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Molecular subtyping of European swine influenza viruses and scaling to high-throughput analysis

Emilie Bonin 1,2,5, Stéphane Quéguiner 1,2, Cédric Woudstra 3, Stéphane Gorin 1,2, Nicolas Barbier 1,2, Timm C. Harder 4, Patrick Fach 3, Séverine Hervé 1,2 and Gaëlle Simon 1,2*

Abstract

Background: Swine influenza is a respiratory infection of pigs that may have a significant economic impact in affected herds and pose a threat to the human population since swine influenza A viruses (swIAVs) are zoonotic pathogens. Due to the increasing genetic diversity of swIAVs and because novel re assortants or variants may become enzootic or have zoonotic implications, surveillance is strongly encouraged. Therefore, diagnostic tests and advanced technologies able to identify the circulating strains rapidly are critically important.

Results: Several reverse transcription real-time PCR assays (RT-qPCRs) were developed to subtype European swIAVs in clinical samples previously identified as containing IAV genome. The RT-qPCRs aimed to discriminate HA genes of four H1 genetic lineages (H1av, H1hu, H1huΔ146–147, H1pdm) and one H3 lineage, and NA genes of two N1 lineages (N1, N1pdm) and one N2 lineage. After individual validation, each RT-qPCR was adapted to high-throughput analyses in parallel to the amplification of the IAV M gene (target for IAV detection) and the β-actin gene (as an internal control), in order to test the ten target genes simultaneously on a large number of clinical samples, using low volumes of reagents and RNA extracts.

Conclusion: The RT-qPCRs dedicated to IAV molecular subtyping enabled the identification of swIAVs from the four viral subtypes that are known to be enzootic in European pigs, i.e. H1avN1, H1huN2, H3N2 and H1N1pdm. They also made it possible to discriminate a new antigenic variant (H1huN2Δ146–147) among H1huN2 viruses, as well as reassortant viruses, such as H1huN1 or H1avN2 for example, and virus mixtures. These PCR techniques exhibited a gain in sensitivity as compared to end-point RT-PCRs, enabling the characterization of biological samples with low genetic loads, with considerable time saving. Adaptation to high-throughput analyses appeared effective, both in terms of specificity and sensitivity. This new development opens novel perspectives in diagnostic capacities that could be very useful for swIAV surveillance and large-scale epidemiological studies.

Keywords: Influenzavirus, Subtyping, Hemagglutinin, Neuraminidase, Pig, High-throughput real-time RT-PCR, LightCycler®1536, Surveillance, Diagnosis

Background

Swine influenza (SI) is a highly contagious viral respiratory infection of pigs that has become enzootic in areas densely populated with this species [1]. It is responsible for significant economic losses in affected herds due to morbidity, stunted growth and secondary infections.

Swine influenza A viruses (swIAVs) are also of public health concern due to their zoonotic potential. Although human infections remain subclinical or commonly produce only mild symptoms, swIAVs may be responsible for fatal cases and/or contribute, e.g. by reassortment, to pandemics, as illustrated by the pandemic that occurred in 2009 [2, 3].

swIAVs are enveloped viruses belonging to the genus Influenzavirus A, family Orthomyxoviridae. Their genome consists of eight segments of single-stranded RNA
The development and harmonization of diagnostic tools rapidly are critically important. Nostic tests and advanced technologies able to identify circulating strains are strongly encouraged [8]. Therefore, diagnosis tests and advanced technologies able to identify circulating strains rapidly are critically important. The development and harmonization of diagnostic tools for detection (M gene) and subtyping (HA and NA genes) of European swIAVs have been initiated in some European countries, especially through the concerted action “European surveillance network for influenza in pigs (ESNIP)” [8, 19]. One issue, following a multi-center study using a panel of reference strains from several countries, was the selection of the best-performing primers/probe sets and amplification protocols that could be used in reverse transcription real-time PCR assays (RT-qPCRs) for the detection of HA and NA genes from the different genetic lineages [20]. In this paper, we describe RT-qPCRs aimed specifically at identifying HA and NA genes from European swIAVs, taking into account local strains isolated recently in France. Following an original approach, we also report their adaptation to high-throughput analyses, in parallel to the amplification of the IAV M gene and the β-actin gene, which provide a comprehensive tool for detection and subtyping of European swIAVs in large biological sample sets.

Methods

Clinical samples and virus strains

Nasal swabs (MW950Sent2mL Virocult®, Kitvia, Labarthe-Inard, France) were collected from herds in France by veterinary practitioners or ANSES personnel, in the context of virological diagnosis passive surveillance programs or specific epidemiological investigations. They were taken from pigs during outbreaks of acute respiratory disease. The swabs were mixed vigorously and supernatants were stored at −70 °C until analysis. Some lung samples were also obtained from necropsy of fatal cases. Total viral RNAs were extracted from 200 μL of nasal swab supernatants or 20–30 mg of lung homogenate using the NucleoSpin RNA or Nucleospin 8 RNA kits (Macherey-Nagel, Hoerdt, France) or the RNeasy Mini Kit (Qiagen, Courtaboeuf, France). IAV genome detection was performed by M gene RT-qPCR using one of the two ready-to-use commercial kits previously validated by the French National Reference Laboratory (NRL), i.e. the LSI VetMAX™ Swine Influenza A−A/H1N1/2009− included kit (Life technologies, Carlsbad, CA, USA) or the ADIAVET™ SIV REALTIME kit (Bio-X Diagnostics, Rochefort, France). Both steps, i.e. viral RNA purification and IAV genome detection, followed the instructions provided by RT-qPCR manufacturers [21].

Virus isolation was attempted from positive clinical samples in Madin Darby canine kidney (MDCK) cells, according to standard procedures [22]. Reference strains, previously selected as representative of the main viral subtypes and lineages encountered in European pigs over the last few years, were also retrieved from the collection of the French NRL or kindly provided by the European surveillance network for influenza in pigs.
(ESNIP) [8, 23]. A/California/04/2009 (H1N1pdm) was obtained from the French National Reference Center for Influenza, Institut Pasteur, Paris.

Initial subtyping

Molecular subtyping

Initially, swIAV-positive samples and/or virus isolates were subjected to molecular subtyping using conventional RT-PCR assays, as previously described [19, 23]. One multiplex RT-PCR enabled the specific detection of HA genes of the H1av (clade 1C [10]), H1hu (clade 1B.1 [10]) and H3 lineages, while another enabled the amplification of NA genes of the N1 and N2 lineages. Since 2010, H1 and N1 genes from H1N1pdm have been screened using ready-to-use commercial kits previously validated by the French NRL for Swine Influenza [21], according to the manufacturer’s instructions. The H1pdm gene (clade 1A.3.3.2 [10]) was detected using the real-time RT-PCR LSI VetMAX™ Swine Influenza A-A/H1N1/2009-H1 detection kit (Life technologies, Carlsbad, CA, USA) or the ADIAVET™ A/H1N1(2009) REALTIME kit (Bio-X Diagnostics, Rochefort, Belgique). The N1pdm gene was detected using the real-time RT-PCR LSI VetMAX™ Swine Influenza A-A/H1N1/2009-N1 detection kit (Life technologies, Carlsbad, CA, USA).

Antigenic subtyping

Some virus strains were further propagated in 9-day-old specific pathogen-free (SPF) embryonated chicken eggs and subjected to antigenic characterization using hemagglutination inhibition (HI) tests, according to standard procedures [22]. Allantoic fluids were tested with hyperimmune sera produced in SPF pigs against reference strains representative of the different subtypes: A/Sw/Cotes d’Armor/0388/2009 (H1av,N1), A/Sw/Flandres/1/1998 (H3N2), A/sw/Scotland/410440/94 (H1hu,N2), A/Sw/Cotes d’Armor/113/2006 (H1hu,N2), A/Sw/Cotes d’Armor/0070/2010 (H1hu,N1), A/Sw/Cotes d’Armor/0186/2010 (H1hu,N2), A/Sw/Sarthe/0255/2010 (H1N1pdm) and A/Sw/France/22–130212/2013 (H1hu,N2Δ146–147) [23, 24].

Sequencing

The subtype and lineage of some selected swIAVs and some positive field samples were determined and/or confirmed by sequencing the HA and NA gene segments. The viral genome was reverse transcribed using universal primers [25] and SuperScript II Reverse Transcriptase (Life technologies, Carlsbad, CA, USA) following the manufacturer’s instructions. Full length HA and NA genes were amplified using Platinum Taq DNA Polymerase High Fidelity (Life technologies, Carlsbad, CA, USA) and in-house designed primers (sequences available on request). Amplified products were then separated in agarose gel and purified with the QIAquick Gel Extraction Kit (Qiagen, Hilden, Germany). Both strands of the amplicons were sequenced with the same primers used for the amplification, as well as with additional internal primers (sequences available on request). Sequencing reactions were performed using the BigDye Terminator v3.1 Cycle Sequencing Kit (Life technologies, Carlsbad, CA, USA) on an automatic DNA sequencer ABI 3130 Genetic Analyzer (Life technologies, Carlsbad, CA, USA). Manual editing of sequences and assembly refinement were done with Vector NTI Advance 11.0 software (Life technologies, Carlsbad, CA, USA). The Influenza Research Database (IRD) was screened with BLASTN2 to identify sequences closely related to the HA and NA sequences.

