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### Workshop: Infectious Bronchitis (IB) in the Brazilian Poultry Industry

# Infectious Bronchitis Virus in Asia, Africa, Australia and Latin America - History, Current Situation and Control Measures

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#### ■ Keywords

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#### **ABSTRACT**

#### **INTRODUCTION**

Infectious bronchitis virus (IBV) is ubiquitous in most parts of the world where poultry are reared and is able to spread very rapidly in non-protected birds. It is shed via both the respiratory tract and the faeces and can persist in the birds and faeces for several weeks or months. Although strict biosecurity and working with a one-age system are essential control measures, normally vaccination is an essential tool to increase the resistance of the chickens against challenge with IBV strains. This paper presents an overview of the history and current situation of IBV in Asia, Africa, Australia and Latin America, including the control measures necessary to control it.

#### HISTORY OF INFECTIOUS BRONCHITIS VARIANTS

For many years it was widely believed that the first variants of infectious bronchitis virus (IBV) occurred in the early 1950s as Jungherr et al. (1956) in the USA showed that the Connecticut (Conn) isolate of 1951 neither cross neutralised nor cross protected with the original Massachusetts (Mass) isolate from the early 1940s. However, a retrospective study (Jia et al., 2002), using monoclonal antibodies and molecular analysis of part of the S1 subunit of the spike glycoprotein (S) gene, identified non-Mass IBVs among US isolates as early as the 1940s. Since the 1950s there have been many reports of other IB variants in USA (reviewed by Fabricant, 1998), some of which are found worldwide. However, most countries are now known to have their own indigenous variants as well, and this paper will consider the situation outside USA and Europe.

IBV appeared in Latin America by the 1950s and the first reported isolate from that continent was of the Mass serotype in Brazil (Hipólito, 1957), although isolation of the first variant (Arkansas [Ark]) was not reported in that county until some 10 years later (Branden & Da Silva, 1986). In a study carried out in the mid 1990s, IB isolates of at least 5 different antigenic types were found in commercial chickens of all types throughout Brazil, but mainly in the major poultry producing area of the south (Di Fabio et al., 2000). More recently, several different genotypes have been identified in Brazil by analysis of either the S1 (Villarreal et al., 2007a; Villarreal et al., 2007b; Montassier et al., 2008) or the nucleoprotein (N) genes (Abreu et al., 2006), but protection studies have not been performed. It is important to remember that in Brazil at that time, as in most other parts of the world, the only live attenuated IB vaccines licensed for use were of the Mass serotype and protection

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studies (Cook et al., 1999; Di Fabio et al., 2000) showed that the currently available Mass vaccine provided inadequate protection against some of these new variants. Hidalgo et al. (1976) reported the first isolation of IB (Mass serotype) in Chile in 1975 and variants were reported for the first time there some 10 years later (Hidalgo et al., 1986). By the mid 1980s, Infectious Bronchitis (IB) was reported as a serious problem in commercial chicken flocks and novel variants, as well as the Mass and Conn serotypes, were isolated from broiler and layer flocks and again, Mass vaccines were found to provide poor protection against challenge with these variants (Cubillos et al., 1991). Elsewhere in Latin America a variant IB was isolated from commercial chickens in Honduras in 1997 and poor protection against it using Mass vaccines was demonstrated (Cook et al., 1999). Following the isolation of the Ark serotype in Mexico in the early 1990s, the use of molecular methods then identified IB variants unique to that country in commercial chickens (Escorcia et al., 2000; Callison et al., 2001; Gelb et al., 2001). They were shown by neutralisation tests to be different from Mass or Conn serotypes, but in vivo protection studies were not performed. Rather surprisingly, IBV variants do not appear to have been reported in Argentina until very recently, when Rimondi et al. (2009), using only molecular techniques, detected 3 unique genotype clusters (in addition to Mass and Conn), one of which was closely related to isolates from Brazil. Similar techniques had been used a few years earlier to identify for the first time a genetically unique IB variant in Colombia (Alvarado et al., 2005).

