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# Title

Pandemic dynamics of COVID-19 using epidemic stage, instantaneous reproductive number and pathogen genome identity (GENI) score: modeling molecular epidemiology

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Authors Bandoy, DJ Darwin R Weimer, Bart C

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- 1 Title:
- 2 Pandemic dynamics of COVID-19 using epidemic stage, instantaneous reproductive number
- 3 and pathogen genome identity (GENI) score: modeling molecular epidemiology
- 4 5
- 6 **Running Head:** Integrated molecular epidemiology and modeling
- 7
- 8 Authors:
- 9 DJ Darwin R. Bandoy<sup>1,2</sup> and Bart C. Weimer<sup>1\*</sup> 10

### 11 Affiliations:

- <sup>1</sup>University of California Davis, School of Veterinary Medicine, 100K Pathogen Genome Project,
   Davis, CA 95616, USA; <sup>2</sup>University of the Philippines Los Baños, College of Veterinary
   Medicine, Department of Veterinary Paraclinical Sciences, Laguna 4031, Philippines
- 15
- 16 17

18 \*corresponding author: <u>bcweimer@ucdavis.edu</u>; +1-530-760-9550

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### 22 Key words:

- 23 reproductive number, COVID-19, corona virus, infectious disease, genomic epidemiology 24
- 25

### 26 **Research in context**

27 Reproductive number is (R) an epidemiological parameter that defines outbreak transmission 28 dynamics. While early estimates of R exist for COVID-19, the sample size is relatively small 29 (<2000 individuals) taken during the early stages of the disease in China. The outbreak is now a 30 pandemic and a more comprehensive assessment is needed to guide public health efforts in 31 making informed decisions to control regional outbreaks. Commonly, R is computed using a 32 sliding window approach, hence assessment of impact of intervention is more difficult to 33 estimate and often underestimates the dynamic nature of R as the outbreak progresses and 34 expands to different regions of the world. Parallel to epidemiological metrics, pathogen whole 35 genome sequencing is being used to infer transmission dynamics. Viral genome analysis 36 requires expert knowledge in understanding viral genomics that can be integrated with the rapid 37 responses needed for public health to advance outbreak mitigation. This study establishes 38 integrative approaches of genome sequencing with established epidemiological outbreak 39 metrics to provide an easily understandable estimate of transmission dynamics aimed at public 40 health response using evidence-based estimates.

41

#### 42 Added value of this study

43 Estimates of R are dynamic within the progression of the epidemic curve. Using the framework 44 defined in this study with dynamic estimates of R specific to each epicurve stage combined with 45 whole genome sequencing led to creation of a novel metric called GENI (pathogen genome 46 identity) that provides genomic evolution and variation in the context of the outbreak dynamics. 47 The GENI scores were directly linked and proportional to outbreak changes when using disease 48 incidence from epicurve stages (index, takeoff, exponential, and decline). By simulating short 49 and standard (2 day and 7 day, respectively) serial intervals, we calculated instantaneous R 50 followed by a global comparison that was associated with changes in GENI. This approach 51 quantified R values that are impacted by public health intervention to change the outbreak

trajectory and were linked to case incidence (i.e. exponential expansion or decelerating) by country. Integrating viral whole genome sequences to estimate GENI we were able to infer circulation time, local transmission, and index case introduction. Systematic integration of viral whole genome sequences with epidemiological parameters resulted in a simplified approach in assessing the status of outbreak that facilitates decisions using evidence from genomics and epidemiology in combination.

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### 59 Implications of all the available evidence

This study created a framework of evidence-based intervention by integrating whole genome sequencing and epidemiology during the COVID-19 pandemic. Calculating instantaneous R at different stages of the epicurve for different countries provided an evidence-based assessment of control measures as well as the underlying genomic variation globally that changed the outbreak trajectory for all countries examined. Use of the GENI score translates sequencing data into a public health metric that can be directly integrated in epidemiology for outbreak intervention and global preparedness systems.

