Immunohistochemical Detection of HCV Proteins in Liver Tissue

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Abstract

Detection and localization of HCV in liver tissue are vital for diagnostic purposes and clinical management of HCV-infected patients, as well as for the elucidation of viropathological mechanisms. The fragility of HCV RNA and the low levels of viral expression in infected tissues are a constant limitation in molecular assays for HCV characterization. HCV antigen detection, by immunohistochemistry, in liver biopsies is an attractive option for precise localization and quantification of viral proteins with direct access to histological patterns. We describe here a study using a novel immunohistochemical method effective on fixed, archived specimens, including liver biopsies and surgical resection samples. The initial protocol uses a biotin-detection system but can also be used in a polymer-detection system. This protocol offers easy, precise, and strong staining resolution with distinct patterns consistent with the liver pathology, irrespective of the viral HCV genotype examined. This approach provides applications for diagnosis as well as for exploratory pathological studies.

Key words: HCV, immunohistochemistry, envelope protein, chronic infection, hepatocellular carcinoma.

1. Introduction

The HCV genome is a single-stranded RNA molecule of about 9600 bp. It encodes for a large polyprotein that is processed by host and virus proteases into several structural and nonstructural viral proteins (Envelope 1/2, Capsid, p7and NS2, NS3, NS4A/B, and NS5A/B).

The detection of HCV replicative intermediates or virus antigen may be helpful for diagnosis or clinical management of
patients with HCV infection, and it is of crucial importance for monitoring patients before and after HCV-related liver transplantation. HCV replication level generally seems to be relatively low in infected liver, hampering the detection of HCV particles directly in the liver (1, 2). Detection methods based on HCV RNA amplification, like in situ PCR or in situ hybridization, do not always detect HCV and can lead to conflicting results concerning localization of viral particles (3), but they remain an interesting tool when used to complement classical methods (4, 5). These methods are probably limited by rapid RNA degradation in tissues and the difficulty of designing efficient probes that overcome the high variability of HCV genomes.

Detection of HCV antigens by immunochemistry in liver biopsies is therefore an interesting option that allows both localization and quantification of viral proteins.

Quite a few antibodies have been raised against hepatitis C antigens and fulfill the conditions for use in immunochemistry (IHC). So far, except for the promising commercial TORDJI 22 and TORDJI 32, no commercial antibody allows specific, reproducible, and efficient staining (6–8), one reason explaining why IHC has failed to become established in routine experiment in diagnostic labs. Among those tested for IHC are antibodies raised against nonstructural proteins (NS3, NS4, and NS5) (9, 10) and core or envelope protein (10, 11).

Antibodies raised against envelope protein E2 provided the best efficiency for IHC techniques (11, 12). Even though the region encoding envelope protein encompasses the hypervariable region I (HVR 1) of HCV, the overall conformation of E2 seems to be quite conserved (13). In our experience (12), anti-E2 antibody D4.12.9 detected the E2 protein from all genotypes tested, including genotypes 1 to 5, indicating that this antibody is suitable for HCV detection irrespective of its precise genotype. In our personal experience (12), the staining location for anti-E2 immunostaining is mainly cytoplasmic, although occasional perimembrane staining (including cytoplasmic and nuclear membrane) occurs. Staining pattern is mainly coarse granular with a microvesicular pattern (see Fig. 3.1).

