Molecular Characterization and Detection of Infectious Bronchitis Virus

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Abstract
This thesis deals with the molecular characterization and detection of infectious bronchitis virus (IBV), an important pathogen that causes heavy losses in the poultry populations worldwide. The aim of the research was to better understand the molecular characteristics of the virus and to investigate the factors behind the continuous emergence of new genetic variants and the occurrence of outbreaks. The studies showed that the viral genome is under a continuous evolution, due to mutations, strong selective pressure and recombination events. These forces lead to a wide genetic diversity and the generation of new variants of this virus. The viral genes encoding the spike, replicase and nucleocapsid proteins can be considered the main genomic regions, which are indicating the evolution processes of IBV. The various strains contain specific structural and functional motifs in their genes and the alterations in these motifs may affect the infection biology of the virus. The constant emergence of new variants in Sweden is likely due to the introduction of novel IBV strains from other European countries through the import of poultry products, or by the continuous migration of wild birds.

The in silico investigations of the spike glycoprotein coding regions of the Massachusetts and QX-like genotypes demonstrated molecular differences between these genotypic variants. It is hypothesized that the genetic diversity in the spike gene of IBV and of other avian coronaviruses, as well as of human, bat, and other animal coronaviruses could be associated with the adaptation and host specificity of these infectious agents.

The data obtained by molecular characterization approach was also used for the development of a new molecular method for the improved detection and genotyping of this virus. A microarray platform was developed for the simultaneous detection and rapid typing of IBV variants. This assay provides a practical tool for better diagnosis, for studying the effectiveness of vaccination and for performing large-scale epidemiological studies.

Keywords: Infectious bronchitis virus, IBV, coronavirus, spike glycoprotein, mutation, selective pressure, recombination, genetic diversity, bioinformatics, multiplex, diagnosis, VOCMA, microarray

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Dedication

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and Little Princess Samreen
for their
Endless Love
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List of Publications

This thesis is based on the work contained in the following papers, referred to by Roman numerals in the text:


IV Öhrmalm, C., Abro, S. H., Baule, C., Zohari, S., Bálint, A., Renström, L. H. M., Blomberg, J. and Belák, S. Variation-tolerant capture multiplex assay (VOCMA) for the simultaneous detection of avian infectious bronchitis virus genotypes using multiplex RT-PCR and Luminex technology (Manuscript).

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**Abbreviations**

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<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>AGPT</td>
<td>Agar gel precipitation test</td>
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<tr>
<td>AI</td>
<td>Aliphatic index</td>
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<td>Conn</td>
<td>Connecticut</td>
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<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<tr>
<td>cDNA</td>
<td>Complementary deoxyribonucleic acid</td>
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<tr>
<td>dNS</td>
<td>Non-synonymous substitution</td>
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<td>dS</td>
<td>Synonymous substitution</td>
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<td>E</td>
<td>Envelope</td>
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<td>ELISA</td>
<td>Enzyme-linked immunosorbent assay</td>
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<td>HI</td>
<td>Haemagglutination inhibition</td>
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<td>IBV</td>
<td>Infectious bronchitis virus</td>
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<td>IFA</td>
<td>Immunofluorescence assay</td>
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<td>IPA</td>
<td>Immunoperoxidase assay</td>
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<td>LRR</td>
<td>Leucine-rich repeat</td>
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<tr>
<td>M</td>
<td>Membrane</td>
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<td>Mass</td>
<td>Massachusetts</td>
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<td>mRNA</td>
<td>Messenger RNA</td>
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<td>MFI</td>
<td>Median fluorescence intensity</td>
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<td>N</td>
<td>Nucleocapsid</td>
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<td>NJ</td>
<td>Neighbor-Joining</td>
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<td>Nsp</td>
<td>Non-structural protein</td>
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<tr>
<td>nt</td>
<td>Nucleotide</td>
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<tr>
<td>ORFs</td>
<td>Open reading frames</td>
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<td>PL&lt;sup&gt;pro&lt;/sup&gt;</td>
<td>Papain-like protease</td>
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<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
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<td>RDP</td>
<td>Recombination detection program</td>
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<tr>
<td>RdRp</td>
<td>RNA-dependent RNA polymerase</td>
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<td>RT-PCR</td>
<td>Reverse transcriptase polymerase chain reaction</td>
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<td>Abbreviation</td>
<td>Description</td>
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<td>S</td>
<td>Spike</td>
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<td>SPF</td>
<td>Specific pathogen-free</td>
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<td>sgRNA</td>
<td>Subgenomic RNA</td>
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<td>UTRs</td>
<td>Untranslated regions</td>
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<td>VNT</td>
<td>Virus neutralization test</td>
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<td>VOCMA</td>
<td>Variation-tolerant capture multiplex assay</td>
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<td>VT</td>
<td>Variation-tolerant</td>
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1 Introduction

1.1 History of infectious bronchitis

Infectious bronchitis (IB) is a highly contagious disease of serious economic importance in the poultry industry worldwide. The first report of IB by Schalk and Hawn referred to a highly contagious disease in young chicks with respiratory symptoms in North Dakota, USA in 1931 (Schalk & Hawn, 1931). Pathogenic alterations in the upper respiratory tract of the birds were prominent; hence the disease was named “infectious bronchitis of young chicks”. Five years later, it was demonstrated that the causative agent of this disease is a virus, which was named Infectious Bronchitis Virus (IBV) (Beach & Schalm, 1936). After the initial description of infectious bronchitis, many cases of the disease were reported in the United States (Jungherr et al., 1956; Hitchner et al., 1966; Johanson et al., 1973; Snyder & Marquardt, 1984; Fabricant, 1998; Hitchner, 2004). Thereafter and to date, a wide range of different IBV serotypes and genotypes have been detected around the world (Jackwood et al., 1997; de Wit et al., 2011).

