

1

Selection for sucrose + mutants.

Oct. 26, 1947.

Pupate 10ml. 2% sucrose 1% peptone, ~~at 1% yeast~~ with  
Bran-Creat Purple Indicator. Broz. as indicated, in series.

A. Y53. 10/26/47. ① 5P.

	2 P 27	5 P 28
1.	st. discoloration	-
2.	+	±
3.		
4.		
5.		
6.		

B. 58-161. ① 10/26/47. 5P.

1.	++	++
2.	-	-
3.		
4.		
5.		
6.		

(-)

IA3 was moved into IA4 P30  
and left overnight at room temperature. Later turned alkaline. Not sucer +  
p4. IA4 was +. Struck out on sucrose EMIB. {<sup>25°</sup><sub>37°</sub>} both Sucer -

Y105

Y106.

I abandoned A6.

Selecta for  $\alpha$ -methylglucoside mutants.

2

Oct. 26, 1947.

1%  $\alpha$ -methylglucoside (ER) broth, autoclaved together. BCP rods.

A. 453. ① 5P

growth  
only fair  
at 24 hr

2P27

- ↓ =

5P28.

↓

B. 58-161 ①. 5P.

- ↓ =

↓

abandoned A 6.

5.

"Transmission" of alkaligenesis.

Oct. 26, 1942.

Glucose + 2% peptone. Vary glucose in methyl-red indicator to establish the critical level. K-12. 3P26. In duplicate.

Glucose: 2% 1% .5% 2% 1%  
Methyl red destroyed.

3P27. Growth + fermentation +. No color changes. Test pH in 1 series.

pH:	acid to MR	++	++	++	±	-
" " BCP.	++	++	++	++	±	-

Use glucose .3%; BromCresol Purple. Prepare series of tubes  $\pm$  2% peptone. Incub. each with K-12. P28.

(B) (-glucose).

11A29. pH B. = ca. 6.8. pH A. < 5.2.

Add eq 8ml B to A1. pH ca. 6.0.

Retain sample in refrigerator for comparison  
to detect alkali production.

- A1. A30. Alkali produced. Transmit to A2.  
i.e. mix cultures.
- (A2) A3. Alkali produced Add to (A3)
- (A3) A6. Alkali produced.... Start A4
- (4) A10. Alkali produced.  
end. Expt.

10/25/47.

Grew Y53 from 2x25 ml 1/25 ft. YP broth and from 3x10 ml YP agar slants 20 hrs. old. Suspended in 10 ml 1/10 phosphate buffer at pH 6.0. Add 20 mg HN2 nitrogen mustard [ $\text{Cl}(\text{Et})_2\text{NCH}_2\text{Cl}$ ] in 10 ml buffer. Let stand 30 minutes. Sediment cells and replace with peptone 10 ml. 1% inoc 1 ml samples into 25 ml YB broth and incubate to prepare inoculum for mutant detection. "spread 1 ml as check on survival  $\rightarrow$  ca.  $10^5$ . Estimate.

$$PS = \log_{10} 10^5 / 10^5 = 5.$$

Many mucoids noted!

	Plates	Colonies/plate	Total colonies	Suspected mutants
10/28/47.	40	250	10,000	-1 not Mal- -2 (mucoid). -

	T1	T4	T6	T7	Lac	Natr.	-3
-1 (W1)	S+	R	S	S	-	T-L-B, ±	-: mutant!
-2	R	R	S	S <sup>2</sup>	-		
-3 (W3)	S'	R	S	S	-	? Hal-	"popillae n fist streaks. Pectinate".
Y105	R	R	R?	R	+		
Y106	R	R	R?	R <sup>1</sup> plaque	+		
X-12	S	R	R?	S	+		

10/29/47. Spread similar populations on galactose EMB plates. 11 AM.  
(colony dimorphism noted •  S/R?)

Of ca.  $40 \times 250 = 10,000$  colonies, no galactose + were noted. However, one colony which was unusually light purple was noted + specified for further study = 4-4. : Not Gal-.

