Laboratory diagnosis of SARS

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The emergence of new viral infections of man requires the development of robust diagnostic tests that can be applied in the differential diagnosis of acute illness, or to determine past exposure, so as to establish the true burden of disease. Since the recognition in April 2003 of the severe acute respiratory syndrome coronavirus (SARS-CoV) as the causative agent of severe acute respiratory syndrome (SARS), enormous efforts have been applied to develop molecular and serological tests for SARS which can assist rapid detection of cases, accurate diagnosis of illness and the application of control measures. International progress in the laboratory diagnosis of SARS-CoV infection during acute illness has led to internationally agreed World Health Organization criteria for the confirmation of SARS. Developments in the dissection of the human immune response to SARS indicate that serological tests on convalescent sera are essential to confirm SARS infection, given the sub-optimal predictive value of molecular detection tests performed during acute SARS illness.

Keywords: laboratory diagnosis; RT–PCR; coronavirus; immune response

1. INTRODUCTION

Recognition of the newly described SARS-CoV followed from its detection in clinical material from humans affected with SARS in 2003 (Drosten et al. 2003; Peiris et al. 2003a; Rota et al. 2003). The identification of the virus and its relationship to human disease were confirmed using Koch’s postulates modified for viral diseases and was achieved through an international network of laboratories working under the coordination of the WHO (Kuiken et al. 2003). Experimental work has indicated that the SARS-CoV can be recovered from several organs in infected animals, indicating a disseminated infection, which parallels the observational experience of many of the laboratories involved in handling clinical samples from SARS cases (Peiris et al. 2003b). The main site of replication, pathology and recovery of the virus during human infection is considered to be the lower respiratory tract (Nicholls et al. 2003). This is consistent with the most important route of human-to-human transmission being through respiratory secretions, although outbreaks of infection that involved dissemination of virus excreted in faeces have also been described (Peiris et al. 2003b). Accurate laboratory diagnosis of SARS-CoV was essential to ensure appropriate individual patient management, local infection control and public health measures which were critical in halting the global spread of the first serious new threat to the human population in the twenty-first century.

One of the difficulties of accurate clinical diagnosis of SARS is the relatively long incubation period after infection (mean 6–7 days but ranging up to 10–14 days), before the onset of clinical symptoms and the relatively non-specific nature of the initial illness presentation. Early symptoms include fever, chills, non-specific malaise and myalgia compared with more florid respiratory symptoms which develop later during illness associated with pulmonary infiltrates in the lungs (Donnelly et al. 2003). Summary analyses of published case series indicate that between 25% and 75% of cases demonstrate gastrointestinal symptoms of diarrhoea, nausea and vomiting as illness progresses (Jernigan et al. 2004). The wide range of recognized gastrointestinal disturbance in different case series may be a reflection of the fact that the earliest compilations of data of the disease did not fully recognize this clinical feature. Overall, the major clinical symptoms of respiratory and enteric disease caused by the SARS-CoV in humans are analogous to disease syndromes caused by several animal coronaviruses in their natural hosts.

2. CLINICAL VIROLOGY

Clinical studies on SARS have shed light on the diagnostic usefulness of different samples at different times during illness, summarized in figure 1. In many viral illnesses, virus shedding is greatest during the early symptomatic phase of illness around the onset of symptoms, e.g influenza (Hayden et al. 1998). However, with SARS-CoV, virus excretion is comparatively low during the initial phase of illness and it is necessary to use very sensitive tests that are able to detect low levels of viral nucleic acid during the first days of illness. The mainstay of diagnosis during the illness phase of SARS has involved the use of RT–PCR to detect the SARS-CoV nucleic acid.
amplified directly from clinical samples. RT–PCR protocols were developed with unprecedented speed as a result of the efforts of the WHO collaborative laboratory network. Samples of different body fluids such as blood, respiratory secretions, urine, stool and lung tissue from suspected and probable cases of SARS were analysed. The end point of detection for the SARS-CoV was similar to that found in previously described protocols for detection of known human coronaviruses (Vabret et al. 2001).