Primers and probes for HA and NA RT-qPCRs

Sets of primers (forward and reverse) and the probe were designed to specifically amplify the different HA and NA genes from European swIAVs, on the basis of alignments of nucleotide sequences of viruses isolated since 2000 and retrieved from the IRD, GISAID (Global Initiative on Sharing All Influenza Data) or Genbank databases. Melting temperatures and/or basic properties of oligonucleotides were approximated using OligoCalc [26] or Vector NTI Advance sequence analysis and design software (Thermo Fisher Scientific, Waltham, MA, USA) (Table 1). Whereas ready-to-use commercial kits have previously been validated for H1pdm and N1pdm detection [21], new sets of primers and the probe specific to these two genes were designed in our study in order to include H1pdm and N1pdm RT-qPCRs in the high-throughput PCR array. Thus, the H1av, H1hu, H1pdm and H3 sets specifically amplify HA genes from the European H1av,N1, H1hu,N2, H1N1pdm and H3N2 lineages, respectively. The N1 set matches with NA genes from both the H1av,N1 and the H1N1pdm lineages whereas the N1pdm set specifically amplifies the NA gene from the H1N1pdm lineage. The N2 primers and probe can detect NA genes from both the H1hu,N2 and the H3N2 lineages. In addition, an H1huΔ146–147 set was manually designed to identify the novel antigenic variant that has emerged among H1hu,N2 viruses following antigenic drift [18]. Alignments comprised full length HA sequences of H1hu,N2 and H1huΔ146–147, as well as various H1hu,N1 reassortant viruses. Thus, the H1huΔ146–147 set design took into account a deletion of 6 nucleotides and a mutation located in the receptor binding site of the H1hu,N2Δ146–147 variant (manuscript in preparation). Finally, M and β-actin sets were included, since in-house RT-qPCRs for swIAV and reference gene detection were used, respectively, as controls in high-throughput PCR assays. All RT-qPCRs used standard
TaqMan DNA probes except those for H1pdm, N1pdm and H1huΔ146–147 which used highly specific DNA probes with conjugated minor groove binder (MGB) groups at the 3’-end. The fluorophore covalently attached to the 5’-end of the oligonucleotide probe was 6-carboxyfluorescein (FAM), hexachlorofluorescein (HEX) or cyanine5 (Cy5) (Table 1). All standard TaqMan probes used black hole quencher (BHQ1) at the 3’-end. Primers and labelled standard TaqMan probes were purchased from Sigma-Aldrich (Saint-Louis, MO, USA). MGB-labelled probes were from Life technologies (Carlsbad, CA, USA).

**Real-time RT-PCR assays for European swIAV subtyping**

Similarly to virus detection by M gene RT-qPCR, the HA/NA subtyping methods were assessed on RNA extracted from amplified viruses or clinical samples using the NucleoSpin RNA kit (Macherey-Nagel, Hoerdt, France) or the RNeasy Mini Kit (Qiagen, Courtaboeuf, France) following the manufacturer’s instructions. Real-

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**Table 1 Primers and probes**

| Target Gene | Primer/Probe | Sequence, labeling | Location | Reference Sequence (accession number) |
|-------------|--------------|--------------------|----------|--------------------------------------|
| H1av | H1av_Fo | gaaggggatgacaggaatga | 1063–1084 | A/Sw/Cotes d’Armor/0388/2009 (KC881265) |
| | H1av_Re | caatahtgaccttctggtgc | 1178–1201 | |
| | H1av_Pr | (HEX)-tcctggtcagctcagaaat-BHQ1 | 1126–1150 | |
| H1hu | H1hu_Fo_1 | gaggaggagctggccagtaaaatgtgtagag [i] | 1090–1109 | A/Sw/Cotes d’Armor/0113/2006 (AM503902) |
| | H1hu_Fo_2 | gattggtcgggtgacagat | 1064–1109 | |
| | H1hu_Re_1 | acctacgcttgtgtaatgttttcctc-3' | 1184–1204 | |
| | H1hu_Re_2 | ttctgatcagaaaa-GAG-3' | 1184–1232 | |
| | H1hu_Pr | (FAM)-cagggatgctgctgtagaag-BHQ1 | 1120–1144 | |
| H1huΔ146–147 | H1hu_dif_Fw | agttgagttcagctcagagaaat | 367–398 | A/Sw/France/22-130212/2013 (KJ128323) |
| | H1hu_dif_Rv | acctacgcttgtgtaatgttttcctc-3' | 1184–1204 | |
| | H1hu_var_FAM | acctacaacatcagaaaa-GAG-3' | 337–356 | |
| | H1hu_var_CY5 | cctgttttcctcagggaaggga-3' | 254–278 | |
| H1pdm | H1pdm_Fo | gggcattgacaggaatga | 582–603 | A/California/04/2009 (FJ966082) |
| | H1pdm_Re | cctcactctctctctctctctctctctctc-ttgccagaaatgtgta | 689–714 | |
| | H1pdm_Pr | (FAM)-atcagctgctgctgtagaag-BHQ1 | 666–685 | |
| H3 | H3_Swine_Fw | cttgatggrggamaytgcaca | 223–243 | A/Sw/France/59–120031/2012 (KC345622) |
| | H3_Swine_Rv | gcacatcagaaaa-GAG-3' | 337–356 | |
| | H3_Swine_CY5 | cctgttttcctcagggaaggga-3' | 254–278 | |
| N1 | N1.3_F | agrccctgtgctgctggaatga | 1255–1274 | A/Sw/Germany/SIV04/2008 (FN429079) [20] |
| | N1.3_R | accgtctgctgctgctg-3' | 1363–1380 | |
| | AV N1.3 FAM | cctgttttcctcagggaaggga-3' | 1306–1328 | |
| N1pdm | N1pdm_Fo | gggacagacaatactctcttaaaagc | 1144–1171 | A/California/04/2009 (FJ966082) |
| | N1pdm_Re | ttcagctgctgctgctgctg-3' | 1220–1243 | |
| | N1pdm_Pr | (FAM)-atcagctgctgctgtagaag-BHQ1 | 1184–1206 | |
| N2 | N2_1367F | agttcagctgtgctgctg-3' | 1305–1325 | A/Sw/Bakum/8602/1999 (EF409258) [20] |
| | N2_1468R | ttgcagagtctgctgctgctg-3' | 1397–1420 | |
| | AV N2_1444 HEX | (HEX)-ctctacggctagctgctgctgctgctgctgctg-3' | 1357–1382 | |
| M | SIV-Forw | atytggacyagtgggagcagcat | 24–47 | A/Sw/France/22-130212/2013 (KM267912) [30] |
| | SIV-Rev | tcctgcattcttgctgctgctg-3' | 101–124 | |
| | M64_FAM | (FAM)-ctctacggctagctgctgctgctgctgctgctg-3' | 74–93 | |
| | | (HEX)-atcagctgctgctgctgctgctgctgctgctgctgctgctgctg-3' | 74–93 | |

*From the first nucleotide of the coding sequence, except for M (from the first nucleotide of the segment)*
time RT-PCR assays were run either on Chromo4 (Bio-Rad Laboratories, Hercules, CA, USA) or MX3005P (Stratagene, Agilent Technologies, La Jolla, CA, USA). Each RT-qPCR mixture contained 5 μL of RNA extract and 20 μL of master mix containing 1X GoTaq Probe qPCR Master Mix, dUTP (2X) (Promega, Fitchburg, WI, USA), 2X GoScript RT Mix for 1-step RT-qPCR (50X) (Promega, Fitchburg, WI, USA) as well as primers and probe in different concentrations depending on the target gene. Thus, the master mix contained 800 nM of each primer and 100 nM of probe for H1av, H3, N1 and N2, 800 nM of each primer and 140 nM of probe for H1hu, 400 nM of each primer and 500 nM of probe for H1huΔ146–147, and 800 nM of each primer and 250 nM of probe for H1pdm and N1pdm. All RT-qPCRs were run as singleplex assays, except N1 and N2 that were run as a duplex assay. The cycling conditions used for H1av, H1hu, H3 and N1/N2 were: 15 min reverse transcription at 45 °C, 2 min of denaturation at 95 °C and 42 cycles of 15 s at 95 °C and 1 min at 56 °C. The program used for H1huΔ146–147, H1pdm and N1pdm (with MGB probe) was 15 min reverse transcription at 45 °C, 2 min of denaturation at 95 °C, and 40 cycles of 15 s at 95 °C and 1 min at 60 °C.

**Evaluation of real-time RT-PCR subtyping performance**

Several panels incorporating virus strains and/or clinical samples were constituted to evaluate the performance of the RT-qPCRs developed for the amplification of the H1av, H1hu, H1huΔ146–147, H3, N1 and N2 genes.

A first panel (panel 1) of 42 swIAV isolates was set up to assess the analytical specificity of the RT-qPCRs designed for H1av, H1hu, H3, N1 and N2 gene amplifications. Among these virus strains, 32 belong to one of the four European enzootic lineages, i.e. 12 H1avN1, 10 H1huN2, 8 H1N1pdm and 2 H3N2, whereas 10 others are isolates originating from reassortments between enzootic viruses, i.e. 5 H1huN1 and 5 H1avN2 (Table 2). RNA extracts were diluted in RNase free-water to adjust the quantification cycle (Cq) value to 24–27 when tested with one of the M gene RT-qPCRs described above, in order to reduce the risk of cross-reactions between H1av and H1hu.

Panel 2 was assembled to test the specificity of the H1huΔ146–147 gene RT-qPCR, i.e. its ability to differentiate the H1 gene of the antigenic variant H1huN2Δ146–147 virus among HA genes previously identified as belonging to the H1hu lineage. This panel consisted of RNA extracts obtained from 46 swIAV strains previously identified as H1huN2 viruses (44/46) or H1huN1 viruses (2/46) (Additional file 1).

The analytical sensitivity of the assays was evaluated by testing 10-fold serial dilutions (from 10⁻² to 10⁻⁹) of RNA extracted from reference strains (Table 3). The range of linearity, the coefficient of linear regression (R²) and the efficacy were then calculated to evaluate the performance (Table 4). The repeatability of each RT-qPCR was estimated by testing, in 5–6 independent assays, the RNA extract dilution immediately below the last dilution in which the target gene was detected by the assay, and calculating the inter-assay coefficient of variation (CV) (Table 5).

Panels 3 and 4 were set up to evaluate the diagnostic abilities of the various RT-qPCRs, i.e. their potential to identify HA and NA genes directly from clinical samples previously shown to contain swIAV genome (M gene RT-qPCR positive). Panel 3 included 60 RNA extracts obtained from 59 nasal swab supernatants and one lung tissue homogenate (Table 6). The swIAV genome was previously subtyped in 37 nasal swabs and the lung sample (11 samples were positive for H1avN1, 4 for H1huN2, 15 for H1huN2, 2 for H1huN1, 1 for H3N2, 4 for H1N1pdm, and 1 for both H1av and H1hu genes mixed with N2), whereas in 22 nasal swab supernatants the swIAV genome was only partially subtyped using conventional RT-PCRs (18 samples tested positive for H1av but not for N1 or N2; 3 samples tested negative for any HA but were found positive for N2, and 1 sample was found to be negative for HA and NA). Panel 4 was specifically assembled to assess the diagnostic ability of the H1huN1Δ146–147 RT-qPCR. It was composed of 78 RNA extracts, i.e. 56 obtained from nasal swab supernatants and 22 from swIAV strains, that were previously found either positive for H1hu but not sequenced, or positive for N2 but in which HA was not identified, or in which neither HA nor NA were identified (see Additional file 2).

