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Letter to the Editor

Liver injury with autoimmune features after vaccination against SARS-CoV-2: The verdict is still open

A R T I C L E   I N F O

Keywords
Autoimmune hepatitis
Drug induced liver injury
SARS-CoV-2 vaccine

Dear Editor,

We read with interest recent reports describing acute hepatic injury following vaccination against SARS-CoV-2 [1]. The clinical expression of the hepatic involvement ranges from mild hepatitis to acute liver failure requiring liver transplantation [2], and biochemical, serological and histological features are typical of autoimmune hepatitis (AIH). We herein describe suspected SARS-CoV-2 vaccine-related liver injury with autoimmune features occurring in three young patients, in whom other causes of acute hepatitis, such as hepatitis A, B, C, E, drug induced liver injury, alcohol, have been excluded after accurate clinical, biochemical, and virological assessment. Detailed liver biopsy findings for each case are presented in Table 1.

The first patient is a 30-years-old woman with Hashimoto thyroiditis and family history of autoimmune diseases (ADs) who received two doses of the Moderna vaccine in September and October 2021. In November 2021 she complained of asthenia, hyporexia, arthralgia and dark urine. Alanine transaminase (ALT) levels were 60-fold upper limit of normal (ULN), total bilirubin was 33.5 µmol/L, IgG was 1642 mg/dL and antinuclear antibodies (ANAs) tested positive at 1:640 titer with homogeneous pattern, while anti-smooth muscle antibody (ASMA), anti-SLA/LP, anti-LKM and anti-LC1 tested negative. A liver biopsy showed periportal lympho-plasmocytic infiltrate with interface hepatitis, without significant fibrosis (Table 1). The patient received high-dose N-acetylcysteine, while she refused steroid treatment. ALT levels decreased to normal in 8 weeks; one month later they were still moderately elevated (up to 3-fold), then normalized again spontaneously. The second patient is a 21-years-old girl with no significant medical history. She performed routine blood tests on a regular basis and family history of autoimmune diseases (ADs) who received two doses of the Moderna vaccine in May 2021, with peak in November, with IgG 1749 mg/dL, total bilirubin 15.4 µmol/L. ANAs were positive at 1:640 titer with homogeneous pattern, while ASMA, anti-SLA/LP and anti-LKM were negative. Liver biopsy showed mild portal lymphoplasmacytic infiltrate with mild-moderate interface hepatitis, without significant fibrosis (Table 1). HLA was A1, B8, DR3. Immunosuppressive treatment was declined, and transaminase levels remained altered (around 3-fold) in the following months. The third patient is a 21-years-old girl with no significant medical history. She came to our attention in November 2021 for persistent 2-fold ALT increase since March 2021 found on routine blood test, before the first vaccinal cycle. In November ALT were 2-fold ULN, ASMA were positive at 1:320 titer, ANAs at 1:80 titer with homogeneous pattern, IgG were 1502 mg/dL, and liver biopsy showed perportal lympho-plasmocytic and lobular inflammatory infiltrate and focal piecemeal necrosis with moderate periportal fibrosis (Table 1). HLA was A1, B8, DR3. A diagnosis of genuine AIH was made, but therapy was declined due to patient hesitancy. In December, a few days after the booster dose (Moderna), she developed a hepatitis flare with ALT levels up to 12-fold ULN, so steroid therapy was initiated with rapid normalization of ALT. Retrospectively, she was noted to have a slight ALT increase (peak 4.7 x ULN) two weeks after the second dose of the Pfizer vaccine in May 2021, with subsequent return to a 2-fold ULN during following months.