1.2 Clinical features of infectious bronchitis

Clinical cases of IB are associated with respiratory, reproductive, digestive and renal infections in domestic poultry and in various other avian species (Cavanagh, 2005). The disease is clinically manifested by coughing, sneezing, tracheal coarse crackles, nasal discharge, decrease of feed intake and conversion, loss of body weight, swollen sinuses, increased water intake, wet droppings, depression, lethargy and poor growth in broilers. In layers the disease, “false layer syndrome”, affects egg quality (thin, rough, fragile, misshapen egg shells and thin watery egg) and causes decrease in egg production. In some cases, the virus infection may cause severe damage to the
oviduct and result in decreased or permanent loss of egg production (Otsuki et al., 1990; Fabricant, 1998; Cavanagh, 2003; Cavanagh & Naqi, 2003; Cavanagh & Gelb, 2008; Worthington et al., 2008). IB infections may lead to mortality up to 20-30% or higher at five to six weeks of age in chicken flocks (Ignjatovic et al., 2002; Seifi et al., 2010). The mortality can increase due to immunosuppression, mycoplasma and other secondary bacterial infections caused by various accompanying infectious agents, e.g. *Escherichia coli*, *Ornithobacterium rhinotracheale* and *Bordetella avium* (Hopkins & Yoder, 1984; Matthijs et al., 2003; Cavanagh & Gelb, 2008). The mortality rate can be as low as 1% and chickens may recover rapidly, if the infections are produced by mildly virulent strains and are not associated with secondary bacterial infections (Cavanagh & Gelb, 2008).

### 1.3 Host specificity

IBV infects a wide range of avian species, especially those reared close to domesticated poultry, for example domestic fowl, partridge, geese, pigeon, guinea fowl, teal, duck and peafowl (Cavanagh, 2005; Cavanagh, 2007). In different hosts, the virus exhibits considerable similarities in its genome. For example, a virus that was isolated from teal and peafowl shared 90-99% sequence related to IBV (Liu et al., 2005). Evidence based on the nucleotide sequences of viruses isolated from samples of ducks, whooper swans, turkeys and pheasants have also shown high similarity to IBV (Breslin et al., 1999; Guy, 2000; Cavanagh et al., 2001; Jonassen et al., 2005; Hughes et al., 2009).

### 1.4 Mode of transmission

IBV is a highly infectious pathogen, and the infected birds usually develop clinical signs very rapidly, within 36-48 hours. The virus replicates primarily in the upper respiratory tract, leading to viraemia, and then spreads to other organs (Raj & Jones, 1997). Usually, the virus is present in high concentrations in the upper respiratory tract during the first 3-5 days post infection (Cook, 1968; El-Houadfi et al., 1986). In general, a large amount of virus is detected in tracheal mucus and faeces during the acute and recovery phases of the disease. In some cases IBV persists as a latent infection, and the carrier birds continue to shed virus particles via faeces. The virus is transmitted horizontally by the contaminated feed, water or faeces. Infected birds shed the virus continuously in the environment and contaminate their surroundings, such as equipment, eggs, also working personnel and trucks, among others, which are the major sources of indirect transmission to different regions (Ignjatović &
Wild birds may play a crucial role as reservoirs and long-distance carriers of IBV (Chen et al., 2009; Hughes et al., 2009).

1.5 Infectious Bronchitis Virus

1.5.1 Morphology and genome organization

IBV belongs to the order Nidovirales, family Coronaviridae, genus Gammacoronavirus (Gonzalez et al., 2003). The enveloped viral particles are round and pleomorphic in shape. The virions are approximately 120 nm in diameter and contain club-shaped surface projection called spikes, which are 20 nm in size (Cavanagh & Gelb, 2008). The positive sense RNA genome is approximately 27.6 kb in size and is encompassing 5’ and 3’ untranslated regions (UTRs) with a poly(A) tail (Figure 1) (Boursnell et al., 1987; Ziebuhr et al., 2000; Mo et al., 2012). A major part of the genome is organized as two overlapping open reading frames (ORFs), 1a and 1b, which are translated into large polyproteins 1a and 1ab through a ribosomal frame shift mechanism. The remaining part of the genome consists of regions coding for the main structural proteins spike (S), envelope (E), membrane (M) and nucleocapsid (N). Two accessory genes have been described, ORF3 and ORF5, that express accessory proteins 3a & 3b and 5a & 5b, respectively (Lai & Cavanagh, 1997; Pasternak et al., 2006).

