

Coclastogenic Effects of Sodium Arsenite on Chinese Hamster Ovary Cells (CHO9) and Primary Human Fibroblasts (VH25)

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ABSTRACT The coclastogenic effects of sodium arsenite (SA) at its non-cytotoxic concentrations of 1.5 μ M and 3 μ M were studied in combination with three different types of known clastogenic agents namely ultra violet radiation (UVB, 280-320nm), methyl methanesulphonate (MMS) and mitomycin C (MMC) at varying doses/concentrations. The end points studied were the frequencies of induced sister chromatid exchanges (SCEs) and micronuclei (MN) in Chinese hamster ovary cells (CHO9) and primary human fibroblast cells (VH25). In general post treatment with SA produced synergistic increases in clastogenicity with all the three clastogens. Very few exceptions were found where increases in clastogenicity was not synergistic. In CHO9 cells post treatment with SA resulted in increased synergistic effects in the frequencies of SCEs at the lower dose/concentrations of UVB, MMS and MMC. In VH25 cells post treatment with SA produced synergistic effects in MMS and MMC induced damage in all the treatments. A similar synergistically increased frequency in MN's at the higher doses in UVB and at the lower concentrations of MMC. A non synergistic increase in MMS after post treatment with SA were observed in CHO cells. In VH25 cells, MMS in all combinations produced a synergistic effect after post treatment with SA. In general, these results demonstrate the coclastogenicity of SA with UVB, MMS and MMC treatments in CHO9 and VH25 cells.

INTRODUCTION

Arsenic is a known human carcinogen and an environmental pollutant (IARC monographs 1980, 1987) known to cause adverse health effects such as liver injury, neurotoxicity, increase the risk of cancers of skin, lung, bladder and liver (IARC monographs 1980; Bates et al. 1992; Chen et al. 1992; Chiou et al. 1995). Sodium arsenite (SA) can act as comutagen due to its ability to inhibit the activities of thiol containing enzymes (Sunderman 1979), such as DNA ligase (Li and Rossman, 1989b) resulting in defective DNA replication, repair, recombination and joining of single- and double-stranded DNA breaks (Lasko et al. 1990).

Earlier studies had reported that inorganic arsenicals can inhibit (a) DNA repair such as the removal of UV-induced thymidine dimers in human cells (Okui and Fujiwara 1986), (b) DNA ligase II activity in Chinese hamster V79 cells (Li

and Rossman 1989b) and (c) repair of N-methyl-N-nitrosourea (MNU) - induced lesions measured by incorporation of dNMPs into damaged DNA template or by interfering with the ligation step (Li and Rossman 1989a). Jha et al. (1992) reported an interference in DNA replication-dependent processes by SA, thus leading to chromatid type of aberrations and SCEs as well as potentiating X-rays and UV induced chromosomal damage in human peripheral lymphocytes and primary human fibroblasts.

In vitro studies (Jacobson-Kram and Montalbano 1985; Kochar et al. 1996) showed that arsenic was clastogenic and induced sister chromatid exchanges in mammalian cells. Several other studies (Jha et al. 1992; Burgdorf et al. 1977; Wen et al. 1981) also showed an enhanced sister chromatid exchanges in human lymphocytes due to arsenic exposure.

Post treatment with SA synergistically increases cytotoxicity, mutagenicity and clastogenicity of DNA crosslinking agents, X-rays, UV-light and alkylating agents in rodent and human cells (Okui and Fujiwara 1986; Jha et al. 1992; Lee et al. 1985; Lee et al. 1986a; Lee et al. 1986b; Lee-Chen et al. 1993, 1994), similar to the enhancing effects of many DNA repair

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inhibitors on the DNA damage induced by known mutagens (Downes et al. 1983).

