Autocrine growth induced by transferrin-like substance in bladder carcinoma cells

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Summary Ample evidence confirms that certain cancer cells have the capacity to produce multiple peptides as growth factors and that expression of their receptor may act in tumour cell paracrine and/or autocrine loop mechanisms, either by extracellular release of the growth factor or by the tumour itself. To study the possibility of an autocrine growth mechanism in bladder carcinoma, we investigated the ability of various bladder carcinoma cell lines to proliferate in serum-free medium. A rat bladder carcinoma cell line, BC47, demonstrated exponential and density-dependent growth in serum-free medium. Furthermore, conditioned medium from BC47 cells induced growth-stimulating activity for BC47 cells themselves. Purification and further characterization of this activity was performed by chromatographic methods, SDS-PAGE and N-terminal amino acid analysis. Finally, we have identified that a transferrin-like 70-kDa protein is found to be the main growth-promoting factor in this conditioned medium. In addition, specific antibodies against transferrin and the transferrin-receptor inhibit the in vitro growth of this cell line. Our data suggest that this transferrin-like factor possibly acts as an autocrine growth factor for cancer cells.

Keywords: autocrine growth factor; bladder carcinoma; rat bladder carcinoma cell line (BC47); serum-free culture; transferrin

The relatively autonomous nature of malignant cells has been known for many years, i.e. they require fewer exogeneous growth factors for optimal growth than do their counterparts. To explain this phenomenon, it has been suggested that cells could become malignant by the endogeneous production of polypeptide growth factors acting on their producer cells via functional external receptors, allowing phenotypic expression of the peptide by the same cell that produces it. This process has been termed ‘autocrine secretion’ (Sporn and Todaro, 1980; Sporn and Roberts, 1985). There is now much circumstantial and direct evidence to support the original hypothesis. Many types of tumour cells release polypeptide growth factors into their conditioned medium when grown in cell culture, and these same tumour cells often possess functional receptors for the released peptide.

The presence of either serum components or substances such as hormones and growth factors is essential to maintain cells in culture. In analysing the regulatory mechanism of growth factors, the importance of cells that can proliferate in chemically defined, serum-free medium without supplements is increasing, because such cells are likely to synthesize factors with growth-stimulating activity (Messing et al., 1984; Matsuda et al., 1989) that can be isolated easily in the absence of extrinsic growth factors. Recently, it was demonstrated that bladder carcinoma cells were able to secrete a variety of autocrine factors, such as tumour-derived adhesion factor (Akaogi et al., 1994) and granulocyte colony-stimulating factor (Tachibana et al., 1995). The present study was undertaken to investigate the ability of various bladder carcinoma cell lines to proliferate in serum-free medium, the purification of growth-stimulating activity in conditioned medium and the possibility of an autocrine growth mechanism in bladder carcinoma.

MATERIALS AND METHODS

Cell lines and culture conditions

Four human bladder carcinoma cell lines (KU1, KU7, T24 and NBT) were used (Tachibana, 1982). A rat bladder carcinoma cell line, BC47, which was induced in inbred ACI/N rats by exposure to N-buty1-N-butanol (4) nitrosamine, was also used. All cell lines were maintained in minimal essential medium (MEM) with 10% fetal bovine serum. Cells were trypsinized, washed three times with Ca2+- and Mg2+-free Hanks’ balanced salt solution, counted and then plated into 25-cm2 tissue culture flasks at densities ranging from 1×104 to 3×105 cells cm-2 in 5 ml of serum-free medium. The medium consisted of a 1:1 mixture of Dulbecco’s modified Eagle medium (DMEM)/Ham’s F-12 medium containing hydrocortisone (50 nM), prostaglandin E1 (25 ng ml-1), 3,3’,5’–triiodo-L-thyronine (5 pm), L-glutamine (0.292 mg ml-1) and sodium selenite (10 nM) with reference to the report of Messing et al. (1982). The cells were incubated at 37°C in a humidified atmosphere of 5% carbon dioxide and 95% air. Cells were removed from the flasks by trypsinization, and viable cells (trypan blue dye exclusion) were counted in a haemocytometer (Bürker-Türk) on days 1–5. The medium was not removed and cells were not refreshed during the experiments.

