Long-term survival after liver transplantation for alcoholic liver disease

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Abstract
Currently, alcoholic cirrhosis is the second leading indication for liver transplantation in the United States and Europe. The quality of life and survival after a liver transplantation (LT) in patients with alcoholic liver disease (ALD) are similar to those in patients with other cirrhosis etiologies. The alcoholic relapse rate after a LT varies from 10%-50%, and these relapse patients are the ones who present a reduced long-term survival, mainly due to cardiovascular diseases and the onset of de novo neoplasms, including lung and upper aerodigestive tract cancer. Nearly 40% of ALD recipients resume smoking and resume it early post-liver transplantation (LT). Therefore, our pre-and post-LT follow-up efforts regarding alcoholic liver disease should be focused not only on alcoholic relapse but also on treating and avoiding other modifiable risk factors such as tobacco. The psychiatric and psychosocial pre-LT evaluation and the post-LT follow-up with physicians, psychiatrists and addiction specialists are important for reversing these problems.

INTRODUCTION
Excessive alcohol consumption causes approximately 2.5 million deaths per year and is responsible for almost 4% of mortality worldwide. Alcohol has been associated with nearly 60 types of diseases and is the third leading risk factor for disease and disability worldwide. Furthermore, excessive alcohol consumption contributes to multiple social problems, including violence, child neglect and ab-
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Table 1 Primary indication for liver transplantation in Europe and the corresponding survival

| Indication for LT | Patients (n) | From 1998 to 2009 |
|------------------|-------------|-------------------|
|                  |             | 1 yr  | 5 yr  | 10 yr |
| Alcoholic cirrhosis | 15019 | 86%  | 73%  | 59%  |
| Acute hepatic failure | 6507  | 70%  | 64%  | 58%  |
| Cirrhosis virus C | 10753 | 80%  | 65%  | 53%  |
| Cirrhosis viral C and alcoholic | 1790 | 85%  | 69%  | 54%  |
| Cirrhosis virus B | 4187 | 83%  | 74%  | 68%  |
| Hepatocarcinoma and cirrhosis | 9122 | 83%  | 62%  | 49%  |
| Cholestatic disease | 9114 | 87%  | 78%  | 70%  |
| Autoimmune cirrhosis | 1892 | 85%  | 76%  | 67%  |
| Hemochromatosis | 468 | 76%  | 66%  | 53%  |

Adapted from the European Liver Transplant Registry[10]. LT: Liver transplantation.

Alcoholic liver disease (ALD) is the main cause of cirrhosis in Western countries and contributes to one third of the mortality associated with liver cirrhosis. Furthermore, ALD is the second most common indication for liver transplantation (LT) in the United States and Western Europe[2-7], accounting for about 40% of transplants in Europe and 20% of transplants in the United States[2-7].

If we analyze the medium- and long-term survival of a transplant patient, there is no doubt that the recipients with an alcoholic etiology have great results, with a European global 5-year survival rate of 73% and a 10-year survival rate of 59%[2,3], rates that are superior to those for recipients with other etiologies (Table 1)[9]. Therefore, we can infer that ALD is a good indication for LT[9]. However, these excellent results are diminished when a harmful alcohol consumption relapse occurs. These relapses and their possible consequences on the transplant and on the survival and quality of life of the patient, as well as possible actions to avoid relapses, are discussed below.

An evidence-based approach was used for this review. MEDLINE search was performed to September 2013 using the following MeSH terms: liver transplantation, alcohol-related disorders, alcohol-induced disorders, drug abuse, substance abuse, tobacco, and neoplasm. Searches were limited to English language articles. References of suitable articles were searched for other appropriate articles.

