Long term follow-up and outcome of liver transplantation from hepatitis B surface antigen positive donors

Roberto Ballarin, Alessandro Cucchetti, Francesco Paolo Russo, Paolo Magistri, Matteo Cescon, Umberto Cillo, Patrizia Burra, Antonio Daniele Pinna, Fabrizio Di Benedetto

Roberto Ballarin, Paolo Magistri, Fabrizio Di Benedetto, Hepatopancreatobiliary Surgery and Liver Transplant Unit, University Hospital “Policlinico”, University of Modena and Reggio Emilia, 41124 Modena, Italy

Alessandro Cucchetti, Matteo Cescon, Antonio Daniele Pinna, Department of Medical and Surgical Sciences, DIMEC, S. Orsola-Malpighi Hospital, Alma Mater Studiorum, University of Bologna, 40138 Bologna, Italy

Francesco Paolo Russo, Umberto Cillo, Patrizia Burra, Gastroenterology/Multivisceral Transplant Unit, Department of Surgery, Oncology and Gastroenterology, Padua University Hospital, 35128 Padua, Italy

Paolo Magistri, Department of Medical and Surgical Sciences and Translational Medicine, Sapienza, University of Rome, 00185 Rome, Italy

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Correspondence to: Roberto Ballarin, MD, PhD, Hepatopancreatobiliary Surgery and Liver Transplant Unit, University Hospital “Policlinico”, University of Modena and Reggio Emilia, Via del Pozzo 71, 41124 Modena, Italy. ballarinroberto@hotmail.com 
Telephone: +39-59-4224740

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Abstract

Liver transplant for hepatitis B virus (HBV) currently yields excellent outcomes: it allows to rescue patients with an HBV-related advanced liver disease, resulting in a demographical modification of the waiting list for liver transplant. In an age of patient-tailored treatments, in liver transplantation as well the aim is to offer the best suitable graft to the patient who can benefit from it, also expanding the criteria for organ acceptance and allocation. With the intent of developing strategies to increase the donor pool, we set-up a multicenter study involving 3 Liver Transplant Centers in Italy: patients undergoing liver transplantation between March 03, 2004, and May 21, 2010, were retrospectively evaluated. 1408 patients underwent liver transplantation during the study period, 28 (2%) received the graft from hepatitis B surface antigen positive (HBsAg)-positive deceased donors. The average follow-up after liver transplantation was 63.7 mo [range: 0.1-119.4; SD ± 35.8]. None Primary non-function, re-liver transplantation, early or late hepatic artery thrombosis occurred. The 1-, 3- and 5-year graft and patient survival resulted of 85.7%, 82.1%, 78.4%. Our results suggest that the use of HBsAg-positive donors liver grafts is feasible, since HBV can be controlled without affecting graft stability. However, the selection of grafts and the postoperative antiviral therapy should be managed appropriately.
Outcome of LT from HbsAg donors

Key words: Liver transplantation; Hepatitis B virus; Hepatitis B surface antigen; Hepatocellular carcinoma; Organ allocation; Organ procurement; Multicenter study

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Core tip: With the intent of developing strategies to increase the donor pool, we set up a multicenter study involving 3 Liver Transplant Centers in Italy between March 2004 and May 2010. 1408 patients underwent liver transplantation during the study period, and 28 received the graft from hepatitis B surface antigen positive (HBSAg)-positive deceased donors. None primary non-function, re-liver transplantation, early or late hepatic artery thrombosis occurred. Our results show that transplantation of grafts from deceased HBSAg positive donors is feasible and this represents a way to expand the donor pool, especially in the high-endemic areas where a large proportion of patients are highly viremic and HBeAg positive.

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INTRODUCTION

Epidemiology of hepatitis B and hepatocellular carcinoma

Hepatitis B virus (HBV) prevalence is different from a geographical region to another (Figure 1): currently, in Northern Europe, United States, Canada and Australia it ranges from 0.1% to 2%, while in central and Eastern Europe, as well as in Mid East, India, Central and Southern America, it is between 3% and 7%. Finally, the highest incidence, ranging from 10% to 20%, is registered in Africa and Easter Countries.

Notably, the incidence of hepatocellular carcinoma (HCC) in the same regions mirrors the prevalence of HBV. In Europe, Japan and North America HBV is responsible for 10%-15% of HCC cases, while conversely, in Asia and Africa, HBV is associated to 70% of cases. According to several studies, the relative risk of developing a tumor is close to 100-fold in HBV carriers vs non-carriers[1].

Liver transplantation for HBV

Liver transplant for HBV currently yields excellent outcomes, but in 1983, before the introduction of HBV immune globulin (HBIG) and antiviral therapy, a United States National Institute of Health consensus conference recommended against transplant for HBV because of the poor outcomes from severe recurrent liver disease. The first studies showed HBIG and HBIG plus lamivudine to improve graft and patient survival[2]. Subsequently, successful suppression of HBV DNA before transplant by Adefovir resulted in improved pre- and posttransplant survival[3]. More recently, the use of the more potent antiviral agent, entecavir, entirely prevented post-transplant recurrence, even in some patients with prior lamivudine resistance[4]. Whereas the original protocols utilized a lifetime administration of HBIG to maintain a blood titer high enough to prevent reinfection, and this was supplemented with lamivudine and now more potent antiviral agents, newer protocols have reduced the time of administration of the HBIG to 1 year with continued antiviral administration indefinitely after, or even use Entecavir or Tenofovir as a single agent to achieve an undetectable pretransplant viral load and maintain this indefinitely afterward[5].

