Molecular Epidemiology of Severe Acute Respiratory Syndrome–Associated Coronavirus Infections in Taiwan

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Background. In 2003, Taiwan experienced a series of outbreaks of severe acute respiratory syndrome (SARS) and 1 laboratory-contamination accident. Here we describe a new phylogenetic analytical method to study the sources and dissemination paths of SARS-associated coronavirus (SARS-CoV) infections in Taiwan.

Methods. A phylogenetic analytical tool for combining nucleotide sequences from 6 variable regions of a SARS-CoV genome was developed by use of 20 published SARS-CoV sequences; and this method was validated by use of 80 published SARS-CoV sequences. Subsequently, this new tool was applied to provide a better understanding of the entire complement of Taiwanese SARS-CoV isolates, including 20 previously published and 19 identified in this study. The epidemiological data were integrated with the results from the phylogenetic tree and from the nucleotide-signature pattern.

Results. The topologies of phylogenetic trees generated by the new and the conventional strategies were similar, with the former having better robustness than the latter, especially in comparison with the maximum-likelihood trees: the new strategy revealed that during 2003 there were 5 waves of epidemic SARS-CoV infection, which belonged to 3 phylogenetic clusters in Taiwan.

Conclusions. The new strategy is more efficient than its conventional counterparts. The outbreaks of SARS in Taiwan originated from multiple sources.

Severe acute respiratory syndrome (SARS) is caused by SARS-associated coronavirus (SARS-CoV) [1–4]. The first known outbreak of SARS occurred in China’s Guangdong province during November 2002 [5]. By 7 August of the following year, SARS had spread to >30 countries, affecting 8096 people and resulting in 774 deaths worldwide [6]. In Taiwan, the first SARS case was diagnosed on 14 March 2003 [7, 8]. This index case involved a Taiwanese businessman who had visited Guangdong province during 5–21 February of that year. After he returned to Taiwan, he transmitted the disease to his wife, his son (SARS-CoV strain TW1), and the doctor who treated his son (SARS-CoV strain TW3).

On 15 March, 7 employees of a Taiwanese construction company flew from Hong Kong to Beijing; 4 of them developed SARS symptoms on 26 March, several days after returning to Taiwan [9]. Also on 26 March, a man residing at the Amoy Gardens housing complex in Hong Kong flew to Taiwan; the following day, he took a train from Taipei to Taichung City to visit his younger brother. The visitor returned to his Hong Kong home on 28 March after having experienced fever during the preceding evening. His younger brother (patient TWC), who developed symptoms on 31 March, became Taiwan’s first SARS-related fatality.

On 6 April, a Taiwanese woman (patient TW-HP1) suffering from fever and coughing that continued for several days visited the emergency room (ER) at mu-
nicipal hospital HP; she was transferred to another hospital on 9 April. Seven HP employees, including a laundry worker who was identified as the index case, eventually developed SARS, resulting, on 24 April, in a shutdown of all operations of hospital HP [7]. In all, 137 probable SARS cases and 26 HP-related fatalities resulted from this single nosocomial infection. Patient TW-HP1 had not traveled outside Taiwan during the preceding 12 months, but, on 27 March, she and the visitor from Hong Kong had taken the same train; their seats were in separate cars (numbers 3 and 5), yet the train ride constitutes their only possible point of contact.

On 28 April, the government of Taiwan imposed mandatory quarantines on all air travelers from China, Hong Kong, Singapore, Macau, and Toronto; however, nosocomial SARS infections continued to be reported in many hospitals islandwide [5]. The hospitals that experienced the most-severe outbreaks are listed in figure 1; in the present study, they are referred to by initials—"HP," "JC," "KC," "GD," and "YM." According to Taiwan’s Center for Disease Control (CDC), 346 of the 664 probable SARS cases that have been reported to the World Health Organization (WHO) were confirmed by reverse-transcriptase polymerase chain reaction (RT-PCR) and/or neutralizing-antibody tests [10]. Previously, Yeh et al. had studied the molecular epidemiology of SARS infection in Taiwan and had concluded that the origin of the Taiwanese SARS epidemic was mainly either Hong Kong or Guangdong, rather than Beijing [11]; in addition, they found that the SARS-CoV isolated from the younger brother (i.e., patient TWC) of the visitor from Hong Kong was not clustered with other isolates from hospital HP [11, 12]. Because a complete genome sequence from the nasopharyngeal aspirate from patient TWC (strain TC1) was available, we decided to reexamine both (1) the source of infection at hospital HP and (2) the paths of dissemination of SARS among hospitals in Taiwan.

