Cell cycle synchronization and BrdU incorporation as a tool to study the possible selective elimination of ErbB1 gene in the micronuclei in A549 cells

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Abstract

Lung cancer often exhibits molecular changes, such as the overexpression of the ErbB1 gene that encodes epidermal growth factor receptor (EGFR). ErbB1 amplification and mutation are associated with tumor aggressiveness and low response to therapy. The aim of the present study was to design a schedule to synchronize the cell cycle of A549 cell line (a non-small cell lung cancer) and to analyze the possible association between the micronuclei (MNs) and the extrusion of ErbB1 gene extra-copies. After double blocking, by the process of fetal bovine serum deprivation and vincristine treatment, MNs formation was monitored with 5-bromo-2-deoxyuridine (BrdU) incorporation, which is an S-phase marker. Statistical analyses allowed us to infer that MNs may arise both in mitosis as well as in interphase. The MNs were able to replicate their DNA and this process seemed to be non-synchronous with the main cell nuclei. The presence of ErbB1 gene in the MNs was evaluated by fluorescent in situ hybridization (FISH). ErbB1 sequences were detected in the MNs, but a relation between the MNs formation and extrusion of amplified ErbB1 could not be established. The present study sought to elucidate the meaning of MNs formation and its association with the elimination of oncogenes or other amplified sequences from the tumor cells.

Key words: A549 cells; Cell cycle synchronization; BrdU incorporation; ErbB1; Micronucleus

Introduction

Gene amplification in a tumor has been associated with poor prognosis and chemotherapy resistance. Cytogenetically, gene amplification can be observed in two different structures, namely, double-minute (DM) and homogeneously staining region (HSR). DM is a small extrachromosomal structure that is paired, acentric, and atelomeric and can be formed by fragments of chromosomes, chromatin particles, and oncogenes. HSR is a region in any chromosome that fails to display the typical banding patterns after trypsin-Giemsa staining (1,2). Oncogene and multi-drug resistance genes are often overexpressed due to their amplification (3). ErbB1 (encoding epidermal growth factor receptor [EGFR] protein) gene amplification, for example, has been found predominantly in cases of lung cancer, as well as in cases of gliomas, breast, and ovarian cancers. Some features like proliferation, survival, induction of angiogenesis, invasion, metastasis, and activation of transcription factors in the tumor cells are because of the amplified ErbB1 (4).

Amplified genes in DM have been detected inside micronuclei (MNs). One example includes the selective elimination of c-myc through MNs by HL-60 (promyelocytic leukemia) and COLO 320 (colon carcinoma) cell lines that leads to the reduction in tumorigenicity (5). HL-60 cells can also be differentiated by eliminating c-myc amplification through MNs (6). Moreover, the neuroblastoma cell lines spontaneously eliminated amplified c-myc gene by MNs formation, leading to a loss of the malignant phenotype (7). Another example is the relationship between the selective elimination of cyclin-dependent kinase 4 (CDK4) sequences and liposarcoma differentiation (8). Micronucleus (MN) formation can be induced by drugs that cause DNA damage, and these treatments usually affect the biology of cancer cells (9,10).

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MN frequency has been extensively used as a biomarker of genomic instability and chromosomal damage. This structure usually contains whole chromosomes or chromatin fragments. MN assay has been used in studies of population biomonitoring (11,12), evaluation of the genotoxic potential of the compounds (13,14), studies of the cancer chemopreventive agents (15), and evaluation of the biological effects in aquatic organisms exposed to chemical pollution (16).

A549, a non-small lung carcinoma cell line, after the treatment with vincristine and aphidicolin showed an increased rate of nuclear abnormalities, including MNs formation (17). A549 cells contain 3.4 copies of ErbB1 gene that encodes EGFR, which is a 170-kDa protein with an extracellular domain, a transmembrane domain, and an intracellular tyrosine kinase domain (18-20). EGFR is frequently overexpressed in many cancer types, including lung cancer, and has been directly associated with cancer progression (21-23).

Studies regarding the loss of amplified oncogenes by MN expulsion can contribute to elucidate the possible functions of MN formation in cancer cells. Thus, this study aimed to standardize a protocol for cell cycle synchronization to further analyze the possible association of MN formation with the extrusion of ErbB1 gene extra-copies in A549 cells.