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#### 68 Abstract

Background: Global spread of COVID-19 created an unprecedented infectious disease crisis that progressed to a pandemic with >180,000 cases in >100 countries. Reproductive number (R) is an outbreak metric estimating the transmission of a pathogen. Initial R values were published based on the early outbreak in China with limited number of cases with whole genome sequencing. Initial comparisons failed to show a direct relationship viral genomic diversity and epidemic severity was not established for SARS-Cov-2.

Methods: Each country's COVID-19 outbreak status was classified according to epicurve stage (index, takeoff, exponential, decline). Instantaneous R estimates (Wallinga and Teunis method) with a short and standard serial interval examined asymptomatic spread. Whole genome sequences were used to quantify the pathogen genome identity score that were used to estimate transmission time and epicurve stage. Transmission time was estimated based on
 evolutionary rate of 2 mutations/month.

Findings: The country-specific R revealed variable infection dynamics between and within outbreak stages. Outside China, R estimates revealed propagating epidemics poised to move into the takeoff and exponential stages. Population density and local temperatures had variable relationship to the outbreaks. GENI scores differentiated countries in index stage with cryptic transmission. Integration of incidence data with genome variation directly increases in cases with increased genome variation.

87 **Interpretation:** R was dynamic for each country and during the outbreak stage. Integrating 88 the outbreak dynamic, dynamic R, and genome variation found a direct association between 89 cases and genome variation. Synergistically, GENI provides an evidence-based transmission 90 metric that can be determined by sequencing the virus from each case. We calculated an 91 instantaneous country-specific R at different stages of outbreaks and formulated a novel metric 92 for infection dynamics using viral genome sequences to capture gaps in untraceable 93 transmission. Integrating epidemiology with genome sequencing allows evidence-based 94 dynamic disease outbreak tracking with predictive evidence. 95

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 Weimer laboratory.

98

### 99 Introduction

100 Outbreaks are defined by the reproductive number (R)<sup>1,2</sup> a common measure of

101 transmission. Probability of further disease spread is evaluated based on the threshold value

102 with likely expansion for values >2 and decline with values of <1. R is the main component for

103 computing the needed proportion of the population to be vaccinated based on herd immunity<sup>3</sup>.

104 The expansion of COVID-19 was determined with the earliest estimate of R = 2.2 (95% CI, 1.4

105 to 3.9) using serial intervals for 424 patients in Wuhan. China<sup>4</sup>. Recalculation with 2033 cases 106 estimated R = 2.2 to 3.6<sup>5</sup>. However, estimates of R for other countries where cases were found 107 as the outbreak grew in China were not done routinely and currently a fixed estimate R is used 108 based on the refined estimate from China. However, this is falling short in predicting the spread 109 of the pandemic and expansion within individual locations, suggesting that R is not likely to be 110 constant and likely to be dynamic for each outbreak location that results in underestimates of 111 the spread rate. This limitation is hindering epidemic dynamics as previously noted due to the 112 parameter is context specific and dynamic<sup>1,2</sup>. Hence, there is a need to rapidly estimate country 113 specific R values during the epidemic. This will provide global comparisons of expansion at 114 each location.

115 The Wallinga and Teunis method for R estimation requires input of outbreak incidences and 116 the serial interval (i.e. the period between the manifestation of symptoms in the primary case 117 and the onset of symptoms in secondary cases)<sup>6</sup>. This approach was implemented in a web 118 resource to estimate R during epidemics<sup>7</sup>. A key advantage of the approach is the ease of 119 production of credible intervals compared to other maximum likelihood estimation approaches. 120 Yet to be done is integration of viral genetic variation with R estimates but one study found that 121 there was no obvious relationship between R, severity of the epidemic and COVID-19 genome 122 diversity<sup>20</sup>.

