

June 26, 1951.

Program: streak out W-1 on EMV B agar. ^{Point} Velect Transfer to plain and to selective medium. When V_1^R clones are detected, take plain inoculum for second streaking. Repeat until V_1^R colonies are obtained by indirect selection without exposure to specific selective agent.

P26. Streak out W-1.

A27. Restreak single colonies. Make test ~~transfer~~ ^{Point} Transfer

A. Preliminary test (whole culture not recently isolated: more likely to produce clones from V_1^R in inoculum).

Transfer somewhat random.

T₁. 1. 21 V_1^R } 6 congruences, presumably clones
T₁. 2. 13 V_1^R }

1. Pool pickings from homologous sites on plain transfer plate. Restreak.

^{Point}
B } Single colony platings. Transfer to P₁, T₁ 1, 2
C }
D }

B. 6 clones. 14 V_1^R ; 14 V_1^R

C 7 clones 30, 29.

D 1? clone. 12, 12

A29. Pool pickings from homologous sites and restreak on EMV Lac. ^{Point} Transfer to plain, T1 agar.
save original cultures

A 15? clones; ca 25 mutants Restreak from 5 sites

B 13? " ca 40 " " " " (without pooling)

C 12 " all with thick sheels 18 muts.

D 1 21

A3-4. 14 clones; 37 mutants. Pool from 5 sites into ca. 1 ml.
 Plate serially to further dilutions.
 Suspend B, C experiment.

D transfer to D11; 2 T1. 7/3/51. Hold in Refrig.

7/4/51. ~~Transfer D3~~

D3: 3-4 clones (?) ca 20 mutants.
 Plate as D4:

A4. Transfer to series of T1 plated at 1, 3, 6" spreading.

This procedure is obviously not working ~~probably~~ properly, presumably because of confusion between mutants within an inoculum to a plate, and those which occur during the growth of colonies on the plate. The samples from homologous sites are probably too small to have a reasonable probability of carrying an identified clone to the next inoculum.

The procedure should be modified as follows: A broth culture should be spread or streaked on plain agar, and permitted to grow for just a few generations. The thinly grown plate should then be ~~transferred~~ ^{printed to} selective and plain agar, and the clones identified. The homologous site should then be inoculated to a small volume of broth, which is then restreaked in the same manner.

See Allebury
 Fed. Proc. 1952

Hermit Leilage

June 26, 1951

A W1177 x 1632

EMS Lac.

B W1635 x 1632

Count from plates: (Lac)

A.	+	-
	17	30
	2	27
	13	33
	14	26
	46	116

B

+	-
187	18

"B" B Lac + } selections
 "C" B Lac - }

Picks "A" at random to EMS Lac. B 100+
 Paint unconservative selections obvious 48-

Transfer to EMS Lac, Mal, MH, sm; TI for scoring

	Lac	Mal	S	MH	TI	Bal	Xyl
1	-	-	R	-	S	-	-
2	+	-	R	-	S	-	-
3	-	-	R	-	S	-	-
4	-	-	R	-	R	-	-
5	+	-	R	-	S	-	-
6	-	+	R	-	S	-	-
7	-	-	R	-	R	-	-
8	-	-	R	-	R	-	-
9	-	-	R	-	S	-	-
10	-	-	R	-	R	-	-
11	-	-	R	-	S	-	-
12	-	-	R	-	S	-	-
13	-	-	R	-	R	-	-
14	-	-	R	-	S	-	-
15	-	-	R	-	R	-	-
16	-	-	R	-	R	-	-
17	-	-	R	-	R	-	-
18	+	+	S	+	S	+	+
19	-	-	R	-	S	-	-
20	-	-	R	-	S	-	-
21	+	+	R	+	S	+	+
22	-	-	R	-	S	-	-
23	+	-	R	-	S	-	-
24	+	-	R	-	S	-	-
25	+	+	S	+	S	+	+
26	+	-	R	-	S	-	-
27	+	+	S	+	S	+	+
28	+	-	R	-	S	+	-
29	+	-	R	-	S	-	-
30	+	-	R	-	S	-	-

A

	Lac	Mal	SV	MHE	TI	Gal	Xgl
31	+	-	R	-	S	-	-
32	+	+	S	+	S	+	+
3	-	+	R	-	R	-	-
4	-	-	R	-	S	-	-
5	+	-	S	-	S	-	-
6	+	-	R	-	S	-	+
7	-	-	R	-	R	-	+
8	+	-	R	-	S	-	-
9	-	-	R	-	S	-	-
40	+	-	R	-	S	-	-
41	+	+	R	-	S	-	-
2	-	+	R	-	S	-	-
3	-	+	R	-	S	-	-
4	-	-	R	+	R	-	+
5	-	-	R	-	R	-	-
6	-	-	R	-	R	-	-
7	-	-	R	-	S	-	-
8	-	-	R	-	S	-	-
9	+	-	R	-	S	-	-
50	+	+	S	+	S	+	+
51	+	-	R	+	S	-	-
2	-	-	R	-	S	-	-
3	+	+	S	+	S	+	+
4	+	-	R	-	S	+	+
5	+	+	S	+	S	+	+
6	-	-	S	-	S	-	+
7	-	+	R	+	S	-	+
8	-	-	R	-	S	-	-
9	-	-	R	-	R	-	-
60	+	-	R	-	S	-	-
61	-	-	R	-	R	-	-
2	-	-	R	-	R	-	-
3	+	-	R	+	R	-	+
4	+	+	R	+	S	-	+
5	+	+	S	+	S	+	+
6	-	-	S	-	S	-	-
7	-	-	S	-	S	-	-
8	-	-	S	-	R	+	-
9	+	+	S	+	S	+	+
70	-	-	S	-	S	-	-
71	+						
2	+						
3	+						
4	+						
5	+						
6	*						

	lac	Mal	S	MTR	TI	gal Xgl	Xgl
71	+	✓ -	R	-	S	-	-
72	+	✓ +	S	+	R	+	+
3	+	✓ -	X	-	S	+	-
4	+	✓ -	S	+	S	+	+
5	+	✓ -	R	-	S	-	-
6	+	✓ +	S	-	R	-	-
7	+	✓ +	S	+	S	+	+
8	-	✓ -	R	-	R	-	-
9	+	✓ +	S	+	S	+	+
80	+	✓ -	R	-	S	-	-
81	+	✓ -	X	-	S	-	-
2	-	✓ -	R	-	S	-	-
3	+	✓ +	S	+	S	+	+
4	+	✓ +	S	+	S	+	+
5	-	✓ -	R	+	R	+	+
6	-	✓ -	S	-	R	-	-
7	-	✓ -	R	-	R	-	-
8	-	✓ -	R	-	R	-	-
9	-	✓ -	R	-	R	-	-
40	+	✓ +	R	-	S	+	+
11	-	✓ +		-	R	-	-
2	-	✓ -		-	S	-	-
3	+	✓ +	S	+	S	+	+
4	+	✓ +	S	-	S	+	-
5	+	✓ +	S	+	S	+	+
6	+	✓ -	R	-	R	+	-
7	+	✓ -	R	-	R	-	-
8	-	✓ -	R	-	R	-	-
44	+	✓ +	S	-	S	+	-
100	+	✓ +	S	+	S	+	+

