JVI Accepted Manuscript Posted Online 22 April 2020 J. Virol. doi:10.1128/JVI.00537-20 Copyright © 2020 Verhagen et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license.

- Phylogeography and antigenic diversity of low pathogenic avian influenza H13 and H16 1
- viruses 2

3

7

16

18

20

21

22

23

25

- Josanne H. Verhagen^{a,b}#, Marjolein Poen^a, David E. Stallknecht^c, Stefan van der Vliet^a, 4
- Pascal Lexmond^a, Srinand Sreevatsan^d, Rebecca L. Poulson^c, Ron A.M. Fouchier^a, Camille 5
- Lebarbenchon^{c,e} 6
- ^a Erasmus Medical Center, Department of Viroscience, Rotterdam, The Netherlands 8
- ^b Linnaeus University, Department of Biology and Environmental Science, Kalmar, Sweden 9
- ^c Southeastern Cooperative Wildlife Disease Study, College of Veterinary Medicine, 10
- Department of Population Health, University of Georgia, Athens, Georgia, USA 11
- ^d Michigan State University, College of Veterinary Medicine, Department of Pathobiology 12
- and Diagnostic Investigation, East Lansing, Michigan, USA 13
- 14 ^e Université de La Réunion, UMR Processus infectieux en milieu insulaire tropical (PIMIT),
- 15 Saint Denis, La Réunion, France
- Running title: Genetic and antigenic variation avian influenza virus 17
- #Address correspondence to Josanne H. Verhagen, josanne.verhagen@lnu.se 19

24

Word count: Abstract (239), Importance (151), Text (4500)

Abstract

27

28

29

30

31

32

33

34

35

36

37

38

39

40

41

42

43

44

26

Low pathogenic avian influenza viruses (LPAIVs) are genetically highly variable and have diversified into multiple evolutionary lineages that are primarily associated with wild bird reservoirs. Antigenic variation has been described for mammalian influenza viruses and for highly pathogenic avian influenza viruses that circulate in poultry, but much less is known about antigenic variation of LPAIVs. In this study, we focussed on H13 and H16 LPAIVs that circulate globally in gulls. We investigated the evolutionary history and intercontinental gene flow based on the hemagglutinin (HA) gene and used representative viruses from genetically distinct lineages to determine their antigenic properties by hemagglutination inhibition assays. For H13 at least three distinct genetic clades were evident, while for H16 at least two distinct genetic clades were evident. Twenty and ten events of intercontinental gene flow were identified for H13 and for H16 viruses, respectively. At least two antigenic variants of H13 and at least one antigenic variant of H16 were identified. Amino acid positions in the HA protein that may be involved in the antigenic variation were inferred, and some of the positions were located near the receptor binding site of the HA protein, as they are in the HA protein of mammalian influenza A viruses. These findings suggest independent circulation of H13 and H16 subtypes in gull populations as antigenic patterns do not overlap and contribute to the understanding of the genetic and antigenic variation of LPAIV naturally circulating in wild birds.

46

47

45

Importance

- Wild birds play a major role in the epidemiology of low pathogenic avian influenza viruses 48
- (LPAIVs) from which these viruses are occasionally transmitted—directly or indirectly—to 49
- 50 other species, including domestic animals, wild mammals and humans, where they can cause

52 LPAIVs in wild birds is poorly understood. Here, we investigated the evolutionary history, intercontinental gene flow, and the antigenic variation among H13 and H16 LPAIVs. The 53 54 circulation of the subtypes H13 and H16 seems to be maintained by a narrower host range, in particular gulls, than for the majority of LPAIV subtypes and may therefore serve as a model 55 for evolution and epidemiology of H1-H12 LPAIVs in wild birds. The findings suggest that 56 57 H13 and H16 LPAIVs circulate independently of each other and emphasize the need to investigate within clade antigenic variation of LPAIVs in wild birds. 58 59 60 **Keywords:** avian viruses, influenza, evolution, epidemiology, ecology, antigenic variation, seabird 61

subclinical to fatal disease. Despite a multitude of genetic studies, the antigenic variation of

Introduction

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

62

Wild birds of the orders Anseriformes (mainly ducks, geese and swans) and Charadriiformes (mainly gulls, terns and waders) play a major role in the epidemiology of low pathogenic avian influenza viruses (LPAIVs). LPAIVs are characterized into subtypes based on their surface proteins hemagglutinin (HA, H1-H16) and neuraminidase (NA, N1-N9), e.g. H13N6. Ducks play an important role in the epidemiology of most LPAIV subtypes. However, birds of the order Charadriiformes—in particular gulls— are the major reservoir for subtypes H13 and H16 (Table S1) (1-4). High prevalence of H13 and/or H16 LPAIVs has been observed in juvenile gulls at breeding colony sites (5-7) and in adults during spring and/or fall migration (8, 9). H13 and H16 viruses have a global distribution. Since first detection in 1977, H13 viruses have been detected in North America, South America, Europe, Asia, Africa and Oceania. Since their first detection in 1975, H16 viruses have been detected in North America, South America, Europe and Asia. The spatial isolation of host populations has shaped LPAIV evolution and led to the independent circulation of different virus gene pools between Western and Eastern hemispheres (10). Yet, some pelagic gull populations connect multiple continents through seasonal migration and overlapping distributions and could facilitate rapid and long-distance dispersal of LPAIV genomes (2, 9, 11-14). For instance, great black-backed gulls (Larus marinus) migrate between Europe and the east coast of North America, and LPAIVs consisting of both North American as well as Eurasian genes have been isolated from this species (12). Upon intercontinental gene flow, i.e. the movement of genes between the different continents, some LPAIV genes seem to have become established in the population, e.g. H6 (15). Influenza A viruses (IAV) belong to the family Orthomyxoviridae and are negative sense single-stranded RNA viruses with a segmented genome. The genome consists of eight

88

89

90

91

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

segments encoding 12 proteins or more, including the surface proteins HA and NA. The HA protein of IAV is a major determinant for virus binding to cells and subsequent cell entry and for generation of IAV-specific antibodies, and thus subjected to strong selective pressure (16). Indeed, in wild birds—in particular mallards (Anas platyrhynchos)—LPAIV infection dynamics seem to be shaped between LPAIV subtypes partially by pre-existing homo- or heterologous antibodies (17). Furthermore, within other host systems, evasion of IAV-specific antibodies by IAVs—so called antigenic variation—has been described for seasonal human IAVs (18, 19), swine IAVs (20-22), equine IAVs (23) and for highly pathogenic avian influenza viruses (HPAIVs) that circulate in poultry (24, 25). Despite numerous studies on the genetic variation of LPAIVs in wild birds, the antigenic variation within LPAIV subtypes that circulate in wild birds is barely investigated (26, 27). To better understand LPAIV epidemiology in gulls, we investigated the global distribution of H13 and H16 LPAIVs and the antigenic variation of a representative subset of H13 and H16 LPAIVs. Based on the sequencing of HA genes of 84 viruses, and hemagglutination inhibition assays, we showed that intercontinental H13 and H16 gene flow occurred frequently, and that H16 genetic lineages did not form antigenic clusters, suggesting that clade-defining mutations were not in critical epitopes (i.e. part of the antigen that binds to specific antibodies). In contrast, the H13 genetic clades partially corresponded with the antigenic variation of H13 LPAIVs, suggesting part of the clade-defining mutations were in critical epitopes. **Results**