Figure 1. A schematic genome organization of the infectious bronchitis virus

1.5.2 Untranslated regions

The small structural motifs of the IBV genome comprise 5’ and 3’ untranslated regions (UTRs), which mediate physical interactions between UTRs, viral encoded replicase proteins and host cellular proteins (Li et al., 2008).
1.5.3 Replicase proteins
Two-thirds of the genome consists of ORF1a and ORF1b that encode polyproteins 1a and 1ab, respectively, and contribute to formation of the replication and transcription complex (Imbert et al., 2008). These polyproteins are post-translationally cleaved to generate 15 non-structural proteins (Nsp2-16), comprising a main protease called 3C-like protease, a papain-like protease (PL pro), an RNA-dependent RNA-polymerase (RdRp) and other non-structural proteins (van Hemert et al., 2008).

1.5.4 Spike glycoprotein
The spike glycoprotein of all coronaviruses contains four domains that are involved in anchoring of the S protein into the lipid bilayer of the virion. The IBV S gene consists of 1162 amino acids, and is cleaved into two sub-units, the N-terminal S1 subunit (535 amino acids) and the C-terminal S2 subunit (627 amino acids). The S1 subunit contains virus neutralization and serotype-specific antigenic determinants that are responsible for binding to the host cell, neutralization and immune response (Koch et al., 1990; Schultze et al., 1992; Ignjatovic & Galli, 1994; Johnson et al., 2003). The nucleotide sequence variation in the spike gene may result in a lower cross protection between serotypes. The high variation in the nucleotide sequences of spike gene can change the protection ability of a vaccine or immunity (Cavanagh, 2003; Cavanagh & Gelb, 2008). The S2 sub-unit contains a fusion peptide-like region and two heptad regions approximately 100 to 130 Å in length (771-879 amino acid in IBV) that are involved in oligomerisation of the protein and entry into susceptible host cells (Tripet et al., 2004; Guo et al., 2009; Shulla & Gallagher, 2009).

1.5.5 The 3a and 3b proteins
Gene 3 contains two ORFs (ORF3a and ORF3b) that are functionally tricistronic in nature (Liu et al., 1991). ORF3a and ORF3b contain highly conserved nucleotide sequence regions (Mo et al., 2012), not only in IBV but also in other gammacoronaviruses (Cavanagh et al., 2001; Cavanagh et al., 2002).

1.5.6 Envelope protein
The envelope protein is a small integral membrane protein associated with the envelope of the virions (Smith et al., 1990; Liu & Inglis, 1991). It has been demonstrated that the E protein is essential for virus assembly (Maeda et al., 1999). Mutations in the amino acid sequence of the E protein in coronaviruses
may considerably affect the assembling of the virus in cells (Fischer et al., 1998).

1.5.7 Membrane protein
The largest portion of the integral membrane protein is embedded within the lipid bilayer, which maintains the structural integrity (Godet et al., 1992). The M protein is responsible for the organization and assembling of the virus particle by interactions with other structural proteins (Vennema et al., 1996; Hogue & Machamer, 2008).

1.5.8 The 5a and 5b proteins
Gene 5 is encompassing two ORFs (ORF5a and ORF5b) that encode the 5a and 5b proteins, respectively (de Vries et al., 1997). It has been demonstrated that these proteins are functionally bicistronic (Pendleton & Machamer, 2005).

1.5.9 Nucleocapsid protein
The N protein consists of 409 amino acids. This protein is involved in a variety of functions such as viral packing, assembly, viral core formation, signal transduction and modulating host cell processes (Drees et al., 2001; He et al., 2004; You et al., 2007).

1.6 Post-transcriptional and post-translational modifications and structural motifs
The post-transcriptional and post-translational modifications such as cleavage, palmitoylation, N-glycosylation, phosphorylation and structural motifs Leucine-rich repeat have important role in virus biology. Therefore, there is need to explore these modifications and structural motifs especially by application bioinformatics, to identify the differences in the spike glycoprotein of the virus variants, in order to bridge a platform for the in-vivo and in-vitro studies, which is part of scope of the present thesis.

1.6.1 Cleavage site motif
The spike glycoprotein contains a protease cleavage site motif that is involved in cleavage of the S1 and S2 sub-units during viral maturation (Cavanagh et al., 1986). The cleavage site motif of IBV comprises one or two pairs of basic amino acids that is cleaved by host cell serine proteases (Cavanagh et al., 1992). The serine proteases of host cell catalyze hydrolyses reactions that cause cleavage of peptide bonds (Voet & Voet, 1990).
1.6.2 Palmitoylation

Palmitoylation is the binding of organic molecules, e.g., palmitic acid, to the cysteine residues of membrane proteins (Linder, 2000). Palmitoylation plays an important role in localization and transport at the sub-cellular level, in protein-protein interactions, and in various physiological characteristics of proteins in the virus (Dunphy & Linder, 1998; Dietrich & Unger mann, 2004). The carboxyl-terminal cysteine peptides of viral membrane glycoproteins act as crucial palmitoylation sites (Ponimaskin & Schmidt, 1995; Petit et al., 2007). In coronaviruses, palmitoylation of viral glycoproteins affects the fusion to cellular membranes, viral assembly and infection of cells (Bos et al., 1995; Petit et al., 2007).

1.6.3 N-glycosylation

The N-glycosylation characteristics of viral glycoprotein are associated with changes of virulence and cellular tropism (Li et al., 2000). Glycosylation in the spike and membrane glycoproteins of coronaviruses is involved in fusion, receptor binding and antigenic characteristics (Alexander & Elder, 1984; Braakman & van Anken, 2000; de Haan et al., 2003; Wissink et al., 2004). Variation in N-glycosylation sites can affect the interaction with receptors, and thus render a virus more susceptible to host innate immune responses and lower recognition by antibodies, affecting virus replication and infectivity (Meunier et al., 1999; Land & Braakman, 2001; Slater-Handshy et al., 2004; Vigerust & Shepherd, 2007).

