

1413/10

	1	2	3	4	5	6	7	8	9	10	
1			Counts on D (from B ₁)								
2											
3											
4			A	B	C	D					
5											
6											
7											
8		10	170	0	112	3					
9											
0		20	~5000	21	~10,000	67					
1		30	~12000	128	~2.10 ⁴	167					
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Counts on D (from B₁)

A B C D

10 170 0 112 3
 20 ~5000 21 ~10,000 67
 30 ~12000 128 ~2.10⁴ 167

B (Gal, sum)

A B C D

Gal. 10 170 0 112 3
 20 20 21 35 67
 30 30 128 167

Gal. 2 isolated and tested

% from D (from B₁)

Gal+ lact+ T_i

A 10 0/100 12/100
 B 20 0/25 0/25
 B 30 1/100 19/100
 C 10 1/101 7/100
 D 10 0/3 0/3
 D 20 0/75 0/75
 D 30 0/100 18/100

Gal + Jun^R colonies from A30 and C30:
all Gal + Lac+ (with one only exception).

A30 C30

T_1^1 20/30 36/45

T_{1p}^1 16/30 33/45

No clearcut difference in contribution from σ parent between
A and C, i.e. Jun^R pretreated and non pretreated with ϕ 's.
tested out of ether

Protective Blending

11414

May 1 1958

REF.

Purpose: Can one blend in media so supplemented that pairing within the F⁻ state will not be interrupted. (Hyp. of progressive pairing vs. progressive entry).

System W3060 ♂. W3064 ♀. look for recovery of bac.

Pulse: 1 minute. ~~to~~ Dilute incubate for 12 minutes post pulse.
Chill in cooled water, 42, or serum broth. Blend in white grounds.

3 PM. ORC: W3060(3X) + W3064(30X) ~~to~~ .1 ml each 1 minute 37%.

add 10 ml warmed broth*, 1/100 in broth, incubate 12 minutes. (several tubes)

A. Add DNP to 10⁻³M

B. Add 1/10 chilled broth

C. Add 1/10 chilled 20% serum

Blend all 3 cultures.

AA: ~~to~~ Dilute (1/100) + plate

BA. D+p.

CA. D+p.

AB: incubate 20 minutes, d+p.

BB. Chill hours,

CB Chill hours

AC: ~~incubate 20 minutes, dilute,~~
~~incubate 20 minutes, plate~~

warm 20 minutes, plate

warm 20 min. plate

AD. incubate hours, dilute,
incubate 20 minutes, plate.

CC ~~incubate~~ incubate 20 minutes, d.p.

D. Add DNP 10⁻³M. Do not blend.

DA. incubate 20 minutes, d+p. (blend?)

DB incubate 20 minutes, d, inc. 20 min., plate.

6.30 - use uv!

Need: Minc B, for scoring

* broth = salts at pH 6.2
glucose - asparagine
per Fisher + Hayes.

May 1st 1958

1414

Protective blending.

3060, 3064, conc. 3x and 30x resp. : .1 ml each.
1 minute pulse at 37°; add 10 ml BGA warmed, dilution
~~from~~ 1/50 in BGA, incubate 12 minutes.

A. dilute 0.1 + 9.9 chilled, blend & plate 0.05 on min. ft B₁.

B. " " BGA warm, incubate 30', blend & plate.

C. add DNP (dinitrophenol) 1/50^{*}, incubate 5', blend :

C₁: remove sample of 0.1, dilute blend & plate.

C₂: incubate 30', dilute, blend & plate.

C₃ incubate 30', dilute, incubate 30', blend and plate

D. add DNP 1/50, incubate 30', dilute in ~~optimal~~ warm BGA

D₁: blend & plate sample

D₂: incubate 30', blend & plate.

* from 1/20 master solution. Final dil. wanted 1/1000

Plate counts:

A	B	C ₁	C ₂	C ₃	D ₁	D ₂
1, 0	0, 1	0, 1	0, 0	0, 0	1, 0	1, 0

too low. Need to do oxams!

Supercell by F⁺

14/4C.

Phase 6 May 1958.

