Insertion of the v-Ha-ras Oncogene Induces Differentiation of Calcitonin-producing Human Small Cell Lung Cancer

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Abstract

Human small cell lung cancers (SCLC) and cell lines derived therefrom are phenotypically heterogeneous concerning neuroendocrine differentiation. Unlike most SCLC tumors and cell lines that express poorly differentiated neuroendocrine phenotypes, the SCLC cell line DMS 53 exhibits mature endocrine differentiation features, including unusually high expression of the gene for the peptide hormone, calcitonin (CT). We now report that introduction of the viral Harvey ras (v-rasH) oncogene into DMS 53 cells via retroviral infection, with resultant constitutive expression, results in increased features of neuroendocrine differentiation. 7–10 d after infection the cells demonstrated altered morphology, increased CT secretion, increased CT gene expression, markedly diminished cellular proliferation, and nearly abolished methylcellulose cloning efficiency. This response of DMS 53 cells to v-rasH is unlike the tumor progression effects we have previously observed in other SCLC lines. Significantly, the differentiation response that follows expression of the virally introduced v-rasH oncogene in DMS 53 cells is similar to that of neoplastic neuroendocrine cell lines derived from adrenal pheochromocytes and thyroid C cells. The effects of constitutive v-rasH expression in DMS 53 SCLC cells and other neuroendocrine cell lines suggest an important role for rasH or related genes in neuroendocrine differentiation.

Introduction

Human small cell lung cancer (SCLC) in cell culture exhibits a range of phenotypes that may reflect both the differentiation state of the bronchial cell of origin and ongoing progression of events during tumor development. Cultures of SCLC hence represent important human cell systems to explore molecular events that regulate phenotypic alterations of normal and neoplastic epithelial cells.

Most SCLC lines, termed classic SCLC, have properties similar to tumors in patients, including high expression of markers typical of early neuroendocrine cell development and a modest degree of endocrine differentiation (1, 2). A significant number of other SCLC lines, characterized as biochemical variant or morphologic variant SCLC, have decreased levels of these same endocrine differentiation markers, increased proliferative indices, amplification of the c-myc gene, and growth patterns and biochemical markers more typical of non-SCLC (1, 2). The SCLC line DMS 53 is unique because it demonstrates more differentiated endocrine features than typically found in classic and variant cells. These include high levels of specific small polypeptide hormones in addition to expression of early neuroendocrine markers (3, 4).

We have been interested in the genetic and biochemical signals that mediate differentiation of these cell phenotypes in the SCLC lineage and recently have shown that when the viral Harvey ras oncogene (v-rasH) is expressed via retroviral infection in variant SCLC, these cells acquire features typical of large cell lung cancer cell lines including adherent growth, increased growth rate, augmented cloning efficiency, resistance to the polyamine analogue α-difluoromethylornithine, and further loss of neuroendocrine markers (5). However, when classic SCLC cells similarly express high levels of v-rasH, their growth characteristics, drug sensitivity, and neuroendocrine markers were not affected (5).

SCLC cells with more mature endocrine features were not initially studied, and hence pose an interesting question with regard to differentiation responses within the SCLC cell lineages. In nonpulmonary neuroendocrine neoplastic cell lines from well-defined endocrine tissues, increasing ras gene expression or microinjection of its p21 protein product results in acquisition of a more differentiated state (6–9). Such results

Figure 1. Northern blot hybridization of 8 μg of total cytoplasmic RNA isolated from uninfected DMS 53 SCLC (lane J), 1504A helper virus–infected DMS 53 SCLC (lane 2), and Ha-MSV-infected DMS 53 SCLC (lane J) probed with (A) an Sst I-Pst I fragment (730 bp) containing only the v-rasH p21 coding region (Oncor) and (B) β-actin. Note high expression of v-rasH in Ha-MSV infected DMS 53 cells (lane J). β-Actin signals demonstrate equivalent loading in each lane.

This paper was presented in part at the International Conference on Hormones, Growth Factors, and Oncogenes in Pulmonary Carcinoma, 30 August–2 September 1987, Hanover, NH, and published in abstract form (1987. Lung Cancer 3:110).

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Received for publication 19 August 1988 and in revised form 28 February 1989.

