Abstract

Self-renewing cancer cells are the only cell types within a tumor that have an unlimited ability to promote tumor growth, and are thus known as tumor-propagating cells, or tumor-initiating cells. It is thought that targeting these self-renewing cells for destruction will block tumor progression and stop relapse, greatly improving patient prognosis. The most common way to determine the frequency of self-renewing cells within a tumor is a limiting dilution cell transplantation assay, in which tumor cells are transplanted into recipient animals at increasing doses; the proportion of animals that develop tumors is used to calculate the number of self-renewing cells within the original tumor sample. Ideally, a large number of animals would be used in each limiting dilution experiment to accurately determine the frequency of tumor-propagating cells. However, large scale experiments involving mice are costly, and most limiting dilution assays use only 10-15 mice per experiment.

Zebrafish have gained prominence as a cancer model, in large part due to their ease of genetic manipulation and the economy by which large scale experiments can be performed. Additionally, the cancer types modeled in zebrafish have been found to closely mimic their counterpart human disease. While it is possible to transplant tumor cells from one fish to another by sub-lethal irradiation of recipient animals, the regeneration of the immune system after 21 days often causes tumor regression. The recent creation of syngeneic zebrafish has greatly facilitated tumor transplantation studies. Because these animals are genetically identical, transplanted tumor cells engraft robustly into recipient fish, and tumor growth can be monitored over long periods of time. Syngeneic zebrafish are ideal for limiting dilution transplantation assays in that tumor cells do not have to adapt to growth in a foreign microenvironment, which may underestimate self-renewing cell frequency. Additionally, one-cell transplants have been successfully completed using syngeneic zebrafish and several hundred animals can be easily and economically transplanted at one time, both of which serve to provide a more accurate estimate of self-renewing cell frequency.

Here, a method is presented for creating primary, fluorescently-labeled T-cell acute lymphoblastic leukemia (T-ALL) in syngeneic zebrafish, and transplanting these tumors at limiting dilution into adult fish to determine self-renewing cell frequency. While leukemia is provided as an example, this protocol is suitable for determining the frequency of tumor-propagating cells using any cancer model in the zebrafish.

Video Link

The video component of this article can be found at https://www.jove.com/video/2790/

Protocol

1. DNA microinjection of zebrafish embryos

1. Linearize rag2::cMyc and rag2::GFP DNA constructs by digesting 10μg of each plasmid with NotI at 37°C overnight.
2. Phenol:chloroform extract and precipitate the plasmids, and resuspend each in 20μL H₂O.
3. Determine the concentration of the DNA by running a 1:1, 1:5 and 1:10 dilution of each digested plasmid on a 1% agarose gel with 20μL, 10μL and 5μL of a high-range DNA ladder. This type of quantification is generally more accurate than a spectrometer reading.
4. Dilute the DNA to a final concentration of 60ng/μL in 1M KCl+0.5X TE for injection into CG1-strain (or other syngeneic strain) zebrafish embryos using 30ng/μL rag2::cMyc + 30ng/μL rag2::GFP.
5. Injections should be performed as previously demonstrated. Briefly, male and female zebrafish are set up in mating chambers the night before injection. On the day of injection, DNA is loaded into the injection needle, the pressure is adjusted so that a bubble with a 50μm diameter is expelled (30pg DNA), which is measured by a stage micrometer. Then, embryos are injected at the one-cell stage, with the DNA being injected directly into the cell, and not into the yolk. Survival of the injected CG1 embryos and larvae is generally poor, and only 5% of fish will successfully incorporate the transgene; thus, >600 embryos should be injected to ensure that some fish will develop leukemia.
6. Store the injected embryos at 28.5°C, and remove the dead embryos after 24hr. Animals are then transferred to large tanks at 5 days of life and raised as previously described.
2. Screening for primary T-ALL in zebrafish larvae

1. Approximately 28 days after injection, anesthetize larval zebrafish by adding 200μL of 4ng/mL Tricane-S to 25mL of fish system water. Examine the larvae for development of fluorescently-labeled T-ALL by using an epifluorescence microscope at 20X magnification, using a 485/20 excitation wavelength and 530/25 emission filter to detect GFP fluorescence (Figure 1).

2. Separate tumor positive larvae from those that are negative. Negative larvae can be examined at a later time point, once tumors may have grown larger, to ensure that no T-ALL positive fish were missed.

3. Hold an adult (>60 days old) syngeneic zebrafish ventral side up, and inject the cell suspension into the peritoneal cavity (Figure 3C). Data from limiting dilution cell transplantation assays from three different primary T-ALLs are shown in Figure 3D. For these experiments, over 220 animals were transplanted, which can be easily accomplished by one person within several hours. DNA microinjection into embryos from syngeneic fish often results in a low survival rate of the injected embryos, with <60% of embryos surviving for 24 hours. Additionally, survival of the injected syngeneic fish is generally poor during larval development, with often <20% surviving for 28 days. It is therefore important to inject a large number of embryos in order to create transgenic fish that will develop T-ALL.

