

Fermentation tests.  
Mutants.

434

Spread out mustard-treated Y40 (see 426) on  
EMB-Galactose + a few maltose, sucrose, glycerol plates  
and examine for mutants. Ca 400 per plate.

Bacteria: 36. # of plates. (mucoid found)

∴ ca 15,000 colonies examined.

No fermentation mutants.

~~Holtoe~~ Mammotol 2 plates + 2 streak plates.  
ca 1000 examined.

2 very small M+ colonies noted. Streak out on fresh medium.  
all M+.

Glycerol - slow utilization. — compare passage on glycerol  
+ penicillin.

Sucrose - very slow, but definite utilization

Fermentation contents -  
enrichment cultures.

435

Feb. 25, 1947.

Dissolve 50 ml NB + 2% sugar + Brom cresol purple A25. Y53.

A26      A27      A3      A4      P5

<sup>symbols and acidity</sup>

glucose

+++

glycerol

++

Y81.

growth heavier than glucose <sup>in % black</sup>  
Y53

lactose

-

-

+ <sup>++</sup> ~~++~~ colonies

sucrose

-

-

- all white colonies!

\* growth inc. standing at room temp.

Streak out some apparently gly- colonies from 434 +  
compare with atypical gly+ — (b).

Y84.

a) —

~~Y80.~~

b) +

Compare a, b + gly+ enrichment culture above.

EMB

Y80

-

No evidence of papillae.

(from  
glucose alone)

Y53 gly ±

+

Y53 gly+ + or ++

different from Y53?

on BCP - medium is not changed in color, cells show slightly different shades (+, ± or pinkish, - more translucent or violet.)

on gly-BCP broth - gly-, gly± and gly+ all show slow acid + gas.

(see over)

1. Enrichment for gly +

---

A5 - streak from old tubes to new gly CMV

A8 - scoring OK, as before!

Resistance materials - cross test.

436

Streak out susp. of

$N$  Hager + Malibran.  $10^{-4}$ .

Streptomyces 5 u/ml

M.G. str. B.G.

Y77      ++ \*    -    - !

Y78      -      ++

\* dye is decolorized.

after 3 days, several hundred colonies  
appeared on the streak.      Y79.

streak Y77 over Y78/MG. to determine if decolorization same  
effect of dye. No evidence of stimulation of previously streaked  
( $Mg^+$ ) culture. Probably due to pH change.

# Inversion Detection

437

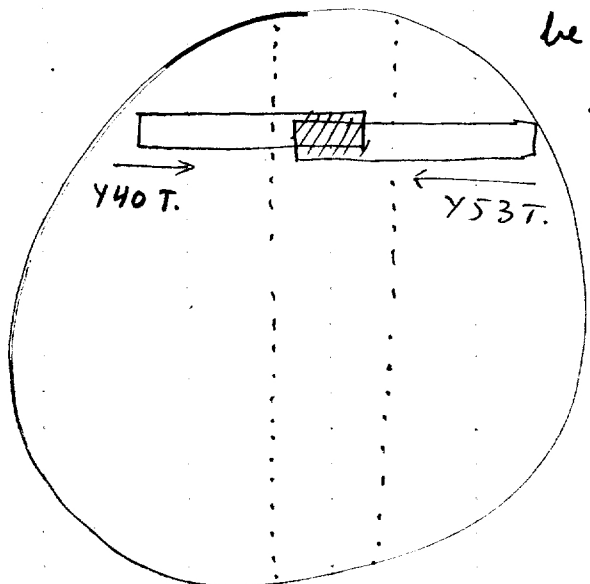
1 MAR 1964

See 437a for summary.

Streak out Y40, Y53 (T from 426) on NA plates.

P1. Pick single colony and streak on NA, overlapping a streak of the other type in the center of the plate, and mix well in center. This can then

be picked after growth and inoculated onto minimal.



P3.

Test combinations by suspending growth from center in water, <sup>1 ml</sup> and streaking pouring on minimal plates. (i.e. 84 tests!) ca  $10^9$  microdunum.

Tests log prototrophs

1	1+	13	1+	27	2	39	1
2	2	14	1+	28	1+	40	1
3	2	15	1+	29	2	41	1
4	1-	16	1+	30	1	42	1
5	1.5	17	1+	31	1+		
6	1.5	18	1+	32	1+		
7	2	19	1-	33	1		
8	1	20	1.5	34	1		
9	1	21	1+	35	1		
10	1+	22	1+	36	1		
11	1	23	1+	37	1		
12	1	24	1+	38	1		
		25	2				

39  
40  
41  
42.

1-6 also tested by streaking  
loopful on T(c) plates.  
colonies

0  
3  
1  
2  
4  
5  
6  
7  
10  
1  
1

84 tests - no inversions

Streak technique not as reliable as desired!

