False-positive PCR results linked to administration of seasonal influenza vaccine

T. Curran,1 C. McCaughey,1 J. Ellis,2 S. J. Mitchell,1 S. A. Feeney,1 A. P. Watt,1 F. Mitchell,1 D. Fairley,1 L. Crawford,1 J. McKenna1 and P. V. Coyle1

Correspondence
T. Curran
tanya.curran@belfasttrust.hscni.net

1Regional Virus Laboratory, Microbiology Dept, Kelvin Building, Royal Victoria Hospital, Belfast BT12 6BA, UK
2Respiratory Virus Unit, Virus Reference Department, Microbiology Services, Health Protection Agency, Colindale, London NW9 5HT, UK

False-positive PCR results usually occur as a consequence of specimen-to-specimen or amplicon-to-specimen contamination within the laboratory. Evidence of contamination at the time of specimen collection linked to influenza vaccine administration in the same location as influenza sampling is described. Clinical, circumstantial and laboratory evidence was gathered for each of five cases of influenza-like illness (ILI) with unusual patterns of PCR reactivity for seasonal H1N1, H3N2, H1N1 (2009) and influenza B viruses. Two 2010 trivalent influenza vaccines and environmental swabs of a hospital influenza vaccination room were also tested for influenza RNA. Sequencing of influenza A matrix (M) gene amplicons from the five cases and vaccines was undertaken. Four 2009 general practitioner (GP) specimens were seasonal H1N1, H3N2 and influenza B PCR positive. One 2010 GP specimen was H1N1 (2009), H3N2 and influenza B positive. PCR of 2010 trivalent vaccines showed high loads of detectable influenza A and B RNA. Sequencing of the five specimens and vaccines showed greatest homology with the M gene sequence of Influenza A/Puerto Rico/8/1934 H1N1 virus (used in generation of influenza vaccine strains). Environmental swabs had detectable influenza A and B RNA. RNA detection studies demonstrated vaccine RNA still detectable for at least 66 days. Administration of influenza vaccines and clinical sampling in the same room resulted in the contamination with vaccine strains of surveillance swabs collected from patients with ILI. Vaccine contamination should therefore be considered, particularly where multiple influenza virus RNA PCR positive signals (e.g. H1N1, H3N2 and influenza B) are detected in the same specimen.

INTRODUCTION

High-throughput molecular testing is increasingly becoming the mainstay of clinical virology laboratories, with many laboratories solely employing amplification-based techniques for viral respiratory infection diagnosis. Many virology laboratories delivering molecular influenza diagnosis during the 2009–2010 pandemic, and in the following 2010–2011 influenza season, employed an automated and algorithmic approach to testing, facilitating faster and greater clinical specimen throughput.

During the 2009–2010 influenza pandemic the Regional Virus Laboratory, Belfast, rationalized screening of respiratory specimens from patients presenting with influenza-like illness (ILI) meeting the case definition to influenza A [matrix (M) gene], H1N1 (2009) [haemagglutinin (HA) gene] and the housekeeping gene RNase P (to determine specimen quality). During the winter of 2010–2011, a broader approach to initial influenza screening was required, due to the co-circulation of influenza with other respiratory viruses causing ILI. The 2010–2011 algorithm tested for influenza A (M gene), H1N1 (2009), influenza B, respiratory syncytial virus (RSV) and RNase P. Samples with discrepant results were further tested with a subtyping panel of real-time PCR assays targeting H1N1 (2009) neuraminidase (NA), seasonal H1N1 and H3N2 HA.

The vast majority of samples received from patients with ILI during the pandemic and the 2010–2011 influenza season tested positive for one influenza subtype. However in both seasons, a small number of specimens collected from community cases of ILI, were positive for both influenza A H3N2 and influenza B in addition to either seasonal H1N1 or H1N1 (2009) viruses, giving rise to concerns of specimen contamination during laboratory testing. The repeated
amplification of the same target sequence, resulting in accumulation of amplified product within the laboratory environment, can be a major source of contamination in molecular testing, and multiple control methods are routinely employed to control this (Aslanzadeh, 2004; Borst et al., 2004; Hartley & Rashitchian, 1993). However, further laboratory analysis of these samples and influenza vaccine, in addition to environmental swabbing of an influenza vaccination area, confirmed source contamination.

