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Cell-based reporter assays for measurements of antibody-mediated cellular cytotoxicity and phagocytosis against SARS-CoV-2 spike protein

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The COVID-19 pandemic caused by SARS-CoV-2 infections has led to excess deaths worldwide. Neutralizing antibodies (nAbs) against viral spike protein acquired from natural infections or vaccinations contribute to protection against new- and re-infections. Besides neutralization, antibody-mediated cellular cytotoxicity (ADCC) and phagocytosis (ADCP) are also important for viral clearance. However, due to the lack of convenient methods, the ADCC and ADCP responses elicited by viral infections or vaccinations remain to be explored. Here, we developed cell-based assays using target cells stably expressing SARS-CoV-2 spikes and Jurkat-NFAT-CD16a/CD32a effector cells for ADCC/ADCP measurements of monoclonal antibodies and human convalescent COVID-19 plasmas (HCPs). In control samples (n = 190), the specificity was 99.5% (95%CI: 98.4–100%) and 97.4% (95%CI: 95.1–99.6%) for the ADCC and ADCP assays, respectively. Among 87 COVID-19 HCPs, 83 (sensitivity: 93.1%, 95%CI: 87.8–98.4%) showed detectable ADCC (titer range: 7.4–1721.6) and ADCP activities (titer range: 4–523.2). Notably, both ADCC and ADCP antibody titers positively correlated with the nAb titers in HCPs. In summary, we developed new tools for quantitative ADCC and ADCP analysis against SARS-CoV-2, which may facilitate further evaluations of Fc-mediated effector functions in preventing and treating against SARS-CoV-2.

1. Introduction

Coronavirus disease 2019 (COVID-19) pandemic caused by the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2) continues and has caused 518 million cases and 6.3 million deaths as of May 13, 2022 (World Health Organization, 2022). The spike protein of SARS-CoV-2 mediates the viral cellular binding and entry via interaction with the ACE2 receptors. SARS-CoV-2 spike proteins contain the furin cleavage site (RRAR motif), which allows S1/S2 cleavage during virus packaging and is required for cell-to-cell virus transmission (Hoffmann et al., 2020). Immune-response against spike proteins, particularly for the neutralization antibodies (nAbs), essentially contribute to the protection acquired from vaccination and natural infection recovery (Baum et al., 2020; Chi et al., 2020; Feng et al., 2021; Gilbert et al., 2022; Khoury et al., 2021). In addition to the neutralization effects, spike-specific antibodies may also eliminate viruses or virus-infected cells via antibody-dependent cellular cytotoxicity (ADCC), phagocytosis (ADCP), or complement-dependent cytotoxicity (CDC) (Ji et al., 2019; Tay et al., 2019; Zohar and Alter, 2020).

So far, several assays for analysis of activities of antibodies in neutralizing SARS-CoV-2 have been developed, including authentic virus-based cytopathogenic effect (CPE) inhibition or plaque reduction (PRNT) tests and pseudotyped lentivirus (LV) or vesicular stomatitis virus (VSV) based neutralization tests (Case et al., 2020; Manenti et al., 2020; Xie et al., 2020; Zhang et al., 2022). However, convenient and high-throughput methods for measurements of ADCC and ADCP activities in serum or plasma samples are still limited. The nuclear factor of activated T-cells (NFAT) signaling pathway is essential for Fcγ receptors
Purified mAbs were obtained from cell culture supernatants using the CHO transfection kit (Thermo Scientific, A29129) following previously described methods as target cells and developed NFAT-reporter cell-based assays for measurements of ADCC and ADCP activities of antibodies. The performance of new assays was systematically evaluated in several mAbs and human plasma samples.

2. Materials and methods

2.1. Monoclonal antibodies

A total of 10 reported SARS-CoV-2 anti-spike mAbs were tested for ADCC and ADCP activities (Baum et al., 2020; Cerutti et al., 2021; Chi et al., 2020; Pinto et al., 2021; Sauer et al., 2021; Tortorici et al., 2021). All mAbs were produced in ExpiCHO cells by transient transfection using a modified EIRBdMie vector (EIRB-Mie-derived) containing dual promoters for human IgG light chain and heavy chain. The coding sequences of the variable regions of 44A8, NTD4–8, BD368–2, COVA2–15, S2M11, REGN10933, S2P6 were synthesized in the human IgG1 backbone (Generalbiol, Anhui, China) according to the published literature. The 36H6, 83H7, and 85F7 were developed in our lab, and their variable regions cDNA sequences were synthesized in the human IgG1 backbone (Generalbiol, Anhui, China) ac-

