

Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy (Review)

Goh ET, Andersen ML, Morgan MY, Gluud LL

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[Intervention Review]

# Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

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## ABSTRACT

#### Background

Hepatic encephalopathy is a common complication of cirrhosis which results in poor brain functioning. The spectrum of changes associated with hepatic encephalopathy ranges from the clinically 'indiscernible' or minimal hepatic encephalopathy to the clinically 'obvious' or overt hepatic encephalopathy. Flumazenil is a synthetic benzodiazepine antagonist with high affinity for the central benzodiazepine recognition site. Flumazenil may benefit people with hepatic encephalopathy through an indirect negative allosteric modulatory effect on gamma-aminobutyric acid receptor function. The previous version of this review, which included 13 randomised clinical trials, found no effect of flumazenil on all-cause mortality, based on an analysis of 10 randomised clinical trials, but found a beneficial effect on hepatic encephalopathy, based on an analysis of eight randomised clinical trials.

#### Objectives

To evaluate the beneficial and harmful effects of flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy.

#### Search methods

We searched The Cochrane Hepato-Biliary Group Controlled Trials Register, CENTRAL, MEDLINE, Embase, Science Citation Index Expanded, and LILACS; meeting and conference proceedings; and bibliographies in May 2017.

#### Selection criteria

We included randomised clinical trials regardless of publication status, blinding, or language in the analyses of benefits and harms, and observational studies in the assessment of harms.

#### Data collection and analysis

Two review authors extracted data independently. We undertook meta-analyses and presented results using risk ratios (RR) with 95% confidence intervals (CI) and I<sup>2</sup> values as a marker of heterogeneity. We assessed bias control using the Cochrane Hepato-Biliary Group domains; determined the quality of the evidence using GRADE; evaluated the risk of small-study effects in regression analyses; and conducted trial sequential, subgroup, and sensitivity analyses.

Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

#### Main results

We identified 14 eligible randomised clinical trials with 867 participants, the majority of whom had an acute episode of overt hepatic encephalopathy. In addition, we identified one ongoing randomised clinical trial. We were unable to gather outcome data from 2 randomised clinical trials with 25 participants. Thus, our analyses include 842 participants from 12 randomised clinical trials comparing flumazenil versus placebo. We classified one randomised clinical trial at low risk of bias in the overall assessment and the remaining randomised clinical trials at high risk of bias. The duration of follow-up ranged from a few minutes to two weeks, but it was less than one day in the majority of the trials.

In total, 32/433 (7.4%) participants allocated to flumazenil versus 38/409 (9.3%) participants allocated to placebo died (RR 0.75, 95% CI 0.48 to 1.16; 11 randomised clinical trials; low quality evidence). The Trial Sequential Analysis and the one randomised clinical trial assessed as low risk of bias (RR 0.76, 95% CI 0.37 to 1.53) found no beneficial or harmful effects of flumazenil on all-cause mortality. The methods used to evaluate hepatic encephalopathy included several different clinical scales, electrophysiological variables, and psychometric tests. Flumazenil was associated with a beneficial effect on hepatic encephalopathy when including all randomised clinical trials (RR 0.75, 95% CI 0.71 to 0.80; 824 participants; 9 randomised clinical trials; low quality evidence), or just the trial at low risk of bias (RR 0.78, 95% CI 0.72 to 0.84; 527 participants). The Trial Sequential Analysis supported a beneficial effect of flumazenil on hepatic encephalopathy. The randomised clinical trials included little information about causes of death and little information on non-fatal serious adverse events.

#### Authors' conclusions

We found low quality evidence suggesting a short-term beneficial effect of flumazenil on hepatic encephalopathy in people with cirrhosis, but no evidence of an effect on all-cause mortality. Additional evidence from large, high quality randomised clinical trials is needed to evaluate the potential benefits and harms of flumazenil in people with cirrhosis and hepatic encephalopathy.

## PLAIN LANGUAGE SUMMARY

#### Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

#### Background

#### What is hepatic encephalopathy?

Cirrhosis is a chronic disorder of the liver. People with cirrhosis may develop hepatic encephalopathy, a condition which results in poor brain functioning. In some people, there are obvious clinical features of disturbed brain functioning (overt hepatic encephalopathy); these changes may be short-lived or persist for long periods of time. In other people, there are no obvious clinical changes but some aspects of brain function, such as attention and the ability to perform complex tasks are impaired when tested (minimal hepatic encephalopathy). The reason people develop hepatic encephalopathy is complex but changes in brain neurotransmitters, which are the chemical messengers which allow nerve cells to communicate with one another, may play a role. The neurotransmitter gamma aminobutyric acid (GABA) is responsible for slowing or inhibiting brain activity and is thought to play a particularly important role.

#### What is flumazenil?

Flumazenil is a medicine that acts on one of the GABA receptors in the brain to modify its effects on these specialised cells and so may benefit people with hepatic encephalopathy. It has to be given into a vein (intravenous) and its effects do not last for more than a few hours.

#### **Review question**

We investigated the use of flumazenil for the treatment of hepatic encephalopathy in people with cirrhosis by reviewing clinical trials in which people were randomly allocated to treatment with flumazenil or an inactive dummy/placebo or no specific intervention.

#### Search date

We searched medical databases and conducted manual searches in May 2017.

#### Study funding sources

Five of the included randomised clinical trials received support from pharmaceutical companies.

#### Study characteristics

We included 14 randomised clinical trials with 867 participants. All randomised clinical trials compared intravenous infusion of flumazenil versus an inactive placebo (dummy infusion, e.g. a salt solution). The duration of treatment ranged from 10 minutes to 72 hours. Ten randomised clinical trials included participants with overt hepatic encephalopathy; three included participants with minimal hepatic encephalopathy; and one randomised clinical trial included participants with overt or minimal hepatic encephalopathy.

## Key results

The analyses showed no effect of flumazenil on all-cause mortality (deaths of any cause) compared with placebo. People who received flumazenil were more likely to recover from their hepatic encephalopathy than people given a placebo. We found little information about serious side effects.

## Quality of the evidence

Overall, the evidence for the effect of flumazenil on hepatic encephalopathy was of low quality; only one randomised clinical trial included had a low risk of bias.

## SUMMARY OF FINDINGS FOR THE MAIN COMPARISON [Explanation]

Flumazenil versus placebo for people with cirrhosis and hepatic encephalopathy

Patient or population: people with hepatic encephalopathy Setting: hospital Intervention: flumazenil

Comparison: placebo

Outcomes	Anticipated absolute	effects* (95% Cl)	Relative effect (95% Cl)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Risk with placebo	Risk with flumazenil				
All-cause mortality follow-up: range 1 day to 2 weeks	Study population 93 per 1000	70 per 1000 (45 to 108)	RR 0.75 (0.48 to 1.16)	842 (11 RCTs)	$\oplus \oplus \bigcirc \bigcirc$ Low <sup>1,2</sup>	The only RCT with lo risk of bias found no e fect of flumazenil on a cause mortality (RR 76,95% CI 0.37 to 1.53 . The Trial Sequenti Analysis found insufficient evidence to su port or refute an intervention benefit/harm
Hepatic encephalopa- thy	Study population		RR 0.75 (0.71 to 0.80)	824 (9 RCTs)	$\oplus \oplus \bigcirc \bigcirc$ Low <sup>1,2</sup>	The only RCT with low risk of bias reports a beneficial effect of flumazenil on hepat encephalopathy (RR of 78, 95% CI 0.72 to 0.84; Barbaro 1998 . The Trial Sequent tial Analysis found that flumazenil was associated with a beneficial cial effect on hepat

Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

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	933 per 1000	700 per 1000 (662 to 746)				encephalopathy (Figure 1). The methods used to assess this out- come varied consider- ably (Table 1) and the duration of follow-up was very short in the majority of RCTs.
Serious adverse events	See comment	See comment	Not estimable	842 (11 RCTs)	⊕⊕⊖⊖ Low <sup>1,2</sup>	All-cause mortality was the only serious ad- verse event reported for both the interven- tion and control group (Table 6). The narrative text in 4 RCTs described that causes of death in- cluded liver failure, pro- gressive liver disease, and infections

\*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% Cl).

CI: confidence interval; RCT: randomised clinical trial; RR: risk ratio.

## **GRADE** Working Group grades of evidence

High quality: We are very confident that the true effect lies close to that of the estimate of the effect.

Moderate quality: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

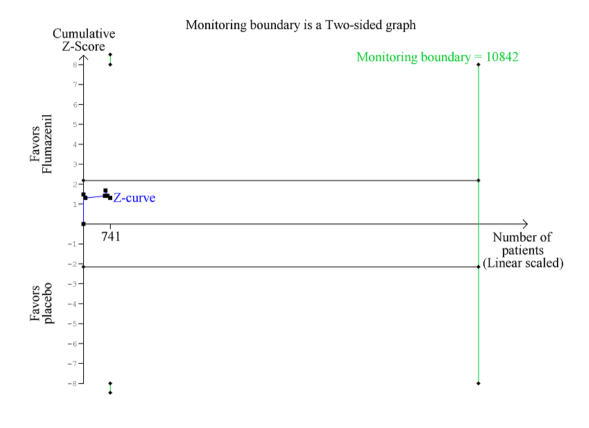
Low quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect.

Very low quality: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

<sup>1</sup> Downgraded due to risk of bias: only one RCT had a low risk of bias.

<sup>2</sup> Downgraded due to imprecision: wide confidence intervals.

Figure 1. Trial Sequential Analysis of randomised clinical trials evaluating flumazenil versus placebo for people with hepatic encephalopathy. The outcome is all-cause mortality. The original meta-analysis included 11 randomised clinical trials with 842 participants. The Trial Sequential Analysis ignored three randomised clinical trials due to insufficient information size (Cadranel 1995; Gyr 1996; Zhu 1998). The analysis was made with alpha 3%, power 90%, relative risk reduction 20%, assumed control risk 10%, and diversity 10%. The blue line (Z-curve) corresponds to the cumulative meta-analysis, the black horizontal line is the conventional boundary (3% level of significance), and the inward sloping green line is the Trial Sequential Monitoring Boundary. Futility boundaries are ignored because the information is insufficient. The analysis found no evidence to support or refute a beneficial or harmful effect of flumazenil on mortality.



## BACKGROUND

## **Description of the condition**

Hepatic encephalopathy is the term used to describe the spectrum of neuropsychiatric changes that can occur in people with cirrhosis. The joint guideline from the European Association for the Study of the Liver (EASL) and the American Association for the Study of Liver Diseases (AASLD) defines hepatic encephalopathy as brain dysfunction associated with liver insufficiency or portal systemic shunting (EASL/AASLD guideline 2014a; EASL/AASLD guideline 2014b).

Clinically apparent or*overt* hepatic encephalopathy manifests as a neuropsychiatric syndrome encompassing a wide spectrum of mental and motor disorders (Weissenborn 1998; Ferenci 2002). Events such as gastrointestinal bleeding, infection, and alcohol misuse can trigger this so-called acute or episodic hepatic encephalopathy. Fifty per cent of instances occur with no obvi-

ous precipitating cause. Episodes may recur. Between episodes, people may return to their baseline neuropsychiatric status or retain a degree of impairment (Bajaj 2010). Less frequently, people present with persistent neuropsychiatric abnormalities, which are always present to some degree, but which may vary in seriousness. The clinical features of overt hepatic encephalopathy ranges from subtle alterations in personality, intellectual capacity and cognitive function to more profound alterations in motor function and consciousness leading to deep coma (Weissenborn 1998). Other abnormalities include impaired psychometric performance (Schomerus 1998; Randolph 2009), disturbed neurophysiological function (Parsons-Smith 1957; Chu 1997), altered cerebral neurochemical/neurotransmitter homeostasis (Taylor-Robinson 1994), reductions in global and regional cerebral blood flow and metabolism (O'Carroll 1991), and changes in cerebral fluid homeostasis (Haussinger 2000). In general, the degree of impairment in these parameters increases as the clinical condition worsens. Minimal hepatic encephalopathy (in the older literature 'subclinical' or 'latent') refers to people with cirrhosis who are 'clinically normal,' but who show abnormalities in neuropsychometric or neurophysiological performance (Ferenci 2002). Hepatic encephalopathy, whether minimal or overt, is associated with impairment in the performance of complex tasks, such as driving (Schomerus 1981; Bajaj 2009; Kircheis 2009), and a detrimental effect on health-related quality of life (Groeneweg 1998), safety (Roman 2011), neurocognitive function post-transplantation (Sotil 2009), and survival (Bustamante 1999; D'Amico 2006; Stewart 2007). About 42% of people with cirrhosis are alive one year after their first episode of hepatic encephalopathy but only 23% are alive after three years (Bustamante 1999). Thus, more than 50% die within one year and more than 75% die within three years. Hepatic encephalopathy also poses a substantial burden for the carers of affected people (Bajaj 2011) and on healthcare systems (Poordad 2007; Stepanova 2012).

#### Prevalence of hepatic encephalopathy

The prevalence of hepatic encephalopathy varies. About 10% to 14% of people with cirrhosis have overt hepatic encephalopathy at the time they are first diagnosed as having liver disease (Saunders 1981). In people with decompensated cirrhosis the prevalence of overt hepatic encephalopathy is around 20% (D'Amico 1986; de Jongh 1992; Zipprich 2012). The cumulated incidence of overt hepatic encephalopathy is as high as 40% (Randolph 2009; Bajaj 2011). The prevalence of minimal hepatic encephalopathy varies but may be as high as 50% (Lauridsen 2011).

#### **Diagnosing hepatic encephalopathy**

There is no gold standard for the diagnosis of hepatic encephalopathy. A detailed neuropsychiatric history and examination is important to identify suggestive abnormalities while eliminating other potential causes of similar cerebral changes (Montagnese 2004). The West Haven or Conn criteria are commonly used to assess and grade the mental state (Conn 1977), while the Glasgow Coma Scale is used to grade the level of consciousness (Teasdale 1974). People with hepatic encephalopathy show impairment on a range of psychometric tests. People with minimal hepatic encephalopathy show deficits in attention, visuo-spatial abilities, fine motor skills, and memory while other cognitive functions are relatively well preserved. People with overt hepatic encephalopathy show additional disturbances in psychomotor speed, executive function, and concentration. The Psychometric Hepatic Encephalopathy Score, which employs five paper and pencil tests to assess attention, visual perception and visuo-constructive abilities is widely used in the assessment of psychometric change in people with cirrhosis (Schomerus 1998; Weissenborn 2001). People with hepatic encephalopathy may have a number of neurophysiological abnormalities (Guérit 2009). The electroencephalogram, which primarily reflects cortical neuronal activity, may show progressive slowing of background activity and abnormal wave morphology. The brain responses, or evoked potentials, to stimuli such as light and sounds may show abnormal slowing or abnormal wave forms. Other potential diagnostic techniques include the Critical Flicker Fusion Frequency (Kircheis 2002), and the Inhibitory Control Test (Bajaj 2008). Blood ammonia concentrations are not routinely measured to diagnose hepatic encephalopathy but they are sometimes monitored in clinical trials.

#### **Description of the intervention**

The pathogenesis of hepatic encephalopathy is complex and incompletely understood. It is associated with a general depression in cerebral function and a shift in balance between inhibitory and excitatory neurotransmission favouring inhibition. Gamma aminobutyric acid (GABA) is the major inhibitory neurotransmitter in the brain; it binds to a specific receptor; the GABA-Acomplex, which also has neurosteroid and benzodiazepine modulatory sites (Butterworth 2016). People with hepatic encephalopathy have been shown to have increased 'GABA tone'. Use of a benzodiazepine receptor antagonist may counteract the increased tone and benefit people with hepatic encephalopathy (Ahboucha 2008).

Flumazenil competitively inhibits the activity at the benzodiazepine recognition site on the GABA-A receptor complex, but lacks major intrinsic pharmacological or behavioural activity (Whitwam 1995). Flumazenil is used to treat benzodiazepine overdose and in the reversal of anaesthesia. Following intravenous administration, flumazenil distributes extensively in the extravascular space with an initial distribution half-life of 4 to 11 minutes and a terminal half-life of 40 to 80 minutes. Maximum plasma concentrations are reached at around 50 minutes. Flumazenil is completely metabolised primarily by hepatic metabolism and has a relatively high hepatic extraction ratio (Amrein 1990). In people

with moderate liver dysfunction, the mean total clearance is decreased to 40% to 60%. In people with severe liver dysfunction, clearance is decreased to 25%. This results in a prolongation of the half-life from 50 minutes in healthy volunteers to 1.3 hours in people with moderate hepatic impairment and 2.4 hours in people with severe hepatic dysfunction (Amrein 1990). Nevertheless, even in people with severe liver disease, its duration of action remains relatively short.

#### How the intervention might work

Glutamate dehydrogenase synthesises GABA from glutamate in presynaptic nerves. GABA binds to a specific receptor, which is embedded in the postsynaptic neural membrane. This receptor is part of a larger GABA-A receptor complex, which also has binding sites for benzodiazepines, barbiturates, and neurosteroids. The binding of any of these ligands opens a chloride channel; the influx of chloride ions results in hyperpolarisation of the postsynaptic membrane and neuro-inhibition. Neurosteroids are synthesised in the brain, primarily in astrocytes, and mediates an increased GABA-A tone, which is associated with hepatic encephalopathy. Neurosteroid synthesis is mediated via activation of translocator protein, a mitochondrial neuroglial cholesterol-transporter protein previously (Butterworth 2016). Translocator protein sites (previously known as 'peripheral-type' or mitochondrial) benzodiazepine receptors. Neurosteroids such as  $3\alpha$ - $5\alpha$ -tetrahydroprogesterone (allopregnanolone) are potent, endogenous, positive allosteric modulators of both the GABA and benzodiazepine sites on the GABA-A receptor complex. Autopsy and imaging studies show consistent upregulation of translocator protein sites in people with hepatic encephalopathy. This upregulation is most likely mediated by ammonia and manganese (Ahboucha 2008), both of which accumulate as a result of hepatocellular failure and portal-systemic shunting of blood. In addition, cerebrospinal fluid and autopsied brains of people with hepatic encephalopathy have increased concentrations of agonist ligands such as diazepam-binding inhibitor and octadecaneuropeptide, which modulate the function of translocator protein sites (Córdoba 2002). People who die from hepatic encephalopathy have increased cerebral concentrations of allopregnanolone, which modulates components of the GABA-A receptors. The neurosteroids may also act synergistically with other neurotoxins such as ammonia and benzodiazepine-like compounds to further modulate GABA-A receptor function. The net effect is an increase in GABA-A tone and neural inhibition. Involvement of the GABA system in the pathogenesis of hepatic encephalopathy is consistent with the increased sensitivity to benzodiazepines observed in these patients (Batki 1987).

Flumazenil is a selective, synthetic benzodiazepine antagonist with high affinity for the benzodiazepine recognition site but is itself devoid of intrinsic activity. This compound functions as an antagonist of positive and negative modulators acting at benzodiazepine recognition sites located in the GABA-A receptor (Vicini 1987). Flumazenil may exert an indirect negative allosteric modulatory effect on GABA-A receptor function by reducing the facilitatory action of the central benzodiazepine receptor on GABA-related opening of the chloride ion-channel and in turn the excessive inhibitory effect.