1. Abbreviations used in this paper: CGRP, calcitonin gene-related peptide; CT, calcitonin; Ha-MSV, Harvey murine sarcoma virus; MTC, medullary thyroid carcinoma cells; SCLC, small cell lung cancers; v-rasH, viral Harvey ras oncogene.

0021-9738/89/07/0194/06 $2.00
Volume 84, July 1989, 194–199
Figure 2. Photomicrographs of uninfected DMS 53 cells (top) and Ha-MSV DMS 53 cells 7 d after infection (bottom). Note morphological alteration from a flat monolayer in uninfected cells (top) to rounded, larger cells growing as mounds of cells (bottom). No morphologic changes were noted in DMS 53 SCLC cells infected with 1504A helper virus alone. ×200.
have been obtained with rat PC 12 pheochromocytoma cells (6–8) and, in our laboratory, with human medullary thyroid carcinoma cells (MTC) (9). SCLC line DMS 53 has phenotypic similarity to MTC in that expression of the calcitonin (CT) gene and resultant CT hormone secretion is extremely high (3, 4). We speculated that the effects of introduction and constitutive expression of \( v-ras^H \) in DMS 53 cells might then differ from the effects of \( v-ras^H \) expression in other SCLC lines, and might instead result in acquisition of differentiation-associated features, such as increased CT gene expression and decreased growth. We now demonstrate that \( v-ras^H \) expression will differentiate DMS 53 SCLC cells. Our findings emphasize further that mutated \( ras \) genes can supply critical functions for altering cell phenotypes in human lung cancer, and that the responses obtained depend on the initial differentiation status of the cell. These data support an important role for \( ras \) or other genes with similar functions in regulating endocrine cell differentiation and show that \( ras \) gene expression in specific types of cultured tumor cells with pronounced neuroendocrine characteristics can cause a decrease in phenotypic features associated with malignancy.

### Methods

**Cell culture and viral infection.** Cells of a well-characterized culture line of human SCLC, DMS 53 (3, 4), were grown at 37°C in Waymouth’s media supplemented with 16% vol/vol of heat-inactivated fetal bovine serum (Gibco Laboratories, Grand Island, NY), 2 mM glutamine, 100 U penicillin, and 100 mg streptomycin sulfate/ml in a humidified CO\(_2\) incubator. The \( v-ras^H \) gene was introduced into DMS 53 cells using our previously described conditions (5, 9–11). Briefly, NIH 3T3 cells, infected with Harvey murine sarcoma virus (Ha-MSV) and 1504A amphotropic murine leukemia helper virus, as well as NIH 3T3 cells infected with 1504A amphotropic murine leukemia helper virus alone, were grown in DME (Gibco Laboratories) supplemented with 9% (vol/vol) heat-inactivated fetal bovine serum. For viral infection, DMS 53 cells were exposed to conditioned medium from the infected NIH 3T3 cells which contained high titers of Ha-MSV and 1504A helper virus or supernatant containing 1504A helper virus alone. Efficiency of infection of DMS 53 SCLC cells was increased by use of Polybrene (Sigma Chemical Co., St. Louis, MO) at 1.0 \( \mu \)g/ml.

**Growth studies.** 10 d after viral infection the cells were harvested by trypsinization and seeded in 25-cm\(^2\) polystyrene flasks at a density of 2.0 \( \times 10^5 \) cells/ml. Cell numbers were determined using an automatic cell counter (Coulter Electronics Inc., Hialeah, FL) and periodically confirmed by hemocytometer counts.

Efficient viral infection and resultant constitutive expression of \( v-ras^H \) occurred after infection of DMS 53 cells with Ha-MSV. Northern blots of total cytoplasmic RNA (8 \( \mu \)g) hybridized with a 730-bp Sst I-Pst I fragment containing only the \( v-ras^H \) coding region showed levels of \( v-ras^H \) mRNA that were 10\(^3\)-fold greater than the levels of endogenous \( c-ras^H \) transcripts detected in uninfected control cells when probed with the above viral probe (Fig. 1) or with a probe for the human \( c-ras^H \) gene (data not shown). Transcripts homologous to the \( v-ras^H \) coding region probe, under our hybridization conditions, were not detected in uninfected control cells nor those infected with helper virus alone.

