ABILITY OF CEA BLOOD LEVELS TO REFLECT TUMOUR BURDEN: A STUDY IN A HUMAN XENOGRAFT MODEL

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Summary.— The relationship of serum carcinoembryonic antigen (CEA) levels to tumour size and antigen content was studied in artificially immune-deprived mice bearing human colonic, breast and lung tumour xenografts. Size was measured as in vivo volume and tumour weight at post-mortem. A multiple implant technique combined with early harvest was used to minimize centrilobular tumour necrosis. CEA was extracted from resected tumours with perchloric acid. A radioimmunoassay using chemical precipitation was used to estimate CEA in blood samples. A correlation was found between CEA blood levels and tumour size in half the tumour lines, in contrast to a recent report (Lewis & Keep, 1981). The CEA content was found to be constant for one tumour line but not another. The possibility that central necrosis in xenograft tumours may account for the discrepancies is discussed. There may be serious limitations for the use of xenograft tumour models for studying the biology of CEA.

Carcinoembryonic antigen (CEA) (Gold & Freedman, 1965) remains one of the most useful tumour-associated substances to monitor malignant disease. Unfortunately, its assay in the blood provides only a crude guide to tumour behaviour; there are two main reasons for this. First, CEA is a heterogeneous large glycoprotein which is difficult to define in precise chemical and immunological terms with many of the available reagents. Second, regulation of CEA levels in cancer patients is subject to numerous poorly understood biological variables with respect to its production and release by tumours, and the mode by which the host metabolizes and clears it from the circulation.

Of fundamental importance to the clinical application of any marker is its ability to provide a reliable index of tumour mass. Unfortunately, CEA may have certain failings in this respect. It is true that there is usually an increase in CEA in individual patients as disease advances (Steward et al., 1974; Di Saia et al., 1975; Khoo & Mackay, 1976; Cove et al., 1979; Dent & McCullough, 1980 and that many patients with advanced malignancies are likely to have high levels of CEA (Chu & Nemoto, 1973; Steward et al., 1974; Barrelet & Mach, 1975; Vincent et al., 1975; Khoo & Mackay, 1976; Di Saia et al., 1977; Borthwick et al., 1977; Martin et al., 1977; Waalkes et al., 1980; Wanebo 1980) which may correlate with the stage of the disease (Martin et al., 1977; Cove et al., 1979; Khoo et al., 1979; Khoo & Mackay, 1976; Gropp et al., 1980; Joyce et al., 1980) but there are wide discrepancies in all series, which are difficult to explain. Thus, there appears to be a need for more precise information on the subject. Unfortunately the inherent variability of clinical studies militates against uniformity, but an

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experimental study under laboratory conditions could have distinct advantages. An investigation is now described, using a model tumour system consisting of immune-deprived mice bearing human CEA-producing tumours.

**MATERIALS AND METHODS**

**Human tumours.**—Human tumours already established in transplant passage to immune-deprived mice were screened for their CEA content and ability to produce high titres of CEA in the blood of host mice. Four colorectal (HK1, 6, 7, 9), one breast (S32) and one lung (p246) were selected for use. The number of previous passages ranged from 7 to 16. Plasma CEA levels ranged from 40 to 306 ng/ml. The histological characteristics of the primary tumours were retained in the xenografts, except that stroma was much reduced and predominantly of mouse origin, as is well described by Warenius (1979). The karyotypes were in all cases human; only occasional murine chromosomes being encountered. There was a considerable range of modal number, from 40 to 80 chromosomes.

**Immune-deprived mice.**—CBA/LAC mice were used throughout. Preparation involved thymectomy at 4–6 weeks of age, and 3 weeks later, whole-body irradiation (9 Gy at 60 cGy/min; 200 kV X-ray machine) followed by reconstitution within 2 h by an i.v. injection of $5 \times 10^6$ syngeneic marrow cells (Miller et al., 1963). The mice were suitable for xenografting from 14 days after irradiation and reconstitution.

**Grafting techniques.**—Under ether general anaesthesia, 2mm$^3$ fragments of tumour were placed into subcutaneous sites in each flank through a single dorsal incision. When it was necessary to vary the bulk of tumour carried by individual mice, multiple implant sites were used, 4 dorsal and 2 ventral, through separate incisions.

