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Proteomic Analysis of the Blood of *Alligator mississippiensis*

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PROTEOMIC ANALYSIS OF THE BLOOD OF *ALLIGATOR MISSISSIPPIENSIS*

A Dissertation

Submitted to the Graduate Faculty of the
Louisianan State University and
Agricultural and Mechanical College
in partial fulfillment of the
requirements for the degree of
Doctor of Philosophy

in

The Department of Chemistry

by

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B.S., University of Nebraska at Kearney, 2003
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December 2011

DEDICATION

This dissertation is dedicated to my grandmothers Fairleen Penn Lightbourne-Smith and Victoria Darville, who I wished could have been here to share this special moment that they had so proudly anticipated. I love you and you both will always be an inspiration.

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First and foremost, I thank GOD the Almighty, who has given me the strength to persevere and see another goal accomplished. This dissertation would not have been possible without the love, support and encouragement of many people.

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LIST OF ABBREVIATIONS

Abbreviation	Name
2D-GE	Two-Dimensional Gel Electrophoresis
2D-PAGE	Two-Dimensional Polyacrylamide Gel Electrophoresis
ACN	Acetonitrile
AFM	Atomic Force Microscopy
AMP	Antimicrobial Peptides
ATCC	American Type Culture Collection
BLAST	Basic Local Alignment Search Tool
BSA	Bovine Serum Albumin
CCA	α -Cyano-4-Hydroxycinnamic Acid
CCD	Charge Coupled Device
CHAPS	3-[(3-cholamidopropyl) dimethylammonio]-1 propanesulfonate
CID	Collision Induced Dissociation
CNBr	Cyanogen Bromide
CRD	Carbohydrate-Recognition Domain
DHB	2,5-Dihydroxybenzoic Acid
DTT	Dithiothreitol
ECD	Electron-Capture Dissociation
EDTA	Ethylenediaminetetraacetate
Endo F	Endo- β -N-Acetylglucosaminidases F
Endo H	Endo- β -N-Acetylglucosaminidases H
ESI	Electrospray Ionization
ESI-Q-TOF	Electrospray Ionization-Quadrupole-Time-of-Flight

ESI-TOF	Electrospray Ionization Time-of-Flight
EST	Expressed Sequence Tag
ETD	Electron-Transfer Dissociation
FA	Formic Acid
Gal-GalNAc	Galactose-N-Acetylgalactosamine
GlcNAc	N-Acetylglucosamine
HPLC	High Performance Liquid Chromatography
HSP	High-Scoring Segment Pairs
HIV-1	Human Immunodeficiency Virus Type 1
IA	Iodacetamide
ID	Inner Diameter
IDA	Information Dependent Acquisition
IEF	Isoelectric Focusing
IgG	Immunoglobulin G
IgM	Immunoglobulin M
IM	Ion Mobility
IM-MS	Ion Mobility Mass Spectrometry
IMS	Ion Mobility Spectrometry
LC	Liquid Chromatography
LC-IMS	Liquid Chromatography-Ion Mobility Spectrometry
LC MS/MS	Liquid Chromatography Tandem Mass Spectrometry
LPS	Lipopolysaccharide
Man	Mannose
MALDI	Matrix Assisted Laser Desorption Ionization

MALDI-TOF-TOF	Matrix-Assisted Laser Desorption Ionization Tandem Time-of-Flight
MBL	Mannose-Binding Lectin
MCP	Microchannel Plate
MIC	Minimal Inhibitory Concentration
MS	Mass Spectrometry
MS/MS	Tandem Mass Spectrometry
MSDB	Mass Spectrometry Protein Sequence Data Base
NCBI	National Center for Biotechnology Information
NMR	Nuclear Magnetic Resonance
OMSSA	Open Mass Spectrometry Search Algorithm
PAGE	Polyacrylamide Gel Electrophoresis
PAS	Periodic Schiff
PDA	Photodiode Array
pI	Isoelectric Point
PMF	Peptide Mass Fingerprinting
PNGaseF	Peptide-N-(N-acetyl- β -glucosaminyl) asparagine amidase
Q/IMS/oa-TOF	Quadrupole/Ion Mobility Separator/Orthogonal Acceleration Time-of-Flight
QTOF	Quadrupole-Time-of-Flight
RPLC	Reversed-Phase Liquid Chromatography
RP-HPLC	Reversed-Phase High-Performance Liquid Chromatography
SA	Sinapinic Acid or 3,5-dimethoxy-4-hydroxycinnamic acid
SCX	Strong Cation-Exchange Chromatography
sDHB	2-hydroxy-5-methoxybenzoic acid

SDS	Sodium Dodecyl Sulfate
SEC	Size-Exclusion Chromatography
TC	T-cell
TCA	Trichloroacetic Acid
TFA	Trifluoroacetic Acid
TFMS	Trifluoromethanesulfonic Acid
TH	T Helper Cell
TOF/TOF	Tandem Time-of-Flight
TSB	Trypticase Soy Broth
TWIMS	Travelling Wave Ion Mobility Spectrometry
UPLC	Ultra Performance Liquid Chromatography

ABSTRACT

My research focus was to investigate alligator blood using mass spectrometry-based proteomics methods to understand their innate immune systems. The first goal was to sequence peptides and proteins from the blood serum and leukocytes using tandem mass spectrometry and *de novo* sequencing. The second goal was to determine the function of these biological molecules and their relationship to the immune system.

One- and two-dimensional gel electrophoresis was used to separate proteins from alligator leukocytes, which were enzymatically digested. The peptides were measured using reversed phase nano-high performance liquid chromatography coupled with tandem mass spectrometry (nano-HPLC-MS/MS) followed with *de novo* sequencing. The results, as described in Chapter 3 show that alligator leukocytes contain proteins that are similar to proteins found in other vertebrates such as mammals and reptiles that are related to immune responses. Isolation of small molecule interferences and peptides exhibiting antimicrobial activity from alligator leukocyte extracts are described in Chapters 4 and 5. Reversed-phase HPLC was used to separate the leukocyte mixture and antimicrobial activity tests were used to determine the active fractions. Interferants, EDTA and spermine were present and showed activity in early fractions. Two major peptides measured at 4.7 and 4.9 kDa in an active fraction were further separated on the basis of their charge, size and shape using ion mobility-mass spectrometry (IM-MS). Due to the limited fragmentation of the peptides using IM-MS, the peptides were isolated and fragmented using MALDI TOF/TOF MS for *de novo* sequencing.

Lectins are a class of carbohydrate selective proteins that are part of the complement immune system. Chapter 6 presents results for a lectin isolated from alligators that have mannan and mannose binding activity. In this study, the monomeric lectin was isolated and enzymatically digested using five different proteases to create small and large peptides which were analyzed by

LC-MS/MS. The peptides were determined via *de novo* sequencing and overlapped to generate the lectin sequence. Lectins may have varying degrees of glycosylation, therefore deglycosylation procedures suitable for mass spectrometry analysis are described in Chapter 7. Conclusions and future directions for the work in this dissertation will be summarized in Chapter 8.

CHAPTER 1. INTRODUCTION

Alligators exhibit strong immune responses when exposed to microorganisms, and are highly resistant to microbial infections.¹⁻³ Therefore, characterization of biological components that make up the immune system of alligators is of interest because these compounds may have therapeutic applications. Antibiotic resistance has become a global public health issue and is increasing at a rate that exceeds the pace of the development of new drugs. Hence, isolation of antimicrobial peptides from alligators may lead to a new class of antibiotics.

Another class of molecules that are part of the immune system is the set of proteins known as lectins, which show promise for their application in cancer therapy, pharmacology and immunology.⁵ The study of lectins isolated from *Alligator mississippiensis*, can provide insight on its structure-function relationship within the crocodile's immune system.

The goal of this research was to use mass spectrometry-based proteomics to isolate and characterize biological molecules in alligator blood and identify specific components that may have medicinal use.

1.1 Biochemistry of the Immune System

The vertebrate immune system is composed of a network of circulating cells and molecules⁷ that recognize and respond to the invasion of foreign pathogens such as viruses, bacteria, fungi and protists.⁸ These molecules include antimicrobial molecules such as lysozymes and defensins, molecules involved in specific recognition of foreign antigens such as immunoglobulins and T-cell receptors, molecules that carry intracellular signals such as cytokines, and molecules that receive and transduce these signals such as cytokine receptors.⁸ Vertebrates have both innate and adaptive immune systems that work together.⁷ The innate immune system responds rapidly and is the first line of defense against pathogens and does not

require previous exposure to exhibit a full immunological response.⁷ In contrast to the innate immune system, the adaptive immune system requires previous exposure to an antigen to exhibit a full immunological response and can take several days to weeks to become completely active.⁷

Most studies on vertebrate immune systems have been performed on mammals⁸ and the evolutionary development of the immune systems of major group of vertebrates, including mammals, birds, amphibians and fish is well known compared to other groups of organisms.⁸ In contrast, little is known about the reptilian immune systems.⁸ A better understanding of reptilian immunology will provide important information on the evolution of both innate and adaptive immune mechanisms. Crocodilians are of interest because they are the only living reptilian archosaurs, which include the dinosaurs, pterosaurs, and crocodilians.⁹

Crocodilians are vertebrates with a complex immune system. The components that make up the reptilian immune system are antimicrobial peptides, macrophages, heterophils, neutrophils, basophils, eosinophils, phagocytic B cells, and proteins of the complement system. Like other vertebrates, crocodilians also have an innate and adaptive immune system.^{7, 10} Crocodilians thrive in microbe containing environments but exhibit a strong resistance to infections.

Alligator serum has been shown to have antibacterial,³ antiviral,² and antiamoebacidal properties.¹¹ Antibacterial activity has also been observed in crocodile serum, particularly *Crocodylus siamensis*.^{12, 13} In addition, the leukocytes of *Alligator mississippiensis* have also shown to produce a broad antimicrobial spectrum.¹ The alligator complement system, which is part of the innate immune system, has also been shown to be effective against gram-positive bacteria.^{3, 14} When alligator serum was compared to human serum, the alligator serum was effective against different strains of Gram-positive bacteria, unlike human serum which had no antibacterial activity. It was proposed that the complement is also responsible for antiviral

activity from the alligator serum. Human T-cells were infected with human immunodeficiency virus type 1 (HIV-1) and when incubated with alligator serum potent antiviral activity was observed.² These studies suggest that crocodylians have a strong innate immune system.

1.1.1 Innate Immune System

The innate immune system comprises different molecules and cells including lysozymes, proteins of the complement system, non-specific leukocytes, and antimicrobial peptides.⁷ Lysozymes are enzymes that cause bacteria cells to be lysed by the hydrolysis of their cell wall.¹⁵ Lysozymes have been isolated from several reptilian organisms such as, lizards,¹⁶ turtles,¹⁶ crocodiles,¹⁷ and alligators⁶. The complement system consists of various proteins found in plasma that kill bacteria via lysis or opsonization.⁷ Lysis entails the complement proteins rupturing the bacterial membrane therefore killing the invading bacteria.¹⁸ Opsonization is the process by which opsonin proteins found in blood serum, bind to the bacterial membrane allowing the bacteria to be recognized by macrophages. The macrophages then engulf the bacteria through phagocytosis.⁷

There are three different pathways to the complement immune system: classical, alternative, and lectin.¹⁹ The classical pathway is activated by the immunoglobulins, immunoglobulin G (IgG) and immunoglobulin M (IgM) to activate an immune response.^{7,19} The alternative pathway does not require antibodies but is activated by molecules such as viruses or lipopolysaccharide (LPS) that are found on the surface of bacteria.¹⁸ Finally, the lectin pathway is activated by mannose sugars of proteins that are on the surface of bacteria.¹⁹ The complement immune system has been characterized in the American alligator¹⁴ and is believed to be responsible for antiviral activity exhibited by alligator serum.²

There are non-specific leukocytes in reptiles including eosinophils, herterophils, basophils, monocytes, and macrophages.⁷ Limited information is known about the function of

eosinophils in reptiles; however, in mammals they play a key role in the defense against parasitic infections.¹⁹ Heterophils are involved in the inflammatory response in reptiles and are also responsible for suppressing microbial invasion.²⁰ Basophils contain immunoglobulins on their surface and when triggered by an antigen releases histamine.²¹ Monocytes and macrophages are phagocytic cells that are responsible for processing and releasing antigens as well as releasing cytokines.¹⁹ Cytokines are regulatory proteins that are released by the cells of the immune system to generate an immune response.²²

An important component of the innate immune system is antimicrobial peptides and proteins. Antimicrobial peptides are found in the host defense system and exhibit antimicrobial activity.²³ Antimicrobial molecules are typically amphipathic and cationic small peptides less than 10 kDa in mass.²⁴ However, they can also be anionic peptides²⁵ or proteins.²⁶ Antimicrobial peptides can be linear and α -helical or cysteine containing and β -sheet. The linear peptides typically consist of 12–25 residues and no cysteines.²⁴ The β -sheet peptides have several antiparallel β -strands and are stabilized with up to six disulfide bonds.²⁴ There are also antimicrobial peptides that are rich in specific residues, such as tryptophan,²⁷ proline, and/or arginine^{28, 29} and histidine.³⁰

There are two major antimicrobial peptide families found in vertebrates: defensins and cathelicidins. Defensins are antimicrobial peptides that are rich in arginine residues and have a characteristic β -sheet fold and six disulfide linked cysteines.²³ They are cationic peptides that bind to microbes via electrostatic interactions.³¹ Defensins have been found in mammals and birds³² and recently the first reptilian defensin was discovered in the European pond turtle *Emys orbicularis*.³³ Defensins are 38-42 residues and found in cells and tissues involved in the host defense system. In many animals the highest concentration of defensins are found in the

granules, where the leukocytes are stored.²³ Defensins exhibit antibacterial, antifungal, and antiviral activity.²³

Another family of antimicrobial peptides is the cathelicidins. Cathelicidins are linear molecules that range in size from 12 to 80 amino acids and, unlike the β -defensins, they lack disulfide bridges.³⁴ Cathelicidins have antifungal³⁵ and antibacterial activity.³⁶ They have been isolated from fish,³⁷ birds,³⁸ mammals,³⁹ and reptiles.⁴⁰ Cathelicidins are produced in the myeloid cells in the bone marrow and stored in the neutrophil granules.⁴¹ Cathelicidins have also been found in monocytes, epithelial cells of the skin, respiratory tract, urogenital tract, as well as T and B lymphocytes.⁴¹

The membranes of microbes and multicellular animals differ in that microbes, particularly bacteria, have an outer membrane surface that is composed of lipids with negatively charged phospholipid head groups. The outer membranes of plants and animals are composed of lipids that do not have a net negative charge (Figure 1-1); the negatively charged head groups are positioned towards the cytoplasm.^{42, 43}

The primary model that explains the mechanism of antimicrobial peptides is the Shai-Matsuzaki-Huang model.^{43, 44} The Shai-Matsuzaki-Huang Model postulates that the peptide interacts with the pathogen membrane, displacing the lipids and disrupting the membrane structure. In some cases the peptide may also penetrate the cell.⁴⁴ The membrane of multicellular animal cells contains cholesterol, which reduces the activity of the antimicrobial peptide via peptide interaction with the cholesterol or stabilization of the lipid bilayer. In addition to this proposed mechanism, studies suggest that there are other mechanisms involved. These mechanisms include 1) depolarization of the bacterial membrane leading to death,⁴⁵ 2) creation of holes in the cell causing cellular leakage,⁴⁶ 3) activation of processes that cause cell death (e.g. hydrolases which destroys the cell wall),⁴⁷ 4) disruption of the membrane function by

rearrangement of the lipids on the outer cell membrane,⁴³ and 5) destruction of the internal cellular components after internalization of the antimicrobial peptide.⁴⁸

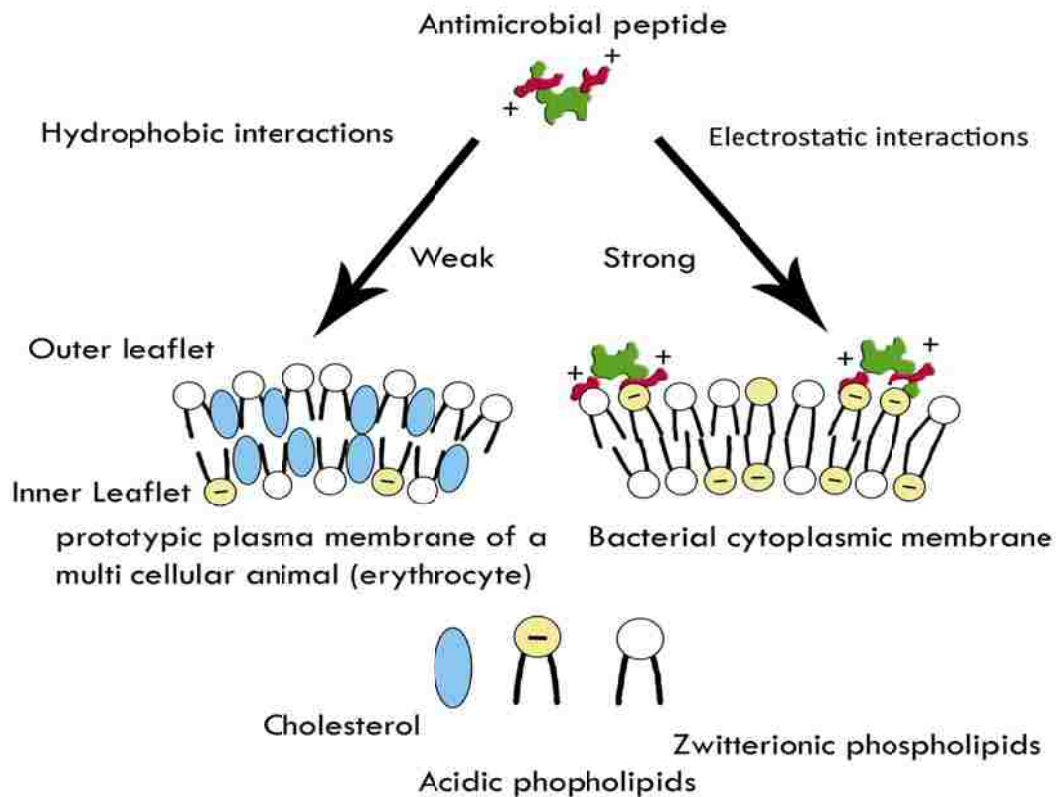


Figure 1-1. Comparison of antimicrobial peptide interaction with animal and bacterial membranes.

Some antimicrobial peptides have been tested for pharmaceutical use, primarily as a topical treatment. The first antimicrobial peptide to undergo clinical trial was the magainin peptide isolated from frog skin.⁴⁹ The magainin peptide was developed as a topical treatment and progressed to Phase III clinical trials to treat diabetic patients suffering from foot ulcers.^{49,50} The study showed the magainin peptide to be equally as effective when compared to the oral antibiotic ofloxacin.⁵¹ A variety of antimicrobial peptides are currently being developed as potential antibiotics as indicated in Table 1-1.

The broad antimicrobial spectrum of the antimicrobial peptides allows them to be used for diverse applications. However, a major drawback of antimicrobial peptides for clinical use is the level of toxicity. The level at which the antimicrobial peptides are effective *in vivo* are usually toxic.⁵² Other factors are stability and immunogenicity.⁴⁹

Table 1-1. Antimicrobial peptides available for therapeutic use.^{49, 53}

Peptide	Pharmaceutical Name	Origin	Mode of Application	Application	Stage
Magainin 2	Pexiganan	African clawed frog skin	Topical	Foot ulcers	Completed Phase III. (Not approved by FDA)
Indolicidin	Omiganan (MBI-226)	Synthetic analog of indolicidin	Topical	Catheter infection	Phase III
Indolicidin	MBI-594AN	Cow erythrocytes	Topical	Acne	Phase III
Protegrin	Isegranin (IB-367)	Pig leukocytes	Oral	Mucositis	Phase III
Histatin	P113P113D	Human	Oral	Gingivitis	Phase II
Heliomycin	Heliomycin	Tobacco budworm	Systemic	Antifungal	Preclinical
Lactoferricin	Lactoferricin	Human	Systemic	Antibacterial	Preclinical
Bactericidal permeability increasing protein histatin	XMP.629	Human	Systemic	Meningococcal meningitis	Phase III

Another group of molecules that play an important role in vertebrate innate immune systems is lectins. Lectins have been studied for more than 100 years in plants and have recently been recognized for their importance in animals.⁵⁴ Lectins are proteins that can recognize carbohydrates that are endogenous to the animal or those on microbes.⁵⁵ They have been

classified based on their carbohydrate ligands, biological processes, subcellular location, and dependence on divalent cations.⁵⁵ They have a wide range of structures, but carbohydrate-binding activity is limited to a particular region of the protein known as the carbohydrate-recognition domain (CRD).⁵⁶

Animal lectins are grouped into two structural families: C-type and S-type.⁵⁶ C-type lectins are found in serum, extracellular matrix, and plasma membranes. They require calcium for binding and bind to a variety of sugars. S-type lectins are found intracellularly and require sulfhydryl for binding to β -galactosides.⁵⁷

Animal lectins possess a wide variety of functions such as cellular growth regulation, mediation of endocytosis, intracellular routing of glycoconjugates, and urate transport.⁵⁸ However, the major function of lectins is to behave as recognition molecules within the immune system.⁵⁸ These roles within the immune system include: direct defense, cell recognition and trafficking, immune regulation, and prevention of autoimmunity.⁵⁸ Different applications of lectins have been demonstrated such as immobilization onto columns for affinity separation of glycoproteins, glycopeptides and oligosaccharides,⁵⁹ selective agglutination,⁶⁰ and blood typing.⁶¹ Lectins show promise for their application in agriculture, cancer therapy, pharmacology, and immunology.⁵

1.1.2 Adaptive Immune System

The adaptive immune system becomes activated following the innate immune response. There are two adaptive immunity responses: cell-mediated and humoral adaptive immunity.⁷ Cell-mediated immunity involves T-cells that are responsible for regulating antibody production. T-cells can differentiate into two types of cells: cytotoxic T-cell (TC) or T helper cell (TH). TC can rapidly kill bacterial or viral infected cells through apoptosis. TC regulates other immune cells. TH helps control other immune cells. T-cells also release cytokines that affect the humoral

response (the immune responses mediated by antibodies).⁷ Humoral adaptive immunity relies on B-cells that recognize antigens even before being processed.¹⁹

1.1.3 Study of the Immune System

To study the immune system of vertebrates, serological assays are typically used, in which *in vitro* reactions between antigen and serum antibodies are studied.⁶² There are many different types of serological assays including agglutination, precipitation, immunoassays, immunofluorescence, fluorescence-activated cell sorting analysis, and lymphocyte function.⁶² These assays help elucidate the immune system's functional and regulatory properties through lymphocyte function measurements and the responses of B- and T-cells as well as antibodies.

Another approach to studying the immune system is hematology, which is the study of blood, including white blood cells, red blood cells, hemoglobin, and platelets.⁶² Hematological tests allow for observation of live blood (whole blood that is unaltered and unstained) with microscopy. Using this test, microbial activity in the blood and its potential effects can be measured. In addition, the white blood cells can be quantified, which can provide insight on how the immune system is functioning. There is also the standard blood microscopy technique in which the blood is stained and fixed; however, staining kills the blood cells.

Another technique for studying the immune system is genomics and proteomics, which is discussed in Section 1.2. Genomics and proteomics provide a more comprehensive view of the immune system and its function as compared to serological assays and hematology tests.

1.2 Chemical Analysis Methods for Proteomics

Proteomics is the approach used to understand complex biological systems by analyzing protein expression, function, modifications, and interactions.⁶³ Proteomics is related to genomics, which is the study of the genetic make-up of an organism.⁶⁴ An important step towards understanding an organism's biology is to determine its genome sequence. However, the genome

sequence is not enough to provide information on complex cellular processes; the complement of proteins associated with a particular genome is essential to this understanding.⁶⁵

Proteomics is complimentary to genomics and provides an additional component to the understanding of biological systems. However, there are significant challenges in proteomics such as limited sample quantity, sample degradation, broad dynamic range ($>10^6$ fold for protein abundance), post-translational modifications, and disease changes.⁶⁶ Protein concentrations typically exceed the dynamic range of a single analytical instrument or method necessitating the use of one or more dimensions of separation.⁶³

Proteomics can be divided into three major branches: structural proteomics,⁶⁷ expression proteomics,⁶⁸ and functional proteomics.⁶⁸ Structural proteomics involves determining the structures of proteins, such as its three-dimensional shape (secondary and tertiary structure) and its amino acid sequence (primary structure).⁶⁷ A commonly used approach for protein sequencing is Edman degradation. Edman degradation was developed by Pehr Edman in 1950 and is one of the oldest and most developed techniques for protein sequencing.⁶⁹ However, Edman sequencing chemistry has largely been replaced by mass spectrometry for protein sequencing and identification.^{70, 71} Protein secondary and tertiary structures can be characterized by X-ray crystallography and nuclear magnetic resonance (NMR) spectroscopy.⁶⁷

Expression proteomics involves the quantitative and qualitative analysis of proteins under different conditions.⁶⁸ This approach allows disease-specific proteins to be identified by comparing the entire proteome between two samples. Proteins that are over-expressed or under-expressed can be identified and characterized. The techniques commonly used for expression proteomics are two-dimensional gel electrophoresis,⁷² multi-dimensional chromatography with mass spectrometry,⁷³ and micro-array techniques.⁷⁴ Two-dimensional electrophoresis suffers from limitations such as the large dynamic range of protein expression in biological systems and

difficulty in analyzing proteins that are post-translationally modified.⁶⁸ Limitations of microarray technology include the sensitivity of the arrays for detection of low abundance genes⁷⁵ as well as its inability to measure post-translational modifications.⁷⁶ Unlike 2-D electrophoresis and microarray techniques, multi-dimensional chromatography with mass spectrometry can measure post-translational modifications in expression proteomics.⁷⁷

Functional proteomics is an approach to analyze and understand the properties of macromolecular networks involved in cells.⁶⁸ Proteins and their specific roles in metabolic activities can be identified.

1.2.1 Sample Purification

Blood plasma and serum are commonly used biological fluids for proteomic analysis because many cells release a portion of their content into the plasma when damaged or upon cell death. There are approximately 10,000 proteins present in human serum⁷⁸ and many proteins of interest are present in low abundance. Plasma comprises ~97% high abundance proteins including albumin (57–71%) and immunoglobulins (8–26%).⁷⁹ Hence plasma samples are difficult to analyze directly and purification is required. In addition, proteomic samples usually contain a large amount of contaminants such as lipids, nucleic acids, and surfactants.⁸⁰ The dynamic range of expressed proteins is greater than six orders of magnitude⁸¹ and protein mixtures can be quite complex and contain proteins with different solubility, hydrophobicity and hydrophilicity, pI, and molecular masses.⁸² Therefore, sample purification is necessary before mass spectrometry analysis.

Separation methods such as affinity-based techniques, chromatography and centrifugation have been employed.⁸³ It is important that the proteins of interest are well resolved with limited sample purification steps to avoid sample loss. Affinity-based techniques have been established for removal of albumin and IgG using immobilized antibodies that are selective

against albumin⁸⁴ and protein A or protein G that selectively capture IgG on the columns.⁸⁵ The removal of salts and other contaminants from protein samples has been accomplished using precipitation with centrifugation.⁸⁰ Commonly used precipitation methods include acetone, trichloroacetic acid (TCA), ammonium sulfate, and chloroform/methanol precipitation.

1.2.2 Digestion

Protein digestion can be performed using three common approaches: in-gel,^{86, 87} in-solution,⁸⁸ or solid phase.^{89, 90} In an in-gel digestion, the proteins are separated on a 1- or 2-D gel and the gel bands are excised for chemical or proteolytic digestion.⁸⁷ A major advantage of in-gel digestion is that it removes detergents and salts that can be interferences in the mass spectrometer.⁸⁷ However, a limitation to this method is the loss of peptides during in-gel digestion through binding to the polyacrylamide.⁸⁷ Another approach that has been developed is in-solution digestion which entails digesting proteins directly in buffers or solvents such as ammonium bicarbonate or acetonitrile.⁹¹ This approach is advantageous in that low abundance molecules that may otherwise be lost in the gel can be detected. A drawback for this method includes longer incubation times due to lower enzymatic concentrations.⁹² Solid phase is another digestion approach which includes immobilization of an endoprotease on a solid support; examples include monolithic columns for trypsin digestion⁹³ and microfluidic devices with integrated trypsin digestion.⁹⁰ Solid phase digestion offers advantages of increased digestion rates, reduced interferences from trypsin autolysis products, low sample consumption, and fast response.^{94, 95} A limitation to this method is the use of organic solvents which improve digestion efficiency but can damage the immobilized enzyme.⁹⁶

Digestion efficiency can be improved by cysteine reduction before digestion. The disulfide bonds are reduced with a reagent such as dithiothreitol (DTT) and an alkylation reaction is performed with iodoacetamide to prevent new disulfide bridges from forming.⁹⁷

Many of the endoproteases and chemicals cleave proteins at specific amino acids generating peptide fragments of varying lengths. Peptide fragments between 6–20 amino acids are best for MS analysis and protein database searching.⁹⁸ Endoproteases and chemicals used for protein analysis are indicated in Table 1-2. The most commonly used endoprotease for proteomic analysis is trypsin. Trypsin cleaves at lysine and arginine residues, unless followed by a proline residue in the C-terminus direction.⁹⁹ Trypsin has good activity in both in-gel and in-solution digests.

Table 1-2. Chemicals and proteases used for enzymatic and chemical cleavage. Cleavage with the endoproteases only occurs if the residue after the cleavage site is not proline, except for Asp-N.¹⁰⁰

Endoproteases	Cleavage Specificity
Trypsin	K, R
Glu-C	E, D
Lys-C	K
Asp-N	D
Arg-C	R
Chymotrypsin	W, Y, F, L, M
Chemical Agents	Cleavage Specificity
70% Formic acid	D
Cyanogen bromide	M
2-nitro-5-thiocyanobenzoate, pH 9 ¹⁰¹	C
Hydroxylamine, pH 9 ¹⁰²	N, G
Iodobenzoic acid	W

Many proteins contain a significant number of lysine and arginine residues that are spaced sufficiently in the sequence so that trypsin produces fragments that are a suitable length for MS analysis. Another complementary enzyme used is Glu-C which cleaves at the carboxyl

side of glutamate residues.^{103, 104} In the presence of selected buffers such as sodium phosphate, it can cleave at both the glutamate and aspartate residues. Other proteases listed in Table 2 with cleavage specificities are useful for producing peptides of varying lengths depending on how many cleavages occur. This is useful for obtaining additional sequence information, especially if the protein is unknown.

There are also non-specific endoproteases such as pepsin as well as endoproteases with broad specificities such as chymotrypsin that are useful for producing multiple overlapping peptides that can increase sequence coverage.⁹⁸ Proteins can also be cleaved with cyanogen bromide, formic acid, and hydroxylamine. Cyanogen bromide is the most commonly used for protein cleavage; it cleaves specifically at methionine residues.¹⁰⁵

1.2.3 Separations

Two separation methods used in proteomics are gel electrophoresis and liquid chromatography (LC). Gel electrophoresis is a sensitive method for separating and identifying proteins in a gel matrix such as agarose or polyacrylamide.⁸⁰ Agarose is typically used to separate larger macromolecules such as nucleic acids and polyacrylamide is typically used to separate proteins. Polyacrylamide gel electrophoresis can be used to determine the size, isoelectric point and purity of proteins.⁸⁰ The gel pores are made by crosslinking of the polyacrylamide with bis-acrylamide to form a network of pores that allows the molecules to move through the gel matrix like a sieve. The gel pore size is determined by the acrylamide monomer concentration.¹⁰⁶

Gel electrophoresis separates molecules based on the differences in migration velocity of ions in the gel under the influence of an electric field. The migration velocity is the product of the electrophoretic mobility and the applied electric field. The electrophoretic mobility is proportional to the ion charge and inversely proportional to the frictional forces. The frictional

forces depend on the analyte size and the viscosity of the solvent. Smaller analytes have a greater mobility and migrate farther down the medium in a given time. Polyacrylamide gel electrophoresis (PAGE) is used to separate proteins and peptides based on their size. Sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE) is the most commonly used gel based technique for separating proteins. SDS is used to denature the proteins and gives the protein an overall net negative charge.¹⁰⁷

Two-dimensional gel electrophoresis (2D-GE) is used to separate proteins based on their charge and mass.^{107, 108} The first dimension separates proteins based on their net charge using isoelectric focusing.¹⁰⁸ Protein separation is performed in a pH gradient; the proteins migrate to their isoelectric point (pI), which is the pH where the protein has a net charge of zero. A protein's pI is determined by the type and number of acidic and basic residues it contains.^{107, 108} The second dimension in 2D-GE separates proteins based on their mass and is usually performed in a SDS gel (SDS-PAGE). 2D-GE has difficulty resolving large proteins, or those with extreme pI or hydrophobicity, and suffers from lack of reproducibility.¹⁰⁹

Liquid chromatography (LC) is a technique used to separate components on a stationary phase using a liquid mobile phase. Reversed-phase high-performance liquid chromatography (RP-HPLC) separates proteins and peptides by hydrophobicity.¹¹⁰ It is one of the most powerful and commonly used liquid chromatography techniques.¹¹¹ Commonly used hydrocarbon ligands for reversed-phase resins include C₄ and C₁₈.¹¹⁰ C₄ is commonly used for polar proteins and C₁₈ is most used for peptides.

Ultra performance liquid chromatography (UPLC) uses smaller particles as well as high speed and peak capacity (the number of peaks that can be resolved per unit time).¹¹² Compared to conventional HPLC columns which are packed with 3.5 to 5 μm particles, UPLC columns are

packed with 1.7 μm particles.¹¹³ Smaller particles shorten the analyte's diffusion path which improves separation efficiency, speed, and resolution.^{114, 115}

Ion exchange is another form of chromatography which involves the separation of proteins and peptides based on their charge.¹¹⁰ A cationic or anionic resin is used and proteins or peptides of opposite charge are retained due to charge attraction. Hydrophilic-interaction chromatography separates proteins based on their hydrophilic properties; hence the stationary phase is polar.¹¹⁶ Another separation technique is affinity chromatography, which separates proteins and peptides based on their specific ligand-binding affinity.¹¹⁰ There are two fractions collected from affinity separation, the unbound and the bound proteins and peptides. The analyses and detection of low abundant proteins, primarily in plasma, can be difficult due to the presence of high abundant proteins such as albumin, immunoglobulins, and transferrin.¹¹⁷ Therefore, affinity-based approaches can be used to remove high abundant proteins or low abundant proteins can be enriched.¹⁰⁹

Due to the complexity of the protein samples, one-dimensional separation techniques are usually insufficient and multi-dimensional separations are employed. In multi-dimensional separation, two or more separation techniques are coupled together to improve the resolving power. These separation techniques are orthogonal to one another.

1.2.4 Mass Spectrometry

Mass spectrometry (MS) is an analytical technique used for measuring the mass and chemical structure of molecules and is widely used for proteome analysis. A variety of ionization techniques can be used for mass spectrometry but the most commonly used techniques for the analysis of biomolecules are electrospray ionization (ESI)¹¹⁸ and matrix assisted laser desorption ionization (MALDI).¹¹⁹⁻¹²¹ MALDI uses a matrix that absorbs laser energy and aids in ionization of the analyte. The ions generated are typically singly charged. ESI can also be used to analyze

biomolecules. Unlike MALDI, its ions are produced from solution. After the ions are formed they are transferred into a mass analyzer by an electric field where they are separated according to their mass-to-charge ratio. Two stages of mass separation can be coupled (either in space or in time) to obtain additional information of the sample being analyzed which is known as tandem mass spectrometry (MS/MS).¹²² Tandem mass spectrometry is used to determine peptide sequences from proteins.⁸³ A peptide is separated from a mixture of peptides in the first stage of mass spectrometry and dissociated by collision with an inert gas. The generated fragments are separated in the second stage of mass spectrometry.^{83, 123}

There are three MS proteome analysis approaches: 1) bottom-up proteomics, 2) shotgun proteomics, and 3) top-down proteomics. In the bottom-up approach the protein mixture is separated usually by 1 or 2-dimensional electrophoresis and the individual protein bands or spots are cut and digested with an enzyme such as trypsin resulting in peptides. The peptides are analyzed by mass spectrometry using peptide mass fingerprinting or tandem mass spectrometry (MS/MS) to create sequence tags for database searching.¹²⁴⁻¹²⁶ Some of the major advantages of using the bottom-up approach are the ability to obtain high-resolution separations and a comprehensive coverage of proteins. It's the most widely used technique in proteomics,¹²⁷ hence several bioinformatics tools are available. In addition, proteins can be separated in a complex mixture before digestion so there is a greater chance of identification. The drawback of this approach is the limited dynamic range¹²⁸ and difficulty separating membrane proteins.^{129, 130}

In shotgun proteomics, a mixture of intact proteins is enzymatically digested and separated using strong cation-exchange chromatography (SCX) followed by reversed-phase liquid chromatography (RPLC).^{109, 131} The separated peptides are subjected to tandem mass spectrometry and database searching.¹³¹ A major advantage of this technique is that thousands of proteins can be identified in a single analysis and the technique is better suited to membrane

proteins. However, limitations of this technique includes the need for complex mixtures to be purified prior to separation,¹³² limited dynamic range,⁶³ and bioinformatics challenges to identify peptide and protein sequences from a large number of acquired spectra.⁶³

In the top-down approach, intact proteins are separated by gel electrophoresis or HPLC before being introduced into the mass spectrometer.^{124, 126} The mass of the protein is measured and tandem mass spectrometry is used to generate sequence tags (a short sub-sequence of a peptide sequence) for database searching. Alternately, *de novo* sequencing (an approach to determining a peptide sequence without prior knowledge of the sequence) can be performed.¹²⁶ Top-down sequencing can be used to locate and characterize post-translational modifications, determine the complete protein sequence, and minimize time-consuming preparation steps such as separation and digestion procedures. Conversely, spectra generated by multiply charged proteins can be very complex and bioinformatics tools still needs more developing for protein identification. In addition, top-down sequencing does not work well with intact proteins greater than 50 kDa.

Separation can also be performed in the gas phase using ion mobility mass spectrometry (IM-MS).¹³³⁻¹³⁵ Ion mobility is a gas-phase technique that separates ions based on their ability to migrate through a gas in the presence of an electric field.^{133, 134} The mobility is dependent on the collision cross section of the ion. An ion mobility spectrometer consists of a gas filled cell where ions travel under the influence of an electric field.¹³³ Ions with larger diameters undergo more collisions with the buffer gas, hence their passage through the drift cell is slower, whereas smaller molecules undergo fewer collisions and pass through the drift cell more rapidly.¹³³ When coupled with MS both the mass to charge and size to charge ratio of the ions can be determined. Both MALDI and ESI can be used as an ionization source for IM-MS.¹³⁶ However, many IM studies reported in literature of peptides and proteins describe the use of an ESI source coupled

with IM.¹³⁷⁻¹³⁹ IM can also be coupled with tandem MS for additional peptide and protein information.¹⁴⁰

IM-MS separations can obtain comparable resolutions to HPLC and CE.¹³⁴ LC-MS, a more commonly used proteomics technique is advantageous in that it increases the dynamic range,¹⁴¹ however, coupling LC to MS limits optimization of LC and LC can take minutes to hours for separation which limits sample throughput.¹⁴² IM offers two major advantages over LC techniques; it reduces separation time and the post-ionization separation provides information on the product.¹³⁴ Another major advantage of IM-MS is that it adds two separation dimensions for different chemical classes,^{143, 144} for example peptides, DNA, oligonucleotide and lipids or conformational classes such as α -helix and random coil.¹⁴⁵ However, IM-MS has its shortcomings for proteomic applications, specifically poor sensitivity and limited peak capacity.¹³⁴ Despite the limitations of IM-MS for proteomics it has still been proven to be a useful method for proteomics,¹³⁴ metabolomics,¹⁴⁶ and glycomics.¹⁴⁷

1.2.5 Protein Bioinformatics

Bioinformatics is the approach used to analyze large numbers of genes and proteins¹⁴⁸ and is important for the elaboration of mass spectrometry data due to the large amount of data produced.¹⁴⁹ It is used in proteomics to provide functional analysis and mining of data sets.¹⁴⁸ Peptide and protein data can be interpreted via peptide mass fingerprinting, database searching, or *de novo* sequencing.

Peptide mass fingerprinting (PMF) is a high-throughput protein identification method.¹⁵⁰ In this method the protein is purified and cleaved using an enzymatic or chemical approach to generate peptides. The peptides are analyzed via ESI or MALDI mass spectrometry and a peptide mass fingerprint (the masses of the intact peptides in the sample) is obtained. The mass fingerprint is compared to theoretical cleavages of protein sequences in databases and the protein

matches are scored based on the best match.^{150, 151} Several programs have been developed for peptide mass fingerprinting including MassSearch, Mowse, MS-FIT, PepMAPPER, PepSea, PeptideSearch, ProFound and PeptIdent.¹⁵² Peptide mass fingerprinting will only yield hits for proteins that are in a sequence database.

An alternate approach to database searching is mass spectral matching. This entails matching the experimental spectrum to a library of previously obtained MS/MS data.¹⁵³ This method is a fast and precise means to identifying peptides whose proteome has been previously identified. Its major limitation is its inability to be used for identifying or discovering new peptides.

Another approach for database searching compares experimental spectra to theoretical spectra to identify peptides in the protein database. Theoretical tandem mass spectra are produced from fragmentation patterns that are known for a specific series of amino acids. Search engines that are used for database searching includes Mascot,¹⁵⁴ SEQUEST,¹⁵⁵ X!TANDEM,¹⁵⁶ Open mass spectrometry search algorithm (OMSSA),¹⁵⁷ SONAR,¹⁵⁸ ProbID,¹⁵⁹ PeptideProphet¹⁶⁰ and OLAV-PMF.¹⁶¹ Two commonly used search engines include Mascot and SEQUEST. SEQUEST uses a cross-correlation score to match hypothetical spectra to experimental spectra¹⁵⁵ whereas Mascot uses a probability score that indicates the probability of whether or not a spectral match was random.¹⁵⁴ When some commonly used search engines were compared, including SEQUEST and Mascot, Mascot proved to be able to better discriminate between a correct and incorrect hit as compared to SEQUEST.¹⁶² An overall evaluation showed that Mascot outperformed the other algorithms used in the study, which included PeptideProphet, Spectrum Mill, SONAR and X!TANDEM.

Mascot is based on probability scoring and the lowest probability is the best match. The match significance criteria depend on the size of the database. The score is reported as $-10\log(P)$, where P is the probability. Hence, the best match has the highest score.

Database searching offers several advantages including high-throughput, robustness and well annotated proteins (detailed information on each protein). Despite these advantages there are also some disadvantages that are associated with database searching including false positive identification due to selection of background peaks, unidentified peptides due to post translational modifications, scoring a longer peptide that may be from a lower quality MS/MS spectrum (low signal-to-noise ratio) with a higher score than a shorter peptide from a higher quality MS/MS spectrum (high signal-to-noise ratio), and, most importantly, it is impossible to identify a peptide that is not part of a protein in the database.¹⁶³

De Novo sequencing is an approach to identifying peptides without database searching, for example for a species whose genome has not been previously sequenced. It is also used to identify post-translational modifications. The *de novo* approach determines peptide sequences using information such as the type of fragmentation method, for example collision induced dissociation (CID),¹⁶⁴ electron-transfer dissociation (ETD),¹⁶⁵ or electron-capture dissociation (ECD),^{164, 166} the type of enzyme used, as well as any chemical modifications. Some commonly used *de novo* sequencing programs are PEAKS¹⁶⁷ Mascot Distiller⁹⁷, Lutefisk,¹⁶⁸ PepNovo,¹⁶⁹ and SHERENGA.¹⁷⁰ Tandem mass spectrometry data can also be searched against expressed sequence tag (EST) databases to identify peptides and proteins for organisms without complete genomes. ESTs are nucleotide sequences (200 to 500 nucleotides long) that are generated by sequencing either one or both ends of an expressed gene originating from specific tissues.¹⁷¹ These nucleotide sequences are translated into protein sequences for protein identification from tandem mass spectra.

Under CID, peptides fragment along the peptide backbone and fragment ions generated from the N-terminus of the peptide are labeled a, b, and c, whereas fragments generated from the C-terminus of the peptide are labeled x, y, and z.¹⁷² *De novo* spectra generated from low energy CID gives only partial peptide ion coverage because of its backbone cleavage specificity; in low energy CID spectra c, x, z and a-type fragment ions are not observed.¹⁷² Hence, it is usually beneficial to collect peptide spectra from other fragmentation methods such as ETD or ECD. Some of the limitations associated with CID include overlapping fragment ion peaks (which can cause incorrect peak assignment), low signal for some of the ions in the CID spectra, difficulty identifying post-translational modifications, and the inability to differentiate between the amino acids leucine and isoleucine.¹⁶³ The ETD and ECD techniques can be used which can differentiate between leucine and isoleucine as well as identify post-translational modifications. ECD also produces less specific backbone cleavage as compared to CID; therefore, more extensive sequence information can be obtained on proteins.¹⁶⁴ However, a limitation of ETD and ECD is their inability to produce good quality data with shorter peptides, such as those generated from tryptic digests.¹⁶³

Basic local alignment search tool (BLAST) is a search algorithm that is used to compare sequence similarities between experimentally determined nucleotide or protein sequences with nucleotide or protein databases.¹⁷³ This approach is useful for the identification of proteins from organisms that have unsequenced genomes.^{174, 175} A BLAST alignment pairs each amino acid in the queried sequence to those in another sequence from a protein database. BLAST begins a search by indexing short character strings (amino acid sequences) within the peptide query by their starting position in the query. The “word size” (length of the amino acid sequence) for a protein-to-protein sequence comparison is typically three. The BLAST software then searches the database to look for matches between the indexed “words” from the queried peptide to

character strings within the sequence in the database. Whenever a word match is found, BLAST then extends the sequence (using the database sequence) in the forward and backward direction to create an alignment. The BLAST score value increases as long as the alignment matches and will begin to decrease once it encounters mismatches.^{176, 177}

The BLAST results are quantified by comparing them to the expect value (E-value). The E-value threshold represents the number of times a good match is expected to occur by chance and is proportional to the size of the database. BLAST determined E-values that are greater than the threshold E-value are considered significant. The higher the similarity between the queried sequence and the sequence in the database the lower the E-value is. This can be seen in Equation 1-1,

$$E = K * m * n * e^{-\lambda S} \qquad \text{EQUATION 1-1}$$

where K is a constant (scaling factor), m is the length of the query sequence, n is the length of the database sequence, λ is the decay constant from the extreme value distribution (scales for the specific scoring matrix used) and S is the similarity score.¹⁷⁶

1.3 Research Objectives

The objective of this research is to characterize the biological molecules isolated from the *Alligator mississippiensis* immune system and identify antimicrobial molecules. To achieve this goal, the specific aims of the project were (1) to isolate and characterize the proteins of the leukocytes, (2) to purify and identify antimicrobial peptides from whole blood, and (3) to isolate and determine the primary sequence of lectin from the serum. The first set of experiments, described in Chapter 3, used gel electrophoresis and multidimensional gel electrophoresis to separate a mixture of isolated leukocyte proteins for mass spectrometry detection and sequence identification. A second set of experiments, described in Chapters 4 and 5, used a combination of

separation techniques for purification of a novel antimicrobial peptide. Although antimicrobial peptides have been identified and characterized in numerous organisms, antimicrobial peptides have not yet been identified in alligators. For the third set of experiments described in Chapter 6, a new lectin from *Alligator mississippiensis* was identified using *de novo* sequencing with different enzymatic digests and characterized. Chapter 7 describes deglycosylation approaches and compares sample clean-up procedures that can be applied to glycoproteins for mass spectrometry analysis.

CHAPTER 2. EXPERIMENTAL*

This chapter contains a description of the instruments and methods used for analyzing peptides and proteins isolated from the blood of alligators. A summary of the instruments and techniques used for this research, including gel-based protein separations, liquid chromatography, ion mobility, mass spectrometry, and proteomic bioinformatics is described. A detailed description of the experimental parameters and procedures used for the separation techniques as well as the electrospray ionization time-of-flight (ESI-TOF), matrix-assisted laser desorption ionization tandem time-of-flight (MALDI-TOF/TOF) and electrospray ionization-quadrupole-time-of-flight (ESI-QTOF) mass spectrometers is given.

2.1 Blood and Leukocyte Collection

Adult alligators were captured at night by boat at J. D. Murphree State Wildlife Refuge in Port Arthur, Texas, with the use of a spotlight and a cable harness and housed at the Rockefeller State Wildlife Refuge in Grand Chenier, Louisiana. The animals were kept in fiberglass-lined concrete tanks approximately 5 m long and 2 m wide, with 50% dry bottom and 50% water of approximate 0.2 m depth. The temperature was maintained at 31 °C. Several alligators up to 1.5 m in length were housed in a single tank. The alligators were fed formulated dry pellets four times per week and the cages were cleaned five times per week.

Blood samples were collected from the supravertebral branch of the internal jugular vein using a heparinized 38 mm long 18 gauge needle and a 60 mL syringe^{178, 179} and transferred to 250 mL plasma bottles containing 25 mL of 500 mM EDTA. This method of blood collection has been approved by the McNeese State University Animal Care and Use Committee and is routinely used to collect blood from wild and captive adult alligators (See Appendix D).

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Leukocytes were isolated from the whole blood by differential sedimentation. Whole blood samples were mixed with one volume of 5% dextran and the erythrocytes were allowed to settle for approximately two hours. The leukocytes were separated manually from the top phase using a transfer pipette.¹

The leukocytes were centrifuged for 15 min. at 1500 *g* and the leukocyte pellet was resuspended in approximately one volume of 10% acetic acid and stored at -80°C until needed. Prior to analysis, the leukocyte pellets were homogenized in a Dounce homogenizer and then centrifuged at 20,000 *g* for 30 min. The clear supernatant was transferred to 1 kDa concentrator tubes (Centriprep, Millipore Corp., Billerica, MA) and centrifuged at 5000 *g* to concentrate the sample and remove the acetic acid. The concentrated extract and precipitate were resuspended in 10 mL of 0.1% acetic acid. The resuspended extract was centrifuged at 20,000 *g* and the clear supernatant was used in the gel electrophoresis and HPLC analysis.