Phylogeographic structure and intercontinental gene flow

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134

135

136

Phylogenetic analyses supported that the H13 HA was structured in three major genetic lineages (A-C; Figure 1, S1 and S2). The time to the most recent common ancestor (tMRCA) of the H13 HA gene was dated in 1927 (± 95% HPD (highest posterior density): [1920-1934]). The tMRCA of viruses of clade A (1963 [1958-1966]) was older than the ones of clade B (1975 [1974-1976]) and C (1977 [1976-1978]). Our analyses support that the geographic origin of H13 viruses of clade B and C could be North America and Europe, respectively (posterior probabilities for the geographic origin of the most recent common ancestor [MRCA]: 1 for clade B and 1 for clade C). For clade A, limited historical data of viruses from different locations as well as low posterior probability (0.62) precludes a conclusion on the geographic origin of the MRCA. Since the first isolation of an H13 IAV from a gull in 1977, 20 potential events of intercontinental gene flow were identified (indicated with 1-20 in Figure 1, S3 and Table 2). Clade A supports the maintenance of H13 in European gulls, with evidence of multiple introductions to North America and Asia (events #3, #5, #6, #7, and #10), and a reverse introduction from North America to Asia (event #8). Clade C was also composed mainly of viruses circulating in Europe, with evidence of multiple introductions to North America (events #12, #15, #19) and Asia (events #13, #16, #17). The introduction of clade C H13 HA in North America (event #19) was followed by an introduction to South America (event #20). Evidence for intercontinental gene flow among North American H13 IAV occurred among eastern and western North American isolates (event #3, #12, #15 and #19). Clade B was composed almost exclusively of viruses circulating in North America, although one gene flow event to South America occurred recently (event #11). The H16 HA was structured in at least two major genetic lineages (Figure 2, S4 and S5). The MCC tree was structured in three main clades (A-C, Figure S5), while the ML tree

provided support for only two main genetic clades (A and B/C merged, Figure S4). The

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

160

161

tMRCA of the H16 HA gene was dated in 1924 [1914-1932]. Clade A included only viruses from Europe and was dated in 1977 [1975-1980]; clade B included only viruses from North America with a time to the tMRCA estimated in 1969 [1967-1971]. Our analyses supported that the geographic origin of clade A and B was Europe and North America, respectively (posterior probabilities for the geographic origin of the MRCA: 0.99 for clade A, 1 for clade B). The tMRCA of clade C was estimated 1965 [1962-1968]. Clade C may have arisen in Europe (posterior probabilities for the geographic origin of the MRCA: 0.87) and consisted of viruses of mixed origin, i.e. Europe, Asia and North America. Since the first isolation of an H16 IAV from a black-legged kittiwake (Rissa tridactyla) in 1975, ten intercontinental gene flow events were identified for viruses of clade C (indicated with 1-10 in Figure 2, S6 and Table 3). As for the H13 subtype, strong support for gene flow between Europe and North America was found, in particular from North-Western European countries: Denmark to North-eastern America (Delaware, New Hampshire, Ouebec), and Iceland to Newfoundland (events #6 and #10). Evidence for intercontinental gene flow among North American H16 IAV occurred among eastern and western North American isolates (event #3, #6, #8 and #10). In particular, intercontinental gene flow #8 seems to have been maintained in North America after its initial introduction in 2006 [2005-2006], for at least ten years, and may have replaced clade B of H16 HA (Figure 2). High rates of nucleotide substitution obtained for the H13 HA genetic lineages were consistent with those previously reported for H4, H6 and H7 subtypes circulating in wild ducks (Table 4). However, the nucleotide substitution rate of clade B—that consists exclusively of North American IAV—was lower than mean rates and HPD obtained for the other two H13 clades. The mean d_N/d_S rate obtained for the three H13 genetic clades were comparable to those previously reported for other subtypes and suggests that the HA was

under strong purifying selection (Table 4). Nonetheless, a slightly higher d_N/d_S rate obtained

Downloaded from http://jvi.asm.org/ on October 7, 2020 by guest

for clade B and C as compared to other lineages suggests that they may be subjected to a more neutral selection. The mean nucleotide substitution and d_N/d_S rates for the H16 gene were also consistent with H13 HA as well as with H4, H6 and H7 subtypes from wild ducks. However, H16 clade C (European mixed)- that consisted of viruses of a geographically more mixed origin – had slightly lower nucleotide substitution rates and higher d_N/d_S rates than clade A (European) and clade B (North American) (Table 4).

168

162

163

164

165

166

167

Antigenic diversity between H13 and H16 LPAIV

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

185

169

As expected from two different HA subtypes, the H13 and H16 viruses formed two separate antigenic variants. The H13 and H16 viruses were generally well separated, forming groups on opposite sides of the antigenic map (Figure 3, Table 5). A total of nine amino acid positions within/near the receptor binding site of the HA were identified that differed consistently between H13 and H16 viruses (based on alignments of 338 H13 and 192 H16 HA indicated in Table 6), of those, amino acid position 145 was located in the 130-loop, 200 and 208 in the 190-helix and 231 and 233 in the 220-loop of the receptor binding site of the HA (HA numbering based on (28, 29). Of those, amino acid position 233 was listed previously as being involved in differences in receptor-binding site between HAs originating from Laridae and Anatidae (30). Additionally, the amino acid at position 196 differed between H13 (valine [V]) and H16 (aspartic acid [D]) viruses; this position may contribute to receptor binding specificity as identified previously based on crystal structures of H5 and H13 LPAIV (31). Due to non-specific cross-reactivity, two H13 viruses (i.e. HEGU/AK/458/85 and HEGU/AK/479/85) had unexpected high titers against H16 antisera (Table 5) and were therefore positioned in the center of the map and served to pull H13 and H16 together.

186

Downloaded from http://jvi.asm.org/ on October 7, 2020 by guest

Antigenic diversity among H13 LPAIV

188

189

190

191

192

193

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

211

187

The representative H13 viruses formed at least two different antigenic variants (Figure 3, Table 5). The viruses of H13 clades A and B were genetically distinct (Figure 1) but were antigenically similar (Figure 3), based on the H13 clade A antisera cross-reacting with H13 clade B viruses and vice versa. In contrast, H13 clade C viruses reacted poorly—if at all with antisera that were raised against clade A and B viruses, and, conversely, antisera against clade C viruses rarely reacted with substantial titers with viruses of clade A and B. Thus, H13 clade A/B and H13 clade C viruses formed two different antigenic variants. The antigenic diversity of H13 clade A/B combined is about the same as the antigenic diversity of the H13 clade C. One H13 clade B virus, i.e. LAGU/DB/1370/86, could not be placed well in the map due to HI titers of 40 or lower (Table 5).

To gain insight into the molecular basis of the antigenic variation between H13 clade A/B and C, amino acids that differed consistently among the different clades of H13 viruses were indicated (based on the alignment of 338 H13, Table 6). A total of four amino acid positions within/near the receptor binding site of the HA were identified that differed consistently for clade A, B and/or C. Of those, amino acids at positions 149 and 254 differed consistently between clade A/B and C. Viruses belonging to clade C—except a single virus from South America that had a arginine (R) at position 149—had a deletion at position 149 (previously identified using a smaller dataset as position 154 (12)), in contrast to viruses of clade A or B that had an aspartic acid (D), glutamic acid (E), asparagine (N) or serine (S) at this position. The correlation between the antigenic distance of H13 representative viruses from A/gull/MD/704/1977 (H13N6) (clade A)—the first detected H13 virus—and the number of HA1 amino acid substitutions from A/gull/MD/704/1977 was 0.87 and was statistically significant (P < 0.0001, Pearson correlation).

Antigenic diversity among H16 LPAIV

214

215

216

217

218

219

220

221

222

223

224

225

226

227

228

229

230

212

213

The representative H16 viruses formed at least one antigenic variant (Figure 3 and Table 5). The genetically distinct H16 clades A, B and C did not form separate antigenic clusters in the map, which reflects the raw HI data as there are no patterns for any of the four H16 antisera tested that correspond to the genetic lineages. The antigenic diversity of the H16 viruses is within eight antigenic units, with BHGU/NL/1/07 being on the edge of this antigenic space (i.e. low titers to all sera). The antigenic diversity of H16 clade A/B/C is about the same as the antigenic diversity of the H13 clade A/B combined and similar to the antigenic diversity of the H13 clade C. Though clade A, B and C did not form separate antigenic clusters in our analysis, amino acids that differed consistently among the different clades of H16 viruses were indicated (based on the alignment of 192 H16 HA, Table 6). A total of three amino acid positions within/near the receptor binding site of the HA were identified that differed consistently among the three H16 clades and were not associated with antigenic variation. The correlation between the antigenic distance of the representative viruses from A/Black-headed gull/TM/13/76 (H16N3) (clade C)—one of the first detected H16 viruses—and the number of HA1 amino acid substitutions from A/Black-headed gull/TM/13/76 was 0.67 and was statistically significant (P = 0.003, Pearson correlation).

