

Prevalence Of Non-Alcoholic Fatty Liver Disease Using Non-Invasive Techniques Among Children, Adolescents, And Youths Living With Hiv

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Running head: Non-alcoholic fatty liver disease in perinatally HIV-infected children

Summary: Based on non-invasive imaging techniques, the prevalence of non-alcoholic fatty liver disease is high among children and youths acquiring HIV perinatally compared to controls. Scores based on clinical and analytical parameters do not identify individuals at risk.

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Abstract

Objective: The prevalence of subclinical liver abnormalities is high among people living with HIV, but data regarding perinatally HIV-infected children and adolescents (PHIV) are scarce. Non-invasive image techniques offer an opportunity to address non-alcoholic fatty liver disease (NAFLD) in a population in which the scores validated for adults have not been tested.

Design: Prospective cross-sectional study including PHIV and uninfected controls.

Methods: Non-invasive imaging techniques for the diagnosis of NAFLD and/or fibrosis were performed, and four scores to predict NAFLD were evaluated.

Results: Seventy-six participants (59.2% women) with a median of 19 years old (IQR: 15.5 – 25.6) were included, 38 were PHIV and 38 were age and sex-matched controls. All HIV participants were on ART at the moment of inclusion, and 86.8% were virologically suppressed. A total of 11 PHIV and three controls were diagnosed with NAFLD (28.9% vs. 7.9%; $p=0.02$) by non-invasive imaging techniques. The performance of scores based on clinical and analytical parameters was very poor. Although non-significant, overweight was more common among participants with NAFLD, who had a significantly higher BMI. Differences in HIV-related parameters between the groups were non-significant, except for the CD4/CD8+ T-cells ratio, decreased among PHIV diagnosed with NAFLD ($p=0.04$).

Conclusions: The prevalence of NAFLD was high (28.9%) among PHIV, and only partially explained by overweight and metabolic syndrome defining factors. The scores based on clinical and analytical parameters did not accurately identify subjects at risk. Therefore, liver ultrasound assessment should be considered for the screening of NAFLD among PHIV in routine clinical practice.

Keywords: Non-alcoholic fatty liver disease (NAFLD), Human immunodeficiency virus (HIV), Antiretroviral therapy (ART), transient elastography, ultrasound, vertical transmission, children, adolescents

Introduction

Since the introduction of antiretroviral therapy (ART), HIV infection has turned into a chronic condition [1]. As a result of this increase in life expectancy, the quality of life of people living with HIV is threatened by comorbidities, including liver, kidney, cardiovascular disease, or cancer [2,3]. Worldwide, the liver disease remains one of the major causes of morbidity and mortality among people living with HIV [4,5]. However, thanks to the efficacy of direct-acting antivirals (DAA) to treat the hepatitis C virus (HCV), the burden of diseases associated with this virus is decreasing dramatically in areas with access to treatment. Instead, the weight of non-alcoholic fatty liver disease (NAFLD) and steatohepatitis on chronic liver disease is increasing.

NAFLD is defined as a fat accumulation higher than 5% in the liver, encompassing different stages of abnormal liver ranging from simple steatosis to non-alcoholic steatohepatitis (NASH) and cirrhosis [3,6]. The increase in NAFLD keeps pace with the increasing prevalence of obesity, insulin resistance [7,8] and other components of the metabolic syndrome [9,10], and is linked to mitochondrial dysfunction [11] and the cytopathic effect of HCV infection [12]. Some reports suggest that NAFLD/NASH will turn into the first cause of liver disease in the general population, both for children and adults, and the first cause of liver transplant in western countries in years to come [3,10]. However, the diagnosis of NAFLD/NASH is challenging, and although we have very sensitive and specific non-invasive methods for steatosis and fibrosis diagnosis in our daily clinical practice, to date, liver biopsy is considered the gold standard [13]. Scores combining clinical and laboratory parameters, such as the hepatic steatosis index (HSI) [14] or triglycerides and glucose (TyG) [15], have been validated in adults for screening and are able to predict NAFLD in the adult population. The AST to platelet ratio index [16] and the fibrosis-4 index and NAFLD fibrosis score [17] are recommended to detect fibrosis. None of these scores has been validated for children [18,19].

New, non-invasive imaging techniques as transient elastography (T.E.) or point shear wave elastography (p-SWE) offer the possibility to stratify the degrees of steatosis [13,20], assessing hepatic rigidity in a quantitative way and can be combined with the mentioned scores to improve accuracy. Their main limitation is a low sensitivity for the diagnosis of early stages of NAFLD/ NASH, according to the European Association for the Study of the Liver [21]. However, up today, the use of non-invasive imaging techniques is recommended for the diagnosis and follow-up both in adults and children [16,21,22].

Among people living with HIV, data suggest an increased prevalence of NAFLD [23–29] ranging from 28.8 [30] to 48.0% [31] when the diagnosis is based on imaging techniques to 57.1 [32] to 72.6% [25] when based on liver biopsy. Although the pathogenesis of NAFLD/NASH is unclear and most probably multifactorial, in the context of HIV infection, authors have suggested a potential deleterious effect of the chronic inflammation and activation of the immune system secondary to the virus [28,33–35] and its treatment [36,37]. In the unique population of perinatally HIV-infected children who face lifelong exposure to

antiretroviral treatment and its metabolic consequences, and the deleterious effects of the virus itself [18] these phenomena have been described since birth [38–40]. However, data addressing the prevalence of NAFLD in this population are scarce [41]. The reliability of non-invasive assessment of NAFLD, including clinical scores in the population of adults living with HIV, has been the focus of intense research, but their use among children and youths has not been established [17]. Although hypertransaminasemia is quite common during follow-up in people living with HIV, its significance remains unknown [40].

The aim of this study was to determine the prevalence of subclinical liver abnormalities based on the use of non-invasive image techniques (TE/p-SWE) in perinatally HIV-infected children and youths (PHIV). Secondary objectives included the description of clinical, epidemiological, virological, and immunological determinants and evaluation of clinical scores for the diagnosis of NAFLD among children and youths.