1.6.4 Phosphorylation

Protein phosphorylation plays a crucial role in the regulation of functional activities in microorganisms (Ingrell et al., 2007). For IBV, two phosphorylated clusters (Ser190 & Ser192 and Thr378 & Ser379) that were identified by a mass spectroscopic method, are located in the middle and C-terminal regions of the N protein (Chen et al., 2005). The C-terminal phosphorylation clusters are associated with the differentiation of viral RNA from non-viral RNA (Spencer et al., 2008).

1.6.5 Leucine-rich repeat

The leucine-rich repeat (LRR) domain is contained in microbial proteins and is associated with innate immunity (Huang et al., 2008). Beyond innate immunity, LRR-containing proteins are associated with various cellular processes including apoptosis, ubiquitin-related processes and nuclear mRNA transport (Kobe & Kajava, 2001; Wei et al., 2008).
1.7 Genetic forces

The genetic forces such as mutations, recombination and selective pressure play significance role in the viral genome evolution (Zhang et al., 2006; Liu et al., 2007). In order to better understand the contribution of these forces in genetic diversity of IBV, these were outlined as subjects of thesis.

Recombination commonly occurs between two or more viruses infecting the same cell. It is believed that a high rate of recombination events occurs in the genomes of non-segmented RNA viruses (Makino et al., 1986), such as in IBV and in other coronaviruses (Lim et al., 2011; Thor et al., 2011; Jackwood et al., 2012). In RNA viruses, a unique copy choice mechanism during polymerase activity allows a high efficiency of RNA recombination (Liao & Lai, 1992). Moreover, recombination, along with the polymerase error rate, is involved in molecular mechanisms that are likely to be responsible for changing tissue and host species tropism of the virus (Graham & Baric, 2010).

In the case of coronaviruses, mutations and selective pressure in genes, especially in hypervariable regions, enable the virus to cross the species barrier and adapt to new host species, hence contributing to viral evolution (Zhang et al., 2006; Liu et al., 2007). It has been demonstrated that during the multiple passages of IBVs in embryonated eggs, mutations and recombination result in strong selective pressure in the S1 gene, leading to the generation of attenuated viruses (Liu et al., 2007).

1.8 Serotypes and genotypes of Infectious Bronchitis Virus

A number of IBV serotypes, e.g. Arkansas, Connecticut (Conn), Massachusetts (Mass), 4/91 or 793B, D274, H120, Italy-02, Baudette, California, Georgia 98 and QX have been identified and reported around the globe (Lee & Jackwood, 2001; Jackwood et al., 2003; Sjaak de Wit et al., 2011). Serotyping of IBV strains is usually carried out by test systems based on chicken-induced IBV serotype-specific epitope(s) (Koch et al., 1990; Keeler et al., 1998). The appearance of a large number of IBV variants in different regions is a major constraint for the practical application of serotyping (Sjaak de Wit et al., 2011). It is important to note that serotyping of IBV is becoming less and less common today, since the serological tests require a panel of serotype specific antibodies, which is not easy to obtain. Thus, recently the typing of the virus is mostly performed by various molecular methods as many types of the virus are diagnosed by molecular methods.

Genotyping of IBV is based on the amplification of the highly variable sequence region of the S1 gene by reverse transcriptase polymerase chain reaction (RT-PCR), followed by sequencing (Jackwood et al., 1992; de Wit,
The molecular diagnostic assays revealed that different genotypes, such as Massachusetts, 4/91, D274, Italy-02 and European QX-like, are circulating in European countries (Figure 2) such as Belgium, Denmark, France, Hungary, Italy, Germany, The Netherlands, Poland, Russia, Slovenia, Spain, Sweden and UK (Davelaar et al., 1984; Parsons et al., 1992; Farsang et al., 2002; Jones et al., 2005; Bochkov et al., 2006; Domanska-Blicharz et al., 2006; Worthington & Jones, 2006; Dolz et al., 2008; Worthington et al., 2008; Benyeda et al., 2009; Handberg et al., 2009; Valastro et al., 2010; Krapez et al., 2011; Abro et al., 2012a; Abro et al., 2012b).

Figure 2. A phylogenetic tree, based on the comparative analysis of S1 gene sequences of different strains obtained from various countries, is showing the occurrence of different genotypes of IBV in Europe.
1.9 Detection of Infectious Bronchitis Virus

