Medications To Treat Alcoholism

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Advances in neurobiology support the development of medications to treat alcoholism by modifying the activity of specific chemical messengers (i.e., neurotransmitters) in the brain. Among the most promising new medications is acamprosate, which appears to decrease the intensity of craving after a person has stopped drinking. Naltrexone (ReVia™) has been shown to decrease alcohol consumption, although its practical effectiveness may be compromised by poor patient compliance and other factors. Ondansetron shows promise for decreasing drinking and increasing abstinence rates among early onset alcoholics, who respond poorly to psychosocial treatment alone. Researchers are investigating whether the use of specific medications in combination can further enhance their effectiveness. Additional research is needed to determine how medications interact with different psychosocial factors and treatments.

KEY WORDS: drug therapy; AOD (alcohol and other drug) dependence; calcium acetylhomotaurinate; naltrexone; AOD craving; AOD abstinence; drug efficacy; antagonists; opioids; glutamate; serotonin; serotonin uptake inhibitors; dopamine; serotonin receptors; buspirone; combination drug therapy; patient compliance; literature review

Although psychosocial therapies help many alcohol-dependent persons reduce alcohol consumption and maintain abstinence, 40 to 70 percent of patients resume drinking within 1 year of such treatment (Swift 1999). Advances in neuroscience research in the past decade suggest the possibility of developing medications to improve the efficacy of concurrent psychological or behavioral addiction therapies (Litten et al. 1996). These advances include research that implicates specific neurotransmitter systems (see textbox, p. 100) in the development of addiction, suggesting that medications that modify the activity of these systems may interfere with the development of alcoholism. In addition, some alcoholics may be biologically predisposed to alcohol dependence. Those persons may benefit from medications designed to correct or ameliorate the biochemical abnormalities that presumably underlie their susceptibility.

This article reviews recent efforts to develop medications to both reduce the desire to drink and promote abstinence. Each section begins with a brief explanation of the probable role played by a specific neurotransmitter system in the development of addiction to alcohol and other drugs (AODs). These neurotransmitter systems include opioids, glutamate, serotonin (5HT), and dopamine. Medications that affect the function of these neurotransmitters to influence alcoholism are discussed, focusing on those medications that have shown the most promise in clinical trials on human subjects. Finally, the article recommends areas for future research.

Opioid Antagonists

Opioid peptides are a class of neurotransmitters that produce physiological effects similar to those of morphine and

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histories of alcoholism (i.e., high-risk alcoholic relatives, subjects with family consumption of two drinks per day).

In a recent study (King et al. 1997), genetic susceptibility to alcoholism. Litten and Allen (1998). One factor producing alcoholism have lower levels of tension, fatigue, and confusion.

Antagonists at the mu opioid receptor may also affect alcohol consumption by suppressing general consummatory behaviors without altering motivational or approach behaviors (for technical terms not defined in the text, see glossary, pp. 100) (Boyle et al. 1998). In addition, researchers (see Gianoulakis 1998) have proposed other mechanisms for the effects of opioid peptides on alcohol consumption, including modulation of the body’s hormonal stress response.

Human laboratory studies exploring the effects of naltrexone on alcohol-induced mood and craving have produced equivocal results (reviewed in Litten and Allen 1998). One factor that may contribute to the discrepancy among research results is a person’s genetic susceptibility to alcoholism. In a recent study (King et al. 1997), nonalcoholic social drinkers (average consumption of two drinks per day) consumed an alcoholic beverage 3 to 4 hours after taking naltrexone. Compared with subjects who did not have any alcoholic relatives, subjects with family histories of alcoholism (i.e., high-risk subjects) experienced less of alcohol’s stimulant effects but reported increased tension, fatigue, and confusion.

Consistent with the above results, persons at genetically high risk for developing alcoholism have lower levels of β-endorphin and demonstrate a more pronounced increase in β-endorphin levels in response to alcohol administration compared with persons who do not have alcoholic relatives. Similarly, naltrexone’s propensity to reduce alcohol intake in humans is greater in persons who have higher beta-endorphin levels (Gianoulakis et al. 1996).

The Food and Drug Administration has approved naltrexone for the treatment of alcohol dependence based largely on evidence from two randomized, double-blind, placebo-controlled studies. In the first of these studies (Volpicelli et al. 1992), recently abstinent male patients undergoing psychosocial alcoholism treatment who received 50 mg/day of naltrexone1 for 12 weeks reported less craving than subjects who received psychosocial treatment alone. In addition, the patients who received naltrexone maintained abstinence for longer periods and exhibited significantly reduced relapse rates than the patients who did not receive naltrexone (i.e., only 23 percent of naltrexone-treated patients compared with 54 percent of placebo-treated patients experienced relapse). Among patients who sampled alcohol while still in treatment, only 50 percent of the naltrexone-treated patients relapsed, compared with 95 percent of the patients who received a placebo (Volpicelli et al. 1992).

