A TUMOUR-ASSOCIATED ANTIGEN FROM THE PLEURAL EFFUSION OF PATIENTS WITH SQUAMOUS-CELL CARCINOMA OF LUNG

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Summary.—A fraction showing tumour-associated antigenic properties has been isolated from pleural effusions of patients with squamous-cell carcinoma of the lung. Purification of the material was accomplished by ion-exchange and affinity chromatography, and by immunoabsorbents. The antigenic activity was monitored by its inhibitory capacity in a specific complement-dependent cytotoxic system. The final fraction has a mol. wt. of \( \sim 1.7 \times 10^5 \), as judged by gel filtration on Sephadex G200, and the main component appears to be a glycoprotein with N-acetyl-D-glucosamine groups. The most purified antigen preparation exhibited a highly selective capacity to inhibit in the cytotoxic assay and to bind, when labelled with \( ^{125}\text{I} \), to 2 specific antisera. The active fractions isolated from pleural effusions fully crossreacted with fractions prepared from squamous-cell carcinoma extracts. CEA and bacterial antigens were not detected in the material, and the presence of \( \alpha \)-fetoprotein, HLA and blood-group antigens may be ruled out on account of their respective molecular weights.

In recent years a number of papers have been published dealing with tumour-associated antigens (TAA)s isolated from human bronchogenic cancers (Holllinshead et al., 1974; Frost et al., 1975; Granlund and Ritts, 1976; McCoy et al., 1977; Kelly and Levy, 1977). The results showed a confusing variety in biochemical details (molecular-weight range, and electrophoretic mobility) as well as the TAA-activity (types and numbers of specificities, and crossreactions) which were probably due to the lack of standardized extraction and purification techniques, the frequent use of crude preparations, and the difference in activity-assessing tests. However, it generally emerged from the body of investigations that TAA(s) appear to occur in pulmonary cancers and might be obtained in a purified form by suitable fractionation methods.

Starting material for most of the present purification experiments was the pleural effusion from patients with squamous-cell carcinoma of the lung. Pleural effusions avoided damaging extraction procedures, and the volumes allowed work with the same material over a longer period of time with repetition and improvement of methods of preparation.

MATERIALS AND METHODS

Antisera.—Anti-lung-carcinoma sera were raised in rabbits by cell suspensions prepared from biopsy tissues of squamous-cell carcinoma patients. The first injection (0.5 ml of packed cells at a time) was given i.m. into the thigh followed by 3 i.v. injections at weekly intervals. No adjuvants were used. The rabbits were bled 10 days after the last injection by heart puncture under anaesthesia. This antiserum was code-named BL. An anti-normal-lung serum was produced in the same way. Other antisera were raised by fractions of pleural effusions. Four injections, containing 1–5 mg protein each, were given i.m. or i.d. respectively, with Freund's complete adjuvants, over a period of 4 weeks. The antiserum used in the present study was
prepared with a fraction absorbed by a Con A column (cf. Results Section). It was designated DU.

Antiseras to acute lymphocytic leukaemia (ALL), to chronic lymphocytic leukaemia (CLL) and to normal human lymphocytes, were produced in rabbits in a similar way to that described above, with cells which had been kept in liquid N₂.

A polyvalent antiserum to human total serum proteins raised in the pig, an antiserum to human foetal proteins raised in the rabbit and several precipitating antiseras to species immunoglobulins were obtained from Biogenzia Lemania SA, Switzerland.

Absorptions of antiseras and antigens.—Absorptions were carried out with cells or finely minced tissues, and/or with normal human sera (NHS) or antiseras crosslinked by glutaraldehyde. Absorptions by cells were performed v/v for 45 min at room temperature, and for 45 min at 4°C. Absorptions by crosslinked materials were done 3 times v/v for 1 h at 4°C.

Tissue-culture cells.—Cells were provided by our own tissue-culture department. E14 cells are a line derived from a squamous-cell carcinoma of the human lung. The cyto genetic properties of the established line have recently been reported (Fischer and Vetterlein, 1977). For control tests, batches were used of cell lines from an osteosarcoma, a spindle-cell sarcoma, a melanoma and foetal-lung fibroblasts, all of human origin.

