

Jan 5 *Micrococcus cryophilus*

NA. 3 hr old - from old broth culture

- 1 Sch fixation Guinea 10, 12 hydrol underhydrolized
- 2 Sch fix Guinea 15, 18 hydrol slightly underhydr.
- 3 Sch fix Guinea 18, 20 hydrol ok, mixed
- 4 O_3O_4 ~~Sch~~ 3 min, Sch fix ^{Guinea} 18, 20 hydrol
little different than without
 O_3O_4

Jan 6 *M. cryophilus*

old ~~broth~~ culture on NSA 3 hr incubation
Sch fix - Guinea new acid
5 10, 20 min hydrol 60° 10 little over, 20 over
7 0, 3 " " all look alike
8 5, 10 " " "

~~NA~~ seems to give better slides

old broth culture on D-O 3 hr incubation
Sch fix - Guinea new acid

6 10, 20 min hydrol 60° 10 little over, 20 over
7 0, 3 " " all look alike
10 5, 10 " " "

Jan 7

nothing on slide

M. cryophilus

11 on NSA - little growth
hydroly 0, 5 min at 63°

12 on NSA - little growth
hydroly 10, 20 min at 63°

13 on Yeast - little growth

hydroly 0, 5 min 63°

14 on Yeast - little growth

hydroly 10, 20 min 63°

Jan 8

M. cryophilus
NSA- gives better slides

on NSA little growth - at 25° for 15 hrs

11 hydrol 0,5 min at 64°

12 hydrol 10,20 min at 64°

plates then at 30° for 1 hr

15 10, 12

16 14, 16

17 4, 6

18 8, 10

on Yeast little growth - at 25° for 15 hrs

13 hydrol 0,5 min at 64°

14 hydrol 10, 20 min at 64°

plate then at 30° for

Jan/2

E coli on NSA

	uncentrifuged cells incubated	
19	0 hrs hydrol	0, 3 min
20	0 hrs "	5, 10 min
23	$\frac{1}{2}$ hr "	0, 3
24	$\frac{1}{2}$ hr "	5, 10
27	1 hr "	0, 3
28	1 hr "	5, 10
31	1 $\frac{1}{2}$ hr "	0, 3
32	1 $\frac{1}{2}$ hr "	5, 10
35	2 "	0, 3
36	2 "	5, 10
37	2 $\frac{1}{2}$ "	0, 3
38	2 $\frac{1}{2}$ "	5, 10

49

ng/duty

centrifuged cells incubated:

21	0 hrs hydrol	0, 3 min
22	0 hrs "	5, 10 min
25	$\frac{1}{2}$ hr "	0, 3
26	$\frac{1}{2}$ hr "	5, 10
29	1 hr "	0, 3
30	1 hr "	5, 10
33	1 $\frac{1}{2}$ "	0, 3
34	1 $\frac{1}{2}$ "	5, 10
37	2 "	0, 3
38	2 "	5, 10

Stam 4-9

Stam

4-9

(1/10) cells mixed 12 $\frac{30}{10}$ to 2 $\frac{10}{10}$

1:1:10

Plate on NSA. Inc. Stam.

Antifungus took 10 mins. (to concentrate)

4:1

Hyd. 0
 19 } 0 S! cells very dark mitach. (N)
 20 } 2 G, (N).
 5 G, N. Gray background.
 10 Nuclei well stained but fuzzy function?

H 344.

21-23 24. (Cross.) - Hyd. 0
 S, N-negative
 no bridges noted

Hyd 10
 W. but moderately
 light. No bridges
 cells rather small.

out

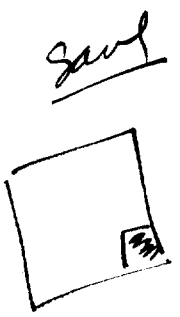
22. cells scattering.
 two types evident.
 No bridges. Almost
 homogeneous stain.

as above.
 somewhat sharper
 but would not
 be adequate for
 bridges.

23 cells sparse
 dense red globules
 in many (P1) cells
 (nuclei?) but background
 also deep blue.

better dispersion
 of cells.