It is clear that viral load increases in respiratory samples in the second week of SARS illness (Peiris et al. 2003b), and that the viral load is greatest in samples taken from lower in the respiratory tract (Drosten et al. 2003), peaking around day 10, with the peak of viral detection in faeces coming slightly later. SARS-CoV RNA was detected in only 32% of individuals in nasopharyngeal aspirates at initial presentation (mean 3.2 days after illness onset), but in 68% at day 14 (Peiris et al. 2003b), and in over 90% of faecal samples collected in the second week of illness, peaking around days 15–17 (Chan et al. 2004). Quantification indicated that viral load in respiratory secretions peaked at day 10 with a geometric mean titre of $1.9 \times 10^7$ copies ml$^{-1}$. The clinical features of SARS illness therefore appear to be a good reflection of the body compartments/fluids in which SARS virus has been detected or recovered, with a clear time course. The detection of virus replication in different body compartments over several weeks, before resolution or progression to death, underlies the suggested use of different clinical samples to detect virus at different times after illness onset (figure 1). More recently, analysis of sequential samples of plasma from patients with SARS during early illness using PCR indicates that there is an early peak of viraemia, with up to 70% of samples containing detectable virus in the first few days after onset of illness (figure 2; Grant et al. 2003). This suggests that a viraemic phase is then most probably followed by increasing virus replication in the lower airways and gastrointestinal tract. Taken together, these observations indicate that sampling to detect SARS-CoV in the first week after onset of illness should involve the simultaneous collection and analysis of different clinical samples, including respiratory samples from as low in the respiratory tract as is practicable, blood, faeces and urine. Detection of virus in the second week after the onset of illness is actually more likely, given the higher viral load, and should also involve sampling from multiple sites.

### 3. MOLECULAR DETECTION

Despite reasonably high rates of detection of virus in clinical samples, and good analytical sensitivity of the tests themselves, the predictive value of molecular diagnostic tests in the early stages of illness are still sub-optimal as they cannot rule out the presence of SARS-CoV. This is partly a reflection of the variable viral load in clinical samples, particularly in respiratory samples, which are most likely to be taken from the upper respiratory tract where the priority is to minimize aerosol generation when sampling to prevent infection of healthcare workers. This and the fact that viral replication does not appear to peak until some time after the onset of disease may result in sub-optimal samples. Obtaining a clear diagnosis may be difficult when the disease symptoms are least specific.

Parallel testing of samples for other infectious agents such as influenza, *Mycoplasma pneumoniae*, *Legionella pneumophila* and human metapneumovirus, which are capable of causing a similar clinical syndrome, is essential in the differential diagnosis early after disease onset and
may help to exclude SARS, particularly in returning travellers from countries where SARS is considered likely to re-emerge from an animal reservoir, although co-infection of SARS with other respiratory pathogens can occur (Poutanen et al. 2003). If the presence of an alternative diagnosis is to be used as the justification for discontinuing SARS-specific isolation procedures, the diagnosis should be based on tests with a high predictive value and the clinical illness should be fully explicable by the alternative diagnosis. Testing of multiple sequential samples increases the reliability of laboratory diagnosis, and reduces the likelihood of false-positive results, which is always of concern when using sensitive molecular diagnostic techniques. These findings underlie the current stringent WHO recommendations about the confirmation and quality control of SARS laboratory diagnosis: ‘laboratories performing SARS specific PCR tests should adopt strict criteria for confirmation of positive results, especially in low prevalence areas where the predictive values might be lower’ (Galen & Gambino 1975). This guidance includes the current requirement for detection of virus by RT–PCR in two different samples (e.g. respiratory and faecal), or sequential samples from the same body site on different days and robust confirmatory strategies. Examples of laboratory-acquired infection that occurred in Southeast Asia in 2003–2004, leading to extensive deployment of healthcare resources for contact tracing and quarantine, emphasize the necessity of stringent biosafety considerations in laboratories diagnosing SARS.

4. VIRUS TARGETS FOR DIAGNOSIS

Initial diagnostic work focused on the molecular detection of SARS-CoV RdRp(Pol) gene, because the Pol gene sequences were the first available (Drosten et al. 2003), and the Pol region of the coronavirus genome is well conserved across all coronaviruses. The use of detection probes involving degenerate primer sets that can detect all known coronaviruses (Stephensen et al. 1999) remains a useful screening approach, because this allows the deployment of a pan-corona molecular strategy, which will detect all known human coronaviruses, some of which may possibly cause diseases that overlap with the clinical syndrome of SARS. This approach can be run in parallel with RT–PCRs which are absolutely specific for SARS-CoV (Yam et al. 2003). The sequence conservation in the Pol region across all coronaviruses is such that diagnostic SARS-CoV tests based on the Pol region of the genome should, as part of a validation process, exclude detection of 229E (group 1 coronaviruses) and OC43 (group 2 coronaviruses) to of a validation process, exclude detection of 229E (group 1 coronaviruses) and OC43 (group 2 coronaviruses) to