O’Malley and colleagues (1992) obtained comparable results in the second study using a similar design but including both male and female alcoholic subjects. An additional important finding in this study was the interaction between naltrexone use and the type of psychosocial therapy employed. The cumulative rate of abstinence was highest for patients who received naltrexone in conjunction with generalized supportive psychotherapy. However, the combination of naltrexone and a more intensive therapeutic program designed to teach specific coping skills was more effective at preventing relapse among patients who sampled alcohol.

Further research has demonstrated that naltrexone-treated abstinent alcoholics report diminished craving for alcohol (O’Malley et al. 1996) and those who sample alcohol after a period of abstinence experience less mood elevation from the alcoholic beverage than non-naltrexone-treated patients (Volpicelli et al. 1995). Unfortunately, the protective effects of naltrexone appear to dissipate over time; O’Malley and colleagues (1996) found no difference in abstinence rates between naltrexone- and placebo-treated patient groups 5 months after naltrexone treatment was terminated.

The major obstacle to the practical clinical effectiveness of naltrexone appears to be poor patient compliance. In a 3-month followup study, Volpicelli and

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1This is the standard dose of naltrexone (under the brand name Trexan®) used for treating opiate addiction.
colleagues (1997) found modest improvement in alcohol consumption (measured as percent drinking days) only among patients who took their daily dose of naltrexone on at least 90 percent of the study days. The large-scale United Kingdom Collaborative study of Chick and co-workers (reviewed in Litten and Allen 1998) found no significant differences between naltrexone and the placebo on various measures of alcohol consumption over 12 months except among patients who exhibited at least an 80-percent compliance rate. The level of supervision necessary to ensure patient compliance may be difficult to achieve outside of specialized treatment facilities.

Further large-scale clinical trials are needed to establish naltrexone’s clinical effectiveness. Research also is needed to determine if naltrexone would be more effective at doses greater than 50 mg/day or if the treatment period were prolonged beyond 12 weeks of daily medication. In addition, particular subpopulations or types of alcoholics may be differentially responsive to naltrexone, especially alcoholics with strong family histories of alcoholism. For example, higher levels of craving measured when treatment was initiated were associated with greater treatment response (Jaffe et al. 1996), suggesting that naltrexone may be more effective in patients with the most severe alcoholism.

**Specific Glutamate Antagonists**

The excitatory neurotransmitter glutamate, acting largely at the N-methyl-D-aspartate (NMDA) receptor, may contribute to some of alcohol’s effects (e.g., intoxication, cognitive impairment, and some symptoms of withdrawal). The NMDA-receptor antagonist acamprosate appears to reduce the intensity of craving after drinking cessation, particularly when the patient is exposed to situations or environments associated with previous alcohol use, where the risk of relapse is greatest (Spanagel and Ziegglansberger 1997).

Acamprosate has been shown to decrease alcohol consumption in rats without interfering with total fluid and food intake (Litten et al. 1996). Few laboratory studies have directly tested acamprosate’s effects on alcohol-related behaviors in humans. However, a magnetic resonance imaging (MRI) study showed reduced glutamatergic activity in certain brain regions in humans following acamprosate administration (Bolo et al. 1998), providing support for the therapeutic potential of acamprosate in the treatment of alcoholism.

Large-scale clinical studies support the effectiveness of acamprosate in treating alcoholism. An analysis of pooled data from a series of randomized, double-blind, placebo-controlled trials at several centers throughout Europe involving more than 3,000 alcoholic subjects indicated that twice as many patients remained abstinent from alcohol for up to 1 year after treatment compared with patients who received only a placebo (Swift 1999). For example, in a 12-week study, Lhuinire and colleagues (1990) showed that acamprosate (1.3 g/day) was more efficacious than placebo in the treatment of recently detoxified alcoholics, with the greatest improvement observed in the most severe alcoholics. Drawbacks of this study included the lack of standardized criteria for diagnosing alcoholism and the use of non-standardized blood chemistry analyses to detect recent alcohol consumption.