Cells from patients.—Cell batches of acute lymphocytic leukaemia (ALL) of chronic lymphocytic leukaemia (CLL) and of human lymphocytes from normal individuals had all been stored in liquid N₂.

Pleural effusions.—Samples from patients with squamous-cell carcinoma of the lung were kindly supplied by Lainz Hospital, Vienna. They were obtained by sterile puncture followed by immediate centrifugation to remove all cellular and particulate material. The clear samples were stored at —20°C until use.

Tumour and E14 extracts.—Cell extracts prepared by phosphate-buffered saline (PBS) were sonicated and centrifuged at 48,000 rev/min for 1 h.

Monitoring of TAA activity.—The widely used test is based on the inhibition of a C³-dependent cytotoxic antiserum. For the present study it was applied as described previously (Wolf and Steele, 1975). Briefly, test fractions were incubated overnight at 4°C with a specific antiserum suitably absorbed and diluted. After that, ⁵¹Cr-labelled target cells followed by diluted guinea-pig complement were added and the reaction mixture was incubated at 37°C for 45 min. Ice-cold medium was then added, the samples were centrifuged to remove the cell debris, and an aliquot of the supernatant was counted in a Gamma Scintillation Spectrometer (Packard). The radioactivity released was expressed as a percentage of the radioactivity released by the antiserum only, and was used as parameter for the relative inhibitory capacity of the fraction.

Labelling of antigen with Naⁱ²⁵I.—A modification of the chloramine-T method of Hunter (1974) has been used (Wolf et al., 1976) without attempting to obtain a product of high specific radioactivity. To a 3 ml flask, containing 25 µl of antigen (10–20 µg protein)
were added 25 µl phosphate buffer (0.1 M, pH 7.4), 10 µl Na$^{125}$I (0.5 mCi), and, under constant stirring, 25 µl chloramine-T (4 mg/ml phosphate buffer). After 6 min, the reaction was stopped by 25 µl of sodium metabisulphite (12 mg/ml PBS) and 100 µl KI (100 mg/ml PBS), containing 1% bovine serum albumin, were then added. In order to separate the tagged protein (5–10 µCi/µg protein) from the free iodine the mixture was immediately passed through a column of Sephadex G 15 which had been equilibrated with PBS and coated with a few drops of 5% bovine serum albumin.

Saturation assay with $^{125}$I-antigen.—Purified TAA preparations, labelled with $^{125}$I-Iodine were tested for binding capacity to various specific and unspecific antisera. Ten or 20 µl of the labelled antigen and 50 µl of antiserum at various dilutions were mixed in small conical tubes (Sarstedt GmbH) and incubated at 4°C. Twenty-four hours later, 100 µl were added of a precipitating goat antiserum specific for the immunoglobulins of the first antiserum in order to separate bound from free antigen. After further 3 h at 4°C, the samples were centrifuged, the supernatants were completely removed and the precipitate in the test tube was counted in a Gamma Scintillation Spectrometer. Results were expressed as a percentage of the total counts added to each tube and the maximal uptake was used as parameter for comparisons.

RESULTS

The specificity of the TAA-monitoring system

The C'-dependent cytotoxic system consisted of the tissue-culture cells E14 (derived from a squamous-cell carcinoma of the human lung) as target cells, and the BL antiserum produced with cell suspensions prepared from a surgical specimen of squamous-cell carcinoma of the lung. A brief account of this system has recently been published (Wolf, 1977).

The BL antiserum was highly cytotoxic for the E14 cells, but, a more detailed characterization (Fig. 1) showed that antibodies to normal human serum proteins and to normal human lung tissue had to be removed by absorption in order to render the antiserum specific for the TAA of squamous-cell carcinoma. Table I demonstrates that, in the absorbed form, the BL antiserum in fact killed preferentially the specific E14 cells, showing little cytotoxicity to unspecific normal and tumour cells of human origin. It was therefore assumed that the absorbed BL antiserum would also be specifically neutralized (inhibited) by soluble TAA of squamous-cell carcinoma origin, and that, combined with the E14 cells, it would provide a specific system for the assessment of TAA-activity during fractionation procedures.

