Diversity of Parvovirus 4–like Viruses in Humans, Chimpanzees, and Monkeys in Hunter–Prey Relationships

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During 2010–2011, we investigated interspecies transmission of partetetraviruses between predators (humans and chimpanzees) and their prey (colobus monkeys) in Côte d’Ivoire. Despite widespread infection in all species investigated, no interspecies transmission could be detected by PCR and genome analysis. All sequences identified formed species- or subspecies (chimpanzee)-specific clusters, which supports a co-evolution hypothesis.

Since 2005, new paroviruses have been discovered in the following groups: humans (parovirus 4 [PARV4]), bats (Eidolon helvum parovirus 1), and other mammals (cows, pigs, wild boars, and sheep; Hong Kong virus) (1–5). Phylogenetic analysis suggests that these paroviruses form a separate novel genus, with the proposed name of Partetetravirus, within the subfamily Parvovirinae. Globally, 3 genotypes of PARV4 have been found to infect humans (6,7). Recently, PARV4-like viruses have also been described in chimpanzees and gorillas (8). Researchers have suggested that partetetraviruses have co-diverged with their hosts during mammalian evolution. Strains described so far have shown restricted sequence diversity within their host-specific clusters. However, the highly restricted sequence diversity of circulating variants of PARV4 also suggests that the virus has emerged and spread in the human population relatively recently. To clarify whether interspecies transmission is possible for primate PARV4-like viruses, as has been shown for other paroviruses (9), we investigated samples in a setting where transmission of certain simian viruses between these species has been documented (10,11). We analyzed samples from wild chimpanzees (Pan troglodytes verus) in the Taï National Park, Côte d’Ivoire; their prey, red colobus monkeys (Piliocolobus badius) and black-and-white colobus monkeys (Colobus polykomos); and humans who hunt colobus monkeys in the same region.

The Study

Chimpanzee and monkey samples were obtained as described from 2002–2007 (10,11). Human volunteers, recruited during a broad study of primate-borne zoonoses (samples collected between June 2006 and January 2007), ranged in age from 7 to 95 years, and all lived adjacent to the primate habitat. All human participants acknowledged eating bushmeat; most (74%) also reported butchering animals, and a small group (8%) admitted hunting bushmeat. Written informed consent forms were signed by all participants. Ethical approval was obtained from the Institut Pasteur Côte d’Ivoire and the Ministère de la Santé of Côte d’Ivoire. Cross-contamination of samples was avoided by using disposal materials and maintaining a strict safety regime for sampling humans and animals. Samples from different species were handled separately throughout the process, from sampling to analysis.

DNA from 17 chimpanzees (lung, spleen, or liver), 30 red colobus monkeys (buffy coat, blood, bone marrow, intestine, spleen, liver, lung, or muscle), and 15 black-and-white colobus monkeys (buffy coat, liver, intestine, or muscle) and 700 humans (blood), was prepared by using commercial DNA extraction kits (QIAGEN, Hilden, Germany). Partetetravirus generic quantitative real-time PCR (qPCR) was used to screen the samples as described (1). Viral DNA from positive samples was amplified by using primers for conserved regions spanning human variants and other partetetraviruses described in pigs and cows (2).

PARV4-like viruses were detected by qPCR in 7 (41%) of 17 chimpanzees tested. Seven (23%) of 30 red colobus and 2 (13%) of 15 black-and-white colobus monkeys were positive for PARV4-like viruses. The availability of only 1 sample per individual animal limited the analysis of viral tissue distribution (Table).

In a recent study of 91 Old World monkeys tested, none exhibited PARV4-like virus seroreactivity, whereas 63% of chimpanzees (P.t. troglodytes) and 18% of

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DISPATCHES

| Sample origin                        | Blood | Bone marrow | Intestine | Spleen | Muscle | Liver | Lung | Total |
|--------------------------------------|-------|-------------|-----------|--------|--------|-------|------|-------|
| Red colobus monkey (Piliocolobus badius) | 1/14 (7)† | 2/6 (33) | 0/1 | 0/3 | 2/2 (100) | 1/2 (50) | 1/2 (50) | 7/30 (23) |
| Black and white colobus monkey (Colobus polykomos) | 2/10 (20)‡ | – | 0/1 | – | 0/3 | 0/1 | – | 2/15 (13) |
| Wild chimpanzee (Pan troglodytes verus) | 0/1‡ | – | – | 5/13 (38) | – | – | 2/3 (67) | 7/17 (41) |
| Human§ | 12/700 (2) | – | – | – | – | – | – | 12/700 (2) |