REF: B

Pulse W3064 x W3870 4 minutes (to t=4). Dilute 4:100. Incubate in BSA
1 0.2ml (40x) 0.2ml

2
3 at 37° to t=12m. Parallel (for C) W3064 only.

at
t=12
4
5
6 A. Blend sample and plate (plate = 1:100 diln. in chilled water, spread 0.1 ml DTherm
7 (max entry of base)
8 plate cell.)
9 (successful recomb.)

10 B. Reincubate and plate at t=60.

1 C. To W3064 del. add ¹³⁰⁶ W3062 plate at t=60

W1306 = F⁺ M V₁ V₆.
made s.r.i. 6/17/57.

2
3
4 D. after blending A, add ¹³⁰⁶ W3062 Plate immediately

5 E. " at t=24

6 F. " at t=60

7
8
9
10 G. " at t=60.

(continued mating control)

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Re-entry stimulated by $F^+ \sigma^{80}$?

1414B

3 May 1958

REF: 1414 sec C

? Might reentry be stimulated by adding $F^+ \sigma^{80}$

Pulse W3064 + W3890 (ORC) 4 minutes (to $t=4$). Dilute 4:100 in
(20x cultures) .2ml .2ml

B5A to $t=12$ m. 37°

A. Blend sample and plate (plate = 1:100 in chilled water and spread
at $t=12$. (check non-entry of *lac*; vs. plate recombination; at time).
0.1ml on D.B. sm.)

B. Unblended sample. plate at $t=35$ m. (successful mating)

C. W3890 + W3064 s/ σ^{80} W3890. Add W6 at $t=12$. Plate at $t=24$, $t=32$.
(check W6 x W3064 crossing)

D. A + W6 (= .5ml 10x ORC W6 per 10ml sample). Plate at $t=12$ (D0);
intended $t=22$ but did C instead; $t=32$ (D20).

Df D20, shows an effect of W6 addn. it is open to objection of further
Hfr x F⁻ mating from $t=12$ for 20 mins., allowing of some injection of
lac. Alternatively, (more interesting) (a) W6 facilitates re-pairing of
preinjected genome; (b) prophage facilitates mating of W6!
Intention: look for *lac* reentry by replica plating

		No <i>lac</i> ⁺	W6 <i>lac</i> ⁺
A	35, 30	No <i>lac</i> ⁺	No <i>lac</i> ⁺
B	~200;	42 <i>lac</i> ⁺ / 400	✓
C2:	no protopho.		✓
D0:	34, 37	No <i>lac</i> ⁺	D20: 59, + 1 <i>lac</i> ⁺⁺ ; 13 probable <i>lac</i> ⁺ <i>cat</i> ⁻ .

∴ A shows early TL⁺ B full entry D is effect of reentry of *lac*.

Replica plate to Strac B, : why ~10% *lac*⁺? May be counting *cat*⁺.

Reincubate. an effect is indicated but The objection may still hold that further
mating has taken place. *lac* scoring m.g. Stroke out! of A, B, D0, D20, .

(over)

Summary - from individual shuffles on λ lac:

A: 6 lac⁺/60

B: 5 lac⁺/70

C - no prototrophs

D-0: 0/30

D-20: 8/120 lac⁺

The addition of W6 and further incubation give lac⁺ recombinants.

1414B

A

A

B = $\frac{5}{70}$

May 7 19 58

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B₁

DO = all Lac⁻

D 20 = $\frac{8}{120}$ Lac⁺

May 7 1955

a

REF:

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+

D20



19

REF:

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141413

D0 - versus D20 shows lacerity. Is this due to

- (1) remaining of $Hfr \times F^-$ despite blending + dilution
- (2) $F^+ \times F^-$ owing to preferentialization of F^-
- (3) Resumption of pairing in interrupted $Hfr \times F^-$ under influence of F^+ .

suggest: use W1306 ($H^- F^+ V_1^r V_6^r$) to distinguish (2).

control: reincubate after blending + dilution to distinguish (1).