Nuclei clear, cytoplasm
 is not. No
 bridges visible



Need better counterstain
 (or longer in Eosin?)
 0-hydr. from NSA is useless
 exc. for S.

2/3/54

H344

19 hydrolysed 0 and 2 minutes
20 hydrolysed 5 and 10 minutes

clusters of v. large
and intermediate
cells.

~~2/3/54~~

from NSA

H244

(P1 & P2) young cells. Plate +
fix at intervals

2/4/54

- 21 both spread on plate and allowed to dry
hydrolysed 0, 10 minutes
22 incubated 30 minutes, 0, 10 minutes
hydrolysis
23 incubated 45 minutes; 0, 10 minutes
hydrolysis - stain OK. fix?

2/5/54, no cells
too light

- X24 H244 - 0, 10 minutes hydrolysis - from NSA
25 H245 - 0, 10 minutes hydrolysis - from NSA

✓ Next by 11 from (lac) to NSA at 12 noon,
fixed 2:30 PM
✓ Next by 11 well disp. large cells
✓ Insuff. hydrolysis?

WB95 & WB95B

PEN aerated

< A

B

+ Sm

10 g/ml

120'

#97

P $\cdot 10^7$

Z $\cdot 10^5$

120

100

80

60

40

20

0

100

200

300

0

100

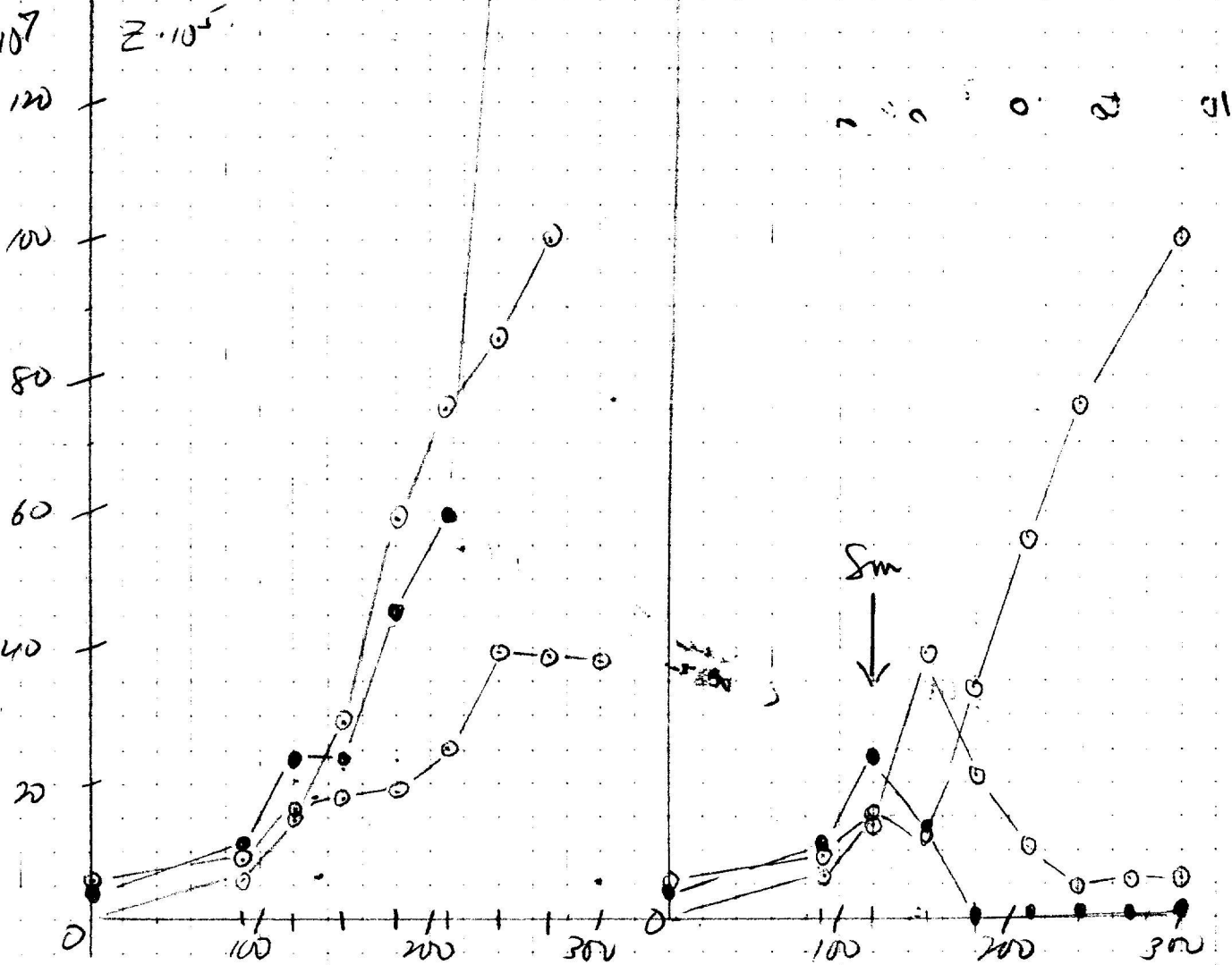
200

300

Sm



10 20 30 40 50



W1895 x W1956

1.0 ml each $\rightarrow$ 10 ml $\left\{ \begin{array}{l} A \\ B \end{array} \right.$
 (12 hr stand)
 PEN culture

