Epidemic history of hepatitis C virus infection in two remote communities in Nigeria, West Africa

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We investigated the molecular epidemiology and population dynamics of HCV infection among indigenes of two semi-isolated communities in North-Central Nigeria. Despite remoteness and isolation, ~15 % of the population had serological or molecular markers of hepatitis C virus (HCV) infection. Phylogenetic analysis of the NS5b sequences obtained from 60 HCV-infected residents showed that HCV variants belonged to genotype 1 ($n = 51; 85 \%$) and genotype 2 ($n = 9; 15 \%$). All sequences were unique and intermixed in the phylogenetic tree with HCV sequences from people infected from other West African countries. The high-throughput 454 pyrosequencing of the HCV hypervariable region 1 and an empirical threshold error correction algorithm were used to evaluate intra-host heterogeneity of HCV strains of genotype 1 ($n = 43$) and genotype 2 ($n = 6$) from residents of the communities. Analysis revealed a rare detectable intermixing of HCV intra-host variants among residents. Identification of genetically close HCV variants among all known groups of relatives suggests a common intra-familial HCV transmission in the communities. Applying Bayesian coalescent analysis to the NS5b sequences, the most recent common ancestors for genotype 1 and 2 variants were estimated to have existed 675 and 286 years ago, respectively. Bayesian skyline plots suggest that HCV lineages of both genotypes identified in the Nigerian communities experienced epidemic growth for 200–300 years until the mid-20th century. The data suggest a massive introduction of numerous HCV variants to the communities during the 20th century in the background of a dynamic evolutionary history of the hepatitis C epidemic in Nigeria over the past three centuries.

INTRODUCTION

Hepatitis C virus (HCV) infection is a serious public health problem worldwide. HCV causes chronic infection in 70–80 % of infected people. Chronic HCV infection may progress to chronic hepatitis and cirrhosis, leading in many cases to severe complications including hepatocellular carcinoma (HCC) and death (Itskowitz, 2007). An estimated 170 million people or ~3 % of the world’s population are chronically infected with HCV; and 3–4 million people are newly infected each year, with most of these cases occurring in Africa (EASL, 1999; WHO, 1999).

HCV is a positive-strand, ssRNA virus classified in the genus *Hepadnavirus* of the family *Flaviviridae* (Shepard et al., 2005). The HCV genome consists of ~9600 nt and encodes a single long polyprotein with the following gene order: 5’-C-E1-E2-p7-NS2-NS3-NS4A-NS4B-NS5A-NS5B-3’ (Lindenbach & Rice, 2005). The HCV genome displays considerable sequence divergence. HCV has been classified into six genotypes, 1–6, with each genotype further subdivided into subtypes displaying different geographical distributions worldwide (Simmonds et al., 2005). For example, HCV subtype 5a and all subtypes of genotype 6 are mainly restricted to South Africa and South East Asia, respectively (Simmonds et al., 1993), while subtypes 1a, 1b, 2b and 3a are distributed globally (Smith et al., 1997). A new genotype 7 has been recently registered in the HCV databases, but the full report on this isolate has not been published (Nakano et al., 2012).

In Africa, divergent strains of HCV genotypes 1 and 2 were found in Ghana, Guinea Conakry, Burkina Faso, Benin Republic and Guinea Bissau, suggesting that these strains have been endemic in the West African subregion (Candotti et al., 2003; Jeannel et al., 1998; Markov et al., 2009; Ruggieri et al., 1996; Wansbrough-Jones et al., 1998). Recently, analysis of the epidemic history of HCV infections traced the modern HCV lineages in West Africa back to the 17th–20th
centuries (Markov et al., 2009; Pouillot et al., 2008b). This estimate, however, does not take into account Nigeria, the country with more than three-quarters of the population of West Africa, which occupies a geographically central position within this African subregion. Identifying HCV circulating in Nigeria is important for guiding HCV infection control initiatives, understanding the evolutionary dynamics and natural history of HCV infection among indigenous West Africans, and predicting the future burden of HCV-related disease in Africa.

