Hepatitis C virus infection acquired in childhood

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Abstract Hepatitis C virus (HCV) infection occurs less frequently in children than in adult patients, and the natural history, prognosis, and clinical significance of HCV infection in children are poorly defined. We report here a descriptive follow-up of the clinical course, biochemical data, and viral markers observed in 37 children with anti-HCV. Ten patients included in the study tested persistently negative for serum HCV-RNA (group 1) and 27 patients tested persistently positive (group 2). In group 1, serum alanine aminotransferase (ALT) was normal in all patients, while two patients had non-organ-specific autoantibodies. In group 2, serum ALT was elevated in 13 of 27 patients, and five patients had non-organ-specific autoantibodies. HCV genotype 1a and 1b were the most prevalent among HCV-RNA-positive patients. Twenty liver biopsies were carried out on 17 patients in our series (mean evolution time, 11.2 years; range, 3–21 years). The liver specimens showed mild necroinflammatory changes in most patients, and fibrosis was absent or low grade. Two HCV-RNA-positive patients became persistently HCV-RNA negative. Of the 26 children investigated, 7 (one in group 1, six in group 2) had a co-infection with hepatitis G virus.

Conclusion Most children chronically infected with HCV were asymptomatic and presented only mild biochemical evidence of hepatic injury. Autoimmunity in the form of non-organ-specific autoantibodies was common. HCV in children induced mild changes in the liver with a low level of fibrosis and at a low rate of progression.

Keywords Anti-liver/kidney microsomal antibody · Hepatitis C virus · Infectious hepatitis · Liver fibrosis · Nonorgan-specific autoantibodies

Abbreviations

ALT alanine aminotransferase
AMA anti-mitochondrial antibody
ANA anti-nuclear antibody
Anti-HBc antibodies to HBV core antigen
Anti-HBs antibodies to HBV surface antigen
Anti-HCV antibodies to HCV
Anti-HEV antibodies to Hepatitis E virus
Anti-HGV antibodies to Hepatitis G Virus
Anti-LKM liver/kidney microsomal antibody
GPCA gastric parietal cell antibody
HBV hepatitis B virus
HCV hepatitis C virus
HGV-RNA hepatitis G Virus-RNA
HIV human immunodeficiency virus
NOSAs non-organ-specific autoantibodies
SMA anti-smooth muscle antibody
Introduction

HCV (hepatitis C virus) infection occurs less frequently in children than in adult patients. The natural history, prognosis and clinical significance of HCV infection are poorly defined in childhood [11]; in contrast, HCV infection in adults presents a high degree of chronicity, with up to 50% of all HCV-infected adults developing progressive liver disease [21]. Results from prospective studies show that 20–30% of chronically infected adults develop compensated and eventually decompensated cirrhosis or hepatocellular carcinoma, or both, within 20 years of the initial infection [1].

The lower prevalence of HCV infection in children and the fact that most patients undergo antiviral drugs treatment result in very limited knowledge of the natural outcome of chronic HCV infection acquired at early ages of life. Thus, immediate goals in the investigation of HCV infection should be to characterize current epidemiology and to describe the pathogenesis and course of hepatitis C in children and adults [20]. To understand the evolution of HCV infection could be itself an important surrogate endpoint for the evaluation of infected young patients with prognostic implications and therapeutic consequences.

The aim of this study was to determine the clinical features and long-term evolution of HCV infection in a group of children who had never received treatment with antiviral drugs.

Patients and methods

Thirty-seven children (16 females, 21 males) with positive antibodies to hepatitis C (anti-HCV) were investigated retrospectively. These patients were followed-up for a period of 5 years. None had received treatment with antiviral drugs for viral hepatitis or had a history of intravenous drug abuse. All subjects made regular visits to our outpatient clinic, and the serum levels of alanine aminotransferase (ALT), albumin, prothrombin time, anti-nuclear antibodies (ANA), anti-mitochondrial antibodies (AMA), anti-smooth muscle antibodies (SMA), liver-kidney anti-microsomal antibodies type I (LKM), anti-gastric parietal cells antibodies (GPCA), rheumatoid factor, thyroxine (T4), thyroid stimulating hormone (TSH), anti-thyroid antibodies, anti-HCV and HCV-RNA were determined at least on five consecutive occasions at 1-year intervals. Hepatitis B virus (HBV) surface antigen (HBsAg), antibodies to HBV surface antigen (anti-HBs), antibodies to HBV core antigen (Anti-HBc), human immunodeficiency virus (HIV) and antibodies to hepatitis E (anti-HEV) were investigated in all patients during one visit. Hepatitis G virus-RNA (HGV-RNA) and antibodies to HGV (anti-HGV) were determined in 26 patients. Genotypes of HCV were performed in 21 viremic children.