AIH is a persistent, fluctuating inflammation of the liver observed in genetically predisposed subjects after exposure to an initiating, mostly unknown, factor [3]. Vaccines have been proposed as potential triggers [4,5]. The association of SARS-CoV-2 vaccination and AIH has been increasingly reported: a systematic review recently reported 32 patients [4,5]. The first patient is a 21-years-old girl with no significant medical history. She performed routine blood tests on a regular basis and family history of autoimmune diseases (ADs) who received two doses of the Moderna vaccine in May 2021, with peak in November, with IgG 1749 mg/dL, total bilirubin 15.4 µmol/L. ANAs were positive at 1:640 titer with homogeneous pattern, while ASMA, anti-SLA/LP and anti-LKM were negative. Liver biopsy showed mild portal lymphoplasmacytic infiltrate with mild-moderate interface hepatitis, without significant fibrosis (Table 1). HLA was A1, B8, DR3. Immunosuppressive treatment was declined, and transaminase levels remained altered (around 3-fold) in the following months. The third patient is a 21-years-old girl with no significant medical history. She came to our attention in November 2021 for persistent 2-fold ALT increase since March 2021 found on routine blood test, before the first vaccinal cycle. In November ALT were 2-fold ULN, ASMA were positive at 1:320 titer, ANAs at 1:80 titer with homogeneous pattern, IgG were 1502 mg/dL, and liver biopsy showed perportal lympho-plasmocytic and lobular inflammatory infiltrate and focal piecemeal necrosis with moderate periportal fibrosis (Table 1). HLA was A1, B8, DR3. A diagnosis of genuine AIH was made, but therapy was declined due to patient hesitancy. In December, a few days after the booster dose (Moderna), she developed a hepatitis flare with ALT levels up to 12-fold ULN, so steroid therapy was initiated with rapid normalization of ALT. Retrospectively, she was noted to have a slight ALT increase (peak 4.7 x ULN) two weeks after the second dose of the Pfizer vaccine in May 2021, with subsequent return to a 2-fold ULN during following months.

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In Supplementary material we reviewed the available case reports of AIH like liver injury until June 2022. The search was performed using PubMed, entering as keywords ‘COVID vaccine’, ‘SARS-CoV-2 vaccine’, “Autoimmune hepatitis” and “Liver injury”, last access in June 2022. A
agree on the rarity of the phenomenon and its generally favorable
described their features and outcomes [2]. All the available evidence
course was favorable in almost all cases, even if in three reports (6%) the
case leading to hepatitis relapse requiring oral steroids. The clinical
Cases associated with azathioprine (<6 months from the last dose, IQR 7-26) and a high proportion of patients
previously altered ALT levels and liver fibrosis indicated a long-standing
after the Moderna booster a striking increase of ALT was observed,
early recognition and to provide guidance for adequate management.
outcome and emphasize the need for increased awareness, to promote
funic central nervous system infection.
derwent to liver transplantation. One patient died of suspected oppor-
tinality [10] .

total of 31 reports was identified, covering a total of 52 cases of AIH-like
liver injury after COVID-19 vaccination. The median age is 61 years
(IQR 41-71), with a prevalence of female sex (N=35/52, 67%). Most
relevant comorbidities are autoimmune diseases (ADs) (N=11/52, 21%) and
dliver diseases (N=12/52, 23%). Most of the reports involve mRNA
vaccines (N=43/52, 83%). Onset time is generally quite short (median
15 days from the last dose, IQR 7-26) and a high proportion of patients
present an acute hepatitis with jaundice (N=43/52, 40%). Autoantib-
odies useful for the diagnosis of AIH were often found (N=35/45, 78%,
of which ANAs 31/45, 69%, ASMA in 11/45,24%, and anti-SLA/LP and
other ADs reactivation after SARS-CoV-2 vaccination [8] : this mecha-
nism has been postulated since the early presentation with detectable
pathogenic autoantibodies [8] . Other proposed mechanisms are
bystander activation and epitope spreading [6] as well as molecular
mimicry [7] . Interestingly, some reports suggest the possibility of
SARS-CoV-2 Spike protein expression within hepatocytes after vacci-
nation [9] and the presence of an immune infiltrate characterized by
activated cytotoxic CD8 T-cells with SARS-CoV-2 Spike-protein speci-
ificity [10].

