CHAPTER 5

Efficacy and Safety of the Anthrax Vaccine

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Introduction

The ongoing debate over the safety and efficacy of the anthrax vaccine is extremely complex. It is possible, however, to categorize the issues and concerns with Anthrax Vaccine Immunization Program as either clinically related or administrative policy related, then address the two categories separately.\(^1\) An important aspect of the clinically related issues is to determine if the anthrax vaccine, Anthrax Vaccine Adsorbed (AVA), hereafter to be referred to as “the vaccine,” is safe and provides effective protection against the effects of exposure to anthrax spores. This review is intended as a clinical assessment based on data in the published, peer-reviewed medical literature and medical textbooks.\(^2\)

If medical personnel determine the vaccine is clinically safe and effective, then senior level policy makers can make a policy decision to vaccinate the Defense Department personnel based on intelligence estimates and relative risk assessments related to the potential use of anthrax spores as a biological weapon. Clinicians and service medical corps officers do not set policy, nor do they have the authority to order vaccination of all personnel.

Military commanders and supervisors should have pertinent clinical facts and information about anthrax and the vaccine, written in lay terms, to serve as a working reference for use to educate those within their chain of command. This includes having an analysis of the major objections opponents to vaccination raise. The research methods employed for this chapter include a review of the peer-reviewed medical literature, medical textbooks, press releases, and internet world-wide-web sites presenting
information and opinions both for and against vaccination.\textsuperscript{3} Due to time and space limitations, this chapter is not intended to be an exhaustive review on the use of anthrax as a biological weapon.\textsuperscript{4} More detailed reviews and discussions of the evidence related to the risk of the use of anthrax as a biological weapon and the policy decision to vaccinate Defense Department personnel are presented in the chapter by Davis and Winegar in this book and several other sources.\textsuperscript{5}

**Pathophysiology of Human Anthrax Infections**

The organism, *Bacillus anthracis*, exists in the soil in dormant spores and can be found throughout the world.\textsuperscript{5} The spores are able to exist in the soil for years under the right conditions, such as cool, dry climates with adequate protection from sunlight.\textsuperscript{7} Grazing animals consume the spores which germinate and multiply inside the animal, causing disease, eventually leading to death. After the animal’s body decomposes, and the anthrax bacteria is exposed to the air. Oxygen in the air stimulates the bacteria to sporulate and new spores are then released into the environment, either being deposited into the soil or spread by birds and insects.\textsuperscript{8}

The word “anthrax” comes from the Greek word *anthrakis*, meaning “coal,” refers to the coal-black skin lesions caused when anthrax bacilli infect the skin.\textsuperscript{9} Incidents of anthrax infections of both humans and beasts have occurred throughout history. The fifth plague against Egypt, recorded in the Book of Exodus, may have been an outbreak of anthrax.\textsuperscript{10} Ancient Greeks, Romans, and Hindus also describe diseases associated with anthrax infection of humans.\textsuperscript{11}

Certain groups of people such as veterinarians and workers in the goat-hair or wool industries have been identified as having a higher risk of contracting anthrax. During the 1800s, anthrax was a significant agricultural and industrial problem.\textsuperscript{12} Indeed, another name for inhalation anthrax is “woolsorters’ disease.”\textsuperscript{13} Exposure to anthrax spores in the work place is effectively controlled through animal vaccination programs, good animal husbandry, vaccinating workers at risk for exposure, and improving working conditions. Such efforts have virtually eliminated
anthrax as an occupational hazard in the United Kingdom since 1940.\textsuperscript{14} In the United States, human anthrax is extremely rare with only 224 cases reported over 50 years.\textsuperscript{15} There has never been a case of human-to-human transmission of anthrax reported, leading most to conclude that anthrax is not contagious.\textsuperscript{16}

In humans, the anthrax bacillus causes three types of infections: cutaneous, inhalation, and gastrointestinal. Ninety-five percent of human anthrax infections are cutaneous. Spores enter through a break in the skin and germinate to form anthrax bacilli, leading to a localized infection. A vesicle then forms and ruptures to produce the characteristic coal-black lesion. Cutaneous anthrax is easily treated with antibiotics and the lesions heal without scarring. Most patients survive and develop an immunity against anthrax.\textsuperscript{17} If left untreated, the mortality rate is between 10 and 20 percent.\textsuperscript{18}

Inhalation anthrax occurs in five percent of human anthrax infections and is caused when spores enter through the lungs, lodging in the alveoli, the microscopic air sacs where oxygen exchange with the blood occurs. The anthrax spores may reside in the lung alveoli for several weeks before germinating.\textsuperscript{19} Macrophages, cells designed to consume foreign bacteria as part of the body’s immune system, engulf the spores and then migrate from the lungs to lymph nodes in the chest. Inside the macrophages, the spores germinate, growing into mature anthrax bacilli. The bacilli multiply and eventually erupt from the macrophages, spreading throughout the bloodstream.

The initial symptoms of inhalation anthrax signal germination of the spores into mature bacilli and are similar to any common upper respiratory tract infection. Since the symptoms are so non-specific, diagnosis at this point is not possible unless there is reason clinically to suspect anthrax exposure. After a few days, the symptoms subside for a brief period, typically 12 to 24 hours. This latent period is followed by an explosive period of severe symptoms, shock, and cardiovascular collapse, leading rapidly to death. During this final phase, massive numbers of anthrax bacilli circulate in the blood throughout the body, releasing deadly toxins.

Once initial symptoms develop, nearly 100 percent of all cases of inhalation anthrax are fatal (usually within three days), even with aggressive treatment using antibiotics and supportive intensive medical care. Therefore, if a potential exposure to inhalation anthrax is suspected,
treatment must be initiated immediately before any symptoms occur. Treatment should be continued either until the possibility of anthrax exposure is excluded or no more dormant spores are left in the lungs (believed to be approximately 60 days). To develop inhalation anthrax, the subject must inhale a minimum number of spores. The number of spores required to kill at least 50 percent of subjects has been reported to be 8,000 to 10,000 but may range from as few as 2,500 to 55,000. The late 2001 anthrax attacks in the United States, using the United States Postal Service, and the subsequent analysis of what causes a lethal dose shows that the lethal dose varies from person to person and that original estimates may have been based on insufficient data. Occupational studies of unvaccinated goat-hair and wool workers demonstrated they inhaled over 500 anthrax spores each day but they did not develop inhalation anthrax. Prior to September 11, 2001, there had been no cases of inhalation anthrax reported in the United States since 1978, and only 18 cases in the previous 80 years. Since September 11th and until January 2002, there were 5 reported deaths, and 22 total cases of persons harmed by the mail-delivered anthrax attacks.

Without deliberate aerosolization (such as during attack with a biological weapon), it is extremely rare for there to be a sufficient concentration of spores in the inhaled air to cause disease, even if there are large amounts of spores deposited on surfaces or in the soil. Studies indicate that secondary aerosolization typically will not stir up enough spores from contaminated soil or surfaces to achieve sufficient concentrations in inhaled air to cause disease. Therefore, decontamination of large areas and soil is usually not indicated and the presence of residual anthrax spores may not necessarily hinder military operations, as some imply.

Gastrointestinal anthrax results from consuming animal products or meat contaminated with anthrax spores. The initial infection occurs either in the mouth and throat or in the intestines. As in inhalation anthrax, macrophages engulf the spores which germinate, forming bacteria that enter the blood stream. The bacteria multiply and release toxins, leading to death in 50 percent of cases. Gastrointestinal anthrax is the rarest form of anthrax infection and has not been reported in the United States.

Rarely, anthrax may also infect the central nervous system, causing hemorrhagic meningitis. This form of anthrax infection does not represent a separate way for anthrax to infect humans. It is actually a
complication of cutaneous anthrax, caused by anthrax bacilli spreading through the blood or lymphatic systems to infect the brain and spinal cord.\textsuperscript{28} This complication is not frequently seen with inhalation or gastrointestinal anthrax, probably because patients die before meningeal infection by anthrax bacilli occurs. Meningeal anthrax is almost always fatal.

\section*{Anthrax Vaccine Adsorbed}

The human body normally fights infection two ways: by producing antibodies that circulate in the blood which recognize and attach to foreign proteins, called antigens, and by special cells (such as macrophages) that engulf (called phagocytosis) the bacteria to kill and digest them. Usually these two processes work together. Antibodies bind to antigens on invading bacteria to mark the bacteria. This attracts macrophages to the bacteria so the macrophages may phagocytize them. Antibodies also bind to circulating antigens, produced and released by bacteria into the blood stream, to neutralize their effect.

Anthrax bacilli that are able to cause disease inhibit both parts of the immune process. First, they produce protective antigen and two toxins called edema factor and lethal factor. The toxins couple with protective antigen and penetrate into the patient’s cells where antibodies in the blood cannot get to them to neutralize their toxic effect. The protective capsule formed by anthrax bacilli in the blood inhibits phagocytosis. As a result, the body’s defenses are rendered ineffective. Understanding these basic concepts are important in order to understand the strategy of treatment regimens and vaccination programs.

