Induction of reactive oxygen intermediates in human monocytes by tumour cells and their role in spontaneous monocyte cytotoxicity

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Summary The present study examined the ability of human monocytes to produce reactive oxygen intermediates after a contact with tumour cells. Monocytes generated oxygen radicals, as measured by luminol-enhanced chemiluminescence and superoxide anion production, after stimulation with the tumour, but not with untransformed, cells. The use of specific oxygen radical scavengers and inhibitors, superoxide dismutase, catalase, dimethyl sulphoxide and deferoxamine as well as the myeloperoxidase inhibitor 4-aminobenzoic acid hydrazide, indicated that chemiluminescence was dependent on the production of superoxide anion and hydroxyl radical and the presence of myeloperoxidase. The tumour cell-induced chemiluminescent response of monocytes showed different kinetics from that seen after activation of monocytes with phorbol ester. These results indicate that human monocytes can be directly stimulated by tumour cells for reactive oxygen intermediate production. Spontaneous monocyte-mediated cytotoxicity towards cancer cells was inhibited by superoxide dismutase, catalase, deferoxamine and hydrazide, implicating the role of superoxide anion, hydrogen peroxide, hydroxyl radical and hypohalite. We wish to suggest that so-called ‘spontaneous’ tumoricidal capacity of freshly isolated human monocytes may in fact be an inducible event associated with generation of reactive oxygen intermediates and perhaps other toxic mediators, resulting from a contact of monocytes with tumour cells.

Keywords: monocyte; tumour cells; interactions; reactive oxygen intermediates; cytotoxicity

Macrophages are prominent in the cellular infiltrates surrounding the tumour and can constitute more than 50% of the total tumour mass (Leek et al, 1996). Mononuclear cell infiltration is regarded as a manifestation of the host response against the growing tumour (Zembala and Buckle, 1989). The anti-tumour effect of macrophages may be due to their cytotoxic/cytostatic activity, in which several toxic mediators including tumour necrosis factor α (TNF-α), reactive nitrogen intermediates (RNI) and reactive oxygen intermediates (ROI) are involved. The presence of TNF-mRNA has been demonstrated in tumour infiltrating macrophages (TIM), which is taken as evidence for the local production of this and perhaps other cytokines with anti-tumour activities (Beissert et al, 1989). In humans, there is no evidence for the local production of oxygen radicals in the tumour bed, however, the role of ROI in the anti-tumour response in humans has been indirectly implicated by observations that myeloperoxidase-deficient individuals have an increased incidence of neoplasms (Lanza et al, 1987). The direct evidence for the importance of ROI in the killing of tumour cells comes from the cell-free systems generating hydrogen peroxide or superoxide anion (O₂⁻) (Ioannidis and de Groot, 1993; Palomba et al, 1996). TIM from mouse mammary carcinomas are able to produce ROI (Mantovani et al, 1992). An increased production of ROI by monocytes was observed in patients with cancer (Trulson et al, 1988, 1994), but the stimulus responsible for their generation has not been established. In contrast to murine macrophages, freshly isolated human monocytes exhibit considerable spontaneous cytotoxicity towards tumour cells (Davies and Edwards, 1992; Martin and Edwards, 1993), but its origins remain largely unclear. Davies and Edwards (1992) demonstrated that cytotoxic factors were released by monocytes cultured with K562 tumour cells, whereas monocytes and tumour cells cultured alone released factor(s) which promoted K562 cells growth. Furthermore, inhibitors of ROI added to K562 cells enhanced their growth. This led us to explore the possibility that spontaneous cytotoxic activity of monocytes may, at least partly, be due to ROI production induced by tumour cells in the coculture.

The present study shows that stimulation of monocytes with tumour, but not untransformed, cells induces the production of ROI, as determined by the generation of chemiluminescence (CL) and O₂⁻. The CL response was associated with O₂⁻ and OH⁻ production and was dependent on the presence of myeloperoxidase. This study also demonstrates that O₂⁻, hydrogen peroxide, OH⁻ and probably hypohalite(s) are involved in monocyte-mediated spontaneous cytotoxicity. This may imply that this ‘spontaneous’ cytotoxicity is probably an inducible event that occurs after contact of monocytes with tumour cells.