Thus, though numerous studies have indicated comutagenic, clastogenic and cytotoxic properties of SA, substantial evidence to show its coclastogenic effects at non-cytotoxic concentrations is lacking. It is well established, that UVB radiation (280-320nm), monofunctional alkylating agent methyl methanesulphonate (MMS) and polyfunctional alkylating agent mitomycin C (MMC) are clastogenic and it was proposed to investigate whether SA can potentiate the clastogenicity of these agents. Special emphasis was given to the incubation conditions and cytotoxicity, as it is important to define biological effects which occur at low, non-cytotoxic concentrations of arsenite. The endpoints studied were the frequency of sister chromatid exchanges (SCEs) and frequency of chromosomal aberrations assessed as induced micronuclei (MN). MN assay was selected because its frequencies appeared to be correlated with the production of acentric fragments (Ramalho et al. 1988; Littlefield et al. 1989).

MATERIALS AND METHODS

Cell Culture: A) Chinese hamster ovary (CHO9) cells, with a cell cycle time of about 12h, were grown in Ham's F-10 medium supplemented with 15% fetal calf serum and antibiotics in a humidified incubator at 37°C and in an atmosphere of 5% CO₂ in air.

B) Primary human fibroblasts (VH25), with a cell cycle time of about 24h, were grown in Ham's F-10 medium supplemented with 15% fetal calf serum and antibiotics, in humidified incubator at 37°C and 2.5% CO₂ in air.

Chemicals Used: The chemicals used were MMS (Eastman Kodak, Rochester, NY), MMC (Kyowa, Japan) and SA (CAS Number 7784-46-5; Sigma) and were dissolved in phosphate-buffered saline (PBS) just before treatment.

Experimental Design: Treatment and experimental protocols to analyse the frequency of sister chromatid exchanges (SCEs) and micronuclei (MN) were as reported earlier from this laboratory (Natarajan et al. 1983; Dulout and Natarajan 1987).

A) Chinese hamster ovary (CHO9) cells

Treatment with SA: Cells were grown for one cell cycle (~12h) in a complete medium with 5-bromodeoxyuridine (BrdU, 5µM; Sigma). They were further allowed to grow with SA at two different concentrations (1.5µM, 3µM) for one

cell cycle time and then were fixed. These concentrations were found to be non-cytotoxic in cell survival studies.

UV Radiation

The UVB (280-320nm) fluence rate was determined using IL700A spectro-radiometer, International light Inc. Newburyport, MA, with a cosine-corrected SEE 400 detector and a WBS 320 filter. Cells were exposed to UVB at a fluence of 4.5j/m²/sec through a Mylar sheet (to eliminate wavelengths shorter than 290nm). The three doses selected were 100j, 250j and 500j.

a) UVB Treatment: The cells were washed with phosphate buffered saline (PBS) and then irradiated in PBS (10ml). After irradiation, the cells were allowed to grow in complete medium with BrdU for 24h and then were fixed.

b) UVB + SA Treatment: Cells were washed with PBS, then irradiated in PBS (10ml) and allowed to grow in complete medium with BrdU and SA at two different concentrations (1.5mM, 3mM) for 24h and then were fixed.

Treatment with Alkylating Agents

a) Methyl methanesulfonate (MMS): Cells were grown for 12h in the presence of BrdU, washed with PBS and treated for 1h with MMS (0.12, 0.25 and 0.5mM). They were washed and allowed to grow in complete medium with BrdU for further 12h and then fixed.

b) MMS + SA: Cells were grown for 12h in the presence of BrdU, washed with PBS and treated for 1h with MMS (0.12, 0.25, 0.5 and 1.0mM). They were washed with PBS and then allowed to grow in a complete medium with BrdU and SA at two different concentrations (1.5 mM, 3 mM) for 12h and then fixed.

c) Mitomycin C (MMC) and d) MMC + SA: The same protocol as above was followed. The concentrations of SA i.e. 0.03, 0.06, 0.12 and 0.24mM were used.

B) Primary human fibroblasts (VH25)

Treatment with SA: Confluent cells were subcultured and grown for 24h in a complete medium with BrdU. They were further allowed to grow in the complete medium with BrdU, with the addition of SA (at two different concentrations 1.5µM, 3µM) for 24h and then fixed.