**QUALITY OF LIFE AFTER LT**

Quality of life involves physical, mental and social well-being, including working life, and is considered a survival indicator that even surpasses traditional indicators[9]. In the cirrhotic patient, a decrease in physical, psychological and intellectual capabilities occurs alongside liver function impairment[9]. Therefore, it is logical to believe that those patients with advanced liver disease might have a significant reduction in their quality of life. Therefore, the question we might ask is the following: does LT improve the quality of life in these patients? To answer this question, numerous studies that evaluated the quality of life after LT have been performed[10-17]. However, it is not easy to extrapolate the results of these studies due to the heterogeneity of the post-transplantation follow-up times and the instruments that were used to evaluate the different spheres comprising the quality of life[10-17]. In general, studies reveal a significant short-term improvement in the quality of life with no differences observed between ALD and non-alcoholic liver disease[10-18]. Notably, although ALD patients seem less likely to be involved in structured social activities during the post-LT phase than the patients who were transplanted as a result of other etiologies, the ALD patients return to society to lead active and productive lives[10]. Few studies analyzed the quality of life long-term. As a representative study, the study by Ruppert et al[15], that included a 12-year follow-up after LT does not show a progressive loss of quality of life in these patients after the first year of LT[15].

Regarding job reinsertion, the age at the time of the LT, the duration of the pre-transplant disability and the physical and general health status of the patient are the factors that correlate more with employment[14,20-22]. Globally, approximately half of the LT patients return to work[15,23], with no differences between the ALD patients and those with the remaining etiologies[13,23].

**ALCOHOLIC RELAPSE/RECIDIVISM AFTER LT**

We must recall that, although LT effectively restores the physiological function of the liver and reverses the complications of portal hypertension, LT does not treat the underlying alcoholism. Alcoholism is a life-long disease that is often characterized by episodes of a relapsing-remitting pattern of alcohol use despite the physical, psychological and social consequences, wherein the probability of long-term sobriety becomes robust only after 5 years of sustained abstinence[24,25].

**Dimension of the problem**

Addiction specialists define relapse as the prolonged resumption of heavy alcohol intake and distinguish this harmful drinking behavior from so-called slips, which are defined as sporadic drinking episodes followed by the reestablishment of abstinence[26]. This definition of alcoholic relapse is in contrast to that by most transplant centers that consider any alcohol consumption after LT to be unacceptable and define recidivism as any use of alcohol after LT[28]. Most of these episodes of alcohol abuse are effectively diagnosed with interviews and validated self-reporting questionnaires[29,30].

Reviews summarizing the post-transplantation alcoholic relapse rates note differences across studies ranging from 10%-95%, likely due to several factors, including variations in the study methodology, the definition and assessment of relapse and the duration of the follow-up (Table 2)[26,31-60]. In general, the risk that alcoholic
Table 2 Alcohol relapse (any use) after liver transplantation for alcohol liver disease

