Therefore, the new H7N9 viruses were highly pathogenic to chickens when compared with the early H7N9 virus and could transmit among chickens by contact.

The biological features of H7N9 virus and its pandemic potential have caused global concern (8). The early H7N9 viruses lacked the basic HA cleavage site, exhibited low pathogenicity, and caused mild or no disease in poultry (9). The cleavage site in HA protein of the isolates we analyzed were KGKRTAR’G or KRKRTAR’G. They had high pathogenicity and replication in chickens and could transmit among chickens by contact. Therefore, these new H7N9 viruses could cause a pandemic among poultry and humans in China.

Molecular evolution showed that Q1 was a triple reassortant virus (H5, H7, and H9 subtypes) consisting of Yangtze River Delta A and B lineages of H7N9 and GSGD96 lineage of H5N1. The Q26 and Q39 viruses were both double reassortant avian influenza viruses (H7 and H9 subtype), as was the early H7N9 virus (Figure; online Technical Appendix Table 1, Figure). Therefore, the 3 H7N9 viruses we isolated have 2 kinds of insertions in the cleavage sites and were likely derived from different lineages of H7N9 viruses, or even from different subtypes that were co-circulating in southern China during 2016–2017.

Acknowledgments
We thank the Key Laboratory of Zoonosis, Ministry of Agriculture; the National and Regional Joint Engineering Laboratory for Medicament of Zoonosis Prevention and Control; the Key Laboratory of Animal Vaccine Development, Ministry of Agriculture; and the Key Laboratory of Zoonoses Control and Prevention of Guangdong.

This work was supported by grants from the National Key Research and Development Program of China (2016YFD0500207), the National Natural Science Foundation of China (U1501212), the Natural Science Foundation of Guangdong Province (2016A030308001), and Basic Research (Discipline Layout) of Shenzhen (JCYJ20160323163102764).

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Rabies and Distemper Outbreaks in Smallest Ethiopian Wolf Population
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DOI: https://doi.org/10.3201/eid2312.170893
Widespread deaths recently devastated the smallest known population of Ethiopian wolves. Of 7 carcasses found, all 3 tested were positive for rabies. Two wolves were subsequently vaccinated for rabies; 1 of these later died from canine distemper. Only 2 of a known population of 13 wolves survived.

Canine diseases pose a growing threat to wildlife species of conservation concern worldwide. Although extensive oral vaccinations have eliminated rabies virus (RABV) from wild carnivore populations in some developed countries (1), elsewhere, the challenges to controlling diseases in endangered wildlife are many and persistent. Massive outbreaks of rabies and, more recently, canine distemper have repeatedly decimated populations of Ethiopian wolves (Canis simensis) in the Bale Mountains of Ethiopia, where more than half of a global population of ≥500 wolves live (2,3). Extensive efforts to control RABV in the reservoir population of sympatric domestic dogs have proved insufficient. Therefore, reactive vaccination of Ethiopian wolves, carried out in response to an outbreak in wolves, has been the primary mechanism to curtail mortality in the affected wolf populations in the Bale Mountains (4).

The fragile status of the Bale population highlights the conservation value of the other remaining, much smaller, wolf populations scattered throughout the highlands of Ethiopia. Models predict these small populations to be particularly vulnerable to disease outbreaks (5); however, no outbreaks had been detected outside Bale, either because they went unnoticed, because in small populations outbreaks die out before causing a major epizootic event, or both. We report consecutive rabies and canine distemper outbreaks among Ethiopian wolves in Delanta, in the Wollo highlands.

This group of wolves is the smallest extant wolf population; 13 wolves in 3 family packs lived in the remaining 20 km² of Afroalpine habitat in 2015. The first wolf carcass was detected in late June 2016; by early September, 7 deaths had been confirmed. RABV infection was identified as the cause of death in all 3 of the carcasses tested, as well as in samples from 1 domestic dog concurrently found dead within wolf habitat (Table). A vaccination intervention was initiated in September 2016, when only 3 wolves were known to be alive; 1 adult male (>2 years of age) and 1 subadult female (1–2 years of age) were trapped (7) and parenterally inoculated with Nobivac Rabies (Merck Animal Health, Madison, NJ, USA) (4). In December 2016, the female wolf was found dead and tested positive for canine distemper virus (CDV) (Table); CDV was also detected in a dog carcass found concurrently in the vicinity of the wolf range. In late May, the vaccinated male was still alive and was observed until at least April 2017 with an unknown adult female.

