Research Article
Viability of Airborne Tumor Cells during Excision by Ultrasonic Device

Masakazu Hashimoto, Tsuyoshi Kobayashi, Hirotaka Tashiro, Shintaro Kuroda, Yoshihiro Mikuriya, Tomoyuki Abe, Yuka Tanaka, and Hideki Ohdan

Department of Gastroenterological and Transplant Surgery, Hiroshima University Hospital, Hiroshima, Japan

Correspondence should be addressed to Tsuyoshi Kobayashi; tsukoba@hiroshima-u.ac.jp

Received 19 November 2016; Revised 26 February 2017; Accepted 23 March 2017; Published 9 April 2017

1. Introduction

In recent years, surgery using a laparoscope has been compared with the traditional laparotomy, showing that its curative outcome is not inferior. Laparoscopic surgery has become less invasive due to the improvements in laparoscopic equipment and the accumulation of clinical experience. The advantages of laparoscopic surgery leave no doubt and will continue to develop in the future.

However, peritoneal dissemination and port-site metastasis were reported in laparoscopic surgery for the first time in 1978 [1] and have since been reported many times in such procedures as gastroenterological surgery [2, 3], gynecological surgery [4], and surgery of the urinary organs [5], with an incidence of almost 1%.

This problem with the laparoscopic tool is thought of as one cause of relapse, and it seems that it is necessary to further evaluate the ultrasonically activated scalpel (UAS) used to resect the tumor and the lymph nodes. Previously, there was a report that no viable carcinoma cells exist in the mist that the tool creates [6]. However, this evaluation may not be sufficient, because the viability of airborne cells was assessed only by electron micrograph.

We examined whether living cancer cells exist in the mist that is dispersed by the UAS, for the evaluation of the safety of laparoscopy-assisted surgery.

2. Materials and Methods

2.1. Cell Line and Culture Conditions. Human hepatoma cell line HepG2 was obtained from the American Type Culture Collection (ATCC). The cells were grown in Dulbecco's Modified Eagle Medium (DMEM; Invitrogen, Carlsbad, CA, USA) containing 10% fetal calf serum (Invitrogen, Carlsbad, CA, USA) in a humidified atmosphere of 95% air and 5% CO₂ at 37°C.

2.2. Animal and Tumor Preparation. We used 20-week-old NOD-Cg-Rag1<sup>tm1Mom</sup>IL2rg<sup>tm1wjl/Sj</sup> mice for the tumor model. HepG2 cells were injected subcutaneously into the flank...
of the mice at $1 \times 10^6$ cells/mouse. After about one month, the tumor size was about 2 cm and was used for the following experiment. All animal experiments were performed in accordance with the guidelines set by the United States National Institutes of Health (1996).

2.3. Collection of Dispersed Cells. The tumor was excised directly by UAS using Harmonic Laparoscopic Coagulation Shears (LCS) (Johnson and Johnson, New Brunswick, NJ, USA) and PowerStar scissors (Johnson and Johnson, New Brunswick, NJ, USA). This instrument was used in high power mode (“LCS high”), high power mode with a suction unit (“LCS high + suction”), and low power mode (“LCS low”), and these were compared to excision using PowerStar bipolar scissors (“PowerStar”). Resection of the tumor mass, which was excised from mice, was performed in a 50 mL Falcon tube and the mist was collected in 3 mL of DMEM, and they were isolated to three experiments system. As a control, the same procedure was conducted using the cultured cell line. After collection, samples were stored at 37°C to ensure that the cell survival rate would not decrease and used for the following procedure as quickly as possible.

