Hypervascular nodule in a fibrotic liver overloaded with iron: identification of a premalignant area with preserved liver architecture.
António Sá Cunha, Jean-Frédéric Blanc, Hervé Trillaud, Victor de Ledinghen, Charles Balabaud, Paulette Bioulac-Sage

To cite this version:
António Sá Cunha, Jean-Frédéric Blanc, Hervé Trillaud, Victor de Ledinghen, Charles Balabaud, et al.. Hypervascular nodule in a fibrotic liver overloaded with iron: identification of a premalignant area with preserved liver architecture.. Comparative Hepatology, BioMed Central, 2005, 4, pp.5. 10.1186/1476-5926-4-5. inserm-00090595

HAL Id: inserm-00090595
https://www.hal.inserm.fr/inserm-00090595
Submitted on 1 Sep 2006

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Hypervascular nodule in a fibrotic liver overloaded with iron: identification of a premalignant area with preserved liver architecture

António Sá Cunha, Jean-Frédéric Blanc, Hervé Trillaud, Victor De Ledinghen, Charles Balabaud* and Paulette Bioulac-Sage

Background: The presence of a hypervascular nodule in a patient with liver disease is highly suggestive of a hepatocellular carcinoma [1]. Increased iron stores in patients with HCC developed on a non-cirrhotic liver is well documented [2-5]; iron stores are seldom depleted at the time of the discovery of the HCC [6]. Few cases of premalignant nodules associated with HCC have also been reported under these circumstances [7,8]. In a fibrotic liver overloaded with iron, we report a case of a hypervascular and hyposiderotic nodule with premalignant features, but with a normal architecture.

Case presentation

General data
A 55 year old man with idiopathic refractory anaemia was addressed to our Unit for the cure of a recently appeared 3.3 cm hypervascular liver nodule in segment II (November, 2003). Physical examination was normal, including BMI. Liver function tests were as follows: ASAT = 53 IU/L (Normal = 40); ALAT = 58 IU/L (N = 50); Bilirubin = 44

Abstract

Background: The presence of a hypervascular nodule in a patient with cirrhosis is highly suggestive of a hepatocellular carcinoma.

Case presentation: A 55 year old man with idiopathic refractory anaemia was addressed for the cure of a recently appeared 3.3 cm hypervascular liver nodule. The nodule was not visible on the resected fresh specimen, but a paler zone was seen after formalin fixation. The surrounding liver was fibrotic (METAVIR score F3) and overloaded with iron. However, the paler zone, thought to be the nodule, had in fact a normal architecture, was less fibrotic, and contained some "portal tract-like structures" (but with arteries only); moreover, this paler area was devoid of iron, contained less glycogen and was characterized by foci of clear hepatocytes.

Conclusion: In spite of the absence of architectural distortion, and a normal proliferative index, the possibility of premalignancy or malignancy should be considered in this type of hypervascular and hyposiderotic nodule, occurring in the context of an iron overloaded liver.
µmol/L (N = 17); PT = 70% (N = 70–100); V = 65% (N = 70–100); RBC = 2.9 × 10^6 cells/μl; Ht = 22.5% and Hb = 7.3 gm/dl; WBC = 4.6 × 10^3 cells/μl; and platelets = 210 × 10^9 /l. Ferritinemia was 1891 ng/l (N < 300), transferrin saturation was 100% (N < 40), iron concentration was 290 µmol/g (assessed by MRI, N < 36). AFP in blood was within the normal range. The patient, of Italian origin, was C282 Y -/-, H63D +/-, and S65C -/-, with no family history of iron overload. Markers for viral and autoimmune diseases were negative. Blood glucose was normal. He used to smoke 30 cigarettes per day; but had stopped for the last 8 years. He drank alcohol only occasionally.

The treatment of his refractory anaemia consisted in blood transfusion (total of 10 packs), Desferoxamine, and Deferiprone.