2.2 Liquid Chromatography

In this work, reversed-phase liquid chromatography was used. Both capillary and nano capillary high performance liquid chromatography (HPLC) as well as nano ultra performance liquid chromatography (UPLC) were used.

2.2.1 Nano HPLC

Nano capillary HPLC (nanoLC) uses columns that have an inner diameter (ID) between 50 and 100 μm and flow rates below approximately 120 nL/min.¹⁸⁰ In this work, nanoLC was coupled to the mass spectrometer described in Section 2.6.4. Typical columns used for nanoLC have lengths of 50-150 mm with 3-5 μm particle diameters. For improved chromatographic separation, acids, including trifluoroacetic acid (TFA) and formic acid (FA), are added to the solvents at 0.1%. TFA is best for offline LC because it provides higher peak capacity with narrower peak width. However, if the LC is coupled online with the mass spectrometer, FA is

preferred because the high surface tension of the TFA solution makes spray formation difficult and trifluoroacetate can form strong ion pair complexes with peptide cationic sites causing signal suppression.¹⁸¹

The LC separations were performed on a LC Packings/Dionex capillary and nano HPLC system. The HPLC system is equipped with a low pressure mixing system, micropump, flow splitting unit that allows flow rates from 50 nL/min to 200 μ L/min, a sample injector, and UV detector. A Vydac C₁₈ 75 μ m ID \times 150 mm, 5.0 μ m column was used. The solvents were water in 0.1% FA and acetonitrile in 0.1% FA. The column flow rate was 200 nL/min and a linear gradient was used for separation.

2.2.2 Capillary HPLC

Capillary HPLC is separation on a column that has an ID between 100 and 300 μ m and flow rate that is several μ L/min. A capillary HPLC (Agilent 1200 series) was used both online with the Agilent mass spectrometer (see Section 2.6.3 below) and offline with a photodiode array (PDA) detector over a wavelength range between 190 and 400 nm. The capillary HPLC is equipped with a binary and gradient pump, diode array detector, autosampler and thermostat. In this work, a C₁₈ 2.1 mm ID \times 50 mm, 5.0 μ m column (Zorbax) and a C₁₈ 4.6 mm ID \times 250 mm, 5.0 μ m column (Vydac) were used. The solvents were water in 0.1% FA and acetonitrile in 0.1% FA. The column flow rate was 300 μ L/min and a linear gradient was used for separation.

2.2.3 Nano Ultra Performance Liquid Chromatography

Nano ultra performance liquid chromatography (nano UPLC) uses columns that typically have an ID between 75 and 1000 μ m and flow rates between 200 nL/min and 100 μ L/min. A UPLC system (Waters nanoAcquity) was used online with the Waters Synapt instrument described in Section 2.6.5. The system is equipped with a binary solvent manager, sample manager, a column heating compartment and a UV detector. In this study, an C₁₈ column

(Acquity) packed with 1.7 μm particles, 5 cm long \times 2.1 mm ID was used. The samples were diluted in 0.1% FA and eluted using a gradient with 3–85% 0.1% FA in acetonitrile (v/v) for 32 minutes. The column flow rate was 500 nL/min and 214 nm UV absorption was recorded.

2.3 Ion Mobility

Another separation technique used in this work was ion mobility (IM). The ion mobility cell was built into a Q-TOF mass spectrometer (Waters Corporation), details of this mass spectrometer are found in Section 2.6.2. The peptide samples were diluted in 0.1% FA and injected on the UPLC system at a flow rate of 500 nL/min. The peptides were separated using a gradient of 3–40% B over 30 minutes. Solvent A was 95% water and 5% acetonitrile containing 0.1% FA. Solvent B was 80% acetonitrile and 20% water containing 0.1% FA. The peptides were ionized using a nano-electrospray source and analyzed via a quadrupole mass analyzer before being introduced to travelling wave ion mobility cell. In travelling wave ion mobility spectrometry (TWIMS) pulsed voltage waves guide the ions through the drift cell via stacked ring electrodes.¹⁸² As the waves move, the ions are pushed for a short distance before moving over the wave. Smaller ions undergo fewer collisions with the buffer gas so travel faster in the traveling wave's gradient, whereas larger ions undergo more collisions with the buffer gas slowing them down therefore requiring a longer time to be moved by the traveling waves.^{140, 182} A pressure of 0.5 bar of nitrogen was maintained in the IM cell. Each set of experiments was run in triplicate.

2.4 Gel Electrophoresis

Both one- and two-dimensional gel electrophoresis was used to separate protein mixtures prior to digestion and mass spectrometry analysis. One-dimensional gel electrophoresis was performed using a small format cell (BioRad mini-PROTEAN). The main components of this gel apparatus are a gel cassette assembly, electrode assembly, clamping frame, mini tank and a lid

(see Figure 2-1). The electrode assembly can hold gels 8 cm (W) × 7.3 cm (H). In this work, precast polyacrylamide gels were used (BioRad Laboratories).

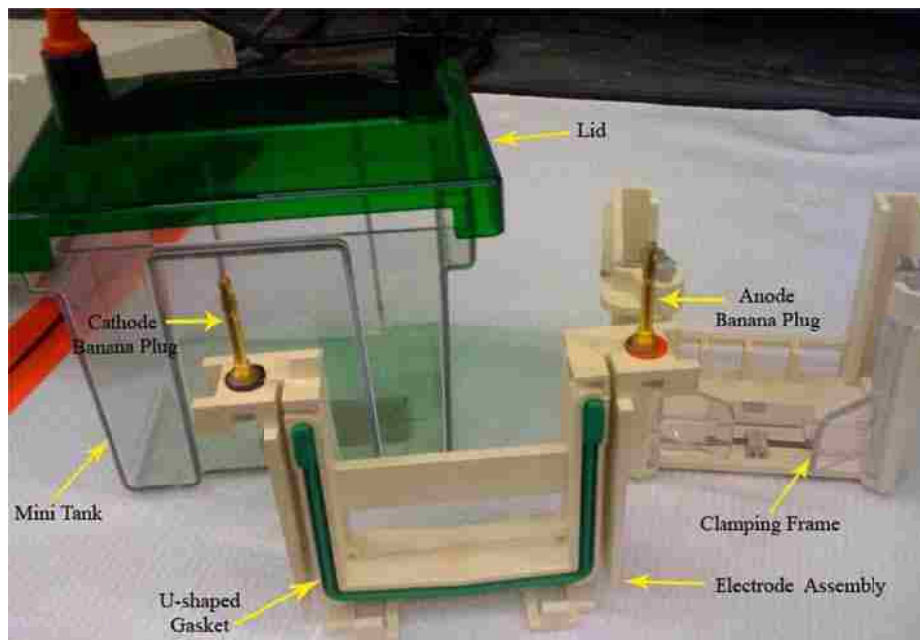


Figure 2-1. Assembly components of the mini-PROTEAN 2D gel electrophoresis cell.

Two-dimensional electrophoresis was performed using either the small format or a standard format electrophoresis system (Protean II XL, Bio-Rad) (see Figure 2-2). For isoelectric focusing, sample was dissolved in sample buffer containing 8 M urea, 2% 3-[(3-cholamidopropyl) dimethylammonio]-1 propanesulfonate (CHAPS), 50 mM dithiothreitol (DTT) and 1% carrier ampholytes. Isoelectric focusing was carried out in an isoelectric focusing cell (Protean, Bio-Rad) at 250 V for 20 min and 14,000 V-h (rapid ramping). SDS-PAGE electrophoresis was performed in slab gels (18.3×19.3 cm) with 1.00 mm thickness. The electrophoresis was conducted at 200 V for 40 min.

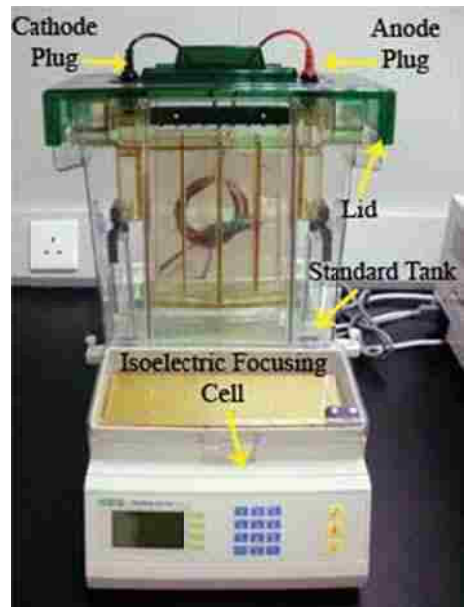


Figure 2-2. Protean II XL large format gel electrophoresis system and isoelectric focusing cell.

Gels can be prepared as a single continuous percentage or as a gradient %T through the gel; examples of commonly used gradient gels include 4–15% and 4–20%. Gradient gels have the advantage that proteins with a wide molecular weight range can be separated and the protein bands are narrower. In this work, 4–20% gels were used unless otherwise indicated.

Another component to the gel electrophoresis is the buffer system. There are two types of buffer systems: continuous and discontinuous.¹⁸³ A continuous buffer system has the same buffer at a constant pH and concentration. The gel used is typically made of one continuous %T. A discontinuous buffer system has a different buffer in the gel solution and the solution for the electrode assembly, and a stacking gel is also added to the resolving gel. The stacking gel has a lower acrylamide concentration and pH, which gives the gel larger pores allowing the proteins in the sample to be concentrated before entering the resolving gel. The resolving gel has a higher acrylamide concentration and higher pH which gives the gel narrow pores. This gives better protein resolution and separation.

The sample buffer system used in this work was a discontinuous buffer system that incorporated sodium dodecyl sulphate (SDS) and a reducing agent, β -mercaptoethanol, in the buffer. SDS is an anionic detergent that denatures and binds to proteins. The polypeptides become rod-like in shape with a negative charge.¹⁸⁴ Because all the proteins have the same charge, they separate according to their molecular mass. The β -mercaptoethanol is used to reduce the disulfide bridges in the proteins so that they can assume the random-coil configuration required for optimum separation.¹⁰⁶

To resolve proteins with similar molecular weights, two-dimensional polyacrylamide gel electrophoresis (2D-PAGE) was performed. 2D-PAGE is a powerful proteomics separation tool as it is capable of separating thousands of proteins in a typical cell.¹⁸⁵ The first dimension, isoelectric focusing (IEF) was performed in an isoelectric focusing cell (Bio-Rad Protean IEF). IEF is usually performed in an immobilized pH gradient strip.¹⁸⁶ The pH gradients used in this work were 3–10 and 3–6. Prior to the second dimension, the proteins separated by IEF underwent an equilibration step. The proteins were reduced and alkylated using DTT and iodoacetamide and SDS was added to give the proteins a negative charge. This allowed the separation based on size in the second dimension. The second dimension was performed using the Bio-Rad Protean II 2-D electrophoresis cell.

The gels were stained to visualize the proteins. Several stains are commercially available: Coomassie blue R-250, Bio-safe Coomassie, Sypro ruby fluorescent stain, and silver stain.^{106, 185} The choice of stain used depends on several factors such as sensitivity, ease of use, and type of imaging equipment available. Coomassie R-250 stain (wool dye with a red hue) is the least sensitive: it requires 40 ng of protein for detection.¹⁸⁵ Bio-safe Coomassie stain is prepared with Coomassie G-250 (wool dye with a green hue) and can be used to detect as little as 10 ng of protein.¹⁸⁵ The most sensitive stains are Sypro ruby fluorescent and silver stains which both can

be used to detect as little as 1 ng of protein.¹⁸⁵ In this work, Bio-safe Coomassie or silver stains were used. Both stains are mass spectrometry compatible, meaning that the stains have high sensitivity for protein detection while minimizing crosslinking of the protein to the gel, which can inhibit recovery for mass spectrometry.¹⁸⁷

There are also specific stains that can be used for detecting modified proteins such as glycoproteins. The most commonly used methods for detecting glycoproteins in polyacrylamide gels involves detection by periodic Schiff (PAS) staining using chromogenic detection, for example acid fuchsin dye.¹⁸⁸ Another approach utilizes Pro-Q Emerald 300 fluorescent dye, that links to glycoproteins using periodic Schiff conjugation.¹⁸⁹ During the staining process the glycols in the glycoproteins are oxidized to aldehydes with periodic acid. The dye then reacts with the aldehydes on the glycoproteins and a fluorescent conjugate is generated.¹⁸⁹ This stain can be used to detect as little as 0.5 ng of glycoprotein and is compatible with mass spectrometry analysis. This glycoprotein stain was used in the initial characterization of lectin proteins studied in this work.

Gel images were captured using a camera (Gel Doc XR System with Quantity One 1-D analysis software, BioRad) which is capable of imaging Coomassie stained, silver stained, and fluorescent gels.

2.5 Edman Sequencing

Edman sequencing involves labeling the N-terminal residue of a protein or peptide with phenyl isothiocyanate. The peptide bond adjacent to the modified residue is cleaved via mild acid hydrolysis, leaving the rest of the protein intact. The terminal amino acid residue that has been modified is identified by liquid chromatography. The process is repeated successively for the residues in the peptide sequence until a complete sequence is determined.⁶⁹ Protein bands were excised from SDS polyacrylamide gels and submitted to the University of Texas Medical

Branch, Protein Chemistry Core laboratory for N-terminal sequencing using a protein sequencer (LC 494, Applied Biosystems, Foster City, California).

2.6 Proteolysis of Isolated Proteins

Proteolysis generates peptides from proteins using either enzymatic or chemical cleavage. Proteolysis experiments were performed for peptide sequencing. Digestions were performed both in-gel and in-solution. Prior to proteolytic digestions, reduction and alkylation was performed to reduce the disulfide bonds for more efficient digestion. For reduction and alkylation a reaction buffer containing 25 mM NH_4HCO_3 was mixed with 10 mM dithiothreitol (DTT). To a vial containing approximately 100 μg of lyophilized protein in 25 mM NH_4HCO_3 , 10 μL of DTT solution was added and the sample mixture was heated for one hour at 57°C. The vial was removed from the heat block and allowed to cool to room temperature and 40 μL of 55 mM of iodacetamide (IA) solution was added and incubated at room temperature in the dark for 30 minutes.

Several enzymatic and chemical proteolytic agents were used: trypsin, α -chymotrypsin, Glu-C, Asp-N, Lys-C, cyanogen bromide (CNBr), and formic acid. The enzymatic digestions were performed in a heat block at 37°C, the CNBr digestions were performed at room temperature in a fume hood, and the FA digestions were performed at 37°C.

For the enzymatic digestions, a 0.2 $\mu\text{g}/\mu\text{L}$ solution of trypsin was prepared in 1 mM HCl and stored for a maximum of four weeks at -20°C. The reaction solution contained 1 μL of trypsin (0.2 μg), 9 μL of 40 mM NH_4HCO_3 and 9% acetonitrile by volume. For enzymatic digestions using Lys-C, Asp-N and Glu-C, the preparations were as follows: 0.1 $\mu\text{g}/\mu\text{L}$ of Lys-C in a 1:20 ratio (v/v), 0.04 $\mu\text{g}/\mu\text{L}$ of Asp-N in a 1:50 ratio and 0.1 $\mu\text{g}/\mu\text{L}$ of Glu-C in a 1:20 ratio. For CNBr digestions a 1 M solution was prepared in acetonitrile (ACN) and stored at -20°C. The CNBr solution contained 5 μL of 5% TFA/50% FA and 5 μL of water (v/v).

2.7 Mass Spectrometry

In this work, four commercial mass spectrometers were used: an ESI quadrupole-time-of-flight (QTOF), a MALDI tandem time-of-flight (TOF/TOF), an ESI time-of-flight, and a quadrupole/ion mobility separator/orthogonal acceleration time-of-flight (Q/IMS/oa-TOF) mass spectrometer

2.7.1 MALDI TOF/TOF

The MALDI TOF/TOF mass spectrometer used in this work was a Bruker UltrafleXtreme. A diagram of this instrument is shown in Figure 2-3. The Bruker TOF/TOF mass spectrometer is equipped with a Nd:YAG laser (wavelength, 355 nm; repetition rate 1 kHz) with beam homogenization (Smartbeam). It is also equipped with delayed extraction, a collision cell, linear and reflectron flight tubes.¹⁹⁰

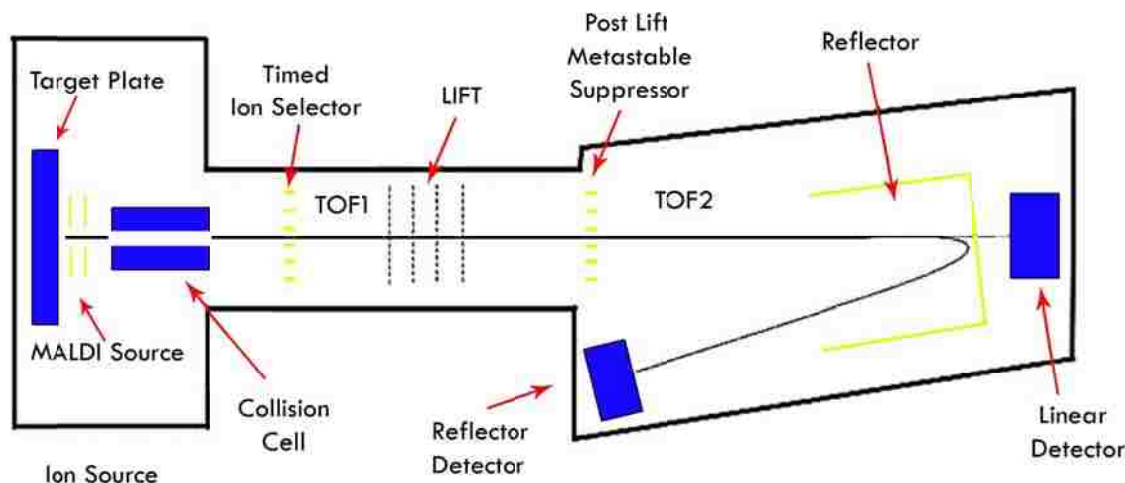


Figure 2-3. TOF/TOF mass spectrometer schematic.

This instrument can be operated in both linear and reflectron mode and it can be operated in MS and MS/MS mode. The instrument can achieve a mass resolution $\geq 40,000$ from 500 to 4000 Da and for intact proteins, with a mass accuracy < 1 ppm.¹⁹¹

The targets provided with the instrument have an integrated target transponder, which is read by the instrument allowing it to recognize the type of target plate being used for analysis. This instrument can use several different types of target plates however, for purposes of this study a 384 spot stainless steel plate was used.

The proteins and peptides were mixed on the 384 spot stainless steel plate using a 1:1 ratio of sample and saturated matrix solution. The matrices were 2-hydroxy-5-methoxybenzoic acid (sDHB), 2,5-dihydroxybenzoic acid (DHB), and 2,4-dimethoxy-3-hydroxycinnamic acid (sinapinic acid, SA) for proteins, and α -cyano-4-hydroxycinnamic acid (CHCA) for peptides. Structures and applications of the matrices used are found in Table 2-1. The protein samples were analyzed in linear positive ion mode and 500 shots averages were collected using a mass range from 30–100 kDa. For peptides, the spectra were obtained in reflector positive ion mode, using a mass range from 1–5 kDa. Masses were determined using FlexAnalysis software (Version 2.0, Bruker Daltonics).

Table 2-1. Commonly used matrices in MALDI for biomolecules.¹⁹²

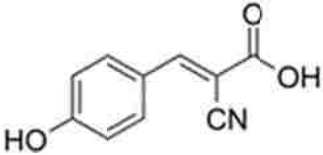
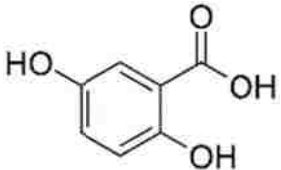
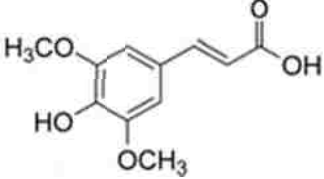
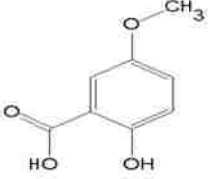
Matrix	Structure	Mass (Da)	Laser λ (nm)	Applications
α -cyano-4-hydroxycinnamic acid (CHCA)		189	337/355	Peptide/proteins mass < 10 kDa
2,5-dihydroxybenzoic acid (DHB)		154	337/355	Carbohydrates & Proteins mass < 5 kDa

Table 2-1. cont'd.

trans-3,5-dimethoxy-4-hydroxycinnamic acid (SA)		224	266/337/355	Peptide/proteins mass > 10 kDa
2-hydroxy-5-methoxybenzoic acid (sDHB)		168	337/355	Carbohydrates & Proteins mass < 5 kDa

2.7.2 LC-TOF Mass Spectrometer

An ESI-TOF MS (Agilent 6210) was used with a liquid chromatography HPLC system. The HPLC system used with this instrument was described in Section 2.2.2. This instrument consists of an electrospray source, a nebulizer and nebulizer gas inlet, capillary, skimmer, octopole ion guides, ion pulser for the orthogonal acceleration TOF, reflectron mirror, and a microchannel plate (MCP) detector. The capillary separates the spray chamber from the ion optics and has an applied potential from -4000 to 150 V. The ions are focused by lenses and enter the TOF mass analyzer. The ions pass into the TOF ion pulser and a high voltage pulse is applied to accelerate the ions into the flight tube. The ions are reflected by a reflectron into the detector. This instrument was used in this study to obtain intact protein mass measurements and has a mass accuracy of < 2 ppm and a mass resolution > 13,000 measured at m/z 2722. The limit of detection for this instrument using ESI is in the range of 1–10 ng/mL. Mass spectra were acquired from 600 to 3000 m/z using ESI at 4200 V. The spectra were extracted and deconvoluted using Analyst software (Version 1.5, Agilent).

2.7.3 ESI-Q-TOF Mass Spectrometer

The mass spectrometer used in this work for peptide sequencing was a hybrid quadrupole time of flight tandem mass spectrometer (Applied Biosystems QSTAR XL). A diagram of the instrument is shown in Figure 2-4. The mass spectrometer consists of an ESI source, three

quadrupoles (focusing quadrupole, q_0 ; mass selection quadrupole, Q_1 ; and collision quadrupole, q_2), an orthogonal TOF, reflectron ion mirror, and a MCP detector. This instrument was operated in MS/MS mode for these studies. The instrument has an orthogonal pulser that pulses the ions out at repetition rate of several kHz using an applied electric field of 10 kV.

To perform MS/MS experiments the instrument software (Analyst) was used to 1) select the three most intense peaks for fragmentation, 2) select three different collision energies (25, 32 and 38 eV), and 3) a repetition rate ranging between 3 to 20 kHz for pulsing the ions into the TOF MS.

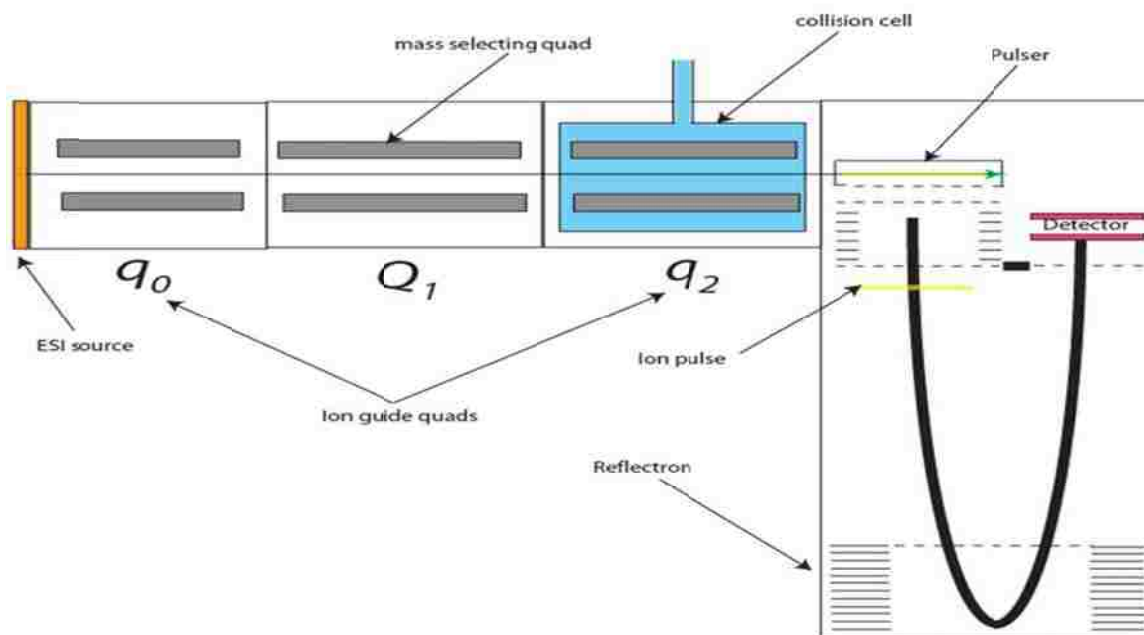


Figure 2-4. Q-TOF mass spectrometer schematic.

2.7.4 ESI-Q/IMS/TOF Mass Spectrometer

A hybrid quadrupole time-of-flight instrument with a travelling wave ion mobility cell was used in this work (Waters Synapt; Figure 2-5). This instrument has an ESI source, ion guide, quadrupole, ion mobility section, and an orthogonal time-of-flight. The ion mobility section is after the mass selection quadrupole and consists of a linear quadrupole ion trap which is used to

accumulate ions before the mobility separation; the accumulated ions are then released into the ion mobility cell.¹⁴⁰ The traveling wave transfer ion guide is used to transport the mobility-separated ions into the oa-TOF for mass analysis. A nano UPLC system is interfaced to a nanospray ionization source of the mass spectrometer. This instrument has a resolution of 15,000 and a mass accuracy < 10 ppm.

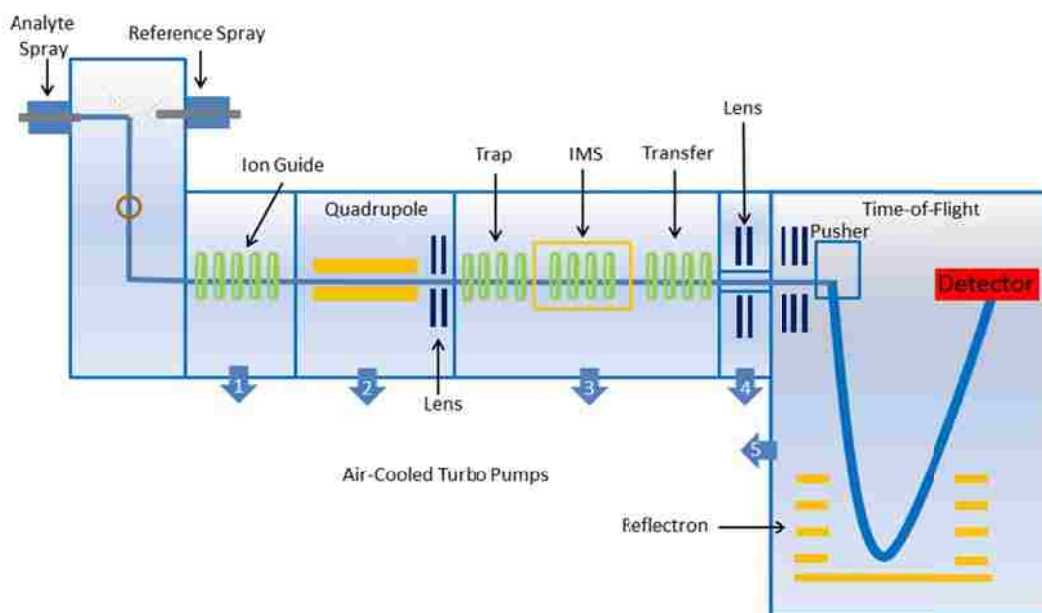


Figure 2-5. Schematic of the hybrid quadrupole/ion mobility separator/orthogonal acceleration time-of-flight.

In this study the instrument was operated in positive ion electrospray mode. The quadrupole was operated in RF mode in order to record full mobility spectra and the data was acquired using MassLynx software version 4.1. The peptide sample analysis on this instrument was performed at Waters Corporation, Milford, MA.

2.7.5 Protein Bioinformatics

Two approaches were used to sequence proteins: database searching and *de novo* sequencing. In the first method the tandem mass spectra from digested proteins were used to determine peptide sequences that were compared to protein sequences in databases. In this work,

the Mascot search engine was used for protein identification. The database search engine calculates peptide fragment masses *in silico* using the enzyme specificity for each protein sequence in the database. These calculated fragment masses are compared with the measured fragment masses and a score is calculated for comparison.^{152, 154} Search engines can also extract partial sequence information from the spectrum based on the ion series. The nomenclature for fragment ions was proposed by Roepstorff and Fohlman¹⁹³ and later modified by Biemann¹⁹⁴ to designate peptide fragment ions. When the charge is retained on the N-terminus the ions are represented by the symbols a, b, and c and when the charge is retained on the C-terminus the ions are represented by the symbols x, y and z (Figure 2-6). The most common ions produced are b and y-ions.¹⁹⁵

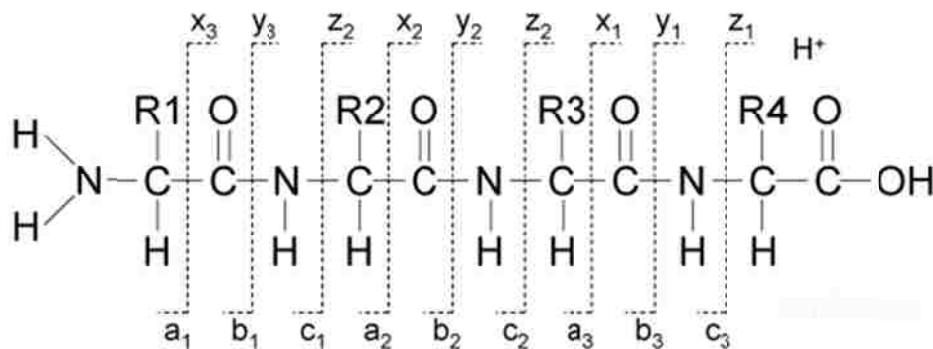


Figure 2-6. Common ions produced from peptide fragmentation.

To obtain sequence information from the tandem mass spectrum the molecular ion of the peptide is identified, which is the precursor ion in CID spectra followed by identifying the sequence ion series. The mass difference between the fragment ion peaks is determined and the masses of the 20 common amino acids are considered.

Mascot is based on the Mowse algorithm¹⁹⁶ which takes into account how many peptides of a given mass are in the database and also takes the protein size into consideration. Mascot also

calculates the probability that the observed match between the experimental data and a protein sequence is random.¹⁵²

The second approach, *de novo* sequencing with database searching, is necessary for organisms that do not have a large number of proteins in the sequence database. One source for protein database information is the genome sequence of an organism. Protein databases can be constructed using the translated protein sequence. Three *de novo* sequencing software tools were used in this work: Mascot Distiller,¹⁹⁷ PEAKS,¹⁶⁷ and BioTools. Mascot Distiller (Version 2.4, Matrix Science, Boston, MA) generates peak lists for *de novo* sequencing. The sequencing software fits each peak in the mass spectrum to a hypothetical amino acid combination of the same mass. A list of amino acid combinations corresponding to each peak is generated from the best fit data.¹⁹⁷

The second software used, PEAKS (Version 5.2, Bioinformatics Solutions, Waterloo, ON), applies four processes: preprocessing, candidate computation, refined scoring, and global and positional confidence scoring. First, the raw MS/MS data is smoothed and deconvoluted from doubly and triply charged ions to singly charged ions, and peak centering (taking the centroid of the first peak of the isotope distribution and removal of the other isotopic peaks) is performed. Second, sequences of all possible combinations of amino acids for the selected precursor mass are determined. The fragment ion series a, b, c, x and y are used to determine the peptide sequences. Next, the best sequences are scored using a stringent scoring scheme where the ion mass error tolerance is stricter. Last, PEAKS generates a confidence score for each of the top-scoring peptide sequences.¹⁶⁷

BioTools (Version 2.2, Bruker Daltonics, Billerica, MA) uses MS/MS spectra from MALDI-TOF/TOF to generate sequence tags of various lengths and scores. Proposed sequences are calculated from the masses in the MS/MS spectrum using sequence tags. Sequence tags are

produced using the fragment masses and information added by the user to the software, such as terminal amino acids and enzymes used. The suggested sequence tags are then extended to produce a list of all peptide sequences that could fit the MS/MS data. The sequences are scored based on the number of matched ions. Several candidate sequences may be calculated with similar scores; in this case the proposed sequences need to be manually fitted to the MS/MS data. However, if the peptide sequence produced complete fragmentation in the MS/MS spectrum one peptide sequence may be the best fit.¹⁹⁸

The peptide sequences found by *de novo* sequencing are used in protein database searches in order to find proteins with similar sequences. The protein databases used in this work were the National Center for Biotechnology Information (NCBI) database, Mass Spectrometry Protein Sequence Database (MSDB), and Swissprot. NCBI is a non-identical protein and nucleic acid database that has sequences taken from multiple sources including Swissprot, the Protein Data Bank, and translations from annotated coding regions in the GenBank. One of the major advantages of NCBI is that it is frequently updated; therefore, this database was used the most in this work. MSDB is also a non-identical protein database that comprises a number of other protein databases; however, it is not updated as frequently. Swissprot is a non-redundant protein database that provides a high level of annotations such as protein function and post-translational modifications.

Proteins were also determined from peptide sequences using Basic Local Alignment Search Tool (BLAST). BLAST is used for comparing high-scoring segment pairs (HSP) in nucleotide or protein sequences to sequences in nucleotide or protein databases. HSP is a region between a pair of sequences, either nucleotide or amino acid, that share high level of similarity.^{174, 199} BLAST can also provide evolutionary relationships for the species under consideration, which shows how two species are related with respect to their evolutionary

descent. For this work, BLAST was used to compare peptide sequences generated from alligator leukocyte and lectin proteins to proteins in the protein database. For BLAST searching, Swiss-Prot and NCBI databases were used to search peptide sequences using all taxonomies. The expect threshold was set to 20,000 and the score matrix used was PAM30.

To look for protein sequence similarities DIALIGN and CLUSTALW software were used. The DIALIGN software uses both local and global alignment features by either aligning all residues in each sequence or aligning only residue segments that are closely related.²⁰⁰ CLUSTALW allows for the alignment of multiple protein sequences in order to determine similarities and differences.²⁰¹

2.8 X-Ray Crystallography

In this work, small molecules were isolated from alligator leukocyte extracts and submitted for structural characterization using an X-ray diffractometer with charge coupled device (CCD) area-detector (Bruker Kappa Apex II, Madison, WI, USA). The X-ray diffractometer provides accurate and precise measurements of the entire three-dimensional structure of a molecule, including the bond angles and distances as well as crystal structure.

2.9 Deglycosylation

Deglycosylation is the removal of a sugar from a glycogen, especially a glycoprotein. The oligosaccharides can be removed using either an enzyme or a chemical reagent. There are several enzymes available for deglycosylation of N- and O-linked oligosaccharides; these include peptide-N-(N-acetyl- β -glucosaminyl)asparagine amidase (PNGaseF), endo F and endo H (endo- β -N-acetylglucosaminidases F and H) for removal of N-linked oligosaccharides,²⁰² as well as O-glycanase for removal of only galactose-N-acetylgalactosamine (Gal-GalNAc) O-linked oligosaccharides.²⁰³ A commonly used enzyme is PNGaseF, which was used in this work. PNGase cleaves between the innermost N-acetylglucosamine (GlcNAc) and asparagine residues

of high mannose (Man) content and complex oligosaccharides from N-linked glycoproteins (Figure 2-7).^{202, 204} This method is advantageous due to its high specificity and its nondestructive nature that allows the oligosaccharides to be recovered for further analysis.

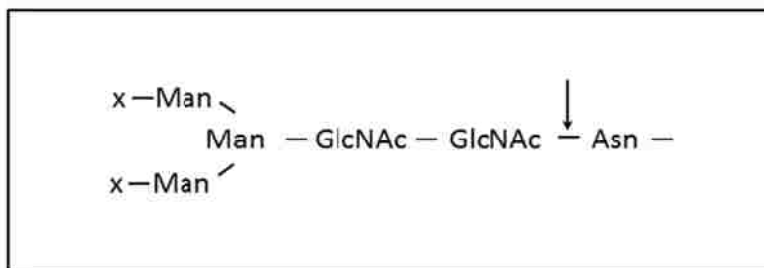


Figure 2-7. Illustration of PNGaseF cleavage location in glycoproteins. (x = H or sugars)

Another deglycosylation procedure used in this work was trifluoromethanesulfonic acid (TFMS) deglycosylation. TFMS is a commonly used chemical reagent that is used to remove oligosaccharides from glycoproteins.²⁰⁵ Protein amide bonds are stable to TFMS whereas glycosidic bonds are not. The chemical reaction is carried out in anhydrous conditions in TFMS without a solvent in the presence of a scavenger that protects the amino acids. During the solvolysis reaction, cleavage of the glycosidic bonds occurs and the protein remains intact. It is important to ensure that the reaction is performed in the absence of water to prevent cleavage of peptide bonds. TFMS is advantageous when compared to enzymatic deglycosylation because carbohydrates can be removed from a glycoprotein regardless of its structure or its linkage (N- or O-linkage).²⁰⁵

2.10 Radial Diffusion Assay

Antimicrobial activity can be monitored using growth inhibition assays. The antimicrobial peptides isolated in this work were tested at various stages using a radial diffusion assay.^{206, 207} Bacteria were incubated overnight at 37°C in trypticase soy broth (TSB) (Sigma

Aldrich) that was previously autoclaved at 120°C for 20 minutes. The TSB was prepared using 3.0 mg of TSB medium dissolved in 100 mL of nanopure water. A 20 μ L volume of bacteria was added to 15 mL of the TSB medium containing 1% (wt/vol) low-electroendomosis-type agarose (Sigma Aldrich). The previously prepared TSB medium containing agarose (agar) was heated in a water bath at 42°C before use. The agar was poured into an agar plate to form a uniform layer and allowed to solidify. Evenly spaced holes were punched using a sterilized pipette tip connected to a vacuum for suction to create wells. Added to individual holes was 5 μ L of control sample (0.1% acetic acid) and polymyxin B, which was used as the standard and isolated peptide samples (antimicrobial peptides). The samples were added in triplicate and the plates were incubated for 3 hours at room temperature followed by an overnight incubation at 37°C. After incubation, the size of the clear zone around each well was measured. The size of a clear zone around the well indicates the antimicrobial activity of the peptides. The absence of a clear zone indicates no antimicrobial activity.

2.11 Reagents and Chemicals

Ammonium bicarbonate and cyanogen bromide (CNBr) were obtained from Sigma Aldrich (St. Louis, MO). The chemicals trifluoroacetic acid (TFA) and acetic acid were purchased from Fisher Scientific. The MALDI matrix α -cyano-4-hydroxycinnamic acid (CHCA) was obtained from Sigma and 3, 5-dimethoxy-4-hydroxycinnamic acid (sinapic acid), DHB, and sDHB were obtained from Fluka. Trypsin, α -chymotrypsin, endoprotease Asp-N from *Pseudomonas fragi*, endoprotease Glu-C from *Staphylococcus aureus*, and endoprotease Lys-C from *Lysobacter enzymogenes* were obtained from Sigma-Aldrich (St. Louis, MO, USA). Dithiothreitol (DTT), iodacetamide, HPLC grade acetonitrile, formic acid, and glu-fibrinopeptide standards (Sigma-Aldrich) were used without further purification.

Laemmli sample buffer, 10× SDS-tris-glycine buffer, β-mercaptanol, tris-HCl gradient gel, Coomassie blue stain (Bio-Safe), and molecular weight marker standards were purchased from Bio-Rad (Hercules, CA, USA). Dithiothreitol, iodacetamide, and glu-fibrinopeptide were purchased from Sigma-Aldrich (St. Louis, MO, USA). Trifluoromethanesulfonic acid (TFMS), pyridine, and toluene were purchased as a kit from Prozyme (Hayward, CA, USA).

The bacteria strains used in this study were obtained from American Type Culture Collection (ATCC) (Manassas, VA). The following American Type Culture Collection (ATCC) bacterial strains were used: *Escherichia coli* (35218), *Shigella flexneri* (12022), *Enterobacter cloacae* (23355), *Klebsiella oxytoca* (33496), *Staphylococcus aureus* (51153), and *Staphylococcus epidermidis* (29887). Trypticase soy broth was purchased from Voigt Global Distribution (Broth 23400, Lawrence, KS) and agarose Type I was purchased from Sigma.

CHAPTER 3. PROTEOME ANALYSIS OF THE LEUKOCYTES FROM THE AMERICAN ALLIGATOR (*ALLIGATOR MISSISSIPPIENSIS*) USING MASS SPECTROMETRY*

The purpose of the research described in this chapter was to investigate proteins related to the alligator immune system. This chapter describes the use of mass spectrometry in conjunction with gel electrophoresis and liquid chromatography to determine peptide sequences from American alligator (*Alligator mississippiensis*) leukocytes and to identify similar proteins based on homology. Proteins from leukocyte extracts were separated using two-dimensional gel electrophoresis and the major bands were excised, digested, and analyzed with on-line nano LC MS/MS to generate peptide sequences. The sequences generated were used to identify proteins and infer their function based on similarity to previously identified proteins. Similar proteins were identified based on matching two or more peptides from the alligator protein to similar proteins by searching against the NCBI database using Mascot and Basic Local Alignment Search Tool (BLAST). For those proteins with only one peptide matching, the phylum of the organism corresponding to the matching protein was considered.

3.1 Introduction

A large amount of anecdotal evidence exists to suggest that crocodylians are resistant to bacterial, fungal and viral infections²⁰⁸⁻²¹⁰ and have the ability to fight a variety of diseases that may pose human health risks.^{211, 212} Several studies have shown that the serum of alligators exhibit innate immunity against fungi, viruses and various bacterial species.^{208, 213, 214} It has been reported that alligator leukocyte extracts exhibit broad-spectrum antimicrobial activity against important human pathogens such as *C. albicans*, *S. faecalis* and *E. coli*.²¹⁵ It is anticipated that elucidation of the alligator blood proteome will aid in identifying these antibacterial, antifungal

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and antiviral peptides believed to be responsible for these properties, which in turn could lead to the development of new antibiotics.

Antimicrobial peptides are typically found in granulocytic leukocytes and are part of the innate immune system;²¹⁶ these antimicrobial peptides are active against a wide range of microorganisms such as fungi, bacteria, and viruses.^{42, 217} Various classes of antimicrobial peptides have been isolated from organisms such as bacteria, plants and animals²¹⁸⁻²²⁰ and are characterized by their amino acid sequence and structure.²¹⁷ The majority of the antimicrobial peptides assume linear α -helical or disulfide-stabilized β -sheet conformations^{24, 217} and are typically rich in lysine and arginine; however, some are also rich in proline, histidine or tryptophan residues.²¹⁷ These antimicrobial peptides are typically less than 10 kDa in molecular mass and are cationic and amphipathic.^{24, 217} However, antimicrobial properties have been reported for anionic peptides as well as for proteins.²⁴

Mass spectrometry (MS) and tandem mass spectrometry (MS/MS) in combination with multi-dimensional separations have become powerful techniques for peptide and protein identification.^{221, 222} Mass spectrometry-based approaches have been extensively employed for the identification and characterization of proteins in blood components such as leukocytes,^{33, 223, 224} serum,²²⁵ platelets,^{226, 227} plasma²²⁸ and red blood cells.²²⁹ For example, online LC MS/MS was used to identify 1444 proteins in human serum.²³⁰ In other studies, the human blood plasma proteome has been characterized using multi-dimensional separation techniques along with MS/MS and database searching to identify over a thousand proteins.^{125, 221, 228}

One of the major challenges in the study of the alligator blood proteome is the limited information available regarding the reptilian genome and proteome.^{33, 231} For proteomic analysis of species with limited genomic and proteomic data, *de novo* sequencing can be used to determine protein sequences.¹⁷⁵ In *de novo* sequencing, proteins are isolated and enzymatically

digested and the resulting peptides are analyzed by tandem mass spectrometry. The sequence is determined by observing the mass differences between fragment peaks in the tandem mass spectra which are generated by low energy collision-induced dissociation (CID) of the peptide of interest.²³² The amino acid sequence can be determined from the N-terminus using the b-ion series or from the C-terminus using the y-ion series.²³³ *De novo* sequencing has been used to identify a number of proteins from species with limited representation in protein databases.²³⁴⁻²³⁷

In the present study, proteins were isolated from the leukocytes of the American alligator (*Alligator mississippiensis*) and separated by one or two dimensional gel electrophoresis. One of the challenges in this study is limited proteome data for the alligator hence peptide sequences were determined using *de novo* sequencing with database searching and BLAST search to identify similar proteins. Using this approach 43 proteins were identified that represent abundant proteins from the alligator leukocyte.

3.2 Experimental

Adult alligators were captured for blood samples and leukocytes were isolated as described in Section 2.1. One-dimensional polyacrylamide gel electrophoresis (PAGE) was performed using precast 4–20% polyacrylamide gradient gels (8.6 × 6.8 cm) on a small format gel electrophoresis system. Leukocyte extracts were dissolved in sample diluting buffer (Tris-HCl at pH 6.8, containing SDS, glycerol, 2-mercaptoethanol, and bromphenol blue dye) and boiled for 3 min prior to loading on the gel. A 10 µl sample volume was loaded on each individual lane and in a separate lane, 10 µl of a protein molecular weight standard solution was loaded. Electrophoresis was conducted for 1 hr at 100 V using tris-glycine (pH 8.3) as the gel running buffer. The gels were then stained overnight with Coomassie blue to aid in visualizing the proteins bands. The gels were rinsed twice with distilled water for 10 min each time to

remove excess stain. The molecular weights of the proteins were estimated using protein molecular weight standards.

Two-dimensional electrophoresis was performed using either the small format or a standard format electrophoresis system. Additional two-dimensional PAGE was also performed by Kendrick Lab Inc. For isoelectric focusing, leukocyte extract was dissolved in sample buffer containing 8 M urea, 2% 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate (CHAPS), 50 mM dithiothreitol (DTT) and 1% carrier ampholytes. The linear IEF gel strips (7 cm), pH 3–10 were rehydrated with sample overnight. Isoelectric focusing was carried out in an isoelectric focusing cell at 250 V for 20 min and 14,000 V-h (rapid ramping). SDS-PAGE electrophoresis was performed in slab gels (18.3 × 19.3 cm) with 1.00 mm thickness. The electrophoresis was conducted at 200 V for 40 min. The gels were stained with ProteoSilver Plus silver stain overnight to visualize the protein bands.

The bands from the 1-D gels were excised and cut into $\sim 1 \text{ mm}^3$ cubes, and the Coomassie blue was washed away with 200 mM NH_4HCO_3 and 40% acetonitrile (2 × 30 min). For silver stained spots, the ProteoSilver Plus stain was removed using commercial destaining solutions.²³⁸ After destaining, the gel pieces were dried in a vacuum centrifuge, rehydrated with 20 μL of 0.4 μg trypsin in 40 mM ammonium bicarbonate containing 9% acetonitrile to completely cover the gel pieces. The tubes were then incubated overnight at 37 °C.^{239, 240} Peptides were extracted from gel pieces with 60% acetonitrile and 40% water containing 0.1% trifluoroacetic acid (TFA).

Liquid chromatography electrospray tandem mass spectrometry (LC MS/MS) was used to analyze the peptides extracted from the gel spots. The tryptic peptides were dissolved in 50 μL of 0.1% formic acid (FA) and 10 μL of the digest was injected with a microplate autosampler onto a 0.3 × 1 mm trapping column (PepMap C₁₈) on a nano LC system at a flow rate of 10 $\mu\text{L}/\text{min}$. After sample loading, the trapping column was washed with 0.1 % FA at a flow rate of 5

$\mu\text{L}/\text{min}$ for an additional 5 min. The peptides were then eluted onto a $75\ \mu\text{m} \times 15\ \text{cm}\ \text{C}_{18}$ column (Biobasic Vydac) and separated using a gradient of 5–40% B over 60 min with a flow rate of 200 nL/min. Solvent A was 95% water and 5% acetonitrile containing 0.1% FA. Solvent B was 80% acetonitrile and 20% water containing 0.1% FA

The effluent was directed to the quadrupole time-of-flight mass spectrometer (QSTAR XL) and ionized using a nano electrospray source at a voltage of 2.5 kV. The mass spectrometer was operated in information dependent acquisition (IDA) mode. Three collision energies (25, 38, and 50 eV) were selected for fragmentation of the peptides.

The peptide sequences were generated from MS/MS data using Mascot Distiller software. The software determined amino acid sequences from MS/MS data using peak fitting and isotope distribution.^{241, 242} The *de novo* sequencing results were verified manually.

De Novo peptide sequences were subjected to database searching using Mascot¹⁵⁴ and BLAST to find proteins that contain sequences matching with sequences determined by *de novo* sequencing. For BLAST searching, Swiss-Prot and NCBI databases were used to search peptide sequences using all taxonomies. The expect threshold was set to 20,000 and the score matrix used was PAM30.

3.3 Results and Discussion

One-dimensional electrophoresis of the leukocyte extracts revealed six major bands ranging from 10 to 75 kDa. The bands marked in Figure 3-1 were excised, digested with trypsin and analyzed by LC-MS/MS.

