

DTIC DOCUMENTATION PAGE

AD-A222 703

ELECTE
MAY 25 1990
D

1b RESTRICTIVE MARKINGS
NA

3. DISTRIBUTION / AVAILABILITY OF REPORT
Distribution Unlimited

2b DECLASSIFICATION / DOWNGRADING SCHEDULE
NA

4 PERFORMING ORGANIZATION REPORT NUMBER(S)

5. MONITORING ORGANIZATION REPORT NUMBER(S)
NA

6a NAME OF PERFORMING ORGANIZATION
University of Kansas

6b. OFFICE SYMBOL
(If applicable)
NA

7a. NAME OF MONITORING ORGANIZATION
Office of Naval Research

6c. ADDRESS (City, State, and ZIP Code)
Lawrence, KS 66045

7b ADDRESS (City, State, and ZIP Code)
800 N. Quincy St.
Arlington, VA 22217-5000

8a. NAME OF FUNDING / SPONSORING ORGANIZATION
Office of Naval Research

8b. OFFICE SYMBOL
(If applicable)
ONR

9 PROCUREMENT INSTRUMENT IDENTIFICATION NUMBER
N00014-86-K-0862

8c ADDRESS (City, State, and ZIP Code)
800 N. Quincy St.
Arlington, VA 22217-5000

10 SOURCE OF FUNDING NUMBERS
PROGRAM ELEMENT NO PROJECT NO TASK NO WORK UNIT ACCESSION NO
61153N RR04106 NR 44g016

11 TITLE (Include Security Classification)
Development of Synthetic Catalysts for Peptide Bond Cleavage
Synthesis and Complete Kinetic Analysis of Compounds 6A, 7A, 8A

12 PERSONAL AUTHOR(S)
Mertes, Kristin Bowman, P.I.

13a. TYPE OF REPORT
Final

13b TIME COVERED
FROM 8/86 TO 2/90

14. DATE OF REPORT (Year, Month, Day)
May 15, 1990

15. PAGE COUNT
6

16 SUPPLEMENTARY NOTATION

COSATI CODES		
FIELD	GROUP	SUB-GROUP

17 SUBJECT TERMS (Continue on reverse if necessary and identify by block number)
Enzyme mimics, Supramolecular, Carboxypeptidase A,
Polyammonium macrocycles, Synthetic catalysts,
Peptide Bond Cleavage, Kinetic Analysis, Ring Systems

19 ABSTRACT (Continue on reverse if necessary and identify by block number)
Synthetic mimics for carboxypeptidase A will be synthesized and the structural and chemical factors responsible for catalytic peptidase activity will be probed. Dicitopic macrocyclic receptors have been designed which incorporate the salient features of the enzyme analog, namely high affinity complex formation, general base and general acid catalysis, and covalent catalysis. Once synthesized the resulting macrocycle-metal ion complexes should non-specifically promote the hydrolysis of C-terminal peptide bonds. The initial macrocycles will have several types of coordination sites: nitrogen-containing heterocycles, ammonium and ether oxygens. One side of the ditopic receptor will preferentially bind zinc(II) ion, the other the peptide substrate.

DISTRIBUTION STATEMENT A
Approved for public release;
Distribution Unlimited

20. DISTRIBUTION / AVAILABILITY OF ABSTRACT
 UNCLASSIFIED/UNLIMITED SAME AS RPT. DTIC USERS

21. ABSTRACT SECURITY CLASSIFICATION
(U)

22a NAME OF RESPONSIBLE INDIVIDUAL
Dr. N. Mafron

22b. TELEPHONE (Include Area Code)
202/696-4760

22c. OFFICE SYMBOL
ONR

May 15, 1990

FINAL REPORT ON CONTRACT N00014-86-K-0862

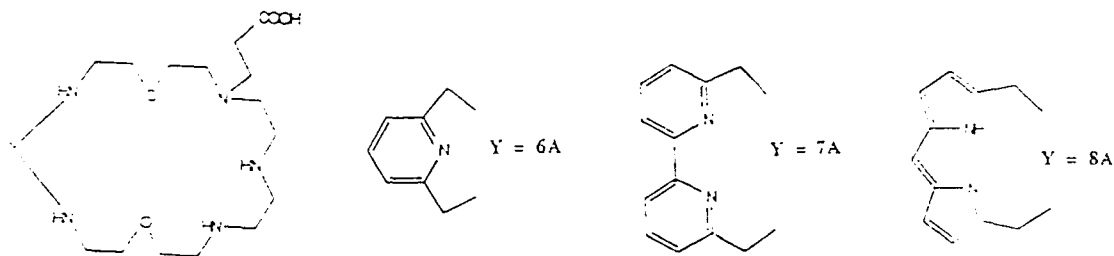
PRINCIPAL INVESTIGATOR: Kristin Bowman Mertes

