The Post-Surgical Inflammatory Response Provokes Enhanced Tumour Recurrence: A Crucial Role for Neutrophils

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Abstract
Background/Aims: Peritoneal trauma activates a cascade of peritoneal defence mechanisms responsible for postoperative intra-abdominal tumour recurrence. After peritoneal trauma, inflammatory cells and soluble factors are present in the abdominal cavity and can be captured in lavage fluids. The present study evaluated which component enhances intra-abdominal tumour recurrence. Furthermore, we evaluated which inflammatory cells are present and studied the influence of anti-neutrophil serum (ANS) on peritoneal tumour recurrence. Methods: In a peritoneal trauma model in rats, postoperative lavage fluids were collected and separated into cellular and supernatant components. Both components were injected in naïve rats together with CC531s colon carcinoma cells. In a second experiment, rats were treated with one or three doses of ANS. Results: Intraperitoneal injection of naïve recipients with inflammatory cells or supernatant resulted in significant tumour recurrence. Severe peritoneal trauma provoked significant intra-abdominal neutrophil influx which could be prevented by ANS. Treatment with one dose did not affect blood cell counts and significantly reduced tumour recurrence. Treatment with three doses of ANS decreased blood lymphocytes, monocytes, and neutrophils and induced tumour load. Conclusions: Neutrophils play a crucial role in postoperative adhesion and growth of spilled tumour cells after surgical peritoneal trauma. Prevention of peritoneal neutrophil influx reduces local tumour recurrence.

Introduction

Locoregional tumour recurrence of colorectal carcinomas remains an important complication of potentially curative surgical intra-abdominal interventions. In a clinical study, Busch et al. [1] suggested an association between recurrent tumour disease and extent of surgical injury. It has also become evident from experimental studies [2, 3] that enhanced tumour cell adherence and tumour growth are inevitable repercussions of surgical peritoneal trauma. The pathogenesis of the processes responsible for postoperative intra-abdominal tumour recurrence is only partly clarified. In a previous study [4], we demonstrated that within a few hours after infliction of peritoneal trauma, factors in the abdominal cavity could be captured in lavage fluid which enhance tumour recurrence in naïve, non-operated recipients. The inflammatory reaction after surgery is not only responsible for the wound-healing process, but also induces tumour recurrence. During this inflammatory response, perito-
neal lymphocytes, submesothelial monocytes, neutrophils, and mesothelial cells act under the control of locally expressed cytokines, chemokines, and adhesion molecules [5, 6].

In the present study, we focus our attention on the individual capacity of inflammatory peritoneal cells and soluble factors to ascertain which element is mainly responsible for enhanced tumour recurrence. Secondly, assuming that post-traumatic intra-abdominal influx of neutrophils is an important factor in the dynamic cascade of peritoneal defence, possibly responsible for local tumour recurrence, prevention of polymorphonuclear neutrophil (PMN) influx might influence this process. Therefore, a second experiment was performed to evaluate whether post-traumatic intra-abdominal PMN influx could be reduced by treatment with anti-neutrophil serum (ANS) and, if so, whether this reduction could influence postoperative tumour development.

**Materials and Methods**

**Animals**

Female inbred WAG rats of reproductive age weighing 140–180 g were obtained from Harlan-CPB (Austerlitz, The Netherlands). They were bred under specific-pathogen-free conditions, kept in a standard laboratory environment (temperature 20–24°C, relative humidity 50–60%, 12-hour light/dark cycles), and fed with standard rat food and water ad libitum. The experimental protocol was approved by the Animal Experiments Committee under the national Experiments on Animals Act and adhered to the rules laid down in this national law that serves the implementation of ‘Guidelines on the Protection of Experimental Animals’ by the Council of Europe (1986), Directive 86/609/EC.

