Acquired colour vision deficiency in patients receiving digoxin maintenance therapy

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Background/aims: Disturbances of colour vision are a frequently reported sign of digoxin toxicity. The aim of this study was to investigate the incidence of acquired colour vision deficiency in elderly hospitalised patients receiving maintenance digoxin therapy.

Methods: 30 patients (mean age 81.3 (SD 6.1) years) receiving digoxin were tested using a battery of colour vision tests (Ishihara, AO Hardy Rand Rittler plates, City tritan test, Lanthony tritan album, and the Farnsworth D15). These were compared to an age matched control group. Serum digoxin concentrations were determined from venous blood samples.

Results: Slight to moderate red-green impairment was found in approximately 20–30% of patients taking digitals, and approximately 20% showed a severe tritan deficiency. There was no correlation between colour vision impairment and serum digoxin level.

Conclusions: Formal colour vision testing of elderly patients taking digitals showed a high incidence of colour deficiency, suggesting that impairment of retinal function can occur even at therapeutic drug levels. As a result, colour vision testing in this population would have limited value for the detection of drug toxicity.

Digoxin remains a frequently prescribed cardiac glycoside used in the treatment of supraventricular dysrhythmias and congestive cardiac failure. Cardiac glycosides have a narrow therapeutic index, and there is considerable overlap in serum concentration of the drug between patients with and without toxicity. Although the clinical diagnosis of digoxin toxicity has fallen substantially over the past 20–30 years, it remains a common medical problem, particularly in the elderly, where it is often difficult to diagnose. Features of digoxin toxicity are usually non-specific and consist of cardiac and non-cardiac effects. Non-cardiac manifestations include symptoms of fatigue, anorexia, nausea, vomiting, headaches, and confusion. Disturbances of vision—for example, blurring, central scotomas, glare effects, and altered colour perception, are a less common but more specific presenting complaint. Symptomatic colour vision disturbances—for example, xanthopsia, are found in up to 15% of intoxicated patients, although formal testing reveals a much higher incidence of colour deficiency. Impairment in both the tritan and red-green (R/G) axes have been found. Using the Farnsworth-Munsell (FM) 100 hue test, several studies have reported a strong positive correlation between the total error score and serum digoxin level, suggesting that colour vision assessment may be used as a predictor of toxicity.

The aims of the present study were to use rapid screening tests for tritan and red-green colour defects to investigate the incidence of acquired colour deficiency in a population of elderly patients receiving maintenance digoxin therapy, and furthermore, to reassess the relation between serum digoxin concentration and colour vision impairment.

Subjects and methods
Colour vision was assessed in 30 hospitalised patients (10 males and 20 females, mean age 81.3 (SD 6.1) years) who were receiving oral digoxin maintenance therapy. The daily dose varied from 62.5 µg to 250 µg (mean 127.1 (44.9) µg), and the mean duration of treatment was 32.4 (62.8) months. Patients with glaucoma, diabetes, retinal/optic nerve disease, significant cataract, and congenital colour deficiency were excluded, as were patients taking concomitant medication known to affect colour perception. The visual acuity of all subjects was 6/12 or better. A battery of pseudoisochromatic plates and a hue discrimination test were used for colour vision assessment. The tests used and their scoring criteria were as follows:

Ishihara pseudoisochromatic plates (38 plate edition)—Plates 1–17 were used on all patients to identify red-green colour deficiency. Patients with greater than three errors were classified as failing the test.

AO Hardy Rand Rittler (HRR) plates—The HRR test identifies and grades the severity of red-green and tritan defects. The defect can be classified as slight or significant (moderate or severe) depending on the number of errors.

City tritan test—This test has five plates. Three plates have small colour differences, identifying slight tritan defects, and two plates have large colour differences to identify severe tritan deficiency.

Lanthony tritan album—The tritan album has six plates. Severity of tritan deficiency (slight, moderate or severe) is determined by the number of errors.

Farnsworth D15—This hue discrimination arrangement test uses 15 coloured caps which are arranged in sequence from a fixed pilot cap. The results are plotted on a hue circle diagram, which allows tritan defects, red-green defects or generally poor hue discrimination to be identified.

Colour vision testing was carried out in good daylight conditions, supplemented when necessary with artificial lighting. Thirty age matched control subjects (eight males and 22 females, mean age 76.8 (11.5) years), who were also hospital inpatients but were not taking digoxin, were examined with the same test battery. The serum digoxin level was determined by immunoassay on samples of venous blood taken at least 6 hours after the last dose.