4. Pellet the cells by centrifuging the 96-well plate at 2000xg for 10 minutes.

3. Isolation and purification of fluorescently-labeled primary leukemia cells

1. When GFP-positive leukemia cells have overtaken >50% of the animal, sacrifice the fish by adding 1mL of 4ng/mL Tricane-S in a petri dish containing 9mL fish system water. Approximate 28 days after transplant, examine the transplanted fish with an epifluorescence microscope using a 485/20 excitation wavelength and 530/25 emission filter to detect GFP fluorescence (Figure 1).

2. Place the fish in a new petri dish containing 1mL 0.9X PBS with 5% Fetal Bovine Serum to buffer the cells. Macerate the fish with a razor blade, pipet the mixture to dissociate large cell clumps, then pass the cells through a 40μm mesh strainer into a 50mL tube. It is not necessary to dissociate all tissue clumps into single cells.

3. Pipet 500μL of the cells to a 4mL polystyrene tube. Add 1μL 1mg/mL propidium iodide (PI), which will label dead cells. Vortex, then keep the cells on ice.

4. Sacrifice a wild-type CG1 fish, macerate and strain in the same way as above to collect normal cells, which serve as a carrier for the tumor cells during transplantation. Add 4mL 5% FBS in 0.9X PBS to the 50mL tube for a total volume of 5mL.

5. Count the normal cells, dilute to 3x10^5 cells/mL in 5% FBS in 0.9X PBS, then pipet 100μL of the 96-well plate of a 96-well plate.

6. Using a Fluorescence Activated Cell Sorter (FACS), sort GFP positive, PI negative tumor cells (Figure 2A,B) into the 96-well plate containing blood cells at the following doses: 10 cells/well into 48 wells, 100 cells/well into 24 wells, 1000 cells/well into 12 wells, and 10,000 cells/well into 6 wells. Additionally, 10,000 cells should be sorted into a well without normal blood cells, and reanalyzed using FACS to assess the purity and viability of the sorted cells (Figure 2C).

4. Limiting dilution transplantation into adult zebrafish

1. Approximately 28 days after injection, examine the transplanted fish with an epifluorescence microscope using a 485/20 excitation wavelength and 530/25 emission filter to detect GFP fluorescence. Record the number of positive fish per total number of fish injected.

2. Re-examine the fish every 14 days for up to 90 days and record the number of positive fish. When a tumor-positive fish become moribund, sacrifice the animal by Tricane-S overdose.

3. Separate tumor positive larvae from those that are negative. Negative larvae can be examined at a later time point, once tumors may have grown larger, to ensure that no T-ALL positive fish were missed.

5. Analysis to determine leukemia-initiating cell frequency

1. Approximately 28 days after transplant, examine the transplanted fish with an epifluorescence microscope using a 485/20 excitation wavelength and 530/25 emission filter to detect GFP fluorescence. Record the number of positive fish per total number of fish injected.

2. Re-examine the fish every 14 days for up to 90 days and record the number of positive fish. When a tumor-positive fish become moribund, sacrifice the animal by Tricane-S overdose.

3. Input the results into a table, as shown in Figure 3D. Upload the data into the web-based ELDA (Extreme Limiting Dilution Analysis) statistical software at http://bioinf.wehi.edu.au/software/elda/index.html, which uses the frequency of tumor positive and negative animals at each transplant dose to determine the frequency of self-renewing leukemia cells within the primary T-ALL.

4. Monitor T-ALL positive fish at least once at week using the epifluorescence microscope to track tumor growth.

6. Representative Results:

DNA microinjection into embryos from syngeneic fish often results in a low survival rate of the injected embryos, with <60% of embryos surviving for 24 hours. Additionally, survival of the injected syngeneic fish is generally poor during larval development, with often <20% surviving for 28 days. It is therefore important to inject a large number of embryos in order to create transgenic fish that will develop T-ALL.

As an example, CG1 strain embryos were injected with rag2::cMy+rag2::GFP. The transgenes co-segregate in the developing embryo, so that every cell that expresses GFP will also express cMy. Larvae were screened for primary T-ALL after 28 days (Figure 1). Previous work has shown that approximately 5% of injected fish will have GFP-positive T-cells within their thymus (Figure 1), and 100% of these fish will develop leukemia as the T-cells become transformed. While a small proportion of zebrafish may have a more advanced leukemia that is very easy to detect at day 28, most will only have a GFP-positive thymus (Figure 1), so larvae should be examined individually under high magnification to ensure that all positive fish are selected. GFP-positive T-ALL cells will overtake >50% of the animal between days 45-70.