2.4. Cell Counting

2.4.1. Viable Cell Counting. The number of viable cells was determined by trypan blue dye exclusion using a microscope.

2.4.2. Immunofluorescence Staining. A coverslips previously coated with fibronectin was placed in a 35 mm dish with DMEM, and the surgical mist that had been collected was added. After incubation overnight, the tumor cells were fixed in 4% paraformaldehyde and then washed with PBS and permeabilized in 0.1% Tween-20 (Sigma-Aldrich, St. Louis, MO, USA). After being blocked with 10% bovine serum albumin, the cells were incubated with anti-Ep-CAM antibody (EBA1) (Santa Cruz Biotechnology Inc., Dallas, TX, USA), which was detected with fluorochrome-coupled secondary antibody (Alexa Fluor 488 Goat Anti-Mouse IgG; Molecular Probes Invitrogen, Carlsbad, CA, USA). The cells were observed under a fluorescence microscope (BZ-8000) and analyzed with BZ-II Analyzer software (KEYENCE Japan, Osaka, Japan).

2.5. Flow Cytometric Analysis. The number of cancer cells generated in a mist among the $2 \times 10^4$ viable cells collected previously was determined by flow cytometry. To distinguish the epithelial cells from blood cells, HepG2 cells were stained with anti-Ep-CAM antibody and anti-CD45 FITC (eBioscience, Inc., San Diego, CA, USA) monoclonal antibodies. We counted the number of cells stained for Ep-CAM. Dead cells were excluded from the analysis by propidium iodide staining. All analyses were performed using a FACSCalibur cytometer (BD Bioscience, Mountain View, CA, USA).

2.6. Statistical Analysis. Quantitative data are represented as the mean ± SE. Comparisons between data within the same experiments were analyzed using the Student's $t$-test. A $p$ value of <0.05 was considered statistically significant. All statistical analyses were performed using the JMP 10 software for Windows (SAS Institute Japan).

3. Results

3.1. Quantification of Viable Cells from Tumor Excision. Each tumor was excised with LCS high, LCS high + suction, LCS low, and PowerStar methods, and the tumor cells from the site were collected and counted (Figure 1). LCS high excision resulted in $5.6 \pm 2.9 \times 10^4$ cells/mL, LCS high + suction in $0.4 \pm 0.2 \times 10^4$ cells/mL, LCS low in $2.3 \pm 0.9 \times 10^4$ cells/mL, and PowerStar in $0.8 \pm 0.4 \times 10^4$ cells/mL. When removing the tumor using LCS high, a large amount of cell-containing mist was generated and thus many tumor cells were collected.

3.2. Immunofluorescence Staining. The number of cells excised by LCS high was significantly greater than that by LCS high + suction, LCS low, and Power Star methods. Based on the number of viable cells released, the methods can be ordered: LCS high (28.9 ± 8.7 cells/high power field [HPF]), LCS high + suction (8.0 ± 8.0 cells/HPF), LCS low (2.7 ± 1.9 cells/HPF), and PowerStar (0.7 ± 0.3 cells/HPF) (Figure 2).

3.3. Flow Cytometric Analysis. There was a significant difference between the number of cells obtained by LCS high and LCS high + suction, and between LCS high + suction and powerstar: LCS high: (28.5 ± 24.7/2.0 × 10^4 cells), LCS high + suction: (7.1 ± 14.8/2.0 × 10^4 cells), LCS low: (4.2 ± 7.1/2.0 × 10^4 cells), and powerstar: (0.5 ± 0.8/2.0 × 10^4 cells) (Figure 3).
Surgery Research and Practice

Figure 2: Immunofluorescence staining (anti-Ep-CAM antibody). In LCS high, LCS high + suction, LCS low, and PowerStar methods, viable cancer cells were detected. The LCS high resulted in significantly more viable cancer cells in comparison with LCS high + suction, LCS low, or PowerStar (*p < 0.05). The white bar is 50 μm.

4. Discussion

This study confirmed that carcinoma cells existed in the mist generated when a UAS (LCS) was used to remove the tumor. We also showed that cell viability and cell adhesion were preserved in these cells.