| Ref. | Study design | Patients (n) | Year | Follow-up mean or median (mo) | Relapse rate |
|------|--------------|--------------|------|------------------------------|-------------|
| Bird et al[36] | Retrospective | 18 | 1990 | 84 | 17% |
| Kumar et al[64] | Retrospective | 52 | 1990 | 25 | 12% |
| Gish et al[36] | Retrospective | 29 | 1993 | 24 | Prospective |
| Knechtle et al[36] | Retrospective | 32 | 1993 | Not stated | 13% |
| Berlakovich et al[27] | Retrospective | 44 | 1994 | 78 | 32% |
| Howard et al[36] | Retrospective | 20 | 1994 | 43 | 95% |
| Krom et al[44] | Retrospective | 30 | 1994 | Not stated | 13% |
| Osorio et al[36] | Retrospective | 43 | 1994 | 21 | 19% |
| Gerhardt et al[36] | Retrospective | 41 | 1996 | 47 | 49% |
| Tringali et al[36] | Retrospective | 58 | 1996 | 27 | 21% |
| Zibari et al[36] | Retrospective | 29 | 1996 | Not stated | 7% |
| Coffman et al[36] | Prospective | 91 | 1997 | Not stated | 20% |
| Anand et al[36] | Retrospective | 39 | 1997 | 25 | 13% |
| Everson et al[36] | Retrospective | 42 | 1997 | Not stated | 17% |
| Foster et al[36] | Retrospective | 63 | 1997 | 49 | 21% |
| Lucey et al[36] | Retrospective | 50 | 1997 | 63 | 34% |
| Stefanini et al[36] | Retrospective | 18 | 1997 | Not stated | 27% |
| Fabrega et al[36] | Retrospective | 44 | 1998 | 40 | 18% |
| Tang et al[36] | Retrospective | 56 | 1998 | 24 | 50% |
| Yates et al[36] | Retrospective | 43 | 1998 | 21 | 19% |
| Gledhill et al[36] | Retrospective | 24 | 1999 | 14 | 25% |
| Pageaux et al[36] | Retrospective | 53 | 1999 | 32 | 42% |
| Pereira et al[36] | Retrospective | 56 | 2000 | 30 | 50% |
| Burra et al[36] | Retrospective | 34 | 2000 | 40 | 33% |
| Jain et al[36] | Retrospective | 185 | 2000 | 94 | 20% |
| D’imartini et al[36] | Prospective | 36 | 2001 | 12 | 38% |
| Gish et al[36] | Retrospective | 61 | 2001 | 83 | 20% |
| Mackie et al[36] | Retrospective | 46 | 2001 | 25 | 53% |
| Bellamy et al[36] | Retrospective | 123 | 2001 | 84 | 13% |
| Karmann et al[36] | Retrospective | 49 | 2001 | 36 | 21% |
| Bravata et al[36] | Retrospective | 313 | 2001 | Not stated | 32% |
| Pageaux et al[36] | Retrospective | 128 | 2003 | 54 | 31% |
| Jabbour et al[36] | Retrospective | 111 | 2004 | 44 | 15% |
| Cuadrado et al[36] | Retrospective | 54 | 2005 | 99 | 26% |
| Bjoromsson et al[36] | Retrospective | 103 | 2005 | 31 | 33% |
| Kelly et al[36] | Retrospective | 90 | 2006 | 67 | 31% |
| Piltzmann et al[36] | Retrospective | 300 | 2007 | 89 | 19% |
| Karim et al[36] | Retrospective | 80 | 2010 | Not stated | 10% |
| Schmieding et al[36] | Retrospective | 300 | 2011 | 84 | 27% |
| Rice et al[36] | Retrospective | 300 | 2013 | 78 | 16% |

recipients return to any alcohol use after LT is between 10%-50% with 8-year follow-ups[28,31-66]. More specifically, between 20 and 50% of the patients who received a liver transplant for end-stage ALD acknowledge some alcohol use in the first 5 years after LT, and 10%-15% will resume heavy drinking[28,31,62]. This finding compares favorably to post-treatment relapse rates as high as 80%-95% in treatment studies of alcoholics without ALD[36].

In a meta-analysis performed in 2008 on the risk of recurrence of substance use after solid organ transplantation that included 54 studies, 50 of which were on LT, it was concluded that the relapse rate of alcohol consumption after LT was 5.6 cases per 100 patients/year, and the relapse rate of excessive consumption was 2.5 cases per 100 patients/year[27]. Additionally, the authors concluded that it was possible that these cumulative incidence rates would become stable at some point that could not be established because few of the studies had a post-transplant follow-up over 7-8 years[27].

Being able to determine the threshold of initiation of alcohol consumption after a liver transplant would be of great clinical and therapeutic utility because this knowledge would allow us to plan specific interventions more accurately. DiMartini et al[36] have described four different patterns of alcohol consumption depending on the starting date, quantity and duration as follows: (1) Minimum consumption over a long period; (2) Early consumption that progresses rapidly to moderate consumption; (3) Early consumption that progresses continuously to a harmful consumption; and (4) Moderate consumption with a late start. These results indicate that we should maintain surveillance after the first year post-LT, despite the fact that the rates for the initiation of consumption generally attenuate over time post-transplantation, probably due to the increase in the stability of sobriety over time[64].

Consequences
The impact of alcohol use on the patient is not entirely clear. The available literature suggests that abusive drinking leads to a decrease in both graft and patient survival and may also lead to the lack of therapeutic compliance.

Adherence to immunosuppressant medication: In LT, adherence to the immunosuppressant treatment and any other drugs medically prescribed is crucial for positive short- and long-term results in the transplanted patients because non-adherence to these measures might lead to graft rejection and failure[66]. Reviews summarizing the nonadherence rates post-transplantation note differences across studies ranging from 3%-47%, probably due to several factors, including variations in study methodology, definitions and the small number of patients included in the studies[63-65].