Anthrax Vaccine Adsorbed is the Food and Drug Administration (FDA) approved and licensed vaccine for use to immunize humans against anthrax infection. The strain of anthrax bacteria used to make the vaccine lacks the ability to make the protective capsule (cannot prevent the body’s defensive macrophages from phagocytizing the bacteria) and is unable to produce disease in humans. There are no live bacteria and no intact cells in the vaccine, so it is impossible to get infected with anthrax bacteria from the vaccine. The vaccine consists of protective antigen isolated from these attenuated (unable to produce disease) anthrax bacteria.\textsuperscript{29} Protective
antigen has been shown to be the essential antigen for provoking the immune response against anthrax in both animals and humans. Every type of anthrax vaccine developed that has been demonstrated as effective in immunizing test subjects against anthrax involves the use of protective antigen as the primary agent to trigger the immune response.\textsuperscript{30}

After injection, the vaccine stimulates the individual’s immune system to produce antibodies against protective antigen which protect the individual from future infections by anthrax bacilli. After vaccination, it takes the individual some time to develop enough immunity to confer protection and one dose may or may not be fully protective.\textsuperscript{31} Therefore, a non-immunized person exposed to aerosolized anthrax spores, in addition to immediate vaccination with the anthrax vaccine, requires treatment with antibiotics to prevent disease.

Formaldehyde (up to 0.02 percent) is used as a stabilizer in the vaccine and benzethonium chloride (0.0025 percent) as a preservative.\textsuperscript{32} The FDA has approved the use of formaldehyde in these trace amounts as a preservative.\textsuperscript{33} The use of formaldehyde as a preservative is actually quite common and has been done for the past 40 years. For example, tetanus toxoid, given to all school children in the United States, contains trace amounts of formaldehyde, yet it has been used safely for decades to induce immunity in millions of people by stimulating the production of antibodies against tetanus.\textsuperscript{34}

The anthrax vaccine, does not contain, nor has it ever contained, squalene as an additive. Squalene is a substance sometimes used to increase the potency of certain vaccines.\textsuperscript{35} Squalene occurs naturally in humans and is a precursor in the synthesis of cholesterol.\textsuperscript{36} Squalene is also found in large amounts in deep-sea shark liver. There are currently several health food supplemental products on the market containing squalene. Proponents claim squalene improves the delivery of oxygen to cells and facilitates the clearance of metabolic toxins.\textsuperscript{37}

Recent reports have stated that newly developed tests have detected trace amounts of squalene in Anthrax Vaccine Adsorbed and other commonly used vaccines.\textsuperscript{38} Previous tests were only able to detect the presence of squalene in parts per million, but the newer, more sensitive tests are able to measure the presence of squalene down to the parts per billion. The concentration of squalene detected in the anthrax vaccine, diphtheria vaccine, and tetanus toxoid, using the newer tests, is about ten
parts per billion. The normal concentration of squalene circulating in human blood is many times higher, about 250 parts per billion, suggesting the presence of trace amounts of squalene in the anthrax vaccine is not clinically significant. The presence of trace amounts of squalene in the anthrax vaccine and in the other vaccines may be a normal bi-product of the production process.  

There have been articles in the press attempting to draw a connection between the use of the vaccine, and Gulf War Syndrome, claiming the agent causing Gulf War Syndrome is squalene. These press reports claim veterans suffering from Gulf War Syndrome have antibodies to squalene in their blood which they got from the anthrax vaccine. Others have gone so far as to charge the Defense Department may have secretly added squalene to lots of the vaccine, used for inoculating troops to increase its efficacy. They claim, without presenting any evidence, that anthrax vaccine, vial labels may have been altered and that lack of documentation in personal shot records suggest a cover-up.

History of Production

Merck, Sharp & Dohme developed the first anthrax vaccine for use in humans during the 1950s to protect workers routinely exposed to anthrax spores. Clinical trials performed in the late 1950s and published in 1962 demonstrated that the vaccine was effective in preventing cutaneous anthrax. Later, the Department of Defense (DOD) approached the state of Michigan to manufacture anthrax vaccine for military personnel. DOD chose the state of Michigan because there was little profit potential to motivate private industry to manufacture a vaccine that would not be used in the general public, and Michigan had extensive experience manufacturing other vaccines such as rabies vaccine.

Therefore in 1970, Michigan Biological Products Institute (hereafter call “The Institute”) began to produce Anthrax Vaccine Adsorbed for DOD. This is essentially the same vaccine as initially produced by Merck, Sharp & Dohme, except that the current anthrax vaccine is more potent and more pure, due to some minor differences in production technique. In 1970, the National Institute of Health’s Division of Biologics Standards
licensed the vaccine, and then in 1972 transferred the license, along with oversight and regulatory authority, to the FDA.\textsuperscript{45} Licensing was based on data collected during studies using both the older anthrax vaccine and Anthrax Vaccine Adsorbed to protect workers at risk from infection. A 1962 study based on the older, less potent vaccine measured its effectiveness in protecting wool mill workers at risk for both cutaneous and inhalation anthrax. A later Centers for Disease Control and Prevention (CDC) study of the current vaccine, conducted for over a decade, showed how effectively it prevented cutaneous anthrax in workers at risk from infection. Of note, it was demonstrated that there was a low risk of serious side effects.\textsuperscript{46} As tensions in the Persian Gulf mounted in early 1990, the U.S. Defense Department asked Michigan Biological Products Institute to dramatically increase the production rate of the vaccine.\textsuperscript{47} The Institute informed DOD it would not be able to meet production expectations with the facilities it possessed at that time. The Institute then worked out a plan to upgrade its production facilities with DOD funding and presented the plan to the FDA in 1995, and the FDA approved the facility upgrade plans. Between 1995 and 1997, the FDA performed several inspections of the Institute’s facilities used to produce rabies vaccine and plasma derivative products. During these inspections, the FDA found numerous discrepancies with policies and procedures, record keeping, analytical laboratories, quality control practices, raw materials handling, filling and packaging, and storage, warehousing, and distribution.\textsuperscript{48} It must be noted that none of these production facilities, nor any of the FDA’s findings, involved the production, safety, or quality of vaccine. In March 1997, the FDA sent the Institute a letter indicating the FDA would begin procedures to revoke the Institute’s license due to lack of adequate progress to address the discrepancies noted during the inspections of the facilities used to produce rabies vaccine and plasma derivative products, unrelated to the production of anthrax vaccine.

In the mean time, Michigan Biological Products Institute had applied to the Food and Drug Administration to upgrade its anthrax vaccine production facilities to meet the increased demand resulting from the DOD anthrax vaccine immunization program. The Food and Drug Administration approved the planned upgrade and, in January 1998, the Institute voluntarily stopped production of the vaccine in order to begin the
FDA-approved renovations to the vaccine production facilities. It is important to note that the stoppage of production of the vaccine is completely unrelated to the discrepancies noted during FDA inspections of rabies vaccine and plasma derivative products production facilities and is completely unrelated to the FDA’s letter of intent to revoke the Michigan Biological Products Institute’s license.

In 1997, Secretary of Defense Cohen made the decision to implement the anthrax vaccine immunization program to vaccinate all military personnel using lots of vaccine already on hand. Since supplies of the vaccine were limited, the immunization program was divided into three phases. Completion depended on the production and release to DOD of additional lots of vaccine after the production facilities were upgraded. As of early 2002, only the first phase has been implemented, meaning only personnel at risk for exposure to inhalation anthrax in high risk areas (i.e., Korea and the Persian Gulf) will be vaccinated.

Part of the Secretary’s directive was that each lot of the vaccine would be completely re-tested using FDA testing procedures to reconfirm potency, safety, purity, and sterility. Each lot had to pass such supplemental testing before it would be administered to Defense Department personnel. The lots undergoing supplemental testing had already passed FDA certification, been released by the FDA for sale, and been purchased by the Department of Defense.

Eight lots underwent supplemental testing for potency and were released before a problem with the potency test itself was discovered in the Fall of 1998. Since then, the potency testing difficulties have been corrected and the test is now working according to specifications. However, the FDA will not release any additional lots until it is satisfied with the quality of the vaccine and has approved necessary potency test amendments implemented to correct the earlier potency testing problems.

