EFFECTS OF LOCAL HYPERTERMIA ON THE PHARMACOKINETICS OF MISONIDAZOLE IN THE ANAESTHETIZED MOUSE

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Summary.—The effects of sodium pentobarbitone anaesthesia, the presence of a tumour, and local hyperthermia to a tumour-bearing leg, on the pharmacokinetics of MISO in the mouse are reported. Analysis of MISO and its metabolite Ro 05-963 was by high-performance liquid chromatography. The plasma kinetics of MISO were largely unaffected by any of these treatments, but hyperthermia substantially reduced tumour concentrations of the drug.

The effects of tumour site and size on unheated-tumour drug concentrations were also studied, and an increase in tumour size was shown to decrease tumour MISO levels, but to different degrees according to whether implanted in the leg or flank.

Uniformity of MISO distribution throughout heated and unheated tumours was examined, and levels were found to be constant within tumours. The presence of a temperature detector in heated tumours did not affect their drug concentration.

The hypoxic cell radiosensitizer misonidazole (1-(2-nitroimidazol-1-yl)-3-methoxy propan-2-ol; Ro 07-0582 from Roche Laboratories; NSC-261037; MISO), which is currently undergoing clinical trials, has been shown to exhibit selective cytotoxicity to hypoxic cells (Mohindra & Rauth, 1976; Brown, 1975; Sutherland et al., 1976). It has been further demonstrated that this cytotoxicity is enhanced by hyperthermia both in vitro (Bleehen et al., 1976; Stratford & Adams, 1977) and in vivo in the mouse (Bleehen et al., 1977; George et al., 1977). This in vivo work in the mouse has involved anaesthetizing the animal, usually with sodium pentobarbitone. Recently there has been a growing awareness that the side effects of anaesthesia can grossly affect the end-points of some experiments, for instance Saffan (the veterinary equivalent of Althesin) has been reported to increase the growth delay of a solid tumour treated with Melphalan (Peacock & Stephens, 1978). Sodium pentobarbitone itself has been reported to alter the radiosensitivity of various tumours and normal tissues, sometimes rendering them radioresistant (Sheldon et al., 1977; Keizer & Van Putten, 1976) and sometimes more radiosensitive (Horney et al., 1977; Hendry, 1978). It has also been reported to have no influence on radiosensitivity (Douglas & Fowler, 1976; Paterson & Matthews, 1951). It is therefore necessary to check the effect of an anaesthetic in any system where it has been used, and the first part of the present paper describes our investigations into the effect of sodium pentobarbitone on the pharmacokinetics of MISO.

The effect of variation in temperature on drug pharmacokinetics had not been widely studied when the subject was reviewed by Ballard in 1974, with the conclusion that both hyperthermia and hypothermia may in some instances influence the pharmacokinetic behaviour of a drug. For example, the rate of binding of $^{32}$P Thiotepa to perfused hind limbs of dogs was shown to be higher at tissue tempera-
tures of 37.2–38.9°C than at 23.9–26.1°C (Rochlin et al., 1961). More recently a
detailed study of Adriamycin in whole-
body hyperthermic (42.5°C) and control
rabbits (Mimnaugh et al., 1978) revealed
no difference in clearance rate of the
parent drug, but significantly higher con-
centrations of Adriamycin and its meta-
bolites in the skeletal muscle and duodenum
of hyperthermic animals. In this paper we
report investigations into the effect of
local hyperthermia on the pharmaco-
kinetics of MISO by studying plasma and
tumour concentrations of MISO and its
O-demethylated metabolite, Ro 05-9963.
We also report on the influence of tumour
size and site on MISO and Ro 05-9963
concentrations, on the uniformity of dis-
tribution of MISO within tumours and on
the effect of mechanical disturbance by
the temperature probe detectors on MISO
concentrations in tumours.

EXPERIMENTAL

Animals

Adult BALB/c mice were obtained from
the breeding colony at NIMR (Mill Hill,
London) and were housed in plastic cages on
sawdust bedding. They were fed PRD nuts
(Labsure Animal Diets, Poole, Dorset) and
allowed water ad lib. Mice weighed 20–25 g.

