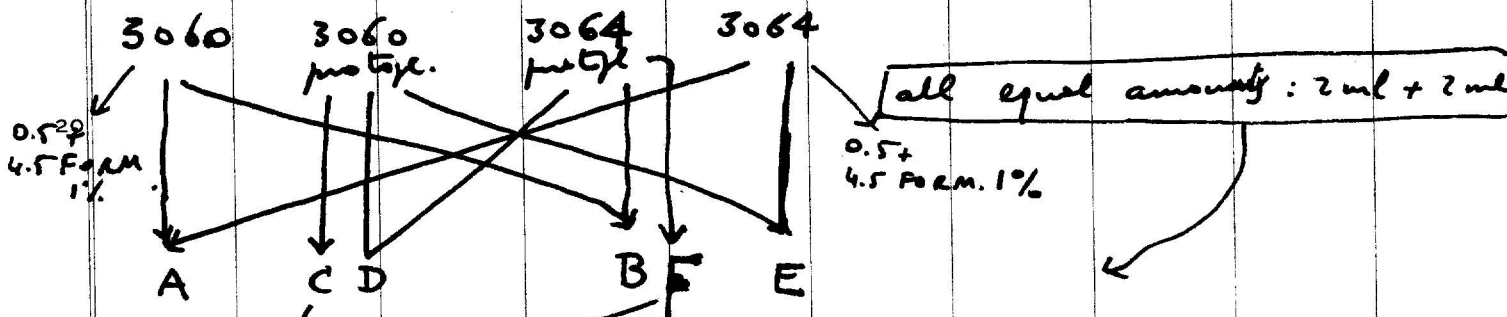


DATE: March 26.

	1	2	3	4	5	6	7	8	9	10
12.20	3060	3064	cult. → rotator 1+10 ml (from rotated cult.)							
13.20	1 ml	+ 9 ml L ₂	+ 10 ⁴ penicillin	→ protoplasts (5 ml.)						
15.50	Centrifuge, resuspend protoplasts in L ₂ medium without penicillin									
10			Also 14.50	3060, 3064, 1+10 ml to rotator.						
			15.50	centrif., resuspend in 5 ml L ₂ medium.						



16.40

C, F: 0.5 + 4.5 ml water; 1 ml → poured into agar.

A, B, D, E: 30' rotator → 0.5 + 4.5 ml water → min St B₁
 ↓
 60' " → 0.5 + 4.5 ml penicillin → agar inoculation.

⇒ Same.

0.5 ml. titrated of protoplast susp: 0.05 ml Form 10% added.

Microscope counts:

- 3060 : 400 × 10⁶ / ml
- 3064 : 940 × 10⁶ / ml
- 3060 protoplasts : 2.2 × 10⁷ / ml - no rods.
- 3064 " : 2.7 × 10⁷ / ml "

3060 protoplasts: agglutinated before centrif. After resus. centrif., pipetting up and down releases agglutination.

DATE: 3/29/58

REF: 1405-II

	1	2	3	4	5	6	7	8	9	10
Counts:			<u>30'</u>		<u>60'</u>					
			Sucrose	B, St	Sucrose	B, St				
	Exp. A		15	33	54	66				
	B		0	0	0	0				
	D		0	0	0	0				
		E	0	1	1	1				

Poured agar plates from C (3060): 278.
 F (3064): 72.

Conclusions: 3060 has reverted to F+. The same single colony i/plate has been used as in exp. 1403. Hence low numbers of recombinants.

Proportion of viable cells in protoplast preparations:
 3060: 1.3×10^{-5}
 3064: 0.3×10^{-5} .

DATE: 4/14/58.

REF: 1405-III

PROTOPLASTS.

Exp 1405-I is repeated, but after bursting in DW, mixtures are either plated or kept in the waterbath until completion of 60' since the beginning of the expt. In addition, a pulse of 10' is given to the mixture.

9.05 am. 1 ml 3060 overnight aerated + 9 ml broth. → Rot.
 10.30 1 ml 3060 expon. + 9 ml I₂
 12.10 coarse granular growth. Agitated, 10³ Pen U/ml added.
 14.40 centrifuged,

Also: 14.20 3064 2 ml + 10 ml Broth.
 14.40 Centrifuged.

