cerebral edema and a \textit{P. falciparum} parasitemia level of 0.7%. The patient died 1 day later (day 2).

A blood sample obtained from the patient on day 1 in France showed a doxycycline concentration of 195 \textmu g/mL plasma. This concentration, which was determined by liquid chromatography coupled with tandem mass spectrometry, was compatible with a last doxycycline uptake 1 day before diagnosis (day –1). The finding of the expected doxycycline plasma concentration, together with assurances (colleague’s statements and collective intake of doxycycline) that the patient had followed the drug regimen, was sufficient to suggest prophylaxis failure in a treatment-compliant patient.

The \textit{P. falciparum} sample obtained from the patient on arrival in France was evaluated for in vitro susceptibility to doxycycline, but the evaluation was unsuccessful. The number of copies of \textit{PftetQ} and \textit{Pfmdt} genes were evaluated relative to the single-copy \textit{P. falciparum} \textit{b}-tubulin gene (\textit{Pf\textit{\textit{b}-tubulin}}), as previously described (7,8). The sample was assayed in triplicate. The 2$^{-\Delta\Delta C_t}$ method (where \( C_t \) indicates cycle threshold) was used and adapted to estimate the number of copies of \textit{Pfmdt} and \textit{PftetQ} by using the formula \( DDC_t = (C_t(\textit{PftetQ} ~or ~\textit{Pfmdt}) - C_t(\textit{Pf\textit{\textit{b}-tubulin}}))_{\text{Sample}} - (C_t(\textit{PftetQ} ~or ~\textit{Pfmdt}) - C_t(\textit{Pf\textit{\textit{b}-tubulin}}))_{\text{Calibrator}} \).

Genomic DNA extracted from 3D7 \textit{P. falciparum}, which has a single copy of each gene, was used as calibrator sample; \textit{Pf\textit{\textit{b}-tubulin}} served as the control housekeeping gene. The experiment was assayed twice. The sample had 2 copies of \textit{PftetQ} and \textit{Pfmdt} genes, which suggested decreased in vitro susceptibility of the sample to doxycycline (8,9). The genotyping of \textit{PftetQ} sequence polymorphisms was done by using conventional methods with the primers \textit{PfTetQ} forward (5’-TCACGACAAATGTGCTAGATAC-3’) and \textit{PfTetQ} reverse (5’-ATCATCATTTGTGGTGGATAT-3’), as previously described (10). Two \textit{PftetQ} KYNNNN motif repeats were found in the sample; <3 KYNNNN motif repeats are predictive of in vitro \textit{P. falciparum}–resistant parasites with an IC\textsubscript{50} of >35 mM (odds ratio 15) (10). The 2 copies of \textit{Pfmdt} and the 2 KYNNNN motif repeats have been shown to be associated with parasites in vitro resistance to doxycycline (9,10). The association of doxycycline resistance (prophylactic failure with statement of correct intake and the presence of an expected concentration) with increased \textit{Pfmdt} copies and decreased \textit{PftetQ} KYNNNN motif repeats suggest that these molecular markers are predictive markers of doxycycline resistance that can be used for resistance surveillance.

\begin{enumerate}
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\section{Avian Gyrovirus 2 DNA in Fowl from Live Poultry Markets and in Healthy Humans, China}

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To the Editor: In 2011, a chicken anemia virus (CAV)-related sequence, designated avian gyrovirus 2 (AGV2), was first identified in serum samples from diseased chickens in Brazil (1). During the same year, a human gyrovirus (HGyV) sequence that had high identity to AGV2 was detected in the skin of humans in France (2). As with CAV, 3 open reading frames (ORFs) for encoding viral proteins (VP) 1–3 (2) overlapped in genome of AGV2. Recently, HGyV/AGV2 has been detected in Hong Kong in chicken meat for consumption by humans, in human blood samples from donors in France, and in HIV-positive persons and organ transplant recipients in Italy and the United States (3–5). However, the epidemiology, host range, transmission route, and pathogenesis of AGV2 remain poorly understood. Bullenkamp et al. found that AGV2 VP3 protein, like CAV VP3, can induce apoptosis of tumor cells (6). Also, Abolnik et al. reported the detection in Southern Africa of AGV2 in brain tissue of chickens that showed severe neurologic signs (7). These findings highlight the potential pathogenesis of AGV2.