The duration of infection was calculated as the interval between the presumed date of infection and the date of the last visit to the clinic. Liver biopsies were obtained in 17 patients. Repeated biopsies were performed in three patients.

Viral markers for HBV, HCV and HIV were tested by third generation ELISA (Axysym; Abbott Diagnostics, Chicago, Ill.). Anti-HEV was detected by a commercially available ELISA (Bioelisa HEV IgG; Biokit, Barcelona, Spain). HGV-RNA and anti-HEV antibodies were detected using commercial tests (Roche Diagnostics, Mannheim, Germany). HCV-RNA was detected by PCR (AmpliCor HCV PCR test, Roche Diagnostics), and HCV genotyping was performed by a second generation line probe assay (INNO-LIPA HCV; Innogenetics, Ghent, Belgium).

Liver biopsies were performed percutaneously and under the ultrasound guidance of an experienced operator. All liver biopsy specimens (>12 mm long) were fixed in formalin, embedded in paraffin stained with hematoxylin-eosin and Masson’s trichrome and then observed by a single experienced pathologist (AM). Histological necroinflammatory activity and fibrosis were scored separately. The degree of histological necrosis with inflammatory activity was scored using the three parameters of the histological activity index (HAI) developed by Knodell et al. [12]; these consist of: (1) piecemeal necrosis and bridging necrosis (score: 0–10), (2) lobular cytolyisis (score: 0–4) and (3) portal inflammation (score: 0–4). Fibrosis was scored independently using the following system: F0, absence of fibrosis; F1, fibrous expansion of portal areas; F2, portal to portal bridging fibrous tracts; F3, portal-central bridging fibrous septa; F4, cirrhosis (bridging fibrous septa with parenchymal nodules). This score is a modification of the fibrosis scoring system of Knodell’s method and introduces one additional degree of fibrosis. Macrovesicular steatosis was graded semiquantitatively using a modified score from Brunt et al. [5]. Alternative and additional diagnoses identified in the liver biopsy were also recorded.

Results

Mean chronological age for the 37 patients (21 males and 16 females) at the last follow-up evaluation was 20.1 years (range: 9–30 years). Most of the subjects were diagnosed among patients who were screened for HCV infection after receiving transfusions of blood products for heart surgery, hematological diseases or casual injuries before 1992. Thus, most of the patients included in the study presented with other diseases that were unrelated to HCV infection: 16 patients had a congenital heart disease, seven had various hematological disorders, three had neurological diseases,
Table 1 Characteristics and biologic data of children with HCV infection

| Age (years) at the follow-up evaluationa | Patients negative for HCV-RNA (n=10) | Patients positive for HCV-RNA (n=27) |
|----------------------------------------|--------------------------------------|--------------------------------------|
| 18 (9–26)                              |                                      | 20 (9–30)                            |
| Duration of the HCV infection (years) in 36 patients at the follow-up evaluationa | 14 (6–23)                            | 16 (5–30)                            |

Route of infection
- Parenteral: 10
- Vertical: 0
- Unknown: 0

Alanine aminotransferase (U/l)
- Normal (n): 10
- ≤2 × UNL (n)b: 0
- 2–4 × UNL (n): 0
- >4 × UNL (n): 0

Non-organ specific antibodies
- Antinuclear antibody: 0
- Anti-mitochondrial antibodies: 1
- Anti-smooth muscle antibodies: 0
- Liver-kidney anti-microsomal antibody type 1: 1
- Gastric parietal cell antibodies: 0

HCV genotype
- 1a: 8
- 1b: 12
- 2a: 1
- Unknown: 6

HBV: 0
HIV: 1
HGV-RNA/Anti-HGV (n=26): 1/0
Anti-HEV: 0

Values are given as the mean, with the range presented in parenthesis.

One female patient presented persistent positive AMA for 3 years, which then normalized and remained negative during the subsequent 9-year follow-up period. Throughout this period the patient was asymptomatic, and liver tests were normal. Two years after the laboratory tests were negative for antibodies, the patient presented with psoriasis. Another patient had persistent high titers (1:620) of anti-LKM. Fifteen years after acquiring the infection this patient lost the anti-HCV but the anti-LKM remained positive for two more years. During the follow-up this patient did not present liver abnormalities or other diseases.

