Video Article Improved Visualization of Lung Metastases at Single Cell Resolution in Mice by Combined *In-situ* **Perfusion of Lung Tissue and X-Gal Staining of** *lacZ***-Tagged Tumor Cells**

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Abstract

Metastasis is the main cause of death in the majority of cancer types and consequently a main focus in cancer research. However, the detection of micrometastases by radiologic imaging and the success in their therapeutic eradication remain limited.

While animal models have proven to be invaluable tools for cancer research¹, the monitoring/visualization of micrometastases remains a challenge and inaccurate evaluation of metastatic spread in preclinical studies potentially leads to disappointing results in clinical trials². Consequently, there is great interest in refining the methods to finally allow reproducible and reliable detection of metastases down to the single cell level in normal tissue. The main focus therefore is on techniques, which allow the detection of tumor cells *in vivo*, like micro-computer
tomography (micro-CT), positron emission tomography (PET), bioluminescence or techniques for *in vivo* monitoring of primary tumor growth and metastasis in different osteosarcoma models. Some of these techniques can also be used for *ex vivo* analysis of metastasis beside classical methods like qPCR⁵, FACS⁶ or different types of histological staining. As a benchmark, we have established in the present study the stable transfection or transduction of tumor cells with the *lacZ* gene encoding the bacterial enzyme β-galactosidase that metabolizes the chromogenic substrate 5-bromo-4-chloro-3-indolyl-beta-D-galactopyranoside (X-Gal) to an insoluble indigo blue dye⁷ and allows highly sensitive and selective histochemical blue staining of tumor cells in mouse tissue ex vivo down to the single cell level as shown here. This is a low-cost and not equipment-intensive tool, which allows precise validation of metastasis⁸ in studies assessing new anticancer therapies⁹⁻¹¹. A limiting factor of X-gal staining is the low contrast to *e.g.* blood-related red staining of well vascularized tissues. In lung tissue this problem can be solved by *in-situ* lung perfusion, a technique that was recently established by Borsig *et al.*12 who perfused the lungs of mice under anesthesia to clear them from blood and to fix and embed them *in-situ* under inflation through the trachea. This method prevents also the collapse of the lung and thereby maintains the morphology of functional lung alveoli, which improves the quality of the tissue for histological analysis. In the present study, we describe a new protocol, which takes advantage of a combination of X-gal staining of *lacZ-*expressing tumor cells and *in-situ* perfusion and fixation of lung tissue. This refined protocol allows high-sensitivity detection of single metastatic cells in the lung and enabled us in a recent study to detect "dormant" lung micrometastases in a mouse model¹³, which was originally described to be non-metastatic¹⁴.

Video Link

The video component of this article can be found at <http://www.jove.com/video/4162/>

Protocol

1. *In-situ* **Lung Perfusion and Fixation**

- 1. Anesthetize mice by i.p. injection of ~150 μl phosphate buffered saline (PBS) containing 112.5 mg/kg (body weight) ketamine, 16.5 mg/kg xylazine, and 15 mg/kg acepromazine (or by another anesthesia containing appropriate pain killers).
- 2. When reflexes of the mouse are no longer observed, fix it on the operating table back-down and wet the fur with 70% Ethanol to slick the hairs down.
- 3. Open the skin from the abdomen up to the neck and pull it to the sides or remove it.
- 4. Open the peritoneum and the thorax to an ample size. Prevent ruptures of big vessels (*e.g.* the vena jugularis) to avoid reduced efficiency of the subsequent perfusion.
- 5. Cut the vena cava caudalis underneath the liver.
- 6. Use a 20 ml syringe equipped with a 21G 24G needle to inject slowly 10-15 ml PBS into the right ventricle of the beating heart until the lung is completely white and the heart stops beating (asystole).
- 7. Pinch off the blood vessels above the liver and the diaphragm with a vascular clamp.
- 8. Use a 10 ml syringe equipped with a 21G 24G needle to inject ~2 ml of 3% paraformaldehyde (PFA) in PBS into the right ventricle.

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- 9. Uncover the trachea outside of the thorax. Use the same syringe to inject ~3 ml of 3% PFA in PBS cranial (outside of thorax) into the trachea until the lung is inflated. Immediately after removal of the needle, pinch off the trachea caudal of the puncture with a vascular clamp and wait for 10 min to allow fixation.
- 10. Transect the blood vessels above the vascular clamp, remove the clamp and inject ~2 ml PBS into the right ventricle to remove excess PFA.
- 11. Puncture the lower parts of all lung lobes with a needle, remove the vascular clamp at the trachea and inject 3-5 ml of a solution of PolyFreeze embedding medium/PBS 1:1 into the trachea caudal of the first puncture until the PFA in the lung lobes is replaced by the solution.
- 12. Remove the lung carefully by pulling the heart with a forceps and by transecting connective tissue ligaments, blood vessels and the trachea. Keep the heart connected to the lungs.
- 13. Depending on how to proceed with the lung tissue analysis continue with the protocols outlined in sections 2 or 3. If you want to do both analyses, remove one lung lobe and proceed as described under 3. Process the remaining lobes connected to the heart as described under 2.

