Cognitive impairment and psychiatric symptoms in normal tension glaucoma (NTG) - A naturalistic study

Mohammad Z Mustafa*, Tej Rane-Malcolm¹, Andrew J Tatham³, Bal Dhillon¹, Amanda Cotton², Premal Shah³ and Roshini Sanders³

¹Princess Alexandra Eye Pavilion, Chalmers Street, Edinburgh, United Kingdom
²Royal Edinburgh, Department of Psychiatry, Edinburgh, United Kingdom
³Queen Margaret Hospital, Department of Ophthalmology, Dunfermline, United Kingdom

Abstract

Purpose: To examine the cognitive and neuropsychiatric characteristics of patients with Normal Tension Glaucoma (NTG).

Methods: A cross-sectional observational study of 99 patients with glaucoma and untreated intraocular pressure consistently <22 mmHg on phasing. All subjects underwent cognitive assessment and psychiatric symptom screening using the mini mental state examination (MMSE), adult verbal learning test (AVLT) and general health questionnaire (GHQ). All subjects had comprehensive ophthalmic examination and documentation of past medical and psychiatric history.

Results: Subjects had a mean age of 70.5 years and 59 (59%) were female. 57 (57%) had an advanced visual field defect in the worse eye, 19 (19%) a moderate defect and 23 (23%) a mild defect. There were a high proportion of patients with impaired MMSE, AVLT or GHQ scores. 28 (28%) had significant short-term memory impairment on AVLT, and 28 (28%) scored 5 or more on the GHQ suggesting psychiatric caseness. 12 (12%) scored 24 or less on the MMSE. 43 subjects underwent CT or MRI neuroimaging, with 13 (30%) showing age excessive generalised cerebral atrophy and 9 (20%) showing excessive small vessel disease. 41 patients had systemic hypertension, 28 ischaemic heart disease, 21 diabetes, 18 migraine, 13 transient ischaemic attack and 3 Raynaud’s disease.

Conclusion: Patients with NTG had a high prevalence of cognitive impairment and psychiatric symptoms identified using common screening tools, suggesting a possible shared intraocular pressure independent mechanism of disease. Ophthalmologists should be aware of the possibility of undiagnosed cognitive impairment and psychiatric morbidity in some patients with NTG.

Introduction

Normal tension glaucoma (NTG) is defined as a form of primary open angle glaucoma (POAG) where intraocular pressure (IOP) is consistently less than 22 mmHg. While this definition has limitations, due to population differences in what constitutes a “normal” pressure and diurnal and day-to-day fluctuation in IOP, it is likely that the relative importance of IOP and non-IOP related mechanisms of glaucomatous damage vary between those that develop glaucoma at low and high IOP. Worldwide the prevalence of NTG varies. In people of European ancestry, NTG is thought to affect as many as 50% of patients suffering with POAG, while in the Japanese population NTG accounts for up to 92% of glaucoma [1,2].

The diagnosis of NTG can be challenging and, perhaps due to the persistent perception that raised IOP and glaucoma are synonymous, patients may present late. Previous studies have shown a disproportionate number of patients with NTG are registered blind. For example, we previously examined a 20-year period of blind registrations in Fife, a region of Scotland with a population of 400000, and found those with NTG were over represented in blind registrations [3,4]. The management of NTG is also challenging. Although studies such as the Collaborative Normal Tension Glaucoma Study have shown that medical or surgical reduction of IOP reduces the risk of disease progression [5], some patients’ progress despite IOP reduction suggesting susceptibility to non-IOP mediated retinal ganglion cell damage [6]. Glaucoma is a neurodegenerative disease characterised by changes to the optic nerve head, however there is growing evidence that the entire visual pathway is affected [7]. This raises the possibility that patients with glaucoma may exhibit non-ocular signs of neurodegeneration. Several studies investigating this theory have suggested an association between glaucoma and neurodegenerative diseases such as dementia [8,9]. For example, the recent Three-City-Alenior Cohort study found a high prevalence of glaucoma in patients with dementia, especially NTG [10]. An association between glaucoma and Alzheimer’s disease (AD) has also been described, and it has been proposed that both conditions involve neuro-inflammation, decreased cerebrospinal fluid pressure and biomechanical stress and that B-amyloid may have a toxic effect on RGC apoptosis [11-13]. There is also evidence that vascular dysfunction contributes to glaucomatous optic neuropathy, with systemic vascular diseases including stroke and ischaemic vascular disease, as well as vasospastic conditions such as Raynaud’s disease and migraine, associated with a higher risk of NTG [14-17].