In September 1998, the state of Michigan sold the Michigan Biological Products Institute facilities with the vaccine licensing rights to BioPort as part of an effort to privatize government programs and cut costs. In late 1999, BioPort completed the renovations and applied to the FDA for inspection and certification of the new production facilities. As of early 2002 BioPort continues to have problems with its renovated facility and still has not received FDA certification.
In the meantime, BioPort has started producing new lots of anthrax vaccine.\textsuperscript{55} It is important to note that the new lots produced by BioPort have not been certified or released for sale by the FDA, have not been purchased by DOD, and have not been administered to anyone. Furthermore, the Defense Department will not purchase these lots to begin phase two of the immunization program until the BioPort facility passes the FDA inspection and the FDA has tested, certified, and released the new lots for distribution.\textsuperscript{56}

BioPort has a total of 32 lots of vaccine in storage for DOD produced before the production facilities were shut down for renovations. In February 1998, the FDA inspected these lots of vaccine, and the Institute voluntarily quarantined ten lots. These will remain quarantined until testing confirms adequate sterility, potency, and quality to the FDA’s satisfaction.\textsuperscript{57} In addition, one other lot was permanently quarantined due to questions regarding sterility and will not be used. Another, 14 lots were tested at random and found not to contain any squalene.\textsuperscript{58}

Most recently, the rate of vaccination of military personnel via the immunization program has been reduced due to the dwindling supply of vaccine. As of May 2000, 17 lots of anthrax vaccine, all produced by Michigan Biological Products Institute before the renovations began, have passed Food and Drug Administration certification tests and passed re-certification tests as ordered by the Secretary of Defense.\textsuperscript{59} Defense officials have pointed out on numerous occasions in the media and in sworn testimony that only these 17 lots have been used for the immunization program.

Defense officials had hoped that BioPort would have obtained FDA approval quickly and new lots tested and released by now.\textsuperscript{60} The FDA has identified 30 deficiencies that need to be rectified before it grants certification of the vaccine. No new lots produced by BioPort are available for the immunization program, forcing a slowdown in the pace of the program. In addition, some members of Congress who are dissatisfied with BioPort’s situation are beginning to urge DOD to consider designing a government-owned, contractor-operated (termed “GOCO”) vaccine production facility.\textsuperscript{61}

In summary, it should be noted that the current vaccine is a FDA licensed, non-experimental vaccine. It is more potent and more pure than, but otherwise identical to, the earlier version of the vaccine produced in the 1950s by Merck, Sharp & Dohme. The Institute voluntarily stopped production of the vaccine in order to upgrade production facilities, not due
to the results of any FDA inspections. The Institute then sold its anthrax vaccine production facilities to BioPort.

The lots of vaccine currently in use for the immunization program were produced before building renovations began, were tested and certified by the FDA for release and distribution, and re-tested by DOD before administration. No lots produced by BioPort have been used yet by DOD to vaccinate its personnel. DOD-mandated supplemental testing, BioPort’s voluntary quarantine of lots previously released by the Food and Drug Administration, and the modifications implemented to improve the quality of testing for potency demonstrate the intense level of interagency scrutiny that exists to ensure that DOD’s immunization program attains the highest possible levels of safety for DOD personnel.

**Efficacy**

Brachman, et al. published a controlled study using the original anthrax vaccine produced by Merck, Sharp & Dohme and supplied by the U.S. Army Chemical Corps in 1962.62 The study looked at how effectively the vaccine prevented anthrax in a population of wool mill workers considered to be at risk for contracting anthrax. Historically, about one percent of these workers contracted cutaneous anthrax annually. To do the study, volunteers were divided into two groups -- one group received the vaccine and the other received a placebo (an inactive substance used as a control that looks like the vaccine but is harmless and has no biological effect). The vaccination schedule used in the study matches the current Food and Drug Administration-approved schedule for vaccinations using Anthrax Vaccine Adsorbed. The rate of occurrence of anthrax in the vaccinated group was compared to the rate of occurrence in the group that received the placebo and all other workers not participating in the study.

During the study period there were 26 cases of anthrax. One case of cutaneous anthrax appeared in a fully vaccinated individual. Twenty-three cases of anthrax appeared in unvaccinated workers and two in partially vaccinated (meaning they did not complete the series of immunizations) workers. No cases of inhalation anthrax occurred in vaccinated or partially
vaccinated workers although five cases of inhalation anthrax occurred in unvaccinated workers during the study period. Four of these cases were fatal. The frequency of occurrence of inhalation anthrax was not sufficient to determine any statistical significance for how effective the vaccine was in preventing inhalation anthrax.63

As already pointed out, the vaccine used in the Brachman study was also a protective antigen vaccine similar to Anthrax Vaccine Adsorbed but less potent and less pure since it contained more cell fragments. Since the mechanism to produce immunity is the same for both vaccines, Brachman’s study results are relevant when discussing the issue of efficacy of the vaccine. In addition, other surveillance studies using the vaccine, completed since the publication of Brachman’s study, confirm Anthrax Vaccine Adsorbed’s efficacy in preventing anthrax in humans.64

Between 1962 and 1974, the Centers for Disease Control and Prevention (CDC) collected data measuring the occurrence of anthrax in workers at risk for infection who had been vaccinated with Anthrax Vaccine Adsorbed versus non-immunized workers. The study also tracked any adverse reactions to the vaccine.65 During this period, an additional 27 cases of cutaneous anthrax were identified, three in partially immunized workers who had only received one or two doses. There were no cases of anthrax in the fully immunized workers.66 A total of 7,000 workers received more than 16,000 doses of the vaccine.67 The efficacy data from the Brachman study, using the original protective antigen vaccine, and the CDC study, using the vaccine, were eventually used during the licensing procedures for it.68

Between 1974 and 1989, it is estimated that an additional 68,000 doses of Anthrax Vaccine Adsorbed were administered to at risk individuals.69 There were no cases of cutaneous anthrax in vaccinated individuals although there continued to be reported cases of cutaneous anthrax in unvaccinated people at risk. In addition, the rate of adverse side effects remained low, comparable to rates cited in the FDA-required package insert that accompanies each vial of the vaccine.70 Due to the increasing rarity of anthrax infections, the fact that workers at risk for exposure to anthrax spores are immunized, and improvements in working conditions, any additional field studies of anthrax vaccine are unlikely.71 In conclusion, the clinical data collected over several decades indicate that
the vaccine is very effective in preventing cutaneous anthrax and, potentially, inhalation anthrax in humans.\textsuperscript{72}

Fortunately, inhalation anthrax in humans is very rare even among unvaccinated workers routinely exposed to anthrax spores. Improvements in the work place plus use of the vaccine in workers at risk for exposure to anthrax spores have essentially eliminated the occurrence of inhalation anthrax.\textsuperscript{73} But, the rareness of this disease also means it is not possible to collect enough data in humans to determine if the vaccine would prevent inhalation anthrax in humans. In order to do a study in humans, one would have to take volunteers, divide them into two groups, vaccinate one group with the vaccine, the other with a placebo, then expose both groups to lethal doses of aerosolized anthrax spores, and track how many in each group contract the disease. Obviously, such a study would be unethical.

Numerous animal studies have been performed to measure the effectiveness of Anthrax Vaccine Adsorbed to prevent inhalation anthrax. Granted, there is always a possibility that results in one species of animals cannot be assumed to represent potential results in another species. For example, animal studies suggest that some species are more difficult to immunize against anthrax infections, using Anthrax Vaccine Adsorbed, than others. In guinea pigs, the vaccine seems to confer variable protection against certain strains of anthrax, suggesting possible species-dependent differences in the guinea pig’s immune system. Guinea pigs seem especially sensitive to one particular strain of anthrax, called the Ames strain, even after they are fully immunized with the vaccine.

On the other hand, Anthrax Vaccine Adsorbed confers excellent protection in rabbits and non-human primates against the Ames strain, providing near 100 percent protection even after as few as two inoculations, including situations where they are exposed to several times the lethal dose of anthrax spores. Moreover, inhalation anthrax infections in non-human primates closely resemble inhalation anthrax infections in humans.\textsuperscript{74} Based on the animal studies results and the absence of cutaneous and inhalation anthrax in fully immunized individuals exposed to anthrax spores, it is reasonable to conclude that the vaccine prevents inhalation anthrax in humans.\textsuperscript{75}
Safety and Side Effects

The side effects and adverse reactions recognized as caused by the vaccine tend to be grouped into four main categories: mild local reactions, moderate local reactions, severe local reactions, and systemic reactions. Mild local reactions are defined by tenderness and redness in an area less than 1 to 2 cm in diameter and occur about 30 percent of the time. Moderate local reactions are identified by an area of response greater than 5 cm in diameter and occur about 4 percent of the time. Severe local reactions are characterized by extensive swelling (edema) of the arm and forearm in which the vaccine was administered. These occur less frequently than moderate reactions. In general, the rate of local reactions is about twice as high in women than men. Systemic reactions are characterized by fever, chills, nausea, and body aches and occur in less than 0.2 percent of vaccinations. Allergic reactions are even less common, being reported in only one per 100,000 doses.

Normally it takes three doses of the vaccine before an individual begins to develop an immune response and seem to correlate with the observation that reactions to subsequent doses tend to be stronger. Individuals who have had cutaneous anthrax or who have severe local or systemic reactions to the vaccine are not to receive the vaccine. In a study conducted from 1962 to 1974, the CDC tracked the occurrence rates of reactions during the administration of more than 16,000 doses to over 7,000 individuals. The results of this study are the rates reported on the informational package insert accompanying each vial as required by the FDA.