**Tumour**

The tumour CC531s is a moderately differentiated, weakly immunogenic colonic adenocarcinoma induced in WAG rats by 1,2-dimethylhydrazine [7]. It is transplantable in syngeneic WAG rats. The tumour is maintained as a cell culture in RPMI 1640 medium (RP MI) supplemented with 5% fetal calf serum (virus and Mycoplasma screened), 1% penicillin (5,000 U/ml), 1% streptomycin (5,000 U/ml), and 1% L-glutamine (200 mmol). RPMI and all supplements were obtained from Life Technologies (Breda, The Netherlands). The cells were passaged once a week using trypsin (0.05%) and EDTA (0.02%). Before use in vivo, the tumour cells were harvested from stationary cultures by gentle trypsinization (5 min, 37°C), centrifugation (5 min, 700 g), and resuspension in RPMI, providing cell suspensions with a viability >90%. Tumour CC531s is relatively insensitive to chemotherapy, but is sensitive to the effects of biological response modifiers.

**Gathering of Lavage Fluid for Passive Transfer Experiment**

Under isoflurane anaesthesia, in 14 rats a laparotomy was performed: exposure and rubbing of the uterine horns and a 5-cm part of the small intestine with surgical Medipress gauze inflicted subsequent trauma to the peritoneum [4]. Rubbing was performed with a device enabling the application of a constant pressure of 120 g/m². In this way, a standardized degree of peritoneal trauma can be inflicted. The abdomen was closed in one layer with silk 2-0 sutures. After 5 h, a second laparotomy was performed, during which the abdominal cavity was lavaged with 5 ml RPMI. After massaging the abdomen, the remaining fluid was aspirated, pooled, and kept on ice until further processing.

**Experimental Design of Passive Transfer (Experiment 1)**

The collected post-trauma lavage fluid was centrifuged, and the cell pellet was resuspended to the original volume with RPMI and then divided into a ‘supernatant’ containing soluble components produced after surgical trauma and a ‘cellular’ component containing the different cell types present in the abdominal cavity after surgical trauma.

Subsequently, 24 rats were divided into three groups: one group served as control group receiving RPMI; the second group was acceptor for the supernatant of the post-trauma lavage fluid, and the third group was acceptor for the cellular component. Of all three components, 3 ml was injected intraperitoneally together with 0.5 million CC531s cells (in 0.5 ml RPMI) without opening the abdominal cavity. In this way, the factors contained in the different components represented the mediators after surgical abdominal trauma, without inflicting additional trauma (fig. 1).

**Effect of ANS Treatment on Cell Content in Peritoneal Cavity and Blood (Experiment 2)**

To investigate the influence of ANS treatment on the intra-abdominal neutrophil count and on the rat immune system, the following procedures were performed. Under isoflurane anaesthesia, 65 rats underwent a laparotomy. In 5 rats no peritoneal trauma was inflicted. In 20 rats, standardized severe peritoneal trauma was inflicted without treatment. In 20 rats, severe peritoneal trauma was inflicted (on day 0) in rats treated with three intraperitoneal doses of ANS (1 ml/kg) on days –1, 0, and +1. In 20 rats severe peritoneal trauma was inflicted after a single intraperitoneal injection of ANS on day –1. After 5, 72, 96, and 144 h, 5 rats of each group were operated for the second time. During this second laparotomy, the abdominal cavity was lavaged with 5 ml RPMI. After massaging the abdomen, the remaining fluid was aspirated and individually kept on ice until further processing. Blood samples were obtained as well, by cardiac puncture.

The collected lavage fluid samples were separated in a supernatant component and a cellular component by centrifugation (1,500 rpm, 5 min). The cellular component was resuspended in RPMI, the total cell amount was determined, and haematoxylin-
and eosin-stained cytocentrifuge slides were made for cell differentiation. At ×100, 100 cells were counted in duplicate and classified into granulocytes (neutrophils, eosinophils, basophils, and mast cells) and lymphoid cells (mononuclear phagocytes and lymphocytes). Total blood leucocyte counts were determined with a micro cell counter, and duplicate differential counts were carried out on May-Grünwald- and Giemsa-stained blood smears.