The epidemiology of HCV infection in Nigeria is not well understood. Most studies of hepatitis C in Nigeria have focused on serological characterization of selected population groups, e.g. prison inmates, patients with diabetes mellitus, blood donors, HIV-infected persons, patients with chronic renal failure and those with sickle cell anaemia, for whom risk for HCV infection in urban areas of Nigeria is variable. Previous studies established a broad HCV-seroprevalence rate, ranging from 1.9% among apparently healthy individuals with a family history of diabetes in Plateau State or among HIV-positive patients in Lagos (Balogun et al., 2010; Nwankiti et al., 2009; Onakewhor & Okonofua, 2009). HCV infections have been shown to play a significant role in the aetiology of chronic liver disease and HCC in Nigeria (Laraba et al., 2010; Mustapha et al., 2007). Despite HCV endemicity, association with HCC and importance of HCV genotypes for clinical management (Hnatyszyn, 2005; Zeuzem, 2004), only three small-scale studies have been made to genetically characterize HCV isolates from Nigeria (Agwale et al., 2004; Mellor et al., 1995; Oni & Harrison, 1996). As a result, an extremely limited number of HCV sequences from Nigeria are available in any repository of nucleotide sequences. Genotype distribution and genetic characterization of HCV strains circulating in different African countries, including Nigeria, are critical for designing universally efficacious vaccines against HCV infection, predicting the sensitivity of diagnostic assays and understanding the evolutionary history and molecular epidemiology of HCV in Africa.

Here, we present the first comprehensive phylogenetic and evolutionary analysis of HCV variants obtained from 60 residents of two rural, remote communities in North Central Nigeria.

**RESULTS**

**HCV prevalence**

Samples (n=519) were collected from the study sites (341 from village 1 and 178 from village 2) and tested for anti-HCV and for HCV RNA by the NS5b real-time PCR. A total of 73 samples (55 from village 1 and 18 from village 2) were found to be anti-HCV-positive and 60 samples were PCR-positive (47 samples from village 1, 13 from village 2), with 55 samples being both anti-HCV- and PCR-positive. Thus, a total of 78 samples were identified that had serological or molecular markers of HCV infection. An overall high prevalence of 15% of HCV infection in these communities represents one of the highest estimates reported in Nigeria (Balogun et al., 2010; Nwankiti et al., 2009; Onakewhor & Okonofua, 2009). Of the 60 PCR-positive individuals, males (n=33) and females (n=27) were almost equally represented. Although, the anti-HCV seroprevalence was slightly higher in men than women (sex ratio=1.2), this difference is not statistically significant. The majority (63.3%) of the infection was found in individuals between the ages of 20 and 40 years.

**Phylogenetic relationships among HCV variants**

Considering the rural location and relative isolation, it was expected that these two small communities would have limited exposure to HCV and should have had a single or only a few HCV strains circulating among the indigenous residents. However, analysis of the NS5b sequences identified HCV variants belonging to genotypes 1 (n=51; 85%) and 2 (n=9; 15%) (Fig. 1). Furthermore, all genotype 1 and 2 variants segregated into subtypes, some of which were not previously characterized (Fig. 1). These findings indicate that the two communities were infected with several HCV strains, suggesting either numerous exposures or exposure to all these HCV lineages in bulk from limited infection events.

The number of HCV genotype 1 variants in these communities was fivefold greater compared with the number of genotype 2 variants. The NS5b sequence analysis did not reveal any community-specific clustering and showed extensive phylogenetic intermixing of these sequences with other HCV isolates from different African countries and strains found worldwide. Consistent with the NS5b phylogenetic analysis, genetic diversity of 17.3 and 16.2% was observed for the genotype 1 and 2 variants, respectively. The high genetic diversity suggests a long evolutionary history. However, the presence of two genotypes and several subtypes as well as intermixing with HCV variants found worldwide strongly suggest that many, if not all, of these variants were introduced rather than evolved in the two communities. Although many variants could not be classified into any known subtype, there was a trend for the variants to form clusters.