Treatment with Alkylating Agents: The experimental protocol was the same as that for CHO9 cells, except for the increased cell cycle time of 24h instead of 12h.

Cell Fixation and Slide Preparation: The cells were fixed in methanol: acetic acid (3: 1) and the slides were prepared by standard procedures and stained by Fluorescence plus giemsa (FPG) technique (Perry and Wolff, 1974) to analyse sister chromatid exchanges (SCEs). Fifty cells from every treatment were scored and each experiment was repeated twice.

CB-MN Assay (Cytochalasin Blocked Micronucleus Assay): The experimental protocol described by Dulout and Natarajan (1987) was adopted to analyse the induced micronuclei *in situ*. Briefly, the cells were grown as monolayer in 24 × 24 mm cover glasses set in 9-cm Falcon petri dishes divided into three wells. Thus, each treatment could be replicated three times in the same petri dish. All other cell culture conditions were the same as above. After the treatment with the mutagens, the cells were washed with PBS and grown in a complete medium containing Cytochalasin B (Sigma; 3µg/ml) in the presence of SA for a further period of one cell cycle time. The frequency of micronuclei was estimated in the cytochalasin blocked binucleate cells. The cover glasses were fixed in 50% methanol with two changes, air dried and stained with Acridine Orange (AO) solution (30µg/ml) for 3-5 min. With AO the cytoplasm gives bright orange fluorescence and the nucleus as well as the micronuclei gives bright yellow/green when observed under a fluorescent Zeiss Axioskop microscope (Carl Zeiss) with FITC/TRITC filter. A total of 1000 binucleated cells per dose were analysed.

The concentrations used for the treatment, as well as fixation times, were determined on the basis of results obtained from our pilot experiments.

All experiments were repeated two times and the mean values with standard deviation are given in the results. The data were statistically analysed by using student's (one-tailed) t-test to estimate the level of significance. Both for SCE and MN analysis, the background value (control) was subtracted from the total.

RESULTS

CHINESE HAMSTER OVARY CELLS

i. UVB and SA Treatment

(a) Sister Chromatid Exchanges: A dose dependent increase in the frequency of SCEs

after treatment with UVB alone and in combination with SA was observed as shown in Table 1. The frequency of SCEs (14.5 per cell) induced by the combined treatment of 100j UVB with 3µM SA was close to the additive frequency of SCEs (14.2) induced by individual treatments of UVB (7.7) and SA (6.5). At the highest dose of UVB (500j) a synergistic effect (37.0 per cell), 21.3% more than the additive value (30.5) of UVB and SA was observed. (The term synergistic effect is used to mean that it is higher than the additive value of the effect induced independently by two different agents used in an experiment). Even though the treatment of UVB at 250j with 3mM SA produced an increase in the frequency of SCEs (25.0 per cell) over the value of any one single treatment yet it was less than the additive frequency (27.6) induced by the individual treatments. The synergistic effect of SA was found only with the highest dose of UVB employed.

Table 1: Frequency of SCEs/cell following UVB/SA treatment in CHO9 cells

Treatment	Observed value	S.D. (±)	Expected additive value	Increase over expected value
control	14.7	3.1		
SA3µM	6.5	3.9		
100j	7.7	4.9		
100j+SA	14.5	7.8	14.2	+0.3 (+2.1%)
250j	21.1	8.3		
250j+SA	25.0	7.8	27.6	-2.6 (-9.4%)
500j	24.0	4.7		
500j+SA	37.0	2.0	30.5	+6.5 (+21.3%)

(b) Micronuclei (MN): There was a dose related increase in the frequencies of MN after treatment with either UVB or SA. Frequency of MN in UVB treated cells increased by post treatment with SA (Table 2). The increased frequencies produced by the combined treatment at the lower doses, 100j and 250j UVB with 1.5mM (82.3 and 108.6) and 3mM (95.8 and 148.6) of SA were less than the additive values (130.8 and 139.6 with 1.5mM SA, and 173 and 181.8 with 3mM SA) of the individual treatments. However in the combined treatment at the highest dose of 500j UVB with 1.5mM (195.2 per cell) and 3mM (253.7 per cell) of SA synergistic effect of 8.3 and 14.0%, were observed. Thus, only at 500j a synergistic effect of SA was observed.

