

ういたのし

MICROCOPY RESOLUTION TEST CHART NATIONAL BUREAU OF STANDARDS-1963-A

	_	
A	D	

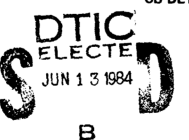
AD-A142 069

CRDC-TR-84013

DYE FLUORESCENCE ANALYSIS FROM BACTERIAL METABOLISM

by A. Peter Snyder David B. Greenberg Theresa T. Wang

RESEARCH DIVISION CB DETECTION & ALARMS DIVISION



April 1984

06

13

012

約4

US Army Armament, Munitions & Chemical Command Aberdeen Proving Ground, Maryland 21010

DISTRIBUTION STATEMENT A Approved for public released Distribution Unlimited

FILE COPY 3110

Disclaimer

The findings in this report are not to be construed as an official Department of the Army position unless so designated by other authorizing documents.

Disposition

For classified documents, follow the procedures in DOD 5200.1-R, Chapter IX, or DOD 5220.22-M, "Industrial Security Manual," paragraph 19. For unclassified documents, destroy by any method which precludes reconstruction of the document.

Distribution Statement

Approved for public release; distribution unlimited.

REPORT DOCUMENTATION PAGE	READ INSTRUCTIONS BEFORE COMPLETING FORM
REPORT NUMBER 2. GOVT ACCESSION NO	
CRDC-TR-84013 AD_4/42	467
TITLE (and Subtitie)	5. TYPE OF REPORT & PERIOD COVERED
	Technical Report
DYE . ORESCENCE ANALYSIS FROM	December 1982-March 19
BACTEI AL METABOLISM	5. PERFORMING ORG. REPORT NUMBER
AUTHOR(a)	8. CONTRACT OR GRANT NUMBER(.)
A. Peter Snyder	
David B. Greenberg	
Theresa T. Wang	
PERFORMING ORGANIZATION NAME AND ADDRESS	10. PROGRAM ELEMENT, PROJECT, TASK AREA & WORK UNIT NUMBERS
Commander	AREA & WORK UNIT NUMBERS
Chemical Research and Development Center ATTN: DRSMC-CLB-R (A)	1L161101A91A
Aberdeen Proving Ground, Maryland 21010	ILIUIIOIAJIA
CONTROLLING OFFICE NAME AND ADDRESS	12. REPORT DATE
Commander	April 1984
Chemical Research and Development Center	13. NUMBER OF PAGES
ATTN: DRSMC-CLJ-IR (A) Aberdeen Proving Ground, Maryland 21010	20
4. MONITORING AGENCY NAME & ADDRESS(II dillerent from Controlling Office)	
	UNCLASSIFIED
	15a. DECLASSIFICATION/DOWNGRADING SCHEDULE
	NA NA
6. DISTRIBUTION STATEMENT (of this Report) Approved for public release; distribution	4
	unlimited.
Approved for public release; distribution	unlimited.
Approved for public release; distribution	unlimited.
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the abetract entered in Block 20, 11 different fo	unlimited.
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the ebetrect entered in Block 20, 11 different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse eide if necessary and identify by block numbe	unlimited.
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the ebstrect entered in Block 20, 11 different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse side if necessary and identify by block number Induced fluorescence Fluores	unlimited.
Approved for public release; distribution D. DISTRIBUTION STATEMENT (of the obstreet entered in Block 20, 11 different for D. SUPPLEMENTARY NOTES D. KEY WORDS (Continue on reverse side if necessary and identify by block number Induced fluorescence Diacetylfluorescein Lipase	unlimited.
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the obstreet entered in Block 20, 11 different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse elds if necessary and identify by block number Induced fluorescence Fluores Diacetylfluorescein Lipase	unlimited.
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the obstrect entered in Block 20, if different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse elds if necessary and identify by block number Induced fluorescence Fluores Diacetylfluorescein Lipase Bacterial metabolism	unlimited. man Report) */ scent product
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the obstreet entered in Block 20, 11 different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse side if necessary and identify by block number Induced fluorescence Fluores Diacetylfluorescein Lipase Bacterial metabolism 1. AMETRACT (Continue on reverse olds H necessary and identify by block number A method based upon induced fluor the rapid detection and characterization of	unlimited. Tom Report) "" scent product prescence is proposed for of viable microorganisms
Approved for public release; distribution DISTRIBUTION STATEMENT (of the obstreet entered in Block 20, 11 different is USUPPLEMENTARY NOTES USUPPLEMENTARY NOTES NEY WORDS (Continue on reverse side if necessary and identify by block number Induced fluorescence Fluores Diacetylfluorescein Lipase Bacterial metabolism AMETRACT (Continue on reverse side if necessary and identify by block number A method based upon induced fluor the rapid detection and characterization of In this technique, a nonfluoresceing dye is	unlimited. mom Report) "" scent product "" rescence is proposed for of viable microorganisms s metabolized intracel-
Approved for public release; distribution Approved for public release; distribution Distribution STATEMENT (of the obstract entered in Block 20, 11 different in Supplementary notes Supplementary notes Automation on reverse side if necessary and identify by block number Induced fluorescence Fluores Diacetylfluorescein Lipase Bacterial metabolism Additional metabolism A method based upon induced fluo the rapid detection and characterization of In this technique, a nonfluorescing dye in lularly by an organism through an enzyme-	unlimited. mom Report) "" scent product "" rescence is proposed for of viable microorganisms s metabolized intracel- specific reaction. This
Approved for public release; distribution Approved for public release; distribution Distribution STATEMENT (of the observed entered in Block 20, 11 different for Supplementary notes Supplementary notes Automation on reverse elde if necessary and identify by block number Induced fluorescence Fluores Diacetylfluorescein Lipase Bacterial metabolism Amethod based upon induced fluo the rapid detection and characterization of In this technique, a nonfluorescing dye is lularly by an organism through an enzyme- produces a fluorescent product which when	unlimited. wom Report) Scent product prescence is proposed for of viable microorganisms s metabolized intracel- specific reaction. This desorbed can be detected
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the obstract entered in Block 20, 11 different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse elds if necessary and identify by block number Induced fluorescence Fluores: Diacetylfluorescein Lipase Bacterial metabolism 4. AMSTRACT (Continue on reverse elds H necessary and identify by block number A method based upon induced fluorescence f	unlimited. wom Report) Scent product prescence is proposed for of viable microorganisms s metabolized intracel- specific reaction. This desorbed can be detected
Approved for public release; distribution 7. DISTRIBUTION STATEMENT (of the obstreet entered in Block 20, 11 different for 8. SUPPLEMENTARY NOTES 9. KEY WORDS (Continue on reverse elds if necessary and identify by block number Induced fluorescence Fluores: Diacetylfluorescein Lipase Bacterial metabolism 1. ABSTRACT (Continue on reverse elds H measures) and identify by block number A method based upon induced fluo the rapid detection and characterization of In this technique, a nonfluorescing dye is lularly by an organism through an enzyme- produces a fluorescent product which when The technique has been applied successful	unlimited. wom Report) Scent product prescence is proposed for of viable microorganisms s metabolized intracel- specific reaction. This desorbed can be detected

