UNCLASSIFIED

AD NUMBER

AD469383

NEW LIMITATION CHANGE

TO

Approved for public release, distribution unlimited

FROM

Distribution authorized to U.S. Gov't. agencies and their contractors; Administrative and Operational Use; Jul 1965. Other requests shall be referred to the Army Biological Laboratories, Fort Detrick, MD 20701.

AUTHORITY

BDRL, per Dept Army ltr, 28 Sep 1971

THIS PAGE IS UNCLASSIFIED

SECURITY MARKING

The classified or limited status of this report applies to each page, unlass otherwise marked. Separate page printents MUST be marked accordingly.

THIS DOCUMENT CONTAINS INFORMATION AFFECTING THE NATIONAL DEFENSE OF THE UNITED STATES WITHIN THE MEANING OF THE ESPIONAGE LAWS, TITLE 18, U.S.C., SECTIONS 793 AND 794. THE TRANSMISSION OR THE REVELATION OF ITS CONTENTS IN ANY MANNER TO AN UNAUTHORIZED PERSON IS PROHIBITED BY LAW.

NOTICE: When government or other drawings, specifications or other data are used for any purpose other than in connection with a definitely related government procurement operation, the U. S. Government thereby incurs no responsibility, nor any obligation whatsoever; and the fact that the Government may have formulated, furnished, or in any way supplied the said drawings, specifications, or other data is not to be regarded by implication C. otherwise as in any manner licensing the holder or any other person or corporation, or conveying any rights or permission to manufacture, use or sell any patented invention that may in any way be related thereto.

AD

TECHNICAL MANUSCRIPT 235

69

0

60

ො

C

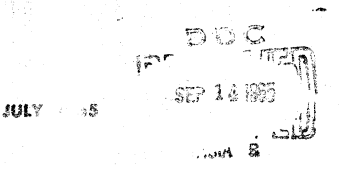
T

 $\sum_{i=1}^{n}$

C

FLUORESCENT CELL-COUNTING ASSAM OF YELLOW FEVER VIRUS

Nicholas Mahon



Best Available Copy

BOLGOICAL LABORATORIES FORT DETRICK

U.S. ARMY BIOLOGICAL LABORATORIES Fort Detrick, Frederick, Maryland

TECHNICAL MANUSCRIPT 235

FLUORESCENT CELL-COUNTING ASSAY OF YELLOW FEVER VIRUS

Nicholas Hahon

Aerobiology Division DIRECTORATE OF BIOLOGICAL RESEARCH

1

Project 10522301A084

July 1965

Best Available Copy

ACKNOWLEDGMENT

The author gratefully acknowledges the technical assistance of Kenneth Cooke during certain phases of the work.

ABSTRACT

A quantitative assay of infective yellow fever virus particles was developed that is based on the enumeration of cells containing fluorescent viral antigen 24 hours after infection of coverslip McCoy cell cultures. The rapidity and efficiency of virus adsorption onto cell monolayers was markedly enhanced by centrifugation of virus inoculum. By this procedure, 95% of virus was adsorbed within 15 minutes and a proportionality was demonstrated between the number of fluorescent cells and the volume of inoculum.

The incubation period of 24 hours was established from observations on the sequential development of viral antigen within cells and fluorescent cell counts. The relationship between virus concentration and cell-infecting units of virus was linear; the distribution of fluorescent cells in cell monolayers was random. The sensitivity of the fluorescent cell-counting assay was comparable to that of the method of intracerebral inoculation of mice.

A quantitative linear relationship was demonstrated between the quantity of virus neutralized and dilutions of antiserum.

I. INTRODUCTION

Since the original demonstration by Coons et al.¹ that mumps virus antigen may be detected microscopically by staining with fluorescent antibody, the procedure has been successfully employed to visualize viral antigen in cell cultures infected by a wide variety of viruses. Recently, a quantitative assay of infective virus particles was Coveloped that is based on the enumeration of fluorescent cells containing viral antigen. This technique has been standardized for only a limited number of viruses that represent a few major groups, e.g., myxoviruses,² adenoviruses,³ herpesviruses,⁴ psittacosis agents,⁵ and poxviruses.^{5,*} In assays of these viruses, the fluorescent cellcounting technique was shown to be sensitive, precise, reproducible, and rapid (results usually obtained within 24 hours). The usefulness of the technique has been enlarged further with the development of a fluorescent cell-counting neutralization test to detect and to measure quantitatively serum-neutralizing antibody.⁷ The feasibility of extending the fluorescent cell-counting technique to the assay of arboviruses was investigated in view of its marked advantages and potential in virus studies.

This report describes the development and standardization of the fluorescent cell-counting technique for the quantitative assay of yellow fever virus.

