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Effectiveness of brief alcohol interventions in primary care populations (Review)

Kaner EFS, Beyer FR, Muirhead C, Campbell F, Pienaar ED, Bertholet N, Daeppen JB, Saunders JB, Burnand B

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[Intervention Review]

Effectiveness of brief alcohol interventions in primary care populations

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ABSTRACT

Background

Excessive drinking is a significant cause of mortality, morbidity and social problems in many countries. Brief interventions aim to reduce alcohol consumption and related harm in hazardous and harmful drinkers who are not actively seeking help for alcohol problems. Interventions usually take the form of a conversation with a primary care provider and may include feedback on the person's alcohol use, information about potential harms and benefits of reducing intake, and advice on how to reduce consumption. Discussion informs the development of a personal plan to help reduce consumption. Brief interventions can also include behaviour change or motivationally-focused counselling.

This is an update of a Cochrane Review published in 2007.

Objectives

To assess the effectiveness of screening and brief alcohol intervention to reduce excessive alcohol consumption in hazardous or harmful drinkers in general practice or emergency care settings.

Search methods

We searched the Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE, and 12 other bibliographic databases to September 2017. We searched Alcohol and Alcohol Problems Science Database (to December 2003, after which the database was discontinued), trials registries, and websites. We carried out handsearching and checked reference lists of included studies and relevant reviews.

Selection criteria

We included randomised controlled trials (RCTs) of brief interventions to reduce hazardous or harmful alcohol consumption in people attending general practice, emergency care or other primary care settings for reasons other than alcohol treatment. The comparison

Effectiveness of brief alcohol interventions in primary care populations (Review)

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group was no or minimal intervention, where a measure of alcohol consumption was reported. 'Brief intervention' was defined as a conversation comprising five or fewer sessions of brief advice or brief lifestyle counselling and a total duration of less than 60 minutes. Any more was considered an extended intervention. Digital interventions were not included in this review.

Data collection and analysis

We used standard methodological procedures expected by Cochrane. We carried out subgroup analyses where possible to investigate the impact of factors such as gender, age, setting (general practice versus emergency care), treatment exposure and baseline consumption.

Main results

We included 69 studies that randomised a total of 33,642 participants. Of these, 42 studies were added for this update (24,057 participants). Most interventions were delivered in general practice (38 studies, 55%) or emergency care (27 studies, 39%) settings. Most studies (61 studies, 88%) compared brief intervention to minimal or no intervention. Extended interventions were compared with brief (4 studies, 6%), minimal or no intervention (7 studies, 10%). Few studies targeted particular age groups: adolescents or young adults (6 studies, 9%) and older adults (4 studies, 6%). Mean baseline alcohol consumption was 244 g/week (30.5 standard UK units) among the studies that reported these data. Main sources of bias were attrition and lack of provider or participant blinding. The primary meta-analysis included 34 studies (15,197 participants) and provided moderate-quality evidence that participants who received brief intervention consumed less alcohol than minimal or no intervention participants after one year (mean difference (MD) -20 g/week, 95% confidence interval (CI) -28 to -12). There was substantial heterogeneity among studies ($I^2 = 73\%$). A subgroup analysis by gender demonstrated that both men and women reduced alcohol consumption after receiving a brief intervention.

We found moderate-quality evidence that brief alcohol interventions have little impact on frequency of binges per week (MD -0.08, 95% CI -0.14 to -0.02; 15 studies, 6946 participants); drinking days per week (MD -0.13, 95% CI -0.23 to -0.04; 11 studies, 5469 participants); or drinking intensity (-0.2 g/drinking day, 95% CI -3.1 to 2.7; 10 studies, 3128 participants).

We found moderate-quality evidence of little difference in quantity of alcohol consumed when extended and no or minimal interventions were compared (-14 g/week, 95% CI -37 to 9; 6 studies, 1296 participants). There was little difference in binges per week (-0.08, 95% CI -0.28 to 0.12; 2 studies, 456 participants; moderate-quality evidence) or difference in days drinking per week (-0.45, 95% CI -0.81 to -0.09; 2 studies, 319 participants; moderate-quality evidence). Extended versus no or minimal intervention provided little impact on drinking intensity (9 g/drinking day, 95% CI -26 to 9; 1 study, 158 participants; low-quality evidence).

Extended intervention had no greater impact than brief intervention on alcohol consumption, although findings were imprecise (MD 2 g/week, 95% CI -42 to 45; 3 studies, 552 participants; low-quality evidence). Numbers of binges were not reported for this comparison, but one trial suggested a possible drop in days drinking per week (-0.5, 95% CI -1.2 to 0.2; 147 participants; low-quality evidence). Results from this trial also suggested very little impact on drinking intensity (-1.7 g/drinking day, 95% CI -18.9 to 15.5; 147 participants; very low-quality evidence).

Only five studies reported adverse effects (very low-quality evidence). No participants experienced any adverse effects in two studies; one study reported that the intervention increased binge drinking for women and two studies reported adverse events related to driving outcomes but concluded they were equivalent in both study arms.

Sources of funding were reported by 67 studies (87%). With two exceptions, studies were funded by government institutes, research bodies or charitable foundations. One study was partly funded by a pharmaceutical company and a brewers association, another by a company developing diagnostic testing equipment.

Authors' conclusions

We found moderate-quality evidence that brief interventions can reduce alcohol consumption in hazardous and harmful drinkers compared to minimal or no intervention. Longer counselling duration probably has little additional effect. Future studies should focus on identifying the components of interventions which are most closely associated with effectiveness.

PLAIN LANGUAGE SUMMARY

Effectiveness of brief alcohol interventions in primary care populations

What is the aim of this review?

Effectiveness of brief alcohol interventions in primary care populations (Review)
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We aimed to find out whether brief interventions with doctors and nurses in general practices or emergency care can reduce heavy drinking. We assessed the findings from 69 trials that involved a total of 33,642 participants; of these 34 studies (15,197 participants) provided data for the main analysis.

Key messages

Brief interventions in primary care settings aim to reduce heavy drinking compared to people who received usual care or brief written information. Longer interventions probably make little or no difference to heavy drinking compared to brief intervention.

What was studied in the review?

One way to reduce heavy drinking may be for doctors and nurses to provide brief advice or brief counselling to targeted people who consult general practitioners or other primary health care providers. People seeking primary healthcare are routinely asked about their drinking behaviour because alcohol use can affect many health conditions.

Brief interventions typically include feedback on alcohol use and health-related harms, identification of high risk situations for heavy drinking, simple advice about how to cut down drinking, strategies that can increase motivation to change drinking behaviour, and the development of a personal plan to reduce drinking. Brief interventions are designed to be delivered in regular consultations, which are often 5 to 15 minutes with doctors and around 20 to 30 minutes with nurses. Although short in duration, brief interventions can be delivered in one to five sessions. We did not include digital interventions in this review.

Search date

The evidence is current to September 2017.

Study funding

Funding sources were reported by 60 (87%) studies. Of these, 58 studies were funded by government institutes, research bodies or charitable foundations. One study was partly funded by a pharmaceutical company and a brewers association, another by a company developing diagnostic testing equipment. Nine studies did not report study funding sources.

What are the main results of the review?

We included 69 controlled trials conducted in many countries. Most studies were conducted in general practice and emergency care. Study participants received brief intervention or usual care or written information about alcohol (control group).

The amount of alcohol people drank each week was reported by 34 trials (15,197 participants) at one-year follow-up and showed that people who received the brief intervention drank less than control group participants (moderate-quality evidence). The reduction was around a pint of beer (475 mL) or a third of a bottle of wine (250 mL) less each week.

Longer counselling probably provided little additional benefit compared to brief intervention or no intervention.

One trial reported that the intervention adversely affected binge drinking for women, and two reported that no adverse effects resulted from receiving brief interventions. Most studies did not mention adverse effects.

Quality of the evidence

Findings may have been affected because participants and practitioners were often aware that brief interventions focused on alcohol. Furthermore, some participants could not be contacted at one-year follow-up to report drinking levels. Overall, evidence was assessed as mostly moderate-quality. This means the reported effect size and direction is likely to be close to the true effect of these interventions.

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

Brief intervention compared to no or minimal intervention for people with hazardous or harmful alcohol consumption						
<p>Patient or population: people with hazardous or harmful alcohol consumption as identified by a screening tool Setting: primary care (directly accessible to participant, no referral required), mostly high income countries Intervention: brief intervention Comparison: no or minimal intervention</p>						
Outcomes	Anticipated absolute effects* (95% CI)		Relative effect (95% CI)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Risk with control	Risk with Brief intervention				
Quantity of drinking (g/week) at 12 months	The mean quantity of drinking (g/week) at 12 months was 238 g/week	MD 20.08 g/week lower (28.36 lower to 11.81 lower)	-	15197 (34 RCTs)	⊕⊕⊕○ MODERATE ¹	Heterogeneity was substantial (73%) but not unexplained; interventions differed in content and delivery. The direction of effect favoured the intervention in 82% of the studies
Frequency of drinking (no. binges/wk) at 12 months	The mean frequency of drinking (no. binges/wk) at 12 months was 0.98 binges/week	MD 0.08 binges/week lower (0.14 lower to 0.02 lower)	-	6946 (15 RCTs)	⊕⊕⊕○ MODERATE ¹	
Frequency of drinking (no. days drinking/wk) at 12 months	The mean frequency of drinking (no. days drinking/wk) at 12 months was 2.73 drinking days/week	MD 0.13 drinking days/week lower (0.23 lower to 0.04 lower)	-	5469 (11 RCTs)	⊕⊕⊕○ MODERATE ¹	

Intensity of drinking (g/drinking day) at 12 months	The mean intensity of drinking (g/drinking day) at 12 months was MD 0.18 g/drinking day lower (3.09 lower to 2.73 higher)	-	3128 (10 RCTs)	⊕⊕⊕○ MODERATE ¹
Adverse effects	Only five trials reported adverse effects. No participants experienced any adverse effects in two trials; one trial reported that the intervention increased binge drinking for women; and two trials reported adverse events related to driving outcomes but concluded they were equivalent in both study arms	-	(5 RCTs)	⊕○○○ VERY LOW 12

* **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% CI).

CI: Confidence interval; RR: Risk ratio; OR: Odds 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

¹ High levels of performance bias due to difficulties with blinding participants and providers

² Downgraded due to inconsistency and imprecision; very few studies provided data, and reporting is inconsistent and data unavailable in a format conducive to meta-analysis

BACKGROUND

Description of the condition

Excessive drinking is a significant cause of mortality, morbidity and social problems in many countries, with a greater global cost to health than for tobacco (WHO 2014). The true impact of alcohol on the health of individuals and the wider community is difficult to estimate because of the many effects resulting from alcohol use, including increased levels of violence, accidents and suicide (GBD 2017). The heavy burden that alcohol use imposes on health, and its significant economic consequences, has led to national and international programmes and policies that seek to reduce consumption levels and so reduce a primary cause of avoidable ill health (Bailey 2011; UK Government 2012).

The impetus for a preventive approach to alcohol problems has been reinforced by epidemiological research. At a population level, most alcohol-related harm is not due to drinkers with severe alcohol dependence but attributable to a much larger group of excessive (hazardous and harmful drinkers) whose consumption exceeds recommended drinking levels (Anderson 1991). Therefore, at a population level, the greatest impact on alcohol-related problems can be made by addressing interventions aimed for excessive rather than dependent drinkers (McGovern 2013).

Description of the intervention

Early identification and secondary prevention of alcohol problems, using screening and brief interventions in primary care, has long been advocated as a strategy to reduce excessive drinking and is the focus of a great deal of research (O'Donnell 2014).

Brief intervention is grounded in social-cognitive theory and typically incorporates some or all of the following elements:

- feedback on the person's alcohol use and any alcohol-related harm clarification as to what constitutes low risk alcohol consumption;
- information on the harms associated with risky alcohol use;
- benefits of reducing intake;
- advice on how to reduce intake;
- motivational enhancement; analysis of high risk situations for drinking and coping strategies; and
- the development of a personal plan to reduce consumption.

Brief intervention is typically structured according to the FRAMES approach which involves practitioners: giving Feedback on the person's intake, impressing the Responsibility for change onto them, offering Advice, listing a Menu of options for behavioural change, having an Empathic approach and building Self-efficacy in the person receiving the brief intervention (Miller 1994).

Some brief intervention trials have included motivational interviewing (Rollnick 1995) or lifestyle counselling approaches. Although forms of brief intervention vary among studies (Heather 1995), core features in primary care are delivery by generalist healthcare workers, targets excessive (hazardous and harmful) drinkers who tend not to be seeking help for alcohol problems, and aims for reduced consumption and alcohol-related harms. Brief interventions in primary care have focused less frequently on dependent drinkers because these people often need more intensive treatment than is available routinely and are likely to require a goal of total abstinence.

How the intervention might work

Excessive drinking can be identified routinely in general practice and emergency care. People are often asked about alcohol consumption during new patient registrations, general health checks, specific disease clinics (e.g. hypertension, diabetes) and other health screening procedures. This identification process is often referred to as screening and typically involves asking a relatively small number of standardised questions about alcohol consumption (e.g. quantity, frequency and intensity of use) and any associated effects using a validated questionnaire or screening tool. Screening and brief alcohol intervention in routine primary care typically occurs opportunistically - when the main purpose of the appointment is something other than help with drinking.

The brief intervention must be delivered within the limited time frame of a standard consultation (typically 5 to 15 minutes for general practitioners (GPs), or up to 30 minutes for nurses). It also needs to fit in with routine practice (e.g. initial screening plus either referral to a practice colleague or later return for intervention). However, brief intervention trials have evaluated a wide range of activity. The shortest of these is a single 5 to 10 minute session of structured advice delivered by GPs or nurses. More intense interventions can provide multiple sessions of motivational interviewing or some other form of counselling, accompanied by repeated follow-up and delivered by primary healthcare workers. Other variations relate to the type of population being treated, the amount of training and support received by therapists, the theoretical basis underlying the intervention, and the use of accompanying written material.

Why it is important to do this review

Although previous reports and reviews have indicated beneficial outcomes of screening and brief intervention for excessive drinkers, crucial questions remain concerning its impact in routine practice and applicability to the broader population (Agosti 1995; Bien 1993; Moyer 2002; NHS CRD 1993; Poikolainen 1999; UK Government 2012; Wilk 1997). Whilst there appears to be little doubt that screening and brief intervention with excessive drinkers

can work successfully in research settings where intervention delivery and follow-up is carefully managed (Flay 1986), there has been uncertainty about extrapolation to real world routine primary care (Heather 2014; Holder 1999; Kaner 2001). However, if health professionals are to be encouraged to adopt and administer brief interventions in routine practice, it is necessary to establish a realistic effect size for brief intervention delivered in clinically-relevant contexts.

The term 'hazardous and harmful drinkers' contains a number of subgroups (e.g. young people, older people, and ethnic minorities). Little is known about how these subgroups respond to brief intervention in primary care. Differential loss of participants from early brief intervention trials led to a call for caution in generalising results to routine practice (Edwards 1997) but types of participants who were lost remains unclear. Therefore, there is need to characterise the types of drinkers for whom brief interventions have a positive impact and any subgroups who have not been represented in the trials to date.

This is an update of our 2007 review (Kaner 2007). An update was necessary because many trials have been conducted since initial publication and relevant developments have occurred in the wider literature; consequently we have added 42 new trials and conducted new subgroup analyses. Firstly, specific tools to measure efficacy and effectiveness have been published (Gartlehner 2006; Koppelaar 2011), which supersede the scale used in the 2007 review. Secondly, the concepts of screening reactivity and assessment reactivity have been defined (McCambridge 2011). These definitions describe how the very process of screening for alcohol consumption or having the effects of drinking assessed may influence reported drinking behaviour, independent of any further brief intervention input. Furthermore, recent literature contains a bigger subgroup of trials that focus on emergency care rather than general practice-based primary care, whereas previously the vast majority of the trials took place in primary care. Finally, interventions in more recent trials are often based on counselling techniques such as motivational interviewing rather than simpler advice-based input.

An important development has been in the digitisation of interventions to reduce hazardous and harmful alcohol consumption, using technologies such as websites and smart phone apps. These interventions (and their comparison with face-to-face brief interventions) are the focus of another Cochrane Review (Kaner 2017), and were excluded from this review.

OBJECTIVES

Main objective

To assess the effectiveness of screening and brief alcohol intervention to reduce excessive alcohol consumption in hazardous or harmful drinkers in general practice or emergency care settings.

Secondary objectives

Specific questions addressed by this review were:

1. Are brief interventions superior to minimal or no intervention?
2. Are extended brief interventions containing more or longer sessions superior to no intervention or to standard brief interventions?

METHODS

Criteria for considering studies for this review

Types of studies

Randomised controlled trials and cluster-randomised controlled trials were eligible for inclusion.

Types of participants

People who routinely presented to general practice, emergency care or other primary care settings for a range of health problems, whose alcohol consumption was identified by a screening tool as being excessive, or who had experienced harm as a result of their drinking behaviour. Studies that recruited participants who were seeking treatment specifically for an alcohol problem or who were mainly dependent on alcohol were excluded.

We defined primary care as all immediately accessible, general healthcare facilities. People needed to be able to access services on demand rather than through a specialist referral, and services needed to cover a broad range of problems. Participants recruited in emergency departments and trauma centres were included if this was the first contact following the emergency event.

Types of interventions

Experimental condition: Brief intervention comprised a single session and up to a maximum of five sessions of verbally-delivered information, advice or counselling that was designed to achieve a reduction in risky alcohol consumption, alcohol-related problems, or both (Babor 1994).

Control conditions: screening or assessment only, usual care for the presenting condition or written information such as a health or alcohol education leaflet (described as minimal intervention).

Psychology-based counselling aimed at reducing alcohol consumption or alcohol-related problems that was unlikely to occur in routine practice, for reasons of length or intensity, were referred to as extended intervention. We defined extended interventions as those that consisted of more than five sessions or total combined session durations was more than 60 minutes.

Interventions specifically aimed at people who were dependent on alcohol were excluded. Digital interventions (e.g. websites, smart phone apps or computer programmes) were excluded because these were investigated in another Cochrane Review (Kaner 2017).

Types of outcome measures

Primary outcomes

The primary outcome was consumption of alcohol. This was most often reported as:

1. Self-reported or other reports of drinking quantity (e.g. drinks per week).
2. Self-reported or other reports of binge drinking frequency (e.g. number of binges per week).
3. Self-reported or other reports of drinking frequency (e.g. drinking days per week).
4. Self-reported or other reports of drinking intensity (e.g. number of drinks per drinking day).
5. Self-reported or other reports of drinking within recommended limits (e.g. government recommended limits).

Although limits vary among countries, practitioners tend to use the national government recommended limits as a guide. Although not specified in the protocol, we also noted the following consumption outcomes if these were reported.

- Proportion of heavy drinkers (in Kaner 2007 a common example was > 35 units/week, but these definitions have reduced recently).
- Proportion of binge drinkers.

Secondary outcomes

1. Levels of laboratory markers of reduced alcohol consumption (e.g. serum gamma-glutamyltransferase (GGT), mean corpuscular volume (MCV)).
2. Alcohol-related harm to drinkers or others affected (e.g. via questionnaires such as the drinking problems index).
3. Patient satisfaction measures.
4. Health-related quality of life.
5. Economic measures including use of health services.

Search methods for identification of studies

Electronic searches

Current update searches

We searched the following databases from 2005 to September 2017:

- Cochrane Central Register of Controlled Trials (CENTRAL) (Issue 8, 2017; searched 25 September 2017);
- Cochrane Database of Systematic Reviews (CDSR) (Issue 9, 2017; searched 25 September 2017);
- Database of Abstracts of Reviews of Effects (DARE) (Issue 2, 2015; searched 25 September 2017);
- MEDLINE (Ovid, 1966 to September week 2, 2017; searched 21 September 2017);
- Embase (Ovid 1980 to 2017 week 38; searched 21 September 2017);
- PsycINFO (Ovid 1840 to September week 3 2017; searched 21 September 2017);
- CINAHL (EBSCO, 1982 to 25 September 2017);
- Science Citation Index Expanded (SCI-EXPANDED) (Web of Science, 2005 to 25 September 2017);
- Conference Proceedings Citation Index - Science (CPCI-S) (2005 to 25 September 2017);
- Social Sciences Citation Index (SSCI) (Web of Science, 2005 to 25 September 2017);
- Emerging Sources Citation Index (ESCI) (Web of Science, 2015 to 25 September 2017);
- NHS-EED (Wiley, issue 2 of 4, 2015; searched 25 September 2017).

Search strategies are reported in [Appendix 1](#); [Appendix 2](#); [Appendix 3](#); [Appendix 4](#); [Appendix 5](#); [Appendix 6](#); [Appendix 7](#); [Appendix 8](#); and [Appendix 9](#).

We applied no language or publication restrictions.

Previous searches

We searched the following sources from the earliest available date to 2006:

- Cochrane Drug and Alcohol Group specialised register (February 2006);
- Cochrane Central Register of Controlled Trials (CENTRAL);
- MEDLINE (Ovid, 1966 to 2005);
- Embase (Ovid, 1980 to 2005);
- PsycINFO (Ovid, 1840 to 2005);
- CINAHL (EBSCO, 1982 to 2005);
- Social Sciences Citation Index (SSCI) (Web of Knowledge, 1970 to 2005);
- Science Citation Index (SCI) (Web of Knowledge, 1970 to 2005);
- Cochrane Effective Practice and Organisation of Care Group specialised register (2005);
- Alcohol Education and Research Council (AERC) alcohol library, searched 2005;

- HEED (searched 3 December 2014 - no access for the September 2017 search); and
- Alcohol and Alcohol Problems Science Database, ETOH (1972 to 2003, after which the database was discontinued).

Searching other resources

We handsearched the reference lists of included studies and relevant systematic reviews. We contacted key informants and experts to enquire about unpublished work and ongoing research, particularly through links with the International Network on Brief Interventions for Alcohol and other drugs (INEBRIA).

We also searched clinicaltrials.gov (25 September 2017).

We searched organisational websites for reports of eligible trials on 26 September 2017:

- USA Department of Health and Human Services, Substance Abuse and Mental Health Services Administration (SAMHSA), National Registry of Evidence Based Programs and Practices (NREPP);
- SAMHSA Screening Brief Intervention and Referral to Treatment (SBIRT);
- Information on Drugs and Alcohol (IDA);
- Alcohol Concern;
- Drug and Alcohol Findings;
- International Network on Brief Interventions for Alcohol and Other Drugs (INEBRIA); and
- National Treatment Agency for Substance Misuse.

Data collection and analysis

Selection of studies

Two review authors independently assessed titles and abstracts in EndNote (EndNote 2015). If the title, abstract and keywords did not yield enough information to ascertain potential for inclusion then the full paper was retrieved.

We initially piloted the inclusion criteria on six retrieved papers for the original review (Kaner 2007). Two review authors independently assessed the study eligibility for both the original review and this update. Any disagreement was resolved by discussion and consensus, or adjudication by a third review author.

Data extraction and management

Two review authors independently extracted data using a piloted data extraction form. We extracted citation information, participants' characteristics (e.g. age, gender, baseline alcohol consumption), intervention descriptions (e.g. number, content and frequency of brief intervention sessions), setting and outcome data to RevMan 5 (Review Manager 2014). We also extracted methodological information to enable critical appraisal. Where data were

missing or unclear we emailed study authors to request clarification or further data.

In the protocol, we specified the use of an intention-to-treat analysis as a criterion of quality. However, this was revised in light of current *Cochrane Handbook* guidance (Higgins 2011b). Intention-to-treat analysis is usually understood to mean that participants were analysed in the groups to which they were randomised, regardless of the treatment they actually received. However, it is also sometimes understood to imply that all participants were included regardless of whether their outcomes were actually collected, which requires imputation of missing outcomes. Rather than using intention-to-treat analysis as a quality criterion, we attempted to extract data for participants in the groups to which they were randomised, regardless of the treatment they actually received, i.e. corresponding to the more widely agreed definition of intention-to-treat analysis.

Outcome data on quantity of alcohol consumed in a specific time period were converted to grams per week for each study. Drinks and units were converted to grams using either a conversion factor reported in the relevant paper or, if none was reported, using the conversion factor appropriate for the country where the study was conducted (Furtwaengler 2013; Gual 1999; Heather 2006; Miller 1991). Months were converted to weeks by multiplying by 52/12. Drinking intensity, drinking days, drinking sessions and occasions were all assumed to be equivalent to drinking days. For laboratory markers of gamma-glutamyltransferase (GGT) (Israel 1996; Romelsjö 1989; Scott 1990; Wallace 1988), microkats/litre were converted to international units/litre (IU/L) by multiplying by 60. Where relevant, values from analyses that involved adjustment for missing data (e.g. through the imputation of baseline values for participants lacking follow-up data) were used in preference to unadjusted values.

Assessment of risk of bias in included studies

Two review authors independently assessed potential risk of bias resulting from the trial design according to Cochrane's 'Risk of bias' tool as described in the *Cochrane Handbook* (Higgins 2011). This is a two-part tool, addressing the following domains.

- Sequence generation and allocation concealment (selection bias).
- Blinding of participants and providers (performance bias).
- Blinding of outcome assessor (detection bias).
- Incomplete outcome data (attrition bias).
- Selective outcome reporting (reporting bias).
- Other source of bias.

The first part of the tool involves describing what was reported to have happened in the study. The second part of the tool involves assigning a judgement relating to the risk of bias in terms of low, high or unclear risk. We used the criteria indicated by the *Cochrane Handbook* adapted to the addiction field to make these judgments (see Appendix 10).

Any discrepancies between review authors were resolved by discussion to achieve consensus.

Measures of treatment effect

We calculated the mean difference (MD) and standard deviation (SD) between the value of the outcome measure at 12 months following the brief intervention and the corresponding value following the control intervention for each continuous outcome. If standard deviations of final values were not available, the change score (i.e. the difference between the final and initial value of the outcome measure) was used if its standard deviation was available. If no standard deviations were available, these trials were omitted from the primary analysis but included in a sensitivity analysis using imputed standard deviations.

The risk difference (RD) with 95% confidence intervals were calculated for dichotomous outcomes because 95% CIs are intuitively clearer and this was consistent with the approach applied previously (Kaner 2007).

Unit of analysis issues

We extracted a direct estimate of the desired treatment effect and its standard error where analyses accounted for clustering in cluster-randomised trials. We assigned imputed standard deviations to the treatment and control groups, such that the standard error of the treatment effect estimated by the weighted mean difference method in RevMan 5 (Review Manager 2014) was the same as that reported in the analysis accounting for clustering. If the analysis did not account for the cluster design, we extracted the number of clusters randomised to each intervention, the average cluster size in each intervention group, and the outcome data for all participants in each intervention group.

A design effect was estimated using an external estimate of the intra-cluster coefficient (ICC). In this way we inflated the variance of the effect estimate. In the case of dichotomous outcomes, this involved reducing the total number of participants and the number of participants with events, whilst keeping the proportion with events fixed. It was then possible to enter data to RevMan 5 (Review Manager 2014), and combine cluster-randomised trials with individually randomised trials in the same meta-analysis.

Dealing with missing data

We contacted study authors to obtain missing data and seek clarification where appropriate. Studies with missing standard deviations or for which the number of participants in each arm was not reported were excluded from the main analysis for the associated continuous measure. These studies were included in a sensitivity analysis, using imputed values for the standard deviations or the number of participants in each arm.

Assessment of heterogeneity

The magnitude of heterogeneity among trials was assessed using the I^2 statistic (Higgins 2002; Higgins 2003). The statistical significance of heterogeneity was assessed using P values derived from χ^2 tests (Deeks 2001).

Assessment of reporting biases

We assessed whether studies appeared to have incomplete reporting bias by noting in the risk of bias assessments whether the reported outcomes matched methods sections or any published protocols. We made every effort to minimise publication bias by searching a wide range of databases and sources of grey literature and not restricting by language or publication status. We constructed funnel plots (plots of the effect estimate from each study against the effect standard error) to assess potential for bias related to the size of the trials, which could indicate possible publication bias.

Data synthesis

The weighted mean difference method was used to estimate pooled effect sizes and 95% CI, if sufficient studies reporting the outcome were available. Most trials reported weekly or monthly alcohol consumption and few reported drinking frequency or intensity. Hence, the meta-analysis of alcohol quantity consumed per week provided most information and constituted the primary meta-analysis. Statistical analyses were performed using RevMan 5 (Review Manager 2014). Because the populations and interventions evaluated by the trials were so heterogeneous, it was deemed more appropriate to use a random-effects model for all analyses (DerSimonian 1986). Random-effects meta-regression modelling was conducted using the `metareg` command in Stata version 14.1 (Stata 2015). Meta-regression was used to assess any differences in calculated effect associated with the publication date of studies, baseline consumption of participants, duration of treatment, and efficacy/effectiveness score.

For dichotomous outcomes (participant classified as a heavy or binge drinker), RDs and 95% CIs were calculated and pooled in a random-effects meta-analysis using Mantel-Haenszel test weighting.

We addressed variable risk of bias using sensitivity analysis (see Sensitivity analysis).

If trials had more than one control arm and the various control arms were very similar (e.g. Sommers 2013), the results for these arms were combined by calculating weighted means of continuous outcomes and summing dichotomous outcomes; likewise for very similar treatment arms (D'Onofrio 2012).

'Summary of findings' tables

We used the GRADE approach to assess the quality of the evidence. GRADE takes into account issues related to both internal

and external validity, such as directness of results (Atkins 2004; Guyatt 2008; Guyatt 2011). The 'Summary of findings' tables present main review findings in a transparent and simple tabular format. In particular, 'Summary of findings' tables provide key information concerning the quality of evidence, the magnitude of effect of the interventions examined and the sum of available data on the main outcomes.

The GRADE system uses the following criteria for assigning grades of evidence.

- **High:** we are very confident that the true effect lies close to that of the estimate of the effect.
- **Moderate:** 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:** our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect.
- **Very low:** we have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of effect.

We used [GRADEpro GDT 2015](#) to import data from [Review Manager 2014](#) for the main outcomes of quantity of drinking (g/week), frequency of drinking (days per week and binges per week) and intensity of drinking (drinks/drinking day) for each of the comparisons (brief intervention versus minimal or no intervention, extended intervention versus minimal or no intervention, extended intervention versus brief intervention) ([Summary of findings for the main comparison](#); [Summary of findings 2](#); [Summary of findings 3](#)).

Subgroup analysis and investigation of heterogeneity

We carried out subgroup analyses to address the effects of each of the following:

Applicability issues - population characteristics, setting and mode of intervention

Where reported, we recorded the gender, age and ethnicity of included participants to assess how applicable brief interventions are to different groups of people presenting to primary care. We conducted subgroup analyses for the primary outcome (alcohol consumption at 12-months follow-up) based on gender and according to adolescents or young adults versus with other age groups. We also examined study settings (general practice or emergency care). We also investigated the modality of the intervention (whether it was reported as advice or counselling-based).

Variability in treatment exposure, control condition, year of publication, baseline consumption and follow up time scales

We calculated a measure of treatment exposure as the sum of the duration of the initial brief intervention plus the total duration of

all booster sessions, in minutes. If a range of durations was given then we used the mean. If duration was not reported, we assumed the brief intervention to take 5 to 10 minutes, with a mean of 7.5 minutes. We carried out a meta-regression analysis to look for any association between the impact of treatment on the quantity of alcohol consumed at 12 months and the level of treatment exposure. We performed this analysis separately for trials that compared a brief intervention with minimal or no intervention, and for trials that compared an extended intervention with minimal or no intervention. Only results from the comparison of extended intervention versus minimal or no intervention were included in meta-regression analysis for trials that included three intervention arms (minimal or no intervention, brief intervention and extended intervention).

The definition of excessive drinking has changed over time (e.g. [NHMRC 2009](#); [UK Department of Health 2016](#)), particularly regarding the threshold for entry into trials. To examine the impact of this and any other temporal factors, trials were classified by year of publication and meta-regression analysis performed to look for any relationship between publication date and the primary outcome measure. This enabled examination of whether the intervention effect had diminished as the definition of excessive drinking has reduced. A meta-regression analysis was also conducted on whether the primary outcome measure was related to the level of consumption at baseline. Because baseline consumption varied over time, meta-regression analysis informed examination of changes when adjustment was made for year of publication. We analysed studies added for this update separately from studies included in the previous review version ([Kaner 2007](#)). This analysis was planned to illustrate if primary outcome measure findings were compatible.

The primary analyses reported outcomes at 12 month follow-up, reflecting the large number of studies with information at this time point, and our interest in investigating robust changes in drinking behaviour rather than shorter term or transient changes. Where sufficient information was available (specifically, for quantity of alcohol consumed and frequency of drinking), analyses were undertaken based on other follow-up times. The minimum follow-up time was six months.

Effectiveness and efficacy

Efficacy trials tend to take place in tightly controlled research environments. They typically recruit a more homogenous group of participants than effectiveness trials. The former involve practitioners or interventionists who are likely to have more skills in alcohol intervention or behaviour change work than generalists working in routine primary care. They may occur in specialist healthcare or university settings and are often well-resourced, supported or closely monitored to ensure that interventions are delivered precisely as intended. Conversely, effectiveness trials are closer to a real world situation and are more representative of routine clinical practice. These tend to have a broader range of partici-

pants, involve clinicians who routinely work in primary care and allow more flexibility in the way the intervention is delivered. We developed a scale based on the work of [Shadish 2000](#) to categorise included trials along a spectrum of efficacy to effectiveness.

Two review authors independently classified each trial (see [Table 1](#)). If an item appeared to be partially clinically representative on any item, then we gave a midpoint score (either 1 or 0.5 as applicable). If the study authors did not report data relating to a particular item, then we allocated an intermediate score to limit bias in the trial toward the effectiveness or efficacy domain. We resolved disagreements concerning classification by discussion to achieve consensus.

We summed all items for each study to provide an efficacy/effectiveness score of 0 to 12. If a study scored highly it was likely to be highly clinically relevant and was considered to be an effectiveness trial with high external validity. Conversely, if a trial scored very low, it was highly research relevant and considered to be an efficacy trial with high internal validity. We plotted the effect of brief intervention compared to minimal or no intervention on the quantity of alcohol consumed, as estimated from random-effects meta-analysis, against the efficacy/effectiveness score. We performed meta-regression analysis to assess whether this treatment effect was related to the efficacy/efficiency score. We also categorised trials as effectiveness or efficacy trials based on score above or below the median and performed subgroup analysis.

Sensitivity analysis

Sensitivity analyses were performed based on the following characteristics.

1. Risk of bias: the primary meta-analysis was repeated in analyses:

- i) including only studies at low risk of bias due to allocation concealment; and

- ii) excluding studies at high risk of bias due to incomplete outcome data (i.e. attrition bias). We did not conduct a sensitivity analysis on the basis of risk of bias due to blinding because it is not possible to mask the nature of the intervention to providers or participants.

2. Missing standard deviations: we imputed the median standard deviation of the relevant outcome from other trials to both treatment and control groups of those with missing SDs.

3. Comparison of outcomes from cluster and individually randomised trials: a sensitivity analysis was performed to investigate the robustness of the conclusions, especially of the effect of varying assumptions about the magnitude of the ICC.

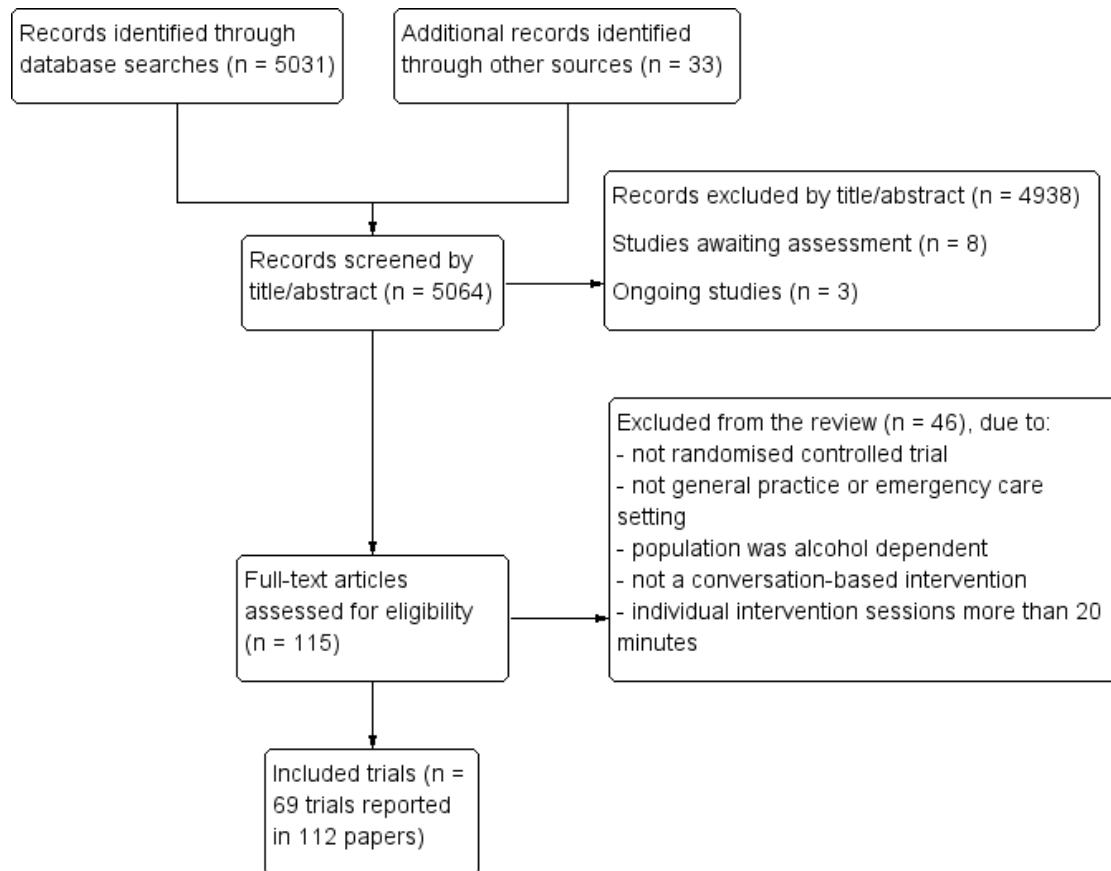
RESULTS

Description of studies

Results of the search

Searches identified 4004 potentially relevant records which were screened by title and abstract for eligibility. We retrieved 165 records for full text assessment. We added 42 studies (73 reports) for this update. The previous version of this review included 29 studies (39 reports) ([Kaner 2007](#)). This review included 69 studies (112 reports) ([Figure 1](#); [Characteristics of included studies](#)),

Figure 1. Study flow diagram.



We contacted 10 authors to request missing data or ask for clarification for this update.

Included studies

Population

The 69 included trials randomised a total of 33,642 participants (median 378, IQR 152 to 599). The mean percentage of male participants was 70%. Among the 45 trials that reported participants' age, the mean age was 40 years (SD 11.18). Eight trials focused on adolescents, young adults, or both (D'Amico 2008; Mertens 2014; Monti 2007; Segatto 2011; Spirito 2004; Spirito 2011; Tait 2004; Walton 2010). Four trials focused on older adults (aged > 55 years, aged > 60 years, or aged > 65 years) (Ettner 2014; Fleming 1999; Moore 2011; Watson 2013). Half the included trials (51%) reported participants' ethnicity; most participants were Caucasian (n = 28 trials, 78%).

In many cases, potential participants were excluded from trials if

they were heavily alcohol dependent, already on an alcohol treatment programme, or had been in the previous year. However, some trials did not specify any exclusion criteria and included a proportion of participants who may have been dependent drinkers. These trials were included where most participants were not identified as being dependent on alcohol and the intervention was not aimed at dependent drinkers.

All participants were screened for eligibility into trials. Screening methods included general health questionnaires (such as the Health and Habits Survey), which sometimes incorporated alcohol consumption questions, and established alcohol screening tools such as CAGE, AUDIT or MAST. Some trials used a combination of these tools and determined alternative inclusion criteria to increase the likelihood of picking up relevant participants. Most trials administered the screening tool by telephone or in a clinic as soon as the person had registered for their appointment; one study administered the questionnaire by telephone following the intervention. There were difference in alcohol consumption inclusion criteria among trials, for example by number of units per week,

screening tool score, level of binge, or high intensity drinking. The mean overall baseline consumption for participants in the 32 trials that reported these data was 244 g/week (SD 119) (about 30 UK units). We included 13 trials that reported baseline consumption for men only (or recruited men only) and also reported the number of men randomised (Aalto 2000; Altisent 1997; Babor 2006; Beich 2007; Córdoba 1998; Díez 2002; Fleming 1997; Huas 2002; McIntosh 1997; Richmond 1995; Rubio 2010; Scott 1990; Wallace 1988). In these trials, the mean baseline consumption was 350 g/week (around 44 UK units). The corresponding value for the nine trials that reported consumption for women was 190 g/week (around 24 UK units). Mean baseline consumption differed between older studies included in the original review and the more recent studies included for this update. Previously, the overall mean baseline consumption in 21 trials reporting these data was 313 g/week (about 39 UK units). Only 11 trials added for this update reported overall mean baseline consumption, which was 181 g/week (about 23 UK units).

We included 14 trials that reported baseline measures of frequency of drinking which could be converted to days drinking per week; the mean value was 2.07 days/week (Aalto 2000; Bernstein 2010; Cherpitel 2009; Daepfen 2007; Fleming 1997; Fleming 2010; Helstrom 2014; Monti 2007; Noknoy 2010; Rubio 2010; Schaus 2009; Senft 1997; Soderstrom 2007; Spirito 2004). Baseline intensity of drinking was reported in 11 trials, in which the mean baseline value was 69 g/drinking day.

There was substantial heterogeneity among trials in terms of the mechanisms of screening participants for inclusion and in the content of both control and intervention arms.

Setting

Most trials (n = 34) took place in the USA (Babor 2006; Bazargan-Hejazi 2005; Beckham 2007; Bernstein 2010; Blow 2006; Curry 2003; D'Amico 2008; Désy 2010; D'Onofrio 2008; D'Onofrio 2012; Ettner 2014; Field 2010; Fleming 1997; Fleming 1999; Fleming 2004; Fleming 2010; Helstrom 2014; Kunz 2004; Longabaugh 2001; Maisto 2001; McDevitt-Murphy 2014; Mello 2008; Monti 2007; Moore 2011; Ockene 1999; Schaus 2009; Senft 1997; Soderstrom 2007; Sommers 2006; Sommers 2013; Spirito 2004; Spirito 2011; Walton 2010; Woolard 2013); 10 in the UK (Crawford 2004; Crawford 2010; Drummond 2009; Drummond 2014; Heather 1987; Kaner 2013; Lock 2006; Scott 1990; Wallace 1988; Watson 2013); six in Spain (Altisent 1997; Córdoba 1998; Díez 2002; Fernández 1997; Rodríguez 2003; Rubio 2010); four in Australia (Dent 2008; Lane 2008; Richmond 1995; Tait 2004); two each in Canada (Israel 1996; McIntosh 1997), Finland (Aalto 2000; Seppa 1992) and Sweden (Romelsjö 1989; Tomson 1998); and one each in Denmark (Beich 2007), France (Huas 2002) Germany (Bischof 2008), Poland (Cherpitel 2009), Switzerland (Daepfen 2007), South Africa (Mertens 2014), Kenya (L'Engle 2014), Brazil (Segatto

2011) and Thailand (Noknoy 2010).

Many interventions (n = 38, 55%) were delivered in general practices (Aalto 2000; Altisent 1997; Babor 2006; Beckham 2007; Beich 2007; Bischof 2008; Córdoba 1998; Curry 2003; D'Amico 2008; Díez 2002; Drummond 2009; Ettner 2014; Fernández 1997; Fleming 1997; Fleming 1999; Fleming 2004; Heather 1987; Helstrom 2014; Huas 2002; Israel 1996; Kaner 2013; L'Engle 2014; Lock 2006; Maisto 2001; McIntosh 1997; Mertens 2014; Moore 2011; Noknoy 2010; Ockene 1999; Richmond 1995; Romelsjö 1989; Rubio 2010; Scott 1990; Senft 1997; Seppa 1992; Tomson 1998; Wallace 1988; Watson 2013) and 27 (39%) were carried out in emergency departments (Bazargan-Hejazi 2005; Bernstein 2010; Blow 2006; Cherpitel 2009; Crawford 2004; Crawford 2010; D'Onofrio 2008; D'Onofrio 2012; Daepfen 2007; Dent 2008; Désy 2010; Drummond 2014; Field 2010; Kunz 2004; Longabaugh 2001; Mello 2008; Monti 2007; Rodríguez 2003; Segatto 2011; Soderstrom 2007; Sommers 2006; Sommers 2013; Spirito 2004; Spirito 2011; Tait 2004; Walton 2010; Woolard 2013). Two studies took place in college health clinics (Fleming 2010; Schaus 2009), one in a public sexual health clinic (Lane 2008) and one in a veterans' affairs medical centre (McDevitt-Murphy 2014). The studies by Lane 2008 and McDevitt-Murphy 2014 were included because although they did not take place in general practice clinics, the interventions were available without referral to large groups of people. One study reported findings for two primary care settings and two other settings; only data from primary care settings were included in the meta-analyses (Díez 2002).

Interventions

Brief intervention

Most studies (n = 61) compared brief intervention with minimal or no intervention. Of these, five also included an extended intervention arm (Aalto 2000; Bischof 2008; Longabaugh 2001; Maisto 2001; Richmond 1995), and eight included two minimal or no intervention arms (Bernstein 2010; Blow 2006; Cherpitel 2009; Daepfen 2007; D'Onofrio 2012; Heather 1987; Richmond 1995; Sommers 2013). Two studies included two intervention arms with identical content but delivered by different health professionals (Babor 2006; McIntosh 1997). Three studies included two substantively different intervention arms (Dent 2008; D'Onofrio 2012; Walton 2010). Four studies delivered minimal intervention which was sometimes described as a control condition and sometimes as an intervention condition (Heather 1987; Kaner 2013; Richmond 1995; Sommers 2006). One study compared an extended intervention with a brief intervention (Spirito 2011). Four studies compared only an extended intervention with minimal or no intervention (Israel 1996; L'Engle 2014; Monti 2007; Moore 2011).

All interventions provided feedback on the screening outcome plus structured advice about potential risks of heavy drinking and ways to reduce consumption. Feedback and structured advice took several formats: described as brief Intervention (and assumed to be based on FRAMES where not reported) (n = 27); based on or informed by motivational interviewing, Motivational Enhancement Therapy (MET), or Brief Negotiated Interview (BNI) (n = 32); or based on Cognitive Behavioural Therapy (CBT) techniques (n = 2). Some were backed up by diaries or exercises for the participant to complete at home, and follow-up telephone calls.

A single brief intervention session was evaluated in 29 studies (Bazargan-Hejazi 2005; Beckham 2007; Blow 2006; Cherpitel 2009; Córdoba 1998; Crawford 2004; Crawford 2010; D'Onofrio 2008; Daepfen 2007; Désy 2010; Díez 2002; Drummond 2014; Fernández 1997; Field 2010; Kaner 2013; Kunz 2004; Lane 2008; Lock 2006; Longabaugh 2001; Maisto 2001; McDevitt-Murphy 2014; Mertens 2014; Ockene 1999; Rodríguez 2003; Scott 1990; Segatto 2011; Spirito 2004; Spirito 2011; Walton 2010). In the remaining studies, there were between two and five sessions, where individual sessions varied from one to a maximum of 60 minutes. General practitioners, nurse practitioners, health psychologists or trainee health psychologists administered the interventions.

Extended interventions

Extended interventions also provided feedback and structured advice but comprised either more than five sessions or more than 60 minutes in total. Extended interventions were based on motivational interviewing (five studies), MET (two studies), multiple FRAMES sessions (two studies), or CBT (one study) approaches. Extended interventions were evaluated in 10 trials (Aalto 2000; Bischof 2008; Israel 1996; L'Engle 2014; Longabaugh 2001; Maisto 2001; Monti 2007; Moore 2011; Richmond 1995; Spirito 2011), in which the total duration was greater than 60 minutes and the number of sessions delivered to participants ranged from two to seven. The total duration of extended intervention sessions ranged from 60 minutes to 180 minutes..

Total treatment exposure

Total treatment exposure was calculated as a combination of the initial session plus any additional sessions. Treatment duration in the intervention arm ranged from less than five minutes (Babor 2006; Huas 2002) to 60 minutes (McIntosh 1997) of advice or counselling. The median duration was 25 minutes and IQR 7.5 to 30.0 minutes. Treatment duration in the control group was up to 10 minutes (Díez 2002; Rodríguez 2003). In the extended intervention conditions, the treatment exposures ranged from 65 minutes to 175 minutes .

Control group content

Five categories of control condition were reported. Participants received:

1. screening only; or
2. screening and assessment only; or
3. usual care - this was usually not described further but was assumed to be care for the presenting condition or usual advice about alcohol consumption; or
4. general health advice or minimal advice about alcohol, comprising general health information or very limited alcohol-related information which often included an instruction to cut down drinking; or
5. a leaflet with either general health and lifestyle advice or more specific information about the risks of hazardous alcohol consumption.

Some trials provided control participants with both usual care and a leaflet. One trial did not include a control condition but compared extended intervention with brief motivational intervention (Spirito 2011).

Efficacy/effectiveness scores

Efficacy/effectiveness scores ranged from 4.5 (Beckham 2007; Fleming 2004; Romelsjö 1989) to 12 (Lock 2006). The median was 8.5 and IQR 7 to 10.5 (Table 2).

Reporting of outcomes

The included studies reported many different measures of the primary outcome. We could not include 22 studies in meta-analyses (Aalto 2000; Bazargan-Hejazi 2005; Beckham 2007; Crawford 2010; D'Amico 2008; Dent 2008; Désy 2010; Díez 2002; Drummond 2009; Heather 1987; Kunz 2004; L'Engle 2014; Lane 2008; McDevitt-Murphy 2014; Mello 2008; Mertens 2014; Noknoy 2010; Rodríguez 2003; Segatto 2011; Sommers 2006; Tait 2004; Woolard 2013), either because outcomes were not reported at 12 months, or outcome measures differed from those prespecified for this review (such as AUDIT score).

Quantity of alcohol consumed in a specified time period

Quantity of alcohol consumed in a specified time period (usually a week or a month) was reported in 51 studies that compared a brief intervention with minimal or no intervention. Of these, 27 studies reported quantity and corresponding standard deviation at 12 months (Altisent 1997; Beich 2007; Bernstein 2010; Blow 2006; Córdoba 1998; Crawford 2004; Daepfen 2007; D'Onofrio 2008; D'Onofrio 2012; Drummond 2014; Fleming 1997; Fleming 1999; Fleming 2004; Fleming 2010; Helstrom 2014; Kaner 2013; Lock 2006; Maisto 2001; Richmond 1995; Rubio 2010; Schaus 2009; Scott 1990; Senft 1997; Soderstrom 2007; Sommers 2013; Wallace 1988; Watson 2013). The authors of one study provided unpublished data on the corresponding standard deviation (Curry 2003). A further six studies reported the

change between baseline and the end of follow-up (change score) in the quantity of alcohol consumed in a specified time period and the corresponding standard deviation (Bischof 2008; Fernández 1997; Field 2010; Huas 2002; Ockene 1999; Romelsjö 1989). We included 34 studies in the primary meta-analysis comparing the effects of a brief intervention with minimal or no intervention on the quantity of alcohol consumed per week, reported at 12 months follow-up. Two studies reported the quantity of alcohol consumed per week both as assessed by structured interview and as reported on a self-completed questionnaire; we used the interview data (Scott 1990; Wallace 1988).

Five studies reported the final values of the quantity of alcohol consumed in a specified time period at 12 months but without corresponding standard deviations (Babor 2006; Cherpitel 2009; Ettner 2014; McIntosh 1997; Spirito 2004). These studies could not be included in the primary meta-analysis but were included in a sensitivity analysis with imputed standard deviations.

Six studies compared an extended intervention with minimal or no intervention and reported quantity of alcohol consumed at 12 months (Bischof 2008; Israel 1996; Maisto 2001; Monti 2007; Moore 2011; Richmond 1995). Three of these studies also compared an extended intervention to a brief intervention (Bischof 2008; Maisto 2001; Richmond 1995). One study, which compared an extended to a brief intervention, was included in a meta-analysis with imputed standard deviations (Spirito 2011).

The units of alcohol used in each trial with the conversion factor used to convert to grams of alcohol are presented in Table 2.

Frequency of drinking (number of drinking sessions in a specified time period)

We included 15 trials that compared brief intervention to no or minimal intervention and reported frequency of drinking in terms of number of binge drinking occasions each week or each month at six or 12 months (Blow 2006; Daepfen 2007; D'Onofrio 2008; D'Onofrio 2012; Fleming 1997; Fleming 1999; Fleming 2004; Fleming 2010; Helstrom 2014; Longabaugh 2001; Ockene 1999; Rubio 2010; Spirito 2004; Soderstrom 2007; Schaus 2009). Two trials reported this outcome for extended versus no or minimal intervention (Longabaugh 2001; Monti 2007).

Frequency of drinking in terms of number of days drinking each week or each month was reported by 11 trials at 6 or 12 months (Bernstein 2010; Cherpitel 2009; Crawford 2004; Curry 2003; Daepfen 2007; Field 2010; Fleming 2010; Helstrom 2014; Maisto 2001; Senft 1997; Spirito 2004). Longabaugh 2001 reported the number of days drinking but the number of participants assessed was not reported. Two trials reported this outcome for extended versus no or minimal intervention (Maisto 2001; Monti 2007), and one for extended versus brief intervention (Maisto 2001).

Intensity of drinking (amount of alcohol consumed in a drinking session)

We included 10 trials that compared brief intervention to no or minimal intervention and reported intensity of alcohol consumption in terms of number of drinks per occasion at 12 months (Bernstein 2010; Cherpitel 2009; Crawford 2004; Curry 2003; Daepfen 2007; Helstrom 2014; Maisto 2001; Schaus 2009; Senft 1997; Spirito 2004). One trial reported this outcome for extended intervention versus no or minimal intervention and versus brief intervention (Maisto 2001).

Laboratory markers

We included seven studies that compared brief intervention to no or minimal intervention and reported gamma-glutamyltransferase (GGT) (Aalto 2000; Beckham 2007; Noknoy 2010; Romelsjö 1989; Scott 1990; Tomson 1998; Wallace 1988). Romelsjö 1989 reported change scores whereas the other studies reported final values; only three studies reported at 12 months. One study reported this outcome for extended versus no or minimal intervention (Israel 1996).

One trial reported mean corpuscular volume (MCV) for this comparison (Seppa 1992).

