Effect of tumour associated tissue eosinophilia on survival of women with stage IB carcinoma of the uterine cervix

P B Bethwaite, L J Holloway, M-L Yeong, A Thornton

Abstract

Aims—To examine the survival of a group of women with stage IB invasive carcinoma of the uterine cervix, divided according to the expression of tumour associated tissue eosinophilia (TATE).

Methods—Histological material from 81 women with stage IB squamous and adenosquamous cervical carcinomas before radiotherapy was assessed for the extent of tissue stromal eosinophilia, quantified using antibodies to human major basic protein.

Results—Twenty eight (38%) of the cases demonstrated TATE of over 30 eosinophils/mm², with 12 (16%) having greater than 100 eosinophils/mm². Eleven women in the series developed distant spread or recurrent pelvic disease, this group having a stromal eosinophil density significantly less (13.8/mm²) than the remainder (69.9/mm²) (p = 0.03). The actuarial five year survival rate for women with a tumour eosinophil density over 30/mm² was 92% compared with 70% with a density under 30 mm², with a significant difference in the survival curves for these two groups (p = 0.03).

Conclusions—As a univariate parameter, a tumour associated tissue eosinophilia of at least modest proportions is associated with statistically improved survival in women with stage IB cervical carcinomas.

(J Clin Pathol 1993;46:1016–1020)

Solid tumours are associated with a variable pattern of stromal inflammatory cell infiltration, usually characterised by a lymphoplasmacytic response. A heavy infiltrate is associated with improved prognosis in a number of tumour sites.1 Eosinophilic leucocytes are sometimes observed as a component of the stromal infiltrate, and a tumour associated tissue eosinophilia (TATE) is well described in a range of tumour types and sites, including carcinoma of the cervix,2–8 lung,5 gastrointestinal tract,10 11 and in transitional cell carcinoma of the bladder.12 Previous work has suggested that TATE may also be associated with a better outlook.8

In the uterine cervix around 3% of cases show an intense TATE. These tumours, with a stromal eosinophil density of over 100/mm², are easily recognised in routine histological sections.6 Some workers have reported an improved survival with “intense TATE”4 6 11 while others find no apparent association.2 7 14

More common, yet less well characterised, are cervical carcinomas with a mixed inflammatory infiltrate, which includes a lesser proportion of eosinophilic leucocytes, usually admixed with chronic inflammatory cells. This “moderate TATE” group with a tissue eosinophil density between 30 and 100/mm² has not been included before when examining the effect of stromal eosinophilia on survival.

Methods

Women with FIGO (International Federation of Gynaecology and Obstetrics) stage IB invasive carcinomas of the uterine cervix were abstracted from a larger series of women with invasive cervical cancers. This series was previously reviewed to examine the effect of tumour mucin production on prognosis, and the details are published elsewhere.15 In summary, the series was drawn from a clinicopathological register of 186 cases of invasive cervical cancer, stage IB and above, presenting to a regionally based gynaecology oncology service between 1980 and 1987.

The series included 90 stage IB tumours (62 squamous cell carcinomas, 19 adenosquamous carcinomas, eight adenocarcinomas and one mesonephroid tumour). The 81 squamous and adenosquamous tumours were selected for the study. As radiotherapy is known to induce TATE, eight cases were excluded for study as only histological material after radiotherapy was available. The histological material before radiotherapy was reviewed from the remaining 73 cases and two representative blocks were selected from each case and the haematoxylin and eosin stained sections studied. The degree of the lymphocytic or plasmacytic stromal response was graded dichotomously as either minimal to scant, or moderate to heavy.

To assist with accurate quantification of eosinophil tissue density, sections were cut from the selected blocks and the avidin-biotin-peroxidase complex (ABC) method was used to demonstrate eosinophil major basic protein (MBP) immunohistochemically.16 Endogenous peroxidase was blocked with hydrogen peroxide, periodic acid, and potassium borohydride. Digestion in 0.1% trypsin was followed by overnight incubation at 4°C with the primary polyclonal antibody, anti-human MBP, donated by Dr Gleich, Mayo Clinic, at a dilution of 1 in 4000.
Visualisation was achieved by 3,3'-diaminobenzidine tetrahydrochloride (DAB, Sigma No D-5637), with haematoxylin counterstaining. The number of positively staining cells in the densest region, usually at the advancing tumour edge, in 40 microscopic fields at 400 times magnification were counted. The field areas were determined using a calibrated grid and the eosinophil density expressed as number per square millimetre of tissue.

Lowe has defined intense TATE as an eosinophil density over 100/mm², roughly corresponding to those cases reported by various workers as cervical carcinoma with pronounced stromal eosinophilia. Decreasing the threshold to 30 eosinophils/mm², however, includes tumors with a moderate infiltrate, which are often not initially recognised in routine sections. These cutoff points were chosen a priori for the current study.

The tissue eosinophil density was compared among the major clinical and pathological subgroups using the Mann-Whitney U test. Detailed survival information was available from the register; survival was calculated from the date of diagnosis to the date of last follow up, intercurrent death, or cervical cancer related death. Survival was analysed using the product limit method of Kaplan and Meier, with univariate parameters assessed using the log-rank test.