On the evening of 10 December 2003, a medical researcher who regularly worked in a biosafety level–4 laboratory started to feel feverish [13]. Although he had recently spent 4 days (7–10 December) in Singapore, an epidemiological investigation indicated that he had contracted SARS from a laboratory-contamination accident on 6 December. According to his description, the SARS-CoV isolates that he had handled in the laboratory included strain HKU-39849 [14] and other clinical isolates that have not been well characterized. The nasopharyngeal aspirate from this medical researcher was included in the present study.

The size of the SARS-CoV genome has been measured as 29.7 kb [15, 16]. A comparative analysis of 14 SARS-CoV isolates has identified 2 distinct genotypes, which can be differentiated on the basis of 4 single-nucleotide variations (SNVs)—C:C:G:C versus T:T:T:T—in the variable regions of the SARS-CoV genome [17]. Furthermore, the genotype with the T:T:T:T SNVs has been associated with infections originating in Hotel M in Hong Kong [8, 17]. Currently, phylogenetic analyses of SARS-CoV require complete genome sequences [5, 11, 17]; however, because of the

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**Figure 1.** Epidemiological curve of confirmed cases of severe acute respiratory syndrome (SARS) in Taiwan, which Taiwan’s Center for Disease Control validated by use of either reverse-transcriptase polymerase chain reaction or serological test. Arrows indicate dates of outbreaks of nosocomial infection in different hospitals and of diagnoses of SARS in several key patients.
limited number of specimens available for complete genome sequencing, some researchers have used the SARS-CoV spike gene for this purpose; but most results have been less than satisfactory [18, 19]. Therefore, the objectives of the present study were (1) to use 20 complete SARS-CoV sequences to combine several variable regions of a SARS-CoV for phylogenetic analysis, in order to develop a simpler tool; (2) to use a different set of sequences from 80 SARS-CoV isolates to validate the method, by 3 different phylogenetic analytical methods; and (3) to apply this proposed tool to elucidate the origin and paths of dissemination of SARS-CoV infections, on the basis of samples collected from hospitals in Taiwan.

PATIENTS, MATERIALS, AND METHODS

Patients. Serum, sputum, or throat-swab specimens from 19 patients with SARS associated with 5 nosocomial infections and 1 laboratory-contamination accident were collected for the present study. For each patient, data on the date of onset of the disease and on possible sources of infection were gathered by trained interviewers (table 1). We also downloaded, for our phylogenetic tree analyses, 20 complete Taiwanese SARS-CoV sequences from GenBank; for 10 of these, SARS-CoV strains have been described elsewhere [11].

Epidemiological investigation. To track the origins of SARS cases in Taiwan, we used SARS-treatment reports written by physicians and submitted by their respective hospitals to Taiwan’s CDC. We also sent trained interviewers to all hospitals that reported nosocomial infections, to gather additional assessment data. We based our investigation on the WHO definition of SARS [20].

Analysis of SARS-CoV sequence variation, and selection of variable regions for phylogenetic analysis. Complete nucleotide sequences from 20 SARS-CoV isolates available from GenBank were aligned by the BioEdit program [21]; the sequence variations were analyzed by the SimPlot program (http://sray.med.som.jhmi.edu/RaySoft/SimPlot). Genomic sequences from civet-cat SARS-CoV strains SZ-3 and SZ-16 were used as standards for comparison [19]. Sequence-variation-distance plots were generated by use of an 800-bp window, a 200-bp step, and a Jukes-Cantor correction. Initially, 4 variable regions (SC18, SC22-23, SC27, and SC28), which contain SNVs of different SARS-CoV genotypes [17], were combined for phylogenetic analysis. Because the resultant phylogenetic tree was not satisfactory, we added another 2 variable regions (SC10 and SC20) based on the sequence-variation plot (figure 2).