Alcohol-related harm

A measure of alcohol-related consequences or harm (e.g. Drinker Inventory of Consequences - DrInC; Alcohol Problems Questionnaire - APQ) was reported by 21 studies (Blow 2006; Cherpitel 2009; D'Onofrio 2012; Drummond 2009; Drummond 2014; Fleming 2010; Heather 1987; Helstrom 2014; Kaner 2013; Lane 2008; Longabaugh 2001; McDevitt-Murphy 2014; McIntosh 1997; Mello 2008; Monti 2007; Romelsjö 1989; Schaus 2009; Spirito 2004; Walton 2010; Watson 2013; Woolard 2013).

Secondary outcomes

The percentage of heavy or binge drinkers at 12 months was reported by 19 studies (Altisent 1997; Beich 2007; Cherpitel 2009; Córdoba 1998; Curry 2003; Daepfen 2007; D'Onofrio 2008; Drummond 2014; Ettner 2014; Fernández 1997; Fleming 1997; Fleming 1999; Fleming 2004; Kaner 2013; Rubio 2010; Scott 1990; Sommers 2013; Wallace 1988; Walton 2010). Definitions of binge drinking varied, and were based on exceeding government recommendations, AUDIT scores, or a trial-defined number of drinks per week (for heavy drinking) or occasion (for binge drinking). Older studies tended to define higher limits (e.g. > 35 units/week by Wallace 1988) than more recent studies (e.g. > 18 drinks/week by Rubio 2010).

Two studies reported patient satisfaction measures (Drummond 2009; Kaner 2013), and six studies reported measures of health status or quality of life (Crawford 2004; Daepfen 2007; Drummond

2014; Kaner 2013; Lock 2006; Tait 2004). Two studies reported costs (Drummond 2009; Ettner 2014) and two studies reported economic evaluations (Lock 2006; Watson 2013). Nine studies reported measures of healthcare utilisation (Babor 2006; Crawford 2004; Dent 2008; Désy 2010; Ettner 2014; Fleming 1997; Fleming 2010; Kaner 2013; Tait 2004).

Reporting of outcomes by gender

Two studies reported on male and female participants in separate papers (Aalto 2000; Scott 1990). Seven studies reported on male and female participants separately in the same paper (Beich 2007; Richmond 1995; Rubio 2010; Scott 1990; Senft 1997; Seppa 1992; Wallace 1988). Data on men and women in the same trial were combined for the primary meta-analysis.

Two studies reported the final values of the quantity of alcohol consumed in a specified time period by gender but did not report the number of men and women assessed (Fleming 1997; Ockene 1999). Therefore, these studies could not be included in the meta-analysis of quantity of alcohol consumed in a specified time period, subgrouped by gender.

Outcomes reported at several follow-up time points

Many studies reported at several time points, although not all outcomes were reported at all time points. The TrEAT study reported after follow-up of one and four years (Fleming 1997). The HEALTH study reported at six-months and one-year follow-up (Ockene 1999). Richmond 1995 reported outcomes at six months and one year. Curry 2003 reported outcomes at three months and one year.

If outcomes were reported at several time points, data for one-year follow up were used in the meta-analyses if available.

Cluster-randomised trials

We included eight cluster-randomised trials (Babor 2006; Córdoba 1998; Drummond 2014; Ettner 2014; Huas 2002; Kaner 2013; Lock 2006; Ockene 1999). Córdoba 1998 and Lock 2006 randomised 33 and 40 general practices respectively with averages of 6.9 and 2.0 participants per practice. Huas 2002 randomised 88 primary care physicians with an average of 4.8 participants per physician, and Ettner 2014 recruited 31 primary care providers with a total of 1186 participants. Kaner 2013 recruited 34 general practices with a target of 31 participants per practice. Ockene 1999 randomised four internal medicine practice sites with an average of 111 participants per site. Babor 2006 recruited five managed care organisations, which provided 15 practices. Drummond 2014 recruited nine emergency departments with an average of 135 participants per site. Clustering was accounted for in analyses by five studies (Babor 2006; Drummond 2014; Kaner 2013; Lock 2006; Ockene 1999). For the three studies that did not account for clustering (Córdoba 1998; Ettner 2014; Huas 2002),

we extracted reported standard deviations for treatment and control groups and inflated them, assuming an ICC of 0.06 to allow for the clustered design (Higgins 2011a; Lock 2006).

Study funding sources

Most studies reported sources of funding (n = 60, 87%). With two exceptions, funding was provided by government institutes, research bodies or charitable foundations. One study was partly funded by a pharmaceutical company and a brewers association (Wallace 1988) and another by a company developing diagnostic testing equipment (Fleming 2004).

Excluded studies

We excluded 46 studies. The most common reason for exclusion was that participants were not recruited from primary care settings (n = 22). Other reasons included that the intervention was not conversation-based (n = 6), the intervention or comparator did not fit our prespecified criteria (n = 5) and the trial reported implementation rather than effectiveness outcomes (n = 2). We excluded two trials from the original review for this update. Chang 1997 compared brief intervention to alcohol treatment referral, which is not an analogous control group with the included trials. Participants in another trial (Gentilello 1999) were inpatients and the trial is included in a Cochrane Review of brief interventions in patients admitted to hospital.. See Excluded studies.

Studies awaiting classification

Searches identified eight studies which await classification (Arnaud 2017; Cherpitel 2016; Crawford 2015; Diaz Gomez 2017; Mello 2016; Monti 2015; Nadkarni 2017; Walton 2015; Characteristics of studies awaiting classification).

Mello 2016 and Monti 2015 recruited adult participants from emergency departments in the USA and administered brief motivational interventions. Monti 2015 reported improvement in the intervention group and Mello 2016 reported no clinically important difference between the groups at follow-up. Four studies recruited adolescents or young adults from emergency departments and administered brief interventions based on motivational interviewing or brief negotiation interviews (Arnaud 2017; Cherpitel 2016; Diaz Gomez 2017; Walton 2015). Two of these (Diaz Gomez 2017; Walton 2015) reported a significant improvement in the intervention group (although one (Walton 2015) noted it was a small effect) and two reported that consumption in both groups was reduced (Arnaud 2017; Cherpitel 2016).

Nadkarni 2017 recruited adult men in primary care centres in India and administered a multi-session (extended) intervention called Counselling for Alcohol Problems (CAP), reporting a significant improvement in the intervention group.

Crawford 2015 recruited adults from a primary care sexual health clinic in the UK and administered very brief advice followed by the

opportunity to see an alcohol health worker for a FRAMES-based intervention; no clinically important difference was seen between the groups.

These studies will be assessed for inclusion in a future update.

Ongoing studies

Three potentially eligible studies are ongoing. One is a UK trial of adolescents aged 14 to 17 who present to the emergency department and are randomised to a face-to-face intervention, a control arm, or a smart phone intervention (ISRCTN45300218). A second trial is recruiting young adults (aged 18 to 35 years) from a Swiss emergency department and administering brief motivational interviewing to the intervention group (ISRCTN13832949). The third is a trial taking place in adult women (aged over 18 years) presenting to Brazilian primary care clinics (RBR-65262c). Participants in the intervention group are to receive two brief inter-

vention sessions, each lasting 20 to 30 minutes.

These studies will be assessed for inclusion in a future update.

Risk of bias in included studies

The main sources of bias were attrition and performance bias. Performance bias occurred because it is not possible to blind participants or providers to a conversation-based intervention. Otherwise, very few trials were at high risk of bias. Many domains were assessed as unclear risk of bias because there was insufficient information reported in study reports to make a clear decision. This was largely because many included studies were published before CONSORT reporting guidelines appeared in 2010 (Schulz 2010).

Risk of bias and reasons for judgements are presented in [Characteristics of included studies](#) and graphic summaries in [Figure 2](#) and [Figure 3](#).

Figure 2. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.

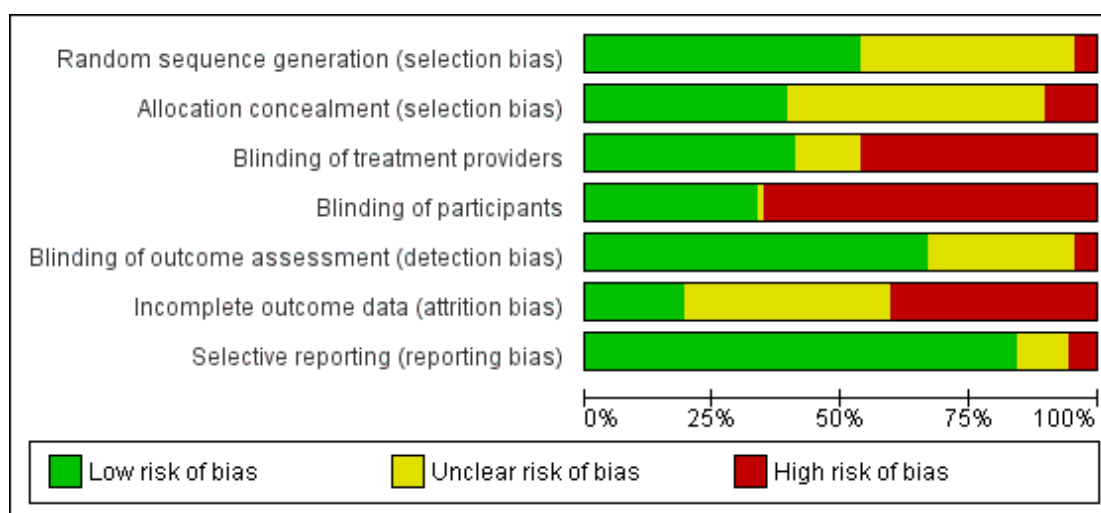


Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.



Allocation

Generation of randomisation sequence

The risk of selection bias due to randomisation methods was judged to be low in 37 trials (54%) (Aalto 2000; Altisent 1997; Babor 2006; Beckham 2007; Bernstein 2010; Bischof 2008; Blow 2006; Crawford 2004; Crawford 2010; Daepfen 2007; Dent 2008; Désy 2010; Drummond 2009; Drummond 2014; Ettner 2014; Fernández 1997; Field 2010; Fleming 1997; Fleming 2010; Israel 1996; Kaner 2013; L'Engle 2014; Lock 2006; Maisto 2001; Mello 2008; Monti 2007; Moore 2011; Noknoy 2010; Ockene 1999; Rubio 2010; Schaus 2009; Scott 1990; Segatto 2011; Sommers 2013; Tait 2004; Walton 2010; Watson 2013).

Risk of selection bias was assessed as high in three trials (4%) (Bazargan-Hejazi 2005; D'Amico 2008; Tomson 1998). Participants in these trials were assigned to treatment arms alternately or according to date of birth, or researchers changed the allocation process during the trial.

Randomisation methods were unclear in 29 trials.

Concealment of allocation

The risk of selection bias due to allocation concealment methods was judged to be low in 25 trials (36%) (Altisent 1997; Babor 2006; Beich 2007; Bernstein 2010; Bischof 2008; Córdoba 1998; Dent 2008; D'Onofrio 2008; Drummond 2014; Ettner 2014; Field 2010; Fleming 1997; Fleming 1999; Fleming 2010; Kaner 2013; L'Engle 2014; Lock 2006; Maisto 2001; Moore 2011; Ockene 1999; Romelsjö 1989; Scott 1990; Segatto 2011; Sommers 2013; Watson 2013).

Risk of bias was judged to be high in seven trials (10%) because participant allocation to arms was predictable, or those delivering the intervention opened a sealed envelope (no detail of numbering or other feature designed to protect allocation concealment) (Aalto 2000; Bazargan-Hejazi 2005; D'Amico 2008; Fernández 1997; Israel 1996; Longabaugh 2001; Rodríguez 2003).

Risk of allocation concealment bias was unclear in 37 trials.

Blinding

Due to the nature of the advice- or counselling-based interventions used in trials, it was not possible to blind either the participants or care providers. The automatic judgement of high risk of bias in this domain is considered by some as a limitation of the tool and, consequently, we judged trials to be at low risk of bias if they explicitly described a mechanism to overcome one of these biases. Detection bias was also affected by the fact that outcomes were self-reported by participants.

However, 28 trials (41%) were designed (either through cluster-randomisation or other design feature) so providers delivering the intervention had no contact with control participants, meaning there was no contamination between arms. These 28 studies were assigned low risk of bias for provider blinding (Babor 2006; Blow 2006; Cherpitel 2009; Córdoba 1998; Crawford 2004; Crawford 2010; Curry 2003; D'Onofrio 2012; Désy 2010; Drummond 2014; Ettner 2014; Fleming 1997; Fleming 1999; Fleming 2004; Fleming 2010; Helstrom 2014; Kaner 2013; Lock 2006; Moore 2011; Noknoy 2010; Ockene 1999; Schaus 2009; Scott 1990; Segatto 2011; Spirito 2011; Tomson 1998; Wallace 1988; Woolard 2013).

Researchers reported attempt to blind participants in 22 trials (32%) and were assigned low risk of bias for participant blinding at enrolment (Bazargan-Hejazi 2005; Curry 2003; D'Amico 2008; D'Onofrio 2008; D'Onofrio 2012; Daepfen 2007; Drummond 2009; Fleming 1997; Fleming 1999; Fleming 2010; Moore 2011; Noknoy 2010; Ockene 1999; Richmond 1995; Romelsjö 1989; Scott 1990; Segatto 2011; Spirito 2011; Tomson 1998; Wallace 1988; Walton 2010; Woolard 2013). Participants were blinded either because they did not know there was an alternate arm, or because the alcohol focus was masked during screening and assessment by other health-related questions.

Outcome assessors were reported to be blinded in 46 trials (67%) and were judged at low risk of detection bias (Babor 2006; Bazargan-Hejazi 2005; Bernstein 2010; Bischof 2008; Cherpitel 2009; Crawford 2004; Crawford 2010; D'Onofrio 2008; D'Onofrio 2012; Daepfen 2007; Drummond 2009; Drummond 2014; Ettner 2014; Fernández 1997; Field 2010; Fleming 1997; Fleming 1999; Fleming 2004; Fleming 2010; Heather 1987; Israel 1996; Kaner 2013; Kunz 2004; L'Engle 2014; Lane 2008; Lock 2006; Longabaugh 2001; Mello 2008; Mertens 2014; Monti 2007; Moore 2011; Noknoy 2010; Ockene 1999; Richmond 1995; Rodríguez 2003; Romelsjö 1989; Rubio 2010; Scott 1990; Senft 1997; Sommers 2006; Sommers 2013; Spirito 2004; Spirito 2011; Wallace 1988; Walton 2010; Woolard 2013).

We assessed three trials (4%) to be at high risk of detection bias because they stated that the outcome assessor was not blinded (Beich 2007; Dent 2008; Segatto 2011). We assessed 20 studies at unclear risk of bias for blinding.

Incomplete outcome data

We assessed 28 trials (40%) at high risk of attrition bias. Of these, 24 studies reported over 30% losses to follow-up (Aalto 2000; Altisent 1997; Bazargan-Hejazi 2005; Beich 2007; Cherpitel 2009; Córdoba 1998; Crawford 2004; D'Onofrio 2012; Dent 2008; Drummond 2014; Fernández 1997; Field 2010; Lock 2006; Richmond 1995; Rodríguez 2003; Schaus 2009; Scott 1990;

Seppa 1992; Soderstrom 2007; Sommers 2006; Sommers 2013; Spirito 2011; Tait 2004; Tomson 1998). Four studies reported completers' data only (Babor 2006; Blow 2006; Désy 2010; Kunz 2004).

The risk of attrition bias was judged to be unclear in 28 studies. Losses to follow-up were not reported according to study arm in five studies (Beckham 2007; Helstrom 2014; Israel 1996; Longabaugh 2001; McIntosh 1997). Reasons for losses to follow-up were not reported in 18 studies (Bernstein 2010; Curry 2003; Daepfen 2007; D'Onofrio 2008; Fleming 2004; L'Engle 2014; Lane 2008; Maisto 2001; McDevitt-Murphy 2014; Mello 2008; Mertens 2014; Monti 2007; Noknoy 2010; Romelsjö 1989; Senft 1997; Spirito 2004; Watson 2013; Woolard 2013). There were unexplained differences between losses to follow-up in the intervention and control arms in five studies (D'Amico 2008; Drummond 2009; Ertner 2014; Fleming 1997; Moore 2011).

Selective reporting

We assessed four trials (6%) at high risk of reporting bias because some aspect of reporting changed between protocol or methods section and reporting of results (Babor 2006; Bernstein 2010; Field 2010; Moore 2011). Seven were judged to be unclear and the rest low.

Effects of interventions

See: [Summary of findings for the main comparison](#) Brief intervention compared to no or minimal intervention for people with hazardous or harmful alcohol consumption; [Summary of findings 2](#) Extended intervention compared to no or minimal intervention for people with hazardous or harmful alcohol consumption; [Summary of findings 3](#) Extended intervention compared to brief intervention for people with hazardous or harmful alcohol consumption

All forest plots are presented with trials listed in chronological order.

I. Brief intervention versus minimal or no intervention

See [Summary of findings for the main comparison](#).

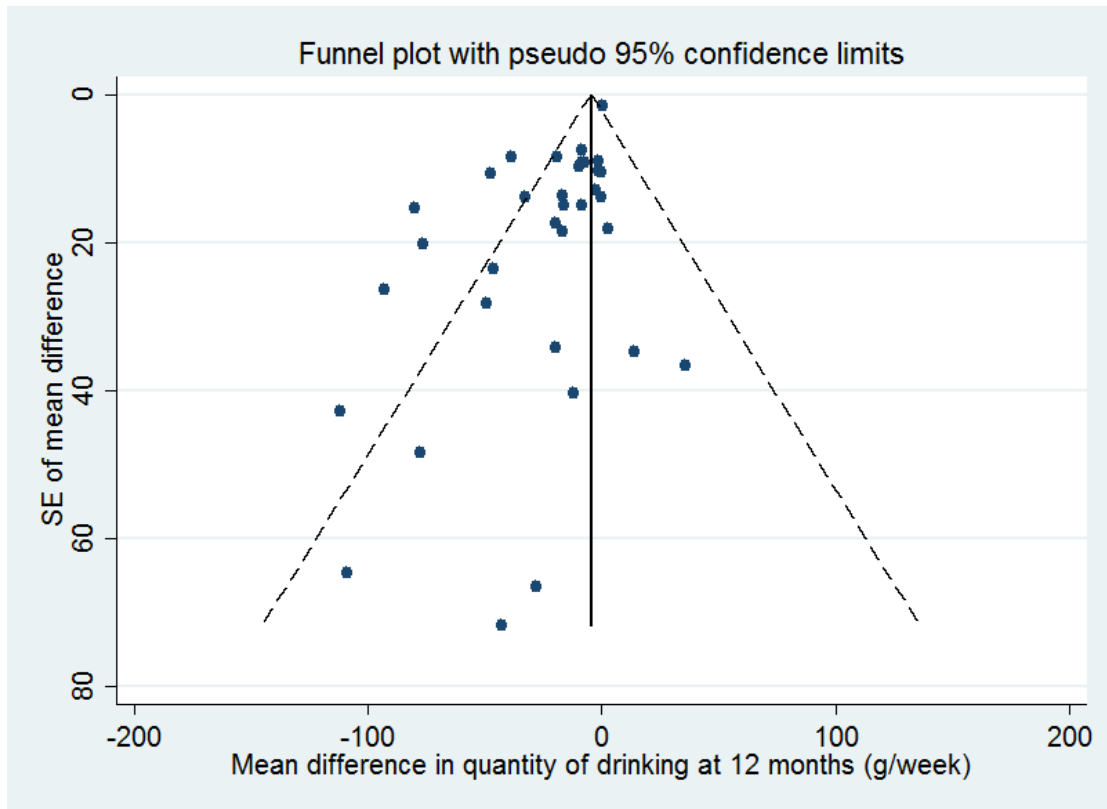
Quantity of alcohol consumed per week (Analyses I.1 to I.16)

Primary meta-analysis (Analysis 1.1)

The primary meta-analysis included 34 trials that reported consumption at 12 months ([Analysis 1.1](#)). A total of 15,197 participants were included in this analysis (median 413, range 64 to 1493), with a mean age of 43 years (SD 8.86). Seven studies in the primary meta-analysis included men only (Altisent 1997; Córdoba 1998; Crawford 2010; Díez 2002; Drummond 2009; Fernández 1997; Huas 2002); one included women only (McDevitt-Murphy 2014); and 26 trials recruited both men and women. Participants' ethnicity was reported in 19 trials (Bernstein 2010; Blow 2006; Córdoba 1998; Curry 2003; D'Onofrio 2008; D'Onofrio 2012; Drummond 2014; Field 2010; Fleming 1997; Fleming 2004; Fleming 2010; Helstrom 2014; Kaner 2013; Maisto 2001; Ockene 1999; Schaus 2009; Senft 1997; Soderstrom 2007; Sommers 2013). With the exception of four studies, most participants (68%) were white.

After 12 months, participants receiving a brief intervention drank a mean 20 g alcohol a week less than those with minimal or no intervention (95% CI 12 to 28; moderate-quality evidence; $I^2 = 73%$). The funnel plot ([Figure 4](#)) indicated asymmetry, suggesting the possibility of under-reporting of results with little or no evidence of an intervention effect.

Figure 4. Funnel plot for quantity of drinking at 12 months, with pseudo 95% confidence limits.



Sensitivity analyses to account for risk of bias and missing standard deviations (Analysis 1.2; Analysis 1.3; Analysis 1.4)

A sensitivity analysis restricted to 19 trials assessed at low risk of bias for allocation concealment showed very similar results to the primary meta-analysis (Analysis 1.2). Participants receiving brief intervention drank 20 g/week less than those with minimal or no intervention (95% CI 10 to 30). Heterogeneity was substantial ($I^2 = 74\%$). Of the 19 trials, 14 were also at low risk of bias for randomisation; five were assessed at unclear risk.

A sensitivity analysis that omitted the 15 trials at high risk of bias due to incomplete outcome data also showed very similar results to the primary meta-analysis (Analysis 1.3). Participants receiving brief intervention drank 21 g/week less than those with minimal or no intervention (95% CI 10 to 32). Heterogeneity was substantial ($I^2 = 80\%$).

A sensitivity analysis included five additional trials (compared to the primary meta-analysis) which did not report standard deviation. We imputed a median standard deviation of 167 g/week to both treatment and minimal or no intervention groups (Babor 2006; Cherpitel 2009; Ettner 2014; McIntosh 1997; Spirito 2004; Analysis 1.4). The addition of these trials made little difference;

pooling of all 39 trials showed that participants who received brief intervention drank 19 g/week less than those with minimal or no intervention (95% CI 12 to 27).

Impact of cluster randomisation (Analysis 1.6)

The six cluster-randomised trials (2588 participants) from the primary meta-analysis showed broadly similar results (MD -25 g/week, 95% CI -43 to -8) to the 28 individually randomised trials (12,609 participants) (MD -19 g/week, 95% CI -28 to -10; Córdoba 1998; Drummond 2014; Huas 2002; Kaner 2013; Lock 2006; Ockene 1999; Analysis 1.5). A sensitivity analysis showed that results from the cluster-randomised trials (MD -20 g/week, 95% CI -34 to -7) were very similar to the individually randomised trials (Analysis 1.6). We assumed an intra-cluster correlation coefficient (ICC) of 0.32 (as found in Lock 2006, unpublished data) for the two cluster-randomised trials that did not allow for cluster-randomisation in their analyses (Córdoba 1998; Huas 2002),

Impact of participant characteristics (Analysis 1.7; Analysis 1.8; Analysis 1.9)

Only 11 trials reported sufficient information (mean, standard deviation and number of participants assessed by treatment arm) by gender on the primary outcome measure at 12 months to permit meta-analysis subgrouped by men and women (Analysis 1.7). Both men (3486 participants) and women (1350 participants) appeared to experience significant benefits of brief intervention (MD for men -42 g/week, 95% CI -65 to -20; for women -30 g/week, 95% CI -59 to -2). The difference between men and women was not statistically significant.

We repeated Analysis 1.7, excluding four trials that recruited only men, to reduce confounding (Analysis 1.8). Results were similar: both men and women experienced significant benefits of brief intervention, with no significant difference in the treatment effect between men and women.

Three trials in the primary meta-analysis (1638 participants) recruited only adolescents or young adults (defined as maximum 21 years, 24 years and 25 years in Bernstein 2010; Fleming 2010; Schaus 2009, respectively). We separated these studies from the trials that did not impose age restrictions (13,559 participants). The treatment effect was smaller in the younger population (MD -7 g/week, 95% CI -17 to 3 compared to -23 g/week, 95% CI -32 to -13 for all adults; Analysis 1.9). However, all trials in younger people were relatively recent and meta-regression showed that, having adjusted for year of publication of the trial, there was no evidence that the intervention effect was lower in the trials of adolescents or young adults, compared to trials in adults ($P = 0.58$).

Impact of setting (Analysis 1.10)

We included 10 trials (6386 participants) in the primary meta-analysis that took place in an emergency department (Bernstein 2010; Blow 2006; Crawford 2004; D'Onofrio 2008; D'Onofrio 2012; Daepfen 2007; Drummond 2014; Field 2010; Soderstrom 2007; Sommers 2013). These studies showed less impact on alcohol consumption than trials based in general practice settings (8811 participants) (MD -10 g/week, 95% CI -18 to -2 versus -26 g/week, 95% CI -37 to -14; Analysis 1.10). There was no evidence

that the treatment effect differed between the sets of trials after adjusting for year of publication in meta-regression analysis.

Impact of intervention modality (Analysis 1.11)

A total of 20 trials tested advice-based interventions (8243 participants). These trials showed greater impact on alcohol consumption than the 12 counselling-based intervention trials (5537 participants, MD -33 g/week, 95% CI -46 to -20 versus -0.2 g/week, 95% CI -3 to 3; Analysis 1.11). Two trials were not included in this subgroup analysis because they contained arms that included both intervention types (Drummond 2014; Kaner 2013). There was no evidence that the treatment effect differed between the sets of trials after adjusting for year of publication in meta-regression analysis.

Impact of alcohol-related content in control condition (Analysis 1.12)

Half the trials in the primary meta-analysis provided some kind of alcohol-related advice or leaflet to control arm participants (Analysis 1.12). The mean difference in the 16 trials that provided alcohol-related content in the control arm (6591 participants) was lower (-13 g/week, 95% CI -23 to -3) than in 18 trials (8606 participants) where control group participants received no alcohol content (e.g. general health information, screening or assessment only) (-24 g/week, 95% CI -36 to -12). However, the test for subgroup differences was not significant.

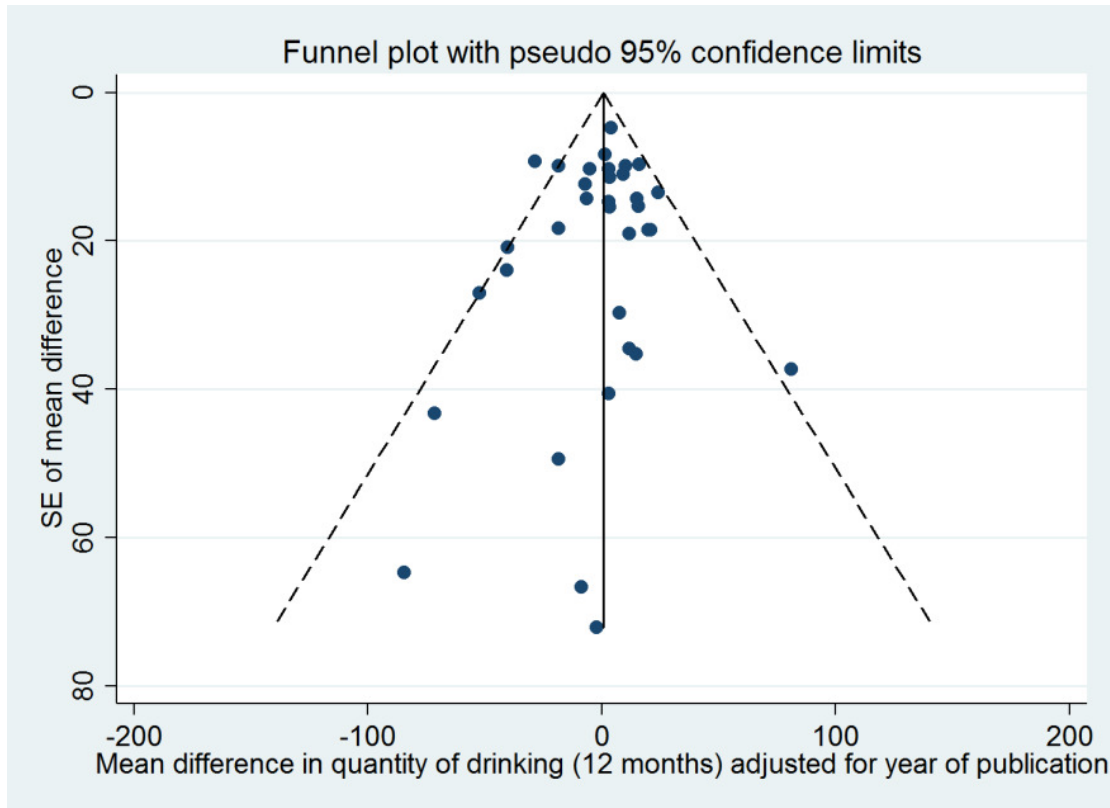
Impact of publication date and baseline consumption

Meta-regression showed that for every year going forward in time, the mean difference in consumption between brief intervention and minimal or no intervention groups decreased by 2.3 g/week (95% CI 1.3 to 3.4; Figure 5). There was a smaller difference in consumption between intervention and control participants in more recent trials. The residual heterogeneity ($I^2 = 42\%$) was notably lower than in the unadjusted analysis. Meta-regression adjustment of the funnel plot based on year of publication provided less evidence of asymmetry than the unadjusted funnel plot (Figure 4; Figure 6).

Figure 5. Meta-regression of quantity of drinking at 12 months on year of publication of trial.

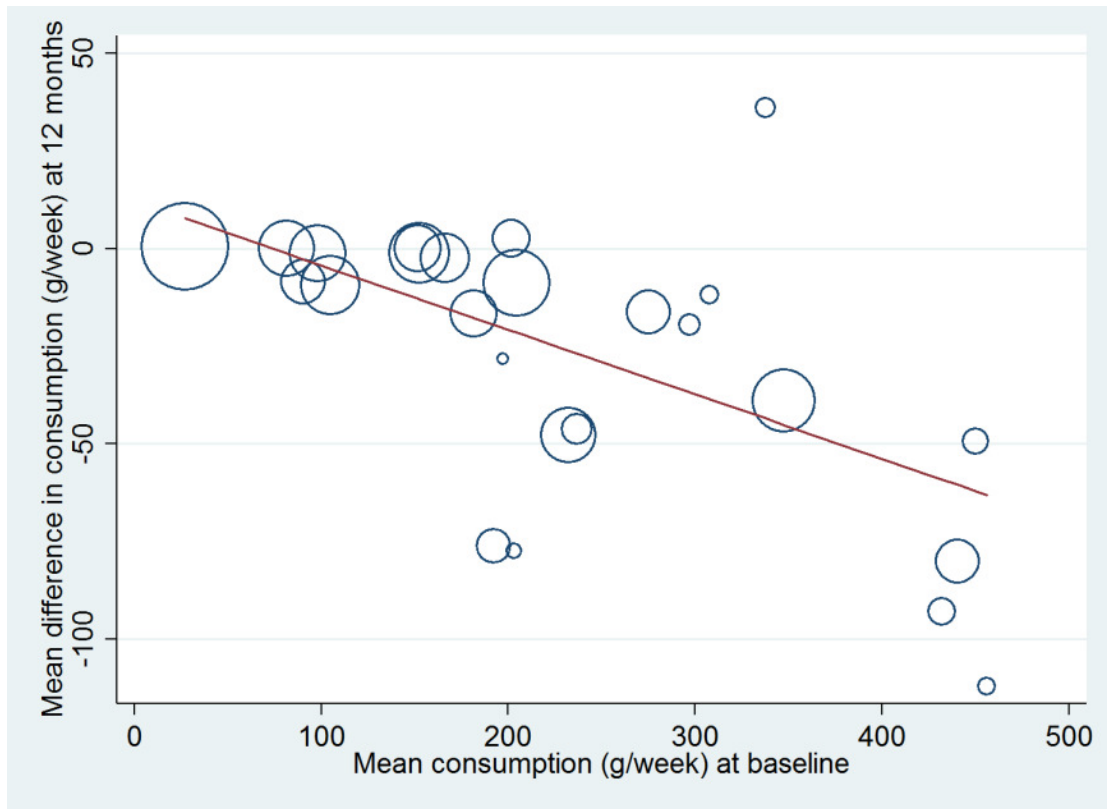


Figure 6. Funnel plot for quantity of drinking at 12 months, adjusted for year of publication, with pseudo 95% confidence limits.



We investigated the association of baseline consumption with effectiveness in a second meta-regression analysis (Figure 7). Results showed that for every g/week increase in baseline consumption, the mean difference in consumption between brief intervention and minimal or no intervention participants increased by 0.16 g/week (95% CI 0.10 to 0.23). There was a bigger difference in consumption between intervention and minimal or no intervention participants where baseline consumption was greater. The residual heterogeneity ($I^2 = 39\%$) was notably lower than in the unadjusted analysis. Including both baseline consumption and year of publication in the meta-regression further reduced residual heterogeneity ($I^2 = 29\%$). The mean difference between the brief intervention and minimal or no intervention arms decreased by 0.10 g/week (95% CI 0.03 to 0.18) for each increase of 1 g/week in baseline consumption, and increased by 1.5 g/week (95% CI 0.1 to 2.9) for each year increment in year of publication.

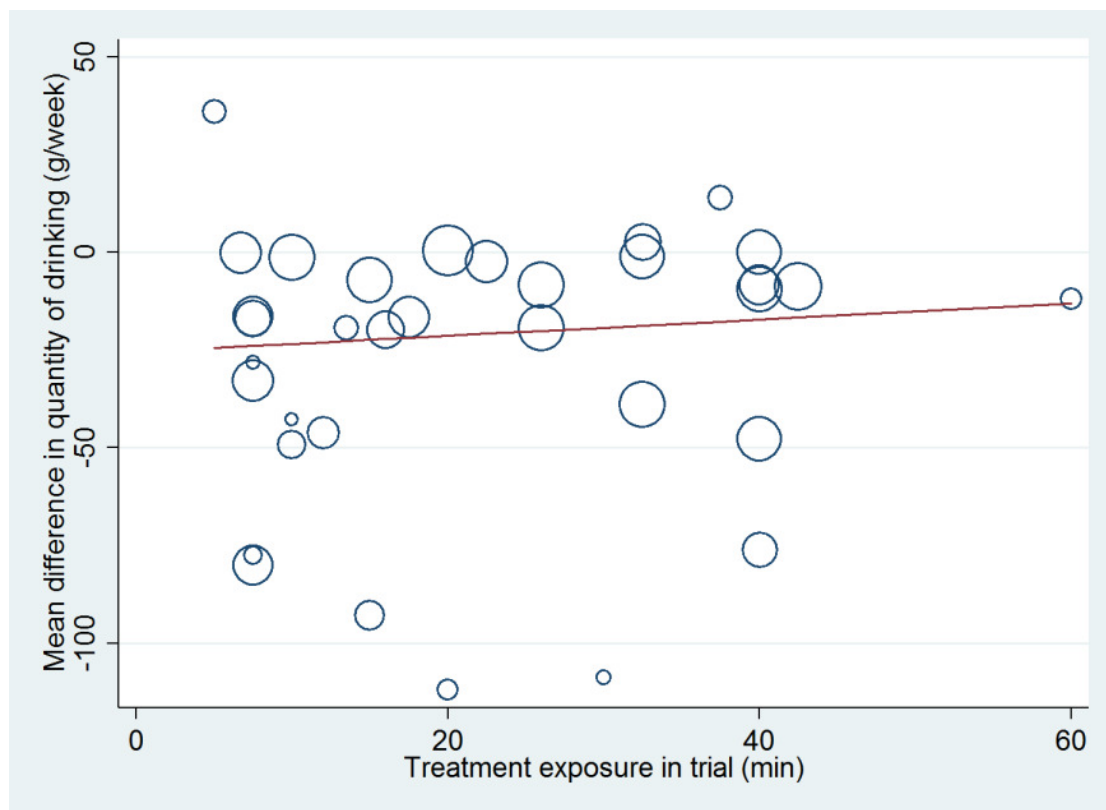
Figure 7. Meta-regression of quantity of drinking at 12 months on baseline consumption.



Treatment exposure

Meta-regression suggested little association between quantity of alcohol consumed at 12 months and increasing treatment exposure. Results showed an increase in the mean difference between the brief intervention and minimal or no intervention arms of 0.2 g/week (95% CI -0.5 to 0.9; $P = 0.57$, for each increase of 1 minute in the treatment exposure; [Figure 8](#)).

Figure 8. Meta-regression of quantity of drinking at 12 months on treatment exposure (mean duration of intervention for the participants in the trial), for trials comparing brief intervention with control.



Impact of follow-up time (Analysis 1.3; Analysis 1.14)

A sensitivity analysis by length of follow-up suggested no decay in impact of the intervention over the first 12 months. Only two trials reported data at eight to nine months, and only one at each 36 and 48 months. The pooled effect on consumption at six months (10,313 participants, MD -22 g/week, 95% CI -32 to -12) was similar to 12 months (15,197 participants, MD -20 g/week, 95% CI -28 to -12; Analysis 1.13). Results were similar when the analysis was restricted to studies that reported outcomes at both 6 months and 12 months (Analysis 1.14).

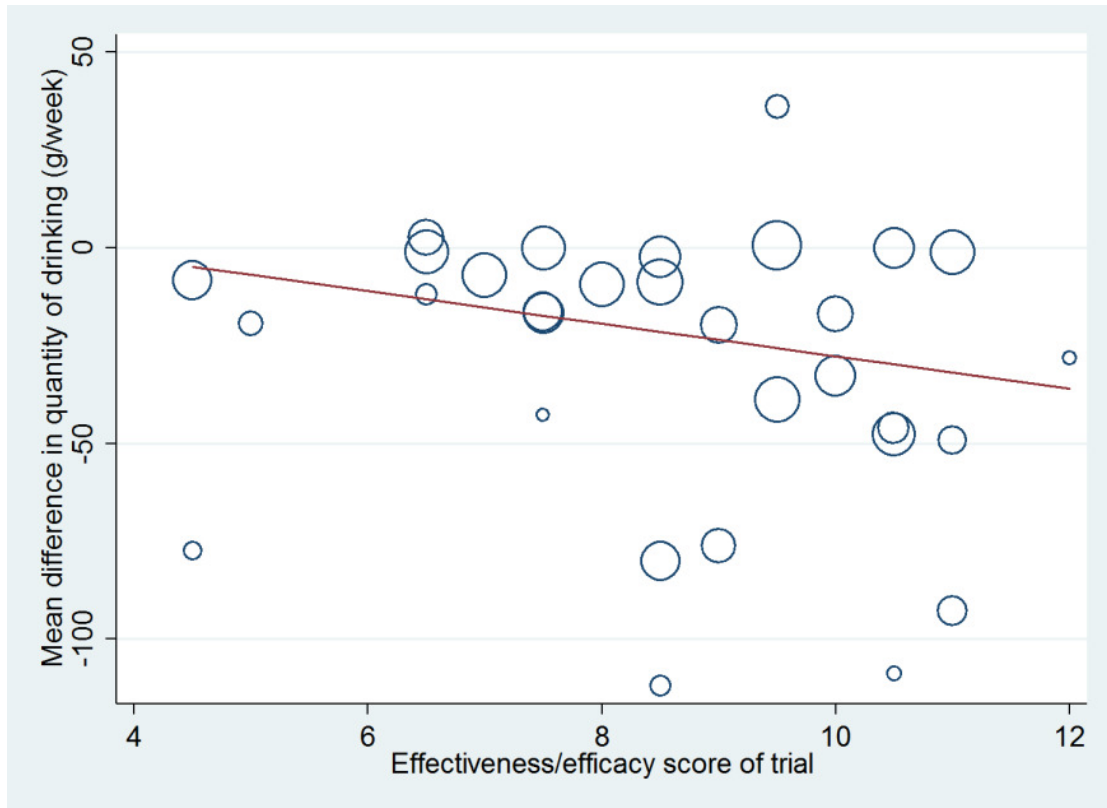
We excluded 16 studies (3816 participants) from the primary meta-analysis that reported at less than 12 months. One study (414 participants) reported at more than 12 months (Aalto 2000). Many trials reported that alcohol consumption reduced in both intervention and control group participants, with greater reduction in the intervention group, but the difference did not reach statistical significance. This suggests these studies followed the same trend as reported in the meta-analysis and would be unlikely to

change the conclusion.

Efficacy and effectiveness score (Analysis 1.15)

There was no significant difference between the pooled findings of the effectiveness trials and the pooled findings of the efficacy trials in terms of the quantity of alcohol consumed at 12 months (Analysis 1.15). The 16 effectiveness trials (7091 participants) showed that intervention participants drank 27 g/week (95% CI 13 to 41) less than minimal or no intervention participants. The 18 efficacy trials (8106 participants) also showed a benefit of brief intervention; participants in the intervention group drank 14 g/week less than minimal or no intervention group participants (95% CI 5 to 24). Meta-regression analysis showed that with each increase in unit of effectiveness efficacy score, the mean difference between intervention and minimal or no intervention increased by 4.1 g/week (95% CI -1.3 to 9.5). Trials that were more clinically representative tended to demonstrate greater effectiveness (Figure 9).

Figure 9. Meta-regression of quantity of drinking at 12 months on effectiveness/efficacy score of trial. Lower scores indicate greater efficacy.



Frequency of drinking and frequency of binge drinking (Analyses 1.16 to 1.19)

Binge drinking frequency (binges/week at 12 month) was reported in 15 trials (6946 participants) (Blow 2006; Daepfen 2007; D’Onofrio 2008; D’Onofrio 2012; Fleming 1997; Fleming 1999; Fleming 2004; Fleming 2010; Helstrom 2014; Longabaugh 2001; Ockene 1999; Rubio 2010; Schaus 2009; Soderstrom 2007; Spirito 2004; Analysis 1.16). Results showed moderate-quality evidence of a very small impact on binge drinking frequency (MD -0.08 binges/week, 95% CI -0.14 to -0.02). We analysed 11 trials (5469 participants) that reported the number of days drinking/week (Bernstein 2010; Cherpitel 2009; Crawford 2004; Curry 2003; Daepfen 2007; Field 2010; Fleming 2010; Helstrom 2014; Maisto 2001; Senft 1997; Spirito 2004). Analysis results indicated a very small impact (MD -0.13 days/week, 95% CI -0.23 to -0.04; Analysis 1.18; moderate-quality evidence). There was little difference between results at 6 months and 12 months for both outcomes (Analysis 1.17 and Analysis 1.19 respectively).

Intensity of drinking (Analysis 1.20)

Drinking intensity (g/drinking day, or a unit that could be converted to g/day) was reported by 10 trials (3128 participants) (Bernstein 2010; Cherpitel 2009; Crawford 2004; Curry 2003; Daepfen 2007; Helstrom 2014; Maisto 2001; Schaus 2009; Senft 1997; Spirito 2004). Results showed moderate-quality evidence of no impact at 12 months (MD -0.2 g/drinking day, 95% CI -3.1 to 2.7; Analysis 1.20).

Heavy and binge drinkers (Analysis 1.21; Analysis 1.22)

Percentages of heavy or at risk drinkers at 12 months was reported in 18 trials (7623 participants) (Altisent 1997; Beich 2007; Cherpitel 2009; Córdoba 1998; Curry 2003; D’Onofrio 2008; Daepfen 2007; Drummond 2014; Ettner 2014; Fernández 1997; Fleming 1997; Fleming 1999; Fleming 2004; Kaner 2013; Rubio 2010; Sommers 2013; Wallace 1988; Walton 2010). However, the definition of heavy drinking varied among trials. The difference between the brief intervention and minimal or no intervention arms in the percentage of heavy drinkers at 12 months was -9%

(95% CI -13 to -4; [Analysis 1.21](#)), although there was substantial heterogeneity ($I^2 = 77%$).

A total of 10 trials (4456 participants) reported the percentage of binge drinkers at 12 months ([Beich 2007](#); [Bischof 2008](#); [Curry 2003](#); [Ettner 2014](#); [Fleming 1997](#); [Fleming 1999](#); [Rubio 2010](#); [Scott 1990](#); [Sommers 2013](#); [Walton 2010](#)). Overall, results showed a significant reduction in the percentage of binge drinkers in the brief intervention group compared to the minimal or no intervention group (RD -7%, 95% CI -12 to -2; [Analysis 1.22](#)). There was substantial heterogeneity ($I^2 = 76%$).

Secondary outcomes

Laboratory markers of drinking ([Analysis 1.23](#); [Analysis 1.24](#))

We included three trials (1166 participants) that reported GGT at 12 months ([Romelsjö 1989](#); [Scott 1990](#); [Wallace 1988](#)). Analysis showed no significant difference between brief intervention and minimal or no intervention; there was no heterogeneity among trials (MD -0.9 IU/L, 95% CI -3.9 to 2.1, $I^2 = 0%$; [Analysis 1.23](#)). In two trials which reported GGT by gender, there was no significant difference in the effect size between men and women ([Scott 1990](#); [Wallace 1988](#); [Analysis 1.24](#)). [Seppa 1992](#) (95 participants) reported MCV and reported no significant difference between brief intervention and minimal or no intervention, both overall (MD 0.6 fl, 95% CI -1.6 to 2.8) and for each gender separately.

Alcohol-related harms

We assessed 20 studies that reported a measure of alcohol related harm ([Blow 2006](#); [Cherpitel 2009](#); [D'Onofrio 2008](#); [D'Onofrio 2012](#); [Drummond 2009](#); [Drummond 2014](#); [Fleming 2010](#); [Helstrom 2014](#); [Kaner 2013](#); [Lock 2006](#); [Longabaugh 2001](#); [McDevitt-Murphy 2014](#); [Mello 2008](#); [Monti 2007](#); [Romelsjö 1989](#); [Schaus 2009](#); [Spirito 2004](#); [Walton 2010](#); [Watson 2013](#); [Woolard 2013](#)). However, this outcome could not be analysed because of the large number of scales used to measure different types of harm. Sixteen studies reported no significant difference between the intervention and control groups in the chosen measure of alcohol-related problems or harm. [Blow 2006](#) reported a reduction in DrInC score in all groups. Two studies ([Kaner 2013](#); [Schaus 2009](#)) reported a short-term change. [Kaner 2013](#) reported fewer problems in the brief counselling group compared to the patient information leaflet group using the Alcohol Problems Questionnaire (APQ) at six months only. [Schaus 2009](#) reported fewer problems in the intervention group using the Rutgers Alcohol Problem Index (RAPI) at six months and nine months.

Patient satisfaction and quality of life measures

Only two studies reported patient satisfaction ([Drummond 2014](#); [Kaner 2013](#)). [Drummond 2014](#) reported no significant difference between groups. [Kaner 2013](#) stated that participants in the brief counselling group reported greater satisfaction than those in the patient information leaflet group.

Adverse effects

Five studies reported whether adverse effects were experienced by trial participants ([Beich 2007](#); [Bischof 2008](#); [Sommers 2006](#); [Sommers 2013](#); [Watson 2013](#)). [Beich 2007](#) reported that for women in the intervention group, binge drinking increased on average (ARR -30%, 95% CI -47 to -9). However, [Blow 2006](#) explicitly stated that younger adult women were most likely to decrease heavy episodic drinking after the intervention. Two trials reported that no adverse effects were experienced by any of the trial participants ([Bischof 2008](#); [Watson 2013](#)). Two trials reported adverse driving events but concluded there were no more in the intervention group than the control group ([Sommers 2006](#); [Sommers 2013](#)).

Health-related quality of life

[Daepfen 2007](#) reported a marginal significant improvement in the physical composite scale of SF-36 in the brief alcohol intervention group compared to the control groups. [Tait 2004](#) reported a significantly greater improvement in the intervention group in well-being as measured by the GHQ-12. Four studies reported no significant differences between intervention and control groups in quality of life measures ([Crawford 2004](#); [Drummond 2009](#); [Kaner 2013](#); [Lock 2006](#)).

Economic evaluations

Six studies either reported an economic evaluation in the trial report ([Drummond 2009](#); [Kunz 2004](#); [Lock 2006](#)) or as a separate publication ([Babor 2006](#); [Crawford 2004](#); [Fleming 1997](#)). Most suggested that the intervention was likely to be cost-effective. This finding is consistent with the results of a recent systematic review of cost-effectiveness, which reported strong evidence that brief intervention programs in primary care are a cost-effective option for tackling alcohol misuse ([Angus 2014](#)).

2. Extended intervention versus minimal or no intervention

See [Summary of findings 2](#).

Quantity of alcohol consumed per week (Analysis 2.1)

The impact of an extended intervention versus minimal or no intervention on consumption of alcohol at 12 months was reported by six trials (1296 participants) (Bischof 2008; Israel 1996; Maisto 2001; Monti 2007; Moore 2011; Richmond 1995). Most used the principles of motivational interviewing in the counselling intervention; Israel 1996 used cognitive behavioural therapy. These trials showed moderate-quality evidence of a reduction in consumption at 12 months (MD -14 g/week, 95% CI -37 to 9; Analysis 2.1). The level of heterogeneity in these findings was moderate ($I^2 = 41\%$). However, there were insufficient studies to allow further investigation of possible reasons for this heterogeneity.

Frequency of drinking and frequency of binge drinking (Analysis 2.2; Analysis 2.3)

At 12 months, two trials (456 participants) reported the frequency of binge drinking, providing moderate-quality evidence of very little difference between the groups (MD -0.08 binges/week, 95% CI -0.28 to 0.12; Longabaugh 2001; Monti 2007; Analysis 2.2). Maisto 2001 and Monti 2007 reported moderate-quality evidence for a small reduction in drinking days per week (MD -0.45, 95% CI -0.81 to -0.09; two studies, 319 participants; Analysis 2.3).

Intensity of drinking (Analysis 2.4)

Maisto 2001 recruited 158 participants and compared extended versus minimal intervention at 12 months. Results indicated little impact (MD -9 g/drinking day, 95% CI -26 to 9; Analysis 2.4; low-quality evidence).

Heavy and binge drinkers (Analysis 2.5; Analysis 2.6)

Moore 2011 (521 participants) reported the percentage of heavy drinkers at 12 months (Analysis 2.5). Bischof 2008 and Moore 2011 (777 participants) reported numbers of binge drinkers at 12 months (Analysis 2.6). There was very little difference between the extended intervention and minimal intervention arms for both outcomes.

Secondary outcomes

Laboratory markers of drinking (Analysis 2.7)

Israel 1996 reported GGT at 12 months. There was no significant difference between extended intervention and minimal intervention (MD -5.7 IU/L, 95% CI -20.3 to 8.9; Analysis 2.7). However, this result was based on only 59 participants.

Alcohol-related harms

Only Monti 2007 reported alcohol-related harms. There was no significant difference between groups in Rutgers Alcohol Problem Index score.

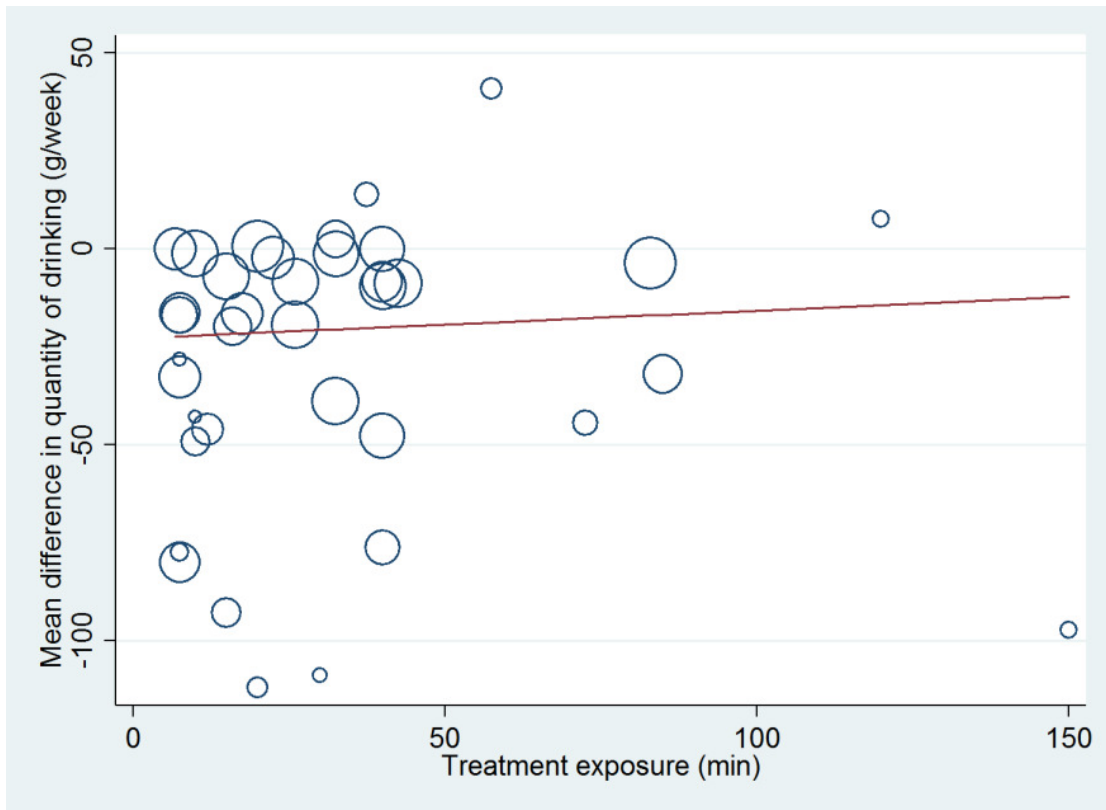
Other secondary outcomes

No studies reported patient satisfaction or health-related quality of life outcomes for this comparison.

Meta-regression of quantity of alcohol consumed on treatment exposure, including both brief and extended interventions

The meta-regression of quantity of alcohol consumed at 12 months on treatment exposure, reported above for trials which compared brief intervention with minimal or no intervention, was repeated, adding trials that assessed an extended intervention. This provided greater statistical power for the meta-regression, not only through inclusion of three extra trials (Israel 1996; Monti 2007; Moore 2011), but also through inclusion of trials with greater treatment exposure. Specifically, for the three trials that included both a brief intervention and an extended intervention (Bischof 2008; Maisto 2001; Richmond 1995), only the extended intervention results were compared with minimal or no intervention. As with the analysis restricted to the comparison of brief intervention with minimal or no intervention, meta-regression suggested no evidence of association between quantity of alcohol consumed at 12 months and increasing treatment exposure (an increase in the mean difference between the brief intervention and minimal or no intervention arms of 0.1 g/week (95% CI -0.3 to 0.5, $P = 0.72$, for each increase of 1 minute in the treatment exposure; Figure 10).

