Avian metapneumovirus: A five-plex digital droplet RT-PCR method for identification of subgroups A, B, C, and D

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End-point and real-time avian metapneumovirus (AMPV) RT-PCRs have been developed to detect one or two of the four recognized subgroups (A, B, C, and D) simultaneously or for broad range AMPV detection. Current subgroup specific tests target variable areas of the genome which makes these PCRs sensitive to specificity defects as recently documented. In the current study, a single five-plex digital droplet RT-PCR targeting the conserved viral polymerase gene of AMPV, which is less prone to genetic drift, has been designed. This digital droplet RT-PCR was capable of identifying each of the four AMPV subgroups. Each subgroup was identified according to a specifically assigned fluorescent amplitude. Specificity, which was tested including three AMPV strains, non-AMPV avian viruses and closely related human respiratory viruses, was 100%. The specific limit of detection for extracted viral RNA was estimated between 1 and 3 copies/µL. This tool simplifies the number of tests required for AMPV genotype diagnostics and should be theoretically less effected by viral genome evolution due to its target region. Ultimately, application of this test will contribute to an improved understanding of the global geographic distribution and subgroup host range of field strains.

KEYWORDS
avian metapneumovirus, digital droplet, identification, five-plex, method

Introduction

Since their first detection in South Africa in the late 1970s, avian metapneumoviruses (AMPV), family Pneumoviridae, genus Metapneumovirus (1), have now been detected in most regions of the world (2). AMPV fundamentally cause respiratory and reproductive diseases in poultry and have had serious economic impact on the poultry industry. To date, based on genetic and antigenic differences, four subgroups are described (AMPV-A, B, C, and D) (3–8) and two genetic lineages of AMPV-C (8) have been identified. Recently, two divergent AMPVs from a monk parakeet (9) and a gull (10) were described and do not cluster genetically with any of the four previously described subgroups. The principal host species for AMPVs are turkeys, chickens and ducks, however, they have
also been isolated from pheasants (11), and been detected in guinea fowls (12), or wild water fowls (13) by serological and molecular methods, respectively.

Concerning field detection of AMPVs in the three principal poultry species (turkeys, chickens and ducks), all subgroups have been detected in turkeys (14–17), although to date, the subgroup C viruses detected in this species have been of a lineage unique to North America (US AMPV-C). In domestic Muscovy and Pekin ducks, only AMPV-Cs of a Eurasian lineage have been reported (18, 19). In chickens, AMPV A, B and an AMPV-C have been described (20–24), however the lineage of the latter is yet to be defined. Recently, details on the extent of infection of the four recognized subgroups and lineages of AMPV in turkeys, chickens and Muscovy ducks under controlled experimental conditions have been published (25). Interestingly, apart from confirming the subgroup / host relationships described above, this study showed that AMPV-C viruses of both US and Eurasian lineage could be re-isolated from chickens, and AMPV-C viruses of Eurasian lineage from turkeys despite a lack of viral RNA detection using a well-characterized real-time RT-PCR (26) targeting the hypervariable SH gene. These results demonstrate that replication of AMPVs in a “non-conventional host” may modify the viral consensus sequence, especially in hypervariable regions of the genome, and thus go undetected by molecular tests, such as those currently available, that target these regions.

To date, both classical and real-time RT-PCRs have been developed for either broad range (3, 27) or subgroup specific detection. Concerning real-time RT-PCR tests that are subgroup specific, tests exist in duplex for subgroups A and B (26, 28, 29), in triplex for subgroup C (both lineages) and the new identified parakeet and gull AMPVs (30) and in individual tests for subgroups C and D (26). All these current subgroup specific molecular tests are based in the hypervariable G or SH genes except for that detecting the AMPV gull virus which is in the polymerase L gene.