MTR

b2

	Lac	Mal	MH	S	TI	X	Gal
1	+	+	+	R		+	+
2		+	+	R		+	
3		+	+	R		+	
4		+	+	R		+	
5		+	+	R		+	
6		+	+	R		-	
7		+	-			-	
8		+	+	R		+	
9		+	+	R		+	
10		+	-	R		-	
11		-	-	R		-	
12		+	+	R	R	+	
13		+	+		R	+	
14		+	-			+	
15		+	-			+	
16		+	+			+	
17		+	-			+	
18		+	-			+	
19		+	+			+	
20		-	-	R		+	
21		+	+			+	
22		+	+			+	
23		+	+			+	
24		-	-			+	
25		+	+	R		+	
26		+	+			+	
27		+	+			+	
28		+	+			+	
29		+	+		R	+	
30		+	+			+	
31		+	+			-	
32		+	+	R		+	
33		+	+			-	
34		-	+	R		+	
35		+	+			+	
36		+	+	R		+	
37		+	+			+	
38		+	+			+	
39		+	+			+	
40		+	+	R		-	
41		+	+			-	
42		+	+			+	
43		+	+			+	
44		+	-			+	
45		+	+			+	
46		+	+			+	
47		+	+			+	
48		+	+		R	+	
49		+	+		R	+	
50		+	+	R		+	

	Lac	Mal	MH	S	TI	X	Gal
51	+	+	+	R		+	+
52		+	+			+	
53		+	+			+	
54		+	+			+	
55		+	+			+	
56		+	+	R		+	
57		+	+	R		+	
58		+	+			+	
59		+	+			+	
60		+	+			+	
61		+	+			+	
62		+	+	R		+	
63		+	+	R		+	
64		+	+	R		-	
65		+	+			+	
66		+	+			+	
67		+	+			+	
68		+	+			+	
69		+	+			+	
70		+	+	R		+	
71		-	+			+	
72		+	+			+	
73		+	+			+	
74		+	+			+	
75		+	+	R		+	
76		+	+			+	
77		+	+			+	
78		+	+			+	
79		+	+			+	
80		-	+	R	R	+	
81		+	+			+	
82		+	-			+	
83		+	+			+	
84		+	+	R	R	+	
85		-	+			+	
86		+	+			+	
87		+	+			+	
88		+	+			+	
89		+	+			+	
90		+	+			+	
91		+	+			+	
92		+	+			+	
93		-	+			+	
94		+	+			+	
95		+	+	R		+	
96		+	+			+	
97		+	+			+	
98		+	+			+	
99		+	+			+	
100		+	+			+	

Repeat M- conjugations
 W1111 x W1632.

852 b.

July 17, 1951.

Cross on D: MBB, (sm) agar.

Test "prototypes" by plate transfer to D: BB, (sm)

Tests very clear.

Plate	M-	Total
1	8	101
2	24 1	27
3	4	84
4	2	85
5	5	87
	<hr/> 20	<hr/> 384

Also pick 20 at random for M+. Pickups on EM B lac.

Pick to EM B.

M- 7 Lac+ 13 Lac-

M+ 3 + 17 -

not significantly different.
 M- grew very well in plating
 media.

✓ Isolates: M- all grew on BMB, not BB. ✓
 M+ " " " " ✓

on EM B Maltose M- 3 Malt+ (also lact+) others -
 M+ all 17 -

Selections, possible vac -

852 C

	lac	Mal	S	MH	TI	Gal	Xyl
1	-	+	S	-	S	+	
2	+	+	S	+	R	+	
3	-	+	S	+	R	+	
4	+	+	R ^{lac}	+	R	+	
5	- ⁺	+	R	+	R	+	
6	- ⁺	+	S	+	R	+	
7	-	+	R	+	R	+	
8	-	+	S	+	S	+	
9	-	+	S	+	S	+	
10	-	+	S	+	S	+	
11	-	+	S	+	S	+	
12	+	+	S	+	S	+	
13	-	+	S	-	S	+	
14	+	+	R ^{lac}	+	S	+	
15	-	+	R	+	R	+	
16	+	-	R ^{lac}	+	R	+	
17	-	-	R	+	S	+	
18	-	+	S	+	S	+	
19	-	+	S	+	S	+	
20	-	+	S	+	S	+	
21	- ⁺	+	S	+	R	+	
22	+	+	R	+	R	+	
23	-	+	R	-	R	+	
24	+	+	R	+	S	+	
25	+	+	S	+	S	+	
26	-	-	R	-	R	+	
27	-	+	R	+	S	+	
28	-	+	S	+	S	+	
29	-	+	S	+	S	+	
30	+	+	R	+	S	+	
31	-	-	S	-	S	+	
32	+	+	S	+	S	+	
33	-	+	S	+	S	+	
34	+	+	S	+	S	+	
35	-	+	R	-	S	+	
36	-	+	R	+	S	+	
37	+	+	R	+	S	+	
38	-	+	R	+	S	+	
39	+	+	R	+	S	+	
40	-	+	S	+	R	+	
41	+	+	R	+	S	+	
42	+	+	S	+	R	+	
43	-	+	S	+	R	+	
44	+	+	S	+	R	+	
45	- ⁺	+	R	+	S	+	
46	+	+	R	+	S	+	
47	- ⁺	+	R	+	R	+	
48	+	+	S	+	R	+	

Counted 852C Lar - only.

852C'

Mal	S	M/R	TI
+	S	-	S
+	S	+	R
+	R	+	R
+	S	+	S
+	S	+	S
+	S	+	S
+	S	+	S
+	S	-	S
+	S	+	R
-	R	+	S
-	R	-	R
-	S	-	S
+	S	+	S
+	R	-	S
+	S	+	S
+	S	+	S
+	S	+	S
+	S	+	R
+	S	+	R

Selected isolations

852
D

June 29, 1951.

Cross B EM⁺lac. Point 3 plates to L175 lac⁻, 174, 17H for

specific selections. #1, 2 rather mixed. (use kind support surface)

#3 OK.

Pick all ^{lac-}Mal⁻ and MH⁻ from #3 plates (including overlaps).

Count. Lac: 10 - 163 + / 173

Mal ~~29~~ 30 } 7? overlaps
MH 21 }

lac-MH 1? ov.
lac-Mal 0?

SR : 27 : 8 Mal⁺
19 Mal⁻

From the interaction of the - lac, MH, Mal selections, in comparison with unselected sets of B, possible linkage relationships may be detected.

Summary

Lac	Mal	S	MHE	TI	A	B	Adj
-	+	R	+	R			1 1
-	+	R	+	S	1 11		
-	+	R	-	R	3 2 11		
-	+	R	-	S	4 3 11		1 1
-	+	S	+	R		1111	4 3
-	+	S	+	S		1111 III	7 6
-	+	S	-	R		11	2 2
-	-	R	+	R	3 2 11		
-	-	R	+	S			1 1
-	-	R	-	R	22 11 1111 1111 1111 1		1 1
-	-	R	-	S	36 26 1111 1111 1111 1111 1		1 1
-	-	S	+	R			1 1
-	-	S	+	S	1 11		
-	-	S	-	S	2 1 1		
-	-	S	-	S			
+	+	R	+	R		11	2 2
+	+	R	+	S	1 2 11	111 111 111 1	16 13 13
+	+	R	-	R			
+	+	R	-	S	1 2 11		5 4 5
+	+	S	+	R	1 1 1	111	
+	+	S	+	S	1 16 111 111 111 1	111 111 111 111 111 111 1	
+	+	S	-	R	1 1 1		8 7
+	+	S	-	S	1 2 11	111 111	
+	-	R	+	R	1 1 1	1	1 1
+	-	R	+	S	1 1 1	111 1	6 5
+	-	R	-	R	1 1 1		
+	-	R	-	S	1 1 1		3
+	-	S	+	R	11 19 111 111 111 111 111	111	
+	-	S	+	S	1 1 1	11	2
+	-	S	-	R	1 1 1	111	3

1111 1111 1111 1111 1111

1000

3

Adjusted (to 100)
Summary used
in CSH 1951

f-1
210

lac Mal S MHC TI

-	+	R	+	R		1
-	+	R	+	S	1	
-	+	R	-	R	3	
-	+	R	-	S	4	1
-	+	S	+	R		3
-	+	S	+	S		6
-	+	S	-	R		
-	+	S	-	S		2
-	-	R	+	R	3	
-	-	R	+	S		1
-	-	R	-	R	22	1
-	-	R	-	S	36	
-	-	S	+	R		
-	-	S	+	S		
-	-	S	-	R	1	
-	-	S	-	S	2	
+	+	R	+	R		2
+	+	R	+	S	1	13
+	+	R	-	R		
+	+	R	-	S	1	
+	+	S	+	R	1	4
+	+	S	+	S	9	45
+	+	S	-	R	1	
+	+	S	-	S		7
+	-	R	+	R	1	1
+	-	R	+	S	1	5
+	-	R	-	R	1	
+	-	R	-	S	11	3
+	-	S	+	R		
+	-	S	+	S		2
+	-	S	-	R		
+	-	S	-	S	1	3

*

*

A	% + S	28	21	15	17	67
B	"	85	84	72	73	88

July 3, 1951.