Collection and concentration of conditioned medium

Conditioned medium was collected from ongoing cultures of BC47 cells growing in serum-free medium, when they reached 90% confluence, was centrifuged at 4000 r.p.m. for 10 min to remove cellular debris and was stored at –80°C. When a sufficient quantity had been collected, it was thawed, pooled and concentrated using an ultrafiltration cell (model 8050, Amicon, WR Grace, Danvers, MA, USA) and a filter with a 5000 molecular weight cut-off. Specimens were sterilized by passage through a membrane filter with 0.45-μm pores and were stored at –80°C until use.

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Cell growth in concentrated serum-free conditioned medium

BC47 cells were plated at $1 \times 10^4$ cells cm$^{-2}$ in serum-free medium supplemented with fivefold-concentrated BC47-conditioned medium (10% of the final volume) in 25-cm$^2$ tissue culture flasks. Growth curves were obtained as described above.

Effect of conditioned media collected from other bladder carcinoma cell lines on growth of BC47 cells in serum-free medium

We investigated whether conditioned media collected from NBT, T24 and KU1 cell lines could stimulate the growth of BC47 cells in serum-free medium. Cell growth was assayed by thymidine incorporation assay as described in the section entitled Proliferation assay.

Effect of medium replenishment on growth of BC47 cells in serum-free medium

BC47 cells ($3 \times 10^4$ cells cm$^{-2}$) were plated into serum-free medium in 25-cm$^2$ tissue culture flasks and were incubated at 37°C. The medium was replaced with fresh medium at 24 h intervals. Sham treatment by removing and returning the medium was also done at 24 h intervals (Masuda et al, 1988). Cell counting was performed as described above.

Gel filtration of conditioned medium

Two hundred microlitres of the 50-fold concentrated BC47-conditioned medium was applied to a $1.0 \times 30$-cm Superose 12 precolumn (Pharmacia LKB Biotechnology, Uppsala, Sweden) equilibrated with 0.05 M Tris-HCl (pH 8.6). The column was eluted at room temperature with 0.05 M Tris-HCl (pH 8.6) at a flow rate of 30 ml h$^{-1}$, and the absorbance of the eluate was monitored at 280 nm. Two-millilitre fractions were collected, sterilized by passage through a membrane filter with 0.22-µm pores and stored at −80°C until assay for biological activity.

Proliferation assay

BC47 cells ($3.2 \times 10^4$ cells per well) were dispersed into 96-well round-bottom plates with 175 µl of serum-free medium and were incubated for 24 h at 37°C in a humidified atmosphere of 5% carbon dioxide and 95% air. Then 25 µl of [3H]thymidine (20 µCi ml$^{-1}$) was added to each well immediately after the addition of 25 µl aliquots of the fractions obtained by gel filtration, and culture was continued for a further 6 h. Subsequently, the medium was removed, the cells were harvested by cell harvester, and the radioactivity was analysed by a liquid scintillation counter (Beckman LS 9800; Beckman Instruments, Fullerton, CA, USA). Results are expressed as a percentage of the c.p.m. obtained in control culture.

Anion exchange chromatography

After using ultrafiltration to concentrate the biologically active fractions obtained by gel filtration, the retentate was applied to a 1.6 x 10 cm HiLoad Q Sepharose HP precolumn (Pharmacia) equilibrated with 0.05 M Tris-HCl (pH 8.6) at room temperature. Bound material was eluted with a continuous 130-ml linear 0–0.5 M sodium chloride gradient in 0.05 M Tris-HCl (pH 8.6), and the eluate was monitored at 280 nm. Five-millilitre fractions were collected at a flow rate of 150 ml h$^{-1}$ at room temperature, sterilized by passage through a membrane filter with 0.22 µm pores and assayed for biological activity as described above.

SDS-PAGE

The most active fraction prepared by gel filtration and anion exchange chromatography, together with the fractions before and behind the active one, were separated by SDS-PAGE as reported previously (Laemmli, 1970). Samples were heated at 95°C for 3 min with 1.5% dithiothreitol before application to a 7.5% polyacrylamide gel. The resulting gel was stained by silver staining kit (Pharmacia). Molecular weight markers were used for protein markers (Pharmacia) with phosphorylase-b (94 kDa), albumin (67 kDa), ovalbumin (43 kDa), carbonic anhydrase (30 kDa), trypsin inhibitor (20 kDa) and α-lactalbumin (14 kDa).