#### Why it is important to do this review

The resource utilisation associated with the management of people with hepatic encephalopathy continues to escalate (Poordad 2007; Stepanova 2012). The costs of treatment and rehabilitation are increasing year on year (Neff 2010). The Identification of effective interventions which will facilitate the management of people with hepatic encephalopathy is clearly important. A number of randomised clinical trials have assessed the effects of flumazenil in people with cirrhosis and hepatic encephalopathy (Klotz 1989; Hermant 1991; Cadranel 1995; Gooday 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999; Dursun 2003). The numbers of the randomised clinical trials and of the included participants is small and several studies used a cross-over design, which hampers the analyses of clinical outcomes such as mortality and morbidity. A meta-analysis undertaken in 2002 found that use of flumazenil may be associated with clinical and electroencephalographic improvement in people with cirrhosis and hepatic encephalopathy (Goulenok 2002). The previous versions of this review included 13 randomised clinical trials (Als-Nielsen 2001; Als-Nielsen 2004) and found no beneficial effect of flumazenil on all-cause mortality, but a potential beneficial effect on manifestations of hepatic encephalopathy. We have updated this review based on current recommendations (Gluud 2017).

## OBJECTIVES

To evaluate the beneficial and harmful effects of flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy.

## METHODS

## Criteria for considering studies for this review

#### Types of studies

We included randomised clinical trials regardless of their publication status, language, or blinding in our primary analyses. If, during the selection of trials, we identified observational studies (i.e. quasi-randomised studies, cohort studies, or patient reports)

that reported adverse events caused by, or associated with, the interventions in our review, we included these studies for a review of the adverse events. We did not specifically search for observational studies for inclusion in this review, which is a known limitation.

#### **Types of participants**

Randomised clinical trials evaluating participants with cirrhosis and hepatic encephalopathy, irrespective of the aetiology and severity of the underlying liver disease. Included participants could have overt or minimal hepatic encephalopathy. If we identified trials including subsets of relevant participants with cirrhosis as well as participants without cirrhosis, we planned to exclude these trials in sensitivity analyses.

#### **Types of interventions**

Flumazenil at any dose, duration, or mode of administration versus placebo or no intervention.

#### Types of outcome measures

We assessed all outcomes at the maximum duration of follow-up.

#### **Primary outcomes**

All cause-mortality.

• Hepatic encephalopathy (number of participants without improved manifestations).

• Serious adverse events defined as any untoward medical occurrence that resulted in death, was life-threatening, required hospitalisation, led to prolongation of the existing hospitalisation, or resulted in persistent or significant disability (ICH-GCP 1997). We analysed serious adverse events as a composite outcome (Gluud 2017).

#### Secondary outcomes

• Non-serious adverse events defined as any adverse event that did not fulfil the criteria for a serious adverse event.

• Health-related quality of life.

#### Exploratory outcomes

• Number Connection Test results.

## Search methods for identification of studies

The last search update was May 2017.

#### **Electronic searches**

We searched The Cochrane Hepato-Biliary Group Controlled Trials Register (Gluud 2017), Cochrane Central Register of Controlled Trials (CENTRAL; 2017, Issue 2) in the Cochrane Library (searched May 2017), MEDLINE Ovid (1946 to May 2017), Embase Ovid (1974 to May 2017), Science Citation Index Expanded (Web of Science; 1900 to May 2017) (Royle 2003), and LILACS (Bireme; 1982 to May 2017) using the strategy described in Appendix 1. We did not have access to Chinese or Japanese databases. We plan to search both Chinese and Japanese databases in future updates, should they become available to us via the Cochrane Hepato-Biliary Group.

## Searching other resources

We searched the reference lists of papers identified in the electronic searches and wrote to authors of the identified clinical trials and relevant pharmaceutical companies. We searched the conference proceedings of the European Association for the Study of the Liver (EASL), the United European Gastroenterology Week (UEGW), the American Gastroenterological Association (AGA), the American Association for the Study of Liver Diseases (AASLD), the International Society for Hepatic Encephalopathy and Nitrogen metabolism (ISHEN), the World Health Organization (WHO) online trial meta-register (apps.who.int/trialsearch/), and Google Scholar using the search terms cirrhosis AND flumazenil.

## Data collection and analysis

#### Selection of studies

Three review authors (ETG, MYM, and LLG), working independently, read the updated electronic searches, performed additional handsearches, and listed potentially eligible trials. All authors read the potentially eligible trials and participated in the final selection of those to be included in the analyses. For trials reported in more than one publication, we selected the paper reporting the longest duration of follow-up as the primary reference. We listed details of all included trials in a 'Characteristics of included studies' table and listed all excluded studies with the reason for their exclusion in a 'Characteristics of excluded studies table'.

## Data extraction and management

All review authors participated in data extraction and at least two review authors independently evaluated each randomised clinical trial. We resolved disagreements through discussion and sought key unpublished information that was missing from published trial reports, through correspondence with the primary investigators of included randomised clinical trials.

Where we were not able to gather sufficient data (number of events and participants) from the text and tables of the included reports of randomised clinical trials or from correspondence with investigators we attempted to extrapolate data , where possible from any contained graphical material.

We gathered data on the following:

• Trials: design (cross-over or parallel), setting (number of clinical sites), country of origin, inclusion period;

• Participants: mean age, proportion of men, type of hepatic encephalopathy, proportion with cirrhosis, proportion with alcoholic liver disease, proportion with viral hepatitis;

• Interventions: type, dose, duration of therapy, mode of administration;

• Outcomes: outcomes assessed, criteria used in the assessment of hepatic encephalopathy.

## Assessment of risk of bias in included studies

We assessed bias control using the domains described in the Cochrane Hepato-Biliary Group Module (Gluud 2017) and classified the risk of bias for separate domains as high, unclear, or low. We also included an overall assessment of bias control as described below:

#### Allocation sequence generation

• Low risk of bias: sequence generation achieved using computer random number generation or a random number table. Drawing lots, tossing a coin, shuffling cards, or throwing dice were adequate if performed by an independent person not otherwise involved in the trial.

• Unclear risk of bias: not described.

• High risk of bias: sequence generation method was not random.

#### Allocation concealment

• Low risk of bias: used a central and independent randomisation unit or similar adequate method (e.g. serially numbered opaque sealed envelopes) to ensure that the allocation sequence was unknown to the investigators (Hrobjartsson 2001; Savovic 2012a; Savovic 2012b).

Unclear risk of bias: allocation not described.

• High risk of bias: allocation sequence was likely to be known to the investigators who assigned the participants.

#### Blinding of participants and personnel

• Low risk of bias: blinding of participants and personnel performed adequately using a placebo. We defined lack of blinding as not likely to affect the evaluation of mortality.

• Unclear risk of bias: insufficient information to assess blinding.

• High risk of bias: no blinding or incomplete blinding.

#### Blinding of outcome assessors

• Low risk of bias: blinding of outcome assessors performed adequately using a placebo. We defined lack of blinding as not likely to affect the evaluation of mortality (Hrobjartsson 2001; Savovic 2012a; Savovic 2012b).

• Unclear risk of bias: there was insufficient information to blinding.

• High risk of bias: no blinding or incomplete blinding.

#### Incomplete outcome data

• Low risk of bias: missing data were unlikely to make treatment effects depart from plausible values. The investigators used sufficient methods, such as intention-to-treat analyses with multiple imputations or carry-forward analyses to handle missing data.

• Unclear risk of bias: there was insufficient information to assess missing data.

• High risk of bias: the results were likely to be biased due to missing data.

#### Selective outcome reporting

• Low risk of bias: the trial reported clinically relevant outcomes (all-cause mortality, hepatic encephalopathy, and serious adverse events). If we had access to the original trial protocol, the outcomes selected should have been those described in the protocol. If we obtained information from a trial registry (such as www.clinicaltrials.gov), we only used that information if the investigators registered the trial before inclusion of the first participant.

• Unclear risk of bias: predefined relevant outcomes were not reported fully or the reporting was unclear.

 High risk of bias: one or more predefined outcomes were not reported.

#### For-profit bias

• Low risk of bias: the trial appeared free of industry sponsorship or other type of for-profit support.

 Unclear risk of bias: no information on clinical trial support or sponsorship.

• High risk of bias: the trial was sponsored by industry or received other support (such as provision of study drugs).

#### Other bias

• Low risk of bias: the trial appeared free of other biases including: medicinal dosing problems or follow-up (as defined below).

• Unclear risk of bias: the trial may or may not have been free of other domains that could put it at risk of bias.

• High risk of bias: there were other factors in the trial that could put it at risk of bias such as the administration of inappropriate treatments being given to the controls (e.g. an inappropriate dose) or follow-up (e.g. the trial included different follow-up schedules for participants in the allocation groups).

#### **Overall bias assessment**

• Low risk of bias: all domains were low risk of bias using the definitions described above.

• High risk of bias: one or more of the bias domains were of unclear or high risk of bias.

#### **Measures of treatment effect**

We analysed dichotomous data using risk ratios (RR) and continuous data using mean differences (MD), both with 95% confidence intervals (CI).

#### Unit of analysis issues

Due to the fluctuating nature of hepatic encephalopathy and the nature of our primary outcomes, we included randomised clinical trials using a parallel-arm design and the first treatment period from cross-over trials.

#### Dealing with missing data

We extracted data on all participants randomised to allow intention-to-treat analyses. We planned to undertake analyses to evaluate the influence of missing data (Higgins 2008) including worstcase scenario analysis, and extreme worst-case and best-case scenario analyses (Gluud 2017). However, we did not identify randomised clinical trials with missing outcome data.

#### Assessment of heterogeneity

We assessed heterogeneity through visual inspection of the forest plots and expressed heterogeneity as  $I^2$  values using the following thresholds: 0% to 40% (unimportant), 40% to 60% (moderate), 60% to 80% (substantial), and greater than 80% (considerable). We included the information in our 'Summary of findings' tables.

#### Assessment of reporting biases

For meta-analyses with at least 10 randomised clinical trials (metaanalysis evaluating all-cause mortality), we planned to prepare funnel plots and regression analyses of funnel plot asymmetry (Harbord 2006). However, our analyses included fewer than 10 randomised clinical trials.

#### Data synthesis

We performed the analyses using Review Manager 5 (RevMan 2014), STATA (Stata 14), and Trial Sequential Analysis (TSA 2011).

#### **Meta-analysis**

We performed random-effects and fixed-effect meta-analyses. The estimates of the random-effects and fixed-effect meta-analyses were similar for all analyses. Therefore, we assumed that any smallstudy effects had little effect on the intervention effect estimate. For random-effects models, precision decreased with increasing heterogeneity and CIs widened correspondingly. Accordingly, the random-effects model provided the most conservative (and a more correct) estimate of the intervention effect. Accordingly, we report the results of our analyses based on random-effects meta-analyses.

#### **Trial Sequential Analysis**

We performed Trial Sequential Analysis to evaluate the risk of errors and to evaluate futility (Wetterslev 2008; Thorlund 2011; Wetterslev 2017). We defined the required information size as the number of participants needed to detect or reject an intervention effect. We set alpha to 3% and power to 90% in all analyses. We used the model-based diversity and repeated all analyses with a diversity increased by 10%. Due to the lack of randomised clinical trials assessed at low risk of bias, we were only able to conduct the analyses with inclusion of all randomised clinical trials (regardless of bias control).

**All-cause mortality:** based on one large randomised clinical trial evaluating flumazenil (Barbaro 1998), and cohort studies evaluating the prognosis of people with hepatic encephalopathy (EASL/AASLD guideline 2014a; EASL/AASLD guideline 2014b), as well as the results of our meta-analysis, we set the relative risk reduction to 10%, the control group risk to 20%, and increased diversity to 10%.

**Hepatic encephalopathy:** based on the Cochrane Hepato-Biliary Group recommendations, we conducted the analysis with a relative risk reduction of 20% (corresponding to the upper 95% CI) and reduced the control group risk from the observed 93% to 60% (set lower than the observed control group risk).

Serious adverse events: we were only able to identify serious adverse events that were fatal. Accordingly, our analysis of all-cause mortality and serious adverse events included the same numbers.

#### Subgroup analysis and investigation of heterogeneity

We performed subgroup analyses to investigate heterogeneity in randomised clinical trials based on the:

- Type of hepatic encephalopathy;
- Inclusion of participants with cirrhosis or acute liver failure
- The trial design;

• Duration of follow-up.

Only one randomised clinical trial had a low risk of bias in the overall assessment.

#### Sensitivity analysis

We planned to perform worst-case scenario analyses, but none of the trials reported missing outcome data.

## 'Summary of findings' table

We used GRADEpro (GRADEpro 2008) to generate a 'Summary of findings' table with information about outcomes, risk of bias, and results of the meta-analyses. We used the GRADE system to evaluate the quality of the evidence for outcomes reported in the review considering the within-study risk of bias (methodological quality), indirectness of evidence, diversity (heterogeneity), imprecision of effect estimate, and risk of publication bias.

## RESULTS

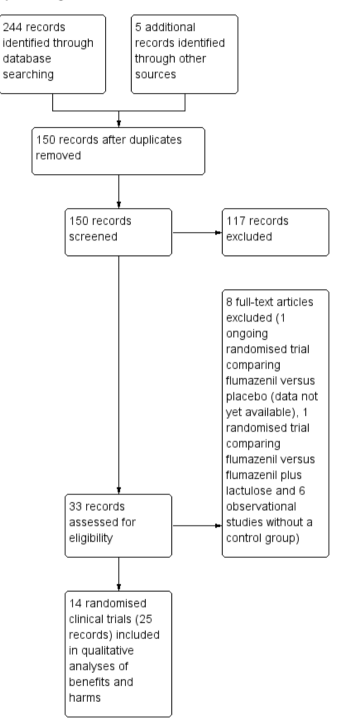
#### **Description of studies**

We included 14 randomised clinical trials in our analyses of benefits and harms (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009). We identified one randomised clinical trial which is currently ongoing (Yale 2014). We are unable to determine if this ongoing randomised clinical trial will be eligible for inclusion in the review. We excluded one randomised clinical trial and 2 non-randomised studies from the primary analyses, but included them in the evaluation of harms (Marsepoil 1990; Kapczinski 1995<u>;</u> Jia 1999). For additional information, see Characteristics of included studies; Characteristics of excluded studies; and Characteristics of ongoing studies tables.

#### **Results of the search**

We identified 249 references in the electronic and manual searches (Figure 2). After exclusion of duplicates and references to papers that did not describe clinical trials assessing benzodiazepine receptor antagonists for hepatic encephalopathy, we retrieved 33 references for further assessment. Fourteen randomised clinical trials described in 25 references fulfilled our inclusion criteria (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009). In addition, we identified one ongoing randomised clinical trial with an estimated completion date of June 2017 (Yale 2014).

Figure 2. Study flow diagram for identification and selection of randomised clinical trials.



Two trials were letters (Klotz 1989; Hermant 1991), and 12 were full-text articles (Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009). The language of the publications was English (Klotz 1989; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003), Chinese (Zhu 1998; Li 2009), or French (Hermant 1991).

## **Included studies**

Six trials used a parallel group (Hermant 1991; Gyr 1996; Zhu 1998; Lacetti 2000; Dursun 2003; Li 2009), and eight used a crossover design (Klotz 1989; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999). In 3 cross-over trials, only people classified as non-responders participated in the second cross-over treatment period (Pomier-Layrargues 1994; Cadranel 1995; Barbaro 1998). Six of the cross-over trials included a washout period, which ranged from 3 minutes to 1 week (Pomier-Layrargues 1994; Gooday 1995; Van der Rijt 1995; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999).

The investigators diagnosed hepatic encephalopathy using clinical scores, electrophysiological techniques, and psychometric tests (Table 1; Table 2; Table 3). Nine randomised clinical trials included participants with overt hepatic encephalopathy (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Van der Rijt 1995; Barbaro 1998; Zhu 1998; Lacetti 2000; Li 2009). Six of these randomised clinical trials described precipitating events; the most common was gastrointestinal bleeding (Table 4). Three randomised clinical trials included participants with minimal hepatic encephalopathy (Gooday 1995; Amodio 1997; Giger-Mateeva 1999), while one included participants with either minimal or overt hepatic encephalopathy (Dursun 2003). The majority of the participants in the 14 included studies had cirrhosis although 30 (33%) of the 90 participants included in two of the trials had fulminant hepatic failure (Van der Rijt 1995; Li 2009).

Recent ingestion of benzodiazepines was a stipulated exclusion criterion in 10 randomised clinical trials with exclusion periods ranging from 3 days to 3 months (Hermant 1991; Pomier-Layrargues 1994; Gooday 1995; Van der Rijt 1995; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Table 5). Nine randomised clinical trials tested for blood/urine benzodiazepines at baseline (Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999; Lacetti 2000); three randomised clinical trials stipulated negative screening for blood/urine benzodiazepines as an inclusion criterion (Hermant 1991; Van der Rijt 1995; Lacetti 2000), while the proportion who tested positive for benzodiazepines in the other 6 randomised clinical trials ranged from 1.9% to 21%. The remaining five randomised clinical trials either stated that they did not measure blood/urine benzodiazepines or else did not mention testing (Table 5).

All randomised clinical trials compared intravenous flumazenil versus placebo. The daily dose of flumazenil ranged from 0.2 mg to 6.5 mg and the total dose from 0.2 mg to 19.5 mg. The duration of treatment ranged from 10 minutes to 72 hours. Five randomised clinical trials evaluated the intervention effects at the end of the intervention (Klotz 1989; Hermant 1991; Amodio 1997; Giger-Mateeva 1999; Dursun 2003). Four randomised clinical trials evaluated clinical uticals at maximum of 24 to 72 hours after the intervention (Pomier-Layrargues 1994; Cadranel 1995; Van der Rijt 1995; Lacetti 2000). The remaining five randomised clinical trials followed participants from 4 days to 4 weeks (Gooday 1995; Gyr 1996; Barbaro 1998; Zhu 1998; Li 2009).

The trials involving participants with overt hepatic encephalopathy defined overall improvement of hepatic encephalopathy based on a clinical assessment of mental status and the electroencephalogram (Table 1). The trials involving participants with minimal hepatic encephalopathy based their assessment of overall improvement on a subjective assessment of 'alertness' (Giger-Mateeva 1999); Number Connection Test results (Dursun 2003), the Simple Reaction Time test results (Gooday 1995), or electroencephalography (Van der Rijt 1995).

#### **Excluded studies**

We excluded one cross-over randomised clinical trial evaluating cognitive function and anxiety in people with alcohol-related or non-alcohol-related cirrhosis (Kapczinski 1995), one randomised clinical trial evaluating flumazenil alone or with lactulose (Wu 2001), and seven observational studies (Grimm 1988; Bansky 1989; Marsepoil 1990; Devictor 1995; Ozyilkan 1997; Golubovic 1999; Jia 1999). None of the excluded studies reported data that allowed analysis of serious adverse events.