Material and Methods

Cell culture

A549 cell line was obtained from the American Type Culture Collection (ATCC) and was maintained in Dulbecco's modified Eagle's minimum essential medium (DMEM, Sigma, USA). It was supplemented with 10% fetal calf serum (FCS; Cultilab, Brazil). Then, this cell line was cultured in a 37°C humidified incubator in an atmosphere of 5% CO₂.

Mitotic index

The cells were seeded on coverslips in 35-mm dishes at 3x10⁴ cells/dish. At different times (according to the assays specified in the Results section), the cells were fixed with 3.7% formaldehyde for 30 min. Then, the cells were washed with phosphate-buffered saline (PBS) and permeabilized with 0.5% Triton X-100 for 10 min, washed again and treated with ribonuclease (RNase; 10 mg/mL) for 30 min. They were incubated with 1% bovine albumin (Sigma, USA) for 1 h. Then, anti-phospho-histone H3 (Upstate, USA) was added and the cells were incubated in a humidified chamber for 3 h. The samples were also incubated with secondary antibody fluorescein isothiocyanate (FITC)-conjugated for 1.5 h. Nuclei were counterstained with propidium iodide (PI) and 1000 cells/slide were counted using a fluorescence microscope (Nikon, EDF-3, Japan). The images were obtained using a confocal laser-scanning microscope (LSM 510, Zeiss, Germany).

Nuclear aberrations and lamin-B labeling

Cells were seeded on coverslips in 35-mm dishes at 3x10⁴ cells/dish. After the double blockade by gradual FCS removal followed by vincristine treatment (detailed in the results), the cells were fixed with 3.7% formaldehyde for 30 min, washed again with PBS and permeabilized with 0.5% Triton X-100 for 10 min. RNase (10 mg/mL) was added for 30 min. In order to observe the nuclear envelope, the samples were incubated with anti-lamin B antibody (Santa Cruz, USA) for 15 h in a humidified chamber. We added secondary antibody (FITC-conjugated) for 1.5 h. Nuclei were counterstained with PI and 1000 cells/slide were counted using a fluorescence microscope (Nikon, EDF-3). The images were obtained using a confocal laser-scanning microscope (LSM 510, Zeiss).

5-bromo-2-deoxyuridine (BrdU) incorporation

After BrdU incorporation (depending on the protocols as shown in the Results section), at different times, the cells were fixed with cold methanol for 30 min, followed by 3.7% formaldehyde for 10 min. The different time durations of BrdU incorporation during synchronization is detailed in the Results section. The cells were permeabilized with 0.5% Triton X-100 for 10 min and washed with PBS. RNase was added for 30 min. We then added anti-BrdU antibody+nuclease (GE Healthcare, UK) for 40 min. The samples were washed with PBS and secondary antibody FITC-conjugated was added for 1 h. Nuclei were counterstained with PI and 1000 cells/slide were counted using a fluorescence microscope (Nikon, EDF-3). The images were obtained using a confocal laser-scanning microscope (LSM 510, Zeiss).

Characterization of the cell cycle in A549 cells

The times of each phase of the cell cycle of A549 cells was determined after making some modifications in the protocol as proposed by Uzbekov et al (24). Briefly, Gap 2 (G2) was evaluated through the frequencies of mitotic cells labeled with anti-BrdU after 1 h of BrdU pulse. The frequencies were determined by fixing cells in intervals of 30 min for 7 h. The first labeled mitotic cells were observed after 3 h. The minimum length of G2 is considered to be the summation of this time plus the time of the BrdU pulse (G2min = 4 h). The maximum length of G2 is considered to be the summation of the time when the frequency of mitotic cells labeled with anti-BrdU is maximum and the period of the BrdU pulse after excluding a half of the mitotic time (see below). We found that G2max = 7 h. Thus, according to Uzbekov et al. (25): G2 = (G2min + G2max/2) – (time of BrdU incorporation). G2 = (4 + 7/2) –1 = 4.5 h.

