Langerhans cells (LC) are an essential component of cutaneous immunological defence mechanisms (Halliday & Muller, 1984). They are bone-marrow derived cells (Stingl et al., 1980), which in the epidermis form a continuous network of cells linked to each other via their dendritic processes (Halliday et al., 1986). The role of this network is unknown, but it presumably aids the trapping of foreign antigens as these cells have been demonstrated to bind epidermal antigens (Shelley & Juhlin, 1977). Following antigen-binding LC migrate via dermal lymphatics to the local lymph nodes where they function as antigen-presenting cells, thereby initiating an immune response against the antigen (Silberberg-Sinakin & Thorbecke, 1980; Streilein & Bergstresser, 1980; Stingl et al., 1978).

Since LC link the epidermis to the systemic immune system, they may be an important component of immunological defence against skin tumours. Recent observations of increased numbers of LC in human skin tumours (McArdle et al., 1986) support such a role for LC. We have also demonstrated that the chemical carcinogen 7,12-dimethylbenz(a)anthracene (DMBA) depletes LC from mouse skin during the period while tumours became macroscopically visible (Muller et al., 1985). In these experiments some of the tumours regressed as the LC repopulated the epidermis.

Some chemical carcinogens induce tumour growth by themselves e.g. DMBA, whereas there are other chemicals which are not complete carcinogens, and these have been divided functionally into initiators and promoters. The two-stage model of carcinogenesis where treatment with both an initiator and a promoter are required to induce tumour growth has been reviewed by Slaga (1984). The initiation phase is an irreversible event, requiring a single application of the initiator, while promotion is reversible, with repeated treatments required, which may be delayed for up to one year following initiation (Slaga, 1984). In this investigation, the two-stage model of carcinogenesis in mouse skin has been utilized to further define the role of LC in carcinogenesis.

BALB/c mice were treated with either 1, 2, or 3 weekly topical applications of 20 μl promoter or initiator to the dorsal surface of each ear. The promoters assessed were 0.1% croton oil (Sigma, Lot 43F-0415; Roe & Peirce, 1961); 0.005% 12-O-tetradecanoylphorbol 13-acetate (TPA, Sigma, Lot 34F-0682; Verma et al., 1978); and 0.005% teleocidin (a gift from Dr Fujiki; Fujiki & Sugimura, 1983), the vehicle in each case being acetone. The initiators used were 10% urethane (Sigma, Lot 102F-0300) in acetone (Graff et al., 1953); 0.5% chrysene (Sigma, Lot 84F-3397) in equal parts of lanoline and liquid paraffin (Scribner, 1973); and 0.25% benz(a)anthracene (Sigma, Lot 129C-0520) in acetone (Scribner, 1973). Controls were treated with acetone alone.

One week following the final treatment, mice were killed by cervical dislocation and their ears were excised for LC quantitation in epidermal sheets by adenosine triphosphatase (ATPase) staining as described previously (Halliday et al., 1986). LC were visualised by light microscopy, and the numbers present in 6 fields were counted for each ear per mouse. The size of the field was determined using a graticule, and the number of LC mm⁻² of epidermis calculated for each mouse. A total area of 8.15 mm² was counted per mouse.

By light microscopy, LC were observed in the epidermis of control BALB/c mouse ears as brown ATPase-positive cells linked to each other via dendritic processes. The LC density was within the range 345–554 cells per mm², which is similar to that previously observed in the dorsal trunk of this mouse strain (Muller et al., 1985). There was no difference in LC density between the control group of mice treated with solvent for 1 week, and the groups treated with the tumour initiators urethane, chrysene, or benz(a)anthracene for 2 or 3 consecutive weeks still did not cause any alteration in the number or morphology of ATPase-positive LC; in contrast the tumour promoters croton oil, TPA, and teleocidin significantly decreased the LC to levels which were similar to those observed after treatment for 1 week (Table I). The tumour initiators examined have no promoter activity (Slaga et al., 1982), and were used at concentrations which have previously been shown to be effective in the two-stage model of tumour-induction (Graff et al., 1953; Scribner, 1973). Thus LC are affected by tumour promoters, but not initiators.

LC were also identified using a Philips 410 electron microscope based on their well characterised ultrastructure and presence of the unique Birbeck granule (Birbeck et al., 1961). LC were frequently observed in control, urethane, chrysene, and benz(a)anthracene-treated epidermis. These initiators did not discernibly alter LC ultrastructure. In contrast, LC were difficult to find in promoter-treated skin; e.g. in one specimen only a single LC was observed in the epidermis of a croton oil-treated mouse, and this showed no features of ultrastructural damage. It has been demonstrated by electron microscopy that under some circumstances the ATPase marker may be modulated from the LC plasma membrane without depleting the cells from the epidermis (Aberer et al., 1981). However, electronmicroscopic examination of promoter and initiator-treated skin confirmed our results obtained by ATPase staining, indicating that the tumour promoters had not modulated ATPase from the LC surface, but had depleted these cells from the epidermis.