inter en en en en en en

Ċ,

•

UNCLASSIFIED SECURITY CLASSIFICATION OF THIS PAGE (When Date Entered)

20. ABSTRACT (Contd)

Bacillus globigii through the enzyme lipase and the nonfluorescing dye diacetylfluorescein. By relating the effect of bacterial and substrate activity to the initial rate of fluorescence generation, standard curves have been obtained from which both age and bacterial concentration can be estimated.

2

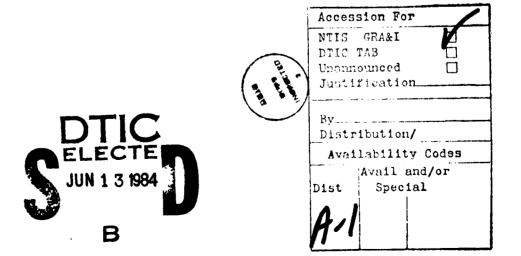
SECURITY CLASSIFICATION OF THIS PAGE(When Data Entered)

PREFACE

The work described in this report was authorized under Project 1L161101A91A, CB Detection and Alarms. This work was started in December 1982 and completed in March 1983.

The use of trade names in this report does not constitute an official endorsement or approval of the use of such commercial hardware or software. This report may not be cited for purposes of advertisement.

Reproduction of this document in whole or in part is prohibited except with permission of the Commander, Chemical Research and Development Center, ATTN: DRSMC-CLJ-IR (A), Aberdeen Proving Ground, Maryland 21010. However, the Defense Technical Information Center and the National Technical Information Service are authorized to reproduce the document for United States Government purposes.



Blank

والمتعالم والمراجع والمراجع والمراجع والمتعاد والمتعاد والمتعال والمعالية والمعالية والمعالية والمعالية والمعا

343434343

.

CONTENTS

Page

 $\sim \sim \sim$

1.		7
2.	EXPERIMENTAL METHODS AND MATERIALS	8
2.1 2.2	Bacterial Preparation and Handling	8 8
3.	RESULTS AND DISCUSSION	8
4.	CONCLUSIONS	12
	LITERATURE CITED	15
	DISTRIBUTION LIST	17

-...

Blank

۳.,

٠.