II. METHODS

A. VIRUS

A plasma suspension of the Asibi strain of yellow fever virus was used throughout this study. Infective plasma was obtained from a rhesus monkey that had been inoculated intraperitoneally with 6 x 10^4 mouse intracerebral LD₅₀ (MICLD₅₀) units of virus and bled 3 days later. The plasma was divided into 1-ml portions, sealed in glass vials, and choosed at -60 C. The plasma suspension of virus had a titer of 10^{8-1} MIGLE₅₀.

. wroublished results.

B. CELL LINE AND CULTIVATION

The McCoy cell line was used for the assay of virus. The origin of the cell line, the procedures, and the medium used for cell cultivation have been reported previously.⁵ Preliminary tests employing high multiplicities of virus to cells indicated that approximately 99% of the cells were susceptible to infection by yellow fever virus.

C. ROTOR CHAMBER INSERTS

Centrifugation was employed routinely to adsorb virus inoculum onto coverslip cell cultures. To withstand the high centrifugal force required to sediment yellow fever virus, a rotor chamber insert (Figure 1) was designed to fit into the arm of a swinging-bucket type SW 25.1 rotor.* Each rotor chamber insert accommodated a 15-mm coverslip cell culture and as much as 1 ml of inoculum. The chamber insert was sealed with a threaded lid. Two slots in the lid facilitated the removal of the chamber insert with forceps from the arm of the rotor after centrifugation. Each arm of the rotor held four chamber inserts (Figure 2). Rotor chamber inserts were made of Delrin** a plastic that retained its structural integrity after repeated autoclaving for 30-minute intervals at 121 C. Both nylon and Lexan*** polycarbonate resin were also suitable materials.

D. VIRUS ASSAY

Virus dilutions were prepared in maintenance medium consisting of mixture 199 and 5% calf serum and introduced in 0.2-ml volumes directly onto coverslip cell cultures held in sterile rotor chamber inserts. Determinations were generally made in triplicate. Routinely, virus adsorption was carried out in a model L Preparative Ultracentrifuge* at 15,000 rpm (19,642 to 29,432 x g, depending on the distance of the chamber insert in the arm of the rotor from the axis of rotation) for 15 minutes at 25 C. Because the plastic rotor chamber inserts were slightly toxic for cell cultures after prolonged contact, coverslip cell cultures

* Beckman Instruments, Inc., Spinco Division, Palo Alto, California. ** Confimm Glass Co., Wilmington, Delaware. *** Concral Electric Co., Schenectady, N.Y.

Est Station Cont 1

Figure 1. Rotor Chamber Insert for Holding Coverslip Cell Cultures.



Figure 2. Alignment of Rotor Chamber Inserts in Arm of SW 25.1 Swinging-Bucket Rotor.

7

were removed and inserted into sterile flat-bottomed glass vials $(18 \times 100 \text{ mm})$. One ml of maintenance medium was added to each vial and the coverslip cell cultures were then incubated at 35 C for 24 hours. Coverslip cell cultures were rinsed twice with cold phosphate buffered saline (PBS), fixed with cold (-60 C) acetone, and either prepared immediately for immunofluorescent staining and cell counting or stored at -60 C. Fixed cell cultures stored at -60 C for as long as 7 weeks showed no appreciable decrease in fluorescence on staining.

E. ANTISERUM

For the preparation of yellow fever virus antiserum, rhesus monkeys were inoculated intraperitoneally with 1 ml of yellow fever vaccine.* Ten days later, the animals were challenged by injection of 1 ml of virulent yellow fever virus. A second challenge dose was administered 2 weeks later; surviving monkeys were bled 2 weeks after the last inoculation. Yellow fever virus antiserum was conjugated with fluorescein isothiocyanate by the method of Riggs et al.⁸ The conjugate was adsorbed twice with mouse liver and chick embryo powders.

F. IMMUNOFLUORESCENT PROCEDURES

The direct fluorescent antibody technique was used to obtain immunofluorescence of infected cells. The staining procedure has been described elsewhere.⁵ Coverslip cell cultures were examined for fluorescent cells with an American Optical microscope equipped with a Fluorolume illuminator, model 645, Corning no. 5840 and Schott BG-13 exciter filters, and E. K. no. 24 barrier filter. At 430 X magnification with the optical system employed, 1064 microscopic fields were contained in the area of a 15-mm coverslip. Fifty microscopic fields were examined for fluorescent cells in each coverslip cell culture. To calculate the number of cell-infecting units (CIU) of virus per ml, the average number of fluorescent cells per field was multiplied by the number of fields per coverslip, the reciprocal of the dilution of virus inoculum, and a volume factor (for conversion to ml).

* The National Drug Company, Philadelphia, Pa.