CONTRACT TITLE: Development of Synthetic Catalysts for Peptide Bond Cleavage: Synthesis and Complete Kinetic Analysis of Compounds 6A, 7A, 8A

START DATE: 6 August 1986

RESEARCH OBJECTIVE:

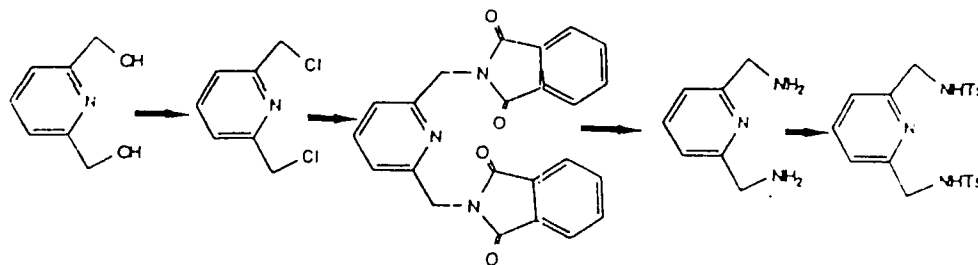
The synthesis of ditopic receptors 6A, 7A, 8A, designed by considering the important interactive features of carboxypeptidase A (CPA) was the goal of this project. Three basic macrocyclic ligands with site specificity for zinc(II) incorporation and functionalized podando groups for covalent catalysis were attempted. Two of these were synthesized and examined for hydrolase activity. The design of the macrocycles was to allow for the critical assessment of the importance of the interactive sites within the natural enzyme, from the general acid catalysis provided by the arginine and tyrosine residues and the zinc(II) ion to the general base role of the Glu-270 residue.



ACCOMPLISHMENTS:

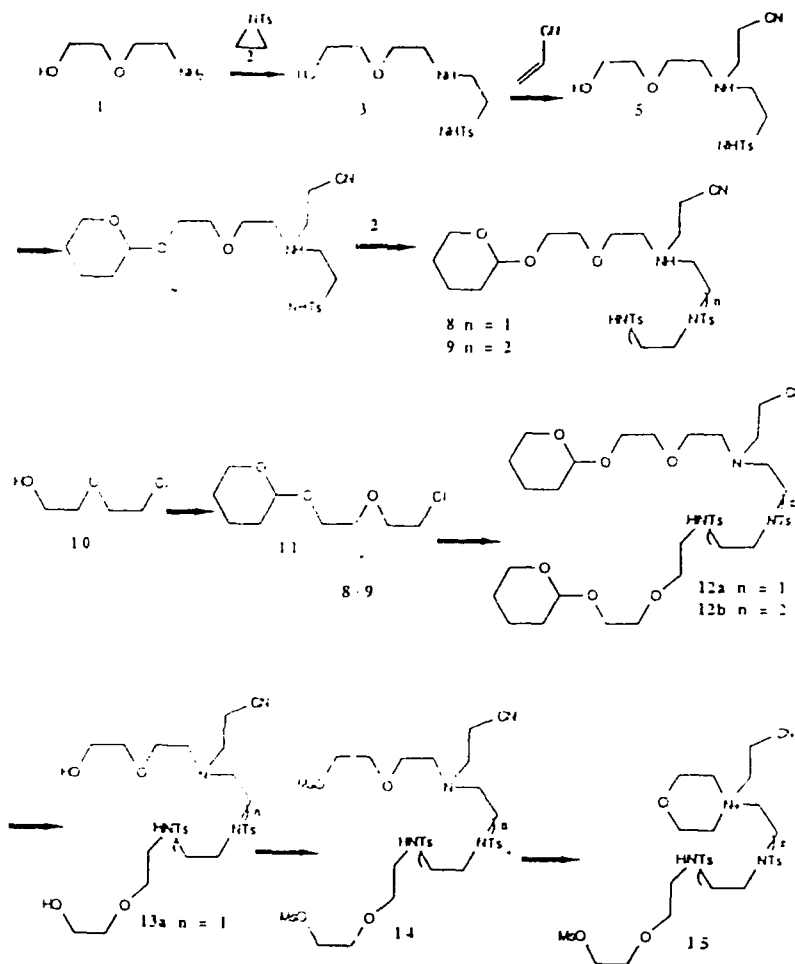
Synthesis

The synthetic efforts could be divided into the synthesis of the two halves of the molecule, the "eastern" and "western" half. The synthesis of the "western" half was relatively straightforward for compound 6A and was accomplished in the first year of the contract (Scheme 1):