123 COVID-19 has reached global spread in all continents except Antarctica and was defined to be a pandemic by the World Health Organization (WHO) in March 2020<sup>8-10</sup>. The outbreak 124 125 dynamics are different between countries as well as varying within individual countries. In part 126 this is due to varying and diverse healthcare systems, socio-cultural contexts, and rigorous 127 testing. Considering the lack of containment globally, except in Singapore, Hong Kong, and 128 Taiwan, we hypothesized that previously calculated R values do not provide reliable estimates 129 because they are more dynamic than is being considered and that influx of new cases and viral 130 mutation are likely sustaining expansion. While viral sequencing is occurring, it is not being

effectively integrated with epidemiological information because there is no existing frameworkfor that to systematically occur.

133 In spite of no clear path for deep integration of viral variation the current pandemic has 134 demonstrated the public health unity for sharing COVID-19 whole genome sequences with an 135 unprecedented openness. By quickly sharing the genome sequences it enables investigation of 136 the genome variation during the outbreak using multiple approaches and samples of the virus 137 genome space. It is approaching a viral population scale, which provides additional information 138 that cannot be gleaned with few sequences. Prior work established the value of estimating 139 transmission dynamics of rapidly evolving RNA viruses and highlights the capability to infer 140 transmission during outbreaks coupled with pathogen genomes<sup>11,12</sup>. This approach was 141 validated in EBOV and MERS. Each virus variant is separated by only several mutations yet produces new dynamics during the outbreak<sup>13,14</sup>. Rapidly evolving pathogens undergo genome 142 143 sequence mutation, selection pressure, random drift and stochastic events between infected 144 individuals<sup>11</sup>. Even small changes in the genome enable transmission that is determined by 145 accounting for the mutations between isolate sequences. It is recognized that the COVID-19 146 genome is changing over the outbreak but there is controversy about the impact and specifics of 147 the exact mutations. In this study, we used incidence data to derive R and compared country 148 specific COVID-19 infection dynamics with viral population genome diversity. By incorporating 149 R, epidemic curve timing, and viral genome diversity we created a systematic framework that 150 deduced how viral genome diversity can be used to describe epidemiological features of an 151 outbreak before new cases were observed. This was done by creating a genome diversity 152 metric that was directly and systematic integrated to provide context and allowed quantification 153 of the infection dynamics globally that are divergent from the early estimates with genomic 154 evidence. We call this approach pathogen genome identity (GENI) scoring system. Using GENI 155 differentiated each stage of the outbreak. It also indicated cryptic local transmission from

156 surveillance systems. This a defining advantage of using sequences as previous cryptic

157 transmission can be inferred in the genomic sequences.

158

### 159 Methods

Incidence data is based on daily Chinese CDC and WHO situations reports as compiled by
the Center for Systems Science and Engineering (CSSE) by the John Hopkins University
(Baltimore, MD, USA) that was accessed on March 1, 2020<sup>15</sup>. We constructed epidemic curves
or epicurves from the incidence data and classified country status accordingly. We defined four
groups that characterize increasing expansion with a decline phase.

165 The extracted time series case data served as the input for determining instantaneous 166 reproductive number on a daily basis to effectively capture dynamic changes due to new 167 detected cases and reduction of cases due to social distancing and nonpharmaceutical 168 interventions. The prior value for R was selected at 2 and prior standard deviation of 5 to allow 169 fluctuations in reporting of cases in the exponential phase. As there is limited access to 170 epidemiological data of case, parametric with uncertainty (offset gamma) distributional estimate 171 of serial interval was used. A mean of 2 and 7 days, with standard deviation of 1 was used to 172 capture short and standard serial interval assumptions using 50 samplings of serial interval distribution. The Wallinga and Teunis method, as implemented by Ferguson<sup>7</sup> is a likelihood-173 174 based estimation procedure that captures the temporal pattern of effective reproduction 175 numbers from an observed epidemic curve. R was calculated using the web application EpiEstim App (https://shiny.dide.imperial.ac.uk/epiestim/)<sup>7</sup>. The descriptive statistics were used 176 177 to compute mean and confidence intervals of the instantaneous reproductive number. 178 GENI score was anchored on the principle of rapid pathogen evolution between 179 transmission events. This requires defining a suitable reference sequence of the outbreak, 180 which is on the early stages the sequence nearest to the timepoint of the index case. For the 181 case of COVID-19, the reference sequence is Wuhan seafood market pneumonia virus isolate 182 Wuhan-Hu-1 NC 045512.2<sup>16</sup>. Publicly available virus sequences were retrieved from GISAID

