

A molecular deletion map of the Y chromosome long arm defining X and autosomal homologous regions and the localisation of the HYA locus to the proximal region of the Yq euchromatin

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ABSTRACT

41 Y-linked DNA probes that detect sequences on the Y chromosome long arm have been used to analyse genomic DNA from a series of 23 patients with deletions of Yq. Southern blot analysis has differentiated 15 distinct breakpoints, which divide Yq into 14 mapping intervals. From the pattern of DNA sequences present in each patient, it has been possible to produce a congruent deletion map, with the exception of two cases which are not compatible with the consensus order. These patients can be explained by the presence of inversion polymorphisms on Yq in the general population or by complex rearrangements induced during the formation of the deleted chromosomes. The distribution of sequences on the Y long arm has defined distinct regions of homology with autosomes, the Y short arm and the long and short arms of the X. A number of the patients have been typed for the presence or absence of H-Y antigen (as determined by the cytotoxic T-cell assay) and it has been possible, from analysis of informative cases, to assign the locus to the proximal region of the Yq euchromatin.

INTRODUCTION

The failure of the differential segment of the Y chromosome (which includes the entire long arm) to undergo recombination at meiosis precludes the use of genetic analysis to develop maps of this sex chromosome. Therefore, it has been necessary to rely upon the use of patients with cytogenetically visible abnormalities of the Y to develop physical deletion maps^{1–5} and to assign genetic functions to sub-regions of the chromosome. Such analysis has indicated that a locus encoding or controlling the expression of the H-Y antigen⁶ and that loci affecting tooth size^{7,8} and spermatogenesis^{9,10} are located in the euchromatic region of the Y long arm.

Of particular value in developing deletion maps of Yq are individuals who possess a chromosome isodicentric for Yp, where a breakpoint on Yq can be identified. At the cytogenetic level, the differentiation of breakpoints in these cases is limited. However, analysis of genomic DNA from these patients with cloned Yq sequences allows the definition of a number of deletion intervals, thus permitting more precise mapping and genotype-phenotype correlations.

In this paper we have used cloned Yq DNA sequences (isolated from Y chromosome-specific libraries) to analyse DNA from patients with structural abnormalities affecting the Y long arm. This analysis has differentiated 15 breakpoints amongst 23 patients and has permitted the assignment of Yq sequences to 14 mapping intervals. H-Y antigen typing of a number of these patients has allowed the assignment of the HYA locus to a region covering the proximal segment of the Yq euchromatin. Further, the patterns of homology of these sequences to other chromosomes has allowed further elucidation of the organisational structure of the euchromatic region of Yq.

RESULTS

Isolation of Yq DNA sequences

Single-copy DNA sequences (isolated from either a flow-sorted chromosome Y-specific library or Y cosmids derived from a Y-only cell hybrid library) which map to Yq were identified by hybridization to the panel illustrated in figure 1. The key members of this panel are the Y-only somatic cell hybrid 7631, ED (a patient with two X chromosomes and two dicentric iso Yq chromosomes with a breakpoint in Yp at Yp11.2) and WC (a patient with a single X chromosome and a monocentric iso Yp chromosome generated by centric fusion). Sequences from the Y were identified by hybridization to Y-only hybrid DNA, those located on proximal Yp by hybridization to both WC and ED DNA, those located on distal Yp by hybridization to WC (and not ED) DNA and those located on Yq by hybridization to ED but not WC DNA. Any X homologies were indicated by hybridization to the DNA of the X-only somatic cell hybrid Horl X. Several of the probes detect homologies in other regions of the Y, some on the X and others on autosomes.

