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.
15. Transfer to 37C incubator

Notes:

A: Serum concentration can be reduced to 1% to reduce spreading.
B: Less Cell-Tak can be used for coating Mat-Tek dishes (a minimum of 15 ul per well is sufficient).
C: All surgical instruments should be dry heat sterilised (180C for 2h), or soaked in 80% ethanol (for 30 min) and dried in the hood, before use.

Biological transfection of cochlear cultures:

Equipment:

Gene Gun (Bio-Rad #164-2431)
Helium cylinder with regulator
Gene gun holder (home made)
Gene gun barrels (home made)
Filters (cell culture inserts with 3.0 um pore size PET track-etched membrane; BD 353092)

Procedure:

(NB. Wear safety glasses at all times, and try not to shoot yourself in the foot or anywhere else)

Remove bullets from fridge and allow them to warm up to room temp.
Connect up gun to helium cylinder, set pressure to 120-140 psi, and purge gun with helium (i.e., shoot it a few times!).
Load bullets into cartridge (bullet) holder, place holder in gun, place gun in stand.
Fire one shot onto filter paper to check dead centre.
Remove all medium from the culture well.
Place 3.0 um filter over the well with the membrane resting on the plastic surrounding the well, and put the dish plus filter in the gene gun stand with the culture centred.
Check pressure is between 120 and 140 psi, and fire!!
Remove dish from stand, add fresh medium, and check gold particle distribution using dissecting scope.
Replace culture in incubator.

Notes:

A. Cartridge (bullet) holders and barrels are sterilized in ethanol (805, 30 min) and dried before use. If using several constructs sonicate the barrels and clean them well before using a new construct.