III. RESULTS

A. STANDARDIZATION OF THE FLUORESCENT CELL-COUNTING TECHNIQUE

The efficiency of centrifugation for the adsorption of yellow fever virus onto coverslip cell monolayers was determined at different centrifugal speeds. Virus adsorption was also carried out with inoculated cell cultures maintained in a stationary position. The results in Table 1 show that comparable assay values were obtained with centrifugal speeds of 10,000 to 22,500 rpm. Assay values were significantly lower at centrifugal speeds below 10,000 rpm as well as at stationary incubation.

To determine whether the position of the rotor chamber inserts in fully loaded rotor arms affected the efficiency of virus adsorption onto cell monolayers, virus inoculum was centrifuged at 15,000 rpm for 15 minutes. The centrifugal force from the proximal to the distal chamber insert ranged from 19,642 to 29,430 x g. Results in Table 2 attest to the efficiency of virus adsorption onto coverslip cell cultures in all rotor chamber inserts at the defined conditions of centrifugation. The same experiment was carried out at 10,000 rpm for 15 minutes; the centrifugal force from the proximal to the distal chamber insert ranged from 8730 to 13,080 x g. That adsorption of virus was inefficient at this condition was reflected by a progressive decrease in assay values corresponding to the proximity of the chamber insert to the axis of rotation.

The rate of virus adsorption onto coverslip cell cultures was determined during centrifugation at 15,000 rpm. Virus inoculum was introduced onto coverslip cell cultures held in rotor chamber inserts and centrifuged for designated intervals of time. To determine the quantity of unadsorbed virus, residual inoculum was introduced onto new coverslip cell cultures and centrifuged at 20,000 rpm for 15 minutes. The results indicate that 50% of the virus inoculum was adsorbed in 5 minutes and approximately 95% in 10 minutes. The amount of virus adsorbed during centrifugation periods of 15, 20, and 30 minutes was similar to that of the 10-minute value. In stationarey cultures, 30% of virus inoculum was adsorbed at 35 C for 2 hours and 22% of virus inoculum was adsorbed at 25 C for 4 hours. In view of the results obtained from these experiments, centrifugation at 15,000 rpm for 15 minutes was selected as the condition for routine adsorption of yellow fever virus onto cell monolayers.

Speed,ª/ rpm	<u>Centrifugation</u> Centrifugal Force, gravity	CIU ^{b/} per ml
2,000	523	5.3 x 10 ⁵
3,000	1,177	3.1×10^{7}
5,000	3,270	5.8 x 10^7
10,000	13,080	1.1×10^{8}
15,000	29,430	1.0×10^{8}
20,000	52,320	1.1 x 10 ⁸
22,500	65,400	1.0×10^{8}
	Stationary Incubation	
Temperature,	Hours	CIU per ml
C		
35	2	4.1×10^{7}
25	4	3.1×10^7

TABLE 1. EFFICIENCY OF CENTRIFUGATION VS. STATIONARY INCUBATION ON THE ADSORPTION OF YELLOW FEVER VIRUS ONTO MCCOY CELL MONOLAYERS

a. Centrifuged for 15 minutes at 25 C.

b. Cell-infecting units of virus.

TABLE 2. EFFICIENCY OF CENTRIFUGATION ON YELLOW FEVER VIRUS ADSORPTION ONTO MCCOY CELL MONOLAYERS IN ROTOR CHAMBER INSERTS ALIGNED AT DIFFERENT DISTANCES FROM THE AXIS OF ROTATION

Distance from Axis of Rotation, cm	Centrifugal Force, <u>a</u> / gravity	10 ⁸ CIUE/ per ml
7.8	19,642	1.4
9.1	22,889	1.4
10.4	26,160	1.5
11.7	29,430	1.3

a. Contrifugation at 15,000 rpm for 15 minutes, 25 C.

b. Gell-infecting units of virus.

The efficiency of centrifugation for promoting virus adsorption from different volumes of inoculum is shown in Table 3. The results revealed a proportionality between the number of fluorescent cells and volume of inoculum. No proportionality was noted when similar volumes of inoculum were introduced onto coverslip cell cultures that were then maintained in a stationary position at 35 C for 2 hours. The rate of cell-virus contact appeared to be independent of the volume of inoculum when centrifugal force was employed.

Volume, ml	Fluorescent Cells Per	r 50 Fields	10 ⁸ CIUE/ per ml
0.1	78	· ·	1.4
0.2	154	• •	1.5
0.5	360		1.5
1.0	711		1.5

TABLE 3. PROPORTIONALITY BETWEEN VOLUME OF INOCULUM AND CELL-INFECTING UNITS OF YELLOW FEVER VIRUS

a. Cell-infecting units of virus.