(supplementary Table 1) with whole genome variant determination using Snippy v4.6.0<sup>17-19</sup>. The 183 184 average mutation per isolate was divided to the total epidemic curve days to derive a daily 185 epidemic mutation rate and scaled to a monthly rate. We calculated the average nucleotide 186 change per month to be 1.7 (95% CI 1.4-2.0), which was within boundaries of another estimate 187 with the substitution rate of  $0.9 \times 10^{-3}$  (95% CI 0.5-1.4  $\times 10^{-3}$ ) substitutions per site per year<sup>20</sup>. 188 We derived a transformed value of this rate before integrating it with epidemiological 189 information. The output from the variant calling step was then used to determine GENI score by 190 calculating the nucleotide difference. The basis for GENI score cutoffs to estimate transmission 191 dates are derived from accepted evolutionary inference of mutation rates of COVID-19.

192 We defined four epidemic curve (epicurves) stages to provide a clear method to define 193 increases in the outbreak. The 'index stage' is characterized by the first report (index case) or 194 limited local transmission indicated by intermittent zero incidence creating undulating epicurve. 195 Secondly, which is distinctly different from stage 1, is the 'takeoff stage' in which the troughs are 196 almost at same level of the previous peak and no longer touches zero, suggesting sustained local 197 transmission. The 'exponential stage' is characterized by the classical hockey stick like sharp 198 uptrend where the outbreak is moving quickly and large number of new cases are emerging. The 199 last stage is 'decline' and is noted when the outbreak has reached the peak and cases being 200 reported are lower than the peak, which will ultimately result in few to no new cases being 201 reported, yet viral circulation is likely still occurring.

#### 202 Results

We determined the outbreak dynamics of pandemic COVID-19 by classifying each country's status according to epicurve stages with a framework of a) index b) takeoff c) exponential d) decline as a clear method that can be used to benchmark metrics that include R and viral genome diversity. First, we calculated R using the instantaneous method using two serial intervals (2 and 7 days; Table 1). As of March 1, 2020, this framework defined global epicurves of COVID-19 outbreaks as gaining momentum globally with 52 countries were in the index stage. Three countries were in the exponential stage and five countries in the takeoff stage (Figure 1). China was the only country that reached the peak of the epicurve and characterized to be in the decline stage - decreasing cases. At this point there was no evidence of any other country near the decline stage and some countries that were poised to move into the takeoff and exponential phase.

214 Instantaneous R sensitively described real-time shifts of COVID-19 incidence captured 215 within each epicurve stage (Figure 2). The decline stage in China was reflected by a decrease 216 in R estimates in the latter stages the outbreak and relative to the early estimates: 1.6 (95 % CI 217 0.4-2.9) and 1.8 (95 % CI 1.0-2.7) for 2- and 7-days serial interval, respectively. Superspreading 218 events inflated R estimates seen in exponential stage that was observed in South Korea: 2.8 219 (95% CI 0.6-5.3) and 25.6 (95 % CI 3.0-48.2) for 2- and 7-days serial interval, respectively. 220 Efficient disease control was instituted in Singapore enabling it to remain in the index stage 221 while Japan was moving to the takeoff stage characterized by increased R estimates 3.6 (95% 222 CI 0.4-7.3) 2.2 (95% CI 1.3-3.0) for 2- and 7-days serial interval, respectively. The R estimates 223 overlaps for all exemplar country outbreak stages in the two serial interval scenarios, suggesting 224 that the transmission could be as short as 2 days. These estimates were relatively lower than 225 previously reported, bringing to light possibility of transmission in the incubation period that is 226 associated with rapidly expanding outbreaks, which is currently being observed in many 227 European countries.