Oct. 28, 1947. Test the cultures indicated on the following EMB:

	Y53	Y87	58-161	Y53	Y87	58-161	
Lactose	++	18 hrs. ++	++	-	-	-	→ ✓
Rhamnose	-	-	-	-	-	alb	+ ✓ + + peptid! some
Inulin	-	-	-	-	-	alb	*
Melibiose	+	+	+	+	+	+	✓ *
$\alpha$ -methyl glucoside	-	-	-	-	-	-	-
Cellobiose	-	-	-	alb	alb	alb	- pap? - ± ± **
Dulcitol	-	-	-	-	+	+	✓
Sorbitol	++	±	+	++	+	+	✓
Sucrose	-	-	-	alb	alb	alb	-
Ethyl Butyrate	inhibited		-	-	-	-	produce alb
Lactose	-	-	+	-	-	++	

on  $\text{CaCO}_3$  (.1%) peptone agar:

Lactose Minor clearing around lact colonies. Not distinct enough!  
Ethyl Butyrate (add "sterile" to hot medium.)

Cellobiose  $\alpha$ -methyl glucoside Sucrose Melibiose

w-1

w-3.

\* Streaks not uniformly dark: lighter in center. However, this is rather a thin plate.

\*\* streaks out the papilla.

P 31. Y53 is Dulcitol negative; Y87 + 58-161 Dulcitol +.

[dulcitol is related to galactose!]

~~w-1.~~ Cellobiose slow +  
10/31/47.  $\alpha$ -methyl glucoside -  
Sucrose -

Cellobiose - . Papillae (?) as 58-161 and Y87?

Streaked again

Carbon source utilization

$T(m)$  + indicated carbohydrate + BM for 58-161  
 $G = \text{glucose } 1\%$  (1%) + TLR for Y53...  
 $g = \text{glucose } .05\%$

16 hrs. 37°. O G g Maltose, + d.m.g. | d.m.g. + g + G. Celloolose + g + C |  
W-1 - ++ + + + ± ++ ++ + + + + + + + + + + + + + +

Y53. - ++ ++ + ± ++ + + + + + + + + + + + + + + + + + +  
W-1 ~~Sucrose + g.~~  
± ++ + + + +

Y53. ± ++ + +

-BM	+BM	G	Lac	Suc
O	O			

58-161 - - ++ + + ±

48 hrs. 487. - - + + ± -  
W-1 O g M M + d.m.g. + g + G C C + G Suc + G + g  
++ +

Y53 - + + ++ ++ + + ± + + + + + + + + + + + + + + + +

BM	G	Lac	Suc
-	++	++	±

487 - ++ + ±

Use .05% in testing sugar utilization in future growth experiments.

Bacillus in 58-161. 11457 HRL.

P 29. inoc 58-161 into YP broth. 2 P 30 Harvest and suspended in 1 ml  
M/10 phosphate, pH 6.0. Add 10 mg. to each of two tubes -  
Hpd.

a. 2:28 PM - 3:02 + 8 min centrif. 34 min.

~~2:48 PM~~ b: 3:02 - 3:17 + 15 min.

at 3:02 PM dilute to peptone  
Centrifuge 10 min. Resuspend in peptone  
Inoc 1 ml each into 25 ml YP. Spread 1 ml samples to  
assay killing.

Killing a)  $\gg 10^4$

b)  $> 10^4$  survived. PS < 5. Spread out +

Retire survivors also!

Spread on Bac, Dal. and Mal.

Galactose. 83 plates. Sharp division into large + small colonies  
noted. ca. 300 large + 900. small noted. Noted  
noted on Bac or Malt plates bac + Malt plates are v. crowded &  
"uniform" colonies - ca. 1000/plate. i.e. some cells inhibited by  
galactose?? Only large could be scored: 25,000 colonies. Ca 7  
most likely possibilities. Also isolate G<sup>R</sup> and G<sup>S</sup> colonies +  
test as contaminations? 7:1-7.

Lactose: 37 plates. Ca 800 on each plate, scoreable = 30,000.  
Almost 1 lac- per plate noted. 12 selected for further study. 7:11-30  
(1/1000)

Maltose 63 plates ca 55,000 colonies scoreable.

8 apparent Mal - noted.

7:41-50.

Total tests: 110,000

Reversion of Lac + Mal in W-1 and W-3.