Since Anthrax Vaccine Adsorbed was licensed in 1970, there have been numerous reviews documenting the occurrence of side effects attributable to it. An independent civilian advisory panel met in 1985 to review the results of the 1962 to 1974 CDC study. The panel reported that only a few systemic side effects had occurred of which all resolved. Local reactions were typically mild and also resolved. From 1974 to 1989, over 68,000 doses of vaccine were administered to persons.
considered at risk for contracting anthrax. This would include goat hair workers, laboratory personnel, livestock handlers, and veterinarians. Yet, after more than 30 years of use, no long-term side effects have been reported in association with this vaccine.84

Since 1973, the U.S. Army Medical Research Institute of Infectious Diseases (USAMRIID) at Fort Detrick, Maryland, has actively followed 1,590 workers who have received more than 10,000 doses of Anthrax Vaccine Adsorbed, again with no reported long-term or chronic side effects. Only 4 percent reported local reactions and only 0.5 percent had any type of systemic reactions. All reactions resolved without any lost work time.85 Another study conducted by the Canadian Armed Forces reported that in 547 individuals who received the vaccine, rates of reaction were less than the rates listed on the package insert. There were no long-term effects except for one individual who reported a persistent nodule at the injection site.86

In addition to the CDC study, the USAMRIID study, and the Canadian study, there are three other separate studies on Anthrax Vaccine Adsorbed, examining the rate of occurrence of adverse reactions. In 1997, the Pittman study reported on 508 subjects who were actively followed after they received the vaccine. Local reaction rates were roughly the same as reported by other studies, but Pittman noted a much higher rate of systemic reactions. Twenty-nine percent were classified as mild and 14 percent were classified as moderate to severe. Another study, conducted at Tripler Army Medical Center in Hawaii, reported a rate of mild systemic effects of 43 percent and moderate to severe in 5 percent out of a total of 536 individuals vaccinated. Both studies are significant in that they report moderate to severe systemic reactions much higher than the 0.05 percent to 0.2 percent usually reported, and they differentiate between mild and moderate to severe systemic reactions.87

The third study is an ongoing Department of Defense study which reported in May of 1999 that out of 223,000 individuals vaccinated, 42 experienced adverse side effects which were reported to the FDA and CDC. Of these, seven either missed more than one day of work or required hospitalization. None of these studies note any long-term or chronic adverse effects attributable to the vaccine and none question the safety of it in their conclusions.88 In addition, there have been no cases of
anaphylactic reactions (severe, potentially life threatening, systemic allergic reactions) reported due to its administration.\(^89\)

There have been multiple review panels, including panels hosted by the FDA, the CDC, the World Health Organization, and the Armed Forces Epidemiological Board. Most recently, a civilian panel of 21 experts from several major medical and research centers led by Dr. Thomas V. Inglesby convened to assess the risk that anthrax could be used as a biological weapon agent. The panel also developed a consensus on the care and management of victims of an anthrax biological weapon attack and examined the safety and efficacy of the vaccine. The panel’s results were published in May 1999 in the *Journal of the American Medical Association*.\(^90\)

The panel concluded that the likelihood that anthrax could be used in a terrorist attack is high. The panel also reported that its investigation of the clinical data on the use of Anthrax Vaccine Adsorbed showed no serious adverse effects have been causally related to Anthrax Vaccine Adsorbed, and it reached a consensus for recommending treatment protocols to care for anthrax victims. The panel also recommended that research should be devoted to developing a next-generation anthrax vaccine which requires fewer doses to immunize humans. The panel’s findings correlate with the findings of numerous other review panels examining the medical literature published on the vaccine which confirm the clinical safety and its efficacy in humans.\(^91\)

In 1990, the FDA and CDC launched the Vaccine Adverse Events Reporting System (hereafter termed “the reporting system”). This is a passive reporting system, meaning success depends on medical personnel, patients, and families taking the initiative to file reports. As of 23 August 2000, 1,859,666 doses of the vaccine had been administered to 463,027 personnel with 945 reports submitted to the reporting system. Of these reports, 492 were determined to be actually due to the vaccine--374 were less than serious, 111 reported a loss of more than 24 hours of duty, and 7 were hospitalized for allergic inflammatory reactions at the injection site. All symptoms resolved and there were no permanent side effects.\(^92\)

In addition to the FDA reviews of the reporting system data, DOD convened the Anthrax Vaccine Executive Committee composed of non-government medical experts. This group meets periodically to review the reports from the Vaccine Adverse Events Reporting System.\(^93\) Since its
first meeting in 1990, the committee has not identified any unexpected patterns of adverse events among the reports submitted to the reporting system.\(^94\) The committee continues to meet every six weeks to review data reported on the vaccine.\(^95\)

To date, the Anthrax Vaccine Executive Committee has concluded it is not possible to attribute to the vaccine, all the symptoms reported to the reporting system. But, for the sake of argument, if one assumes that all the reports could be linked causally to the vaccine, the rate of adverse reactions, including serious or severe ones, is still less than 0.03 percent. This is below the rate of 0.05 percent reported by other studies and well below the rate of 0.2 percent listed in the vaccine product information package insert. By way of comparison, the hepatitis B vaccine, required for all health care workers, has a systemic reaction rate five times greater than that observed due to the anthrax vaccine.\(^96\)

Based on reporting System data, the FDA has concluded that it has no concerns about the safety of the vaccine and “continues to view it as safe and effective for individuals at risk of exposure to anthrax.”\(^97\)

In all there have been at least 13 studies conducted in humans assessing the safety of the present anthrax vaccine or its precursor protective antigen vaccine, including those discussed in this paper, covering almost 50 years of clinical experience.\(^98\) The clinical evidence accumulated is consistent from study to study and demonstrates that the vaccine is safe and effective. Anthrax Vaccine Adsorbed quite possibly has undergone more scrutiny that any other vaccine developed for human use, yet it continues to find endorsement in medical textbooks, in the medical peer-reviewed literature, and in sworn testimonies given before Congressional panels as a safe and reliable vaccine against human anthrax infections.\(^99\)

**Understanding the Arguments Against Anthrax Vaccine Adsorbed**

It is possible to group the concerns over the vaccine as follows: concerns over the vaccine’s safety and efficacy; concerns regarding proper indications for use; concerns over its manufacture and ingredients; concerns
regarding the lack of FDA certification of BioPort’s renovated manufacturing facility; concerns regarding licensure (including whether or not the vaccine is investigational in nature); concerns over lack of published peer-reviewed clinical trials; concerns over the development of strains of anthrax that are resistant to vaccination; the apparent high rate of occurrence of symptoms in vaccinated individuals at Dover Air Force Base, Delaware; concerns over whether the threat justifies the use of the vaccine since weaponized anthrax has not yet been used against our armed forces. Most of the evidence addressing these concerns has already been presented.

As already discussed, licensure of the anthrax vaccine and the information for the package insert was based on both the Brachman study and data collected by the CDC over a ten-year period on the use of the vaccine in workers considered at risk for exposure to anthrax spores. Furthermore, the Brachman study is not the only place where data on the safety and efficacy of the vaccine has been published. As already noted in this paper, there have been numerous other studies on the safety and efficacy of the vaccine conducted over several years involving tens of thousands of human subjects. Although not all of these studies are individually published as peer-reviewed articles, the data collected by these studies has been examined by review panels and published in several articles that have undergone the peer-review process. Therefore, stating that the Brachman study contains the “published adverse reaction rates” without acknowledging these other sources of data in the peer-reviewed literature is misleading.

Another concern raised by opponents to the immunization program is that adversaries might develop strains of anthrax that are resistant to the vaccine. Some base this on the fact that strains of anthrax have been developed that are resistant to antibiotics. Also, there have been reports that anthrax strains have been developed that may render the Russian-developed live attenuated vaccine ineffective. Neither of these reports mean that a strain of anthrax has been produced that is resistant to the present U.S. vaccine.

First, it must be pointed out that developing resistance to antibiotics is not the same as developing resistance to vaccines. Antibiotics (biochemicals produced in nature or synthesized in laboratories that are toxic to bacteria) are completely different from antibodies (complex proteins produced by the inoculated individual’s immune cells) that result
from vaccination. Bacteria commonly develop resistance to antibiotics through several naturally occurring mechanisms, resulting in the antibiotic (such as penicillin or tetracycline) no longer being toxic to the bacteria.

Anthrax Vaccine Adsorbed, however, induces the inoculated individual to produce antibodies against protective antigen, which also is a protein. In order for anthrax to develop a resistance to the vaccine, the bacteria’s genetic code for protective antigen would have to be altered in such a way so the bacteria produces an altered version of protective antigen that the antibodies cannot recognize, but the protective antigen would still have to retain its functional ability to combine with the host’s cells and the other anthrax toxins (which are also proteins made by anthrax bacteria) to produce disease.

