Prevalence, Pattern, and Risk Factors for Retinal Vascular Occlusions in a Tertiary Hospital in Jos, Nigeria
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Objective: Retinal vascular occlusions are the second most common retinal vascular diseases following diabetic retinopathy. They are associated with several systemic and ocular pathologies and are significant causes of visual loss. The aim of this study was to determine the prevalence, pattern, and risk factors for retinal vascular occlusions in a tertiary hospital in Jos, Nigeria.

Methods: A retrospective review of all cases of retinal vascular occlusions seen in the eye clinic from January 2011 to December 2014.

Results: Three thousand eight hundred twenty-one new patients were seen during the study period, and there were 52 (1.36%) cases of retinal vascular occlusions. Only the case files of 44 (84.6%) patients were available and were analyzed. These were made up of 17 (39%) males and 27 (61%) females with a mean age of 53.5 years. These had a total of 46 retinal vascular occlusions comprised of 37 (80.4%) retinal vein occlusions, 8 (17.4%) central retinal artery occlusions, and a case (2.2%) of combined central retinal artery and vein occlusion (Figure 1).

Conclusion: Retinal vascular occlusions are uncommon in Jos. They are associated with significant systemic pathologies and lead to severe visual loss. A comprehensive management of patients with retinal vascular occlusions is, therefore, necessary to correct associated diseases or predisposing risk factors that could lead to local recurrences. Early diagnosis and treatment of these associated systemic disorders in the general population will also help in reducing the incidence of these conditions and resultant blindness.

REFERENCES

Table 1: Systemic and ocular associations

<table>
<thead>
<tr>
<th>Type of VO</th>
<th>HT (n)</th>
<th>DM (n)</th>
<th>HIV (n)</th>
<th>SCD (n)</th>
<th>Glaucoma (n)</th>
<th>Others (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRVO</td>
<td>17 (53)</td>
<td>4 (36.4)</td>
<td>-</td>
<td>-</td>
<td>3 (33.3)</td>
<td>-</td>
</tr>
<tr>
<td>BRVO</td>
<td>4 (13)</td>
<td>2 (18.2)</td>
<td>-</td>
<td>-</td>
<td>3 (33.3)</td>
<td>-</td>
</tr>
<tr>
<td>HRVO</td>
<td>8 (25)</td>
<td>4 (36.4)</td>
<td>-</td>
<td>-</td>
<td>2 (22.2)</td>
<td>-</td>
</tr>
<tr>
<td>CRAO</td>
<td>3 (9)</td>
<td>1 (9)</td>
<td>2 (100)</td>
<td>1 (100)</td>
<td>1 (11.1)</td>
<td>6 (100)</td>
</tr>
<tr>
<td>CRAO+CRVO</td>
<td>1 (100)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>32 (100)</td>
<td>11 (100)</td>
<td>2 (100)</td>
<td>1 (100)</td>
<td>9 (100)</td>
<td>6 (100)</td>
</tr>
</tbody>
</table>


Figure 1: Sex distribution of retinal vascular occlusions. HRVO: Hemiretinal vein occlusion; CRVO: Central retinal vein occlusion; BRVO: Branch retinal vein occlusion; CRAO: Central retinal artery occlusion

Figure 2: Sex distribution of retinal vascular occlusions. HRVO: Hemiretinal vein occlusion; CRVO: Central retinal vein occlusion; BRVO: Branch retinal vein occlusion; CRAO: Central retinal artery occlusion

Abstracts

VITREO-RETINA

Mobile Phones for Retinopathy of Prematurity Screening in Lagos, Nigeria, Sub-Saharan Africa
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Introduction: Retinopathy of prematurity (ROP) was initially thought to be rare in Sub-Saharan Africa. The first reported cases in Nigeria were noted in blind schools study with an incidence of 0.5%. Another study in Ibadan, Southwest Nigeria noted a prevalence of 5.5%. A more recent study in Port Harcourt, Southern Nigeria, and the middle belt region recorded prevalence rates of 47.2% and 89.6%, respectively. Reasons given for the rarity of ROP included the nonsurvival of the preterms due to inadequate infrastructures. The other reason given was the nonavailability of trained personnel to detect the disease. Some eye doctors would not screen because of the nonavailability of treatment modalities. However, recent developments have shown that more preterms are delivered, and facilities for special baby care are becoming readily available. A sample of opinions among Neonatologists in Nigeria revealed that screening for ROP is not done in their centers. The aim of this study was to describe the use of mobile phone technology in the screening of ROP in Lagos, Nigeria, Sub-Saharan Africa.

