Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Detection of baloxavir resistant influenza A viruses using next generation sequencing and pyrosequencing methods

Mira C. Patel a, Vasilyi P. Mishin a, Juan A. De La Cruz a,b, Anton Chesnokov a, Ha T. Nguyen a,b, Malania M. Wilson a, John Barnes a, Rebecca J.G. Kondor a, David E. Wentworth a, Larisa V. Gubareva a,b

a Influenza Division, National Center for Immunization and Respiratory Diseases, Centers of Disease Control and Prevention, Atlanta, GA, USA
b Battelle Memorial Institute, Atlanta, GA, USA

1. Introduction

A new anti-influenza drug, baloxavir marboxil (baloxavir) was approved for treatment of influenza A and B infections in Japan and the U.S. in 2018, and since then it has entered markets in numerous countries. Baloxavir binds to polymerase acidic (PA) subunit of viral RNA polymerase and hinders the cap-dependent endonuclease activity, which is crucial for generation of capped RNA primers for viral transcription (Noshi et al., 2018). Influenza A viruses have been shown to develop resistance to baloxavir by acquiring substitutions at one of the highly conserved residues in the PA catalytic site - isoleucine 38 (I38X). Influenza A viruses carrying I38T (threonine) or I38S (serine) in PA can be readily selected in cell culture in presence of baloxavir (Chesnokov et al., 2020; Noshi et al., 2018). Notably, natural polymorphisms at PA residue 38 (PA-38) are rare among influenza A viruses (Gubareva et al., 2019; Omoto et al., 2018; Stevaert et al., 2013). Substitutions at PA-38 produce a varying effect on baloxavir susceptibility, depending on the substituted amino acid and virus subtype. Valine (V) does not seem to affect baloxavir susceptibility, while leucine (L) conferred 10-fold reduced susceptibility in A(H1N1)pdm09 and ~3-fold in A(H3N2) viruses (Gubareva et al., 2019; Noshi et al., 2018; Omoto et al., 2018, CDC unpublished data). The amino acids threonine (T), methionine (M) or phenylalanine (F) confer 10- >50-fold reduced susceptibility to baloxavir (Omoto et al., 2018) and detected in viruses collected from ~2 to 23% of baloxavir recipients infected with influenza A viruses (Hayden et al., 2018; Hirotsu et al., 2019). Virological surveillance conducted in Japan during 2018–19 season detected viruses carrying PA-38 substitutions with frequency of 2.3% in A(H1N1)pdm09 and 8.0% in A(H3N2) viruses, while no I38-substituted influenza B virus was reported (https://www.niid.go.jp/niid/images/flu/resistance/20191227/dr18-1).

* Corresponding author. Influenza Division, National Center for Immunization and Respiratory Diseases, Centers of Disease Control and Prevention, 1600 Clifton Road, MS-H17-5, Atlanta, GA, 30329-4027, USA.
E-mail address: lgubareva@cdc.gov (L.V. Gubareva).
https://doi.org/10.1016/j.antiviral.2020.104906
Received 22 May 2020; Received in revised form 15 July 2020; Accepted 5 August 2020
Available online 14 August 2020
0166-3542/Published by Elsevier B.V. This is an open access article under the CC BY license (http://creativecommons.org/licenses/by/4.0/).
2. Material and methods

2.1. Viruses

Influenza A viruses used in this study were submitted to the WHO Collaborating Centre for Surveillance, Epidemiology and Control of Influenza at the CDC by U.S. PHLs and other laboratories participating in GISRS (Table S1). A reverse genetically engineered influenza A(H1N1) pdm09 virus containing I38F, was kindly provided by WHO Collaborating Centre for Reference and Research on Influenza at Melbourne (Koszalka et al., 2019). Additionally, A(H1N1)pdm09 viruses carrying either I38S or I38T were selected by culturing A/Illinois/08/2018 (I38) in Madin-Darby canine kidney (MDCK) cells for five passages in presence of baloxavir (Chesnokov et al., 2020).

Complete genome sequences of both derivative viruses were analyzed by NGS. Besides I38S, no additional substitutions were detected in the I38S carrying virus; while the genome of I38T virus had one additional substitution, HA-K119N (Chesnokov et al., 2020). A(H1N1)pdm09 viruses were propagated in MDCK cells, whereas A(H3N2) viruses were propagated in MDCK-SAT1 cells (kindly provided by M. Matrosovich). Prior to use in this study, all viruses were propagated by infecting cells at a low multiplicity of infection (MOI) and incubating for short period of time, except two viruses containing I38V, A/California/153/2016 and A/Hawaii/89/2016, were propagated using procedure for routine surveillance to produce high hemagglutination titers (https://www.who.int/influenza/gisrs_laboratory/manual_diagnosis_surveillance_influenza/en/). (Table S1).

To evaluate accuracy of detection and quantitation of viral subpopulations by pyrosequencing, we prepared subtype-specific artificial mixtures of two or three viruses carrying different amino acids at PA-38. The mixtures were prepared using virus stock preparations, and ratios were determined using NGS analysis.

2.2. Next generation sequencing

Viral RNA was extracted using QIAamp Viral RNA Mini Kit (Qiagen) and codon-complete influenza genome was amplified using Uni/Inf primer set and Super-Script III One-Step RT-PCR with Platinum Taq High Fidelity enzyme (Invitrogen). Indexed paired end libraries were generated using Nextera XT Sample Preparation Kit (Illumina) following the manufacturer’s protocol. Illumina MiSeq was used to generate sequences that were analyzed by the IRMA approach (Shepard et al., 2016). The current cut-off for reportable SNP for depositing NGS data to public databases (GISAID and GenBank) and for routine surveillance purposes is set at 20–25%, although sequence data are stored and can be re-analyzed to provide a more detailed information on viral quasispecies composition. To analyze artificially prepared virus mixtures, RNA extraction, PCR amplification and library preparation were carried out three times independently to evaluate any possible variation that the sequencing process may have on PA-38 variant detection.

