CHROMOSOME BANDS INDUCED IN HUMAN AND SYRIAN HAMSTER CELLS BY CHEMICAL CARCINOGENS

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Received 3 April 1974. Accepted 25 April 1974

Summary.—After 24 hours of treatment of human peripheral leucocyte cultures or Syrian hamster embryonic secondary cultures with some known chemical carcinogens at concentrations that produce transformation in vitro of hamster cells, the chromosomes of both types of cells exhibited typical G banding. Chromosome bands did not result with urethane, a compound which does not produce cytotoxicity or cause transformation by direct exposure of cells. Non-carcinogenic chemicals which do not inhibit cell multiplication also failed to produce bands. Cytogenetic analysis following removal of the carcinogens from the cultures indicated non-banded uniformly stained chromosomes. It is concluded that metaphases with chromosome bands are more likely to be the result of nonspecific toxicity rather than being related to the carcinogenic properties of the chemicals.

During the past 3 years a number of techniques have been developed for distinguishing between morphologically similar chromosomes according to the banding pattern which formed after special treatments of fixed chromosomes (Caspersson, Johnanson and Modest, 1970; Sumner, Evans and Buckland, 1971; Drets and Shaw, 1971; Seabright, 1971). The bands, generally referred to as G or Q, represent specific regions of the chromosomes that become swollen and show transverse ridges which take up and bind dyes. Recently, it has been reported that the addition of some chemicals to living cells results in identical chromosomal bands without the requirement for a special post-fixation treatment. Treatment of cells with actinomycin D (Hsu, Pathak and Shafer, 1973) or tetracycline (Meisner, Chuprevich and Inhorn, 1973) during the G2 phase of the cell cycle resulted in G bands, whereas ethidium bromide or nogalamycin (Hsu, Pathak and Shafer, 1973) apparently produced bands the reverse of the G type and referred to as R bands. Hydroxyurea produced the typical chromosomal cross bands when the cells were treated during the DNA synthesis or G2 periods (Popescu and DiPaolo, 1974). Thus, diverse chemicals can induce chromosomal banding in fixed preparations or in pretreated living cells. We now report that some chemical carcinogens produce chromosome bands in human and hamster cells in culture and that non-carcinogens do not.

MATERIALS AND METHODS

Seven known chemical carcinogens were tested: 7,12-dimethylbenz(a)-anthracene (DMBA), benzo(a)pyrene (BP), aflatoxin B1 (AFB1), n-acetoxy-2-fluorenylacetamide (N-acetoxy-FAA), N-methyl-N'-nitro-N-nitrosoguanidine (MNG), 4-nitroquinoline-1-oxide (4NQO) and ethylecarbamate (urethane). Three non-carcinogens tested were: pyrene (Py), phenanthrene (Ph) and perylene (Pe).

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BP, DMBA and urethane were purchased from Eastman Kodak Co.; Ph, Py and Pe from K & K Laboratories; N-acetoxy-FAA from Starks Assoc., Inc.; AFB from Calbiochem and MNNG from Aldrich Chemical Co. The chemicals were dissolved in acetone (10 mg/ml) and diluted with warm complete medium to make stock solutions. Further dilutions were made with fresh complete medium to obtain the concentrations needed. Final concentrations of carcinogens used were those which, with the exception of urethane, produced neoplastic transformation of hamster cells in culture (DiPaolo, Donovan and Nelson, 1969; DiPaolo, Nelson and Donovan, 1972).

Secondary passages of Syrian hamster embryo cultures derived from whole foetuses (DiPaolo et al., 1969) and human peripheral leucocyte cultures obtained from venous blood were used. Blood was harvested with heparin (1000 i.u./ml, Upjohn Co.) and the white cells from 25–30 ml of blood collected by sedimenting for 1.5 h at room temperature. Plasma was centrifuged for 5 min at 800 rev/min in an International clinical centrifuge and resuspended at 1 × 10^6 cells/8 ml 1640 medium supplemented with 15% foetal bovine serum. Phytohaemagglutinin P (0.01 ml/culture) was added and cells were incubated at 37°C and 5% CO₂ for 72 h.

