# Within-host diversity improves phylogenetic and transmission reconstruction of SARS-CoV-2 outbreaks

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- Abstract Accurate inference of who infected whom in an infectious disease outbreak is critical
- <sup>17</sup> for the delivery of effective infection prevention and control. The increased resolution of
- <sup>18</sup> pathogen whole-genome sequencing has significantly improved our ability to infer transmission
- <sup>19</sup> events. Despite this, transmission inference often remains limited by the lack of genomic
- <sup>20</sup> variation between the source case and infected contacts. Although within-host genetic diversity is
- <sup>21</sup> common among a wide variety of pathogens, conventional whole-genome sequencing
- <sub>22</sub> phylogenetic approaches exclusively use consensus sequences, which consider only the most
- <sup>23</sup> prevalent nucleotide at each position and therefore fail to capture low frequency variation within
- samples. We hypothesized that including within-sample variation in a phylogenetic model would
- help to identify who infected whom in instances in which this was previously impossible. Using
   whole-genome sequences from SARS-CoV-2 multi-institutional outbreaks as an example, we
- whole-genome sequences from SARS-CoV-2 multi-institutional outbreaks as an example, we
- show how within-sample diversity is partially maintained among repeated serial samples from
   the same host, it can transmitted between those cases with known epidemiological links, and
- the same host, it can transmitted between those cases with known epidemiological links, and how this improves phylogenetic inference and our understanding of who infected whom. Our
- <sup>30</sup> technique is applicable to other infectious diseases and has immediate clinical utility in infection
- <sup>31</sup> prevention and control.
- 32
- 33 Introduction
- <sup>34</sup> Understanding who infects whom in an infectious disease outbreak is a key component of infection
- prevention and control (*Didelot et al., 2012*). The use of whole-genome sequencing allows for
- <sup>36</sup> detailed investigation of disease outbreaks, but the limited genetic diversity of many pathogens
- often hinders our understanding of transmission events (*Campbell et al., 2018*). As a consequence
- of the limited diversity, many index case and contact pairs will share identical genotypes, making

<sup>39</sup> it difficult to ascertain who infected whom.

Within-sample genetic diversity is common among a wide variety of pathogens (Mongkolrattan-40 othai et al., 2011: Lieberman et al., 2016: Dinis et al., 2016: Leitner, 2019: Popa et al., 2020), This 41 diversity may be generated *de novo* during infection, by a single transmission event of a diverse 42 inoculum or by independent transmission events from multiple sources (Worby et al. 2014) The 43 maintenance and dynamic of within-host diversity is then a product of natural selection, genetic drift and fluctuating population size (*Didelot et al. 2012*) The transmission of within-host varia-45 tion between individuals is also favored as a large inoculum exposure is more likely to give rise to 46 infection (Murphy et al., 1984: Han et al., 2019: Lee et al., 2022: Sender et al., 2021: Spinelli et al., 47 2021: Trunfig et al., 2021). The amount of within-sample diversity transmitted from index-case to 48 contact is determined by the bottleneck size (Zwart and Elena, 2015), with stringent bottlenecks 49 limiting the number of genotypes transmitted from the host to the recipient, and wide bottlenecks 50 allowing for the transmission of higher levels of genetic diversity (Worby et al., 2014). 51 Phylogenetic analysis provide information regarding the structure of the genetic diversity among 52 pathogen isolates. Moreover, pathogen phylogenetic trees can be used as input for many down-53 stream analysis, including inference of transmission events, population size dynamics or estima-54 tion of parameters of epidemiological models (Didelot et al., 2018). Most genomic and phyloge-55 netic workflows involve either genome assembly or alignment of sequencing reads to a reference 56 genome. In both cases, conventionally the resulting alignment exclusively represents the most 57 common nucleotide at each position. This is often referred to as the consensus sequence. Al-58 though genome assemblers may output contigs (combined overlapping reads) representing low 59 frequency haplotypes, only the majority contigits kept in the final sequence. In a mapping approach, 60 a frequency threshold for the major variant is usually pre-determined, under which a position is 61 considered ambiguous. The lack of genetic variation between temporally proximate samples and 62 the slow mutation rate of many pathogens results in direct transmission events sharing exact se-63 guences between the hosts when using the consensus sequence approach. For instance, the sub-64 stitution rate of SARS-CoV-2 has been inferred to be around 2 mutations per genome per month 65 (Harvey et al., 2021). Given its infectious period of 6 days (Byrne et al., 2020), most consensus 66 sequences in a small-scale outbreak will show no variation between them. This lack of resolution 67 and poor phylogenetic signal complicate phylogenetic inference, limiting the downstream analysis and conclusions that can be extracted from the phylogenetic tree. Previous work has shown the 60 advantages of using within-host diversity to infer transmission events compared to using consen-70 sus sequences (Wymant et al., 2018: De Majo et al., 2018). Aside from transmission inference, the 71 use of the within-host pathogen genetic data directly within phylogenetic inference will improve 72 any downstream analysis using a phylogenetic tree as starting point. 73 We hypothesize that the failure of consensus sequence approaches to capture within-sample 74 variation arbitrarily excludes meaningful data and limits pathogen phylogenetic and transmission 75 inference, and that including within-sample diversity in a phylogenetic model would significantly 76 increase the evolutionary and temporal signal and thereby improve our ability to infer infectious 77 disease phylogenies and transmission events. 78 We tested our hypothesis on multi-institutional SARS-CoV-2 outbreaks across London hospitals 79 that were part of the COVID-19 Genomics UK (COG-UK) consortia (COVID-19 Genomics UK (COG-UK)) 80 2020). Technical replicates, repeated longitudinal sampling from the same patient, and epidemio-81 logical data allowed us to evaluate the presence and stability of within-sample diversity within the 82 host and in independently determined transmission chains. We also evaluated the use of within-83 sample diversity in phylogenetic analysis by conducting outbreak and phylogenetic simulations 84 of sequencing data using a phylogenetic model that accounts for the presence and transmission

of sequencing data using a phylogenetic model that accounts for the presence and transmission
 of within-sample variation. We show the effects on phylogenetic inference of using consensus se-

<sup>87</sup> quences in the presence of within-sample diversity, and propose that existing phylogenetic models

can leverage the additional diversity given by the within-sample variation and reconstruct the phy-

logenetic relationship between isolates. Lastly, we show that by taking into account within-sample

- <sup>90</sup> diversity in a phylogenetic model we improve the temporal signal in SARS-CoV-2 outbreak analysis.
- <sup>91</sup> Using both phylogenetic outbreak reconstruction and simulation we show that our approach is
- superior to the current gold standard whole-genome consensus sequence methods.

