Telomeric PNA FISH and COFISH on Metaphase Chromosomes

**Required Solutions/Reagents**

**FISH & COFISH**

**Colcemid**  
(Roche 10295892001, 10 ug/mL, stored at 4°C)

**Fixative**  
Make fresh.  
3:1 MeOH and glacial acetic acid.

**Blocking Reagent**  
(Roche 11096176001)  
Make a 10% stock, dissolved in maleic acid buffer:  
100 mM malic acid  
150 mM NaCl  
Adjust pH to 7.5 (20°C) with NaOH.  
Store at 4°C.

**Hybridizing Solution**  
*Make fresh, but can be stored at 4°C for 2 weeks if you have excess.*  
10 mM Tris-HCl pH 7.2  
70% formamide (from the deionized stock)  
0.5% blocking reagent (from 10% stock)

**Hybridization Wash #1**  
10 mM Tris-HCl pH 7.2  
70% formamide  
(Optional) 0.1% BSA, must be dissolved before adding formamide.

**Hybridization Wash #2** (Alternatively, PBS can be used)  
0.1 M Tris-HCl pH 7.2  
0.15 M NaCl  
0.08% Tween-20  
Add DAPI to 2nd of 3 washes:  
FISH – 1:3,000 DAPI from 0.5 mg/mL stock  
COFISH – 1:750 DAPI from 0.5 mg/mL stock

**PNA probes from PNA Bio Inc.**  
**FITC-TelC (F1009)**  
TelC-FITC: FITC-OO-CCCTAACCCTAACCCTAA 3’  
Make 100μM stock in formamide. Heat 10 min at 60°C and vortex occasionally.  
Make 10 μl aliquots and keep at -80°C for long-term storage.  
Store thawed aliquots at 4°C (in the dark).  
Dilute 1:1000* and heat the dilution for 5 min at 60°C before use.
Cy3-TelG (F1006)
TelG-Cy3: Tam-OO-TTAGGTTAGGGTTAGGG 3’
Make 50µM stock in formamide. Heat 10 min at 60°C and vortex occasionally.
Make 10 µl aliquots and keep at -80°C for long-term storage.
Store thawed aliquots at 4°C (in the dark).
Dilute 1:1000* and heat the dilution for 5 min at 60°C before use.

* Adjust dilutions based on signal intensity and/or background.

DAPI
Reconstitute at 0.5 mg/ml, store at -20°C in the dark (Sigma D-9542)

ProLong Gold Antifade Reagent
(Life Technologies P36934)

Frosted Slides
(Fisher Scientific 12-550-11)

Microscope Cover Glass
(Fisherbrand 12-545-M 24x60)

COFISH

BrdU/BrdC
BrdU (MP Biomedicals 100166) stock concentration: 10 mM in H2O
BrdC (Sigma B5002) stock concentration: 10 mM in H2O
Make 1000x working stock solution of BrdU/BrdC by mixing 3 parts BrdU to 1 part BrdC (final concentrations of 7.5 mM of BrdU and 2.5 mM of BrdC.)
Store at -20°C covered in aluminum foil.

RNase A
(Sigma R5000)
Stock solution: 50 mg/mL in 10 mM Tris-HCl pH 7.2
Heat inactive 10 min at 80°C. Store at -20°C.

Hoechst 33258
(Life Technologies H21491, 100 mg)
Stock solution: 10 mg/ml in H2O. Store at 4°C covered in aluminum foil.

Exonuclease III & buffer
(Promega M1811)
Methanol/acetic acid fixed metaphase spreads

1. Split cells 36-48 hours before harvest to be ~50% confluent at time of harvest.
2. For COFISH, add BrdU:BrdC 12-16 hours before harvest (for a little less than one cell cycle to avoid double labeling).
3. Incubate 30 min – 2 hours in regular medium with 0.1 μg/ml of colcemid for human cells (e.g. 0.05 μg/ml for RPE1 cells, 0.1 μg/ml for HeLa cells) and fast-growing mouse cells, and 0.2 μg/ml for slower-growing mouse cells – the effects of the colcemid should be obvious at time of harvest (rounded, refractile cells with blebby membranes).
3. Harvest cells by trypsinization, neutralize trypsin by adding media, and spin down (5 min at 1000 rpm). Be gentle during harvesting, many of the dividing cells tend to lift off. Cells that have floated off during the harvest can be recaptured by spinning down the wash supernatant as well. You can also use the conditioned media to neutralize the trypsin so that you can recover any floating mitotic cells.
4. Remove supernatant completely, resuspend in 5 ml of 0.075 M KCl (pre-warmed to 37°C), be gentle to prevent lysis, this step swells the cells.
5. Incubate for 15 to 30 min at 37°C, invert tubes every 5-10 min to keep cells suspended. (Concentration of KCl, incubation time and temperature can be tested to keep the cells intact during swelling.)
6. Spin the cells down, 5 min at 1000 rpm in a benchtop centrifuge. (A couple of drops of fixative, approx 500μL, can be added before spinning, if you’re having problems with a lot of cell debris when you drop metaphase spreads.)
7. Decant the KCl, resuspend cells fully in the small volume of KCl that was left (by tapping).
8. Drop by drop add 1 ml of cold fixative while the cells are slowly and gently mixed on a vortex (<1000 rpm).
9. Fill to 10 ml with the fixative and store at 4°C o/n or longer; cells can be kept at this stage indefinitely.
10. When ready to drop, spin the cells down (1000rpm).
11. Resuspend cells in 1ml new fixative (may vary depending on cell number), or the old fixative left in the tubes.
12. Place a few slides in cold water.
13. In the thermotron, drop the resuspended cells from a couple of inches above the end of the wet slide tilted at a 45° angle, wash the nuclei with fresh fixative (drop fixative across the slide with a bulb and Pasteur pipette; the nuclei and chromosomes should not wash off.)
14. **If not using thermotron:** for FISH, place wet paper towels on top of a heating block set to 80°C. Place slides after dropping metaphases for 1 min on the humidified 80°C heating block.
15. **If not using thermotron:** for COFISH, place wet paper towels on top of a heating block set to 42°C. Place slides after dropping metaphases for 1 min on humidified 42°C heating block.
16. Check the slides under a regular light microscope for spreading efficiency. You should see many nuclei (all of the cytoplasmic membranes should be washed away or barely visible.) Well spread metaphase chromosomes should look like small black dots at low magnification. The arms of the chromosomes should be visible at higher magnifications. If the nuclei are too crowded or too sparse, you may need to dilute or concentrate your sample and drop the slide again.
17. Let the slides dry o/n, protected from light if intended for COFISH.
COFISH