During the follow-up span of this study, another female patient became persistently anti-HCV negative 12 years after she was infected. At the time of writing, eight patients remain anti-HCV-positive and HCV-RNA-negative, and they show no evidence of liver disorders.

Group 2

HCV-RNA was detectable in the blood of 27 patients, two of whom became HCV-RNA-negative. The age of exposure to the infection and the type of exposure could not be established in one patient; following the detection of HCV in the blood serum, this patient was followed for 11 years and was positive for HCV up to and including the last follow-up visit.

All patients, with one exception, were asymptomatic. The exception had been diagnosed with leukemia in 1987 and received transfusions of blood products on several occasions. She underwent chemotherapy and had a cholestatic hepatitis; liver tests revealed persistent abnormalities. Anti-HCV and HCV-RNA were detected in 1991. The patient presented portal hypertension and progressive liver failure, and when she was 8 years old, in 1993, she received a liver transplant.

The evolution of serum ALT levels is shown in Table 1. Serum levels of albumin and prothrombin time were abnormal only in the patient who developed hepatic failure.

The presence of non-organ-specific autoantibodies (NOSAs) was detected in five patients. One patient (genotype 1a) diagnosed with a single ventricle and Fontan surgery developed persistent anti-LKM (titers 1:160–1:620) and the rheumatoid factor throughout the evolution of the disease. Liver biopsy could not be obtained in this patient. Another male patient (genotype 1b) had positive ANA for 2 years, which subsequently normalized. This patient had increased ALT, and the liver biopsy showed minimal necrosis with inflammatory changes (HAI: 2) and no fibrosis. A third patient (genotype 1b) had persistent PGCA during her follow-up period (10 years), with slightly elevated ALT levels, mild necrosis with inflammatory changes (HAI: 4) and minimal fibrosis (stage 1). A fourth patient (genotype 1b) with AST levels greater than fourfold the upper normal level presented SMA during the last 2 years of her follow-up (HAI: 6). The patient diagnosed...
with Turner syndrome and IgA deficiency (genotype 1b) developed throughout the years a seronegative polyarticular rheumatoid arthritis, psoriasis and celiac disease. She had intermittently positive ANA.

The levels of T4 and TSH were normal in all patients, and no patient developed antithyroid antibodies.

Abdominal ultrasounds did not show significant abnormalities. Only the patient who presented cirrhosis had changes associated to portal hypertension.

Liver biopsies were performed in viremic patients who presented altered hepatic function tests and for whom parental consent had been obtained. Seventeen patients underwent 20 liver biopsies. Mean evolution time of the infection at the time of biopsy was established in 16 patients (mean: 11.2 years; range: 3–21 years). The results on necroinflammatory activity and fibrosis are shown in Table 2.

A second hepatic biopsy was performed in three patients after 5, 12 and 14 years, respectively. The minimal necroinflammatory activity observed in the first biopsy specimens of these three children remained unchanged in the second biopsy. One of the patients progressed from F0 to F1 fibrosis. A second patient never presented with fibrosis, and both samples of the third patient had minimal fibrosis (F1).

During the follow-up period, 2/27 patients became persistently HCV-RNA-negative (for 9 and 10 years, respectively). Both patients showed minimal necrosis and inflammatory activity with minimal fibrosis (F1) in the biopsy performed before HCV-RNA clearance. One patient of Group 1 was HIV-positive, while one patient of Group 1 and six patients of Group 2 were HGV-positive.

Viral coinfections observed in both groups are expressed in Table 1.

### Discussion

In this study, chronic hepatitis C infection was silent in most of the children, and there was little biochemical evidence of liver disease in these patients. Only one girl with portal hypertension and liver failure developed symptoms related to the infection. Consequently, our data suggest that HCV infection may be currently underdiagnosed in children and, moreover, that young patients could become a potential source of infection. We therefore strongly recommend that children falling in risk groups be screened for HCV infection.

Of the 37 pediatric patients in this series, seven (19%) had NOSAs. These antibodies are highly prevalent in subjects exposed to the HCV, and a positive test result for SMA, GPCA and ANA is part of the natural course of chronic HCV infection in adults and children [7]. Lenzi et al. [13] reported a higher prevalence of NOSAs in anti-HCV-positive adult patients than in normal controls (25 vs. 6%). Stroffolini et al. [22] found positive NOSAs in 36.9% of 502 subjects with HCV-RNA-positive chronic hepatitis. Muratori et al. [16] detected positive NOSAs in 16 of 47 children (34%) with chronic hepatitis C, and anti-LKM was present in 11% of the children.

