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Subclinical avian hepatitis e virus infection in layer flocks in the united states

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Highlights

- Circulation of avian HEV was determined in healthy commercial layer farms in the
 United States.
- A fluorescent microbead immunoassay (FMIA) was developed for detecting anti-avian
 HEV IgY antibodies.
- Prevalence of avian HEV infection was high among clinically healthy laying hens.
 - Different avian HEV genotype 2 strains were circulating within a farm.

Abstract

The objective of this study was to determine patterns of avian HEV infection in
naturally infected chicken farms. A total of 310 serum samples and 62 pooled fecal samples
were collected from 62 chicken flocks on seven commercial in-line egg farms in the
Midwestern United States and tested for avian HEV circulation. Serum samples were tested
for presence of anti-avian HEV IgY antibodies by a fluorescent microbead immunoassay
(FMIA) which was developed for this study. The FMIA was validated using archived
samples of chickens with known exposure ($n = 96$) and compared to the result obtained with
an enzyme-linked immunosorbent assay (ELISA) based on the same capture antigen. There
was an overall substantial agreement between the two assays ($\kappa = 0.63$) with earlier detection
of positive chickens by the FMIA ($P = 0.04$). On the seven farms investigated, the overall
prevalence of anti-avian HEV IgY antibodies in serum samples from commercial chickens
was 44.8% (20-82% per farm). Fecal samples were tested for avian HEV RNA by a nested
reverse-transcriptase PCR. The overall detection rate of avian HEV RNA in fecal samples
was 62.9% (0-100% per farm). Sequencing analyses of partial helicase and capsid genes
showed that different avian HEV genotype 2 strains were circulating within a farm. However,

45	no correlation was	found between	avian HEV RNA	detection and e	gg production.	egg weigh

- 46 or mortality. In conclusion, avian HEV infection is widespread among clinically healthy
- 47 laying hens in the United States.

- 49 Keywords: Avian hepatitis E virus (avian HEV); Chickens; Fluorescent microsphere
- 50 immunoassay, RT-PCR, Subclinical infection.



Introduction

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52	Hepatitis E virus (HEV) are non-enveloped, single-stranded RNA viruses with an
53	icosahedral capsid symmetry that belongs to the family <i>Hepeviridae</i> (Meng et al., 2012).
54	HEV has been identified in several animal species (Meng, 2013) and, based on host tropism
55	and genetic relatedness, strains genetically characterized thus far can be classified into two
56	genera: genus Orthohepevirus (all mammalian and avian hepatitis E virus [avian HEV]
57	isolates) and genus <i>Piscihepevirus</i> (cutthroat trout virus) (Smith et al., 2014). Currently, four
58	avian HEV genotypes have been described in chicken flocks worldwide (Johne et al., 2014).
59	Genotype 1 has been identified in Australia and Korea, genotype 2 is present in North
60	America, genotype 3 is present in Europe and China and, more recently, a novel putative
61	genotype 4 has been detected in Hungary and Taiwan (Marek et al., 2010; Banyai et al.,
62	2012; Kwon et al., 2012; Hsu and Tsai, 2014a). The avian HEV genome contains three open
63	reading frames (ORFs): ORF1 encodes non-structural viral proteins, ORF2 encodes the
64	capsid protein and ORF3 encodes a small multifunctional phosphoprotein (Meng et al.,
65	2012).
66	
67	A HEV-related sequence was first detected in chickens with big liver and spleen
68	disease in Australia (Payne et al., 1999). A similar disease manifestation, designated as
69	hepatitis-splenomegaly syndrome (HSS), was first associated with avian HEV in North
70	America (Haqshenas et al., 2001). In both broiler breeder hens and egg-type chickens ranging
71	from 30 to 72 weeks of ages, HSS is clinically characterized by increased mortality and
72	decreased egg production. On gross examination, blood-tinged fluid can frequently be
73	observed in the coelomic cavity and livers and spleens are typically both enlarged (Meng and

