A human natural antibody to adenocarcinoma that inhibits tumour cell migration

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Summary We characterized a natural human antibody to adenocarcinomas and investigated the biological role of this Ab/Ag complex in cancer expansion. Human monoclonal antibodies (HuMAbs) were generated with hybridoma fusion methods using regional nodal lymphocytes of colon carcinoma patients. Among 1036 HuMAbs, only one, termed SK1, an IgM, was adenocarcinoma specific in the immunohistochemical study. The antigen recognized by SK1 (Ag-SK1) was a glycoprotein with a molecular weight of 42–46 kDa. The expression of Ag-SK1 on carcinoma cells varied according to the cell growth periods but was independent of cell cycle stage as elucidated by two-colour fluorescence-activated cell sorter (FACS) analysis. A dot-blots analysis showed that the concentration of Ag-SK1 per total protein differed considerably among eight colon carcinoma cells examined and that the difference was closely correlated with the invasion capacity of the cells as assessed by a microchemotaxis assay. Furthermore, up to 87% of cell migration was inhibited by SK1 in a dose-dependent manner. These data suggested that Ag-SK1 is metabolized and expressed on highly invasive carcinoma cells. In addition, it appears that, although rare, some patients do mount an anti-cancer antigen response in their draining lymph nodes. A HuMAb such as SK1 may be a good candidate for the treatment of cancer invasion and metastasis.

Keywords: human monoclonal antibody; natural antibody; colon cancer; invasion; immunotherapy

MATERIALS AND METHODS

Cell lines

Carcinoma cell lines were all purchased from ATCC (Rockville, MD, USA). A human lymphoblastoid cell line fusion partner, sHFP-1, was grown and used as described previously (Glassy, 1989).

Hybridoma generation

Eighteen regional draining lymph nodes from six Dukes’ B colon carcinoma patients were taken aseptically from resected specimens. Nodal lymphocytes were prepared as a single-cell suspension by mechanical dispersion with scalpels as described previously (Glassy, 1987). Approximately 5.0 × 10⁶ to 1.0 × 10⁷ lymphocytes from each lymph node were individually stimulated with a cytokine mixture (Borrebaeck et al. 1988) for 6–7 days following an ‘in vitro stimulation’ procedure (Koda and Glassy, 1990). This cytokine mixture was reported to contain a considerable amount of interleukin (IL)-2 and interferon (IFN)-gamma (Borrebaeck, 1987). After stimulation, one part of the lymphocytes was mixed with two parts of the human lymphoblastoid cell line sHFP-1 and washed with serum-free RPMI-1640 three times. Ordinary cell fusions were individually performed for each lymph node with 50% polyethylene glycol 1500 (Boehringer Mannheim, Germany) as described (Glassy et al. 1983). After the fusions, cells were resuspended in RPMI-1640 medium supplemented with 10% fetal calf serum, glutamine and 0.2 mM hypoxanthine/0.2 μM amethopterin/32 μM thymidine (HAT: Sigma, St Louis, MO, USA) at 5 × 10⁶ cells ml⁻¹. Cells were then plated at 1.0 × 10⁵ per well in 96-well microtitre plates without the use of feeder layer cells. Usually, 2–4 plates were prepared from one lymph node.

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EIA analysis

Approximately 2–5 weeks after the fusion, the wells were screened microscopically for hybridoma growth. The supernatants of wells showing positive hybridoma growth were first assayed for producing human immunoglobulin by conventional ELISA, using a 96-well assay plate precoated with goat anti-human IgM or IgG (Jackson ImmunoResearch, West Grove, PA, USA). The immunoreactive huMABS were then identified by cell-EIA on a panel of cell lines immobilized on Falcon flexible assay plates (Glassy and Suri, 1985). Briefly, logarithmic phase cells were collected with 0.02% EDTA in calcium- and magnesium-free Dulbecco–phosphate-buffered saline (PBS) (D-PBS), washed twice in D-PBS, resuspended and aliquoted at 2 × 10⁶ cells per well of flat-bottom 96-well plates. These plates were then placed overnight in a 37°C drying oven and then stored until use. After blocking the plate with 0.5% bovine serum albumin (BSA)/D-PBS, 50 ml of test supernatants were added to the wells, incubated for 1 h at room temperature. Washed three times with 0.5% Tween 20/D-PBS and incubated for an additional 30 min with an affinity-purified horseradish peroxidase (HRP)-conjugated goat anti-human IgM (Jackson ImmunoResearch). Following washings, the plates were developed with orthophenylenediamine (OPD) plus hydrogen peroxide in citrate buffer. pH 5.0. Reactions were stopped with 2.5 m sulphuric acid solution, and were read at 492 nm with an enzyme immunoassay (EIA) reader. The positive huMABS were subjected to immunostaining of several human cell lines to eliminate the false-positive clones inherent in cell EIA (Koda et al. 1990).