Tumours

The tumour used throughout this work is
the subline of the EMT6 tumour now desig-
nated EMT6/Ca/VJAC, and it has been
described in detail elsewhere (Twentyman &
Bleehen, 1975). The tumour is maintained in
alternating in vivo/in vitro passages, regu-
larly returning to frozen stock. Tumours were
implanted using cells harvested from the 2nd
to 5th in vitro passage after an in vivo passage.
An inoculum of 10⁸ cells in 0.1 ml growth
medium was used. Leg tumours were pro-
duced by i.m. inoculation in the lower leg, and
flank tumours by inoculation intradermally
(i.d.) in the previously plucked skin of the
flank.

Tumour volumes

Leg tumours.—Volumes were calculated
after taking caliper measurements of 2
mutually perpendicular diameters at right
angles to the axis of the leg, taking the mean
of these and calculating the volume as for a
sphere, then subtracting the volume of
normal leg. This method was found to be
acceptable for tumours greater than 6-0 mm
in mean diameter. (Any tumour in which the
difference between diameters exceeded 1-0
mm was discarded.) Leg tumours usually
reached a mean diameter of about 8-0 mm by
Days 7 or 8, and for “small” leg tumours the
mean diameters were restricted to 7.75–8.5
mm, that is, 90–170 mm³. “Large” leg
tumours were usually grown by Days 10–11
and the mean diameters were restricted to
10–11.25 mm, equivalent to a volume of
135–550 mm³. For “very large” leg tumours
the mean diameters were restricted to 12–0–
13.75 mm, i.e. 700–1100 mm³.

Flank tumours.—Volumes were calculated
by measuring 3 mutually perpendicular axes
of the usually ovoid tumour, and calculating
the volume according to the method of
Watson (1976). Tumours usually arrived at a
volume of about 100 mm³ by Day 9. “Small”
flank tumours were used in the volume range
60–140 mm³ and “large” ones in the range
350–550 mm³.

Drugs

Supplies of MISO and its O-demethylated
metabolite Ro 05-9963 (1-(2-nitrimidazole-
1-yl)-2,3 propandiol) were provided by Roche
Laboratories (Welwyn Garden City, Herts).
MISO was routinely dissolved in Hanks’
balanced salt solution at 25 mg/ml for the
high dose (1.0 g/kg) and 2.5 mg/ml for the
low dose (100 mg/kg). The solutions were
injected i.p. such that a 25g mouse received
1.0 ml. MISO solutions were prepared daily
and protected from light.

Sodium pentobarbitone (sodium 5-ethyl-
5(1-methylbutyl)-barbiturate; Sagatal) was
obtained from May and Baker Ltd (Dagen-
ham, Essex) as a 60mg/ml solution and diluted
1 in 10 in Hanks’ before injection. 60 mg/kg
was the standard initial dose for animals not
also receiving MISO and 40 mg/kg for those
treated with MISO. This reduced dose is
adequate, because MISO increases the anaes-
thetic effect of Sagatal. Animals being heated
usually require at least one repeat dose of
about 15 mg/kg to maintain anaesthesia for
the full heating time of 60 min. When used,
Sagatal was always administered 10 min before the injection of MISO.

**Heating**

Leg tumours were heated by immersion in a temperature-controlled circulating water-bath (Grant Instruments, Barrington) with the water bath temperature controlled to ±0.1°C. The water bath temperature was monitored with a mercury thermometer calibrated against a thermometer standardized by the National Physical Laboratory. Mice were supported in a specially designed jig with only the tumour-bearing leg immersed. Heating was started 10 min after injection of Sagatal and immediately after injection of MISO. In all experiments the duration of heating was 60 min at a water bath temperature of 44°C. This produced an intra-tumour temperature of 43.0±0.6°C for small leg tumours. Temperature monitoring of the tumours, performed in one set of experiments only, was by means of a 27-gauge needle thermistor probe in association with a direct reading electric thermometer (Light Laboratories, Brighton). The needle was inserted into the flank above water level and passed down to the tumour s.c. in order to minimize errors in temperature reading due to conduction along the needle. The same apparatus fitted with a special blunt temperature probe was used to measure rectal temperatures.

