THE EFFECT OF PERIOPERATIVE ANALGESIA WITH OMNOPON AND PARECOXIB ON THE ENDOCYTIC ACTIVITY OF MURINE PHAGOCYTES ON THE MODEL OF TUMOR SURGERY

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We used the model of surgical tumor removal to compare the effect of anesthesia with opioid analgesic omnopon and selective cyclooxygenase-2 inhibitor parecoxib on the endocytic activity of phagocytes of different localization sites. 50 C57/black mice were transplanted with Lewis lung carcinoma in the hind paw pad. After 22 days, the tumor paw was amputated. Analgesics (omnopon 10 mg/kg, parecoxib – 20 mg/kg) were administered 30 min before the operation and once per day for 3 days after the surgery. Assessment of the endocytic activity of phagocytes was performed by FACS analysis before the surgery, at days 1 and 3 after the surgery. It was found that parecoxib analgesia maintained the endocytic activity of blood and spleen phagocytes in the postoperative period. At day 3 after the surgery in parecoxib-treated animals phagocytic activity of splenic granulocytes were 2.2 times higher compared to that in the group receiving opioid analgesia. Phagocytic indices of monocytes in parecoxib-treated mice were also 1.6 and 2.5 times higher for blood and spleen monocytes, respectively. Thus, parecoxib analgesia maintained the activity of blood and spleen phagocytes in mice after the surgical tumor removal at a much higher level as compared with the omnopon analgesia.

Keywords: phagocytic activity, perioperative analgesia, parecoxib, opioid drugs.

INTRODUCTION

Surgical excision is the mainstay of treatment for potentially curable solid tumours, but metastatic disease remains the most prevalent cause of cancer-related death in these patients. Recent studies demonstrated that surgical and traumatic injury profoundly affects both innate and adaptive immune responses [4]. The immune suppression following an excessive inflammatory response after surgery can create conditions for tumor immune-evasion, thus, promoting the metastatic process.

In addition, the prolonged inflammation promotes a spread of tumor cells [2]. Thus, minimalization of postoperative inflammation could reduce the risk of relapse and meta-
stasis. Phagocyte system is essential for both development and resolution of inflammatory response [7]. During the onset of inflammation, phagocytes polarize to pro-inflammatory phenotype with substantial production of reactive oxygen species and pro-inflammatory cytokines. These lead to the elimination of inflammatory triggers and switch to anti-inflammatory phenotype of phagocytes with a subsequent resolution of inflammation. Potent endocytic activity, which is among the features of this phenotype, serves for the clearance of neutrophils dying of netosis, and tissue debris which is the result of inflammatory cells destruction. A decrease in the endocytic activity of phagocytes may be the reason for chronization of inflammation, thus, promoting tumor progression [10].

Surgical stress factors that suppress immune cells functions, include tissue trauma, pain, and premedication pharmaceuticals, such as anaesthetic drugs and opioid analgesics. Despite opioids are often used for treatment of surgical and cancer pain, recent studies suggest that opioid administration have inhibitory effects on antibody production, natural killer cell activity, cytokine expression, and functional activity of phagocytes [4, 8]. Consequently, there is a search of alternative analgesics which can at least partially substitute opioids for perioperative pain relief in tumor surgery in order to maintain immune function of cancer surgical patients on high level. In this respect, cyclooxygenase-2 (COX-2) inhibitors are of a particular interest as they were also reported to influence cancer incidence [3, 13].

In this study, we aimed to compare the effect of the perioperative analgesia with opioid drug omnopon and selective COX-2 inhibitor parecoxib on the endocytic activity of phagocytes using a murine model of surgical tumor excision.

MATERIALS AND METHODS

50 males of C57/black mice (18–22 g, 1.5 months old) were used. Animals were housed in an animal care facility of the National Cancer Institute, Ukraine. All procedures with animals were performed in accordance with the principles of humanity as it was written in “General principles of animal experimentation” approved by the National Congress on Bioethics (Kyiv, 2001–2007) and in accordance with Council directive of 24 November 1986 on the approximation of laws, regulations and administrative provisions of the Member States regarding the protection of animals used for experimental and other scientific purposes (86/609/EEC). The Bioethical Committee protocol No. 60 of the National Cancer Institute was approved in June 16, 2015. Lewis lung carcinoma (LLC) cell line was used as an experimental tumor model. LLC cells were kindly provided by the Bank of Cell Line of R. E. Kavetsky Institute of Experimental Pathology, Oncology and Radiobiology, NAS of Ukraine. LLC cells were transplanted subcutaneously into the right hind paw in the amount of 4 × 10^5 cells per mouse. After the tumor cell transplantation experimental animals were randomized by weight and assigned to 3 groups (15 animals per group): experimental group I (received for perioperational analgesia 10 mg/kg of opioid drug omnopon), experimental group II (received for perioperational analgesia 20 mg/kg of COX-2 inhibitor parecoxib) and control group (received equivalent volume of saline). 5 mice were leaved without tumor transplantation and used as an intact control.