Therefore, we question whether the consumption or abuse of substances pre-LT increases the risk of non-adherence to immunosuppressant treatment[63,66] and whether an alcoholic relapse is associated with pre-LT alcohol use[66,62,63]. Berlakovich et al[36] studied the effect of alcohol consumption on adherence and found that the patients who relapsed (15 of the 118 transplanted ALD patients) had a non-adherence rate that was not different from that of the patients who did not relapse. This finding was also demonstrated in a study performed in our hospital, where there was no association between the adherence to drug treatment and the presence or absence of alcoholic relapse in a series of transplanted ALD patients[66]. We believe that this concept is endorsed by the meta-analysis of Dew et al[27], which showed a lack of association. Specifically, these authors observed that European studies presented a lower non-adherence rate to immunosuppressant treatment compared to the North American studies, despite presenting significantly higher relapse rates of harmful alcohol consumption[27].

Thus, the lack of adherence seems to be linked to the personality of the patient, the acknowledgement of their
disease, the complexity of the medical prescriptions, the presence of family support and the doctor-patient relationship, more so more than to alcohol consumption.\(^{67,68}\)

**Liver graft:** Resuming alcohol consumption after LT may damage the graft because of poor compliance with immunosuppressive drug treatment and alcohol-related liver injury. Graft loss from recurrent disease related to alcohol use is rare.\(^{69,70}\) Globally, graft dysfunction related to relapse ranges from 0%-17%, although deaths related to relapse range from 0%-5%\(^{80,81}\). There are few studies on the severity of the liver lesions associated with alcoholic consumption after LT on ALD patients\(^{80,82-84}\). Rice et al\(^{85}\) found that alcohol relapse is associated with advanced fibrosis on biopsy. In contrast, our histologic study revealed only mild hepatic changes directly attributable to alcohol.\(^{86}\) As reported by Pageaux et al\(^{87}\), fatty changes and pericellular fibrosis represented the most relevant histological findings in patients who resumed heavy alcohol intake.

In contrast, several studies have shown that ALD LT patients have a lower rejection risk compared to other LT indications, suggesting an inhibitory effect of alcohol over some components of the immune response.\(^{27-29,76-78}\) We have corroborated this finding and observed a lower incidence of acute rejection among patients who relapse to alcohol consumption compared to abstemious patients.\(^{30}\)

**Long-term survival:** Jain et al\(^{31}\) observed that the 5-year post-transplantation survival rate was significantly lower for transplanted ALD patients compared to transplanted non-alcoholic liver disease patients, mainly due to cardiovascular events and de novo neoplasms, especially of the aerodigestive tract, which suggests that immunosuppression by itself is not an initiation factor for malignant changes.\(^{30}\) We reached a similar conclusion after we evaluated the alcoholic relapse risk in a series of transplanted ALD patients and the influence of a relapse on survival.\(^{32}\) In our case, the 5-year survival rate was similar between relapers and non-relapers (92.9% vs 92.4%, respectively), but after 10 years, the survival rate decreased significantly in the relapse patients (45.1% vs 85.5%), with malignant tumors and cardiovascular events the main cause of death in these patients (Figure 1)\(^{33}\). In addition, tobacco consumption was observed in all the patients with an alcoholic relapse and in only one quarter of the abstemious patients, which might explain the higher mortality rate due to cardiovascular events and neoplasms in these patients; this finding has been observed in other studies, as will be discussed later. However, the transplanted ALD patients are potentially affected not only by alcohol consumption but also by liver diseases with other etiologies. This finding has been shown in a recent study in which excessive alcohol consumption had a negative impact on long-term survival after LT regardless of the indication.\(^{82}\)

Despite these results, it is important to distinguish from among the relapers those who are “slip” drinkers (mild alcohol consumption that is usually isolated or self-limited) and those who are “heavy” drinkers (a long period of alcohol consumption with a loss of control) because the former have a better survival rate compared to the latter.\(^{83}\)

### IDENTIFICATION OF THESE PATIENTS

Because of the above findings, it is important to identify relapse patients, but it is more important to prevent this relapse by identifying the patients at risk.