Comparison of conventional and proposed strategies for phylogenetic analysis. Three methods—neighbor joining (NJ), Fitch and Wagner parsimony (Pars), and maximum likelihood (ML)—were used for comparison of the conventional (i.e., complete genome) and proposed strategies. The proposed strategy for phylogenetic analysis entailed the deletion of conserved domains and the combination of minimal variable regions. The MEGA2 and Phylip3.6 software packages were used to construct the phylogeny [22–24]; and 80 SARS-CoV sequences down-

Table 1. Demographic data and possible sources of infection in cases of severe acute respiratory syndrome (SARS) that were used for molecular epidemiological study.

| Patient (sex; age, years) | Date, in 2003, of onset of SARS | Source of infection |
|--------------------------|----------------------------------|--------------------|
| TW-HP1 (F; 47)           | 11 April                         | Transmission during train ride (took same train as visitor from Hong Kong) |
| TW-HP2 (M; 41)           | 28 April                         | Nosocomial infection (visited emergency room of hospital HP) |
| TW-HP3 (F; 37)           | 29 April                         | Nosocomial infection (was a patient in hospital HP) |
| TW-HP4 (M; 31)           | 29 April                         | Family contact (wife was a nurse at hospital HP) |
| TW-JC2 (F; 38)           | 1 May                            | Nosocomial infection (was radiography technician at hospital JC) |
| TW-KC1 (M; 54)           | 15 May                           | Family contact (relative was a patient with SARS, at hospital KC) |
| TW-KC3 (F; 42)           | 20 May                           | Nosocomial infection (was a patient in hospital KC) |
| TW-PH1 (M; 60)           | 19 May                           | Nosocomial infection (visited hospitals KC and PH) |
| TW-PH2 (F; 51)           | 23 May                           | Nosocomial infection (provided care to a patient in hospital PH) |
| TW-GD1 (F; 26)           | 21 May                           | Nosocomial infection (provided care to a patient in hospital GD) |
| TW-GD2 (M; 75)           | 29 May                           | Nosocomial infection (was a patient in hospital GD) |
| TW-GD3 (M; 73)           | 2 June                           | Nosocomial infection (was a patient in hospital GD) |
| TW-GD4 (M; 75)           | 6 June                           | Nosocomial infection (was a patient in hospital GD) |
| TW-GD5 (M; 78)           | 4 June                           | Nosocomial infection (was a patient in hospital GD) |
| TW-YM1 (F; 47)           | 8 June                           | Nosocomial infection (provided care to a patient in hospital YM) |
| TW-YM2 (F; 67)           | 8 June                           | Nosocomial infection (provided care to a patient in hospital YM) |
| TW-YM3 (F; 90)           | 8 June                           | Nosocomial infection (was a patient taken care of by both TW-YM1 and TW-YM2, in hospital YM) |
| TW-YM4 (M; 86)           | 8 June                           | Nosocomial infection (was a patient in hospital YM) |
| SCVJ (M; 44)             | 10 December                      | Laboratory-contamination accident |
Figure 2. Analysis of nucleotide-sequence variation of 20 isolates of human severe acute respiratory syndrome–associated coronavirus (SARS-CoV), by the SimPlot program. Two civet-cat SARS-CoV strains were used as references. The X-axis indicates the nucleotide location of the SARS-CoV genome; the Y-axis indicates the rate of nucleotide-sequence variation. Sequence differences are plotted with a window of 800 nt and 200-nt steps. The 6 variable regions—SC10, SC18, SC20, SC22-23, SC27, and SC28—used for the combined phylogenetic analyses are in boldface type.
RT-PCR and sequencing. RNA was extracted from serum, sputum, or throat-swab specimens by QIAamp viral-RNA mini-kits (Qiagen). RT-PCR primer pairs for the 6 SARS-CoV variable regions are listed in table 2. RT-PCR was performed in a single-tube reaction (Qiagen) using primers from each region. The PCR thermocycler program consisted of pre-denaturing for 4 min at 95°C; 35 cycles of denaturing for 30 s at 95°C, annealing for 30 s at 52°C, and initial extension at 68°C for 3 min; and final extension at 68°C for 10 min. The PCR product was gel-purified for DNA sequencing by use of an ABI PRISM 3700 DNA Analyzer (Applied Biosystems). For the laboratory-contamination case, multiple primers [17] were used for complete genome sequencing. Superscript III RT (Invitrogen) was used for production of cDNA, and a RACE kit (Roche) was used to amplify the 5′ and 3′ ends of cDNA. Both strands of the PCR product were analyzed, and the resultant sequences were assembled by use of the SeqMan II software package (version 5.0; Lasergene).