PEN 10ml
 PEN + DM 10ml } aerate 37°

#74

W1895 •

W1956 ○

sm^R Lec[±] ○

$\times 10^7$

A

B

60

50

40

30

20

10

0

105
200

100

60

120

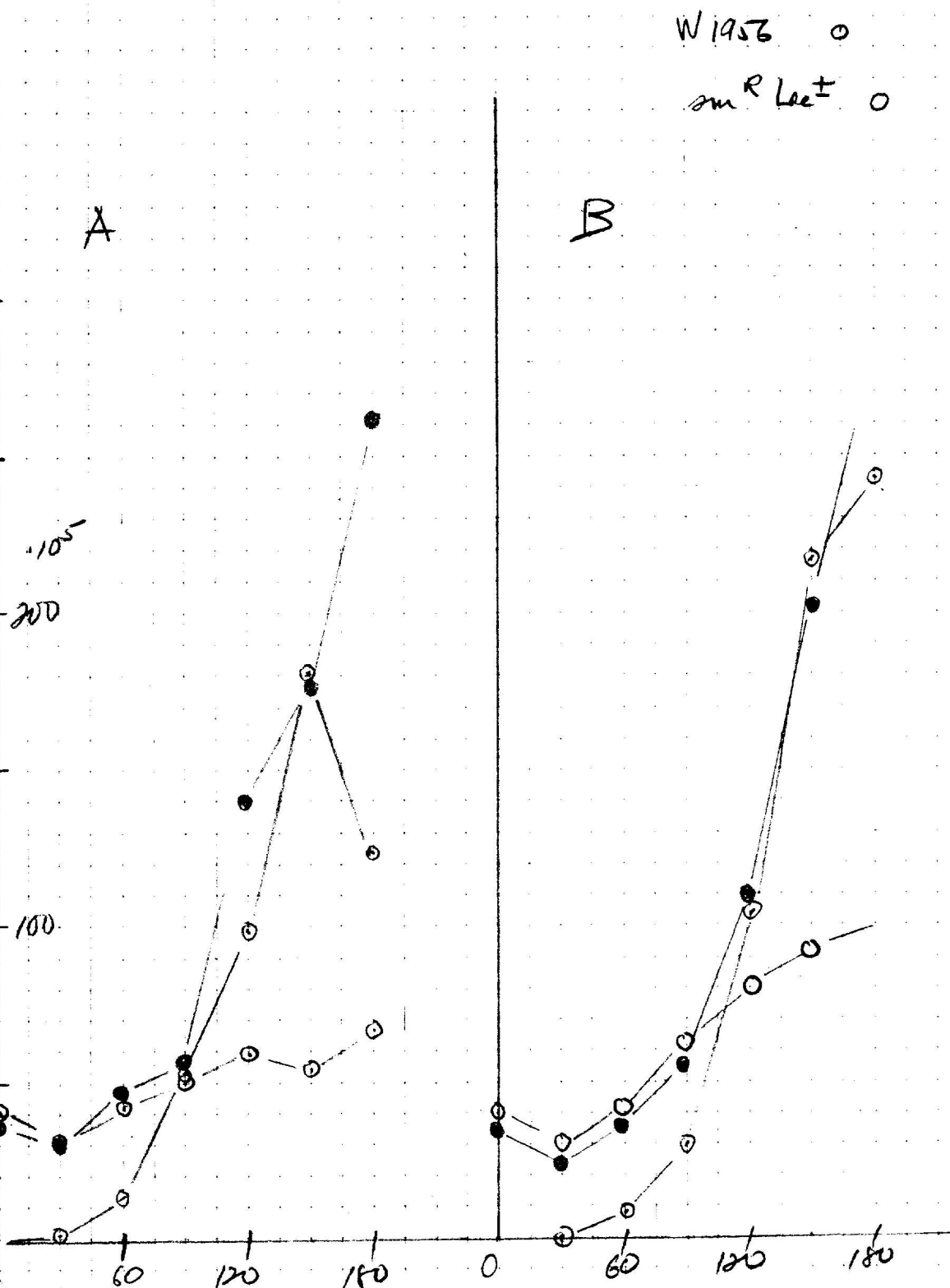