#### 93 Results

# Sampling, demographics and metadata

Between March 2020 and November 2020, 451 healthcare workers, patients and patient contacts 95 at the participating North London Hospitals were diagnosed at the Camelia Botnar Laboratories 96 with SARS-CoV-2 by PCR as part of a routine staff diagnostic service at Great Ormond Street Hospi-97 tal NHS Foundation Trust (GOSH). A total of 289 isolates were whole-genome sequenced using the 98 Illumina NextSeg platform, which resulted in 522 whole-genome sequences including longitudinal and technical replicates (Supplementary file 1). The mean participant age was 40 years old (median 100 38.5 years old, interguartile range (IOR) 30-50 years old), and 60% of the participants were female 101 (Supplementary file 2). All samples were SARS-CoV-2 positive with real time oPCR cycle threshold 102 (C) values ranging from 16 to 35 cycles (Supplementary file 2). The earliest sample was collected 103 on 26th March 2020, while the latest one dated to 4th November 2020 (Figure 1-figure Supplement 1a) A total of 291 samples had self-reported symptom onset data for which the mean time 105

<sup>106</sup> from symptom onset to sample collection date was 5 days (IQR 2-7 days, *Figure 1—figure Supple-*

*ment 1*b). More than 90% of the samples were taken from hospital staff, while the rest comprised patients and contacts of either the patients or the staff members (Supplementarv file 2).

# <sup>109</sup> Genomic analysis of SARS-CoV-2 sequences

A total of 454 whole-genomes with mean coverage higher than 10x were kept for further analysis. 110 resulting in an average coverage across isolates of 2457x (Figure 1—figure Supplement 2). Allele 111 frequencies were extracted using the pileup functionality within *bcftools* (*Danecek et al.*, 2011) with 112 a minimum base and mapping quality of 30, which represents a base call error rate of 0.1%. Vari-113 ants at low frequency at positions where the mapped reads support more than one allele were 114 coined as minor or low-frequency variants. Variants were filtered further for read position bias 115 and strand bias. Only minor variants with an allele frequency of at least 2% were kept as puta-116 tive variants. Samples with a frequency of missing bases higher than 10% were excluded, keeping 117 350 isolates for analysis. The mean number of low frequency variants was 12 (median 3, IOR 1.00 – 118 9.75), although both the number of variants and its deviation increased at high C, values (*Figure 1*— 119 figure Supplement 3). 120

## <sup>121</sup> Within-sample variation in technical replicates

To understand the stability of within-sample variation and minimize spurious variant calls, we sequenced and analyzed technical replicates of 17 samples. Overall, when the variant was present in both duplicates the correlation of the variant frequencies was high ( $R^2 = 0.9$ , *Figure 1*a right). The high correlation was also maintained at low variant frequencies (*Figure 1*a left).

Minor variants were less likely to be shared when one or more of the paired samples had low viral load. These discrepancies may appear because of amplification bias caused by low genetic 127 material, base calling errors due to low coverage, or low base quality. The mean proportion of 128 discrepant within-sample variants between duplicated samples was 0.39 (sd = 0.29) although this 129 varied between duplicates (Figure 1—figure Supplement 4). C. values in RT-PCR obtained during 130 viral amplification are inversely correlated with viral load (*Tom and Ming, 2020*). The proportion of 131 shared intra-host variants was negatively correlated with C, values in a logistic model (estimate=-132 0.78, p-value=0.008), with higher C, values associated with a lower amount of shared intra-host 133 variants (*Figure 1*c). The number of within-sample variants detected also increased with C value. 134 as well as the deviation in the number of variants between duplicates (*Figure 1*d). This could be 136 explained either by an increase in the number of spurious variants at low viral loads (Tonkin-Hill 136



**Figure 1. Genomic analysis of technical duplicates before filtering. a** Allele frequency comparison between technical replicates for all frequencies (right) and for frequencies up to 1% (left). Colors represent the C<sub>t</sub> value for the sample. **b** Proportion of shared minor variants between technical replicates in relation to the C<sub>t</sub> value. **c** Total number of minor variants in relation to the C<sub>t</sub> value. Lines linked two technical replicates. Each sequence has a different color, with sequences from the same patient having a different shade of the same color.

**Figure 1—figure supplement 1.** Collection date distribution and time from symptom and days from symptom onset

Figure 1—figure supplement 2. Sample mean coverage distribution

Figure 1—figure supplement 3. Effects of C, value on whole-genome sequencing data

**Figure 1—figure supplement 4.** Proportion of shared minor variants between technical replicates using different filters of allele frequency

- 137 et al., 2021), biased amplification of low level sub-populations minor rare alleles (McCrone and
- Lauring, 2016), or due to the accumulation of within-host variation through time, as late stages of
- <sup>139</sup> infection are usually characterized by high C, values (low viral load).
- Based on these results, only samples with a  $C_r$  value equal or lower than 30 cycles were consid-
- ered, which resulted in 249 samples kept for analysis. Additionally, only variants with a frequency
- higher or equal than 2% were used. For the filtered dataset, 414 out of 29903 positions were poly-
- morphic for the consensus sequence, while the alignment with within-sample diversity had 1039
   SNPs. Of these, 699 positions had intra-host diversity, of which 78% (549/699) were singletons. The
- <sup>144</sup> SNPs. Of these, 699 positions had intra-host diversity, of which 78% (549/699) were singletons. The <sup>145</sup> majority of samples (207/249, 83%) contained at least 1 position with a high quality within-host vari-
- majority of samples (207/249, 83%) contained at least 1 position with a high quality within-host va
   ant. and the median amount of intra-host variants per sample was 2 (IOR 1-4.5).

#### <sup>147</sup> Within-sample variation in epidemiologically linked samples

Given the limited genomic information in the consensus sequences, epidemiological data is often 148 necessary to infer the directionality of transmission. We categorized our samples within the fol-149 lowing groups: samples that a) did not have any recorded epidemiological link, b) were from the 150 same hospital (possibly linked), c) samples that were part of the same department within the same 151 hospital (probable link), d) samples that had an epidemiological link within the same department 152 of the same hospital (proven link), e) were a longitudinal replicate from the same patient and f) a 153 technical replicate from the same sample. 154 We tested the concordance between epidemiological and genomic data by determining the SNP 155

We tested the concordance between epidemiological and genomic data by determining the SNP distance between pairs of samples with epidemiological links and without them. Pairs of samples from the same hospital, department, epidemiologically linked, or longitudinal and technical replicates had a lower SNP distance (were more closely related) than those samples that did not have any relationship, although this difference was small in the case of pairs of samples from the same hospital (*Table 1*).