Analysis of the nucleotide-signature patterns of different waves of epidemic SARS infection. To perform the analysis of nucleotide-signature patterns, representative SARS-CoV strains with complete sequences were chosen from each wave of epidemic SARS infection in Taiwan. In addition, cases that were of unclear origin were also included in the analysis. The nucleotide sequence of SARS-CoV strain Urbani [16] was used as a prototype for the comparison.

RESULTS

Development of a proposed strategy for phylogenetic analysis of SARS-CoV isolates. Pairwise comparisons were used to analyze sequence variations among 20 SARS-CoV isolates. The results show that the 3′ region of the viral genome had the highest sequence variation, especially near the junction of replicase 1b and the spike genes (figure 2). Six variable regions—SC10, SC18, SC20, SC22-SC23, SC27, and SC28—were chosen for testing the proposed strategy for phylogenetic-tree analysis; the total length of the sequences in these 6 regions was 5287 nt. Using 3 phylogenetic analytical methods, we compared, for 80 SARS-CoV isolates, the topology and robustness of the phylogenetic tree loaded from the GenBank database were used for the comparison. A bootstrap analysis of 100 replicates was used to compare the robustness of the NJ and Pars trees generated by the conventional strategy versus that of the NJ and Pars trees generated by the proposed strategy [25]. For the ML method, the P value for branch length was calculated by use of the hidden Markov model, before the conventional and proposed strategies were compared [26].

Table 2. Primers used for reverse-transcriptase–polymerase chain reaction analysis of 6 variable regions of the severe acute respiratory syndrome–associated coronavirus (SARS-CoV) genome.

| Region (np), primer paira | Fragment used for combined analysisa, np (length) | Coding region |
|-------------------------|-----------------------------------------------|---------------|
| SC-10 (8964–10100)      |                                               |               |
| SC10F: 5′-GGATGCTATGGGCAAACCTGTGCC-3′  |
| SC10R: 5′-GGACAGTATACTGTGTCATCCAAC-3′  |
| SC-18 (17002–18124)     |                                               |               |
| SC18F: 5′-CACTCCAAGGACCACCTGGTACTG-3′  |
| SC18R: 5′-CGGTAGGTCATGTCCTTTGGTATG-3′  |
| SC-20 (18914–20154)     |                                               |               |
| SC20F: 5′-GGTTGTGAAGTCTGCATTGCTTGC-3′  |
| SC20R: 5′-CCTCTAAGTCTCTGCTCTGAGTAA-3′  |
| SC22-23                 |                                               |               |
| SC22F: 5′-GGTGTGTAAGTCTGCATTGCTTGCG-3′  |
| SC22R: 5′-CCTCTAAGTCTCTGCTCTGAGTAA-3′  |
| SC-27 (26007–27112)     |                                               |               |
| SC27F: 5′-CAATCGACGGCTCTTCAGGAGTTG-3′  |
| SC27R: 5′-CTCTGCTATGGTAACCTGGAAGTC-3′  |
| SC-28 (27018–28166)     |                                               |               |
| SC28F: 5′-AGACACGGCCGTCAGAAGGAGTTG-3′  |
| SC28R: 5′-ATGCCGCGGGGCACTACGTGTTT-3′  |

NOTE. np, nucleotide position; ORFs, open reading frames.

* Nucleotide residues of Tor2 SARS-CoV.
Figure 3. Neighbor-joining (NJ) trees of 80 isolates of severe acute respiratory syndrome–associated coronavirus (SARS-CoV) that were downloaded from GenBank, by use of (left) the conventional strategy, which uses the complete genome sequence, and (right) the proposed strategy, which uses sequences from 6 SARS-CoV variable regions. Numbers at nodes are bootstrap values (%), and those at 3 nodes are circled: node a, between civet-cat SARS-CoV and human SARS-CoV; node b, between SARS-CoV subgroups A1 (early phase) and A2 (middle phase); and node c, between SARS-CoV subgroups A and B (late phase) [5]. The scale bar indicates the genetic distance, estimated on the basis of Kimura’s 2-parameter substitution model [24].
generated by the proposed strategy, which uses only these 6 regions, versus the topology and robustness of the tree generated by the conventional strategy, which uses the complete genome sequences. As shown in figure 3, the topology of the NJ tree generated by use of the complete genome sequence was almost identical to that of the NJ tree generated by use of a combination of the 6 variable regions; similar results were obtained for the Pars trees and the ML trees (data not shown).