**Tumour measurements.**—Tumour bulk was assessed as volume in most studies. When multiple implants were made, tumours often coalesced, making volume measurement impossible. These tumours were excised at post-mortem and weighed. Volumes were measured as $\pi D^2/6$, $D$ being the mean of two diameters at right angles (Nowak et al., 1978). This assumes that the tumour is approximately spherical. Tumours which failed to retain a spherical or slightly ellipsoidal shape were therefore excluded. Measurement of depth diameter was omitted because it is subject to greater experimental error (Dethlefsen et al., 1968). The accuracy of this formula was checked from a calibration curve of volume measurements against the weight of tumour obtained after excision; this indicated a slight tendency to over-estimate actual tumour volume, in part due to the thickness of the skin. No attempt was made to correct for this because the relative values were unlikely to be significantly in error.

**Collection of blood samples.**—Individual blood samples were usually collected by venesection via the infra-orbital sinus. When mice were to be killed cardiac puncture was used. Fine glass pipettes coated with 360 µg tripotassium EDTA were used and ~200 µl withdrawn each time. Plasma was separated and stored at −70°C until assayed.

**CEA assay.**—For a number of reasons it was necessary to devise a radioimmunoassay especially for this study. Only small volumes of blood could be removed from mice without risk of serious hypovolaemia. The automated assay in clinical use required large plasma samples. Since with the available equipment, samples of 50 µl were found to be the minimum before unacceptable errors from the manual pipetting manoeuvres occurred, most of the mouse sample would be required for a single assay. It was thus necessary to devise a more sensitive assay.

Chemical precipitation for separating the antisera-bound from free CEA was chosen, because it is simpler and cheaper than a double-antibody method, and because mouse anti-CEA/CEA complexes are unlikely to reduce the amount of antigen available for binding in this type of assay (Stevens et al., 1978).

The assay was set up in the conventional way by Dr M. G. Ormerod and Miss N. Neylon. The CEA used was isolated from hepatic metastases of a human colonic carcinoma and purified to satisfy criteria described by Westwood & Thomas (1975) and Westwood et al. (1978). Anti-CEA sera were raised in a rabbit by the method described by Ormerod (1978). CEA was labelled with $^{125}$I using chloramine T by Mr M. Capp. All dilutions were in phosphate-buffered saline (0-05M phosphate, 0-15M NaCl, pH 7-15).
containing 1 g/l bovine serum albumin (PEB). Additional precipitants were 11-5% polyethylene glycol (PEG) and pooled human albumin.

A dilution curve of rabbit CEA antiserum against 125I-CEA diluted to give about 2 x 10⁴ cts/min/ng CEA was obtained. Doubling dilutions of the antiserum in PEB were made, beginning at 1:400. Duplicate 50 µl volumes of diluted antiserum, labelled CEA and pooled normal mouse plasma were mixed and incubated at 37°C overnight. Fifty µl stored human albumin and 1 ml 11-5% PEG solution were added and stored at 4°C for 30 min. The tubes were then centrifuged and the radioactivity of the pellet estimated. A

titration curve (Fig. 1) indicated that ~90% of the labelled 125I-CEA reacted with the antiserum. A dilution of 1:5000 is shown to react with 50% of the maximum amount of 125I-CEA. This dilution was used throughout.

For each assay a standard curve (Fig. 2) was carried out using 50 µl solutions with known amounts of unlabelled CEA, 1-0 ng 125I-labelled CEA and 1:5000 dilution of rabbit antiserum.

Specificity.—Ormerod (1978) has previously reported that the rabbit antisera used in this study do not react with these antigenic determinants of CEA which are shared with the non-specific cross-reacting antigen (CEX). A dilution curve of a highly concentrated
CEA-containing pooled mouse plasma sample conformed to the standard curve, indicating the purity of CEA produced in the mouse.

**Precision.**—The mean calibration with its 95% tolerance limits is shown in Fig. 2. The precision throughout the dose range is determined from these 95% tolerance limits by taking half this difference divided by the nominal dose to give the relative dose error at this point. These dose errors, plotted against dose to determine the region of minimum error in the assay calibration curve revealed a minimum dose error of 12.5% at 60 ng/ml. At the extreme ranges of the assay this was 68 and 78%.