Inhibition experiments were performed with the absorbed BL antiserum at a final dilution of 1 : 20. For the inhibition tests, dilutions were prepared of all fractions according to protein content. Usually concentrations between 1 µg/ml and 100 µg/ml of protein were examined. The most purified antigen fractions showed a 100% inhibition at 100 µg protein/ml. The

TABLE I.—Tittrations of Specific BL Antiserum with Various $^{51}$Cr-labelled Cells

| BL antiserum | $^{51}$Cr cells | Maximal cytotoxicity (% of releasable label)* |
|--------------|-----------------|-------------------------------------------|
| Not absorbed | E14             | 70†                                       |
|              | Foetal lung fibroblasts (F 2000) | 35                                      |
| Absorbed on normal human serum and normal lung tissue | E14 | 30                                     |
|              | Foetal lung fibroblasts (F 2000) | 13                                      |
|              | Melanoma (MEL 364) | 8                                       |
|              | Osteosarcoma (2 T) | 2.5                                     |
|              | Spindle-cell sarcoma (FCHT) | 3                                       |
|              | Normal human lymphocytes | 0                                       |

* Releasable label is the radioactivity released by freezing and thawing.
† Mean values of 2 sets of duplicates. (A background of 10% is from all figures.)
Pleural effusion (Tumour extract) (50)
dialysed/Tris-HCl, 0·1m, pH 6·5
QAE-Sephadex A 50

\[
\begin{array}{ccccccc}
0·0m & 0·1m & 0·2m & 0·3m & 0·4m & 0·5m & (NaCl) \\
(20) & (0·5) & (0·5) & (5) & (10) & (1·0) & \\
(0%) & (0%) & (5%) & (30%) & (5%) & (0%) & \\
+ & + & + & + & + & + & \\
\end{array}
\]

Con A column, PBS, pH 7·2

not absorbed, absorbed, eluted by 50 mm glucoside
(2·3) (1·0)
(5%) (25%)
++ +

[A] 125I-labelled

Wheat-germ-lectin column
PBS, pH 7·2 (1·0)

not absorbed, discarded, eluted by 100 mm glucosamine
(0·35) (0·05)
(10%) (20%)
++ +

Sephadex G 200
PBS, pH 7·2

\[
\begin{array}{ll}
1·5-1·8 \times 10^5 & <6·5 \times 10^4 \\
\text{mol. wt.} & \text{mol. wt.} \\
\end{array}
\]

absorbed on anti-human serum proteins and anti-lung tissue antisera (protein not measurable)
(15%)
++ +

125I-labelled uptake by antiserum

Fig. 2.—Flow diagram of the purification procedure. Average protein yields (mg) estimated by Folin-Ciocalteus phenol reagent, and % inhibition in the 51Cr assay are given in brackets. + inhibition at 100 µg/ml; ++ inhibition at 10 µg/ml; +++ inhibition at 1 µg/ml.

The preparation of the antigen

Antigenic fractions were chiefly prepared from pleural effusions, although control extracts of squamous-cell carcinoma tissue and of E14 cells were fractionated in the same way. The flow chart of the fractionation procedure is shown in Fig. 2. The starting material was dialysed against Tris/HCl buffer, 0·1m, pH 6·5, and the sample (3–5 ml containing up to 150 mg protein) was applied to a QAE-Sephadex A 50 column (1·9 x 12 cm) equilibrated with the same Tris/HCl buffer. Stepwise elution was performed with increasing NaCl starting from zero concentration up to 0·5m in Tris buffer. Activity was found in fractions eluted by 0·2–0·4m NaCl, but most of it was recovered in fractions eluted by 0·2m or 0·3m NaCl. Active fractions were concentrated by polyethylene glycol, dialysed against phosphate-buffered saline.
(PBS) pH 7.2, and layered on to a Con A-Sepharose (Pharmacia) column (1.0 × 8.5 cm) equilibrated with PBS, pH 7.2. The column was washed with PBS and the absorbed material eluted with 0.05M α-methyl-D-glucoside in PBS. Most of the activity was recovered in the bound material, which again was dialysed and concentrated.