Previously, it was shown that these cells were unable to survive because of damage to the cell membrane. This was shown by electron microscopy, in a study on the existence of carcinoma cells in surgical mist examined in vitro [6]. We proved that living cells exist among the dispersed material by three methods in the current study. Immunofluorescence staining sufficiently confirmed cell viability and adhesive ability. Moreover, when observed over time, these cells had a proliferation rate similar to the original cell line (data not shown). The number of carcinoma cells was determined by the exclusion of dead cells and blood cells, using a flow cytometric method. In every detection and counting method we used, there was a similar result, which is the fact that the LCS high power excision method results in a danger of scattering many carcinoma cells.

In a clinical study, a randomized controlled study with a mean follow-up of 21 months compared port-site metastases and recurrence rates of laparoscopically resected colon carcinomas (44 cases) with conventional open colectomies (47 cases). This study showed no wound- or port-site metastases, with similar recurrence rates in both groups: 16.1% versus 15% [7]. The rate of peritoneal dissemination associated with port-site metastasis is not increased by the use of laparoscopic procedures, as reported in a randomized trial comparing laparoscopic-assisted colectomy and open colectomy for the treatment of nonmetastatic colon cancer [8]. This means that the dissemination is related to the advanced nature and the biological behavior of the tumor, rather than to the laparoscopic technique itself [9, 10].

However, there are some hypotheses for the mechanism by which peritoneal dissemination and port-site metastasis happens, and Castillo and Vitagliano reported that the following may all play a role: (a) tumor-related factors, (b) wound-related factors (including the immune response), and (c) surgery-related factors [11]. Depressed immune function may also contribute to tumor recurrence and metastasis. Overall immune function is diminished in the perioperative period because of several mediating factors, including anesthetic agents, opioids, surgical trauma, blood transfusions, temperature changes, pain, and psychological stress [12]. In animal models, surgical trauma has been shown to reduce natural killer cell activity and promote tumor metastasis [12, 13]. It was thought that these clinical reports may not apply to laparoscopic surgery, which results in a smaller wound and could maintain immunity.

UAS were first introduced and widely used in laparoscopic surgery and robot assisted surgery, and they are a quite useful surgical apparatus in the surgical field [14, 15]. UAS is effective in not only laparoscopic surgery but also conventional open surgery. Advantages of UAS over conventional electrosurgical instruments include reduced operating times, less intraoperative blood loss, and reduced
leakage from the cut surface of organs in a variety of surgeries such as cholecystectomy, thyroidectomy, gastrectomy for gastric cancer, colorectal surgery, and pancreatic and hepatic resections [16–22]. It seems that it is important to understand the characteristics of this device used regardless of the type of surgery; this is true in laparoscopic surgery or laparotomy.

As a method for preventing dissemination, we showed that we can decrease the number of cells by changing from the high power mode of the LCS to low power and by using suction adjacent to the excision site. In clinical settings, when excising a tumor, it is thought that the number of carcinoma cells that are dispersed into the intraperitoneal cavity can be decreased by reducing the power used or using suction.

The present study had several limitations. Avoiding direct manipulation of the tumor, which is the main precaution surgeons must take during oncologic surgery, was not considered in this study. This study focused only on the airborne viable cancer cells released during cutting of tumors using LCS, not on clinical practice in detail. We believe that further studies, using animal models, are needed.

5. Conclusions

In conclusion, this study demonstrates that airborne viable cancer cells can be released during tumor ablation with the UAS. Moreover, it was shown that the number of cells released can be decreased by changing from the high power mode of the LCS to the low power mode and by applying suction adjacent to the excision site.

Conflicts of Interest

The authors declare that they have no conflicts of interest concerning this article. The authors received a research grant from Johnson & Johnson K.K. Medical Company.

Acknowledgments

The authors thank Dr. M. Hattori (Hiroshima University, Japan) for his assistance with the statistical analysis. This work was carried out with the kind cooperation of the Analysis Center of Life Science and Analysis Center of Life Science and the Research Facilities for Laboratory Animal Science, Natural Science Center for Basic Research and Development (N-BARD), Hiroshima University, Hiroshima, Japan.

References


