The role of catalytic iron in asbestos induced lipid peroxidation and DNA-strand breakage in C3H10T\textsuperscript{1/2} cells

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Summary

The involvement of catalytic iron in \textit{in vitro} activities of crocidolite asbestos has been investigated. Exposure of C3H10T\textsuperscript{1/2} cells to either the UICC crocidolite standard reference sample or a non fibrous (milled) derivative resulted in an increase of thiobarbituric acid reactive substances. This catalytic activity was inhibited by pretreatment with the iron chelator desferrioxamine. The effect of this activity on cellular DNA was measured in an assay based on the production of DNA-strand breaks. Increased levels of DNA-strand breaks were detected in cultures treated with both the milled and UICC crocidolite. Inclusion of desferrioxamine with the asbestos inhibited DNA-strand breakage. It is concluded the catalytic iron present on the dust is capable of damaging both lipid and DNA and that this could be an important mechanism in asbestos pathogenicity.

Asbestos has been shown to increase the incidence of malignant mesothelioma and bronchogenic carcinoma following human exposure to fibre (Wagner et al., 1966; Selkoff & Lee, 1980). Investigations on experimental animals have also shown that asbestos minerals induce mesotheliomas when injected or implanted into the pleural cavity (Wagner et al., 1973; Stanton et al., 1977). The main conclusion from these studies was that the size and shape of the asbestos fibres were the most important factors in determining the incidence of mesothelioma. The exact mechanism of asbestos toxicity is not known and has been further complicated by conflicting reports in the literature of its activity in systems designed to detect genotoxicity.

In gene mutation assays asbestos has produced negative results (Chamberlain & Tarmy, 1977; Reiss et al., 1982) while in one study it was found to be weakly mutagenic towards the hypoxanthine-guanine phosphoribosyl transferase (HGPRT) locus of Chinese hamster lung cells (Huang, 1979). Similarly, other investigators have reported increases in sister chromatid exchanges (Livingston et al., 1980; Babu et al., 1981), while others have found no increase (Price-Jones et al., 1980). Asbestos fibres have been shown to cause morphological transformation in some systems, for example Syrian hamster embryo cells (Hesterburg & Barrett, 1984) but not C3H10T\textsuperscript{1/2} cells (Brown et al., 1983). A number of other studies have demonstrated an increase in chromosomal aberrations including breaks, fragmentation and aneuploidy (Sinnock & Seabright, 1975; Huang et al., 1978).

One hypothesis which has been proposed for the carcinogenicity of asbestos is that these substances induce the production of oxygen free radicals which may damage DNA and augment carcinogenesis (Mossman & Landesman, 1983). Asbestos has been shown to stimulate the production of oxygen free radicals from polymorphonuclear leucocytes (Doll et al., 1982). Recently, Weitzmann and Graceffa (1984) have demonstrated that suspensions of asbestos catalyse the formation of hydroxyl radicals from hydrogen peroxide.

Asbestos has appreciable amounts of iron within its lattice structure which can act as a catalyst for the generation of active oxygen radicals by an iron mediated Haber–Weiss reaction (Eberhardt et al., 1983) and induce DNA strand scission (Kasai & Nishimura, 1984). An important end result of oxygen free radical damage is the peroxidation of polyunsaturated fatty acids which can lead to the formation of malondialdehyde. We have previously reported increases in lipid peroxidation with cells and found the iron content was responsible for the oxygen free radical-like damage (Turver et al., 1985).

To investigate the role of free radicals in the action of asbestos, we have examined the production of malondialdehyde in cell cultures following treatment with asbestos and studied the effect of iron chelating agents on its production. A parallel study was also conducted to find whether asbestos had any effect on cell DNA strand breakage. Again scavengers of iron were added to test the hypothesis of an association between iron content and DNA-strand breakage.

Materials and methods

Cell cultures

The C3H10T\textsuperscript{1/2} cell line (Reznikoff et al., 1973) was received from Dr W.J. Harris, Inveresk Research International, Musselburgh, Scotland, UK. Cells were cultured in Dulbecco’s modification of Eagle’s medium containing 10% heat inactivated foetal bovine serum, streptomycin (50 \textmu g\textsuperscript{mL}^{-1}), penicillin (50 \textmu U\textsuperscript{mL}^{-1}) and 3.6 g\textsuperscript{L}^{-1} sodium bicarbonate. Cultures were incubated at 37°C in an atmosphere of 8% carbon dioxide in air. Tissue culture medium, serum reagents, and sterile plastic were obtained from Flow Laboratories, Irvine, Scotland and Gibco Europe, Paisley, Scotland, UK.

Chemicals

\textit{6}\textsuperscript{3H}-Thymidine specific activity 26 Ci\textsuperscript{mmol}^{-1} was supplied by Amersham International PLC, UK. Desferrioxamine mesylate was supplied by Ciba Laboratories, Horsham, UK. Other organic chemicals were from Sigma Chemical Company, Poole, Dorset, UK.