Liver transplant for hepatitis B virus (HBV) currently yields excellent outcomes: it allows to rescue patients with an HBV-related advanced liver disease, resulting in a demographical modification of the waiting list for liver transplant. In a review of the Scientific Registry of Transplant Recipients (SRTR) database of registrants to the liver transplant list in the United States from 1985 to 2006, the overall number of registrants for HBV began declining after 1998 when oral antiviral therapy was first introduced[5]. Of the main indications for transplant owing to HBV (advanced liver disease, acute liver failure, and HCC), only HCC was increasing in number; registrants for advanced liver disease was declining most rapidly. This trend should continue; the data suggest that those with an early response to antiviral treatment with Tenofovir for acute severe reactivation of HBV have improved non-transplant survival (57% vs 13% for placebo-treated patients)[5].

However, antiviral therapy did not influence survival for those with acute liver failure owing to de novo HBV infection in a North American cohort of patients with acute liver failure[7]. It will likely be at least another decade until the incidence of HCC owing to HBV-induced liver disease begins to significantly decline, and this in part will be owing to treatment of HBV (as well as immunization of populations that began in the early 1990s). Eventually the choice of treatment to prevent HBV reinfection must take into account treatment efficacy, patient adherence, and cost.

Extended criteria for organ acceptance

The unmatched demand and supply rate between organs for transplantation is well known. As a matter of fact, we observed during the last decade a similar annual rate of donors in Europe and United States, while an increase of the “demand” for liver transplantation has been reported, in terms of new patients added in the waiting lists, longer mean waiting time and drop-out rate. Moreover, the lack of organs led to the exclusion from the waiting list of many
patients who can benefit from a transplant\textsuperscript{[8,9]}. In an age of patient-tailored treatments, in liver transplantation as well the aim is to offer the best suitable graft to the patient who can benefit from it. In Europe and in the United States is estimated that almost 10% to 30% of patients listed for liver transplant dies before organ availability\textsuperscript{[8]}. In the United States status I patients i.e., patients entering in the waiting list at the highest medical urgency, reported a 12 folds increased risk of death while on the list compared with those entering at the two lowest categories of urgency\textsuperscript{[10]}. Data from Scandinavia between 1990 and 2001 show that the mortality rate among patients waiting for liver transplant was 16%, while 27% of patients listed for a highly urgent liver transplantation failed to get the graft\textsuperscript{[11]}. For many patients with a severe clinical status needing urgent transplant, the so-called marginal organ donor can provide a chance of cure. Patients that never obtained a transplant due to their clinical characteristics may as well benefit from a marginal donor, overcoming the problem of organ shortage.

The terms extended donor or expanded donor (ECD) mean changes in donor acceptability criteria, which not justifies the negative connotations of these terms. Although criteria to select organs for donation were revised and modified over years, this evolution did not affect neither patients’ nor organs’ survival. Characteristics of donor and recipient, together with allocation scheme, organ procurement and transplant procedure define the “ideal organ”. Moreover, marginal donors can allow to obtain comparable survival rates when an appropriate allocation is ruled out.

Figure 1 Geographic distribution of chronic hepatitis B virus infection-worldwide (2005). For multiple countries, estimates of prevalence of hepatitis B surface antigen (HBsAg), a marker of chronic hepatitis B virus infection, are based on limited data and may not reflect current prevalence in countries that have implemented childhood hepatitis B vaccination; prevalence may vary within countries. Source: Centers for Disease Control and Prevention (http://www.cdc.gov).

Criteria and terms for certified suitability of organ donors: Assumptions and operational strategies in Italy

In 2001 a national commettee of experts nominated by the Italian National Transplant Centre (Centro Nazionale Trapianti-CNT) released a document for all personnel involved in the evaluation process of potential organ donor. The Commitee was made up of infectious disease experts, immunologists, clinical experts, surgeons, coordinators, anatomopathologists, medical examiners and oncologists. During the preparation phase, which lasted one year, the text underwent a series of changes and supplements, resulting in a final version shared with the scientific community and approved by the Italian National Transplant Centre as technical annex (guidelines) to the Ministry Decree of August 2, 2002\textsuperscript{[12]}.

These Guidelines focus on two main aspects: (1) The definition of acceptable/unacceptable risks for donor suitability or single organ utilization; and (2) the establishment of practical steps for the risk evaluation process.

The first aim was to identify the different risk levels and as a result five risk levels have been defined: (1) unacceptable risk; (2) increased but acceptable risk; (3) calculated risk; (4) not assessable risk; and (5) standard risk.

Unacceptable risk: The donor classified under this category should be excluded from donation and no organ can be used for transplantation. For example, HIV1 or 2 positive donors fall into this category, as well as HBsAg and HDV contemporaneous seropositivity. Neoplastic diseases represents an unacceptable risk
with the following exceptions: carcinoma in situ, basal cell carcinoma, cutaneous squamous cell carcinoma without metastases, carcinoma in situ of the cervix, carcinoma in situ of vocal cords, urothelial papillary carcinoma (T0 according to the TNM classification). Eventually, systemic infections caused by agents for which treatments are not feasible and documented prior disease must also be considered as exclusion criteria.

