

An outbreak of human coronavirus OC43 infection and serological cross-reactivity with SARS coronavirus

David M Patrick MD¹, Martin Petric PhD¹, Danuta M Skowronski MD¹, Roland Guasparini MD², Timothy F Booth PhD³, Mel Krajden MD¹, Patrick McGeer MD PhD⁴, Nathalie Bastien PhD³, Larry Gustafson MD², Janet Dubord MD², Diane MacDonald MSc¹, Samara T David MSc³, Leila F Srouf MD³, Robert Parker MD², Anton Andonov PhD³, Judith Isaac-Renton MD¹, Nadine Loewen MD², Gail McNabb BSc¹, Alan McNabb BSc¹, Swee-Han Goh PhD¹, Scott Henwick MD⁵, Caroline Astell PhD⁶, Jian Ping Guo PhD⁴, Michael Drebot PhD³, Raymond Tellier MD⁷, Francis Plummer MD³, Robert C Brunham MD¹

DM Patrick, M Petric, DM Skowronski, et al. An outbreak of human coronavirus OC43 infection and serological cross-reactivity with SARS coronavirus. *Can J Infect Dis Med Microbiol* 2006;17(6):330-336.

BACKGROUND: In summer 2003, a respiratory outbreak was investigated in British Columbia, during which nucleic acid tests and serology unexpectedly indicated reactivity for severe acute respiratory syndrome coronavirus (SARS-CoV).

METHODS: Cases at a care facility were epidemiologically characterized and sequentially investigated for conventional agents of respiratory infection, SARS-CoV and other human CoVs. Serological cross-reactivity between SARS-CoV and human CoV-OC43 (HCoV-OC43) was investigated by peptide spot assay.

RESULTS: Ninety-five of 142 residents (67%) and 53 of 160 staff members (33%) experienced symptoms of respiratory infection. Symptomatic residents experienced cough (66%), fever (21%) and pneumonia (12%). Eight residents died, six with pneumonia. No staff members developed pneumonia. Findings on reverse transcriptase-polymerase chain reaction assays for SARS-CoV at a national reference laboratory were suspected to represent false positives, but this was confounded by concurrent identification of antibody to N protein on serology. Subsequent testing by reverse transcriptase-polymerase chain reaction confirmed HCoV-OC43 infection. Convalescent serology ruled out SARS. Notably, sera demonstrated cross-reactivity against nucleocapsid peptide sequences common to HCoV-OC43 and SARS-CoV.

CONCLUSIONS: These findings underscore the virulence of human CoV-OC43 in elderly populations and confirm that cross-reactivity to antibody against nucleocapsid proteins from these viruses must be considered when interpreting serological tests for SARS-CoV.

Key Words: *Coronavirus; Human coronavirus OC43; Outbreak; Respiratory infection; SARS-CoV; SARS coronavirus; Severe acute respiratory syndrome*

Une flambée de coronavirus OC43 humain et une réactivité croisée sérologique avec le coronavirus du SRAS

HISTORIQUE : Pendant l'été 2003, une flambée de troubles respiratoires a été étudiée en Colombie-Britannique, au cours de laquelle des tests des acides nucléiques et une sérologie ont indiqué, contre toute attente, une réactivité au coronavirus du syndrome respiratoire aigu sévère (CoV-SRAS).

MÉTHODOLOGIE : Les cas d'un établissement de soins ont subi une caractérisation épidémiologique et un examen séquentiel afin de dépister les agents classiques d'infection respiratoire, le CoV-SRAS et d'autres CoV humains. La réactivité croisée sérologique entre le CoV-SRAS et le CoV humain-OC43 (CoVH-OC43) a fait l'objet d'un examen par titrage de zones de peptides.

RÉSULTATS : Quarante-vingt-quinze des 142 résidents (67 %) et 53 des 160 membres du personnel (33 %) ont eu des symptômes d'infection respiratoire. Les résidents symptomatiques souffraient de toux (66 %), de fièvre (21 %) et de pneumonie (12 %). Huit résidents sont morts, dont six de pneumonie. Aucun membre du personnel n'a souffert de pneumonie. On soupçonnait que les résultats des titrages de RT-PCR du CoVH-SRAS à un laboratoire de référence national comportaient des faux-positifs, mais ce facteur était confondu par le dépistage sérologique concomitant d'anticorps à la protéine N. Des tests subséquents par RT-PCR ont confirmé le CoVH-OC43. La sérologie des convalescents a permis d'écartier le SRAS. Il était frappant de souligner que les sérums ont démontré une réactivité croisée contre les séquences de peptides nucléocapsides communes au CoVH-OC43 et au CoV-SRAS.

CONCLUSIONS : Ces résultats soulignent la virulence du CoVH-OC43 dans les populations âgées et confirme qu'il faut envisager une réactivité croisée à l'anticorps contre les protéines de nucléocapside de ces virus au moment d'interpréter les tests sérologiques du CoV-SRAS.