2.2. Human samples for assay validations

Human convalescent COVID-19 plasma samples (n = 87) were collected from recovered COVID-19 patients (one sample for each patient) from the Xinglin Branch of the First Affiliated Hospital of Xiamen University. All involved COVID-19 HCP samples were collected in the convalescent phase (≥14 days from illness onset or diagnosis, range 14–122 days) and showed detectable anti-SARS-CoV-2 RBD antibodies in a double-sandwich immunoassay (Wantai Biological Pharmacy). Among these samples, 70 were collected between 14 and 90 days (early convalescent phase), and 17 were collected between 91 and 122 days (late convalescent phase) since illness or diagnosis onset. For specificity evaluation, 190 stored plasma samples pre-dating the SARS-CoV-2 outbreak (collected in 2019) were used. All samples were heat-inactivated for pseudovirus (PsV) neutralization tests and ADCC/ADCP assays. Informed consent was obtained from each subject. The study was approved by the institutional review board of the School of Public Health of Xiamen University following the Declaration of Helsinki.

2.3. Target cells with stable expression of SARS-CoV-2 spike

The codon-optimized full-length spike cassette (SFLwt) of SARS-CoV-2 (referring to MN908947.3) was synthesized (Generalbiol, Anhui, China). A furin-site mutated spike cassette (SFLfk, changed RRAR to GSAR) was generated via site-directed mutagenesis based on SFLwt. Both SFLwt and SFLfk were cloned into the EIRB-Mie vector, which is a modified PiggyBac vector with IRES-driven dual-selection markers of red-fluorescent protein (mRuby3) and Blasticidin S-resistance gene (Blasticidin S deaminase, Bsd). The Expi293F Cells (Thermo Scientific, A14528, 293 F) were co-transfected with the full-length spike expression vector (EIRB-Mie-SFLwt or EIRB-Mie-SFLfk) and a PiggyBac transposase expression vector (at a ratio of 4:1) using the PEI transfection system (BIOHUB, 78PEI25000–5 G). Then, blasticidin was supplied at 30 μg/mL in the culture medium for selections of cells with successful transfection since 24 h after transfection. Cells were cultured at 37 °C in a stackable CO2 (8%) incubator shaker (Kühner AG, SMX1503C). Subsequently, cells that were stably integrated with the transfected cassettes were obtained via continuous blasticidin selection and flow cytometry (FACS) cell sortings (mRuby3-activated).

2.4. Effector cells

The Jurkat-Lucia™ NFAT and Jurkat-Lucia™ NFAT-CD16 (hereafter designated Jurkat-NFAT-hCD16) cells were purchased from InvivoGen. Both cells were derived from the human Jurkat cell line by stable integration of an NFAT-inducible Lucia luciferase reporter (InvivoGen). The former is the parental reporter cell line and the latter having an additional expressing of human Fc receptor CD16A (FcRIIA; V158 allotype). For the generation of Jurkat-NFAT-hCD32 cells, the parental Jurkat-NFAT cells were stably transduced with a lentiviral vectored human CD23A (FcRIIA; H131 allotype) expression cassette. The CD23A cDNA is linked with an IRES-mRuby3-P2A-Bsr cassette (synthesized by Generalbiol, Anhui, China) and cloned into the plV-EF1a-MCS-IRES-Bsd vector. The presence of IRES-driven expression of mRuby3 and Bsd enable convenient positive-selections via blasticidin supplement and FACS-sorting. The Jurkat-NFAT-hCD16 and Jurkat-NFAT-hCD32 cells were cultured in SuperCulture serum-free lymphocyte medium (DAKEWEID) with Normocin (10 μg/mL), Blasticidin (10 μg/mL), and Zeocin (10 μg/mL) at 37 °C in a stackable CO2 (8%) incubator shaker.