Analysis of patients with Y long arm breakpoints using Yq DNA sequences

The ideograms in figure 2 present a schematic summary of the aberrant Y chromosomes revealed by cytogenetic analysis of our panel of patients. Two categories of dicentric iso Yp chromosome have been distinguished amongst these patients with breakpoints (based on the banding pattern of the Y described by Magenis¹⁸) in Yq11.21 and Yq11.22. In addition, a patient with the derivative

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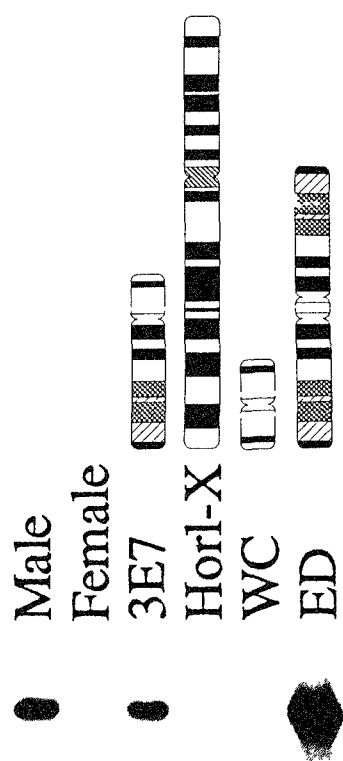


Figure 1 Mapping panel for the rapid regional assignment of Y probes. Hybridization of probes to this panel allowed their localization to the long or short arm of the Y chromosome and identified any X or autosomal homologies. The figure shows the hybridization of the probe GMGY21 which maps to the long arm of the Y. ED has two isodicentric Yq chromosomes with a breakpoint at Yp11.1. WC has a monocentric isoYp chromosome. 3E7 and Horl X are Y only and X only somatic cell hybrids on a mouse background.

5 from a 5 Y translocation, a patient with the derivative 14 of a 14 Y translocation, a patient with a ring Y, a patient with a Yq deletion with a breakpoint at Yq 11.23 and three patients with the derivative X from X Y translocations have contributed to the collection of Yq breakpoints. In all, the panel consists of 23 patients. The data from the analysis of this panel with the full set of Yq DNA sequences is summarised in table 1. The Yq sequence scored by probe 50f2 is the E fragment.

From the profile of DNA sequences in these patients it is possible to construct a linear series of breakpoints delineating 14 mapping intervals labelled YQ1–YQ14. This is shown in

Table 1 The pattern of Yq sequences present in members of the deletion panel

	W	A	D	K	M	D	J	C	F	F	C	I	K	R	Z	J	D	M	P	F	H	J	J
	C	M	F			B	G	O	B	F	C	T	M	S	A	C	M	N	M	W	M	H	L
					M																		
GMGY4C	+																						
CMPY20	+																						
CMPYF1																							
GMGY6																							
pDP320a																							
GMGY3																							
OX3																							
OX5																							
CRI232																							
STB14																							
YEX8																							
GMGY19																							
GMGY37																							
GMGY38																							
GMGY13																							
C1B																							
GMGY29																							
GMGY30																							
PVY64																							
M1A																							
GMGY15																							
GMGY33																							
GMGY12																							
50f2E																							
PVY65																							
COS40T																							
GMGY36																							
GMGY39																							
GMGY26																							
cCMPY3																							
OX2																							
OX7																							
GMGY10																							
GMGY14																							
GMGY20																							
GMGY21																							
GMGY18																							
OX1																							
cCMPY4																							
GMGY1																							
GMGY28																							

(+) indicates the presence of a Yq signal and () indicates the absence of a Yq signal. () indicates not determined.

figure 3. Two patients (DF and K M) do not fit into this map, both have distal Yq sequences in the absence of those located in a more proximal position. These data expand and revise the previously published³ analysis of Yq breakpoints, altering the intervals defined in that study.

In addition to detecting Y specific DNA fragments, several of these probes also have X or autosomal homologies and some detect sequences which map to other regions of the Y chromosome. These data are summarised in table 2. Some of these homologies are apparent at low stringency only, while others are retained at high stringency washing conditions.