MATERIALS AND METHODS
Isolation of cell populations

Human peripheral blood mononuclear cells were isolated from EDTA–blood of healthy donors on a standard Ficoll/Isopaque
Monocytes were separated from mononuclear cells by counterflow centrifugal elutriation with a JE-5.0 elutriation system equipped with a 5-mL Sanderson separation chamber (Beckman, Palo Alto, CA, USA) as previously described (Zembala et al., 1994a). Isolated monocytes were 90–96% pure, as judged by flow cytometry analysis (FACScanV, Becton-Dickinson, Mountain View, CA, USA), using anti-CD14 (Leu-M3; Becton-Dickinson) monoclonal antibody (Zembala et al., 1994a) and contained 2–4% NK cells, as judged by staining with anti-CD56 monoclonal antibody (Dako, Glostrup, Denmark). The cells were suspended in culture medium RPMI-1640 (Biochrom, Berlin, Germany) with gentamycin (25 μg·ml⁻¹, Biochrom), glutamine (2 mm, Gibco, Paisley, UK) and 10% fetal calf serum (FCS) (Biochrom).

The following human cell lines were used: HPC-4 (human pancreatic adenocarcinoma) (Siedlar et al., 1995), CaOV (ovarian carcinoma), normal human skin fibroblasts, peritoneal mesothelial cells (obtained in our laboratory), and established cell lines DeTa (colon carcinoma), Hu 1703 (transitional cell bladder carcinoma), BC3726 (uroepithelium) (Zembala et al., 1994a), Hep G2 (hepatocellular carcinoma) and Caco-2 (colon adenocarcinoma) were obtained from Professor V Colizzi (II Università di Roma, Italy). Cell lines carcinomas) and Caco-2 (colon adenocarcinoma), normal human skin fibroblasts, peritoneal mesothelial cells (obtained in our laboratory), and established cell lines DeTa (colon carcinoma), Hu 1703 (transitional cell bladder carcinoma), BC3726 (uroepithelium) (Zembala et al., 1994a), Hep G2 (hepatocellular carcinoma) and Caco-2 (colon adenocarcinoma) were obtained from Professor V Colizzi (II Università di Roma, Italy). Cell lines were regularly tested for Mycoplasma sp. contamination using standard PPLO medium (Difco, Detroit, MI, USA).

Chemiluminescence

CL was used to detect activated oxygen species (Ernst et al., 1984). Various tumour cells/monocytes ratios were used to establish conditions for optimal CL response. As a result, 1×10⁴ monocytes and 5×10⁵ tumour cells were resuspended in 100 μl of culture medium and added to the vials (Berthold, Vienna, Austria). Then, 300 μl of 2 mM luminol (5-amino-2,3-dihydro-1,4-phthalazine-dione, Sigma, St. Louis, MO, USA) in Krebs-Ringer buffer with Mg²⁺ and Ca²⁺ was added. The vials were placed in the measuring chambers of a Multi-Biolumat (Berthold), kept at 37°C and the CL response was continuously recorded. The results were calculated as integrals (cumulative counts, cc) of the response recorded during 300 min. Data were expressed as the percentage of control response or CL index, calculated as:

$$\text{CL index} = \frac{\text{cumulative counts of tumour-stimulated monocytes}}{\text{cumulative counts of unstimulated monocytes}}$$

In some experiments, phorbol myristate acetate (PMA) (100 ng·ml⁻¹, Sigma) was added to monocytes or tumour cell suspension and the CL response was recorded.

Measurement of O₂⁻ production

Superoxide production was determined by the cytochrome c reduction assay as described by Pick and Mizel (1981). Briefly, monocytes (1×10⁴ per well) and tumour cells (1×10⁴–3×10⁵ per well) were resuspended in 160 μM solution of ferricytochrome c (Sigma) in phenol-red-free Hanks’ balanced salt solution (HBSS), and plated in 96-well flat bottom tissue culture plates (Nunc, Roskilde, Denmark) in a final volume of 100 μl per well. Wells supplemented with 300 U ml⁻¹ of superoxide dismutase (SOD) (Sigma) were used as blanks. Cytochrome c reduction was measured after 2 h of incubation at 550 nm in an Elisa reader (Labsystems Multiscan Plus, Labsystems, Finland) using a 450-nm reference filter (Martin and Edwards, 1993). Results were expressed as nM per 1×10⁵ cells per 2 h.