**Measurement of plasma and tumour concentrations of MISO and Ro 05-9963**

At appropriate times after MISO injection, mice were bled by cardiac puncture, using diethyl ether anaesthesia where necessary. Tumours were dissected out directly after cardiac puncture. Blood and tumour samples were put on ice immediately after collection. Plasma was obtained by centrifugation (2000 g for 10 min) of heparinized whole blood, and stored at −20°C, as were tumour samples. Plasma and tissue homogenate (10% to 20% w/v in distilled water) were analysed by reversed-phase high-performance liquid chromatography (HPLC) as previously described (Workman et al., 1978). This technique allows the specific assay of MIS and the O-demethylated metabolite, Ro 05-9963.

**Estimation of pharmacokinetic parameters**

For up to 6 or 8 h after a dose of 1·0 g/kg of MISO, the elimination of MISO from the plasma closely approximates to first-order kinetics. The apparent elimination rate constant (K_e) is given by the slope of the plot of log MISO concentration against time, and was estimated by the method of least squares linear regression. The apparent half-life (t_1/2) is given by \(1n2/K_e\). The area under the curve (AUC) of a drug has been widely used as a measure of tissue exposure. The AUCs of plasma or tumour MISO and Ro 05-9963 concentrations were estimated by Simpson's rule. When the AUC for both compounds is converted from \(\mu g.h/ml\) to \(\mu mol.h/ml\) the AUCs can be summed to give the total nitroimidazole AUC.

**RESULTS**

**Effect of Sagatal on MISO pharmacokinetics**

The dose of Sagatal used to anaesthetize mice also treated with 1·0 g/kg MISO (e.g. in heating experiments) is 40 mg/kg. The effect of this dose on MISO pharmacokinetics was investigated in non-tumour-bearing mice. Fig. 1 shows the
TABLE I.—Effects of Sagatal on the pharmacokinetics of MISO in normal and pheno-barbitone-pretreated male mice after a 1·0 g/kg dose of MISO

| Pretreatment | Sagatal anaesthesia | Apparent MISO $t_1$ (h) | AUCo-8h† $\mu$mol h/ml | Total 2-nitroimidazole |
|--------------|---------------------|------------------------|-------------------------|------------------------|
| A None       | None                | 2·36                   | 15·80                   | 17·55                  |
|              | 40 mg/kg            | 2·12                   | 17·04                   | 18·96                  |
| B Phenobarbitone | None                | 1·09                   | 8·17                    | 10·34                  |
| 5 days at 100 mg/kg | 40 mg/kg            | 1·08                   | 7·90                    | 9·99                   |

95% confidence limits in parentheses (5 animals per time point).
† No significance tests possible for AUC.

time courses of plasma MISO and Ro 05-9963 after 1·0 g/kg MISO in control and Sagatal-treated male mice, and Table I shows the $t_1$ values and AUCs (Part A). There is no significant difference between the $t_1$ values ($P > 0·1$). Similar results were obtained in repeat experiments and also in female animals.

Previous studies have shown that the $t_1$ can be shortened by phenobarbitone pre-treatment with 100 mg/kg daily for 5 days before MISO treatment (Workman, 1979). Part B in Table I shows that Sagatal has no effect in the induced mice. We also demonstrated that Sagatal had no effect on tumour concentrations of MISO and Ro 05-9963. For example, MISO concentrations in small flank tumours 40 min after injection were 359 ± 23 (2 × s.e.) µg/g in control mice and 385 ± 18 µg/g in mice treated with Sagatal (n = 5 in both cases; $P > 0·1$).

