In vitro differentiation of rhabdomyosarcomas induced by nickel or by Moloney murine sarcoma virus

P. Nanni1,2, G. Azzarello1, L. Tessarollo2,4, C. De Giovanni1,2, P. -L. Lollini1,5, G. Nicoletti1,5, K. Scotlandi1, L. Landuzzi1, M. Panozo2,4, E. D'Andrea2,4, S. Schiaffino3 & L. Chieco-Bianchi2,4

1Institute of Cancerology, University of Bologna; 2Interuniversity Center for Cancer Research (CIRC); 3Institute of General Pathology, CNR Unit for Muscle Biology and Physiopathology, University of Padua; 4Institute of Oncology, University of Padua, and 5National Institute for Cancer Research, Genoa, Section of Bologna, Italy.

Summary In vitro cultures and clonal derivatives have been established from rat rhabdomyosarcomas induced by Moloney-Murine Sarcoma Virus (MSV) or by nickel sulfate; differentiation ability has been studied as expression of desmin, embryonic and adult myosin isoforms, a-actin isoforms and cellular fusion. The two rhabdomyosarcoma models showed different levels of myogenic differentiation. Multinucleated myotube-like structures were frequently observed in cultures derived from nickel-induced tumours. Desmin was present in 50–80% of cells and embryonic myosin in up to 10%. In MSV-tumour-derived cultures and in their metastases or clonal derivatives two cell types are present in different ratios: spindle-shaped cells, adherent to plastic surfaces, and rounded cells, loosely attached or floating free in the medium. These showed features of myogenic differentiation (10–80% desmin-positive cells), but embryonic myosin expression and production of multinucleated myotube-like structures were very rare events. Cultures from autochthonous lymph node and lung metastatic cells showed similar patterns of differentiation. Retinoic acid increased differentiated features (myotube formation and embryonic myosin expression) only in nickel-induced rhabdomyosarcoma cells. The two models described here mimic the heterogeneity in differentiation pattern found among human rhabdomyosarcomas. Myogenic differentiation ability was retained at a good level by nickel-induced tumours, whereas it was strongly impaired in MSV-induced tumours.

Inoculation of Moloney-murine sarcoma virus (MSV) in newborn rats can induce rhabdomyosarcomas which grow progressively and consistently metastasise to the regional lymph nodes and lungs (Lasneret, 1967; Perk et al., 1968). Rhabdomyosarcomas can be induced in rats also by injection of nickel (Gilman, 1962), an established carcinogen/mutagen in human and animals (Sen & Costa, 1985; Tomatis et al., 1989).

A few data on differentiative ability of these two experimental rhabdomyosarcoma models are reported in the literature (Hildebrand et al., 1980; Altmannsberger et al., 1985; Azzarello et al., 1987; Babai et al., 1988; Borrione et al., 1988). Moreover, cell cultures have been rarely established only from nickel-induced rat rhabdomyosarcomas (Pot-Deprun et al., 1983). A better characterisation and comparison of cell cultures from nickel- and MSV-induced rhabdomyosarcomas could lead to the establishment of an interesting animal model.

We therefore derived cultures and clones from MSV- or nickel-induced rat rhabdomyosarcomas and from autochthonous metastases, to compare in vitro the differentiation ability of cells transformed by these two carcinogenic agents and to evaluate the dynamics and the modulation of the myogenic differentiation process.

Materials and methods

Tumour induction

Wistar/Furth rats were obtained from Dr G. Parmiani, Istituto Tumori, Milano, Italy, and maintained thereafter through inbreeding under conventional conditions. The murine sarcoma virus, Moloney isolate, originally obtained from Dr J.B. Moloney, was maintained by in vivo serial passages in 1–2 week old Balb/c mice. Cell-free extracts from pooled neoplastic tissue were prepared as previously described (Colombatti et al., 1975). The preparation used for the present study had an in vitro titre of 2 x 10^12Focus Forming Units (FFU) ml−1 (Hartley & Rowe, 1966), when tested on SC-1 cells: rhabdomyosarcomas were induced by i.m. injection of 10^10 FFU into the thigh of newborn rats. Rhabdomyosarcomas were also induced by i.m. injection of 10 mg of NiS2 suspended in about 0.5 ml olive oil into the thigh of adult (250 g) rats (Borrione et al., 1988). Animals were handled according to the European guidelines.