Surgical removal of tumor was performed on 22nd day after inoculation. Mice were anesthetized by ketamine (25 mg/kg intraperitoneally), tumor paw was ligated and amputated on the level of knee joint. Analgesic drugs (or saline in control group) were injected
intraperitoneally 30 min before the tumor excision and once a day for 3 days after the surgery (4 injections in total). 5 mice per group were euthanized by cervical dislocation at 3 time-points: before the surgery, 1st and 3rd day after surgery. Blood and spleens were taken for the evaluation of monocytes and granulocytes phagocytic activity.

Phagocytic activity was assessed by FACS analysis. Briefly, 50 µ of blood or spleenocytes’ suspensions (2×10^6 cells/ml) were mixed with 40 µ of FITC-labeled *Staphylococcus aureus* Cowan I (1×10^7 cells/ml) and incubated at 37 °C for 30 min. Then 2 ml of cold erythrocyte lysis solution containing EDTA was added for 10 min to stop the reaction, and then cells were washed twice with phosphate buffered saline. Results were assessed using FACSCalibur flow cytometer and CellQuest software (Becton Dickinson, USA). Granulocytes or monocytes were gated according to forward and side scatter. Phagocytosis percentage was measured as a percentage of fluorescence emitting cells (*S. aureus* engulfed cells) in the respective gate, and phagocytic index was measured as a geometric mean fluorescence of these cells (which represents the mean number of bacterial cells engulfed by one phagocyte).

Statistical analysis was conducted in Statistica 10 software (StatSoft Inc., USA) using the one-way analysis of variance (ANOVA) and Wilcoxon test. Differences with type I error <0.05 were considered as statistically significant.

RESULTS AND DISCUSSION

Surgical trauma and pharmaceuticals used markedly perioperative reduce the activity of the immune cells, including phagocytes, such as neutrophils, monocytes and macrophages [5]. Phagocytes of peripheral blood and spleen play a key role in the resolution of postoperative inflammation, thus reducing the risk of relapse and metastases. Peripheral blood phagocytes eliminate apoptotic cells and residual tumor cells which can spread to circulation after the surgery. Circulating antigens are also filtered and eliminated in spleen by splenic phagocytes [1].

In the present study, we observed a decrease in phagocytosis percentage of granulocytes in peripheral blood at 1st day after the surgical removal of transplanted tumor with its subsequent recovery at 3rd day (Fig. 1). This can be explained by surgical trauma and blood loss with further reparative process. Analgesia with omnopon, however, notably depresses this postoperative recovery of phagocytosis. Thus, phagocytosis percentage of peripheral blood granulocytes in control and parecoxib treated mice to postoperative 3rd day recover almost to preoperative levels, while in omnopon treated mice its continue to decrease and at day 3 was 20 % lower compared to control group (p<0.05). Postoperative decrease in the phagocytic activity of blood granulocytes was also more pronounced in the omnopon group (Fig. 1, A) where at 1st day after the surgery phagocytic index was 2.3 times lower compared to parecoxib group (p<0.05). All these findings are consistent with data indicating that morphine can affect the myeloproliferation and inhibit functional activity of neutrophils (reactive oxygen species production, complement receptor synthesis and phagocytosis) [2].

Proportion of splenic granulocytes with phagocytic capacity decreased significantly after the surgery in control group and omnopon group while in parecoxib group it changed only slightly throughout the perioperative period (Fig. 1, B). In addition, phagocytic activity of these cells at 3rd day post-surgery was 2.2 times higher in parecoxib treated animals compared to omnopon analgesia group (p<0.05).
Fig. 1. Phagocytic activity of murine granulocytes before and after surgical removal of tumor: phagocytosis percentage of granulocytes of peripheral blood (A) and spleen (B); phagocytic index of granulocytes of peripheral blood (C) and spleen (D). t0 – before the surgery; t1 – 1st day after the surgery; t2 – 3rd day after the surgery. The values presented are means ± SEM (n = 5); * – p<0.05 compared to omnopon-treated group