Whether these cases represent genuine AIH triggered by vaccine or
toxic vaccine-induced liver injury is at present a matter of debate,
which only a longer follow up could figure out: recurrent flares of ALT or
persistence of ALT and IgG alterations will suggest typical AIH and need for
prolonged immunosuppression, whereas spontaneous and complete
biochemical and histological resolution will indicate vaccine-induced
liver injury. Cases of genuine AIH reactivation after vaccination em-
phasize the need for close follow-up after vaccination, particularly when
the immunosuppressive treatment has not been introduced yet. As
of today, we are unable to prove the direct role of vaccination in the
induction of hepatic damage, but we cannot disprove it either. Rigorous
population-based studies and active pharmacovigilance are urgently
needed to assess beyond reasonable doubt incidence and clinical sig-
nificance of such observations. Until then, the final verdict should
remain open.

| Case I (11 portal spaces) | Case II (12 portal spaces) | Case III (12 portal spaces) |
|---------------------------|---------------------------|---------------------------|
| Periportal or perisepal interface hepatitis | Severe (continuous around > 50% of portal areas). Moderate plasma cells infiltrate | Mild/moderate (focal, most portal areas). Scattered plasma cells and eosinophils |
| Confluent necrosis | Zone 3 necrosis in some areas | Absent |
| Focal (spotty) lytic necrosis, apoptosis and focal inflammation | Five to ten foci per 10X objective, perivenular necroinflammatory infiltrate | Two to four foci per 10X objective, perivenular necroinflammatory infiltrate |
| Portal inflammation | Marked, all portal areas | Mild, focal inflammation |
| Fibrosis | No fibrosis | Fibrous expansion of some portal areas, without septa |
| Other features | Mild bile duct injury | Focal ductular hyperplasia, no rosettes |
| | | Mild biliary regression, focal rosettes |

Table 1
Liver biopsy findings among three patients.
Data are presented according to Ishak grading and staging.

This data are widely consistent with a recent retrospective study
(ahead of print), which collected data from 18 countries on 87 cases and
described their features and outcomes [2]. All the available evidence
agree on the rarity of the phenomenon and its generally favorable
outcome and emphasize the need for increased awareness, to promote
early recognition and to provide guidance for adequate management.

Among our patients, the first two had stringent temporal correlation
between vaccine and hepatic injury and a previously healthy liver has
been documented, so direct causal correlation should be suspected.
Interestingly, the kinetics of biochemical hepatic damage was quite
different (graphic presentation in Supplementary material): hyperacute and self-limiting for the first patient, less acute but persistently
abnormal and fluctuating for the second one. For the third patient,
alter liver function tests were present before vaccination, but only
after the Moderna booster a striking increase of ALT was observed,
which was rapidly controlled with steroid treatment. In the latter case,
previously altered ALT levels and liver fibrosis indicated a long-standing
process, pointing to pre-existing AIH. Of note, as well as this “genuine
AIH”, even the first two patients had a preexistent AD, highlighting
predisposition to AIH-like liver injury after vaccination. Efe et al. re-
ported that 28% of the AIH-like-liver-injury patients had been diagnosed
with other AD before liver injury onset, of which most common are
autoimmune thyroiditis (14%), inflammatory bowel diseases (3%) and
sarcoidosis (3%) [2]. The significant proportion of patients with other
ADs developing AIH-like liver injury suggest genetic predisposition,
such as impaired clearance of nucleic acids, TLR polymorphism, HLA
haplotype [7]. Various authors propose that vaccination may not
generate new ADs, but rather triggers long-lasting latent autoimmunity
[6,7]. The strong inflammatory response induced by pattern recognition
receptors could act like a relapse trigger of a latent AIH, as shown for
other ADs reactivation after SARS-CoV-2 vaccination [8]: this mecha-
nism has been postulated since the early presentation with detectable
pathogenic autoantibodies [8] . Other proposed mechanisms are
bystander activation and epitope spreading [6] as well as molecular
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Authors contributions

MF co-wrote the original manuscript and edited the final submission; ML assisted with conceptualization and edited the final submission; LM provided patient care, co-wrote the original manuscript and edited the final submission (Fig. 1).

Declaration of Competing Interest

No conflict of interest to declare

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