Figure 10. Meta-regression of quantity of drinking at 12 months on treatment exposure (mean duration of counselling for the participants in the trial), for trials comparing brief or extended intervention with control.



3. Extended intervention versus brief intervention

See [Summary of findings 3](#).

Quantity of alcohol consumed per week (Analysis 3.1; Analysis 3.2)

We included three trials (552 participants) in the meta-analysis comparing alcohol consumption at 12 months for extended versus brief interventions ([Bischof 2008](#); [Maisto 2001](#); [Richmond 1995](#)). There was no difference in consumption between participants who received an extended intervention and those who received a brief intervention, although the associated confidence interval was very wide (MD 2 g/week, 95% CI -42 to 45; [Analysis 3.1](#); low-quality evidence). The extended interventions varied. [Bischof 2008](#) provided four half hour telephone sessions based on motivational interviewing. [Maisto 2001](#) provided one 30 to 45 minute motivational interview followed by two shorter sessions. [Richmond 1995](#) provided five sessions totaling over 60 minutes.

We performed sensitivity analysis that included an additional trial ([Spirito 2011](#)) and imputed unreported standard deviations ([Analysis 3.2](#)). This made little difference to the result (MD 1 g/week, 95% CI -39 to 41).

Frequency and intensity of drinking (Analysis 3.3; Analysis 3.4)

No trials reported the number of binges per week for extended versus brief intervention. [Maisto 2001](#) reported frequency in terms of number of drinking days per week and intensity in terms of drinks per drinking day at 12 months for this comparison ([Analysis 3.3](#); [Analysis 3.4](#)). There was low-quality evidence of little difference between extended and brief interventions for both outcomes (MD -0.5 drinking days/week, 95% CI -1.2 to 0.2; MD -1.7 g/drinking day, 95% CI -18.9 to 15.5; 147 participants).

Heavy and binge drinkers (Analysis 3.5)

Two trials reported numbers of binge drinkers at 12 months ([Bischof 2008](#); [Spirito 2011](#)). Pooled data showed no statistically significant difference in the percentage of binge drinkers between extended and brief interventions (RD 2%, 95% CI -8 to 12; 339 participants; [Analysis 3.5](#)).

No trials that compared extended and brief interventions reported numbers of heavy drinkers.

Secondary outcomes

No trials that compared extended and brief interventions reported laboratory markers at 12 months. The trials did not report any measures of alcohol-related harm, patient satisfaction or quality of life.

ADDITIONAL SUMMARY OF FINDINGS *[Explanation]*

Extended intervention compared to no or minimal intervention for people with hazardous or harmful alcohol consumption						
Patient or population: people with hazardous or harmful alcohol consumption Setting: primary care (directly accessible to participant, no referral required) Intervention: extended intervention Comparison: no or minimal intervention						
Outcomes	Anticipated absolute effects* (95% CI)		Relative effect (95% CI)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Risk with control	Risk with Extended intervention				
Quantity of drinking (g/week) at 12 months	The mean quantity of drinking (g/week) at 12 months was 236 g/week	MD 14.43 g/week lower (37.41 lower to 8.54 higher)	-	1296 (6 RCTs)	⊕⊕⊕○ MODERATE ¹	
Frequency of drinking (no. binges/wk) at 12 months	The mean frequency of drinking (no. binges/wk) at 12 months was 1.3 binges/week	MD 0.08 binges/week lower (0.28 lower to 0.12 higher)	-	456 (2 RCTs)	⊕⊕⊕○ MODERATE ¹	
Frequency of drinking (no. days drinking/week) at 12 months	The mean frequency of drinking (no. days drinking/week) at 12 months was 2.1 drinking days/week	MD 0.45 drinking days/week lower (0.81 lower to 0.09 lower)	-	319 (2 RCTs)	⊕⊕⊕○ MODERATE ¹	
Intensity of drinking (g/drinking day) at 12 months	The mean intensity of drinking (g/drinking day) at 12 months was 76.6 g/day	MD 8.51 g/day lower (25.69 lower to 8.67 higher)	-	158 (1 RCT)	⊕⊕○○ LOW ¹²	

* **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% CI).

CI: Confidence interval; **RR:** Risk ratio; **OR:** Odds 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

¹ High risk of performance bias due to difficulties with blinding participants and providers

² Imprecision suggested by small number of trials/participants

Extended compared to brief intervention for people with hazardous or harmful alcohol consumption						
Patient or population: people with hazardous or harmful alcohol consumption Setting: primary care (directly accessible to participant, no referral required) Intervention: extended intervention Comparison: brief intervention						
Outcomes	Anticipated absolute effects* (95% CI)		Relative effect (95% CI)	No of participants (studies)	Quality of the evidence (GRADE)	Comments
	Risk with brief intervention	Risk with Extended				
Quantity of drinking (g/week) at 12 months	The mean quantity of drinking (g/week) at 12 months was 251 g/week	MD 1.54 g/week higher (42.01 lower to 45.10 higher)	-	552 (3 RCTs)	⊕⊕○○ LOW ¹²	
Frequency of binge drinking (no. binges/wk) at 12 months - not measured	-	-	-	-	-	
Frequency of drinking (no. days drinking/week) at 12 months	The mean frequency of drinking (no. days drinking/week) at 12 months was 2.82 days drinking/week	MD 0.51 drinking days/week lower (1.21 lower to 0.19 higher)	-	147 (1 RCT)	⊕⊕○○ LOW ²³	
Intensity of drinking (g/drinking day) at 12 months	The mean intensity of drinking (g/drinking day) at 12 months was 70 g/drinking day	MD 1.7 g/drinking day lower (18.86 lower to 15.46 higher)	-	147 (1 RCT)	⊕⊕○○ VERY LOW ¹²³	

* **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% CI).

CI: Confidence interval; **RR:** Risk ratio; **OR:** Odds 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

¹ Imprecision suggested by very wide confidence intervals

² High risk of performance bias due to difficulties blinding participants and providers

³ Imprecision suggested by small number of trials/participants

DISCUSSION

Summary of main results

We included 69 studies (33,642 participants), of which 34 studies (15,197 participants) provided data for our primary outcome. At 12 months, when compared with minimal or no intervention, brief intervention reduced the quantity of alcohol consumed by 20 g/week (95% CI 12 to 28), which equates to two to three UK standard units. We undertook three sensitivity analyses that indicated the effect is robust. Excluding trials that did not report adequate allocation concealment, including trials where standard deviations were imputed, and assuming different intra-cluster correlation coefficients for cluster-randomised trials, all showed a similar degree of benefit to the primary analysis. A funnel plot showed some evidence of publication bias but this diminished after adjustment for year of publication (Figure 4).

Most participants were male (70%). Ethnicity was poorly reported, but in those trials that provided these data, most participants were white (70%). Mean baseline consumption was about 310 g/week, but this varied among trials from about 90 to 460 g/week (11 to 57 UK standard units).

There was substantial heterogeneity among trials in terms of setting (general practice or emergency care settings), populations enrolled, screening instruments used, baseline consumption of alcohol, and the brief intervention and minimal or no intervention conditions delivered. Therefore, the statistical heterogeneity in the meta-analyses was not surprising. Subgroup analyses showed that heterogeneity of findings was restricted to individually randomised trials and trials with high treatment exposure.

Further meta-analyses demonstrated very small significant impact of brief intervention in terms of days of drinking and binge days per week, amounting to a reduction of one day per two months and no significant impact on amount consumed per drinking day. The percentage of heavy and binge drinkers at 12 months showed a small but significant decrease of 9% and 7% respectively at 12 months in the intervention groups compared to participants receiving minimal or no intervention.

Further meta-analyses suggested that people receiving an extended intervention may reduce their consumption compared to participants receiving minimal or no intervention at 12 months, but provided no evidence that extended interventions reduce consumption any more or less than brief interventions. These meta-analyses were less robust. They were based on much smaller groups of participants than the primary meta-analysis and may be confounded by the fact that attendance at multiple sessions was not always reported (i.e. participants may not all have received a full 'extended' intervention). Extended interventions differed from brief interventions not only in terms of contact time with participants but also because they were more likely to involve counselling (shorter interventions may 'draw on' counselling techniques but are likely to be too short to make full use of them).

We assessed the impact of both duration and type (advice or counselling) of intervention on drinking outcomes. Firstly, we performed a meta-regression, which found little evidence of a link between treatment exposure time and reported alcohol consumption. This seems to support the meta-analysis suggesting that there is little difference in impact between brief and extended intervention approaches. Secondly, we carried out a subgroup analysis according to whether interventions were advice- or counselling-based. This suggested that counselling-based interventions, despite being more intensive and providing more contact with participants, may be associated with smaller reductions in consumption than advice-based interventions. Taken together, these analyses suggest little evidence of a dose response effect, either in terms of longer duration or higher intensity of intervention.

Clinical importance

At baseline, the mean level of consumption of participants was 244 g/week (equivalent to 30 UK standard units), although this differed according to year of publication (older studies 313 g/week versus newer studies 181 g/week). The mean reduction provided by a brief intervention reported here of 20 g/week (2.5 UK units) would mean that many participants would continue to drink at a level that would be considered hazardous according to recommendations in most countries. However, this reduction is still likely to be beneficial at an individual level, given the number of disease conditions at least partly attributable to alcohol and given that the higher the baseline level of drinking, the stronger the effects of any given reduction in terms of all-cause mortality (Rehm 2013). At a population level, any reduction is likely to have a significant impact on health, quality of life and healthcare resource use, given that over 5% of the global burden of disease is attributable to alcohol (WHO 2014).

Effectiveness/efficacy

There was no significant difference between trials classified as effectiveness and efficacy trials in the effect of brief intervention on the quantity of alcohol consumed, although meta-regression suggested that trials with higher scores on our scale (i.e. more clinically representative or 'real world' trials) may show more intervention effect. This difference is counterintuitive. One would expect that higher scoring trials would result in smaller effect sizes because they are less tightly controlled, but there may be some interaction with publication date or some other feature of the trials. Other tools (e.g. Gartlehner 2006; Wieland 2017) have been developed to more precisely measure and assess efficacy/effectiveness status and we plan to carry out comparative work using these tools.

This lack of evidence of any difference in the current review may indicate insensitivity in our descriptive classification tool. In some papers, authors did not report information relating to certain items. In these cases, we ascribed a mid-value score for that item so as not to tip the study toward the efficacy or effectiveness domain.

This may have reduced variation in the final scores (there were not many extreme scores, particularly on the efficacy side of the scale) and led to clustering of trials toward the middle of this scale. However, although the trials were skewed toward the effectiveness domain, they were distributed along a continuum. Moreover, it is highly unlikely that there would be any pure efficacy studies, since the trial protocol would need to be acceptable and relevant to clinicians (and ethics committees) before it could be enacted in health service contexts. It is possible that any intervention effect may be related to individual factors, which were combined in the overall efficacy/effectiveness score. It is beyond the scope of this review to investigate this further.

Screening

It was clear in this review (as in other work) that many trials reported reductions in alcohol consumption in minimal or no intervention arms at follow-up. It is difficult to assess if this might be due to regression to the mean or an impact of the screening itself. One trial reported that a significant proportion of participants reduced their drinking between screening and assessment, and so the brief intervention was delivered to some participants who were no longer eligible for such an approach (McIntosh 1997). Therefore, it is possible that screening alone might represent an impetus for change (McCabridge 2011). This should be further investigated in future work.

Publication date

Several plausible reasons might explain why more recent trials tend to demonstrate less impact on alcohol consumption than older trials. Firstly, the definition of excessive drinking has reduced over time in terms of consumption per week. In the earliest published trial included in this review (Wallace 1988), the inclusion criteria stipulated at least 35 and 21 units/week for men and women respectively. However, a more recent study recruited participants drinking at least 21 units/week (men) or 14 units/week (women) (Helstrom 2014). Mean baseline consumption in brief alcohol intervention trials has correspondingly reduced over time. In the original review, mean baseline consumption was 313 g/week (around 39 UK units/week), whereas in newly added studies for this update it was 181 g/week (around 23 UK units/week). Typically, trials did not report the upper cut-off point for exclusion in terms of consumption levels but these differences between baseline characteristics of participants in older versus more recent trials demonstrates the lowering of inclusion criteria over the years. Moreover, many recent trial participants have been enrolled with consumption levels only just above recommended drinking levels. Consequently, with lower baseline alcohol consumption, there is less change required before participants reach a lower risk drinking level.

Secondly, minimal or no intervention conditions in more recent trials are more likely to provide some kind of alcohol-related advice

or information, which might highlight alcohol consumption to control group participants and encourage them to decrease their consumption, reducing the difference in effect between intervention and control groups. A subgroup analysis by content of control condition appeared to support this, suggesting a larger difference between intervention and control group where no alcohol-related information is provided (Analysis 1.12).

Finally, most earlier trials took place in general practice-based primary care, whereas more recently published trials were more likely to take place in emergency departments, where the effect size appears to be smaller (Analysis 1.10).

Overall completeness and applicability of evidence

This review suggests that brief interventions are effective at reducing alcohol consumption compared to minimal or no intervention in hazardous and harmful drinkers presenting to general practice or emergency care. This finding does not apply to low risk or alcohol dependent drinkers, or necessarily to other health or social care settings. This update saw the inclusion of a greater proportion of trials which took place in emergency departments, which enabled us to formally compare effectiveness between emergency care and general practice. Although the raw subgroup analysis suggests a difference in effectiveness between settings, this result is confounded by publication date since so many of the emergency care trials are recent.

Whereas the previous published version of our review reported there was no evidence of a significant effect for women, the addition of further data has changed this conclusion, although there are still only seven trials reporting data separately for 1350 women. This analysis suggests that brief intervention has a significant impact on weekly alcohol consumption compared to women experiencing minimal or no intervention conditions. In the context of the reported closing of the gap between hazardous and harmful alcohol consumption by women compared to men, this is an important finding (Slade 2016). The significant effect for men also stands from the original review.

Only four trials were based in countries not defined as high income by the WHO, so this evidence cannot be assumed to apply to countries where the structure or functioning of primary care differs, or where less liberal attitudes to alcohol consumption may exist. Most of the participants in trials were also Caucasian and so any future evaluations should specifically aim to enrol a wider diversity of participants.

Quality of the evidence

The direction of evidence is broadly consistent. Most studies in the primary meta-analysis (82%) reported a reduction in consumption

for brief intervention compared to minimal or no intervention participants.

We downgraded the evidence quality assessment to moderate due to a systematic risk of bias in the studies. Blinding is notoriously difficult in trials of complex conversation-based interventions, because the intervention cannot easily be disguised from participants, and this is something that is unlikely to change. Indeed, it has been suggested that GRADE assessments may not adequately describe the evidence base of complex interventions and an extension is required to adequately reflect them (Movsisyan 2016). Empirical research has shown that trials where the participant and treatment provider are not blinded may be more likely to report significant effects of the intervention (Schulz 1995). Attrition bias also tends to be high in these types of trials. If participants who dropped out of the brief intervention groups had higher alcohol consumption than those who did not, our estimated reduction in alcohol consumption due to brief intervention would be an overestimate of the real effect.

Nevertheless, the estimated reduction in the quantity of alcohol consumed per week was sufficiently marked that the real effect is likely to be a reduction in alcohol consumption. Furthermore, the random-effects model which we used assumes that the effect of treatment is different in different populations and that the estimated reduction in alcohol consumption of 20 g/week is the mean treatment effect, averaged over all populations. Therefore, the findings provide strong evidence that brief interventions are effective in many populations.

As in the original review, an analysis had been planned linked to the level of loss to follow-up, but was not possible. Loss to follow-up was defined in different ways across the studies, and it was not always clear whether allowance had been made for loss to follow-up in the reported results. However, an analysis that excluded trials at high risk of bias due to incomplete outcome data gave similar results to those from the primary meta-analysis. Further research should elucidate how to reduce this effect.

Potential biases in the review process

We made every effort to locate all existing eligible trials, regardless of publication status or language, and we searched grey literature and references in addition to bibliographic databases to reduce publication bias. We contacted authors to clarify the reporting of data used in the meta-analyses where necessary. We further assessed potential bias from selective reporting by comparing the outcomes proposed in the protocols (where available) or methods sections with those in the final study report. The results of these assessments appear in the risk of bias tables ([Characteristics of included studies](#)).

We worked hard to ensure that the inclusion criteria were applied consistently to potentially eligible studies. The term 'brief intervention' is used to describe many different types of interventions, and so we further defined the inclusion criteria in terms of number

and duration of intervention sessions. It was sometimes difficult to tell whether the setting fit our definition of 'primary care', and in some studies there was a mixture of hazardous, harmful and potentially dependent drinkers. The eligibility of all the studies was assessed by two of the review authors independently, but where they were uncertain the study was always discussed by two authors (EK, FB) to make the final decisions as consistent as possible.

One potential weakness of the studied outcomes is that they are predominantly self-reported, so may be susceptible to social desirability bias. Many of the studies made efforts to minimise this effect and a recent trial suggests that for questions about alcohol consumption this may not be a big problem (albeit the participants were students, not the general public) (Kypri 2016).

The addition of 42 studies to the updated review provided a lot of extra data. Some of these data were added to existing analyses, but others enabled us to carry out subgroup analyses which were not proposed in the original protocol, often because they could not have been foreseen at the time. For example, brief interventions were rarely carried out in emergency departments 10 years ago, but many of the newly added studies were, enabling subgroup analysis of brief interventions in general practice versus emergency care. Although it can introduce bias to carry out subgroup analyses which were not pre-specified in the review protocol, we feel that each of the additional subgroup analyses in this update is backed up by findings in the wider literature.

Agreements and disagreements with other studies or reviews

Several reviews have been published since the previous version of this review was published in 2007. A systematic review of reviews of brief interventions in primary care assessed 24 systematic reviews containing a total of 56 trials (O'Donnell 2014). The review concluded that although there was good evidence of effectiveness of these interventions, there were gaps in terms of particular populations (e.g. women, younger and older drinkers, ethnic minorities) and the optimum length and frequency of such interventions. Since then, further reviews have addressed some of these issues.

Another review of seven systematic reviews suggested that 15 minutes of brief intervention was better than usual care or longer input, and that more sessions were better than single session (Álvarez-Bueno 2015). Our review suggests little extra impact from longer duration, but this indicates that splitting that longer duration into multiple sessions may increase the effect. A systematic review addressed alcohol interventions (including digital or telephone delivered) across settings aimed at women, or reporting by gender (De Paula Gebara 2013). Only three (of 36) trials took place in primary care and their results were mixed. Two reviews addressed brief interventions in a younger population. One included trials of adolescents and young adults aged between 11 and 30 years and found a significant reduction in consumption and alcohol-related problems for up to one year, which did not appear

to vary across participant demographics, intervention length, or intervention format (Tanner-Smith 2015). A second reviewed included seven trials along with practice projects, a pilot study and an observational study of adolescents and young adults aged between 12 and 25 years in emergency care and reported inconclusive results, although it also reported that six trials found reductions in alcohol use for all participants (Diestelkamp 2016). Four trials reported reductions in consumption or harms but none reported both. Another review of brief interventions in emergency care included 28 trials (comprising digital and non face-to-face interventions as well as face-to-face brief interventions) and reported a small significantly significant reduction in consumption at 12 months (Schmidt 2016). Schmidt 2016 also carried out moderator analyses for intervention type (face-to-face versus not), duration of intervention (< 15 minutes versus > 15 minutes), intervention deliverer (emergency department staff versus external), study quality (low risk of bias versus high) and control ingredients (treatment as usual, leaflet, unspecific advice, personalised alcohol feedback) and reported that none of these features altered the direction or significance of the meta-analysis results.

The systematic review referenced in our original review reported a -38 g/week reduction in alcohol consumption attributed to brief intervention, as did our previous meta-analysis (Bertholet 2005). This update reports a reduced effect size resulting from brief alcohol intervention, which may be due to the enrolment of less heavy drinkers into these types of trials as well as increasing alcohol content in control conditions. This update agrees with most other reviews in reporting a small but significant impact of brief interventions on hazardous and harmful drinkers in primary care settings.

Brief interventions are increasingly being enhanced or replaced by digital interventions (Kaner 2017), which may prove less costly to deliver and maintain (after development work is complete). It is plausible that digital interventions may have a greater reach to populations who do not engage with primary care, since people will be able to access them through a computer or mobile device anywhere or at any time. However, it is also possible that digital delivery will not suit some people and some may be less likely to complete an intervention if there is no personal encouragement. Nevertheless, digital input may be able to support practitioners' work in helping participants reduce their alcohol consumption by providing additional self-assessment, self care or self-monitoring resources. Kaner 2017 provides comparison and further discussion of digital versus practitioner-delivered interventions.

AUTHORS' CONCLUSIONS

Implications for practice

This review provides moderate-quality evidence that brief interventions delivered in primary care reduce alcohol consumption

in hazardous and harmful drinkers by an average of 20 g/week (the equivalent of approximately two to three UK standard drinks) compared to controls receiving usual care, screening or assessment only, or minimal alcohol advice. Whilst the effect sizes due to brief alcohol intervention are relatively small, because of the high proportion of the population who attends primary care, this is likely to result in a positive public health or population level benefit. The effect is robust and was found up to one year following the brief intervention. Brief interventions appear to be effective for both men and women but there was insufficient evidence to make any judgements about differential effectiveness in different ethnic groups. There was a suggestion of a reduced effect in younger people compared to the general population aged 18 years or more, and in emergency care compared to general practice, but this disappeared when adjusted for the overall reduction in effect size due to newer trials. Longer or more intensive intervention appeared to have little effect in significantly improving outcomes. The effects of brief alcohol interventions are typically found in trials that report outcomes in terms of weekly alcohol consumption. It is not clear why the impact seems smaller for frequency or intensity of drinking, although, within a weekly (7-day) frame of reference, the scope for variability in days drinking or number of high intensity ('binge') events is relatively low. A further challenge in this field is the very wide range of different outcome measures, which are reported over greatly varying periods of time. There were too few trials reporting many of these outcome measures to provide sufficient statistical power to detect potentially small effects.

Implications for research

Although brief interventions seem to be effective at reducing hazardous and harmful consumption in adult men and women primarily based in high income countries, there is a clear need for more evaluative research on brief interventions with younger people, from cultural minority groups and also in low and lower-middle income countries. Moreover, there is some suggestion that screening alone may result in alcohol consumption reduction and this should be investigated further. A quarter of the included trials in this review could not contribute to the meta-analysis, either because they did not report a measure of consumption that could be converted to g/week of alcohol, or because the follow-up time was too short (< 6 months). There is an urgent need for standardisation of outcomes in alcohol trials and future trials should report longer follow-up times to aid understanding of the duration of effect. Future studies should aim to capture adverse effects of brief interventions. A further methodological improvement would be to address the issues of high attrition bias within these interventions and to enable GRADE to make a more sophisticated judgement of quality in trials of complex conversation-based interventions. Finally, future research direction should focus on implementation issues, including a more precise specification of brief intervention components.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Aalto 2000

Methods	Parallel group RCT: extended brief intervention versus brief intervention versus active control. ITT: outcome data were based on imputation of baseline values to participants lost to follow-up	
Participants	Setting: Finland; primary care clinic. Participants 20 to 60 years, consuming ≥ 280 g absolute ethanol/week or CAGE ≥ 3 for men, ≥ 190 g absolute ethanol/week or CAGE ≥ 2 for women; excluded if severe psychiatric disease, or at least one detox treatment, or alcohol dependence, or alcohol-related disease; screened by self-administered health questionnaire including CAGE and quantity-frequency consumption questions. Number randomised = 414; 71% male; mean age = 41.6 years; 18% comprehensive school, 7.3% vocational school, 12.7% college or university; 18.7% working/studying, 13.6% unemployed, 6% retired. At baseline: mean drinking amount per week = 286 g for men, 165.5 g for women, 259.8 g overall; mean drinking times per week = 2.2 for men, 2.1 for women; mean usual drinking amount per occasion = 139.2 g for men, 85.8 g for women; mean CAGE = 3.2 for men, 2.8 for women	
Interventions	Group A (N = 149) received brief intervention from GP or nurse at baseline, 2, 6, 12, 18, 24 and 30 months. Intervention was 10 to 20 minutes based on FRAMES according to the needs of individual participants. Group B (N= 137) received the same intervention less frequently: at baseline, 12 and 24 months. Group C (N = 128) received advice to reduce drinking and contact their GP in the event of health problems; were not told about 36 month follow-up	
Outcomes	Mean drinking amount per week; drinking times per week; usual drinking amount per occasion; CDT, AST, ALT, GGT, MCV Assessed at 36 months (blood tests additionally assessed at each brief intervention)	
Funding source	Not reported.	
Declaration of interests	Not reported.	
Notes	Lahti project. Loss to follow-up: <ul style="list-style-type: none"> • Group A: 61/149 (41%). • Group B: 55/137 (40%). • Group C: 55/128 (43%). 	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Aalto 2000 (Continued)

Random sequence generation (selection bias)	Low risk	“Randomisation was performed by each participating general practitioner: before a participant arrived for a feedback session, the general practitioner drew a card from a mixed pack that included equal numbers of A, B, and C signs to refer the participant to the group which the participant was to be allocated” (p. 1682)
Allocation concealment (selection bias)	High risk	GP could see allocation and it was possible to subvert.
Blinding of treatment providers	High risk	GP provided intervention to all arms therefore risk of contamination between arms was high
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Unclear whether outcome assessors were blinded.
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Unclear risk	Did not specify exact alcohol outcome measures in the methods

Altisent 1997

Methods	Parallel group RCT: brief intervention versus active control ITT: no.
Participants	Setting: Spain; primary care clinic. Participants aged 15 to 75 years with a weekly consumption of > 280 g for men and > 168 g for women; excluded if current treatment for alcohol problems or hepatologic problems or concomitant diseases requiring alcohol abstinence or MALT > 11; screened by MALT scale for alcohol dependence. Number randomised = 139; 100% male; mean age = 45 years; 47% comprehensive school, 19% vocational school, 33% college/university; 49% working/studying, 36% unemployed, 15% retired. At baseline: mean weekly alcohol consumption = 57 units (1 unit = 8 g alcohol)
Interventions	Intervention group (N = 75) received five minutes general advice from GP with support material plus a five-visit program over the year. NB, 21 participants were subsequently excluded; 54 received intervention treatment. Control group (N = 64) received a single session of brief advice from GP. NB, 19 were subsequently excluded; 45 received the control treatment

Altisent 1997 (Continued)

Outcomes	Percentage reduction in alcohol consumption; MALT test; Goldberg score; per cent drinking < 35 units/week Assessed at 12 months.
Funding source	Este trabajo se realizo con la ayuda proporcionada por la Beca FS 93/0882. [This work was conducted with support provided by scholarship/grant number FS 93/0882]
Declaration of interests	Not reported.
Notes	Loss to follow-up: <ul style="list-style-type: none"> • Intervention group: 20/54 (37%). • Control group: 15/45 (33%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	States only that assignment was made by simple randomisation ("aleatorizacion simple", p. 122)
Allocation concealment (selection bias)	Low risk	Consecutively numbered, sealed envelopes ("los sobres cerrados y numerados", p. 122)
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported in results.

Babor 2006

Methods	Cluster RCT: brief intervention (two arms delivered by different professionals) versus usual care; 15 practices randomised Used a pre-post repeated measures group design. ITT: yes.
Participants	Setting: USA; family or internal medicine clinics in a managed care organisation (MCO) Participants ≥ 18 years were screened with Health Assessment Survey including AUDIT

	<p>and selected if they scored 8+ (men) or 7+ (women) Number of clinics randomised = 15; number of participants randomised = 50,411, 60% male; mean age followed up at 3 months (n = 1379) = 45.9 years At baseline: mean drinks per week (defined by AUDIT questions 1 to 2) = 13.8 (men); 8.7 (women)</p>	
Interventions	<p>Group P (N = 17,257 randomised, 1151 sampled for follow-up) received brief intervention (according to their level of drinking - see below for detail) from physician or physician assistant Group S (N = 15,938 randomised, 1124 sampled for follow-up) received brief intervention from specialist, i.e. mostly nurses Group C (N = 17,216 randomised, 1955 sampled for follow-up) was the control group and received "patient care as usual" (not defined) Within each group treatment allocated by drinking level: Zone I (modified AUDIT 7 to 15 for women and men aged > 65 years; 8 to 15 for men < 65 years) participants received simple advice (3 to 5 minutes) and an information brochure; Zone II (AUDIT 16 to 19) received advice and a more extensive self-help manual; Zone III (AUDIT > 19) specified advice and referral to specialty care for alcohol assessment and treatment Advice comprised:</p> <ul style="list-style-type: none"> • feedback of screening scores and any problems or symptoms identified; • discussion of the participant's risk level; • connection of drinking risk to actual or potential problems and information about safer drinking limits; • commitment to a goal of cutting back or stopping; and • presentation of the brochure and encouragement to reach the established goal. 	
Outcomes	<p>Number of drinks per week; per cent at risk drinkers who had changed drinking at three months; health care utilisation (all days, outpatients visits, inpatient days, emergency department visits, ADM-related days (mental health) Assessed at 3 and 12 months.</p>	
Funding source	<p>This research was supported by grant no. 029620 from the Robert Wood Johnson Foundation</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>Cutting Back programme. Payment: the MCOs were reimbursed; participants received no payment Loss to follow-up: 81/575 (14%) overall (not reported by arm)</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random numbers from table generated by independent statistician (p. 625)
Allocation concealment (selection bias)	Low risk	Allocation by independent statistician - not a member of the team (p. 625)

Babor 2006 (Continued)

Blinding of treatment providers	Low risk	Cluster-randomised trial and all participants from a particular clinic received the same intervention, so therefore risk of contamination between arms was low
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Computer-assisted telephone interview by independent survey organisation - interviewers not aware of the study condition (p. 625)
Incomplete outcome data (attrition bias) All outcomes	High risk	Only reported data from those who completed, not those randomised
Selective reporting (reporting bias)	High risk	Only primary outcome fully reported.

Bazargan-Hejazi 2005

Methods	Parallel group RCT: brief intervention versus usual care. ITT: yes.
Participants	Setting: USA; emergency department. Participants Aged ≥ 18 years, English or Spanish speaking, excluded if received alcohol counselling in last year, or cognitive impairment precluded informed consent, or medical treatment prevented them being interviewed, or in policy custody; screened with CAGE ≥ 1 Number randomised = 295; 80% male; mean age = 38.8 years; 64% Black, 30% Hispanic, 6% other; 84% high school or above, 16% less than high school diploma At baseline: mean number of drinks per day = 3.29; number drinking at least 6 drinks per occasion at least weekly = 186
Interventions	Both groups received CAGE for screening plus 30 minute baseline assessment questionnaire Both groups received a health packet that contained a variety of health information (not restricted to alcohol) Intervention group (N = 139) received a brief negotiated semi-scripted interview lasting 15 to 20 minutes from peer educators, which integrates elements of motivational interviewing and readiness to change Control group (N = 142) received usual care (undefined). These participants received a referral to local treatment programs only if the participant requested one of the physicians or if the treating physician made an independent diagnosis and referral
Outcomes	Percentage moving to a lower risk category (based on AUDIT); percentage scoring > 7 on AUDIT Assessed at 3 months.

Funding source	Supported by grants from the Center for Minority Health and Health Disparities, National Institute on Alcohol Abuse and Alcoholism (U24AA11899-05) and the National Institutes of Health, National Center for Research Resources and Research Centers in Minority Institutions (G12-RR03026-16)	
Declaration of interests	Not reported.	
Notes	Payment: USD 10 for baseline assessment, USD 25 for follow-up interview session Loss to follow-up: <ul style="list-style-type: none"> • Intervention group: 51/139 (37%). • Control group: 45/142 (32%). 	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"Each of the three health promotion advocates performed random allocation for their own enrollees, assigning the first participant by a flip of a coin, and alternating status thereafter" (p. 69)
Allocation concealment (selection bias)	High risk	Alternate allocation is predictable.
Blinding of treatment providers	High risk	Health promotion advocates delivered the intervention and interacted with participants from both arms of the study
Blinding of participants	Low risk	"Alcohol information was not included in the health packet [received by all participants] to avoid contamination" (p. 69)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Enrollees were not followed up by the same health promotion advocate who assessed them initially. Participants were notified not to reveal their group assignments to project staff at any time" (p. 69)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Unclear risk	Number of participants for follow-up in each AUDIT subgroup was not specified. However, we did not include these subgroups in analyses, so this aspect did not introduce bias

Beckham 2007

Methods	Parallel group RCT: brief intervention versus usual care. ITT: no. Participants were “given the opportunity to assess themselves” for hazardous alcohol use with AUDIT, i.e. self-selecting not comprehensive sample
Participants	Setting: USA; rural community healthcare centres. Participants: Aged ≥ 18 years with AUDIT 8+; 64% had some psychiatric diagnoses; excluded if pregnant or suicidal; screened with AUDIT Number randomised = 28; 46% male; 97% Caucasian; age range = 19 to 55 years Baseline information for total sample not reported.
Interventions	Both groups received AUDIT for screening and 30 minute baseline assessment Motivational interviewing group (N = 12 assessed) received one 45 to 60 minute motivational interviewing session, also utilising FRAMES, from a nurse practitioner Control group (N = 13 assessed) received usual care; after six week follow-up received GGT results and local referral for low-cost local alcohol treatment
Outcomes	Drinks per day (1 drink = 12 ounces of beer, 5 ounces of wine, 1.5 ounces of spirits or hard liquor); GGT Assessed at 6 weeks.
Funding source	Not reported.
Declaration of interests	“No relationship exists between the author and any commercial entity or product mentioned in the article that might represent a conflict of interest. No inducements have been made by any commercial entity to submit the manuscript for publication”, p. 110
Notes	Loss to follow-up not reportable: number randomised to each arm at baseline not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	“Participants were randomly assigned to the intervention or control group using a table of random numbers” (p. 106)
Allocation concealment (selection bias)	Unclear risk	Allocation concealment not described.
Blinding of treatment providers	High risk	Not blinded; no reported attempt to reduce bias.
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Blinding of outcome assessor not described.

Beckham 2007 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up not reported.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported in results.

Beich 2007

Methods	Parallel group RCT: brief intervention versus inactive control ITT: yes.
Participants	Setting: Denmark; general practice surgery. Participants aged 18 to 64 years, scheduled to see 39 general practitioners, AUDIT score between 8 and 21 (those with AUDIT > 12 were screened for dependency and referred if necessary); excluded if they reported weekly consumption > 35 drinks, had severe acute illness, or reading disability or spoke a foreign language, or illiterate, or had mental or physical impairment, or inebriated, or currently being treated for an alcohol use disorder, or pregnant; screened with AUDIT Number randomised = 906; 67% male; other characteristics not reported for sample At baseline: usual weekly consumption reported > 35 drinks = 24% of men, 17% of women; previous weekly consumption reported > 35 drinks = 38% of men, 32% of women; weekly binge drinking (defined as 6+ units) = 41% of men, 21% of women; monthly binge drinking = 82% of men, 69% of women
Interventions	Both groups received AUDIT for screening, which also provided many baseline data, and a secondary baseline questionnaire (time not reported) Intervention group (N = 442) received one 10 minute session from the GP based on the 'drink less' protocol used by the WHO collaborative study on brief interventions, including feedback on present drinking, advice on reducing drinking with suggestions on how to do it, a self-help booklet, and an open invitation for a follow-up consultation at the earliest convenience Control group (N = 464) received no feedback or intervention
Outcomes	Mean number of drinks (12 g alcohol) in a "usual" week, mean number of drinks consumed last week, number of participants usually consuming over the recommended limit (21 units for men, 14 units for women), number of participants binge drinking (defined as 6+ units) at least once a week; all reported separately for men and women Assessed at 12 months.
Funding source	This work was supported in part by grants from Alkoholpolitisk Kontaktudvalg (Danish Ministry and Board of Health), Forskningsfonden (Association of County Councils in Denmark), and Fonden til Laegevidenskabens Fremme
Declaration of interests	Not reported.
Notes	Loss to follow-up: <ul style="list-style-type: none"> ● Intervention group: 204/442 (46%). ● Control group: 165/464 (36%).

Beich 2007 (Continued)

<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation not described.
Allocation concealment (selection bias)	Low risk	Screening questionnaires went into a sealed envelope which had a symbol on the front which had to be uncovered scratch card style (p. 594)
Blinding of treatment providers	High risk	"Blinding was not feasible, either for participants and GPs, or for outcome assessment and statistical analysis" (p. 594)
Blinding of participants	High risk	"Blinding was not feasible, either for participants and GPs, or for outcome assessment and statistical analysis (p. 594)
Blinding of outcome assessment (detection bias) All outcomes	High risk	Blinding was not feasible, either for participants and GPs, or for outcome assessment and statistical analysis" (p. 594)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported in results.

Bernstein 2010

Methods	Parallel group RCT: brief intervention versus standard-assessed active control versus minimally assessed active control ITT: no.
Participants	Setting: USA; paediatric emergency department within level 1 trauma centre. Participants: Aged 14 to 21 years who reported bingeing (\geq five drinks in two hours for males or \geq four drinks in two hours for females), and/or high risk behaviours in conjunction with alcohol use, and/or AUDIT \geq 4 for those aged 14 to 17 years or \geq 8 for those aged 18 to 21 years; screened with adolescent version of AUDIT; excluded if they: <ul style="list-style-type: none"> • were not alert and oriented to person, time, and place; • could not be interviewed separately from accompanying family; • planned to leave area in next three months; • could not provide reliable contact information; • were currently in a residential substance abuse treatment facility; • were in custody or institutionalised;

	<ul style="list-style-type: none"> presented for rape exam or psychiatric evaluation for suicide precautions; or if parents opted out for those aged < 18 years, or could not speak English, Spanish, Haitian Creole, or Cape Verdean Creole. <p>Number randomised = 853; 45% male; 2.0% American Indian/Alaskan, 1.4% Asian, 51.3% Black/African American, 19.2% Hispanic/Latino, 0.5% Native Hawaiian/Pacific Islander, 25.6% White; 25% in employment; 21% attending school, 57% high school graduate or equivalent, 22% dropped out of high school; mean age of total sample not reported</p> <p>Baseline information for total sample not reported.</p>	
Interventions	<p>Participants were screened with a youth and young adult survey incorporating AUDIT; AC (standard assessed control) and I (intervention) groups received an assessment battery (duration not reported) including a 30-day Timeline Followback calendar, Adolescent Injury Checklist, Drinking and Driving Scale, and other more general health-related questionnaires</p> <p>Intervention group (N = 283) received 20 to 30 minute motivational interview from trained peer educators, plus one 5 to 10 minute telephone booster session 10 days post-enrolment</p> <p>Standard assessed control group (N = 284) received the same assessment as the intervention group, and a brief written handout containing advice about alcohol risks, a list of community resources and adolescent treatment facilities, and appointments to return at 3 and 12 months</p> <p>Minimally assessed control group received the same brief written handout as standard assessed control group, and an appointment to return in 12 months</p>	
Outcomes	<p>Mean number of drinking days per month, mean number of drinks per drinking day, mean number of drinks per week, maximum number of drinks per day</p> <p>Assessed at 3 months (intervention and assessed control groups only) and 12 months</p>	
Funding source	Supported in part by NIAAA P60AA13759, NIAAA Youth Alcohol Prevention Center at BU- 2006-2009; funding = USD 2.5 million (direct)	
Declaration of interests	Not reported.	
Notes	<p>Project RAP: Reaching Adolescents for Prevention.</p> <p>Payment: USD 10 at enrolment, USD 35 at subsequent follow-up visits</p> <p>Loss to follow-up at 3 months:</p> <ul style="list-style-type: none"> Intervention group: 81/283 (29%). Assessed control group: 87/284 (31%). <p>Loss to follow-up at 12 months:</p> <ul style="list-style-type: none"> Intervention group: 76/283 (27%). Assessed control group: 75/284 (26%). Minimally assessed control group: 88/286 (31%). 	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Bernstein 2010 (Continued)

Random sequence generation (selection bias)	Low risk	“Randomisation was based on computer-generated lists, blocked to balance assignment after every nine subjects and stratified by age group (14 to 17 and 18 to 21 years)” (p. 892)
Allocation concealment (selection bias)	Low risk	A two-stage assignment was carried out using a double opaque envelope system. The first envelope indicated randomization to either minimal assessment (MAC) or assessed status (I or AC). The second sealed envelope (inside the first) distinguish between assessed controls and those participants eligible for an intervention (AC vs. I) (p. 892)
Blinding of treatment providers	Unclear risk	Appears that peer educators who delivered the intervention only saw intervention participants but this is not clear
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	“Participants were cautioned not to discuss their enrolment procedures or allocation with research assistants who conducted their follow-up” (p. 892)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported, but reasons not given.
Selective reporting (reporting bias)	High risk	Abstinence was recorded as first primary outcome, but this was not reported in results

Bischof 2008

Methods	Parallel group RCT: 'full care' telephone-based brief intervention versus stepped care telephone-based brief intervention versus inactive control ITT: yes.
Participants	Setting: Germany; 84 primary care practices. Participants: Aged 18 to 64 years with average consumption > 20/30 g alcohol per day for women/men within the last four weeks, or regular heavy drinking episodes (“binge drinking”) defined as > 60/80 g alcohol for women/men on at least two occasions within the last four weeks; included if assessed as alcohol dependent; excluded if they had an acute or terminal illness, or severe drug dependence, or no telephone, or did not understand German, or unable to read, or were in alcohol treatment, or reported no alcohol consumption in the last four weeks; screened with AUDIT and LAST (Luebeck

	Alcohol dependence and abuse Screening Test) Number randomised = 408; 68% male; mean age not reported for total sample Baseline characteristics not reported for total sample.	
Interventions	<p>All three groups received a health questionnaire incorporating AUDIT and LAST for screening; this served as baseline assessment</p> <p>Full care group (N = 131) received computerised feedback immediately after assessment; followed by one 30 minute session of brief intervention counselling by telephone by trained psychologists based on motivational interviewing and structured elements of behaviour change counselling straight after assessment; followed by three similar booster sessions at 1, 3 and 6 months</p> <p>Stepped care group (N = 138) received the same computerised feedback as full care immediately after assessment but no intervention at this point; they received a maximum of three 30 to 40 minute brief intervention sessions at 1, 3 and 6 months. If stepped care participants reported a reduction of alcohol consumption below the inclusion criteria for the trial and indicated high self-efficacy to maintain the change, no further contact was made until the 12 month follow-up</p> <p>Control participants (N = 139) received a booklet on health behaviour</p>	
Outcomes	Grams per day of alcohol, percentage meeting binge criteria. Assessed at 12 months.	
Funding source	“Funding for this study was provided by German Federal Ministry of Research and Education grant no. 01 EB 0121. The German Federal Ministry of Research and Education had no further role in study design; in the collection, analysis and interpretation of data; in the writing of the report; or in the decision to submit the paper for publication”, p. 250	
Declaration of interests	Conflict of interest: None.	
Notes	<p>SIP (Stepped Interventions for Problem drinkers)</p> <p>Loss to follow-up:</p> <ul style="list-style-type: none"> • Full care group: 15/131 (11%). • Stepped care group: 6/138 (4%). • Control group: 13/139 (9%). <p>We contacted the authors, who supplied missing data.</p>	
Risk of bias		
Bias	Authors’ judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	“Randomisation of subjects to one of the three groups was carried out prior to the telephone interview using sealed cards drawn by study staff from one container” [analogous to card shuffle described in <i>Cochrane Handbook</i> as low risk of selection bias] (p. 245).

Bischof 2008 (Continued)

Allocation concealment (selection bias)	Low risk	Final study participants were randomly allocated, using coloured cards in sealed, non-transparent envelopes (drawn envelopes were not put back), to either the control group or to one of two intervention groups (p. 245)
Blinding of treatment providers	High risk	“Due study design, blinding could not be conducted for the individual study participant and the staff member providing the counselling session” (p. 245)
Blinding of participants	High risk	“Due to our study design, blinding could not be conducted for the individual study participant and the staff member providing the counselling session” (p. 245)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	“A blinded personal interview was conducted by study staff who had no contact with the participant prior to the outcome assessment which mainly was assessed by telephone” (p. 247)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported in results.

Blow 2006

Methods	Parallel group RCT: brief intervention and tailored booklet versus brief intervention and generic booklet versus tailored booklet only (active control) versus generic booklet only (active control) ITT: yes.
Participants	Setting: USA; emergency department level 1 trauma centre. Participants: Aged ≥ 19 years and injured, displaying at risk or heavy episodic drinking. At risk drinking was defined as: <ul style="list-style-type: none"> • ≥ 15 drinks per week for men aged < 65 years; • ≥ 12 drinks per week for women aged < 65 years and men aged ≥ 65 years; and • ≥ 9 drinks per week for women aged ≥ 65 years in the previous 3 months. Heavy episodic drinking was defined as <ul style="list-style-type: none"> • ≥ 5 drinks/occasion on > 4 occasions in the last month (i.e. weekly episodes) for men < 65 years; • ≥ 4 drinks per occasion on > 4 occasions in the last month for men ≥ 65 years and all women.

	<p>Excluded if:</p> <ul style="list-style-type: none"> • severely injured (e.g. unconscious); • in need of immediate life-saving procedures; • blood alcohol level > 200 mg/dL; • self-inflicted injury, sexual assault, overdose, poisoning, near-drowning, chronic injury without specific event associated with re-injury; • pregnant; • prisoner; or • non-English speaking. <p>Screened with computerised health survey incorporating AUDIT-C and DrInC; participants also received an alcohol salivary test before intervention, which proceeded once blood alcohol concentration reached 100 mg/dL or less</p> <p>Number randomised = 575 (but following data are for 494 participants who were followed-up); 71% male; mean age 27.8 years; 86% White, 6% Black, 8% other race; 80% some college education or higher</p> <p>At baseline: mean number of drinks per week = 21.</p>
Interventions	<p>Participants were screened with a health survey incorporating AUDIT-C and DrInC which served as baseline assessment (duration was not reported)</p> <p>Tailored manual/brief advice group (N = 129) received brief intervention before leaving the emergency department from a research social worker, during which the tailored booklet was reviewed with the participant. The tailored booklet was generated according to responses to the computerised screening tool</p> <p>Tailored manual/no brief advice group (N = 121) received a tailored booklet from the research social worker who told them that, based on their responses to the health screen, they scored as at-risk for hazardous drinking and should review the booklet</p> <p>Generic manual/brief advice group (N = 124) received brief intervention before they left the emergency department from a research social worker, during which the generic booklet was reviewed with the participant. The generic booklet was identical to the tailored booklet in length, content and graphics but included standard rather than tailored text/graphics</p> <p>Generic manual/no brief advice group (N = 120) received the generic booklet from the research social worker who told them that, based on their responses to the health screen, they scored as at-risk for hazardous drinking and should review the booklet</p> <p>Duration of intervention was not reported.</p>
Outcomes	<p>Mean number of drinks per week, number of binge drinking episodes (> 5 drinks for men, > 4 drinks for women) in previous month, DrInC score (alcohol-related consequences)</p> <p>Assessed at 3 and 12 months.</p>
Funding source	<p>This study was supported by NIAAA grant AA111629.</p>
Declaration of interests	<p>Not reported.</p>
Notes	<p>Participants willing to complete the survey were entered into a monthly draw for USD 100. Participants were paid USD 20 for three month telephone follow-up interview and USD 30 for 12 month telephone follow-up</p> <p>Loss to follow-up: 81/575 (14%) (not reported according to study arms)</p> <p>Data (including baseline data) were reported only for participants who were followed up</p>

Blow 2006 (Continued)

<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"A computer program automatically determined eligibility for the RCT and randomly assigned participants to one of four intervention conditions" (Blow 2006, p. 570). "To prevent the possibility that staff could unwittingly manipulate assignment to intervention condition, participants were randomly assigned to conditions via the computer using an urn randomisation technique" (Blow 2009, p. 487)
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of treatment providers	Low risk	Cluster randomised - treatment providers only had contact with one arm and only delivered one intervention, reducing the chance of cross contamination
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	High risk	Reported data only from those who completed not those randomised
Selective reporting (reporting bias)	Low risk	Outcomes stated in methods were reported.

Cherpitel 2009

Methods	Parallel group RCT: brief negotiated interview versus assessment only active control versus screening only active control ITT: yes.
Participants	Setting: Poland; emergency department. Participants: Aged ≥ 18 years not presently being treated for an alcohol-related problem, willing to give informed consent to be randomised into one of three groups, willing to provide contact information for at least two individuals who would always know participant's whereabouts, screened as consuming ≥ 11 drinks per week for men (≥ 6 for women); or ≥ 4 drinks on one occasion for men (≥ 3 for women); or ≥ 1 on the RAPS (Remorse, Amnesia, Perform, Starter) tool. Participants who reported drinking an

	average of more than six drinks per day at baseline on both quantity/frequency screening and 30 day timeline follow back were excluded from the analysis Number randomised = 446; 85% male; mean age not reported for total sample Baseline characteristics not reported for total sample.	
Interventions	<p>Participants were screened with the RAPS tool and quantity/frequency questions. The assessed group and brief negotiated interview groups underwent assessment (duration not reported) as follows: blood alcohol concentration measure with a breathalyser, self-reported drinking in the previous six hours using the Timeline Followback, consequences of drinking using the Short Inventory of Problems SIPs +6, stage of change using the Readiness to Change Ruler, risk-taking assessed with questions from Eysenk and Jackson, and sensation seeking assessed with questions adapted from Zuckerman</p> <p>Brief negotiated interview group (N = 147) received a 15 to 20 minute brief negotiated interview from a trained emergency department nurse, which included engagement and permission, feedback, information and norms, decisional balance and pros and cons, readiness to change, menus of options, and prescription for change. A list of AA groups and specialised services for alcohol treatment and counselling was provided to the participant</p> <p>Assessed group (N = 152) received the same list of AA groups and specialised alcohol services</p> <p>Screened group (N = 147) received no assessment or intervention</p>	
Outcomes	Per cent exhibiting at-risk drinking, per cent with RAPS > 1, drinking days per week, drinks per drinking day, maximum drinks on one occasion, number of negative consequences Assessed at 3 and 12 months.	
Funding source	This research was supported by NIAAA grant R21 AA 016081-01.	
Declaration of interests	Not reported.	
Notes	<p>Loss to follow-up at three months:</p> <ul style="list-style-type: none"> ● Brief negotiated interview group: 26/147 (18%). ● Assessed group: 23/152 (15%). ● Screened group: 12/147 (8%). <p>Loss to follow-up at 12 months:</p> <ul style="list-style-type: none"> ● Brief negotiated interview group: 60/147 (41%). ● Assessed group: 53/152 (35%). ● Screened group 55/147 (37%). 	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Randomisation process not described.

Cherpitel 2009 (Continued)

Allocation concealment (selection bias)	Unclear risk	“Participants were first randomised to the screen-only or assessment condition by the study interviewer, who drew an envelope [does not specify sealed or opaque] with the condition assignment. The envelope of those receiving an assessment contained a second envelope, which was opened by the interviewer following assessment to determine whether the participant was assigned to the intervention condition” (p. 984)
Blinding of treatment providers	Low risk	Nurse who delivered the intervention only saw intervention participants
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Assessment interviewer was blind to group status (p. 986).
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	RAPS+1 not reported at three months, but because we focused on 6 and 12 month outcomes, there was little impact

Crawford 2004

Methods	Parallel group RCT: brief intervention versus active control. ITT: yes.
Participants	Setting: UK. Participants: Aged ≥ 18 years, English speakers, resident in greater London, alert and oriented, consuming > 8 units in any one session at least once a week for men or > 6 units for women, or believing their attendance at accident and emergency is related to alcohol; excluded if already in contact with alcohol services, or requesting help with alcohol problems; recruited from accident and emergency; screened by PAT. Number randomised = 599; 78.1% male; mean age = 44 years (range 18 to 90 years). At baseline: mean units consumed during drinking session = 21.2
Interventions	Experimental group (N = 287) received a 30 minute session with an experienced alcohol worker to discuss current and previous drinking in a manner tailored to the subject, plus a health information leaflet containing contact details for national and local alcohol support agencies. Control group (N = 312) received the leaflet only.

Outcomes	Mean weekly units consumed, mean units consumed per drinking day, mean proportion days abstinent all assessed at six and 12 months; mean number of attendances at local emergency department, mean EQ-5D single score both assessed at 12 months only; mean score on GHQ assessed at six months only	
Funding source	“We are grateful to the Alcohol and Education Research Council who funded the study”. (p.1338)	
Declaration of interests	“We declare that we have no conflict of interest”. (p.1338)	
Notes	Loss to follow-up: <ul style="list-style-type: none"> • Experimental group: 98/287 (34%). • Control group: 117/312 (37.5%). 	
<i>Risk of bias</i>		
Bias	Authors’ judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	“Computer generated list of random numbers” (p. 1335).
Allocation concealment (selection bias)	Unclear risk	“Opaque envelopes marked with a unique patient identification number prepared according to randomisation list” (p. 1335)
Blinding of treatment providers	Low risk	Participants re-attended with alcohol health worker to receive brief intervention; therefore, no contamination
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	“Follow-up interviews were done either by telephone or in person by a researcher blinded to allocation status” (p. 1335)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in the methods were reported.