For further purification 2 different methods were used.

(A) In order to obtain peak distinction in gel filtrations (in spite of low protein content of the fractions at that stage), the antigen was labelled with $^{125}$Iodine and the protein fraction, after passing a Sephadex G 15 gel (cf. Labelling) was layered on to a Sephadex G 200 column (0.9 × 30 cm) equilibrated and eluted with PBS, pH 7.2. The elution profile showed 2 $^{125}$I-peaks, the first appearing shortly in front of human IgG and the second shortly after human albumin.

(B) By the second method, the active fraction of the Con A column was applied to a column of wheat-germ lectin (WGL) coupled to Sepharose 6MB (Pharmacia). This column (2 ml volume) was equilibrated and washed with PBS, and the absorbed material, eluted by N-acetyl-D-glucosamine (100 mM in PBS) was dialysed very carefully (in order to remove all glucosamine) and then concentrated. The glycoproteins absorbed by the wheat germ lectin appeared to possess high specific inhibitory activity in the cytotoxic assay. The preparation was iodinated and examined for uptake by antisera in the saturation assay (Table II), or it was submitted to successive absorption on crosslinked antisera to normal human proteins and to normal human lung tissues.

**Table II.—Saturation Analysis Carried out with Various Antisera and $^{125}$I-labelled Antigens**

| $^{125}$I Antigen (origin and fraction type) | Antiserum* | Maximal uptake of $^{125}$I antigen (%)§ (Bound/Total)†‡ |
| --- | --- | --- |
| **Pleural Effusion** | | |
| Con A-bound material | BL, not absorbed | 35 (34;31;36;38;35;35) |
| | absorbed | 20 (18;19;21;22) |
| | DU, absorbed | 30 (25;29;30;35) |
| | anti-normal lung tissue, not absorbed | 8 (7;8) |
| | anti-human foetal protein, not absorbed | 10 (9;11) |
| | anti-CLL-cells, not absorbed | 13 (10;17;19;16) |
| | anti-ALL-cells, not absorbed | 3 (3;3) |
| | anti-normal human lymphocytes, not absorbed | 3 (3;3) |
| **Sephadex G 200** | | |
| Fraction: 1.5–1.7 × 10$^4$d | BL, absorbed | 25 (21;24;25;29) |
| < 6.5 × 10$^4$d | DU, absorbed | 70 (69;70;75;68;69;76) |
| | BL, absorbed | 2 (2;2) |
| | DU, absorbed | 8 (8;8) |
| **Wheat germ lectin-bound material** | BL, absorbed | 15 (14;13;18;19) |
| | DU, absorbed | 55 (55;54;52;53) |
| **Squamous-cell carcinoma (cells)** | | |
| Con A-bound material | BL, absorbed | 15 (13;15;19;14) |
| | DU, absorbed | 25 (26;28;24;23) |
| **Material not bound** | BL, absorbed | 0 (0;0) |
| | DU, absorbed | 0 (0;0) |
| **E14 (cultured cells)** | | |
| Con A-bound material | BL, absorbed | 12 (12;10;14;13) |
| | DU, absorbed | 20 (19;20;23;21) |

* Antisera were titrated in 5-fold dilutions from 1:20 to 1:62,500. Maximal uptake occurred at dilutions 1:20–1:100.
† In most cases the total was 20,000 ct/min. In case of the G 200 fractions it was 6000 ct/min.
‡ Background of 6–10% subtracted before calculation.
§ Mean values of duplicates of 1–3 tests (first column), and values of single tubes (in brackets).
isolated tissue. Culture cells and both selves recovered from tumour cells of normal human serum and lung.

