The prevalence of pseudoexfoliation syndrome in Chinese people

A L Young, W W T Tang, D S C Lam

Background: Pseudoexfoliation syndrome (PXS) is regarded as rare in people of Chinese ethnicity but the prevalence of this condition is not known. This epidemiology study was conducted to assess the prevalence of PXS in cataract patients and to report the clinical features present.

Methods: Prospective descriptive study conducted in the period from March 1999 to May 2001 in ophthalmology departments in cluster hospitals serving a population of about 1.2 million. 500 consecutive patients aged 60 or above attending the general ophthalmic clinics with a presumed diagnosis of cataract were recruited. A detailed examination including biomicroscopy, intraocular measurement, and gonioscopy were performed on all patients. All positive PXS cases were documented photographically.

Results: 500 patients were examined. They ranged from the ages of 60 to 91 years old, with a male to female ratio of 1:2. 40% suffered from hypertension while 24% were known diabetics. Only two positive cases (0.4%) of PXS were identified in the study population. 18% of all eyes were found to have narrow angles (defined as grade 0 to 2 by Shaffer grading). Nuclear sclerosis was the single most common type of lens opacity.

Conclusion: PXS is a rare condition in Chinese people. A prevalence rate of 0.4% in patients aged 60 or above was identified in this hospital based epidemiology study. To the best of our knowledge, this was the first study conducted in a Chinese population to examine the prevalence of PXS.

Pseudoexfoliation syndrome (PXS) was first described in 1917 by Linberg in a Finnish population. The clinical diagnosis is made by the presence of typical pseudoexfoliation material (PXM) on the anterior capsule surface. In addition to PXM, other features include endothelial pigmentation, loss of pupillary ruff, iris transillumination, Sampaolesi’s line, and pigment deposition in angle and/or posterior subcapsular (PSC). PXS was classified morphologically as follows: nuclear sclerosis (NS), cortical (C), and posterior subcapsular (PSC). PXS was diagnosed clinically by the presence of typical PXM on anterior lens capsule or at the pupil border, with or without Sampaolesi’s line and pigment deposition in angle and/or corneal endothelium. All positive cases were photographed.

RESULTS

Of 500 patients recruited, 315 (63%) were female and 185 (37%) were male; ages ranged from 60 to 91 years old. Approximately three quarters of all participants were over 65 years old (fig 1).

Forty per cent of patients had hypertension while 24% had diabetes mellitus. The prevalence of hypertension in the local general population had been reported to be 48% among those aged 70 or above, while diabetes mellitus in those older than 65 was reported to be about 25%. The participants in our study were: England (4%), Germany (4.7%), Norway (6.3%), Eskimos (0%), Russia (12%), Finland (22%), Iceland (29%), Greece (16.1%), Australia (0.98%), and Iran (9.6%). The clinical, microscopic, and ultrastructural features of PXS in Chinese patients has been described, but its prevalence has not been previously examined. Although a weak association between PXS and brown irides was described, it is generally considered as rare in Chinese people. The primary aim of our study was to assess the prevalence of PXS in a Chinese population and to report the clinical features present. In addition, information on angle status and cataract types is documented.

Abbreviations: PXM, pseudoexfoliation material; PXS, pseudoexfoliation syndrome
The study appeared to be a reasonable representation of the age matched general population.

The distribution of the morphological types of cataract observed is illustrated in figure 2. The most common presentation was mixed cataract with both nuclear and cortical elements (50% of all cases) while NS was the single most common morphology type (present in 99%). Cortical and posterior subcapsular opacities were observed in 69% and 19% respectively.

Eighteen percent of all eyes were found to have narrow angles. The percentage of narrow angles observed was higher in females than males, at 22% and 12% respectively. The difference was statistically significant ($\chi^2$ of independence: $\chi^2 = 16.62; df = 1; p < 0.0001$).

Only two out of 500 patients (0.4%) were found to have PXS. Both were male and aged 64 and 66 years old respectively. They were clinically diagnosed by the presence of typical PXM on the peripheral anterior lens capsule. Both patients had only unilateral ocular involvement, with the absence of any additional features. The angles were open, and intraocular pressure and cup:disc ratios were found to be normal. Both patients had nuclear sclerosis without any phacodonesis.

**DISCUSSION**

Although all cases of PXS in this study were men, no definite sex predominance has been previously shown. Clinical PXS may present unilaterally, as observed in the two positive cases of this study. Our two cases did not have glaucoma at the time of diagnosis. As PXS has an increased predilection for both open and narrow angle glaucoma, intraocular pressure in both eyes should be regularly monitored. Acute angle closure glaucoma is common in Hong Kong and narrower angles were considered to be related. In this study, 18% of all included participants had narrow angles, which is much higher in comparison with a prevalence of 3.8% of narrow angles reported in the Framingham study.

Nuclear sclerosis was the commonest type of lens opacity observed, followed by cortical and posterior subcapsular opacities. The results are similar to findings reported in the Beaver Dam Eye Study and Framingham Eye Study. In contrast, cortical opacities were most prevalent in African populations.

Cataracts are known to be more common in PXS. Ascorbic acid is significantly reduced in the aqueous of cataract patients with PXS. One may therefore consider the possible association between oxidative stress and UV exposure in PXS and hence the reason for the association between the two. The prevalence of PXS in patients with cataract had been reported to be 17.7%, 23.5%, and 28.7% in Turkey, Portugal, and Spain respectively. In contrast, our prevalence of 0.4% (similar to Eskimos—possibly related to a common ancestry) is much lower in comparison.

One must stress that precise figures on prevalence can only be obtained by studying population groups and that the current small scale hospital based study on patients with cataracts would lead to bias. In addition, the true figure could have been underestimated, as the clinical signs in PXS are known to be subtle in the early stages. Furthermore, it is regarded difficult to detect PXS in patients with cataracts. In conclusion, PXS is a rare condition in Chinese people with a prevalence rate of 0.4% in patients with cataract aged 60 or over. To the best of our knowledge, this was the first study conducted in a Chinese population to assess the prevalence of PXS.

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