**HCV variants from West and Central Africa**

The diversity of the HCV variants observed in the two communities was compared to HCV genotype 1 and 2 strains identified in other West and Central African countries (Fig. 2). HCV NS5b sequences from Central Africa had the tendency to form compact and distinct clusters. A few of these sequences intermixed with sequences from West Africa (Fig. 2). It should be noted, however, that among 141 sequences from Central Africa,
136 were obtained from Cameroon, with only five sequences being from two immediate neighbouring countries (Table 1). Conversely, 107 sequences from West Africa originated from six countries within the West African subregion (Table 1). These sequences were more diverse and grouped with many of the variants determined in this study, thus indicating a genetic relatedness of HCV variants identified among the two communities and HCV strains broadly circulating in West Africa. However, the distribution of HCV variants between genotypes 1 and 2 in these communities in Nigeria is very different from the other West African countries. In the Nigerian communities, 85% of HCV variants belong to genotype 1, while only 15% of HCV variants recovered from the other West African countries belong to this genotype.

**Intra-host HCV heterogeneity**

The presence of two HCV genotypes, several classified/unclassified subtypes and numerous HCV variants raise questions about the origin and maintenance of HCV diversity in these two communities. To investigate genetic connections among these variants in the population, quasispecies analysis of HVR1 was conducted using high-throughput pyrosequencing of 43 HCV genotype 1 and six genotype 2 samples. The small size of the communities and
circulation of many HCV variants suggests a potential for frequent mixed infections. However, analysis of quasi-species revealed a very limited intermixing of HCV quasi-species among residents (Figs 3 and 4).

There were six groups of relatives included in the analysis: group 1 – kng005 (20 years, female), kng006 (40 years, male) and kng007 (33 years, female); group 2 – kng021 (22 years, male), kng022 (45 years, male), kng23 (14 years, male) and kng024 (12 years, male); group 3 – kng036 (60 years, male) and kng037 (4 years, male); group 4 – kng020 (30 years, female) and kng59 (32 years, male); group 5 – kng025 (40 years, female) and kng026 (50 years, female); and group 6 – kng047 (44 years, male) and kng048 (35 years, male). HCV intra-host variants were found clustered together among relatives within groups 1, 2, 3, 5 and 6 (Figs 3 and 4). Although HCV variants identified from two relatives of group 4, kng020 and kng059, segregated into two distinct branches (Fig. 1), some intra-host variants found in kng020 intermixed with variants from kng059 (Fig. 3). Additionally, HCV intra-host variants from kng020 were also found to be intermixed with variants from kng025 (Fig. 3). These findings indicate a probable intra-familial HCV transmission or infections of relatives from a common source. However, clusters of genetically close intra-host variants were also identified among unrelated residents. For example, clusters of HCV variants from kng001 (15 years, male), kng002 (10 years, female), kng003 (28 years, female) and kng004 (28 years, male), or kng031 (60 years, female), kng032 (45 years, female), kng033 (30 years, female) and kng039 (50 years, male) were found among non-relatives. It is important to note that kng039 did not reside in the same community with kng031, kng032 and kng033. The identification of genetically close HCV variants among males and females of a broad age range (4–60 years) residing in different communities suggests a complex pattern of HCV dissemination within the communities, with the intra-familial transmission being common.