An IBV associated with swollen head syndrome and causing severe problems throughout southern Africa was isolated in the early 1980s (Morley & Thomson, 1984), confirmed as a variant, and shown to be poorly protected against by Mass vaccines (Cook *et al.*, 1999). The only other report of the detection of IBV variants in sub-Saharan Africa is the recent report by Ducatez *et al.* (2009), who detected a novel IBV in Nigeria and Niger, antigenically and genetically distinct from other known IBVs. However, no association with disease was demonstrated and there is no information on the ability of currently available vaccines to protect against it.

IB variants have been recognised in Egypt since the 1950s (Sheble *et al.*, 1986; Eid, 1998) with the isolation of an IB variant shown by neutralisation tests to be closely related to the Dutch variant D3128 and subsequently variants related to Mass, other Europeans IBVs and one related to an Israeli variant

have been identified by genome analysis in that country (Abdel-Moneim *et al.*, 2006). In the early 1980s the unusual enterotropic variant, know as IB "G" was isolated in Morocco (El-Houadfi & Jones, 1985). More recently, Bourogaa *et al.* (2009) used both molecular methods and cross neutralisation tests to identify as variants IB isolates from Tunisia, which are closely related to ones found in Europe.

Variant IBVs have been recognised in Israel since at least the mid 1990s (Meir et al., 1998; Callison et al., 2001; Meir et al., 2004) on the basis of both virus neutralisation tests and molecular techniques, and protection studies have shown that Mass vaccines provide inadequate protection against some of these novel variants. In Jordan, the use of RT-PCR enabled European IB variants D274 and 4/91 (793B) to be detected (Roussan et al., 2008), but since the primers used were designed to detect only specific variants, it is possible that others are present in that country. Similar methods have been used to identify 4/91 in Iran (Seyfi Abad Shapouri et al., 2004).

In the Indian subcontinent, antibodies to several "American" (Muneer *et al.*, 1987a) and "European" (Ahmed *et al.*, 2007) IB variants have been demonstrated in Pakistan, but virological studies have still to be performed. An IBV isolated in India in the early 2000s from cases of nephrosis, was reported to have a unique S1 sequence, indicating it to be different from other known IBVs (Bayry *et al.*, 2005).

In Asia, many studies have been performed in different countries in recent years. In Malaysia where IBV was first isolated in 1967, variants have been present since at least 1979 (reported by Lohr, (1988). More recently, molecular epidemiological studies of IBVs isolated in Malaysia and Singapore showed that, whilst some were of the Mass serotype (probably identical to the H120 vaccine), others were similar to IBVs reported from China and Taiwan (Yu et al., 2001). These studies led the authors to suggest that IB variants have existed in Asia for some time. This finding was substantiated by Zulperi et al. (2009) who used sequence and phylogenetic analysis to study two variants isolated in Malaysia, 10 years apart. One was similar to several Chinese variants whilst the other was characterised as unique to Malaysia, but no protection studies were performed. IB has been a problem in Thailand since the 1950s, despite the use of many different IB vaccines and a recent molecular study by Pohuang et al. (2009) has identified two groups of IB variants in Thailand by phylogenetic analysis of the SI gene. Group I appeared to be unique to that country, whilst Group II showed a



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close relationship to Chinese IBVs, including variant A2 (see below).

IB variants have been associated with disease outbreaks in Korea since at least the mid 1980s (Song et al., 1998). Initially, Mass vaccines were successful in controlling disease, but since 1990 IB outbreaks have been experienced in well-vaccinated flocks with an increased incidence of renal problems. Song et al. (1998) classified 40 IB isolates as Mass plus four local genotypes, one of which was not only the predominant type, but in pathogenicity studies, it caused 50% mortality in SPF chicks inoculated at day-old. More recent studies (Lee et al., 2004) have extended this work and reported further genetic diversity amongst Korean IB variants isolated from diseased flocks; some of which are indigenous to Korea, whilst others share genetic relationships to IB variants from other countries in the region (Lee et al., 2008). It is suggested that Korean IBVs are continually evolving (Jang et al., 2007).

Mase et al. (2004) carried out a detailed analysis of Japanese IB variants by looking at the N-terminus of the S1 glycoprotein and identified three major genetic groups not found in other countries. One group, present in Japan since at least the 1960s, may be found only in Japan, whilst the other two, which are more recent, are related to Chinese and Taiwanese variants (see below). These groups were distinct from those found in Europe or USA. However, the 4/91 serotype has recently been isolated in Japan (Mase et al., 2008; Shimazaki et al., 2008).