¹University of British Columbia Centre for Disease Control; ²Fraser Health Authority, Vancouver, British Columbia; ³Public Health Agency of Canada, Winnipeg, Manitoba and Ottawa, Ontario; ⁴Kinsmen Laboratory of Neurological Research, University of British Columbia, Vancouver; ⁵Surrey Memorial Hospital, Surrey; ⁶British Columbia Cancer Agency's Genome Sciences Centre, British Columbia; ⁷Hospital for Sick Children, Toronto, Ontario

Correspondence and reprints: Dr David M Patrick, University of British Columbia Centre for Disease Control, 655 West 12th Avenue, 2nd Floor, Vancouver, British Columbia V5Z 4R4. Telephone 604-660-3199, fax 604-660-0197, e-mail david.patrick@bccdc.ca

Received for publication August 4, 2006. Accepted September 22, 2006

On July 5, 2003, the World Health Organization (WHO) announced that human transmission of severe acute respiratory syndrome coronavirus (SARS-CoV) had ceased (1). Many drew the inference that the virus, whose causal role for SARS had been established (2-5), had been driven back into nature. From its emergence in November 2002, SARS was characterized as a lower respiratory syndrome involving pneumonia and/or acute respiratory distress syndrome (6,7). Transmission to others occurred most easily from severely affected patients, and was rare without fever (8).

Hence, a report of reactive tests for SARS-CoV on respiratory specimens submitted during an outbreak of generally mild respiratory infection accompanied by limited mortality in the elderly residents of a British Columbia long-term care facility during the summer of 2003 was unexpected (9). In the present article, the outbreak is described, the initial and subsequent laboratory findings are summarized, and the probable basis of serological cross-reactivity responsible for early findings is elucidated.

METHODS

Epidemiological methods

A case definition was established as "any resident or staff member who experienced symptoms of respiratory infection – rhinitis, sore throat, fever, cough or pneumonia – after July 1, 2003". Assessment of the extent and timing of symptoms was performed by chart review for residents of the facility and for patients transferred to hospital, and by interview of health care professionals. Records of staff absenteeism were examined, and interviews with ill workers were conducted. The accumulated findings were entered into a database (Microsoft Access, Microsoft Corporation, USA). Tabulation and construction of epidemic curves were accomplished in Microsoft Excel (Microsoft Corporation, USA).

Specimen collection

Nasopharyngeal, nasal and lower respiratory specimens were collected from a subset of symptomatic residents and transported to the University of British Columbia Centre for Disease Control (BCCDC) laboratories within 24 h. Specimens from the lower respiratory tract, stools and autopsy materials were obtained where available. Sera were collected from a subset of symptomatic residents in various stages of presentation. Sera collected before the 21st day following symptom onset were considered acute; after the 28th day, they were considered convalescent.

Microbiological testing

Respiratory specimens were tested for adenovirus, parainfluenza virus, respiratory syncytial virus, enterovirus and influenza virus by immunofluorescence microscopy and isolation in cell culture (10). Serological testing was carried out for *Mycoplasma pneumoniae*, *Chlamydomphila pneumoniae* and *Chlamydomphila psittaci*. Respiratory specimens were cultured for bacterial pathogens and, together with urine specimens, were evaluated for *Legionella* antigen using standard methods (10).

Nucleic acid-based testing

The reverse transcriptase-polymerase chain reaction (RT-PCR) assays for SARS-CoV were conducted at the National Microbiology Laboratory (NML) in Winnipeg, Manitoba, the BCCDC and the British Columbia Cancer Agency's Genome Sciences Centre (Vancouver, British Columbia); the results are summarized in Table 1. A detailed description of assays is available

as a technical appendix through personal communication with the authors. Of note, the 'pancoronavirus' PCR (BCCDC) targeted a 220-base pair region of the polymerase 1b gene that is highly conserved among CoVs (11). Amplicons from this assay were sequenced for phylogenetic analysis. RT-PCR was also performed for human metapneumovirus (hMPV) at the NML (12).

Serological methods

Methods used at the NML and BCCDC are summarized in Table 1. The BCCDC diagnostic algorithm required that specimens reactive by ELISA be confirmed using the neutralizing antibody test.

Assessing the potential for cross-reactivity

Multiple 10 amino acid peptides incorporating all the sequences in the SARS-CoV and human CoV (HCoV)-OC43 nucleocapsid proteins were prepared on derivatized cellulose membranes using a robotic peptide synthesizer (Autospot ASP 222, Intavis Bioanalytical Instruments, Germany) as previously described (13). The peptides overlapped by eight residues, with each subsequent peptide on a membrane being shifted by two amino acids toward the C-terminal end. There were 220 peptide spots prepared from the sequence of the HCoV-OC43 nucleocapsid protein and 207 spots from the corresponding SARS-CoV protein.