Shivaprasad, 2013). A significant proportion of chicken flocks worldwide are seropositive to

avian HEV, even though seropositive flocks do not necessarily suffer from HSS (Meng and

Shivaprasad, 2013). Thus far, no clear association between avian HEV strains and
pathogenicity has been established (Marek et al., 2010). Avian HEV isolates obtained from
healthy chickens were able to induce liver lesions but did not produce clinical HSS in
specific-pathogenic-free chickens (Billam et al., 2009). However, apparently healthy
chickens with reduced egg production have been shown to be positive for avian HEV RNA i
the absence of other pathogens, suggesting that subclinical infection with avian HEV may
impair egg production in broiler breeders (Sprygin et al., 2012). Therefore, additional studies
are needed to elucidate the significance of avian HEV infections in clinically healthy laying
hens.

Current methods to demonstrate avian HEV infection include conventional and real-time reverse transcriptase (RT) PCR assays for detection of avian HEV RNA (Sun et al., 2004; Bilic et al., 2009; Troxler et al., 2011) and commercial and in-house enzyme-linked immunosorbent assays (ELISA) and agar gel immunodiffusion (AGID) tests for detection of anti-avian HEV antibodies (Huang et al., 2002; Morrow et al., 2008; Zhao et al., 2013; Hsu and Tsai, 2014b). ELISAs based on the ORF2 capsid protein have been widely adapted and are commonly used in serological surveys (Peralta et al., 2009; Kwon et al., 2012; Zhao et al., 2013; Liu et al., 2014). Fluorescent microbead immunoassays (FMIA) are increasingly being used in veterinary serology (Wagner et al., 2011; Langenhorst et al., 2012; Gimenez-Lirola et al., 2014). An advantage of this new technology is the ability to screen for antibodies against multiple pathogens simultaneously in one reaction well using a small amount of sample, thus saving time, labor and reagents. In addition, FMIAs may have improved sensitivity compared to conventional serological assays, such as ELISAs (van Gageldonk et al., 2008).

The objective of this study was to evaluate the avian HEV infection dynamics,
including fecal avian HEV RNA shedding and prevalence of anti-avian HEV antibodies, in
healthy layer hen flocks in the Midwestern United States. For serology purposes, an FMIA to
detect anti-avian HEV IgY antibodies was developed, validated and compared with an
indirect ELISA.

Material and methods

Farm selection and characterization

Seven commercial egg farms, designated A through G, volunteered to participate in this study. The final number of participating farms was determined based on responses to an email sent to the farms' managers and/or veterinarian-in-charge, and to on-farm availability to provide the specified samples and production data. Participating farms were located in four U.S. Midwestern states: Iowa, Illinois, Michigan and South Dakota. At the time of sample collection Iowa represented the largest U.S. egg producer and Michigan was ranked seventh. The number of sampled farms did not reflect the state's egg production. Each farm had an egg production facility wherein eggs were collected from multiple chicken houses and transported to an egg processing room via a collection of belts and conveyors. These in-line operations produced washed and graded whole shell eggs for retail and/or food service customers. Chickens on each farm were vaccinated against Marek's disease, infectious bronchitis, Newcastle disease, avian influenza, fowl pox, *Mycoplasma gallisepticum*, and *Salmonella* Enteritidis.

Samples were collected between October and December 2013. At the time of sample collection, chickens in flocks on participating farms were healthy and none had clinical signs associated with HSS. A flock consisted of all chickens housed in one building. All flocks on