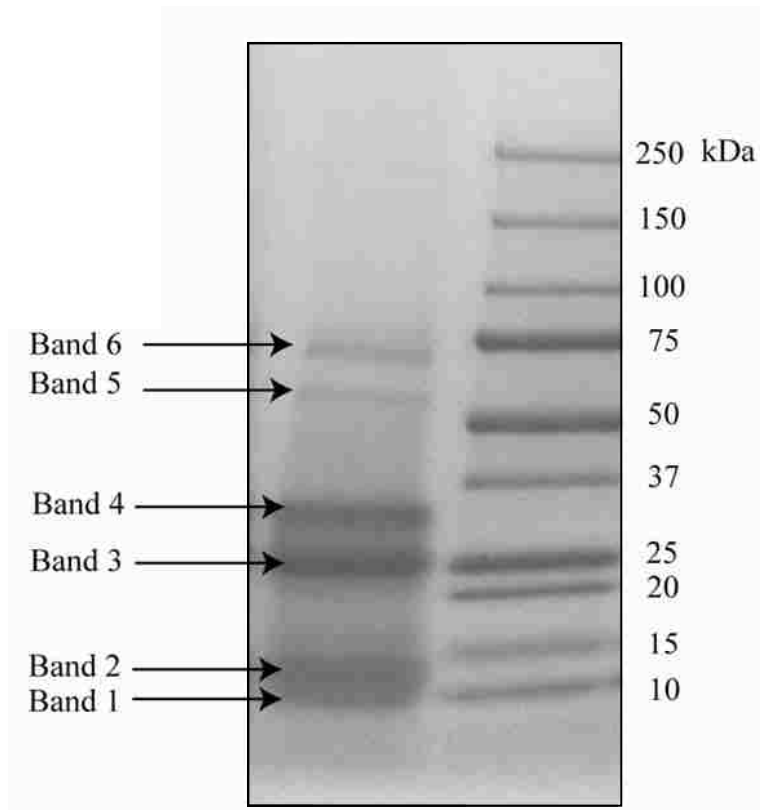


Figure 3-1. Alligator leukocyte extract separated on 1D gel electrophoresis and stained with Coomassie blue. Band numbers are referenced in Tables 1 and 2.

Figure 3-2 shows a two dimensional separation of the leukocyte extracts stained with mass spectrometry compatible silver stain. The number of proteins visible on the 2D gel from the alligator leukocyte was lower than a typical cell extract due to acetic acid denaturation of some proteins. The majority of the protein spots were observed in the pI range of approximately 5–7 and molecular weight range of approximately 20 to 200 kDa. Thirty-seven of the most intense well resolved spots (labeled on the 2D gel in Figure 3-2) were selected for further analysis and excised, digested and analyzed by LC MS/MS.

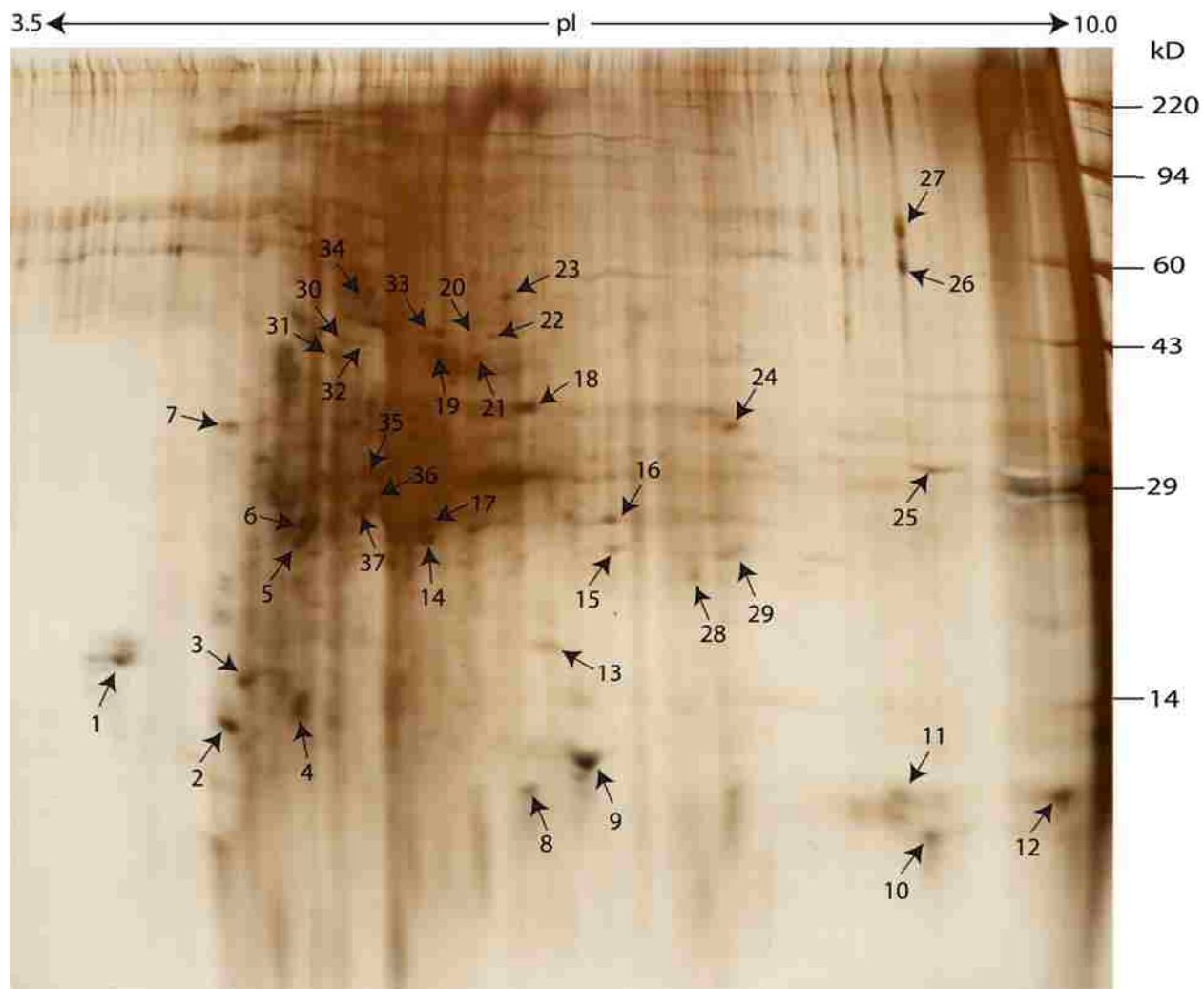


Figure 3-2. 2D separation of alligator leukocyte stained with Proteosilver stain for mass spectrometry analysis. Approximately 400 μg of leukocyte protein was loaded and separated on a 2D large gel format.

Figure 3-3 shows a selected separation and associated mass spectra that illustrates the method of peptide sequences determination. The LC separation in Figure 3-3a shows the total ion chromatogram for the digest of Band 1. Figure 3-3b shows an example of a mass spectrum acquired at 33.3 minutes as indicated with an asterisk. Tandem MS data were collected and peptide sequences were determined by *de novo* sequencing. Figure 3-3c illustrates the peptide fragment series that was found using *de novo* from the MS/MS data acquired from Band 1 of

Figure 3-1. A similar strategy was followed for the other labeled bands from Figure 1 as well as the protein spots in Figure 3-2.

For the characterization of peptides and proteins in the gel bands and spots, a three-step strategy was adopted. First, the peptides from the digest were sequenced using *de novo* sequencing as outlined above. Second, the sequences were searched against the full SwissProt and NCBI databases using Mascot to identify similar proteins. Third, a BLAST search was performed for *de novo* peptide sequences that did not match sequences in the database search of step two.

The Mascot search results are summarized in Table 3-1. The matched peptides shown in the table were identified from either the 1D or 2D gel as indicated in the table. The proteins included in the table had at least two peptide matches and sequence coverage ranging from 1 to 37%. The proteins polyubiquitin, hypothetical protein LOC100158585, ubiquitin, hypothetical protein, hypothetical protein 4732456N10, endonuclease P1, myosin alkali light chain, myosin regulatory light chain, actin, β -actin, actin-2, talin 1, chain A, chicken annexin V {complex with Ca^{2+} }, alkaline phosphatase-like enzyme, lysozyme C, ras-related protein O-Krev, annexin I, hemoglobin and vimentin-4 proteins were identified.

The *de novo* sequenced peptides that didn't match sequences in the database were subjected to a BLAST search to identify proteins containing similar sequences. The results of the BLAST search are summarized in Table 3-2 (and Appendix A, Table 1).

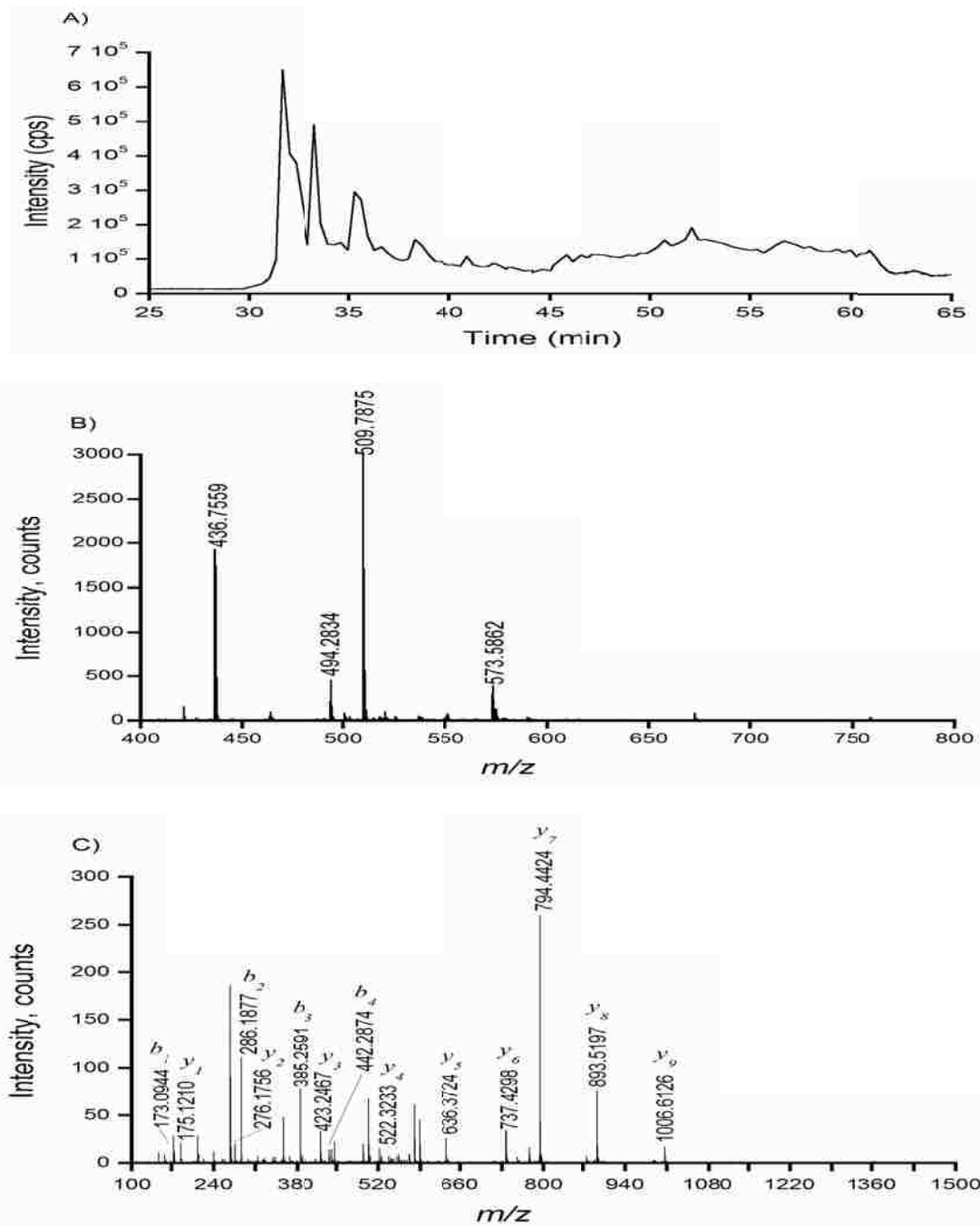


Figure 3-3. Mass spectrometry data for peptide sequencing. Figure (a) shows the total ion chromatogram of all the mass spectra recorded during the gradient LC separation as a function of time. The * in Figure (a) indicates the trace for a particular ion eluted at approximately 33.3 minutes. Figure (b) shows the mass spectrum of the peptides eluted during the time indicated in Figure (a). Figure (c) shows the MS/MS plot of the peptide ion highlighted in Figure 2 (b). The mass differences between the y-ions series is used to predict the amino acid series. The y-ion series is written in the direction from carboxy to amino terminus direction, going from left to right.

Table 3-1. Proteins identified in leukocyte extract using Mascot search. Proteins were identified by Mascot search of MS/MS data from the leukocyte peptide digest with databases, NCBI, MSDB and SWISS-PROT. NCBI accession numbers are provided in table.

Band/Spot No.	m/z	Charge	Matched Peptides	Name	Function	Accession No.	Mascot Score	% Coverage
B1	383.2	+2	MQIFVK	Polyubiquitin	a	BAA23486	340	8
	520.3	+2	EGIPPDQQR					
	534.8	+2	ESTLHLVLR					
	541.8	+2	TLSDYNIQK					
	508.6	+3	IQDKEGIPPDQQR					
B1	894.5	+2	TITIEVEPSDTIENVK	Ubiquitin	a	AAG22093	368	32
	520.3	+2	EGIPPDQQR					
	526.3	+2	NSTLHLVLR					
	534.8	+2	ESTLHLVLR					
	541.3	+2	TLSDYNIQK					
B3	508.6	+3	IQDKEGIPPDQQR	Hypothetical protein LOC100158585	b	NP_001121485	70	3
	895.0	+2	TITNEVEPSDTIENVK					
	902.0	+2	TITKEVEPSDTIENVK					
	405.2	+2	LASYLDK					
	533.3	+2	AQYEDLAKK					
B3	515.3	+2	VLDELTAR	Hypothetical protein	b	XP_511487	218	3
	579.3	+2	QGVEADVNGLR					
	651.4	+2	ALEEANADLEVK					
	453.3	+3	QSVEADINGLRR					
B2	565.8	+2	NLASAVS DLLK	Hypothetical protein 4732456N10	b	NP_808385	187	6
	871.8	+3	AVSSAIAHLLGEVAQGNENYTGIAAR					
3	639.3	+2	LALDIEIATYR	Hypothetical protein 4732456N10	b	NP_808385	187	6
1	651.8	+2	SLNLDSIIAEVK					
	738.4	+2	FLEQQNKVLETK					
1	518.9	+3	LIGGHALS DAESWAK	Endonuclease P1	c	P24289	82	11
	565.6	+3	LANWINEIHGSEIAK					
2	518.9	+3	LIGGHALS DAESWAK	Endonuclease P1	c	P24289	255	25
	565.6	+3	LANWINEIHGSEIAK					
	567.9	+3	VSDSSLSENHAEALR					
	851.5	+2	VSDSSLSENHAEALR					
4	665.5	+4	FLVHFIGDMTQPLHDEAYAVGGNK	Endonuclease P1	c	P24289	72	11
	518.9	+3	LIGGHALS DAESWAK					
5	565.6	+3	LANWINEIHGSEIAK	Endonuclease P1	c	P24289	100	11
	518.9	+3	LIGGHALS DAESWAK					
7	565.6	+3	LANWINEIHGSEIAK	Endonuclease P1	c	P24289	72	11
	518.9	+3	LIGGHALS DAESWAK					
9	565.6	+3	LANWINEIHGSEIAK	Endonuclease P1	c	P24289	381	20
	518.9	+3	LIGGHALS DAESWAK					
11	565.6	+3	LANWINEIHGSEIAK	Endonuclease P1	c	P24289	72	11
	518.9	+3	LIGGHALS DAESWAK					
12	565.6	+3	LANWINEIHGSEIAK	Endonuclease P1	c	P24289	98	11
	518.9	+3	LIGGHALS DAESWAK					
2	498.3	+2	HVLVTLGEEK	Myosin alkali light chain	d	AAA48979	181	30
	513.2	+2	EAFQLFDR					
	686.8	+2	ALGQNPTNAEVMK					
	575.2	+3	VFDKEGNGTVMGAEIR					
3	518.2	+2	ELLTTMGDR	Myosin regulatory	e	1805343A	189	18

^a Removal of damaged or unnecessary proteins and controlling of other cellular processes

^b Unknown function

^c Hydrolyzes only single stranded DNA and RNA without specificity for bases

^d Binds actin and modulates myosin motor function

^e Affect myosin motor function and kinetics

Table 3-1 cont'd.

				light chain				
	614.8	+2	LNGTDPEDVIR					
	708.3	+2	FTDEEVDELYR					
6	499.7	+2	DLTDYLMK	Actin	^f	<u>ACD44586</u>	188	14
	895.9	+2	SYELPDGQVITIGNER					
14	527.3	+2	TALAPSTMK	Actin	^f	<u>ACF32719</u>	79	4
17	566.7	+2	GYSFTTAAER	β-actin	^f	<u>ABB91894</u>	255	37
	581.3	+2	EITALAPSTMK					
	895.9	+2	SYELPDGQVITIGNER					
	855.4	+3	GYSFTTAAER					
19	473.2	+2	AVFPSIVGR	β-actin	^f	<u>CAA27396</u>	81	7
	652.0	+3	VAPEEHPVLLTEAPLNPK					
33	400.2	+2	AVFPSIVGR	Actin-1	^f	<u>P02578</u>	275	18
	505.9	+3	IWHHTFYNELR					
	652.0	+3	VAPEEHPVLLTEAPLNPK					
	499.7	+2	DLTDYLMK					
	566.7	+2	GYSFTTAAER					
	581.3	+2	ELTALAPSTMK					
20	473.2	+2	AVFPSIVGR	β-actin	^f	<u>ABD48797</u>	139	10
	505.9	+3	IWHHTFYNELR					
	652.0	+3	VAPEEHPVLLTEAPLNPK					
35	499.7	+2	DLTDYLMK	Actin	^f	<u>AAA20641</u>	103	13
	652.0	+3	VAPEEHPVLLTEAPLNPK					
	855.7	+3	LCYDALDFEQEMQTAASSSLEK					
36	499.7	+2	DLTDYLMK	Actin-2	^f	<u>P45885</u>	164	14
	505.9	+3	IWHHTFYNELR					
	652.0	+3	VAPEEHPVLLTEAPLNPK					
	654.3	+3	YPIEHGIITNWDDMEK					
37	505.9	+3	IWHHTFYNELR	Actin	^f	<u>Q99023</u>	119	9
	652.0	+3	VAPEEHPVLLTEAPLNPK					
	499.7	+2	DLTDYLMK					
37	378.2	+2	LLLAVVK	Chain A, Crystal Structures of Chicken Annexin V in Complex with Ca ²⁺	^g	<u>1YII_A</u>	102	5
	501.3	+2	VLTEILASR					
B2	565.8	+2	NLASAVSDLLK	Talin 1	^h	<u>XP_002192479</u>	141	1
	871.8	+3	AVSSAIAHLLGEVAQGNENYTGIAAR					
9	539.9	+2	GTSIFGLAPSK	Alkaline phosphatase-like	ⁱ	<u>NP_001096792</u>	114	4
	714.9	+2	DKQNLVQAWQAK					
12	877.4	+2	NTDGSTDYQILQINSR	Lysozyme C	^j	<u>LZFER</u>	114	10
13	747.9	+2	INVNEIFYDLVR	Ras-related protein O-Krev	^k	<u>P22123</u>	95	6
18	701.4	+2	GVDEATHIILTK	Annexin I	^l	<u>CAA39971</u>	141	8
	851.9	+2	GLGTDEDLNEILASR					
31	578.3	+2	FGGPGTASRPS	Vimentin-4	^m	<u>P24790</u>	151	6
	627.7	+2	TRSYVTSTRTYSLG					
	561.2	+2	DLYEEEMR					
12	510.8	+2	VLASFGAEVK	Hemoglobin β chain	ⁿ	<u>P02130</u>	131	19
	560.3	+2	KFIVDLWAK					
	566.7	+2	FHVDPENFK					

^f Highly conserved proteins that are involved in various types of cell motility and are ubiquitously expressed in all eukaryotic cells

^g Exocytosis

^h Cytoskeletal protein in lymphocyte involved in connections of major cytoskeletal structures to the plasma membrane

ⁱ Causes the chemical reactions that remove phosphates from a number of different molecules

^j Cell derived leukocyte that is part of the immune system to fight against infectious diseases and foreign material

^k Controls a spectrum of diverse cellular processes

^l Ca²⁺ dependent phospholipid binding proteins with potential anti-inflammatory activity

^m Filament protein

ⁿ Involved in oxygen transport from the lung to the various peripheral tissues

Table 3-2. Proteins identified at single peptide level using BLAST. *De novo* sequenced peptides obtained from a gel digest were searched using BLAST and proteins were matched based on sequence similarity and evolutionary relationship using the NCBI database. The top E-values with the closest related organisms are reported.

Band Number	<i>m/z</i>	Charge	<i>De novo</i> Sequence	Protein	Function	Accession Number	Protein Length	Organism	E-Value
B4	1060.6	+2	EINLSPDSTSAVVSGLMVATK	Fibronectin	^a	<u>P11722</u>	1256	Gallus gallus	2e-11
B4	536.3	+2	YEVSVYALK	Fibronectin	^a	<u>P11722</u>	1256	Gallus gallus	0.33
B4	701.3	+2	YQINQQWER	Fibronectin	^a	<u>Q91740</u>	2481	Xenopus laevis	0.013
B1	613.9	+2	LIALLEVLSQK	Filamin-C	^b	<u>Q14315</u>	2725	Homo sapiens	0.019
B1	487.3	+2	VYGPGE	Filamin-C	^b	<u>Q14315</u>	2725	Homo sapiens	36
B2	736.4	+2	AQQVSQGLDLLTAK	Vinculin	^c	<u>P12003</u>	1135	Gallus gallus	5e-05
B2	739.4	+2	LGQMTDQ+ADLR	Vinculin	^c	<u>P12003</u>	1135	Gallus gallus	0.006
B2	655.9	+2	TVTAMDVVYALK	Histone H4	^d	<u>P70081</u>	103	Gallus gallus	6e-04
B2	628.7	+2	LITKAVSASK	Histone H1	^e	<u>P09987</u>	218	Gallus gallus	0.49
B6	833.9	+2	FSGSGSGTDFFTIS	Ig K chain V-I region Lay	^f	<u>P01605</u>	108	Homo sapiens	5e-06

^a Cell adhesion, cell motility, opsonization, wound healing & cell shape maintenance

^b Plays a central role in muscle cells, probably by functioning as a large actin-cross-linking protein

^c Involved in attachment of the actin-based microfilaments to the plasma membrane; may also play roles in cell morphology & locomotion

^d Plays a central role in transcription regulation, DNA repair, DNA replication and chromosomal stability; a mixture of histones H2B and H4 has antimicrobial activity against Gram-positive bacterium *M.luteus*

^e Condensation of nucleosome chains into higher order structures

^f Used by the immune system to identify and neutralize foreign objects, such as bacteria and viruses

Table 3-2 cont'd.

B6	751.4	+2	VFGGGTKLTVL QPK	Ig A chain V-III region LOI	^g	<u>P80748</u>	111	Homo sapiens	7e-05
B2	510.8	+2	VLASFGEAVK	Hemoglobin, β	^h	<u>P02130</u>	146	Alligator mississippiensis	0.27
B2	500.8	+2	PISGDPK	Microtubule-actin cross- linking factor 1	ⁱ	<u>Q9UPN3</u>	5430	Homo sapiens	54
B6	522.8	+2	IMSIVDPNR	α -actinin-1	^j	<u>P05094</u>	893	Gallus gallus	0.1
B5	506.2	+3	QEYDESGPSIVHR	γ -Actin	^k	<u>Q5JAK2</u>	375	Rana lessonae	3e-05
B1	683.9	+2	STDYGILQINSR	Lysozyme C	^l	<u>P00698</u>	147	Gallus gallus	6e-04
B1	503.3	+2	WDAW+ALK	Acyl-CoA-binding protein	^m	<u>Q9PRL8</u>	86	Gallus gallus	1.7
B4	509.8	+2	MLPMQK	Myeloid protein 1	ⁿ	<u>P08940</u>	326	Gallus gallus	36
B1	565.3	+2	LVTDVQEAVR	Proactivator polypeptide (Containing Saposin-A, B, C & D)	^o	<u>Q13035</u>	518	Gallus gallus	0.084
B2	646.9	+2	ISMPDFDLNLK	Neuroblast differentiation-associated protein AHNAK	^p	<u>Q09666</u>	5890	Homo sapiens	0.001
B2	772.0	+2	AVASAAAALVLK	Talin-1	^q	<u>P54939</u>	2541	Gallus gallus	0.026

^g Activates the alternative complement pathway by binding to the short consensus repeat domain 3 (SCR3) of factor H

^h Involved in oxygen transport from the lung to the various peripheral tissues

ⁱ Cross-linking actin to other cytoskeletal proteins

^j Anchor actin to a variety of intracellular structures

^k Highly conserved protein involved in various types of cell motility

^l Enhances the activity of immunoagents

^m Displace diazepam from the benzodiazepine recognition site located on the GABA type A receptor. Diazepam binding inhibitor (32-86) has antibacterial properties

ⁿ Cell derived leukocyte that is part of the immune system to fight against infectious diseases and foreign material

^o Lysosomal degradation of sphingolipids takes place by the sequential action of specific hydrolases

^p Required for neuronal cell differentiation

^q Cytoskeletal protein in lymphocyte involved in connections of major cytoskeletal structures to the plasma membrane

Table 3-2 cont'd.

B2	525.8	+2	LGTFLEN	Histone acetyltransferase p300	^r	<u>Q09472</u>	2414	Homo sapiens	36
B4	821.9	+2	V+QQQADDAEER	Putative tropomyosin α -3 chain-like protein	^s	<u>A6NL28</u>	223	Homo sapiens	0.006
B6	587.3	+2	PP+PPARAA	Protein bassoon	^t	<u>Q9UPA5</u>	3926	Homo sapiens	17
B5	506.8	+2	PSLPSGVD	Stromelysin-1	^u	<u>P08254</u>	477	Homo sapiens	12
B6	583.8	+2	AAGEIIAIPRR	Envelope glycoprotein gp160	^v	<u>Q89607</u>	852	HIV-2 B_EHO	0.011
B1	831.4	+2	NSWGTSWGEGDYFR	Dipeptidyl-peptidase 1 (Cathepsin C)	^w	<u>Q60HG6</u>	463	Macaca fascicularis	5e-07
B5	684.9	+2	FRTTMLQDSI	Origin recognition complex subunit 2	^x	<u>Q75PQ8</u>	576	Rattus norvegicus	0.014
B6	586.8	+2	LP EQGT SSR	Cadherin EGF LAG seven-pass G-type receptor 2	^y	<u>Q9QYP2</u>	2144	Bos Taurus	0.36
B6	593.3	+2	AILYNYWDK	Complement component c3	^z	<u>CAC69535</u>	401	Crocodylus niloticus	0.50
B6	506.8	+2	NEALIALLR	Plastin-2	^{aa}	<u>P13796</u>	627	Homo sapiens	0.79

^r Functions as histone acetyltransferase and regulates transcription via chromatin remodeling

^s Binds to actin filaments in muscle and non-muscle cells

^t Involved in the organization of the cytomatrix at the nerve terminals active zone (CAZ) which regulates neurotransmitter release

^u Can degrade fibronectin, laminin, gelatins of type I, III, IV, and V; collagens III, IV, X, and IX, and cartilage proteoglycans

^v The surface protein gp120 (SU) attaches the virus to the host lymphoid cell by binding to the primary receptor CD4

^w Plays a role in the generation of cytotoxic lymphocyte effector function; the central coordinator for activation of many serine proteases in immune/inflammatory cells

^x Component of the origin recognition complex (ORC) that binds origins of replication

^y Receptor that may have an important role in cell/cell signaling during nervous system formation

^z Plays a central role in the complement system and contributes to innate immunity

^{aa} Actin-binding protein; plays a role in the activation of T-cells

Peptide sequences found using BLAST was used in a search of both the SwissProt and NCBI databases. Of the peptides found by *de novo* sequencing, 81% matched similar proteins in the two databases. The proteins were identified at single peptide level by observing the E-value, which is a statistical value indicating sequence similarity and potential evolutionary relationship.²⁴³ A low E-value indicates a statistically significant match. The peptides listed in Table 3-2 are those returned from the BLAST search having the lowest three E-values (the complete Table can be found in Appendix A). Notably, the organisms associated with the low E-value peptides are close in homology to alligators, for example *Chelonia caretta* (turtle), *Caiman crocodilus* (caiman) and *Gallus gallus* (chicken). Eleven of the thirty-two *de novo* sequences had chicken protein sequences in the top three hits. Also, there were several identified proteins that are associated with leukocytes and the immune system. However, the identification of hemoglobin β suggests contamination from red blood cells during isolation of the leukocytes from the whole blood. Other peptides in the table matched proteins that have not, to date, been associated with the immune system.

To assess the quality of the protein matches, several factors were considered. First, the number of peptides matching the protein was considered. For example, three of the *de novo* sequences in Table 3-2 matched the protein fibrinectin, which increases the confidence level that a similar protein is present in the alligator leukocyte. The percentage of matching residues was also considered. The *de novo* sequenced peptides matching fibrinectin had 100% sequence alignment with that protein. However, the sequence coverage was only 3%.

Last, a BLAST search was performed using a limited taxonomy containing birds and reptiles. Limiting the taxonomy search reduced the random matches and resulted in lower E-values for the protein matches compared to the full database search. This data is shown in the supplementary material (Appendix A, Table 2).

The proteins were grouped based on their functionality. Protein functions were divided into six groups: other, cytoskeletal proteins, immune proteins, enzymes, DNA/synthesis proteins, and unknown. Proteins involved in the cytoskeletal system, immune system, and other make up the three most abundant groups of peptides with approximately 37%, 23% and 23% respectively.

De novo sequencing coupled with Mascot and BLAST searches aided in the identification of 43 proteins that are similar to the proteins isolated from the alligator's leukocyte extract. To identify and characterize the function of the proteins derived from the *de novo* sequenced peptides, two approaches were employed. First, if a protein was identified from more than one *de novo* sequenced peptide in the same gel band, the protein and its function were assigned. Second, *de novo* sequenced peptides with only one peptide matching a protein, the phylum of the matched protein was considered. From the three highest E-values from the BLAST search, seen in Table 3-2 (for the complete table, refer to Appendix A), E-values less than one were first considered and the organisms associated with the protein hit were classified by biological phylum and only animals were considered. Matched proteins that are from closely related species, e.g. reptiles and birds, were moved to the top of the list. Of the *de novo* sequenced proteins, 29% matched invertebrates 39% mammals, 18% birds, 6% amphibians, 5% fish and 3% reptiles. Although proteins from invertebrates are highly represented, it should be noted that 81% of the database proteins are associated with invertebrates. The taxonomic distribution of vertebrate proteins in the Swiss-Prot database is 50% mammal, 25% fish, 10% bird, 9% amphibian, and 6% reptile.²⁴⁴ This can be compared to 55% mammal, 27% bird, 8% amphibian 6% fish, and 4% reptile, thus bird proteins are overrepresented in the matched proteins and fish proteins are underrepresented. This is consistent with evolutionary reports that birds have a close phylogenetic relationship to crocodilians.²⁴⁵

A putative function was assigned to the *de novo* sequenced proteins identified based on the function of the matched proteins. The majority of the proteins identified in the alligator leukocyte are common to proteins identified in leukocytes of other class of animals.

A range of proteins related to the immune system were identified. The identified immune system proteins were fibrinectin, histone H4, Ig kappa chainV-1 region, Ig lambda chain V-III region, lysozyme C, acyl-CoA-binding protein, myeloid protein, cathepsin C, complement component c3, ubiquitin, and polyubiquitin. The protein ubiquitin was identified with 32% sequence coverage, the largest observed. This is consistent with previous reports demonstrating that ubiquitin is a highly conserved protein in eukaryotes.²⁴⁶ Ubiquitin has several functions such as the removal of damaged or unnecessary proteins and controlling other cellular processes such as antigen processing, apoptosis, immune response and inflammation.²⁴⁷ Although it has several functions, it is related to the immune system. A *de novo* sequenced protein matched lysozyme (*Chelonia mydas*) from residue 50 to 61. Lysozyme was one of the first antibacterial proteins isolated from insects and is also a component of the humans innate immune system.²⁴⁸ The complement C3 protein also plays a role in innate immunity in organisms^{248, 249} and plays a major role in the activation of the complement system, which helps to remove pathogens.²⁴⁹ A *de novo* sequenced protein matched histone (*Gallus gallus*), a class of proteins that are involved in DNA packaging; however, they have been reported to behave as antimicrobial agents.²⁵⁰

Several cytoskeletal proteins, myosin alkali light chain, myosin regulatory light chain, β -actin, actin-2, actin-1, annexin V, annexin I, ras-related protein O-krev, vimentin-4, talin 1, filamin-C, vinculin, microtubule-actin cross-linking factor 1, α -actinin-1, actin, plastin-2, and putative tropomyosin γ -3 chain-like protein were identified. These were the major group of proteins identified and play an important role in the function of leukocytes. Actin is the most

abundant protein in all eukaryotic cells²⁵¹ and is responsible for muscle contraction and cell motility and works in conjunction with myosin protein through actin-myosin interactions.^{251, 252} Examples of cell movements include invasion of tissue by white blood cells to fight infection and migration of cells involved in wound healing.²⁵² Actin has multiple functions but is also related to the immune system; it regulates the integrin function, which is important for inflammation and immunity. Other identified cytoskeletal proteins, talin, α -actinin, plectin-2, and filamin-C, also play a role in the actin cytoskeletal system.²⁵³⁻²⁵⁶ Variations of myosin and tropomyosin proteins work along with actin to assist in phagocytosis of microorganisms.²⁵⁷ Annexins have various functions such as regulation of membrane organization and of Ca^{2+} concentrations within cells as well as participation in phagocytosis.²⁵⁸ Vinculin, another cytoskeleton protein, is necessary for cell movement and changes in cell shape.²⁵⁹ Vimentin, a filament protein, is involved in structural processes such as wound healing and is also found to be secreted by macrophages.²⁶⁰ Also identified in this study was a ras-related protein; these proteins are GTPases that are a part of the Ras superfamily of proteins and are involved in regulating cell behavior such as cell growth and action.²⁶¹

A class of enzymes known as endonucleases was also identified with high Mowse scores. These enzymes are commonly found in bacteria and behave like an immune system: bacteria use these enzymes to cut DNA from foreign material destroying them.²⁶² Although endonucleases are typically found in bacteria, alligator leukocyte may contain proteins that are similar to endonuclease protein and play a similar role. Alkaline phosphatase is a key enzyme involved in the process of dephosphorylation. It is also found to be involved in the protection against endotoxins, which may be lethal after an infection.²⁶³

Three matched proteins are hypothetical proteins from the western clawed frog, (*Xenopus tropicalis*), house mouse (*Mus musculus*), and chimpanzee (*Pan troglodytes*). The *Pan*

trogodyte's hypothetical protein had four matching peptide sequences suggesting that a similar protein is expressed in the alligator leukocyte. However, the function of these hypothetical proteins is not known and their relationship to the immune system cannot be determined.

There were proteins identified from the Mascot search that had a score greater than 57 but only one matching peptide: lysozyme C and ras-related protein O-krev (Table 3-2). These proteins cannot be identified with the same degree of confidence as the other proteins that had more than one peptide matching the protein. The proteins characterized as other have not been previously identified in leukocytes of other organisms. However, the E-values are high which suggests limited sequence identity and evolutionary relationship.

3.4 Summary

In the work described in this Chapter, proteins from the leukocytes of the American alligator (*Alligator mississippiensis*) were identified and their functions were characterized by adopting a general proteomics approach. A three-step strategy was performed to identify similar proteins in the gel bands and spots: *de novo* sequencing, Mascot search, and BLAST search. Mascot search results identified eighteen proteins with sequence similarity to the alligator leukocyte proteins. The identified proteins are common among eukaryotes and are associated with the immune system. The BLAST search results showed that 81% of *de novo* sequenced proteins matched similar proteins in the database. Among the identified proteins, the majority were related to the cytoskeletal system.

The BLAST search identified proteins most closely related to vertebrates. Proteins originating from birds, particularly chicken, were overrepresented in the matched proteins and correlated with immune related proteins reported for other species. Fish species was the least represented species among the matched proteins.

CHAPTER 4. SMALL MOLECULE INTERFERENCES IN *ALLIGATOR MISSISSIPPIENSES* LEUKOCYTE EXTRACT

In this chapter, experiments in which small molecule interferants in the leukocyte extracts from *Alligator mississippiensis* were separated and purified using reversed phase liquid chromatography are described. Seven bacterial species including *Escherichia coli*, *Enterobacter cloacae*, *Shigella flexneri*, *Klebsiella oxytoca*, *Salmonella*, *Staphylococcus aureus*, and *Staphylococcus epidermidis* were assessed for antimicrobial activity. The active fractions were characterized using tandem mass spectrometry and X-ray crystallography. The results from the mass analysis and X-ray crystallography suggest that the component exhibiting antimicrobial activity was ethylenediaminetetraacetate (EDTA), the anticoagulant agent used during the collection of the alligator blood. Tandem mass spectrometry data along with accurate mass measurements suggests that the other small molecule interferant is spermine.

4.1 Introduction

Antimicrobial drug resistance is a growing problem. Numerous bacteria such as *Staphylococcus aureus*, *Salmonella*, *E. coli*, *Pseudomonas aeruginosa*, and *Streptococcus pneumoniae* have developed resistance to antibiotics.²⁶⁴ One of the most resistant bacteria affecting hospital settings is *Staphylococcus aureus*: over 50% of its bacterial strains have become resistant to penicillin.^{264, 265} Due to this problem, new antibiotics with broad spectra of activity must be developed.

Studies have shown that when extracts of alligator leukocytes from blood are exposed to bacterial pathogens, the growth of the bacteria is inhibited.²⁶⁶ Several anticoagulant agents that prevent blood clotting are available, such as EDTA, heparin, citrate, and oxalate. EDTA is a chelating agent that synergistically inhibits growth of bacteria with antimicrobial agents²⁶⁷ such as sulfamethoxazole, trimethoprim,²⁶⁸ penicillin G, ampicillin, tetracycline, and

chloramphenicol.²⁶⁹ Studies have shown that EDTA can also inhibit bacterial growth independently.²⁷⁰ EDTA enhances the effectiveness of antibiotics against bacteria, particularly gram-negative bacteria, and chelates Ca^{+2} and Mg^{+2} which disrupts the bridge between them and lipopolysaccharide (LPS) in the bacterial outer membrane.^{267, 271}

Peptides and small molecules have the potential for therapeutic use. Small molecules are less costly and easy to synthesize and are stable, therefore longer half-life.²⁷² However, small molecules can potentially interact with multiple targets and accumulate in tissues causing toxicity.²⁷² Unlike small molecules, peptides are more specific and can be chemically modified to make the peptide less susceptible to proteolysis *in vivo*.²⁷²

Another small molecule that exhibits antimicrobial activity is spermine. Spermine is a biological polyamine that is found to be widely distributed in animal tissues.²⁷³ It has antibacterial activity, providing resistance to a wide range of microorganisms such as *S. aureus*, *S. albus*, and *B. anthracis*.^{273, 274} Spermine was first isolated from semen and exhibits antimicrobial activity against gram-positive microorganisms.²⁷⁵

In this chapter, we report that EDTA and spermine are present and isolated from leukocyte extracts of alligators. These molecules can potentially cause interferences with the activity measurements.

4.2 Experimental

Leukocyte extracts were obtained from the blood of alligators as described in Section 2.1. In this study, which was the first studies performed on isolation of peptides from alligator leukocytes, 0.1 mM EDTA was used as the blood anticoagulant.

A Bradford protein assay was used to determine the concentration of proteins in the alligator leukocyte extracts. A 2 mg/mL quantity of bovine serum albumin (BSA) was used to prepare five BSA dilutions from 125 to 2000 $\mu\text{g}/\text{mL}$. A 0.1% acetic acid buffer was used to

perform the dilutions. Each standard and leukocyte protein extract was pipette into clean test tubes and mixed with the protein dye reagent. The protein samples were assayed in triplicate and incubated for 5 min before UV-Vis analysis. Absorbance was measured at 595 nm for both the protein standard and leukocyte protein extracts. The concentration of the leukocyte protein mixture was determined using a standard Bradford curve.²⁷⁶

The leukocyte extract was tested for antimicrobial activity before chromatographic separation. A radial diffusion assay was used to monitor bacterial growth inhibition.²⁷⁷ Leukocyte extracts and chromatographic fractions of the leukocyte extracts were pipetted into wells on the activity plate, incubated overnight at 37°C, and zones of bacterial growth inhibition were measured the following day.

Alligator leukocyte extracts were separated by reverse phase liquid chromatography (RP-LC) and peak fractions were monitored with a photodiode array (PDA) detector and manually collected in 2 minute intervals. For PDA detection, the UV spectrum was recorded between 190 and 400 nm. The fractions were tested for antimicrobial activity and the active fractions were lyophilized and re-suspended in 0.01% (v/v) aqueous acetic acid. The fractions that exhibited antimicrobial activity were injected onto the C₁₈ reversed phase (RP) column using a gradient of 10–90% acetonitrile over 40 minutes for further purification and analyzed using ESI-TOF mass spectrometry.

The purified fraction that exhibited antimicrobial activity was submitted for structural characterization using X-ray crystallography. Tandem mass spectrometry (MS/MS) using the nano ESI-Q-TOF was also performed to obtain structural information on the molecules.

4.3 Results and Discussion

4.3.1 Separation of Leukocyte Extract

The alligator leukocyte was separated into 35 fractions using C_{18} reversed-phase chromatography (Figure 4-1). The fractions from the leukocyte extract were collected and the initial fractions collected within the first six minutes (labeled F2 and F3) were tested for antimicrobial activity.

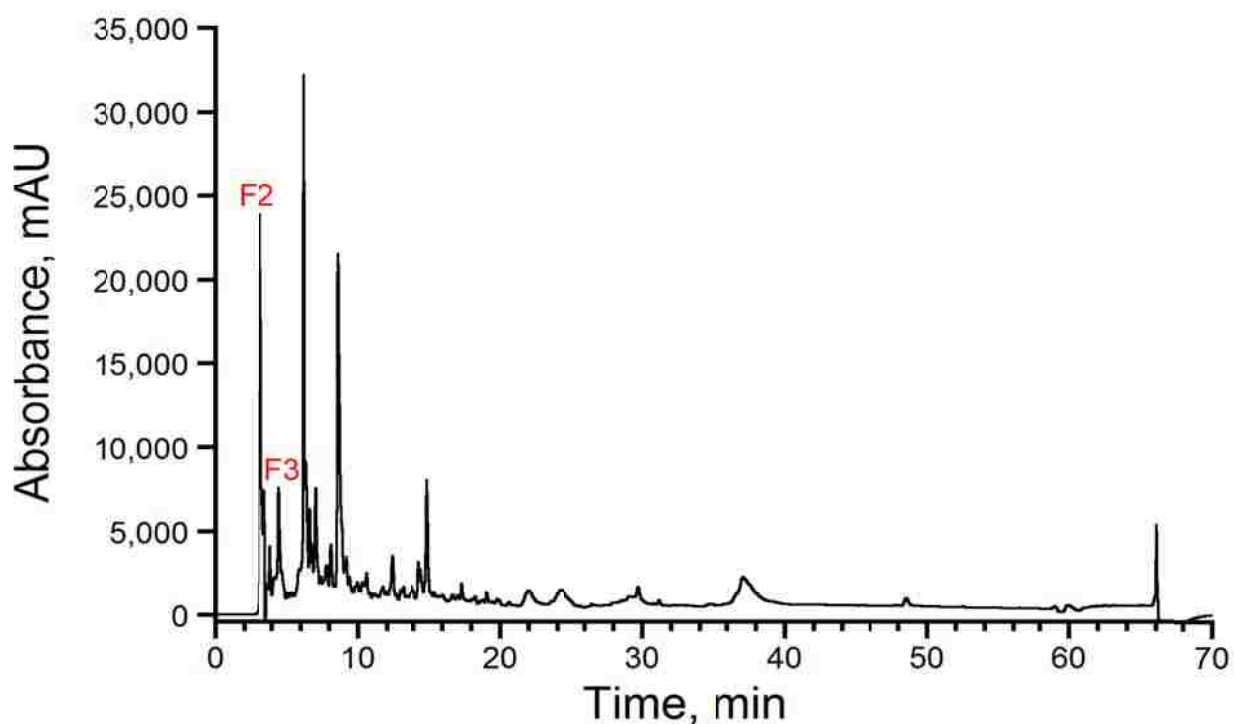


Figure 4-1. Separation of alligator leukocyte using reversed-phase separation and monitored between 190 and 400 nm.

The collected fractions F2 and F3 were further purified using RP chromatography as seen in Figures 4-2 (A) and (B).

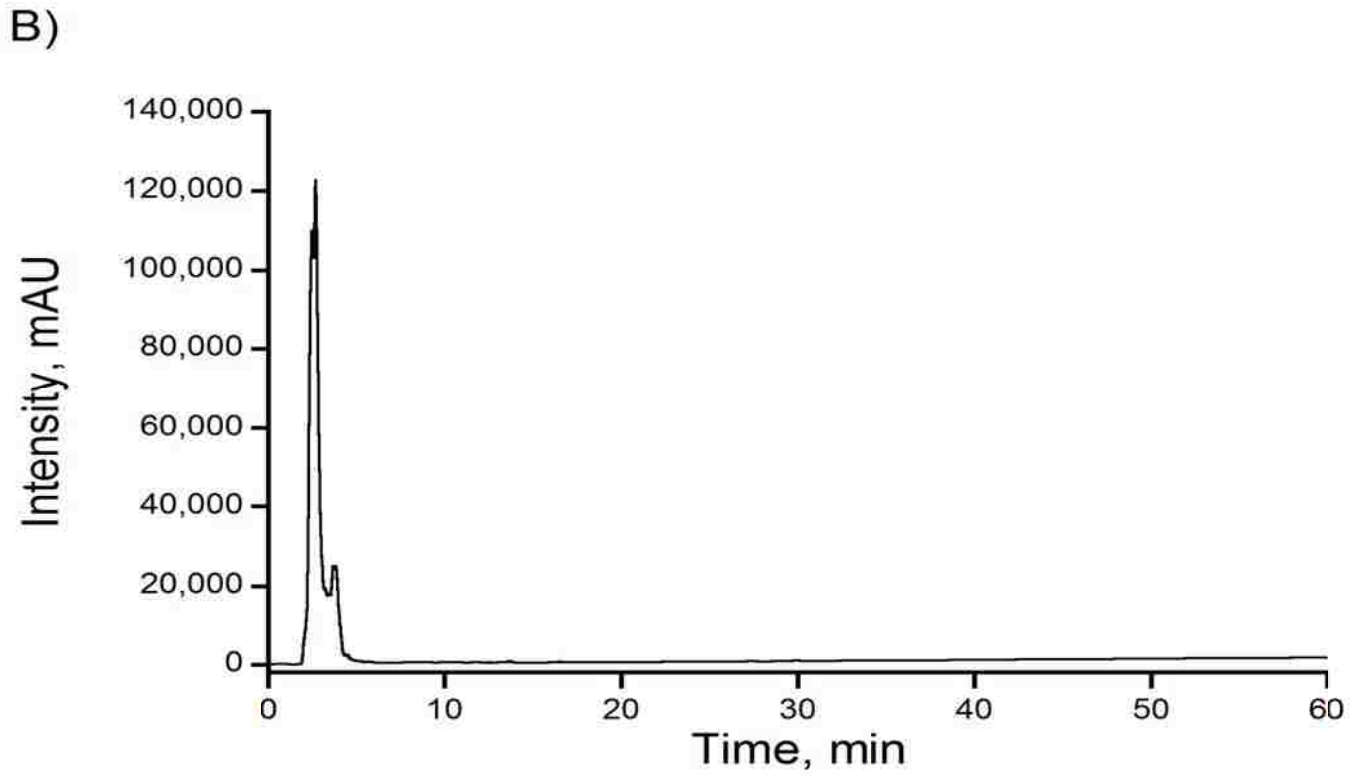
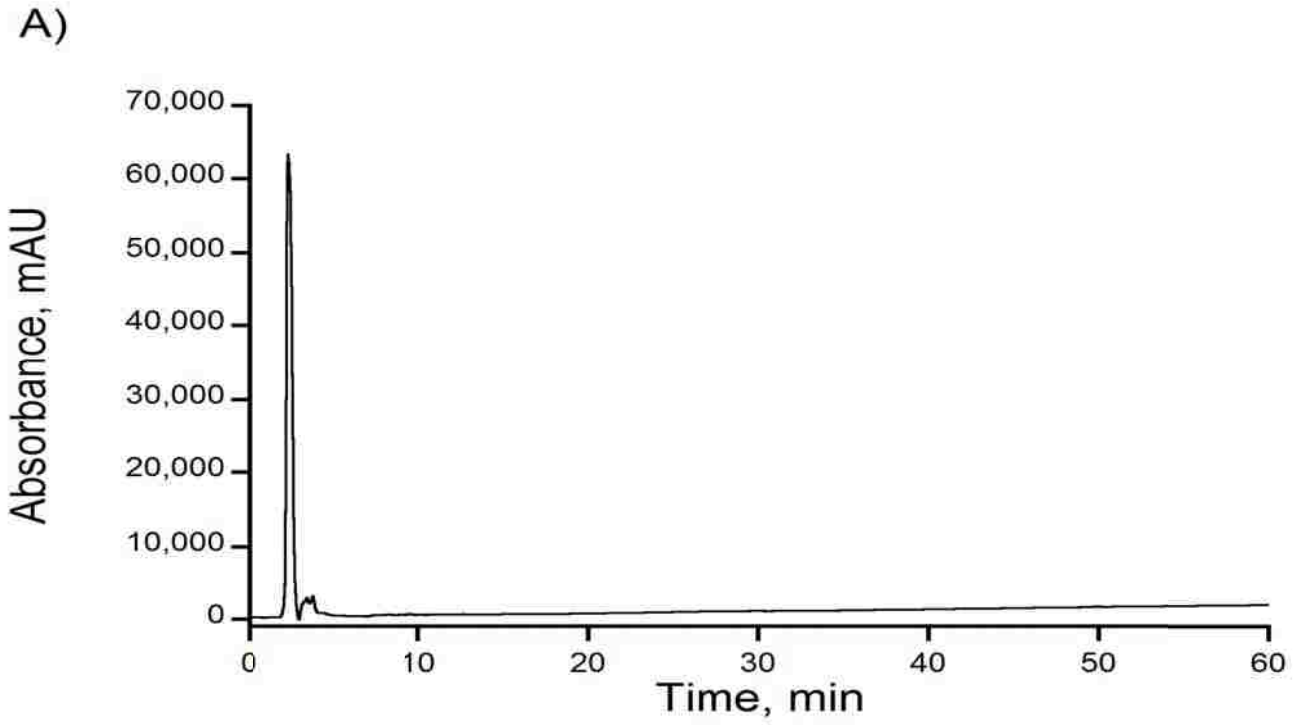


Figure 4-2. (A) Purification of Fractions 2 and (B) Fraction 3 using reversed-phase separation.