To compare the robustness of trees generated by the proposed method versus that of trees generated by the conventional method, we focused on the bootstrap values for 3 bifurcation nodes between SARS-CoV clusters: node “a,” between civet-cat SARS-CoV and human SARS-CoV; node “b,” between SARS-CoV subgroups A1 and A2; and node “c,” between SARS-CoV subgroups A and B. In the NJ trees, the bootstrap values for nodes a, b, and c were, respectively, 100%, 100%, and 63% for the conventional method versus 96%, 83%, and 66% for the proposed method (figure 3); similar results were obtained for the Pars trees and the ML trees (data not shown).

Table 3. Bootstrap and P values for trees generated by use of the complete genome and by use of 6 variable regions of sequences of severe acute respiratory syndrome–associated coronavirus, by 3 traditional analytical methods.

| Sequence(s) used for phylogenetic analysis | Neighbor-joining method | Parsimony method | Maximum-likelihood method, P |
|-------------------------------------------|-------------------------|------------------|-----------------------------|
| Complete genome (29.7 kb)                 | Node a 100              | Node a 100       | Node a <.01                 |
|                                           | Node b 100              | Node b 100       | NS                          |
|                                           | Node c 63               | Node c 56        | NS                          |
| 6 Variable regions (5.3 kb)               | 96                      | 93               | <.01                        |
|                                           | 83                      | 78               | <.01                        |
|                                           | 66                      | 67               | <.01                        |

NOTE. NS, not significant.

As shown in figure 5, there were 27 SNVs and 1 dinucleotide

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|                                           | 66                      | 67               | <.01                        |

NOTE. NS, not significant.

As shown in figure 5, there were 27 SNVs and 1 dinucleotide
deletion among 14 SARS-CoV isolates; 12 of the 27 SNVs were nonsynonymous changes. Distinctive SNVs were identified for each wave of epidemic SARS infection in Taiwan—at nucleotide position 3165 for wave 1; at nucleotide positions 3852, 11493, and 26477 for both wave 3 and wave 4; and at nucleotide positions 26203 and 27812 for wave 4. The nucleotide-signature pattern between the strain from patient SCVJ (the laboratory-contamination case) and strain TWC consisted of 2 SNVs (at nucleotide position 16325–26600) and 1 dinucleotide deletion (at nucleotide position 27808–27809).

DISCUSSION

When, for phylogenetic analysis, we combined sequences from different variable regions of the SARS-CoV genome, we assumed that no dual infection or recombination between the SARS-CoV subgroups had occurred. Because SARS is an acute infectious disease, the odds that it will be contracted from 2 subgroups are very low. Furthermore, the strategy of combining different genomic regions for phylogenetic analysis has been used in molecular epidemiological investigations of the origins of HIV-2. In those studies, the evolutionary history of a simian immunodeficiency virus/HIV-2 lineage was reconstructed by use of a combination of partial gag and env sequences; the method increased the accuracy of the phylogenetic analysis [28].

It is noteworthy that the SimPlot analysis demonstrated that the 3' region of the viral genome, especially near the junction of replicase 1b and the spike genes (encoding protein S), had the greatest sequence variation (figure 2). The S protein is...
Figure 5. Signature patterns of single-nucleotide variations in 5 waves of epidemic severe acute respiratory syndrome–associated coronavirus (SARS-CoV) infection in Taiwan. Nucleotides are numbered on the basis of the complete genome sequence of SARS-CoV strain Urbani [16]. Deletions are denoted by X’s. *, Isolate with 24-nt deletion at nucleotide position 26132–26155; **, amino acid residues of SARS-CoV nonstructural protein (Nsp) and open reading frame (Orf) [16].

considered to be the most important target for the humoral and cellular immune responses to SARS-CoV [15, 16].