In order to predict the post-LT alcoholic relapse risk with a high degree of accuracy, it is necessary to acknowledge the risk factors that have a strong correlation, which has not yet been achieved. In this regard, numerous studies have identified factors related to the risk of post-LT alcoholic relapse, such as alcohol dependence, an age less than 40 years at the time of the transplantation, a lack of family and social support, a family history of alcoholism, personality or psychiatric disorders, previous abstinence or substance abuse failures, younger age at LT, and the refusal of further rehabilitation before the LT.\(^{27,30,37,40,42,43,54,78,83-87}\) However, this association has not yet been corroborated in other studies.\(^{26,30,37,54,73,84-86}\) For this reason, Kotlyar et al\(^{88}\) decided to perform a critical review of the literature on LT in ALD candidates and concluded that patients with a lack of social support, active smoking, psychotic or personality disorders or a pattern of nonadherence should be listed only with reservation, and those who have a diagnosis of alcohol abuse as opposed to alcohol dependence may make better transplant candidates. Finally, the most controversial among these risk factors is the 6-mo pre-LT period of abstinence, about which many studies have reported a high predictive power regarding relapse,\(^{27,36,54,69-71}\) while others have not found such a correlation.\(^{49,79,89,90,92}\)

Most LT centers in Europe and the United States require a minimum of 6 mo of abstinence before being included in the waiting list. This common practice is based on two points: first, the possibility of improving liver function and possibly avoiding the LT, and second,
the higher alcoholic relapse rate reported in patients with a period of abstinence less than 6 mo\textsuperscript{[115,116]}. Both points have been discussed. Veldt et al\textsuperscript{[94]} demonstrated that those with irreversible ALD were identified with 3 mo of abstinence; out of 74 patients with a Child-Pugh C liver function, the percentage of patients with improvement after 1, 2 and 3 mo of abstinence was 23%, 40% and 66%, respectively, and the remaining 33% did not show improvement at a 1-year follow-up. Furthermore, it has not been proven that this 6-mo abstinence period improves survival after LT\textsuperscript{[95]}. Considering all of this, and although improved post-LT abstinence rates have been documented with a longer pre-LT abstinence period, a cut-off point has not yet been established\textsuperscript{[96,97]}. Therefore, the pre-LT alcohol abstinence period could be shortened for some patients because this factor by itself is a poor indicator of post-LT relapse. In addition, some patients, especially those with a high Model for End-Stage Liver Disease score, have a considerable risk of mortality during the 6 mo abstinence period\textsuperscript{[98,99]}. In conclusion, a thorough assessment by a trained alcoholism and addiction professional, rather than defined sobriety periods, should be the tool used to assess the future risk of alcoholic relapse in the alcoholic patient.