Methods

Study design and participants

We carried out a prospective longitudinal study in two tertiary hospitals in Madrid, Spain. PHIV followed up in the Spanish National Network of Children and Adolescents Living with HIV (CoRISpe) were included. Uninfected siblings, partners, and adolescents who attended for post-exposure prophylaxis were included as controls, matched by gender and age (± 2 years). HIV participants and uninfected controls were recruited at Hospital Universitario La Paz and Hospital General Universitario Gregorio Marañón in Madrid (Spain) from June 2018 to December 2020. All patients included in the CoRISpe cohort undergo serology at inclusion and during follow-up according to clinical criteria, and all patients included in the study were screened for a full panel of diseases with liver involvement, including viral infections. Additional exclusion criteria for participants and controls were history of HBV or HCV infection or co-infection, acute infections or opportunistic diseases, or chronic inflammatory diseases at the moment of inclusion, and reporting $>30g$ in men and $>20g$ in females of alcohol intake. The study was approved by the Ethics Committee of participating hospitals and conducted according to the Declaration of Helsinki. All participants signed informed consent before inclusion in the study.

Demographic, clinical, analytical, and virological data

Sociodemographic and clinical data were collected from medical records. Virological and immunological data, including HIV-specific variables and ART history, were collected for PHIV from the CoRISpe database. CDC 2014 classification was used to categorize HIV disease stages. HIV viral load was considered suppressed when < 50 copies/mL. HBV and HCV participants were excluded, and no other hepatic diseases were discarded. Liver enzymes (AST, ALT, and GGT) were considered elevated according to laboratory ranges: ALT > 35 UI/L, AST and GGT when > 40 UI/L. Fasting total cholesterol was considered elevated when > 200 mg/dL and triglycerides when >150 mg/dL. Weight, height, waist, and

hip measurements were taken according to a unified protocol. Body Mass Index (BMI) was automatically calculated according to the formula: $BMI = \text{weight (kg)} / \text{height}^2 \text{ (m)}$. All measurements were z-score adjusted. The standard deviations for BMI according to age and gender were calculated using the 2010 Spanish Growth charts [42]. Overweight was defined as a BMI z-score between +1 and +2 SD, and obesity was defined as BMI z-score ≥ 2 SD. For adults ≥ 18 years of age, we used the WHO definition for overweight and obesity based on a BMI ≥ 25 and ≥ 30 respectively [43].

Non-invasive techniques for NAFLD diagnosis

All patients included in the study underwent both transient elastography (with CAP) and Shear wave elastography, as well as hematology and biochemistry, serological and virological studies. Ultrasound elastography measurements were performed in a supine position with the right arm maximally abducted, after ≥ 6 hours of fasting. Two trained clinicians, blind to HIV status, performed the ultrasound studies according to a unified protocol.

Liver fibrosis was evaluated by means of transient elastography (TE), FibroScan [44], as well as Shear Wave Elastography. Liver elasticity/stiffness was quantified based on the speed of ultrasound waves when passing through the liver. Point Shear Wave Elastography (pSWE) uses an internal acoustic radiation force impulse or ARFI, measuring the speed of shear wave propagation through the hepatic parenchyma at a fixed point or ROI. It was performed with a 4-9 MHz convex probe in a Samsung Prestige RS80 ultrasound scanner [45]. Ten valid measurements were acquired at 2-4 cm in-depth in the liver. The measured speed is given in m/s and converted to kPa for tissue stiffness estimation [46]. Then, the median is calculated, as well as the interquartile range. Normal values have been defined for children [47]. The risk for advanced fibrosis was stratified by APRI (< 0.5 mild fibrosis), advanced fibrosis p-SWE and/or TE Liver fibrosis was defined according to the Metavir Scale for the TE as the absence of fibrosis F0-F1 (< 7.6 kPa), moderate fibrosis F2 (7.6-9.5 kPa), advanced fibrosis F3 (9.5-14 kPa) and F4 (14 kPa); and for p-SWE as the absence of fibrosis F0-F1 (< 5.7 kPa), moderate fibrosis F2 (> 7.6 kPa), advanced fibrosis F3 (< 11.6 kPa) and F4 (> 11.6 kPa).

Both pSWE and the Controlled Attenuation parameter (CAP) by TE were used to establish NAFLD diagnosis. CAP evaluates the ultrasonic attenuation in the liver at 3.5 MHz at depth 25 to 65 mm using FibroScan and represents a noninvasive assessment of liver steatosis. CAP values in dB/m were reported as the median of 10 acquisitions, and the cut-off point of 248 dB/m defined by Karlas et al [48] was used to define the presence of steatosis. The criteria defined by Dasaranthy S et al. [49] and Brill F et al. [50] were used for the diagnosis of steatosis by pSWE: increased parenchymal echogenicity of the liver, hepatic vein blurring, portal vein blurring and visualization of the diaphragm.

Clinical and analytical scores

Four scores validated to predict liver abnormalities in the adult population were calculated for study participants, and their accuracy to predict fibrosis/NAFLD was compared to non-invasive imaging techniques. As there are no specific pediatric cut-off points, the ones described for adults were used. Triglyceride Glucose Index (TyG), Hepatic Steatosis Index (HSI), AST to Platelet Ratio Index (APRI), and Fibrosis 4 (FIB-4) were calculated automatically. TyG, an index to determine insulin resistance and able to identify individuals at risk for NAFLD, was calculated according to the equation: $TyG = \ln [Fasting\ triglyceride\ (mg/dl) \times Fasting\ glucose\ (mg/dl)]/2$. An index of 4.49 or larger indicates a patient is likely to suffer from insulin resistance and an index of 8.38 or larger indicates steatosis. HSI, an index used to identify candidates for liver study, was calculated using the equation: $HSI = 8 \times ALT/AST + BMI (+ 2\ if\ type\ 2\ diabetes, + 2\ if\ female)$. For interpretation, NAFLD was diagnosed >36 , ruled out with a value <30 , while values between 30- 36 are considered inconclusive. Two scores for fibrosis evaluation were performed: $APRI = AST\ (IU/L) / AST\ upper\ normal\ limit\ (IU/L) / Platelet\ count\ (10^9/L) \times 100$. When $APRI < 0.5$, fibrosis can be ruled out. FIB-4 was also used to evaluate liver fibrosis: $FIB\ 4 = (Age \times AST) / (Platelet\ count \times \sqrt{ALT})$. A value >2.67 suggests fatty liver, and > 3.25 suggests cirrhosis by HCV; a value <1.3 is relevant to exclude significant fibrosis. All index and interpretations were performed using MDCalc (<https://www.mdcalc.com/>).