4.3.2 Antimicrobial Activity

Results from this study showed that Fractions 2 and 3 exhibited antimicrobial activity against *Escherichia coli*, *Shigella flexneri*, *Salmonella*, *Staphylococcus aureus*, and *Staphylococcus epidermidis* (Table 4-1). When tested against *Klebsiella oxytoca*, a gram-negative bacterium, neither of the fractions exhibited activity. However, growth inhibition was observed for Fraction 2 against *Enterobacter cloacae*, a gram-negative bacterium, but not for Fraction 3. The largest antimicrobial activities from Fractions 2 and 3 were observed for *Staphylococcus epidermidis*, a gram-positive bacterium; zones of inhibition were 17 and 14 mm respectively. The lowest activity was observed in *Salmonella*, a gram-negative bacterium, at 5 and 4 mm respectively.

Table 4-1. Antibacterial activity of Fractions 2 and 3 from leukocyte extract against bacterial species tested using radial diffusion assay.

Bacterial Strain	Zone of Inhibition (mm)	
	(F2)	(F3)
<i>Escherichia coli</i>	10.0	5.0
<i>Enterobacter cloacae</i>	7.0	0
<i>Shigella flexneri</i>	6.0	5.0
<i>Klebsiella oxytoca</i>	0	0
<i>Salmonella</i>	5.0	4.0
<i>Staphylococcus aureus</i>	15.0	10.0
<i>Staphylococcus epidermidis</i>	17.0	14.0

Both fractions exhibited antimicrobial activity towards both gram-negative and gram-positive bacteria; however growth inhibition was highest for gram-positive bacteria.

The negative control, 0.01% acetic acid, showed no antibacterial activity and the positive controls, polymyxin B and synthesized EDTA and spermine showed antibacterial activity against the tested microbes.

4.3.3 ESI-MS Analysis

ESI-MS was used to analyze fractions F2 and F3 as seen in Figure 4-3 (A) and (B). Because of the high mass accuracy of the ESI-TOF instrument it was possible to assign chemical formulas to the peaks present in the mass spectrum. The ion at m/z 203.2250 was observed in both fractions F2 and F3 and was the most abundant ion present. The molecular formula, $C_{10}H_{26}N_4$, for the molecule measured at m/z 203.2250 was confirmed with a mass accuracy of 6 ppm.

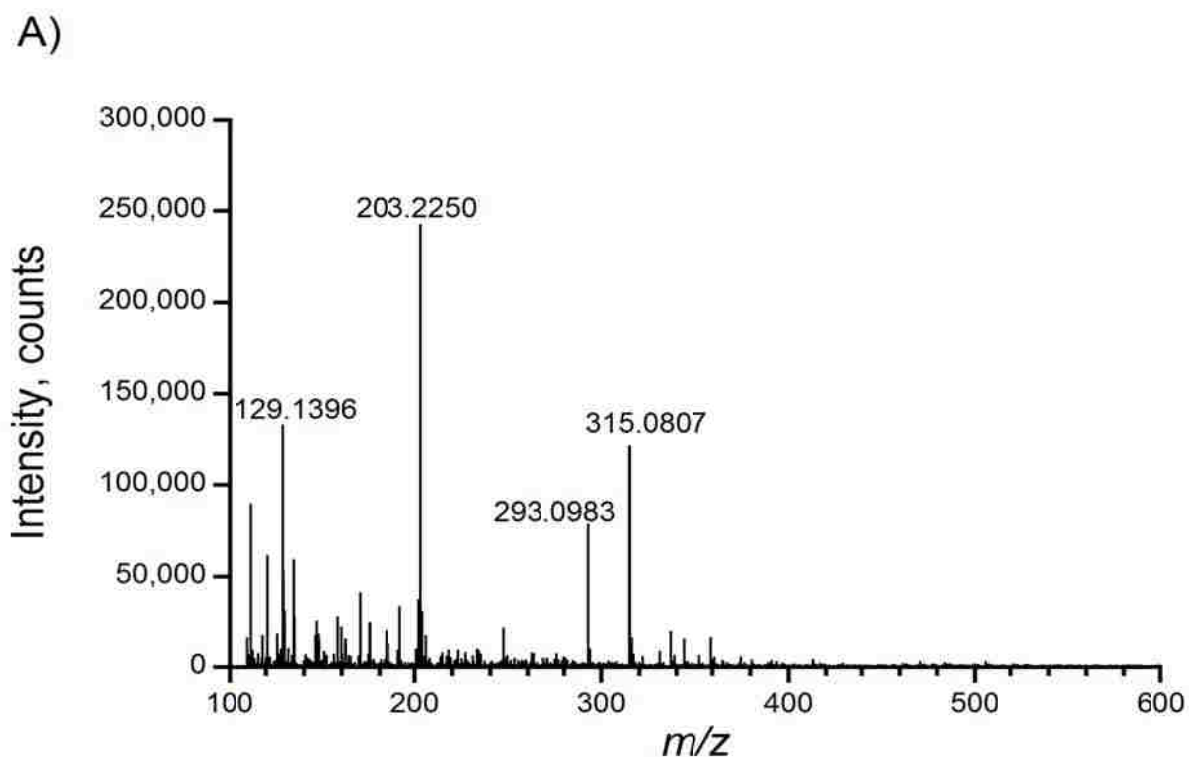
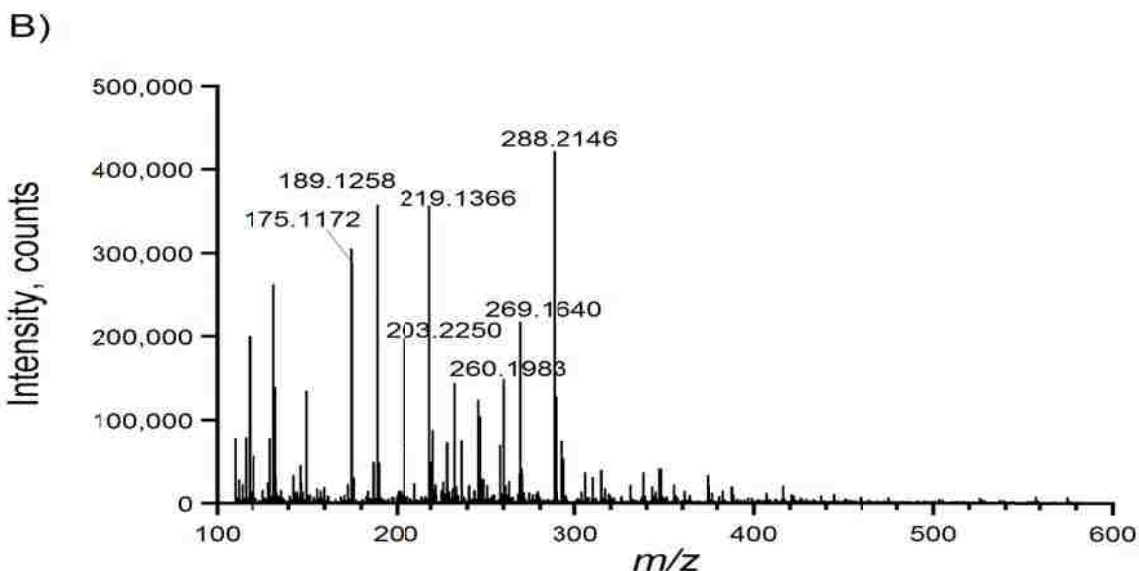


Figure 4-3. ESI-MS of fractions (A) F2 and (B) F3 isolated from alligator leukocyte extract.

Figure 4-3. cont'd.



Another intense peak in Fraction 2 was at m/z 315.0807 with a mass accuracy of 3 ppm, corresponding to molecular formula $C_{10}H_{16}N_2O_8Na$. However, this molecule was absent in Fraction 3. This suggests that this molecule could have contributed to the antimicrobial activity of Fraction 2 because growth inhibition was observed against *Enterobacter cloacae* from Fraction 2 but not from Fraction 3.

Previous studies have shown that antimicrobial components isolated from plants, insects and mammals are often peptides with 12–60 amino acids and have masses less than 10 kDa.²⁴ The results from this study show that the antibacterial properties of the fractions are either 1) produced by small molecules with masses less than 400 Da, 2) produced by larger molecules that are present at concentrations below the mass spectrometer detection limit, or 3) produced by small and large molecules acting synergistically (as noted above) with the larger molecule not detected in the mass spectrometry analysis.

Tandem MS of the small molecules identified from MS analysis was performed to identify their structure. Figure 4-4 shows the MS/MS data for the product ions 315.1 and 203.2Da.

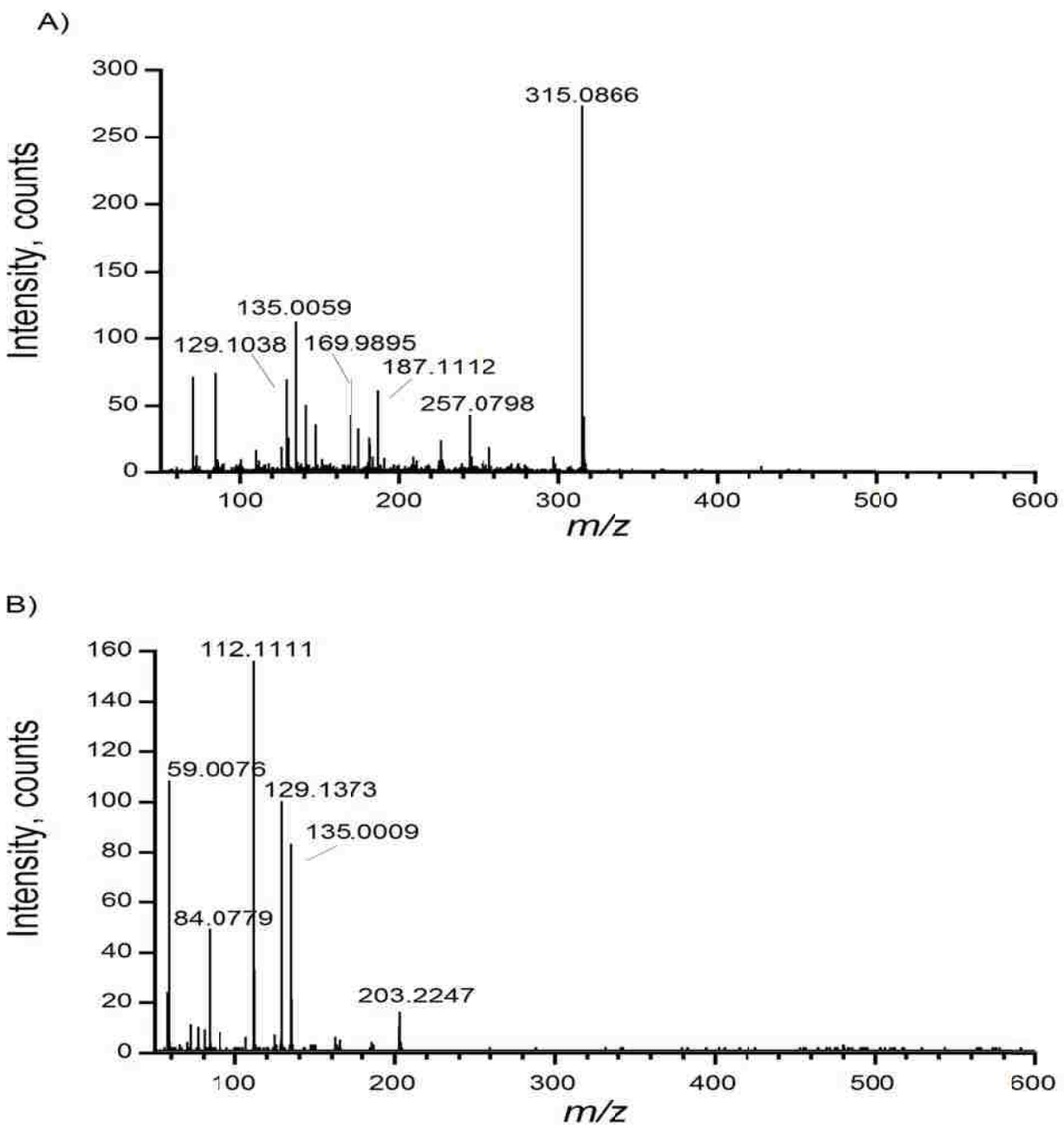


Figure 4-4. Tandem mass spectrometry data of (A) the m/z 315.1 ion from Fraction 2 and (B) the 203.2 ion from Fractions 2 and 3.

The MS/MS results for the ion at m/z 203.2250 showed fragments previously reported for spermine, $C_{10}H_{26}N_4$, by Q-TOF analysis.²⁷⁸ These fragments are based on the cleavages of the carbon and nitrogen chain, as seen on Figure 4-5.

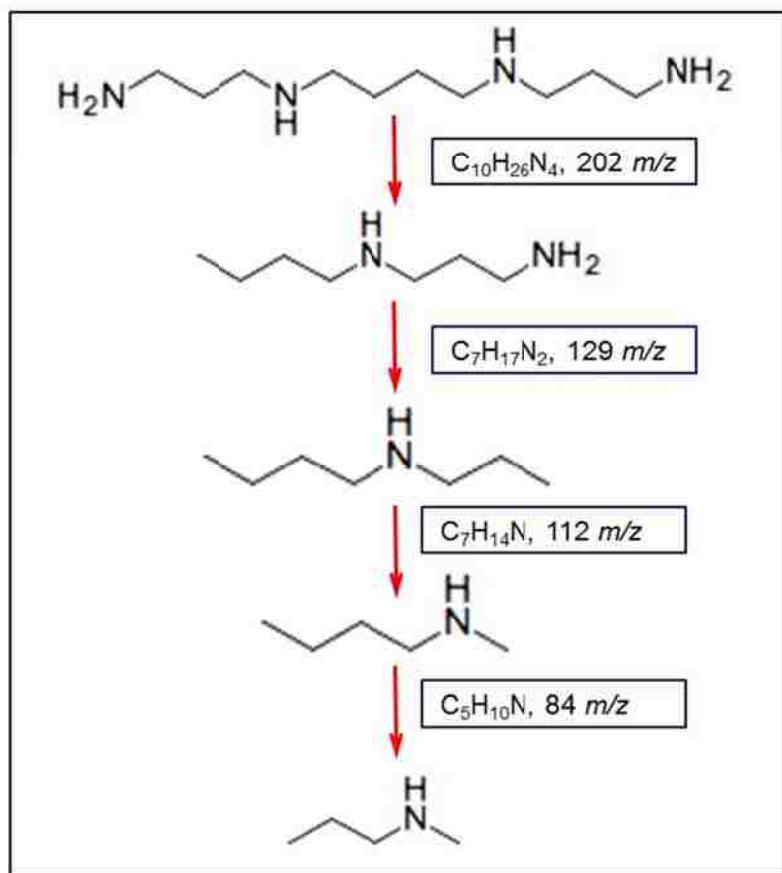


Figure 4-5. Fragmentation of spermine.

The fragments observed from the ion at m/z 203.2250 in Figure 4-4 (B), for example 84.1 112.1, and 129.1, are characteristic of the fragmentation pattern of spermine ion. The tandem MS data are consistent with the molecular formula obtained from the MS accurate mass results. Results from the MS/MS spectrum of the ion at m/z 315.1 ion did not allow for structure assignment.

4.3.4 X-ray Crystallography

Fraction 2 was dried and formed crystals that were analyzed using X-ray crystallography. The crystallographic data was searched in the Cambridge Structural Database for organic and organometallic compounds and eight hits of possible structures were generated, including phenylammonium nitrate ($C_6H_8N_2O_3$), EDTA disodium salt ($C_{10}H_{18}N_2Na_2O_{10}$)ⁿ, 3a,4,7,7-Tetramethyl- (3aR,4S,6aR,10aS)-perhydronaphtho(8a, 1-b)furan-2-one ($C_{16}H_{26}O_2$), 7-Iodo-2,9-dimethyl-4,4-ethylenedioxy-11-oxatricycloundec-2-ene ($C_{14}H_{19}I_1O_3$), 7-Methoxy-4,9,13-trimethyl-6-oxa-1,13:10,11-diepoxytricyclotridec-3-en-5-one ($C_{16}H_{20}O_5$), catena-((μ_2 -Benzene-1,3-dicarboxylato)-aqua-(propane-1,3-diamine)-copper(ii)monohydrate) ($C_{11}H_{16}Cu_1N_2O_5$)_{n,n}(H₂O)), catena-(tetra-sodium bis(μ_2 -methylenediphosphonato)-di-copper(ii)dehydrate) (($C_2H_4Cu_2O_{12}P_4^{4-}$)_n,4n(Na¹⁺),2n(H₂O)), and 3-(4-Methoxybenzylidene)-2-oxo-2,3-dihydro-1H-imidazo(1,2-a)pyridiniumbromide ($C_{15}H_{13}N_2O_2$). Based on the crystal structure, seen on Figure 4-6, and the crystal's space group values, the structure was a best match for EDTA disodium salt.

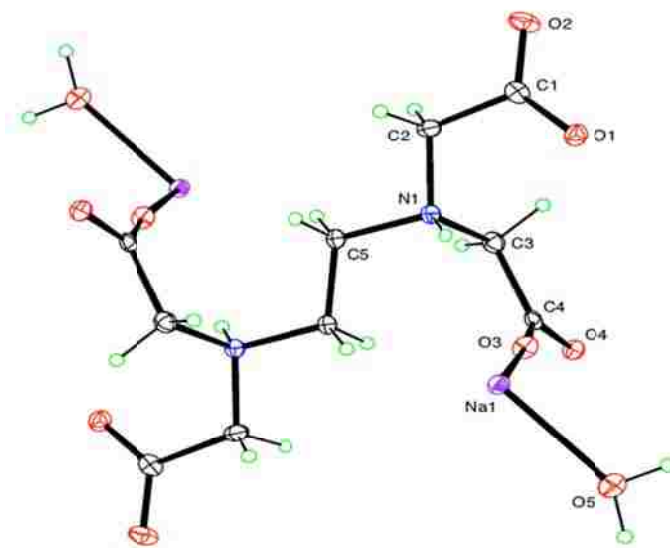


Figure 4-6. Crystal structure of EDTA disodium salt obtained from alligator leukocyte extract.

4.4 Summary

In the work described in this Chapter, leukocyte extracts were subjected to purification and characterization by LC ESI-MS, MS/MS and X-ray crystallography analysis. The results from this study suggest that EDTA, a blood anticoagulant, in the presence of a protein might contribute to inhibition of bacterial growth. In addition, previous studies have shown that EDTA contributes to bacterial growth inhibition at both high and low concentrations.²⁷⁰ An alternate explanation is that the antimicrobial activity resulted from a molecule that was not detected in the mass spectrometry analysis. RP separation was capable of reducing the complexity of the leukocyte protein mixture; however, the chromatographic fractions were still not sufficiently pure to identify the specific molecules that contributed to antimicrobial activity. Additional off-line RP separations assisted in small molecule purification and antimicrobial activity was identified for fractions collected within the first six minutes. The masses of the small molecules detected in the active fractions were measured by ESI-MS and the small molecules were identified by empirical formula calculations based on accurate mass measurement. A molecule, from the active fractions, F2 and F3 at m/z 203.2 had the formula, $C_{10}H_{26}N_4$ with 6 ppm mass error. In addition, tandem MS analysis revealed fragmentation patterns for this ion that matched the molecule spermine. Another molecule from fraction F2 at m/z 315.1 (3 ppm accuracy) had the formula $C_{10}H_{16}N_2O_8Na$ and X-ray crystallography was used for further characterization where results from the structural database identified the structure from the active fraction as EDTA disodium salt.

The results from this study suggest that EDTA may not be a suitable anticoagulant agent for isolation of blood proteins when testing for antimicrobial activity. As a result, for future studies the anticoagulant agent heparin was used as a substitute. The other small molecule that was identified was spermine, which is typically found in all animal cells and is usually excreted

from semen and pancreatic tissue. Blood contains various proteins that are secreted or shed from cells and tissues throughout the body.⁷⁸ Therefore, it may not be possible to eliminate spermine as an interferant, hence for the studies described in the following chapters, small molecules eluting within the first six minutes from the leukocyte extracts were not considered as potential candidates for antimicrobial molecules.

CHAPTER 5. ISOLATION AND *DE NOVO* SEQUENCING OF ANTIMICROBIAL PEPTIDES FROM LEUKOCYTES OF THE AMERICAN ALLIGATOR (*ALLIGATOR MISSISSIPPIENSIS*)

The goal of the research in this chapter was to isolate novel peptides with antimicrobial activity from American alligator blood (*Alligator mississippiensis*). Reversed-phase high performance liquid chromatography (RP-HPLC) was used to separate and purify peptides from leukocyte extracts and the HPLC fractions were checked for activity. A partially purified fraction containing two major peptides at 4.7 and 4.9 kDa was tested for antimicrobial activity against *E. coli*, *E. cloacae*, and *K. oxytoca*. The peptides were analyzed using ion mobility spectrometry mass spectrometry. Matrix assisted laser desorption ionization time-of-flight/time-of-flight mass spectrometry (MALDI-TOF/TOF) was used to generate MS/MS data for the peptides and *de novo* sequencing was used to determine their sequence. The 4.7 kDa peptide was aligned to other antimicrobial peptides and showed 36% sequence identity with Pilosulin 4 and the 4.9 kDa peptide showed 33% sequence identity to SpStrongylocin 2.

5.1 Introduction

Increasing bacterial resistance to antibiotics has become the driving force behind the development of new anti-infectives to replace or supplement conventional antibiotics.²⁷⁹ Antimicrobial proteins and peptides from natural sources show promise as new antibiotics. Antimicrobial peptides (AMP) are important components of the innate immune system that have been isolated in plants,²⁸⁰ vertebrates,²⁸¹ and invertebrates²⁸² and have potential for pharmaceutical use. Most antimicrobial peptides are small (<10 kDa), cationic, lysine and arginine containing molecules; however, some anionic peptides have also been reported.²⁵ Cationic proteins and peptides have an affinity for microbe cell surfaces, which have negatively charged groups. This attraction is dependent on electrostatic interactions and their tertiary

structure.²⁸³ Cationic proteins and peptides vary in their size, structure, and mechanism of action.²⁸³

In some vertebrates such as mammals and reptiles, antimicrobial proteins and peptides are found in the blood, particularly in the neutrophils. There are several classes of antimicrobial proteins and peptides associated with neutrophils, such as lactoferrin, cathelicidins, calprotectin and defensins.²⁸³ Defensins are small cationic peptides around 4 kDa with a broad antimicrobial activity against bacteria, fungi, viruses, and parasites.²⁸⁴ β -defensins serve as effector molecules of innate immunity, providing an initial defense against infectious agents,²⁸⁵ and have been identified in bovine neutrophils,²⁸⁶ turtle³³ and avian leukocytes,²⁸⁷ and human plasma.²⁸⁸ Leukocytes in the blood of crocodylian species have a range of antimicrobial, antifungal, and antiviral activity,^{1, 289, 290} but little is known about antimicrobial peptides in *Alligator mississippiensis*.

There is limited information about reptile antimicrobial peptides. To date there are very few reptilian antimicrobial peptide sequences identified among the nearly 400 that have been reported from other sources.^{33, 36} The *Alligator mississippiensis* leukocytes have a broad range of activity towards bacteria, viruses and fungi.¹ Some studies have been performed to characterize the proteins and peptides from alligator leukocytes;²⁹¹ however, none have been sequenced.

De Novo sequencing is a common approach for mass spectrometry based sequencing of peptides and proteins that are not in a protein sequence database. *De Novo* sequencing entails identifying peptide sequences without any previous knowledge of the protein sequence.¹⁶⁴

In this study, we isolated two novel peptides that exhibit antimicrobial activity from the leukocytes of the American alligator (*Alligator mississippiensis*) using reversed-phase chromatography separation. The peptides were sequenced using *de novo* sequencing and compared to other known antimicrobial peptides.

5.2 Experimental

5.2.1 Isolation and Extraction of Antimicrobial Peptides

Blood was drawn from farm and wild alligators ranging in size from 5 to 12 feet for isolation of leukocytes using an approach described in Section 2.1.

5.2.2 Microbial Strains

All bacterial strains were obtained from American Type Culture Collection (ATCC). The following bacterial strains were used: *Escherichia coli* (ATCC 43827), *Enterobacter cloacae* (ATCC 23355), and *Klebsiella oxytoca* (ATCC 33496).

5.2.3 Antimicrobial Activity

The antimicrobial activity of the leukocyte extracts and the peptide fractions from HPLC eluents were determined using an antimicrobial assay.²⁷⁸ Trypticase soy broth containing 1% agarose was sterilized and 15 mL aliquots were transferred to centrifuge tubes. The nutrient broth was inoculated with 20 μ L of a log phase culture of bacteria and added to petri dishes. Wells were cut into the petri dish using a sterilized pipette with a vacuum line attached for suction. Five microliters of leukocyte extract dissolved in aqueous 0.1% acetic acid (v/v) was added to each well and allowed to diffuse at room temperature for 3 hours. The plates were then incubated at 37°C overnight and the zones of bacterial growth inhibition were measured. The antibiotic polymyxin B at 2×10^{-5} M was used as a positive control and aqueous 0.1% acetic acid (v/v) was used a negative control.

5.2.4 Peptide Separation

Leukocyte extract in 0.1% acetic acid was centrifuged to remove any insoluble residue. A 10 μ L volume of concentrated sample was injected with an autosampler system onto a 0.3×1 mm trapping column on a LC system (Agilent) at a flow rate of 0.3 ml/min. The sample was eluted onto a 2.1×30 mm, 3.5 μ m particle size C₁₈ column (Zorbax) and separated using a

gradient of 5 to 60% B (80% acetonitrile (v/v) and 20% water (v/v) containing 0.1% formic acid (v/v)) over 70 min with a flow rate of 0.3 mL/min. The eluent was monitored by UV absorbance from 190 to 480 nm. A fraction collector was used to collect the eluent at timed intervals for further characterization and activity studies.

5.2.5 Ion Mobility Mass Spectrometry

The peptide samples were diluted in 0.1% formic acid and injected on the nanoAcquity ultra pressure liquid chromatography (UPLC) system at a flow rate of 500 nL/min. The peptides were separated using a gradient of 3–40% B over 30 minutes. The peptides were ionized using a nano-electrospray source, operated in positive ion mode and analyzed via a quadrupole mass analyzer before being introduced to the ion mobility spectrometry (IMS) on the ESI-Q/IMS/TOF MS. The quadrupole was operated in RF mode in order to record full mobility spectra and the data was processed using MassLynx software. A pressure of 0.5 bar of nitrogen was maintained in the ion mobility (IM) cell.

5.2.6 MALDI Mass Spectrometry

Alligator leukocyte fractions were mixed on a stainless steel plate using a 1:1 ratio of 1 μ L sample and 1 μ L saturated matrix including α -cyano-4-hydroxycinnamic acid (CCA) and 2-hydroxy-5-methoxybenzoic acid (sDHB) matrix solution containing 0.1% v/v TFA for molecular mass determination. The sample and matrix solution was air-dried at room temperature and analyzed in TOF mode on the MALDI TOF/TOF MS. Mass spectra were recorded in reflector positive ion mode with external calibration using a standard calibration mixture. Spectra were 500 shot averages collected using a mass range from 420–5000 Da. Masses were determined using FlexAnalysis software. MS/MS data was obtained by LIFT/CID fragmentation. MS/MS experiments were carried out in reflectron mode and more than 5000 laser shots were collected. The MS/MS data were processed in FlexAnalysis.

5.2.7 ESI Mass Spectrometry

The fraction containing the two peptides of interest was also analyzed on the ESI-QTOF. The peptide sample was diluted twofold with 50% ACN solution containing 0.1% FA (v/v). MS/MS experiments were performed on the selected parent ion and the collision energies were varied from 25 to 60 eV in 5 eV intervals.

5.2.8 *De Novo* Sequencing and Sequence Analysis

De Novo sequencing of the peptide fragments generated using MALDI MS/MS was performed with BioTools. The *de novo* sequencing results were verified manually. The peptide sequences produced by *de novo* sequencing were aligned with β -defensins peptide sequences from other vertebrates for sequence comparison.

The alligator antimicrobial peptide sequence was compared to other antimicrobial peptides in the Antimicrobial Peptide Database.²⁹² The alligator antimicrobial peptide was also aligned to find the most similar peptides in the database.

5.3 Results and Discussion

5.3.1 Antimicrobial Activity of Leukocyte Extracts

The antimicrobial activity of the alligator leukocyte extracts was tested against three species of bacteria. The leukocyte extracts exhibited antimicrobial activity against gram negative bacterium *E.coli*, *K. oxytoca*, and *E.cloacae* with approximately 0.5 mm zone of inhibition.

5.3.2 Peptide Purification

Several RP separations were run to separate and collect the active peptides. The separation of the peptide and protein mixture from the alligator leukocytes produced more than 12 fractions (Figure 5-1). The fraction collected at 15 min at 20% acetonitrile showed antimicrobial activity against the tested bacteria. The deconvoluted masses from the ESI data

(Figures 5-1B and 5-1C) showed at least three peptides present in the active fraction at 4618.3, 4745.4, and 4961.9 m/z .

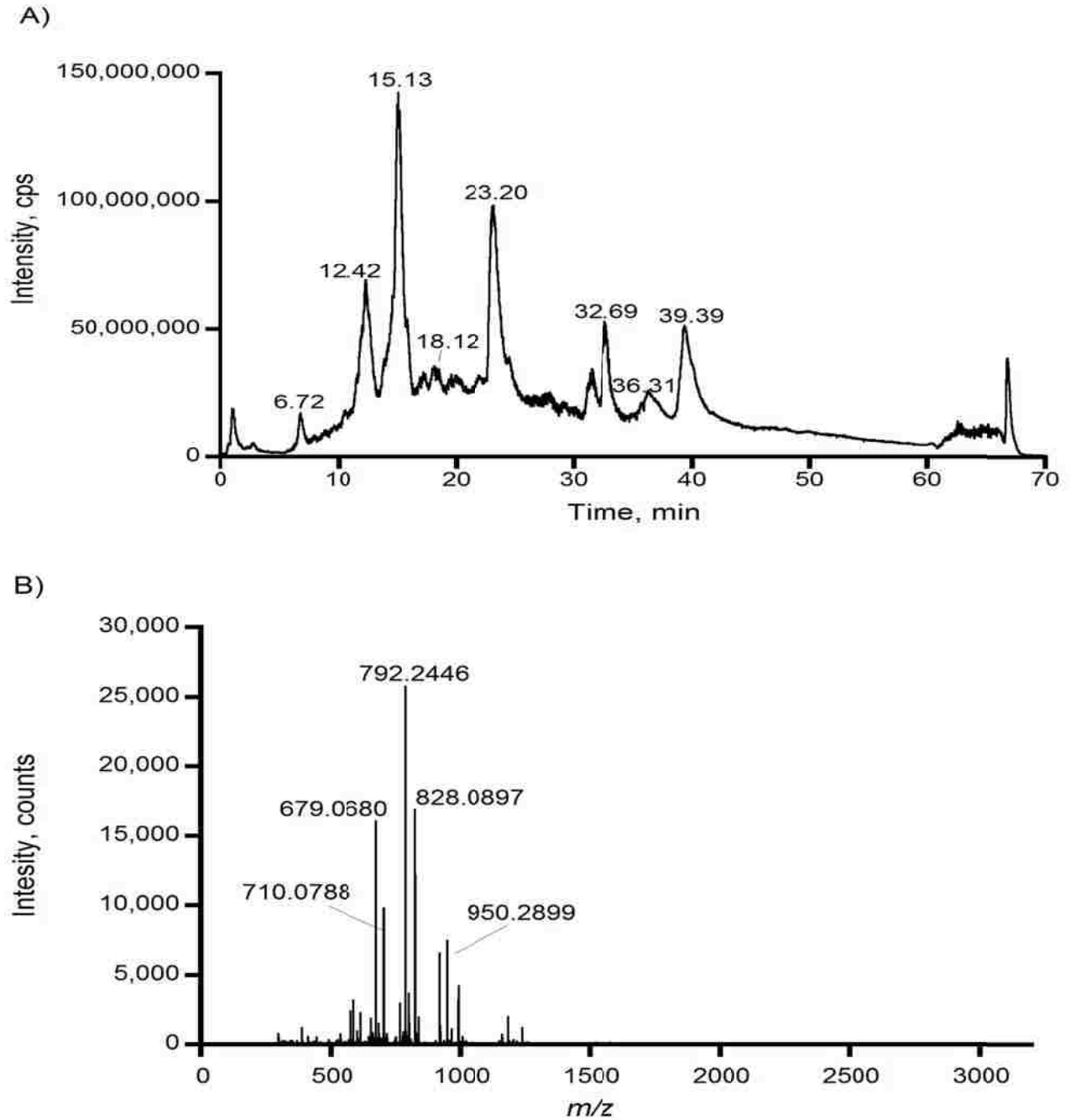
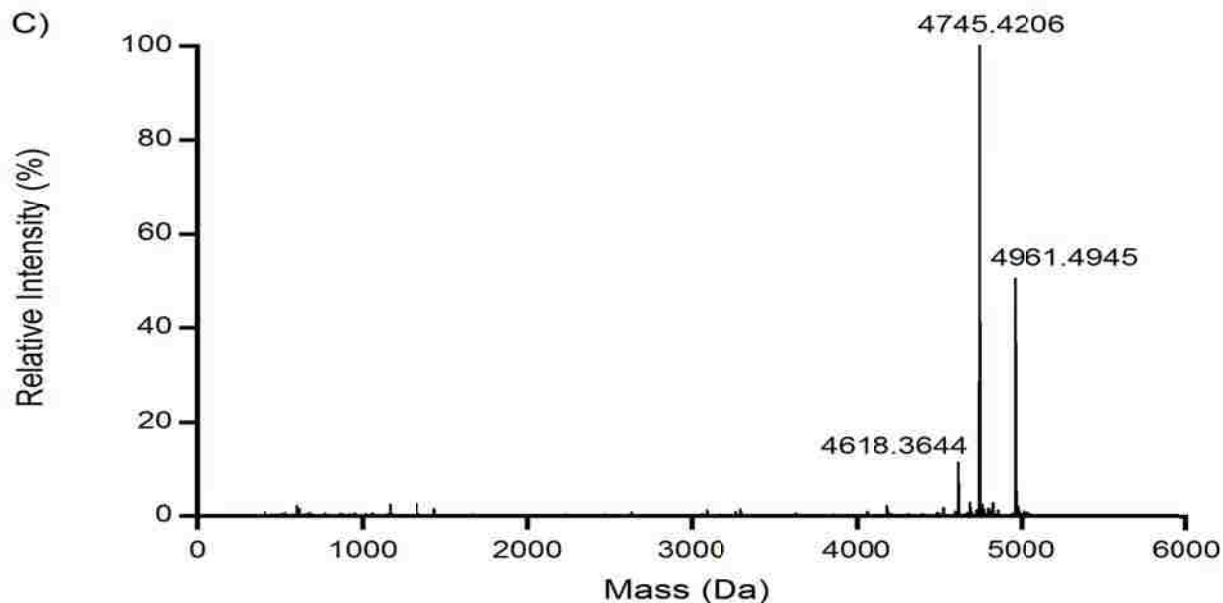


Figure 5-1. (A) The leukocyte extract in 0.1% acetic acid was separated using RP-HPLC on a C₁₈ Vydac column. (B) Mass spectrum of the biologically active fraction eluted at 20% acetonitrile. (C) Deconvoluted mass spectrum of the peptides eluted at 15% acetonitrile.

Figure 5-1. cont'd.



In addition to the major peptides identified in the active fraction, there were other minor impurities in the lower mass range which may suggest the presence of other antimicrobial peptides in lower quantities.

5.3.3. Ion Mobility Mass Spectrometry of the Active Peptides

The two peptides from the alligator leukocyte extract at 4745.4 and 4961.9 Da have a mass difference of ~200 Da; therefore, nano liquid chromatography-ion mobility spectrometry mass spectrometry (LC-IMS) was used to separate them based on their charge, size and shape. The base peak chromatogram seen in Figure 5-2 corresponds to multiple peptides with similar retention times, indicating that these two peptides coelute. Figures 5-3 and 5-4 show plots of m/z versus drift time and m/z versus retention time of the ions separated by ion mobility.

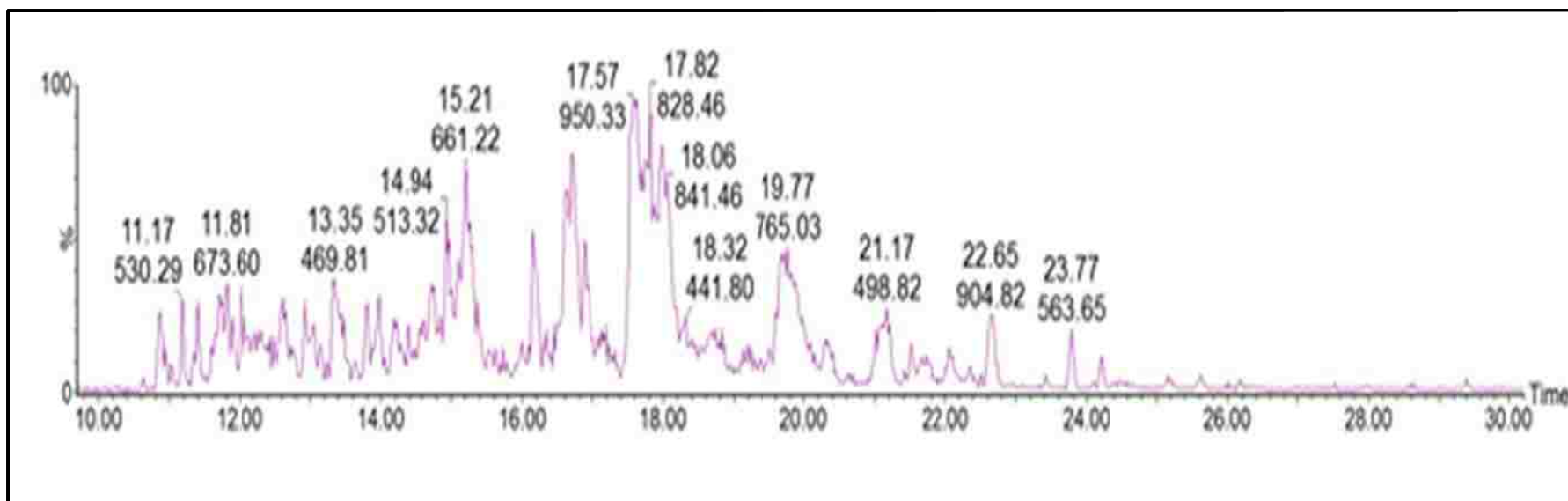


Figure 5-2. Base peak chromatogram of the nanoflow LC-IMS experiment for a mixture of peptides from alligator leukocyte extracts. The values above the peaks represent both the elution time and m/z value for ions measured at the respective times.

(A)

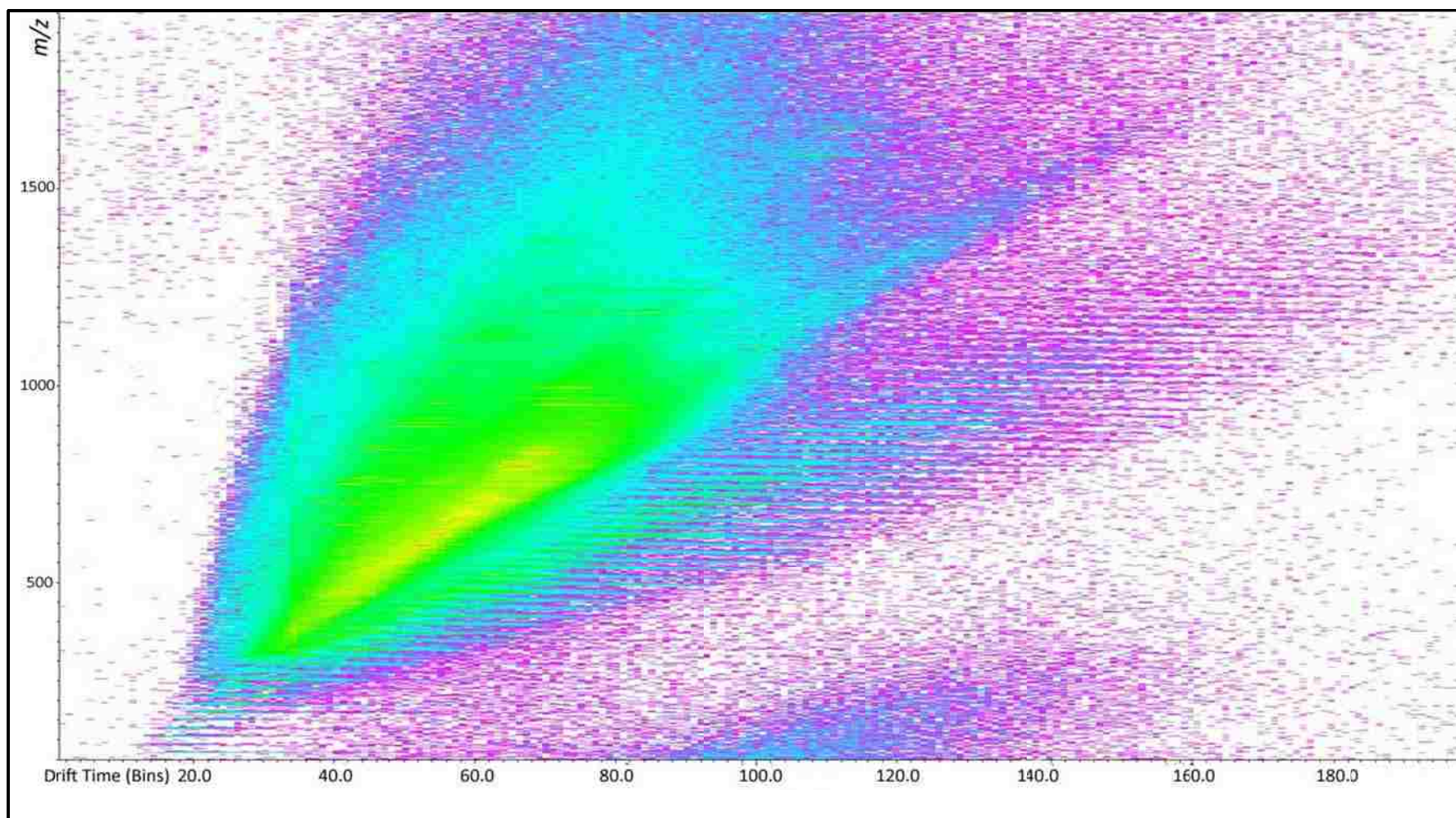
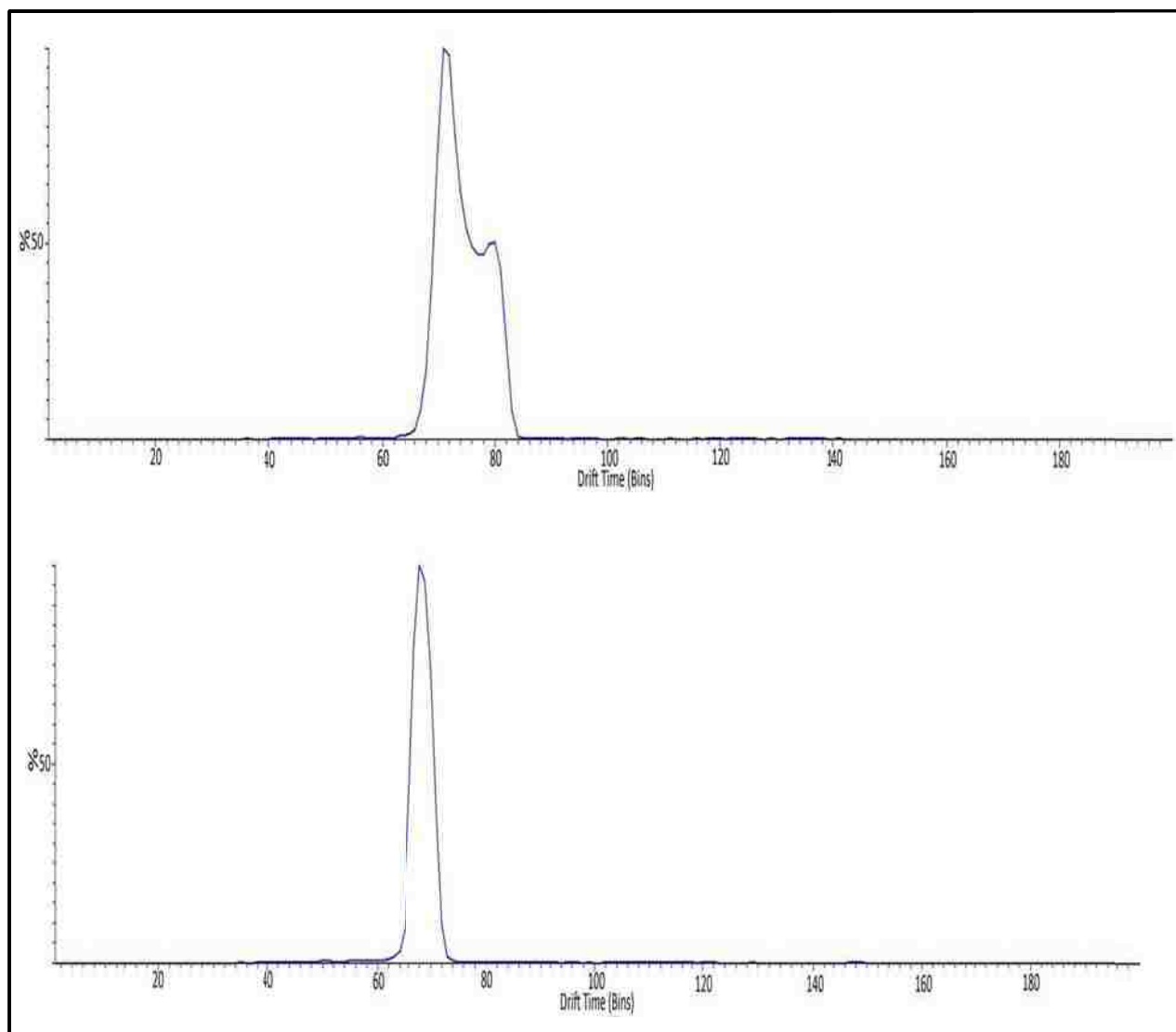


Figure 5-3. (A) 2D Plot of m/z versus ion arrival time for active peptides from leukocyte extract. (B) Ion mobility drift times.

Figure 5-3. cont'd.

(B)



The two peptides, from the RP fraction exhibiting antimicrobial activity, with similar retention times between 17.5 and 18 min (seen in Figure 5-4B and C) were separated in the IM cell based on their ion mobility drift times as seen in Figure 5-4A. The peptide ions were measured at $m/z = 827.8$ with a charge state of +6 and $m/z = 949.9$ with a charge state of +5. These results indicate that these two peptides can be separated on the basis of their net charge.

