PROTEIN BIOTOXINS OF MILITARY SIGNIFICANCE

Jiří Patočka¹, Ladislav Středa²

University of Defence, Faculty of Military Health Sciences, Czech Republic: Department of Toxicology¹ and University of South Bohemia, Faculty of Health and Social Studies, Czech Republic: Department of Radiology and Toxicology¹
State Office for Nuclear Safety, Czech Republic: Department for Control of the Prohibition of Chemical Weapons²

Summary: There is a spectrum of several threat agents, ranging from nerve agents and mustard agents to natural substances, such as biotoxins and new, synthetic, bioactive molecules produced by the chemical industry, to the classical biological warfare agents. The new, emerging threat agents are biotoxins produced by animals, plants, fungi, and bacteria. Many types of organisms produce substances that are toxic to humans. Examples of such biotoxins are botulinum toxin, tetanus toxin, and ricin. Several bioactive molecules produced by the pharmaceutical industry can be even more toxic than are the classical chemical warfare agents. Such new agents, like the biotoxins and bioregulators, often are called mid-spectrum agents. The threat to humans from agents developed by modern chemical synthesis and by genetic engineering also must be considered, since such agents may be more toxic or more effective in causing death or incapacitation than classical warfare agents. By developing effective medical protection and treatment against the most likely chemical and mid-spectrum threat agents, the effects of such agents in a war scenario or following a terrorist attack can be reduced. Toxin-mediated diseases have made humans ill for millennia. Unfortunately, the use of biological agents as weapons of terror has now been realized, and separating naturally occurring disease from bioterroristic events has become an important public health goal. The key to timely identification of such attacks relies on education of primary care physicians, first responders, and public health officials.

Key words: Biotoxin; Terrorism; Toxic protein; Ricin; Abrin; Viscumin; Volkensin; Modeccin; Conotoxin; Botulinum toxin; Clostridium perfringens toxins; Diphteriae toxin; Staphylococcus toxin; Shigatoxin; Verotoxin; Cholera toxin; Tetanus toxin; Microcystin

Introduction

Toxin-mediated diseases have made humans ill for millennia. They also have been used in beneficial ways. Unfortunately, the use of biological agents as weapons of terror has now been realized, and separating naturally occurring disease from bioterroristic events has become an important public health goal. The key to timely identification of such attacks relies on education of primary care physicians, first responders, and public health officials.

Natural toxins are toxic compounds produced by living organisms. There are principle toxic substances of poisonous animals, plants, microorganisms and other forms of life. The manner of their toxic effect can be very different in individual substances. According to the organ toxicity we distinguish neurotoxins, hepatotoxins, nephrotoxins, hemotoxins etc. For example neurotoxins are toxic agents or substances that inhibit, damages or destroys the tissues of the nervous system, especially neurons, the conducting cells of your body’s central nervous system. Neurotoxic effects can include behavior changes, seizures, as well as wide range of effects, including death. According to the chemical structure it is possible to divide all natural toxins (biotoxins) into nonprotein and protein compounds.

Chemical Weapons Convention (CWC) (1993) includes toxins as chemical agents, and specifically includes toxins in its control regime along with other highly toxic chemicals. There are these protein toxins: Abrin, botulinum toxins, Clostridium perfringens toxins, Corynebacterium diphteriae toxin, microcystins, Staphylococcus aureus toxin, ricin, shigatoxin, and tetanus toxin. On the „control list“ of the Australian Group, voluntary association of 33 countries established in 1985, also conotoxins, verotoxin, cholera toxin, modeccin, volkensin and viscumin are enregistered. Biotoxins have been employed in warfare and in terrorist activities.

¹The 33 states participating in the Australia Group are Argentina, Australia, Austria, Belgium, Bulgaria, Canada, Cyprus, Czech Republic, Denmark, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Japan, Luxembourg, the Netherlands, New Zealand, Norway, Poland, Portugal, Romania, Slovakia, South Korea, Spain, Sweden, Switzerland, Turkey, United Kingdom, and United States. The European Commission also participates. Several other countries, including Russia, Ukraine, India, and China, have national export controls for some, but not all, of the items on the group’s lists.
attacks (61). Botulinum toxin produces a descending flaccid paralysis. Staphylococcal enterotoxin B produces a syndrome of fever, nausea, and diarrhea and may produce a pulmonary syndrome if aerosolized. *Clostridium perfringens* epsilon-toxin could possibly be aerosolized to produce acute pulmonary edema. Ricin intoxication can manifest as gastrointestinal hemorrhage after ingestion, severe muscle necrosis after intramuscular injection, and acute pulmonary disease after inhalation (21). All these biotoxins may be perilous if will misuse by terrorists (67). A brief review of nonprotein biotoxins has been published recently (51). This review covers peptide toxins which are on the „control list” of chemical weapons. Each toxin is briefly characterized chemically, pharmacologically and toxicologically. Their natural sources, availability, stability, and military potential are discussed.