In this context, the development of a multiplex reaction PCR fully based on a non-hypervariable region of the genome that is capable of detecting and differentiating the four recognized subgroups found in poultry would be beneficial. Theoretically, this could be addressed using recent real-time PCR machines that now have the capability of reading up to four different fluorescent wavelengths. In reality however, real-time PCRs that differentiate more than three targets are rare due to the level of competition between primers, probes and target molecules within the reaction mixture. The technology of droplet digital PCR (ddPCR), based on the amplification of target molecules of the same mixture but distributed into thousands of reaction droplets, reduces this competition and results in an improved capacity for multiplexing (31). ddPCR is based on end-point PCR reactions and uses droplet reader machines capable of reading several fluorescent wave lengths. Different positive targets can be separated into specific clusters using probes labeled with different fluorescent dyes but also by using probes at different concentrations that are labeled with the same dye. The method of generating specific clusters by varying probe concentrations is based on the fact that the fluorescent amplitude value is directly linked to the number of probe molecules in the PCR reaction mixture. Combining these two techniques means that many specific clusters can be obtained even with a two-color reader.

The current study describes the development of a five-plex RT-ddPCR for the identification of AMPV-A, B, C or D and the housekeeping gene Glyceraldehyde-3-phosphate dehydrogenase (GAPDH). For the amplification of AMPV molecules, two common primers and four subgroup specific probes were used which targeted a small region of the conserved viral polymerase gene. Distinction between subgroups was achieved on the principle of varying probe concentrations.

Materials and methods

Ethic statements

Tracheal swab samples were taken during previous animal experiments performed at Anses Ploufragan (25). These animal experiments were approved by Anses Ploufragan ethical committee (committee number C2EA-016/ ComEth ANSES/ENV/AUPEC) and authorized by French Ministry for higher education and research (application number 14-024, statement number 08/04/14-7). Chickens were raised and humanely euthanized in agreement with EU directive number 2010/63/UE.

Viruses

Twenty-eight AMPV field strains that had been isolated on Vero cells as previously described (32), three AMPV vaccine strains, 35 AMPV positive tracheal swab samples from previously performed experimental trials in birds (27) and 35 non-avian metapneumovirus strains were used as test samples. The subgroup assignment of AMPVs had previously been determined using existing well-characterized diagnostic methods (26) and Sanger sequencing. A reference virus of each subgroup was also tested (AMPV-A 85051, AMPV-B 86004, AMPV-C 99178, and AMPV-D 85035) for which the...
TABLE 1 Details of the five-plex RT-ddPCR AMPV primers and probes.

| Name             | Target                             | Sequence 5′-3′ | L ORF position | Amplicon size |
|------------------|------------------------------------|----------------|----------------|---------------|
| PanAMPV L1Fwd    | AMPV-A, B, C, and D                | CACAGAGTCTATTYTGCTGG | 1,980–2,001    | 177 pb        |
| PanAMPV L1Rev    | AMPV-A, B, C, and D                | TCCACACCTTTTGGRCACCA | 2,140–2,158    |               |
| AMPVA L1Pro      | AMPV A                             | FAM-TCTGCGTCTA-ZEN-TGTGCTAGATTCCACC-IABkFQ | 2,065–2,089 |               |
| AMPVB L1Pro      | AMPV B                             | FAM-CTATAACAC-ZEN-CCATCTGCTGACGGATGATIABkFQ | 2,086–2,114 |               |
| AMPVC L1Pro      | AMPV C                             | HEX-AGACATGCC-ZEN-CCTCCAGAAACAGAAGGAGT-IABkFQ | 2,044–2,072 |               |
| AMPVD L1Pro      | AMPV D                             | FAM-CTGCTCAGG-ZEN-TATGTGTGCTATGTGCAATTC-IABkFQ | 2,071–2,100 |               |
| PanGAPDH Fwd     | GAPDH                              | TGAGTAGTAGTTAGGATCCACT | 198 pb |               |
| PanGAPDH Rev     | GAPDH                              | GCCAGGCGAGTTGGTGTCG |               |               |
| PanGAPDH Pro     | GAPDH                              | HEX-TGAGCGCCA-ZEN-GCCTTCTCCATGG-AIBkFQ |               |               |

*AMPV reference strain DQ666911.
*AMPV reference strain AB548428.
*AMPV reference strain HG934338.
*AMPV reference strain HG934339.

