The control of abstinence in the treatment of alcohol dependence: the use of acamprosate in relapse prevention

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Abstract

Treatment with acamprosate is a valid tool to complement psychotherapy as it does not cause addiction, abuse or withdrawal of its suspension and does not interfere with other medications that patients often holics must take. To evaluate the effectiveness, our study evaluated the effects of Acamprosate compared to γ-hydroxybutyrate in clinical-physiological and social health in a way indicator of a possible therapeutic success in terms of abstinence from alcohol and social reintegration. A total of 36 patients were observed, of which 5, 4 men and 1 woman at the Drug Addiction Service (Servizio Tossicodipendenza, Ser.T) Alcamo, and 31, 21 men and 10 women at the Ser.T of Palermo. As regards the craving, during the period of treatment with acamprosate, there has been a change, in the sense of reduction, of craving for alcohol: if before therapy was in 68% of cases, medium-high, it becomes after 3-4 months of therapy in low-nil in 89% of patients observed. It has been recorded that, after 3-4 months after receiving acamprosate, the clinical picture of the patient is greatly improved by referring to biological markers. In particular, the strong point seems to be the ability for the user to experience a new sense of normalcy and to remove the desire for significant periods of alcohol.

Introduction

Alcoholism can also deal with drug treatments. This is the message that emerges from the press conference of presentation of Campral©, trade name of acamprosate, a specifically neuromodulator indicated in the maintenance of abstinence in alcohol-dependent patients.1 Alcoholism is a disease characterized by: craving, loss of control, tolerance and physical dependence.2 Aim of the study was to clinically prove efficacy, influence and usefulness of acamprosate in reducing craving during abstinence in alcoholics. To evaluate the real effectiveness, our study evaluated the effects of acamprosate, compared to γ-hydroxybutyrate (GHB). Through an antagonistic action of glutamatergic hyperactivity that characterizes alcohol abstinence, acamprosate reduces the craving, the incidence and severity of relapses in alcoholic patients.3 It is available in tablets of 333 mg and the recommended dosage is 2 tablets 3 times a day. Concomitant use of alcohol has no effect on the acamprosate pharmacokinetics, suggesting that patients can continue treatment even during relapse periods without health risks.4 There are no signs of tolerance, dependence, withdrawal syndrome; the percentages of relapses after stopping treatment are very low. Unlike other antacrating drugs, acamprosate presents an over time growing effectiveness.

Materials and Methods

The structures that have joined the study were Drug Addiction Service (Servizio Tossicodipendenza, Ser.T) of Alcamo and the Palermo’s district 11. In total, from May 2011 to March 2012 36 patients were observed, of which 5 (4 men and 1 woman) at the Sert of Alcamo and 31, including 21 men and 10 women at the Ser.T of Palermo. The patients for whom we have adopted the treatment with acamprosat are 22.2%. General data such as sex, age, education, employment and clinical data and the possible positive and/or the pattern of diseases related to addiction have been reviewed. Of the sample, 69.5% were male, while women represented only 30.5%. Of the sample, 52.8% were aged between 35 and 50 years, 30.5% over 50 years, while only 16.7% of the sample between 25-35 years. Patients already attending the Ser.T are treated with GHB and are subjected for the first time treated with acamprosate, and have different expectations of the drug: succeed to terminate therapy with GHB, stop the use of alcohol, overcome the withdrawal symptoms and cease the use of drugs. Of patients, 58.3% revealed frequent and intense relapses during treatment with GHB. Before starting treatment with acamprosate all patients underwent medical and toxicological assessment (clinical examination, medical history and general toxicology, urinalysis, blood alcohol test, psychological examination) and were informed about the acamprosate characteristics and action mechanism.
Results

From data obtained by medical records of patients already being treated with GHB peaks of aspartate amino-transferase (AST) of 920 U/L, alanine transaminase (ALT) 670 U/L and γ-glutamyltransferase (GGT) 1700 U/L, we encountered. Weekly, through medical examination and an interview with a psychologist, we evaluated the clinical course of the patient, the patient’s response to acamprosate, any impact resulting from the reduction of the use of alcohol by the use of GHB and possible side effects. Permanently suspended the use of GHB, the patient continued treatment with only acamprosate: to this point we have carried out new surveys, recording the data obtained after 3-4 months from suspension of GHB and after the end of therapy. The percentage of relapse during treatment with GHB was 55.6%, while with acamprosate only 13%. According to the latest findings obtained after 6-8 months after treatment with acamprosate, the marker’s values are returned in standard: the average of AST was reduced from 77 to 17 U/L, those of ALT 43.3 to 21 U/L, and that of GGT from 356.8 to 61 U/L. Almost all the sample believed acamprosate subjectively more effective than GHB (obviously those who have experienced the two drugs).

The reason for the preference of acamprosate to GHB is given by the lesser dependence towards the drug and the alcohol decreased desire. The majority of patients experience a more relaxed, understood as a reduction in anxiety, nervousness as well as increased ability to concentrate. The alcohol craving during treatment with acamprosate decreased in subjects clearly treated: if before treatment it was medium-high in 68% of cases, it became a low-void 3-4 months in 89% of patients observed. The success rate of treatment with acamprosate was very high at 86%, but should be considered an abandonment of therapy (14%) caused by the appearance of side effects such as bloating medium-high (1 case), ulcerative (1 case), sexual dysfunction (2 cases), and psychiatric disorders (that may already exist).

Discussion

Acamprosate is a drug that acts on the glutamatergic system; it seems useful not only in the control of mood but also in improving the withdrawal symptoms and decrease the number of relapses. Through an antagonistic action of glutamatergic hyperactivity that characterizes abstinence from alcohol, acamprosate reduces the craving, the incidence and severity of relapse in alcoholics subjects. Acamprosate is also useful to reduce or cancel the appearance of acquired stimuli leading to the alcohol dependence in time: in this sense inhibition of glutamatergic transmission by the acamprosate is able to reduce the severity of abstinence’s syndrome. On the other hand, its use must have a sufficient duration to allow neuronal excitability normalization in the most enduring possible: the treatment, in fact, is recommended for one year. In any case, the use can be continued even in the face of relapses, with the aim to reduce the frequency or severity.

Conclusions

The hope for the future is that a growing experience by clinicians can facilitate, over time, an optimal selection of suitable patients for treatment with acamprosate.

References