Chapter 1

Literature Review

1.1. Vaccines

With the increasing emergence of antibiotic resistant infectious organisms more emphasis should be placed on prevention of infection and disease via immunization and less on post-infection antibiotics. (Shearer and Clerici, 1997). The principle of vaccine development is to alter a pathogen or its toxin in such a way as to render them harmless without removing their ability to illicit a similar immune response as that of the same but unmodified pathogen. Since the antigenic determinants are still present on the vaccine, the antibodies generated by the adaptive immune response after vaccination would still be able to recognize the active pathogen during subsequent natural infection. The secondary antibody response would be mounted and the infectious particles would be brought under control faster and removed more effectively.

There are a few requirements that an ideal vaccine would have to satisfy. i) Elicitation of neutralizing antibodies that reacts with all of the pathogen’s strains and subtypes. ii) Induction of strong humoral and cytotoxic immune responses. iii) Induction of immune responses that recognize latently infected cells. iv) Mimic natural infection and induce local immunity at all sites of entry into the host. vi) Safety, showing no toxic effects or possibility of reversion to wild-type infection. vii) The effect of the vaccine must be long-lasting following a single inoculation. viii) Must be economically feasible, thus must be produced cost-effectively and have long-term stability in storage, excluding the use of cold-chains.

This chapter will briefly review the principles of different vaccine approaches, their respective advantages, as well as the major problems associated with each. Also included at the end of this chapter is an introduction into African Horse Sickness Virus (AHSV) Viral Protein (VP) 7 and its possible use as a subunit antigen presentation system.
1.2. Types of Vaccines

1.2.1. Inactivated Vaccines

Inactivated vaccines are infectious agents that cannot replicate and have thus lost their infectivity through damage to their nucleic acid by various agents. The method by means of which the nucleic acid is inactivated should not affect the antigenic specificity of the protein surface of the respective organisms. Common methods for the inactivation of these pathogens include heat-inactivation, irradiation (Huter et al., 2000), chemical treatment, especially with formaldehyde (Bahnemann, 1990) and ultrasonication (Melamed et al., 1991). The more frequently preferred formaldehyde method of inactivation has successfully been used in several cases, including the first worldwide commercialized production of a vaccine against Hepatitis A (André, 1995). It has been found that these inactivation treatments might, however, result in reducing or altering the vaccine’s antigenic character and could be responsible for the loss of relevant immunogenic epitopes (Duque et al., 1989; Ferguson et al., 1993). A further complication occurs when it was found that certain bacteria only generate specific protective antigens on their outer membrane in specific in vitro growth conditions (Deneer et al., 1989). Also, with pathogens that aggregate it has been found that a miniscule proportion of the infectious agents in the core of the aggregate may "survive" the inactivation treatment and produce the disease, which the vaccine is supposed to prevent (Brown, 1993).

Inactivated vaccines normally elicit a narrow immune response spectrum. Their inability to infect cells results in good humoral immunity but limited, if any, cell-mediated - or mucosal immunity. In spite of this, inactivated vaccines still is one of the two major models for vaccine success, the other being live attenuated vaccines (Hilleman, 2000).

1.2.2. Live Attenuated Vaccines

Live attenuated vaccines are infectious pathogen strains that have lost their virulence or disease causing properties. These attenuated vaccines retain their ability to infect and replicate in cells and are capable of inducing strong, long-lasting immunity, cellular and humoral, similar to that induced by the natural infection (Ruprecht, 1999). Live attenuated vaccines can be delivered via the natural route of infection, thereby providing better protection at the site of entry of the pathogen. It is important to induce mucosal immunity since most pathogenic organisms are either restricted to these areas or need
to transit them during the early steps of infection (Levine et al., 1983). These vaccines, because of their replicating ability within the host, generally induce a longer duration of immunity (Ellis, 2001). However, over-attenuation will limit replication, which might result in mild, non-protective immune response.

The ability of live attenuated vaccines to infect and replicate also cause the main problem, especially if under-attenuation occurred. There exists the possibility that the pathogen might back-mutate after in vivo replication and revert back to the lethal wild-type strain (Appel, 1978; Minor et al., 1986). This is a particular problem in immunorepressed individuals where even a weakened pathogen can cause the disease. Another disadvantage of live tissue culture grown vaccines is the risk of contaminating extraneous viruses in the tissue culture serum being used (Thornton, 1986). One popular example involved the introduction of bovine viral diarrhea into vaccines grown in bovine cells (Wensvoort et al., 1988).

Despite all the disadvantages and safety concerns, the importance of live vaccines is best highlighted by the fact that ten of the seventeen licensed viral-based vaccines in the United States of America are live vaccines (Hilleman, 2000).

1.2.2.1. Biological Attenuation

Biological attenuation, though not very well understood, is frequently attained through passage through atypical host animals or cultured cells (Vaughn et al., 1996). The unfavorable growth conditions, which could also be induced by mutagenic chemicals (Levine et al., 1987), give rise to a random series of attenuated or mutant pathogens. These new mutant strains are constantly monitored and selected for antigenic retention and loss of virulence. A combination vaccine for African horse sickness, which contains eight of the nine distinct serotypes, was attained through biological attenuation. Three of the eight serotypes were plaque attenuated by passage in cell culture, whereas the other five strains were attenuated by serial intracerebral passage in adult mice (Taylor et al., 1992).

Biological attenuation can also be achieved by selecting for mutants according to their growth properties at different temperatures (Treonor et al., 2000). These cold adapted temperature-sensitive pathogens are unable to grow efficiently at elevated temperatures and are consequently less virulent than the wild-type. Also, co-infection of cells with two or more different strains of a virus with
segmented genome could lead to the production of reassortant viruses that are less virulent than the wild-type strain, which could be used as a vaccine (Rennels et al., 1996). This same principle is also the cause for concern when vaccination occurs with a biologically attenuated serotype combination vaccine. Reassortment of the genomic segments may occur between the different serotypes during replication in the vaccinated host, which might result in progeny virus with novel characteristics, including new virulence characteristics (Nutall et al., 1992).

1.2.2.2. Genetic Attenuation

Recent advances in recombinant DNA technology and knowledge have made it possible to attenuate pathogens through the genetic manipulation of specific genes responsible for toxicity. This method is more preferred than biological attenuation. More stability is attained in the phenotype through extensive modification or deletions to the pathogenic genome to reduce virulence. By creating multiple modifications or deletion of entire genes the possibility of back-mutation to the virulent form is decrease (Kit et al., 1987).

It has been shown that attenuating genes can be in the structural (Farrell et al., 1994), non-structural (Liang et al., 1997) or even in the noncoding region of the pathogen’s genome (Kuhn et al., 1992). Generally non-essential genes are deleted that may inhibit sufficient replication of the pathogen, thereby prohibiting it from causing the disease. Viruses can also be attenuated by making recombinant chimeric viruses consisting of complementary but not fully compatible genes from related viruses (Kuhn et al., 1991).

Another way to genetically attenuate pathogens would be to delete or mutate genes that are necessary for in vivo replication. Viruses can be rendered replication-incompetent by propagating them in vitro in a cell line transfected with the essential gene (Reddy et al., 1999; Zakhartchouk et al., 1999). When introduced into an individual, the virus would be able to infect yet not replicate inside the host since it would not have access to the essential gene (Farrel et al., 1994). A variation of this approach is applied with bacterial vaccines that are engineered to be metabolite-dependent. Here the metabolite-dependent bacterial vaccine is grown in vitro in the presence of the required metabolite. After injection into an individual, the bacterium will continue propagating as long as the individual takes supplements of the metabolite. Once the supplementation stops, the bacteria die (Donnenberg et al., 1991).
It has also been shown that the establishment of persistent infection with retroviruses can be prevented by the mutation or deletion of the integrase gene (Vogel et al., 1993).

1.2.3. Nucleic Acid Vaccines

1.2.3.1. DNA Vaccines

It has been shown that naked DNA encoding vaccine antigens can be transfected in vivo into cells (Wolff et al., 1990). The produced antigens are either secreted to elicit a humoral immune response or they can be associated on the cell surface to elicit a cellular immune response. Only small quantities of proteins being expressed are required for effective priming of the immune response. The antigens are continuously produced in the cells, even for several months after DNA immunization (Yankauckas et al., 1993), which is probably responsible for inducing long-term immunity. Priming with DNA vaccines followed by boosting with the antigenic protein itself induce higher levels of protection than with either DNA immunization or protein immunization individually (Amara et al., 2001).

The attractiveness of this approach is in its simplicity. Expression of the gene is coupled to a strong upstream promoter capable of expressing the gene in mammalian cells. Usually the cytomegalovirus immediate/early promoter is used (Donnelly et al., 1997). The introduction of introns into the plasmid, resynthesizing genes to remove cryptic splice sites and improving codon biases have all been shown to improve the expression levels of coded antigens (Vinner et al., 1999). A requirement for DNA vaccines to elicit a strong humoral immune response is the presence of unmethylated CpG sequences in the plasmid backbone. These dinucleotides induce murine B cells to proliferate and secrete immunoglobulins (Krieg et al., 1995; Davis et al., 1998). Adjuvants are unnecessary with DNA vaccines containing the CpG sequences.

An advantage of DNA immunization where viral antigens are concerned is that the DNA encoded antigens’ post-translational modifications are exactly the same as that of a natural viral infection. Also, the viral antigens are being presented efficiently on the MHC molecules for T-cell recognition (Ulmer et al., 1996) It is also possible to manipulate and shift the immune response to predominantly either a humoral or cytotoxic immunity by the co-administration of plasmid encoding cytokine (Kwissa et al., 2003) or interleukin genes (Kim et al., 1997). Mucosal immune responses can be generated by delivery of the DNA plasmids to the mucosal surfaces (Fyan et al., 1993; Kuklin et al., 1997) and it has been
reported that DNA vaccines can induce immune responses in neonates even in the presence of passive antibodies (Lewis et al., 1999).