2.3. Pyrosequencing

Viral RNA was extracted from 100 μl of sample using MagNA Pure Compact Nucleic Acid Isolation Kit I (Roche Diagnostics). A SuperScript III One-Step RT-PCR system with Platinum Taq High Fidelity enzyme (Invitrogen) was used for cDNA synthesis and amplification. Type A-specific primers, forward InfA-F58 and reverse InfA-R280biot, were used for RT-PCR (Table S2). Subtype-specific reverse primers (InfH1-R1141biot and InfH3-R1126biot) were used to generate long amplicons (Table S2). Pyrosequencing reactions were performed on PyroMark Q96 ID (Qiagen) instrument as described by the manufacturer’s protocol. Both sequence analysis (SQA) and allele quantitation (AQ) modes of the instrument were utilized for analysis of substitutions at PA-38. Two different nucleotide dispensation orders, cyclic (GCAT)G and customized GCAAGC TTC(GCAT)G, were used to perform pyrosequencing by
SQA mode. SNP analysis by AQ mode was conducted to determine the proportion of each PA-38 variant in a mixture. The dispensation order for AQ analysis was generated by PyroMark software depending on the targeted SNP sequence. A 10% cut-off was used to determine the detection of an SNP (Deyde et al., 2009).

RNA extraction followed by RT-PCR and pyrosequencing, was carried out three independent times to calculate proportions (%) of each PA-38 variant in artificially prepared virus mixtures. All pyrograms were visually inspected by the operator. Pyrograms were interpreted as: “pass” if variant sequences at PA-38 and downstream sequence were correctly determined; “fail” if PA-38 variant sequences and downstream sequence could not be correctly interpreted; “indeterminant” if PA-38 variant sequences could not be conclusively determined, but the downstream sequence was correctly identified.

3. Results

3.1. Optimization of pyrosequencing to detect PA-38 variants in the double mixtures

A panel of ten seasonal influenza A viruses with and without amino acid substitutions at PA-38 was assembled for this study (Table S1). This panel comprised of viruses detected by surveillance, selected in vitro, or generated using reverse genetics. Viruses were tested using, 1) type A-specific primers for RT-PCR amplification, 2) subtype-specific sequencing primers, and 3) the cyclic order of nucleotide dispensation (GCA)₃₆G₃₆, as described previously (Methods and Table S2) (Chesnokov et al., 2020; Gubareva et al., 2019). With this approach, amino acids I, F, M, L, S, T, or V at PA-38 were readily identified (data not shown).

It is common for respiratory specimens collected from drug-treated patients to contain a mixture of sensitive and resistant viruses. Indeed, viruses with and without amino acid substitution at PA-38, e.g., I38 T/I, were detected in baloxavir-treated patients (Takahata et al., 2019a, 2019b; Uehara et al., 2019). Although rare, virus culturing can also result in emergence of PA polymorphic variants, including at PA-38 (Chesnokov et al., 2020; Gubareva et al., 2019).

We wanted to investigate whether PA-38 variants in the mixtures could be reliably identified using pyrosequencing assay. To this end, we prepared artificial mixtures containing two viruses, one of which had I38 (wildtype) (Table 1). Using cyclic dispensation (SQA mode), we tested five different double mixtures of A(H1N1)pdm09 subtype. Notably, only one, containing I/F, was identified correctly after visual inspection of pyrogram and passed the test (Table 1). Pyrograms for other mixtures - I/L, I/S, I/T, or I/V - failed quality assessment due to misalignment of downstream sequence peaks (see a representative pyrogram for I/T in Fig. 1A) and could not be interpreted by an operator. Therefore, we wanted to know whether the assay performance can be improved by using a customized order of nucleotide dispensation. Based on analysis of PA gene sequence alignment for both the subtypes (not shown) and an approach previously described (Levine et al., 2011), we customized the dispensation by including nine non-cyclic nucleotides prior to four rounds of cyclic dispensions, which should result in synchronized extension of downstream sequence reads for the variants. Addition of A and T at positions four and eight of customized dispensation order was derived: GCAAGCTTC(GCAT)₆G₆, as described previously (Methods and Table S2) (Chesnokov et al., 2012). Thus, the following dispensation order was derived: GCAAGCTTC(GCAT)₆. When the single variants were tested using this customized dispensation, all different amino acids at PA-38 were easily identified in both subtypes (representative pyrograms in Fig. 1B and C). For all five double mixtures of A(H1N1)pdm09 tested, visual inspection of pyrograms was required to pass the test and confirm the presence of both the variants (representative pyrogram in Fig. 1D, Table 1).

Next, we tested double mixtures of A(H3N2) viruses using both cyclic and customized nucleotide dispensions. Regardless of dispensation order used, both PA variants in the mixtures were correctly identified (Table 1). Because customized dispensation produced good results for both virus subtypes, it was used in all subsequent experiments.