**Chemical classification of protein toxins**

All peptide toxins are created by one of more linear or cyclic polypeptide chains. That polypeptide is constructs from more amino acids connected by means of peptide linkage.

Proteins are large macromolecules composed of one or more peptide chain. It is possible to say that polypeptides composed from less number of amino acids are designated as peptides or polypeptides, from the larger number as proteins. The crossover between peptides and proteins is not defined exactly and is unsubstantial from practice point of view.

**Biotoxins as chemical weapons of warfare and terrorism**

Biotoxins can be employed in warfare and in terrorist attacks. In this era, it is imperative that health care providers are familiar with illnesses caused by these agents. Botulinum toxin produces a descending flaccid paralyisis (50). Staphylococcal enterotoxin B produces a syndrome of fever, nausea, and diarrhea and may produce a pulmonary syndrome if aerosolized. *Clostridium perfringens* epsilon-toxin could possibly be aerosolized to produce acute pulmonary edema. Ricin, abrin and other plant protein toxin intoxication can manifest as gastrointestinal hemorrhage after ingestion, severe muscle necrosis after intramuscular injection, and acute pulmonary disease after inhalation. Health care providers should be familiar with the medical consequences of toxin exposure, and understand the pathophysiology and management of resulting illness (21).

**Plant toxins**

Plant toxic proteins belong to a group of phytoxins, which inhibit the protein synthesis of eukaryotic cells. The toxins of this group are glycoproteins with molecular weights of about 60 kDa which consist of two subunits linked into a dimer by a disulfide bond. One of the subunits is lectin with sites for carbohydrate binding (B-chain), and the other subunit is specific N-glycosidase (A-chain), which modifies the 28S rRNA-60S ribosomal subunit (23). The group of proteins known as the lectins was first recognised in plant seeds and they bind to specific sugars. Although lectins, in general, are not very toxic, there are some relationships between lectins and toxins (66). They may also serve as recognition markers in cellular differentiation and act as immunotoxin. Only A-chain is toxic due to inhibiting protein synthesis. The A-chain catalytically inactivates 60S ribosomal subunit by removing adenine from positions 4 and 324 of 28S rRNA. Ribosome-inactivating proteins (RIP) have been identified in many plants and some of them are very toxic, but only in connection with B-chain. The A-chain is carrier of toxicity, but B-chain binds to cell surface receptors and facilitate a transport of the A-chain across the cell membrane. The A-chain is not active until it internalized by the cell, where halts protein synthesis(63). Each toxin molecule can disable approximately 2000 polysomes per minute, enough to eventually kill the cell. The most known plant toxic proteins of this type are ricin, abrin, modessin, viscumain and volkensin (47,51).

**Ricin**

Ricin is a protein produced by the castor oil plant, *Ricinus communis*. It is native to tropical Africa, but has naturalised sub-tropical and temperate areas as well. The whole of the plant is poisonous, containing the toxin ricin, which reaches the highest levels in the seeds. The seeds also contain a purgative oil, the triglyceride of ricinoleic acid. The seeds have been used in folk medicine against many diseases for centuries.

Ricin is the compound responsible for toxicity of the seeds of *R. communis* (58). Ricin has been known as a poison for years, usually through livestock deaths. One to three seeds may be fatal to a child; two to four may be poisonous to an adult, while eight may be fatal. A fatal ingested dose for human is about 1 µg /kg (10). Because the alimentary tract destroys lots of ricin, it is much more potent when administered paraenterally. The toxically active A-chain of ricin is a 267 amino acid globular protein and is classed as an N-glycosidase. The A-chain is protein containing 8 alpha helices and 8 beta sheets (29,39). The B-chain is composed of 262 amino acid residues and is classed as a lectin (29). The B chain has an affinity for bindig to galactosides (58) and possesses two galactose binding sites that are attracted to galactose containing glycoproteins at the cell surface. The A- and B-chain glycoproteins are linked by a disulfide bond located at residue 259 of the A-chain and residue 4 of the B-chain (38). Ricin is a glycoprotein with carbohydrate side chains in the form of mannose-rich N-linked oligosaccharides and particularly binds to mannose receptors of cells of the reticuloendothelial system. The specific sites with potential for binding of high mannose carbohydrate chains of ricin are at asparagines 10 and 236 of the A-chain, and asparagines 95 and 135 of the B-chain (58).
The toxic effects of ricin are essentially the result of the action of the A-chain, which inactivates the ribosomes of the cell. It is thought that the B-chain serves to bind to galactose-containing and/or mannose-containing structures at the cell surface thus allowing the A-chain to enter the cell (44). By this way the protein is internalized, meaning that it is taken into the cell as a toxin-binding site complex in vesicles. The B-chain of ricin facilitates the escape of the A-chain by binding to the endosomal membrane allowing the A-chain to pass through the membrane. Then, the B-chain dissociates from the A-chain by breaking the disulfide bond. The A-chain is thus delivered to the cytoplasm where it is taken up by the Golgi apparatus and transported to the endoplasmic reticulum (44).