**Reproducibility.**—Because batches of completed experiments were assayed together on the same assay day, and because the relative rather than absolute values of the CEA estimations were important, the inter-assay variation was irrelevant. Samples of known value were included in each assay and intra-assay variability was ~10%.

**Tissue CEA.**—CEA was extracted from freshly resected specimens using perchloric acid by the method of Khoo et al. (1973). CEA levels were measured by the double-antibody radioimmunoassay method of Laurence et al. (1972) which is used for measuring plasma levels in hospital patients at the Ludwig Institute. Although there were obvious variations in precision at individual CEA titres between this assay and the mouse plasma assay described above, there were no discernible qualitative differences.

**Statistics.**—The significance of the analysis of data was calculated using Spearman's test of rank correlation coefficient.

**RESULTS**

**CEA blood levels related to tumour mass**

CEA rarely became detectable in the blood of mice bearing the human tumours until the total tumour volume reached ~0.6 ml. As individual tumours reached

![Graph](image)

**Fig. 3.**—Relation of plasma CEA to tumour volume in colon tumours. Rank correlation HK1; 0.92, P < 0.01. HK9; 0.32, P > 0.05.

| Tumour | No. of mice | Tumour volume range (ml) | Tumour mass range (g) | CEA blood levels range (ng/ml) | Correlation: CEA levels vs CEA levels | Correlation: Tumour volume (or mass) vs CEA levels | P |
|--------|-------------|--------------------------|-----------------------|-----------------------------|---------------------------------------|----------------------------------|----|
| Colon  |             |                          |                       |                             |                                       |                                  |     |
| HK1    | 9           | 0.7–2.2                  |                       | 20–270                      | 0.92                                  | < 0.01                           |    |
| HK6    | 20          | 1.1–3                    |                       | 20–250                      | 0.26                                  | NS                               |    |
| HK7    | 7           | 1.1–2.3                  |                       | 20–110                      | 0.82                                  | < 0.05                           |    |
| HK9    | 25, 11      | 0.8–3, 0.2–8.1           |                       | 0–220, 20–640               | 0.32                                  | NS                               |    |
| Breast |             |                          |                       |                             |                                       |                                  |     |
| S32    | 35          | 0.5–5                    |                       | 20–620                      | 0.44                                  | < 0.05                           |    |
| Lung   |             |                          |                       | 0–850                       | 0.38                                  | NS                               |    |
| p246   | 18          | 0.4–2.4                  |                       |                             |                                       |                                  |     |

Table.—Correlation of tumour size with blood CEA level (Spearman’s rank correlation)
1·5 ml central necrosis set in. Therefore readings were usually taken 4–6 weeks after transplantation, when most of the tumours had grown to 0·5–1·0 ml. Mice bearing tumours of volume < 0·5 ml, or tumours which were frankly necrotic, ulcerated or infected, were discarded.

The results of the bilateral subcutaneous flank volume study are summarized in the Table, and illustrated in Fig. 3. The wide scatter of CEA values within each tumour group was of particular interest, and in only half was there a correlation with tumour volume HK1 (Fig. 3) \( P < 0·01 \), HK7 \( P < 0·05 \) and S32 \( P < 0·05 \).

The greatest discrepancies were found in HK6, where the largest tumours were associated with the lowest plasma CEA levels, whereas in p246 and HK9, mice with relatively small tumour loads occasionally had high levels of circulating CEA. Among HK6, HK9 and p246, it appeared that the highest CEA levels occurred in mice bearing tumour burdens in the mid range. It was considered that some discrepancies might have occurred because of centrlobular necrosis in the larger tumours. However, although post-
mortem examination confirmed this in certain cases, low CEA levels were found in association with some large tumours in which centrilobular necrosis was not excessive.

The multiple implant study was designed to minimize the effect of centrilobular necrosis. A much wider range (0.5–8 g) could thus also be attained. The results again revealed a wide variation, but an overall linear correlation \((P < 0.05)\) (Fig. 4). One notable discrepancy was in a mouse carrying one of the largest tumour loads, which had a particularly low plasma level of CEA.

The relationship of tumour CEA content, tumour mass and CEA blood levels

HK9 and HK6 tumours were used for this investigation. The HK9 study was an extension of the multiple-implant investigation described above, whereas the HK6 multiple implants were set up specially. (The plasma CEA estimations for this particular group were rendered invalid through technical fault.)