To determine the length of cell cycle and S phase, the cells were incubated with BrdU and fixed at intervals of 3 h for 36 h. The frequencies of labeled cells increased from 36.7% at the beginning of the experiment to 88.8%
after 27 h \((x = 0.888)\). This time duration was considered to calculate the length of the cell cycle (Lcc), where Lcc=S = 27. The length of S phase (S) was calculated from the initial frequency of positively labeled BrdU cells, where \(S = 0.367\).

Thus, according to Uzbekov et al. (24): \(S/Lcc = 0.367/0.888 = 0.413\). So, \(S = 0.413\) Lcc. Once Lcc=S = 27, we found that: Lcc = 46 h and \(S = 19\) h.

The time of the mitosis (M) was calculated using the mean of the mitotic index (2.45% as per the our results of mitotic index). According to Uzbekov et al. (25), M is denoted by the mean of the mitotic index multiplied by the time of the whole cell cycle. Thus: \(M = 0.0245 \times 46 = 1\) h 7 min.

Finally, the duration of Gap1 (G1) phase = Lcc–G2–M. So, \(G1 = 21.3\) h.

**Fluorescent in situ hybridization (FISH)**

According to the manufacturer’s instructions, FISH protocol, with some modifications, was performed using the kit ZytoVision SPEC EGFR/CEN 7 Dual Color probe (Germany) (26). At the end of the protocol, the samples were washed in PBS for 1 min and incubated with TOPRO-3 for 20 min. The images were obtained using a confocal laser-scanning microscope (LSM 510, Zeiss).

**Statistical analysis**

The statistical analyses were performed using the test of equality of two proportions. It enabled us to evaluate if the proportion of responses of two variables and/or their levels were statistically significant. In order to further complement the descriptive analysis of the quantitative values, we applied the confidence interval to the mean.

**Results**

**Cell cycle synchronization**

The A549 cell cycle was determined based on the labeled mitoses method using BrdU pulse, kinetics of BrdU incorporation, and mitotic indexes. Cell cycle phases were calculated according to Uzbekov et al. (24). The mean cell cycle lasted 46 h with G1 corresponding to 21.5 h, S phase to 19 h, G2 to 4.5 h, and mitosis to 1 h and 7 min, on average. Based on these data, a schedule was designed to attempt the synchronization of A549 cells (Figure 1).

The FCS was gradually removed every 24 h until it reached 0% in the culture medium. We anticipated that after 50 h of FCS privation, the majority of the cells would be arrested in G0/G1 phase. The medium was then supplemented with 10% FCS for 44 h so that the cells could progress in cell cycle. Vincristine (0.1 \(\mu\)g/mL) was added to the medium for 6 h to block the serum-deprived synchronized cells in subsequent mitosis (Table 1). The time of vincristine treatment was determined based on the length of A549 cell cycle.

Other concentrations of the drug were tested (0.05 and 0.5 \(\mu\)g/mL; Table 1). However, the concentration of 0.1 \(\mu\)g/mL was chosen, because of the higher frequency of c-mitosis until T6, and the absence of c-mitosis in T18, T24, and T48, denoting the progression in cell cycle after drug removal. Vincristine at 0.5 \(\mu\)g/mL also induced high frequency of nuclear abnormalities. We considered c-mitosis as mitosis blocked by vincristine. Morphological characteristics of c-mitosis are disorganized spindles and the condensed chromosomes are unorganized in the equatorial plate (Figure 2).

Micronucleated, binucleated, and multinucleated cells were counted at different recovery times, together with mitosis and c-mitosis (mitosis blocked by vincristine) (Table 1). The cells arrested in M phase after the treatment with vincristine were considered to be synchronized. We found 33.8% of c-mitosis in T6 compared to 18% in T0. The percentage achieved in T6 indicated the frequencies of synchronized cells. Figure 2 illustrates the persistence of vincristine activity on the microtubules of A549 cells 6 h after its removal from the culture medium.

We observed multinucleated and binucleated cells in all groups and there was a gradual increase in frequencies of nuclear aberrations over the recovery times. The frequency of the micronucleated cells was higher in T18 (7.4%) compared to the non-synchronized cells (basal frequency of micronucleated A549 cells = 1.48%) and to the other recovery times (5.8% in T24 and 6.2% in T48). Examples of such alterations are shown in Figure 3. The labeling of the nuclear lamina led to confirmation of the micronuclei integrity.