An adversary intent on producing a strain of anthrax resistant to Anthrax Vaccine Adsorbed would, therefore, need to possess highly sophisticated and very expensive genetic engineering capabilities. Needless to say, any genetics program intended to alter anthrax to change the characteristics of protective antigen would be a monumental undertaking and well beyond the reach of most potential adversaries. Not surprisingly, there is no documentation that a strain of anthrax consistently resistant to the vaccine in all species has been produced.\textsuperscript{105}

At Dover Air Force Base, Delaware, the number of individuals reporting adverse reactions after inoculation with the anthrax vaccine appears to exceed the rate one would expect based on the published literature. A list of many of the symptoms reported can be found on the world-wide-web.\textsuperscript{106} There are several problems, however, trying to make a connection between these symptoms and the vaccine. First of all, there is no discernible pattern to the symptoms. The time of onset between vaccination and the onset of symptoms is highly variable, ranging from a few hours to months. The listings on the web site do not indicate if these patients got better except in one or two cases.

From a statistical perspective, after almost 40 years of clinical experience with Anthrax Vaccine Adsorbed, plus several studies documenting its safety, why would there be this sudden cluster of cases at Dover? By way of contrast, the U.S. Army Medical Research Institute of Infectious Diseases tracked 1,590 individuals who received 10,451 doses of the vaccine, over several years, documenting rates of adverse events no higher than those listed in the FDA package insert and no loss of duty.\textsuperscript{107} With no
recurrent pattern of symptoms, and no consistent temporal relation of the development of symptoms to inoculation with the vaccine it is extremely difficult to claim the cases at Dover prove the vaccine is the cause. In addition, the rate of occurrence of any disease (for example thyroid disease) in vaccinated personnel at Dover is equal to or less than the rate of occurrence of the same disease in unvaccinated individuals, and the rate of occurrence of individual symptoms in personnel vaccinated is no higher than the rate expected when vaccinating personnel with any other vaccine, further complicating claims that the anthrax vaccine caused the symptoms.

Without a doubt it would be wrong to trivialize the symptoms these patients are experiencing. The symptoms are very real and must be addressed in a compassionate, professional manner. But, the fact that these individuals are having symptoms and the fact that they received shots does not prove that the vaccine caused the symptoms. In contrast, it is more likely these individuals would have developed the symptoms from which they currently suffer even if they had not received the vaccine.

One writer goes so far as to suggest there should not be any vaccinations of Defense Department personnel until an anthrax-based biological weapon is actually used even though he acknowledges that historical precedence exists to justify concern that anthrax could be used as a political tool. This approach ignores the ready availability of anthrax spores and the potential for weaponization. There are limitations associated with constant use of personal protective gear, with gathering intelligence to provide advanced warning of an attack, and no means exists to detect reliably that an attack with anthrax is occurring. Considering that international conventions historically have failed to prevent proliferation of biological weapons and that it takes time for an individual to develop an immunity against anthrax after vaccination, it becomes apparent that waiting until an attack is imminent before immunizing personnel would not only be ineffective but dangerous. Consequently, immunization against anthrax before an attack becomes imminent is still our best proactive defense to protect personnel from attacks using anthrax-based biological weapons.

The debate over the anthrax vaccine has led to introduction of a bill in Congress that would suspend the DOD immunization program. Much of the language of the bill cites language similar to the language found in the
Hersack

media and on the internet. The bill would also prohibit the gathering of any data whatsoever on adverse effects potentially related to the vaccine.

This prohibition is indeed unfortunate for three reasons. First, there is nothing unethical about collecting data while the immunization program is in effect. Second, DOD would not be able to collect the vast amount of valuable data that could be used to resolve the issues and concerns that led to the introduction of the bill. Third, the bill ignores the relative risks of not vaccinating DOD personnel (including the real risk that military personnel could be attacked with an anthrax weapon and the lethality of inhalation anthrax) versus the large amount of clinical data documenting the vaccine’s safety and efficacy.

There are also allegations that the Defense Department is not doing enough to document the occurrence of reactions or side effects due to vaccination. These allegations are highly critical of the effectiveness and accuracy of the Vaccine Adverse Events Reporting System (hereafter, called the “reporting system”) data claiming that the rate of reporting to the reporting system by DOD health care workers is low. Some allege health care workers and physicians were ordered not to report any but the most severe reactions and not to report any symptoms or reactions not specifically listed in the FDA package insert for the vaccine. There is no source or substantiating documentation given for this allegation.

The purpose of the reporting system is to gather ongoing, long-term data on potential adverse reactions due to vaccines that were not identified during limited clinical trials. For example, if the incidence of a particular reaction or severe systemic effect occurring is one in a million, then several hundred thousand or even several million doses may have to be administered before that reaction would be observed. In reality, such extensive, long-term studies are not possible during clinical trials. Furthermore, if evidence exists that the vaccine will prevent more disease and save more lives than any harm caused by the vaccine, it may be regarded as unethical to withhold the vaccine from market to conduct long-term studies. The FDA instituted its reporting system to continue to collect data over the long-term, after vaccines are released for sale, to look for extremely rare adverse effects even though initial studies indicate a vaccine is safe and effective.

The Vaccine Adverse Events Reporting System is a passive reporting system, meaning that individuals must take the initiative to file a report.113
There is no one that actively calls or surveys vaccinated individuals to see if they developed any symptoms. In some cases this could be a disadvantage, leading to low reporting rates. Also, it is not possible just by using the reporting system data to establish that a particular vaccine actually caused an event, but, through the identification of possible trends over the long-term, this data is useful to direct new clinical studies to establish causality.

Several facets have been built into the reporting system to facilitate gathering all the facts. For example, anyone, including patients and families, may report any symptom suspected to be due to a vaccine. In addition, medical personnel are routinely reminded through extensive educational programs about the reporting system and the need to report. Furthermore, medical personnel are required to report adverse effects due to vaccines to the manufacturer who are then required to report those.\textsuperscript{114}

The Vaccine Adverse Events Reporting System receives over 12,000 reports of possible adverse reactions to vaccines each year. Fifteen percent are considered serious, including those that are life-threatening, result in hospitalization, missed work, or permanent disability. It should be noted that for some childhood vaccines, more reports of potential adverse effects from the vaccine are filed each year than the number of reported cases of the disease the vaccine is designed to prevent.\textsuperscript{115} This cumulative evidence suggests that, contrary to the criticisms of anthrax vaccine opponents, the reporting system is highly successful.

The Defense Department has reiterated to medical personnel that they should report any events they feel may be due to the anthrax vaccine to the reporting system. There is no documentation that the DOD instructed medical personnel to file a report only when they observe the side effects and reactions listed in the FDA package insert.\textsuperscript{116} Instead, DOD encourages all medical personnel to report all events potentially thought to be related to Anthrax Vaccine Adsorbed and requires them to report to the reporting system all adverse reactions potentially associated with the vaccine resulting in hospitalization or loss of more than 24 hours duty.\textsuperscript{117} Additionally, in 1999, the Air Force Surgeon General directed that any adverse events even suspected by medical personnel to be related to the vaccine will be reported to the reporting system.\textsuperscript{118} The allegations that DOD physicians are prohibited from filing reports to the reporting system
on any potential vaccine-related event are completely unfounded, ignoring the fact that such prohibitions may be illegal.

In spite of the extensive documentation of evidence addressing the safety and efficacy of Anthrax Vaccine Adsorbed in humans, concerns over its use continue to be propagated in the media. Many press release articles confuse facts, combine separate facts, or report facts in such a way as to be potentially misleading. For example, an extensive article published in the *Phoenix New Times* states that anthrax vaccine production has been halted due to problems with the new BioPort production facilities. Actually, the Michigan Biological Products Institute voluntarily halted production in order to renovate the facility, then later sold it to BioPort. After the sale, BioPort completed renovations but has had problems obtaining FDA certification of the renovated facility. While both facts are true, they are not directly, nor causally, related to each other as the *Phoenix New Times* article implies. This article also attempts to raise completely unsubstantiated concerns that there could be birth defects if a man who received the vaccine fathers a child.

Another example is an Associated Press article entitled “Food and Drug Administration inspection cites problems in vaccine production.” The article correctly states the problems are with FDA certification of the renovated facilities, required before new batches of the anthrax vaccine may be sold, but the last sentence of the article gives an unrelated fact that several anthrax vaccine lots failed FDA potency testing. The article does not clarify that these were older lots, none of which have been used by the Defense Department and are not related to the problems with FDA certification on BioPort’s more recently renovated production facilities. This could lead one not familiar with the facts to believe there is a direct relationship between the recent inspections, lots of the vaccine that failed potency testing, and the lots currently in use by DOD where, in fact, no such direct relationship exists.

Understanding the debate over the present vaccine, in large part involves understanding the internet’s effect on public opinion and the challenges the internet presents to those seeking a scientifically-rigorous opinion. The informational world-wide-web sites on anthrax have some potential value, but they can also be the source of significant confusion and misinformation. Some individuals on the Internet go so far as to raise questions regarding the legality of anthrax vaccine immunization program,
implying military personnel are duty bound to disobey it and that the DOD immunization program is a violation of their civil rights. It is important to note that many links found on internet web sites opposed to the immunization program connect to on-line source documents that do present the facts regarding Anthrax Vaccine Adsorbed. These facts are consistent with the clinical and historical evidence presented above and match the information presented on the DOD anthrax informational website. The difference is how the facts are interpreted and represented, often taking and quoting documents out of context, and inserting subjective opinions and editorial comments.

