Establishment and characterisation of a new tumorigenic cell line with a normal karyotype derived from a human breast adenocarcinoma

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Summary A new cell line (CAL51) was isolated from a malignant pleural effusion of a woman with metastatic breast cancer. These cells grow in continuous culture and exhibit the morphological, ultrastructural and immunohistochemical features of epithelial cells of mammary origin. They are tumorigenic in nude mice and clone in soft agar. Oestrogen receptors are not detected. CAL51 consists of a homogeneous population of cells with normal chromosomes even after the use of high resolution banding. Cytogenetic analysis of the cells from the tumour induced by CAL51 in the nude mouse confirmed the normality and the stability of the karyotype. All breast cancer cell lines established to date present abnormal karyotypes; CAL51 cell line may be more informative than cell lines with aberrant karyotypes for investigating essential genetic differences between normal and malignant mammary gland cells.

Breast adenocarcinomas are among the most frequent cancers in women. These generally slow-growing tumours present a wide spectrum of clinical courses that sometimes defy all attempts at prognosis and treatment. In vitro models are required for the study of these cancers, and several cell lines have already been established and characterised (Lafargues & Ozzello, 1958; Soule et al., 1973; Cailleau et al., 1974; Engel et al., 1978; Whitehead et al., 1983; Yamane et al., 1984; Chu et al., 1985; Vandewalle et al., 1987). Nevertheless, owing to the heterogeneity and the diversity of mammary cancers, a great number of cell models is necessary to understand the reasons for this diversity and the effect of anticancer drugs on tumour cells. Cytogenetic studies of mammary adenocarcinoma cell lines are essential for comprehension of the pathogenesis of these cancers (Trent, 1985; Gebhart et al., 1986). The implication of chromosomal alterations in these pathologies has opened a new and promising route towards better knowledge of these cancers (Cervenka & Koulischer, 1973). Chromosomal alterations are generally numerous, and markers often demonstrate hyperploidy in these cancers (Sandberg, 1980).

Demonstration of the minimum genetic alterations indispensable for cell transformation is difficult, and might be easier on cells with a karyotype closer to normal. Sandberg and Wolman mentioned the existence of such cells, but most of their results concerned karyotype studies without chromosome banding (Sandberg, 1980; Wolman, 1983).

CAL51, corresponding to a cell population with a 'normal' karyotype, is a new mammary adenocarcinoma cell line derived from the malignant pleural effusion of a patient treated at the Antoine-Lacassagne Cancer Center (Nice, France).

Materials and methods

Patient and cell culture

This 44-year-old woman was seen in January 1984 for progressive breast cancer. After chemotherapy, she underwent bilateral mastectomy with resection of positive axillary nodes; histology diagnosed invasive adenocarcinoma with extensive intraductal involvement. Oestrogen and progesterone receptors were both negative. After radiotherapy and chemotherapy, she presented in August 1985 with cutaneous, osseous and hepatic metastases and bilateral pleural effusion. On 4 October 1985, thoracentesis produced numerous clumps of cancerous cells that were cultivated.

The cells were maintained in Dulbecco's modified minimum essential medium with Earle's salts (D MEM) (Boehringer, France SA) supplemented with 0.6 μg ml⁻¹ bovine insulin, 5 × 10⁻¹⁰⁵ ml⁻¹ transferrin, 2 mmol L-glutamine, 10% fetal calf serum, 400 U ml⁻¹ penicillin, and 200 μg ml⁻¹ streptomycin. Cells were stored in an incubator at 37°C in a 5% CO₂ atmosphere.

Doubling time

At passages 15 and 50, 12 × 10⁶ cells were plated in Falcon plastic flasks (25 cm²) containing complete culture medium. Cells were harvested with a solution of 0.05% trypsin in phosphate-buffered saline (PBS) and were counted every day for 11 days starting 24 h after plating.

Electron microscopy

The cell layer was scraped with a rubber policeman to obtain groups of cells. After centrifugation, the residue was fixed twice with 1.25% cold glutaraldehyde and 2% osmic acid at room temperature for 30 min. The cells were then dehydrated by rinses of increasing concentrations of alcohol and propylene oxide. The residue was embedded in Epon. Sections were prepared with an LKB Nova ultramicrotome; grids were stained by the LKB Ultrastainer and the sections were examined with a Philips CM 12 (20 keV) electron microscope.

Chromosomal analysis

Cytogenetic examination of CAL51 cells was performed at around passage 13. Cells in the exponential phase of growth were treated with colchicine (Seromed, Biopro) for 1/2 h at a final concentration of 1 μg ml⁻¹. Cells were then trypsinised, washed and treated for 45 min with a hypotonic solution (0.075 M KCl, EGTA 0.2 g l⁻¹, hyaluronidase 25 IU for 20 ml, Hepes buffer 4.8 g l⁻¹ in distilled water). Cells were then fixed in acetic acid:methanol (1:3) and dropped onto grease-free, cooled slides for chromosome counting and examination. B bands were obtained by heat denaturation of the chromosomes according to the method of Dutrillaux and Lejeune (1971). Xenografted CAL51 cells were plated and studied in vitro in the same manner.

