length determinations using the Genescan software on an ABI 373A automatic sequence. Southern blotting of DRPLA Southern blots contains a PCR with one primer labelled with fluorescent dye. For the statistical comparison of the distributions of CAG copies for the three samples, the p value from the exact procedure for contingency tables was calculated using StatXact's Monte Carlo approach with 1,000,000 replications. Thus the width of the 99% confidence interval for the p value could be calculated to 0-001 indicating that the p value given is at least to the second decimal.

The distribution of repeat sizes in the DRPLA gene in 47 male and female controls (94 chromosomes) varies between seven and 25 CAG copies with a maximum for [CAG]_min in 29% of the chromosomes. In two cases (4%) homozygosity has been observed.

In 35 patients with questionable diagnoses of SCA1 without expansions in the ataxin gene, seven to 21 CAG copies in the DRPLA gene have been found, including 26 of 70 alleles with [CAG]_min (37%). No significant differences in the repeat frequency of the DRPLA gene between controls and patients with ataxia could be ascertained. For seven patients only one allele could be amplified, indicating 20% homozygosity. This indicates that one CAG allele is too small to be amplified in a small number of cases. Insufficient amplification of an expanded repeat appears improbable since DNA analyses of two affected persons, kindly provided by Dr T Warner and A Harding, London, have been successful.

Investigating the DNA of 73 patients suspected of having HD but negative for the mutation in the IT15 gene, alleles ranging from [CAG]_min in the DRPLA gene were identified. Two homozygous DNA samples (3%) were present. Surprisingly, in this collection the second most frequent allele was [CAG]_min, the most frequent being [CAG]_max (30% of chromosomes). Twenty-six of the 146 chromosomes investigated (18%) contain 10 CAG copies representing about 36% of the patients with symptoms of HD. Controls and patients shared a common ethnic background and, therefore, differences between populations could not account for this result. Furthermore, the allele distribution in our controls corresponds to data of the white population, namely the alleles [CAG]_20 and [CAG]_18 being overrepresented in the Japanese population, too.

Allele distributions in the DRPLA gene in the three groups are summarised in the figure. Neither in our patients with chorea nor in patients with symptoms of ataxia could an expansion of the CAG repeat on chromosome 12, to explain the affected status, be found. Our data confirm the low frequency of the DRPLA mutation in the European population. Interestingly, the incidence of allele [CAG]_max is significantly increased in patients with a questionable diagnosis of HD. The exact p value for the comparison of the distributions of CAG copies is calculated as p = 0.003 which indicates a significant discrepancy between the samples. This result may be due to linkage disequilibrium between the disease and a mutation on chromosome 12 or, possibly, a further gene causing a progressive neurodegenerative disorder is located in the DRPLA region. Such a role of genes with polymorphic markers are necessary to define a high risk haplotype for this genetically uncharacterised disease. Investigations of pedigrees with affected persons with the allele [CAG]_max may detect the presence of a gene responsible on chromosome 12.

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Detection of Y mosaicism in patients with Turner’s syndrome

The very high intrauterine loss rate in Turner’s syndrome and the observation that there is a higher ratio of mosaic karyotypes to monosity X in liveborns compared with aborted fetuses has led to the speculation that all liveborn patients are mosaic for a critical cell line.

Experimental evidence has shown that 70 to 80% of patients with Turner’s syndrome retain the maternal X chromosome. Therefore, there are the theoretical possibility of Y mosaicism occurring in 35 to 40% of Turner’s syndrome patients.

This has not been shown in cytogenetic studies where only 5% of cases have Y mosaicism and another 3% have unidentified markers. However, low level mosaicism may be missed by conventional cytogenetic techniques and over 100 cells would need to be examined to exclude 5% mosaicism with 99% confidence. This fact is also a limitation in the use of FISH which has an established role in the identification of fragments.

Identification of Y chromosomal mosaicism is of clinical relevance because of the high risk of tumour formation in the dysgenetic gonad components with either or structurally abnormal Y chromosomes present, which may be as high as 30%. Page has hypothesised the presence of a ‘gonadoblastoma’ locus which is probably located in interval 4b–5.

Several studies on relatively small numbers of patients have investigated Y mosaicism in Turner’s syndrome using Southern blotting with Y-specific probes, PCR with one or two Y primer sets, and even Southern blotting of amplified DNA and have suggested that occult Y mosaicism occurs in 0 to 33% of patients. In view of this lack of consensus, we have used PCR with a panel of Y primers from the whole length of the Y chromosome to determine the frequency of occult Y mosaicism. As a part of a total ascertaining study, 100 patients with Turner’s syndrome were recruited from the four growth clinics and from adult Turner’s syndrome clinics in Scotland. The patients had the following karyotypes: 45,X/46,XX in 70/35; 45,X/46,XX(Xr) in 10, 45,X/46,XX in seven, 45,X/46,XY in three, 45,X/46,Xi(Y)p in one, 45,X/46,X M in nine, and other karyotypes in eight.

