Effect of electrohydraulic and extracorporeal shock waves on gastrointestinal cancer cells and their response to cytotoxic agents

A Warlters, D L Morris, A Cameron-Strange, W Lynch

Abstract
Electrohydraulic and extracorporeal shock waves were used to treat the colorectal and gastric cancer cell lines LIM 2412 and MKN45. The effect on viability, cell proliferation, and the action of antitumour drugs was studied. Results showed that electrohydraulic and extracorporeal shock waves were cytotoxic to all cell lines and also caused considerable inhibition of cell proliferation. A significant additional reduction in cell viability was achieved by shock waves in cancer cells treated with methotrexate, 5-fluorouracil, and vincristine. These data indicate that shock waves may be worthy of further evaluation in the treatment of gastrointestinal cancers.

Despite continuing scientific research into the detection and treatment of gastrointestinal cancers, colon cancer remains the commonest cause of cancer deaths in non-smokers in many parts of the world: one in 25 Australians is expected to develop cancer of the bowel. There has been no major improvement in the overall five year survival rate in the last 40 years. The search for new treatments or methods of improving the results of existing therapy has included studies of chemosensitisation by temperature and radiotherapy.

In 1985, Russo et al first showed that high energy shock waves (HESW) suppressed tumour growth both in vitro and in vivo. Since then, several other groups have shown that HESW cause delayed cell growth and cell death in both normal and malignant cells in vitro. The effect of HESW on gastrointestinal cancers has received little attention: however, a group in the UK have shown some effect in a gastric cancer cell line. We are investigating the response of the colonic cancer cell line LIM 2412, as well as the gastric carcinoma cell line, MKN45, to extracorporeal shock waves (ECSW) and electrohydraulic shock waves (EHSW).

These forms of shock waves have been shown to alter cell membrane permeability, expose DNA, and damage mitochondrial structures. The possible effect of HESW on the sensitivity of cancer cells to cytotoxic drugs may be by membrane damage, allowing higher intracellular drug concentrations, or by effects on intracellular organelles, or a combination of these.

Methods
CELL CULTURE
LIM 2412, kindly donated by the Ludwig Insti-
All experiments were performed in triplicate and viability was assessed using trypan blue exclusion after 24 and 72 hour incubations.

PROLIFERATION ASSAYS

1×10^5 cells which had received either EHSW or ECSW treatment were aliquoted into 96 well microtitre plates. Fifty μl of each drug, at the concentrations outlined above, were then added to each group of shocked cells. All experiments were done in triplicate. Plates were then incubated for 72 hours and cells were pulsed with tritiated thymidine 1 μCi/well for 18 hours.

After this, cells were harvested and counted on a beta counter. The percentage inhibition of control was calculated as follows:

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\text{% Inhibition} = \frac{\text{DPM sample} \times 100}{\text{DPM control}}
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STATISTICAL TESTS

A two way ANOVA was done on viability data to assess significance.

Statistical significance of proliferation assays was determined by the proportional difference of means using Student's t test.

Results

EFFECT OF EHSW ON CELL VIABILITY AND RESPONSE TO CYTOTOXIC AGENTS

EHSW significantly reduced cell viability in LIM2412 (p<0.001) and MKN45 (p<0.01) up to 40% and 10% respectively (Fig 1). An additional 10–20% (p<0.05) reduction in LIM2412 viability was seen when 5-FU and VCR were present after HESW treatment, as shown in Table I.

There was little additional effect of 5-FU at concentrations higher than 0.1 mg/ml for both cell lines. However, there seemed to be a greater sensitivity to shock waves (Table I). MTX failed to have any further effect on either cell line and these data are not included in detail.

ECSW EFFECT ON CELL VIABILITY AND RESPONSE TO CYTOTOXICS

ECSW had no significant effect on either LIM2412 or MKN45 cell viability after 24 hour incubation, however, at 72 hours after treatment, LIM2412 cells showed a 10% (p<0.001) decrease in cell viability (Fig 2).

MTX and VCR failed to achieve any additional effect on viability of either cell line after shock wave treatment (data not included). 5-FU increased LIM2412 cell kill by approximately an additional 10% (p<0.05).

EHSW EFFECT ON CELL PROLIFERATION AND RESPONSE TO CYTOTOXIC AGENTS

EHSW had an appreciable effect on both MKN45 and LIM2412 cell proliferation in the presence of 5-FU and VCR (Figs 3 and 4). LIM2412 proliferation seemed to be sensitised to 5-FU and VCR after HESW treatment. This sensitisation was only observed in cells that had received higher shock wave doses (20 shocks, p<0.01).

MKN45 cell proliferation was significantly delayed in the presence of 5-FU (p<0.001) after treatment with 10 EHSW, compared with unshocked cells (Fig 5). Unlike LIM2412, MKN45 showed no response to VCR after shock wave treatment.

Inhibition of LIM2412 or MKN45 cell growth was not observed when MTX was present (data not presented).

ECSW EFFECT ON CELL PROLIFERATION AND RESPONSE TO CYTOTOXIC AGENTS

LIM2412 cell growth was unchanged by ECSW and/or exposure to 5-FU, MTX, or VCR.

MKN45 cell proliferation, however, was significantly inhibited after ECSW treatment (p<0.001). Figures 6 and 7 show that by treating the cells with either 5-FU or MTX, an increased level of growth inhibition can be achieved (p<0.05).
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**Discussion**

Our investigations indicate that in vitro both EHSW and ECSW cause an increase in cell death and decrease in cell proliferation of colonic and gastric cancer cells. This finding is supported by the viability and proliferation studies, which showed a decrease in cell viability and an inhibition of cell proliferation after shock wave treatment. Colonic cancer cells seemed more susceptible to EHSW than gastric cancer cells.

Shock waves have been associated with causing nuclear membrane and cellular membrane damage and attenuating nuclear chromatin. 5-FU acts by altering the RNA of cells, as well as decreasing the surface charge and transmembrane potential of tumour cells. Our results display an increased response in both LIM 2412 and MKN45 cells to 5-FU toxicity after EHSW and ECSW treatment. This enhancement of sensitivity to 5-FU may be related to the cytopathological changes in cells after exposure to shock waves, as outlined above. Both colonic and gastric cancer cells’ response to 5-FU was greater than responses to either MTX or VCR.

MKN45 cells seemed to be more susceptible to MTX than LIM 2412 cells, whereas VCR exerted a greater cytotoxic effect on LIM 2412 than on MKN45. It is already well established that different cell lines display varying responses to antitumour agents. There remain many unknowns – the optimal timing of shock waves to chemotherapy and subsequent treatment, the dosage of shock waves and how often this should be repeated.

Preliminary results in the in vitro effect of EHSW and ECSW in colonic and gastric cancer look promising: future work should include in vivo studies. The proposal of sensitising cells to chemotherapy by an accurately applied regional technique clearly could have merit.

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