(6553363) (2002026) (2002027) (2002026) (2002027) (2002027)

and the second statement of the second statement of the second statement of the second statement of the second

and the state of the

DYE FLUORESCENCE ANALYSIS FROM BACTERIAL METABOLISM

1. INTRODUCTION

A methodology for rapid detection and identification of microorganisms has long been of concern to the medical, pharmaceutical, and food processing fields. Because of this sustained interest, the area has remained active and has seen significant advances over the classical time-consuming protocols of standard plate counting, membrane filtration, or the multiple-tube fermentation procedures. However, even today these techniques can require as much as 1 to 3 days; hence, continued research is still indicated before a rapid, generalized microbial characterization scheme is to be realized.

Various earlier developments have focused upon techniques in immunofluorescence, 1^{-6} enzyme amplification, 7,8 light detection and ranging (LIDAR), 9 mass spectrometry, 10 bacteriophage lysis, 11-13computer assisted probabilistic methods, 14-18 the recently developing field of gel ferrography 19, and the double fluorescent DNA staining method coupled to flow cytometry. 20 Each of these techniques is limited by disadvantages such as laborious and time-consuming laboratory preparation and sample handling, long observation times, and nonspecificity with respect to bacterial characterization and subsequent identification.

The present investigation, which we believe to offer the possibilities of speed, sensitivity, and specie identification to the process of microorganism detection is embraced in the following equation:

nonfluorescent +	enzyme	supplied	fluorescent	+ metabolites (1)
substrate	(in bacteria)	cofactors	product	+ metabolites (1)

This method combines several attractive experimental features. These include

- (a) exploitation of the particular enzymatic/metabolic machinery of the cell by an added substrate in which a predetermined reaction takes place,
- (b) the product fluorescence which can be used to indicate a positive cellular response, and
- (c) rapid data collection and analysis under 30 minutes.

Equation 1 presents various modes of biochemical strategy in order to uniquely characterize a given bacterium or small group of bacteria. The technique pinpoints the identity of a particular enzyme that is indigenous to the bacteria themselves through several

^{*}J. K. Marquis, A. P. Russell, and V. C. Westcott. Boston University Medical Center. Personal communication, 1983.

experimental avenues. In one instance, for the same bacterial enzyme a series of chemically similar substrates (e.g., esters) can be utilized to generate unique reaction rates and, thus, associated product fluorescence. Another scheme involves sequentially using a series of substrates in order to trigger the various potential enzymatic reactions that can occur within the bacterium. The present experimental format exploits the former procedure.

The first-order nature of the experimental fluorescence-producing chemistry in this work suggests the Michaelis-Menten data analysis approach as most attractive. Precedence for applying Michaelis-Menten kinetics to enzymatic processes in vivo (i.e., reactions occurring in the intact, living cell) has been previously documented in the literature, 21,22 although the theory, strictly speaking, was initially developed for enzymatic processes occurring in vitro.

2. EXPERIMENTAL METHODS AND MATERIALS

2.1 <u>Bacterial Preparation and Handling</u>.

Bacillus subtilis var niger (Bacillus globigii, BG), the test organism, was grown on a nutrient agar slant for 24 hours at 37° C and stored at 4°C. One loopful of the inoculum culture from the slant was spread evenly over the agar surface of the nutrient agar plates with a sterile glass rod. After 16 to 18 hours incubation at 37° C, the growth layer was scraped from the agar by adding 10 ml of sterile 0.1 M Tris, 0.85% NaCl, pH 8.0, in each plate. The cell suspension was diluted to 75 ml with the buffer and was stored at 4° C. Viable cell counts of the bacterial samples in the first set of experiments (vide infra) were made by plating serial dilutions using sterile distilled water as diluting blank , tryptose agar as plating agar, 37° C as incubation temperature, and 36 hours as incubation time.

Nutrient agar and tryptose agar were purchased from DIFCO Laboratories, Detroit, Michigan.

2.2 <u>Bacterial Assays</u>.

The specific reaction examined was the hydrolysis of the substrate diacetylfluorescein $(5.0 \times 10^{-5} \text{M})$ by microbial lipase. Normally, methylcellosolve²³⁻²⁵ and acetone²⁷ are the solvents used to dissolve diacetylfluorescein with subsequent buffer dilution; however, we found early in our work that acetone provides approximately twice the activity as methylcellosolve. A typical assay solution consisted of 2.0 ml of the dye solution at a prepared dye concentration²³⁻²⁶ and 0.1 ml of the bacterial suspension added at the beginning of the assay. A Farrand fluorimeter, Model MK-2, was used to excite diacetyl-fluorescein at 469 nm and to monitor the emission at 519 nm with excitation and emission slits of 1.0 and 2.5 nm, respectively.