One of the most comprehensive world-wide-web sites opposing administration of the vaccine is entitled “Anthrax Vaccine Links and Information” and provides an extensive list of links to other related sites. Included are links to sites with copies of Congressional testimonies, Government Accounting Office reports, summaries of the symptoms reported by personnel at Dover Air Force Base who received anthrax shots, press releases, and other documents of interest.

To illustrate the importance of presentation, one title to a link claims the FDA admits it has never received data on the vaccine’s effects on long-term health, potentially leading some to believe the data does not exist. This link connects to a letter the FDA wrote to the Executive Director for Veterans for Integrity in Government. The letter responds to a series of questions, including whether or not any studies on the long-term health effects of the anthrax vaccine have been performed. It states the data has not been submitted to the FDA but adds the vaccine had (at the time of writing) been used for more than 28 years in veterinarians, laboratory personnel, industrial workers, and FDA inspectors. The clear intent of the answer is that the studies on the long-term health effects of the vaccine have been performed and the data does exist, but it has not been formally submitted to the FDA.

There is also confusion between present DOD vaccine and the British version of the vaccine. Another link on the “Anthrax Vaccine Links and Information” internet web site announces there have been British reports of outbreaks of Gulf War Syndrome after “recent” anthrax vaccinations. It references a British article entitled “Anti Bio-weapon Vaccine for troops Fails Safety Tests” from an independent British newspaper which reports newly produced lots of the British anthrax vaccine failed safety tests.
article cites concerns from British Persian Gulf War veterans that the British version of the anthrax vaccine may have caused Gulf War Syndrome and that further use of the vaccine may cause more to develop symptoms. They claim many fell ill after recent vaccinations, but the article provides no substantiating information. The article further alleges the lots used had expired, and the shelf life had been extended several times.\textsuperscript{126}

This link is misleading because the article is about the British version of anthrax vaccine. The British version of anthrax vaccine is not the U.S. vaccine. It is not produced either by Michigan Biological Products Institute or BioPort, does not require Food and Drug Administration licensure, and is not used in the United States. Furthermore, the FDA and British regulatory systems are completely separate. Yet the adversarial Internet site contains no statements to make this distinction.

Many who are opposed to the DOD immunization program have attempted to connect the vaccine with Gulf War Syndrome in spite of the fact that no such causal relationship has ever been demonstrated.\textsuperscript{127}

Other links seem intended to provoke an emotional response, such as one with photographs of injection sites with signs of local reactions entitled “Painful Anthrax injection site photos…OUCH!” Others refer to the numerous cases of individuals at Dover Air Force Base who claim to have developed symptoms after receiving the vaccine, claiming this proves the DOD really knows that the vaccine is not safe. There are also links to support groups and on-line chat rooms where those opposed to the immunization program may discuss their views or tell their story.

Interestingly, there are also links to other sites opposed to the use of other types of vaccines (e.g., hepatitis B vaccine) or all vaccines in general. This suggests that those opposed to the use of anthrax vaccine are part of a larger movement opposed to the use of all vaccines.

At the forefront of the opposition to the DOD’s anthrax vaccine program is an emergency room physician from Maine named Dr. Meryl Nass. She has written a number of articles and testified before Congress several times against the anthrax vaccine program and is regarded by opponents to the Anthrax Vaccine Immunization Program. For example, one link on the “Anthrax Vaccine Links and Information” site makes the claim that DOD officials really do know the anthrax vaccine is not safe. It turns out the link leads to an unpublished article written by Dr. Nass about an informational meeting for 100 physicians at Fort Detrick in May 1999
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where issues about the anthrax vaccine were discussed. Her article implies that military physicians asking policy questions about Anthrax Vaccine Adsorbed and Vaccine Adverse Events Reporting System proves the Defense Department does in fact know Anthrax Vaccine Adsorbed is not safe. She ends by admonishing the readers to contact their congressional representatives.

Dr. Nass also has her own informational world-wide-web home page about anthrax. The “Anthrax Vaccine Links and Information” site lists her credentials, which includes three years experience studying the anthrax outbreak in Zimbabwe. She is quoted as saying as many as ten percent of those receiving Anthrax Vaccine Adsorbed have gotten sick, although there is no explanation of what that means--whether the symptoms were mild, moderate, severe, localized, or systemic.

In 1999, Dr. Nass published an article reviewing the anthrax vaccine and its potential protective value against a biological attack with weaponized anthrax. While extensively researched and documented, she cites sources that are of questionable veracity. For example, she alleges the Defense Department may have attempted to increase the potency of Anthrax Vaccine Adsorbed by secretly adding squalene, citing herself as the source by referring to a letter she wrote to the Army Surgeon General in May 1998. Based on this allegation, she implies a potential connection between Gulf War Syndrome and Defense Department vaccination programs, including Anthrax Vaccine Adsorbed, which is the basis for most of the concern with the safety and efficacy of Anthrax Vaccine Adsorbed.

She also states in this article “the present human anthrax vaccine probably provides only limited protection for troops facing a BW (sic) attack by anthrax.” She bases this assertion on the lack of controlled studies in humans that investigate the clinical effectiveness of the vaccine against inhalation anthrax during a biological attack. Yet, she presents no clinical data of her own to substantiate her claim that the vaccine may not be effective in preventing inhalation anthrax after a biological attack. In other words, in her opinion, the more than 30 years of clinical data from field trials of anthrax vaccine in workers exposed to anthrax, the absence of inhalation anthrax in the workplace since 1978, and the animal studies that demonstrate Anthrax Vaccine Adsorbed’s effectiveness in preventing inhalation anthrax are not enough to conclude that the present vaccine may
prevent inhalation anthrax after an attack. Instead, she implies the only way to justify using the vaccine to protect against inhalation anthrax would be to design and conduct a study in which humans are deliberately exposed to aerosolized anthrax spores.

No vaccine is perfect, meaning that none is 100 percent safe and effective. But, as has been presented, the clinical evidence suggests that the current anthrax vaccine is safe and effective—probably safer with lower rates of side effects than other vaccines in use today. Even though the present vaccine is safe and effective, that does not mean there is no room for improvement regarding vaccinations against anthrax. The requirement for six inoculations does create a significant logistical problem especially as supplies of currently re-tested and approved lots of vaccine run low. But, the current requirement for six inoculations is in accordance with the FDA ruling and probably won’t change unless studies are done to confirm the vaccine provides protection with fewer doses. Newer vaccines that require fewer doses to confer immunity have been developed but have not been approved for use by the FDA. The 30 percent rate of occurrence of local reactions ideally could be lower, although this rate is already lower than other vaccines currently required by DOD.

A major challenge is how to demonstrate an individual has developed adequate immunity against anthrax after vaccination either with the present or a newer anthrax vaccine without exposing the individual to aerosolized anthrax spores. As discussed previously, it is not ethical to expose individuals to aerosolized anthrax spores to see if the vaccine prevents development of inhalation anthrax. Using animal models may or may not be useful since species differ in their sensitivity to anthrax and differences in their immune systems may alter the efficacy with which anthrax vaccines confer immunity. Measuring the level of antibodies an individual has circulating in the blood against protective antigen has been shown to be a very unreliable measurement of immunity against anthrax. The next best approach would be to develop a test that could be administered to the individual to indicate the degree of immunity. Currently no such test exists, which is one reason why the Food and Drug Administration recommends a series of six shots of Anthrax Vaccine Adsorbed with an annual booster. With such a test, individuals could be screened and only those with inadequate immune responses would require
supplemental inoculations, potentially decreasing the required number of doses of vaccine.

It should be reiterated, however, that even though there is room for improvement, none of these issues negate the current value and effectiveness of the vaccine. The risk from taking the vaccine is far less than the risk of being a target in a bio terrorist event or biological warfare attack.

Dr. Nass believes there should be more emphasis on using alternatives to vaccination with Anthrax Vaccine Adsorbed to protect troops from anthrax. For example, she suggests there should be more emphasis on the use of protective equipment. The problem with this approach is, due to the lack of real time detection capability, there is no way for personnel to know when they need to wear the protective equipment, meaning they would have to wear it continually to be effective. Dr. Nass also expresses concern that immunizing troops against anthrax may provoke an adversary to simply pick another biological agent. As previously discussed, other biological agents are more difficult to weaponize and the likelihood that other agents would be used in an attack instead of anthrax is much lower.

Lieutenant General (retired) James T. Scott recently wrote an editorial which, arguably, does more to place the entire controversy over DOD’s anthrax vaccination into proper perspective than any other work examined in this paper. He states that both sides share the blame for escalating this debate out of proportions. Officials from the Defense Department could have done better stating their case for a comprehensive vaccination program in peacetime. DOD’s credibility had already been damaged by how it handled the Agent Orange and Gulf War Syndrome issues. This problem is exacerbated by the chronic under-funding of the military health care system which is eroding away what little confidence beneficiaries may have in military health care and the Defense Department failed to anticipate the effect the Internet would have on spreading dis-information campaigns against the Anthrax Vaccine Immunization Program.

