Genetic mutations play a central role in evolution. For a significantly beneficial mutation, a one-time mutation event suffices for the species to prosper and predominate through the process called “monophyletic selective sweep.” However, existing methods that rely on counting the number of mutation events to detect selection are unable to find such a mutation in selective sweep. We here introduce a method to detect mutations at the single amino acid/nucleotide level that could be responsible for monophyletic selective sweep evolution. The method identifies a genetic signature associated with selective sweep using the population genetic test statistic Tajima’s D. We applied the algorithm to ebolavirus, influenza A virus, and severe acute respiratory syndrome coronavirus 2 to identify known biologically significant mutations and unrecognized mutations associated with potential selective sweep. The method can detect beneficial mutations, possibly leading to discovery of previously unknown biological functions and mechanisms related to those mutations.

**IMPORTANCE** In biology, research on evolution is important to understand the significance of genetic mutation. When there is a significantly beneficial mutation, a population of species with the mutation prospers and predominates, in a process called “selective sweep.” However, there are few methods that can find such a mutation causing selective sweep from genetic data. We here introduce a novel method to detect such mutations. Applying the method to the genomes of ebolavirus, influenza viruses, and the novel coronavirus, we detected known biologically significant mutations and identified mutations the importance of which is previously unrecognized. The method can deepen our understanding of molecular and evolutionary biology.

**KEYWORDS** evolution, selective sweep, genomes, influenza, ebolavirus, SARS-CoV-2
Viruses undergo adaptive genetic mutations within individual hosts and populations of the hosts. Establishing infection in new hosts often requires numerous adaptive changes, such as receptor specificity adjustments (3–5) and overcoming host antiviral defenses (5–7). An example of diversifying selection is the evolution of the hemagglutinin (HA) gene of influenza A virus to evade the host’s immune response, leading to antigenic drift (8–10). Polyphyletic convergent evolution can be observed in the emergence of drug-resistant mutations; for example, the same mutation in the neuraminidase (NA) gene of influenza A virus arose independently in patients treated with oseltamivir (11–13). When a mutation is substantially advantageous for viral replication or transmission, strains with the mutation will reproduce more effectively than other
strains, outcompeting those without the mutation (14, 15). This is what we call “monophyletic selective sweep” in this study.

Many methods for detecting signatures of natural selection in genomic data were first applied using viral data because of the rapid evolution of viruses. Calculation of relative ratios of nonsynonymous and synonymous mutations (dN/dS) is a popular way to detect diversifying selection and purifying selection (16, 17). Detection of polyphyletic convergent evolution can be accomplished by studying biased frequencies in protein substitution patterns (18, 19). These methods count the number of mutation events through an evolutionary pathway. Yet a mutation responsible for monophyletic selective sweep occurs only once in an evolutionary pathway. When there is a significantly beneficial mutation, a one-time mutation event must suffice for the viral strain to prosper and dominate an entire population (20). Therefore, the methodology of counting the number of mutation events cannot detect such a mutation leading to monophyletic selective sweep. Methods of detecting selective sweep are available for entire genes or alleles (21), and a few tools to detect a specific region or mutation causing selective sweep are also available (18, 22–25). However, because they do not consider the phylogenetic relationship of strains with a beneficial mutation, they could fail to detect monophyletic selective sweep.

Here, we introduce a method to detect mutations at the single amino acid/nucleotide level that could be responsible for monophyletic selective sweep, Tajima’s D-based identification of mutation associated with monophyletic selective sweep (DMAMS). The method detects mutations associated with monophyletic selective sweep with a statistical test using Tajima’s D, which is a scaled ratio of two measures of genetic diversity—the mean number of pairwise differences and the number of segregating sites—to test neutral evolution through a site-frequency spectrum (26). A negative Tajima’s D value suggests recent selective sweep or population expansion, whereas a positive value suggests balancing selection or sudden population contraction. In this study, we establish our algorithm, DMAMS, which detects a genetic signature (a certain amino acid or nucleotide at a certain position) associated with selective sweep by identifying a monophyletic population with the specific genetic signature that has a large-magnitude negative Tajima’s D value. We further demonstrate the application of the algorithm to ebolavirus, influenza A virus, and severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) to determine whether known biologically significant mutations and unrecognized mutations are associated with potential selective sweep through evolution.

RESULTS

Algorithm and concept of DMAMS. Figure 2 illustrates the DMAMS algorithm. Details of the algorithm are described in Materials and Methods. Briefly, DMAMS determines if there is a monophyletic cluster of strains that share a specific mutation, scanning all 20 amino acids at every position. All nodes in the phylogenetic tree are scanned for each amino acid at each site for clustering. When it finds a cluster of monophyletic strains with a certain amino acid at a certain position that is not shared with strains outside of the monophyletic cluster, Tajima’s D for the cluster is computed to check potential selective sweep. Tajima’s D values are also computed for internal nodes, comparing the values at nodes in the cluster with the values at other nodes outside the cluster to see the cluster-specific effect.

Population growth and selective sweep are known to have similar effects on Tajima’s D (26). The purpose of DMAMS is to detect selective sweep of strains with a certain mutation in a monophyletic subpopulation. Because monophyletic selective sweep is a result of population expansion of a subgroup that consists of monophyletic strains with a certain genetic signature (Fig. 1), differentiation of selective sweep from population expansion is not required in DMAMS. Furthermore, DMAMS will not be affected by changes in an entire population size. When a whole population has expanded and monophyletically clustered strains have shared a certain mutation just
by chance, DMAMS will not detect the mutation as associated with selective sweep. That is because DMAMS does not only calculate Tajima’s $D$ values for nodes inside the cluster and in the rest of the tree are also computed. There are two criteria to detect a cluster under monophyletic selective sweep: one is an absolute criterion that Tajima’s $D$ of the cluster is significantly low, and the other is a relative criterion that Tajima’s $D$ values at each node in the cluster are significantly smaller than the values at each node in the rest part of the tree. When both criteria are met, the genetic signature of the monophyletic cluster is considered to have an association with potential selective sweep.