*Each sample represents 1 animal or person; PARV4, parvovirus 4; –, no sample.
†Blood or buffy coat.
‡Buffy coat.
§Dried blood spot on filter paper.

| Sample material, no. PCR positive/no. tested (%) |
|-----------------------------------------------|
| Blood Bone marrow Intestine Spleen Muscle Liver Lung Total |
| Red colobus monkey (Piliocolobus badius) | 1/14 (7)† | 2/6 (33) | 0/1 | 0/3 | 2/2 (100) | 1/2 (50) | 1/2 (50) | 7/30 (23) |
| Black and white colobus monkey (Colobus polykomos) | 2/10 (20)‡ | – | 0/1 | – | 0/3 | 0/1 | – | 2/15 (13) |
| Wild chimpanzee (Pan troglodytes verus) | 0/1‡ | – | – | 5/13 (38) | – | – | 2/3 (67) | 7/17 (41) |
| Human§ | 12/700 (2) | – | – | – | – | – | – | 12/700 (2) |

In 12 (1.7%) of the 700 human blood samples, PARV4 genotype 3 was amplified. The median age of infected persons was 11 years (mean 20.8 years, range 7–75 years). As discussed previously (12), the broad age range suggests that PARV4 genotype 3 in Africa has alternative routes of infection from genotypes 1 and 2 that are found in Europe. Near full-length nucleotide sequences from viral genomes were generated from samples from 4 persons (GenBank accession nos. JN798193–196, 1 chimpanzee (JN798203), and 1 black-and-white colobus monkey (JN798211). Partial sequences were derived from 7 persons (JN798192, JN798197–201), 3 chimpanzees (JN798204–206), and 4 red colobus monkeys (JN798207–210).

On the basis of phylogenetic analyses of the 20 sequences obtained, we could not detect any interspecies transmission; all sequences formed host species–specific clusters (Figure). Sequence diversity of genotype 3 viruses from humans in the defined region in our study was 0.9% over 1,423 nt positions, which is in accordance with what has been published (12). One additional near full-length genotype 3 sequence from Africa was included in the phylogenetic analysis; it clustered with human sequences found in this study. Compared with sequences from human PARV4 (AY622943) isolates, sequences from isolates from chimpanzees differed by 19% over 4,771 nt and formed a distinct cluster, together with sequences derived from chimpanzees of other subspecies. PARV4-like virus from P.t. verus (JN798203) from this study and P.t. troglodytes (HQ113143) from Cameroon (8) differed by 10%, suggesting subspecies-specific PARV4-like viruses within chimpanzees of 2 different subspecies from distinct habitats. Black-and-white colobus monkey viruses exhibited a closer relationship to viruses from humans than did PARV4-like viruses from red colobus monkeys. This finding corresponds to a previous phylogenetic analysis in which the genomic relationship of different species showed that colobus monkeys separated earlier from piliocolobus monkeys during evolution (13). The distance between PARV4-like viruses from piliocolobus monkeys to those of colobus monkeys supports these data.

To evaluate whether underlying infections were present in the hunters (chimpanzees and humans) originating from the prey, a red colobus PARV4-like virus–specific qPCR was designed. Samples from humans, chimpanzees, and black-and-white colobus monkeys, which previously tested positive in the generic partetravirus qPCR, were retested. Although chimpanzees consumed immense quantities of red colobus meat and organs (~45 kg/year for adult males [4]), we could not detect red colobus PARV4-like virus in any of the 17 chimpanzees analyzed. Similarly, no red colobus PARV4-like virus DNA was discovered in humans, although in this particular region the red colobus monkey is the most hunted and eaten primate (15). These results support the hypothesis that PARV4-like viruses are species specific, notwithstanding constant high exposure to infectious materials.

Conclusions

We demonstrated that partetravirus infection is widespread in monkeys, chimpanzees, and humans in West Africa. However, the PARV4-like viruses seem to remain species specific, despite continuous opportunities for interspecies transmission. The data presented here suggest that the risk for zoonotic transmission of PARV4-
like viruses from nonhuman primates in West Africa is low. Nonetheless, parvovirus evolution has pointed toward occasional cross-species transmissions (9). This observation, coupled with the frequent intimate contact between bushmeat hunters and their prey, compels the continued vigilance for cross-species transmission of these viruses and others with the intention of mitigating the risk posed by novel introductions of viral zoonoses.