Due to constant evolution, there is continuous emergence of new IBV variants all over the world. For example, different genotypes such as Massachusetts, 4/91, D274, Italy-02 and European QX-like are currently circulating in Europe. Various methods are used for the detection and identification of IBV. The most common assays for routine diagnosis are virus isolation, haemagglutination inhibition (HI), enzyme-linked immunosorbent assay (ELISA), immunoperoxidase assay (IPA), virus neutralization test (VNT), immunofluorescence assay (IFA), agar gel precipitation test (AGPT), RT-PCR and real-time RT-PCR (de Wit, 2000). The majority of the conventional diagnostic assays in fact are time- and material-consuming, laborious, costly and provide relatively low specificity and sensitivity (King & Hopkins, 1983; Mockett & Cook, 1986; Karaca & Naqi, 1993; Naqi et al., 1993; De Wit et al., 1995; de Wit et al., 1997; de Wit, 2000). Various real-time RT-PCR assays have been recently elaborated in veterinary medicine for the diagnosis of a variety of infectious pathogens (Belák, 2007). Accordingly, a range of real-time RT-PCR assays has been developed for the specific detection of IBV using various approaches, such as the TaqMan technology. In the general real-time RT-PCR technique is highly sensitive and it can be applied not only for the rapid detection and identification of IBV, and even to quantify genomic RNA in clinical samples (Callison et al., 2006). On the other hand, sometimes even the real-time RT-PCR assay may lead to poor results due to limitations in detecting the divergent or the new virus variants. In addition, being based on the conserved 5’ UTR or the nucleocapsid genes, the amplicons obtained by diagnostic PCR assays are not suitable for determining the types of the viruses. The virus typing, in majority of the cases is based on the comparative analysis of nucleotide sequences of the variable S gene.

Considering that there are no reliable and practical assays available today, that can simultaneously detect and type IBV variants; the development of such an assay was contemplated in the present work. By applying variation-tolerant detection chemistry and the Luminex technology, we have developed a new tool for the rapid detection and accurate identification of various genotypes of IBV, including newly emerging variants of this virus.
2 Aims of the studies

The overall objectives of the present studies were to characterize various strains of IBV, to determine genetic diversity and to investigate the factors driving the evolution of this virus. A further task was, to develop a molecular diagnostic assay, in order to improve the detection and typing of IBV variants causing disease outbreaks in Sweden and in other European countries.

The specific aims were:

- To perform molecular characterization of IBV isolates from Sweden based on the comparative sequence analysis of the complete spike gene, in order to better understand the epidemiology and the factors behind the occurrence of new outbreaks (Paper I).
- To study the full-length genome of the European QX-like isolate of IBV, detected in Sweden, by sequencing and phylogenetic analysis (Paper II).
- To compare the molecular characteristics of isolates belonging to the classical Massachusetts and emerging QX-like genotypes of IBV in order to determine differences in potential functional and structural motifs of the spike gene that may relate to the biological properties of the virus. In addition, a comparative analysis of the complete spike gene was performed based on sequences of avian and mammalian coronaviruses (Paper III).
- To develop a specific assay for the simultaneous detection and typing of the QX-like, Italy-02, M41, 4/91 and D274 genotypes of IBV, which are currently circulating in Europe (Paper IV).
3 Materials and methods

3.1 Samples and screening for Infectious Bronchitis Viruses

Clinical samples (trachea, bronchi, ceaca) were obtained from different outbreaks in Sweden (Papers I, II, III and IV). The D274, Italy-02 and 4/91 isolates kindly provided from the Netherlands, Slovenia and France (Merial) were used in the study (Paper IV). The samples were screened by real-time RT-PCR assay as described by Callison et al. (2006) for the presence of IBVs (Papers I and IV).

3.2 Isolation of Infectious Bronchitis Viruses

The real-time RT-PCR positive IBV samples were propagated in specific pathogen-free (SPF) embryonated hen’s eggs. Virus isolation was performed by inoculation of 9–11 day old eggs with 200 µl of 10% tissue homogenates and incubation at 37 °C for 72 hours. The allantoic fluid was harvested, and isolates were stored at -20 °C for the further use.

3.3 RNA extraction, synthesis of cDNA, PCR amplification and sequencing

The viral RNA was extracted using the TRIzol reagent according to the manufacturer’s instructions (Invitrogen, Carlsbad, USA) (Papers I, II, and IV). Synthesis of cDNA was performed with gene specific primers using SuperScript II reverse transcriptase (Invitrogen™ Life technologies, Carlsbad, USA) as recommended by the manufacturer (Papers I and II). PCR amplification and sequencing were performed for the S gene and also for fragments covering the whole genome (Papers I, II and IV). Different pairs of primers were used for amplification and sequencing of the full-length genome
as reported by Liu et al. (2009) (Paper II). The sequencing reactions were performed by using Big Dye terminator sequencing kit (Applied Biosystems, Foster City, CA), as recommended by the manufacturer (Papers I, II, and IV). For sequence analysis, the sequence data set was pair wise edited and aligned with the software Lasergene DNASTAR (Madison, USA) (Papers I, II and IV).

3.4 Recombination analysis

The recombination analyses were performed on the complete gene sequences using the Recombination Detection Program (RDP v3.44) (Martin et al., 2010). Different methods available in the program were applied in order to compare and accurately determine the positions of the recombination hot spots and to differentiate closest fragment of non-recombinant sequences (Ohshima et al., 2007). The marker positions were indexed for the determination of recombinant clusters (Papers I and II).

3.5 Selection pressure

Evidence of selective pressure in individual genes was examined by using the SNAP (Korber et al., 2000) services available at web server http://hcv.lanl.gov/content/sequence/SNAP/SNAP.html (Papers I and II). The differences between synonymous (dS) and non-synonymous (dNS) amino acid substitutions were calculated in order to determine the substitution rate in the analyzed gene(s) individually (Papers I and II).