3. RESULTS AND DISCUSSION

Two sets of experiments were conducted over a 1-month period using the same suspension in order to observe the effect of bacterial age on the significant parameters. In one study in which the substrate concentration was held constant, the bacterial concentration was varied and the initial reaction rate was determined via the tangent to the initial portion of the fluorescence generation curve. Figures 1 and 2 reflect these results. The plot presented in Figure 1 shows a marked contrast in the differentiation of the hydrolysis rates of bacterial diacetylfluorescein above 5.0×10^4 cells/ml, which correlates well with the data of Pinteno and Findl²⁷ as shown in Figure 3. Their work made use of the enzymatic hydrolysis of fluorescein-di- β -galactopyranoside with Escherichia coli neotype. The normalized comparison between the two studies suggest similar kinetics.

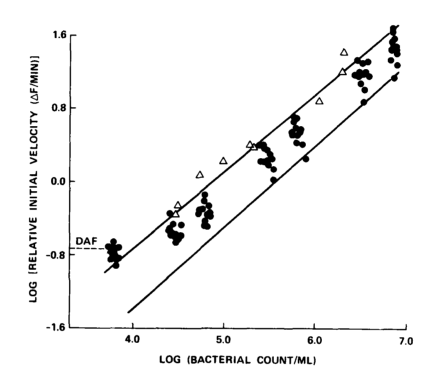
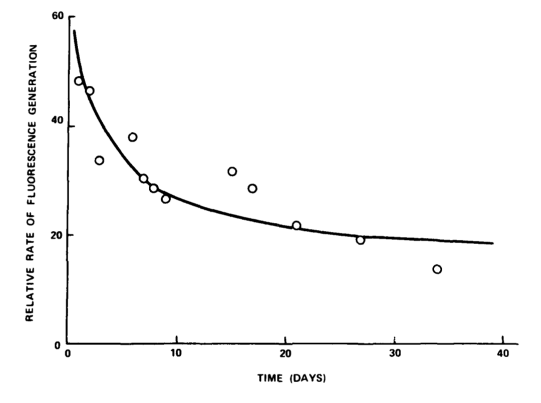
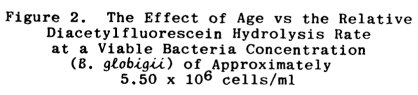
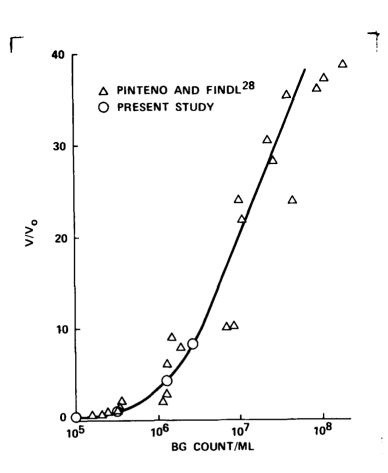


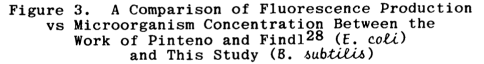
Figure 1. Log Plot of the Concentration of an Aging B. globigii (BG) Suspension vs the Relative Diacetylfluorescein (DAF) Hydrolysis Rate

The lines represent the maximum and minimum experimentally determined limits. The points designated (Δ) represent experiments on a separate 1 to 3 day old BG suspension. The point at which the rate of spontaneous DAF hydrolysis and the bacterial detection limit of sensitivity coincides is noted at the lower left.









Bacterial age was not addressed in the Pinteno and Findl report; whereas, The BG from this work was taken from 1- to 2-day old suspension data.

As the bacterial suspension aged, a pronounced sensitivity in age delineation is observed above 5.0 x 10^4 cells/ml. Figure 2 shows a plot of age vs the relative hydrolysis rate of diacetylfluorescein for a concentration of approximately 5.0 x 10^6 cells/ml. An apparently exponential decay of the fluorescein emission is observed. This effect is interesting in that, despite the decrease in metabolic activity of the cell suspension with time, the viable B. subtilis population, as shown in Figure 1, appears to remain constant. Possible explanations of this anomaly include (a) genetic repression of the lipase-producing cellular machinery, (b) intracellular degradation of the metabolic machinery, and (c) cell wall modification due to aging. Therefore, the log plot of Figure 1 becomes generally useful for determining both the bacterial concentration and age. For the present buffer and detection system, Figure 1 shows clearly that the limit of bacterial concentration, in which the signal-to-noise ratio is twice that of the spontaneous hydrolysis of diacetylfluorescein, is 3.0×10^4 cells/ml, approximately. If this graph is to be a generally useful and predictive tool for a particular bacterial concentration and age, it must be independent of

microbial suspensions for that same microorganism. This thesis is apparently supported by a separate B. subtilis suspension of the same age (1 to 3 days) whose data (Δ) is superimposed upon the graph of the figure. These results indicate that for a sample analysis one need only measure the relative velocity of an enzymatic *in vivo* reaction in order to obtain the bacterial concentration from the previously prepared standard curve. At the same time, a rough bacterial age can be estimated.