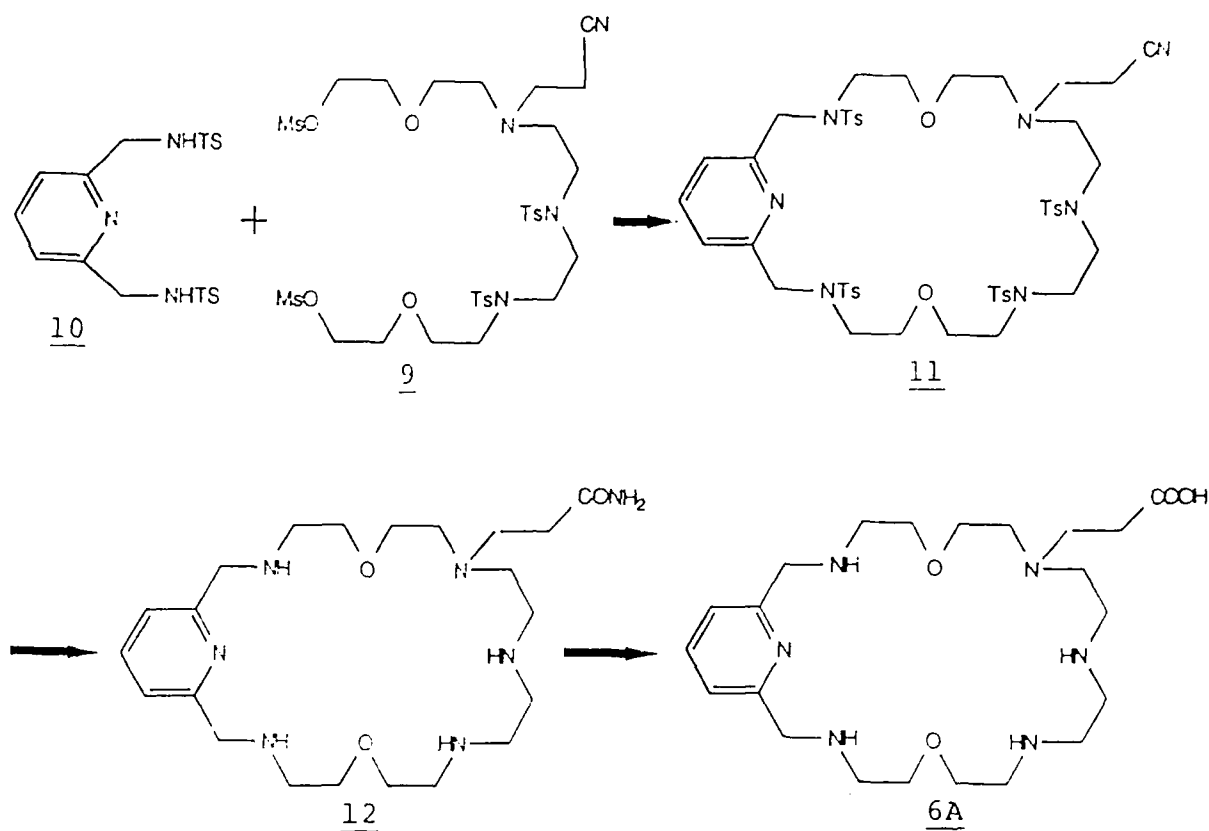
Scheme 1

In the synthesis of the "eastern" half of the molecule several synthetic problems were encountered. One of the major difficulties was a polymerization reaction where more than one unit of the tosylaziridine added to the azoxy chain. This was further complicated by difficulties in separating the three major isomers. In proceeding from 7 to 8 the tosyl aziridine reacted further with the triamine product to give tetraamines and higher analogs. A second problem was found to occur in the mesylation of 13a, in which 14 was lost due to an undesired ring closure leading to 15.



These problems were solved in year two by using a convergent sequence as shown in Scheme 2. Here 3 and 6 are combined to give 7. This avoided the difficulties in extension of 5, which could not be controlled.