# Inversion Tests: Summary.

437A.

cryp. material. tests:

		Cumul. Yield Tests.
426.	MN <sub>2</sub> fit.	Y40 × Y53T; Y53 × Y40T, by st. tests., 0 + 0, 20 tests.
433.	MN <sub>2</sub> heat.	do. 17 0
437.	"	Y40T × Y53T. ni 0 only, X magas; 2 × 42 = 84 121 0
508	X-ray	Y40T × Y53T. ni 0 only. magas. 2 × 14 = 28. 149 0

Time of prototroph initiation.

438

Pour 440x453 plates in T(0). To exp. add also  $10^{-6}$  R-12 cultured + washed similarly in order to compare rates of colony development.

See 445.

Y65 } test for inversion.  
Y68 }

939.

Plate Y65 x Y40

Y68 x Y68.

in  $T(O)$  +  $T(O_1)$ .

no prototropes shall!

∴ not due to suppression of X in  $B_1$ - $B_4$  regions.

Segregation of *Gly*-

440

March 4, 1947 -

Y80 x Y81. cf. 435.

$\beta_1^- \rightarrow \beta_1^+$  as usual. Pick & plate on EMBlac + EMBlgly agar.  
o  $\beta_1^+$

Lac-R Lac-S Lac+R Lac+S  
 $\beta_1^-$  Lac-R -S +R +S.  
 $\beta_1^+$

Dly segn.

Y80 x Y80.

Dly

Lac

V

+  
+  
+  
+  
+  
+  
+  
+  
+  
+

-  
-  
-  
-  
-  
-  
+  
-  
-

R  
S  
R  
S  
R  
S  
S  
S  
R  
R  
S  
R

Dly

Lac

V.

-  
-  
-  
-  
-  
-

R  
R  
S  
R  
R  
R  
S  
K

\*

[scoring maybe  
inadequate]

when ref again

4 hours later, series

B had gotten con-  
siderably darker!

A.

in min.

Lac

V

T(0)

T(B<sub>1</sub>)

C.

B<sub>1</sub>

-  
-  
-  
-  
-  
-  
-  
-  
+

-  
-  
-  
-  
-  
+  
-  
-

R  
S  
R  
S  
S  
R  
R  
S  
R  
R  
S  
R

D.  
[

-  
?  
+  
-  
-  
-  
-  
-  
-  
B<sub>1</sub>

-  
-  
-  
-  
-  
-  
-  
+  
-  
-  
R  
S  
S  
R  
R  
S  
S  
R  
S

B<sub>1</sub>

?  
-  
-  
-  
?

Note: B<sub>1</sub><sup>+</sup>: 25/25 Dly +  
B<sub>1</sub><sup>+</sup> + B<sub>1</sub><sup>-</sup>: 44/46 Lac +. Dly + close footage? These results are difficult  
to reconcile with the map!

\*

Lac, V  
segregation.

440

March 6, 1947

Lac-  $V^R$  Lac-  $V^S$  Lac+  $V^R$  Lac+  $V^S$

$B_1^+$     ~~15~~ ~~9~~ 0 1 25    Faulty segregation of Lac +?

all bly+

$B_1^-$

bly- 24 19 5

bly+ 1 (1?)

This scoring of bly, done at 2 days,  
is not borne out by readings at 3 days,  
when in both groups, all were bly+,  
with some variation in intensity.

Note: 24 Lac- : 1 Lac+ in  $B_1^+$  (95%!)  
46 Lac- : 5 Lac+ in  $B_1^-$ .

$$1) B_1^- / B_1^+ \quad \begin{array}{r} 25 \\ 41 \end{array} \quad \begin{array}{r} 24 \\ 40 \end{array} \quad \begin{array}{r} 1 \\ 4 \end{array} \quad \begin{array}{r} 25 \\ 45 \end{array} \quad \begin{array}{r} \\ \hline 64 & 6 & 70 \end{array} \quad \begin{array}{r} X^2 = .04 \\ .02 \\ .04 \\ .03 \\ \hline .13 \end{array}$$

compare to standard:

$$\begin{array}{r} 18 & 24 & 7 & 1 & 25 \\ 51 & 44 & 19 & 6 & 70 \end{array} \quad \chi^2 = \frac{13^2}{51} + \frac{13^2}{19} = \frac{3.3}{8.9}$$

a) old data: 72% Lac- ||     $\chi^2 = \frac{36}{18} + \frac{36}{7} = 7$      $p = < .008$ .     $\frac{3.3}{12.2}$   
 b) new data 64% Lac- ||    efficiency of Lac+

Peculiar segregation may explain peculiarities of bly segregation.