Primer design

Based on a nucleotide alignment using MEGA software version 7 (34) of all available AMPV full length sequences on the database, a zone in the L gene (position 1980–2140) that allowed for the design of one single forward and one single reverse primer that were common to all subgroups and for the design of subgroups specific probes was selected (Table 1). Primers and probes were evaluated in silico using OligoAnalyzer Tool (https://eu.idtdna.com/pages/tools/oligoanalyzer) for secondary structures and self- and heterodimers. Specificity was investigated using Primer-BLAST tool. Probes for AMPV-A, B and D used 5′-6-carboxyfluorescein (FAM) and probes for AMPV-C and an endogenous reference gene, the house keeping gene GAPDH (Table 1) used 5′-hexachloro-fluorescein phosphoramidite (HEX).

Digital droplet RT-PCR

The RT-ddPCR platform used enables the detection of two fluorophores: FAM in channel 1 (CH1) and HEX in channel 2 (CH2). Based on implementation of varying probe concentrations, a five-plex reaction was possible by assigning AMPV-A, B and D AMPV targets to CH1 and GAPDH to CH2. RT-ddPCR assays were performed in a 20 µl final volume in RNase free water using the One-step RT-ddPCR Advanced Kit for Probe (Bio-Rad, France, reference 1864022), containing 5 µl of Supermix, 2 µl of Reverse transcriptase, 1 µl of 300 mM DTT, 1 µl of RNA template and primers at final concentrations of 900 nM. For CH1, 300 mM of the AMPV-A probe, 200 mM of the AMPV-B probe, and 100 mM of the AMPV-D probe was used per

Viral RNA extraction

Viral RNA was extracted from all of the above samples as described previously (27) using QIAamp Viral RNA Mini Kit (Qiagen, France, reference 52906).
TABLE 2: Results of the five-plex RT-ddPCR AMPV with viruses used for specificity assessments.

| Family            | Genus                | Species                      | Virus strains | Positive/Total |
|-------------------|----------------------|------------------------------|---------------|----------------|
| Pneumoviridae     | Metapneumovirus      | Avian metapneumovirus        | AMPV A (T, V) | 10+/10         |
|                   |                      |                              | AMPV B (T, C, V) | 12+/12         |
|                   |                      |                              | AMPV C (T, D) | 7+/7           |
|                   |                      |                              | AMPV D (T) | 2+/2           |
|                   | Human metapneumovirus|                              | HMPV A1, A2, B1, B2 | 0+/4           |
|                   | Human respiratory virus|                             | RSV A<sub>H</sub>, A2, B1 | 0+/3           |
| Orthomyxoviridae  | Alphainfluenzavirus  |                              | H1N1, H5N2, H5N3, H7N1, H7N3 | 0+/7           |
| Paramyxoviridae   | Newcastle disease virus|                             | PMV1 | 0+/2           |
|                   | Avian paramyxovirus |                              | PMV2-4, 6-9, 11 | 0+/8           |
| Adenoviridae      | Aviadenovirus        |                              | Fowl adenovirus type 1 | 0+/1           |
|                   | Suadenovirus         |                              | VEH | 0+/1           |
| Birnaviridae      | Avibirnavirus        |                              | IBDV serotype 1.2 | 0+/2           |
| Picornaviridae    | Tremovirus           |                              | Avian encephalomyelitis virus | 0+/1           |
| Reoviridae        | Orthoreovirus        |                              | Avian reovirus | 0+/1           |
| Coronaviridae     | Gamma coronavirus    |                              | IBV, TCoV | 0+/4           |
| Herpesvirida      | Iltovirus            | Gallid alphaherpesvirus 1     | ILTV | 0+/1           |

T, turkeys; C, chickens; D, ducks; V, vaccine.

reaction. For CH2, the concentration of probes was 300 mM for AMPV-C and 100 mM for GAPDH. The reaction mixture was then loaded into the well of an eight-channel disposable droplet generator cartridge and water-oil emulsified into 20,000 droplets with QX200 Droplet Generator (Bio-Rad, France). After emulsion, the droplets were transferred to a 96-well PCR plate for the RT-ddPCR reaction (C1000 Touch Thermal Cycler, Bio-Rad, France). The cycling conditions were as follows: 50°C for 60 min (RT reaction), 95°C for 10 min, followed by 40 cycles of 95°C for 30 s, 54°C for 30 sec, and 60°C for 60 s. A final cycle of 98°C for 10 min was performed followed by a hold temperature of 12°C. Reactions were read in a QX200 Droplet Reader (Bio-Rad, France). The results were analyzed by QuantaSoft software (version 1.7.4 Bio-Rad, France).