3.2. Comparison of PA-38 variant proportions in the double mixtures using pyrosequencing and NGS analysis

The proportions of PA-38 variants were determined in eight artificial

### Table 1

| Subtype       | Virus name and amino acid at PA-38 | Codon | PA-38 mixture | Pyrosequencing, SQA results | Percentages of detected variants (Mean ± SD) | NGS | Pyrosequencing, AQ results |
|---------------|-----------------------------------|-------|---------------|-----------------------------|---------------------------------------------|-----|---------------------------|
|               |                                   |       |               | Cyclic          | Custom         | Short amplicon | Long amplicon |
| H1N1pdm09     | A/Illinois/08/2018-I38            | ATT   | I/F           | pass            | pass           | 76.7 ± 3.7    | 76.3 ± 1.2    | –              |
|               | RG-A/Perth/261/2009-I38F         | TTT   | I/L           | fail            | pass           | 23.3 ± 3.7    | 23.7 ± 1.2    | –              |
|               | A/Illinois/08/2018-I38            | ATT   | I/S           | fail            | pass           | 54.6 ± 0.5    | 45.4 ± 1.2    | –              |
|               | A/Illinois/08/2018-138L           | CTT   |               |                |                | 54.6 ± 1.2    | 54.6 ± 1.2    | –              |
|               | A/Illinois/08/2018-138            | ATT   | I/T           | fail            | pass           | 42.2 ± 0.1    | 47.3 ± 1.3    | –              |
|               | A/Illinois/08/2018-138T           | AGT   |               |                |                | 45.2 ± 1.2    | 45.2 ± 1.2    | –              |
|               | A/Illinois/08/2018-138             | ACT   |               |                |                | 52.7 ± 1.3    | 52.7 ± 1.3    | –              |
|               | A/Illinois/08/2018-138             | ATT   | I/V           | fail            | pass           | 54.3 ± 5.8    | 71.0 ± 0.8    | –              |
|               | A/California/153/2016-138V        | GCT   |               |                |                | 57.7 ± 0.5    | 57.7 ± 0.5    | –              |
| H3N2          | A/Louisiana/50/2017-I38           | ATA   | I/M           | pass            | pass           | 64.3 ± 0.9    | 57.7 ± 0.5    | –              |
|               | A/Louisiana/49/2017-I38M          | ATG   |               |                |                | 64.3 ± 0.9    | 57.7 ± 0.5    | –              |
|               | A/Louisiana/50/2017-I38           | ATA   | I/T           | pass            | pass           | 68.9 ± 5.5    | 76.5 ± 2.2    | –              |
|               | A/Bangladesh/3007/2017-I38T       | ACA   |               |                |                | 76.5 ± 2.2    | 76.5 ± 2.2    | –              |
|               | A/Louisiana/50/2017-I38           | ATA   | I/V           | pass            | pass           | 56.4 ± 0.2    | 60.0 ± 2.4    | –              |
|               | A/Hawaii/89/2016-138V             | GTA   |               |                |                | 43.6 ± 0.2    | 40.0 ± 2.4    | –              |

AQ: allele quantitation, RG: reverse genetically engineered virus, SD: standard deviation, -: not tested.

* Letters in boldface indicate the nucleotide change compared to wildtype codon for isoleucine (I).

* Pyrosequencing was performed in SQA mode using cyclic (GCA)₃₆₆G₃₆ and customized GCAAGCTTC(GCAT)₆ nucleotide dispensation orders to detect PA-38 variants in the mixtures. All the pyrograms were visually inspected by the operator: pass = both variants were correctly identified; fail = variant sequences could not be determined.

* Results are derived from three independently carried out RNA extraction, RT-PCR and pyrosequencing; and three independent RNA extraction, PCR amplification and library preparation for NGS.

* Dispensation order for AQ analysis was automatically generated by PyroMark software depending on the targeted sequences and a position of SNP.
mixtures by both, NGS analysis and pyrosequencing (Table 1). For the latter assay, testing was done using AQ mode, which requires knowledge of targeted sequences and a position of SNP. Of note, NGS and pyrosequencing were performed independently, including RNA extraction and downstream steps. Considering this experimental set-up, the proportions determined by both assays correlated well with differences ranging 0.4–8%, except for two mixtures containing I38V, which showed difference of 17–24% (Table 1). This discrepant outcome required further investigation. Among the viruses used to prepare mixtures, only two viruses carrying I38V were grown differently (see Methods). It is known that infection of cells at high MOI and prolonged incubation can lead to accumulation of defective interfering (DI) particles. Such DI particles often contain defective gene segments encoding polymerase proteins (Dimmock and Easton, 2014). The defective gene segments mostly retain the gene termini but lack the middle region. Notably, NGS method utilizes universal primers to 5′ and 3′ termini of all gene segments, which results in amplification of DNA amplicon is short (223 bp), and the targeted nucleotide triplet (PA-38) resides close to the gene terminus. Therefore, we wanted to test whether use of a long amplicon would improve the assessment of virus sub-populations by pyrosequencing. To this end, we used new subtype-specific reverse primers, InfH1-R1141biot and InfH3-R1126biot, to generate long amplicons (1084 and 1069bp, Fig. 1. Representative pyrograms and software readouts for influenza A(H1N1)pdm09 viruses containing one or two PA-38 variants. Pyrosequencing was carried out in SQA mode using either (A) cyclic (GCAAGCTTC(GCAT)₆ nucleotide dispensation orders. (D) Arrow points to a nucleotide “C” identified at the second position of the triplet encoding the amino acid at PA-38; indicating the presence of 38 T. (E) Pyrogram of 38I/T mixture by AQ mode with nucleotide percentages provided by the PyroMark ID software (blue box). Expected sequences were based on NGS analysis. Underlined nucleotides indicate codon for PA-38. Lower case nucleotides indicate the single nucleotide polymorphism (C and T) at the second position of codon for PA-38. Highlighted readout is determined by the software: Blue = pass; Yellow = check (visual inspection by the operator is required to interpret the sequence and determine pass or fail); Red = fail.
respectively) (Table S2). Indeed, sequencing of the long amplicons yielded a decreased proportion of I38V in both mixtures (Table 1), bringing the results of pyrosequencing closer to those of NGS. Of note, use of a short DNA amplicon for pyrosequencing is preferred as it is likely to improve testing results for samples with low virus load.