3060 ♂, 3064 resuspended in $\left. \begin{matrix} 1.2 \\ 4 \end{matrix} \right\} 1 \text{ ml } I_2; = 10 \times$
 0.3 ml in water bath.

3060 ♂ 10x → 0.1 ml + 0.9 ml DW → 0.1 on B lac.

→ 0.1 + 1.9 ml formalin 1% → Count (all burst-out?)

3064 10x → 0.5 } X
 → 0.5 }

→ 0.1 + 1.9 ml formalin 1% → Count = 9×10^8 /ml

X and 3/4 (left in I₂ of 3060 ♂) incubated in water bath for 10', then: diluted 20x with I₂ medium warmed.

(X) → 0': 1 ml + 9 ml DW → plate 0.05 and 0.01 on min St B₁.
 ↓ + 10 ml broth, water bath. (= 0/60') → plate $\left. \begin{matrix} 0.05 \\ 0.01 \end{matrix} \right\}$

10' water bath: same as above → = 10
 = 10/50.

20' " " "

30' " " "

45' " " "

60' " 1/10 DW → = X-60 (pl. 0.05, 0.01)

→ + 10 ml broth → = 60/0 (pl. 0.05, 0.01)

DATE:

REF:

1405-III

♂ - Counts of viable cells in protoplast suspension:

From suspension obtained after cult. -

0.1 + 0.9 DW, → 0.1 EMB lac plate. 246 col.

From same susp. del 40x in I_2 , → 0.1 EMB lac. 4 col.

↓
30' incub w. l. → " (dried up)

↓
60' " → " 67 col.

The initial No. of cells which were protoplasted is unknown. but is of the order $2-5 \cdot 10^8$ /ml. Hence fraction surviving is of the order 10^{-5} .

	0	10	20	30	45	60	minutes
	X-0	X-10	X-20	X-30	X-45	X-60	
0.05 ml	0	191	∞	∞	∞	∞	
0.01 ml	0	41	362	~800	~720	760	
	0/60	10/50	20/40	30/30	45/15	60/0.	
0.05 ml	11	178	∞	∞	∞	∞	
0.01 ml	0	21	244	434	~900	609	← should be multiplied x2 to five figures comparable to X

