

Characterization of Hepatitis C Virus Circulating among Injecting Drug Users (IDU) in Kilifi County, Kenya.

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Abstract

Background: Hepatitis C virus is a major global health problem estimated to infect over 170 million people globally with the most common route of infection being injecting drug use (IDU). Treatment for HCV infection has traditionally been shown to be genotype specific; however the available drugs are still expensive and out of reach in many developing countries. To improve on monitoring, there is need to continuously document the genotypic burden and epidemiology in different populations arises.

Objectives: This study aimed to determine the circulating genotypes and link the data to the socio-demographics of injecting drug users in Kilifi County along the Kenyan coastline.

Methods: Using a random sampling method, this was a cross-sectional prospective study conducted among 127 injecting drug users, whereby ethical clearance was sought from Kenyatta National Hospital/ University of Nairobi Ethical and Research review committee (KNH/UON-ERC), and Reference number P366/07/2017 on 25 th September 2017. Serology for HCV was done followed by nucleic acid amplification and eventual genotyping. Socio-demographic data was collected using questionnaire administered at the sites.

Results: A total of 28 (23 males and 5 females) samples out of 127 samples were positive for HCV giving a prevalence of 22.1%. The modal age group was 25- 49 years. Of the positive samples, 11 were amplified by PCR, all from the male IDUs. Prevalent genotypes included genotype 1a (13%) and genotype 4a (87%). Both genotype 1a and 4a were only found in males.

Conclusion: Although it is the first time to report HCV4a in Kilifi town and Mtwapa as compared to Watamu and Malindi which had both HCV1a and HCV4a. Tourists born in countries where HCV prevalence is high are supposed to be screened before accessing entry to Kilifi County. No clinical trial was conducted. The study reveals the burden of HCV infection among IDUs in Kilifi County. The Government should formulate policies for intervention on testing and treatment of HCV in Kilifi County targeting IDUs in order to minimize spread to other populations.

Background

Hepatitis C virus causes of liver cancer such as [hepatocellular carcinoma](#) (HCC) and [lymphomas](#) in humans (Ferri 2015 and Rusyn 2014). Hepatitis C is a global health problem, over 130 million people are chronically infected yearly ([Amal 2015](#)). Studies show that every year, approximately 308,000 deaths occur due to liver cancer while up to 758,000 deaths are estimated to occur due to liver cirrhosis (Seyed and Farshid 2014). Approximately 80% of HCV infected persons will develop chronic hepatitis. Eleven percent will progress to liver cirrhosis for the period of 20-year time interval. Life-threatening will be as a result of liver failure which leads to hepatocellular carcinoma (Te and Jensen 2010, Longo *et al.*, 2011). Blood products, needle or syringe sharing among members of intravenous drug parties or undergoing a needle stick by health workers leads to transmission of HCV. Other risk factors are high-risk sexual behaviors, tattooing, reused and unsterilized dental and surgical instruments, and unsterilized laboratory

equipment (Samimi-Rad *et al.*, 2012). About 80% of HCV infected persons are asymptomatic. Infected persons with acute HCV exhibit symptom ranging from fever, joint pain and jaundice. Asymptomatic individuals are more difficult to identify (Lozano *et al.*, 2012; Lucas *et al.*, 2014).

Most IDUs with persistent infection are unaware of the infection, screening programs to identify patients will be required to prevent silent progression of the disease (Fabienne and Jay 2017; Mohd *et al.*, 2013; Lavanchy, 2011). In most case HCV is often first diagnosed in late stage. Due to slow and silent onset, many patients are unaware of their infection and at least 40% cases remain undetected. Chronic hepatitis C infection is difficult to assess, because it is frequently subclinical (Te and Jensen, 2010). Patients with chronic hepatitis C are at risk of cirrhosis and hepatocellular carcinoma and their contacts (especially injecting drug users, IDUs and commercial sex workers CSWs) at risk of acquiring the infection through exposure to the virus. Given this situation, it appears that there is no clear understanding of the contextual factors that continue to fuel the upsurge in HCV infections among the key populations such as IDUs and CSWs (Kenya drug laws and human rights 2018). Molecular genotyping of HCV identified in patients is necessary, for proposing therapeutic options (Longo *et al.*, 2011). It is assumed that some genotypes are more common in certain areas or groups of people. The high number of chronically infected individuals, the burden of disease and the absence of a vaccine indicates that treatment will form part of the control of the disease (Korir, 2013). The aim of this study was to determine the prevalence, social characteristics and HCV genotypes circulating among IDUs in rehabilitation centers in Kilifi County Kenya.

