Surveillance for highly pathogenic influenza A viruses in California during 2014–2015 provides insights into viral evolutionary pathways and the spatiotemporal extent of viruses in the Pacific Americas Flyway

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We used surveillance data collected in California before, concurrent with, and subsequent to an outbreak of highly pathogenic (HP) clade 2.3.4.4 influenza A viruses (IAVs) in 2014–2015 to (i) evaluate IAV prevalence in waterfowl, (ii) assess the evidence for spill-over infections in marine mammals and (iii) genetically characterize low-pathogenic (LP) and HP IAVs to refine inference on the spatiotemporal extent of HP genome constellations and to evaluate possible evolutionary pathways. We screened samples from 1496 waterfowl and 1142 marine mammals collected from April 2014 to August 2015 and detected IAV RNA in 159 samples collected from birds (n = 157) and pinnipeds (n = 2). HP IAV RNA was identified in three samples originating from American wigeon (Anas americana). Genetic sequence data were generated for a clade 2.3.4.4 HP IAV-positive diagnostic sample and 57 LP IAV isolates. Phylogenetic analyses revealed that the HP IAV was a reassortant H5N8 virus with gene segments closely related to LP IAVs detected in mallards (Anas platyrhynchos) sampled in California and other IAVs detected in wild birds sampled within the Pacific Americas Flyway. In addition, our analysis provided support for common ancestry between LP IAVs recovered from waterfowl sampled in California and gene segments of reassortant HP H5N1 IAVs detected in British Columbia, Canada and Washington, USA. Our investigation provides evidence that waterfowl are likely to have played a role in the evolution of reassortant HP IAVs in the Pacific Americas Flyway during 2014–2015, whereas we did not find support for spill-over infections in potential pinniped hosts.

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INTRODUCTION

Highly pathogenic (HP) H5 influenza A viruses (IAVs) of the Goose Guangdong lineage were first detected in China in 1996.1 During the subsequent two decades, viral descendants that share ancestry at the hemagglutinin gene segment have evolved through reassortment with viruses of numerous neuraminidase (NA) subtypes, resulting in the emergence and intercontinental spread of novel HP IAVs designated as belonging to clade 2.3.4.4.2–10 In autumn 2014, Goose Guangdong lineage clade 2.3.4.4 HP IAVs were detected in wild, captive and domestic birds in North America.5,6,11 Purportedly, clade 2.3.4.4 HP H5N8 IAVs were introduced into North America via migratory waterfowl.11 Although definitive proof for this hypothesis remains elusive, this proposed mechanism of introduction is consistent with (i) intercontinental migratory movements of potential waterfowl hosts,12–14 (ii) previous evidence for viral dispersal between East Asia and North America by wild birds,15 (iii) the widespread detection of HP H5N8 and descendant viruses in wild birds in the Pacific Americas Flyway during 2014–2015,16 (iv) a lack of pathobiological effects or barriers to infection in experimentally inoculated waterfowl17 and (v) the circulation of low-pathogenic (LP) viruses in wild waterfowl sampled in Alaska that share recent common ancestry with reassortant viruses descended from Eurasian lineage HP H5N8.18,19 Therefore, it is likely that waterfowl contributed to the introduction and dissemination of clade 2.3.4.4 HP IAVs in North America.

While surveillance for IAVs in waterfowl sampled in the Pacific Flyway has provided information on the prevalence and evolution of

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In this study, we explore the prevalence, evolution and spread of HP clade 2.3.4.4 IAVs in the Pacific Americas Flyway by using surveillance data collected in California before, concurrent with and subsequent to the previously reported outbreak of 2014–2015. Specifically, we evaluate the prevalence of HP clade 2.3.4.4 IAVs in waterfowl relative to previous reports and assess the evidence for spill-over infections in pinnipeds. Furthermore, we genetically characterize both LP and HP IAVs recovered through our sampling effort to refine inference on the spatiotemporal extent of HP clade 2.3.4.4 genome constellations in California during the outbreak and to evaluate possible evolutionary pathways. Our results supplement previous summaries of surveillance efforts for clade 2.3.4.4 HP IAVs in North America and extend inference relative to viral evolution. This information may be useful to better understand the role of wildlife in the maintenance and evolution of clade 2.3.4.4 IAVs in North America. Furthermore, our results may inform surveillance strategies and biosecurity recommendations for regulatory agencies, poultry producers and other stakeholders that are concerned with HP IAVs in the United States of America.