Consider

if back water is desired

Segregation of drug-resistances

441

A ~~Y77 x Y78~~  
Y77 x Y78. - in T/0) OK!

Streptomycin 54/ml  
Malcilate 20. 100v/ml

B Y53 x Y78 OK! ~~#6 Lac - 1 Lac +.~~  
as above, but with

C. Y40 x Y77 OK! no F

A. Y77 x Y78.	Lac	V	str	M.G.	Lac	V	str	M.G.
	+	R		R	-	R		R
<small># per cent</small> <small>blue color</small>	+	RS		R	-	S		R
	+	S		R?	-	S		R
	-	R		R?	-	R		R
	+	R		R?	+	R		R
	+	R		R	+	R		R
	-	R		-	-	R		R

? probably S.

scoring uncertain

due to selection of  
resistant residue

M.G. resistance  
 $10^{-4}$ . lac lac+MR lac+Ms lac-HR lac-Hs.

C.	2	1	8	2
<u>total</u>	0	2	16	3

scoring not certain

Lac+S x	2	3	24	5
Lac-R				

ind. cont.

absence of resistant. If valid, indicates linkage to B4.

should score on minimal plates to avoid selection for  
resistant contaminants.

P3. - mix YB cultures of Y77 ( $Mg^1$ ) and Y78 ( $St_5^{-1}$ ).  $30^\circ$   
also use mixture of mut. plating technique of 441.

1.	Mix culture in $Mg + St_5$	$A^6$ $0, 0$	$A^8$ 1 colony;
2	Y77	"	2 colonies; <sup>as</sup> 3.; 4 colo.
3	Y78	"	1 colony ca 100 minute clearings, <sup>as</sup> colo.
4	Y77 + Y78	"	ca 10 v. small <sup>A - as 3.</sup> <sup>B &gt; 10 large colonies</sup> innumerable minute clearings
			A6 - incubate at $35^\circ$ .

Indeterminate whether the mult. resistant colonies represent recombinations.

Compare 4 (cross)  $\bar{c} \bar{c}$  mutations of  $Sm^S Mg^R$  to  $Sm^R Mg^R$

There may be some synergism in view of the lag by before colonies are detectable!

[See Y84  $\times$  Y79. and plate in brilliant green. 7  
+ streptomycin.]

Resistance steps; resistance to streptothrin.

443

March 4, 1947.

Recd. 127,000 u. streptothrin from Woodruff, Thruhs.  
non-strike ampule.

Suspended in 2.7 ml 95% alcohol for 3 hours Add  
10.3 ml sterile H<sub>2</sub>O → ca 10,000  $\frac{u}{ml}$  in 20%  
u/100ml = 1ml alcohol. Dilute further as required.

100u	1
Y78	T
Y53	T

500	5
Y78	T
Y53	1-200

strike Y78 on 5u agar for —  
Y82 - streak out on 5u agar

1000	10
Y78	
Y53	

Y83 - streak on 10u agar  
Y84 - do.

10,000	100
Y78	0
Y53	0

Y78      Streptothrin  
on 5u agar gives colonies but not so large  
as Y82.

Y83, Y84 OK on 10u agar.

See 452.

Y40 x Y53. Mix cells in agar pour on small plate.

$$\begin{array}{r} 0: \quad 11 \\ \quad 15 \\ \quad 9 \\ \quad 10 \\ \quad 15 \\ \hline 60 \end{array}$$

m = 12.0

$$\begin{array}{r} B: \quad 12 \\ \quad 11 \\ \quad 14 \\ \quad 12 \\ \quad 9 \\ \quad 13 \\ \quad 6 \\ \hline 12 \end{array}$$

m = 11.1

$$\begin{array}{r} B_1: \quad 124 \\ \quad 136 \\ \hline 130 \end{array}$$

$m = \frac{100}{130}$

more turbid.

A).

B)

20 B tested: 18 B<sup>+</sup>2 B<sup>-</sup> (Lac-R; Lac+R)~~+/-~~

5/20	364
2/20	445
5/50	452
12/90	13%

C. Seg. of Lac, V<sup>R</sup> in 0, B<sub>1</sub>:

0:	-R	-S	+R	+S
24	<del>11</del>	<del>9</del>	<del>1</del>	0
B <sub>1</sub>	22	<del>15</del>	14	0

	-R	-S	+R	+S
0	17	10	9	0
B <sub>1</sub>	16	12	13	0
51.	11	11	11	11

None 0 plate, streak out 11-12 mm surface. These colonies appeared at same time as prototrophs (24 hours) and were of comparable size.

## Lac - V segregation.

446

March 6, 1947.