3.3. Analysis of A(H3N2) virus isolates containing mixture of PA-38 variants

Next, using pyrosequencing we tested three A(H3N2) virus isolates containing PA-38 polymorphic variants I/M, I/T, or I/K, as determined by NGS analysis conducted during routine surveillance (Chesnokov et al., 2020; Gubareva et al., 2019). For A/Massachusetts/04/2019 (passage S2), the proportion of I38M variant determined by NGS and pyrosequencing was similar, 27% vs. 25% (Table 2). For the second isolate, A/Bangladesh/3007/2017 (passage S2), NGS detected ~10% lower proportion of I38T variant (39% vs 50%). It is worth noting that I38-substituted viruses identified in both isolates were successfully purified by conventional limiting dilution and tested phenotypically. Substitutions I38M and I38T conferred 11- and 116-fold reduced baloxavir susceptibility, respectively (Gubareva et al., 2019). Notably, neither assay detected I38-substituted variants in respective clinical specimens, suggesting that the PA-38 mutants were selected during virus culturing. For the third isolate, A/Hawaii/28/2017 (passage S2), there was an apparent discordance between the results of NGS and pyrosequencing. While NGS analysis detected I38K variant in both instances, clinical specimen (14%) and the virus isolate (27%); pyrosequencing did not detect it in either sample (Table 2). In addition, presence of I38K in the virus isolate could not be ascertained using Sanger sequencing (Table S2, data not shown). Lysine (K) can be encoded by nucleotide triplets AAG or AAA. For A/Hawaii/28/2017, the nucleotide substitution was detected by NGS analysis at the second position in the triplet, ATA → AAA. We attempted to recover the I38K-containing virus by limiting dilution. A total of 62 biological virus clones were generated from the S2 virus isolate and tested using pyrosequencing, but all clones showed only ATA at PA-38 (wildtype). It was shown that the substitution of a nonpolar isoleucine at 38 with a polar amino acid (T or S) conferred greater baloxavir resistance, respectively (Gubareva et al., 2019). Moreover, pyrograms showed only ATA at PA-38 (wildtype). On the other hand, the customized dispensation nucleotide dispensation orders was suitable for identifying different amino acid substitutions at PA-38 of seasonal influenza A viruses. However, the cyclic dispensation was not optimal for detecting amino acid substitutions at PA-38 of seasonal influenza A viruses.

Table 2
PA-I38-substituted variant proportions in influenza A(H3N2) virus samples determined using NGS analysis and pyrosequencing.

| Virus name | PA GISAID accession # | Sample | Percentage of PA-I38-substituted variant (Mean ± SD) |
|------------|------------------------|--------|---------------------------------------------------|
|            |                        |        | NGS Pyrosequencing (AQ)                            |
| A/Massachusetts/04/2019 | EP11362029 clinical isolate | ND | ND |
| A/Bangladesh/3007/2017  | EP1107334 clinical isolate | T | 49.7 ± 1.2 |
| A/Hawaii/28/2017        | EP1106149 clinical isolate | M | 24.8 ± 0.7 |
|                        | EP1107334 clinical isolate | K | 27 (27) |
|                        | EP11016323 clinical isolate | K | 27 (27) |

ND: not detected (below limit of detection), S2: passage 2 in MDCK-SIAT1 cells.

3.4. Detection of PA-38 variants in triple mixtures using pyrosequencing

Reportedly, virus samples collected from baloxavir-treated patients occasionally contain three PA-38 variants (Takashita et al., 2019a, 2019b; Uehara et al., 2019). Therefore, we prepared artificial virus mixtures containing three different PA variants and tested them using NGS and pyrosequencing (Table S3). In three instances, presence of all PA-38 variants was correctly identified upon visual inspection of pyrograms, which were deemed “pass”. For two other mixtures, not all PA-38 variant sequences could be conclusively determined and were deemed “indeterminant” (Table S3). Due to consecutive changes at more than two nucleotides of the same codon, built-in algorithm of the PyroMark software did not support SNP analysis of triple mixtures tested here using AQ mode. Overall, it was more challenging to determine PA-38 variants in triple mixtures of A(H1N1)pdm09 subtype. While all variants in the A(H1N1)pdm09 triple mixture containing I/S/T were identified, the other two mixtures (I/L/T and I/F/T) could not be resolved using pyrosequencing method described here. Visual inspection of pyrogram of the triple mixture I/L/T suggested the presence of ATT and ACT, while presence of CTT in the mixture was not evident. Similarly, it was difficult to ascertain the presence of all three PA-38 variants in the mixture I/F/T (data not shown).

4. Discussion

Over the past few years, NGS analysis has become the primary method of influenza virological surveillance in the U.S., including drug resistance monitoring (Jester et al., 2018). Availability of viral genome sequences is especially desirable as it facilitates the establishment of molecular markers associated with decreased susceptibility for new drugs. Conversely, pyrosequencing assay is useful for detecting previously established markers of drug resistance by generating short, targeted sequence readouts. Once optimized, pyrosequencing can provide a formidable method to test hundreds of samples in a timely manner. Here we showed that pyrosequencing using both cyclic and customized nucleotide dispensation orders was suitable for identifying different amino acid substitutions at PA-38 of seasonal influenza A viruses. However, the cyclic dispensation was not optimal for detecting I38-substituted A(H1N1)pdm09 viruses present in a mixture with the wildtype (I38) virus. On the other hand, the customized dispensation improved the assay performance in detecting PA-38 variants in double mixtures. The improvement, however, was not sufficient to identify all variants in triple mixtures. Because double or triple mixtures are anticipated in specimens collected from baloxavir-treated patients, it is prudent not to rely on software sequence readout, but to verify pyrogram by visual inspection. In some instances of triple mixtures, even visual inspection of pyrograms could not ascertain the presence of all the PA-38 variants and therefore the result of these mixtures was deemed “indeterminant”. This limitation of the assay may not be critical for surveillance purposes, but it could complicate the interpretation of testing outcomes needed to make informed clinical care decisions. Moreover, pyrosequencing has an inherent limitation due to use of dATP analog deoxyadenosine-β-thio triphosphate in the reaction as it produces a higher peak on the pyrogram than the other dNTPs (Harrington et al., 2013). This needs to be kept in mind, when visually inspecting pyrograms.