MATERIALS AND METHODS

Ethics statement

Sampling procedures that involved live animals as reported in this study were reviewed by the Institutional Animal Care and Use committees at the University of California (UC), the California State University and the US Geological Survey (USGS) (the UC permits No. 18034 and No. 19559; the California State University permit No. 2014-2; and the USGS permit WERC-2015-01), and were conducted under authorizations granted by the following permits: US Fish and Wildlife Service Special Use Permit No. 81620-14-009; US Fish and Wildlife Service Scientific Collection permit No. MB191637-0; USGS Federal Bird Banding Permits No. 21142 and No. 22911; California Department of Fish and Wildlife Scientific Collection Permits No. 3855, No. 5000 and No. 8090; and National Oceanic and Atmospheric Administration National Marine Fisheries Service Marine Mammal Protection Act permit No. 18786-01.

Sampling

Cloacal swabs, oropharyngeal swabs and/or fecal samples were collected from live waterfowl in Solano County, California, from July through August in both 2014 and 2015, and from hunter-harvested waterfowl in Solano and Butte counties, California, from October 2014 through January 2015 (Figure 1). Live birds were captured in baited swim-in traps or by using rocket nets or were sampled by swabbing freshly deposited feces. Hunter-harvested waterfowl samples were collected at check stations. Swabs were collected using sterile polyester-tipped applicators, placed in vials containing 2 mL of cold virus transport medium (VTM: Medium 199 with Earle’s salts, L-glutamine and sodium bicarbonate plus 2 mU/L penicillin G, 200 mg/L streptomycin, 2 mU/L polymyxin B, 250 mg/L gentamicin, 0.5 mU/L nystatin, 60 mg/L ofloxacin, 200 mg/L sulphamethoxazole and 0.5% bovine serum albumin), and transported on ice to the laboratory where they were stored at −80 °C on the day of collection until processing. Swabs were placed individually in vials with VTM except for 129 samples from live-sampled birds, which contained pooled cloacal and oropharyngeal swabs.

From April 2014 to August 2015, pinnipeds that stranded along the Central Coast of California, as identified through the California Marine Mammal Stranding Network, and that were hospitalized for rehabilitation at The Marine Mammal Center in Sausalito, California, were sampled for IAVs as part of routine health examinations upon entry to the rehabilitation facility (Figure 1). Species sampled included California sea lions (Zalophus californianus), Guadalupe fur seals (Arctocephalus townsendi), northern elephant seals (Mirounga angustirostris), northern fur seals (Callorhinus ursinus), Pacific harbor seals (Phoca vitulina) and Steller sea lions (Eumetopias jubatus). Nasal and rectal swabs were collected from each animal, and the swabs were placed individually in vials containing VTM. Samples were refrigerated up to 1 week before shipping to UC Davis. Upon receipt at the UC Davis laboratory, samples were either further processed immediately or stored at −80 °C.

Screening for IAVs

Viral RNA was extracted from all swab samples using MagMAX-96 Al/ND Viral RNA Isolation Kits (Ambion/Applied Biosystems, Foster City, CA, USA). RNA was screened using a two-step real-time reverse transcriptase-PCR (rRT-PCR) that targeted the IAV matrix gene.26 Any sample that did not yield a cycle threshold (Ct) value ≤45 was considered negative for IAV. Matrix gene rRT-PCR-positive samples were further screened using a HP clade 2.3.4.4 H5-specific rRT-PCR that utilized the following primers and probe: H5 EA+1462 (5′-TGG GTA ATG GTT GTT TCG AG-3′); H5 EA-1565 (5′-ATT GCT TCT TCT GAA TAC TTA GGG-3′) and H5EA 1487 probe (6-FAM) 5′-TCA CAA ATG/ZEN/TGA TAA CGA ATG TAT GGA TCT TCT GAA TTT TGT GAG C-3′ (IBFQ). The AgPath ID RT-PCR kit (Life Technologies, Carlsbad, CA, USA) and the AB 7500 Fast instrument (Life Technologies) were used to screen for HP clade 2.3.4.4 H5 IAV by applying the following thermocycling protocol: 45 °C for 10 min; 95 °C for 10 min; and then 40 cycles of 94 °C for 10 s, 57 °C for 10 s and 72 °C for 5 s. VTM samples that were identified as putatively positive for HP clade 2.3.4.4 H5 IAVs were sent to the California Health and Food Animal Safety Laboratory and to the US Department of Agriculture National Veterinary Services Laboratory for further testing. Virus isolation was attempted on all matrix-positive, HP H5 rRT-PCR-negative samples by

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Virus isolation was attempted on all matrix-positive, HP H5 rRT-PCR-negative samples by
inoculating 200 μL of VTM into the allantoic cavities of two 11-day-old embryonated chicken eggs (Charles River, North Franklin, CT, USA) and incubating at 37.8 °C for 3 days or until embryo death. Allantoic fluid from each pair of eggs was collected and pooled, and 50 μL of this fluid was processed for RNA extraction and rRT-PCR testing for the IAV matrix gene target as described above. Virus isolates were defined as any sample that yielded allantoic fluid with a Ct value ≤ 45 for the IAV matrix gene after being cultured in specific pathogen-free eggs.