Determination of hydrogen peroxide production

Hydrogen peroxide detection was based on the horseradish peroxidase-dependent oxidation of phenol red, as described by Pick and Mizel (1981). Briefly, cells were washed and resuspended in assay solution containing 0.56 mM of phenol red (Sigma) and 20 U ml⁻¹ horseradish peroxidase (Sigma) in phenol-red-free HBSS, and seeded in tissue culture plates in a final volume of 100 μl per well. After incubation for 2 h, the reaction was stopped by the addition of 10 μl of 1 N sodium hydroxide per well and absorbance was read at 600 nm. Wells with sodium hydroxide added at the beginning of testing were used as blanks. Results were expressed as nM per 1×10⁵ cells per 2 h.

Flow cytometry and cell sorting

Monocytes (2×10⁶) were stimulated with HPC-4 cells (5×10⁶) for 40 min at 37°C. After washing, the cells were stained with anti-CD14 monoclonal antibody (Becton-Dickinson) for 20 min at 4°C. Then, cells were washed and resuspended in 5 ml of phosphate-buffered saline (PBS). Cell sorting was performed with FACS Vantage Cytometer (Becton-Dickinson) equipped with a Power Macintosh 7600/120 computer using a Cell Quest v. 3.0 software. The ion laser Innova Enterprise II (Coherent, Santa Clara, CA, USA) operating at 488 nm was used as a light source. After setting CD14⁺ (monocytes) and CD14⁻ (tumour cells) gating, sorting was performed using a 70 μm nozzle tip with a drop drive frequency of 25 kHz, three drops envelopes and ‘normal-R’ sort mode. Sorted cells were collected into water-cooled (constant temperature circulator, Neslab Instruments, Portsmouth, NH, USA) polystyrene Falcon 2057 tubes (Becton-Dickinson), precoated with FCS to avoid plastic charging and cell attachment to the well. The purity of sorted cells was checked by flow cytometry and it exceeded 95%.

Inhibitors of ROI production

The following blockers and scavengers of oxygen-free radicals (all from Sigma) were used at the indicated concentrations: superoxide dismutase (SOD, 100–600 U ml⁻¹), catalase (CAT, 100–5000 U ml⁻¹), dimethyl sulphoxide (DMSO, 50–200 mM), deferoxamine mesylate (25–200 mM) and the inhibitor of myeloperoxidase 4-aminobenzoic acid hydrazide (ABAH, 50–150 μM).

Cytotoxicity assay

To obtain a significant cytotoxicity, monocytes were cultured with tumour cells at 1:0.4 ratio (Davies and Edwards, 1992). Monocytes (5×10⁴ per well), HPC-4 (2×10⁴ per well) or their mixtures were cultured for 20 h in the presence or absence of ROI inhibitors or scavengers. Then, the culture medium was removed.
and 100 µl per well of MTT (2 mg ml⁻¹, 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide, Sigma) dye solution was added. Cell proliferation was assessed by reduction of MTT, as previously described (Zembala et al., 1993; see also Hruby and Beck, 1997). Percentage of cytotoxicity was calculated according to the formula previously described (Hruby and Beck, 1997):

\[
\frac{OD(\text{monocytes + tumour cells}) - OD(\text{monocytes alone})}{OD(\text{tumour cells alone})} \times 100
\]

The effect of inhibitors on tumour cell survival was calculated as:

\[
\frac{OD(\text{tumour cells + inhibitor})}{OD(\text{tumour cells alone})} \times 100
\]

**Statistical analysis**

Significance analysis was performed using an unpaired Student’s t-test. P-values < 0.05 were considered significant.