180

0

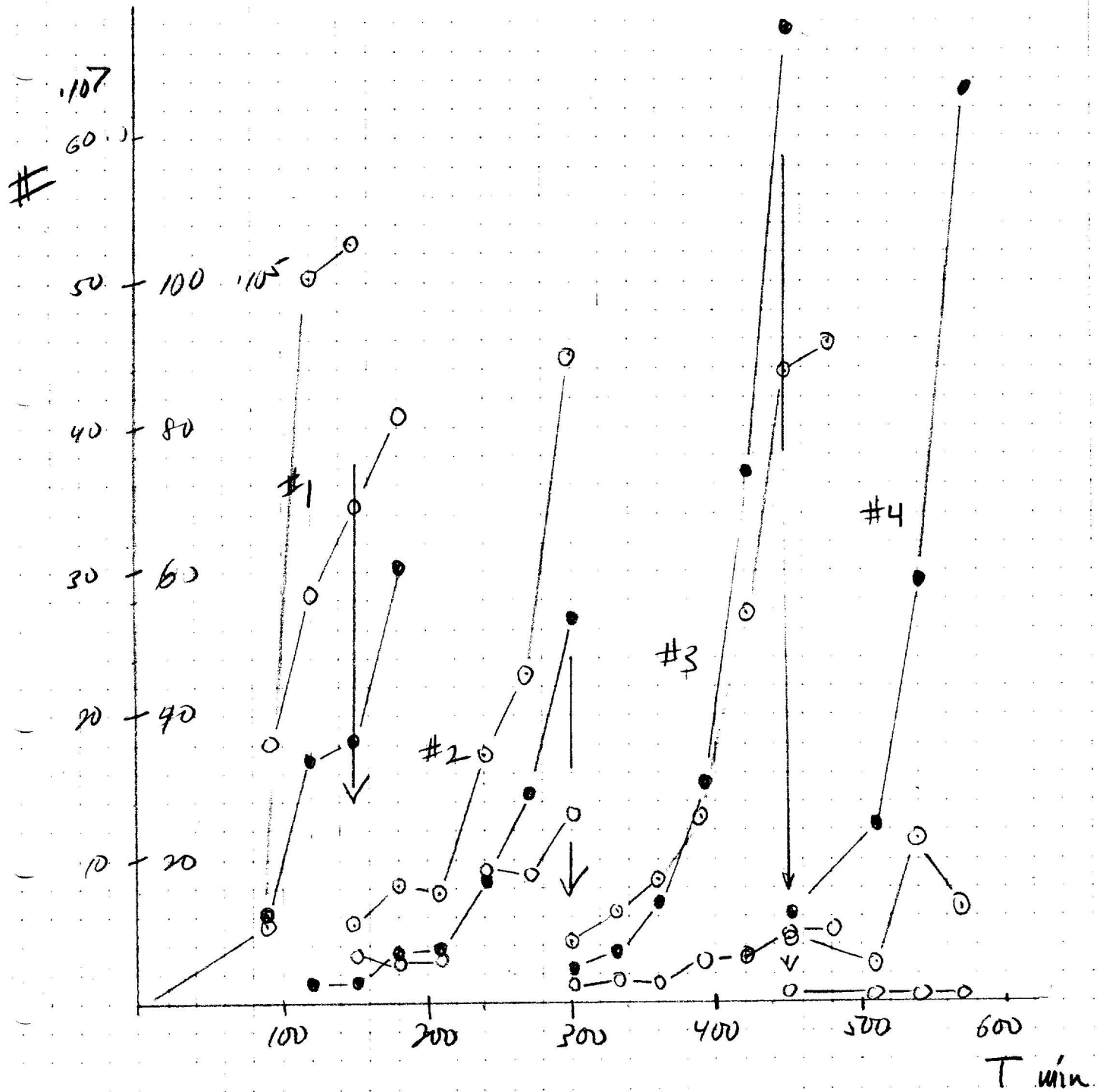
60

120

180



W1895 & W1956 turbidostat PEN+DM + Th 0.2 g/ml #79
 W1956 in initial X₀



#96

turbidostatic growth

PEN ASSAY

