Investigator:_	A-Biphagee
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Ciarance	LIE IN	aciophage-Colc	ANY FORTHING UTIL (GIVI-CFU) ASSAY
Experiment #	þ	(Survival #2	Source of Irradiation: Radionuclide
Mice Sex, Strain,	Age:_	SW, F, 5-6 WK	injection
Type of Irradiation	on:	Chronic	•

Animals per group: 3

To determine GH-CFC response in viouse bone marrow 7 days following or single injection of so-different amount of 17Sn-DTPA Aims: Summary of Results:

#### Brief Procedure:

- Inject animals in groups of 3 with desired activity of Sn-117m intravenously through lateral tail
- 2) Sacrifice each mouse on optimal day by cervical dislocation and sterilize using 70% EtOH and immediately move it into laminar flow hood.
- 3) Remove both femurs carefully using sterile instruments and clean the attached tissue thoroughly.
- Flush the bone marrow with 2% Horse Serum in Dulbecco's Modified Eagles Medium (2% HS-DMEM) 4) using 21G needle and syringe.
- Separate the mononuclear cells by density gradient procedures using Histopaque. 5)
- Plate the desired number of cells (cell suspension) in mixture of 60% HS-DMEM and 1.7 ml 0.6% bacto agar solution in the presence of 9.2 U (New Sigma Unit) of GM-CFS.
- 7) Keep the plated petri dishes for 20 min. in laminar flow hood and move them into incubator with 5% CO<sub>2</sub> and 95% air, at 37°C.
- 8) Count the granulocyte macrophage colonies on 7th day.

### Sn-117m injections:

Group# Probe#	Date of injection	Activity injected (KBd/g)	Date Sacrificed	# of days	Remarks if any		
Cl	4/24/98	0	19/01/98	7	0.9% Nacl	injected	control
C2	9/24/98	0	10/01/98	7	ч	<b>લ</b>	ч
1	9/34/98	57	10/01/98	7			
2	9/24/98	117	10/01/98	7			
3	9/24/98	178	10/01/98	7			
4	9/24/98	224	10/01/98	7		-	
5							
6							
7							
8							

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## Preparing Media and Agar.

Culture Medium (Double Strength, 2X): 13.37g (1 pack) of DMEM powder (Gibco, Cat # 12100-038) + 490 ml deionized water + 16 μl of L-asparagine (Gibco Cat # 12416-012) at a concentration of 5 μg/μl + 150.4 μl of DEAE dextran (mol. wt. = 2x106, intrinsic viscosity = 0.7) at a concentration of 1 μg/μl (Sigma Cat # D-9885) + 10 ml of penstrep (Gibco Cat # 600-5070, 5,000 units/ml pen, 5,000 μg/ml streptomycin) + 3.7 g of NaHCO<sub>3</sub> (Gibco Cat # 11810-025).

Culture Medium (2X) with 60% Horse Serum: Add 60% Horse Serum to 2X DMEM

Wash Medium: i) Mix equal amounts of culture medium and sterile deionized water. ii) Add 2% HS

Agar: Prepare 0.6% agar by adding 0.6 g Difco Bacto agar (Difco Cat # 0140-15-4) to 100 ml boiling deionized water. Autoclave on liquid cycle for 20 min.

Comments If any:

## Flushing Bone marrow:

 Remove both femurs from each mouse and place them in a test tube containing wash medium kept in ice, if the femur can not be flushed immediately.

2) Flush the marrow from each femur by aspirating 3 ml of Wash Medium through the femur 5 times with a 21G needle/3 ml syringe in a 50 ml conical centrifuge tube. Follow with two flushes with 1 ml of fresh Wash Medium.

3) Spin the cells at 1200 rpm for 5 minutes at 4°C, decant, break up the cell pellet, resuspend the cells in 5 ml of cold Wash Medium, and vortex the cell suspension.

## Comments If any:

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### **Counting the Cells:**

Add 10µ1 of cell suspension to 20 ml of Isotone II in a coulter cup and count the cells using coulter counter. Calculate total # of cells in each group.