Intraperitoneal Tumour Cell Adhesion and Growth after Treatment with ANS (Experiments 3 and 4)

A reproducible animal experimental model was used [4]. Briefly, under isoflurane anaesthesia and aseptic conditions, a laparotomy was performed by a midline incision of 5 cm. Both uterine horns were exposed, not touched or rubbed with surgical Medipress gauze, and sutured to the lateral peritoneum both proximally and distally using Surgilene 6-0 sutures. In this way, a standardized degree of peritoneal trauma was inflicted. To induce peritoneal metastases, 0.5 × 10^6 CC531s tumour cells, in 1 ml RPMI, were injected intraperitoneally before closing the abdomen. The abdomen was closed in two layers with 5-0 polyglycolic acid and 2-0 silk sutures.

In experiment 3, 9 rats underwent severe peritoneal trauma without treatment. Nine rats underwent severe peritoneal trauma which were treated with an intraperitoneal injection of ANS on day –1. Before closing the abdomen 0.5 × 10^6 CC531s tumour cells, in 1 ml RPMI, were injected intraperitoneally in all rats.

In experiment 4, similar groups were formed, with the difference that the treated group received intraperitoneal ANS injections on days –1, 0, and +1. Again, before closing the abdomen, 0.5 × 10^6 CC531s tumour cells were injected into the abdominal cavity.

Evaluation of Tumour Load

Three weeks after injection of CC531s tumour cells, all rats were sacrificed, and the intraperitoneal tumour load was scored semi-

Fig. 1. Experimental design of the passive transfer experiment.
quantitatively at the following peritoneal sites: right uterine horn, left uterine horn, subcutaneously (at the site of the operative scar), parietal peritoneum (on the lateral abdominal wall sides, where no uterine horns were fixed), kidney, liver, retroperitoneum, and omentum. Scoring was performed by two blinded observers, using a tumour scoring system derived from the peritoneal cancer index (Steller score) [4, 10], ranging from 0 to 5 per abdominal site. For each rat, the score at all peritoneal sites was summarized, from which a mean total tumour load per rat could be estimated.

**Statistics**

Statistical analysis was performed using one-way ANOVA tests to determine overall differences. If an ANOVA test was significant at the 5% level, the Newman-Keuls post hoc test was carried out to make a comparison between groups. Data are expressed as mean ± SEM. For tumour load, median and range were calculated, and statistical analysis was performed using the non-parametric Kruskal-Wallis analysis of variance to determine overall differences, followed by the non-parametric Mann-Whitney U test to compare differences between groups. Statistical significance was defined as p < 0.05.

**Results**

**Effect of Post-Traumatic Lavage Fluid on Peritoneal Tumour Load**

After intraperitoneal injection of the lavage fluid samples collected after surgical trauma, a diffuse peritoneal tumour load was found in all groups (fig. 2). When compared with the control group (RPMI), the cellular factors caused significantly enhanced tumour recurrence (p < 0.01), as well as the supernatant (p < 0.05). Injection of tumour cells with RPMI alone resulted in a median total Steller score of 0 (range 0–3), whereas injection with the supernatant or cells resulted in Steller scores of 1 (range 0–5) and 3 (range 0–5), respectively. Injection with the cellular component of the lavage fluid also resulted in a significantly higher tumour load as compared with the supernatant (p < 0.05).

**Cell Content in Peritoneal Cavity and Blood after Surgery and the Effect of ANS Treatment**

There was a significant increase in total intra-abdominal cell count after infliction of severe peritoneal trauma up to 144 h after the operation (fig. 3). Figure 3 also shows that intraperitoneal administration of three doses of ANS as well as one dose of ANS significantly decreased the total intra-abdominal cell count after infliction of severe peritoneal trauma for at least 96 h postoperatively (p < 0.01). After treatment with one dose of ANS, the total cell count increased earlier than after treatment with three doses, the differences being significant at 96 h (p < 0.01) (fig. 3). A marked increase of the percentage of neutrophils was seen in the severely traumatized group (p <
This increase was seen till 96 h postoperatively (fig. 4). Treating the rats with three doses as well as one dose of ANS did annul this increase (fig. 4).