Statistical analysis

Categorical variables were presented as total counts and percentages (%), and continuous variables appeared as median and interquartile ranges (IQR). Chi-square test or Fisher test, as appropriate, were performed to compare categorical variables and U Mann-Whitney test to compare continuous variables. Due to sample size restrictions, we did not perform multivariate analysis to assess determinants for NAFLD. A p-value less than 0.05 was considered statistically significant. All analyses were conducted in IBM-SPSS Statistics Version 25.0 (Armonk, NY: IBM Corp).

Results

General characteristics of the study population

Globally, 76 participants (59.2% women) with a median age of 19.0 (IQR: 15.5 – 25.6) years old were included in our study. Of them, 38 were PHIV, and 38 were age and sex-matched HIV-negative controls. The sociodemographic and clinical characteristics of both groups are shown in Table 1. Participants living with HIV had been diagnosed at a median of 19.2 (IQR: 3.6 – 56.4) months of age. All PHIV were on ART at the moment of inclusion in the study, and 86.8% of them were virologically suppressed. Overall, the most used recent combination (44.7%) was 2 Nucleoside Reverse Transcriptase Inhibitors (NRTI) + 1 Integrase Strand Transfer Inhibitors (INSTI), followed by an 18.4% using 2 NRTI + 1 Protease Inhibitor (PI) and a 15.8% using 2 NRTI + 1 Non-Nucleoside Reverse Transcriptase Inhibitors (NNRTI).

The median number of ART regimens received since diagnosis was 4.5 (IQR: 3.0 – 8.8). Half of PHIV had been exposed to AZT, 45% to stavudine (d4T) and over 36% to didanosine (ddI). Main HIV-related variables are shown in Table 2, according to NAFLD diagnosis.

NAFLD in PHIV

By means of non-invasive imaging techniques, a total of 11 PHIV participants were diagnosed with steatosis, vs. only three of the HIV-negative controls (28.9% vs. 7.9%; $p=0.02$). Comparisons between participants with and without NAFLD, including data regarding HIV-related parameters, are shown in Table 2. Overall, eleven PHIV were diagnosed with NAFLD by imaging techniques, at a median age of 17.9 years (IQR: 11.3 – 26.6), ranging from 9 to 33 years old. Characteristics of PHIV diagnosed with NAFLD are shown in Table 3. Seven were women and only two presented overweight (BMI >25). Their immunological situation was good, with only one participant having a CD4 Nadir below 200 cells/ml. The median number of previous ART regimens was 5 (IQR: 3 – 9), with five participants with previous exposure to ddI and d4T and eight to AZT. Fibrosis (stage F2 or higher) was detected in 3 (7.9%) PHIV and none of the HIV-negative controls. The three HIV-infected patients with fibrosis were male, aged 18.1, 19.2, and 27.8 years respectively. All were F2 according to the Metavir Scale (9.7, 8.2 and 8.8 KPa respectively) and only the last one had steatosis (patient #10, Table 3). The BMI of the other two patients was 22.5 and 22.7, respectively, CD4/CD8 ratio was greater than 1 in both cases with a CD4 nadir >200 cells/mL. None of them had an altered APRI or FIB4 index. No other risk factors for fibrosis were identified.

When compared to PHIV non presenting NAFLD, no differences were found regarding age, gender or ethnicity. Although non-significant, overweight was more common among participants with NAFLD, who had a significantly higher BMI. Differences in HIV-related parameters between the groups were non-significant, except for the CD4/CD8+ T-cells ratio, which decreased among those diagnosed with NAFLD ($p=0.04$). Time from diagnosis, CDC HIV stage, number of regimens and ART regimens were not associated with the prevalence of NAFLD.

Performance of the clinic-analytical scores for the screening of NAFLD in PHIV

Overall, no differences in laboratory parameters such as glucose, aminotransferases, cholesterol, triglycerides, platelets or bilirubin values were observed among PHIV, independent of the NAFLD diagnosis. Regarding screening scores, results were normal for most HIV-infected participants, presenting only 5 out of 30 PHIV with HIS score available with an altered index (NAFLD diagnosis confirmed by imaging techniques in three) and 2 out of 31 with an altered APRI (none of them had fibrosis by TE). FIB-4 and TyG scores were normal in all cases.

Discussion

In this exploratory study using non-invasive imaging techniques, the prevalence of NAFLD among youths living with HIV since childhood was surprisingly high (28.9%) compared to an uninfected cohort. The fact that no clear relation to overweight and the metabolic syndrome defining factors could be established, together with the extremely poor performance of the scores based on clinical and analytical parameters, are worrisome, as identification of patients at risk remains extremely challenging among youths.

Studies in industrialized countries suggest that NAFLD represents the most common chronic liver disease in children, with a prevalence ranging from around 10% in children, 17% in teenagers, to 40-70% among obese children [51]. Among adults in the general population, the numbers go up to 30% in adults and 80% among obese patients [52]. Compared to these data, the prevalence of NAFLD in our series, integrated mainly by teenagers and youths, is two times higher than expected by age group [51,52] supporting previous results in PHIV [53]. The first hypothesis to explain these findings is the presence of HIV infection and ART treatment, as underlined by the fact that NAFLD prevalence was much lower among the recruited uninfected controls. Mitochondrial dysfunction can be caused by ART and has been described to contribute to the pathogenesis of NAFLD and lead to ROS production, lipid peroxidation and cytokine release that induce liver inflammation and fibrosis [54]. This hypothesis is in line with the limited studies addressing the prevalence of NAFLD among PHIV [26,40], based on abnormal non-invasive markers [16] and non-invasive imaging techniques [41], suggesting abnormalities in around half of the studied populations [36,41]. The sample size of the mentioned studies is small, and methodologies cannot be compared. None of the studies is based on biopsies, which is meant to be the gold standard for diagnosis. However, these findings are worrisome, and especially considering that compared to other westernized countries, the Mediterranean region is usually considered a low-risk area for metabolic disorders [55]. According to the literature, the expected prevalence of NAFLD/NASH in the area would be lower compared to the one reported in areas such as North America [56–58]. Although results are worrisome, the longitudinal evolution of fatty liver is unknown, and studies have proven that changes in diet and behavior can induce disease regression [58]. Prompt diagnosis and management may prevent disease progression, as the rate of success is higher when lifestyle changes are promoted early in life [59,60]. The hypothesis of a direct relationship between liver damage and HIV-infection plus ART treatment in this cohort is plausible and has been pointed out in previous studies [27,61]. However, the fact that apart from the CD4/CD8 ratio, no HIV-related variables were independently associated with NAFLD, does not support the role of chronic inflammation or metabolic abnormalities secondary to ART as an underlying cause. Advanced age, years from HIV infection [28], higher viral load, increased liver enzymes and long exposure to NRTIs [29] and especially the use of ddI/d4T, have been previously described as risk factors for fibrosis progression among people living with HIV [35]. However, none of these factors were identified as risk factors for NAFLD in our cohort, nor time since diagnosis or ART exposure. The etiology of NAFLD is not well understood and is most probably multifactorial,