The incubation period, defined here as the interval between virus inoculation and the appearance of recognizable quantities of viral antigen in cell cultures suitable for enumeration of infected cells. was established from sequential observations on the development of viral antigen and fluorescent cell counts. Coverslip cell cultures were inoculated in the prescribed manner and fixed after designated intervals of incubation at 35 C. Within 16 hours after inoculation, a few cells containing faint cytoplasmic fluorescent foci were noted. Fluorescence was more diffuse and granular in appearance at 20 and 22 hours after infection. At 24 and 26 hours, fluorescent cells were easily discerned. Fluorescence was intense and filled the cytoplasm and its clongated extensions (Fig. 3). That a second cycle of infection had occurred was suggested by the presence of varied degrees of fluorescence in cell cultures examined at 30, 44, and 48 hours after inoculation. Some elongated cells contained minute fluorescent foci; others exhibited diffuse fluorescence throughout the cytoplasm. In cells that had rounded, fluorescence was intense. In general, the development and appearance of fluorescence in infected McCoy cells was similar to that described in infected hamster kidney cells.⁹ At 16 to 20 hours, counts of fluorescent cells were one-fifth to one-tenth the uniform counts found at 22 to 26 hours; after this period (30 to 48 hours) they were 3 to 4 times higher. Based on these observations and findings, the optimal period for incubation of cell cultures inoculated with yellow fever virus was established as 24 hours.



Figure 3. Fluorescent Yellow Fever Virus Antigen in McCoy Cells 24 Hours after Infection. Magnification: X 215.

B. QUANTITATIVE EVALUATION OF THE ASSAY

A linear relationship was demonstrated between twofold dilutions of virus over a range of 1.8 log units and the number of cell-infecting units of virus (Fig. 4). These data suggest that each fluorescent cell was the consequence of infection by a single infective virus particle.

In a single experiment, 10 determinations were made to estimate the precision of the fluorescent cell-counting assay for yellow fever virus. Coverslip cell cultures were infected with a standard quantity of virus inoculum and incubated in the prescribed manner. The number of cell-infecting units of virus per ml of inoculum ranged from 1.1×10^8 to 1.6×10^8 with a mean of 1.2×10^8 . The standard deviation of 0.18 compared favorably with those obtained in similar studies with the psittacosis agent⁶ and variola virus.*

The mode of distribution of fluorescent cells on a coverslip cell monolayer was determined by examing 200 random microscopic fields. The frequencies of fields containing fluorescent cells correspond closely to the theoretical frequencies (Fig. 5). The X^2 test of possibless of fit of the experimental data to the theoretical Poisson distribution gave a probability of approximately 0.75 at d.f. = 8. Fluorescent cells were randomly distributed in infected cell monolayers.

The sensitivity of the fluorescent cell-counting procedure was compared with that of intracerebral inoculation of mice. The results in Table 4 indicate that there was no significant difference in sensitivity between the two assay procedures. Assays showed less variation, however, when performed by the fluorescent cell-counting technique than by the method of intracerebral inoculation of mice.

* Eshon, unpublished results.

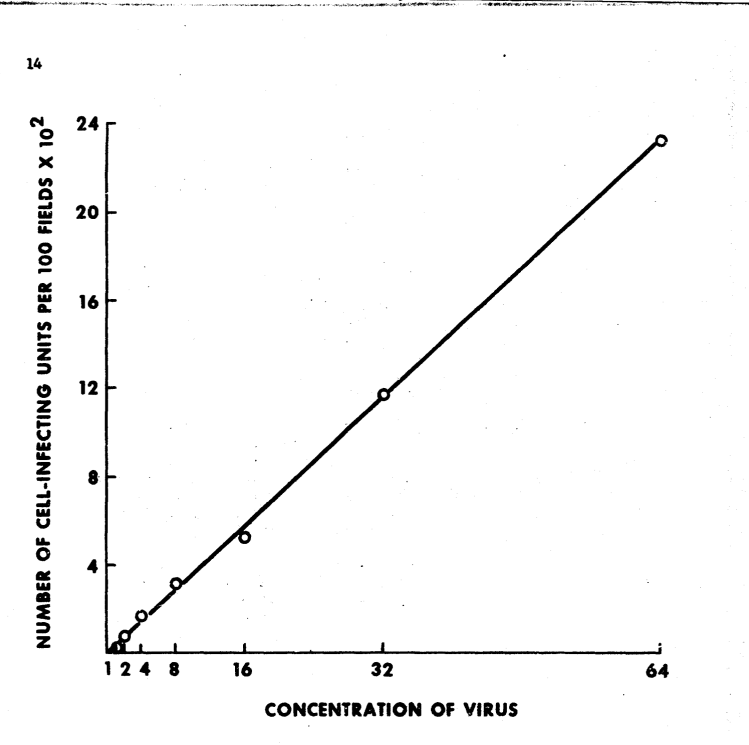


Figure 4. Linear Function Between Concentration of Yellow Fever Virus and the Number of Cell-Infecting Units.