### Sequencing of virus isolates

Full-genome sequencing was attempted on RNA that was extracted from all virus isolates at either UC Davis, the Icahn School of Medicine at Mount Sinai (ISMMS) or the USGS Alaska Science Center. For genomic sequencing at UC Davis and ISMMS, sequencing was attempted on RNA extracts that were provided by UC Davis for 31 viral isolates and three diagnostic original swab samples for which Ct values ≤ 45 were obtained by rRT-PCR screening for HP clade 2.3.4.4 H5 IAVs. For each isolate, 2 μL of RNA template was amplified using SuperScript III One-Step RT-PCR System with Platinum Taq High Fidelity (Invitrogen) and the following primers: MBTuni-12′-ACG CGT GAT CAG CAA AAG CAG G-3′ at 0.1 μM, MBTuni-12(M) 5′-ACG CGT GAT CAG CRA AAG CAG G-3′ at 0.1 μM and MBTuni-13 5′-ACG CGT GAT CAG TAG AAA CAA GG-3′ at 0.2 μM. Thermocycling conditions were 42 °C for 60 min; 5 cycles of 94 °C for 15 s, 45 °C for 30 s and 68 °C for 3 min; and 30 cycles of 94 °C for 15 s, 57 °C for 30 s and 68 °C for 3 min. Following product visualization and verification (5 μL on a 1.0% agarose gel), excess deoxynucleotide triphosphates and primers were removed using ExoSAP-IT (USB Corporation, Cleveland, OH, USA). PCR products were quantified using a Quant-iT dsDNA HS Assay Kit (Invitrogen) and prepared for sequencing using a Nextera XT DNA library preparation kit (Illumina, Inc., San Diego, CA, USA). Indexed libraries were pooled and sequenced on an Illumina MiSeq using a 500-cycle
Table 1. Summary of surveillance sampling of wild waterfowl and pinnipeds in California for HP clade 2.3.4.4 H5 influenza A viruses in wild waterfowl during 2014–2015

<table>
<thead>
<tr>
<th>Species</th>
<th>Pre-outbreak (April–August 2014)</th>
<th>Outbreak (September 2014–March 2015)</th>
<th>Post-outbreak (April–August 2015)</th>
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<tr>
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<td>n</td>
<td>MA+ (%)</td>
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<td>Waterfowl*</td>
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<td>American green-winged teal (Anas crecca)</td>
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<td>American wigeon (Anas americana)</td>
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<td>Bufflehead (Bucephala albeola)</td>
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<td>Common goldeneye (Bucephala clangula)</td>
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<td>Gadwall (Anas strepera)</td>
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<td>Mallard (Anas platyrhynchos)</td>
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<td>Northern pintail (Anas acuta)</td>
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<td>Northern shoveler (Anas clypeata)</td>
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<td>Ruddy duck (Oxyura jamaicensis)</td>
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<td>Marine mammals*</td>
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<td>California sea lion (Zalophus californianus)</td>
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<td>Nasal swabs</td>
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<td>Rectal swabs</td>
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<td>Pacific harbor seal (Phoca vitulina richardi)</td>
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<td>Nasal swabs</td>
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Abbreviations: highly pathogenic, HP; clade 2.3.4.4 HP H5 or intercontinental lineage A positive samples, icA+; matrix-positive samples, MA+; not tested, NT; number of samples collected, n; virus isolation positive samples, VI+.

*All waterfowl samples are swabs collected from the cloaca only unless otherwise specified.

**MA+ defined as Ct value ≤ 45 using real-time reverse transcriptase-PCR (rRT-PCR) for matrix gene target (Runstadler et al.26).

***icA+ defined as having Ct value < 40 using rRT-PCR for clade 2.3.4.4 hemagglutinin gene target (see Materials and Methods).

†VI+ defined as collected allantoic fluid with Ct value ≤ 45 using rRT-PCR for matrix gene target (Runstadler et al.26).

‡Negative rRT-PCR results targeting matrix gene product were not included in the table for the following species: American coot (Fulica americana; n = 2); Blue-winged teal (Anas discors; n = 1); Cackling goose (Branta hutchinsii; n = 1); Canada goose (Branta canadensis); cloacal swab only (n = 1); Cinnamon teal (Anas cyanoptera; n = 6); Gadwall (Anas strepera); combined cloacal/oropharyngeal swab (n = 1); Greater white-fronted goose (Anser albifrons; n = 19); Lesser scaup (Aythya affinis; n = 3); Mallard (Anas platyrhynchos; oropharyngeal swab only (n = 2); fecal swab (n = 25); Ring-necked duck (Aythya collaris; n = 11); Ross's goose (Chen rossii; n = 1); Snow goose (Chen caerulescens; n = 5); and Wood duck (Aix sponsa; n = 8).

§Diagnostic icA+ samples not subjected to VI at UC Davis; therefore, results were reported excluding icA+ samples.