Purification of huMAB SK1

To stabilize a clone secreting a large amount of immunoglobulin, the hybridoma secreting huMAB SK1 was cloned by four independent series of limiting dilutions (one cell per well). The best clone recovered secreted approximately 1.2–1.5 μg ml⁻¹ 1.0 × 10⁶ 24 h⁻¹ for at least 2 months in continuous culture and doubled in quantity every 20–26 h. The hybridoma was cultured in serum-free medium, S-Clone (Sanko Junyaku, Tokyo, Japan); and 101 of culture supernatant was collected. Proteins were then precipitated by the salting-out method using 45% saturated ammonium sulphate. The precipitate was then dialysed thoroughly against Ca²⁺ and Mg²⁺ free D-PBS. The solution was then passed through an affinity column in which goat anti-human IgM monoclonal antibody was affinity to CNBr-Sepharose 4B (Pharmacia, Tokyo, Japan) at 1 mg g⁻¹ gel. After washing the column with Ca²⁺ and Mg²⁺ free D-PBS, human IgM was eluted with 6 M guanidine hydrochloride. The effluent was immediately dialysed against D-PBS, and the concentration of human IgM was adjusted to 100 μg ml⁻¹ with D-PBS supplemented with 1% human albumin.

Antigen characterization

One half of the target cells immobilized on to assay plates were treated with several enzymes to estimate the antigen nature recognized by the huMABS. Periodate treatment was performed with 5 mM periodic acid (Sigma) in sodium acetate buffer, pH 4.5, at room temperature for 45 min. Neuraminidase (Sigma) was diluted to 1 unit ml⁻¹ with citrate phosphate buffer, pH 5.0, and incubated with cells overnight at 37°C. Trypsin at 0.1% concentration, pH 7.6, was incubated with cells at 37°C for 20 min. After washing, ordinary cell EIA was performed, and the alterations in reactivity, if any, were evaluated.

Invasion assay

The invasion capacity of colon carcinoma cell lines was determined using a 24-well microchemotaxis chamber (Kurashiki Boseki, Okayama, Japan) in quadruplicate assay (Aznavoorian et al. 1990). A polycarbonate membrane with an 8-μm pore size was precoated with an extracellular matrix mixture in D-PBS (40 μg ml⁻¹ of type 4 collagen, 10 μg ml⁻¹ fibronectin and 5 μg ml⁻¹ laminin, all from Cosmo Bio, Tokyo, Japan) by soaking overnight in this mixture at 4°C. After removal of the solution, RPMI-1640 supplemented with 10% fetal calf serum (FCS) was added to the unit. In the upper chamber, 10⁵ cells per well were seeded, whereas, in the lower chamber, 5 μg ml⁻¹ laminin, fibronectin and type 4 collagen were added as chemoattractants. After 36 h of incubation at 37°C in a 5% carbon dioxide humidified incubator, the membranes were fixed with cold acetone for 8 min and stained with Meyer’s haematoxylin. The number of cells that migrated to the lower side of the membrane was microscopically counted and evaluated. In some experiments either purified huMAB SK1 or irrelevant human IgM MAb was added to the upper chamber at various concentrations to inhibit the cell migration with antibodies.