2.5. FACS and western blot analyses

FACS analyses were used to determine the expression levels of human CD16 or CD32 in Jurkat-NFAT-hCD16 or Jurkat-NFAT-hCD32 cells. In brief, cells were incubated with PE/Cyanine7 anti-human CD16 antibody (BioLegend, 302016, for Jurkat-NFAT-CD16 cells) or FITC CD32/FCGR2/Fc gamma RIIb antibody (Sino Biological, 10374-MM01-F, for Jurkat-NFAT-hCD32) for 30 min at 4 °C. After twice washes with PBS buffer, stained cells were subjected for FACS analyses using LSRFortessaX-20 (BD Biosciences). The cell lysates of 239 F, SFLwt, and SFLfk (1 × 10^6 cells) were prepared and subjected to western blots using HRP-conjugated mouse antibodies specific to the S1 subunit and S2 subunit (developed in our lab).

2.6. ADCC and ADCP assays

Antibodies or plasmas were pre-diluted in a reaction medium of IMDM medium containing 10% heat-inactivated FBS (Gibco, heating to 56 °C for 30 min). Aliquots (20 μL) of pre-diluted antibodies or plasmas were incubated with of SFLwt or SFLfk target cells (2 × 10^6 cells in 90 μL reaction medium) in 96-well plates for 1-hour at 37 °C in 5% CO2 incubator. Subsequently, Jurkat-NFAT-hCD16 or Jurkat-NFAT-hCD32 effector cells (2 × 10^5 cells in 90 μL) were added into the mixtures for further incubation at 37 °C in 5% CO2 incubator for 18-hour. Finally, 20 μL supernatants from co-cultures were mixed with 50 μL pre-diluted Coelenterazine substrate (Topscience, T6808, 10 μM, 1:4000 diluted with Opti-MEM medium) in a 96-well plate for immediate bioluminescence measurements using a multimode microplate reader (Spark 20 M, Tecan). The luciferase (Luc) induction fold (ADCC or ADCP Luc induction fold) was calculated by the fold change of relative light unit (RLU) values of the test well to control well (mock buffer reaction with target and effector cells).
2.7. Quantitative and statistical analyses

The four-parameter logistic (4-PL) model was used for fitting dose-dependent responses (ADCC or ADCP Luc induction fold) of antibodies or human plasma. The cut-off value for a positive response in ADCC or ADCP assay was defined as the Luc induction fold ≥ 2. For quantitative analyses, the parameter of EC₅₀ (for mAbs) or ED₅₀ (for plasma or serum samples), which were defined as the minimal concentration (EC₂) or the maximal dilution fold (ED₂), to achieve a positive reaction (Luc induction fold=2), was calculated following the 4-PL fitting equation. The Gaussian function was used for frequency distribution determination. Linear regression models and the Pearson coefficient were used for correlation analyses. All statistical tests and data visualization were conducted using GraphPad Prism (version 9.3.1).

3. Results

3.1. Characterizations of cells for ADCC and ADCP assays

As the presence of the furin site, the SARS-CoV-2 spike may be cleaved by endogenous furin-like enzymes and result in the disassociation of S1 and S2 subunits. We designed two constructs, the SFLwk and SFLfk (Fig. 1 A), to generate cell lines expressing SARS-CoV-2 spike proteins as target cells. Both constructs contain full-length (aa 1–1273) SARS-CoV-2 spike protein-expressing cassettes. Being different from the SFLwt, the furin site was mutated in the SFLfk. Two cell lines expressing spike proteins were developed via stable transfections of the SFLwt and SFLft into 293 F cells. FACS analyses (Fig. 1 B), for either staining with S1 or S2 antibodies, demonstrated the successful expressions of spike proteins on the two cell lines. Both cell lines presented an over 95% positive rate, as evidenced by the integrated mRuby3 expression and S2 proteins on the two cell lines. Moreover, western blots revealed furin-cleaved spike derivates (furin-cleaved S1 and S2 subunits) were presented in SFLwt cell lines but not in SFLfk cell lines. In contrast, the intact spike protein levels appeared to be higher in SFLfk than that in SFLwt cell lines. These results supported the two cell lines could be used as target cells in ADCC or ADCP assays. For effectors, two Jurkat-derived NFAT reporter cells, Jurkat-NFAT-hCD16 and Jurkat-NFAT-hCD32, were used for ADCC and ADCP assays, respectively. The expressions of hCD16 or hCD32 receptors were evidenced by FACS analyses (Fig. 1 D).