At the endoplasmatic reticulum, ricin inactivates the cell ribosomes by elimination of adenine in specific RNA sequences on the 28S ribosomal subunit (33). A single A-chain molecule is capable of deactivating every ribosome in the cell thus halting protein synthesis and culminating in cell death. It is important to note that the toxin is specific for eukaryotic ribosomes. A hairpin loop on the 28S rRNA containing the tetranucleotide loop GAGA is the most likely target for attack by ricin on the ribosome, however it is thought that the ribosome conformation is an important factor in recognition by the protein (33). Therefore, ricin is not a nucleotide sequence-specific protein.

Ricin is the only toxin to exist naturally in large quantities. It is a byproduct of castor oil production and ricin isolation is a simple and cheap separation. Easy preparation and low price might make this toxin attractive to poor country. For nations or terrorists who lack the money to spend on nuclear weapons and other high-tech killing instruments, toxin warfare offers horrific appeal: biological/toxin weapons are cheap, easy to make, and simple to conceal. Even small amounts, if effectively used, could cause massive injuries and make millions suffer (54). Ricin’s significance as a potential biological warfare toxin relates in part to its wide availability. Worldwide, one million tons of castor beans are processed annually in the production of castor oil and in the waste is five percent ricin by weight. The toxin is also quite stable and extremely toxic by several routes of exposure, including the respiratory route (53).

In biological warfare it is expected that ricin would be released as a toxic cloud. It could also be injected into specific persons as a terrorist or sabotage weapon. Additionally, ricin is easy to produce and is stable. The toxic effects of ricin occur because it kills the cells of the body that it contacts when it is taken into the body. Upon inhaling an adequate amount of ricin, death of persons affected would be expected in 36-48 hours because of difficulty breathing and circulatory system effects. Ingested ricin is expected to cause internal bleeding, death of vital organs and death of the individual. Injected ricin causes death by major organ failure.

The prediction of symptoms to be expected is based on animal studies and accidental human exposures, which were not fatal. Symptoms would probably vary depending on whether ricin was inhaled, ingested or injected. About three hours after inhaling ricin, the symptoms expected are cough, tightness of the chest, difficulty breathing, nausea and muscle aches. This would progress to a severe inflammation of the lungs and airways, increased difficulty breathing, cyanosis and death in 36-48 hours from failure of the breathing and circulatory systems. Ingestion of ricin would be expected to cause nausea and vomiting, internal bleeding of the stomach and intestines, failure of the liver, spleen and kidneys and death of the individual by collapse of the circulatory vessels. No specific affects on the lungs and airways would be expected. If injected, ricin causes marked death of muscles and lymph nodes near the site of injection and probable failure of major organs and death of the individual (52).

Abrin

Abrin is a potent toxin that has been isolated from the seeds of Abrus precatorius (or Rosary pea). Its use as a tool for research was described in 1972 by Sharon and Lis (60). Abrin exists in two forms, abrin a and abrin b. Both are composed of two chains, an A-chain and a B-chain. A disulfide bond between Cys247 of the A-chain and Cys8 of the B-chain links the A and B chains. The A-chain is 251 residues and is divided into 3 folding domains. The A-chain catalytically inactivates 60S ribosomal subunits by removing adenine from positions 4 and 324 of 28S rRNA therefore inhibiting protein synthesis. The B-chain is a galactose specific lectin that facilitates the binding of abrin to cell membranes (7,45). The B-chain of both forms of abrin consists of 268 amino acid residues and shares 256 identical residues. Comparison of their sequences with that of the ricin’s B-chain shows that 60% of the residues of abrin’s B-chain are identical to those of the ricin’s B-chain and that two saccharide-binding sites in ricin B-chain identified by a crystallographic study are highly conserved in abrin B-chain (30).