**High-throughput real-time RT-PCR procedure**

High-throughput real-time RT-PCR amplifications (M, β-actin, H1av, H1hu, H1huΔ146–147, H1pdm, H3, N1, N1pdm and N2) were conducted on a LightCycler 1536 real-time PCR system (Roche, Meylan, France). A Bravo automated liquid handling platform equipped with a chiller and a PlateLoc thermal microplate sealer (Agilent Technologies, La Jolla, CA, USA) was used as a microplate dispenser. Each RT-qPCR mixture contained 1 μL of RNA extract and 1 μL of master mix containing 1X RealTime ready DNA Probes Master (Roche, Meylan, France), 2X GoScript RT Mix for 1-step RT-qPCR (50X) (Promega, Fitchburg, WI, USA), 800 nM of each concerned primer, with the exception of the M gene forward primer that was included at a final concentration of 400 nM, and 250 nM of standard or MGB-labelled TaqMan probe. The one-step real-time RT-PCR program involved a 30 min reverse transcription of RNA at 45 °C, followed by a 2 min denaturation step at 95 °C, and lastly 40 cycles of 0 s (i.e. a pulse) at 95 °C and 1 min at 58 °C.
**Table 2** Specificity of the real-time RT-PCR assays developed for the identification of swIAV HA and NA genes originating from the European enzootic lineages H1av-N1, H1hu-N2, H1N1pdm and H3N2

| Virus strains (Panel 1) | Lineage known from conventional RT-PCRs and commercial H1pdm and N1pdm RT-qPCRs[^a] | M-gene RT-qPCR[^b] (Cq-value) | Real-time RT-PCRs for molecular subtyping[^c] (Cq-value) | HA subtyping | NA subtyping |
|------------------------|---------------------------------------------------------------------------------------|-----------------------------|-----------------------------------------------------|--------------|---------------|
| **Subtype and lineage** | **Name**                                                              |  **H1[^av]** |  **H1[^hu]** |  **H3** |  **N1** |  **N2** |
| H1[^av]                | A/Sw/Asnie/0054/10                                                               | 25.48          | 24.16        | no Cq  | no Cq  | 24.61  | no Cq  |
|                        | A/Sw/Ille et Vilaine/0187/11                                                      | 26.87          | 26.92        | no Cq  | no Cq  | 24.45  | no Cq  |
|                        | A/Sw/Ille et Vilaine/0208/11                                                      | 24.6           | 23.45        | no Cq  | no Cq  | 19.05  | no Cq  |
|                        | A/Sw/Morbihan/0294/11                                                             | 27             | 25.88        | no Cq  | no Cq  | 24.86  | no Cq  |
|                        | A/Sw/Finistere/0307/11                                                            | 27.12          | 30.23        | no Cq  | no Cq  | 28.64  | no Cq  |
|                        | A/Sw/LoireAtlantique/0405/11                                                      | 26.9           | 29.84        | no Cq  | no Cq  | 28.03  | no Cq  |
|                        | A/Sw/France/41–120137/12                                                          | 24.33          | 19.84        | no Cq  | no Cq  | 27.55  | no Cq  |
|                        | A/Sw/France/72–120183/12                                                          | 25.12          | 25.16        | no Cq  | no Cq  | 25.27  | no Cq  |
|                        | A/Sw/France/29–120326/12                                                          | 26.34          | 36.15        | no Cq  | no Cq  | 27.68  | no Cq  |
|                        | A/Sw/France/22–120340/12                                                          | 24.29          | 23.40        | no Cq  | no Cq  | 22.70  | no Cq  |
|                        | A/Sw/France/56–120452/12                                                          | 25.75          | 30.63        | no Cq  | no Cq  | 25.91  | no Cq  |
|                        | A/Sw/France/53–130065/13                                                          | 26.65          | 33.03        | no Cq  | no Cq  | 30.16  | no Cq  |
|                        | A/Sw/Cotes d’Armor/0102/08                                                       | 25.38          | 23.13        | no Cq  | no Cq  | 23.62  | no Cq  |
|                        | A/Sw/Cotes d’Armor/0186/10                                                       | 26.56          | 24.92        | no Cq  | no Cq  | 26.69  | no Cq  |
|                        | A/Sw/Morbihan/0213/11                                                             | 24.26          | 24.37        | no Cq  | no Cq  | 23.98  | no Cq  |
|                        | A/Sw/Morbihan/0599/11                                                             | 24.6           | 26.17        | no Cq  | no Cq  | 24.06  | no Cq  |
|                        | A/Sw/France/37–120,345/12                                                        | 24.34          | 27.26        | no Cq  | no Cq  | 24.91  | no Cq  |
|                        | A/Sw/France/0074/11                                                              | 26.81          | 30.80        | no Cq  | no Cq  | 29.33  | no Cq  |
|                        | A/Sw/France/22–110,153/11                                                        | 26.87          | 32.06        | no Cq  | no Cq  | 28.08  | no Cq  |
|                        | A/Sw/Finistere/0186/11                                                            | 26.63          | 31.08        | no Cq  | no Cq  | 29.74  | no Cq  |
|                        | A/Sw/Ille et Vilaine/0346/11                                                      | 25.6           | 26.83        | no Cq  | no Cq  | 25.26  | no Cq  |
|                        | A/Sw/Ille et Vilaine/0415/11                                                      | 24.05          | 28.07        | no Cq  | no Cq  | 26.43  | no Cq  |
|                        | A/Sw/Maine et Loire/0589/11                                                       | 24.65          | 23.19        | no Cq  | no Cq  | 23.27  | no Cq  |
|                        | A/Sw/France/56–120177/12                                                         | 25.05          | 30.97        | no Cq  | no Cq  | 30.66  | no Cq  |
|                        | A/Sw/France/22–120255/12                                                         | 25.68          | 29.93        | no Cq  | no Cq  | 27.52  | no Cq  |
|                        | A/Sw/France/29–120,258/12                                                        | 27.17          | 31.01        | no Cq  | no Cq  | 29.21  | no Cq  |
|                        | A/Sw/France/22–130032/13                                                         | 24.54          | 23.30        | no Cq  | no Cq  | 25.82  | no Cq  |
|                        | A/Sw/Cotes d’Armor/0046/08                                                       | 25.85          | 28.06        | no Cq  | no Cq  | 24.46  | no Cq  |
|                        | A/Sw/Cotes d’Armor/0070/10                                                       | 24.59          | 28.80        | no Cq  | no Cq  | 24.54  | no Cq  |
|                        | A/Sw/Morbihan/0163/10                                                             | 26.12          | 32.22        | no Cq  | no Cq  | 26.53  | no Cq  |
|                        | A/Sw/France/56–110255/10                                                         | 25.63          | 29.24        | no Cq  | no Cq  | 32.04  | no Cq  |
|                        | A/Sw/France/22–120067/12                                                         | 26.46          | 31.69        | no Cq  | no Cq  | 26.66  | no Cq  |
|                        | A/California/04/09                                                               | 25.92          | 27.76        | no Cq  | no Cq  | 27.76  | no Cq  |
|                        | A/Sw/Sarthe/0255/10                                                               | 26.26          | no Cq         | no Cq  | no Cq  | 26.51  | no Cq  |
|                        | A/Sw/Sarthe/0262/10                                                               | 25.32          | no Cq         | no Cq  | no Cq  | 26.02  | no Cq  |
|                        | A/Sw/Cotes d’Armor/110466/10                                                     | 25.15          | no Cq         | no Cq  | no Cq  | 27.35  | no Cq  |
Table 2  Specificity of the real-time RT-PCR assays developed for the identification of swIAV HA and NA genes originating from the European enzootic lineages H1avN1, H1huN2, H1N1pdm and H3N2 (Continued)

| Virus strains (Panel 1) | M gene RT-qPCRb (Cq-value) | Real-time RT-PCRs for molecular subtypingc (Cq-value) |
|------------------------|-----------------------------|-----------------------------------------------|
| Subtype and lineage     | HA subtyping                | NA subtyping                                 |
| HA lineage             |                             |                                               |
| of the most            |                             |                                               |
| antigenically          |                             |                                               |
| related reference      |                             |                                               |
| strain (from HI tests) |                             |                                               |
|                        | HA                          | NA*                                          |
| A/Sw/Haute-Loire/0578/11 | 26.42                      | no Cq                                        |
| A/Sw/France/18–120158/12 | 27.29                      | no Cq                                        |
| A/Sw/France/18–120333/12 | 24.04                      | no Cq                                        |
| A/Sw/France/71–130116/13 | 24.68                      | no Cq                                        |
| A/Sw/Flandres/1/98     | 24.98                      | no Cq                                        |
| A/Sw/France/59–120031/12 | 26.1                       | no Cq                                        |
| H3                     |                             |                                               |
| H3                     |                             |                                               |
| N2                     |                             |                                               |
| A/Sw/France/18–120158/12 | 28.44                      | no Cq                                        |
| A/Sw/France/18–120333/12 | 29.73                      | no Cq                                        |
| A/Sw/France/71–130116/13 | 24.37                      | no Cq                                        |
| A/Sw/France/59–120031/12 | 25.58                      | no Cq                                        |

aN1av means that the conventional N1 RT-qPCR was positive but the N1pdm commercial RT-qPCR was negative; bCommercial kits [21]; cHA RT-qPCRs were run as monoplex procedures whereas NA RT-qPCRs were run in duplex. Cq: quantification cycle.

Evaluation of high-throughput RT-qPCR performance

Panel 5 was established to evaluate the analytical specificity of the real-time RT-PCRs (including new in-house H1pdm and N1pdm RT-qPCRs) when using the high-throughput RT-qPCR procedure (see Additional file 3). This panel included of RNA extracts obtained from 88 swIAVs and 30 nasal swab supernatants. The isolates as well as the viruses present in clinical samples were first subtyped using RT-qPCRs run under low-throughput conditions. Serial 10-fold dilutions (10\(^{-1}\) to 10\(^{-8}\)) of RNA obtained from 8 reference strains were prepared in water to evaluate the analytical sensitivity of each RT-qPCR when run on the LightCycler® 1536 (Table 7). The M gene copy number present in each dilution point was measured using a duplex M/β-actin gene RT-qPCR as described separately [27]. Each dilution point was tested in duplicate.