Cells culture and cloning

Cells derived from two nickel-induced rhabdomyosarcomas (NI-1 and NI-2), and from two MSV-induced tumours (MSV-1 and MSV-2) and autochthonous metastases were adapted to grow in Dulbecco's MEM (DMEM) supplemented with 100 U ml−1 penicillin, 100 μg ml−1 streptomycin and either 10% foetal calf serum (FCS) (proliferation medium) or 2% horse serum (HS) (differentiation medium). Differentiation medium was previously found to enhance myogenic differentiation in some myogenic model systems (Nanni et al., 1986; Dym & Yaffe, 1979). Cells were routinely subcultured approximately 1–2 times a week, at dilutions from 1:3 to 1:8, and incubated in an atmosphere with 5% CO2 at 37°C. With the purpose to study cultures as close as possible to the originating in vivo-grown populations, cells between the 5th and 15th in vitro passage were used throughout the study. Cell cultures have been further propagated reaching the 30th in vitro passage, without loss of proliferative ability or appearance of peculiar features. Both nickel- and MSV-derived rhabdomyosarcoma cell cultures are tumorigenic when injected subcutaneously into nude mice or 2–3 week old syngeneic rats.

Clones were isolated either with cloning cylinders from a Petri dish containing sparse colonies 14 days after seeding of 15–30 cells cm−2 or by picking up individual colonies grown in 0.33% agar 14 days after seeding in 60 mm Petri dish of 1,000–3,000 nickel-induced and 3,000–30,000 MSV-induced rhabdomyosarcoma cells.

DNA extraction and Southern blot analysis

High-molecular-weight DNA was prepared by cell lysis, proteinase K digestion, extraction with phenol-chloroform, and
precipitation with ethanol (Sambrook et al., 1989). DNA (10 μg) was digested with the appropriate restriction endonuclease, electrophoresed through 0.8% agarose gel, denaturated, neutralized and transferred to Hybond-N (Amersham, England) filters according to Southern (1975). Filters were hybridised overnight to a DNA probe labelled by random priming with 32P-dCTP (Feinberg & Vogelstein, 1983) at 42°C in 50% formamide, washed at 65°C in 0.1 × standard saline citrate (SSC: 0.15 M NaCl, 0.015 M Na citrate), and 1% sodium dodecyl sulfate (SDS) for 1 h, and exposed to Kodak X-Omat S films (Kodak, Rochester, NY) at −80°C with intensifying screens. The 450 bp v-mos probe used in this study was derived by PstI digestion from the pMSV-31 plasmid (Jones et al., 1980).

Monoclonal antibodies and evaluation of differentiation

The anti-desmin monoclonal antibody was purchased from Boehringer-Mannheim (Mannheim, Germany). Monoclonal antibodies against α-sarcomeric and α-smooth muscle actin isoforms were purchased from Sigma Chemical Co., St Louis, USA, and from Sclavo, Siena, Italy. Myosin isoforms were specifically stained with the following antibodies: BF-G6 (embryonic), BA-D5 (type 1), SC711 (type 2A), BF-F3 (type 2B). The reactivity of these antibodies against rat rhabdomyosarcomas has been previously characterised (Azzarello et al., 1987; Borrione et al., 1988).

