Chapter 3

Visualization of Adult Stem Cells Within Their Niches Using the Drosophila Germline as a Model System

Annekatrin König and Halyna R. Shcherbata

Abstract

The germaria of the fruit fly Drosophila melanogaster present an excellent model to study germline stem cell–niche interactions. Two to three adult stem cells are surrounded by a number of somatic cells that form the niche. Here we describe how Drosophilae germaria can be dissected and specifically immuno-stained to allow for identification and analysis of both the adult stem cells and their somatic niche cells.

Key words Drosophila, Germarium, Ovary, Adult stem cells, Stem cell niche, Germline, Ovarian soma, Immunostaining

1 Introduction

Adult stem cells usually reside in the stem cell niche, a unique physiological microenvironment that helps stem cells to carry on self-renewing divisions throughout the lifetime of an organism. The niche includes cellular and noncellular elements that can be divided into one of the two main mechanistic types—physical contacts and diffusible factors [1]. Close contacts include tight junctions, adherens junctions, gap junctions, the Notch signaling pathway, the basement membrane, and extracellular matrix proteins. Diffusible factors, which are secreted by niche cells and travel over varying distances to keep stem cell identity, often affect transcription. Stem cells must be anchored to the niche through cell–cell interactions so that they will stay both close to niche factors that specify self-renewal and far from differentiation stimuli.

Presently the existence of a stem cell niche has been demonstrated for mammalian adult stem cells in the hematopoietic, epidermal, neural, and intestinal systems. However, the stem cell niches involved in maintenance of adult mammalian tissues and particularly their role in cancer development remain complex, poorly defined, and difficult to study in vivo [2].
The *Drosophila* ovarian stem cell niche is very well characterized and has been used for many years to unravel the complex stem cell–niche interactions. The insights gained from these studies led to a better understanding of how stem cells work: in addition to cell–cell interactions [3] between stem and niche cells, a variety of signaling pathways involved in stem cell control were described [4–13].

The easily identifiable and analyzable cells in the *Drosophila* germline niches and the sophisticated genetic tools that are available in *Drosophila* make it an ideal system for studying stem cell–niche interactions [14]. The paired ovaries of the adult female fly each consist of 16–20 ovarioles that contain developing egg chambers. Located at the anterior of every ovariole is the germarium, where two to three stem cells are held by 5–7 cap cells and are in contact with other somatic cells (see Fig. 1b). By asymmetric division, the adult stem cells give rise to both new stem cells and differentiated cells that will become the egg. The differentiated germline cells are surrounded by somatic escort cells that are another important component of the stem cell niche [4, 15]. More posteriorly, follicle cells that are generated by specific stem cells encapsulate the differentiating germline [16]. The individual ovarioles are held together by the
terminal filaments. A variety of different markers makes it possible to nicely immunostain and analyze number, localization, shape, and interactions of the individual cells (see Fig. 2b). In addition, the development of the stem cell niche itself can be directly observed in developing larvae and early pupae where the cap cells divide and terminally differentiate (see Figs. 1a and 2b) [17].

In this chapter we show how to visualize adult stem cells in their niches in adult female Drosophila.

## 2 Materials

### 2.1 Fly Husbandry

1. Standard cornmeal agar food (recipes can be found at [http://fly.bio.indiana.edu/](http://fly.bio.indiana.edu/)).
2. Yeast paste: Dry yeast should be mixed in 5% propionic acid (see Note 1).

### 2.2 Ovary Dissection

1. Ice block for immobilization of the flies.
2. Sharp tweezers.
4. Stereomicroscope for dissection.
5. Pasteur pipettes.
## 2.3 Fixation and Washing

1. Fixing solution: 4 % formaldehyde in phosphate-buffered saline (PBS) \((see \text{ Note 2})\).

2. Nutator.

3. PBT: 0.2 % Triton X in PBS.

## 2.4 Antibodies

1. Blocking solution: 0.2 % bovine serum albumin, 5 % normal goat serum in PBT \((see \text{ Note 3})\).

2. Primary antibodies: Many antibodies are available to study germline–niche interactions; some of these are listed in Table 1. Dilute primary antibodies in blocking solution and store at 4 °C \((see \text{ Notes 4–8})\).