The first study involving an African horse sickness virus DNA vaccine was reported by Romito et al., 1999. The outer capsid protein viral protein 2 (VP2) generates the main determinant of serotype-discriminatory neutralising-specific immune responses for African horse sickness. (Huismans et al., 1987; Roy et al., 1996; Scanlen et al., 2002). A horse was injected with plasmid DNA containing the AHSV serotype 3 VP2 gene downstream of a cytomegalovirus immediate-early enhancer/promoter. Viral protein 2-specific antibodies were generated but neutralisation titres were low.

Although naked DNA vaccines have been shown to function without a protective delivery system, a problem that occurs is that the majority of the plasmid is degraded before entering the nucleus of a cell and initiating transcription of the gene of interest, thereby lowering expression levels. It is possible to use, for instance, the alphavirus envelope proteins to encapsulate and deliver the nucleic acids directly to the cells (Pushko et al., 1997; Schultz-Cherry et al., 2000). Chemical approaches have also been tested, including using various polymers and liposomes to encapsulate the DNA (Ellis, 2001).

There are other major concerns with DNA vaccines for which there have been no solution to date. The major problem is that the plasmid DNA might integrate into the genome of a multiplying host cell, and thereby activate a host protooncogene or deactivate a suppressor gene, which could lead to cancer (Xiong et al., 1997). There also exists the risk that the plasmid could induce an autoimmune response to the host cell’s DNA.

1.2.3.2. RNA Vaccines

An alternative to DNA vaccination is the use of RNA vaccines. The RNA fragments are less stable in vivo than the DNA plasmids used for vaccination. Subsequently the expression of the antigens is short-lived and therefore, an overall less effective approach than DNA vaccination. Expression levels can be boosted through the incorporation of a gene encoding an alphaviral replicase together with the gene encoding the foreign antigen, which would enable the RNA to self-replicate in the transfected cell (Tubulekas et al., 1997). This approach has been shown to generate strong immune responses (Zhou et al., 1994). Like DNA vaccines, these RNA fragments can also be delivered naked or encapsulated in
either non-replicative viral particles (Zhou et al., 1995) or liposomes (Martinon et al., 1993; Conry et al., 1995).

RNA vaccines, due to their inherent low stability, generally are problematic when prepared and administered. Despite these technical and other mentioned problems, the main attraction of RNA vaccines, unlike DNA vaccines, is their inability to integrate into the genome of the host cell and possibly cause cancer.

1.2.4. Live Recombinant Vaccine Vehicles

Non-pathogenic or attenuated microorganisms, be they viral or bacterial, can be engineered as vectors for expressing foreign antigenic polypeptides from other non-related pathogens (Yilma et al., 1988). Insertion of cloned genes from pathogens into the genomes of other live organisms occurs so that during infection with the recombinant vector, the surface proteins produced by the cloned genes are presented on the infected cells by the MHC complex to the T-lymphocytes (Ellis 2001). Alternatively the expressed proteins are broken up and presented to elicit a cytotoxic T-lymphocyte response. Antibodies are produced that will neutralize the pathogen from which the surface proteins were cloned.

It is also possible to express the foreign antigen on the surface of the recombinant vector. This will have the desired effect of also stimulating the humoral immune response (Liljeqvist et al., 1999). Heterologous cell-surface display in the context of live bacterial vaccines have been described for both Gram-positive (Fischetti et al., 1996; Stähl et al., 1997) and Gram-negative (Georgiou et al., 1993) bacteria. The heterologous use of viral systems have shown to elicit strong immune responses against the foreign antigen (Dalsgaard et al., 1997) and have the advantage of targeting the recombinant viral particles to specific cells within the host (Ohno et al., 1997). Another advantage of live recombinant vaccine vectors is that even though the host might already have immunity against the vector itself, a strong immune reaction can still be elicited against the added antigenic polypeptides it carries (Zakhartchouk et al., 1999).

A problem that might occur is that one of these "harmless" vectors might have lethal consequences in certain individuals, especially those that are immunocompromised. This problem, however, can be overcome by the incorporation of cytokines or immunomodulatory genes into the vector (Kurilla et al., 1993; Sambhi et al., 1991). The expression of these genes might enhance the immune response to the
pathogen. Another solution would be the use of related viruses from other species that are replication-defective in mammals (Fries et al., 1996; Somogyi et al., 1993). These vectors would be able to infect mammalian cells, express the transgene that would elicit its own immune response, yet would be unable to replicate.

The use of live recombinant vehicles seems promising for vaccine purposes. One of the best examples and greatest successes of a vectored vaccine is vaccinia virus carrying the rabies virus glycoprotein gene (Brochier et al., 1994). The ability of thermal stable vaccinia virus to infect foxes by oral ingestion provided the basis for a campaign to reduce rabies infection in foxes in Western Europe. The campaign started in Belgium during 1989 by distribution of vaccine immunized bait from airplanes. This strategy completely eliminated rabies virus infection from Belgium’s fox population by 1993. Vaccinia virus was also used as live recombinant vehicle for the delivery of AHHSV serotype 4 VP2 for immunization in horses. The immunization approach was successful in inducing a protective immune response against a challenge with a lethal dose of AHHSV-4 (Stone-Marschat et al., 1996).

There are, however, still several problems associated with this delivery system (Medina et al., 2001). These include the possibility of reversion back to the virulent phenotype, the stability of the recombinant phenotype, horizontal gene transfer to the environment that were not apparent in the small-scale trials and a possible reduction in efficacy of the vaccine because of prior exposure to the carrier. Finally, as mentioned before, the host's genetic factors might influence the vaccine efficacy.

1.2.5. Antibody-based Vaccines

1.2.5.1. Passive Vaccination Strategies

Passive vaccination is the injection of preformed antibodies into a host to treat an infection. It is especially effective where pathogenic circulating toxins are produced and a high titre of specific antibodies is required (LeClaire et al., 2002). Various expression systems can be used to produce monoclonal antibodies, including bacteria (Laden et al., 2002), plants (Hiatt et al., 1989; Stoger et al., 2002), yeast, and mammalian cells (Little et al., 2000).

The increase in knowledge concerning the genetics and structure of the immunoglobulins and advancements in recombinant DNA technology has permitted the genetic manipulation of antibody
molecules. Chimaeric antibodies can now be created to suit their respective antigens and functions (Boulianne et al., 1984; Better et al., 1988). Research into the use of fragments derived from antibody molecules has also advanced. It has been shown that these smaller fragments have better penetration of solid tumours and rapid clearance abilities, compared to whole antibodies (Huston et al., 1993). These fragments can also be fused to other molecules, such as cytokines (Boleti et al., 1995), metal-binding proteins (George et al., 1995), toxins or drugs (Huston et al., 1993) to expand on their medical applications.

The absence of any viral or bacterial material is a great advantage, although the expression of these antibodies causes a huge problem. The chosen expression system has to be optimized for every different antibody because of their differences in amino acid sequence (Verma et al., 1998). The optimality of the expression system would also depend on the required quantity and the required purity of the final product. Another problem is that with complex antigens different epitopes maybe be instumental in stimulating various aspects of the immune response required for effective protection (Adorini et al., 1979). Since monoclonal antibodies are specific only for a single specific epitope, a pathogen could easily elude an immune response. Also a mutagenic slight alterations to the specific epitope may cause the failure of the antibody.

1.2.5.2. Anti-idiotype Antibody Vaccines

Another application for antibodies in vaccination studies is the anti-idiotype antibody. These antibodies are created by using neutralizing antibodies against the virus as the immunogen in an animal host. The newly generated antibodies induced against the first antibodies (anti-idiotype antibodies) should resemble the epitope on the virus that is susceptible to the neutralizing antibody. This anti-idiotype, therefore, when used to immunize another host, should elicit neutralizing antibodies against itself and by analogy against the infectious virus itself (Dalgleish et al., 1988).

Immunization with anti-idiotype antibodies has been shown to confer protection in animal disease models (Poskitt et al., 1991), although the same problems as with passive vaccination strategies are found with this vaccination strategy.
1.2.6. Subunit Vaccines

1.2.6.1. Particulate and Fusion Vaccines

Subunit vaccines are based on the principle that it is possible to elicit the same immune response against specific proteins from an infectious organism, as from a natural infection. This approach focuses on the use of whole surface proteins as possible vaccines since they are the first proteins the immune system normally comes into contact with and against which antibodies are generated (Burnette, 1991). It is important to have prior knowledge of the pathogenic components that are involved in inducing a protective immunity.

Recombinant DNA technology has further broadened the potential uses of subunit vaccines. Genetic fusion of foreign antigenic regions from one organism to a subunit protein from another organism, which has suitable immunogenic properties, is now possible (Uhlen et al., 1992). Fusion proteins, therefore, have the combined properties of the original gene products. This allows for the generation of immune responses against foreign antigens that are being stabilized and presented on carrier proteins. Depending on the chosen epitopes, both the T- and B-cell mediated immune responses can be generated (Löwenadler and Lycke, 1994). Care should be taken not to alter the structure of the carrier protein with the insertion of the epitope amino acids.

These proteins can be produced en mass utilizing different expression systems. The expression system for a specific protein’s production will depend on the type of organism the protein was originally derived from. This is done to ensure that the protein undergoes the correct posttranslational modification (Dertzbaugh, 1998). Many techniques have been developed for the extraction and purification of fusion proteins. A good example of purifying the proteins from a solution would be affinity handles. It is possible to add affinity fusion partners, which enable purification by affinity chromatography (Nygren et al., 1994). Site-specific enzymatic cleavage can remove the affinity handles after purification (Nilsson et al., 1997). Some of the expressed proteins, like viral envelope glycoproteins, insert themselves into the host cell’s membrane. The only way to harvest these proteins is by lysis of the cell, which prolongs and increases the cost of production. Techniques have been developed to remove the transmembrane anchors of these membrane-bound proteins so that they can be secreted into the medium without lysis of the host production cells and subsequently easily harvested from the medium (Kowalski et al., 1993).
Virus-like particles (VLP) can also be used as subunit vaccine or as carriers for epitopes from other viruses (Roy, 1996). These particles are attained from protein-protein self-assembly in cells expressing the required envelope proteins (Kunkel et al., 2001). The particles seem to induce a more effective immune response than denatured or soluble proteins (Netter et al., 2001). The reason for this enhanced immunity might be linked to the fact that VLPs are presenting confirmation epitopes to the immune system, which mimics a natural infection.