In a similar study, Pailie and colleagues (1995) found acamprosate at doses of both 2.0 g/day and 1.3 g/day to be superior to a placebo in the treatment of alcohol-dependent patients who drank an average of approximately 17 standard drinks per day. The percent of patients continuously abstinent was highest among those taking 2 g/day of acamprosate, lowest among those taking a placebo, and at an intermediate level among patients taking 1.3 g/day of acamprosate. The major improvements in this study, compared with that of Lhuinire and colleagues (1990), was the use of standardized diagnostic criteria to identify alcoholism and better laboratory standardization for monitoring alcohol consumption by blood chemistry, despite the relative insensitivity of such methods as an indicator of transient alcohol consumption.

In some trials, the 2-to-1 ratio of abstinence rates among acamprosate-treated patients compared with placebo-treated patients persisted for at least 1 year following cessation of treatment, although overall relapse rates had increased during the interval. At the end of 48 weeks of treatment, Sass and colleagues (1996) found abstinence rates of 43 percent among acamprosate-treated subjects compared with 21 percent among placebo-treated subjects. At the end of an additional 48 weeks without medication, abstinence rates were 39 percent and 17 percent, respectively (Sass et al. 1996). Similar results of effectiveness for acamprosate in the treatment of alcoholics have been reported by others (Whitworth et al. 1996; Geerlings et al. 1997).

**Serotonergic Medications**

Serotonin plays a role in regulating diverse physiological functions, including body rhythms, sleep, and mood states, as well as appetitive and consummatory behaviors (Lovinger 1997). At least 15 subtypes of the serotonin receptor exist, some of which have potential roles in the development of alcohol-use disorders through multiple interactions with other neurotransmitter systems. The effects of serotonergic agents in the treatment of alcoholism have been well studied over the past two decades. This section examines the effects of general and specific serotonergic medications that show promise for alcoholism treatment.

**Selective Serotonin Reuptake Inhibitors**

Selective serotonin reuptake inhibitors (SSRIs) are medications that interfere with the removal of serotonin from the synapse following release of the neurotransmitters (see textbox). Various
SSRIs are used to treat psychiatric conditions, including mood disorders (e.g., depression) and anxiety disorders. The effects of SSRIs are not limited to specific serotonin-receptor subtypes.

Alcohol consumption has been shown to alter serotonin-receptor function in humans and animals, and early studies have indicated that administration of SSRIs can suppress alcohol consumption in animals (Lovinger 1997). Activation of specific serotonin-receptor subtypes can stimulate dopaminergic activity in the nucleus accumbens (see glossary, p. 106), thereby potentially contributing to alcohol's rewarding effects (Lovinger 1997). However, the relationship between serotonin-reuptake inhibition, food intake, and fluid consumption is complex (Gill and Amit 1989). Thus, the effect of SSRIs on alcohol consumption may be attributable to generalized decreases in consummatory behavior rather than to a reduction in alcohol's reinforcing effects (Swift 1999).

Despite encouraging results of animal studies (Lovinger 1997), the findings of clinical trials using SSRIs for the treatment of alcoholics without co-occurring psychiatric disorders have been equivocal. Although earlier studies showed that SSRIs can reduce alcohol consumption in alcohol abusers (Naranjo et al. 1990, 1992; Naranjo and Sellers 1989), more recent long-term studies have not confirmed these results (Gorelick and Paredes 1992; Kranzler et al. 1995). Recent studies are mostly negative, although some studies suggest that SSRIs treated with buspirone may exhibit some improvement in both craving and drinking behavior (reviewed in Malec 1996).

Predicated on the findings of human laboratory studies showing that alcoholics with an early onset of disease may be deficient in serotonin (5HT) (Buydens-Branchey et al. 1989; Fils-Aime et al. 1996), Kranzler and colleagues (1996) examined the effects of the SSRI fluoxetine (Prozac) on alcohol drinking among this subtype of alcoholic. The researchers presumed that fluoxetine would compensate for the functional deficit in early onset alcoholics, thereby resulting in a cessation of alcohol consumption. Surprisingly, however, Kranzler and colleagues (1996) found that fluoxetine appeared to increase alcohol consumption among early onset alcoholics, suggesting that the relationship between serotonergic dysfunction and the onset of alcoholism is neither simple nor direct.

Although a review of pharmacological agents for the treatment of alcoholics with co-occurring psychiatric disorders is outside the scope of this article, evidence suggests that SSRIs are effective for treating alcoholics with co-occurring depression. In the first controlled double-blind trial of an SSRI in alcoholics with co-occurring depression, Cornelius and colleagues (1997) found that both depressive symptoms and the level of alcohol consumption decreased significantly in patients who were administered fluoxetine compared with patients who were taking a placebo.