A W1177 x 1632 Cross on 1. D(10) + BMB, sm for
B W1635 x 1632 segregation of M+/-

July 7. B' B, grown together 1:5 4 hours 2. EMS Lac for linkage data.
n D(10)

July 3: A: numerous colonies on BMB, sm.
B 1 " 6 plates!

Repeat B, B'. July 6.

A. Test, by velvet transfer, M segregation from A.
Not all scoreable.

3. 83/83 M+
2. 7/48 M- } 12 M- ✓
1. 4/62 M- } all are

do isolate 12 M+ ✓) 14 M-.

B' ⁹/₁₀ colonies. Test on +, - M.

all are M+.

M is +

both TL
and
- 5 -

Suggestion: As "B", is Lac linked to TL?

Test by crossing BMS^R x TL^S

on TL sm agar
and studying segregation
of TL, Lac.

(see over)

Lac character of M+, M- selection

B'	M+	6 Lact	2 Lac-
A	M+	12:	3 Lact 9 Lac-
	M-	11:	2 Lact 9 Lac-

Linkage of Lac to M? Not supported by these data.

July 6, 1951.

W1634 (Cellulose-fermenter). Irradiated 8 secs on EMB lac
25 x 50 = 1250 colonies.

Some sectorials undoubtedly ignored. Many morphological sectorial cols.

Spot Lac- on EMB Glu for further purification.

By transfer tests:

13 lac- all are Sucrose Cellulose Xylose Galactose Maltose
and glucose positive

Save 1. lac- 854 1

July 12, 1951.

20 x 50 = 1000 on EMB glucose No definite mutants
15 x 40 = 600 " " " " "

7/16. Check Vaughn's cultures 776-835, 837, 839, 840, 841 1-5
s.o., EMB glu, cell, suc, lac

7/17. all are ++ or +. #1,3 probably most generally suitable
= 835 (Vaughn 129) = W 1647
= 839 (Vaughn 168) = W 1648

7/17. a 1634
b 1647 - uv resistant
c 1648

854-(2-4)

a. 3 possible flu- from 12 x 400 = ca 5000 ✓

b. 1 " " 8 x 100 = 800 ✓ (+ second try will!)

c. 1 " " = 854-5

All 4 mutants grow rather poorly. Are negative on
galactose, glucose, xylose, mannitol, maltose, lactose, sucrose and
cellulose

see over

(1648??)

W1647. UV

1 Glu - mut on 5 plates, variably crowded

7 Lac - or ±

①. Glu - Cl - Suc + Mal - Lac -
Xyl + Gal +

Designation of W1647 almost certainly
correct, but conceivably was
substituted ~~for~~ W1648. Designate
1647a for immediate source of Glu - mutant
(W1677)

Penicillin effects on K12

July 8, 1951.

24hr. K12 culture .5 / 10 ml fresh Penmassay + Penicillin as indicated.

- 1. No penicillin
- 2. 1 unit / ml
- 3. 5 units. / ml

* P9. Filter 1, then 2; 3 then W1177 (4) as controls.

	1	2	3	4
A. Plate .1 ml on EM3 lac	✓st	st 1+	1 col. lac+	
B. " .1 ml with W1177 on EM3 EMS lac			2 minute lac-	
C. Inoc. 10 ml Penmassay	✓st	✓st	Turbid	Turbid
D. " " + W1177.	st	1 minute lac-, r?		
AA etc. same but N10. (W1177)	probable leak			
AA	++	++		
BB (W677)	++	++		
CC	2 lact		st.	
DD	7 lact	7 lact	st	

Possible "protogenic"? in 3B, 2D
restreak in D(0)

This experiment n.g. :
leaky jacket & filter
used for 1 and 2

With 1-5 units, no morphological effects were noted.

Penicillin effects.

July 11, 1951.

P12.		K12, 1ml + penicillin in Penassay tubes	P12.
1	A 0	++	+++
2	B 50	++	Normal size + motility.
3	C 50	++	many filaments, some white granules
4.	D 1200/ml	±	inhibited growth.

Filter 1, 2, 3 through separate 2-4 lb Mandel filters. Pass turbid (ca 10^8) W1177 broth culture through sterility. Plate this filtrate on EMBAc (= E) and inoculate 1ml into Penassay (= F).

		1	2	3
A.	Inoculate filtrates 1ml, Penassay	-	-	K12 only ++
B	" " " " + W1177. Plate after 24h.	0	0	++ K12
C	Plate " .1 " EMBAc	0	0	0
D	" " " EMS Lac + W1177	0	0	0
E		0	0	0
F		-	K12 only ++	K12 only ++

A few particles per ml ~~survive~~ survive filtration, if penicillin has been added. No unique genetic activity is noted. However, higher titres of FA, and heat-inactivation are needed to parallel Salmonella findings. W1177 control satisfactory.

Also plate filtrates into serum agar (+ penicillin?)

F: streak out on EMBAc + or - serum to find W1177.

75a Penicillin K12 10% inoculum. 24h. 37°
" R.T.

- A. Sediment and Filter supernatant. Mandler 816 filter
B. Follow with untreated W1177 $>10^8$ (10^7 /ml) control.

1. Plate A on serum agar; 0.1ml
- 2 " " nutrient " "
- 3 " " EMB lac 0.1 "
- 4 Proc 1ml samples into Penassay.
- 5 " " " " + .5ml W1177
- 6 Plate B on EMB lac ser
- 7 Proc 1ml Penassay.

Conclusion: Filter not satisfactory. 6 showed ca 50 lac-
1,2,3 each showed ca 400-500 (lac+). 4 all turbid.

Refilter refrigerated suspensions, 1416 Mandler repeats
all sterile. No prototrophs from 5, 7

Conclusion: Penicillin may result in slight
inactivation & lethality of K-12, but no consistent
effects noted. No genetic effects whatever noted.

5. showed ^{minute} small colonies on EMS lac after 3-5 days
 Pick and streak out on EMS ~~lac~~ lac, D/O. - 3 days: no growth on ^{latter}
 [Probably T-L+ reversion growing as carryover]. not as former.
 8 Accor W1649 + 85Tb 1ml in Penassay.
 9 " W1632 " " " " Wash + plate heavily EMS lac
 No colonies

 #

Conclusions: No evidence of genetic activity in penicillin treatments of K12.

F2 Crosses

July 6, 1951.

Cross in D TLB, sm W677x

A W1367

No yield

~~See 889~~

B W1362

No yield.

C W1490

2 colonies / 2 plates - 1 grew out: B, -

D W1368 x W677

D' (do., grown together 6-8 hours).

7/17 D: Transfer plates to D (sm TLB₁), EMS Mal sm TB₁.
 Only 1 out of about 60 grew on EMBB₁.

~~D' Most grew on EMBB₁, few on TLB₁.! (paste mixing of plates)~~

Strain out auxotrophs on EMB lac to purify for further characterization. 8 saved.

#1 - T-L-B₁ - S^R Lac-

W1649.

all 8 are T-L-(B₁-) S^R lac - Mal - Xyl - MHE - Gal -

(S^R Mutant ??)

7/19/51. E. W1649 x ~~W1638~~ W1632 on EMS lac, Mal, MHE.

