persisting macular pigment. (B) Late phase fluorescein angiography demonstrating parafoveal hyperfluorescence. (C) Spectral-domain optical coherence tomography scan through fovea with visible microcystoid spaces (arrow) in inner nuclear layer and outer plexiform layer. (D) Overlay of all segmented microcystoid spaces (white dots) against the area of increased parafoveal reflectance on blue light reflectance image. Note the two temporal microcytoid spaces that are present beyond the area of increased reflectance (arrows).

Table. Summary of the clinical and demographic features of identified patients.

ID: Patient identification number; CFT: Average central foveal thickness on ETDRS grid with Heidelberg Explorer; OCT: Optical coherence tomography; HC: Hyporeflective cavities; EZ: ellipsoid zone; ONL: outer nuclear layer.