Amino acid analysis

After SDS-PAGE under reducing conditions, the protein bands were transferred to a polyvinylidene difluoride membrane. The amino (N)-terminal amino acid sequence of the 70-kDa protein band that corresponded to the biological activity was analysed with a gas-phase protein sequencer (PPSQ-10; Shimadzu, Kyoto, Japan). Its amino acid sequence analysis was performed by Tokyo Research Laboratories, Kyowa Hakko Kogyo (Tokyo, Japan).

Antibody studies

For the antibody studies, the mouse anti-human transferrin monoclonal antibody (Cosmo Bio, Tokyo, Japan) was used. This antibody is a mouse IgG molecule that has been shown to bind human transferrin specifically. BC47 cells were plated at $1 \times 10^4$ cells in 5 ml of serum-free medium added at various concentrations of antibodies (5 µg ml$^{-1}$, 10 µg ml$^{-1}$ and 20 µg ml$^{-1}$) in 25-cm$^2$ tissue culture flask. The cells were incubated at 37°C in a humidified atmosphere of 5% carbon dioxide and 95% air. Twenty-four hours and 48 h later, bromodeoxyuridine was added at a final concentration of 30 µg ml$^{-1}$ per flask, and the cells were reincubated for another hours. After bromodeoxyuridine labelling, the cells were trypsinized, washed three times with phosphate-buffered saline, and DNA-synthesizing cells were determined by the DNA-bromodeoxyuridine double-staining method using flow cytometry as reported previously (Tachibana et al, 1991). The same experiment was made with various amounts (5 µg ml$^{-1}$, 15 µg ml$^{-1}$ and 30 µg ml$^{-1}$) of the mouse anti-human transferrin receptor monoclonal antibody (Oncogene Science, NY, USA). Also, we investigated the effect of anti-human transferrin antibody on the growth-stimulating activity in BC47-conditioned medium. BC47 cells were plated at $1 \times 10^4$ cells cm$^{-2}$ in serum-free medium supplemented with both fivefold-concentrated BC47-conditioned medium and anti-human transferrin antibody (20 µg ml$^{-1}$) in 25-cm$^2$ tissue culture flasks. As a control, anti-human IgG antibody was used. To eliminate the possibility that the anti-human transferrin antibody was toxic to the cells, the same experiment was performed with KU7 cells that could grow well in serum-free medium, independent of plating cell density. Growth curves were obtained as described above.
RESULTS

Growth of bladder carcinoma cell lines in serum-free medium

Five bladder carcinoma cell lines were cultured in serum-free medium (Figure 1). As demonstrated by cell counts after a 24 h incubation period, the plating efficiency of three cell lines, with the exception of KU1 and BC47, were considerably reduced in serum-free medium. KU1 cells could survive without proliferation until 48 h, however, it could not be cultivated thereafter. On the other hand, BC47 cells grew well when plated at a density of $3 \times 10^4$ cells cm$^{-2}$, but this cell line did not grow when plated at a lower density (Figure 2). Thus, it seems likely that the growth of BC47 cells in this serum-free medium was dependent on the plating cell density. Conversely, KU7 cells demonstrated vigorous growth in serum-free medium, regardless of plating cell density. T24 cells survived without proliferation during an experiment, while NBT cells could not be cultivated in serum-free medium. Fivefold-concentrated BC47-conditioned medium was found to stimulate the proliferation of BC47 cells plated in serum-free medium at $1 \times 10^5$ cells cm$^{-2}$ (Figure 2). However, conditioned media from the other human bladder carcinoma cell lines NBT, T24 and KU1 did not exhibit a growth-stimulating activity on BC47 cell comparable to the effect of conditioned medium collected from BC47 (Figure 3). Moreover, when compared with control cultures, a significant decrease in cell numbers was observed by changing the medium at 24-h intervals during incubation for a period of 72 h (Table 1). These experiments suggest that growth-stimulating activity of conditioned medium collected from BC47 is not derived from a pre-existing substance in serum-free medium but is secreted by only BC47 cells growing in serum-free culture.