including genetic, behavioral, diet-related, and inflammatory factors. While the association between NAFLD and metabolic syndrome is beyond question [29,34,62] and has been described also in the context of HIV infection [35], several authors describe the presence of steatosis in patients not presenting any of the metabolic syndrome defining factors [63–67]. Although in this study the differences in terms of the prevalence of overweight among PHIV with and without NAFLD was non-significant, BMI was significantly higher in participants with NAFLD. Probably, this reflects how the reduced sample size impaired our ability to analyze weight as a dichotomous variable. However, as most scores include biochemical and clinical parameters related to BMI, these subjects would never be identified as subjects at risk when classical scores are used. Similarly, hypertransaminasemia can be present or not at diagnosis [31,62]. This issue is even more challenging between children, adolescents and youth patients, in which scores have not been validated. In children, in fact, even the definition of metabolic syndrome is controversial [68,69].

Imaging techniques are not the gold standard for diagnosis of NAFLD but allow a non-invasive approach and permit close follow-up of patients, including monitoring disease regression. The main limitations are a low sensitivity for early stages of the disease, as changes suggestive of steatosis cannot be appreciated if the fat content is lower than 20-30%, and the interobserver variability, partially overcome with the use of standardized parameters such as p-SWE or CAP. Cirrhosis associated with NAFLD is hard to detect, and there are no ultrasound markers for steatohepatitis. Small participants may require the use of an adapted probe. On the other side, the adipose panniculus of participants with overweight might lead to low image quality. Both fibrosis and steatosis can modify the attenuation and dispersion of ultrasound waves, and this is in fact one of the limitations of the imaging techniques when evaluating the liver. None of these limitations should interfere with its sensitivity for the identification of higher-risk patients with no metabolic syndrome, and thus, to our view these techniques are ideal among individuals living with HIV. Furthermore, our results suggest a very low sensitivity of usual NAFLD scores, with no concordance with imaging diagnosis, which induces us to recommend against the use of scores for children and youths. Together with the limitation regarding diagnosis, which did not include liver biopsy and thus cannot be considered certain, the main limitation of our study is the small sample size. The reduced number of participants included impaired our ability to analyze HIV-related risk factors. However, despite the reduced strength of the study, we found statistically significant differences regarding prevalence between cases and uninfected controls.

As genetic and lifestyle-related factors might determine the risk for NAFLD, controls were recruited among siblings and uninfected partners, to achieve a comparable cohort in terms of epidemiological, genetic and lifestyle-related factors, leaving HIV status as the only differentiating factor. However, the potential effect of intrauterine ART exposure is always a concern. The pubertal status assessment was not included in the protocol and therefore puberty could not be assessed as a potential confounder for NAFLD diagnosis [70]. Data regarding alcohol consumption rely only on clinical anamnesis.

Despite all limitations, the unexpected higher prevalence of NAFLD observed in this cohort of PHIV, together with an 8% of fibrosis are extremely worrisome, and comparable to previous studies [53]. These findings emphasize the need to systematically screen for liver disease among PHIV, regardless of age or the presence of risk factors. According to our results, the absence of metabolic factors or abnormal liver enzymes does not rule out the possibility of NAFLD. Due to the bad performance of classic scores in this population, non-invasive ultrasound techniques are nowadays the best option for screening, reserving biopsy for diagnosis confirmation when progression is suspected. Furthermore, non-invasive imaging techniques could be informative during the follow-up, and to evaluate long-term evolution. There is an urgent need to include liver assessment into routine clinical practice, to achieve a prompt diagnosis before the disease progresses to fibrosis or steatohepatitis. The development of new and more sensitive diagnostic procedures, as well as biomarkers to predict progression to severe forms of the disease is mandatory.

Conclusion

Larger and longitudinal studies addressing the evolution of liver disease in PHIV are needed. Clinicians should be aware of the risk and consider the need for screening. According to our results, the performance of scores based on clinical and analytical parameters for the identification of patients at risk is poor among youths. Despite their limitations, imaging techniques should be considered when available. As specific therapeutic measures are under research, intensifying prevention of metabolic risk factors since childhood seems mandatory to avoid future comorbidities.

Authorship

I.C., A.L., C.B., S.A., M.L.N. and T.S. performed the research. T.S., M.L.M., A.O., S.A., designed the research study, L.E., M.J.M. C.B., M.L.N., and C.D. recruited patients and performed anthropometric measurements. A.L. and A.O. realized ultrasonographic measurements. I.C, S.A. and T.S. analyzed the data. I.C. and T.S. wrote the original draft. A.O., A.L., L.E., M.J.M., C.B., M.L.M., C.D., S.A. and M.L.N. critically discussed and revised the final version of the paper. All authors have read and approved the final manuscript.

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References

1. Smith CJ, Ryom L, Weber R, Morlat P, Pradier C, Reiss P, et al. Trends in underlying causes of death in people with HIV from 1999 to 2011 (D:A:D): A multicohort collaboration. *The Lancet*. 2014;384(9939):241–8.
2. Deeks SG, Lewin SR, Havlir DV. The end of AIDS: HIV infection as a chronic disease. *The Lancet*. 2013;382(9903):1525–33.
3. Van Welzen BJ, Mudrikova T, El Idrissi A, Hoepelman AIM, Arends JE. A Review of Non-Alcoholic Fatty Liver Disease in HIV-Infected Patients: The Next Big Thing? *Infectious Diseases and Therapy*. 2019;8(1):33–50.
4. Eyawo O, Franco-Villalobos C, Hull MW, Nohpal A, Samji H, Sereda P, et al. Changes in mortality rates and causes of death in a population-based cohort of persons living with and without HIV from 1996 to 2012. *BMC Infectious Diseases*. 2017;17(1):174.
5. The data collection on adverse events of anti-HIV drugs (D:A:D) study group, Smith C, Sabin CA, Lundgren JD, Thiebaut R, Weber R, et al. Factors associated with specific causes of death amongst HIV-positive individuals in the D:A:D study. *AIDS*. 2010;24(10):1537–48.
6. Goh GBB, McCullough AJ. Natural History of Nonalcoholic Fatty Liver Disease. *Digestive Diseases and Sciences*. 2016;61(5):1226–33.
7. Aboud M, Elgalib A, Kulasegaram R, Peters B. Insulin resistance and HIV infection: A review. *International Journal of Clinical Practice*. 2007;61(3):463–72.
8. Araujo S, Bañón S, Machuca I, Moreno A, Pérez-Elías MJ, Casado JL. Prevalence of insulin resistance and risk of diabetes mellitus in HIV-infected patients receiving current antiretroviral drugs. *European Journal of Endocrinology*. 2014;171(5):545–54.
9. Rinella ME. Nonalcoholic fatty liver disease a systematic review. *JAMA*. 2015;313(22):2263–73.