To understand the distribution of shared low frequency variants among different groups of samples, we performed a pairwise comparison of all samples and calculated the proportion of

#### Table 1. SNP distance between pairs of samples.

Sample relationship	Estimate (95%Cl)	p-value
None	11.04 (10.94 - 11.15)	Reference
Hospital	9.78 (9.48 - 10.09)	$\leq 1 \times 10^{-4}$
Department	5.15 (4.54 - 5.83)	$\leq 1 \times 10^{-4}$
Epidemiological	1.5 (1.22 - 1.78)	$\leq 1 \times 10^{-4}$
Longitudinal duplicates	0 (0 - 0.2)	$\leq 1 \times 10^{-4}$
Technical replicate	0 (0 - 0.2)	$< 1 \times 10^{-4}$

shared within-sample variants (shared variants divided by total variants in the pair) within groups 163 with epidemiological links and without them. The proportion of shared within-host variants was 164 higher between technical replicates, longitudinal duplicates, epidemiologically linked samples, and 165 samples taken from individuals from the same department when compared to pairs with no epi-166 demiological links, although the range of this probability was large (Figure 2. Figure 2—figure Sup-167 *plement 1*). The probability of sharing a low frequency variant was inferred using a logistic regres-168 sion model (Figure 2—figure Supplement 2). There was a tendency for the probability to increase 169 with variant frequency, but the association was not strong (Odds ratio 1.8, 95% CI 0.9 - 3.5, p=0.08). 170 The probability of sharing a low frequency variant for samples with no epidemiological links was 171  $9.5 \times 10^{-6}$  (95% Cl 8.8  $\times 10^{-6}$  –  $1.02 \times 10^{-5}$ ). Samples from the same hospital did not have a probability 172 significantly higher than those without any link  $(3.3 \times 10^{-3}, 95\% \text{ Cl} 2.7 \times 10^{-3} - 4.03 \times 10^{-3})$ . On the 173 other hand, pairs from the same department, with epidemiological links, replicates or technical 174 replicates all had a significantly higher probability of sharing a low frequency variant when com-175 pared to those pairs with no link (all Wald test p-values < 0.001). The inferred probabilities for pairs 176 from the sample department was 1.4% (95% CI 0.9% – 2.1%), which increased to 5% for pairs with 177 epidemiological links (95% Cl 4.2% - 6.4%). For longitudinal replicates, the probability was inferred 178 to be 38% (95% CI 35% - 41%), and were shared between multiple time points (Figure 2—figure 179 Supplement 3). Technical replicates were estimated to have the highest probability (70%, 95% CI 180 64% - 76%). 181



**Figure 2. Probability of sharing within-host variants in sample pairs.** The probability of variants shared between pairs of samples calculated as the number of low frequency variants in both samples divided by the total number of variants between the pair. Colors grouped samples by their relationship. Points represent the mean probability a variant is shared between all pairwise samples within a group and allele frequency. Error bars show the 95<sup>th</sup> and 5<sup>th</sup> percentiles.

Figure 2—figure supplement 1. Allele frequency comparison in pairwise sample pairs.

Figure 2—figure supplement 2. Probability that minor variants are shared.

Figure 2—figure supplement 3. Dynamics of low frequency variants in longitudinal duplicates.

- <sup>182</sup> Within-host diversity model outperforms the consensus model in simulations
- <sup>183</sup> The effect of within-sample diversity in phylogenetic inference was tested by evaluating the accu-
- racy in the reconstruction of known phylogenetic trees using a conventional phylogenetic model
- and a model that accounts for within-sample variation.
- The presence of within-sample diversity was coded in the genome alignment using existing
- 187 IUPAC nomenclature (IUPAC-IUB Joint Commission on Biochemical Nomenclature (JCBN), 1984).
- For the consensus sequence alignment, only the 4 canonical nucleotides were used (*Figure 3*a,b),
- while the proposed alignment retained the major and minor allele information as independent
- character states (*Figure 3*c,d).



#### Figure 3. Model of within-host diversity.

Proposed evolutionary model of within-host diversity in genomic sequences. Uppercase letters represent the major variant in the population, while lowercase letters indicate presence of a minor variant alongside the major one. **a**, **c** Genome sequences where some positions show within-sample variation (top), represented by a major allele (big size letter) and a minor one (smaller size), as well as its representation in the alignment (bottom). **b**, **d** Models of nucleotide evolution. Character transitions are indicated by arrows. **a** Consensus sequence, where only the major allele is represented in the alignment. **b** Model of nucleotide evolution using the consensus sequence, with four character states representing the four nucleotides. **c** Sequence with within-sample variation, represented by an uppercase letter for the major allele and a lower case letter for the minor allele. **d** Model of nucleotide evolution with 16 character states accounting for within-sample variation.

In order to evaluate the differences in tree inference with and without the inclusion of within-191 sample diversity, we simulated genome alignments for 100 random trees using a phylogenetic 192 model where both major and minor variant combinations were considered, resulting in a total of 193 16 possible states (*Figure 3*d). In the proposed model, transitions and transversions between the 19 four nucleotides in the population occur in the following steps: first a minority variant evolves at 195 low frequency, then the minor variant increases its frequency to become the majority nucleotide, and finally the variant is fixed (Figure 3d), with all the steps being reversible. Therefore, within-197 host evolutionary dynamics are modelled by explicitly considering base change as a process of 198 minor variant evolution and eventual fixation. The substitution rates chosen for the simulations. 190 as shown in Supplementary file 4, were selected to reflect a slow rate of minor variant evolution 200 and a fast rate at which minor variants are lost or fixated in the population, which in turns results 201 in a highly dynamic landscape of within-sample variation, with the four canonical nucleotides 100 202

<sup>203</sup> times more likely to be present than low frequency variants.

From the simulated genomes, two types of alignments were generated: a consensus sequence, where only the major allele was considered (*Figure 3*a); and an alignment that retained the major and minor allele information as independent character states (*Figure 3*c). From the simulated alignments, RaxML-NG was used to infer phylogenetic trees (*Kozlov et al., 2019*). The consensus sequence was analyzed with a GTR+ $\gamma$  model, while the PROTGTR+ $\gamma$  model was used in order to accommodate the extra characters of the alignment with within-sample diversity and major/minor variant information.

The two models were evaluated for their ability to infer the known phylogeny that included 211 within-host diversity. The estimated phylogenies were compared to the known tree using differ-212 ent measures to capture dissimilarities in a variety of aspects relevant to tree inference (Supple-213 mentary file 3). For all the metrics employed, the phylogenies inferred explicitly using within-host 21/ diversity as independent characters approximated better to the initial tree than the one using the 215 consensus sequence (Figure 4). Additionally, the transition/transversion rates inferred by the phy-216 logenetic models accounting for within-host diversity accurately reflect the rates used for the sim-217 ulation of genomic sequences (Supplementary file 4, 5 and 6). 218



#### Figure 4. Similarity scores for inferred trees.

Comparison of the phylogenetic trees inferred using simulated sequences from known random starting trees and different phylogenetic models. Colors differentiate the metrics used for the comparison.