125	each farm were sampled ($n = 5$ to 11, average $n = 8$). Production data available for each flock
126	included average hen-day egg production (%), weekly mortality (%), and egg weight.
127	
128	Sample collection
129	A total of 310 serum samples were collected from 62 flocks divided among seven egg
130	farms. From each flock, blood samples were taken from each of five chickens arbitrarily
131	selected from different locations within a house. Blood samples were acquired by
132	venipuncture of the brachial vein in 5 ml serum tubes ($n = 25$ to 55/farm; average $n = 42$). In
133	addition, one pooled fecal sample was obtained from each house by collecting and pooling 2
134	g of fresh droppings from manure pits at five arbitrarily selected sites.
135	
136	Blood and fecal samples were collected on the same day from all flocks on a farm,
137	labeled with the house number, placed in insulated boxes with ice packs (4 °C) and shipped
138	on the day of collection to the Iowa State University Veterinary Diagnostic Laboratory (ISU-
139	VDL), Ames, Iowa. Upon arrival, blood was centrifuged at $1500 \times g$ for 10 min to harvest
140	serum and aliquotted into 4 mL plastic tubes. Fecal samples were homogenized and an
141	aliquot of 0.4 g was resuspended in 4 ml PBS, vigorously vortexed, and centrifuged at 1500
142	$\times g$ for 10 min. All samples were stored at -80 °C until testing. Shipping and storage
143	conditions were similar for all samples.
144	
145	Fluorescent microbead immunoassay (FMIA) development
146	Experimental serum samples
147	Ninety-six serum samples from 36 specific-pathogen-free chickens from a previous avian
148	HEV study (Billam et al., 2009) were used as positive and negative controls to develop the
149	FMIA. Briefly, 24 6-week-old chickens were intravenously inoculated with avian HEV

genotype 2. Blood samples were collected before inoculation and weekly thereafter for a total
of four weeks. Twelve chickens where sham-inoculated and served as negative controls.
Seroconversion started at day post-inoculation (dpi) 14 and at dpi 28 all inoculated chickens
had seroconverted to avian HEV as previously determined by an in-house ELISA (Billam et
al., 2009). A total of 36 negative control samples (all samples collected on dpi 0) and 60
samples (dpi 7-28) from chickens infected experimentally were tested. Specifically, negative
control samples were used for estimating diagnostic specificity and 24 serum samples from
experimentally infected chickens collected on dpi 21 ($n = 12$) or 28 ($n = 12$) were used for
estimating diagnostic sensitivity of the FMIA.
Antigen and conjugation to carboxylated paramagnetic microbeads
A truncated recombinant ORF2 capsid protein of avian HEV expressed in Escherichia
coli and purified by affinity chromatography as described previously (Haqshenas et al., 2002)
was used as antigen for the FMIA. Fluorescent microsphere coupling was performed using a
method described previously (Gimenez-Lirola et al., 2012) by addition of 25 μg of the avian
HEV recombinant ORF2 capsid protein to 2.5×10^6 carboxylated-fluorescent microbeads
(bead region 64, Luminex Corp., Austin, TX, USA).
<u>FMIA</u>
The assay was performed as described previously (Gimenez-Lirola et al., 2012) with
exception that a 1:2000 dilution of a biotin-conjugated goat anti-chicken IgY Fc (Gallus
Immunotech) was used. Samples were analyzed using a Luminex-100 flow cytometer
(Luminex) at default settings set by the manufacturer. Median fluorescence intensity (MFI) of
the reporter signal estimated from 50 beads was used for the data analysis. A set of internal
standard controls described as high positive control, low negative control and cut off control,

175	were selected after initial analysis and subsequently included on each plate. Additionally, a
176	blank control (serum diluent) was included on each plate. The MFI data was corrected for
177	background levels by subtracting the blank control MFI from the sample MFI (MFI-
178	Bkg). Results were presented as antibody index (MFI-Bkg sample/MFI-Bkg cut off serum).
179	Samples with an index value below 0.9 were considered negative and those above 1.1 were
180	considered positive. Values from 0.9 to 1.1 were considered inconclusive.
181	
182	Cut-off value determination
183	Sensitivity and specificity were evaluated using receiver operator characteristic (ROC) curve
184	analysis. The optimal cut-off points determined to be a sample MFI value ranging between
185	1008 and 1297 giving an overall diagnostic sensitivity and specificity of 100%. The
186	cumulative area under the ROC curve (AUC) indicated that the aHEV ORF2-based FMIA
187	was 100% accurate. The analysis was performed using GraphPad Prism v. 6.01 (GraphPad
188	Software).
189	×O
190	Assay reproducibility
191	The reproducibility of the FMIA was evaluated by utilizing six different chicken sera. The
192	coefficient of variation (CV) was used to evaluate the intra- and inter-assay variation. The
193	inter-assay CV determined by each sample tested in three different runs on different
194	occasions ranged from 0.57% to 7.83%; while the intra-assay CV determined by three
195	replicates tested within the same run ranged from 0.74% to 7.99%, indicating that the results
196	were reproducible.
197	
198	Comparison of FMIA and ELISA