**Saturation Analysis**

Fractions which proved to be good inhibitors in the cytotoxic system, along with non-inhibiting control fractions, were labelled with $^{125}$I and tested in a saturation-type radioimmunoassay to provide additional evidence for their specificity by being selectively taken up by the specific antiserum BL. It was, furthermore, investigated whether the BL antiserum (raised by tumour cells) and the DU antiserum (raised by pleural-effusion fractions) would exhibit similar specific properties in this test, and whether crossreactions would occur between both antisera and fractions from pleural effusions, tumour-cell extracts and tissue-culture cell extracts.

It may be gathered from Table II that the labelled antigens were taken up by both antisera, BL and DU, the antiserum DU being, generally, more avid than antiserum BL. It will also be noted that the antigen uptake was selective. Unspecific antisera reacted much less with the antigen preparations, in particular the antiserum specific for normal human lymphocytes did not bind antigen at all, although it exhibited a good cytotoxic titre with normal lymphocytes (unpublished data). Besides, control preparations, e.g. the low-molecular-weight component obtained from the Sephadex G 200 column, and the material which was recovered in the void volume of the Con A-columns, both did not attach themselves to either antiserum. There was also a wide crossreaction between the antigens isolated from squamous-cell carcinoma tissue, pleural effusions, and the E14 culture cells and both antisera. No evidence was found for a stronger uptake of antigens from tumour cells by the BL antiserum as compared with the DU antiserum. These results thus appear to prove that the pleural effusions contain the same antigens as occur on, and can be extracted from, squamous-cell carcinoma cells. They also confirm the specificity of the $^{51}$Cr test by showing that the uptake of labelled antigens corresponds principally to their inhibitory capacity.

**DISCUSSION**

The BL antiserum (produced with squamous-cell carcinoma cells of a patient) after exhaustive absorption on normal human serum proteins and normal lung tissue, still exhibited a cytolytic effect on the E14 target cells, which could be selectively inhibited by certain isolated and purified materials. These inhibiting substances may, therefore, contain specific antigens, associated with squamous-cell carcinoma cells, which do not occur in sera and lung tissues of normal individuals (or only on very low levels).

Biochemically, the antigen would appear to be a glycoprotein with N-acetyl-glucosamine groups, which are the sugar groups specifically bound by wheat-germ lectin. To judge by elution profiles from gel-filtration columns, the molecular weight of the most purified fractions was slightly higher than that of IgG (i.e. $\sim 1.7 \times 10^5$ dalton). After passing the WGL-column the specific inhibitory activity of the antigen preparation had increased considerably, an observation perhaps due to an effect similar to that described by Clemetson et al. (1976) who were able to separate normal transplantation antigens from tumour-specific antigens in extracts from a murine mastocytoma on columns of wheat-germ lectin.

During the course of the separation procedure, it was observed that, apart from fractions containing most of the inhibitory activity, some others of different constitution exhibited small inhibitory effects. This may be due to imperfect
separation, or to different molecules carrying the same active groups. It may also indicate genuinely dissimilar, weakly crossreacting substances, and confirm the existence of various distinct TAA-activities which were recently reported by Veltri et al. (1977).

The molecular weight found for the described isolated antigen is close to that of carcinoembryonic antigen (CEA) (Tillack et al., 1974). But, by radioimmunoassay, kindly performed by Mr B. Chaput (Hoffman La Roche, Vienna) several crude and purified fractions of the present series were all negative for CEA. The presence of α-fetoprotein and of HLA antigens can also be ruled out, because of their respective molecular weights, and so may the involvement of blood-group antigens (mol. wt. 2–3 × 10^8). Finally, contamination by bacterial antigens from pleural infections could not be detected in the antigen fractions by an antiserum raised to a wide bacterial spectrum. Some caution, concerning possible cross-reactions of antigens from certain other tumours, appears to be, nevertheless, appropriate. Like the anti-CLL serum that shows a relatively high antigen uptake in the saturation assay, the BL antiserum produced a relatively high cytotoxic titre with Ewing sarcoma cells. Studies of crossreactions of the antisera with such “unrelated” antigens, and with “related” antigens such as TAAs from lung cancers of different histopathological origin, as well as the suitability of the isolated antigen as a diagnostic marker for lung cancer in clinical trials are now in progress.

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