Here we report this novel cause of contamination and discuss its significance.

**METHODS**

**Clinical samples.** During January 2009 to February 2011, 11 033 specimens were tested for influenza in the Regional Virus Laboratory, Belfast. Both combined nasal and throat swabs in addition to sputum samples in either Nuclisens lysis buffer (bioMérieux) in 2009–2010 or eNat medium (Copan) in 2010–2011 were received. Specimens were received from hospitals and general practitioner (GP) practices in Northern Ireland and also from the 29 GP practices participating in the Northern Ireland GP sentinel surveillance scheme.

**Molecular analysis.** Following nucleic acid extraction of 200 μl sample using the QIAsymphony platform (Qiagen), specimens were tested by real-time Taqman PCR according to the seasonal testing algorithms. These assays included generic influenza A (M gene), influenza B (HA gene), H1N1 (2009) (HA gene), seasonal H1N1 (HA gene) and H3N2 (HA gene) (Health Protection Agency, 2009a, d, e; van Elden et al., 2001). The 10 μl PCRs consisted of 1 x Superscript III Platinum One step qRT-PCR (Invitrogen) with 0.4 μM primers, 0.4 μM Taqman probe and 2 μl extracted specimen. All real-time assays were carried out on a LightCycler 480 real-time machine (Roche) using the following cycling conditions: 50 °C 15 min, 95 °C 5 min, followed by 45 cycles of 95 °C 10 s and 60 °C 1 min. Positive and negative controls were included in each PCR assay run. Specimens with crossing threshold (Ct) values ≤ 40 were reported as positive.

Specimens in which an unusual pattern of PCR positivity for multiple targets was observed were repeat tested. Any specimens with the same unusual pattern of positivity were selected for further investigation. Specimens were re-extracted, restested in all real-time PCR assays in addition to two H5N1 specific assays (Health Protection Agency, 2009b, c; WHO, 2007) and underwent sequence analysis.

**Detection of RNA in influenza vaccine.** Trivalent 2010 vaccine from two manufacturers (Wyeth and Sanofi Pasteur), were obtained from the hospital Pharmacy and Occupational Health departments. Both trivalent vaccines contained an A/California/7/2009 (H1N1)-like virus, an A/Perth/16/2009 (H3N2)-like virus and a B/Brisbane/60/2008-like virus as recommended by the World Health Organization.

Two hundred and fifty microlitres of each were extracted using the QIAasyphony platform (Qiagen) and subsequently tested for influenza A and B viral RNA.

**Sequencing.** Partial sequencing of the M gene was carried out on two of the specimens (14701/09 and 12821/10) with unusual patterns of PCR positivity, in addition to both vaccines, using the BigDye Terminator v3.1 Cycle sequencing kit (Applied Biosystems). Primers targeting a 153 bp region were used: forward primer 5’-CTTGTATAT-TGGGATTCTTGA-3’ and reverse primer 5’-ATCGTAACATCCA-CAGCA-3’ (primer positions 813–833 and 948–966 respectively on the M gene: GenBank accession number CY009317). Products were purified with the Montage Seq® sequencing reaction cleanup kit (Millipore) prior to sequencing on a 3100 capillary sequencer (Applied Biosystems). Sequences were analysed for homology using NCBI BLAST analysis. Additional PCR assays (not shown) targeting longer regions of the influenza M gene failed to generate products for sequencing.