Methods

Study site and study design

This study was conducted in Kilifi County located within the Kenyan coastline. The design was cross sectional. Participants were recruited from harm reduction/rehabilitation centers within Malindi, Watamu, Kilifi town and Mtwapa {note that Kilifi town and Mtwapa is one centre splited into two called Muslim Welfare Association (MEWA)} located at Kilifi County.

Sample collection, transportation and storage

After informed consent, Socio-demographic data was captured using questionnaire which was administered at the facilities. The data captured included, name of centers, date of birth, residential area, marital status, religion, occupation and the level of education. Further five milliliters of venous blood was collected into EDTA vacutainer tubes. They were then transported in dry ice from rehabilitation Centers to KEMRI laboratories for analysis.

Serological tests

Viral RNA was extracted from 170 µl of serum by using Zymo Research Kit (Irvine CA, U.S.A.) according to the manufacturer's instructions. The RNA pellets were resuspended in 60 µl of TE buffer. The RNA concentrations were measured in ng/µl using NanoDrop spectrophotometer (ThermoFisher Scientific, Wilmington, Delaware, USA).

Polymerase Chain Reaction (RT-PCR) and Sequencing reaction

Nested RT-PCR was carried out using Zymo Research PCR Kit (Irvine CA U.S.A.) and GeneAmp PCR system 9700 (Applied Biosystems, Foster City, CA, USA). In reverse transcription and PCR reactions, both 5'UTR universal primers: forward KY80 (5'-GCA GAA AGC GTC TAG CCA TGG CGT-3') and reverse KY78 (5'-CTC GCA AGC ACC CTA TCA GGC AGT-3') were used. 35 cycles of PCR (94°C for 30 seconds, 54°C for 1 minute, and 68°C for 2 minutes) were performed HCV PCR products of 244 bp were visualized, and electrophoresed on 1.5% agarose gel (Fabienne and Jay 2017; (Medhat *et al.*, 2014)). The positive amplicons were purified using the ZR DNA sequencing clean up kit (Zymo, Irvine, CA, USA) according to the manufacturer's protocol to eliminate unincorporated primers and dNTPs (Lynnette *et al.*, 2019). The purified amplicons were sequenced with Big Dye kit, and the ABI PRIS M 3100 genetic analyzer (Applied Biosystems) according to manufacturer's instructions.

Sequence analysis

Samples were aligned with the representative sequences for each major genotype and subtype selected from the HCV database and Gene Bank using the Multiple Sequence Alignment Program, ClustalW. Homology and evolutionary distance pair-wise comparisons for percent nucleotide were made. (Medhat *et al.*, 2014). The phylogenetic analysis of HCV isolates was performed with MEGA7.0.14 software.

Statistical Methods

Statistical Package for the Social Sciences (SPSS) version 20 software (SPSS Inc., Chicago, IL, USA) was used for statistical analysis. Non-parametric Mann-Whitney U test was used to compare between genotypes with respect to quantitative variables. Chi square was used to analyze association between social demographics and circulating HCV genotypes. A p-value of less or equal to 0.05 was considered statistically significant (Valentine *et al.*, 2018).

Results

Demographic characteristics of study participants

A total of 127 IDUs out of 4587 registered patients were recruited for the study {107 (84.2%) were male and 20 (15.7%) were female}. The samples in each site were as follows; Malindi, (n =35), Watamu (n =43), and Kilifi and Mtwapa (n =49) respectively. The mean age for the three centers was found to be 33.7 years.

Prevalence of HCV among study participants

Anti-HCV positive included 10 IDUs from Kilifi, 4 from Watamu and 14 from Malindi making a total of 28 out of 127 participants. Twenty three IDUs (82.1%) were male while 5 (17.9%) were women. Of the 28 IDUs, 11 (8.7%) male were confirmed to be HCV positive using PCR as compared to none for female IDUs. A total 28 participants screened, seroprevalence was established at 22.1%. With the highest prevalence found in males (18.1%; n=23) and low in females at (3.9%; n=5) (P=0.004). The seroprevalence in Malindi was 40%, Watamu 9.3% and Kilifi and Mtwapa 20.4% respectively. Table 1

