Induction of Interferon and Erythropoietic Differentiation in Cells Transformed by Friend Virus

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Two lines of Friend virus (FV)-transformed mouse spleen cells have been analyzed in respect to their interferon production capacity: neither F4 cells, which liberate infectious FV when kept under tissue culture conditions, nor the thymidine kinase-deficient B8 cells, which do not produce significant amounts of FV, release detectable amounts of autogenous interferon into cell supernatants. However, interferon is produced in these cells in amounts comparable to that in L-929 cells when interferon induction is initiated with UV-inactivated Newcastle disease virus. Conversely poly(I)-poly(C), a potent interferon inducer in L-929 cells, proved ineffective in this capacity in F4 or B8 cells. When erythropoietic differentiation is induced in these cells by dimethyl sulfoxide, no autogenous interferon production occurs, but with NDV-induction a four- to fivefold increase of interferon production is observed. A similar elevation of interferon production is achieved during 5-bromodeoxyuridine stimulation of differentiation in the thymidine kinase-deficient B8 cells. The refractiveness against poly(I)-poly(C) displayed in unstimulated cells is not overcome at any stage of differentiation, indicating major differences of Newcastle disease virus and poly(I)-poly(C) induction mechanisms.

When transformed by Friend virus (FV), mouse spleen lymphocytes can be propagated under tissue culture conditions (9, 21). A cloned line (F4) of FV-transformed mouse spleen cells produces infectious virus (17) containing both components of the FV complex (8), the spleen focus-forming virus, and the murine leukemia virus-Friend (7, 20). Addition of dimethyl sulfoxide (ME₂SO) to the culture medium of these transformed cells induces their differentiation along the erythropoietic pathway (9, 17). The 5-bromo-2'-deoxyuridine (BrdU)-resistant cell-line B8 of the same origin likewise can be induced to differentiate by ME₂SO but also by high concentrations of BrdU (6, 16). This BrdU-induced hemoglobin synthesis in B8 cells (but not the ME₂SO induced) is accompanied by a transient burst in FV synthesis on the second day of BrdU treatment (18). Exogenous interferon inhibits this virus induction (Ostertag, Crozier, and Swetly, manuscript in preparation) as well as the FV release in F4 cells (22), demonstrating that these FV-transformed cells are sensitive to interferon. The synthesis of hemoglobin which is induced during the erythroid differentiation remains unaffected by the presence of interferon (22). Cells to synthesize and release interferon is investigated by using different inducers, Newcastle disease virus (NDV) or poly(I)-poly(C).

Erythropoietic differentiation, stimulated in F4 cells by addition of ME₂SO and in B8 cells by ME₂SO or by BrdU, occurs concurrently for more than 70% of the cells (6) and permits for the first time a study of interferon inducibility at the different stages of cellular differentiation.

MATERIALS AND METHODS

Cells and viruses. The FV-transformed cloned mouse spleen cell line F4 and the BrdU-resistant mutant B8 have been obtained from W. Ostertag, and their isolation and properties are published (6, 16, 21). The cells are cultured in plastic dishes in Basal Eagle medium supplemented with 15% fetal bovine serum, double amounts in amino acids and vitamins, and a fourfold amount of glutamine. Incubation is at 37°C in humidified air containing 5% CO₂.

NDV (California strain) grown in chicken eggs has been obtained from G. Bodo. Hemagglutination has been determined with chicken erythrocytes. For interferon induction UV-irradiated virus has been used. L-929 cells are kept in Basal Eagle medium containing 10% calf serum (inactivated for 30 min at 56°C). L-929 cells serve as controls in the interferon induction experiments with NDV and poly(I)-poly(C) and are used for the plaque reduction test of vesicular stomatitis virus.
Vesicular stomatitis virus, strain Indiana, has been grown in chicken eggs and plaque assayed on L-292 cells.

**Reagents.** ME₄SO (purchased from Merck, Darmstadt; gas chromatography grade) is added without treatment to the culture media where indicated at concentrations between 140 to 250 mM. BrdU has been purchased from Sigma Chemical Co., St. Louis. It is dissolved in 0.2 N NaOH at 20 mg/ml. The final concentration after adding it to B8 cells for stimulation of erythroid differentiation is 0.8 mM; the final concentration to suppress erythroid differentiation in F4 cells is 25 μM.