In the second series of experiments, diacetylfluorescein concentrations were varied from 2.4 x 10^{-5} M to 4.8 x 10^{-8} M while, again, monitoring the rate of reaction. The viable B. subtilis bacterial concentration remained constant during the course of this study; hence a standard of 5.5 x 10^6 cells/ml assay solution was used. Because the intracellular hydrolysis of diacetylfluorescein obeys Michaelis-Menten kinetics,²⁸ the enzyme parameter, K_M was able to be evaluated for each daily set of runs from Hanes-Woolf plots.^{28,29} Since it was first postulated and then observed (see Figure 1) that lipase activity on diacetylfluorescein decreases as the bacterial suspension ages, the use of the Michaelis-Menten parameter was included in our investigation. Over the 1-month course of this study, relative K_M values varied only modestly (0.6 to 1.6 x 10^{-5} M) in a random fashion, with an average value of $1.17 \times 10^{-5} M$. This is to be expected because the Michaelis-Menten theory dictates that K_M is independent of the enzyme concentration;²⁸ therefore, the data suggest that the parameter might be used to more completely identify a specific bacterial-enzyme system. The data representing the separate BG suspension superimposed on Figure 1 yielded a K_M value of 1.38 x 10⁻⁵M, which is in reasonable agreement with the value cited above. It is of further interest to note that in other studies of intracellular diacetylfluorescein hydrolysis, K_M values of 3.6 x 10^{-5} M for fibroblast L cells, ²² 2.9 x 10^{-6} M for cultured mouse lymphoma cells, ²¹ and average intracellular fluorescein concentrations of 4.4 x 10^{-5} M and 2 x 10^{-5} M for leukemic and normal mouse lymphocytes, 30respectively, were reported. To the best of our knowledge, these represent the only quantitative Michaelis-Menten analyses of experiments with living mammalian and bacterial cells. $K_{\rm M}$ values of 3.6 x $10^{-5} {\rm M}$ and 7.0 x 10^{-6} M were reported for the cell-free extracts of the cultured mouse lymphoma cells mentioned above and an in vitro solution of porcine pancreas lipase, 23,31 respectively. At a K_M value of 1.17 x 10^{-5} M, the B. subtilis hydrolyzed diacetylfluorescein is within an order of magnitude of that found in the mammalian cells and the in vitro lipase experiment.

4. CONCLUSIONS

These encouraging early results suggest that further study concerning this technique as a valid approach to microorganism determination is warranted. From this work there emerges various parameters that appear to be useful in the determination of both concentration and age of a bacterium. These include the observed reaction rate and the Michaelis-Menten constant. In addition, the method suggests that sequential substrate usage could permit further refining of the process of uniquely identifying microorganisms. The characterization of a mixed bacteria sample, however, is significantly more germane. By the very nature of enzyme reactions, the substrates themselves shed light on the kind of bacteria present. Thus, a quick antibiotic screen and/or biochromatographic techniques utilizing different retention times for different microorganisms* could also be used in bacterial separation and determination. Greater bacterial resolution could be achieved by coupling a low-power source of laser excitation with a microfluorimeter in a microscopic stage system. This is the subject of continuing work by the authors.

云れとたららららい、西日はもうとうかい

The second s

*D. B. Greenberg, University of Cincinnati, and G. Janauer, State University of NY at Binghamton. 1983.