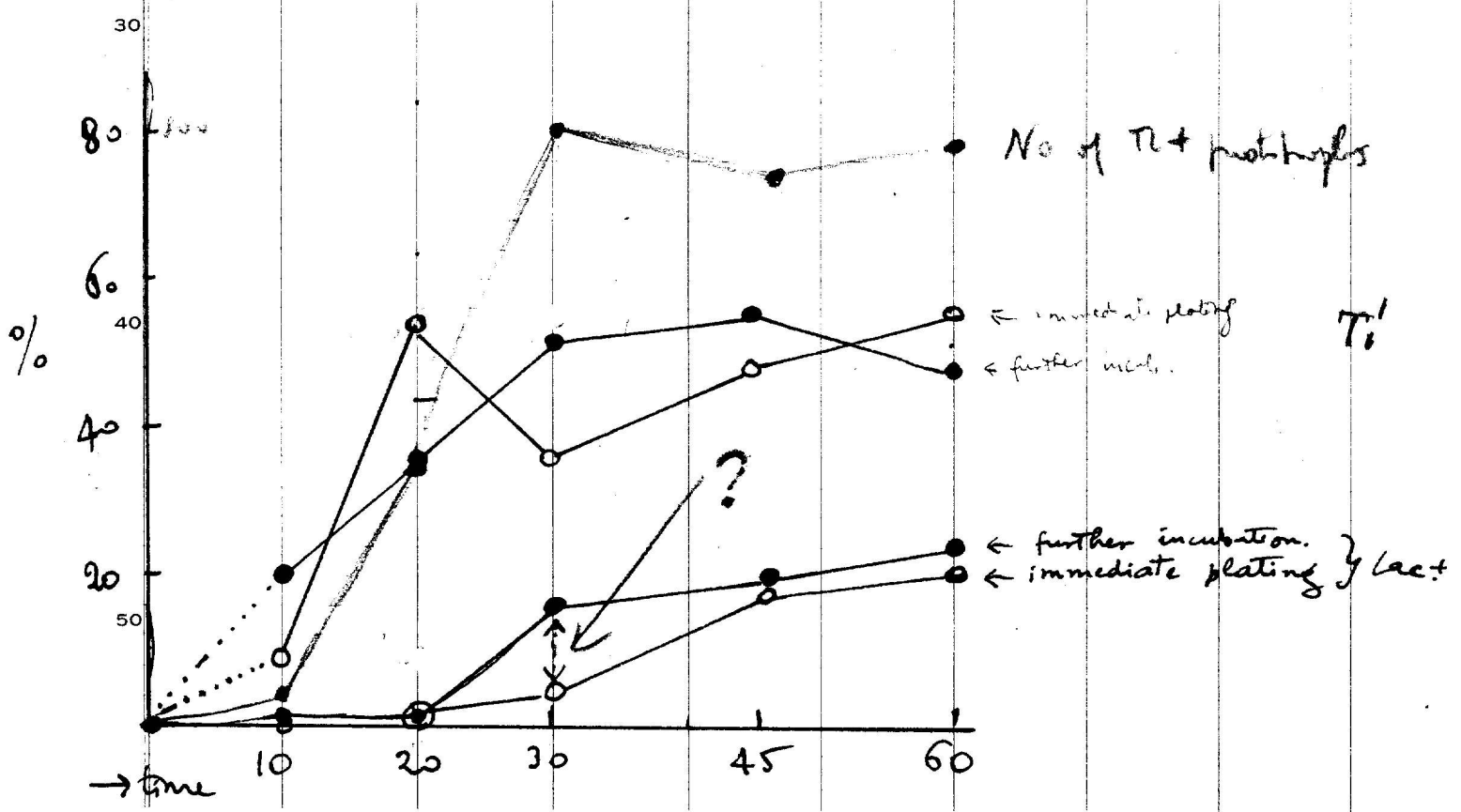
50

addition of water
 Before: After 4/17/58.

REF: 1405-3.

	1	2	3	4	5	6	7	8	9	10
TIME	0/60		10/50		20/40		30/30		45/15	60/0
Gal+	0/12		0/71		0/50		0/50		1/50	1/50
Lact	4/12		2/71		1/50		8/50		10/50	12/50
π_1^{10}	8/12		7/71		18/50		26/50		28/50	22/50

	X-10	X-20	X-30	X-45	X-60.
Gal+	0/90	0/50	0/50	2/50	0/50
Lact	0/90	1/50	2/50	9/50	10/50
π_1^1	4/90	27/50	14/50	24/50	29/50.



DATE: April 14.

REF: 1405-IV

Repeat of 1405-II: Crossings between protoplasts and non protoplasts.

10.30 a.m. 1 ml 3060, 3064 + 10 ml L₂.

11.50 { 3060 poorly grown, agglutinated.
 { 3064 well grown, 3-500 · 10⁶/ml.
 Penicillin 10⁴ units added.

13.35 cultures of 3060, 3064 started: 1 + 9 ml broth

15.00 3060, 3064 cultures and 3060, 3064 protoplasts centrifuged, resusp in L₂: 3064 + 3064 ♂ 10 ml, 3060 ♂ 2 ml, 3060 ♀ 5 ml.

15.35 Cross:

	A	B	D	E
3060	0.5	0.5	-	-
3060 ♂	0.5	-	0.5	0.5
3064 ♂	-	0.5	0.5	-
3064	0.5	-	-	0.5

3060

0.5

0.5

-

-

3060 ♂

~~0.5~~

-

0.5

0.5

3064 ♂

-

0.5

0.5

-

3064

0.5

-

-

0.5

- water bath.

Algo:

Viability Test

C: 3060 ♂: 0.1 ml + 0.9 water → 0.05 Blue
 → 0.01 Blue

F 3064 ♂ " " "

16.40.