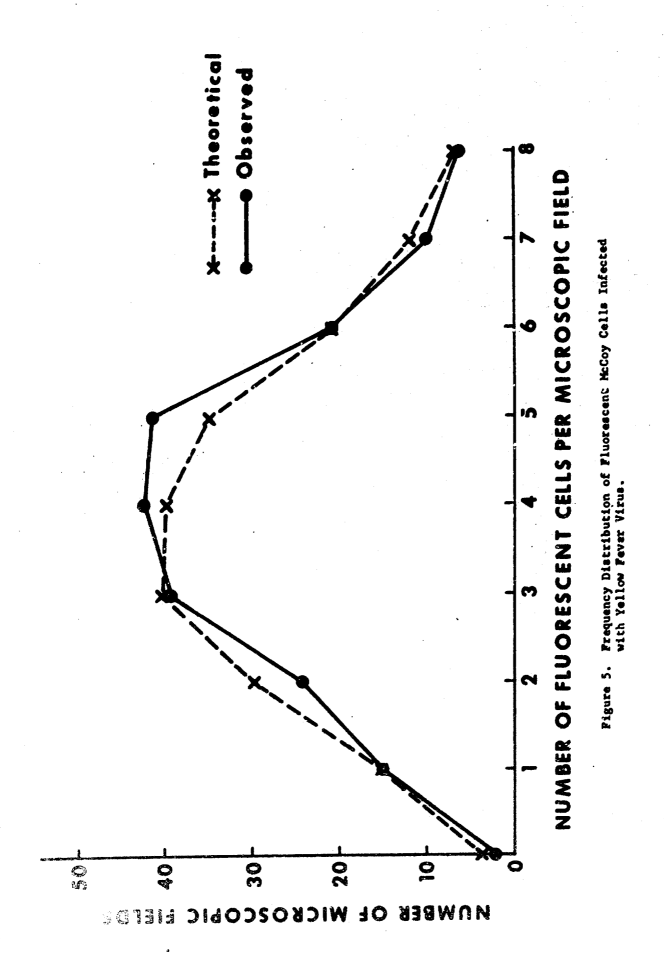


TABLE 4. COMPARISON OF TWO PROCEDURES FOR THE ASSAY OF YELLOW FEVER VIRUS

Assay Procedure	Mean	SD	SE, mean
Fluorescent cell-counting	8.14/	±0.07	±0.08
Intracerebral inoculation of mice	8.1 <u>b</u> /	±0.67	±0.22

a. Reciprocal of cell-infecting units of virus (log₁₀) per ml based on 10 titrations, determined in 24 hours.

 Reciprocal of LD₃₀ (log₁₀) mouse intracerebral units of virus per ml based on 12 titrations, 14-day observation period.