**Increased but acceptable risk:** This category includes organs that can be used in case of urgency or particular clinical conditions of recipients. In these cases, even when the evaluation process shows the presence of pathogens or transmissible disease, organ utilization is allowed in the light of a risk benefit assessment. Patients struck by fulminant hepatitis, or retransplants for liver primary non function, or patients who underwent hepatectomy for trauma with complete organ function loss are included in this category.

**Calculated risk:** Includes all cases where the presence of a specific pathogen or a serological status of the donor (HBsAg⁺, or anti-HCV⁺ or HBCAb⁺) is compatible with transplantation recipients with the same disease or serological status, independently from recipient's health conditions.

**Not assessable risk:** Includes cases for which the evaluation process does not allow an appropriate risk assessment for transmittable diseases for lack of one or more assessment elements (e.g., failure to collect an accurate medical history for lack of relatives, unavailability of microbiology data despite a well-grounded suspicion of infectious pathology).

**Standard risk:** Includes cases for which the evaluation process did not identify any risk factor for transmittable disease. It is the most frequent condition in the assessment of donors and grafts.

The national guidelines also identify some special conditions that concern two main aspects, namely neoplastic and infectious risks.

About infections, special attention should be paid to the following cases: donor with HCV infection; donor with HBV infection (HBsAg positivity); donors with antcore IgG antibodies against B virus (HBCAb). In such cases the guidelines impose the adoption of the following procedures.

**HBsAg positive donor:** If a donor turns out to be HBsAg positive, transplantation is allowed in a HBsAg positive recipient, after informed consent, provided that the following conditions are met: (1) the donor has a negative HDV antigen, negative IgM anti HDV antibodies, negative IgG anti HDV antibodies or with a titre < 1:100 or below the significant level according to the assay used; the absence of IgM anti HDV does not exclude delta virus chronic infection; (2) the liver recipient is not co-infected by delta virus; and (3) the patient follow-up can be monitored on the basis of a common national protocol established by the National Transplant Centre and to record data on a National Registry.

**HBsAg negative donor:** If the recipient is HBsAg negative, he has no anti-HBV antibodies or has a protective anti-HBsAg titre (≥ 10 mUI/mL), transplantation can be performed, after informed consent, when the following conditions are met: (1) the donor has a negative HDV antigen, negative IgM anti HDV antibodies, negative IgG anti HDV antibodies or with a titre < 1:100 or below the significant level according to the used assay; and (2) the patient follow-up can be monitored on the basis of a common national protocol established by the National Transplant Centre and to record data on a National Registry.

As a supplement to these measures, the Italian National Transplant Centre has deemed as proper to support further transplant network health workers, through adhoc developed information tools and an expert task force (second opinion) for evaluation of doubtful cases.

**Study design**

With the intent of developing strategies to increase the donor pool, we set-up a multicenter study involving 3 Liver Transplant Centers in Italy: the Universities of Modena, Bologna and Padova. The study was approved by the institutional review boards at each center. Patients undergoing liver transplantation between March 2004, and May 2010, were retrospectively evaluated. Among 1408 patients who underwent liver transplantation during the study period, 28 (2%) received the graft from HBsAg-positive deceased donors. All subjects were informed of the possible risks, consented to enter the study and signed a written form. For each HBsAg case we collected general clinical features and data regarding the transplantation, including MELD score and ischemia time. Then we retrospectively analyzed post-operative data, namely immunosuppressive therapy, histological evidence of HBV recurrence and antiviral therapy, and episodes of acute rejection.

The Italian regulations issued by the CNT allow HBsAg positive HDV negative recipients, HBCAb positive HDV negative patients, and HBV negative subjects with severe end-stage liver disease and a low life expectancy, to receive grafts from HBsAg positive HDV negative donors. Liver biopsy during organ procurement drives the evaluation on graft status, together with the serovirological complete assessment of HBV and HCV status, including HBV DNA. Moreover, Ishak score ≤ 1 and low inflammation, together HDV negative test in both donor and recipient, are required. HBV viral load, liver function test and age are not considered as exclusion criteria.

We performed liver biopsies routinely pre- and
the 28 recipients were female (median age at operative setting was adjusted to maintain a plasma concentration between 5 and 12 ng/mL. Steroids were started at a dose of 20 mg daily, then tapered down and discontinued within 6 mo.

The median body mass index (BMI) at the time LT was 25.3 (range: 19-34; SD ± 3.2).

Nineteen patients had hepatocellular carcinoma (67.9%) with 13 cases (68.4%) resulting within the Milan criteria, whereas 6 patients (31.6%) were outside Milan and inside UCSF criteria.

The UNOS status was 2A in 5 patients (17.8%), 2B in 15 patients (53.6%), and 3 in 8 patients (28.6%).

**Donor characteristics**

Donor characteristics were reported on Table 3 and the overall serological state of the recipient/donor is shown in the Table 4.