DNA was extracted from peripheral lymphocytes of all patients by the same normal female (CEC, karyotype 46,XX) and PCR was carried out with nine sets of primers from the Y chromosome: PABY from the pseudoautosomal region which amplifies X and Y specific products of different sizes, SRY,11 amelogenin12 which amplifies X and Y specific products with a size difference, Y chromosome13,14 YSR1,15 SRY,16 and SY160,17 Y1.1.18 An additional primer set was used in patients who were positive with Y1.1.1, SY146 and SY160 were amplified in a multiplex reaction. Such PCR products were resolved on 1–2% agarose gels stained with orange G bromide stained agarose gels with both normal and normal female controls on every gel.

Southern blotting of amplified DNA was also carried out on 84 of the patients who had been shown to be negative on direct visualisation of PCR products. Three primer sets were used initially, SY1, Y centromere, and SRY, and a fourth SY110,13 5,7,19 9,20 from the SRY gene was used on positive cases. Great care was taken to prevent contamination either by PCR products or genomic DNA when setting up the PCR reaction or running gels. All the experiments were carried out by the same female (46,XX karyotype). Separate pipettes were used for setting up the PCR and running gels, autoclaved solutions were used, and all solutions were subject to UV light for 10 minutes before use. The PCR products were run in a separate laboratory one floor below that used for setting up the PCR. All duplicate were resolved in two different DNA samples extracted from fresh blood samples. In five other cases different DNA samples were used.

Forty-six normal females were also screened using the SY1 primer set. In 40 of these the DNA was extracted by female laboratory staff and in six by a male scientist. Probes used for the Southern blotting were gene cleaned PCR products amplified from normal males. These were labelled with 3PdCTP by the random labelled method.

Parental origin of the normal X chromosome was determined either by use of Southern blotting with DNA from the parents. In 40 by PCR with primers from a repeat sequence in the androgen receptor gene, as previously described. Concurrence of X inactivation patterns in patient 9 assisted in the identification of the normal X chromosome.

Nine patients were positive for one or more Y regions on direct visualisation of ethidium bromide stained gels with the results shown in Table 1 along with the percentages of the primers used. Three of these were already known to have a Y chromosome cytogenetically and were positive for all regions tested (patients 3, 4, and 5, table 1). One of these subjects (patient 3) had a prepubertal
gonadectomy at the age of 6 years and was found to have a gonadoblastoma in the dysgenetic gonads. Patient 1 presented at the age of 17 years with primary amenorrhoea and was found to have a 45,X line and also a second cell line containing a small unidentifiable marker. She had no features of virilisation. She was positive for all primer sets tested as were the gonads which were removed after the identification of the Y material. Patient 2 had a 45,X/46,X,+M karyotype. She had no virilisation features and no secondary sexual characteristics. Gonadectomy was carried out and no evidence of gonadoblastoma was found in the streak gonads.

Patient 6 had a 45,X/46,X(Y)p karyotype and presented at birth with ambiguous genitalia. Prophylactic gonadectomy was performed at the age of 1 year and a testis was identified on the right and a streak gonad on the left with normal female internal organs. There was no trace of tumour found in the gonads. On PCR testing this patient had positive Y sequences down to and including vY146 (interval 6). Patient 7 was originally thought to have a 45,X karyotype. She had no signs of virilisation. The karyotype was reviewed in the light of the PCR result and a total of 95 cells were then counted before an (Yq) was found in 10% of cells. FISH analysis was also carried out but again was negative until the third review of the slides. She underwent gonadectomy at the age of 9 years and no evidence of tumour was found on histology.

Patient 8 was reported to have a 45,X karyotype and had the features of Turner's syndrome with no additional features of note. She developed a primary lung carcinoma with cerebral metastases at the age of 16 years and died two years later. She had no evidence of ovarian pathology on CT scan. She had only distal Yq sequences present on PCR analysis.

Patient 9 had a 45,X/46,X(Xq) karyotype and has features of Turner's syndrome but had spontaneous puberty. She is positive for distal Yq sequences.

After the finding of distal Yq sequences in patients 8 and 9, DNA from the mothers of both these patients was analysed for Y sequences. The mother of patient 8 was also positive for the same distal Yq sequences but not any other Y sequences and the mother of patient 9 was negative for all sequences tested.

Using the method of Southern blotting of amplified DNA, no patients were consistently positive for either sY58 or Y centromere sequences. Seventeen (17/84, 20%) Turner patients were positive for SRY1 sequences. The signals obtained varied in intensity and were only visible after overnight exposure. Fourteen of the normal females (14/46, 30%) also showed positive signals using this primer set and the signal intensity varied in intensity. Only one of the samples extracted by a male scientist showed a positive signal.

A separate experiment using XES10/11 primers was then carried out for those patients and controls positive with the SRY1 primers. This set of primers is within the coding region for SRY but both primers are outside the HMG box. None of the samples showed positive signals. Serial dilution of male DNA showed that this method was able to detect 0.0001 µg male DNA.