**RESULTS**

**Induction of chemiluminescent response of human monocytes by tumour cells**

Monocytes were incubated with various tumour cells in the presence of luminol, and the CL was recorded in a chemiluminometer. Figure 1A shows that monocytes incubated with cancer cells (HPC-4 or DeTa) gave rise to the CL response, which reached a peak around 100–120 min, whereas monocytes kept in the medium yielded negligible response. The addition of normal human uroepithelial cells induced low level of CL. The response to tumour cells was much slower than that seen after stimulation of monocytes with PMA (Figure 1B), which peaked at 3–8 min, whereas during the first 30 min monocytes stimulated with HPC-4 cells generated little CL. Also, no significant CL was recorded when unstimulated and PMA-stimulated tumour cells were used. Tumour cell lines differed in their ability to evoke CL response. As shown in Table 1, a strong response was induced by HPC-4, DeTa and Hep G2 cell lines. Other cancer cells, Caco-2 and Hu 1703, elicited a moderate CL response, whereas BC 3726 and CaOV induced a weak response. In contrast, normal uroepithelial cells, peritoneal mesothelial cells or fibroblasts (Table 1) did not stimulate significant CL. It was concluded that the CL response of monocytes was the result of their direct interaction with tumour cells, and that the response was different, but characteristic, for the given cell line used.

**Identification of cells responsible for CL generation**

Because, potentially, both monocytes and cancer cells can produce ROI, attempts were made to establish which cell type was responsible for the CL response. Monocytes were stimulated with tumour cells for 40 min, stained with anti-CD14 monoclonal antibody and sorted out into CD14⁺ and CD14⁻ populations. As control, unstimulated monocytes were also sorted (‘dummy sorting’). These procedures were timed in such a way to measure CL around the peak of response. Table 2 shows that CD14⁺ cells that were in contact with tumour cells gave a significantly higher CL response than ‘dummy sorted’ CD14⁻ cells. CD14⁺ cells showed a background CL. This clearly indicates that monocytes and not tumour cells are the source of luminol-dependent CL.

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**Figure 1** The kinetics of the CL response of monocytes stimulated with tumour cells. (A) Monocytes (1 × 10⁶) were mixed with tumour cells (5 × 10⁶) HPC-4 (pancreatic carcinoma), DeTa (colon carcinoma) and normal uroepithelial cells (HCV 29). After addition of luminol, CL was recorded over 300 min. (B) Monocytes were stimulated with PMA (1 ng ml⁻¹) or HPC-4 cells and CL recorded during 30 min. Unstimulated or PMA-stimulated HPC-4 cells were also used. Results calculated as integrals and expressed in cc (cumulative counts) are also indicated. A typical experiment, one of four, is shown.

**Table 1** Chemiluminescent response of monocytes stimulated with different cell lines

| Cell line | CL index (mean ± s.d.) | Range | Number of experiments |
|-----------|------------------------|-------|----------------------|
| HPC-4     | 26.5 ± 17.5            | 9.0–58.0 | 10                   |
| Hep G2    | 25.3 ± 6.7             | 20.7–33.0 | 4                    |
| DeTa      | 14.1 ± 9.7             | 4.7–32.5 | 7                    |
| Caco-2    | 7.2 ± 5.7              | 2.0–5.3 | 4                    |
| Hu 1703   | 6.4 ± 3.5              | 3.8–10.4 | 3                    |
| BC 3726   | 4.2 ± 3.1              | 1.3–7.6 | 3                    |
| CaOV      | 4.1 ± 0.4              | 3.7–4.5 | 2                    |
| HCV 29    | 2.7 ± 1.0              | 1.4–4.1 | 5                    |
| Mesothelial cells | 2.8 ± 0.2 | 2.8–3.1 | 2                    |
| Skin fibroblasts | 2.6 ± 0.6 | 2.0–3.3 | 4                    |

Monocytes (1 × 10⁶ cells in 50 µl of culture medium) were stimulated with tumour cells (5 × 10⁶ cells in 50 µl of culture medium) in Krebs-Ringer buffer with 2 mM luminol, and chemiluminescence (CL) was recorded during 300 min. The data were expressed as index of CL, as described in Materials and methods.
six different experiments are shown.

Table 2 Chemiluminescence of monocytes (MO) sorted after contact with tumour cells

| Cells                             | Integrals (cc) |
|-----------------------------------|----------------|
| MO, medium/unsorted               | 0.90 x 10^4    |
| MO, medium/sorted CD14+           | 0.75 x 10^4    |
| MO, tumour cells/sorted           | 7.49 x 10^4    |
| CD14-                             | 0.08 x 10^4    |

Figure 2 - Effect of ROI inhibitors on CL response of monocytes stimulated with HPC-4 cells. SOD, catalase (CAT), deferoxamine (DEF), dimethyl sulphoxide (DMSO) and ABAH were added to the mixture of monocytes and HPC-4 cells. The results are expressed as the percentage of control response of monocytes and HPC-4 cells without inhibitors. Means ± s.e. of six different experiments are shown.