Data analysis

Droplet clusters were evaluated in 2D plots as shown in Figure 1 using QuantaSoft software version 1.7.4 Bio-Rad, France). A cut-off amplitude threshold, set at ~6,000 for CH1 and 3,200 for CH2, distinguished positive droplets (one or more than one copy of the target) from negative droplets (devoid of target molecules). Ten or more positive droplets was considered as a positive result. The mean amplitude values for positive clusters in CH1 and CH2 for test samples were compared with those of AMPV reference strains to assign the AMPV subgroup. An absolute quantification of the clusters was performed.

Probe interference test

Each probe used collectively in the five-plex RT-ddPCR was tested individually in its own simplex test. AMPV-B, C, and D were tested at 10^4-10^2 TCID<sub>50</sub>/ml and AMPV-A at 10^4-10^1 TCID<sub>50</sub>/ml. Composition of individual reactions and cycling conditions were identical to those described for the five-plex. The ratio of extracted viral RNA log<sub>10</sub> copies/µl obtained by simplex RT-ddPCR compared to five-plex RT-ddPCR was evaluated. Comparison of performance of multiplex and simplex RT-PCR reactions are not documented by published guidelines but a bias above 25% has previously been reported to be significant (35). Thus, a result below 25% (absolute value) between the simplex and five-plex reactions for each subgroup would signify no interference between probes when combined into one single reaction. A Pearson’s correlation coefficient was used to determine the correlation between the methods along the range.

Specificity

The specificity of the method was evaluated by testing the 28 AMPV field strains, the three attenuated vaccine strains, the 35 AMPV-positive tracheal swabs, four human metapneumoviruses (HMPV), three respiratory syncytial virus (RSV) and various avian viruses of others families. Details of these are shown in section “Viruses” and in Table 2. For AMPV samples, the mean and standard deviation (SD) of
TABLE 3  Comparison between five-plex and simplex RT-ddPCR.

| Viral titer<sup>a</sup> | AMPV-A | AMPV-B | AMPV-C | AMPV-D |
|------------------------|--------|--------|--------|--------|
|                        | Five-plex<sup>b</sup> | Simplex<sup>b</sup> | Bias   | Five-plex<sup>b</sup> | Simplex<sup>b</sup> | Bias   | Five-plex<sup>b</sup> | Simplex<sup>b</sup> | Bias   |
| 4                      | 3.27   | 3.46   | 5.49   | 2.87   | 3.10   | 7.42   | 2.79   | 2.85   | 2.11   | 2.57   | 2.90   | 11.38 |
| 3                      | 2.27   | 2.48   | 8.47   | 1.84   | 2.03   | 9.36   | 1.86   | 1.89   | 1.59   | 1.60   | 1.89   | 15.34 |
| 2                      | 1.24   | 1.46   | 15.07  | 0.91   | 1.12   | 18.75  | 0.89   | 0.85   | −5.72  | 0.61   | 0.92   | 33.70 |
| 1                      | 0.44   | 0.41   | −7.32  |        |        |        |        |        |        |        |        |        |

<sup>a</sup> Log<sub>10</sub>(TCID<sub>50</sub>/ml).

<sup>b</sup> Log<sub>10</sub> (absolute copies/µl of extracted viral RNA), Bias expressed in % [calculation: 100–(five-plex result/Simplex result) * 100].

the amplitude values in the CH1 and CH2 channels were calculated. Non-parametric Kruskal-Wallis test was applied to confirm statistical differences between the amplitude values of the different subgroups.

**Sensitivity and linearity**

A dilution series was prepared for AMPV reference viruses by diluting viral suspension in a supernatant of specific pathogen-free (SPF) turkey tracheal swabs. Five replicates of the dilution series were performed (two independent assays, three repeats for the first assay and two repeats for the second). The determination of the limit of detection (LoD) and the limit of quantification (LoQ) were based on these samples. The LoD was the last dilution in which all replicates were positive and the LoQ was the last dilution in which all replicates showed a coefficient of variation (CV) in inter-assays of less than 25%. In ddPCR, the absolute quantification can be estimated from the number of positive droplets to the total number of droplets and their known volume with an optimal precision of concentration at 1.6 molecules per droplet.