HCV staining patterns appear to differ slightly according to the pathological status of the liver tissue. We observed a very strong staining of hepatocyte membrane, cytoplasm, and perinuclear regions in liver from patients with active HCV-related cirrhosis (Intense plasma and nuclear membrane staining was observed in cases with high inflammatory activity.) In noncirrhotic and nontumoral tissues, anti-E2 staining intensity increased with hepatitis fibrosis state. In HCV-related tumors, staining was exclusively detected within regeneration nodules and confined to hepatocytes whose morphology remained unchanged. Staining appeared in one of two distinct patterns: trabecular throughout the hepatic parenchyma or only in isolated cells.
2. Materials

1. Normal source of samples: Transparietal biopsies or surgical biopsies.
2. Nature of samples: Frozen or formalin-fixed human liver.
3. Xylene dilutions in water.
4. Absolute ethanol dilutions in water: 95%, 70%.
5. Methanol solution (0.3% hydrogen peroxide).
6. Vector antigen unmasking solution (Vector Laboratories, Burlingame, CA).
7. Phosphate-buffered saline (PBS) solution (Sigma).
8. Skimmed milk.
9. Bovine serum albumin (BSA, Sigma).
10. Primary mAb, D₄₁₂₉ at 0.2 μg/mL.
11. Secondary antibody, Vectastain® ELITE ABC PEROXY-
DASE KIT Rabbit IGG (Vector).
12. DAB Substrate kit (Vector).
14. Mounting medium (DAKO Mounting medium, Dako,
North America).
15. Slide racks and trays (Fisher Scientific).
16. Microwave oven.
17. Microscope cover glass.
18. Light microscope.

3. Method

1. Deparaffinize tissue (see Note 1) sections in xylene for 10 min
(twice).
2. Rehydrate the tissue in graded ethanol concentrations (100%,
95%, and 70% for 5 min each) and proceed immediately to
step 3.
3. Block endogenous peroxidase activities by incubation in
methanol solution for 30 min at room temperature (see
Note 2).
4. For unmasking, place the slides in a microwave oven for
15 min in antigen unmasking solution. Set the microwave
power high enough to bring the solution to a boil, and then
reduce power so that the solution continues boiling for the
required time. Allow the slides to cool down for 30 min in
the same unmasking solution (see Note 3).
5. Incubate with PBS solution (5% skimmed milk and 0.1%
BSA) for 1 h at room temperature. Do not rinse.
6. Incubate sections overnight with the primary mAb at 4°C.
Sections without primary antibody can be used as controls.
7. Wash in PBS three times, 5 min each.
8. Incubate for 30 min with secondary antibody at 5 μl/mL in
PBS solution (0.1% BSA).
9. Wash in PBS three times, 5 min each.
10. Amplify signal at 37°C for 45 min (Vectastain peroxydase kit);
use PBS solution (0.1% BSA).
11. Wash in PBS three times, 5 min each.
12. Incubate tissue sections with the DAB substrate at room tem-
perature for 5 min,
13. Wash in PBS three times, 5 min each.
14. Counterstain the sections with Mayer’s hematoxylin.
15. Dehydrate the section through successive ethanol baths (70% ethanol, 95% ethanol, absolute ethanol, and xylene, 5 min in each solution).


4. Notes

1. Sensitivity of detection is significantly increased by use of fresh-frozen tissue.

2. Alternatively, a method based on polymer detection can be used with D₄₁₂₉ in IHC. It generally offers same sensitivity and a slightly better resolution because it eliminates endogenous biotin interference, the main source of nonspecific background staining. The method differs from the classic protocol in step 3, where we incubate 10 min in Dako peroxidase block solution (DakoCytomation EnVision + Dual Link System Peroxidase, Dako, France) and rinse gently with distilled water. For step 8, apply peroxidase-labeled polymer (DakoCytomation EnVision + Dual Link System Peroxidase) to cover specimen. Incubate for 30 min and rinse the slides with the buffer solution provided. Eventually, apply substrate-chromogen solution (DakoCytomation EnVision + Dual Link System Peroxidase) to cover the specimen, add DAB, and incubate for 2–10 min (optimal incubation time can be determined by verifying signal intensity under the microscope). Rinse gently with distilled water from a wash bottle (do not focus flow directly on tissue). Proceed with step 13.

3. During microwave treatment, ensure that solution level is sufficient to cover the tissue section throughout the treatment. Check solution level every 3–5 min.

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