The only subunit vaccine, which is also the only recombinant expressed vaccine of the seventeen licensed viral-based vaccines in the United States of America, is that of hepatitis B (Hilleman, 2000). Valenzuela et al. (1982) found that Saccharomyces cerevisiae expressed hepatitis B virus surface antigen (HbsAg) assembled into subviral particles similar to that of the hepatitis B virus. Since then it has been found that extensive modifications can be made to the primary structure of HBcAg without impairing particle assembly, even insertions as large as 238 amino acids (Beterams et al., 2000). These highly immunogenic particles can effectively serve as carriers for the presentation of other peptide sequences.

Not only proteins that form the outer capsid or membrane of the organism can be used as subunit vaccines. Many bacteria secrete toxins that are responsible for pathogenesis and tissue damage (Shewan et al., 1982). It is possible to use these extracellular toxins not only as subunit vaccines to induce effective immunity, but also as carriers for epitopes from other organisms (Rauly et al., 1999).

There are several advantages to subunit vaccines. They cannot replicate and are non-infectious, nor is any inactivation agents required that might alter the immunogenicity of the protein. As mentioned before, it is possible to introduce specific T- and B-cell epitopes into the subunit proteins thereby directing and enhancing the immune response. Another advantage, when compared to inactivated vaccine systems, is the absence of irrelevant protein components, which decreases the antigenic competition in the vaccinated host.

As well as advantages, there are problems associated with the use of subunit vaccines. Envelope proteins might not induce a sufficiently strong cell-mediated response against virus infected cells. Disease might be caused by a great variety of viruses and their strains (Wareing and Tannock, 2001). Many viruses change frequently through antigenic drift (Jong et al., 1999). There might be contaminants in growth media (Thornton, 1986; Wensvoort et al., 1988). The major problem with subunit vaccines is the size of the antigenic insertions that can be made without the alteration of the
protein structure (Roy, 1996). It is only possible to insert small antigenic regions without disturbing the overall protein structure.

1.2.6.2. Synthetic Peptide Vaccines

Another type of subunit vaccine is the synthetic peptide vaccine. Synthetic peptide vaccines are based on the principle that it is possible to generate neutralizing antibodies against the intact virus particle using small defined peptides representative of the antigens found on the viral envelope or surface proteins (Van Regemortel, 2001). These peptides mimic the 8 to 12 long amino acid fragments that are normally displayed on the cell surface by major histocompatibility complex (MHC) molecules for T-cell recognition (Nelson et al., 1992). They are also able to generate B-cells that are less conformation-dependent and which recognize linear epitopes. Depending on the epitopes from the pathogen that are being used, either the humoral or the cellular branch of the immune system can be predominantly stimulated. It has also been found that a better and broader immune response is generated when different multiple epitopes are used as synthetic peptide vaccines (Tam, 1996). Multiple copies of a single epitope can also be synthesized, which leads to a stronger immune response subsequent to injection (Coéffier et al., 2001).

These peptides are strongly immunogenic when delivery is coupled with either a strong adjuvant (Hsu et al., 1996) or fused with a carrier protein (Kaumaya et al., 1992). When delivered alone, however, synthetic peptides are rapidly cleared in vivo and are subsequently not very immunogenic (Ben-Yedidia et al., 1997). Also, it has been found that synthetic peptide vaccines are less efficient at generating humoral immune responses than cellular immune responses. The reason for this is that the free flexible B-cell linear peptides raise antibodies that do not bind optimally to the sequence in the way that they do when it is present in a more rigid structure within the native protein molecule (Jemmerson, 1987; Jackson et al., 2000). The synthetic peptide might also form secondary structures by means of disulfide bonds, which might lower the immunogenicity of the vaccine (Simard et al., 1997). Research into the chemical stabilization of the peptides in solution is being conducted with the hope of improving the immunogenicity (Cabezas et al., 2000).

Despite all the disadvantages, this method of vaccination offers the ideal in safety, specificity, quality and economy since specific synthetic peptides are generated without any pathogen material (Ertl et al., 1996). Consequently there are no infectious particles still drifting around and no contamination with
harmful materials from the cells. However, it must be mentioned that these synthetic peptides are only economically feasible as long as the total length of the peptide does not exceed fifty amino acids.
1.3. African Horse Sickness Virus

African horse sickness is an infectious, but non-contagious gnat (*Culicoides* sp.)-transmitted disease, which affects species of the equidae. The disease, which is endemic to sub-Saharan Africa, causes a high mortality in horses, often exceeding 90%. It also occurs in donkeys, zebras and goats, but at a considerably lower death rate (Borden *et al.*, 1971). The causative agent is a dsRNA virus of the *Orbivirus* genus, belonging to the family *Reoviridae* and is closely related to bluetongue virus (BTV), which is the prototype virus of this genus (Basak *et al.*, 1996).

Ten dsRNA segments of different sizes compose the viral genome, three large, designated L1-L3, three medium, M4-M6, and four small, S7-S10. This genome encodes at least ten proteins, seven of which are structurally related (VP1 to VP7) and four non-structural proteins (NS1 to NS3A) (Van Staden and Huismans, 1991). The morphology of purified AHSV particles is essentially identical to that of BTV particles (Burroughs *et al.*, 1994). The seven structural proteins are organised in a two-layered capsid. The genome is packaged in a highly ordered icosahedral inner core composed of two major proteins, VP3 and VP7, and three minor structural proteins VP1, VP4, and VP6. The outer capsid, which is removed when the virus infects the cell, is composed of VP2 and VP5 (Roy *et al.*, 1994). The inner core is also able to infect certain types of cells without the aid of the outer capsid proteins (Mertens *et al.*, 1996).
1.3.1. African Horse Sickness Virus Viral Protein 7

African horse sickness virus VP7, which is coded for by the AHSV S7 dsRNA segment, has 349 amino acids. The sequence is highly hydrophobic, containing a large number of alanines, methionines, and prolines (Roy et al., 1991). There are nine serotypes for AHSV and VP7 has been found to be highly conserved between these serotypes. Three VP7 monomers (M, 38K) spontaneously organise into a trimeric structure of which the three subunits are non-covalently bound. The main outer surface components of the core particle are VP7 trimers. Two hundred and sixty of these VP7 trimers are arranged on a VP3 scaffold forming a T=13 lattice. Co-expression of the VP7 and VP3 genes of AHSV in a baculovirus expression system results in the formation of empty core-like particles (CLP) that resemble normal cores. (Maree et al., 1998).

The VP7 monomer is composed of two distinct domains, a bottom and a top domain. The bottom domain (residues 1-120 and 250-349) forms a complex of hydrophobic loops and nine α-helices, five in the N-terminal portion and four in the C-terminal region. The more hydrophilic top domain
(residues 121-249) is folded into anti-parallel β-strands connected by β-turns (Basak et al., 1996). It is one of the helices from the bottom domain of the VP7 trimer that forms a flat surface that interacts with the VP3 scaffold. The domains of the three monomers are twisted anti-clockwise around the three-fold axis of the trimer, allowing the top domain of one monomer to rest on the C-terminal region of the adjacent monomer within the same trimeric subunit (Grimes et al., 1995). The last 16 residues at the C-terminal have been shown essential for trimer interaction (Le Blois, et al., 1993). Additions of extra amino acids to the C-terminus inhibit core formation (Le Blois, et al., 1993; Monastyrskaya et al., 1997).

![Diagram](image)

**Figure 1.2.** Diagrammatic representation of (a) the AHSV VP7 protein with the three domains and (b) a VP7 trimer.

As mentioned in Section 2, the virion, lacking the outer capsid, is still able to infect certain types of cells. Blue tongue virus VP7 attaches to a cell surface receptor and allows the BTV core to pass
through the cell’s membrane. This is mediated by the RGD (Arg-168–Gly-169–Asp-170) motif (Grimes et al., 1995). In AHVS VP7 this motif is replaced by an AGQ (Ala-167–Gly-168–Gln-169) motif, which probably does not fulfill the same function as that of the RGD motif. African horse sickness virus VP7 does however, contain a RGD motif on a highly flexible loop, amino acid segment 175-180, at amino acid position 178 to 180. This loop is located deeper within the core than that of the easily accessible BTV motif, at the lower part of the top domain. It has been shown though that the RGD motif can still bind integrins effectively (Basak et al., 1996).

A unique feature of AHVS VP7 is the ability of the VP7 trimers, when expressed in a recombinant baculovirus, to spontaneously aggregate in infected insect cells (Spodoptera frugiperda) into large, flat, hexagonal crystals (Chuma et al., 1992; Burroughs et al., 1994). This same phenomenon is found when AHVS VP7 is expressed in BHK21 cells (Wade-Evans et al., 1997). Infected cells have been found to contain between one to three crystals per cell. These hexagonal crystals have an average diameter of about 6 μm and are not found when expressing BTV VP7 which is completely soluble (Oldfield et al., 1990). Such large structures are unique to AHVS within the Orbivirus genus. Also, these structures have a highly ordered lattice that is consistent with a trimeric subunit structure and is probably due to hydrophobic interactions between the trimers (Chuma et al., 1992; Basak et al., 1996).