Results
Plate tests
Subjects receiving digoxin showed a significant number of red-green and tritan colour vision errors (Tables 1 and 2). The Ishihara test identified red-green deficiency in 23.3% of subjects. A greater number failed the red-green plates of the AO HRR test (20% with slight defects and 13.3% showing significant red-green defects). By contrast, no control subject failed the Ishihara or the red-green grading plates of the HRR.
Although tritan errors were common in controls, these defects were typically only slight or occasionally moderate (Table 2). By contrast, approximately 20% of subjects in the digoxin group showed a severe tritan defect (Table 2). Figure 1 shows the mean number of red-green and tritan errors recorded in control and digoxin groups. These differences were statistically significant. However, there was no correlation between serum digoxin concentration and the number of red-green errors ($r^2 = 0.0009$, $p = 0.87$), tritan errors ($r^2 = 0.0175$, $p = 0.49$) or total errors ($r^2 = 0.001$, $p = 0.67$) (Fig 2). Similarly, there was no correlation between the total number of errors and duration of treatment ($r^2 = 0.003$, $p = 0.83$).

Using the D15 panel, 26.7% of the digoxin group showing characteristic tritan confusion errors, with 30% showing an overall poor hue discrimination (Fig 3). No patient showed confusions along the deutan or protan red-green axes. Although control subjects commonly made non-specific arrangement errors, there were no errors along the tritan confusion axis.

**DISCUSSION**

Simple screening tests for colour deficiency revealed a high incidence of colour vision impairment in a population of elderly hospitalised patients on maintenance digoxin therapy compared to an age matched control group of hospital inpatients. None of the patients reported any subjective alteration of colour perception. Although 10% of patients were found to have a serum drug concentration of $>2.6$ nmol/l, the mean serum digoxin level for the group was in the middle of the therapeutic range and no patient had clinically defined digoxin toxicity. Both red-green and tritan errors were identified. Slight to moderate red-green impairment was found in approximately 20–30% of patients, and approximately 20% showed a severe tritan deficiency. These data suggest that a subclinical impairment in visual function can occur even at therapeutic doses of digoxin. There have been several case reports in the literature of overt visual disturbance—for example, photopsia and reduced acuity, in patients with blood digitalis levels within the therapeutic range. In all cases discontinuation of therapy was followed by complete resolution of symptoms. The reasons for the eye’s susceptibility to the untoward effects of digitalis are not fully understood. There is evidence that the retina is the primary site of drug toxicity. Several retinal cells express digitalis sensitive isoforms of sodium-potassium ATPase (Na$^+$K$^+$ ATPase)—for
example, photoreceptors, Müller cells, and the retinal pigment epithelium. Clinical electoretinography and in vitro cell studies have shown that toxic levels of digoxin lead to rod and cone dysfunction, with cones being affected to a greater extent. These effects were reversed following dose reduction or cessation of therapy.

All the patients in the present study were elderly, and advanced age appears to be a contributory factor in general for drug toxicity, including for digoxin. The basis for the increased susceptibility to the toxic effects of digitals in the elderly is unclear, although it may be the consequence of altered renal function, electrolyte disturbances, or changes in the density or character of the drug receptor. The compromised vascular status of the digoxin treated cohort may have also contributed to the observed disturbance in colour vision. It has been suggested that the effects of digoxin on colour vision may be continuous, commencing with a slight impairment of hue discrimination and progressing initially to a tritan defect, and then to a red-green deficiency as toxicity increases. The colour vision defect most commonly seen in the present study was in the tritan axis. It is not clear whether these defects reflect a particular vulnerability of the short wave (S) cone pathway to digoxin or the lower density of S cones. Advanced age is known to adversely affect blue-yellow discrimination as a result of reduced transmission of short wavelengths of light through the ocular media. Slight or moderate tritan errors were common in age matched controls, however severe defects were only observed in patients taking digoxin. The finding of red-green defects in patients taking digitals in the absence of toxicity was surprising, as red-green deficiency has previously been reported only in severe digitalis intoxication.

However, severe defects were only observed in patients taking digoxin. The finding of red-green defects in patients taking digitals in the absence of toxicity was surprising, as red-green deficiency has previously been reported only in severe digitalis intoxication. By contrast with previous studies on younger populations of patients taking digitalis, no correlation was found over, the high incidence of colour vision impairment in the elderly in the absence of toxicity was surprising, as red-green deficiency has previously been reported only in severe digitalis intoxication.

In summary, elderly patients receiving maintenance digoxin therapy showed a high incidence of colour vision impairment. Disturbances in both the tritan and red-green axes were present, in the absence of a subjective alteration of colour perception. By contrast with previous studies on younger populations of patients taking digitals, no correlation was found between colour vision loss and serum drug concentration. These findings suggest that routine colour vision testing in elderly patients receiving digoxin would have limited value for the detection of drug toxicity.

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