Serology outcome and demographic characteristics

The highest number of injecting drug users was in the age group of 31-35(39%),it also had a higher number of IDUs who tested positive 8 (20.5%) as compared to other age brackets. In the study, 35 IDUs sampled were married, 68 single and 24 divorced. (OR=1.0, 3.0, 2.0. P = 0.065, 0.330 and CI: 0.9-9.7, 0.5-8.6 respectively). Those of the Muslim faith had higher number of HCV infections, 16 compared to Christians, 12 and pagans zero. (OR= 1.0, 1.4 and P= 0.497, 0.309 respectively). In education, Lower primary had 13, Upper primary 6 and Secondary and Tertiary 9 (OR=1.0, 0.8, 2.1, P = 0.682, 0.146. CI: 0.3-2.3) for upper primary, CI: 0.8-5.8 for secondary and tertiary respectively). Tout/drivers/riders group of IDUs had highest HCV infection 12, (OR=1), Beach boys were 5, Fishermen were 3 and other IDUs with different occupations were 7 respectively. (OR=1.0, 1.3, 0.8, 0.9. P= 0.688, 0.712, 0.831 CI: 0.4-4.3, 0.2-3.1, 0.3-2.6 respectively). Table 2

Polymerase chain reaction (PCR) and genotyping

The 28 samples which were positive were subjected to PCR, 11 (39.3%) were confirmed positive (all males) and none of the 5 female IDUs was positive. The 11 samples were then subjected for sequencing and results gave 3 out of the 11 samples as HCV Genotype 1 subtype a (HCV1a) while 8 were HCV Genotype 4 subtype a (HCV4a). Table 3

Phylogenetic tree construction

With reference of Full-genome consensus sequences for different HCV genotypes (including genotypes 1, 2, 3, 4, 5 and 6), there was only genotype 1 and 4 (Figure 1). The rest there was no evident on the tree because there were no associated isolates. Maximum composite likelihood algorithms were utilized, and phylogenetic trees were constructed by the neighbor-joining method. The reliability of different phylogenetic groupings was evaluated by using the bootstrap re-sampling test from the MEGA program (1,000 bootstrap replications). (Medhat *et al.*, 2014; Charlotte *et al.*, 2019).

Discussion

In this study we have managed to establish a higher prevalence of HCV in male injectors 107 (84.2%) than females 20 (15.7%). We have also shown a changing trend of HCV genotype distribution with genotype 4a becoming more prevalent in this region as opposed to the traditional genotypes 1a, genotype 1a can be seen in Watamu and Malindi but not in Kilifi, it also shows that it is the first time to report HCV4a in Kilifi. Hepatitis C patients were found to be illiterate, it shows that lack of knowledge about the disease and its precautions become a strong cause for disease. In this study, there is an increase of HCV infection as compared to the study which was done by (Mwatela *et al.*, 2015) in the entire Coast which showed a prevalence of 16.7%. We have shown that there is increased diversity along the coastal strip centers like Mtwapa which is part of Kilifi town both are under Muslim welfare association (MEWA).

Genotyping of hepatitis C virus (HCV) is considered an important tool for epidemiological and clinical studies and valuable marker for disease progression and response to antiviral therapy (Amen *et al.*, 2017). However, development of antiviral drugs, vaccines, and genotyping assays has a major impact on HCV this is because HCV has high levels of genetic diversity

(Cuypers *et al.*, 2015; Medhat *et al.*, 2014; Vanessa *et al.*, 2018). In this study, we have confirmed that only two genotypes circulating in Kilifi County; this can form a basis for more targeted control for the 2 subtypes in the region. Hepatitis C Virus diversity is very essential in response to antiviral therapy (Medhat *et al.*, 2014; Tawhida *et al.*, 2015). More severe liver disease and more elevated liver enzymes have been highly associated with HCV genotype 1 (Chakravarti *et al.*, 2011). The endemicity of HCV genotypes in some areas is reversed with multiplicity and diversity of subtypes and strains. A good example is origin of genotypes 1 and 2 to West Africa and genotype 4 to Central Africa (James *et al.*, 2014). This can be done by using a phylogenetic analysis to revealed two monophyletic clusters (bootstrap value, ~87) containing HCV infected patients in Kilifi County from which a partial 5'UTR sequence was available.