1.3.2. African Horse Sickness Virus Viral Protein 7 Vaccine

African horse sickness virus VP7 crystals have been shown to be highly immunogenic when used as a subunit vaccine and are able to elicit a strong immune response in a mouse model against subsequent viral infections (Wade-Evans et al., 1997). The attained response was shown to be crystal-structure dependent since the same amount of heat denatured VP7 protein could not elicit the same level of protection. The passive transfer of antibodies from immunised mice failed to protect non-immunised mice against a subsequent viral challenge. This indicates that antibodies were not primarily responsible for the protection generated in the VP7 crystal vaccinated mice. It is possible that a protective T-cell response was generated with vaccination though further studies are required to confirm this (Wade-Evans et al., 1998).

The use of BTV CLP consisting of VP3 and VP7 trimers has also been investigated as a vaccine delivery system for foreign epitopes (Roy, 1996). Chimearic VP7 particles were created by the insertion of 15aa, representing a Bovine Leukemia Virus (BLV) glyco protein (gp) 51 epitope, into the
Ala 145 site of VP7. The Ala 145 site was chosen for insertion since it forms an exposed loop on the CLP's surface. The production of stable CLPs after co-expression of the chimeric protein with VP3, indicates that the inserted epitope does not disturb the structural stability of the VP7 trimers. Preliminary data from subsequent immune response tests indicated that the epitope is presented effectively and could, possibly, elicit humoral immune responses.
1.4. Aims and Strategy of this Study

The aim of this study is to investigate the structural constraints associated with utilising Viral Protein 7 of African Horse Sickness Virus as a particulate, multiple peptide vaccine delivery system. To effectively fulfil the purpose of this study, the aim can be broken down into three individual parts.

1. To enable AHHSV9 VP7 to act as a particulate, multiple peptide vaccine delivery system, multiple cloning sites for the insertion of peptides first have to be created. The first aim is to investigate the effect on structure and solubility of AHHSV9 VP7 by the insertion of three multiple cloning sites.

Strategy:

- Introduction of site specific mutations into VP7
- Expression of newly constructed mutant genes in a baculovirus expression system
- Solubility and structural analysis of expressed proteins

2. The purpose in constructing a VP7 protein with three multiple cloning sites, is for it to be able to accommodate and efficiently present multiple epitopes to the immune system. The second aim is generated from this: investigating the effect on structure and solubility of modified AHHSV9 VP7 by the insertion of peptides at two of the multiple cloning sites.

Strategy:

- Cloning of the peptides into two sites of modified AHHSV VP7
- Expression of newly constructed mutant genes in a baculovirus expression system
- Solubility and structural analysis of expressed proteins

3. The major disadvantage of virus like particles as antigen carriers, is the limitation in the size of the peptides that can be inserted. In order to utilise the full potential of the VP7 particles effectively as an
antigen presentation system, an investigating was launched into the effect on structure and solubility of modified AHSV9 VP7 by the insertion of different size antigenic sequences.

**Strategy:**

- Cloning of the AHSV4 VP2 stretches in the 177 amino acid site of modified AHSV VP7
- Expression of newly constructed mutant genes in a baculovirus expression system
- Solubility and structural analysis of expressed proteins
Chapter 2

The Creation of the Three Site VP7 Construct and the Characterisation of the Structural Features and Stability.

2.1. Introduction

In order to utilise the VP7 crystals as a multiple antigen presentation system, areas have been identified within the VP7 sequence into which the foreign antigens can be inserted. The more hydrophilic top domain of AHSV VP7 is exposed on the surface of the assembled hexagonal crystals. This domain contains a very small number of hydrophilic regions that are potential target sites for the insertion of foreign peptides or epitopes. A particular promising site is the RGD segment contained in a highly flexible loop (amino acid position 175 to 180) in the top domain. Amino acids position 144/145 as tested by Roy, 1996 and amino acid position 200/201 (Maree, 2000 unpublished data) are both hydrophilic areas that are on exposed loops and should provide little if any structural hindrance to the VP7 trimeric structure when modifications are made. These areas should also not affect the trimer-trimer interactions that are necessary for the formation of the VP7 crystals.

![Diagram](image)

Figure 2.1. Diagrammatic representation of the accessible regions on a VP7 trimer.
A recombinant VP7 construct, VP7mt144/200 (Fig. 2.2), has already been created by the insertion of two multiple cloning sites into aa position 144/145 and 200/201 (Riley, 2003). Restriction enzyme sites *SmaI*, *EcoRI* and *XhoI* were inserted at the 144/145 site whereas restriction enzyme sites *HindIII*, *XbaI* and *SaiI* constitutes the multiple cloning site at aa position 200/201. These primarily hydrophilic insertional modifications increased the solubility of the VP7 protein and weakened the trimer-trimer interactions but did not seem to prevent crystal formation. It was, however, noted that the increase in solubility of these recombinant VP7 molecules produced crystals with a more “ruffled” appearance, instead of the smooth hexagonal crystal-like appearance as formed by wild-type VP7 (Riley, 2003).

![Diagram](image)

**Figure 2.2.** Diagrammatic representation of VP7mt144/200 (Riley, 2003)

The VP7mt144/200 construct served as template for the creation of a construct with three multiple cloning sites at aa position 144/145, 177/178 and 200/201. The extra multiple cloning site would be advantageous if this construct is to be used for a multiple antigenic subunit vaccine. Experiments were also carried out to study the effect of sonication on the crystal structure of VP7. This investigation was launched in order to find a suitable method to disrupt the aggregation of the assembled VP7 crystals. An extensional study for the possible use of recombinant VP7 as a vaccine was done to evaluate the stability of the recombinant VP7 under different storage conditions.
2.2. Materials and Methods

Materials

Life Technologies provided the oligonucleotide primers, TaKaRa Ex Taq™ polymerase, as well as the Bac-to-Bac™ baculovirus expression system and Cellfectin™. Grace’s insect medium and foetal calf serum were purchased from Highveld Biological, whereas Rainbow™ protein molecular weight marker was purchased from AEC Amersham. Roche Diagnostics provided the restriction endonucleases, RNase A, T4 DNA ligase, DNA molecular weight marker II (MWII) and the High Pure™ DNA PCR Product and Plasmid Purification Kits. Perkin Elmer Biosystems ABI PRISM™ provided the Big Dye Terminator Cycle Sequencing Ready Reaction Kit. The Density Marker Beads were acquired from Pharmacia Biotechnology. The NERC Institute of Virology and Environmental Microbiology supplied the Sf9 cells. The VP7mt144/200 plasmid construct was obtained from Me. J. E. Riley. Merck and Sigma-Aldrich provided all the other chemicals.

2.2.1. Polymerase Chain Reaction

Polymerase chain reaction was used to introduce the site-specific mutation at amino acid position 177 in AHSV VP7mt144/200 (Riley, 2003). All PCR reactions were carried out using TaKaRa Ex Taq™ polymerase (Life Technologies) and consisted of 10 ng template DNA, 100 pmol of each primer (table 2.1), 3 µl 2.5 mM dNTP mix, 5 µl 10 x TaKaRa Ex Taq™ polymerase buffer (250 mM TAPS pH 9.3, 500 mM KCl, 20 mM MgCl₂, 10 mM 2-mercaptoethanol), 5 units (U) TaKaRa Ex Taq™ polymerase enzyme and made up to a final volume of 50 µl with ddH₂O. A GeneAmp PCR System 9600 (Perkin Elmer) was used for the PCR reactions.

The amplification conditions for both the reaction were as follows: 1 cycle at 94°C for 2 min; 25 cycles 93°C for 45 sec, 63°C for 1 min, 72°C for 2 min; 1 cycle at 72°C for 10 min.

2.2.2. Agarose Gel Electrophoresis

DNA samples were analyzed by the electrophoretic separation of DNA in agarose gels containing 0.5 µg/ml ethidium bromide. 1% - 4% agarose gels were used depending on the sizes of the fragments to be separated. TAE buffer (40 mM Tris-HCl, 20 mM NaAc, 1 mM EDTA, pH 8) was used as buffer
and in the making of the gel. The samples were mixed with a loading buffer (1% bromophenol blue, 1% xylene cyanol, 1xTAE buffer, 50% glycerol) prior to loading on the gel. Results were visualized by UV fluorescence.

Table 2.1. Primer sequences used in PCR site-directed mutagenesis of VP7mt144/200

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2.2.3. Purification of Amplified DNA Fragments

DNA fragments of interest were purified via the High Pure™ PCR Product Purification Kit (Roche Diagnostics)

2.2.4. Restriction Enzyme Digestion

The restriction endonucleases that were used in the cloning procedure during the preparation of DNA fragments or in the characterization and identification of recombinant DNA plasmids were used according to the specifications of the manufacturers (Roche Diagnostics).
2.2.5. Dephosphorylation

The digested pFastBac plasmid was dephosphorylated by incubating the linearised DNA in a total reaction volume of 20 μl in the presence of 0.5 U of calf intestinal alkaline phosphatase and 2 U of 10 x dephosphorylation buffer (0.5 M Tris-HCl, 1 mM EDTA, pH 8.5) at 37 °C for 20 min.

2.2.6. Ligation

Sticky-blunt end ligations were carried out at 12 °C for 16 h. A high, 10:1, ratio of insert: vector molecules was used in a total volume of 20 μl, which also contained 2 U of T4 DNA ligase and 2 U of 10 x ligation buffer (660 mM Tris-HCl, 50 mM MgCl2, 10 mM dithio-erythritol, 10 mM ATP, pH 7.5).