Immunohistochemical staining

A panel of frozen sections (4 μm thick) of human tissues was stained with huMAB SK1 using a standard avidin–biotin peroxidase complex (ABC) method. When deparaffinized formalin-fixed sections were used, they were soaked in 0.1 M citrate buffer (pH 6.0) and treated with microwaves twice for 10 min as described previously (Shi et al. 1991; van den Berg et al. 1993). Frozen sections were fixed with cold acetone for 8 min before the blocking procedures. Sections were first treated with 0.03% hydrogen peroxidase in methanol for 30 min, washed, then blocked with 1% BSA/D-PBS plus 5 μg ml⁻¹ of goat anti-human IgM (Jackson ImmunoResearch) at room temperature for 1 h. After washing with D-PBS, huMAB SK1 diluted to 5 μg ml⁻¹ with 0.5% BSA/D-PBS was applied and incubated overnight in the moist chamber at 4°C. After washing, the sections were incubated with biotinylated goat anti-human IgM (Jackson ImmunoResearch) for 1 h at room temperature. They were then washed and incubated with avidin–biotinylated peroxidase complex (ABC staining kit from Dako Japan, Tokyo, Japan) for 1 h at room temperature. After several more washings, slides were reacted with diaminobenzidine substrate for 10 min, rinsed, counterstained with haematoxylin and mounted.

Flow cytometry

Single cell suspensions of colon carcinoma cell lines were treated with 3.7% formaldehyde solution for 10 min at room temperature. After removal of the solution, cold acetone was added and incubated on ice for 5 min, washed, then incubated in 5% FCS/D-PBS for 1 h. Following washings, SK1 or human polyclonal IgM (each at 5 μg ml⁻¹) was applied and incubated on a shaker for 1 h at room temperature, washed, then appropriately diluted FITC-conjugated goat anti-human IgM (Jackson ImmunoResearch) was added and incubated for an additional 30 min at room temperature.
After washing, they were passed through a nylon mesh (200 inch-1 mesh), stained with propidium iodide (PI; 20 µg ml-1, Sigma) and then analysed with FACScan (Becton Dickinson Japan, Tokyo, Japan). The signal from fluorescein isothiocyanate (FITC) corresponding to Ag-SK1 levels was measured with a 530-nm bandpass filter. The PI signal corresponding to the DNA content was measured with a 585-nm bandpass filter in conjunction with a 640-nm longpass filter. The doublet G2/M phase cells that resemble singlet G2/M phase cells were eliminated by setting a gate on the PI pulse intensity vs the PI pulse width blotting field.

Dot-blot analysis

Cell lysates in 0.01 M Tris-HCl (pH 7.2) and 0.15 M sodium chloride (Tris buffered saline: TBS) were prepared from logarithmic phase cells (3 days after subculture) using an ultra-sonicator. After centrifugation at 3000 g for 10 min. the supernatants were collected and the total protein concentration was determined using a Bio-Rad Protein Assay kit (Nippon Bio-Rad Laboratory, Tokyo, Japan) following the standard procedure. The protein concentration was adjusted to 100 µg ml-1, and serially diluted samples were fixed on a nitrocellulose membrane using a Bio-Dot Microfiltration Apparatus (Bio-Rad). The membrane was then air dried and soaked in the blocking solution (50 mM Tris, 2% BSA, 0.25% gelatin, 0.154 M sodium chloride) for 3 h. The HuMAb SK1 at 5 µg ml-1 in 0.5% BSA/TBS was then added and incubated at room temperature for 1 h. It was washed again and incubated with HRP-conjugated goat anti-human IgM (Jackson) for 30 min. After washing, the reactivity was visualized with a peroxidase substrate kit (Bio-Rad).

SDS-PAGE and Western blot

Cell lysates prepared with an ultra sonicator were mixed with a sample buffer (final concentration: 0.06 M Tris-HCl, 2% SDS, 10% glycerol) and heated at 95°C for 4 min. Acrylamide gel electrophoresis was performed with a Bio-Rad mini-Protein II slab gel apparatus using precast 4–15% linear gradient gels (Bio-Rad) under non-reducing conditions. Separated proteins were then transferred onto a nitrocellulose membrane using a Bio-Rad mini trans-blot apparatus. The blotted filters were stained with HuMAb SK1 as described above.