Poly(I)·poly(C) has been purchased from Miles Laboratories; DEAE-dextran was from Pharmacia; and neomycin was from C. H. Boehringer, Ingelheim.

**Stimulation of erythropoietic differentiation and hemoglobin synthesis.** To cells growing at a density of 0.5 x 10⁴ to 10⁶ cells/ml, ME₄SO is added on day 1 at a concentration of 1%. Cells are then transferred in daily intervals to fresh medium containing ME₄SO at concentrations: 1.25% on day 2, 1.5% on day 3, and 1.75% on days 4 and 5. The cell density throughout the stimulation period is maintained at 10⁶ to 1.5 x 10⁹ cells/ml. The BrdU stimulation of differentiation in BrdU-resistant B8 cells is carried out in cells which have been maintained for a period of 8 weeks in medium without BrdU. Cells are plated then at a density of 10⁵ cells/ml and exposed to medium with 0.8 mM BrdU with daily changes of medium through a 5-day stimulation period. The percentage of cells involved in erythroid differentiation and synthesizing hemoglobin has been determined by methanol fixation of cells on slides and staining with benzidine-Wright-Giemsa. A minimum of 10⁴ cells has been scored for the ratio of benzidine reactive cells to viable cells in culture.

**Production of interferon in the FV-transformed cells.** Production of interferon in F4 and B8 cells has been assayed in supernatants of cultures kept either without ME₄SO or in presence of ME₄SO. Medium which has supported cellular growth for at least 24 h has been centrifuged to remove cells (800 × g, 10 min), and cell debris and supernatant virus (82,000 × g, 50 min) were adjusted to pH 2.5 with 1 N HCl and dialyzed against 0.1 M glycine-hydrochloride buffer, pH 2.5, for 5 days at 4 C. After adjustment to neutral pH, the amount of interferon has been determined by its capacity to inhibit the cytopathic effect of vesicular stomatitis virus in L-929 mouse cells. One unit of interferon is defined as the reciprocal of the highest dilution of a sample that reduces vesicular stomatitis virus plaque formation in a L-929 cell culture by 50%. The plaque assay has been standardized with an NIH reference standard mouse interferon preparation.

**Induction of interferon production.** Two types of interferon-inducing agents have been applied: UV-inactivated NDV and poly(I)·poly(C) with DEAE-dextran and neomycin.

The NDV induction of interferon has been carried out by removing the medium from cells and exposing them to aliquots of 0.6 hemagglutinating units of NDV per 10⁶ cells for 1 h at 37 C, in a volume of 0.5 ml of phosphate-buffered saline. For poly(I)·poly(C) induction, a spectrum of inducer concentrations ranging from 0.5 to 40 μg of poly(I)·poly(C), 10 to 500 μg of DEAE-dextran, and 300 μg of neomycin in phosphate-buffered saline per 10⁶ cells has been applied. Mouse L-929 cells serve as controls in the interferon induction experiments. In these cells interferon induction is optimal when 10⁴ cells are exposed to 4 μg of poly(I)·poly(C), 100 μg of DEAE-dextran, and 300 μg of neomycin for 30 min at 25 C. For the poly(I)·poly(C) induction, the spleen cells are grown as described above and centrifuged at 1,000 x g for 5 min, and the resulting cell pellets are exposed to various concentrations of poly(I)·poly(C) at 25 C for 30 min. Controls are obtained by incubating cells in phosphate-buffered saline without inducers.

After the incubation period the cells are washed four times with medium without serum and further incubated at 37 C in serum-containing medium with the appropriate amounts of ME₄SO or BrdU for interferon induction in differentiating cells. Cell-free supernatants are tested for interferon after 24 h as described above. Cell counts and benzidine staining of cells are carried out to relate interferon production to cell number and to cell differentiation.

Cell numbers are determined by counting the viable cells after trypan blue staining in a hemocytometer.