Partial purification of growth-stimulating activity produced by BC47 cells

When concentrated BC47-conditioned medium was separated by gel filtration, the peak of growth-stimulating activity was observed in the high-molecular-weight fraction (Figure 4). In the anion-exchange chromatography, growth-stimulating activity was bound to the column at pH 8.6 and was eluted from the column by approximately 0.26 M sodium chloride (Figure 5). The lower-molecular-weight fractions with a minor activity by gel filtration were also

Table 1 Effect of medium change on the growth of BC47 cells in serum-free medium

| No. of cells per flask after a 72-h incubation period ($\times 10^4$ cells) |
|------------------|------------------|
| Control          | 2.42 ± 0.10      |
| Sham treatment   | 2.19 ± 0.08      |
| Medium change    | 1.80 ± 0.21      |

BC47 cells were plated at $3 \times 10^4$ cells cm$^{-2}$ tissue culture flasks. The medium was replaced with fresh medium at 24-h intervals. Sham treatment by removing the medium and then returning it was performed at 24-h intervals. All studies were performed in duplicate. Number of cells were expressed as mean ± s.d. *Not significant; **P < 0.01 compared with the control.
analysed in the same manner; however, no growth-stimulating activity could be detected. The biologically active fraction obtained from gel filtration was applied to SDS-PAGE. As demonstrated in Figure 6, a light band at 70 kDa, which was close to the major dark band, was seen. When the active fraction from anion-exchange chromatography was also applied to SDS-PAGE, this 70-kDa band comigrated with the dark band. However, the 70-kDa band was considered to correspond to growth-stimulating activity because of an increase in its staining intensity according to purification.

**Structural analysis of the purified 70-kDa protein**

In order to examine the relationship between the purified 70-kDa protein and other proteins, its N-terminal amino acid sequence was analysed with an automated sequencer. The N-terminal sequence up to the 17th amino acid residue was determined to be S-A-G-W-N-I-P-I-G-L-L-Y-X-D-L-P-E., in which X seemed to be C (Cys). As a result of an investigation of the homology using the protein database, this sequence was found to be completely identical to the sequence of amino acid residues 125–141 from the N-terminus of human transferrin (MacGillivray et al, 1982; Yang et al; 1984).

**Inhibition of transferrin-like growth-stimulating activity by antibodies against human transferrin and transferrin receptor**

To investigate the possibility of an autocrine mechanism in this system, monoclonal antibodies against human transferrin and transferrin receptor were tested for their effect on the proliferation of BC47 cells. Figure 7 showed that a decrease in the proportion of cells in S-phase was seen in a dose-dependent fashion by adding the anti-human transferrin antibody. Also, as shown in Figure 8, the decrease in the number of cells in S-phase of the cell cycle was manifested by adding the anti-human transferrin receptor antibody. As demonstrated previously, BC47-conditioned medium stimulated the proliferation of BC47 cells plated at $1 \times 10^4$ cells.
The growth of BC47 cells in serum-free medium demonstrated considerable dependence upon a high cell density. This was presumably not only because of technical factors such as the reduced plating efficiency but also because of an inherent density-dependence of the growth of BC47 cells in serum-free medium. Also, the number of proliferating BC47 cells decreased by periodic medium exchange. This phenomenon has been explained by the production of molecules that have a mitogenic effect and are transmitted between neighbouring cells. Cell proliferation apparently occurs when these substances reach a sufficient concentration to be stimulatory when bound to the target cells. The supernatant of growing cell cultures is therefore likely to contain not only growth-promoting factors (Cross and Dexter, 1991) but also various secretory factors (Lioatta et al, 1986; Koshikawa et al, 1992; Akaogi et al, 1994). The present study also showed that culture of BC47 cells in serum-free conditions provided a suitable system for investigating such growth factors. We have shown that BC47 cells are able to secrete a 70-kDa molecule, which has growth-stimulating activity on BC47 cells themselves, into the chemically defined synthetic medium without transferrin. In addition, the N-terminal amino acid sequence of this molecule is homologous to the sequence of amino acid residues 125–141 of human transferrin, except for lacking amino acid residues 1–124 from N-terminus. BC47 is a rat bladder carcinoma cell line; however, strong homology is claimed to exist between the amino acid sequence of human and non-human transferrins (Bowman et al, 1988). Consequently, the bioactivity of human and rat transferrin is thought to be very similar, and we used monoclonal antibody to human transferrin and transferrin receptor in the blocking experiment.