**Simulation of evolution and performance of DMAMS.** To test the performance of DMAMS, evolution of a viral genome was simulated, and we applied DMAMS to the data generated by the simulations. Details of the simulations are given in Materials and Methods. Phylogenetic trees for virtual viral strains from our simulations reproduced characteristic features of five evolutionary scenarios: neutral evolution, diversifying selection, purifying selection, polyphyletic convergent evolution, and monophyletic selective sweep (Fig. 1). The trees showed unbiased distribution of mutations in a designated site in the neutral evolution model, a wide variety of mutations on the designated site in the diversifying selection model, a conserved polymorphism on the designated site in the purifying selection model, multiple independent occurrences of a designated mutation in the designated site in the polyphyletic convergent evolution model, and rapid expansion of a monophyletic subpopulation of strains with the designated mutation in the designated site in the monophyletic selective sweep model.

Algorithms named SLAC, FUBAR, and MEME for analyzing diversifying selection using different assumptions and statistical models (27, 28) detected selection on the designated site in 35% to 79% of the diversifying selection simulations (Fig. 3A). SALC and FUBAR can also be utilized to detect purifying selection. They detected purifying selection on the designated site in 81% to 82% of the purifying selection simulations. Another algorithm for convergent evolution named DEPS (18) detected selection for the designated mutation in the designated site in 59% of the polyphyletic convergent evolution model, and rapid expansion of a monophyletic subpopulation of strains with the designated mutation in the designated site in the monophyletic selective sweep model.

Algorithms named SLAC, FUBAR, and MEME for analyzing diversifying selection using different assumptions and statistical models (27, 28) detected selection on the designated site in 35% to 79% of the diversifying selection simulations (Fig. 3A). SALC and FUBAR can also be utilized to detect purifying selection. They detected purifying selection on the designated site in 81% to 82% of the purifying selection simulations. Another algorithm for convergent evolution named DEPS (18) detected selection for the designated mutation in the designated site in 59% of the polyphyletic convergent evolution simulations. DMAMS detected the beneficial mutation that had been accumulated polyphyletically by convergent evolution in only 10% of our simulations. Since DMAMS was developed to detect selective sweep of strains with a certain mutation in a monophyletic subpopulation, the sensitivity of DMAMS was not high enough.
FIG 3 Detection of selection by different methods for different evolutionary scenarios. (A) Proportion of detection of selection at a specific site by various tools for 100 simulations under the neutral evolution, diversifying selection, purifying selection, polyphyletic convergent evolution, and monophyletic selective sweep models. Population size, 200 for all simulations. For the monophyletic selective sweep model, the selection coefficient by a beneficial genetic signature is 2.0. (B) Proportion of detection of selection at a specific site by DMAMS for 100 simulations under the neutral evolution and monophyletic selective sweep models. An entire population size is either decreasing (contraction) or increasing (growth). For the monophyletic selective sweep model, the selection coefficient by a beneficial genetic signature is 2.0. Details of the population change in the simulations are described in Materials and Methods. (C) Proportion of detection of selection at a specific site by DMAMS for 100 simulations under the neutral evolution and monophyletic selective sweep models using the evolutionary model with parameters obtained from the HA gene of influenza A virus. Population size, 200. For the monophyletic selective sweep model, the selection coefficient by a beneficial genetic signature is 2.0. (D) Proportion of detection of selection at a specific site by DMAMS for 100 simulations under the monophyletic selective sweep model according to the selection coefficient. Population size, 200. (E) Proportion of detection of selection at a specific site by DMAMS for 100 simulations under the monophyletic selective sweep model according to the effective population size. The selection coefficient by a beneficial genetic signature is 2.0. (F) Proportion of detection of selection at a specific site by DMAMS for 100 simulations under the monophyletic selective sweep model according to the sampling size. Population size, 400. The selection coefficient by a beneficial genetic signature is 2.0.
to detect a beneficial mutation under polyphyletic convergent evolution. Those tools, SLAC, FUBAR, MEME and DEPS, detected selection in 3% to 17% of the monophyletic selective sweep simulations. Other tools to detect a region under selective sweep, SweeD and OmegaPlus, also did not detect the beneficial mutation in the monophyletic selective sweep simulations. This is likely because their algorithms are not suitable for detecting selective sweep that has caused a selection advantage in monophyletic strains. Details are presented in Discussion. In contrast, DMAMS detected selective sweep by the designated mutation in the designated site in 97% of the monophyletic selective sweep simulations. The false-positive rate of DMAMS was 2% (positivity in the neutral evolution simulations).

Sensitivity and specificity of DMAMS in different evolutionary scenarios and population models. We then tested the performance of DMAMS for a viral population whose effective population size has expanded or contracted because Tajima’s D is strongly affected by changes in population size. When a whole viral population has expanded and monophyletically clustered strains in a subpopulation have shared a certain mutation just by chance, DMAMS did not detect the mutation as associated with selective sweep (i.e., few false positives, as shown in the left side of Fig. 3B). Also, when a whole viral population size has contracted but monophyletically clustered strains have predominated because of monophyletic selective sweep by a certain beneficial mutation, DMAMS detected the mutation as associated with selective sweep with high sensitivity (i.e., few false negatives, as shown in the right side of Fig. 3B). The good performance was achieved because DMAMS not only calculates Tajima’s D of the cluster of strains with a mutation of interest ensuring selective sweep but also compares Tajima’s D values for nodes inside the cluster and the values for nodes in the rest of the tree to see the effect that is specific in the monophyletic cluster of strains with the certain mutation.

We also simulated the evolution of a viral genome assuming a general time reversible (GTR) nucleotide substitution model with gamma distribution of evolutionary rates among sites using parameters obtained from data of the HA gene of human influenza A virus, along with a deleterious effect for most nonsynonymous mutations, to reproduce a more realistic evolution of the viral genome. DMAMS still detected a beneficial mutation as associated with selective sweep in 90% of the monophyletic selective sweep simulations (i.e., true positives) and detected a nonbeneficial mutation in only 1% of the neutral evolution simulations (i.e., false positives) (Fig. 3C).