Medications That Affect Specific Serotonin-Receptor Subtypes

Buspirone. Serotonin receptors of the subtype designated 5HT, may be involved in general consummatory behavior as well as the development of alcohol intoxication and alcoholism (Lovinger 1997). Buspirone (Buspar), which has partial agonist activity at the 5HT receptor, is commonly prescribed to treat anxiety disorders. Findings of reduced drinking in rats (Lovinger 1997) have led to clinical studies in human alcoholics. Results of controlled human studies are mostly negative, although some studies suggest that alcoholics treated with buspirone may exhibit some improvement in both craving and drinking behavior (reviewed in Malec 1996). Evidence suggests that buspirone's mechanism of action in reducing alcohol-related behaviors may result from reductions in anxiety, rather than through reduction in the rewarding effects of alcohol per se, and that any potential benefit from this medication is probably limited to alcoholics with independent co-occurring anxiety disorders (Malec et al. 1996).

Ritanserin. Serotonin 5HT receptors may play a role in alcohol's rewarding effects and also contribute to the development of acute physical withdrawal symptoms (Lovinger 1997). Laboratory research demonstrates that certain 5HT antagonist can significantly suppress alcohol intake by rats without affecting water consumption, although some of these medications can affect food consumption (Myers and Lankford 1998; Overstreet et al. 1997). Antagonists at the 5HT receptor have also been shown to reduce alcohol intake by modulating the effects of dopamine in the nucleus accumbens (Lovinger 1997).

The effectiveness of ritanserin as a treatment for alcohol dependence was recently tested in a rigorous multicenter, placebo-controlled clinical trial in which 423 alcoholics received up to 5 mg/day concurrently with cognitive behavioral therapy. All study participants had significantly reduced their alcohol consumption by the end of the study period. However, no difference in drinking level was observed between patients receiving ritanserin and those who had received the placebo. Although higher ritanserin doses may have been more effective, high doses of 5HT receptor antagonists may increase the potential for adverse side effects, particularly irregular heartbeat (Johnson et al. 1996).

Ondansetron. The most promising serotonergic medication for treating alcoholism appears to be the 5HT antagonist ondansetron. Animal studies demonstrate that the 5HT receptor enhances the rewarding effects of alcohol, perhaps in part by increasing the alcohol-induced release of dopamine in the nucleus accumbens (Lovinger 1997). Consistent with this hypothesis, antagonists at the 5HT receptor have been shown not only to block dopamine activity but also to reduce alcohol intake in a variety of animals (for a review see Johnson and Cowen 1993).

In humans, ondansetron reduced some of alcohol's positive subjective effects including self-reported desire to drink (Johnston and Cowen 1993; Swift et al. 1996). In a 6-week preliminary clinical trial, Sellers and colleagues (1994) found that 0.5 mg/day, but not 4 mg/day, was efficacious in treating moderate alcohol abusers. Researchers (Johnson et al. 1999) recently have presented a much more effective, high doses of 5HT receptor antagonists may increase the potential for adverse side effects, particularly irregular heartbeat (Johnson et al. 1996).

Depressed mood can occur as a symptom of alcoholism. Each subject in this study met standard diagnostic criteria for co-occurrence of both alcoholism and a clinical depressive disorder independently.
larger double-blind clinical trial of ondansetron’s effectiveness as adjunctive treatment to cognitive-behavioral therapy in both early and late onset alcoholics. Preliminary analysis of results from that study showed that ondansetron was significantly superior to a placebo in reducing alcohol drinking and increasing abstinence rates among early onset but not late onset alcoholics. Previous data have shown that earlier age of onset for alcoholism is associated with certain biochemical measures thought to indicate serotonin dysfunction (Swann et al. 1999). This finding is consistent with literature positing serotonin dysfunction among early onset alcoholics (Buydens-Branchey et al. 1989; Linnola and Virkkunen 1994). It is, therefore, plausible that ondansetron’s treatment effect may be associated with its ability to ameliorate serotonergic abnormality in this subtype of alcoholic (see figure below).

Intriguingly, compounds that facilitate serotonin function, even though neurochemical analyses suggest a serotonin-deficient state, either have no effect or may worsen drinking behavior of early onset alcoholics (Kranzler et al. 1994a,b).