Results

Among the 73 cases studied, 67 were staged FIGO stage IB, the remaining six as IB occult. The age range of the patients was 25 to 76 years, with a mean of 43.7 years. Fifty-six cases (77%) were non-mucin producing squamous cell carcinomas, while the remaining 23% were adenosquamous tumours, as already defined by us to include "covert" mucin producing squamous cell tumours.15

The women were treated as follows: routine hysterectomy alone n = 2; Wertheim's hysterectomy alone n = 16; preoperative caesium/Wertheim's hysterectomy n = 43; hysterectomy/postoperative radiation n = 9; radiation only n = 3. Details of the protocols are given elsewhere. Pelvic lymph node sampling was undertaken in 64 women, with 17 (27%) having histologically confirmed nodal disease. Six of 73 women had evidence of subsequent distant metastatic disease (lung n = 3, brain n = 1, bone marrow n = 1, cervical lymph nodes n = 1) or recurrent pelvic disease n = 5.

Quantification of eosinophil infiltration for all cases showed a range from 0 to 542 eosinophils/mm² (mean (SD) 61.5 (115.5). Twenty eight (38%) cases had an eosinophil density over 30/mm², with 12 (16%) having greater than 100/mm².

Table 1 shows that there was no significant difference in stromal eosinophil density among different tumour subgroups defined on the basis of histological type, tumour size, depth of invasion or presence of vascular space invasion. Younger women seemed to have a higher stromal eosinophil density compared with older patients, but the difference did not reach significance. There was no significant difference in eosinophil density in tumours with or without pelvic lymph node metastases, but tumours which subsequently developed pelvic or distant disease had a significantly lower density than the remainder.

![Figure 1](http://jcp.bmj.com/)

*Figure 1 Quantification of tissue associated tumour eosinophil density among cases, divided according to occurrence of subsequent distant metastatic disease or tumour recurrence (mean density marked with a bar).
Follow up of the series ranged from one month to 7-75 years, with a mean of 5-2 years. Thirteen women died of cervical cancer related events, the five year survival for the series being 78.9%. Six women died of other causes (other malignancies n = 3, motor vehicle accident n = 2, alcoholic liver disease n = 1); these intercurrent deaths were censored in the survival analysis. The crude five year survival rate for women with a tumour eosinophil density over 100/mm² (intense TATE) was 93% compared with 75% with a density under 100/mm², although the difference in the two survival curves was not significant (log rank χ² test 1.87, p = 0.17). When comparing the outcome of women with moderate and intense TATE (>30 eosinophils/mm²) with the remainder, however, the survival experience was significantly different (log rank χ² test 4.23, p = 0.03). The crude five year survival rate for women with a tumour eosinophil density over 30/mm² was 92% compared with 70% with a density under 30/mm² (fig 2).

The survival of women whose tumours had minimal, rather than moderate or pronounced lymphoplasmacytic stromal inflammatory response, is compared in fig 3. While there seems to be a small difference in five year survival rates, the survival curves for these two groups of women were not significantly different (log-rank χ² 0.64, p = 0.42).

Table 2. Summary of previous work on survival of women with cervical carcinomas demonstrating intense stromal eosinophilia.

<table>
<thead>
<tr>
<th>Author/ year</th>
<th>No (%) with intense stromal eosinophilia</th>
<th>Survival with intense stromal eosinophilia</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ayhan et al 1992*</td>
<td>29 **(26)</td>
<td>5 y survival 86%</td>
<td>No significant effect of intense TATE on survival</td>
</tr>
<tr>
<td>Lowe 1988†</td>
<td>34† (3.3)</td>
<td>5 y survival 100%</td>
<td>Suggestion of improved outlook compared with a general unselected series</td>
</tr>
<tr>
<td>(i) UK</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(ii) Malawi</td>
<td>13† (3.0)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kapp &amp; Volsi 1983*</td>
<td>14‡ (3.4)</td>
<td>11/12 (92%) stage IB/2A free of disease</td>
<td>Earlier stage disease and suggestion of improved survival with intense TATE</td>
</tr>
<tr>
<td>Bostram and Hart 1981!</td>
<td>3‡ (4.5)</td>
<td>4/4(100%) stage IB free of disease</td>
<td>Survival with intense TATE comparable with cervical cancer in general</td>
</tr>
<tr>
<td>(i) Connecticut</td>
<td>5** (4.3)</td>
<td>4/5 (80%) alive at 5 y</td>
<td>No effect of intense TATE on survival</td>
</tr>
<tr>
<td>Sridhu et 1970†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ballar et al 1966†</td>
<td>33** (11.9)</td>
<td>5 y survival 49% of 51% for remainder of series</td>
<td>Suggestive but not significant improvement in survival with intense TATE</td>
</tr>
<tr>
<td>(i) South West England</td>
<td>16** (4.5)</td>
<td>5 y survival 56% of 40% for remainder of series</td>
<td></td>
</tr>
</tbody>
</table>

*subjective grading into mild, moderate, and intense eosinophilic infiltrate.
†stromal eosinophils > 100 / mm².
‡stromal eosinophils > 25% of stromal inflammatory cells.
**subjective grading into "conspicuous" and "inconspicuous" categories.
***"heavy stromal eosinophilia"—not otherwise defined.