Hfr $\approx$ F⁻
1 : 1Hfr < F⁻
1 : 10

N. 106

Z. 105

600

500

400

300

200

100

270

285

300

315

330

270

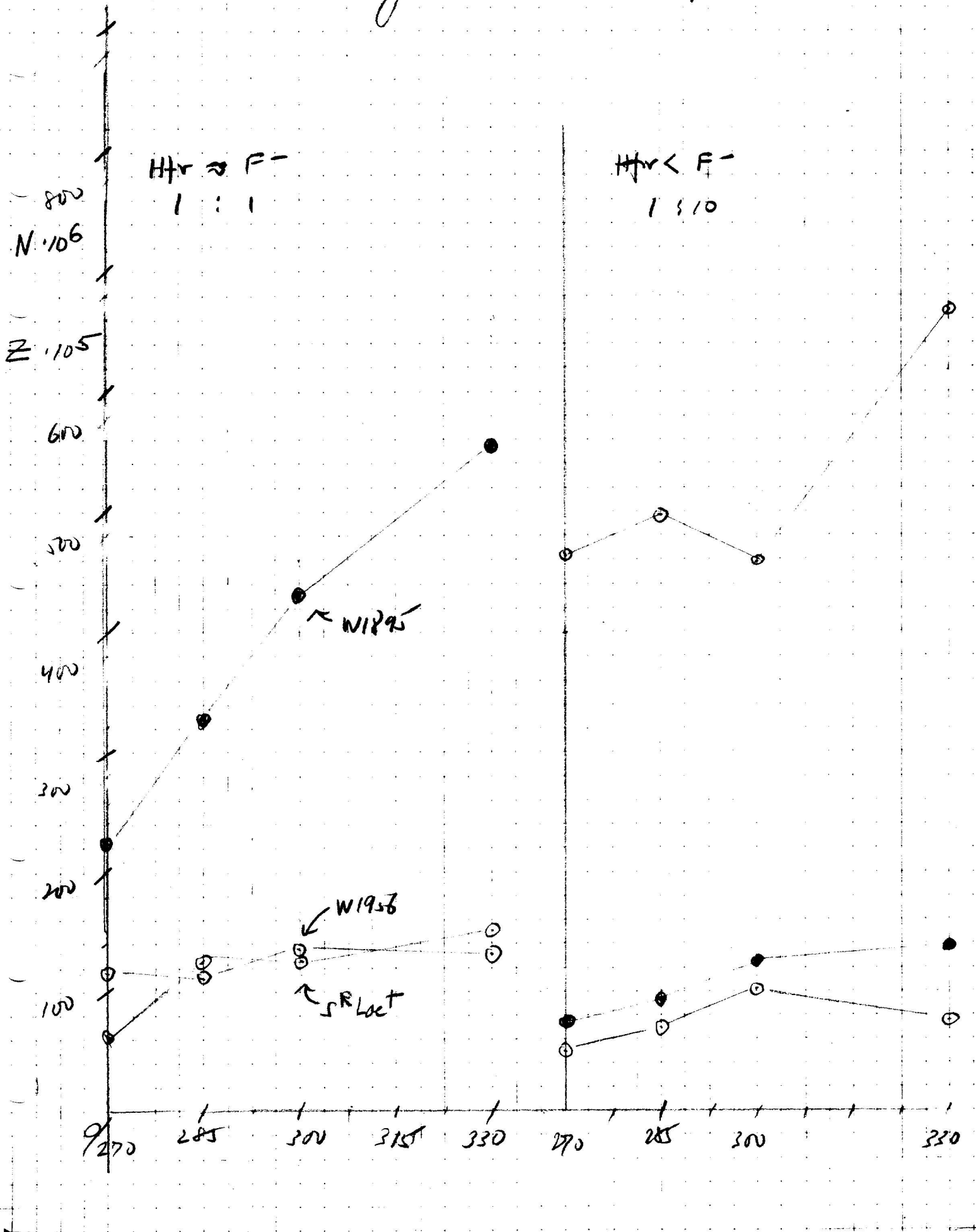
285

300

330

W1895

W1956