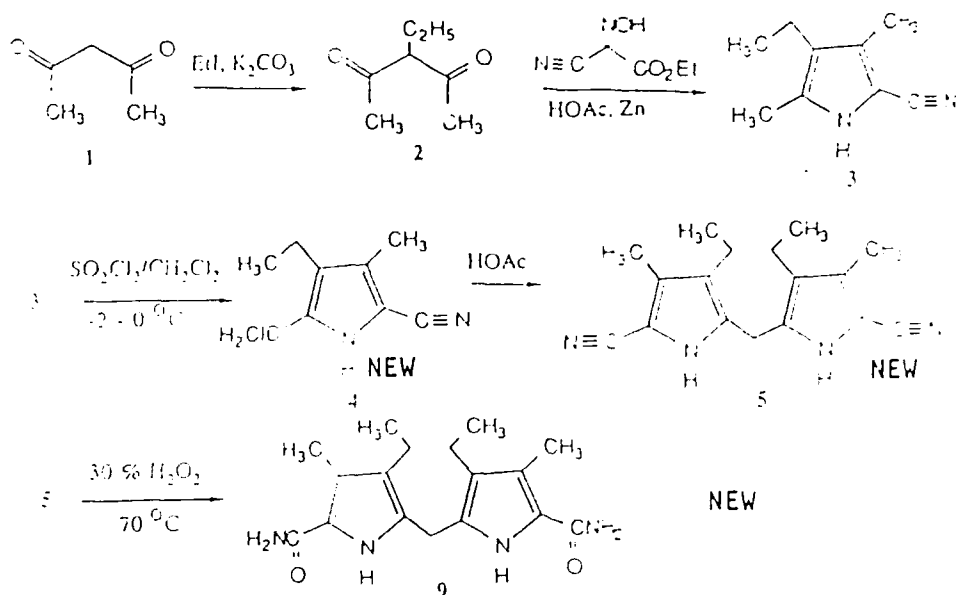
Accession For	
NTIS GRA&I	<input checked="" type="checkbox"/>
DTIC TAB	<input type="checkbox"/>
Unannounced	<input type="checkbox"/>
Justification	
By _____	
Distribution/	
Availability Codes	
Dist	Avail and/or Special
A-1	



Scheme 4.

Testing for catalytic activity was completed on **6A** and **7A** and their precursor amides (e.g., **12**, Scheme 4) using the hydrolysis of *O*-(*trans*-*p*-chlorocinnamoyl)-*L*- β -phenyllactic acid at 80 °C and pH 7.0. The results are shown in Table 1. Only one of the compounds showed any evidence of catalytic activity, **6A**. These studies and related studies in phosphoryl transfer catalysis indicate that steric effects play a crucial role in the catalytic activity of these macrocycles. Hence, less sterically hindered substrates are more desirable for testing these compounds.

The synthesis of compound **8A** was not achieved due to the extreme sensitivity of the dipyrromethane portion of the macrocycle. Several new pyrrole and dipyrromethanes were synthesized (Scheme 5), however, and their chemistry is currently being investigated.



Scheme 5

While the compounds synthesized were not catalytically active, a variety of new synthetic techniques were achieved. The critical point in these studies to be made is that ring size must be optimal and steric hindrance minimal for any reaction to occur. These findings are corroborated by modeling studies in phosphoryl transfer reactions, where the macrocycles are only active for 21- and 24- membered ring systems, and catalysis is greatly reduced in the presence of podando side chains.

PUBLICATIONS AND REPORTS:

Gu, K.; Mertes, K. B.; Mertes, M. P., Strategy for the Synthesis of Unsymmetrical N-Substituted Polyazamacrocycles, *Tetrahedron Lett.* 1989, **30**, 1323-1326.

Table 1. First Order Rate Constants (k_{obsd}) for the Hydrolysis of O-(trans-p-chlorocinnamoyl)-L- β -phenyllactic Acid at pH 7 and 80 °C.

macrocycle	metal	k_{obsd} (min^{-1} , $\times 10^5$)
none	none	3.68
none	Zn ²⁺	4.99
24pyCONH2, 10	none	4.02
24pyCONH2, 10	Zn ²⁺	4.24
24pyCOOH, 6A	none	3.72
24pyCOOH, 6A	Zn ²⁺	5.82
27dipyCONH2, 11	none	2.96
27dipyCONH2, 11	Zn ²⁺	2.45
27dipyCOOH, 7A	none	2.50
27dipyCOOH, 7A	Zn ²⁺	2.65