To avoid saturation of positive droplets by target viral RNA molecules, a range from 10<sup>4</sup> to 10<sup>1</sup> TCID<sub>50</sub>/ml was selected to evaluate the linearity of the method. 2D plots from absolute quantification (log<sub>10</sub> viral RNA copies/µl) against the viral titer (log<sub>10</sub> TCID<sub>50</sub>/ml) for the four reference viruses were generated and the coefficient of determination (R<sup>2</sup>) were calculated for each subgroup.

**Results**

**Probe interference test**

Assays were done to compare the performance of probes in simplex and five-plex reactions by determining the absolute quantification of extracted viral RNA (log<sub>10</sub> copies/µl) from reference viruses of each subgroup. All biases between the two methods were ≤25% (absolute values) except for AMPV-D at a viral titer of 10<sup>2</sup> TCID<sub>50</sub>/ml (Table 3). To determine the correlation of the two methods, a Person’s correlation coefficient for dynamic range was calculated and was equal to 0.995 (p-value <0.0001). Thus, significant correlation was observed between the simplex and the five-plex tests.
### Specificity

The specificity of the five-plex RT-ddPCR assay was calculated to be 100% based on 31 AMPVs representing the four recognized subgroups of AMPV (Table 2). All the strains were correctly assigned to their specific subgroup based on their amplitude values. For CH1, a mean amplitude value of 15,312 was obtained for subgroup A (standard deviation (SD) 677, \(N = 10\)), 11,681 for subgroup B (SD 777, \(N = 12\)), and 8,482 for subgroup D (SD 30, \(N = 2\)) (Supplementary Table S1). For CH2, mean amplitude values were 4,114 for GAPDH (SD 116, \(N = 23\)) and 5,649 for subgroup AMPV C’s of both lineages (SD 171, \(N = 7\)) (Supplementary Table S1). Results of Kruskal-Wallis test showed significant differences between subgroup clusters (p-value \(<0.0001\)) for CH1 and CH2 amplitude (Figure 2). Probes were specific to their subgroup and did not cross-react. No droplet was observed in non-specific clusters. No AMPV-positive results were obtained from the 35 non-AMPV viruses, including the closely related HMPV and RSV (Table 2). Concerning the AMPV-positive tracheal swab samples, all were detected and assigned to their expected subgroup. Additionally, six tracheal swab samples from non-infected SPI birds (turkeys, ducks and chickens, two samples per species) were screened and were only GAPDH-positives (Table 4). All test samples were also positive for the GAPDH.

### Sensitivity and linearity

The estimation of the LoD was the same for the four subgroups and was equal to \(10^1\) TCID\(_{50}\)/ml, corresponding to one to three copies/\(\mu\)l of extracted viral RNA (Table 5).

For ddPCR, the number of target molecules per droplet is important for accurate calculation of the concentration of initial target molecules by Poisson correction statistics. A range from \(10^3\) to \(10^1\) TCID\(_{50}\)/ml was selected to evaluate the linearity of the method. A 2D plot of viral titer (log\(_{10}\) TCID\(_{50}\)/ml) against absolute quantification (log\(_{10}\) copies/\(\mu\)l) was generated for the four reference viruses. The \(R^2\) of the linear regression model showed a linear correlation from \(10^4\) to \(10^1\) TCID\(_{50}\)/ml for subgroup A (\(R^2 = 0.9971\)) and from \(10^4\) to \(10^2\) TCID\(_{50}\)/ml for subgroups B, C and D (\(R^2 = 0.9990, 0.9999,\) and 1, respectively) (Supplementary Figure S1).

These data were correlated with the sensitivity results showing a decrease of precision when the mean number of positive droplets was very low (Table 3). The range between \(10^2\) and \(10^1\) TCID\(_{50}\)/ml for subgroups B, C, and D was detectable but not quantifiable due to a high percentage of variation in the results.