### Dopamine Antagonists

Although medications that directly affect dopamine function would appear hypothetically to be promising candidates for alcoholism treatment, experimental results have not been encouraging (Di Chiara 1997). In animals, a variety of dopamine antagonists, such as the antipsychotic medication haloperidol (Haldol®) and tiapride, were shown to attenuate low-dose alcohol-induced hyperactivity, whereas higher doses of alcohol were associated with sedation. In human alcoholics, tiapride was demonstrated in a randomized, placebo-controlled clinical trial to be superior to a placebo in increasing abstinence, self-esteem, and life satisfaction. Drinking levels also were significantly lower in the tiapride-using patients compared with the placebo group (Shaw et al. 1994). However, similar to other dopaminergic medications, an important problem with these medications is their potential to produce neurological side effects such as movement disorders (Tamion et al. 1990). Earlier research suggests that tiapride may be useful in alleviating alcohol-withdrawal symptoms (Swift 1999). However, tiapride may increase the likelihood of seizures, a common complication of withdrawal, and would probably need to be prescribed with anticonvulsants (Delmeire 1980). Thus, the side-effect profile of dopamine antagonists may limit their widespread use.

### New Vistas

### Opioid Antagonists

The mu receptor antagonist nalmefene has been shown to increase abstinence in a preliminary placebo-controlled, double-blind study of 21 alcoholics (Mason et al. 1994). Nalmefene is also less likely than naltrexone to produce the adverse side effect of liver damage. Consequently, researchers and clinicians anxiously await results of subsequent large-scale clinical trials of nalmefene in alcoholic populations.

Opioid-peptide receptors other than the mu receptor have been implicated in some alcohol-related behaviors (Herz 1997). Research suggests that antagonists of the delta opioid receptor (e.g., naltriben and naltrindole) may suppress alcohol consumption in animals (Froehlich et al. 1991). Further animal and human studies are needed.

### Calcium Channel Antagonists

Calcium has many physiological functions throughout the body and is required for nerve cell activity. Medications that interfere with the passage of calcium into cells (i.e., calcium channel blockers) are prescribed to treat high blood
Combination Treatments

Interactions among neurotransmitter systems suggest that combining therapeutic agents for the treatment of alcoholism may be an effective strategy. One potentially promising treatment would combine acamprosate and naltrexone. While naltrexone reduces craving derived from alcohol’s positive rewarding effects (Volpicelli et al. 1995), acamprosate diminishes craving associated with the period following acute withdrawal (Spanagel and Ziegglinsberger 1997). Thus, the combination of naltrexone and acamprosate might not only make it easier for a patient to abstain from alcohol use but also might prevent a“slip” from turning into a relapse. Several important issues must be resolved with any medication combination, however, including optimal dosing and timing, duration of treatment, and management of potential adverse interactions. NIAAA is currently funding clinical trials to establish the effectiveness of naltrexone and acamprosate, both alone and in combination, for the treatment of alcoholism.

Conclusions

Of the medications reviewed, acamprosate’s potential appears to be the most widely established, although naltrexone remains the approved choice in the United States. Results of ongoing clinical trials suggest that naltrexone and acamprosate, both separately and in combination, may usher in a new era of medication treatments with multiple neurotransmitter targets. In addition, preliminary data suggest that ondansetron offers new hope for the treatment of early onset alcoholics, who generally tend to respond poorly to psychosocial treatment alone.

Nonetheless, important challenges remain. Based on the realization that alcoholics constitute varying subtypes with differing biological and psychosocial contributions to the disease, researchers must rigorously test medication development among these subgroups. Differential subgroup efficacy would not only provide an important therapeutic advantage to those who might benefit but also could offer information on the neurochemical factors mediating that aspect of the disease. Optimally, this would enable medications to be targeted toward the type of alcoholics who would most benefit from their use.

Additionally, few pharmacotherapy studies vary the dose of the medication studied. Without dose-ranging studies, the response ceiling may not be achieved nor would the benefits to adverse event relationship be characterized thoroughly. This could also lead to premature discarding of a potential therapeutic medication simply on the basis that the chosen dose was ineffective. Finally, the development of neuroimaging and molecular techniques promise new tools for both characterizing and charting the progress of disease and for understanding the site-specific actions of medications associated with treatment response. The coming decade promises more powerful tools for characterizing drug effects and for closing the gap between animal models of addiction and the human condition.

References

Boile, N.; Nedelec, J.F.; Muzet, M.; De Witte, P.; Daigrou, A.; Durbin, P.; and Machet, J.P. Central effects of acamprosate: Part 2. Acamprosate modifies the brain in-vivo proton magnetic reso-
nance spectrum in healthy young male volunteers. Psychiatry Research 82:115–127, 1998.

