

# JOURNAL OF MEDICAL GENETICS

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Papers should conform to one of the following categories. *Original contributions* on clinical or laboratory aspects of medical genetics in man and on related animal studies. *Case reports* with particularly instructive clinical or genetic features: to be not longer than 1000 words of text, two or at most three figures, one table (if necessary), and eight references. *Short reports* of unusual cases: to be not longer than 500 words or point form description with a clinical photograph and partial karyotype, if appropriate, and no more than two or three reference. Single case reports will usually only be considered in one of these forms. *Review articles* will generally be by invitation, but suggestions from authors wishing to prepare a review article will be welcome. *Annotations, Hypotheses, Preliminary communications, and Technical notes* will also be considered, as will *Short communications* giving information on new translocations, chromosome identification by banding techniques, and second and third findings of important haemoglobins. Contributions to the *Correspondence* and *Question and answer* columns will be welcomed.

All contributions should be accompanied by a summary giving the main results and conclusions. Typescripts should be double spaced with wide margins. One page proof will be sent to the author submitting the paper and alterations on the proof, apart from printer's errors, are not permitted. Twenty-five free reprints will be supplied and further reprints may be ordered when the proof is returned.

*Figures* should be kept to a minimum and should be numbered consecutively in Arabic numerals. Legends should be typed on a separate sheet. Photographs should be on glossy paper and diagrams should be drawn on stout white paper. Photographs of karyotypes do not reproduce well. Chromosomes should be cut out and stuck onto stout paper. Any lettering should be indicated on a separate transparent overlay. Pedigrees should use squares and circles. Generations should be numbered with Roman and individuals with Arabic numerals; members belonging to the same generation should be horizontally aligned.

*Tables* should not be included in the body of the text, but should be typed on separate pages and numbered with Arabic numerals.

*References* should conform precisely to the style current in this Journal. Authors are responsible for the *accuracy* and *completeness* of their references as these will not be checked by the Editorial Office.

*Nomenclature.* Authors should refer to the following publications.

(1) Chromosomes: ISCN. An international system for human cytogenetic nomenclature (1978). *Birth Defects* 1978; XIV:No 8. Also in *Cytogenet Cell Genet* 1978;21: 309-404.

(2) Dermatoglyphs: Penrose L S. Memorandum on dermatoglyphic nomenclature. *Birth Defects* 1968;4:No 3.

(3) Enzymes: WHO Scientific Group. Standardization of procedures for the study of glucose-6-phosphate dehydrogenase. *WHO Tech Rep Ser* 1967;No 366.

(4) Blood coagulation: International Committee of Haemostasis and Thrombosis (Graham J B *et al*). A genetic nomenclature for human blood coagulation. *Thromb Haemostas* 1973;30:2-11.

(5) Loci: Conventional nomenclature should be used, with lower case lettering as appropriate (for example, Race RR, Sanger R. *Blood groups in man*. 6th ed. Oxford, London: Blackwell, 1975; and Giblett ER. *Genetic markers in human blood*. Oxford, London: Blackwell, 1969).

*SI units.* The units in which the authors' work was measured should be cited first followed by either the SI units or the traditional units. This does not apply to Tables, but here a conversion factor should be added as a footnote.

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# Announcements

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*Journal of Medical Genetics*, 1982 **19**, 80

## CANCER FAMILY STUDY GROUP

Following a meeting chaired by Professor David Harnden at the Department of Cancer Studies in Birmingham, it was decided to hold regular meetings and exchange information on all aspects of familial cancer. Besides those present at the meeting, it was felt that others might be interested in the problem of cancer in a familial setting in both children and adults. It is hoped that this group will facilitate the study of these families by biological scientists, geneticists, and clinicians.

Anyone who would like further details please contact the Secretary, Dr R A Cartwright, Yorkshire Regional Cancer Organisation, Cookridge Hospital, Leeds LS16 6QB, West Yorkshire.

## HUMAN CYTOGENETIC NOMENCLATURE

At an open meeting of cytogeneticists held during the 6th International Congress of Human Genetics in Jerusalem, a new Standing Committee for Human Cytogenetic Nomenclature was elected to serve for a period of 5 years. The members are Martin Bobrow (Europe), Maimon Cohen (USA and Canada), Uta Francke (USA and Canada), David Harnden (Europe), Patricia Jacobs (USA and Canada), Grant Sutherland (Asia and Australasia), and Jacob Wahrman (Middle East and Africa). The committee would like to receive comments on ISCN (1978) and ISCN (1981) to help with future revisions. Contact Professor David Harnden (Chairman), Department of Cancer Studies, University of Birmingham, Birmingham B15 2TJ, UK.