Cells were harvested, counted, and centrifuged at 400 g for 10 min onto glass slides. Cytocentrifuge slides were immediately fixed with methanol:acetone (3:7) at −20°C and stained in an indirect immunofluorescence assay. Slides were examined under a Reichert Biowar microscope equipped for phase contrast and fluorescence. At least 300 cell elements (either mono- or multinuclear) in random fields were scored at 312.5 × for determining the percentage of stained cells. To evaluate the percentage of multinucleated cells, after washing off the unbound fluorescein-conjugated second antibody (Sera-Lab, Bicester, UK), cell nuclei were stained with ethidium bromide (100 μg ml⁻¹ in phosphate-buffered saline) for 5 min. After extensive washings and mounting, slides were examined under a Reichert Biowar microscope equipped for phase contrast and green-red fluorescence. At least 200 nuclei in random fields were scored at 1250 ×.

All trans-retinoic acid (Aldrich, Milwaukee, USA) was stored at −20°C as a 1 mM stock solution in ethanol; it was added at 1 μM final concentration in proliferation medium to cultures 24 h after seeding of 0.2 × 10⁶ cells in 25 cm² flasks. After 3 days, medium was changed with retinoic acid-containing differentiation medium and cultures incubated for additional 4 days. Controls with ethanol-containing medium were performed in parallel. At the end of experiment, myotube formation was evaluated by double-blind count of myotube-like structures on 15 random fields (phase contrast, × 100) of control and treated cultures: myotubes number was then multiplied by the ratio between flask and observed surfaces to have an estimate of total number. Cell yield was then determined and cytocentrifuge samples prepared and stained for embryonic myosin as reported above.

Results

Nickel-induced rhabdomyosarcomas

Cultures from nickel-induced tumours (NI-1 and NI-2) consisted of polygonal or spindle-shaped cells and multinucleated myotube-like structures (Figure 1). Desmin and embryonic and adult myosin, specifically expressed during myogenic differentiation of normal and tumour cells (Altmannsberger et al., 1985; Nanni et al., 1986; Schiaffino et al., 1986; Eusebi et al., 1986; Dias et al., 1987; Kelland et al., 1989), were studied by immunofluorescence technique in monolayer or cytocentrifuge samples. All cells showed expression of vimentin molecules. The anti-desmin antibody stained near 50–80% of the cells and a very strong positivity was evident in myotube-like structures (Figure 2a). Myotube-like structures

Figure 1 In vitro morphology of nickel-induced rhabdomyosarcoma cell cultures. a, NI-1; b, NI-2. Phase contrast, × 100.

Figure 2 Expression of markers of myogenic differentiation in NI-1 rhabdomyosarcoma cell cultures. a, desmin; b, embryonic myosin (BF-G6). Immunofluorescence, × 500.
showed a strong positivity for embryonic myosin (Figure 2b) whereas most small cells were negative; quantitative data are reported in Figure 3. The expression of type 1 or type 2 myosin isoforms was also studied: positive cells were found only rarely (<0.1%). Expression of α-sarcomeric actin was found sporadically, whereas α-smooth muscle actin was observed in 10–40% of cells.

NI-1 and NI-2 cells grown in 4 days in proliferation medium and then shifted to differentiation medium (DMEM supplemented with 2% horse serum) and cultured for additional 7 days showed impressive increase in myotubes (Figure 4); expression of embryonic myosin was also increased (Figure 5). A parallel slight increase in α-sarcomeric actin-positive cells was observed as well (data not shown).

In order to rule out any influence of heterogeneity on differentiation, NI-1 clones were isolated from adherent or agar cultures and studied for the expression of desmin and embryonic myosin (Figure 3). Most clones showed a differentiation phenotype similar to that of parental cells. However, we also obtained two clones (NI-1/1 and NI-1/4) which showed morphological pattern and marker expression attributable to a less differentiated phenotype: a sporadic induction of myotubes and of embryonic myosin-positive cells when cultured in differentiation medium was observed.

A clone with a differentiation pattern similar to that of parental cells (NI-1/B) was chosen for further study on the modulation of differentiated features. A kinetic study of NI-1/B cells showed an increase in myotube formation and in embryonic myosin expression during culture in differentiation medium (Figure 5).