199	In order to evaluate the FMIA, a subset of the experimental $(n = 48)$ and the field $(n = 48)$
200	= 56) serum samples were tested for the presence of specific anti-avian HEV IgY antibodies
201	by an in-house ELISA described elsewhere (Billam et al., 2009) using the same antigen used
202	in this study. For comparison, the results were presented as an antibody index as specified for
203	the FMIA test (sample OD or MFI-Bkg/cut off serum mean OD or MFI-Bkg).
204	
205	RNA extraction and avian HEV RNA detection
206	RNA extractions from the fecal samples were performed using the QIAamp Viral
207	RNA Mini kit (Qiagen). Extracts were subsequently used for detection of the partial helicase
208	and capsid genes of avian HEV as described previously (Sun et al., 2004) in nested RT-PCR
209	reactions. Briefly, for the helicase gene detection, external primer set 5'-
210	TGTTATYACACCCACCAARACGYTG-3', and 5'-CCTCRTGGACCGTWATCGACCC-
211	3'; and internal primer set 5'-GCCACGGCTRTTACACCYCAYGT-3', and 5'-
212	GACCCRGGRTTCGACTGCTT-3' were used. For the capsid gene, external primer set 5'-
213	TCGCCYGGTAAYACWAATGC-3', and 5'-GCGTTSCCSACAGGYCGGCC-3'; and
214	internal primer set 5'- ACWAATGCYAGGGTCACCCG -3', and 5'-
215	ATGTACTGRCCRCTSGCCGC -3' were used. PCR products were examined on a 1%
216	agarose gel and amplicons with the expected size (386 bp for the helicase gene and 242 bp
217	for the capsid gene) were excised and purified with the QIAquick Gel Extraction Kit
218	(Qiagen).
219	
220	Sequencing and phylogenetic analyses
221	Sequencing of at least two avian HEV capsid and two helicase RT-PCR positive
222	samples from each farm was performed directly on both strands at the Iowa State University
223	DNA Facility, Ames, Iowa, USA (Applied Biosystems 3730xl DNA Analyzer). Sequences

were aligned with published data using BLAST at the national Centre for Biotechnology Information (NCBI)¹. Sequences were compiled using Lasergene software and the Clustal V alignment algorithm (DNAStar). Identical nucleotide sequences were represented as one sequence and used in phylogenetic analysis. For sequence analysis, the 361 bp sequences of the helicase gene and 232 bp of the capsid gene were compared to each other and to sequences of other avian HEV isolates representing genotypes 1 (GenBank accession numbers AM943647 and JN597006), 2 (GenBank accession numbers AY535004, EF206691, and EU919187), 3 (GenBank accession numbers AM943646 and GU954430), and 4 (GenBank accession numbers JN997392 and KF511797). Sequences reported herein were deposited in the GenBank database under the accession numbers KJ495790 to KJ495804 and KJ914879 to KJ914889.