Streak out papillae of W-1, W-3 and Y530 on Lac and on

Nov. 2, 47 Melt. EMB agar. Note that on original plate, W-3 had some papillated, some non-papillated colonies.

W-1. Lac. All Lac-. Papillae in streak?

Mal. alkali. All Mal-.

W-3. Lac All -.

Mal All -!

} Hold for  
restreaking

K

Results A6.

W-1. Revert for lac found. Verify and number W-33.

W-3-(Mal) all -; papillae are not Mal+!

W-3-(Lac) + and -. Verify. ✓ 1034

Mastard killing: 58-161.

9

Add 10 mg mustard to 10 ml 58-161, assay =  $3 \times 10^9$ , in phosphate  
M/10 pH 6.0. Dilute 1:100 at intervals and spread (on sucrose EM3 which  
is available). 1 ml. Colony count = survivors/ml  $\times 10^{-3}$

0 time assay =  $3 \times 10^9$

60 mins assay =  $2 \times 10^6$ . PS = 3+. for 60 mins.

Assume 10% survive for each 20 mins. at .1% in phosphate buffer.

Use .2% and 30 mins. treatment.

Nov. 2, 1947.  $S^S$

$S^R$

p1 - Disc small and large colonies of 58-161 on galactose ETIB plates of Expt. 7 into YP 25 ml. Incubating lit.

p2. Treat  $\epsilon$ .2% MN2 in phosphate buffer 40 min. Add 1 ml susp. to 25 ml YP for further incubation.

Incubate  $S^S$  on left (sinistral) side of incubator.

Expt. 8. Lac. 70 pl.  $\times$  20 col.  $\rightarrow$  1400 tests  $\rightarrow$  12 colo. mutants. more all lac +.  
ca 50% of colonies are lac - (is microcolonies?).

Pick test. 10-1.

Pick colonies from a Gal plate mix. with the untreated suspension. Pick 3 colonies to slants W30, W31, W32.

Malt.  $16 \times 40 = 640$  tests.

1:1 dimorphism of very dark and less dark colonies. Both are + and have a sheen. (Corresponds to above??)

Gal.  $23 \times 30 = 700$ . No mutants.  
some contaminated

$S^S$ . Lac.  $0.3 \times 100 = 6500$ . No mutants.

Malt.  $17 \times 80 = 1360$  No mutants.

Gal.  $13 \times 50 = 700$  No mutants.

11.

Nov. 3, 1947. Characterization of Mutants of Exp. 7.

A. Galactose mutants - 1st. streak on galactose EMBS.

7-1.	Majority of light purple colonies Pick to glucose slants.	B4	w-2.
7-2.	Gal± as in 7-1. No Gal+	B4	w-4
7-3	do.	B4	w-5
7-4.	Majority of illegible, v. small colonies. Dozens of heavy streaks, a few Gal+, larger colonies. Test on Lac: Lac - alkaline. throw out.		
7-5	as 7-1.	B4	w-6
7-6.	Do.	B4	w-7.
7-7.	As 7-2.	B4.	w-8.

B. Lactose mutants.

7-11.	Typical lac - .	not B-H-! <sup>n.g. in 16h.</sup>	w-9
7-12.	Do. Papilliforme		w-10
7-13	Do. Pap.		w-11
7-14.	Do. + two lac+ colonies.		w-12
7-15	Do.		w-13
7-16.	Do.		w-14
7-17	Do.		w-15
7-18	Do. Colonies smallest. lac+ in heavy streaks		w-16
7-19	Do. Weak utilization?		w-17
7-20	Do.		w-18.

7-21.	do.			
7-22	do.			
<u>7-MALTOSE</u>		what is the nutrition of these creatures?		
			w-19	
			w-27	
7-41.	Majority are light purple. Some Mal+.	BM		w-20
7-42.	Mal+.			
7-43.	do 41.	BM		w-21
7-44.	All Mal +	BM		w-22
7-45	do.	BM		w-23
7-46.	do.	BM		w-24
7-47	do.	BM		w-25
7-48	do.	BM		w-26.

Test units. by comparison of  $T(B+M) \approx T_0$ .