From water bath, A, B, D, E:
 0.5 + 4.5 ml water → 0.05 1/10 DW → 0.05 ml
 must B, (1/10)

Algo, on agar plate: A, B, E:

0.05 from test tube

0.05 1/10 dil. in L₂

0.05 1/100 dil in L₂.

D: 0.05 from test tube.

DATE:

REF: 1405-IV.

1	2	3	4	5	6	7	8	9	10
Viability tests:									
				C	0.05	44 col	3060 ♂:		
					0.01	13 "			
				F	0.05	168	3064 ♂.		
					0.01	145. ?			

Microscopy: C 2-3 protoplasts per small square, many of which
ghosts. = $5 \cdot 10^7$ /ml.
F 8-9 $160 \cdot 10^6$ /ml.

3060 ♂: Viable $2 \cdot 10^{-4}$;
3064 ♂: " $2 \cdot 10^{-4}$.

	A	B	D	E
Sucrose B ₁				
1/1	∞	∞	~1000, small	∞
1/10	∞	∞ irreg	not met	∞
1/100	2-5000	∞ irreg	m.m.	~500
Sm B ₁				
1/1	∞	32	2	~2000
1/10.	~2000	0	0	162

Note. 3060 is not prototrophic, therefore crosses on Suc B₁ are valid.

DATE:

4/19/58

REF:

1405-4

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Summary of pyrogatory.

♂	♀		min sub B ₁			micro B ₁		
			gal+	lac+	T ₁	gal+	lac+	T ₁
-	-	A	0/50	8/50	22/50	26/50	22/50 [⊕]	
-	♂	B	2/50	10/50	27/50	24/50	-	
♂	♂	D	-	-		24/50 [⊕]	28/50 [⊕]	
♂	-	E	2/100	13/100	58/100	0/50	9/50	26/50

Also:

From D: streaks on EMB lac.

⊕ many mixed.

Later note: W3060 showed erratic growth on B₁ mediums. ? question of a usable auxotrophic marker in this strain. Should be reviewed in the fall.

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DATE:

PROTOPLAST CROSSES.

REF:

1405-5

Repetition of 1405-4.

3060, 3064 ♂ as in experiment 1409-2, but diluted 5x with L_2 . 3060, 3064 rods in L_2 , 2x concentrated, from overnight rotated cultures.

	1	2	3	4	5	6	7	8	9	10		
		♂	♀	Plated on Suc B ₁		Suc B ₁ St		B ₁	B ₁ St			
M	♂	♂		$\frac{1}{1}$	0.02; $\frac{1}{10}$	0.02	$\frac{1}{1}$	0.02	$\frac{1}{1}$	0.02	$\frac{1}{1}$	0.02
N	♂	Normal		$\frac{1}{100}$	0.02		$\frac{1}{100}$	0.02	$\frac{1}{100}$	0.02	$\frac{1}{100}$	0.02
O	Norm	♂		$\frac{1}{1}$	0.05		$\frac{1}{1}$	0.05	$\frac{1}{1}$	0.05	$\frac{1}{1}$	0.05
P	Norm	Norm.		$\frac{1}{100}$	0.02		-		$\frac{1}{100}$	0.02	$\frac{1}{100}$	0.02

Mixtures for crosses set up in equal amounts at 12.20. After 60' incubation in water bath, dilutions (with L_2) and platings.

Viability tests of protoplasts: suspensions employed in crosses diluted 1+10 ml H_2O , \rightarrow $\left. \begin{array}{l} 0.01 \\ 0.05 \end{array} \right\}$ B lac.
(2 hrs after preparation of suspensions)

← Drop of flamed alcohol

Counts: 0.01 # 3060: 70 col.
0.01 # 3064: 136 col.

Microscope counts on protoplasts: (after stay at room temp):
♂ 3060: $52 \cdot 10^6$ /ml (but many rods!)
♂ 3064: $392 \cdot 10^6$ /ml no rods.

Viability of protoplasts: $1.6 \cdot 10^{-3}$ for # 3060 ♂
 $0.3 \cdot 10^{-3}$ for # 3064 ♂.