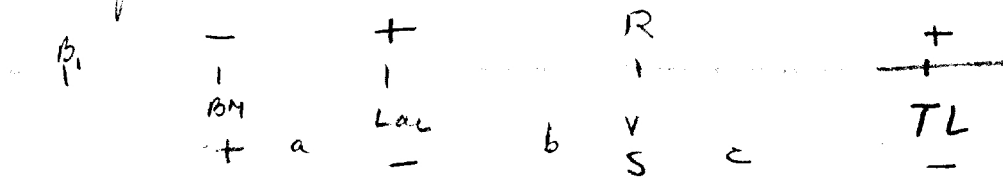
Use colonies from Expt. 437  
(grown together - chance for selection) [Y40 x Y53 (T) by plate number].

Plate #		Lac - R	Lac + R	Lac - S	Lac + S.	$\Sigma$	* V scoring not dependable -
9	10	2	5	1	0	8	
20		4	4	2	0	10	phage apparently
*	27	40-30	9	6	0	0 { 15	very low titer.
*	22	40-30	8	10	0	0 } 18	no lysis by itself
*	3		8	3	0	0	
			3	1	0	0	4
			7	3	3	0	13
			1	2	1	0	4
			6	2	2	0	10
			4	2	0	0	6 Lac - = 135/214
			6	1	0	1	8 = 63%
			1	1	2	0	4 compare = 70% per.
			2	1	1	0	4 V <sup>R</sup> = 190/214
			6	3	2	1	1L = 85%
			3	2	0	0	5 comp. = .74 per.
			13	7	1	0	21
			5	14	2	0	21
"			1	3	0	0	4
"			11	1	3	0	15
"			1	1	1	0	3
"			5	1	0	0	6
"			7	4	1	0	12
			113	77	22.	2	/214.

The agreement of the Lac+R; Lac-S classes with exp from previous  
expts is very poor. Reexamine crosses of aberrant cultures.

There is a shift from lac-S to lac+R

On map basis



$$\begin{array}{c} < \text{lac-} \quad \text{lac-} < \quad > \text{a} \\ > \text{v}^R \quad \text{lac-} < \quad < \text{c} \end{array}$$

(Lac. more than lac+)

$$\begin{array}{l} a = +R \\ b = -R \\ c = -S \end{array} \quad \left[ \begin{array}{l} \text{-1 would not be augmented by the} \\ \text{elimination of } c? \end{array} \right]$$

or, another interpretation, is that ~~the~~  
the previous strokes were revised for T or for L, i.e. a longer distance  
in the interval v-(T or L)

∴ compare types  $c$  at short or long interval  
(i.e. low or high  $-S$ ) biochemically.

card. 446.

#	Loc-R	Loc+R	Loc-S	Loc+S.	$\Sigma$
15.	3	9	1	1	14
33	2	1	2	0	5
34	5	0	10	0	5
30	6	2	2	0	10
31	7	7	4	0	18
24	3	3	1	0	7
32	7	3	1	0	11
35	7	2	1	0	10
1	6	4	4	0	14
36	4	2	2	0	8
16	12	10	3	0	25
26	14	9	3	0.	26

$$\frac{83-76}{83} \cdot \frac{52^{53}}{106} \cdot \frac{22^{21}}{53} \cdot \frac{1}{76} / \frac{151}{27}$$

$$\frac{106-113}{106} \cdot \frac{77^{76}}{129} \cdot \frac{22^{27}}{44} \cdot \frac{2}{3} / 214$$

$$\frac{189}{189} \cdot \frac{129}{129} \cdot \frac{44}{44} \cdot \frac{3}{3} / 365$$

$$\chi^2_2 = \frac{49}{83} + \frac{49}{106} + \frac{1}{53} + \frac{1}{76} + \frac{9}{20} + \frac{9}{27} = 1.86 \quad p = .4.$$

cf. 445.  
compare  $\Sigma$  remainder  
forget.

These samples agree.

Homogeneity??

.59 | 1.86

.46

.02

.01

.45

.33

analysis of  
446 vs 359 summarized.

-R +R -S +S.  $\Sigma$

189	129	44	3	365
100	55	50	4	209
289	184	99	7	574

$$\chi^2 = \frac{25}{105} + \frac{25}{184} + \frac{144}{67} + \frac{144}{117} + \frac{196}{36} + \frac{196}{63} + \frac{1}{4} + \frac{1}{3} \dots$$

$$= .2$$

$$.1$$

$$2.2$$

$$1.2$$

$$5.4 \\ 3.1$$

$$6.2 \\ 3.3$$

$$12.7 = \chi^2_3$$

$$8.5 \\ 4.2 = \chi^2_2 \quad p = .04$$

it is the difference in the frequency of  
-S which differentiates the distributions.

category	194 - 189	129	318
	95 - 100	55	60
	289	184	473

should be 194:124  
95:60

$$\chi^2_1 = \frac{25}{189} + \frac{25}{124} + \frac{25}{60} = \text{etc.}$$

Hendry of segregation types.

