Characterization of monoclonal antibodies to bovine enteric 
coronavirus and antigenic variability among Quebec isolates

Brief Report

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Summary. Twenty monoclonal antibodies (MAbs) were prepared against the 
Mebus strain of bovine enteric coronavirus, 14 of them reacting with the pe-
plomeric S (gp 100) glycoprotein. Competition binding assays allowed the def-
inition of at least 4 distinct antigenic domains for the S glycoprotein, designated 
as A, B, C, and D; epitopes associated to neutralizing activity being located in 
sites A, B, and C. One MAb directed to the hemagglutinin HE (gp 140/gp 65) 
glycoprotein inhibited the hemagglutinating activity of the virus, but had no 
neutralizing activity. Comparison of Quebec enteropathogenic BCV isolates 
using polyclonal antiserum and MAbs directed to the S glycoprotein confirmed 
their close antigenic relationship, but also revealed the occurrence of at least 
three distinct antigenic subgroups. Antigenic domain D was highly conserved 
among BCV isolates, as well as non-neutralizing epitopes assigned to antigenic 
domains A and C. The Minnesota strain of turkey enteric coronavirus could 
be distinguished from BCV isolates by MAbs directed to epitopes of antigenic 
domain C, whereas human coronavirus HCV-OC 43 could be distinguished by 
MAbs directed to epitopes of antigenic domain A. The porcine hemagglutinating 
encephalomyelitis virus could be distinguished from the other hemagglutinating 
coronaviruses by neutralizing epitopes located on antigenic domains A, B, and 
C.

Bovine coronavirus (BCV) is a member of the family Coronaviridae, which 
replicates in the differentiated enterocytes of the small and large intestines, 
causing severe diarrhea in newborn calves [18], winter dysentery (hemorrhagic 
enteritis) or chronic shedding in adult cattle [4, 21]. The virus is also responsible
for upper respiratory tract illness in growing calves [20]. The virion contains four major structural proteins: the 52 k nucleocapsid (N) protein, the 24–26 k transmembrane or matrix (M) glycoprotein, the 180–200 k spike (S) glycoprotein, and the 125–140 k hemagglutinin/esterase (HE) glycoprotein [14, 17]. The S glycoprotein often is posttranslationally cleaved by host-cell proteases into two 100 k fragments, S1 and S2, respectively, corresponding to the N- and C-terminal subunits [1, 10]. The HE glycoprotein is a disulphide-linked dimer of 65 k subunits and is associated to additional small granular projections located at the base of the typical large bulbous peplomers of the hemagglutinating coronavirions [9, 10, 17]. Both S and HE interact with receptors on the cell surface [23, 24] and trigger the immune system eliciting the production of neutralizing antibodies [8, 11, 26].

On the basis of serological studies, BCV has been classified in the subgroup of hemagglutinating coronaviruses, including the hemagglutinating encephalomyelitis virus of swine (HEV), the human respiratory coronavirus HCV-OC 43, the turkey coronavirus (TCV), and the various strains of mouse hepatitis virus [8, 9, 25]. These viruses all share antigenic determinants located on their homologous structural proteins [14]. Comparison of the nucleotide sequences of the structural genes and molecular hybridization studies with cDNA probes confirm this antigenic classification [1, 3, 22, 28, 30]. Although there exist BCV strains that can be distinguished for their pathogenicity, there is still some controversy as to the existence of distinct BCV serotypes. Polyclonal antisera and monoclonal antibodies (MAbs) that were raised against a BCV strain (S2) isolated in Scotland detected only minor strain variations among five cell culture-adapted strains of BCV by both indirect immunofluorescence (IIF) and hemagglutination inhibition tests (HAI) [12]. Similar findings were also reported by counterimmunoelectrophoresis and immunodiffusion tests [5]. More recently, anti-S MAbs were found to distinguish between vaccine and wild-type BCV strains in mildly denaturating Western and neutralization assays [15]. Two neutralizing epitopes appear to be conserved in virulent and avirulent strains, while two epitopes of the avirulent strains are not detected in the wild-type strains; non-neutralizing epitopes appear to be conserved among the various isolates [12, 15]. We report here the production of MAbs against the Mebus strain of BCV and the detection of antigenic plurality among bovine coronaviruses isolated from dairy cattle in Quebec through the reaction of MAbs against the S glycoprotein.