2.2.7. Preparation of Competent *E.coli* Cells

The standard CaCl2 method described by Cohen *et al* (1972) was used to prepare the *E.coli* XI1 Blue competent cells used in all pFastBac plasmid transformations. 1 ml of an overnight XI1 Blue culture was used to inoculate 100 ml Lauria-Bertani (LB) medium (1% NaCl (w/v), 1% tryptone (w/v), 0.5% yeast (w/v)). Cells were grown at 37°C with shaking to logarithmic (log) phase (OD550 = 0.5). The cells were collected by centrifugation at 5000 rpm for 5 min at 4°C. The pellet was then gently resuspended in 50 ml of freshly made ice-cold 50 mM CaCl2, centrifuged again at the above-mentioned conditions and resuspended in 5ml of 50 mM CaCl2. The cells were kept on ice for 1 h before sterile 15% (v/v) glycerol was added and the cells frozen away at −70°C.

2.2.8. Transformation of Competent Cells

The ligation mixture was added to 100 μl of competent cells and allowed to absorb for 30 min on ice. The mixture was then subjected to a 90 sec heat shock at 42°C and left to cool on ice for 2 min. 900 μl LB was added, followed by an incubation period of 1 h at 37°C with shaking. Aliquots of 100 μl were then plated out on LB agar plates (1.2% agar in LB medium) containing 12.5 μg/ml tetracycline hydrochloride (tet) and 100 μg/ml ampicillin (amp). Plates were then incubated overnight at 37°C.
2.2.9. Plasmid DNA Isolation and Purification

The Birnboim and Doly (1979) alkaline lysis method was used for the isolation of plasmid DNA. Selected colonies from the agar plates were inoculated into 5ml of LB medium, containing the appropriate antibiotics (amp and tet), and grown for 16 h at 37°C with shaking. 3ml of each culture was harvested by bench top centrifugation at 15000 rpm for 1 min after which the pellets were resuspended in 100μl of solution 1 (50 mM glucose, 10 mM EDTA, 25 mM Tris, pH 8) and incubated at room temperature for 5 min. Cell lysis and chromosomal DNA denaturation were achieved by adding 200μl of freshly prepared alkaline-SDS buffer (0.2N NaOH, 1% SDS), followed by a 5 min incubation on ice. 150 μl cold 3 M NaAc (pH 4.8) was added resulting in the reannealing of the plasmid DNA and the precipitation of the genomic DNA, proteins and high molecular weight RNA. After a 10 min incubation period on ice, the supernatant, containing the plasmid DNA, was collected after centrifugation at 15000 rpm for 10 min at 4°C. Two volumes of 96% ethanol was added to the supernatants and incubated at -20°C for 30 min to precipitate the plasmid DNA. This was followed by 10 min centrifugation at 15000 rpm and subsequent washing of the DNA pellet with 80% ethanol. The clean DNA pellet was resuspended in UHQ.

2.2.10. Nucleotide Sequence Determination

Automated nucleotide sequence determination was done using the ABI PRISM™ Big Dye Terminator Cycle Sequencing Ready Reaction Kit (Perkin Elmer). Reactions consisted of 250-500 ng double-stranded plasmid template DNA, 3.2 pmol primer, 4 μl Terminator Ready Reaction Mix and were made up to a total volume of 10 μl with ddH2O. The subsequent PCR reaction was done in a GeneAmp PCR System 9600 (Perkin Elmer) with heated lid and consisted of 25 cycles under the following conditions: 10 sec at 96°C, 45 sec at 50°C and 4 min at °C.

Ethanol precipitation was done to remove unincorporated dye terminators and salts. A pre-made precipitation mix (50 μl 99.9% EtOH, 10 μl ddH2O and 2 μl 3 M NaAc pH 4.8) was added to the 10 μl sequence reaction and incubated on ice for 10 min. This was followed by a 20 min centrifugation step at 13200 rpm. The pellet was washed with 250 μl 70% EtOH, 25°C, centrifuged again for 15 min at 13200 rpm and dried. Samples were then resuspended in 3 μl sequencing loading buffer (5:1 ration of deionised formamide to 25 mM EDTA pH 8 containing 50 mg/ml dextran blue) before denaturation at 95°C for 2 min, followed by cooling on ice. Sample analyses was done on a ABI PRISM 377...
sequencer by loading 1.5 µl of the prepared reactions on a 4% denaturing polyacrylamide gel and running the gel for 7 h at 1.6 kV. Sequences were analysed by using the ABI PRISM Sequencing Analysis™ and Sequence Navigator™ programs.

2.2.11. Hydropathy Predictions

The ANTHEPROT package (Geourjon et al., 1991; Geourjon and Deleage, 1995) and the Hopp and Woods predictive method (Hopp and Woods, 1981; Hopp and Woods, 1983) were used to calculate the hydropathy plots of the VP7 mutants.

2.2.12. Cells and Media

*Spodoptera frugiperda* cell clone Sf9 was used to propagate wild type and recombinant baculoviruses. The cells were grown in suspension culture consisting of Grace’s medium supplemented with 10% foetal calf serum and antibiotics (penicillin, streptomycin, fungizone) at 27°C.

2.2.13. Transposition

In order to transpose the newly synthesized VP7 recombinants, the recombinant pFastBacs were transformed into *E.coli* DH10Bac™ cells (Life Technologies). These cells contain the bacmid and the helper plasmid (Bac-to-Bac™ Expression system Manual) necessary for the transposition reaction to occur. The DMSO method described by Chung and Miller (1988) was used to make the *E.coli* DH10Bac cells competent. 100 ml LB medium was inoculated using 1 ml from a 5 ml overnight culture of DH10bac cells. This new culture was grown to early log phase (OD$_{550}$ = 0.5). Cells were collected by centrifugation at 5000 rpm for 5 min at 4°C. The pellet was then gently resuspended in 10 ml of ice-cold TSB (0.5% NaCl (w/v), 1.6% tryptone (w/v), 1% yeast (w/v), 10% PEG (w/v), 5% DMSO (w/v), 10 mM MgCl$_2$, 10 mM MgSO$_4$) and incubated for 20 min on ice before transposition.

Recombinant pFastBac plasmid DNA was added to 200 µl of the competent DH10Bac cells and incubated on ice for 30 min. TSBG (TSB plus 20 mM glucose) was added followed by an incubation period at 37°C for 4 h and plating out on agar plates containing Kanamycin (50 µg/ml), Gentamycin (7 µg/ml), Tetracycline (10 µg/ml), isopropyl-β-D-thiogalactopyranoside (IPTG, 40 µg/ml) and 5-bromo-4chloro-3indolyl-β-D-galactopyranoside (X-gal, 300 µg/ml). The plates were incubated for 48 h after which the white colonies were inoculated into 5ml LB medium.

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2.2.14. Isolation of Composite Bacmid DNA

The isolation of composite bacmid DNA proceeded according to the Baculovirus Expression System Manual (Life Technologies). The cells were collected from the overnight cultures by centrifugation and resuspended in Solution 1. This was followed by the addition of alkaline-SDS buffer and incubation at room temperature for 5 min. Cold 2.5 M NaAc, pH 5.5, was added and samples were left on ice for 10 min. After centrifugation the DNA was precipitated with isopropanol and incubated on ice for 10 min. The DNA pellet was washed with 80% ethanol and air-dried for 10 min, before being dissolved in sterile UHQ.

2.2.15. Transfection

Six well (35 mm) tissue culture plates were seeded with Sf9 cells at 1.0x10^6 cells per well in 2 ml Graces medium with antibiotics, after having been grown in a 75 ml suspension culture. The cells were allowed to seed for 1 h at room temperature. For each transfection, a solution was made by the addition of the bacmid minipreparation DNA to Graces medium, containing no antibiotics or serum. A second separate solution was prepared by the addition of 6µl of Cellfectin™ reagent (Life Technologies) to Graces medium, also containing no antibiotics or serum. The two solutions were combined and incubated at room temperature for a 45 min time period. Each seeded well was washed twice with 2 ml of Graces medium containing no serum or antibiotics. Graces medium (without serum or antibiotics) was added to the combined solutions A and B to a final volume of 1 ml. The wash medium was removed from the wells and the cells were overlaid with the lipid-DNA complexes formed within the combined solution A and B. The six well plates were incubated for a period of 5 h at 27°C, followed by the removal of the transfection mixture and the addition of Graces medium containing serum and antibiotics. The cells were then incubated in a moist environment at 27°C for a further 96 h. The medium containing the recombinant virus was removed and stored at 4°C until further use.

2.2.16. Large-Scale Recombinant Protein Expression and Isolation

75 cm³ Tissue culture flasks were seeded at 1.0x10^7 with S. frugiperda cells before being infected with the appropriate recombinant baculovirus at a M.O.I. of 5-10 pfu/cell. After a 72 h incubation period, the infected cells were harvested by centrifugation at 3000 rpm for 5 min at 4°C. The pellet was washed with 1 x phosphate buffered saline (PBS) (137 mM NaCl, 2.7 mM KCl, 4.3 mM
Na$_2$HPO$_4$.2H$_2$O, 1.4 mM KH$_2$PO$_4$, pH 7.3), resuspended in 1 ml TNN lysis buffer (50 mM Tris-HCl pH 8, 150 mM NaCl, 0.5% Nonidet P-40) and incubated for 30 min on ice.

2.2.17. SDS-Polyacrylamide Gel Electrophoresis

A 15% SDS-PAGE gels were made as described by Laemmli (1970). The separating gels consisted of 0.375M Tris-HCl pH 8.8, 0.1% SDS, 30% acrylamide and 0.8% bisacrylamide. Polymerisation of the gels occurred by the addition of 0.008% (v/v) tetra-methyl-ethylene-diamine (TEMED) and 0.08% (w/v) ammonium peroxysulfate. The stacking gel contained the same except for 0.125 M Tris-HCl pH 6.8 and 0.1% SDS. TGS buffer (0.025 M Tris-HCl pH 8.3, 0.192 M glycine, 0.1% SDS) was used for the protein electrophoresis which occurred using a Vertical Slab Gel SE 400 unit (Hoefer Scientific Instruments). The harvested Sf9 cells containing the protein samples were added to equal volumes of 2 x protein solvent buffer ((PSB) 0.125 M Tris-HCl pH 6.8, 4% SDS, 20% glycerol, 10% 2-mercaptoethanol) and denatured for 3 minutes at 95°C, before loading on the gel. The gel was run for a period of 16 – 20 h at 120V.