Blank

K

LITERATURE CITED

1. Halmann, M., Velan, B., and Sery, T. Appl. Environ. Microbiol. 34, 473 (1977).

2. Drow, D. L., Maki, D. G., and Manning, D. D. J. Clin. Microbiol. 10, 442 (1979).

3. Wolters, G., Kuijpers, L. P., Kacaki, J., and Schuurs, H.W.M. J. Infect. Dis. 136, s311 (1977).

4. Goldman, M. Ann. N. Y. Acad. Sci. 177, 439 (1971).

5. Danielsson, D., Nathorst-Windahl, G., and Salden, T. Ibid., 23.

6. Pavlova, M. T., Beauvais, E., Brezenski, F. T., and Litsky, W. Appl. Microbiol. 23, 571 (1972).

7. Hinsberg, W. D., Milby, K. H., Lidofsky, S. D., and Zare, R. N. Proc. SPIE. J. A. Gelbwachs, ed. 286, 132 (1981).

8. Imsaka, T., and Zare, K. N. Anal. Chem. 51, 2082 (1979).

9. Betz, H., Buhay, H., Fenters, J., and Vana, S. ARCSL-CR-81039. May 1981. UNCLASSIFIED Report.

10. Anhalt, J. P. Anal. Chem. 47, 219 (1975).

11. Kellenberger, G. and Kellenberger, E. Virology 3, 275 (1957).

12. Pijper, A. J. Path. Bact. 57, 1 (1945).

13. Smith, P. B., and Cherry, W. B. J. Inf. Dis. 96, 34 (1955).

14. Feltham, R.K.A., and Sneath, P.H.A. J. Gen. Microbiol. 128, 713 (1982).

15. Kelley, R. W., and Kellogg, S. T. Appl. Environ. Microbiol. 35, 507 (1978).

16. Smith, P. B., Gavin, T. L., Isenberg, H. D., Sonnenwirth, A., Taylor, W. I., Washington, J. A. II, and Balows, A. J. Clin. Microbiol. \$, 657 (1978).

17. Friedman, R. B., and MacLowry, J. Appl. Microbiol. 26, 314 (1973).

18. Friedman, R. B., Bruce, D., MacLowry, J., and Brenner, V. Am. J. Clin. Pathol. 60, 395 (1973).

19. Russell, A. P., DeMaria, A., Johns, M., and Westcott, V. C. J. Tribology (1983).

20. Van Dilla, M. A., Langlois, R. G., Pinkel, D., Yajko, D., and Hadley, W. K. Science 220, 620 (1983).

21. Rotman, B., and Papermaster, B. W. Proc. Nat'l. Acad. Sci. 55, 135 (1966).

22. Sernetz, M. Fluorescence Techniques in Cell Biology, p. 243. Springer-Verlag, Berlin. 1973.

23. Guilbault, G. G., and Hieserman, J. Anal. Chem. 41, 2006 (1969).

24. Kramer, D. N., and Guilbault, G. G. Ibid., 35, 588 (1963).

25. Guilbault, G. G., and Kramer, D. N. Ibid., 36, 409 (1964).

26. Mitz, M. A., Blanchard, G. C., and Deacon, T. E. U. S. Army Biological Laboratories. DA-18-064-CML-2842. July 1963. UNCLASSIFIED Report.

27. Pinteno, F., and Findl, E. Naval Research Contract N00014-78-C-0713. February 1981. UNCLASSIFIED Report.

28. Segel, I. H., ed. Biochemical Calucations. 2nd Ed. Wiley-Interscience. 1975.

29. Segel, I. H., ed. Enzyme Kinetics. Chapter 4. Wiley-Interscience. 1975.

30. Szollosi, J., Kertai, P., Somogyi, B., and Damjanovich, S. J. Histochem. Cytochem. 29, 503 (1981).

31. Sherman, W. R., and Stanfield, E. F. Biochem. J. 102, 905 (1967).

DISTRIBUTION LIST 2

Names	Coples	Names	Copie
CHEMICAL RESEARCH AND DEVELOPMEMNT	CENTER	Federal Emergency Management Agency	
		Office of Research/NPP	
ATTN: DRSMC-CLB (A)	1	ATTN: David W. Bensen	1
ATTN: DRSMC-CLB-C (A)	1	Washington, DC 20472	
AFTN: DRSMC-CLB-PO (A)	1		
ATTN: DRSMC-CLB-R (A)	1	HQDA (DAMA-CSS-C)	1
ATTN: DRSMC-CLB-R(M) (A)	1	Washington, DC 20310	
ATTN: DRSMC-CLB-R(S) (A)	1		
ATTN: DRSMC-CLB-T (A)	1	HQ Sixth US Army	
ATTN: DRSMC-CLC-B (A)	1	ATTN: AFKC-OP-NBC	1
AFTN: DRSMC-CLC-C (A)	1	Presidio of San Francisco, CA 94129	
ATTN: DRSMC-CLC-E (A)	1		
AFTN: DRSMC-CLF (A)	1	Commander	
ATTN: DRSMC-CLJ-IL (A)	2	DARCOM, STITEUR	
ATTN: DRSMC-CLJ-IR (A)	1	ATTN: DRXST-STI	1
ATTN: DRSMC-CLJ-M (A)	1	Box 48, APO New York 09710	
ATTN: DRSMC-CLN (A)	1	•	
ATTN: DRSMC-CLT (A)	1	Commander	
ATTN: DRSMC-CLW-C (A)	1	USASTCFEO	
ATTN: DRSMC-CLW-P (A)	1	ATTN: MAJ Mikeworth	1
ATTN: DRSMC-CLY-A (A)	1	APO San Francisco 96328	•
ATTN: DRSMC-CLY-R (A)	6		
		Commander	
COPIES FOR AUTHOR(S)		USA Nuclear & Chemical Agency	
ATTN: DRSMC-CLB-R (A)	8	ATTN: MONA-WE	1
RECORD COPY: DRSMC-CLB-A (A)	1	7500 Backlick Rd, Bidg 2073	•
	•	Springfield, VA 22150	
DEPARTMENT OF DEFENSE		op ng 1010, 11 22130	
		Army Research Office	
Administrator		ATTN: DRXRO-CB (Dr. R. Ghirardelli)	1
Defense Technical Information Cent	or	P.0. Box 12211	•
ATTN: DTIC-DDA-2	er 2	Research Triangle Park, NC 27709	
Cameron Station, Building 5	۷.	Nesedicii il langie Fark, NG 27709	
Alexandria, VA 22314		OFFICE OF THE SURGEON GENERAL	
Director		Commander	
Defense intelligence Agency		USA Medical Bioengineering Research	
ATTN: DB-4G1	1	and Development Laboratory	
Washington, DC 20301		ATTN: SGRD-UBD-AL, Bldg 568	1
-		Fort Detrick, Frederick, MD 21701	
Commander			
USASED, USAINSCOM		Commander	
ATTN: IAFM-SED-III	1	USA Medical Research Institute of	
Fort Meade, MD 20755	-	Chemical Defense	
		ATTN: SGRD-UV-L	1
DEPARTMENT OF THE ARMY		Aberdeen Proving Ground, MD 21010	•
HODA (DAMO-NCC)	1		
	•		