19 May 9th, 1958.

REF: 1415/3

	1	2	3	4	5	6	7	8	9	10
1	W 3060 (Esther's transfer) : 1 ml + 7.5 ml Penney, 8 cult.									
2	W 3064 " : 0.5 ml + " 10 "									
3	3 hrs rotation. Spun, resusp. in water 1 ml, pooled different									
4	cultures, distributed each strain in equal amounts in									
5	two tubes, spun again, each ^{strain on} tube resusp in 0.5 ml.									
6	(Concentration about $80 \times$) of D(aspartic) *, and the other									
7	in D(aspartic + azide 10^{-4}).									
8	* : to 10 ml of D(M) : .1 of 20% glucose and									
9	.02 of 10 mg/ml aspartic acid. pH 7.0.									
10	Azide and non azide parents incubated at 37° for									
11	5', then:									
12	(AB) Parents in non-azide D(asp) mixed, 0.2 ml + 0.2 ml									
13	and pulsed for 2½ minutes, then diluted as follows:									
14	A. .1 ml in 10 ml D(asp)									
15	B. same + azide .1 of 1% solution.									
16	(CD) Parents in D(asp) <u>azide</u> 10^{-4} , 0.2 ml + 0.2 ml,									
17	and pulsed for 2½ minutes, then diluted as follows:									
18	C. .1 ml in 10 ml D(asp)									
19	D. same + azide .1 ml of 1% solution.									
20	(E ₁) Unpulsed recombination control 10^{-2} : .05 ^{of each} parents, $80 \times$ conc.									
21	to 10 ml of D(asp) at 37°, 40'. Dilute 1/100, plate .05.									
22	(E ₂) Unpulsed recomb. control 10^{-4} : .05 of 1/100 dilution of									
23	each parent to 10 ml of D(asp) at 37° for 40'. plate .05									



A and C : after 40' from pulse, dilute $\frac{1}{100}$ in water, plate .05 in minifm B,

B and D :
 B_1, D_1 : .1 ml to 10 ml water, plate as above
 B_2, D_2 : .1 ml to D (asp) 10 ml, 40' in water bath, plate .05 on minifm B.

Plate recombination

10^{-4} dilution of parental suspensions, .025 of each to minifm B, plate

Counts : photometry at 650 m μ of $\frac{1}{100}$ dilution of concentrated parents :

3060	50%	= 240×10^6 /ml
3064	52%	= 320×10^6 /ml

10^{-7} dilution, .05 on B lac i

3060 :	100, 139
3064 :	159, 164

Original conc. suspensions : 24×10^9 /ml; 32×10^9 /ml resp.



1415/3

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1 2 3 4 5 6 7 8 9 10

Plate counts : Numbers were too high for accurate counts.

A₁ : ~ 800

B₁ : same

B₂ : same

C₁ : ~ 500

D₁ : ~ 500

D₂ : ~ 800

E₁ : 250

E₂ : 140.

Plate recomb : 0, 0-

Conclusions : too many recombinants in the unpulsed controls. for any valid conclusions to be drawn. Due to use of D (aspartic) instead of BGA? Azide is however almost ineffective. Try higher concentrations.



19

May 15 1958 -

REF:

1415/4

	1	2	3	4	5	6	7	8	9	10
			AZIDE : DOSE							
1										
2										
3										
4	W 3060	from frj.	2 ml +	7.5 ml	Pen	1 st rot		Spun, conc	3x in Pen	
5										
6	W 3064		1 ml +	7.5 ml	Pen	..			conc 30x "	
7										
8										
9	<u>Mating mixture:</u>		0.2 ml +	0.2 ml,	1' pulse			^{in Pen assay} Dilution		
0	in BGA new (pH 6)		0.1 + 10	&	again	0.1 + 10			and	
1	the same in Pen assay				incubated for another 12'				then	
2	stopped by chilling.									
3	1 ml added from under		<u>chilled tube</u>		Pen		to 0.1 ml of Azide conc:			
4			BGA							
5			0, 1/10000	1/3000	1/1000	1/300	1/60	final conc		1/10 of that indicated
6	final	0	10 ⁻⁵	10						
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9										

May 23/58.

REF:

14/5/5

19

AZ^{10E}, DNP

W 3060 1 ml (fig) + 7.5 ml Pen } (430' Rotation
W 3064 0.5 ml + 7.5 ml Pen

Mating mixture : 1:1 ratio ♂:♀.

incubate 12', chill for 10' → plate 1/10 1/100 1/1000
and add to 0.9 ml of mating mixture: 0.05

A	0.1 ml of DNP M/20	= M/200
B	0.03 "	M/700
C	0.1 " M/200	M/2000
D	0.1 of azide 10%	Azide 1%
E	0.03 " }	3%
F	0.1 " 1%	1%
G	0.1 water	<u>Control</u>

incubate
30'

then drill - Sample and plate .05 of dilutions
1/10, 1/100, 1/1000 of all.