The treatment of NI-1/B cultures with retinoic acid, a known inducer of differentiation (Reiss et al., 1986; Gabbert et al., 1988; Waxman et al., 1988), caused a significant increase in myotube formation and in embryonic myosin-positive cells, without affecting growth rate (Figure 6).

**MSV-induced rhabdomyosarcomas**

We derived in vitro cultures from two MSV-induced rhabdomyosarcomas (MSV-1 and MSV-2), from metastatic lymph nodes (MSV-1-LNM and MSV-2-LNM) and from lung metastases (MSV-1-UMP1, MSV-1-UMP2, MSV-1-UMP3, MSV-2-MP). In all cultures a few spindle-shaped cells attached to the surface were observed along with many rounded cells, loosely attached or floating free in the medium (Figure 7).

A large set of clonal derivatives was obtained from all cultures: clonality was assessed by Southern blot analysis using a v-mos specific probe. As an example, molecular analysis of MSV-1 derived cell clones is reported in Figure 8: besides the 12 Kb germline c-mos fragment present in all cells, one to three additional bands corresponding to clonally integrated MSV proviruses were detected using the EcoRI restriction endonuclease (which does not cut within the proviral DNA). This restriction pattern suggests an oligoclonal origin of MSV tumours, as shown in a different viral system (D'Andrea et al., 1987). Moreover, equimolarity of proviral and germline bands suggested that one to three copies of MSV provirus were present in the same cell. As expected, DNA restriction with SstI, which cuts within the provirus LTR, disclosed a single additional 5.3 Kb proviral band with

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**Figure 3** Expression of desmin and embryonic myosin in nickel-induced rhabdomyosarcoma cultures and clonal derivatives. Clones derived from adherent and agar cultures are designated by letters or figures, respectively.

**Figure 4** Induction of myotube-like structures in NI-1 a,b,c, and NI-2 d,e,f cells cultured 4 days in proliferation medium a,d and 7 additional days in differentiation medium (b,e: bright field, ×25; c.f: myotubes, phase contrast, ×100). The percentage of multinucleated cells in NI-1 and NI-2 cultures increased from 0.8% and 1.7% at 4 days to 17.3% and 10.7% at 11 days.
an intensity, when compared to the 2.8 Kb germline c-mos, proportional to the number of proviruses detected after EcoRI restriction.

MSV-1 and MSV-2 cell cultures and clonal derivatives showed variable levels of desmin production and a very low expression of embryonic myosin and \(\alpha\)-smooth muscle actin (for either marker no positive cell was ever detected in most cultures) (Figure 9). Myotube-like structures were never observed in MSV-1 and MSV-2, and no cell expressing slow or fast myosin or \(\alpha\)-sarcomeric actin was found.

To investigate whether cells derived from autochthonous metastatic nodules showed a peculiar differentiation pattern, we compared the expression of vimentin, desmin and embryonic myosin along with the formation of multinucleated myotube-like structures in cultures derived from lymph node and lung metastases: no peculiarity was observed (Figure 9).

Clonal derivatives obtained from MSV-induced rhabdomyosarcomas showed the presence of the two morphological cell types observed in the parental cultures, even if at variable ratios. All clonal derivatives of MSV-1 cells appeared to lack both desmin and embryonic myosin expression (Figure 9), but were characterised by a \(v\)-mos restriction pattern comparable to that of the parental line, thus confirming their rhabdomyosarcomas nature. Clones isolated from MSV-2 cultures retained a percentage of desmin-positive cells similar to that of parental cells. Therefore, the loss of marker expression found for MSV-1 clones is not likely to be due exclusively to cloning procedure.

Culture in differentiation medium failed to induce any increase in myotube-like structure formation and in desmin and embryonic myosin expression (Figure 10). Analogous results were obtained when retinoic acid was added to the cultures (data not shown).