The membranes were blocked overnight at 4°C with 5% skim milk. They were then incubated with serum samples diluted 100-fold in buffer for 1 h at 37°C. After washing, membranes were treated with horseradish peroxidase-conjugated goat antihuman immunoglobulin G (Sigma, USA). The membranes were developed with enhanced chemiluminescence reagents and the positive spots were visualized on a Bio-Rad Fluor-S Multi-Imager (Bio-Rad Laboratories, USA).

RESULTS

Epidemiology

The outbreak occurred in a long-term care facility located in British Columbia. Residents were elderly or disabled adults in need of intermediate or extended care. From July 1 to August 22, 2003, 95 of 142 residents (67%) and 53 of 160 staff members (33%) experienced respiratory tract infections. The median age of affected residents was 83.7 years, with a range of 43.6 to 98.7 years. The epidemic curve is depicted in Figure 1. Peak incidence occurred between July 20 and 28, and the outbreak had tapered off by August 22, 2003. A floor-by-floor pattern of spread was not discernible, but a common area on the ground floor was used by most residents. Symptoms, findings and outcomes in residents and staff are summarized in Table 2. Most had mild clinical findings. Of the residents who met the case definition, 66% experienced cough, 21% fever, 66% rhinitis and 12% pneumonia. Twelve residents were admitted to hospital, eight for pneumonia. There were eight deaths among patients with respiratory symptoms. During the preceding six months, there was a mean of three deaths per month. None of the 53 affected staff members died, was hospitalized or developed pneumonia.

Microbiology findings

No pathogenic organisms were identified on routine testing from any of the residents. The exception was the finding of hMPV in respiratory specimens from four of 17 residents tested by RT-PCR. Molecular or serological testing for SARS-CoV was performed on specimens from 51 of

TABLE 1
Listing of diagnostic methods and key study findings

PCR assay	Laboratory	Gene target(s)	Results
RT-nested PCR for SARS-CoV	NML	Membrane glycoprotein and polymerase	Specimens from 9 of 40 patients were positive (two on repeat extraction)
TaqMan RT-PCR (Applied Biosystems, USA) for SARS-CoV	NML	Polymerase and nucleocapsid	3 of 9 specimens were positive
RT-PCR for SARS-CoV	BCCDC	Nucleocapsid	0 of 66 specimens from 39 residents were positive
Roche LightCycler PCR (Roche Diagnostics, USA) for SARS-CoV	BCCDC	Polymerase 1b	0 of 40 specimens from 25 residents were positive
'Pancoronavirus' PCR	BCCDC	Conserved region of polymerase 1b	14 of 39 residents positive in one or more specimen
RT-PCR for HCoV-OC43	BCGSC	Nucleocapsid and matrix	3 of 3 specimens positive
RT-PCR for HCoV-OC43	NML	Polymerase and nucleocapsid	4 specimens positive
Serological assay	Laboratory	Antigen/method	Results
ELISA for SARS-CoV antibody	NML	Infected cell lysate, baculovirus expressed N protein	6 of 35 acute specimens reactive for antibody to SARS-CoV by ELISA, neutralizing antibody or both. Three others with weak N band by WB. No positives by neutralizing antibody on convalescent testing
IFA for SARS-CoV antibody	NML	Infected cells	
Neutralizing antibody to SARS-CoV	NML	Plaque reduction on Vero E6 cells	
WB	NML	Baculovirus expressed N protein*	
Euroimmun indirect immunofluorescence (Medizinische Labordiagnostika, Germany)	BCCDC	<i>Escherichia coli</i> expressed HIS-tagged SARS-CoV nucleocapsid protein	0 of 35 specimens positive. Indeterminate NA assay of one in eight found in six patients. No confirmed positives with convalescent serology
WB	BCCDC	Infected cell lysate and recombinant N protein	
NA to SARS-CoV	BCCDC	Microneutralization test on Vero E6 cells	

*Data from references 3 and 7. BCCDC British Columbia Centre for Disease Control; BCGSC British Columbia Cancer Agency's Genome Sciences Centre; HCoV Human coronavirus; IFA Immunofluorescent antibody; NA Neutralizing antibody; NML National Microbiology Laboratory; RT-PCR Real-time polymerase chain reaction; SARS-CoV Severe acute respiratory syndrome CoV; WB Western blot

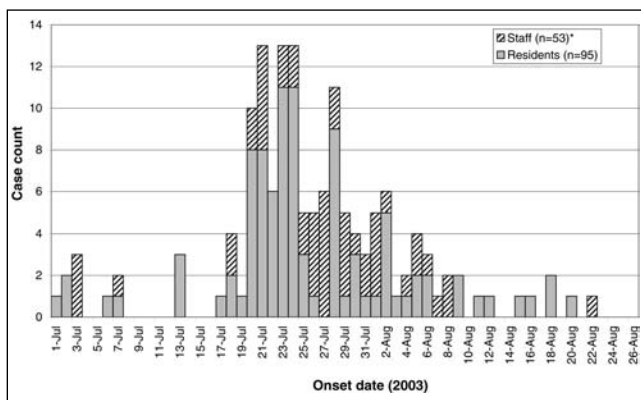


Figure 1) Epidemic curve: Onset of respiratory illness in a long-term care facility. *Onset dates for two staff members are missing

95 affected residents, the results of which are summarized in the final column of Table 1.