**Origin of micronuclei**

As described above, we found MNs after recovery of vincristine treatment and, then, we tried to determine the origin of these MNs (mitotic or interphasic) by making use of the pulses of BrdU incorporation as a tool. Figure 4A shows the examples of cells with nuclei and micronuclei that incorporated BrdU (BrdU+). Two schedules of BrdU incorporation have been designed: BrdU-1, wherein the pulse of BrdU occurred when the synchronized cells were in the S phase; and BrdU-2, wherein the pulse of BrdU occurred 1 hour before withdrawal of vincristine (Figure 4B). The cells were fixed after 0, 6 and 18 h of recovery. We analyzed frequencies of interphase nuclei, c-mitoses, and MNs. The results are reported in Table 2.

With regard to BrdU-1, cells in T0 showed 40.1% BrdU labeled nuclei and cells in T6 and T18 showed 42.1% and 47.3% labeled nuclei, respectively. As expected, T6 showed the highest frequency of c-mitosis with BrdU incorporation, while T18 showed the lowest frequency of c-mitosis BrdU+. In T0 and T6, the frequencies of c-mitosis BrdU+ were higher than c-mitosis BrdU-. During BrdU incorporation, synchronized cells were in the S phase and in T0, they were blocked in c-mitosis with BrdU labeling. We evaluated around 100 MNs for each group.
and 43% of them were found to be positive to BrdU incorporation in T0. T6 and T18 showed 23% and 36% of positive MNs, respectively. These BrdU-labeled MNs in T18 were originated probably during mitosis. The MNs without BrdU labeling were originated from cells that were not in the S phase at the BrdU pulse and these cells were considered to be non-synchronized.

In the next set of experiments considering BrdU-2, it was anticipated that the synchronized cells would be arrested in mitosis, but we found such cells that incorporated BrdU and, therefore, these cells were non-synchronized. As expected, c-mitosis in T0 did not show BrdU labeling. The analysis of the cell population showed an increased frequency of BrdU+ MNs. T0, T6 and T18 showed 25%, 34% and 47% of BrdU labeling into MNs, respectively. The BrdU+ MNs observed in T0 and T6 couldn’t be induced by vincristine and they probably originated in interphase. Again, T18 also had enrichment of interphasic MNs, although some of them were probably formed in mitosis. The results can be viewed in Table 2.

These experiments showed that MNs could originate in mitosis as well as in interphase. Then, we chose T18 for the later experiments because of the higher frequency of MNs achieved at this recovery time.

**Characterization of micronuclei content**

ErbB1 gene amplification in lung cancer cells is a very common observation. We evaluated if selective oncogene elimination was possible in the MNs shown in the ErbB1 gene. We also analyzed the cells submitted to synchronization protocol (T18 group) and compared them with the cells cultured in normal medium, supplemented with 10% FCS (control group).

Figure 5 shows cells with examples of MNs positive or negative to ErbB1 labeling (that is ErbB1+ and ErbB1−, respectively). We observed 21.11% and 14.44% of MNs

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**Table 1.** Effects of double blockade (FCS deprivation + 0.05, 0.1 or 0.5 μg/mL vincristine) on A549 cells.