The aim of the current study was to examine the cognitive and psychiatric morbidity in some patients with NTG.

Correspondence to: Mohammad Z Mustafa, Princess Alexandra Eye Pavilion, Chalmers Street, Edinburgh, EH3 9HA, United Kingdom, Tel: 0131 5361000; E-mail: mohammadmustafa@nhs.net

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neuropsychiatric characteristics of a naturalistic clinical series of patients with NTG using previously validated psychiatric tools. Identifying disease associations may provide further insight into disease mechanisms. Cognitive impairment also has implications for treatment concordance so it is possible that identification of cognitive or psychiatric symptoms could potentially help improve outcomes.

Methods
This was a cross-sectional observational study including 132 consecutive patients with a diagnosis of NTG attending the ophthalmology department at NHS Fife, Scotland in the year 2009. NTG was defined by the presence of glaucomatous optic neuropathy on slit lamp biomicroscopy identified by a glaucoma specialist (R.S.) and an untreated IOP consistently <22 mHg using Goldmann applanation tonometry. Ninety-nine of these patients consented to undergo cognitive assessment and psychiatric symptom screening. Of the remaining 33 patients, 8 lacked capacity to consent due to advanced cognitive impairment requiring long-term care. Fifteen were serial non-compliers with appointments and medications and had high fixation errors at visual field testing. Six declined to take part in the study.

Subjects provided written informed consent before participation and the study conformed to the tenets of the Declaration of Helsinki. The Fife ethics committee granted ethical approval prospectively.

All patients with a possible diagnosis of glaucoma referred from community optometrists in Fife, Scotland were streamlined to the NHS specialist glaucoma service comprising higher surgical trainees, specialist glaucoma nurses and hospital optometrists under the supervision of a consultant glaucoma specialist (R.S.). At the initial visit all patients had a full ophthalmic examination, including visual acuity, refraction, detailed stereoscopic biomicroscopy of the anterior segment, gonioscopy, IOP measurement with Goldmann applanation tonometry, dilated fundus examination, stereoscopic photography of the optic disc and standard automated perimetry (SAP) using the 24-2 Swedish Interactive Threshold Algorithm standard strategy of the Humphrey Field Analyser II (Carl Zeiss Meditec, Dublin, CA, USA). A comprehensive medical history was also taken including history of smoking, migraine, Raynaud’s disease, hypertension, ischaemic heart disease, stroke, transient ischaemic attack, peripheral vascular disease and diabetes. Routine haematological indices were evaluated including full blood count, ESR, plasma viscosity, urea and electrolytes, glucose, cholesterol, lipid profile, B12 and folate levels.

Glaucoma was defined by the presence of glaucomatous changes to the optic nerve head or retinal nerve fibre layer on stereoscopic biomicroscopy or stereophotographs. The study only included patients with repeatable glaucomatous defects on SAP and those with open angles on gonioscopy. We excluded patients with secondary glaucoma, including pseudoexfoliation and pigmented glaucoma. To establish the presence of NTG, patients with glaucomatous changes and an untreated IOP <22 mmHg at presentation underwent phasing. Phasing consisted of 2 hourly IOP measurements between 8 am to 4pm. Patients with IOP measurements <22 mmHg at all times points were deemed to have NTG.

Patients with non-glaucomatous optic neuropathies were excluded. We also excluded patients with blood disorders such as polycythaemia and multiple myeloma, those with carotid atherosclerotic disease and those with visual field loss secondary to stroke or other neurological or other non-glaucomatous ocular and non-ocular pathologies.

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Results
The 99 subjects included in the study had a mean (± 12 years) age of 70.5 ± 0.5 years. The demographic and clinical characteristics of
patients included in the study are shown in Table 1. Fifty-nine (59%) of participants were female (Table 1). One hundred and sixty eight eyes (85%) had vision better than 6/12 at presentation. Twenty-three (23%) had mild glaucomatous visual field loss in the worse eye, 19 (19%) moderate loss and 57 (57%) severe visual field loss. There were no significant differences in gender (p<0.73) and age (p<0.38) between the 3 groups of glaucoma severity.