The mechanism of toxic action of abrin is identical to that of ricin (53) but the toxicity of abrin in mice is 75 times higher that of ricin (0.04 µg/kg for abrin compared to 3 µg/kg for ricin). The diagnosis, clinical features, treatment, protection, prophylaxis and so on is also the same for both abrin and ricin intoxications (46, 48).

Viscumin

Viscumin (Mistletoe lectin I, ML I), inevitable to the family of RJP’s, was identified in the late 1980’s as the main pharmacologically-active ingredient of mistletoe (Viscum album) extract and is largely responsible for its toxicity (31). It is comparable in toxicity to ricin and acts by the same mechanism. When viscumin binds to its target cell, protein synthesis in that cell is interrupted as a result of the A-chain’s enzymatic activity, like a ricin. This interruption induces a cellular stress response, which triggers the release of cytokines by the target cell and, at high viscumin con-
centrations, apoptosis of the cell. The association of A- and B-subunits is predominantly hydrophobic in nature.

**Volkensin**

Volkensin is a lectin from *Adina volkensii* (the kilyambiti plant) that is comparable in toxicity to ricin and that acts by the same mechanism (a ribosome-inactivating protein) (6). The plant is a relatively unattractive and toxic succulent plant found in Africa that appears to be of little interest. However, it has proven useful as a research reagent in neurology because of its ability to be taken up and transported by certain types of nerve. There may be pressure to develop commercial sources for the research community (52).

**Modeccin**

Modeccin is a lectin from the roots of *Adenia digitata* an African succulent plant that is comparable in toxicity to ricin (43) and acts by the same mechanism (46,56). The plant does not seem to have any significant uses, such as a food or medicine and so is not available in quantities comparable to abrin, let alone ricin. However, the seed does seem to be readily available. The subunits of modeccin were isolated (subsequently referred to as modeccin 4B), purified from the roots of *Adenia digitata* by affinity chromatography on Sepharose 4B (17). They are an A subunit (mol. wt. 26 000), which inhibits protein synthesis, and a B subunit (mol.wt. 31 000), which binds to cells. A second form of modeccin, not retained by Sepharose 4B, was purified by affinity chromatography on acid-treated Sepharose 6B: this form is subsequently termed modeccin 6B. Modeccin 6B has a molecular weight indistinguishable from that of modeccin 4B, and consists of two subunits of mol.wts. 27 000 and 31 000, joined by a disulphide bond. The subunits were not isolated because of their high insolubility in the absence of sodium dodecyl sulphate. As compared with modeccin 4B, modeccin 6B is slightly less toxic to animals, does not agglutinate erythrocytes, and is a more potent inhibitor of protein synthesis in a lysate of rabbit reticulocytes, giving 50% inhibition at the concentration of 0.31 μg/ml (2).

**Animal toxins**

There are numerous protein toxins produced by many different animal sources, such as snakes, scorpions, spiders, insects, frogs, sea anemones etc. Only conotoxins, the toxic principle of marine snails of the genus *Conus*, are presented on the control list of the Australian Group.

**Conus toxins (conotoxins)**

These are toxic peptides produced by the fish-hunting marine snails of the genus *Conus*. The conotoxins (the snails produce a mixture of toxic substances) are used to paralyse the fish being attacked (32). For centuries members of the Conidae family have been collected for their unique and intricately designed shells. Only during the last few decades have Cone shells become an exciting area for scientific research. Cone shells are marine snails and are found in reef environments throughout the world. They prey upon other marine organisms, immobilising them with unique venoms. They can be dangerous for human, too. There have been 30 recorded cases of human envenomation by fish-eating cone shells, in some cases fatal (15).

The composition of the venom differs greatly between species and between individual snails within each species, each optimally evolved to paralyse it’s prey. The active components of the venom are small peptides toxins, typically 12–30 amino acid residues in length. They have very tight conformations by multiple disulfide bridges. These patterns of disulfide bridge help to define a number of structural classes of conotoxin. Today several tens of conotoxins is known and these are divided into four group. Alpha-conotoxins are the shortest and have only two disulfide bridges. Mu-conotoxins, omega-conotoxins, delta-conotoxins, and kappa-conotoxins contain three disulfide bridges between different cysteine residues of peptide chain (42).