To the opponents who are also service members, Scott writes that it is time to find out the facts. He states service members concerned over the immunization program should be sure the information they possess is based on solid facts. He admonishes them to ask themselves if they are only concerned with the safety and efficacy of Anthrax Vaccine Adsorbed or if their concerns run much deeper -- that their opposition to the Anthrax
The current anthrax vaccine is a licensed vaccine and has been demonstrated to be clinically safe and effective for preventing inhalation anthrax after exposure to anthrax spores. Based on the findings of the 1985 advisory review panel examining the safety and efficacy of the vaccine, the FDA categorized the vaccine as a “Category 1 (safe, effective, and not misbranded) vaccine.” In spite of the existing documentation of its safety and efficacy, DOD continues to ask outside consultants and panels to review the evidence documenting the safety and efficacy of the vaccine. For example, the Defense Department asked the Institute of Medicine to review all available data on Anthrax Vaccine Adsorbed. One would be hard pressed to identify another vaccine in use today that has undergone more scrutiny than Anthrax Vaccine Adsorbed.

There are significant issues with Anthrax Vaccine Adsorbed that should be addressed, including the current dosage regimen, the inability to specifically measure the level of immunity an individual may already possess, and the occurrence of local reactions in 30 percent of those who are vaccinated. In spite of these issues, there is no clinical evidence that DOD’s Program is considered to be at risk for exposure to anthrax. The risk of serious adverse reactions or permanent injury from the anthrax vaccine is no higher than (and, in fact, is probably lower than) that for any other vaccine commonly in use in the general population today. In
contrast, the risks to military personnel from the threat of attack with an anthrax-based biological weapon, plus the high lethality of inhalation anthrax, far outweigh the risks associated with vaccination.

The large number of doses of vaccine required to establish immunity, plus the annual requirement for a booster, creates significant problems in terms of logistics and costs for the Defense Department to complete the Anthrax Vaccine Immunization Program and vaccinate all DOD personnel, especially in light of dwindling supplies of vaccine. Ideally, a reliable test to measure an individual’s immunity against anthrax should be developed. To ease the burden, only personnel expected to deploy to areas where the risk for potential use of weaponized anthrax is highest should be vaccinated. Military personnel not expected to deploy to these areas are at no greater risk for exposure to weaponized anthrax spores than the general population of the United States and need not be vaccinated. This is consistent with consensus panel recommendations that there is no requirement to vaccinate the entire population of the U.S. since the risk of exposure to weaponized anthrax for any given community within the U.S. is extremely low.143

The U.S. Army Medical Research Institute of Infectious Diseases completed pre-clinical research on a next-generation anthrax vaccine several years ago. The new recombinant vaccine is now in advanced clinical development. Unfortunately, Food and Drug Administration approval of a new vaccine is still several years away. In the meantime, long-term data collection studies should continue in order to document further the safety of Anthrax Vaccine Adsorbed and attempt to identify extremely rare adverse effects which may only become apparent after millions of doses of vaccine have been administered. The Defense Department should also continue with programs to provide long-term follow-up to individuals claiming to have developed symptoms after receiving the anthrax vaccine. These patients’ symptoms are real, and they deserve compassionate, professional medical care.

Continuance of the anthrax vaccine program should include an aggressive, active educational and informational program designed to address concerns at all levels, from the top leadership down to the installation level. The Defense Department web-site and its links to other service-specific web-sites are excellent but passive, meaning they depend
on people going to these sites to get the facts. What is needed is an active education program where information is actively taken out to the troops.

DOD programs actively promoting education of all military personnel, using the information on the internet web-site, could significantly alleviate the suspicions and doubts currently surrounding the anthrax vaccine immunization program. Commander and supervisor involvement at every level of command is essential to begin rebuilding the confidence military personnel should have in their chains of command. Commanders and supervisors also should be aware of the biased nature of informational internet web-sites opposed to the program, emphasizing to their personnel the importance of basing any conclusions about the vaccine or the anthrax vaccine immunization program on all the facts. Such proactive educational efforts should prove useful to reverse any negative trends and perceptions emanating from DOD’s handling of the Agent Orange and Gulf War Syndrome issues. The Anthrax Vaccine Immunization Program should be viewed as an opportunity for the Department of Defense to demonstrate its commitment to maintaining the health and safety of service personnel while countering any threat to our nation’s security from anthrax-based mass-casualty weapons.

Notes

1. This is the approach used in an article recently published by Mazzuchi, et al. In this article the authors emphasize that the decision to immunize is a command policy decision even though maintaining the health of the service members is the primary objective. (John F. Mazzuchi, Robert G. Claypool, Kenneth C. Hyams, David Trump, James Riddle, Relford E. Patterson, Sue Bailey, “Protecting the Health of U.S. Military Forces: A National Obligation,” *Aviation, Space, and Environmental Medicine* 71, no. 3 (March 2000), 260-265.)

2. “Peer-reviewed” refers to the process major medical journals use to decide if submissions are worthy of publication. Normally, the lead investigator submits a manuscript for consideration to the editorial board. The editorial board then selects members of the board (unknown to the author of the manuscript) to review the article to see if it meets stringent criteria such as scientific process, experimental design, analysis of the data, discussion, and conclusions. Some journal editorial boards “blind” the editorial
reviewers and authors from each other so the reviewers and authors do not know each other to make the review process more objective. The peer-review process is considered to be the most effective means of assuring quality publications in the medical literature. It should be noted that medical textbooks are not peer-reviewed although there is an editor to whom writers of the individual book chapters submit their manuscripts. Therefore, publishing in a textbook is not considered to be as significant as publication in a peer-reviewed journal.

3. For more detailed information, the reader is referred to the Defense Department’s informational world-wide-web site addressing anthrax vaccination at http://www.anthrax.osd.mil/. This site includes several links to papers, covering a variety of issues related to the anthrax vaccine. There are also several web sites outlining the reasons against the Defense Department’s anthrax vaccination program. The most prominent and complete with numerous links to other sites is http://www.dallasnw.quik.com/cyberella/index.htm. It is important to note that these sites present nearly identical historical, clinical, and factual information. Where these sites differ is how they interpret the information and what conclusions they draw.

4. Numerous extensive reviews of the disease process of anthrax, the vaccine, and the threat weaponized anthrax poses to United States military personnel already exist in the literature. Many are cited in this chapter.

5. Numerous countries, including those who are signatories of the Biological Weapons and Toxins Convention (including the former Soviet Union and Iraq) are known to have offensive biological weapons development programs, including development of weapons using anthrax as the agent. Major D. L. Clement, in an interesting study, concludes that overt use of tactical biological agents on the battlefield is unlikely due to difficulty in hiding the identity of the attacker and the risk of overwhelming response. Biological attacks against United States forces overseas are more likely to be on a small scale by terrorist groups. He identifies anthrax as the ideal biological warfare agent and concludes its use by terrorists or covert operators (such as special forces) against U.S. forces either in the U.S. or overseas is highly plausible, especially during deployments. (Major David Lee Clement, “A Determination of the Military Significance of Modern Biological Warfare,” Master’s Thesis, U.S. Army Command and General Staff College, Ft. Leavenworth, KS, (1993), 70, 79.) Inglesby, in his article published in the Journal of the American Medical Association (Thomas V. Inglesby, et al., “Anthrax as a Biological Weapon,” Journal of the American Medical Association 281, no. 18 (12 May, 1999), 1735-1745.), states anthrax is one of the most serious agents that could be used as a biological weapon, presenting a clinical discussion of anthrax to demonstrate why it would make such an effective weapon. (See also Mazzuchi, et al., 261.) For more information related to the threat anthrax poses as a
potential biological weapon, the reader is referred to the United States Air Force Counterproliferation Center’s world-wide-web site at http://www.au.af.mil/au/awc/awcgate/awc-cps.htm, which is updated regularly and contains numerous links to other important sites on this topic.


13. Holmes, 897.


15. Inglesby, 1736.

16. Brachman, 940.
17. Ibid., 943.
18 Ibid., 942.


22. Ibid., 1744.

23. Ibid., 1736-1737.


27. “Hemorrhagic meningitis” refers to inflammation of the protective coverings of the brain and spinal cord with associated bleeding.

28. Ibid., 819.

29. There are two different types of anthrax vaccine in existence today for human use. (‘An Assessment of the Safety of the Anthrax Vaccine: A Letter Report,’’ (30 March 2000), n.p.; on-line, Internet, 12 August 2000, available from http://www.nap.edu/html/anthrax_vaccine/.) Vaccines manufactured by filtering and purifying protective antigen (such as Anthrax Vaccine Adsorbed) are used in the West, primarily by the U.S. and the United Kingdom. The former Soviet Union manufactured an anthrax vaccine using live attenuated (weakened) anthrax spores. This type of vaccine is not available in the West. Although the efficacy of the live, attenuated spore vaccine has been reported to be higher than protective antigen-based vaccines (Hedlund, 64, and Meryl Nass, “Anthrax Vaccine,” *New Vaccines and New Vaccine Technology* 13, no. 1 (March 1999), 187-
208.), an obvious concern related to using this type of vaccine is that the live spores, although weakened, could still cause anthrax. (Inglesby, 1740.)

