Epidemiological studies have suggested that a high fat diet and the metabolic activity of gut flora are major factors contributing to development of colon cancer (Reddy, 1981). Studies in rats have showed that bile acids such as deoxycholic acid and lithocholic acid enhance tumour formation in the colon (Reddy, 1981), stomach (Koborn et al., 1984) and liver (Cameron et al., 1981). Moreover, bile acids have been found to have initating (Kelsey et al., 1979) or enhancing (Kaibara et al., 1984) effects in transformation of cultured cells. We have been attempting to clarify the mechanism of these biological effects by studying DNA lesions in cultured cells exposed to bile acids. Previously, we found that various bile acids induced DNA single strand breaks, the extent of which depended on their degree of incorporation into cultured cells, and that most of the single strand breaks (up to 80%) were associated with protein-DNA cross linking. We also found that the induction of single strand breaks was effectively suppressed by topo-isomerase inhibitors such as novobiocin (Nov), nalidixic acid (Nal), oxolinic acid (Oxl) and coumermycin A (Kaneko et al., 1986). To examine whether these inhibitors also inhibited cell transformation, we investigated their effects on promotion of transformation by lithocholic acid (LC) in Balb/3T3 cells. Here we show that Nal and Oxl efficiently reduced the promotion of transformation by LC initiated by 3-methylcholanthrene (MC) in BALB/3T3 fibroblasts.

The BALB/3T3, A31-1-1 in vitro transformation system was used. This system has been well characterized (Kakunga, 1973) and used as a model system in two-step carcinogenesis experiments (Kennedy & Little, 1978). The cells were grown in Eagle's minimum essential medium supplemented with 10% heat-inactivated foetal calf serum. For transformation, inocula of 1 x 10⁴ cells of the A31-1-1 BALB/3T3 cell line (passage 3 after receipt as a gift from Dr T. Kuroki) were seeded into ten 60 mm cell culture dishes per experimental group. After 24 h, the cultures were treated with MC (Spectrum Chem, MFG. Corp., Redondo Beach, Ca.) for 72 h at 37°C. Then the medium was changed, and culture was continued for 4 weeks in the presence or absence of LC (Sigma Chem. Co., St Louis, Mo.) with or without Oxl or Nal (Sigma Chem. Co.) as indicated. The medium was renewed twice a week. Type III foci (Reznikoff et al., 1973) in each plate were scored as transformants with the aid of a dissecting microscope. The criteria adopted for them are that cells are highly condensed, heavily piled upon one another, and randomly oriented. In addition, areas of patchy, dense growth without cross-linking were often observed but they were clearly distinct from the above foci and were not scored. MC and LC were dissolved in dimethylsulfoxide (Pierce, Rockfield, II). Nal and Nov were dissolved in distilled water and Oxl was dissolved in 40 mM NaOH. The promoter and inhibitors were present throughout the expression period after the treatment with MC. Four dishes per group seeded with 200 cells were treated in parallel to assess the effects of the various treatments upon colony forming efficiency. Colonies in test dishes were scored after incubation for one week.

Five experiments were carried out to determine the effects of Nal and Oxl on MC-induced transformation and LC-induced promotion of transformation, together with confirmation of LC-induced promotion (6-mercaptopurine (6-MP) inhibition, 1984) and three of these experiments (A, B and C) are shown in Table I. As shown in expts. A, B and C, Oxl and Nal clearly suppressed transformation promoted by LC. Surprisingly, the suppression was frequently to below the level of transformation induced by MC only. In most experiments, these inhibitors also suppressed transformation induced by MC only. On the other hand, novobiocin, which also suppressed induction of DNA strand breaks by LC, little affected LC-induced promotion of transformation when added at 75 μM, which was the highest concentration possible because of its high toxicity (Table II). LC alone (40 μM) induced transformation on prolonged treatment (expts. A, B and C), but not on a single treatment (data not shown). Therefore, it may have weak initiating activity in BALB/3T3 cells, as it does in hamster embryo cells (Kelsey et al., 1979). The transformation induced by prolonged treatment with LC was also suppressed by Nal (expt. C) or Oxl (expt. B). Thus, the prokaryotic topo-isomerase inhibitors Nal and Oxl, but not Nov, suppressed transformation induced by MC or LC, and also promotion by LC after initiation by MC in BALB/3T3 cells as well as suppressing LC-induced DNA strand breaks in cultured human fibroblasts (Kaneko et al., 1986).