(after a 2 hr stay at room temperature; rods may have increased)

DATE:

REF: 1405-5

	1	2	3	4	5	6	7	8	9	10
		<u>Plate counts</u>								
♂ ♀ ♂ ♀	(M)	Sucrose B ₁ 0.02: ~2000			Suc B ₁ ft 0.02: ~800		min B ₁ 0.02: many small and 158 large		min st B ₁ 0.02: many small col.	
		$\frac{1}{10}$ 0.02: 172 + many smaller colonies.								
♂ normal	(N)	302			374		255		314	
Normal ♂	(O)	10 ⁴ ?			10 ⁴ ;		~1500		~500 large + many small only.	
Normal Normal	(P)	~3000					~3000		~2000.	
Counts:	3060 ♂	many small ones (~2000).								
	3064 ♂	no growth, except for 1 colony								
		Picking								

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DATE:

4/19/58.

REF:

1405-6

Repeat of exp 1405-3 to test further difference between immediate and non immediate plating on Lac at 30' (see question mark in previous graph).

3060 3, resuspended in L_2 10x conc.; pour with 3060 overnight rotated culture.

Mixtura in equal amounts. Also, viability test of 3060 5: DW $\frac{1}{10}$ \rightarrow 0.02 on EMB Lac. (465 colonies counted).

From 3060 3 x 3064, 10' pulse then dilution $\frac{1}{100}$ in prewarmed L_2 , and further incubation in w.c. for 25, 30, 35'.

After such times, dilution in DW 1+10, incubation for 10', then addition of 10 ml Penafloy 2x and:

A) immediate plating

B) further incubation for until completion of 65' from end of pulse, then plating.

Platings: 0.1 & 0.02 on min St B₁.

By mistake, 35' has been kept at room temperature.

	25'		30'		35'	
	A	B	A	B	A	B
	NON INC.	INC.	NON INC.	INC.	NON INC.	INC.
0.1	141	88	194	146	261	232
0.02	23	10	27	19	46	35

DATE:

4/23/58.

REF:

1405-7

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OSMOTIC SHOCK ON RODS

3060, 3064 o.z.c. resuspended in 5x conc. with T_2 medium.

Mix in equal amounts for a 5' pulse in water bath.

Dilute $1/100$ in T_2 medium, warmed.

After 10'; 20'; 30' dilute $1/10$ in DW, incubate 5', add 2x Penicillin in equal amounts and:

A) plate 0.05 on min stB₁

B) incubate further until completion of 50' since dilution in $T_2/100$, plate 0.05

Plate counts:

1-2000 colonies in each except for 10'A (≈ 100) and 20'A (≈ 300).

Gal

lac

 T_1

10' A

1/50

B

22/50

20' A

0/50

B

11/50

30' A

4/50

B

13/50

DATE:

REF:

1405-8

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CROSSES ♂ × ♂.

2323, 3064 protoplasts 2^h30' incub. (1 ml + 9 ml from o.r.c.).Resuspended in T₂ medium at 10 × conc.10 ~~0.5~~ Mixed equal amounts, then diluted 1/100 after 10' pulse in T₂ medium. Incubated 50', then spread on min B₁ & sucrose B₁:0.05 from untl. & 1/10, 1/100, 1/1000 ^{and 1/10,000} dilutions in T₂ medium

From parental suspensions: → 1/10 dilt. water → 0.05 B lac

→ 1/100 T₂ → 0.05 on suc B₁↓
1/100 T₂↓
1/100 T₂↓
1/10 T₂↓
1/10 T₂

→ 0.05 sucrose Penellay

→ same.

and microscopic counts

Microscope counts: 2323 ♂: 4.2 × 10⁹/ml.3064 ♂: 3.2 × 10⁹/ml10⁻⁵10⁻⁶

3064

10⁷

15

2.10⁸/ml viable

2323

320

10

= 6.10⁸/ml viable

Viability tests:

3064

~ 10,000

2323

~ 3600

Crosses:

Counts:

2323: clean

3064: many small colonies.