Methods	Parallel group RCT: brief intervention and health information leaflet versus health information leaflet only ITT: yes.	
Participants	Setting: UK, emergency department. Participants: Aged > 18 years presenting after an episode of deliberate self harm; consume > 8 (men) or > 6 (women) units of alcohol in a drinking session at least weekly. Excluded if unable or unwilling to provide verbal consent to participate (non-English language or impaired consciousness), or no fixed address in greater London, or already in contact with alcohol misuse services, or made a specific request for help with alcohol problem at the time of assessment Number randomised = 103; 51% male; mean age = 37.2 years, age range 18 to 65 years At baseline: participants reported drinking up to a mean of 21 units of alcohol per drinking session	
Interventions	Participants were screened with the PAT (Paddington Alcohol Test) which served as baseline assessment Experimental group (N = 51) received approximately 30 minutes of assessment and discussion of current and previous drinking based on the FRAMES approach from an alcohol nurse specialist (ANS). Following the appointment the ANS had the option of referring the participant for further help, such as individual alcohol counselling or detoxification services. The ANS took a patient-centred and non-confrontational approach. Participants also received the information leaflet "Think about Drink", which contains information about the health damaging effects of excessive alcohol consumption, details of benchmarks for recommended limits and contact details for national alcohol help lines Control group (N = 52) received the information leaflet only	
Outcomes	Units of alcohol per drinking session, mean AUDIT score. Assessed at 3 and 6 months.	
Funding source	This study was funded by St Mary's Paddington Charitable Trust	
Declaration of interests	Declaration of Interest: None.	
Notes	GBP 15 shopping voucher. Loss to follow-up: <ul style="list-style-type: none"> • Experimental group: 17/51 (33%). • Control group: 11/52 (21%). 	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	An independent team used random number tables. Regular checks were made to ensure that eligible participants were allocated according to the sequence on the randomisation list (p. 1822)

Crawford 2010 (Continued)

Allocation concealment (selection bias)	Unclear risk	Pre-prepared sealed opaque envelopes contained identical pieces of card - either an appointment card for an alcohol nurse specialist or a blank card (p. 1822)
Blinding of treatment providers	Low risk	Experimental arm was randomised to a separate appointment with alcohol nurse specialist who did not see control group participants, therefore little chance of cross contamination
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"All follow-up data were collected and managed by a researcher who was masked to allocation status" (p. 1823)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Curry 2003

Methods	Parallel group RCT: brief motivational message plus booster telephone calls versus usual care ITT: paper reported multiple imputation to impute outcome data for non-respondents; unpublished data supplied to the reviewers was based on 222/333 (67%) of those randomised
Participants	Setting: USA, primary care clinic. Participants: (no age restrictions), presenting for pre-booked appointment; screened by telephone interviews to select those with AUDIT score ≤ 15 and consuming ≥ 2 alcoholic drinks per day in past month (chronic drinking), or ≥ 2 episodes of binge drinking (≥ 5 drinks) in past week (binge drinkers), or ≥ 1 episode of driving after ≥ 3 drinks; excluded if alcoholic, pregnant, terminally ill, or cognitively impaired. Number randomised = 333; 65% male; mean age 46.9 years; 16% unemployed; 91% post-high school education; 68% income > \$35,000 per year; 80% Caucasian. Number assessed = 222 (67%). At baseline: mean drinking amount = 166 g/week; 42% chronic drinkers; 33% binge drinkers
Interventions	Intervention group (N = 166) received: a) a brief motivational message of 1 to 5 minutes from the primary care physician during the planned routine visit; b) self-help manual; c) written personalised feedback; d) up to three telephone counselling calls over 10 weeks by a psychology graduate Control group (N = 167) received usual care (no intervention or any information about

Curry 2003 (Continued)

	their participation in the study in their notes)	
Outcomes	Drinks per week, drinking days per week, binges per week, grams of alcohol per drinking day, percentage of binge drinkers, percentage of heavy drinkers (average of > 1 drink per day for women or > 2 drinks per day for men Assessed at 3 and 12 months.	
Funding source	This study was supported by National Institute on Alcohol Abuse and Alcoholism Grant RO1 AA09175 (to Susan J. Curry, Principal Investigator)	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: Intervention group: 66/166 (40%). Control group: 45/167 (27%). Analyses of frequency and intensity of drinking are based on unpublished data on 222 cases Analysis of quality of alcohol consumed/week are based on published means and unpublished SDs	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Method of allocation concealment not described.
Blinding of treatment providers	Low risk	Treatment providers only interacted with intervention participants
Blinding of participants	Low risk	During recruitment participants were blinded to the focus of the study on alcohol
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Unclear if outcome assessor was blinded (p. 157).
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported per arm but not reasons. Uneven follow-up between arms; used multiple imputation which uses a regression-type approach that can remove bias that is due to differential non-response if the imputation model contains variables that are good predictors of the outcome and of non-response

Curry 2003 (Continued)

Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.
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Córdoba 1998

Methods	Cluster RCT: brief intervention versus active control; 74 primary care practices were randomised, average cluster size = 6.9. ITT: unclear.	
Participants	Setting: Spain, primary care practices. Participants: Men aged 14 to 50 years with either weekly alcohol consumption over 35 units, or over 10 units on any given day at least once a month (1 unit = 8 g); excluded if alcohol dependency or previous advice to reduce drinking or chronic pathology/treatment requiring > 3 months abstinence or CAGE score > 1; recruited from general practice; screened by lifestyle questionnaire with embedded CAGE Number randomised = 546 (all data reported for 229 followed up participants); 100% male; mean age 36.5 years; 100% Hispanic; 70.1% married; 78.7% middle-lower or lower social status; 95.6% employed; 64.1% further education At baseline: mean weekly alcohol consumption = 54.0 units; CAGE = 1 for 63.2% of participants (data given for 229 heavy throughout-week drinkers only)	
Interventions	Both groups were screened using a lifestyle questionnaire incorporating CAGE questions, this served as baseline assessment Intervention group (N = 104 participants) received from the GP 15 minutes cognitive-behavioural therapy consisting of a self-informative booklet including day diary for registration of consumption, individualised agreement of consumption targets, and offer of follow-up and support Control group (N = 125) received from the GP five minute 'simple advice' which reproduced usual care	
Outcomes	Per cent of participants cutting down to under 35 units per week Assessed at 12 months	
Funding source	Not reported	
Declaration of interests	Not reported	
Notes	EBIAL trial Loss to follow-up not reported by arm	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation is unclear

Córdoba 1998 (Continued)

Allocation concealment (selection bias)	Low risk	All research workers in one centre conducted the same intervention, and they were not aware of the details of the other intervention; allocation unit was the primary care practice (p.563)
Blinding of treatment providers	Low risk	Cluster randomised: treatment providers only had contact with one arm and only delivered one intervention, reducing the chance of cross contamination
Blinding of participants	High risk	No reported attempt to blind participants
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Overall loss to follow-up 49%
Selective reporting (reporting bias)	Low risk	Outcomes described in the methods section are reported

D'Amico 2008

Methods	Parallel RCT: motivational interview versus control. ITT: no.
Participants	Setting: USA; primary care clinic providing free care to underserved populations, described as uninsured (working poor, homeless, runaway and high-risk youth) Participants: 12 to 18 years who reported alcohol consumption or drug use and consequences due to use, English-speaking, able to return for 15 to 20 minute appointment Number randomised = 64 (of which 42 completed and are in the analysis, following percentages are of 42 completers); 48% male; mean age of total sample not reported; 85.7% Hispanic/Latino, 9.5% African American, 4.8% White; 42.9% completed high school, 42.9% didn't complete high school, 14.3% don't know; 31.0% attended at least some college, 52.4% didn't attend at least some college, 16.7% don't know (some are below college age) At baseline: 78.6% reported alcohol use in the last 30 days; 100% reported alcohol use in their lifetime
Interventions	Participants were screened with CRAFFT plus six filler questions on health, and both groups completed a baseline survey (duration not reported) MI group (N = 38 randomised, 22 completers in analysis) received a 15 to 20 minute motivational interviewing intervention plus a 5 to 10 minute booster telephone call one month after intervention from case managers who worked in the mental health division of the clinic. The motivational interviewing session focused on assessing motivation to change (five to seven minutes), enhancing motivation for change (5 to 7 minutes), and

D'Amico 2008 (Continued)

	making a plan (5 to 7 minutes). The booster call briefly reiterated what the teen had discussed in the Project CHAT session, reviewed their goals and whether they had been able to implement any of the strategies they had discussed during the session, and revised goals as necessary Control group (N = 26 randomised, 20 completers in analysis) were screened and assessed; no detail is given about their care
Outcomes	Number of days consumed alcohol in the last month, how many drinks consumed, number of days consumed more than three drinks Assessed at 3 months.
Funding source	Work on this article was supported by a grant from the National Institute on Drug Abuse (R21DA018854) to Elizabeth J D'Amico
Declaration of interests	Not reported.
Notes	Project CHAT. Participants received USD 15 for baseline survey; USD 25 for 3-month survey; USD 15 for completion. During the study incentive was increased to USD 30, and 22 teens received this payment Loss to follow-up: MI group: 22/38 (42%). Control group: 20/26 (23%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"Initially the youth were randomised on a one-to-one basis... as the trial progressed, we recognised that dropout rates were unequal between groups... thus, to maximise power, we altered the allocation schedule such that the probability of being allocated to the intervention was higher." i.e. the researchers changed the randomisation procedure (p. 56)
Allocation concealment (selection bias)	High risk	As above, researchers changed allocation procedure whilst the trial was in progress to alter the allocation
Blinding of treatment providers	Unclear risk	Unclear whether treatment provider has contact with both intervention and control participants
Blinding of participants	Low risk	Screening questionnaire had filler questions to mask recruitment criteria (p. 57)

D'Amico 2008 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not described.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Differential in missing data across groups. This is one of the few studies that did not exclude hard to reach populations such as homeless people
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

D'Onofrio 2008

Methods	Parallel RCT: brief negotiated interview versus scripted discharge instructions ITT: yes.
Participants	Setting: USA, emergency department. Participants: ≥ 18 years who reported consuming in excess of NIAAA limits, or whose emergency department visit related to an injury associated with alcohol use; excluded if non-English speaking, AUDIT > 19, or using drugs daily, or currently enrolled in a substance abuse treatment program, or seeking treatment for an acute psychiatric complaint or hospitalised for a psychiatric problem in the past year, or critically ill or injured, or cognitively impaired Number randomised = 500 (of which 494 in analysis, following percentages are of 494); 68% male; mean age of total group not reported; 67% White, 20% Black, 11% Hispanic, 2% Other; 47% high school or less, 36% some college, 17% college degree or more Baseline characteristics for total sample not reported.
Interventions	Participants were screened with an 18-item health screening questionnaire, including multiple health factors and the NIAAA quantity/frequency questions embedded. Both groups received baseline assessment (duration not reported), including alcohol consumption for the past 30 days as measured by Time Line Follow-Back method, and questions concerning drinking-related consequences such as drinking after driving, injury sustained while drinking, and arrests. Readiness to change was assessed by the Contemplation Ladder, and the Short Form Health Survey was used to assess health status in mental and physical domains BNI group (N = 250 randomised, 247 analysed) received brief negotiated interview from one of a group of trained emergency practitioners - a manual-guided intervention using techniques based on motivational interviewing, brief advice, and behavioral contracting and designed to be delivered in less than ten minutes. The four primary steps were: (1) raise the subject of alcohol; (2) provide feedback by reviewing the participant's screening data, make a connection between alcohol and the visit/illness or injury if possible, review National Institute on Alcohol Abuse and Alcoholism guidelines for low-risk drinking; (3) enhance motivation with motivational interviewing techniques; and (4) negotiate and advise by summarising the participant's reasons for change and negotiating a drinking goal. Participants were then asked to complete and sign a drinking agreement The control group (N = 250 randomised, 247 analysed) received scripted discharge in-

	<p>structions read by the emergency practitioner, designed to be less than one minute in length (although actual mean length was 1.4 minutes). This included a statement recommending that the participant decrease alcohol intake and, if appropriate, use seatbelts, exercise regularly, and stop smoking. A handout was provided with more information related to all identified health risks</p>
Outcomes	<p>Average drinks per week, binge episodes in last month, proportion over NIAA guidelines, number who have driven after drinking, number arrested or pulled over for driving under the influence, number who had motor vehicle crash whilst intoxicated, number who were injured while drinking one or more times, contact with legal system, missed 1 workday in past 30 days Assessed at 12 months.</p>
Funding source	<p>Funded by National Institute on Alcohol Abuse and Alcoholism grant R01 AA12417-01A1 (Dr D'Onofrio), National Institute on Drug Abuse grant K23 DA15144 (Dr Pantalon), and Robert Wood Johnson Generalist Physician Faculty Scholar Award (Dr Fiellin)</p>
Declaration of interests	<p>None (says all conflicts must be reported but none were specified)</p>
Notes	<p>Project ED Health. USD 20 at intake visit (from research associate); USD 40 at six months by post; USD 50 at 12 months by post 95% follow-up at six months; 92% at 12 months.</p>

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Participants were randomly assigned with 100 block randomisation but does not say how the sequence was generated
Allocation concealment (selection bias)	Low risk	"Research associates were provided with sealed, opaque envelopes for each of the 500 randomised study identification numbers. The identification number appeared on the outside, and the assigned treatment condition was specified inside the envelope" (p. 744)
Blinding of treatment providers	Unclear risk	Treatment provider provides both intervention and control, but control statement is scripted and reported duration of control arm suggests contamination is unlikely
Blinding of participants	Low risk	"The screen included questions related to smoking, exercise, and seatbelt use to mask alcohol as the central focus" (p. 744)

D'Onofrio 2008 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	“Interviews were conducted by research associates blinded to subject treatment assignment” (p. 744)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Follow-up rates were high and comparable, but reasons for loss to follow-up not reported
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

D'Onofrio 2012

Methods	Parallel RCT: brief negotiated interview (BNI) plus booster versus BNI versus scripted discharge instructions (SC) versus scripted discharge instructions with no assessment (SC-no assessment) ITT: yes.
Participants	Setting: USA, emergency department. Participants: Aged ≥ 18 years who reported alcohol consumption exceeding the NIAAA's low-risk limits (“hazardous drinkers”) or whose index emergency department visit was related to an injury associated with alcohol use (“harmful drinkers”), screened with NIAA quantity frequency questions; excluded if non-English speaking, or AUDIT > 19, or using drugs daily, or currently enrolled in a substance abuse treatment program, or seeking treatment for an acute psychiatric complaint or hospitalised for a psychiatric problem in the past year, or critically ill or injured, or cognitively impaired Number randomised = 889; 72% male; mean age of total group not reported; 63% White, 22% Black, 13% Hispanic, 2% Other; 40% high school or less, 42% some college, 14% college degree or more (missing data in SC-no assessment group) Baseline characteristics for total sample not reported.
Interventions	Participants were screened with a 17-item health screening questionnaire, including multiple health factors and the NIAAA quantity/frequency questions embedded. The BNI+booster, BNI and SC groups received a baseline assessment (details not given) BNI + booster group (N = 295) received brief negotiated interview from trained emergency practitioners - a manual-guided intervention using techniques based on motivational interviewing, brief advice, behavioural contracting and designed to be delivered in less than 10 minutes. The four primary steps were: 1) raise the subject of alcohol; 2) provide feedback by reviewing the participant's screening data, make a connection between alcohol and the visit/illness or injury if possible, review NIAAA guidelines for low-risk drinking; 3) enhance motivation using motivational interviewing techniques; and 4) negotiate and advise by summarising the participant's reasons for change and negotiating a drinking goal. Participants were then asked to complete and sign a drinking agreement. They received an additional telephone booster intervention at 1 month from a trained primary care nurse: if participants had reduced their drinking to low risk they received reinforcement and relapse prevention; if they were still above low-risk limits the nurse attempted to negotiate a new agreement and consider a change in the future. Coping skills and stressors were assessed and motivational strategies for change were offered

	BNI group (N = 297) received BNI only. SC group (N = 148) received no further screening beyond the general health questionnaire or intervention targeting alcohol use SC-NA group (N = 149) received no assessment or intervention
Outcomes	Mean number of drinks in the past seven days, mean number of binge drinking days in the past 28 days, reported driving after drinking more than three drinks at least once in the past 12 months, Short Inventory of Problems score, Injury Behaviour Checklist score Assessed at 6 and 12 months.
Funding source	Funded by National Institute on Alcohol Abuse and Alcoholism grant 1R01AA14963
Declaration of interests	None (says all conflicts must be reported but none were specified)
Notes	USD 20 at index visit and at the 6 month and 12 month assessments Loss to follow-up at 6 months (for primary outcomes which were collected by interactive voice response): BNI + booster group: 121/295 (41.0%). BNI group: 118/297 (39.7%). SC group: 75/148 (50.1%). Loss to follow-up at 12 months (for primary outcomes which were collected by interactive voice response): BNI+booster group: 126/295 (42.7%). BNI group: 116/297 (39.1%). SC group: 62/148 (41.9%). SC-NA group: 75/149 (50.3%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Participants were randomly assigned with 100 block randomisation but doesn't say how the sequence was generated
Allocation concealment (selection bias)	Unclear risk	"Research associates were provided with sealed envelopes for each of the study identification numbers" (p.183). In the previous study by the same authors (D'Onofrio 2008), researchers were provided with sealed, opaque, numbered envelopes
Blinding of treatment providers	Low risk	Control group were assessed by researcher and then received nothing more; clinicians administered the same intervention to all intervention participants, a research nurse administered the booster; therefore the chances of contamination between arms are

D'Onofrio 2012 (Continued)

		low
Blinding of participants	Low risk	Quantity and frequency questions recommended by the National Institute on Alcohol Abuse and Alcoholism to identify hazardous and harmful drinkers were embedded in a 17-item general health questionnaire. They controlled for social desirability bias by using IVR to collect consumption data (p. 184)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Research associates collecting data were blind to study group assignment, many data were collected by interactive voice response (p. 184)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Daepfen 2007

Methods	Parallel RCT: brief alcohol intervention versus control with assessment versus control without assessment ITT: unclear.
Participants	Setting: Switzerland, emergency department. Participants: ≥ 18 years who were admitted to emergency department for an injury; excluded if history of alcohol-related treatment over the last 12 months, or clinically intoxicated, or medical condition that precluded a face-to-face interview, or if not qualifying as hazardous drinker over the last 30 days Number randomised = 987; 78% male; mean age = 36.7 years. Baseline data not reported for total sample.
Interventions	Participants were screened with a 10-item lifestyle questionnaire including three alcohol-related questions. BAI and CA groups received a 30 minute assessment Brief Alcohol Intervention (BAI) group (N = 310) received a single motivation style brief intervention session lasted approximately 15 minutes and covered six steps: (1) to thank the participant for participation and provide reassurance about confidentiality; (2) to provide feedback about participant's alcohol use compared to similar measures for men and women in the Swiss community and ask the participant's opinion of the feedback; (3) to ask the participant to explore the pros and cons of their alcohol use; (4) to use a 1 to 10 scale to explore participant's importance and readiness to change their drinking patterns; (5) to ask if the participant feels ready to set an objective and provide positive reinforcement about the ability to achieve this objective; and (6) to give each participant written material, including their AUDIT score, drinking pattern percentiles compared

Daeppen 2007 (Continued)

	to the Swiss community and their drinking pattern objectives Control with assessment (CA) group (N = 342) received assessment and “usual care” (not defined) Control no assessment (C) group (N = 335) received “usual care” only
Outcomes	Per cent hazardous drinkers, mean number of days drinking per week in previous 12 months, mean number of drinks per occasion in last 12 months, mean number of binge drinking occasions per month in last 12 months, mean number of drinks in last seven days, mean AUDIT score, mean SF-12 score (mental and physical) Assessed at 12 months.
Funding source	Funded by grant no. 3200-067949 of the Swiss National Science Foundation. The Swiss National Science Foundation is the government agency funding scientific research in Switzerland
Declaration of interests	Not reported.
Notes	Loss to follow-up: BAI: 74/310 (24%). CA: 65/342 (19%). C: 78/335 (23%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	“Each participant was randomised and assigned to experimental or control group based by block of ten participants on a computer-generated allocation method” (p. 1226)
Allocation concealment (selection bias)	Unclear risk	Not described.
Blinding of treatment providers	High risk	“A group of five female and two male research assistants were recruited and selected to do the screening, assessment and BAI at baseline, depending on their ability to apply research procedures and to conduct BAI... Research assistants conducted the motivational style BAI right after the assessment” (p. 1228)
Blinding of participants	Low risk	“One of the goals of the trial was to blind subjects assigned to the control groups to minimize possible intervention effects. The subjects in both control groups were told that the trial focused on general health behaviours, including alcohol use” (p. 1228)

Daeppen 2007 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	“Research assistants at follow up were blinded to the participant assignment, and because this interview conducted no evaluation of the BAI they were unaware from which group the participants came” (p. 1228)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but reasons not reported by arm.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Dent 2008

Methods	Parallel RCT: motivational interview (extended) versus brief intervention versus standard care; only brief intervention and standard care groups used because the motivational interview was not comparable to other trials, and very few participants received the intervention ITT: yes.
Participants	Setting: Australia, emergency department. Participants: Aged ≥ 18 years presenting to emergency department for any reason; excluded if inability or refusal to give informed consent (including lack of communication in English, intoxication preventing coherent answers to questions), or previous enrolment and inability to be followed up (e.g. travellers, no phone contact) Number randomised = 468; 78% male; mean age of total sample not reported Baseline data not reported for total sample.
Interventions	Participants were screened with PAT (Paddington Alcohol Test), and this served as baseline data BI group (N = 159) received a semi scripted informative discussion from a trained nurse or doctor, median five minutes (range 1 to 30 minutes), including the risks of high-risk drinking, a definition of a standard drink, advice on safe drinking, tips on cutting down and incorporated empathy and encouragement. A purpose-designed pamphlet reinforcing the information discussed was also provided SC group (N = 161) received standard care, i.e. no counselling unless the clinician would have accessed addiction services as part of clinical duty or the participant requested counselling MI group (N = 148) received an appointment at a specialist drug and alcohol centre A separate control group (not randomised) was comprised of non-consenting participants who screened positive
Outcomes	Self-reported most drinks in a day, number of participants exceeding daily alcohol limit once per week, emergency department attendances Assessed at 1 and 3 months.
Funding source	An Alcohol Education and Rehabilitation Foundation Grant supported the present study

Dent 2008 (Continued)

Declaration of interests	Not reported.	
Notes	Loss to follow-up: BI group: 83/159 (52%). SC group: 95/161 (59%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer generated block-randomisation process (p. 122).
Allocation concealment (selection bias)	Low risk	Sequentially-numbered, opaque, sealed envelope (p. 122).
Blinding of treatment providers	High risk	"Group allocation was not blinded to the participant, clinicians performing interventions or the researchers conducting follow-up interviews" (p. 123)
Blinding of participants	High risk	"Group allocation was not blinded to the participant, clinicians performing interventions or the researchers conducting follow-up interviews" (p. 123)
Blinding of outcome assessment (detection bias) All outcomes	High risk	"Group allocation was not blinded to the participant, clinicians performing interventions or the researchers conducting follow-up interviews" (p. 123)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Drummond 2009

Methods	Parallel RCT: stepped care brief intervention versus brief advice ITT: yes.
Participants	Setting: Wales, primary care practices. Participants: Aged ≥ 18 years presenting to primary care clinic and screening AUDIT ≥ 8 , may also have diagnosis of alcohol use disorder, or be drinking above Department of Health guidelines (> 21 units per week or > 8 units per day), must live within commuting distance of PC practice, and not have received treatment for alcohol use disorder in

	<p>previous 180 days; excluded if primary drug dependence other than alcohol, or severe mental or physical illness, or severe cognitive impairment or legal issues that might interfere with follow-up</p> <p>Number randomised = 112; 100% male; mean age = 42.7 years.</p> <p>At baseline: mean AUDIT score = 13.7.</p>	
Interventions	<p>Stepped Care group (N = 54) received:</p> <p>Step 1. 40-minute session of behaviour change counselling from a trained practice nurse. Each participant was invited for follow-up with the same nurse 28 days after the initial session. Participants who had consumed > 21 units of alcohol in any one week or > 10 units in any one day during the 28-day period were referred to step 2</p> <p>Step 2. All participants referred to step 2 were offered a maximum of four 50-minute sessions of motivational enhancement therapy conducted by a trained alcohol counsellor. Sessions were held at the primary care practice and scheduled for one per week over a four week period. Each participant was invited for a follow-up with the practice nurse 28 days after the final session. Participants who had consumed > 21 units in any one week or > 10 units in any one day in the 28-day period were referred to step 3</p> <p>Step 3. Participants were referred to the local community alcohol team for specialist intervention. There was no limit on duration or intensity of treatment, which could encompass, where necessary, detoxification, inpatient treatment, outpatient counselling, relapse prevention and drug therapy. The clinical protocol included a caveat that any participant in the intervention group who needed urgent referral to step 3 at any stage could be referred without going through intermediate care steps</p> <p>Minimal intervention group (N = 58) received a five minute directive advice session from a practice nurse addressing the need for them to reduce their alcohol consumption. Participants also received a short self-help booklet outlining the consequences of excessive alcohol consumption and including details on where to seek help locally for alcohol problems</p>	
Outcomes	<p>Total drinks in previous 180 days, mean drinks per drinking day, per cent days abstinent, alcohol problems questionnaire (APQ), quality of life (SF-12), health utility (EQ-5D), mean costs</p> <p>Assessed at 6 months.</p>	
Funding source	<p>The study was funded by the Wales Office for Research and Development. All of the authors are independent of the funding body</p>	
Declaration of interests	<p>Declaration of interest: None.</p>	
Notes	<p>Loss to follow-up:</p> <p>Stepped Care: 15/54 (28%) (52 received step 1, 17 received step 2, 1 received step 3)</p> <p>Minimal intervention: 6/58 (10%) (all received intervention)</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomisation was conducted remote from the research centres by the trial ran-

Drummond 2009 (Continued)

		domisation service at the University of York" (p. 449)
Allocation concealment (selection bias)	Unclear risk	"Block randomisation with seeded random elements was used to minimise the possibility of subversion" (p. 449)
Blinding of treatment providers	High risk	Treatment providers not blinded.
Blinding of participants	Low risk	Participants completed "an AUDIT questionnaire embedded within a general lifestyle questionnaire" (p. 449)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Follow-up was conducted by a researcher masked to the allocated group" (p. 449)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up is reported with reasons. Loss to follow-up is 10% versus 28% but stepped care design means that not all participants received all steps
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Drummond 2014

Methods	Cluster RCT: patient information leaflet versus brief advice versus brief lifestyle counselling, average cluster size = 135 ITT: yes.
Participants	Setting: England, emergency departments. Participants: Aged ≥ 18 , who were screened positive on an alcohol screening test, sufficiently alert and orientated to provide informed consent, living within the catchment area of the ED, and being able to speak, read or write English sufficiently well to complete study questionnaires; excluded if already seeking alcohol treatment, or participating in another study of alcohol interventions, or having sustained a severe injury, or suffering from a serious mental health problem, or grossly intoxicated, or being of no fixed abode; screened by one of three alcohol screening tools: the modified Single Alcohol Screening Question (M-SASQ), or the FAST Alcohol Screening Test, or a modified version of Paddington Alcohol Test (SIPS-PAT) (EDs were randomly assigned to one of the three screening approaches) Number randomised: 1204; 65% male; mean age = 34.6 years, 88% white, 65% continued with education after the age of 16 years, over 45% current smokers At baseline: mean AUDIT score = 12.4 (SD 6.9).
Interventions	Brief Lifestyle Counseling (BLC) group (N = 395) received the patient information leaflet (PIL) and brief advice (BA) from emergency department staff. The following

	<p>day (or as soon as possible thereafter), they received 20 minutes of lifestyle counselling alcohol intervention based on the 'How much is too much?' intervention pack used for the PIL group, delivered by Alcohol Health Workers (AHW) with specialist training and experience in alcohol motivational interventions</p> <p>B) group (N = 403) received five minutes of brief advice about drinking using the SIPS brief advice tool ('Brief Advice About Alcohol Risk') developed for the trial and based on the 'How much is too much?' intervention pack. Following brief advice, the PIL was delivered in the same manner as in the minimal intervention group</p> <p>PIL group (N = 406) received simple clinical feedback using a standard script that their test result indicated they were drinking above the government's 'safe' drinking levels, and were given a PIL - the Department of Health's 'Drinking and You: How Much is Too Much?' leaflet, including information on local alcohol services where further help could be sought by the patient themselves</p>	
<p>Outcomes</p>	<p>AUDIT status (score of < 8 versus \geq 8) on the extended item AUDIT questionnaire; average number of drinks per day using the quantity-frequency questions of the extended AUDIT; alcohol related problems using the Alcohol Problems. Alcohol Screening and Brief Intervention in Emergency Care Questionnaire (APQ) , readiness to change using a modified Readiness Ruler, patient satisfaction using a modified version of the Patient Satisfaction Questionnaire measured (at 12 months only) Assessed at 6 and 12 months.</p>	
<p>Funding source</p>	<p>The study was funded by the UK Department of Health. The Department of Health, proposed the general study design, but the details of the methodology were determined by the study team. The sponsor had no role in data collection, data analysis, interpretation of the results or writing the report. The corresponding author had full access to the data and had final responsibility for the decision to submit for publication</p>	
<p>Declaration of interests</p>	<p>Competing interests: all authors had financial support from the Department of Health in England (Alcohol Policy Unit) for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work</p>	
<p>Notes</p>	<p>Loss to follow-up at 6 months: Patient information leaflet: 114/406 (28%). Brief advice: 99/403 (25%). Brief lifestyle counselling: 128/395 (32%). Loss to follow-up at 12 months: Patient information leaflet: 142/406 (35%). Brief advice: 108/403 (27%). Brief lifestyle counselling: 144/395 (36%). Payment: GBP 10 voucher for baseline assessment.</p>	
<p><i>Risk of bias</i></p>		
<p>Bias</p>	<p>Authors' judgement</p>	<p>Support for judgement</p>

Drummond 2014 (Continued)

Random sequence generation (selection bias)	Low risk	"Randomisation was conducted using a remote secure randomisation service (p. 3)
Allocation concealment (selection bias)	Low risk	Cluster randomised trial.
Blinding of treatment providers	Low risk	Treatment providers not blinded but cluster randomisation means little chance of contamination
Blinding of participants	High risk	Participants were not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Researchers conducting 6 and 12 month follow up were blinded to the participants allocated treatment condition and efforts were made to prevent participants from inadvertently revealing the intervention they received" (p. 3)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported.

Désy 2010

Methods	Parallel RCT: SBIRT versus usual care. ITT: no.
Participants	Setting: USA; emergency department. Participants: ≥ 18 years presenting to emergency department with non life-threatening injuries; excluded due to age, language, level of consciousness or acuity; screened with NIAA quantity/frequency questions and CAGE. Number randomised = 94; 60% male; mean age 39 years, age range 19 to 69 years; 77% White, 13% African American, 9% Hispanic, 1% Other ethnicity; education and employment status not reported. Baseline data not reported for total sample.
Interventions	SBIRT group (N = 49) received one session of 5 to 10 minutes of motivational counselling from an emergency department staff nurse, plus educational brochures and a list of community resources where further intervention and treatment could be sought, plus special discharge instructions that provided resources for follow-up care and community services appropriate to their alcohol use risk level. Control group (N = 42) received usual care.
Outcomes	Mean number of drinks per week, mean drinking days per week, per cent with recurring emergency department visits Assessed at 3 months.

Funding source	This work was supported by a grant from the National Highway Traffic Administration	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: SBIRT group: 23/49 (47%). Control group: 22/42 (52%).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Before the start of the study, subject numbers were computer generated in blocks of 100" (p. 541)
Allocation concealment (selection bias)	Unclear risk	"Before the start of the study, subject numbers were... placed in sequentially numbered sealed enrolment packets" (p. 541)
Blinding of treatment providers	Low risk	"Nurses... were blinded to the group assignment for all study subjects" (p. 541)
Blinding of participants	High risk	Participants were not blinded, screening questions concern alcohol consumption (National Institute on Alcohol Abuse and Alcoholism quantity and frequency questions and the CAGE questionnaire)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up was high compared to other studies, although participants commonly considered 'hard-to-reach' (such as homeless or non-local people) were not excluded
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported.

Methods	Parallel RCT: brief intervention versus control. ITT: unclear.	
Participants	Setting: Spain, recruited from university hospital or urban general practice or rural general practice or industrial occupational health clinic Participants: men between 18 to 65 years with a weekly alcohol consumption of 21 to 95 units (1 unit = 8 g); screened by an evaluation survey with drinking questions embedded with general health questions; excluded if alcohol dependence (defined as alcohol consumption > 95 units per week) or psychiatric disorders. Number randomised = 1022; 100% male; mean age 42.4 years (of those evaluated not randomised). At baseline: mean weekly consumption = 47.1 units; % risk drinkers (> 35 units/wk) = 62%	
Interventions	Intervention group (N = 592) received the evaluation survey (10 minutes) plus a self-help manual containing methods to evaluate their drinking and its effects on their lifestyle, and guidelines for consumption, with an extra 10 minutes of advice and explanation of the manual Control group (N = 430) received the evaluation survey only (10 minutes), with no comment or advice	
Outcomes	Weekly alcohol consumption; per cent at-risk drinkers (> 35 units per week); all reported by setting at six months for the university hospital, urban general practice, and rural general practice, and 12 months for the industrial occupational health clinic	
Funding source	Not reported.	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: Intervention group: 111/592 (19%). Control group: 84/430 (20%). Extracted data for C Urbano and C Rural groups only, others were not based in primary health care. Loss to follow-up for these two groups: Intervention group: 49/255 (19%). Control group: 43/229 (19%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Random allocation was performed by order of entering the study, controlling by age (p. 17)
Allocation concealment (selection bias)	Unclear risk	Not reported.

Díez 2002 (Continued)

Blinding of treatment providers	High risk	Not blinded; no reported attempt to reduce bias.
Blinding of participants	High risk	Not blinded; no reported attempt to reduce bias.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up is reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Ettner 2014

Methods	Cluster RCT: brief intervention plus telephone boosters versus usual care ITT: yes.
Participants	Setting: USA, primary care clinic. Participants: aged > 60 years invited by letter, who had consumed at least one alcoholic beverage in the past three months and was physically and cognitively well enough to participate; excluded if severe cognitive impairment, or terminally ill or deceased, or residing in a skilled nursing facility, or moving out of the area in the next year, or did not speak English, or no longer a patient of the physician, or did not drink alcohol; patients could also be excluded at the discretion of the physician; screened with CARET (an updated and revised version of the Alcohol-Related Problems Survey) Number randomised = 1186; 66% male; 96% White, < 1% Black, < 1% Asian/Pacific, 1.4% American Indian; 3% less than high school, 10% high school graduate, 27% some college, 25% college graduate, 35% graduate degree At baseline: mean drinks per week = 13.6 (SD 8.0).
Interventions	Intervention group (N = 546) received a mailed personalised patient report, an educational booklet on alcohol and aging, a drinking diary to track alcohol consumption, and, depending on the individual patient's reported alcohol-associated risks (as identified on the CARET screening tool), up to 13 'tip sheets' (e.g. on drinking sensibly, sleep, preventing falls and fractures, gout, etc.). Their physician received a copy of the patient report attached to the front of the patient's notes, and used it in the patient's appointment to discuss the patient's drinking and associated risks, and to advise them to reduce consumption if appropriate. Patients then received telephone calls from a health educator who answered questions about the written materials and engaged in the following five steps: (a) assessment and direct feedback; (b) negotiation and goal setting; (c) behavioural modification techniques; (d) self-help-directed bibliotherapy, and; (e) follow-up and reinforcement Control group (N = 640) received usual care, which could include alcohol-related advice

Outcomes	At-risk drinking, drinks per week, recollection of alcohol-related discussions, health care utilisation, costs	
Funding source	This project was funded by National Institute on Alcohol Abuse and Alcoholism (NIAAA) Grant 1RO1AA013990 (Principal investigator: Susan L. Ettner). Alison A. Moore's time was additionally supported by NIAAA Grants R01 AA15957 and K24 AA15957 (Principal investigator: Alison A. Moore). O. Kenrik Duru's time was supported by National Institute on Aging Grants 5P30AG021684-12 and 5K08AG033630-05	
Declaration of interests	Conflict of interest: all authors declared no potential conflicts of interest	
Notes	Project SHARE. Loss to follow-up at 12 months: Intervention group: 107/546 (20%). Control group: 30/640 (5%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"We aimed to match each participating physician to another with the same specialty and clinic site. Physicians from each of the matched pairs were randomly assigned by a statistician who drew random numbers from a uniform [0,1] distribution for the pair; the physician having the lower number was assigned to the intervention group. The patient's treatment assignment was then based on the random assignment of the patient's primary care physician" (p. 448)
Allocation concealment (selection bias)	Low risk	Cluster randomised.
Blinding of treatment providers	Low risk	"Randomisation at the level of participating physicians was used so that physicians would not see patients in both the intervention and control arms of the study, thereby avoiding contamination effects" (p. 448)
Blinding of participants	High risk	Not blinded; no reported attempt to reduce bias.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Research assistants, blinded to treatment allocation, entered all data collected" (p. 448)

Ettner 2014 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up is reported with reasons, but big differential between arms
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported.

Fernández 1997

Methods	Parallel RCT: brief intervention versus control. ITT: unclear.	
Participants	Setting: Spain, general practice. Participants: Men aged 18 to 64 years consuming > 21 International Units per week; screened by alcohol consumption questionnaires; excluded if other drug consumption or psychiatric disorders or previous attendance at specialised alcohol dependence programs Number randomised = 152; 100% male; mean age = 40.3 years; 73.5% employed, 15.9% unemployed, 8.6% retired, 2.0% studying; 1.3% higher education level, 17.8% standard, 78.9% lower	
Interventions	Intervention group (N = 67) received 10 minutes counselling backed up by didactic material Control group (N = 85) received no intervention.	
Outcomes	Number of participants with weekly intake \geq 35 IU at 6 to 18 months; number of participants with weekly intake \geq 21 IU at 6 to 18 months	
Funding source	Este estudio ha sido financiado parcialmente por la ayuda de investigacion FIS 94/0179 [This study was funded partially by the help of FIS 94/0179 research]	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: Intervention group: 29/67 (43%). Control group: 35/85 (41%).	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Block randomisation (p. 128).
Allocation concealment (selection bias)	High risk	Doctor carried out allocation at the appointment (p. 128).
Blinding of treatment providers	Unclear risk	Not clear whether the same provider delivers both intervention and control arms

Fernández 1997 (Continued)

Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The nurse who performed the assessment was unaware of allocation (p. 129)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Field 2010

Methods	Parallel RCT: brief motivational intervention versus treatment as usual + assessment ITT: unclear.
Participants	Setting: USA, trauma unit. Participants: Aged ≥ 18 years who presented with an injury, demonstrated orientation to person, place and time, and identified themselves as white, Hispanic or black, screened with stepwise procedure: (i) clinical indication of acute intoxication or alcohol use or positive blood alcohol concentration; (ii) self-reported drinking six hours prior to injury; (iii) at-risk drinking as per NIAAA guidelines (7 drinks per week for women, 14 drinks per week for men; > 4 drinks per day in men, > 3 drinks per day in women); (iv) positive on CAGE questionnaire, 40 minute assessment procedure; excluded if they spoke neither English nor Spanish, or no identifiable residence, or under arrest or in police custody at admission, or actively suicidal or psychotic, or victims of sexual assault, or their medical condition precluded face-to-face interview, or Glasgow Coma Score ≥ 14 that did not resolve prior to discharge Number randomised = 1493; 82% male; mean age of total sample not reported; 45% White, 19% Black, 36% Hispanic
Interventions	BMI group (N = 737) received brief motivational intervention lasting 15 minutes from a trained health educator; the primary components consist of acknowledging the patient's responsibility for changing drinking, encouraging the patient to explore the pros and cons of drinking, assessing importance, confidence and readiness to change drinking behaviour, reinforcing the patient's sense of self-efficacy and providing support for any efforts or intention to quit drinking or reduce harm associated with drinking, including injury TAU + group (N = 756) received treatment as usual, which included referral to a drug and alcohol counsellor or other appropriate hospital services, and a patient handout regarding the effect of alcohol
Outcomes	Volume consumed per week, maximum amount consumed per occasion, percentage days abstinent, percentage days heavy drinking Assessed at 6 and 12 months.

Field 2010 (Continued)

Funding source	This work was supported by a grant (R01 013824; principle investigator: R. Caetano) from the National Institute on Alcohol Abuse and Alcoholism to the University of Texas School of Public Health. The lead author would like to acknowledge the support of the NIH Health Disparities Loan Repayment Program funded by the National Center of Minority Health and Health Disparities	
Declaration of interests	Declarations of interest: none.	
Notes	MARIA study USD 25 for baseline assessment and USD 50 for 6 month and 12 month follow-up assessments Loss to follow-up: BMI: 317/737 (43.0%). TAU+: 309/756 (40.9%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomised... using a permuted block design (block size 6) to ensure approximately equal distribution of patients according to their race/ethnicity" (Field 2010, p. 65)
Allocation concealment (selection bias)	Low risk	"Study clinicians were blinded to patient randomisation prior to completion of baseline assessment. Treatment assignment was generated offsite and was provided to study clinicians in sealed opaque envelopes" (Roudsari 2009, p. 287)
Blinding of treatment providers	High risk	Although study clinicians were blinded to patient randomisation prior to completion of the baseline assessment, they then had contact with both arms (Field 2010, p. 65)
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Research staff blind to treatment assignment conducted follow-up assessments" (Field 2010, p. 66)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	High risk	Authors specified typical quantity consumed as a primary outcome but did not

report these data

Fleming 1997

Methods	Parallel RCT: brief physician advice versus health booklet control ITT: only in 1 sub analysis paper.
Participants	Setting: USA. Participants: patients with regularly scheduled physician appointments, aged 18 to 65 years, consuming > 14 drinks per week for men and > 11 drinks per week (1 drink = 8 g) for women; screened by health screening survey provided by reception personnel containing four sets of parallel questions on exercise, smoking, weight and alcohol use; excluded if pregnant, under 18, over 65, had attended an alcohol treatment program in previous year, reported symptoms of alcohol withdrawal in previous year, advice in previous three months from GP to change alcohol use, consumption > 50 drinks per week, reported symptoms of suicide. Number randomised = 774; (following numbers are for those assessed): 62% male; 92% White, 1% Hispanic, 4% African American, 3% other; 41% high school or less, 40% some college, 20% college degree or more. At baseline: mean consumption in previous seven days = 19.0 drinks; number of binge drinking episodes (defined as > 5 drinks for men or > 4 drinks for women on one occasion) in previous 30 days = 5.5; binge drinkers in previous 30 days = 78%; excessive drinkers (defined as > 20 drinks per week for men or > 13 drinks per week for women) in previous 7 days = 43%
Interventions	Intervention group (N = 392) received booklet on general health issues, and scheduled to see their personal physician for two 15 minute advice sessions (one brief intervention and one reinforcement session) one month apart, and a workbook containing feedback regarding current health behaviours, review of prevalence of problem drinking, adverse effects of alcohol, worksheet on drinking cues, drinking agreement and diary cards (based on MRC trial). Participants received a follow-up telephone call from the clinic nurse two weeks after each meeting with GP. Control group (N = 382) received a health booklet on general health issues, instructed to address health concerns in their usual manner
Outcomes	Mean drinks in previous seven days; binge drinking (defined as > 5 drinks for men or > 4 drinks for women on one occasion); excessive drinking (defined as > 20 drinks per week for men or > 13 drinks per week for women); assessed at 6 and 12 months (further paper gives 48 month data); health care utilisation (hospital days and emergency department visits), changes in health status (smoking status, depression, motor vehicle crashes, unintentional injuries)
Funding source	This work was supported by grant AA 08512-01A from the National Institutes of Health, National Institute on Alcohol Abuse and Alcoholism, and was assisted by the Wisconsin Research Network, Madison, and the Wisconsin Institute of Family Medicine, Milwaukee
Declaration of interests	Not reported.

Fleming 1997 (Continued)

Notes	<p>TrEAT trial Participants paid USD 50, practices paid USD 300. Loss to follow-up: Intervention group = 39/392 (10%). Control group = 12/382 (3%). Separate papers on 48 month data; cost-benefit analysis; subgroup analysis of women of childbearing age; subgroup analysis of young adults</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer generated random sequence (p. 1040).
Allocation concealment (selection bias)	Low risk	Physicians were not told which of their patients were in control group
Blinding of treatment providers	Low risk	Researchers carried out baseline assessment; physicians (treatment providers) only saw the intervention participants and were not told which patients were in the control group, therefore little chance of contamination
Blinding of participants	Low risk	"The survey was designed as a general lifestyle questionnaire to... minimise the intervention effect of the alcohol questions" (p. 1040)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up included telephone interview by researcher not assigned to subject's clinic (p. 1040)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up 10% in intervention group and 3% in control group
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported.

Fleming 1999

Methods	Parallel RCT: physician delivered counselling versus health booklet ITT: yes.
Participants	Setting: USA, primary care clinic. Participants: patients > 65 years with regularly scheduled appointments at community

	<p>based primary health care clinics; consuming > 11 drinks for men or > 8 drinks (1 drink = 8 g) per week for women, or ≥ 2 positive responses to CAGE, or binge drinking (defined as ≥ 4 drinks per occasion for men two or more times in the last three months, or ≥ 3 drinks per occasion for women); screened using modified Health Screening Survey; baseline assessment 30 minute face-to-face interview in primary care clinic conducted by trained researcher; excluded if attendance at an alcohol treatment programme or reported symptoms of alcohol withdrawal in the last year, or physician advice received in previous three months to change alcohol use, or consumption > 50 drinks per week, or reported thoughts of suicide.</p> <p>Number randomised = 158; 66% male; age range = 65 to 75 years; 26% college educated At baseline: mean weekly alcohol consumption = 16.0 drinks; mean binge drinking episodes (defined as ≥ 4 drinks per occasion for men or ≥ 3 for women) in previous 30 days = 3.7; binge drinkers in previous 30 days = 44.9%; excessive drinkers (defined as > 20 drinks per week for men and > 13 for women) in previous seven days = 29.7%</p>	
Interventions	<p>Intervention group (N = 87) received general health booklet, scheduled to see their personal physicians for two 10 to 15 minute sessions one month apart (1 brief intervention, 1 reinforcement session) received follow-up phone call from nurse two weeks after each session and a workbook containing feedback on patients health behaviours, review of problem drinking prevalence, reasons for drinking, adverse consequences of alcohol, drinking agreement, drink diary cards. Used same protocols as Medical Research Council trial and Project TrEAT</p> <p>Control group (N = 71) received a general health booklet only</p>	
Outcomes	<p>Mean drinks in previous seven days; binge drinking episodes (defined as > 4 drinks per occasion for men or > 3 for women) in previous 30 days; percentage of participants binge drinking in previous 30 days; percentage of participants drinking excessively (defined as > 20 drinks per week for men and > 13 for women) in previous seven days</p> <p>Assessed at 3, 6, 12 months.</p>	
Funding source	<p>This work was supported by NIAA grant 3R01 AA08512-0351, the American Academy of Family Physicians, and the Dean Foundation for Health Research and Education</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>GOAL trial Physicians paid USD 250. Loss to follow-up: Intervention group: 9/87 (9%). Control group: 4/71 (6%).</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.

Fleming 1999 (Continued)

Allocation concealment (selection bias)	Low risk	Physicians were not told which of their patients were in control group (p. 379)
Blinding of treatment providers	Low risk	Researchers carried out baseline assessment; physicians (treatment providers) only saw the intervention participants and were not told which patients were in the control group, therefore little chance of contamination
Blinding of participants	Low risk	Patients completed a modified Health Screening Survey with alcohol questions embedded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up telephone interviews done by physician not linked to patient's clinic (p. 379)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up is reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Fleming 2004

Methods	Parallel RCT: brief primary care clinician advice versus usual care ITT: yes.
Participants	<p>Setting: USA, primary care clinics.</p> <p>Participants: Aged 30 to 60 years who were on medication for diabetes or hypertension with CDT level > 2.5%, or ≥ 50 drinks in the previous 30 days for men or ≥ 30 for women; identified by patient database search, screened by medical records review and telephone interview; baseline interview conducted at clinic participating in the study excluded if current symptoms of alcohol withdrawal, or participation in alcohol treatment programme in previous 12 months; recruited from existing study of %CDT test in 8 primary care clinics.</p> <p>Number randomised = 151; 45% male; mean age = 48.7 years; 88% white, 8% black, 4% other; high school or less = 41%, college degree = 21%, technical degree = 18%, advanced degree = 19%.</p> <p>At baseline: current alcohol abuse = 7.3%, current alcohol dependence = 9.9%, mean drinks in previous 30 days = 33.2, percentage heavy drinkers (defined as ≥ 30 drinks in previous 30 days for men or ≥ 25 for women) = 39.1%, mean frequency of binge drinking (defined as ≥ 5 drinks in one occasion for men or ≥ 4 for women) in previous 30 days = 2.6</p>

Interventions	Intervention group (N = 81) received two 15 minute sessions (one month apart) from nurse practitioners or physician assistants and two five minute follow-up phone calls from the office nurse. Sessions (based on TrEAT) followed a scripted workbook reviewing prevalence of problem drinking, adverse effects of alcohol, %CDT test result, drinking diary cards and a drinking agreement in the form of a prescription. Control group (N = 70) received a general health booklet and were told by the researcher to contact the physician with any health concerns	
Outcomes	Mean percentage of heavy drinkers, mean drinks in previous 30 days, mean frequency of binge drinking, proportion of subjects who reduced %CDT Assessed at two months, four months and 12 months with change scores reported	
Funding source	This study was funded by Axis Shield (CT-C0003), by the National Institute on Alcohol Abuse and Alcoholism (R01 AA8512-03) and by grants from the Department of Family Medicine, University of Wisconsin. [Axis Shield: "A pioneering organisation focused on the development and manufacturing of important and innovative in vitro diagnostic tests for use in clinical laboratories"]	
Declaration of interests	Not reported.	
Notes	Loss to follow-up at 12 months: Intervention group: 11/92 (12%). Control group: 5/75 (7%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Allocation concealment not reported.
Blinding of treatment providers	Low risk	Nurse practitioners or physician assistants carried out the brief intervention but had no contact with control patients, so little chance of contamination
Blinding of participants	High risk	Participants not blinded (p. 632).
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Data were collected by a researcher blind to group assignment (p. 632)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up is reported but not reasons.