Dusts

The UICC samples of asbestos (Timbrell & Rendall, 1971) were used, and a sample of the UICC crocidolite was ball milled for 8 h (Brown et al., 1978) to provide a low cytotoxic dust. All dust samples were weighed and autoclaved dry and resuspended in culture medium by sonication immediately before addition to cell cultures.

Measurement of lipid peroxidation

Lipid peroxidation was measured using the thiobarbituric acid (TBA) test for malondialdehyde using methods previously described (Gavino et al., 1981). This method
detects products of lipid free radical damage, particularly malondialdehyde, but also includes other lipid oxidation products and the term malondialdehyde was used synonymously with thiobarbituric acid reactive substances (TBARS). The experiments used cell cultures maintained in 56 cm² Petri dishes which were grown to confluence before treatment with dust for 24 h. The role of iron was studied using desferrioxamine at a dose that was previously determined not to be toxic to the cells. A standard curve was constructed using known amounts of MDA generated by the acid hydrolysis of 1,1,3,3-tetramethoxypropane.

DNA-strand breakage

The method was based on that described by Collins et al. (1982) especially suitable for measuring dust induced DNA damage. C3H10T1/2 cells were inoculated into 8 chamber multislides (Lab-Tek division of Miles Laboratories) each chamber receiving 1 × 10⁴ cells in 0.3 ml of medium. These cultures were then incubated for 2 days with 6-[³H]dT at 0.2 μCi ml⁻¹ and grown to confluence. After this time they were washed and reincubated in non-radioactive medium. Two hours later this medium was replaced by medium containing 10 mM hydroxyurea and 10 μM 1-p-D-arabinobiosylfuransosylcytosine (AraC, Sigma Chemical Co.).

After 1 h suspensions of dust or a solution of 4-nitroquinoline-1-oxide as positive control was added, these solutions being made in medium containing the above inhibitors. After 24 h the chambers were removed from the slides and the monolayers washed in medium. In some experiments desferrioxamine was added with dust at a non-toxic dose and DMEM was replaced by MEM to reduce the effect of ferric iron present in the DMEM. The cells were then lysed in alkali by gently pipetting 50 μl of alkali sucrose (5% w/v sucrose, 0.3 M NaOH, 0.5 M NaCl) onto each slide. After 15 min at 4°C the alkali cell lysate was brought to pH 4.5 with 15 μl of 2M acetic acid. A 10 mm diameter nitrocellulose disc (BA85 Schleicher & Schuell) was placed over each slide followed by a similarly sized GF glass fibre disc (Whatman) to absorb the lysis solution. The plastic chamber unit previously removed from the slide was inverted and placed over the disc with a 25 g weight to exert even pressure during the transfer of DNA.

After 1 h at room temperature the chamber, weight and glassfibre disc were removed. Fifty μl of a solution containing 0.24 Uml⁻¹ deoxyribonuclease-S (Calbiochem) in sodium acetate buffer (0.03 M, pH 4.5 with ZnSO₄ at 30 μM) was added to each nitrocellulose disc. The slides were then incubated at 45°C for 45 min, a second glass fibre disc was then placed on each filter to absorb the supernatant. The nitrocellulose discs were treated with TCA, washed and dried. All the discs were then placed in scintillation vials with 2 ml of scintillant and counted. The percentage of radioactivity released from the nitrocellulose filter by the nuclease treatment is a measure of single-stranded DNA resulting from unwinding at the sites of damage.

Results

Effect of crocidolite asbestos on lipid

The results in Figure 1 show that crocidolite asbestos is effective in stimulating the accumulation of TBARS in cultures of C3H10T1/2 cells. The effect of milling crocidolite is also shown and had no effect on the production of TBARS. Asbestos contains iron in its lattice structure which could participate in the production of TBARS. To investigate this hypothesis cultures of C3H10T1/2 cells were treated with asbestos which had been pretreated with desferrioxamine, a specific iron chelator. The results in Figure 1 show that desferrioxamine significantly reduced TBARS formation in both asbestos treated and control cultures. In the same experiment milled crocidolite was also effective in stimulating the accumulation of TBARS.

Effect of crocidolite asbestos on DNA

The alkaline strand separation method was used to detect the accumulation of DNA breaks produced by asbestos. The effect of fibre size and exposure time on DNA strand breakage is presented in Figure 2. When C3H10T1/2 cells were incubated with either milled crocidolite or the parent UICC crocidolite for only 2.5 h there was no significant increase in DNA-strand breakages as measured by the quantity of radioactive label released from the nitrocellulose-bound DNA following SI-nuclease treatment. In contrast, when these same concentrations were extended to 24 h all concentrations produced significant breakage compared with the control. Also, the ability of milled crocidolite to induce strand breakage indicates the phenomenon is largely independent of fibre morphology. The carcinogen 4-nitroquinoline-1-oxide was used as a positive control and induced DNA-strand breakages in a dose-dependent manner confirming the assay was responsive to chemically-induced DNA-damage.