ii. MMS and SA treatment

(a) Sister Chromatid Exchanges: A dose

Table 2: Frequency of micronuclei following UVB/SA treatment in CHO9 cells (per 1000 Binucleate cells)

Treat-ment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	34.4	11.1		
SA1.5uM	81.2	18.8		
100j	49.6	3.7		
100j+SA	82.3	14.0	130.8	- 48.5 (-37.1%)
250j	58.4	9.0		
250j+SA	108.6	15.6	139.6	- 31 (-22.2%)
500j	99.1	5.6		
500j+SA	195.2	3.1	180.3	+ 14.9 (+8.3%)
SA3uM	123.4	20.1		
100j+SA	95.8	18.0	173.0	- 77.2 (-44.6%)
250j+SA	148.6	14.9	181.8	- 33.2 (-18.3%)
500j+SA	253.7	2.0	222.5	+ 31.2 (+14.0%)

related increase in the frequency of SCEs was observed when cells were treated either with MMS or with SA individually as shown in Table 3. There was a moderate synergistic increase ranging from 19.8 to 26.2% in the frequency of SCEs in all the combined treatment with MMS and SA except in one combination with highest doses of both the chemicals, and this may probably be due to toxic effect resulting from that combination.

Table 3: Frequency of SCEs/cell following MMS/SA treatment in CHO9 cells

Treat-ment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	16.4	3.9		
SA1.5uM	0.6	3.1		
MMS.25	25.6	7.2		
MMS+SA	31.5	6.6	26.2	+ 5.3 (+20.2%)
MMS.5	65.0	9.2		
MMS+SA	82.8	11.8	65.6	+ 17.2 (+26.2%)
SA3uM	5.2	6.0		
MMS.25+SA	36.9	12.0	30.8	+ 6.1 (+19.8%)
MMS.5+SA	65.6	11.1	70.2	- 4.6 (-6.5%)

MMS concentration is in mM

(b) Micronuclei (MN): A dose related increase in the frequency of MN was observed after treatment with MMS or SA when used singly or in combination as presented in Table 4. The increase in the MN frequencies in all the combinations showed an effect that was lower than the additive value of two single treatments.

iii. MMC and SA

(a) Sister Chromatid Exchanges: A dose dependent increase in the frequency of SCEs

Table 4: Frequency of micronuclei following MMS/SA treatment in CHO9 cells (per 1000 Binucleate cells)

Treat-ment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	33.5	5.5		
SA1.5uM	78.5	9.4		
MMS.12	97.3	44.7		
MMS+SA	132.9	7.8	175.8	- 42.9 (-24.4%)
MMS.25	110.6	13.7		
MMS+SA	156.5	15.3	189.1	- 32.6 (-17.2%)
MMS.5	247.5	11.3		
MMS+SA	272.5	17.6	326.0	- 53.5 (-16.4%)
SA3uM	134.4	10.1		
MMS.12+SA	112.3	7.3	231.7	- 119.4 (-51.5%)
MMS.25+SA	227.2	54.7	245.0	- 17.8 (-7.3%)
MMS.5+SA	248.0	2.4	381.9	- 133.9 (-35.1%)

MMS concentration is in mM

after treatment with MMC or SA singly and in combination was observed, as shown in Table 5.

A synergistic effect at the 0.12mM and 0.24mM MMC and a less than additive value at 0.03mM MMC+SA (1.5mM and 3mM) and in 0.24mM MMC+SA 3mM were observed.