### Discussion

Multiplex PCRs are difficult to develop mainly because of the competition between the molecules in the same reaction mixture, however, when one PCR reaction mixture containing many different molecules can be separated into thousands of individual reactions (droplets in the case of ddPCR), then competition is drastically reduced without losing the multiplexing capabilities of the reaction as a whole. Here, a five-plex RT-ddPCR for the detection and differentiation of any one of the four recognized AMPV subgroups has been achieved using an amplitude-based multiplexing approach with double-quenched probes at different final concentrations coupled with the same dye (31) to achieve sufficient cluster separation for clear identification of each subgroup. Amplitude reads on CH1 corresponding to FAM labeled probes were more extensive on the value scale than those of HEX, and for this reason, three targets (AMPV-A, B and D) were separated using this fluorochrome. Only two targets were used with HEX (AMPV-C and GAPDH). Ten clearly defined clusters were obtained: one for each AMPV subgroup (four in total), one for each AMPV subgroup in association with GAPDH (four in total), one for GAPDH only and a final cluster for negative droplets absent of target molecules (Figure 1).

Prior to the current study, all but one subgroup specific AMPV PCR have targeted open reading frames (ORFs) of high nucleotide variability between subgroups such as the SH or G ORFs (26, 28–30, 36) because sequences specific to each subgroup in these regions are in abundance. However, the problem with this approach is that hypervariable sequences are subject to frequent nucleotide change implicating potential mismatch with primers and/or probes, as has been previously evoked (25). The current RT-ddPCR has been therefore developed using a low-variability region in the L ORF (4, 34) located between nucleotide positions 5′ 1,980 and 5′ 2,158 to overcome the above-mentioned limitations. Other conserved ORFs in the AMPV genome, such as that coding for the nucleocapsid protein (N), which are more abundant as viral RNA molecules in

### Table 4: Detection of AMPV in tracheal swabs from previous experimental trials.

| Virus | Species | Turkey | Duck | Chicken |
|-------|---------|--------|------|---------|
| AMPV-A | 5/5 | / | 5/5 |
| AMPV-B | 10/10 | / | / |
| AMPV-C | 2/2 | 3/3 | / |
| AMPV-D | 10/10 | / | / |
| Negative | 0/2 | 0/2 | 0/2 |

Results expression: positives number detected/total number screened; /, not available.
TABLE 5 Limits of detection and quantification.

| Sub-group | Viral titer<sup>a</sup> | Assay 1<sup>b</sup> | Assay 2<sup>b</sup> | Mean +/- SD (CV) | Copies/µl | R² |
|-----------|-------------------------|---------------------|---------------------|-----------------|-----------|----|
|           | Rep 1 | Rep 2 | Rep 3 | Rep 1 | Rep 2 | |
| A         | 4     | 3.27  | 3.25  | 3.39  | 3.23  | 3.21 | 3.27 +/- 0.07 (2.2) | 0.9971 |
|           | 3     | 2.32  | 2.28  | 2.30  | 2.25  | 2.18 | 2.27 +/- 0.05 (2.4) |
|           | 2     | 1.26  | 1.28  | 1.25  | 1.16  | 1.25 | 1.24 +/- 0.05 (3.9) |
|           | 1     | 0.46  | 0.53  | 0.51  | 0.36  | 0.32 | 0.44 +/- 0.09 (20.8) |
|           | 0     | N     | N     | N     | N     | N   | |
| B         | 4     | 2.83  | 2.87  | 2.89  | 2.87  | 2.89 | 2.87 +/- 0.03 (0.9) | 0.9990 |
|           | 3     | 1.81  | 1.80  | 1.81  | 1.84  | 1.92 | 1.84 +/- 0.05 (2.6) |
|           | 2     | 0.88  | 0.89  | 0.94  | 0.94  | 0.91 | 0.91 +/- 0.03 (3.4) |
|           | 1     | 0.26  | 0.36  | 0.18  | 0.15  | 0.40 | 0.27 +/- 0.11 (41.4) |
|           | 0     | N     | N     | N     | N     | N   | |
| C         | 4     | 2.84  | 2.82  | 2.70  | 2.79  | 2.79 | 2.79 +/- 0.05 (1.9) | 0.9999 |
|           | 3     | 1.84  | 1.91  | 1.85  | 1.85  | 1.83 | 1.86 +/- 0.03 (1.7) |
|           | 2     | 0.81  | 1.00  | 0.78  | 0.90  | 0.94 | 0.89 +/- 0.09 (10.3) |
|           | 1     | -0.10 | -0.22 | -0.30 | -0.10 | -0.40 | -0.22 +/- -0.13 (58.7) |
|           | 0     | N     | N     | N     | N     | N   | |
| D         | 4     | 2.57  | 2.56  | 2.57  | 2.58  | 2.58 | 2.57 +/- 0.01 (0.3) | 1.0000 |
|           | 3     | 1.55  | 1.58  | 1.62  | 1.59  | 1.65 | 1.60 +/- 0.04 (2.3) |
|           | 2     | 0.51  | 0.67  | 0.59  | 0.67  | 0.59 | 0.61 +/- 0.07 (11.5) |
|           | 1     | -0.15 | 0.15  | 0.15  | -0.15 | 0.04 | 0 +/- 0.15 |
|           | 0     | N     | N     | N     | N     | N   | |