IB variants have been recognised in Taiwan since at least the mid 1960s and two distinct lineages have been identified, as well as Mass and IBVs related to those reported in neighbouring countries (Wang & Tsai, 1996; Wang & Huang, 2000; Huang *et al.*, 2004). The failure of Mass vaccines to provide adequate protection led to the development of vaccines from indigenous strains (Huang & Wang, 2006).

For many years little was known of the situation regarding IB variants in China, but the fact that Mass type vaccines were used successfully suggests that variants may not have been a problem before the 1980s. However, by the mid 1990s, this was clearly no longer that case and in the last decade many published reports have revealed the diversity of IBVs causing disease in that country. By means of *in vivo* studies and antigenic typing using monoclonal antibodies and cross-neutralisation tests, Wu *et al.* (1998) identified highly pathogenic IBV variants in China associated both with respiratory disease and nephritis, and showed that the H120 vaccine provided poor protection against the

challenge with these isolates. In a recent study of 26 IBVs isolated between 1985 and 2008 from a variety of disease conditions in the Guangxi region of China, Wei et al. (2009) identified 4 clusters based on RT-PCR analysis of the N gene. They were grouped into 7 serotypes by neutralisation tests, but there was poor correlation between the results of genotyping and serotyping. In an analysis of the genome of 26 IB variants isolated from kidneys, proventriculus, and oviduct in different areas of China between 1995 and 2004, Liu et al. (2006) identified Mass type IBVs plus five genotypes apparently found only in China and cocirculating there. One of these (genotype A2) was subsequently shown to be the dominant indigenous type in China (Liu et al., 2009). Other genotypes showed close relationships with either Korean or Taiwanese IB variants, and one was closely related to an Australian isolate (Liu et al., 2006). Cuiping et al. (2007) identified the 4/91 variant in China, along with the Australian T strain, as well as one variant indigenous to China. Thus the diversity of IB variants in China is now well established, some being restricted to and co-circulating in that country, whilst others show similarities with IB variants identified in other countries in that region.

Molecular characterization using a part of the S gene of 91 IBV strains isolated between 1998 and 2002 from chickens in Russia showed the complexity of the Russian IBV situation (Bochkov *et al.*, 2006). The main group of isolates (38 viruses) belonged to the Mass genotype circulating in Russia since the early 1970s. A second group of 22 strains were of known European genotypes D274, 4/91, B1648, 624/I and Italy-02. Two isolates from very distant geographic locations in Russia (Far East and the European part) clustered together with Chinese strains of the QXIBV genotype. The remaining 27 Russian isolates were divided in 11 novel genotypes.

In Australia, where IBV has always evolved independently from the rest of the world due to its geographical isolation (Ignjatovic *et al.*, 2006), many different IB variants have been isolated and characterised since the early 1960s (Cumming, 1963), and *in vivo* protection studies were performed with these variants (Klieve & Cumming, 1988). Using both monoclonal antibodies directed against the major IBV proteins and sequencing studies, several distinct lineages have now been recognised (Ignjatovic *et al.*, 1997; Ignjatovic *et al.*, 2006), all different from those found elsewhere. Interestingly, in the nearby country of New Zealand, IB problems were uncommon before



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the 1970s, when IB variants were first reported (Lohr, 1988). It was initially believed, using cross-neutralisation tests, that IB variants had evolved independently of those reported in Australia or US, and at least 4 different variants were identified (Lohr, 1976, 1977). However, sequencing of the S1 gene has recently revealed genetic relationships between these early IB isolates and ones made since 2000, and phylogenetic analysis also showed that they are more closely related to Australian than to European or North American isolates (McFarlane & Verma, 2008).

The above account highlights the large number of IB variants that exist worldwide; some being unique to a particular area; others having a more general distribution. The origin of these variants is not clear, but Shieh *et al.* (2004), reporting the close relationship of isolates from Taiwan and Japan to IBVs found in Australia and USA, suggested that the Asian variants are recombinants; their S gene being derived from Australian variants and the N genes from US strains.