The high sensitivity in detecting a genetic signature causing monophyletic selective sweep by DMAMS was achieved when the selection coefficient was 200% (i.e., strains with the designated genetic signature were twice as likely to produce offspring in the next round as were strains without the genetic signature) (Fig. 3A to C). The sensitivity of DMAMS depends on the degree-of-fitness increase by the genetic signature (Fig. 3D). The sensitivity of DMAMS also depends on the effective population size and sampling size (Fig. 3E and F). The increase in Tajima’s D value is attributed to the low sensitivity of DMAMS in evolutionary scenarios with a small selection coefficient and small effective population size (see Fig. S1A to C in the supplemental material). Changing the parameter setting for “minimum size of cluster to test” in DMAMS might restore sensitivity in some scenarios (Fig. S1D). On the other hand, the specificity of DMAMS was robust over those different evolutionary scenarios and settings in DMAMS (Fig. S2).

Application of DMAMS to viral data. We applied DMAMS to real viral genetic/genomic data. As described in the introduction, DMAMS was developed to detect not sites with frequent mutations but genetic signatures associated with monophyletic selective sweep. Ebolavirus has been introduced into human populations several times (29), and the largest outbreak occurred in West Africa in 2014 to 2015 (30). A mutation at position 82 of the glycoprotein (GP) gene of the virus was reported under positive selection using the calculation of dN/dS (i.e., diversifying selection) (31). The mutation could affect its infectivity, which may have led to the large outbreak (31, 32). The diversifying selection on the site was detected using not only viral genetic data during the
2014-2015 outbreak but also historical data since 1976 (31). Although repeated jumps of the host species barrier from a natural host to humans made the possibly adaptive mutation at position 82 of the GP gene evident, whether the mutation affected the spread of the disease in 2014 to 2015 remains inconclusive (33). In fact, when we analyzed sequence data collected only during the 2014-2015 outbreak, neither SLAC nor DEPS detected positive selection on the site because of the limited number of mutation events on the site in the single outbreak (Table 1). In contrast, DMAMS identified a monophyletic cluster with the genetic signature 82V in the GP gene (Fig. S3A) and found an association with potential selective sweep. The result provides supporting evidence for the importance of the mutation in the spread of the disease during the outbreak. DMAMS also found other genetic signatures associated with monophyletic selective sweep during the 2014-2015 outbreak (Table 1). Among them, 111C in the nucleoprotein (NP) gene and 759G in the RNA-dependent RNA polymerase encoded in the L gene were reported to affect the replication efficiency shown by molecular experiments (34).

We next tested if drug-resistant mutations of influenza A virus were associated with monophyletic selective sweep because of the global transmission and spread of drug-resistant strains among human populations (35–37). Oseltamivir-resistant mutations in the influenza A virus have been reported mainly in patients given the antiviral (12, 38), and calculation of dN/dS using sequence data of the NA gene of influenza A(H1N1) pdm/09 virus revealed diversifying selection on a site responsible for the drug sensitivity (39). However, in the 2007-2008 season, a strain of human H1N1 virus with a specific oseltamivir-resistant mutation, 275Y in the NA gene, emerged, and its monophyletic descendants spread to people, including those without a history of antiviral use (36, 40). We analyzed genetic data of the virus collected in that single season, because the effective population size of influenza A virus fluctuates from season to season (41). We identified both a monophyletic cluster with the genetic signature 275Y and sporadic occurrence of strains with the mutation (Fig. S3B). Whereas DEPS identified a significantly high frequency of 275Y substitutions, DMAMS showed that Tajima’s $D$ of the monophyletic cluster of strains with the mutation was not smaller than that of the other strains without the mutation (Table 1). The result suggests that there was no association between the monophyletically clustered drug-resistant strains and selective sweep at a population level, which seems reasonable because the antiviral was not administered to most people for prophylaxis or treatment.

We also utilized DMAMS to investigate intrahost viral evolution (i.e., quasispecies). Rogers et al. reported a case of influenza infection by analyzing the mutational spectrum of an H3N2 influenza A virus population sampled from an immunocompromised patient who had shed virus over a 21-month period (42). The patient was infected with quasispecies of drug-sensitive and -resistant viruses during the course of treatment with neuraminidase inhibitors. Although the study by Rogers et al. simply investigated the time course trend of frequency of drug-resistant mutations, DMAMS can take advantage of their data because they used PacBio single-molecule sequencing, which yielded read lengths long enough to perform phylogenetic analysis. It is important to mention that there must be an expansion of population size in intrahost evolution of the virus. Still, DMAMS is not affected by such changes in an entire population size, as shown in our simulation (Fig. 3B).

Two mutations responsible for resistance against a neuraminidase inhibitor, 119V and 222V encoded in the NA gene, emerged during the treatment. Unfortunately, the number of long reads covering the region including position 119 was not sufficient to perform phylogenetic analysis; we focused on position 222. At the beginning of the appearance of the drug-resistant 222V mutation, strains with the mutations did not form a phylogenetic cluster (Fig. S3C). Two weeks after treatment began, the proportion of strains with the drug-resistant mutations increased (42). DMAMS found that the 222V mutation was associated with a monophyletic cluster under potential selective sweep, while the site showed a dN/dS of $<1$ (Table 1 and Fig. S3D). The results indicate
| Virus          | Gene                  | Time           | No. of sequences (after exclusion of identical sequences) | Genetic signature of interest | Diversifying selection | Statistical significance | Convergent evolution with statistical significance | Cluster with genetic signature of interest | Low Tajima's D (less than 1.5) for the cluster | Smaller Tajima's D for strains in the cluster compared with strains without the signature | Statistical significance | Other genetic signatures associated with the same cluster | Detected genetic signatures associated with potential selective sweep (other than genetic signature of interest and other signatures in the same cluster) |
|---------------|-----------------------|----------------|-------------------------------------------------------------|------------------------------|------------------------|-------------------------|---------------------------------------------|-----------------------------------------------|-----------------------------------------------|-------------------------------------------------|-----------------------------|------------------------------------------------------|------------------------------------------------------------------------------------------------|
| Ebolavirus    | Whole genome          | 2014–2015      | 718                                                         | GP-82V                       | No                     | NA                      | No                           | Yes                                           | Yes                                           | Yes                                              | Yes                                          | Yes                                                  | VP35-330L, L-759G, NP-111C, VP35-249L, L-399P, L-1645S, 45N, 78E, 82S, 86A, 139R, 188M, 249K, 267I, 287I, 329E, 367L, 393V, 451S, 453T, 146K, 219T |
| H1N1 influenza A virus | Neuraminidase gene | 2007–2008      | 188                                                         | 275Y                         | Yes                    | No                      | Yes                           | Yes                                           | Yes                                           | No                                                | NA                                          | 354G                                                  |                                                                |
| H3N2 influenza A virus | Neuraminidase gene | Intrahost quasispecies (at the commencement of treatment) | 66                            | 222V                         | No                     | NA                      | No                           | No                                            | NA                                            | NA                                                | NA                                          | NA                                                    | None                                                                |
|               |                       | Intrahost quasispecies (2 wks after the treatment)     | 63                            | NA                           | NA                     | No                      | Yes                           | Yes                                           | Yes                                           | Yes                                              | Yes                                          | 242I                                                  | None                                                                |
| SARS-CoV-2    | Whole genome          | January–June 2020 | 2,309                                                      | S-614G                       | No                     | NA                      | No                           | Yes                                           | No                                            | NA                                                | NA                                          | None                                                  | ORF1a-265L, ORF3a-57Q, N-203K, N-204R                             |

