Hepatitis B virus genotype E detected in Brazil in an African patient who is a frequent traveler

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Abstract

Genotype E of hepatitis B virus (HBV) has not been described in Brazil and is found mainly in Africa. Genotype A is the most prevalent in Brazil, and genotypes B, C, D, and F have already been reported. We report here an HBV genotype E-infected patient and some characterization of surface (S) protein, DNA polymerase (P) and precore/core (preC/C) coding regions based on the viral genome. The patient is a 31-year-old black man with chronic hepatitis B who was born and raised in Angola. He has been followed by a hepatologist in São Paulo, Brazil, since November 2003, and he is a frequent traveler to Latin America, Africa, and Europe. In 2003, he was diagnosed with HBV infection and started treatment with lamivudine with the later addition of adefovir dipivoxil. No known risk factor was identified. Serologically, he is HBsAg and anti-HBe positive, but HBeAg and anti-HBs negative. DNA sequence analysis of the S/P region confirmed that this patient is infected with genotype E, subtype ayw4. The preC/C region showed G1896A and G1899A mutations but no mutations in the basal core promoter. Nucleotide substitutions common in genotype E were also observed (C1772, T1858 and A1757). Although this is not an autochthonous case and there is no evidence of further spread, the description of this case in Brazil highlights the current risk of viral genotypes spreading with unprecedented speed due to constant travel around the world.

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Key words
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Hepatitis B virus (HBV) is a major cause of chronic liver disease worldwide. This virus is a small DNA virus with a partially double-stranded 3.2-kb genome (1) and has been classified into eight genotypes (A-H) based on sequence divergence over the entire genome. The genotypes show a distinct and well-defined geographic distribution (2). Genotype E of HBV has not been described in Brazil and is found mainly in Africa. Genotype A is the most prevalent in Brazil, and genotypes B, C, D, and F have already been reported.

In the present study, we describe the clinical and epidemiological data of an African patient infected with HBV genotype E which...
A 31-year-old black man with a history of chronic hepatitis B infection has been followed by a hepatologist in São Paulo, Brazil, since November 2003, when he was diagnosed with HBV infection. This patient was born and raised in Angola and, as a lawyer, he frequently travels to Latin America, Africa, and Europe. Initial treatment consisted of lamivudine but, due to low compliance, adefovir dipivoxil was added. The patient suffers from morbid obesity and systemic arterial hypertension. No known risk factor was identified. Currently, he shows normal serum levels of hepatic enzymes (alanine aminotransferase, aspartate aminotransferase, alkaline phosphatase, and \( \gamma \) glutamyltransferase). Serologically, he is HBsAg and anti-HBe positive, but HBeAg and anti-HBs negative. Viral load determined by real time PCR was 291 IU/mL or 3,832.66 copies/mL in August 2006, when the consulting physician also asked for genotype and antiviral resistance tests.

HBV DNA was extracted from 200-µL plasma samples using the QIAamp DNA mini-kit (QIAGEN, Hilden, Germany) according to manufacturer instructions. HBV DNA was detected by nested PCR amplifying the overlapping surface antigen (S) and DNA polymerase (P) genes, as previously described (3,4). For the detection of precore/core mutants, HBV DNA was amplified using the primers described by Takahashi et al. (5). HBV genotyping was carried out by direct sequencing of the amplified product using an ABI3130 DNA sequencer analyzer and the SeqScape Software V2.5 (Applied Biosystems, Foster City, CA, USA), which compares the sequence with a database of sequences from the different genotypes, was detected in Brazil.

The patient provided informed written consent to perform all studies described here and sequences were deposited in Genbank (accession numbers EF151263 for the S/P region and EF153194 for the precore/core region).