いる

Commander	
US Army Environmental Hygiene Agency	
ATTN: HSHB-O (B. Donovan)	2
Aberdeen Proving Ground, MD 21010	
US ARMY MATERIEL DEVELOPMENT AND READINESS COMMAND	
Commander	
HQ, DARCOM	
AFIN: DRCED (BG Robinson)	1
5001 Eisenhower Ave	
Alexandria, VA 22333	
· · · · · ·	
Commander	
USA Materiel Development and	
Readiness Command	
ATTN: DRCSF-P	1
5001 Elsenhower Ave	
Alexandria, VA 22333	
Project Manager Smoke/Obscurants	
AT IN: DRCPM-SMK-S	3
Aberdeen Proving Ground, MD 21005	
,	
Commander	
USA Foreign Science & Technology Center	
ATTN: DRXST-MT3	1
220 Seventh St., NE	
Charlottesville, VA 22901	
······································	
Director	
USA Materiel Systems Analysis Activity	
ATTN: DRXSY-MP	1
ATTN: DRXSY-CR (Mr. Metz)	1
Aberdeen Proving Ground, MD 21005	
•	
Commander	
USA Missile Command	
Redstone Scientific Information Center	
ATTN: DRSMI-RPR (Documents)	1
Redstone Arsenal, AL 35898	
-	
Director	
DARCOM Field Safety Activity	
ATTN: DRXOS-C	1
Charlestown, IN 47111	

Commander USA Natick Research and Development Laboratories ATTN: DRDNA-0 1 ATTN: DRDNA-IC 1 ATTN: DRDNA-IM 1 ATTN: DRDNA-ITF (Dr. Roy W. Roth) 2 Natick, MA 01760 US ARMY ARMAMENT, MUNIFIONS AND CHEMICAL COMMAND Commander USA Armament, Munitions and Chemical Command ATTN: DRCSM-ASN (R) 1 ATTN: DRCSM-IRW (R) 1 Rock Island, IL 61299 Commander USA Dugway Proving Ground ATTN: Technical Library (Docu Sect) 1 Dugway, UT 84022 US ARMY ARMAMENT RESEARCH AND DEVELOPMENT CENTER Commander USA Armament Research and **Development Center** ATTN: DRSMC-LCA-L (D) AFTN: DRSMC-LCE-C (D) ATTN: DRSMC-LCU-CE (D) ATTN: DRSMC-SCA-T (D) ATTN: DRSMC-SCM (D) AFTN: DRSMC-SCP (D) ATTN: DRSMC-SCS (D) ATTN: DRSMC-TDC (D) (Dr. D. Gyorog) ATTN: DRSMC-TSS (D) 2 ATTN: DRCPM-CAWS-AM (D) 1 Dover, NJ 07801 Armament Research and Development Center USA AMCCOM ATTN: DRSMC-TSE-OA (Robert Thresher) 1 National Space Technology Laboratories NSTL Station, MS 39529 Commander USA AMCCOM

18

ATTN: DRSMC-QAC-E (A)