447.

March 7, 1947

Recover 446 - 22, 25 + 27. in order to ascertain suitability of  
disproportions in ratios. Compare segregation of lac + v = 440, 453 stand.  
In previous test:

A	22:-	8:10:0:0
B	27:-25	9:6:0:0
C	25:-27	5:14:2:0

Lac, V.

D. 440 x 453.

	-R	+R	-S	+S
--	----	----	----	----

A	47	19	9	1
B	44	20	7	1
B	26	12	16	0
	13	16	0	
C	7	5	2	1
	80	36	32	2

D. -	30 ✓	21 ✓	19	0 ✓
O <sub>2</sub>	10	7	5	0
N <sub>2</sub>	6	5	8	2
M.B.	4	4	6	0
S.H.	2	1	2	0

D. total: 52. 38 39 0

This is homogeneous with  
the previous tabulation:  
100:55:50:4.

Total

132 74 71 2 279.

$$\chi^2 = \frac{100}{37} + \frac{100}{62} = 4.3$$

Compare A = D..	47	19	9 + 1	0	76	$\frac{4}{21} + \frac{4}{36} = .28$
	52	38	39	0	129	$\frac{64}{18} + \frac{64}{31} = \underline{5.6}$
	99	57	49		205	$\chi^2 = 10.1$

Second.

$$p = .007$$

Exposure  $R \bar{=} S.$

58	66	10	76
98	90	39	31
156	49	129	205

$$\begin{aligned}\chi^2 &= 64 \left( \frac{1}{58} + \frac{1}{98} + \frac{1}{10} + \frac{1}{31} \right) \\ &= 64 ( .017 \quad .010 \quad .055 \quad .032 ) \\ &= 64 (.114) \\ &= 7.3 \qquad p = .007.\end{aligned}$$

---

Therefore the total discrepancy is due to a difference in the proportion of  $V^R$  cultures. In practice, this means a deficiency of  $-S$  cultures in the new group.

-R +R -S +S.

A. 22      37      20      7      1

B. 25      25      13      16      0

C. 27      7      5      2      1

76      38      25      2

D.      —      30      22      19      0  
 O<sub>2</sub>      10      7      5      0  
 N<sub>2</sub>      2      1      2      0  
 SH      6      5      6      1  
 MB      4      4      5      1

$\Sigma$ .      52      39      37      2      447  
 69      38      31      2

This is the "abnormal" distribution.  
 1 plate showed 13:14:0:0.  
 (perhaps scoring of V<sup>R</sup> was  
 not proper.) Recheck! ✓

Plating plate

compart. —

A	6	2	0	0
	6	14	6	0
	8	1	3	0
B	17	3	4	1
	4	1	6	0
	12	10	8	1
C	9	2	2	0
	7	5	2	1

89	41	130
100	40	140
189	81	270

Caution! Be careful of scoring V's.

Interpretation of 447.

448

Everyone  $T^- \rightarrow T^+$  in 447 lines + standard.

March 7, 1947

Using cells from Exp. 447, plate the Y53 components at a  $10^{-2}$  dilution into LB<sub>1</sub> agar.

Y53	tube 6	10 <sup>-2</sup> dil.
(Y53-422)	tube 7	30 sec.
25	10	"
22	1-2	

There is no stable differentiation, apparently due to scoring difficulties.

Plating medium for recombination

449

March 7, 1947

factors: buffer: phosphate, citrate, acetate, phthalate. More alkaline.

RH: thioglycollate; ascorbic acid; pyruvate-lactate;  $O_2$ ; Methylene Blue. Hydrogen??

B<sub>1</sub>? - Try on BM + TP.

T(0) - V40 x Y53 mix in agar & pour on pre-poured plates.

1. Controls

2. "O<sub>2</sub> atmosphere" least turbid plates

3 N<sub>2</sub> atmosphere turbid plates; colonies very small, countable once rare.

4 Na thioglycollate 2 mg =  
2.2 mg =

5. Ascorbic acid 1% no colonies (pH 4-5) No turbidity

6 Methylene Blue 100<sup>-4</sup> Turb. like  
10<sup>-5</sup> " "

The number of apparent colonies is correlated inversely with the gross turbidity, and was best in O<sub>2</sub>, most in N<sub>2</sub>. S19 had no apparent effect, however. There is, however, no marked increase of colonies in the O<sub>2</sub> atmosphere plates, but those which do appear are larger.

See 441. Test Y78. on EMB-Lactose. cf. 453.