### Table 1

A subset of antibodies that are useful to study germline stem cell niche interactions is shown

<table>
<thead>
<tr>
<th>Protein recognized</th>
<th>Name of the antibody</th>
<th>Raised in</th>
<th>Antibody source</th>
<th>Used to mark in the germarium</th>
<th>Dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Armadillo</td>
<td>N2 7A1</td>
<td>Mouse, IgG2a</td>
<td>DSHB Iowa</td>
<td>Cell-Cell adhesion between cap cells and between stem cells and cap cells</td>
<td>1:50</td>
</tr>
<tr>
<td>Bag of marbles (BAM)</td>
<td>BamC (cytoplasmic)</td>
<td>Mouse, Rat</td>
<td>D. McKearin</td>
<td>Differentiating germline cysts, not in germline stem cells</td>
<td>1:1,000</td>
</tr>
<tr>
<td>E-Cadherin, extracellular domain</td>
<td>5D3</td>
<td>Mouse, IgG2b</td>
<td>DSHB Iowa</td>
<td>Cell-Cell adhesion between cap cells and between stem cells and cap cells</td>
<td>1:50</td>
</tr>
<tr>
<td>Engrailed</td>
<td>4D9</td>
<td>Mouse, IgG1</td>
<td>DSHB Iowa</td>
<td>Cap cells</td>
<td>1:50</td>
</tr>
<tr>
<td>Held out wings (HOW)</td>
<td>HOW</td>
<td>Rabbit, Rat</td>
<td>T. Volk</td>
<td>Germline stem cells, cystoblasts</td>
<td>1:1,000</td>
</tr>
<tr>
<td>Hts/Adducin-like</td>
<td>1B1</td>
<td>Mouse, IgG1</td>
<td>DSHB Iowa</td>
<td>Spectrosomes and fusomes</td>
<td>1:50</td>
</tr>
<tr>
<td>Lamin C</td>
<td>LC28.26</td>
<td>Mouse, IgG1</td>
<td>DSHB Iowa</td>
<td>Cap cells</td>
<td>1:50</td>
</tr>
<tr>
<td>Phosphorylated Mothers against Dpp (pMAD)</td>
<td>pMad</td>
<td>Rabbit</td>
<td>E. Laufer</td>
<td>Germline stem cells</td>
<td>1:5,000</td>
</tr>
<tr>
<td>Traffic jam</td>
<td>TJ</td>
<td>Guinea pig</td>
<td>D. Godt</td>
<td>Escort cells, cap cells</td>
<td>1:3,000</td>
</tr>
<tr>
<td>Vasa</td>
<td></td>
<td>Rat</td>
<td>P. Lasko</td>
<td>Germline cells</td>
<td>1:1,000</td>
</tr>
</tbody>
</table>

Scientists who generated a particular antibody are named as source. Their addresses are available from flybase (http://flybase.org/). DSHB Developmental Studies Hybridoma Bank at the University of Iowa.
3. Secondary antibodies: Conjugated Alexa fluor goat anti-mouse, goat anti-rabbit, or goat anti-rat from Molecular Probes, diluted 1:500 in blocking solution (Molecular Probes); store at 4 °C (see Notes 8 and 9). The secondary antibodies have to be chosen with respect to the laser lines of the available microscope.

2.5 DNA Staining and Mounting
1. DAPI solution: Make a 100× DAPI solution (1 mg/ml) and store aliquots at −20 °C. For staining, dilute in PBS (see Note 10).
2. Glycerol: 70 % Glycerol, 3 % n-propyl gallate (NPG) (see Note 11).
3. Tungsten needles.

2.6 Analysis
Laser scanning confocal microscope is used for analysis.

3 Methods

3.1 Dissection
All steps are carried out at room temperature unless otherwise stated. During all incubations and washes, the Eppendorf tubes are placed on a nutator.

3.1.1 Adult Ovaries
1. Immobilize 5–10 female flies by putting them on an ice block.
2. The ovaries are positioned in the abdomen of the fly and are simple to find in well-fed individuals (see Note 12). Dissect the flies in 1× PBS using a stereomicroscope, and hold the fly with one pair of tweezers at the thorax. Carefully open the cuticle at the posterior end of the animal with another pair of tweezers. If necessary, gently push the abdomen to squeeze out the paired ovaries. Remove all remnants of guts and cuticle and place the ovaries in an Eppendorf tube using Pasteur pipettes (see Note 13).

3.1.2 Larval Ovaries
1. Pick up late third instar larvae from the wall of the food vial or bottle.
2. Select a female larva and hold with a pair of tweezers at the anterior end.
3. The larval ovaries are located in the fat body. Cut off the larval head and hold the posterior end of the remaining larval body with one pair of tweezers. Carefully now invert the larvae by pulling it over the tweezers with another pair of tweezers. Remove cuticle and guts and transfer the fat body into an Eppendorf tube or a 24-well plate (see Note 14).
3.1.3  Fixation

1. Add fixing solution and incubate for 10 min. Remove the fixing solution carefully and wear protective gloves when handling the fixative.

