## Mouse cochlear culture preparation

## Supplies:

DMEM/F12 medium (Sigma D8062)
Foetal calf serum (any supplier)
Cell-Tak (BD Bioscience, BD 354240)
HBSS (GIBCO 14025-050)
Hepes 1M (Sigma, H0887)
0.1 M sodium bicarbonate pH 8.0 (sterile filtered)
Mice (postnatal days 0-3)
Mat-Tek dishes (MatTek Corporation, Cat # P50G-0-14-F)
Ampicillin (Sigma A-9518; 10 mg/ml in H2O, sterile filtered)
90 mm and 35 mm diameter sterile plastic petri dishes (any supplier)

## Equipment:

Horizontal lamina air hood Dissecting microscope with fibre optic light source Dissecting instruments CO2 incubator

## Procedure:

- 1. Prepare medium: 93 ml DMEM/F12, 7 ml FCS, 100 ul ampicillin (10mg/ml).
- 2. Prepare Hepes buffered Hanks' Balanced salt solution (HBHBSS): Add 5 ml 1 M Hepes to 500 ml HBSS.
- 3. Prepare Cell Tak coated Mat Tek dishes: To 20 ul Cell Tak add 300 ul 0.1 M sodium bicarbonate pH 8.0; immediately add 60 ul to each well and spread. Replace lid; do not let the Cell Tak dry out. Wash 2 with HBHBSS before adding tissue (see below).
- 4. Kill mouse pups via approved method.
- 5. Surface sterilize pups by immersion in 80% ethanol for 6 min (3 changes, 2 min each).
- 6. Cut of heads and drop into 90 mm diameter dish containing HBHBSS (if using mutants remember to snip the tail and freeze for subsequent genotyping).
- 7. Bissect heads in two along the mid-saggital plane.
- 8. Transfer half heads to 90 mm diameter dish with HBHBSS.
- 9. Remove brain, pop out the cochleae from half heads, separating them from the vestibular bit of the labyrinth and place in 35 mm diameter dish with HBHBSS.
- 10. Remove cartilagenous capsule and transfer (with forceps) the still coiled cochlea complete (if possible) with stria vascularis to a 35 mm diameter dish with HBHBSS.
- 11. Remove (unwind) the stria, and separate the cochlear coil (GER/LER complex) from the mesenchymal modiolar tissue without unwinding or stretching the epithelium.
- 12. Cut cochlear coils into basal and apical 'halves' (cochlear is only 1.5 to 1.75 coils at this stage) with a pair of sharp needles.
- 13. Transfer basal and apical coils (with a serum prewetted, bent glass Pasteur pipette, or with a curette [a small spoon]) to a dish of clean HBHBSS.
- 14. Transfer coils to the medium filled wells (200 ul per well) of the Mat-Tek dishes with a curette, making sure the coils are sunny-side (hair-cell side) up before sliding them out of the spoon onto the substrate.