∥Not all samples were subjected to virus isolation on account of logistical considerations (e.g., insufficient sample volume).

(1) Negative rRT-PCR results targeting matrix gene product were not included in the table for paired nasal and rectal swab samples (processed separately) for the following species: Guadalupe fur seal (Arctocephalus townsendi; n = 24); northern elephant seal (Mirounga angustirostris; n = 175); northern fur seal (Callorhinus ursinus; n = 30); and Steller sea lion (Eumetopias jubatus; n = 1).
reagent kit with paired-end reads. Sequence reads were assembled with Geneious 9.1.334 using reference data for IAVs that were obtained from GenBank35 to map reads. Consensus sequences were verified using FlAnotation. The GenBank accession numbers for 28 complete and 3 partial IAV genomes that were sequenced at the USGS Alaska Science Center as part of this study are KY828600–KY828841.

**Phylogenetic analyses**

IAV sequences that originated from samples collected in East Asia (Japan, South Korea and Taiwan) and North America (Canada and United States) from 2010 to 2016 were downloaded from the National Center for Biotechnology Information Influenza Virus Resource database in December 2016. Only full-length or near-full-length genomic segments that were associated with a full date of collection (DD/MM/YYYY) were included. Sequences for IAVs that were generated as part of this study were aligned with reference sequences using MUSCLE v3.8.31,30 inspected visually in Geneious v7.1.5 (North America Biomatters Inc., Newark, NJ, USA) and trimmed to remove nucleotides that were outside the coding region.

Maximum-likelihood trees using the GTR+GAMMA substitution model in RAxML v8.1.1631 were reconstructed for each segment by using publicly available sequences and the sequences that were generated for this study, as described above (sequences per segment: PB2 = 1749; PB1 = 1806; PA = 1802; H5 = 372; NP = 1735; N1 = 248; N2 = 335; N8 = 602; M = 1719; and NS = 1760). Trees were visualized in FigTree v1.4.3 (http://tree.bio.ed.ac.uk/software/figtree/) and randomly down-sampled to reduce the number of sequences for Bayesian molecular clock phylogenetic analysis (~300 taxa). The downsampling strategy preserved all clade 2.3.4.4 isolates from East Asia and North America, and concurrently circulating LP IAV isolates from the Pacific Americas Flyway that originated from samples collected from 2014 to 2016 (PB2 = 345, PB1 = 353, PA = 340, H5 = 338, NP = 366, N1 = 248, N2 = 355, N8 = 306, M = 362 and NS = 370).

Bayesian phylogenetic trees for each segment were reconstructed using BEAST v1.8.3,32,33 At least four independent markov chain Monte Carlo chains of 40–80 million generations each were run using a GMRF Bayesian skyline coalescent tree prior and uncorrelated lognormal distribution to produce 10 000 trees. To ensure that markov chain Monte Carlo chains converged on the same optimal tree, runs were visually inspected in Tracer v1.6.0 (http://tree.bio.ed.ac.uk/software/tracer/). After removing 10%–30% burn-in from each chain, the trees were combined to produce a single maximum clade credibility tree per genomic segment.

**Molecular dating analysis**

The time of most recent common ancestry (tMRCA) and 95% highest posterior densities (95% HPDs) for clades of interest (see Results) was inferred from the nodes of the maximum clade credibility tree as inspected in FigTree v1.4.3. We also considered the posterior probabilities that were associated with each branch node when estimating timing of ancestry. Estimates of tMRCA were only generated for nodes with a posterior probability ≥ 0.90. Estimates of tMRCA for bifurcating nodes with lower support (<0.90) were not calculated, as these phylogenetic relationships were considered to be unresolved.

**RESULTS**

**IAV detection and subtypes of isolates**

From April 2014 through August 2015, we screened 3779 swab samples that had been collected in California from 2638 individual animals (1496 waterfowl and 1142 pinnipeds) for IAVs (Table 1). Samples originated from three sampling periods that were defined by our sampling methodology and related to the previously described outbreak of HP clade 2.3.4.4 H5 IAV in the Pacific Americas Flyway: pre-outbreak (April–August 2014); outbreak (September 2014–March 2015); and post-outbreak (April–August 2015).

IAV RNA was detected in 157 waterfowl samples (10.5% of birds sampled; 157/1496) and two swab samples that had been collected from pinnipeds (0.2% of pinnipeds sampled; 2/1142; Table 1). HP clade 2.3.4.4 H5 IAV RNA was identified in three cloacal swab samples (0.2% of birds sampled; 3/1496) collected in Butte (n = 2) and Solano (n = 1) counties originating from hunter-harvested American wigeon (Anas americana) sampled on 17 January 2015 (Table 1). When only considering samples that were collected from 20 December 2014 to 1 February 2015, the same time period that was used in a previous summary of surveillance efforts for HP clade 2.3.4.4 IAVs in the Pacific Americas Flyway,16 we detected RNA for clade 2.3.4.4 IAVs in 3/374 samples (0.8%) collected from hunter-harvested waterfowl (data not shown). Whole-genome sequencing was successfully performed on RNA that had been extracted from a single diagnostic sample, and was verified as belonging to a HP clade 2.3.4.4 IAV of the H5N8 subtype (strain name: A/ American wigeon/California/UCD58P/2015(H5N8)).