In the example provided, zebrafish were sacrificed at 65 days of life, and leukemia cells were isolated and FACS sorted for limiting dilution transplantation. Single cells that were propidium iodide negative and GFP-positive (Figure 2) were sorted into a 96-well plate. Reanalysis was done on 10,000 sorted cells in order to assess viability (ideally >95%) and purity (ideally >92%, Figure 2). Transplanted T-ALLs generally arise before 28 days, but some tumors may take more than 60 days to develop. In this example, at day 28, early stage tumors were seen as an area of GFP-positive cells near the injection site (Figure 3B), while some T-ALLs were more progressed, having expanded to fill the peritoneal cavity (Figure 3C). Data from limiting dilution cell transplantation assays from three different primary T-ALLs are shown in Figure 3D. For these experiments, over 220 animals were transplanted, which can be easily accomplished by one person within several hours. Data analysis using ELDA software showed that, on average, 1 in 135 T-ALL cells were self-renewing leukemia-propagating cells (Figure 3E).
Figure 1. Zebrafish larvae injected at the one-cell stage with \textit{rag}:-\textit{cMyc}+\textit{rag}:-\textit{eGFP} were screened for primary leukemia growth 28 days post-injection. Fish (*) had GFP-positive T-cells within the thymus; this fish will develop T-ALL as the T-cells become transformed over time. This stage is very common at day 28. One fish (**) had an advanced T-ALL that has spread into the soft tissue. The remaining two fish are negative for tumor growth. The image was taken at 16X magnification.

Figure 2. Fluorescently-labeled T-ALL cells were sorted from diseased animals and used in the limiting dilution cell transplantation assay. First, a gate was drawn to select single cells (upper left panel), then propidium iodide negative cells are selected (middle left panel). Finally, a gate was drawn for the GFP-positive cells (lower left panel).
drawn to select only the GFP-positive leukemia cells for sorting (lower left panel). Sorted cells should be reanalyzed to assess viability and purity before transplant (right panels).

**Figure 3.** Zebrafish were examined for leukemia growth 28 days after transplant. Fish are either tumor negative (A), have a small tumor growing at the injection site (B), or have a progressed leukemia (C). The images were taken at 7X magnification. The total number of leukemia-positive fish per total number of fish transplanted at each dilution is recorded, as in (D). The data are input into the web-based ELDA program to calculate the number of self-renewing leukemia cells and the upper and lower 95% confidence intervals (E).

**Discussion**

A major strength of using zebrafish in cancer research is that large numbers of animals can be used at relatively low cost. This is especially important in limiting dilution cell transplantation assays, where the proportions of transplanted animals that develop tumors to the total number transplanted are used to determine tumor-initiating cell frequency. In the method presented here, over 70 transplant recipient animals are used per assay, providing an accurate estimate of tumor-propagating cell number. Both the number of animals used and the doses of tumor cells transplanted should be optimized for a given cancer model; for example, if initial experiments show tumor-initiating cells are rare, useful transplant doses may be 100,000, 50,000, 10,000 and 1,000 cells per transplant.

Syngeneic zebrafish strains are useful in limiting dilution analysis. However, these strains are not yet commonly used in all laboratories, and some zebrafish cancer models exist only in allogenic fish. Limiting dilution cell transplantations can be performed in wild-type strains that have undergone immune ablation, although the self-renewing cell frequency may be calculated as 10-100 fold higher than if syngeneic zebrafish were used. It is important to note that greater doses of cells should be used for transplants into non-immune matched, irradiated recipients, as some tumor cells will not survive transplant into a foreign microenvironment. Additionally, many tumors will begin to regress after 21 days when the recipient animal's immune system recovers, so animals should be assessed for tumor growth every 7 days after transplant.

The methods presented here are suitable for use with any zebrafish cancer model, and have thus far been used to determine tumor-initiating cell frequency in both T-cell acute lymphoblastic leukemia, and a solid tumor model, rhabdomyosarcoma. These tumors were fluorescently labeled by co-injection of embryos with both an oncogene and fluorescent marker; because the DNA co-segregates, any cell expressing the oncogene will also express the fluorescent protein. While fluorescence is recommended because it facilitates tracking of tumor growth without the need to sacrifice the animal and provides a straightforward way to isolate tumor cells by FACS, it is possible to label tumor cells with antibodies to tumor-specific makers to facilitate their purification, though antibody staining in zebrafish may require optimization.

Finally, once the incidence of tumor-propagating cells has been determined for a given tumor type, it is possible to use these assays to identify the mechanisms that govern their cell frequency. For example, primary tumor cells can be treated with pathway specific inhibitors before limiting dilution transplant, or transplanted fish themselves may be dosed with drugs. Additionally, the genetics of the tumors themselves may be manipulated by injecting the one-cell stage fish with a gene of interest, so that the resulting primary tumors will over-express the gene. In these experiments, any effect on self-renewing cell frequency will be observed in the limiting dilution assays.
Disclosures

No conflicts of interest declared.

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References