Neither the way in which LC acts on inducing DNA strand breaks and transformation nor the way in which Oxl and Nal act on suppressing them are known. The following observations suggest that topo-isomerase participate in these processes: First, Oxl, Nal and Nov have been reported to inhibit 4'- (9-acridinylamino)-methanesulfon-m-anisidide (m-AMSA) induced DNA-protein cross linking mediated by topo-isomerase II (Pommier et al., 1984, Nelson et al., 1984). Moreover, we found previously that they also inhibited DNA-protein cross linking induced by LC in cultured fibroblasts (Kaneko et al., 1986). Second, in a preliminary experiment we found that in a cell free system topo-isomerase assayed by measuring relaxation of closed circular DNA (unpublished data). Nal may inhibit LC-induced processes by reducing the level of ATP, since it was found to interfere with oxidative phosphorylation in isolated rat liver mitochondria and to inhibit ATP synthesis (Gallagher et al., 1986). However, its effect may not be mediated by decrease in the ATP level, because DNA strand breaks in cultured fibroblasts were increased 2- to 3-fold in the presence of an uncoupler such as oligomycin or dinitrophenol (unpublished results). In addition, it was shown that Nal and Oxl inhibit topo-isomerase II in a cell free system from HeLa cells at high concentration (Miller et al., 1981) but that Nal does not inhibit topo-isomerase from D. melanogaster (Hae & Brutlag, 1980). Based on these

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for
for
None
113.5+ 8.9
LC
107.8+ 6.8
LC+Oxl
110.3+ 8.1
MC(1 μg ml⁻¹)
None
62.8+ 8.5
LC
65.8+ 3.5
Oxl
70.0+ 5.4
LC+Oxl
69.0+ 1.6
B.
None
None
73.5+ 8.8
LC
96.5+ 9.2
Oxl
110.3+ 9.5
LC+Oxl
98.2+ 6.8
MC(1 μg ml⁻¹)
None
76.8+ 3.0
LC
76.0+ 13.1
Oxl
76.5+ 7.3
LC+Oxl
68.7+ 7.4
C.
None
None
77.0+ 5.9
LC
69.8+ 3.3
Nal
81.0+ 5.0
LC+Nal
68.3+ 9.3
MC(1 μg ml⁻¹)
None
28.8+ 5.0
LC
24.5+ 3.0
LC+Nal
51.0+ 10.0
MC(5 μg ml⁻¹)
None
30.0+ 8.1
LC
27.0+ 1.9
Nal
19.8+ 3.7
LC+Nal
45.8+ 7.0

Table I Effects of oxolinic acid and nalidixic acid on lithocholic acid-induced promotion of transformation initiated by 3-methylcholanthrene

| Treatment | Plating efficiencya | No. of type III foci/dishb | Fractions of dishes without foci | Frequency of transformation |
|-----------|---------------------|---------------------------|---------------------------------|---------------------------|
| Expt.     | Initiation          | Promotion                 | /200                            | /10 × 10⁻⁴                |
| A.        | None                | None                      | 104.0± 8.8                      | 0.0                       | 0.0                       |
|           | LC                  | 96.5± 9.2                 | 0.4± 1.0                        | 8                         | 0.86± 2.10                |
|           | Oxl                 | 110.3± 9.5                | 0.0                             | 10                        | 0.0                       |
|           | LC+Oxl              | 98.2± 6.8                 | 0.4± 0.5                        | 6                         | 0.84± 1.10                |
|           | MC(1 μg ml⁻¹)       | None                      | 76.8± 3.0                       | 0.8± 0.8                  | 4                         | 2.16± 2.14                |
|           | LC                  | 76.0± 13.1                | 0.3± 0.5                        | 7                         | 0.82± 1.30                |
|           | Oxl                 | 76.5± 7.3                 | 0.3± 0.5                        | 3                         | 2.36± 2.02a               |
|           | LC+Oxl              | 68.7± 7.4                 | 0.6± 0.7                        | 3                         | 4.9± 4.1a                 |