#### **CURRENT SITUATION**

As discussed, it is becoming clearer that many countries have to deal with many types of IBV. At the moment genotyping is by far the most used system and has largely replaced serotyping and protectotyping. Does this create a problem? The preferred typing system depends on the goal (e.g. selection of vaccination programmes, or epidemiological studies), available techniques, experience, field situation and costs.

Classification systems are divided into two major groups: functional tests, that regard the biological function of a virus, and non-functional tests, that look at the viral genome (De Wit, 2000). Typing by functional tests results in protectotypes and antigenic types (serotypes and epitope-types). Tests that look at the genome result in genotypes. With protectotyping, the complete immune response of a chicken against an IBV strain is measured. For the field, grouping of IBV strains into protectotypes is the most important system from a practical point of view, because it provides direct information about the efficacy of a vaccine. Strains that induce protection in chickens against each other belong to the same protectotype. Protectotyping is laborious and expensive, and requires high level facilities for performing vaccination-challenge studies and SPF

Serotyping is based on the reaction between an IBV strain and chicken-induced IBV serotype-specific

antibodies. Two strains (A and B) are considered to be of the same serotype when two-way heterologous neutralisation titres (antiserum A with virus B, and antiserum B with virus A) differ less than 20-fold from the homologous titres (antiserum A with virus A, antiserum B with virus B) in both directions (Hesselink, 1991). Serotyping becomes less practical when more IBV types are detected in a certain area, as every serotype needs its own neutralization test. For new strains that appear to be different, an antiserum has to be produced in SPF birds. As mentioned before, more and more countries have to deal with an increasing number of variants, which decreases the practicability of serotyping.

Grouping of strains based on genetic characterisation of (a part of) the genome results in genotypes. Methods include sequencing, detection of genotype-specific parts of the genome by RT-PCR, or determination of the position of enzyme cleavage sites (De Wit, 2000). Genomic information is objective and provides essential information for epidemiological studies. The part of the gene that encodes the S1 subunit of the spike glycoprotein is most frequetly used for genotyping. The S1 subunit of the S protein is the main inducer of protective immunity and carries most of the virus neutralizing epitopes, including serotypespecific epitopes, which are usually conformationdependent (Mockett et al., 1984; Cavanagh & Davis, 1986; Koch et al., 1990; Cavanagh et al., 1992). A change of a few (percentage of) amino acids in the S1 protein can already result in a change of serotype (Cavanagh et al., 1992) due to a change in a virusneutralizing epitopes, whereas other larger percentages of mutations at other parts of S1 might not result in relevant change in antigenicity of the virus. On the other hand, IB viruses of different serotypes and genotypes not only have different epitopes, but also share common epitopes, which are of importance in cross-immunity (Cavanagh et al., 1992; Cavanagh et al., 1997) and cell-mediated immune responses (Boots et al., 1992; Ignjatovic & Galli, 1995). These aspects of the IB virus result in a disadvantage of the use of genotyping in the field, as the direct translation of information about usually of a part of the genome (usually a part of the S gene) of an IBV strain into biological function or antigenicity of the virus is not possible or is not without risk. Despite these limitations, a few papers have reported that the S1 gene sequence (part of 700 nucleotides) comparison was a better predictor of immune challenge in chickens than serotyping by virus neutralization (Ladman et al., 2006).



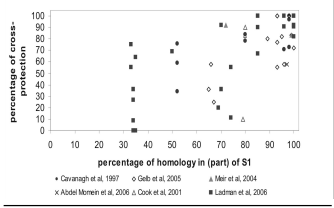
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Whether this is a general rule is unknown, as only a small number of strains and vaccines have been compared, and also different parts of the S1 gene were used for the comparison of the homology. But in general, a lower homology in the sequence of the S1 subunit of 2 strains (e.g. a vaccine and a field strain) means that there is more chance that relevant mutations have occurred, what might result in a lower cross-protection.