2.2.18. Coomassie Brilliant Blue Staining

After electrophoresis, the protein bands were visualised, using Coomassie brilliant blue stain (0.125% Coomassie brilliant blue, 50% methanol) for a period of 20 minutes. Progressive destaining occurred by placing the stained gel in a solution containing 5% ethanol and 5% acetic acid until only the minimum background remained. The gel was then stored in H$_2$O.

2.2.19. Sucrose Gradient Studies

One fifth to half of the whole cell lysates resuspended in TNN lysis buffer (Section 2.2.16.) were loaded onto a 50%-70% discontinuous sucrose gradient (sucrose w/v, 50 mM Tris-HCl, 150 mM NaCl, pH 8). The samples were centrifuged at 12000 rpm for 75 min at 4°C, with a Sorvall AH-650 rotor. Fractions, consisting of 20 drops per fraction, were collected from the bottom of the gradient, resulting in 11 - 12 fractions depending on the temperature of the sucrose. The pellet was resuspended in 200 µl PBS. Protein content was precipitated from a sample consisting of 200 µl sucrose fractions by the addition 1200 µl PBS, followed by 45 min of centrifugation at 5000 rpm at 4°C. Quantification of the protein content was done by the Sigma Gel™ software program (Jandel Scientific) which measure the relative band intensities on a SDS-PAGE gel.
2.2.20. Density Analysis

The approximate densities of the proteins were calculated using Density Marker Beads (Pharmacia Biotechnology). 500 µl of the whole cell lysate was loaded onto a 50%-70% discontinuous sucrose gradient. Another 50%-70% discontinuous sucrose gradient was loaded with Density Marker Beads before undergoing isopycnic centrifugation: 40000 rpm for 18 h at 4°C, using a Beckmann SW55 TI rotor. The sucrose gradient containing the lysate was fractionated and analysed as described in section 2.2.19. The gradient containing the Density Marker Beads was used as density measurement reference.

2.2.21. Sonication

Protein aggregation disruption was studied with a Sonifier® Cell Disrupter B-30 unit (Branson Sonic Power Company). The sucrose gradient aggregated proteins in the pellet of a preceding sucrose gradient, suspended in PBS, were mostly subjected to 25 pulses of 20% pulsed duty cycle at 4000 Hz with a 1 min cooling-down period after every 5 pulses. The samples were then subjected to further solubility studies (2.2.19) before being analysed on a SDS-PAGE gel (2.2.17).

2.2.22. Trimerization Assay

Trimerization assays were conducted by first purifying the proteins according to the method described in section 2.2.19. The second and third fractions were pooled before undergoing 25 pulse sonication. The sonicated, unboiled protein samples were mixed with 10% glycerol and analysed on a 10% SDS-PAGE gel (2.2.17).

2.2.23. Storage Stability Studies

Sucrose gradient purified proteins, suspended in ddH₂O, were subjected to different storage conditions. This included flash-freezing the samples in liquid N₂, overnight storage at 4°C, and freeze drying the samples in a Freezemobile Twin 6 (Virtis) until. Flash-freezed samples were defrosted on ice and freeze-dried samples were re-hydrated by the addition of ddH₂O before subsequent solubility studies (2.2.19) were performed. This was followed by analysis on a SDS-PAGE gel (2.2.17).
The second PCR reaction amplified the second segment, consisting of the nucleotides from amino acid position 177 up to the 3' end of the VP7mt144/200 construct template. In this reaction primers amplified the segment with the addition of Stul and BssHII sites to the 5' end and a KpnI site to the 3' end of the segment (Table 2.1.). The amplified segment, as well as the pFastBac-VP7mt144/177 construct, underwent separate Stul and KpnI digestions. The cleaned PCR product was then ligated into the prepared pFastBac-VP7mt144/177 construct to create the final three site AHSV VP7 construct; VP7mt144/177/200, size 5995 bp.

**Figure 2.3.** Primer binding positions and PCR amplification of the two segments cloned into pFastBac (Fig. 2.4).
2.2.24. Scanning Electron Microscopy (S. E. M.)

Purified recombinant VP7 crystals were resuspended and left for 15 min in a fixative solution (0.15 M Na₂HPO₄, 0.1% glutaraldehyde) before being filtered through a 0.22 μm nylon filter. Three subsequent washing steps with 0.075 M Na₂HPO₄ followed with a 10 min waiting step between each. The crystal covered filters were then dehydrated for 15 min consecutively in 50%, 70%, 90% and three times in 100% ethanol. Critical point drying with liquid CO₂ further dehydrated the samples before being placed on an aluminium stub and spatter coated with gold-beladium in a S. E. M. autocoating unit E5200 (Polaron Equipment Ltd.). Samples were then viewed in a Jeol 840 S. E. M.

2.3. Results

The recombinant VP7 construct, VP7mt144/200 (Riley 2003), as provided by Me. J. E. Riley, served as template for the creation of a three site construct, designated VP7mt144/177/200. The site between amino acid 177 and 178 of AHHSV-9 was chosen as a target for the insertion of restriction endonuclease sites, since, as stated before, this is in the middle of a the highly flexible RGD containing loop. Restriction endonuclease sites PstI (codes for Leu and Gln), StuI (codes for Arg and Pro) and BssHI (codes for Ala and Arg) were inserted. Fifty percent of the inserted amino acids into the hydrophilic loop are hydrophobic and could possibly affect the structure of the loop and subsequently the trimer-forming abilities of these recombinant VP7 molecules.

2.3.1. Construction of Cloning Site 177

A two-step cloning procedure, as summarised in figures 2.3. and 2.4., was required for the creation of the VP7mt144/177/200 construct. In the first step a PCR reaction was used to amplify the first segment that stretches from the 5’ start of the gene up to amino acid position 177, using the VP7mt144/200 construct as template. The primers were designed to amplify the fragment adding a 5’ BgII overhang and a 3’ PstI and StuI overhang, as shown in Table 2.1. The obtained PCR product was subjected to a double separate digestion with the BgII and StuI restriction endonucleases and was cleaned before being cloned directly into the prepared pFastBac plasmid vector, between the BamHI and StuI sites. The product’s BgIII site is an isoschizomere of pFastBac’s BamHI site, thus binding to it and simultaneously destroying both the sites. This intermediate construct, size 5335 bp, was designated pFastBac-VP7mt144/177.
Figure 2.4. Construction of pFastBac-VP7mt144/177/200
2.3.2. Nucleotide Sequence Determination and Hydropathy Predictions

The primers listed in table 2.1. were used during automated DNA sequencing (section 2.2.10) to determine the nucleotide sequence of the newly constructed VP7mt144/177/200. Overlapping sequences were generated from which the complete gene sequence could be deduced. The sequencing not only confirmed the insertion of the of the 18 nucleotides constituting the three restriction endonuclease sites between amino acid position 177 and 178, but also showed the presence of three restriction endonuclease sites at both amino acid site 144 and 200, acquired from the VP7mt144/200 construct (Appendix A).

The derived sequence also supported the accuracy with which the PCR reaction was performed. No new mutations were generated with the amplification of the two fragments from the VP7mt144/200 template. It should, however, be noted that during the construction of the VP7mt144/200 construct a mutation at nucleotide position 30/31 occurred. This resulted in a CG sequence being switched to a GC
sequence and a subsequent change in amino acid 5 from wild-type arginine to alanine. As discussed in Riley 2003, this mutation should have little to no affect on the protein’s structural formation.

The sequences were aligned and compared using ClustalX version 1.81 (Higgins and Sharp, 1988; Higgins et al., 1996), shown in Appendix A. Subsequently, the obtained sequence was further analysed using the ANTHEPROT package (Geourjon et al., 1991; Geourjon and Deleage, 1995) to calculate the hydropathy plots using the Hopp and Woods predictive method (Fig. 2.6) (Hopp and Woods, 1981; Hopp and Woods, 1983). The hydrophilic character of each amino acid is based on the property of its side chain to be more soluble in water than in an apolar solvent. As can be seen in figure 2.6 (a), VP7 is a highly hydrophobic protein. There are only a few hydrophilic regions, which are exposed on the surface of the folded VP7 trimeric structures and on the surface of the hexagonal crystals. The insertion of six amino acids, leu-gln-arg-pro-ala-arg, at the 177/178 amino acid site significantly increased the hydrophilicity of this particular region. This is expected since when added together, the residual hydrophilic character of the six amino acids equals +2.5 (Table 2.2). It should be noted the increase in hydrophilicity is located at the C-terminal of the insert, the contribution of the strong hydrophilic arginines. An even larger increase in hydrophilicity is shown at the 200/201 amino acid site. Only two of the inserted amino acids (leu and val) are hydrophobic and the hydrophilicity is more evenly spread over the region, more efficiently cancelling the effect of these two hydrophobic amino acids. There is, however, an overall slight increase in hydrophobicity at the 144/145 site. This effect is mostly generated by the amino acids at the N-terminal of the insert. The overall impact these changes have on the structure of the protein could, at this stage, not be predicted.
Table 2.2. Hydrophilic character of the amino acids added to VP7 (Hopp and Woods, 1981; Hopp and Woods, 1983).