Also: incubate for another 30' after 1/100 dilution
in warmed broth. then plate undil, and 1/10 -

X

mating mixture incubated until the end for a total
of 60' then plated: .05 of 1/10, 1/100, 1/1000

All plating on minifun B,



5/23/1958

REF: 1415/5

19

	1	2	3	4	5	6	7	8	9	10	
			1 st time			2 nd time					
		1/10	1/100	1/1000		1/1	1/10				
1											
2											
3											
4											
5	A	∞	~500	57		~1000	92	} DNP	M/200		
6											
7	B	∞	1000	118		"	137			M/700	
8											
9	C		∞	212		"	251	} Azide	M/2000		
0											
1	D	~1000	19	2		~	81		}	1%	
2	E	∞	103	7		"	67				3%
3											
4	F	∞	~500	54		"	132		1%		
5											
6	G	∞	∞	154		"	166				
7											
8											
9	X	∞	~1000	135							
0											
1	O	∞	112	29							
2											
3											

* big & small, plate record

Conclusions

Matings probably stopped by Az 3%, DNP M/200 or more. Action apparently reversible, but new recruitment may have taken place (although G 1/1000 and G 2 1/10 show equal numbers): no. of prototrophs have considerably increased and it is unlikely that they were all blocked before TL at the 12'.



19

5/28/58

REF:

1415/6

	1	2	3	4	5	6	7	8	9	10
1	AZIDE RESISTANTS.									
2										
3										
4	W 3060 ;		W 3947		(=3060 Az ^R)					
5	W 3064 ;		W 3935		(=3064 Az ^R)					
6										
7										
8										
9										
0										
1										
2										
3										
4										
5										
6										
7										
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2										
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6										
7										
8										
9										
0										

1^{hr} cultures, spun, resusp. 2x conc.

Mating mixtures: 2 ml + 2 ml.
using iced suspensions -

- A ♂_s + ♀_s
- B ♂_s + ♀_r
- C ♂_r + ♀_s
- D ♂_r + ♀_r

Each mixture distributed to 4 tubes, 0.9 ml each.

- Adding:
- tube 1 0.1 ml water
 - 2 Azide 3% 0.1 ml, at 0'
 - 3 " " at 5'
 - 4 " " at 12'

At 40', dilutions 1/5 in chilled water, blending, plating of .05 on D (from B₁). Same for 1/10 dilution from the latter -

5. Plate recombinants



19

May 31st, 1958.

REF:

1415/6

	1	2	3	4	5	6	7	8	9	10
		Plate count		← Azk 0	← Azk 0	← Azk 12				
1										
2		1	2	3	4	5				
3										
4	A	∞	1	45	211	(24)				
5										
6	B	∞	3	73	271	18				
7										
8	C	∞	7	112	359	2				
9										
0	D	∞	16	254	~440	3				
1										
2										
3										
4	A''	(78)	0	0	19	miss				
5										
6	B''	(140)	0	2	-					
7										
8	C''	(140)	0	13	13					
9										
0	D''	(96)	0	8	26					
1										
2										
3										
4										
5										
6										
7										
8										
9										
0										
1										
2										
3										
4										
5										
6										
7										
8										
9										
0										

↑
x factor 10--



	1	2	3	4	5	6	7	8	9	10
1										
2		λ	λR	$\lambda V_1 \lambda_2$	$\lambda V_2 \lambda$					
3	Bact. / Ph									
4										
5	λ^s	S	S	S	S					
6										
7	λ^r	R	R	S	S					
8										
9	λ^h	R	R	S	S					
0	$\lambda(H)^s$	R	S	R	S			isolated once only		
1	$\lambda(H)^h$	R	R	S	S					
2										
3										
4	λ^a λR_1	R	R	S	S					
5										
6	half	R	R	R	R					
7										
8										
9										
0										
1										
2										
3										
4										
5										
6										
7										
8										
9										
0										
1										
2										
3										
4										
5										
6										
7										
8										
9										
0										

isolated once only

= ~~number~~ of infection carried phage

= λR_2



19

5/29/58

REF:

1415/7

	1	2	3	4	5	6	7	8	9	10
--	---	---	---	---	---	---	---	---	---	----

AZIDE RES-: OPTIMAL AZIDE CONC.