*Concentrations used were as follows: LC, 40 μM; Oxl, 100 μM; Nal, 0.5 mM; Values are means ± s.d. for 4 dishes (plating efficiency) or 10 dishes (transformation); Differences from MC(1 μg ml⁻¹) (P<0.001 for exp. A, P<0.01 for exp. C) by t-test; Differences from MC(1 μg ml⁻¹) plus LC (P<0.001 for both exp. A and C and P<0.01 for exp. B); Differences from MC(1 μg ml⁻¹) (P<0.01); Differences from MC(5 μg ml⁻¹) (P<0.001); Differences from MC(5 μg ml⁻¹) (P<0.02); Differences from MC(5 μg ml⁻¹) plus LC (P<0.001).

Table II Effects of novobiocin on lithocholic acid-induced promotion of transformation initiated by 3-methylcholanthrene

| Treatment | Plating efficiencyb | No. of type III foci/dishb | Fractions of dishes without foci | Frequency of transformation |
|-----------|---------------------|---------------------------|---------------------------------|---------------------------|
| Initiation | Promotiona          | /200                      | /10 × 10⁻⁴                      |
| None      | None                | 104.0± 8.8                | 0.0                             | 0.0                       |
|           | LC                  | 96.5± 9.2                 | 0.4± 1.0                        | 8                         | 0.86± 2.10                |
|           | Nov                 | 97.8± 2.5                 | 0.0                             | 10                        | 0.0                       |
|           | LC+Nov              | 100.8± 10.0               | 4.8± 7.5                        | 0                         | 9.5± 14.9                |
|           | MC(1 μg ml⁻¹)       | None                      | 73.5± 6.9                       | 1.0± 1.3                  | 5                         | 2.7± 3.5                 |
|           | LC                  | 74.5± 4.3                 | 20.1± 3.9                       | 0                         | 54.0± 10.5                |
|           | Nov                 | 71.3± 10.2                | 4.4± 3.3                        | 1                         | 12.3± 28.6                |
|           | LC+Nov              | 62.3± 4.5                 | 17.2± 6.7                       | 0                         | 55.2± 21.5                |

*Concentration used were: LC, 40 μM; Nov, 75 μM; As foot-note to Table I.

findings we propose that LC, Oxl and Nal may act in the following way on the induction of DNA strand breaks and transformation and on their suppression: First, LC might interfere with some function of topoisomerase and this interference might induce DNA-protein cross linking. Second, LC-induced DNA-protein cross linking might play a role in LC-induced promotion of transformation. Third, the interaction between LC and topoisomerases might be inhibited directly or indirectly by Oxl or Nal, or cells treated with Oxl or Nal might be able to repair LC-induced damage. Experiments using a cell-free system are necessary to investigate these possibilities and will help in determining whether topoisomerases actually participate in transformation induced by MC alone.

Recently, there has been increasing interest in the function of topoisomerases. There are suggestions (Wang, 1985) that topoisomerase I is essential for RNA transcription and that topoisomerase II is essential to DNA replication. Also, it is reported that topoisomerase II is a major constituent of nuclear scaffold (Earnshaw et al., 1985) and is essential to mitosis (Holm et al., 1985). Modification of topoisomerase II by an antineoplastic agent such as m-AMSA was reported to induce preferential cleavage at a DNase I hypersensitive region (~270 bp upstream of Ori) of SV40 chromatin in
monkey kidney cells (Yang et al., 1985) and sister chromatid exchange in Chinese hamster cells (Dillehay et al., 1987). On the other hand, recently, the effect of 4β-phorbol 12,13-dibutyrate on differentiation of HL-60 cells was found to be suppressed by inhibitors of topoisomerase II such as Nov and m-AMSA and activation of topoisomerase II due to phosphorylation by protein kinase C was suggested to be involved in the differentiation process (Sahyoun et al., 1986).

Thus, topoisomerases are considered to be important not only in differentiation but also in carcinogenesis after activation or inactivation. But there have been no previous suggestions that topoisomerases participate in promotion of carcinogenesis or in the whole process of carcinogenesis. Therefore, it is important to examine whether they are essential for induction of transformation by MC and induction of promotion by LC.

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