- also 441A - Lactose - on Lac → glycerol white + black colonies +  
 Y78 (<sup>Lac</sup>  
58-161 Sm<sup>2</sup>) ++ pale. gly. count as Lac some blue blocks  
 58-161 +++ - greenish ! - (±??)  
 Y80 (58-161-V<sub>1</sub>-N<sub>1</sub>-) -

Note, 480 on v. long incubation turns a faint violet color, à la sucrose  
 from previous expts & now

Y53	++		Y80	-
Y81	++		58-161	±
Y40	++			
Y78	++			
		relative amts.		

58-161 is slow, but will score vs. 480. see 453.

Y80 x Y81.

Gly. segregation:

March 9, 1947.

A.

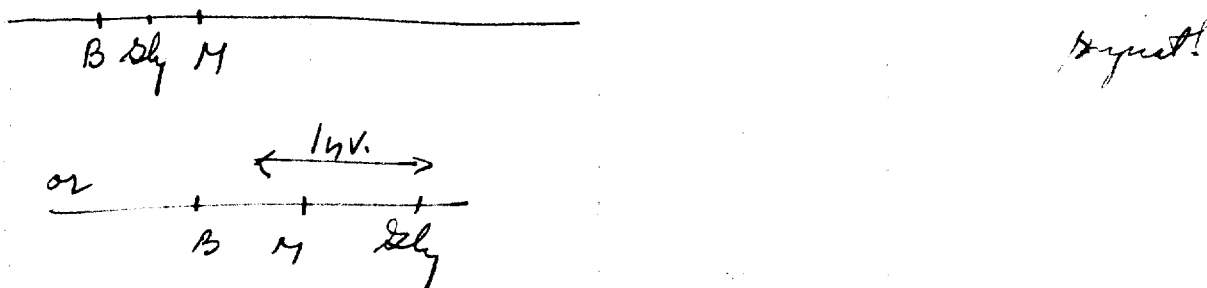
Repeat 440

See 451A.

Y80 x Y81.

T(0).	16/16 Gly+	12/14 Lac-	86%
T(B <sub>1</sub> )	13/13 Gly+	58/72 Lac-	78%

~~2~~ Note, this denotes absolute linkage of Gly to B4. This is predictable either on the mapping: written Y80 x Y53  
nor Y81 x Y40  
unpublished!



Note (also) the segregation for Lac may be disturbed. Compacting occurs.  
data for Y40 x Y53: (71% Lac-)  $p = .06$

$$\bar{e} \text{ Y40} \quad p < .025$$

$$(\text{this} + \text{Y40}) \bar{e} 71\% \quad p = <.001$$

$$\text{Y40} \bar{e} 71\% \quad p = .008$$

$$B_+^+/B_-^- \quad p = .3$$

Sexual segregation.

Mauls 9/1947.

B.  $Y80 \times Y53$ . Test  $B_1^+$  + ( $B_1^- + B_2^+$ ) on Sly; lac. These plates were allowed to stand 3 days before testing.

a.  $B_1^+$  27 Lac- : 9 Lac+ 35/35 Sly+

b.  $B_1^-$  50 Lac+ : 16 Lac+ scoungumentari: 1++ ; 30+/31  
77 Lac- : 25 Lac+ ca 75% lac-

C. ~~B.~~  $Y81 \times Y40$ . Test on lac | 6/16 16/16 +  
(out & out)

$B_1^+$	104 Lac- : 43 Lac+
$B_1^-$	28 : 13
	132 : 56 188

70% lac-

This experiment is not homogeneous with the earlier  $Y40 \times Y53$  ones.

D.  $Y53 \times Y40$ .  
 $B_1^+$  | -R -S +R +S → stand  $\chi^2 = 3.2$   
 33 32 28 25 14 18 0 →  $p = .2$  late wt  
 $B_1^-$  | 18 19 12 15 13 10 } 44 →  $\chi^2 = 3.5$  →  $p = .2$

51. 40 27 1 119 g. standard.  $p = .06$

91 : 28 76% lac-

Segregations of B-

452

March 9, 1947.

See 445

Y40 x Y53 in B. lotus medium. Rick colonies to H<sub>2</sub>O, +  
moi. longerful in T(0) + T(B). 50 isolates.

50 isolates. At 12 hours, 5 diff. B-  $\frac{1}{2}$  did not grow in either  
way V size. 5 at first.

	Lac	V
1	+	R
2	+	R
3	-	R
4	-	R
5	-	R.

P13 - 5 grew on B, not O.  
Ancest, 1-4 contained primarily  
-S, E + R as well. 5 was uniformly  
-R. see 456.

To summarize B-

~~-R - S + R + S.~~

-R + R - S + S.

III I III 0 1  $\therefore 0-S/12$ .