10. Smith SK, Perito ER. Nonalcoholic Liver Disease in Children and Adolescents. *Clinics in Liver Disease*. 2018;22(4):723–33.
11. Blas-García A, Apostolova N, Esplugues JV. Oxidative Stress and Mitochondrial Impairment After Treatment with Anti-HIV Drugs: Clinical Implications. *Current Pharmaceutical Design*. 2011;17(36):4076–86.
12. Yoon EJ, Hu K-Q. Hepatitis C Virus (HCV) Infection and Hepatic Steatosis. *International Journal of Medical Sciences*. 2006;3(2):53–6.
13. Papatheodoridi M, Cholongitas E. Diagnosis of Non-alcoholic Fatty Liver Disease (NAFLD): Current Concepts. *Current Pharmaceutical Design*. 2019;24(38):4574–86.
14. Fennoun H, El Mansouri S, Tahiri M, Haraj NE, El Aziz S, Hadad F, et al. Interest of hepatic steatosis index (HSI) in screening for metabolic steatopathy in patients with type 2 diabetes. *The Pan African medical journal*. 2020;37:270.
15. Zhang S, Du T, Zhang J, Lu H, Lin X, Xie J, et al. The triglyceride and glucose index (TyG) is an effective biomarker to identify nonalcoholic fatty liver disease. *Lipids in Health and Disease*. 2017;16(1):15.
16. Kapogiannis BG, Leister E, Siberry GK, Van Dyke RB, Rudy B, Flynn P, et al. Prevalence of and progression to abnormal noninvasive markers of liver disease (aspartate aminotransferase-to-platelet ratio index and Fibrosis-4) among US HIV-infected youth. *AIDS*. 2016;30(6):889–98.
17. Castera L, Friedrich-Rust M, Loomba R. Noninvasive Assessment of Liver Disease in Patients With Nonalcoholic Fatty Liver Disease. *Gastroenterology*. 2019;156(5):1264–1281.
18. Siberry GK. Preventing and Managing HIV Infection in Infants, Children, and Adolescents in the United States. *Pediatrics in Review*. 2014;35(7):268–86.
19. Hartley A, Santos Ferreira DL, Anderson EL, Lawlor DA. Metabolic profiling of adolescent non-alcoholic fatty liver disease. *Wellcome Open Research*. 2018;3:166.
20. Sasso M, Beaugrand M, de Ledinghen V, Douvin C, Marcellin P, Poupon R, et al. Controlled attenuation parameter (CAP): A novel VCTETM guided ultrasonic attenuation measurement for the evaluation of hepatic steatosis: Preliminary study and validation in a cohort of patients with chronic liver disease from various causes. *Ultrasound in Medicine and Biology*. 2010;36(11):1825–35.
21. Bailey SS, Youssfi M, Patel M, Hu HH, Shaibi GQ, Towbin RB. Shear-wave ultrasound elastography of the liver in normal-weight and obese children. *Acta Radiologica*. 2017;58(12):1511–8.

22. Drescher HK, Weiskirchen S, Weiskirchen R. Current Status in Testing for Nonalcoholic Fatty Liver Disease (NAFLD) and Nonalcoholic Steatohepatitis (NASH). *Cells*. 2019;8(8):845.
23. Sterling RK, Smith PG, Brunt EM. Hepatic steatosis in human immunodeficiency virus: A prospective study in patients without viral hepatitis, diabetes, or alcohol abuse. *Journal of Clinical Gastroenterology*. 2013;47(2):182–7.
24. Kapoor N, Audsley J, Rupali P, Sasadeusz J, Paul TV, Thomas N, et al. A gathering storm: HIV infection and nonalcoholic fatty liver disease in low and middle-income countries. *AIDS*. 2019;33(7):1105–15.
25. Morse CG. Fatty liver disease in HIV: Common, underappreciated, and understudied. *AIDS*. 2017;31(11):1633–5.
26. Cervo A, Shengir M, Patel K, Sebastiani G. NASH in HIV. *Current HIV/AIDS Reports*. 2020;17(6):601–14.
27. Kaspar MB, Sterling RK. Mechanisms of liver disease in patients infected with HIV. *BMJ Open Gastroenterology*. 2017;4(1):e000166.
28. Maurice JB, Patel A, Scott AJ, Patel K, Thursz M, Lemoine M. Prevalence and risk factors of nonalcoholic fatty liver disease in HIV-monoinfection. *AIDS*. 2017;31(11):1621–32.
29. Guaraldi G, Squillace N, Stentarelli C, Orlando G, D’Amico R, Ligabue G, et al. Nonalcoholic fatty liver disease in HIV-infected patients referred to a metabolic clinic: Prevalence, characteristics, and predictors. *Clinical Infectious Diseases*. 2008;47(2):250–7.
30. Lui G, Wong VWS, Wong GLH, Chu WCW, Wong CK, Yung IMH, et al. Liver fibrosis and fatty liver in Asian HIV-infected patients. *Alimentary Pharmacology and Therapeutics*. 2016;44(4):411–21.
31. Vuille-Lessard É, Lebouché B, Lennox L, Routy JP, Costiniuk CT, Pexos C, et al. Nonalcoholic fatty liver disease diagnosed by transient elastography with controlled attenuation parameter in unselected HIV monoinfected patients. *AIDS*. 2016;30(17):2635–43.
32. Lemoine M, Barbu V, Girard PM, Kim M, Bastard J-P, Wendum D, et al. Altered hepatic expression of SREBP-1 and PPAR γ is associated with liver injury in insulin-resistant lipodystrophic HIV-infected patients. *AIDS*. 2006;20(3):387–95.
33. Serrano-Villar S, Pérez-Elías MJ, Dronda F, Casado JL, Moreno A, Royuela A, et al. Increased risk of serious non-AIDS-related events in HIV-infected subjects on

antiretroviral therapy associated with a low CD4/CD8 ratio. PLoS ONE. 2014;9(1):e85798.