Low detection of COVID-19 was observed in representative countries in the index stage with low R values (<2) that can be attributed to effectiveness of social distancing intervention (i.e. Hong Kong) or under detection for countries with limited testing (i.e. United States) (Figure 3a). Sustained local transmission was occurring in five countries that were progressing into takeoff stage (Japan, Germany, Spain, Kuwait and France) as measured by R values (>2) (Figure 3b). The magnitude of spread was apparent with relatively higher R estimates (>10) in Italy, Iran and South Korea, which demonstrated sudden surges in incidence due to prior undetected clusters in part but other factors may contribute to this observation (Figure 3b). This significantly
increased the instantaneous R estimates versus other methods of estimation but allows a more
obvious depiction of the surge of cases allows differentiation of the takeoff stage from
exponential stage.

239 We further examined the value of computing country-specific instantaneous R by 240 comparing different temperature range (tropical versus temperate) and population density. 241 Population density of key cities (Table 2) and the higher temperature range values were used 242 for selected countries; however, no direct link was observed. Increases in the South Korean 243 outbreak was associated with a secretive religious group Shinsheonji (73% cases of COVID-19 244 in South Korea) located mainly in Daegu with a lower population density 883/km<sup>2</sup> as compared 245 to the rest of the areas with an outbreak<sup>21</sup> and likely explain the outbreak expansion in the early 246 epicurve. Religious beliefs that modify health seeking behavior particularly reporting clinical 247 signs of COVID-19 combined with continued large group gathering prevented early detection of 248 the outbreak. While most countries (Table 2) have cooler temperatures (10-6°C), Singapore's 249 temperature higher indicated that local transmission occurred at higher temperatures and 250 suggests that temperature shifts will not likely change transmission. These commonly accepted 251 environmental and behavioral activities did not explain the epicurve. This led to the hypothesis 252 that the viral genomic variation underpinned changes in cases during outbreaks in each country.

253 We determined the relationship of epicurve stage with viral genetic variation using a 254 metric that merges absolute genome variation with the rate of genome change to create the 255 GENI metric that anchored population genome diversity with the rate of evolution for the SARS-256 Cov-2. To examine how the viral genome diversity was associated with the epicurve stages we 257 first examined the index stage (Singapore) and the exponential (South Korea). Integration of 258 GENI scores successfully distinguished the index from exponential stage (Figure 4). An 259 increase in GENI scores was associated with exponential stage with a median score of 4. 260 suggesting that the viral diversity and rate of mutation played was directly proportional to case

261 increases during this stage. Singapore (index stage) effectively controlled the disease before 262 becoming exponential had a GENI median score of 2. This was found in multiple time points 263 during the outbreak were multiple mutation events were directly associated with increases in 264 cases. While China is in the decline stage the retrospective association with R, cases, and 265 GENI provided longitudinal evidence of multiple expansion in cases with mutation events in the 266 viral genome, especially early in the epicurve. The repeated viral mutations and epicurve 267 expansion were associated in each time point over 3 months, in three countries, and in three 268 different outbreak stages. This finding is useful in integrating virus genome diversity and 269 evolution into assessment of outbreak status in an outbreak between countries but also within 270 the epicurve when combined into a triad with instantaneous R estimates. The proportionality of 271 GENI scores with the epicurve stage indicates its value in determining the outbreak status and 272 the importance of generating population scale genome sequence resources.

273 A framework to merge epidemiology and population genomics was derived from this 274 study as a systematic method for molecular epidemiology (Fig. 5). It requires dynamic 275 measurements be taken for R and longitudinal efforts to determine each virus whole genome 276 sequence. Using this triad of measurements accurately and guickly provided insight to measure 277 outbreak progress but also provides an evidence-based method for interventions. This study 278 demonstrated an advancement of how to use population genomics in a viral situation where the 279 mutation rate is fast and the genome diversity of the population is extraordinarily high. GENI 280 provided a missing method that defines how to use viral genome mutation dynamics and 281 genome population diversity, which is only observable using large numbers of genomes, that 282 occurs during an outbreak.