Results

Analytical performances of the real-time RT-PCRs for H1av, H1hu, H1huΔ146–147, H3, N1 and N2 gene amplification

The analytical performances of the novel RT-qPCRs developed for European swIAV subtyping in addition to the H1pdm and N1pdm commercial kits previously validated were first evaluated in low-throughput analyses.

Analytical specificity

Analyses of 42 swIAV isolates from panel 1 showed perfect concordance between the results from H1av, H1hu, H3, N1 and N2 RT-qPCRs and those expected from conventional, i.e. end-point RT-PCRs and/or sequencing data, indicating 100% analytical specificity for the assays (Table 2). Despite non negligible viral loads, i.e. Cq-values around 25 in M gene RT-qPCR, none of these assays showed any undesired amplification of genes from influenza viruses other than those expected. According to plan, the N1 RT-qPCR was able to detect NA genes from the H1avN1 and H1N1pdm lineages, while the N2 RT-qPCR amplified NA genes of viruses from the H1huN2 and H3N2 lineages.

Among the 46 swIAV strains included in panel 2 and previously identified as viruses that bear an H1 hu gene, 25/44 H1 huN2 strains were detected by the H1 huΔ146–147 RT-qPCR indicating they were variant viruses, in accordance with HA gene sequencing (see Additional file 1). All other H1 huN2 strains did not exhibit the 6 nt-deletion encountered in the variant and were not detected by specifically designed RT-qPCR H1 huΔ146–147, showing that there were no false-positive results. By contrast, the two H1 huN1 viruses were not detected even though they also showed the 146–147 amino acid deletion, due to a mutation in the MGB probe target region. Indeed, sequencing revealed that these viruses retained a thymine residue (T) in probe position 12, instead of having mutated into a cytosine residue (C) as compared to the parental H1 hu gene. These reassortant viruses, 100% identical in their HA genes, were isolated from the same farm 2.5 months apart.

Analytical sensitivity

Real-time RT-PCRs developed for HA or NA subtyping of previously detected swIAVs exhibited lower sensitivity levels than the M gene assays used for the detection, except those amplifying the H1 huΔ146–147 and the H3 genes (Table 3). Thus, assays conducted on 10-fold dilutions of reference strains indicated that the detection limits
Table 3: Analytical sensitivity of the real-time RT-PCRs for H1av, H1hu, H3, N1 and N2 on ten-fold dilutions of RNA extracted from swIAVs of different subtypes

| Virus strain | Virus stock dilution | M gene RT-qPCR\(^a\) (Cq-value) | Real-time RT-PCR\(^b\) for molecular subtyping (Cq-value) |
|--------------|----------------------|---------------------------------|-------------------------------------------------------|
| H1\(_av\)N1 | A/Sw/Cotes d'Armor/0388/2009 | 10\(^{-1}\) 12.52 16.10 no Cq nt no Cq 15.52 no Cq |
|              |                      | 10\(^{-2}\) 16.09 19.52 no Cq nt no Cq 18.75 no Cq |
|              |                      | 10\(^{-3}\) 20.16 22.77 no Cq nt no Cq 22.46 no Cq |
|              |                      | 10\(^{-4}\) 23.83 25.85 no Cq nt no Cq 25.71 no Cq |
|              |                      | 10\(^{-5}\) 27.03 28.86 no Cq nt no Cq 28.18 no Cq |
|              |                      | 10\(^{-6}\) 29.79 32.32 no Cq nt no Cq no Cq no Cq |
|              |                      | 10\(^{-7}\) 33.36 36.26 no Cq nt no Cq no Cq no Cq |
|              |                      | 10\(^{-8}\) 38.59 no Cq no Cq nt no Cq no Cq no Cq |
|              |                      | 10\(^{-9}\) no Cq no Cq no Cq no Cq no Cq no Cq no Cq |
|              |                      | 10\(^{-10}\) no Cq no Cq no Cq no Cq no Cq no Cq no Cq |
| H1\(_hu\)N2 | A/Sw/Scotland/410440/1994 | 10\(^{-1}\) 13.47 no Cq 15.18 nt no Cq no Cq 16.90 |
|              |                      | 10\(^{-2}\) 17.15 no Cq 18.53 nt no Cq no Cq 19.92 |
|              |                      | 10\(^{-3}\) 20.43 no Cq 21.99 nt no Cq no Cq 23.54 |
|              |                      | 10\(^{-4}\) 23.79 no Cq 25.19 nt no Cq no Cq 26.60 |
|              |                      | 10\(^{-5}\) 27.25 no Cq 28.06 no Cq no Cq 28.68 |
|              |                      | 10\(^{-6}\) 30.11 no Cq 30.44 no Cq no Cq 30.91 |
|              |                      | 10\(^{-7}\) 33.05 no Cq 33.36 no Cq no Cq 33.67 |
|              |                      | 10\(^{-8}\) 37.48 no Cq 37.10 no Cq no Cq 37.72 |
|              |                      | 10\(^{-9}\) no Cq no Cq no Cq no Cq no Cq no Cq no Cq |
|              |                      | 10\(^{-10}\) no Cq no Cq no Cq no Cq no Cq no Cq no Cq |
| H1\(_hu\)N2Δ146–147 | A/Sw/France/22–130212/2013 | 10\(^{-1}\) 10.98 nt 11.88 12.68 nt no Cq 11.48 |
|              |                      | 10\(^{-2}\) 14.67 nt 15.26 16.10 nt no Cq 14.38 |
|              |                      | 10\(^{-3}\) 17.82 nt 18.66 20.07 nt no Cq 17.78 |
|              |                      | 10\(^{-4}\) 21.3 nt 21.77 22.95 nt no Cq 20.92 |
|              |                      | 10\(^{-5}\) 25.64 nt 25.03 26.53 nt no Cq 24.54 |
|              |                      | 10\(^{-6}\) 28.06 nt 29.83 no Cq 30.06 no Cq |
|              |                      | 10\(^{-7}\) 31.54 nt 33.36 no Cq 33.67 no Cq |
|              |                      | 10\(^{-8}\) 35.16 nt 37.03 no Cq 37.34 no Cq |
|              |                      | 10\(^{-9}\) no Cq nt 39.63 no Cq 39.99 no Cq |
|              |                      | 10\(^{-10}\) no Cq nt no Cq no Cq nt no Cq |
| H3N2 | A/Sw/Flandres/1/1998 | 10\(^{-1}\) 13.13 no Cq no Cq nt 17.045 no Cq 16.865 |
|              |                      | 10\(^{-2}\) 16.7 no Cq no Cq nt 19.88 no Cq 19.725 |
|              |                      | 10\(^{-3}\) 20.21 no Cq no Cq nt 23.55 no Cq 23.275 |
|              |                      | 10\(^{-4}\) 23.57 no Cq no Cq nt 27.075 no Cq 26.67* |
|              |                      | 10\(^{-5}\) 26.9 no Cq no Cq nt 30.235 no Cq 30.12 |
|              |                      | 10\(^{-6}\) 30.07 no Cq no Cq nt 33.53 no Cq 33.42 |
|              |                      | 10\(^{-7}\) 32.7 no Cq no Cq nt 35.33 no Cq 35.23 |
|              |                      | 10\(^{-8}\) 37.04 no Cq no Cq nt 37.86 no Cq 37.76 |
|              |                      | 10\(^{-9}\) no Cq no Cq no Cq nt 39.99 no Cq 39.9 |
|              |                      | 10\(^{-10}\) no Cq no Cq no Cq nt no Cq no Cq |

\(^{a}\) Cq quantification cycle, nt not tested. \(^{b}\) M gene RT-qPCRs were run on one replicate whereas molecular subtyping RT-qPCRs were run on duplicates (Thermocycler MxPro – Mx3005P). Thus, Cq-values for molecular subtyping RT-qPCRs are mean Cq-values between duplicates, except numbers indicated in italics with an asterisk (*). Cq-values indicated in bold represent detection limits. \(^{\dagger}\) Commercial kits [21].
Based on M gene RT-qPCR Cq-values would be >35 for H1huΔ146-147 and H3, around 33 for H1avN2, around 27 for N1, between 24 and 28 for N2 depending on the virus subtype, and around 24 for H1hu. Based on calculation of the ranges of linearity, it appeared that all genes were correctly detected from the first (10⁻¹) strain dilution, i.e. at Cq-value <13.5, except the H3 gene for which the lowest limit was estimated at the second (10⁻²) dilution, at a Cq-value equivalent to 16.7 (Table 4). Between the limits of the range of linearity, the slopes of the standard curves varied from -3.266 to -3.3483. The coefficients of linear regression (R²) varied from 0.9957 to 0.9997 and the efficiencies were calculated between 96.81% and 104.16%. Repeatability and intermediate precision, assessed by calculating the inter-assay CVs, gave satisfactory results (Table 5). The inter-assay CVs were all <10%, ranging from 1.20 for the H1av RT-qPCR to 6.51 for the N2 RT-qPCR run on the H3N2 strain.

**Diagnostic abilities of the RT-qPCR assays targeting the H1av, H1hu, H1huΔ146-147, H3, N1 and N2 genes of European swIAVs**

Nasal swabs from animals naturally infected with H1avN1, H1huN2 or H3N2 swIAVs, as well as with H1huN2 or H1huN1 reassortant viruses (panel 3), were detected positive for both HA and NA corresponding genes in 100% of cases tested for both genes (Table 6). Additional samples tested either only in the simplex HA assays or in the N1/N2 multiplex assay were also qualified positive according to the expected HA or NA gene. Inclusion of 4 samples positive for H1N1pdm confirmed N1 amplification without any H1av gene amplification. Sample 130103-2, which was previously known to contain both the H1av and H1hu genes together with the N2 gene, was also proved here to contain the N1 gene, enabling us to confirm the hypothesis of a mixture of H1avN1 and H1huN2 viruses in this sample. Analyses of 18 samples previously partially subtyped as “H1avN?” confirmed the H1 av subtype in 9/18 cases. Others did not give a positive signal, probably due to limiting genome amounts (7/11 were samples with M gene RT-qPCR Cq-values >32, while the H1 av RT-qPCR detection limit was estimated around M gene Cq-value = 33). Conversely, the N1 gene was identified in 10/18 H1avN? samples, even in samples where M gene Cq-value >27, which was estimated as the detection limit. The N2 gene was not amplified in any of these samples. Panel 3 also comprised 3 H?N2 samples. The N2 gene was detected in all of them, whereas the H1hu gene was identified in 2/3. The last one (sample 130277–3) was shown to have an H1hu gene after propagation in cell culture (data not shown). Finally, a sample (130133–5) that was left fully unsubtyped using conventional RT-PCRs was here classified as containing the H1av, H1hu and N2 genes, confirming the ability to detect virus mixtures using these RT-qPCRs.