34. Crum-Cianflone N, Dilay A, Collins G, Asher D, Campin R, Medina S, et al. Nonalcoholic fatty liver disease among HIV-infected persons. *Journal of Acquired Immune Deficiency Syndromes*. 2009;50(5):464–73.
35. Vodkin I, Valasek MA, Bettencourt R, Cachay E, Loomba R. Clinical, biochemical and histological differences between HIV-associated NAFLD and primary NAFLD: A case-control study. *Alimentary Pharmacology and Therapeutics*. 2015;41(4):368–78.
36. Pillaye JN, Marakalala MJ, Khumalo N, Spearman W, Ndlovu H. Mechanistic insights into antiretroviral drug-induced liver injury. *Pharmacology Research and Perspectives*. 2020;8(4):e00598.
37. Jones M, Núñez M. Liver toxicity of antiretroviral drugs. *Seminars in Liver Disease*. 2012;32(2):167–76.
38. Sainz T, Serrano-Villar S, Díaz L, Isabel González Tomé M, Dolores Gurbindo M, Isabel de José M, et al. The CD4/CD8 ratio as a marker T-cell activation, senescence and activation/exhaustion in treated HIV-infected children and young adults. *AIDS*. 2013;27(9):1513–3.
39. Sainz T, María Álvarez-Fuente †, Navarro ML, Díaz L, Rojo P, Blázquez D, et al. Subclinical Atherosclerosis and Markers of Immune Activation in HIV-Infected Children and Adolescents: The CaroVIH Study. *Journal of Acquired Immune Deficiency Syndromes*. 2014;65(1):42–9.
40. Pokorska-Śpiewak M, Stańska-Perka A, Popielska J, Ołdakowska A, Coupland U, Zawadka K, et al. Prevalence and predictors of liver disease in HIV-infected children and adolescents. *Scientific Reports*. 2017;7(1):12309.
41. Rubio A, Monpoux F, Huguon E, Truchi R, Triolo V, Rosenthal-Allieri M-A, et al. Noninvasive Procedures to Evaluate Liver Involvement in HIV-1 Vertically Infected Children. *Journal of Pediatric Gastroenterology and Nutrition*. 2009;49(5):599-606.
42. Carrascosa A. Aceleración secular de crecimiento en España. *Estudios Españoles de Crecimiento 2010. Población autóctona y población inmigrante. Endocrinología y Nutrición*. 2014;61(5):229–33.
43. Obesity and overweight What causes obesity and overweight? Global Health Observatory (GHO) More on obesity More on nutrition Fact sheet on malnutrition Data. 2021. Available from: <https://www.who.int/news-room/fact-sheets/detail/obesity-and-overweight>

44. Gu J, Wang W, Zou Z, Huang F, Fang C, Li X, et al. Emerging trends and new developments in transient elastography: A bibliometric and cocitation analysis from 1999 to 2017. *Canadian Journal of Gastroenterology and Hepatology*. 2019;2019:3280657.
45. Barr RG, Wilson SR, Rubens D, Garcia-Tsao G, Ferraioli G. Update to the Society of Radiologists in Ultrasound Liver Elastography Consensus Statement. *Radiology*. 2020;296(2):263–74.
46. Honda Y, Yoneda M, Imajo K, Nakajima A. Elastography techniques for the assessment of liver fibrosis in non-alcoholic fatty liver disease. *International Journal of Molecular Sciences*. 2020;21(11):40–39.
47. Mjelle AB, Mulabecirovic A, Havre RF, Rosendahl K, Juliusson PB, Olafsdottir E, et al. Normal liver stiffness values in children: A comparison of three different elastography methods. *Journal of Pediatric Gastroenterology and Nutrition*. 2019;68(5):706–12.
48. Karlas T, Petroff D, Sasso M, Fan JG, Mi YQ, de Lédinghen V, et al. Impact of controlled attenuation parameter on detecting fibrosis using liver stiffness measurement. *Alimentary Pharmacology and Therapeutics*. 2018;47(7):989–1000.
49. Dasarathy S, Dasarathy J, Khiyami A, Joseph R, Lopez R, McCullough AJ. Validity of real time ultrasound in the diagnosis of hepatic steatosis: A prospective study. *Journal of Hepatology*. 2009;51(6):1061–7.
50. Bril F, Ortiz-Lopez C, Lomonaco R, Orsak B, Freckleton M, Chintapalli K, et al. Clinical value of liver ultrasound for the diagnosis of nonalcoholic fatty liver disease in overweight and obese patients. *Liver International*. 2015;35(9):2139–46.
51. Clemente MG, Mandato C, Poeta M, Vajro P. Pediatric non-alcoholic fatty liver disease: Recent solutions, unresolved issues, and future research directions. *World Journal of Gastroenterology*. 2016;22(36):8078–93.
52. Schwimmer JB. Clinical advances in pediatric nonalcoholic fatty liver disease. *Hepatology*. 2016;63(5):1718–25.
53. Aepfelbacher JA, Balmaceda J, Purdy J, Mattingly A, Zambell K, Hawkins K, et al. Increased Prevalence of Hepatic Steatosis in Young Adults with Lifelong HIV. *Journal of Infectious Diseases*. 2019;220(2):266–9.
54. Nassir F, Ibdah JA. Role of mitochondria in nonalcoholic fatty liver disease. *International Journal of Molecular Sciences*. 2014;15(5):8713–42.