Figure 4—figure supplement 1. Similarity scores for inferred trees with different rates.

Figure 4—figure supplement 2. Similarity scores for inferred trees from coalescent simulations.

As different pathogens are likely to show different dynamics of within-host variation and the 219 rates used for the simulations will inevitably affect the improvement of using the 16-state model. 220 we simulated genomes with different parameters. As expected, choosing rates that promote an 221 abundant and stable landscape of low frequency variation (rate of minor variant acquisition of 222 20, and rates of variant switch and lost of 1) made the 16-state model to perform better than the 223 model using consensus sequences, which improved as the proportion of low frequency variants 224 decreased (Figure 4—figure Supplement 1). Conversely, in simulations using a lukes-Cantor DNA model, and therefore without any low frequency variation, both models showed similar results 226 (Figure 4—figure Supplement 1). 227

To understand the effects of genetic linkage between sites in the phylogenetic model due to the clonal relationships between genomes, we evaluated another set of simulations where the starting tree was generated using the coalescent model, which increases the correlation between sites. For all metrics used, the model using low frequency variants inferred phylogenies more similar to the starting coalescent tree than those inferred using the consensus sequence (*Figure 4—figure Supplement 2*).

We further assessed the effect of within-host diversity in phylogenetic inference by simulating
 pathogen evolution throughout the time frame of infectious disease outbreaks (*De Maio et al.,* 2018). We simulated outbreaks using TransPhylo (*Didelot et al., 2017*) with a host population vary-

ing between 10 and 15 hosts, no recombination, complete sampling of the outbreak and selecting 237 epidemiological parameters to match the transmission dynamics of SARS-CoV-2. For each out-238 break, we simulated the evolution and transmission of the pathogen population within each host 239 with varving values of mutation rates and transmission bottlenecks using fastsimcoal2 (Excoffier 240 et al., 2013) as previously described by De Majo et al. (2018). We compared the resulting phylo-241 genetic trees to the real outbreak phylogeny using the Kuhner-Felsenstein distance (Kuhner and 242 Felsenstein, 1994). Even though using consensus sequences performed better than a random 243 distribution of trees, using within-host diversity outperformed the consensus sequence in all in-244 stances (Figure 5). The phylogenies inferred using within-host diversity were more similar to the 245 real outbreak phylogeny for wider bottleneck sizes, with the best performance when no bottleneck 246 was present. As expected, both the consensus sequences and the sequences reflecting within-247 sample diversity were more informative at higher mutation rates, even though the consensus se-248 guence only showed improvement with a mutation rate of  $10^{-3}$  mutations per base per generation 240 cycle (Figure 5). 250





#### <sup>251</sup> Within-host diversity improves the resolution in SARS-CoV-2 phylogenetics

Genome sequences collected at different time points are expected to diverge as time progresses. 252 resulting in a positive correlation between the isolation date and the number of accumulated mu-253 tations (temporal signal) (*Rieux and Balloux, 2016*). The alignment with consensus sequences and 254 the one reflecting within-sample variation were used to infer two different phylogenetic trees (Fig. ure 6—figure Supplement 1). Longitudinal samples in the phylogeny inferred using within-host 256 diversity reflected the expected temporal signal, with an increase in genetic distance as time pro-257 gressed between the longitudinal pairs in a linear model (coefficient 2.24, 0.59 - 3.88 95% CL p = 258 0.019. Figure 6—figure Supplement 2). The difference in C, value among longitudinal duplicates 259 was not correlated with higher genetic distances (coefficient 1.62, -0.66 - 3.91, 95% CL p = 0.2). 260 Similarly, we analyzed the number of low frequency variants within outbreaks by counting the 261 number of within-sample variants for each isolate belonging to a specific outbreak and inferred 262 their change through time taking the earliest isolate date as the starting point of the outbreak. In 263 general, as the outbreaks progressed the number of low frequency variants increased (coefficient 264 0.16, 0.06 - 0.27 95% Cl, p = 0.003,  $r^2$  = 0.19, Figure 6—figure Supplement 3). 265 We analyzed the impact of using within-sample variation on the temporal structure of the phy-266

logeny by systematically identifying clusters of tips in the phylogenetic tree with an identical con sensus sequence and no temporal signal. We then performed a root-to-tip analysis using the tree

inferred with intra-sample diversity. Only clusters with more than 3 tips were used for the root-to-

tip analysis. The majority of clusters (10/11) showed a positive correlation between the distance of

the tips to the root and the collection dates, demonstrating a significant temporal signal between

samples when there was none using the conventional consensus tree (*Figure 6*).



# Figure 6. Previously uninformative clusters present temporal signal when using within-sample diversity.

A set of 11 outbreak clusters (one per panel, each plotting the root to tip distance in number of substitutions per genome against time) in which all samples had identical consensus genomes sequences (and therefore no temporal signal). Blue colors indicate those regressions that after utilizing within sample diversity now have a positive slope (temporal signal), and red shows those regressions that have a negative slope (misleading or false positive temporal signal).

Figure 6—figure supplement 1. Phylogenetic trees for SARS-CoV-2.

Figure 6—figure supplement 2. Genetic distance between longitudinal samples.

**Figure 6—figure supplement 3.** Number of low frequency variants within outbreaks as the outbreak progresses.

To illustrate the downstream application of the improved phylogenetic resolution, we inferred 273 a time-calibrated phylogeny from the phylogeny inferred using the 16-character state model with 274 the collection dates of the tips using BactDating (Didelot et al., 2018) (Figure 7-figure Supple-275 ment 1) and calculated the likelihood of transmission events within potential epidemiologically 276 identified outbreaks using a Susceptible-Exposed-Infectious-Removed (SEIR) model (Lekone and 277 Finkenstädt, 2006; Eldholm et al., 2016). The SEIR model was parameterized with an average la-278 tency period of 5.5 days (Xin et al., 2022), an infectious period of 6 days (Byrne et al., 2020), and 279 a within-host coalescent rate of 5 days as previously estimated for SARS-CoV-2 (Wang et al., 2020). 280 The likelihood of transmission was calculated for every pair of samples, while the Edmonds algo-281 rithm as implemented in the R package RBGL (Carey et al., 2021) was used to infer the graph with 282

the optimum branching (Figure 7c,d; Figure 7—figure Supplement 2).