Koszalka and colleagues recently published a study, where they explored utility of pyrosequencing for rapid detection of PA-I38X variants in influenza A and B viruses (Koszalka et al., 2020). There were
several differences in the assay design compared to our study (different cyclic dispensation order, biotinylation of forward primer, generation of shorter amplification (100bp), and use of DNA plasmids to prepare artificial PA-38 variant mixtures). The authors showed that pyrosequencing using AQ mode could estimate proportions of variants in known artificial mixtures containing I/T and I/M, but not I/F; and triple mixtures were not tested. Because of the intuitive nature of the pyrosequencing assay, it is possible to create hypothetical scenarios to determine the outcome of a sequence readout. Applying such an approach made us question whether testing unknown samples carrying two to three different variants would yield conclusive data. Taken together, we believe that a customized dispensation order of nucleotides provides an improvement in resolving samples carrying mixtures of PA-38 variants compared to cyclic dispensation orders (Gubareva et al., 2019; Koszalka et al., 2020). Our data indicate that DI particles containing defective PA gene segments may affect the proportion of a viral population detected by pyrosequencing compared to NGS analysis, at least in artificially prepared mixtures. However, this is unlikely to affect testing of clinical specimens. One limitation of our study is not propagating I38V containing viruses using a low MOI and incubating for short period of time as was done for other viruses.

Every sequencing technique has limitations and may introduce biases or errors (e.g., general increase of errors toward the end of reads for Illumina) (Schrömer et al., 2016). In our study, NGS analysis detected the presence of I38K variant in A/Hawaii/28/2017 (H3N2) virus; while pyrosequencing, Sanger sequencing and limiting dilution did not. To the best of our knowledge, viruses carrying I38K as a dominant variant have not been reported by global surveillance or detected after baloxavir exposure. However, virus specimen from a baloxavir-treated patient was reported to contain I38K as a mixture (I38 T/K/I) in Japan during 2018–19 season (Takashita et al., 2019a). Recently Jones and colleagues have shown that introduction of polar, positively charged lysine (K) at PA-38 reduced the polymerase activity by 50% compared to wildtype (I38) in minirepliicon assay (Jones et al., 2020). However, they apparently did not investigate the effect of this amino acid substitution on virus properties. To improve the current knowledge, it would be desirable to generate a reverse genetically modified virus and determine effect of I38K on virus fitness and baloxavir susceptibility. This information would benefit the interpretation of PA-38 sequencing results for clinical care use. In future experiments, viruses with recently reported PA-38 substitutions (e.g., I38N) could be derived using reverse genetics and tested by our assay to further show its utility (Ince et al., 2020; Imai et al., 2020).

It is worth noting that the frequency of baloxavir resistance detection among circulating viruses remains low in the U.S. (https://www.cdc.gov/flu/weekly/weeklyarchives2019-2020/Week13.htm). However, PA-38 substitutions were detected in 2.3% of A(H1N1)pdm09 and 8.0% of A(H3N2) viruses collected during 2018–19 season of Japan, with the majority of mutant viruses being isolated from baloxavir-treated children (https://www.niid.go.jp/niid/images/flu/resistance/20191227/dr18-19e20191227-1.pdf,Takashita et al., 2019a). The widespread use of influenza antiviral can lead to emergence of resistant viruses and is a serious public health concern.

In the past, it was suggested that the widespread use of adamantanes, incited by the fear of ‘bird flu’ and the SARS coronavirus facilitated emergence of adamantante-resistant viruses in Asia in 2003–2004 (Bright et al., 2005) and their global spread in the following years (Bright et al., 2006; Deyde et al., 2007). Therefore, it is conceivable that severe influenza seasons and ongoing SARS-CoV-2 pandemic can also lead to rampant use of available antivirals, including baloxavir, which in turn may cause the emergence and spread of drug-resistant viruses. In view of this possibility, it would be prudent to strengthen laboratory capabilities to allow early detection of drug resistance emergence.

In conclusion, various genotypic assays have been used to detect drug resistance markers in influenza viruses, including baloxavir resistant viruses (Gubareva et al., 2019; Koszalka et al., 2020; Nakauchi et al., 2020). All methods have their advantages and disadvantages and decision which assay to use depends on the community needs and laboratory capabilities.

Declaration of competing interest

None.

Acknowledgements

We are thankful to the U.S. Association of Public Health Laboratories (APHL) and other laboratories participating in World Health Organization Global Influenza Surveillance and Response System (WHO-GISRS) for productive collaboration with the CDC Influenza Division and submission of influenza viruses. The authors would like to thank Dr. Aaron Hurt from Melbourne WHO Collaborating Centre for Reference and Research on Influenza for kindly providing reverse genetically modified baloxavir-resistant reference viruses. We would like to acknowledge the valuable contributions of Thomas Stark and other colleagues from the Virology, Surveillance and Diagnosis Branch of the CDC Influenza Division.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.antiviral.2020.104906.

Funding source

This study was supported by the Influenza Division of Centers for Disease Control and Prevention(CDC). The study does not reflect the official position of the CDC but personal opinions of authors.