One double-blind, cross-over trial involved participants with cirrhosis who were liver transplant candidates (Kapczinski 1995). The objective of the trial was to evaluate the differential effects of flumazenil versus placebo on cognitive function and anxiety in 10 people with alcohol-related cirrhosis, 10 people with non-alcoholrelated cirrhosis, and 10 healthy volunteers. None of the included participants had evidence of overt hepatic encephalopathy. The investigators evaluated a range of psychometric tests and reported the results as group mean values. The trial report did not provide information about the number of participants with abnormal test results, but stated that participants with cirrhosis performed worse

than people in the control group on several tests including verbal recall, and on reaction time tasks. Treatment with flumazenil had no effect on the test results, but induced anxiety in the participants with non-alcoholic cirrhosis.

Two prospective non-randomised observational studies involved 47 participants with cirrhosis and overt hepatic encephalopathy (Marsepoil 1990; Jia 1999). The first study included 25 participants with alcohol-related cirrhosis and acute hepatic encephalopathy, 13 of whom received flumazenil in a dose of 0.2 mg intravenously every 10 minutes until clinical improvement up to a maximum total dose of 2 mg followed by a continuous maintenance infusion of 0.3 mg. per hour for 48 hours.(Marsepoil 1990). The second study included 22 participants with cirrhosis and overt hepatic encephalopathy (Jia 1999), 12 of whom received flumazenil while the remaining 10 received a traditional Chinese medicine, both infused intravenously. The trial report stated that two participants in the flumazenil group died of liver failure. The report did not mention deaths in the control group.

A randomised clinical trial involving 20 participants with cirrhosis and overt hepatic encephalopathy, 12 of whom were treated with repeated bolus injections of flumazenil combined with lactulose administered as a retention enema while a further eight participants received flumazenil alone (Wu 2001). The study authors defined improvement of hepatic encephalopathy as a three-point or greater reduction in the Conn Score within six hours after the administration of the interventions. Seven (58%) of the 12 participants in the flumazenil plus lactulose group and 4 (50%) of the eight participants in the flumazenil alone group showed improvement. There were no reported deaths or adverse events. We excluded five additional observational studies, involving 59 participants, 12 (20%) of whom were children (Grimm 1988; Bansky 1989; Devictor 1995; Ozyilkan 1997; Golubovic 1999). Two studies included participants with fulminant hepatic failure (Grimm 1988; Devictor 1995), One study, involving 14 participants with cirrhosis and overt hepatic encephalopathy reported an improvement in mental status in 71% of participants within minutes of receiving flumazenil which lasted for one to two hours; six participants died (Bansky 1989). Another study, including 10 participants with cirrhosis and severe hepatic encephalopathy reported transient improvements in the manifestations of hepatic encephalopathy in 80% of the included participants; six died within one year (Golubovic 1999). A third study evaluated the effects of incremental intravenous boluses of flumazenil in 11 participants with cirrhosis in whom baseline somatosensory evoked potentials were abnormal; four (36%) showed a improvement in evoked potentials with flumazenil (Ozyilkan 1997). One prospective study, involving 15 adults and two children, reported transient improvement in hepatic encephalopathy, after administration of flumazenil, in four (44%) of nine participants with fulminant hepatic failure and five (63%) of eight with cirrhosis (Grimm 1988). In a study undertaken exclusively in children with fulminant liver failure, awaiting emergency liver transplantation, flumazenil had a transient beneficial effect on arousal in one child (Devictor 1995).

## **Risk of bias in included studies**

We assessed the risk of bias based on published information and on additional information from the trial investigators (Figure 3).

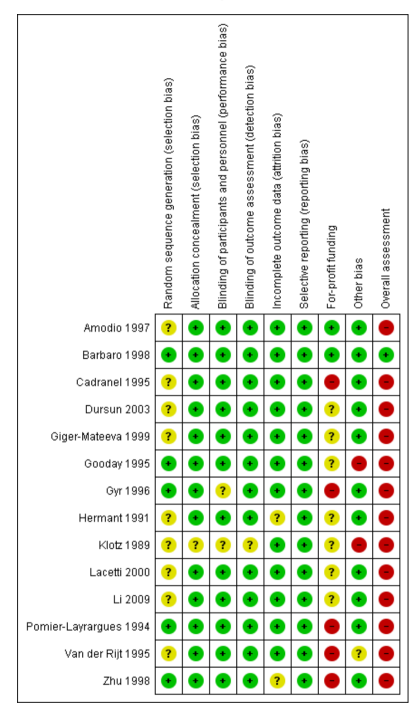


Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

## Allocation

We classified 4 randomised clinical trials with adequate allocation sequence generation and allocation concealment at low risk of selection bias (Gooday 1995; Gyr 1996; Barbaro 1998; Zhu 1998). The remaining 10 trials used an adequate method to conceal the allocation, but they did not describe the allocation sequence generation (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Van der Rijt 1995; Amodio 1997; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009).

#### Blinding

We classified all randomised clinical trials as having a low risk of performance and detection bias as they were doubleblind and placebo-controlled with blinding of participants, personnel, and outcome assessors (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009).

## Incomplete outcome data

Two randomised clinical trials gave the impression that there were no missing outcome data although this was not specifically stated (Hermant 1991; Zhu 1998). The remaining trials had no missing outcome data and included all participants in the reported analyses (Klotz 1989; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009).

#### Selective reporting

We did not have access to protocols and were, therefore, unable to evaluate any potential differences between outcomes described in protocols compared with trial publications. All randomised clinical trials included a description of the outcomes all-cause mortality and hepatic encephalopathy (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009). Therefore, we classified all trials at low risk of reporting bias.

## For-profit funding

Pharmaceutical companies provided financial or other support for 5 of the randomised clinical trials (Pomier-Layrargues 1994; Van der Rijt 1995; Cadranel 1995; Gyr 1996; Zhu 1998). Seven trials did not provide information about funding (Klotz 1989; Hermant 1991; Gooday 1995; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009). The remaining 2 trials did not receive support from pharmaceutical companies (Amodio 1997; Barbaro 1998).

#### Other potential sources of bias

One randomised clinical trial simplified the intervention regimen and assessment of outcomes after the inclusion of 9 of 18 participants (Van der Rijt 1995). We classified this trial at unclear risk of other bias and the remaining trials at low risk of bias (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003).

#### **Overall risk of bias**

We classified one randomised clinical trial at low risk of bias for all domains (Barbaro 1998), and the remaining trials at high risk of bias (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003; Li 2009).

#### **Effects of interventions**

See: Summary of findings for the main comparison Flumazenil versus placebo for people with cirrhosis and hepatic encephalopathy

The total number of participants was 867. Three cross-over randomised clinical trials did not report outcomes for the first intervention period (Gooday 1995; Amodio 1997; Giger-Mateeva 1999);we received information about the number of participants and all-cause mortality rates during the first allocation period for one of these trials (Gooday 1995), but not for the remaining two (Amodio 1997; Giger-Mateeva 1999). We were able to gather data from the first allocation period for the 5 remaining cross-over trials and required data from all of the parallel-arm trials. Accordingly, our analyses included 842 participants.

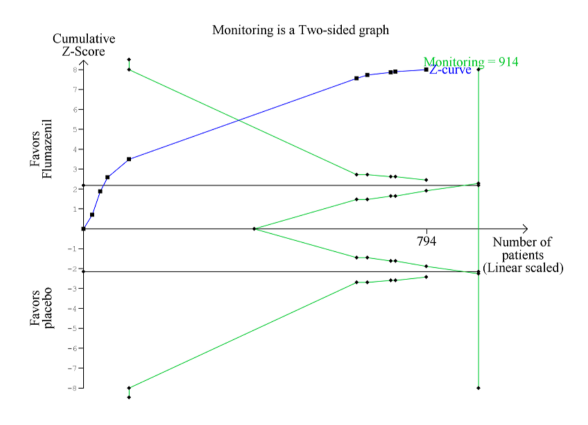
#### **Primary outcomes**

#### All-cause mortality

In total, 32/433 participants allocated to flumazenil versus 38/409 participants allocated to placebo died (RR 0.75, 95% CI 0.48 to 1.16; 11 trials;  $I^2 = 0\%$ ; Analysis 1.1). The trial classified as low risk of bias found no beneficial or detrimental effect of flumazenil on all-cause mortality (RR 0.76, 95% CI 0.37 to 1.53; Analysis 1.2). There was no evidence of small-study effects (P = 0.31). The Trial Sequential Analyses ignored three randomised clinical trials

due to insufficient information indicating that we have insufficient evidence to support or refute an effect of flumazenil on all-cause mortality (Cadranel 1995; Gyr 1996; Zhu 1998; Figure 4). The trials including participants with minimal hepatic encephalopathy did not report any deaths (Gooday 1995; Dursun 2003; Analysis 1.1). There were no differences between trials involving participants with cirrhosis compared with trials involving participants with cirrhosis or fulminant hepatic failure (Analysis 1.11). Additional analyses showed no differences between trials using a crossover or a parallel-arm design (Analysis 1.3); or between trials with short-term or long-term (> 1 day) follow-up (Analysis 1.4).

Figure 4. Trial Sequential Analysis of randomised clinical trials evaluating flumazenil versus placebo for people with cirrhosis and hepatic encephalopathy. The outcome is hepatic encephalopathy. The original metaanalysis included 11 randomised clinical trials with 824 participants. The Trial Sequential Analysis is made with alpha 3%, power 90%, relative risk reduction 20%, assumed control risk 60%, and diversity 10%. The blue line (Z-curve) corresponds to the cumulative meta-analysis, the black horizontal line is the conventional boundary (3% level of significance), and the inward sloping green line is the Trial Sequential Monitoring Boundary. The analysis found that the Z-curve crossed the monitoring boundary before reaching the diversity-adjusted required information size of 914 participants.



#### Hepatic encephalopathy

Analysis of the data on 824 participants involved in nine randomised clinical trials showed that flumazenil was associated with a beneficial effect on hepatic encephalopathy (RR 0.75, 95% CI 0.71 to 0.80;  $I^2 = 0\%$ ; Analysis 1.5). The analysis only included 10 participants with minimal hepatic encephalopathy. The trial classified as having a low risk of bias found a beneficial effect of flumazenil on hepatic encephalopathy (RR 0.78, 95% CI 0.72 to 0.84; Analysis 1.6). In the Trial Sequential Analysis, the Z-curve crossed the monitoring boundary (see Figure 4 for additional information). The analysis found that the diversity adjusted information size was 914 participants. The information size was 1028 participants when we increase the diversity to 20% (the model based diversity was 0%). The subgroup analyses showed no differences between subgroups of trials including participants with cirrhosis or fulminant hepatic failure (Analysis 1.9), or trials stratified by their design (Analysis 1.7), or their duration of follow-up (Analysis 1.8).

#### Serious adverse events

We were only able to conduct analyses for fatal serious adverse event (Table 6). In the largest randomised clinical trial (Barbaro 1998), 13 nonresponders in the flumazenil group and 17 nonresponders in the placebo group died within 3 to 4 days (range 2 to 6). The causes of death were septic shock (n = 20); hypovolaemic shock (n = 8) and lactic acidosis (n = 2), but information was not provided on the number of deaths by cause in each group. In a smaller randomised clinical trial, 4 of 28 participants allocated to flumazenil and 5 of 21 participants allocated to placebo died within 4 weeks of the trial (Gyr 1996). One participant in the placebo group died with respiratory failure during the course of the trial, but the causes of death in the remaining eight participants were not provided. Three trials reported all-cause mortality in the flumazenil and control groups without providing information about the cause of death (Zhu 1998; Lacetti 2000; Li 2009). None of the included participants experienced seizures.

#### Secondary outcome measures

None of the included trials assessed health-related quality of life. We were unable to conduct a meta-analysis of non-serious adverse events. Four randomised clinical trials reported that none of the included participants experienced non-serious adverse events (Hermant 1991; Barbaro 1998; Lacetti 2000; Dursun 2003). In one trial, four participants in the flumazenil group experienced nausea, vomiting, flushing, or irritability (Gyr 1996); the total number of participants with the individual adverse events was not described. Two participants in one trial had transient palpitations, but the intervention group was not specified (Zhu 1998). One

cross-over trials reported that one in 10 participants felt drowsy, possibly after flumazenil infusion (Giger-Mateeva 1999).

#### **Exploratory outcomes**

We were able to include Number Connection Test results from one randomised clinical trial with 40 participants. The trial found a MD of -3.79 seconds (95% CI -32.14 to 24.56; Analysis 1.10).

#### 'Summary of findings' table

As shown in the Summary of findings for the main comparison, we downgraded the strength of the evidence to low based on methodological concerns.

## DISCUSSION

#### Summary of main results

This review included 14 randomised clinical trials published between 1989 and 2009. The primary meta-analyses showed no beneficial or detrimental effect of flumazenil on all-cause mortality, but it showed a potential short-term beneficial effect of flumazenil on the manifestations of hepatic encephalopathy. We found little evidence suggesting that flumazenil was associated with serious adverse effects, However, the reporting of serious and non-serious adverse events was generally incomplete or unclear, and our analyses of adverse events may be subject to outcome reporting bias. Based on methodological concerns, we classified the strength of the evidence as low. Therefore, our review remains inconclusive.

## Overall completeness and applicability of evidence

Flumazenil is a short-acting specific benzodiazepine antagonist which acts by inhibiting activity at the benzodiazepine recognition site on the GABA/benzodiazepine receptor complex. Its oral bioavailability is poor and hence intravenous administration is necessary (Brogden 1991). The main indication for flumazenil is the reversal of benzodiazepine overdose or prolonged anaesthesia. Based on pharmacological studies, the onset of its effect is rapid and its duration of action is short (Brogden 1991). In healthy people, the half-life is 50 minutes. Thus, repeated low intravenous doses or continuous infusion is needed if the clinical situation requires a longer lasting effect (Hood 2014). Hepatic encephalopathy is characterised by an increase in GABA-A tone which is the rationale for use of flumazenil in this condition.

The most important outcomes for people with cirrhosis and hepatic encephalopathy include mortality, morbidity, adverse events, and health-related quality of life (Bajaj 2011). We found no detrimental or beneficial effect on all-cause mortality or adverse events

but a potential short-term effect on the manifestations of hepatic encephalopathy. There was no reported information on healthrelated quality of life. However, the applicability of the evidence will be limited because of the need for intravenous administration. Most trials evaluated single or repeated bolus injections (Klotz 1989; Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000). One trial assessed the effect of a 72-hour infusion of flumazenil (Van der Rijt 1995), but the investigators changed the intervention to bolus injections because participants found the three-day infusion too stressful. None of the trials compared different doses or different modes of administration. We found no effect of the dose of flumazenil on the estimated effect on all-cause mortality or hepatic encephalopathy. However, based on the limited number of events and trials, important clinical differences may have been overlooked. The half-life of flumazenil is prolonged up to 2.4 hours in people with moderate to severe hepatic decompensation, hence providing some prolongation of action (Amrein 1990). However, we found no further or additional effects of flumazenil when analysing trials with more than 24-hour follow-up.

The majority of the included randomised clinical trials enrolled participants with cirrhosis and an acute episode of hepatic encephalopathy although the severity varied between trials. Episodes of hepatic encephalopathy often develop in response to a precipitating event such as gastrointestinal bleeding, which was the most common precipitant identified in the included trials in this review. Identification and treatment of precipitating factors is key to the management of affected people (EASL/AASLD guideline 2014a; EASL/AASLD guideline 2014b). We did not have sufficient data to assess potential difference between precipitated and non-precipitated hepatic encephalopathy. None of the included participants had surgically created or transjugular intrahepatic portosystemic shunts. Likewise, the review contained very little information about people with recurrent or persistent (chronic) hepatic encephalopathy. Two trials included a small number of participants with fulminant hepatic failure (Van der Rijt 1995; Li 2009). This condition is infrequent in clinical practice. We were unable to gather data that allowed us to evaluate any differential effects of flumazenil on hepatic encephalopathy associated with acute or chronic liver failure. However, subgroup analyses based on aggregated data found no difference between trials including or not including participants with fulminant hepatic failure. Nevertheless, it is likely that the pathophysiology of hepatic encephalopathy in participants with acute liver failure differs from that associated with cirrhosis. Three trials enrolled people with minimal hepatic encephalopathy (Gooday 1995; Amodio 1997; Giger-Mateeva 1999), while a further trial included participants with both minimal and low-grade acute hepatic encephalopathy (Dursun 2003). In these four trials, the objective appears to be more mechanistic than therapeutic, but based on our subgroup analysis, we found no difference in outcomes between overt and minimal hepatic encephalopathy. However, the diagnostic end points for hepatic encephalopathy in these trials were very different, namely clinical assessment versus psychometry, and the number of trials was small so that statistical differences may be overlooked.

Prior intake of benzodiazepines may influence the effects of flumazenil. Ten randomised clinical trial stipulated prior ingestion of benzodiazepines as an exclusion criterion (Hermant 1991; Pomier-Lavrargues 1994; Gooday 1995; Van der Rijt 1995; Amodio 1997; Barbaro 1998; Zhu 1998; Giger-Mateeva 1999; Lacetti 2000; Dursun 2003), while nine trials undertook baseline screening for benzodiazepines (Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Van der Rijt 1995; Gyr 1996; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999; Lacetti 2000). Three randomised clinical trials stipulated negative testing for benzodiazepines as an inclusion criterion (Hermant 1991; Van der Rijt 1995; Lacetti 2000). In the remaining six trials, in which baseline testing was undertaken, it was unclear whether the results were available at the time of randomisation or whether participants were included irrespective of the results; in two of these trials, none of the participants tested positive (Amodio 1997; Giger-Mateeva 1999), while in the remaining four trials, between 1.9% and 21.4% of participants tested positive (Pomier-Layrargues 1994; Cadranel 1995; Gyr 1996; Barbaro 1998). Thus, the exclusion of participants based on reports of non-ingestion of benzodiazepines is clearly unreliable. In all four of these trials, the assessment of outcomes in relation to the presence/absence of benzodiazepines were related to the response to flumazenil rather than to intervention allocation (Pomier-Layrargues 1994; Cadranel 1995; Gyr 1996; Barbaro 1998). All four showed that the majority of participants who responded to flumazenil did not have detectable circulating benzodiazepines whereas some non-responders had measurable quantities of these substances in their blood. The trials concluded that the presence of benzodiazepines was not predictive of participants' responses to flumazenil. We did not find this information in 5 trials, which should be considered when evaluating their results (Klotz 1989; Gooday 1995; Zhu 1998; Dursun 2003; Li 2009).

Administration of flumazenil by sublingual lozenge and topical cream has also been tested, albeit very selectively (Rye 2012); the preparations are not generally available nor applicable for use in most clinical settings and is not tested in clinical trials evaluating participants with hepatic encephalopathy.

On the basis of this review, the use of flumazenil in the management of hepatic encephalopathy would be limited. It might be of value in people with severe hepatic encephalopathy not responding to usual management to facilitate procedures or to allow assessment of cognitive status in potential transplant candidates if other conditions, such as hypoxic injury, are suspected.