The cell culture-adapted Mebus strain [18] of BCV was obtained from the American Type Culture Collection (ATCC VR 874), Rockville, MD. The prototype BCV was originally isolated in bovine fetal kidney cells from diarrhea fluid of calf [18]. Our virus strain had been passaged five times in human rectal tumour (HRT-18) cells in the presence of 10 U/ml bovine pancreatic trypsin, as previously described [6, 8]. Twelve other BCV isolates were recovered from clinical cases of epidemic diarrhea in newborn calves in Quebec dairy herds from January 1987 to May 1990. No commercial BCV vaccine had been applied
MAbs against bovine enteric coronavirus

in these herds during the year preceding these outbreaks. We received the Minnesota strain [6] of TCV from Dr. B. S. Pomeroy of the University of Minnesota, St. Paul, MN, and HCV-OC43 [14] from Dr. P. J. Talbot, Institute Armand-Frappier, Laval, Qc, Canada. The 67N strain [14] of HEV was obtained from the ATCC (VR-740). All these hemagglutinating coronaviruses were serially propagated in HRT-18 cells in the presence of trypsin [6, 8].

Under the following conditions, the yield of viral production after three to five successive passages was similar for the different isolates as revealed by IIF [8] or indirect immunoperoxidase [2], and calculation of the infectivity titers (ranged between 10^{5.5} and 10^{6.5} TCID 50/ml). The extracellular virions of the various BCV isolates were purified by differential and isopycnic ultracentrifugation on continuous 20 to 55% (v/v) sucrose gradients, as previously described [6]. Electron microscopic examination of purified viral preparations revealed that most viral particles possessed the double fringe of surface projections [9].

Monoclonal antibodies to the Mebus strain of BCV were produced from BALB/c mice, as previously described [7]. Hybrid cells were cultured in HAT-medium containing 10% fetal calf serum and 10% Boehringer Mannheim-condimed H1 medium (supernatant of a mouse lymphoma cell line supplemented with 1 mmol/l oxaloacetate, 1 mmol/l sodium pyruvate, 0.2 μg/ml insulin, 10 ng/ml PMA, pH 7.6) in replacement of feeder layers of mouse macrophages. The supernatant from growing hybridomas was screened for anti-BCV antibody production by both IIF and ELISA. Antibody producing hybridomas were terminally diluted twice before inoculation into peritoneal cavity of pristane (2,6,10,14-tetramethylpentadecane) primed mice. Purification of immunoglobulins (IgG) from ascitic fluid, and initial characterization of MAbs and their titration by virus neutralization, ELISA, or HAI tests were also done as previously described [7].

Administration of BALB/c mice with either purified whole virus or NP 40-denaturated virus prepared from the supernatant fluids of BCV-infected HRT-18 cells induced high specific humoral response. Titers of serum from mice providing the immune cells ranged from 1280 to 5120 by IIF, while titers higher than 256,000 were obtained by ELISA. Of 64 hybridoma cell lines initially secreting anti-BCV MAbs, as determined by IIF and ELISA, 20 could be subcloned, serially propagated, and used to produce ascitic fluids. Log_{10} ELISA titers of clarified ascitic fluids ranged from 2.9 to 6.2.

The polypeptide specificity of the anti-BCV MAbs purified from ascitic fluids was determined by Western immunoblotting [7, 8]. The four major proteins described previously for BCV isolates [14, 17] were identified by using homologous hyperimmune rabbit serum. Under the conditions used, 14 hybridomas produced MAbs that recognized the S glycoprotein. These MAbs reacted either with gp 200 or gp 100, or both protein species (data not shown). Only one hybridoma produced MAbs reacting to the HE glycoprotein. In addition,
four MAbs reacted specifically to the N protein and another reacted to the M glycoprotein.

In order to correlate biological functions with polypeptide specificities, anti-BCV MAbs were studied for their ability to neutralize the virus and to inhibit its hemagglutinating activity. Characterization of the isotypes of the MAbs showed that most antibodies belonged to the IgG 2A, IgG 2B or IgG 1 subclass. Neutralization studies revealed that five of the MAbs have a strong neutralizing activity (VN titers ranging between 2560 and 10240), whereas three others had a lower neutralizing activity (VN titers ranging from 160 to 640). All these MAbs were identified as anti-gp100 MAbs by Western immunoblotting (Table 1). Six of the anti-S MAbs were devoid of neutralizing activity; they were directed either to the gp 100 or gp 200 glycoproteins. None of the MAbs to the N or the M structural proteins neutralized virus infectivity in vitro. The monoclonal BCA 3 that reacted to gp 125–140 inhibited the HA activity of the virus, but did not show neutralizing activity.