Fleming 2004 (Continued)

Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.
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Fleming 2010

Methods	Parallel RCT: brief intervention versus health booklet. ITT: yes.
Participants	Setting: USA, college primary care clinics. Participants: ≥ 18 years, presenting at student health centre and reporting > 50 drinks or ≥ 8 heavy drinking episodes in previous 28 days (≥ 5 standard 14 g drinks) for men, or > 40 drinks or ≥ 6 heavy drinking episodes (≥ 4 standard 14 g drinks) for women in the previous 28 days, screened with general lifestyle questionnaire containing alcohol consumption questions, received 30 to 45 minutes assessment; excluded if pregnant, or had attended an alcohol-treatment program in the previous year, or reported symptoms of alcohol withdrawal in the last 12 months, or received advice from their physician in the previous three months to change their alcohol use, or drank more than 200 drinks in the previous 28 days, or reported symptoms of suicide Number randomised = 986; 49% male; mean age 21 years; 91% White; 19% freshman, 17% sophomore, 22% junior, 26% senior, 16% graduate Baseline characteristics for total sample not reported.
Interventions	BI group (N = 493) received two 15 to 20 minute face to face brief intervention sessions one month apart from a trained physician or nurse practitioner. The first reviewed the participant's current health habits and put self-reported alcohol use at the baseline in a normative context of peer drinking rates and consequences, and examined the subject's expectancies of alcohol use. In addition, the clinician discussed the subject's peak BAC and its consequences, pointed to the financial costs associated with abusive drinking and estimated alcohol calories consumed while drinking. The clinician facilitated the subject's contemplation of his/her life goals and how they would be affected by changes in alcohol use. The student's readiness to change was evaluated and a plan to reduce alcohol consumption was formulated. Subjects were offered drinking cards as a way to self-monitor drinking rates. In addition, three work sheets titled 'Identify Reasons to Make Drinking Changes,' 'Strategies to Success' and 'Alcohol and Decision-Making' were offered to students to consider at home. The second visit reviewed drinking since the first visit and discussed work sheets taken home after the first intervention. The clinician explored the subject's discrepancy between goals and drinking behaviour and ambivalence about drinking change, and focused on harms reduction and alternatives to drinking. The student and the clinician agreed on goals for alcohol use, general health and career. Ways to reward oneself for cutting down on drinking were discussed. Participants received a follow-up phone call or email two weeks after the first visit and one month after the second visit. 88% completed all four parts, 8% received one physician visit, 4% received no physician intervention Control group (N = 493) received the same health booklet as the intervention group, plus usual care

Fleming 2010 (Continued)

Outcomes	Mean drinks in previous 28 days, mean heavy drinking days in previous 28 days, mean drinking days in previous 28 days, percentage of subjects with at least one hospitalisation or emergency department visit or urgent care visit or admission to local detoxification unit in previous six months, mean RAPI score (Rutgers Alcohol Problem Index) Assessed at 6 and 12 months.	
Funding source	This research was supported by National Institute on Alcohol Abuse and Alcoholism grant R01 AA014685-01 (Michael F. Fleming, principal investigator); the Child and Family Research Institute, Vancouver, British Columbia, Canada (Elizabeth M. Saewyc, principal investigator); the Michael Smith Foundation for Health Research (Elizabeth M. Saewyc, principal investigator); and the Institute for Population and Public Health, Canadian Institutes for Health Research grant CPP 86374 (Elizabeth M. Saewyc, chair in Applied Public Health)	
Declaration of interests	Not reported.	
Notes	CHIPs (College Health Intervention Projects) study. USD 200 for completing the required procedures. Loss to follow-up: 96% completed either six month or 12 month follow-up; 88% completed both	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised using a computer-generated allocation method (separately for men and women) (p. 26)
Allocation concealment (selection bias)	Low risk	Each clinician had control and experimental participants in his or her practice with no identifiers available to recognise controls
Blinding of treatment providers	Low risk	Researchers carried out baseline assessment; physicians (treatment providers) only saw the intervention participants and were not told which patients were in the control group, therefore little chance of contamination
Blinding of participants	Low risk	"The health screening survey was designed as a general lifestyle questionnaire to increase student acceptance of the research procedures and to minimize the intervention effect of the alcohol questions" (p. 26)
Blinding of outcome assessment (detection bias)	Low risk	Follow-up was carried out by researchers not assigned to the students' clinic, and

Fleming 2010 (Continued)

All outcomes		they were blinded to group status (p. 26)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up is reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Heather 1987

Methods	Parallel RCT: Drinking Reasonably And Moderately with Self control scheme (DRAMS) versus simple advice versus non-intervention control group ITT: unclear.	
Participants	<p>Setting: Scotland, urban teaching primary care practices.</p> <p>Participants: Aged 18 to 65 years with weekly alcohol consumption > 35 units for men or > 20 units for women (1 unit = 8 g), or clinical suspicion by GP of dependence; screened by health questionnaire with alcohol consumption questions embedded. Patients with high consumption or provoking clinical suspicion were given a 10-question DRAMS medical questionnaire by GP from which any positive response indicated eligibility; excluded if the Brief Edinburgh Alcohol Dependence Schedule showed evidence of late dependence, had known liver disease or severe mental illness, were receiving antidepressant medication, were of subnormal intelligence, were dependent on opiate drugs or were pregnant.</p> <p>Number randomised = 104; 75% men; mean age = 36.4 years (range 18 to 64, SD = 12.2)</p> <p>At baseline: mean consumption in previous month = 194.4 units; mean Michigan alcoholism score = 7.2 (SD = 5.9)</p>	
Interventions	<p>DRAMS group (N = 34): DRAMS kit contained a four-page introductory leaflet for GPs, a patient record card for recording patient details, results of blood tests, self-monitored alcohol consumption and a medical questionnaire with a checklist of ten medical complications, adverse social consequences and signs of physical dependence, a two week self-monitoring drinking diary card for use by patient, a 59 page self-help book, a pocket-sized and abbreviated version of the a self-help manual for controlled drinking. Responses to 10-item medical questionnaire were entered on the medical record card. The patient was handed the drinking diary card and asked to fill it in honestly, a follow-up consultation in two weeks was arranged. At follow-up the results of the blood tests and the drinking diary card were reviewed with the patient and if the existence of a drinking problem was confirmed, the doctor advised the patient to try to control the amount consumed. The patient was then introduced to the self help book and encouraged to decide on a realistic plan of action based on measures suggested in the book and using further diary sheets. Additional appointments were made at which the patients medical condition and progress at cutting down were reviewed, using the results of further blood tests</p> <p>Advice group (N = 32): were informed that their drinking could be harmful and were given strong advice to cut down from GP but no specific targets given and no follow-up consultations regarding their alcohol problem were arranged.</p>	

Heather 1987 (Continued)

	Control group (N = 38): doctor explained that the study would involve a blood test and an assessment interview, but made no specific reference to treatment or drinking and arranged no follow-up consultations in connection with their alcohol problem	
Outcomes	Units of alcohol consumed in previous month, units of alcohol consumed in heaviest month of the previous six; control of drinking problems factor score; GGT and mean corpuscular volume, collaterals (individual known to the patient) reports of patients change in drinking Assessed at 6 months.	
Funding source	Not reported.	
Declaration of interests	Not reported.	
Notes	DRAMS trial Loss to follow-up: DRAMS group: 5/34 (15%). Advice group: 2/32 (6%). Control group: 6/38 (16%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Unclear how randomisation sequence was generated.
Allocation concealment (selection bias)	Low risk	Unclear how allocation was concealed.
Blinding of treatment providers	High risk	Treatment providers not blinded (p. 359).
Blinding of participants	High risk	Patients were screened with a general health questionnaire but were told that the study was about the way people's drinking changes over time (p. 359)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The follow-up interviewers were blind to patients' study groups (p. 359)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up completely recorded with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Methods	Parallel RCT: telephone care management and standard care versus standard care ITT: no.
Participants	<p>Setting: USA, primary care clinic.</p> <p>Participants: those who endorsed heavy drinking over the past week (> 21 drinks for men and > 14 drinks per week for women or for men over the age of 65), or any episodes of binge drinking, and had adequate hearing to participate in scheduled telephone assessments. Screened by AUDIT-C; excluded if they reported active suicidal ideation, or had received specialised addiction treatment during the three months prior, or exhibited symptoms of alcohol dependence or repeated use of illicit drugs in the last year, or experienced current delusions or hallucinations, or had current symptoms of post-traumatic stress disorder (PTSD), or a history of mania</p> <p>Number randomised = 146; 98% male, mean age = 57 years, 55% white, 37.4% employed</p> <p>At baseline: mean drinks per day = 3.2 (SD = 2.4), drinking days per week = 6, drinks per drinking day = 5.1, binge days per month = 10 (SD = 9.9)</p>
Interventions	<p>Telephone Care Management group (TCM, N = 68) received standard care and a TCM session at three, six and nine months. The TCM intervention was based on a treatment manual developed during a pilot phase of this study and includes elements of motivational enhancement, decisional balance, education about alcohol misuse, and development of an individualised behaviour change plan. TCM is consistent with the stepped care model. Behavioural health specialists, two nurses trained in motivational approaches to substance use and brief interventions for addictions, maintained regularly scheduled telephone contact to develop a treatment plan, monitor treatment effectiveness and adverse effects, assess and encourage treatment adherence, and offer support and education. The content of individual sessions included alcohol use monitoring, support and education, and individualised education about at-risk drinking and information about common comorbidities (e.g., depressive symptoms). The TCM manual also includes the use of an addiction management algorithm, that is, a telephone adaptation of a brief intervention developed in previous studies. The algorithm provides guidelines for clinical decision making regarding referral to specialty addictions treatment when needed (e.g. following a significant increase in alcohol use) and close collaboration with the patient's physician. Workbooks logging treatment goals and progress were mailed to participants after each session</p> <p>Standard care (SC, N = 71) group received information and brief advice about the risks associated with alcohol misuse and suggestions to decrease alcohol use. Also received a letter informing them of recommended drinking limits and a description of standard drinks, as is standard practice for all patients endorsing heavy drinking during the assessment</p>
Outcomes	Drinks per day, number of days drinking, drinks per drinking day, binges per month, problems due to alcohol (SIP), motivation to change Assessed at 4, 8 and 12 months.
Funding source	Not reported.
Declaration of interests	The authors declare no conflict of interest.

Helstrom 2014 (Continued)

Notes	Loss to follow up not reported by arm. Seven could not be contacted following baseline or withdrew from the study	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Not described.
Blinding of treatment providers	Low risk	Interventionist only had contact with intervention group, so low chance of contamination between groups. The primary care doctor had contact with both groups but was supposed to give the same advice to both, so this did not affect risk of bias
Blinding of participants	High risk	Not blinded; no reported attempt to reduce bias.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not blinded; no reported attempt to reduce bias.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up not described by arm and only reasons given were "unable to follow-up or withdrew"
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Huas 2002

Methods	Cluster RCT: brief intervention versus usual care, average cluster size = 4.8 ITT: unclear.
Participants	Setting: France, general practice. Participants: aged 18 to 65 years who were consuming > 28 glasses per week and on ≥ 5 days per week; screened by MAST and declared consumption of alcohol; excluded if MAST ≥ 3, or history of alcohol dependence, or in treatment for alcohol problems. Number randomised = 541; 100% male; mean age = 51.8 years.
Interventions	Intervention group (number not reported) received 10 minutes intervention focused on reducing alcohol consumption to < 28 glasses per week. Patients with physical or biological symptoms were invited back every three months, otherwise they returned at one year.

Huas 2002 (Continued)

	Control group (number not reported) received usual care.
Outcomes	Mean drinks per week. Assessed at 1 year.
Funding source	Etude realise grace a un contrat CNAM-INSERM, et au soutien des laboratoires Liphia. [Study completed through a CNAM-INSERM contract, and support of Liphia laboratories. CNAM=National Conservatory of Arts and Trades; INSERM = National Institute of Health and Medical Research]
Declaration of interests	Not reported.
Notes	Number of participants assessed = 419, associated with 88 physicians

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported.
Allocation concealment (selection bias)	Low risk	Cluster randomised.
Blinding of treatment providers	Unclear risk	Not blinded; no reported attempt to reduce bias.
Blinding of participants	High risk	Not blinded; no reported attempt to reduce bias.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Israel 1996

Methods	Parallel RCT: Cognitive behavioural counselling (extended) versus brief advice (control) . ITT: no.
Participants	Setting: Canada, primary care practices. Participants: drawn from those attending clinic who answered positively to at least one of four trauma questions and consumed ≥ 90 drinks in previous four weeks (average > 3 drinks per day), or consumed ≥ 5 drinks per day for ≥ 8 days in previous four weeks,

	<p>or CAGE \geq 2; screened by trauma questionnaire followed by alcohol consumption questionnaire followed by CAGE questionnaire; excluded if severe physical dependence on alcohol, or serum gamma glutamyl transferase (GGT) activity > 2 standard deviations above the mean, or treatment for emotional or psychiatric problems, or regular attendance at Alcoholics Anonymous, or current substance abuse (other than alcohol).</p> <p>Number randomised = 105; age range = 30 to 60 years.</p> <p>At baseline (for those assessed): mean alcohol consumption in previous four weeks = 145.2 drinks; serum GGT = 56.9 U/l</p>	
Interventions	<p>Brief counselling group (number not reported) received a pamphlet with guidelines for achieving abstinence or acceptable drinking, were counselled on cognitive behavioural techniques (30 minutes) to achieve abstinence or sensible drinking as their goal. Acceptable drinking was the consumption of 12 drinks per week as a maximum, provided it did not exceed four drinks (three for women) on any single day and provided that not more than one drink was consumed in one hour. Patients in this group were asked to be alcohol-free when retiring at night. Patients were given reactive strips to detect alcohol in saliva when retiring, and asked to keep records of alcohol consumption and presence or absence of alcohol in saliva on retiring. They were advised of changes in their GGT values, and the significance of the test in relation to alcohol consumption was explained. Cognitive behavioural counselling techniques were those developed by Sanchez-Craig et al for self-referred problem drinkers and were delivered by a nurse ratiore recruited for study. Patients also received one 20 minute session every two months for one year, during which patients were retested for GGT before each visit and were informed of any change in values. The nurse further reviewed the strategy to achieve the patient's goal. Participants were informed of their new GGT values and their significance at each session</p> <p>Advice group (number not reported) received advice to reduce their consumption and were handed a pamphlet with guidelines for achieving abstinence or acceptable drinking. They were also informed of their GGT values, and the significance of the test in relation to alcohol consumption was explained</p>	
Outcomes	<p>Mean alcohol consumption in previous four weeks; serum GGT; psychosocial problems Assessed at 12 months.</p>	
Funding source	<p>This work was supported by the National Institute on Alcohol Abuse and Alcoholism (ROI-AAO9331)</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>Loss to follow-up not recorded by treatment arm, 30% overall. Missing data: gender, number randomised to each arm, baseline data for all randomised participants: requested, not available</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Independent process.

Israel 1996 (Continued)

Allocation concealment (selection bias)	High risk	“Nurse opened a sealed envelope” (p. 1445)
Blinding of treatment providers	High risk	Provider not blinded.
Blinding of participants	Low risk	Screening and baseline questionnaires were broad and not focused on alcohol (p. 1445)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up questionnaires were administered “by a research assistant who was not aware of group assignment” (p. 1445)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up not reported by treatment arm > 30% overall
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Kaner 2013

Methods	Cluster RCT: patient information leaflet (PIL) versus PIL plus brief advice (BA) versus PIL plus BA plus brief lifestyle counselling (BLC), average target cluster size 31 ITT: yes.
Participants	Setting: UK, primary care practices. Participants: ≥ 18 years, alert and oriented, resident within 20 miles of the practice, and able to understand English sufficiently to complete study questionnaires, screened with FAST (Fast Alcohol Screening Test) ≥ 3 or positive M-SASQ (Modified Single Alcohol Screening Question); excluded if already involved in an alcohol research study, or specifically seeking help for alcohol problems, or severely injured or unwell, or had a serious mental health problem, or were grossly intoxicated, or had no fixed abode Number randomised = 34 clusters, 756 patients; 62% male; mean age 45 years; 92% White; 55% education post 16 years; 34% degree or equivalent professional qualification At baseline: mean AUDIT = 12.7.
Interventions	The interventions were cumulative. PIL group (N = 251, all received) received simple feedback on their screening outcome and a 16 page patient information leaflet (‘How much is too much?’) describing the effects of alcohol on health and well being, showing the number of units contained in popular alcoholic drinks, and listing two internet help sites, local alcohol treatment agencies, and details of a national Drinkline number BA group (N = 251, 250 received) received PIL plus five minutes of structured brief advice from practice staff based on the ‘How much is too much?’ BI programme. In addition to providing specific details about the health and social consequences of hazardous and harmful drinking, patients were shown a sex-specific graph, which indicated that their drinking exceeded that of most of the population, and a list of benefits that would result from reduced drinking. Thereafter, patients were taken through a menu of techniques to help reduce drinking and asked to consider a personal target for an achievable reduction

	<p>in drinking</p> <p>BLC group (N = 254, 143 received) received PIL plus BA and were asked to make an appointment for a follow-up consultation within two weeks, which comprised a 20 minute session of brief lifestyle counselling from practice staff based on a condensed form of motivational interviewing called health behaviour change. The patients first described their typical drinking day and then rated the importance of changing their drinking and their confidence about changing their drinking on a ten point scale (where a higher number indicated greater importance or confidence and vice versa). The practitioner then worked with these ratings to establish why they were at the current level and how they might be increased to a higher point before eliciting both pros and cons of drinking and finally working through a six step plan to help reduce drinking levels</p>	
Outcomes	<p>Proportions of participants with negative AUDIT < 8 at six months; mean AUDIT scores, experience of alcohol related problems (alcohol problems questionnaire), health utility (EQ-5D), satisfaction, service utilisation, and patients' motivation to change drinking behaviour (readiness to change) as measured by a modified readiness ruler at 12 months Assessed at 6 and 12 months.</p>	
Funding source	<p>EK was part funded by Fuse the Centre for Translational Research in Public Health. Fuse is a UKCRC Public Health Research Centre of Excellence, and funding comes from the British Heart Foundation, Cancer Research UK, Economic and Social Research Council, Medical Research Council, and the National Institute for Health Research. This study was funded by the Department of Health</p>	
Declaration of interests	<p>All authors had financial support from the Department of Health in England (Alcohol Policy Unit) for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work</p>	
Notes	<p>SIPS trial.</p> <p>Practices received GBP 3000 (staged). Screening and brief intervention was incentivised as GBP 1 per participant screened (EUR 1.26 or USD 1.85, all conversions at 2008 exchange rates), GBP 8 per brief advice, and GBP 32 per brief lifestyle counselling.</p> <p>Participants received GBP 10 voucher after baseline assessment and at each follow-up interview</p> <p>Loss to follow-up:</p> <p>BLC group: 51/254 (20%).</p> <p>BA group: 46/251 (18%).</p> <p>PIL group: 61/251 (24%).</p> <p>We contacted the authors, who supplied missing data.</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Secure remote randomisation service" (p. 2).

Kaner 2013 (Continued)

Allocation concealment (selection bias)	Low risk	Cluster randomised so all patients in each practice received the same intervention
Blinding of treatment providers	Low risk	Cluster randomisation meant that treatment providers interacted with only one arm, so risk of contamination low
Blinding of participants	High risk	Screening was by alcohol tools; control group received a leaflet specifically about drinking
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Researchers who were blinded to allocated intervention contacted the participants (p. 3)
Incomplete outcome data (attrition bias) All outcomes	Low risk	To assess the impact of missing data on the primary outcome, we carried out multiple imputation using the ICE procedure in STATA
Selective reporting (reporting bias)	Unclear risk	Protocol specifies proportion of patients drinking within recommended levels; outcome paper says negative AUDIT

Kunz 2004

Methods	Parallel RCT: brief intervention versus health information pack. ITT: unclear.
Participants	Setting: USA, emergency department. Participants: English or Spanish speaking, aged > 18 years with CAGE score ≥ 1 , having used alcohol in previous 12 months; excluded if they received alcohol counselling in previous year, or signs of cognitive impairment, or physically unable to take part as a consequence of medical treatment, or in police custody. Number randomised = 294; (following data are for those assessed), 81% male; mean age = 41.7 years; 70% African American, 30% Hispanic; 44% high school or more education At baseline (for those assessed): mean weekly consumption in previous three months = 34.1 drinks; binge drinkers in previous three months (defined as > 6 drinks on one occasion for men and > 4 for women) = 92%; mean AUDIT score = 20.1; injured as a result of drinking in previous three months = 27%
Interventions	Intervention group (N = 151) received action plans from researchers according to their self-reported levels of readiness to change: seek more information about drinking, think more about negative consequences of drinking, or lower their drinking per day, per week and per occasion. Participants received a copy of their action plan, a packet of health information and a reminder about a follow-up session. Control group (N = 143) received the packet of health information only

Kunz 2004 (Continued)

Outcomes	Mean weekly alcohol consumption; percentage binge drinkers in previous month, AUDIT score Assessed at 3 months.	
Funding source	Research for this study was supported by grants R01 AA13167 and U24 AA11899 from the National Institute of Alcohol Abuse and Alcoholism (NIAAA) and the National Center for Minority Health Disparities, National Institutes of Health	
Declaration of interests	Not reported.	
Notes	Participants received USD 10 for baseline and USD 20 for follow-up Loss to follow-up: Intervention group: 61/151 (40%). Control group: 39/143 (27%). Missing data: baseline data for all randomised participants: requested, no reply. Screening, baseline and follow-up questionnaires were not masked as in TrEAT (Fleming 1997) and GOAL (Fleming 1999) trials (participant blinding)	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No information available on sequence generation.
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of treatment providers	High risk	Treatment providers not blinded.
Blinding of participants	High risk	Screening, baseline and follow-up questionnaires were not masked (p. 365)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Patients assessed initially by one health promotion advocate were re-interviewed by another at 3 month follow-up (p. 365)
Incomplete outcome data (attrition bias) All outcomes	High risk	Data (baseline and follow-up) reported only for those not lost to follow-up
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported.

Methods	Parallel RCT: brief intervention versus equal attention nutrition control ITT: yes.
Participants	Setting: Kenya, drop-in centres (DIC) for AIDS, Population, Health, and Integrated Assistance (APHIA) II project Participants: women who self-reported being female sex workers, were registered as an APHIA program participant, aged ≥ 18 years, lived in Mombasa and planned to reside there for the next 12 months, moderate risk drinkers defined as those who scored 7 to 19 on AUDIT, had a laboratory-confirmed negative result for gonorrhoea, chlamydia, and trichomoniasis at enrolment; excluded if < 7 on AUDIT Number randomised = 818. Baseline characteristics not reported for whole population.
Interventions	Intervention group (N = 410) received one-on-one sessions lasting 20 minutes on average from nurse counsellors; based on the WHO Brief Intervention for Alcohol Use and contained elements of stages of change and social cognitive health behaviour change theories. Motivational interviewing techniques, focused on goal-setting and increasing self-efficacy for changing behaviour, the provision of positive feedback and encouragement for change, and use of counselling noted and noting stage of change for alcohol reduction were essential intervention elements. In the first counselling session that occurred at the enrolment visit, all participants were presented with their AUDIT screening results. Sessions then included discussion of risks and consequences from drinking, soliciting participants' commitment to reduce drinking, identifying the goal of reduced drinking or abstinence, developing a habit-breaking plan, discussing high-risk situations and coping strategies, and providing feedback and encouragement. Nurse counsellors used a flip chart that included locally-designed illustrations such as physiological consequences of alcohol use and depictions of risky situations relevant to female sex workers, and recorded notes from each counselling session on a data, assessment, and plan form, to track discussions across sessions and record the date for the next session Equal attention nutrition control group (N = 408) received monthly 20-minute sessions from the nurse counsellors, based on Kenyan National Guidelines on nutrition and HIV/AIDS. Sessions included assessment of women's nutritional status, addressed nutritional needs for women and their children and other key groups including people living with HIV and/or taking antiretroviral medication, and included development and monitoring of a nutrition care plan
Outcomes	AUDIT and behavioural interview. Assessed at 6 and 12 months.
Funding source	Supported by the Public Health Evaluation (PHE) component of the President's Emergency Plan for AIDS Relief (PEPFAR), PHE #KE09.0235. Funding was provided through the US Agency for International Development (USAID), under the terms of AID-623-A-11-00007
Declaration of interests	The authors have no conflicts of interest to disclose.
Notes	Loss to follow-up: Intervention group: 37/410 (9%). Control group: 29/408 (7%). No direct payment but could receive condoms.

<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"A statistician not otherwise involved in the study generated the randomization sequences using the random function RANUNI in SAS" (p. 447)
Allocation concealment (selection bias)	Low risk	Statistician produced written assignments sealed in individual tamper-evident opaque envelopes. "The envelopes were fully protected until the site coordinator confirmed the prospective participants' eligibility, obtained written informed consent, and collected all baseline data" (p. 447)
Blinding of treatment providers	High risk	"Given the nature of the intervention, study participants and site staff could not be masked to treatment allocation" (p. 447)
Blinding of participants	High risk	"Given the nature of the intervention, study participants and site staff could not be masked to treatment allocation" (p. 447)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Study investigators and analysts were masked until data handling and analysis decisions were finalized (p. 447)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported; only reason given was "missed appointment"
Selective reporting (reporting bias)	Unclear risk	Had intended to report AUDIT at all time points but do not. They explain that this is because of overlaps in reporting due to time frame component of AUDIT. Therefore alcohol frequency and binge frequency responses from interviews are reported instead

Lane 2008

Methods	Parallel RCT: brief intervention versus screening only control ITT: yes.
Participants	Setting: Australia, public sexual health clinic. Participants: Aged > 16 years, ≥ 8 on AUDIT, or AUDIT-3 = 3 or 4; excluded if they had a language or literacy problem, mental health problem or were too sick to participate Number randomised = 184. Baseline AUDIT score: 8-12 = 99 (54%), 13-19 = 58 (31%), $\geq 20 = 27$ (15%)
Interventions	Brief intervention group (N = 87) received 5 to 10 minutes advice, guided by the Drink-less handy card Control group (N = 97) received screening only (AUDIT and basic demographical information)
Outcomes	AUDIT score, questions relating to any changes in alcohol consumption, having any recent treatment for alcohol problems, and recall and acceptability of being asked about alcohol and getting advice from a research nurse at their visit to the clinic Assessed at 3 months.
Funding source	The Drug and Alcohol Council, New South Wales Health, provided funding for the study. The funding body had no part in the conduct or design of the design
Declaration of interests	Competing interests: none.
Notes	Loss to follow-up: Brief intervention group: 21/87 (24%). Control group: 30/97 (31%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Pre-coded consent forms in sealed envelopes, but not opaque or numbered
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up data were directly entered into SPSS v.15 by the telephone interviewer who was blind to group status
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but no reasons.

Lane 2008 (Continued)

Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.
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Lock 2006

Methods	Cluster randomised controlled trial: brief intervention versus standard advice, average cluster size = 2 ITT: yes.	
Participants	Setting: UK, general practice. Participants: Aged ≥ 16 years with AUDIT score ≥ 8 for men or ≥ 7 for women; excluded if current major physical or psychiatric illness, or severely alcohol dependent, or severely brain damaged or mentally impaired. Number randomised = 127; 50% male; mean age = 44.1 years; 72% employed, 5% unemployed, 15% retired, 3% students; 3% primary school, 6% some secondary school, 47% completed secondary school, 21% technical or trade certificate, 23% university or tertiary education. At baseline: mean weekly units consumed = 24.6; mean AUDIT score = 10.5	
Interventions	Intervention group (N = 67) received 5 to 10 minute intervention using the 'drink-less' protocol. This involved structured advice including: standard drink units, recommended low-risk consumption levels, benefits of cutting down drinking, tips on helping patients reduce consumption, advice on how to set goals, determine action and review progress. Also received a self-help booklet/diary to take away. Control group (N = 60) received standard treatment comprising nurses advice on cutting down drinking and UK Government Health Education Authority leaflet entitled 'Think About Drink', which contained daily benchmark guides for adult men and women and basic advice on alcohol	
Outcomes	AUDIT score, mean drinks per drinking day, mean units per week, Drinking Problems Index, SF-12 physical health, health related quality of life measured by the SF-12 Assessed at 6 and 12 months.	
Funding source	Funding for this project was provided by an NHS Executive (Northern & Yorkshire) Research and Development Regionally Commissioned Project Grant	
Declaration of interests	Not reported.	
Notes	Loss to follow-up at 12 months: Intervention group: 31/67 (46.2%). Control group: 18/60 (30.0%). Number of patients assessed = 78 in 40 GP practices.	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Lock 2006 (Continued)

Random sequence generation (selection bias)	Low risk	Computer generated allocation (p. 428).
Allocation concealment (selection bias)	Low risk	Randomisation was carried out by a member of the team not involved in recruitment or training, all patients in each practice received the same intervention (p. 428)
Blinding of treatment providers	Low risk	Cluster randomised so that treatment providers interacted with only one arm
Blinding of participants	High risk	Patients knew that alcohol advice was being evaluated (p. 429)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome measurement was done by researchers who were blind to patient's study group (p. 429)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Longabaugh 2001

Methods	Parallel RCT: brief intervention versus brief intervention plus booster versus standard care ITT: yes.
Participants	Setting: USA, emergency department. Participants: English or Spanish speaking, living less than one hour from hospital, aged ≥ 18 years, presenting to emergency department with an injury that did not require hospitalisation; either breath alcohol positive (BAC ≥ 0.003 mg/dl) in emergency department, or reported having ingested alcohol in six hours previous to injury, or AUDIT ≥ 8 ; excluded if homeless, or under arrest, or psychiatric disorders, or previous diagnosis of alcohol dependence or abuse Number randomised = 539; 78% male; mean age = 27 years (SD = 9); 72% white, 14% Latino/Hispanic, 10% black, < 1% Asian, < 1% Native American, 3% other; 72% employed At baseline: AUDIT = 12.8; DrInC lifetime negative consequences score = 15.6; mean self-reported alcohol-related injuries in previous year = 1.6
Interventions	BI group (N = 182) received brief intervention by specially trained clinician: one 40-60 minute session based on motivational interviewing. Began with open-ended questions about the patient's injury and a possible connection between injury and alcohol use. If the patient assessed said that there was no connection between the present injury and alcohol use, the interventionist broadened the focus to include negative effects the patient had

	<p>identified on the DrInC as attributed to drinking. The interventionist listened reflectively, provided affirmations, brief summaries and elicited self-motivational statements. To increase awareness of the pros and cons of the target behaviour, patients were assisted in assessing how their alcohol use compared with a national sample, how much money they spent on alcohol, how many calories they consume because of alcohol use, and positive and negative consequences that may be related to their alcohol use. Patients were assisted in determining whether or not they wanted to change any behaviours and, if so, which behaviours they wished to target. In regard to the target behaviour chosen, patients filled out a decisional balance form indicating the pros and cons of change. Last, with the assistance of the interventionist, the patient completed a change worksheet that listed: (1) "The changes I want to make are"; (2) "The most important reasons why I want to make these changes are"; (3) "The steps I plan to take are"; (4) "The ways other people can help me are"; (5) "I will know my plan is working if" and; (6) "Some things that could interfere with my plan are." The patient was given a copy of the change plan at the end of the session.</p> <p>BIB group (N = 169) received brief intervention (as above) plus a booster session with specially trained clinician 7 to 10 days after BI. Based on motivational interviewing. Participants were encouraged to discuss post-discharge experiences, particularly in regard to the change plan they had made in the first session. Then they were provided with additional information about their use of alcohol in an effort to help strengthen the change plan. Based upon reported post-discharge experiences and discussion during the booster session, patients could alter change plans to reflect their experience since the first session.</p> <p>SC group (N = 188) received the customary care that they otherwise would have received in the absence of the research protocol. This consisted of treatment for their injury by the emergency department staff</p>	
Outcomes	<p>Number of heavy drinking days per week; alcohol related injuries; negative consequences from drinking (DrInC) Assessed at 1 year.</p>	
Funding source	<p>This study was funded by National Institute on Alcohol Abuse and Alcoholism grant 5R01 AA09835; Richard Longabaugh Principal Investigator</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>Loss to follow-up not reported by arm.</p>	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Unclear how randomisation sequence was generated.
Allocation concealment (selection bias)	High risk	Interventionist opened a sealed envelope that revealed treatment assignment (p. 807)

Longabaugh 2001 (Continued)

Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	All data were collected by independent research assistants who were blind to patient treatment assignment (p. 807)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up not reported per arm.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Maisto 2001

Methods	Parallel RCT: brief advice versus motivational enhancement versus standard care ITT: unclear.
Participants	<p>Setting: USA, primary care clinics.</p> <p>Participants: Aged ≥ 21 yrs with ≥ 8 on AUDIT, or ≥ 16 standard drinks per week for men or ≥ 12 for women (1 standard drink = 0.6 oz ethanol = 14 g); screened by a lifestyle survey containing general health, stress and alcohol-specific questions including AUDIT; excluded if other drug abuse or dependence, or alcohol withdrawal for previous year, or participation in substance abuse treatment in last year, or unstable psychiatric status and life expectancy < 2 years, or having an active malignancy with the exception of non melanoma skin cancer.</p> <p>Number randomised = 301; 70% men; mean age = 45.6 years; 77% white, 22% black, 1% other; 89% at least high school; 60% employed.</p> <p>At baseline (for previous 30 days): mean days abstinent = 16.3; mean drinks = 75.3; mean number of days consumed one to six drinks = 10.0; mean drinks per drinking day = 5.7; mean Alcohol Dependence Scale score = 5.2</p>
Interventions	<p>Brief advice group (N = 100) received one 10 to 15 minute session from a trained interventionist giving feedback from baseline results and implications for participant's drinking, and advice on goal for reducing or stopping drinking, and a booklet on the effects of alcohol. In this condition, interventionists were trained to focus on delivering feedback to the participant and guiding selection of a drinking goal.</p> <p>Motivational enhancement group (N = 101) received one 30 to 45 minute session, focused on delivery of feedback of the baseline assessment data and the setting of alcohol use goals, and two 15 to 20 minute 'booster' sessions after two and six weeks, which were designed to provide a formal check-up on the participant's progress, from a trained interventionist. Throughout ME sessions, the interventionist used empathy, reflective listening and other techniques to enhance the participant's motivation to change their alcohol use and the participant was encouraged to elaborate and discuss his or her thoughts related to alcohol use and its consequences. The participant was given considerable freedom and time to discuss and select drinking goals. They received the same booklet on the effects of alcohol after the first session.</p>

	Standard care group (N = 100) received no intervention from research interventionists but, as in all other conditions, selected data (the AUDIT score, if positive; alcohol consumption, if high risk; systolic blood pressure if ≥ 200 ; diastolic blood pressure if ≥ 100 ; GGT levels at least five times higher than normal; blood glucose if 350 or higher) from baseline assessments was forwarded to GP who was not discouraged from acting. The inclusion of items on health-related habits other than drinking allowed for the feedback to be placed in a more general health context. Physicians did not receive any instructions about use of interventions for the participant's alcohol use	
Outcomes	Mean days abstinent; mean drinks; mean days consumed one to six drinks; mean drinks per drinking day Assessed at 1, 3, 6, 9 and 12 months.	
Funding source	This research was supported by National Institute on Alcohol Abuse and Alcoholism grant AA1029. Dr Joseph Conigliaro is supported by an Advanced Career Development Award from the Department of Veterans Affairs (VA) HSR&D Service (CD-97324-A) and by a Robert Wood Johnson Foundation Generalist Physician Faculty Scholar Award (#0315000). Dr Conigliaro is also a core member of the VISN-4 Mental Illness Research, Education and Clinical Center and the Pittsburgh Geriatric Research, Education and Clinical Center. Dr Kraemer is supported by a Mentored Clinical Scientist Development Award from the National Institute of Alcohol Abuse and Alcoholism (5K01-AA00235)	
Declaration of interests	Not reported.	
Notes	ELM trial Loss to follow-up: Brief advice group: 26/100 (26%). Motivational enhancement group: 28/101 (28%). Standard care group: 15/100 (15%). Separate paper on elderly as subgroup.	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Project coordinator used random numbers table (p. 606).
Allocation concealment (selection bias)	Low risk	The schedule was kept in an envelope in a locked drawer and was used only by the project coordinator (p. 606)
Blinding of treatment providers	Unclear risk	Physicians saw control patients and interventionists administered interventions so no risk of contamination, but physicians were forwarded details of AUDIT score (if positive) and alcohol consumption (if high) and were not discouraged from advising patients

Maisto 2001 (Continued)

Blinding of participants	High risk	Not blinded. Completed AUDIT for screening therefore alcohol focus clear
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not clear whether outcome assessor was blinded.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but not reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

McDevitt-Murphy 2014

Methods	Parallel RCT: personalised drinking feedback plus motivational interviewing versus feedback only ITT: unclear.
Participants	Setting: USA, Veterans Affairs Medical Centre primary care clinic Participants: combat veterans who screened positive on AUDIT for hazardous drinking; excluded if they had schizophrenia or were in residential substance abuse treatment Number randomised = 68; 91.2% male, mean age = 32.31 years (SD 8.84); 64.7% White, 27.9% Black
Interventions	Motivational interviewing and personalised drinking feedback (PDF) group (N = 35) received PDF in a one hour session where the clinician proceeded through the pages of a PDF packet and used the material to engage the veteran in a motivational interview-style discussion about the content. The session concluded with the clinician querying the impact of this information and engaging the veteran in collaborative goal-setting. The personalised drinking feedback included: deployment related phenomena (personalised feedback about combat experiences and post-deployment adjustment, PTSD and depression, personalised feedback about veterans own symptoms); coping (general information about coping styles, feedback about veteran's own typical coping style); substance misuse; normative feedback based on age-and gender-based norms; psycho-education and personalised feedback about blood alcohol content, binge drinking and risks associated with veteran's own drinking, personal risk factors for alcohol dependence; PTSD-alcohol misuse, information about the "vicious cycle" relationship between PTSD-related symptoms and alcohol misuse, included personalised information about the veteran's motives for drinking Feedback only group (N = 33) received detailed (PDF) packets, provided in written format with 30 minutes to read followed by a chance to ask questions
Outcomes	Mean drinks per week, mean drinking days per week, past month binge days, drinks per drinking day (from TLFB), recent consequences of alcohol use using DrInC Assessed at 6 weeks and 6 months.

McDevitt-Murphy 2014 (Continued)

Funding source	This work was conducted with support from National Institute of Alcohol Abuse and Alcoholism Grant K23AA016120 to Meghan E McDevitt-Murphy, and with support from the Office of Research and Development, Memphis Veterans Affairs Medical Center, and the Tennessee Board of Regents, through the Center for Applied Psychological Research	
Declaration of interests	Not reported.	
Notes	Loss to follow up: MI & PDF group: 3/35 (8.5%). FO group: 2/33 (6%). Payment: USD 50 for baseline, USD 25 for each follow-up.	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but no reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

McIntosh 1997

Methods	Parallel RCT: brief physician advice versus physician delivered cognitive behavioural strategies versus nurse delivered cognitive behavioural strategies ITT: unclear.
Participants	Setting: Canada, family practice clinic. Participants: aged > 15 years who had consumed ≥ 4 standard drinks on any day in the previous month, or responded positively to ≥ 1 CAGE question. Number randomised = 159; 52% male; mean age = 31.1 years; 50% employed full time, 16% employed part time, 34% unemployed

Interventions	<p>Group 1 (N = 40) received two 30 minute sessions two weeks apart from a physician (not their own) using cognitive behavioural strategies and giving advice on sensible drinking, helping participants understand the function of alcohol within their daily activities, and developing a plan and drinking goals. At the first session they received booklets containing this information and diary sheets to record drinking.</p> <p>Group 2 (N = 66) received the same intervention and materials as group 1 but from a nurse ratiore rather than a physician.</p> <p>Group 3 (N = 53) received five minutes of advice from their own family physician based on the patient's initial self-reports of drinking; physicians gave specific information on a standard drink, the limits of moderation, sensible drinking, and avoiding risky situations. Possible connections between drinking and the health concerns that prompted the office visit were identified. This advice was reinforced with a handout</p>
Outcomes	<p>Mean monthly quantity frequency of drinking; problem/problem free drinker (participants were described as problem free if they had no more than one of: insomnia, headache, nausea, cramps, diarrhoea, palpitations, shakiness, sweats, poor memory, poor concentration, mood or personality changes and feeling sluggish)</p> <p>Assessed at 3, 6 and 12 months.</p>
Funding source	<p>This project was funded by the National Health Research and Development Program, Project no 6603-1355-DA. We acknowledge the support of the Drug Dependency Services Division, Department of Health, and the Cape Breton Regional Hospital in Sydney, Nova Scotia</p>
Declaration of interests	<p>Not reported.</p>
Notes	<p>Loss to follow-up: not reported by arm; 1/159 (1%) at 3 months, 11/159 (7%) at 6 months, 16/159 (10%) at 12 months overall.</p> <p>Group 1 and group 2 are aggregated in our analysis as they are an identical intervention</p>

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of treatment providers	Unclear risk	Not blinded but each treatment provider delivered only one kind of intervention therefore limited chance of contamination
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.

McIntosh 1997 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss-to follow-up reported as 10% overall and not by study arm
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Mello 2008

Methods	Parallel RCT: brief telephone intervention versus standard care ITT: yes.	
Participants	Setting: USA, screened in emergency department, intervention delivered at home via telephone Participants: aged ≥ 18 years who were non-critically injured (in the first year only motor crash victims but after that all injured patients), discharged to home, had a telephone, English-speaking, using alcohol at risky levels (defined as NIAAA quantity frequency guidelines, either > 14 drinks per week for men (> 7 for women) or ≥ 5 drinks per occasion for men (≥ 4 for women) screened by research assistant using a health survey during ED visit; excluded if suicidal or in police custody Number randomised = 285; 61% male; 72% White, 87% non-Hispanic; 75% employed Baseline data not reported for total sample.	
Interventions	BI group (N = 140) received 30 minute scripted brief intervention delivered by telephone within five days of emergency department visit by trained counsellors, which aimed to mobilise the participant's own resources to bring about changes needed to reduce alcohol-related risky behaviours. The counsellor's role is to be reflective, to provide an atmosphere that will support and enhance the participant's motivation for change and that will lead the participant to initiate and persist in behavioural change efforts. Two weeks later, they received an additional 15 minute brief intervention booster session. 69% of BI participants received booster SC group (N = 145) received no study interventions.	
Outcomes	Mean change in AUDIT score, binge drinking, impaired driving measures - assessed at three months Difference in AUDIT-C, alcohol related injuries, differences in DrInC (Drinkers Inventory of Consequences) score - assessed at 12 months	
Funding source	The study was funded by the Centers for Disease Control and Prevention's (CDC's) National Center for Injury Prevention and Control (R49/CCR1232280; Mello, principal investigator)	
Declaration of interests	None (says all conflicts must be reported but none were specified)	
Notes	DIAL study. Participants received a total of USD 70 for baseline and follow up assessments Loss to follow-up at three months: BI group: 10/140 (7%). SC group: 2/145 (1%).	

Mello 2008 (Continued)

	Loss to follow-up at 12 months: BI group: 18/140 (13%). SC group: 18/145 (12%).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Participants were randomised by computer assignment to either treatment or standard care group" (p. 45)
Allocation concealment (selection bias)	Unclear risk	Research staff opened an envelope that contained allocation assignment (p. 45)
Blinding of treatment providers	High risk	Not blinded and contamination risk is high because treatment provider talks to all participants on the phone
Blinding of participants	High risk	All patients received briefing that the intervention is about alcohol use and driving behaviours
Blinding of outcome assessment (detection bias) All outcomes	Low risk	All data was collected by a research assistant blinded to treatment condition (p. 45)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but not reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Mertens 2014

Methods	Parallel RCT: brief motivational intervention plus referral list versus usual care plus referral list ITT: no.
Participants	Setting: South Africa, primary health care clinic. Participants: patients aged 18 to 24, who screened positive to heavy alcohol or illicit and non medical drug use with a single question: "In the past year how many times have you had three or more drinks on one occasion" (for women) or "In the past year, how many times have you had five or more drinks on one occasion" (for men); excluded if too ill to participate or no phone Number randomised = 403; (in assessed population) mean age = 21 years; 49% black, 51% mixed race; 8% education grade six or less, 79% some high school, 13% completed high school, 74% unemployed

Interventions	BMI group (N = 206) received brief motivational intervention for alcohol and drug misuse at the Cape Town Drug Counselling Center (CTDCC). The training manual was Rollnick's Health Behavior Change: A Guide for Practitioners (Rollnick et al 1999). Average intervention length was 10 minutes plus referral resource list for drinking and drug use Usual Care group (N = 197) received minimally enhanced usual care plus resource list	
Outcomes	ASSIST score, prevalence of at risk use of alcohol, cannabis, methamphetamine, sedatives, methaqualone and heavy drinking Assessed at 3 months.	
Funding source	Funded by the National Institute on Drug Abuse R21DA022557.	
Declaration of interests	Conflict of interest statement: none declared.	
Notes	Loss to follow-up: BMI group: 16/206 (7.8%). Usual Care group: 24/197 (12.2%). Payment: voucher for cell phone time or a local supermarket, worth ZAR 50 (approximately USD 6)	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not described.
Allocation concealment (selection bias)	Unclear risk	Sealed envelope but not described as numbered or opaque (p. 431)
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Research interviewers conducting follow-up were blinded to randomisation status" (p. 431)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but only reason given is "unable to be contacted"
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Methods	Parallel RCT: motivational intervention plus feedback plus booster versus feedback only ITT: unclear.
Participants	Setting: USA, emergency department. Participants: aged 18 to 24 years who were treated in level I trauma centre and (a) had a blood alcohol concentration (BAC) greater than 0.01% according to a biochemical test, or (b) reported drinking alcohol in the six hours prior to the event that caused their visit, or (c) scored ≥ 8 on AUDIT; screened with mini-mental status examination to assess readiness for consent and assessment; excluded if they did not speak English, had a self-inflicted injury or were in police custody Number randomised = 198; 68% male; mean age of total sample not reported; 65% White, 13% Hispanic, 6% Black, 2% Asian, 2% American Indian, 12% Other or multiple race Baseline characteristics for total sample not reported.
Interventions	MI group (N = 98) received 30 to 45 minutes motivational interviewing in the emergency department from counsellors which incorporated open-ended exploration, personalised feedback, support for self-efficacy and discussion about patients' alcohol use and associated risky behaviours. Session components included establishing rapport, assessing motivation for change, enhancing motivation and establishing goals for change. A worksheet was used to outline the patient's reasons for change and barriers to change. A graphic personalised feedback report was derived from the baseline assessments and included normative information about consumption, summaries of the patient's alcohol-related consequences and indicators of risk, and was discussed in the MI session to enhance motivation. If the patient was interested in discussing change, a goal worksheet containing reduction and cessation strategies was used to facilitate this process. Patients received copies of the worksheets and feedback report, and handouts about alcohol risks and local treatment facilities. The one month booster (20 minutes) started with an assessment (30 day TLFB) of the patient's past-month drinking. The counsellor then reviewed the patient's goals from the first session and inquired about progress. Goals were adjusted or new goals set. For the three month booster (25-30 minutes), patients completed a short assessment battery of alcohol consumption and problem measures described above. A new feedback sheet about the patient's recent behavior relative to their baseline report was generated and reviewed, and a copy mailed to the patient. Progress toward goals since the first booster was reviewed and new goals were generated and discussed FO group (N = 100) received the same baseline assessment and computer-generated personalised feedback report as those in MI, and counsellors spent one to three minutes introducing the report as providing information about how much the patients drink and what happens when they drink, and how they compare to others their age. Conversation with the counsellor was minimal, but any questions or concerns were addressed appropriately. Patients received the same handouts as in MI. At the three month assessment participants were mailed a new feedback sheet
Outcomes	Mean days drinking in past month, mean heavy drinking days in past month, mean drinks per week in past month, RAPI (Rutgers Alcohol Problem Index) score, number of alcohol related injuries in the past year Assessed at 6 and 12 months.

Monti 2007 (Continued)

Funding source	This investigation was supported by research grant AA09892 from the National Institute on Alcohol Abuse and Alcoholism and by a Department of Veterans Affairs Senior Career Research Scientist Award to Peter M Monti
Declaration of interests	Not reported.
Notes	USD 25 gift certificate for participating at baseline, USD 15 cash for 1 month booster, USD 30 for three month booster, \$40 each for six and 12 month follow-ups, \$30 bonus if all follow-ups were completed within one month of due date Loss to follow-up: MI group: 19/98 (19%). FO group: 14/100 (14%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was carried out by the project coordinator using a random numbers table (p. 1235)
Allocation concealment (selection bias)	Unclear risk	Implementation of randomisation sequence unclear.
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Research assistants conducting follow-up were blind to intervention condition (p. 1235)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but not reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Moore 2011

Methods	Parallel RCT: brief intervention versus general health booklet ITT: yes.
Participants	Setting: USA, primary care clinics. Participants: ≥ 55 years, English or Spanish speaking, presenting for any reason to participating clinicians and reporting consumption of at least one alcoholic drink in the previous week; screened with the Comorbidity Alcohol Risk Evaluation Tool (CARET) ; excluded if they could not hear the screening questions, were too ill to participate or

	<p>had received treatment for an alcohol use disorder in the prior three months.</p> <p>Number randomised = 631; 71% male; mean age = 68.4 years, age range 55 to 89 years; 88% White, 9% Hispanic, 3% Other ethnicity; 23% high school or less, 31% some college, 46% college degree or more; 74% retired or homemaker, 26% employed full or part time</p> <p>At baseline: mean 15.2 drinks in previous seven days; 34% had a heavy drinking day in previous seven days (≥ 4 drinks per day); 24% binge drinkers (≥ 4 drinks per occasion)</p>
Interventions	<p>Intervention group (N = 310) received one session of one to five minutes from the primary care provider, followed by three sessions of 40 minutes, 20 minutes and 20 minutes respectively from a health educator. At the baseline visit, the research assistant informed intervention group participants that the study focused on healthy drinking behaviours and they were asked to sign a second informed consent form describing additional procedures. Before seeing the primary care practitioner (PCP), the research assistant handed each participant a personalised report and drinking diary to keep track of alcohol use. A similar version of the report was given to the PCP. The reports outlined participants' alcohol-associated risks identified by the CARET (e.g. drinking two drinks daily and taking a sedative) and potential consequences (e.g. increased sedation and falls). The PCP gave the participant oral and written advice (in prescription-style format on an alcohol education booklet). At two, four and eight weeks after the baseline visit, a health educator called participants to provide additional feedback and counselling using motivational interviewing techniques to help reduce alcohol consumption.</p> <p>Control group (N = 321) received a health information booklet outlining recommended behaviours for alcohol use, nutrition, exercise, medication use and smoking. Research assistants encouraged participants to read the booklet and discuss it with their PCPs</p>
Outcomes	<p>Per cent scoring one to seven on CARET, per cent with one or more heavy drinking days in past seven days, number of drinks in past seven days, CARET risk score range</p> <p>Assessed at 3 and 12 months.</p>
Funding source	<p>This study was supported by R01 AA013937, K23 AA00270, and K24 AA15957 (awarded to Dr Moore) from the National Institute of Alcohol Abuse and Alcoholism, P30AG021684 from the National Institute on Aging (Dr Moore), P30MH082760 from the National Institute of Mental Health (Dr Tang), the Medical Student Training in Aging Program, funded by the National Institute on Aging (T35 AG026736), and the John A. Hartford Foundation (Ms Gill and Ms Chen), Special Fellowship in Advanced Geriatrics, Veterans Affairs Greater Los Angeles Healthcare System (Dr Lin). Data management activities were supported by the UCLA Claude Pepper Older Americans Independence Center funded by the National Institute on Aging (P30AG028748). The sponsor provided financial support for the study only and had no role in the design and conduct of the study; the collection, management, analysis and interpretation of the study; or in the preparation, review or approval of the manuscript</p>
Declaration of interests	<p>Declarations of interest: none.</p>
Notes	<p>Healthy Living As You Age (HLAYA) trial; elderly drinkers.</p> <p>Loss to follow-up at 3 months:</p> <p>Intervention group: 64/310 (21%).</p> <p>Control group: 12/321 (4%).</p>

	Loss to follow-up at 12 months: Intervention group: 88/310 (28%). Control group: 22/321 (7%).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Assignment information was prepared "at the study coordinating center using a computer-generated set of random numbers" (p. 113)
Allocation concealment (selection bias)	Low risk	"Research assistants used consecutively numbered, sealed opaque envelopes" (p. 113)
Blinding of treatment providers	Low risk	Treatment providers were not involved in randomisation or allocation and were alerted only when they had intervention patient (although they had appointments with all participants)
Blinding of participants	Low risk	"To mask the true purpose of the study to potential participants before randomisation occurred, questions on seat belt use, exercise, diet and smoking were asked" (p. 114)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Research assistants, blinded to treatment allocation, conducted all baseline and follow-up outcome assessments" (p. 114)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported with reasons, but large difference between control and intervention group
Selective reporting (reporting bias)	High risk	Mentions re-administering the 7-day alcohol timeline follow-back, questions on self-rated health status, ability to perform strenuous activities (p. 114) but does not report in results

Methods	Parallel RCT: motivational enhancement therapy versus usual care ITT: unclear.
Participants	Setting: Thailand, primary care clinics. Participants: 18 to 65 years presenting to primary care clinic and self-completing AUDIT ≥ 8 ; excluded if alcohol-dependent (according to DSM-IV criteria, as applied by a physician), or history of any liver disease, or history of regular alcohol drinking starting early in the morning, or recent consumption of extremely high amounts per day (> 120 g for men or > 80 g for women), or neurological disease and psychiatric disorders, or pregnant Number randomised = 117; 91% male; mean age = 37 years; 52% primary school or less, 25% secondary school, 7% graduate or higher, 16% unknown At baseline: mean 6.39 drinks per drinking day or 15.15 drinks per week during the previous month; mean 4.75 drinks per drinking day or 11.92 drinks per week during the previous week; mean AUDIT = 17.4; 59% and 43% of participants had hazardous drinking behaviour (men > 4 drinks per day, women > 3 drinks per day) during the previous month and week respectively
Interventions	Motivational Enhancement Therapy (MET) group (N = 59) received three 15 minute sessions of MET from a trained nurse according to project MATCH protocol (a project which aimed to match participant characteristics to their treatment), starting with an evaluation of the patient's ability to change his drinking habits according to the stage of change. For participants in the pre-contemplation stage, the main technique was feedback, using reflection and questioning skills to elicit self-motivational statements. If change was contemplated, the study nurse would work with the participant's ambivalence using a pros and cons technique. If in the determination stage, options on how to reduce drinking behaviour were provided. At the same time, an empathic counselling style and encouragement of the participant's self-efficacy were used to support change in drinking behaviour. Subsequently, each participant's readiness to change drinking behaviour was assessed. If in the determination stage, options on how to reduce drinking behaviour were provided. Participants were then asked to commit themselves to try and change their behaviour and to negotiate appropriate goals. A plan was made with measurable goals in changing drinking behaviour during the action phase. Relapse prevention procedures were used during the maintenance phase Control group (N = 58) received usual care (not detailed).
Outcomes	Mean number of drinks per drinking day during previous week, per cent of participants engaging in hazardous drinking (men > 4 , women > 3 drinks) per drinking day during previous week, mean number of drinks per week during previous week, per cent of participants engaging in hazardous drinking per week during the previous week, mean number of binge drinking episodes during previous week, mean number of times drunk during previous month, GGT Assessed at 6 weeks (i.e. at time of last interview), 3, 6 months
Funding source	This research was supported by grants from the Thai Health Promotion Foundation
Declaration of interests	Not reported.

Noknoy 2010 (Continued)

Notes	Loss to follow-up: MET group: 3/59 (5%). Control group: 7/58 (12%).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomisation... was carried out... using a standard randomisation table" (p. 264)
Allocation concealment (selection bias)	Unclear risk	"Randomisation codes were distributed to each PCU in sealed envelopes" (p. 264)
Blinding of treatment providers	Low risk	Control group complete assessment only and nurse interventionist saw only intervention group so no risk of contamination
Blinding of participants	Low risk	Subjects randomised into the control condition "were told that the trial focused on health behaviours, which included questions on smoking, exercise, eating behaviour, weight and alcohol use" (p. 265)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"The study interviewers at follow-up visits were not aware of the assignment allocation of the study participants" (p. 265)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but not reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Ockene 1999

Methods	Cluster RCT: special intervention versus usual care, average cluster size = 111. ITT: yes.
Participants	Setting: USA, primary care centres. Participants: 21 to 70 years consuming > 12 standard drinks per week or ≥ 5 standard drinks on ≥ 1 occasion in previous month for men, or > 9 standard drinks per week or ≥ 4 standard drinks on ≥ 1 occasion in previous month for women (1 standard drink = 12.8 g alcohol); screened by Health Habits Survey with embedded CAGE; excluded if pregnant, or planning to move away from the area within one year, or did not have telephone, or already participating in alcohol intervention programme, or psychiatric disorder.