Table 5: Frequency of SCEs/cell following MMC/SA treatment in CHO9 cells

Treat-ment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	16.4	3.9		
SA1.5uM	0.6	3.1		
MMC.03	3.8	4.8		
MMC+SA	4.2	4.5	4.4	- 0.2 (-4.5%)
MMC.12	4.3	4.6		
MMC+SA	9.2	4.5	4.9	+ 4.3 (+87.7%)
MMC.24	14.1	6.9		
MMC+SA	16.7	7.5	14.7	+ 2.0 (+13.6%)
SA3uM	5.2	6.0		
MMC.03+SA	6.1	7.0	9.0	- 2.9 (-32.2%)
MMC.12+SA	11.0	7.3	9.5	+ 1.5 (+15.8%)
MMC.24+SA	15.4	7.4	19.3	- 3.9 (-20.2%)

MMC concentration is in mM

(b) Micronuclei (MN): A dose dependent increase in the frequencies of MN was obtained after treatment with MMC or SA used singly or in combination as shown in Table 6. A synergistic increase ranging from 13.0 to 105.7% in the frequencies of MN was observed in all the combinations with all doses of MMC and SA over the expected additive values. However the highest doses of MMC + SA (0.24mM + 3mM) resulted in a lower frequency of induction of MN which may be due to toxic effect of the combination treatment (expected value/observed value, 224.3/114.1).

Table 6: Frequency of micronuclei following MMC/SA treatment in CHO9 cells (per 1000 Binucleate cells)

Treatment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	33.5	5.5		
SA1.5 μ M	78.5	9.4		
MMC.03	34.5	44.7		
MMC+SA	232.5	7.8	113.0+	119.5 (+105.7%)
MMC.12	67.3	13.7		
MMC+SA	241.7	15.3	145.8	+ 95.9 (+65.8%)
MMC.24	89.9	11.3		
MMC+SA	235.6	17.6	168.4	+ 67.2 (+39.9%)
SA3 μ M	134.4	10.1		
MMC.03+SA	216.3	7.3	168.9	+ 47.4 (+28.1%)
MMC.12+SA	228.0	54.7	201.7	+ 26.3 (+13.0%)
MMC.24+SA	114.1	2.4	224.3	- 110.2 (-49.1%)

MMC concentration is in mM

PRIMARY HUMAN FIBROBLASTS

i. MMS and SA treatment

(a) *Sister Chromatid Exchanges*: A dose-related increase in the frequency of SCE was observed when cells were treated either with MMS or with SA individually and in combination. As shown in Table 7, a synergistic increase ranging from 5.9 to 88.5% over the expected additive values in the frequency of SCE in all the combined treatments was observed.

Table 7: Frequency of SCEs/cell following MMS/SA treatment in VH25 cells

Treatment	Observed value	S.D. (\pm)	Expected additive value	Increase over expected value
control	10.5	3.4		
SA1.5 μ M	5.3	5.0		
MMS.25	6.9	4.6		
MMS+SA	16.5	7.1	12.2	+ 4.3 (+35.2%)
MMS.5	29.4	10.9		
MMS+SA	58.1	27.1	34.7	+ 23.4 (67.4%)
SA 3 μ M	6.1	4.6		
MMS.25+SA	24.5	4.4	13.0	+ 11.5 (+88.5%)
MMS.5+SA	37.6	10.1	35.5	+ 2.1 (+5.9%)

MMS concentration is in mM

(b) *Micronuclei (MN)*: A dose dependent increase in the frequency of MN was obtained after treatment with MMS, only at the low concentrations (0.25 and 0.5mM). At the highest dose of MMS (1.0mM), the increase in the frequency of MN was not dose related most probably due to toxic effect of the combined treatment. As presented in Table 8, the frequency

of MN in MMS treated cells was increased by post treatment with SA. However, the frequencies of MN (38.1 and 36.2) in the combined treatments (0.25mM MMS with 1.5mM SA and 3mM SA respectively) were less than the respective additive values (71.0 and 78.2). At the combination of higher concentrations (0.5 and 1.0mM) with SA (1.5 and 3 μ M) a synergistic increase ranging from 2.9 to 48.8% over the additive value was noticed.