So far, little is known about AGV2 in mainland China among chickens and humans. Because live poultry markets (LPMs) play a critical role in the transmission of poultry pathogens to humans, we used PCR to investigate the presence of AGV2 in chickens (54 feather shaft samples) from 4 LPMs in Yangzhou and in 178 human blood samples from healthy persons living in Yangzhou. The DNA from the feather shafts and human blood were extracted as previously described (8). PCR was performed by using the following 2 primers: AGV2_F 5′-CGTGTCCGCCAGCAGAAACGAC-′3 and AGV2_R 5′-GGTAGAAGCCAAAGCGTCCACGGA-′3. The PCR targets partial VP2 and VP3 genes that have an expected size of 346 bp. The parameters of the PCR were as follows: 1 cycle at 95°C for 5 min; then 30 cycles at 94°C for 30 s, 64°C for 30 s, and 72°C for 30 s; and 1 cycle at 72°C for 10 min. PCR showed that a band with the size of ≈346 bp could be amplified in 10 of 54 chicken feather samples and in 2 of 178 human blood samples.

We confirmed the AGV2 specificity of these PCR-amplified bands by direct sequencing using the Sanger method. The sequence assay showed that the 12 sequences identified here had 98.3%–100% homology to each other and 92.2%–99.1% aa identity to AGV2 samples previously deposited in GenBank (see Figure legend for accession numbers). The positive rates for samples from the 4 LPMs tested were 25%, 12.5%, 15.8%, and 20%; the positive rate for the 178 human blood samples was 1.1%. The low positive frequency of AGV2 in human blood detected in this study is consistent with that found by investigation in other countries (3,4). Because the limit of detection of PCR in this study was estimated to be 2.7 copies of AGV2 DNA using dilutions of a plasmid with partial AGV2 sequence, we determined that the copy number of AGV2 in the 2 positive human blood samples was 2.7 × 10^3 copies/mL plasma.

We also constructed a phylogenetic tree using the neighbor-joining method (1,000 bootstrap replications) with MEGA6 (9). The tree analysis revealed that the 12 AGV2 isolates we identified and 7 AGV2 isolates from GenBank clustered into 2 subgroups on the basis of the PCR amplified fragment (Figure). The 12 AGV2 sequences we identified clustered together with gyrovirus sequences detected in ferret and human blood samples in subgroup I, and the prototype sequence Ave3 was located in subgroup II. The 12 AGV2 showed ≈92.2%–93% aa identity to Ave3, and <99.1% homology with isolates CL33, G13, and 915F06007 detected in ferret and human samples.
AGV2 sequences also showed ≈93%–93.9% identities to ACV2 sequence that was previously identified in human fecal samples from mainland China (GenBank accession no. JQ690763). The China sequence also clustered with Ave3 in subgroup II. These findings indicate that ≥2 subgroups of AGV2 are circulating in mainland China.

Our results demonstrate the presence of AGV2 in LPMs and human blood in mainland China. The amplification and analysis of partial AGV2 sequences was the major limitation in our method. The high homology between sequences identified in LPMs and human blood indicates the LPMs are a potential source for AGV2 in humans. Unlike our 12 conserved AGV2, AGV2 identified by Santos et al. in southern Brazil varied <15.8%, and these variants of AGV2 were mainly detected in diseased chickens (8). However, little is known about the molecular epidemiology of these AGV2 variants in other countries. More recently, Varela et al. reported the detection of AGV2 in poultry vaccines, indicating the potential role of contaminated vaccines in the spread of AGV2 (10). Future studies should investigate the large geographic distribution of AGV2 and monitor the variants, the host range, and the associated diseases.

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