5. DIAGNOSTIC DEVELOPMENTS

As might be expected in the first several months after the emergence of a new human pathogen, there has been an explosion of diagnostic developments, particularly in the commercial sector. One of the difficulties in validating new diagnostic tests is the availability of clinical material because over 90% of the cases worldwide occurred in Southeast Asia. It is likely that incremental gains in sensitivity of SARS-CoV PCR tests will occur over the next few years as there is increasing use of real-time PCR platforms capable of detecting multiple targets and concurrent or multiplexing of SARS-specific and pan-corona tests. Greater gains in sensitivity may also come from techniques that concentrate the biological sample before processing for nucleic acid extraction (Grant et al. 2003; Chan et al. 2004).

6. VIRUS PROPAGATION

It is fortunate that the SARS-CoV virus, in contrast to many animal coronaviruses, can be cultured easily in a variety of continuous cell lines, including FRhK and Vero E6 cells, produces a recognizable and distinct widespread CPE (figure 3), and grows well at 33 °C and 37 °C. This has allowed the recovery of infectious virus from affected individuals, which in turn expedited sequencing of the entire virus genome (Marra et al. 2003). Moreover, the development of infectivity assays has allowed quantification of virus infectivity and development of neutralization assays (plaque reduction neutralization tests are shown in figure 4). Infectious virus has not been recovered beyond three weeks after illness onset even though virus species at least during infection in cell culture. Products of non-structural genes, such as the RdRp(Pol), responsible for replication and transcription of the viral genome, are needed in smaller amounts than structural genes, such as the NC protein involved in assembly of the virions. Consequently, sgRNA for the NC gene should be more abundant than sgRNA for the Pol gene in infected cells. This feature of coronavirus biology may be relevant for improving its detection in clinical practice. There are considerable differences in the concentration of viral RNA fragments in infected cells, with several log-fold increases in the amount of NC (mRNA) RNA in infected cells, compared with the transcripts of Pol genes (Thiel et al. 2003). This finding suggests that there may be some diagnostic advantages to targeting NC genes for molecular detection as well as other genes, to improve the overall sensitivity of detection, because the amount of viral template will be much higher, if clinical material contains virus infected cells as well as whole virus. Using clinical samples spiked with a mixture of SARS-CoV virus and infected cells, it is evident that detection of NC genes does provide some additional sensitivity (table 1). Several laboratories have developed diagnostic PCRs for the detection of other regions of the genome, particularly the viral NC gene (Emery et al. 2004). This approach is consistent with the observation that targeting the SARS-CoV NC region improved the sensitivity of detection more than a 100-fold in experimentally infected animals (Kuiken et al. 2003).
Table 1. Detection of SARS-CoV in spiked simulated clinical specimens using different molecular detection strategies.

<table>
<thead>
<tr>
<th>sample type</th>
<th>number of samples</th>
<th>2Bp/4Bma (block)</th>
<th>N5/N6 primers$^d$ (block)</th>
<th>N5/N6 primers$^c$ (light cycler)</th>
<th>N primers$^f$ (light cycler)</th>
<th>SARS-specific Pol primers$^g$ (light cycler)</th>
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<tr>
<td>SARS-CoV spiked samples</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>faecal$^a$</td>
<td>38</td>
<td>15 (39.5%)</td>
<td>28 (73.7%)</td>
<td>32 (84.2%)</td>
<td>32 (84.2%)</td>
<td>30 (78.9%)</td>
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<tr>
<td>plasma</td>
<td>15</td>
<td>13 (86.7%)</td>
<td>14 (93.3%)</td>
<td>15 (100%)</td>
<td>15 (100%)</td>
<td>15 (100%)</td>
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<tr>
<td>respiratory$^b$</td>
<td>35</td>
<td>34 (97.1%)</td>
<td>27 (77.1%)</td>
<td>28 (80.0%)</td>
<td>30 (85.7%)</td>
<td>28 (80%)</td>
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<tr>
<td>urine</td>
<td>9</td>
<td>7 (77.8%)</td>
<td>8 (88.9%)</td>
<td>9 (100%)</td>
<td>9 (100%)</td>
<td>9 (100%)</td>
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<tr>
<td>total spiked</td>
<td>97</td>
<td>69 (71.1%)</td>
<td>77 (79.4%)</td>
<td>84 (86.6%)</td>
<td>86 (88.7%)</td>
<td>82 (84.5%)</td>
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<tr>
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<tr>
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<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
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<tr>
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<td>5</td>
<td>0</td>
<td>0</td>
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<td>2</td>
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<tr>
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<td>0</td>
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<td>0</td>
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<td>4</td>
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<td>overall sensitivity (%)</td>
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<td>79.4%</td>
<td>86.6%</td>
<td>88.7%</td>
<td>84.5%</td>
<td></td>
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<td>overall specificity (%)</td>
<td>57.5%</td>
<td>100%</td>
<td>100%</td>
<td>92%</td>
<td>85.2%</td>
<td></td>
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<tr>
<td>positive predictive value</td>
<td>80.2%</td>
<td>100%</td>
<td>100%</td>
<td>97.7%</td>
<td>95.3%</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Twelve out of 38 faecal samples were co-inoculated with rotavirus.