Discussion

Our study showed that cultures obtained from nickel-induced rat rhabdomyosarcomas appear more differentiated than those from MSV-induced rhabdomyosarcomas.

The presence of myotube-like structures in nickel-induced rhabdomyosarcomas along with the expression of embryonic myosin are suggestive of a more conserved differentiation ability, in comparison to MSV-induced tumour cells. The ability to form myotube-like structures has been reported also in rat rhabdomyosarcomas induced by nickel (Pot-Deprun et al., 1983) or by dimethylbenzanthracene (Gerharz et al., 1989).

In our cultures derived from MSV-induced rhabdomyosarcomas and their clonal derivatives, two main morphological components were evident; expression of desmin was variable, whereas expression of myosin or fusion were very rare events. Cell cloning of MSV-1 yielded an even less differentiated phenotype: all clones studied lacked both desmin and embryonic myosin expression. We have no explanations for this decreased expression: it might be that an intense clonal
expansion of a low desmin-expressing population lead to a further differentiation alteration or that some interaction among different subpopulations play a role in differentiation process. However, this is not a general phenomenon, since clones of MSV-2 retained a differentiation pattern similar to that of parental cells.

MSV-induced tumours frequently showed autochthonous lung and lymph node metastases. The relationship between differentiation and metastatic ability is still under debate (Dexter, 1977). Some reports suggest that a higher differentiation is associated to a higher metastatic capacity (Bennett et al., 1986). In the autochthonous system studied here, cells derived from metastatic nodules did not show a peculiar pattern of expression of myogenic differentiated structures, even when evaluated in a kinetic test.

The differences observed in the two rhabdomyosarcoma systems described here might be due to several factors. A possibility is that the two transforming agents interact in vivo with distinct populations of muscle cells: skeletal muscle fibres and satellite cells (myogenic precursor cells in mature muscle) for nickel, and foetal myoblasts for MSV. This might be relevant in determining the differentiation pattern (Schwab & Luger, 1980; Yablonka et al., 1987). The nature of the target cell of chemical and viral carcinogens in the induction of rhabdomyosarcoma is still controversial. Alternatively, the different pattern of differentiation found in cultures of nickel- and MSV-induced rhabdomyosarcomas might be due to intrinsic differences in the oncogenic potential and/or in the mechanism of tumour induction displayed by the two carcinogens. In fact, a differential role of cytoplasmic and nuclear oncogenes in acquisition of the transformed phenotype, subversion of the proliferation control, and interference with the expression of differentiation programs has been proposed.

Figure 7 In vitro morphology of MSV-induced rhabdomyosarcoma cell cultures. a, MSV-1; b, MSV-1-LNM. Phase contrast, × 100.

Figure 8 Southern blot analysis of MSV integration pattern in MSV-induced rhabdomyosarcoma cells and their clonal derivatives. Ten μg of DNA from normal rat kidney (N), MSV-1 cells (M), and clonal derivatives were digested with restriction endonucleases, separated by agarose gel electrophoresis, blotted on nylon filters, and hybridised to 32P-labelled v-mos probe. Molecular weights of c-mos germline fragments (12 Kb or 2.8 Kb) and provirus DNA (5.3 Kb) are reported on the left.
(Alema & Tatò, 1987). Oncogenes, like src, block differentiation directly, through selective regulation of transcription, which is independent from disruption of the proliferative control. On the other hand, nuclear oncogenes block differentiation by an indirect mechanism, via uncontrolled cell proliferation. Thus, differences in differentiation programs of the two experimental systems might be due to the involvement of different oncogenes in tumour induction: v-mos could act directly while the chemical carcinogen would act indirectly through alterations of genes likely linked to the proliferation control program.

The results reported here suggest that cell lines from nickel- or MSV-induced rat rhabdomyosarcomas represent a useful model for studying the expression and modulation of differentiation in well differentiated (nickel) and very poorly differentiated (MSV) myogenic tumour cells.

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