SR Lac⁺

HFZF -
1011

800

600

400

200

100

270

285

300

330

N. 106
Z. 106
150

100

50

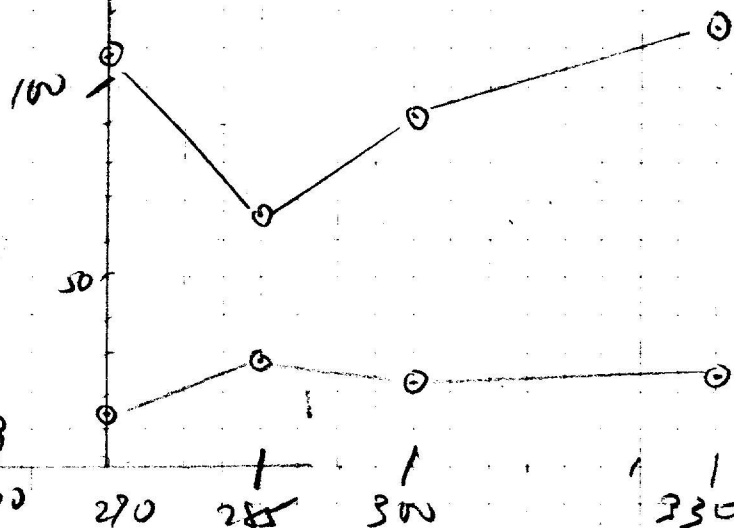
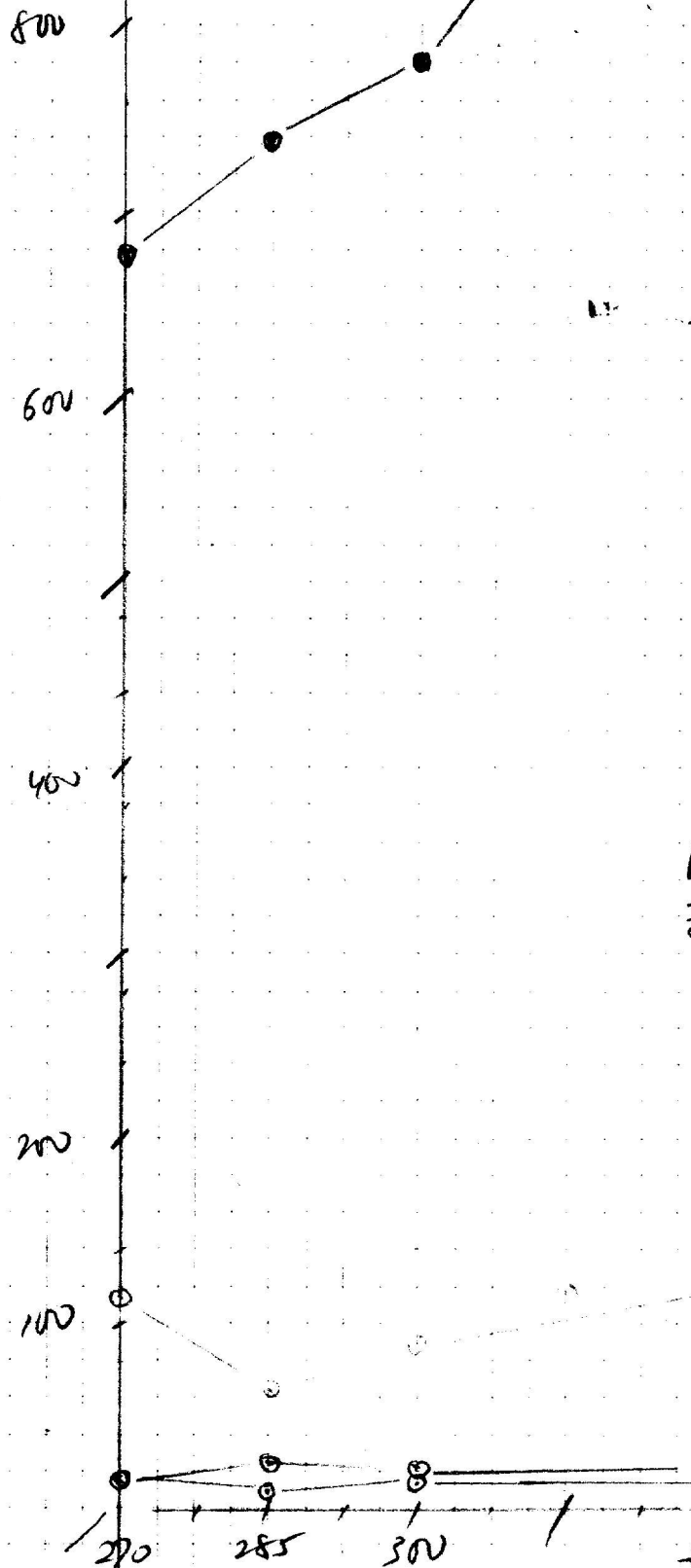
270