*Phylogenetic trees for the analyses can be found in Fig. S3 in the supplemental material. NA, not available.

*dN/dS of > 1 at the site of interest analyzed using SLAC was regarded as diversifying selection.
that the drug exerted evolutionary pressure at an individual host level to induce not
diversifying selection but monophyletic selective sweep caused by the mutation.
Another genetic signature, 242I, was found in the same cluster under potential selec-
tive sweep along with 222V (Table 1). Acquisition of a drug-resistant mutation is known
to lead to a reduction in viral growth efficiency (43). The 242I mutation warrants further
investigation; it may have the potential to compensate for a decrease in viral growth
efficiency by a drug-resistant mutation.

Finally, we applied DMAMS to the data of SARS-CoV-2. SARS-CoV-2 appeared at the
end of 2019 and caused a worldwide pandemic in 2020 (44). A recent study reported
an increase in frequency of a certain mutation, D614G in the S gene encoding spike
protein; the mutation potentially increases the fitness of the virus, as shown by clinical
and experimental data (45, 46). Still, the significance of the mutation for public health
in the real world is controversial (47–49). Using sequence data of the virus published
by July 2020 from all over the world, a phylogenetic tree of the virus showed multiple
substitution events between D and G on the site and a monophyletic cluster with the
genetic signature 614G (Fig. S3E). The site was not under diversifying selection eval-
uated by SLAC or under convergent evolution toward G detected by DEPS (Table 1). In
addition, DMAMS did not show the 614G-specific population growth of monophyletic
strains (Table 1). Actually, not only the monophyletic strains with the genetic signature
but also strains outside of the cluster without the genetic signature had large-magni-
tude negative Tajima’s D values, −2.79 and −2.80, respectively. The result reflects a
rapid growth in population size of the novel virus by a global pandemic in a naive pop-
ulation. There was no significant difference in Tajima’s D values between strains with
and without the 614G mutation.

DISCUSSION

Here, we report an approach to detecting genetic signatures associated with mono-
phyletic selective sweep. Our method simply uses genetic sequence information as
data and can identify possibly beneficial mutations even if the mutation event
occurred only once throughout an evolutionary pathway. We also showed a proof-of-
concept application for viral genetic data.

Previously developed tools to detect selective sweep do not consider monophyletic
subpopulations with beneficial genetic signatures (22–25). In addition, they were
developed to detect a gene or a subregion of a gene under selective sweep; they could
not specifically identify a specific site associated with the selective sweep. Furthermore, most of such tools, including SweeD and OmegaPlus tested in the pres-
et study, were built for a diploidy genome (24, 25). They assume accumulation of al-
leles with characteristic genetic signatures by recombination, and they search the
region under selective sweep by checking linkage disequilibrium. Therefore, they are
not applicable to our idea of monophyletic selective sweep, that is, the expansion of a
monophyletic subpopulation caused by a one-time beneficial mutation. Looking at
this from the opposite perspective, DMAMS cannot be applied to data that include
sequences generated by recombination. This is because DMAMS depends on recon-
struction of the evolutionary path by a phylogenetic tree. The phylogenetic tree using
whole sequences cannot reflect the true evolutionary path when recombination has
occurred.

Another tool to detect convergent evolution, DEPS (18), was not sensitive enough
to detect a mutation under directional evolution that is shared only among monophy-
letic strains (Fig. 3A). Our algorithm, DMAMS, can identify a mutation associated with
monophyletic selective sweep even when the mutation event has occurred only once
in an evolutionary pathway. Still, it should be noted that DMAMS is not sensitive
enough to detect a beneficial mutation shared among polyphyletic strains (i.e., poly-
phyletic convergent evolution). Such a mutation is not the target for DMAMS. In other
words, DMAMS is not a tool to replace existing tools for detecting (polyphyletic)
convergent evolution but is a complementary tool to evaluate the significance of mutations shared only among monophyletic strains.

Because DMAMS utilizes a site-frequency spectrum of genetic sequences by calculating Tajima’s D, it can detect genetic signatures not only of deduced amino acids in a coding region but also nucleotides in both coding and noncoding regions. With increasing understanding of the functional roles of genetic sequences themselves, including regulation of DNA/RNA stability, transcription and translation efficiencies, and interaction with other molecules (50–53), methods of identifying genetic signatures responsible for selective sweep at the nucleotide level should be of great interest. Unfortunately, because genetic data of noncoding regions were not abundant compared to those of coding regions, we could not find a good example to show this advantage of our method.