A total of 86 LP IAV isolates were recovered from waterfowl that had been sampled throughout the entire April 2014–August 2015 sampling period, while none were isolated from either nasal or rectal swab samples collected from pinnipeds. From the 86 LP IAV isolates that originated from waterfowl, we amplified and genetically sequenced 57 partial or complete IAV genomes. Sequencing indicated that the three isolates for which genetic data were obtained represented mixed infections. The combined subtypes of 51 LP IAVs for which we obtained full genomes and for which we did not detect evidence of mixed infections included H1N1 (n = 7), H2N5 (n = 1), H2N6 (n = 1), H3N8 (n = 5), H4N6 (n = 10), H4N8 (n = 3), H5N2 (n = 3), H6N1 (n = 1), H7N3 (n = 2), H7N4 (n = 1), H8N4 (n = 1), H10N2 (n = 1) and H11N9 (n = 15). The predominate combined subtype of LP IAV genomes from isolates recovered from waterfowl samples collected during the pre-outbreak period was H1N1 (6/9 or 67%), during the outbreak period was H5N2 (3/10 or 30%) and during the post-outbreak period was H11N9 (15/32 or 47%).

**Phylogenetic analyses**

Phylogenetic analyses revealed that sequences for all eight gene segments of strain A/American wigeon/California/UCD58P/2015 (H5N8) were clustered in monophyletic clades with other HP clade 2.3.4.4 IAVs that had been detected in western North America during 2014–2015 (Figures 2 and 3; Supplementary Figures S1–S8). However, PB1 and PA gene segment sequences for A/American wigeon/California/UCD58P/2015(H5N8) were both of North American lineages, in contrast to other HP clade 2.3.4.4 H5N8 IAVs that had been previously reported from California, A/chicken/California/15-004912/2015(H5N8) and A/turkey/California/K1500169-1.2/2015 (H5N8), which were composed of completely Eurasian origin gene segments (Figures 2 and 3; Supplementary Figures S1–S8). The PB1 gene segment sequences for strains A/American wigeon/California/UCD58P/2015(H5N8), A/American wigeon/Oregon/AH0012525/2015 (H5N8) and A/Canada goose/Oregon/AH0012452/2015(H5N8) formed a monophyletic clade that shared recent common ancestry with 14 H11Nx LP IAVs (including a mixed infection) and a single H11N9 LP IAV that circulated among mallards (Anas platyrhynchos) that had been sampled in California as part of this study (posterior probability = 1.00; tMRCA: 13 June 2014, 95% HPD: 28 October 2013–14 November 2014; Figure 2; Supplementary Figure S2).
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Figure 2 Bayesian phylogenetic tree depicting the inferred genetic relationship among PB1 gene segment sequences for isolates that originated from wild waterfowl in California during the period from 2014 to 2015, highly pathogenic (HP) clade 2.3.4.4 H5N8 viruses (red), LP viruses (black), and reassortant HP H5N8 viruses (light blue). The bracket indicates the portion of the phylogeny that is presented in panel B. Inferred evolutionary distance is indicated by the scale bar. (A) A partial PB1 phylogeny to depict inferred genetic ancestry among strain A/American wigeon/California/UDC585P/2015 (H5N8), other reassortant HP H5N8 viruses that have been detected in North America and other LP influenza A viruses that were isolated in California as part of this study or elsewhere as part of concurrent surveillance efforts. Color is used to differentiate branch tips for HP reassortant H5N8 viruses (light blue) and LP viruses (black). Asterisks are used to indicate genetic sequences that were produced as part of the current study. Blue bars at nodes depict 95% highest posterior densities, which can be interpreted relative to the provided timescale. The posterior probability for the node depicting inferred common ancestry between reassortant HP H5N8 influenza A viruses and LP viruses detected in mallards sampled in California as part of this study is shown. Refer to Supplementary Figure S2 for the complete phylogenetic tree with tip labels.

Similarly, these three reassortant HP clade 2.3.4.4 H5N8 IAVs formed a monophyletic clade in our phylogeny for the PA gene segment, sharing common ancestry with four reassortant HP clade 2.3.4.4 H5N1 IAVs that had been detected in British Columbia, Canada and Washington, USA, as well as eight H11Nx LP IAVs (including a mixed infection) that had been detected in mallards sampled in California for this study (posterior probability = 1.00; tMRCA: 25 May 2014, 95% HPD: 22 November 2013–8 October 2014; Figure 3; Supplementary Figure S3). This strongly supported clade of PA gene segment sequences also appeared to share ancestry with sequences for LP H1N1 IAVs that had been detected in mallards in Alaska during autumn 2014; however, a Bayesian posterior probability (0.72) that was lower than our a priori threshold precluded us from inferring that this relationship was resolved with confidence and thus estimating the tMRCA (Figure 3).