The type(s) of free radical(s) generated by monocytes stimulated with tumour cells

To establish which type of ROI was associated with generation of CL by monocytes, the blockers or scavengers of free oxygen radicals were used. SOD, which dismutates O_o^- to hydrogen peroxide, deferoxamine, which prevents HO formation, and DMSO, scavenger of OH_., diminished CL response of monocytes stimulated with HPC-4 cells (Figure 2). This indicates that mainly O_o^- and OH_., were detected by luminol-dependent CL. Surprisingly, catalase, which decomposes hydrogen peroxide to water and oxygen, had no effect on CL generation, indicating that hydrogen peroxide was not a major type of ROI evoking CL. Also, no synergistic effect of SOD and catalase was observed (data not shown). The CL response was myeloperoxidase-dependent because it was markedly inhibited by ABAH. This indirectly indicated that luminol-enhanced CL of monocytes stimulated with tumour cells is dependent on O_o^- and OH_. production and the presence of myeloperoxidase.

Demonstration of O_o^2-, but not hydrogen peroxide, production by monocytes stimulated with tumour cells

The cytochrome c reduction test was used to measure the level of O_o^2- production in the culture of monocytes with tumour cells within 30–180 min. Addition of tumour cells at a low ratio (1:0.1–1:0; monocytes:tumour cells respectively) increased O_o^2- production above the background level (release of O_o^2- by unstimulated monocytes) (Figure 3), whereas the admixture of higher numbers of HPC-4 resulted in a decreased response. HCV 29 cells, incapable of stimulating CL in monocytes (Figure 1, Table 1), were used as control.

When monocytes were cultured with tumour cells for 40 min and then sorted into CD14+ and CD14- populations, it was observed that the highest amount of O_o^2- was produced by CD14+ cells (3.3 nM ± 0.08 per 1 x 10^5 cells, mean of three different experiments), whereas CD14- cells generated 1.06 nM and ‘dummy sorted’ CD14+ cells – 0.90 nM of O_o^2-. In these experiments, the mean level of O_o^2- found in the co-culture was 1.44 nM. The production of O_o^2- by CD14+ cells was probably connected with the presence of small numbers of contaminating monocytes in this population because HPC-4 cells alone, unstimulated or stimulated with PMA (Figure 1B), LPS or fMLP (data not shown) did not produce O_o^2-. This again clearly points to CD14+ cells as the source of O_o^2- in the co-culture. In contrast, hydrogen peroxide was spontaneously produced by both monocytes and, even more, by tumour cells. The level of hydrogen peroxide was not increased in cocultures (data not shown), in keeping with the absence of catalase effect on the generation of CL by monocytes.

The effect of ROI scavengers on the inhibition of tumour cell growth by monocytes

Freshly isolated human monocytes exhibit considerable cytotoxicity towards tumour cells. To assess a possible role of ROI in this phenomenon, several ROI inhibitors were added to monocytes cultured with HPC-4 tumour cells at the ratio 1:0.4. Under these conditions, cytotoxicity, as assessed by the MTT test, in untreated cultures was approximately 33%. It was significantly lower when cells were treated with SOD, catalase and deferoxamine (Figure 4A). Also, ABAH diminished cytotoxicity. These inhibitors had no effect on the proliferation of tumour cells cultured alone (Figure 4B). This implies that hydrogen peroxide, O_o^2- and OH_., were involved in monocyte-mediated cytocidal activity.

DISCUSSION

The generation of ROI is thought to be one of the mechanisms responsible for the anti-tumour effect of activated macrophages (Ding et al, 1988; Jaattela and Wissing, 1993). It is also proposed that the cytotoxic effect of TNF produced by monocytes is due to the generation of ROI within target cells, which in turn activate...
values of three independent experiments

HPC-4 cells cultured alone (monocytes (5 × 10^4 per well)) and HPC-4 cells (2 × 10^5 per well) (A), or to HPC-4 cells cultured alone (B). After 24 h of incubation, MTT was added for a further 2 h as described in Materials and methods. The results are expressed as the percentage of growth inhibition of tumour cells by monocytes (A) or tumour cell survival (B). The data represent the mean values of three independent experiments ± s.e.