Boile, A.E.L.; Stewart, R.B.; Macenski, M.J.; Spiga, R.; Johnson, B.A.; and Mesich, R.A. Effects of acute and chronic doses of naltrexone on ethanol self-administration in rhesus monkeys. Alcoholism: Clinical and Experimental Research 22(2):359–366, 1998.

Buydens-Branchey, M.; Branchey, M.; and Noumair, D. Age of alcoholism onset, I. Relationship to psychopathology. Archives of General Psychiatry 4:225–230, 1989.

Cornelius, J.R.; Salloum, L.M.; Ehrler, J.G.; Jarrett, P.J.; Cornelius, M.D.; Perel, J.M.; Thase, M.E.; and Black, A. Fluoxetine in depressed alcoholics: A double-blind, placebo-controlled trial. Archives of General Psychiatry 54(8):701–705, 1997.

De Beun, R.; Loehmann, A.; and De Vry, J. Conditioned taste aversion and place preference induced by the calcium channel antagonist nimodipine in rats. Pharmacology, Biochemistry & Behavior 54:657–663, 1996.

Delmeire, F. Taptridal. Acta Psychiatria Belgica 80:191–201, 1980.

De Chiaro, G. Alcohol and dopamine. Alcohol Health & Research World 21(2):108–114, 1997.

Fils-Aime, M.L.; Eckardt, M.J.; George, D.T.; Brown, G.L.; Mefford I.; and Lennola M. Early-onset alcoholics have lower cerebrospinal fluid 5-hydroxyindoleacetic acid levels than late-onset alcoholics. Archives of General Psychiatry 53(3): 211–216, 1996.

Freuhi, J.C. Opoid peptides. Alcohol Health & Research World 21(2):97–136, 1997.

Freuhi, J.C.; Zweifel, M.; Harts, J.; Lumen, L.; and Li, T.K. Importance of delta opioid receptors in maintaining high alcohol drinking. Psychopharmacology 103:467–472, 1991.

Geeelho, P.J.; Ansons, G.; and Van den Brink, W. Acamprosate and prevention of relapse in alcoholics. European Addiction Research 3(3):129–137, 1997.

Gianoulakis, C. Alcohol-seeking behavior. The roles of the hypothalamic-pituitary-adrenal axis and the endogenous opioid system. Alcohol Health & Research World 22:202–210, 1998.

Gianoulakis, C.; De Waele, J.P.; and Thuandavayil, J. Implication of the endogenous opioid system in excessive ethanol consumption. Alcohol 13:19–23, 1996.

Gill, K., and Amit, Z. Serotonin uptake blockers and voluntary alcohol consumption. A review of recent studies. Recent Developments in Alcoholism 7:225–248, 1989.

Gorelick, D.A., and Paredes, A. Effect of fluorox-
tine on alcohol consumption in male alcoholics. Alcoholism: Clinical & Experimental Research 16(2): 261–265, 1992.

Herz, A. Endogenous opioid systems and alcohol addiction. Psychopharmacology 129:99–111, 1997.

Jaffe, A.J.; Rounsaville, B.; Chang, G.; Schot-
tenfeld, R.S.; Meyer, R.E.; and O’Malley, S.S. Naltrexone, relapse prevention, and supportive ther-
apy with alcoholics: An analysis of patient treatment.

Some calcium channel blockers used to treat high blood pressure include nifedipine (e.g., Adalat) and verapamil (e.g., Calan).
matching. Journal of Consulting & Clinical Psychology 64:1044–1053, 1996.

JOHNSON, B.A., AND COWEN, P.J. Alcohol-induced reinforcement: Dopamine and 5-HT3 receptor interactions in animals and humans. Drug Development Research 30:153–169, 1993.

JOHNSON, B.A.; JASINSKI, D.R.; GALLOWAY, G.P.; KRANZLER, H.; WEINREB, R.; ANTON, R.F.; MASON, B.J.; BOHN, M.J.; PETTINATI, H.M.; RAWSON, R.; CLYDE, C. Ritalin use in the treatment of alcohol dependency: A multi-center clinical trial. Psychopharmacology 128:206–215, 1996.

JOHNSON, B.A.; ROACHE, J.D.; JAVORS, M.A.; DI CLEMENTE, C.C.; CLONINGER, C.R.; PRIHODA, T.; BORDNICK, P.S.; AND ATT-DAOUD, N. Ondansetron and alcohol consumption: Preliminary analysis. Alcoholism: Clinical & Experimental Research 20:853–858, 1996.