Population dynamics

In order to investigate the origin and spread of HCV genotype 1 and 2 variants in these Nigerian communities we used the Bayesian coalescent framework implemented in BEAST. Each genotype was analysed separately. The inferred dates for the most recent common ancestor (tMRCA) for genotypes 1 and 2 were 1335 Common Era (C.E.) [95 % highest posterior density (95 % HPD): 1127–1529] and 1721 C.E. (95 % HPD: 1645–1791), respectively. Both genotypes experienced epidemic growth (Figs 5 and 6). The effective population size of genotype 1 increased progressively during the 18th–20th centuries and reached a plateau by the mid-20th century (Fig. 5). The genotype 2 expansion began during the mid-18th century and stabilized in the mid-20th century (Fig. 6). The genotype 2 expansion began during the mid-18th century and stabilized in the mid-20th century (Fig. 6). The genotype 2 showed a rapid exponential expansion between ~1920 and 1960. In support of these observations, Fu’s Fs test rejected the null hypothesis of neutrality and constant population for HCV genotype 1 ($P=0.0001$) and genotype 2 ($P=0.0006$) variants. Collectively, the data suggest that HCV
Table 1. GenBank sequences used in this study

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*HCV-1, HCV genotype 1, HCV-2, HCV genotype 2.*
Fig. 3. Maximum-likelihood tree of intra-host HVR1 variants identified in 43 individuals infected with HCV genotypes 1. All sequences from a single individual are shown using the sample identification code. The three samples that share variants have been identified with the same colour. Dotted circles represent members of the same family. Bootstrap values higher than 60% are indicated at the major nodes.

Fig. 4. Maximum-likelihood tree of intra-host HVR1 variants identified in six individuals infected with HCV genotypes 1. All sequences from a single individual are shown using the sample identification code and same colour. Dotted circle represent members of the same family. Bootstrap values higher than 60% are indicated at the major nodes.
variants of both genotypes identified in the Nigerian communities experienced population expansion for about 200–350 years.

**DISCUSSION**

Nigeria has a high burden of chronic hepatitis C and HCC associated with HCV infection (Balogun et al., 2010; Laraba et al., 2010; Mustapha et al., 2007; Nwankiti et al., 2009). The rate of HCV infection was estimated to range from 1.9 to 14.5% in the country (Balogun et al., 2010; Nwankiti et al., 2009; Onakewhor & Okonofua, 2009). This study was carried out in two remote communities in North Central Nigeria. Despite the relative isolation of these communities, ~15% of the population experienced HCV infection. The observation of the high HCV prevalence in the villages lends additional support to high level of endemicity of HCV in Nigeria.

Owing to the isolation of these small communities, it was expected that only a few HCV strains would have been introduced and transmitted within the population. However, phylogenetic analysis of the NS5b sequences from 60 isolates obtained in this study revealed the presence of many distinct HCV variants and subtypes belonging to genotypes 1 and 2 (Fig. 1). HCV genotype 1 was detected in 85% and genotype 2 in 15% of all PCR-positive specimens. This finding is in agreement with three previous studies from Nigeria that showed predominance of HCV genotype 1, with variants of genotypes 2 and 4 being reported as minority genotypes (Agwale et al., 2004; Mellor et al., 1995; Oni & Harrison, 1996).

Although isolates in this study were collected from individuals living in closed communities, all HCV NS5b sequences were different from each other. Phylogenetic analysis showed extensive intermixing of the HCV variants identified among residents of the Nigerian communities and reported from other countries of West Africa (Fig. 2), indicating their common genetic history. These findings suggest that the heterogeneous HCV variants did not independently evolve within these remote communities but rather were introduced. Due to limited contacts with the general population, it is conceivable that few opportunities have been available for extraneous transmission, thus indicating that the introduction of the many HCV strains to the communities occurred in bulk. Alternatively, there is a remote possibility of the incremental introduction and accumulation of HCV variants but that would take a long period of time.

Analysis of HVR1 quasispecies showed that over a half of 49 HCV-infected members of the communities shared closely related HCV variants (Figs 3 and 4), thus revealing a complex pattern of potential HCV transmissions. The high HCV prevalence in these closed communities and the presence of many HCV lineages should have generated frequent opportunities for mixed infections. However, with the exception of kng020 sharing HCV variants with kng025 and kng059 (Fig. 3), no intermixing was observed among HCV quasispecies from different residents. Consistent with this finding, mixed HCV infections were rarely observed among such highly exposed groups as injection drug users, who experience only transient infections with more than one HCV variant (Viazov et al., 2000, 2010). These observations suggest a significant interference among HCV
lineages that thwart their co-existence in the same host. HCV superinfection exclusion is a possible molecular mechanism contributing to such interference (Tscherné et al., 2007).