**Figure 7. Within-sample variation improves resolution of infectious disease outbreaks.** Effect of using low frequency variants in phylogenetic inference. **a** Maximum likelihood phylogeny using the consensus sequences (left) and the alignment leveraging within-sample variation. Replicates of the same sample share the same color. Sample IDs are coded as follows: SF, for staff members; P, for patients; and PC, for patient contacts. **b** Transmission network inferred using within sample variation. Edge width is proportional to the likelihood of direct transmission using a Susceptible-Exposed-Infectious-Removed (SEIR) model. Colored edges represent the Edmunds optimum branching and thus the most likely chain. **c** Heatmap of the likelihood of direct transmission between all pairwise pairs of samples using a SEIR model. Vertical axis is the infector while the horizontal axis shows the infectee.

Figure 7—figure supplement 1. Time calibrated phylogenetic trees for SARS-CoV-2. Figure 7—figure supplement 2. Phylogenetic and transmission for SARS-CoV-2 outbreaks.

Figure 7 represents an example of an outbreak involving 4 hosts, with one patient, one patient 284 contact, and two hospital staff members. All samples have one technical replicate, while patient 285 sample also has two serial samples (which were removed for transmission inference). The ML tree 286 inferred using the consensus sequences (Figure 7a, left) shows that most isolates have the exact 287 same consensus sequence. Although this suggests that all isolates belong to the same outbreak. 288 the similarity between sequences precludes exact transmission inference. However, the ML tree 289 inferred using sequences with low frequency variants correctly clusters technical and longitudinal 290 replicates, and groups the isolates in distinctive sets that better inform transmission inference (Fig-291 ure 7b,c). We applied the same analysis to other potential outbreaks and obtained similar results 292 (Figure 7—figure Supplement 2). 293

#### 294 Discussion

Detailed investigation of transmission events in an infectious disease outbreak is a prerequisite for 295 effective prevention and control. Although whole-genome sequencing has transformed the field of 296 pathogen genomics, insufficient pathogen genetic diversity between cases in an outbreak limits the 297 ability to infer who infected whom. Using multi-hospital SARS-CoV-2 outbreaks and phylogenetic 298 simulations, we show that including the genetic diversity of subpopulations within a clinical sample 299 improves phylogenetic reconstruction of SARS-CoV-2 outbreaks and determines the direction of 300 transmission when using a consensus sequence approach fails to do so. 301 The majority of samples sequenced harbored variants at low frequency. However, most vari-302 ants were not consistently called in technical replicates, suggesting they were spurious or unreli-303 able. Within-sample variation was less consistent between paired technical replicates with lower 304 viral load (higher C,). This is likely to be a consequence of low starting genetic material giving rise 305 to amplification bias during library preparation and sequencing. Establishing a cut-off for high C. 306 values is therefore important to accurately characterize within-host variation. In our study, we ex-307 cluded samples with a C. value higher than 30 cycles based on the diagnostic PCR used at GOSH. 308

Since C, values are only a surrogate for viral load and are not standardized across different assays 309 (Evans et al., 2021), appropriate thresholds would need to be determined for other primary PCR 310 testing assays. Similarly, variant calls at very low frequency were less likely to be present in both 311 technical replicates. These variants at low frequency are thus potentially not genuine and the re-312 sult of sequencing and variant calling errors. For our work, we removed any variants with an allele 313 frequency lower than 2%. Until sequencing and variant calling technologies improve for low fre-31 guency variants, technical replicates will remain essential for the study of pathogen within-host 31! diversity in order to distinguish genuine variation from sequencing noise. The effect of this noise 316 on phylogenetic inference will depend on the signal-to-noise ratio and the amount of variation 317 already present in the consensus sequences. Spurious low frequency variation will likely affect 318 only the branch length estimation in phylogenetic inference by adding potentially erroneous calls. 310 unless there is presence of batch bias which could artificially cluster epidemiologically unrelated 320 isolates together. 321

The generation, maintenance and evolution of subpopulations within the host reflect evolu-322 tionary processes which are meaningful from phylogenetic and epidemiological perspectives. Sub-323 populations within a host can emerge from three mechanisms; de novo diversification in the host. 324 transmission of a diverse inoculum, or multiple transmission events from different sources. If the 325 subpopulations are the result of de novo mutations, nucleotide polymorphisms within the subpop-326 ulations accumulate over time and may therefore result in a phylogenetic signal useful for phyloge-327 netic inference. In our data, longitudinal samples taken at later time points were demonstrated to 328 accrue genomic variation. Although this pattern can be confounded by decreasing viral load as in-329 fection progresses, C. values in our dataset were not correlated with a higher genetic distance, and 330 clusters in our data containing both longitudinal and technical replicates also corroborate these 331 results. Transmission of a diverse inoculum also gives rise to phylogenetically informative shared 332 low frequency variants, as our results show that transmission pairs are more likely to share vari-333 ants at low frequency. The effect of multiple transmission events in the phylogeny depends on the 334 relatedness of both index cases and the bottleneck size in each transmission event 335

Paired samples with epidemiological links and from the same department shared a higher pro-336 portion of low frequency variants and were located closer in the consensus tree than samples with 337 no relationship. These patterns suggest that the distribution of low frequency variants is linked to 338 events of epidemiological interest. The fact that technical duplicates shared more within-host di-339 versity than longitudinal replicates of the same sample suggests that much of the variation within 340 hosts is transitory. Therefore, within-host diversity may be relevant on relatively short time scales. 341 which is precisely where consensus sequences lack resolution. Combining the data derived from 342 fixed alleles in the consensus sequences and transient within-sample minor variation enables an 343 improved understanding of the relatedness of pathogen populations between hosts. 344

The effects of neglecting within-host diversity in phylogenetic inference were analyzed by us-345 ing simulated sequences under a phylogenetic model that reflects the presence and evolution of 346 within-host diversity. We compared a conventional consensus phylogenetic model and a model 347 that leverages within-sample diversity, and evaluated their ability to infer the known phylogeny. 348 Our proposed phylogenetic model incorporates within-sample variation by explicitly coding ma-3/10 ior and minor nucleotides as independent characters in the alignment. We demonstrated that 350 phylogenies inferred using the conventional consensus sequence approach were unresolved and 351 unrepresentative of the known structure of the simulated tree. However, sequences that included 352 within-host diversity showed higher resolution that resulted in phylogenetic trees more similar to 353 the simulated phylogeny. As other mutational models, our 16-state model assumes independence 354 between sites in the alignment. This assumption can be violated due to the presence of genetic link-355 age, which can be caused by multiple biological processes, such as clonal relationships between 356 microorganisms, recombination or selection of co-evolving sites. To increase the amount of ge-357 netic linkage due to clonal relationships between organisms, we repeated our simulations using 358 a coalescent model to create the starting tree, and confirmed that the 16-state model still outper-359

formed the conventional consensus sequence in the presence of high linkage. Other sources of

<sup>361</sup> genetic linkage are not accounted for, and their inclusion in phylogenetic inference is out the scope

of this work.