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Figure 2.6. Hydrophilicity Plots of (a) VP7 and (b) VP7mt144/177/200 according to the Hopp and Woods Predictive method (Hopp and Woods, 1981; Hopp and Woods, 1983.).
2.3.3. Baculovirus Expression

The Bac-to-Bac™ expression system was used for the expression of the all the proteins. *E. coli* DH10Bac™ cells, containing the baculovirus genome, were transformed with the recombinant pFastBac transfer vector containing the recombinant VP7 genes (2.2.13). During the four-hour growth period, transposition of the recombinant gene of interest occurred from the pFastBac plasmid into the genome of the baculovirus. The cells were plated out on enriched agar plates containing X-gal and IPTG, which enabled blue/white selection of positively transposed colonies containing recombinant bacmid DNA. The recombinant bacmid DNA was subsequently isolated (2.2.14) and transfected into Sf9 insect cells (2.2.15) for the generation of recombinant baculovirus expressing the gene of interest. Recombinant baculovirus was recovered and used to prepare viral stocks for subsequent viral infections, which occurred at a M.O.I. of 5-10 pfu/cell.

Expression of the proteins of interest was verified by means of SDS-PAGE gel electrophoresis (2.2.17). The effect of the 18 amino acid size increase of the expressed recombinant protein (39 kDa) can be clearly noted when compared with wild-type VP7 (37 kDa) on a 15% SDS-PAGE gel (Fig. 2.7). No significant changes to the expression levels, which might have been caused by the site directed mutations in the recombinant protein, can be detected.

![Figure 2.7. A 15% SDS-PAGE analysis of VP7mt144/177/200. Lane 1 represents the protein size marker. Lane 2 and 3 contain proteins from a mock infection and wild-type baculovirus-infected cells respectively. Lane 4 contains expressed VP7 and lane 5 contains the lysate of cells infected with baculovirus expressing the recombinant VP7mt144/177/200.](image)
2.3.4. Solubility and the Effect of Sonication on VP7 Aggregation

Solubility studies were performed to investigate whether the hydrophilic changes in the modified VP7mt144/177/200 mutant, as discussed in 2.3.2., indeed has any significant contribution to the destabilisation of its crystal forming abilities. To study the effect the modification had on solubility, SF9 cells were infected respectively with recombinant baculovirus stocks expressing VP7 and VP7mt144/177/200. After a 72 h incubation period, post-infected cells were harvested and lysed. Whole cell lysates were loaded on a 50%-70% discontinuous sucrose gradient and centrifuged as described in 2.2.19. Fractions were collected from the bottom of the gradient, resulting in 11 - 12 fractions. All particulate protein structures were selected and collected by means of the centrifugation conditions described in section 2.2.19, ensuring purity of the VP7 recombinant proteins, if not quantity. A sample of each fraction was SDS-PAGE gel (2.2.19). The relative protein band intensities on the SDS-PAGE gels were measured and quantified as percentages of the total specified protein content across all the fractions by the Sigma Gel™ software program (Jandel Scientific). The experiments were repeated several times to exclude inherent possibilities of variation.

Figure 2.10 shows the different protein particle distribution of the VP7 and the VP7mt144/177/200 constructs. The majority of the VP7 and VP7mt144/177/200 proteins are recovered from the pellet of the sucrose gradient. It should be noted though that approximately 56% of the VP7mt144/177/200 proteins (Fig. 2.10 b) are located in the pellet, whereas only approximately 42% of the VP7 proteins (Fig. 2.10 a) are located in the pellet. The free VP7 crystal portion peak in the second fraction from the bottom of the sucrose gradient, with decreased amounts of smaller particulate structures in the middle section and finally increasing amounts of the smallest particles, possibly soluble free VP7 trimers, in the top part of the gradient. The free VP7mt144/177/200 protein particles, however, only peak in the third fraction from the bottom of the sucrose gradient, with a decreasing amount of smaller particulate structures in the middle section of the gradient. There is no real indication of soluble, small particles or free trimers in the hydrophilic top part of the VP7mt144/177/200 sucrose gradient.

VP7mt144/177/200 proteins were also subjected to isopycnic centrifugation along with Density Marker Beads (2.2.20). The results, as presented in figure 2.9, clearly show that the VP7mt144/177/200 proteins actually occur in the first three fractions of the 50%-70% sucrose gradient when density centrifugation, and not size selection, is applied. The proteins in these three fractions, when compared to the Density Marker Beads, correspond to an approximate buoyant density of 1.108 g/ml to 1.133 g/ml.
The pellets of the sucrose gradients from both the constructs were subjected to sonication. This was aimed at disrupting the purified aggregates in the pellet and increasing the yield of purified protein that might be used for vaccination purposes (2.2.21). The samples underwent 25 pulses of sonication before, again, being loaded on a 50%-70% discontinuous sucrose gradient for further solubility studies (2.2.19). The attained results were compared with the construct’s sucrose gradient in the absence of sonication. As can be seen in figure 2.11 a, VP7 pellet sonication disrupts the aggregation between the crystals, resulting in an overall increase, when compared to unsonicated VP7, in the percentage proteins contained in each fraction. An intense peak of protein crystals is noted in the third fraction from the bottom. This fraction seems to constitute approximately 44.8% of the total amount of proteins that were loaded on the gradient, whereas this fraction only constitutes approximately 12% of the total protein content in the absence of sonication. There is a rapid decline in the percentage protein located in the fourth fraction. A final increase in the ninth to eleventh fractions, peaking at fraction nine, illustrates the presence of soluble free and aggregated trimeric VP7 structures in the soluble fraction of the sucrose gradient. The pellet of this sonicated VP7 sample still contains approximately 5% of the total protein start content.

The results attained from VP7mt144/177/200 in the absence of sonication compared to sonicated VP7mt144/177/200 are illustrated in figure 2.11 b. Sonicated Vp7mt144/177/200 protein particles have the same sedimentation profile as unsonicated VP7mt144/177/200, although the percentage particles in each fraction is higher in the absence of a pellet. Also, there is an increase in smaller particles when sonicated, as illustrated by the presence of the proteins in fractions eight and nine. This tendency to revert from large aggregated protein masses to smaller particles was further investigated by the sonication of additional samples of VP7mt144/177/200 at more than 25 pulses. Figure 2.12 shows the results of stability tests that were done on VP7mt144/177/200 after no sonication, 25 pulses, 50 pulses, 100 pulses and finally, exhaustive sonication (approximately 1900 pulses). A clear shift from large particulate structures at low sonication to more soluble small particles or trimeric structures at high sonication can be observed.

2.3.5. Trimerization Assays

A trimerization assay to confirm the formation of trimeric structure by VP7mt144/177/200 was performed similar to that described by Limn et al., 2000. Large particulate structures were sonicated to reduce the size of the assembled particles. These unboiled samples were loaded unto a 10% SDS
PAGE gel for analysis (Section 2.2.22). It can be seen that there is a greater tendency by VP7mt144/177/200 to form free monomers (39 kDa), even when non-denaturing conditions are being applied (Fig. 2.8). An extremely faint band of 117 kDa can be detected indicating the presence of the trimeric VP7mt144/177/200.

2.3.6. Storage Stability Studies

Cold-chain storage of vaccines cannot always be maintained, since it is expensive and not very practical, especially in third world countries. It is thus important to test the effects different storage conditions have on the stability of VP7mt144/177/200, if it is to be used as a subunit vaccine. The second and the third fractions of VP7mt144/177/200 sucrose gradients were pooled and subjected to three different storage methods. This included flash-freezing the samples in liquid N₂, overnight storage at 4°C, and freeze drying the samples (2.2.23). After subsequent sample preparation, the samples were subjected to sucrose gradient sedimentation analyses (2.2.19). The results, in figure 2.13, were compared to control VP7mt144/177/200 proteins that were fractionated on sucrose gradients directly after cell lysis. At the bottom of the gradient, the stored samples have the same protein distribution in each fraction as the control, but only at much lower levels. Towards the top of the gradient, the stored samples have a higher percentage of small particles in the soluble fraction than the control. The most conspicuous result is the high levels of VP7mt144/177/200 protein that was found in the pellet of the stored samples. Where the control have approximately 56% of the total protein amount in the pellet, the stored samples, that were taken from the second and third fractions of a sucrose gradient, have higher portions of their total protein content in the pellet. The freeze-dried sample has approximately 53% in the pellet, the flash-freezed sample has approximately 68% in the pellet, with approximately 80% of the sample that was left at 4°C occurring in the pellet.

2.3.7. Scanning Electron Microscopy (S.E.M.)

Protein samples collected from the second and third fractions were pooled, fixed, mounted and spatter coated with gold-beladium particles for investigation under the S.E.M. Unmodified VP7 protein crystals were used as control. The wild-type VP7 crystals appeared exactly as first described by Chuma et al., 1992 (Fig. 2.14 a and b). They are flat, smooth, hexagonal structures with a mean size of approximately 8 μm² in diameter. The structures formed by VP7mt144/177/200 proteins seem to be concentration dependent. The structures formed by a low concentration of the recombinant VP7mt144/177/200 protein have a rough, flat and an almost circular appearance (Fig. 2.14 c and d).
The units are approximately the same size as the 8 μm crystals formed by wild-type VP7. When high concentrations of VP7mt144/177/200 proteins are viewed, the units seem decompressed and have a mean size of approximately 10 μm in diameter (Fig. 2.14 e). The units are connected with web-like tentacles/ cables (Fig. 2.14 f). This study was extended by using VP7mt144/177/200 freeze dried samples for S.E.M. study (Fig. 2.15 a, b and c). The units from this sample had the same structure and texture as high concentration samples that did not undergo the drying process.

A further study was conducted by S.E.M. examination of sonicated wild-type VP7 samples (Fig. 2.15 e and f) and sonicated VP7mt144/177/200 samples (Fig. 2.15 d). Twenty-five pulses of sonication reduced the VP7 crystals to large fragments of the hexagonal crystals, with chance whole hexagonal crystal units still to be found. The same amount of sonication pulses on the VP7mt144/177/200 samples reduced the VP7mt144/177/200 large units (as described above) to small fragments of nondescript shape and a mean size of approximately 1 μm. No web-like tentacles/ cables were observed between the small fragments.