We showed that the sensitivity of DMAMS depends on the magnitude of the selection coefficient given by a beneficial mutation. We defined the selection coefficient in our simulations as the likelihood to produce offspring in the next round of generation (Fig. 3D). In real data of viral evolution, the definition of generation is context dependent, such as “a single round of replication in a cell” or “transmission from one host to another,” and those are difficult to observe. Therefore, magnitudes of selection coefficient are hard to be inferred using real viral genetic data. It is important to note that no detection of mutations associated with selective sweep by DMAMS does not necessarily mean that there is no selective sweep, especially when the degree of fitness by a beneficial mutation is not very high.

Our method is limited by the fact that DMAMS cannot detect causal relationships between genetic signatures and selective sweep but simply detects an association between them. Theoretically, DMAMS cannot distinguish a true beneficial mutation causing monophyletic selective sweep apart from “hitchhiking” mutations that were introduced simultaneously with the beneficial mutation (54, 55). There is also a caveat that the data set for DMAMS should include only strains circulating in the same environment. For example, when DMAMS detects a mutation that was found only in a specific geographical area as a mutation associated with potential selective sweep, it is possible that the expansion of the subpopulation is not caused by selective sweep by the certain mutation but is due to external factors that are different between the two areas, such as size and density of the host population, host genetic factors, and environmental conditions. Mutations identified by DMAMS should be further investigated for possible confounding with other factors.

As we have seen, the emergence of mutation and its fixation process can be observed in real time, particularly with rapidly evolving viruses (2, 56). DMAMS can identify genetic signatures associated with monophyletic selective sweep. However, it cannot prove a causal relationship between mutation and selective sweep. Further studies are required to show the biological significance of identified mutations. Even though DMAMS cannot yield direct evidence of the evolutionary significance of a certain mutation, it is a useful tool, like genome-wide association study, to identify a candidate mutation that is potentially important (57). DMAMS has the potential to reveal previously unrecognized biological functions and mechanisms, thereby deepening our understanding of molecular and evolutionary biology.

MATERIALS AND METHODS

The DMAMS algorithm. Aligned genetic (DNA or RNA) sequence data and their associated rooted phylogenetic trees were used to detect genetic signatures associated with monophyletic selective sweep. Figure 2 illustrates the schematic principle of the Tajima’s D-based identification of mutation associated with monophyletic selective sweep (DMAMS) algorithm. First, a rooted phylogenetic tree was scanned for a monophyletic cluster with a genetic signature that is rarely shared with strains outside the cluster. Genetic signature can be either an encoded amino acid or a nucleotide at a certain position. DMAMS scans all 20 amino acids (or 4 nucleotides) at every position of sequence data. In addition, all nodes in the phylogenetic tree are scanned for each amino acid (or each nucleotide) at each site for clustering. The minimum cluster size to test can be determined at the user’s discretion. In the present paper, the parameter was set at 5% for the minimum cluster size for all analyses unless otherwise noted. When the proportion of strains with a genetic signature in a monophyletic subgroup was higher than 90%
and the proportion of strains with the genetic signature in the rest of the data set was smaller than 50%, the monophyletic subgroup was regarded as a cluster with the genetic signature. Although ideally 100% of the monophyletically clustered subpopulation shares the same beneficial mutation, there might be further mutations resulting in the loss of the beneficial mutation in some strains in the subpopulation. Therefore, we set a 10% margin. For example, when “the minimum size of cluster to test” is 5%, the minimum percentage required for the detection of a potentially beneficial mutation by DMAMS is (5% × 90%) = 4.5%. We regarded that as small enough to detect selective sweep in its early stage.

When a monophyletic cluster with a certain genetic signature was detected, Tajima’s D values were also computed for nodes inside the cluster and for nodes in the rest of the tree, excluding the outgroup used for rooting the phylogenetic tree. The nodes for the calculation of Tajima’s D should include at least the minimum size of strains for a cluster. Similar inferential methods to measure fitness (i.e., selection coefficient) of strains in a phylogenetic clade are discussed elsewhere (10, 23, 58, 59). We set two criteria to detect a monophyletic cluster under potential selective sweep. One is an absolute criterion that Tajima’s D of the cluster is lower than −1.5 (with approximately 90% confidence [26]), and the other is a relative criterion that Tajima’s D values at each node in the cluster are significantly smaller than the values at each node in the rest of the tree. The absolute criterion is to detect potential selective sweep for monophyletically clustered strains with a genetic signature. The relative criterion is to see their cluster-specific selective sweep, excluding the effect of changes in effective population size of an entire population. When both criteria were met, the genetic signature of the cluster was considered to have an association with potential selective sweep. For the relative criterion, comparison of Tajima’s D values at nodes in the cluster with the values at nodes outside the cluster was made with the Mann-Whitney U test. A P value of less than 0.05 was considered statistically significant after adjustment for false discovery using the Benjamini-Hochberg procedure (60). The algorithm of DMAMS was implemented in a script we developed, which is available at https://github.com/yukifuruse1217/DMAMS.

Simulation of evolution. The evolution of viral genes was simulated with a constant population size of 50 to 400 or under an exponential growth or contraction population model. The effective population size increases from 10 to 400 at 1,000 rounds of generation in the slow growth model, it increases from 10 to 400 at 100 rounds of generation in the rapid growth model, and it decreases from 1,000 to 200 at 1,000 rounds of generation in the contraction model. The virtual gene was composed of 3,000 nucleotides, and the whole region of the gene had an open reading frame (i.e., 1,000 deduced amino acids). In a given round, each strain produced a number of offspring to populate the next round depending on the fitness of the strain determined by the selection coefficient. Nucleotide mutation (substitution) occurred at a rate of 0.002 per site per round; the substitution rates among four nucleotides were the same (i.e., Jukes-Cantor model). Five evolutionary models for simulations to change fitness according to mutation at a designated site were included: neutral evolution, diversifying selection, purifying selection, polyphyletic convergent evolution, and monophyletic selective sweep.