**Table 4** Performance of real-time RT-PCRs for H1av, H1hu, H1huΔ146-147, H3, N1 and N2 on ten-fold dilutions of RNA extracted from swIAVs representative of the different subtypes

| Virus strain | Criteria | Real-time RT-PCR for molecular subtyping | H1av | H1hu | H1huΔ146-147 | H3 | N1 | N2 |
|--------------|----------|------------------------------------------|------|------|--------------|----|----|----|
| A/Sw/Cotes d’Armor/0388/2009 (H1avN1) | Range of linearity | **<12.52 to 33.36** | **<12.52 to 27.03** | | | | | |
| | Slope  | **−3.2918** | **−3.266** | | | | | |
| | R² | 0.9986 | 0.9957 | | | | | |
| | Efficacy | 101.27% | 104.16% | | | | | |
| A/Sw/Scotland/410440/1994 (H1huN2) | Range of linearity | **<13.47 to 23.79** | **<13.47 to 23.79** | | | | | |
| | Slope  | **−3.3483** | **−3.2725** | | | | | |
| | R² | 0.9998 | 0.9957 | | | | | |
| | Efficacy | 98.91% | 102.10% | | | | | |
| A/Sw/France/22–130212/2013 (H1huN2Δ146-147) | Range of linearity | **<10.98 to 25.64** | **<10.98 to >35.16** | | | **<10.98 to 28.06** | | |
| | Slope  | **−3.281** | **−3.4008** | | | **−3.266** | | |
| | R² | 0.9997 | 0.9992 | | | 0.9988 | | |
| | Efficacy | 101.74% | 96.81% | | | 102.39% | | |
| A/Sw/Flandres/1/1998 (H3N2) | Range of linearity | 16.7 to 32.7 | **<13.13 to 23.57** | | | **<13.13 to 23.57** | | |
| | Slope  | **−3.2964** | **−3.297** | | | **−3.297** | | |
| | R² | 0.9982 | 0.998 | | | 0.998 | | |
| | Efficacy | 101.08% | 101.07% | | | 101.07% | | |

*The range of linearity is given as the interval of Cq-values obtained from M gene RT-qPCR on diluted samples (Thermocycler MxPro – Mx3005P). R²: coefficient of linear regression. HA RT-qPCRs were run as simplex assays; N1 and N2 RT-qPCRs were run in duplex.*
The diagnostic ability of the H1\textsubscript{hu\Delta 146−147} RT-qPCR was evaluated by testing 78 samples of mainly H1N2 or H7N2 subtypes included in panel 4. The H1\textsubscript{hu\Delta 146−147} gene was identified in 3/15 H1\textsubscript{huN2} isolates (not sequenced previously), in 16/46 nasal swab supernatants in which the H1\textsubscript{huN2} virus was identified, as well as in 1/6 H1\textsubscript{huN7} samples (Additional file 2). Although the specificity of the H1\textsubscript{hu\Delta 146−147} was previously demonstrated, these results were further validated by sequencing the HA genes of the three isolates that tested positive for the H1\textsubscript{hu\Delta 146−147} gene as well as the HA gene of strain A/Sw/France/29–150034/15 that tested negative. Sequencing confirmed that the three positive strains contained the 6 nt deletion by contrast to the other one that contained a H1\textsubscript{hu} gene without this deletion (Genbank accession numbers KJ128334, KY241154, KY241115 and KY241115). Thus, 20/78 viruses or clinical samples previously identified as H1\textsubscript{hu}-positive were rapidly identified as having an H1\textsubscript{hu\Delta 146−147} gene.

### Analytical performances of the HA and NA RT-qPCRs in high-throughput analyses

Real-time RT-PCRs for H1\textsubscript{av}, H1\textsubscript{hu}, H1\textsubscript{hu\Delta 146−147}, H3, N1 and N2 gene amplifications, as well as RT-qPCRs designed to amplify the IAV M gene and the porcine \( \beta \)-actin genes were included in order to check the amount and quality of the genetic material and/or demonstrate any potential PCR inhibition. These last RT-qPCRs were previously validated in low-throughput analyses [21, 27]. Thus, 10 RT-qPCR assays were run together on a LightCycler\textsuperscript{1536} to analyze the 118 samples from panel 5, i.e. 83 swIAVs, 5 virus mixtures and 30 nasal swab supernatants, all previously analyzed in low-throughput PCR assays (see Additional file 3). As expected, all samples were detected positive for the M gene. RNA extracts from virus strains exhibited M gene Cq-values ranging from <5 to 16.69, whereas extracts from clinical samples exhibited higher Cq-values, which mostly ranged between 14 and 30. Only 2/118 samples (1.7%) (both RNA extracts from MDCK cell culture supernatants) were found negative for the \( \beta \)-actin gene.

All HA and NA RT-qPCRs exhibited 100% specificity, as no undesirable cross-reaction or false-positive results were obtained, irrespective of the nature of the sample, i.e. RNA extracts from virus stock containing high amounts of genetic material, or RNA extracts from nasal swab supernatants. As expected, the H1\textsubscript{av} RT-qPCR amplified H1 genes originating from the H1\textsubscript{avN1} lineage, i.e. H1 genes from H1\textsubscript{avN1} enzootic viruses and from H1\textsubscript{avN2} reassortants. The H1\textsubscript{hu} RT-PCR amplified H1 genes from H1\textsubscript{huN2} viruses, H1\textsubscript{hu\Delta 146−147} variants and H1\textsubscript{huN1} reassortants. By contrast, the H1\textsubscript{hu\Delta 146−147} RT-PCR only detected the H1\textsubscript{hu\Delta 146−147} antigenic variants. In accordance with the results obtained in low-throughput analyses, it did not detect H1\textsubscript{huN1} reassortants bearing H1\textsubscript{hu} genes with one or two amino acid deletions close to the receptor-binding site (RBS) (called H1\textsubscript{huN1\Delta 146−147} and H1\textsubscript{huN1\Delta 147}). The H1pdm RT-qPCR amplified HA genes of viruses from the H1N1pdm lineage but also any other H1 gene, and the H3 RT-qPCR amplified HA from H3N2 viruses only. The N1 RT-qPCR amplified N1 genes from both the H1\textsubscript{avN1} and the H1N1pdm lineages, whereas the N1pdm RT-qPCR amplified the N1 genes from H1N1pdm viruses specifically. The N2 RT-qPCR amplified NA genes from all HN2 viruses. Thus, all samples were fully subtyped according to expected results, including the virus mixtures, except one of them in which the N2 gene was not amplified in parallel to the H1\textsubscript{av}, H1\textsubscript{hu} and N1 genes, as expected. As observed for the M gene, the Cq-values obtained for the HA and NA RT-qPCRs were lower on RNAs extracted from virus isolates as compared to clinical samples. In several cases, the Cq-value was too low to be precisely determined (<5). A sample without any viral RNA but rather consisting of water was added in each PCR array as a negative control and no unspecific detection was observed (data not shown).

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### Table 5

| Real-time RT-PCR\(^a\) for molecular subtyping | RNA extract from (virus strain – name and subtype) | RNA extract dilution (\( \approx \) M gene Cq-value) | Mean Cq-value | SD | Inter-assay CV (%) |
|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|---------------|----|-------------------|
| H1\textsubscript{av}                           | A/Sw/Cotes d’Armor/0388/2009 (H1\textsubscript{avN1}) | \( 10^{-4} \) (\( \approx 23.83 \)) | 24.50          | 0.9 | 1.20              |
| H1\textsubscript{hu}                           | A/Sw/Scotland/410440/1994 (H1\textsubscript{huN2}) | \( 10^{-3} \) (\( \approx 20.43 \)) | 25.63          | 0.39 | 1.52              |
| H1\textsubscript{hu\Delta 146−147}            | A/Sw/France/22–130212/2013 (H1\textsubscript{huN2\Delta 146−147}) | \( 10^{-4} \) (\( \approx 21.3 \)) | 22.80          | 0.71 | 3.12              |
| H3                                            | A/Sw/Flanders/1/1998 (H3N2)                      | \( 10^{-2} \) (\( \approx 16.7 \)) | 18.60          | 0.31 | 1.68              |
| N1                                            | A/Sw/Cotes d’Armor/0388/2009 (H1\textsubscript{huN1}) | \( 10^{-4} \) (\( \approx 23.83 \)) | 25.64          | 0.37 | 1.46              |
| N2                                            | A/Sw/Scotland/410440/1994 (H1\textsubscript{huN2}) | \( 10^{-3} \) (\( \approx 20.43 \)) | 24.78          | 1.19 | 4.79              |
| N2                                            | A/Sw/Flanders/1/1998 (H3N2)                      | \( 10^{-3} \) (\( \approx 20.21 \))^\(*\) | 28.56          | 1.86 | 6.51              |

