Introduction
The following protocol outlines how to coat alvetex® membranes with Basement Membrane Extract to facilitate and enhance attachment and differentiation of both normal and transformed anchorage dependent cells. Example data shown herein was obtained using this protocol to grow HepG2 hepatocytes in alvetex® for 7 days in 6-well inserts (AMS.AVP004-32) in 6-well plate format.

Method:
1. Prepare alvetex® for coating by first treating with 70% ethanol followed by two PBS washes as described in the relevant product information leaflet. Leave alvetex® in the second PBS wash until ready to apply the Basement Membrane Extract solution.
2. Dilute Basement Membrane Extract to a concentration of 0.8 mg/ml (1 in 10 dilution) using appropriate cell culture media (e.g. MEM for the example below). Handle the reagents on ice, using pre-chilled pipette tips to perform the dilution and subsequent application onto alvetex®.
3. Aspirate the second PBS wash from alvetex® disc and carefully pipette 350 µl of the diluted Basement Membrane Extract solution onto each disc. Replace plate lids and leave to stand for 1-2 hours at room temperature.
4. Remove excess fluid from alvetex® in well insert format by gently tapping the plate or Petri dish on the worktop. Check that no residual fluid is hanging from the base of the well inserts. Aspirate to remove any residual coating agent from the bottom of the wells. If using alvetex® in 12-well plate format, tilt the plate and gently aspirate any excess fluid from the edge of the wells.
5. Prepare cells for seeding in the appropriate culture media and seed directly on the wet Basement Membrane Extract coated alvetex® membrane in the volumes relevant to the alvetex® product format. Allow the cells to settle for 30-90 minutes in an incubator (5% CO₂, 37 °C) before flooding with media.
Example: Growth of HepG2 Hepatocyte Cell Line in Basement Membrane Extract Coated Alvetex®

Cell Culture details:
HepG2 cells (ATCC, HB-8065) were routinely maintained in T-75 flasks. HepG2 complete media consisted of: MEM media (Gibco, 21090) supplemented with 10 % v/v FBS, 2 mM L-glutamine and 100 U/ml Penicillin/Streptomycin. Alvetex® 6-well inserts (AMS.AVP004-32) in 6-well plates, were coated with Basement Membrane Extract as described above. Cells were seeded at a density of 1 x 10^6 cells in 150 µl media suspension per disc and were left to settle for 60 minutes in an incubator (5 % CO₂, 37 °C). Media was carefully added to each sample well (9 ml per well). Cultures were maintained for 7 days, with media changed on days 3 and 5.

Results:
Pre-coating of alvetex® discs with Basement Membrane Extract resulted in enhanced infiltration of cells into the scaffold compared with control cultures in untreated alvetex®. Cells were seen to occupy the entire depth of the scaffold after 7 days of growth in Basement Membrane Extract -coated discs, while cells grown in untreated alvetex® occupied only the upper half of the scaffold. These findings indicate that pre-treatment of alvetex® with extracellular matrix products is able to enhance the growth of appropriate cell types into the 3D structure.

Uncoated alvetex® control

Basement Membrane Extract -coated alvetex®

Figure 1. Brightfield micrographs at low (10x) and high (40x) magnification showing HepG2 cells cultured for up to 7 days on 22 mm diameter alvetex® discs presented in 6-well insert (AMS.AVP004-32) in well insert holders in 6-well plate format. Cells were fixed, sectioned and counterstained with haematoxylin and eosin.