Findings from nucleic acid testing for SARS-CoV

Specimens from nine patients had a signal on one or more RT-PCR assays at the NML and were suspected of being false positives. These included two of the patients who had tested positive for hMPV by PCR at the NML. Analysis of the sequences of amplicons from three suspected false-positive PCR specimens at the NML (lung, stool and respiratory) showed homology to sequences of SARS-CoV (14,15). No specimens were positive for SARS-CoV by RT-PCR at the BCCDC.

Findings from nucleic acid tests for other CoVs

Table 1 summarizes the findings from nucleic acid tests for other CoVs. RT-PCR amplicons from specimens of nine of the 14 patients obtained by the 'pancoronavirus' assay were of concentrations adequate for sequencing. The following 163 base pair sequence internal to the primers was obtained from 12 specimens from these nine patients (including three who had tested positive for hMPV by PCR at the NML): TCGT-GCTATGCCAAACCTACTACGTATTGTTAG-TAGTTTGGTATTAGCCCCGAAAACATGAGACATGTTGCTCGCAAAGCGATAGGTTTTATCGACTTGC-GAATGAATGCGCACAAGTTTTGAGTGAAATTGT-TATGTGTGGTGGCTGTTATTATGTTAAGCCT. This sequence has a homology of over 99% (162 of 163) to the corresponding region from HCoV-OC43, but of only 51.2% to the corresponding region of SARS-CoV (Figure 2) (16). Three of these specimens were independently tested by RT-PCR targeted to the matrix and nucleocapsid genes at the British Columbia Cancer Agency's Genome Sciences Centre, and the amplicons were sequenced. Analysis of the sequences (GenBank accession number AY382775-7) showed that they were 98.5% homologous to the corresponding sequences of the HCoV-OC43 genes (16,17). HCoV-OC43 sequences in specimens from four patients were also identified by the NML.

Findings from serological testing

Acute and convalescent sera were collected from 35 affected residents (Table 1). For all nine patients for whom antibody to SARS-CoV were detected on preliminary testing of acute specimens by any of the above methods, no significant neutralizing

TABLE 2
Spectrum of symptoms in affected residents and staff

Symptom or finding	Residents (n=95)		Staff (n=53)	
	n	%	n	%
Fever	20	21.1	11	20.8
Dyspnea	8	8.4	4	7.5
Cough	63	66.3	18	34.0
Sore throat	23	24.2	34	64.2
Runny nose	63	66.3	36	67.9
Myalgia	9	9.5	20	37.7
Headache	2	2.1	22	41.5
Vomiting	2	2.1	2	3.8
Diarrhea	6	6.3	8	15.1
Rash	0	0.0	1	1.9
Fatigue	9	9.5	21	39.6
Nausea	2	2.1	6	11.3
Chills	1	1.1	2	3.8
Pneumonia	12	12.6	0	0.0
Hospitalized (x-ray confirmed)	8	8.4	0	0.0
Clinically diagnosed	4	4.2	0	0.0
Hospitalized without pneumonia*	4	4.2	0	0.0
Deaths†	8	8.4	0	0.0

*Reasons for hospitalization among those admitted without pneumonia included gastric bleed (n=1), pulmonary embolus (n=1) and exacerbation of chronic obstructive pulmonary disease (n=1); †Two residents with congestive heart failure/pneumonia, two with pneumonia, one with chronic obstructive pulmonary disease/pneumonia, one with pulmonary embolism, one with ruptured aortic aneurism, and one admitted for palliative care

antibody titres were detected in the acute or convalescent sera by microneutralization test at the BCCDC or by repeat plaque reduction neutralization assay at the NML. In six patients, neutralizing activity was detected in the serum at a dilution of 1:8, but this was not repeatable. HCoV-OC43-like virus was detected in specimens from three of these six cases. (In this test, neutralizing activity at a dilution of 1:8 has been detected sporadically in healthy controls. In contrast, activity in convalescent sera from SARS patients ranges between dilutions of 1:32 and 1:1024). There was no evidence of seroconversion when acute and convalescent sera were tested in parallel by the virus neutralization test.

Correlation of clinical and laboratory findings

Clinical and laboratory findings for all 18 patients whose specimens on initial or subsequent testing by any method had evidence of SARS-CoV are displayed in Table 3. Residents in whom SARS-CoV was potentially indicated by initial nucleic acid or serological tests expressed a range of symptoms, from mild disease through to lobar pneumonia, which were not consistent with the known profile of SARS.

