Video Article

In vivo Imaging and Therapeutic Treatments in an Orthotopic Mouse Model of Ovarian Cancer

Alexis B. Cordero1, Youngjoo Kwon1, Xiang Hua2, Andrew K. Godwin1
1Department of Medical Oncology, Women’s Cancer Program
2Transgenic Mouse Facility, Fox Chase Cancer Center

Correspondence to: Andrew K. Godwin at Andrew.Godwin@fccc.edu

URL: http://www.jove.com/index/Details.stp?ID=2125
DOI: 10.3791/2125
Citation: Cordero A.B., Kwon Y., Hua X., Godwin A.K. (2010). In vivo Imaging and Therapeutic Treatments in an Orthotopic Mouse Model of Ovarian Cancer. JoVE. 42.
DOI: 10.3791/2125
URL: http://www.jove.com/index/Details.stp?ID=2125

Abstract

Human cancer and response to therapy is better represented in orthotopic animal models. This paper describes the development of an orthotopic mouse model of ovarian cancer, treatment of cancer via oral delivery of drugs, and monitoring of tumor cell behavior in response to drug treatment in real time using in vivo imaging system. In this orthotopic model, ovarian tumor cells expressing luciferase are applied topically by injecting them directly into the mouse bursa where each ovary is enclosed. Upon injection of D-luciferin, a substrate of firefly luciferase, luciferase-expressing cells generate bioluminescence signals. This signal is detected by the in vivo imaging system and allows for a non-invasive means of monitoring tumor growth, distribution, and regression in individual animals. Drug administration via oral gavage allows for a maximum dosing volume of 10 mL/kg body weight to be delivered directly to the stomach and closely resembles delivery of drugs in clinical treatments. Therefore, techniques described here, development of an orthotopic mouse model of ovarian cancer, oral delivery of drugs, and in vivo imaging, are useful for better understanding of human ovarian cancer and treatment and will improve targeting this disease.

Protocol

I. Preparation of Ovarian Tumor Cells

1. Grow ovarian cancer cell lines expressing luciferase in culture. Sources of luciferase can be Firefly, Renilla, or other species. Cells expressing fluorescent proteins can also be used. For this demonstration we use an ovarian cancer cell line, OVCAR5, expressing firefly luciferase.
2. Harvest cells using routine cell culture technique.
3. Keep the cell suspension (10,000 cells/μL) in phosphate buffered saline (PBS) on ice until time of injection.

II. Intrabursal Injection

This procedure requires assistance from a second person. All surgical procedures are conducted under aseptic conditions. This includes wearing surgical attire and using sterile surgical instruments, syringe, and needles.

1. Prepare the anesthetic solution by mixing 3 mL of ketamine hydrochloride (100 mg/mL), 1.6 mL of xylazine hydrochloride (100 mg/mL), 1.5 mL of acepromazine (10 mg/mL), and 20 mL of 0.9% sodium chloride.
2. Check the animal’s identification number and observable health. Anesthetize the animal with prepared ketamine-xylazine-acepromazine anesthesia via intraperitoneal (i.p.) injection with a dose volume of 8–9 mL/kg body weight (BW).
3. Confirm that the animal is under an acceptable plain of anesthesia by performing a toe pinch with forceps or fingers to the animal’s hind paws. If there is pedal reflex, wait for a deeper plain of anesthesia until the animal is unresponsive to this procedure.
4. Lay the animal dorsal side up on a sterile gauze pad with its head facing away and its tail facing towards you. The point of incision is located to the left or right of the midline and above the ovaries. Shave or wet the fur with 70% alcohol to the incision point.
5. Lift the wetted skin using forceps and make a small incision with the scissors at the dorsomedial position and directly above the ovarian fat pad. The ovarian fat pad should be visible beneath the surface of the peritoneal wall. Fat pad is easily recognizable by its white color in contrast to the dark pink tissue surrounding it.
6. Gently lift the peritoneal wall lining and make a small incision as described above (5).
7. Place a sterile soaked saline gauze pad on the midline adjacent to the incision. Locate the ovarian fat pad and gently pull it out and rest it onto the gauze. Stabilize the ovary by clamping the fat pad with a bulldog clip. Under a dissecting microscope, position the ovary as to allow for the insertion of the needle (30 gauge, G) into the oviduct tubule bend leading to the bursa. When the needle is inserted into the proper position, it should be visible under the bursa.
8. Gently push the plunger of the syringe to inject 5 μL of cell suspension between the bursa and the ovary while the syringe is positioned to injection site. This step requires two people. One person pushes the plunger while the other person maintains positioning of needle. Remove the needle quickly to seal the puncture site but gently enough not to tear the bursa and tubule. The bursa should appear to be slightly distended with proper injection.
9. Release the fat pad from the bulldog clip and gently replace the reproductive tract and fat pad back into the peritoneal cavity. Gently close the body wall by pulling the upper peritoneal lining over the lower lining. Close the skin with surgical staples or wound clips.
10. Place the recovering animal back in its cage and provide a safe heat source to avoid hypothermia and speed up recovery. Monitor the breathing rate and ease, the return of muscle tone, and the ability to voluntarily move. These are all good indicators of the progression towards recovery. Staples or wound clips can be removed 7 or more days post surgery.