The median age was 52.6 years (range: 13-79, SD ± 16.9). 13 donors were female (46.4%) while 15 donors were male (53.6%). The death causes are reported on the Table 3. The average body mass index (BMI) of donors was 25.3 (range: 19-34; SD ± 3.2). Nineteen patients had hepatocellular carcinoma (67.9%) with 13 cases (68.4%) resulting within the Milan criteria, whereas 6 patients (31.6%) were outside Milan and inside UCSF criteria.

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Data on pre-perfusion histologic features of the biopsies are shown in Table 3. Most of the HBsAg positive grafts had a HAI inflammatory grade between 0-2 (71.4%), followed by an HAI inflammatory grade between 3-4 (28.6%). None of the grafts used had an HAI inflammatory grade score ≥ 5.

In particular, 6 donors (21.4%) had a grading score 0; 7 donors (25%) had a grading score 1; 7 donors (25%) had a grading score 2; 4 donors (14.3%) had a grading score 3; 4 donors (14.3%) had a Grading score 4. All the grafts had a fibrosis stage ≤ 1.

We performed a standard antiviral prophylaxis in all patients, independently from serovirological profile.

All HBsAg-positive recipients were on antiviral treatment with nucleos(t)ide analogues before liver transplantation and continued the same antiviral therapy with the addition of HBV-specific immunoglobulins (HBIG) after liver transplantation. The HBsAg-negative recipients began a similar combined treatment after LT, with lamivudine (LMV) and HBIG.

HBIG administration consisted of 10000 IU during the anhepatic phase, then 5000 IU every day for the first month, subsequently 5000 IU weekly for the second month and finally 5000 IU every 3-4 wk to maintain an anti-HBs titre above 250 IU/mL. This is the standard regimen of the transplant centers and it is applied even to HBV patients receiving an HBsAg-negative graft. Tacrolimus administration in the post-operative setting was adjusted to maintain a plasma concentration between 5 and 12 ng/mL. Steroids were started at a dose of 20 mg daily, then tapered down and discontinued within 6 mo.

**Statistical analysis**

We reported continuous data as mean ± SD, and then compared those data by using the 2-side Student’s t test. The χ² test with Yates’ correction, or Fisher’s exact test when appropriate, was used to compare groups for categorical variables. Survival of grafts and patients were evaluated using the Kaplan-Meier method and compared with the log-rank test. The statistical significance was accepted for P < 0.05. All the statistical analysis were performed using SPSS® 19.0.

**DISCUSSION**

**Recipient characteristics**

Four out of 28 recipients were female (median age at liver transplantation: 57.6 years, range: 26-67). Data were collected from liver transplantation until the last follow-up visit and the average follow-up after liver transplantation was 63.7 mo (range: 0.1-119.4; SD ± 35.8). Recipient characteristics were reported on Tables 1 and 2.

HBV related cirrhosis, with or without HCC, was the indication for liver transplantation in 27 patients (Table 1), while 1 patient was transplanted due to secondary biliary cirrhosis.

The five HBsAg-negative patients showed serological evidence of past HBV infection. The MELD score (Model of End Stage Liver Disease) was applied to stage their liver disease status. In case of HCC, an extra score based on HCC stage was added, according to the centre (or regional) allocation policy.

Patients were transplanted after an average of 452 d on waiting list (range: 37-1962; SD ± 394) and at the time of liver transplantation presented an average MELD biochemical score of 15.6 (range: 7-33; SD ± 6.5) and an average MELD score correction (depending from other clinical variables) of 26.8 (range 11-39; SD ± 7.2).

The median body mass index (BMI) at the time LT was 25.3 (range: 19-34; SD ± 3.2).

Nineteen patients had hepatocellular carcinoma (67.9%) with 13 cases (68.4%) resulting within the Milan criteria, whereas 6 patients (31.6%) were outside Milan and inside UCSF criteria.

Table 1 describes different downstaging treatments for each patient.

The UNOS status was 2A in 5 patients (17.8%), 2B in 15 patients (53.6%), and 3 in 8 patients (28.6%).
For 14 (50%) grafts the staging was 0. Macrosteato-
tosis of the grafts are reported on the Table 3.

**Operative factors**

Cold ischemia time was in an average of 429 min
(range: 255-632) and the warm ischemia time (WIT)
was around 39.7 min (range: 30-55). The average
hemat loss was 2307 mL (range: 300-13000). The
mean length of stay in the Intensive Care Unit (ICU)
was 5.5 d (range: 0-22), while the average Hospital
stay was 21.4 d with a range from 6 to 143.

**Clinical outcome**

None primary non-function (PNF), re-LT, early or
late hepatic artery thrombosis occurred after liver
transplantation.

Two (7.1%) patients who received an HBsAg-
positive donor liver had acute cellular rejection with
a total of 1 event respectively for each patient.

Biliary complication occurred in seven patients
(25%); in particular five biliary stenosis and two biliary
leakages.

Five patients (17.9%) developed a major infection,
2 patients (7.1%) had an Hepatitis C recurrence.

Recurrent of HBV infection, confirmed histological-
ly, occurred in 4 (14.3%) patients who received HBsAg positive grafts. The mean time of
onset of HBV recurrence was 2.1 (± 1.4) mo.