3.2. Optimizations of experimental conditions for ADCC and ADCP assays

To explore the optimized experimental conditions, we used an RBD-targeting mAb h83H7 as a proof-concept positive control to test the effector cell density, the E:T (cell number ratio of effectors to targets) ratio, and the reaction time. The ADCC Luc fold inductions of 3-fold serial dilutions of h83H7 were tested with Jurkat-NFAT-hCD16 at various cell densities (5 × 10⁴ cells/well, 1 × 10⁵ cells/well, or 2 × 10⁵ cells/well) in combination with SFLfk cells at different E:T ratios (1:1, 2:1, or 4:1). As shown in Fig. 2 A, Jurkat-NFAT-hCD16 using at 2 × 10⁵ cells/well with the same density of target cell (1:1) resulted in higher Luc induction folds. At this condition, the h83H7 mAb generated a dose-dependent response with an upper plateau of 10–14 Luc induction folds when detected at 18-hour or later (Fig. 2 B). In contrast, little or even no response was observed for an irrelevant control mAb (an HBsAg specific humanized mAb). In the same condition, h83H7 also exhibited dose-dependent ADCP activity in the Jurkat-NFAT-hCD32/SFLfk reporter system, though the effects appeared to be weaker than that in the ADCC test (Fig. 1 C). To shorten the turnaround time, bioluminescence detections in further experiments of this study were performed at 18-hour after co-cultures of the effector and target cells.

3.3. Evaluations for the ADCC and ADCP activities of representative mAbs

Besides the h83H7, additional nine mAbs were further tested for

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Fig. 1. Characterizations of the target and effector cells for ADCC and ADCP assays against SARS-CoV-2 spike protein, (A). Schematics of constructs for generations of SFLwt and SFLfk cells. The SFLwt contains a codon-optimized gene encoding a full-length SARS-CoV-2 spike (aa 1–1273, identical to the spike protein derived from MN908947.3). The only difference between the SFLwt and SFLfk was the furin site is mutated (RRAR changed into GSAS). (B). Flow cytometry analyses of Exp293F (293 F) cells with stable expressions of SFLwt or SFLfk by antibodies specific to the S1 subunit (left panel), S2 subunit (middle panel), or mRuby3 selection marker (right panel). (C). Western blots for SARS-CoV-2 spike proteins in cell lysates of SFLwt and SFLfk cells. S, intact spike protein; S1, furin-cleaved S1 subunit; S2, furin-cleaved S2 subunit. (D). Flow cytometry analyses for expressions of hCD16 and hCD32 in Jurkat-NFAT-hCD16 (left panel) and Jurkat-NFAT-hCD32 (right panel) effector cells, respectively.
their activities in ADCC and ADCP assays. Among these mAbs, the 4 A8 and NTD4–8 are NTD-specific mAbs. The S2P6 recognizes the stem helix in the C-terminus (aa 1148–1152) of the S2 subunit. The remaining mAbs, including REGN10933, BD368–2, COVA2–15, h36H6, h85F7, and S2M11, are all RBD-targeting mAbs. All of these mAbs showed neutralization activities in previous studies. In ADCC and ADCP tests, most mAbs presented detectable and dose-dependent activities targeting either the SFLwt or SFLfk cells (Fig. 3). To quantitative assessments of neutralization activities in previous studies. In ADCC and ADCP tests, most mAbs presented detectable and dose-dependent activities targeting either the SFLwt or SFLfk cells (Fig. 3). To quantitative assessments of their potencies, the minimal concentrations to achieve 2-fold RLU increase of these mAbs were calculated using the 4-PL fitting curves. Overall, the EC2 values of these mAbs inversely correlated with their maximal Luc induction folds (values of the upper plateaus in fitting curves) in either ADCC (R2 = 0.448, p = 0.002) or ADCP (R2 = 0.230, p = 0.044) assays (Fig. S1). However, some prozone effects were noted, particularly for the ADCP assay (Fig. 3B). Therefore, we considered the EC2 as the preferred parameter for quantitative comparisons of the potencies of mAbs in the two assays. In ADCC tests, the 4A8, BD238–2, COVA2–15, h36H6, h85F7, and S2M11 showed comparable potencies (<4-fold difference) in targeting the SFLwt and SFLfk cells. However, the NTD4–8, 2P6, and REGN10933 mAbs resulted in markedly higher activities when using the SFLwt as the target cells. In ADCP tests, the 4A8, NTD4–8, COVA2–15, h85F7, and S2M11 showed similar potencies (<4-fold difference) in targeting the SFLwt and SFLfk cells. However, the h36H6, BD368–2, 2P6, and REGN10933 mAbs showed higher activities (>4-fold difference) when using the SFLwt as the target cells. These data suggested the potential differences of some mAbs in mediating FcyR activation when targeting the furin-cleaved and uncleaved SARS-CoV-2 spikes, which may be attributed to their epitope difference. As the SFLwt cell line expressed the natural form of the SARS-CoV-2 spike and presented a better performance for reporter assays in mAbs tests, we selected it as the target cell in subsequent studies.