Table S1

| Subtype | Virus name | Amino acid at PA-38 | Codon | GISAID # EPI-JSL | Source | Reference |
|---------|------------|---------------------|-------|-----------------|--------|-----------|
| H1N1pdm09 | A/Illinois/08/2018 | I | ATT | 315855 | Surveillance | Gubareva et al., 2019 |
| RG-A/Perth/261/2009, I38F | F | TTT | N/A | Reverse Genetics | Koszalka et al., 2019 |
| A/Illinois/37/2018 | L | CTT | 315856 | Surveillance | Gubareva et al., 2019 |
| A/Illinois/08/2018 | S | AGT | 365522 | In vitro selection | Chesnokov et al., 2020 |
| A/Illinois/08/2018 | T | ACT | 348120 | In vitro selection | Chesnokov et al., 2020 |
| A/California/153/2016 | V | GTT | 241691 | Surveillance | Gubareva et al., 2019 |
| H3N2 | A/Louisiana/50/2017 | I | ATA | 315857 | Surveillance | Gubareva et al., 2019 |

(continued on next page)
Table S1 (continued)

| Subtype            | Virus name | Amino acid at PA-38 | Codon | GISAID # EPI_ISL_ | Source | Reference                  |
|--------------------|------------|---------------------|-------|-------------------|--------|----------------------------|
| A/Louisiana/2017   | M          | ATG                 |       | 315858            | Surveillance | Gubareva et al., 2019       |
| A/Bangladesh/2007  | T          | ACA                 |       | 286069            | Surveillance | Chensovok et al., 2020      |
| A/Hawaii/2016      | V          | GTA                 |       | 23918i            | Surveillance | Gubareva et al., 2019       |

N/A: Not available.

* Letters in boldface indicate the nucleotide change compared to wildtype codon for isoleucine (I).

References

Bright, R.A., Medina, M.J., Xu, X., Perez-Oronoz, G., Wallis, T.R., Davis, X.M., Povinelli, L., Cox, N.J., Klomov, A.I., 2005. Incidence of adamantane resistance among influenza A (H3N2) viruses isolated worldwide from 1994 to 2005: a cause for concern. Lancet 365, 1175-1181.

Bright, R.A., Shay, D.K., Shu, B., Cox, N.J., Klomov, A.I., 2006. Adamantane resistance among influenza A viruses isolated early during the 2005-2006 influenza season in the United States. J. Am. Med. Assoc. 295, 691-694.

Chesnokov, A., Patel, M.C., Mishin, V.P., De La Cruz, J.A., Lollis, L., Nguyen, H.T., Tsuchiya, K., Hayden, F.G., Uehara, T., Watanabe, A., 2019. Baloxavir marboxil in influenza A(H3N2) viruses circulating in the United States during the 2016/17 and 2017/18 seasons. Euro Surveill. 24, 1800666.

Deyde, V.M., Shu, T.G., Trujillo, A.A., Okomo-Adhiamb, M., Garten, R., Klomov, A.I., Gubareva, L.V., 2010. Detection of molecular markers of drug resistance in 2009 pandemic influenza A (H1N1) viruses by pyrosequencing. Antimicrob. Agents Chemother. 54, 1102-1110.

Deyde, V.M., Okomo-Adhiamb, M., Shu, T.G., Wallis, T.R., Fry, A., Dharan, N., Klomov, A.I., Gubareva, L.V., 2009. Pyrosequencing as a tool to detect molecular markers of resistance to neuraminidase inhibitors in seasonal influenza A viruses. Antivir. Res. 81, 16-24.

Deyde, V.M., Xu, X., Bright, R.A., Shaw, M., Smith, C.B., Zhang, Y., Shu, Y., Gubareva, L.V., Cox, N.J., Klomov, A.I., 2007. Surveillance of resistance to adamantanes among influenza A(H1N1) and A(H3N2) viruses isolated worldwide. J. Infect. Dis. 196, 249-257.

Dimmock, N.J., Easton, A.J., 2014. Defective interfering influenza virus RNAs: time to reevaluate their clinical potential as broad-spectrum antivirals? J. Virol. 88, 5217-5227.

Gubareva, L.V., Mishin, V.P., Patel, M.C., Chesnokov, A., Nguyen, H.T., De La Cruz, J., Spencer, S., Campbell, A.P., Sinner, M., Reid, H., Garten, R., Katz, J.M., Fry, A.M., Barnes, J., Wentworth, D.E., 2019. Assessing baloxavir susceptibility of influenza viruses circulating in the United States during the 2016/17 and 2017/18 seasons. Euro Surveill. 24, 1800666.

Harrington, C.T., Lin, E.I., Olson, M.T., Eshleman, J.R., 2013. Fundamentals of pyrosequencing. Arch. Pathol. Lab Med. 137, 1296-1303.

Hayden, F.G., Sugaya, N., Hirotsu, N., Lee, N., de Jong, M.D., Hurt, A.C., Ishida, T., Sekino, H., Yamada, K., Portsmouth, S., Kawaguchi, K., Shishido, S., Araki, M., Tsuikiya, K., Uehara, T., Watanabe, A., Baloxavir Marboxil Investigators, G., 2018. Baloxavir marboxil for uncomplicated influenza in adults and adolescents. N. Engl. J. Med. 379, 913-923.