Effect of the presence of a tumour on MISO pharmacokinetics

Previous studies have shown that solid tumours may inhibit the metabolism of drugs (Sladek et al., 1978). In view of this, a series of experiments was carried out on male mice with and without small leg tumours, using low-dose MISO (100 mg/kg); the low dose was used since any small difference due to the presence of a tumour would probably be more apparent at the low dose. Tumours were not heated, but mice were anaesthetized with Sagatal. MISO plasma concentrations were measured at 30min intervals for 3 h after injection, and data from one experiment are shown in Fig. 2, illustrating that there is no observable effect due to the presence of the tumour. The apparent $t_1$ values are
Table II.—Apparent MISO t½ after a 100 mg/kg dose of MISO to anaesthetized male mice, with and without "small" leg tumours.

| Expt | t½ (h) Without tumour | t½ (h) With tumour |
|------|-----------------------|-------------------|
| A    | 0.73                  | 0.67              |
|      | (0.51-1.28)           | (0.55-0.88)       |
| B    | 0.61                  | 0.67              |
|      | (0.56-0.69)           | (0.62-0.73)       |
| C    | 0.60                  | 0.69              |
|      | (0.48-0.79)           | (0.58-0.86)       |

95% confidence limits in parentheses. 3 animals per group.

given in Table II, experiment B being the one illustrated in Fig. 2. There is no significant difference between t½ values, \( P > 0.1 \) in all cases. The t½ values quoted here for the low dose (100 mg/kg) are shorter than those shown in Table I for 1-0 g/kg. This is due to the dose-dependence of MISO pharmacokinetics (Workman, in preparation). Comparison between high-dose experiments also showed that there were no differences due to the presence of a tumour.

Effect of local heat on plasma and tumour levels of MISO and Ro 05-9963

The effect of heat on plasma and tumour levels of MISO and its metabolite was investigated in male animals with small leg tumours (see experimental section) which were heated for 1 h at a waterbath temperature of 44°C. As part of this study we also investigated the effects of MISO with and without local heating on the core temperature of anaesthetized mice.

High dose studies

1-0 g/kg MISO was injected i.p. into anaesthetized animals at the start of heating. Unheated animals were also anaesthetized. In all three experiments plasma MISO levels were similar in locally heated and control animals for up to 6 h. Later plasma levels were rather higher in the heated animals, so that plasma MISO t½ values tended to be somewhat higher in locally heated animals. However, this difference was only significant \( P < 0.02 \) in one experiment for which the data are illustrated in Fig. 3 and pertinent pharmacokinetic parameters are summarized in Table III. In the other experiments the t½ values for locally heated and control animals respectively were 2.44 and 1.97 h in one, and 3.09 and 2.44 h in the other. In neither experiment was the difference significant \( P > 0.1 \). In all 3 experiments the plasma MISO AUCs were similar for locally heated and control animals, and Ro 05-9963 levels were not affected.

All experiments showed a lower tumour MISO and Ro 05-9963 level in heated than unheated tumours, the difference being greater at the earlier times (up to 3 h) (Fig. 3). This produced a lower AUC for heated tumours, an effect which is described in greater detail below.

Low-dose studies

In view of the difference in MISO pharmacokinetics at high and low doses, the effect of local hyperthermia was
Table III.—Effect of 1 h local heat to the tumour on plasma and tumour concentrations of MISO and Ro 05-9933 after a 1.0 g/kg dose of MISO

|                  | Plasma | Tumour |
|------------------|--------|--------|
|                  | AUC₀₋₉₉ h | Total 2-NITROIMIDAZOLE | AUC₀₋₉₉ h | Total 2-NITROIMIDAZOLE |
| Control          | 1.83 (1.67-2.02) | 18.95 | 1.82 | 20.77 | 7.65 | 0.14 | 7.79 |
| Heat             | 2.70 (2.49-3.08) | 21.27 | 1.71 | 22.98 | 5.73 | 0.11 | 5.84 |
| Heat/Control AUC₀₋₉₉ | 1.12 | 1.12 | 0.75 | 0.75 |

95% confidence limits in parentheses. 3 animals per time point.

Examined at the low dose of MISO. The experiment was repeated exactly as above but using a 100 mg/kg dose of MISO and measuring plasma and tumour levels of MISO and Ro 05-9963 at 0.5, 1, 2, 3 and 4 h only after MISO administration, as levels fall below the detectable limit at later times. The time courses of both measured drugs are shown in Fig. 4 and are seen to be very similar in heated and control animals. The apparent MISO t₅₀ for heated and control animals are 1.02 h (0.84-1.28 h) and 0.95 h (0.77-1.24 h) respectively, and there is no significant difference between these values (P > 0.1). It was found that tumour levels of MISO and Ro 05-9963 were undetectable in all cases.