Distribution List for Annual and Final Reports

1. Put a cover page (Form DD 1473) on your report and attach a copy of the distribution list. Mail one copy of the report to each person on the contractor subset list attached on which your name appears. The other subset list is for your information only. Please don't forget to attach this distribution list to your report - otherwise the folks below think they have mistakenly received the copy meant for the Molecular Biology Program and forward it to us.
2. Mail two copies to (include a DTIC Form 50 with these two copies too)
Administrator
Defense Technical Information Center
Building 5, Cameron Station
Alexandria, VA 22314
3. Mail one copy to each of the following:
 - (a) Dr. Michael Marron
ONR Code 1141
Molecular Biology Program
800 N. Quincy Street
Arlington, VA 22217-5000
 - (b) Administrative Contracting Officer
ONR Resident Representative
(address varies - see copy of your grant)
 - (c) Director,
Applied Research Directorate
ONR Code 12
800 N. Quincy Street
Arlington, VA 22217-5000
 - (d) Director
Office of Naval Technology
Code 22
800 N. Quincy Street
Arlington, VA 22217-5000
 - (e) Director
Chemical and Biological Sci Div
Army Research Office
P. O. Box 12211
Research Triangle Park, NC 27709
 - (f) Life Sciences Directorate
Air Force Office of Scientific Research
Bolling Air Force Base
Washington, DC 20332
 - (g) Director
Naval Research Laboratory
Technical Information Div, Code 2627
Washington, DC 20375

AMZEL, L. Mario
Department of Biophysics
Johns Hopkins School of Medicine
725 North Wolfe Street
Baltimore, MD 21205

ANDERSEN, Niels H.
Department of Chemistry
University of Washington
Seattle, WA 98195

ARNOLD, Frances H.
Dept of Chemical Engineering
California Institute of Technology
Pasadena, CA 91125

AUGUST, J. Thomas
Department of Pharmacology
Johns Hopkins Medical School
725 North Wolfe Street
Baltimore, MD 21205

BEVERIDGE, David L
Department of Chemistry
Wesleyan University
Hall-Altwater Laboratories
Middletown, CT 06457

BRAMSON, H. Neal
Department of Biochemistry
Univ of Rochester Medical Center
601 Elmwood Avenue
Rochester, NY 14642

BRUCE, Thomas C.
Department of Chemistry
University of California-Santa
Barbara
Santa Barbara, CA 93106

CASE, Steven T.
Department of Biochemistry
Univ of Mississippi Medical Center
2500 North State Street
Jackson, MS 39216-4505

CHANG, Eddie L.
Bio/Molecular Engineering
Naval Research Laboratory
Code 6190
Washington, D.C. 20375-5000

CHRISTIANSON, David W.
Department of Chemistry
University of Pennsylvania
231 South 34th Street
Philadelphia, PA 19104-6323

CORDINGLEY, John S.
Department of Molecular Biology
University of Wyoming
Box 3944 University Station
Laramie, WY 82071

DeGRADO, William F.
E. I. du Pont de Nemours & Co
Central R & D, Experimental Station
P. O. Box 80328
Wilmington, DE 19880-0328

EVANS, David R.
Department of Biochemistry
Wayne State Univ School of Medicine
540 E. Canfield Street
Detroit, Michigan 48201

FEIGON, Juli F.
Department of Chem & Biochemistry
UCLA
405 Hilgard Avenue
Los Angeles, CA 90004-1569

FICHT, Allison R.
Dept of Med Biochem & Genetics
Texas A&M University
College Station, TX 77843

FRAUENFELDER, Hans
Department of Physics
University of Illinois
Urbana, IL 61801

GABER, Bruce
Naval Research Laboratory
Bio/Molecular Engineering Branch
Code 6190
Washington, DC 20375

GETZOFF, Elizabeth D.
Scripps Clinic & Research Foundation
Department of Molecular Biology
10666 North Torrey Pines Road
La Jolla, CA 92037

GOODMAN, Eugene M.
Biomedical Research Institute
University of Wisconsin
P. O. Box 2000
Kenosha, WI 53141

HO, Pui Shing
Department of Biochemistry and
Biophysics
Oregon State University
Corvallis, OR 97331

HOGAN, Michael E.
Baylor Center for Biotechnology
4000 Research Forest Drive
The Woodlands, TX 77381

HONIG, Barry
Columbia University
Dept of Biochem and Molec Biophys
630 West 168th St.
New York, NY 10032

HOPKINS, Paul B.
Department of Chemistry
University of Washington
Seattle, WA 98195