The largest cluster contained 87% of HCV sequences were identified as genotype 4, with isolates M01, M02, M05, M07, M23, K20, W10, and W14. This isolates clusters with isolates from: Portugal, France, Egypt, Cyprus, Southeast Asia, Middle East and Saudi Arabia. On the other hand genotype 1a Isolates were K08, K14 and W17 Clusters with isolates from: America, Japan, Indonesia, China and India. Genotype diversity is particularly high in China and many Southeast Asian countries and also in Western Europe and Australia, perhaps as a result of population immigration from Africa and/or Asia

(International migration data 2012). In countries like these, vaccine efficacy at a population level would be dependent on the generation of cross reactive immunity; an alternative approach could also be the development of vaccines hosting different immunogens. In this study, HCV genotype 4a and genotype 1a is predominant among the population of drug users, this suggest that there is a need for continuous monitoring of these for better treatment outcomes that entirely depend on HCV genotyping results. With global travel, increasing of tourist to Kilifi Count and the geographic zone of this region dictates the distribution of this genotypes in this region (Rong et al., 2014).

Due to lack of vaccine and effective therapy, the prevention of HCV infection has been a great challenge to developing country and owns one-fifths of the world's population (Tanaka et al., 2011). World Health Organization Assembly approved on 2016 the Global Health Sector Strategy to eliminate hepatitis infection by 2030, (Assembly WHOS-NWH 2016). It introduced global targets for the care and management of HCV including 90% reduction in new cases of chronic hepatitis C, 65% reduction in hepatitis C deaths, and treatment of 80% of eligible people with chronic hepatitis C infections (WHO 2016). To achieve these goals, the country need to develop national policies based on reliable epidemiological evidence (Saraswat et al., 2015). In Kilifi County, Screening populations at risk and Counseling program to IDUs with history of sexual contact or sharing of items should be done to stop spread of HCV in the region. Tourists born in countries where HCV prevalence is high are supposed to be screened before accessing entry to Kilifi County. However, data are often outdated and conflicting, making evidence-based policy and resource allocation difficult.

Conclusions

In this study, the prevalence of HCV in the area seems to be increasing as compared to previous studies. It is also the first time to report HCV4a in Kilifi town as compared to Watamu and Malindi which had both HCV1a and HCV4a. The study reveals the burden of HCV infection among IDUs in Kilifi County. The Government should formulate policies for intervention on testing and treatment HCV in Kilifi County targeting IDUs in order to minimize spread to other populations.

List Of Abbreviations

Bp	Base pair
CI	Confidence interval
CSW	Commercial Sex workers
dNTPS	Nucleoside triphosphate
EDTA	Ethylenediaminetetraacetic acid
DNA	Deoxyribonucleic acid

HCC	hepatocellular cancer
HCV	Hepatitis C virus
HCV1a	Hepatitis C virus Genotype one Subtype a
Hcv4a	Hepatitis C virus Genotype Four Subtype a
IDUs	Injecting Drug Users
KANCO	Kenya AIDS NGOs Consortium
KEMRI	Kenya Medical Research
MEWA	Muslim Education Welfare Association
n	Participants who are either positive or Negative
n/N	Percentage of participants
N	Number of participants
OR	odds ratio
P=	P Value
PCR	Polymerase Chain Reaction
RNA	Ribonucleic acid
RT-PCR	Ribonucleic acid
SPSS	Statistical Package for the Social Sciences
TE buffer	Buffer used to solubilize DNA or RNA, while protecting it from degradation
μl	microliters
5'UTR	Five prime untranslated region
WHO	World Health Organization
ZR	Zymo Research

Declarations

Ethical Consideration

Ethical clearance was sought from Kenyatta National Hospital/ University of Nairobi Ethical and Research review committee (KNH/UON-ERC), Reference number P366/07/2017 on 25th September 2017.

Consent

Consent to participate is attached together with the worksheet and Manuscript

Availability of Data and Materials

It is available in IDUs worksheet that I have attached.

Competing Interest:

There was no competing interest in this project.

Funding

The project was funded by Children investment fund foundation (CIFF). File Number: E- 2008. The owner of the project frontier to buy for me reagents since I was a student, I dint have a source of funds for my project.

Author's Contributions

M. O. R. He is the author, his work was to collect samples, do laboratory work (both serological and molecular work) and writing of manuscript.