The 3 algorithms most commonly used in molecular phylogenetic analyses are the NJ, ML, and Pars methods; we used all 3 to obtain tree topologies. In terms of robustness, a bootstrap value of 70% is often cited as a cutoff for a reliable cluster [25]. Although the bootstrap values for nodes a and b in the NJ and Pars trees generated by the conventional method were higher than those in the NJ and Pars trees generated by the proposed method, the difference was not statistically significant. In contrast, the bootstrap values for node c in the Pars trees generated by the conventional and proposed methods were 56% and 67%, respectively, and the corresponding bootstrap values in the NJ trees were 63% and 66%. This finding was confirmed by the ML trees; the P value for node c in the tree generated by the conventional method was not statistically significant, and that for node c in the tree generated by the proposed method was <0.01. Accordingly, the tree generated by the proposed method was more reliable; also, because the proposed method requires only 7 RT-PCR reactions to perform the analysis, it is less time-consuming and more efficient than the conventional method. To facilitate other laboratories’ future molecular epidemiological studies of outbreaks of SARS, we have made the nucleotide-sequence–alignment file of 80 SARS-CoV reference strains available on our center’s Web site (http://www.ym.edu.tw/aids/Molepi/).

In the present study, we used 39 Taiwanese SARS-CoV isolates, including 20 downloaded from the GenBank database, to trace the origin and the path of dissemination of SARS-CoV infections that occurred in Taiwan during 2003. Phylogenetic analyses demonstrated that the Taiwanese SARS-CoV strains were distributed in 3 clusters—B1, B2, and B3—which differ from clusters 1–3 reported by Yeh et al. [11]; the latter clusters were not defined on the basis of a bootstrap value, whereas we used a bootstrap value of 70% as the cutoff to define our clusters B1–B3. Therefore, our cluster B1 contains both cluster 1 and cluster 2 of Yeh et al., and their cluster 3 was divided into 2 clusters—B2 and B3—in our study.

On the basis of the epidemiological data for the patients with SARS who were considered in the present study, it is clear that Taiwan experienced 5 waves of epidemic SARS infection during 2003. The first and second waves were in different phylogenetic clusters, suggesting that they had different origins. Although both the second and the third waves were in the B2 cluster, the SARS-CoV isolates in the third wave had their own nucleotide-signature pattern (figure 5). Neither the first wave nor the second wave led to serious outbreaks, but the third wave—originating with a resident of Amoy Gardens who visited Tai-
Figure 6. Two possible paths of dissemination—between hospitals on Taiwan Island and a hospital on Penghu Island—of severe acute respiratory syndrome–associated coronavirus (SARS-CoV) infection. Numbers in shaded circles indicate the temporal order of nosocomial infections; numbers in parentheses indicate dates at onset of the infections.

It is important to note that TWC, the SARS-CoV strain isolated from patient TWC, clustered with WHU, an isolate from Wuhan City, China, in cluster 1 but did not cluster with either strain TC1 (direct sequencing of a sample from patient TWC) or strain TWC3 (direct sequencing of a sample from patient TW-HP1) (figures 3 and 4). In addition, there was a 7-nt difference between TWC and TC1 (figure 5). If we assume that strain CUHK-AG01 represents the first-generation SARS-CoV in the transmission link, then both strain TC1 and strain TWC3 were the second-generation, and TWH was the third-generation. According to the SNV-based analysis (figure 5), the number of nucleotide changes in the SARS-CoV genome per number of intermediate hosts was extremely low (<1 nt change/host). Because it has also been shown that no or very limited nucleotide changes occur in SARS-CoV sequences from either cultures or primary clinical specimens [11, 30], we can tentatively conclude that strain TWC is a laboratory contaminant and did not originate from patient TC1.