![Figure 1. Phylogenetic tree. Phylogenetic analysis obtained with the MEGA3 software of a 593-bp fragment of hepatitis B virus (HBV) DNA from the index case (S gene - GenBank accession number EF151263), 29 genotype E African sequences and two sequences of each of the other HBV genotypes. The tree was constructed by the neighbor-joining method and only bootstrap values higher than 80 are shown. A wooly monkey HBV sequence was used as the outgroup. GenBank accession numbers as well as the original countries of all HBV isolates are indicated. The length of the horizontal bar indicates the number of nucleotide substitutions per site.](image-url)
thereby generating a score of similarity. Sequences were submitted to BLAST in the GenBank database. A phylogenetic tree was constructed with the MEGA3 software (6) using a distance method (neighbor joining) and robustness of the tree was assessed by 1000 bootstrap replicates. The amino acid sequence obtained from the nucleotide sequence was also analyzed to assess subtype and possible drug resistance mutations in the HBV sequence.

Initial analysis of the nucleotide sequence of the S/P genomic region of HBV with the SeqScape Software V2.5 (Applied Biosystems) showed that it clustered with genotype E. Since this genotype has never been reported in Brazil, we performed a more extensive characterization of this virus. Phylogenetic analysis of the S/P region compared with reference isolates from GenBank representing all known HBV genotypes confirmed that it was classified into genotype E (Figure 1), and the HBsAg subtype was identified as ayw4 based on the presence of Arg122, Lys160, Leu127, and Ser140. Val to Leu mutation at position 209, present in all genotype E HBV sequences, was also found in this patient. No other amino acid mutations were found within the S protein and no drug resistance-related mutations were found in the HBV DNA polymerase protein. When submitted to BLAST in the GenBank, the sequence from the S/P region showed 99.07% similarity to several HBV genotype E sequences.

Mutation V209L is also found in genotype Aa, a discrete subtype within genotype A (7). Subgroup Aa has some amino acids differing from group Ae and was primarily found in sub-Saharan Africa. It was suggested that subgroup Aa may have evolved in indigenous populations of African countries. Later, this genotype was also found in South America (8,9), suggesting that this high proportion of subgroup Aa may be due to the intense slave trade from Africa to South America during the colonial period. In another study in a city with a large percent of African descendants (Salvador, BA, Brazil), genotype A was also the most prevalent in the local population, followed by genotypes D and F, but no case of genotype E was identified (10). The Afro-Brazilian population has a higher prevalence of HBV markers compared to other ethnic groups (11) which might be related to a higher HBV infection rate among the first black inhabitants that were brought to Brazil and/or to the low socioeconomic level of that population. The lack of other genotype E cases in Brazil and South America is very intriguing, since this genotype has a high prevalence in Africa and may also have come with the African slaves, besides subgroup Aa.

The low sequence diversity of genotype E in isolates from African patients over a large geographical area suggests a short evolutionary history in humans (12,13). Indeed, genotype E has never been found within the Americas (14) or in the Afro-Brazilian population (8,10,11,15), a fact suggesting the emergence of genotype E as a human pathogen from the mid to the late 19th century.

Analysis of the precore/core region showed G1896A and G1899A precore mutations but no mutations in the basal core promoter (BCP). In another study with genotype E-infected patients, precore mutation was detected in 75% of the patients without HBeAg, in contrast to only 25% of the patients with HBeAg. All the strains described so far have T1858 (16), as also does the index case described here. This substitution allows the G1896A mutation and prevents the production of HBeAg. BCP mutation was detected in a few patients previously described and there was no relation to the presence of HBeAg (16,17). Other substitutions common to all genotype E sequences were also found (A1757 and C1772) in this patient. Mutation G1757A abolishes COUP TFI and HNF-1 nuclear factors binding to BCP in experiments in vitro (18) and may influence the clinical characteristics of genotype E, probably being related to the mild
evolution observed in this case.

We described here the major features of the first HBV genotype E case found in São Paulo State, Brazil. This genotype has never been reported in Brazil. It was recently found in 2 Argentinian sisters whose mother is from the Dominican Republic (19). In Brazil, genotypes A, B, C, D, and F have already been described and their distribution varies among Brazilian regions (4,10,20), suggesting an influence of the immigration pattern for each region. Further molecular epide-

References

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