Figure 4 Effect of ROI inhibitors on the growth of HPC-4 cells cultured with or without monocytes. SOD (300 U ml⁻¹), catalase (CAT, 500 U ml⁻¹), deferoxamine (DEF, 100 μM) and ABAH (100 μM) were added to co-cultures of monocytes (5 × 10^4 per well) and HPC-4 cells (2 × 10^5 per well) (A), or to HPC-4 cells cultured alone (B). After 24 h of incubation, MTT was added for a further 2 h as described in Materials and methods. The results are expressed as the percentage of growth inhibition of tumour cells by monocytes (A) or tumour cell survival (B). The data represent the mean values of three independent experiments ± s.e.

pre-existing toxic mediators, e.g. proteases (Wong and Goeddel, 1989). There is also direct evidence that generation of O₂⁻ and hydrogen peroxide in cell-free systems causes tumour cell killing (Ioannidis and de Groot, 1993; Palomba et al, 1996). In humans, there is indirect evidence that spontaneous cytotoxicity of monocytes towards tumour cells may be associated with their ability to produce ROI (Davies and Edwards, 1992; Martin and Edwards, 1993). Although an altered production of ROI by monocytes of cancer patients has been observed (Trulson et al, 1988, 1994; Hara et al, 1991), it is unclear which cells, and how, are stimulated for ROI production.

The present studies addressed the question of whether monocytes can be directly activated by tumour cells for ROI generation. We are aware of only one report on the ability of K562 (chronic myelogenous leukaemia) cells to stimulate peripheral blood mononuclear cells for a CL response (Ernst et al, 1984). In this report, monocytes were indirectly implicated as a source of CL, because phagocyte-depleted mononuclear cells were poor producers. Our study clearly demonstrates that monocytes and not tumour cells are the source of CL because CD14⁺, but not CD14⁻, cells sorted out from the co-culture produced CL and released O₂⁻. The role of natural killer (NK) cells in the tumour cell-induced CL response is unlikely because contamination of elutriated monocytes with CD56⁺ cells is low, approximately only 2% CD14⁺ cells show expression of CD56 (Rothe et al, 1996), and, finally, cell populations highly enriched for NK activity do not respond in target cell- and latex-induced CL (Ernst et al, 1984). Tumour cell-induced CL response peaked around 2 h. These kinetics are distinct from the rapid CL response seen after the activation of monocytes with PMA (Figure 1B), lipopolysaccharide (LPS), latex or zymosan, which lasts for 1–15 min (Trulson et al, 1988; Martin and Edwards, 1993). Monocytes respond poorly to normal cells, such as skin fibroblasts, uroepithelial and mesothelial cells, compared with cancer cells. This may suggest that monocytes are able to discriminate between transformed and normal cells. This is in keeping with previous findings regarding the ability of monocytes to release TNF and nitric oxide after stimulation with cancer, but not untransformed, cells (Zembala et al, 1994a, b; Hasday et al, 1990), and other observations on the capacity of activated monocytes to lyse tumour, but not non-neoplastic, cells (Galligioni et al, 1993). The molecular mechanism(s) by which monocytes can distinguish cancer cells remains largely unknown, but the role of phosphatidylserine (Utsugi et al, 1991) or hyaluronic acid (Pericle et al, 1996) on tumour cells and CD44 determinants on monocytes (Zembala et al, 1994b) have been implicated.

Why various tumour cells possess different capacities to induce the CL response is unclear. Distinct surface structures of tumour cells were found to be responsible for monocyte activation for TNF production (Jänicke and Männel, 1990), but their molecular structure has not been identified. Our preliminary observations indicate that the presence of some adhesion molecules, including CD44, and expression of hyaluronic acid on tumour cells, which is known to vary considerably (van Muijen et al, 1995), may be important.