Effect of local heating on core temperature

Measurements of rectal temperature, as a measure of core temperature, have been made on unanaesthetized and MISO-treated anaesthetized heated and unheated mice at an ambient temperature of 26.5°C. Unanaesthetized, untreated mice have a rectal temperature of about 38–38.5°C, whereas the temperature of anaesthetized (40 mg/kg Sagatal) mice treated with 1.0 g/kg MISO drops by an average of 5°C during the first hour and then steadily rises during the next 9 h so that it has returned to normal by 10 h after drug administration. Although a dose of 60 mg/kg Sagatal (which produces a similar level of anaesthesia to 40 mg/kg Sagatal in conjunction with 1.0 g/kg MISO—see “Drugs” section) also causes a temperature drop of about 5°C, this effect is relatively short-lived, the minimum temperature being recorded 40 min after the drug is given, and by 2 h after the

![Time vs Plasma Nitroimidazole Concentration](image-url)

**Fig. 4.**—Effect of local hyperthermia of the tumour-bearing leg on low-dose MISO levels in plasma. ○—○ Unheated plasma MISO. ●—● Heated plasma MISO. ▼ Unheated plasma Ro 0-9963. ▼ Heated plasma Ro 05-9963. Lines are fitted by least squares regression.
temperature is within 0.5°C of that of untreated controls. The effect of 1.0 g/kg MISO on unanaesthetized animals is almost the same as on anaesthetized ones, the temperature dropping by an average of 4°C in the first hour. The time scale of recovery is similar to that for anaesthetized mice: normal by 10 h after treatment following a steady rise from 2 h after treatment. The temperatures of locally heated, anaesthetized mice also treated with 1.0 g/kg MISO show more variation between animals, and they rise to a maximum after about 30 min of heating (from 39.5 to 40.5°C) when further anaesthetic is required and the temperature drops to between 38.5 and 39.5°C. It then rises gradually or remains approximately steady for the remaining heating time.

Thus the core temperatures of heated and unheated mice after treatment with 1.0 g/kg MISO are very different during the hour of heating, differences of up to 7°C being quite likely. After heating, the temperatures of heated animals drop, so that the discrepancy between core temperatures of heated and unheated mice diminishes.

**Effect of heat, size and site on high dose MISO levels in tumours**

In these experiments MISO levels were measured in tumour and plasma at 2 selected times only, 1 and 3 h after injection of 1.0 g/kg of MISO. The effect of heat for 1 h after the injection of MISO in large and small tumours was examined, and also the effect of tumour size in both leg and flank, using 3 sizes of leg tumour and 2 of flank tumour. Small leg and flank tumours were of similar volume, as were large tumours in both sites (see “Experimental” section for sizes used). No flank tumours were heated, but all mice were anaesthetized with Sagatal. All experiments demonstrated the same trend and the results of one are summarized in Table IV. The MISO tumour: plasma ratios are seen to be consistently lower in heated than in control tumours (P < 0.01 for 1 h data and P < 0.001 for 3h data). Also, in both flanks and legs, large tumours have a lower ratio than small ones (P < 0.001 for both times) but no further decrease is seen in very large leg tumours. The tumour: plasma ratios in unheated small leg and flank tumours were similar, as might be expected, but the same increase in size decreased the tumour: plasma ratio more in leg tumours than in flank tumours (P < 0.001 for 1 h data and P < 0.01 for 3h data). Similarly, the effect of heat on large leg tumours appeared to be proportionately greater than on small leg tumours, but this was significant at 3 h only (P > 0.2 for 1h data and P < 0.01 for 3h data).

The probabilities quoted above were calculated by carrying out an analysis of variance on log tumour:plasma ratios, considering firstly the effects of heat and size and their interaction, and then the effects of site and size and their interaction. Results of this analysis are given in Table V to illustrate the relative significance of the effects. The effect of size is the most significant, having the greatest F ratio.

