A 35-year-old woman presented to Retina clinic at Farabi Eye Hospital in September 2011, with complaint of blurred vision from several years ago. She did not have any history of trauma, retinal or choroidal vascular disease, or prior retinal or choroidal inflammation. Her best-corrected visual acuity was 20/25 in her right eye and 20/20 in her left eye with no significant refractive error.

Slit lamp examination of anterior segment was unremarkable. Dilated fundus examination of both eyes revealed RPE mottling changes with subretinal yellowish white plaque in macular area. Intraocular pressures (IOP) were measured 14 mmHg and 16 mmHg in the right and left eye, respectively, using Goldmann applantation tonometer.

SD-OCT of the right eye revealed a cup-shaped focal choroidal excavation in the macula with detachment between retinal pigment epithelium (RPE) and sensory retina layers. Disturbances of the outer retinal layers within the choroidal excavation area were obvious. SD-OCT also showed subretinal hyper reflective material. The retinal layers from the retinal nerve fiber layer to the outer plexiform layer were essentially intact.

SD-OCT of the left eye revealed disturbance of the RPE and photoreceptors layers and small subretinal fluid without obvious excavation.

Fundus autofluorescence images of the both macular region showed an outer circumferential ring of hyperfluorescence. Moreover in the left eye, central hypofluorescent areas were seen. Fluorescein angiography (FA) of the right eye revealed irregular hyperfluorescence in early phase corresponding to the RPE attenuation over the macular area that remained without change until late phases. In the left eye, there were hyperfluorescence areas in early phases. During mid to late phases, more hyperfluorescent areas in consistent with staining of materials were obvious. No leakage was observed in both eyes.

Indocyanine green angiography (ICG) of both eyes was near normal without any hyper-permeability in late stages, though mild dilation of submacular choroidal vessels was seen. Electroretinogram (ERG) was normal but Electrooculogram (EOG) was demonstrated a sever loss of the light response (low Arden ratio). The patient was followed up annually without any treatment, at each follow-up imaging was performed (OCT, FA). No change was observed in the course of the disease.
DISCUSSION
Best vitelliform macular dystrophy (VMD) also known as best disease is an autosomal dominant retinal dystrophy caused by heterozygous mutations in the bestrophin 1 gene [1]. Affected individuals show a solitary or multifocal vitelliform lesion (yolk-like) in the macula with good visual acuity in childhood [2]. Histopathologic evaluations, Spectral domain Optical coherence tomography (SD-OCT) and fundus autofluorescence (FAF) indicated lipofuscin accumulations at the level of the Retinal pigment epithelium (RPE) and also revealed atrophy and disruption of the RPE/photoceptor complex [1-3].

In 2006, FCE was first reported as a peculiar OCT finding in otherwise healthy eye by Jampop et al., as a choroidal posteriorly excavated zone without evidence of staphyloma or any scleral changes at that area [4]. Since that time, numerous reports have been raised about the choroidal excavation and association between focal choroidal excavation (FCE) and Choroidal neovascularization [5,6]. Epstein–Barr virus infection [7] Best disease [3] Polyposidal choroidal vasculopathy [8] and Central Serous Chorioretinopathy [9,10]. Margolis et al., demonstrated two type of choroidal excavation: excavations that involved the outer retina up to external limiting membrane without separation between RPE and photoreceptor (conforming type) and excavations that involved the RPE with separation between the outer retina and the RPE within it (nonconforming) [11]. The aetiology of FCE is still unknown, but it has been suggested that FCE is a congenital abnormality or may be multifactorial [12]. FCE have also divided in two patterns by Wakabayashi et al., one that involved the outer retinal layers up to the external limiting membrane (ELM), and other that involved only the retinal pigment epithelium (RPE) [13].

Several theories have been postulated about the aetiology of FCE, including [7,10,11]: failure of the choroidal development in the embryonic stage, microstaphyloma, focal choroidal atrophy due to congenital or acquired choroiditis, EBV infection and contraction of focal choroidal scarring. Nevertheless the aetiology of afore mentioned disease is currently unknown and further prospective study on FCE is necessary to clarify its aetiology, clinical course, and visual prognosis. Ellaban et al., reported that FCE was seen in 7.8% of eyes with central serous chorioretinopathy (CSR) [10]. Also, Lee et al., found that CSR was present in 24.4% of FCE eyes [14].

Suzuki et al., stated that an intervening hyporeflective space in the nonconforming FCE may represent serous retinal detachment (SRD) due to improper pumping action of RPE at the FCE [9]. They also stated that each of conforming or nonconforming FCE may be appeared after resolution of subretinal fluid.

Conforming FCE may progress to nonconforming FCE with separation of the photoreceptor tips from the apical surface of the RPE due to tension stress on outer retina, resulting in accumulation of fluid in subretinal space. These phenomena accompany with loss of vision due to shedding of photoreceptors in that space like a chronic CSR [11]. According to Margolis et al., classification, our patient had a nonconforming choroidal excavation. Given the our patient’s history, persistent metamorphopsia without significant change in vision over time, It seems that FCE in our case was nonconforming type from the long period of time and SRD in this case is not a secondary event. This may support this hypothesis that FCE size and shape would remain stable [14].

CONCLUSION
To the best of our knowledge, our patient is the second report of choroidal excavation in Best vitelliform macular dystrophy. Finally there may also be an association or a Causal relationship between best disease and FCE, which requires further studies to confirm.

REFERENCES

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