O. O. E. He supervised laboratory work, assisted in field work, review manuscript writing and data analysis.

B. K.P. His work was to review manuscript and administrative work.

O. K. A. His work was to review manuscript and laboratory work analysis.

M. G. J. His work was field work and laboratory analysis

R. K. V. His work was to assist in laboratory analysis.

M. S. C. He assisted in funding of the project, field work and administrative work.

L. W. R. He supervised laboratory work, review manuscript writing and data analysis.

All authors have read and approved this manuscript for publication.

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Tables

Table 1

Variable	Rehabilitation Centers			Overall	P Value
	KILIFI, n=49	WATAMU, n=43	MALINDI, n=35		
	n (%)	n (%)	n (%)	n (%)	
Rapid Test					
Negative	39 (79.6)	39 (90.7)	21 (60.0)	99 (78.0)	0.004
Positive	10 (20.4)	4 (9.3)	14 (40.0)	28 (22.1)	
PCR					
HCV1a	0	1 (33.3)	2 (66.7)	3 (27.3)	0.121
HCV4a	5 (100.0)	2 (66.7)	1 (33.3)	8 (72.7)	

The table above reveals the Prevalence of Hepatitis C Virus (HCV) which is 22.1% and its Genotypes among study participants in Kilifi County.

Table 2

Variable	N	(n/N) %	Positive n (%)	Negative n (%)	OR (95%CI)	P value
Age						
16-20 years	3	66.7	2 (66.7)	1 (33.3)	1.0	
21-25 years	12	25.0	3 (25.0)	9 (75.0)	0.2 (0-2.6)	0.171
26-30 years	29	10.3	3 (10.3)	26 (89.7)	0.1 (0-0.8)	0.011
31-35 years	39	20.5	8 (20.5)	31 (79.5)	0.1 (0-1.6)	0.070
36-40 years	21	33.3	7 (33.3)	14 (66.7)	0.3 (0-3.3)	0.265
41-45 years	14	7.1	1 (7.1)	13 (92.9)	0.04 (0-0.9)	0.014
46-50 years	7	57.1	4 (57.1)	3 (42.9)	0.7 (0-11.3)	0.778
51-55 years	2	0	0	2 (100.0)	-	0.136
Gender						
Male	107	21.5	23 (21.5)	84 (78.5)	1.0	
Female	20	25.0	5 (25.0)	15 (75.0)	1.2 (0.4-3.7)	0.729
Marital status						
Married	35	11.4	4 (11.4)	31 (88.6)	1.0	
Single	68	27.9	19 (27.9)	49 (72.1)	3.0 (0.9-9.7)	0.065
Divorced	24	20.8	5 (20.8)	19 (79.2)	2.0 (0.5-8.6)	0.330
Religion						
Muslim	77	0	16 (20.8)	61 (79.2)	1.0	0.497
Christian	46	26.1	12 (26.1)	34 (73.9)	1.4 (0.6-3.2)	
Pagan	4	20.8	0	4 (100.0)	-	0.309
Education						
Lower primary and below	65	20.0	13 (20.0)	52 (52.5)	1.0	
Upper primary	36	16.7	6 (16.7)	30 (30.3)	0.8 (0.3-2.3)	0.682
Secondary and above	26	34.6	9 (34.6)	17 (17.2)	2.1 (0.8-5.8)	0.146
Occupation						
Tout/driver/riders	55	21.8	12 (21.8)	43 (78.2)	1.0	
Beach boy	19	26.3	5 (26.3)	14 (73.7)	1.3 (0.4-4.3)	0.688
Fisherman	17	17.6	3 (17.6)	14 (82.4)	0.8 (0.2-3.1)	0.712
Other	35	20.0	7 (20.0)	28 (80.0)	0.9 (0.3-2.6)	0.837

The table above shows correlation between Serological outcome and demographic characteristics of Participants in Kilifi County.

Key: N= Number of participants, n= participants who are either Positive or negative (n/N) = Percentage of participants, OR= Odds Ratio, CI = Confidence Intervals.