(A)

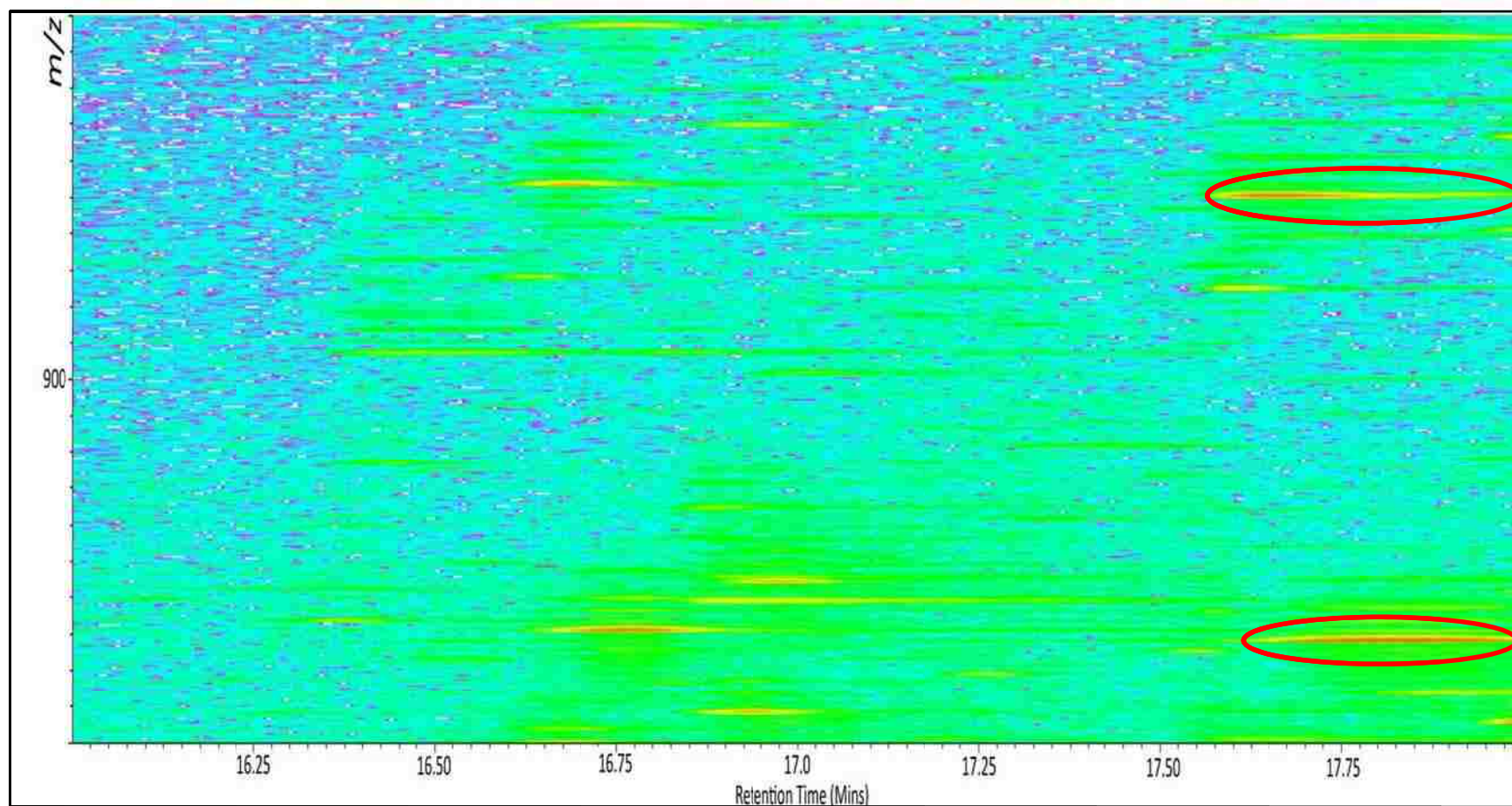
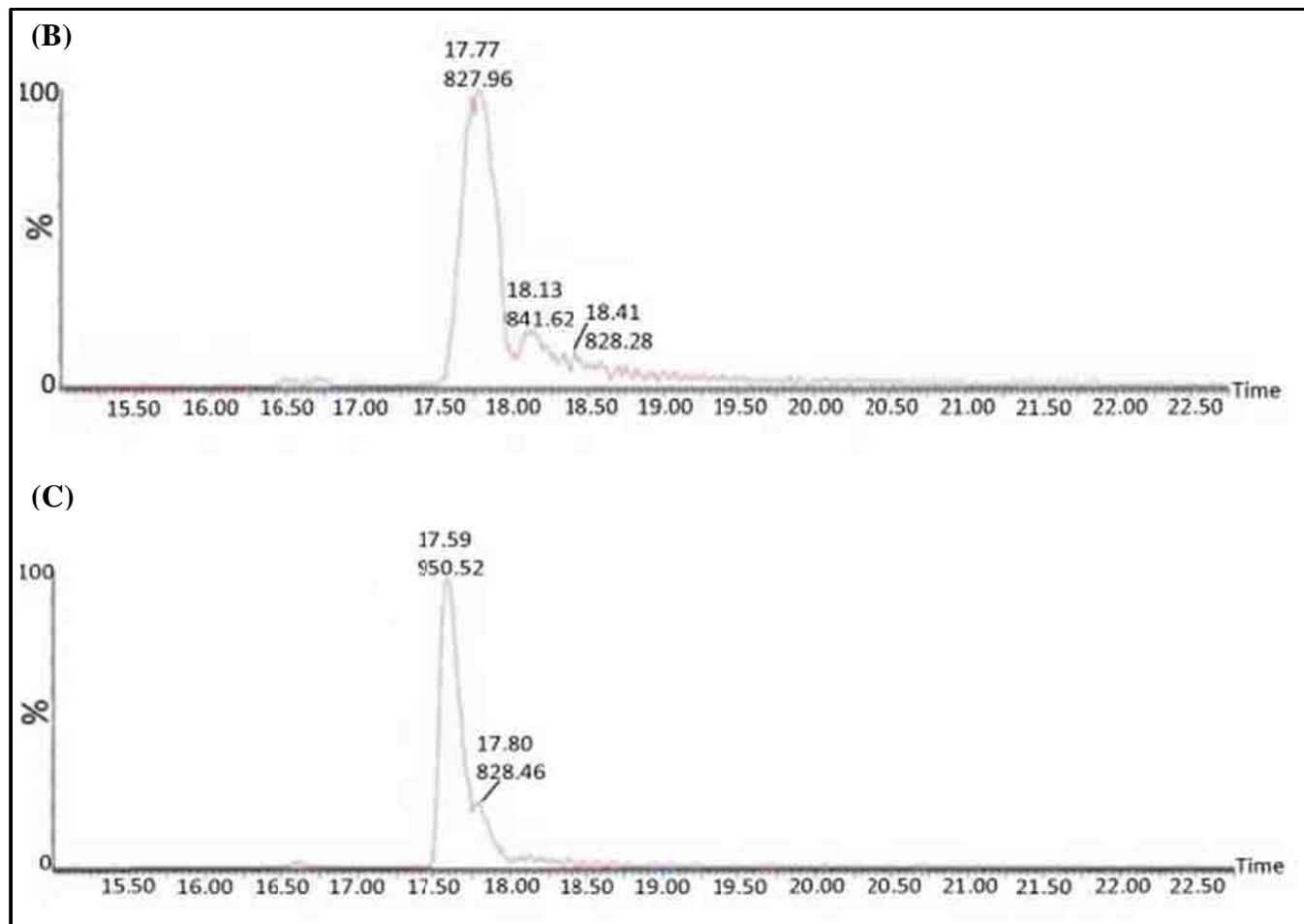


Figure 5-4. (A) 2D plot of m/z versus retention time representing the coupling of nanoLC to ion mobility separation. Circled in red is the separation of the 4.7 (top) and 4.9 kDa (bottom) peptide. (B) Extracted ion chromatograms recorded for the 4.7 kDa peptide and (C) 4.9 kDa peptide.

Figure 5-4. cont'd.



The peptides measured at 4745.4126 and 4961.4771 Da that were separated using IMS were subjected to MS fragmentation. However, insufficient fragments from the parent ions were generated; therefore, the MS data could not be used for *de novo* sequencing.

5.3.4 MALDI MS/MS of Peptides

Due to limited fragmentation from the ion-mobility spectrometry-mass spectrometry analysis, the leukocyte fraction exhibiting antimicrobial activity was analyzed using MALDI MS/MS. The two major peaks were observed at 4.7 and 4.9 kDa (Figure 5-5A), which were in agreement with the ESI-MS data (Figure 5-1). The peptides at 4.7 and 4.9 kDa were selected for

MALDI LIFT-MS/MS analysis (Figures 5-5B and 5-6C) and the primary structure was obtained for the 4.7 kDa peptide via *de novo* sequencing. The peptide sequence RDHRTKLLKKFDKSMMPCEKCKESKDFKKNKKTWECGA was determined for the 4.7 kDa peptide using both automated and manual *de novo* sequencing. Fragment ions predicted for the 4.7 kDa peptide sequence were confirmed in the mass spectrum shown in Table 5-1.

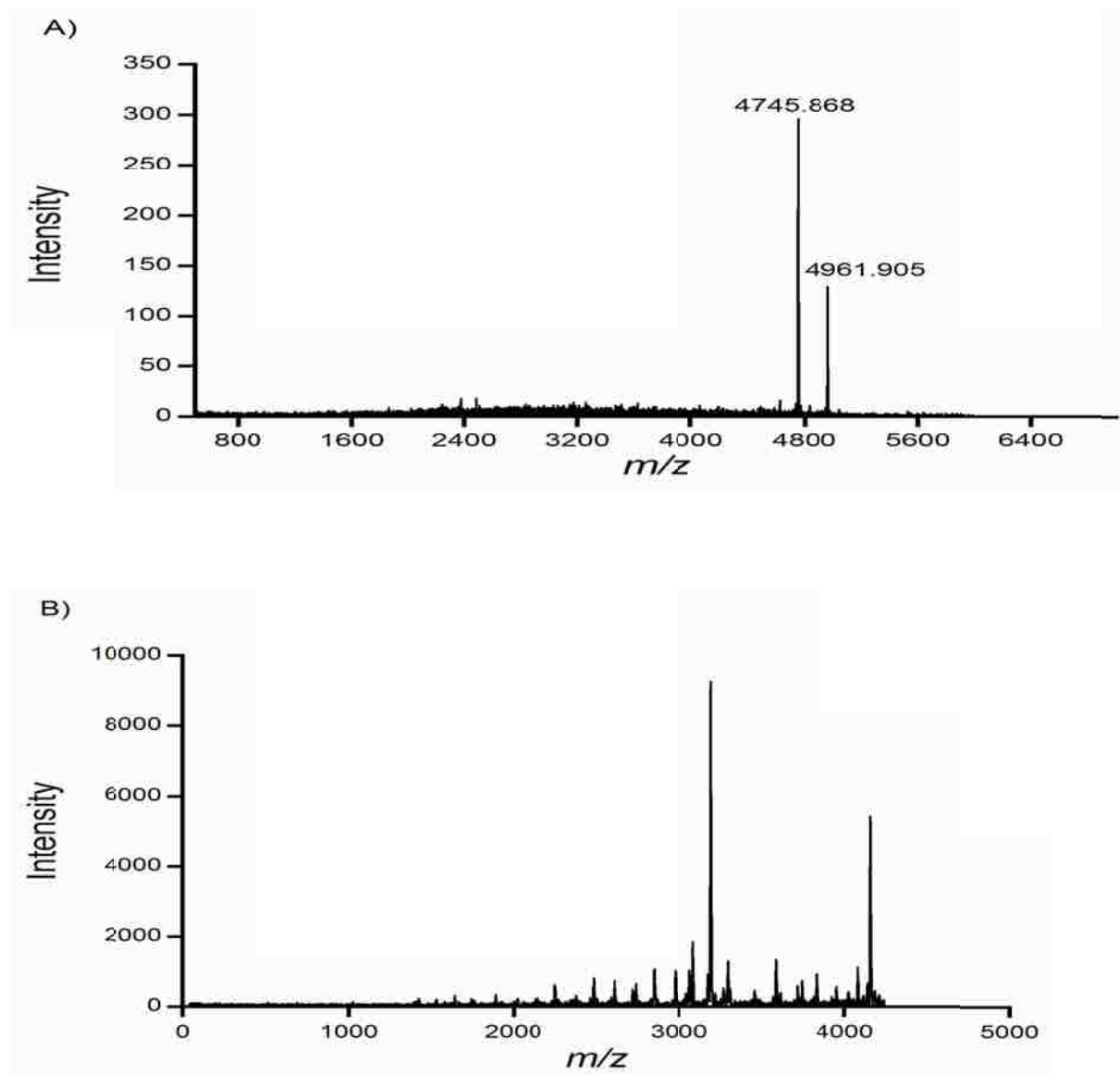


Figure 5-5. (A) MALDI spectrum of leukocyte fraction exhibiting antimicrobial activity. (B) MALDI LIFT-TOF/TOF spectrum of the 4.7 kDa peptide and (C) 4.9 kDa peptide.

Figure 5-5. cont'd.

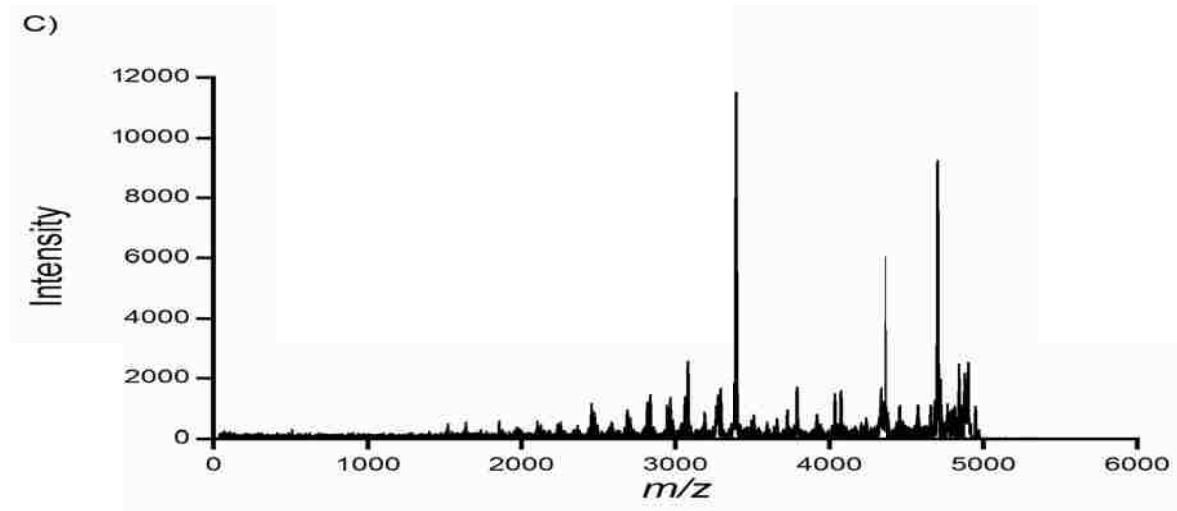


Table 5-1. Fragment ions predicted and confirmed in the mass spectrum for the 4.7 kDa peptide.

N-term	Residue	y-series	b-series	C-term
1	R	4746.3921	157.1084	39
2	D	4590.2910	272.1353	38
3	H	4475.2641	409.1942	37
4	R	4338.2052	565.2954	36
5	T	4182.1041	666.3430	35
6	K	4081.0564	794.4380	34
7	K	3952.9614	922.5330	33
8	L	3824.8665	1035.6170	32
9	K	3711.7824	1163.7120	31
10	K	3583.6874	1291.8069	30
11	F	3455.5925	1438.8754	29
12	D	3308.5241	1553.9023	28
13	K	3193.4971	1681.9973	27
14	S	3065.4022	1769.0293	26
15	M	2978.3701	1900.0658	25
16	M	2847.3296	2031.1103	24
17	P	2716.2892	2128.1630	23
18	E	2619.2364	2257.2056	22
19	C	2490.1938	2360.2148	21
20	E	2387.1846	2489.2574	20
21	K	2258.1420	2617.3524	19
22	C	2130.0471	2720.3615	18
23	K	2027.0379	2848.4565	17
24	E	1898.9429	2977.4991	16
25	S	1769.9003	3064.5311	15
26	K	1682.8683	3192.6261	14
27	D	1554.7733	3307.6530	13
28	F	1439.7464	3454.7215	12
29	K	1292.6780	3582.8164	11
30	K	1164.5830	3710.9114	10
31	N	1036.4880	3824.9543	9
32	K	922.4451	3953.0493	8
33	K	794.3502	4081.1442	7
34	T	666.2552	4182.1919	6
35	W	565.2075	4368.2712	5

Table 5-1. cont'd.

36	E	379.1282	4497.3138	4
37	C	250.0856	4600.3230	3
38	G	147.0764	4657.3445	2
39	A	90.0550	4728.3816	1

Mass tolerance: 0.5 Da

MS/MS tolerance: 0.5 Da

Bold represents ions found in the mass spectrum.

Peptides in the low mass range of the spectrum were absent due to inefficient fragmentation of the 4.7 kDa peptide. However, the fragments in the higher m/z range correlated with the “b” and “y” ions in the mass spectrum, and allowing the determination of the peptide sequence from the C- to N-terminus.

The 4.9 kDa peptide was sequenced in a similar manner to the 4.7 kDa peptide and the sequence obtained for this peptide is

MGAKTKWKRFRGLDTCVFFLLSCKKHCCLLHHTKGRNKTKAAV.

The peptide primary structure was determined based on the fragment ions in Table 5-2. Fragments in the mass spectrum that were not present in either the b- or y-ion series for the 4.9 kDa peptide were predicted by the *de novo* software.

Table 5-2. Fragment ions predicted and confirmed in the mass spectrum for the 4.9 kDa peptide.

N-term	Residue	y-series	b-series	C-term
1	M	4961.6449	132.0478	43
2	G	4830.6044	189.0692	42
3	A	4773.5829	260.1063	41
4	K	4702.5458	388.2013	40
5	T	4574.4509	489.2490	39
6	K	4473.4032	617.3439	38
7	W	4345.3082	803.4233	37
8	K	4159.2289	931.5182	36
9	R	4031.1339	1087.6193	35
10	F	3875.0328	1234.6877	34
11	R	3727.9644	1390.7889	33
12	G	3571.8633	1447.8103	32
13	L	3514.8418	1560.8944	31
14	D	3401.7578	1675.9213	30
15	T	3286.7308	1776.9690	29
16	C	3185.6832	1879.9782	28
17	V	3082.6740	1979.0466	27
18	F	2983.6056	2126.1150	26
19	F	2836.5371	2273.1834	25
20	L	2689.4387	2386.2675	24
21	L	2576.3847	2499.3516	23
22	S	2463.3006	2586.3836	22
23	C	2376.2686	2689.3928	21
24	K	2273.2594	2817.4877	20
25	K	2145.1644	2945.5827	19

Table 5-2. cont'd.

26	H	2017.0695	3082.6416	18
27	C	1880.0106	3185.6508	17
28	C	1777.0014	3288.6600	16
29	L	1673.9922	3401.7440	15
30	L	1560.9081	3514.8281	14
31	H	1447.8241	3651.8870	13
32	H	1310.7651	3788.9459	12
33	T	1173.7062	3889.9936	11
34	K	1072.6586	4018.0886	10
35	G	944.5636	4075.1100	9
36	R	887.5421	4231.2111	8
37	N	731.4410	4345.2541	7
38	K	617.3981	4473.3490	6
39	T	489.3031	4574.3967	5
40	K	388.2554	4702.4917	4
41	A	260.1605	4773.5288	3
42	A	189.1234	4844.5659	2
43	V	118.0863	4943.6343	1

Mass tolerance: 0.5 Da

MS/MS tolerance: 0.5 Da

Bold represents ions found in the mass spectrum.

5.3.5 ESI MS/MS of Peptides

ESI MS/MS experiments were performed to collect corresponding sequence information on the peptides from the fraction exhibiting antimicrobial activity, to overlap with the peptide sequences obtained from the MALDI TOF/TOF experiments. The ion measured at m/z 940.5 (5+ charge), was subjected to MS/MS fragmentation and data was acquired using different collision energies. The spectrum generated using 45 eV collision energy shows the following fragmentation pattern in Figure 5-6.

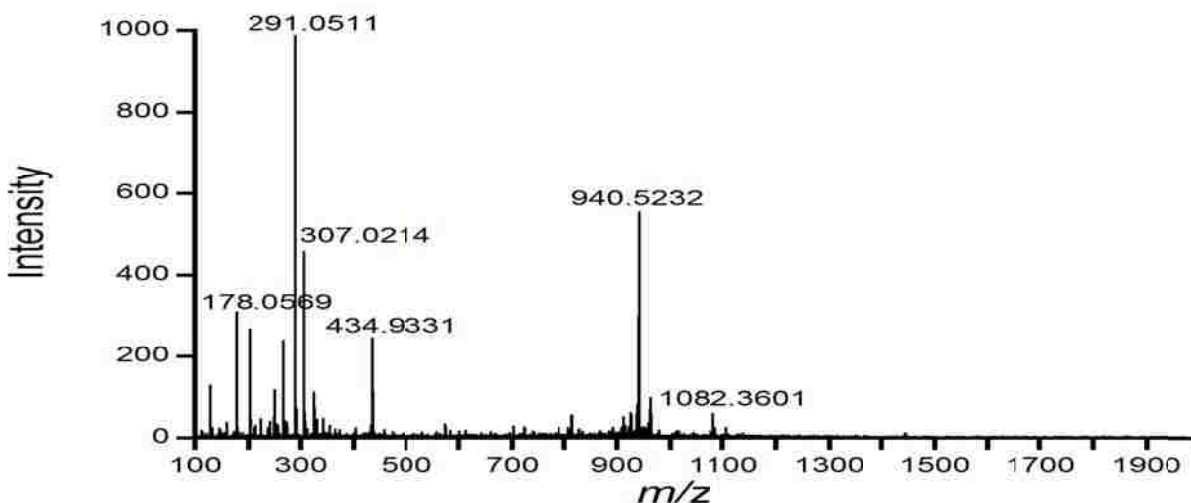


Figure 5-6. MS/MS of on the ion measured at m/z 940 with a 5+ charge.

Incomplete sequence information was determined from the spectrum due to the lack of fragmentation of the parent ion. Fragmentation with lower collision energies (e.g. 25 eV) and higher collision energy (e.g. 60 eV) were the same. This data suggests that 1) the size of the intact peptide is too large to fragment using low energy CID and 2) disulfide bond(s) may be present which will stabilize the peptide structure and make it difficult to fragment.

5.3.6 Alligator Peptide Sequence Analysis

The alligator peptide sequence generated from the 4.7 kDa peptide was searched against the antimicrobial peptide database for sequence similarity and it showed 36% sequence similarity to Pilosulin 4, an antimicrobial peptide from an Australian ant and 33% sequence homology to NA-CATH, a snake cathelicidins. The similarities between the antimicrobial peptides are seen in Figure 5-7.

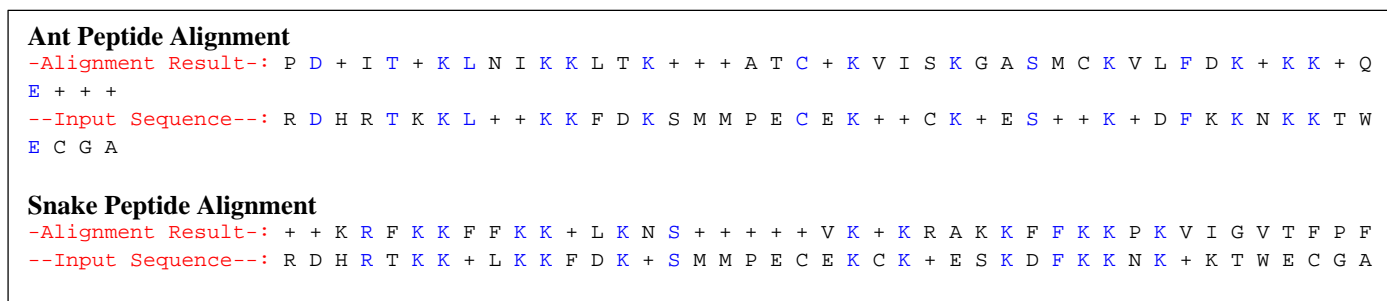


Figure 5-7. Sequence comparison among antimicrobial peptides with alligator 4.7 kDa peptide. Residues highlighted in blue represent identity in the sequence.

The alligator peptide sequence has low sequence homology when compared to antimicrobial peptides from other species, including snakes, from the antimicrobial peptide database. This suggests that the alligator sequence may function as an antimicrobial peptide based on its antimicrobial characteristics but its sequence does not share strong homology to other reptilian antimicrobial peptides.

The 4.7 kDa peptide has a theoretical isoelectric point (pI) of 9.63 which is close to the average pI of antimicrobial peptides which is 9.26.²⁹³ However, a high positive net charge is also required for antimicrobial peptides to carry out their function. The 4.7 kDa peptide sequence has a net charge of +8 at pH 5.5, indicating that it will also be cationic under *in vivo* conditions near physiological pH. The cationic nature of antimicrobial peptides is conserved and it may be essential for the initial electrostatic attraction to the phospholipid membrane of bacteria and fungi.²⁴

There are seven hydrophobic residues at positions 8, 11, 15, 16, 28, 35, and 39 and the peptide may form an alpha helix. The total net charge, hydrophobic ratio, and Boman index for the peptides were calculated using the antimicrobial peptide prediction server. The antimicrobial peptide prediction server predicts whether a new peptide sequence has the potential to be antimicrobial based on principles such as size, net charge, hydrophobic percentage and residue composition. The system can also perform simple structural predictions, for example if hydrophobic residues occur at every two to three residues in the peptide sequence an amphipathic helix will be predicted.²⁹²

The Boman index is an estimate of the antimicrobial peptide's potential to bind to other proteins.²⁹⁴ A lower index value (≤ 1) indicates that the peptide will most likely have more antibacterial activity with little side effects *in vivo* whereas a higher index value indicates that the peptide is multifunctional.²⁹⁴ The sequence determined for the peptide measured at 4.7 kDa has a Boman index of 3.8 kcal/mol. It also has an even number of cysteines which suggests that it may form a disulfide-bond linked beta structure,²⁹² similar to defensins. The percent hydrophobicity of this peptide is 25%, which is similar to that for defensins;²⁹⁵ suggesting that the peptide may have antimicrobial activity.²⁹⁶

The sequence similarity search for the 4.9 kDa peptide showed 33% sequence identity to SpStrongylocin 2 from sea urchin and Brevinin-2PTa from frog. The sequences of the peptides can be seen in Figure 5-8.



Figure 5-8. Sequence similarity between antimicrobial peptides with alligator 4.9 kDa peptide. Highlighted in blue represent identical peptides.

Like the 4.7 kDa peptide, the sequence similarity with the alligator 4.9 kDa peptide to other peptides in the database is low.

The 4.9 kDa peptide sequence has a net charge of +13 at pH 5.5 and a theoretical pI of 10.26. The peptide has 19 hydrophobic residues, 2 Val, 5 Leu, 3 Phe, 4 Cys, 1 Met, 3 Ala, and 1 Trp and the percent hydrophobicity of the peptide is 44%. The peptide sequence of the 4.9 kDa peptide contains an even number of cysteines; therefore it may form a disulfide-bond linked beta structure similar to β -defensins. The Boman index for this peptide is 1.68 kcal/mol, which is similar to defensins,²⁹⁵ suggesting that the peptide may have antimicrobial activity.²⁹⁴

5.4 Summary

Peptides and proteins were extracted from alligator leukocyte and the mixture was tested for antimicrobial activity. The extract was separated, each fraction tested for antimicrobial activity, and the peptides from the active fraction were sequenced using mass spectrometry and *de novo* sequencing. Based on ESI and MALDI MS measurements, the masses of the predominant peptides in the active fraction were found to be 4.7 and 4.9 kDa. The peptides were

not separated by RP-HPLC; hence ion-mobility mass spectrometry was used to separate the peptides by their size, mass, and charge for MS analysis and sequencing. Due to limited fragmentation from IM-MS, MALDI MS/MS was used with *de novo* sequencing to determine the sequence of the 4.7 kDa peptide. The putative sequence has 39 residues and a net charge of +8 at pH 5.5 and the 4.9 kDa peptide has a putative sequence containing 43 residues and a net charge of +13 at pH 5.5. Based on the peptide's primary structures, size, cationicity, and its predicted antimicrobial activity; the 4.7 and 4.9 kDa peptide properties are consistent with the characteristics of antimicrobial peptides.

CHAPTER 6. ISOLATION AND DETERMINATION OF THE PRIMARY STRUCTURE OF A LECTIN PROTEIN FROM THE SERUM OF THE AMERICAN ALLIGATOR (*ALLIGATOR MISSISSIPPIENSIS*)*

In this chapter mass spectrometry in conjunction with *de novo* sequencing for determining the amino acid sequence of a 35 kDa lectin protein isolated from the serum of the American alligator (*Alligator mississippiensis*) that exhibits calcium-dependent binding to mannose is described. The protein was isolated from alligator serum using mannose affinity chromatography and the N-terminal sequence was determined using Edman degradation. Enzymatic digestion with trypsin, α -chymotrypsin, Lys-C, Glu-C, and Asp-N proteases was used to generate peptide fragments for analysis by liquid chromatography tandem mass spectrometry (LC-MS/MS). Separate analysis of the protein digests with multiple enzymes extended the protein sequence coverage. *De novo* sequencing was accomplished using Mascot Distiller and PEAKS software and the sequences were searched against the NCBI database using Mascot and BLAST to identify homologous peptides. A 35 kDa protein was identified as an alligator lectin.

6.1 Introduction

Animal lectins comprise three major families: S-type lectins, P-type lectins, and C-type lectins.²⁹⁷ S-type lectins (also known as galectins) are found in the cytoplasm and nucleus of cells and sometimes on cell surfaces.²⁹⁸ This family of lectins is involved in cell adhesion and plays an important role in normal development of multicellular animals.²⁹⁸ P-type lectins target lysosomal enzymes in their subcellular compartment,²⁹⁹ and they differ from other lectins in that they have the ability to recognize phosphorylated mannose residues. The C-type lectins bind to carbohydrates in a calcium-dependent manner and are grouped into three classes: endocytic lectins, collectins, and selectins.²⁹⁸

*The work reported in the chapter has been submitted to *Comparative Biochemistry and Physiology – Part B: Biochemistry & Molecular Biology*. The protein sequence data reported in this paper will appear in the UniProt Knowledgebase under the accession number P86928.

Endocytic lectins are found in cell membranes and are specific for different sugars, for example mannose, which is found on the cell surface of microorganisms.²⁹⁸ Endocytic lectins act as opsonins, binding to the cell surface carbohydrates on microorganisms, stimulating phagocytosis by leukocytes, and eventually death. Collectins function in a similar way to endocytic lectins; they bind to microorganisms that have mannose glycans on their surfaces, leading to lysis of the pathogen and activation of the innate immune system.²⁹⁷ Selectins direct the movement of leukocytes to sites of inflammation.³⁰⁰ The importance of animal lectins is increasingly recognized and studied because of their critical roles in host defense and potential for novel medical applications.³⁰¹

Among the family of animal lectins, the C-type lectin known as mannose-binding lectin (MBL) has been well characterized in humans and some other animals.^{302, 303} Mannose-binding lectin is a calcium-dependent protein found in the serum of mammals.³⁰⁴ This protein, which is generated in the liver, tags the surface of bacteria making them susceptible to destruction.³⁰⁴ The C-type lectin consists of a collagen-like domain which interacts with cells that have been targeted by the innate immune system.³⁰⁵ Binding of MBL to target cells activates the serum complement component of the immune system, which is an important element of innate immunity³⁰⁵ and consists of a set of plasma proteins that attack extracellular pathogens.³⁰⁶ Complement proteins play an important role in innate immunity. The innate immune system can identify foreign material by recognizing pattern receptors, such as mannose, that are found on the surface of pathogens.^{306, 307}

Another family of animal lectins that bind to carbohydrates in a calcium-dependent manner is intelectins.³⁰⁷ Intelectins exhibit high sequence homology and have been identified in many organisms including human,³⁰⁸ mice,³⁰⁹ frogs,³¹⁰ trout,³¹¹ and ascidians.³¹² The principal function of intelectins is to participate in cell differentiation,³¹³ apoptosis,³¹⁴ and recognize tumor

antigens.³¹⁵ However, they also recognize pathogens and bacterial components and have roles in innate immunity.³⁰⁷ Hence, they behave similarly to C-type lectins.

Reptilian lectins such as those from lizards³¹⁶ and snakes³¹⁷ have been isolated and characterized. However, limited information is available for alligator lectins. Alligators have a potent immune system and their serum produces a broader antibacterial activity compared to other vertebrates such as humans.³¹⁸ Hence, characterization of lectins from alligator serum may provide a better understanding of the structure-function relationship of their immune system.

Lectins from different species such as *Turbo (Lunella) coreensis*,³¹⁹ *Bubalus bubalis*,³²⁰ *Halocynthia roretzi*,³¹² and *Salmo salar*³²¹ have been isolated and identified using Edman degradation and cDNA cloning.³²²⁻³²⁴ However, mass spectrometry is a more commonly used tool for protein sequencing and identification.³²⁵ Tandem mass spectrometry (MS/MS) in conjunction with separation techniques is a commonly used approach to sequence and characterize proteins derived from complex mixtures.³²⁵ A recent study showed the use of mass spectrometry to characterize the primary sequence of a galactose-binding lectin from the seeds of a legume plant.³²⁶ In addition to this study, several other studies have been conducted that demonstrate the utility of mass spectrometry for identifying lectin sequences.³²⁷⁻³²⁹

One of the major challenges in the study of alligator proteins is the limited information available on the reptilian genome and proteome. For proteomic analysis of species with limited proteomic data, *de novo* sequencing can be used to determine protein sequences.³³⁰ In typical *de novo* sequencing methods, proteins are isolated and enzymatically digested, and the resulting peptides are analyzed by tandem mass spectrometry. The sequence is determined by observing the mass differences between fragment peaks in the tandem mass spectra that are generated by fragmentation of the peptide of interest.³³¹ *De novo* sequencing was previously used to sequence peptides from proteins of the American alligator leukocytes.³³²

In the present study, mannose affinity separation was used to isolate a 35 kDa lectin protein from the serum of the American alligator (*Alligator mississippiensis*). Liquid chromatography ESI-MS and MALDI-MS was used to obtain the mass of the intact protein. The protein was digested using several different proteases. Peptide sequences were determined using *de novo* sequencing. Database searching and BLAST search was also used to match peptides with more than 50% homology. Using mass spectrometry and different enzymes independently, peptides of different lengths were generated and aligned to determine the sequence of the American alligator lectin.

6.2 Experimental

Proteins were isolated from alligator serum with affinity chromatography. Mannan agarose (3 mL) was equilibrated with 3 mL of mixing buffer (10 mM Tris-HCl, pH 7.8, 1.25 M NaCl) and allowed to settle into a column prepared from a Pasteur pipette plugged with cotton. Fresh alligator serum (5 mL) was mixed with 5 mL of loading buffer (20 mM Tris-HCl, pH 7.8, 2.5 M NaCl). The diluted serum was allowed to filter through the column and the column was washed with 10 volumes of loading buffer. The proteins were eluted with 5 mL of elution buffer (10 mM Tris-HCl, pH 7.8, 1.25 M NaCl, 2 mM EDTA). The isolated lectin protein was transferred to 10 kDa centrifugal concentrator tubes and centrifuged at 7500 g to concentrate and desalt the sample. A portion of the concentrate was used for mass analysis before enzymatic digestion.

The collected lectin samples were analyzed by reverse phase liquid chromatography (RP-LC) ESI time-of-flight mass spectrometry (Agilent 6210) and matrix-assisted laser desorption ionization time-of-flight/time-of-flight tandem mass spectrometer to obtain the mass of the intact protein. For LC/MS, 10 µg of sample obtained from affinity separation was dissolved in 10 µl of water containing 0.1% formic acid (FA) and loaded onto a C₁₈ reverse phase column 5 µm 250 ×

mm (Vydac) which had been equilibrated with 0.1% FA in water. A linear gradient from 0% to 60% of B over 60 minutes was used for protein separation. Solvent A was water containing 0.1% FA and solvent B was acetonitrile containing 0.1% FA. The eluent from the LC column was directed towards the mass spectrometer and mass spectra were acquired from 600 to 3000 m/z using ESI at 4200 V. The spectra were extracted and deconvoluted using Analyst software.

The protein samples for MALDI analysis were deposited on a ground-steel target using a 1:1 ratio (v/v) of sample and saturated matrix solution. The matrices used were a 9:1 mixture of 2,5-dihydroxybenzoic acid and 2-hydroxy-5-methoxybenzoic acid (sDHB) and 2,4-dimethoxy-3-hydroxycinnamic acid (sinapinic acid, SA). The samples were analyzed in positive ion mode and 500 shots were collected. The mass spectra were collected using a mass range from 20 to 90 kDa. Exact masses from the spectra were determined using Bruker FlexAnalysis software.

One-dimensional SDS-PAGE was performed using pre-cast 4–20% polyacrylamide gradient tris-HCl gels on the small format gel electrophoresis system. The samples were mixed 1:1 v/v with sample buffer containing 62.5 mM tris-HCl at pH 6.8, 2% SDS, 25% glycerol (w/v), 0.01% w/v bromophenol blue, and, in both the presence and absence of 10% 2-mercaptoethanol (v/v), heated at 95°C in a water bath for 5 minutes before loading on the gel. Samples were run at 100 V for 1 h using tris-glycine (pH 8.3) as the gel running buffer. Gels were stained with Coomassie blue for 1 h and rinsed twice with distilled water for 10 min (each time) to remove the stain. The gel images were captured using the Gel Doc XR System and with BioRad Quantity One 1-D analysis software. Samples were excised from the SDS polyacrylamide gel and submitted for N-terminal sequencing by Edman degradation using a protein sequencer (Model 494, Perkin Elmer Applied Biosystems).

A purified alligator lectin (40 μg) was incubated in the presence of 40 μL of mannan-, mannose-, D-lactose-, N-acetylgalactosamine-, N-acetylglucosamine-, and β -D-glucose-agarose.

After 30-minute incubation at ambient temperature, the samples were centrifuged and a 20 μ L aliquot of the supernatant (containing the unbound protein) was subjected to SDS-PAGE. One-dimensional SDS-PAGE was performed and the gels were stained with Coomassie blue. The experiment was repeated three times. The density of the bands, in pixels, was evaluated and compared using ImageJ software.³³³

To obtain maximum sequence coverage, separate aliquots of lectin protein obtained from affinity separation were subjected to several different enzymatic digestions. The protein fractions were dissolved in a 100 mM ammonium bicarbonate buffer adjusted to the pH range in which each endoprotease was most active. The protein was reduced using 50 mM DTT in 25 mM ammonium bicarbonate buffer and incubated at 37°C for one hour. Following reduction, the protein was alkylated with 50 mM iodoacetamide in 25 mM ammonium bicarbonate buffer for 45 minutes at room temperature in the dark. The in-solution digestions were performed separately using trypsin, α -chymotrypsin, Lys-C, Glu-C, or Asp-N according to the manufacturer's protocol. Briefly, digestion with trypsin was performed using a 1:20 ratio of enzyme to substrate in 100 mM ammonium bicarbonate, pH 8.5; with α -chymotrypsin using a 1:20 ratio at pH 8.5; with Lys-C using a 1:20 ratio at pH 8.5; with Glu-C using a 1:20 ratio at pH 7.8; and with Asp-N using a 1:50 ratio at pH 8.5. All digestions were run at 37°C overnight. The enzymatic digests were dried in a vacuum centrifuge and resuspended in 0.1% FA for LC separation.

Bands from the one-dimensional gels were excised and cut into \sim 1 mm cubes, and the stain was removed with 200 mM NH_4HCO_3 and 40% acetonitrile (2 \times 30 min). After destaining, the gel pieces were dried in a vacuum centrifuge, rehydrated with 20 μ L of 0.4 μ g trypsin in 40 mM ammonium bicarbonate containing 9% acetonitrile to completely cover the gel pieces. The tubes were then incubated overnight at 37°C.³³⁴ A similar approach was used for digesting the protein gel pieces using α -chymotrypsin, Lys-C, Glu-C, and Asp-N proteases.^{335, 336} Peptides

were extracted from gel pieces with 60% acetonitrile and 40% water containing 0.1% trifluoroacetic acid (TFA). CNBr digestions were performed overnight with 1 M CNBr at room temperature in the dark. The following day the CNBr digestion was dried completely using a speed-vac and resuspended in water containing 0.1% FA for ESI analysis. FA digestions were performed overnight at 37°C.

Liquid chromatography electrospray tandem mass spectrometry (LC MS/MS) was used to analyze the peptides derived from the in-gel and in-solution digests. The peptides were dissolved in 20 μ L of 0.1% FA and 10 μ L of the digest was injected onto a 0.3 \times 1 mm trapping column (PepMap) on a nano-LC system (Ultimate,) at a flow rate of 10 μ L/min. The peptides were eluted onto a 75 μ m \times 15 cm C₁₈ column (Biobasic) and separated using a gradient of 5 - 40% B over 90 minutes with a flow rate of 200 nL/min. The peptides were analyzed via a quadrupole time-of-flight mass spectrometer (QSTAR) and ionized using a nano-electrospray source at a voltage of 2.5 kV. The mass spectrometer was operated in information-dependent acquisition (IDA) mode. Three collision energies (25, 38, and 50 eV) were selected to fragment the peptides.

The peptide sequences were generated from MS/MS data using Mascot Distiller and PEAKS software. The Mascot Distiller software determined amino acid sequences from MS/MS data using-peak fitting and isotope distribution²⁴¹ and PEAKS software computes peptides for whose ions matches as many of the most abundant peaks in the spectrum.¹⁶⁷ The *de novo* sequencing results were also verified manually. *De novo* peptide sequences were searched using Mascot and BLAST for protein identification based on homology searches. For Mascot and BLAST search NCBI database were used to compare to peptide sequences derived from all taxonomies.

Multiple sequence alignments were performed using DIALIGN, CLUSTALW, and BLAST described in Section 2.7.5. The 35 kDa alligator lectin sequence was aligned and compared with four other vertebrate lectin proteins.

6.3 Results and Discussion

6.3.1 Isolation of the 35 kDa Alligator Lectin

SDS-PAGE analysis under non-reducing conditions (Figure 6-1a) of the isolated alligator lectin proteins showed five major bands between 35 and 250 kDa, which suggests a mixture of proteins or oligomers. By contrast the SDS-PAGE under reducing conditions showed only two bands that corresponded to 35 and 70 kDa (Figure 6-1b), suggesting that some of the bands were the result of oligomers. The band detected at 70 kDa could be another protein or a dimer of the 35 kDa protein.

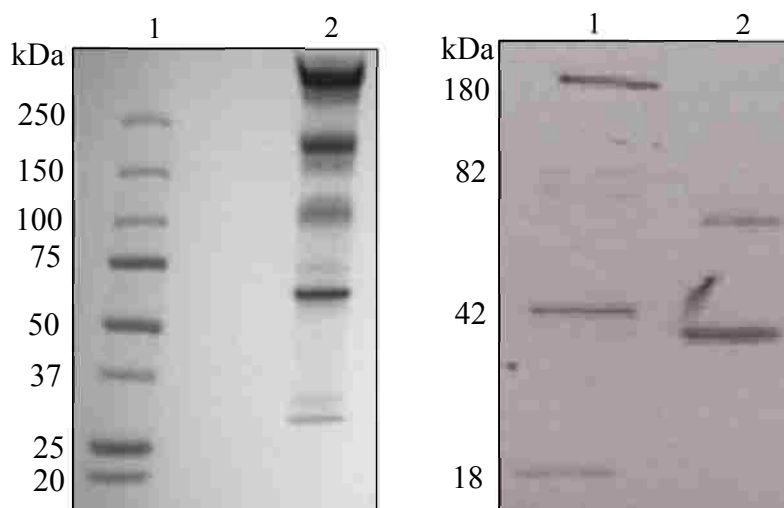


Figure 6-1. SDS-PAGE analysis of non-reduced and reduced alligator lectin protein on a 4–20% gel. (A) Lane 1, molecular weight marker; lane 2, non-reduced alligator lectin. (B) Lane 1, molecular weight marker; lane 2, reduced alligator lectin.

LC ESI-MS and MALDI-MS was performed to obtain the intact mass of the lectin protein and oligomers. Figure 6-2a shows a LC separation of the crude extract of the alligator lectin and Figure 6-2b shows mass spectrum of the major peak that eluted at R_T between 32 and

35 minutes. The deconvoluted mass spectrum of the major peak reveals the average masses of the eluted lectin protein to be 34.6, 69.3, 103.9, and 138.6 kDa, consistent with the monomer, dimer, trimer and tetramer of the 35 kDa protein.

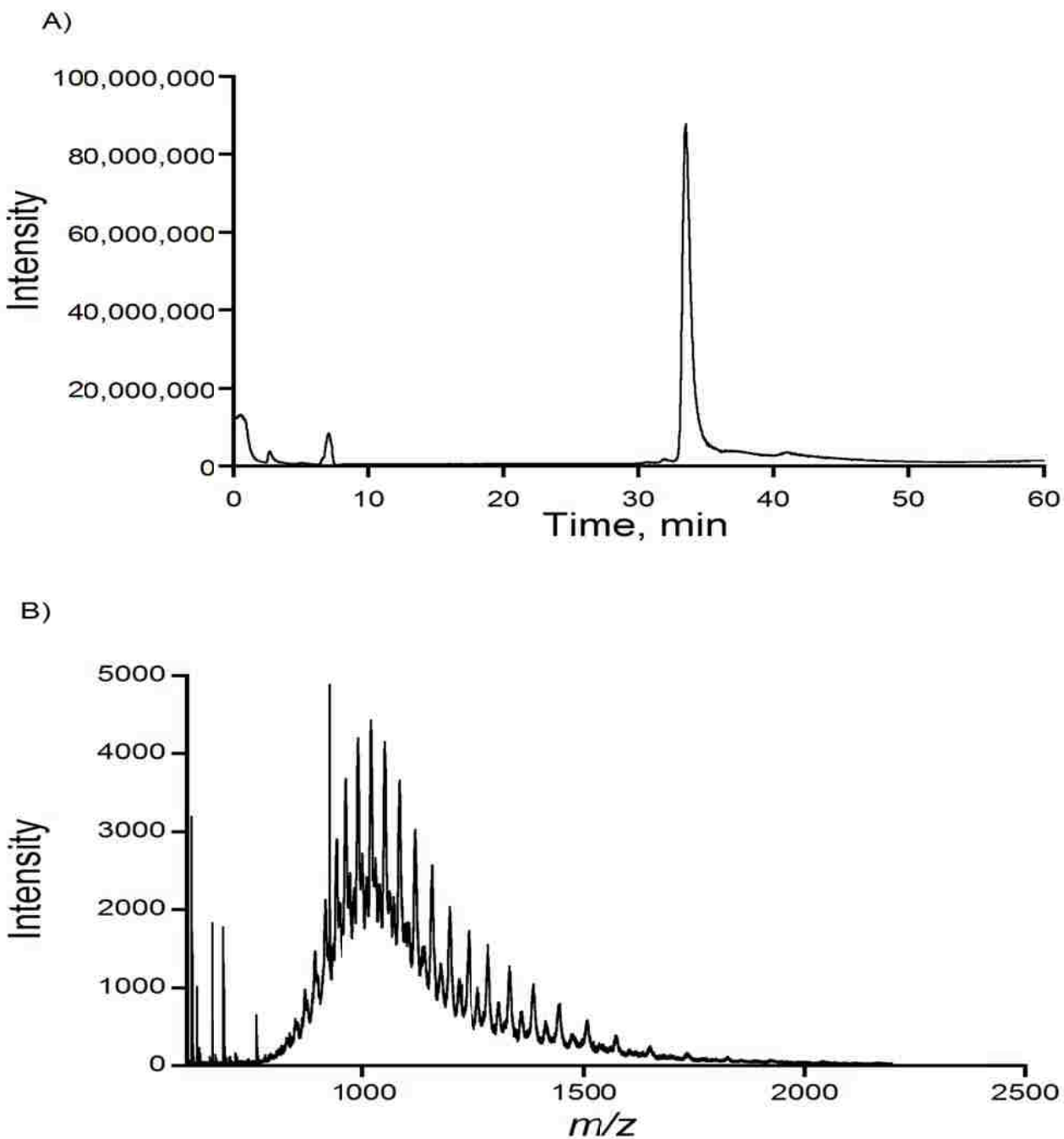


Figure 6-2. (A) The chromatogram of alligator lectin eluted from a C_{18} reversed phase column. The lectin protein was eluted between 32 and 35 minutes. (B) The mass window of the isotopic distribution for the protein masses.

Figure 6-3 shows a MALDI mass spectrum of the crude extract, the observed masses from MALDI-MS analysis were 34.7, 69.3, 104.0, and 138.6 kDa.

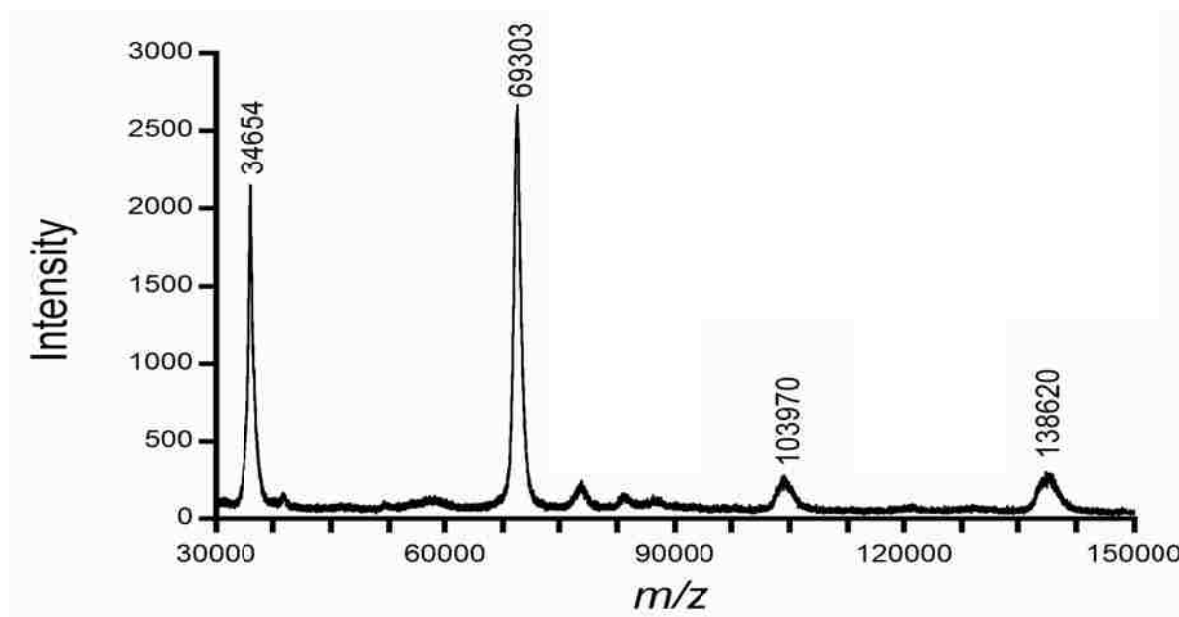


Figure 6-3. Linear mode (+) MALDI mass spectrum showing the average masses of the oligomers from the 35 kDa alligator lectin.

6.3.2 Sequence Determination of the 35 kDa Alligator Lectin.

To sequence the isolated protein, Edman sequencing and mass spectrometry were used. Edman degradation of the 35 kDa molecular weight gel band of the alligator protein produced the N-terminal sequence NNQLALKLAATGGSTNXLPA, where position X was undetermined. The unidentified amino acid at position X is potentially due to the amino acid being glycosylated (O-linked glycosylation) which can not be detected by Edman sequencing.³³⁷

Peptides were obtained from enzymatic digestion of the 35 kDa gel band and the isolated lectin sample using the different endoproteases as described above. Trypsin produced ten peptides of various lengths from the 35 kDa protein. To collect additional sequence information from overlapping peptides, the endoproteases Lys-C, Asp-N, and Glu-C were used. Endoprotease Lys-C produced five peptides, Glu-C digest produced nine peptides, and seven

peptides were produced from Asp-N digest. A less specific endoprotease, α -chymotrypsin, was also used to obtain additional overlapping peptides. The endoprotease α -chymotrypsin produced seven fragments that overlapped with fragments generated from trypsin, Asp-N, and Lys-C. There was no additional sequence coverage obtained from the CNBr and FA digest.

Peptides obtained from the in-solution and in-gel digests of the 35 kDa band were *de novo* sequenced. Figure 4 shows mass spectra for a Glu-C and tryptic digest of the protein to illustrate peptide sequence determination. Figures 6-4a and 6-4b show examples of mass spectra acquired at 57 and 58 min, respectively, in the separation of the Glu-C and tryptic digests. Tandem MS data were collected and peptide sequences were determined via *de novo* sequencing. Figures 6-4c and 6-4d illustrate the peptide fragment series generated from the MS/MS data acquired from the ion 492.8 in Figure 4a and 743.8 in Figure 6-4 b. The sequences were searched against the NCBI protein database using Mascot and no protein matches were found. The *de novo* sequenced peptides were subjected to a BLAST search to identify proteins with similar sequences. Peptide sequences found with BLAST exhibited strong similarity to intelectin-1 and intelectin-2 proteins from mammals. Some of the lectin peptide sequences showed a high degree of similarity to the intelectin peptides, with few amino acid differences; these regions of similarity appear to be conserved among the lectins. The *de novo* sequenced peptides that did not match peptides in the protein database with a high degree of similarity appear to be regions of the alligator lectin sequence that are unique.