## Quality of the evidence

The review is limited because of methodological and statistical is-

sues relating to the included randomised clinical trials. Seven of the 14 included trial used a cross-over design (Pomier-Layrargues 1994; Cadranel 1995; Gooday 1995; Van der Rijt 1995; Amodio 1997; Barbaro 1998; Giger-Mateeva 1999). While this design is suitable for evaluating interventions which are predicted to have a temporary effect on chronic stable conditions, it is not suitable for evaluating interventions which have a short-term effect in unstable conditions (Rosenkranz 2015). The majority of trials in this review included participants with an acute episode of hepatic encephalopathy, and thus, a cross-over design is not appropriate. The problem was further compounded since some of the trials used a modified design with cross-over of only the non-responders to the second treatment period. Therefore, inclusion of both periods in our analyses would introduce potential bias and would have posed statistical problems. We included data from the first period of the seven cross-over trials, obtaining the data which correspond to those available from the more suitable parallel-armed randomised clinical trials. The drawback of the strategy is loss of information from the second period. A further problem with the cross-over design is that it precludes an assessment of risk of relapse.

Only one trial was at low risk of bias in the overall assessment (Barbaro 1998). The trial included the largest number of participants and had a weight of 39% in the analysis of mortality and 56% in the analysis of hepatic encephalopathy. We found no difference between this trial and the remaining trials with a high risk of bias for the two outcomes based on the test for subgroup differences; exclusion of the trial did not change the overall conclusion. We found no evidence of publication bias or other small-study effects and between-trial heterogeneity was negligible. Nevertheless, the CIs were wide. Although these findings support the quality of the evidence, we still have potential problems with the use of the cross-over design. We classified the quality of the evidence as low. While most trials reported mortality and hepatic encephalopathy, the quality of the reporting of both non-fatal serious and non-serious adverse events was low. We looked for additional information about harms in observational studies, but were unable to retrieve and analyse adverse events in the studies identified.

#### Potential biases in the review process

We attempted to minimise possible selection biases by using a comprehensive search strategy. Searches in electronic databases were combined with extensive handsearches. In addition, we also searched conference proceedings and abstract books. We think it likely that we have not missed published trials, but we cannot exclude the possibility that we have missed unpublished trials. The intervention is not high-profile and it is possible that negative trials, particularly if small, will not have appeared as abstracts at conferences or been published in full. However, our meta-regression analyses showed no evidence of publication bias or other dissemination biases.

## Agreements and disagreements with other studies or reviews

One meta-analysis undertaken in 2002 (Goulenok 2002) included 6 randomised clinical trials with 641 participants (Pomier-Layrargues 1994; Cadranel 1995; Van der Rijt 1995; Groeneweg 1996; Gyr 1996; Barbaro 1998). The mean percentages of people with clinical improvement (5 trials) were 27% in treated groups and 3% in placebo groups. We believe that two of these involve the same population of participants (Groeneweg 1996; Gyr 1996). The first trial included 49 participants with mild to moderate (Grade I to III) hepatic encephalopathy (Gyr 1996), while the second trial, which is described by the authors as an ancillary study, reported electroencephalography data from 32 of the original 49 participants (Groeneweg 1996). We excluded the second trial from this review.

The previous version of this review included 13 randomised clinical trials with 805 participants (Als-Nielsen 2004). This earlier review found no effect of flumazenil on all-cause mortality based on an analysis of 10 trials and a beneficial effect on hepatic encephalopathy based on an analysis of 8 trials with a risk difference of 0.28 (95% CI 0.20 to 0.37). We have updated the review by inclusion of two additional randomised clinical trials in the analyses of benefits, and the addition of one previously included randomised clinical trial and two observational studies in our assessment of harms. Our analysis of all-cause mortality included 13 trials and the analysis of hepatic encephalopathy included 9 trials. In agreement with the previous review, we found no effect on allcause mortality and a beneficial effect on hepatic encephalopathy.

## AUTHORS' CONCLUSIONS

## Implications for practice

This review includes randomised clinical trials evaluating the treatment of hepatic encephalopathy. The analyses found some evidence that flumazenil may be associated with a short-term effect on hepatic encephalopathy, but no beneficial effect on important clinical outcomes such as all-cause mortality, serious adverse events, or health-related quality of life. Likewise, we are unable to determine the risk of non-fatal serious or non-serious adverse events based on the available evidence.

#### Implications for research

We used the EPICOT format to define the implications of our review for research (Brown 2006). Overall, the evidence is insufficient and additional evidence from randomised clinical trials is needed to evaluate whether flumazenil has a clinically relevant effect on hepatic encephalopathy.

Evidence (what is the current state of the evidence?): this review includes 14 randomised clinical trials and found low quality evi-

dence that flumazenil may have a beneficial short-term effect on hepatic encephalopathy. The evidence concerning all-cause mortality, non-fatal serious adverse events, and non-serious adverse events is insufficient.

**P**articipants (what is the population of interest?): the largest body of evidence evaluated people with cirrhosis and an acute episode of overt hepatic encephalopathy. Only a relatively small proportion had minimal hepatic encephalopathy and chronic overt hepatic encephalopathy; very few had acute liver failure.

Interventions (what are the interventions of interest?): flumazenil.

Comparisons (what are the comparisons of interest?): placebocontrolled randomised clinical trials.

Outcomes (what are the outcomes of interest?): all-cause mortality, hepatic encephalopathy, and adverse events; evidence evaluating the effect on health-related quality of life is also needed.

Time stamp (date of literature search): May 2017.

## A C K N O W L E D G E M E N T S

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\* Indicates the major publication for the study

## CHARACTERISTICS OF STUDIES

## Characteristics of included studies [ordered by study ID]

## Amodio 1997

Methods	Double-blind, single-centre, placebo-controlled RCT.	
NICHIOUS	<b>Cross-over design:</b> all participants underwent both intervention periods (receive flumazenil and placebo)	
Participants	<ul> <li>13 participants with cirrhosis with no evidence of overt hepatic encephalopathy but with abnormal brainstem evoked potentials (5 participants) or prolonged Number Connection Test times (6 participants), or both at baseline corresponding to a diagnosis of minimal hepatic encephalopathy</li> <li>Mean ± SD age: flumazenil/placebo: 54 ± 7 years.</li> <li>Proportion of men: 77%.</li> <li>Aetiology of cirrhosis: alcohol 77%; hepatitis B/C 15%.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): 0%.</li> </ul>	
Interventions	Intervention comparison: intravenous bolus flumazenil 1 mg followed by 4 boluse 0.5 mg every 30 minutes versus placebo (saline) Total dose of flumazenil: 3 mg. Washout period: 72 hours before cross-over to the alternative arm. Cointerventions: none described.	
Outcomes	Outcomes included in meta-analyses: none.	
Neuropsychiatric assessment	<ul><li>Baseline and post infusion:</li><li>Brainstem auditory evoked potentials;</li><li>Number Connection Test.</li></ul>	
Inclusion period (date)	Not described.	
Country	Italy.	
Notes	<b>Included data:</b> RCT did not describe outcomes for first intervention period. Therefore, we were unable to include the trial in our meta-analyses. The study report includes 2 tables containing data for the 5 participants with abnormal evoked potentials at baseline and the 6 participants with abnormal Number Connection Test times. There were no change in the group mean variables after flumazenil	

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described.
Allocation concealment (selection bias)	Low risk	Concealed drug containers.

## Amodio 1997 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data.
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	Low risk	Funding from the Italian Liver Foundation.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.
Barbaro 1998		

Methods	Double-blind, multi-centre, placebo-controlled RCT. <b>Cross-over design:</b> investigators crossed-over participants who did not respond to in- tervention during first period (remained in Grade III or IVa coma) to alternative inter- vention
Participants	527 participants with cirrhosis and overt hepatic encephalopathy (Grades III or IVa; Table 2), admitted to an intensive care unit. Diagnostic criteria corresponded to acute hepatic encephalopathy. Precipitating factors are described (Table 4). Mean $\pm$ SD age (grade III/IVa): flumazenil: 56 $\pm$ 11.5/53 $\pm$ 12 years; placebo: 48 $\pm$ 20/ 55 $\pm$ 13.5 years Proportion of men: 69%. Aetiology of cirrhosis: alcohol 40%; hepatitis B/C 59%. Proportion testing positive for benzodiazepines at baseline (Table 5): 10/527 (1.9%) participants.
Interventions	<ul> <li>Intervention comparison: intravenous infusion flumazenil 1 mg given over 3 to 5 minutes versus placebo (isotonic saline)</li> <li>Total dose of flumazenil: 1 mg.</li> <li>Cointerventions: lactulose 30 mL every 6 hours via nasogastric tube; antibiotics were given to 22 participants with sepsis in the flumazenil group and 8 in the placebo group</li> </ul>
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed for a maximum of 4 days after randomisation.

## Barbaro 1998 (Continued)

Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Coma grade at baseline (Table 2);</li> <li>Modified Glasgow Coma Scale (Table 2) assessed at 10 minutes before and evolution 10 minutes after the intervention for a maximum of 3 hours;</li> <li>Continuous electroencephalography recorded 15 minutes before and 10 minutes after the infusion (Table 3).</li> </ul>	
Inclusion period (date)	January 1993 to December 1997.	
Country	Italy.	
Notes	<b>Included data:</b> Serum benzodiazepines were detected in 10 participants (4 with Grade III and 6 with Grade IVa coma). The published paper provides no information about the distribution of these participants to flumazenil or placebo during the first intervention period. Therefore, we were unable to exclude these participants from the analyses. The trial reported on several serious adverse events (Table 6).	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated sequential list of block-randomised assignments
Allocation concealment (selection bias)	Low risk	Concealed ampoules of flumazenil and placebo.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel us- ing placebo.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment using placebo.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants included in analyses
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	Low risk	No information provided.
Other bias	Low risk	No other biases.
Overall assessment	Low risk	Low risk of bias.

Cadranel 1995

Methods	Double-blind, single-centre, placebo-controlled RCT. <b>Cross-over design:</b> participants who did not respond after 10 minutes during the first period received the alternative intervention		
Participants	<ul> <li>14 participants with cirrhosis experiencing 18 separate episodes of acute hepatic encephalopathy classified as Grade II to IV (Table 2). Precipitating factors are described (Table 4).</li> <li>Mean ± SD age: whole group 54.8 ± 7.7 years.</li> <li>Proportion of men: 71%.</li> <li>Aetiology of cirrhosis: alcohol 71%; hepatitis B/C 29%.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): 3/14 (21.4%) participants.</li> </ul>		
Interventions	Intervention comparison: continuous intravenous infusion flumazenil 0.1 mg/mL at 1 mL/minute flumazenil versus placebo (sodium edetate 1 mg). Investigators stopped the infusion after 10 minutes if participants showed improvement in electroencephalography or coma grade Total dose of flumazenil: 1 mg. Cointerventions: none described.		
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed after maximum of 3 days.		
Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Clinical assessment of mental status (Table 2) assessed at baseline and within 100 minutes after infusion;</li> <li>Electroencephalography graded using a 5-point scale (Table 3) assessed at baseline and within 10 minutes after infusion.</li> </ul>		
Inclusion period (date)	May 1988 to May 1990.		
Country	France.		
Notes	<b>Included data:</b> the trial included 14 participants who between them experienced 18 episodes of acute hepatic encephalopathy. 1 participant entered the trial once and 1 entered the trial 3 times. We included data from the first intervention period in our analyses. The published report described the number of participants who died after the second treatment period only (Table 6).		
Risk of bias			
Bias	Authors' judgement Support for judgement		

Dias	Ruthors Judgement	Support for Judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Concealed drug vials.

## Cadranel 1995 (Continued)

Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Outcome data were complete.
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in original pro- tocol or information in trial registries
For-profit funding	High risk	Hoffmann-La Roche Ltd. supplied flumazenil and placebo.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

## Dursun 2003

Methods	Double-blind, single-centre, placebo-controlled, parallel-arm RCT
Participants	<ul> <li>40 participants with cirrhosis and hepatic encephalopathy classified as subclinical (corresponding to minimal; 10 participants) or overt Grade I to III (30 participants; Table 2). Type of overt hepatic encephalopathy (acute or chronic) not specified Mean ± SD age: flumazenil: 44.5 ± 12.9 years; placebo: 43.7 ± 11.9 years.</li> <li>Proportion of men: 73%.</li> <li>Aetiology of cirrhosis: alcohol 0%; hepatitis B/C 100%.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): investigators did not screen for benzodiazepines.</li> </ul>
Interventions	Intervention comparison: intravenous infusion flumazenil 1 mg/hour for 5 hours versus placebo (saline) administered similarly Total dose of flumazenil: 5 mg. Cointerventions: lactulose 30 mL 6-hourly
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), serious adverse events (Table 6), and Number Connection Test assessed after a maximum of 5 hours
Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Clinical assessment of mental status (Table 2) assessed at baseline and every 30 minutes after infusion for a maximum of 5 hours;</li> </ul>

## **Dursun 2003** (Continued)

	<ul> <li>Glasgow Coma Score (Table 2) assessed at baseline;</li> <li>Electroencephalography (Table 3) assessed at baseline and 1 hour after infusion;</li> <li>Number Connection Test assessed at baseline and every 30 minutes after infusion for a maximum of 5 hours;</li> <li>Blood ammonia concentrations assessed at baseline.</li> </ul>
Inclusion period (date)	December 1999 to January 2002.
Country	Turkey.
Notes	<b>Included data:</b> the trial report included information about participants with minimal and overt hepatic encephalopathy. We have analysed these 2 groups separately

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Concealed drug containers.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants are included in analyses
Selective reporting (reporting bias)	Low risk	Trial describes clinically relevant outcomes. We had no access to information about out- comes described in the original protocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

Giger-Mateeva 1999

Methods	Double-blind, single-centre, placebo-controlled RCT. <b>Cross-over design:</b> investigators crossed over all participants to the alternative interven- tion			
Participants	10 participants with cirrhosis and no clinical evidence of overt hepatic encephalopathy; 5 participants had minimal hepatic encephalopathy based on the finding of either abnormal visual evoked potentials or Number Connection Test results <b>Age (range):</b> 40 to 60 years. <b>Proportion of men:</b> 80%. <b>Aetiology of cirrhosis:</b> alcohol 30%; hepatitis B/C 70%. <b>Proportion testing positive for benzodiazepines at baseline (Table 5):</b> 0%.			
Interventions	Intervention comparison: intravenous infusion flumazenil 1 mg over 2 minutes versus placebo Total dose of flumazenil: 1 mg. Washout period: 4 hours. Cointerventions: none reported.			
Outcomes	Outcomes included in meta-analyses: none.			
Neuropsychiatric assessment	<ul> <li>At baseline and post infusion:</li> <li>Visual evoked potential at baseline and every 8, 16, 24, 32, and 40 minutes after infusion;</li> <li>Visual reaction time at baseline and 5, 10, and 20 minutes after infusion;</li> <li>Auditory reaction time at baseline and 5, 10, and 20 minutes after infusion;</li> <li>Number Connection Test at baseline and 10 minutes after infusion.</li> </ul>			
Inclusion period (date)	Not described.			
Country	The Netherlands.			
Notes	<b>Included data:</b> trial did not include separate information about the first allocation period. Therefore, we were unable to include the trial in our meta-analyses			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Not described.		
Allocation concealment (selection bias)	Low risk	Concealed drug containers.		
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.		

## Giger-Mateeva 1999 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data.
Selective reporting (reporting bias)	Low risk	The trial describes clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

# Gooday 1995

Methods	Double-blind, single-centre, placebo-controlled RCT. <b>Cross-over design:</b> all participants were crossed over to alternative intervention
Participants	<ul> <li>10 participants with cirrhosis and subclinical (corresponding to minimal) hepatic encephalopathy diagnosed based on a score on the Digit Symbol Substitution test of &lt; 1 SD of the age-matched normative mean</li> <li>Mean age ± SD: 53.9 ± 7.4 years.</li> <li>Proportion of men: 80%.</li> <li>Aetiology of cirrhosis: alcohol 60%; hepatitis B/C 20%.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): apparently not performed (not specifically stated).</li> </ul>
Interventions	Intervention comparison: intravenous infusion flumazenil 0.2 mg over an unspecified time versus placebo (saline) Total dose of flumazenil: 0.2 mg. Washout period: 1 week. Cointerventions: none described.
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality and serious adverse events (Table 6) assessed at end of the intervention.
Neuropsychiatric assessment	At baseline: • Digit Symbol Substitution Test. At baseline and post infusion: • Simple reaction time; • Complex reaction time; • Auditory Verbal Learning Test; • Digit Symbol Substitution Test;

## **Gooday 1995** (Continued)

	• Digits forward and backwards. Duration of follow-up and timing of tests not described.	
Inclusion period (date)	Not described.	
Country	UK.	
Notes	<b>Included data:</b> we received additional (unpublished) information about the trial meth- ods and number of participants allocated to flumazenil/placebo during the first alloca- tion period via email in 2003 when conducting the previous version of this review. The trial did not evaluate the number of participants with an overall improvement in hepatic encephalopathy. Therefore, we were unable to include the trial in our analyses of this outcome measure	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random number table.
Allocation concealment (selection bias)	Low risk	Serially numbered opaque sealed envelopes used in administration of concealed drug containers
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data.
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	High risk	Primary investigators described a signifi- cant drug by order and group by drug by order interaction
Overall assessment	High risk	High risk of bias.

Gyr 1996

Methods	Double-blind, multi-centre, parallel-arm, placebo controlled RCT
Participants	<ul> <li>49 participants with cirrhosis and chronic overt hepatic encephalopathy (Grades I to III; Table 2).</li> <li>Mean age ± SD: flumazenil: 55.5 ± 9.4 years; placebo: 53.6 ± 10.3 years.</li> <li>Proportion of men: 69%.</li> <li>Aetiology of cirrhosis: alcohol 51%; hepatitis B/C 35%.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): 11% in flumazenil group; 5% in placebo group.</li> </ul>
Interventions	Intervention comparison: intravenous boluses of flumazenil 0.4 mg, 0.8 mg, and 1 mg at 1-minute intervals followed by a 3-hour infusion of flumazenil 1 mg/hour versus placebo (saline) Total dose of flumazenil: 5.2 mg. Cointerventions: none reported.
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed after a maximum of 4 weeks.
Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Clinical assessment of mental status (Table 2) at baseline and every 30 minutes for 5 hours then every 1 hour until 12 hours post infusion;</li> <li>Continuous (20 minutes) electroencephalography immediately after the infusion and then at 2 hours 40 minutes, 3 hours, and 7 hours 40 minutes post infusion (Table 3).</li> </ul>
Inclusion period (date)	Not reported.
Country	Switzerland (primary), France, Germany, Italy, Canada, the Netherlands, the UK, and Korea
Notes	<b>Included data:</b> authors reported intention-to-treat analyses including all participants randomised and a per-protocol analysis excluding protocol violators (25 participants). We included data on all participants in our analyses

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated list of random num- bers.
Allocation concealment (selection bias)	Low risk	Sealed envelopes used in double-blind ad- ministration of flumazenil and placebo
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Blinding of participants and personnel.