Findings from peptide assay for cross-reactivity

Eight patient sera from the outbreak were analyzed (Table 4). Five of these had tested weakly reactive by the recombinant nucleocapsid protein ELISA for SARS-CoV at the BCCDC, but had no detectable neutralizing antibody activity (patients 12, 13, 18, 36 and 44). Two of these and one other (patients 13, 44 and 50) were found to have antibody to SARS-CoV by the NML, and two sera that were nonreactive for SARS-CoV antibody on retesting in any laboratory were further analyzed for reactivity to the peptides of the N proteins of SARS-CoV and CoV-OC43.

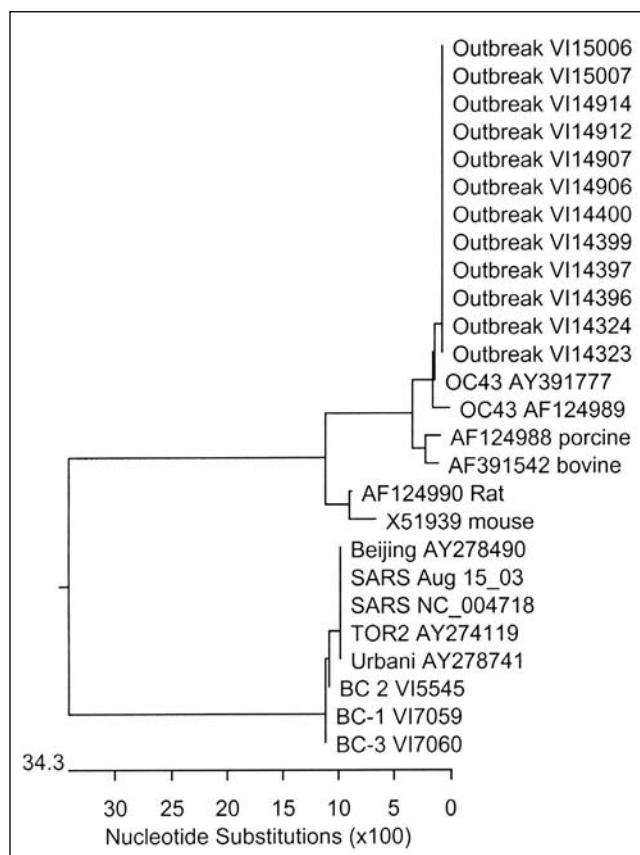


Figure 2) Phylogenetic tree from sequencing of the pan-coronavirus polymerase 1b gene region. Image reproduced with permission from the University of British Columbia Centre for Disease Control Molecular Services

Four of the sites recognized by the sera shared homology between SARS-CoV and HCoV-OC43, and these have been designated S2 to S5 and 43-2 to 43-5, respectively. Where two or three peptides in a row were recognized, the overall sequence was documented (ie, 12 amino acids were listed in the case of two in a row; 14 for three in a row). Serum from patient 48 that was not reactive by SARS-CoV nucleocapsid protein ELISA was only reactive for peptide site 43-5. Conversely, the sera from patients that were weakly reactive by the BCCDC or NML ELISA (patients 13, 44 and 50) were reactive to all peptides expressed by both SARS-CoV and HCoV-OC43. Sera from the other patients were reactive for one to three of the four homologous peptides. These findings show that there are epitopes on the nucleocapsid protein at which antibodies to SARS-CoV and HCoV-OC43 may cross-react.

Public health measures

Once the outbreak was reported, routine influenza-control measures were instituted. After findings suggesting possible involvement of SARS-CoV were reported, these measures were enhanced, and SARS-type respiratory and contact precautions were exercised around symptomatic cases (18). This was to assure that measures shown to limit the spread of SARS-CoV (19,20) were in place until or unless it could be demonstrated that the agent responsible for the facility outbreak was not SARS-CoV.

TABLE 3
Correlation of laboratory and clinical findings for 18 patients with an initial assay positive for severe acute respiratory syndrome coronavirus (SARS-CoV)

Patient number	NML SARS-CoV PCR	BCCDC HCoV-OC43 PCR	NML SARS-CoV serology acute/conv	BCCDC SARS-CoV serology acute/conv	hMPV by PCR	Fever	URI symptoms	Pneumonia	Disposition
12	+ Nasal	- Nasal	ND/-	ND/Ind		+	+	+	Hospitalized
13	+ Stool	- NP	+/-	Ind/Ind			+		Recovered
14	+ Nasal	- Nasal	ND/-	-/-			+		Recovered
17	+ NP	+ NP	-/ND	-/ND			+	+	Died
18	ND	ND	-/-	Ind/Ind			+		Recovered
19	ND	- Stool	WB+/-	-/-			+		Recovered
21	+ Nasal	+ Nasal	-/-	-/-	+		+		Recovered
26	+ Autopsy	+ Autopsy	ND	ND	+	+	+		Died
27	+ Nasal	+ Nasal	-/-	-/-			+		Recovered
29	ND	ND	+/-	-/-			+	+	Recovered
30	ND	ND	WB+/-	-/-			+		Recovered
32	+ Nasal	+ NP	+/-	-/-			+		Recovered
35	- Nasal	+ Nasal	+/-	-/-	+		+		Recovered
36	- Stool	- Stool	-/-	Ind/Ind		+	+		Recovered
44	ND	ND	+/-	Ind/Ind			+		Recovered
46	+ Nasal	+ Nasal	-/-	-/-			+		Recovered
47	ND	ND	WB+/-	-/-			+		Recovered
50	ND	ND	+/-	-/-			+		Recovered