Further studies at passage 17 for CAL51 and at passage 9 for xenografted CAL51 cells using thymidine synchronisation
and BrdU incorporation (Viegas-Pequignot & Dutrillaux, 1978) were developed with some differences since an incubation of 18–20 h with a final concentration of 10 μg of BrdU per ml was required to study replication banding pattern and high resolution banding. ISCN T, C and Q-banding (ISCN, 1985) were applied on some preparations.

**Immunological studies**

 Peroxidase labelling was performed to reveal the oestrogen receptors and the epithelial nature of the cells. CAL51 cells were cultured on glass slides for both experiments, then fixed. After indirect immunocytochemistry (peroxidase antiperoxidase), cells were stained with Harris’ haematoxylin.

**Oestrogen receptors** Cells were fixed with 3.7% formaldehyde in PBS, then with pure methanol and acetone. The specific monoclonal antibody used was H222SP gamma from the ER-ICA kit (Abbott).

**Epithelial nature of the cells** Cells were fixed with acetone. An antikeratin monoclonal antibody (KLI-Immunotech) diluted to 1:50 and an anti-epithelial membrane antigen (DAKO, EMA) diluted to 1:25 were used. The same experiments were performed on paraffin sections of the primary tumour of this patient and on a section of a tumour induced in the nude mouse by CAL51.

**Clonogenic growth in soft agar**

Cells were cultured as described by Salmon et al. (1978). In brief, cells were suspended on 0.3% agar in enriched Connaught Medical Research Laboratories Medium 1066 (Grand Island Biological Co., Grand Island, NY, USA) with 15% horse serum (Flow Laboratories, Puteaux, France) in 35 mm Petri dishes containing an underlayer of 0.5% agarose in culture medium. Cells were then incubated at 37°C in a humidified atmosphere of 5% CO₂ and 95% air. Plates were examined with an inverted phase microscope 21 days after plating. Positive controls were obtained in the same conditions using melanoma cell line CAL1, which clones in soft agar (Coudri et al., 1983).

**Tumorigenicity**

Six-week-old athymic nude mice with the Swiss genetic background (IFCA Credo Laboratories, France) were used. Some 2 × 10⁶ cells in 0.2 ml Ringer lactate solution were inoculated s.c. into both flanks of three mice.

**Results**

**Morphology by light microscopy**

After culturing, epithelial cells appeared rapidly. Figure 1 shows the CAL51 cells as they appeared under light microscopy. These heterogeneous cells (small, polyhedral cells or larger, more rounded cells) showed anchorage-independent growth. They presented visible nuclear abnormalities. The first passage was performed one month after initiation of culture. Cells were maintained in culture until passage 40, then conserved in liquid nitrogen.

**Doubling time**

CAL51 cells grow exponentially up until day 8, when they reach a plateau phase. The population doubling time was measured during the period of exponential growth: 45 h at passages 15 and 50.

**Electron microscopy**

CAL51 cells all have a large nucleus with irregular contours and a large nucleolus. The cytoplasm contains abundant ribosomes (free or organised in ergastoplasmic cisternae),

![Figure 2](image)

**Figure 2** Electron micrograph of cell line CAL51: a, zonula adhaerens (ZO) (x 17,000); b, microfilaments (MF) (x 17,000).

![Figure 1](image)

**Figure 1** Photomicrograph of cell line CAL51 (x 70).
mitochondria, rather numerous Golgi bodies and some lysosomes. The cells are often contiguous and have zonula-adhaerens junctions (Figure 2a). Certain cells contain numerous bundles of microfilaments organised around electron-free cavities (Figure 2b). Their apical poles are covered with microvilli suggesting glandular luminae.

Chromosomal analysis

T, R, Q and C-banding  Cytogenetic analysis of cell line CAL51 revealed normal diploidy; the average number of chromosomes per cell analysed was 46 (Figure 3). The karyotype was normal: no structural abnormality was seen by R-banding. Cytogenetic analysis of xenografted CAL51 cells plates in vitro gave the same results. Some metaphases were incomplete, and others carried one or two abnormal chromosomes. Since these anomalies were not recurrent, it is very likely that they were generated from cells with normal karyotype at late passages. The tetraploid metaphases or prometaphases had no additional anomalies. R and T-banding (Dutrillaux & Couturier, 1981), allowing an identification of the polymorphism which may exist on the short arms of the acrocentrics, clearly showed that the parental chromosomes were present for pairs 13, 14, 15, 21 and 22 (Figure 4). Q-bands confirmed this interpretation and exhibited a polymorphism for chromosome 3. Finally, chromosomes 9 could be differentiated by their heterochromatin.