**RESULTS**

No autogenous interferon production occurs in FV-transformed cells. The aim of this part is a correlation between continuous release of infectious FV from the transformed cells and production of autogenous interferon. Cells of line F4 produce 10⁴ spleen focus-forming units/10⁶ cells and 10⁴ to 10⁵ PFU/10⁶ cells of murine leukemia virus as tested in the XC assay. On the other hand the BrdU-resistant B-8 cells produce only 1 to 5 spleen focus-forming units/10⁴ cells and no detectable leukemia virus component in the XC-test (1, 15, 19).

Both types of cells are plated either under conditions of exponential cell growth (0.5 x 10⁴ to 1.5 x 10⁶ cells/ml) or under conditions of low mitotic activity (2.5 x 10⁴ to 3 x 10⁶ cells/ml). Supernatant medium is tested after 24 h of incubation for interferon as described above. No interferon activity has been found (Table 1) in the supernatants of growing or resting cells, nor in FV releasing or non-producer cells.

ME₄SO or BrdU stimulation of cell differentiation does not lead to autogenous interferon production. When F4 cells or B8 cells are exposed to 1 to 2% ME₄SO for 4 to 5 days, more than 70% of the cells react with benzidine, indicating their erythroid character, and 20 to 30% of the soluble cytoplasmic protein fraction constitutes hemoglobin (17, 22). Parallel to the progress in cell differentiation a reduction in the
TABLE 1. Autogenous interferon production in F4 and B8 cells either unstimulated or at various stages of erythroid differentiation and the interferon induction by NDV and poly(I)-poly(C)

<table>
<thead>
<tr>
<th>Differentiation stimulating agent</th>
<th>Interferon inducer</th>
<th>Interferon production (U/10^6 cells) in cell line:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>F4</td>
</tr>
<tr>
<td>None</td>
<td>None</td>
<td>&lt;10</td>
</tr>
<tr>
<td>ME2SO (0.14 to 0.25 M) (1 to 5 days)</td>
<td>None</td>
<td>&lt;10</td>
</tr>
<tr>
<td>BrdU (10^{-2} M) (1 to 5 days)</td>
<td>None</td>
<td>NT</td>
</tr>
<tr>
<td>None</td>
<td>Poly(I)-poly(C)</td>
<td>&lt;10</td>
</tr>
<tr>
<td>ME2SO (0.14 to 0.25 M) (1 to 5 days)</td>
<td>NDV</td>
<td>5,250</td>
</tr>
<tr>
<td>BrdU (10^{-2} M) (1 to 5 days)</td>
<td>Poly(I)-poly(C)</td>
<td>&lt;10</td>
</tr>
<tr>
<td>None</td>
<td>Poly(I)-poly(C)</td>
<td>NT</td>
</tr>
</tbody>
</table>

* NT. Not tested.

uptake of low-molecular-weight metabolites and a decrease in DNA, RNA, and protein synthesis are observed (6). FV synthesis increases 10-fold in F4 cells in the early stages of stimulation and decreases again from day 3 on (22). The BrdU stimulation of B8 cells leads to a lower level of hemoglobin synthesis (5%) of soluble cytoplasmic proteins and about 40% of benzidine-positive cells), to a similar reduction of membrane permeability and macromolecular synthesis and to a transient induction of FV synthesis on day 2 to 3. With this diversity of events preceding the onset of differentiation, it seemed worthwhile to look for autogenous interferon induction during the course of ME2SO or BrdU treatment. Supernatants harvested in daily intervals after stimulation of differentiation have been assayed for interferon activity. Neither stimulation of F4 or B8 cells with ME2SO nor of B8 cells with BrdU leads at any stage of this differentiation period to detectable autogenous interferon production (Table 1).