$^b$ Fourteen out of 35 respiratory samples were co-inoculated with influenza A virus.

$^c$ Stephensen et al. (1999).

$^d$ SARS-specific NC primers: N5: 5′ GGTGACGGGCAAAATGAAAAGGC 3′ (for primer); N6: 5′ ATGAGGAGCGAGAAGGCTGGAC 3′ (reverse primer).

$^e$ Probe for light cycler assay: NC LCR probe: 5′ LC red 640-ATTTGGCCACCGCAATCC-phosphate 3′; NC Flu probe: 5′ GAGCCTTGAATACACCCAAAGACC-fluorescein 3′.

$^f$ SARSNP fpr1/SARSNP rpr1/SARSNP prb1 (Kuiken et al. 2003).

$^g$ SARSTM fpr1/SARSTM rpr1/SARSTM prb1 (Kuiken et al. 2003).
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7. SEROLOGICAL ASSAYS

As the virus produced a CPE in FRhK cells, virus-infected cells were used as an antigen substrate before the aetiological agent had been described. Seroconversion using IF was the earliest serological test used to detect SARS (Peiris et al. 2003a). Use of IF indicated that seroconversion took place ca. 10 days after illness onset, but might not actually be evident in all cases until ca. 28 days after onset, as ca. 10–20% of individuals did not have detectable antibodies until after day 21 (Peiris et al. 2003b). The late seroconversions noted may reflect the fact that many patients were treated with high-dose steroids, which is likely to have delayed the antibody response, although this cannot be firmly concluded from the clinical datasets available. The only available laboratory method for excluding the diagnosis of SARS-CoV infection is to obtain a negative result on serological testing of a convalescent phase serum at 28 days after onset of symptoms. It is therefore essential for understanding the true disease burden to have robust and reliable serological tests.

The development of SARS ELISA tests followed rapidly after the identification of the virus, and the use of virally infected cells to prepare antigen for indirect ELISAs for the detection of SARS antibodies (IgM and IgG) has been possible because of the ability to culture virus to reasonable titre, and to use virus-infected cells as a source of antigens (figure 5; Ksiazek et al. 2003; Kuiken et al. 2003).

Analysis of antibody responses to SARS-CoV has so far shown limited cross reactivity with antibodies to human group 1 or group 2 coronaviruses. Although full evaluations are not complete yet and further data are required, some limited cross reactivity to group 1 animal coronaviruses has been noted (Ksiazek et al. 2003). The method of preparation of antigens and the formulation of serological tests may impact substantially on the ability to detect any cross-reacting antibody. Understanding the impact of serological responses to other human coronaviruses on antibody response to SARS-CoV is important because it will affect the specificity of tests and conclusions about exposure to SARS-CoV in the absence of clinical illness. Serological data are developing rapidly and early data suggest that high levels of neutralizing antibodies are formed after SARS infection and last for at least several months after infection. The use of neutralizing antibody tests (such as shown in figure 4) indicates that antibody to SARS may also cross neutralize related animal viruses, perhaps with a slightly lower titre, and this is taken as a

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Figure 3. Mock infected Vero E6 cells (a) compared with CPE of SARS in Vero E6 cells (b) at 4 days after infection.

Figure 4. Neutralizing antibody tests. Infectivity (a) and plaque reduction assays (b,c) for the detection of SARS-neutralizing antibodies. Virus-infected cells are shown in the first two wells of (a), indicating clear plaque formation in Vero E6 cells 4 days after inoculation, with mock infected cell control in the third well. Infection of virus in the presence of increasing dilution of SARS convalescent serum is shown in (b) and indicates total inhibition of virus growth at day 4 after inoculation with an IC₅₀ serum titre of ca. 1 in 320. Non-SARS serum with no neutralizing activity is shown in (c).
suggestion of more than one serogroup of SARS-CoV (Zheng et al. 2004).