Of the 37 children participating in our study who manifested past or active HCV infection, two had serum anti-LKM. This incidence is lower than that observed by Muratori et al. (11%) [16] and Bortolotti et al. (10.3%) [4] in children

| Table 3 Hepatic fibrosis in biopsies from children with HCV infection |
|-------------------------|------------------|------------------|------------------|------------------|
|                         | Vogt et al. [23] | Hoshiyama et al. [9] | Guido et al. [8] | El-Raziky et al. [6] | This report |
| Number of patients      | 17               | 38               | 112              | 26              | 17          |
| Mean duration of the HCV infection (year) | 21.2±4.6 | 7.1±2.8 (n=23) | 12.9±3.1 (n=15) | 8.04±5.3 | Non-determined | 11.2±5.6 |
| Fibrosis                |                  |                  |                  |                  |             |
| No/low grade            | 14/2            | 23               | 15               | 25/81           | 20/4        | 7/9        |
| Severe/cirrhosis        | 0/1             | 0                | 0                | 5/1             | 1/1d        | 0/1e       |

*a Some patients were co-infected with HBV and/or had secondary iron overload
*b Two patients with congestive heart failure
*c Patient with Anti-HBs, anti-HBc and anti-HEV antibodies
*d This 12-year-old cirrhotic patient was not thalassaemic and HBV markers were negative
*e This leukemic patient was infected with HCV and was also receiving chemotherapy concurrently
with HCV infection. A possible explanation for this finding is the higher age and longer mean duration of HCV infection in the patients included in our series in comparison to the children in the other two studies. The prevalence reported by Lenzi et al. (1.3% [13]), Stroffolini et al. (2.25% [22]) and Reddy et al. (0% [18]) in adults support this hypothesis. Anti-LKM is an immunomarker of type 2 autoimmune hepatitis [2] that tends to present at younger ages and to affect mainly children. Our data suggest that age may be a factor contributing to the presence of anti-LKM1 in children infected with HCV. Moreover, we observed that one of our non-viremic patients had persistent antibodies after the test for anti-HCV became negative.

Liver specimens from the children with hepatitis C showed mild necroinflammatory changes and a low level of fibrosis. In most patients, fibrosis was absent or low grade. Only one patient, as mentioned above, developed cirrhosis and liver failure over a period of 6 years. This leukemic patient had other additional risk factors for liver damage that may have affected this outcome. In this respect, Hoshiyama et al. [9] observed that chronic hepatitis C is more frequent among children with hepatitis C infection following blood transfusions for malignant disease than in patients with hepatitis C following blood transfusions for open heart surgery.

We have compared our results with those obtained from other pediatric series [6, 8, 9, 23]. Table 3 summarizes the data on fibrosis obtained from liver biopsies of 210 pediatric patients (above-mentioned studies). Fibrosis was not detected or was low grade in 200 patients (95%), and severe fibrosis or cirrhosis was detected in ten patients (5%). Minola et al. [15] and Poynard et al. [17] demonstrated an inverse correlation between age at HCV infection and progression to cirrhosis that suggests that chronic HCV infection in childhood induces mild changes in the liver with a low level of fibrosis and a low rate of progression. However, it has been observed that mild chronic hepatitis C can be a progressive disease in adult patients [3, 19] and that the long evolution of infection acquired in childhood could increase the risk of fibrosis. Thus, some children infected early in life, in whom chronic disease has a mild liver expression, might develop fibrosis in adulthood.

It is noteworthy that 6/27 of the children with HCV infection in our series had a co-infection with HGV (HGV-RNA-positive) or had a past HGV infection (anti-HGV-positive). This prevalence of HGV infection is higher than that found in healthy children (6%) in Spain [10]. These data suggest that HCV-infected children must be considered to be a risk group for HGV infection since HCV and HGV are both parenterally transmitted. Moreover, no differences were observed in biochemical and histological findings in HCV-infected children with and without a concomitant infection with HGV, which suggests that superimposed HGV infection does not influence the course of HCV infection.

Conversely, our data do not support that HEV, which can be transmitted by a parenteral route [24], is prevalent in children who had undergone blood transfusions (prevalence rate: 0%). However, this low value may result from the low prevalence of HEV infection in Spain (2.8%) [14].

Conclusion

Most children chronically infected with HCV are asymptomatic and present only mild biochemical evidence of hepatic injury. Spontaneous clearing of the virus occurs occasionally. Autoantibodies are common in HCV patients. The natural history of chronic hepatitis C in children differs from that in adults since HCV infection is relatively benign, induces mild changes in the liver with a low level of fibrosis and a low rate of progression and is rarely associated with severe or decompensate liver disease.

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