Whereas HK6 demonstrated a clear linear correlation \((P < 0.05)\) of tumour CEA content with tumour mass, considerable disparity existed in HK9 (Fig. 5) indicating that CEA concentration in xenograft tumours does not remain constant. The relationship of tumour CEA content to plasma levels in HK9 was even more disparate. The possibility had to be considered that where there was wide discrepancy between tumour mass and CEA blood levels, there may have been a corresponding discrepancy in tumour CEA content, but this was clearly not the case.

There is no indication, therefore, that tumours with low or high concentrations were especially associated with corresponding plasma CEA levels.

DISCUSSION

Previous xenograft investigators have pointed out that tumours must reach a certain size before CEA becomes detectable in the blood (Primus et al., 1973; Mach et al., 1974; Sordat et al., 1974; Miwa et al., 1976; Lewis & Keep, 1981). Indeed, CEA was as rare in mice bearing small tumours in the present study that it became policy not to measure plasma levels until the total tumour bulk was \(~1.5~\text{ml}\). Such a size represents \(~5\%\) of the total weight of the tumour-bearing host, which in comparison to malignancies in patients is enormous. In contrast, relatively small tumours in patients are capable of producing very high levels of circulating CEA. This curious observation has also been made for other markers, such as human chorionic gonadotrophin (hCG) (Kameya et al., 1975) and \(\alpha\)-foeto-protein (AFP) (Raghavan et al., 1980) produced by xenograft tumours. The reason cannot be wholly explained by assay insensitivity, since small concentrations should be detectable in spite of measurement being inaccurate. It would seem logical that CEA would only become detectable when the amount produced and released by the tumour exceeds that eventually metabolized. However, although it has been shown that CEA is rapidly cleared by the liver in mice (Thomas et al., 1976; Thomas & Hems, 1975) the possibility of a threshold level for CEA clearance has not yet been specifically investigated.

The possibility that there may be a change in CEA synthesis during growth should seriously be considered. Certainly, as tumours grow centrilobular necrosis sets in, and the CEA concentration is naturally smaller when this becomes extensive (Mach et al., 1974). It is not surprising that Lewis & Keep (1981) should find in their single tumour line that CEA concentrations vary widely, because their series produced tumours of vastly differing size. The linear correlation which was found in the HK6 tumours in the present investigation may be because the tumours were harvested early, before serious centrilobular necrosis could have occurred. The possibility that necrosis may be an important factor affecting CEA blood levels has been investigated by the
author in a separate study (Quayle, 1982) in which it was found that when necrosis occurred rapidly and extensively, considerable increases in CEA titres were frequent. Nevertheless, the fact that the tumour mass vs tumour CEA correlation did not exist in HK9, in spite of similar conditions, indicates that factors other than necrosis are responsible.

A further anomaly identified in this study is the finding that tumour size does not always correlate with plasma CEA. Again this cannot wholly be explained on the basis of tumour necrosis because, if it were, the correlation in the multiple-implant study, which was harvested early, should have been closer than in volume studies.

Until recently, Miwa et al. (1976) were the only workers to investigate the relationship between blood CEA and tumour xenograft size. Their claim for a direct correlation does not, however, stand up to close scrutiny, mainly because the number of samples is small (5) and the spread of levels and tumour size uneven. Stragand et al. (1980) and Lewis & Keep (1981) failed to demonstrate any correlation in their more recent detailed studies on a single tumour line, but the validity of their observation should perhaps be viewed with some circumstances since they were using tumours which, in some cases, were as large as 8-8 and 7-14 g respectively, which would be expected to contain a considerable degree of central necrosis. Indeed Lewis & Keep (1981) specifically stated that "central necrosis was a consistent feature" in their tumour line. Since the present study has indicated behavioural differences between tumour lines, it is conceivable that the two tumour lines used by these authors may not be representative. A further factor, again acknowledged by Lewis & Keep (1981), was that their assay for CEA by a double antibody technique may have been incapable of determining the presence of CEA molecules masked by murine immunoglobulins.

The variable ability of CEA to reflect tumour burden is not wholly unexpected, since a review of the extensive literature indicates that levels of CEA in the blood are affected by numerous other biological variables. Even so, because the validity of xenograft tumour models for such investigations remains in doubt, this failing of CEA may have been exaggerated.

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