	<p>Number randomised = 530; 64.7% male; mean age = 43.9 years; 94.6% White, 5.4% non-white; less than high school level = 8.6%, high school graduate or some college = 51.0%, college graduate or more = 40.4%.</p> <p>At baseline: mean drinks per week = 17.8.</p>
Interventions	<p>SI group (N = 274) participants were told that at their next regularly scheduled appointment their providers probably would discuss one of the health issues that was asked about in their lifestyle interview. They were given the same booklet as the UC participants. The SI providers received 2.5 hours training in the patient-centred alcohol intervention program during which the change in orientation required to work with high-risk drinkers (where the goal may be reduction in drinking) vs alcohol dependent patients (where the goal is abstinence) was emphasised. The SI providers were asked to carry out the brief 5 to 10 minute patient centred alcohol counselling sequence at the time of a regular visit with patients identified as high-risk drinkers. Counselling focused on the number of drinks per week, binge drinking, or both, depending on the participant's problem area (s). The SI providers were instructed to request that the patient set a follow-up visit to review progress. The SI office sites also had a limited office support system designed to assist the busy primary care provider in carrying out the intervention. Although implemented by Project Health RAs, the system was designed to be incorporated easily into usual office procedures and includes the RA affixing the following to the chart of the high risk drinker: the lifestyle interview summary sheet, which reports the participant's alcohol history (drinks per week, history of binge drinking, family history of alcohol abuse); the intervention algorithm to remind the physician of the counselling sequence taught in the training sessions; patient education materials in the form of the tip sheets for the providers' use with patients.</p> <p>UC group (N = 256) received a health booklet that included advice on general health issues and were told to address any health questions with their providers. The UC providers were encouraged to identify and intervene with patients with alcohol-related issues to whatever extent they thought appropriate. All providers were encouraged to attend the weekly conference series in which the approach to the patient with alcohol problems was presented biannually as part of a two year curriculum</p>
Outcomes	<p>Mean drinks per week, mean binge drinking episodes (defined as > 5 drinks on one occasion for men and > 4 for women), change in weekly drinking levels and binge drinking episodes</p> <p>Assessed at 6 and 12 months.</p>
Funding source	<p>This project was supported by grant 5-R01-AA09153 from the National Institute on Alcohol Abuse and Alcoholism</p>
Declaration of interests	<p>The authors have not been engaged in any financial or personal conflicts of interests as a result of the research described in this manuscript</p>
Notes	<p>HEALTH trial</p> <p>Loss to follow-up:</p> <p>SI group: 39/274 (14%).</p> <p>UC group: 46/256 (18%).</p> <p>Number of participants assessed = 445 in 4 practice sites.</p> <p>12 month outcome data reported in separate paper, but baseline data for all randomised</p>

Ockene 1999 (Continued)

	participants reported only in Ockene 1999.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Sequence generated using random number generator in SA statistical software (p. 2199)
Allocation concealment (selection bias)	Low risk	Cluster randomised so all patients in a practice received the same intervention
Blinding of treatment providers	Low risk	Cluster randomised so treatment providers only interacted with one arm
Blinding of participants	Low risk	Alcohol questions were embedded with other health-related questions to minimise the intervention effect of the alcohol questions (p. 2199)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up interviews were conducted by blinded RAs (p. 2199)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow up similar across both groups and reasons reported (only refused or not available)
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Richmond 1995

Methods	Parallel RCT: alcohol screen versus minimal intervention versus no intervention versus no assessment
Participants	Setting: Australia, general practice. Participants: 18 to 70 years consuming > 35 standard drinks per week for men or 21 for women; Screened by MAST and Ph (physical dependence on alcohol) score; no exclusion criteria listed Number randomised = 378; 57% male; mean age = 37.7 years; 74% employed; 67% beyond secondary school. At baseline: mean weekly consumption = 36.8 units.
Interventions	Alcohol screen (AS) group (N = 96) received five consultations: 1) five minutes where patients were given self-help manual and day diary to monitor their consumption; 2) one week later, patients had a 15 to 20 minute consultation where a personalised approach

Richmond 1995 (Continued)

	<p>to patient education using a flip-over unit displaying 12 pictorial and text prompts was used, and patients were counselled about recommended limits, problems associated with excessive drinking, alternate activities; 3) one month later, patients had 5 to 25 minute consultation to reinforce and support new drinking habits; 4) and 5) were both five minute sessions for further support.</p> <p>Minimal intervention (MI) group (N = 96) received five minutes brief advice and self-help manual.</p> <p>No intervention (NI) group (N = 93) received no intervention.</p> <p>No assessment (NA) group (N = 93) received no intervention and no assessments (for comparison with NI group to test whether the assessments make a difference)</p>	
Outcomes	<p>Mean weekly consumption; mean quantity-frequency consumption; GGT</p> <p>Assessed at 6 and 12 months.</p>	
Funding source	<p>The study was funded by the Drug and Alcohol Research and Education Committee of the Department of Health, Housing, Local Government and Community Services as part of the National Campaign Against Drug Abuse and by the National Health and Medical Research Council</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>Alcohol screen trial</p> <p>Loss to follow-up:</p> <p>AS group: 32/96 (33%).</p> <p>MI group: 26/96 (27%).</p> <p>NI group: 30/93 (32%).</p> <p>(NA group was not assessed at 1 year).</p>	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Patients were not randomly assigned but allocated to one of the four study groups in weekly blocks, the order of blocks being randomly determined for each GP (p. 121)
Allocation concealment (selection bias)	Unclear risk	Method of allocation concealment unclear.
Blinding of treatment providers	High risk	GPs delivered both Alcoholscreen and minimal intervention.
Blinding of participants	Low risk	Alcohol questions were disguised among questions relating to weight, smoking and exercise. Study was described to patients as being about lifestyles

Richmond 1995 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Low risk	“Follow-up assessment was carried out at 6 and 12 months... by a research assistant blind to study group” (p. 121)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Rodríguez 2003

Methods	Parallel RCT: brief intervention versus minimal intervention. ITT: yes.	
Participants	Setting: Spain, emergency department. Participants: Aged > 18 years, had been involved in a traffic crash in previous six hours and BAC \geq 0.2 g/L; excluded if unable to speak Spanish, non-resident, very severe medical, psychiatric or social conditions; acute intoxication, patient not sober at discharge; screened by alcohol-on-site saliva test. Number randomised = 85; 88% male; median age = 26 years. Baseline AUDIT-C = 4.9.	
Interventions	BI group (N = 40) received 15 to 20 minute intervention based on FRAMES, model of change and motivational interviewing, and an information leaflet. MI group (N = 45) received five minute empathic advice and the same information leaflet	
Outcomes	AUDIT-C positive or negative, percentage participants who reduced consumption, percentage reduction in hazardous drinkers Assessed at one year.	
Funding source	Esta investigacion esta parcialmente subvencionada por el Fondo de Investigaciones Sanitarias (01/0903) y por el Sevei Catala de Transit [This research was partially funded by the Health Research Fund (01/0903) and by the Catalan traffic service]	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: BI group: 12/40 (30%). MI group: 16/45 (36%). Not included in meta-analysis due to unusable outcome data - discussed narratively	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Rodríguez 2003 (Continued)

Random sequence generation (selection bias)	Unclear risk	Randomly distributed by weeks (p. 193).
Allocation concealment (selection bias)	High risk	Randomly distributed by weeks; predictable sequence.
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-ups were performed blind by phone.
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Romelsjö 1989

Methods	Parallel RCT: brief intervention versus brief advice. ITT: unclear.
Participants	Setting: Sweden, primary care district health centres. Participants: 18 to 64 years, consuming at least 40 g 100% ethanol per day for men and 30 g for women, or drinking in the morning at least every second time when drinking alcohol, or having difficulties restricting drinking at least every second time when drinking alcohol, or CAGE \geq 3, or elevation of GGT; screened by mailed general health questionnaire incorporating alcohol consumption questions and CAGE, and a telephone interview followed by a health examination; excluded if inpatient care for alcoholism or alcohol psychosis in previous three years, or care at an inebriate's institution in previous three years, or other substance abuse in previous three years, or ongoing treatment or need for treatment for a mental disorder, or severe somatic disease, or other potential causes of elevated GGT. Number randomised = 83; 84% male; mean age = 46.3 years (range = 21 to 64 years); 86% employed. At baseline: GGT = 2.0 Ukat/l; daily alcohol consumption = 29.1 g 100% ethanol; 'problems index' = 11.3
Interventions	Intervention group (N = 41) were advised by their GP to cut down their alcohol consumption by the next visit, and if this seemed difficult, to abstain from alcohol. Elevated GGT values and changes in GGT over time were used in discussion with patients in a biofeedback approach. Patients were asked to visit the laboratory again so that the new GGT values were at the GP's disposal at the visit. The GP treated other conditions when necessary, and tried to get a good picture of the patient's social and psychological condition and to give support. The time interval between visits was according to the

	GP's judgement. The mean number of visits to the GP was about three. Control group (N = 42) were told by their general practitioner that they should cut down on alcohol consumption and that a follow-up examination was planned for approximately one year later. They had earlier been informed by the nurse that their alcohol consumption was elevated or, by letter, that their GGT was elevated
Outcomes	Change in GGT; change in self-reported alcohol consumption; change in a combined measure of alcohol problems 'problem index' Assessed at one year.
Funding source	This work was supported by a grant from the Swedish Ministry of Health and Social Affairs, the Commission for Social Research (Project No D84/212:1) and by the Stockholm County Council
Declaration of interests	Not reported.
Notes	Loss to follow-up: Intervention group = 5/41 (12%). Control group = 6/42 (14%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomisation was performed by the project leader according to a randomisation scheme unknown to the GPs; each patient was randomised to reach an equal number of patients from each group per GP. . distribution for all GPs was considered... then the total distribution was considered. .. if this distribution was equal, ultimately the choice was made by coin flipping" (p. 1320)
Allocation concealment (selection bias)	Low risk	Allocation sent in a closed envelope to the GP, which was opened only after the first visit. The rationale for this, which worked well, was that the attitude or behaviour of the GP towards the patient should not be affected by knowledge of allocation (p. 1320)
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	Low risk	It was attempted to conduct the interviews as a rather open talk on health and social questions; alcohol questions were embedded in other health-related questions (p.

Romelsjö 1989 (Continued)

		1320)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessment was carried out in a lab.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but only reasons are “drop-out”.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Rubio 2010

Methods	Parallel RCT: brief intervention plus health leaflet versus health leaflet ITT: yes.
Participants	Setting: Spain, primary care clinic. Participants: 18 to 65 years, AUDIT = 8 to 15 or reported binge drinking (men \geq 5 and women \geq 4 standard drinks per occasion one or more times in the past month); screened with AUDIT; excluded if alcohol dependent (AUDIT > 15), or refused to take part, or pregnant, or planning to move out of the area within the year, or no telephone, or already participating in an alcohol intervention program, or Axis I psychiatric disorder that prevented participation (according to physician's judgement) Number randomised = 752; 65% male; mean age of total sample not reported; 88% employed, 4% unemployed, 8% homemaker; 58% high school or less, 38% some college, 4% college degree or more Baseline data not reported for total sample.
Interventions	BI group (N = 371) received two 10 to 15 minute counselling sessions four weeks apart from the primary care physician using a scripted workbook, which included a review of alcohol-related health effects, a pie chart displaying the frequency of different types of at-risk drinkers, a list of methods for cutting down drinking, a treatment contract, and cognitive behavioural exercises. An office nurse contacted the patients two and eight weeks after the initial counselling sessions to reinforce the face-to-face sessions. Participants also received the same booklet as the control group Control group (N = 381) received a booklet on general health issues and were instructed to address any health concerns in their usual manner
Outcomes	Mean number of binge drinking episodes in the last 30 days, mean number of drinks in previous seven days, mean number of participants binge drinking in last 30 days, mean number of participants reporting excessive drinking in the previous seven days (> 18 drinks per week for men and > 13 for women); all reported separately by gender Assessed at 12 months.
Funding source	Funding: This research was supported by Grants FCM/03 and FCM/04 (Fundacion Cerebro y Mente) and Instituto de Salud Carlos III, Centro de Investigación en Red de Salud Mental, CIBERSAM

Rubio 2010 (Continued)

Declaration of interests	Conflict of Interest: none of the authors have any conflicts of interest associated with the work presented in this manuscript	
Notes	Loss to follow-up: BI group: 31/371 (8%). Control group: 47/381 (12%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomised by a computer generated allocation method (men and women were randomised separately) (p. 73)
Allocation concealment (selection bias)	Unclear risk	Implementation of sequence not reported.
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	High risk	Not blinded. Screened by AUDIT.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Conducted by blinded researchers not assigned to the subject's clinic (p. 73)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Schaus 2009

Methods	Parallel RCT: brief motivational intervention and educational brochure versus educational brochure ITT: yes.
Participants	Setting: USA, university primary care centre. Participants: students ≥ 18 years who reported drinking ≥ 5 (for men) or ≥ 4 (for women) drinks containing alcohol in a row on at least one occasion during the past two weeks, screened with an 11-item health questionnaire including quantity frequency alcohol question; excluded if pregnant, or planning to leave the university within 12 months, or already enrolled in an alcohol or drug treatment program, or had consumed more than 200 drinks in the past 30 days, or had a BAC greater than 0.35 on any day within the past 30 days Number randomised = 363; 48% male; mean age not reported for total sample; 77% White, 5% Black, 11% Hispanic, 2% Asian, 1% American Indian or Alaskan Native,

	<p>3% Other, 1% prefer not to respond; 26% freshman, 21% sophomore, 28% junior, 17% senior, 8% graduate/post baccalaureate</p> <p>At baseline: mean number of days drinking in past 30 days = 8.6, number of days episodic (5/4) drinking = 5.2, mean BAC = 0.08, mean peak BAC = 0.15</p>	
Interventions	<p>BMI group (N = 181, 174 completed intervention) received two 20-minute BMI sessions from health centre staff (physicians, physician assistants, nurses) combining patient-centred MI techniques and cognitive-behavioural skills training, and written individual normative feedback based on the participant's baseline assessment. The focus of the first BMI session was to establish rapport between the participant and the provider by focusing on positive lifestyle concerns of the patient. The second session maintained the student-centred focus of MI and stressed the alcohol skills training components of the BASICS program. At completion of the second BMI session, participants received the participant feedback document for future reference, along with an alcohol prevention educational brochure entitled 'Drinking: What's Normal, What's Not'</p> <p>Control group (N = 182) received the brochure only.</p>	
Outcomes	<p>Mean typical BAC, mean peak BAC, mean number of drinks per sitting, mean number of drinks per week, mean number of times drunk in a typical week, mean peak number of drinks in a sitting, mean number of days heavy episodic drinking (met 5/4) in month, mean RAPI (Rutgers Alcohol Problem Index) score; all in previous 30 days</p> <p>Assessed at 3, 6, 9 and 12 months.</p>	
Funding source	<p>This research was supported by National Institute on Alcohol Abuse and Alcoholism grants 1 U18 AA015673-01 and U01 AA14741</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>USD 30 at baseline, USD 10 each at three, six, nine months; USD 40 at 12 months</p> <p>Loss to follow-up at 12 months: BMI: 70/181 (39%). Control: 57/182 (31%).</p>	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Subjects were block randomised using SPSS to either the control or the intervention group, where the order of the interventions varied randomly within each block (p. 132)
Allocation concealment (selection bias)	Unclear risk	"The group assignment was placed into a sealed envelope by the data manager and was not available to those recruiting subjects until after informed consent was obtained" (p. 132)

Schaus 2009 (Continued)

Blinding of treatment providers	Low risk	Control groups were allocated to health providers who received no training in BI (a different group of people to those who administered brief intervention), so no chance of cross-contamination
Blinding of participants	Unclear risk	Participants filled in a healthy lifestyle questionnaire as well as alcohol questions at screening, but all participants received an alcohol-related educational brochure (p. 132)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Participants completed follow-up documents online; unclear whether data assessors could see group allocation
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Measures stated in the methods were reported in the results.

Scott 1990

Methods	Parallel RCT: brief intervention versus control. ITT: yes.
Participants	Setting: England, general practice. Participants: 17 to 69 years, consuming > 350 g alcohol for men and > 168 g for women in previous week; screened by Health Survey Questionnaire including quantity frequency measure of alcohol; excluded if consumption > 1050 g/week for men or 560 g/week for women, or previous advice to cut down during the previous year. Number randomised = 226; 68% male; mean age = 44.7 years. At baseline: mean alcohol consumption for previous week (from interview) = 526 g for men, 293 g for women; mean quantity frequency drinking for previous week (from HSQ) = 439 g for men, 247 g for women; binge drinkers = 43% for men (defined as consumption of 140 g on at least two occasions in previous three months), 18% for women (defined as consumption of > 14 units on at least two occasions in previous three months)
Interventions	Intervention group (N = 113) asked at the end of the assessment interview to make an appointment with their own general practitioner. Advice lasted ten minutes and consisted of feedback of the assessment interview and results of blood tests, information on the risks of excessive drinking, information on the benefits of drinking less, information on how the patient's weekly alcohol consumption compared with that of the general population using a histogram, and advice to reduce alcohol consumption to below four units per week. Advice was supplemented with a self-help booklet (the 'Cut Down on Drinking' booklet) designed for the study.

Scott 1990 (Continued)

	Control group (N = 113) received no advice from their general practitioner, except at their own request	
Outcomes	Change in weekly alcohol consumption. Assessed at one year.	
Funding source	The pilot study was supported by the Alcohol and Education Research Council and the main study by the Mental Health Foundation	
Declaration of interests	Not reported.	
Notes	'Cut Down on Drinking' trial. Loss to follow-up: Intervention group = 33/113 (29%). Control group = 43/113 (38%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random numbers table used to randomise patients (p. 891).
Allocation concealment (selection bias)	Low risk	Allocated at assessment interview before making an appointment with doctor (p. 891)
Blinding of treatment providers	Low risk	Researcher carried out baseline assessment and only intervention patients were asked to make an appointment with GP so low risk of contamination between groups
Blinding of participants	Low risk	No mention was made that the assessment interview was about alcohol consumption, and alcohol questions were embedded amongst other health-related questions
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up interview done by a researcher blinded to patient allocation
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Methods	Parallel RCT: motivational intervention and educational brochure versus educational brochure only ITT: no.	
Participants	Setting: Brazil, emergency room (ER). Participants: Aged 16 to 25 years, admitted to ER up to six hours after last alcohol use, permanent residents in Uberlandia, able to read and understand consent forms, screened with multiple choice questionnaire; excluded if no permanent address in the city, or unable to interview, or psychotic or mental disorder, or cognitive damage, or under arrest, or undergoing treatment at addiction care centres, or refusal to participate (due to physical pain, fatigue or lack of interest), or alcohol use > 6 hours prior to ER visit Number randomised = 175; 90% male; mean age = 21.8 years; 71% Caucasian; 1% illiterate, 49% junior high, 47% senior high, 3% university or college	
Interventions	MI group (N = 87) received one 45-minute motivational interview from a trained senior psychologist, followed by a three-page educational brochure on risks of alcohol consumption and possible ways or tips to consider reduction or to avoid problems related to alcohol abuse EB group (N = 88) received the same educational brochure plus maximum five minute discussion with trained junior psychologist researchers	
Outcomes	Mean number of days of alcohol use, mean number of days of light use (one to four units/day), both in previous three months Assessed at 90 days.	
Funding source	Not reported.	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: MI group: 13/87 (15%). EB group: 13/88 (15%).	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"A lottery system was employed" (p. 227).
Allocation concealment (selection bias)	Low risk	The lottery system was performed "by ER personnel not linked to the clinical trial in order to avoid selection bias" (p. 227)
Blinding of treatment providers	Low risk	One psychologist was responsible for administering intervention and different researchers were responsible for screening and "control" condition, therefore no

Segatto 2011 (Continued)

		chance of cross-contamination
Blinding of participants	Low risk	“Patients were blinded to the intervention applied” (p. 228)
Blinding of outcome assessment (detection bias) All outcomes	High risk	“The person who conducted the follow-up was not a baseline recruiter or interventionist, nor was intervention condition masked” (p. 228)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Senft 1997

Methods	Parallel RCT: brief intervention versus usual care. ITT: yes.
Participants	Setting: USA, primary care clinics. Participants: ≥ 21 years, total AUDIT score = 8 to 21, or sum of AUDIT frequency & quantity item scores ≥ 5 , or ≥ 6 drinks (defined as 10 oz beer or 4 oz wine or 1 oz liquor) per occasion at least weekly; screened by questionnaire including AUDIT; excluded if pregnant. Number randomised = 516; 71% male; mean age = 42.4 years; 82% white, 18% non-white; some college or more = 59.5%. At baseline: mean AUDIT score = 10.6; mean weekly drinking days = 3.4; mean drinks per drinking day = 4.9; binge drinkers (defined as ≥ 6 drinks per occasion at least weekly) = 28%
Interventions	Intervention group (N = 260) received a 30-second message from the primary care clinician and a 15-minute session with a health counsellor immediately following the medical visit. Clinicians included primary care physicians, nurse practitioners, and physicians' assistants. Screeners cued clinicians to deliver a brief message, based on the following script, to patients in their exam rooms: “Thank you for filling out the AUDIT questionnaire. I'm concerned that your drinking may pose a risk to your health. I recommend that you consider cutting down on your drinking and I'd like you to meet briefly with the health counsellor today after our visit”. Health counsellors were trained in principles of motivational interviewing. Components of the counselling session included: gathering additional information about the quantity and frequency of the subject's alcohol use and giving feedback by comparison to national norms; explaining the acute and chronic effects of alcohol use and teaching the patient ways to estimate blood alcohol level; recommending no more than three drinks daily for men, no more than two drinks daily for women, and no alcohol on at least two days per week. Abstinence was mentioned as the only sure method of eliminating health risks from alcohol and “zero” alcohol was recommended for driving; suggesting some options to help achieve lower-risk drinking.

Senft 1997 (Continued)

	If the patient was receptive, a low-risk drinking plan was created, building the patient's confidence that he or she could succeed. All intervention group patients, whether or not they agreed to the counselling session, were offered a packet of printed materials. Control group (N = 256) received usual care only.
Outcomes	Number of standard drinks or total standard ethanol content units in previous three months; drinking pattern (steady, episodic, or combination); peak blood alcohol concentration Assessed at 6 and 12 months.
Funding source	This research was supported by grant number R01 AA08976 from the National Institute on Alcohol Abuse and Alcoholism
Declaration of interests	Not reported.
Notes	Loss to follow-up: Intervention group: 64/260 (25%). Control group: 41/256 (16%). This study did not exclude participants told by their GP to avoid alcohol

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Method of allocation concealment not reported.
Blinding of treatment providers	High risk	GPs saw all participants and delivered a 30 second message giving advice to cut down to intervention group
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Follow-up telephone interviews done by interviewers who were unaware of subjects' study group status (p. 465)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but no reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Seppa 1992

Methods	Parallel RCT: brief intervention versus control. ITT: unclear.
Participants	Setting: Finland, general practice. Participants: adults with erythrocyte mean cell volume (MCV) ≥ 100 fl, ≥ 2 positive answers to Molmo Modified Michigan Alcoholism Screening Test and macrocytosis for which no other aetiology was found, medical history giving clues to alcohol abuse or an alcohol-induced increase in serum gamma glutamyltransferase (GGT) (≥ 50 U/L) were taken into account Number randomised = 178; 79% male; mean age = 53.2 years. At baseline: mean MCV = 101.6 fl; GGT = 145.4 U/L.
Interventions	Intervention group (N = 92) were asked to visit the doctor (GP) at intervals of three months for one year to monitor their MCV value and long-term alcohol consumption. The sessions were brief: the patients were asked about their alcohol consumption, were informed about the results of their blood test, and encouraged to minimise consumption. Control group (N = 86) received no intervention, but were invited by mail to contact their doctor one year later
Outcomes	Self-report of whether alcohol consumption was reduced; patient history for past year; MCV values Assessed at 12 months.
Funding source	Not reported.
Declaration of interests	Not reported.
Notes	Loss to follow-up: Intervention group = 51/92 (55%). Control group = 32/86 (37%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No information on how sequence was generated.
Allocation concealment (selection bias)	Unclear risk	No information on concealment of allocation.
Blinding of treatment providers	High risk	Doctors saw patients from both arms.
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	No information available on blinding.

Seppa 1992 (Continued)

Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Soderstrom 2007

Methods	Parallel RCT: brief personalised motivational intervention versus brief information and advice ITT: yes.
Participants	Setting: USA, Shock Trauma Centre. Participants: ≥ 18 years and English speaking, with intact cognition and hospital length of stay ≥ 12 hours, reporting (1) any positive response to an item of the CAGE questionnaire, or (2) drinking two or more times per week with total weekly drinking of eight or more drinks for women and 15 or more drinks for men, or (3) drinking two to four times a month or less with typical daily consumption of four or more drinks for women and five or more drinks for men, or (4) drinking six or more drinks on one occasion, weekly, daily or almost daily, screened with CAGE and AUDIT; excluded if unstable contact information, or legal issues (i.e. probation or parole), or a severe alcohol use problem considered inappropriate for a brief intervention, or a severe drug use problem Number randomised = 497; 85% male; mean age of total sample not reported; 63% White Baseline characteristics of total sample not reported.
Interventions	Personalised motivational intervention (PMI) group (N = 250) received (i) a 15 to 20 minute motivational interview from trained clinical psychology doctoral graduates who explored information obtained in the intake interview to help the subject evaluate his or her drinking patterns, assess motivations and barriers to change, explore motivation to modify drinking behaviours and support any initiative to reduce or eliminate excessive drinking; (ii) a personalised feedback letter that summarised participant intake data and provided feedback based on responses to the questionnaires regarding problem dimensions, as well as stage of change, decisional balance, self-efficacy, risk taking, stress and social support with some recommended actions, sent within one week of the subject's discharge; (iii) a phone call (duration not reported) from same interventionist after several days to reinforce the interview and discuss the letter, where attempts were made to elicit a specific change plan (according to the participant's stage of change) regarding drinking behavior; (iv) a further phone call (duration not reported) four weeks after trauma centre discharge, where the same interventionist checked the change plan, re-assessed motivation, discussed problem solving difficulties and barriers, supported self efficacy and reinforced the need and motivation to modify drinking behaviour Brief information and advice (BIA) group (N = 247) received (i) advice (duration not reported) from trained clinical psychology doctoral graduates to reduce their drinking to reduce future risk of injury; (ii) a two-page handout that described causes of injury, the role of alcohol and injury, definition of a drink, types of drinkers, alcohol consumption in the United States, alcohol-related problems, and tips on how to reduce drinking or receive help; (iii) a phone call (duration not reported) two weeks after discharge from

Soderstrom 2007 (Continued)

	the same interventionist asking how they were doing, if they had a chance to read the handout and if they had any questions
Outcomes	Mean number of drinks, mean number of binges, both within the last 90 days Assessed at 6 and 12 months.
Funding source	Supported by the National Institute on Alcoholic Abuse and Alcoholism (grant 2 RO1 AA09050-04A2)
Declaration of interests	Not reported.
Notes	Extended versus brief. Loss to follow-up at 12 months for outcome 1: 249 (50%) in total (unclear per arm) Loss to follow-up at 12 months for outcome 2: 248 (50%) in total We contacted the authors, who supplied missing data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Generation of randomisation sequence not reported.
Allocation concealment (selection bias)	Unclear risk	"After the intake was completed the intervention specialist opened a sealed envelope which indicated the subject's group assignment" (p. 1103)
Blinding of treatment providers	High risk	Not blinded and delivered both arms of trial therefore increased chance of contamination
Blinding of participants	High risk	Not blinded. Screening tools alcohol focused.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Unclear whether outcome assessor was blind to group allocation
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Unclear risk	Primary outcome not specified and not all measures listed in the methods are reported in the results

Methods	Parallel RCT: brief counselling versus simple advice versus usual care ITT: unclear.
Participants	Setting: USA, level 1 trauma centres. Participants: 18 to 45 years, English speaking with intact cognition, hospitalised within 24 hours of an injury after a motor vehicle crash, BAC \geq 10mg/dL, potential for discharge from hospital to home within four weeks after entry into the study; screened by daily review of the emergency department (ED) or trauma service admitting logs; excluded if they had attended an alcohol treatment program in the past year, or evidenced signs and symptoms of alcohol withdrawal, or had received advice from their health care provider in the past three months to reduce their alcohol use, or drank more than 150 g of alcohol (12 standard drinks) per day, or scored \geq 2 on any or all of the three alcohol-dependence items on AUDIT Number randomised = 187; 78% male; mean age = 29.03 years; 89% White At baseline: mean BAC = 165.18 mg/dL.
Interventions	BC group (N = 63) received (i) 20 minute brief counselling based on WHO model, TrEAT, FRAMES and reflective listening, using a printed self-help manual, from a nurse clinician hired for the study, on alcohol consumption and its consequences, recommended drinking limits, feedback about the participant's drinking behaviours to make the connection between drinking and the injury, pros and cons of changing drinking behaviours and how the participant's life might be different with a change in drinking, a realistic drinking agreement was negotiated based on the discussion; (ii) a 20 minute booster phone call by the same interventionist one month after discharge to review the agreement, explore risky situations, and devise coping mechanisms, drinking goals were re-negotiated SA group (N = 68) received (i) five minutes of feedback, using a printed self-help manual, from a nurse clinician hired for the study, that they were drinking in ways that put their health at risk and advice about current health behaviours and adverse effects of alcohol. The most important part of the simple advice intervention was to discuss "low risk" drinking and to review the menu of alternative choices: abstinence or cutting back to low risk drinking; (ii) a five minute booster phone call by the same interventionist one month after discharge, repeating the same information Control group (N= 56): no details reported.
Outcomes	Mean BAC, mean number of standard drinks in previous month, number of binges in previous month Assessed at 3, 6 and 12 months.
Funding source	Funded by the Centers for Disease Control and Prevention, National Center for Injury Control and Prevention, R49/CCR-510153
Declaration of interests	Not reported.
Notes	Loss to follow-up at 12 months: BC group: 26/63 (41%). SA group: 39/68 (57%). Control group: 22/56 (39%).

Sommers 2006 (Continued)

<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described.
Allocation concealment (selection bias)	Unclear risk	Not described.
Blinding of treatment providers	Unclear risk	Unclear if individual nurse clinicians delivered all arms.
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Interviewers were blinded to participant condition during follow-up interviews" (p. 525)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Sommers 2013

Methods	Parallel RCT: brief intervention group versus contact control group versus non-contact control group ITT: yes.
Participants	Setting: USA, emergency department. Participants: Aged 18 to 44 years admitted to emergency department between 8am and 2am on randomly selected days, drove a motor vehicle two days per week, screened positive on two risky driving and two hazardous drinking items in the three minute screen (at least two of the following in the past month: four standard alcoholic drinks on a typical drinking day, five drinks per occasion for women and six for men, 11 drinks per week for women and 14 for men); excluded if no identifiable residence or contact phone number, or were under arrest at the time of emergency department visit, or were unable to participate in an English language screen due to language or cognitive barriers, or were admitted to the hospital as an inpatient, or had a history of daily alcohol use suggesting alcohol dependence on AUDIT Number randomised = 476; 70% male; age range = 23 to 34.5 years; 62% non-White; 57% employed full-time, 17% employed part-time, 26% unemployed; 17% less than high school, 38% high school graduate, 45% more than high school Baseline data not reported for total sample.

Interventions	<p>BI group (N = 150) received (i) a 20 minute SBIRT intervention from a nurse clinician employed for the study, based on FRAMES and reflective listening, on both reducing risky driving practices and reducing alcohol (a drinking typology - dependent, heavy, and low-risk drinkers), the early consequences of heavy drinking, and tips to cut down on drinking. They received written feedback adapted from the TrEAT protocol on their risky driving and hazardous alcohol behaviours, and the interventionist discussed pros and cons of behaviours and negotiated a contract for change; (ii) a 20 minute booster telephone call 10 to 14 days after discharge the interventionist discussed and renegotiated the contract</p> <p>CC group (N = 162) received same 20 minute assessment as BIG group but no intervention</p> <p>NC group (N = 164) received no assessment or intervention.</p>	
Outcomes	<p>Number of participants drinking ≥ 5 drinks per drinking day in previous three months, median maximum number of drinks within six hour period, median number of drinks in a typical week, median number of drinks in a typical day, number of participants having ≥ 1 binge per month, number of participants with AUDIT > 10; also driving-related outcomes</p> <p>Assessed at 3, 6, 9 and 12 months.</p>	
Funding source	<p>This work was funded by a grant (R49/CCR-523225) from the National Center for Injury Prevention and Control, Centers for Disease Control and Prevention (PI:MSS)</p>	
Declaration of interests	<p>Not reported.</p>	
Notes	<p>An SBIRT study.</p> <p>USD 20 at enrolment; USD 20 for baseline, three, six, nine month interviews, USD 40 for 12 month interview</p> <p>Loss to follow-up:</p> <p>BIG group: 53/150 (35%).</p> <p>CCG group: 53/162 (33%).</p> <p>NCG group: 43/164 (26%).</p> <p>We contacted the authors, who supplied missing data.</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation by computer software (p. 1754).
Allocation concealment (selection bias)	Low risk	"Enrolment packets of the same size and thickness were placed in opaque envelopes, sealed and locked in a drawer in consecutive order by participant number" (p. 1754)

Sommers 2013 (Continued)

Blinding of treatment providers	High risk	The assessment, brief intervention, and booster intervention were delivered by 4 nurse clinicians employed by the study team; i.e. treatment providers had contact with all participants
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	“Interviewers were blinded to condition during follow-up interviews” (p. 1755)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30% overall.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods were reported.

Spirito 2004

Methods	Parallel RCT: motivational interview and handout versus brief advice and handout (“standard care”) ITT: no.
Participants	Setting: USA, emergency department. Participants: 13 to 17 years and English speaking, presenting to emergency department with evidence of alcohol in blood, breath, or saliva, or reported drinking alcohol in the six hours before the injury that required treatment in the emergency department, assessment 45 minutes, received intervention once BAC < 0.1 and/or they could pass mental state examination; excluded if suicidal, or in police custody, or had suffered serious traumatic injury requiring admission Number randomised = 152; 64% male; mean age = 15.6 years; 72% White, 7% African American, 3% Asian/East Indian, 17% Hispanic, 1% Native American; 28% middle school, 57% high school, 3% high school graduate or equivalent, 12% dropped out or expelled Baseline data not reported for total sample.
Interventions	MI group (N = 78) received (i) handout on avoiding drinking and driving; (ii) list of substance abuse treatment referral services; (iii) personalised feedback sheet; (iv) information sheets on negative effects of alcohol; (v) 35 to 45 minute motivational interview with study researchers trained in MI. MI protocol included the following six components: (1) emphasis on personal responsibility for change and that any decisions made about changing their drinking would be up to them; (2) exploration of motivation for drinking and review of potential negative consequences; (3) personalised normative assessment feedback about their pattern of alcohol use and risks; (4) imagining their future if they continued to drink the same way versus if they decided to make a change in their drinking behaviour; (5) establishing goals with regard to drinking; and (6) anticipating barriers to accomplishing the goals. In addition to being empathic and avoiding argumentation,

	<p>the interventionists made an effort to develop a discrepancy between the teen's current drinking behavior and current and longer-term goals, and to support the teen's sense of self-efficacy about making changes. They recommended follow-up with primary care provider</p> <p>SC group (N = 74) received (i) the same handout on avoiding drinking and driving; (ii) list of substance abuse treatment referral services (iii) five minute brief advice session from emergency department physicians advising participants to stop drinking. They recommended follow-up with primary care provider</p>
Outcomes	<p>Mean days per month drinking, mean drinks per occasion, mean days per month bingeing (≥ 5 drinks per occasion), mean number of alcohol-related injuries, percentage reporting alcohol-related problems</p> <p>Assessed at 3, 6 and 12 months.</p>
Funding source	Supported by grant number AA09892 from the National Institute on Alcohol Abuse and Alcoholism
Declaration of interests	Not reported.
Notes	<p>USD 20 gift certificate on completion of intervention, USD 10 at three months, USD 15 at six months, USD 25 at 12 months</p> <p>Only received intervention if well enough to be discharged.</p> <p>Loss to follow-up: MI group: 14/78 (18%). SC group: 14/74 (19%).</p>

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Used a random numbers table but randomisation took place after baseline assessment (p. 397)
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of treatment providers	High risk	Research interventionist delivers both arms.
Blinding of participants	High risk	No reported attempt to blind participants.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Research assistants... were blind to treatment group assignment" (p. 398)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but not reasons.

Spirito 2004 (Continued)

Selective reporting (reporting bias)	Unclear risk	Two items from the ADQ were very highly correlated so they only reported one of them
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Spirito 2011

Methods	Parallel RCT: individual motivational interview versus individual motivational interview and family check up ITT: not reported.
Participants	Setting: USA, emergency department. Participants: Aged 13 to 17 years, English or Spanish speaking, presenting to urban level 1 trauma centre with positive BAC using blood, breath or saliva, or self-reported drinking in six hours previous to emergency department visit, approached only when BAC < 0.1% or they passed a mental status examination; screened by BAC or self-reported drinking within six hours of visit (Adolescent Drinking Questionnaire); excluded if suicidal, or had experienced serious traumatic injury Number randomised = 125; 46% male; mean age of total sample not reported; 66% White, 2% Black, 2% Asian American or East Indian, 28% Hispanic, 2% mixed race At baseline: mean days per month drinking = 3.4, mean drinks per occasion = 5.3, mean days per month drinking high volume (≥ 5 drinks per occasion) = 2.6
Interventions	IMI group (N = 63) received (i) 45 to 60 minute interview from interventionists with masters degrees in counselling and psychology that included an emphasis on personal responsibility, exploration of motivation for drinking and review of potential negative consequences, personalised normative assessment feedback, establishing goals regarding drinking, and anticipating barriers to accomplishing goals, such as peer pressure to drink. In addition to being empathic and avoiding argumentation, the interventionists made an effort to develop a discrepancy between the adolescent's current drinking and longer-term goals and to support the adolescent's sense of self-efficacy. (ii) Participants' parents were given follow-up information regarding substance use treatment services and received monthly booster brochures on parenting IMI + FCU group (N = 62 randomised, 50 received intervention) received their intervention from interventionists with masters degrees in counselling and psychology: (i) the same 45 to 60 minute interview (participant only). (ii) Additionally, the family received a one hour videotaped family assessment task (called "FAsTask6"), which consists of the parent(s) and the adolescent discussing family beliefs regarding alcohol, marijuana, cigarette, other drug use and other topics, such as curfew. All FAsTask videos were coded separately by the treatment provider and a second rater to provide information to be used in the feedback session. (iii) A week after the family completed the FAsTask, parents received one hour feedback (self-assessment, support and clarification, feedback, and development of a plan for change) derived from the FAsTask and from self-report measures that assessed family stress, parent substance use, and beliefs regarding substance use. Parent motivation for change, change options, and specific steps for making positive changes in family relationships and parenting were discussed, including potential barriers to change. (iv) Participants' parents were given the same follow-up information regarding substance use treatment services and received monthly booster brochures on parenting

Outcomes	Mean days per month drinking, mean drinks per occasion, mean days per month with high volume drinking (≥ 5 drinks per occasion), number of participants reporting high volume drinking (≥ 5 drinks per occasion); all for previous three months Assessed at 3, 6 and 12 months.
Funding source	This study was supported by grant AA013385 from the National Institute on Alcohol Abuse and Alcoholism. The study sponsor (the National Institutes of Health) had no role in (1) the study design; (2) the collection, analysis, and interpretation of data; (3) the writing of the report; and (4) the decision to submit the paper for publication
Declaration of interests	Financial Disclosure: none reported.
Notes	Brief intervention vs extended intervention USD 20 gift certificate for adolescent for IMI completion; USD 50 for parents for FCU completion Loss to follow-up at 12 months: IMI group: 16/63 (25%). IMI + FCU group: 26/62 (42%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Patients were randomly assigned but method of sequence generation not reported
Allocation concealment (selection bias)	Unclear risk	"Patients were then randomly assigned ... based on condition assignments contained in sealed envelopes" (p. 270)
Blinding of treatment providers	Low risk	Both arms receive initial intervention (so no contamination) and then booster is a separate visit which comprises a discussion between the family
Blinding of participants	Low risk	Both arms receive initial intervention so no differential knowledge of intervention between the groups
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Research assistants... were masked to treatment group assignment" (p. 270)
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30% overall.

Spirito 2011 (Continued)

Selective reporting (reporting bias)	Unclear risk	Prespecified “feeling drunk” outcome was highly correlated with “high volume drinking days” so former was omitted from results
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Tait 2004

Methods	Parallel RCT: brief intervention versus usual care. ITT: unclear.	
Participants	Setting: Australia, emergency department. Participants: 12 to 19 years, presenting to emergency department with involvement of alcohol or other drug use, screening “via the hospital ED information system and by liaison with medical staff” (no more details provided); no exclusion criteria specified Number randomised = 127; 55% male; mean age = 16.7 years; 83% born in Australia; 17% in full time work Baseline characteristics not reported for total sample.	
Interventions	BI group (N = 60) received (i) an interview (duration not reported) with a named counsellor (researcher, social worker or health worker), aimed at facilitating attendance at a community AOD treatment agency. They identified and discussed negative consequences of alcohol/drug use, impediments to reduce negative consequences and high risk (trigger) situations; offered advice about appropriate services that were available in his/her local area and the type of intervention that would be offered by the external agency. Possible impediments to treatment were discussed with the adolescent (e.g. need for a counsellor of the same gender). (ii) Participants received a reminder telephone call prior to attending their appointment UC group (N = 67) received “usual hospital care” (details not specified)	
Outcomes	Attendance at community treatment agency by four months (reported at 4 and 12 months), hazardous alcohol use (AUDIT-3 ≥ 5 for men or ≥ 4 for women) at 12 months only, GHQ-12, FAD (family assessment device) scores Assessed at 4 and 12 months.	
Funding source	This study was funded by Healthway, the West Australian Health Promotion Foundation. The Alcohol Education and Rehabilitation Foundation funded the collection of ED data and subsequent analyses	
Declaration of interests	Not reported.	
Notes	Loss to follow-up at 12 months: BI group: 22/60 (37%). UC group: 18/67 (27%). We contacted the authors, who supplied missing data.	
<i>Risk of bias</i>		
Bias	Authors’ judgement	Support for judgement

Tait 2004 (Continued)

Random sequence generation (selection bias)	Low risk	Computer generated randomisation codes (p. 366).
Allocation concealment (selection bias)	Unclear risk	“Codes were held in numbered, sealed envelopes by the interviewer” (p. 366)
Blinding of treatment providers	Unclear risk	Unclear whether interventionist researcher had contact with both groups
Blinding of participants	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not reported.
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30% overall.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Tomson 1998

Methods	RCT: brief intervention versus usual care. ITT: unclear.
Participants	Setting: Sweden, general practice health centre. Participants: 25 to 54 years with GGT > 0.89 microkatal/L; screened by health check including GGT; excluded if chronic alcoholic. Number randomised = 222. The following data is for N = 75, who were not excluded and then were followed up: 81% male; mean age = 45.2 years; 73% blue collar, 27% white collar. At baseline: mean weekly consumption given at baseline only for intervention group = 337 g; mean S-GGT = 1.7 microkatal/L
Interventions	Intervention group (N = 100) had an appointment with a nurse to establish whether the laboratory finding could be explained by excess alcohol intake. The nurse had experience from outpatient care of alcoholics and made a ‘general lifestyle and health survey interview’. The assessment also included self-reported alcohol consumption and the CAGE questionnaire. Patients responding “yes” to two or more of the four questions were classified as positive. The nurse discussed those with an unclear elevation of GGT with the GP, who sometimes recommended further examination and/or laboratory investigations. Participants classified as high consumers of alcohol (280 and 210 g 100% alcohol per week for men and women, respectively) were offered two more consultations with the nurse during the intervention year. The consultations focused on lifestyle in general and alcohol consumption in particular. Factors that facilitated or made controlled drinking more difficult were discussed in an empathic way. GGT was used as a biochemical feedback at follow-up.

	Control group (N = 122) recommended an appointment with the GP. The participants were asked about diseases and ongoing medication. They were told that the most common reasons for an elevated GGT had to do with eating and drinking habits and drug use. They were asked about their own thoughts concerning the reason for their elevated GGT. When the spontaneous response was that their alcohol consumption was a plausible explanation, the GP accepted this without comment. In the other cases the GP also made a clinical and an extended laboratory investigation. Those classified as high consumers were not given any advice about how to reduce their drinking, but were told that a follow-up was planned in one to two years	
Outcomes	GGT measured at one and two years. Data about sickness allowance days one year before, during, and one year after the intervention, and notations in the social services records were collected. The number of visits to the health centre for reasons other than alcohol intervention (intervention group), days of hospital care, and deaths up to two years after intervention were followed Assessed at one and two years.	
Funding source	Not reported.	
Declaration of interests	Not reported.	
Notes	Loss to follow-up: Intervention group: 70/100 (70%). Control group: 77/122 (63%). Randomisation (n = 222) then assessment & exclusion (leaving n = 75 who actually had the intervention or control sessions)	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Date of birth used for randomisation (as reported in Kristensen 1983)
Allocation concealment (selection bias)	Unclear risk	No information on how allocation was concealed.
Blinding of treatment providers	Low risk	Different treatment providers interacted with the different arms (nurse for intervention group, doctor for control group: "the GP was anxious not to use the consultation with the control group as an intervention but to try to rule out an elevated GGT due to a serious disease" - p. 189)
Blinding of participants	Low risk	Recruitment letter to participants does not discuss alcohol as a reason for attendance but just that elevated blood test needed a follow-up

Tomson 1998 (Continued)

Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Not clear whether outcome assessors were blinded.
Incomplete outcome data (attrition bias) All outcomes	High risk	Loss to follow-up > 30%.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Wallace 1988

Methods	RCT: brief intervention versus control. ITT: unclear.
Participants	Setting: UK, general practice. Participants: 17 to 69 years, had consumed ≥ 35 units of alcohol in previous week for men and 21 units for women, or had ≥ 2 positive CAGE responses; screened by self-administered health survey questionnaire including quantity/frequency scale and modified CAGE; excluded if serious illness, or received medical advice about drinking in the previous year, or requested help from nurse with drinking, or GGT > 150 IU/L. Number randomised = 909; 71% male; mean age = 42.4 years. At baseline: alcohol consumption for previous week (from interview) = 55.0 units; alcohol consumption according to questionnaire (quantity/frequency) = 44.1 units; GGT = 23.0 IU/L
Interventions	Intervention group (N = 450) contacted by their general practitioner and asked to attend for a brief interview. After an assessment interview about the pattern and amount of alcohol consumption and evidence of alcohol related problems and dependence (obtained by using the brief Edinburgh alcohol dependence scale), patients were shown a histogram based on figures from a national survey of drinking habits to illustrate how their weekly consumption compared with that of the general population. Advice was given about the potential harmful effects of their current level of alcohol consumption, backed with the information booklet 'That's the Limit'. Men were advised to drink not more than 18 U/week and women not more than 9 U/week. Where there was evidence of dependence on alcohol general practitioners were encouraged to advise abstinence. Patients were given a drinking diary, the front cover of which was a facsimile of an ECIO prescription with the words "Cut Down on your Drinking!" The last page contained a guide to the alcohol content (in U) of a range of drinks. An initial follow up appointment one month later was offered to all patients; subsequent appointments at 4, 7 and 10 months were at the discretion of the general practitioner. During these sessions, the patient's drinking diary was reviewed and feedback given on the results of blood tests indicating evidence of damage due to alcohol Control group (N = 459) received no advice from their general practitioner about drinking except at their own request or if there was evidence that their alcohol consumption had already resulted in substantially impaired liver function (γ -glutamyltransferase activity exceeding 150 IU/L)

Outcomes	Stated alcohol consumption in the seven days before interview; consumption as assessed by the health survey questionnaire; changes in the values of laboratory indicators of alcohol consumption Assessed at 6 and 12 months.
Funding source	The study was generously supported by grants from the Alcohol and Education Research Council, the Brewers' Society, the Medical Council on Alcoholism, the Medical Research Council, and Merck Sharp and Dohme Ltd
Declaration of interests	Not reported.
Notes	Loss to follow-up: Intervention group: 87/450 (19%). Control group: 74/459 (16%).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not reported.
Allocation concealment (selection bias)	Unclear risk	Not reported.
Blinding of treatment providers	Low risk	GPs only interviewed intervention participants and were not involved in recruitment or baseline assessment
Blinding of participants	Low risk	Recruitment and screening described as "lifestyle and health survey interview" (p. 664)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Nurse conducting interviews were unaware of patient allocation (p. 664)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Methods	Parallel RCT: computer-administered brief intervention versus therapist-administered brief intervention versus brochure on alcohol and violence ITT: unclear.
Participants	Setting: USA, emergency department. Participants: 14 to 18 years, presenting to emergency department for medical illness or injury with normal vital signs (including trauma patients after initial stabilisation), reporting past-year alcohol consumption and aggressive behaviour; screened with computerised self-complete survey plus AUDIT-C, CRAFFT and Problem Oriented Screening Instrument for Teenagers (POSIT); excluded if victim of acute sexual assault, or displaying suicidal ideation, or altered mental status precluded consent, or medical instability (i.e. abnormal vital signs). Number randomised = 726 ;44% male; mean age 16.8 years; 56% African American, 39% White, 7% Hispanic, 5% Other ethnicity; 10% dropped out of school. At baseline: 52% past year binge drinking (≥ 5 drinks), 48% AUDIT-C \Rightarrow 3, 45% POSIT alcohol score ≥ 2
Interventions	Computer-administered brief intervention (CBI) group (N = 237) had one session (median 29 minutes) on an interactive multimedia computer program developed for the study and viewed on tablet laptops with touch screens and audio delivered through headphones, to ensure participant privacy. Therapist-administered brief intervention (TBI) group (N = 254) received one session (median 37 minutes) from a research therapist who utilised a tablet laptop to provide personalised feedback from the screening and baseline surveys as well as age- and sex-specific normative information. Adolescents completed computerised checklists identifying reasons to stay away from drinking and fighting. Using a preprogrammed algorithm, the computer selected a set of role-play scenarios based on the participant's risk behaviours, and the therapist guided the participant Control group (N = 235) received a brochure containing information on alcohol and violence with community resources including phone numbers
Outcomes	Per cent with AUDIT-C ≥ 3 , per cent binge drinking (≥ 5 drinks per occasion), per cent with POSIT (Problem Oriented Screening Instrument for Teenagers) ≥ 2 Assessed at 3, 6 and 12 months.
Funding source	This project was supported by National Institute on Alcohol Abuse and Alcoholism (NIAAA) grant 014889
Declaration of interests	Not reported.
Notes	SafERteens study; teenagers Payment: USD 1 for screening; USD 20 for baseline assessment; USD 25 for three month assessment; USD 30 for six month assessment; USD 35 for 12 month assessment; USD 111 in total. Loss to follow-up at three months: CBI group: 32/237 (14%). TBI group: 39/254 (15%). Control group: 29/235 (12%). Loss to follow-up at six months: CBI group: 28/237 (12%).

Walton 2010 (Continued)

	<p>TBI group: 45/254 (18%). Control group: 27/235 (11%). Loss to follow-up at 12 months: CBI group: 36/237 (15%). TBI group: 50/254 (20%). Control group: 33/235 (14%). We contacted the authors, who supplied missing data.</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was based on computer-generated algorithm (p. 528)
Allocation concealment (selection bias)	Unclear risk	Used "numbered sealed envelopes" (p. 528)
Blinding of treatment providers	High risk	Not blinded.
Blinding of participants	Low risk	"Participants were blinded to condition assignment until after the baseline assessment" (p. 528)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Follow-up staff were blinded to baseline condition assignment" (p. 528)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Loss to follow-up reported with reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Watson 2013

Methods	RCT: stepped care versus minimal intervention. ITT: yes.
Participants	<p>Setting: England and Scotland, general practices.</p> <p>Participants: Aged ≥ 55 years, scored ≥ 8 on AUDIT, provided contact details on the screening form, were residing in a stable place of residence, lived within commutable distance of the primary care centre; excluded if they had received treatment for substance use, excluding nicotine, in the previous 90 days, were already seeking help for alcohol use, had any outstanding legal issues likely to lead to imprisonment, suffered from severe mental or physical illness likely to preclude active participation in treatment or follow-up</p> <p>Number randomised = 529; 80.3% male; mean age = 62.83 years (SD = 5.83); 35.3%</p>

	<p>employed, 52.3% retired, 1.4% home worker, 0.2% student, 2.7% seeking work; 63.3% continued education after school, 41.8% degree or equivalent qualification At baseline: mean drinks per day = 3.39 (SD = 2.21), mean AUDIT-C score = 8.26 (SD = 2.22), mean drinking problems index score = 2.86 (SD = 3.12)</p>	
Interventions	<p>Stepped care (N = 266) consisted of three consecutive steps, in which progression between steps was dependent upon the outcome of each previous step Step 1: one 20 minute session of behavioural change counselling (based on motivational interviewing) was delivered by the practice/research nurse. The counselling was protocol guided. Four weeks after randomisation, the participant was contacted by the nurse and a short telephone assessment was made regarding the participant's alcohol consumption in the previous four weeks using the extended AUDIT-Consumption (three item) (AUDIT-C). If the participant was still consuming alcohol at hazardous levels, a referral was made to step 2 of the intervention Step 2: MET intervention by a trained therapist in the primary care environment. Three weekly 40 minute sessions; protocol guided; addressed six basic principles of increasing motivation for change. Four weeks after the final MET session, the nurse contacted the participant and a short telephone assessment was made regarding the participant's alcohol consumption in the previous four weeks using the extended AUDIT-C. If the participant was still consuming alcohol at hazardous levels, a referral was to be made to step 3 of the intervention Step 3: a referral to the local specialist alcohol services to receive specialist intervention including, as necessary, detoxification, inpatient care, outpatient counselling, group therapy, relapse prevention treatment or medication Minimal intervention group (N = 263) received a five minute brief advice intervention with the practice nurse or research nurse involving feedback of the results of the screening and discussion regarding the health consequences of continued hazardous alcohol consumption. The participant also received a brief self-help booklet, 'Safer drinking - a self help guide', outlining the consequences of excessive alcohol consumption and providing information on sources of help for drinking problems locally and nationally</p>	
Outcomes	<p>Mean drinks per day, AUDIT-C score, drinking problems index. Assessed at 6 and 12 months.</p>	
Funding source	<p>The research reported [here] was funded by the HTA programme as project number 06/304/142</p>	
Declaration of interests	<p>Declared competing interests of authors: none.</p>	
Notes	<p>Loss to follow-up: Stepped Care group: 31/266 (12%). Minimal intervention: 35/263 (13%).</p>	
<i>Risk of bias</i>		
Bias	Authors' judgement	Support for judgement

Watson 2013 (Continued)

Random sequence generation (selection bias)	Low risk	"The generation of the randomisation sequence and subsequent treatment allocation were performed by an independent, secure, remote, telephone randomisation service based at the University of York" (p. 7)
Allocation concealment (selection bias)	Low risk	"The generation of the randomisation sequence and subsequent treatment allocation were performed by an independent, secure, remote, telephone randomisation service based at the University of York" (p. 7)
Blinding of treatment providers	High risk	"Owing to the nature of the intervention and the pragmatic aim of the evaluation, treatment allocation, once determined, was not concealed from the participant or the professional delivering the intervention (p. 7)
Blinding of participants	High risk	"Owing to the nature of the intervention and the pragmatic aim of the evaluation, treatment allocation, once determined, was not concealed from the participant or the professional delivering the intervention" (p. 7)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Participants self-completed postal questionnaires.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up is reported but not reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

Woolard 2013

Methods	Parallel RCT: brief intervention versus standard care. ITT: yes.
Participants	Setting: USA, emergency department. Participants: ≥ 18 years, reported alcohol use in the prior month and marijuana use in the prior year; screened by the Wellness questionnaire; excluded if they were admitted to the hospital, or in treatment for substance use or psychiatric disorders, or were not English speaking, or were in police custody or were unable to consent or too medically unstable to participate in the research Number randomised = 515; 83.4% male; 68% White, 17% Hispanic

Interventions	<p>BI group (N = 249) received two counselling sessions based on the principles of MI as described by Miller and Rollnick (2002) and the MATCH MET treatment manual. The first BI session lasted between 20 and 60 minutes, and the second 15 to 40 minutes. The goal of the first brief intervention session was to engage the participant in reflection upon the pros and cons of alcohol and marijuana use. The intervention included: feedback; emphasis on personal responsibility for change; advice with permission; a menu of alternative change options; an empathic interventionist style; and fostering patient self-efficacy. The interventionist provided direct feedback concerning the participant's alcohol and marijuana use compared to community norms. The participant and interventionist developed a change plan that addressed changes the participant wanted to make, usually focused on reduction of substance use. If the participant was not ready to change substance use, the interventionist worked with the participant to increase motivation for change. The interventionists discussed with participants the pros and cons of alcohol and marijuana use and also explored their conjoint use, and the effect that conjoint use had on the reported pros and cons. The focus of the second BI session was to review and reinforce the change plan for those who had one, or to increase motivation for change and create a change plan with those who had not made a change plan in the first session</p> <p>Standard care group (N = 266) received routine emergency care for their presenting medical complaint during and after completing the baseline assessment. All study participants were offered information on local treatment resources for substance misuse either following the BI or after the baseline assessment for the SC group</p>	
Outcomes	<p>Days alcohol use in past 30 days, days alcohol binge use in past 30 days, alcohol negative consequences</p> <p>Assessed at 3 and 12 months.</p>	
Funding source	<p>This study was supported by grant funding from NIAAA (1R01AA13709-01A1)</p>	
Declaration of interests	<p>There are no conflicts of interest by any author.</p>	
Notes	<p>Payment: USD 25 for baseline and three month assessments and USD 50 for 12 month assessments</p> <p>Loss to follow up:</p> <p>Brief intervention group: 43/249 (17%).</p> <p>SC group: 46/266 (17%).</p>	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method of sequence generation not described.
Allocation concealment (selection bias)	Unclear risk	Closed envelopes but does not specify numbered or opaque (p. 1734)
Blinding of treatment providers	Low risk	Interventionist only delivered one style of intervention and did not come into contact

		with control participants
Blinding of participants	Low risk	“Participants were not informed of the hypotheses being tested” (p. 1734)
Blinding of outcome assessment (detection bias) All outcomes	Low risk	RAs “blinded to the participant’s original treatment condition” (p. 1734)
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Loss to follow-up reported but not reasons.
Selective reporting (reporting bias)	Low risk	Outcomes specified in methods are reported.