**Uniformity of MISO distribution in heated and unheated tumours**

It has been shown that cell survival is higher in the centre than the periphery of heated tumours with 1.0 g/kg MISO (Bleehen et al., 1977). Although it seemed likely that a higher temperature at the periphery accounted for the lower survival there, it was possible that a non-uniform distribution of MISO throughout the tumour might have contributed to this phenomenon. Experiments were therefore carried out using male mice with leg tumours given 1.0 g/kg MISO and either heated for 1 h in a 44°C water bath or left at room temperature. All mice were anaesthetized with Sagatal. Tumours were excised 1 h after injection with MISO, i.e. immediately after heating when appropriate, and carefully dissected into peripheral and central areas which were assayed separately. Data from one experiment are shown in Table VI. They
TABLE IV.—Effect of heat, size and site on tumour/plasma ratios of MISO for a 1·0 g/kg dose of MISO

| Time (h) | Control | Heat |
|----------|---------|------|
|          | Tumour (µg/g) | Plasma (µg/ml) | Tumour (µg/g) | Plasma (µg/ml) |
| Small leg tumours (90–170 mm³) |
| 1       | 326·5 ± 44·3 | 621·6 ± 31·6 | 52·5 | 252·1 ± 42·8 | 609·8 ± 22·1 |
| (n=10)  |         |         | (n=10) |         |         |
| 3       | 139·1 ± 28·7 | 281·7 ± 28·3 | 49·4 | 138·7 ± 39·9 | 338·8 ± 35·3 |
| (n=10)  |         |         | (n=10) |         |         |
| Large leg tumours (350–550 mm³) |
| 1       | 98·9 ± 43·3 | 739·4 ± 65·2 | 13·4 | 47·6 ± 15·0 | 815·0 ± 74·5 |
| (n=10)  |         |         | (n=10) |         |         |
| 3       | 84·3 ± 21·3 | 456·6 ± 58·6 | 18·5 | 28·8 ± 13·8 | 437·5 ± 43·4 |
| (n=10)  |         |         | (n=10) |         |         |
| Very large leg tumours (700–1100 mm³) |
| 1       | 118·4 ± 37·8 | 924·6 ± 77·8 | 12·8 | nd          |            |
| (n=9)   |         |         | (n=9)  |         |         |
| 3       | 125·5 ± 33·2 | 508·8 ± 45·7 | 24·7 | nd          |            |
| (n=8)   |         |         | (n=8)  |         |         |
| Small flank tumours (60–140 mm³) |
| 1       | 387·0 ± 57·8 | 665·7 ± 73·3 | 58·1 | nd          |            |
| (n=10)  |         |         | (n=10) |         |         |
| 3       | 219·9 ± 40·3 | 432·1 ± 50·5 | 50·9 | nd          |            |
| (n=10)  |         |         | (n=10) |         |         |
| Large flank tumours (350–550 mm³) |
| 1       | 285·1 ± 69·1 | 874·1 ± 58·2 | 30·3 | nd          |            |
| (n=8)   |         |         | (n=8)  |         |         |
| 3       | 179·9 ± 59·5 | 511·5 ± 110·5 | 35·2 | nd          |            |
| (n=8)   |         |         | (n=8)  |         |         |

Figures are mean ± 2 s.e. of n determinations.
nd = not determined.

TABLE V.—Results of analysis of variance on log tumour/plasma ratios for 1h and 3h data from Table IV

| Effect | 1h data | 3h data |
|--------|---------|---------|
|        | F ratio | P       | F ratio | P       |
| Heat   | 8·2     | <0·01   | 22·9    | <0·001  |
| Size   | 138·9   | <0·001  | 93·0    | <0·001  |
| Heat × size | 1·51   | >0·2   | 10·2   | <0·01   |
| Site   | 18·4    | <0·001  | 13·0    | <0·001  |
| Size   | 81·0    | <0·001  | 64·6    | <0·001  |
| Site × size | 13·5  | <0·001 | 11·7    | <0·01   |

show that there is no significant difference (P > 0·1) in MISO concentrations between the centres and peripheries of tumours having had the same treatment, i.e. either heated or controls. However, the difference reported above between heated and unheated tumours is again seen. There is a significant difference in tumour concentrations, for both peripheral and central portions, between heated and unheated tumours (P < 0·01).