## Gyr 1996 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data.
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	High risk	Support from Hoffmann-La Roche Ltd.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

## Hermant 1991

Methods	Double-blind, parallel-arm, single-centre, placebo-controlled RCT	
Participants	<ul> <li>12 participants with cirrhosis and an acute episode of hepatic encephalopathy defined as Grade IIIa with severely abnormal electroencephalography changes, but a Glasgow Coma Score of &lt; 12 (Table 2).</li> <li>Proportion of men: not reported.</li> <li>Mean age ± SD: whole group 58.2 ± 5.4 years.</li> <li>Aetiology of liver disease: not reported.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): 0%.</li> </ul>	
Interventions	Intervention comparison: intravenous infusion flumazenil 0.2 mg/kg over 10 minutes versus placebo (saline) Total dose of flumazenil: 0.2 mg/kg. Cointerventions: none described.	
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality (Table 6) and serious adverse events (Table 6).	
Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Glasgow Coma Scale (Table 2);</li> <li>Electroencephalography (Table 3).</li> <li>The timing of assessments post infusion was not described.</li> </ul>	
Inclusion period (date)	Not described.	
Country	France.	

# Hermant 1991 (Continued)

Notes	Included data: the trial report did not specifically state the number of participants	
INDICS		
	with (or without) improvement in hepatic encephalopathy separately for the allocation	
	groups. Therefore, we were unable to include the trial in the analysis of this outcome	
	measure	
	Article published in French (full translation available).	

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Concealed drug containers.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Trial report gave the impression that there were no missing outcome data although this was not specifically stated
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

## Klotz 1989

Methods	Double-blind, cross-over, single-centre, placebo-controlled RCT
Participants	2 participants with cirrhosis and stable hepatic encephalopathy (Grade III). Description corresponded to chronic overt hepatic encephalopathy although this was not specifically stated <b>Proportion of men:</b> not reported. <b>Mean age:</b> not reported. <b>Aetiology of liver disease:</b> alcohol 100%.

## Klotz 1989 (Continued)

	<b>Proportion testing positive for benzodiazepines at baseline</b> (Table 5): apparently not performed (not specifically stated).
Interventions	Intervention comparison: intravenous infusion flumazenil 1 mg over 1 minute versus placebo Total dose of flumazenil: 1 mg. Washout period: not specified. Cointerventions: not reported.
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality and serious adverse events (Table 6) assessed for a maximum of 2 hours post interventions.
Neuropsychiatric assessment	Clinical assessment of mental status: assessed after 2 hours (no specific score; timing not specified)
Inclusion period (date)	Not described.
Country	Germany.
Notes	<b>Included data:</b> investigators described the design as cross-over but did not provide data from the first intervention period. Therefore, we were unable to include the trial in our analyses

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment not described.
Blinding of participants and personnel (performance bias) All outcomes	Unclear risk	Trial described as double blind and placebo controlled. However, trial only reported as a letter and the type of placebo (or mode of administration) is not described
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Trial described as double blind and placebo controlled. However, trial only reported as a letter and the type of placebo (or mode of administration) is not clearly described
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants included in analyses
Selective reporting (reporting bias)	Low risk	Trial gave impression that participants sur- vived although this was not specifically stated. We had no access to information

#### Klotz 1989 (Continued)

		about outcomes described in the original protocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	High risk	Trial only included 2 participants.
Overall assessment	High risk	High risk of bias.
Lacetti 2000		
Methods	Double-blind, single-centre, par	illel-arm placebo-controlled RCT
Participants	<ul> <li>54 participants with cirrhosis and acute hepatic encephalopathy (Grade III or IV). Precipitating factors are described (Table 4).</li> <li>Mean age ± SD: flumazenil: 59.6 ± 6.0 years; placebo: 57.7 ± 5.4 years.</li> <li>Proportion of men: 54%.</li> <li>Aetiology of cirrhosis: hepatitis B/C 100%.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): 0%.</li> </ul>	
Interventions	Intervention comparison: intravenous infusion flumazenil 0.4 mg/mL at 10 mL/minute for 5 minutes versus placebo (saline) Total dose of flumazenil: 2 mg. Cointerventions: lactulose enemas; branch-chain amino acids.	
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed for maximum of 24 hours after intervention.	
Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Clinical assessment of mental status (score not specified) assessed at baseline;</li> <li>Glasgow Coma Score (Table 2) assessed at baseline and every 30 minutes for the first 6 hours and then every 6 hours for 24 hours post infusion.</li> </ul>	
Inclusion period (date)	January 1997 to December 1997	
Country	Italy.	
Notes	<b>Included data:</b> we included data on all participants in our analyses. <b>Notes about the design:</b> investigators repeated the intervention once after 3 hours in non-responders (no improvement in neurological status) or immediately if they detected an improvement followed by a relapse. The report did not include information about the number of participants who received a second infusion. In the results section of the report the investigators stipulate that the second infusion was the same as the one first received	
Risk of bias		
Bias	Authors' judgement	Support for judgement

### Lacetti 2000 (Continued)

Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Double-blind administration of flumazenil and placebo.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel us- ing placebo.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment using placebo.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants were included in the analyses
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

#### Li 2009

Methods	Double-blind, single-centre, parallel-arm RCT.
Participants	<ul> <li>72 participants with overt hepatic encephalopathy (Grade III or IV) associated with cirrhosis (65%) or fulminant hepatic failure (35%). Diagnostic criteria for participants with cirrhosis corresponded to acute hepatic encephalopathy</li> <li>Mean age ± SD: flumazenil: 55.4 ± 6.6 years; placebo: 56.8 ± 7.9 years.</li> <li>Proportion of men: 63%.</li> <li>Aetiology of cirrhosis: not reported.</li> <li>Proportion testing positive for benzodiazepines at baseline (Table 5): apparently not performed (not specifically stated).</li> </ul>
Interventions	Intervention comparison: slow intravenous injection flumazenil 0.5 mg followed by intravenous infusion of flumazenil 1 mg of over 30 minutes versus placebo (saline) Total dose of flumazenil: 1.5 mg. Cointerventions: lactulose enemas, L-ornithine L-aspartate
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed after a maximum of 2 weeks.

## Li 2009 (Continued)

Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Clinical scale (not specified) assessed at baseline;</li> <li>Glasgow Coma Score (Table 2) assessed at baseline and after 2 hours;</li> <li>Electroencephalography assessed at baseline and after 2 hours.</li> </ul>	
Inclusion period (date)	May 2006 to July 2008.	
Country	China.	
Notes	<b>Included data:</b> the trial report did not provide separate information on participants with cirrhosis and participants with acute liver failure. Therefore, we conducted a sensitivity analysis excluding this trial	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Double-blind administration of flumazenil and placebo.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants included in the analyses
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in original pro- tocol or information in trial registries
For-profit funding	Unclear risk	No information provided.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

Pomier-Layrargues 1994

Methods	Double-blind, single-centre, cross-over, plac <b>Cross-over design:</b> investigators only crosse IV coma, 24 hours after the first study perio	ed over participants who remained in Grade
Participants	21 participants with cirrhosis and acute hepatic encephalopathy (Grade IV). Precipitating factors are described (Table 4). Mean age ± SD: flumazenil: 52.7 ± 5.4 years; placebo: 57.4 ± 9.0 years. Proportion of men: 81%. Aetiology of cirrhosis: alcohol 62%; hepatitis B/C 5%. Proportion testing positive for benzodiazepines at baseline (Table 5): 4/21 (19%) participants.	
Interventions	Intervention comparison: intravenous infusion flumazenil 2 mg over 5 minutes versus placebo (saline) Total dose of flumazenil: 2 mg. Washout period: 24 hours. Cointerventions: lactulose 30 mL 4 times daily.	
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed after a maximum follow-up of 24 hours.	
Neuropsychiatric assessment	<ul> <li>Baseline and post infusion:</li> <li>Modified Glasgow Coma Scale (Table 2) undertaken at baseline and every 15 minutes for up to 5.5 hours post infusion;</li> <li>Continuous electroencephalography (Table 3) 15 minutes before and 15 minutes after the infusion.</li> </ul>	
Inclusion period (date)	March 1988 to February 1992.	
Country	Canada.	
Notes	Included data: we only included data from the first treatment period in our analyses	
Risk of bias		
Bias	Authors' judgement Support for judgement	
Random sequence generation (selection bias)	Low risk	Random numbers.
Allocation concealment (selection bias)	Low risk	Blinded administration of flumazenil or placebo.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.

## **Pomier-Layrargues 1994** (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants included in analyses
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	High risk	Technical assistance from Hoffmann-La Roche Ltd., Canada and Nutley, NJ, USA
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.
Van der Rijt 1995		
Methods	Double-blind, single-centre, cross-over, placebo-controlled RCT <b>Cross-over design:</b> all participants (except 2 who underwent transplantation) received flumazenil and placebo	
Participants	18 participants with hepatic encephalopathy secondary to acute liver failure (28%) or cirrhosis (82%), who had an arterial blood ammonia > 30 $\mu$ mol/L, and an abnormal electroencephalography despite at least 24 hours of treatment with a low protein diet and lactulose alone or with neomycin. Precipitating factors are described (Table 4). <b>Mean age ± SD:</b> whole group 48.56 ± 14.67 years. <b>Proportion of men:</b> 39%. <b>Aetiology of cirrhosis:</b> alcohol 38%; hepatitis B/C 15%. <b>Proportion testing positive for benzodiazepines at baseline (Table 5):</b> 0%.	
Interventions	<ul> <li>Intervention comparison 1:</li> <li>First 9 participants: intravenous infusion flumazenil 0.1 mg/minute over 10 minutes; 4 hours later given a bolus injection flumazenil 0.5 mg followed by a continuous infusion of flumazenil 0.25 mg/hour for 3 days versus infusion vehicle alone</li> <li>Total dose of flumazenil: 19.5 mg.</li> <li>Washout period: 24 hours.</li> <li>Intervention comparison 2:</li> <li>Second 9 participants: intravenous infusion of flumazenil 0.1 mg/minute over 10 minutes versus infusion vehicle alone</li> <li>Total dose of flumazenil: 19.5 mg.</li> <li>Washout period 24 hours.</li> <li>Total dose of flumazenil: 19.5 mg.</li> <li>Washout period 24 hours.</li> <li>Cointerventions: protein restriction, lactulose alone or with neomycin.</li> </ul>	

## Van der Rijt 1995 (Continued)

Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed after a maximum of 3 days.
Neuropsychiatric assessment	<ul> <li>First 9 participants: baseline and post infusion: <ul> <li>Blood ammonia concentration assessed at baseline;</li> <li>Mental status assessed clinical scale (Table 2) at baseline and after 15 minutes and then 24, 48, and 72 hours;</li> <li>Eectroencephalography, conventional and spectral grading (Table 3) recorded at baseline and after 15 minutes and then 24, 48, and 72 hours.</li> </ul> </li> <li>Second 9 participants: baseline and post infusion: <ul> <li>Blood ammonia concentration assessed at baseline;</li> <li>Mental status assessed clinical scale (Table 2) at baseline and after 15 minutes;</li> <li>Electroencephalography, conventional and spectral grading (Table 3) recorded at baseline and after 15 minutes;</li> </ul> </li> </ul>
Inclusion period (date)	February 1987 to February 1990.
Country	The Netherlands.
Notes	<b>Included data:</b> Two patients were withdrawn on day one of the study to undergo liver transplantation thus only 16 people took part in the full cross-over study. The study involved people with hepatic encephalopathy associated with cirrhosis and with acute liver failure; the trial data were not provided separately for these two groups so no separate analysis can be performed by aetiology of the hepatic encephalopathy. We only include data from the first treatment period in our analyses

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Used concealed drug containers.
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No missing outcome data and all partici- pants were included in analyses
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information

## Van der Rijt 1995 (Continued)

		about outcomes described in original pro- tocol or information in trial registries
For-profit funding	High risk	Support provided by Hoffmann-La Roche B.V., Mijdrecht, The Netherlands
Other bias	Unclear risk	Investigators simplified the intervention regimen after inclusion of the first 9 par- ticipants as the second period of the 72- hour infusion was too demanding for par- ticipants. The effect that this has on bias control was unclear
Overall assessment	High risk	High risk of bias.

## Zhu 1998

Methods	Double-blind, single-centre, parallel-arm, p	lacebo-controlled RCT
Participants	cipitating factors are described (Table 4). Mean age ± SD: flumazenil: 62.2 ± 2.7 yea Proportion of men: 69%. Aetiology of cirrhosis: alcohol 80%; hepat	
Interventions	Intervention comparison: intravenous infusion flumazenil 1 mg over 5 minutes versus placebo (saline) Total dose of flumazenil: 1 mg. Cointerventions: intravenous branched-chain amino acids.	
Outcomes	<b>Outcomes included in meta-analyses:</b> mortality, hepatic encephalopathy (Table 1), and serious adverse events (Table 6) assessed for a maximum of 2 weeks (until death or discharge)	
Neuropsychiatric assessment	<ul><li>Baseline and post infusion:</li><li>Clinical assessment of hepatic encephalopathy (Table 2).</li></ul>	
Inclusion period (date)	April 1995 to March 1996.	
Country	China.	
Notes	Included data: all participants were included in the analyses.	
Risk of bias		
Bias	Authors' judgement	Support for judgement

## Zhu 1998 (Continued)

Random sequence generation (selection bias)	Low risk	Random numbers with stratified block ran- domisation.
Allocation concealment (selection bias)	Low risk	Administration of concealed drug contain- ers with sealed, opaque, serially numbered envelopes
Blinding of participants and personnel (performance bias) All outcomes	Low risk	Blinding of participants and personnel.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Blinding of outcome assessment.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Missing outcome data not described.
Selective reporting (reporting bias)	Low risk	Trial described clinically relevant out- comes. We had no access to information about outcomes described in the original protocol or information in trial registries
For-profit funding	High risk	Roche supplied the flumazenil.
Other bias	Low risk	No other biases.
Overall assessment	High risk	High risk of bias.

RCT: randomised clinical trial; SD: standard deviation.

# Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Bansky 1989	Prospective study including 14 participants with cirrhosis and overt hepatic encephalopathy. The investigators reported an improvement in mental status in 71% of participants within minutes of receiving intravenous flumazenil lasting for 1 to 2 hours. Participants also received lactulose . Six participants died. The study was excluded as it did not include a control group
Devictor 1995	Prospective study evaluating 7 children with fulminant hepatic failure awaiting emergency liver transplantation. The investigators reported that flumazenil injection led to a transient improvement in mental status in 1 child but had no effect on mental status in the remaining six. The study was excluded as none of the participants had hepatic encephalopathy associated with cirrhosis and it did not include a control group

(Continued)

Golubovic 1999	Prospective study including 10 participants with alcohol-related cirrhosis and overt hepatic encephalopathy clas- sified as grade IV based on an assessment of mental status, electroencephalography, and visual evoked responses. The investigators reported an improvement in mental status in 8/10 participants. Six participants died within 1 year. The study was excluded as it did not include a control group
Grimm 1988	Prospective study including 17 participants (2 children) with hepatic encephalopathy associated with acute liver failure (9 participants) or cirrhosis (8 participants). Cointerventions included lactulose, branched-chain amino acids, antibiotics, diuretics, histamine-receptor antagonists, human albumin, and fresh frozen plasma. Transient improvement in the manifestations of hepatic encephalopathy was seen following flumazenil in 4 (44%) participants with fulminant hepatic failure and 5 (63%) with cirrhosis. Mortality was not reported. This study was excluded as it did not include a control group
Jia 1999	Open, single-centre, non-randomised study which is included in the sensitivity analyses of serious adverse events. The study involved 22 participants with cirrhosis and overt hepatic encephalopathy (Grades I-III using West Haven criteria) recruited between April 1996 and September 1997 Intervention comparison: 12 participants received an intravenous bolus of flumazenil 0.5 mg followed by an intravenous infusion of flumazenil in a dose of 1.0 mg over 4 hours for an unspecified period of time. The remaining 10 participants received Xing-Nao-Jing, a traditional Chinese medicine also given as an intravenous infusion Total dose of flumazenil: 1.5 mg. Outcomes: The study report stated that 2 participants in the flumazenil group died of liver failure but the time of death in relation to the intervention was not specified and study did not specifically state if there were any deaths in control group. The article was published in Chinese but a translation was available. The study was excluded as it did not contain a control group
Kapczinski 1995	Double-blind, cross-over, placebo-controlled, single-centre randomised clinical trial including 20 liver transplant candidates with cirrhosis. The trial is included in the sensitivity analyses of serious adverse events. The main objective of trial was to evaluate the differential effects of flumazenil on cognitive function and anxiety in people with alcohol-related (10 participants) or non-alcohol-related (10 participants) cirrhosis. None of the included participants had evidence of overt hepatic encephalopathy. The investigators evaluated a range of psychometric tests and reported the results as group mean values. No information was provided about the number of participants with abnormal test results <b>Proportion of men:</b> 60%. <b>Mean ± SD age:</b> alcohol-related cirrhosis: 47.7 ± 10.5 years; non-alcoholic cirrhosis: 48.4 ± 11.7 years <b>Proportion testing positive for benzodiazepines at baseline:</b> not tested <b>Intervention:</b> intravenous infusion flumazenil 0.1 mg/minute for 10 minutes then 0.05 mg/minute for 20 minutes versus placebo (saline) <b>Total dose of flumazenil:</b> 2 mg. <b>Washout period:</b> 60 minutes. <b>Outcomes:</b> The investigators reported changes in psychometric tests for participants with alcohol-related or non-alcohol-related cirrhosis without providing an overall estimate of numbers with (or without) improved manifestations. The trial did not report any deaths or serious adverse events The study was excluded because none of the participants had hepatic encephalopathy
Marsepoil 1990	Open, single-centre, prospective, non-randomised study included in the sensitivity analyses of serious adverse events at 48 hours. The study involved 25 participants with alcohol-related cirrhosis and acute hepatic en- cephalopathy, 13 of whom received flumazenil. The proportion of men and the mean age of participants was not reported <b>Proportion testing positive for benzodiazepines at baseline:</b> not mentioned

#### (Continued)

	<ul> <li>Intervention: intravenous bolus of flumazenil 0.2 mg every ten minutes until improvement in clinical status up to a maximum total dose of 2 mg followed by a continuous maintenance infusion of 0.3 mg. per hour for 48 hours</li> <li>Total dose of flumazenil: maximum 16.4 mg.</li> <li>Outcomes: .The Investigators reported that the mortality rates were similar in the flumazenil and control groups, but did not provide information on the number of participants who died. Published in French but a translation was available.The study was excluded as it was not randomised</li> </ul>
Ozyilkan 1997	Prospective study evaluating the effect of 30-minute, incremental intravenous boluses of flumazenil in 11 participants with cirrhosis (6 stage 0,4 stage one, one stage 2 hepatic encephalopathy) in whom baseline somatosensory evoked potentials were abnormal. Four patients (36%) showed a clear improvement in evoked potentials with flumazenil. Mortality was not described. The study was excluded as it did not include a control group
Wu 2001	Randomised clinical trial comparing intravenous flumazenil plus lactulose enemas versus flumazenil alone. A total of 20 participants (18 men) with cirrhosis and hepatic encephalopathy were included. The maximum dose of flumazenil was 9 mg. The dose was adjusted based on the clinical effect. Investigators assessed hepatic encephalopathy based on Conn Criteria and defined an improvement from Grade IV to I within 6 hours as clinically significant. None of the included participants died and all 12 in the flumazenil plus lactulose group and all 8 in the flumazenil group showed improved manifestations of hepatic encephalopathy. The study was excluded as there was no placebo arm

# Characteristics of ongoing studies [ordered by study ID]

#### Yale 2014

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Trial name or title	Treatment of Hepatic Encephalopathy with Flumazenil and Change in Cortical Gamma Aminobutyric Acid Levels in MRS [magnetic resonance spectroscopy]
Methods	Randomised clinical trial.
Participants	Participants with non-alcoholic cirrhosis and hepatic encephalopathy
Interventions	Flumazenil and placebo.
Outcomes	Recovery from hepatic encephalopathy and change in cortical gamma aminobutyric acid levels
Starting date	November 2014.
Contact information	Deanna Martin, deanna.martin@yale.edu and Amanda Brennan amanda.brennan@yale.edu.
Notes	Estimated completion date: June 2017.