Hirotsu, N., Sakaguchi, H., Sato, C., Ishibashi, T., Baba, K., Omoto, S., Shishido, T., Tsuikiya, K., Hayden, F.G., Uehara, T., Watanabe, A., 2019. Baloxavir marboxil in

Table S2

Primer sets used for RT-PCR, pyrosequencing and Sanger sequencing for analysis of PA-38 codon.

| Name of primer | Sequence (5' to 3') |
|----------------|---------------------|
| RT-PCR         | InfA-5BS             | GCAAATGCTTC(GCAT) |
|                | InfA-R280biot        | biot-ATGCTTCAGCC |
|                | InfH1-R1141biot      | biot-ATGCTTTCTGTGCCATAT |
|                | InfH3-R1126biot      | biot-ATGCTTTCTGTGCAAGAG |
| Pyrosequencing  | InfH1-F91            | GAAACTAATAGTTTGGTCG |
|                | InfH3-F95            | GACACAAATTTGCAGCC |
| Sanger sequencing | InfH3-F70            | GAAAAAGCGAATGAAAGGT |

Table S3

Assessment of PA-38 variants in artificially prepared influenza A virus mixtures.

| Subtype          | Virus name and amino acid at PA-38 | PA-38 mixture | Codon | Percentages of variants by NGS (Mean ± SD) | Pyrosequencing, SQA results |
|------------------|-----------------------------------|---------------|-------|------------------------------------------|----------------------------|
| H1N1pd09         | A/Illinois/08/2018-I38             | I/L/T         | ATT   | 47.9 ± 1.6                                | indeterminant              |
|                  | A/Illinois/03/2018-I38             | C/T/T         | CT    | 28.4 ± 0.8                                |                            |
|                  | A/Illinois/08/2018-I38             | I/F/T         | TT    | 28.6 ± 0.7                                | indeterminant              |
|                  | A/Illinois/03/2018-I38             | I/F/T         | TTT   | 6.5 ± 0.7                                 |                            |
|                  | A/Illinois/08/2018-I38             | I/S/T         | ATT   | 57.7 ± 0.1                                | Pass (I/S/T)               |
|                  | A/Illinois/08/2018-I38             | I/S/T         | AGT   | 20.2 ± 0.7                                |                            |
|                  | A/Illinois/08/2018-I38             | I/S/T         | ACT   | 22.1 ± 0.8                                |                            |
|                  | A/Illinois/2017-I38                | I/M/T         | ATG   | 22.9 ± 0.9                                | Pass (I/M/T)               |
|                  | A/Illinois/2017-I38                | I/M/T         | M     | 50.9 ± 1.2                                |                            |
|                  | A/Bangladesh/2007/2017-I38         | V/M/T         | GTA   | 21.4 ± 1.0                                | Pass (V/M/T)               |
|                  | A/Illinois/2017-I38                | V/M/T         | M     | 40.2 ± 2.7                                |                            |
|                  | A/Bangladesh/2007/2017-I38         | V/M/T         | A     | 38.3 ± 2.0                                |                            |

RG: reverse genetically engineered virus.

* Letters in boldface indicate the nucleotide change compared to wildtype codon for isoleucine (I).

† Results are derived from three independent RNA extraction, PCR amplification and library preparation.

‡ Pyrosequencing was performed in SQA mode using customized nucleotide dispensation order GCAAGCTTC(GCAT)4 to detect variants at PA-38 in the mixtures. All the pyrograms were visually inspected by the operator: pass = all variants were correctly identified; indeterminant = all variants could not be conclusively identified.
Japanese pediatric patients with influenza: safety and clinical and virologic outcomes. Clin. Infect. Dis. https://doi.org/10.1093/cid/ciz908.

Imai, M., Yamashita, M., Sakai-Tagawa, Y., Ieisutuki-Horimoto, K., Kiso, M., Murakami, J., Yasuhara, A., Takada, K., Ito, M., Nakajima, N., Takahashi, K., Lopes, J.S.S., Dutta, J., Khan, Z., Kriti, D., van Bakel, K., Tokita, A., Hagiwara, H., Izumida, N., Kuroki, H., Nishino, T., Wada, N., Koga, A., Adachi, E., Jubish, D., Hasegawa, H., Kawoaka, Y., 2020. Influenza A variants with reduced susceptibility to baloxavir isolated from Japanese patients are fit and transmit through respiratory droplets. Nat Microbiol 5, 27–33.

Ince, W.L., Smith, F.R., O'Rear, J.J., Thomson, M., 2020. Treatment-emergent influenza virus PA substitutions independent of those at I38 associated with reduced baloxavir susceptibility and virus rebound in trials of baloxavir marboxil. J. Infect. Dis. https://doi.org/10.1093/infdis/jiaa164.

Jester, B., Schwerzmann, J., Mustaqué, D., Aden, T., Brammer, L., Humes, R., Shult, P., Shahangian, S., Gubareva, L., Xu, X., Miller, J., Jernigan, D., 2018. Mapping of the US domestic influenza virologic surveillance landscape. Emerg. Infect. Dis. 24, 1300–1306.

Jones, J.C., Pascua, P.N.Q., Fabrizio, T.P., Marathe, B.M., Seiler, P., Barman, S., Webby, R.J., Webster, R.G., Govorkova, E.A., 2020. Influenza A and B viruses with reduced baloxavir susceptibility display attenuated in vitro fitness but retain ferret transmissibility. Proc. Natl. Acad. Sci. U. S. A. 118, 8593–8601.

Koszalka, P., Farrerke, R., Mitraud, E., Vijaykrishna, D., Hult, A.C., 2020. A rapid pyrosequencing assay for the molecular detection of influenza viruses with reduced baloxavir susceptibility due to PA/I38X amino acid substitutions. Influenza Other Respir Viruses 14, 460–464.

Koszalka, P., Tilman, D., Ro, M., Vijaykrishna, D., Hult, A.C., 2019. Baloxavir marboxil susceptibility of influenza viruses from the Asia-Pacific, 2012-2018. Antivir. Res. 164, 91–96.