NML serology was denoted as '+' if positive by ELISA, neutralizing antibody assay or both. BCCDC British Columbia Centre for Disease Control; Conv Convalescent; hMPV Human metapneumovirus; Ind Indeterminate (screening ELISA weakly positive but confirmatory neutralization assay negative); ND Not done; NML National Microbiology Laboratory; NP Nasopharyngeal specimen; PCR Polymerase chain reaction; URI Upper respiratory infection; WB+ Serology positive only by Western blot (ELISA and indirect fluorescent antibody negative on same specimen)

DISCUSSION

We have characterized an outbreak of respiratory illness due to HCoV-OC43. The observed attack rate of 67% and case fatality rate of 8% underscore the pathogenic potential of HCoVs in frail populations. This adds to other observations underscoring that CoVs other than SARS-CoV may be responsible for a broader spectrum of disease than coryza alone (21-23).

The fact that no staff members were severely affected was clearly incongruent with tests indicating the possible presence of SARS-CoV. British Columbia had experienced three importations and one occupational transmission of SARS during the original global outbreak (24). None of the four SARS patients were epidemiologically connected to the facility where this outbreak took place.

While initial RT-PCR findings at the NML were suspected to be false-positive because they did not fit the epidemiological picture, early signals on serological testing meant that SARS precautions were prudent as the investigation progressed. The outbreak was observed for 60 days, during which no severe illness among staff and no change toward a pattern reminiscent of SARS occurred. Subsequent molecular and serological investigations conducted at three laboratories ruled against a role for SARS-CoV and in favour of HCoV-OC43 as the etiological agent for this outbreak. Consultation among the BCCDC, Health Canada and the WHO concluded that the outbreak was not SARS, and was unlikely to have involved SARS-CoV.

In 2003, the SARS diagnosis was in its infancy. Several reports have emphasized the increased risk of false-positive SARS tests in the postoutbreak period because of very low pretest likelihood and discussed the potential for cross-reactivity (25,26). The WHO called for further standardization of SARS testing. Since then, several reports (27-31) have made it clear

that cross-reactivity on serological testing may occur, especially when the target is the nucleocapsid protein. Such cross-reactivity has been demonstrated to nucleocapsid proteins of other CoVs but also to human interleukin-11. These studies confirm that antibodies to spike protein are more specific and generally recommend the use of more than one antigen in confirmatory testing or the use of a virus neutralization test.

In the present study, most sera evaluated for cross-reactivity recognized homologous peptide sequences on HCoV-OC43 and SARS-CoV nucleocapsid protein, establishing that these cases were indeed producing cross-reacting antibodies. The immune response to the peptides was most likely a broad-spectrum response that the patients had evolved as a result of repeat infections by different CoVs throughout their lives. Hence, serological tests based on reactions to the SARS-CoV nucleocapsid protein may inadvertently be interpreted as reactive due to cross-reactivity with antibodies from a current or preceding CoV infection, and it is not surprising that Western blot and ELISA assays based on that moiety should have given false-positive results. Because the epitopes to which neutralizing antibodies are directed are believed to be on the spike protein, such cross-reactions may be less likely during the neutralizing antibody test.

The issue of serological cross-reactivity is also compelling when interpreting reports on the seroprevalence of SARS-CoV in animal and human hosts (32,33) and reports of apparent sporadic SARS-CoV infection (34).

With respect to the nucleic acid-based tests used by the NML in these investigations, false-positive results driven by intrinsic test performance or amplicon contamination remain most likely. The reference laboratory had processed hundreds of specimens during the preceding SARS outbreak. Depletion of original specimens made it impossible to completely confirm

TABLE 4
Reactivity of sera with severe acute respiratory syndrome coronavirus (SARS-CoV) and human CoV-OC43 peptides

Case number	S2	43-2	S3	43-3	S4	43-4	S5	43-5	SARS-CoV ELISA-OD BCCDC	SARS-CoV ELISA NML
3	+	+	-	-	+	-	+	+	0.060	neg
12	+	+	-	+	+	+	-	+	0.825	neg
13	+	+	+	+	+	+	+	+	0.255	1:800
18	+	+	+	-	+	+	+	+	0.243	neg
36	+	-	+	-	+	+	-	+	0.435	neg
44	+	+	+	+	+	+	+	+	0.320	neg
48	-	-	-	-	-	-	-	+	0.029	neg
50	+	+	+	+	+	+	+	+	0.065	1:3200

For SARS nucleocapsid protein, the sequences recognized are:

S2=GLPNNTASWFTA
 S3=KMKELSPRWYFYLLGT
 S4=SQASSRSRSSRSRGNRSR
 S5=SAFFGMSRIGME

S3 homology with 43-3

S4 homology with 43-4

S5 homology with 43-5

S2 homology with 43-2

*+ indicates patient serum reactive to the indicated peptide spot.