The other three specimens were referred to the Respiratory Virus Unit, Health Protection Agency, Colindale, for further sequence analysis. Viral RNA was extracted from specimens using the NucliSens easyMag system (bioMérieux). Viral RNA was amplified using primers targeting the influenza A M gene and the amplicons were directly sequenced using the PCR primers AMF Forward (5’-GAGTCTTCTA-ACMGAGGTTGAAGCTA-3’, nucleotides 17–43 in CY009317) and AMR Reverse (5’-GGGCACGGTGAGCGTRAA-3’, nucleotides 204–222) (Ellis & Curran, 2011). PCR clean-up and sequencing reactions were performed by the Genomic Services Unit at the Department for Bioanalysis and Horizon Technologies, Health Protection Agency. PCR amplicons were prepared for sequencing by using Ampure Magnetic beads on the Biomek NxP robot (Beckman Coulter). Sequencing was performed as described, with clean-up of products with CleanSeq magnetic beads on the Biomek robot and automated sequence detection on a 48-capillary ABI 3730 genetic analyser.

**Environmental swabbing.** Thirty-eight environmental swabs were taken in October 2010 in a Belfast hospital Occupational Health vaccination area following a single session where 123 health care workers had received influenza trivalent vaccine. All swabs were pre-moistened in tap water. A premoistened-only swab was also tested as a negative control. Swabs were set down on tables in the area in addition to streaking the swabs over various lengths (ranging from 2 cm to 1 m). Each surface was individually swabbed four times. Swabs were immediately placed in 500 μl Nuclisens lysis buffer, nucleic acid extracted and tested for influenza.

**RNA surface survival time.** An RNA survival experiment was undertaken to estimate the stability and durability of influenza vaccine RNA on surfaces. A 1 in 1000 dilution of the Wyeth vaccine was made in tap water, applied evenly using cotton wool onto a laminate flooring material and allowed to air dry, stored in a non-laboratory domestic setting. Over the subsequent 9 weeks, a swab of a 4 cm trail was taken at each of eight time points. These swabs were placed in Nuclisens lysis buffer, extracted and tested by real-time PCR as described.

**RESULTS**

**Cases**

During the period from June 2009 to February 2011, a total of 11 033 specimens were tested for influenza A, 8187 of which were also tested for influenza B (a subset of samples; influenza B results were tested due to rationalization of the 2009–2010 testing algorithm for hospital and community specimens). Nineteen per cent (2105/11 033) of specimens were positive for influenza A, of which 18 % (385/2105) were received from sentinel influenza GP practices and 82 % (1720/2105) from non-sentinel GP practices and hospitals. The prevalence of influenza B from 2009–2011 was 3 % (227/8187), of which 14 % (32/227) were received from sentinel influenza GP practices and 86 % (195/227) from non-sentinel GP practices and hospitals. Six specimens were positive for both H1N1 (2009) and influenza B.

During 2009–2011, an unusual pattern of influenza A H1N1 (2009), H3N2, seasonal H1N1 and influenza B virus detection...
by PCR was observed in five specimens, which were re-extracted and repeat tested (two H5N1 real-time assays were also included in this repeat testing). The results are shown in Table 1. Four specimens collected and tested in 2009 were positive for seasonal H1N1, H3N2 and influenza B, but negative for H1N1 (2009) and H5N1 viruses. A fifth specimen collected and tested in 2010 was positive for H1N1 (2009), H3N2 and influenza B viruses. This specimen was negative for seasonal H1N1 and H5N1 viruses. The age of the patients ranged from 16 to 76 years, with a 3 : 2 ratio of women to men. The 2009 specimens were from patients attending two different GP practices. Specimens 13584/09 and 13810/09 were taken 1 day apart in September 2009 in GP practice 1. Specimens 14136/09 and 14710/10 were taken 7 days apart the following month in GP practice 2. Specimen 12821/10 was taken in September 2010 in GP practice 3. None of the five patients had received that year’s influenza vaccine prior to or on the day of sampling. There was no social or geographical link between the cases. All specimens had been taken in areas in the practices where both clinical sampling and influenza vaccination were regularly carried out.

Detection of RNA in influenza vaccine

Both 2010 trivalent vaccines were extracted and tested in all influenza Taqman real-time PCR assays. Influenza RNA from both trivalent vaccines was detected in all of the real-time PCR assays except those for detection of seasonal H1N1 and H5N1. Ct values ranging from 11.79 to 13.79 were observed, representing a significant load, in the order of an estimated 10^{10} copies ml^{-1} of target (based on general extrapolation of Ct 28 equivalent to 10^{5} copies ml^{-1} and a log value equatable to 3.32 Ct).