3.6 Analysis of functional and structural motifs

As reported in Paper III, N-glycosylation sites were predicted using services available at web server http://www.cbs.dtu.dk/services/NetNGlyc. The potential phosphorylation sites were determined by using the website http://www.cbs.dtu.dk/services/NetPhos. The cleavage site motif analysis was performed by the traditional approach using the amino acid position in the sequence of the spike glycoprotein, as previously described by (Parker & Masters, 1990; Jackwood et al., 2001). LRR regions were identified by LRR finder, available at http://www.lrrfinder.com/result.php. The primary structures of the spike glycoprotein were predicted by http://expasy.org/tools. The secondary structures were predicted by the GOR4 method using services at http://npsa-pbil.ibcp.fr/cgi-bin/npsa_automat.pl?page=npsa_gor4.html. The palmitoylation sites were determined with the medium threshold frequency using web domain http://csspalm.biocuckoo.org/prediction.php.
3.7 Phylogenetic analysis

As described in Papers I, II and III, phylogenetic analysis was performed using sequences generated in these studies, as well as sequence data downloaded from the GenBank database. Phylogenetic relationship was determined using both parsimony and nucleotide distance methods. The nucleotide distance matrix between sequences was used to construct a phylogeny by Neighbor-Joining (NJ) using MEGA 4 (Tamura et al., 2007). The NJ method was applied because it is relatively fast and can be suitable for large data sets (Tamura et al., 2004). The sequences were compared by using the gamma Tamura–Nei model (Tamura & Nei, 1993). Bayesian analyses were performed by using MrBayes 3.1 software (Ronquist & Huelsenbeck, 2003).

3.8 Variation-tolerant capture multiplex assay (VOCMA)

As reported in Paper IV, synthetic targets were designed as an alternative positive control for each genotype: M41, D274, 4/91, Italy-02, QX-like, IBV_Capsid (consensus). The hybridization events of variation tolerant (VT) were designed using the NucZip algorithm (Öhrmalm et al., 2010; Öhrmalm et al., 2012).

The amplification of target nucleic acid by VOCMA, executed as single-tube one-step multiplex amplification, was performed using long specific VT primer-probes and short generic primers, with the biotinylated generic second primer. The extracted nucleic acid or synthetic ssDNA target was added directly to a mastermix one-step RT-PCR mixture. Specific synthetic 5 amino-C12 modified detection probes for the different IBV genotypes were designed and coupled to xMAP carboxylated colour-coded microspheres (Luminex Corp., Austin TX, USA) according to the manufacturer’s instructions (Luminex Corporation, Austin TX, USA). The biotin-labeled amplified template was mixed with hybridization buffer of each probe-coupled xMAP bead. The IBV VOCMA has six unique beads, each coupled with either six specific VT detection probes of 48-62 nt (long probes) or of 33-36 nt (short probes). The mixture was treated at 95 °C for 2 min, followed by hybridization at 50 °C for 30 min with shaking at 600 rpm on a Thermostar (BMG LabTech; Offenburg, Germany) microplate incubator. After a short centrifugation, 40 µL of supernatant was discarded and a mixture of 38µL 3M TMAC-TE hybridization buffers + 2 µL of Streptavidin-R-phycoerythrin (QIAGEN, Hilden, Germany) was added. The tubes were further incubated at 50 °C for 15 min, before analysis for internal bead and R-phycoerythrin reporter fluorescence on the Luminex®200™ flow meter (Luminex corporation, Austin
The quantity of biotinylated target that hybridized to the probe-linked beads was measured as Median Fluorescence Intensity (MFI).
4 Results and discussion

4.1 Characterization and genetic diversity of the S gene of IBV (Papers I and III)

Outbreaks of infectious bronchitis have been detected in Sweden between 1995 and 2010. Twenty samples originating from these outbreaks were selected for molecular characterization based on analysis of sequences of the complete S gene (Paper I). The viruses collected in 1995-1999 showed nucleotide sequence difference of varying degree 18.7-21% in comparison to the isolates from 2009-2010. The isolates from the 1990s shared lower identities to the viruses collected in the 2000s, reflecting the genetic signature of new variants. Considerable sequence variation was found in the region encompassing the S1 part of the S gene. In the S2 subunit, regions of sequence variation were found interspaced with regions of high conservation, also contributing to overall diversity of the S gene. The variation in nucleotide sequences of the S1 region may lead to alter the virus characteristics such as epitopes and receptor binding abilities. It has been reported that comparative analysis of the hypervariable region of the S1 nucleotide sequences is a suitable tool for the discrimination of various IBV field variants (Kusters et al., 1987; Kwon et al., 1993; Wang & Huang, 2000). There were nucleotide insertions and deletions at different locations in the spike gene of isolates from the 2000s in comparison to isolates from the 1990s. The nucleotide insertions, deletions and point mutations in the S gene contribute to evolution of IBV (Kusters et al., 1990; Lai, 1992; Kottier et al., 1995). The significant substitutions in the S1 region warrant further investigation to ascertain their relevance, if any, in the virus biology.

Substitutions in the coding sequences are likely attributable to strong positive selection pressures in the spike gene. Strong selective constraints could affect the primary and secondary structures of the gene, which may lead to alteration of genetic and molecular features of the virus. It has been reported
that positive selection in IBV can lead to emergence of new strains that are capable of escaping the immune system (Dolz et al., 2008).