EMS Lac	95%+
Mal	" +
MHE	90%+

Hybrid direct recombinants also give aberrant linkage results.

Material from 853 W1635 x W1632.
852

EMSMR 41- 359 total

Strobes from D10) to EMS lac.

Also collect lac- from EMS lac plates

Repurify all lac-.

Plate count
 EMS lac. 18 - : 187+
 MH 19 - : 55 total
 From D10) 174 - : 245 total
 strobes to
 EMS lac

Read points from lac+ and from lac- separately.

- Transfer all lac+ to MH, Mal.
- Transfer 50 lac+ also to Xyl, T6, T1
- Transfer 28 lac- to MH Mal T6 T1 Xyl

3/16/52

Embryonic? : lac, Mal are independent

LAC +	Laz	Mal	MHL	EMB Xyl	S	T1	T6
1			-	-		R	R
2			-	-			R
3							R
4							R
5		-		±	R -		R
6			-	-	R -	R	R
7							R
8							R
9		-	-	±	R -		R
10			-	-			R
11							25 R
12							26 R
13					R +		27 R
14							28 R
15			-				29 R
16			-	-			30 R
17					R -		31 R
18							32 R
19							33 R
20							34 R
21						R	3 R
22	-						
23	-	-	(+)	±	R -		
24				-			35 R
25		-		-			36 R
26		-	-				
27			-				
28			-				
29			-				
30			-				
31			-	-			
32							
33		-	-	-	R -		
34							
35							
36		-	-	-	R -		
37							
38							
39							
40				±	R -		
41			±		R -		
42							
43		+	-	(+)?	R -		
44							
45				-			
46			-				
47							
48							
49							
50							

Malon
EMB

only few lac - 06 crosses.

ALL EXCEPT
13 MAL-

2

	LAC	MAL	MTL	EMB XYL	EMS S	LAC	T ₁	T ₆
51								
52			-					
53		-	-	±	R ⁻	R	•	R
54					R(s) ⁻	R	•	R
55	-	±				<u>R</u>		
56								
57								
58								
59								
60								
61					R ⁺			
62	+	-	?	-	R ⁻			
63		-						
64								
65								
66		-				R	•	R
67		-	-	-	R ⁻			
68								
69			-	-				
70					R ⁺			
71	?		-	-				
72								
73								
74				-		R ^{unc}		
75	-		+ Mucoid	-	R ⁻	R^{unc}		
76		-	-	-				
77			-	-				
78			-	-				
79						<u>R</u>		
80	-							
81	?					R	•	R
82								
83		-	-	-	R ⁻			
84	+							
85			-	-				
86						R	•	R
87								
88								
89								
90			-	-	R ⁻			
91			-					
92								
93								
94		-	-	-				
95	-		-	-				
96								
97								
98								
99								
100			-		R ⁻			

3

	LAC	MAL	MTL	EMB XYL	EMS LAC S R ⁺	T ₁ (R)	T ₆
101							
102							
103	-						
104							
105							
106							
107							
108				+			
109							
110							
111							
112							
113							
114			-				
115							
116							
117							
118			-	-			
119							
120							
121							
122					R ⁺		
123							
124							
125							
126							
127			-+	-+			
128							
129							
130							
131		-					
132							
133			-				
134							
135		-	-	-	R ⁻		
136		-	-	-	R ⁻		
137							
138							
139							
140							
141						R	.. R
142							
143							
144							
145							
146	-						
147		-					
148				+			
149					R ⁺	R	.. R
150							

4

	LAC	MAL	MTL	XYL	EMS LAC S	T ₁	T ₆
151							
152		-	-	±			
153							
154			-	-			
155						R	• R
156			-				
157							
158		-+			R ⁺		
159							
160		-					
161			-	-	R ⁺		
162							
163						R ^{lac-}	
164							
165			-	-+			
166							
167							
168	-+						
169							
170	-	-+				R ^{lac-}	
171			-	-		R	• R
172			-	-			
173							
174							
175			-+	-+			
176							
177	•						
178							
179							
180			-	-+			
181							
182							
183							
184							
185		+	-	-			
186							
187	-			-	R ^{lac-}		
188			-	-			
189						R	• R
190							
191							
192							
193		-	-	-	R ⁺		
194						R	• S
195							
196							
197							
198	-						
199							
200			-	-			

5.

	LAC	MAL	MTL	XyL	EMS LAC S	T ₁	T ₆
201							
202		+			R ⁻		
203			-	+			
204							
205	- ✓					<u>R</u>	
206					R ⁻		
207							
208							
209			-	+			
210							
211			-	-			
212		-	-	-			
213							
214							
215						R lac ⁻	
216			-	-			
217	+ ✓						
218							
219					R ⁻		
220							
221							
222					R ⁻	R	✓ R
223		+			R ⁻		
224							
225							
226							
227						R lac ⁺	
228							
229		-	+		R ⁻	R	✓ R
230					R ⁺		
231					R ⁺		
232							
233		-			R ⁻		
234							
235							
236							
237			-				
238					R ⁺		
239		-	-		R ⁺		
240	+ ✓			(+)	R ⁻	R lac ⁻	
241							
242							
243					R ⁻	R lac ⁻	
244							
245							
246							
247							
248							
249							
250							

	LAC	MAL	MTL	XyL	S	T ₁	T ₆	
16								
1						R	S	37
2			-	-	R MAL+		S	
3						R	S	
4						R	S	
5							S	
6							R	42
7						R	S	
8							S	
9							S	
10							S	
11						R R	S	4
12						R	S	
13							S	
14		-	-	-			S	
15		-					S	
16		-	-	-	R -		S	52
17							S	
18							R	
19						R	S	
20			-	-			S	5
21					R MAL+		R	51
22						R R	S	52
23						R R	R	53
24						R	S	54
25							S	55
26		-	-	-	R -		S	56
27			-	-			S	57
28							R	58

Lact

Lac -

July 10, 1951

P10

1. Inoc. 1 colony W-1 to Penmassay, USA slant
~~845~~ spread .1 and (0.1) ml on EMB lac. Assay .1 ml @ T1.
930 A12

4P Print to ± T1 agar.

A13: Read:

- a 46 clones; 31 singles; (1 plate) 21 singles (second plate).
- b 10 clones; 6 singles (on plate only)

Pick from b sites to fresh broth.

spread .001 ml; streak. show 4 P11 - 1030 07
Print 10²⁰ P11.

A14 .001 ml: 15 clones 13 singles ①
5 " ②

③ Pick 5 sites.

N15 Plate at various dilutions

Shows incubation:

Print to T1.

- Assays of successive isolates.
110 .1 ml 91.5 mic.
25 x 10⁶ 1 ml ca 500.
310 .01 ml ca 1000.
411 38/582

A streak: 1 clone in nearly confluent portion. Restreak directly,
also streaking and compare cross-streak with T1, as against
random isolated colonies. Pick 1 clone = 1

B 10⁻⁵ Semi confluent. 9 clones; no singles.
Pick 3 clones = 2-4 streakout
pool to Penmassay.

See over:

1. Random tests unstruck colonies. / T1.

1.	10 S.	0 R
2.	"	"
3.	25 S	5 R
4.	8 S	2 R

∴ successful isolation of V_1^R by indirect selection

2. Dilute 858-4 10^{-8} plate .1 ml. 3 plates.

#2 unstruck - too weak. Purify T1 after 9 hours (purify colonies)

1.	12 V_1^R	301	total
3.	16	261	"

38 / 562

Pick best isolated homologous colonies.

✓ each of 4 V_1^R
Random V_1^S .

Test against T1 and replate to purify.

Store single colony as 858-5...
Inoc + transfer broth to test stability

Note In final plating a few plaques of contaminant phage (identical as T1) were found. However, no evidence of T1 was found in the broth series, or in selection line. It seems unlikely that this could interfere with achieved result. However, see 859 for repetition of general expt.

Gradient Selection: W-112 / T1.