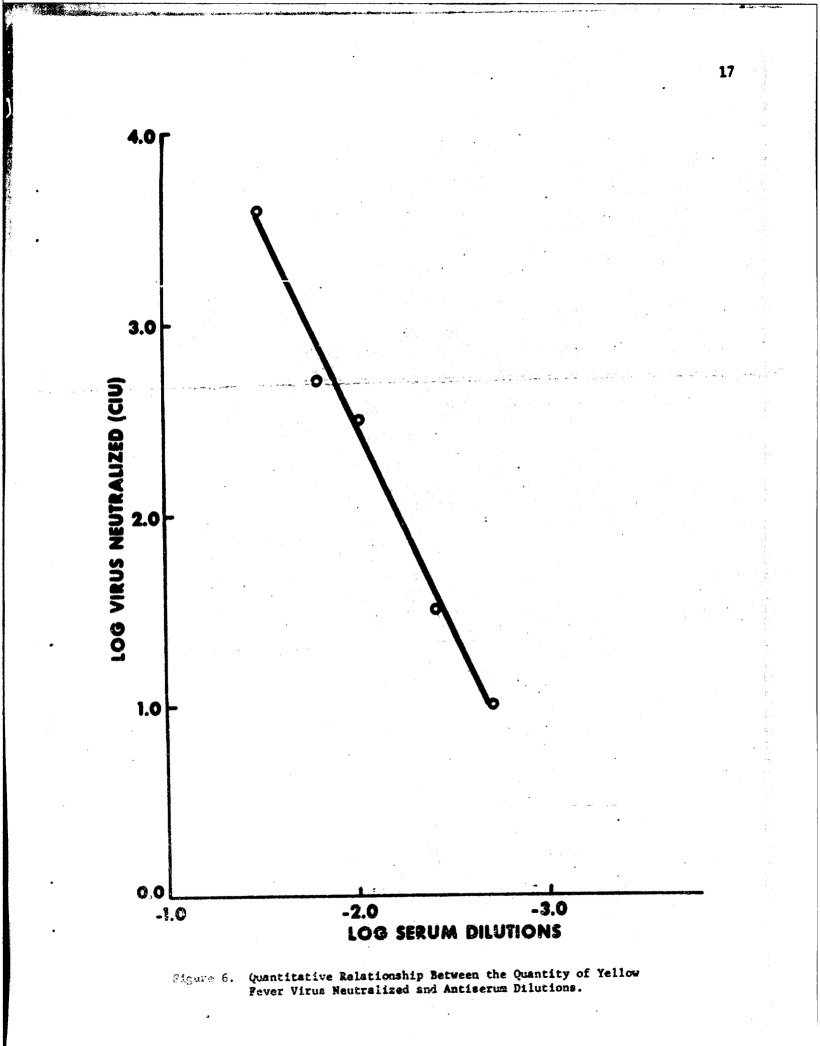
C. QUANTITATIVE VIRUS-SERUM RELATIONSHIP

Box titrations were carried out by mixing equal volumes of different concentrations of virus suspension and serial twofold dilutions of virus antiserum. Virus-serum mixtures were incubated at 35 C for 1 hour and introduced onto coverslip cell cultures in accord with the standard procedure. A linear function was obtained between the quantity of virus neutralized and antiserum dilutions (Fig. 6). For each 0.5 log unit (3.2-fold) increase of virus concentration, the quantity of antiserum required for neutralization increased approximately 0.3 log unit (2-fold).

IV. DISCUSSION

The use of centrifugal force to promote virus adsorption onto coverslip cell cultures is an important factor that has contributed directly to the successful development of the fluorescent cell-counting technique for the quantitative assay of infectious yellow fever virus particles. The fact that centrifugation of virus inoculum increases the rapidity and efficiency of virus adsorption was previously demonstrated in the development of the fluorescent cell-counting assay for the psittacosis agent⁵ and variola virus.* Because of the large size of these infectious particles, a centrifugal force of only 500 x g was required to bring approximately 99% of them in contact with the cell monolayer. With

* Hahon, unpublished results.



eastern equine encephaloxyelitis virus, an infectious agent that approaches the size of yellow fever virus, Gey et al.¹⁰ employed centrifugation at more than 100,000 x g for 1 ..our to expose harvested HeLa cell cultures to a suspension of the virus. Evidence that centrifugation enhanced virus adsorption was obtained from the early destructive effect of the virus on cell cultures. Noncentrifuged control cell cultures, exposed to the same amount of virus suspension, showed only the beginning of a cytopathic effect in an equivalent period of time.

The sedimentation and efficient adsorption of yellow fever virus onto coverslip cell monolayers with the aid of high centrifugal speeds was feasible by the use of plastic rotor chamber inserts that were designed to hold coverslips. Approximately 95% of virus was adsorbed from inoculum within 15 minutes at 25 C with centrifugal force that ranged from 19,000 to 29,000 x g, whereas only 30% of virus was adsorbed from inoculated cell cultures held in a stationary position at 35 C for 2 hours. The latter result approximated the findings of Litwin¹¹ on the adsorption of yellow fever virus in stationary cell cultures held at 35 C for 1 hour. Although only 10 to 20% of the cells could be initially infected with a high inoculum¹¹ centrifugation of high virus concentrations of inoculum resulted in the infection of approximately 99% of the cells in monolayers. In addition, with the aid of centrifugation, a proportionality was demonstrated between the volume of inoculum and the number of fluorescent cells. The relative efficiency of virus adsorption as a function of the volume of inoculum is relevant to the detection of virus particles from dilute suspensions. Centrifugation of virus inoculum also increased the sensitivity of the fluorescent cell-counting assay so that it was comparable to that of the method of intracerebral inoculation of mice. The advantages of the fluorescent cell-counting technique for the assay of yellow fever virus over that of the latter procedure is in its greater precision, rapidity, specificity, and ease of making replicate determinations.

The concomitant increase in intensity of fluorescent viral antigen in infected cells and the uniformity of fluorescent cell counts at 22 to 26 hours after inoculation served to establish this interval as the length of the incubation period. The initial growth cycle of yellow fever virus lasts 14 to 15 hours after infection, and virus is liberated for at least 10 hours thereafter.¹¹ Ideally, the incubation period should be terminated before newly synthesized virus particles are released to prevent secondary infection of cells. Because the accumulation of viral antigen in infected cells was insufficient to permit the accurate enumeration of fluorescent cells at 15 hours after inoculation, the incubation period was extended past the time of virus release. This extension period, from the time of virus release (15 hours) to the end of the incubation period (24 hours), did not affect the accuracy of fluorescent cell counts because it was too short to permit the visualization of fluorescent viral antigen in secondarily infected cells. The demonstration that a quantitative relationship exists between the quantity of yellow fever virus neutralized and dilutions of antiserum provides a basis for the development of a fluorescent cell-counting neutralization test to measure serum-neutralizing antibodies. For research and diagnostic purposes, the advantages of the test would be similar to that offered by the fluorescent cell-counting assay of virus. The assay of yellow fever virus by the procedure described is applicable to the quantitative assay of other arboviruses and those viral agents of small dimensions.