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GenBank accession numbers of the generated sequences: human PARV4, JN798192–JN798202; P. t. verus PARV4-like, JN798203–JN798206; P. badius PARV4-like, JN798207–JN798210; C. polykomos PARV4-like, JN798211.

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References

1. Adlhoch C, Kaiser M, Ellerbrok H, Pauli G. High prevalence of porcine Hokovirus in German wild boar populations. Virol J. 2010;7:171. http://dx.doi.org/10.1186/1743-422X-7-171
2. Lau SK, Woo PC, Tse H, Fu CT, Au WK, Chen XC, et al. Identification of novel porcine and bovine parvoviruses closely related to human parvovirus 4. J Gen Virol. 2008;89:1840–8. http://dx.doi.org/10.1099/vir.0.2008/000380-0
3. Jones MS, Kapoor A, Lukashov VV, Simmonds P, Hecht F, Delwart E. New DNA viruses identified in patients with acute viral infection syndrome. J Virol. 2005;79:8230–6. http://dx.doi.org/10.1128/JVI.79.13.8230-8236.2005
4. Canuti M, Eis-Huebinger AM, Deijs M, de Vries M, Drexler JF, Oppong SK, et al. Two novel parvoviruses in frugivorous new and old world bats. PLoS ONE. 2011;6:e29140. http://dx.doi.org/10.1371/journal.pone.0029140
5. Tse H, Tsoi HW, Teng JL, Chen XC, Liu H, Zhou B, et al. Discovery and genomic characterization of a novel ovine partetravirus and a new genotype of bovine partetravirus. PLoS ONE. 2011;6:e25619. http://dx.doi.org/10.1371/journal.pone.0025619
6. Fryer JF, Delwart E, Bernardin F, Luke PW, Lukashov VV, Baylis SA. Analysis of two human parvovirus PARV4 genotypes identified in human plasma for fractionation. J Gen Virol. 2007;88:2162–7. http://dx.doi.org/10.1099/vir.0.82620-0
7. Simmonds P, Douglas J, Bestetti G, Longhi E, Antinori S, Parravicini C, et al. A third genotype of the human parvovirus PARV4 in sub-Saharan Africa. J Gen Virol. 2008;89:2299–302. http://dx.doi.org/10.1099/vir.0.2008/001180-0
8. Sharp CP, LeBreton M, Kantola K, Nana A, Díjof J, Djoco CF, et al. Widespread infection with homologues of human parvoviruses B19, PARV4, and human bocavirus of chimpanzees and gorillas in the wild. J Virol. 2010;84:10289–96. http://dx.doi.org/10.1128/JVI.01304-10
9. Shackelton LA, Parrish CR, Truyen U, Holmes EC. High rate of viral evolution associated with the emergence of carnivore parvovirus. Proc Natl Acad Sci U S A. 2005;102:379–84. http://dx.doi.org/10.1073/pnas.0406765102
10. Leendertz FH, Junglen S, Boesch C, Formenty P, Couracey-Hymann E, Courgnaud V, et al. High variety of different simian T-cell leukemia virus type 1 strains in chimpanzees (Pan troglodytes verus) of the Tai National Park, Côte d’Ivoire. J Virol. 2004;78:4352–6. http://dx.doi.org/10.1128/JVI.78.8.4352-4356.2004
11. Leendertz FH, Zirkel F, Couracey-Hymann E, Ellerbrok H, Morozov VA, Pauli G, et al. Interspecies transmission of simian foamy virus in a natural predator–prey system. J Virol. 2008;82:7741–4. http://dx.doi.org/10.1128/JVI.00549-08
12. Panning M, Kobbe R, Vollbach S, Drexler JF, Adjei S, Adjei O, et al. Novel human parvovirus 4 genotype 3 in infants, Ghana. Emerg Infect Dis. 2010;16:1143–6. http://dx.doi.org/10.3201/ eid1607.100025
13. Roos C, Zinner D, Kubatko LS, Schwarz C, Yang M, Meyer D, et al. Nuclear versus mitochondrial DNA: evidence for hybridization in colobine monkeys. BMC Evol Biol. 2011;11:77. http://dx.doi.org/10.1186/1471-2148-11-77
14. Boesch C, Boesch H. Hunting behavior of wild chimpanzees in the Tai National Park. Am J Phys Anthropol. 1989;78:547–73. http://dx.doi.org/10.1002/ajpa.1330780410
15. Refisch J, Koné I. Impact of commercial hunting on monkey populations in the Tai region, Côte d’Ivoire. Biotropica. 2005;37:136–44. http://dx.doi.org/10.1111/j.1744-7429.2005.03174.x

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