| Nuclear aberrations           | T0   | T6   | T18  | T24  | T48  |
|-------------------------------|------|------|------|------|------|
| 0.05 μg/mL vincristine        |      |      |      |      |      |
| c-mitoses                     | 21.1 | 10.1 | 0.4  | 0.2  | 0.1  |
| Total mitoses                 | 23.1 | 12.5 | 1.3  | 3.4  | 2.3  |
| Micronucleated cells          | 0.9  | 1.0  | 8.1  | 4.4  | 4.7  |
| Binucleated cells             | 1.2  | 1.1  | 7.3  | 5.8  | 5.9  |
| Multinucleated cells          | 0.2  | 2.5  | 11.6 | 10.8 | 6.2  |
| 0.1 μg/mL vincristine         |      |      |      |      |      |
| c-mitoses                     | 18.0 | 33.8 | 0.0  | 0.0  | 0.0  |
| Total mitoses                 | 19.7 | 35.0 | 1.2  | 1.1  | 1.8  |
| Micronucleated cells          | 0.7  | 0.7  | 7.4  | 5.8  | 6.2  |
| Binucleated cells             | 2.1  | 2.1  | 5.8  | 9.2  | 10.3 |
| Multinucleated cells          | 0.2  | 2.7  | 13.2 | 16.1 | 14.6 |
| 0.5 μg/mL vincristine         |      |      |      |      |      |
| c-mitoses                     | 14.4 | 31.5 | 1.0  | 1.8  | 0.4  |
| Total mitoses                 | 15.0 | 32.2 | 2.1  | 3.1  | 1.2  |
| Micronucleated cells          | 0.6  | 0.3  | 10.8 | 15.4 | 23.7 |
| Binucleated cells             | 1.4  | 1.2  | 7.0  | 6.8  | 8.0  |
| Multinucleated cells          | 0.1  | 0.1  | 24.9 | 39.0 | 45.0 |

Results are reported as percent of 1000 cells counted. FCS: fetal calf serum; T: times of recovery after vincristine removal (0, 6, 18, 24 and 48 h).
ErbB1+ in control cells and in the T18 group, respectively. But these differences were not statistically significant. The quantification of ErbB1+ MNs in control cells and in the T18 group indicated that there was no selective elimination of the ErbB1 gene in MNs of synchronized cells.

Discussion

BrdU incorporation for 1 h resulted in 36.75% of A549 cells BrdU+ and they represented the S phase frequency in the cell population. Chang et al. (27) studied the A549 cell cycle by cytometry and reported that the percentage of cells in the S phase was 32.1 after 1 day and 38.9% after 2 days in culture. They also showed that 54% of the cells were in G1 phase, indicating the longer duration of this phase in A549 cells.

The drugs that block the cell cycle at specific points have been used to obtain synchronized cells. However, the methods should be standardized for each cell type taking into consideration the duration time of cycle phase and the cell behavior after treatment. Many drugs were used to cause cell cycle arrest: a) thymidine, aphidicolin, mimosine, hydroxyurea, and 5-fluorodeoxyuridine were used to cause cell cycle arrest in the S phase; b) N-acetyl-leucyl-leucyl-norleucinal (ALLN), nocodazole, colchicine and colcemide in mitosis; and c) lovastatin and Hoechst 768159 in G1 phase (methods reviewed by Uzbekov, (28)).

Synchronization of tumor cell population is difficult because tumor cells have the ability to avoid the effects of

Figure 2. Cytoskeleton analyses in A549 cells submitted to double blockade. Upper panels show the integrity of microtubules and microfilaments of A549 cells in interphase (upper left) and in mitosis (upper right), in the control cells. Lower panels show A549 cells in T6 with microtubule network disorganization in interphase (lower left) and in c-mitosis (lower right) after treatment with vincristine. Images are projections of different slices obtained by laser scanning confocal microscopy and show microtubule in blue and microfilaments in green. Nuclei were counterstained with propidium iodide (red).
many drugs. One such example is the expression of genes responsible for the multidrug resistance (MDR) phenotype. Our group could not find efficient conditions of colchicine treatment to induce cell cycle arrest in A549 cell line (data not shown). Instead, vincristine was the alternative used to block the cell cycle in mitosis.

As observed at T6, with an increase in the frequency of c-mitosis, even after the removal of vincristine, it still continued to act in the cells. After 18 h of recovery, all the cells progressed in the cell cycle because there was no evidence of c-mitosis. Keeping in mind that A549 cells could show mechanisms of drug resistance against vincristine and other drugs, which is common in cancer cell lines, 33.8% of synchronized cells could be considered relevant for this study (29). This possible mechanism of drug resistance in A549 cells is supported by the negative results obtained with colchicine treatment. This rate of synchronized cells was supported by statistical analysis.

The data of synchronized cells allowed us to infer that...
many A549 cells evaded the mitotic checkpoint. In this case, despite the microtubules disassembly, the cells progressed in the mitosis, increasing the frequencies of micronucleated, binucleated, and multinucleated cells (30). The A549 mitotic checkpoint is considered to be functional. Weitzel and Vandre´ (31) and Masuda et al. (32) showed that the treatment of A549 cells with drugs that interfere with the spindle dynamic were responsible for causing the cell cycle arrest in mitosis.