## 283

#### 284 **Discussion**:

Public health response is proportional to the severity and transmission dynamics of an
infectious disease outbreak. This requires epidemiological metrics that can be used as decision
criteria, and ideally, they can be used to assess impact of the intervention. In this work we

288 determined that R is much more dynamic in the COVID-19 pandemic than previously 289 appreciated by country as well as over the outbreak within each country (Fig 2-3). The 290 instantaneous R estimation with a serial interval of 2 was extremely sensitive to shifts in the 291 epicurve during the index phase (Fig 2-3). Singapore is an excellent example of effectively 292 controlling and containing the COVID-19 outbreak. They previously designated a response system called Dorscon (Disease Outbreak Response System Condition)<sup>22</sup> providing a 293 294 systematic approach to control so that they have not moved past the index phase. In contrast, 295 most other countries in this phase are poised to move into the takeoff phase (Fig 3). The 296 transition into the takeoff phase signified a transition from a 2-day serial interval to a 7-day serial 297 interval that was more sensitive to shifts in the epicurve. 298 While estimates of R alone is insightful in retrospect, gaps in epidemiological surveillance 299 due to several factors creates blind spots that hindered the ability to determine interventions. To 300 overcome this limitation, we merged GENI estimates based on whole genome sequence 301 variation and mutation rate with the epicurve and R and provided a predictive triad of 302 measurement that resulted in insight that accurately refined case expansion (Fig. 4). Each 303 phase of the outbreak was characterized with mutations that led to new cases in established 304 outbreaks by case definition. The merged information indicate that China found variation in the 305 viral sequence much earlier than the outbreak cases increased. Independent of the phase 306 framework merging sequence variants with the epicurve found that new cases were observed in 307 the same timeframe as new sequence variants were found. Previous studies that the

308 relationship of genomic diversity with epidemic severity (i.e. R) found no clear link<sup>20</sup>. However,

309 by merging instantaneous R, the epicurve stage, and the GENI index it is clear that a link exists

310 for each country examined that resulted in a direct link between outbreak dynamics and the

311 absolute genomic mutation with the mutation rate. The GENI index provides a basis to examine

312 imported cases or locally spreading, both of which addressed this current work using

313 established metric - R and novel integration of viral whole genome sequences to define changes

314 in the sequence that are directly linked to increases in cases. This leads to an epidemiological 315 metric that is scientifically robust and at the same time can convey complex biological properties 316 to enable an efficient characterization of an outbreak in combination. Transforming complex 317 pathogen characteristics was made usable to public health and medical field using the GENI 318 score as a complete merged information set with other characteristics of the outbreak. 319 Previous outbreaks, such as Ebola, employed state of the art analysis using phylodynamics that is anchored on the genetic evolution<sup>13</sup>. Inference such as time to most recent common 320 321 ancestor allowed estimation of outbreak origin, population size, and R - yet this was not 322 integrated into the outbreak dynamics and stage of advancement in the outbreak. This type of 323 analysis is possible because genomic sequences carry temporal signals and when used in 324 context with sample from different timepoints, previous divergence can be determined. The 325 GENI score includes these signals and expands their use by merging them with the outbreak 326 dynamic using the population genome variation as well as the mutation rate. 327 This inherit information is not limited to viruses. Another recent example in a bacterial setting 328 was the cholerae outbreak in Haiti wherein the phylogenetic analysis resolved the origin of the 329 pathogen<sup>23</sup>. However, for this analysis to succeed, a substantial database of genome 330 sequences is needed, collected across time and geographic location to enable placement in a 331 phylogenetic context. As outbreaks as bound to happen in the future, investment in cataloguing 332 the genomic space of pathogens is as ever important<sup>24,25</sup>. It is critical to obtain COVID-19 333 sequences from humans as well as other animals that have zoonotic potential, as was demonstrated previously with zoonotic *Campylobacter* species<sup>26,27</sup>. Creating sequence 334

repositories of pathogens is critical and underway for various pathogens<sup>25</sup> as well as COVID-