## DATA AND ANALYSES

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 All-cause mortality	11	842	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.48, 1.16]
1.1 Overt hepatic encephalopathy	10	822	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.48, 1.16]
1.2 Minimal hepatic encephalopathy	2	20	Risk Ratio (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
2 All-cause mortality and bias control	12	844	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.48, 1.16]
2.1 Low risk of bias	1	527	Risk Ratio (M-H, Random, 95% CI)	0.76 [0.37, 1.53]
2.2 High risk of bias	11	317	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.43, 1.31]
3 All-cause mortality and trial design	12		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
3.1 Cross-over	6	592	Risk Ratio (M-H, Random, 95% CI)	0.69 [0.35, 1.34]
3.2 Parallel-arm	6	252	Risk Ratio (M-H, Random, 95% CI)	0.81 [0.45, 1.44]
4 All-cause mortality and duration of follow-up	11		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
$4.1 \leq 1 \text{ day}$	5	176	Risk Ratio (M-H, Random, 95% CI)	0.85 [0.38, 1.87]
4.2 > 1 day	6	666	Risk Ratio (M-H, Random, 95% CI)	0.71 [0.42, 1.21]
5 Hepatic encephalopathy	9	824	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.71, 0.80]
5.1 Overt hepatic encephalopathy	9	814	Risk Ratio (M-H, Random, 95% CI)	0.73 [0.67, 0.80]
5.2 Minimal hepatic encephalopathy	1	10	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.41, 1.39]
6 Hepatic encephalopathy and bias control	9	824	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.71, 0.80]
6.1 Low risk of bias	1	527	Risk Ratio (M-H, Random, 95% CI)	0.78 [0.72, 0.84]
6.2 High risk of bias	8	297	Risk Ratio (M-H, Random, 95% CI)	0.69 [0.61, 0.78]
7 Hepatic encephalopathy and trial design	9		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
7.1 Cross-over	4	584	Risk Ratio (M-H, Random, 95% CI)	0.77 [0.72, 0.83]
7.2 Parallel-arm	5	240	Risk Ratio (M-H, Random, 95% CI)	0.69 [0.59, 0.79]
8 Hepatic encephalopathy and duration of follow-up	9		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
$8.1 \leq 1 \text{ day}$	4	164	Risk Ratio (M-H, Random, 95% CI)	0.71 [0.60, 0.83]
8.2 > 1 day	5	660	Risk Ratio (M-H, Random, 95% CI)	0.72 [0.62, 0.84]
9 Hepatic encephalopathy and acute liver failure	9		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
9.1 Cirrhosis	7	734	Risk Ratio (M-H, Random, 95% CI)	0.76 [0.71, 0.82]
9.2 Acute liver failure or cirrhosis	2	90	Risk Ratio (M-H, Random, 95% CI)	0.62 [0.47, 0.80]
10 Number Connection Test	1		Mean Difference (IV, Random, 95% CI)	Totals not selected
11 All-cause mortality and acute liver failure	11	842	Risk Ratio (M-H, Random, 95% CI)	0.75 [0.48, 1.16]

# Comparison 1. Flumazenil versus placebo

11.1 Participants with	9	752	Risk Ratio (M-H, Random, 95% CI)	0.71 [0.45, 1.14]
cirrhosis				
11.2 Participants with	2	90	Risk Ratio (M-H, Random, 95% CI)	1.06 [0.31, 3.62]
cirrhosis or acute liver failure				

#### Analysis I.I. Comparison I Flumazenil versus placebo, Outcome I All-cause mortality.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: I All-cause mortality

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Overt hepatic encephalopathy					
Barbaro 1998	I 3/265	17/262	-	39.0 %	0.76 [ 0.37, 1.53 ]
Cadranel 1995	1/9	2/5		4.2 %	0.28 [ 0.03, 2.35 ]
Dursun 2003	0/16	0/14			Not estimable
Gyr 1996	4/28	5/21		13.6 %	0.60 [ 0.18, 1.97 ]
Hermant 1991	0/6	0/6			Not estimable
Lacetti 2000	6/28	5/26		17.1 %	.   [ 0.39, 3.22 ]
Li 2009	5/39	4/33	_ <b>_</b>	12.7 %	1.06 [ 0.31, 3.62 ]
Pomier-Layrargues 1994	0/11	0/10			Not estimable
Van der Rijt 1995	0/9	0/9			Not estimable
Zhu 1998	3/13	5/12		13.4 %	0.55 [ 0.17, 1.83 ]
Subtotal (95% CI)	424	398	•	100.0 %	0.75 [ 0.48, 1.16 ]
Total events: 32 (Flumazenil), 38 (Plac	ebo)				
Heterogeneity: $Tau^2 = 0.0$ ; Chi <sup>2</sup> = 2.0	)5, df = 5 (P = 0	.84); I <sup>2</sup> =0.0%			
Test for overall effect: $Z = 1.28$ (P = 0	0.20)				
2 Minimal hepatic encephalopathy					
Dursun 2003	0/4	0/6			Not estimable
Gooday 1995	0/5	0/5			Not estimable
Subtotal (95% CI)	9	11			Not estimable
Total events: 0 (Flumazenil), 0 (Placeb	0)				
Heterogeneity: not applicable					
Test for overall effect: not applicable					
Total (95% CI)	433	409	•	100.0 %	0.75 [ 0.48, 1.16 ]
Total events: 32 (Flumazenil), 38 (Plac	ebo)				
	,				
			0.01 0.1 1 10 100		
			Favours flumazenil Favours placebo		(Continued

Study or subgroup	Flumazenil	Placebo	Risk Ratic M- H.Random,955		( Continued) Risk Ratio M- H.Random.95%
	n/N	n/N	H,Kandom,95; Cl	70	H,Kandom,95% Cl
Heterogeneity: $Tau^2 = 0.0$ ; Ch	$i^2 = 2.05$ , df = 5 (P = 0	0.84); l <sup>2</sup> =0.0%			
Test for overall effect: $Z = 1.28$	B (P = 0.20)				
Test for subgroup differences:	Not applicable				
			0.01 0.1 1 10	) 100	
			Favours flumazenil Favou	urs placebo	

#### Analysis 1.2. Comparison I Flumazenil versus placebo, Outcome 2 All-cause mortality and bias control.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: 2 All-cause mortality and bias control

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random,955 Cl
I Low risk of bias					
Barbaro 1998	13/265	17/262	-	39.0 %	0.76 [ 0.37, 1.53 ]
Subtotal (95% CI)	265	262	•	39.0 %	0.76 [ 0.37, 1.53 ]
Total events: 13 (Flumazeni), 17 ( Heterogeneity: not applicable Test for overall effect: $Z = 0.78$ (F	× ,				
2 High risk of bias Cadranel 1995	1/9	2/5		4.2 %	0.28 [ 0.03, 2.35 ]
Dursun 2003	0/20	0/20			Not estimable
Gooday 1995	0/5	0/5			Not estimable
Gyr 1996	4/28	5/21		13.6 %	0.60 [ 0.18, 1.97 ]
Hermant 1991	0/6	0/6			Not estimable
Klotz 1989	0/1	0/1			Not estimable
Lacetti 2000	6/28	5/26	-	17.1 %	.   [ 0.39, 3.22 ]
Li 2009	5/39	4/33	_ <b>-</b>	12.7 %	1.06 [ 0.31, 3.62 ]

(Continued  $\dots$ )

Study or subgroup	Flumazenil	Placebo	Risk Rati		( Continued) Risk Ratio M-
	n/N	n/N	H,Random,95 Cl	%	H,Random,95% Cl
Pomier-Layrargues 1994	0/11	0/10			Not estimable
Van der Rijt 1995	0/9	0/9			Not estimable
Zhu 1998	3/13	5/12		13.4 %	0.55 [ 0.17, 1.83 ]
Subtotal (95% CI)	169	148	•	61.0 %	0.75 [ 0.43, 1.31 ]
Total events: 19 (Flumazenil), 2	I (Placebo)				
Heterogeneity: Tau <sup>2</sup> = 0.0; Chi	<sup>2</sup> = 2.05, df = 4 (P = 0	.73); l <sup>2</sup> =0.0%			
Test for overall effect: Z = 1.01	(P = 0.31)				
Total (95% CI)	434	410	•	100.0 %	0.75 [ 0.48, 1.16 ]
Total events: 32 (Flumazenil), 3	8 (Placebo)				
Heterogeneity: Tau <sup>2</sup> = 0.0; Chi	<sup>2</sup> = 2.05, df = 5 (P = 0	.84); l <sup>2</sup> =0.0%			
Test for overall effect: $Z = 1.28$	(P = 0.20)				
Test for subgroup differences: (	$Chi^2 = 0.00, df = 1 (P = 1)$	= 0.98), I <sup>2</sup> =0.0%			
			0.01 0.1 1	0 100	
			Favours flumazenil Favo	ours placebo	

## Analysis 1.3. Comparison I Flumazenil versus placebo, Outcome 3 All-cause mortality and trial design.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: 3 All-cause mortality and trial design

I Cross-over Barbaro 1998 Cadranel 1995 Gooday 1995 Klotz 1989 Pomier-Layrargues 1994 Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: Z = 1.11 (P = 0.2 2 Parallel-arm Dursun 2003	df = 1 (P = 0.	n/N 17/262 2/5 0/5 0/1 0/10 0/9 <b>292</b> .38); I <sup>2</sup> =0.0%	H,Random,95% Cl	90.3 % 9.7 % <b>100.0 %</b>	H,Random,9 Cl 0.76 [ 0.37, 1.53 ] 0.28 [ 0.03, 2.35 ] Not estimable Not estimable Not estimable Not estimable <b>0.69 [ 0.35, 1.34 ]</b>
Barbaro 1998 Cadranel 1995 Gooday 1995 Klotz 1989 Pomier-Layrargues 1994 Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: Z = 1.11 (P = 0.2 2 Parallel-arm	1/9 0/5 0/1 0/11 0/9 <b>300</b> 00) , df = 1 (P = 0.	2/5 0/5 0/1 0/10 0/9 <b>292</b>	•	9.7 %	0.28 [ 0.03, 2.35 ] Not estimable Not estimable Not estimable Not estimable
Cadranel 1995 Gooday 1995 Klotz 1989 Pomier-Layrargues 1994 Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: Z = 1.11 (P = 0.2 2 Parallel-arm	1/9 0/5 0/1 0/11 0/9 <b>300</b> 00) , df = 1 (P = 0.	2/5 0/5 0/1 0/10 0/9 <b>292</b>		9.7 %	0.28 [ 0.03, 2.35 ] Not estimable Not estimable Not estimable Not estimable
Gooday 1995 Klotz 1989 Pomier-Layrargues 1994 Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: Z = 1.11 (P = 0.2 2 Parallel-arm	0/5 0/1 0/11 0/9 <b>300</b> xo) . df = 1 (P = 0.	0/5 0/1 0/10 0/9 <b>292</b>	-		Not estimable Not estimable Not estimable Not estimable
Klotz 1989 Pomier-Layrargues 1994 Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: Z = 1.11 (P = 0.2 2 Parallel-arm	0/1 0/11 0/9 <b>300</b> co) , df = 1 (P = 0.	0/1 0/10 0/9 <b>292</b>	-	100.0 %	Not estimable Not estimable Not estimable
Pomier-Layrargues 1994 Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: Z = 1.11 (P = 0.2 2 Parallel-arm	0/11 0/9 <b>300</b> odf = 1 (P = 0.	0/10 0/9 <b>292</b>	-	100.0 %	Not estimable Not estimable
Van der Rijt 1995 <b>Subtotal (95% CI)</b> Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: $Z = 1.11$ (P = 0.2 2 Parallel-arm	0/9 <b>300</b> po) df = 1 (P = 0.	0/9 <b>292</b>	-	100.0 %	Not estimable
Subtotal (95% CI) Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: $Z = 1.11$ (P = 0.2 2 Parallel-arm	<b>300</b> bo) df = 1 (P = 0.	292	•	100.0 %	
Total events: 14 (Flumazenil), 19 (Placeb Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> = 0.76, Test for overall effect: $Z = 1.11$ (P = 0.2 2 Parallel-arm	oo) , df = 1 (P = 0.		•	100.0 %	0.69 [ 0.35, 1.34 ]
Heterogeneity: $Tau^2 = 0.0$ ; $Chi^2 = 0.76$ , Test for overall effect: $Z = 1.11$ (P = 0.2 2 Parallel-arm	df = 1 (P = 0.	.38); I <sup>2</sup> =0.0%			
Darsan 2005	0/20	0/20			Not estimable
	0/20	0/20			Not estimable
Gyr 1996	4/28	5/21		24.0 %	0.60 [ 0.18, 1.97 ]
Hermant 1991	0/6	0/6			Not estimable
Lacetti 2000	6/28	5/26	-	30.1 %	.   [ 0.39, 3.22 ]
Li 2009	5/39	4/33		22.3 %	1.06 [ 0.31, 3.62 ]
Zhu 1998	3/13	5/12		23.6 %	0.55 [ 0.17, 1.83 ]
Subtotal (95% CI)	134	118	•	100.0 %	0.81 [ 0.45, 1.44 ]
Total events: 18 (Flumazenil), 19 (Placeb Heterogeneity: $Tau^2 = 0.0$ ; $Chi^2 = 1.16$ ,	df = 3 (P = 0.	.76); I <sup>2</sup> =0.0%			
Test for overall effect: $Z = 0.73$ (P = 0.4	,	- 0.72) 12 -0.000			
Test for subgroup differences: $Chi^2 = 0$ .	13, dt = 1 (P =	= 0.72), 14 =0.0%			

# Analysis I.4. Comparison I Flumazenil versus placebo, Outcome 4 All-cause mortality and duration of follow-up.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

#### Comparison: I Flumazenil versus placebo

Outcome: 4 All-cause mortality and duration of follow-up

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
$  \leq  $ day					
Dursun 2003	0/20	0/20			Not estimable
Gyr 1996	4/28	5/21		44.4 %	0.60 [ 0.18, 1.97 ]
Hermant 1991	0/6	0/6			Not estimable
Lacetti 2000	6/28	5/26	-	55.6 %	.   [ 0.39, 3.22 ]
Pomier-Layrargues 1994	0/11	0/10			Not estimable
Subtotal (95% CI)	93	83	+	100.0 %	0.85 [ 0.38, 1.87 ]
Test for overall effect: Z = 0.41 2 > 1 day		17/2/2		E / 2 0/	
	(P = 0.68)				
Barbaro 1998	13/265	17/262		56.3 %	0.76 [ 0.37, 1.53 ]
Cadranel 1995	1/9	2/5		6.1 %	0.28 [ 0.03, 2.35 ]
Gooday 1995	0/5	0/5			Not estimable
Li 2009	5/39	4/33		18.3 %	1.06 [ 0.31, 3.62 ]
Van der Rijt 1995	0/9	0/9			Not estimable
Zhu 1998	3/13	5/12		19.3 %	0.55 [ 0.17, 1.83 ]
Subtotal (95% CI)	340	326	•	100.0 %	0.71 [ 0.42, 1.21 ]
Total events: 22 (Flumazenil), 28 Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup> Test for overall effect: Z = 1.26 Test for subgroup differences: Cl	= 1.34, df = 3 (P = 0) (P = 0.21)	,			

0.01 0.1 1 10 100 Favours flumazenil Favours placebo

## Analysis 1.5. Comparison I Flumazenil versus placebo, Outcome 5 Hepatic encephalopathy.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

#### Outcome: 5 Hepatic encephalopathy

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratic
	n/N	n/N	H,Random,95% Cl		H,Random, C
l Overt hepatic encephalopath	у				
Barbaro 1998	199/265	253/262		74.9 %	0.78 [ 0.72, 0.84 ]
Cadranel 1995	4/10	8/8		0.8 %	0.43 [ 0.21, 0.90
Dursun 2003	9/16	4/ 4		2.2 %	0.58 [ 0.38, 0.89
Gyr 1996	21/28	21/21	-	8.1 %	0.76 [ 0.61, 0.95
Lacetti 2000	6/28	12/26	<b>.</b>	0.6 %	0.46 [ 0.20, 1.06
Li 2009	22/39	31/33	+	4.8 %	0.60 [ 0.45, 0.80
Pomier-Layrargues 1994	8/11	10/10		2.7 %	0.74 [ 0.50, 1.09
Van der Rijt 1995	5/9	7/9	-+	0.9 %	0.71 [ 0.36, 1.41
Zhu 1998	10/13	12/12	-	3.9 %	0.78 [ 0.57, 1.08
Subtotal (95% CI)	419	395	•	<b>98.9</b> %	0.73 [ 0.67, 0.80
Total events: 284 (Flumazenil), 3	368 (Placebo)				
Heterogeneity: $Tau^2 = 0.00$ ; Ch	$i^2 = 9.02$ , df = 8 (P =	0.34); I <sup>2</sup> =I I%			
Test for overall effect: $Z = 6.60$	(P < 0.00001)				
2 Minimal hepatic encephalopat	thy				
Dursun 2003	3/4	6/6	-+	1.1 %	0.75 [ 0.41, 1.39
Subtotal (95% CI)	4	6	•	1.1 %	0.75 [ 0.41, 1.39
Total events: 3 (Flumazenil), 6 (I	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 0.91	(P = 0.36)				
Total (95% CI)	423	401	•	100.0 %	0.75 [ 0.71, 0.80
Total events: 287 (Flumazenil), 3	374 (Placebo)				
Heterogeneity: Tau <sup>2</sup> = 0.00; Ch	$i^2 = 9.01$ , df = 9 (P =	0.44); l <sup>2</sup> =0%			
Test for overall effect: Z = 8.68	(P < 0.00001)				
Test for subgroup differences: C	$Chi^2 = 0.01, df = 1 (P = 1)$	= 0.93), l <sup>2</sup> =0.0%			