3.4. Measurements of ADCC and ADCP activities in human convalescent plasmas

We further investigated the performance of two assays in detecting ADCC and ADCP activities mediated by polyclonal antibodies acquired from natural infections. First, we evaluated the specificities of the assays using 190 plasma samples collected from healthy individuals in the pre-pandemic era. As shown in Fig. 4A, 1 and 5 samples were positive (resulted in >2 Luc induction fold, at 1:4 dilution) in ADCC and ADCP tests, respectively. The specificity was 99.5% (95%CI: 98.4–100%) for the ADCC assay and was 97.4% (95%CI: 95.1–99.6%) for the ADCP assay. Of all 87 COVID-19 HCPs we tested, 83 and 81 samples showed positive reactions in ADCC and ADCP tests (at 1:4 dilution). The sensitivities were 95.4% (95%CI: 91.0–99.3%) and 93.1% (95%CI: 87.8–94.8%) for ADCC and ADCP assays, respectively. Detailedly, the sensitivities of the ADCC and ADCP assays were 97.1% (68/70) and 95.7% (67/70) among samples collected in the early convalescent phase; and were 88.2% (15/17) and 82.4% (14/17) among samples collected in the later convalescent phase.

We further assessed the ADCC and ADCP antibody titers (ED2) of these HCPs via serial dilution measurements of each sample (Fig. S2, from 1:8–1:17496). These HCPs presented ADCC antibody titers ranging from 7.4 to 1721.6 (for 83 ADCC positive samples) and ADCP titers ranging from 4 to 523.2 (for 81 ADCP positive samples). Overall, the ADCC (Fig. 4C, R2 = 0.819) and ADCP (Fig. 4D, R2 = 0.792) antibody titers of these HCPs positively correlated with their Luc induction fold
measure at 1:8 dilution. Moreover, the antibody titers determined in the ADCC, ADCP, and PsV neutralization assays were significantly correlated (Fig. 4E to G).

4. Discussions

Antibodies against SARS-CoV-2 spike acquired from natural infections or vaccine immunization play antiviral roles through multiple
functions, including the Fab-mediated neutralization and the Fc-mediated immune opsonization. Most previous studies focused on the neutralization functions of spike antibodies, which are essentially required for protection against COVID-19. Beyond neutralization, in vivo therapeutic studies revealed the Fc-mediated effector functions are also critical for viral control, as evidenced that the LALA Fc loss-function mutations significantly diminished the efficacity of spike antibodies to protect animals from lethal SARS-CoV-2 challenges (Ullah et al., 2021; Winkler et al., 2021). By contrast, the GAALIE mutated S309 mAb (VIR-7832) with enhanced Fc-effector function showed improved clearance of SARS-CoV-2 infected cells linking to boost T-cell immunity (Cathcart et al., 2022). Several studies revealed the presence of ADCC and ADCP activities of anti-spike antibodies in plasmas of COVID-19 patients (Adeniji et al., 2021; Natarajan et al., 2021; Tso et al., 2021), and inadequate ADCC or ADCP response appeared to be associated with severe disease and mortality (Bahnan et al., 2021; Brunet-Ratnasingham

Fig. 4. Quantitative measurements of ADCC and ADCP activities of human plasma samples. Distributions of bioluminescence inductions of 190 samples (at 1:4 dilution) from pre-COVID-19 healthy populations in ADCC (A) and ADCP (B) assays. (C) Correlation analyses of the ADCC (C) or ADCP (D) titers (ED$_2$, maximum dilution fold to achieve ≥2-fold induction determined by the 4-PL fitting curve) and bioluminescence induction folds (at 1:8 dilution) in detecting 87 HCP samples from COVID-19 patients. (E) Comparison of the ADCC and ADCP titers (ED$_2$) of 87 HCPs. Correlations between the titers (Y-axis) of ADCC (F) or ADCP (G) of 87 HCPs with their neutralizing antibody titer (X-axis).
Moreover, several ADCC assays for SARS-CoV-2 antibodies were established based on the determinations of targeted cell loss using NK cell lines or PBMC (Beaudoin-Bussières et al., 2021; Chen et al., 2021; Garvin et al., 2021). However, due to the lack of robust assays for quantitative assessments, current understandings regarding antibodies’ ADCC and ADCP activities remain limited, particularly for their role in linkage with COVID-19 disease and vaccine protection efficacy.

