

Puzzling Origins of the Ebola Outbreak in the Democratic Republic of the Congo, 2014

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Maganga et al. (1) and Naccache et al. (GenBank numbers [KP271018](#), [KP271019](#), and [KP271020](#)) recently reported the genome sequences of the Zaire Ebolavirus (ZEBOV) that caused the 2014 outbreak in the Democratic Republic of the Congo (COD). In contrast to the virus sampled from the ongoing outbreak in West Africa, the sequences from COD (COD/2014/Boende-Lokolia, COD/2014/Lomela-Lokolia16, COD/2014/Lomela-Lokolia17, and COD/2014/Lomela-Lokolia19) are phylogenetically close to the ZEBOVs isolated during the 1995-1996 outbreaks in equatorial Africa (Fig. 1A). Importantly, however, such sequence similarity is far greater than expected given the tree topology and higher rate of ZEBOV evolution determined previously (2, 3). In particular, regression analyses of root-to-tip genetic distances against time of sampling show that the COD sequences deviate from the generally clock-like evolution of other ZEBOVs (Fig. 1B to D). In particular, the COD viruses sampled in 2014 are no further distant from the root of the tree than those from 1994 to 1996, indicating that they have evolved at a lower rate than other ZEBOVs.

There are several possibilities to account for this markedly slower evolution. Anomalously low evolutionary rates have previously been shown to be artifacts resulting from the inadvertent release of viruses archived in laboratories, with the strain of human influenza A virus that led to the 1977 epidemic as a high-profile example (4). However, such an explanation is clearly implausible in this case, particularly because the COD outbreak occurred in a remote geographic region. More likely is that this lineage of ZEBOV was maintained in an animal host population characterized by a lower replication rate and hence fewer opportunities for mutation, such that the ecology of Ebola is more com-

plex than generally envisaged. Previous PCR and serological studies have suggested that various species of fruit bat are the natural hosts of ZEBOV, with some nonhuman primates potentially acting as intermediate hosts (5-7). It is therefore possible that the COD/2014 lineage is maintained endemically in a different bat species with a lower population that roosts at lower densities or in as-yet-unknown animal hosts. Finally, it is possible that the COD/2014 lineage replicates and/or mutates at an intrinsically lower rate, which could be conferred by those mutations fixed prior to its emergence (Fig. 1A, smaller box). Further studies of virus genetics and growth kinetics could provide insights into the viability of this hypothesis. Contamination during genome sequencing, which could also generate such data, is unlikely, because similar viral sequences were produced in two different laboratories (1) (GenBank numbers [KP271018](#), [KP271019](#), and [KP271020](#)). Given the persisting health threat posed by ZEBOVs, revealing the exact origins of this outbreak clearly merits rigorous investigation.