2. Wash the ovaries three times for 15 min each with PBT (see Notes 15 and 16).

3.2  Antibody Staining

1. Add blocking solution and incubate for 1 h.

2. Remove the blocking solution and add primary antibody solution. Incubate overnight at 4 °C.

3. Remove the antibody solution (see Note 5) and wash the ovaries three times for 15 min each with PBT.

4. Block again in blocking solution for 1 h.

5. Incubate the ovaries in secondary antibody solution overnight at 4 °C or for 3 h at room temperature.

6. Remove the secondary antibody solution and wash with PBT twice for 15 min. Add DAPI solution and incubate for 10 min.

7. Remove the DAPI solution and wash three times for 15 min with PBT.

8. Remove as much PBT as possible and add a few drops of glycerol to the ovaries.

   (a) Adult ovaries: Place the ovaries on a slide and use tweezers and tungsten needles to separate the individual ovaries and to remove the mature eggs.

   (b) Larval ovaries: Place the fat bodies on a slide and locate the larval ovaries. Carefully remove remnants of the fat body.

9. Place a coverslip on top of the samples and analyze using a confocal microscope.

4  Notes

1. The yeast paste should have a “peanut butter-like” texture. The propionic acid helps to avoid fungal or bacterial contamination.

2. Prepare the solution fresh from a 16 % stock solution at room temperature.

3. Goat serum is used in the blocking solution if the secondary antibody was produced in goat. If you have to use secondary antibodies that were generated in another animal, use other serums from the appropriate animal.

4. Primary antibodies: Apart from several monoclonal antibodies, that are available from the Developmental Studies Hybridoma Bank, a variety of different polyclonal rabbit, goat, sheep, and
guinea pig antibodies have been made by different labs (see Table 1). However, make sure not to use goat serum in the blocking solution if the primary antibody is goat derived.

5. When analyzing GFP-marked clonal cells, you may use an anti-GFP antibody to better visualize the GFP.

6. Stability of primary antibodies: Some primary antibodies can be reused a couple of times, whereas others can be used only once. Dilution and stability of every antibody or antibody batch have to be tested separately.

7. If the antibody staining shows a high level of nonspecific background, it may be pre-absorbed with fixed embryos: incubate fixed embryos overnight with the antibody solution. Use this antibody solution and use it for staining your sample.

8. To avoid bacterial contamination and to extend stability of the antibody solution 0.05 % of sodium azide can be added.

9. Choose an antibody that targets the animal in which the primary antibody was produced. Conjugated Alexa fluor antibodies that were raised to target different animals and that have sufficiently different emission spectra can be combined to immunostain different antigens at a time. Additionally, if the primary antibodies are from different antibody subclasses (IgM or IgG subclasses) secondary antibodies specific to the antibody subclass can be used to discriminate the patterns. Sensitivity and/or cross-reactivity can vary. We have had good experiences with Alexa 568 goat anti-mouse (emits red light), combined with for example Alexa 488 goat anti-rabbit/rat (emits green light) and Alexa 633 goat anti-rat/rabbit (emits far-red light).

10. If the available confocal microscope does not have a UV laser illumination system that is necessary to detect DAPI-stained DNA, you may use propidium iodide to stain the nuclei instead. Propidium iodide staining: Incubate the ovaries for 15 min in PBS containing 2 μg/ml propidium iodide. Staining with DAPI allows you to use three other secondary antibodies emitting green, red, and far-red light in parallel with DNA staining, whereas propidium iodide emits red light itself.

11. Add NPG to the glycerol and vortex. If the NPG will not dissolve, heat the solution at 37 °C overnight.

12. Oogenesis is highly dependent on the individual’s environment. Therefore, the flies should be “fattened” on wet yeast prior to dissection for at least 2 days and should also be kept in a community with males. However, when analyzing ovarian phenotypes it is recommended to collect and stain wild-type and mutant females at different timepoints and in several independent experiments.
13. Depending on the antibody used, the immunostaining protocol can vary. If the used antibody is also staining the peritoneal muscle sheath, it is necessary to destroy the sheath around the ovarioles by sucking them up and down several times in a Pasteur pipette.

14. The larval fat body that contains the ovaries will not sink to the ground, but floats in the solution. It is therefore recommended to check under the stereomicroscope that the fat bodies are not washed away when adding or removing liquids from the sample.

15. The ovaries should be fixed as fast as possible after dissection to preserve the cellular structures. It is important not to exceed or shorten the fixation time to avoid poor immunostaining.

16. Upon all incubation and washing steps make sure to add an amount of liquid that is sufficient to allow the ovaries to float in the tube or the plate upon gentle rocking. Furthermore, when removing solutions from the tube do not pull up the ovaries into the pipette and do not damage the ovaries. The ovaries should stay intact until the very end of the procedure since the individual germaria are otherwise very easily lost.

References


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