The peptides from the different enzymatic digests produced overlapping data that were aligned to create the primary sequence of the protein (Figure 6-5). Partial protein sequences were generated when peptide sequences from the different digests were overlapped. Multiple alignments of the 35 kDa alligator lectin peptides with peptides of other lectin proteins provided sequence positioning of the alligator lectin peptides.

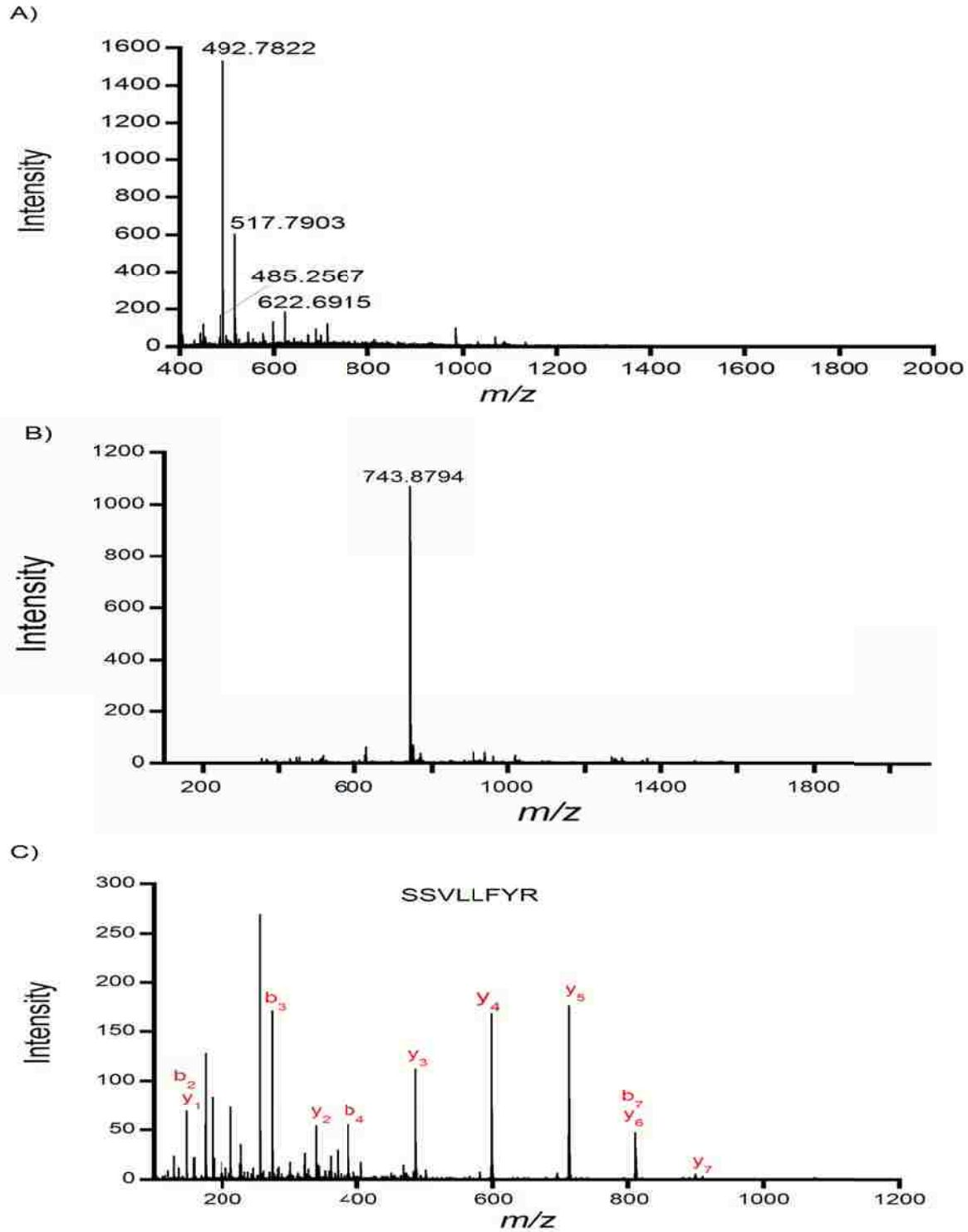
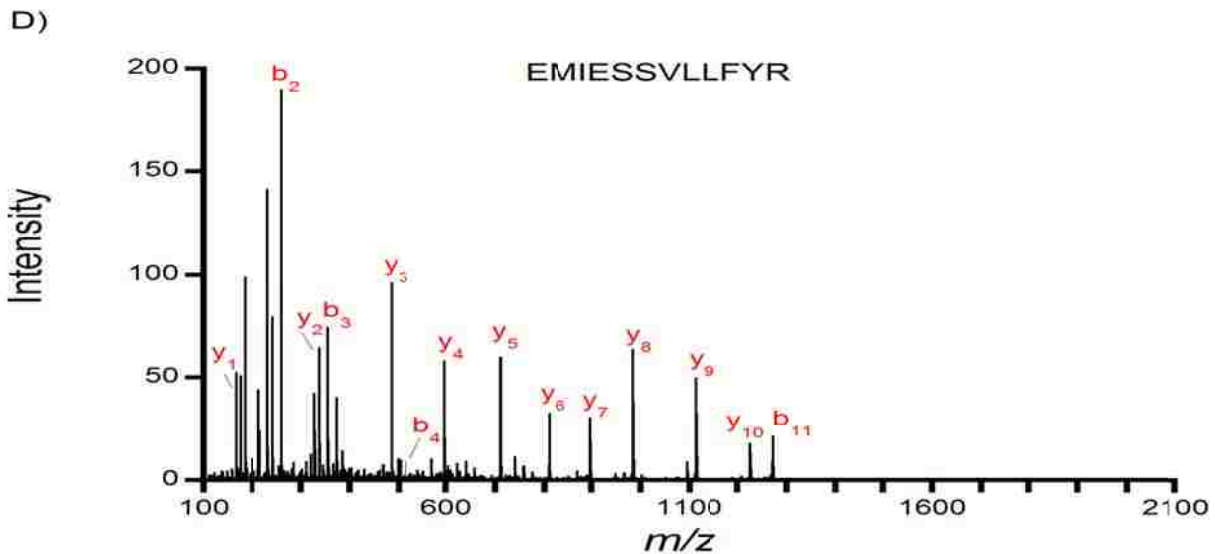


Figure 6-4. Mass spectrometry data for peptide sequencing illustrating (A) an example of peptides eluted from the Glu-C digest at 57 minutes, (B) peptides eluted from trypsin digest at 59 minutes, (C) an MS/MS plot of the ion 492.8 generated from the mass spectrum of the Glu-C digest, and (D) an MS/MS plot of the ion 767.7 generated from the mass spectrum of the trypsin digest.

Figure 6-4. cont'd.



The chymotryptic peptide LAATGGSTNSLPA of the 35 kDa protein determined by mass spectrometry and *de novo* sequencing overlapped with residues 8 to 20 from the Edman sequence and identified the missing residue as serine.

The digests from the 70 kDa gel band generated some peptides that show sequence identity to peptides from the 35 kDa protein. This suggests there is some sequence similarity between the 35 kDa and 70 kDa protein from the SDS-PAGE gel. However, unique peptide sequences were identified, suggesting that this is a different protein rather than the dimer of the 35 kDa protein.

Overall, approximately 98% of the 35 kDa protein was sequenced and found to consist of 313 residues; however, 2% of the protein was not sequenced. The missing sequence may correspond to a short tryptic peptide that was *de novo* sequenced from both the in-gel and in-solution digest. This short tryptic peptide did not overlap with the other enzymatic peptide fragments; hence its position in the sequence could not be determined.

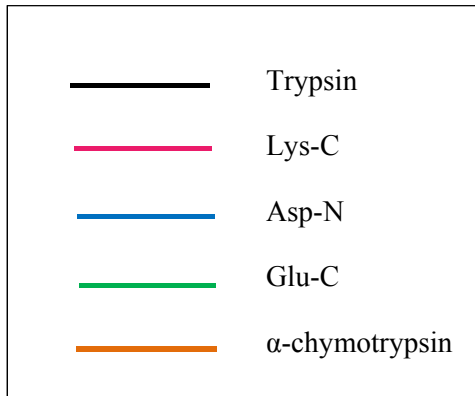
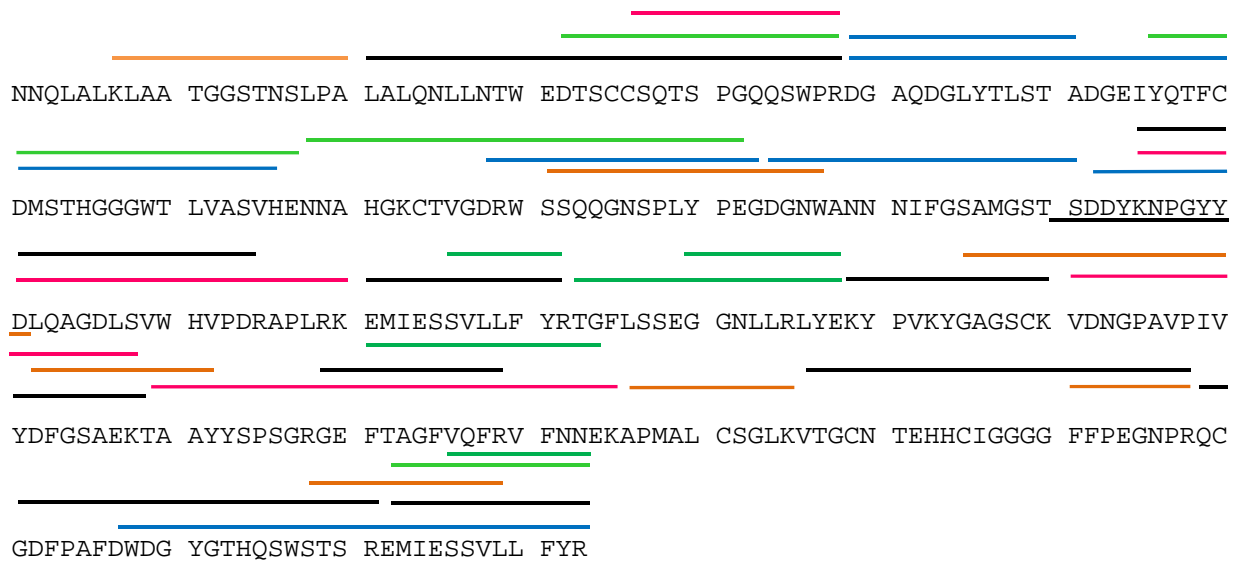
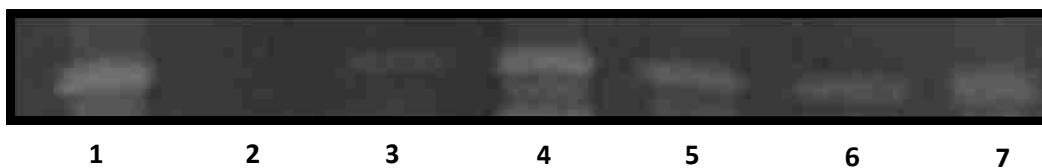


Figure 6-5. Primary structure of the 35 kDa lectin protein isolated from American alligator assembled from different endoprotease digestions. The peptide sequences were generated using ESI-MS/MS. Peptides obtained from the different enzymes was highlighted using different colors.

6.3.3 Binding Activity of Alligator Lectin

The alligator lectin-binding activity was determined for four carbohydrates, and the results are shown in Figures 6-6A and 6-6B.

A)



B)

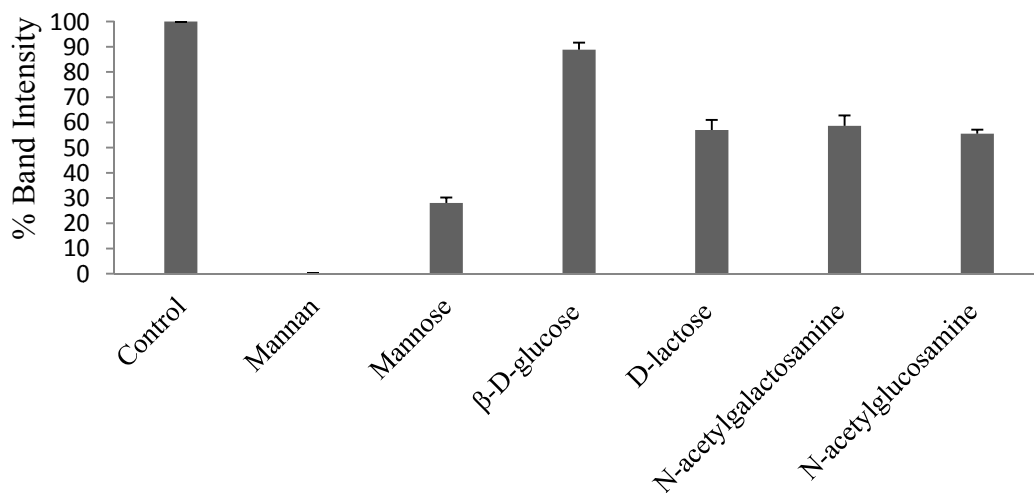


Figure 6-6. (A) The isolated alligator lectin was incubated with agarose-conjugated mannan (Lane 2), mannose (Lane 3), β -D-glucose (Lane 4), D-lactose (Lane 5), N-acetylgalactosamine (Lane 6), and N-acetylglucosamine (Lane 7). The samples were centrifuged and the supernatants were subjected to 1D gel analysis. The control (Lane 1) was alligator lectin with underivatized agarose. (B) The image of the gel bands were quantified in ImageJ, a Java-based image processing program.

The alligator lectin binds strongly to agarose-conjugated mannose and mannan, indicating little or no unbound protein remaining in the supernatant. This produced a relatively low gel band intensity of 28% for mannose and no band was observed in the presence of mannan. Weaker binding activity of the alligator lectin was observed with β -D-glucose, D-lactose, N-acetylglucosamine, and N-acetylgalactosamine; therefore, there was less interaction and a relatively stronger band intensity of 89%, 57%, 56%, and 59% respectively, compared to the control (100%).

6.3.4 Comparison of the 35 kDa Alligator Lectin Sequence with Other Animal Lectins

The alligator lectin sequence was compared with other calcium-dependent animal lectins, specifically mannose-binding lectins and intelectins, from six other species: *Homo sapiens*, *Mus musculus*, *Rattus norvegicus*, *Pan troglodytes*, *Gallus gallus*, and *Xenopus tropicalis* using DIALIGN software. A sequence comparison for the lectin of the above organisms to the alligator lectin sequence resulted in 13%, 3%, 1%, 5%, and 4% sequence similarity, respectively.

Sequence comparison of the alligator lectin sequence to intelectin-1 from *Homo sapiens*, *Mus musculus*, *Rattus norvegicus*, and *Xenopus tropicalis*, exhibited 57%, 59%, 60%, and 57% similarity respectively (Figure 6-7). A sequence comparison of the 35 kDa alligator lectin protein was also made with intelectin-2 from *Homo sapiens* (58% identity) and *Mus musculus* (59% identity). The degree of similarity of alligator lectin protein sequence to intelectin-2 sequence (Appendix B, Figure 1) is close to that of the intelectin-1. Both mannose-binding lectin and intelectin proteins bind in a calcium-dependent manner.³⁰⁷ However, sequence comparisons suggest that the lectin protein isolated here is more closely related to intelectins rather than to mannose-binding lectins.

For the human intelectin-1, residue 154 is the start of the consensus sequence (Asn-X-Thr/Ser) indicating a site of N-linked glycosylation. However, this site is not present in the alligator lectin sequence, indicating that it lacks N-glycosylation at this site. There are several possible sites of glycosylation in the alligator lectin sequence, for example at the serine and threonine residues, suggesting that O-linkage is possible. BLAST search of the alligator lectin sequence showed a conserved fibrinogen-related domain near the N-terminus region of the sequence between residues 50 to 80. The fibrinogen related domain has been shown to be similar to the calcium-dependent carbohydrate-recognition domains.³³⁸ Both mouse and human intelectin-1 sequences have a fibrinogen domain between residues 32 to 251 and 32-255

respectively. A comparison of the alligator lectin sequence to both mouse and human intelectin-1 fibrinogen domain sequences showed a high degree of sequence similarity (63% and 61% respectively).

Mouse	MTQLGFLFLFIMVATRG-----CSAAEENLDTNRWGNSSFFSSL	37
Rat	MTQLGFLFLFLIVATRG-----GSAAKEDLETNKGTHSFFDSL	37
Human	MNQLSFLFLFLIATTRG-----WSTDEANTYFKEWTCSSSPSL	37
Frog	MLVYSLLVFALGFAPAGHAGSCEQASISEKKEKILNLLACWTEKTDNNGSRFSGSPTGDM	60
Alligator	NNQLALKLAATGGSTNX-----LPALALQNLNTWEDTSCCSQ	38
	. : : : . : .	
Mouse	P---RSCKEIKQEHTKAQDGLYFLRTKNGVIYQTFCDMTTAGGGWTLVASVHENNMRGKC	94
Rat	S---RSCKEIKEENTGAQDGLYFLRTENGVIYQTFCDMTTAGGGWTLVASVHENNMGGKC	94
Human	P---RSCKEIKDECPSAFDGLYFLRTENGVIYQTFCDMTSGGGWTLVASVHENDMRGKC	94
Frog	SYGYRSCNEIKSSDPSAPDGIYTLATEHGESYQTFCDMTTNGGGWTLVASVHENNMFGKC	120
Alligator	T----SPGQQSWPRDGAQDGLYTLSTADGEIYQTFCDMSTHGGGWTLVASVHENNAHGKC	94
	. * : . * * : * * * . * * * * * : : * * * * * * * * : * * *	
Mouse	TVGDRWSSQQGNRADYPEGDGNWANYNTFGSAEAATSDDYKNPGYFDIQAENLGIWHVVP	154
Rat	TVGDRWSSQQGNRADYPEGDGNWANYNTFGSAEGATSDDYKNPGYFDIQAENLGIWHVVP	154
Human	TVGDRWSSQQGSKAVYPEGDGNWANYNTFGSAEAATSDDYKNPGYFDIQAENLGIWHVVP	154
Frog	TVGDRWSTQQGNTLQNPEDGNWANYATFGLPEGATSDDYKNPGYFDIQAENLGIWHVVP	180
Alligator	TVGDRWSSQQGNSPLYPEGDGNWANNIFGSAMGSTSDYKNPGYFDLQAGDLSVWHVVP	154
	*****:*** . ***** * * . . :*****:*** * :* :*****:	
Mouse	KSPLHN--WRKSSLLRYRTFTGFLQHLGHNLFGLYKYPVKYGEKGCWTDNGPALPVVYD	212
Rat	NSPLHS--WRNSSLLRYRTFTGFLQHLGHNLFGLYQKYPVKYGEKGCWTDNGPALPVVYD	212
Human	KSPMQH--WRNSSLLRYRTDTGFLQTLGHNLFGLYQKYPVKYGEKGCWTDNGPVPVVYD	212
Frog	NTPMFn--WRNSSILRYRTQNSFFTEEGNLFELYKYPVKYDYGKCLADNGPAVPPVYD	238
Alligator	RAPLRKEMIESSVLLFYR--TGFLSSEGGNLLRLYEKYPVKYGAGSCKVDNGPAVPIVYD	212
	. : * : . . * : * * . . * : * * * : * : * * * . * . * . * * * : * : * * *	
Mouse	FGDARKTASYSPSGQREFTAGYVQFRVFNNEERAASALCAGVRVTGCNTEHHCIGGGGFF	272
Rat	FGDAQKTASYSPYQKEFTAGFVQFRVYNNERAASALCAGMKVTGCNSEAHCIGGGGFF	272
Human	FGDAQKTASYSPYQREFTAGFVQFRVFNNEERAANALCAGMRVTGCNTEHHCIGGGGYF	272
Frog	LGSAEKTSSLYSPNGRNEFTAGFVQFRVVNTEKAALALCPGVKVKGCNAEHHCIGGGGYF	298
Alligator	FGSAEKTAAYSPSGRGEFTAGFVQFRVFNNEKAPMALCSGLKVTGCNTEHHCIGGGGFF	272
	: * . * . * * : * * * * : * * * * * * * * * * * * . * . * . * * * : * * * * * * * * *	
Mouse	PEGNPVQCGDFASFDWDGYGTHNGYSSSRKITEAAVLLFYR	313
Rat	PEGNPLQCGDFGAFDWDNGYGTHIGYSSSREITEAAVLLFYR	313
Human	PEASPOQCGDFSGFDWDSGYGTHVGYSSSREITEAAVLLFYR	313
Frog	PEGSPRQCGDFSAFDWDGYGTHAGWSASKEITEAAVLLLYR	339
Alligator	PEGNPRQCGDFPAFDWDGYGTHQSWSTSREMISSVLLFYR	313
	** . . * * * * * . * * * . * * * * . : * : * : * : * : * * * * * *	

Figure 6-7. The *Alligator mississippiensis* 35 kDa lectin amino acid sequence was compared to intelectin-1 protein of *Homo sapiens*, *Mus musculus*, *Rattus norvegicus*, and *Xenopus tropicalis*.

Previous reports show that many vertebrate lectins contain homologous amino acid sequences.³⁰³ The 35 kDa alligator lectin sequence shows a high degree of similarity to

intelectins; however, intelectin proteins are animal lectins that typically have an affinity to galactose- and galactofuranose-binding C-type lectins and require calcium for carbohydrate-binding activity.³⁰⁷ Although intelectins have been previously reported in other organisms, few studies have identified their binding specificity. In this study, the 35 kDa protein identified as alligator lectin had an affinity to mannose with a calcium-binding dependence. A previous report indicated that a rainbow trout intelectin had a binding specificity for both N-acetylglucosamine (GlcNAc) and mannose saccharides.³¹¹ The alligator lectin could be a similar intelectin-like protein that has a binding specificity for mannose residues.

Intelectin proteins are oligomeric; for example, human intelectin-1 is a trimeric protein.³³⁹ ESI and MALDI analysis (Figures 6-2b and 6-3) suggests that the alligator lectin is an oligomeric protein that exists primarily as a 35 kDa monomer *in vitro*.

Mammalian intelectins usually have at least 10 conserved cysteine residues³³⁹ for example as in human intelectin-1 (Figure 6-6); the 35 kDa alligator lectin sequence has nine cysteine residues. It has cysteine residues at positions 35 and 36 at the N-terminus region that are not present in the intelectin sequences of other vertebrates. Also, the alligator lectin sequence lacks a cysteine at position 42 that is conserved in the other organisms. These cysteine residues can influence tertiary protein structure, as well as carbohydrate-recognition specificity and oligomer formation and structure. When comparing the sequence to those derived from other vertebrates, the lack of a cysteine residue at position 41 in the alligator lectin sequence, as well as the different locations of the other cysteine residues may result in lower thermal stability of the protein.

In addition to the presence of the cysteine residues that allow for oligomers to form via disulfide bonding, intelectins also have carbohydrate recognition domains.³⁴⁰ The calcium-dependent carbohydrate-recognition domains are similar to the fibrinogen-like domain.

Fibrinogen-like proteins typically contain fibrinogen domains at the C-terminal region in both vertebrates and invertebrates.^{338, 341} An example of a human fibrinogen-like protein is ficolin. Although ficolin does not show sequence homology to C-type lectins or intelectins, it plays a vital role in innate immunity in a manner similar to the mannose-binding lectin.³⁴² There are two types of ficolins found in humans, ficolin- α which is a 326 amino acid protein and has a fibrinogen terminal domain at residues 109 to 326, and ficolin- β , a 313 amino acid protein with a fibrinogen terminal domain at residues 96 to 313.³³⁸ The alligator lectin sequence has low sequence similarity to both the fibrinogen domains in human ficolin- α (16% similarity) and ficolin- β (9% similarity) (Appendix B, Figure 2), suggesting that its carbohydrate-recognition domain is different to that found in ficolin proteins. BLAST results for the alligator lectin showed that it has a fibrinogen-related domain between residues 50 and 80, suggesting that its carbohydrate recognition domain is near the N-terminal region of its sequence as previously shown in the *Halocynthia roretzi* lectin from ascidian plasma.³¹²

Binding studies were conducted with several carbohydrates leading to the conclusion that the alligator lectin exhibits strong selectivity for both mannan and mannose, which functions as a recognition molecule in the complement system. The alligator lectin showed some binding activity towards D-lactose suggesting that the lectin may also bind to terminal galactose as observed with intelectins from other organisms. The weak binding of the lectin observed in the presence of β -D-glucose, N-acetylglucosamine, and N-acetylgalactosamine, suggests that the lectin has limited binding specificities for these carbohydrates.

6.4 Summary

Sequence data collected from a 35 kDa protein isolated from the serum of *Alligator mississippiensis*, using a mannose affinity column, indicated that it is a lectin protein comprising approximately 313 amino acids. Mass spectrometry analysis of different enzymatic digests, from

both in-gel and in-solution, was used to generate peptide sequences by MS/MS and *de novo* sequencing as well as database searching with Mascot and BLAST. Using sequence alignment tools, peptides from the lectin protein were aligned with lectin proteins from other vertebrates and peptides from the different digests were overlapped to assemble the protein sequence.

The primary structure of the alligator lectin exhibited more than 50% similarity to human, mouse, rat and frog intelectin-1 protein and less than 15% homology to mannose-binding lectin from other vertebrates. Although the alligator lectin showed a higher degree of similarity to intelectin protein, the differences of the conserved cysteine residues between the alligator and other vertebrate intelectins may result in structural differences. Strong homology to the fibrinogen-like domain of the alligator lectin was observed near the N-terminus region. There were no putative N-glycosylation sites identified in the alligator lectin sequence; however it has potential O-linkages. The isolated alligator lectin exhibited strong binding affinities toward mannose and mannan as compared to other tested carbohydrates.

CHAPTER 7. A MASS SPECTROMETRY APPROACH FOR THE STUDY OF DEGLYCOSYLATED PROTEINS*

The work described in this chapter focuses on preparatory steps to remove salts and buffers for mass spectrometry (MS) analysis after enzymatic or chemical deglycosylation. In this work, the glycosylated protein fetuin and a lectin protein isolated from the serum of *Alligator mississippiensis* were used to evaluate methods for desalting samples after an enzymatic or chemical deglycosylation. Precipitation and dialysis were used to prepare the deglycosylated samples for MS analysis. Both the precipitation and dialysis methods were suitable for sample preparation prior to analysis by matrix assisted laser desorption ionization (MALDI) MS.

7.1 Introduction

Proteomics is the study of the structure and functions of proteins in an organism or tissue, and mass spectrometry is now a routinely used method for protein identification in complex mixtures.¹²⁵ Prior to a typical mass spectrometry analysis, separation techniques such as two-dimensional gel electrophoresis and high performance liquid chromatography (HPLC) are used to reduce the complexity of protein mixtures.^{125, 325}

Some proteins are post-translationally modified to help regulate gene expression, protein turnover, and cellular structure.³⁴³ Glycosylation is a common post-translational modification that is involved in biological functions such as immune recognition and inflammation.³⁴⁴ There are two major types of protein carbohydrate linkages: N- and O-linked glycosylation. With N-glycosylation, the glycan is attached to an asparagine residue followed by any amino acid other than proline, which is linked to a serine or threonine residue. During O-glycosylation, the glycan is attached to a serine or threonine residue. Glycoproteins have been characterized by mass spectrometry, but this is challenging due to the heterogeneity of the carbohydrate moieties and

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the complexity of the resulting mass spectra.³⁴⁵ Glycosylation of the protein may also limit the extent of proteolytic digestion. To avoid this, the N- and/or O-linked carbohydrates can be removed prior to analysis.^{346, 347}

Enzymatic or chemical removal of the glycan can be used for analysis of the protein without the attached oligosaccharides. An enzymatic approach using peptide: N-glycosidase (PNGase F)²⁰⁴ and a chemical approach using trifluoromethanesulfonic acid (TFMS) have been used to remove the oligosaccharides.^{205, 348} PNGase F is an enzyme that removes N-linked oligosaccharides, leaving both the protein and oligosaccharides intact for further analysis. There is no enzyme that cleaves O-linked oligosaccharides. TFMS is a non-specific deglycosylating agent that cleaves both N-linked and O-linked oligosaccharides to leave the intact protein.³⁴⁹

The proteins used in this study were enzymatically or chemically deglycosylated followed by matrix assisted laser desorption ionization (MALDI) MS. The steps for both chemical and enzymatic deglycosylation have been reported previously,^{205, 350, 351} but the deglycosylated product requires cleanup prior to MALDI mass spectrometry. We have developed important sample desalting procedures that follow deglycosylation prior to mass spectrometry analysis, which are not clearly outlined in many procedures. This study focuses on the treatment of deglycosylated proteins for mass spectrometry analysis.

7.2 Experimental

7.2.1 Materials

Fetuin, a plasma glycoprotein produced by the liver, was obtained from Sigma-Aldrich and the lectin protein was isolated from the blood plasma of *Alligator mississippiensis*. Blood from seven American alligators was collected, heparin added for anticoagulation, and the blood allowed to clot overnight at ambient temperature. The serum was collected and the lectin was isolated using mannan-agarose affinity chromatography.

7.2.2 Enzymatic Deglycosylation

Proteins containing oligosaccharides attached through N-linked glycosylation were deglycosylated as described in Section 2.9. Approximately 100 μg of protein was dissolved in 18 μl of nanopure water. Denaturing buffer (2 μl), containing 0.6 mg/mL SDS and 0.5 mg/mL DTT, was added and heated at 100°C for 10 minutes to increase the rate of deglycosylation. The sample was incubated on ice at 0°C for five minutes and centrifuged for 10 sec. To the denatured sample, a non-ionic detergent (NP-40) was added to prevent loss of enzyme activity. The PNGaseF enzyme was added to the sample and it was incubated at 37°C for four hours.^{204, 352} The intact protein was isolated from the mixture via protein precipitation, which removed the buffers that can interfere with the MALDI signal. Acetone or trichloroacetic acid (TCA) was used to precipitate the protein.³⁵³

7.2.3 TCA Precipitation

TCA precipitation was accomplished by the addition of one volume of 60% TCA. The sample was gently vortexed and incubated on ice overnight. The sample mixture was centrifuged at 10,000 g for 30 min and the supernatant was removed. Chilled acetone was added to wash the precipitate and it was centrifuged again at 10,000 g.⁸⁰ This procedure was repeated twice. The precipitate was air-dried and suspended in aqueous 0.1% trifluoroacetic acid (TFA) or 0.1% formic acid (FA) for MALDI analysis.

7.2.4 Acetone Precipitation

Acetone precipitation was achieved by the addition of four volumes of acetone to the protein sample, followed by an overnight incubation on ice. The sample was centrifuged at 10,000 g for 30 min at 4°C. The supernatant was removed and the precipitate was air dried for 30 min, then suspended in the same buffer as used above in the TCA precipitation.

7.2.5 Chemical Deglycosylation

In-solution deglycosylation was performed using a deglycosylation kit (Prozyme) containing trifluoromethanesulphonic acid (TFMS). TFMS removes most glycan types from glycoproteins and leaves the protein intact.²⁰⁵ Briefly, approximately 0.3 mg of protein in a vial was dried overnight using a lyophilizer. The dried protein was capped with a septum and placed on a dry ice-ethanol bath for 20 seconds to reduce the temperature of the reaction. A mixture of TFMS and toluene solution was prepared and 50 μl was slowly added using a gas-tight syringe. Precautions were taken to prevent the dried protein from being exposed to moisture due to the potential for hydrolysis of the peptide bonds.^{205, 354} The vials containing the protein with TFMS mixture were incubated at -20°C for five minutes then shaken briefly to help dissolve the protein. The solvation process was repeated and followed by a four-hour incubation at -20°C .

7.2.6 Dialysis

Dialysis was performed prior to mass spectrometry analysis. After the four-hour incubation, the vials containing the samples were placed in a dry ice-ethanol bath for 20 seconds, and then 150 μl of pyridine solution was added to each vial. The vials were incubated on dry ice for five minutes, and then in a dry ice-ethanol bath for 15 minutes. A 400 μl quantity of 0.5% ammonium bicarbonate was added to each vial to neutralize the sample. Following deglycosylation, the protein was isolated using dialysis. Each sample was added to a dialysis cassette (Thermo Scientific, Rockford, IL, USA) and dialyzed against 10 mM ammonium formate for two hours at room temperature. The buffer was changed and dialysis was continued for an additional two hours. After a final buffer change, the sample was dialyzed overnight at room temperature.

7.2.7 Mass Spectrometry

Glycoproteins and deglycosylated proteins were analyzed in linear MS mode with the Bruker UltrafleXtreme MALDI mass spectrometer. The glycoproteins and the deglycosylated proteins (after dialysis or precipitation) were mixed on a steel target using a 1:1 ratio of sample and saturated matrix solution. The matrices used were 2-hydroxy-5-methoxybenzoic acid (sDHB) and 2,4-dimethoxy-3-hydroxycinnamic acid (sinapinic acid, SA). The samples were analyzed in positive ion mode and 500 shot averages were collected using a mass range from 30 to 100 kDa. Masses were determined using FlexAnalysis software.

7.3 Results and Discussion

Two methods were used for deglycosylation: PNGaseF and trifluoromethanesulfonic acid. The N- and O-linked glycoprotein protein, fetuin, was enzymatically deglycosylated to remove N-linked oligosaccharides and chemical deglycosylation was used to remove both N- and O-linked oligosaccharides. Fetuin is a glycoprotein with 359 amino acids containing N-linked glycosylation sites at Asn 99, 156 and 176, and numerous O-linked glycosylation sites.³⁵⁵ As seen in Figure 7-1a, a mass of 45.3 kDa was measured for fetuin before deglycosylation which is consistent with previous studies.³⁵⁶ The deglycosylated fetuin was analyzed with MALDI after deglycosylation with PNGaseF and no further cleanup. No signal was observed possibly due to buffers used during the deglycosylation. Using acetone and TCA precipitation independently, the protein was separated from the PNGaseF released oligosaccharides. The deglycosylated protein from the pellet of fetuin gave a 39.5 kDa in the mass spectrum and was compared with the glycosylated fetuin using MALDI and a mass shift of 5.8 kDa was observed in the mass spectrum (Figure 7-1b). Both acetone and TCA precipitation yielded sufficient protein recovery for MALDI analysis and the MALDI results were comparable.

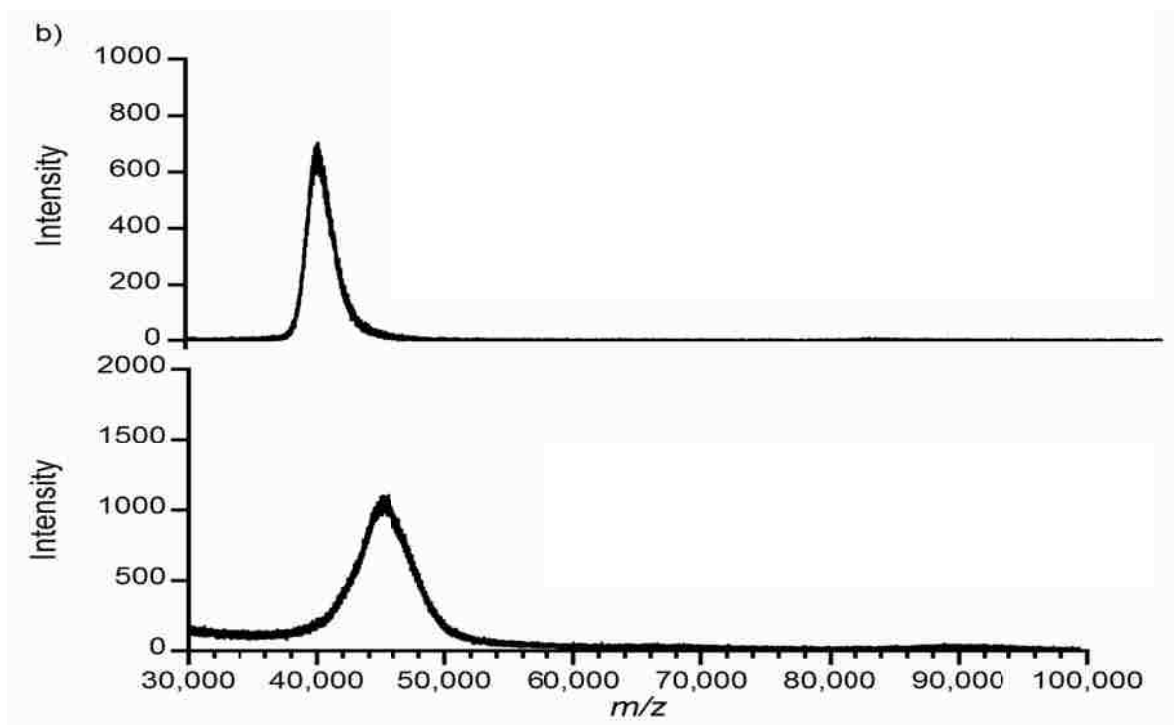
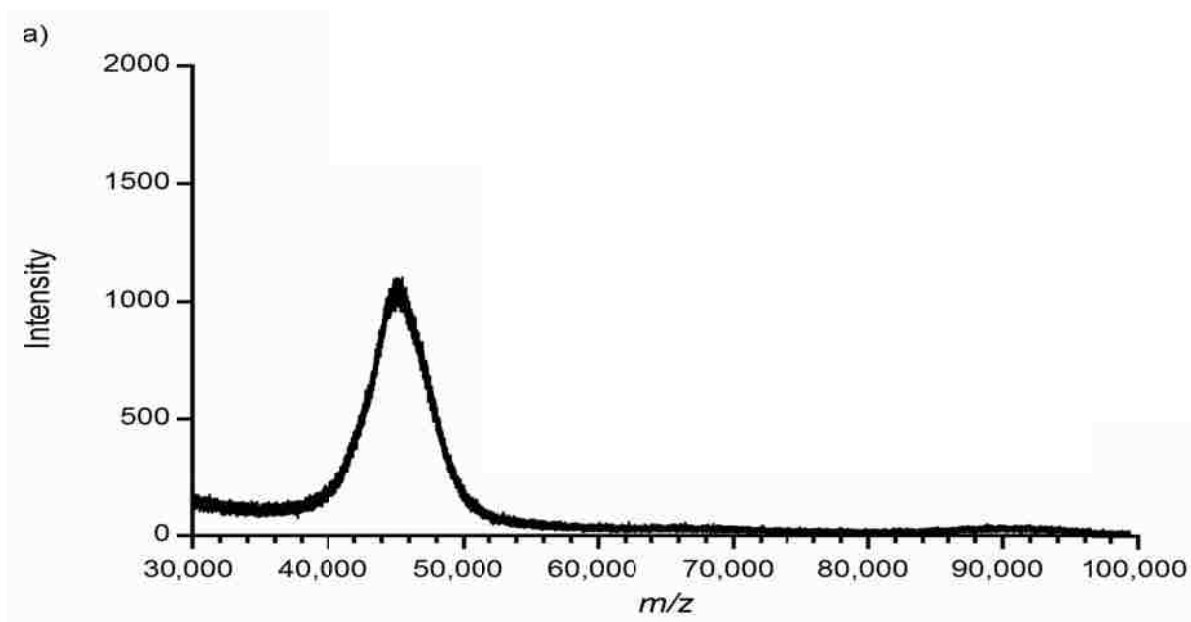
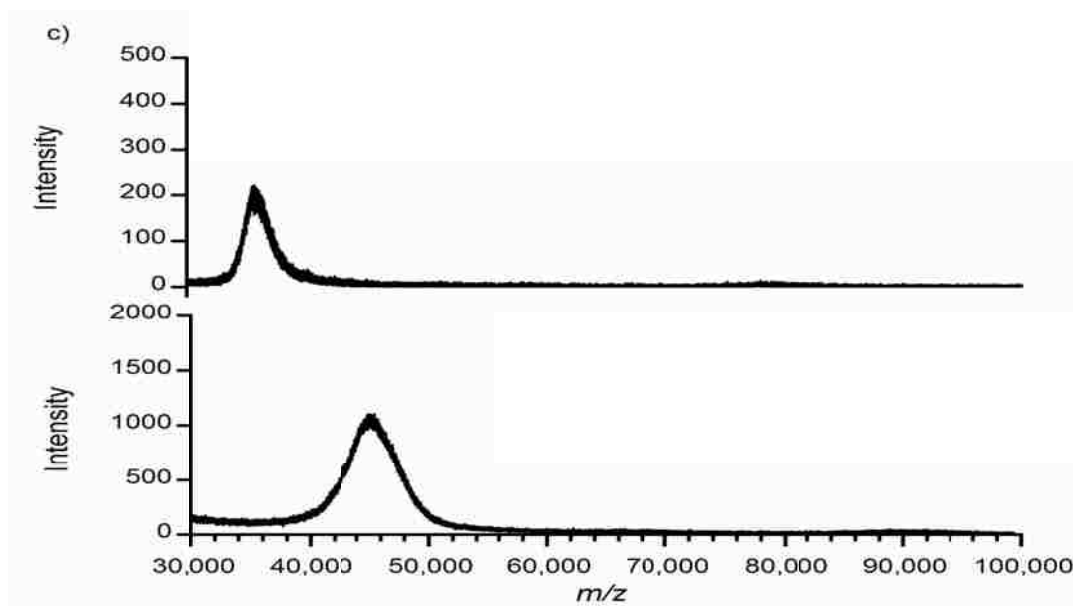


Figure 7-1. MALDI mass spectrometry of fetuin using SA matrix: (a) MALDI spectrum of intact fetuin, (b) MALDI spectra comparing glycosylated and deglycosylated fetuin after acetone precipitation, (c) fetuin after TFMS deglycosylation followed with dialysis.

Figure 7-1. cont'd.



The chemical deglycosylation method using trifluoromethanesulfonic acid was also used to treat the fetuin protein. Following the deglycosylation, a MALDI analysis showed no signal in the protein mass range. However, following dialysis (which was used to desalt and concentrate the deglycosylated protein) a signal was observed. Compared to the precipitation methods, dialysis was more time-consuming and the protein signal observed was lower (Figure 7-1c). However, it removed interfering salts and produced a MALDI signal, although it was not as intense as the signal obtained from the precipitated deglycosylated protein.

This approach was applied to an alligator lectin protein. The non-specific deglycosylation method using TFMS as described above was used to remove oligosaccharides from the protein. The intact protein mass was measured at 34.6 kDa before deglycosylation. After deglycosylation and dialysis a mass of 32.2 kDa was observed in the MALDI mass spectrum (Figure 7-2).

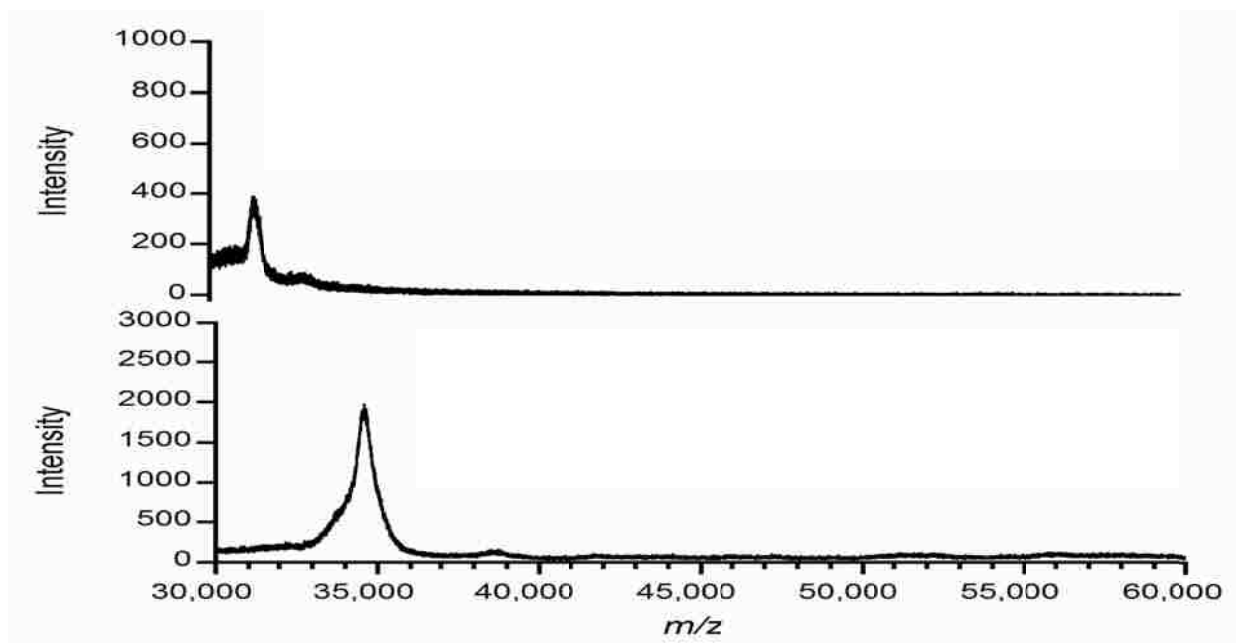


Figure 7-2. Mass spectrometry data before (above) and after (below) deglycosylating the lectin protein using SA matrix.

The mass shift suggests the loss of oligosaccharides and the less intense signal indicates lower protein recovery from the dialysis procedure.

7.4 Summary

Appropriate sample preparation is necessary to obtain mass spectrometry data after deglycosylation. To demonstrate the necessity for sample cleanup prior to mass spectrometry, we have used a glycosylated protein, fetuin, and a lectin protein isolated from *Alligator mississippiensis* serum. Protein precipitation with acetone and TCA was used following deglycosylation to concentrate and desalt the proteins. Alternately, dialysis was used to purify the protein from the other unwanted substances, as demonstrated with the chemical deglycosylation procedure. The precipitation and dialysis methods resulted in efficient desalting and protein concentration. However, the precipitation methods produced a higher protein signal as compared to dialysis.

CHAPTER 8. CONCLUSIONS AND FUTURE DIRECTIONS

In this dissertation, mass spectrometry based proteomics was used to study the alligator blood proteome. Proteins and peptides found in the leukocyte extracts from the alligators were identified using gel electrophoresis techniques, liquid chromatography, and mass spectrometry sequencing, including bottom-up and top-down sequencing. Several peptides and proteins with immunological functions, including lectin and antimicrobial peptides were isolated and identified. The significance of this work lies in that it provides insight on the function and evolution of both innate and adaptive immune systems of crocodylians.

In Chapter 3, the proteome of the leukocytes was investigated because they are one of the most important cells in the body to participate at the site of an infection and play an important role in innate immunity. A 1- and 2-D gel electrophoresis system was used to separate leukocyte proteins, and enzymatic digest of the protein bands and spots using trypsin was applied to cleave the proteins to peptides for peptide sequencing. Using Mascot, a comparison was made between the experimentally observed fragment ions and the predicted fragments for all hypothetical peptides of a particular mass. From the MASOT search, 17 proteins were identified based on sequence homology. For MS/MS data that did not match any proteins in the database, peptide sequences were determined using *de novo* sequencing. Using the *de novo* sequencing results, 23 proteins were identified based on peptide matching using BLAST.

To further investigate peptides from the acidic leukocyte extracts that exhibited antimicrobial activity, a series of separations and antimicrobial tests were performed. Using reversed-phase chromatography, the leukocyte extracts (which consisted of a mixture of peptides and proteins) were separated, collected, and tested for antimicrobial activity. The initial eluents collected within the first 10 min exhibited activity and were further characterized using mass spectrometry and X-ray crystallography for identification, which was described in Chapter 4.

Small molecules less than 500 Da were identified, and the results indicated that the anticoagulant, EDTA, which was used during the blood collection, may have contributed to the antimicrobial activity. Due to the antimicrobial activity interferences from EDTA used in the work in Chapter 4, heparin was used as an alternative anticoagulant for the studies described in Chapter 5.

To investigate the specific peptides and proteins from the alligator leukocytes that exhibit antimicrobial activity, peptides and proteins isolated from alligator leukocyte extracts were separated by reversed phase chromatography as described in Chapter 5. The antimicrobial activity of the peptides from the chromatographic fractions was tested for growth inhibition of various microbes to determine the antimicrobial spectrum of the peptides. Among the microbes studied, growth inhibition was observed for *E. coli*, *E. cloacae* and *K. oxytoca* indicating that the antimicrobial peptides from the alligator are active against gram-negative bacteria. Within a fraction that exhibited antimicrobial activity, two peptides at masses 4.7 and 4.9 kDa were identified using mass spectrometry. Due to the close mass of the peptides, ion mobility was used to determine whether the peptides can be separated based on their charge, size and shape. The separated peptide was analyzed using tandem mass spectrometry; however, limited sequence information was determined due to incomplete fragmentation. Therefore, the peptides were subjected to MALDI MS/MS for sequencing. Peptide sequences were determined using *de novo* and manual sequencing. From this work, the first antimicrobial peptides were isolated and partially characterized from *Alligator mississippiensis* and showed to have a net charge of +8 and +13 respectively at pH 5.5, indicating that they are cationic. Based on size and charge, these peptides exhibit strong characteristics similar to previously identified antimicrobial peptides.

In Chapter 6, another class of immunological proteins, lectins, was studied which provides insight on the structure-function relationship of another class of proteins involved

within crocodile's innate immune system. A new calcium-dependent binding lectin protein was isolated from the serum of the *Alligator mississippiensis* by affinity chromatography using mannose columns. The sequence of this novel class of proteins from the blood serum of the alligator was determined with mass spectrometry and Edman sequencing and was used for homologous comparison with other lectins. Molecular masses of the isolated lectin were determined by ESI-MS and were found to be oligomeric which are consistent with the monomer, dimer, trimer and tetramer of the 35 kDa monomer lectin protein. The monomeric protein was enzymatically digested using four different proteases, creating small and large peptides, which were overlapped to enhance protein sequence coverage. The primary sequence of a novel lectin protein isolated from *Alligator mississippiensis* was determined, and its binding specificity to selected carbohydrates was studied.