The proposed phylogenetic model used for the simulations did not include direct base transitions and transversions, but rather modelled base change as a process of minor variant acquisition and fixation. Therefore, a base change is composed of the following steps: first a minor variant is gained; then the minor variant increases in frequency and becomes the majority variant; and finally the new variant is fixed. In this way, within-host evolution is partially included in the model as a process of minor variant evolution and eventual lost or fixation. As shown in the simulations, this process of within-host evolution is also captured when the minor bases are simply incorporated as additional states in the Markov chain without explicitly limiting the possible transitions.

We complemented the phylogenetic simulations with tree inference of outbreaks simulated us-371 ing TransPhylo (*Didelot et al.*, 2017). We parameterized the simulations to reflect the transmission 372 dynamics of SARS-CoV-2, including a generation time of 5 days and a sampling time of 7 days. Given 373 this parameters, most simulated outbreaks lasted less than a month. We then simulated genetic 374 sequences within the outbreak using fastsimcoal2 (Excoffier et al., 2013) as previously described 375 by **De Majo et al.** (2018). Using a mutation rate of  $5 \times 10^{-6}$  mutations per base per replication cycle. 376 as previously described for SARS-CoV-2 and other betacoronaviruses (Sender et al., 2021; Amicone 377 et al., 2022), and varying bottleneck sizes, we showed that tree inference using within-sample diver-378 sity improves as the transmission bottleneck widens, although even at low bottleneck sizes trees 370 inferred using within-sample diversity are more accurate than those inferred using consensus se-380 quences. Similarly, using varying mutation rates and a constant bottleneck size of 10 pathogens. 381 we showed that tree inference was more accurate as mutation rates increased, although inference 382 using consensus sequences improved only at a very high mutation rate of  $10^{-3}$  mutations per base 383 per cycle, which has mostly been observed in some HIV studies (*Cuevas et al., 2015*). Together, 384 our simulations show that at the short time frame of disease transmission, phylogenetic inference 385 using alignments that contain information regarding within-sample diversity outperform phyloge-386 nies inferred with consensus sequences, even at parrow transmission bottlenecks and very low 387 mutation rates. Since TransPhylo simulates phylogenetic trees alongside the outbreak simulation. 388 we could directly compare our inferred phylogenies with the known simulated trees. However, al-389 though phylogenetic trees can inform transmission inference, phylogenetic trees themselves and 390 transmission trees are not interchangeable. Nevertheless, increasing the resolution of phyloge-391 netic trees can improve inference of transmission chains and calculation of the likelihood of trans-302 mission events 393

Previous studies have addressed the use of within-host variation to infer transmission events. 304 Wymant et al. (2018) employed a framework based on phylogenetic inference and ancestral state 395 reconstruction of each set of populations detected within read alignments using genomic windows. 306 Our study extends this work by coding genome-wide diversity within the host directly in the align-397 ment and the phylogenetic model. *De Majo et al.* (2018) proposed direct inference of transmission 308 from sequencing data alongside host exposure time and sampling date within the bayesian frame-399 work BEAST2 (*Bouckgert et al., 2014*). Our approach is focused on directly improving the temporal 400 and phylogenetic signal of whole-genome sequences, and it's especially suited for use in applica-401 tions and analysis that employ a phylogenetic tree as input to infer transmission (Didelot et al., 402 2017). 403 Apart from transmission inference, phylogenetic trees can be used to infer many parameters 404

Apart from transmission inference, phylogenetic trees can be used to infer many parameters of epidemiological interest, such as R<sub>0</sub> or the effective population size. In our work, we showed that the temporal signal of clusters where all isolates had the same sequences increased with the inclusion of within-sample diversity, which in turns allows better inference of phylogenetic trees. When analyzing specific outbreaks, we showed that groups of samples without genetic differences were clustered apart from other isolates of the outbreak, providing additional information on genetic relationships that could be used for transmission inference or to better understand the genetic

- structure of the outbreak. Even though transmission inference can be improved with epidemio-
- <sup>412</sup> logical data such as collection dates even when all isolates have the same genetic sequences, such

data can't provide information regarding how samples cluster within the outbreak. Additionally,

the order of collection dates not always correspond to the order of infection.

Future work will extend this model by including allele frequency data in addition to indepen-415 dent characters for major and minor variants. Moreover, to limit the number of character states 416 we only allowed two variants at each position. Transmission inference of pathogens with high lev-417 els of within-host diversity. for instance as observed in HIV. could benefit from including more than 418 two alleles. In those cases, the number of possible character state combinations would be too large. 410 and therefore other methods such as phyloscanner *Wymant et al.* (2018) could resolve transmis-420 sion events more accurately. However, it's important to note that the low frequency of a third allele 421 could result in more sequencing and mapping errors which could in turn bias phylogenetic infer-422 ence and genomic analysis. Phylogenetic models that explicitly include dynamics of within-sample 423 variation and sequencing error may further improve phylogenetic inference or allow researchers 424 to better estimate parameters of interest, including R0, bottleneck size, transmissibility and the 425 origin of outbreaks. 426

In line with conventional consensus sequencing approaches, we used a reference sequence for genome alignment and variant calling. Although widely used, one limitation of this approach is a potential mapping bias causing some reads to reflect the reference base at low frequencies at a position where only a variant should be present. Although we applied stringent quality filtering, we cannot rule out the persistence of some false positive minor variants. Using genome graphs to map to a reference that encompasses a wider spectrum of variation may alleviate this problem, and could be an interesting addition to pathogen population genomic analysis.

Our results demonstrate that within-sample variation can be leveraged to increase the reso-434 lution of phylogenetic trees and improve our understanding of who infected whom. Using SARS-435 CoV-2 hospital outbreaks and simulations, we show that variants at low frequencies are consistent 436 within sample replicates, phylogenetically informative and are more often shared among epidemi-437 ologically related contacts. By coding within-sample variation directly in the alignment the ad-438 ditional genetic information can be easily incorporated in phylogenetic inference, facilitating its 439 application within existing epidemiology pipelines and public health infrastructure. We propose 440 that pathogen phylogenetic models should accommodate within-host variation to improve the un-441 derstanding of infectious disease transmission and aid infection control measures. 442