Hamster cells in log growth were treated with chemicals for 4, 8 or 24 h. Human leucocytes were also treated for the same time intervals before terminating the culture at 72 h. After completion of chemical treatment, Colemide at a concentration of 0.02 and 0.04 μg/ml medium for human and hamster cells respectively, was added for 3 h. Subsequently, the cells were collected (the hamster cells were first detached using a rubber policeman) by centrifuging the cultures at 800 rev/min for 5 min, resuspended in 0.075 mol/l KCl (human cells) or in a solution of one part complete medium and 2 parts distilled water (hamster cells), and incubated for 10–12 min at 37°C. After the hypotonic treatment, the cells were fixed 3 times in 3:1 (methanol-acetic acid). Slides were made by the air dry method and chromosomes were stained with Giemsa (1 ml Gurr's Giemsa R66 to 50 ml phosphate buffer at pH 6.9–7.0) for 3–5 min, washed with distilled water, dried and mounted in permount. Two hundred metaphases were examined. Only those chromosomes with well-defined dark bands separated by negative zones were considered as banded metaphases. Each chemical was tested in at least 4 separate experiments but the data presented are limited to 1 experiment using the cells from one source.

**RESULTS**

With the concentrations used, all carcinogens except for urethane produced bands in chromosomes of living human and hamster cells; non-carcinogenic chemicals were ineffective in producing bands (Table). The cross bands occurred only after 24 h of treatment, while 4 or 8 h of treatment with the same concentrations failed to produce chromosome bands. The incidence of metaphases with chromosomes exhibiting bands varied with the carcinogen and in general neither human leucocytes nor Syrian hamster cells were more susceptible although the highest incidence of banded metaphases (80%) occurred when DMBA was applied to human leucocytes.

An effect of the carcinogens in both human and hamster cells was a partial inhibition of the cell multiplication, except for urethane. Non-carcinogens had little or no effect on cell division. For example, the average number of Syrian hamster cells in control dishes (50 mm

**Table—Percent of Metaphases Exhibiting Chromosome G Bands after Treatment with Various Chemicals**

| Chemical     | Human leucocytes | Syrian hamster fibroblasts |
|--------------|------------------|---------------------------|
| BP (1·0)     | 65               | 50                        |
| DMBA (0·1)   | 80               | 60                        |
| MNNG (0·1)   | 20               | 30                        |
| 4NQO (0·01)  | 28               | 35                        |
| AFB (1·0)    | 65               | 30                        |
| N-Acetoxy-FAA (2·5) | 30 | 20                        |
| Urethane (50·0) | 0             | 0                         |
| Py (10·0)    | 0                | 0                         |
| Ph (20·0)    | 0                | 0                         |
| Pe (10·0)    | 0                | 0                         |
| Control      | 0                | 0                         |
FIG. 1a.—G bands of human chromosomes produced by post-fixation trypsin treatment.

FIG. 1b.—G bands of chromosomes from human leucocyte cultured cells produced by prefixation with BP treatment (1 μg/ml for 24 h).
Fig. 1c.—Syrian hamster chromosomes with fine bands after N-acetoxy-AAF prefixation treatment (5 µg/ml for 24 h). Metaphases with this type of band also occurred after non-carcinogen treatment.

Fig. 1d.—Syrian hamster banded metaphase with multiple structural aberrations (arrows) after 24 h AFB1 treatment (1 µg/ml) before fixation.
diameter) doubled during a 24 h period (4.7 × 10^4 to 9 × 10^4), urethane treated cells exceeded this number (12.3 × 10^4) and compounds such as AFB_1 and N-acetoxy-AAF reduced the average cell number (6 × 10^4). Non-carcinogens did not influence cell number. Bands occurred after carcinogen treatment of living cells, even in metaphases surrounded by cytoplasm or in those with pronounced contraction. Although the bands obtained with each carcinogen were not analysed band by band, it is apparent that the bands have a configuration similar to the G type (Fig. 1a, b). Nevertheless, it is possible that subtle variations may have occurred with different agents.

Chromosomes prepared from non-treated human lymphocyte cultures or chemically treated human or hamster cultures by the above technique and stained in phosphate buffered Giemsa for 3.0–3.5 min occasionally had entire metaphases with a faint banding pattern of a fuzzy nature.