A direct relationship exists between the avidity and the maximal amount of MAb bound to a given amount of antigen [13, 19]. The anti-S MAbs were titrated in an indirect ELISA and the avidities were estimated using the plateau level absorbance values; they varied between 0.80 and 1.50 (Fig. 1). At a concentration of 0.01 µg of gammaglobulin per well, only MAbs BCA 2, BCF 4, BCB 5, BCB 1, and BCA 3 were able to saturate those viral antigens coated. The MAbs BCA 7, BCH 1, BCH 5, and BCH 8 did not reach a plateau level within the range studied, thus suggesting significantly lower relative avidities for the S glycoprotein.

In order to determine whether the anti-S MAbs could be divided into clusters recognizing the same immunogenic region of the peplomer protein, competition immunoassays were conducted according to Deregt et al. [11]. Affinity purified antibodies were conjugated to D-biotin N-succinimide ester (molecular ratio 90:1) according to the method described by Kendall et al. [16]. Conjugation had no apparent adverse effect on the binding properties of ten of the purified anti-S MAbs and endpoint titers ranged from approximately $10^4$ to $10^5$. To define overlapping antibody-binding sites, competition was considered to be positive only if it occurred reciprocally and over a range of several log_{10} dilutions; each antibody was used both as competitor and biotinylated probe [11, 19].

The results of the competition binding experiments conducted with these ten anti-S MAbs are summarized in Table 1. By their reactivity in competition binding assays with biotinylated MAbs, anti-S MAbs fell into four competitive binding groups, A, B, C, and D, and between individual members of each group the competition was generally strong and reciprocal. An epitope map showing the main antigenic domains delineated on the BCV peplomeric S glycoprotein was deduced from the competition binding experiments and is illustrated in Fig. 2. Epitopes associated with the neutralizing activity are located on sites A, B, and C. Antigenic domains A and B appeared to be associated with a strong neutralizing activity and were recognized by MAbs BCF 4, BCB 1, and BCH 5,
| Competitor antibody | Competition with biotinylated probe antibody$^b$ |
|---------------------|---------------------------------------------|
| Clone no. | Ig isotype | SN$^a$ titers | BCA 2 | BCH 8 | BCA 1 | BCH 3 | BCA 3 | BCB 5 | BCF 4 | BCB 1 | BCA 7 |
| BCA 2 | G1 | 320 | ++++ | ++++ | ++++ | ++++ | - | - | - | - | - |
| BCH 8 | G1 | 640 | ++++ | ++++ | ++++ | ++++ | - | - | - | - | - |
| BCA 1 | G 2a | <20 | ++++ | ++++ | ++++ | ++++ | + | - | - | - | - |
| BCH 3 | G 2a | <20 | ++++ | ++++ | ++++ | ++++ | + | - | - | - | - |
| BCA 3 | G 1 | 10240 | - | - | - | ND | ++++ | ++++ | - | - | - |
| BCB 5 | G 2b | 10240 | - | - | + | ND | ++++ | ++++ | - | - | - |
| BCH 5 | G 2a | <20 | - | - | - | - | + | - | ++++ | ++++ | - |
| BCF 4 | G 2a | 10240 | - | - | - | - | - | - | ++++ | ++++ | - |
| BCB 1 | G 2a | 10240 | - | - | - | - | - | - | ++++ | ++++ | - |
| BCA 7 | G 2b | <20 | - | - | - | ND | - | - | - | - | + - + |

$^a$ Log$_{10}$ of the highest dilution of ascitic fluid neutralizing 100 TCID$_{50}$ of virus

$^b$ Percentage competition: ++++ > 75%, ++ 25 to 75%, - < 25% and if it occurred over fewer than three ten-fold dilutions. By definition, binding of each conjugate to viral antigen in the absence of competitor represents 100% binding (0% competition), while the binding of the conjugate following prior incubation with excess of the same unlabeled MAb represents 0% binding (100% competition) [11]. The percentage of competition usually reached a plateau level over a range of three to four ten-fold dilutions

ND Not done
Fig. 1. Comparative avidity of monoclonal antibodies directed against the BCV (Mebus strain) peplomeric S glycoprotein. Serial ten-fold dilutions of purified antibody were incubated in the wells of microtiter plates containing absorbed purified BCV virions (0.5 µg of protein). After incubation for 90 min at 37 °C, binding to the viral antigen was detected by the addition of peroxidase-labeled goat anti-mouse IgG. The resulting A 450 value was read and used to rate MAbs for avidity.