The classical method for accurate ADCC quantification is the chromium-51 ($^{51}$Cr) release assay, requiring specialized equipment and radioactive materials (Brier et al., 1975; Nelson et al., 2001). Besides, several non-radioactive methods were also developed based on the measurements of cell deaths of the target cells via released lactate dehydrogenase or fluorescent stains with various dyes (calcine-AM, BCECF, CSFE, or EuTFA) (Alrubayyi et al., 2018; Broussas et al., 2013; Gillissen et al., 2016; Gómez-Román et al., 2006; Kolber et al., 1988; Neri et al., 2001; Patel and Boyd, 1995; Roden et al., 1999). The pH-sensitive dyes, which preferentially accumulate in acidic organelles such as phagosomes and present pH-dependent fluorescence (Kamen et al., 2019; Mikay et al., 2009), are commonly used in ADCC assays. However, these methods require flow cytometers or high-resolution microscopes that limit their usage. The NFAT reporter system is an alternative approach to characterize the Fc-effector functions of antibodies via measurements on NFAT pathway activation, which is the downstream action of Fc engagement of Fc gamma receptors in both ADCC and ADCP processes. Here, we developed robust and convenient ADCC and ADCP assays using well-characterized target cells that stably expressed SARS-CoV-2 spike and Jurkat-NFAT-CD16a/CD32a effector cells. We also establish methods and indexes for quantitatively analyzing the potencies of mAbs (Fig. 3) as well as the ADCC and ADCP antibody titers in human plasmas (Fig. 4). As the targets for assays, the SFLwt cells exhibited better performance than SFLk in some tested mAbs though there was no marked difference noted in spike expressing levels for the two cells. It is possible that partial furin cleavage of the spike trimer may facilitate the engagement of IgG-Fc and Fc receptors. On the other hand, although all test mAbs were expressed with the same human-IgG1 backbone, their activities in ADCC and ADCP assays appeared to be distinctly varied, possibly attributed to the affinity difference or epitope diversity.

The two assays developed in our study can also be used in detecting the antibody titers of ADCC and ADCP in human plasmas. Our data demonstrated the specificities of the assays in SARS-CoV-2 native populations (Fig. 4 A). Most of the HCP samples from recovered COVID-19 patients presented detectable antibodies with ADCC and ADCP activities, showing titers correlating with that determined in neutralization tests (Fig. 4). As the Ab level is the well-documented immune marker inversely associated with COVID-19 risk and directly associated with vaccine efficacy (Garcia-Beltran et al., 2021; Khoury et al., 2021), it is reasonable to assume the ADCC and/or ADCP antibody titers may also serve as surrogates of anti-SARS-CoV-2 immunity. However, further studies using the two assays to explore the significance and clinical value in vaccinated populations are certainly required.

In summary, our study developed cell-based reporter assays to evaluate the ADCC and ADCP activities of the mAbs and human blood samples against the SARS-CoV-2 spike. The good performance of the two assays as documented by our data supports their further applications both in basic and clinical studies.

**CRediT authorship contribution statement**

**Yuting Hong:** Investigation, Methodology, Data curation, Formal analysis, Writing – original draft. **Huilin Guo:** Investigation, Methodology, Data curation, Validation. **Min Wei:** Investigation, Methodology. **Yali Zhang:** Methodology, Validation. **Mujin Fang:** Visualization. **Tong Cheng:** Writing – review & editing. **Zhiyong Li:** Resources. **Shengxiang Ge:** Formal analysis, Supervision. **Xiangyang Yao:** Resources, Conceptualization, Writing – review & editing, Funding acquisition. **Quan Yuan:** Conceptualization, Project administration, Writing – review & editing, Funding acquisition. **Ningshao Xia:** Supervision, Writing – review & editing.

**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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**Data Availability**

Data will be made available on request.

**Appendix A. Supporting information**

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.jviromet.2022.114564.

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