LITTEN, R.Z.; ALLEN, J.; AND FERTIG, J. Psychopharmacology: A bridge for alcohol problems: A review of research with focus on developments since 1991. Alcoholism: Clinical & Experimental Research 20(5):859–876, 1996.

MALEC, T.S.; MALEC, E.A.; AND DONGIER, M. Efficacy of buspirone in alcohol dependence: A review. Alcoholism: Clinical & Experimental Research 20:853–858, 1996.

MASON, G.A.; REVZANI, A.H.; GRADY, D.R.; AND GABRUTT, J.C. The subchronic effects of the TRH analog TA-0910 and bromocriptine on alcohol preference in alcohol-prefering rats: Development of tolerance and cross-tolerance. Alcoholism: Clinical & Experimental Research 18:1196–1201, 1994.

MYERS, R., AND LANKFORD, M. Action of the 5-HT3 antagonist amperozide on alcohol-induced polikiloerha in rats. Psychopharmacology, Biochemistry & Behavior 59:91–95, 1998.

NARANJO, C.A., AND SELLERS, E.M. Serotonin uptake inhibitors attenuate ethanol intake in problem drinkers. Recent Developments in Alcoholism 7:255–266, 1989.

NARANJO, C.A.; KADLEC, K.E.; SANHUEZA, P.; WOODLEY-REMS, D.; AND SELLERS, E.M. Fluoxetine differentially alters alcohol intake and other consummatory behaviors in problem drinkers. Clinical Pharmacology & Therapeutics 47:490–408, 1990.

NARANJO, C.A.; POULOS, C.X.; BRENNER, K.E.; AND LANCOT, K.L. Citalopram decreases desirability, liking, and consumption of alcohol in alcohol-dependent drinkers. Clinical Pharmacology & Therapeutics 51:729–739, 1992.

O’MALLEY, S.S.; JAFFE, A.J.; CHANG, G.; SCHOTTELENDS, R.S.; MEYER, R.E.; AND ROUNSAVILLE, B. Naltrexone and coping skills training for alcohol dependence: A controlled study. Archives of General Psychiatry 49:881–887, 1992.

O’MALLEY, S.S.; JAFFE, A.J.; RODE, S.; AND ROUNSAVILLE, B.J. Experience of a “slip” among alcoholics treated with naltrexone or placebo. American Journal of Psychiatry 153:281–283, 1996.

OVERSTREET, D.; MCArTHUR, R.; REVZANI, A.; AND POST, C. Selective inhibition of alcohol intake in diverse alcohol-prefering rat strains by the 5-HT3 antagonists amperozide and FG 5984. Alcoholism: Clinical and Experimental Research 21:1484–1454, 1997.

PAILE, F.M.; GUELH, J.D.; PERKINS, A.C.; ROVER, R.J.; STERL, L; AND PAROT, P. Double blind randomized multicentre trial of acamprosate in maintaining abstinence from alcohol. Alcohol and Alcoholism 30:239–247, 1995.

RUSH, C.R.; AND PAZZAGLIA, P.J. Pretreatment with isradipine, a calcium-channel blocker, does not attenuate the acute behavioral effects of ethanol in humans. Alcoholism: Clinical & Experimental Research 22:539–547, 1998.

SASS, H.; SOYKA, M.; MANN, K.; AND ZIEGLÄNSBERGER, W. Relapse prevention by acamprosate. Results from a placebo-controlled study on alcohol dependence [published erratum appears in the December 1996 issue of Archives of General Psychiatry 53(12):1097]. Archives of General Psychiatry 53:673–680, 1996.

SELLERS, F.M.; TONEATTO, T.; ROMACH, M.K.; SOMER, G.R.; SOBELL, L.C.; AND SOBELL, M.B. Clinical efficacy of the 5-HT3 antagonist ondansetron in alcohol abuse and dependence. Alcoholism: Clinical & Experimental Research 18:879–885, 1994.

SHAW, G.K.; WALLER, S.; MAJUMDAR, S.K.; ALBERTS, J.L.; LATHAM, C.J.; AND DUNN, G. Tiapride in the prevention of relapse in recently detoxified alcoholics. British Journal of Psychiatry 165:515–523, 1994.

SPANAGEL, R., AND ZIEGLÄNSBERGER, W. Anticraving compounds for ethanol: New pharmacological tools to study addictive processes. Trends in Pharmacological Sciences 18:54–58, 1997.

SWANN, A.C.; JOHNSON, B.S.; CLONINGER, C.R.; AND CHEN, Y.R. Relationships of plasma tryptophan availability to course of illness and clinical features of alcoholism: A preliminary study. Psychopharmacology 143:380–384, 1999.