<sup>a</sup> Log<sub>10</sub> (TCID<sub>50</sub>/ml).
<sup>b</sup> Log<sub>10</sub> (absolute copies/µl) of extracted viral RNA; R², Co-efficient of determination of the linear regression curve; Rep, replicate number; SD, standard deviation; CV, co-efficient of variation expressed in %; N, not detected; bold line: limit of detection; underlined line: limit of quantification.

an infected cell due to their proximity to the 3′ leader end of the genome (37, 38), were initially considered as targets for the design. However, no suitable regions could be found.

As described, the main focus of the current study was on the multiplexing aspect of the RT-ddPCR and not on its quantification capabilities. However, one advantage of developing ddPCRs is that absolute quantities of target molecules are calculated automatically without the need for the development of reference standards. Quantities in ddPCR are calculated using the “Poisson distribution model” based on the total number of droplets in a given reaction volume and the number of positive droplets within this population. This type of quantity estimation is not established by referring to a concentration curve generated using RNA transcripts that may give imprecise measures due to the initial method of dosage. Thus, quantities of target molecules in the initial sample as calculated by ddPCR could be interpreted to be more accurate; however, the efficiency of the RT-PCR also needs to be taken into account. The LoDs of the five-plex RT-ddPCR developed in the current study were in the same order as those described previously for subgroup specific AMPV real-time RT-PCR tests (26, 28). However, as described above, the differences in the methods of quantification by RT-ddPCR and real-time RT-PCR means that comparisons should be treated with caution.

In conclusion, an end-point RT-ddPCR has been developed that is capable of identifying AMPV-A, B, C and D in a single reaction. This multiplex RT-ddPCR is based on a highly conserved region of the genome and thus should have less risk of being affected by viral genome evolution and the impact it has on the specificity of such tests. A diagnostic test of such type with greater durability in terms of specificity will also help improve the accuracy when assessing the distribution of AMPV-A, B-C and D in the field. Further developments of this method could include the ability to detect co-infections with different AMPV wild type viruses or of wild type virus and vaccine strains, both unreported to date. Finally, because the primers used for this test also match the L gene sequences of the two newly identified gull and parakeet AMPVs, it may be possible to extend this five-plex RT-ddPCR to a seven-plex using machines offering more numerous fluorescent color options.
Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Author contributions

EL: study design, experiments, analyses, and writing of the manuscript. SB: statistical analysis of data. CA: experiments. CC and NE: critical analysis and contribution to manuscript text. PB: supervision of experiments and manuscript text. All authors reviewed and approved the manuscript.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fvets.2022.1058294/full#supplementary-material

SUPPLEMENTARY FIGURE S1
Dose-response of reference viruses for each tested AMPV subgroup (A, B, C, and D). Viral titer of each dilution of each reference virus is represented on X-axis (expressed in log_{10} TCID_{50}/ml). Corresponding values obtained by the five-plex RT-ddPCR AMPV are represented on Y-axis (expressed in log_{10} copies/μl). The coefficient of determination (R^2) of the linear regression model for each subgroup is indicated.
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