55. Tosti V, Bertozzi B, Fontana L. Health Benefits of the Mediterranean Diet: Metabolic and Molecular Mechanisms. *Journals of Gerontology - Series A Biological Sciences and Medical Sciences*. 2018;73(3):318–26.
56. Torres MCP, Aghemo A, Lleo A, Bodini G, Furnari M, Marabotto E, et al. Mediterranean diet and NAFLD: What we know and questions that still need to be answered. *Nutrients*. 2019;11(12):2971.
57. Katsagoni CN, Georgoulis M, Papatheodoridis G v., Fragopoulou E, Ioannidou P, Papageorgiou M, et al. Associations between Lifestyle Characteristics and the Presence of Nonalcoholic Fatty Liver Disease: A Case-Control Study. *Metabolic Syndrome and Related Disorders*. 2017;15(2):72–9.
58. Romero-Gómez M, Zelber-Sagi S, Trenell M. Treatment of NAFLD with diet, physical activity and exercise. *Journal of Hepatology*. 2017;67(4):829–46.
59. Anania C, Massimo Perla F, Olivero F, Pacifico L, Chiesa C. Mediterranean diet and nonalcoholic fatty liver disease. Vol. 24, *World Journal of Gastroenterology*. 2018;24(19):2083–94.
60. Jennison E, Patel J, Scorletti E, Byrne CD. Diagnosis and management of non-alcoholic fatty liver disease. *Postgraduate Medical Journal*. 2019;95(1124):314–22.
61. Sudjaritruk T, Bunupuradah T, Aurpibul L, Kosalaraksa P, Kurniati N, Sophonphan J, et al. Nonalcoholic fatty liver disease and hepatic fibrosis among perinatally HIV-monoinfected Asian adolescents receiving antiretroviral therapy. *PLoS ONE*. 2019;14(12):e0226375.
62. Nishijima T, Gatanaga H, Shimbo T, Komatsu H, Nozaki Y, Nagata N, et al. Traditional but not HIV-related factors are associated with nonalcoholic fatty liver disease in asian patients with HIV-1 infection. *PLoS ONE*. 2014 Jan 31;9(1):e87596.
63. Younes R, Bugianesi E. NASH in Lean Individuals. *Seminars in Liver Disease*. 2019;39(1):86–95.
64. Fracanzani AL, Petta S, Lombardi R, Pisano G, Russello M, Consonni D, et al. Liver and Cardiovascular Damage in Patients With Lean Nonalcoholic Fatty Liver Disease, and Association With Visceral Obesity. *Clinical Gastroenterology and Hepatology*. 2017;15(10):1604-1611.
65. Kim D, Kim WR. Nonobese Fatty Liver Disease. Vol. 15, *Clinical Gastroenterology and Hepatology*. 2017;15(4):474–85.
66. Albhaisi S, Chowdhury A, Sanyal AJ. Non-alcoholic fatty liver disease in lean individuals. *JHEP Reports*. 2019;1(4):329–41.

67. Phipps M, Wattacheril J. Non-alcoholic fatty liver disease (NAFLD) in non-obese individuals. *Frontline Gastroenterology*. 2019;11(6):478–83.
68. DeBoer MD. Assessing and managing the metabolic syndrome in children and adolescents. *Nutrients*. 2019;11(8):1788.
69. Weihe P, Weihrauch-Blüher S. Metabolic Syndrome in Children and Adolescents: Diagnostic Criteria, Therapeutic Options and Perspectives. *Current obesity reports*. 2019;8(4):472–9.
70. Suzuki A, Abdelmalek MF, Schwimmer JB, Lavine JE, Scheimann AO, Unalp-Arida A, et al. Association Between Puberty and Features of Nonalcoholic Fatty Liver Disease. *Clinical Gastroenterology and Hepatology*. 2012;10(7):786–94.

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Table 1. Characteristics of participants living with perinatally acquired HIV and HIV-negative controls. Continuous variables are expressed as medians and interquartile rates. Frequencies are expressed as percentages (%). BMI: body mass index, y: years, NAFLD: non-alcoholic fatty liver disease, KPa: Kilopascals, kg: kilograms, cm: centimeters.

Parameters	PHIV (n=38)	HIV-negative controls (n=38)	<i>p-value</i>
Age			0.16
Median (IQR), y	18.0 (14.9 - 24.2)	21.93 (14.9 – 27.0)	
Range	6.9 – 33.5	6.1 – 33.7	
Women (%)	22 (57.9)	23 (60.5)	0.82
Origin (%)			0.02
Spain	24 (63.2)	32 (84.2)	
Latinamerica	5 (13.2)	4 (10.5)	
Sub Saharan Africa	8 (21.1)	0 (0.0)	
Asia	1 (2.6)	2 (5.3)	
Weight, kg	54.0 (44.6 – 60.3)	60.7 (48.9 – 72.3)	0.04
Height, cm	158.0 (152.0 – 168.0)	169.3 (152.9 – 180.0)	0.02
BMI	20.3 (18.4 – 23.6)	22.0 (19.4 – 23.5)	0.27
Overweight ($1 < z\text{-score} < 2\text{DS}$ or BMI > 25)	6 (15.8)	7 (18.4)	0.76
Obesity (≥ 2 SD z-score or BMI > 30)	1 (2.6)	0 (0.0)	0.31
Waist, cm	71.0 (65.0 – 77.0)	75.3 (68.1 – 81.0)	0.25

Hip, cm	87.5 (79.0 – 90.8)	90.0 (77.1 – 94.0)	0.44
Waist-hip ratio	0.86 (0.79 – 0.91)	0.87 (0.83 – 0.92)	0.57
NAFLD diagnosis by imaging techniques (%)	11 (28.9)	3 (7.9)	0.02
Fibrosis (%)	3 (7.9)	0 (0)	0.21
FibroScan result, KPa	5.0 (4.2 – 5.6)	4.1 (3.6 – 5.1)	0.01

Table 2. Characteristics of participants living with perinatally acquired HIV with and without NAFLD.