The average follow-up was 63.6 mo (range:
0.1-119.4). The 6 deceased patients died not for
the Hepatitis B recurrence but for different reasons.
In particular, the cause and time of death were
respectively: 1 patient for severe sepsis (0.4 mo),
1 patient for cardiac arrest (0.1 mo), 1 patient for
HCV recurrence (11.8 mo), 2 patients for HCC recurrence
(3.5 and 13 mo, respectively) and one patient for
Merkel cell carcinoma (45 mo).

The 1-, 3- and 5-year graft and patient survival
resulted of 85.7%, 82.1% and 78.4% (Figure 2).

**Read-out**

Liver transplantation is an established therapeutic
modality for patients with end-stage liver disease
or/and hepatocellular carcinoma. However, in recent
years the number of patients needing a transplant
increase overcoming the supply: as a result, the mean
waiting time is now longer than before, with higher
mortality rates of patients waiting for an organ. It is
estimated that 15% to 20% of patients on the waiting
list die each year without receiving a suitable organ.

Several strategies have been developed by
transplant physicians to face this increased demand:
innovative ways of expanding the donor pool are the
use of split and live donor LT. Another approach is the
use of organs from "less-than-perfect donors"; also
called "suboptimal donors". Non-heart-beating donors

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**Table 1 Recipient characteristics at the time of liver transplantation**

| Case | Age | Gender | ABO | BMI | Indication | Real MELD | MELD correct | UNOS | Waiting List(d) | Year LT | HCC criteria | Downstaging type | No. |
|------|-----|--------|-----|-----|------------|-----------|-------------|------|----------------|---------|-------------|-----------------|-----|
| 1    | 62  | M      | 0   | 27  | HCC/HBV   | 26        | 36          | 2A   | 37             | 2007   | MILAN IN     | LOC(1)         | 1   |
| 2    | 65  | M      | B   | 21  | HCC/HBV   | 16        | 39          | 3    | 522            | 2007   | MILAN IN     | LOC(1) + SUR(1) | 2   |
| 3    | 54  | M      | A   | 22  | HCC/HBV   | 10        | 33          | 3    | 513            | 2007   | MILAN IN     | LOC(1)         | 3   |
| 4    | 45  | M      | A   | 24  | HCC/HBV   | 14        | 34          | 2B   | 419            | 2007   | MILAN OUT    | LOC(3)         | 4   |
| 5    | 62  | M      | 0   | 29  | HCC/HBV   | 12        | 33          | 3    | 445            | 2008   | MILAN OUT    | LOC(2)         | 5   |
| 6    | 33  | M      | 0   | 28  | HCC/HBV   | 12        | 35          | 2B   | 515            | 2008   | MILAN IN     | LOC(1)         | 6   |
| 7    | 45  | M      | A   | 23  | HCV/HBV   | 24        | 24          | 2A   | 101            | 2005   | MILAN IN     | LOC(1)         | 7   |
| 8    | 64  | M      | B   | 27  | HBV/HCV   | 33        | 33          | 2A   | 478            | 2005   | HBV/HCV     | LOC(2)         | 8   |
| 9    | 26  | M      | A   | 23  | HBV      | 23        | 23          | 2B   | 742            | 2005   | MILAN IN     | LOC(2)         | 9   |
| 10   | 64  | M      | A   | 22  | HCC/HBV   | 11        | 23          | 2B   | 144            | 2005   | MILAN IN     | LOC(2)         | 10  |
| 11   | 65  | F      | B   | 25  | HCC/HBV   | 12        | 24          | 2A   | 195            | 2006   | MILAN IN     | LOC(2)         | 11  |
| 12   | 56  | M      | A   | 24  | HCC/HBV   | 10        | 24          | 2B   | 217            | 2006   | MILAN OUT    | LOC(5)         | 12  |
| 13   | 61  | F      | 0   | 23  | HCC/HBV   | 14        | 30          | 2B   | 356            | 2006   | MILAN IN     | LOC(2)         | 13  |
| 14   | 59  | M      | A   | 24  | HBV      | 30        | 30          | 2B   | 118            | 2007   | MILAN IN     | LOC(1)         | 14  |
| 15   | 48  | F      | A   | 27  | CBS      | 21        | 33          | 2A   | 82             | 2007   | MILAN IN     | LOC(2)         | 15  |
| 16   | 65  | M      | A   | 24  | HCC/HBV   | 8         | 25          | 2B   | 358            | 2009   | MILAN IN     | LOC(2)         | 16  |
| 17   | 57  | M      | A   | 30  | HCC/HBV   | 14        | 39          | 2B   | 576            | 2009   | MILAN OUT    | LOC(7)         | 17  |
| 18   | 55  | M      | B   | 28  | HCC/HBV   | 18        | 26          | 2B   | 38             | 2009   | MILAN IN     | LOC(3)         | 18  |
| 19   | 65  | M      | A   | 24  | HCC/HBV   | 7         | 25          | 2B   | 371            | 2009   | MILAN IN     | LOC(3)         | 19  |
| 20   | 63  | M      | A   | 24  | HCC/HBV   | 16        | 25          | 2B   | 1962           | 2010   | MILAN IN     | LOC(2)         | 20  |
| 21   | 60  | M      | A   | 34  | HCC/HBV   | 10        | 27          | 2B   | 330            | 2010   | MILAN OUT    | LOC(1) + SUR(1) | 21  |
| 22   | 61  | M      | A   | 24  | HBV      | 11        | 11          | 3    | 855            | 2010   | HBV/HCV     | LOC(1)         | 22  |
| 23   | 67  | M      | 0   | 19  | HBV      | 16        | 16          | 2B   | 742            | 2004   | MILAN IN     | LOC(2)         | 23  |
| 24   | 55  | M      | A   | 24  | HBV      | 17        | 17          | 3    | 127            | 2004   | MILAN IN     | LOC(2)         | 24  |
| 25   | 60  | F      | 0   | 29  | HBV/HCV  | 17        | 17          | 2B   | 961            | 2004   | MILAN IN     | LOC(1)         | 25  |
| 26   | 55  | M      | A   | 26  | HCC/HBV   | 10        | 20          | 3    | 748            | 2005   | MILAN IN     | LOC(1)         | 26  |
| 27   | 54  | M      | A   | 27  | HCC/HBV   | 13        | 19          | 3    | 134            | 2006   | MILAN OUT    | LOC(2)         | 27  |
| 28   | 67  | M      | 0   | 22  | HCC/HBV   | 12        | 29          | 3    | 575            | 2009   | MILAN IN     | LOC(1) + SUR(1) | 28  |