232

233

231

Discussion

234

235

236

We investigated the evolutionary history and intercontinental gene flow based on the hemagglutinin (HA) gene of H13 and H16 LPAIV and selected representative viruses from

238

239

240

241

242

243

244

245

246

247

248

249

250

251

252

253

254

255

256

257

258

259

260

261

genetically distinct lineages to determine their antigenic properties by HI assays. H13 formed at least three distinct genetic clades as suggested previously based on smaller datasets (9, 32-35), while H16 formed at least two distinct genetic clades. Twenty and ten events of intercontinental gene flow were identified for H13 and for H16 viruses, respectively. At least two antigenic variants of H13 and at least one antigenic variant of H16 were identified. The presence of different antigenic variants among viruses of a single LPAIV subtype is in contrast to previous findings based on antigenic characterization of LPAIV H3 (26), and implies that antigenic variation within LPAIV subtypes occurs.

The frequency of intercontinental gene flow of the HA gene of H13 and H16 viruses was similar to the HA gene of H6 viruses, but lower than for internal genes (2, 27, 36, 37). Previously, intercontinental gene flow has been described extensively for the H6 HA genes, while no intercontinental gene flow was detected for the H4 and H7 subtypes (15, 38). For the H6 subtype, gene flow has been described ten times with four established genes during a period of 31 years (1975-2006; (15)). Also, evidence for intercontinental gene flow among North American H13 and H16 genes occurred among eastern and western North American LPAIVs in contrast to eastern North American LPAIVs only as reported previously (39). Given the relatively high number of intercontinental flow of IAV internal genes by shorebirds and gulls (2, 27, 36, 37), one may have expected a higher gene flow of gull-associated H13 and H16 HA genes, compared to e.g. H6. However, a higher intercontinental gene flow only was apparent with H13 (i.e. 20 events during a period of 35 years). This may suggest i) broader host range, host population size and/or host distribution of H13 than H16, and/or ii) local H13-specific herd-immunity is lower than H16-specific herd immunity and therefore less limiting establishment opportunities in host populations of H13, and/or iii) higher environmental survival of H13 than of H16, and/or iv) introduced H13 HA genes may be less

affected by strong subtype-dependant competition with endemic HA genes (e.g. with respect

263

264

265

266

267

268

269

270

271

272

273

274

275

276

277

278

279

280

281

282

283

284

285

286

to linkage to NS1 and NP as these contain most gull-specific features (33)) than introduced H16 genes. Interestingly, no H13 or H16 gene flow was described from Asia to Europe, which is in contrast to e.g. HPAIV H5 viruses that have been introduced from Asia to Europe several times (40, 41). The relatively low frequency of detection of intercontinental gene flow of H13 or H16 genes out of North America and in particular Asia, relative to Europe, may be due to a bias in IAV surveillance and sequencing (i.e. number of available IAV sequences from gulls isolated in Europe is higher than from North America and in particular Asia). Antigenic diversity of LPAIV depends partially on the host population size and

structure. In this study, both H13 and H16 LPAIV formed at least three or two distinct genetic clades respectively that did not or only partially corresponded with antigenic clusters. The H16 genetic clades did not form antigenic clusters, suggesting that clade-defining mutations were not in critical epitopes. In contrast, the H13 genetic clades partially corresponded with the antigenic variation of H13 LPAIV, suggesting that part of the clade-defining mutations were in critical epitopes. Also, given that the H13 antigenic space is larger than the antigenic space covered by H16 viruses, the host population of H13 may be larger and more widely distributed than the host population of H16 LPAIV, facilitating the circulation of more than one antigenic variant of a single LPAIV subtype. Strong genetic and antigenic divergence between two co-circulating lineages could be the product of a very large host meta-population size and relatively rare cross-species transmission rate (42). Globally, viruses of the H13 subtype seem to be more common than viruses of the H16 subtype (2, 4), which is consistent with the finding that H13 LPAIV consists of multiple antigenic variants. Besides increased host population size and host distribution, prolonged virus survival may shape LPAIV epidemiology and evolution. Antigenic diversity within H13 LPAIV may be shaped by amino acid substitutions near the receptor binding site of the HA protein. In this study, we found evidence that amino acids or deletions at positions 149 and 254 of the HA protein may be

288

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

involved in antigenic diversity among H13 strains. In addition, position 149 could be involved in H16 LPAIV antigenic diversity as all H16 viruses had a deletion at this position and H16 clade A, B and C were antigenically similar.

Co-circulating and newly introduced H13 or H16 LPAIV can be either antigenically similar or antigenically different. In the Northern hemisphere, H13 and H16 IAV subtypes circulate most extensively on breeding colonies in hatch-year birds at the end of summer and early fall (5-7). In black-headed gulls (which in Europe are one of the main host for H13 and H16 LPAIV), infection with H13 or H16 result in strong protection against reinfection with the same virus, however susceptibility to infection with the other subtype or with another strain of the same subtype is unknown (43, 44). Our findings support the independent longterm maintenance and co-circulation of at least two genetically distinct lineages of H13 and of H16 in Eurasia. This pattern is similar to the one that has been described for the H3 IAV subtype in ducks in North America (42). Our analysis showed that these genetically distinct co-circulating lineages may belong to the same antigenic variant. Here, we found evidence that genetically distinct co-circulating H13 or H16 LPAIV on a black-headed gull breeding colony site in the Netherlands may be either antigenically different (e.g. H13 clade A virus A/BHGU/NL/7/2009 (H13N2) and H13 clade C virus A/BHGU/NL/20/2009 (H13N2) or antigenically similar (e.g. H16 clade A A/BHGU/NL/10/2009 (H16N3) and A/BHGU/NL/21/2009 (H16N3) and H16 clade C A/BHGU/NL/26/2009 (H16N3). Similar, intercontinental gene flow occurred with HA genes that were antigenically similar to local circulating viruses (i.e. H16 clade C viruses that were genetically closely related to SB/DB/172/06 and SB/DB/195/06 versus local circulating H16 clade B viruses), and HA genes that were antigenically different from local circulating viruses (i.e. H13 clade C viruses, genetically closely related to LAGU/NJ/AI08-0714/08 versus local circulating H13 clade B viruses.

313

314

315

316

317

318

319

320

321

322

323

324

326

327

328

329

330

331

332

333

334

335

336

Antigenic variation within a LPAIV subtype at the clade level (i.e. H13 clade A/B combined versus H13 clade C) was described here, yet less is known about antigenic variation within genetic clades of H13, H16 or other LPAIV subtypes. For H13, genetic diversity within clades seemed stable—e.g. viruses of clade A, B or C, collected over three decades were antigenically closely related—suggesting no major genetic differences; this is in contrast to the few mutations needed for antigenic change in seasonal human IAV. Similarly, a study on antigenic variation of H3 LPAIV isolated in North America suggested that genetically diverse viruses were antigenically stable (26). Major antigenic changes in seasonal human IAV were due to amino acid substitutions immediately adjacent to the receptor binding site (18); this could potentially also explain antigenic variation between antigenically different viruses of H13 clade A/B combined and clade C (i.e. amino acid positions 149 of the HA). Future work on antigenic variation of LPAIV should include within clade genetic and antigenic variation.