Aberdeen Proving Ground, MD 21010

۱

```
Commander
USA Technical Detachment
US Naval EOD Technology Center
Indian Head, MD 20640
US ARMY TRAINING & DOCTRINE COMMAND
Commandant
USA Infantry School
ATTN: CTDD, CSD, NBC Branch
Fort Benning, GA 31905
Commandant
USA Missile & Munitions Center
    and School
ATTN: ATSK-CM
ATTN: ATSK-TME
Redstone Arsenal, AL 35898
Commander
USA Logistics Center
ATTN: ATCL-MG
ATTN: DLSIE
Fort Lee, VA 23801
Commandant
USA Chemical School
ATTN: ATZN-CM-C
ATTN: ATZN-CM-AFL
ATTN: ATZN-CM-TPC
Fort McClellan, AL 36205
Commander
USAAVNC
ATTN: ATZQ-D-MS
Fort Rucker, AL 36362
Commander
USA infantry Center
ATTN: ATSH-CD-MS-C
Fort Benning, GA 31905
Commander
USA Training and Doctrine Command
ATTN: ATCD-N
Fort Monroe, VA 23651
```

Commander USA Armor Center ATTN: ATZK-CD-MS ATTN: ATZK-PPT-PO-C Fort Knox, KY 40121

Commander 1 USA Combined Arms Center and Fort Leavenworth ATTN: ATZL-CAM-IM Fort Leavenworth, KS 66027 US ARMY TEST & EVALUATION COMMAND Commander USA Test & Evaluation Command 1 ATTN: DRSTE-CT-T Aberdeen Proving Ground, MD 21005 DEPARTMENT OF THE NAVY Chief of Naval Research 1 AFTN: Code 441 1 800 N. Quincy Street Arlington, VA 22217 Project Manager 1 Theatre Nuclear Warfare Project Office AFTN: PM-23 (Dr. Patton) 1 ATTN: TN-09C Navy Department Washington, DC 20360 t Commander 2 Naval Explosive Ordnance Disposal 2 Technology Center AFTN: AC-3 Indian Head, MD 20640 Commander Naval Surface Weapons Center 1 Code G51 Dahlgren, VA 22448 Chief, Bureau of Medicine & Surgery 1 Department of the Navy AFTN: MED 3C33 Washington, DC 20372 Commander Naval Air Development Center ATTN: Code 2012 (Dr. Robert Helmbold) Warminster, PA 18974 US MARINE CORPS 1 Commandant HQ, US Marine Corps AFTN: Code LMW-50 Washington, DC 20380

1

1

1

1

1

1

1

1

Commanding General Marine Corps Development and Education Command ATTN: Fire Power Division, D091 1 Quantico, VA 22134 DEPARTMENT OF THE AIR FORCE

Department of the Air Force Headquarters Foreign Technology Division ATTN: TQTR Wright-Patterson AFB, OH 45433

ASD/AESD Wright-Patterson AFB, OH 45433

AFAMRL/IS ATIN: COL Johnson Wright-Patterson AFB, OH 45433

AFAMRL/HE ATTN: Dr. Clyde Reploggie Wright-Patterson AFB, OH 45433

AFWAL/FIEEC (Wendell Banks) Wright-Patterson AFB, OH 45433

HQ AFSC/SDZ ATTN: CPT D. Riediger Andrews AFB, MD 20334

HQ, AFSC/SDNE Andrews AFB, MD 20334

HQ, AFSC/SGB Andrews AFB, DC 20334

HQ, NORAD ATTN: J-3TU Peterson AFB, CO 80914

HQ AFTEC/TEL Kirtland AFB, NM 87117

USAF TAWC/THL Eglin AFB, FL 32542

AFATL/DLV Eglin AFB, FL 32542

USAF SC ATTN: AD/YQ ATTN: AD/YQO (MAJ Owens) Eglin AFB, FL 32542 AD/XRO Eqlin AFB, FL 32542 USAFSAM/VN Deputy for Chemical Defense ATTN: Dr. F. Westey Baumgardner Brooks AFB, TX 78235 AMD/RDTK ATTN: LTC T. Kingery Brooks AFB, TX 78235 AMD/RDSM Brooks AFB, TX 78235 AMD/RD SX Brooks AFB, TX 78235

1

1

1

1

1

1

1

1

1

1

1

1

2

OUTSIDE AGENCIES

1

1

1

1

1

1

1

1

1

1

1

1

Battelle, Columbus Laboratories AITN: IACTEC 505 King Avenue Columbus, OH 43201

Toxicology Information Center, JH 652 National Research Council 2101 Constitution Ave., NW Washington, DC 20418

US Public Health Service Center for Disease Control ATTN: Logging Control Officer Mrs. M. Brocato (W.L.Webb) Atlanta, GA 30333

Director Central Intelligence Agency AFTN: AMR/ORD/DD/S&F Washington, DC 20505

ADDITIONAL ADDRESSEE

Commandant Academy of Health Sciences, US Army AFFN: HSHA~CDH AFFN: HSHA~IPM Fort Sam Houston, FX 78234

.

۰.) بر 7-84 A RANKI (CO)