2. Visualization of *lacZ***-tagged Metastatic Tumor Cells in Entire Lung Lobes**

- 1. Place the lung in a plastic cup with tight closing lid and fix it with 2% formaldehyde in PBS at room temperature for 30-60 min.
- 2. Remove fixation solution and wash thoroughly 3 times with PBS.
- 3. Add at least 10 ml freshly prepared 5-Bromo-4-chloro-3-indolyl-β-D-galactoside (X-Gal) staining solution (X-Gal stock solution [40 mg/ml in Dimethylformamide] 1:40 diluted in Basic Staining Solution; pH 7.1 (**Table 1**); protect from light).
- 4. Put a piece of gauze on top of the swimming lung to keep it completely in the solution and place the lid only loosely on the pot to allow exchange of air.
- 5. Incubate at 37 °C for 3-5 hr protected from light.
- 6. Remove X-Gal solution, rinse once with PBS to remove residual X-Gal solution (optional) and add 4% PFA.

3. Visualization of *lacZ***-tagged Metastatic Tumor Cells in Lung Cryosections**

- 1. Prefill a labeled embedding mold to 1/3 with undiluted PolyFreeze embedding medium. Put the lung lobe on top and fill up with embedding medium until the lobe is completely covered. Try to prevent bubbles.
- 2. Incubate the embedded lungs at 4 °C for 20 min.
- 3. Then freeze the embedded lungs slowly in a mixture of dry ice and Isopentane and store at -80 °C.
- 4. Cut 7-10 μm cryosections on a cryostat and mount them on SuperFrostPlus microscope slides.
- 5. Incubate the sections immediately with X-Gal staining solution at 37 °C for 24 hr in a humidified chamber.
- 6. Rinse the slides in PBS 2 times for 5 min and then briefly in distilled water.
- 7. Counter-stain with nuclear fast red for 10-30 sec, rinse in distilled water and mount the slides with Immu-Mount.

4. Representative Results

Asai *et al.* reported in the original article on the Dunn-derived LM8 OS mouse model that s.c. primary tumors derived from the parental Dunn cells, different from those derived from the highly metastatic LM8 sub cell line, do not spontaneously form detectable lung metastases in syngeneic C3H mice¹⁴. With the here reported techniques, we reinvestigated the formation of metastases in the Dunn/LM8 mouse OS models. We took advantage of stable *lacZ*-transduced Dunn and LM8 cells and of a protocol for *in-situ* lung perfusion and fixation in mice. Representative images of perfused and non-perfused lungs of mice subcutaneously injected with *lacZ*-transduced and non-transduced control Dunn and LM8 cells are shown in **Figure 1**. In mice injected with control Dunn cells, macroscopic and microscopic metastases remained undetectable in non-perfused and perfused lungs (**Figure 1A**, *i-iv*). But, interestingly, in mice injected with Dunn-*lacZ* cells, X-gal staining revealed blue micrometastatic foci of single cells or small cell clusters (< 0.1 mm) on the surface of non-perfused lungs (**Figure 1A**, *vi*). *In-situ* perfusion and fixation of the lungs further improved the detectability of Dunn-*lacZ* micrometastases (**Figure 1A**, *viii*). However, outgrowth to macroscopic foci was not observed (**Figure 1A**, *v*, *vii*).

In mice injected with control LM8 cells, translucent, barely detectable macrometastatic foci larger than 0.1 mm in diameter were recognized in non-perfused lungs (**Figure 1B**, *i*). Perfusion of the lung (**Figure 1B**, *iii*) did not improve the detection of the foci. However, in mice injected with LM8-*lacZ* cells multiple X-Gal stained blue macro- (**Figure 1B**, *v*) and micrometastases (**Figure 1B**, *vi*) were detected on the surface of non-perfused organs. Moreover, perfusion of the lungs further improved the detectability of macro- and micrometastases (**Figure 1B**, *vii*-*viii*). Consequently, micro- and macrometastases became visible at a higher density and a larger number, mainly due to the translucency of the perfused tissue in which foci underneath the organ surface became also visible.