Localisation of the HYA locus using the Yq panel

Expression of the H Y antigen, a minor histocompatibility antigen which causes rejection of male to female grafts within inbred strains of mice, has been mapped in humans previously to the Y chromosome long arm or centromeric region⁶. In an attempt to delineate this interval more precisely, H Y antigen status, using the cytotoxic T cell assay⁶, was determined for informative members of the deletion panel. The data for patients WC, JC, DM and FW are summarised in table 3. It was not possible to type all individuals because of MHC restrictions of the H-Y specific T cell clones available. Analysis of the informative cases indicates that the region of the Y responsible for H Y antigen expression corresponds to the intervals between CEN and YQ8 as indicated in figure 3. From the cytogenetic analysis of these patients this places the locus in Ycen Yq11.21.

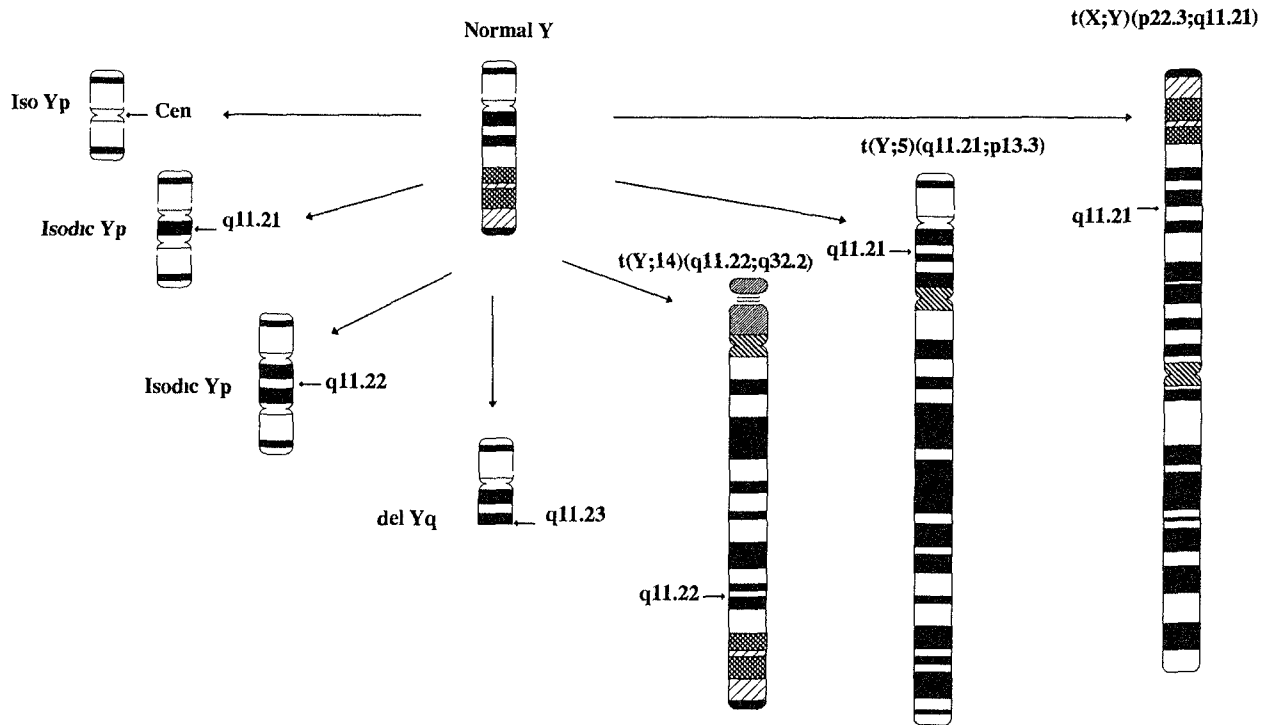


Figure 2. The figure summarises the categories of Y abnormalities found amongst the patients of the deletion panel. DG (ring Y) is not illustrated.

DISCUSSION

Molecular analysis of these patients with probes that detect sequences on Yq has revealed the existence of 15 breakpoints and made possible the construction of a Yq deletion map in which Yq probes have been assigned to 14 intervals. In so doing, the precision of karyotype-phenotype correlations in patients with Y abnormalities will be improved and further enhanced as similar Yq maps assembled by others^{4,19-22} are integrated into a unitary description of Yq.