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Statistical analysis

Summary statistics including normality were calculated for all the farms. For each flock actual egg production, mortality and egg weights were compared to the breed standards. Breed standard values were defined and provided by the genetic companies for each week of the egg chicken production cycle for the specific genetic line utilized and were valid for the year in which samples were collected. Breed standards are publically shared by all genetic companies and are applicable for any commercial farm located in the U.S. Statistical analysis of the data among farms was performed by one-way analysis of variance (ANOVA). Pearson's correlation test was used to correlate presence of avian HEV RNA in feces with egg production, egg weight, and hen mortality. A kappa index (κ) was performed to determine the agreement of positive and negative results between ELISA and FMIA. The strength of agreement was scored as follows: ≤0, poor; 0.01 to 0.2, slight; 0.21 to 0.4, fair;

¹ http://www.ncbi.nlm.nih.gov

248	0.41 to 0.60, moderate; 0.61 to 0.80, substantial; and 0.81 to 1, almost perfect. $P < 0.05$ was
249	set as the statistically significant level. The statistical analyses were performed using the SAS
250	v.9.2 (SAS Institute).
251	
252	Results
253	Evaluation and validation of the FMIA
254	The earliest detection of avian HEV-specific IgY antibodies was detected at dpi 14 in
255	29.2% (7/24) avian HEV infection. On dpi 21 and onwards all 24 chickens were positive for
256	anti-avian HEV IgY antibodies.
257	
258	To further evaluate the avian HEV FMIA, the same antigen used in the FMIA was
259	coated on 96-wells ELISA plates. Internal FMIA controls, 48 experimental serum samples,
260	and 56 field serum samples were tested by the ELISA. There was a substantial agreement
261	between ELISA and FMIA when using the ORF2 recombinant capsid antigen ($\kappa = 0.63$).
262	Although there was no difference for the overall detection rates between assays (43.4%
263	[53/122] FMIA vs. 36.0% [44/122] ELISA; $P = 0.052$), when considering only the
264	experimental samples subset, FMIA showed a higher detection rate at 14 dpi (7/12 vs. 1/12, P
265	= 0.04) (Table 1).
266	
267	Production data
268	Egg production, mortality and egg weights were compared to breed standards.
269	Average egg production and average mortality data for participating flocks on each of the
270	seven investigated farms are shown in Fig. 1. A summary of median, maximum and
271	minimum mortality, egg production and egg weights for each farm are shown in Fig. 2.

273	Overall performance of all farms at the time of sample collection was satisfactory and
274	mortality rates were mostly within an acceptable range when compared to breed standards
275	(average \pm SD, 0.13 \pm 0.06) with exception of Farm B that presented an average 10%
276	increase in mortality rates ($P < 0.05$) (Fig. 2). Maximum reported weekly mortality was
277	0.27% at 87 weeks of age on Farm B, 0.27% at 81 weeks of age on Farm E, and 0.27% at 109
278	weeks of age on Farm G (Fig. 1).
279	
280	Egg production met or exceeded standard egg production curves in 5 of 7 farms (Figs.
281	1 and 2). Specifically, egg production on Farms A and B was on average 4 to 7% lower when
282	compared to the breed standards while Farm C presented on average a 3.5% higher egg
283	production ($P < 0.05$) (Fig. 2). On Farm A, egg production in 92 to 97 week old flocks was
284	on average 11% less than expected and on Farm B, egg production decrease was on average
285	18% from 87 to 94 weeks of age (Fig. 1). On Farm D, there was a 12% drop in egg
286	production in the 89 week old flock; however the overall egg production was on average 5%
287	higher than the breed standard in the other flocks (Fig. 1). Lack of egg production from hens
288	on Farm B at 77 weeks of age and on Farm G at 76 weeks of age was due to induction of
289	molt prior to onset of a second production cycle, a routine procedure used by commercial egg
290	producers in the Midwestern United States.
291	
292	Egg weights were within an acceptable range when compared to breed standards with
293	exception of a flock in Farm G had a maximum decrease of 10.09% when compared to the
294	standard (Fig. 2). In all other flocks the maximum egg weight decrease was lower than 5.00%
295	(average \pm SD, 0.97 \pm 2.67) with exception of Farm F that exceed the breed standards (Fig.
296	2).
297	