RESULTS

Colon carcinoma reactive antibody

A total of 1036 immunoglobulin-secreting hybridomas were generated from the 18 lymph nodes obtained from six Dukes' B colon carcinoma patients. The cell ELA and the immunocytotoxic chemical stainings revealed that 16 antibody-containing supernatants were reactive to colon carcinoma cells and such hybridomas were generated by each of the patients investigated. However, the immunocytotoxic chemical stainings of a panel of cultured cell lines with these antibodies showed that most of the selected HuMAbs were not specific to carcinoma but were multi-reactive, even to non-malignant cells (Koda et al. 1990). Among these HuMAbs, SK1 showed the most restricted specificity to carcinoma cell lines (Chang et al. 1993). This carcinoma-preferred reactivity was confirmed by immunohistochemical study in which SK1 reacted mainly with malignant tissues originating in the epithelium (Table 1; Figure 1). In many of the cases the observed immunohistochemical staining was heterogeneous, as clearly seen in Figure 1B and C.

Overall, 107 out of 109 colorectal cancer cases scored as Ag-SK1 positive, as were 10 out of 11 stomach cancers and four out of four pancreatic cancers (Table 1). Breast cancer scored positive in five out of eight cases, representing approximately 62%. Only one case out of 30 normal colorectal mucosa tissues tested scored as Ag-SK1 positive. Weak or trace stainings were observed with liver (two out of seven), pancreas (two out of four) and kidney (one out of four) tissues. All the rest were negative. The overall reactivity pattern was clearly adenocarcinoma-restricted.

Partial Ag-SK1 analysis

The HuMAb SK1 lost most reactivity to target cells treated with neuraminidase, trypsin or periodate (data not shown), suggesting that the antigen nature of Ag-SK1 was a glycoprotein. The approximate molecular weight of Ag-SK1 was a two-chain structure of 42–46 kDa, as seen by SDS-PAGE with non-reducing conditions, followed by Western blot (Figure 2). Normal colon mucosa of ulcerative colitis and fibroblast cells were negative for this two-chain structure whereas both primary colorectal carcinoma tissue and a colorectal cancer that metastasized to the liver were Ag-SK1 positive (Figure 2).

Ag-SK1 expression on cancer cells

Analysis of colonies of HT-29 cells, a carcinoma of the colon, indicated that Ag-SK1 expression was regulated by either cell

| Tissues       | Number of tissues |
|---------------|------------------|
| Colorectal cancer | 107              |
| Colorectal mucosa   | 1                |
| Stomach cancer     | 10               |
| Stomach mucosa     | 0                |
| Pancreatic cancer  | 4                |
| Pancreas           | 0                |
| Liver              | 0                |
| Lung cancer        | 2                |
| Lung               | 0                |
| Kidney             | 0                |
| Brain              | 0                |
| Melanoma           | 0                |
| Sarcoma            | 0                |
| Breast cancer      | 5                |
| Prostate           | 0                |
| Skin               | 0                |
| Oesophagus         | 0                |
| Bladder            | 0                |
| Testis             | 0                |
| Ovary              | 0                |
| Uterus             | 0                |
| Spleen             | 0                |
| Thyroid            | 0                |
| Heart              | 0                |
| Bone marrow        | 0                |
| Eye                | 0                |
| Nerve              | 0                |
| Skeletal muscle    | 0                |
| Salivary gland     | 0                |

Table 1 Immunohistochemical staining of human tissues with HuMAb SK1

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proliferation or cell density. This is shown in Figure 3. Figure 3A shows the degree on immunostaining by individual cells, and 3B shows a small colony. The outer peripheral region, typically a three- to five-cell-thick layer, of a larger colony of cells (Figure 3C and D) shows good immunostaining, whereas the cells in the centre of the colony are poorly, if at all, stained. The outer peripheral cells of the colony are actively growing, moving outward and therefore expressing Ag-SKI. On the other hand, the inferior cells of the colony are neither growing nor moving, and are Ag-SKI negative.