July 17, 1951.

Inoculate single colony broth cultures 11 AM. = 859-1

P17 Plate EMBA₂ ca 5 hours. Print to T1 plates
0.1 ml 5 clones; Total: 16; 12

N18. Pick clones to Pennassay. 3-8 PM. ^B 10⁻², ^C 10⁻³, 10⁻⁴ ml plated on EMBA₂
8 PM Print to T1 plates

A19. A. 23 clones, 1-3 singles. Pool clones from
B 1 clone 2 singles B and C to
C 1 clone 1 single. Pennassay. 859-3

N19. 2 hour tube (probably ca 3 x 10⁸) plated out on EMBA₂
12 N19 - 10⁻³ 10⁻⁴ 10⁻⁵
A B C

9³⁰ A20. C: (ca 600 cells) No V, R

B: 2 clones; 1, 0 singles } Pool 2 from B and
A: 16 clones; 2, 0 singles. } 2 from A to Pennassay
859-4.

10A21. 10⁻⁶ dilutions. 3 plates. After 1 hour, single replica transfer
A Strains out 4 of the V, R.
B 12 mutants; 793 total Test against T1. V, R ✓
C Place isolated V, R in hand.
Inoculate serial transfers in both, 5A, B,

858 and 859. Each transferred serially
through 10 5ml tubes Remassay
with loopful inocula.

8/3/51. Plate out on EMB glu. Replica to EMBLac. T1.
all colonies were V₁R. W1485 control lysed on T1 plates.

Total counts:

858
<hr/>
290
277
269
<hr/>
836

859.
<hr/>
131
150
166
<hr/>
447

Replica Efficiency

860

July 24, 1951.

A. Compare fresh and old growth for clones.

1. Inoc 58-161 MEMB (24) 9 AM.

2. Spread " " " 3:15 PM.

325. Replica to T1 agar, pairs.

B. Dilute 58-161 10^{-5} Spread .1 ml, let dry.

① Replica to series of plain agar.

② Assay 10^{-7} ml for count.

③ As ①, but 10^{-5} ml original

Dilution	Original	Replica (removed)							
		1	2	3	4	5	6	7	
② 10^{-7}	0	99	91	7	1				
1 10^{-6}	(1000)	$\pm 10^3$	42	19	12	10	6		
3 10^{-5}	(10,000)	$\neq ca 10^4$	331	171	108	109	87		

\therefore About 10-25% of initial cells are removed.

5-10% deposited on first replica, with indicated "decay" thereafter.

8/30/61 - see experiments with E. Klein on efficiency of colony transfer.

Singlets and clones
Replica plating

861

July 25, 1951.

2¹⁵ P.M.

W112 (859-1)

1a. 1 ml + T1 assay 190

b. 0.2 ml + T1 assay. 90 ~~78~~

2a. ~~0.2 ml~~ .1 ml Replica to 2 T1 plates immediately

b .02 " " " "

3a .1 " " after incubation 2¹⁵ -

b .02 " "

2a	15	4	0) no sign of clones.
2b	2	0	6	

3a	73	65	53) many clones.
3b	14	10	5	

2bb 11 8 4, 2 replica swabs from 2b after incubation.

Use 2bb and 3b for published figures of replicated clones

7/27. Nonclonal replica.

c1 .1 ml w-1 culture transfer (EMZ) as initial

c2 1 ml (sed + conc to .1 ml → 10:1)

July 28, 1951.

Use technique of (respiring microcolony)

2-wk old culture H226. 10^{-6} dil.

Control: .1 ml spread over plate; .01 ml in line
exptl. .01 ml in line 11:30 AM. Respired after 7 1/2 hours
mostly segregated on H.S. 7 PM.
EMBS Lac.

Repeat, new H226. 7/29. Mostly segregated. By replica, almost all
lac- are prototrophs. ^{1 aux. lary} noted = 862-1
24868

7/30. New H226. (Hyophil) 10^{-6} .01 ml.
from single colony. 10 AM. - 3 PM.
5 hours.

Control plating .1 ml: 166 23

.01 ml brush. 29 4

Probably 2x too many cells per brush, but many likely

From replica plates, ^{most} ~~some~~ lac- are auxotrophic; ~~most prototrophic~~

Replica lac- "segregate" selection to EMS Lac

Clones

- 1 4 v 5- v v v -----
- 2 6 v
- 3 w - - - - v - w -
- 4 w v v v v v v v v v
- 5 v v v v v v v v v v
- 6 v v v v v v v v
- 7 v v v v v v v v
- 8 v v v v v v v v v v
- 9 v v v v v v v v
- 10 v v v v v v v v
- 11 v v v v v v v v
- 12 v v v v w v v v
- 13 w w v
- 14 w w w v v v v
- 15 w w w w w v v v
- 16 w w v v v v v v v v
- 17 w - - v v v v
- 18 v v v v v v v v
- 19 v v v v v v v v
- 20 v v
- 21 v v v v v v v v
- 22 v v v v v v v v
- 23 v v v v v v v v
- 24 v v v v v v v v
- 25 v v v v v v v v
- 26 v v
- 27 w w w w w w w w
- 28 v v v v v v v v

- 29 VVVV -W V
 30 V V W V - V V V V V V V
 31 V V -W V
 32 WW W -
 33 VVV V W
 34 VV V
 35 WW V V
 36 V V
 37 W W WW V
 38 W W W V
 39 V V
 40 WWV - WW-
 41 W V-
 42 WWW
 43 W-
 44 V WW W
 45 W -W
 46 ~~46~~ ^WW ^WV V W W V W W V
 47 ~~WWWWWW~~
 48 W ^WW W V V V V V V
 49 W W W - - - W V
 50 VVV W V V W V
 51 W ^W- W - - - V V V W
 52 V V V -
 53 ^WV V W
 54 V - -
 55 VVV - -

56 v v v v v

57 $\overline{v}v$ v \overline{v} vv

58 vvov

59 ov

— clares

15, 7, 24, 10, 17, 16, 15, 21, 19, 21, 14

many mixed clares.

Pick possible new —
from vacv clares as best possible. Pick to EMBS Mal

all picked were Mal+.

52 Mal_v, phototrophic

all others anoxygenic.

all δ gl⁺ except 29, 57.

Probably most of the Lac- were previous segregants

see 867

July 30, 1951

A. K1 QT-h- }
B. K1 h2-arg } p^{ts.}

Add put to all plates, EMS Lac
1 plate per x.

1. A x W-1
2. B x W-1
3. A x H245
4. B x H245
5. A x H290
6. A x W1606 (as DM-)

K1 QT-h-
= put-multicam-hist-
K1 h2-arg
= put hist arg.

8/2/51. all plates barren. Other crosses with
W1606, H245, H290 were fully fertile.

Maas says put + reversions are futile.

July 29, 1951.

- Inactivate on EMB agar 6-20 sec.
- Inactivate washed suspensions 2... 20 sec. inoc
1/2 ml into Punnassay. Use 4 sec treatment

7/30 Plate on EMB lac

ca 300 tested for auxotrophy by replica: all X+.

No lac, or Mal mutants on ca 20 plates. Many colonies
showing morphological characteristics.

ca 300 more tests for auxotrophs. ~~1 doubtful: revertants~~

10 x 350 No auxotrophs detected!
= 3500.

Total, ca. 4000 tests!

Plate on EMB/Mal sim. [E. Woodworth records show W1647].

Variable types, some Mal±. All Lac+.

$\frac{1}{2}$
 $\frac{2}{3}$

9/20/51. Problem adopted by E. Cahn.

W1606 x H293; W1675 x H290
S^D x S^R

~~865~~
865

Mostly lact. Replica to EMSlac, EMSlac sm. for S^S.
ca ~~200~~ 200 colonies - grew primarily on
EMS Lac, ± sm.

Aug. 7, 1957 ~~W1606 x~~

B W1675 x H290.

