

Anticonvulsant Screening Program

Test 76 Results - In-vitro Hippocampal Slice Culture Neuroprotection Assay (NP)

ASP ID: 330044 U Screen ID: 1

Solvent Code: DMSO Solvent Prep:

Test Date: 13-Jan-2009

Reference: 439:27

Summary of NP Assay: Kainic acid

- Test Result: No Neuroprotection
- ADD compounds evaluated: 330044 330045

Note: This experiment is run at two different concentrations of candidate drug against a fixed concentration of excitotoxin. If multiple candidates from the same participant source are scheduled for NP screening we will test compounds in pairs whenever possible.

Comments:

TEST 76: *in vitro* HIPPOCAMPAL SLICE CULTURE NEUROPROTECTION ASSAY

Compound 1 : ADD Number: 330044 Batch U Screen ID: 1 Date Started: 01-13-2009

Compound 2 : ADD Number: 330045 Batch U Screen ID: 1 Date Completed: 01-30-2009

References: 439 : 27

Excitotoxin: KA Insult Duration: 4 Hours Solvent: DMSO

Primary Screen Results: No neuroprotection observed

EXPERIMENT IMAGES & WELL DESCRIPTION

A1	KA 20µM	A2	KA 20µM +	A3	KA 20µM +
			330044 10µM+ 330045 10µM		330044 10µM+ 330045 10µM
					
					
B1	KA 20µM	B2	KA 20µM +	B3	KA 20µM +
			330044 100µM+ 330045 100µM		330044 100µM+ 330045 100µM

PRIMARY SCREEN EXPERIMENT DESCRIPTION

The "Primary Screen Experiment" is a qualitative assessment of the ability of a compound to prevent excitotoxic cell death. Organotypic hippocampal slice cultures are treated with N-methyl-D-aspartate (NMDA) or kainic acid (KA) to induce neuronal cell death. Propidium iodide (PI), a membrane-impermeant compound, is included in all wells of the culture plate. Dying cells have compromised cell membranes, thus PI may diffuse into the cell, intercalate with DNA and fluoresce. Thus, the intensity of the PI fluorescence is proportional to the amount of cell death in the individual slices. Hippocampal slice cultures are treated with the excitotoxin alone, or where indicated above, with the excitotoxin and either one or two investigational compounds at the concentrations indicated. If neuroprotection occurs as a consequence of the added compound, slice cultures will have a visibly reduced fluorescent intensity when compared to the slice cultures that have been treated with the excitotoxin alone.