Numerous other gene segments for LP IAVs that were recovered from waterfowl samples collected in California during 2014–2015 shared relatively recent common ancestry with HP clade 2.3.4.4 reassortant viruses previously reported in North America. The PB1 gene of A/mallard/California/360/2014(H4N6) shared relatively recent common ancestry with viruses that had been recovered in Alaska in September 2014 and reassortant HP clade 2.3.4.4 H5N1 IAVs recovered in British Columbia, Canada and Washington, USA in December 2014–February 2015 (posterior probability = 1.00; tMRCA: 22 October 2013, 95% HPD: 23 May 2013–27 February 2014; Figure 4; Supplementary Figure S2). Similarly, strain A/gadwall/California/COL039/2014(H6N1) shared recent common ancestry at the N1 NA gene segment with other H6N1 LP IAVs that had been detected in Alaska in 2015 and HP clade 2.3.4.4 reassortant H5N1 IAVs isolated in British Columbia, Canada, and Washington, USA, during the 2014–2015 outbreak (posterior probability = 0.99; tMRCA: 5 December 2013, 95% HPD: 26 March 2013–13 July 2014; Figure 4; Supplementary Figure S9). N5 gene segment sequences for two isolates recovered from an American wigeon (A/American wigeon/California/LS257/2014(H5N2)) and a northern shoveler (Anas clypeata; A/northern shoveler/California/LDC188/2014(H5N2)) also clustered in strongly supported clades with LP H5N2 subtype IAVs that had been recovered from waterfowl sampled in Alaska that shared common ancestry with reassortant HP clade 2.3.4.4 H5N1 IAVs isolated in British Columbia, Canada, and Washington, USA, during the 2014–2015 outbreak (posterior probability = 1.00; tMRCA: 9 January 2014, 95% HPD: 8 February 2014–6 September 2014; Figure 4; Supplementary Figure S8). The isolate A/northern shoveler/California/LDC188/2014(H5N2) also appeared to be genetically closely related at the N2 NA gene segment to HP clade 2.3.4.4 H5N2 IAVs isolated from wild and domestic birds throughout the Pacific Northwest and Central USA during the 2014–2015 outbreak; however, a relatively low Bayesian posterior probability (0.31) precluded us from inferring that this relationship was resolved with confidence and estimating the tMRCA (Supplementary Figure S10). Gene segment sequences for other IAVs that had been isolated from waterfowl as part of this study clustered with those for IAVs recovered from wild birds throughout the Pacific Americas Flyway and other areas of North America (Figures 2 and 3; Supplementary Figures S1–S10).
Figure 3 Bayesian phylogenetic trees depicting the inferred genetic relationship among PA gene segment sequences for isolates originating from wild waterfowl in California from 2014–2015, highly pathogenic (HP) clade 2.3.4.4 H5 influenza A viruses detected in North America during the concurrent outbreak and other reference sequences for wild bird low-pathogenic (LP) influenza A viruses isolated from North America and East Asia. (A) The complete Bayesian phylogeny. Eurasian viral lineages are color-coded and identified as follows: red = intercontinental lineage A HP H5N8 that was first introduced into North America; light blue = reassortant HP H5N8; dark blue = reassortant HP H5N1; and green = reassortant HP H5N2. The bracket indicates the portion of the phylogeny that is presented in panel B. Inferred evolutionary distance is indicated by the scale bar. (B) Partial PA phylogeny to depict inferred genetic ancestry among strain A/ American wigeon/California/UCD58P/2015(H5N8), other reassortant HP H5N1 and H5N8 viruses detected in North America and other LP influenza A viruses that were isolated in California as part of this study or elsewhere as part of concurrent surveillance efforts. Color is used to differentiate among branch tips for HP reassortant H5N1 (dark blue), HP reassortant H5N8 (light blue) and LP influenza A viruses (black). Asterisks are used to indicate genetic sequences produced as part of the current study. Blue bars at nodes depict 95% highest posterior densities, which can be interpreted relative to the provided timescale. The posterior probability of a node that depicts inferred common ancestry between reassortant HP H5N1 and HP H5N8 influenza A viruses with LP viruses detected in mallards sampled in California as part of this study is shown. Refer to Supplementary Figure S3 for the complete phylogenetic tree with tip labels.