**Imaging results**

Ultrasound examination showed a hypoechoic ovoid nodule (3.2 × 1.9 cm) in segment II. MRI showed a hypointense non-tumoral liver on T1- and T2-weighted images due to iron overload. In comparison, the nodule was hyperintense on T1- and T2-weighted images. After gadolinium injection, the nodule was hyperintense on T1-weighted images and remained so in the portal phase (Fig. 1).

*Figure 1*

Hyperintensity of the nodule on T1- and T2-weighted images (left), which remains so after gadolinium injection (right), on T1-weighted image in the portal phase.
Liver pathology

On December 2003, a left lobe heptectomy was performed laparoscopically. Follow-up was uneventful. The resected specimen was carefully sliced but no nodule was found on the fresh specimen, and in the expected area. However, after formalin fixation, a 2 cm in diameter paler area was identified (Fig. 2, arrow). All slices were routinely processed. The following stainings were performed: H&E, trichrome, Perls, reticulin, PAS, and several immunostains (CD34, cytokeratins 7 and 19, CRBP1 and α-SMA; the latter for identification of quiescent and/or activated hepatic stellate cells [9]). The liver was fibrotic (METAVIR score F3) (Figs. 3a, 4a) with an iron overload 3 + (according to Searle score), mainly in hepatocytes of zones 1 and 2 (Fig. 5a). Liver iron concentration was 286 μmol/g (n < 36), and the iron concentration / age ratio was 5.1. Small foci of clear cells devoid of iron were also observed (not shown).

The paler zone, poorly limited from the adjacent parenchyma, was strikingly different. The architecture was preserved but the area was far less fibrotic (METAVIR score F1; Figs. 3b, 4b), with less iron (Fig. 5b), less glycogen (Fig. 6a), and with foci of clear hepatocytes (Figs. 6a, 6b). In these clear areas, hepatocytes were slightly bigger and occasionally displayed in two cell-thick plates. In these areas, as elsewhere, reticulin network, as well as Ki-67 (Mib-1) and CD34 were normal (not shown). One of the

---

**Figure 2**
Formalin fixed specimen: a flat and slightly clearer area is visible in the expected zone (arrow).
Figure 3
Septal fibrosis in non tumoral liver (a), contrasting with absence of fibrosis in the nodule (b). Masson’s trichrome.
Figure 4
Septal fibrosis in non tumoral liver (a), contrasting with absence of fibrosis in the nodule (b). Reticulin staining.
Figure 5
Iron overload in non tumoral liver (a), contrasting with less iron in the nodule (b) Perls staining.
Figure 6
PAS staining: (a) foci of clear hepatocytes (arrow) close to the border between the non tumoral PAS positive zone, on the left side, and the PAS negative nodule on the right side of the photograph; (b) a clear focus in the nodule.
most striking findings was the presence of different types of portal tracts: some were normal (Fig. 7), whereas others contained mainly ductules (Fig. 8) and others arteries (Fig. 9). Regarding the number of CRBP1 and α-SMA positive cells, no obvious differences were seen between the fibrotic and non-fibrotic parts of the liver. CRBP1 positive cells seemed to contain few lipid droplets.

Discussion
The mechanism accounting for the major hepatic iron overload is possibly multifactorial, including refractory sideroblastic anemia and blood transfusion, although the patient received only a limited number (<10) of blood transfusions. An associated hereditary iron overload such as transferrin receptor 2 haemochromatosis in this Italian patient cannot be ruled out.

Premalignant lesions have previously been described in iron overloaded patients in the absence of cirrhosis, although these lesions were discovered in the clinical context of a HCC [7,8]. To our knowledge, this is the first reported case of a premalignant area mimicking by imaging a HCC, but exhibiting microscopically a still well-preserved architecture, in an otherwise fibrotic liver.