Through mutual typing of patients CO, FF, RS, JC and DM we have been able to cross-reference to a limited extent our Yq intervals to the deletion map of David Page's laboratory (Douglas Vollrath, personal communication and manuscript in preparation), and, consequently to the original map of the Y produced by Vergnaud et al.⁴ The Vergnaud map intervals describing Yq are shown aligned against our own intervals in figure 3. Patients CO, FF and RS have breakpoints in interval 5 of the Vergnaud map, and patients JC and DM in interval 6. Thus the boundary between intervals 5 and 6 of the Vergnaud map occurs between patients JC and DM on the distal side, and patients FF and RS on the proximal side. Interval 4B defines the centromere and interval 7 marks the start of the distal heterochromatic repeats. From this, it is possible to assign our intervals YQ1–YQ6 as sub-divisions of the Vergnaud interval 5 and the distal segment of YQ8 to interval YQ14 as sub-divisions of interval 6 since patients JH and JL do not contain any of the heterochromatic repeats of distal Yq (corresponding to interval 7). At this point we cannot assign interval YQ7 to interval 5 or 6 and similarly, for the proximal part of interval YQ8. The order of breakpoints defined by the two maps for the patients typed in common using

Table 2. Summary of homologies detected by Yq probes

No Homologies	Yq X	Yq Autosomal	Yq Autosomal/X	Yq-Yp
GMGY6 (2)	PDP320A (2)	GMGY4C (cen)	CMPY20 (cen/Yp)	50I2 (8)
YEX8 (3)	GMGX3 (3)	OX3 (3)	GMGY37 (3)	GMGY39 (10)
GMGY13 (3)	OX5 (3)	OX5 (3)	GMGY38 (3)	GMGY26 (10)
GMGY30 (5)	cri232 (3)	C1B (3)	PVY64 (5)	cCMPY3 (10)
GMGY15 (6)	STS (3)	50I2 (8)	GMGY12 (8)	OX2 (10)
GMGY33 (7)	GMGX19 (3)	OX2 (10)	PVY65 (8)	OX7 (10)
COS40T (9)	m1a (5)	OX7 (10)	GMGY39 (10)	GMGX10 (11)
GMGY36 (9)	GMGX10 (11)	cCMPY4 (13)	cCMPY3 (10)	GMGY14 (11)
		GMGY28 (14)		OX1 (12)
		CMPY21 (12)		
		GMGY29 (4)		
		GMGY26 (10)		
		GMGY21 (12)		
		GMGY1 (13)		
		GMGY20 (11)		
		GMGY18 (12)		

The numbers in brackets refer to Yq deletion intervals YQ1–YQ14. The category Yq autosomal/X contains probes whose homologies have not yet been assigned to autosomes or the X chromosome.

a completely non-overlapping set of markers is identical, except that we have been able to resolve patients DM and JC as separate breakpoints.

The development of a molecular deletion map of Yp has been achieved in a similar way by studying X-Y interchange in XX male patients¹². Here too, apparent exceptions to the consensus order (as with patients K-M and DF for the Yq map) were discovered and could reflect inversion polymorphisms of the Y in the general population or rearrangements induced during the formation of these aberrant Y chromosomes. In this context, Donlon and Muller²³ have provided evidence of Yq sequences