285

300

330

as to left - expanded scale
N scale = Z scale



Nuclear Staining Technic

Materials:

Buffer KH_2PO_4 M/15, 0.9078 gm./100 ml. use 4 ml. in about 200 ml.
 Na_2HPO_4 M/15, 0.947 gm./100 ml. use 6 ml. distilled water

Abopon mounting material: Glyco Products Co., a water-soluble resin.

Saturated aqueous mercuric chloride (sublimite): use 2 parts in 1 part absolute ethyl alcohol.

1 ml. conc. HCl plus 9 ml. distilled water makes approx. N/1.

Method:

- J.M.*
1. Fix bacteria (in hanging drop or on sector of agar) in 2% osmium tetroxide for 2 1/2 to 3 minutes. Do not dry cells until after fixation.
 2. Spread on coverslips or make impression smear. Dry 10 seconds.
 3. Robinson uses additional fixation for 1 1/2 min. in HgCl_2 , followed by washing in several changes of 95% alcohol and water.
 4. Place slide in cold N/HCl for 60 seconds.
 5. Hot HCl (about 60°C.) for 10 minutes. *12-14*
 6. Cold HCl for about 20 seconds.
 7. Wash several times in buffer, pH 7. *5 drops + 2 drops 1% MIB/10 ml buffer.*
 8. Stain for 30 minutes in Giemsa, diluted 1-100 or 1-20 with buffer. (Ordinary commercial Giemsa usually requires stronger soln.)
 9. Wash briefly in buffer.
 10. Mount in water or Abopon (diluted 2 parts in 1 part water).

Chabaud's Fixative

EtOH 80%	60 ml
Phenol	15 gm
Formalin	5 ml
Acetic Acid	2 ml

"N/1 HCl" = 1 vol conc HCl + 9 vol H₂O.

For further examination

19: generally too dense

A/B suggests that dipping flattens cells out which may be advantageous for some purposes, but may distort shape and engender S? Relation to G is not clear.