Fig. 2. Antigenic map of BCV (Mebus strain) peplomeric S glycoprotein deduced from competitive binding assays. A, B, C, and D Four antigenic sites defined by the anti-S (gp 100) monoclonal antibodies. The MAbs defining each site are indicated in the corresponding circle. Antigenic site B is partly related to antigenic sites A and C via common non-neutralizing epitopes (shadowed). The neutralizing MAb BCH 1, an antibody of apparent low avidity, could not be assigned to a particular antigenic group since a reciprocal competition experiment could not be done due to the lack of biotinylated probe. However, the neutralizing epitope defined by this MAb has been found to be conserved among all BCV isolates analysed and for this reason was assigned to another antigenic site. The antigenic sites on the peplomeric glycoprotein of BCV were designated according to the nomenclature used by Deregt et al. [11] and Vautherot et al. [27].
and MAbs BCB 5 and BCA 3, respectively. The anti-S MAbs having a weak neutralizing activity (BCA 2, BCH 8) or devoid of neutralizing activity (BCA 1 and BCH 3) defined antigenic domain C. The anti-S reactivity of MAbs BCA 7 could not be blocked by the other MAbs and for this reason it was suggested to be directed against a distinct non-neutralizing antigenic domain. Furthermore, antibodies BCA 1 and BCH 3, and BCH 5, which were assigned to antigenic domains C and A, showed reciprocal competition with monoclonal BCA 3 assigned to antigenic domain B. This suggests the presence of overlapping regions among major antigenic domains of the S glycoprotein that are involved in the neutralizing activity.

The BCV anti-S MAbs were tested for their cross-reactivity with other hemagglutinating coronaviruses and were applied to analyse bovine coronaviruses isolated from dairy cattle in Quebec. In preliminary experiments using polyclonal rabbit antisera, the relationship among HCV-OC-43, HEV, TCV, and BCV was confirmed by IIF and Western immunoblotting; by HAI and neutralization assays, only TCV and BCV appeared to be serologically related [8]. Furthermore, using polyclonal antisera, there was no evidence from this study to warrant distinction of the various Quebec BCV isolates into classical serotypes (data not shown).

The cross-reactivity of BCV anti-S MAbs was tested in an indirect ELISA [7]. Twice the amount of MAb giving maximal binding to the Mebus strain of BCV was used for antigenic comparison. As illustrated in Fig. 3, binding to the heterologous virus strains was expressed as a percentage of the absorbance values obtained with the reference virus. Overall, at least three antigenic groups could be recognized among the BCV isolates studied. Two Quebec isolates of BCV (BCQ 4 and BCQ 718) reacted similarly to the Mebus strain with the 12 MAbs tested. The MAbs BCH 6, BCH 1, and BCA 7 did not show significant variability among the various Quebec BCV isolates, and bound intensively to the three other hemagglutinating coronaviruses tested. The MAbs BCB 1, BCF 4, BCB 5, and BCA 3 reacted with a very low intensity against eight of the Quebec isolates. Among the latter, six reacted strongly with the other MAbs. The BCV isolates BCQ 9, BCQ 12, and BCQ 2070, and HEV exhibited no reactivity to the four previous MAbs and to MAbs BCH 3, BCA 1, BCH 8, and BCA 2. Four of the anti-BCV MAbs (BCH 5, BCB 1, BCF 4, and BCH 3) did not react against HCV-OC-43. The Minnesota strain of TCV reacted strongly with most of the anti-BCV MAbs; this virus appeared to differ from the reference BCV strain only by its lower reactivity to MAbs BCH 3, BCH 8, and BCA 2 (50% of the value obtained with the homologous virus).