LITERATURE CITED

- Coons, A.H.; Snyder, J.C.; Cheever, F.S.; Murray, E.S. 1950. Localization of antigen in tissue cells: IV. Antigens of rickettsiae and mumps virus. J. Exp. Med. 91:31-38.
- Wheelock, E.F.; Tamm, I. 1961. Enumeration of cell-infecting particles of Newcastle disease virus by the fluorescent antibody technique. J. Exp. Med. 113:301-316.
- 3. Philipson, L. 1961. Adenovirus assay by the fluorescent cellcounting procedure. Virology 15:263-268.
- 4. Goodheart, C.R.; Jaross, L.B. 1963. Human cytomegalovirus. Assay by counting infected cells. Virology 19:532-535.
- 5. Hahon, N.; Nakamura, R.M. 1964. Quantitative assay of psittacosis virus by the fluorescent cell-counting technique. Virology 23:203-208.
- 6. Carter, G.B. 1965. The rapid detection, titration, and differentiation of variola and vaccinia viruses by a fluorescent antibody-coverslip coll monolayer system. Virology 25:659-662.
- 7. Hahon, N.; Cooke, K.O. 1965. Fluorescent cell-counting neutralization test for psittacosis. J. Bacteriol. 89:1465-1471.
- Riggs, J.L.; Seiwald, R.J; Burckhalter, J.H.; Downs, C.M.; Metcalf, T.G. 1958. Isothiocyanate compounds as fluorescent labeling agents for immune serum. Amer. J. Pathol. 34:1081-1097.
- 9. McGavran, M.H.; White, J.D. 1964. Electron microscopic and immunofluorescent observations on monkey liver and tissue culture cells infected with the Asibi strain of yellow fever virus. Amer. J. Pathol. 45:501-517.
- Gey, G.O.; Bang, F.B.; Gey, M.K. 1954. Responses of a variety of normal and malignant cells to continuous cultivation, and some practical applications of these responses to problems in the biology of disease. Ann. N. Y. Acad. Sci. 58:57(439.
- 11. Litwin, J. 1964. Growth of 17D yellow fever virus and factors influencing its transmission within cell cultures in vitro. Acta Pathol. Microbiol. Scand. 61:605-618.