325

Materials and Methods

Viruses. The HA sequences of H13 (n=64) and H16 (n=20) viruses isolated from wild birds in North America (n=39 and n=5, respectively) and Europe (n=25 and n=15, respectively) between 1976 and 2010 were determined at the University of Minnesota (Saint Paul, Minnesota, USA) and at the Department of Viroscience of the Erasmus Medical Center (Rotterdam, the Netherlands). Details on virus isolates including GenBank accession numbers are summarized in Table S2 and S3; details related to the Sanger sequencing methodology are available upon request. The HA sequences were supplemented with full-length nucleotide sequences of the HA gene of H13 and H16 viruses isolated from wild birds between 1975 and 2017 and downloaded from GenBank (https://www.ncbi.nlm.nih.gov). The full dataset

338

339

340

341

342

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

359

360

361

included sequences of H13 (n=519) and H16 (n=276) HA genes and was biased towards virus strains collected since 2000 due to increased surveillance and sequencing since 2000. Of this full dataset, viruses representing the genetically distinct clades were selected (n=44; H13 clade A, B, C and H16 clade A, B, C; see the Results section for clade definition) to investigate the antigenic diversity of H13 and H16 viruses. Of those viruses, viruses that were genetically most divergent were selected (n=10) to generate ferret antisera (Table 1). The antigenic properties of all representative viruses (n=44) were analysed in hemagglutination inhibition (HI) assays using the panel of ten ferret antisera. Genetic analyses. The nucleotide sequences of the coding region of H13 and H16 HA were

aligned with the program CLC 8.0 (CLC bio, Aarhus, Denmark). Neighbor-Joining trees were then generated, with 1000 bootstraps, in order to assess the overall genetic structure of the H13 (n=519) and H16 (n=276) HA sequences. To lower the bias in species and geography (e.g. black-headed gulls (Chroicocephalus ridibundus) from the Netherlands and glaucouswinged gulls (Larus glaucescens) from Alaska), duplicate sequences (i.e. identical sequences of the same host species, location and date) were identified with Mothur 1.39.5 (45) and removed, resulting in final alignments of H13 (n=338) and H16 (n=192) HA. To identify the genetic structure of H13 and H16 virus subtypes Maximum-likelihood trees with 1000 bootstraps were generated with the software PhyML 3.1 (46). The general time reversible (GTR) evolutionary model, an estimation of the proportion of invariable sites (I) and of the nucleotide heterogeneity of substitution rate (a) was used as selected by Model Generator 0.85 (47). To investigate the evolutionary history of H13 and H16 virus subtypes Bayesian Markov Chain Monte Carlo coalescent analyses were performed. The temporal structure of the dataset was assessed with the program TempEst 1.5.3 (48). Both datasets showed a positive correlation between genetic divergence and sampling time and appear to be suitable

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

384

385

386

Downloaded from http://jvi.asm.org/ on October 7, 2020 by guest

for phylogenetic molecular clock analyses. Time to the most recent common ancestors (MRCA) as well as geographic ancestral states (i.e. continent), and their associated posterior probabilities were obtained based on the method described by Lemey et al. with the program BEAST 1.10.1 (49, 50). A strict molecular clock model was selected as relaxed clock models (uncorrelated exponential and uncorrelated lognormal) resulted in low effective sample sizes (ESS < 200) in spite of high chain length (>200 million states). In all simulations a Bayesian skyline coalescent tree prior (51) was selected. The Shapiro-Rambaut-Drummond-2006 nucleotide substitution model was selected (52), and has been used in population dynamic studies of other IAV subtypes (15, 38, 42, 53). Overall, a similar methodology was used as in previous studies on IAV evolutionary dynamics of subtypes H4, H6 and H7 (15, 38, 54). Analyses were performed with two independent chain lengths of 100 million generations sampled every 1000 iterations; the first 10% of trees were discarded as burn-in. Substitutions rates based on independent analyses of the major H13 and H16 clades were obtained using the program BEAST 1.10.1. Nonsynonymous substitutions (d_N) and synonymous substitutions (d_S) rates were obtained using the single likelihood ancestor counting method implemented in HyPhy (55). Computations were performed with the Datamonkey webserver (56, 57). **Antisera.** Post-infection antisera were prepared upon nasal inoculation of ferrets (> 1 year of age, male, two ferrets per virus) with virus (cultured on embryonated chicken eggs, per ferret 10^6 - 10^7 median egg infectious dose (EID₅₀)/100 µl) and blood collection by exsanguination 14 days later. An overview of antisera used in this study is provided in Table 1. Antisera were pre-treated overnight at 37°C with receptor-destroying enzyme (Vibrio cholerae neuraminidase), followed by inactivation for 1 hr at 56°C before use in HI assays.

Antigenic analyses. HI assays were performed according to standard procedures (58). The HI

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

407

408

409

410

411

titer is expressed as the reciprocal value of the highest serum dilution that completely inhibited hemagglutination. To investigate antigenic variation among and within H13 and H16 viruses, antigenic cartography methods were used as described previously (19). Briefly, antigenic cartography is a method to analyse and visualize HI assay data. The titers in an HI table can be thought of as specifying target distances between antigens and antisera. In an antigenic map, the distance between antigen point A and antiserum point S corresponds to the difference between the log₂ value of the maximum observed titer to antiserum S from any antigen and the titer of antigen A to antiserum S. Modified multidimensional scaling methods are used to arrange the antiserum and antigen points in an antigenic map to best satisfy the target distances specified by the HI data (18). Because antigens are tested against multiple antisera, and antisera are tested against multiple antigens, many measurements can be used to determine the position of the antigens and antisera in an antigenic map, thus improving the resolution of the HI data. **Ethics statement.** This study was approved by the independent animal experimentation ethical review committee Stichting DEC consult (Erasmus MC permit 122-98-01, 122-08-04 and 15-340-03) and was performed under animal biosafety level 2 (ABSL-2) conditions. Animal welfare was monitored daily, and all animal handling was performed under light anesthesia (ketamine) to minimize animal discomfort. Data availability. Sequences are available in GenBank under accession numbers KF612922 to KF612965, KR087564, KR087572, KR087577 to KR087595, KR087597 to KR087601, KR087604 to KR087615, and MK027211 and MK027212.

Acknowledgements

1	1	3
7	1	J

424

412

414	This work was funded by the Swedish Research Council Vetenskapsrådet [2015-03877],
415	National Institute of Allergy and Infectious Diseases (NIAID), National Institutes of Health
416	(NIH), Department of Health and Human Services, under Contract No.
417	HHSN266200700007C and HHSN272201400008C. CL is supported by a 'Chaire mixte :
418	Université de La Réunion – INSERM'. The funding agencies did not have any involvement in
419	the study design, implementation, or publishing of this study and the research presented
420	herein represents the opinions of the authors, but not necessarily the opinions of the funding
421	agencies. We gratefully acknowledge the following researchers for sharing, preparing virus
422	isolates and sequences amongst others: Scott Krauss, Janice C. Pedersen, Shinichiro
423	Enomoto, Justin D. Brown, Jonathan Runstadler, Nichola Hill, Nicola Lewis, Alexander

Shestopalov, Neus Latorre-Margalef, and Jonas Waldenström.

References

425

- Olsen B, Munster VJ, Wallensten A, Waldenstrom J, Osterhaus AD, Fouchier RA. 426 1.
- 2006. Global patterns of influenza a virus in wild birds. Science 312:384-8. 427
- 2. Wille M, Robertson GJ, Whitney H, Bishop MA, Runstadler JA, Lang AS. 2011. 428
- Extensive geographic mosaicism in avian influenza viruses from gulls in the northern 429
- hemisphere. PLoS One 6:e20664. 430
- 431 3. Lang AS, Lebarbenchon C, Ramey AM, Robertson GJ, Waldenstrom J, Wille M.
- 2016. Assessing the Role of Seabirds in the Ecology of Influenza A Viruses. Avian 432
- 433 Dis 60:378-86.
- 434 Zhang Y, Aevermann BD, Anderson TK, Burke DF, Dauphin G, Gu Z, He S, Kumar
- S, Larsen CN, Lee AJ, Li X, Macken C, Mahaffey C, Pickett BE, Reardon B, Smith T, 435
- Stewart L, Suloway C, Sun G, Tong L, Vincent AL, Walters B, Zaremba S, Zhao H, 436
- 437 Zhou L, Zmasek C, Klem EB, Scheuermann RH. 2017. Influenza Research Database:
- An integrated bioinformatics resource for influenza virus research. Nucleic Acids Res 438
- 439 45:D466-D474.
- 440 5. Verhagen JH, Majoor F, Lexmond P, Vuong O, Kasemir G, Lutterop D, Osterhaus
- AD, Fouchier RA, Kuiken T. 2014. Epidemiology of influenza A virus among black-441
- 442 headed gulls, the Netherlands, 2006-2010. Emerging Infectious Diseases 20:138-41.
- 443 6. Velarde R, Calvin SE, Ojkic D, Barker IK, Nagy E. 2010. Avian influenza virus H13
- circulating in ring-billed gulls (Larus delawarensis) in southern Ontario, Canada. 444
- 445 Avian Dis 54:411-9.
- 446 7. Graves IL. 1992. Influenza viruses in birds of the Atlantic flyway. Avian Diseases
- 36:1-10. 447

14.