The principal mode of transmission resulting in such a complex pattern of transmission is not clear. Findings of genetically close HCV quasispecies among relatives indicate that intra-familial transmission is a common route of HCV spread in the communities. It was previously suggested that sharing of equipment used for parenteral practices is an important route of HCV transmission in rural regions of Africa (Ndong-Atome et al., 2009; Njouom et al., 2003). However, such parenteral modes as body scarification, clitoridectomy using unsterilized instruments, circumcision, shaving by local barbers, that are known to be responsible for HCV transmission in Africa (Adewole et al., 2009; Ocama & Seremba, 2011; Pépin et al., 2010), could not be responsible for the introduction of HCV genotypes 1 and 2 strains to these villages but could play a role in maintaining modern lineages of the virus. We can only speculate about the possible introduction and transmission events that occurred many years ago.

The existence of heterogeneous HCV variants reflects a long-term presence and diversifying evolution of this virus and may be associated with a particularly extensive transmission mode in West Africa. A high prevalence of HCV genotype 4 lineages in Egypt had been linked to massive treatment campaigns against schistosomiasis between 1920 and 1980 (Frank et al., 2000). Studies from other African countries have argued that the HCV epidemic was driven iatrogenically, mostly by campaigns against trypanosomiasis, yaws and syphilis (Njouom et al., 2007; Pépin, Labbé, 2008; Pouillot et al., 2008b). Given this scenario, massive iatrogenic interventions could have jump-started the HCV epidemic in the Nigerian population. However, our data provide no direct indication on the events that could have led to such high prevalence of divergent HCV variants in the two communities.

Bayesian coalescence analysis indicated that the tMRCA for HCV genotype 1 and 2 lineages in the two Nigerian populations existed approximately 675 and 286 years ago, respectively. The tMRCA for genotype 1 variants identified in this study pre-dates the tMRCA for HCV genotype 1 reported globally (Ferraro et al., 2008; Lampe et al., 2010; Magiorkinis et al., 2009; Pépin et al., 2010). This finding is consistent with many suggestions that HCV genotype 1 originated in Africa and spread to other parts of the world (Markov et al., 2009; Simmonds et al., 1993, 2005). The tMRCA for HCV genotype 2 variants found in the Nigerian communities is similar to the reported estimate for other countries in West Africa (Pouillot et al., 2008a). It should be noted, however, that analysis of sequences linked to a single date of collection used in this study could result in estimating tMRCA inaccurately. The timescale approximation for HCV epidemic may vary considerably depending on the estimation of evolutionary rate (Lampe et al., 2010). The rate of $5 \times 10^{-4}$ substitutions per site per year used in the present analysis represents the best estimate of HCV NS5b evolutionary rates consistent with estimates made in several studies (Pouillot et al., 2008a; Pybus et al., 2001; Tanaka et al., 2002).