In the neutral evolution model, every strain has the same fitness regardless of mutations. In the diversifying selection model, strains increase the fitness when any nonsynonymous mutations occur at a designated position. In the purifying selection model, strains decrease the fitness when any nonsynonymous mutations occur at a designated position. In the polyphyletic convergent evolution model, strains increase the fitness when a designated nonsynonymous mutation occurs at a designated position. In the monophyletic selective sweep model, strains exhibit high fitness throughout generations as long as they possess a designated genetic signature at a designated position. The degree to increase fitness (selection coefficient) by the designated mutation subject to change to investigate the sensitivity of DMAMS, ranging between 1.044 and 2.0. For example, when the selection coefficient is 2.0, strains with the designated genetic signature are twice as likely to produce offspring in the next round than strains without the genetic signature. Simulations were run 100 times for each setting and stopped at 2,000 to 4,000 rounds of generation for neutral evolution, diversifying selection, and purifying selection models. Simulations were stopped when the proportion of strains with a designated beneficial genetic signature exceeded 30% of the entire population for the polyphyletic convergent evolution and monophyletic selective sweep models.

In addition, simulations that are more realistic in reproducing viral evolution were run. We acquired parameters for viral evolution using data of the HA gene of human influenza A (H3N2) virus between 2009 and 2018, assuming a GTR nucleotide substitution model plus gamma distribution for evolutionary rates among sites. Also, in the simulation, we assumed that nonsynonymous mutations decrease fitness except for a designated beneficial mutation. The all-evolutionary simulations were implemented in a script we developed, which is available at https://github.com/yukifuruse1217/DMAMS.

Tests for evolutionary selection using simulated data. Genetic sequences of strains at the final round of simulation were sampled along with the original sequence before introducing any mutation to root the phylogenetic tree. Phylogenetic trees were constructed using a maximum-likelihood method with the Jukes-Cantor model or the GTR plus gamma distribution model using MEGA7 (61). The sequence data and phylogenetic tree were used to test diversifying selection by SLAC (27), FUBAR (62), and MEME (63), to test purifying selection by SLAC and FUBAR, to test convergent evolution by DEPS (18), and to test selective sweep by SweeD3.0 (25), OmegaPlus (24), and DMAMS.

Viral genetic data. Genetic sequences of the whole genome of ebolavirus collected in West Africa between 2014 and 2015 and the NA gene of human H1N1 influenza A virus from the Northern Hemisphere in the 2007-2008 season were obtained from GenBank via Virus Variation Resource (https://www.ncbi.nlm.nih.gov/genome/viruses). All genetic sequences of the whole genome of SARS-CoV-2 were obtained from GenBank via SARS-CoV-2 Resources (https://www.ncbi.nlm.nih.gov/sars-cov-2/), accessed on 6 July 2020.
Genetic sequences of the NA gene for quasispecies of human H3N2 influenza A virus in a single patient were obtained from SRA (accession no. PRJNA253584) (42). Reads generated by PacBio were mapped to a reference genome [A(H3N2)/California/7/2004], and mapped reads were trimmed to include genetic signatures of interest (amino acid position 222) so that the final data have the maximized number of sequences.

Sequence data including ambiguous nucleotides were removed for further analyses. Strains with identical nucleotide sequences in the same data set were also excluded for two reasons. (i) Tajima’s D is a statistic to indicate change in population size and/or existence of selection. Having identical sequences in a data set will not give any additional information on them. (ii) Because DMAMS calculates Tajima’s D values at internal nodes of the phylogenetic tree, it will be impossible to compute the values when a node contains a cluster composed only of identical sequences.

Phylogenetic trees were constructed using a maximum-likelihood method based on a GTR substitution model using MEGA7 (61) for influenza virus and RAxML (64) for ebolavirus and SARS-CoV-2. The trees were rooted using an old strain of the same virus. Sequence data, along with the phylogenetic tree, were tested using SLAC, DEPS, and DMAMS to detect sites under evolutionary selection.

Data availability. All genetic data analyzed in the study were acquired from public databases, GenBank and SRA, as described above. The source code for DMAMS and evolutionary simulations developed in the present study can be found at GitHub (https://github.com/yukifuruse1217/DMAMS).

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.
FIG S1, EPS file, 0.7 MB.
FIG S2, EPS file, 0.01 MB.
FIG S3, TIF file, 0.9 MB.

ACKNOWLEDGMENTS
I thank Yoshio Koyanagi for his support.

This work was supported by a Grant-in-Aid for Scientific Research on Innovative Areas (grant number JP19H04832), a Grant-in-Aid for Scientific Research (grant number JP19K07576), and a Grant-in-Aid for Scientific Research on Promotion of Joint International Research (grant number JP19KK0204) from the Japan Society for the Promotion of Science, by a Research Program on Emerging and Re-emerging Infectious Diseases (grant number 19fk0108108h0001) from the Japan Agency for Medical Research and Development, by the Leading Initiative for Excellent Young Researchers (grant number 16809810) from the Ministry of Education, Culture, Sports, Science and Technology in Japan, and by the Future Development Funding Program (grant number not available) from Kyoto University Research Coordination Alliance.

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
I declare no conflicts of interest.