Detection of anti-avian HEV IgY antibodies		Detection	of anti-avia	an HEV IgY	antibodies
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The detection rates of anti-avian HEV IgY antibodies are shown in Fig. 1 and Table 2.
All seven investigated chicken farms had at least one seropositive animal. Considering all
farms, the overall detection rate of anti-avian HEV IgY antibodies was 44.8% (145/310). The
proportion of positive animals ranged from 20% (5/25, Farm F) to 82.0% (41/50, Farm B) per
farm. Considering all the examined animals by rank of age, seropositive animals were
detected at any age, but chickens older than 50 weeks were found to have higher
seropositivity rates compared to 16-26 and 31-45 weeks old chickens ($P < 0.05$).

Detection of avian HEV RNA and correlation with production data

The detection rates of avian HEV RNA are shown in Fig. 1 and Table 2. The overall detection rate of avian HEV RNA in the pooled fecal samples of the seven studied farms was 62.9% (39/62) considering the combined results of both RT-PCR assays utilized (capsid gene, 32/62; helicase gene, 38/92). The majority of flocks (31/39) were RT-PCR positive for both capsid and helicase genes. However; on Farm B, the 69 week old flock was only positive for the capsid gene, and the 91week old flock was only positive for the helicase gene. On Farm C, the flocks with layer hens at 16, 23, 23, 43, 94 and 103 weeks of age were positive only for the helicase gene.

All tested samples from Farms E (n = 7) and F (n = 5) were negative for avian HEV RNA which coincided with the lowest detection of anti-avian HEV IgY antibodies among all tested farms, 22.9% and 20%, respectively (Table 2). Avian HEV RNA was detected in 40% (4/10) of the samples on Farm G and in 75-100% of the samples on Farms A, B, C, and D (Table 2). No correlation was found between avian HEV RNA detection and hen-day egg production (P = 0.09), weekly hen mortality (P = 0.37) or egg weight (P = 0.15).

323	
324	Sequence and phylogenetic analyses
325	At least two avian HEV RNA positive samples were selected for sequencing of both a
326	361 bp region in the helicase gene and a 232 bp region of capsid gene in each farm. The
327	sequences obtained were compared with the reference strains of each avian HEV genotype.
328	All sequences clustered within avian HEV genotype 2 previously identified in the USA (Fig.
329	3).
330	
331	Sequence analysis of the helicase gene from Farms A to G presented 82.3-99.7% of
332	nucleotide sequence identity. Within a farm, nucleotide sequence identities varied from 85.2-
333	99.7%. Between sequences described here and the USA prototype avian HEV, the nucleotide
334	sequence identify was 84.9-88.6% and it was 84.9-97.2% to the avirulent avian HEV-VA
335	strain. Sequence identities with avian HEV genotype 1, 3 and 4 isolates varied between 76.5-
336	82.5%.
337	×O
338	Sequence analysis based on the capsid gene revealed similar results. Among farms,
339	nucleotide sequence identity of avian HEV strains varied between 80.6-97.0%. Within a
340	farm, nucleotide sequence identities varied from 85.2-97.3%. Between sequences described
341	here and the USA prototype avian HEV and avirulent avian HEV-VA strains, there was 85.1-
342	88.9% and 79.7-90.1% nucleotide sequence identity, respectively. Sequence identities with
343	avian HEV genotype 1, 3 and 4 isolates varied between 73.8-77.9%.
344	
345	Discussion
346	Serological surveys have shown a worldwide distribution of avian HEV infection,
347	although clinical cases of HSS seem to occur infrequently (Meng and Shivaprasad, 2013).