Figure 2. Five year Kaplan-Meier actuarial survival curves for women with stage IB cancer of the uterine cervix, stratified according to degree of tumour associated tissue eosinophilia.

Figure 3. Five year Kaplan-Meier actuarial survival curves for women with stage IB cancer of the uterine cervix, stratified according to degree of stromal lymphocytic or plasmacytic inflammatory cell infiltration.

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Eosinophils were characterised over 140 years ago and their presence in tissue tumour sections has been described in case reports since the turn of the century. This association was distinguished from tumour related blood eosinophilia, characteristic of a number of carcinomas and soft tissue tumours, and usually associated with late stage disease and widespread metastases. We now recognise that tumour related blood eosinophilia and tumour stromal eosinophilia are often independent events and that while the former is associated with a poor outlook, the latter is sometimes a predictor of a good outcome.

Intense TATE has been extensively reported in cervical carcinoma, usually associated with invasive squamous cell carcinomas of the large non-keratinising type. The proportion of such tumours is variably reported to range from 3% to 26%, depending on the criteria used to define "intense" eosinophilic stromal infiltration. As demonstrated in this series, the use of special techniques identifies a greater proportion of tumours with moderate TATE, which may not be conspicuous in routinely stained sections, especially if eosinophils are admixed with a dense infiltrate of lymphocytes or plasma cells.

Previous workers have investigated the prognostic effect of intense TATE in cervical carcinomas. The results of recent series are summarised in table 2. Because of the small number of cases with intense TATE in most series, formal survival analysis techniques have not been applied. Instead, comments on the prognostic effect of intense TATE have been confined to comparing, either with or without a parametric statistical test, the crude five year survival rate of this group with the remainder. Three of the six studies undertaken since 1965 suggest that intense tumour stromal eosinophilia is associated with an improved crude five year survival, the remainder concluding there is no survival advantage. The current study is the first to use formal survival analysis techniques combined with quantification of the stromal eosinophilia, demonstrating a significantly improved survival associated with lesser degrees of eosinophilic infiltration than assessed before.

In our data there is, not surprisingly, a strong correlation between the degree of stromal eosinophil and lymphoplasmacytic cell infiltration (table 1). This latter variable has been shown to be associated with improved survival and there may be some confounding by this effect in our eosinophil survival data. In our data, however, an eosinophilic infiltrate predicts a significantly improved survival, while a lymphoplasmacytic infiltrate does not, although the former variable was assessed more rigorously than the latter. Eosinophils kill a wide range of helminth parasites, especially in their larval stage, and there is a body of experimental work which shows that eosinophils can kill tumour cells in vitro, possibly through the release of cationic proteins and the generation of toxic hypohalonic acids by eosinophil peroxidase.

In addition, eosinophils synthesize CD4 and HLA-DR and may act as antigen presenting cells and stimulate local immune reactions. Furthermore, stromal eosinophilia potentiates the effect of radiation treatment. Dalal and coworkers have shown that a cervical tumour stromal eosinophilia of over 20 cells/mm² is associated with a significantly better tumour radiation response, as defined by a greater than 50% reduction in tumour size following treatment. They hypothesised this may be an effect of the induction of tissue oedema by eosinophil products, with enhancement of radiation damage through increased free radical formation.

The factors which determine the variability in tumour stromal eosinophilia are unknown. In this study no difference was noted in eosinophil density between small and large tumours, defined either by surface dimension or depth of invasion. Tumour necrosis, as exemplified by the phenomenon of post irradiation eosinophilia, may be a factor, although tumour necrosis was not a conspicuous feature in most cases examined in this series. Recent interest has centred on variable tumour oestrogen receptor phenotypes, with one receptor type appearing to show increased binding to eosinophils. Further work to define this and other cytokine mediated differences in cervical eosinophil recruitment is needed.

Many factors have been identified as having prognostic value in women with low stage cervical carcinoma, including tumour size, depth of invasion, lymphatic spread, vascular space invasion, occult tumour mucin production and local recurrence following primary treatment. The current study has not examined stromal eosinophilia in relation to these other variables, an exercise which would require larger numbers for meaningful multivariate analysis. But we have shown that, as a univariate parameter, a stromal eosinophilia of at least modest proportions is associated with a significantly improved survival in stage IB carcinomas. We do not advocate formal quantification of tumour stromal eosinophil infiltration in routine assessment of invasive cervical carcinomas. Our current practice is to examine several high magnification fields in haematoxylin and eosin stained sections, and where more than a scattered eosinophil is present, roughly corresponding to at least our moderate TATE group, we append a comment to the histology report along with the other routinely described prognostic indices.


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doi: 10.1136/jcp.46.11.1016

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