58-161.	$\frac{g}{g}^R$		
		w-28	
		w-29	

# Galactose inhibition

Nov. 3. Deminhibition on galactose was noted in 7.  
 $G^S + G^R$  were streaked out on EMB.

	$G^S$	$G^R$
Bac	++	++
Gal	++ only sl. smaller than.	++
Mal	++	++
Zulu.	± (slow)	± (slow).

The galactose effect was not reproduced here, nor in the plates streaked from cultivated  $G^S, G^R$ , in expt. 10. These were, however, a different batch. Test m:

galactose 1%	pyrrole 1%
galactose 1%	N2 amine 1%
galactose 1%	N2 Tane 1%

Dulcitol.

13

Nov. 3 '47.

K-12 ferments dulcitol very weakly. Grows into broth + compare. In 16 hours, dulcitol broth is vigorously fermented by 58-161.

A. from broth

B. from plate (a slow colony). } Both form only "weak" colonies.

Streak A. again. - Slow + as before!

?? Reduction of Methylene Blue ??

Nov. 4, 1947.

EMB - Gal 1% - agar 1½ %

- A. Peptone 1%
- B. " 0.3%
- C. N-2-Amine "B" 1%
- D. N-2-Tartrate 1%.

Shows out  $\text{G}^R + \text{G}^S$  colonies. Py.

AS.

- A No dimorphism
- B Large + small colonies. Not as marked as C.
- C Large + small colonies.
- D. no dimorphism.

4  
Nov. 24, 1947.

Mutant Rec.

Treat a single colony culture of  
58-161.  $\in \text{HN2}$  .2% for A) 5 min. B) for 30  
Lac mutant plating.

do. W-1 for galactose mutant plating.  
Incubate 24 hours before plating per current technique

A. 58-161 (treated) - 50  $\mu\text{l}$   $\times$  70 cols. on Lac = 3500 tests

No mutants

B. W-1.

35  $\mu\text{l}$   $\times$  100 cols on Gal = 3500 tests

No mutants.

Reduced sugar utilization.

Nov. 4, 1947.

See 14A. for Treatments.

Add ~~flask~~ ca  $10^{10}$  cells to various of acetone) to:

		A5	A6	A7	A9
58+161 :	1. 100 ml T(m) + (BM.)	±	+	-	-
	2. 100 ml T(m) + glucose .05 %	+++	-	-	-
	3. do. cellobiose	±	-	-	-
	4. do sucrose	±	-	-	-
	5. do sucrose 200ml.	±	-	-	-
	6. do. sucrose 200ml.	±	-	-	-
	7. do. 2-methylglucoside.	±	-	-	-
W-1.	8. do. sucrose TLB, 300ml.	±	-	-	-

- no change  
ditto

ditto.

abandon experiment on Nov. 9.

Nov. 5, 1947.

## U.-V. Killing Rate.

 $(10^8 \text{ cells})$ 

Hanovia lamp. 6" from plates. Spread 1 ml of grown cultures of 58-161 on EMB lactose plates. Irradiate as indicated.

t.	colonies
0	$> 10^5$
5 sec.	$> \cancel{10}^3$
10 sec.	112
15 sec.	ca 100.
20 sec	50
30 sec	28
1 M.	20
2 M	7
6 M	3.

There is certainly a break in the killing curve between 5 and 10 seconds, or else, the survivors of higher doses represent large clumps.  
10 sec. is a convenient irradiation time which has a  $\mu S = \log \frac{10^8}{10^2} = 6$ .

One "weak" bac colony noted on 10 second plate.

Streak + compare with subs.

$\beta$ -methyl glucoside

Nov. 4, 1947.

Sample from H.A. Hardy. (mp 106-108°)

1) 1% in EMB.	w-1	-	A6. dark spot - Residue. $\rightarrow$
	y53	-	
	y10	-	all - .
	58-161	-	
	w-22	-	

P5

2) .05% as C-source in T(m)+BM. 58-161 mol. P5.

	A6.	A7	A10.
1) $\beta$ -methylgl.	-		+
2) $\alpha$ -methyl..	-	++	-
3) $\alpha + \beta$	-	++	+
4) glucose	++	++	++
5) $\beta$ +glucose	++		+++

Nov. 6, 1947.