An approach to analyze proteins that have been glycosylated also was evaluated. In Chapter 7, preparatory steps used after deglycosylation procedures and before mass spectrometry analysis were described. Here, glycosylated proteins, fetuin and alligator lectin, were deglycosylated and methods for sample desalting and concentrating, such as precipitation and dialysis were used. The dialysis and precipitation methods were both suitable for sample preparation prior to MS analysis. A comparison between precipitation and dialysis procedures showed that precipitation methods produce higher protein signal, indicating a larger protein recovery.

One of the future directions is to investigate other properties of the peptides, other than bacterial properties. Some antimicrobial peptides have shown to have anti-inflammatory, antifungal, antiviral and anti-tumor activity, so these properties should be investigated. Prior work showed that the crude leukocyte extract exhibited antimicrobial activity, so it is hypothesized that the activity may be due to multiple peptides. Therefore, the alligator leukocyte

extract should be further investigated for other antimicrobial peptides or proteins, other than the two peptides isolated in this work and their antimicrobial activity should be characterized, including the minimal inhibitory concentration (MIC) and the antimicrobial spectrum.

The interaction of alligator antimicrobial peptides in the presence of microbes should also be investigated using a technique such as atomic force microscopy (AFM). This study can help provide an understanding of how alligator antimicrobial peptides affect bacterial cells. This effect may be based on the different models proposed as discussed in Section 1.1.1., for example, the carpet model, where peptides cover the membrane like a carpet and destroy the bacteria lipid bilayer.³⁵⁷ By reducing the AFM cantilever size to approximately 1000 times smaller than the normal size, the imaging speed can be increased without destroying the bacteria.³⁵⁸ The bacteria can be immobilized on poly-L-lysine coated cover slides (which will provide stronger cell adhesion) and imaged in aqueous solution to keep the cells alive. Alligator antimicrobial peptide can be added to the liquid droplet comprised of bacteria, at a concentration above the minimal inhibitory concentration (MIC) to acquire images.³⁵⁹ It is also necessary to evaluate the antimicrobial peptides presented in this dissertation for their suitability in commercial products, such as topical creams. They can be added to existing products to enhance the antimicrobial activity of the product and/or they can potentially be used as an antiseptic cream for cuts or an antifungal cream for infections such as athlete's foot.

The research in this dissertation also showed that lectin proteins are present and can be isolated from alligators. Therefore, additional studies should be performed to further characterize the identified lectin in this work, such as investigating the temperature and pH effect, and the effect of the identified alligator lectin on bacteria induced hemagglutination activity; which will indicate whether or not the lectin protects against bacterial induced erythrocyte damage. Because carbohydrates play a vital role in biological processes and diseases, lectins can be used to

identify certain diseases. Depending on the specificity of the alligator lectin it can potentially be used as a means to identify a particular disease state. Finally, antibodies against the alligator lectin discussed in this work can be prepared and used for isolation of lectins from other crocodilian species. This will provide more insight on lectins and their function within crocodilian species.

Lastly, an exhaustive MS-based proteomics study should be performed to map all the proteins involved in the alligator's immune system. The immune system is complex and made of several types of cells and proteins that play different roles in fighting infections. Therefore, a more comprehensive study of the structure-function relationship of the immune related proteins will help provide a better understanding of crocodilian immunity as well as provide more insight on the evolutionary development of reptile's immune system.

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APPENDIX A. PROTEINS MATCHED FROM *DE NOVO* SEQUENCING

The peptide sequences in the table were generated from MS/MS data using *de novo* sequencing. *De novo* peptide sequences were subjected to database searching using BLAST to find proteins that contain sequences matching with the determined *de novo* sequences. The results were used to identify biological components that comprise the alligator leukocyte proteome (Chapter 3, Section 3.3).

Table A.1. Proteins identified at single peptide level using BLAST. *De novo* sequenced peptides obtained from gel digest were searched using BLAST and proteins were matched based on sequence similarity and evolutionary relationship using the SwissProt and MSDB database. The top three E-values are reported.

Band Number	<i>m/z</i>	Charge	<i>De novo</i> Sequence	Protein	Accession Number	Protein Length	Organism	E-Value				
B4	1060.6	+2	EINLSPDSTSAVVSGLMVATK	Fibronectin	P11722.3	1256	Gallus gallus	2e-11 (1 st hit)				
			EINLSPDSTSAVVSGLMVATK									
			EINLSPDSTSAVV+GLMVATK						Q91289.1	1328	Pleurodeles waltl	9e-11 (2 nd hit)
			EINLSPDSTS+VSGLMVATK	Fibronectin	P04937.2	2477	Rattus norvegicus	7e-10 (3 rd hit)				
B4	536.3	+2	YEVSVYALK	Fibronectin	P07589.4	2478	Bos taurus	0.33 (1 st hit)				
			YEVSVYALK									
			YEVSVYALK						P11722.3	1256	Gallus gallus	0.33 (2 nd hit)
			YEVSVYALK						P11276.3	2477	Mus musculus	0.33 (3 rd hit)
			YEVSVYALK						P04937.2	2477	Rattus norvegicus	0.33 (4 th hit)
			YEVSVYALK						Q91289.1	1328	Pleurodeles waltl	0.33 (5 th hit)

Table A.1. cont'd.

			YEVSVYALK	Fibronectin	Q28275.2	522	Canis lupus familiaris	0.33 (6 th hit)
			YEVSVYALK	Fibronectin	Q28377.2	522	Equus caballus	0.33 (7 th hit)
			YEVSVYALK	Fibronectin	P02751.4	2386	Homo sapiens	0.33 (8 th hit)
			YEV+VYALK	Fibronectin	Q91740.1	2481	Xenopus laevis	1.9 (9 th hit)
			Y+VSVYA	Collagen alpha-1(XII) chain	Q60847.3	3120	Mus musculus	117 (10 th hit)
			Y VS+YALK	RNA 2-thiouridine synthesizing protein B	NP_240337.2	95	Buchnera aphidicola	117 (11 th hit)
			YEVSVY	Neuronal acetylcholine receptor subunit β-4	NP_434693.1	495	Rattus norvegicus	117 (12 th hit)
			YEVSVY	Neuronal acetylcholine receptor subunit β-4	NP_000741.1	498	Homo sapiens	117 (13 th hit)
			YEVSVY	Neuronal acetylcholine receptor subunit β-4	NP_683746.1	495	Mus musculus	117 (14 th hit)
B4	701.3	+2	YQINQQWER					
			YQINQQWER	Fibronectin	P07589.4	2478	Bos taurus	0.013 (1 st hit)
			YQINQQWER	Fibronectin	P11276.3	2477	Mus musculus	0.013 (2 nd hit)
			YQINQQWER	Fibronectin	Q91740.1	2481	Xenopus laevis	0.013 (3 rd hit)
			YQINQQWER	Fibronectin	P04937.2	2477	Rattus norvegicus	0.013 (4 th hit)
			YQINQQWER	Fibronectin	P02751.4	2386	Homo sapiens	0.013 (5 th hit)
			INQQWE	Protein lava lamp	NP_525064.1	2779	Drosophila melanogaster	27 (6 th hit)
			YQINQQ	DNA topoisomerase 2- associated protein pat1	NP_595976.2	754	Schizosaccharomyces pombe	65 (7 th hit)
			YQINQQ	Dynammin-like protein C	XP_645576.2	904	Dictyostelium discoideum	65 (8 th hit)
			YQINQQ	Ornithine	YP_672340.1	326	Escherichia coli 536	65

Table A.1. cont'd.

			carbamoyltransferase				(9 th hit)
			Protein Cpn60	NP_849007.1	526	Cyanidioschyzon merolae	65 (10 th hit)
			Thymidylate kinase	NP_077850.1	230	Ureaplasma parvum	65 (11 th hit)
			Ornithine carbamoyltransferase	NP_757196.2	334	Escherichia coli O6	65 (12 th hit)
			Exodeoxyribonuclease V gamma chain	ZP_07124340.1	1122	Escherichia coli K-12	65 (13 th hit)
B1	613.9	+2	LIALLEVLSQK				
			LIALLEVLSQK	NP_001449.3	2725	Homo sapiens	0.019 (1 st hit)
			LIALLEVLSQK	NP_001074654.1	2726	Mus musculus	0.019 (2 nd hit)
			LIALLEVLSQK	Q80X90.2	2602	Mus musculus	0.019 (3 rd hit)
			LIALLEVLSQK	CAT00728.1	2647	Mus musculus	0.019 (4 th hit)
			LIALLEVLSQK	NP_001104026.1	2647	Homo sapiens	0.019 (5 th hit)
			LIALLEVLSQK	NP_001448.2	2602	Homo sapiens	0.019 (6 th hit)
			+IALL+VLSQ	A5DIV5.2	461	Pichia guilliermondii	5.1 (7 th hit)
			L+ALLEVLS QK	NP_892006.2	8797	Homo sapiens	5.1 (8 th hit)
			LI+LLEVLS	AAK83382.1	1678	Mus musculus	9.2 (9 th hit)
B1	487.3	+2	VYGPVGVEPR				
			+YGPVGVEP	YP_076110.1	205	Symbiobacterium thermophilum	15 (1 st hit)
			+YGPVGVEP	YP_076123.1	194	Symbiobacterium thermophilum	15 (2 nd hit)
			VYGPVGVE	NP_001449.3	2725	Homo sapiens	36 (3 rd hit)
			VYGPVGVE	NP_001074654.1	2726	Mus musculus	36 (4 th hit)
			VYGPVGVE	P13466.1	857	Dietyostelium discoideum	36 (5 th hit)
			GPGVEPR	B9J8B4.1	217	Thermosipho africanus TCF52B	49 (6 th hit)

Table A.1. cont'd.

			VYGP VEP	50S ribosomal protein L25	B7IFM6.1	217	Thermosipho africanus TCF52B	49 (7 th hit)
B2	736.4	+2	AQQVSQGLDLLTAK AQQVSQGLDLLTAK	Vinculin	P12003.4	1135	Gallus gallus	5e-05 (1 st hit)
			AQQVSQGLD+LTAK	Vinculin	P85972.1	1066	Rattus norvegicus	7e-04 (2 nd hit)
			AQQVSQGLD+LTAK	Vinculin	P26234.4	1135	Sus scrofa	7e-04 (3 rd hit)
			AQQVSQGLD+LTAK	Vinculin	Q64727.4	1066	Mus musculus	7e-04 (4 th hit)
			AQQVSQGLD+LTAK	Vinculin	P18206.4	1134	Homo sapiens	7e-04 (5 th hit)
			QV+QGLD L AK	Macrophage colony-stimulating factor 1 receptor	Q9I8N6.1	977	Danio rerio	3.8 (6 th hit)
B2	739.4	+2	MLGQMTDQVADLR MLGQMTDQVADLR	Vinculin	P85972.1	1066	Rattus norvegicus	2e-05 (1 st hit)
			MLGQMTDQVADLR	Vinculin	P26234.4	1135	Sus scrofa	2e-05 (2 nd hit)
			MLGQMTDQVADLR	Vinculin	Q64727.4	1066	Mus musculus	2e-05 (3 rd hit)
			MLGQMTDQVADLR	Vinculin	P18206.4	1134	Homo sapiens	2e-05 (4 th hit)
			LGQMTDQ+ADLR	Vinculin	P12003.4	1135	Gallus gallus	0.006 (5 th hit)
			MLGQM +QV	Uncharacterized protein SE_2353	Q8CQQ7.1	300	Staphylococcus epidermidis ATCC 12228	6.8 (6 th hit)
			L QMT D +ADLR	Pre-mRNA-processing ATP-dependent RNA helicase	Q2HAD8.1	1064	Chaetomium globosum	6.8 (7 th hit)
B2	655.9	+2	TVTAMDVVYALK TVTAMDVVYALK	Histone H4	P83865.2	103	Litopenaeus vannamei	6e-04 (1 st hit)
			TVTAMDVVYALK	Histone H4	P82888.2	103	Olithodiscus luteus	6e-04 (2 nd hit)
			TVTAMDVVYALK	Histone H4	P02310.2	103	Tetrahymena pyriformis	6e-04 (3 rd hit)
			TVTAMDVVYALK	Histone H4	Q6WV74.3	103	Mytilus chilensis	6e-04 (4 th hit)
			TVTAMDVVYALK	Histone H4	Q8MTV8.3	103	Aplysia californica	6e-04 (5 th hit)

Table A.1. cont'd.

TVTAMDVVYALK	Histone H4	P50566.2	103	<i>Chlamydomonas reinhardtii</i>	6e-04 (6 th hit)
TVTAMDVVYALK	Histone H4	P91890.3	103	<i>Trichogramma cacoeciae</i>	6e-04 (7 th hit)
TVTAMDVVYALK	Histone H4	P91882.3	103	<i>Diadromus pulchellus</i>	6e-04 (8 th hit)
TVTAMDVVYALK	Histone H4	P70081.3	103	<i>Gallus gallus</i>	6e-04 (9 th hit)
TVTAMDVVYALK	Histone H4	Q43083.3	103	<i>Pyrenomonas salina</i>	6e-04 (10 th hit)
TVTAMDVVYALK	Histone H4	P91849.3	103	<i>Apis mellifera</i>	6e-04 (11 th hit)
TVTAMDVVYALK	Histone H4	P62786.2	103	<i>Triticum aestivum</i>	6e-04 (12 th hit)
TVTAMDVVYALK	Histone H4	P59259.2	103	<i>Arabidopsis thaliana</i>	6e-04 (13 th hit)
TVTAMDVVYALK	Histone H4	P08436.2	103	<i>Volvox carteri</i>	6e-04 (14 th hit)
TVTAMDVVYALK	Histone H4	Q6WV72.3	103	<i>Mytilus trossulus</i>	6e-04 (15 th hit)
TVTAMDVVYALK	Histone H4	P62776.2	103	<i>Holothuria tubulosa</i>	6e-04 (16 th hit)
TVTAMDVVYALK	Histone H4	P69151.2	103	<i>Tetrahymena pyriformis</i>	6e-04 (17 th hit)
TVTAMDVVYALK	Histone H4	P35057.2	103	<i>Solanum lycopersicum</i>	6e-04 (18 th hit)
TVTAMDVVYALK	Histone H4	P35059.2	103	<i>Acropora Formosa</i>	6e-04 (19 th hit)
TVTAMDVVYALK	Histone H4	Q27443.2	103	<i>Ascaris suum</i>	6e-04 (20 th hit)
TVTAMDVVYALK	Histone H4	Q9U7D0.3	108	<i>Mastigamoeba balamuthi</i>	6e-04 (21 st hit)
TVTAMDVVYALK	Histone H4	Q41811.3	103	<i>Zea mays</i>	6e-04 (22 nd hit)
TVTAMDVVYALK	Histone H4	P04915.2	103	<i>Physarum polycephalum</i>	6e-04 (23 rd hit)
TVTAMDVVYALK	Histone H4	P84040.2	103	<i>Drosophila melanogaster</i>	6e-04 (24 th hit)
TVTAMDVVYALK	Histone H4	P40287.1	118	<i>Entamoeba histolytica</i>	6e-04

Table A.1. cont'd.

			TVTAMDVVYALK	Histone H4	Q76NW2.1	108	Dictyostelium discoideum	6e-04 (25 th hit)
			TVT+MDVVYALK	Histone H4	P27996.2	103	Solaster stimpsoni	0.003 (27 th hit)
			TVTA+DVVYALK	Histone H4	P80737.3	103	Blepharisma japonicum	0.011 (28 th hit)
			TVTA+DVVYALK	Histone H4	P62790.2	104	Sterkiella nova	0.011 (29 th hit)
			TVTA+DVVYALK	Histone H4	Q27765.3	103	Styela plicata	0.011 (30 th hit)
			TVTA+DVVYALK	Histone H4	P90516.1	89	Blepharisma japonicum	0.011 (31 st hit)
			TVTA+DVVYALK	Histone H4	P62792.2	103	Phanerochaete chrysosporium	0.011 (32 nd hit)
			TVTA+DVVYALK	Histone H4	P80739.3	107	Moneuplotes crassus	0.011 (33 rd hit)
B2	628.7	+2	LITKAVSASK					
			LITKAVSASK	Histone H1.5	P43276.2	223	Mus musculus	0.49 (1 st hit)
			LITKAVSASK	Histone H1.03	P08285.2	224	Gallus gallus	0.49 (2 nd hit)
			LITKAVSASK	Histone H1.10	P08286.3	220	Gallus gallus	0.49 (3 rd hit)
			LITKAVSASK	Histone H1.11	P08287.2	225	Gallus gallus	0.49 (4 th hit)
			LITKAVSASK	Histone H1.01	P08284.2	219	Gallus gallus	0.49 (5 th hit)
			LITKAVSASK	Histone H1	P09987.2	218	Gallus gallus	0.49 (6 th hit)
			LITKAVSASK	Histone H1.11R	P08288.2	219	Gallus gallus	0.49 (7 th hit)
			LITKAV+ASK	Histone H1	P16403.2	213	Homo sapiens	2.9 (8 th hit)
			LITKAV+ASK	Histone H1.3	P02251.1	213	Cryptolagus cuniculus	2.9 (9 th hit)
			LITKAV+ASK	Histone H1.4	P10412.2	219	Homo sapiens	2.9 (10 th hit)
			LITKAV+ASK	Histone H1.3	P16402.2	221	Homo sapiens	2.9 (11 th hit)

Table A.1. cont'd.

			LITKAV+ASK	Histone H1.1	P02253.1	104	Bos taurus	2.9 (12 th hit)
			LITKAV+ASK	Histone H1.2	P15865.3	219	Rattus norvegicus	2.9 (13 th hit)
			LITKAV+ASK	Histone H1.2	P15864.2	212	Mus musculus	2.9 (14 th hit)
			LITKAV+ASK	Histone H1.4	P02252.1	73	Oryctolagus cuniculus	2.9 (15 th hit)
			LITKAV+ASK	Histone H1.4	P43274.2	219	Mus musculus	2.9 (16 th hit)
			LITKAV+ASK	Histone H1	P09426.3	218	Anas platyrhynchos	2.9 (17 th hit)
			LITKAV+ASK	Histone H1.5	P16401.3	226	Homo sapiens	2.9 (18 th hit)
			LITKAV+ASK	Histone H1	P43277.2	221	Mus musculus	2.9 (19 th hit)
			LI KAVSASK	Histone H1	P84408.3	55	Salmo salar	9.3 (20 th hit)
			LI KAVSASK	Histone H1	P06350.2	207	Oncorhynchus	9.3 (21 st hit)
B6	833.9	+2	FSGSGSGTDFFTISR					
			FSGSGSGTDFFTIS	Ig K chain V-I region Lay	P01605.1	108	Homo sapiens	5e-06 (1 st hit)
			FSGSGSGTDFFTIS	Ig K chain V-I region WAT	P80362.1	108	Homo sapiens	5e-06 (2 nd hit)
			FSGSGSGT+FTFTIS	Ig K chain V-I region OU	P01606.1	108	Homo sapiens	9e-06 (3 rd hit)
			FSGSGSGTDFFTISR	Ig K chain V-III region HIC	P18136.1	129	Homo sapiens	2e-05 (4 th hit)
			FSGSGSGTDFFTISR	Ig K chain V-III region Ti	P01622.1	109	Homo sapiens	2e-05 (5 th hit)
			FSGSGSGTDFFTISR	Ig K chain V-III region SIE	P01620.1	109	Homo sapiens	2e-05 (6 th hit)
			FSGSGSGTDFFTISR	Ig K chain V-III region HAH	P18135.1	129	Homo sapiens	2e-05 (7 th hit)
			FSGSGSGTDFFTISR	Ig K chain V-III region WOL	P01623.1	109	Homo sapiens	2e-05 (8 th hit)
			FSGSGSGTDFFTISR	Ig K chain V-III region GOL	P04206.1	109	Homo sapiens	2e-05 (9 th hit)
B6	751.4	+2	VFGGGTKLTVLQPK					

Table A.1. cont'd.

	VFGGGTKLTVL QPK	Ig A chain V-III region LOI	P80748.1	111	Homo sapiens	7e-05 (1 st hit)
	VFGGGT+LT+LGQPK	IA-like polypeptide 1	P20764.2	209	Mus musculus	6e-04 (2 nd hit)
	VFGGGTKLTVLG	Ig A-1 chain V regions MOPC	P01724.1	129	Mus musculus	0.006 (3 rd hit)
	VFGGGTKLTVLG	Ig A chain V-I region BL2	P06316.1	130	Homo sapiens	0.006 (4 th hit)
	VFGGGTKLTVLG	Ig A chain V-V region DEL	P01719.1	108	Homo sapiens	0.006 (5 th hit)
	VFGGGTKLTVLG	Ig A chain V-VI region EB4	P06319.1	131	Homo sapiens	0.006 (6 th hit)
	VFGGGTKLTVLG	Ig A chain V-II region VIL	P01711.1	111	Homo sapiens	0.006 (7 th hit)
	VFGGGTKLTVLG	Ig A-1 chain V region H2020	P01726.1	129	Mus musculus	0.006 (8 th hit)
	VFGGGTKLTVLG	Ig A-1 chain V region S43	P01727.1	129	Mus musculus	0.006 (9 th hit)
	VFGGGTKLTVLG	Ig A-1 chain V region S178	P01725.1	110	Mus musculus	0.006 (10 th hit)
	VFGGGTKLTVLG	Ig A chain V-VI region SUT	P06317.1	111	Homo sapiens	0.006 (11 th hit)
B2	510.8	+2	V(I/L)ASFGEAVK			
	VLASFGEAVK	Hemoglobin, β	P02131.1	146	Caiman crocodilus	0.27 (1 st hit)
	VLASFGEAVK	Hemoglobin, β	P02130.1	146	Alligator mississippiensis	0.27 (2 nd hit)
	VL+SFGEAVK	Hemoglobin, ϵ	P02128.2	147	Gallus gallus	1.6 (3 rd hit)
	VL+SFGEAVK	Hemoglobin, ρ	P02127.2	147	Gallus gallus	1.6 (4 th hit)
	VL SFGEAVK	Hemoglobin, ϵ	Q28496.3	147	Microcebus murinus	2.1 (5 th hit)
	VLASFGEAV	Hemoglobin, β	P02129.1	146	Crocodylus niloticus	2.1 (6 th hit)
	VL SFGEAVK	Hemoglobin, ϵ	P19759.2	147	Otolemur crassicaudatus	2.1 (7 th hit)
	VL SFGEAVK	Hemoglobin, β	P13274.1	146	Chrysemys picta bellii	2.1 (8 th hit)

Table A.1. cont'd.

VL SFGEAVK	Hemoglobin, β	P83133	146	<i>Dipsochelys dussumieri</i>	2.1 (9 th hit)
VL SFGEAVK	Hemoglobin, β	P02125.1	146	<i>Ciconia ciconia</i>	2.1 (10 th hit)
VL SFGEAVK	Hemoglobin, β	P14524.1	146	<i>Turdus merula</i>	2.1 (11 th hit)
VL SFGEAVK	Hemoglobin, β	P08851.1	146	<i>Accipiter gentilis</i>	2.1 (12 th hit)
VL SFGEAVK	Hemoglobin, β	P02116.1	146	<i>Ara ararauna</i>	2.1 (13 th hit)
VL SFGEAVK	Hemoglobin, β	P07411.1	146	<i>Vultur gryphus</i>	2.1 (14 th hit)
VL SFGEAVK	Hemoglobin, β	P15165.1	146	<i>Apus apus</i>	2.1 (15 th hit)
VL SFGEAVK	Hemoglobin, β	P08261.1	146	<i>Larus ridibundus</i>	2.1 (16 th hit)
VL SFGEAVK	Hemoglobin, β	P21668.1	146	<i>Psittacula krameri</i>	2.1 (17 th hit)
VL SFGEAVK	Hemoglobin, ϵ	Q95238.3	147	<i>Propithecus verreauxi</i>	2.1 (18 th hit)
VL SFGEAVK	Hemoglobin, β	P02120.1	146	<i>Anseranas semipalmata</i>	2.1 (19 th hit)
VL SFGEAVK	Hemoglobin, β	P10782.1	146	<i>Phalacrocorax carbo</i>	2.1 (20 th hit)
VL SFGEAVK	Hemoglobin, ϵ	P08223.2	146	<i>Eulemur fulvus fulvus</i>	2.1 (21 st hit)
VL SFGEAVK	Hemoglobin, β	P07406.1	146	<i>Passer montanus</i>	2.1 (22 nd hit)
VL SFGEAVK	Hemoglobin, β	P68061.1	146	<i>Aegypius monachus</i>	2.1 (23 rd hit)
VL SFGEAVK	Hemoglobin, β	P02122.1	146	<i>Aquila chrysaetos</i>	2.1 (24 th hit)
VL SFGEAVK	Hemoglobin, ϵ	Q28338.3	147	<i>Cheirogaleus medius</i>	2.1 (25 th hit)
VL SFGEAVK	Hemoglobin, β	P02109.3	147	<i>Didelphis virginiana</i>	2.1 (26 th hit)
VL SFGEAVK	Hemoglobin, β	P02121.1	146	<i>Phoenicopterus ruber ruber</i>	2.1 (27 th hit)

Table A.1. cont'd.

			VL SFGEAVK	Hemoglobin, β	Q98905.3	147	Geochelone carbonaria	2.1 (28 th hit)
			VL SFGEAVK	Hemoglobin, β	P82113.1	146	Stercorarius maccormicki	2.1 (29 th hit)
B2	500.8	+2	LSSPISGDPK					
			PISGDPK	Microtubule-actin cross-linking factor 1	Q9UPN3.3	5430	Homo sapiens	54 (1 st hit)
			PISGDPK	Microtubule-actin cross-linking factor 1, isoform 4	Q96PK2.2	5938	Homo sapiens	54 (2 nd hit)
			PISGDPK	Microtubule-actin cross-linking factor 1	Q9QXZ0.1	5327	Mus musculus	54 (3 rd hit)
			SP+SGDPK	TonB-dependent heme receptor A	P44523.1	744	Haemophilus influenza	129 (4 th hit)
			L SPIS DP	M-phase inducer phosphatase 1-B	P30309.1	550	Xenopus laevis	175 (5 th hit)
			L SPIS DP	M-phase inducer phosphatase 1-A	P30308.1	550	Xenopus laevis	175 (6 th hit)
			LSSP I+G+PK	Hemoglobin subunit β -3	P02136.2	147	Rana catesbeiana	175 (7 th hit)
			L SPIS DP	M-phase inducer phosphatase 3	P30311.1	572	Xenopus laevis	175 (8 th hit)
			L SPIS DP	M-phase inducer phosphatase 2	P30310.1	599	Xenopus laevis	175 (9 th hit)
B6	522.8	+2	IMSIVDPNR					
			IMSIVDPNR	α -actinin-1	Q3B7N2.1	892	Bos taurus	0.1 (1 st hit)
			IMSIVDPNR	α -actinin-1	Q2PFV7.1	892	Macaca fascicularis	0.1 (2 nd hit)
			IMSIVDPNR	α -actinin-1	Q7TPR4.1	892	Mus musculus	0.1 (3 rd hit)
			IMSIVDPNR	α -actinin-1	P05094.3	893	Gallus gallus	0.1 (4 th hit)
			IMSIVDPNR	α -actinin-1	P12814.2	892	Homo sapiens	0.1 (5 th hit)
			IMSIVDPNR	α -actinin-1	Q9Z1P2.1	892	Rattus norvegicus	0.1 (6 th hit)
			IMS+VDPN	α -actinin-4	A5D7D1.1	911	Bos taurus	6.2 (7 th hit)
			IMS+VDPN	α -actinin-4	Q9QXQ0.2	911	Rattus norvegicus	6.2 (8 th hit)
			IMS+VDPN	α -actinin-4	P57780.1	912	Mus musculus	6.2 (9 th hit)

Table A.1. cont'd.

			IMS+VDPN	α -actinin-4	Q5RCS6.1	911	Pongo abelii	15 (10 th hit)
			IMS+VDPN	α -actinin-2	P20111.1	897	Gallus gallus	15 (11 th hit)
			IMS+VDPN	α -actinin-4	O43707.2	911	Homo sapiens	15 (12 th hit)
			IMS+VDPN	α -actinin-4	Q90734.1	904	Gallus gallus	15 (13 th hit)
B5	506.2	+3	QEYDESGPSIVHR					
			QEYDESGPSIVHR	γ -Actin	P20359.2	375	Emericella nidulans	3e-05 (1 st hit)
			QEYDESGPSIVHR	Actin	P53455.2	375	Ajellomyces capsulatus G186AR	3e-05 (2 nd hit)
			QEYDESGPSIVHR	POTE ankyrin domain family member E	Q6S8J3.3	1075	Homo sapiens	3e-05 (3 rd hit)
			QEYDESGPSIVHR	Actin	Q0PGG4.1	375	Bos grunniens	3e-05 (4 th hit)
			QEYDESGPSIVHR	β -Actin	P84856.1	361	Chlorocebus pygerythrus	3e-05 (5 th hit)
			QEYDESGPSIVHR	Chimeric POTE-actin protein	A5A3E0.2	1075	Homo sapiens	3e-05 (6 th hit)
			QEYDESGPSIVHR	γ -Actin	Q5JAK2.1	375	Rana lessonae	3e-05 (7 th hit)
			QEYDESGPSIVHR	Actin, cytoplasmic 2	Q6P378.1	375	Xenopus tropicalis	3e-05 (8 th hit)
			QEYDESGPSIVHR	K-actin	Q9BYX7.1	375	Homo sapiens	3e-05 (9 th hit)
			QEYDESGPSIVHR	Actin	Q2U7A3.1	375	Aspergillus oryzae	3e-05 (10 th hit)
			QEYDESGPSIVHR	β -Actin	P84336.1	375	Camelus dromedarius	3e-05 (11 th hit)
			QEYDESGPSIVHR	β -Actin	Q6NVA9.1	375	Xenopus tropicalis	3e-05 (12 th hit)
			QEYDESGPSIVHR	β -Actin	Q7ZVI7.2	375	Danio rerio	3e-05 (13 th hit)
			QEYDESGPSIVHR	Actin, cytoskeletal 3	Q25379.1	172	Lytechinus pictus	3e-05 (14 th hit)
			QEYDESGPSIVHR	Actin, cytoplasmic	Q964E1.1	376	Biomphalaria obstructa	3e-05 (15 th hit)

Table A.1. cont'd.

QEYDESGPSIVHR	Actin, cytoplasmic	P12716.1	376	Pisaster ochraceus	3e-05 (16 th hit)
QEYDESGPSIVHR	Actin-3	P41340.1	376	Limulus polyphemus	3e-05 (17 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic A3	P04829.3	376	Bombyx mori	3e-05 (18 th hit)
QEYDESGPSIVHR	Actin-2	P26197.1	377	Absidia glauca	3e-05 (19 th hit)
QEYDESGPSIVHR	Actin-2	P92176.1	376	Lumbricus	3e-05 (20 th hit)
QEYDESGPSIVHR	Actin	P91754.1	372	Lumbricus rubellus	3e-05 (21 st hit)
QEYDESGPSIVHR	Actin, cytoskeletal 3A	P53474.1	376	Strongylocentrotus purpuratus	3e-05 (22 nd hit)
QEYDESGPSIVHR	Actin, cytoskeletal 3B	P18499.1	376	Strongylocentrotus purpuratus	3e-05 (23 rd hit)
QEYDESGPSIVHR	Actin-3	P41113.1	376	Podocoryna carnea	3e-05 (24 th hit)
QEYDESGPSIVHR	Actin, cytoskeletal 2A	Q07903.1	376	Strongylocentrotus purpuratus	3e-05 (25 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic A3a	Q25010.1	376	Helicoverpa armigera	3e-05 (26 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q964E3.1	376	Biomphalaria alexandrina	3e-05 (27 th hit)
QEYDESGPSIVHR	γ -Actin	P63256.1	375	Anser anser anser	3e-05 (28 th hit)
QEYDESGPSIVHR	Actin	P18602.1	327	Artemia species	3e-05 (29 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic	P92179.2	376	Biomphalaria glabrata	3e-05 (30 th hit)
QEYDESGPSIVHR	β -Actin	P68142.1	375	Takifugu rubripes	3e-05 (31 st hit)
QEYDESGPSIVHR	Actin	P53461.1	376	Halocynthia roretzi	3e-05 (32 nd hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q93131.1	375	Branchiostoma floridae	3e-05 (33 rd hit)
QEYDESGPSIVHR	Actin, cytoskeletal 1	P53465.1	376	Lytechinus pictus	3e-05 (34 th hit)
QEYDESGPSIVHR	Actin	P10365.1	375	Thermomyces lanuginosus	3e-05

Table A.1. cont'd.

					(35 th hit)
QEYDESGPSIVHR	Actin-1	O18499.1	376	Saccoglossus kowalevskii	3e-05 (36 th hit)
QEYDESGPSIVHR	β -Actin	Q91ZK5.1	375	Sigmodon hispidus	3e-05 (37 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic type 8	P53506.1	376	Xenopus laevis	3e-05 (38 th hit)
QEYDESGPSIVHR	Actin-1	P92182.1	376	Lumbricus terrestris	3e-05 (39 th hit)
QEYDESGPSIVHR	γ -Actin	Q8JJB8.1	375	Triakis scyllium	3e-05 (40 th hit)
QEYDESGPSIVHR	Actin-15A	P10990.1	376	Strongylocentrotus franciscanus	3e-05 (41 st hit)
QEYDESGPSIVHR	γ -Actin	Q9UVW9.1	375	Acremonium chrysogenum	3e-05 (42 nd hit)
QEYDESGPSIVHR	Actin, cytoskeletal 2	P53466.1	376	Lytechinus pictus	3e-05 (43 rd hit)
QEYDESGPSIVHR	Actin-1/2	P41112.1	376	Podocoryna carnea	3e-05 (44 th hit)
QEYDESGPSIVHR	Actin, cytoskeletal	P53464.1	376	Heliocidaris tuberculata	3e-05 (45 th hit)
QEYDESGPSIVHR	Actin, adductor muscle	Q26065.1	376	Placopecten magellanicus	3e-05 (46 th hit)
QEYDESGPSIVHR	Actin, cytoskeletal 1B	P53473.1	376	Strongylocentrotus purpuratus	3e-05 (47 th hit)
QEYDESGPSIVHR	β -Actin	P48975.1	375	Cricetulus griseus	3e-05 (48 th hit)
QEYDESGPSIVHR	Actin-15B	P69004.1	376	Strongylocentrotus franciscanu	3e-05 (49 th hit)
QEYDESGPSIVHR	Actin, muscle	P17304.1	376	Aplysia californica	3e-05 (50 th hit)
QEYDESGPSIVHR	β -Actin	P60711.1	375	Rattus norvegicus	3e-05 (51 st hit)
QEYDESGPSIVHR	Actin, muscle-type	Q25472.1	378	Molgula oculata	3e-05 (52 nd hit)
QEYDESGPSIVHR	Actin	Q8X119.1	375	Exophiala dermatitidis	3e-05 (53 rd hit)
QEYDESGPSIVHR	Actin-42A	P02572.3	376	Drosophila melanogaster	3e-05 (54 th hit)

Table A.1. cont'd.

QEYDESGPSIVHR	β -Actin	P79818.2	375	<i>Oryzias latipes</i>	3e-05 (55 th hit)
QEYDESGPSIVHR	β -Actin	P83750.1	375	<i>Cyprinus carpio</i>	3e-05 (56 th hit)
QEYDESGPSIVHR	Actin Cyl, cytoplasmic	P69002.1	376	<i>Heliocidaris erythrogramma</i>	3e-05 (57 th hit)
QEYDESGPSIVHR	Actin-1	P30163.1	376	<i>Onchocerca volvulus</i>	3e-05 (58 th hit)
QEYDESGPSIVHR	Actin-2	P30162.1	376	<i>Onchocerca volvulus</i>	3e-05 (59 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q93129.1	375	<i>Branchiostoma belcheri</i>	3e-05 (60 th hit)
QEYDESGPSIVHR	Actin, non-muscle	P17126.1	376	<i>Hydra vulgaris</i>	3e-05 (61 st hit)
QEYDESGPSIVHR	β -Actin	O93400.1	375	<i>Xenopus laevis</i>	3e-05 (62 nd hit)
QEYDESGPSIVHR	Actin-2	O18500.1	376	<i>Saccoglossus kowalevskii</i>	3e-05 (63 rd hit)
QEYDESGPSIVHR	Actin, clone 403	P18603.1	376	<i>Artemia</i> species	3e-05 (64 th hit)
QEYDESGPSIVHR	β -Actin	P15475.1	376	<i>Xenopus borealis</i>	3e-05 (65 th hit)
QEYDESGPSIVHR	Actin, cytoskeletal 1A	P53472.1	376	<i>Strongylocentrotus purpuratus</i>	3e-05 (66 th hit)
QEYDESGPSIVHR	β -Actin	O42161.1	375	<i>Salmo salar</i>	3e-05 (67 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q964E0.1	376	<i>Biomphalaria tenagophila</i>	3e-05 (68 th hit)
QEYDESGPSIVHR	Actin	O13419.1	375	<i>Botryotinia fuckeliana</i>	3e-05 (69 th hit)
QEYDESGPSIVHR	β -Actin	P53486.1	375	<i>Takifugu rubripes</i>	3e-05 (70 th hit)
QEYDESGPSIVHR	Actin-5	P41339.1	376	<i>Limulus polyphemus</i>	3e-05 (71 st hit)
QEYDESGPSIVHR	γ -Actin	Q9URS0.1	375	<i>Penicillium chrysogenum</i>	3e-05 (72 nd hit)
QEYDESGPSIVHR	Actin, cytoskeletal	P53463.1	376	<i>Heliocidaris erythrogramma</i>	3e-05 (73 rd hit)
QEYDESGPSIVHR	Actin	P50138.1	375	<i>Puccinia graminis</i>	3e-05 (74 th hit)

Table A.1. cont'd.

QEYDESGPSIVHR	Actin-1/3	P10983.2	376	<i>Caenorhabditis elegans</i>	3e-05 (75 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic type 5	P53478.1	376	<i>Gallus gallus</i>	3e-05 (76 th hit)
QEYDESGPSIVHR	β -Actin B	P53485.1	375	<i>Takifugu rubripes</i>	3e-05 (77 th hit)
QEYDESGPSIVHR	Actin-3	Q03342.1	309	<i>Echinococcus granulosus</i>	3e-05 (78 th hit)
QEYDESGPSIVHR	Actin, muscle	P12717.1	376	<i>Pisaster ochraceus</i>	3e-05 (79 th hit)
QEYDESGPSIVHR	β -Actin	P29751.1	375	<i>Oryctolagus cuniculus</i>	3e-05 (80 th hit)
QEYDESGPSIVHR	Actin-2	P10984.3	376	<i>Caenorhabditis elegans</i>	3e-05 (81 st hit)
QEYDESGPSIVHR	Actin, muscle	Q25381.1	172	<i>Lytechinus pictus</i>	3e-05 (82 nd hit)
QEYDESGPSIVHR	Actin-6	P53459.1	373	<i>Diphyllobothrium dendriticum</i>	3e-05 (83 rd hit)
QEYDESGPSIVHR	Actin-11	P41341.1	376	<i>Limulus polyphemus</i>	3e-05 (84 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q964E2.1	376	<i>Biomphalaria pfeifferi</i>	3e-05 (85 th hit)
QEYDESGPSIVHR	Actin	Q24733.1	33	<i>Dictyocaulus viviparus</i>	3e-05 (86 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic type 5	P53505.1	376	<i>Xenopus laevis</i>	3e-05 (87 th hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q00215.1	375	<i>Styela plicata</i>	3e-05 (88 th hit)
QEYDESGPSIVHR	Actin	O17320.1	376	<i>Crassostrea gigas</i>	3e-05 (89 th hit)
QEYDESGPSIVHR	Actin-4	P10986.2	376	<i>Caenorhabditis elegans</i>	3e-05 (90 th hit)
QEYDESGPSIVHR	Actin-1	P35432.1	375	<i>Echinococcus granulosus</i>	3e-05 (91 st hit)
QEYDESGPSIVHR	Actin, cytoplasmic	O17503.1	375	<i>Branchiostoma lanceolatum</i>	3e-05 (92 nd hit)
QEYDESGPSIVHR	Actin, cytoplasmic	Q964D9.1	376	<i>Helisoma trivolvis</i>	3e-05 (93 rd hit)

Table A.1. cont'd.

			QEYDESGPSIVHR	Actin, muscle	Q00214.1	379	Styela plicata	3e-05 (94 th hit)
			QEYDESGPSIVHR	Actin-5C	P10987.4	376	Drosophila	3e-05 (95 th hit)
			QEYDESGP+IVHR	Actin-1	P04751.1	377	Xenopus laevis	2e-04 (96 th hit)
			QEYDE+GPSIVHR	α -Actin-1	P53479.1	377	Cyprinus carpio	2e-04 (97 th hit)
			QEYDE+GPSIVHR	α -cardiac actin	P04751.1	377	Xenopus laevis	2e-04 (98 th hit)
			QEYDE+GPSIVHR	Actin, α cardiac	P53480.	377	Takifugu rubripes	2e-04 (99 th hit)
B1	683.9	+2	STDYGILQINSR					
			STDYGILQINSR	Lysozyme C	P84492.1	130	Chelonia mydas	6e-04 (1 st hit)
			STDYGILQINSR	Lysozyme C	Q7LZ13.1	129	Tragopan satyra	6e-04 (2 nd hit)
			STDYGILQINSR	Lysozyme C	Q7LZQ1.3	131	Pelodiscus sinensis	6e-04 (3 rd hit)
			STDYGILQINSR	Lysozyme C	P24364.1	129	Lophura leucomelanos	6e-04 (4 th hit)
			STDYGILQINSR	Lysozyme C	P81711.1	129	Syrmaticus soemmerringii	6e-04 (5 th hit)
			STDYGILQINSR	Lysozyme C	P00702.2	147	Phasianus colchicus colchicus	6e-04 (6 th hit)
			STDYGILQINSR	Lysozyme C	P00703.2	147	Meleagris gallopavo	6e-04 (7 th hit)
			STDYGILQINSR	Lysozyme C	Q7LZP9.1	129	Lophophorus impejanus	6e-04 (8 th hit)
			STDYGILQINSR	Lysozyme C	P49663.1	130	Phasianus versicolor	6e-04 (9 th hit)
			STDYGILQINSR	Lysozyme C	P00698.1	147	Gallus gallus	6e-04 (10 th hit)

Table A.1. cont'd.

			STDYGILQINSR	Lysozyme C	P00705.1	147	Anas platyrhynchos	6e-04 (11 th hit)
			STDYGILQINSR	Lysozyme C	P19849.1	129	Pavo cristatus	6e-04 (12 th hit)
			STDYGILQINSR	Lysozyme C	P00701.2	147	Coturnix japonica	6e-04 (13 th hit)
			STDYGILQINSR	Lysozyme C	Q7LZQ3.1	129	Crax fasciolata	6e-04 (14 th hit)
			STDYGILQINSR	Lysozyme C	P37156.1	125	Tachyglossus aculeatus aculeatus	6e-04 (15 th hit)
			STDYGILQINSR	Lysozyme C	P24533.1	129	Syrnaticus reevesii	6e-04 (16 th hit)
			STDYGILQINSR	Lysozyme C	Q7LZT2.1	129	Tragopan temminckii	6e-04 (17 th hit)
			STDYGILQINSR	Lysozyme C	P00707.1	129	Ortalis vetula	6e-04 (18 th hit)
			STDYGILQINSR	Lysozyme C	P22910.1	129	Chrysolophus amherstiae	6e-04 (19 th hit)
			STDYG+LQINSR	Lysozyme C	P00704.2	129	Numida meleagris	0.003 (20 th hit)
			STDYG+LQINSR	Lysozyme C	P00699.1	129	Callipepla californica	0.003 (21 st hit)
			STDYG+LQINSR	Lysozyme C	P00700.2	129	Colinus virginianus	0.003 (22 nd hit)
			STDYGIL+INSR	Lysozyme C	P00706.1	129	Anas platyrhynchos	0.004 (23 rd hit)
			STDYGIL+INSR	Lysozyme C	Q7LZQ2.1	129	Aix sponsa	0.004 (24 th hit)
B1	503.3	+2	WDAWDALK					
			WDAW+ALK	Acyl-CoA-binding protein	Q9PRL8.1	86	Gallus gallus	1.7 (1 st hit)
			WDAW+AL	Peroxisomal 3,2-trans-enoyl-CoA isomerase	Q9WUR2.2	391	Mus musculus	13 (2 nd hit)
			WDAW+AL	Peroxisomal 3,2-trans-enoyl-CoA isomerase	Q5XIC0.1	391	Rattus norvegicus	13 (3 rd hit)

Table A.1. cont'd.

			WDAW+AL	Peroxisomal 3,2-trans-enoyl-CoA isomerase	O75521.4	394	Homo sapiens	13 (4 th hit)
			WDAW+ LK	Acyl-CoA-binding protein	P12026.2	87	Sus scrofa	18 (5 th hit)
			WDAW+ LK	Acyl-CoA-binding protein	Q8WN94.3	87	Oryctolagus cuniculus	18 (6 th hit)
			WDAW+ LK	Acyl-CoA-binding protein	P07108.2	87	Homo sapiens	18 (7 th hit)
			WDAW+ LK	Acyl-CoA-binding protein	P07107.2	87	Bos taurus	18 (8 th hit)
B4	509.8	+2	IGTMLPMQK					
			I TMLPM QK	Peptidoglycan hydrolase flgJ	Q914P4.1	400	Pseudomonas aeruginosa	15 (1 st hit)
			MLPMQK	Myeloid protein 1	P08940.2	326	Gallus gallus	36 (2 nd hit)
			IG MLPM K	Arginine biosynthesis bifunctional protein argJ	Q71Z77.1	398	Listeria monocytogenes str. 4b F2365	49 (3 rd hit)
			IG MLPM K	Arginine biosynthesis bifunctional protein argJ	Q8Y6U2.1	398	Listeria monocytogenes	49 (4 th hit)
			G+MLPM+K	Replicase large subunit	P18339.2	1609	Tobacco mild green mosaic virus	49 (5 th hit)
B1	565.3	+2	LVTDVQEAVR					
			LVTDVQEAVR	Proactivator polypeptide (Containing Saposin-A, B, C & D)	O13035.1	518	Gallus gallus	0.084 (1 st hit)
			LVTDVQEA+R	Hypothetical protein	YP_001608779.1	525	Bartonella tribocorum CIP 105476	1.2 (2 nd hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q21XJ9.1	278	Rhodoferrax ferrireducens	17 (3 rd hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q2SVN0.1	335	Burkholderia thailandensis E264	17 (4 th hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q39GW5.1	319	Burkholderia species 383	17 (5 th hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q0BFQ0.1	319	Burkholderia ambifaria AMMD	17 (6 th hit)
			LVT DVQEAV	Aliphatic sulfonates	Q3JSR6.1	335	Burkholderia pseudomallei 1710b	17

Table A.1. cont'd.

				import ATP-binding protein ssuB					(7 th hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q62K56.2	335		Burkholderia mallei	17 (8 th hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q1BWL4.1	319		Burkholderia cenocepacia AU 1054	17 (9 th hit)
			LVT DVQEAV	Aliphatic sulfonates import ATP-binding protein ssuB	Q63TW1.1	327		Burkholderia pseudomallei	17 (10 th hit)
B2	646.9	+2	ISMPDFDLNLK						
			ISMPDFDLNLK	Neuroblast differentiation-associated protein AHNAK	Q09666.2	5890		Homo sapiens	0.001 (1 st hit)
			ISMPDFD++	DNA polymerase III subunit alpha	Q1RKF9.1	1172		Rickettsia bellii RML369-C	5.1 (2 nd hit)
			ISMPDFD++	DNA polymerase III subunit alpha	Q68VX1.1	1180		Rickettsia typhi	5.1 (3 rd hit)
			ISMPDFD++	DNA polymerase III subunit alpha	Q92GB2.1	1181		Rickettsia conorii	5.1 (4 th hit)
			ISMPDFD++	DNA polymerase III subunit alpha	Q4UK40.1	1207		Rickettsia felis	5.1 (5 th hit)
			ISMPDFD++	DNA polymerase III subunit alpha	O05974.2	1182		Rickettsia prowazekii	5.1 (6 th hit)
			ISMPDFD++	DNA polymerase III subunit alpha	O51526.2	1147		Borrelia burgdorferi	5.1 (7 th hit)
			ISMPDFD++	DNA polymerase III subunit alpha	O83675.1	1170		Treponema pallidum	5.1 (8 th hit)
			ISMPDFD++	DNA polymerase III subunit alpha	Q9RX08.1	1335		Deinococcus radiodurans	5.1 (9 th hit)
			ISMPDFD++	DNA polymerase III subunit alpha	P57332.1	1161		Buchnera aphidicola	5.1 (10 th hit)
			ISMPDFD	DNA polymerase III subunit alpha	Q9HXZ1.1	1173		Pseudomonas aeruginosa	6.8 (11 th hit)
B2	772.0	+2	AVASAAAALVLK						
			AVASAAAALVLK	Talin-1	P26039.1	2541		Mus musculus	0.026 (1 st hit)
			AVASAAAALVLK	Talin-1	P54939.2	2541		Gallus gallus	0.026 (2 nd hit)

Table A.1. cont'd.

			AVASAAAALVLK	Talin-1	Q9Y490.3	2541	Homo sapiens	0.026 (3 rd hit)
			AVA+AAA LVLK	Talin-2	Q9Y4G6.4	2542	Homo sapiens	3.8 (4 th hit)
			AVA+AAA LVLK	Talin-2	Q71LX4.2	2375	Mus musculus	3.8 (5 th hit)
			AVA AAALVLK	Phycocyanin-645 α -1 chain	P23815.1	70	Chroomonas species	6.8 (6 th hit)
B2	525.8	+2	LGTFLENEK					
			LGTFLE+E	Glutamyl-tRNA reductase	Q2FTL0.1	424	Methanospirillum hungatei JF-1	36 (1 st hit)
			LGTFLEN	Histone acetyltransferase p300	Q09472.2	2414	Homo sapiens	36 (2 nd hit)
			L TFLEN K	Probable nucleoporin C890.06	Q9URX8.3	1315	Schizosaccharomyces pombe	87 (3 rd hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	B0BBM4.1	354	Chlamydia trachomatis L2b/UCh-1/proctitis	117 (4 th hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	B0B7F9.1	354	Chlamydia trachomatis	117 (5 th hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	Q25419.1	359	Chlamydomphila felis	117 (6 th hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	Q5L612.1	359	Chlamydomphila abortus	117 (7 th hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	Q823E0.1	359	Chlamydomphila caviae	117 (8 th hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	Q9PKF1.1	354	Chlamydia muridarum	117 (9 th hit)
			TFL+NEK	UDP-3-O-[3-hydroxymyristoyl] glucosamine N-acyltransferase	O84245.1	354	Chlamydia trachomatis	117 (10 th hit)

Table A.1. cont'd.