## 443 Materials and Methods

# 444 Model for within-host diversity

To test the accuracy of different models at inferring known phylogenies, 100 random phylogenetic 445 trees with 100 tips each were generated using the function *rtree* within the R package *ape* (*Par*-446 adis et al 2004) Whole-genome alignments were simulated from the random 100 phylogenies 447 with the function SimSeg of the R package phangorn (R Core Team and R Foundation for Statistical Computing, 2021: Schliep, 2011) using a model with 16 character states that represent the combinations of the 4 nucleotides with each other as minor and major alleles (*Figure 3*d). Three 450 substitution rates for the model were considered: a rate at which minor variants evolve, equal to 1: 451 the rate at which minor variants are lost, leaving only the major nucleotide at that position, equal 452 to 100; and the rate at which minor/major variants are switched, equal to 200. This rates result in 463 fixed bases (A, C, G, and T) being 100 times more frequent than low frequency bases. A different 454 set of simulations was performed using rates that promote a high rate of low frequency variation 455 by having a lower rate of variant loss and switch (rates 1, 10, 10 for minor variant evolution, loss 456 and switch. respectively); a low amount of low frequency variation by increasing the rates of variant 457 switch and loss (1, 10, 100); and using a lukes-Cantor model of sequence evolution and therefore 450 resulting in no minor variants. 459

- Two types of alignments were generated from the simulated genomes: a consensus sequence, where only the major allele was considered; and an alignment that retained the major and minor allele information as independent character states. RaxML-NG (*Kozlov et al., 2019*) was used to infer phylogenetic trees. The consensus sequence was analyzed with a GTR+ $\gamma$  model, while the PROTGTR+ $\gamma$  model was used for the alignment with intra-host diversity and major/minor variant information. Several metrics were used to compare the 200 inferred phylogenetic trees with their respective
- starting phylogeny from which the sequences were simulated (Supplementary file 3). We chose 467 metrics available in R suitable for unrooted trees, using the option 'rooted=FALSE' where appro-468 priate. The Robinson-Foulds (RF) distance (Robinson and Foulds, 1981) calculates the number of 460 splits differing between both phylogenetic trees. For the weighted Robinson-Foulds (wRF), the dis-470 tance is expressed in terms of the branch lengths of the differing splits. The Kuhner-Felsenstein 471 distance (Kuhner and Felsenstein, 1994) considers the edge length differences in all splits, regard-472 less of whether the topology is shared or not. Last, the Penny-Steel distance or path difference 473 metric (Steel and Penny, 1993) calculates the pairwise differences in the path of each pair of tips. 474 with the weighted Penny-Steel distance (wPS) using branch length to compute the path differences. 475
- All functions were used as implemented in the package phangorn (Schliep, 2011) within R (R Core
- Team and R Foundation for Statistical Computing, 2021).

# 478 Outbreak simulations

- Disease outbreaks of size between 10 and 15 hosts were simulated using TransPhylo (Didelot et al., 470 2017), with a mean generation time of 5 days and a mean sampling time of 7 days, both parameters 480 with standard deviation of 1 day (Wang et al., 2020; Hart et al., 2022). To ensure that the outbreak 481 ends, the negative binomial distribution for the offspring number was set with a mean of 1 and a 482 dispersion parameter of 0.5, resulting in a basic reproductive number ( $R_0$ ) of 1. To simplify the sim-483 ulations, all hosts from the outbreak were sampled. A total of 20 outbreaks were simulated. The 484 population evolution within and between hosts was simulated using fastsimcoal2 (Excoffier et al., 485 2013) as previously described by De Maio et al (De Maio et al., 2018), where transmissions are incor-486 porated as population migrations with a given bottleneck size and populations evolve with a given 487 mutation rate per generation time. Sequences were simulated for a within-host population size of 1000 and a genome size of 1000bp. To understand the effect of transmission bottleneck size in 489
- 480 phylogenetic inference, varving values of bottleneck size were used along a constant mutation rate
- $_{491}$  of  $5 \times 10^{-6}$  mutations per base per generation cycle. Additionally, sequences were simulated at dif-
- 492 ferent mutation rates with a constant bottleneck size of 10 pathogens. Sequences with the varying
- <sup>493</sup> bottleneck sizes and mutation rates were simulated using the same 20 simulated outbreaks. Phy-
- logenetic trees were inferred from the alignments using RaxML-NG as previously described. The
- resulting trees were time-calibrated using the additive uncorrelated relaxed clock model (ARC) as implemented in BactDating (*Didelot et al., 2018*). The root of the outbreak was inferred as part of
- <sup>496</sup> implemented in BactDating (*Didelot et al., 2018*). The root of the outbreak was inferred as part of <sup>497</sup> the dating model. The inferred trees were compared to the known simulated phylogenies using
- the dating model. The inferred trees were compared to the known simulated phylogen
- <sup>498</sup> the Kuhner-Felsenstein distance (*Kuhner and Felsenstein, 1994*).

# 499 Amplification and whole-genome sequencing

SARS-CoV-2 real-time gPCR confirmed isolates from London hospitals were collected as part of the 500 routine diagnostic service at Great Ormond Street Hospital NHS Foundation Trust (GOSH) (Storey 501 et al., 2021) and the COVID-19 Genomics UK Consortium (COG-UK) (COVID-19 Genomics UK (COG-502 (IK) 2020) between March and December 2020, in addition to epidemiological and patient meta-603 data (Supplementary file 2). Multiple types of samples were collected: isolates from different pa-504 tients: longitudinal replicates, where multiple isolates were collected from the same patient at 505 different time points; and technical replicates, where multiple sequencing runs were performed 506 from the same biological isolate. SARS-CoV-2 whole-genome sequencing was performed by UCL 507 Genomics, cDNA and multiplex PCR reactions were prepared following the ARTIC nCoV-2019 se-508

quencing protocol (Tyson et al., 2020). The ARTIC V3 primer scheme (ARTIC Network, 2021) was 509 used for the multiplex PCR, with a 65°C, 5 min annealing/extension temperature. Pools 1 and 2 mul-510 tiplex PCRs were run for 35 cycles. 5uL of each PCR were combined and 20uL nuclease-free water 511 added. Libraries were prepared on the Agilent Bravo NGS workstation option Busing Illumina DNA 512 prep (Cat. 20018705) with unique dual indexes (Cat. 20027213/14/15/16). Equal volumes of the 513 final libraries were pooled, bead purified and sequenced on the Illumina NextSeq 500 platform 514 using a Mid Output 150 cycle flowcell (Cat. 20024904) (2 x 75bp paired ends) at a final loading 515 concentration of 1.1pM. 516

## <sup>517</sup> Whole-genome sequence analysis of SARS-CoV-2 sequences

Raw illumina reads were quality trimmed using Trimmomatic (*Bolger et al., 2014*) with a minimum
 mean quality per base of 20 in a 4-base wide sliding window. The 5 leading and trailing bases of
 each read were removed, and reads with an average quality lower than 20 were discarded. The
 resulting reads were aligned against the Wuhan-Hu-1 reference genome (GenBank NC\_45512.2,
 GISAID EPI\_ISL\_402125) using BWA-mem v0.7.17 with default parameters (*Li and Durbin, 2010*).
 The alignments were subsequently sorted by position using SAMtools v1.14 (*Li et al., 2009*). Primer
 sequences were masked using ivar (*Grubaugh et al., 2019*).