W-30 represents a series of Lac+ colonies which constitute 50% of the colonies found upon spreading mustard-treated 58-161 or lactose EMB. 12 colonies derived from the untreated inoculum were all Lac+. The inoculum was derived from a single large colony ( $\delta^R$ ) of a type constituting about  $\frac{1}{4}$  the colonies formed on a galactose-N2 Amine B-EMB plate (see Expt. 7). In Expt. 10, the ratio of Lac+:Lac- was:

	+	-	
plate 1.	18	10	
plate 2.	10	4	
plate 3.	8	6	
plate 4	12	16	
	48	36	84

A large fraction of both these types was found on each of the 70 plates of the experiment. (10)

- a) Pick 5 colonies from plate Expt. 7 in attempt to reproduce the expt.
- b) Pick + streak out 5  $\overset{\text{Lac}^+}{\text{colonies}}$  from Expt. 10 to determine stability of these Lac+: all plates show all Lac+
- c) Set up selection Expts. between these Lac+ and Lac- in broth.

On the Lac+ plates, W-30 forms broad flat colonies. These seem to engulf Lac- colonies which they may contact.



On maltose, two colony types are seen. Initially, they differ only in that some are less intensely colored on EMB than the others. Later they develop large lobes of gummy material which projects from the surface.



been noted.

On galactose no peculiarities had

C. Selection Expt.

A. Streak Lac+ colonies 2P6 into YP broth 25 ml.

B. Do. + 1 loopful of Lac- (W30) suspension.

Streak out duplicate lac plates for initial assay.

A: all colonies lac+ (Lac- < 1/100)

B: ditto.

		P7.
C. Streak out A	<del>+</del>	A 6: all +
D. B		A 6: all +
		all +
A2	Streak out P7.	all +
B2	" "	all +

The combination: W-30 and the population of Lac+ found on these plates does not seem to satisfy selective hypotheses. The Lac+ may be resistant to the hypothetical inhibitor. 58-161 (18-A) should be used instead.

D. Selection Expt. 11/10/47. Mix 1 ml broth culture of 18A1 with .05 ml similar suspension of W-30 from s lamb in YP broth. Streak out initially, etc.

①.	18D-A1	Streak 18A1 on EMB-Lac	A 11
	- B1	18A1 + W-30 on EMB-Lac.	ca 100:1 + : -

N1R	A2 +	all +
	B2 -	ca 20:1, + : 1

A12      B3 -      ca 20:1 + : 1

This selection expt. does not explain original findings.

Nov. 6, 1947.

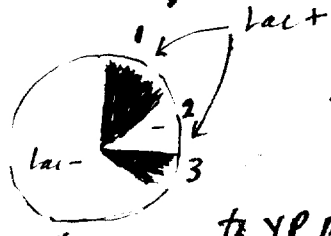
Spread .1 ml 58-161 culture on EMB-Malt and Lac plates.  
Incubate each for 10secs. in R. Smith's Hanover lamps.  
ca. 160/plate

Lac (10) Nothing noted on 9 plates. See below for 10th. \*

Malt (10) No mutants or sectors noted.

~~No mutants or sectors noted.~~

\* A colony was found of the following appearance.



This suggests further a delayed effect of the mutagenic agent.

Picks from each of the 4 distinguishable zones to YP Broth & incubate. When grown spread on and streak out.

				Lac EMB.
1	W-35	W-38	38	A10.
2	W-36	W-35	35	
3	W-37	W-37	37	1. Ca 50:1 Lac+ : Lac- . No evidence
4	W-38.	W-36	36.	of sectoring of the +.

2. Ca 50:1 Lac- : Lac+ . No sectoring

3. Ca 100:1 Lac+ : Lac. No sectoring.

4. All Lac- .

Streak out and transfer to slants.

Nov. 6, 1947.

P3. streak out Y53 on EMB-lactose - 3% peptone.

A6. Papillae well developed. Pick papillae to small H<sub>2</sub>O and streak out.

#P8. All of 20 plates pure Lac + exc. for 1 which was streaked from a mixed colony. ∴ Lac + is a stable reversion in most cases.

Nov. 8, 1947.