Methods: Ethical Committee approved the program before the commencement of the study. Preterms with birth weight of <1.5 kg or gestational age of <32 weeks are eligible for screening according to laid down criteria. In conjunction with the neonatologist, topical tropicamide (0.5%) and phenylephrine (2.5%) were used to dilate the pupils. Examination was carried out in dim illumination. A lid speculum (Sauer Premature Infant Speculum Denville, New Jersey, USA) was used to part the lids. Indirect ophthalmoscopy was used to examine the fundus to ensure there were no missed diagnoses. Iphone 5S by Apple Inc. Cupertino, California, United States with 20 diopter lens (Volk) was used to examine the fundus. A mobile phone app, FILMIC Pro (developed by Cinegenix, Seattle, USA) was launched in video mode. The camera flash served as the source of illumination, and its intensity was controlled by the App. The 20 diopter lens was used to capture the image of the retina which was picked up by the camera system of the mobile phone [Figure 1]. After a thorough examination, the video was examined and snapshot of still images taken [Figure 2]. Another mobile phone app, Aviary (Adobe Inc., New York, USA) was used to edit the picture.

Another mobile phone app, Aviary (Adobe Inc., New York, USA) was used to edit the picture.
Introduction: The number of people with type 2 diabetes is increasing in every country. Eighty percent of people with diabetes live in low- and middle-income countries.\textsuperscript{1-3} Prevalence rates of diabetes mellitus in Nigeria are estimated to be between 8% and 10%.\textsuperscript{4} The International Diabetes Federation reported that Nigeria has the highest number of diabetics in Africa.\textsuperscript{5} Diabetic retinopathy (DR) is now a significant cause of blindness in developing countries such as Nigeria.\textsuperscript{6,7} A majority of diabetics are unaware of their diabetic status, those who are aware do not realize the importance of consistent follow ups with the physicians, and most of the physicians are not trained to detect and refer sight-threatening DR for timely treatment.\textsuperscript{8} People with DR can be treated to prevent visual impairment and blindness. Most endocrinologists and general physicians are not trained to screen for DR. Indirect ophthalmoscopy is too cumbersome for general ophthalmologists, direct ophthalmoscopy is inadequate, and fundus photography is beyond the reach of poor economies. Mobile phones technology combined with a 20 diopter lens appear to be less expensive and can be taught to endocrinologists, general physicians, and resident doctors. The aim of this study is to describe the use of mobile phone technology in the screening for DR in poor resource setting of Nigeria.

Methods: Patient population: The diabetic patients attending the University College Hospital and the Lagos University Teaching Hospital. All consenting patients had their pupils dilated with tropicamide 0.5% and phenylephrine 10% before the examination. The iPhone system is

Results and Discussion: The flash of the phone camera serves as the illumination, and its intensity is controlled by the mobile phone app - FILMIC pro. The distance between the phone and the 20 diopter lens is much shorter than what obtains in indirect ophthalmoscopy [Figure 1]. A video recording of the examination is preferable. The video can subsequently be reviewed to choose the appropriate frame(s) for editing.

Mobile phones technology appears to be useful in screening for ROP. It is good for documentation, patient education, and for teaching in poor resource settings where Retcam is not available. It also enhances telemedical services in remote areas where retina surgeon may not be readily available. Previous workers have stressed the use of mobile phone fundus photography in poor resource settings.\textsuperscript{4-6}

Conclusion and Recommendation: It is recommended that more studies be done in the area to perfect the method.

REFERENCES

The Role of Electrophysiological Tests in the Management of Retinal Diseases
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Introduction: Electrophysiological tests (EPTs) are a group of tests that may be used to ascertain the integrity of the visual system and its pathways beyond standard clinical examination. They were first developed in the mid-19th century and introduced in clinics in the mid-20th century.[1] Objectives are to assess function of visual pathway from the photoreceptors to visual cortex.

Objectives of this Presentation: (1) To demonstrate the importance of EPT in the management of retinal diseases. (2) To enumerate some important EPT available.