The specificity of the SARS-CoV antibody response has allowed seroprevalence studies to be undertaken using IF, which have concluded that there was little spread of SARS-CoV in the general population in Hong Kong, based on blood donor screening (Donnelly et al. 2003). However, up to 40% of humans who are market traders of live animals or who are restaurant workers preparing exotic meat of the putative wild animal reservoir (members of the viverrid, mustelid and canid families), showed evidence of exposure to SARS viruses, which has been taken to support the zoonotic origin of the SARS-CoV (Guan et al. 2003). Screening of archived healthy adult sera in Hong Kong (Zheng et al. 2004) taken before the SARS outbreak indicated that a few had detectable antibody to the SARS-CoV, suggesting pre-existing evidence of exposure to a related virus.

One of the difficulties of screening individual sera, whether from cases of illness, for serosurveillance or for contact tracing, is the sensitivity and specificity of ELISAs or IF assays, which typically have sensitivities and specificities between 90% and 98%. This is true for almost all ELISAs used to screen human sera for many viral diseases, and usually leads to an algorithm of a screening assay followed by a confirmatory assay. Invariably, a small proportion of reactive sera will not be true positives after the first ELISAs. To improve the certainty of diagnosis, a serological strategy needs to be adopted, involving a second tier of tests (figure 6) with or without an additional second serum to test for seroconversion. Many laboratories have adopted a neutralization test as a ‘gold standard’ confirmatory assay, with typical neutralizing antibody titres of between several hundred and several thousand detectable at 28 days after onset of illness. The rise in neutralizing antibody may not exactly parallel the rise in total antibody detection, and may be somewhat slower to develop. However, tests that use virus-infected cells, or live virus, as required for whole-cell lysate ELISA assays, IF or neutralization tests, require the growth of virus (figures 4 and 5), which in turn requires a biosafety level 3 laboratory and prevents the tests being used widely. It is likely that trends in serological assay development will be towards the use of recombinant antigen ELISAs and finding surrogate methods for neutralization tests such as receptor binding assays, which may be a safer alternative for the serological diagnosis of SARS. The limited data available internationally so far suggest good correlation between recombinant protein ELISAs, Western blots and IF results (Wu et al. 2004), but much more evaluation will be required to fully understand the relationships between antibodies to different coronaviruses.

There remain many unanswered questions about the nature of serological responses to infection with the SARS-CoV, despite the astonishing rapidity of development of robust diagnostic tests. The next few years will

Figure 5. Detection of antibodies to SARS-infected cell antigen by dot blot (a)(i,ii) and ELISA (b). (a) Sera from SARS-positive UK probable case (Kuiken et al. 2003) at (i) day 7 acute, and (ii) day 28 convalescent, is shown in dot blot assay formats indicating the detection of low levels of antibody early after illness onset and seroconversion at day 28. (b) ELISA reactivity of serial dilutions of serum taken from: (i) acute SARS day 7; (ii) day 28 convalescent SARS; and (iii) day 28 after influenza A illness presenting with a clinical syndrome fulfilling the WHO case definition for ‘probable SARS’.

Figure 6. Suggested algorithm for serological testing of SARS.
undoubtedly see the unravelling of the relationship between neutralizing and functional antibody and total antibodies to specific virus proteins, the duration and magnitude of antibody response in the context of disease protection and a comparison of antibody response in children and adults. An attempt to understand serological relationships between different SARS-CoV viruses as well as between SARS-CoV and other human and non-human coronaviruses will benefit our understanding of the biology of coronaviruses as a whole, and assist understanding of the severity of SARS disease in humans. A very significant side effect of SARS-related research is likely to be much more focus on the burden of illness as a result of human coronaviruses and their role in acute respiratory and gastrointestinal infections, a neglected backwater of human virology.

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GLOSSARY

CPE: cytopathic effect

ELISA: enzyme linked immunosorbent assay

IF: immunofluorescence

NC: nucleocapsid

RdRp(Pol): RNA-dependent RNA polymerase

RT–PCR: reverse transcription–polymerase chain reaction

SARS: severe acute respiratory syndrome

SARS-CoV: severe acute respiratory syndrome coronavirus

sgRNA: sub-genomic RNA

WHO: World Health Organization