Favours flumazenil Favours placebo

# Analysis I.6. Comparison I Flumazenil versus placebo, Outcome 6 Hepatic encephalopathy and bias control.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: 6 Hepatic encephalopathy and bias control

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random, C
I Low risk of bias					
Barbaro 1998	199/265	253/262	•	74.9 %	0.78 [ 0.72, 0.84 ]
Subtotal (95% CI)	265	262	*	74.9 %	0.78 [ 0.72, 0.84 ]
Total events: 199 (Flumazenil), 2	253 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 6.75$	(P < 0.00001)				
2 High risk of bias					
Cadranel 1995	4/10	8/8		0.8 %	0.43 [ 0.21, 0.90 ]
Dursun 2003	3/4	6/6		1.1 %	0.75 [ 0.41, 1.39 ]
Dursun 2003	9/16	4/ 4	-+-	2.2 %	0.58 [ 0.38, 0.89 ]
Gyr 1996	21/28	21/21	-	8.1 %	0.76 [ 0.61, 0.95 ]
Lacetti 2000	6/28	12/26	<u> </u>	0.6 %	0.46 [ 0.20, 1.06 ]
Li 2009	22/39	31/33	+	4.8 %	0.60 [ 0.45, 0.80 ]
Pomier-Layrargues 1994	8/11	10/10		2.7 %	0.74 [ 0.50, 1.09 ]
Van der Rijt 1995	5/9	7/9		0.9 %	0.71 [ 0.36, 1.41 ]
Zhu 1998	10/13	12/12	-	3.9 %	0.78 [ 0.57, 1.08 ]
Subtotal (95% CI)	158	139	•	25.1 %	0.69 [ 0.61, 0.78 ]
Total events: 88 (Flumazenil), 12	21 (Placebo)				
Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup>	= 6.48, df = 8 (P = 0	.59); I <sup>2</sup> =0.0%			
Test for overall effect: Z = 5.77	(P < 0.00001)				
Total (95% CI)	423	401	•	100.0 %	0.75 [ 0.71, 0.80 ]
Total events: 287 (Flumazenil), 3	( )				
Heterogeneity: $Tau^2 = 0.00$ ; Ch		0.44); l <sup>2</sup> =0%			
Test for overall effect: $Z = 8.68$					
Test for subgroup differences: C	$2hi^2 = 2.69, df = 1 (P = 1)$	$= 0.10), 1^2 = 63\%$			

Favours flumazenil Favours placebo

## Analysis 1.7. Comparison I Flumazenil versus placebo, Outcome 7 Hepatic encephalopathy and trial design.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: 7 Hepatic encephalopathy and trial design

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
Cross-over					
Barbaro 1998	199/265	253/262	•	94.6 %	0.78 [ 0.72, 0.84 ]
Cadranel 1995	4/10	8/8		1.0 %	0.43 [ 0.21, 0.90 ]
Pomier-Layrargues 1994	8/11	10/10		3.4 %	0.74 [ 0.50, 1.09 ]
Van der Rijt 1995	5/9	7/9		1.1 %	0.71 [ 0.36, 1.41 ]
Subtotal (95% CI)	295	289	•	100.0 %	0.77 [ 0.72, 0.83 ]
Total events: 216 (Flumazenil), 2	78 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$ ; $Chi^2$	= 2.64, df = 3 (P = 0	0.45); I <sup>2</sup> =0.0%			
Test for overall effect: $Z = 7.17$	(P < 0.00001)	,			
2 Parallel-arm	(				
Dursun 2003	12/20	20/20	+	15.9 %	0.61 [ 0.43, 0.87 ]
Gyr 1996	21/28	21/21	-	37.6 %	0.76 [ 0.61, 0.95 ]
Lacetti 2000	6/28	12/26	<b>_</b>	3.2 %	0.46 [ 0.20, 1.06 ]
Li 2009	22/39	31/33	-	23.8 %	0.60 [ 0.45, 0.80 ]
Zhu 1998	10/13	12/12	-	19.6 %	0.78 [ 0.57, 1.08 ]
Subtotal (95% CI)	128	112	•	100.0 %	0.69 [ 0.59, 0.79 ]
Total events: 71 (Flumazenil), 96	(Placebo)				
Heterogeneity: $Tau^2 = 0.00$ ; Ch	i <sup>2</sup> = 4.28, df = 4 (P =	$0.37$ ); $ ^2 = 6\%$			
Test for overall effect: $Z = 5.02$					
Test for subgroup differences: C	· /	$= 0   6    ^2 = 49\%$			
iest for subgroup differences. C		0.10), 1 = 1770			

Favours flumazenil Favours placebo

# Analysis I.8. Comparison I Flumazenil versus placebo, Outcome 8 Hepatic encephalopathy and duration of follow-up.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

#### Comparison: I Flumazenil versus placebo

Outcome: 8 Hepatic encephalopathy and duration of follow-up

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
$  \leq  $ day					
Dursun 2003	12/20	20/20	-	21.6 %	0.61 [ 0.43, 0.87 ]
Gyr 1996	21/28	21/21	•	55.5 %	0.76 [ 0.61, 0.95 ]
Lacetti 2000	6/28	12/26	<b>.</b> _	4.1 %	0.46 [ 0.20, 1.06 ]
Pomier-Layrargues 1994	8/11	10/10		18.8 %	0.74 [ 0.50, 1.09 ]
Subtotal (95% CI)	87	77	•	100.0 %	0.71 [ 0.60, 0.83 ]
Total events: 47 (Flumazenil), 63	3 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$ ; Chi <sup>2</sup>	· /	$(42) \cdot 1^2 = 0.0\%$			
0 ,		J.HZ), I =0.076			
Test for overall effect: $Z = 4.08$	(P = 0.000044)				
2 > 1 day					
Barbaro 1998	199/265	253/262	•	56.8 %	0.78 [ 0.72, 0.84 ]
Cadranel 1995	4/10	8/8		3.9 %	0.43 [ 0.21, 0.90 ]
Li 2009	22/39	31/33	+	18.8 %	0.60 [ 0.45, 0.80 ]
Van der Rijt 1995	5/9	7/9		4.4 %	0.71 [ 0.36, 1.41 ]
Zhu 1998	10/13	12/12	-	16.1 %	0.78 [ 0.57, 1.08 ]
Subtotal (95% CI)	336	324	•	100.0 %	0.72 [ 0.62, 0.84 ]
Total events: 240 (Flumazenil), 3	311 (Placebo)				
	· · · ·				
Heterogeneity: $Tau^2 = 0.01$ ; Ch	$i^2 = 5.58$ , df = 4 (P =	0.23); l <sup>2</sup> =28%			
Heterogeneity: Tau <sup>2</sup> = 0.01; Ch Test for overall effect: $Z = 4.31$		0.23); l <sup>2</sup> =28%			
	(P = 0.000016)				

Favours flumazenil Favours placebo

# Analysis I.9. Comparison I Flumazenil versus placebo, Outcome 9 Hepatic encephalopathy and acute liver failure.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: 9 Hepatic encephalopathy and acute liver failure

Study or subgroup	Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
Cirrhosis					
Barbaro 1998	199/265	253/262	•	78.1 %	0.78 [ 0.72, 0.84 ]
Cadranel 1995	4/10	8/8		0.9 %	0.43 [ 0.21, 0.90 ]
Dursun 2003	12/20	20/20		3.6 %	0.61 [ 0.43, 0.87 ]
Gyr 1996	21/28	21/21	-	9.1 %	0.76 [ 0.61, 0.95 ]
Lacetti 2000	6/28	12/26		0.7 %	0.46 [ 0.20, 1.06 ]
Pomier-Layrargues 1994	8/11	10/10		3.1 %	0.74 [ 0.50, 1.09 ]
Zhu 1998	10/13	12/12	-	4.5 %	0.78 [ 0.57, 1.08 ]
Subtotal (95% CI)	375	359	•	100.0 %	0.76 [ 0.71, 0.82 ]
Total events: 260 (Flumazenil), 3	336 (Placebo)				
Heterogeneity: Tau <sup>2</sup> = 0.00; Ch	$i^2 = 6.05$ , df = 6 (P =	0.42); I <sup>2</sup> = I %			
Test for overall effect: Z = 7.83	(P < 0.00001)				
2 Acute liver failure or cirrhosis					
Li 2009	22/39	31/33	-	84.7 %	0.60 [ 0.45, 0.80 ]
Van der Rijt 1995	5/9	7/9		15.3 %	0.71 [ 0.36, 1.41 ]
Subtotal (95% CI)	48	42	•	100.0 %	0.62 [ 0.47, 0.80 ]
Total events: 27 (Flumazenil), 38	3 (Placebo)				
Heterogeneity: Tau <sup>2</sup> = 0.0; Chi <sup>2</sup>	P = 0.21, df = 1 (P = 0.21)	0.64); l <sup>2</sup> =0.0%			
Test for overall effect: Z = 3.56	(P = 0.00037)				
Test for subgroup differences: C	2.27, df = 1 (P =	= 0.13), 1 <sup>2</sup> =56%			

0.02 0.1 1 10 50

Favours flumazenil Favours placebo

#### Analysis 1.10. Comparison | Flumazenil versus placebo, Outcome 10 Number Connection Test.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: 10 Number Connection Test

Study or subgroup	Flumazenil	Pla	acebo			C	Me Differer	ean hce		Mean Difference
	Ν	Mean(SD)[seconds]	Ν	Mean(SD)[secor	nds]	IV,Ra	ndom	,95% CI		IV,Random,95% Cl
Dursun 2003	20	86.56 (47.37)	20	90.35 (44.05)						-3.79 [ -32.14, 24.56 ]
				Fa	-50 wours fl	-25 umazenil	0	25 Favours	50 placebo	

# Analysis I.II. Comparison I Flumazenil versus placebo, Outcome II All-cause mortality and acute liver failure.

Review: Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy

Comparison: I Flumazenil versus placebo

Outcome: II All-cause mortality and acute liver failure

Flumazenil	Placebo	Risk Ratio M-	Weight	Risk Ratio
n/N	n/N	H,Kandom,95% Cl		H,Random,95% Cl
13/265	17/262		39.0 %	0.76 [ 0.37, 1.53 ]
1/9	2/5		4.2 %	0.28 [ 0.03, 2.35 ]
0/20	0/20			Not estimable
0/5	0/5			Not estimable
4/28	5/21		13.6 %	0.60 [ 0.18, 1.97 ]
0/6	0/6			Not estimable
6/28	5/26		17.1 %	1.11 [ 0.39, 3.22 ]
0/11	0/10			Not estimable
		0.01 0.1 1 10 100		
	n/N 13/265 1/9 0/20 0/5 4/28 0/6 6/28	n/N         n/N           13/265         17/262           1/9         2/5           0/20         0/20           0/5         0/5           4/28         5/21           0/6         0/6           6/28         5/26	M- H,Random,95% Cl 13/265 17/262 1/9 2/5 0/20 0/20 0/5 0/5 4/28 5/21 0/6 0/6 6/28 5/26 0/11 0/10	M- H,Random,95% n/N n/N Cl 13/265 17/262 42 % 0/20 0/20 42 % 0/20 0/20 42 % 0/5 0/5 4/28 5/21 13.6 % 0/6 0/6 13.6 % 0/6 0/6 17.1 %

Favours flumazenil Favours placebo

(Continued ...)

Study or subgroup	Flumazenil n/N	Placebo n/N	Risk Ratio M- H,Random,95% Cl	Weight	( Continued) Risk Ratio H,Random,95% Cl
Zhu 1998	3/13	5/12		13.4 %	0.55 [ 0.17, 1.83 ]
Subtotal (95% CI)	385	367	•	87.3 %	0.71 [ 0.45, 1.14 ]
Total events: 27 (Flumazenil), 34	1 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$ ; $Chi^2$	<sup>2</sup> = 1.71, df = 4 (P = 0	.79); I <sup>2</sup> =0.0%			
Test for overall effect: $Z = 1.40$	(P = 0.16)				
2 Participants with cirrhosis or	acute liver failure				
Li 2009	5/39	4/33		12.7 %	1.06 [ 0.31, 3.62 ]
Van der Rijt 1995	0/9	0/9			Not estimable
Subtotal (95% CI)	48	42	+	12.7 %	1.06 [ 0.31, 3.62 ]
Total events: 5 (Flumazenil), 4 (I	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.09$	(P = 0.93)				
Total (95% CI)	433	409	•	100.0 %	0.75 [ 0.48, 1.16 ]
Total events: 32 (Flumazenil), 38	3 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$ ; $Chi^2$	<sup>2</sup> = 2.05, df = 5 (P = 0	.84); l <sup>2</sup> =0.0%			
Test for overall effect: $Z = 1.28$	(P = 0.20)				
Test for subgroup differences: C	$Chi^2 = 0.34$ , $df = 1$ (P =	= 0.56), l <sup>2</sup> =0.0%			
			0.01 0.1 1 10 100		
			Favours flumazenil Favours placebo		

# ADDITIONAL TABLES

Table 1. Definition of improved manifestations of hepatic encephalopathy

Trial	Type of hepatic encephalopathy	Neuropsychiatric assessment	Definition of overall improve- ment
Amodio 1997	Minimal		Investigators did not define or as- sess the number of participants with an overall improvement
Barbaro 1998	Overt	Mental status assessed using a clinical scale (Table 2), Modified Glasgow Coma Scale (Table 2), and electroencephalography (Table 3).	Improvement in clinical scores or electroencephalography.
Cadranel 1995	Overt	Mental status assessed using a clinical scale (Table 2), and electroencephalography (Table 3).	Improvement in clinical score or electroencephalography.

Dursun 2003	Minimal or overt		ical score and Number Connec-
Giger-Mateeva 1999	Minimal	Number Connection Test and brainstem auditory evoked re- sponse	Not defined. The investigators in- cluded a post-hoc subjective as- sessment of alertness
Gooday 1995	Minimal	Simple and complex reaction time, verbal memory, psychomo- tor speed, short-term and work- ing memory	Improvement in psychomotor speed evaluated using change in reaction time; Investigators did not define or assess the number of participants with overall improve- ment.
Gyr 1996	Overt	Mental status (Table 2), and elec- troencephalography (Table 3).	Clinically relevant improvement defined as a 2-point improvement in clinical score at any time during treatment compared with base- line. The investigators also re- ported improvement defined us- ing the clinical scale score (mean for all individual observations)
Hermant 1991	Overt	Glasgow Coma Scale (Table 2), and electroencephalography.	A 2-point improvement in the Glasgow Coma Score and elec- troencephalography
Klotz 1989	Overt	Clinical assessment (score not de- scribed).	Improvement in clinical status
Lacetti 2000	Overt	Mental status (scale not specified) and Glasgow Coma Scale (Table 2).	Investigators originally classified participants as Grade III to IV coma. Method of assessment not stipulated. The trial report de- fined 'clinically relevant improve- ment' as primary outcome de- fined as a 3-point improvement in Glasgow Coma Score
Li 2009	Overt	Glasgow Coma Scale (Table 2), and electroencephalography	Improvement in the Glasgow Coma Score of $\geq 3$ points.
Pomier-Layrargues 1994	Overt	Modified Glasgow Coma Score (Table 2), and electroencephalog-raphy.	Improvement in $\geq 2$ items on modified Glasgow Coma Score within 1 hour after the end of treatment

## Table 1. Definition of improved manifestations of hepatic encephalopathy (Continued)

#### Table 1. Definition of improved manifestations of hepatic encephalopathy (Continued)

Van der Rijt 1995	Overt	Clinical scale (Table 2).	$A \ge 1$ -point decrease in severity of hepatic encephalopathy.
Zhu 1998	Overt	Clinical scale (Table 2).	Overall improvement in hepatic encephalopathy based on clinical grade

RCT: randomised clinical trial.

### Table 2. Neuropsychiatric assessment scales

Scale (Grippon 1988) used in Cadra	nel 1995; Barbaro 1998.
Ι	Euphoria or depression, mild confusion, slowness, disorder in sleep rhythm
II	Drowsiness, inappropriate behaviour, accentuation of stage I
III	Stupor; participant sleeps most of the time, but is rousable; inco- herent speech; marked confusion
IVa	Coma, co-ordinated response to painful stimuli.
IVb	Coma, hyperextension, and pronosupination after painful stimuli
IVc	Coma, no response to painful stimuli.
V	Clinical decerebration.
Scale (Fitz 1998) used in Dursun 20	03.
Subclinical	Normal examination with subtle changes in psychometric or Number Connection Tests
Ι	Impaired attention, irritability, depression, or personality changes
II	Drowsiness, behavioural changes, sleep disorders, and poor memory
III	Confusion, disorientation, somnolence, and amnesia.
Scale (Jones 1988) used in Gyr 1996	
-	Clinical assessment criteria consisted of the anamnestic criterion: disorders of sleep pattern (insomnia, hypersomnia, inversion of sleep rhythm) in combination with assessment of the level of con-

Scale (Grippon 1988) used in Cadranel 1995; Barbaro 1998.

	sciousness (1 to 4 as described below). Score items weighted so major disturbances of consciousness (portal systemic encephalopa- thy stage III and IV) were associated with scores of $\geq$ 11. Portal systemic encephalopathy stage II defined as scores of 5 to 10 and stage I of 3 to 4
1	Light disturbance of consciousness if $\geq 1$ of following symptoms were present: drowsiness (tendency to fall asleep but wake up spon- taneously or in response to normal voice or light), intermittent or permanent disorientation, retardation of ability to perform mental tasks (serial subtractions of sevens), mood disorder, inappropriate behaviour
2	Somnolence (arousable to physical stimuli such as mild prodding or shaking only)
3	Stupor (localised motor response to pain).
4	Coma (unarousability, no or unlocalised motor reactions to painful stimuli)
Scale (no reference provided in paper) used in Van der R	lijt 1995.
1	Presence of $\geq 2$ of following abnormalities: inverted sleep pattern, disturbed memory, impaired calculation (serial sevens), slowness of speech, or flapping tremor
2	Presence of $\geq 2$ of following: lethargy, time disorientation, or flapping tremor
3	Presence of $\geq 2$ of following: a state in which person had to be stimulated repetitively to open his/her eyes or execute commands, disorientation in terms of place and disorientation with respect to person
4	Coma.
Scale (Conn 1977) used in Zhu 1998.	
1	Trivial lack of awareness, euphoria or anxiety, shortened attention span, impaired performance of addition or subtraction
2	Lethargy or apathy, minimal disorientation for time or place, sub- tle personality change, inappropriate behaviour
3	Somnolence to semistupor, but responsive to verbal stimuli; con- fusion; gross disorientation
4	Coma.