Evidence indicates a first outbreak of rabies, overlapping or followed soon after by a canine distemper outbreak. Confirmation of disease in contemporarily recovered dog carcasses is consistent with a pattern of transmission from reservoir domestic dogs to their wild relatives (as observed in the Bale Mountains [8]), with disastrous consequences for the small Delanta population, which harbored <20 wolves before the epizootic events. Although the larger Bale wolf population has recovered from epizootic events in the past (2,9), smaller populations are expected to be less resilient, a factor exacerbated by their virtual isolation from other wolf populations. Modeling has predicted a high extinction risk if Ethiopian wolf populations are affected by consecutive epizootic events over a short period of time (5). The combined exacerbated effects of RABV and CDV infection were first described in 2010 in the Bale Mountains (3).

Although the loss of Afroalpine habitats is bound to determine the fate of Ethiopian wolf populations (2 extinctions were recorded in areas of a similar size to that of Delaware during 1999 and 2010) (2), incursions of infectious diseases can drive local extinctions. Preemptive vaccination, in combination with actions to protect the habitat of this specialized predator, could greatly reduce the risk of populations becoming extinct, even if a relatively low proportion of the wolves is successfully vaccinated (4). Recently, SAG2, an oral rabies vaccine, was successfully tested in Ethiopian wolves (10), and a CDV parenteral vaccination trial is ongoing, with positive

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**Table.** Characteristics of and test results for Ethiopian wolf and domestic dog carcasses recovered in Delanta, Ethiopia, 2016*

<table>
<thead>
<tr>
<th>Date found</th>
<th>Species</th>
<th>Age and sex</th>
<th>Postmortem</th>
<th>Tested for RABV†</th>
<th>Tested for CDV‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jun 26</td>
<td>Ethiopian wolf</td>
<td>Juvenile female</td>
<td>No</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Jun 28</td>
<td>Ethiopian wolf</td>
<td>Juvenile male</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Jul 07</td>
<td>Ethiopian wolf</td>
<td>Adult male</td>
<td>Yes</td>
<td>Positive</td>
<td>No</td>
</tr>
<tr>
<td>Jul 11</td>
<td>Ethiopian wolf</td>
<td>Juvenile female</td>
<td>Yes</td>
<td>Positive</td>
<td>No</td>
</tr>
<tr>
<td>Jul 18</td>
<td>Ethiopian wolf</td>
<td>Juvenile female</td>
<td>No</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Aug 12</td>
<td>Ethiopian wolf</td>
<td>Adult male</td>
<td>No</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Sep 01</td>
<td>Domestic dog</td>
<td>Adult male</td>
<td>Yes</td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Sep 07</td>
<td>Ethiopian wolf</td>
<td>Adult female</td>
<td>Yes</td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Nov 27</td>
<td>Domestic dog</td>
<td>Adult male</td>
<td>Yes</td>
<td>Negative</td>
<td>Positive</td>
</tr>
<tr>
<td>Dec 27</td>
<td>Ethiopian wolf</td>
<td>Adult female</td>
<td>Yes</td>
<td>Negative</td>
<td>Positive</td>
</tr>
</tbody>
</table>

*CDV, canine distemper virus; NA, not applicable; RABV, rabies virus.
†Rabies diagnostic reverse transcription PCR was performed as described previously (6).
‡CDV diagnostic reverse transcription PCR was performed as described previously (3).
preliminary results. We propose proactive vaccination of Ethiopian wolves across their distribution as an effective and urgently needed strategy to protect the species from extinction. This program should be part of an integrated disease control plan that also includes controlling disease in domestic dogs, limiting contact between dogs and wolves, and conducting policy and education interventions to reduce the size and roaming behavior of local dog populations (2).

Acknowledgments
We thank the Ethiopian Wildlife Conservation Authority and Environment and Forest and Wildlife Protection and Development Authority (Amhara National Regional State) for support and permission to work in Delanta. We thank Delanta and Gubalaftu Waredas and Wolf Ambassadors, and Leigh Thorne and Daisy Jennings for excellent technical assistance.

The work was funded by the Born Free Foundation and the Wildlife Conservation Network. The work undertaken by Animal and Plant Health Agency is funded by a grant (SEV3500) from the UK Department for Environment, Food and Rural Affairs, Scottish and Welsh Governments.

The animal care and use protocols for the ethical handling of Ethiopian wolves were approved by the Ethiopian Wildlife Conservation Authority and the University of Oxford’s Local Ethical Review Process (Zoology ERC; case no. ZERC040905) and adhere to the United Kingdom’s ASPA regulations (1986).

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High Abundance and Genetic Variability of Atypical Porcine Pestivirus in Pigs from Europe and Asia

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