REFERENCES
1. Loewe L, Hill WG. 2010. The population genetics of mutations: good, bad and indifferent. Philos Trans R Soc Lond B Biol Sci 365:1153–1167. https://doi.org/10.1098/rstb.2009.0317.
2. Frost SDW, Magalis BR, Kosakovsky Pond SL. 2018. Neutral theory and rapidly evolving viral pathogens. Mol Biol Evol 35:1348–1354. https://doi.org/10.1093/molbev/msy088.
3. Taubenberger JK, Kash JC. 2010. Influenza virus evolution, host adaptation, and pandemnic formation. Cell Host Microbe 7:440–451. https://doi.org/10.1016/j.chom.2010.05.009.
4. Li W, Zhang C, Sui J, Kuhn JH, Moore MJ, Luo S, Wong SK, Huang IC, Xu K, Vasilieva N, Murakami A, He Y, Marasco WA, Guan Y, Choe H, Farzan M. 2005. Receptor and viral determinants of SARS-coronavirus adaptation to human ACE2. EMBO J 24:1634–1643. https://doi.org/10.1038/sj.emboj.7600640.
5. Letko M, Seifert SN, Olival KJ, Plovworth RK, Munster VJ. 2020. Bat-borne virus diversity, spillover and emergence. Nat Rev Microbiol 18:461–471. https://doi.org/10.1038/s41579-020-0394-z.
6. Sawyer SL, Emerman M, Malik HS. 2004. Ancient adaptive evolution of the primate antiviral DNA-editing enzyme APOBEC3G. PLoS Biol 2:e275. https://doi.org/10.1371/journal.pbio.0020275.
7. Sauter D, Kirchhoff F. 2019. Key viral adaptations preceding the AIDS pandemic. Cell Host Microbe 25:27–38. https://doi.org/10.1016/j.chom.2018.12.002.
8. Suzuki Y. 2006. Natural selection on the influenza virus genome. Mol Biol Evol 23:1902–1911. https://doi.org/10.1093/molbev/msl050.
9. Smith DJ, Lapedes AS, de Jong JC, Bestebroer TM, Rimmelzwaan GF, Osterhaus ADME, Fouchier RAM. 2004. Mapping the antigenic and genetic evolution of influenza virus. Science 305:371–376. https://doi.org/10.1126/science.1097211.
10. Neher RA, Bedford T, Daniells RS, Russell CA, Shraiman BI. 2016. Prediction, dynamics, and visualization of antigenic phenotypes of seasonal influenza viruses. Proc Natl Acad Sci U S A 113:E1701–E1709. https://doi.org/10.1073/pnas.1525578113.
11. Monto AS, McKimm-Breschkin JL, Macken C, Hampson AW, Hay A, Klimov A, Tashiro M, Webster RG, Aymard M, Hayden FG, Zambon M. 2006. Detection of influenza viruses resistant to neuraminidase inhibitors in global surveillance during the first 3 years of their use. Antimicrob Agents Chemother 50:2395–2402. https://doi.org/10.1128/AAC.01339-05.
12. Kiso M, Mitamura K, Sakai-Tagawa Y, Shiraishi K, Kawakami C, Kimura K, Hayden FG, Sugaya N, Kawaoka Y. 2004. Resistant influenza A viruses in...
children treated with oseltamivir: descriptive study. Lancet 364:759–765. https://doi.org/10.1016/S0140-6736(04)16934-1.

13. Wang B, Dwyer DE, Blyth CC, Soedjono M, Shi H, Kessor A, Ratnamohan M, McPhie K, Cunningham AL, Saksena NK. 2010. Detection of the rapid emergence of the H275Y mutation associated with oseltamivir resistance in severe pandemic influenza virus A/H1N1 09 infections. Antiviral Res 87:16–21. https://doi.org/10.1016/j.antiviral.2010.04.002.

14. Steinbrück L, McHardy AC. 2011. Allele dynamics plots for the study of evolutionary dynamics in viral populations. Nucleic Acids Res 39:e4. https://doi.org/10.1093/nar/gkr999.

15. Orr HA. 1998. The population genetics of adaptation: the distribution of factors fixed during adaptation evolution. Evolution 52:935–949. https://doi.org/10.2307/2411226.

16. Suzuki Y, Gojobori T. 1999. A method for detecting positive selection at single amino acid sites. Mol Biol Evol 16:1315–1328. https://doi.org/10.1093/oxfordjournals.molbev.a026042.

17. Yang Z, Nielsen R. 2000. Estimating synonymous and nonsynonymous substitution rates under realistic evolutionary models. Mol Biol Evol 17:32–43. https://doi.org/10.1093/oxfordjournals.molbev.a026236.

18. Kosakovsky Pond SL, Poon AFY, Leigh Brown AJ, Frost SDW. 2008. A maximum likelihood method for detecting directional evolution in protein sequences and its application to influenza A virus. Mol Biol Evol 25:1809–1824. https://doi.org/10.1093/molbev/msn123.

19. Zhang J, Kumar S. 2015. Detection of convergent and parallel evolution at the amino acid sequence level. Mol Biol Evol 14:527–536. https://doi.org/10.1093/oxfordjournals.molbev.a025789.

20. Charlesworth B. 2009. Fundamental concepts in genetics: effective population size and patterns of molecular evolution and variation. Nat Rev Genet 10:195–205. https://doi.org/10.1038/nrg2526.

21. Pavlidis P, Alachiotis N. 2017. A survey of methods and tools to detect recent and strong positive selection. J Bio Res (Thessalon) 24:7. https://doi.org/10.1093/oxfordjournals.molbev.a025789.

22. Klingen TR, Reimering S, Loers J, Mooren K, Klawonn F, Krey T, Gabriel G, Groschke C1, D, Stamatakis A, Alachiotis N. 2013. SweeD: likelihood-based detection of selective sweeps in thousands of genomes. Mol Biol Evol 30:2234. https://doi.org/10.1093/molbev/mst112.

23. Klingen TR, Reimering S, Loers J, Mooren K, Klawonn F, Krey T, Gabriel G, Groschke C1, D, Stamatakis A, Alachiotis N. 2013. SweeD: likelihood-based detection of selective sweeps in thousands of genomes. Mol Biol Evol 30:2234. https://doi.org/10.1093/molbev/mst112.

24. Rambaut A, Pybus OG, Nelson MI, Viboud C, Taubenberger JK, Holmes EC. 2008. The genomic and epidemiological dynamics of human influenza A virus. Nature 453:615–619. https://doi.org/10.1038/nature06945.

25. Rogers MB, Song T, Sebra R, Greenbaum BD, Hamelin M-E, Fitch A, Twaddle A, Cui L, Holmes EC, Boivin G, Ghe Din E. 2015. Intrahost dynamics of antiviral resistance in influenza A virus reflect complex patterns of segment linkage, reassortment, and natural selection. mBio 6:e00246-14. https://doi.org/10.1128/mBio.00246-14.

26. Kojima H, Herlocher LM, Hoffmann E, Matsushita MN, Monto AS, Webster RG, Govorkova EA. 2005. Neuraminidase inhibitor-resistant influenza viruses may differ substantially in fitness and transmissibility. Antimicrob Agents Chemother 49:4075–4084. https://doi.org/10.1128/AAC.49.11.4075-4084.2005.