An additional histological analysis using cryosections of lung tissue confirmed the improved detectability of lung metastases in mice injected with *lacZ*-transduced Dunn and LM8 cells. In mice with primary tumors derived from Dunn-*lacZ* or LM8-*lacZ* cells, unlike in mice with primary tumors of the respective control cells, micrometastases or even single cell foci were recognized in lung sections (**Figure 2**). Moreover, macrometastases were also more clearly visible in mice injected with LM8-*lacZ* cells than in animals injected with the control LM8 cells (**Figure 2C, D**).

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Figure 1. Detection of spontaneous metastases of non-transduced (i-iv) and *lacZ*-transduced (v-viii) Dunn (A) and LM8 (B) cells in nonperfused (i, ii, v, vi) and perfused (iii, iv, vii, viii) lungs in C3H mice. Metastases in representative X-Gal-stained whole lungs (v, vii in A and B) and respective close-ups (vi, viii in A and B) appear blue. Macrometastatic foci (> 0.1 mm) in the organs of mice injected with non-tagged LM8 cells (i-iv in B) are indicated by asterisks. Areas shown in close-ups are representative for entire lungs. Scale bars indicate 1 mm in images of whole lungs and 0.1 mm in close-ups. [Click here to view larger figure](http://www.jove.com/files/ftp_upload/4162/4162fig1large.jpg).

Figure 2. Detection of Dunn and LM8 micrometastases in lung cryosections. Sections of O.C.T. embedded lung tissue were incubated in X-Gal staining solution at 37 °C for 24 hr in a humidified chamber and then counterstained with nuclear fast red. Arrows point to micrometastatic foci recognized in lung sections of mice injected with Dunn-*lacZ* (B) or LM8-*lacZ* cells (D). Micrometastases remained undetectable in lung sections of mice injected with non-transduced Dunn (A) or LM8 cells (C). Scale bars indicate 0.1 mm.

Discussion

The here presented results in the Dunn/LM8 mouse OS model demonstrate the power of the newly established method that combines the X-Gal staining of *lacZ*-tagged tumor cells with the *in-situ* perfusion/fixation of lung tissue. This combination of the two techniques allows high sensitivity detection of micrometastatic lesions down to the single cell level and also improves the visualization of macrometastases on the lung surface (**Figure 1**) as well as in lung sections (**Figure 2**). While the X-Gal staining also allows the detection of (micro)metastases in other organs, *in-situ* perfusion/fixation improves the detectability of metastatic foci in tissues other than the lung only slightly because of the natural blood- and tissue-
related color of these organs¹³. Even if the perfusion is directed t the contrast between X-Gal staining and the natural color of the organ. However, the method is applicable to any type of *lacZ*-tagged tumor cells, will dramatically improve the detectability of lung metastasis down to the level of dormant single-cell micrometastases and enables an easy and reliable quantification of macro- and micrometastases. A limitation of this method and all other techniques that are based on reporter genes, including luciferase and fluorescent proteins, is the stability of the transgene expression. As shown in **Figure 2d**, not all tumor cells within the macrometastatic foci are stained blue, indicating a lack of beta-galactosidase activity. This might be related to necrosis, but it is more likely due to a loss of transgene expression. We observed that the Dunn and LM8 cells are very effective in the down-regulation of *lacZ* and other transgenes even under continuous selection for expression. We therefore switched in recent studies to the murine K12 and K7M2 and the human HOS, 143B and SaOS-2 osteosarcoma cell lines, which all maintained stable *lacZ* expression *in vitro* as well as *in vivo* up to 100% over time.

Once stable *lacZ*-expression is guaranteed, this technique can be applied in studies with gene-manipulated tumor cells, *e.g.* to mechanistically investigate the process of tissue colonization¹³ as well as for the development and testing of new therapies aiming at eradication of metastatic $lesions¹$ 1,16 . Furthermore, it can serve as a benchmark for the improvement of current radiological imaging techniques, such as PET, (micro)CT and MRI, used for early detection of metastatic lesions. In a recent PET study (unpublished) with different tracers we verified the *in vivo* detected lung metastases subsequently *ex vivo* with the described protocol. In an ongoing study with a new small animal micro-CT (SkyScan) we are so far able to detect lung metastases *in vivo* down to a size of 0.5 mm and *ex vivo* down to 0.3 mm, but we aim at a resolution of 0.1 mm. Interestingly, this is the size limit that we set to distinguish macro- from micrometastases with the combined method of *in-situ* perfusion and X-Gal staining. This underlines again the sensitivity and usefulness of this easy and cost-effective technique.

Disclosures

No conflicts of interest declared.

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