\(^a\) HA RT-qPCRs were run as simplex assays; N1 and N2 RT-qPCRs were run in duplex. Cq quantification threshold, SD standard deviation, CV coefficient of variation. Each RNA extract was tested (either on Thermocycler MxPro – Mx3005P or Chromo4) in 6 different assays at the dilution corresponding to 10 times the detection limit of the concerned RT-qPCR (except H3 that was run on a lowest dilution of the H3N2 strain). The RNA extract dilution marked with an asterisk (*) was tested in 5 independent assays.
Table 6 Performance of real-time RT-PCR assays targeting the H1av, H1hu, H3, N1 and N2 genes in porcine clinical samples containing European swIAVs

| Clinical samples (Panel 3) | M gene RT-qPCR (Cq-value) | Real-time RT-PCRs for molecular subtyping* (Cq-value) |
|---------------------------|---------------------------|------------------------------------------------------|
|                           | HA subtyping | H1av | H1hu | H3 | N1 | N2 | NA subtyping |
| H1av                      | N1av         | 120137-4 | 20.6 | 19.19 | no Cq | no Cq | 19.85 | no Cq |
|                           |             | 120183-5 | 18.1 | 19.33 | no Cq | no Cq | 19.25 | no Cq |
|                           |             | 130004-5 | 17.59 | 19.08 | no Cq | no Cq | 17.75 | no Cq |
|                           |             | 130020-4 | 26.43 | 31.38 | no Cq | no Cq | 26.83 | no Cq |
|                           |             | 130028-3 | 26.43 | 22.13 | no Cq | no Cq | 26.98 | no Cq |
|                           |             | 130065-5 | 17    | 23.08 | no Cq | no Cq | 20.73 | no Cq |
|                           |             | 130144-4 | 22.26 | 22.20 | no Cq | no Cq | 22.46 | no Cq |
|                           |             | 130153-6 | 22.46 | 32.33 | no Cq | no Cq | 22.49 | no Cq |
|                           |             | 130157-6 | 27.46 | 24.22 | no Cq | no Cq | 24.25 | no Cq |
|                           |             | 140014-4 | 17.56 | nt    | nt    | nt    | 17.45 | no Cq |
|                           |             | 140016-5 | 19.09 | nt    | nt    | nt    | 19.73 | no Cq |
| H1av                      | N2           | 120057-3 | 17.89 | 17.25 | no Cq | no Cq | no Cq | 19.72 |
|                           |             | 120345-2 | 17.51 | 21.51 | no Cq | no Cq | no Cq | 20.35 |
|                           |             | 130133-3 | 21.18 | 21.67 | no Cq | no Cq | no Cq | 23.54 |
|                           |             | 110214-1 | 28.92 | 27.82 | no Cq | no Cq | nt    | nt    |
| H1hu                      | N2           | 100231-1 | 18.57 | 17.25 | no Cq | 24.79 | no Cq | 21.86 |
|                           |             | 120037-4 | 23.88 | no Cq | 27.79 | no Cq | no Cq | 27.04 |
|                           |             | 120285-6 | 23.37 | no Cq | 25.58 | no Cq | no Cq | 24.45 |
|                           |             | 120424-5 | 21.32 | no Cq | 25.43 | no Cq | no Cq | 24.96 |
|                           |             | 13039-2  | 22.5  | no Cq | 22.66 | no Cq | no Cq | 22.08 |
|                           |             | 130111-5 | 21    | no Cq | 22.79 | no Cq | no Cq | 22.06 |
|                           |             | 130111-6 | 19.6  | nt    | nt    | nt    | no Cq | 22.04 |
|                           |             | 130129-4 | 18.29 | no Cq | 26.63 | no Cq | no Cq | 22.58 |
|                           |             | 130193-6 | 21.95 | no Cq | 25.36 | no Cq | no Cq | 24.11 |
|                           |             | 130129-5 | 17.86 | no Cq | 27.63 | nt    | no Cq | 22.63 |
|                           |             | 130140-5 | 13.74 | no Cq | 20.21 | nt    | no Cq | 16.36 |
|                           |             | 130285-4 | 29.54 | nt    | nt    | nt    | no Cq | 27.57 |
|                           |             | 130420-4 | 22.31 | no Cq | 23.90 | nt    | no Cq | 24.12 |
|                           |             | 130429-6 | 22.42 | no Cq | 24.26 | nt    | no Cq | 21.40 |
|                           |             | 130431-4 | 15.82 | no Cq | 18.31 | nt    | no Cq | 17.41 |
| H1hu                      | N1av         | 110619-8 | 17.28 | no Cq | 21.86 | no Cq | 18.74 | no Cq |
|                           |             | 120067-3 | 27.47 | no Cq | 33.08 | no Cq | 27.47 | no Cq |
| H1av/H1hu                 | N2           | 130103-2 | 21.46 | 28.36 | 23.42 | no Cq | 22.72 | 24.56 |
| H1pdm                     | N1pdm        | 110578-1 | 22.16 | no Cq | no Cq | no Cq | 24.09 | no Cq |
|                           |             | 120158-7 | 16.97 | no Cq | no Cq | no Cq | 19.37 | no Cq |
|                           |             | 120333-1 | 20.08 | no Cq | no Cq | no Cq | 21.70 | no Cq |
|                           |             | 130116-1 | 25.21 | no Cq | no Cq | no Cq | 28.01 | no Cq |
| H3                        | N2           | 120031-1 | 21.83 | no Cq | no Cq | 22.68 | no Cq | 24.46 |
| H1av                      | N?           | 130044-2 | 32.18 | no Cq | no Cq | nt    | 30.37 | no Cq |
|                           |             | 130113-2 | 33.24 | no Cq | no Cq | nt    | no Cq | no Cq |
|                           |             | 130120-3 | 33.91 | 41.43 | no Cq | nt    | 31.64 | no Cq |
In order to go further in evaluating the sensitivity of the methods when run as high-throughput analyses, each RT-qPCR (except the one targeting the β-actin gene) was tested against 10-fold serial dilutions (10$^{-1}$ to 10$^{-8}$) of reference strains (Table 7). Depending on the virus stock and its initial genomic load, the M gene was still detected to the last dilution or to the 10$^{-6}$ or the 10$^{-7}$ dilution. In all cases, it was detected to the 10$^{-5}$ dilution. The H1 av,H1 huΔ146-147 and H1pdm genes were detected in samples exhibiting a corresponding M gene Cq-value until 32, approximately, while the H3 gene detection limit was found to exceed an M gene Cq-value >35. The detection limit for the H1 hu gene varied from 25 to 33 depending on the virus (parental virus, antigenic variant, reassortant or reassortant with deletion), that of N1 from 29 to 31, and that of N2 from 30 to 33. The N1pdm was detected until an M gene Cq-value of 31. The ranges of linearity are given in Table 8. The slopes calculated over these ranges were all (except one) comprised between -3.2 and -3.7, which means the efficacies of the RT-PCRs varied from 85% to 105%. The N2 RT-qPCR run on the reference H3N2 strain was the only one found to be outside these limits, showing an efficacy of 80.51%.

Robustness of the RT-qPCR subtyping tool in low- and high-throughput analyses
The full RT-qPCR subtyping tool began to be used routinely by the French NRL for Swine Influenza from January 2014, as part of the analytical workflow for swIAV surveillance. Thus, nasal swab supernatants previously selected as containing M gene from a swIAV were subjected to H1 av,H1 hu and H3 monoplex assays, to the N1/N2 duplex assay, as well as to commercial H1pdm and N1pdm RT-PCRs. Samples that were found to be positive for H1 hu were subjected to the H1 huΔ146-147 RT-PCR in a second step. Considering the detection limits evaluated above, subtyping was undertaken on samples with M gene Cq-values <35 only. Samples exhibiting M gene Cq-values >35 were considered to be not typable. Looking at the proportions of HA and NA genes that were successfully subtyped in these samples from January 2014 to August 2016, it appeared that both genes were identified in almost all samples with M gene Cq-value <25.

### Table 6 Performance of real-time RT-PCR assays targeting the H1 av,H1 hu, H3, N1 and N2 genes in porcine clinical samples containing European swIAVs (Continued)

| Clinical samples * | swIAV lineage known from conventional RT-PCRs and commercial H1N1pdm RT-qPCRs | Identification | M gene RT-qPCR (Cq-value) | Real-time RT-PCRs for molecular subtyping * (Cq-value) | HA subtyping | NA subtyping |
|-------------------|--------------------------------------------------------------------------------|-----------------|---------------------------|----------------------------------------------------------|---------------|---------------|
|                   |                                                                                |                 |                           | H1 av, H1 hu, H3, N1, N2                                |               |               |
|                   |                                                                                |                 |                           | H1 av                                      | 28.13 | no Cq | nt | 27.85 | no Cq |
|                   |                                                                                |                 |                           | H1 hu                                      | no Cq | no Cq | nt | 24.76 | no Cq |
|                   |                                                                                |                 |                           | H3                                        | 27.33 | no Cq | nt | 28.30 | no Cq |
|                   |                                                                                |                 |                           | N1                                        | 28.30 | no Cq | nt | 24.76 | no Cq |
|                   |                                                                                |                 |                           | N2                                        | 26.51 | no Cq | nt | 25.68 | no Cq |
|                   |                                                                                |                 |                           | N1                                         | no Cq | no Cq | nt | no Cq | no Cq |
|                   |                                                                                |                 |                           | N2                                         | no Cq | no Cq | nt | no Cq | no Cq |
|                   |                                                                                |                 |                           | H7                                         | no Cq | no Cq | nt | no Cq | no Cq |
|                   |                                                                                |                 |                           | N2                                         | 27.58 | nt    | nt | no Cq | 26.27 |
|                   |                                                                                |                 |                           | N1                                         | 26.51 | no Cq | nt | no Cq | 25.68 |
|                   |                                                                                |                 |                           | N2                                         | no Cq | no Cq | nt | no Cq | 24.52 |

*All samples were nasal swab supernatants except the marked one that was a lung sample. *N1av means that the conventional N1 RT-PCR was positive but the N1pdm commercial RT-qPCR was negative. All RT-qPCRs were run as monoplex procedures except N1 and N2 that were run in a duplex. Cq: quantification cycle; nt = not tested. Bold data correspond to additional results obtained with the novel RT-qPCRs as compared to conventional RT-PCRs.
### Table 7: Analytical sensitivity of the real-time RT-PCRs for M, H1av, H1hu, H1huΔ146–147, H1pdm, H3, N1, N1pdm and N2 on ten-fold dilutions of RNA extracted from swIAVs of different subtypes when run on LightCycler®1536