C. W1675 x H294

on EMSlac, ± sm

TL lac-Met-S^D M-lac^rMalv S^S

8/5/51

"1675" proved to be BM-lac^rMet S^D
(probably W1606)

Losses n.g.

July 30, 1951.

M290 x M293

M-Lac^vMal⁻ - ~~TFL~~
^S lac^vMal⁺S^R

a. EMS Lac

b. EMS Mal

a. 20 picked.

11 Lac^v. Others lac⁻ (Lac^v.)

b " "

8 Mal⁻
~~7~~ " + "
 3 Mal^v?
 2

Test ~~the~~ single colonies on
 EMS Lac; EMS Mal / ser.

~~Mal~~

	Mal	S	#
b.	-	S	8
	+	R	5
	Mal ^v	V	2?
	+	S	4
a.	-	S	9
	+	R	1

Mal^v relatively infrequent
 in this cross.

Grow 866-1 on D(Lac) for wiser platings
 on EMS Mal.

Diploid microcolony segregation

August 2, 1954.

See 862 .01 ml; 3×10^{-6} per plate 11³⁰ -

assay	.1 ml	EMB lac		Mal			V
		v	-		+	-	
		14	6		1	0	23
		9	3		9	1	26
							11 ³⁰ - SPM.

ca 200/ml. About 2 clones / plate.

Agar with 2-3 ~~in~~ streaks not respnd.

On EMB lac 6 plates. EMB/Mal 3 plates

clones:	-	v	
		1	1v
	1	1	1v
		1	1+
		1 ¹⁰	1v (+, - also!)
		no segregants	1 mixed v, +.

No useful data but method is substantiated.

8/5/51. H226 A) .01 ml 10^{-6} B) .01 ml 3×10^{-7}

D lac count; many -. (assay: 14v, 17-)

5 readable lac v clones, ca 10%. 2 possible segregants, but crowded.
Streaked EMB Mal. (Both Malt!)

8/6/51. As above. Total 25 plates, mostly too crowded. Mostly lac v
E 47 possible useful clones (5-20) altogether. (5 1/2 h. inc.)
Pick lac- to EMB/Mal. 12: all Malt. (2 Mal v)

8/7/51 F 4 1/2 h. inc. ~~long~~ standard long method - Few colonies
should use 2x. Many lac-!
(over)

G, H.

UV 20 sec.

ca 50% swr.

Results inconclusive. Increase in
Mat~~+~~ in both intact and exposed.

Method probably is unsound.

July 30, 1957 FT

A. H226, grown in D(Lac), no treatment. Auxotrophy noted in replica platings. Nutrients, Mal, S tested in such platings.

H267, grown D(Lac). UV 30 sec. Grew heavily in D(Lac) +

B. B7

C. TLB₁.

Plate on EMB lac, Replica tests for auxotrophs.

B gave numerous auxotroph lac⁺; C very few.

Type		Nutr	lac	Mal	Mal S	S	Mal S	Mal S
A.	1	M	v	v	v	S	S	
	2	TL	v		+v	S	S	} H245 type
	3	TL	v		+v	S	S	
								H294
B	3	M	-	-	-	-	-	R
	4	TL	-	-	-	v	-	R
	5	M	+	-	-	S	-	R
	6	M	v	v	S	v	-	R
	7	M	+	+	S	v	-	R
	8	TL	v	+	S	v	-	R
	9	M	-	-	-	-	-	R
	10	+	+	-	-	-	-	R
	11	TL	-	-	-	-	-	R
	12	M	-	-	-	-	-	R
	13	M	-v	-	S	S	-	R
	14	M	+	+	S	(v)(+)	-	R
	15	MTL	+	v	-	v	-	R
	16	M	+	-	-	-	-	R
	17	M	-	-	-	-	-	R
	18	TL	-	-	-	-	-	R
	19	M	v	-	-	-	-	R
	20	+	-	-	-	-	-	R
	21	M	-	-	-	-	-	R
	22	MTL	+	-	S	(+)	-	R
	23	M	+	-	-	-	-	R
	23	M	+	-	-	-	-	R
C	1	TL	v	-	-	v	v	v
	2	TL	+	-	-	v	v	v
	3	TL	+	-	-	v	v	v

		Mal	S	
A1	M	v?	s	H294
A2	TL	v	s	H295
B1	M	-	R	H291
B2	TL	-	v	
B3	M	v	v	
B4	M	v	s	(H294 type)
C1	TL	v	v	

August 3, 1951.

more 1 drop culture of phage T3 (10^{-3} lysate) to 5ml permassay
incubate overnight.

A
8/3

- 1 Y44
- 2 Y51
- 3 Y44+Y51
- 4 Y44+T3
- 5 Y51 "
- 6 Y44+Y51 + "

clear
"]

#1, 2, 3, 6 plated. all grew without
plaques on EMB.
No protoplasts after 72 hours.

Sediment turbid cultures. Plate on D(0)

Repeat A, with heavier bacterial inocula (ca. 4ml)

1-3 turbid. 4-6 clear. streak out 4-6. Also W1664/3
4-6 remained clear! passages
No true 1664/3 found as survivors (physical tests)

W1663/3 obtained from platings on EMB. = W1679.

Purify by 3 subcultures on EMB and recheck ✓.

- 8/12.
- 1 1679
 - 2 " + T3 ±
 - 3 1679+1665
 - 4 1665
 - 5 1665+T3 Clear.
 - 6 1679+1665+T3. ± turbidity.
- Protoplasts
1 Sediment. Also
2 innoculate 2 and 6
3 into Perm assay.

- a) The parents are not completely stable
- b) No detectable effect. The smears had numerous
○ (L?) forms.

See Notes by E. Cahn

August 29 (ff.) 1957.

a) Diauxotrophs from PF9 (meth-) and PF12 (Leuc SR).

UV 40 sec., resuspended cultures in water from meth. ~~Leuc~~ ^{Leuc in both}
 Wash, noc 1:100 in D(10) + meth + leuc + 1000u/ml penicillin,
 aerate overnight. Plate on ± MB base after 10-12 h.,
 replica to D(ML).

PF 9 ca 15% auxotrophs.

PF 12 ca 5% .

Replicate to EM3 for Recheck.

On Recheck:

PF9. 7/8 and 37/40 = 44/48 OK as auxotroph

PF12 14/20 OK.

See 870a. for Random series.

b) Plate irradiated suspensions from (a) at ↑. PF9, PF12 and
 PF9+12 (uv, grown together). The last showed high counts of prototrophs
 (ca 5000/ml) but moderate counts 500-1000?/ml) were
 seen in the separate cultures also.

Physinia Plate Tests

8709

Group	HC	V. ts	A1	A2	A3	A4	A5	YNA	EMB
PF12	1			+					
	2					+			
	3								
	4						++		
	5				+				
	6					+			
	7				+				
	8					+			
	9				+				
	10				+				
	11				+				
	12					+			
	13				+				
PF9	1			+					
	2					+			
	3				+				
	4				+				
	5				+				
	6				+				
	7				+				
	8						+		
	9				±		+		
	10				+		+		
	11						+		
	12						+		
	13				+				
21						+			
22				+					
23						+			
24				+					
25						+			
26				+				++	
27				+					
28				+					
29			++						
30				+					
31				+					
32				+					
33				+					
41				+			±	++	
42				+				++	
43				+					
44			++	+				+	
45				+					
46				+		±			
47				+		±			
48				+		+			
49				+					
50				+					
53				+					
51									

all but 53

None

See over

±

ERRATIC

all A3 - Rechecked by replica:
all use Trypt. -

Discard these, and all A2 - (presumably added.)
Isol or Val or IV

and PF 26.

Keep other PF numbers and preserve in dry tubes.

September 24, 1951.