The paralytic components of the venom that have been the focus of recent investigation are the alpha, omega and mu-conotoxins. All of these conotoxins act by preventing neuronal communication, but each targets a different aspect of the process to achieve this. The alpha-conotoxins target nicotinic ligand gated channels, the mu-conotoxins target the voltage-gated sodium channels and the omega conotoxins target the voltage-gated calcium channels.

Another class of peptides from Conus venoms are the conantokins, the first peptide antagonists which target to the major excitatory receptors in the vertebrate central nervous system, glutamate receptors. The conantokins selectively inhibit a subtype of glutamate receptor, the N-methyl-

---

### Tab. 1: Individual types of conus toxins and their biological targets.

<table>
<thead>
<tr>
<th>alpha</th>
<th>mu</th>
<th>omega</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nicotinic acetylcholine receptors. The effect is a paralysis similar to that seen with curare.</td>
<td>Sodium channels. This is also the target for saxitoxin and tetrodotoxin and the effects are similar.</td>
<td>Calcium channels associated with nerve impulse transmission at the neuromuscular junction.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>delta</th>
<th>kappa</th>
<th>conantokins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sodium channels. Unlike mu conotoxins, they slow the inactivation of the sodium channel.</td>
<td>Potassium channels. They are also known as shaker peptides because they block a potassium channel known as „Shaker“ and as a result they induce tremors.</td>
<td>NMDA glutamate receptors. This blocks nerve impulses that use glutamic acid rather than acetylcholine as the neurotransmitter.</td>
</tr>
</tbody>
</table>
D-aspartate (NMDA) receptor, which are ligand-gated Ca channels. The conantokins cause rather striking and till this time little explored biological effects (5).

A short outline of all known conus toxins and their targets is summarized in Table I.

Individual conotoxins, even within the same class can vary greatly in lethality towards mammals. Some of the tremor inducing omega conotoxins are not lethal, whereas others of the same group are lethal at low levels. However, they are only toxic in rats and mice when administered intracranially (into the brain). Some alpha conotoxins have lethal doses as low as 25 μg/kg for mouse. This may be an overestimate of toxicity because it is determined from the dose required to kill a mouse in 20 minutes. In addition, it has to be borne in mind that the toxicity of the complex mixture of peptides that is cone snail venom may be much greater than the sum of its parts because of the synergistic interaction between toxins acting on different aspects of neural function. Incidents of cone snails killing people are known to have occurred.

**Microbial toxins**

Protein toxins represent numerous group of toxic compounds, produced by different pathogenic bacteria. These bacterial toxins are in this review: botulinum toxins, *Clostridium perfringens* toxins, *Corynebacterium diphtheriae* toxin, *Staphylococcus aureus* toxins, shigatoxin, verotoxin, cholera toxin, and tetanus toxin.

**Botulinum toxins**

Botulinum toxin is derived from the genus of anaerobic bacteria named *Clostridia*. Seven antigenic types of botulinum toxin exist, designated from A through G (49). They can be identified based on antibody cross reactivity studies. *Clostridium botulinum*, is a familiar bacterium that causes botulism, a form of food poisoning (8). Naturally occurring botulism is the disease that results from the absorption of botulinum toxin into the circulation from a mucosal surface (gut, lung) or a wound. It does not penetrate intact skin. The botulinum toxin into the circulation from a mucosal surface or a wound. It does not penetrate intact skin. The main biological activity of epsilon-toxin is the production of oedema in various organs and cytoskeletal changes (51).

**Clostridium perfringens toxins**

The Gram-positive pathogen *Clostridium perfringens* is a major cause of human and veterinary enteric disease largely because this bacterium can produce several toxins when present inside the gastrointestinal tract. *C. perfringens* food poisoning is one of the most common in the industrialized world (62). The enteric toxins of *C. perfringens* share two common features: 1) they are all single polypeptides of modest ( ~25–35 kDa) size, although lacking in sequence homology, and 2) they generally act by forming pores or channels in plasma membranes of host cells (55). These enteric toxins include *C. perfringens* enterotoxin (CPE), which is responsible for the symptoms of a common human food poisoning and acts by forming pores after interacting with intestinal tight junction proteins. Two other *C. perfringens* enteric toxins, epsilon-toxin (a bioterrorism select agent) and alpha-toxin, cause veterinary enterotoxemias when absorbed from the intestine. *C. perfringens* enterotoxin (beta-toxin) has been shown to be the virulence factor responsible for causing the symptoms of *C. perfringens* food poisoning. Beta-toxin is a single polypeptide chain with a molecular weight of 3.5 kDa that binds to receptors on the target epithelial cells. Through a unique four-step membrane action it finally causes a breakdown in normal plasma membrane permeability properties (4, 55). Also beta- and epsilon-toxins apparently act by forming oligomeric pores in intestinal or extra-intestinal target tissues. Other *C. perfringens* toxins have different effect. *C. perfringens* alpha-toxin is able to lyse erythrocytes via calcium channels activation (41). The main biological activity of epsilon-toxin is the production of oedema in various organs and cytoskeletal changes and plasma membrane functional alteration (13).