1 LOC: Locoregional therapy [transcatheter arterial chemoembolization (TACE) and/or radiofrequency ablation (RITA)]; SUR: Surgery; MELD: Model for end-stage disease; BMI: Body mass index; LT: Liver transplantation; HCC: Hepatocellular carcinoma; CBS: Secondary biliary cirrhosis; HCV: Hepatitis C virus.
and donors older than 65 years belong to such donors, as well as steatotic liver allografts and patients with previous exposure to HBV or HCV. Also the selection HBsAg positive donors represents a way to expand the pool of transplantable grafts.

On the other hand, living donor (LD) LT was adopted in Eastern countries to counterbalance the lack of deceased donors due to cultural reasons. Living donors and split liver transplantation have been used to contrast the donor shortage, but they have failed to significantly decrease the number of patients on the wait list. Those two approaches have ethical issues as well as steatosic and donors older than 65 years belong to such donors, as well as steatotic liver allografts and patients with previous exposure to HBV or HCV. Also the selection HBsAg positive donors represents a way to expand the pool of transplantable grafts.

As a matter of fact, wider acceptance criteria can assure more donors available for transplantation and several guidelines are available to classify donors as standard or ECD [16-20]. Two main categories of ECD can be identified: the first one includes grafts with risk of dysfunction due to direct or indirect liver injury, the second accounts for the risk of disease transmission between donor and recipient.

Table 2 Recipient characteristics  

| Recipient variables | n = 28 |
|---------------------|--------|
| Transplant center (No. of patients) |        |
| Modena | 6 (21.4) |
| Bologna | 16 (57.1) |
| Padova | 6 (21.4) |
| Gender |        |
| Male | 24 (85.7) |
| Female | 4 (14.3) |
| ABO blood group |        |
| Isogroup | 28 (100) |
| 0 | 7 (25) |
| A | 17 (60.7) |
| B | 4 (14.3) |
| Age (yr), mean (range, SD) | 57.6 (26-67, ± 8.7) |
| Body mass index, mean (range, SD) | 25.3 (19-34, ± 3.1) |
| Associated hepatocellular carcinoma |        |
| Meeting Milan criteria | 19 (69.7) |
| Meeting UCSF criteria | 13 (44.4) |
| Correct MELD score, mean (range, SD) | 26.7 (11-39, ± 7.2) |
| UNOS status |        |
| 2A | 5 (17.5) |
| 2B | 15 (53.6) |
| 3 | 8 (28.6) |
| Time waiting list (d), mean (range, SD) | 452 (37-1962, ± 393.5) |
| HBsAg status |        |
| Positive | 23 (82.1) |
| Negative | 5 (17.9) |
| HBV DNA positive at LT | 12 (52.1) |
| HCV co-infection | 4 (14.3) |
| HDV co-infection | 0 |

UCSF: University of California, San Francisco; LT: Liver transplantation; HCV: Hepatitis C virus; HDV: Hepatitis D virus; MELD: Model for end-stage disease.

In the first case should be taken into account that those grafts must be carefully evaluated and transplanted in recipients capable to overcome the increased physiologic stress.

The ECD liver disease transmission risk is broken into 2 separate categories: (1) viral transmission of HCV, HBV, HTLV-1, and HTLV-2; and (2) malignancy transmission. Our previously reported results are consistent with other studies showing that it is safe to allocate grafts from HCV positive donors into HCV positive recipients [21-25]. The HCV positive donor liver must have no evidence of cirrhosis or stage > 1 fibrosis. It is clear that HCV-positive livers should be declassified as ECDs.

HBV scenario: About 2 billion people have serological evidence of present or past HBV infection worldwide, and a prevalence of more than 350 million cases of chronic infection is estimated [26].