O.r.c. cultures of W3060, 3064, 3947, 3935 -
Spun and resuspended in distilled ~~to~~ broth

Mixtures as in exp 1415/6, then 0.9 of mixture per tube,
to which 0.1 of the following conc. of azide in Penassay
were added:

3% , 1% , 0.3% , 0.1% , 0.

Called 1 2 3 4 5 -

45' incubation, then dilution 1/1000, flatys of 0.05
on D (from B.) -

		1	2	3	4	5
A	0	0	7	57	38	
B	0	1	6	18	44	
C	0	1	43	108	~400	
D	1	10	41	118	318	

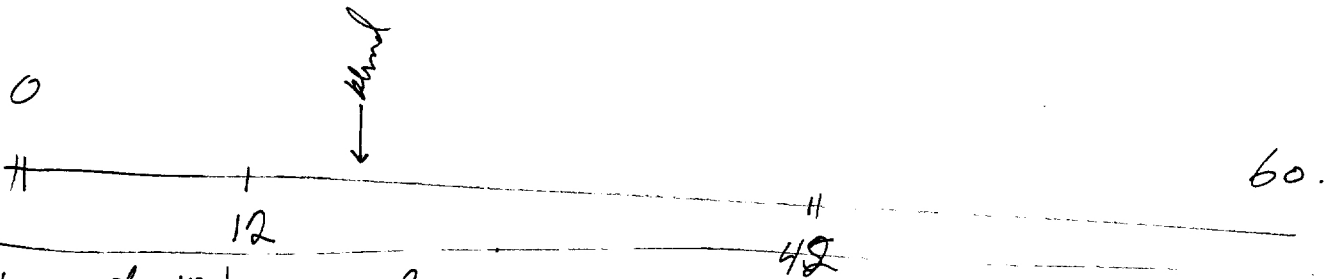
No discrimination.

Pulse — 12 minutes.

- 1. Blend & plate.
- 2. Blend in DNP
- 3. DNP reversibility.

4. Incubate + 30 minutes. Blend + plate.

② add DNP. Blend. Incubate (in DNP) 1 hour and overnight.



1. at 12' d + pl.

2: at 12' add DNP. Incubate 30 minutes in DNP. → 3

3. Blend + plate. 4. Dilute (to remove DNP), Incubate 30 minutes.

5 after 5 minutes, blend. → incubate in DNP {30 minutes} (overnight)

→ dilute incubate 30 minutes

1415

I Effect on maturing

plate after 60 m: = I A - B - C - D. at 10⁵⁵

II Effect on injection: $10^{15} - 11^{15}$ dilute 1:100 in B & A.

↙ plate ~~4~~ k
↘ plate B at 12¹⁵
TV.

5 May.
14/15

3060 - 3064. $\sim \frac{1}{2}$ ml 100x. chilled.

NaN_3 - now have 10 mg/ml. Use at 100x/ml.
Need to dilute 100x at least.

acidine orange.

- ① Do these compounds inhibit mating?
- ② Do they inhibit injection once started? 12 minutes.
- ③ are they reversible.

① mating: add .1 ml cells each parent + .2 ml inhibitor.
pulse 1 minute. dilute ~~1:1000 in~~ a BGA.
warm BGA. Plate 1:10,000 on D smB. at 60 minutes.
955

② Also dilute ① sample 1:100 in BGA. at 12 minutes
add inhibitor. incubate 30 minutes ~~at plate~~ dilute 1:100 blend
& plate. (Anes sample chilled 30 mins.) ~~any some A chilled.~~

③ after 30 minute dilute 1:100, incubate 30 minutes +, then
& plate.

BGA - D)

4 tubes. 1 ml each. 10^{03} PM warmed.
at 10^{15} chilled add 1 ml of inhibitors. - Warm and incubate
+ 2 minutes.