Table 3

Variable	HCV1a (%)	HCV4a (%)	P value
Age	Age		
16-20 years	0	0	0.426
21-25 years	0	0	
26-30 years	0	2 (100.0)	
31-35 years	1 (25.0)	3 (75.0)	
36-40 years	1 (33.3)	2 (66.7)	
41-45 years	0	1 (100.0)	
46-50 years	1 (100.0)	0	
51-55 years	0	0	
Gender			
Male	3 (27.3)	8 (72.7)	-
Female	0	0	
Marital status			
Single	2 (40.0)	3 (60.0)	0.452
Married	0	3 (100.0)	
Divorced	1 (33.3)	2 (66.7)	
Religion			
Pagan	0	0	0.425
Muslim	2 (22.2)	7 (77.8)	
Christian	1 (50.0)	1 (50.0)	
Education			
Lower primary and below	1 (25.0)	3 (75.0)	0.145
Upper primary	0	4 (100.0)	
Secondary and above	2 (66.7)	1 (33.3)	
Occupation			
Tout/driver/riders	2 (40.0)	3 (60.0)	0.488
Beach boy	0	3 (100.0)	
Fisherman	1 (50.0)	1 (50.0)	
Other	0	1 (100.0)	

The table above shows correlation of Polymerase Chain Reaction (PCR) and Genotypes with participant demographic characteristics in Kilifi County. It reveals the percentages of Hepatitis C virus Genotype one Subtype a {HCV1a (%)} and Hepatitis C virus Genotype four Subtype a {HCV4a (%)} among the IDUs in Kilifi County.

Figures

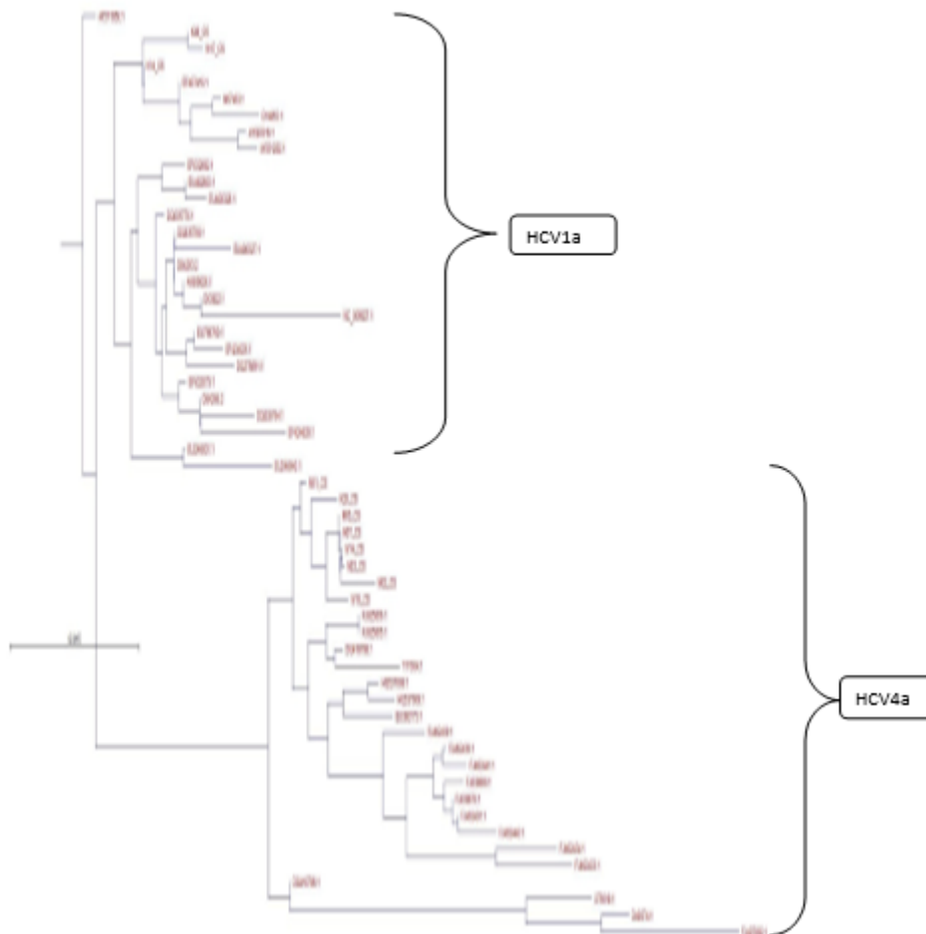


Figure 1

Phylogenetic neighbor-joining trees of Hepatitis C Virus (HCV) 5'UTR partial sequences in Kilifi County, it revealed two monophyletic clusters (bootstrap value, ~87) containing Hepatitis C Virus Genotype one Subtype a (HCV1a) and Hepatitis C Virus Genotype four Subtype a (HCV4a) among infected patients in the County.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- [IDUsWorksheets.xlsx](#)