Using MEGA 7.0 software (http://www.megasoftware.net) with the maximum-likelihood algorithm and a bootstrap of 1,000 replicates, we constructed a phylogenetic tree (online Technical Appendix Figure). All 8 sequences clustered within subtype 4d of HEV. The sequences were similar to each other (95.5%–99.8% similarity in nucleotide sequence) and similar to sequences reported for other cattle (83.3%–85.3%; online Technical Appendix Figure). Moreover, these sequences shared 96.1%–96.6% similarity with a human HEV strain (GenBank accession no. KC163335) from the Yantai Prefecture in 2012 and 95.7%–97.9% similarity with a swine strain (GenBank accession no. KF176351) isolated in Shandong Province the same year.

Our data strongly indicate that HEV infection occurs in yellow cattle and that they could also play a role as a reservoir of HEV. Because these animals serve mainly as a source of food, consumption of undercooked meat from yellow cattle, similar to pork, might also contribute to the transmission of HEV to humans. Additionally, we also detected HEV RNA in 8 of 70 sheep (online Technical Appendix Table 2). Eight sequences from yellow cattle had 95.1%–99.8% nt homology with 8 sheep-derived HEV strains, possibly because mixed raising of domestic livestock is popular in this region. Our finding of high sequence similarity between yellow cattle, sheep, swine, and human populations suggests a complicated interspecies transmission of HEV occurred in this province. Further studies are required to evaluate the contribution of the yellow cattle reservoir to human HEV infection.

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Introgressed Animal Schistosomes Schistosoma curassoni and S. bovis Naturally Infecting Humans

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To the Editor: Schistosomiasis, a disease caused by infection with parasitic worms (schistosomes), is a neglected tropical disease across many parts of the world. Numbers of infected livestock are unknown, but >250 million persons are infected; the greatest number of cases are in sub-Saharan Africa (1). Schistosome eggs are excreted through urine or feces, depending on the species, and hatch into miracidia upon contact with freshwater. Larvae are transmitted to the mammalian host indirectly through a molluscan intermediate host. Goals to eliminate schistosomiasis by 2020 in select countries in Africa have
been proposed by the World Health Organization (http://www.who.int/neglected_diseases/NTD_RoadMap_2012_Fullversion.pdf) and a coalition of partners combating neglected tropical diseases (2).

Selective pressures imposed by natural phenomena and human activities affect the dynamics and distribution of schistosomiasis. For example, ongoing changes in agricultural practices in some regions are creating new water bodies shared by humans and domestic livestock. These anthropogenic changes increase opportunities for mixing of and subsequent exposure to different Schistosoma species, especially those that infect humans and livestock. Such mixing could increase the potential for novel zoonotic hybrid parasites to emerge and become established (3,4).

Since the 1940s, researchers have suspected that natural hybridization within and between human and animal schistosome species occurs in definitive and intermediate hosts. In western Africa, hybrids, predominantly between S. haematobium (human schistosome) and S. bovis or S. curassoni (livestock schistosomes), have been isolated from children and snails (5,6). Hybrids between these livestock-only Schistosoma species have also been reported in cattle and sheep but not in other hosts (7). As demonstrated in the field and under experimental laboratory conditions, neither S. bovis nor S. curassoni, as single species, can fully develop in humans or nonhuman primates (8,9).

We report evidence of a child in Niger who was infected by livestock-specific schistosomes through the hybridization and introgression of S. bovis with S. curassoni. Samples were collected from a 10-year-old girl in Kokourou, Tillaberi region, Niger (14°20′61.50″N, 0°91′94.0″E), as part of longitudinal monitoring and evaluation of national disease control interventions in the area. Tillaberi has an ongoing high prevalence and infection intensity of schistosomiasis, despite more than a decade of high-coverage mass treatment with praziquantel.

Schistosoma eggs were recovered from the child’s urine. Miracidia that hatched from the eggs were stored on a Whatman FTA card (Sigma-Aldrich, St. Louis, MO, USA), providing the opportunity to perform noninvasive molecular characterization on schistosome larvae without any necessity for laboratory passage. Multilocus analyses of mitochondrial and nuclear DNA regions (6) on all 42 larvae collected from the child identified 2 individual miracidia with a livestock S. bovis × S. curassoni hybrid profile with no genetic signal from human-specific S. haematobium. Although no species-specific markers exist for determining parental lineages, internal transcribed spacer (ITS), a powerful marker for detecting introgression, has been used successfully to detect hybridization events across the Schistosoma genus. A partial mitochondrial cox1 sequence for miracidia was identical to that for S. bovis, and the nuclear ITS and partial 18S rDNA sequences were identified as S. curassoni.