III. Oral Gavage Administration...
In Vivo Imaging

We use the Caliper Life Sciences to monitor the behavior of cells injected into intrabursal cavity. Experiments using this system typically have a timeframe of 4~16 weeks from the time of tumor implantation.

Representative Results

Figure 1. In Vivo imaging of ovarian tumor cells in an orthotopic mouse model. OVCAR5 cells expressing luciferase were injected intrabursally into right ovary and imaged over time. Images were taken 13 (left), 17 (middle), and 22 (right) days after the injection using the IVIS Spectrum.
imaging system. Dorsal side is shown in upper panel and ventral side is in lower panel. Note that the peritoneal spread of tumor cells 22 days following injection.

Experiments on animals were performed in accordance with the guidelines and regulations set forth by Fox Chase Cancer Center's Institutional Animal Care and Use Committee.

Disclosures

No conflicts of interest declared.

Discussion

Ovarian cancer is the leading cause of death among all gynecologic malignancies. The high mortality rate of this disease is largely due to its late diagnosis and the lack of reliable diagnostic methods. Furthermore, conventional chemotherapy often encounters chemoresistance and relapse of cancer. Therefore, novel therapeutics is required to effectively target this disease. In development of new therapy targeting human ovarian cancer, it is critical to develop a representative animal model.

An orthotopic animal model has advantages over conventional xenograft models (e.g. subcutaneous or intraperitoneal injections of tumor cells) in that 1) it reproduces the primary site of tumor formation, 2) it represents common site of metastases, and 3) it provides tumor cells to interact with appropriate microenvironment. Rodents have a unique bursal membrane that surrounds the ovary and is continuous with the oviduct. This unique anatomy of rodents allows injection of ovarian tumor cells orthotopically. Intrabursally injected ovarian tumor cells behave similar to human disease, thereby growing within intrabursal membrane and spreading to peritoneal cavity as tumor progresses. Injection of cells stably expressing luciferases or fluorescent proteins also allows tracking behavior of tumor cells in real time. Bioluminescent and/or fluorescent imaging technology makes it possible to repeatedly image tumor cells over an extended period of time and study tumor growth, distribution, and regression in non-invasive manner.

In establishing the orthotopic model, it is critical to quickly remove the needle from the bursa upon injection. Abrupt removal of the needle seals the puncture site and prevents the leakage of injected cells. However, at the same time, the movement should be gentle not to tear the bursa. If leakage occurs during injection, it can cause cells to seed in the abdomen and potentially confound the study. I.e., premature spreading outside the ovary. In this event, the specific animal should be recorded and followed for the development of multiple tumor sites in the peritoneum prior to treatment or alternatively eliminated from the subsequent analyses. Prior to performing oral gavage, it is critical to check the length of the gavage tube by measuring from the tip of the animal's head to the last rib. It is helpful to mark the tube at the animal's nose and do not pass the tube past this mark. Insertion of the tube past this mark can result in perforation to the stomach. Determining the length of the tube is especially important with younger animals or animals weighing under 20 g. Upon insertion of the gavage needle down the esophagus, there should be no resistance or struggle from the animal. It is important not to administer the solution or suspension too fast as this can lead to reflux and subsequently deliver inaccurate dose volume, as well as adding stress to the animal. For comparable imaging results from time to time, it is desirable to keep the time consistent between injection of luciferase substrate and imaging. The time lapse can be determined by taking series of images after substrate injection and observe the kinetics of signals. Development of an orthotopic mouse model of ovarian cancer, oral delivery of drugs, and in vivo imaging are necessary techniques for better understanding of development and spread of ovarian cancer and assessing novel therapeutic regimens that may ultimately improve the outcome of patient with this deadly disease.

Acknowledgements

We thank Dr. Harvey Hensley for his expert technical support. This work benefited from the use of the following Fox Chase Cancer Center (FCCC) facilities: Laboratory of Animal Facility and Small Animal Imaging Facility. This work was supported in part by the Ovarian Cancer SPORE at FCCC/UPenn (P50 CA083638), the FCCC core grant (P30 CA006927), and the Ovarian Cancer Research Fund.

References