Table 2.	Neuropsychiatric assessment scales	(Continued)
----------	------------------------------------	-------------

Scores	Eye opening (E):
	• $4 =$ spontaneous;
	• 3 = to voice;
	• 2 = to pain;
	• 1 = none.
	Verbal response (V):
	• 5 = normal conversation;
	• 4 = disoriented conversation;
	• 3 = words, but not coherent;
	• 2 = no words, only sounds;
	• 1 = none.
	Motor response (M):
	• 6 = normal;
	• 5 = localised to pain;
	• 4 = withdraws to pain;
	• 3 = decorticate posture (an abnormal posture that can
	include rigidity, clenched fists, legs held straight out, and arms
	bent inwards towards the body with wrists and fingers bend and
	held on chest);
	• 2 = decerebrate (an abnormal posture that can include
	rigidity, arms and legs held straight out, toes pointed downward
	head and neck arched backwards);
	• 1 = none.
Grading	• Severe: GCS 3-8 (minimum score 3).
Grading	<ul> <li>Moderate: GCS 9-12.</li> </ul>
	• Mild: GCS 13-15.
Modified Glasgow Coma Scale (Papp	as 1983) used in Pomier-Layrargues 1994; Barbaro 1998.
Scores	1. Verbal ability;
	2. Eye-opening;
	3. Pupillary light reflex;
	4. Corneal reflex;
	5. Spontaneous eye movements;
	6. Oculocephalic reflex;
	7. Motor response; and

#### Table 3. Assessment of electroencephalography changes

Ι

Electroencephalography grading/Fischer classification (Nusinovici 1977 and Spehlman 1991) used in Hermant 1991; Pomier-Layrargues 1994; Cadranel 1995; Barbaro 1998.

8. Pattern of respiration.

Irregular background activity (theta and alpha).

Flumazenil versus placebo or no intervention for people with cirrhosis and hepatic encephalopathy (Review)

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#### Table 3. Assessment of electroencephalography changes (Continued)

II	Continuous theta activity, bursts of delta waves.			
III	Prevalent delta activity; polyphasic transients sharp and slow wave complexes			
IVa	Continuous delta activity; abundant sharp and slow wave com- plexes; electroencephalography reactivity present			
IVb	Slower activity (delta and some polyphasic transients); electroen- cephalography reactivity = 0			
IVc	Discontinuous activity with silent periods.			
V	Flat.			
Electroencephalography grading (Parsons-Smith 1957) used in	n Dursun 2003.			
A	Generalised suppression of alpha rhythm and its frequent replace- ment by faster potentials in all leads. The tracings in this grade are generally flat and featureless			
В	Alpha rhythm very unstable and disturbed by random waves at 5-7 per second over both hemispheres. Rhythms most often seen over temporal lobes. In many cases with underlying fast activity			
С	Alpha rhythm still seen, but disturbed over both hemispheres by medium-voltage 5-6 per second waves. These occur in runs, are not paroxysmal, and do not usually block to eye opening although blocking may occur. Rhythms are particularly well seen over tem- poral and frontal lobes			
D	5-6 per second rhythms seen in grade C are now constant in all areas and replace all other cortical activity recorded on electroen- cephalogram. Appearance of this abnormality in a patient present- ing with only slight neuropsychiatric symptoms is very striking			
E	5 to 6 per second rhythms replaced by frontally preponderant bi- lateral synchronous 2 per second rhythms, which spread back- wards over hemispheres. At times, 6 per second rhythms might reappear, but special features of records are occurrence of these diencephalic discharges			
Electroencephalography grading (Kennedy 1973) used in Gyr 1996.				
0	8 to 12 per second basic rhythm, mean dominant frequency > 8 per second, % theta < 20			

#### Table 3. Assessment of electroencephalography changes (Continued)

1	Sudden shifts between normal alpha frequency (around 9 or 10 per second) and slow substitutes (6-8 per second); mean dominant frequency > 7 per second, % theta > 35
2	Diffuse slow activity posterior alpha rhythm seen occasionally, mean dominant frequency 5 to 7 per second, % theta > 60
3	Dominant slow activity in all areas, mean dominant frequency 3 to 5 per second, % delta 70
4	Bilaterally synchronous, 2-3 per second waves, predominating over frontal lobes and spreading backwards to occipital lobes; oc- casional short-lived appearance of faster rhythms (5 or 6 per sec- ond) or voltage depression, mean dominant frequency < 3 per second, % delta 70

Electroencephalography grading (Markand 1984) used in Van der Rijt 1995.

0	Background activity consisting of alpha rhythm.
1	Alpha rhythm with some scattered theta waves.
2	Background activity of theta activity intermixed with some delta and alpha frequencies
3	Background of delta polymorphic activity of high amplitude with spontaneous variability
4	Delta activity of relatively small amplitude.

## Table 4. Precipitating factors

Trial	Participants (n)	Precipitating factors (n)
Barbaro 1998	527	Gastrointestinal bleeding (352), surgery (95), sepsis (45), dehydration (6), unknown (29)
Cadranel 1995	14	Gastrointestinal bleeding (4), sepsis (7), alcoholic hepatitis (3), portal vein thrombosis (1), viral hepatitis (1), unknown (2)
Lacetti 2000	54	Gastrointestinal bleeding (31), sepsis (7), drugs (11), surgery (1)
Pomier-Layrargues 1994	21	Gastrointestinal bleeding (7), sepsis (2), dehydration (1), surgery (2), none (9), por- tacaval shunting (4)

#### Table 4. Precipitating factors (Continued)

Van der Rijt 1995	18	Hepatitis (5), acute exacerbation in cirrhosis (2), partial hepatectomy (1)
Zhu 1998	25	Gastrointestinal bleeding (13), protein overload (6), infection (2), wounds (1), un-known (3)

n: number of participants.

#### Table 5. Baseline screening for benzodiazepines in trial participants

Trial	Re- quired period free of benzodiazepines before inclusion	Baseline screening for ben- zodiazepines	Screening method and detection level	Negative testing at baseline an inclu- sion criterion	Proportion testing positive for benzo- diazepines at base- line
Amodio 1997	2 weeks	Yes	<ul> <li>Blood</li> <li>Emit-dau</li> <li>technique, Dupont</li> <li>Detection</li> <li>limit 0.3 μg/mL</li> <li>diazepam</li> </ul>	No	0%
Barbaro 1998	4 days	Yes	<ul> <li>Blood</li> <li>Thin-layer</li> <li>chromatography:</li> <li>Detection</li> <li>limit &gt; 11 mg/L</li> </ul>	No	1.9%
Cadranel 1995	Not reported	Yes	<ul> <li>Blood and urine</li> <li>Thin-layer chromatography:</li> <li>Detection limit &gt; 11 mg/L</li> </ul>	No	21.4%
Dursun 2003	3 days	No	Not reported	Not reported	Not reported
Giger-Mateeva 1999	3 months	Yes	<ul> <li>Urine</li> <li>Abbott TDx/ TDxFLx</li> <li>immunoassay</li> <li>Detection</li> <li>limit &lt; 200 ng/mL</li> </ul>	No	0%
Gooday 1995	1 month	No	Not reported	Not reported	Not reported

Gyr 1996	Yes, but length not specified	Yes	<ul> <li>Blood and urine</li> <li>Abbott TDx immunoassay</li> <li>Detection</li> <li>limits: blood 10 ng/ mL 100 ng/mL, urine &lt; 200 ng/mL</li> <li>Post-hoc analysis</li> <li>High-pressure</li> <li>liquid</li> <li>chromatography</li> <li>Detection</li> <li>limits: blood &lt; 50 ng/mL, urine 10 ng/mL to 100 ng/ mL</li> </ul>	No	8.2% on screening tests Flumazenil 11%; placebo 5% 12/49 samples for more sensitive test- ing lost
Hermant 1991	Not reported	Yes	Not reported	Yes	0%
Klotz 1989	Not reported	No	Not reported	Not reported	Not reported
Lacetti 2000	2 weeks	Yes	<ul> <li>Urine</li> <li>Roche KIMS</li> <li>immuno-enzymatic assay</li> <li>Detection</li> <li>limit: not specified</li> <li>Post-hoc analysis</li> <li>High-pressure</li> <li>liquid</li> <li>chromatography</li> <li>Detection</li> <li>limit: not specified</li> </ul>	Yes	0%
Li 2009	Not reported	No	Not reported	N/A	N/A
Pomier-Layrargues 1994	3 days	Yes	<ul> <li>Blood</li> <li>Abbott TDx</li> <li>immunoassay</li> <li>Detection</li> <li>limit 12 ng/mL</li> <li>Post-hoc analysis</li> <li>Gas</li> <li>chromatography-</li> <li>mass spectroscopy</li> <li>Detection</li> <li>limit: 1 ng/mL</li> </ul>	No	19%

## Table 5. Baseline screening for benzodiazepines in trial participants (Continued)

#### Table 5. Baseline screening for benzodiazepines in trial participants (Continued)

Van der Rijt 1995	Recent	Yes	<ul> <li>Blood</li> <li>High-pressure</li> <li>liquid</li> <li>chromatography</li> <li>Detection</li> <li>limit: not specified</li> </ul>	Yes	0%
Zhu 1998	7 days	No	Not reported	Not reported	Not reported

Table 6. Serious adverse events

Trial	Number of participants	Included in analyses of serious adverse events	Data included in pri- mary analysis	Serious adverse events
Amodio 1997	13	No	Cross-over RCT. Data from the first treatment period not described	Publication does not de- scribe any deaths or other seri- ous adverse events
Barbaro 1998	527	Yes	Cross-over RCT. Data from the first treatment period included	Thirteen non-re- sponders in the flumaze- nil group and 17 non re- sponders in the placebo group died 3 to 4 days (range 2-6) after ran- domisation. The causes of dead were septic shock (20 participants); hypo- volaemic shock (8 par- ticipants) and lactic aci- dosis (2 participants) but information was not pro- vided on the number of deaths by cause in each group
Cadranel 1995	14	Yes		One of 12 responders died from septic shock on day 4 and 2 of 6 non-responders died from septic shock (day 2) and lactic acidosis (day 4) but information is not provided on the groups to which they a were al- located

Dursun 2003	40	Yes	Parallel-arm RCT. We included all participants in the analyses	Pub- lication did not describe any deaths or other seri- ous adverse events
Giger-Mateeva 1999	10	No	Cross-over RCT. Data from the first treatment period not described	Pub- lication did not describe any deaths or other seri- ous adverse events
Gooday 1995	10	Yes	Cross-over RCT. Data from the first treatment period included	
Gyr 1996	49	Yes	Parallel-arm RCT. We included all participants in the analyses	Four of 28 participants allocated to flumazenil and 5 of 21 allocated to placebo died within 4 weeks of the trial. One participant in the placebo group died with respiratory failure dur- ing the course of the study. The authors de- scribed participants as having severe liver dis- ease suggesting that the cause of death in the re- maining 8 participants may have been cirrho- sis-related although this is not specifically stated. The investigators classi- fied the remaining ad- verse events viz flushing, nausea, vomiting, and ir- ritability, which were ex- perienced by 4 partici- pants, as non-serious
Hermant 1991	12	Yes	Parallel-arm RCT. We included all participants in the analyses	Pub- lication did not describe any deaths or other seri- ous adverse events
Klotz 1989	2	No	from the first treatment	Publication does not de- scribe any deaths or other seri-

#### Table 6. Serious adverse events (Continued)

			scribed	ous adverse events
Lacetti 2000	54	Yes	Parallel-arm RCT. We include all participants in the analyses	Six of 28 participants in the flumazenil group and 5 of 26 in the control group died. The causes of death were not pro- vided
Li 2009	72	Yes	Par- allel-arm RCT. The in- cluded participants had hepatic encephalopathy associated with cirrho- sis or acute liver fail- ure. Data were not pro- vide separately for the 2 groups	Five of 39 participants in the flumazenil group and 4 of 33 participants in the control group died. The causes of death were not provided
Pomier-Layrargues 1994	21	Yes	Cross-over RCT. Data from the first treatment period were included	Pub- lication did not describe any deaths or other seri- ous adverse events
Van der Rijt 1995	18	Yes	Cross-over RCT. The in- cluded participants had hepatic encephalopathy associated with cirrho- sis or acute liver fail- ure. Data were not pro- vide separately for the 2 groups.Data from the first treatment period were included	Pub- lication did not describe any deaths or other seri- ous adverse events. Two participants with fulmi- nant hepatic failure un- derwent orthotopic liver transplantation on day one of the study
Zhu 1998	25	Yes	Parallel-arm RCT. We included all participants in the analyses	Three of 13 participants in the flumazenil group and 5 of 12 participants in the control group died. The causes of death were not provided

RCT: randomised clinical trial.

# APPENDICES

# Appendix I. Search strategies

Database	Time span	Search strategy
Cochrane Hepato-Biliary Group Con- trolled Trials Register	May 2017.	(((benzodiazepine receptor OR GABA) AND (antagonist* OR blocking agent*)) OR flumazenil OR flumazepil)
Cochrane Central Register of Controlled Trials (CENTRAL) in the Cochrane Li- brary	2017, Issue 2.	<ul> <li>#1 MeSH descriptor: [GABA Antagonists] explode all trees</li> <li>#2 MeSH descriptor: [Flumazenil] explode all trees</li> <li>#3 ((benzodiazepine receptor or GABA) and (antagonist* or blocking agent*)) or flumaze*il</li> <li>#4 #1 or #2 or #3</li> <li>#5 MeSH descriptor: [Liver Cirrhosis] explode all trees</li> <li>#6 MeSH descriptor: [Hepatic Encephalopathy] explode all trees</li> <li>#7 liver cirrhosis or hepatic encephalopathy</li> <li>#8 #5 or #6 or #7</li> <li>#9 #4 and #8</li> </ul>
MEDLINE Ovid	1946 to May 2017.	<ol> <li>exp GABA Antagonists/</li> <li>exp Flumazenil/</li> <li>(((benzodiazepine receptor or GABA) and (antagonist* or blocking agent*)) or flumaze*il).mp. [mp=protocol supplementary concept, rare disease supplementary concept, title, original title, abstract, name of substance word, subject heading word, unique identifier]</li> <li>1 or 2 or 3</li> <li>exp Liver Cirrhosis/</li> <li>exp Hepatic Encephalopathy/</li> <li>(liver cirrhosis or hepatic encephalopathy).mp. [mp=protocol supplementary concept, rare disease supplementary concept, title, original title, abstract, name of substance word, subject heading word, unique identifier]</li> <li>5 or 6 or 7</li> <li>4 and 8</li> <li>(random* or blind* or placebo* or meta-analysis).mp. [mp=protocol supplementary concept, rare disease supplementary concept, title, original title, abstract, name of substance word, subject heading word, unique identifier]</li> </ol>
Embase Ovid	1974 to May 2017.	<ol> <li>exp benzodiazepine receptor blocking agent/</li> <li>exp 4 aminobutyric acid receptor blocking agent/</li> <li>exp FLUMAZENIL/</li> <li>(((benzodiazepine receptor or GABA) and (antagonist* or blocking agent*)) or flumaze*il).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer]</li> <li>1 or 2 or 3 or 4</li> </ol>

## (Continued)

		<ul> <li>6. exp liver cirrhosis/</li> <li>7. exp hepatic encephalopathy/</li> <li>8. (liver cirrhosis or hepatic encephalopathy).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer]</li> <li>9. 6 or 7 or 8</li> <li>10. 5 and 9</li> <li>11. (random* or blind* or placebo* or meta-analysis).mp. [mp=title, abstract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manufacturer]</li> <li>12. 10 and 11</li> </ul>
Science Citation Index Expanded (Web of Science)	1900 to May 2017.	<ul> <li>#5 #4 AND #3</li> <li>#4 TS=(random* or blind* or placebo* or meta-analysis)</li> <li>#3 #2 AND #1</li> <li>#2 TS=(liver cirrhosis or hepatic encephalopathy)</li> <li>#1 TS=(((benzodiazepine receptor or GABA) and (antagonist* or blocking agent*)) or flumaze*il)</li> </ul>
LILACS (Bireme)	1982 to May 2017.	(benzodiazepine receptor or GABA) and (antagonist\$ or block- ing agent\$)) or flumaze\$ [Words] and hepatic encephalopath\$ [Words]

# WHAT'S NEW

Last assessed as up-to-date: 5 May 2017.

Date	Event	Description	
4 May 2017	New search has been performed	Searches updated.	
14 October 2016	New search has been performed	The review methods and analyses are revised based on the recent recommendations of the Cochrane Hep- ato-Biliary Group, the MECIR guidelines, and the Cochrane Handbook for Reviews of Interventions	
14 October 2016	New search has been performed	Title change. Previously, the review was published with the title "Benzodiazepine receptor antagonists for hep- atic encephalopathy"	
9 May 2016	New citation required but conclusions have not changed	We included two additional randomised clinical trials (RCTs) in the analyses of benefits and harms and ex- cluded one RCT from the analyses of benefits. The lat- ter RCT and two additional observational studies are included in analyses of harms	

## CONTRIBUTIONS OF AUTHORS

LLG: drafted review and completed the statistical analyses.

LLG and MYM: validated the extracted data and refined the drafting of the review.

All authors participated in the selection of randomised clinical trials and extraction of data; interpretation of the results and in the critical revision of the review; and approved of the final version before submission.

Peer Reviewers: Manuel Romero-Gómez, Spain; R Todd Frederick, USA.

Contact Editor: Genaro D'Amico, Italy.

Sign-off Editor: Christian Gluud, Denmark.

# DECLARATIONS OF INTEREST

LLG: acted as investigator in studies funded by Norgine, Abbvie, Intercept, and Merck; received funding for travel expenses from Novo Nordisk; and received funding for lectures from Eli Lilly and Norgine.

MYM: no conflicts of interest.

ETG: no conflicts of interest.

MLA: no conflicts of interest.

## SOURCES OF SUPPORT

#### Internal sources

• Copenhagen Trial Unit, Denmark.

#### **External sources**

• No sources of support supplied

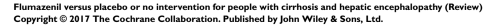
## DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The review has been extensively revised compared to the original protocol and the previously published version of this current review (Als-Nielsen 2004). The changes mainly reflect the current recommendations (Gluud 2017).

In the previous review (Als-Nielsen 2004), the primary outcomes included 'recovery' from hepatic encephalopathy defined as complete resolution of symptoms and 'improvement' of hepatic encephalopathy. We removed outcome 'recovery.' The term may be misleading as episodes may recur. Furthermore, people may show some degree of impairment between episodes (Bajaj 2010). Based on current guidelines (Gluud 2017), we assessed 'improvement' as number of participants 'without improvement of hepatic encephalopathy.' We now include all-cause mortality as a primary outcome rather than 'survival.'

The previous version of the review included data from both periods of cross-over trials. In this review, we only included data from the first treatment period because hepatic encephalopathy is fluctuating condition and because participants may die early in the randomised clinical trials.

We now report the results of meta-analyses using risk ratios, instead of risk differences, and include observational studies to improve our assessment of serious adverse events, Trial Sequential Analyses, and regression analysis (Harbord test) to evaluate the risk of smallstudy effects. The bias assessment is also updated.



## INDEX TERMS

#### Medical Subject Headings (MeSH)

Acute Disease; Antidotes [\*therapeutic use]; Chronic Disease; Flumazenil [\*therapeutic use]; GABA Modulators [\*therapeutic use]; GABA-A Receptor Antagonists; Hepatic Encephalopathy [\*drug therapy]; Randomized Controlled Trials as Topic

#### MeSH check words

Humans