Effect of the presence of a temperature probe on tumour levels of MISO

Having previously reported that the presence of a temperature probe can affect
the survival of tumours treated with MISO and heat (Honess et al., 1978) we investigated the possibility that the bleeding caused by the probe could cause higher MISO levels in monitored than unmonitored tumours. Experiments were carried out on male mice bearing small leg tumours heated for 1 h after administration of 1·0 g/kg MISO. They had been anaesthetized with Sagatal 10 min before receiving MISO. Tumours were excised immediately after heating, and the results indicated no significant difference between monitored and unmonitored tumours (P > 0·1); hence the extra cell killing in the presence of the probe cannot be attributed to higher MISO levels.

**DISCUSSION**

In this paper we have shown that sodium pentobarbitone anaesthesia does not alter the plasma kinetics of MISO in normal mice. We have also shown that the presence of a tumour does not affect the plasma MISO levels. Neither the anaesthetic nor the tumour had a significant effect on the half-life or AUC. Also the oxidative demethylation to the Ro 05-9963 metabolite was unchanged. We therefore proceeded to investigate the effect of local hyperthermia on plasma and tumour kinetics of MISO, being as certain as possible that no differences that might be observed could be attributed to an anaesthetic artefact, anaesthesia being unavoidable in our current method of tumour heating.

Local hyperthermia was found to have very little effect on the plasma pharmacokinetics of MISO. There was, however, a tendency for the plasma MISO t₄ in locally heated animals to be rather longer at the high dose (1·0 g/kg) of MISO, but this was not significant in 2/3 experiments, nor at the low dose (100 mg/kg). The increase in t₄ was due to higher plasma levels at the later times (8–12 h), when no more than 20% of the peak drug concentration remained. As a result, the MISO AUC was not affected. Previous reports have suggested that the toxicity of MISO is related to the AUC (Dische et al., 1977; Workman, 1979). Our finding that hyperthermia does not affect the plasma AUC of MISO is interesting in view of the increased toxicity of the drug by this treatment (Overgaard, 1979; unpublished results). This suggests that other factors may also be involved in the toxicity. Our finding that locally applied hyperthermia, which also caused a systemic temperature rise of between 2 and 3°C, has very little effect on the pharmacokinetics of MISO is essentially similar to that of Mimnaugh (1978) on Adriamycin in rabbits. In that study the core temperature of hyperthermic animals was 2·5°C higher than that of controls.

In contrast to the finding that local heating had little effect on systemic MISO pharmacokinetics, this treatment was found to reduce the MISO tumour levels, differences of up to 70% being found in the early samples. Other studies on tissue uptake of drug at raised temperatures deal with normal tissues, e.g. dog leg (Rochlin et al., 1961) and rabbit skeletal muscle and duodenum (Mimnaugh, 1978) and they both demonstrate a higher level in the heated normal tissue. It seems unlikely that the reduction in MISO concentration could be due to in-

**Table VI.**—Comparison of MISO and Ro 05-9963 levels (μg/g) in central and peripheral areas of leg tumours, heated and unheated, for a 1·0 g/kg dose of MISO

