

REPORT DOCUMENTATION PAGE			Form Approved OMB NO. 0704-0188	
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1. AGENCY USE ONLY (Leave Blank)		2. REPORT DATE		3. REPORT TYPE AND DATES COVERED
4. TITLE AND SUBTITLE			5. FUNDING NUMBERS	
6. AUTHOR(S)				
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES)			8. PERFORMING ORGANIZATION REPORT NUMBER	
9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES) U. S. Army Research Office P.O. Box 12211 Research Triangle Park, NC 27709-2211			10. SPONSORING / MONITORING AGENCY REPORT NUMBER	
11. SUPPLEMENTARY NOTES The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision, unless so designated by other documentation.				
12 a. DISTRIBUTION / AVAILABILITY STATEMENT Approved for public release; distribution unlimited.			12 b. DISTRIBUTION CODE	
13. ABSTRACT (Maximum 200 words)				
14. SUBJECT TERMS			15. NUMBER OF PAGES	
			16. PRICE CODE	
17. SECURITY CLASSIFICATION OR REPORT UNCLASSIFIED	18. SECURITY CLASSIFICATION ON THIS PAGE UNCLASSIFIED	19. SECURITY CLASSIFICATION OF ABSTRACT UNCLASSIFIED	20. LIMITATION OF ABSTRACT UL	

NSN 7540-01-280-5500

Standard Form 298 (Rev.2-89)
Prescribed by ANSI Std. Z39-18
298-102

Enclosure 1

REPORT DOCUMENTATION PAGE (SF298)
(Continuation Sheet)

Generated the 3D structure of the zinc-bound botulinum neurotoxin serotype A in complex with a nanomolar peptidic inhibitor

Developed nonpeptidic inhibitor leads of anthrax lethal factor in collaboration with Dr. Stephen H. Leppla of the PC Program

Developed the most potent, selective, nonpeptidic inhibitor lead of botulinum neurotoxin serotype A in collaboration with Dr. James J. Schmidt and Major Charles Millard of the USAMRIID

Successfully predicted the 4D structure of a SARS cysteine proteinase complexed with its peptide substrate 20 days after the release of the SARS genome by the CDC (released to the Protein Data Bank, PDB code: 1P76)

2. Include results from collaboration with other PI's within the PC program as well as other government agencies

Identified inhibitor leads of cofactor-independent phosphoglycerate mutase in collaboration with Drs. Terrance Leighton and Mark J. Jedrzejas of the PC Program

Identified the mechanism for the solubility of problem of B-DNA binders in collaboration with Dr. Richard Tanaka of the PC Program

Identified potential drug targets of a known antiviral agent in collaboration with Dr. Jeffrey C. Bottaro of the PC Program

Described in previous reports

3. Include filed and issued patents and publications

Discovery of a New Inhibitor Lead of Adenovirus Proteinase: Steps Toward Selective, Irreversible Inhibitors of Cysteine Proteinases, Yuan-Ping Pang, Kun Xu, Thomas M. Kollmeyer, Emanuele Perola, William J. McGrath, Dave T. Green, and Walter F. Mangel, *FEBS Lett.*, 502, 93-97 (2001)

Rational Design of Alkylene-Linked Bis-Pyridiniumaldoximes as Improved Acetylcholinesterase Reactivators, Yuan-Ping Pang, Thomas M. Kollmeyer, Feng Hong, Jong-Cheol Lee, Pamela I. Hammond, Sharie P. Haugabouk, and Stephen Brimijoin, *Chem. Biol.* 10, 491-502 (2003)

Improved Loading and cleavage Methods for Solid-Phase Synthesis Using Chlorotriyl Resins: Synthesis and Testing of a Library of 144 Discrete Chemicals as Potential Farnesyltransferase Inhibitors, Jewn Giew Park, Kevin J. Langenwalter, Carolyn Weinbaum, Patrick J. Casey, and Yuan-Ping Pang, *J. Comb. Chem.* 6, 407-413 (2004)

Nonbonded Bivalence Approach to Cell-Permeable Molecules that Target Specific DNA Sequences, Yuan-Ping Pang, *Bioorg. Med. Chem.*, 12, 3063-3068 (2004)

Three-Dimensional Model of a Substrate-Bound SARS Chymotrypsin-Like Cysteine Proteinase Predicted by Multiple Molecular Dynamics Simulations: Catalytic Efficiency Regulated by Substrate Binding. Yuan-Ping Pang, *Proteins*, 57, 747-757 (2004)

Lowest Level of Theory Required for Reliable Prediction of the 3D Structures of Amide-Containing Molecules Used in Virtual Screening for Drug Leads, James D. Xidos and Yuan-Ping Pang, submitted

Three-Dimensional Model of the Zinc Endopeptidase of Botulinum Neurotoxin Serotype A Complexed with a P4-P3' Substrate Fragment Predicted by Multiple Molecular Dynamics Simulation Docking. Yuan-Ping Pang, submitted

Lowest Level of Theory Required for Reliable Prediction of the 3D Structures of Amide-Containing Molecules Used in Virtual Screening for Drug Leads. James D. Xidos and Yuan-Ping Pang, submitted

EUSCAN: A Computer Program for Determination of Molecular Flexibility of Drug-Like Chemicals with Consideration of Chemical Equivalence. James D. Xidos and Yuan-Ping Pang, in preparation

US Patent Application: Novel Antiviral Drugs: Small-Molecule Inhibitors of the NP3 Serine Protease of Dengue 2 Virus, Stanley Watowich, Andrew Russo, Robert Malmstrom, and Yuan-Ping Pang

4. Wherever possible include information that is relevant to the Department of Defense and biological warfare defense.

We believe that the technology to be developed in this project should yield effective countermeasures to both biological and chemical weapons, specifically effective countermeasures to anthrax, botulinum toxin, SARS coronavirus and West Nile Virus.

Transition Activities:

Commercial product development:

Not yet

DoD/Government product development:

The technologies for developing anthrax countermeasures have been transitioned to NIH/NIAID and ARO. We have received the funding from NIH/NIAID and ARO to further the development of anthrax countermeasures.

The technologies for developing botulinum countermeasures have been transitioned to USAMRIID. We have received the funding from USAMRMC to further the development of botulinum countermeasures.

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