Crosses: as control 3064.

100 bigger colonies in untl.

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1

"

1/10.

Discarded.

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DATE:

4/15

CULTURE PHASE.

REF: 1406

1	2	3	4	5	6	7	8	9	10
9 am.		3060,	1 ml + 10 ml.	broth			Photometer, 650 m μ		
							0.29		
		3064,	1 ml + 10				0.37		
			0.5	"			0.32		
			0.2	"			0.20		

10 a.m. photometer readings. as above.

Alyo: saturated (overnight aerated) broth cultures:

3060 Sat 0.51

3064 Sat. 0.80.

All broths ^{reducedly} taken to density equal to that of 3060 exponential, by diluting with broth. (3064 ^{EXP.} 0.5 ml + 10, dil 8 ml + 2 ml); 3060 Sat 5 ml + 6 ml broth, 3064 Sat 3 ml + 15 ml).

Mixtures in equal amounts, 2 ml + 2 ml to waterbath:

3060 sat	3060 Exp.	3064 sat	3064 Exp.

A
B
C
D

x
x

x
x

x

x

x

x

10' pulse, then dilute 1/100 warm broth; → 0.05 with St B₁

↓ stay in water bath 50' → 1/10 DW, 1 ml flooded St B₁ plates.

0.05 with St B₁

1/10 DW → 0.05 "

0.01 except for A.

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DATE: 4/17/58.

REF: 1406

	1	2	3	4	5	6	7	8	9	10
	After 10'									
			A	B	C	D				
		0.05	482	293	482	320				
10	flooded with line of 1/10 oil.		~2000	~2000	~2000	~2000				
	After 60'									
		0.05	many	many	many	many				
20		1/10 0.05	288	~300	~500	~500				
		" 0.01	not done	59	91	113				
30										
40										
50										



used for picking and testing.

DATE:

CULTURE PHASE

4.19.58

REF:

1406

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SUMMARY

A B C D

♂

1 1 exp exp

10

♀

1 exp 1 exp.

10'

Malt

0/50 0/50 0/50 0/50

Gal +

0/50 0/50 0/50 0/50

Lac +

1/50 0/50 1/50 0/50

20

T_i'

12/50 9/50 10/50 8/50

60'

Malt +

0/50 0/50 0/50 0/50

Gal +

6/50 5/50 2/50 6/50

Lac +

20/50 16/50 12/50 21/50

30

T_i'

30/50 30/50 22/50 29/50

40

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DAT

LEONARDASI, ESTHER M., University of Wisconsin, Madison, Wis.--Fine structure of the Gal locus in *Escherichia coli* B-12.--A distinctive recess (one of a group of closely linked loci separable by recombination) has been assigned to each of ten independently occurring Gal⁻ (galactose nonfermenting) mutants. All the mutants were obtained after UV irradiation except Gal₃ which was spontaneous. Allelism was tested by large scale matings and by transduction analysis.--From phage-linked transduction via HFT λ , where practically every phage particle may result in a transductional event, produced heterogenetic clones when donor and recipient bore distinctive recess. The heterogenotes were unstable, segregating the two input and two crossover classes (Morse, Lederberg and Lederberg 1956). Two cistrons (cis-trans position effect groups) had been demonstrated: trans heterogenotes with members of the same cistron are phenotypically mutant. Galactokinase was missing in mutants of one cistron (Gal 2,8) while UDP transferase activity was absent in the other (Gal 1, 4, 6, 7), a defect corresponding to human congenital galactosemia, (Kalkar, Kurahashi 1957). Gal₃⁻ and Gal₉⁻ are genetically cistronic with both of the foregoing groups; they have not yet been successfully analysed for their enzymatic defect. Among two hundred new mutants, no recess identical with the first ten have recurred. Most have been assigned to either of the first two cistronic groups, but a small number which would represent a fourth genetic group have given only normal galactose-positive heterogenotes with every standard Gal⁻ tested. A few mutants were not transmissible by λ but were found to carry other modifiers including that of hexose metabolism which obscure their relationship to the Gal⁻ group.--An attempt was made to map the Gal markers in linear sequence by their relationship to the Ip locus. Intercrosses of Hfr M⁻ Gal_x⁻ x F⁻ Gal_y⁻ were made on a medium which selected efficiently for M⁺Gal_x⁺Gal_y⁺ recombinants, and these were then scored for the segregation of an Ip marker. However, the linkage of Gal-Ip proved to be not close enough to compensate for the perturbations of segmental loss from the Hfr parent, and the data were inconclusive.