**Corynebacterium diphtheriae toxin**

Diphtheria toxin is an extracellular protein of *Corynebacterium diphtheriae* that inhibits protein synthesis and kills susceptible cells (26). *C. diphtheriae* is responsible for diphtheria. Diphtheria is a contagious, airborne, toxin-producing infection caused by *C. diphtheriae*. It is characterized by the formation of a gray resistant pseudo-membrane in the lining of the mucous membrane of the upper respiratory tract as well as in the tonsils. Certain forms of the disease may be fatal. The global mortality rate for diphtheria is 5% to 10% and may reach 20% among children under 5 and adults over 40. In 1888, Roux discovered the diphtheria toxin secreted by *C. diphtheriae*, an agent of diphtheria. In 1890, the work of von Behring and Kitasato on antibodies to diphtheria antitoxins made it possible to envision their use in treating the disease (22). In 1897, Ehrlich established a standardized diphtheria toxin. These passive serum therapies would soon lead to active immunization. Diphtheria vaccines were first used in France in the 1920s. Mass immunization only began in the 1950s. The diphthe-
ria vaccines used today throughout the world against the pathogenic and lethal effects of the diphtheria toxin are obtained by detoxification of the toxoid with formalin.

**Staphylococcus toxins**

*Staphylococcus aureus* is a spherical bacterium (coccus) which on microscopic examination appears in pairs, short chains, or bunched, grape-like clusters. These organisms are gram-positive. Some strains are capable of producing a highly heat-stable protein enterotoxins, range in size from 19 to 26 kDa, that cause illness in humans. Seven immunologically different forms of *Staphylococcus* enterotoxins is known: A, B, C₁, C₂, C₃, D and E. These toxins are responsible for symptoms of food poisoning that follow consumption of food contaminated by *Staphylococcus* bacteria.

The onset of symptoms in staphylococcal food poisoning is usually rapid and in many cases acute, depending on individual susceptibility to the toxin, the amount of contaminated food eaten, the amount of toxin in the food ingested, and the general health of the victim. The most common symptoms are nausea, vomiting, retching, abdominal cramping, and prostration. Some individuals may not always demonstrate all the symptoms associated with the illness. In more severe cases, headache, muscle cramping, and transient changes in blood pressure and pulse rate may occur. Recovery generally takes two days. However, it is not unusual for complete recovery to take three days and sometimes longer in severe cases.

**Infective dose –** a toxin dose of less than 1.0 µg in contaminated food will produce symptoms of staphylococcal intoxication. This toxin level is reached when *S. aureus* populations exceed 100,000 per gram.

*S. aureus* produce also hemolytically active toxins, so called hemolysins. Three hemolysins, alpha toxin, beta toxin, and delta toxin, are known and each of them exist in more molecular forms. Their mechanism of toxic action is different (25).

Alpha toxin, also known as alpha hemolysin, is one of many virulence factors produced by *Staphylococcus aureus* (16). Alpha toxin is a single polypeptide chain known in four molecular forms with a molecular weight from 26 to 39 kDa. The majority of *S. aureus* strains isolated from humans produce this toxin. Alpha toxin has been shown to be lethal in animals, causing respiratory paralysis, vascular and smooth muscle spasms, and tissue necrosis (64).

Beta toxin (beta-hemolysin) is one of several extracellular proteins produced by *S. aureus*. It is a sphingomyelinase which disrupts the membranes of erythrocytes and other mammalian cells. Despite its characterized mechanism of action, the role of beta-toxin in human and animal disease remains unclear (34).

Delta toxin (delta hemolysin) is heat-stable peptide, molecular weight 5 kDa, composed from 26 amino acid residues. Its molecule create amphiphatic helix with hydrophilic amino acid residues on one side of long axis of molecule, hydrophobic residues on other side and produced lesions in biological membranes very similar to those produced by melittin and Triton X-100 (16).