470

471

472

448 8. Lewis NS, Javakhishvili Z, Russell CA, Machablishvili A, Lexmond P, Verhagen JH, Vuong O, Onashvili T, Donduashvili M, Smith DJ, Fouchier RA. 2013. Avian 449 influenza virus surveillance in wild birds in Georgia: 2009-2011. PLoS One 8:e58534. 450 451 9. Huang Y, Wille M, Benkaroun J, Munro H, Bond AL, Fifield DA, Robertson GJ, Ojkic D, Whitney H, Lang AS. 2014. Perpetuation and reassortment of gull influenza 452 A viruses in Atlantic North America. Virology 456-457:353-63. 453 454 10. Obenauer JC, Denson J, Mehta PK, Su X, Mukatira S, Finkelstein DB, Xu X, Wang J, Ma J, Fan Y, Rakestraw KM, Webster RG, Hoffmann E, Krauss S, Zheng J, Zhang Z, 455 Naeve CW. 2006. Large-scale sequence analysis of avian influenza isolates. Science 456 457 311:1576-80. 11. Van Borm S, Rosseel T, Vangeluwe D, Vandenbussche F, van den Berg T, Lambrecht 458 B. 2012. Phylogeographic analysis of avian influenza viruses isolated from 459 460 Charadriiformes in Belgium confirms intercontinental reassortment in gulls. Arch Virol 157:1509-22. 461 Wille M, Robertson GJ, Whitney H, Ojkic D, Lang AS. 2011. Reassortment of 462 12. 463 American and Eurasian genes in an influenza A virus isolated from a great blackbacked gull (Larus marinus), a species demonstrated to move between these regions. 464 465 Archives of Virology 156:107-15. 13. Ratanakorn P, Wiratsudakul A, Wiriyarat W, Eiamampai K, Farmer AH, Webster RG, 466 Chaichoune K, Suwanpakdee S, Pothieng D, Puthavathana P. 2012. Satellite Tracking 467 468 on the Flyways of Brown-Headed Gulls and Their Potential Role in the Spread of 469 Highly Pathogenic Avian Influenza H5N1 Virus. PLoS One 7:e49939.

Hall JS, Teslaa JL, Nashold SW, Halpin RA, Stockwell T, Wentworth DE, Dugan V,

Ip HS. 2013. Evolution of a reassortant North American gull influenza virus lineage:

drift, shift and stability. Virol J 10:179.

496

473 15. Bahl J, Vijaykrishna D, Holmes EC, Smith GJ, Guan Y. 2009. Gene flow and competitive exclusion of avian influenza A virus in natural reservoir hosts. Virology 474 390:289-97. 475 476 16. Webster RG, Bean WJ, Gorman OT, Chambers TM, Kawaoka Y. 1992. Evolution and ecology of influenza A viruses. Microbiol Rev 56:152-79. 477 Latorre-Margalef N, Grosbois V, Wahlgren J, Munster VJ, Tolf C, Fouchier RA, 478 17. 479 Osterhaus AD, Olsen B, Waldenstrom J. 2013. Heterosubtypic immunity to influenza A virus infections in mallards may explain existence of multiple virus subtypes. PLoS 480 Pathogens 9:e1003443. 481 482 18. Koel BF, Burke DF, Bestebroer TM, van der Vliet S, Zondag GC, Vervaet G, Skepner E, Lewis NS, Spronken MI, Russell CA, Eropkin MY, Hurt AC, Barr IG, de Jong JC, 483 Rimmelzwaan GF, Osterhaus AD, Fouchier RA, Smith DJ. 2013. Substitutions near 484 485 the receptor binding site determine major antigenic change during influenza virus evolution. Science 342:976-9. 486 487 19. Smith DJ, Lapedes AS, de Jong JC, Bestebroer TM, Rimmelzwaan GF, Osterhaus 488 AD, Fouchier RA. 2004. Mapping the antigenic and genetic evolution of influenza virus. Science 305:371-6. 489 490 20. Lorusso A, Vincent AL, Harland ML, Alt D, Bayles DO, Swenson SL, Gramer MR, 491 Russell CA, Smith DJ, Lager KM, Lewis NS. 2011. Genetic and antigenic characterization of H1 influenza viruses from United States swine from 2008. Journal 492 493 of General Virology 92:919-30. 494 21. Marozin S, Gregory V, Cameron K, Bennett M, Valette M, Aymard M, Foni E,

Barigazzi G, Lin Y, Hay A. 2002. Antigenic and genetic diversity among swine

influenza A H1N1 and H1N2 viruses in Europe. J Gen Virol 83:735-45.

518

519

520

27.

Pathogens 3:e167.

497 22. Cong YL, Pu J, Liu QF, Wang S, Zhang GZ, Zhang XL, Fan WX, Brown EG, Liu JH. 2007. Antigenic and genetic characterization of H9N2 swine influenza viruses in 498 China. J Gen Virol 88:2035-41. 499 500 23. Lewis NS, Daly JM, Russell CA, Horton DL, Skepner E, Bryant NA, Burke DF, Rash AS, Wood JL, Chambers TM, Fouchier RA, Mumford JA, Elton DM, Smith DJ. 2011. 501 Antigenic and genetic evolution of equine influenza A (H3N8) virus from 1968 to 502 503 2007. J Virol 85:12742-9. 24. Cattoli G, Milani A, Temperton N, Zecchin B, Buratin A, Molesti E, Aly MM, Arafa 504 A, Capua I. 2011. Antigenic drift in H5N1 avian influenza virus in poultry is driven by 505 506 mutations in major antigenic sites of the hemagglutinin molecule analogous to those for human influenza virus. J Virol 85:8718-24. 507 25. Koel BF, van der Vliet S, Burke DF, Bestebroer TM, Bharoto EE, Yasa IW, Herliana 508 509 I, Laksono BM, Xu K, Skepner E, Russell CA, Rimmelzwaan GF, Perez DR, Osterhaus AD, Smith DJ, Prajitno TY, Fouchier RA. 2014. Antigenic variation of 510 511 clade 2.1 H5N1 virus is determined by a few amino acid substitutions immediately 512 adjacent to the receptor binding site. MBio 5:e01070-14. 26. Bailey E, Long LP, Zhao N, Hall JS, Baroch JA, Nolting J, Senter L, Cunningham FL, 513 514 Pharr GT, Hanson L, Slemons R, DeLiberto TJ, Wan XF. 2016. Antigenic 515 Characterization of H3 Subtypes of Avian Influenza A Viruses from North America. Avian Dis 60:346-53. 516

Krauss S, Obert CA, Franks J, Walker D, Jones K, Seiler P, Niles L, Pryor SP,

migratory birds and evidence of limited intercontinental virus exchange. PLoS

Obenauer JC, Naeve CW, Widjaja L, Webby RJ, Webster RG. 2007. Influenza in

22

544

545

36.