10th International Congress of Genetics, Montreal, 1958.

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1407-2

EML

Lysogeny Variable in Gal⁺ + Lp⁺
yield at 60 mins

Standard protocol: back cross

EML
1407-2

rec parents
together off 20/58

summary

	A	B	C	D	E	F
V ₁ Lac Lp	27	60	49	44	24	0
Gal-	1	1	1	1	1	1
Σ	28	61	50	45	25	1

Σ / % Gal⁺ 28/14 61/51 50/50 45/47 25/27 1/90.7

	A	B	C	D	E	F
Gal+	0	0	0	0	0	0
Gal-	0	0	0	0	0	0
Σ	0	0	0	0	0	0

Σ / % Gal⁺ 0/0 0/0 0/0 0/0 0/0 0/0

Σ / % Gal⁺ 28/14 61/51 50/50 45/47 25/27 1/90.7

Intra:		# Strains unlocked			
Gal	Lp	A	B	E	F
0	0	90	102	62	46
0	1	0	4	3	12
1	0	74	35	71	5
1	1	12	49	67	49



19

May 7th, 1958

REF:

1408/2

	1	2	3	4	5	6	7	8	9	10
1	#	W3908	(F-4 ⁺ + tryp Gal ₆ S ^R)							
2		W3870	Hfr ₂ B ₁ ⁺ Lp ^R .							
3										
4										
5										
6										
7										
8										
9										
0										
1										
2										
3										
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o.r.c. : ml: 0.5 ♂ + 1.0 ♀ - incubated for 1 hr.
Plated on 5 Gal Sm B₁ ; 0.05 undil, 1/10, 1/100
1/1000, 1/10000

Controls : 0.05 of undiluted cultures.

After 40 hrs :
undil : > 1000, Gal + 4 Gal -
1/10 : 56 Gal + 70 Gal -
1/100 : 4 Gal + 3 Gal -
1/1000 : 0

Controls :
W3908 : 3 col.
W3870 : 0 col.



19 May 10th 1958

REF: 1408/3

	1	2	3	4	5	6	7	8	9	10
	Gal - Try linkage									
♀	3908 s ^R Gal _s try - Lp ⁺ .									
♂:	1 W 3367 Lp ^R M-					Media:				
	2A: W 3060 + - Th?					M Gal try p B ₁ Sm				
	2B: W 3870 R - Th?					S Gal Sm B ₁ .				
	3: W 3848 s P Mals					Cross:				
	4: W 3752 s					.5 ♂ + 1.0 ♀, 1 ^R 37°				
	13: W 3782 s M-					.05 of 10 ⁰ , 10 ⁻¹ , 10 ⁻² .				
	21: W 3898 + L- (481 M).									
	yield 10 ⁰	10 ¹	10 ²	Sel ⁺ try ⁺	Gal ⁺ /total	Lp ⁺ /Gal ⁺	Lp ⁺ /Gal ⁰			
1.	0	0	0	-						
2A	∞	154	10	32%						
2B	∞	63	-	25%?	85%	17%				
3	14	0	0	18%						
4	39	0	0	16%						
13	36	miss.	0	4%						
21	17	0	0	245%						

DATE:

5/15/58

REF:

1408/4

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Gal TIMING - Hfr₂

2 ml O₁ v 3370 + 7.5 ml Pen; 1 ml 3908 + 7.5 ml Pen, 1 ml ^{for} not

Ⓐ

Spin ♂ & ♀, resuspend in Pen 3.0, 0.3 μ take respectively
 Mix 0.1 + 0.1, at 37°, after 1' add 10 ml warmed
 broth - (1/50 dil)