The origin of the fourth wave (cluster B3) is still unknown.
The molecular epidemiological data suggest that it originated at hospital JC (patient TW-JC2) and then spread to hospitals KC, GD, and YM, as well as to others. The epidemiological investigations also support this hypothesis: TW-PH1, a patient with SARS who was treated at hospital JC went to Kaohsiung City and received treatment at hospital KC, and, after being treated at hospital KC (table 1), went back to Penghu Island and was hospitalized in hospital PH, where he transmitted the disease to other health-care workers. As shown in figure 6, after combining the epidemiological data with the results of our phylogenetic-tree analysis, we conclude that the fourth wave of epidemic SARS infection progressed in 2 dissemination paths: the first path was from hospital JC to hospital KC in Kaohsiung and then to hospital PH on Penghu Island; the second path was from hospital JC to hospital GD and then to hospital YM. Two cases associated with community outbreaks in Taipei (strains TW10 and TW11) also belong to this cluster.

With regard to the laboratory contamination, the laboratory researcher said that he had used SARS-CoV strain HKU-39849 for his experiment; however, our results indicate that the sequence in patient SCVJ is more closely related to that of strain TWC (figure 5). Because the researcher claimed that he had not obtained strain TWC from Taiwan’s CDC and that he had used many clinical SARS-CoV strains besides HKU-39849, we are continuing our investigation to confirm both (1) whether the virus that he had obtained from Taiwan’s CDC was in fact strain HKU-39849 or strain TWC and (2) which SARS-CoV strain that he handled was the source of the laboratory contamination. Because >30 imported SARS cases have been reported by Taiwan’s CDC but have not yet been analyzed, we plan to use the proposed tool to conduct further analyses, in an attempt to identify their origins. These contact histories will provide valuable information for the control of future SARS infections.

**NUCLEOTIDE-SEQUENCE ACCESSION NUMBERS**

The SARS-CoV nucleotide sequences (6 sequences for each of 18 strains) identified during this research have been deposited in GenBank (accession numbers AY451856–AY451963). The reference sequences (accession numbers) used in our sequence-variation analysis were from the following 20 strains: Urbani (AY278741), CUHK-W1 (AY278554), TOR2 (AY274119), HKU-39849 (AY278491), BJ01 (AY278488), BJ02 (AY278487), BJ03 (AY278490), BJ04 (AY279354), GD01 (AY278489), TW1 (AY291451), TW2 (AY321118), SIN2774 (AY283798), SIN2748 (AY283797), SIN2679 (AY283796), SIN2677 (AY283795), SIN2500 (AY283794), HSR1 (AY323977), CUHK-Su10 (AY282752), Frankfurt1 (AY291315), and GZ50 (AY304495). In addition, 58 SARS-CoV genomes from GenBank were used for comparisons of phylogenetic analyses using the conventional and proposed strategies: TW9 (AY502932), TW8 (AY502931), TW7 (AY502930), TW6 (AY502929), TW5 (AY502928), TW4 (AY502927), TW3 (AY502926), TW2 (AY502925), TW11 (AY502924), TW10 (AY502923), GZ02 (AY390556), ZS-C (AY395003), LC5 (AY395002), LC4 (AY395001), LC3 (AY395000), LC2 (AY394999), LC1 (AY394998), ZS-A (AY394997), ZS-B (AY394996), HSZ-Cc (AY394995), HSZ-Ab (AY394994), HSZ-S2 (AY394993), HSZ-Fc (AY394991), HSZ-E (AY394990), HSZ-D (AY394989), HSZ-Fb (AY394987), HSZ-Cb (AY394986), HSZ-Bb (AY394985), HSZ-S2-A (AY394983), GZ-F (AY394979), GZ-B (AY394978), NS-1 (AY308724), WHU (AY394850), ShanghaiQXC1 (AY463059), ShanghaiQXC2 (AY463060), GD69 (AY313906), FRA (AY310120), SoD (AY461660), Sinol-3 (AY485278), Sinol-11 (AY485277), CUHK-AG03 (AY345988), CUHK-AG02 (AY345987), CUHK-Ag01 (AY345986), PUMC03 (AY357076), PUMC02 (AY357075), PUMC01 (AY350750), GZ50 (AP004495), TW35 (AP006299), TW2 (AP006298), ZMY1 (AY351680), TWY (AP006561), TWS (AP006560), TWK (AP006559), TWI (AP006558), TWH (AP006557), TC3 (AY348314), TC2 (AY338175), and TC1 (AY338174). Two civet-cat SARS-CoV strains (SZ3 [AY304495] and SZ16 [AY304488]) were used as the outgroup of the rooted trees [19].

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