Within 7 d after infection with Ha-MSV, DMS 53 cells developed marked morphological changes. Cells became more rounded and larger, with increased cytoplasm to nucleus ratio, and had a less pyknotic nuclear pattern with prominent nucleoli. While DMS 53 cells normally grow as a flat confluent monolayer, after Ha-MSV infection they grew as rounded mounds of cells (Fig. 2). No morphologic alterations were observed after infection with the amphotropic helper virus alone.

**Results**

Cloning efficiency. DMS 53 cells infected with Ha-MSV and uninfected DMS 53 controls were suspended at a density of 10,000 cells/ml in 1.8% methylcellulose in growth medium. The methylcellulose/cell suspension was layered over a 3% agar base. Colonies were evaluated and scored at day 10. Cloning efficiency was calculated as the number of colonies per plate divided by the number of cells suspended. The data reported are results of three independent experiments, each containing quintuplicate samples.

**Secretion of calcitonin.** Supernatant from Ha-MSV-infected or uninfected DMS 53 cells was centrifuged (1,500 g, 3 min) and the concentration of CT in the supernatants was determined by RIA (12) and normalized to cell number. Culture media not exposed to cells had no detectable CT levels.

**cDNA probes.** CT (pTT1062) and CT gene-related peptide (CGRP) (pTT42) cDNA probes have been described (13). pTT1062 contains only human CT-specific sequences (exon 4). Neither CGRP-specific sequences (exons 5–6), nor sequences common to both CT and CGRP (exons 1–3), are found in this probe. pTT38 contains a Pvu II-Pst I restriction fragment from pTT42 which contains only CGRP-specific sequences including 17 bp of exon 5 and all of exon 6 of the CT gene (13).

Recombinant plasmids containing human \( \beta \)-actin sequences were provided by D. Cleveland (Johns Hopkins University School of Medicine, Baltimore, MD). An Sst-Pst I fragment (730 bp) containing only the \( v-ras^H \) gene p21 coding region was obtained from Oncor (Gaithersburg, MD). All probes were labeled with \(^{32}P\)dCTP by nick translation to a sp act of ~5 \( \times \) 10\(^8\) cpm/\( \mu\)g of DNA.

**RNA hybridization.** Cytoplasmic RNA was extracted by NP-40 lysis in the presence of vanadyl ribonucleoside complex as previously described (12, 14). 8 \( \mu\)g of total or 4 \( \mu\)g of poly A \(^+\) selected cytoplasmic RNA for Northern blot hybridization analysis was electrophoresed in a 6% formaldehyde/1.2% agarose gel and transferred to nitrocellulose. Filters were hybridized in 5 \( \times \) SSPE (1 \( \times \) SSPE = 150 mM NaCl, 10 mM Na phosphate, 1 mM EDTA), 10% formamide, 1% SDS, 60 mg/ml polyethylene glycol 8,000, 5 mg/ml powdered milk, and 200 mg/ml sonicated salmon sperm DNA for 16 h at 42°C. After hybridization filters received a final wash with 0.1% standard saline citrate, pH 7.0/0.1% SDS at 65°C for 30 min. For autoradiography, films were exposed for 24–36 h at ~70°C. Hybridizations and quantitation of signal for CT and \( \beta \)-actin RNAs by densitometry were performed as described (12, 13).
Figure 4. Photomicrographs of methylcellulose/agar plates of uninfected DMS 53 SCLC cells (A) and Ha-MSV-infected DMS 53 SCLC cells (B). Note the presence of multiple large clones of cells in DMS 53 SCLC (A). In contrast, Ha-MSV-infected DMS 53 SCLC (B) shows no clones present. DMS 53 SCLC cells infected with Ha-MSV removed from the methylcellulose suspensions remained viable as assessed by trypan blue dye exclusion. Clones were counted at day 10.
with Ha-MSV. Cell doubling times were increased from 36 to 70 h (Fig. 3). Decreased growth was accompanied by marked reduction in cloning efficiency. Cloning efficiency in methylcellulose-agar was reduced from 8.2 to 0.2% after induction of the v-rasH expression (Fig. 4). DMS 53 cells expressing v-rasH remained viable as assessed by trypan blue staining after their removal from the methylcellulose suspensions.