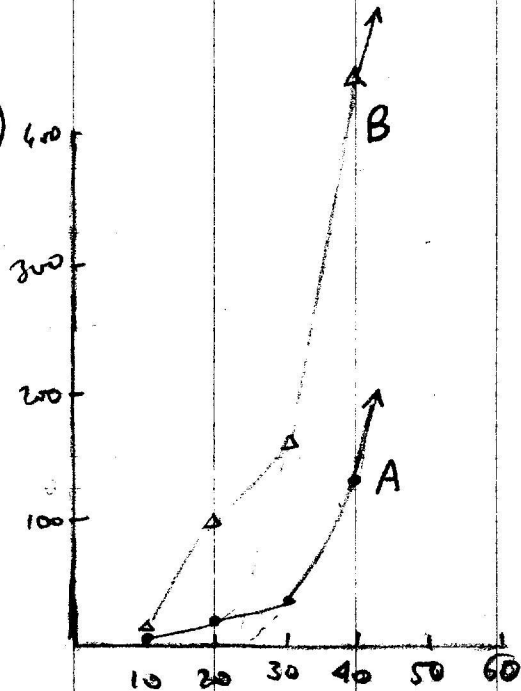
After 10', 20', 30', 40', 50' withdraw 2 ml sample, chill,
 blend, plate on Sgal Jim B, D (Jim B), Mgal Jim B, Tryp.

Ⓑ

Spin ♂ & ♀, resuspend in fresh broth, v .5 ♂ + 5 ♀ + 5 Pen
 Rest: same.

Plate counts D (Jim B) 60

	A	B
10	7 4	33 12
20	18 -	91 116
30	31 31	174 149
40	140 112	418 554
50	(1200) 823	∞
60	(2500)	∞



Entrance of Tryp at 20' ?

Probably data insufficient to answer
 the point

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	1	2	3	4	5	6	7	8	9	10
1		5' Gal for B, Counts								
2										
3										
4										
5			A			B				
6			Gal+/Gal-	Tot	Gal+%	Gal+/Gal-	Tot	Gal+%		
7										
8										
9										
0		10'	2/3	4		2/10	12			
1		20'	missing			17/99	116	12%		
2		30'	0/31	31	0%	5/146	149	3.3%		
3		40'	18/94	112	16%	114/410	554	20%		
4		50'	447/376	823		too many				
5		60'	too many			"				
6										
7										
8										
9										
0										
1										
2										
3										
4										
5										
6										
7										
8										
9										
0										

2
 17
 5
 70
 11
 35
 110
 48
 9.1
 5.7
 1.8
 1.2



19 May 16th, 1958

REF: 1408/5

	1	2	3	4	5	6	7	8	9	10
1										
2	W 3870	repy culture,	2 ml +	7.5 ml	broth	→	1	lv	not.	
3										
4	W 3908	"	1 ml +	3.5		→	1	lv	not.	
5										
6										
7										
8	Spun, resuspended, warmed -		Matry mixture.							
9	2 ml ♂ + 10 ml ♀ + 10 ml broth		in flask.							
10										
1										
2	<u>Controls</u> :		♂	1 ml + 10 ml broth	→	1 ml + 1.5 water	→			
3										
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♂, 1 ml + 10 ml broth → 1 ml + 1.5 water →
→ .05 & .1 plated

♀, .5 ml + .5 ml broth → + 1.5 water → .05 & .1

Plate result: .05 diluted ♂ + .05 diluted ♀, as above. .1 + .1 also

Sampling from matry mixture: by means of warmed 1 ml pipette, to 1.5 ml chilled water, then blended, and amount from .05 .1 to .05 plated on

D (for B, MetA)
B (Gal for B, MetA)

Times : 10', 15', 20', 22', 24', 26', 28', 30', 32', 34', 36', 38', 40', 43', 50', 50' -

Note : 32' is actually 33'10"
34' ~~33'~~ " 34'30"
36' ~~35'~~ " 36'10"
50' 50'35"

From 38th or 36th minute (possibly also 35th), contamination from waterbath through the pipette. Exp. discarded.