	6	5	0	1	12
A ( $\bar{x}$ )	6	3	3	.36	

$$\chi^2 = \frac{4}{3} + \frac{9}{3} = 4.3 \\ p = .14$$

$$\chi^2 = 5 + \frac{16}{9} + \\ = 7 + .03$$

$\therefore$  maps

B. g B 13 M

March 11, 1947.

P11. streaks on single glycol plate (EMB-2%) the following:

	K-12	A12	P12	A13	4P13	A14
1	-	-	+	++	+++	+++
2	58-161	-	-	-	-	+ streaks.
3	Y40	-	-	±	±	+ streaks.
4	Y10	-	-	±	+	
5	Y53	-	-	+	++	
6	Y46	-	-	+	++	
7	Y64	-	-	±	++	
8	Y78	-	±	++	++	
9	Y77	-	-	±	++	
10	Y80	-	±	±	±	translucent, creamy shade, not opaque purple.
11	Y81	-	-	+	++	
12	Y82	-	-	±	++	
13	Y73	-	-	+	+	
14	Y74	-	-	+	+	
15	Y82	-	-	+	++	
16	Y83	+	++	++	++	
17	Y84	-	-	+	++	
18	Y79	-	-	+	++	

4/11/47

on various plates, streak: *Escherichia coli* 20  
*Salmonella* 20  
*Phytophthora* 21  
*Staph aureus*

1  
2  
3  
4  
5

nutr. -

Vor 4/11.

Malachite	H.G.	1	++	✓	-	++	✓	+	✓	+	✓	5.
Brown		10	++	✓	++	✓	++	✓	-	-	-	✓
		50	-	-	-	-	-	±	-	-	-	-
		100	-	-	(- colo.)	-	++	-	-	-	-	trig colo.
Bullard	B.G.											
Brown		100.	-	-	✓	-	✓	-	-	✓	-	✓

*Streptomyces* Sth 1 ++ ✓ + - ✓ + - ✓ + ++ ✓ ++ ✓ ✓

5 ++ ✓ - - ✓ ± ± papillae ± + + - ✓ colonies ✓ O.H.

*Streptomyces* Sm 10 - ✓ (prod colo) - - - ✓ ± + ✓ -

1 - - ✓ + + - + + - ± + ✓ - ✓ colonies ✓ grow large.

5 - - - - - - - ± - - ✓

*Penicillium* sm! 10 ++ ✓ - ✓ - ✓ - ✓ - ✓

sm! 100 ++ ✓ + + - ± + ++ ✓ ✓

Contd N.A. ++ ++ ✓ + + - ± + + - ++ ✓

Mar 10/11. Readings 1st. 11/10 A 12.

6 A 12

9 A 13

Contd. - [kinds of selective case.] : H.G. B.G. Sth Sm Pen

<i>Escherichia</i>	10-50	- < 100	{	10	10	< 1.	-	
<i>Sal</i> 20	50	50 < 100	{	5	5	1-5	> 1	/ 0
<i>S21</i>	> 100	{ < 100	{	> 5	> 5	1-5	> 1	-
<i>Phyto</i>	1-10	{ < 100	{	> 10	> 5	1-5	> 1	-
<i>Staph</i>	< 1	{ < 100	{	5	5	1	1	10

order of activity:

M.G.	S.m	Sth	Pens:	
Steph	Sevn.	Steph		
P.t.	Steph	S 20		
S.m.	S 20	S 21		
S.20	S 21	Senaria		
S.21	Phyto	Phyto	Senaria	

↓ mix:  
resistame

use smaller of following papillae on higher drug rate for higher steps:

S 20/MG 50 on B.G. 100  
M.G. 100.

S.m / Sth 10

S 20 / Sth ~~25~~

S 21 / Sth ~~25~~ on higher Sth.

Steph / Sth 5

S 20  
S 21 | S.m | on higher S.m.  
Steph |

# Resistance mutants

454a

<u>Steph.</u>	M.G.	1 5 10	turbid. ca $10 \text{ Mg}^{\text{R}}/\text{cc}$ ; gd. inhibitor of residue perfusely clear!
<u>S. m.</u>		1 5	ca $10^4 \text{ S.m.}^{\text{R}}$ . ca $10 \text{ S.m.s.}^{\text{R}}$
<u>Proflavine 100.</u>			Turbid.
<u>Seratia</u>	M.G.	10 20 50	clear at $\frac{1}{4}$ hr. clear at 24 hrs. a. ca $20 \text{ Mg}^{\text{R}}$ (ctd?)
<u>S.m.</u>		1 5	clear, no S.m. seen.; some v. small S.m. ca. $10 \text{ mg.}^{\text{R}}$ ? colonies white!
<u>S.th.</u>		100 20	ca 5 S.th. $^{\text{R}}$ / ml col. v. small ca $100 \text{ S.th.}^{\text{R}}/0.1 \text{ ml}$ wide range of color.
<u>Profl.</u>	100		turbid
<u>S.20</u>	B.g.	100	clear, no B.g. $^{\text{R}}$ !
	S.m.	1	turbid
	S.m.	5	turbid; but some possibly S.m. $^{\text{R}}$ , v. small
	Profl.	100	turbid.
<u>S.21</u>	B.g.	100	clear, no B.g. $^{\text{R}}$ .
	S.m.	1	turbid.
	S.m.	5	as S.20. inhibition incomplete.
	Profl.	100	turbid.
<u>Phyto.</u>	M.G.	10	inhibition incomplete
S.th.	"	50 100	inhibition incomplete; some selective clear! ca 100 $\text{mg.}^{\text{R}}$ resist.
S.m.		10 50	incomplete inhibition Clear! ca 100 $\text{mg.}^{\text{R}}$ resist?
Profl.		100	clear! - ca 300 $\text{mg.}^{\text{R}}$ resistant