Comorbidities were common. Forty-one (41%) subjects had systemic hypertension, 28 (28%) ischaemic heart disease, 21 (21%) diabetes, 13 (13%) had a history of transient ischaemic attacks, and 7 (7%) peripheral vascular disease. 18 (18%) suffered from migraine and 3 (3%) had Raynaud’s Syndrome. Thirty-one patients were current smokers.

There were a high proportion of patients with impaired MMSE, AVLT or GHQ scores. Twelve patients (12%) had abnormal MMSEs, 7 with mild, 3 moderate and 2 severe cognitive deficits (Table 2). Ten of the 12 patients (83%) with abnormal MMSE had advanced visual field defects and of 57 patients with advanced field defects, 10 (18%) had abnormal MMSEs.

On the GHQ, 28 patients (28%) scored 5 or more, suggesting a high probability of having a psychiatric diagnosis. The highest scores tended to be in relation to somatic symptoms, followed by anxiety and insomnia then social dysfunction and finally severe depression. On the AVLT, 28% of patients were 2 standard deviations (SD) or less than expected for age and gender adjusted means in terms of short-term verbal memory performance (Figure 1), 25% were 2 SD or less for verbal learning (Figure 2) and 22% were 2 SDs or less for long-term verbal memory (Figure 3). Normally, only 2.5% of the sample would be expected to score 2 SD or less than the age and IQ adjusted population mean.

Twenty one patients (21%) admitted to having had ‘problem with their nerves’ at some stage, 11 (11%) were currently on medication for their ‘nerves’, 9 (9%) had previously seen a psychiatrist, but only 3 (3%) were currently involved with mental health services. Forty-three (43%) patients underwent neurological imaging. Thirteen patients had evidence of generalised cerebral atrophy and nine patients had micro vascular disease in excess of that expected for age. This data was reported and sourced from the specialist neuroradiology reports provided by our regional service.

Discussion

The results from this study suggest a high prevalence of cognitive impairment and psychiatric symptomatology in patients with NTG. In our cohort, 12% of patients had cognitive impairment identified on the MMSE, which is higher than one would expect for healthy subjects of a similar age. A recent meta-analysis indicated a global prevalence of dementia in relation to cognitive impairment of 5%-7% [25], in those aged 60 years or over. Our study suggests that the prevalence of cognitive impairment in patients with NTG may be twice that.

The GHQ showed that more than one in four (28%) of those with NTG had a high probability of having a psychiatric diagnosis. Furthermore, over one quarter of patients had AVLT scores 2 SD or less than the expected age matched population mean. The Rey AVLT test is a sensitive tool that measures a number of key aspects of memory, including short-term verbal memory, verbal learning and long-term verbal memory abilities. The results therefore suggest that there is a higher prevalence of short-term memory impairment in patients with NTG compared to an age matched reference population. [22-24] Moreover as some patients were excluded from the study due to advanced psychiatric illness, this further suggests a high prevalence of psychiatric and cognitive problems in those with NTG.

To our knowledge this is the first study to systematically use validated screening tools to quantify cognitive impairment and psychiatric symptomatology in patients with NTG. Previous studies have however reported a higher prevalence of dementia, cognitive impairment and depression in patients with glaucoma [26,27], and a high prevalence of glaucoma in those with Alzheimer’s disease [13,28,29]. Yochin et al. also reported correlation between depression and increasing glaucoma severity [27]. In the present study we found patients with NTG were more likely to report somatic rather than depressive symptoms using the GHQ. The high prevalence of psychiatric symptoms may also be a result of the impact of glaucoma and glaucoma treatment on quality of life. However, despite the high prevalence of cognitive impairment and psychiatric symptoms, only 3 patients were currently in contact with mental health services. This suggests that ophthalmologists should consider whether some patients with NTG might benefit from formal assessment by mental health services.

Almost 57% of patients included in this study had an advanced
visual field defect in at least one eye at presentation, which is in agreement with previous studies showing patients with NTG are more likely to present with advanced disease [30]. It has also been shown that factors including older age, African ancestry and socioeconomic deprivation are associated with delayed presentation [31-35].