**Shigatoxin**

*Shigella* are the most important organisms can cause dysentery. *Shigella dysenteriae* type 1 (Sd1) is the most virulent of the four serogroups of *Shigella*. Sd1 is the only cause of epidemic dysentery. In addition to bloody diarrhoea, the illness caused by Sd1 often includes abdominal cramps, fever and rectal pain. Less frequent complications of infection with Sd1 include sepsis, seizures, renal failure and the haemolytic uraemic syndrome (HUS) (19). Altered arachidonic acid metabolism has been implicated in the pathogenesis of renal injury in the HUS caused by shigatoxin (59). Approximately 5–15% of Sd1 cases are fatal. Shigatoxin works as enterotoxin, neurotoxin and cytotoxin. It is consist of two protein subunit: Subunit A perform as N-glycosidase which split adenin on ribosomes and inhibits proteosynthesis (40).

**Verotoxin**

The verotoxin, or shiga-like toxin family is a group of closely related toxins produced by certain pathogenic strains of *Escherichia coli*. Verotoxin-producing *E. coli*, especially of serotype O157:H7, cause a zoonotic food or waterborne enteric illness that is often associated with large epidemic outbreaks as well as the HUS, the leading cause of acute renal failure in children (27). These strains are a significant cause of human hemorrhagic colitis. In addition, they are both water borne and food borne and may also be transmitted from person-to-person by the oral-fecal route. In adults, illness caused by verotoxin may last several days. In children and the elderly, the illness can be fatal (20). Structure and mechanism of toxic action of verotoxin is equal shigatoxin but recently new pieces of knowledge were obtained. It has been known for some time that following the intracellular routing of shigatoxin and/or verotoxin to the endoplasmic reticulum and nuclear membrane, the toxins translocate into the cytoplasm and target ribosomes for damage. However, numerous recent studies have shown that these toxins trigger programmed cell death signaling cascades in intoxicated cells. The mechanisms of apoptosis induction by these toxins are newly emerging, and the data published to date suggest that the toxins may signal apoptosis in different cells types via different mechanisms (9).

**Cholera toxin**

*Vibrio cholerae* is a Gram-negative, curved rod bacteria. *V. cholera* is responsible for approximately seven pandemic infections of extreme diarrhea and dehydration across the globe, resulting in millions of death throughout the centuries. *V. cholerae* as pathogen has been with mankind long before we were ever capable of detecting it, or understanding its mechanism of infection. Until recent times, the methods to confront this pathogen, and begin to understand its function were not available.
The biochemistry of cholera toxin has been well characterized over recent years (12,37). Cholera toxin is a heterohexameric protein, composed from five subunits B and one subunit A (AB5, choleragen), responsible for the symptoms produced by *V. cholerae* infection. In the first step of cell intoxication, the B-pentamer of the toxin binds specifically to the branched pentasaccharide moiety of ganglioside GM1 on the surface of target human intestinal epithelial cells.

The normal function of GM1 is not clearly understood but it has been implicated in numerous signal transduction pathways. Cholera toxin and GM1 complex show no major conformational change, but it is theorized about that the A subunit of the cholera toxin is inserted into the cell. This translocation is takes approximately 15 minutes and during this time it is theorized that the A subunit is cleaved along a disulfide bridge between A1 and A2, which keeps the protein inactive (36).

The A1 subunit of cholera toxin is the enzymatically active portion of the protein molecule, and it acts as an ADP-ribosyltransferase. It catalyzes a transfer of an ADP-ribose from an NAD+ to the arginine at location 187 in the alpha chain of the regulatory protein Gs (28). This ribosylation of Gs stabilizes the GTP bound form of the protein, lower its GTPase activity creating a near constitutively on signal for the generation of adenyl cyclase, and therefore elevating cAMP levels. This high cAMP level results in the activation of the sodium pumps in the lumen of the cell through the acAMP dependent kinase pathway, forcing out Na+ ions. The resultant electrochemical imbalance drives out Cl- and H2O to balance the Na+ release. The net flow of fluid is now out of balance and the cells attempt to compensate for the dehydration through removal of fluid from the blood. This is the biochemical process responsible for the clinical characteristics of epidemic cholera infection. (14).