To further understand the proliferative nature of Ag-SKI, we analysed the expression of the antigen on Calu-1 cells, a carcinoma of the lung, each day as the culture expanded. These data are shown in Figure 4. Calu-1 cells were seeded at low density (Figure 4, 1D) and allowed to expand until the cultures were greater than 90% confluent, typically by days 6-7. As can be seen, the antigen is diffuse and spread throughout the cell in very low-density cultures. By day 3, the antigen has ‘rounded up’ and can be seen indenting the nucleus. High-density cultures were Ag-SKI negative by immunostaining. Similar findings were also observed in PANC-1, pancreatic cancer cells (data not shown). These data suggest that Ag-SKI is not always expressed on a given carcinoma cell but is metabolized in each cell depending on the circumstances. Two-colour FACS analysis using HuMAb SK1 and PI showed that antigen expression in each cell is independent of the cell cycle status (Figure 5).

**Dot-blot analysis**

The amount of Ag-SKI per extracted cell protein was assessed by dot-blot analysis (Figure 6, Table 2), where the ratio of Ag-SKI-positive cells was determined by FACS analysis (Table 2). We showed that the concentration of Ag-SKI per protein differs considerably among cell lines, in which SW620, WiDr and LoVo cell lines were considered to express a high amount of Ag-SKI, with the low expression group consisting of Caco-2 and NCI-H716 cell lines. The data shown in Figure 6 were obtained from cells harvested during mid-log growth (between 30% and 50% confluent, usually 3–4 days' culture) when antigen expression was highest.

In FACS analysis, the antigen-positive cells within each cell line were recorded as a discrete subset with more than ten times greater antigen expression in comparison with the negative cells (Figure 5B). The ratio of Ag-SKI-positive cells in each cell line varied in accordance with the harvested time of the cells. However, there was a tendency for cells with high Ag-SKI concentration in dot-blot analysis to have higher antigen-positive ratios in FACS analysis (Table 2).
Human natural antibody inhibiting cancer invasion

Figure 2  Western blot analysis of the antigen recognized by HuMAb SK1. Lanes 1–6 were stained with SK1, whereas lanes 8–11 were stained with irrelevant human IgM as a control. Lanes 1 and 8, human fibroblast cell line; lanes 2 and 9, normal colon mucosa of a patient with ulcerative colitis; lane 3, colon carcinoma cell line SW620; lane 4, colon carcinoma cell line LoVo; lanes 5 and 10, primary colon cancer tissue; lanes 6 and 11, liver metastases tissue of colon cancer; lane 7, molecular weight markers.

Figure 3  Immunocytochemical staining of colony-forming colon carcinoma cell line HT-29. All the cells in the small colony (A) were antigen-SK1 positive, whereas in intermediate or large colonies (B, C), only cells that form the peripheral region of each colony were stained with HuMAb SK1. On a magnified view of C (D), the intensity of staining differed considerably between inner and outer cells. Scale bar 50 μm in figures A, B and D; 500 μm in C.

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Successive changes in antigen expression found in Calu-1 lung carcinoma cell line. On 1–7 days (1D–7D) after the beginning of subculture, cells were stained with HuMAb SK1. A broadly expressed cytoplasmic antigen, which is seen on 1D and 2D, was condensed beside nuclei on 3D (indicated by arrows). The antigen was dispersed again on 4D, and was hardly seen on 7D when cells become more than 90% confluent. Scale bars = 50 μm

**Table 2** Cell invasion capacity and concentration of Ag-SKI

| Cell line   | Number of migrated cells (mean ± s.d.) | Concentration of Ag-SKI assessed by dot-blot analysis | Ag-SKI positive cells determined by FACS (%) |
|-------------|----------------------------------------|-------------------------------------------------------|---------------------------------------------|
| SW620       | 232 ± 52                               | 32x                                                   | >90                                         |
| WiDr        | 155 ± 37                               | 32x                                                   | 40–70                                       |
| LoVo        | 89 ± 23                                | 16x                                                   | 60–80                                       |
| LS174T      | 93 ± 21                                | 8x                                                    | 60–80                                       |
| COLO320DM   | 72 ± 15                                | 8x                                                    | 30–50                                       |
| HT-29       | 22 ± 13                                | 8x                                                    | 40–60                                       |
| NCI-H716    | <10                                    | 4x                                                    | 30–50                                       |
| Caco-2      | <10                                    | 2x                                                    | 20–40                                       |

*Mid-log phase cells (30–50% confluent) were harvested and prepared for analysis. Data indicate maximum dilution ratio of the extracted protein that can be detected by HuMAb SK1 on dot-blot analysis (see Materials and methods). The positive ratio was calculated with Kolgomolov–Smirov statistics.