Recombination events were detected in the spike gene sequences of isolates from 2009-2010 (Paper I). This is an important information, considering the fact that genetic recombination among heterologous IBVs could lead to the emergence of new variants, and result in genetic diversity (Kottier et al., 1995; Lee & Jackwood, 2000; Lim et al., 2011). Recombination is believed to decrease the mutation rate and associated constraints which are responsible for the genetic diversification that results in emergence of new variants of coronaviruses (Worobey & Holmes, 1999).

Taken together, point mutations, strong selective pressure and to some extent recombination events in the spike gene are contributing factors to the generation of new emerging variants and genetic diversity that leads to constant evolution of the virus.

Phylogenetic analysis based on partial S1 and complete S gene sequences revealed four distinct clusters: Massachusetts, QX-like, Italy-02 and 4/91 genotypes. Swedish isolates from 1995 to 1999 clustered together with strains from China, Korea, Spain and Thailand, represented as Massachusetts-type. The Swedish isolates from 2009-2010 formed a group with sequences from France, Italy, the Netherlands, Spain and UK, which belonged to the QX-like genotype (Paper I). The analysis revealed new branches within the QX-like genotype, indicative of the emergence of new strains. Furthermore, this signifies a shift from predominantly classical Massachusetts viruses in 1990s to emerging QX-like viruses dominating in Sweden. QX-like emerging viruses showed close relatedness to QX-like types present in other European countries. Hence, the emergence of these QX-like viruses could be attributed to trade and to importation of chickens and poultry products from other European countries. Alternatively, indirect routes may have contributed to the introduction of the QX-like strains in Sweden. As has been reported, QX-like viruses were detected in France in 2004 close to the border regions of Belgium and in The Netherlands where these viruses were already prevalent (Worthington et al., 2008). The introduction of viruses by wild or migratory birds as further hosts and carriers of IBV still remains to be investigated, although it has been reported that migratory birds play an important role as reservoirs, in transmission and in long-distance carriage of coronaviruses (Chen et al., 2009; Hughes et al., 2009; Muradrasoli et al., 2010).

As reported in Paper III, phylogenetic analysis of the complete S gene revealed variable relationships among the investigated avian and mammalian coronaviruses. The high genetic divergence observed in the coronaviral genomes is likely due to host specificity and continuous mutations in this gene.
Furthermore, this genetic diversity in the spike gene may affect the biology of viruses, as changes favour interspecies transmission and tissue tropism that leads to adaptation to new host species. Previously, a shift of tissue tropism in IBV has been described (Zhou et al., 2004; Liu et al., 2006), resulting in infection of a wide range of avian host species (Cavanagh, 2005).

As described in Paper III, with the help of an *in silico* approach, comparative differences were observed between QX-like and Massachusetts strains, notably, in N-glycosylation sites, palmitoylation domains, phosphorylation peptides, cleavage sites, LRR motifs and primary and secondary structures of the spike glycoprotein. The changes in these molecular and structural characteristics may influence the infection biology of the virus. However, the predicted characteristics need to be further investigated in biological assays to determine if they relate to differences in the behavior of these viruses *in vivo*.

4.2 Analysis of full-length genome sequence of an emerging QX-like isolate of IBV (Paper II)

Recently, QX-like viruses are taking relevance in infections in chicken flocks in most countries of Europe, posing problems for control of avian infectious bronchitis. Thus, Paper II provides novel information about the genome of an European QX-like virus responsible for one such outbreaks. The complete genome of isolate CK/SWE/0658946/10 consists of six genes (27,664 nt) with ten open reading frames (ORFs) flanked by 5' and 3' UTRs. In the comparison of full-length genomes, the isolate was closely related to the sequence of the strain ITA/90254/2005, with lower sequence similarity shown to the classical Massachusetts strain. The sequences of the individual ORF1a, ORF1b, S and M genes shared lower similarities to the corresponding sequences of the classical Massachusetts strain. The lower similarities in the two viral genomes may be the result of high nucleotide substitution rates or to strong selective constraints. The evolutionary pressure analysis of the genome showed strong selective constraints especially in the replicase and spike genes, with a high substitutions rate across the genes. Also, recombination events were observed in the N gene of the isolate. A high rate of mutation and strong selective constraints across the genome contribute to high genetic divergence of the virus.

In addition to the clinical features in flocks infected with QX-like viruses, high sequence similarities and close phylogenetic relatedness was found to the viruses isolated from other European countries. Taken together, these characteristics explain the high genetic relatedness of QX-like viruses
circulating in Europe. Currently, the wide spread of the QX-like viruses in Europe may be associated with intensified poultry husbandry and less-strict border controls on the continent.

The generated sequence data and information will be useful for potential application of generation of an infectious clone, determination of pathogenic nature of the virus, and has relevance for diagnostics and vaccines.

4.3 Development of VOCMA for broad detection and typing of IBV (Paper IV)

The studies described in Papers I, II and III have demonstrated that high sequence variability, strong selective pressure, recombinant events, and variation in molecular motifs, especially in the spike gene are responsible for genetic divergence within IBV. This high genetic diversity poses a problem for diagnostics and especially for the accurate typing of strains involved in new outbreaks. The majority of the currently used diagnostic methods are not able to differentiate and type the different variants of IBV. This may lead to diagnostic problems, especially when multiple genotypes are co-circulating in a region, such as with the co-existence of Mass 41, Italy-02, 4/91, QX-like and D274 recently in Europe.