The hyperarterialized nodule did not correspond to a macroscopically visible nodule, but rather to an ill-defined area with preserved architectural organization.
However, this area was considered as premalignant based on the following arguments: arterial hypervascularization with isolated arteries in the parenchyma [3]; loss of iron [10], and of glycogen; and presence of clear hepatocytes foci [11]. The diagnosis of a focal nodular hyperplasia (FNH)-like nodule as described in cirrhosis [12-14], particularly related to alcohol, seems unlikely due to the loss of iron and to the presence of clear hepatocytes foci. Nonetheless, that diagnosis cannot be ruled out, and should always be kept in mind, especially if a liver transplantation is foreseen. Recently, it has been reported that coexisting iron overload could significantly worsen the course of FNH [15]. Unfortunately, and because no frozen material of the lesion (which was not visible) was available, in this case no specific molecular studies could be carried out to settle that important issue.

Our case strengthens previous observations [7,8] showing that malignancy can overrun cirrhosis in iron overload (Fig. 10). A minor degree (stage) of fibrosis in areas devoid of iron could be a direct consequence of iron loss (less toxicity) and/or related to the malignant process [16,17] in its early phase (as denoted by absence of cellular disorganization, and negativity of the MIB-1 immunostaining). It was concluded that this patient needs a strict surveillance because he may be at risk of recurrence. Indeed, clear hepatocytes foci devoid of iron were also observed outside the nodule [10].

**Figure 8**
On the left side, ductular reaction around a portal tract in the nodule. On the right side, in another portal tract the bile duct is visible but not the portal vein and the artery. CK7 immunostaining.
Conclusion
The possibility of premalignancy or malignancy should be considered even in the absence of cirrhosis, when a nodule is observed in a patient with past or present liver iron overload.

List of abbreviations used
AFP – α-fetoprotein; ALAT – alanine aminotransferase; ASAT – aspartate aminotransferase; BMI – body-mass index; CRBP1 – cellular retinol-binding protein 1; FNH – focal nodular hyperplasia; Hb – hemoglobin; Ht – hematocrit; HCC – hepatocellular carcinoma; MRI – magnetic resonance imaging; PAS – periodic acid Schiff; PT – pro-trombine time; RBC – red blood cells; α-SMA – α-smooth muscle actin; WBC – white blood cells.

Authors’ contributions
A Sá Cunha performed the surgery. JF Blanc collected the references and contributed to the writing. H Trillaud reviewed the MRI. V De Ledinghen, hepatologist, was in charge of the patient. C Balabaud wrote the paper. P Bioulac-Sage interpreted the liver histology and contributed to the writing. All authors read and approved the final manuscript.

Figure 9
An unpaired artery in the nodule. α-SMA immunostaining.
**Iron overload**

- **Cirrhosis**
  - may not occur if iron is depleted at an early stage of the disease

- **Fibrosis**
  - Resistant iron lobular area
  - Proneoplastic nodule
  - HCC

- **Iron free foci**

**Pathways leading to HCC in iron overload:**
- **(a) classical pathway:**
  - Resistant iron lobular area
  - Proneoplastic nodule
  - HCC

- **(b) alternate pathway (rarely observed):**
  - Resistant iron lobular area
  - Proneoplastic nodule
  - HCC