**ALCOHOL AND TOBACCO**

Patients who undergo LT have an unexpectedly high rate of de novo extrahepatic cancer\textsuperscript{[79,99,102]}, including lung and upper aerodigestive tract cancers\textsuperscript{[98,100,101]}, and studies have reported that patients with ALD are particularly affected\textsuperscript{[97,99,102]}. Indeed, these tumors are known to be associated with alcohol intake and smoking because the carcinogenic or co-carcinogenic effects of smoking and drinking might be enhanced by the post-LT immunosuppressive therapy\textsuperscript{[103]}. The purported mechanisms of alcohol-mediated oncogenesis are poorly understood, but these pathways may include the carcinogenic properties of acetaldehyde and/or the inhibition of DNA methylation via the alteration of retinoid processing\textsuperscript{[98,104,105]}. Saigal et al\textsuperscript{[106]} found that patients who underwent LT for ALD appeared to have an increased risk of developing post-transplantation malignancies compared with those who underwent LT for other liver diseases. These authors hypothesized that a tumorigenic action mediated by the immunosuppressive effect of alcohol on natural killer cells could explain this observation. In fact, in the non-immunosuppressed population, alcoholism is associated with an increased risk for several malignancies, including liver and alimentary tract tumors\textsuperscript{[107,108]}. Jain et al\textsuperscript{[95]} observed a higher rate of de novo oropharyngeal and pulmonary neoplasms in transplanted ALD patients than in those with a non-alcoholic disease, similar to Duvoux et al\textsuperscript{[109]}, suggesting the presence of other initiators of malignant changes in addition to immunosuppression. Among the identified risk factors, alcohol and tobacco consumption were highlighted\textsuperscript{[90,99,109]}, data also obtained in our study\textsuperscript{[99]}. Regarding tobacco, nearly 90% of alcoholics smoke\textsuperscript{[110]}, compared to 26.7% of the general population of the United States\textsuperscript{[111]}. Regarding the candidates for LT, approximately 60% are smokers\textsuperscript{[112,113]} and 15%-40% continue to smoke after the LT\textsuperscript{[112,114]}. DiMartini et al\textsuperscript{[114]} found that nearly 40% of ALD recipients resume smoking and resume it early post-LT, increase their consumption over time and quickly become tobacco dependent. In a recent meta-analysis, active smoking was revealed as one of the major risk cofactors, independent of alcoholic relapse, of long-term morbidity and mortality in transplant recipients, either from cardiovascular complications or from de novo neoplasms\textsuperscript{[80]}. This data was confirmed in numerous studies\textsuperscript{[36,109,114-116]}. In an interesting study from a conceptual point of view, Herrero et al\textsuperscript{[115]} showed that smoking withdrawal after LT may have a protective effect against the development of neoplasia. In particular, these researchers observed that patients with a smoking history who continued smoking after the LT, presented a hazard ratio of approximately 20 for the development of neoplasms associated with tobacco (head and neck, lung, esophagus, kidney and urinary tract carcinomas), while the risk of developing these neoplasms was reduced significantly in ex-smokers\textsuperscript{[116]}. Furthermore, as noted earlier, smoking is also a risk for cardiovascular disease, and this is one of the most frequent causes of late mortality after LT\textsuperscript{[106,123]}. Pungpapong et al\textsuperscript{[117]} found a higher rate of vascular complications in LT recipients who had a history of smoking. Those who quit smoking 2 years prior to the transplantation reduced the incidence of vascular complications by 58%. Therefore, our pre- and post-LT follow-up efforts regarding ALD should be focused not only on alcoholic relapse but also on treating and avoiding other modifiable risk factors such as tobacco, not simply because of what was discussed earlier, but because we now acknowledge that tobacco is a risk factor for alcohol abuse. In a recent study on mice, it was observed that the rodents exposed to nicotine tended to ingest alcohol more frequently than those that were not administered such a substance due to a reduction in the dopamine response of the reward-response system in the brain, which thus decreased the pleasurable response to alcohol\textsuperscript{[124]}. Given all the above-mentioned observations, the establishment of control programs and post-LT interventions could perhaps reduce the mortality in these patients, as will be discussed below.

**POST-LT FOLLOW-UP IN ALD**

Several approaches have been evaluated to reduce alcohol recidivism in alcoholic patients after LT, but there is no standardized approach, and the available data are few and often controversial. In some liver transplant centers, alcoholic patients are encouraged to attend support groups, even if the data demonstrating the efficacy of such treatment in this cluster of patients are currently lacking. In a pilot study, Georgiou et al\textsuperscript{[125]} reported that psychological interventions could be a valid approach to enhance motivation in these patients. However, this study was con-
ducted on a limited number of patients, and the efficacy of this intervention on alcohol recidivism after LT was not evaluated. Björnsson et al.\[35\] evaluated the impact of the management of alcoholic patients by addiction psychiatrists, social workers and tutors in the period before LT and reported a 22% prevalence of alcohol recidivism in the treated group vs 48% in the untreated group. The presence of an alcohol addiction unit within a liver transplant center is not usual, but the study of Addolorato et al.\[126\] suggests that it could represent a useful approach to reducing alcohol recidivism after LT. However, objective and accurate indicators of abstinence are required\[127\]. Direct detection in the blood or breath only assesses alcohol intake within the preceding 10-12 h\[39,128\]. Carbohydrate-deficient transferrin (CDT) is an indirect marker that reflects alcohol intake in the previous 1-4 wk\[129,130\]; however, the daily consumption of 60-89 g of ethanol for a period of at least 7-10 d is required for a positive result. Therefore, CDT is inappropriate for the detection of low-to-moderate alcohol intake. Furthermore, wide ranges in sensitivity and specificity of 46%-73% and 70%-100%, respectively, have been reported\[131\]. Nevertheless, a high rate of false-positives with the CDT test has been reported, particularly in patients with severe liver damage\[131\].