PF 12:	19	--- L+	A2	iso	--- Val	--- I.V.	---	0	iso-Val	
	20	A3	A3	del	typ	typ.	del+typ	0	Typ	
	21		HCV	Y ₄	RNA	NZ.		0	??	
	22	A4	A4	hist	thru	glut	red	rop.	0 hist	
PF 9:	23	A2	M+	A2	L.	IL	Val	I+V.	I+V+L	0 IsoVal
	24	A3	A3	del	typ	typ	del+typ	0	Typ	
	25	A4	A4	hist	thru	glut	red	rop.	0 Hist	
	26	---	A1	Lys	Arg	Cyst	0		Cyst	
	27	---	A1	Lys	Arg	Cyst	0		Cyst	
	28	YNA. M+	YNA.	Pur	Pyz	Guan	Aden	Xanth	Hypx	RNA 0

add new

add 11e. th.

(Guanine)

For crossing, try Meth ^{Cyst} ~~---~~ x Leuc Isol.

PF 26 x PF 19.

These cultures prepared by drying by Hershey's method. Also try new developmental experiments. Add suspension directly to granular or powdered $CaCO_3$ gel, previously washed and sterilized in tubes. * Seal off in air unless otherwise indicated in notes.

9/4-5.

- a) Attempt crosses of PF 26 x 19. 1) Plate 2-4 cultures
- b) Grow together in P. assay; c) UV 0.1 sec. on washed cells, maintain P. assay sep. and together.

(Record of date by registered letter.)

Crossing Attempts with *Pseudomonas fluorescens* 810c

Sept 4, 1951. ff.

see 870b.

PF = 26 x 19.
inc plates 3-4 days.

a. Grown separately.

1/4
1 19
2 26
3 26 + 19

0 0
0

5 v. small (cont??) PF??

b. Grown together

1/5
1 19
2 26
3 (26 + 19)
4 (26) + (19)

0
10!
2
1

4 PF
✓ PF.

c. UV 603; grown together

1 19
2 26
3 (26 + 19)
4 (26) + (19)

v. heavy

0

0

1

2

(prizints, uordic)

Not PF. Papillate colonies!

Not PF

Hold in refrigerator for later study.

1/16/51. Strausout
D(0):

12 } a 3"
2 }
3 } b 2
4 }
5 }
6 }
7 } b 3
8 }

all
SS

Pseudomonas??
to be fed.

9 } b 4
10 }

- d.
1. PF9 = M- 6 5 petri dishes.
 2. PF19 UV L.IV.SR 0
 3. PF19 UV + PF9. 3 large + many small → > 100 petri dishes.
 - 4 = 1+2 4 + (1-2) 9 "

Replica to EMS Inc sm: None SR

This result is rather indecisive. ① Repeat with tests for S^R ...
see 769 ② Survey other substrates for stability. Test PF 2, 3, 4, 7, 8
Use PF 12 = L- S^R x PF 28 = M-Gu-

P. fluorescens

9/21/51. Stability tests on auxotrophs.

Grow on NA. 48 hrs. Harvest, wash and suspend in 3± ml.

Plate 1 ml samples on minimal agar. (probably ca 10^9 cells)

^{Prototrophs}
 PF 2 9
 3 0, same background (leucine?)
 4 0 " " (leucine?)
 6 > 100
 7 > 50
 8 10
 12 21
 28 ca 30 (for Duem + Meth⁻)

Nonauxotrophs of choice are ~~PF1~~, PF3, (4)
 Try x PF28

10/2. PF28 (uv) x PF12. SM

PF12	9	7 ^R 3 ^S
PF28 ^{uv}	0	-
grow together	3	3R
separately.	2	3R

No evidence of recombination.

Replecia to EMS ± sm.

Do PF12 homogeneous?

10/6: Yes!!

Experiment should be repeated

10/3

EMB Lac.
 EMB Lac sm.
 SR.

Stoke culture streaked out on
 Colonies replica plated to
 All of ca 50-100 were

The PF12 S^s may have been contaminants

~~10/7~~ Britate Permasey cultures. (PF-12 from colony on sm.)

Expt. completed ca 10/20. Just as above.

PF12	ca 50	all S ^R	By replicate
" 28 ^{uv}	0		
" 12+28 ^{uv}	ca 50.	all S ^R	"

No evidence of recombination!

Pseudomonas fluorescens
Summary of Crossing Expts.

870s.

10/22/51.

Diauxyotroph combinations: L-IV- M-Cys
PF 19 x 26

PF 26 gave a few prototrophs on dense platings.

Mono-diauxyotroph combinations (5th malbar):

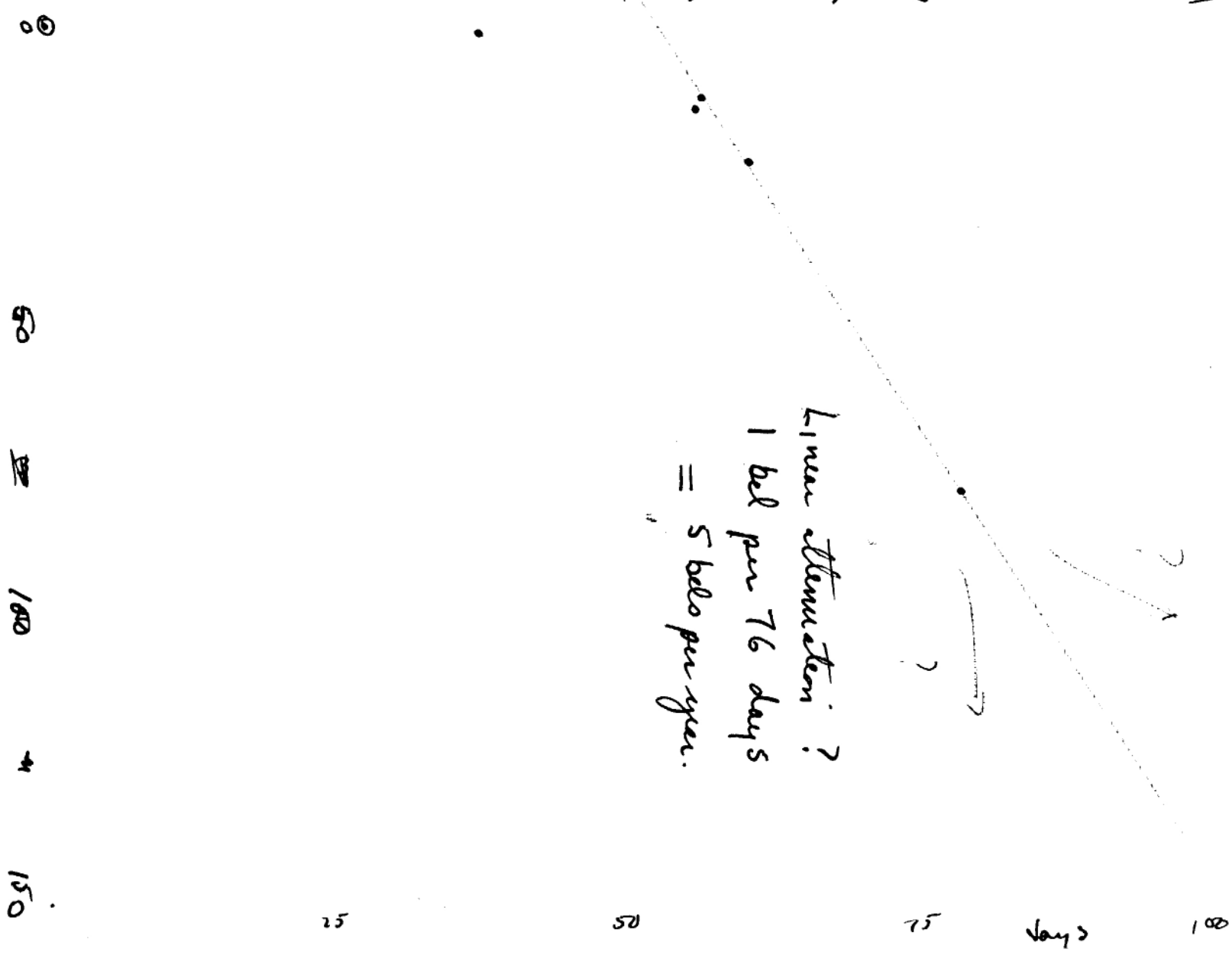
PF 19

Try PF 19 x 28.