AA: Alcoholics Anonymous

ADM-related days: services for conditions related to alcohol, drugs or mental health

ALT: alanine aminotransferase

AST: aspartate aminotransferase

AUDIT: Alcohol Use Disorders Identification Test

AUDIT-C: Alcohol Use Disorders Identification Test Consumption, a brief version of AUDIT

BAC: blood alcohol content CAGE: Cut down, Annoyed, Guilty, Eye-opener, an alcohol screening tool

CDT: Carbohydrate-deficient transferrin, a blood test used to detect heavy alcohol use

CRAFFT: Car, Relax, Alone, Forget, Friends, Trouble, an alcohol screening tool

DRInC: drinker inventory of consequences

FRAMES: Feedback, Responsibility (of the drinker for change), Advice, Menu of options, Empathy (from interventionist), Self efficacy (from the drinker)

GGT: gamma-glutamyltransferase

GP: general practitioner

ITT: intention-to-treat

IMI: individual motivational interview

MALT: Munich alcoholism test

MCV: mean corpuscular volume, part of a blood count

MI: motivational interview

NIAAA: National Institute on Alcohol Abuse and Alcoholism (USA)

PAT: Paddington Alcohol Test, an alcohol screening tool

RA: research assistant

RAPS: Remorse, Amnesia, Perform, Starter, an alcohol screening tool

RCT: randomised controlled trial

SBIRT: screening, brief intervention and referral to treatment

SC: stepped care

TLFB: Timeline Followback, a method of self-assessing alcohol consumption

WHO: World health Organization

Characteristics of excluded studies *[ordered by study ID]*

Study	Reason for exclusion
Academic ED SBIRT 2007	Participants in both arms received the interventions sequentially rather than in parallel
Ameratunga 2011	Not a conversation-based intervention.
Apodaca 2007	Not a brief intervention; the face-to-face element was an explanation of the written materials
Areesantichai 2010	Participants were not recruited from primary care.
Areesantichai 2013	Participants were not recruited from primary care.
Babor 1996	Some participants were not recruited from primary care: unable to separate outcome data
Baker 2014	Participants were not recruited from primary care.
Ball 2007	Participants were not recruited from primary care.
Boekeeloo 2004	Not a conversation-based intervention.
Ceperich 2011	Participants were not recruited from primary care.
Chander 2015	Participants were not recruited from primary care.
Chang 1997	Comparator was alcohol treatment referral (previously included study, Kaner 2007).
Field 2014	A large proportion of participants were dependent drinkers and had been admitted to the trauma centre
Gentilello 1999	Participants were inpatients; excluded if discharged within 24 hours (previously included study, Kaner 2007).
Gwadz 2008	Participants were not recruited from primary care.
Haller 2014	Intervention was practitioner training.
Hansen 2012	Not a conversation-based intervention.
Hansen 2012a	Participants were not recruited from primary care.
Hansson 2007	Participants were not recruited from primary care.
Hilbink 2012	Not a brief intervention.
Houry 2011	Not a conversation-based intervention.
Hyman 2007	Implementation trial.

(Continued)

Ingersoll 2013	Participants were not recruited from primary care.
Ivanets 1991	Some participants were not recruited from primary care: unable to separate outcome data
Jessop 2008	Not a conversation-based intervention.
Kristenson 1983	Participants were not recruited from primary care.
Kulesza 2013	Participants were not recruited from primary care.
Lang 2007	Not an RCT.
Maheswaran 1992	Participants were not recruited from primary care.
Massey 2015	Neither arm was conversation only.
McDevitt-Murphy 2015	Comparator arm was not eligible - active intervention.
Mezzani 2007	Implementation trial.
Milner 2010	Not an RCT.
Monti 1999	A screening tool was not used to recruit participants.
Nayak 2015	Outcomes are presented only by readiness to change scores.
Nilssen 1991	Participants were not recruited from primary care.
Pal 2007	Participants were not recruited from primary care.
Palm 2016	Participants were not screened before randomisation; trial contained non-risky drinkers
Rhodes 2015	Not all participants were risky drinkers.
Shetty 2011	Participants were not recruited from primary care - they attended a clinic for follow-up care at a level 1 trauma centre
Sterling 2017	Not all participants were risky drinkers.
Toft 2009	Participants were not recruited from primary care.
UKATT 2005	Participants were not recruited from primary care.
Walton 2015a	Only admitted inpatients were screened and recruited.
Worden 2011	Participants were not recruited from primary care.

(Continued)

Wutzke 2002	Some participants were not recruited from primary care: unable to separate outcome data
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Characteristics of studies awaiting assessment [ordered by study ID]

Arnaud 2017

Methods	Cluster RCT
Participants	Setting: Germany; children under 18 years who presented to emergency departments of paediatric hospitals with alcohol intoxication
Interventions	Single session brief motivational intervention (manual-based, delivered by staff external to hospital) with telephone booster after six weeks, plus carer consultation
Outcomes	Frequency of binge drinking, number of alcoholic drinks on a typical occasion, and alcohol-related problems using the brief Rutgers Alcohol Problem Index; measured at three and six months
Notes	

Cherpitel 2016

Methods	Parallel design RCT
Participants	Setting: USA, participants were those aged 18 to 30 years who presented to the emergency department, self-identified as of Mexican origin, reported consuming 15 (men) or 8 (women) drinks/week, or 5 (men) or 4 (women) drinks/occasion in previous 28 days
Interventions	Promotores (peer health promotion advocates) delivered brief negotiation interview (20 minutes)
Outcomes	Mean days binge drinking, mean drinking days/week, mean drinks/drinking day; measured at 3 and 12 months
Notes	

Crawford 2015

Methods	Parallel design RCT
Participants	Setting: UK, adults aged 19 years or over presenting to (primary care) sexual health clinic and reporting excessive consumption - 8 (men) or 6 (women) drinks on a single occasion more than monthly
Interventions	Brief advice (2 to 3 minutes) from the treating clinician plus offer of an appointment with an alcohol health worker, who delivered a FRAMES brief intervention for up to 30 minutes
Outcomes	Mean weekly alcohol consumption measured at six months

Crawford 2015 (Continued)

Notes	Project SHEAR (Sexual Health and Excessive Alcohol: Randomised trial)
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Diaz Gomez 2017

Methods	Parallel design RCT
Participants	Setting: France, participants aged 16 to 24 years presenting to emergency department
Interventions	Brief motivational intervention (duration uncertain)
Outcomes	Mean number of drinks, measured at 3 months
Notes	

Mello 2016

Methods	Parallel design RCT
Participants	Setting: USA, adults aged 18 years or over presenting to emergency department and reporting moderate or high-risk alcohol consumption (ASSIST score 11 or greater)
Interventions	Telephone-based brief motivational interviewing, median 40 minutes, maximum of 3 calls
Outcomes	Frequency of binge drinking, mean number of drinks/week, maximum number of drinks/occasion; all in previous 30 days; measured at 12 months
Notes	Project ReDIAL

Monti 2015

Methods	Parallel design RCT
Participants	Setting: USA, participants aged 18 to 65 years who received treatment in emergency department and screened for harmful alcohol use and risky sexual behaviour
Interventions	Brief motivational intervention
Outcomes	Mean number of heavy drinking days, mean drinks/week; measured at 3, 6 and 9 months
Notes	

Nadkarni 2017

Methods	Parallel design RCT
Participants	Setting: India, men aged 18 to 65 years presenting to primary health care clinic who scored 12 to 19 on AUDIT
Interventions	Counselling for Alcohol Problems (CAP); a manual intervention delivered over a maximum of four sessions of 30 to 45 minutes duration (i.e. an extended intervention by the definition of this review)
Outcomes	Remission (AUDIT score > 8), mean drinks/day, per cent days abstinent, per cent days heavy drinking; measured at three months
Notes	

Walton 2015

Methods	Parallel design RCT
Participants	Setting: USA, participants aged 14 to 20 years presenting to emergency department who screened positive for risky drinking on AUDIT
Interventions	Motivational interviewing based brief intervention delivered by therapist (there is also a computer-delivered and a control arm)
Outcomes	Alcohol consumption (conference abstract gives no more detail of type of consumption), measured at three months
Notes	Project U Connect

ASSIST: Alcohol, Smoking and Substance Involvement Screening Test

AUDIT: Alcohol Use Disorders Identification Test

FRAMES: Feedback, Responsibility (of the drinker for change), Advice, Menu of options, Empathy (from interventionist), Self efficacy (from the drinker)

Characteristics of ongoing studies *[ordered by study ID]*

ISRCTN13832949

Trial name or title	Development and evaluation of the efficacy of a brief motivational intervention among young adults admitted in the emergency room while being alcohol intoxicated
Methods	Parallel design RCT
Participants	Setting: Switzerland, adults aged 18 to 35 years attending the emergency room with alcohol intoxication
Interventions	Brief motivational intervention
Outcomes	Heavy drinking days in the last month

ISRCTN13832949 (Continued)

Starting date	1 December 2016
Contact information	Dr Jacques Gaume Lausanne University Hospital Avenue de Beaumont 21 bis - P2, Lausanne 1011 Switzerland
Notes	

ISRCTN45300218

Trial name or title	SIPS Junior
Methods	Parallel design RCT
Participants	Setting: UK, adolescents (14 to 17 years) attending emergency department
Interventions	Smart phone or web-based electronic brief intervention or face-to-face personalised feedback Control group will receive treatment according to the injury that brought them to the emergency department
Outcomes	Total alcohol consumption in previous 28 days
Starting date	March 2014
Contact information	s.coulton@kent.ac.uk
Notes	

RBR-65262c

Trial name or title	Brief intervention for hazardous and harmful women's alcohol consumption
Methods	Parallel design RCT
Participants	Setting: Brazil, women (aged over 18 years) presenting to primary health care clinic who are screened as having risky or harmful alcohol consumption
Interventions	Brief alcohol intervention - two sessions of 20 to 30 minutes each
Outcomes	AUDIT score change
Starting date	1 June 2017
Contact information	Talita Dutra Ponce, talitadp@usp.br
Notes	

AUDIT: Alcohol Use Disorders Identification Test

DATA AND ANALYSES

Comparison 1. Brief intervention versus control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Quantity of drinking (g/week) at 12 months	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
2 Quantity of drinking (g/week) at 12 months, restricted to trials at low risk of bias due to allocation concealment	19	8856	Mean Difference (IV, Random, 95% CI)	-19.98 [-30.45, -9.51]
3 Quantity of drinking (g/week) at 12 months, restricted to trials at low risk of bias due to attrition	19	8496	Mean Difference (IV, Random, 95% CI)	-20.69 [-31.78, -9.60]
4 Quantity of drinking (g/week) at 12 months, with imputation of unknown standard deviations	39	17514	Mean Difference (IV, Random, 95% CI)	-19.35 [-26.90, -11.80]
5 Quantity of drinking (g/week) at 12 months, subgrouped by cluster/individual randomisation	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
5.1 Cluster-randomised	6	2588	Mean Difference (IV, Random, 95% CI)	-25.30 [-42.69, -7.91]
5.2 Individually randomised	28	12609	Mean Difference (IV, Random, 95% CI)	-18.77 [-28.01, -9.53]
6 Quantity of drinking (g/week) at 12 months, subgrouped by cluster/individual randomisation, varying imputed ICC	34	15197	Mean Difference (IV, Random, 95% CI)	-19.41 [-27.59, -11.22]
6.1 Cluster-randomised	6	2588	Mean Difference (IV, Random, 95% CI)	-20.32 [-33.50, -7.14]
6.2 Individually randomised	28	12609	Mean Difference (IV, Random, 95% CI)	-18.77 [-28.01, -9.53]
7 Quantity of drinking (g/week) at 12 months, subgrouped by gender	11	4836	Mean Difference (IV, Random, 95% CI)	-37.42 [-54.39, -20.45]
7.1 Men	11	3486	Mean Difference (IV, Random, 95% CI)	-42.21 [-64.53, -19.89]
7.2 Women	7	1350	Mean Difference (IV, Random, 95% CI)	-30.27 [-58.99, -1.55]
8 Quantity of drinking (g/week) at 12 months, subgrouped by gender, excluding trials of men only	7	4036	Mean Difference (IV, Random, 95% CI)	-33.12 [-51.47, -14.76]

8.1 Men	7	2686	Mean Difference (IV, Random, 95% CI)	-35.16 [-60.51, -9.80]
8.2 Women	7	1350	Mean Difference (IV, Random, 95% CI)	-30.27 [-58.99, -1.55]
9 Quantity of drinking (g/week) at 12 months, subgrouped by adolescents/young adults only versus other	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
9.1 Adolescents/young adults only	3	1638	Mean Difference (IV, Random, 95% CI)	-7.09 [-17.18, 2.99]
9.2 Other trials	31	13559	Mean Difference (IV, Random, 95% CI)	-22.79 [-32.42, -13.16]
10 Quantity of drinking (g/week) at 12 months, subgrouped by general practice/emergency setting	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
10.1 General practice setting	24	8811	Mean Difference (IV, Random, 95% CI)	-25.69 [-37.30, -14.08]
10.2 Emergency care setting	10	6386	Mean Difference (IV, Random, 95% CI)	-9.73 [-17.52, -1.93]
11 Quantity of drinking (g/wk) at 12 months, subgrouped by intervention modality (advice versus counselling)	32	13780	Mean Difference (IV, Random, 95% CI)	-21.01 [-30.06, -11.96]
11.1 Advice-based	20	8243	Mean Difference (IV, Random, 95% CI)	-32.68 [-45.75, -19.60]
11.2 Counselling-based	12	5537	Mean Difference (IV, Random, 95% CI)	-0.17 [-2.96, 2.61]
12 Quantity of drinking (g/week) at 12 months, subgrouped by type of control	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
12.1 No alcohol content	18	8606	Mean Difference (IV, Random, 95% CI)	-24.41 [-36.47, -12.35]
12.2 Some alcohol content	16	6591	Mean Difference (IV, Random, 95% CI)	-12.80 [-22.93, -2.66]
13 Quantity of drinking (g/week), subgrouped by length of follow-up	38		Mean Difference (IV, Random, 95% CI)	Subtotals only
13.1 At 6 months	21	10313	Mean Difference (IV, Random, 95% CI)	-21.56 [-31.56, -11.55]
13.2 At 8 to 9 months	2	352	Mean Difference (IV, Random, 95% CI)	-15.70 [-34.19, 2.79]
13.3 At 12 months	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
13.4 At 36 months	1	277	Mean Difference (IV, Random, 95% CI)	-6.11 [-79.55, 67.33]
14 Quantity of drinking (g/week), subgrouped by length of follow-up (6 and 12 months) and restricted to trials with information at both times	18		Mean Difference (IV, Random, 95% CI)	Subtotals only
14.1 At 6 months	18	9739	Mean Difference (IV, Random, 95% CI)	-21.28 [-31.82, -10.75]

14.2 At 12 months	18	9645	Mean Difference (IV, Random, 95% CI)	-22.25 [-34.04, -10.47]
15 Quantity of drinking (g/week) at 12 months, subgrouped by effectiveness/efficacy	34	15197	Mean Difference (IV, Random, 95% CI)	-20.08 [-28.36, -11.81]
15.1 Effectiveness trials	16	7091	Mean Difference (IV, Random, 95% CI)	-27.06 [-41.49, -12.63]
15.2 Efficacy trials	18	8106	Mean Difference (IV, Random, 95% CI)	-14.37 [-23.97, -4.76]
16 Frequency of binge drinking (number binges/week) at 12 months	15	6946	Mean Difference (IV, Random, 95% CI)	-0.08 [-0.14, -0.02]
17 Frequency of drinking (number binges/week), subgrouped by length of follow-up (6 & 12 months) and restricted to trials with information at both times	9		Mean Difference (IV, Random, 95% CI)	Subtotals only
17.1 At 6 months	9	4354	Mean Difference (IV, Random, 95% CI)	-0.15 [-0.22, -0.08]
17.2 At 12 months	9	4333	Mean Difference (IV, Random, 95% CI)	-0.12 [-0.23, -0.02]
18 Frequency of drinking (number days drinking/week) at 12 months	11	5469	Mean Difference (IV, Random, 95% CI)	-0.13 [-0.23, -0.04]
19 Frequency of drinking (number days drinking/week), subgrouped by length of follow-up (6 & 12 months) and restricted to trials with information at both times	6		Mean Difference (IV, Random, 95% CI)	Subtotals only
19.1 At 6 months	6	3637	Mean Difference (IV, Random, 95% CI)	-0.12 [-0.24, 0.00]
19.2 At 12 months	6	3658	Mean Difference (IV, Random, 95% CI)	-0.16 [-0.28, -0.04]
20 Intensity of drinking (g/drinking day) at 12 months	10	3128	Mean Difference (IV, Random, 95% CI)	-0.18 [-3.09, 2.73]
21 Heavy drinkers at 12 months	18	7623	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.13, -0.04]
22 Binge drinkers at 12 months	10	4456	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.12, -0.02]
23 Laboratory markers - GGT (IU/L) at 12 months	3	1166	Mean Difference (IV, Random, 95% CI)	-0.89 [-3.86, 2.08]
24 Laboratory markers - GGT (IU/L) at 12 months, subgrouped by gender	2	1094	Mean Difference (IV, Random, 95% CI)	-0.03 [-2.52, 2.46]
24.1 Men	2	764	Mean Difference (IV, Random, 95% CI)	-2.08 [-6.10, 1.95]
24.2 Women	2	330	Mean Difference (IV, Random, 95% CI)	1.23 [-1.94, 4.40]

Comparison 2. Extended intervention versus minimal or no intervention

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Quantity of drinking (g/week) at 12 months	6	1296	Mean Difference (IV, Random, 95% CI)	-14.43 [-37.41, 8.54]
2 Frequency of binge drinking (number binges/week) at 12 months	2	456	Mean Difference (IV, Random, 95% CI)	-0.08 [-0.28, 0.12]
3 Frequency of drinking (number days drinking/week) at 12 months	2	319	Mean Difference (IV, Random, 95% CI)	-0.45 [-0.81, -0.09]
4 Intensity of drinking (g/drinking day) at 12 months	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
5 Heavy drinkers at 12 months	1		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6 Binge drinkers at 12 months	2	777	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.07, 0.03]
7 Laboratory markers - GGT (IU/L) at 12 months	1		Mean Difference (IV, Random, 95% CI)	Subtotals only

Comparison 3. Extended versus brief intervention

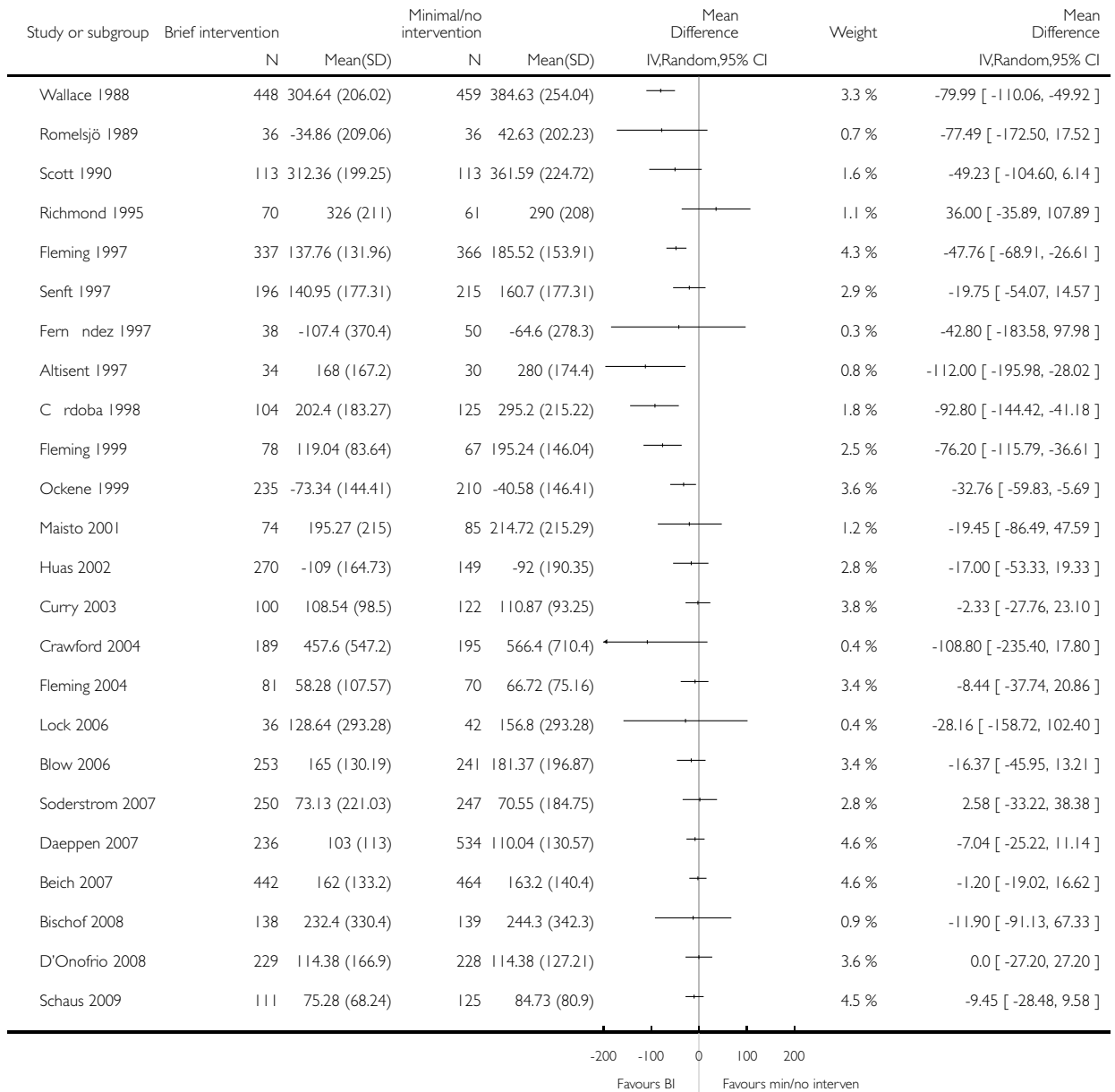
Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Quantity of drinking (g/week) at 12 months	3	552	Mean Difference (IV, Random, 95% CI)	1.54 [-42.01, 45.10]
2 Quantity of drinking (g/week) at 12 months, with imputation of unknown standard deviations	4	635	Mean Difference (IV, Random, 95% CI)	1.13 [-37.00, 41.26]
3 Frequency of drinking (number days drinking/week) at 12 months	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
4 Intensity of drinking (g/drinking day)	1		Mean Difference (IV, Random, 95% CI)	Subtotals only
5 Binge drinkers at 12 months	2	339	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.08, 0.12]
6 Laboratory markers - MCV (fl) at 12 months	1	95	Mean Difference (IV, Random, 95% CI)	0.62 [-1.58, 2.81]
6.1 Men	1	81	Mean Difference (IV, Random, 95% CI)	0.80 [-1.70, 3.30]
6.2 Women	1	14	Mean Difference (IV, Random, 95% CI)	0.0 [-4.57, 4.57]

Analysis 1.1. Comparison 1 Brief intervention versus control, Outcome 1 Quantity of drinking (g/week) at 12 months.

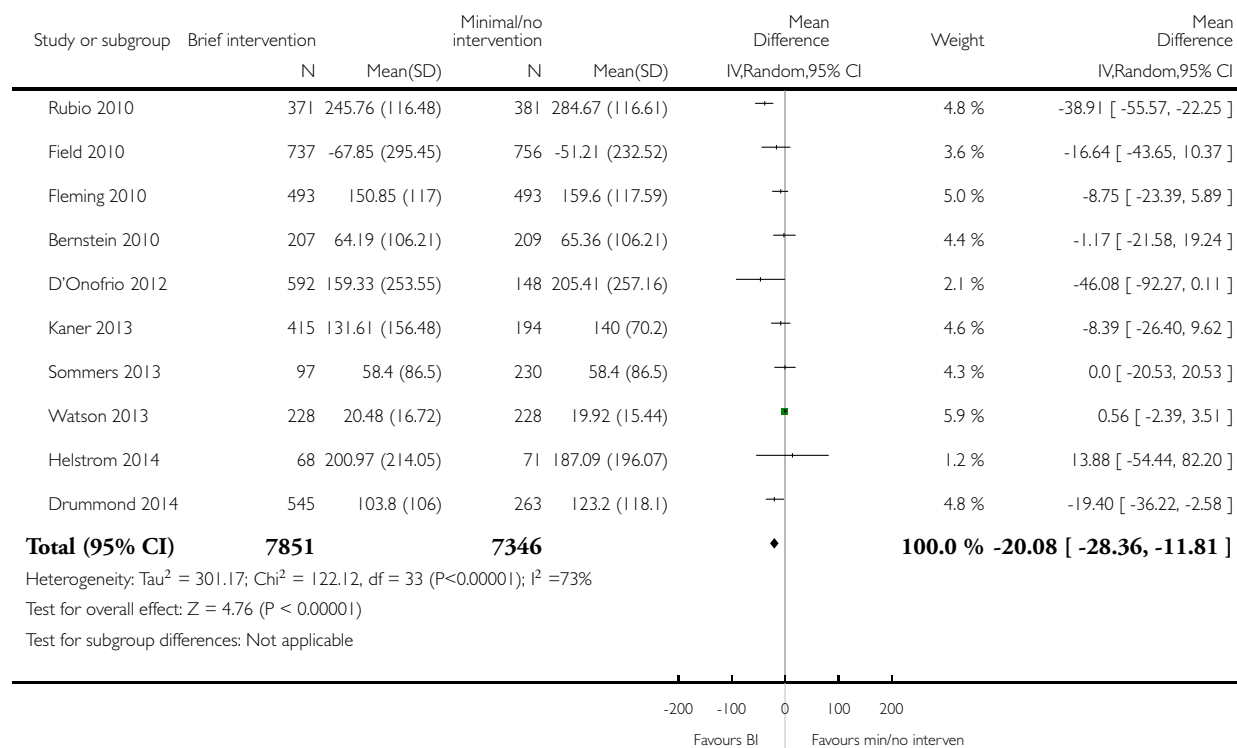
Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 1 Quantity of drinking (g/week) at 12 months



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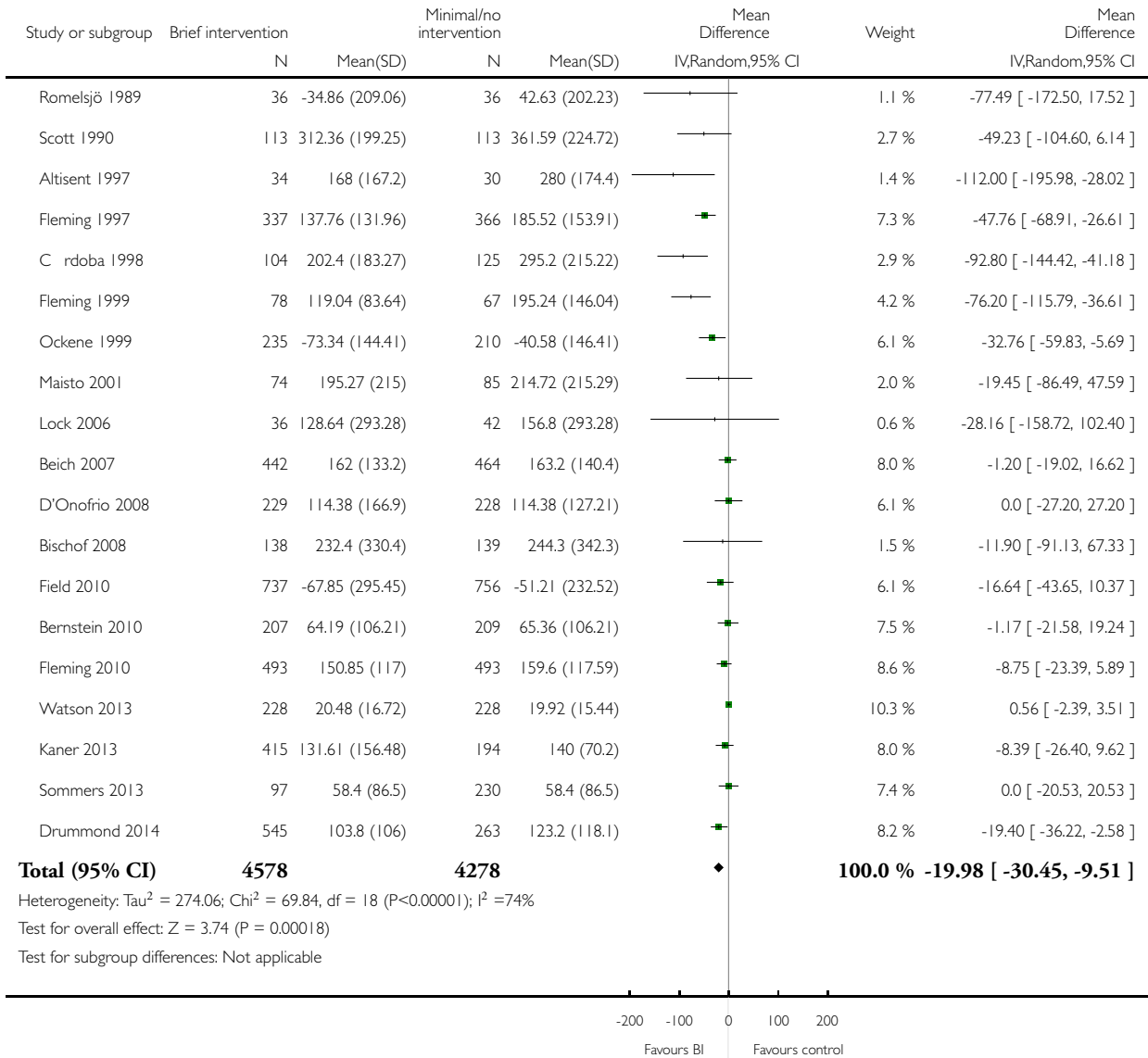


Analysis 1.2. Comparison 1 Brief intervention versus control, Outcome 2 Quantity of drinking (g/week) at 12 months, restricted to trials at low risk of bias due to allocation concealment.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 2 Quantity of drinking (g/week) at 12 months, restricted to trials at low risk of bias due to allocation concealment

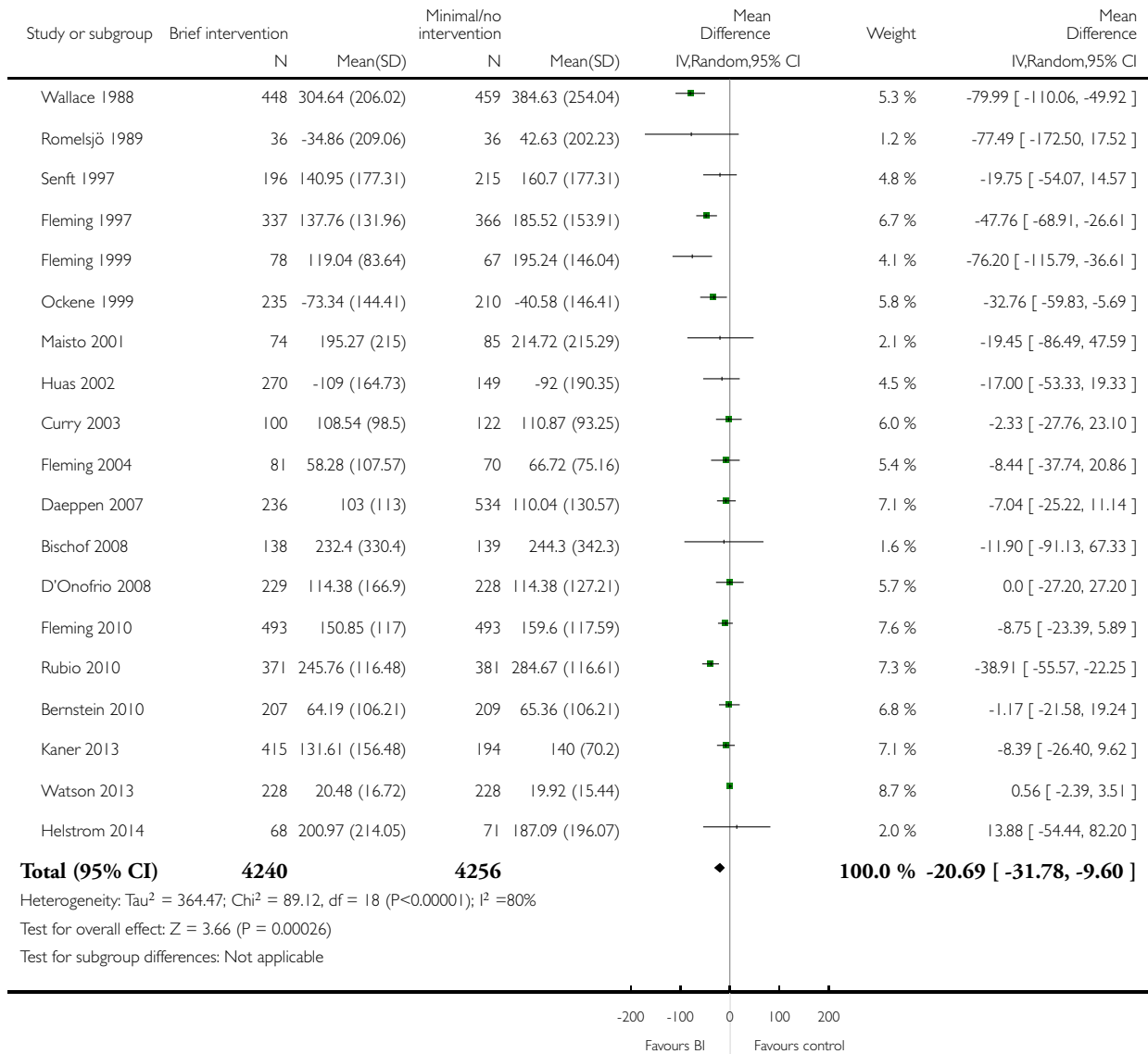


Analysis 1.3. Comparison 1 Brief intervention versus control, Outcome 3 Quantity of drinking (g/week) at 12 months, restricted to trials at low risk of bias due to attrition.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 3 Quantity of drinking (g/week) at 12 months, restricted to trials at low risk of bias due to attrition

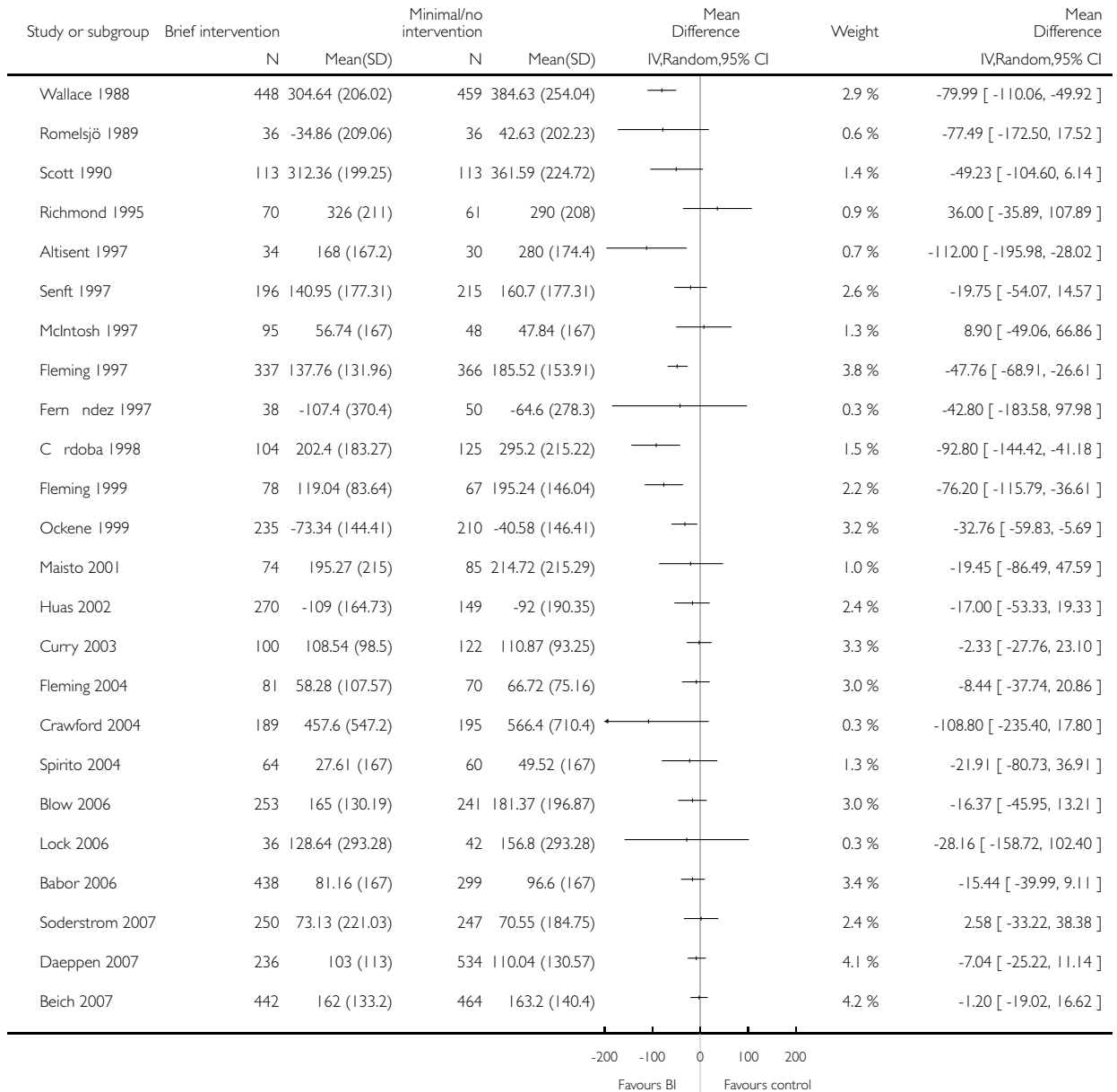


Analysis 1.4. Comparison 1 Brief intervention versus control, Outcome 4 Quantity of drinking (g/week) at 12 months, with imputation of unknown standard deviations.

Review: Effectiveness of brief alcohol interventions in primary care populations

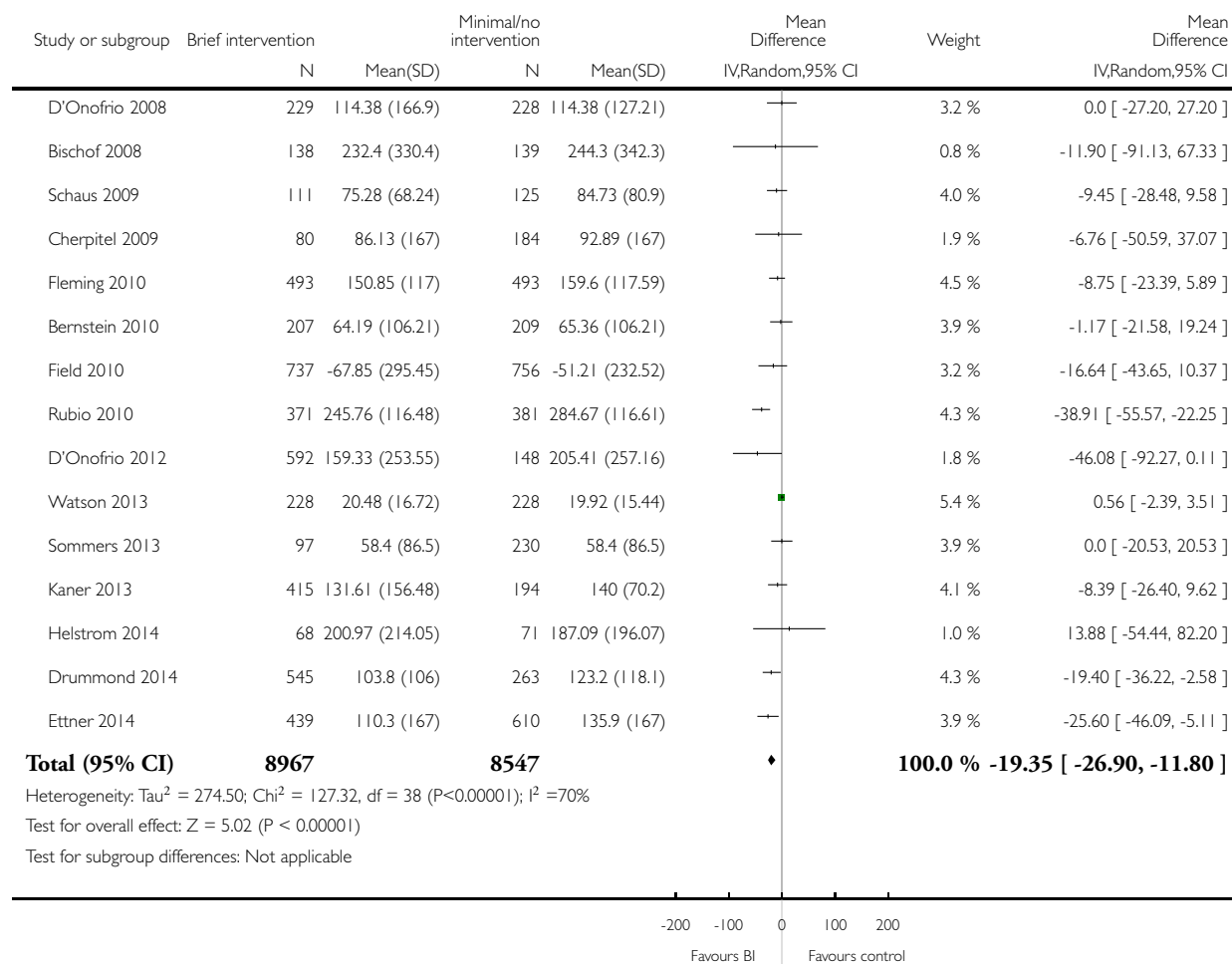
Comparison: 1 Brief intervention versus control

Outcome: 4 Quantity of drinking (g/week) at 12 months, with imputation of unknown standard deviations



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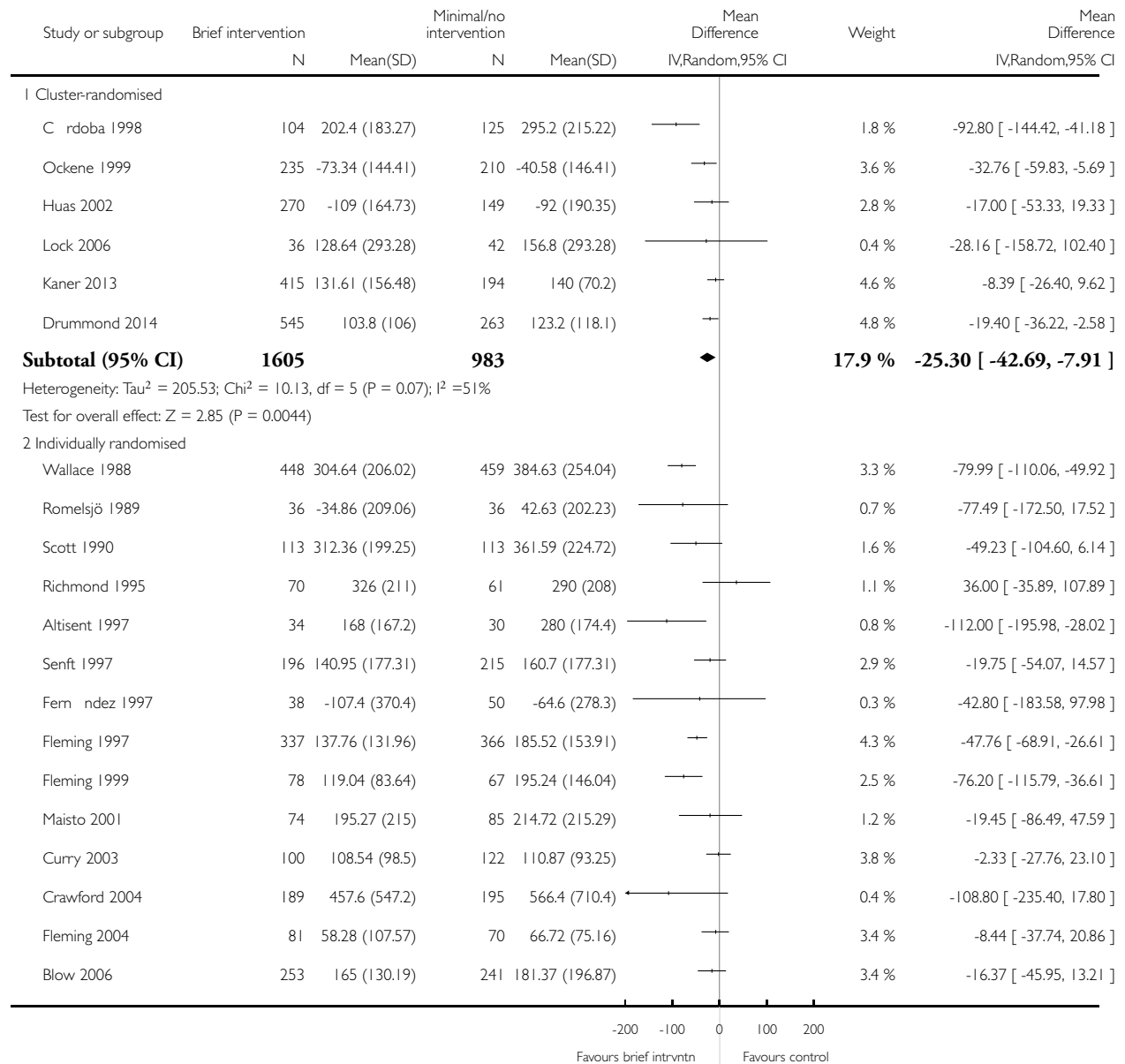


Analysis 1.5. Comparison 1 Brief intervention versus control, Outcome 5 Quantity of drinking (g/week) at 12 months, subgrouped by cluster/individual randomisation.

Review: Effectiveness of brief alcohol interventions in primary care populations

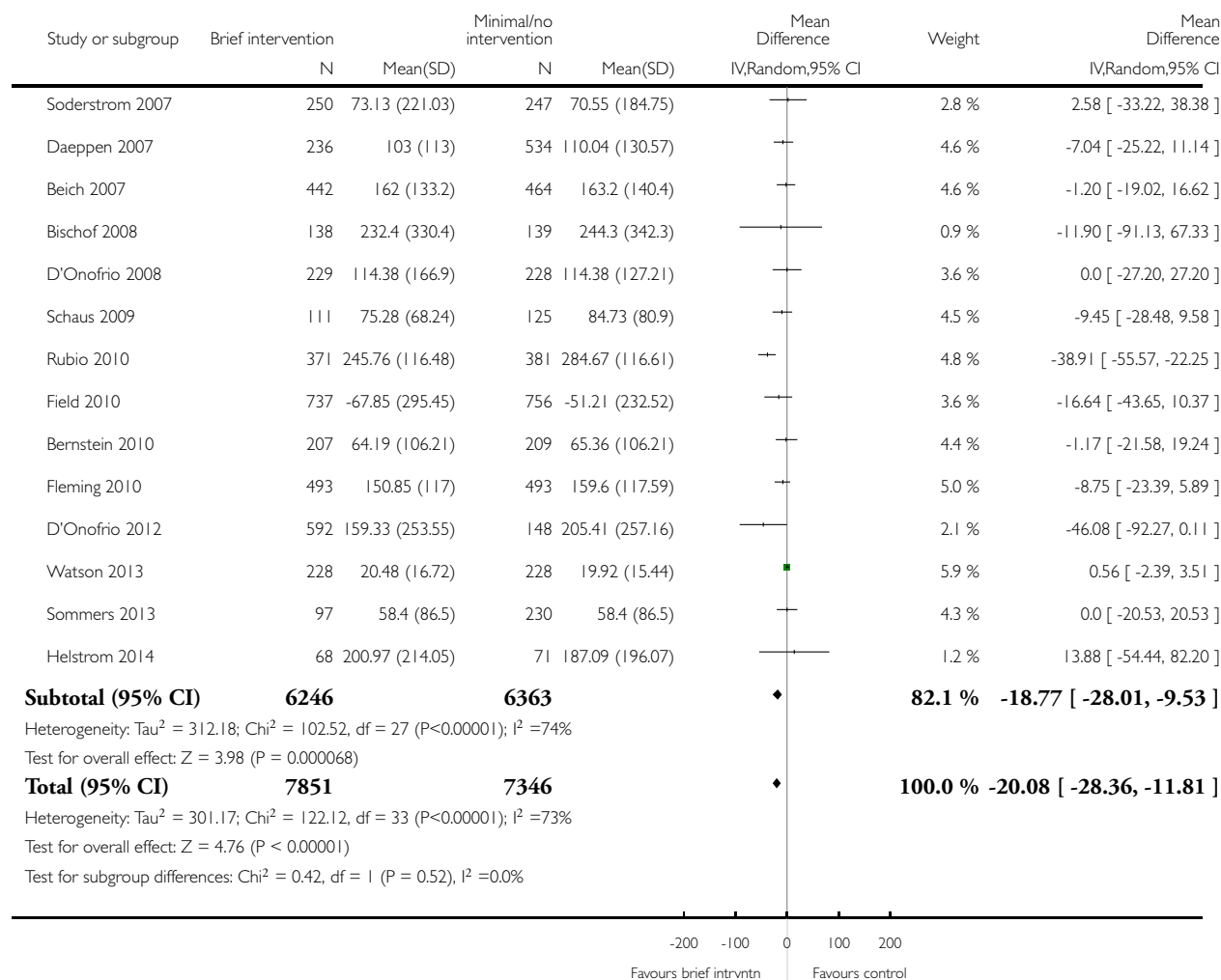
Comparison: 1 Brief intervention versus control

Outcome: 5 Quantity of drinking (g/week) at 12 months, subgrouped by cluster/individual randomisation



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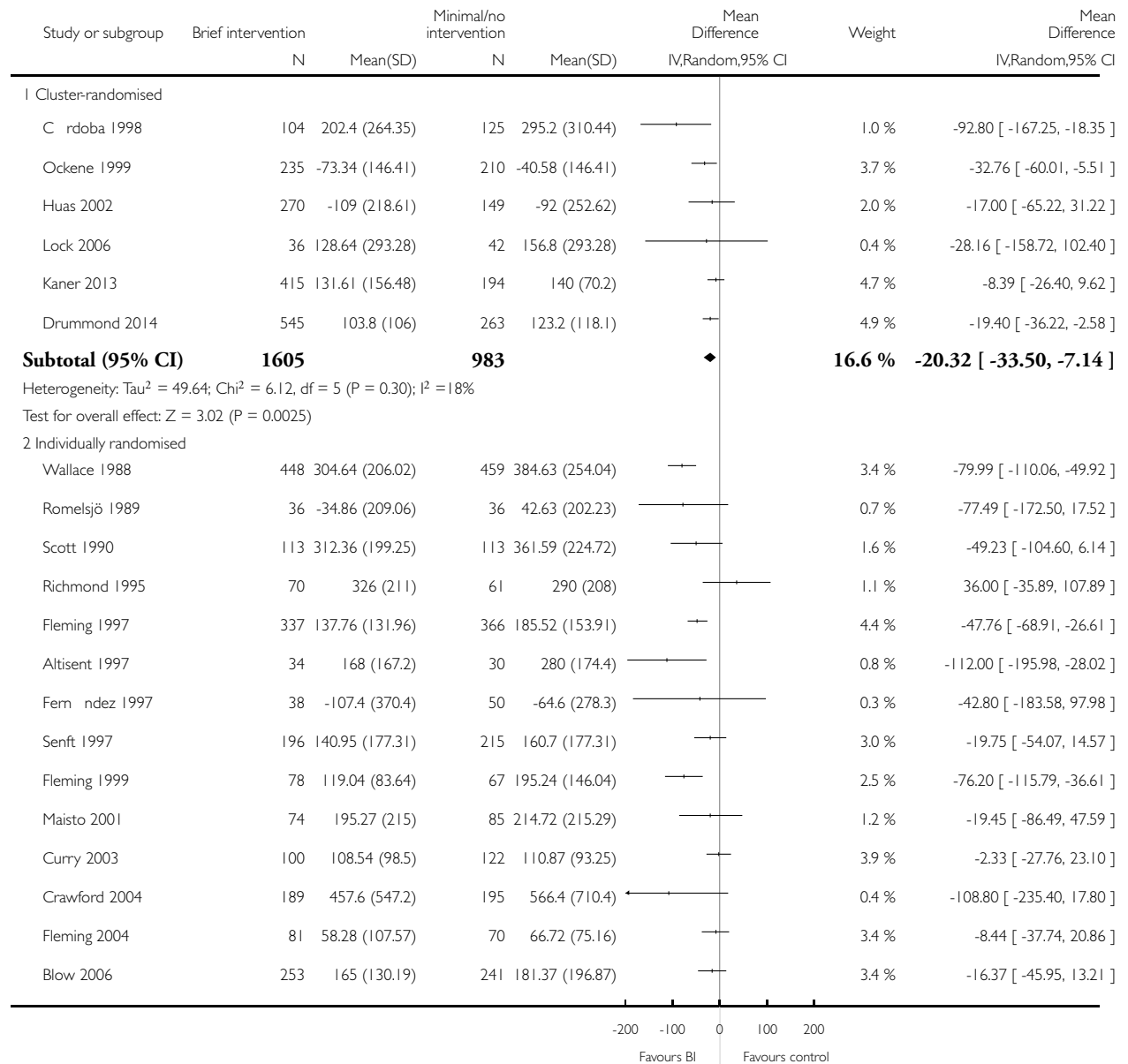


Analysis 1.6. Comparison 1 Brief intervention versus control, Outcome 6 Quantity of drinking (g/week) at 12 months, subgrouped by cluster/individual randomisation, varying imputed ICC.