**Invasion capacity of colon carcinoma cell lines**

The cell invasion capacity was determined with a microchemotaxis assay, and the data are summarized in Table 2. The number of cells that migrated to the lower chamber was high in SW620 and WiDr cell lines, whereas almost no invaded cells were observed in NCI-H716 and Caco-2 cells. When comparing the cell invasion capacity with the amount of Ag-SKI in each cell line, there was a tendency for the cell lines with high invasion capacity to have a higher amount of Ag-SKI (Table 2).

To confirm the hypothesis that Ag-SKI was an invasion- or migration-related molecule of colon carcinomas, we next investigated whether the invasion capacity of these cells was inhibited by the addition of HuMAb SK1 into the upper sides of the chemotaxis chambers in a quadruplicate assay. As shown in Figure 7, only a
limited number of cells migrated to the lower side of the membrane after 36 h of incubation with HuMab SK1. This inhibition was dose dependent and up to 87% of cell migration was blocked with 1 µg ml⁻¹ HuMab SK1 (Figure 8).

DISCUSSION

Data have been accumulating indicating that the overall immune system of cancer-bearing patients is able to recognize autologous tumour antigens specifically. In cellular immunity, the cytotoxic T lymphocyte that specifically targets autologous tumours has been induced in vitro (Barnd et al. 1989; Wright et al. 1989; Topalian et al. 1989; Peoples et al. 1995). In humoral immunity, human monoclonal antibodies against cancers have been primarily generated via hybridoma technology using B lymphocytes from cancer-bearing patients (Yamaguchi et al. 1990; Posner et al. 1991; Hoon et al. 1993; Oka et al. 1994). In spite of the existence of this specific cancer recognition by the patient’s immune system, tumours do grow in a physiological environment, implying that the humoral response was an insufficient contribution to the anti-tumour effect. The basic strategy of cancer biotherapy is to magnify or to modify the innate anti-tumour immune system using a variety of ‘bioweapons’ such as cytokines (Rosenberg et al. 1989; Weiner et al. 1991; Dillman et al. 1993; Ernstoff 1994), gene transfer (Porgador et al. 1993; Tohmatsu et al. 1993; Iwanuma et al. 1995), adoptive transfer of activated killer cells (Rosenberg et al. 1986; Andreesen et al. 1990; Keilholz et al. 1994) and cancer vaccinations (Elliott et al. 1993; Hoover et al. 1993; Houghton, 1995). So far, the majority of these approaches have involved cellular immunity, whereas human antibodies have been used in in vivo diagnosis of malignancies only (Ditziel et al. 1993; Chaudhuri et al. 1994; Ditzel et al. 1994).

The expected anti-tumour effects of human monoclonal antibodies could be listed as follows: (1) to bind the functional molecule in the metastasizing process of cancer cells, such as invasion, destroying basal membrane, or implantation in distant organs, thereby blocking the cancer metastases or expansions; (2) to induce antibody-dependent cell-mediated cytotoxicity (ADCC) or complement dependent cytotoxicity (CDC) against cancer cells; and (3) to induce anti-idiotypic antibody against the administered HuMabs. This second antibody may mimic the tumour-associated antigen(s) and offer an opportunity to induce a cell-mediated immune response specific to tumour cells (Ferrone 1993; Fargerberg et al. 1995).

The current data indicate that the antigen that is recognized by HuMab SK1 is a two-chain molecule with approximate molecular weight of 42–46 kDa, as it appears from the two most prominent bands (Figure 2). Tissue specificity analysis showed that HuMab SK1 preferentially stained adenocarcinoma tissues with minimal to negative staining among non-epithelial tumours and normal epithelium (Table 1). The antigen-positive cells within each colon cancer cell line seemed to represent a discrete subset with different levels of Ag expression, judging from FACS analysis (Figure 5) and from the immunocytochemical staining of HT-29 (Figure 3). There was a variety of Ag-positive ratios observed, depending upon the harvested time of the cells (FACS data from Table 2). Furthermore, there were variations in antigen location and inten-
Horizontally observed the capacity of the cell line to invade the membrane was completely inhibited by the addition of HuMAb SK1 in the upper chamber (right). Scale bars = 200 μm.