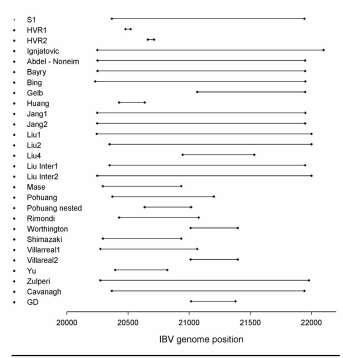
An analysis of several papers reporting the level of homology of the S1 gene or a part of it and the level of cross-protection (Cavanagh et al., 1997; Cook et al., 2001; Meir et al., 2004; Gelb et al., 2005; Ladman et al., 2006) showed that there is a higher chance of good level of cross protection between strains with a high level of homology than between strains with a low homology (Figure 1). However, these data also show that the relation is not very strong. Some strains that differ only by a few percentage points from the other strain in the sequenced part of the genome showed a significant drop in cross-protection (Meir et al., 2004; Abdel-Moneim et al., 2006), whereas there was a high level of cross-protection against other strains with a much lower homology (Meir et al., 2004). Figure 1 also shows the wide variation in the level of cross-protection detected in strains with the same level of homology in comparison with the strains that are used as vaccines. Another limitation that needs to be considered is the different parts of S1 that different laboratories and research groups use for their analyses of level of homology, due to the wide variation of primers used in PCR tests (Figure 2). Several laboratories sequence a part of S1 that includes the highly variable regions (HVR) 1 and 2; others do not. Analysis of different parts of S1 can result in different levels of homology. When the IBV strain tested is the result of a recombination event between different IB genotypes, the examination of different parts of S1 may mean the detection of a different genotype (Wang et al., 1993; Jia et al., 1995; Ammayappan et al., 2008; Dolz et al., 2008). The detected homology with other strains is then very dependent on the part of the genome that is being used for the comparisons of the homology. Using larger parts of S1 for the comparison of strains reduced the risk of finding high levels of homology between strains where there are not.

This leads to the conclusion that genotyping is an excellent tool for epidemiological studies (Figure 3), and that is a convenient, practical tool for typing that best can be used as a screening tool to select potential relevant strains. Especially when there is suspicion in

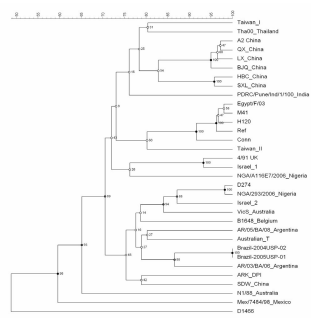
the field that the genotype of recent isolates does not provide accurate information about the true antigenic nature of these IBV isolates, conventional testing (serotyping) and especially *in vivo* studies are required.



**Figure 1** - Correlation between the level of homology in (a part) of the S1 glycoprotein of IBV strains and the level of cross-protection between these strains as reported in six papers.



**Figure 2** - Examples of sequenced parts (place and length) of the IBV spike protein genome and HVR 1 and 2 used by different authors reporting levels of homology between certain IBV strains or the relation between level of homology between a vaccine and a challenge strain and the detected level of crossprotection. The purpose of this overview is not to compare individual (one or more) primer sets as reported by specific authors but to show the wide variation that is being used in IBV research, which can make it difficult to compare results of different workers, as discussed in this paper.



**Figure 3** - Phylogenetic tree of a selection of IBV variants from different parts of the world showing the wide diversity of IBV. The tree is based on comparison of the partial S gene (between nucleotides 20447 and 20924 (numbering compared to the genome of Ark DPI (Ammayappan *et al*, 2008), coding for a part of the S1 glycoprotein including the highly variable regions 1 and 2). The phylogenetic tree analysis was conducted by neighbour-joining method using bootstrap analysis (100 replications). The numbers along the nodes refer to the bootstrap values

#### **CONTROL MEASURES**

IBV is ubiquitous in most parts of the world where poultry are reared, and it is able to spread very rapidly in non-protected birds (De Wit *et al.*, 1998). It is shed via both the respiratory tract and the faeces and can persist in the birds and faeces for several weeks or months. Although strict biosecurity and working with a one-age system are essential control measures, vaccination is usually essential to increase the resistance of chickens against the challenge with IBV strains (Cook, 2008).

