B-scan opthalmic Ultrasonography: a review corroborated with echograms

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Abstract
There are many investigative methods employed by Ophthalmologists to establish diseases of the eyes. An example is ophthalmoscopy. But this cannot explicitly evaluate posterior eye segment in cases of opaque vitreous or vitreous haemorrhages. The resort to B-scan ophthalmic ultrasonography seamlessly aids in non-invasive, cheap, real-time and reproducible diagnosis of ocular pathologies. Some of these ocular pathologies with supportive echograms observed in the course of our practice are reviewed in this study.

Teaching points
1. B-scan ocular ultrasonography is a cheap, readily available and reproducible means of evaluating lesions of the eyes using high frequency transducers.
2. B-scan ocular ultrasonography is a good substitute to ophthalmoscopy in evaluation of posterior segment of the eye in dense vitreous and vitreous haemorrhages.
3. B-scan ocular ultrasonography can readily detect ocular pathologies and this can further be improved with augmentation with dynamic ocular scan.
4. Definitive detection of the various types of ocular detachments is the beauty of B-scan ocular ultrasonography.
5. Contraindication for B-scan ocular ultrasonography is globe rupture.

Introduction
Ocular B-mode ultrasonography (USS) is an essential adjuvant for evaluation of ocular diseases. Its use in ophthalmic evaluation dates to 1950s [1,2]. This is not unconnected with the anatomic accessibility of the eye. The cystic nature of the eye also made it a good candidate for sonographic evaluation, a modality that is based on high frequency sound waves. But the beauty of ocular ultrasonography is its role as a substitute to ophthalmoscopy or as an ancillary to some ophthalmoscopic limitations. For example, it gives instantaneous information on lesions in eyes with opaque ocular media [1,3,4]. This opaque medium would ordinarily have conferred some hindrances to adequate ophthalmoscopic visualization of the posterior segment of the globe. Direct visualization of the retro-bulbar pathologic lesions with ophthalmoscope is also not possible even with clear media [1]. This therefore calls for a substitutive imaging modality like B-scan ocular ultrasonography that will readily characterize most ocular lesions. These lesions range from lens pathology to posterior eye segment pathologies.

Ocular B-scan produces a two-dimensional cross-sectional display of the globe and the orbit [1,3]. It is a cheap, non-invasive, real time and quick imaging modality with good spatial resolution. It could also be performed at the patient's bedside. Moreover, ocular B-scan can be done simultaneously with eye movement and this is referred as dynamic study. This dynamic study is incorporated in distinguishing the different types of ocular detachments as well as mobility of intra-vitreous echoes [1,3] One could also have used computed tomography (CT) and magnetic resonance imaging (MRI) in many ocular and orbital conditions. But they cannot scan in real time and they have limited role in the evaluation of the vitreous, retina and choroid detachments [1,3]. The above pitfalls justify our choice of B-scan ultrasonography in evaluations of ocular pathologies.

Aim
To highlight the usefulness of B-scan ultrasonography in elucidation of ocular pathologies.

Evaluations of ocular pathologies based on affected structures

Pathologies of the anterior eye segment: Moving in anterior posterior direction, the anterior eye segment consists of cornea, anterior chamber, iris, ciliary body, posterior chamber and lens. These are structures that rarely require USS, as they are readily assessed by clinical evaluations, ophthalmoscopy, slit-lamp examination, and ultrason sound bio microscopy. The later uses frequencies as much as 35-50 MHz to evaluate abnormalities like ocular trauma, foreign bodies and opaque elements in the anterior segment [2,3,5-8]. Notwithstanding

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Key words: ocular ultrasonography, b-scan, ophthalmoscopy, vitreous, retina, detachments

Received: April 15, 2019; Accepted: April 25, 2019; Published: April 29, 2019
the core utility of USS in posterior eye segment evaluations, it has also been used to localize intra-ocular foreign body (IOFB) in the anterior chamber thereby defining the size, shape and nature of the IOFB [1]. On USS, small metallic IOFB in the anterior eye segment will appear as a markedly echogenic short linearity that cast comet tail artefact.

**Lens pathology:** Pathologies of the lens of the eyes are known to cause visual impairment or blindness in millions of people. The major lens pathology is cataract which is a degenerative condition. Cataract is defined as any opacity of the lens that results in significant variations in the refractive index of the lens over distances like the wavelength of transmitted light. World Health Organization (WHO) documented 18 million people to be blind from cataract. Common causes of cataract include long-term exposure to the sun’s ultraviolet light, aging, inherited disorders, diabetes mellitus and trauma [9,10].