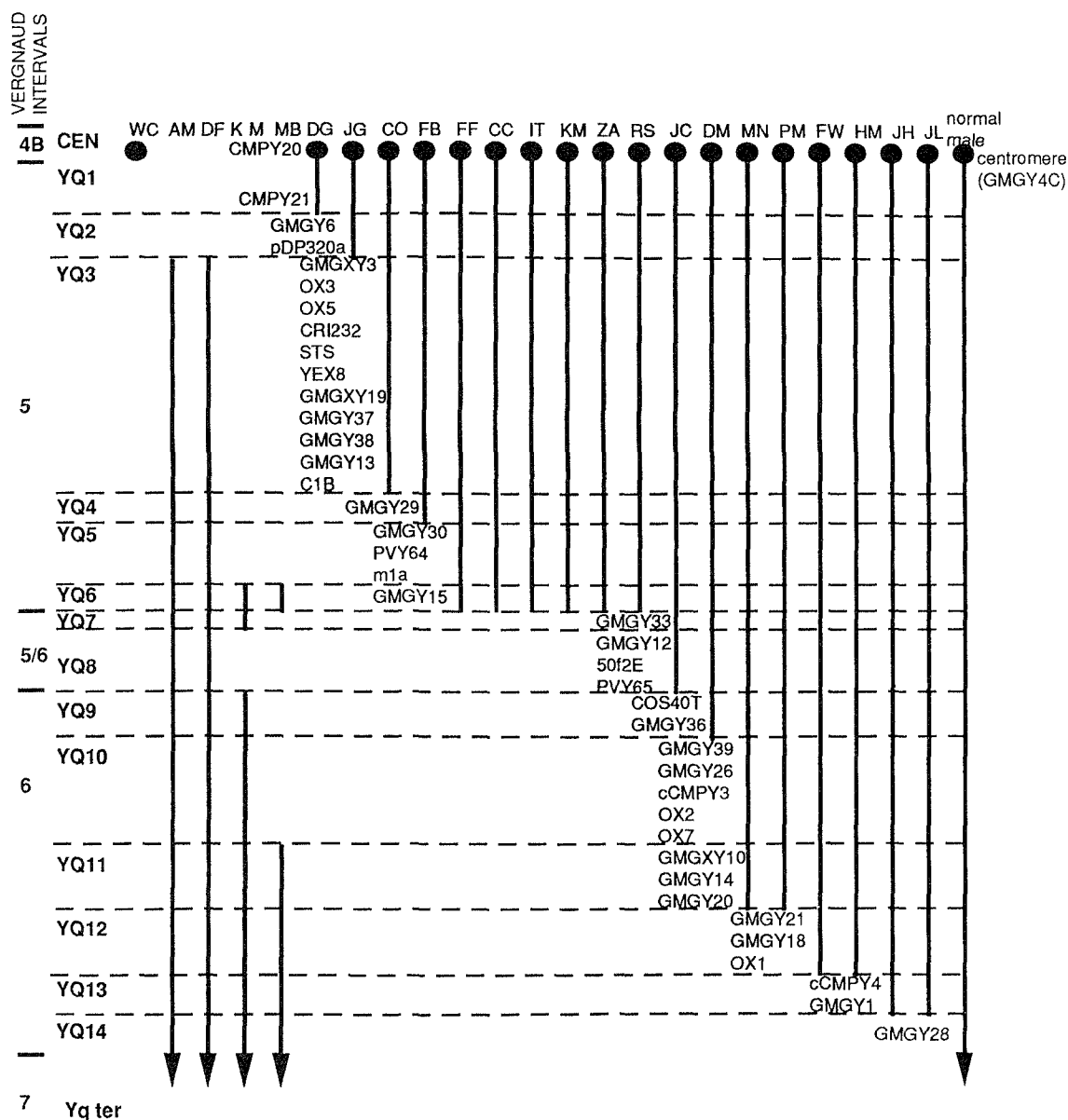


Figure 3 The figure shows a schematic deletion map of Yq derived from the data shown in table 1 and illustrates the position of probes in relation to breakpoints. The deletion intervals are numbered YQ1–YQ14 and are shown aligned against the deletion intervals of the map of Vergnaud⁴

in XX males which they suggest could be generated by pericentric inversion in the paternal Y chromosome