The Bayesian skyline plot (BSP) analysis showed that both genotype 1 and 2 populations experienced expansion (Figs 5 and 6). Fu’s statistical test supports this finding. HCV genotype 1 expansion steadily progressed for over 350 years starting in approximately 1700; the HCV genotype 1 population finally stabilized about the mid-20th century (Fig. 5). Genotype 2 expansion began in the mid-18th century and ended approximately during the mid-20th century (Fig. 6). A similar observation was made for genotype 2 lineages in Guinea-Bissau in West Africa, where the effective population size was shown to have been increasing from the mid-18th century. However, this growth had stabilized by approximately 1900 (Markov et al., 2009). A trend to stabilization was also observed in the genotype 1 and 2 BSPs at about the same time (Figs 5 and 6), indicating potential mitigation of the epidemic growth by the end of the 19th century. However, different from the genotype 2 expansion in Guinea-Bissau, both genotypes 1 and 2 in the Nigerian communities showed expansion before the mid-20th century (Figs 5 and 6). A similarly rapid spread of genotypes 1 and 2 from 1920 to 1950 was also observed in neighbouring Cameroon in Central Africa (Markov et al., 2009). However, this spread was not preceded by epidemic growth as seen in this study. Thus, the Nigerian expansion seems to combine the growth patterns observed in West and Central Africa. During the colonial period, in French speaking areas in Central Africa, unsafe injections practised mostly by mobile medical units for the prevention of sleeping disease early in 20th century were most probably responsible for the rapid HCV expansion in Cameroon (Njouom et al., 2007; Pouillot et al., 2008b). In English-speaking areas in West Africa including Nigeria, medical services have been broadly provided preferentially in stationary facilities since the 19th century (Millar & Foege, 1969) and could have contributed to the expansion during the 19th and mid-20th centuries in Nigeria through unsafe injection practices.

We have recently detected a population expansion of HBV genotype E lineages in these Nigerian communities (Forbi et al., 2010). Strikingly, this expansion occurred during approximately 1970–1980 or approximately 20 years later than the HCV expansion detected here. The tMRCA for the modern HBV genotype E was predicted to have existed approximately in the mid-20th century. The cause for this difference in time of expansion between two viral pathogens that share modes of transmission and co-circulate in the same communities is unknown. The difference in tMRCA indicates that the HCV genotype 1 and 2 lineages are much older than HBV genotype E (Forbi et al., 2010). It is conceivable that the epidemiological processes responsible for the origination of the modern HBV genotype E lineages (Forbi et al., 2010) could have contributed to the
continuation of expansion of the already abundant HCV variants during the mid-20th century.

Six of the nine HCV genotype 2 variants identified in the Nigerian communities share a single branch of the phylogenetic tree with the major cluster of the variants from Cameroon, indicating a strong genetic connection between the genotype 2 strains circulating in both countries. These variants represent a subset of all genotype 2 lineages identified in West Africa (Figs 1 and 2). Recently, it was suggested that the genotype originated in West Africa and spread from the most western part of this region toward the centre of the continent (Markov et al., 2009; Ndjomou et al., 2003; Pouillot et al., 2008b). The detection of genetic similarity between the predominant genotype 2 lineages in Nigeria and Cameroon is consistent with this hypothesis and further suggests that the genotype 2 strains were filtered through Nigeria before reaching Cameroon. Nigeria contributes over three-quarters of the population in West Africa and is a convergence point and centre for active population mixing in Africa. Owing to its geographical location and population density, Nigeria may have critically facilitated HCV transmission from West to Central Africa.

The data suggest that these two Nigerian communities have experienced a massive introduction of multiple HCV variants, which probably occurred around the mid-20th century. The extensive genetic variation among HCV variants observed in this study indicates that the introduction was rather recent because genetic drift would have reduced the number of HCV variants circulating in these small communities over an extended period of time spanning more than one human generation. The prevalence of ~15% is among the highest reported in West and Central Africa (Bekondi et al., 2010; Nagalo et al., 2011). Whether the prevalence and HCV genetic composition in these two communities represents the HCV population circulating in Nigeria as a whole needs further enquiry. This study is the first comprehensive investigation of the epidemic history of HCV in Nigeria revealing significant variation in the evolutionary histories of HCV genotypes 1 and 2 in two remote Nigerian villages that could be critical to understanding the dynamics of the diversity and spread of HCV in West and Central Africa.

**METHODS**

**Ethics statement.** Written informed consent was obtained from all participants involved in this study. The use of specimens in this study was approved by the Institutional Review Board (IRB) of the US Centers for Disease Control and Prevention and the Nigerian Federal Ministry of Health.