In one experiment hamster cells were treated for 24 h with 1 µg of BP/ml medium. The carcinogen was removed and cells allowed to grow for an additional 24 h; when chromosomes were prepared, no banded metaphases were observed. Another type of chromosome band associated with either carcinogen or non-carcinogen and human cells can be described as a series of multiple bands without a distinguishable pattern that in general resembles a coiling chromosome structure (Fig. 1c). This phenomenon was observed following either carcinogen, including urethane, or non-carcinogen treatment. The highest incidence (20%) was following MNNG and the lowest incidence, (2%) was with DMBA, urethane and Ph. The types of chromosomal damage produced by the chemical carcinogens consisted primarily of chromatid breaks and chromosome exchanges (Fig. 1d). The incidence of the chromosomal aberrations was dependent upon concentration and length of exposure of the cells to the carcinogens.

**DISCUSSION**

The banding techniques make possible the accurate identification of individual chromosomes. The techniques have been applied to cancer cytogenetics to determine whether a causal relationship exists between chromosome alterations and malignant transformation (DiPaolo, Popescu and Nelson, 1973; Yamamoto, Rabinowitz and Sachs, 1973), and to establish the origin of chromosomal rearrangements such as chronic granulocytic leukaemia (Rowley, 1973).

Efforts are now being directed towards elucidating the mechanisms of banding formation. Recent studies indicate that the interaction of dye, either quinacrine mustard or Giemsa with DNA, as well as the relationship between DNA and its associated non-histone proteins play an important role in banding formation (Comings et al., 1973; Sumner and Evans, 1973). The significance of the fine type of bands is not clear; however, this type of structure may be the basic configuration upon which the G bands are formed. Although the mechanism of banding in living cells and in fixed preparations may differ, some common feature probably exists since identical patterns result. In fact, fixed chromosome preparations immersed for 1–5 min in a solution of 1 µg/ml of MNNG in Hanks' medium without Ca^{++} and Mg^{++} and stained with buffered Giemsa also resulted in G bands (unpublished data).

The list of diverse compounds capable of producing chromosome bands now includes chemical carcinogens. It is possible that chromosome bands associated with the carcinogens are a reflection of a nonspecific type of chemical toxicity which leads to cell death because of DNA or associated protein changes that cause conformational changes of nucleoprotein (Meisner et al., 1973). Chromosomal banding may be an immediate response to carcinogen treatment because when the cells were grown for an additional 24 h in medium without carcinogen, no banded metaphases were found. With the cells
used in this experiment, partial inhibition of cell multiplication that increased with time occurred after treatment with all carcinogens that caused chromosome bands whereas the non-carcinogens and urethane, neither of which had any effect on cell multiplication, failed to produce bands. Furthermore, urethane treatment actually increased the number of mitoses. In other experiments, the addition of urethane to Syrian hamster cells seeded to form colonies did not result in any transformation or affect colony forming ability (DiPaolo et al., 1972).

Previous studies have shown that the frequency of transformation, a step in the process leading to neoplasia, continues to increase with increasing concentration of chemical carcinogen even after all cells sensitive to the toxic effect of the compound have been killed (DiPaolo et al., 1971a). In fact, if the cells are protected from the toxicity of the compounds used in the study, the transformation frequency may even increase (DiPaolo et al., 1971a, b). Examination of chromosomes of cells transformed by chemical carcinogens reveals that the changes in number of chromosomes or new marker chromosomes, as indicated by banding patterns, are not causally related to the process of carcinogenesis but are reflections of secondary alterations (DiPaolo et al., 1973)

Depending upon the carcinogen concentration used, chromosomal breakage may reflect cell lethality (DiPaolo and Popescu, unpublished data). In the present experiments, breaks and rearrangements occurred following carcinogen treatment which inhibited cell multiplication. The chromosome bands obtained using chemical carcinogens are similar to those reported for compounds such as hydroxyurea and azure B not suspected to be in vivo carcinogens. Furthermore, metaphases with chromosome crossbands were not observed with urethane, a well known in vivo carcinogen. Therefore, the manifestations reported here probably are not related to transformation, nor can they be expected to serve as an accurate index for surveying chemical compounds for possible carcinogenicity.

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