Considering these problems, there is a need for an assay that could simultaneously detect and type different IBV variants. For this reason, we have designed and developed an assay for specific and simultaneous detection of IBV QX-like, Italy-02, M41, 4/91 and D274 using a new technique, termed VOCMA (Öhrmalm et al., 2012).

In order to provide broad-range detection, the assay targets the capsid and hypervariable sequence region of the S gene that facilitates the simultaneous typing of the various viral variants. Being based on the variation tolerance and multiple detection principle, the assay has great potential to target divergent sequences in the genomes of the different variants of IBV. Two sets of genotype specific detection probes (long and short probes) were designed to facilitate the detection of the specific genotype: long probes to create a broader variation tolerance (48-62 nucleotides) and short probes to gain higher specificity (33-36 nucleotides). Both probe-sets revealed high specificity that could distinguish the targeted IBV genotypes from each other. The analytical specificity of the assay was evaluated with the selected set of different genotypic variants. The strong signals detected were specific to their respective targets. The possible cross-reactions in the IBV assay were evaluated and no cross-reactivity or non-specific signals were detected when tested on heterologous avian and bovine coronaviruses. So far, the results
demonstrated a low possibility of false positive reactions that could result from the presence of other coronaviruses in the field samples.

The performance of the assay was evaluated by testing 46 samples for the five targeted IBV genotypes. The detected signals were specific to their respective targets. The positive samples were amplified and sequenced by targeting in the spike gene, and the sequence analysis confirmed detection of the correct genotype by the assay.

The strength of this assay is that it is able to detect and simultaneously genotype a wide range of virus variants of IBV. A weakness is that, the sensitivity of VOCMA is lower, compared to the real-time RT-PCR. Therefore, a considerable amount of virus in the sample is needed for the detection by VOCMA. Further work is required to increase the sensitivity of the assay.

In summary, the developed VOCMA method was found to be useful for the simultaneous detection and typing of IBV strains representing different genotypes. The assay is relatively fast and specific, and it provides a new tool for surveillance and monitoring of IBV infections and for large-scale epidemiological studies.
5 Conclusions

- The Massachusetts-type strains, which were predominant in Sweden in the 1990s, have been replaced by QX-like variants. The emergence of these QX-like viruses is likely the consequence of their introduction from other European countries.

- The complete genome sequence data and the phylogenetic analysis have revealed that the genome of IBV is under a continuous process of evolution, due to mutations, strong selective pressure and recombination events. The data will be useful in further evolutionary studies and for a better understanding of the infection biology of the virus. Furthermore, the information will be helpful for the development of novel diagnostic assays and vaccine candidates.

- This study provided insights showing that functional and structural motifs in the spike glycoprotein are different between the classical Massachusetts and emerging QX-like genotypes. The predicted characteristics have to be further investigated in biological assays, in order to determine if they relate to differences in the behaviour of the viruses under in vivo conditions.

- The multiplex VOCMA assay provides a method for the molecular detection and typing of the diverse IBV genotypes circulating in Europe. Furthermore, this assay is considered to be a practical new tool for the monitoring of IBV infections and for large-scale epidemiological studies.

- The observations concerning the genome construction and evolutionary aspects of IBV variants, as well as the development of a new detection and typing assay, provide information and new possibilities to combat infectious bronchitis, a viral disease of global importance.
6 Future prospects

This thesis has dealt with the characterization and detection of IBV variants currently circulating in Sweden and other European countries. The characterization was intended to provide a better understanding of the molecular biology of the virus and to investigate the factors behind the continuous emergence of new genetic variants of this virus. The possibility to detect the various genotypes present in Europe was improved by the development of a new genotyping assay. While the present research will be helpful for diagnosis, epidemiology, disease monitoring and adoption of effective control measures, further research questions have arisen that need follow-up studies:

- The continuous variation of IBV makes it very difficult to control infectious bronchitis by using live attenuated vaccines for immunization (Cavanagh, 2003). The knowledge gained from this thesis can be useful for the development of new vaccine candidates that would provide adequate immunity, and protect poultry against multiple IBV variants.
- The biological characteristics of different strains of IBV, particularly with regard to virulence, need to be further investigated. In addition, the pathogenesis of IB induced by the currently circulating viral variants needs to be further studied.
- As the egg inoculation studies are time consuming and cumbersome it would be of great value to establish susceptible cell lines and powerful systems of virus production, in order to facilitate studies on the virus and on the infection it causes.
- Further investigations are needed to study tissue tropism, particularly of the new virus variants, in order to better understand the changes in virus behavior and the development of various disease manifestations,
ranging from classical respiratory disease to symptoms associated with the new and emerging strains.

- Continued research is needed on reverse genetics for better understanding of the molecular interactions and for the development of novel IBV vaccines.
- The role of various animal species for transmission of IB or IBV-related coronaviruses, such as wild birds, bats etc., requires further study.

This research would be helpful for improved prevention and for more effective control of this disease.
7 Populärvetenskaplig sammanfattning

Infektöst bronkitvirus (IBV) drabbar höns men även andra fågelarter som lever i nära anslutning till tamhöns. Det har påvisats hos fasaner, pärlhöns, kricka och påfågel. Sjukdomen infektös bronkit (IB) är en för slaktkyckling- och äggproduktion ekonomiskt över hela världen viktig sjukdom förutom att den negativt påverkar fåglarnas välbefinnande.


References


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