Parameter	NAFLD (n=11)	No NAFLD (n=27)	Univariate (p)
Age, y	18.0 (11.3 – 26.6)	18.1 (15.8 – 23.3)	0.32
Women, (%)	7 (63.6)	15 (55.6)	0.65
Origin, (%)			0.19
Spain	9 (81.8)	13 (48.2)	
Latinamerica	2 (18.2)	3 (11.1)	
Sub Saharan Africa	0 (0.0)	8 (29.6)	
Asia	0 (0.0)	3 (11.1)	
BMI	23.3 (21.2 – 25.0)	19.1 (18.4 – 22.5)	0.04
Overweight, z-score +1-2 SD or BMI > 25 (%)	3 (27.3)	3 (11.1)	0.22
Obese, z-score +2 SD or BMI > 30 (%)	1 (9.1)	0 (0.0)	0.11
Waist hip ratio	0.91 (0.87 – 0.94)	0.83 (0.77 – 0.89)	0.01
FibroScan Value (kPa)	4.9 (3.9 – 6.4)	5.0 (4.5 – 5.6)	0.82
Time since HIV diagnosis, y	15.3 (10.1 – 25.7)	15.0 (11.3 – 22.9)	0.64

CDC HIV Stage			0.12
N	1 (9.1)	1 (3.7)	
A	6 (54.5)	5 (18.5)	
B	3 (27.3)	8 (29.6)	
C	1 (9.1)	10 (37.0)	
Unknown	0 (0.0)	3 (11.1)	
HIV VL <50 copies/mL, (%)	11 (100.0)	22 (81.5)	0.13
CD4+ T-cells count	839 (680 - 1420)	845 (678 - 1182)	0.61
CD4+ T-cells Nadir	302 (216 - 360)	393 (197 - 493)	0.32
CD4+ T-cells Nadir < 200 cells/mL	1 (9.1)	6 (22.2)	0.37
CD4/CD8+ T-cells Ratio	0.9 (0.6 - 1.1)	1.3 (1.0 - 1.7)	< 0.05
CD4/CD8 ratio < 1	4 (36.4)	5 (18.5)	0.21
ART regimens since diagnosis	5 (3 - 9)	3 (4 - 8)	0.76
Current ART, (%)			0.09
1 PI, (%)	2 (18.2)	1 (9.7)	
1 PI + 1 INSTI, (%)	0 (0.0)	2 (7.4)	
1 NNRTI + 1 INSTI, (%)	1 (9.1)	0 (0.0)	
2 NRTI + 1 INSTI, (%)	3 (27.3)	14 (51.9)	
2 NRTI + 1 PI, (%)	1 (9.1)	6 (22.2)	
2 NRTI + 1 NNRTI, (%)	4 (36.4)	2 (7.4)	
2 NNRTI + 1INSTI (%)	0 (0.0)	2 (7.4)	
Glucose (mg/dL)	80 (77 - 86)	82 (75 - 84)	0.68
ALT (U/L)	17 (13 - 26)	22.0 (11.8 - 29.3)	0.61

AST (U/L)	18 (15 – 21)	22.5 (15.5 – 30.5)	0.16
Altered Transaminases	0 (0.0)	0 (0.0)	
GGT (U/L)	18.5 (14.0 – 32.3)	17.5 (13.8 – 21.5)	0.65
Cholesterol (mg/dL)	173 (138 - 181)	153 (132 – 174)	0.41
Cholesterol > 200 mg/dL	1 (9.1)	4 (14.8)	0.64
LDL (mg/dL)	101 (75 - 117)	83 (68 – 99)	0.16
HDL (mg/dL)	45 (36 - 55)	50 (44 – 56)	0.16
Triglycerides (mg/dL)	83 (63 - 144)	79 (57 – 122)	0.59
Triglycerides > 150 mg/dL	2 (18.2)	4 (14.8)	0.80
Platelets (cells/mm ³)	311 (228 - 336)	254 (225 – 316)	0.28
Bilirubin (mg/dL)	0.4 (0.4 – 57.0)	0.5 (0.4 – 16.5)	0.81
HSI	33.2 (28.5 – 39.3)	27.8 (26.8 – 34.5)	0.01
Altered HSI	3 (27.3)	2 (7.4)	0.38
APRI	0.2 (0.1 – 0.2)	0.2 (0.2 – 0.3)	0.10
Altered APRI	0 (0)	2 (7.4)	0.66
FIB-4	0.3 (0.2 – 0.4)	0.4 (0.2 – 0.6)	0.48
Altered FIB-4	0 (0)	0 (0)	
TyG	4.4 (4.3 – 4.7)	4.4 (4.2 – 4.6)	0.63
Altered TyG	0 (0)	0 (0)	

Continuous variables are expressed as medians and interquartile rates. Frequencies are expressed as percentages (%). BMI: body mass index, V.L.: viral load, n: number, y: years, ART: antiretroviral treatment; NRTI: Nucleoside Reverse Transcriptase Inhibitors, NNRTI: Non-Nucleoside Reverse Transcriptase Inhibitors, P.I.: Protease Inhibitor, INSTI: Integrase Strand Transfer Inhibitors, ddI: didanosine, d4T: stavudine, AZT: zidovudine, ALT: Alanine Aminotransferase, AST: Aspartate Aminotransferase, GGT: Gamma-glutamyl Transferase, HSI: Hepatic Steatosis Index; APRI: AST to Platelet Ratio Index; FIB-4: Fibrosis-4 score; TyG: Triglycerides and fasting glucose index; ULN: upper limit normality.

Table 3. Characteristics of PHIV diagnosed with NAFLD.

Patient	Gender	Age (y)	Tanner	BMI	Waist hip ratio	Fibrosis	Nadir CD4	CD4/C D8 ratio	CDC Stage	ART (n)	Altered HSI
#1	F	17.8	V	24.2 (N)	0.91	F0-F1	340	0.31	A	4	Und
#2	M	11.3	II	23.4 (O)	0.94	F0-F1	217	1.27	A	5	No
#3	F	10.5	I	25.0 (O)	0.91	F0-F1	402	0.92	N	3	Und
#4	F	9.2	I	17.6 (N)	0.84	F0-F1	328	1.02	A	1	U
#5	F	18.0	V	23.2 (N)	U	F0-F1	360	0.86	A	2	Yes
#6	M	18.0	V	26.8 (O)	1.07	F0-F1	302	0.74	A	4	U
#7	M	26.1	V	21.9 (N)	0.89	F0-F1	158	U	C	9	No
#8	F	26.6	V	21.2 (N)	U	F0-F1	200	U	A	9	Und
#9	F	33.5	V	18.2 (N)	0.87	F0-F1	216	U	B	12	No
#10	M	27.8	V	43.3 (Ob)	U	F2	291	U	B	10	Yes
#11	F	26.0	V	23.3 (N)	U	F0-F1	639	U	B	7	Yes

Continuous variables are expressed as medians and interquartile rates. F: female, M: male, BMI: body mass index, PNPLA3: patatin-like phospholipase domain-containing 3 protein, ART: historical number of antiretroviral treatments used since HIV diagnosis, y: years, n: number, N: normal BMI, O: overweigh, Ob: obesity, Und: Undetermined; U: unknown.