For vaccination of chickens against IBV, both live attenuated and inactivated (usually oil-adjuvanted) vaccines are used. Live vaccines are used particularly in young birds to achieve early protection against challenge, and also for priming of future layers and breeders that are going to be later boosted with the inactivated vaccines. In areas with an increased level of field challenge, live attenuated vaccines are also periodically used during the laying period with the intention of keeping a high level of local protection of the respiratory tract.

Most vaccines used in the world are of the Mass serotype. In several parts of the world, Mass vaccines are the only vaccines allowed, but elsewhere vaccines of one or more other serotypes are permitted. Vaccines of a certain serotype or genotype are normally able to protect well-vaccinated chickens against a homologous challenge. Often, there is partial protection against strains of other protectotypes, serotypes or genotypes that can vary from high to low (Figure 1 and reviewed for the H-strain by Bijlenga et al. (2004). The magnitude and duration of the response to vaccination is dependent on many factors, including age of the chick, levels of maternal immunity, immunogenicity of the vaccine, method of vaccine application, virulence of the field strain challenge, interval between vaccination and challenge, and immunocompetency of the host. Chickens vaccinated under optimal conditions may be immune for many months, and for broilers, this may be life-long (Bijlenga et al., 2004).

It has been shown that vaccination with 2 antigenically distinct live-attenuated vaccines such as Mass and 4/91, can result is a broad cross-protection against many different IBV types (Cook *et al.*, 1999; Terregino *et al.*, 2008). The cross-protection was broader when these vaccines were applied with a 2-week interval than when the vaccines were combined on the same day.

Results of challenge studies and field work have shown that vaccination with a bivalent vaccine containing the Mass and Ark strains (Gelb et al., 1989; Gelb et al., 1991; Gelb et al., 2005) provided, in average, higher level of cross-protection against certain heterologous field strains than other combinations of vaccines, such as Mass together with Conn or Mass with JMK. The Mass and Ark vaccine did not provide significant protection against challenge with another strain (Ladman et al., 2002). It is not known whether the separate application of these 2 IBV strains would have resulted in a higher (or lower) level of cross-protection against the same heterologous challenges.

A well-vaccinated chicken is protected against challenge with a virulent homologous IBV strain. This means that this well-vaccinated chicken is also protected against an early revaccination with a homologous vaccine. Neverthelless, revaccination of young birds, especially broilers, using a vaccine of the same serotype as the first vaccine, has proven to be beneficial under field conditions. This is an indication that the quality of the first vaccination may need careful attention. Whatever live vaccine is used, application is a very critical step. IBV is a sensitive virus that can be



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easily inactivated (Cavanagh & Gelb, 2008), which may produce inadequate vaccination results under field conditions (De Wit et al., 2010). It can be applied by eye- or nasal drop, spray, and drinking water route. It is essential that a high percentage of the birds receive a required dose of the vaccine in the right place. Inadequate uptake of the vaccine may result in no or decreased level of protection, delayed protection, prolonged presence/circulation of the vaccine virus in the flock, resulting in an increased risk of bacterial infections with E.coli or other bacteria (Goren, 1978; Smith et al., 1985; Cook et al., 1986; Matthijs et al., 2003) and even an increase of virulence of the virus (Hopkins & Yoder, 1986)

In order to achieve higher level of protection of commercial layers and parent stock during the laying period, the use of inactivated IBV vaccines after a priming with live IBV vaccines has been shown to be effective against homologous Mass challenges (Gough et al., 1977; Box et al., 1980; Timms & Bracewell, 1983). The efficacy of increasing the level of protection against heterologous challenges in the laying period has rarely been reported, although birds that had been vaccinated twice with a live Mass type vaccine and boosted with a killed oil-emulsion vaccine containing a Mass strain showed no protection against challenge with a Ark type strain (Muneer et al., 1987b).

#### CONCLUSION

The topic of the how to type the large number of IB variants currently found worldwide and relate this to the best vaccination strategy to use to protect against them is clearly complex. Whilst genotyping has advantages of ease and speed, the number and variety of different primers used by different research groups makes interpretation very difficult, and this paper emphasises the need of a standardised method of performing genotyping. Another major point that needs to be remembered is that only the vaccinated chicken decides whether genetic differences as shown by genotyping or serotyping are relevant for the level of cross-protection.

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