Discussion: The various types of electrophysiological testing available include[1-7] (1) electroretinogram (ERG), (2) macular or focal ERG, (3) pattern ERG, (4) electroocoilogram (EOG) measurement of retinal function with standardized eye movements, (5) flash visually evoked cortical potential (Flash VEP), (6) pattern appearance VEP, (7) other nonstandard procedures for recording VEP and ERG, e.g., measurement of interocular beat frequencies, flicker, and sweep VEPs, multi-focal ERG, fast oscillation, (8) electromyogram, and (9) electrystagmography. For measurement of nystagmus and eye movements, among these, the ERG, EOG, and VEP are the most commonly performed tests in clinical practice. Indications for these tests include but not only limited to[1-7] (1) confirmation of neurological or ophthalmological disease, (2) evaluation of the visual pathways in the presence of media opacities, (3) pediatric neurology assessment, (4) unexplained visual loss, (5) drug toxicity and monitoring, (6) assessment of retinal or optic nerve function following trauma, (7) quantitative assessment of visual disease, (8) detection of disease or carrier states of inherited visual disorders, and (9) research. The full field ERG tests the mass electrical response of the retina to photic stimulation, thereby assessing the status of the retina in eye disease. The two components that are most often measured are the a- and b-waves.[1] The a-wave is the first large negative component, followed by the b-wave which is corneal positive and usually larger in amplitude. Changes in these waves indicate different types of retinal disease [Figure 1]. This type of ERG records the mass potential over the entire retina thus unless 20% or greater of the retina is diseased the test will appear normal. To this end, the multifocal ERG was developed to overcome this challenge and is capable of detecting maculopathies and other focal retinal diseases.[1] The pattern ERG is used to detect subtle optic neuropathies. The EOG measures the potential that exists between the cornea and Bruch’s membrane. It is useful in chorioquine and hydroxychloroquine maculopathy and Best’s disease [Figure 2].[1] VEP is an evoked potential caused by a visual response to an electrical or auditory stimulus.

References

Optic disc melanocytoma (ODM) is a rare, benign, pigmented ocular tumor arising from dendritic melanocytes within the optic disc or any part of the uvea.\(^1,2\) A 17-year-old female with painless, nonprogressive poor vision noticed since childhood. At examination, unaided visual acuity was 6/36 and 6/18 in the right eye and left eye, respectively which improved to 6/9 bilaterally with pin hole examination. There was relative afferent pupillary defect in the right eye. Fundoscopy of the right eye revealed a raised pigmented optic disc lesion, about 1.5 disc diameters in size, obscuring the lower disc margins with overriding normal vessels [Figure 1]. The anterior and posterior segments of the left eye were normal. Ocular ultrasound done showed a well-defined homogenous soft tissue mass arising from the choroid in the optic nerve area and projecting into the vitreous cavity. The clinical impression was presumed right ODM with refractive error. The patient was counseled and spectacles prescribed, periodic follow-up and monitoring with fundus pictures were emphasized.

**Discussion:** ODM is seen as a dark brown/black lesion on fundoscopy, it may be flat or slightly elevated with fibrillate extensions which may proceed to the edge of the disc and into the adjacent retina.\(^3\) It is usually unilateral, bilateral cases have been reported, but they are quite rare.\(^4\) Visual acuity in most cases remains normal. In a study by Zografos et al., visual acuity was normal in 70%, subnormal in 27% of the cases.\(^4\) Afferent pupillary defect is an associated finding in 30% of cases.\(^4\) It is likely due to compression of the optic disc fibers and may be seen in the presence of good vision.\(^2\) ODM is commonly described as a stationary lesion.\(^2,3\) It may however enlarge in up to 10–15% of cases,\(^3\) malignant transformation is rare, reported in 1–2% of cases.\(^3\) Useful investigations include ultrasound or computer tomography especially for lesions >0.5 mm,\(^5\) but they cannot differentiate ODM from other raised lesions involving the optic disc. Fundus fluorescein angiography is the hallmark of diagnosis; it shows hypofluorescence due to closely compact and deep pigmentation of the melanocytes.\(^1,2\) Histology shows round, closely apposed cells with normochromic nuclei and distinct borders with abundant cytoplasm.\(^1\) Factors supporting ODM in this patient include the presence of an RAPD, raised pigmented optic disc lesion, nonproliferation of the lesion on follow-up visits, and being a female of African descent. Differential diagnosis includes juxtapapillary choroidal melanoma, juxtapapillary choroidal nevus, metastatic melanoma to the optic disc, adenoma of the RPE, and hyperplasia of the RPE.

**References**

Conclusion: ODM is a rare benign tumor; it should be considered in patients presenting with a pigmented optic disc lesion and regular follow-up with fundus photography is advocated.

REFERENCES