Mononuclear phagocytes were also affected by surgical stress (Fig. 2). Relative quantities of phagocytically active monocytes in peripheral blood were markedly reduced at day 1 after the operation compared to preoperative values in all studied groups (Fig. 2, A), which also could be a consequence of blood loss and migration to inflammation sites. At 3rd day after the surgery phagocytosis percentage of circulating monocytes have increased compared to 1st day levels, however, in omnopon treated mice this recovery reaches only (47.3±4.7)% as compared to (59±3.8)% in control group and (61±4.9)% in parecoxib group (p<0.05). At the same time, the amount of splenic mononuclear phagocytes during postoperative period was higher in parecoxib treated mice (Fig. 2, B) (at 3rd day post-surgery these values were (61.0±3.1)% in parecoxib group compared to (52.5±3.2)% and (45.5±2.9)% in control and omnopon groups, respectively, p<0.05).

Postoperative values of phagocytic indices of both peripheral blood and splenic monocytes were also higher in mice receiving parecoxib analgesia compared to omnopon and saline treated animals. Phagocytic activity in parecoxib group at 3rd day after the surgery was 1.6 times higher for blood monocytes, and 2.5 times higher for splenic monocytes as compared to omnopon group (p<0.05).
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Fig. 2. Phagocytic activity of murine monocytes before and after surgical removal of tumor: phagocytosis percentage of monocytes of peripheral blood (A) and spleen (B); phagocytic index of monocytes of peripheral blood (C) and spleen (D). t0 – before the surgery; t1 – 1st day after the surgery; t2 – 3rd day after the surgery. Values presented are means ± SEM (n = 5); * – p<0.05 compared to omnopon-treated group

Maintenance of high phagocytic activity of neutrophils and monocytes is crucial for the resolution of inflammation in postoperative period. Malfunction of mechanisms regulating inflammation resolution leads to chronization of inflammatory process accompanied by decline of adaptive immune response and promotion of metastasizing [11, 12].

Perioperative analgesia with COX-2 inhibitor parecoxib resulted in preservation of phagocytic function of granulocytes and monocytes on higher levels compared to control animals. COX-2 is one of the key enzymes in the development of inflammation and its overexpression may lead to prolonged inflammatory process [6]. COX-2 product prostaglandin E2 (PGE2) exerts an inhibitory effect on the endocytic activity of phagocytes [12], and is suggested to link chronic inflammation with tumor growth promotion [6, 12]. Inhibition of COX-2 by parecoxib during perioperative period may neutralize negative influence of PGE2 on the phagocytic function.

On contrary, analgesia with opioid drug omnopon led to more potent decrease in the postoperative endocytic activity and quantity of phagocytes in comparison with control and parecoxib treated animals. These results correspond to literature data confirming the negative effect of opiates on the phagocytic activity of monocytes and neutrophils [9, 4].
To summarize, obtained results put forward the COX-2 inhibitor parecoxib as a promising alternative to the opioid drugs in the perioperative analgesia. Further studies may reveal the benefit of parecoxib in the reduction of metastases and the postoperative complications in surgical cancer patients.

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C57/black, которым перевивали карциному легких Льюис в подушечку задней лапы. На 22 сутки лапу с опухолью ампутировали. Анальгетики (омнопон в дозе 10 мг/кг, парекоксб – 20 мг/кг) вводили за 30 мин до операции и 1 раз в сутки в течение 3 дней после операции. Оценку эндоцитарной активности фагоцитов проводили методом проточной цитометрии за сутки до, на 1 и 3 сутки после операции. Было установлено, что применение парекоксб способствует поддержанию эндоцитарной активности фагоцитов крови и селезенки в послеоперационном периоде. На 3 сутки после операции в группе животных, получавших для обезболивания парекоксб, фагоцитарная активность гранулоцитов селезенки была в 2,2 раза выше по сравнению с группой, получавшей опиоидную аналгезию. Фагоцитарные индексы моноцитов при обезболивании парекоксбом также были выше в 1,6 и 2,5 раза выше для моноцитов крови и селезенки соответственно. Таким образом, при аналгезии парекоксбом активность фагоцитов крови и селезенки мышей после операции сохраняется на более высоком уровне по сравнению с применением омнопона.

**Ключевые слова:** фагоцитарная активность, периоперационная аналгезия, парекоксб, опиоидные препараты.