Parental blood was available from all patients except patient 2 and the normal X chromosome was found to be maternal in origin for all the other patients.

We have shown that 9% (9/100) of Turner's syndrome patients from a large unselected group have Y mosaicism, fewer than half of whom were identified by cytogenetic analysis. If patients with known Y material and unidentified markers are excluded from the analyses, -3% (3/87) of patients in this cohort were found to have unexpected Y mosaicism. This is a slightly higher figure than that found in other studies to date (table 2) and probably reflects the fact that in this study multiple primer sets were used.

The present study highlights several different points. Firstly PCR can detect low level mosaicism missed by conventional cytogenetic analysis of relatively large numbers of cells, as in patient 7. Secondly, more than one area of the Y chromosome should be screened. Many of the previous studies have used SRY primers and the use of such primers alone would have missed four of our nine positive cases (patients 2, 7, 8, and 9). Also by using multiple primer sets we were able to define the content of the Y isochromosome in patient 6. This patient may have been thought to be at low risk of tumour formation as she presumably has mainly short arm material on(Yp)). However, using PCR it has been shown that a significant amount of long arm material is present in the isochromosome including the at region 7 (table 1).

Thirdly, Y material may be present even in patients with other mosaic karyotypes (patient 9, table 1). This confirms findings which have been noted in two other patients in different reports. Medeiros et al10 found SRY sequences on PCR with direct visualisation in a patient with a 45,X/46,XX karyotype in streak ovaries and Rosenmund et al9 described a patient with an XY translocation using Southern blotting for those Y material detected in patients 8 and 9 is uncertain as it probably represents only distal Yq sequences. The finding of these distal sequences was confirmed by using a second primer set (sY159) from the same region.

We have so far been unable to ascertain the location of these sequences using FISH but this may be because the sequences are beyond the limit of resolution of this technique. An explanation for the Y long arm sequences present in these patients may be that although it is generally supposed that pairing of the X and Y chromosomes during meiosis only involves the distal third of the X short arm and almost all the Y short arm up to the centromere, studies of the XY hybrid by electron microscopy have shown that it may in fact be much more extensive.25 This hypothesis is supported by the fact that the mother of patient 8 was also positive for the distal Yq sequences thus indicating that these may represent a familial translocation.

Although the repetitive sequences DYZ2 have been thought to map to the heterochromatic region only and thus represent genetically inert material, Young et al9 found these sequences in the non-fluorescent band Yq11 so that they may have some effect. At present, patient 9 has died but had no evidence of gonadoblastoma at the time of death.

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<th>PHRY</th>
<th>SRY</th>
<th>AMEL</th>
<th>CEN</th>
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SB - Southern blotting, cen - centromere.
positive with one set of SRY primers (SRY1) but all were negative for Y centromere and sY85 sequences and a similar percentage of normal female controls (14/46, 30%) were also positive using this method of analysis. Southern blotting of amplified DNA is extremely sensitive and very prone to contamination; however, great care was taken during all our experiments to prevent contamination. The positive results cannot be the result of contamination by genomic DNA as only one locus was affected and positive cases were retested on different DNA samples. Contamination by PCR products is also extremely unlikely, firstly because different pipettes were used to set up the PCR from those used to run out the products, the laboratory used to run out products was situated on a different floor from that used to set up the experiment, and, most importantly, the intensity of the positive signals seen in both the patients and controls varied from person to person.

The explanation for the positive signals in these patients and controls may be that the PCR primers used by Kocova et al2 and ourselves amplify a 270 bp sequence representing a DNA binding motif known as the high mobility group box (HMG box). This motif shares homology with a DNA binding motif present in non-histone proteins related to HMG1 and HMG226 and a DNA binding domain of H/puf (human upstream binding factor)27 and possibly other regions of the genome as well. It is possible that the signals detected in Turner’s syndrome patients and normal female controls represent amplification of partially homologous sequences from these other regions. This theory is supported by our finding that patients and controls previously positive for SRY1 were negative using primer sets XES10/11 which are still within the coding sequence for SRY but are located well outside the HMG box. These patients were negative in spite of the sensitivity of this particular assay which detects down to 0.0001 μg of male DNA. Southern blotting of amplified DNA did not identify any patients not previously picked up using direct visualisation of PCR products and we would therefore not advocate its routine use. However, it may be useful in confirming ambiguous results if great care is taken in the experimental methodology.

The clinical significance of occult Y mos- saicism is unknown but in our experience PCR (without Southern blotting of amplified DNA) is more effective than routine cyto-

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11 Yankowitz J, Neely EK, Haidu K, Norton ML, Rosenberg RG.戈尔川. Genetic and phenotypic analysis for its detection and may be further enhanced by screening multiple tissues.

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Detection of Y mosaicism in patients with Turner's syndrome.

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