Table 8: Frequency of micronuclei following MMS/SA treatment in VH25 cells (per 1000 Binucleate cells)

Treatment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	20.0	3.7		
SA1.5 μ M	39.5	9.1		
MMS.25	31.5	9.8		
MMS+SA	38.1	7.3	71.0	- 32.9 (-46.3%)
MMS.5	165.5	49.5		
MMS+SA	305.0	6.0	205.0	+ 100.0 (+48.8%)
MMS1.0	85.5	5.4		
MMS+SA	143.2	2.7	125.0	+ 18.2 (+14.6%)
SA 3 μ M	46.7	8.3		
MMS.25+SA	36.2	9.1	78.2	- 42.0 (-53.7%)
MMS.5+SA	283.3	6.2	212.2	+ 71.1 (+33.5%)
MMS1.0+SA	136.0	3.2	132.2	+ 3.8 (+2.9%)

MMS concentration is in mM

ii. MMC and SA

(a) *Sister Chromatid Exchanges*: A dose dependent increase in the frequency of SCE was obtained in treatment with MMC and SA. A moderate synergistic effect more than the additive value of single treatments with MMC and SA was observed in all the combined treatments of MMC and SA (Table 9) except in the combinations of 0.06mM MMC with 1.5 and 3mM SA.

Table 9: Frequency of SCEs/cell following MMC/SA treatment in VH25 cells

Treatment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	10.5	3.4		
SA1.5 μ M	5.3	5.0		
MMC.03	30.1	14.7		
MMC+SA	37.5	9.6	35.4	+ 2.1 (+5.9%)
MMC.06	32.2	11.6		
MMC+SA	37.4	10.8	37.5	- 0.1 (-0.3%)
SA 3 μ M	6.1	4.6		
MMC.03+SA	39.5	3.4	36.2	+ 3.3 (+9.1%)
MMC.06+SA	36.6	4.5	38.3	- 1.7 (-4.4%)

MMC concentration is in mM

(b) **Micronuclei (MN):** A dose dependent increase in frequency of MN was also observed in single treatments either with MMC, SA or in combination. A synergistic effect ranging from 74.6 to 176.0% over the additive frequencies was observed in all the combined treatments except in the treatment at the highest dose level of 0.06mM MMC + 3 μ M SA (Table10).

Table 10 : Frequency of micronuclei following MMC/SA treatment in VH25 cells (per 1000 Binucleate cells)

Treatment	Observed value	S.D. (\pm)	Expected additive value	Difference over expected value
control	20.0	3.9		
SA1.5 μ M	39.5	9.1		
MMC.03	7.7	2.1		
MMC+SA	100.4	6.5	47.2	+53.2 (+112.7%)
MMC.06	24.7	6.0		
MMC+SA	177.2	4.2	64.2	+113.0 (+176.0%)
MMC.12	61.1	2.7		
MMC+SA	175.7	4.0	100.6	+75.1 (+74.6%)
SA3 μ M	46.7	8.3		
MMC.03+SA	146.4	8.5	54.4	+92.0 (+169.1%)
MMC.06+SA	26.3	5.8	71.4	-45.1 (-63.2%)

MMC concentration is in mM

DISCUSSION

In CHO9 cells, SA and all the three clastogenic agents studied i.e. UVB, MMS and MMC with different modes of action were found to be effective in producing SCEs and MN. It is known that mutagens such as UVB and alkylating agents (MMS and MMC) require S-dependent repair of induced DNA lesions for the formation of chromosomal aberrations as well as SCEs (Perry and Evans 1975; Kihlman 1977; Natarajan and Obe 1982).

In VH25 cell line also, an effective induction of SCEs and MN with MMS, MMC and SA was observed. However the frequency of SCEs produced by MMS in VH25 cells was lower than that was observed in CHO cells. Similar frequencies of SCEs in both the cell lines were produced by MMC and SA.