![Figure 3](image3.jpg)  
Figure 3  Chromosome number distribution in a sample of 100 CAL51 cells.

![Figure 4](image4.jpg)  
Figure 4  Conservation of heteromorphism of heterochromatic regions. First row, Q-banding; second row, Giemsa staining of the same chromosomes; third row, R-banding of acrocentric chromosomes; fourth row, Q-banding of chromosomes 3 and R-banding after heat denaturation or Brd U incorporation (right) of chromosomes 9.
High resolution banding Apart from the fact that a high proportion of tetraploidy was found in CAL51 cells and in xenografted cells as well, no recurrent abnormalities could be observed in the 35 established karyotypes. After synchronisation and BrdU incorporation, a normal replication pattern was observed (Figure 5). In some prometaphases, a high number of bands per cell could be analysed, and yet, no anomalies could be detected. The only abnormal parameter was that a 18–20 h treatment by BrdU was necessary to obtain a replication banding pattern equivalent to a 7 h treatment in lymphocytes. This indicates a very low cell cycle, specially for the late S and G2-phases.

Immunohistochemical studies

Oestrogen receptors (ER) The results are negative. This is not surprising because the patient's primary tumour was ER negative.

Epithelial nature of the cells Reactivities with antibodies KL1 and anti-EMA were strongly positive (Figure 6a). The same was true for the section of the original tumour and the section of the tumour induced by CAL51 in a nude mouse (Figure 6b).

Cloning in soft agar
CAL51 cells had a cloning efficiency of 5.9%.

Tumorigenicity
Ten days after inoculation of CAL51 cells, four tumours appeared at six injection sites. The xenografts were excised after 1.5 months and put back in culture. The cells obtained were identical to those of cell line CAL51. Histological examination of the grafted tumours diagnosed differentiated adenocarcinoma (Figure 7a). A histologic section of this patient’s tumour is shown in Figure 7b for comparison.

Discussion
Characterisation of CAL51 cells revealed that these epithelial cells are unquestionably of mammary origin. They are tumorigenic in nude mice and clone in soft agar. Because their phenotype is stable in culture and their doubling time is suitable for kinetics studies, they are a valuable model system for the study of mammary adenocarcinoma. Their main originality is their normal karyotype. CAL51 consists of a population of diploid cells with normal chromosomes after metaphase and high resolution bandings. Since a proportion of metaphases exhibited non-clonal anomalies, such as chromosome gains or losses, or structural rearrangements, it is likely that a certain chromosomal instability exists. In some cultures, these cells undergo endoreduplication without additional anomalies. At least for acrocentrics and chromo-
somes 1 and 9 where it could be investigated no loss of heterozygosity was detected, even for chromosome 13 for which losses frequently occur (Gerbault-Seureau et al., 1987; Lundberg et al., 1987). The phenotype of these cells and their high tumorigenicity indicate that they are not normal contaminating cells, which could have been present in the pleural effusion and which can be maintained in vitro. Cytogenetic analysis of the cells from the tumour induced by CAL51 in the nude mouse confirmed the 'normality' and the stability of the karyotype. This fact is not exceptional, and has already been observed by other authors. In his review of human breast cytogenetics, based principally on unbanding cases, Sandberg (1980) stated that diploid metaphases are occasionally observed. Trent (1985) also cited the work by Wolman (1983), who found karyotypically normal diploid cells in primary mammary adenocarcinomas but not in metastatic effusions.

These authors all concluded that acquisition of an abnormal karyotype is indicative of malignant progression rather than initial events in tumorigenesis. As mentioned by Trent (1985) and Smith et al. (1985), it could be questioned whether diploid cells are really representative of the tumoral population. In culture since October 1985, cell line CAL51 has conserved a diploid karyotype with normal chromosomes. They still produce tumours in nude mice, which is an argument to consider that they represent the original cancer.

The genetic alterations responsible for the transformation into cancerous cells should exist in the genome of CAL51 cells, but they are not visible on the chromosomes, even after high resolution banding. Molecular studies might demonstrate these alterations, which may be minimal, at difference with most established cancer cell lines in which, the chromosomal damage is so extensive that the most characteristic aberrations are difficult to single out. CAL51 is an ideal model for investigating essential genetic alterations involved in malignant transformation of mammary gland cells.

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Figure 6 Immunoperoxidase staining of CAL51 with antikeratin antibodies: a, culture on glass slides (×700); b, section of xenografted tumour (×175).

Figure 7 a, Grafted tumour (differentiated adenocarcinoma) (×175). b, Primary tumour of the patient (poorly differentiated carcinoma) (×175).
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