One miracidium from the child had a pure S. haematobium profile, 13 had S. bovis (cox1) × S. haematobium (ITS) profiles, and 5 had S. bovis (cox1) × S. haematobium × S. curassoni (ITS) profiles, suggesting the potential for repeated interactions and cross-pairings among these 3 species. No S. curassoni mitochondrial DNA was found in miracidia. The molecular data suggest that these hybrids were not first generation, but a result of parental and/or hybrid backcrosses. The data confirm the occurrence of bidirectional introgression between Schistosoma species that infect livestock and those that infect humans in Niger. Our data also show that hybrid livestock:livestock schistosomes can directly infect humans without combining with a Schistosoma species that infects humans.

Hybridization of parasites is an emerging public health concern at the interface of infectious disease biology and evolution (3). Our results raise several critical questions regarding the evolution, epidemiology, health effects, and ultimate control of schistosomiasis. Hybrid schistosomes, and, in particular, hybridized livestock:livestock schistosomes infecting humans, could potentially extend intermediate and definitive host ranges; confer altered infectivity, virulence, and drug efficacy; or even potentially replace existing single species (3,4). Hybrid vigor has been observed experimentally for schistosomes, and similar evidence is gathering from experiments with other parasites. Our results strongly indicate that hybrid livestock:livestock schistosomes can infect a human definitive host, even when neither of its single parental livestock species appears compatible. Such novel livestock schistosome hybrids infecting humans could be predicted to spread to wherever suitable intermediate snail hosts are endemic, as has recently been reported for zoonotic Schistosoma hybrids in Corsica (10).

It is imperative to identify and understand the transmission dynamics of introgressed Schistosoma species combinations. The detection of multiple introgressed hybrids with mixed ancestry in a single child suggests that Schistosoma species may be adapting to recent anthropogenic changes. If novel zoonotic hybrid species are playing a role in maintaining and exacerbating schistosome transmission, illness caused by infection, or both, treatments for humans and livestock may have to be adjusted accordingly within a One Health framework (4).

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Anaplasma phagocytophilum

The data used in this study were collected as part of the monitoring and evaluation processes of the Schistosomiasis Control Initiative programs taking place in Niger. Sequences were obtained using the DNA sequencing facilities in the Natural History Museum. Ethical approval for this research was granted by the Niger Ministry of Health Ethical Review Board and by the Imperial College Research Ethics Committee (ICREC_8_2_2, EC no. 03.36, R&D no. 241 03/SB/003E) in combination with ongoing Schistosomiasis Control Initiative activities. All infected children in the study were provided treatment with 40 mg/kg praziquantel.

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Rickettsia raoultii in Dermacentor reticulatus
Ticks, Chernobyl Exclusion Zone, Ukraine, 2010

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To the Editor: The Chernobyl Exclusion Zone (CEZ) surrounds the center of the 1986 Chernobyl nuclear power plant disaster. Preliminary study shows predominance of Dermacentor reticulatus ticks in the CEZ; ticks of other species, such as Ixodes ricinus, are surprisingly rare, even in habitats where they should be relatively common (1). A few reports document presence of Ix. trianguliceps ticks (2,3). Prevalence of pathogens (Anaplasma phagocytophilum, Borrelia burgdorferi s.l., Babesia spp.) in these ticks is higher in the CEZ than in other regions (3,4). One pathogen transmitted by Dermacentor spp. ticks is Rickettsia raoultii, which has been isolated from species of Dermacentor ticks found in Asia (5,6) and since 1999 has also been detected in Europe.

In our study, D. reticulatus ticks were collected by use of the flagging method (1) in the CEZ in September 2010. Ticks were collected from areas where they were known to occur, around the former villages of Korohod (51°16′02′′N; 30°01′04″E) and Cherevach (51°12′44″N; 30°07′45″E) and around Chernobyl city (51°17′04″N; 30°13′25″E). The habitats investigated included open areas and the remnants of farmlands. A total of 201 D. reticulatus ticks, 87 males and 114 females, were collected and investigated (Table).

DNA was extracted by use of the ammonium hydroxide method (7). Isolated DNA was examined for the presence of the Rickettsia sp. citrate synthase gene (gltA) by use of PCR with RpCS.409d and RpCS.1258n primers (8). Positive amplicons were sequenced, and sequences were edited by using AutoAssembler software (Applied Biosystems, Foster City, CA, USA) and compared with GenBank entries by using blastn version 2.2.13 (http://www.ncbi.nlm.nih.gov/blast/download.shtml). All obtained sequences were submitted to GenBank (accession nos. KX056493 and KX056494).