521 28. de Graaf M, Fouchier RA. 2014. Role of receptor binding specificity in influenza A virus transmission and pathogenesis. EMBO J 33:823-41. 522 29. Burke DF, Smith DJ. 2014. A recommended numbering scheme for influenza A HA 523 subtypes. PLoS One 9:e112302. 524 30. Yamnikova SS, Gambaryan AS, Tuzikov AB, Bovin NV, Matrosovich MN, 525 Fedyakina IT, Grinev AA, Blinov VM, Lvov DK, Suarez DL, Swayne DE. 2003. 526 527 Differences between HA receptor-binding sites of avian influenza viruses isolated from Laridae and Anatidae. Avian Diseases 47:1164-8. 528 31. Lu X, Qi J, Shi Y, Wang M, Smith DF, Heimburg-Molinaro J, Zhang Y, Paulson JC, 529 530 Xiao H, Gao GF. 2013. Structure and receptor binding specificity of hemagglutinin H13 from avian influenza A virus H13N6. J Virol 87:9077-85. 531 32. Benkaroun J, Shoham D, Kroyer ANK, Whitney H, Lang AS. 2016. Analysis of 532 533 influenza A viruses from gulls: An evaluation of inter-regional movements and interactions with other avian and mammalian influenza A viruses. Cogent Biology 2. 534 535 33. Tonnessen R, Hauge AG, Hansen EF, Rimstad E, Jonassen CM. 2013. Host 536 restrictions of avian influenza viruses: in silico analysis of H13 and H16 specific signatures in the internal proteins. PLoS One 8:e63270. 537 538 34. Iamnikova SS, Gambarian AS, Aristova VA, L'Vov D K, Lomakina NF, Munster V, 539 Lexmond P, Foucher RA. 2009. [A/H13 and A/H16 influenza viruses: different lines of one precursors]. Vopr Virusol 54:10-8. 540 541 35. Wang ZJ, Kikutani Y, Nguyen LT, Hiono T, Matsuno K, Okamatsu M, Krauss S, 542 Webby R, Lee YJ, Kida H, Sakoda Y. 2018. H13 influenza viruses in wild birds have

undergone genetic and antigenic diversification in nature. Virus Genes 54:543-549.

Swayne DE, Runstadler JA, Happ GM, Senne DA, Wang R, Slemons RD, Holmes

Dugan VG, Chen R, Spiro DJ, Sengamalay N, Zaborsky J, Ghedin E, Nolting J,

546 EC, Taubenberger JK. 2008. The evolutionary genetics and emergence of avian influenza viruses in wild birds. PLoS Pathogens 4:e1000076. 547 Fries AC, Nolting JM, Danner A, Webster RG, Bowman AS, Krauss S, Slemons RD. 37. 548 549 2013. Evidence for the circulation and inter-hemispheric movement of the H14 subtype influenza A virus. PLoS One 8:e59216. 550 38. Lebarbenchon C, Stallknecht DE. 2011. Host shifts and molecular evolution of H7 551 552 avian influenza virus hemagglutinin. Virol J 8:328. 39. Pearce JM, Ramey AM, Ip HS, Gill RE, Jr. 2010. Limited evidence of trans-553 hemispheric movement of avian influenza viruses among contemporary North 554 555 American shorebird isolates. Virus Research 148:44-50. 40. Lycett S, Bodewes R, Pohlmann A, Bank J, Banyai K, Boni M. 2016. Role for 556 migratory wild birds in the global spread of avian influenza H5N8. Science 354:213-557 558 217. 41. Adlhoch C, Gossner C, Koch G, Brown I, Bouwstra R, Verdonck F, Penttinen P, 559 560 Harder T. 2014. Comparing introduction to Europe of highly pathogenic avian 561 influenza viruses A(H5N8) in 2014 and A(H5N1) in 2005. Eurosurveillance 19. 42. Bahl J, Krauss S, Kuhnert D, Fourment M, Raven G, Pryor SP, Niles LJ, Danner A, 562 563 Walker D, Mendenhall IH, Su YC, Dugan VG, Halpin RA, Stockwell TB, Webby RJ, 564 Wentworth DE, Drummond AJ, Smith GJ, Webster RG. 2013. Influenza a virus migration and persistence in North American wild birds. PLoS Pathogens 9:e1003570. 565 566 43. Verhagen JH, Hofle U, van Amerongen G, van de Bildt M, Majoor F, Fouchier RA, 567 Kuiken T. 2015. Long-Term Effect of Serial Infections with H13 and H16 Low-

Pathogenic Avian Influenza Viruses in Black-Headed Gulls. J Virol 89:11507-22.

570

571

587

588

589

590

591

49.

50.

51.

44.

572 45. Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB, Lesniewski RA, Oakley BB, Parks DH, Robinson CJ, Sahl JW, Stres B, Thallinger GG, Van Horn 573 DJ, Weber CF. 2009. Introducing mothur: open-source, platform-independent, 574 575 community-supported software for describing and comparing microbial communities. Appl Environ Microbiol 75:7537-41. 576 46. Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O. 2010. New 577 578 algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. Systematic Biology 59:307-21. 579 47. Keane TM, Creevey CJ, Pentony MM, Naughton TJ, McLnerney JO. 2006. 580 581 Assessment of methods for amino acid matrix selection and their use on empirical data shows that ad hoc assumptions for choice of matrix are not justified. BMC Evol Biol 582 583 6:29. 584 48. Rambaut A, Lam TT, Max Carvalho L, Pybus OG. 2016. Exploring the temporal structure of heterochronous sequences using TempEst (formerly Path-O-Gen). Virus 585 586 Evol 2:vew007.

Brown J, Poulson R, Carter D, Lebarbenchon C, Pantin-Jackwood M, Spackman E,

Shepherd E, Killian M, Stallknecht D. 2012. Susceptibility of avian species to North

American H13 low pathogenic avian influenza viruses. Avian Diseases 56:969-75.

592 inference of past population dynamics from molecular sequences. Mol Biol Evol

sampling trees. BMC Evolutionary Biology 7:214.

finds its roots. PLoS Computational Biology 5:e1000520.

Drummond AJ, Rambaut A. 2007. BEAST: Bayesian evolutionary analysis by

Drummond AJ, Rambaut A, Shapiro B, Pybus OG. 2005. Bayesian coalescent

Lemey P, Rambaut A, Drummond AJ, Suchard MA. 2009. Bayesian phylogeography

593 22:1185-92.

- 594 52. Shapiro B, Rambaut A, Drummond AJ. 2006. Choosing appropriate substitution
- models for the phylogenetic analysis of protein-coding sequences. Mol Biol Evol 595
- 23:7-9. 596
- 597 53. Vijaykrishna D, Bahl J, Riley S, Duan L, Zhang JX, Chen H, Peiris JS, Smith GJ,
- Guan Y. 2008. Evolutionary dynamics and emergence of panzootic H5N1 influenza 598
- viruses. PLoS Pathog 4:e1000161. 599
- 600 54. Lebarbenchon C, Brown JD, Stallknecht DE. 2013. Evolution of influenza A virus H7
- and N9 subtypes, Eastern Asia. Emerg Infect Dis 19:1635-8. 601
- 55. Pond SL, Frost SD, Muse SV. 2005. HyPhy: hypothesis testing using phylogenies. 602
- 603 Bioinformatics 21:676-9.
- 56. Delport W, Poon AF, Frost SD, Kosakovsky Pond SL. 2010. Datamonkey 2010: a 604
- suite of phylogenetic analysis tools for evolutionary biology. Bioinformatics 26:2455-605
- 7. 606
- 57. Kosakovsky Pond SL, Frost SD. 2005. Not so different after all: a comparison of 607
- 608 methods for detecting amino acid sites under selection. Mol Biol Evol 22:1208-22.
- 609 58. Hirst GK. 1943. Studies of Antigenic Differences among Strains of Influenza a by
- Means of Red Cell Agglutination. Journal of Experimental Medicine 78:407-23. 610
- 612 Figure legends

- 613 Figure 1. Maximum clade credibility (MCC) trees for influenza A virus H13 hemagglutinin
- 614 subtype (n= 338). Branches were colored according to most probable geographic origin (red:
- 615 North America; orange: South America; dark blue: Europe; light blue: Asia; green: Oceania;
- gray: not identified). Black node bars represent the 95% highest posterior densities for times 616
- 617 of the common ancestors. Numbers highlight intercontinental gene flow events as detailed in
- Table 2 and Figure S3. Virus strain names and posterior probabilities are detailed in Figure 618