DISCUSSION

In this study, we consistently detected LP IAVs in waterfowl sampled in California during each of three sampling periods during April 2014–August 2015 and found a relatively low apparent prevalence of HP H5 clade 2.3.4.4 IAVs in wild waterfowl sampled in California, identifying clade 2.3.4.4 IAV RNA in three samples that were collected from American wigeon on 17 January 2015. These results are consistent with previous findings, including (1) the detection of LP IAVs in waterfowl sampled in California during both breeding and non-breeding periods, (2) the relatively low prevalence of HP H5 clade 2.3.4.4 viruses in waterfowl sampled in the Pacific Americas Flyway during the 2014–2015 outbreak (0.8% in this study during 20 December 2014–1 February 2015 compared with 1.3% reported by Bevins et al.) (3) the approximate seasonal timing at which HP H5 clade 2.3.4.4 IAVs were first and most frequently detected in California (first detected during the non-breeding season in 2014–2015, specifically 28 December 2014, and most commonly detected 10–17 January 2017) and (4) the most common host (American wigeon) among hunter-harvested waterfowl in which HP IAVs were identified during 20 December 2014–1 February 2015. Thus, our evaluation of the prevalence of HP clade 2.3.4.4 IAVs in waterfowl relative to previous reports supports prior conclusions regarding the proportion of sampled waterfowl hosts that were infected during the outbreak and the likely timing of introduction into California. With regard to the latter relationship, we acknowledge that our interpretation is based on limited sampling of waterfowl in California during the pre-outbreak period.

Pinnipeds that stranded at coastal areas of North America have previously been found to be positive for avian-origin IAVs. However, despite sampling a large number of animals that stranded along the Central Coast of California during 2014–2015, we found limited evidence of exposure to LP IAVs and no evidence for infection with HP H5 clade 2.3.4.4 IAVs. It is unclear whether this result reflects a lack of exposure of marine mammals in California to HP IAVs during the 2014–2015 outbreak period, unidentified exposure biases that could have resulted from sampling stranded animals, poor infectivity of such viruses in pinnipeds or other unidentified factors. Alternatively, as we did not test for antibodies to HP clade 2.3.4.4 IAVs, it is plausible that California pinnipeds that were sampled as part of this study had been previously exposed to such viruses and that our sampling for active viral shedding was not sufficiently broad in time or space to detect evidence of infection. The development and application of a serologic test to assess marine mammal exposure to HP clade 2.3.4.4 IAVs would be helpful for determining whether pinnipeds had been exposed to such viruses.

As per our efforts to assess the spatiotemporal extent of viral lineages and to refine inference regarding evolutionary pathways of novel reassortant viruses by using genetic analyses, the HP single clade 2.3.4.4 virus for which we were able to generate complete genomic data was found to be of the H5N8 subtype. Again, this finding is consistent with a lack of detection of reassortant HP clade 2.3.4.4 H5N1 or H5N2 IAVs in wild or domestic birds in California during the 2014–2015 outbreak. Thus, it is plausible that reassortant HP clade 2.3.4.4 H5N1 and H5N2 viruses emerged at areas north of California and were never introduced to this state. Alternatively, the introduction of these reassortant viruses may have occurred, but this study and other surveillance efforts in California during 2014–2015 lacked sufficient sensitivity to detect such viruses in wild or domestic birds.
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Figure 4 Partial Bayesian phylogenetic trees depicting the inferred genetic relationship among PB1, N1 NA and NS gene segment sequences for isolates that originated from wild waterfowl in California from 2014 to 2015 (indicated with asterisks), highly pathogenic (HP) clade 2.3.4.4 H5N1 influenza A viruses detected in North America during the concurrent outbreak and other reference sequences for wild bird low-pathogenic (LP) influenza A viruses isolated from elsewhere in North America as part of this study further support the hypothesis that viruses that circulated in wild birds in the Pacific Americas Flyway were involved in reassortment events that led to the emergence of novel reassortant HP IAV genome constellations. This finding, similar to information that was previously reported in research on IAVs in waterfowl that had been sampled in Alaska during 2014–2015,18,19 adds to the evidence that wild birds are likely to have played a role in the emergence and spread of novel reassortant HP clade 2.3.4.4 IAVs in North America.

On the basis of the results of our phylogenetic analyses, we propose a plausible evolutionary pathway for the HP clade 2.3.4.4 IAV strain A/American wigeon/California/UCDS8P/2015(H5N8), which was detected in Solano County, California, as part of this study (Figure 5). The phylogenetic position of sequences for six gene segments (PB2, HA, NP, N8, and NS and M) of A/American wigeon/California/UCDS8P/2015(H5N8) in Eurasian viral lineages and within monophyletic clades comprised other HP clade 2.3.4.4 IAVs that had been previously detected in wild and domestic birds in North America during the 2014–2015 outbreak, suggests that this virus most likely followed an evolutionary trajectory that involved reassortment between the HP strain A/Solano/CA/1625/2015(H5N8) in Eurasian viral lineages at the PA gene segment and a lack of detection of HP IAV genome constellations. This is supported through analyses in the current investigation, we acknowledge that numerous alternative pathways of viral evolution are also plausible.