Sequencing

Amplicons from PCR targeting part of the M gene of specimens 14710/09 and 12821/10 in addition to both trivalent influenza vaccines were sequenced. BLAST analysis on NCBI showed the M gene partial sequences from these specimens to have 100% homology to the M gene sequence of A/Puerto Rico/8/1934. Sequence analysis of PCR products generated following amplification of a different region of the influenza A M gene from specimens 13584/09, 13810/09 and 14136/09 generated similar results (Fig. 1).

Environmental swabbing

Environmental swabs from the occupational health vaccination area were tested and the results are shown in Table 2. Influenza RNA was detectable in all swabs in all assays except those for seasonal H1N1 and H5N1.

RNA surface survival time

Eight swabs corresponding to eight different time points were taken of diluted (1 : 1000) vaccine that had been applied and air-dried onto a piece of laminate floor material. The results are shown in Table 3. Diluted vaccine RNA remained relatively static over all time points, and was still detectable at day 66.

DISCUSSION

The 2009 influenza pandemic was the first pandemic to occur since the advent of molecular techniques (McCaughey, 2010). This molecular approach to testing has allowed the introduction of and reliance on a wide range of rapid and sensitive virology assays. However, PCR contamination is often a consequence of amplicon-to-specimen and occasionally specimen-to-specimen contamination within the laboratory (Speers, 2006).

Five specimens collected and tested during 2009 and 2010 demonstrated unusual patterns of influenza virus RNA positivity; the 2009 specimens were positive for seasonal H1N1, H3N2 and influenza B but negative for H1N1

### Table 1. Real-time PCR results of five cases tested 2009–2010

<table>
<thead>
<tr>
<th>Virus (gene target)</th>
<th>13584/09†</th>
<th>13810/09†</th>
<th>14136/09†</th>
<th>14710/09†</th>
<th>12821/10‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Influenza A (M)</td>
<td>34.20</td>
<td>33.97</td>
<td>32.13</td>
<td>36.89</td>
<td>35.07</td>
</tr>
<tr>
<td>Influenza B (HA)</td>
<td>35.67</td>
<td>35.83</td>
<td>39.18</td>
<td>37.17</td>
<td>31.25</td>
</tr>
<tr>
<td>H1N1 (2009) (HA)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>36.28</td>
</tr>
<tr>
<td>H1N1 (2009) (NA)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>34.02</td>
</tr>
<tr>
<td>Seasonal H1N1 (HA)</td>
<td>33.05</td>
<td>32.79</td>
<td>32.81</td>
<td>37.29</td>
<td>–</td>
</tr>
<tr>
<td>H3N2 (HA)</td>
<td>34.24</td>
<td>34.12</td>
<td>34.48</td>
<td>37.84</td>
<td>33.96</td>
</tr>
<tr>
<td>H5N1 (HA)§</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

*–, Negative.
†2009 specimens.
‡2010 specimen.
§Combined results for both H5N1 assays.
(2009), while the 2010 specimen showed a different pattern, being H1N1 (2009), H3N2 and influenza B positive but seasonal H1N1 negative. This appeared to mirror the changes made to the influenza trivalent vaccine over the two years. All specimens were collected in September/October of 2009 and 2010, coinciding with the beginning of the Northern Ireland seasonal influenza vaccination programme; none of the patients had been vaccinated at the time of sampling. All five cases had specimens taken at three different GP practices as part of the Northern Ireland sentinel surveillance programme.

Partial sequencing of the influenza A M gene from the specimens and vaccines showed $\geq 99\%$ homology with A/Puerto Rico/8/1934 reassortant viruses, used to generate high-yield reassortant strains in trivalent influenza vaccines. PCR and sequencing of product was successful where only short regions were targeted. Sequencing of

![Fig. 1. Sequence alignments of PCR products with different regions of the M gene (GenBank accession number CY009317): (a) primer positions 813–833 and 948–966; (b) primer positions 17–43 and 204–222.](http://jmm.sgmjournals.org)
longer regions of the M gene proved unsuccessful, which may reflect fragmentation of the vaccine RNA.