available:

~~St.~~: Sth 100  
Phyt. Sm 50  
Prof 100

St.: Sm 5  
Sth 10  
Thg 5

S20 B.G. S21 Bg  
Sth 10 Sth 10  
Sm 5 Sm 1

Senates Sth 20 }  
50 } excavation.  
100 }

Sm 5  
Mg 50

4/11/47.

As above in N.A. and T(0).

photographs show up as papillae in T(0) streaks, justifying this technique. Throw out N.A. plates.

T(0) colonies in center zone

1	++
2	++
3	+
4	++
5	++
6	++
7	++

Streak out further 48 hrs. colonies of 440T + 453T.

n.g. Too irregular. N.A mix + plate is more reliable.

14/11/57

See 452 for sources.

Colonies from 440 x 453 on Broth were picked to water + T(O), T(B) inoculated. Those reported are 5/50 which grew on T(O) only after 2-3 days. The T(O) and T(B) tubes were both streaked on Mac-V agar: ( $\downarrow$  turbidity)

	T(O)	T(B)
1	-S; +R	-S; +R
2	-S; +R	-S; +R
3	-S; +R	-S; +R
4	-S; +R	-S; +R
5	+R	+R.

Since -S and +R are the parental configurations, the delayed growth [and the original colony formation] might be due to syntrophyism?

∴ streak out T(O) tubes on EMBA to purify.

Test - (a) and + (b) colony of each on B, O medium.

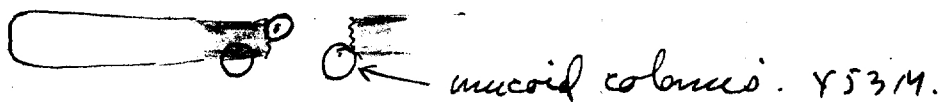
1a	=	=
1b	++	++
2a	=	=
2b	-	-
3a	-	-
3b	-	-
4a	-	-
4b	-	-
5a	=	=
5b	-	-

Expl. 1. Syntrophic colony  
 2. Colony not picked; only intestinal growth in agar. Requires repeating.

19 Mar 1951.

Streak Y53 across T1 on EMB agar.  
Suspend slimy growth from " " intersection of bacteria + phage  
with S. + streak out.

Note, at intersections of bacteria + phage, a zone of coloration of  
the bacteria as if there were there some enzymatic activity!



N21. Pick from red region + from mucoid colonies to water  
and streak out on EMB lac. (Y53/1).

Y53/1 same growth (probably resolants; ~~bac-~~ bac- +, lac+).  
Standard type

1. Y53M - all mucoid. pick one colony + spot on T1 VR  
also streaks out →

2 Y53M<sub>2</sub> = all mucoid. P23. Streaks out →

3 Y53M<sub>3</sub>. all mucoid. Pick to slant and shelf for subsequent  
analysis. Y86

Plate Y40 x Y53 as usual. At. 48 hours pick 5 largest (>) and 5 smallest (<) colonies from each of 7 plates + compare the distributions.

-R -S +R +S.

4	1	0	0
---	---	---	---

4	0	1	0
---	---	---	---

3	0	2	0
---	---	---	---

2	2	1	0
---	---	---	---

2	0	1	0
---	---	---	---

2	0	3	0
---	---	---	---

1	0	4	0
---	---	---	---

In large, +R > -S.

In small +R > -R.

---

18	4	13	0	{	$\chi^2_3 = 5.14$	$p = .16$ .
1		17	1			

But. compare all 3 groups,  
Random selection from these plates gave:  $\chi^2_4 = 12.61$   
 $p = .013$

27	13	13	1.
----	----	----	----

selection may play a role.

221	136	142	6
-----	-----	-----	---

is cumulative data.

Note that both in in large+small selection types, there is a marked deficiency in -S as compared to random selection + cumulative data!