CL is commonly used for the detection of ROI production. However, there is no agreement on the type(s) of the oxygen species responsible for generation of CL. Hydrogen peroxide, O₂⁻, O₃⁻ (singlet oxygen) or OH⁻ is generated during oxidative burst. The decrease of tumour cell-induced CL after treatment with deferoxamine or DMSO points to the role of OH⁻. In the present system, CL was dependent on the presence of myeloperoxidase, as shown by the inhibitory effect of ABAH. The inhibition of CL by SOD indicated that also O₂⁻ is involved. Hydrogen peroxide, O₂⁻, and myeloperoxidase are implicated in CL generated by monocytes stimulated with PMA and fMLP (McNally and Bell, 1996). However, in our hands, CL induced by tumour cells was not dependent on hydrogen peroxide generation because catalase had no effect. Hence, tumour cell-induced CL response of monocytes may be distinct from that induced by PMA or fMLP because of different kinetics and range of oxygen species generated.

The involvement of O₂⁻ in the CL response was proven by a direct demonstration of O₂⁻ production by monocytes stimulated with cancer but not normal epithelial cells. Surprisingly, this production occurred after stimulation of monocytes with rather low numbers of tumour cells. The increased proportion of the latter resulted in a smaller O₂⁻ release, which can be due to a negative (inhibitory or inactivating) influence of tumour cells. It is known that cancer cells may produce several factors that inactivate toxic mediators, including SOD (Sun et al, 1989). Transforming growth factor β (TGF-β), produced by tumour cells, may also be responsible for the inhibition of O₂⁻ release because it blocks signalling pathways, including protein tyrosine kinase (Pazdrak et al, 1995) which is involved in signal transduction for ROI generation (Mytar et al, 1998). Studies are underway to define inhibitory factors of tumour cells which may down-regulate the response of monocytes.

In contrast to murine macrophages, freshly isolated human blood monocytes exhibit considerable spontaneous cytotoxicity towards tumour cells (Davies and Edwards, 1992), and there is indirect evidence suggesting that ROI may be involved (Martin
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and Edwards, 1993). On the basis of our study, it appears that this ‘spontaneous’ cytotoxicity of monocytes may in fact be due to induction of production of several toxic molecules after contact with tumour cells. As a result, generation of TNF (Hasday et al, 1990; Zembala et al, 1994b), RNI (Zembala et al, 1994a) and ROI (this study) may be initiated and be responsible for the inhibition of proliferation and/or destruction of tumour cells by previously non-activated monocytes. The present study indicates that O2, hydrogen peroxide and OH are involved in monocyte-mediated cytotoxicity towards tumour cells. This conclusion is based on the effect of specific ROI inhibitors. Because these inhibitors had no effect on proliferation of tumour cells in the absence of monocytes, it is likely that they down-regulated ROI production by monocytes. The effect of catalase, the specific scavenger of hydrogen peroxide, is somewhat confusing. Because highly toxic hypohalites and OH are generated from hydrogen peroxide (King et al, 1997), the question arises whether inhibitory action of catalase on the cytocidal activity of monocytes may be due to decreased generation of these toxic molecules. The role of hypohalites may be supported by the inhibitory effect of ABAH, which by blocking myeloperoxidase activity prevents their generation. It remains unclear whether all above radicals are effector molecules or whether at least some of them (O2, hydrogen peroxide) act mainly as precursors of other toxic intermediates.

The ability of various tumour cell lines to evoke a CL response of monocytes may perhaps explain why in different systems cytotoxicity of monocytes was found to be ROI-dependent or ROI-independent (Nakabo and Pabst, 1997). The involvement of ROI in the cytotoxic effect of Bacille Calmette-Guérin (BCG) - or PMA-activated murine macrophages (Nathan et al, 1979; Nathan and Cohn, 1980) and LPS- or PMA-activated human monocytes (Klassen and Sagone, 1980; Mavier and Edgington, 1984; Martin and Edwards, 1993; McLachlan et al, 1995) is well established. The present data suggest that contact with tumour cells may act to cytocidal capacity of monocytes and generation of ROI. Hence, ‘spontaneous’ monocyte cytotoxicity may, in fact, be an inducible event. The question arises whether TIM can also be stimulated by cancer cells for generation of ROI in the tumour bed.

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