The most credible explanation of these results is carry-over or detection of vaccine RNA. A recent publication described the contamination of diagnostic swabs in horses who had previously received influenza vaccination (Diallo et al., 2011). While this has not previously been reported for inactivated human influenza vaccines, recipients of intranasal live attenuated vaccine can shed influenza RNA for up to 3 weeks post vaccination and give rise to positive results when tested by PCR (Freed et al., 2007). In addition, contamination of specimens with pertussis vaccine has been documented (Leber et al., 2010). Surprisingly high loads (in the order of 10^10 copies ml^-1) were detectable in both trivalent vaccines. The effect of such high RNA loads in these influenza vaccines has not been documented in the literature. Single-stranded H5N1 RNA has been shown to stimulate Toll-like receptors, increasing the efficiency of antigen presentation (Geeraedts et al., 2008). Therefore vaccine RNA may have an important role in immunogenicity and efficacy of the vaccine. However the presence of such high concentrations of RNA may contribute to the well-documented systemic side effects associated with influenza vaccine, due to cytokine production initiated by the vaccine RNA (Julkunen et al., 2000).

### Table 2. Real-time PCR results of environmental swabbing of occupational health vaccination area

<table>
<thead>
<tr>
<th>Swab Description</th>
<th>C&lt;sub&gt;t&lt;/sub&gt; values*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Set down Table 1†</td>
<td>36.42 ± 0.67 36.36 ± 0.72</td>
</tr>
<tr>
<td>2 2 cm trail Table 1§</td>
<td>34.56 ± 0.82 32.36 ± 1.60</td>
</tr>
<tr>
<td>3 10 cm trail Table 1§</td>
<td>35.46 ± 0.73 33.19 ± 1.04</td>
</tr>
<tr>
<td>4 1 m trail Table 1§</td>
<td>33.43 ± 2.09 31.28 ± 2.76</td>
</tr>
<tr>
<td>5 2 cm trail chair§</td>
<td>36.97 ± 0.35 36.58 ± 1.37</td>
</tr>
<tr>
<td>6 20 cm trail Table 2$</td>
<td>35.56 ± 2.34 30.80 ± 2.81</td>
</tr>
<tr>
<td>7 Floor$</td>
<td>36.11 ± 1.29 37.28 ± 3.24</td>
</tr>
<tr>
<td>8 Tables 2 and 3$</td>
<td>33.86 ± 2.92 30.29 ± 2.82</td>
</tr>
<tr>
<td>9 Extensive Tables 1–3$</td>
<td>31.71 ± 3.41 29.09 ± 3.66</td>
</tr>
<tr>
<td>10 Vaccinated arm#</td>
<td>30.08 ± 2.16 27.12 ± 1.98</td>
</tr>
<tr>
<td>11 Negative control**</td>
<td>–</td>
</tr>
</tbody>
</table>

*Data are means ± sd of four individual swabs. –, Negative.
†Combined result for both H5N1 assays.
§Swabs set down on Table 1 for a few seconds.
$Swabs moved 2 cm, 10 cm, 20 cm or 1 m along Table 1, Table 2 or chair.
||Swabs trailed along floor or Tables 2 and 3.
§Swabs vigorously trailed backwards and forwards for several metres along Table 1, Table 2 and Table 3.
#Skin swabs of vaccinated arm taken immediately post-vaccination.
**Swab moistened with same nuclease-free water as used to premoisten each swab used prior to sampling.