The results showed that the synchronization was efficient in causing enrichment of micronucleated cell population in the cultures. This enrichment could be considered to be a useful tool in studying the expulsion of amplified genes by micronuclei formation. Nevertheless, micronuclei induced by vincristine shown in our results were not spontaneous micronuclei and thus it was not possible to determine if they originated during interphase or mitosis.

Micronuclei observed in T0 and T6 could not be generated in synchronized cells because these cells were supposed to be arrested in c-mitosis. As shown in Table 1, the cells that incorporated BrdU in the S phase probably evaded the cell synchronization according to the presented schedules.

BrdU incorporation allowed us to elucidate the origin of MNs observed in T18, in mitotic and interphasic cells. We observed cells that incorporated BrdU in the S phase probably evaded the cell synchronization according to the presented schedules.

BrdU incorporation allowed us to elucidate the origin of MNs observed in T18, in mitotic and interphasic cells. We observed cells with MN BrdU− and nuclei BrdU+. The MNs were able to replicate their DNA content and this process seemed to be non-synchronic with the main cell nuclei. Okamoto et al. (33) also showed similar results and it was observed that the timing of replication in the MN and the nucleus sometimes were non-synchronic, depending on the MN content.

Some authors have proposed different mechanisms for MN formation (8,34-36). Many previous works have already demonstrated that there are amplified genes of extrachromosomal regions in the MNs of tumor cells. One example is the gene MYC, which was observed in DM, in the hydroxyurea-induced MNs in human colorectal carcinoma (37,38). Villa et al. (39) observed MYC in MNs of leukemia cells and Valent et al. (40) showed that this

![Figure 5. Fluorescent in situ hybridization of ErbB1 gene in A549 cells. A, Control cells (from non-synchronized cell culture) showing usually 3 copies of ErbB1 gene in each nucleus. B, Cells submitted to synchronization protocol after 18 h recovery. Note a micronuclei positive to ErbB1 labeling (red arrow) and micronuclei negative to ErbB1 labeling (white arrow). Multinucleated cells showed a deviation of the usual number of copies of ErbB1. The ErbB1 gene was labeled in green. Nuclei were counterstained with TO-PRO-3 (blue). Images were obtained by laser scanning confocal microscopy.](image-url)
gene, as DM, was found in spontaneous MNs of neuroblastoma in vivo. Hélias-Rodzewicz et al. (8) also associated the elimination of amplified CDK4 sequences with differentiation in liposarcoma.

The frequency of micronuclei positive to BrdU in T18 (36%) showed the importance of this time of recovery in our analysis. This recovery time also showed the highest frequency of micronucleated cells (7.4%). ErbB1 was found to be amplified in many lung cancer cells, especially in A549 cells (about 3 copies/nucleus) (18,26). Thus, we investigated the relation between ErbB1 amplification and its presence in the micronucleated A549 cells.

We compared the frequency of MNs ErbB1+ in the control and T18 groups. Statistical analysis did not show any differences between these groups; thus, we concluded that there was no preferential extrusion of ErbB1 gene by synchronized cells. Shimizu et al. (38) further demonstrated that DM can originate aggregates that get delayed during the chromatid separation in anaphase, eventually leading to MN formation. ErbB1 was not found in the extrachromosomal regions of the A549 cell line in our culture conditions. This result could help explain the low frequency of MNs ErbB1+.

Figure 6 summarizes the results and shows the schedule of synchronization and the possibilities of MN formation in mitosis or interphase obtained by BrdU-1 and BrdU-2 assays in the T18 group. Due to the enrichment of MNs in the T18 group, this time was chosen for evaluating the possible extrusion of extra copies of ErbB1. Nevertheless, our data suggested that there was no selective elimination of this gene in this system.

This research is an important piece of the puzzle that seeks to elucidate the meaning of MN formation and its association with the elimination of oncogenes or other amplified sequences from tumor cells. The synchronization protocol was efficient to cause the enrichment of the MN population in A549 cells and it could be useful in studying MN formation in this cell line.

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