SWIFT, R.M. Drug therapy for alcohol dependence. New England Journal of Medicine 340(19):1482–1490, 1999.

SWIFT, R.M.; DAVIDSON, D.; WHELHAN, W.; AND KUZNETSOV, O. Ondansetron alters human alcohol intoxication. Biological Psychiatry 40:514–521, 1996.

TAMION, F.; PETIT, J.; MASSARI, P.; LEROY, J.; BIGA, N.; AND OKESENHELMER, G. Malignant Neuroleptic Syndrome during tiapride treatment. Journal de Toxicologie Clinique et Experimentale 10:461–467, 1990.

VOLPICELLI, J.R.; ALTERMAN, A.J.; HAYASIDA, M.; AND O’BRIEN, C.P. Naltrexone in the treatment of alcohol dependence. Archives of General Psychiatry 49:876–880, 1992.

VOLPICELLI, J.R.; WATSON, N.T.; KING, A.C.; SHERMAN, C.E.; AND O’BRIEN, C.P. Effect of naltrexone on alcohol “high” in alcoholics. American Journal of Psychiatry 152(4):613–615, 1995.

VOLPICELLI, J.R.; RHINES, K.C.; RHINES, J.S.; VOLPICELLI, L.A.; ALTERMAN, A.J.; AND O’BRIEN, C.P. Naltrexone and alcohol dependence. Role of subject compliance. Archives of General Psychiatry 54(8):737–742, 1997.

WHITWORTH, A.B.; FISHER, F.; LESCH, O.M.; NIMMERREICHTER, A.; OBERBAUER, H.; PLATZ, T.; POTGIETER, W.H.; AND FLEISCHHAER, W.W. Comparison of acamprosate and placebo in long-term treatment of alcohol dependence. Lancet 347:1438–1442, 1996.
GLOSSARY

Agonist: A drug or medication that mimics the effects of a neurotransmitter at a receptor.

Antagonist: A drug or medication that blocks or reverses the effects of a neurotransmitter.

Appetitive behavior: The activities involved in seeking and approaching a rewarding stimulus such as food or alcohol.

Consummatory behavior: The activities involved in contacting and obtaining a rewarding stimulus to attain its effects (i.e., eating or drinking).

Dependence: Addiction to alcohol (i.e., alcoholism) or to other drugs resulting in part from the effects of reinforcement.

Dopamine: A neurotransmitter that helps regulate motivational behavior, among other functions. Release of dopamine into the nucleus accumbens has been associated with the reinforcing effects of alcohol.

Endorphins: A class of opioid peptides, including beta-endorphin.

Glutamate: The primary excitatory neurotransmitter in the brain.

Motivational: Pertaining to mental states or behavior that increase the probability of approaching and attaining a reward (e.g., seeking and consuming alcohol). Motivational behavior includes appetitive and consummatory phases.

Negative craving: An urge or desire to resume alcohol consumption in response to neurological changes induced by heavy drinking.

Neuron: A nerve cell.

Neurotransmitter: A chemical messenger released by an excited or stimulated neuron. After release, neurotransmitters may travel across a synapse and then bind to a receptor on an adjacent neuron, usually triggering a series of chemical and electrical changes in the second cell.

Nucleus accumbens: A brain region affected by many drugs of abuse and implicated in the process of reinforcement.

Opioid peptides: Neurotransmitters with morphine-like effects, including mood elevation. Beta-endorphin and other opioid peptides may contribute to the development of alcoholism by modulating the activity of dopamine in the nucleus accumbens.

Partial agonist: An agent that is only partially effective as an agonist. Partial agonists also can act as antagonists.

Protein: The product of the genetic information encoded in a gene.

Receptor: A protein usually found on the surface of a neuron or other cell that recognizes and binds to neurotransmitters or other chemical messengers.

Reinforcement: A learning process that strengthens the ability of a rewarding stimulus to elicit motivational behavior.

Reward: The goal of motivated behavior, sometimes consciously perceived as a pleasurable state (e.g., the effects of mild alcohol intoxication) or as relief from a painful or uncomfortable state (e.g., amelioration of withdrawal symptoms).

Serotonin: A neurotransmitter that helps regulate multiple physiological functions, including appetitive and consummatory behaviors. Serotonin also may modulate the effects of dopamine in the nucleus accumbens.

Synapse: A microscopic gap separating adjacent neurons where neurotransmitters and receptors cluster.

Withdrawal: A syndrome that may occur following cessation or reduction of heavy drinking. Symptoms may include tremors, agitation, and seizures.