Best Available Copy

DOCU (Security claunification of title - b-dy of abateur	MENT CONTROL DAT		a the overall report is the called.
1 ORIGINATING ACTIVITY (Corporate muthor)			ORT SECTION TY CLASSING A LIGH
U.S. Army Biological Laboratori Fort Detrick, Frederick, Maryla		U1 21 GRC	nclassified
3 REPORT TITLE			
FLUORESCENT CELL-COUNTING ASSAY	OF YELLOW FEVER V	IRUS	
4 DESCRIPTIVE NOTES (Type of repart and inclusiv	e datez)		
5 AUTHOR(5) (Last name, first name, initial)			
Hahon, Nicholas			
S. REPORT DATE	7. TOTAL NO.		70 NO. OF REFS
July 1965		BEES	
LE CONTRACT DE GRANT NO.			
6. раслест NO. 10522301A084	Те	chnical Mar	nuscript 235
¢.	Sb. OTHER RE this report)	PORT NO(S) (A	ny other numbers that may be assigned
d.			
Qualified requestors may obtain Foreign announcement and dissemine Release or announcement to the p	nation of this pu ublic is not auth 12. SPONSORIN U.S. Arm	blication b orized. G MILITARY AC y Biologica	DDC is not authorized
Foreign announcement and dissemi Release or announcement to the p	nation of this pu ublic is not auth 12. SPONSORIN U.S. Arm	blication b orized. G MILITARY AC y Biologica	by DDC is not authorized.
Foreign announcement and dissemi Release or announcement to the p 11. SUPPLEMENTARY NOTES 13 ABSTRACT A quantitative assay of infe	nation of this public is not authors of this public is not authors of the second secon	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par	by DDC is not authorized. TIVITY al Laboratories erick, Maryland, 21701 ticles was
Foreign announcement and dissemi <u>Release or announcement to the p</u> 11. SUPPLEMENTARY NOTES 13 ABSTRACT A quantitative assay of infe developed that is based on the e	nation of this public is not authors of this public is not authors of the second secon	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls containi	by DDC is not authorized TIVITY 11 Laboratories erick, Maryland, 21701 ticles was ing fluorescent
Foreign announcement and dissemin <u>Release or announcement to the p</u> 11 SUPPLEMENTARY NOTES 13 ABSTRACT A quantitative assay of infe developed that is based on the e viral antigen 24 hours after inf	nation of this public is not authors in the second	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls containi ip McCoy ce	by DDC is not authorized TIVITY 11 Laboratories erick, Maryland, 21701 ticles was ing fluorescent ell cultures.
Foreign announcement and dissemi <u>Release or announcement to the p</u> 11. SUPPLEMENTARY NOTES 13 ABSTRACT A quantitative assay of infe developed that is based on the e	nation of this public is not authors in authors in a second state of the second state	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls contains ip McCoy ce nto cell mc	by DDC is not authorized. TIVITY 11 Laboratories erick, Maryland, 21701 ticles was ing fluorescent ell cultures. poolayers was
Foreign announcement and dissemine Release or announcement to the p in supplementary notes a quantitative assay of infe developed that is based on the e viral antigen 24 hours after inf The rapidity and efficiency of v markedly enhanced by centrifugat 95% of virus was adsorbed within	nation of this public is not authors of this public is not authors of the second secon	dication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls contains ip McCoy ce nto cell mo ulum. By t proportion	by DDC is not authorized TIVITY 11 Laboratories erick, Maryland, 21701 ticles was ing fluorescent ell cultures. pholayers was this procedure, hality was
Foreign announcement and dissemine Release or announcement to the p in supplementary notes A quantitative assay of infe developed that is based on the e viral antigen 24 hours after inf The rapidity and efficiency of v markedly enhanced by centrifugat	nation of this public is not authors of this public is not authors of the second secon	dication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls contains ip McCoy ce nto cell mo ulum. By t proportion	by DDC is not authorized TIVITY 11 Laboratories erick, Maryland, 21701 ticles was ing fluorescent ell cultures. pholayers was this procedure, hality was
Foreign announcement and dissemine Release or announcement to the p in supplementary notes a ABSTRACT A quantitative assay of infe developed that is based on the e viral antigen 24 hours after inf The rapidity and efficiency of v markedly enhanced by centrifugat 95% of virus was adsorbed within demonstrated between the number	nation of this public is not authom ublic is not authom is sponsorm U.S. Arm Fort Det: ctive yellow fever numeration of celle ection of coversi- irus adsorption on ion of virus inocu- 15 minutes and a of fluorescent celle hours was established antigen within ce oncentration and of fluorescent cells fluorescent cell-	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls contains ip McCoy ce nto cell mo ulum. By t proportion ls and the shed from c ls and flucture cell-infect in cell mc counting as	by DDC is not authorized. TIVITY 11 Laboratories erick, Maryland, 21701 eticles was ing fluorescent e11 cultures. pholayers was this procedure, hality was to volume of beservations on the morescent cell counts. eting units of virus pholayers was
Foreign announcement and dissemin <u>Release or announcement to the p</u> IS SUPPLEMENTARY NOTES A quantitative assay of infe developed that is based on the e viral antigen 24 hours after inf The rapidity and efficiency of v markedly enhanced by centrifugat 95% of virus was adsorbed within demonstrated between the number inoculum. The incubation period of 24 sequential development of viral The relationship between virus c was linear; the distribution of random. The sensitivity of the	nation of this pui ublic is not auth 12 SPONSORIN U.S. Arm Fort Det ctive yellow fever numeration of cel ection of coversi- irus adsorption or ion of virus inoce 15 minutes and a of fluorescent ce hours was establin antigen within ce oncentration and of fluorescent cells fluorescent cell- rebral inoculation	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls contains ip McCoy ce ato cell mo alum. By t proportion ls and the shed from c ls and fluc cell-infect in cell mo counting as n of mice.	by DDC is not authorized. TIVITY 11 Laboratories erick, Maryland, 21701 eticles was ing fluorescent e11 cultures. pholayers was this procedure, mality was this volume of beservations on the morescent cell counts. eting units of virus pholayers was say was comparable
Foreign announcement and dissemin <u>Release or announcement to the p</u> 11. SUPPLEMENTARY NOTES 13 ABSTRACT A quantitative assay of infe developed that is based on the e viral antigen 24 hours after inf The rapidity and efficiency of v markedly enhanced by centrifugat 95% of virus was adsorbed within demonstrated between the number inoculum. The incubation period of 24 sequential development of viral The relationship between virus c was linear; the distribution of random. The sensitivity of the to that of the method of intrace A quantitative linear relati	nation of this pui ublic is not auth 12 SPONSORIN U.S. Arm Fort Det ctive yellow fever numeration of cel ection of coversi- irus adsorption or ion of virus inoce 15 minutes and a of fluorescent ce hours was establin antigen within ce oncentration and of fluorescent cells fluorescent cell- rebral inoculation	blication b orized. G MILITARY AC y Biologica rick, Frede r virus par ls contains ip McCoy ce ato cell mo alum. By t proportion ls and the shed from c ls and fluc cell-infect in cell mo counting as n of mice.	by DDC is not authorized. TIVITY 11 Laboratories erick, Maryland, 21701 eticles was ing fluorescent e11 cultures. pholayers was this procedure, mality was this volume of beservations on the morescent cell counts. eting units of virus pholayers was say was comparable

2 . • • • • •

.