Often avian HEV is considered as minor causative agent in diagnostic investigations but is frequently regarded as insignificant when another poultry disease has been diagnosed in the same chickens. However, drops in egg production and increased mortality have been reported in clinically healthy chickens infected with avian HEV in the absence of other known pathogens (Sprygin et al., 2012). To further investigate the patterns of avian HEV infection in apparently healthy chickens, fecal samples were tested for HEV RNA by RT-PCR and serum samples were tested for anti-avian HEV IgY antibodies by FMIA. In addition, mortality rates and egg production, parameters usually affected during HSS outbreaks, and egg weight as indirect method to evaluate the liver function (Husbands, 1970), were acquired from the flocks. Egg weight is largely determined by yolk weight (Jaffe, 1964), and yolk is synthesized in the liver hepatocytes.

Although Farm B, which presented the highest detection rates of avian HEV, showed the overall highest increase in mortality and highest decrease in egg production within the farms sampled, no associations between egg production data, mortality rates and detection rates of avian HEV RNA were found when using the cumulative data of all farms. Different management factors and biosecurity measures of each farm, diet, host genetics and presence of other pathogens could partially explain differences between production performances among flocks (Gerber et al., 2014). In fact, a prospective study has found that nearly identical avian HEV nucleotide sequences have been retrieved from broiler breeder flocks over two years, independent of clinical signs (Troxler et al., 2014). In this same study, viral load did not seem to have an effect on pathogenicity (Troxler et al., 2014). Recently, a pilot study using experimentally infected broiler breeder chickens could reproduce more typical macroscopic and microscopic avian HEV lesions than previous studies with young chickens (Park et al., 2015). However, production losses commonly associated with avian HEV

outbreaks have not been reproduced experimentally to date. Further systemic prospective studies addressing avian HEV subclinical infection and its impact on production data on a larger scale are needed.

Testing serum samples from chickens with unknown avian HEV exposure in this study indicated that the seropositive rate of anti-avian HEV IgY antibodies was 44.8% (145/310) in sexually mature chickens at various stages of egg production and that all farms studied were seropositive for anti-avian HEV IgY antibodies with a detection rate ranging from 20% to 82% per farm. This finding is in agreement with previous reports from the U.S. and other countries (Huang et al., 2002; Peralta et al., 2009; Kwon et al., 2012; Zhao et al., 2013). The present study also suggests that the likelihood of being seropositive increases with age, as previously described (Huang et al., 2002; Peralta et al., 2009; Troxler et al., 2014).

Avian HEV RNA was detected in five of the seven farms tested, including in hens close to the end of the egg production (> 100 weeks of age) and in farms with high numbers of avian HEV seropositive chickens. This suggests that the humoral response may not offer complete protection against avian HEV fecal shedding and/or that chickens might be reinfected at different points of the production cycle with different strains of avian HEV. Indeed, sequencing of avian HEV partial helicase and capsid genes showed that different strains of avian HEV existed on the same farm. Similarly, HEV re-infection in seropositive animals has been reported in adult pigs (de Deus et al., 2008). Possible factors that could explain the co-circulation of different avian HEV sequences on a farm include co-mingling animals from different sources, a common food supplier or shared workers between different farms; however, these factors were not investigated in the present study. Although there is no currently available information regarding the cross-protection among different avian HEV

genotypes, vaccination of pigs with recombinant capsid antigens derived from HEV isolated from swine, rat, and avian induced a strong IgG anti-HEV antibody response but only a partial cross-protection to a HEV genotype 3 challenge (Sanford et al., 2012).

Interestingly, avian HEV RNA was not detected in any of the flocks in Farms E and F although anti-avian HEV antibodies were detected in most of the flocks on all farms investigated. The failure in detecting avian HEV genome could be due to a lower amount of viral shedding into the fecal material of chickens without clinical signs. As pooled samples were used for the analysis, the dilution effect could have decreased the viral copies below the detection limit of the assay used. Alternatively as the avian HEV genome shows a high variability (Sprygin et al., 2012), the primers used for avian HEV RNA detection in the present study may have not amplified the viral genome due to mismatches in the sequences. In addition, lack of detection could be due to poor quality of samples and possible degradation of viral RNA, although samples from all farms were processed and tested similarly. It is also worth noting that Farms E and F contained the lowest number of flocks among the farms sampled in the present study which could have decreased the chances of detecting positive samples if the viral shedding was low. A higher number of fresh fecal samples from each flock should be tested to address these issues.