4

3 $\log s = -\log s$

Linear approximation?
 1 bel per 76 days
 = 5 bels per year.



September 4, 1951.

a) See typewritten memorandum

9/5. b) Dry K-12 by various methods to compare viability. (F uses heavy suspensions adapted to peptone 20% from NA 5 units).

- | | | | |
|----|---|----------|-----------------------|
| 1. | CaCl ₂ etc. (Hershey, Brown) | | |
| 2. | In contact with granular silica, | air | 2.5 x 10 ⁵ |
| 3. | " | " | vac. |
| 4. | " | powdered | air |
| 5. | " | " | vac. |

9/18/51 contents 1/10ml with

9/4 c) 6. PF 21

1. gran SiO₂ air
2. " SiO₂ vac.

9/5. Open 1, 2, suspended in 10ml Penassay and titrate. (10⁻⁷)

c1: 2

c2: 17

September 30, 1951.

Resume: Survival.

Date	Days	Concentration [10ml]	pS	Count
9/6/51. Initial	0	3.4×10^9	0	340
9/7	1	11×10^7	1.49	11
9/14	12	1.9×10^7	2.25	1; 187/100
9/15	14	2×10^7	2.23	20/10
9/29	23	1.3×10^7	2.42	13/10; 1/1; 80/100.
11/24 11/24	79	2.4×10^5	3.14 4	1/10 17/100 295/1000

Count: $\left. \begin{matrix} 1.1/10 \\ .1/10 \\ .1/10 \end{matrix} \right\} \begin{matrix} .1 \text{ ml} \\ = 1 \times \\ = 10^{-6} \text{ ml} \\ \text{from } 10 \text{ ml} \end{matrix}$

September 6, 1951.

- A. Broth, direct. 1:1K12, 10ml Permaseay, aerated overnight. AS II AI 1
- B. " , sedimented, resuspended in 1ml (1:10)
 .05ml per tube.
1. Gran silica, Air ###
 2. " " Vac ||
 3. Pds Alumina Air
 4. " " Vac
 5. Hooking CaCl₂ - Vac |
- 9/6/51.

Assay B. (.05/10; etc.).

1.1/10	Count	299
.1/10		376
.1/10		675
.1/10		

(= 10⁻⁶ ml sample) original titer 6.7 x 10⁹ cells/ml.
 (from 10ml.)

Each tube received 3.4 x 10⁹ conc. 10:1
 3.4 x 10⁸ dust broth.

Hold in refrigerator assay sample tubes. Empty tubes into 10ml Permaseay.

including washings from walls. Dilution schedule as above:

(.05/10); 1.1/10; .1/10; .1/10. Plate .1

9/11/51.
 812
 B1

	Count	10x	100x	Assay/ml (with 10ml)	Survival (pS)
A 1	2	15	15	2 x 10 ⁵	1.3
A 2	15	121	15	1.5 x 10 ⁶	.3!
B 1	11	161	11	1.1 x 10 ⁷	1.3
B 2	10		10	10 ⁷	1.3
B 3	0		0	0	-
B 4	0		0	0	-
B 5	14	188	14	1.4 x 10 ⁷	1.3
	340.				

Ca 95% "Process Loss", and attenuation in 24 hours.

9/14/51. B1 112 11 187 1.9 x 10⁷ 2.2

9/15/51 B1 85 20 0 ! ~~2.7~~ ~~2.7~~

September 15, 1951.

Harvest from Pirmessay Pastes. Concentrate in acetone 2°
 ca. 30 / 1.5 = 20:1 Use .05ml per sample = 1ml

a) Titrate initial samples (1.1/10; .1/10; .1/10; .1/plate) ^{initial}

Silica tubes previously labeled. Store in vac. des. over 2 minute.

Initial assay.

	Count	Assay/tube
58-161	34	$.34 \times 10^9$
w-1177	148	1.5×10^9

Until 10/11/51, Silica used was Davian, 05-08-09-216.

New lot received 10/11/51 (Grade); 40-08-09-226

10/11/51.

B. ~~H295~~ H295. Harvest, 10:1, .05 ml / tube. Assay ml $\frac{100 \times 139 \times 10^{-3}}{2} = 3.5 \times 10^9$ per tube

	Count.	Surv.
1. Old Silica	33	
3. New Silica	< 10	
2. Silica Grade 923 Mesh 100-200.	< 10.	
4. Activated Alumina	1-2	

10/12/51 ~~Plate~~ Add tube to 10ml. Plate 10^{-5} ml

Very low survivals. Should be repeated.

Department of Genetics
University of Wisconsin
Madison 6, Wisconsin

January 1, 1952

Preservation of Bacterial Cultures on Silica Gel

This circular was written in response to a number of inquiries. Judging from these, present methods for preservation of bacterial cultures are not entirely satisfactory, and it would be a real contribution to laboratory technique to work out a better one. Unfortunately, I can only suggest a principle that seems very plausible, but that has not yet been empirically justified.

The working principles are (1) that suitably dried bacteria should survive just as well in a sealed tube under air as in vacuo, and (2) that if this is correct, chemically inert desiccants such as anhydrous silica gel could greatly facilitate the practice. The following arrangement has been tried: small vials or tubes are filled nearly full with silica gel (Davison Company, Baltimore; Grade 40, 6-16 mesh). About 1 to 1.5 gms. of silica fits well into the tubes used. The tubes are plugged, then baked in a sterilizing oven at 160-180 C., 2 hours, to dry and sterilize the tubes. These are stored in a desiccator. The bacteria to be preserved are suspended in 2% peptone. About .05 ml. is pipetted directly to the silica, and the end of the tube sealed off. The tubes were then stored in a refrigerator. The water disappears very quickly. To regenerate the cultures, the tubes were broken, and the silica poured into broth. Considerable gas is liberated. After about an hour to allow redispersion, viable counts were made on the broth. The 24-hour survival, in apparently dried condition, was quite high (about 10%), but this was more encouraging than long-term experiments. After four - five months, the survival has been low, of the order of 10^{-5} , and some tubes are inviable. In its present form, the method is not a success, and cannot be recommended for long-term preservation and storage. I think that it could be greatly improved, without complication, by experiments leading to a better suspending fluid, and possibly by drying the cells on a layer of glass bead over the silica. What is most needed is an improved theoretical understanding of the biology of suspended animation in successfully dried cultures.

Despite its shortcomings, the silica gel tubes provide an ideal method for mailing cultures. They are not affected by undue cold in the way agar slants are, and probably ought to be more resistant to high temperatures as well. The mechanical strength of small sealed tubes allows them to be sent with simple padding in an ordinary envelope, and the absence of any liquid minimizes possible hazards from breakage and leakage.

I hope that other workers with suitable facilities to study preservation problems, or with a potential interest in the biophysics of suspended animation may be able to make some use of this suggestion.

Joshua Lederberg

September 25, 1951.

A. K-12 overnight, aerated. Dilute 1:10 (est. 2×10^3 /ml)
 A2 .02 ml Assay original K-12: 5.7×10^9
 A5 .05 ml
 B2 1:1000 (est 2×10^6 /ml). .02 ml

Assay.

B2. ① Ca 2gm. Silica

② Ca .5 gm Silica

A2 ③ " Survival pS

④ " 2gm samples may have been exposed to heat close to cell dummy sealing.

1 6 6/114
 2 16 16/114
 3 875 8.75/114 1.115
 4 > #3 ca 10% 1 ±

A2 unit 1.14 x 10⁶/tube. 11,400 per plate = pS 0.

B2 unit. 114.

9/29/51.