Evaluation of the optic nerve and scleral-choroidal-retinal layer with ultrasound elastography in glaucoma and physiological optic nerve head cupping

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Abstract

Aim: To evaluate the strain ratio of the optic nerve and retina-choroid-sclera (RCS) layers in individuals with physiological optic disc cupping (PC) and glaucoma patients using strain elastography. Material and methods: We evaluated 56 eyes of 56 subjects (20 eyes with glaucoma, 19 eyes with PC, and 17 normal eyes). The strain ratio of orbital fat to optic nerve (SROFON) was calculated as the ratio of the optic nerve to intraconal fat tissue and the strain ratio of orbital fat to retina-choroid-sclera (SROFRCS) was calculated as the ratio of RCS layers to intraconal fat tissue. Results: SROFON was 0.92 in the control group, 1.07 in the PC group and 1.6 in the glaucoma group and a statistically significant difference was present between the three groups (p<0.05). SROFRCS had no statistically significant difference between the three groups. Conclusions: SROFON values could contribute to the differentiation of the patients with glaucoma and PC.

Keywords: glaucoma; ultrasonography; strain elastography; strain ratio; optic nerve.

Introduction

Glaucoma is a chronic, progressive, and asymptomatic anterior neuropathy. In time it can produce damage of the retinal nerve fiber layer (RNFL) and optic nerve, with loss of the related visual field [1,2]. A cup/disc (C/D) ratio (ratio of the cup located in the middle of the optic disc to the optic disc itself) higher than 0.4 raises the suspicion of glaucoma. However, in the healthy population this ratio can range between 0.0-0.9 [2]. Physiological cupping (PC) is diagnosed in cases with high C/D ratio, normal neuroretinal rim shape, normal RNFL thickness, and normal visual field test [2]. It is difficult to know whether the increase of C/D ratio is physiological or due to glaucomatous damage using only biomicroscopic optic disc examination and in practice, further tests are required. Early stages of the C/D ratio changes can also be missed on the biomicroscopic optic disc examination during the follow-up of glaucoma progression [3-5].

Although grayscale ultrasonography (US) is a non-invasive method that has been used to evaluate the eye structures for many years, elastography is a new technique and is not used as widely for the orbita and orbital structures as for other organs [6,7].

Strain elastography (SE, quasi-static elastography, qualitative elastography) requires the application of force with the probe (compression-decompression) to ensure the tissue displacement. SE shows a color-coded map reflecting the compressibility of the tissues. The technique compares the tissue of interest with the surrounding tissues and enables a relative evaluation of stiffness. The tissue for which the stiffness will be evaluated and a reference tissue for comparison are selected and the stiffness calculated accordingly. SE enables both qualitative (color-coded map) and semi-quantitative (strain ratio cal-
culation) evaluation of elasticity [6,8-10]. The basic limitation of this technique is the wide variability of the image and elasticity values as the pressure applied with the probe is examiner-dependent and not standard [11]. The sensitivity of SE vary between 57% and 92% [12,13].

The first study showing that ocular structures could be evaluated with elastography was conducted in 2010 [14]. The main problems in the SE evaluation of ocular structures are the possible influence of the small dimension and ocular fluid content. The semiquantitative nature of SE also creates difficulties [12,15]. Various studies have evaluated the ON using elastographic techniques in glaucoma, optic neuritis, and multiple sclerosis [12,15-18].

In this study our aim was to investigate the ability of SE and of the SR values in differentiating between the optic nerve and scleral-choroidal-retinal (SCR) layers in patients with PC and glaucomatous optic nerve damage.

Material and methods

Patients

This prospective study was conducted between April 2015 and May 2016. Local Institutional Ethics Committee approved the study. A written informed consent form was obtained from all patients before the study. We assumed the two eyes of the same subject would be similar. We therefore only evaluated the right eye in each person. We evaluated 56 eyes of 56 subjects (20 eyes with glaucoma, 19 eyes with PC, and 17 eyes with completely normal findings). Subjects with systemic diseases such as hypertension, diabetes mellitus, or peripheral and central nervous system disorders that caused neuronal damage (multiple sclerosis, cranial vasculitis, etc.) were excluded. All patients underwent a routine eye examination as well as Optic Coherence Tomography (OCT) and visual field tests at the eye clinic before elastography. The physician who performed the SE evaluation was blinded to the condition of the individuals prior to the SE investigation.

US technique

The investigation was conducted using an elastography device (Esaote, MyLab60, Geneva, Italy) with 6-18 MHz linear probe while the patient was in the supine position with the eyelids closed. The investigation was started using grayscale US (average frequency of the transducer 15 MHz) and was continued with SE during minimal eye compression. The most appropriate image was selected by performing compression-decompression 5-6 times until the spring turned green indicating the image was suitable for SE evaluation. The values of the strain ratio (SR) of orbital fat to optic nerve (SROFON) and the strain ratio (SR) of orbital fat to retina-choroidal-retinal (SCR) layers in patients with PC and glaucomatous optic nerve damage.