| Virus strain | Lineage | Name | Virus stock dilution | M gene RT-qPCR (Cq-value) | Real-time RT-PCR for molecular subtyping (Cq-value) | H1av | H1hu | H1huΔ146–147 | H1pdm | H3 | N1 | N1pdm | N2 |
|--------------|---------|------|----------------------|--------------------------|--------------------------------------------------|-------|------|----------------|-------|----|-----|-------|----|
| H1avN1       | A/Sw/Cotes d'Armor/0388/2009 | 10^{-1} | 12.01 | 12.53 | no Cq | no Cq | no Cq | no Cq | 12.33 | no Cq | no Cq | no Cq |
|              |         | 10^{-2} | 14.90 | 15.80 | no Cq | no Cq | no Cq | no Cq | 16.02 | no Cq | no Cq | no Cq |
|              |         | 10^{-3} | 19.46 | 19.84 | no Cq | no Cq | no Cq | no Cq | 20.63 | no Cq | no Cq | no Cq |
|              |         | 10^{-4} | 22.83 | 23.14 | no Cq | no Cq | no Cq | no Cq | 24.35 | no Cq | no Cq | no Cq |
|              |         | 10^{-5} | 26.53 | 26.85 | no Cq | no Cq | no Cq | no Cq | 27.11 | no Cq | no Cq | no Cq |
|              |         | 10^{-6} | 29.52 | 29.72 | no Cq | no Cq | no Cq | no Cq | 30.24 | no Cq | no Cq | no Cq |
|              |         | 10^{-7} | 32.81 | 32.51*| no Cq | no Cq | no Cq | no Cq | 33.74*| no Cq | no Cq | no Cq |
|              |         | 10^{-8} | 35.45 | no Cq | no Cq | no Cq | no Cq | no Cq | 35.34*| no Cq | no Cq | no Cq |
| H1avN2       | A/Sw/Cotes d'Armor/0186/2010 | 10^{-1} | 15.01 | 15.14 | no Cq | no Cq | no Cq | no Cq | 14.32 | no Cq | no Cq | no Cq |
|              |         | 10^{-2} | 18.14 | 18.43 | no Cq | no Cq | no Cq | no Cq | 18.04 | no Cq | no Cq | no Cq |
|              |         | 10^{-3} | 21.72 | 21.96 | no Cq | no Cq | no Cq | no Cq | 22.34 | no Cq | no Cq | no Cq |
|              |         | 10^{-4} | 25.34 | 25.27 | no Cq | no Cq | no Cq | no Cq | 26.12 | no Cq | no Cq | no Cq |
|              |         | 10^{-5} | 28.91 | 27.95 | no Cq | no Cq | no Cq | no Cq | 29.51 | no Cq | no Cq | no Cq |
|              |         | 10^{-6} | 32.76 | 31.62 | no Cq | no Cq | no Cq | no Cq | 32.40 | no Cq | no Cq | no Cq |
|              |         | 10^{-7} | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | 34.31*| no Cq | no Cq | no Cq |
|              |         | 10^{-8} | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq |
| H1huN2       | A/Sw/Cotes d'Armor/0113/2006 | 10^{-1} | 14.52 | no Cq | 16.11 | no Cq | no Cq | no Cq | 15.00 | no Cq | no Cq | no Cq |
|              |         | 10^{-2} | 17.45 | no Cq | 19.33 | no Cq | no Cq | no Cq | 17.60 | no Cq | no Cq | no Cq |
|              |         | 10^{-3} | 22.48 | no Cq | 23.92 | no Cq | no Cq | no Cq | 22.71 | no Cq | no Cq | no Cq |
|              |         | 10^{-4} | 25.87 | no Cq | 27.60 | no Cq | no Cq | no Cq | 26.40 | no Cq | no Cq | no Cq |
|              |         | 10^{-5} | 28.49 | no Cq | 30.10*| no Cq | no Cq | no Cq | 28.89 | no Cq | no Cq | no Cq |
|              |         | 10^{-6} | 33.49 | no Cq | 33.58*| no Cq | no Cq | no Cq | 32.04*| no Cq | no Cq | no Cq |
|              |         | 10^{-7} | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | 34.76 | no Cq | no Cq | no Cq |
|              |         | 10^{-8} | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq |
| H1huN1Δ147   | A/Sw/France/22–130212/2013 | 10^{-1} | 8.59  | 11.90 | 10.61 | no Cq | no Cq | no Cq | 8.58  | no Cq | no Cq | no Cq |
|              |         | 10^{-2} | 12.08 | no Cq | 15.01 | 13.69 | no Cq | no Cq | 13.22 | no Cq | no Cq | no Cq |
|              |         | 10^{-3} | 15.54 | no Cq | 18.64 | 18.07 | no Cq | no Cq | 16.69 | no Cq | no Cq | no Cq |
|              |         | 10^{-4} | 19.05 | no Cq | 22.40 | 21.91 | no Cq | no Cq | 20.91 | no Cq | no Cq | no Cq |
|              |         | 10^{-5} | 22.85 | no Cq | 25.88 | 26.00 | no Cq | no Cq | 24.98*| no Cq | no Cq | no Cq |
|              |         | 10^{-6} | 25.44 | no Cq | 29.72 | 29.21 | no Cq | no Cq | 27.95 | no Cq | no Cq | no Cq |
|              |         | 10^{-7} | 29.78 | no Cq | 32.37*| 32.17*| no Cq | no Cq | 31.09 | no Cq | no Cq | no Cq |
|              |         | 10^{-8} | 32.73 | no Cq | 36.51 | no Cq | no Cq | no Cq | 34.43 | no Cq | no Cq | no Cq |
| H1huN1Δ147   | A/Sw/Cotes d'Armor/0070/2010 | 10^{-1} | 14.85 | no Cq | 18.39 | no Cq | no Cq | no Cq | 16.85 | no Cq | no Cq | no Cq |
|              |         | 10^{-2} | 18.11 | no Cq | 21.66 | no Cq | no Cq | no Cq | 20.45 | no Cq | no Cq | no Cq |
|              |         | 10^{-3} | 22.90 | no Cq | 26.08 | no Cq | no Cq | no Cq | 25.07 | no Cq | no Cq | no Cq |
|              |         | 10^{-4} | 25.71 | no Cq | 28.73 | no Cq | no Cq | no Cq | 27.57 | no Cq | no Cq | no Cq |
|              |         | 10^{-5} | 29.68 | no Cq | 31.39 | no Cq | no Cq | no Cq | 30.95 | no Cq | no Cq | no Cq |
|              |         | 10^{-6} | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | 34.42 | no Cq | no Cq | no Cq |
|              |         | 10^{-7} | 33.47*| no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq |
|              |         | 10^{-8} | No Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq | no Cq |
For $25 \leq \text{Cq-value} <30$, the proportion of unidentified HA and/or NA genes increased slightly. In these samples, the N1/N2 multiplex appeared less sensitive than HA simplex assays, as NA genes were less frequently amplified. In samples with Cq-value $>30$, the proportion of viruses fully or partially subtyped fell to about 50%.

Finally, the diagnostic ability of the high-throughput subtyping RT-qPCR assay was tested on RNAs extracted from 919 nasal swabs collected during a longitudinal study conducted in three pig herds located in Brittany and previously selected as M gene-positive samples [28]. These herds were known to be affected by H1 avN1 and H1 huN2 viruses only. Among them, HA and NA genes were both identified in 697 samples, whereas only the HA or NA gene was detected in 77 others, depending on the apparent amount of virus genome in the samples (Table 9). Almost all samples with an M gene Cq-value $<30$ were fully subtyped (Table 9). In samples with an M gene Cq-value $>30$, the NA gene was more frequently amplified than the HA gene (Table 9). Fourteen samples from one herd were shown to contain virus mixtures and/or reassortant viruses (data not shown).

**Discussion**

Eight real-time RT-PCRs were evaluated for European swIAV molecular subtyping in clinical samples demonstrated to contain IAV genome. They aimed to discriminate HA genes for four H1 genetic lineages (H1 av, H1 hu, H1 huΔ146–147, H1pdm), and one H3 lineage, as well as NA genes of two N1 lineages (N1, N1pdm) and one N2 lineage. Altogether, the RT-qPCRs enabled the identification of swIAVs from the four viral subtypes known to be enzootic in European pigs, i.e. H1 avN1, H1 huN2, H3N2 and H1N1pdm. They also made it possible to quickly identify a new antigenic variant (H1 huN2Δ146–147) among H1 huN2 viruses, as well as reassortant viruses, i.e. H1 huN1 or H1 avN2, and virus mixtures. All assays were optimized to take into account the genetic diversity

| Virus strain | Lineage | Name | Virus stock dilution | M gene RT-qPCR (Cq-value) | Real-time RT-PCR for molecular subtyping (Cq-value) |
|--------------|---------|------|----------------------|--------------------------|--------------------------------------------------|
|              |         |      |                      | H1 av | H1 hu | H1 huΔ146–147 | H1pdm | H3 | N1 | N1pdm | N2 |
| A/Sw/Cotes d’Armor/0190/2006 | H1N1pdm | A/Sw/Sarthe/0255/2010 | A/Sw/Flandres/1/1998 | 10^−1 | 13.31 | no Cq | no Cq | no Cq | 19.30 | no Cq | 17.24 | 16.38 |
| 10^−2 | 17.24 | no Cq | 20.87 | no Cq | no Cq | 19.93 | no Cq | no Cq |
| 10^−3 | 20.86 | no Cq | 23.92 | no Cq | no Cq | 23.79 | no Cq | no Cq |
| 10^−4 | 25.01 | no Cq | 28.30 | no Cq | no Cq | 27.43 | no Cq | no Cq |
| 10^−5 | 28.21 | no Cq | no Cq | no Cq | no Cq | 30.37 | no Cq | no Cq |
| 10^−6 | 31.31 | no Cq | no Cq | no Cq | no Cq | 34.50* | no Cq | no Cq |
| 10^−7 | no Cq | no Cq | no Cq | no Cq | no Cq | 35.28* | no Cq | no Cq |
| 10^−8 | no Cq | no Cq | no Cq | no Cq | no Cq | 36.13* | no Cq | no Cq |
| 10^−9 | no Cq | no Cq | no Cq | no Cq | no Cq | 38.50* | no Cq | no Cq |
| 10^−10 | no Cq | no Cq | no Cq | no Cq | no Cq | 39.38* | no Cq | no Cq |
| 10^−11 | no Cq | no Cq | no Cq | no Cq | no Cq | 40.38* | no Cq | no Cq |

Cq quantification cycle. Cq-values are mean Cq-values between duplicates except numbers indicated with an asterisk that were obtained from a single sample (*). Cq-values highlighted in grey represent detection limits.
encountered among swIAVs isolated in France in recent years, whereas keeping in mind usefulness on an European level. They exhibited a gain in sensitivity as compared to conventional RT-PCRs, allowing the characterization of biological samples with low genetic loads, with considerable time saving. Reagents and amplification procedures were harmonized to run all assays in parallel, using a unique amplification cycle profile irrespective of the equipment.