As mentioned above, the establishment of a deletion map of the Y long arm makes it possible to produce more precise correlations between the various mapping intervals defined by patients and any clinical phenotype caused by deletions in the euchromatic regions of Yq. Deletions involving the euchromatic segment of Yq have suggested that genes important in spermatogenesis and tooth size map to Yq11. In mouse, the *Hya* locus and *Spy*, a locus controlling spermatogenesis, have been found to be closely linked²⁴. Linkage between HYA and AZF in man is also evident^{6,22}, our patients WC (also analysed in the paper by Simpson et al⁶), JC, DM and FW localise HYA to Ycen–Yq11.21. On the basis of azoospermic patients deleted for Yq11, this region seems also to be necessary for successful

spermatogenesis in humans. Great care, however, has to be exercised in interpreting such correlations in order to exclude mosaicism or perturbation of X–Y pairing during meiosis provoked by the abnormal structure of dicentric iso Yp chromosomes as the cause of infertility. Consequently, conclusions of the clinical significance of the euchromatic region of Yq founded on karyotype–phenotype correlations must be based on Yq-chromosomes where mosaicism can be excluded.

The ambiguities introduced by mosaicism into the interpretation of other cell lines typed as negative for H–Y antigen has prevented us from defining a more precise mapping interval for the HYA locus. Nevertheless, the location we have defined for the HYA locus agrees well with that determined by Cantrell et al²⁵. Their case 5 is positive for H–Y antigen and the probe 50f2E and has a breakpoint in interval 6 of the map of Vergnaud et al⁴. This

Table 3. Summary of H-Y Antigen Typing

	HLA Serotype		% lysis with cytotoxic cells specific for				H-Y type
	A	B	A2	A2/H-Y	B7	B7/H-Y	
Exp. 1							
Control Male	2	<u>Z</u> ^a , 62	ND ^b	ND	37 ^c	80	+
Control Female	3	<u>Z</u>	ND	ND	47	1	-
DM	1,2	<u>Z</u>	ND	ND	52	72	+
Exp. 2							
Control Male	3	<u>Z</u> , 13	ND	ND	60	87	+
Control Female	3,24	<u>Z</u>	ND	ND	41	6	-
JC	3	<u>Z</u>	ND	ND	56	72	+
Exp. 3							
Control Male	1,2	7,8	64	29	ND	ND	+
Control Female	2,3	15	62	0	ND	ND	-
FW	1,2	?15	52	25	ND	ND	+
WC	2	12	85	0	ND	ND	-

Prior to H-Y antigen typing, HLA typing was done to determine whether the patients were of A2 or B7 types, since the T-cell clones available for H-Y typing were restricted by either HLA-A2 or HLA-B7. Standard HLA serotyping was performed at the tissue typing laboratory at the Royal Postgraduate Medical School (Hammersmith, London) by Mr Nick Davey and by fluorescence-activated cell sorting (FACS) analysis at the Clinical Research Centre (Harrow, UK) using the HLA-A2 specific monoclonal antibody HB82 (BB7 2) and the HLA B7 (crossreactive on B40) monoclonal antibody HB59 (MB40 2). Expression of HLA A2 and B7 alloantigens identified by T cells was confirmed by cytotoxic T-cell lysis (CTL) experiments in which the patients' cells were also typed for H-Y⁶. Epstein-Barr virus-transformed lines from each of the patient were used as target cells in CTL assays. The cytotoxicity was measured in a ⁵¹Cr release assay as described previously⁶. Control normal male and female cell lines were examined with each experiment.

^a The identity of the HLA restriction molecule used as a restriction element for the detection of H-Y antigen in the CTL assays is underlined in the HLA serotyping columns.

^b ND = not done.

^c Figures underlined in the CTL test columns are those showing significant levels of titrating lysis.

is very similar to the distal limit defined by our patient JC who is also positive for 50f2E and the H-Y antigen, where the breakpoint (by cross-referencing to the common patients typed by ourselves and Douglas Vollrath—see figure 3) has been shown to lie in interval 6. Case 4 of the Cantrell et al.²⁵ study is H-Y antigen negative and has a breakpoint between the centromere and interval 6, probably in interval 5 of the Vergnaud map.