Induction of interferon production succeeds with UV-inactivated NDV but not with poly(I)-poly(C). To relate the failure of interferon production in the FV releasing cells, either to a defect caused by the inability of the transformed cells to synthesize interferon or to a defect of FV to induce it, the following experiment has been performed. F4 or B8 cells are exposed to either of two established interferon inducers: UV-inactivated NDV and the double-stranded polynucleotide poly(I).poly(C). This second inducer is ineffective by itself in another mouse cell line (L-929); however, in the presence of DEAE-dextran (5) and neomycin (2), it induces interferon production in L-929 cells to a level of 5,000 U/10^6 cells. A cell toxic effect is associated with this interferon induction. The results are summarized in Table 1 and represent data from five experiments. For induction of interferon in F4 and B8 cells, the concentration of poly(I)-poly(C) has been varied between 0.5 and 40 μg per 10^6 cells. Neither of the applied concentrations, without or in combination with various amounts of DEAE-dextran and with neomycin, have resulted in an induction of interferon or in significant cell toxicity. In contrast to this inability of poly(I)-poly(C) to induce interferon in the FV-transformed spleen cells, UV-inactivated NDV induces about 5,000 U of interferon/10^6 cells. No significant difference in the interferon production has been found between the FV releasing F4 cells and the B8 cells, which do not produce virus.

The amount of interferon induced by NDV depends on the stage of erythropoietic cell differentiation. Not only hemoglobin synthesis increases but also the release of FV from F4 cells goes up more than 10-fold when ME2SO is added and cells begin to differentiate (22). However, when F4 cells are exposed to 2 x 10^{-8} M BrdU for 2 days before and then during the 5 days of ME2SO treatment, no significant effect on cell growth and viability occurs, but hemoglobin synthesis is inhibited (16). Interferon may be seen also as a specialized cellular product, and thus it appeared interesting to follow the capacity of the cell to respond to NDV induction of interferon at different stages of the erythropoietic differentiation.

The F4 cells have been treated either with 2 x 10^{-8} M BrdU for 2 days and then exposed to ME2SO and BrdU, or without pretreatment exposed to ME2SO. In daily intervals, UV-irradiated NDV was added to aliquots of 2 x 10^6 cells, incubation was carried on for 24 h in ME2SO-containing medium with or without BrdU, and cell virus-free supernatants were assayed for interferon. The percentage of cells involved in erythroid differentiation is determined by benzidine staining. The results are presented in Fig. 1. Interferon production increases fourfold on day 3 of the stimulation period. This time coincides with a strong increase in hemoglobin synthesis in these cells. During the late phase of stimulation, the interferon production decreases again to a level below that of unstimulated cells. The optimum of interferon production fluctuated slightly between days 2 and 3. Treatment of cells with BrdU, which inhibits the hemoglobin synthesis (Fig. 1, lower panel), keeps the NDV-induced interferon production on the level of un-
stimulated cells throughout the period of treatment.

The production of interferon after NDV induction in BrdU-resistant B8 cells has been investigated under the following conditions: treatment for 5 days with (i) ME₂SO, 1 to 2%; (ii) BrdU, 0.8 × 10⁻³ M; (iii) ME₂SO, 1 to 2%, and BrdU, 0.8 × 10⁻³ M. UV-inactivated NDV is added at daily intervals to aliquots of cells, and the interferon activity is determined in the cell supernatants 24 h thereafter.

Stimulation with ME₂SO leads to differentiation of more than 60% of the cells in culture and to patterns of interferon synthesis similar to that in ME₂SO-treated F4 cells (Fig. 2A). Interferon production rises to 16,000 U/10⁶ cells on days 3 and 4 of stimulation. Under these conditions of stimulation, no FV induction occurs in the B8 cells (18). When stimulation of hemoglobin synthesis is carried out with 0.8 × 10⁻³ M BrdU in the absence of ME₂SO (Fig. 2b), 40% of cells are benzidine reactive and the overall rate of hemoglobin synthesis is 5% of the soluble cytoplasmic protein fraction (16). Increase over the basic level of interferon production during BrdU stimulation is lower (threefold increase on day 3) than in B8 cells stimulated by ME₂SO. When ME₂SO and BrdU were added together (Fig. 2C), both the interferon inducibility and the hemoglobin synthesis were higher than with either of the stimulating agents alone, indicating an additive effect.