Although some studies have examined the association between glaucoma and dementia [8, 10-13], few have specifically examined psychiatric and cognitive associations. Yochim BP et al. found 20% of patients with POAG had memory impairment, which is comparable to the 28% found in similarly aged patients with NTG in our study [27]. They also found, from the same sample of patients 22% had impaired executive functioning using the California Verbal Learning Test and just over 12% of patients had mild to moderate depressive symptoms [27]. The GHQ suggested that 28% of our sample had a high probability of having a psychiatric disorder, thus potentially requiring neuropsychiatric assessment. As only 3% of the patients currently had contact with mental health services, it may be advisable for patients presenting with NTG to be routinely screened for mental health problems. There may also be implications on compliance of treatment exacerbated by diminished cognition, memory impairment and possibly poor mental health. Certainly, studies have shown a poor adherence to medications related to psychosocial factors [36].

This study has several limitations. First, due to the limitations of current diagnostic tools, the use of the term NTG is controversial and may not be a distinct disease from POAG. IOP is known to fluctuate and in many patients is highest outside office hours. Although we conducted office hour phasing of IOP for all subjects, it is quite possible that patients had IOP above 21 mmHg at other times or on days when...
they were not attending the hospital for examination. Furthermore, in order to recruit sufficient numbers of patients, the study was commenced in 2009, prior to pachymetry being routinely available in our unit. Unfortunately for this reason corneal thickness data was not available for all patients. However, even if corneal thickness was known, one could argue that as IOP measurements are also affected by factors such as corneal hysteresis, the true IOP remains uncertain [37]. We therefore chose to adopt a pragmatic approach, with phasing of IOP representing current clinical practice. However, the cut off value of 21 mmHg for NTG, which is based on population data for normal IOP ranges, is not applicable for all populations and it is hard to conceive that patients with glaucoma and untreated IOPs of 20 mmHg or 22 mmHg have different diseases. It is however useful to phenotype patients and those with consistently low IOPs may have different factors contributing to glaucomatous damage, compared to those with high pressures. If further studies corroborate an association between cognitive and psychiatric difficulties and NTG, this would support the concept of a possible shared non-IOP dependent neurodegenerative pathological process. Future studies should also include patients with POAG with higher IOPs, to determine whether the high prevalence of cognitive deficit and abnormal mental health is also present in these patients.

The GHQ and MMSE are not diagnostic tests but are screening instruments for general mental health symptoms and for general cognitive impairment. The AVLT is a more specific and sensitive instrument which measures aspects of verbal memory but is not diagnosis specific. Not withstanding this, our study suggests a higher frequency of cognitive deficit and mental health symptoms amongst patients with NTG. These may significantly impact on treatment compliance, and glaucoma-associated visual loss potentially may compound the disability associated with cognitive impairment. If neuropsychiatric morbidities remain undetected it is likely to have a negative impact on ocular and systemic health and wellbeing. Few of the patients in our study were in contact with mental health services and it is likely this is reflective of a wider unmet need in health service provision in patients attending glaucoma clinics in other UK regions.

We suggest that further research is needed to explore associations between glaucomatous optic neuropathy, cognitive decline and psychiatric co-morbidities in order to better understand the relevance of shared pathogenesis and improve patient care.

Summary

What was known before:
• Glaucoma is a neurodegenerative disease.
• Glaucoma is associated with other neurodegenerative diseases such as Alzheimer’s disease.

What this study adds:
• Quantifies psychiatric and cognitive profile of patients with normal tension glaucoma.
• Suggests that NTG patients may need to be routinely screened for neuropsychiatric symptoms with further psychiatric assessment thereafter.
• Cognitive and neuropsychiatric problems could impact on a patient’s compliance with NTG treatment.
• Ophthalmic treatment may have to be modified taking into account these neuropsychiatric findings.

Contributorship statement

Mohammad Z. Mustafa (MM) wrote the article with editing from Andrew J. Tatham (AT), Bal Dhillon (BD) and Roshi Sanders (RS). The study was designed by RS and BD with glaucoma expertise from AT. Tej Rane-Malcolm carried out data collection with cognitive testing carried out by Premal Shah (PS) and Amanda Cotton. Data analysis was carried out by MM and PS. RS was project lead and also carried out all clinical ophthalmic examination.

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Competing interests
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