Degradation of the Newly Synthesized Strand
1. Rehydrate slides in PBS for 5 min at rt. *Perform all steps in coplin jars that are protected from light.*
2. Treat slides with 0.5 mg/ml RNase A (in PBS, DNase free) for 10 min at 37°C.
3. Stain slides with 0.5 µg/ml Hoechst 33258 (Sigma) in 2x SSC for 15 min at rt.
4. Place slides in a shallow plastic tray (*the ones used for DNA gels*) and add enough 2x SSC to just cover the slides. Expose slides to 365 nm UV light at rt (Stratalinker 1800 UV irradiator) for 5.4x10^3 J/m^2 (choose ‘energy’ and enter 5400 on Stratalinker display).
5. Digest the BrdU/BrdC-substituted DNA strands using coverslips on the slides with at least 95 µl of 10 U/µl of Exonuclease III (Promega) in buffer supplied by manufacturer (50 mM Tris-HCl, 5 mM MgCl2, and 5 mM DTT, pH 8.0) at 37°C for 30 min.
6. Wash in PBS for 5 min.
7. Dehydrate in ethanol series: 5 min in 70%, 95%, 100% at rt and air dry slides (slides can be stored at rt in the dark for several days).

Hybridization
1. Place 95 µl of hybridization mix (see recipe) onto coverslips and pick up the coverslip with the metaphase slide (cells down), then turn the slide up.
2. Hybridize for 1.5 – 2 hours with TelG-Cy3 (1:7500), at rt, in the dark, with wet paper towels.
3. Rinse in wash #1 for 2-5 seconds. *Dry slide briefly by dabbing on Kimwipe.*
4. Hybridize for 1.5 – 2 hours with TelC-FITC (1:1000), @ rt, in the dark, with wet paper towels.
5. Take off coverslip, wash slides in hybridization wash #1: 2 times 15 min, on shaker.
6. Wash slides in hybridization wash #2: 3 times 5 min on shaker. To second wash, add 1:750 DAPI from 0.5 mg/ml stock.
7. Dehydrate in ethanol series: 5 min in 70%, 95%, 100% ethanol at rt and air-dry slides.
FISH
(L Lansdorp et al. Hum Mol Gen, 1996, 5, 685-691.)
(The first section can be skipped entirely. Alternatively, rehydration/fixation with formaldehyde/dehydration can be performed to help make the DAPI and telomere signals look more compact and nicer. Additional pepsin treatment strongly increases signal intensity and makes the chromosomal arms skinny.)

Rehydrate MetOH/Acetic acid spreads in PBS (pH 7.0-7.5) for 5 minutes.
Fix in 4% formaldehyde in PBS, for 5-30 min (dilute from 37% commercial formaldehyde).
Wash in PBS 3 times, 5 minutes each.
Perform the following optional steps
a. Treat with pepsin (1 mg/ml) @37°C, 10 min; pepsin prepared fresh in 10 mM glycine pH 2 and warmed up to 37°C.
b. Wash in PBS 2 times, 2 min each.
c. Fix in 4% formaldehyde in PBS, 2 min.
d. Wash in PBS 3 times, 5 min each.
Dehydrate in ethanol series: 5 min in 70%, 95%, 100% at rt and air dry slides.
1. Place 95 µl of hybridization mix (see recipe) onto coverslips and pick up the coverslip with the metaphase slide (cells down), then turn the slide up.
2. Denature on a 80°C hot plate, 3 – 10 min.
3. Hybridize for 1 – 2 hours, at rt, in the dark, with wet paper towels, or overnight at 4°C.
4. Take off coverslip, wash slides in hybridization wash #1: 2 times 15 min, on shaker.
5. Wash slides in hybridization wash #2: 3 times 5 min on shaker. To second wash, add 1:3000 DAPI from 0.5 mg/ml stock (1:500-1000 if without pepsin treatment). 
6. Dehydrate in ethanol series: 5 min in 70%, 95%, 100% ethanol at rt and air-dry slides.

SCORING

Fragile Telomere
score well-separated telomere signals, only score long-arms for mouse cells

Telomere-Sister Chromatid Exchange
score well-separated telomere signals, only score long-arms for mouse cells

SCORING