Decreased growth often accompanies cell differentiation. Our previous studies of human MTC (9) demonstrated that v-rasH infection resulted in diminished growth and induction of a program of differentiation that included a marked increase in expression of the neuroendocrine peptide hormone gene, CT. DMS 53 SCLC cells, which also produce high basal levels of CT, might be expected to increase CT production in response to differentiation. Fig. 5 shows that increased CT secretion occurs after infection with Ha-MSV. Consistent with this product increase, the Ha-MSV-infected cells had increased expression of CT mRNA (Fig. 6). It is interesting that the increase involves only one of the two major mRNA species generated by alternative processing of the CT gene. One of these species, CT mRNA, contains exons 1, 2, 3, and 4 and is elevated threefold. However, the other major mRNA species from the CT gene, which encodes CGRP, and which is derived from exons 1, 2, 3, 5, and 6 (13, 15), did not increase after expression of the virally introduced v-rasH oncogene (Fig. 6).

Discussion

In many cell types, ras gene overexpression or mutation is associated with neoplastic transformation or progression (5, 16–23). However, previous studies in two well-defined neuroendocrine tumor cell lines, rat pheochromocytoma cell line PC 12 (6–8) and human MTC (9), demonstrated that expression of an exogenously introduced ras gene or gene product can induce features of neuroendocrine differentiation. We now show that neoplastic endocrine cells that arise from the complex background of the bronchial epithelium, and which are putative progenitors of the cancer SCLC, differentiate in a fashion similar to human MTC and PC 12 after expression of the virally introduced v-rasH oncogene. We observed acquisition of differentiation-associated features which include a decrease of such tumorigenic properties as the ability of the cells to clone, and a reduced cell growth fraction. In contrast, no phenotypic alterations were observed in DMS 53 SCLC cells solely infected with 1504A amphotrophic helper virus alone. These data suggest that acquisition of differentiation-associated features in response to ras expression may be specific for cells of distinct neuroendocrine phenotypes.

Our results in cell culture appear to correlate with recent observations in vivo regarding ras gene expression and neuroendocrine cell differentiation. For example, high levels of the ras gene protein product p21 have been demonstrated by immunohistochemistry in differentiated neural or neuroendocrine tissue (24). Moreover, increased expression of rasH protein product p21 predicts favorable prognosis in patients with neuroblastoma (25). These results suggest that ras or related genes may have an important role in maintenance of neuroendocrine phenotypes.

The data obtained from the DMS 53 SCLC cell line are especially significant when viewed in the context of our previous work with v-rasH and SCLC (5). SCLC cell lines with partially developed neuroendocrine features, termed classic SCLC, demonstrated no phenotypic alterations of growth, morphology, biochemical markers, or drug sensitivity after the virally introduced v-rasH gene was expressed. Biochemical variant SCLC acquired non-SCLC features and demonstrated features of tumor progression (5). Thus, the response to v-rasH expression seen in SCLC cells appears to be dependent on the phenotypic context of the recipient cell.

The molecular mechanisms by which the ras gene influences endocrine cell differentiation and the behavior of lung cancer cells may relate to the functions of the ras gene products, which may mediate signal transduction events. Our data with DMS 53 SCLC, which shares neuroendocrine features with MTC and PC 12 cells, suggests that ras-mediated transduction events may replace functions lost after transformation of pulmonary endocrine cells. The differential phenotypic responses in SCLC lines after v-rasH expression may reflect the different responses that normal bronchial epithelial cells may exhibit in response to expression of ras or related genes in vivo during cell renewal and differentiation. Elucidation of the specific genetic and biochemical events responsible for the differential responses to expression of virally introduced v-rasH in different SCLC cell lines should provide important information regarding the mechanisms through which alterations of genes such as rasH may promote or interfere with normal differentiation processes. The model systems offered by cultured human lung cancer cells with differential phenotypic responses to the rasH gene provide unique means to pursue these questions.
Acknowledgments

We thank Dr. Paul Celano, Dr. Robert Casero, Dr. Andree de Bustros, and Dr. Nancy Davidson for helpful discussions, Dr. Alan Rein for providing Ha-MSV and helper virus, Dr. Don Cleveland for the human β-actin probe, Ms. Sara Lyles for CT R1As, Ms. Kathleen Wieman for making the figures, and Ms. Sandra Lunds and Ms. Tammy Hess for typing the manuscript.

This work was partly supported by Training Grant NRSA 1532-CA08207-02 (to Dr. Mabry) and R01-CA48081-01 from the National Institutes of Health.

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