At least five categories of sequences mapping to Yq can be distinguished and the establishment of a deletion map has been useful in indicating how these are organised. Figure 4 summarises the arrangement of these sequences on Yq as it is presently understood; (1) sequences which are specific to the Y chromosome, (2) sequences which define a block of homology with Xp22.3-pter, (3) sequences which detect homologies with Xq28, (4) sequences which define a block of homology with Yp and (5) sequences which detect homologies with autosomes. For several sequences it has not yet been determined whether the homologies are to the X or autosomes.

The block of X-Y homology in proximal Yq (homologous to Xp22.3-pter) contains the STS pseudogene (STSP) and several other closely linked sequences²⁶. It is believed to have arisen on Yq as a result of a pericentric inversion in an ancestral Y chromosome leading to the removal of this group of sequences from the X-Y pairing region²⁷. This is supported by the finding that the probe m1a maps to interval YQ5, which is distal to the YQ3 interval containing GMGXY3, STS, GMGXY19 and CRI232. On the short arm of the X there is a similar arrangement which if inverted on the ancestral Y would give rise to the order shown in figure 3. The same findings have been observed by

Number of probes in each interval for each category of homology

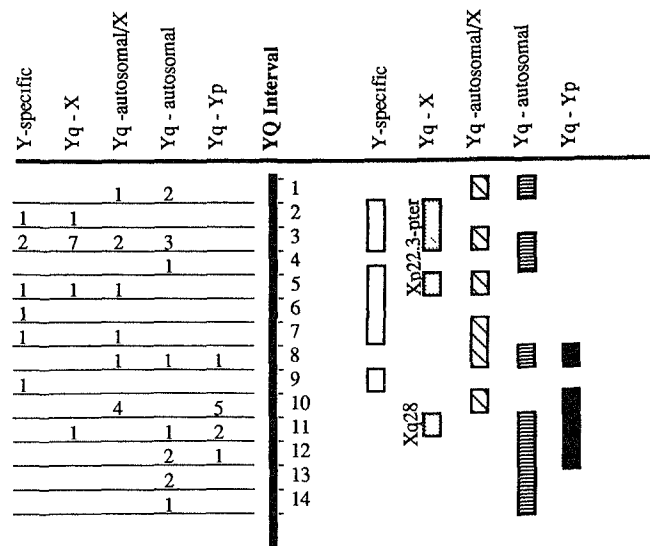


Figure 4. The figure displays schematically the arrangement of the sequences in Yq that detect X, autosomal and Yp homologies, and also sequences that are Y specific.

Bardoni et al.²⁰. Thus these sequences probably represent ancient homologies which date from the ancestral homologues which gave rise to the sex chromosomes.

In contrast, the Yq-Xq28 homology is of more recent origin and includes not only the sequence GMGXY10 but also the sequences DXYS61 and DXYS64^{28,29}. Bardoni et al.²⁰ have shown that DXYS61 and DXYS64 map to the Yq telomere whereas the GMGXY10 locus is located in the distal Yq euchromatin. DXYS61 and DXYS64 sequences are of more recent origin, appearing on the Y between chimpanzee and human, whereas GMGXY10 is older and is likely to have moved to the Y before the divergence of the old world monkeys³⁰. Duplicative transposition from the X is the most likely explanation for the appearance of these sequences on the Y³¹.

The presence of Yq-Yp homologies suggests the occurrence of duplications and intrachromosomal rearrangements (either inversions or transpositions or both) during the evolution of the Y. In this respect, several of the sequences which detect multiple loci may reflect expansion of blocks of sequence by such mechanisms.

Many of the sequences mapping to Yq with autosomal homologies may have colonised the Y by transposition or retroposition, the latter resulting in the deposition of processed pseudogenes. There are a number of examples of Y pseudogenes which include actin³², argininosuccinate synthetase³³, two anonymous transcripts described by Leroy et al.³⁴ and GMGY28 which represents a processed pseudogene derived from a chromosome 2 transcript (unpublished results). The generation of intrachromosomal rearrangements and the importation of sequences from other chromosomes may be important events in initiating and preserving a sex determining role for the Y by preventing pairing and recombination of the differential portion with the X.