27. Wiersinga WJ, Rhodes A, Cheng AC, Peacock SJ, Prescott HC. 2020. Pathophysiology, transmission, diagnosis, and treatment of coronavirus disease 2019 (COVID-19): a review. JAMA 324:782–793. https://doi.org/10.1001/jama.2020.12839.

28. Kerber B, Fischer WM, Gnanakaran S, Yoon H, Theiler J, Abfarletter W, Hengartner N, Giorgi EE, Bhattacharya T, Foley B, Hastie KM, Parker MD, Partridge DG, Evans CM, Freeman TM, de Silva TI, McDanal C, Perez LG, Tang H, Moon-Walker A, Whelan SP, LaBranche CC, Saphire EO, Monterfield DC, Angyal A, Brown RL, Carrillo L, Green LR, Groves DC, Johnson KJ, Keeley AJ, Lindsey BB, Parsons PJ, Raza M, Bowland-Jones S, Smith N, Tucker RM, Wang D, Wyles MD. 2020. Tracking changes in SARS-CoV-2 Spike: evidence that D614G increases infectivity of the COVID-19 virus. Cell 182:812–827.e19. https://doi.org/10.1016/j.cell.2020.06.043.

29. Hou YJ, Chiba S, Halfmann P, Ehre C, Kuroda M, Dinnon KH, Leist SR, Schäfer A, Nakajima T, Takashiki K, Lee RE, Mascenik TM, Graham R, Edwards CE, Tse LV, Okuda K, Markmann AJ, Bartelt D, da Silva A, Margolis DM, Boucher RC, Randell SH, Suzuki T, Gralinski LE, Kawaoka Y, Baric RS. 2020. SARS-CoV-2 D614G variant exhibits efficient replication ex vivo and transmission in vivo. Science 370:1464–1468. https://doi.org/10.1126/science.abe8499.

30. Grubaugh ND, Hanage WP, Rasmussen AL. 2020. Making sense of mutations: what D614G means for the COVID-19 pandemic remains unclear. Cell 182:794–795. https://doi.org/10.1016/j.cell.2020.06.040.

31. Vele E, Hill V, McCrone JT, Price A, Jorgensen D, O’Toule A, Southgate J, Johnson R, Jackson B, Nascimento FF, A, Craig S, Coworkers 2020. Evaluating the effects of SARS-CoV-2 spike mutation D614G on
transmissibility and pathogenicity. Cell 184:64–75.e11. https://doi.org/10.1016/j.cell.2020.11.020.

49. van Dorp L, Richard D, Tan CCS, Shaw LP, Acman M, Balloux F. 2020. No evidence for increased transmissibility from recurrent mutations in SARS-CoV-2. Nat Commun 11:5986. https://doi.org/10.1038/s41467-020-19818-2.

50. Chavali PL, Stojic L, Meredith LW, Joseph N, Nahorski MS, Sanford TJ, Sweeney TR, Krishna BA, Hosmillo M, Firth AE, Bayliss R, Marcelis CL, Lindsay S, Goodfellow I, Woods CG, Gergely F. 2017. Neurodevelopmental protein Musashi-1 interacts with the Zika genome and promotes viral replication. Science 357:83–88. https://doi.org/10.1126/science.aam243.

51. Jopling CL, Yi MK, Lancaster AM, Lemon SM, Sarnow P. 2005. Molecular biology: modulation of hepatitis C virus RNA abundance by a liver-specific microRNA. Science 309:1577–1581. https://doi.org/10.1126/science.1113329.

52. Lim CS, Brown CM. 2018. Know your enemy: successful bioinformatic approaches to predict functional RNA structures in viral RNAs. Front Microbiol 8:2582. https://doi.org/10.3389/fmicb.2017.02582.

53. Hutchinson EC, von Kirchbach JC, Gog JR, Digard P. 2010. Genome packaging in influenza A virus. J Gen Virol 91:313–328. https://doi.org/10.1099/vir.0.017608-0.

54. Fay JC, Wu CI. 2000. Hitchhiking under positive Darwinian selection. Genetics 155:1405–1413.

55. Messer PW, Neher RA. 2012. Estimating the strength of selective sweeps from deep population diversity data. Genetics 191:593–605. https://doi.org/10.1534/genetics.112.138461.

56. Geoghegan JL, Holmes EC. 2018. Evolutionary virology at 40. Genetics 210:1151–1162. https://doi.org/10.1534/genetics.118.301556.

57. Cantor RM, Lange K, Sinsheimer JS. 2010. Prioritizing GWAS results: a review of statistical methods and recommendations for their application. Am J Hum Genet 86:6–22. https://doi.org/10.1016/j.ajhg.2009.11.017.

58. Luksza M, Lässig M. 2014. A predictive fitness model for influenza. Nature 507:57–61. https://doi.org/10.1038/nature13087.

59. Villa M, Lässig M. 2017. Fitness cost of reassortment in human influenza. PLoS Pathog 13:e1006685. https://doi.org/10.1371/journal.ppat.1006685.

60. Benjamin Y, Hochberg Y. 1995. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J R Stat Soc Ser B 57:289–300. https://doi.org/10.1111/j.2517-6161.1995.tb02031.x.

61. Kumar S, Stecher G, Tamura K. 2016. MEGA7: Molecular Evolutionary Genetics Analysis version 7.0 for bigger datasets. Mol Biol Evol 33:1870–1874. https://doi.org/10.1093/molbev/msw054.

62. Murrell B, Moola S, Mabona A, Weighill T, Sheward D, Kosakovsky Pond SL, Scheffler K. 2013. FuBAR: a fast, unconstrained Bayesian approximation for inferring selection. Mol Biol Evol 30:1196–1205. https://doi.org/10.1093/molbev/msw054.

63. Murrell B, Wertheim JO, Moola S, Weighill T, Scheffler K, Kosakovsky Pond SL. 2012. Detecting individual sites subject to episodic diversifying selection. PLoS Genet 8:e1002764. https://doi.org/10.1371/journal.pgen.1002764.

64. Stamatakis A. 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30:1312–1313. https://doi.org/10.1093/bioinformatics/btu033.