**Serum samples and serological testing.** A total of 519 blood samples were randomly obtained in 2007 from asymptomatic indigenes of two small rural communities in Keffi local government area of Nasarawa state in Nigeria, with 341 samples being from village 1 and 178 from village 2. These two villages with a total population of ~2000 inhabitants are ~2 miles apart. The residents have limited access to basic healthcare facilities.

All the studied samples were anonymous with a coding number for analysis. Samples were screened in the field for the presence of HCV antibodies using an immuno-chromatographic rapid assay (Shantha Biotechnics Ltd) following the manufacturer’s instructions. This assay uses recombinant antigen NS3, NS4 and NS5 to detect anti-HCV activity. Blood samples were centrifuged at 11,620 g for 10 min to separate serum from the cells. The serum samples were stored at −20°C until further testing was carried out.

**Nucleic acid extraction and cDNA synthesis.** Total nucleic acid was extracted from all serum specimens using the Roche MagNA Pure LC instrument and MagNA Pure LC Total Nucleic Acid Isolation kit (Roche Diagnostics), and eluted with 50 μl buffer according to the manufacturer’s instructions. RNA was precipitated and reverse-transcribed using both random and specific primers as described previously (Alter et al., 1999). Reverse transcription was carried out for 60 min at 42°C in a total volume of 20 μl × 5× PCR buffer (Roche), 200 pmol of each deoxynucleotide triphosphate, 1 μg random primers μl⁻¹, 25 U AMV reverse transcriptase (Roche) and 40 U RNase inhibitor (Roche), followed by heating at 95°C for 5 min.

**Molecular detection of NS5b gene by real-time PCR.** To identify HCV genotypes and subtypes, a segment of HCV NS5b encompassing positions 8275–8616 was amplified using real-time nested PCR. All amplification reactions were performed using the Fast-start DNA Master plus SYBR Green 1 kit and the Stratagene MX3005p QPCR machine (Agilent Technologies). First-round PCR was conducted using primers NS5B-K1 (5'-TGAGGATTCGCTGTGATGACCCGC-TGCTTTGTA-3') and NS5B-K2 (5'-GGCCGATTCCTTGGTCTGA-TAGCCGTCCGAGAA-3'). The nested reaction was performed with primers NS5B-122 (5'-CTCAACCCTACGTGAGAGACAT-3') and NS5B-R1 (5'-GCTTCTAGGTCGCGCGGTCGCTC-3'). PCR included incubation at 94°C for 2 min and 30 s, followed by 50 cycles of 94°C for 20 s, 45°C for 1 min and 30 s, followed by 50 cycles of 94°C for 20 s, 50°C for 25 s and 72°C for 25 s for 45°C. Nested PCRs with 45 cycles were performed using the same conditions with 2 μl of the first-round product diluted 1:100.

**Sequencing of the NS5b region.** Sequencing was performed using the second-round PCR products and nested primers. Sequencing reactions were conducted using the BigDye v3.1 chemistry sequencing kit (Applied Biosystems), and products were sequenced using an automated sequencer (3130xl Genetic Analyzer; Applied Biosystems). Sequencing PCR involved 25 cycles, each cycle consisting of 96°C for 10 s, 50°C for 5 s and 60°C for 4 min.