The effect of desferrioxamine on strand breakage was investigated. Table 1 shows that pretreatment with desferrioxamine reduced asbestos DNA-strand breakage. The reduction was only small and the inhibition could have been affected by the presence of iron in the DMEM reducing the effectiveness of the desferrioxamine. To remove this possibility a second experiment was conducted in which the cells were treated in MEM which contains no added iron.

Table 1 The effect of desferrioxamine on crocidolite induced DNA-damage in C3H10T1/2 cells as measured by the S1-nuclease digest method

| Treatment | % Release of nitrocellulose-bound DNA by S1-Nuclease |
|-----------|-----------------------------------------------|
| Control   | 34.04 ± 9.73(6) 16.07 ± 4.21(6)*               |
| Crocidolite | 63.20 ± 7.79(5)* 51.61 ± 10.95(4)*             |
| Milled Crocidolite | 70.45 ± 8.38(5)* 52.37 ± 1.75(5) |
The results from this experiment are presented in Figure 3 and demonstrate that desferrioxamine produces significant inhibition of asbestos strand breakages.

**Discussion**

The results above indicate crocidolite asbestos can induce lipid peroxidation and cause DNA damage in C3H10T1/2 cells by a mechanism involving iron. In previous studies asbestos has been shown to induce lipid peroxidation with red blood cells (Gabor & Anca, 1975), phospholipid emulsions (Weitzman & Weitberg, 1985) and isolated microsomes (Gulumian et al., 1983). Others have shown the iron component of crocidolite fibres can catalyse the generation of hydroxyl radicals from hydrogen peroxide (Weitzman & Graceffa, 1984). We have previously shown that the lipid damaging activities of crocidolite is preventable by desferrioxamine treatment (Turver et al., 1985). This has led to the suggestion that asbestos can cause cell toxicity by a modified Haber-Weiss reaction.

The generation of TBARS by crocidolite asbestos was found to be independent of fibre morphology. Milled (non-fibrous) crocidolite asbestos was as least as effective as the UICC sample, indicating a surface-related phenomenon. This effect is further supported again by our earlier observations and those of Weitzman and Graceffa (1984). The iron chelator desferrioxamine significantly reduced asbestos stimulated TBARS accumulation suggesting that the oxidative damage responsible for lipid peroxidation was produced by an iron-catalysed reaction. Amphibole asbestos, such as crocidolite, possesses significant amounts of ferrous and ferrie iron linked into its crystal structure and has previously been shown to generate free radicals from hydrogen peroxide (in non-cellular systems). This type of mechanism has also been shown to damage isolated DNA when incubated with asbestos and hydrogen peroxide (Kasai & Nishimura, 1984).

A number of the toxic and pathogenic activities of asbestos fibres have been shown to be dependent on the size and shape of the dust (Brown et al., 1978; Stanton et al., 1977). Milled crocidolite is significantly less cytotoxic than parent dust towards C3H10T1/2 cells so the increased sensitivity of DNA from treated cells is not caused by an overall toxic phenomenon. The formation of strand breaks was also dependent on the length of incubation, there was little effect at 2.5h while significant damage was seen after 24h. The delay may represent the time for the dust to interact with the cells. In contrast 4-nitroquinoline-N-oxide produced detectable damage after both 2.5h and 24h incubation. Thus there are similarities between asbestos-induced lipid peroxidation and DNA-strand breaks which appear to be a product of surface chemistry rather than fibre morphology. Further insight into the DNA-damaging activity of asbestos was found from the effect of desferrioxamine on DNA-strand breakage. The addition of desferrioxamine to asbestos exposed cells caused a significant reduction in DNA-strand breakage. This was particularly evident when these experiments were carried out in medium containing little or no added iron.

The presence of increased DNA-strand breaks in asbestos-exposed cells would seem to support the idea that asbestos can cause direct genetic damage. The presence of this activity could also explain some earlier reports showing genotoxicity with asbestos (Huang et al., 1978; Livingston et al., 1980). However, there have been reports which have failed to detect DNA-strand breaks using alkaline elution in human cells (Formace et al., 1982). There may be a number of reasons for these variations: these authors used a different assay method; a shorter exposure time and with lower concentrations of dust. In this study milled crocidolite was found to be as active as the UICC crocidolite sample. In contrast, milled crocidolite has been found to be less tumourigenic in animals by intrapleural injection (Wagner et al., 1984). It is possible that a combination of fibre size and
breakage and lipid peroxidation is consistent with the concept that this damage is by a free-radical mechanism. However, these effects were not fibre-size dependent but may still represent an important mechanism in the aetiology of asbestos associated disease.

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