619 S2. 620 Figure 2. Maximum clade credibility (MCC) trees for influenza A virus H16 hemagglutinin 621 622 subtype (n=192). Branches were colored according to most probable geographic origin (red: North America; orange: South America; dark blue: Europe; light blue: Asia; green: Oceania; 623 gray: not identified). Black node bars represent the 95% highest posterior densities for times 624 625 of the common ancestors. Numbers highlight intercontinental gene flow events as detailed in Table 3 and Figure S6. Virus strain names and posterior probabilities are presented in Figure 626 S5. 627 628 Figure 3. Antigenic map of H13 and H16 influenza A viruses (n=44). Different subtypes and 629 genetic clades are indicated with colors (yellow: H13 clade A; orange: H13 clade B; red: H13 630 631 clade C; blue: H16 clade A; purple: H16 clade B; green: H16 clade C). White circles indicate the antisera. Respective virus strains are abbreviated; the full name can be found in Table 5. 632 633 Asterices indicates antigens BHGU/NL/20/09, BHGU/SE/1/06, BHGU/SE/1/03, 634 GBBG/AK/1421/79, BHGU/NL/1/07, HEGU/NY/AI00-532/00 and LAGU/NJ/AI08-0714/08 that had only two numerical HI titers to the tested sera and hence their placement in the map 635 636 is not robust. In this map the distance between the points represents antigenic distance as 637 measured by the hemagglutination inhibition (HI) assay in which the distances between antigens and antisera are inversely related to the log2 HI titer. Each square in the grid of the 638 639 antigenic map equals a two-fold difference in the HI assay. 640 **Tables** 641 642 **Table 1.** Representative viruses selected to generate ferret antisera used to map the antigenic

diversity of H13 and H16 influenza A viruses

	2	÷	
	۶	ς,	
i	7	ŧ.	
	ì	É	
٤	5	5	
Ĺ			
	C)	
Ī	7	ŧ	
	è	₹	
	١,	=	

	1	
Subtype	Clade	Virus strain name
H13	A	A/Gull/Maryland/704/1977 (H13N6)
	A	A/Black-headed gull/Netherlands/2/2007 (H13N6)
	В	A/Ring-billed gull/Georgia/AI00-2658/2000 (H13N6)
	В	A/Gull/Minnesota/1352/1981 (H13N6)
	С	A/Laughing gull/ New Jersey/AI08-0714/ 2008 (H13N9)
	С	A/Great black-headed gull/Astrakhan/1420/1979 (H13N2)
H16	A	A/Black-headed gull/Sweden/2/1999 (H16N3)
	В	A/Herring gull/New York/AI00-532/2000 (H16N3)
	С	A/Black-headed gull/Turkmenistan/13/1976 (H16N3)
	С	A/Black-headed gull/Sweden/5/1999 (H16N3)

- Table 2. Intercontinental gene flow events for influenza A virus H13 hemagglutinin. MRCA: 645
- Most Recent Common Ancestor. HPD: Higher Posterior Density. Event # corresponds to the 646
- 647 numbers indicated in Figure 1 and S3

648

H13 Event Ti		Time of the	Geographic origin of the	Location of					
Clade #		MRCA ± 95%	MRCA (posterior)	introduction					
		HPD							
A	1	1963 [1958-	North America (0.62)	Oceania					

В

C

2	1974 [1972-	North America (0.73)	Europe
	1975]		
3	1988 [1987-	Europe (1)	North America
	1989]		
4	1990 [1988-	Europe (0.82)	South America
	1991]		
5	1996 [1995-	Europe (0.75)	Asia
	1997]		
6	2003 [2003-	Europe (1)	Asia
	2004]		
7	2005 [2004-	Asia (0.48)	North America
	2005]		
8	2009 (2009-	North America (0.9)	Asia
	2010]		
9	2006 [2006-	Europe (0.96)	Asia
	2007]		
10	2011 [2010-	Europe (1)	Asia
	2011]		
11	2013 [2012-	North America (0.96)	South America
	2014]		
12	1987 [1985-	Europe (0.99)	North America
	1988]		

650

13	2002 [2002- 2003]	Europe (1)	Asia
14	2005 [2004- 2005]	Asia (0.55)	North America
15	2010 [2009- 2010]	Europe (1)	North America
16	2004 [2003- 2005]	Europe (0.97)	Asia
17	2013 [2013- 2014]	Europe (0.99)	Asia
18	2014 [2013- 2014]	North America (0.39)	Asia
19	2011 [2010- 2011]	Europe (0.99)	North America
20	2012 [2011- 2012)	North America (0.94)	South America

651 **Table 3.** Intercontinental gene flow events for influenza A virus H16 hemagglutinin. MRCA:

652 Most Recent Common Ancestor. HPD: Higher Posterior Density. Event # corresponds to the

numbers indicated in Figure 2 and S6 653

H16	Event	Time of the	Geographic origin	Location of introduction

Clade	#	MRCA ± 95%	of the MRCA	
		HPD	(posterior)	
С	1	1971 [1968-	Europe (0.97)	Asia
		1972]		
	2	1976 [1976-	Asia (0.71)	Europe
		1976]		
	3	1976 [1972-	Europe (0.86)	North America
		1980]		
	4	1999 [1999-	Europe (1)	Asia
		1999]		
	5	2003 [2002-	Europe (1)	Asia
		2004]		
	6	1999 [1998-	Europe (0.99)	North America
		2000]		
	7	2008 [2007-	Europe (0.99)	Asia
		2009]		
	8	2006 [2005-	Europe (0.97)	North America
	0	2006]	N	
	9	2006 [2006- 2007]	North America (0.55)	South America
	10			North America
	10	2008 [2007- 2009]	Europe (0.63)	norui America
		2007]		

Table 4. Molecular evolution of the HA gene of influenza A virus subtypes H13 and H16 655

Genetic lineage	\mathbf{N}^1	Time period ²	ution rate ³	d _N /d _S	
			Mean	95% HPD	Mean
H13	338	40	3.8	3.6-4.1	0.13
H13 - A	54	39	3.8	2.3-4.9	0.09
H13 - B	76	39	0.8	0.6-1.0	0.18
H13 - C	208	37	5.5	5.0-6.0	0.16
H16	192	41	3.1	2.8-3.4	0.09
H16 - A	56	33	4.5	3.9-5.2	0.10
H16 - B	19	35	4.6	3.9-5.2	0.06
H16 - C	117	40	1.5	1.2-1.8	0.11

¹ number of nucleotide sequences included in the analysis; ² in years; ³ per 10⁻³ substitution / site / 656

657 year; HPD: highest posterior density.

Table 5. Hemagglutinin inhibition data of H13 and H16 influenza A viruses (n=44) 659