**Deep pyrosequencing of the HVR1 region.** Forty-three samples from patients infected with HCV genotype 1 (HCV-1) and six from patients with HCV genotype 2 (HCV-2) were used for quasispecies analysis. This subset was analysed using the next-generation sequencing technology (454/Roche GS FLX platform) on the HVR1 region of the genome. The junction E1/E2 region (309 nt), which contains the HVR1 region, was amplified using the nested PCR protocol described in Ramachandran et al. (2008). Each sample was amplified independently with fusion primers including the 454-primer key (A and B for forward and reverse primers, respectively), a different multiple identifier (MID) for each sample and HCV-specific sequence. The PCR products were purified using Agencourt AmpPure XP (Beckman Coulter). The quality of the amplicons was assessed using 3100 bioanalyser (Agilent Technologies). Purified amplicons were quantified using the Quan-it Picogreen dsDNA assay kit (Invitrogen). PCR amplicons were mixed at equimolar concentrations and diluted to a final concentration of 10 copies molecular ml⁻¹ prior to being subjected to emulsion (em)PCR, which was performed.
following the instructions supplied with the kit and enriched beads were subjected to pyrosequencing (titanium chemistry) using the 454/Roche GS FLX instrument. The original sequence reads (raw data) were processed using the SFFFILE tools. Sequence reads belonging to each sample were identified and separated using MID. Low quality reads were removed. The pyrosequencing files were post-processed with the empirical threshold (ET) error correction algorithm (Skums et al., 2011), which shows very high accuracy in finding true haplotypes, removing false haplotypes and estimating the frequency of true ones. The ET algorithm includes a calibration step using sequence reads from single-clone HVR1 samples, estimating an empirical frequency threshold for indels and haplotypes, and also correcting homopolymer errors using sequence alignment (Skums et al., 2011).

**Phylogenetic analysis.** Preliminary sequence analysis was conducted using SeqMan and MEGALIGN programs from the Lasergene DNA and Protein analysis software (version 8.0, DNASTAR Inc.). The Accelrys GCG Package (Genetic Computer Group, version 11.1-UNIX, Accelrys Inc.) was used for further analysis. Nucleotide sequences were aligned using the GCG multiple alignment program PILEUP. HCV genotypes/subgenotypes were classified based on the NS5b sequence (Smith et al., 1997) and by comparing each sequence with published reference sequences from GenBank. Initial phylogenetic trees were built using the Kimura two-parameter model of nucleotide substitution (Tamura et al., 2004). Phylogenetic trees were constructed using the maximum-likelihood algorithm implemented in DNAML (PHYLIP package, v.3.6). Bootstrap values were calculated using 1000 replicates.

All sequences generated in this study were further subjected to phylogenetic analysis together with published sequences (NS5b region) from West and Central Africa retrieved from GenBank (Table 1).

**Estimation of evolutionary dates and demographic history.** The date of the MRCA of HCV strains from Nigeria was calculated using 362 nt NS5b sequences from the two villages. These sequences belong to genotypes 1 (n=51) and 2 (n=9). Divergence times were calculated using BEAST (ver. 1.6.0) (Drummond & Rambaut, 2007). The GTR substitution model was used with four gamma categories and invariant sites. Codons were partitioned into three partitions with unlinked substitution model and unlinked rate heterogeneity model across the codon partitions. Each sequence alignment was analysed with a strict or relaxed clock with an initial estimate for the rate of substitution of $5 \times 10^{-4}$ (Njouom et al., 2007). Constant size, exponential and expansion growth priors were used. All models were run until the effective sample size for each was greater than 200. In order to compare the models the Bayes factor was estimated using importance sampling of the posterior probability (Newton et al., 1994). The exponential substitution model with an expansion growth prior was chosen because it had the largest Bayes factor; however, none of the models tested was found to be superior to any other model.

BSP analysis was done by segregating the NS5b sequences into the genotypes 1 (n=51) and 2 (n=9) groups. Each dataset was analysed separately using BEAST (ver. 1.6.0) (Drummond & Rambaut, 2007) with a strict clock, a substitution rate of $5 \times 10^{-4}$ and a piecewise-constant skyline model with six groups.

**Population dynamics of HCV in West and Central Africa.** Unbiased estimates of nucleotide diversity were calculated according to Nei (1987) using the program ARLEQUIN (Excoffier et al., 2005). Fu’s Fs test for detecting population growth, was calculated using ARLEQUIN (Excoffier et al., 2005). A P-value of 0.05 or less was considered statistically significant.

**Nucleotide sequence accession numbers.** Sequences from Nigeria reported in this study have been deposited in the National Center for Biotechnology Information database and can be retrieved under accession numbers JQ679028–JQ679087.

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