Figure 5 shows the results of blood differential cell counts in the four groups at different time points. Similar results were found with lymphocyte, monocyte, or neutrophil counts. Treatment with three doses of ANS significantly decreased these cell counts for a period of at least 96 h (p < 0.01). This effect was not seen after treatment with one dose of ANS.

**Intraperitoneal Tumour Development after Treatment with ANS**

As shown in figure 6, a marked tumour load is seen after severe trauma. Treatment with a single dose of ANS significantly reduced the median tumour load in severely traumatized rats (p < 0.01). Surprisingly, intraperitoneal injection of three doses of ANS induced even more tumour load (p < 0.001).

**Discussion**

In this study, a cell-seeding model was used to mimic the clinical situation of tumour cell spill during tumour resection.

Following peritoneal trauma, a variety of cytokines, growth factors, and other inflammatory mediators are produced by activated mesothelial cells and by stamped-inflammatory cells [11]. The production of mesothelial cell and inflammatory cell derived chemokines will
cause an early posttraumatic migration of PMNs and monocytes to the injured peritoneal cavity, in order to promote the peritoneal healing process [12]. However, these mediators and recruited cells not only serve peritoneal healing, but, as shown in a previous study [4], could enhance tumour recurrence in naïve recipients as well. The current study demonstrates that the cellular components of the lavage fluid collected after inflicting surgical trauma, i.e., inflammatory cells, lead to enhanced tumour recurrence. More detailed analysis of the cellular fraction of these lavage fluids revealed a trauma-related influx of PMNs into the abdominal cavity (fig. 4). PMNs use both oxygen-dependent and oxygen-independent processes in killing micro-organisms, but these processes also (further) damage surrounding host tissue [13, 14]. In vitro, adhesion of activated PMNs to a mesothelial monolayer has been shown to induce retraction, gap formation, and detachment, ending with substantial mesothelial cell injury and exposure of extracellular matrix components, thereby already creating a preferential site for adhesion of free tumour cells [15]. A relation has been demonstrated between extent of tissue trauma and tumour recurrence. In order to diminish tissue trauma and tumour recurrence, minimally invasive surgery is promoted and proved promising in rat models [4, 16]. In addition, laparoscopic surgery appears to have less impact on the cellular components of the immune response than laparotomy [3]. A recent meta-analysis [17] reported faster postoperative recovery with laparoscopic surgery for colorectal cancer as compared with open surgery, though in this meta-analysis no differences were seen in recurrence rates. However, only a few studies published long-term results with a favourable trend towards laparoscopy [18].

The coincidence of post-traumatic intra-abdominal PMN influx with tumour cell adhesion and growth, however, is no solid proof for the role of PMNs in these pathogenetic processes. Effective inhibition of tissue injury by PMNs has been achieved [19, 20]. Dovi et al. [21] even showed that wound healing was accelerated in PMN-depleted mice. However, whether inhibition of PMNs affected tumour recurrence has not been investigated. The present study demonstrated that a single intraperitoneal dose of ANS could prevent the influx of PMNs without influencing the blood cell count. This study showed that a selective reduction of post-traumatic PMN influx, without causing systemic immunosuppression, was possible and could indeed significantly lower tumour adhesion and growth. Averting the post-traumatic intra-abdominal PMN influx by intraperitoneal injection of three doses of ANS significantly increased tumour recurrence, at first to our surprise. However, treatment with three doses of ANS also significantly decreased blood lymphocyte, monocyte, and PMN counts, thereby seriously compromising the rat immune system. It is conceivable that this immunosuppression promotes tumour growth. We observed earlier that immunosuppression leads to enhanced growth of the tumour used in the current experiments [22].

In conclusion, these studies demonstrated that the early inflammatory sequelae after surgery promote tumour recurrence and that this effect is mainly based on the cellular component of the inflammatory process. Preventing the postoperative influx of PMNs without affecting the systemic immune response reduced peritoneal tumour recurrence.

References