Suptype						H	13				H	116	
Clade				A	A	В	В	C	C	A	В	C	C
	Virus name	Subtype	Virus abbreviation										
				BHGU/NL/2/07	GULL/ML/704/77	GULL/MN/1352/81	RBGU/GE/AI00-2658/00	GBBG/AK/1420/79	LAGU/NJ/AI08-714/08	BHGU/SE/2/99	HEGU/NY/AI0-532/00	BHGU/SE/5/99	BHGU/TM/13/76
H13 / A	A/Black-headed gull/Netherlands/2/07	H13N6	BHGU/NL/2/07	320	280	80	<10	20	<10	<10	<10	<10	25
	A/Black-headed gull/Netherlands/4/07	H13N6	BHGU/NL/4/07	1280	400	320	<10	35	<10	<10	<10	10	40
	A/Black-headed gull/Netherlands/7/09	H13N2	BHGU/NL/7/09	10	160	<10	<10	<10	<10	10	<10	<10	15
	A/Black-headed gull/Sweden/10/05	H13N6	BHGU/SE/10/05	240	320	40	<10	10	<10	<10	<10	<10	15
	A/Great-black headed gull/Sweden/1/03	H13N6	GBBG/SE/1/03	80	240	20	<10	<10	<10	<10	<10	<10	<10
	A/gull/ML/704/77	H13N6	GULL/ML/704/77	40	240	20	<10	< 20	<10	<10	<10	<10	<10
H13 / B	A/gull/MN/1352/81	H13N6	GULL/MN/1352/81	120	160	320	<10	20	<10	<10	<10	<10	<10
	A/gull/NJ/34/92	H13N6	GULL/NJ/34/92	80	240	80	<10	240	<10	<10	<10	<10	<10
	A/Herring gull/DB/13/90	H13N2	HEGU/DB/13/90	40	140	140	10	25	<10	<10	<10	<10	<10
	A/Laughing gull/DB/1370/86	H13N2	LAGU/DB/1370/86	10	40	<10	10	40	<10	<10	<10	<10	<10
	A/ring-billed gull/GE/AI00-2658/00	H13N6	RBGU/GE/AI00- 2658/00	10	60	40	<u>640</u>	15	<10	<10	<10	<10	<10
	A/ring-billed gull/MN/AI10-1708/10	H13N6	RBGU/MN/AI10- 1708/10	80	200	120	10	10	<10	<10	<10	<10	<10
H13 / C	A/Black-headed gull/Netherlands/1/00	H13N8	BHGU/NL/1/00	35	<10	<10	<10	1280	120	<10	30	<10	30
	A/Black-headed gull/Netherlands/20/09	H13N2	BHGU/NL/20/09	<10	<10	<10	<10	280	<10	<10	<10	<10	35
	A/Black-headed gull/Netherlands/4/08	H13N8	BHGU/NL/4/08	<10	<10	<10	<10	140	80	<10	<10	<10	25
	A/Black-headed gull/Sweden/1/03	H13N8	BHGU/SE/1/03	<10	<10	<10	<10	560	40	<10	<10	<10	<10

Jour	position Clade														
	H13 A	D	A,T,S	A	D,E,N,S	K,Q	K	T	V,L V,I	V	E	S,G	K	S,L S,L	
	H13 B	D	A,T,S	Α	D,N,S	K,R	G,R	T	V,I	T,A	E	S,G	S,R,N,H	S,L	
	H13 C	D	V,A	A	DEL,R	K,R,S	G,R	T,A,V	V,I	T,A,E	E	D,N,S	S,R,G	S,T]
	H16 A H16 B	D E D	T V	S S	DEL DEL	L DEL	G G	E D	D D	E E,?	T T,V	K K	K K,E	E E	
	H16 C	D	V,A	S	DEL	K,DEL	G	E,D	D	E	T	K	K	E	
	665														
\leq															

	A/Black-headed gull/Sweden/1/06	H13N8	BHGU/SE/1/06	<10	<10	<10	<10	120	<10	<10	<10	<10	<10
	A/Black-headed gull/Sweden/1/99	H13N6	BHGU/SE/1/99	10	<10	10	30	160	<10	<10	<10	<10	10
	A/Black-headed gull/Sweden/2/03	H13N8	BHGU/SE/2/03	<10	<10	<10	<10	200	50	<10	<10	<10	10
	A/Great-black headed gull/AK/1420/79	H13N2	GBBG/AK/1420/79	10	35	10	<10	2720	160	10	<10	35	25
	A/Great-black headed gull/AK/1421/79	H13N2	GBBG/AK/1421/79	<10	<10	<10	<10	140	80	<10	<10	<10	<10
	A/Great-black headed gull/AK/591/82	H13N2	GBBG/AK/591/82	<10	40	<10	<10	480	100	<10	<10	40	80
	A/Great-black headed gull/GJ/76/83	H13N2	GBBG/GJ/76/83	<10	<10	<10	<10	320	80	<10	<10	<10	30
	A/Herring gull/AK/458/85	H13N6	HEGU/AK/458/85	30	20	<10	<10	1920	480	70	<10	80	80
	A/Herring gull/AK/479/85	H13N6	HEGU/AK/479/85	140	35	10	<10	1920	640	280	120	280	120
	A/Laughing gull/NJ/AI08-714/08	H13N9	LAGU/NJ/AI08-	<10	<10	<10	<10	320	560	<10	<10	<10	<10
	0 00		714/08										
H16 / A	A/Black-headed gull/Netherlands/5/07	H16N3	BHGU/NL/5/07	35	25	<10	<10	140	<10	960	160	320	640
	A/Black-headed gull/Netherlands/1/07	H16N3	BHGU/NL/1/07	<10	<10	<10	<10	<10	<10	80	<10	<10	40
	A/Black-headed gull/Netherlands/10/09	H16N3	BHGU/NL/10/09	20	80	<10	<10	280	15	1280	160	640	640
	A/Black-headed gull/Netherlands/21/09	H16N3	BHGU/NL/21/09	70	200	20	<10	240	<10	480	<10	240	280
	A/Black-headed gull/Netherlands/3/07	H16N3	BHGU/NL/3/07	100	90	20	<10	100	<10	120	140	60	120
	A/Black-headed gull/Sweden/2/99	H16N3	BHGU/SE/2/99	10	<10	<10	<10	10	<10	960	80	35	380
	A/Black-headed gull/Sweden/8/05	H16N3	BHGU/SE/8/05	<10	<10	<10	<10	10	<10	1280	<10	30	140
H16/B	A/Herring gull/DB/2617/87	H16N3	HEGU/DB/2617/87	<10	<10	<10	<10	<10	<10	<10	120	20	1600
	A/Herring gull/NY/AI0-532/00	H16N3	HEGU/NY/AI0-	<10	<10	<10	<10	<10	<10	<10	320	<10	320
			532/00								_		
	A/Laughing gull/DB/2839/87	H16N3	LAGU/DB/2839/87	<10	<10	<10	<10	<10	<10	160	80	20	1920
H16/C	A/Black-headed gull/Netherlands/26/09	H16N3	BHGU/NL/26/09	10	25	<10	<10	20	<10	30	80	20	1280
	A/Black-headed gull/Sweden/5/99	H16N3	BHGU/SE/5/99	10	<10	<10	<10	70	<10	560	30	1600	400
	A/Black-headed gull/TM/13/76	H16N3	BHGU/TM/13/76	25	30	<10	<10	27.5	<10	50	320	100	4800
	A/environment/Sweden/2/05	H16N3	ENV/SE/2/05	20	30	10	<10	140	30	960	320	1280	640
	A/Little tern/Sweden/1/05	H16N3	LITE/SE/1/05	<10	15	<10	<10	15	<10	10	30	20	1280
	A/shorebird/DB/172/05	H16N3	SB/DB/172/05	<10	<10	<10	<10	30	<10	240	60	200	1280
	A/shorebird/DB/195/06	H16N3	SB/DB/195/06	<10	<10	<10	<10	<10	<10	<10	30	20	560
	A/Slender-billed gull/AK/28/76	H16N3	SBGU/AK/28/76	20	140	10	<10	50	<10	80	160	100	1280

Table 6. Amino acid differences within/near the receptor binding site of the HA protein 661 662 among H13 and H16 subtypes and clades, based on the HA gene of H13 (n=338) and H16 (n=192) LPAIVs, including the 130-loop (position 136-147 according to Burke & Smith 663 2014), 190-helix (200-208) and 220-loop (230-240). DEL, deletion of amino acid. 664

Amino acid position	139	142	145	149	166	176	177	196	198	200	208	217	218	224	231	233
Clade																
H13 A	D	A,T,S	A	D,E,N,S	K,Q	K	T	V,L	V	Е	S,G	K	S,L	K	P	Y
H13 B	D	A,T,S	A	D,N,S	K,R	G,R	T	V,I	T,A	E	S,G	S,R,N,H	S,L	K,N	P,L	Y,
H13 C	D	V,A	A	DEL,R	K,R,S	G,R	T,A,V	V,I	T,A,E	Е	D,N,S	S,R,G	S,T	N,T,K	P	Q Y
H16 A	E	T	S	DEL	L	G	E	D	E	T	K	K	E	E	I	D
H16 B	D	V	S	DEL	DEL	G	D	D	E,?	T,V	K	K,E	E	E	I	D,E
																,N
H16 C	D	V,A	S	DEL	K,DEL	G	E,D	D	E	T	K	K	E	E	I,V	D,
																N