Attempts to induce interferon by poly(I)·poly(C) in F4 and B8 cells at different stages of erythropoietic differentiation have been unsuccessful (Table 1).

![Fig. 1](http://jvi.asm.org/)

**Fig. 1.** NDV induction of interferon in differentiating F4 cells. F4 cells were stimulated by ME₂SO as described, and aliquots of cells were exposed to 0.6 hemagglutinating units of UV-inactivated NDV. After 24 h, interferon activity was determined in the cell supernatants. (Upper panel) Units of interferon produced by 10⁶ cells at different times after ME₂SO stimulation. (Lower panel) Percentage of cells in erythropoietic differentiation as determined by benzidine staining. Symbols: ○, F4 cells stimulated by ME₂SO; ●, F4 cells treated with 2 × 10⁻⁴ M BrdU for 2 days and then exposed to ME₂SO for 6 days in the presence of 2 × 10⁻⁴ M BrdU.

![Fig. 2](http://jvi.asm.org/)

**Fig. 2.** NDV induction of interferon in BrdU-resistant B8 cells during erythropoietic differentiation. Stimulation of differentiation by ME₂SO treatment (A), by treatment with 0.8 × 10⁻³ M BrdU (B), and by treatment with ME₂SO and 0.8 × 10⁻³ M BrdU (C). (Upper panels) Interferon induced by NDV in relation to time of stimulation. (Lower panels) Percentage of cells involved in differentiation as indicated by benzidine staining.
DISCUSSION

The aim of this study is the control of interferon induction in cells undergoing differentiation. In this specific situation differentiation results in the induction of a specialized cellular protein (hemoglobin) and a parallel decrease in DNA, RNA, and overall protein synthesis.

Upon adequate stimulation, interferon is produced as well in highly differentiated cells of reticuloendothelial origin with no capacity of further proliferation, like leukocytes (10, 11), as in cells with a low degree of differentiation and infinite proliferative capacity, like continuous cell lines (4). Spleen cells have been discussed repeatedly as possible in vivo target cells for interferon induction (13, 14). Production of interferon may be viewed as a cellular response to an external stimulus and represents a process analogous to enzyme induction and thus differentiation. Inhibition by actinomycin D of interferon formation (12) is in line with this view. ME₂SO, which induces hemoglobin formation, has no stimulating effect on the interferon production. This implies that induction of cell differentiation and presence of FV are not sufficient for interferon induction.

UV-inactivated NDV induces around 5,000 U of interferon in 10⁴ F4 or B8 cells.

When cells stimulated with ME₂SO for 2 to 3 days are challenged with UV-NDV, the fourfold amount of interferon activity is released. This effect may indicate either a higher potential of differentiating cells to produce interferon or simply an increased receptiveness of ME₂SO-exposed cells to the viral interferon inducer. To distinguish between the two possibilities, ME₂SO stimulation of hemoglobin synthesis has been inhibited in BrdU-sensitive F4 cells by a low dose of BrdU, which does not affect cell proliferation (16). Under these condition the increase in interferon induction is not observed, which indicates that the mere presence of ME₂SO does not facilitate interferon induction but that cells had to evolve in their differentiation.

In BrdU-resistant B8 cells hemoglobin production can be induced by high concentrations of BrdU also in the absence of ME₂SO. The two- to threefold increase of interferon inducibility when cells had been stimulated for 3 days with BrdU gives further support to the concept of linkage between interferon production and the state of cell differentiation. At the final stage of erythropoiesis interferon inducibility regresses.

A special property of these transformed spleen cells seems to be the refractiveness of interferon production mechanism to poly(I)·poly(C) induction. A wide variety of reaction conditions as well as the presence of DEAE-dextran and/or neomycin did not lead to measurable interferon induction.

When differentiating cells are challenged with poly(I)·poly(C) under similar conditions where NDV-mediated induction is elevated, no release of interferon is observed. That these cells remain refractive to the polynucleotide inducer may reflect either a property of the cell membrane (nucleolytic activity) or a fundamental difference in the induction mechanisms by NDV and poly(I)·poly(C) (3), the latter being inactive here.

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LITERATURE CITED