Evaluation in low-throughput analyses showed perfect specificities for H1av, H1hu and H3 RT-qPCRs conducted in parallel as simplex assays. While multiplexing may facilitate the diagnostic procedure, preliminary studies showed that individual RT-qPCR sensitivities were somewhat affected when run in a triplex assay (data not shown). Based on M gene Cq-values, RT-qPCRs appeared slightly less sensitive than assays involving MGB-labelled probes, such as previously validated commercial kits or

| Virus strain | Criteria | M gene RT-qPCR<sup>a</sup> | Real-time RT-PCR for molecular subtyping |
|--------------|----------|-----------------------------|-----------------------------------------|
| A/Sw/Cotes d’Armor/0388/2009 (H1<sub>hu</sub>N1) | Range of linearity<sup>b</sup> | <12.01 to 35.45 | <12.01 to 35.45 |
| | Slope | −3.423 to −3.386 | −3.356 |
| | R<sup>2</sup> | 0.9963 to 0.9971 | 0.9914 |
| | Efficacy | 95.95% to 97.38% | 98.61% |
| A/Sw/Cotes d’Armor/0186/2010 (H1<sub>hu</sub>N2) | Range of linearity<sup>b</sup> | <15.01 to 32.76 | <15.01 to 32.76 |
| | Slope | −3.5616 to −3.2659 | −3.674 |
| | R<sup>2</sup> | 0.9992 to 0.9988 | 0.9962 |
| | Efficacy | 90.89% to 102.39% | 87.15% |
| A/Sw/Cotes d’Armor/0113/2006 (H1<sub>hu</sub>N2) | Range of linearity<sup>b</sup> | <14.52 to 28.49 | <14.5 to 33.49 |
| | Slope | −3.637 to −3.5244 | −3.5025 |
| | R<sup>2</sup> | 0.9988 to 0.9938 | 0.9966 |
| | Efficacy | 88.34% to 92.19% | 92.98% |
| A/Sw/France/22–130212/2013 (H1<sub>hu</sub>N2<sub>Δ146–147</sub>) | Range of linearity<sup>b</sup> | <14.59 to 32.73 | <14.59 to 32.73 |
| | Slope | −3.4642 to −3.3586 | −3.6888 |
| | R<sup>2</sup> | 0.9989 to 0.9992 | 0.9958 |
| | Efficacy | 94.39% to 90.06% | 92.52% |
| A/Sw/Cotes d’Armor/0070/2010 (H1<sub>hu</sub>N1) | Range of linearity<sup>b</sup> | <14.85 to 29.68 | <14.85 to 29.68 |
| | Slope | −3.726 to −3.307 | −3.481 |
| | R<sup>2</sup> | 0.9952 to 0.9903 | 0.995 |
| | Efficacy | 85.52% to 106.1% | 93.76% |
| A/Sw/Cotes d’Armor/0190/2006 (H1<sub>hu</sub>N1<sub>Δ147</sub>) | Range of linearity<sup>b</sup> | <13.49 to 31.31 | <13.49 to 31.31 |
| | Slope | −3.6041 to −3.724 | −3.6323 |
| | R<sup>2</sup> | 0.9979 to 0.9958 | 0.9984 |
| | Efficacy | 89.44% to 85.58% | 88.50% |
| A/Sw/Sarthe/0255/2010 (H1N1pdm) | Range of linearity<sup>b</sup> | <13.31 to 33.50 | <13.31 to 33.50 |
| | Slope | −3.5216 to −3.4394 | −3.4324 |
| | R<sup>2</sup> | 0.9927 to 0.9939 | 0.9987 |
| | Efficacy | 92.92% to 95.32% | 99.59% |
| A/Sw/Flandres/1/1998 (H1N2) | Range of linearity<sup>b</sup> | <12.96 to 34.01 | <12.96 to 34.01 |
| | Slope | −3.485 to −3.505 | −3.8987 |
| | R<sup>2</sup> | 0.9977 to 0.9931 | 0.9942 |
| | Efficacy | 93.62% to 90.06% | 80.51% |

<sup>a</sup>The range of linearity is given as the interval of Cq-values obtained from M gene RT-qPCR on diluted samples. <sup>b</sup>R<sup>2</sup> Coefficient of linear regression
Table 9: Percentages of HA and NA genes identified in nasal swab supernatants with M gene Cq-value <35, either in low-throughput analyses (routine diagnosis performed by the French NRL, January 2014–August 2016) or in high-throughput analyses (longitudinal survey in three herds), according to the M gene Cq-values

| Range of M gene Cq-values | Number of samples | Proportion of HA identified | Proportion of NA identified |
|---------------------------|-------------------|-----------------------------|-----------------------------|
| 30 ≤ Cq < 35              | 87                | 52.88%                      | 20.56%                      |
| 25 ≤ Cq < 30              | 151               | 93.22%                      | 79.19%                      |
| 20 ≤ Cq < 25              | 198               | 97.00%                      | 99.14%                      |
| < 20                      | 106               | 100%                        | 99.22%                      |

| Range of M gene Cq-values | Number of samples | Proportion of HA identified | Proportion of NA identified |
|---------------------------|-------------------|-----------------------------|-----------------------------|
| 30 ≤ Cq < 33              | 189               | 12.17%                      | 33.86%                      |
| 25 ≤ Cq < 30              | 273               | 89.38%                      | 96.34%                      |
| 20 ≤ Cq < 25              | 350               | 98.86%                      | 100%                        |
| < 20                      | 106               | 100%                        | 100%                        |

*Samples tested for H1 av, H1 hu, H1pdm, H3, N1, N1pdm and N2. *Samples tested for H1 av, H1 hu, N1 and N2 only as herds were previously known to be affected solely by H1 N1 and H1 N2 viruses.

other in-house methods aimed at amplifying the M, H1pdm or N1pdm genes [21]. Nevertheless, the detection levels were satisfactory as HA subtyping was nearly 100% successful in clinical samples up to M gene Cq-value = 30. The H1 hu RT-qPCR was the least efficient, probably due to higher genetic variability among H1 hu genes, as compared to the H1 av and H3 genes [9]. The H1 huN1–147 RT-qPCR, run in a second step on H1 huNy-positive samples also demonstrated excellent specificity and sensitivity, leading to rapid discrimination of novel antigenic variants among H1 huN2 viruses. While H1 huN1 reassortants are rare events in most European countries, they are sporadically detected in France [8, 29] and some of them may also have an H1 huN1–147 gene. In this study, two H1 huN1 reassortants exhibiting 2 amino acid deletions at positions 146–147 of the RBS, thus bearing H1 genes genetically and antigenically closer to the H1 huN2 variant than to the parental H1 hu, were not detected by the H1 huN1–147 RT-PCR due to a mismatch within the “H1 huN1 var” MGB probe. As a result, classification of the H1 hu gene from H1 huN1 reassortants as a “A146–147 variant” (or not) would be better confirmed by HA sequencing.

N1 and N2 RT-qPCRs, adapted from protocols previously evaluated at the European level, confirmed the specificities expected from in silico analyses [20]. While designed to amplify N2 genes from both European H1 huN2 and H3N2 lineages, the N2 RT-PCR proved to detect N2 genes from the novel H1 huN2 A146–147 variants. By contrast to HA RT-qPCRs, they were run in a duplex assay because preliminary studies indicated no reduction in sensitivity as compared to corresponding simplex assays (data not shown). Nevertheless, it should be noted that in routine diagnosis, the NA gene was less frequently subtyped than the HA gene in samples with an M gene Cq-value >30, leading to the hypothesis that this difference could be related, at least partially, to multiplexing.

Scaling M and β-actin RT-qPCRs as well as HA (H1 av, H1 hu, H1 huN1–147, H1pdm, H3) and NA (N1, N1pdm, N2) subtyping assays to higher throughput, with concurrent miniaturization of individual reactions, was successful. The LightCycler®1536 system designed for automated high-throughput laboratory workflows proved in this context to provide very high well-to-well homogeneity, good inter-assay reproducibility and low inter-plate variability. The 10 RT-qPCRs, run together onto one microplate using the same amplification procedure, showed very good efficiency. Each RT-qPCR retained its specificity. They all exhibited comparable and acceptable sensitivities. When run on clinical samples with an M gene Cq-value <30, the techniques were able to fully subtype more than 90% of detected swIAV genomes, showing an increased proportion of characterized NA genes in samples with Cq-values of 25–30 as compared to equivalent samples using classical thermocyclers. By contrast, when run on clinical samples with an M gene Cq-value >30, H1 av, H1 hu, N1 and N2 RT-PCRs appeared slightly less efficient than in low-throughput analyses. This could be related to the low volume of RNA extract included into the RT-qPCR mixture when run on LightCycler®1536 system as compared to other thermocyclers (1 μL instead of 5 μL), but this would need to be further investigated and confirmed for other subtyping RT-qPCRs.

Conclusion

The emergence of the pandemic A/H1N1 virus of swine origin in 2009 highlighted the need for global surveillance of influenza A viruses in pigs. Its subsequent introduction into the pig population, its co-circulation with other enzootic swIAVs, and genomic reassortment events have led to an increase in the genetic diversity of swIAVs. Altogether, these RT-qPCR assays provide a rapid and simple genotyping method to identify viruses that infect the European pig population, including a novel H1 huN2 variant identified in France, sporadic HA/NA reassortants and virus mixtures, as a first characterization step before full genome sequencing and/or antigenic subtyping. Screening of individual samples against the 10 target genes in a high-throughput scenario opens novel perspectives in diagnostic abilities, which will be very useful for swIAV surveillance and large-scale epidemiological studies.
Additional files

**Additional file 1:** Specificity of the real-time RT-PCR developed to identify H1_{nu}Δ146-147 antigenic variants among H1_{nu}N_{2} swIAVs (panel 2). (DOCX 18 kb)

**Additional file 2:** Identification of new antigenic variants among H1_{nu}N_{2} swIAVs by real-time RT-PCR targeting the H1_{nu}Δ146-147 gene. (DOCX 22 kb)

**Additional file 3:** Specificity of real-time RT-PCRs for detection and subtyping of swIAVs when run simultaneously as simplex assays on LightCycler®1536. (DOCX 61 kb)

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Availability of data and materials

All relevant information is provided in this current manuscript.

Authors’ contributions

GS, SH and PF designed the project. EB and TH drew the primers and the probes. EB, SQ, CW, SG and NB performed the experiments. EB, SQ, SG, SH and GS analyzed and interpreted the results. EB and GS wrote the manuscript. All authors revised the manuscript, worked together for its improvement, and approved the final version.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

None of the authors of this paper have a financial or personal relationship with other people or organizations that could inappropriately influence or bias the content of the paper. The authors declare that they have no competing interests.

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