**References**

1. Taouli B, Losada M, Holland A, Krinsky G: Magnetic resonance imaging of hepatocellular carcinoma. *Gastroenterology* 2004, 127:S14-S2.
2. Turlin B, Juguet F, Moirand R, Le Quilliec D, Campion JP, Lemoine B, Ramee MP, Brissot P, Deugnier Y: Increased liver iron stores in patients with hepatocellular carcinoma developed on a noncirrhotic liver. *Hepatology* 1995, 22:446-450.
3. Kowdley KV: Iron, hemochromatosis, and hepatocellular carcinoma. *Gastroenterology* 2004, 127:S79-S86.
4. Bioulac-Sage P, Le Bail BL, Winnock M, Balabaud C, Bemard C, Blanc JF, Saric J: Occurrence of hepatocellular carcinoma in nonfibrotic livers. *Hepatology* 2000, 32:1411-1412.
5. Blanc JF, De Ledinghen V, Bernard PH, de Verneuil H, Winnock M, Le Bail B, Carles J, Saric J, Balabaud C, Bioulac-Sage P: Increased incidence of HFE C282Y mutations in patients with iron overload and hepatocellular carcinoma developed in noncirrhotic liver. *J Hepatol* 2000, 32:805-811.
6. Pellise M, Gonzalez-Abad R, Navasa M, Miquel R, Bruguera M: Hepatocellular carcinoma in a patient with hereditary hemochromatosis without cirrhosis). *Gastroenterol Hepatol* 2001, 24:132-134.
7. Blanc JF, De Ledinghen V, Trimoulet P, Le Bail B, Bernard PH, Saric J, Balabaud C, Bioulac-Sage P: Premalignant lesions and hepatocellular carcinoma in a non-cirrhotic alcoholic patient with iron overload and normal transferrin saturation. *J Hepatol* 1999, 30:325-329.
8. Attia A, Blanc JF, Saric J, Balabaud C, Bioulac-Sage P: Premalignant lesions and hepatocellular carcinoma on non cirrhotic liver overloaded with iron). *Gastroenterol Clin Biol* 2000, 24:955-959.
9. Lepreux S, Bioulac-Sage P, Gabbiani G, Sapin V, Houssot C, Roncin B, Balabaud C, Desmouliere A: Cellular retinol-binding protein-I expression in normal and fibrotic/cirrhotic human liver: different patterns of expression in hepatic stellate cells and (myo)fibroblast subpopulations. *J Hepatol* 2004, 40:774-780.
10. Deugnier YM, Charalambous P, Le Quilliec D, Turlin B, Searle J, Brissot P, Powell LW, Halliday JW: Preneoplastic significance of hepatic iron-free foci in genetic hemochromatosis: a study of 185 patients. *Hepatology* 1993, 18:1363-1369.
11. Su Q, Bannasch P: Relevance of hepatic preneoplasia for human hepatocarcinogenesis. *Toxicol Pathol* 2003, 31:126-133.
12. Libbrecht L, Bielen D, Verslype C, Vanbeekvoort D, Piremme J, Nevens F, Desmet V, Roskams T: Focal lesions in cirrhotic explant livers: pathological evaluation and accuracy of pretransplantation imaging examinations. *Liver Transpl* 2002, 8:749-761.
13. Nakashima O, Kurogi M, Yamaguchi R, Miyaaki H, Fujimoto M, Yano H, Kumabe T, Hayabuchi N, Hisatomy J, Sata M, Koijiro M: Unique hypervascular nodules in alcoholic liver cirrhosis: identical to focal nodular hyperplasia-like nodules? *J Hepatol* 2004, 41:992-998.
14. Quaglia A, Tibballs J, Grasso A, Prasad N, Oznoz P, Davies SE, Burroughs AK, Watsonson A, Dhillon AP: Focal nodular hyperplasia-like areas in cirrhosis. *Histopathology* 2003, 42:14-21.
15. Hohler T, Lohe AW, Schirmacher P: Progressive focal nodular hyperplasia of the liver in a patient with genetic hemochromatosis—growth promotion by iron overload? *Dig Dis Sci* 2000, 45:587-590.
16. Theise ND, Lapook JD, Thung SN: A macroregenerative nodule containing multiple foci of hepatocellular carcinoma in a noncirrhotic liver. *Hepatology* 1993, 17:993-996.
17. Park YN, Yang CP, Cubukcu O, Thung SN, Theise ND: Hepatic stellate cell activation in dysplastic nodules: evidence for an alternate hypothesis concerning human hepatocarcinogenesis.* Liver* 1997, 17:271-274.

**Publish with BioMed Central and every scientist can read your work free of charge**

"BioMed Central will be the most significant development for disseminating the results of biomedical research in our lifetime."  
Sir Paul Nurse, Cancer Research UK

Your research papers will be:
- available free of charge to the entire biomedical community
- peer reviewed and published immediately upon acceptance
- cited in PubMed and archived on PubMed Central
- yours — you keep the copyright

Submit your manuscript here:
http://www.biomedcentral.com/info/publishing_adv.asp