Levine, M., Sheu, T.G., Gubareva, L.V., Mishin, V.P., 2011. Detection of hemagglutinin variants of the pandemic influenza A(H1N1) 2009 virus by pyrosequencing. J. Clin. Microbiol. 49, 1307–1312.

Nakache, M., Takashita, E., Fujisaki, S., Shirakura, M., Ogawa, R., Morita, H., Miura, H., Saito, S., Watanabe, S., Omidii, T., Kageyama, T., 2020. Rapid detection of an I38T amino acid substitution in influenza polymerase acidic subunit associated with reduced susceptibility to baloxavir marboxil. Influenza Other Respir Viruses 14, 436–443.

Nakache, M., Ujike, M., Obuchi, M., Takashita, E., Takayama, I., Ejima, M., Obo, K., Konomi, N., Omidii, T., Tashiro, M., Kageyama, T., influenza virus surveillance group of, J., 2011. Rapid discrimination of oseltamivir-resistant 275Y and -susceptible 275H substitutions in the neuraminidase gene of pandemic influenza A/ H1N1 2009 virus by duplex one-step RT-PCR assay. J. Med. Virol. 83, 1121–1127.

Noshi, T., Kizato, M., Taniguchi, K., Yamamoto, A., Omidii, F., Baba, K., Hashimoto, T., Ishida, K., Kushima, Y., Ialtori, K., Kawai, M., Yoshida, R., Kobayashi, M., Yoshinaga, T., Sato, A., Okamatsu, M., Sakoda, Y., Kida, H., Shishido, T., Naito, A., 2018. In vitro characterization of baloxavir acid, a first-in-class cap-dependent endoribonuclease inhibitor of the influenza virus polymerase PA subunit. Antivir. Res. 160, 109–117.

Omoto, S., Speranzini, V., Hashimoto, T., Noshi, T., Yamaguchi, H., Kawai, M., Kawaguchi, K., Uehara, T., Shishido, T., Naito, A., Cusack, S., 2018. Characterization of influenza virus variants induced by treatment with the endoribonuclease inhibitor baloxavir marboxil. Sci. Rep. 8, 9635.

Schirmer, M., D’Amore, R., Ijaz, U.Z., Hall, N., Quince, C., 2016. Illumina error profiles: resolving fine-scale variation in metagenomic sequencing data. BMC Bioinf. 17, 125.

Shepherd, S.S., Meno, S., Bahi, J., Wilson, M.M., Barnes, J., Neumann, E., 2016. Viral deep sequencing needs an adaptive approach: IRMA, the iterative refinement meta-assembler. BMC Genom. 17, 708.

Stevart, A., Dallocchio, R., Desi, A., Pala, N., Rogolino, D., Sechi, M., Naenssens, L., 2013. Mutational analysis of the binding pockets of the diolce acid inhibitor L-742,001 in the influenza virus PA endonuclease. J. Virol. 87, 10524–10538.

Storms, A.D., Gubareva, L.V., Su, S., Wheeling, J.T., Okomo-Adhiambo, M., Pan, C.Y., Reisdorf, E., St. George, K., Myers, R., Wotton, J.T., Robinson, S., Leader, B., Thompson, M., Shannon, M., Klimov, A., Fry, A.M., Group, U.S.A.R.S.W., 2012. Oseltamivir-resistant pandemic (H1N1) 2009 virus infections, United States, 2010-11. Emerg. Infect. Dis. 18, 308–311.

Takashita, E., Ichikawa, M., Morita, H., Ogawa, R., Fujisaki, S., Shirakura, M., Miura, H., Nakamura, K., Kishida, N., Kuwahara, T., Sugawara, H., Sato, A., Akimoto, M., Mitamura, K., Abe, T., Yamazaki, M., Hasegawa, S., Hasegawa, O., Ogadri, T., 2019a. Human-to-Human transmission of influenza A(H1N2) virus with reduced susceptibility to baloxavir, Japan, February 2019. Emerg. Infect. Dis. 25, 2108–2111.

Takashita, E., Kawakami, C., Ogawa, R., Morita, H., Fujisaki, S., Shirakura, M., Miura, H., Nakamura, K., Kishida, N., Kuwahara, T., Ota, A., Togashi, H., Saito, A., Mitamura, K., Abe, T., Ichikawa, M., Yamazaki, M., Watanabe, S., Ogadri, T., 2019b. Influenza A(H1N2) virus exhibiting reduced susceptibility to baloxavir due to a polymerase acidic subunit I38T substitution detected from a hospitalised child without prior baloxavir treatment, Japan, January 2019. Euro Surveill. 24, 1900170.

Tsiatis, A.C., Norris-Kirby, A., Rich, R.G., Hafez, M.J., Cocke, C.D., Ehleman, J.R., Murphy, K.M., 2010. Comparison of Sanger sequencing, pyrosequencing, and melting curve analysis for the detection of KRAS mutations: diagnostic and clinical implications. J. Mol. Diagn. 12, 425–432.

Uehara, T., Hayden, F.G., Kawaguchi, K., Omoto, S., Hult, A.C., De Jong, M.D., Hirotsu, N., Sugaya, N., Lee, N., Baba, K., Shishido, T., Tsujiya, K., Portsmouth, S., Kida, H., 2019. Treatment-Emergent influenza variant viruses with reduced baloxavir susceptibility: impact on clinical and virologic outcomes in uncomplicated influenza. J. Infect. Dis. 221, 346–355.

Zahoor, R., Vanke, L.J., Church, D., Topp, E., Read, R.R., McAllister, T.A., 2012. High-throughput species identification of enterococci using pyrosequencing. J. Microbiol. Methods 89, 174–178.