KAHNE, Daniel
Department of Chemistry
Princeton University
Princeton, NJ 08544

KEMP, Robert G.
Chicago Medical School
Dept of Biological Chemistry
3333 Green Bay Rd.
North Chicago, IL 60064

KHORANA, Gobind H.
Department of Biology
MIT
77 Massachusetts Ave.
Cambridge, MA 02139

KIM, Sangtae
Chemical Engineering
University of Wisconsin
1415 Johnson Drive
Madison, WI 53706

LANSBURY, Peter T.
Department of Chemistry
MIT
Cambridge, MA 02139

LAURSEN, Richard A.
Chemistry Department
Boston University
590 Commonwealth Avenue
Boston, MA 02215

LENZ, Robert W.
Chemical Engineering Department
University of Massachusetts
Amherst, MA 01003

LEWIS, Randolph V.
Molecular Biology Department
University of Wyoming
University Station Box 3944
Laramie, WY 82071

LINDSAY, Stuart M.
Department of Physics
Arizona State University
Temp, AZ 85278

LOEB, George I.
David W. Taylor Research Center
Code 2841
Annapolis, MD 21402-5067

MASILAMANI, Divakar
Biotechnology Department
Allied-Signal Inc.
P. O. Box 1021R
Morristown, NJ 07960

McCONNELL, Harden M.
Stanford University
Department of Chemistry
Stanford, CA 94305

McELROY, Willam D.
Department of Chemistry
University of California - San Diego
La Jolla, CA 92093-0601

MERTES, Kristin Bowman
University of Kansas
Dept of Chemistry
Lawrence, Kansas 66045

NAGUMO, Mark
Bio/Molecular Engineering Branch
Naval Research Laboratory
Code 6190
Washington, DC 20375-5000

OLIVERA, Baldomero M.
Department of Biology
University of Utah
Salt Lake City, UT 84112

PABO, Carl O.
Department of Biophysics
Johns Hopkins University
School of Medicine
Baltimore, MD 21205

PRENDERGAST, Franklyn G.
Dept of Biochemistry & Molec Biol
Mayo Foundation
200 First St. SW
Rochester, MN 55905

PUGH, Jr., Edward N.
Department of Psychology
University of Pennsylvania
3815 Walnut Street
Philadelphia, PA 19104-6196

RACKOVSKY, Shalom R.
Department of Biophysics
University of Rochester
School of Medicine and Dentistry
Rochester, NY 14642

RAJAN, K. S.
Illinois Institute of Technology
Research Institute
10 W. 35th St.
Chicago, IL 60616

REINISCH, Lou
Laser Biophysics Center
Uniformed Services University
4301 Jones Bridge Road
Bethesda, MD 20814

RICH, Alexander
MIT Department of Biology
Cambridge, MA 02139

RICHARDS, J. H.
California Institute of Technology
Division of Chemistry and Chemical
Engineering
Pasadena, CA 91125

ROTHSCHILD, Kenneth J.
Department of Physics
Boston University
590 Commonwealth Avenue
Boston, MA 02215

SCHULTZ, Peter G.
Department of Chemistry
University of California-Berkeley
Bekeley, CA 94720

SEEMAN, Nadrian
Department of Chemistry
New York University
New York, NY 10003

SELSTED, Michael E.
UCLA
Dept of Medicine
37-055 CHS
Los Angeles, CA 90024

SIGMAN, David S.
UCLA School of Medicine
Dept of Biological Chemistry
Los Angeles, CA 90024

SIKES, Steven C.
Department of Biological Sciences
University of South Alabama
Mobile, AL 36688

SINSKEY, Anthony J.
Laboratory of Applied Microbiology
MIT Department of Biology
Cambridge, MA 02139

STEWART, James M.
Department of Chemistry
University of Maryland
College Park, MD 20742

STEWART, John M.
Department of Biochemistry
University of Colorado
Health Science Center
Denver, CO 80262

TURNER, Douglas H.
Department of Chemistry
University of Rochester
Rochester, NY 14627

URRY, Dan W.
Laboratory of Molecular Biophysics
University of Alabama
P. O. Box 311
Birmingham, AL 35294

WAITE, J. Herbert
College of Marine Studies
University of Delaware
Lewes, DE 19958

WARD, Keith B.
Naval Research Laboratory
Code 6030
Washington, DC 20375

WARSHEL, Arieh
Department of Chemistry
University of Southern California
University Park
Los Angeles, CA 90089-0482

WATT, Gerald D.
Dept of Chemistry & Biochemistry
University of Colorado
Campus Box 215
Boulder, CO 80309-0215