**336 19**<sup>18</sup>.

Prior work forewarned the practice of being overly dependent on early estimates of R
 alone<sup>28</sup>. By having the most accurate possible information for a dynamic metric and taking into
 account the complex dynamics that factor in the calculation of R along with merging this the

340 aenomics of the pathogen is a robust and insightful method to assess outbreak dynamics, as 341 demonstrated in this study. Openness and data sharing of incidence reports and sequences at unprecedented scale is being done in this pandemic and it is paving rewards<sup>29</sup>. Leveraging on 342 343 these resources opens unexpected collaboration and avenues for applying relevant 344 bioinformatic and disease modelling skills across the scientific community to solve global public 345 health problems. Examples that hindered this were observed in several countries that led to 346 cryptic spread of the disease in countries. Additionally, lacking the epidemiological infrastructure 347 and genome sequencing capabilities limit this approach that is not acceptable for modern public 348 health. However, without the appropriate technical skills in the performing complicated 349 phylogenetic inference, utility of such innovation will be limited. Establishing a protocol for 350 merging epidemiology and genomics was defined in this work (Fig. 5) and can be instituted 351 globally. 352 353 Conclusion 354 This study integrated population genomics into epidemiological methods to provide a framework 355 for molecular epidemiology. Specifically, this study demonstrated using epicurves, 356 instantaneous R estimates, and GENI specific case increases in COVID-19 are directly 357 associated with viral mutation. It was demonstrated that the pandemic is poised to become 358 larger and that mutation will be associated with the increase in cases. Exemplar outbreaks, such 359 as Singapore, found increases in cases with viral mutations that were effectively controlled. 360 However, other outbreaks had expanding R estimates during the outbreak, as well as numerous 361 viral mutation events. Use of epicurve stages, instantaneous R estimates, and GENI provided a 362 robust and accurate framework to monitor outbreak progression to different stages with direct 363 association between cases and increases in each metric. 364

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- 369

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448		

#### 449 **Figure legends**:

450 Figure 1. Distribution of country classification based on COVID-19 epicurve status.

451

452 Figure 2. Instantaneous reproductive number estimates for different stages of the COVID-19 453 epidemic curve: a) index (Singapore) b) takeoff (Japan) c) Exponential (South Korea) d) decline 454 (China) in short (2 days) and standard (7 days) serial interval. Decelerating stage of epidemic 455 curve results to a reproductive number lower than 2 for both serial intervals, epidemic curve with 456 multiple introductions yields 2-day serial interval with higher reproductive number and 457 exponential serial interval yields higher reproductive number for the 7-day serial interval. Dot (.) 458 the surge in the epidemic curve of China corresponds to the alteration of the case definition of 459 COVID-19 by broadening confirmed cases with pneumonia confirmed with CT (computed 460 tomography) scan. South Korea's higher reproductive number is due to cryptic transmission 461 associated with a secretive cult with altered health seeking behavior.

462

463 Figure 3. Epicurve estimates with different serial intervals. Panel A represents Epicurves and 464 instantaneous R values for index stage countries using 2- and 7-day serial interval. Panel B 465 Global dynamics of COVID-19 using instantaneous estimate of reproductive number with 2-day 466 serial interval. Under preincubation period infectivity scenario, reproductive numbers globally 467 increasing (> 2). Italy's R = 8 is highest due to late detection of infection clusters. This higher R 468 estimate is due to a huge bump in cases combined with diagnostic gap of low-level incidence. 469 The same surge dynamics is seen in South Korea. Global dynamics of COVID-19 using 470 instantaneous estimate of reproductive number with 7-day serial interval. Italy's R value inflates 471 to 57 with the 7-day serial interval assumption and overlaps with the lower threshold of 2 day 472 serial interval R estimate. This estimation depicts a decreasing pattern for countries multiple 473 introductions like Singapore, Hong Kong.