Single-nucleotide variants were identified using the pileup functionality of samtools (Li et al. 525 2009) via the pysam package in Python (https://github.com/pysam-developers/pysam). Variants 526 were further filtered using bcftools (*Danecek et al., 2011*). Only variants with a minimum depth 527 of 50x and a minimum base quality and mapping quality of 30 were kept. Additionally, variants 628 within low complexity regions identified by sdust (https://github.com/lh3/sdust) were removed. 520 Previously identified problematic sites were masked to avoid systematic sequencing errors and 530 phylogenetic bias (*De Majo et al., 2020*). For positions where only one base was present, the min-531 imum depth was 20 reads, with at least 5 reads in each direction. Positions with low frequency 532 variants were filtered if the total coverage at that position was less than 100x, with at least 20 533 reads in total and 5 reads in each strand supporting each of the main two alleles. 534

Two different alignments were prepared from the data. First, an alignment of the consensus sequence where the most prevalent base at each position was kept. Variants where the most prevalent allele was not supported by more than 60% of the reads were considered ambiguous. Additionally, an alignment reflecting within-sample variation at each position as well as which base is the most prevalent and which one appears at a lower frequency by using the IUPAC nomenclature for amino acids (*IUPAC-IUB Joint Commission on Biochemical Nomenclature (ICBN), 1984*).

For the two different alignments, maximum likelihood phylogenies were inferred by using RAXML 541 NG (Kozlov et al., 2019) with 20 starting trees (10 random and 10 parsimony), 100 bootstrap repli-542 cates, and a minimum branch length of  $10^{-9}$ . For the consensus sequence, the GTR model was 543 used. For the alignment reflecting within-host diversity, a model with amino acid nomenclature 544 (PROTGTR) was used. All models allowed for a  $\gamma$  distributed rate of variation among sites. Phy-545 logenetic trees were time-calibrated using the known collection dates and the ARC model within BactDating (*Didelot et al., 2018*). For transmission inference, the dated phylogeny was used with 547 the longitudinal replicates removed by keeping the earliest sampled isolate. The likelihood of trans-548 mission was calculated using a Susceptible-Exposed-Infectious-Removed (SEIR) model (Lekone and E 4 0

550 Finkenstädt. 2006: Eldholm et al.. 2016).

#### **Data availability**

552 Samples sequenced as part of this study have been submitted to the European Nucleotide Archive

<sup>553</sup> under accession PRJEB53224. Sample metadata is included in Supplementary file 1.

### **554** Code availability

- All custom code used in this article can be accessed at
- bttps://github.com/arturotorreso/scov2\_withinHost.git.

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#### **570** Competing interests

- The authors declare no competing interests
- **572** Ethics declarations
- <sup>573</sup> Ethical approval was obtained for all individual studies from which this data was derived.
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# 773 Appendix Figures and Tables

supplementary file 1. Study participants metadata.

**Supplementary file 2.** Sample collection and demographics.

Supplementary file 3. Metrics used for phylogenetic tree comparison.

**supplementary file 4.** Transition/transversion rates and base frequencies of the known simulated tree.

**Supplementary file 5.** Inferred transition/transversion rates and base frequencies when using the consensus sequence. Numbers show the average of 100 simulations.

**Supplementary file 6.** Inferred transition/transversion rates and base frequencies when accounting for within-host diversity. Numbers show the average of 100 simulations



Figure 1—figure supplement 1. Collection date distribution and time from symptom and days from symptom onset.

(a) Distribution of collection dates. (b) Histogram of time from symptom onset to sample collection.



**Figure 1—figure supplement 2. Sample mean coverage distribution.** Density distribution of mean coverage.



Figure 1—figure supplement 3. Effects of C, value on whole-genome sequencing data.

**a** Higher C, values were linked to a higher number of within-sample variation. **b** Correlation between C, value and isolate sequencing mean coverage. Lower coverage was associated to higher C, values ( $R^2 = 0.13$ , t-statistic p-value < 0.001).





Individual plots of shared within-host variants between technical duplicates using increasing thresholds of allele frequency. Colors represent C, value, while the size of the point shows the total number of within-host variants between the two samples.









Probability that low frequency variants are shared inferred with a logistic model with allele frequency and epidemiological relationship as independent variable and whether a variant is shared or not as dependent variable. Y-axis in logarithmic scale for representation.



Figure 2—figure supplement 3. Dynamics of low frequency variants in longitudinal duplicates.

Variant frequency of low frequency variants through time in longitudinal duplicates. Each panel represents a single individual, with variants indicated by dots at each time point. The same variant at different time points is linked by lines. Yellow colors represent variants that are consistently found at each time point, while grey dots show variants that present in the first sampling event but lost in subsequent isolates.





Comparison of the phylogenetic trees inferred using simulated sequences with different transition/transversion rates to reflect different within-host diversity levels. Colors show the different rates of within-host evolution. Light colors represent trees inferred with consensus alignments, while dark colors show trees inferred with the model accounting for within-host diversity.



Figure 4—figure supplement 2. Similarity scores for inferred trees from coalescent simulations.

Comparison of the phylogenetic trees inferred using simulated sequences from known coalescent starting trees and different phylogenetic models. Colors differentiate the metrics used for the comparison.



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#### Figure 6—figure supplement 1. Phylogenetic trees for SARS-CoV-2.

SARS-CoV-2 phylogenetic trees inferred from consensus sequences (left) and an alignment with major and minor variant information (right).



#### Figure 6—figure supplement 2. Genetic distance between longitudinal samples.

The genetic distance in the phylogenetic tree inferred using within-sample diversity increased as the between longitudinal samples progressed. Black line shows the best fit in a linear model, while the blue shaded area represents the 95% Cl.



# Figure 6—figure supplement 3. Number of low frequency variants within outbreaks as the outbreak progresses.

Y-axis shows the number of low frequency variants for each isolate within an outbreak, while the x-axis represents the days since that particular outbreak started. Black line shows the best fit in a linear model, while the blue shaded area represents the 95% CI.





SARS-CoV-2 phylogenetic trees inferred from consensus sequences (left) and an alignment with major and minor variant information (right). Branch lengths are measured in years.







**a-d** Phylogenies of SARS-CoV-2 outbreaks. The branch lengths are in units of substitutions per genome, and the scales are shown under the trees. Colors represent samples from the same individual. Samples with the same name are technical replicates. Left tree of each panel shows the phylogeny inferred with the consensus alignment. Right tree represents the phylogeny inferred using within-sample variation. Heatmap shows the likelihood of direct transmission for each pair of samples in a SEIR model of transmission. Vertical axis is the infector while the horizontal axis shows the infectee.