Cataract could be congenital, juvenile, pre-senile or senile. The latter is the commonest. Cataracts could also be classified as traumatic, immature, mature and hyper mature cataracts. On ophthalmoscopy, cataract is seen as a white reflection with an opaque lens called leukocoria. B-scan is important in cataracts as it evaluates the posterior eye segment which may be difficult with ophthalmoscopy. Such pre-operative B-scan is to rule out possible contraindications to cataract extractions such as retinal detachment or tumours. On B-scan, features of cataracts range from completely opaque hyperechoic lens cortex, intra-lenticular echoes, reflective material beneath the lens capsule, thick hyperechoic posterior capsule of the lens and combined thick hyperechoic anterior and posterior lens capsules. In immature cataracts, scattered opacities are separated by clear zones whereas in mature cataract, the thick echogenicity on the posterior capsule of the lens is a continuous plaque [3].

**Lens dislocation:** Lens dislocation could be traumatic, spontaneous or it could originate from hereditary or connective tissue diseases. Examples of the last aetiologies are Marfan’s syndrome, congenital aniridia, congenital glaucoma, Ehlers –Danlos syndrome, homocystinuria and molybdenum cofactor deficiency [11]. The commonest cause is trauma. Here, blunt force in anterior–posterior direction leads to globe equatorial expansion leading to zonular fibres disruption and lens dislocation [12-14]. Spontaneous lens dislocation can result from myopia. Partial zonular dialysis may lead to partial lens dislocation (subluxation) and complete zonular rupture leads to complete lens dislocation (ectopia lentis) [11]. In partial dislocation, the crystalline lens remains partially attached to one ciliary body and has an oblique axis. In partial lens dislocation, scan is done in different positions as the globulous lens might be in normal position when patient lies supine. In a complete dislocation, the lens sinks into the vitreous body in a dependent position, lying over the retina. It moves during dynamic examination, simulating floating lens sign seen in CT. The delayed complications of lens dislocation include secondary or phacoytic glaucoma, delayed retinal detachment, allergic uveitis, proliferative vitreoretinopathy, cataract and vision loss [11]. The secondary glaucoma resulting from lens dislocation is caused by a synchysis between the iris and the lens, causing the iris to curve forward, and termed iris bombe. This prevents the flow of aqueous humour from the posterior to the anterior chamber. Treatment of choice for lens dislocation is surgical repair, repositioning, explanting, or exchanging the displaced intraocular lens (Figures 1 and 2) [13].

**Pathologies of the posterior eye segment**

**Vitreous pathologies:** Vitreous pathologies are multifarous and they range from vitreous haemorrhages, vitreitis, asteroid hyalosis, persistent hyperplastic primary vitreous, coloboma, vitreous degeneration, posterior vitreous detachment, vitreoschisis, and retrohyaloid haemorrhage.

**Vitreous floaters, vitreous haemorrhages, vitreous degeneration, vitreitis and asteroid hyalosis**

The vitreous cavity is filled with transparent vitreous fluid or gel. The vitreous fluid is an acellular viscous fluid with 99 % water content and the rest consists of type 2 collagen fibrils and hyaluronic acid [3]. However, due to ageing, liquefaction of the vitreous called syneresis occurs. This will lead to ultrasonographic detection of few low-amplitude punctate and linear mobile echoes floating within the
Vitreous haemorrhages are caused by trauma, proliferative vascular retinopathies, coagulopathies, age related macular degenerations, retinal breaks, retinal microaneurysms, posterior vitreous detachment (PVD) and neoplasia. On USS, they appear as reflective intra-gel mobile dot echoes that are dependent and freely moves with dynamic scan. Acute bleed will appear as echo lucent or low reflective echoes. These haemorrhages may clear in 2-4 weeks or become chronic. When haemorrhages become chronic, the dot-like echoes get organized to form fibrinous vitreous membranes of varying reflectivity and devoid of anatomic attachment [3,8,16-18].

Vitreitis may mimic vitreous haemorrhages but the vitreous echoes will be less mobile and may have associated membranous echoes. But other supportive USS findings of ocular inflammation will be visible, and they include retinochoroidal thickening, exudative retinal detachment, optic disc elevation, Tenon's space widening. Asteroid hyalosis is an asymptomatic degenerative condition resulting from calcium soaps accumulation on vitreous fibrils in the eye [17]. It is usually unilateral, and the cause is unknown, but it is common in diabetes mellitus. On USS, the vitreous cavity shows a variable number of discrete mobile bright echoes (reminiscent of the particles in a snow globe) with a classical space between the particles and posterior globe wall (Figure 3) [3,8,16,17].

**Posterior vitreous detachment (PVD):** PVD is an-age related phenomenon whereby the posterior vitreous capsule or hyaloid detaches from the underlying retina [3]. It is also called hyaloids detachment and it can be an incidental finding or pathological. But it is seen in two-thirds of people older than 65 years [19]. PVD can be partial or complete. It can also be focal or broad, single or multiple and its reflectivity depends on aetiology [17]. On USS especially on dynamic scan, it characteristically shows as an undulating membrane that moves freely and should swil away from the region of the optic disc in cases of complete posterior vitreous detachment [3]. Note that the mobility of PVD is more than that of retinal detachment (Figure 4).