The selection criteria of the recipient of HBcAb-positive donors are currently debated, while it has been demonstrated that a lifelong antiviral therapy is needed after transplantation of those grafts [23,27,28]. The majority of chronic HBV infections is nowadays present in the Western Pacific region [29], while a recent survey from Korea showed an overall HBsAg prevalence of 3.7%. This group of ECD is currently underestimated due to the high risk of HBV reactivation and to the paucity of clinical data, and up-to-now they are not used in most of the transplant centers.

Because of the existing shortage of organs, the increased demand for LT, and given the possible implications in terms of extension of the donor pool, the use of HBsAg-positive grafts should be studied to assess safety policies. To date, only a few studies exist regarding the effect of donor HBsAg positivity on survival (Table 5). These available reports yield conflicting results and are limited by small sample sizes and short follow-up [30-38].

Gonzalez-Peralta et al [31] were the first to report a successful LT of an HBsAg-positive graft into HBV negative recipient, who shortly afterwards turns HBsAg positive. Several reports in literature attested...
the use of HBIg and antiviral drugs against HBV such as lamivudine, adefovir dipivoxil, and tenofovir in recipients with HBsAg-positive grafts. Loggi et al. reported a series of 10 HBsAg-positive grafts with HBIg and nucleos(t)ide analogue prophylaxis. In their experience only one patient died due to HCV recurrence over a mean follow-up period of 36.8 mo. In a cohort with 8 patients out of 10 positive for HBsAg after LT, no patient ever had any signs of active HBV hepatitis.

However, there was no comparison of outcomes between HBsAg-positive graft recipients with and without HBIg prophylaxis.

Using comprehensive clinical data from the SRTR database, Li et al. failed to identify any significant association between the use of HBsAg-positive donors and post-transplant graft or patient survival, after adjusting for other predictors of post-transplant survival. Their results demonstrate that HBsAg-positive donors for liver transplantation are safe and comparable in terms of outcomes and long-term survival to the use of HBsAg-negative grafts. Furthermore, other studies clearly showed that using HBIg may improve post-transplant survival in recipients with HBsAg-positive grafts.

Several innovations have been introduced during the last two decades to improve the outcomes of patients receiving LT for HBV-related liver disease, such as the administration of HBIg since the early 1990s and lamivudine in late 1990s. Although there is now a consensus in favor of the use of HBIg in HBV-positive recipients, its application in HBV positive donors is still unclear.

Our study shows that the use of HBsAg positive grafts is a safe procedure when carried out in combination with appropriate antiviral therapy and when the graft fibrosis is ≤ 1 and the grading score is ≤ 4.

The 4 Hepatitis B recurrences that we have followed during the post-LT didn’t influence the graft and patient survival. From our own experience, there were no cases of PNF and the infectious and biliary complications were similar to the cases of HBsAg negative graft recipients.

However, our research shows some relevant limitations: first, even if it represents the major European study, the number of patients is still too low and therefore it doesn’t allow to establish ultimate conclusions. Then, we have chosen to focus only on a descriptive kind of analysis while for the future it will be necessary to perform comparative studies and matched analysis. Second, the lack of a common serial protocol hepatic biopsies has not allowed to examine the histological evolution of these grafts as well as a serial protocol for the dosage of the HBsAg quantification and the HBV-DNA level.

**CONCLUSION**

Despite the small number of cases, our results suggest

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**Table 3 Donor characteristics**

| Case | Age | Gender | ABO Gr. | BMI (kg/m²) | Cause of death | Time ICU (d) | Sodium (mEq/mL) | Vasopressors | Histologic activity index | Graft steatosis |
|------|-----|--------|---------|-------------|---------------|-------------|-----------------|--------------|-------------------|---------------|
| 1    | 59  | M      | 0       | 26          | CH            | 2           | 165             | No           | 1                 | 0%            |
| 2    | 13  | F      | B       | 19          | T             | 21          | 161             | Yes          | 2                 | 0%            |
| 3    | 69  | M      | A       | 24          | CH            | 3           | 152             | No           | 0                 | 0%            |
| 4    | 72  | F      | A       | 27          | CH            | 7           | 150             | No           | 1                 | 0%            |
| 5    | 66  | M      | 0       | 22          | CH            | 5           | 137             | Yes          | 3                 | 1            |
| 6    | 60  | M      | 0       | 26          | CH            | 4           | 158             | No           | 2                 | 0%            |
| 7    | 73  | M      | A       | 26          | CH            | 2           | 151             | Yes          | 2                 | 1%            |
| 8    | 51  | M      | B       | 23          | CH            | 6           | 149             | Yes          | 3                 | 1%            |
| 9    | 54  | M      | A       | 24          | T             | 13          | 160             | Yes          | 2                 | 1%            |
| 10   | 72  | F      | A       | 23          | T             | 2           | 148             | No           | 2                 | 0%            |
| 11   | 60  | F      | B       | 29          | CH            | 8           | 162             | Yes          | 3                 | 1%            |
| 12   | 65  | M      | A       | 29          | CH            | 3           | 140             | Yes          | 4                 | 1%            |
| 13   | 50  | M      | 0       | 24          | CH            | 12          | 141             | Yes          | 2                 | 1%            |
| 14   | 48  | M      | A       | 23          | T             | 2           | 143             | No           | 4                 | 1%            |
| 15   | 26  | M      | A       | 23          | T             | 1           | 145             | Yes          | 1                 | 0%            |
| 16   | 52  | F      | A       | 28          | CH            | 1           | 155             | Yes          | 4                 | 1%            |
| 17   | 79  | F      | A       | 24          | CH            | 19          | 136             | No           | 0                 | 0%            |
| 18   | 46  | F      | B       | 29          | CH            | 2           | 158             | No           | 0                 | 0%            |
| 19   | 61  | M      | A       | 29          | CH            | 3           | 156             | No           | 1                 | 1%            |
| 20   | 50  | M      | 0       | 25          | CH            | 6           | 154             | Yes          | 2                 | 1%            |
| 21   | 44  | F      | A       | 24          | CH            | 6           | 144             | Yes          | 4                 | 1%            |
| 22   | 23  | F      | A       | 21 other    | CH            | 7           | 149             | Yes          | 1                 | 0%            |
| 23   | 57  | F      | 0       | 24          | CH            | 4           | 147             | Yes          | 0                 | 0%            |
| 24   | 59  | M      | A       | 24          | CH            | 1           | 160             | Yes          | 0                 | 0%            |
| 25   | 35  | F      | 0       | 23          | CH            | 1           | 140             | Yes          | 1                 | 0%            |
| 26   | 36  | M      | A       | 25          | T             | 2           | 146             | Yes          | 3                 | 1%            |
| 27   | 23  | M      | A       | 29          | T             | 2           | 147             | Yes          | 0                 | 0%            |
| 28   | 66  | F      | 0       | 27          | CH            | 3           | 151             | Yes          | 1                 | 0%            |

