Posterior microphthalmos and papillomacular fold-associated cystic changes misdiagnosed as cystoid macular edema following cataract extraction

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Purpose: Novel spectral-domain optical coherence tomography (SD-OCT) findings in posterior microphthalmos (PM) include cystoid changes in the papillomacular fold (PMF). These changes may be misdiagnosed as cystoid macular edema (CME) and may trigger unnecessary treatment including intravitreal injections. We report a case that underscores this scenario aiming to increase awareness of this entity among ophthalmologists.

Method: A case report.

Results: A 25-year-old male presented with a history of clear lens extraction in both eyes as a refractive procedure. Postoperatively, he was diagnosed with CME and received five intravitreal bevacizumab injections in each eye as well as oral diamox for persistent cystic spaces noted on OCT. He was referred to our institution with the diagnosis of non-responding CME. A complete ophthalmic evaluation, including ultrasonography and SD-OCT, confirmed the diagnosis of PM and PMF with cystic cavities in the PMF. A fluorescein angiogram disclosed absence of macular leakage or optic nerve head staining indicating that the cystic cavities seen on SD-OCT are not due to postoperative CME but are rather the intrinsic cystic changes commonly seen in eyes with PM and a PMF. These cystic cavities were incorrectly thought to represent postoperative CME, which triggered the unnecessary treatment and lack of response.

Conclusion: Cystic-like cavities are a component of the PMF in PM eyes and may be misdiagnosed as CME, especially in a postoperative setting. This case underscores the importance of understanding the intrinsic features of the PMF in eyes with PM and may help increase awareness among ophthalmologists of such a potentially confusing scenario.

Keywords: macular edema, papillomacular fold, posterior microphthalmos

Introduction

Posterior microphthalmos (PM) is an autosomal recessive ocular condition¹,² characterized by high hyperopia, a normal appearing anterior segment of normal or subnormal dimensions, and a short axial length with posterior segment foreshortening.²,³ A retinal papillomacular fold (PMF)²-⁵ or wrinkles⁴-⁶ is a characteristic finding in most eyes with this condition. In about half of the cases, the PMF harbors cystoid changes. The latter can be either visible on fundus examination or detectable only on optical coherence tomography (OCT) imaging.⁶ These cystic changes may be misdiagnosed as cystoid macular edema (CME) and may trigger unnecessary treatment, including intravitreal injections. We report a case that underscores this clinical finding aiming to increase awareness of this entity among ophthalmologists.
Case report
A 25-year-old otherwise healthy male presented with a history of poor vision in both eyes for many years for which he had been wearing thick hyperopic glasses. Neither his non-consanguineous parents nor his five siblings had poor vision or obvious hyperopia. One year prior to his presentation at our institution, he underwent clear lens extraction with posterior chamber intraocular lens implantation in both eyes as a refractive procedure. His vision did not improve following surgery, and he was diagnosed with postoperative CME, namely, Irvine-Gass syndrome, as the cause of his poor visual outcome. During his postoperative course, the CME appearance persisted for which he received five intravitreal injections of bevacizumab and two transseptal triamcinolone injections for each eye. He was also treated with oral diosux with no improvement and was referred to us for non-responding CME.

On examination, the best-corrected visual acuity was 20/125 in the right eye and 20/160 in the left eye. The intraocular pressure measured 12 mmHg and 16 mmHg in the right and left eyes, respectively. The anterior segment examination was unremarkable in both eyes except for well-centered posterior capsule opacification. The white-to-white horizontal corneal perimeter measurement was 11.5 mm in each eye. The posterior segment examination of both eyes showed mildly dilated and tortuous retinal blood vessels with a crowded optic disc and a horizontal PMF. PM was suspected, so an A-scan ultrasonography was performed, and it disclosed short globes with an axial length of 15.83 mm in the right eye and 15.87 mm in the left eye. Spectral-domain optical coherence tomography (SD-OCT) of the macula in both eyes showed a PMF (categorized as a retinal PMF or wrinkles, characteristic of PM,2-6) recognized during the fundus examination by opthalmologists familiar with this condition. In a previous report, we described five types of folds, based on fundus examination and photography. In addition, based on SD-OCT vertical cuts passing through the PMF, the folds were categorized into one of the following five configurations: 1) an omega-shaped fold defined as a high, inverted U-shaped fold containing one or more large, centrally located, inner nuclear layer cyst(s); 2) a high, inverted U-shaped fold; 3) an inverted V-shaped fold; 4) a low height, dome-shaped fold; or 5) only retinal surface wrinkles.6 We also described some features intrinsic to the PMF, including sparing of the majority of the outer retinal layers, the absence of a foveal depression in most cases with preservation of the inner retinal layers within the PMF and beyond it, and cystic cavities of variable sizes in approximately 50% of the cases.6 These cavities were consistently located in the inner nuclear layer, but they were not always clinically visible. In some cases, the PMFs also harbored cystic cavities located in the ganglion cell layer. The latter were invariably visible on opthalmoscopy.6 The case reported here harbored both a foveal elevation and visible cystic changes, clearly demonstrated in the ganglion layer.

Discussion
Because the anterior segment in PM appears to be of normal or subnormal dimensions, PM may not be easily recognized unless an axial length measurement of the globe demonstrates the axial foreshortening, or the retinal PMF or wrinkles, characteristic of PM.2-6 is recognized during
on SD-OCT, which, together, were mistaken for CME by
the treating ophthalmologist. The cystic cavities in this
case represent, nevertheless, the characteristic intrinsic feature
of the internal structure of the PMF in many eyes with PM.
These cysts can be differentiated from postoperative CME in
at least two ways. First, these are usually located in atypical
layers of the retina for postoperative CME, namely, in the
inner nuclear and ganglion cell layers, while the extracellular
component of postoperative CME typically starts in the outer
plexiform layer, and primarily affects the plexiform layers,
before it may progress to involve the inner retinal layers.7
Second, these cystic-like changes are not associated with
fluorescein angiographic leakage in the macular area or stain-
ing of the optic nerve head, features that characterize typical
postoperative CME. The absence of fluorescein leakage in
PMFs has been well described.3,6,8 In our case, the SD-OCT
features were typical of PMF-associated cystic changes and
the fluorescein angiogram was unremarkable with no macular
leakage or disc staining, supporting the diagnosis of PM with
PMF and excluding postoperative CME. In addition, the lack
of any response to intensive and aggressive treatment as well
as the stability of the cystic cavities on SD-OCT over almost
a year of follow-up also supports a structural finding rather
than postoperative CME.

In conclusion, this case report underscores the importance
of understanding the salient features that may be associated
with the PMF in eyes with PM in avoiding wrong diagnoses
and unnecessary interventions as well as the potential risks
that may be associated with them. Cystic changes seen on
SD-OCT in eyes with PM after having undergone ocular
surgery do not necessarily indicate postoperative CME.
A careful evaluation, including fluorescein angiography, is
critical in resolving the diagnosis.

Disclosure
In relation to this case report, the authors declare that there
are no conflicts of interest.

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as a genetically heterogeneous condition that can be allelic to nanophthal-
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