Coclastogenic Effect of SA with UVB in CHO9 Cells: Arsenic has been shown to enhance the cytotoxicity, mutagenicity and clastogenicity in combination with UV radiation, X-rays, alkylating agents as well as DNA cross-linking compounds in cultured mammalian cells (Hartwig, 1995; Hartmann and Speit, 1996). The cocytotoxic effect of arsenite in excision of pyrimidine dimers has been reported (Okui and Fujiwara 1986) in normal human fibroblasts.

There was a synergistic increase in the frequency of SCEs produced by UVB after treatment with SA at all doses except 250j of UVB. The UVB clastogenicity and the enhancing effect of SA on it may be due to the interference of SA in excision repair.

Earlier studies (Jha et al. 1992; Lee et al. 1985) showed that post treatment with sodium arsenite synergistically increased the UVC induced chromosomal aberrations. Huang et al. (1992) studied the enhancing effect of arsenite on chromosomal aberrations induced by UVC at various points in the cell cycle and reported a coclastogenic effect of SA by treating cells at late G1/S transition with UVC and arsenic. They also suggested that the inhibition of dimer excision might be important than the inhibition of ligation for the expression of coclastogenic effect, as chromatid exchanges were believed to result from misjoining of two double-strand breaks, as observed in the combined treatment of UVC and SA treated in late G1 of CHO cells. This is supported by the results of our study where a synergistic increase in the frequency of SCEs was observed, when UVB treated cells were post treated with SA.

The dose dependent increase in the MN frequency of UVB + SA treated cells probably resulted from the inhibition of DNA ligation by SA as has been earlier reported (Jha et al. 1992; Lee et al. 1985, 1986a, b) using UVC + SA. In CHO9 cells synergistic effect of SA with UVB and MMC in the induced frequency of MN was noticed at higher doses/concentrations of UVB and MMC and no synergistic effect with MMS was noticed.

Coclastogenic Effect of SA with MMS and MMC in CHO9 and VH25 Cells: As in the case of UVB, SA potentiated the effects of MMS and MMC in the induction of SCEs. Post treatment with SA and MMS and MMC treated cells produced synergistic effect at lower concentrations and at the highest concentration the effect was reduced, which may be due to the cytotoxic effect of the combination of MMS or MMC with SA. This is also supported by the report (Hartwig, 1995) that in AS III and other metal ions DNA repair processes were disturbed at low non-cytotoxic concentrations.

However, in VH25 cells a synergistic effect in the production of MN at all doses except in one with MMC and two with MMS after SA was observed. This shows a differential sensitivity

of two cell lines namely CHO9 and VH25, in the production of chromosomal aberrations (MN) when treated with MMS. This difference in response may be related to the known different kinetics of nucleotide or base excision repair capacity of these two cell lines.

Although frequencies of SCEs in both cell lines in MMS treated cells were synergistically increased after post treatment with SA the increase in the frequency of MN was lower than the additive value of two single treatments. This observation shows that there may be different pathways leading to the formation of SCEs and chromosomal aberrations (MN) depending on the type of lesions responsible for these endpoints.

SA post treatment synergistically increased the frequency of SCEs and MN induced by MMC in CHO cells. These results are similar to those reported in the combined treatment of cross-linking agents, Cis-diammine dichloroplatinum (II) (Cis-ptII) and 8-methoxypsoralen (8-MOP) with SA (Lee et al. 1986b) in CHO cells.

As alkylating agents are S-dependent clastogen and induce chromatid type of aberrations and when SA is also present during S phase of the cell cycle, an increase in clastogenicity can be observed. Earlier studies (Jha et al. 1992) had also shown that SA could act as a clastogen when present during S-phase of the cell cycle by interfering with DNA replication process.

Present results suggest that non-cytotoxic concentrations of SA enhances the clastogenicity of UVB, MMS and MMC in CHO9 cells and primary human fibroblasts (VH25). Its coclastogenic effect is S-dependent and not restricted to certain types of DNA lesions alone. This study adds further evidence that SA inhibits DNA repair process and interferes with DNA replication-dependent processes in UVB, MMS and MMC treated cells leading to induction of sister chromatid exchange and micronuclei.

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