MATERIALS AND METHODS

Chromosome specific libraries and cell lines

The chromosome specific libraries and cell lines WC (46,X with a monocentric iso Yp chromosome), ED (48,XX with two dicentric iso Yq chromosomes, AMIR2N (a somatic cell hybrid containing the derivative X chromosome of the following translocation t(X,Y)(p22.3,q11.1)), the Y only somatic cell hybrid 7631 and the X only somatic cell hybrid Horl X have all been described in detail elsewhere.^{1,3,11,12} Single copy DNA sequences were isolated from chromosome specific libraries¹

Patient material

Patient WC has a monocentric iso Yp chromosome. Patients JC, CO, FF, CC, IT, KM, ZA, RS, DM and FB have dicentric iso Yp chromosomes with a breakpoint in Yq11.21. Patients MN, PM, FW, HM and JH all possess dicentric iso Yp chromosomes with a breakpoint in Yq11.22 (see figure 2). Patients AM (from which the hybrid AMIR2N is derived), DF and K M possess the derivative X of X Y translocations (t(X,Y)(p22.3,q11.21)) and patients JG and MB the respective derivatives of a Y 5 (t(Y,5)(q11.21,p13.3)) and a Y 14 (t(Y,14)(q11.22,q32.2) translocations. Patient DG has a ring Y (46,XrY) and patient JL is 46,XYq— with loss of the distal heterochromatic region from Yq11.23. Patients JC, CO, CC, RS, MN, JG, and JL were obtained from Ellen Magenis. K M was obtained from Nick Dennis. FB was obtained from John Pearson. MB was obtained from Diane Curtis.

Preparation of genomic DNA and Southern blot analysis

Preparation of high molecular weight genomic DNA from peripheral blood lymphocytes or lymphoblastoid cell lines, restriction digests and Southern transfers were carried out as described by Maniatis et al.¹³ DNA probes were labelled by oligonucleotide random priming as described by Feinberg and Vogelstein¹⁴ and molecular hybridization under the conditions described by Wahl et al.¹⁵

Determination of H-Y status

Lymphoblastoid cell lines were typed for H-Y antigen using the cytotoxic T cell assay described by Simpson et al.⁶

DNA probes

The probe pDP320a (DYS148) was provided by David Page and has been described by Fisher et al.¹⁶ Probes YEX8, PVY64 and PVY65 were obtained from Peter Vogt. Probes OX1 (DYS27), OX2 (DYS26), OX3 (DYS28), OX5 (DYS30) and OX7 (DYS33) were obtained from Ian Craig. Probe CRI 232 (DXS278) was obtained from Collaborative Research. The steroid sulphatase (STS) probe was obtained from Andreas Ballabio. Probe M1A (DXS31) was obtained from J L Mandel. Probe 50f2 (DYS7) was obtained from Jean Weissenbach. Probe C1B is a cosmid fragment obtained from Kay Taylor. Probe cos40T is a cosmid fragment obtained from Chris Lau. Probes Y46.1 (cCMPY4) and 14.1/IB (cCMPY3) are cosmid fragments obtained from Carole Sargent. Probes GMGY1 (DYS12), GMGY6 (DYS66), GMGY4C (DYS52), GMGY12 (DYS64), GMGY13 (DYS63), GMGY14 (DYS118), GMGY15 (DYS62), GMGY18 (DYS75), GMGY20 (DYS73), GMGY21 (DYS72), GMGY26 (DYS77), GMGY28 (DYS79), GMGY29 (DYS80), GMGY30 (DYS84), GMGY33, GMGY36 (DYS88), GMGY38 (DYS86), GMGY39 (DYS89), GMGX3 (DXY29Y), GMGX19 (DYS74), GMGX10 (DXY37Y), and CMPY21 were isolated from a flow-sorted Y lambda library.¹⁷ Probe 691 # 10 (CMPY20) was isolated from a foetal brain cDNA library (Stratagene 937201).

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