For OC43 nucleocapsid protein, the sequences recognized are:

43-2=GNVVPYYSWFSG
 43-3=NQRQLLPRWYFYLLGT
 43-4=RSAPNSRSTSRSSRA
 43-5=APTAGAFFFFGRLELA

Abbreviations:

ELISA NML: ELISA report from the National Microbiology Laboratory
 ELISA-OD: Optical density reading of ELISA at the British Columbia Centre for Disease Control (usual cut-off is 0.20)

this assumption. The NML has maintained rigorous quality control since then, with no evidence of similar findings. Since 2003, there has been some effort to refine RT-PCR testing for SARS-CoV generally, and most experts now caution against the use of nested methods for routine diagnostic testing (35).

The present investigation underscores the fact that laboratory testing is but one way to form inferences on the etiology of outbreaks, and cannot replace scrupulous clinical and epidemiological observation. When these different approaches lead to inconsistent observations, it is important to remain open to all possible explanations.

ACKNOWLEDGEMENTS: The authors wish to acknowledge the following individuals: Dr Murray Fyfe and Ms Cheryl McIntyre for assistance with epidemiology, interviews and organization of materials; Ms Lisan Kwindt and Ms Natalie Gaudin for assistance with manuscript preparation; Hayley Rees-Howlett for assistance during record review at the long-term care facility; Lisa Chu, who organized the public health nursing response for contact investigation and follow-up; L Liu, E Grudeski, M Traykova-Andonova, J Krishnan, C Ouellette and D Dick (National Microbiology Laboratory) and Kathy Adie, Loretta Janz, A Mak and R Chow (British Columbia Centre for Disease Control) for many hours contributed to resolving diagnostic test findings; and William Campbell for assistance with the peptide spot assay. The authors also express their appreciation for the cooperation of residents and families of the affected facility. Investigations were performed as a component of an outbreak investigation and did not involve human experimentation. The authors have no relevant conflict of interest to declare. Financial support was provided by the home institutions as listed on the title page.

REFERENCES

- World Health Organization. Update 96 – Taiwan, China: SARS transmission interrupted in last outbreak area <www.who.int/csr/don/2003_07_05/en> (Version current at November 6, 2006).
- Peiris JS, Lai ST, Poon LL, et al. Coronavirus as a possible cause of severe acute respiratory syndrome. *Lancet* 2003;361:1319-25.
- Ksiazek TG, Erdman D, Goldsmith CS, et al. A novel coronavirus associated with severe acute respiratory syndrome. *N Engl J Med* 2003;348:1953-66.
- Drosten C, Gunther S, Preiser W, et al. Identification of a novel coronavirus in patients with severe acute respiratory syndrome. *N Engl J Med* 2003;348:1967-76.
- Fouchier RA, Kuiken T, Schutten M, et al. Aetiology: Koch's postulates fulfilled for SARS virus. *Nature* 2003;423:240.
- Drazen JM. SARS – looking back over the first 100 days. *N Engl J Med* 2003;349:319-20.
- Poutanen SM, Low DE, Henry B, et al. Identification of severe acute respiratory syndrome in Canada. *N Engl J Med* 2003;348:1995-2005.
- Varia M, Wilson S, Sarwal S, et al. Investigation of a nosocomial outbreak of severe acute respiratory syndrome (SARS) in Toronto, Canada. *CMAJ* 2003;169:285-92.
- Enserink M. Infectious diseases. Unexplained false alarm may hold lessons. *Science* 2003;302:767.
- Patrick R, Murray P, Baron EJ, Jorgensen JH, Pfaller MA, Tenover FC, Tenover MC. *Manual of Clinical Microbiology*. Washington: ASM Press, 2003.
- Adachi D, Johnson G, Draker R, et al. Comprehensive detection and identification of human coronaviruses, including the SARS-associated coronavirus, with a single RT-PCR assay. *J Virol Methods* 2004;122:29-36.
- Bastien N, Normand S, Taylor T, et al. Sequence analysis of the N, P, M and F genes of Canadian human metapneumovirus strains. *Virus Res* 2003;93:51-62.
- Guo JP, Petric M, Campbell W, McGeer PL. SARS corona virus peptides recognized by antibodies in the sera of convalescent cases. *Virology* 2004;324:251-6.
- Gene sequence of the SARS Coronavirus (Frankfurt Strain). GenBank accession number AY291315. <www.ncbi.nlm.nih.gov/entrez/viewer.fcgi?db=nucleotide&val=31581502> (Version current at November 6, 2006).
- Marra MA, Jones SJ, Astell CR, et al. The Genome sequence of the SARS-associated coronavirus. *Science* 2003;300:1399-404.
- Gene sequence of human coronavirus OC43. GenBank accession number AY391777. <<http://www.ncbi.nlm.nih.gov/entrez/viewer.fcgi?db=nucleotide&val=37702815>> (Version current at November 6, 2006).
- Kamahora T, Soe LH, Lai MM. Sequence analysis of nucleocapsid gene and leader RNA of human coronavirus OC43. *Virus Res* 1989;12:1-9.
- Health Canada. Public health management of SARS cases and contacts: Interim guidelines. <http://www.phac-aspc.gc.ca/sars-sras/pdf/phmanagementofcases12-17_e.pdf> (Version current at November 12, 2006).
- Fisher DA, Chew MH, Lim YT, Tambyah PA. Preventing local transmission of SARS: Lessons from Singapore. *Med J Aust* 2003;178:555-8.
- Riley S, Fraser C, Donnelly CA, et al. Transmission dynamics of the etiological agent of SARS in Hong Kong: Impact of public health interventions. *Science* 2003;300:1961-6.
- Birch CJ, Clothier HJ, Seccull A, et al. Human coronavirus OC43 causes influenza-like illness in residents and staff of aged-care facilities in Melbourne, Australia. *Epidemiol Infect* 2005;133:273-7.
- El-Sahly HM, Atmar RL, Glezen WP, Greenberg SB. Spectrum of clinical illness in hospitalized patients with "common cold" virus infections. *Clin Infect Dis* 2000;31:96-100.
- Vabret A, Mourez T, Gouarin S, Petitjean J, Freymuth F. An outbreak of coronavirus OC43 respiratory infection in Normandy, France. *Clin Infect Dis* 2003;36:985-9.
- Skowronski DM, Petric M, Daly P, et al. Coordinated response to SARS, Vancouver, Canada. *Emerg Infect Dis* 2006;12:155-8.
- World Health Organization. Summary of the discussion and recommendations of the SARS Laboratory Workshop, 22 October 2003. <www.who.int/csr/sars/guidelines/en/SARSLabmeeting.pdf> (Version current at November 6, 2006).