474

475 Figure 4. Relationship of pathogen genome identity (GENI) score with the temporal signal along 476 the epidemic curve. Local transmission is captured by virus mutation as expressed in GENI 477 score values. GENI scores of SARS-COV2 isolates are relative to Wuhan reference strain 478 Wuhan-Hu-1 NC 045512.2. The red line in the China epicurve represents the time before an 479 outbreak was determined yet genome sequences were circulating. The blue shaded curves 480 indicate GENE scores directly overlaid with the outbreak curve. The dotted line represents the 481 common point in time as a reference for visualization. The GENI score and epicurve show 482 similarity except in China as the outbreak advanced to takeoff and exponential the GENI score 483 increased while in the index stage example of Singapore the outbreak was contained and the 484 GENI score remained <2.

485

Figure 5. Integration of genomic and classical epidemiology for outbreak investigation. The foundation of epidemiology is the accurate and timely reporting of cases which enable the calculation of the number. Genomic Identity (GENI) score is formulated from genomic data of pathogens to differentiate imported cases versus local transmission and measure time of cryptic spread. Together these two epidemic values deliver insight that can be directly used for making decision criteria for public health intervention.

492

## **Table 1.** Country-specific Instantaneous Reproductive Number (R) estimates for COVID-19 as

## 495 of March 1, 2020.

		Instantaneous Reproductive Number (R) with different serial intervals		
Country	Cases	2 days	7 days	
Mainland China	79251	1.6	2.1	
South Korea	3150	2.8	25.6	
Italy	1128	8	57.0	
Iran	593	2.8	17.1	
Japan	241	3.6	2.2	
Singapore	102	3.3	1.6	
France	100	2.9	16.9	
Hong Kong	95	2.6	1.6	
Germany	79	3.1	17.2	
United States	70	4.3	1.7	
Kuwait	45	2.6	15.3	
Spain	45	3.7	10.8	
Thailand	42	3.8	1.7	

- **Table 2**. Epidemiological Parameters and instantaneous R estimates. The population density for
- 499 South Korea is based on Daegu where 75% of the cases are reported.

### 

	Reproductive Number (R)	Temperature (°C) during outbreak	Population Density (people/km²)	Interpretation in consideration of the epidemiological curve
Singapore	3.3	32	8136	Imported cases, limited local transmission
France	2.9	10	4300	Imported, Local transmission >1-2 month
Italy	8	10	7200	Imported cases, Local transmission >1 month
United States	4.3	9	8444	Imported cases, Local transmission >2 month
South Korea	2.8	6	883	Imported cases, Local transmission >1-2 month

- 503 **Table 3**. Relationship of Pathogen Genome Identity (GENI) Score derived from mutational
- 504 difference from the index genome (Wuhan isolate of COVID-19 or cluster isolate reference from
- 505 multiple outbreak regions outside of territory).
- 506

Equivalent Pathogen Genome Identity (GENI) score for COVID- 19	Clinical Interpretation and Epidemiological Inference	Notes
0-2	No difference from index case isolate genome or reference, imported case if there is no prior report, indicative of acute transmission <1 month	Reference genome is primarily earliest isolate available.
3-4	recent local transmission (average 1-2 months) if there are no prior report of cases	Subsequent outbreak clusters can serve as sources of introduction hence near neighbor reference has to be selected to generate an accurate GENI score.
>4	sustained local transmission (greater than 2 months) if there is are no prior report of cases	Subsequent outbreak clusters can serve as sources of introduction hence near neighbor reference has to be selected to generate an accurate GENI score.





Figure 2

# Index Stage



### Takeoff Stage





Figure 4