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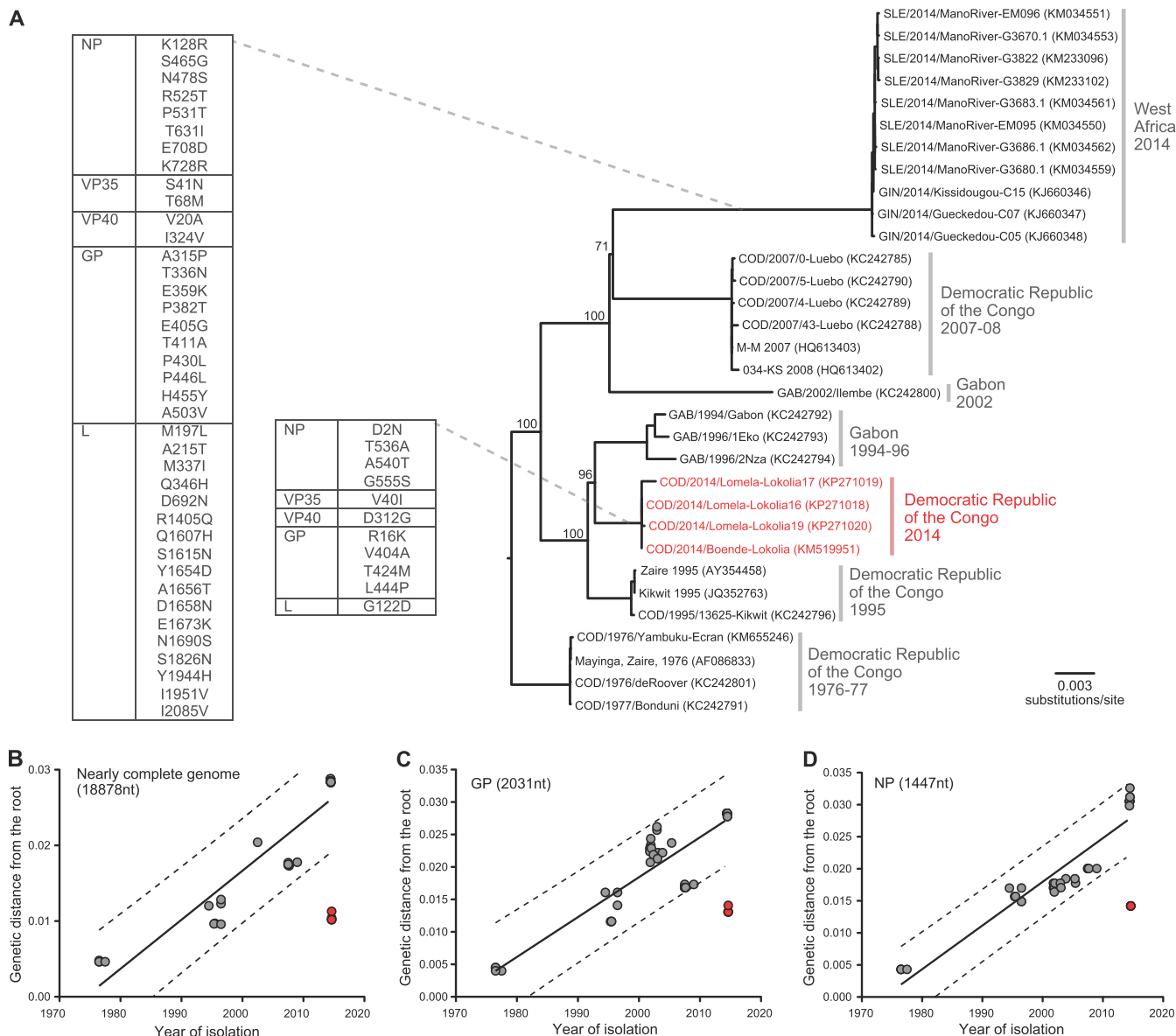


FIG 1 Phylogeny and regression of root-to-tip genetic distances against time of sampling for ZEBOV 1976–2014. (A) Maximum-likelihood phylogeny estimated from the nearly complete genome sequences of 32 ZEBOVs, including those (COD/2014/Boende-Lokolia, COD/2014/Lomela-Lokolia16, COD/2014/Lomela-Lokolia17, and COD/2014/Lomela-Lokolia19; strain names in red) collected from the 2014 outbreak in the Democratic Republic of the Congo (COD). Branch supports (shown at selected nodes) were assessed by bootstrap analysis (1,000 replicates). The boxes on the left show amino acid changes in the NP, VP35, VP40, and GP proteins that have occurred on specific branches, as indicated by dashed lines. Positions are numbered from the first residue of each protein. (B) The genetic distances from the root of the tree to each sequence (tip) are plotted against their year of sampling. Solid and dashed lines are the linear regression fits and the 95% prediction bands for the data points excluding COD/2014 sequences. The COD/2014 ZEBOV sequences (red dots) fall below the expected genetic distance, indicating that they have evolved markedly more slowly. (C and D) Similar observations made using the GP and NP genes, which include more sequences with different time points.

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