Currently, the determination of ethyl glucuronide (EtG), a metabolic product of alcohol, either in the urine or in the hair of patients offers a new, reliable possibility for the detection of alcohol intake\[132-135\]. Urinary EtG (uEtG) remains positive for up to 80 h after alcohol consumption and allows for the detection of very small amounts of ethanol (uptake of < 5 g)\[133,135\] with a sensitivity and specificity of 89% and 99%, respectively\[131\]. However, positive uEtG tests may occur after the accidental consumption of foods containing alcohol, such as chocolate, cake and others. To reduce this problem in the transplant setting, a higher cut-off level for uEtG than what is routinely used (> 0.5 mg/L instead of > 0.1 mg/L) is recommended\[132\].

Furthermore, the detection of EtG in the scalp hair of patients is a powerful tool for monitoring abstinence over a retrospective period of up to 6 mo. Each hair segment of 1 cm in length reflects alcohol consumption over a period of approximately 1 mo. The test has been validated for a maximal hair length of 6 cm\[135\].

Thus, based on the above, we can infer that regular monitoring after LT is critical for determining the ongoing abstinence from tobacco and alcohol and for providing treatment assistance when tobacco or alcohol use are identified. Using a combination of methods (patient interviews by a trained alcoholism and addiction professional connected to the transplant team, independent caregiver reports and biochemical monitoring) provides the greatest yield because every method can add to the number of identified cases\[27\].

As we have discussed previously, patients who undergo LT have an unexpectedly high rate of de novo extrahepatic cancer\[98,99\], including lung and upper aerodigestive tract cancer\[100,102\], and studies have reported that patients with ALD are particularly affected\[98,99,102\]. Therefore,
apart from identifying and acting on predisposing factors for the development of neoplasms (such as tobacco use, solar exposure and infections from oncogenic viruses), intensive tumor screening protocols have been suggested for these patients. Herrero et al\(^9\) concluded that ALD transplant patients, smokers or ex-smokers, should have a further follow-up, including a low-radiation-dose thorax computed tomography (CT) scan, a consultation with an ear, nose and throat specialist and a urine exam, as suggested by Benlloch et al\(^16\), who recommend an annual head and neck cancer screening due to the high risk of this type of cancer. However, only two studies have shown that intensive screening protocols increase survival\(^7,8\). Therefore, at the present time, patient education, mainly to avoid smoking and sun exposure, and periodic clinical follow-ups continue to be the standard of care regarding treatment\(^9\).

**CONCLUSION**

In the last 10 years, ALD is the LT indication that has seen the greatest increase in prevalence\(^9\) as well as in post-LT survival rate compared with other causes of liver disease, although concerns over alcoholic relapse remain. Even though less than 5% of grafts are rejected at 5 years post-LT due to a direct or indirect consequence of alcohol consumption\(^13\), transplanted ALD patients who relapse have an increased long-term mortality due to cardiovascular pathologies and the onset of *de novo* neoplasms.

Much has been discussed regarding the risk factors for relapse, and the most controversial has been, and continues to be, the pre-LT abstinence period. In view of the foregoing, we can say that this period by itself should not be a determining factor to include a patient on the list because many other factors exist; therefore, a good psychiatric and psychosocial evaluation that identifies and addresses such factors before and after the LT is important\(^10,14\).

The major incidence of *de novo* neoplasms in this type of patient could be remedied with the detection of and action on the predisposing factors for the development of neoplasms in addition to the development of more intensive programs for the detection of neoplasm; however, the efficacy of this approach must be demonstrated. What we conclude is that the pre-LT evaluation and the post-LT follow-up in ALD patients should be a multidisciplinary task that includes transplant specialists, psychiatrists and addiction treatment specialists (Figure 2).

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