**Retro-hyaloid haemorrhage:*** It is also called sub-hyaloid haematoma or posterior hyphema. Vitreoretinal adhesions can cause retinal tears or avulsion of a peripheral blood vessel, resulting in vitreous and retro-hyaloid haemorrhage [3]. It can also be due to vitreoschis which is splitting of the posterior cortical vitreous, common in diabetic retinopathy [17]. On USS, retro-hyaloid haemorrhages appear as multiple low reflective dot echoes posterior to PVD which may demonstrate layering of echoes with a straight surface but moves with head movement [17].

**Persistent hyperplastic primary vitreous:** This is also called persistence of fetal vasculature. Persistent hyperplastic primary vitreous (PHPV) is a congenital developmental anomaly arising from failure of regression of embryological primary vitreous and hyaloid vasculature and presenting as leukocoria [3,20]. It is idiopathic, typically unilateral but 10% are bilateral. USS shows as microphthalia, retrolental high reflective thick fibrovascular linear membrane extending from the lens to the optic disc, thin lens with irregular, congenital cataractous posterior capsule and peri-papillary tractional retinal detachment [3, 4,17,21,22]. The pattern of retinal detachment in PHPV is usually funnel shaped suggestive of total retinal detachment. A persistent hyaloid artery may also be present in the centre of retrolental band detected with Doppler scan.

**Pathologies of retina**

Retinal pathologies include retinal detachment, retinal tear, sub-retinal haemorrhages, retinal cysts.

**Retinal detachment**

Retinal detachment (RD) is the detachment of the inner sensory layer of the retina from the outer pigmented layer called retinal pigmented epithelium [3,23]. Basically, RD is categorized as rhegmatogenous RD, tractional RD, and exudative RD. RD could also be total, partial, or focal, depending on its extension. The classical USS features of total RD is a “V” shape echogenic lesion in the vitreous cavity with anterior attachment to the ora serrata and posterior...
attachment to the optic nerve head [3,17]. On the other hand, partial RD will show as an intra-vitreal linear echogenic membrane extending to the optic nerve head, but not across it [3,23]. The leaflets of the membranes in acute RD are thinner and more mobile than in chronic RD. Thus, Chronic RD is often seen as a rigid “triangle sign” [3,15,18,24]. Sometimes, retinal cyst can form in chronic RD. The retina can have an open or tightly closed funnel configuration in cases of complete RD seen in conditions like proliferative vitreoretinopathy, retinopathy of prematurity, PHPV and retinoblastoma. Exudative and tractional RD are often associated with subretinal haemorrhages (Figure 5).

Sub-retinal haemorrhages

Causes of sub-retinal haemorrhages are trauma, age related macular degeneration, intra-papillary haemorrhages, diabetes mellitus, retinal tear, RD and ocular tumours [17,25,26]. On USS, sub-retinal haemorrhages present as echogenic debris beneath an elevated retina. This debris has variable reflectivity due to clotted or lysed blood. The debris might be mobile on dynamic scan or having a shifting contour. It tends to change contour and reduction in height with time due to resolution or breakthrough vitreous haemorrhage [17].

Retinoblastoma

Retinoblastoma is the commonest childhood malignant ocular tumour. It is seen in 1 in 17,000 live births with mean age at presentation of 2 years for unilateral forms and one year in bilateral forms [27]. The tumour arises from the retina and could be exophytic (subretinal) or endophytic (growing towards the vitreous cavity) [17]. On USS, it could present as a single or multiple solid mass with irregular high internal reflectivity due to presence of calcium [17,28]. Internal calcification is very characteristic as it seen in up to 90% of cases (Figure 6).

Choroidal pathologies

Choroidal detachment

Choroidal detachment is caused by trauma, hypotony, corneal ulcers, intra-ocular pressure lowering agents, ocular surgeries and RD. Choroidal detachment can also cause RD [29]. In choroidal detachment fluid accumulates in the supra-choroidal space and this fluid can be echo lucent or echo rich. On USS, choroidal detachment presents as a dome shaped thick membrane in the periphery and it does not extend beyond the equator as its limited by the vortex vein exit inferiorly [17].

Other ocular morphologies

Alterations in size and shape of the ocular globe

The normal antero-posterior diameter of the ocular globe is approximately 22 to 25 mm in adolescents and adults. In patients with long-term myopia, the antero-posterior axis of the globe is lengthened. The globe sometimes develops a thin wall, often manifesting as a pear-shaped sacculation of the posterior pole. This is referred to as a posterior staphyloma [3].

Phthisis bulbi

Phthisis bulbi is the terminal stage of many ocular lesions and this include trauma, infection and surgery. Here, the eye is redundant, microphthalamic, blind, non-functioning, with extensive calcifications. The affected eye has crenated, shrunken globe with extensive calcification and loss of the normal shape. Cataracts, partial lens dislocation, vitreous fibro-vascular membrane and retinal detachment may also be seen [3,4,30].

Conclusion

Ocular B-scan is a cheap, convenient and readily available tool for diagnosis of diseases of the eyes. This is most valuable in diseases of posterior eye segment especially in opaque vitreous or vitreous haemorrhage.

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