LC took more than 6 weeks to return to control values after croton oil treatment (Table II) which is similar to the 8 week time period we have previously observed for LC repopulation of DMBA-treated epidermis (Muller et al., 1985). This long recovery time provides further support that promoters deplete LC from the epidermis rather than modulating ATPase from the plasma membrane. Fünten & Fünten (1983) found that the critical effects of tumour promoters last for at least 2 months in mouse epidermis. As this is similar to the time LC remain depleted

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SHORT COMMUNICATION

Tumour promoters but not initiators deplete Langerhans cells from murine epidermis

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Table I  Langerhans cell densities* in murine epidermis treated weekly with tumour promoters or initiators

| Treatment of mice | 1 Week | 2 Weeks | 3 Weeks |
|-------------------|--------|---------|---------|
|                   | Mean Langerhans cells mm\(^{-2}\) (range) | Mean Langerhans cells mm\(^{-2}\) (range) | Mean Langerhans cells mm\(^{-2}\) (range) |
| Control\(^b\)     | 386 (350-415) – 8 | 385 (345-435) – 8 | 422 (345-530) – 8 |
| Promoters:        |        |         |         |
| croton oil        | 123 (60-235) <0.01 6 | 146 (110-190) <0.01 6 | 132 (105-165) <0.01 6 |
| TPA               | 179 (140-195) <0.01 6 | 140 (105-230) <0.01 6 | 157 (125-200) <0.01 6 |
| teleocidin        | 156 (120-215) <0.01 6 | 129 (110-160) <0.01 6 | 116 (50-175) <0.01 6 |
| Initiators:       |        |         |         |
| urethane          | 380 (310-465) NS 6 | 384 (355-460) NS 6 | 399 (450-495) NS 6 |
| chrysenes         | 395 (355-440) NS 6 | 426 (350-550) NS 6 | 450 (356-560) NS 6 |
| benzo(a)anthracene| 394 (335-425) NS 6 | 453 (393-550) NS 6 | 486 (335-570) NS 6 |

* Determined by staining for ATPase; \(^b\) Treated with acetone alone; \(^c\) Statistical comparison with controls (unpaired Wilcoxon rank sum test; Sokal & Rohlf, 1969); NS: not significant; \(n\): number of mice in group.

Table II  Langerhans cell repopulation following depletion by 3 weekly treatments with croton oil

| Time since final croton oil treatment | Mean Langerhans cells mm\(^{-2}\) (range)* | \(p\) | \(n\) |
|--------------------------------------|----------------------------------------|------|------|
| 1 week                               | 137 (89-179) <0.005 6                  |      |      |
| 3 weeks                               | 231 (179-264) <0.005 5                 |      |      |
| 6 weeks                               | 271 (237-293) <0.005 6                 |      |      |
| Controls\(^a\)                        | 323 (285-384) – 6                     |      |      |

* Determined by staining for ATPase; \(^a\) Statistical comparison with controls (unpaired Wilcoxon rank sum test; Sokal & Rohlf, 1969); \(^b\) Treated with acetone alone; \(n\): number of mice in group.

Following croton oil treatment, LC depletion may be one of the critical steps in tumour promotion.

Croton oil, the first tumour promoter to be discovered (Berenblum, 1941), has been thoroughly investigated and found to be a strong promoter with very little, if any, initiating potential (Klein-Szanto, 1984). It is however a multicomponent mixture of lipids, of which a series of eleven phorbol diesters have been found to be active tumour promoters (Hecker, 1968). The most potent tumour promoter of these phorbol diesters, TPA, was observed to deplete LC from the epidermis in the present study. As croton oil and TPA depleted LC to similar levels (\(p\) not significant) it is likely that the croton oil-mediated depletion of LC was caused by TPA in the croton oil. However a cumulative effect involving the other active phorbol diesters cannot be excluded. Teleocidin, an indole alkaloid, is structurally unrelated to TPA, but is a potent tumour promoter which lacks initiator activity (Fujiki & Sugimura, 1983). As these chemically unrelated promoters have similar effects on LC, LC-depletion may be a general step in the process of tumour promotion in the skin.

The tumour promoter-induced depletion of LC demonstrated in this study has important implications for understanding the process of tumour growth. LC are an essential component of cutaneous immunological defence mechanisms (Halliday & Muller, 1984), and therefore any potential tumour cells may be inhibited from growing into a tumour by the LC presenting tumour-associated-antigens to T cells, thus inducing an anti-tumour immune response. Depletion of LC by a promoter might enable potential tumour cells to grow into a tumour unhindered by an immune response. We have previously shown that sensitization of mice with di-nitrofluorobenzene through skin depleted of LC by treatment with the complete carcinogen DMBA activates specific suppressor T lymphocytes which inhibit subsequent attempts to induce immunity against this antigen (Halliday & Muller, 1986). Hence, upon recovery of LC from the effects of tumour promoters, they may be unable to activate immune defence mechanisms due to the presence of specific suppressor T cells.

Electromicroscopy failed to reveal any degenerating LC in tumour promoter-treated skin. Therefore, it is likely that tumour promoters do not destroy LC but induce their migration from the epidermis. TPA has been shown to activate other cells of the immune system; it modulates the T4 antigen from T lymphocytes (Solbach, 1982), collaborates with anti-T3 antibodies to cause activation and proliferation of T lymphocytes (Hara & Fu, 1985), and can substitute for macrophages during mitogen activation of T lymphocytes (Rosenstreich & Mizel, 1979). Likewise, tumour promoters may activate LC to migrate from the epidermis.

It is concluded that while transformed cells may be inhibited from growing into a tumour by an immune response mounted against tumour-associated-antigens presented via LC to T lymphocytes, depletion of the LC by promoters would abrogate this response, thus enabling the transformed cell to grow unhindered. However, this may not be the only effect of tumour promoters on anti-tumour immunity as TPA has also been shown to suppress macrophage and NK cell tumour cytotoxicity (Keller, 1979). In contrast to our findings with tumour promoters, tumour initiators had no effect on LC. Whether alteration of local antigen-presenting cells occurs in other models of chemical carcinogenesis which involve multiple steps, such as in the liver (Farber, 1984), is unknown, but such an investigation would determine if this is a requirement for tumour growth at other sites.

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