Our phylogenetic analysis also provides support for relatively recent common ancestry at the PB1, PA, N1 NA and NS gene segments among LP IAVs that were recovered from wild and domestic birds during 2014–2015 outbreak in wild birds, independent of domestic poultry.

Phylogenetic clustering of the reassortant HP H5N8 virus that was sequenced as part of this study with other HP clade 2.3.4.4 IAVs that were detected in western North America during 2014–2015 is congruent with previously proposed pathways of viral introduction and evolution within the Pacific Americas Flyway.5,11,18,19,37 In addition, estimates of relatively recent common ancestry of numerous gene segments (i.e., PB1, PA, N1 and NS) of reassortant HP clade 2.3.4.4 H5N1 and H5N8 viruses with LP IAVs from waterfowl sampled in California as part of this study further support the hypothesis that viruses that circulated in wild birds in the Pacific Americas Flyway were involved in reassortment events that led to the emergence of novel reassortant HP IAV genome constellations. This finding, similar to information that was previously reported in research on IAVs in waterfowl that had been sampled in Alaska during 2014–2015,18,19 adds to the evidence that wild birds are likely to have played a role in the emergence and spread of novel reassortant HP clade 2.3.4.4 IAVs in North America.

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as no IAV isolate from California sequenced as part of this study contained more than a single gene segment that shared recent common ancestry with clade 2.3.4.4 HP H5N1 IAVs, which is comparable to data for LP IAVs from Alaska in which no more than two gene segments that shared recent common ancestry with a reassortant HP H5N1 IAV were identified in any given isolate,19 either multiple reassortment events led to the emergence of the HP clade 2.3.4.4 H5N1 genome constellation in North America or the genome constellation that served as a predecessor to this virus, via reassortment with the icA HP H5N8 initially introduced into North America, has not yet been identified. Thus, while this study provides important new data on genetic diversity for IAVs that were circulating among waterfowl in California during 2014–2015 and complements previously published data from Alaska, additional genetic information for wild waterfowl-origin LP IAVs from other locations in the Pacific Flyway during this time period would be useful for estimating the time and place of the emergence of this reassortant HP clade 2.3.4.4 H5N1 IAV genome constellation.

While we did not detect HP clade 2.3.4.4 H5 IAVs in wild waterfowl sampled in California in the summer of 2015 despite sampling IAV-naïve hatch-year birds at the same location where HP H5N8 had been identified 6 months prior, we caution against drawing the conclusion that such viruses have disappeared from this region or elsewhere in North America. Although it has been suggested that HP IAVs are adapted to domestic birds and unable to be maintained in wild waterfowl,18 recent detection of HP clade 2.3.4.4 H5N2 IAVs in wild mallards in Alaska19 and Montana,20 (which have seemingly been absent from domestic birds in North America since mid-2015), widespread detection of HP H5Nx IAVs in wild birds in Europe, Africa and Asia from 2016 to 2017,41 and pathobiological data from laboratory experiments that support the adaption of HP clade 2.3.4.4 H5 IAVs to waterfowl17,42,43 collectively provide evidence that this conceptual model may be inaccurate for numerous contemporary viral lineages descended from HP H5 Goose Guangdong lineage IAV predecessors.44 While information provided in this study and a previous report by Bevins et al.,16 provide data regarding the circulation of clade 2.3.4.4 IAVs in waterfowl in the Pacific Americas Flyway, we do not know the encounter and detection probabilities for such viruses in surveillance efforts, how prevalence varied across space and time during and after the 2014–2015 outbreak and how environmental persistence may contribute to viral ecology of clade 2.3.4.4 IAVs in North America. Thus, continued surveillance for HP IAVs in California and elsewhere in North America is warranted, as is vigilance with regard to adhering to biosecurity recommendations to limit the contact of domestic birds with water, soil or fomites that may have come into contact with wild waterfowl or wild bird feces.

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Figure 5  Proposed evolutionary pathway for influenza A virus strain A/American wigeon/California/UCD58P/2015(H5N8) sequenced as part of this study. Lines representing evolutionary mechanisms are labeled. Silhouettes depict that wild waterfowl were probable hosts for viruses along the proposed evolutionary pathway. The notation ‘mallard/California/H11Nx-like’ refers to an unsampled viral progenitor to H11Nx influenza A viruses that was detected in mallards sampled in California as part of this study sharing common ancestry at the PB1 and PA gene segments with reassortant HP H5N8 viruses. The subtype, host species and location for this progenitor virus are unknown. high ly pathogenic, HP; influenza A virus, IAV; intercontinental lineage A clade 2.3.4.4 H5N8, icA; low pathogenic, LP.
Supplementary Information for this article can be found on the Emerging Microbes & Infections website (http://www.nature.com/emi)