26. Peiris JS, Yuen KY, Osterhaus AD, Stohr K. The severe acute respiratory syndrome. *N Engl J Med* 2003;349:2431-41.
 27. Chan KH, Cheng VC, Woo PC, et al. Serological responses in patients with severe acute respiratory syndrome coronavirus infection and cross-reactivity with human coronaviruses 229E, OC43, and NL63. *Clin Diagn Lab Immunol* 2005;12:1317-21.
 28. Che XY, Qiu LW, Liao ZY, et al. Antigenic cross-reactivity between severe acute respiratory syndrome-associated coronavirus and human coronaviruses 229E and OC43. *J Infect Dis* 2005;191:2033-7.
 29. Cheng M, Chan CW, Cheung RC, et al. Cross-reactivity of antibody against SARS-coronavirus nucleocapsid protein with IL-11. *Biochem Biophys Res Commun* 2005;338:1654-60.
 30. Maache M, Komurian-Pradel F, Rajoharison A, et al. False-positive results in a recombinant severe acute respiratory syndrome-associated coronavirus (SARS-CoV) nucleocapsid-based Western blot assay were rectified by the use of two subunits (S1 and S2) of spike for detection of antibody to SARS-CoV. *Clin Vaccine Immunol* 2006;13:409-14.
 31. Wang Y, Chang Z, Ouyang J, et al. Profiles of IgG antibodies to nucleocapsid and spike proteins of the SARS-associated coronavirus in SARS patients. *DNA Cell Biol* 2005;24:521-7.
 32. Centers for Disease Control and Prevention (CDC). Prevalence of IgG antibody to SARS-associated coronavirus in animal traders – Guangdong Province, China, 2003. *MMWR Morb Mortal Wkly Rep* 2003;52:986-7.
 33. Zhong NS, Zheng BJ, Li YM, et al. Epidemiology and cause of severe acute respiratory syndrome (SARS) in Guangdong, People's Republic of China, in February, 2003. *Lancet* 2003;362:1353-8.
 34. World Health Organization. New case of laboratory-confirmed SARS in Guangdong, China – update 5. <www.who.int/csr/don/2004_01_31/en> (Version current at November 6, 2006).
 35. Chui L, Drebot M, Andonov A, Petrich A, Glushek M, Mahony J. Comparison of 9 different PCR primers for the rapid detection of severe acute respiratory syndrome coronavirus using 2 RNA extraction methods. *Diagn Microbiol Infect Dis* 2005;53:47-55.
-



Hindawi
Submit your manuscripts at
<http://www.hindawi.com>