### Table 3. Vaccine RNA surface survival time (C<sub>t</sub> values)

<table>
<thead>
<tr>
<th>Assay</th>
<th>1 h</th>
<th>10.5 h</th>
<th>24 h</th>
<th>1.5 days</th>
<th>3 days</th>
<th>9 days</th>
<th>20 days</th>
<th>66 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Influenza A (M)</td>
<td>36.83</td>
<td>35.69</td>
<td>36.20</td>
<td>35.35</td>
<td>35.44</td>
<td>35.57</td>
<td>34.79</td>
<td>–</td>
</tr>
<tr>
<td>Influenza B (HA)</td>
<td>37.54</td>
<td>35.64</td>
<td>37.01</td>
<td>35.98</td>
<td>35.30</td>
<td>–</td>
<td>35.13</td>
<td>–</td>
</tr>
<tr>
<td>H1N1 (2009) (HA)</td>
<td>39.06</td>
<td>35.38</td>
<td>35.79</td>
<td>34.36</td>
<td>34.72</td>
<td>–</td>
<td>35.37</td>
<td>36.77</td>
</tr>
<tr>
<td>H1N1 (2009) (NA)</td>
<td>38.17</td>
<td>–</td>
<td>37.44</td>
<td>37.51</td>
<td>37.45</td>
<td>–</td>
<td>–</td>
<td>35.82</td>
</tr>
<tr>
<td>Seasonal H1N1 (HA)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>H3N2 (HA)</td>
<td>&gt;40</td>
<td>37.43</td>
<td>38.18</td>
<td>36.83</td>
<td>35.86</td>
<td>–</td>
<td>37.07</td>
<td>37.75</td>
</tr>
<tr>
<td>H5N1† (HA)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

*, –, Negative.
†Combined result for both H5N1 PCR assays.
The level of RNA in these vaccines clearly has potential to generate environmental contamination. Specimens from all five cases came from GP practices that routinely used the same treatment room for clinical sampling and influenza vaccination. The fact that H1N1 (2009), H3N2 and influenza B (but not seasonal H1N1) RNA was detected in the environmental swabs collected during 2010 from the occupational health vaccination area and the ability to detect vaccine influenza RNA for at least 66 days indicates a significant risk for environmental contamination and subsequent contamination of clinical specimens collected from such locations.

Previous studies have documented the potential risk of false-positive nucleic acid amplification test results when specimens are taken in an area that is contaminated with target DNA (Meader et al., 2008). Therefore it is surprising that this phenomenon appears not to have been recognized for influenza previously. Possibly individual laboratories dismiss these types of results as erroneous because multiple targets are positive. It is important to realize early that vaccine contamination is a potential explanation, especially where there are multiple influenza virus positive results (H1N1, H3N2 and influenza B) in the same specimen. In addition, laboratories employing algorithmic approaches to testing are unlikely to recognize vaccine contamination, leading to incorrect clinical diagnoses, unnecessary laboratory work and unsound epidemiological conclusions.

Controlling this type of contamination would involve a few simple measures. Ideally, separate areas should exist for vaccination and clinical sampling. Staff administering vaccine should wear protective clothing (aprons and gloves) and should not leave the vaccination area. Opened swabs should not be set on surfaces either before or after swabbing. Dropped swabs should be repeated rather than sent to the laboratory. Surfaces in treatment rooms should be regularly cleaned after vaccine use. Previous studies have shown that the use of common household cleaning agents such as 1% bleach, 10% malt vinegar and antiviral wipes has demonstrated that the use of common household cleaning agents regularly cleaned after vaccine use. Previous studies have documented the potential risk of preventing PCR contamination in a clinical laboratory. (Ann Clin Lab Sci 34, 389–396).


Leber, A. & other authors (2010). Vaccines shown to contain PCR-detectable DNA include Pentacel®, Daptacel®, and Adacel®. Detection of Bordetella pertussis DNA in acellular vaccines and in environmental samples from pediatric physician offices. Interscience Conference on Antimicrobial Agents and Chemotherapy (ICAAAC): Boston, USA.

ACKNOWLEDGEMENTS

The authors would like to thank staff at the Health Protection Agency, in particular Richard Allan in the Respiratory Virus Unit, for technical support.

REFERENCES


Leber, A. & other authors (2010). Vaccines shown to contain PCR-detectable DNA include Pentacel®, Daptacel®, and Adacel®. Detection of Bordetella pertussis DNA in acellular vaccines and in environmental samples from pediatric physician offices. Interscience Conference on Antimicrobial Agents and Chemotherapy (ICAAAC): Boston, USA.