Transferrin, the major iron-transporting protein in plasma, is a glycoprotein with a molecular weight of 80 kDa (MacGillivray et al, 1982). It transports ferric iron from the intestine, reticuloendothelial system and liver parenchymal cells to all proliferating cells in the body. Previous experiments strongly indicate that the only function of transferrin in supporting cell proliferation is supplying cells with iron. Numerous in vitro studies have demonstrated the requirement of transferrin for proliferation and/or DNA synthesis, and thus transferrin has been included among the factors required for cell growth in serum-free media (Barnes and Sato, 1980). As discussed on previous reports (Shewale and Brew, 1982; MacGillivray et al, 1983), chemical and physical evidence indicates that the ferric iron bound at each site in transferrin is liganded with 1 or 2 histidyl residues, 2 or 3 tyrosines and a bircarbonato ion that probably interacts with an arginyl side chain. In the case of metal-binding protein and enzymes, in which the locations of binding site residues have been determined, two metal-liganding residues have been found to be no more than four residues apart in the amino acid sequence (Liljas and Rossman, 1974), presumably to form an initial locus for weak association before the formation of the complete binding site by conformational adjustments. On this basis, the only two candidates for components of the binding sites that are sufficiently close in the sequence are considered to be tyrosines 185 and 188 in the N-terminal domain and 514 and 517 in the carboxy-terminal domain of transferrin (Schwale and Brew, 1982). The 'transferrin-like' molecule in serum-free medium conditioned by BC47 cells is likely to contain the above sequence of amino acids; therefore, this molecule is considered to act as an iron-carrier protein similar to native transferrin.

Another line of investigation revealed that transferrin may serve a role as a growth factor, independent of its function as a transporter of iron (May and Cuatrecasas, 1985). Transferrin is considered to be an

**DISCUSSION**

Serum has been used in nutritive media to supply to cultured cells a growth-stimulating activity with a complex, undefined and variable nature. The substitution of serum by standard media supplemented with nutrients and hormones of known composition and concentration simplifies the investigation of all classes of growth factors. Growth factors play a central role in cell transformation (Heldin and Westermark, 1984). In general, as tumours progress to a more malignant phenotype, they become less dependent on serum-derived growth factors for their growth in vitro and begin producing polypeptide growth factors (Rodeck and Herlyn, 1991), suggesting that autocrine growth mechanisms may be involved in malignant transformation. In several cell lines, it has been reported that the production of autocrine growth factors does occur. It has been shown that tumour cell growth can be regulated by normal growth factors, and the products of many viral and cellular onco- genes have been found to be related to growth factors or growth factor receptors (Wong and Passaro, 1989).
Figure 7  DNA and bromodeoxyuridine (BrdU) two-colour analysis of BC47 cells with anti-transferrin antibody by flow cytometer. Bivariate bromodeoxyuridine/DNA (green/red) fluorescence distributions are displayed in a dot-density graph with DNA distribution on axis of abscissa and bromodeoxyuridine fluorescence on axis of ordinate. BC47 cells were plated at $1 \times 10^6$ cells in 5 ml of medium in 25-cm² tissue culture flasks and various concentrations of antibodies were added: (A) none, (B) 5 µg ml⁻¹, (C) 10 µg ml⁻¹, (D) 20 µg ml⁻¹. The figures in each dot graph indicate the percentage of cells with positive staining.
Figure 8 DNA and bromodeoxyuridine (BrdU) two-colour analysis of BC47 cells with anti-transferrin receptor antibody by flow cytometer. Bivariate bromodeoxyuridine/DNA (green/red) fluorescence distributions are displayed in the same dot-density graph as Figure 7. BC47 cells were plated at 1 x 10⁶ cells in 5 ml of medium in 25-cm² tissue culture flasks. Various amounts of antibodies were added. (A) None, (B) 5 μg ml⁻¹, (C) 15 μg ml⁻¹, (D) 30 μg ml⁻¹. The figures in each dot graph indicate the percentage of cells with positive staining.

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The symbols indicate cell lines from: with conditioned (20 10⁴ autocrine actively growing, such as cancer that BC47 dependently themselves.

Growth factors (Trowbridge and Omary, 1981), however, and we confirmed it to be much evidence growth factors and presented at 10⁶ Jg anti-human ml-'). It can be demonstrated that the number of human bladder carcinomas to synthesize transferrin and/or a 'transferrin-like' substance might provide a source of available iron to support localized proliferation of bladder carcinoma cells in vivo in areas not well vascularized. In addition, the additional identification of such autocrine growth factors as the 'transferrin-like' substance and/or transferrin could lead to new therapeutic strategies for the treatment of bladder carcinoma in the future.

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