In the present study, data obtained in the competition binding experiments with BCV anti-S MAbs were in agreement with previous findings reported in cases of an avirulent (BCQ 2) and a virulent (G110) BCV strain [11, 26]. However, a more extensive MAb library is required to establish more definitive spacial and operational mapping of the antigenic determinants carried by the S glycoprotein of the BCV virion. Vautherot et al. [27] have identified a total of
Fig. 3. Cross-reactivity of monoclonal antibodies to the S glycoprotein of the Mebus strain of BCV with Quebec BCV isolates. The cross-reactivity between the various isolates was tested by an indirect ELISA. Twice the amount of MAb giving maximal binding to the Mebus strain was used. Binding to the heterologous virus is expressed as a percentage of the $A_{450}$ value obtained with the homologous virus. □ 50 to 100%; ☐ 25 to 50%; ■ less than 25%

seven antigenic domains for the S glycoprotein of BCV by means of a competitive binding assay; four independent antigenic domains were found to be located on S1, two on S2, and one on gp200. As demonstrated in the present study, most of the neutralizing-mediating determinants appeared to cluster in two major antigenic domains of the S1 subunit, A and B, the latter being subdivided into subsites [27]. Recently, BCV anti-S MAbs BCB5, BCA3, BCF4, and BCB1 that possess a strong neutralizing activity were found to react against the spike glycoprotein S1 subunit, expressed by recombinant baculovirus [29; Parker M.D., pers. comm.].

Comparison of Quebec enteropathogenic BCV isolates using polyclonal antisera and MAbs directed to the S glycoprotein confirmed their close antigenic relationship, but also revealed the occurrence of at least three distinct antigenic subgroups. These subgroups were identified by neutralizing MAbs directed to epitopes of antigenic domains A, B, and C. Antigenic domain D of the S glycoprotein, defined by MAb BCA7, appeared to be highly conserved among Quebec BCV isolates, as well as a non-neutralizing epitope assigned to antigenic domain A. Three Quebec BCV isolates differed from the other by neutralizing and non-neutralizing epitopes assigned to antigenic domain C. Another neutralizing MAb (BCH1), that could not be assigned to the aforementioned antigenic domains, was found to be directed to an epitope highly preserved
among the various BCV isolates, as well as the three other hemagglutinating coronaviruses tested. The Minnesota strain of TCV, which could not be differentiated from BCV by virus neutralization tests using polyclonal rabbit or turkey hyperimmune sera [8], was found to differ from the latter by epitopes of antigenic domain C. These epitopes appeared to be associated with weak neutralizing activity. Epitopes of antigenic domains A and B, associated to a strong neutralizing activity, appeared to be highly conserved among both viruses, thus confirming previous reports on the close antigenic and genomic relationship between the turkey and bovine enteropathogenic coronaviruses [8, 9, 28].

On the other hand, the HCV-OC 43 could be distinguished by neutralizing MAbs directed to epitopes of antigenic domain A. These results are in agreement with previous findings showing that non-neutralizing anti-gp 200 MAbs map a unique site located on the S2 part of the peplomer glycoprotein and react similarly with various hemagglutinating coronaviruses, whereas anti-gp 105 MAbs, that defined at least four independent neutralizing or non-neutralizing domains on the S1 part of the peplomer glycoprotein, react only with the homologous virus [26]. Using polyclonal hyperimmune sera, we were unable to show cross-reactivity by virus neutralization between the human and bovine hemagglutinating coronaviruses, but common epitopes located on their various structural proteins were recognized by Western immunoblotting [14]. Recently, high degrees of homology were demonstrated for the nucleotide sequence of the S and HE genes of both viruses [31].

The porcine hemagglutinating encephalomyelitis virus could be distinguished from the other hemagglutinating coronaviruses tested by neutralizing epitopes located on antigenic domains A, B, and C. These results are also in agreement with those obtained by Vautherot et al. [26] who have demonstrated that HEV can be differentiated from the other hemagglutinating coronaviruses by its reactivity to epitopes located on the various antigenic domains of the S1 part of the peplomer glycoprotein.

Recently, comparison of the S gene sequences of six highly virulent or avirulent BCV isolates have revealed more than 98% homologies in spite of different origins of the viruses [15, 31]. Overall, there were only 45 to 56 nucleotide differences between the virulent and avirulent groups, while there were 6 to 14 differences among four avirulent strains. It has been speculated that substitutions of few amino acids in the putative fusogenic domains and two proline at position 507 and 567 in the antigenic domains may cause altered immunogenic and other functional properties of the S proteins specified by the virulent and avirulent BCV strains [31]. Nucleotide sequence analysis of the portion of the S1 gene corresponding to the two major neutralizing domains are presently in progress in order to establish the significance of the differences observed among the Quebec BCV isolates.

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