30. In addition to Anthrax Vaccine Adsorbed, there have been a number of experimental anti-anthrax vaccines developed but not necessarily tested or released for use in humans. All anti-anthrax vaccines developed in some way center on provoking an immunogenic response to protective antigen. (Hedlund, 64-68.)

31. Ibid., 67, 76.


33. In large doses and with repeated exposure formaldehyde may cause cancer. But there is no evidence that repeated doses of trace amounts of formaldehyde when used as a preservative in vaccines is harmful.


40. A study reported by Asa, et al., claims to have found that squalene antibodies are only found in the blood of people suffering from Gulf War Syndrome. This claim forms the basis of several press reports in the mainstream media accusing that the Department of Defense secretly added squalene to Anthrax Vaccine Adsorbed. (Pamala
B. Asa, Yon Cao, Robert F. Garry, “Antibodies to Squalene in Gulf War Syndrome,” *Experimental and Molecular Pathology* 68, no. 1 (February 2000), 55-64. It is very important to note that this study is poorly constructed and has been refuted due to insufficient numbers of subjects in the study population and lack of sufficient control groups. In addition, the authors themselves caution in the article that the results of their study do not establish that squalene was used in Anthrax Vaccine Adsorbed or any other vaccine during the Persian Gulf War.


42. The first anthrax vaccine was produced in the 1800’s by Louis Pasteur for use in animals. It was the first vaccine ever developed to protect against bacterial infection, making anthrax the first disease for which a vaccine was ever produced. Anthrax is probably the most studied and scrutinized of any bacterial infectious disease process. “Anthrax Vaccine Immunization Program,” n.p., on-line, Internet, 11 February 2000, available from http://www.anthrax.osd.mil/.


48 Ibid.

49. “Anthrax Vaccine Adsorbed Stockpile Analysis – Supplemental Testing Needed

50. Myers, n.p.


57. Ibid.


64. Myers, n.p.

65. Ibid.


73. Federal Register. See also, Lew, 1886. See also, LaForce, 1010.


75. The Food and Drug Administration has not approved Anthrax Vaccine
Adsorbed for use to protect against inhalation anthrax. But, lack of Food and Drug Administration approval for a specific indication does not prohibit the use of a medication for that indication if clinical evidence exists to support it. Food and Drug Administration licensure means that the medical product may be sold commercially in the U.S. Food and Drug Administration approval means testing for that indication has been completed according to Food and Drug Administration specifications in connection with the licensing process. Lack of Food and Drug Administration approval only means that the rigorous testing required by the Food and Drug Administration (paid for by the manufacturer) before it will grant its endorsement has not been completed. If there is reasonable clinical evidence to support using a medication for an indication not approved by the Food and Drug Administration, a physician may prescribe that medication for that use based on the physician’s clinical judgment. Such use is also not considered experimental since the medication is already licensed. In a letter to the Honorable Dan Burton dated 26 November 1999, Melinda K. Plaisier, the Associate Commissioner for Legislation for the Food and Drug Administration discusses the procedures by which lots are released by the Food and Drug Administration for sale and distribution. Regarding indications for use, she states, “The labeling for Anthrax Vaccine Adsorbed does not mention route of exposure (e.g., cutaneous) per se. Use of the vaccine for protection against both cutaneous and inhalation anthrax exposure is not inconsistent with the labeling for Anthrax Vaccine Adsorbed.” She adds that there is no reason for Anthrax Vaccine Adsorbed to be returned to an investigational new drug status when used to vaccinate against inhalation anthrax, especially since the rarity and risk of human inhalation anthrax precludes gathering additional clinical data.


82. Ellenberg, n.p.
88. Ibid., 4.
90. Inglesby, 1735-1736, 1740, 1744.
95. “Safety Review of Anthrax Vaccine (24 April 2000),” n.p.; on-line,


100. Anthrax Vaccine Adsorbed has been administered to thousands of veterinary and laboratory workers, livestock handlers, and workers in the wool sorter and animal hide industry since 1970. In addition, there have been at least four major independent reviews by civilian panels on the safety and efficacy of Anthrax Vaccine Adsorbed. (“Vaccine Safety,” (no-date), n.p.; on-line, Internet, 4 September 2000, available from http://www.anthrax.osd.mil/Site_Files/safety/safety_info.htm.)

101. It should be noted that the strength of clinical evidence would be enhanced if more data were published in the peer-reviewed literature and researchers should be encouraged to submit their data to the peer-review process. (“An Assessment of the Safety of the Anthrax Vaccine: A Letter Report.”, n.p.)

102. Ibid.


105. Dr. Nass cites a study in which different strains of anthrax were tested in guinea pigs vaccinated with Anthrax Vaccine Adsorbed. In the study, nine of 27 strains appeared to be resistant to vaccination. (Nass, “Anthrax Vaccine Safety and Efficacy: Response to the Army Surgeon General Ronald Blanck’s Posting.”) Friedlander, “Anthrax Vaccine: Evidence for Safety and Efficacy Against Inhalation Anthrax,” however, provides a thorough review of several animal studies, including the study cited
by Dr. Nass, in which Anthrax Vaccine Adsorbed is tested against various strains of anthrax in several different animal species. He demonstrates that, while Anthrax Vaccine Adsorbed provided variable protection against certain strains of anthrax in guinea pigs, Anthrax Vaccine Adsorbed provided excellent protection against even the most virulent and seemingly resistant strains (such as the Ames strain) in non-human primates and rabbits, even after just two doses of Anthrax Vaccine Adsorbed. This most likely reflects species-specific differences with the guinea pig’s immune system, making it more difficult to immunize the guinea pig against anthrax compared to other animal species, and is not due to any strain of anthrax developing resistance to the anthrax vaccine. (Friedlander, “Anthrax Vaccine: Evidence for Safety and Efficacy Against Inhalation Anthrax,” 2105-2106.)


108. Dr. Nass reports the results of a survey sent out to members of the 9th Airlift Squadron at Dover Air Force Base. In this report, she admits that the survey results cannot be used to establish that Anthrax Vaccine Adsorbed actually caused the symptoms due to the lack of a control group. Overall 252 surveys were sent out and 139 were completed and returned. She interprets 81 as probably having a systemic reaction due to Anthrax Vaccine Adsorbed. At least six indicated they felt Anthrax Vaccine Adsorbed did not cause their symptoms. Even though she admits there can be no statistical analysis of this data and it is not possible to prove causality, Dr. Nass concludes something must be wrong at Dover because it is just not normal for so many otherwise healthy people at one location to be having symptoms or strange illnesses of one sort or another. A useful analysis would be to compare the rate of occurrence of symptoms and diseases in 9th Airlift Squadron personnel against the rate of occurrence in all unvaccinated personnel at Dover and the U.S. population in general, but this is not provided. Nor is there an attempt to look for recurring patterns of symptoms. Indeed, it appears from the survey results that the patterns of symptoms vary widely, with no two individuals’ symptom patterns matching. (Meryl Nass, “Survey Results of the 9th Airlift Squadron,” (no date), n.p.; on-line, Internet, 27 August 2000, available from http://www.anthraxvaccine.org.)


111. Hedlund, 76.


118. Lieutenant General Hal M. Hornberg, ”Vaccine is Safe, Effective,” Air Force Times 60, Issue 16 (22 November 1999), 55.


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126. Ibid.

127. Several independent nationally renowned scientific groups have addressed this issue and have found no evidence to link anthrax vaccine with illnesses among Gulf War veterans. There have been several unsubstantiated allegations in the media and elsewhere about experimental vaccines that may have contained non-Food and Drug Administration-licensed substances such as squalene. Only the Food and Drug Administration-licensed anthrax vaccines have been used. (“Anthrax Vaccine and the Persian Gulf War,” (no date), n.p.; on-line, Internet, 15 February 2000, available from http://www.anthrax.osd.mil/oldavip/qna/GULFWAR.HTM.)


130. Laughlin, n.p.

132. Ibid., 199.

133. Ibid., 203.


135. New anthrax vaccines have been developed and are ready for clinical testing. But, so far, lack of funding has prevented the performance of clinical trial studies. (LaForce, 1013)


139. Ibid.

140. Ibid.


142. The letter dated 30 March 2000 is the result of a request the Secretary of Defense sent to the Institute of Medicine to assess the safety of Anthrax Vaccine Adsorbed. The Institute of Medicine commented that its assessment is an early step in sorting out the complex issues surrounding Anthrax Vaccine Adsorbed and has started a two year in-depth study which will include a review of all available data from the Department of Defense. (“A Letter Report from the Institute of Medicine (IOM), 30 March 2000” (no date), n.p.; on-line, Internet, 12 August 2000, available from http://www.anthrax.osd.mil/SCANNED/ARTICLES/ltrReportIntro.htm.)

143. Inglesby, 1740.