Due to confusion, comparisons of $\pm 0s$ are not worthwhile.

However Gurnea seems OK.

222 Series complete except 22.

Need time ^(Hcl) since an alcohol effect: angrain's w/ us.

Also. G from unbird from Sues:

study!
 Ey A. - 2 (4m.) G not prom. (G'l?)
 C5A - 2 - N v. char. No cyt. Grapus cytoplasm!
 (blueish + reddish)
 (N?)
 G'l

	axial ratio	size
A1-1 N, S ^{occ}	3-5	$14/2 = 1$ $12/3 = 2$
-2 N! (some cells still have blue cytoplasm)		1 or 2
-3 N.		

A2

C. 4A - 1 Flattened, dest. No S. N faint metach.

4B (5B) N met dehydr. (overstained). Sl. shrinkage but not dest.

5A (4B) Cells sparse. but resemble

5B. (5A) Sl. destruction but less than 1st.

no S.

suggests NASse > NSA to minimize dehydr.

C1. ~~NO~~ ^{NO} ~~variable~~ ^{variable}

23-1

- 2 G distinct over most of slide // dark purple cyt v. faint N?

- 3 Typ. N

permanent S
N ✓ but only in answer accumulation (high salt?)
beautiful terminal S at

III/25.

22

1 —

2 ac G klibs. No G. dark, hom. cytoplasm

3 = 20A1 app.

23

2 V. Prom ^{blue} G. No N? Granular cyt. purple.

3 typ N. Note residual (?)

11/10/53.

15. Compare fixatives. But acid possibly weak.

G not seen after osmic., V. prominent in 15A2-3!

16. ^{B, K} y. E coli B, K12. Fix Schaudinn S, G prominent in both -1, -2.

A. W1177 x W1895 from Penassay, comparing NA $\pm$ salt $\pm$ suc.
Note S only in NSA.

Make number pups - ask best can

W1637.

~~W2333~~

✓ other org...

Fri W2333 HCl series.
Bz effects.
Crosses.

Bz effect

Alcohol effect

data on H₂O x F - persistence
of H₂O in cell?

Repeat

A2.-1

-2

- 3

-3.

A3-1

B1 - 1

B2-1

3/26

Cm 0, 3, 5, 10 wt. series

cells are more eccentric than (W2049) and may have more obvious vesiculate nucleus

3/31

€ (W2049)

10 now OK. Quite distinct from Cm.

cells longer, narrower rather than peripheral chromatin.

3/29/24 slides

$\frac{c2}{0} - 4$ ✓ ✓ ~~weak~~ N (negative + intact) dep. aff.

10: G⁺ but N weak

11-3 0 as above. Some S, weak - staining

1 - 2_H 1

↓ better.

2 - 2_H 2 faint N; some S. dep. cytopl.

10 very (washed out) - that dirty
some signs of transverse walls.

slides 84

3/31/54
5 W2404
6. W2438
W2049

acid or
A too much!

4 stilly red
hazog. purple

v. l. much detail:

5 7m HCl.
Rounded cells
hangerous purple
blobs! usually prob. of 8!

13m sl. washed out
but ~~is~~ differentiation
of nuclei.

6. as 4. Some weak blobs?
(weak acid ~~pro~~ by desolysis $\rightarrow$ blobs?)

13. as 4. Δ may
be negative.

4/14/54 slides (8 D)

Hg/ brevis

B/cu

0 in Hcl.

N.v. dense blue cytopl., N. neg.
dupes? no detail
pale no detail
post detail, clear apt.

3-4

2
5
10

①

2

cells red!!

DATE:

4/14/54

REF:

from NSA

BSI

I didn't think there was anything on the first 1-27, so I did another pair

1 - 0, 2 minutes hydrolysis at 59°

10

Giesea stavi

2 S, 10 minutes hydrolysis - Giesea

20

BSIC -

3 0, 2 minutes hydrolysis - Giesea

4 S, 10 minutes hydrolysis - Giesea

30

BSIC - rev

5 0, 2 minute hydrolysis - Giesea

40

6 S, 10 minute hydrolysis Giesea

50