## **Coulter Counter Parameters:**

Current(I)=500  $\mu$ A Full Scale = 1  $T_L$  = 2.7  $T_u$  = 99.9 Manometer Select = 500  $\mu$ 1

Attenuation= 4
Alarm Threshold = off
Preset Gain = 1
Stirrer control = off

# Multiplication Factor to get total # of cells in 5 ml = 20,000 x Coulter count

Group #	Coulter Count without ZG	Avg	Total # of cells	Coulter Count with 5 drops ZG	Avg	Total # of cells
Cl						O. COAD
C2						
1						
2						
4						
5						
6						
7						
_		1 1				

### Comments If any:

Separating Mononuclear cells and washing the cells:

- Transfer 3.5 ml of Histopaque (Sigma Cat #H8889) into fresh 15 ml tubes (1 tube
- 2) Layer the cell suspension carefully on top of Histopaque and centrifuge at 1500 rpm, 4°C, for 30 minutes.
- 3) Using a Pasteur pipette transfer the mononuclear cells into fresh 15 ml tubes.
- Dilute the cell suspension to 15 ml by adding cold Wash Medium into each tube 4) and spin them at 1200 rpm, 4°C, for 5 min.
- 5) Decant the supernatant, break the pellet, and add 15 ml cold Wash Medium, and spin them again at 1200 rpm, 4°C, for 5 min. Repeat this procedure 2 more times.
- After 3rd wash break the pellet and resuspend in 2 ml Culture medium (2x 6) DMEM) with 60% HS and keep the tubes in dry bath at 37°C.
- Add 20  $\mu l$  of cell suspension to 20 ml of Isotone II in a coulter cup and determine 7) total # of cells in each group using coulter counter.

Coulter Counter Parameters: Same as above

# Multiplication Factor to get # of cells/ml = 2000 x Coulter Count

Group #	Coulter Count without ZG	Avg	# cells per ml	Coulter Count with 5 drops ZG	Avg	# cells per ml
CI				\$504, 5457, 5442	10	10000533
CZ				4553, 4591, 4454	(45)	
1				3616, 3360, 3327	343A	68 68 666
2				2823, 2894, 2891	2869	c738 600
3					2151	430200
4					2299	45 89 333
5						· · · · · · · · · · · · · · · · · · ·
6						· · · · · · · · · · · · · · · · · · ·
7						
8						

Average # of cells per ml =  $5000 \times 2000 = 10,000,333$ 

### DILUTIONS

Dilution A: (1.0x106 cells /ml, Total volume 3.4 ml)

Vol of final cell suspension required = 3400000/cells per ml = 3400000/16000333 = 0.339

1.7 ml Agar + 1·36 ml Medium + 0·339ml Cell Suspension

Dilution B: (3.0x105 cells /ml, Total volume 3.4 ml)

Vol of final cell suspension required = 1020000/cells per ml =1020000/10000333 2 0.101

1.7 ml Agar + 1.60 ml Medium + 0.101 ml Cell Suspension

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### Plating the Cells:

Culture Medium: Maintain four 13mm tubes each containing 4.5 ml of Culture medium in dry bath at 37°C.

Agar: Maintain five 16mm tubes each containing 6.5 ml of Agar in dry bath at 37°C.

1) Warm up dilution tubes (2 or 3 per group) to 37°C in dry bath.

- 2) Warm up Agar (30 ml) and 60% HS in 2x DMEM (30 ml) in separate tubes to 37°C.
- 3) Mark the Six-well plates (3 wells for each dilution for each group) containing 20 u1 of stock GM-CSF (9.2 U) in each well.
- 4) Mix 1.7ml agar + x ml of 2x DMEM with 60% HS + y ml cell suspension + 0.02 ml GMCSF (x +y = 1.7 ml) in a dilution tube.
- 5) Add 1 ml of mixture 4 to each well, mix properly and let it gel for about 30 minutes.
- 6) Repeat steps 4 and 5 for each dilution.

7) Repeat steps 1 to 6 for each group.

- 8) Incubate the cells in an incubator at 37°C and 5% CO<sub>2</sub>, 95% air for 7 days.
- 9) On 8th day of incubation count colonies and determine the survival fractions.

Counting the Colonies: (Inverted at 40X or dissecting at 35X)

10/08/18

Group #	Dose Activity <sup>#</sup> injected	# of cells plated	# CFU-GM counted	Avg	SF
CI		3 ×10 <sup>5</sup>	169, 172, 159	] 160.83	
C2		3×16 <sup>-5</sup>	156, 149, 160		
1		3×105	80, 88, 95	87.66	0.5450
2		3×105	33, 47, 41	40.33	0.2507
3		3×105	34, 23, 21	<del> </del>	0.1616
4		3×105	34, 23, 21 20, i4, 15	16.33	<u> </u>
5					
6					
7					
0					