Spread  $10^8$  cells of a single cell culture of 58-161 from 'P' on maltose + lactose EMB plates. Irradiate each plate ca 10 secs. in Hanovia u.v. lamp at 6". Score at 36 hr.

Maltose. 24 pl.  $\times$  60 cols. = 1440 colonies tested. All colonies +.

Lactose 59 pl.  $\times$  ca. 150 scoreable colonies = 9000 tested.

12 plates showed 14 likely colonies. Some of these are clearly simply mucoid.

①, ② Mucoid.



4. Small white colony ✓

5. a c Stake out 3 compo.  
b

6. M 7. ✓

8. ✓ 9. M



→ no lac-  
m streaking

13. D. 14. O mucoid.  
alt + mucoid

- 3a 1:1 +: - (+) w50  
 3b almost pure (-) w48  
 3c 1:1 ± and - (-) w49

4. pure -. w39.

5. a. 20:1 lac+ : Lac- Purify + : w41.  
 b. 2:1 + : - →  
 c. pure -. w40.

7. pure - w42. (slight utilization?) ✓

8. 1:1 +: - The Lac- is w43.

10. a. 10:1 +: - +, w44  
 b. pure - - w45

11. a. 2:1 +: - (+) w46

b. pure - w47.

12. all +. Not mutant.

$\beta$ - $\beta$ -methyl glucose

Nov. 10, 1947.

Compare 58-161 and 17-1.

	glucose	$\beta$ -Me Gluc.	cellobiose	
17-4. (maltose)	± +++	— —	+ ±	— —
17-1. (maltose)	++ ++	— ±	± ±	— —
w-20.	± +++	— —	— —	— —

24 hrs., 36 h.

def. difference  
maltose contam?

415. do. only glucose is +++. others from - to ±.

∴ 17-1 is only slow utilization.

5 A16: p17

	$\beta$ -Me Gluc.	Cellobiose
17-41	± ✓	± ✓
17-4	± ✓	± ✓
w-20.	± ✓	± ✓
17-1A.	✗ + .	± .

Transfer 17-1A to  
similar series. → much slower  
on  $\beta$ -methyl than on glucose.

U-V. cur.

Nov. 14, 1947.

Irradiate 50 ml of 24 hr. ~~(at 37)~~ W-37 conc. to 10 ml. with washing. 3 ml / 25 ml quartz flask. 40 secs. at 6 in. from Hanovia Lamp, with manual stirring. Spread 1 ml on EMBS.

30 Lec, 20 Gal plates. All too dense (ca.  $10^5$ - $10^6$ /plate).  
(Autoabsorption rather marked!)

1 ml samples irradiated 5 secs more into synthetic minimal.

- glucose
- sucrose
- $\alpha$ -methyl glucoside
- cellulose

}

ellng. after 4 days.

15

Attempts at induced utilization  
of sucrose & other sugars.

Dec. 10, 1947.

Prepare suspensions, ca 10% / ml., of Y10. Inoculate 3 ml at a time 5 secs. in aqua-tg flesh rotated at hood of Hanovia uv lamp. 1/2 ml inocula into 50 ml 1/25 mol flasks containing 1 ml + sugars ± .05% (except glucose .15%) ± glucose .005% (g.) or 4P 10.

	9A 11.	2P 11.	P 6 11	9A 12	P 13	P 16.
1. Sucrose	±					
2. "	±					
3. " g.	++		- + No +	* No +.	±	(+)
4. " g.	++		-			-
5. Ref.	±					
6. Ref.	±				* No +.	
7. " g.	++		++			
8. " g.	++ ±	* No +	++			(+)
9. Celloolose	±					
10. " g.	++		- * No +	* No +.	(do. wss.)	(+)
11. d. M. gluc	-			+ No +.		(+)
12. " g.	+					-
13. Trehalose	+++	—				
14. "	+++	—				
15. Glucose	+++	++				✓
16. g.	+	+				✓
17. —	±	— baseline.				

Experiment terminated 12/24, without the recovery of any plus-variations in this series.  
Y55 (Salicin-plus mutant) should be tried on the beta-glucosides.

\* Streaks out on corresponding medium.

[ .005% glucose is apparently an excessive "boost". Use .001%. ] All these sugars were evidently somewhat broken up by their prolonged auto-fermenting.