|          | Control (n = 10) | Heat (n = 9) |
|----------|-----------------|-------------|
|          | MISO | Ro 05-9963 | MISO | Ro 05-9963 |
| Periphery | mean  | 729 | 24·6 | 354 | 16·9 |
|          | 2 x s.e.| 203 | 5·8 | 91 | 2·7 |
| Centre   | Mean | 694 | 16·3 | 317 | 14·7 |
|          | 2 x s.e. | 160 | 4·7 | 120 | 4·1 |
CREASED oxidative metabolism in the tumour (which is on average 2°C hotter than the maximum core temperature) because correspondingly higher Ro 05-9963 levels might be expected, yet these were found to be unchanged. However, we cannot rule out the possibility that more rapid metabolism by other routes may be responsible. It is also possible that the decreased MISO concentrations in heated tumours are lower due to a reduction in the blood supply. This might be caused by heat-induced oedema constricting the blood vessels in the leg; oedema is brought about within 30 min of immersion in a 44°C water bath. Alternatively, it may be caused by a reduction in blood flow to the heated tumour, associated with a drop in blood pressure caused by sodium pentobarbitone. We did show that in the absence of heat sodium pentobarbitone did not affect tumour MISO concentration, but were unable to compare heated tumours with and without the anaesthetic. Johnson et al. (1976) have measured blood pressure and tumour blood flow at different temperatures in CBA and WHT mice bearing leg tumours under sodium pentobarbitone anaesthesia. Observations were made on blood flow at 20 and 39°C. Whereas at 20°C the anaesthetic caused a slight decrease in blood flow followed by recovery, at 39°C it caused falls of between 5 and 20%. The authors speculate that the effect of sodium pentobarbitone on the blood flow at hyperthermic temperatures suggests that the lowered blood pressure decreases total tumour perfusion. If this is indeed the case in this work, then we have not contrived to avoid all anaesthetic artefacts.

Bleehen et al. (1977) have shown that cell survival, as measured by in vitro plating assay, is lower at the periphery of heated MISO-treated tumours than at the centre. We have shown that there is no difference between the MISO concentrations at the periphery and centre of such tumours. Thus the possibility that lower drug concentrations in hotter parts of the tumour can lower the overall concentration in heated tumours is eliminated. It appears that the difference is for the tumour as a whole, as would be indicated by the impedance of blood flow, by whatever mechanism.

The finding that the tumour drug concentration can be reduced by heating is important for drugs the in vivo cytotoxicity of which is enhanced by hyperthermia, such as MISO itself, cyclophosphamide (unpublished observations) and bleomycin (Marmor, 1979). In these cases, the potential effect must exceed the observed effect, since the heated tumours have received less drug than the controls. This must be considered in evaluating the exploitability of such an enhancement.

The observation that tumour size and implantation site in unheated tumours influence the MISO concentration in those tumours is also significant. It is perhaps to be expected that larger tumours would have lower total levels of drug, because part of their volume is composed of a necrotic centre with no intact blood supply, which will receive drug only by diffusion. Donelli et al. (1977) have shown in intramuscular Lewis lung tumours, for a variety of cytotoxic agents, that drug is mainly to be found in the viable part, with very little in the necrotic parts of the tumour. None the less it is surprising that the same increase in tumour size has more effect on the drug concentration in leg tumours than in flank tumours. This may be due to differences in geometry and hence the relative sizes of necrotic centres in leg and flank tumours with similar calculated volumes. The demonstration that, irrespective of site, large tumours have lower concentrations of drug than small ones may be of importance with regard to reports of investigations with other drugs where small tumours have shown greater sensitivity, estimated by growth delay, than large ones. This phenomenon has been reported for the response of the Lewis lung tumour to cyclophosphamide (Steel & Adams, 1975) and for the response of the P815X2 mastocytoma and the EMT6 tumour to
BCNU (Schenken, 1976; Twentyman, 1978). Perhaps the effect is partly a reflection of the actual tumour drug dose as well as the usually accepted explanation in terms of proliferative state.

The demonstration that MISO levels are the same in monitored and unmonitored tumours shows that the greater cell killing in heated monitored tumours treated with MISO (Honess et al., 1978) cannot be attributed to higher drug levels due to intra-tumoural haemorrhage. It may therefore be caused by better temperature equilibration of the monitored tumours due to conduction along the metal probe within the tumour, or by direct mechanical damage.

The main conclusions from this work are that the pharmacokinetics of MISO are unaffected by sodium pentobarbitone anaesthesia and very slightly affected by hyperthermia at a 44°C water bath temperature for 1 h. However, tumour levels of MISO are reduced. Also tumour levels of MISO in unheated tumours vary according to size and site. It seems unlikely that this phenomenon should be confined to MISO and, if it were shown to occur for other drugs also, it would at least partially account for some of the reports of variation in tumour chemosensitivity with size.

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