B4	821.9	+2	V(I/L)QQQADDAEER					
			V+QQQADDAEER	Putative tropomyosin α -3 chain-like protein	A6NL28.2	223	Homo sapiens	0.006 (1 st hit)
			V+QQQADDAEER	Tropomyosin α -3 chain	Q63610.2	248	Rattus norvegicus	0.006 (2 nd hit)
			QQQADDAE+R	Tropomyosin α -4 chain	P09495.3	248	Rattus norvegicus	0.20 (3 rd hit)
			QQQADDAE+R	Tropomyosin α -4 chain	Q6JRU2.3	248	Mus musculus	0.20 (4 th hit)
			QQQAD+AE+R	Tropomyosin α -4 chain	P67937.3	248	Sus scrofa	1.2 (5 th hit)
			QQQAD+AE+R	Tropomyosin α -4 chain	P02561.2	248	Equus caballus	1.2 (6 th hit)
B6	587.3	+2	IPPKPPARAAR					
			PP+PPARAA	Protein bassoon	Q9UPA5.4	3926	Homo sapiens	17 (1 st hit)
			PPKPPAR	Frizzled-8	O93274.1	581	Xenopus laevis	30 (2 nd hit)
			IPPKPP R	Sodium/hydrogen exchanger 2	Q9UBY0.1	812	Homo sapiens	30 (3 rd hit)
			PPKPPAR	Structure-specific endonuclease subunit slx4	Q7SEJ3.1	1013	Neurospora crassa	30 (4 th hit)
			PP+PPARA	Protein bassoon	O88737.3	3942	Mus musculus	97 (5 th hit)
			PPKPPA A	DNA polymerase subunit γ -1	P54098.1	1239	Homo sapiens	97 (6 th hit)
			PP+PPARA	Protein bassoon	O88778.3	3938	Rattus norvegicus	97 (7 th hit)
P PPAR+AR	Stress response protein NST1	Q5KHY3.1	1353	Filobasidiella neoformans	97 (8 th hit)			
B5	506.8	+2	IPSLPSGVDK					
			PSLPSGVD	Stromelysin-1	P08254.2	477	Homo sapiens	12 (1 st hit)
			PSLPSG+D	Stromelysin-1	Q28397.1	477	Equus caballus	54 (2 nd hit)
			IP LPSGV+	Amino-acid acetyltransferase	A1C6J2.1	718	Aspergillus clavatus	72 (3 rd hit)
B6	583.8	+2	AAGEIIAIPRR					
			AAGEIIAIPRR	Envelope glycoprotein gp160	Q89607.1	852	HIV-2 B_EHO	0.011 (1 st hit)
			AA IIAIPRR	Envelope glycoprotein gp160	P15831.2	859	Human immunodeficiency virus type 2	17 (2 nd hit)

Table A.1. cont'd.

			EIIAIP+R	Porphobilinogen deaminase	A7GBW3.1	290	Clostridium botulinum F str. Langeland	22 (3 rd hit)
			IIAIPRR	Envelope glycoprotein gp160	Q76638.1	857	HIV-2_B_UC1	22 (4 th hit)
B1	831.4	+2	NSWGTSWGEDGYFR					
			NSWGTSWGEDGYFR	Dipeptidyl-peptidase 1 (Cathepsin C)	Q60HG6.1	463	Macaca fascicularis	5e-07 (1 st hit)
			NSWGTSWGE+GYFR	Dipeptidyl-peptidase 1 (Cathepsin C)	Q3ZCJ8.1	463	Bos taurus	3e-06 (2 nd hit)
			NSWGT WGEDGYFR	Dipeptidyl-peptidase 1 (Cathepsin C)	Q5RB02.1	463	Pongo abelii	5e-06 (3 rd hit)
B5	684.9	+2	FRTTMLQDSIR					
			FRTTMLQDSI	Origin recognition complex subunit 2	Q75PQ8.1	576	Rattus norvegicus	0.014 (1 st hit)
			FRTTMLQDSI	Origin recognition complex subunit 2	Q60862.1	586	Mus musculus	0.014 (2 nd hit)
			FRTTMLQDSI	Origin recognition complex subunit 2	Q13416.2	577	Homo sapiens	0.014 (3 rd hit)
			FRTTMLQD I	Origin recognition complex subunit 2	A6QNM3.2	577	Bos taurus	0.20 (4 th hit)
			FRT++LQDS	Origin recognition complex subunit 2	Q91628.1	558	Xenopus laevis	22 (5 th hit)
B6	586.8	+2	LPPEQGTSSR					
			LPPEQGTSSR	Cadherin EGF LAG seven-pass G-type receptor 2	XP_002686199.1	2920	Bos Taurus (flamingo homolog, Drosophila)	0.36 (1 st hit)
			LPPEQGTSSR	Cadherin EGF LAG seven-pass G-type receptor 2	NP_001179860.1	2920	Bos Taurus	0.36 (2 nd hit)
			LPPE+GTTS	26S proteasome regulatory subunit 7, putative	XP_002778882.1	543	Perkinsus marinus ATCC 50983	173 (3 rd hit)
			LPPE+GTT S	Hypothetical protein LELG_03374	XP_001525446.1	716	Lodderomyces elongisporus NRRL YB-4239	173 (4 th hit)
			PPE+GTTS R	Thiamine ABC transporter, permease protein	YP_002827746.1	540	Rhizobium sp. NGR234	232 (5 th hit)
B6	593.3	+2	AILYNYWDK					
			AILYNYWDK	Complement component c3	CAC69535.1	401	Crocodylus niloticus	0.50 (1 st hit)
			ILYNYWD	Collagen-like cell surface-anchored protein ScID	YP_002745264.1	324	Streptococcus equi subspecies Zooepidemicus	24 (2 nd hit)

Table A.1. cont'd.

			ILYNYWD	Collagen-like cell surface- anchored protein ScLD	YP_002745683.1	313	Streptococcus equi subspecies equi 4047	24 (3 rd hit)
			ILYNYWD	Collagen-like cell surface- anchored protein ScLD	YP_002122604.1	336	Streptococcus equi subspecies zoepidemicus MGCS10565	24 (4 th hit)
			ILYNYWD	Collagen-like protein D	ABA41494.1	258	Streptococcus equi subspecies Equi	24 (5 th hit)
			A+LY YWDK	Hypothetical protein Btr_0321	YP_001608779.1	711	Bartonella tribocorum CIP 105476	41 (6 th hit)
			AILYNYW	Complement component c3	NP_990736.1	1652	Gallus gallus	41 (7 th hit)
B6	506.8	+2	NEALIALLR					
			NEALIALLR	Plastin-2	Q6P698.1	624	Danio rerio	0.79 (1 st hit)
			NEALIALLR	Plastin-2	P13796.5	627	Homo sapiens	0.79 (2 nd hit)
			NEALIALLR	Plastin-2	Q61233.4	627	Mus musculus	0.79 (3 rd hit)
			NEALIALL	Plastin-1	A6H742.1	630	Bos taurus	8.3 (4 th hit)
			NEALIALL	Plastin-1	Q14651.2	629	Homo sapiens	8.3 (5 th hit)
			NEALIALL	Plastin-1	Q3V0K9.1	630	Mus musculus	8.3 (6 th hit)
			NEALIALL	Plastin-1	P19179.1	630	Gallus gallus	8.3 (7 th hit)
			NEALIAL LR	Metal-dependent hydrolase	Q6KZZ9.1	220	Picrophilus torridus	20 (8 th hit)

Table A.2. Proteins identified at single peptide level using BLAST. *De novo* sequenced peptides obtained from gel digest were searched using BLAST with a limited taxonomy, containing birds (taxid:8782), crocodiles (taxid:8493), turtles (taxid:8459), tuataras (taxid:8508) and squamates (taxid:8509) using the SwissProt database. The top three E-values are reported.

Band Number	<i>m/z</i>	Charge	<i>De Novo</i> Sequence	Protein	Accession Number	Protein Length	Organism	E-Value	
B4	1060.6	+2	EINLSPDSTSAVVVSGLMVATK						
			EINLSPDSTSAVVVSGLMVATK	Fibronectin	P11722	1256	Gallus gallus	2e-13 (1 st hit)	
			NLSP+S SA	Endoribonuclease Dicer	Q25BN1	1921	Gallus gallus	2.2 (2 nd hit)	
B4	536.3	+2	YEVSVYALK						
			YEVSVYALK	Fibronectin	P11722	1256	Gallus gallus	0.003 (1 st hit)	
			Y+++VYA+K	Collagen α -1(XII)	P13944	3124	Gallus gallus	3.7 (2 nd hit)	
B4	701.3	+2	YVSVYA	Tyrosine-protein kinase BTK	Q8JH64	657	Gallus gallus	12 (3 rd hit)	
			YQINQQWER						
			YQINQ ER	Solute carrier family 41 member 2	Q5ZHX6	573	Gallus gallus	2.1 (1 st hit)	
B4	701.3	+2	INQQW	Thrombin-like enzyme	Q072L7	258	Lachesis stenophrys	2.8 (2 nd hit)	
			INQQW	Serine α -fibrinogenase	Q8JH85	258	Macrovipera lebetina	2.8 (3 rd hit)	
			INQQW	Serine β -fibrinogenase	Q8JH62	257	Macrovipera lebetina	2.8 (4 th hit)	
			QQWER	Zyxin	Q04584	542	Gallus gallus	2.8 (5 th hit)	
			YQ + QQWE	Structural maintenance of chromosomes protein 2	Q90988	1189	Gallus gallus	5.0 (6 th hit)	
			B1	613.9	+2	LIALLEVLVSQK			
IA LLEVLVS	Adenylate cyclase type 9	Q9DGG6				1334	Gallus gallus	2.3 (1 st hit)	
LLEVL+Q	Exportin-7	Q5ZLT0				1087	Gallus gallus	4.1 (2 nd hit)	
LI +LE +LSQK	E3 ubiquitin-protein ligase MIB2	Q5ZLJ9				954	Gallus gallus	5.5 (3 rd hit)	

Table A.2. cont'd.

			AL EV+SQK	Cell division cycle-associated protein 1	Q76190	469	Gallus gallus	5.5 (4 th hit)
B1	487.3	+2	VYGPVGEPR					
			VYGP VE	116 kDa U5 small nuclear ribonucleoprotein component	Q5F3X4	972	Gallus gallus	6.7 (1 st hit)
			VYGP	Homeobox protein Hox-C8	Q9YH13	242	Gallus gallus	29 (2 nd hit)
			YGP VE	Protein sidekick-2	Q8AV57	2177	Gallus gallus	39 (3 rd hit)
B2	736.4	+2	AQVVSQGLDLLTAK					
			AQVVSQGLDLLTAK	Vinculin	P12003	1135	Gallus gallus	5e-07 (1 st hit)
			L+LLTAK	NADH-ubiquinone oxidoreductase chain 3	Q47874	115	Alligator mississippiensis	5.4 (2 nd hit)
			LDLLT K	Probable ATP-dependent RNA helicase DDX10	Q5ZJF6	875	Gallus gallus	9.8 (3 rd hit)
B2	739.4	+2	MLGQMTDQVADLR					
			LGQMTDQ+ADLR	Vinculin	P12003	1135	Gallus gallus	6e-05 (1 st hit)
			Q+TDQVA LR	COUP transcription factor 2	Q90733	410	Gallus gallus	2.3 (2 nd hit)
			MLGQM	Importin-13	Q5ZIC8	958	Gallus gallus	5.5 (3 rd hit)
B2	655.9	+2	TVTAMDVVYALK					
			TVTAMDVVYALK	Histone H4	P70081	103	Gallus gallus	5e-06 (1 st hit)
			TVTAMDVVYALK	Histone H4	P62800	103	Cairina moschata	5e-06 (2 nd hit)
			TVT +D VYALK	Rho guanine nucleotide exchange factor 6	Q5ZLR6	764	Gallus gallus	1.7 (3 rd hit)
			T+T +DVVY	UPF0636 protein C4orf41 homolog	Q5ZI89	1132	Gallus gallus	5.5 (4 th hit)
B2	628.7	+2	LITKAVSASK					
			LITKAVSASK	Histone H1.03	P08285	224	Gallus gallus	0.005 (1 st hit)
			LITKAVSASK	Histone H1.10	P08286	220	Gallus gallus	0.005 (2 nd hit)
			LITKAVSASK	Histone H1.11L	P08287	225	Gallus gallus	0.005 (3 rd hit)

Table A.2. cont'd.

			LITKAVSASK	Histone H1.01	P08284	219	Gallus gallus	0.005 (4 th hit)
			LITKAVSASK	Histone H1	P09987	218	Gallus gallus	0.005 (5 th hit)
			LITKAVSASK	Histone H1.11R	P08288	219	Gallus gallus	0.005 (6 th hit)
			LITKAV+ASK	Histone H1	P09426	218	Anas platyrhynchos	0.028 (7 th hit)
			L+TKAV A	Neurochondrin	Q5ZIG0	702	Gallus gallus	18 (8 th hit)
B6	833.9	+2	FSGSGSGTDFTFITISR					
			FSG GSG +FT	Large fibroblast proteoglycan	Q90953	3562	Gallus gallus	1.7 (1 st hit)
			S SG GT FTF+ R	Docking protein 3	A3R064	426	Gallus gallus	5.4 (2 nd hit)
			SG+GSG+ TF	Protein PRRC1	Q5F310	442	Gallus gallus	13 (3 rd hit)
B6	751.4	+2	VFGGGTKLTVLGQPK					
			TVLGQP	Transmembrane protein 184C	Q5ZMP3	445	Gallus gallus	5.4 (1 st hit)
			GT LTVL Q K	Integrator complex subunit 2	Q5ZKU4	1192	Gallus gallus	13 (2 nd hit)
			VFG +KLT	Protein HIRA	P79987	1019	Gallus gallus	18 (3 rd hit)
B2	510.8	+2	VLASFGEAVK					
			VLASFGEAVK	Hemoglobin subunit β	P02131	146	Caiman crocodilus	0.003 (1 st hit)
			VLASFGEAVK	Hemoglobin subunit β	P02130	146	Alligator mississippiensis	0.003 (2 nd hit)
			VL+SFGGEAVK	Hemoglobin subunit epsilon	P02128	147	Gallus gallus	0.015 (3 rd hit)
			VL+SFGGEAVK	Hemoglobin ρ chain	P02127	147	Gallus gallus	0.015 (4 th hit)
			VLASFGEAV	Hemoglobin β chain	P02129	146	Crocodylus niloticus	0.021 (5 th hit)
			VL SFGGEAVK	Hemoglobin β chain	P13274	146	Chrysemys picta bellii	0.021 (6 th hit)
			VL SFGGEAVK	Hemoglobin β chain	P83133	146	Dipsosaurus dorsalis	0.021 (7 th hit)
			VL SFGGEAVK	Hemoglobin β chain	P02125	146	Ciconia ciconia	0.021 (8 th hit)

Table A.2. cont'd.

			VL SFG _E AVK	Hemoglobin β chain	<u>P14524</u>	146	Turdus merula	0.021 (9 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P08851</u>	146	Accipiter gentilis	0.021 (10 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P02116</u>	146	Ara ararauna	0.021 (11 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P07411</u>	146	Vultur gryphus	0.021 (12 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P15165</u>	146	Apus apus	0.021 (13 th hit)
			VL SFG _E AVK	Hemoglobin β/β' chain	<u>P08261</u>	146	Larus ridibundus	0.021 (14 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P21668</u>	146	Psittacula krameri	0.021 (15 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P02120</u>	146	Anseranas semipalmata	0.021 (16 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P10782</u>	146	Phalacrocorax carbo	0.021 (17 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P07406</u>	146	Passer montanus	0.021 (18 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P68061</u>	146	Aegyptius monachus	0.021 (19 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P02122</u>	146	Aquila chrysaetos	0.021 (20 th hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P02121</u>	146	Phoenicopters ruber ruber	0.021 (21 st hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>Q98905</u>	147	Geochelone carbonaria	0.021 (22 nd hit)
			VL SFG _E AVK	Hemoglobin β chain	<u>P82113</u>	146	Stercorarius maccormicki	0.021 (23 rd hit)
B2	500.8	+2	LSSPISGDPK					
			LSSP ISG+P	Hemoglobin subunit β	<u>P07036</u>	146	Chloephaga melanoptera	2.3 (1 st hit)
			PISGDP	Myosin-binding protein C	<u>Q90688</u>	1272	Gallus gallus	4.1 (2 nd hit)
			LSSP I G+PK	Hemoglobin subunit ϵ	<u>P14261</u>	147	Cairina moschata	13 (3 rd hit)
			LSSP I G+PK	Hemoglobin ρ chain	<u>P02127</u>	147	Gallus gallus	13

Table A.2. cont'd.

							(4 th hit)	
			LSSP I+G+P	Hemoglobin β/β' chain	<u>P08261</u>	146	Larus ridibundus	13 (5 th hit)
			LSSP I+G+P	Hemoglobin β -1 chain	<u>P18993</u>	146	Varanus exanthematicus albigularis	13 (6 th hit)
B6	522.8	+2	IMSIVDPNR					
			IMSIVDPNR	α -actinin-1	<u>P05094</u>	893	Gallus gallus	0.001 (1 st hit)
			IMS+VDPN	α -actinin-2	<u>P20111</u>	897	Gallus gallus	0.15 (2 nd hit)
			IMS+VDPN	α -actinin-4	<u>Q90734</u>	904	Gallus gallus	0.15 (3 rd hit)
			SIVDP NR	PRELI domain-containing protein 1, mitochondrial	<u>Q90673</u>	215	Gallus gallus	5.0 (4 th hit)
B5	506.2	+3	QEYDESGPSIVHR					
			QEYDESGPSIVHR	γ -actin	<u>Q5ZMQ2</u>	375	Gallus gallus	3e-07 (1 st hit)
			QEYDESGPSIVHR	γ -actin	<u>P63256</u>	375	Anser anser	3e-07 (2 nd hit)
			QEYDESGPSIVHR	β -actin	<u>P60706</u>	375	Gallus gallus	3e-07 (3 rd hit)
			QEYDESGPSIVHR	Actin, cytoplasmic type 5	<u>P53478</u>	376	Gallus gallus	3e-07 (4 th hit)
			QEYDE+GPSIVHR	Actin, alpha cardiac muscle 1	<u>P68034</u>	377	Gallus gallus	2e-06 (5 th hit)
			QEYDE+GPSIVHR	α -actin	<u>P08023</u>	377	Gallus gallus	2e-06 (6 th hit)
			QEYDE+GPSIVHR	α -actin-1	<u>P68139</u>	377	Gallus gallus	2e-06 (7 th hit)
			QEYDE+GPSIVHR	α -actin-1	<u>Q90X97</u>	377	Atractaspis microlepidota	2e-06 (8 th hit)
			EYDE+GPSIVHR	Γ -2-actin	<u>P63270</u>	376	Gallus gallus	2e-05 (9 th hit)
B1	683.9	+2	STDYGILQINSR					
			STDYGILQINSR	Lysozyme C	<u>P84492</u>	130	Chelonia mydas	5e-06 (1 st hit)
			STDYGILQINSR	Lysozyme C	<u>Q7LZI3</u>	129	Tragopan satyra	5e-06 (2 nd hit)

Table A.2. cont'd.

STDYGILQINSR	Lysozyme C	<u>Q7LZO1</u>	131	Pelodiscus sinensis	5e-06 (3 rd hit)
STDYGILQINSR	Lysozyme C	<u>P24364</u>	129	Lophura leucomelanos	5e-06 (4 th hit)
STDYGILQINSR	Lysozyme C	<u>P81711</u>	129	Syrmaticus soemmerringii	5e-06 (5 th hit)
STDYGILQINSR	Lysozyme C	<u>P00702</u>	147	Phasianus colchicus colchicus	5e-06 (6 th hit)
STDYGILQINSR	Lysozyme C	<u>P00703</u>	147	Meleagris gallopavo	5e-06 (7 th hit)
STDYGILQINSR	Lysozyme C	<u>Q7LZP9</u>	129	Lophophorus impejanus	5e-06 (8 th hit)
STDYGILQINSR	Lysozyme C	<u>P49663</u>	130	Phasianus versicolor	5e-06 (9 th hit)
STDYGILQINSR	Lysozyme C	<u>P00698</u>	147	Gallus gallus	5e-06 (10 th hit)
STDYGILQINSR	Lysozyme C-1	<u>P00705</u>	147	Anas platyrhynchos	5e-06 (11 th hit)
STDYGILQINSR	Lysozyme C	<u>P19849</u>	129	Pavo cristatus	5e-06 (12 th hit)
STDYGILQINSR	Lysozyme C	<u>P00701</u>	147	Coturnix japonica	5e-06 (13 th hit)
STDYGILQINSR	Lysozyme C	<u>Q7LZO3</u>	129	Crax fasciolata	5e-06 (14 th hit)
STDYGILQINSR	Lysozyme C	<u>P24533</u>	129	Syrmaticus reevesii	5e-06 (15 th hit)
STDYGILQINSR	Lysozyme C	<u>Q7LZT2</u>	129	Tragopan temminckii	5e-06 (16 th hit)
STDYGILQINSR	Lysozyme C	<u>P00707</u>	129	Ortalis vetula	5e-06 (17 th hit)
STDYGILQINSR	Lysozyme C	<u>P22910</u>	129	Chrysolophus amherstiae	5e-06 (18 th hit)
STDYG+LQINSR	Lysozyme C	<u>P00704</u>	129	Numida meleagris	3e-05 (19 th hit)

Table A.2. cont'd.

			STDYG+LQINSR	Lysozyme C	<u>P00699</u>	129	Callipepla californica	3e-05 (20 th hit)
			STDYG+LQINSR	Lysozyme C	<u>P00700</u>	129	Colinus virginianus	3e-05 (21 st hit)
			STDYGIL+INSR	Lysozyme C-3	<u>P00706</u>	129	Anas platyrhynchos	4e-05 (22 nd hit)
			STDYGIL+INSR	Lysozyme C	<u>Q7LZQ2</u>	129	Aix sponsa	4e-05 (23 rd hit)
B1	503.3	+2	WDAWDALK					
			WDAW+ALK	Acyl-coenzyme A binding protein	<u>Q9PRL8</u>	86	<u>Gallus gallus</u>	0.017 (1 st hit)
			WDAW AL	Acyl-CoA-binding domain-containing protein 5	<u>Q5ZHQ6</u>	492	<u>Gallus gallus</u>	0.76 (2 nd hit)
			W+ WDA	β -crystallin B3	<u>P55165</u>	211	<u>Gallus gallus</u>	11 (3 rd hit)
			WDAW	β -crystallin A3	<u>P10042</u>	215	Gallus gallus	11 (4 th hit)
			W+AW+ LK	Acyl-CoA-binding protein	<u>P45882</u>	103	Anas platyrhynchos	11 (5 th hit)
B4	509.8	+2	IGTMLPMQK					
			MLPMQK	Myeloid protein-1	<u>P08940</u>	326	Gallus gallus	0.35 (1 st hit)
			TMLPM	Brachyury protein	<u>P79777</u>	433	Gallus gallus	3.7 (2 nd hit)
			ML MQK	Eukaryotic translation initiation factor 3 subunit	<u>Q5F428</u>	564	Gallus gallus	12 (3 rd hit)
			TM LPMQ	Paired box protein Pax-6	<u>P47237</u>	216	Gallus gallus	12 (4 th hit)
			TM LPMQ	Paired box protein Pax-6	<u>P47238</u>	416	Coturnix japonica	12 (5 th hit)
B1	565.3	+2	LVTDVQEAVR					
			LVTDVQEAVR	Sapoin-A	<u>Q13035</u>	518	Gallus gallus	8e-04 (1 st hit)
			VTDV EA	Cadherin-4	<u>P24503</u>	913	Gallus gallus	13 (2 nd hit)
			LVT+VQ EA	Prolactin-2	<u>P55752</u>	199	Gallus gallus	13 (3 rd hit)

Table A.2. cont'd.

			LVT+VQ EA	Prolactin-1	<u>P55751</u>	199	Alligator mississippiensis	13 (4 th hit)
			VTD VQEA	Cystatin	<u>P08935</u>	111	Bitis arietans	13 (5 th hit)
			LVT+VQ EA	Prolactin-2	<u>P55754</u>	199	Crocodylus novaeguineae	13 (6 th hit)
			LVT+VQ EA	Prolactin-1	<u>P55753</u>	199	Crocodylus novaeguineae	13 (7 th hit)
			VTDVQ+	Fibronectin	<u>P11722</u>	1256	Gallus gallus	18 (8 th hit)
B2	646.9	+2	ISMPDFDLNLK					
			SMPDF	Myosin-Ig	<u>Q5ZMC2</u>	1007	Gallus gallus	7.4 (1 st hit)
			DFDL+L	Zinc finger FYVE domain-containing protein 27	<u>Q5ZL36</u>	406	Gallus gallus	18 (2 nd hit)
			M D+DL LK	Uncharacterized protein C17orf85 homolog	<u>Q5ZM19</u>	604	Gallus gallus	24 (3 rd hit)
			IS+PDF	Collagen α -3(VI) chain	<u>P15989</u>	3137	Gallus gallus	24 (4 th hit)
B2	772.0	+2	AVASAAAALVLK					
			AVASAAAALVLK	Talin-1	<u>P54939</u>	2541	Gallus gallus	2e-04 (1 st hit)
			VASAAAA	Insulin-like growth factor 2 mRNA-binding protein 3	<u>Q5ZLP8</u>	584	Gallus gallus	5.5 (2 nd hit)
			V +AA+ALVL	NADH-ubiquinone oxidoreductase chain 2	<u>Q21398</u>	346	Struthio camelus	7.3 (3 rd hit)
B2	525.8	+2	LGTFLENEK					
			LGTFL N EK	α -type phospholipase A2 inhibitor anMIP	<u>A1XRN2</u>	166	Atropoides nummifer	2.1 (1 st hit)
			LGTFL N	α -phospholipase A2 inhibitor clone 09	<u>B1A4R4</u>	166	Lachesis muta muta	6.7 (2 nd hit)
			LGTFL N	α -phospholipase A2 inhibitor	<u>B1A4R0</u>	166	Lachesis muta muta	6.7 (3 rd hit)
			LGTFL N	Phospholipase A2 myotoxin inhibitor protein	<u>Q8AYA2</u>	166	Bothrops moojeni	6.7 (4 th hit)
			LGTF LEN	Cannabinoid receptor 1	<u>P56971</u>	473	Taeniopygia guttata	8.9 (5 th hit)

Table A.2. cont'd.

B4	821.9	+2	VIQQADDAEER					
			VIQ QQ+D+ EE	Tenascin	<u>P10039</u>	1808	Gallus gallus	0.29 (1 st hit)
			I+Q+ADD EE	Transcription factor SOX-11	<u>P48435</u>	396	Gallus gallus	0.70 (2 nd hit)
			QQ DDAEER	Myosin heavy chain	<u>P29616</u>	1102	Gallus gallus	0.70 (3 rd hit)
			+Q +AD DAEER	Myosin-3	<u>P02565</u>	1940	Gallus gallus	1.7 (4 th hit)
			+Q +AD DAEER	Myosin heavy chain	<u>P13538</u>	1939	Gallus gallus	1.7 (5 th hit)
B6	587.3	+2	IPPKPPARAAR					
			+PPKPPA	G2/mitotic-specific cyclin-B2	<u>P29332</u>	399	Gallus gallus	1.7 (1 st hit)
			PPKP P RAA	Venom protease inhibitor 2	<u>B7S4N9</u>	88	Oxyuranus scutellatus	5.5 (2 nd hit)
			PP+PPAR	Eukaryotic translation initiation factor 3 subunit H	<u>Q5ZLE6</u>	348	Gallus gallus	5.5 (3 rd hit)
			PPKP A+AA	Transcription factor SOX-11	<u>P48435</u>	396	Gallus gallus	5.5 (4 th hit)
			PP PPAR AR	Histone H3-like centromeric protein A	<u>Q6XXM1</u>	131	Gallus gallus	7.4 (5 th hit)
			PARAAR	SCO-spondin	<u>Q2PC93</u>	5255	Gallus gallus	7.4 (6 th hit)
B5	506.8	+2	IPSLPSGVDK					
			IP LPSGV	Coatmer subunit Δ	<u>Q5ZL57</u>	510	Gallus gallus	13 (1 st hit)
			LPS VDK	116 kDa U5 small nuclear ribonucleoprotein component	<u>Q5F3X4</u>	972	Gallus gallus	18 (2 nd hit)
			LPSG+D	Neural cell adhesion molecule 1	<u>P13590</u>	1091	Gallus gallus	32 (3 rd hit)
			IP+LP+G	Hepatocyte nuclear factor 1-α	<u>Q90867</u>	634	Gallus gallus	32 (4 th hit)
B6	583.8	+2	AAGEIIAIPRR					
			AGEI+ IP	Jumonji domain-containing protein	<u>Q5ZHV5</u>	425	Gallus gallus	24 (1 st hit)

Table A.2. cont'd.

IAIPR	4 Lysine-specific demethylase 3A	<u>Q5ZIX8</u>	1325	Gallus gallus	24 (2 nd hit)
IIAIP	Cytochrome c oxidase polypeptide I	<u>079548</u>	533	Gallus gallus	24 (3 rd hit)
IIAIP	Cytochrome c oxidase subunit I	<u>Q94WR7</u>	516	Buteo buteo	24 (4 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O21399</u>	516	Struthio camelus	24 (5 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O03546</u>	337	Rhea americana	24 (6 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O03539</u>	337	Nothoprocta perdicaria	24 (7 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O03521</u>	337	Casuaris bennetti	24 (8 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O03515</u>	337	Apteryx australis	24 (9 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>P18943</u>	515	Gallus gallus	24 (10 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O03524</u>	337	Dromaius novaehollandiae	24 (11 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>P24984</u>	516	Coturnix japonica	24 (12 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>O03554</u>	337	Tinamus major	24 (13 th hit)
IIAIP	Cytochrome c oxidase subunit I	<u>Q79672</u>	514	Pelomedusa subrufa	24 (14 th hit)
AGEII	MBD1-containing chromatin-associated factor 1	<u>Q5ZIE8</u>	1085	Gallus gallus	43 (15 th hit)
IIPRR	Tenascin	<u>P10039</u>	1808	Gallus gallus	43 (16 th hit)
EIIA RR	Protein sidekick-1	<u>Q8AV58</u>	472	Gallus gallus	58 (17 th hit)
AAG IIAI	Alcohol dehydrogenase 1	<u>P49645</u>	375	Gallus gallus	58

Table A.2. cont'd.

								(18 th hit)
			AAG IIAI	Alcohol dehydrogenase 1	<u>P19631</u>	375	Coturnix japonica	58 (19 th hit)
			GEI IPR	Lysozyme C	<u>Q91159</u>	145	Opisthocomus hoazin	58 (20 th hit)
			GEI IPR	DNA damage-binding protein 1	<u>Q805F9</u>	1140	Gallus gallus	58 (21 st hit)
B1	831.4	+2	NSWGTSWGEDGYFR					
			NSWGT WG GY	Cathepsin K	<u>Q90686</u>	334	Gallus gallus	0.008 (1 st hit)
			NSWG WG+ GY	Cathepsin L1	<u>P09648</u>	218	Gallus gallus	0.12 (2 nd hit)
			NSW T WG G+F	Cathepsin B	<u>P43233</u>	340	Gallus gallus	0.29 (3 rd hit)
B5	684.9	+2	FRTTMLQDSIR					
			TML+DS+	Talin-1	<u>P54939</u>	2541	Gallus gallus	7.3 (1 st hit)
			MLQD+I	Hsp90 co-chaperone Cdc37	<u>Q57476</u>	393	Gallus gallus	7.3 (2 nd hit)
			FR+TM L+DS	78 kDa glucose-regulated protein	<u>Q90593</u>	652	Gallus gallus	7.3 (3 rd hit)
			MLQDS	5,6-dihydroxyindole-2-carboxylic acid oxidase	<u>Q57405</u>	535	Gallus gallus	13 (4 th hit)
			LQ+SIR	Coiled-coil domain-containing protein 132	<u>Q5ZKV9</u>	949	Gallus gallus	18 (5 th hit)
			TTML+ +I	Filensin	<u>Q06637</u>	657	Gallus gallus	18 (6 th hit)
B6	586.8	+2	LPPEQGTSSR					
			LPPEQG	Basic protease inhibitor	<u>P00993</u>	110	Caretta caretta	3.0 (1 st hit)
			LPPE G+ SS	Muscle-specific regulatory factor 4	<u>Q01795</u>	242	Gallus gallus	13 (2 nd hit)
			LPPEQ	Abhydrolase domain-containing protein FAM108C1	<u>Q5ZJX1</u>	310	Gallus gallus	18 (3 rd hit)

Table A.2. cont'd.

			LPPEQ	α -actinin-2	<u>P20111</u>	897	Gallus gallus	18 (4 th hit)
			PEQG TS	Dihydropyrimidinase-related protein 2	<u>Q90635</u>	572	Gallus gallus	18 (5 th hit)
			LPPEQ	α -actinin-4	<u>Q90734</u>	904	Gallus gallus	18 (6 th hit)
B6	593.3	+2	AILYNYWDK					
			AILYNY	Complement C3	<u>Q01833</u>	1651	Naja naja	0.85 (1 st hit)
			LYNYW	Ectonucleoside triphosphate diphosphohydrolase	<u>P79784</u>	495	Gallus gallus	1.1 (2 nd hit)
			LY+YW	Fatty acyl-CoA reductase 1	<u>Q5ZM72</u>	515	Gallus gallus	12 (3 rd hit)
			LY NYW+	Monocarboxylate transporter 9	<u>Q5ZJU0</u>	507	Gallus gallus	12 (4 th hit)
B6	506.8	+2	NEALIALLR					
			NEALIALL	Plastin-1	<u>P19179</u>	630	Gallus gallus	0.081 (1 st hit)
			NEA+IA LL	tRNA (cytosine)-methyltransferase NSUN2	<u>Q5ZLV4</u>	796	Gallus gallus	21 (2 nd hit)
			NE+LI L R	Ubiquitin carboxyl-terminal hydrolase 28	<u>Q5ZID5</u>	1047	Gallus gallus	21 (3 rd hit)
			EALI+L	CCR4-NOT transcription complex subunit 10	<u>Q5ZIW2</u>	744	Gallus gallus	39 (4 th hit)

APPENDIX B. LECTIN SEQUENCE

Protein sequencing with MS/MS and *de novo* sequencing yielded almost full coverage of an alligator lectin. The sequence was compared to lectins from other vertebrates using DIALIGN software to investigate sequence homology (Chapter 6, Section 6.4.4).

Figure 1. *Alligator mississippiensis* lectin sequence aligned with *Homo sapiens* and *Mus musculus* intelectin-2.

```

Human      MLSMLRMTMTRLCFLLFFSVATSGCSAAAASLEMLSREFETCAFSFSSLPRSCKEIKERC 60
Mouse      -----MTQLGFLLFIMIATRVCSSAAEEN----LDTNRWGNSF-FSSLPRSCKEIKQED 48
Alligator  -----NNQLALKLAATGGSTNXLPALALQN--LLNTWEDTSCCSQTSPPGQQSWPRD-- 49
              .:* : *   .:   .*   *       :   .: * . .  ::

Human      HSAGDGLYFLRITKNGVVYQTFCDMTSGGGWTLVASVHENDMRGKCTVGDRWSSQQGNKA 120
Mouse      TKAQDGLYFLRITENGVYIYQTFCDMTTAGGGWTLVASVHENNLGRCTVGDRWSSQQGNRA 108
Alligator  -GAQDGLYTLSTADGEIYQTFCDMSTHGGGWTLVASVHENNAHGKCTVGDRWSSQQGNSP 108
              * **** * * :* :*****: : *****: :*:***** .

Human      DYPEGDGNWANYNTFGSAEAATSDDYKNPGYDIQAKDLGIWHVPNKSPM--QHWRNSAL 178
Mouse      DYPEGDGNWANYNTFGSAEGATSDDYKNPGYFDIQAENLGIWHVNNNSPL--HTWRNSSL 166
Alligator  LYPEGDGNWANNNI FGSAMGSTSDDYKNPGYDLQAGDLSVWHVDPDRAPLRKEMIESSVL 168
              ***** * **** .:*****:*:** :* .:*****.:*: . . .* *

Human      LRYRTNTGFLQRLGHNLFGLIYQKYPVKYRSGKWCWVDNGPAIPVVYDFGDAKKTASYYSKY 238
Mouse      LRYRTFTGFLQRLGHNLFGLYQKYPVKYGEKGCWTDNGPAFPVVYDFGDAQKTASYYSKY 226
Alligator  LFYR--TGFLSSEGNLLRLEKYPVKYAGSCKVDNGPAVPVYDFGSAEKTAAYYSKY 226
              * ** **** . * **: :*:***** *.* ***** .:***** .:***:****

Human      GQREFVAGFVQFRVFNNERAANALCAGIKVTGCNTEHHCIGGGGFFPQPKRQCGDFSAF 298
Mouse      GRNEFTAGYVQFRVFNNERAASALCAGVRVTGCNTEHHCIGGGGFFPEFDPEECGDFAAF 286
Alligator  GRGFTAGFVQFRVFNNEKAPMALCSGLKVTGCNTEHHCIGGGGFFPEGNPRQCGDFPAF 286
              *: ** .*:*****:* . ***:*.:*****: .*. :***** .**

Human      DWDGYGTHVKSSCSREITEAAVLLFYR 325
Mouse      DANGYGTHIRYSNSREITEAAVLLFYR 313
Alligator  DWDGYGTHQSWSTSREMIESSVLLFYR 313
              * :***** * ***: *::*****

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Figure 2. *Alligator mississippiensis* lectin sequence aligned with *Homo sapiens* ficolin- α and ficolin- β .

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Ficolin-1      -MELSGATMARGLAVLLVLFHLHIKLNLPQAADT-CPEVKVVGLEGSDKLTILRGCPGLPG 58
Alligator     NNQLALKLAATGGSTNSLPALALQNLLNTWEDTSCCSQTSPPGQQSWPRDGAQDGLYTLST 60
               **:  * * : . : * : **      ** * . . * : . : * * .

Ficolin-1      APGPKGEAGVIGERGERGLPGAPGKAGPVGPKGDRGEKGMERGEKGDAGQSQSCATGPRNC 118
Alligator     ADG-----EIYQTFCDMSTHGGGWTLVASVHENNAHGKCTVGDRWSSQQG-NSPLYPEGD 114
               * *      :      : . * . . . . : : * * : : * . . . . * .

Ficolin-1      KDLLDRGYFLSGWHTIYLPDCRPLTVLCDMDTDGGGWTVFQRRMDGSVDFYRDWAAYKQG 178
Alligator     GNWANNNIFGSAMG----STSDDYKNPGYYDLQAGDLSVWH--VPDRAPLRKEMIESSVL 168
               :  : . . * * . . . . . * : . * . : * : : : . . : : .

Ficolin-1      FGSQLGFEFVLGNDNIHALTAQGSSELRVLDLDFEGNHQFAKYKSFKVADEAEKYKLVLG- 237
Alligator     LFYRTGFLSSEGGNLLRLYEKYPVKYAGSCKVDNGPAVP IYDFGSAEKTAAAYSPSGR 228
               : : * : . . * : * : . : . . . . . . . * * : : * *

Ficolin-1      -AFVGGSSAGNSLTGHNNNFSTKDQDNDVSSNCAEKFQGAWWYADCHASNLNGLYLMP 296
Alligator     GEFTAGFVQFRVFNNEKAPMALCSG-LKVTGCNTEHHCIGGGGFFPEGNPRQCDFPAFD 287
               * . . * . : : : : : : . . * : . * . : . . * :

Ficolin-1      HESYANGINWSAAKGYKYSYKVSEMKVRPA 326
Alligator     WDGYGTHQSWSTSREMIESSVLLFYR---- 313
               : . * . . . * : : : :

Ficolin-2      -MELDRAVGVLGAATLLLSFLGMAWALQAAD--TCPEVKMVGLESDKLTILRGCPGLPG 57
Alligator     NNQLALKLAATGGSTNSLPALALQNLLNTWEDTSCCSQTSPPGQQSWPRDGAQDGLYTLST 60
               : * : . . * . : * * . : * : : : * . . * : . : * * .

Ficolin-2      APGPKGE--AGTNGKRGERGPPGPPGKAGPPGPNAGPEQPCLTGPRTCKDLLDRGHFL 115
Alligator     ADGEIYQTFCDMSTHGGGWTLVASVHENNAHG-KCTVGDRWSSQQGNSPLYPEGD-GNWA 118
               * * : . . . : * . . : . . * : : * : . . * . * * :

Ficolin-2      SG--WHTIYLPDCRPLTVLCDMDTDGGGWTVFQRRVDGSVDFYRDWATYKQGFGRSRLGEF 173
Alligator     NNNIFGSAMGSTSDDYKNPGYYDLQAGDLSVWH--VPDRAPLRKEMIESSVLLFYRTGF 176
               . . : : . . . * : . * . : * : : * . . : : . : * * :

Ficolin-2      WLGNDNIHALTAQGTSELRVLDLDFEDNYQFAKYRSFKVADEAEKYNLVLG--AFVEGSA 231
Alligator     SSEGGNLLRLYEKYPVKYAGSCKVDNGPAVP IYDFGSAEKTAAAYSPSGRGEFTAG-F 235
               . . * : * : . : . . . : . . . * * : : * * * * . *

Ficolin-2      GDSLTFHNNQSFSSTKDQDNDLNTGNCAVMFQGAWWYKNCHVSNLNNGRYLRGTHGSFANGI 291
Alligator     VQFRVFNNEKAPMALCSGLKVTGCNTEHHCIGGGGFFPEGNPRQCDFPAFDWDGYGTHQ 295
               : . * : : : : : . . . . * * . : . . * : . . . .

Ficolin-2      NWKSGKGYNSYKVSEMKVRPA 313
Alligator     SWSTSREMIESSVLLFYR---- 313
               . * . : : * : :

```

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ANIMAL USE PROTOCOL

All investigators and instructors using species currently covered by federal statute and guidelines must complete this form and submit it to the Director of Research Services and Sponsored Programs (Campus Box 90655) for review by the IACUC a minimum of two weeks prior to initiating the protocol. The investigator will be informed by the Director of the IACUC of the committee's action and recommendations. IACUC evaluation of a proposal may require visiting the research site or animal facility to be used.

Date Received by RSSP _____
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Additional Comments Attached
IACUC DIRECTOR W. Owen (date) 22 Oct 09

1. Project Title (or laboratory exercise title): Collection of blood from crocodilians

2. Investigator(s)/Instructor(s): Mark Merchant
Department: Chemistry Phone: _____

3. Time period of protocol: From Oct 2009 to: Oct 2012

4. Attach a description of the experimental protocol and any supporting material to this form.

5. Name of funding source for this proposal: LEQSF

6. Description of Animals:

- Species and Common Name (if any) and number of animals:
American alligator (Alligator mississippiensis)
Broad-snouted caiman (Caiman latirostris)
Yacare caiman (Caiman yacare)
Orinoco crocodile (Crocodylus intermedius)
American crocodile (Crocodylus acutus)
Spectacled caiman (Caiman crocodylus)
Nile crocodile (Crocodylus niloticus)
Saltwater crocodile (Crocodylus porosus)

March 2006

★
(more on back)

- Freshwater crocodile (*Crocodylus johnstonii*)
- Dwarf crocodile (*Osteolaemus tetraspis*)
- Cuvier's dwarf caiman (*Paleosuchus palpebrosus*)
- Scheidt's dwarf caiman (*Paleosuchus trigonatus*)
- Black caiman (*Melanosuchus niger*)
- Slender-snouted crocodile (*Crocodylus cataphractus*)
- Phillipine crocodile (*Crocodylus mindorensis*)
- Morelet's crocodile (*Crocodylus moreletii*)
- Siamese crocodile (*Crocodylus siamensis*)
- New Guinea crocodile (*Crocodylus novaeguineae*)
- Cuban crocodile (*Crocodylus rhombifer*)
- Mugger crocodile (*Crocodylus palustris*)
- African dwarf croc (*Osteolaemus tetraspis*)
- False Gharial (*Tomistoma schlegelii*)
- Gharial (*Gavialis gangeticus*)

Source of animals (if wild, list current collecting permit number):

wild and captive

Location where animals will be housed: McNeese LERC alligator handling facilities

Location where animals will be used: Alligator handling facilities

7. Plan for disposition of sick animals (check all that apply):

Euthanize (state method) _____

Clinical Treatment

Other procedures (specify) cervical dislocation followed by PITING

8. Plan for disposition of dead animals (check all that apply):

Save, Freeze

Save, Refrigerate

Necropsy

Other, (Specify) _____

9. Is there a potential for pain, distress or discomfort to the animals?

Yes

No

If yes, use the appropriate number (see below) to indicate the maximum level of pain/discomfort/distress to be experienced by the animals.

Pain Level: 1

1 = Negligible, 2 = Pain avoided by appropriate drug use, 3 = Short-term discomfort/distress/pain, 4 = Long-term discomfort/distress/pain. (An explanation must be attached to the protocol for a pain level of 4.)

10. Federal law requires a written statement of A. and B. below to be included in protocols for any use of applicable species. Presently excluded from this requirement are rats and mice of the genus Rattus and Mus, birds, and animals used in accepted agricultural practices.

A. "The principal investigator/instructor has considered alternatives to procedures that may cause more than momentary or slight pain or distress to the animal, and has provided a written narrative description of the methods and sources used to determine that alternatives were not available".

B. "The principal investigator/instructor has provided written assurance that these activities do not unnecessarily duplicate previous experiments.

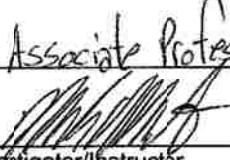
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McNeese State University
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I agree to comply with all Federal, State, and University animal welfare laws and policies during this project. I am aware that the new animal use protocol form must be submitted to RSSP and the IACUC if the protocol described above is altered after it is begun. The information provided is accurate, complete and true to the best of my knowledge.

Name (Print): Mark Merchant
Title: Associate Professor
Signature: 
Investigator/Instructor
Date: 10/09/09

March 2006



MCNEESE STATE UNIVERSITY

Mark E. Merchant, Ph. D.
Associate Professor of Biochemistry
P.O. Box 90455, Lake Charles, LA 70609-0455
Phone (337) 475-5773, Fax (337) 475-5950
mmerchan@mail.mcneese.edu

Chemistry Department

October 10, 2009

McNeese ACUC Committee,

This letter is written to serve as a resubmission of protocols used to collect serum, plasma, or whole blood from alligators (*Alligator mississippiensis*), and other crocodylian species (enumerated on ACUC form).

It is common practice for research personnel to collect blood from the spinal vein of crocodylians (Olson *et al.*, 1975, Zippel *et al.*, 2003). We typically use a 2.75 cm 20 ga. needle. The needle is inserted into the dorsal side of the neck, approximately halfway between the back of the skull and the first set of dorsal scutes. I have bled literally thousands of wild and captive alligators and crocodiles using this method during the past eight years. In addition, we bleed alligators *via* direct cardiac puncture on rare occasions using 26 ga. needles. The requirement to collect several blood samples in a relatively short time frame from small alligators during kinetic studies necessitates the cardiac bleed. We have found that repeated blood collection from the spinal vein of small alligators (< 1 m) sometimes compromises the integrity of the spine of the smaller animals. Cardiac puncture has been described by Lloyd and Morris (1999) and is used widely by other researchers using crocodylians as animal models (Busk *et al.*, 2003, Stevenson *et al.*, 1957).

If you have any questions or need additional information, please contact me at x5773.

Thank you,


Mark Merchant

- Busk M., Overgaard J., Hicks J., Bannet A., Wang T. (2003). Effects of feeding on arterial blood gases in the American alligator *Alligator mississippiensis*. *J. Exper. Biol.* 203 (20), 3117-3124.
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VITA

Lancia Nadina Fallen Darville-Bowleg was born in Freeport, Bahamas, to her parents, Christine Darville and Lancelot Darville Sr. Lancia graduated with a Bachelor of Science degree in 2003 from University of Nebraska at Kearney and a Master of Science degree in chemistry in 2005 from University of Nebraska-Lincoln. During her graduate study, her research was focused on the classification and identification of explosives to be developed into a chemical library for use by the Nebraska State Crime Lab.

In 2005, Lancia enrolled in the doctoral program at Louisiana State University and A&M College, where she joined Dr. Kermit K. Murray's research group. Lancia's research included application of mass spectrometry-based proteomics for identification, determination and characterization of immunological proteins from alligators. As a graduate student, Lancia has worked in collaboration with Dr. Mark E. Merchant and has submitted three articles to date. Her work has been presented at local and regional conferences, where she has given four oral presentations, twelve first-author and two collaborative poster presentations. Lancia's research has also been highlighted in both national and international science journals and magazines as well as local and national news broadcasts.

While attending LSU she received several fellowships and awards: Graduate Alliance for Education in Louisiana (GAELA) supplement, the Charles E. Coates travel award, Teaching Scholar Award for Excellence in Teaching and Outstanding Researcher. In 2009, she became affiliated with the Howard Hughes Medical Institute (HHMI) Math and Science Program as a science mentor where she executed leadership in the development and academic guidance of students interested in STEM areas. Her professional affiliations include the American Chemical Society (ACS), American Society of Mass Spectrometry (ASMS), the National Organization for

the Advancement of Black Chemists and Chemical Engineers (NOBCChE) and Iota Sigma Pi (ISP) National Honor Society of Women in Chemistry. Lancia is currently a candidate for the degree of Doctor of Philosophy in chemistry, which will be conferred at the December 2011 commencement.