Fig 1. Strain elastography of the eye and the calculation of strain ratio of a) the optic nerve and b) the retina-choroid-scleral layers

Fig 2. Strain elastography and Optic Coherence Tomography of the right eye in a) 41-year-old normal female patient (SROFON 0.76, OCT C/D ratio 0.37, and RNFL thickness 109 μm); b) 68-year-old female with physiological optic nerve cupping (SROFON 1.1, OCT C/D ratio 0.80, and RNFL thickness 112 μm); c) 53-year-old female patient with glaucoma (SROFON 1.36, OCT C/D ratio 0.88, and RNFL thickness 72 μm).

Fig 3. Point distribution graph of SROFON values of the 3 groups (Group 1: Control group, Group 2: PC group, Group 3: Glaucoma group).
higher than the control group (p<0.001). b No significant difference between the 3 groups. c Significantly lower than the control group (p<0.001). d Significantly higher than the control group (p<0.001).

The mean age was 61.12±7.1 years in the control group, 60.11±8.6 years in the PC group, and 59.7±12.1 years in the glaucoma group and the female/male ratio was 11/6, 11/8, and 14/6, respectively (p>0.05).

The mean C/D ratio was statistically significantly higher in the PC and glaucoma groups than the control group (p<0.001). However, there was no difference between the PC and glaucoma groups (p>0.05). There was no statistically significant difference between the control and PC groups for mean RNFL thickness on OCT (p>0.05) but this value was statistically significantly lower in the glaucoma group comparing with the PC and control groups (p<0.001). Mean SROFRCS values were highest in the glaucoma group and lowest in the control group. There was a statistically significant difference between the 3 groups (p<0.001). There was no statistically significant difference between the 3 groups for mean ROFRCS values (p>0.05) (Table 1).

In figure 2 we showed a SE and OCT example from each of the 3 groups. The difference between the 3 groups can be seen more clearly in the point distribution graph of the SROFON values of the control group, PC group and glaucoma group (fig 3).

**Discussions**

Elastography, a new US technique, has been largely used for evaluating the prostate, lymph nodes, breast, thyroid, testicles, or kidneys [16]. However, we found only 3 studies in which elastography was used for the study of the optic nerve in glaucoma patients but none of these studies included subjects with PC [12,15,18]. Agliardi et al compared the Shear Wave Elastography[SWE] and optic disk/optic nerve SR values obtained in glaucoma patients with a control group but found no statistically significant difference [18]. Dikici et al detected higher elasticity values in the glaucoma group [12] and Unal et al found higher optic nerve elasticity values in the glaucoma group than the control group [15]. We obtained similar results regarding the elasticity of the optic nerve in glaucomatous patients.

In a recent published study Inal et al [16] found high optic nerve stiffness using both elastographic techniques, SWE and SE, in multiple sclerosis patients comparing with optic neuritis patients. The optic nerve stiffness in patients with unilateral optic neuritis was higher compared with the normal eye in the study of Batur et al using the Acoustic Radiation Force Impulse (ARFI) technique [17].

Glucoma causes irreversible retinal ganglion cell damage. This results in fibrosis, especially in the retinal nerve fiber layer and therefore in the optic nerve [19]. Senility and some nervous system disorders can also compromise the optic nerve [20]. We excluded from the study groups the patients with nervous system disorder and there was no significant difference in the mean age of the groups. So, our opinion is that the stiffness values are directly related to glaucoma.

Burgoyne et al [21] conducted an experimental study in monkeys. They found that glaucoma caused connective tissue damage in the optic nerve even in the early stages. This was followed by recovery with scar formation in the extracellular matrix. They stated that such scar formation could lead to stiffness of the optic nerve head. The high ON stiffness in the glaucoma group in our study can be explained by fibrosis.

The highest SROFON values were in the glaucoma group and the lowest in the control group. Our results indicate that elastography provides meaningful values for differentiating PC and glaucoma changes of the optic nerves.

US in one of the most reliable radiologic methods and has no known significant side effects. Orbital Doppler
US and grayscale US are accepted as safe, easy-to-use and non-ionizing radiation methods. Ocular elastography may cause side effects in the eye and surrounding tissues. It is however safe and quick in evaluating retrobulbar fat tissue and the optic nerve [16].

Our study had certain limitations. The first of these is that SE results can vary by the user. Another limitation is that an eye-specific orbital elastography probe does not exist. We therefore used the most appropriate probe and frequency range for the eye. The third limitation is that the eye has viscoelastic and poroelastic complex compartments. The compression effect may therefore not reach the deep orbital and periorbital structures.

In conclusion, we found that SROFON values differed between PC and glaucoma patients. Elastography may contribute to determining whether changes in subjects with a high C/D value are physiological or glaucoma-related. However, these findings need to be supported by other studies using various elastography techniques on a larger patient series.

Conflict of interest: none

References