### Tetanus toxin

The illness known as tetanus is caused by a neurotoxin produced by the anaerobic bacterium *Clostridium tetani*. It acts upon the presynaptic membranes of both central and peripheral nervous systems to block the release of neurotransmitters. Tetanus toxin is synthesised as a single polypeptide chain of 150 kDa, and undergoes proteolytic cleavage to produce a di-chain toxin consisting of the N-terminal 50 kDa fragment (light chain) linked by a disulphide bond to the 100 kDa carboxy terminal fragment (heavy chain). Fragment C (the C-terminal half of the heavy chain) retains ganglioside binding activity which is essential for the binding of the toxin to neuronal cells (1,24). Tetanus toxin is likewise botulinum toxin a neurotoxin and their molecular structures and mechanism of action are very similar (65).

The LD50 in unvaccinated humans is estimated at <2.5 ng/kg (18). It is a powerful neurotoxin which may be fatal if inhaled or introduced into a wound. It causes muscle rigidity or spasms, paralysis, and death. If contact occurs, flush eyes, skin or wounds thoroughly with water. Seek medical attention, since supportive therapy will be required if symptoms occur. Immune globulin may also be a part of the medical treatment.

### Blue-green algae toxins

Algae are small, often microscopic organisms. Blue-green algae called also cyanobacteria are the most primitive form of life. Only certain species of blue-green algae are capable of producing toxins and even these species are harmless most of the time. The presence of toxic algae is frequent reason for swimming prohibition in swimming pool or bathing in countryside (35). Toxic algae periodically bloom in freshwater aquatic sheet as well as in coastal waters, sometimes poisoning seabirds and marine mammals and interfering with economically important fisheries. These microorganisms are rich source of different very toxic compounds. There are largely neurotoxins and hepatotoxins (53). Only microcystins and nodularin are peptides and only microcystins are on the „control list” of the Australian Group.

### Microcystins

The microcystins are a group of cyclic heptapeptide hepatotoxins produced by a number of cyanobacterial genera, the most notable of which is the widespread *Microcystis* from which the toxins take their name. Microcystins consist of a seven-membered peptide ring, which is made up of five non-natural amino acids and two natural amino acids. It is these two protein amino acids that distinguish microcystins from one another, while the other amino acids are more or less constant between variant microcystins. Using amino acid single letter code nomenclature, each microcystin is designated a name depending on the variable amino acids which complete their structure. The most common and potentially toxic microcystin-LR contains the amino acids leucine (L) and arginine (R) in these variable positions.

Approximately 60 different microcystins was identified till this time. The general stucture of microcystins (Fig. 1) showing the variable amino acid positions “X” and “Y”. The amino acids are delineated in this diagram and num-

![Microcystins](https://example.com/microcystins.png)

**Fig. 1:** The common chemical structure of microcystins. Adda = 3-amino-9-methoxy-2,6,8-trimethyl-10-phenyldeca-4, 6-dienoic acid, D-Glu = D-glutamic acid, Mdha = N-methyldehydroalanine, D-Ala = D-alanine, Masp = D-methylaspartic acid, and X and Y are both variable L-amino acids.
bered according to the microcryst standard nomenclature. R1 and R2 are H in demethylated microcystins. These monocyclic heptapeptides are characterised by some invariant amino acids, including one of unusual structure - 3-amino-9-methoxy-2,6,8-trimethyl-10-phenyldeca-4,6-dienoic acid (Adda) - which is essential for expression of toxicity. (53). Microcystins are chromophore stable, but suffer biodegradation in reservoir waters. The most common form of the family, microcystin-LR (L and R identifying the 2 variable amino acids, in this case leucine and arginine respectively) has an LD50 in mice and rats of 36–122 µg/kg by various routes, including aerosol inhalation. Although human illnesses attributed to microcystins include gastroenteritis and allergic/irritation reactions, the primary target of the toxin is the liver, where disruption of the cytoskeleton, consequent on inhibition of protein phosphatases 1 and 2A, causes massive hepatic haemorrhage. Microcystins are tight-binding inhibitors of these protein phosphatases, with inhibition constants in the nanomolar range or lower. Uptake of microcystins into the liver occurs via a carrier-mediated transport system, and several inhibitors of uptake can antagonise the toxic effects of microcystins. The most effective of these is the antibiotic rifampin, which protects mice and rats against microcystin-induced lethality when given prophylactically and, in some cases, therapeutically (11). More recent experimental evidence shows that microcystins may also act as liver tumor promoters in extremely small amounts. A tumor promoter does not initiate cancer formation but helps a previously developed cancer to survive. Poisoning symptoms may take 30 minutes to 24 hours to appear, depending upon the size of the animal affected and the amount of toxic bloom consumed. Microcystin toxicity may include jaundice, shock, abdominal pain, weakness, nausea and vomiting, rapid and weak pulse and death (3).

Conclusion


References