BMI: Body mass index; CH: Cerebral hemorrhage; T: Trauma; ICU: Intensive care unit.
that the utilization of grafts from deceased HBSAg positive donors, according to our allocation criteria, is feasible and HBV can be controlled with graft stability if selection of grafts and postoperative antiviral treatment are appropriately managed. This way it could be possible to expand the donor pool, especially in the high-endemic areas where a large proportion of patients are highly viremic and HBeAg positive.

Long-term follow-up data and large-scale mul-

| Case | HBSAg | HBsAb | HBCAb | HBeAb | HBV DNA | HDV | HDV RNA | HCV Ab | Therapy pre-LT | Mutation |
|------|-------|-------|-------|-------|---------|-----|---------|--------|---------------|----------|
| R1   | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D1   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R2   | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D2   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R3   | +     | -     | +     | -     | +       | -   | -       | -      | Lam + Adef    | Yes      |
| D3   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R4   | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D4   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R5   | +     | -     | +     | -     | +       | -   | -       | -      | Lam + Adef    | Yes      |
| D5   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R6   | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D6   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R7   | -     | +     | +     | -     | -       | -   | -       | -      | No            | No       |
| D7   | +     | -     | +     | -     | +       | -   | -       | -      | No            | No       |
| R8   | -     | +     | +     | -     | -       | -   | -       | -      | No            | No       |
| D8   | +     | -     | +     | -     | +       | -   | -       | -      | No            | No       |
| R9   | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D9   | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R10  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D10  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R11  | +     | -     | +     | -     | +       | -   | -       | -      | No            | No       |
| D11  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R12  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D12  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R13  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D13  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R14  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D14  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R15  | +     | -     | +     | -     | +       | -   | -       | -      | No            | No       |
| D15  | +     | -     | +     | -     | +       | -   | -       | -      | No            | No       |
| R16  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D16  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R17  | +     | -     | +     | -     | +       | -   | -       | -      | Lam + Adef    | Yes      |
| D17  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R18  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D18  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R19  | +     | -     | +     | -     | +       | -   | -       | -      | Adefovir      | No       |
| D19  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R20  | +     | -     | +     | -     | +       | -   | -       | -      | Lam + Adef    | No       |
| D20  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R21  | +     | -     | +     | -     | +       | -   | -       | -      | NA            | No       |
| D21  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R22  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D22  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R23  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D23  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R24  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | Yes      |
| D24  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R25  | -     | +     | +     | -     | +       | -   | -       | -      | No            | No       |
| D25  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R26  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D26  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R27  | +     | -     | +     | -     | +       | -   | -       | -      | Adef         | Yes      |
| D27  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |
| R28  | +     | -     | +     | -     | +       | -   | -       | -      | Lam           | No       |
| D28  | +     | -     | +     | -     | +       | -   | -       | -      | No            | -        |

R: Recipient; D: Donor; LT: Liver transplant; Lam: Lamivudine; Adef: Adefovir.
ticenter studies are required to confirm our findings.

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Table 5 Literature review

| Ref. | Year | Cases |
|------|------|-------|
| Francillon et al. | 2005 | 3 |
| Jiang et al. | 2011 | 6 |
| Loggi et al. | 2012 | 10 |
| Saidi et al. | 2013 | 92 |
| Choi et al. | 2013 | 8 |
| Li et al. | 2013 | 78 |
| Ju et al. | 2013 | 23 |
| Yu et al. | 2014 | 42 |
| Krishnamoorthy et al. | 2014 | 28 |
| Jeng et al. | 2015 | 14 |
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