Figure 7 Microscopic features of upper and lower sides of the membrane in the chemotaxis chamber. LoVo colon cancer cells migrated to the lower side of the membrane after 36 h of incubation (left arrow). This cell migration was almost completely inhibited by the addition of HuMAb SK1 in the upper chamber (right).

Figure 8 Dose-dependent inhibition of cell migration with HuMAb SK1 added in the upper chamber. The vertical bar indicates a number of migrated cells to the lower side of the membrane per 10^6 cells. The horizontal line indicates concentration of antibodies. No inhibition was observed with irrelevant human IgM MAb. Cell line used was LoVo colon cancer cell line. ■ SK1; ● IgM.

sity in Calu-1 and PANC-1 cells (Figure 4). These data suggest that the antigen is not always expressed on a given cell but is possibly metabolized in each cell depending on the circumstances. A dot-blot analysis and FACS analysis showed that the invasion capacity of eight colon carcinoma cell lines correlates with the Ag-SK1 concentration of each cell line and with the ratio of Ag-SK1-positive cells (Figure 6, Table 2). In addition, cell migration was inhibited by the addition of HuMAb SK1 in a dose-dependent manner. Taken together, these data suggest that HuMAb SK1 recognizes an antigen related to colon carcinoma cell invasion that appears to be not only metabolized according to physiological circumstances but is also expressed on highly invasive cell populations.

Although SK1 did not cross-block with mouse monoclonal antibodies directed to the known invasion- or metastasis-related molecule such as E-cadherin, sialyl Tn, sialyl Le^a, sialyl Le^b, vitronectin receptor alpha-chain, integrin beta 1- or beta 3-chains (data not shown), the possibility that SK1 recognizes an epitope that contains some of these molecules still remains, as the mouse and human MAb might recognize different epitopes on the same molecule. This is an issue to be analysed further.

In the current study, we evaluated 1036 HuMAbs derived from 18 regional lymph nodes of six Dukes' B colon cancer patients. Among them, a high reaction specificity against cancer cells was seen in only one HuMAb, SK1. Although we used the lymphocytes derived from the nearest lymphoid system to the tumour, the appearance frequency of specific HuMAbs was unexpectedly small, suggesting that most of the patients may not raise a humoral immune response to autologous tumours, otherwise the B-cell population that is primed to secrete a specific antibody to tumours is small in number or is not fully activated to be fused (Koda et al., 1990).

The inhibition of in vitro colon cancer invasiveness by HuMAb SK1 required as much as 1 μg ml^-1 of antibody concentration (Figure 8), which is unlikely to be secreted by such small cell
populations of tumour-specific B lymphocytes in regional lymph nodes. This suggests that human natural antibodies to carcinoma, even when they exist, may not play an anti-cancer role effectively under physiological conditions.

A HuMab SK1 is considered to be an augmentation of the natural humoral immune response of cancer-bearing patients. It may encourage successful cancer biotherapy using such HuMabs that exert a direct action upon cancer invasion and metastases. Although the appearance of SK1 was low in frequency (1 out of 1036), its existence does suggest that there may be others also specific to cancer. The task then is to develop appropriate search protocols to prove effectively the entire human immune response repertoire and identify those sequences that identify cancer-specific antigens.

The availability of immunoreagents such as SK1 will not only provide new insights into the working of the anti-cancer human immune response but also give the oncologist clinically useful antibodies for better diagnosis and therapy.

**ABBREVIATIONS**

Ag. antigen: HuMab, human monoclonal antibody; kDa, kilodalton; HRP, horseradish peroxidase; ELA, enzyme immunoassay; PI, propidium iodide; SDS-PAGE, sodium dodecyl sulphate-polyacrylamide gel electrophoresis; FACS, fluorescence-activated cell sorter.

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