![Image of gel electrophoresis](image)

**Figure 2.8.** Trimerization assay of VP7mt144/177/200 analysed on a 10% SDS-PAGE gel. Lane 1 represents the protein size marker. Lane 2 to 8 contain purified VP7mt144/177/200 proteins that underwent 25 pulse sonification before being loaded unboiled onto the gel.
Figure 2.9. Density Marker Beads (a) and protein distribution of VP7mt144/177/200 (b) on a 50%-70% sucrose density gradient using a Beckmann SW55 TI rotor at 40000 rpm for 18 hours at 4°C. The VP7mt144/177/200 protein distribution was analyzed by SDS-PAGE electrophoresis, quantified by the Sigma Gel™ analysis program and converted to the shown graphic form. On the SDS-PAGE gels, lane 1 contains the protein size marker, whereas lanes 2-13 contain the fractions as taped from the bottom to the top of the gradients. Lane 14 represents the protein pellet of the gradient.

Measurements of the Density Marker Beads’ locations were taken and converted into a graph (a). The highlighted yellow area indicates the approximate density of the VP7mt144/177/200 structures.
Figure 2.10 (a), (b). Protein distribution of VP7 (a) and VP7mt144/177/200 (b) on a 50%-70% sucrose gradient using a Sorvall AH-650 rotor at 12000 rpm for 75 min at 4°C, analyzed by SDS-PAGE electrophoresis, quantified by the Sigma Gel™ analysis program and converted to the shown graphic form. On the SDS-PAGE gels, lane 1 contains the protein size marker, whereas lanes 2-13 contain the fractions as collected from the bottom to the top of the gradients. Lane 14 represents the protein pellet of the gradient.
Figure 2.11 (a), (b). Protein distribution of 25 pulse sonicated VP7 (a) and sonicated VP7mt144/177/200 (b) each with a respective unsonicated control. Proteins were centrifuged on a 50%-70% sucrose gradient using a Sorvall AH-650 rotor at 12000 rpm for 75 min at 4°C, analyzed by SDS-PAGE electrophoresis, quantified by the Sigma Gel™ analysis program and converted to the shown graphic form. On the SDS-PAGE gels, lane 1 contains the protein size marker, whereas lanes 2-13 contain the fractions as collected from the bottom to the top of the gradients. Lane 14 represents the protein pellet of the gradient.
Figure 2.12. Protein distribution of sonication range of VP7mt144/177/200 pellets. Samples were centrifuged on a 50%-70% sucrose gradient using a Sorvall AH-650 rotor at 12000 rpm for 75 min at 4°C, analyzed by SDS-PAGE electrophoresis, quantified by the Sigma Gel™ analysis program and converted to the shown graphic form. On the SDS-PAGE gels, lanes 1-12 contain the fractions as collected from the bottom to the top of the gradients.
Figure 2.13. Protein distribution of sucrose gradient purified, unsonicated VP7mt144/177/200 samples stored at different conditions. Samples were centrifuged on a 50%-70% sucrose gradient using a Sorvall AH-650 rotor at 12000 rpm for 75 min at 4°C, analyzed by SDS-PAGE electrophoresis, quantified by the Sigma Gel™ analysis program and converted to the shown graphic form. On the SDS-PAGE gels, lanes 1-12 contain the fractions as collected from the bottom to the top of the gradients.
Figure 2.14. Scanning Electron Microscopy photographs depicting the structures formed by the different protein constructs. (a) and (b) show the hexagonal crystals formed by wild-type VP7. The structures formed by VP7mt144/177/200 at low concentrations are shown in figure (c) and (d), whereas (e) depicts a structure at high concentration, with a wide-angle view depicted in (f).
Figure 2.15. Scanning Electron Microscopy photographs depicting the structures formed by the different protein constructs under different conditions. (a) and (b) show VP7mt144/177/200 structures that had been freeze dried, with a wide-angle view depicted in (c). A wide-angle view of sonicated VP7mt144/177/200 is shown in (d). (e) depicts remaining fragments of VP7 crystals after sonication, with a wide-angle view shown in (f)
2.4. Discussion

African horse sickness virus VP7 monomers are highly hydrophobic and three of these monomers tend to spontaneously organise into trimeric structures through non-covalent interactions. These VP7 trimers, when expressed in a recombinant baculovirus, spontaneously assemble in infected insect cells into large, flat, hexagonal crystals (Chuma et al., 1992; Burroughs et al., 1994). These crystalline structures have a highly ordered lattice that is consistent with a trimeric subunit structure. The more hydrophilic BTV VP7, on the other hand, is completely soluble and does not form these crystal structures (Oldfield et al., 1990). Thus it was deduced that the ability of AHSV VP7 trimers to assemble into crystalline structures could probably be ascribed to hydrophobic interactions between the trimers. Therefore, modifications that affect the solubility of VP7, might influence the formation of the particulate structures.

The creation of cloning sites in VP7 protein added, in total, 6 amino acid insertions at each of the three respective sites. These changes in the construction of VP7mt144/177/200 contributed to the overall hydrophilicity of the protein, the impact of which could not be determined at the early stages before solubility studies. The initial solubility studies made a comparison between the solubility of wild-type VP7 and the recombinant VP7mt144/177/200 construct (Fig. 2.10). The VP7 structures, present in the second fraction from the bottom of the gradient, seemed to be larger and more compact than the structures formed by VP7mt144/177/200, which occurred in the third fraction from the bottom. The absence of small particles in the soluble top portion of the VP7mt144/177/200 gradient indicated an increased tendency towards aggregation between the assembled particles. This was further supported by a significant increase in the percentage aggregated proteins in the pellet of the VP7mt144/177/200 sucrose gradient, when compared to VP7. The most likely possible explanation for this phenomena is that due to the structural stress caused by increased hydrophilicity at three key sites in the top domain of the trimers, the hydrophobic regions in the trimers are more exposed. This causes the trimers and the larger structures formed by them, to aggregate more. The results of the storage stability studies, which will be discussed later in this section, embrace this explanation. As mentioned in 2.3.6., the most conspicuous result was the high levels of VP7mt144/177/200 protein that was found in the pellet. The protein samples used in this particular study were pooled from the second and third fractions, thus did not occur initially in the pellet. Extremely high levels of protein aggregation can be correlated with the extend of time that the samples were left in solution. The freeze dried sample had the shortest
period of time for aggregation, therefore the lowest amount of aggregated proteins. The flash-freeze sample had a longer period for aggregation as defrosting occurred. The sample with the highest amount of aggregated proteins, is the sample that was left to stand overnight at 4°C.

Further studies were conducted on the aggregated proteins found in the pellets of the sucrose gradients. Sonication of the pellets showed that VP7 proteins form stable crystals with a distinct size and density, as illustrated by the presence of approximately 44.8% of the proteins in a single fraction (Fig 2.11 a). This observation was further supported by almost no significant increase in the intermediate particulate structures found in the middle fractions, except for an increase in the proportion of small particulate structures at the top of the gradient. The sonication product of the VP7mt144/177/200 construct resembles the right side of a Gaussian distribution curve. Structures of varying size and density are found with an increase in the soluble trimer proportion with longer sonication periods. This not only indicates instability in the VP7mt144/177/200 construct that is not found in the wild-type VP7, but also that no distinct units are formed anymore.

Scanning electron microscopy confirmed the solubility studies. The VP7mt144/177/200 structures are not smaller than crystals that are formed by wild-type VP7, as thought earlier. The VP7mt144/177/200 structures vary in shape and size, which might account for the difference in distribution during rate zonal centrifugation; than the smooth, flat hexagonal VP7"crystals moving faster through the gradient and reaching density equilibrium sooner. Higher concentrations of VP7mt144/177/200 protein units are connected with web-like tentacles/ cables. These connecting web-like structures are probably the products of aggregation between the units. It also seem that the units themselves are composed of these web-like tentacles/ cables - contradictory to the smooth, hexagonal crystals produced by wild-type VP7 and the structures formed with low concentrations of VP7mt144/177/200. This might be due to aggregational stress that is exerted on these structures, thus physically pulling them apart. All these changes can be ascribed to the changes in solubility, leading to structural stress and the changing of the trimeric unit.

Meyer, 2002, also conducted research on the presentation of neutralising epitopes on the surface of AH SV VP7. A VP7 construct was created with a single multiple cloning site at amino acid position 177. A triple repeat of a single epitope was inserted into this multiple cloning site. In total, twenty-two amino acids were inserted into position 177 with a net hydrophilic character of +8.8 (Hopp and Woods, 1981; Hopp and Woods, 1983). In these studies no distortion of the hexagonal crystalline structure was observed. These results differ from those observed with the VP7mt144/177/200
construct, which only has 18 additional amino acids with a net hydrophilic character of 7.3, as shown in table 2.2 (Hopp and Woods, 1981; Hopp and Woods, 1983). This is probably due to the more “spread-out” distribution of the insertions in the VP7mt144/177/200 construct, in contradiction to the single positional insertion made Meyer.

It is important for vaccination purposes to study the effect of different storage methods on the stability of the vaccine. Not one of the storage methods had a direct effect on the structure of the proteins. It would seem that freeze drying, with the least amount of aggregation, is the best storage method. Studies under the S.E.M. confirmed the absence of any changes to the overall structure of the freeze dried proteins. The only foreseeable problem is the aggregation of the proteins. Sonication might be a solution to the aggregation problem since both the particulate and the individual trimers have potential as a vaccine delivery system. It might even be more advantageous to use fragments and single trimer units since only T-cell mediated immunity was generated and no antibody-based humoral immune response by the injection of whole 6 μm VP7 crystals (Wade-Evans et al., 1998). A possible explanation for the exclusive generation of T-cell mediated immunity might be that the large crystals are taken up by antigen-presenting cells in the body, which breaks down the crystals and only present the epitope regions to the T-helper cells.

Sonication, however, will not stop re-aggregation from occurring. Further studies might have to be conducted in the use of non-polar chemicals for dissolving the dried proteins in before injection.