In conclusion, avian HEV infection was detected directly (RNA) or indirectly (antibodies) in all studied U.S. farms. Different strains of avian HEV genotype 2 were found in different flocks within a farm. There was no clear association between levels of egg production or mortality and detection rates of avian HEV RNA.

Conflict of interest statement

423	None of the authors of this paper has a financial or personal relationship with other
424	people or organizations that could inappropriately influence or bias the content of the paper.
425	
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553	Figure legends
554	
555	Fig. 1. Actual and standard egg production (%), actual and standard weekly mortality (%),
556	anti-avian HEV IgY antibodies detection rate by FMIA (%) (grey bars) and avian HEV RNA
557	detection by nested RT-PCR (red boxes indicate the age group that was found positive for
558	HEV RNA) for flocks on Farm A (<i>n</i> =11 flocks), Farm B (<i>n</i> =10 flocks), Farm C (<i>n</i> =11
559	flocks), Farm D (n = 8 flocks), Farm E (n = 7 flocks), Farm F (n = 5 flocks) and Farm G (n = 10
560	flocks).
561	
562	Fig. 2. Difference (%) of actual mortality (A), egg production (B), and egg weights (C) in
563	Farms A through G when compared to the breed standards. The box-and-whiskers plots show
564	cumulative results of all flocks within a farm. Different superscripts (a,b) indicate significant
565	(P < 0.05) differences between farms for the measured parameter.
566	
567	Fig. 3. Phylogenetic tree based on 361 bp of the helicase gene (A) and 232 bp of the capsid
568	gene (B) of the avian HEV isolates. Sequences that were obtained in this study are indicated
569	in a black box. The phylogenetic tree was constructed by the NJ method implemented in the
570	Lasergene software (DNASTAR). The nucleotide substitution per 100 residues is given;
571	bootstrap values are indicated for the major nodes. Genotype classification represented as
572	proposed by Marek et al. (2010) and Hsu and Tsai (2014).
573	

74	Table 1				
75	Comparison of anti-avian HEV antibodies detection rates with ELISA and FMIA on sample				
76	obtained	d from 12 experimen	ntally infected chick	ens on different days	post infection. Data
77	presente	ed as number of posi	tive chickens/total r	number of chickens te	sted (mean group
78	antibody	y index ± standard d	eviation).		
	Assay	0	7	14	21
	FMIA	0/12 (0.04±0.07)	0/12 (0.03±0.02)	7/12 (2.62±3.08) ^{A*}	12/12 (6.87±3.17)
	ELISA	0/12 (0.20±0.04)	0/12 (0.23±0.08)	$1/12 (0.41\pm0.32)^{B}$	9/12 (1.89±0.93)
30 31 32	0.05) wi	ithin groups.		anusci)	
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Table 2

Detection rates of anti-avian HEV IgY antibodies in serum samples tested by FMIA and avian HEV RNA in fecal samples tested by RT-PCR from Farms A through G. Data presented as number of positive samples/total number of samples tested (prevalence).

	Anti-avian HEV IgY	Avian HEV RNA
Farm	antibody (%)	(%)
A	29/55 (52.7) ^{B*}	10/11 (90.9) ^B
В	41/50 (82.0) ^C	10/10 (100) ^B
C	18/55 (32.7) ^{AB}	9/11 (81.8) ^B
D	24/40 (60.0) ^{BC}	6/8 (75.0) ^B
Е	8/35 (22.9) ^A	0/7 ^A
F	5/25 (20.0) ^A	0/5 ^A
G	36/50 (72.0) ^{BC}	$4/10 (40.0)^{B}$

*Different superscripts (A,B,C) within a column indicate significant differences in detection rates (P < 0.05) among groups.

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