Review: Effectiveness of brief alcohol interventions in primary care populations

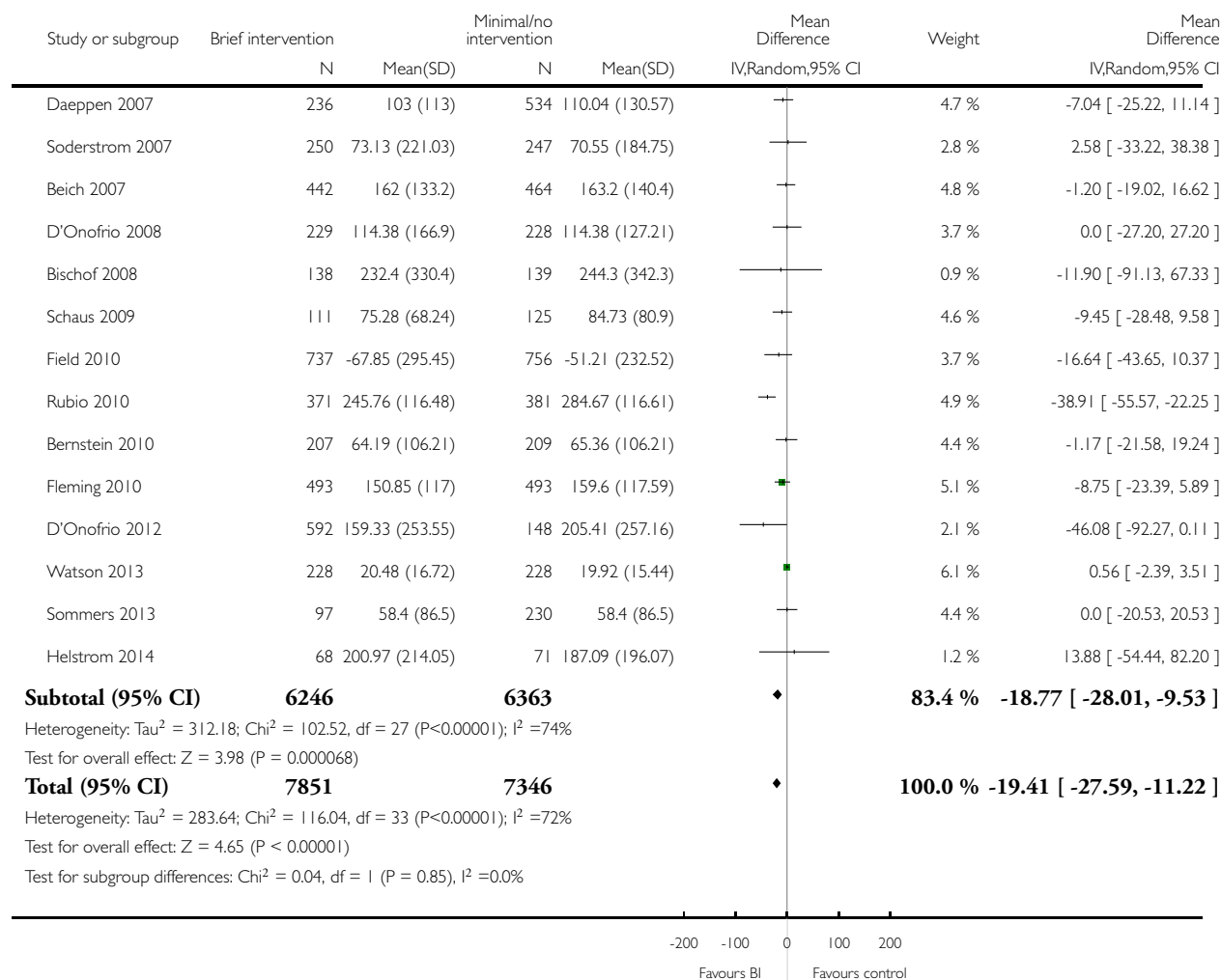
Comparison: 1 Brief intervention versus control

Outcome: 6 Quantity of drinking (g/week) at 12 months, subgrouped by cluster/individual randomisation, varying imputed ICC



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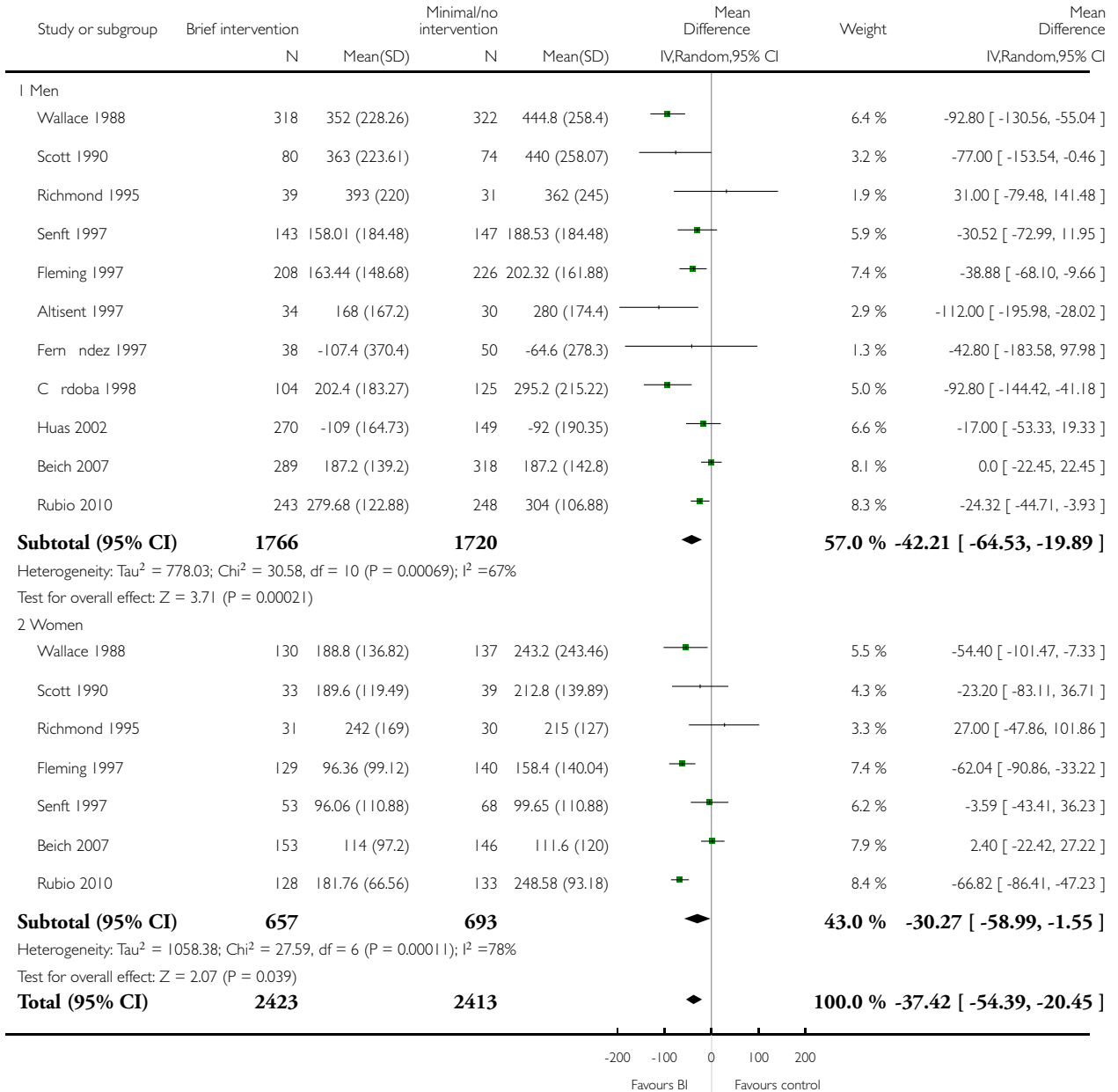


Analysis I.7. Comparison I Brief intervention versus control, Outcome 7 Quantity of drinking (g/week) at 12 months, subgrouped by gender.

Review: Effectiveness of brief alcohol interventions in primary care populations

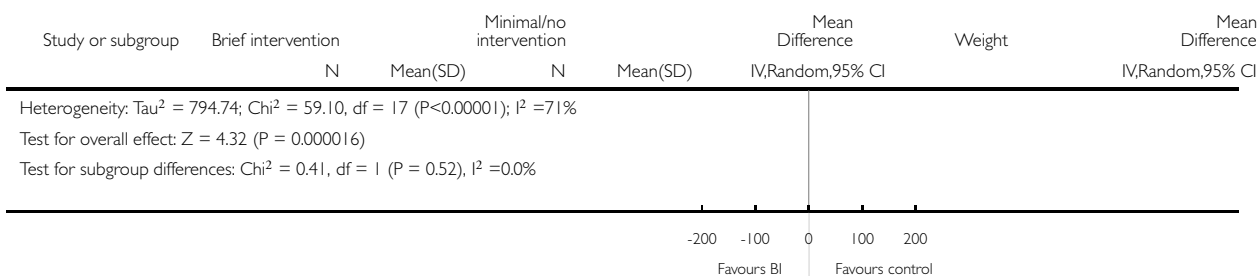
Comparison: I Brief intervention versus control

Outcome: 7 Quantity of drinking (g/week) at 12 months, subgrouped by gender



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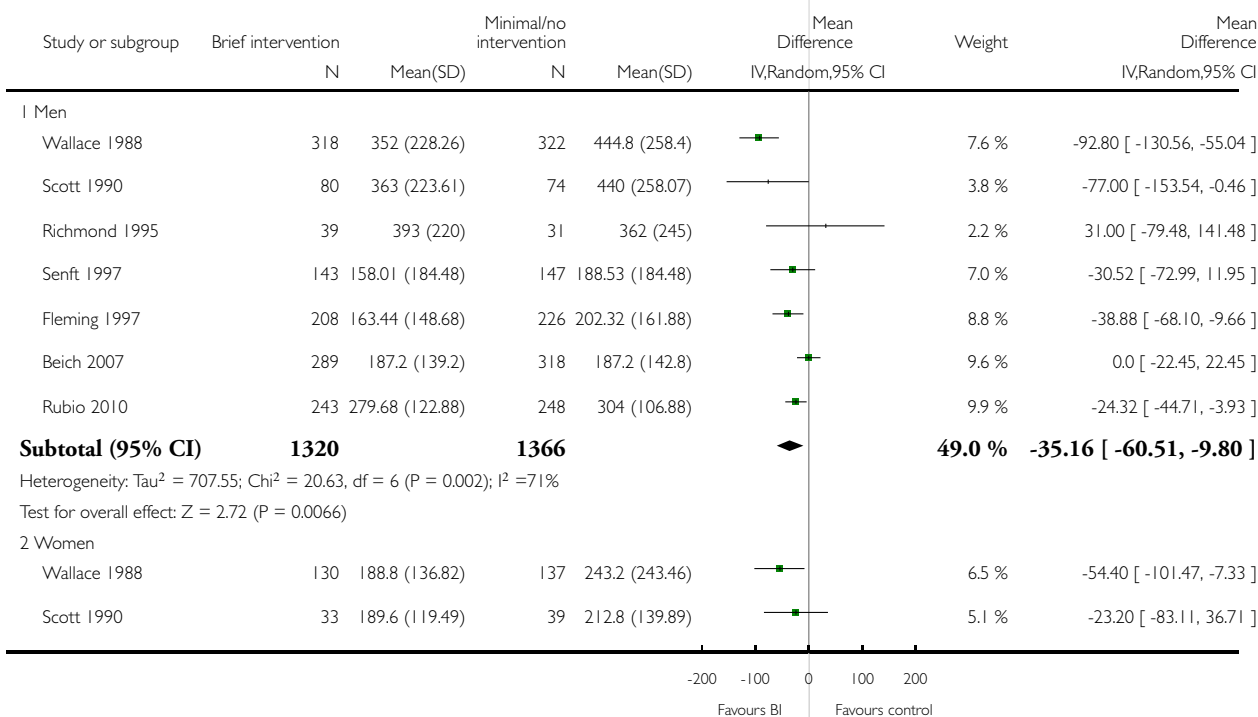


Analysis 1.8. Comparison 1 Brief intervention versus control, Outcome 8 Quantity of drinking (g/week) at 12 months, subgrouped by gender, excluding trials of men only.

Review: Effectiveness of brief alcohol interventions in primary care populations

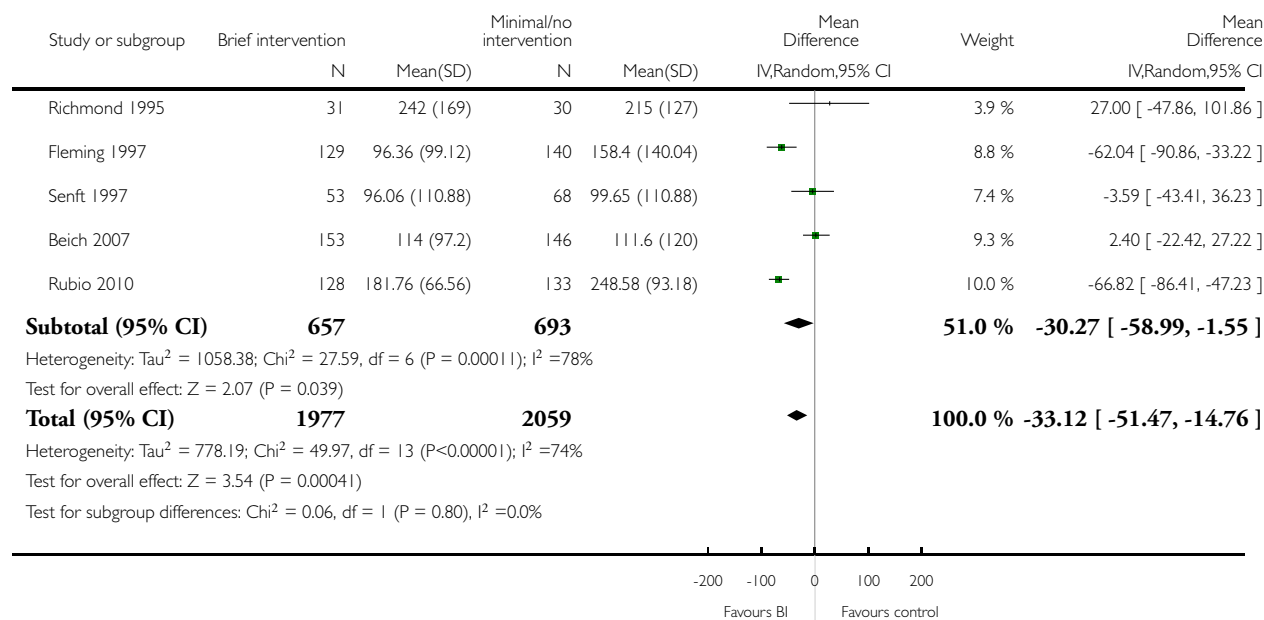
Comparison: 1 Brief intervention versus control

Outcome: 8 Quantity of drinking (g/week) at 12 months, subgrouped by gender, excluding trials of men only



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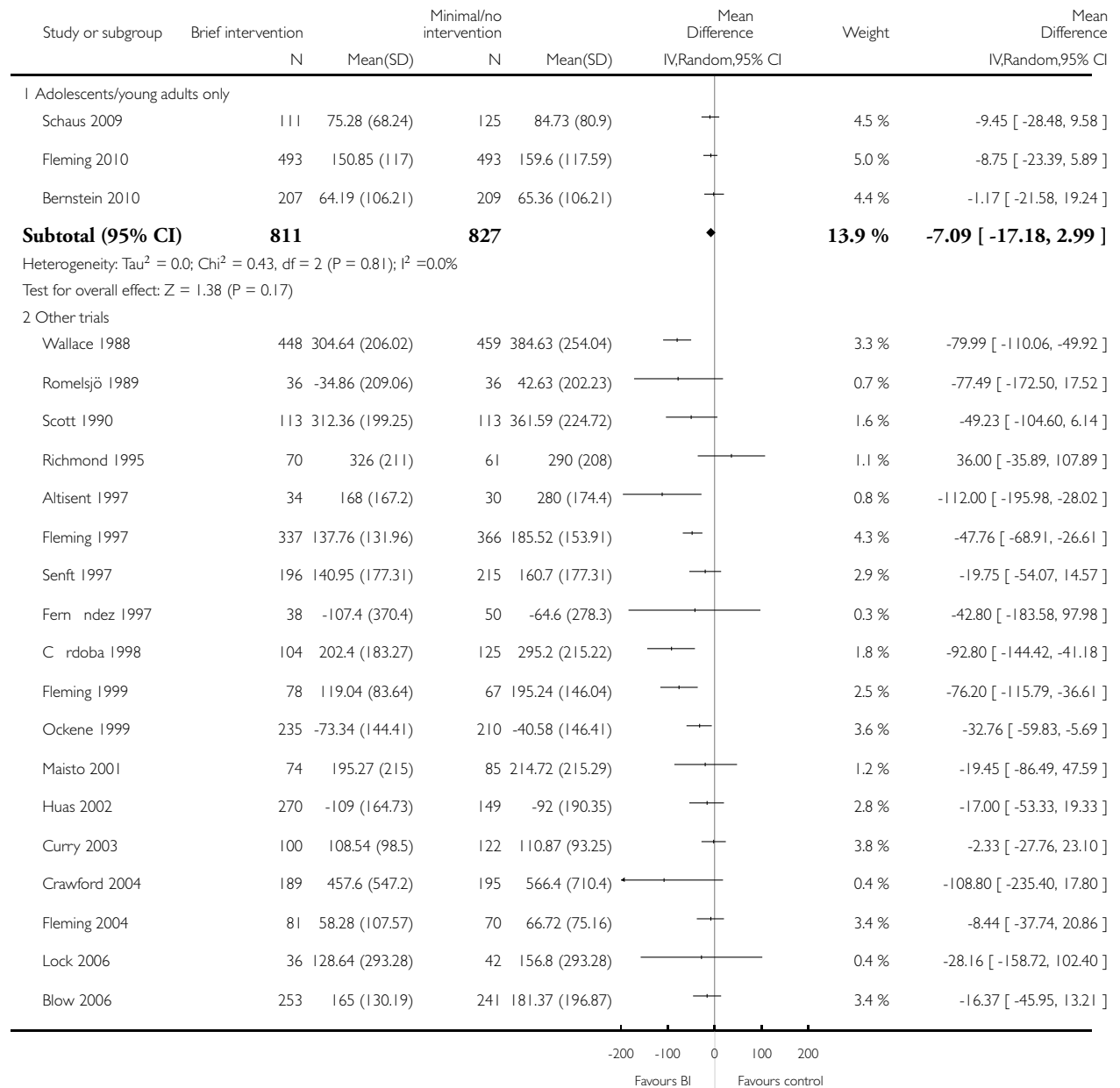


Analysis 1.9. Comparison 1 Brief intervention versus control, Outcome 9 Quantity of drinking (g/week) at 12 months, subgrouped by adolescents/young adults only versus other.

Review: Effectiveness of brief alcohol interventions in primary care populations

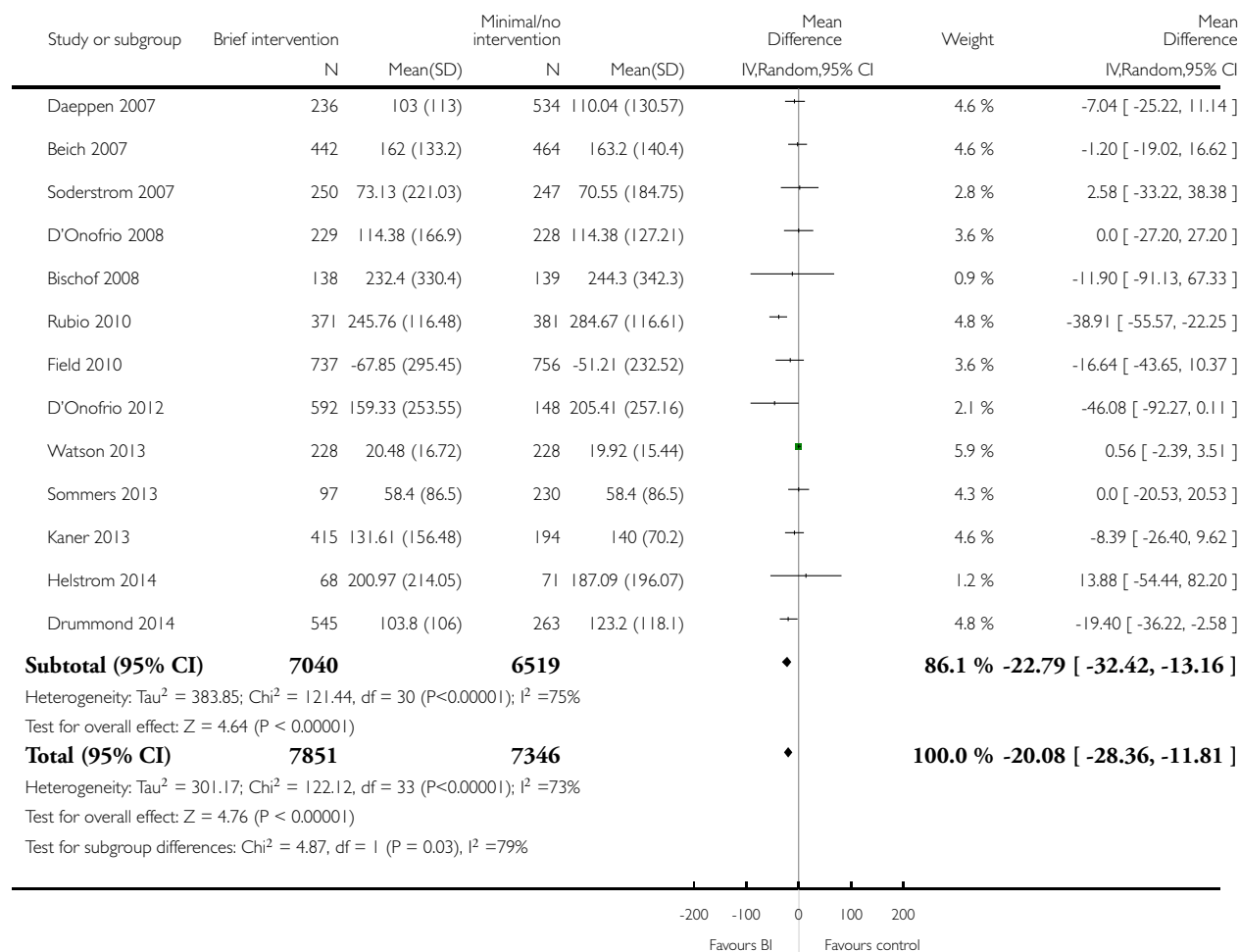
Comparison: 1 Brief intervention versus control

Outcome: 9 Quantity of drinking (g/week) at 12 months, subgrouped by adolescents/young adults only versus other



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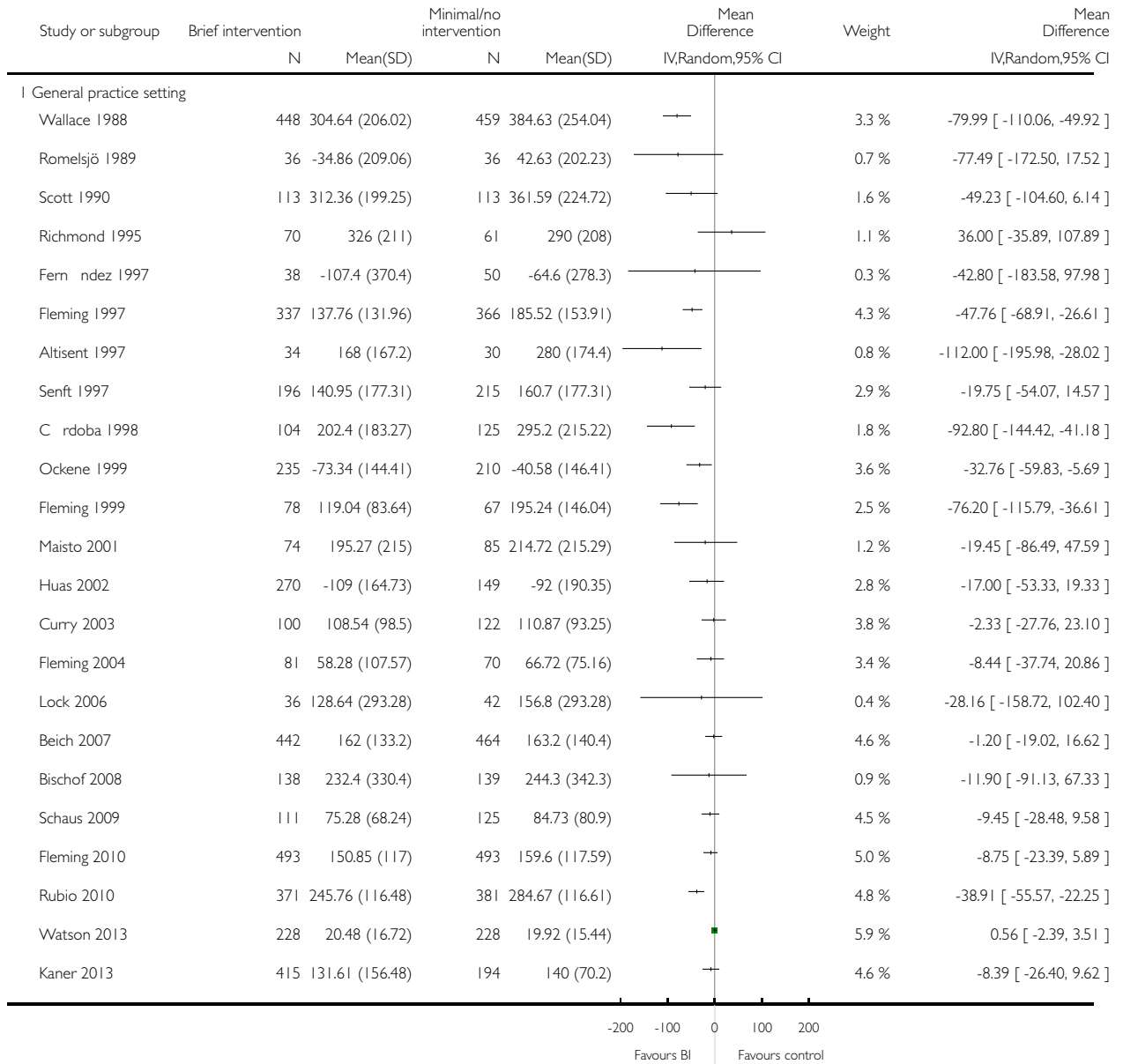


Analysis I.10. Comparison I Brief intervention versus control, Outcome 10 Quantity of drinking (g/week) at 12 months, subgrouped by general practice/emergency setting.

Review: Effectiveness of brief alcohol interventions in primary care populations

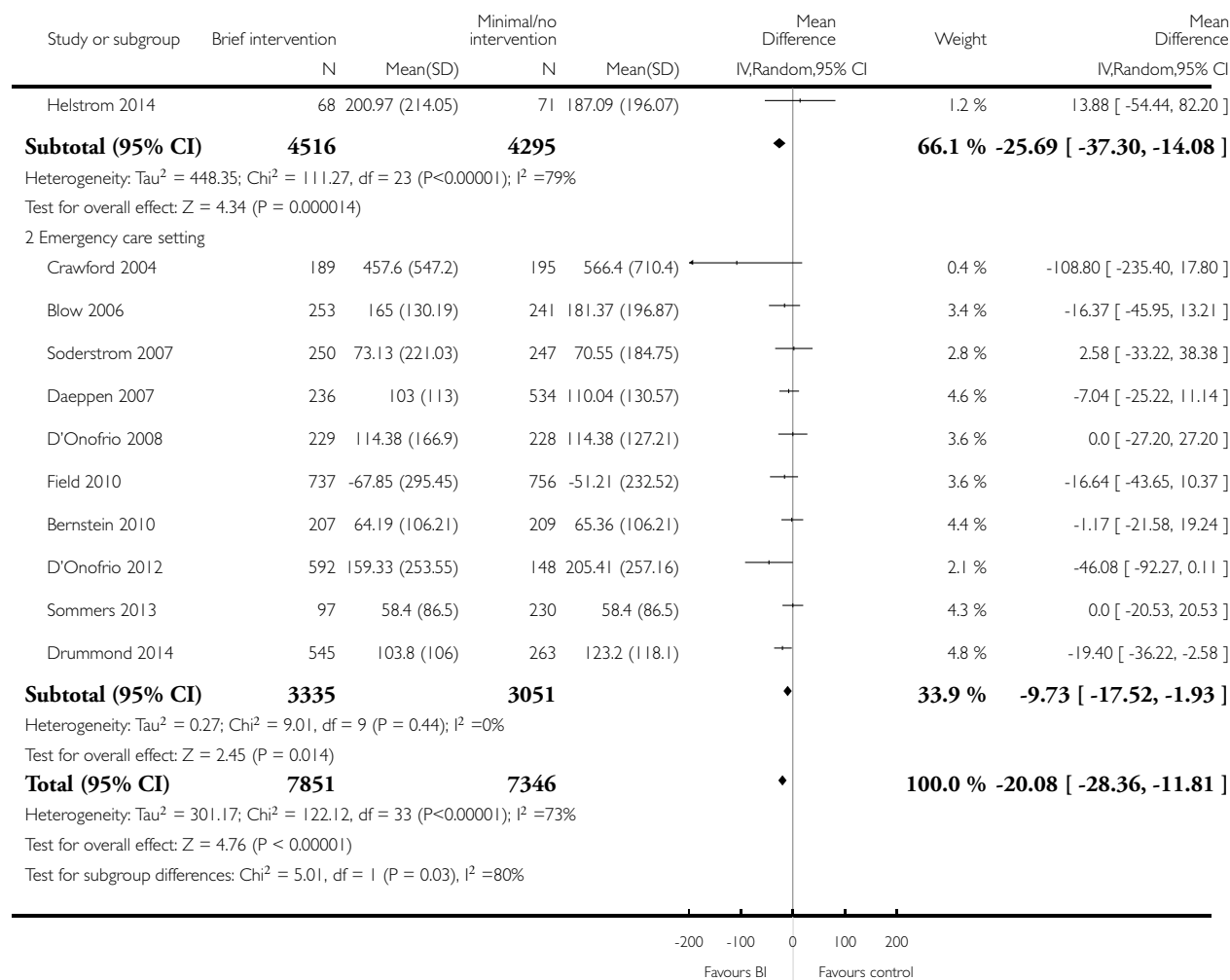
Comparison: I Brief intervention versus control

Outcome: 10 Quantity of drinking (g/week) at 12 months, subgrouped by general practice/emergency setting



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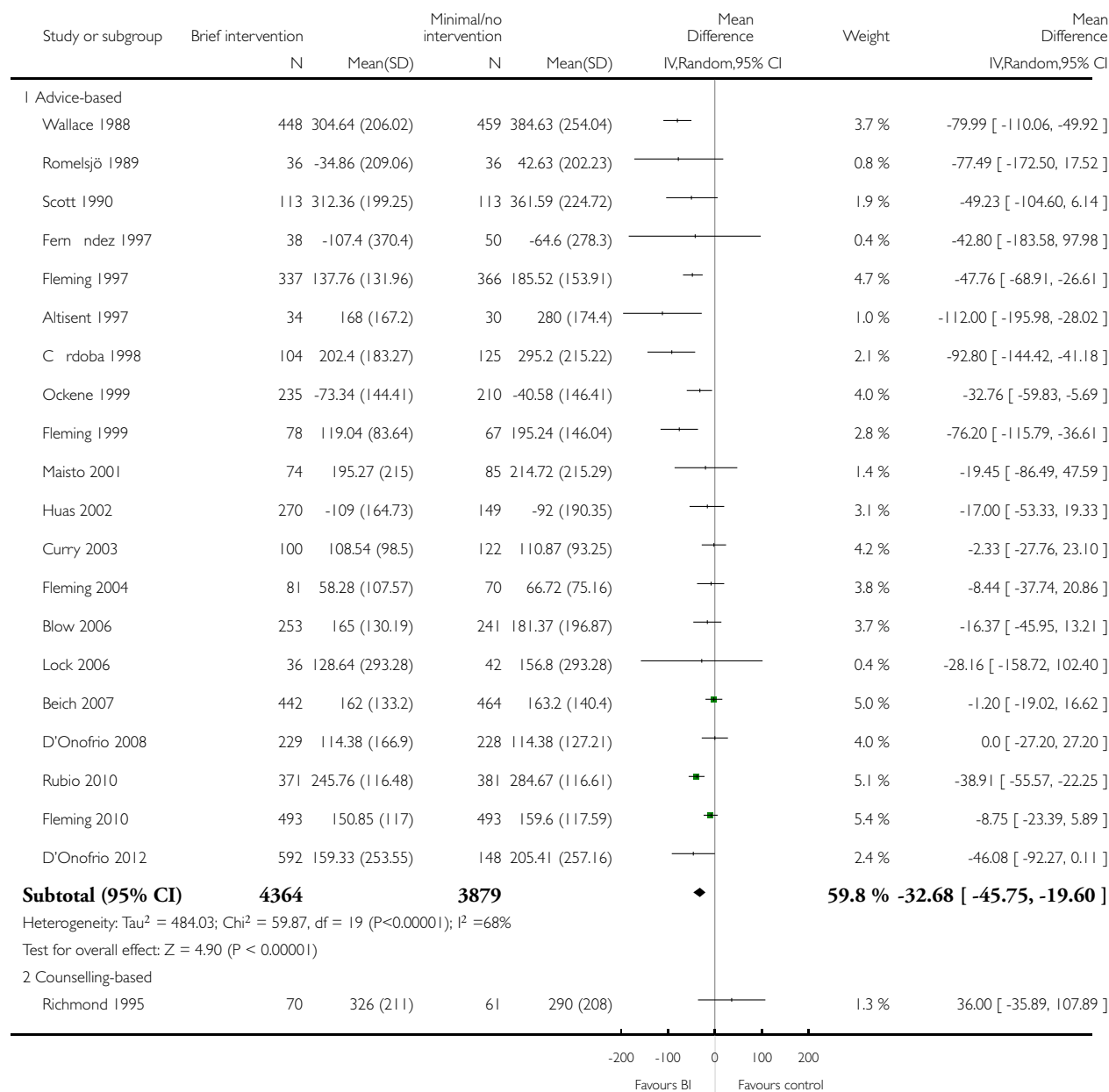


Analysis 1.11. Comparison 1 Brief intervention versus control, Outcome 11 Quantity of drinking (g/wk) at 12 months, subgrouped by intervention modality (advice versus counselling).

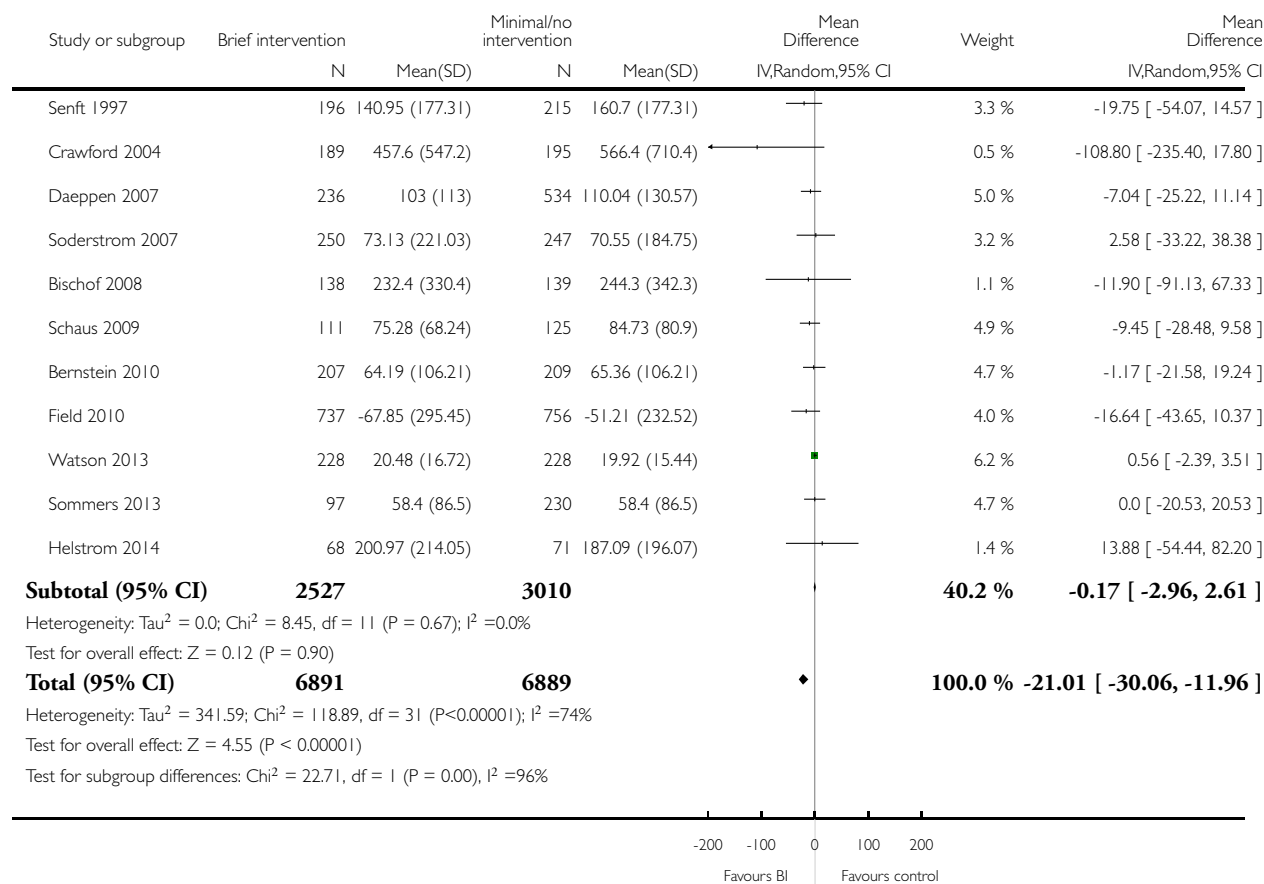
Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 11 Quantity of drinking (g/wk) at 12 months, subgrouped by intervention modality (advice versus counselling)



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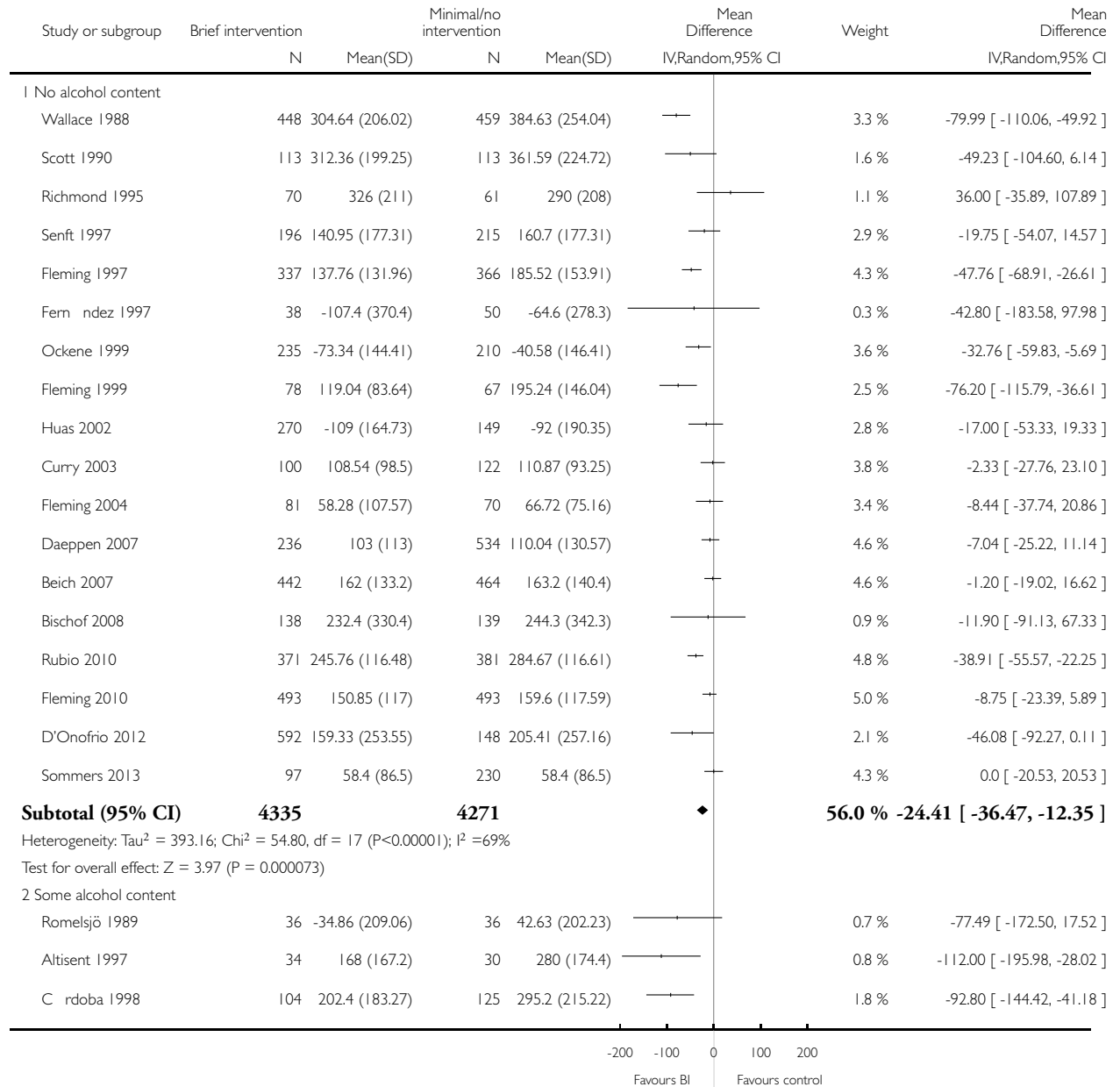


Analysis I.12. Comparison I Brief intervention versus control, Outcome 12 Quantity of drinking (g/week) at 12 months, subgrouped by type of control.

Review: Effectiveness of brief alcohol interventions in primary care populations

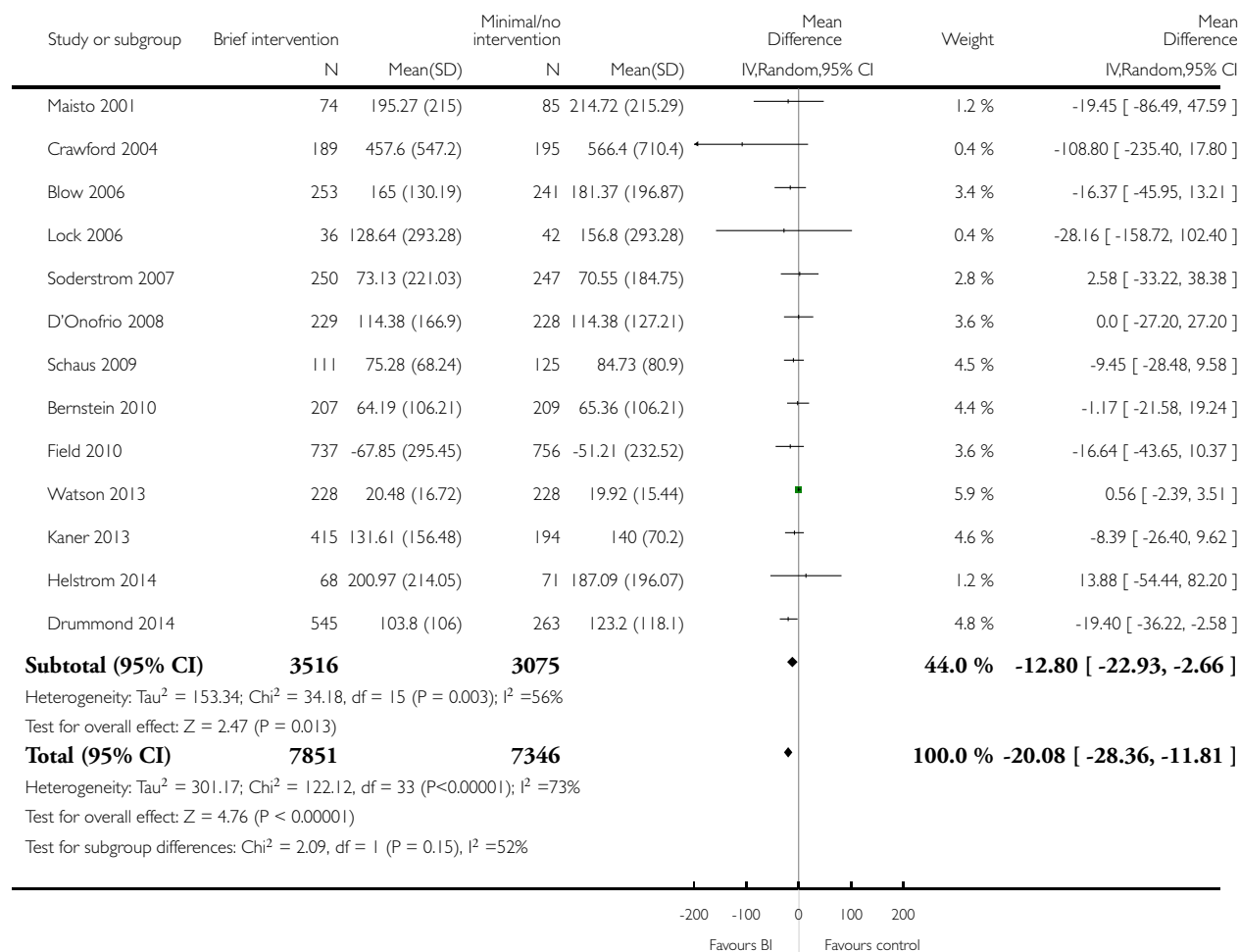
Comparison: I Brief intervention versus control

Outcome: 12 Quantity of drinking (g/week) at 12 months, subgrouped by type of control



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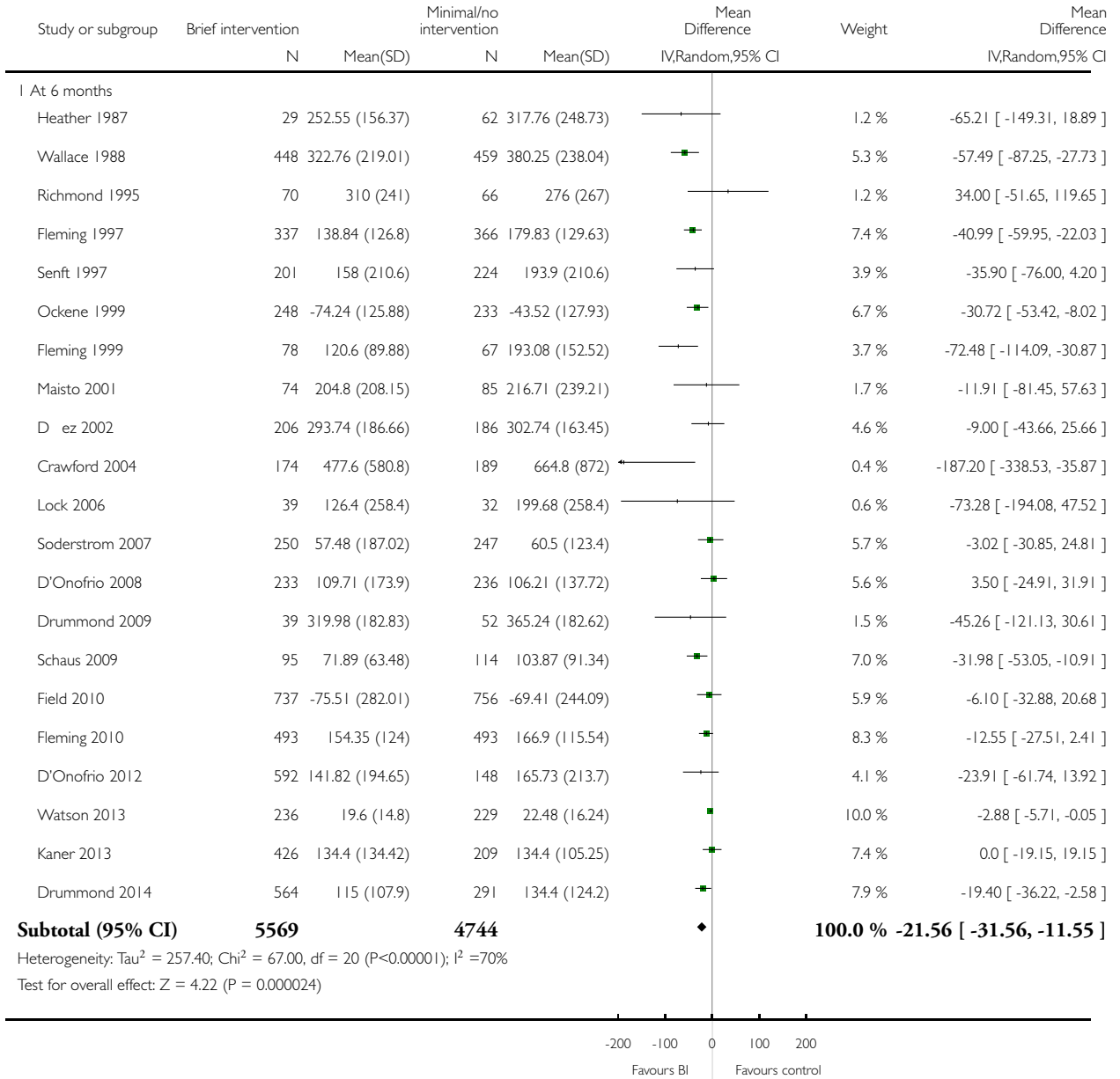


Analysis 1.13. Comparison 1 Brief intervention versus control, Outcome 13 Quantity of drinking (g/week), subgrouped by length of follow-up.

Review: Effectiveness of brief alcohol interventions in primary care populations

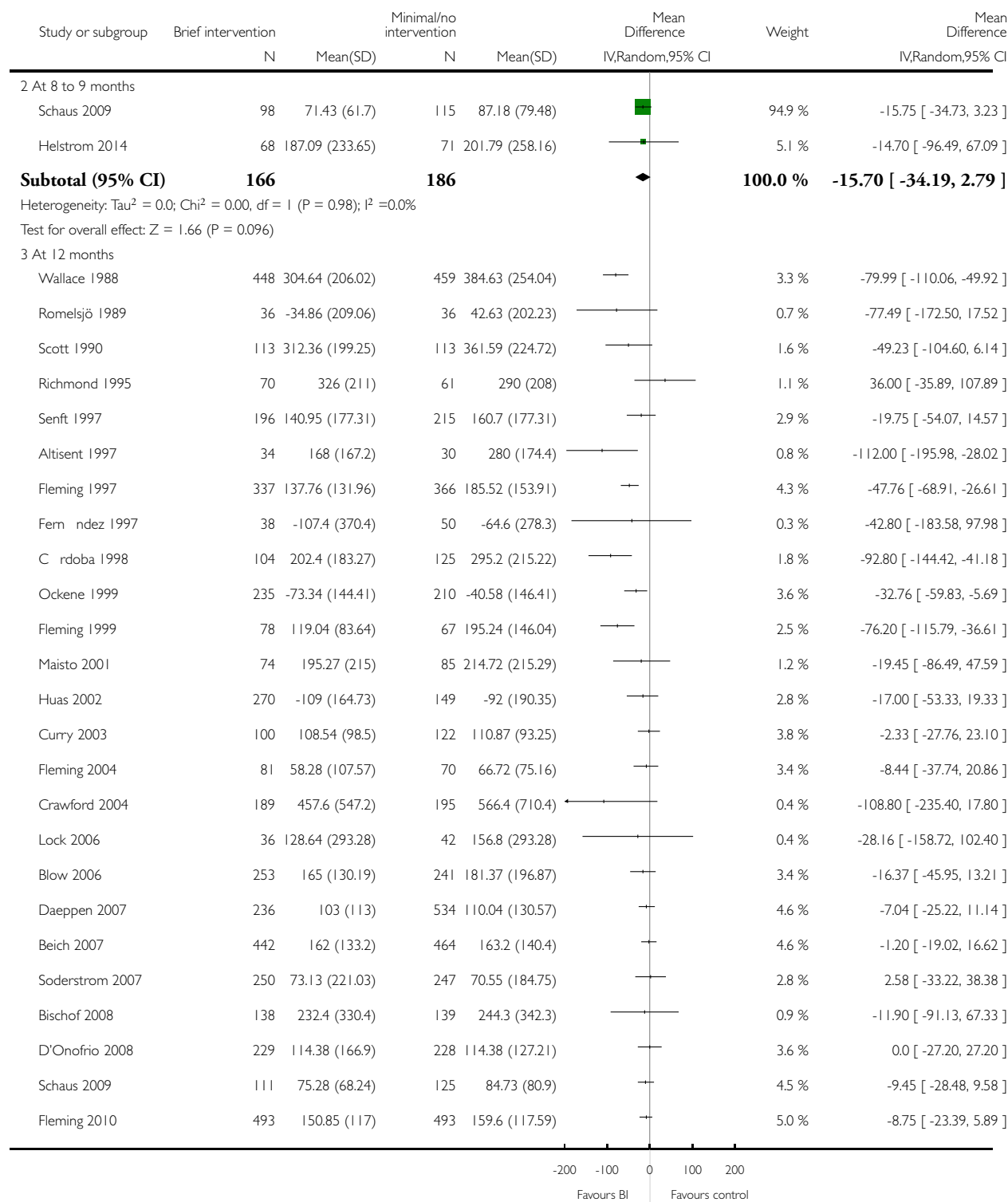
Comparison: 1 Brief intervention versus control

Outcome: 13 Quantity of drinking (g/week), subgrouped by length of follow-up



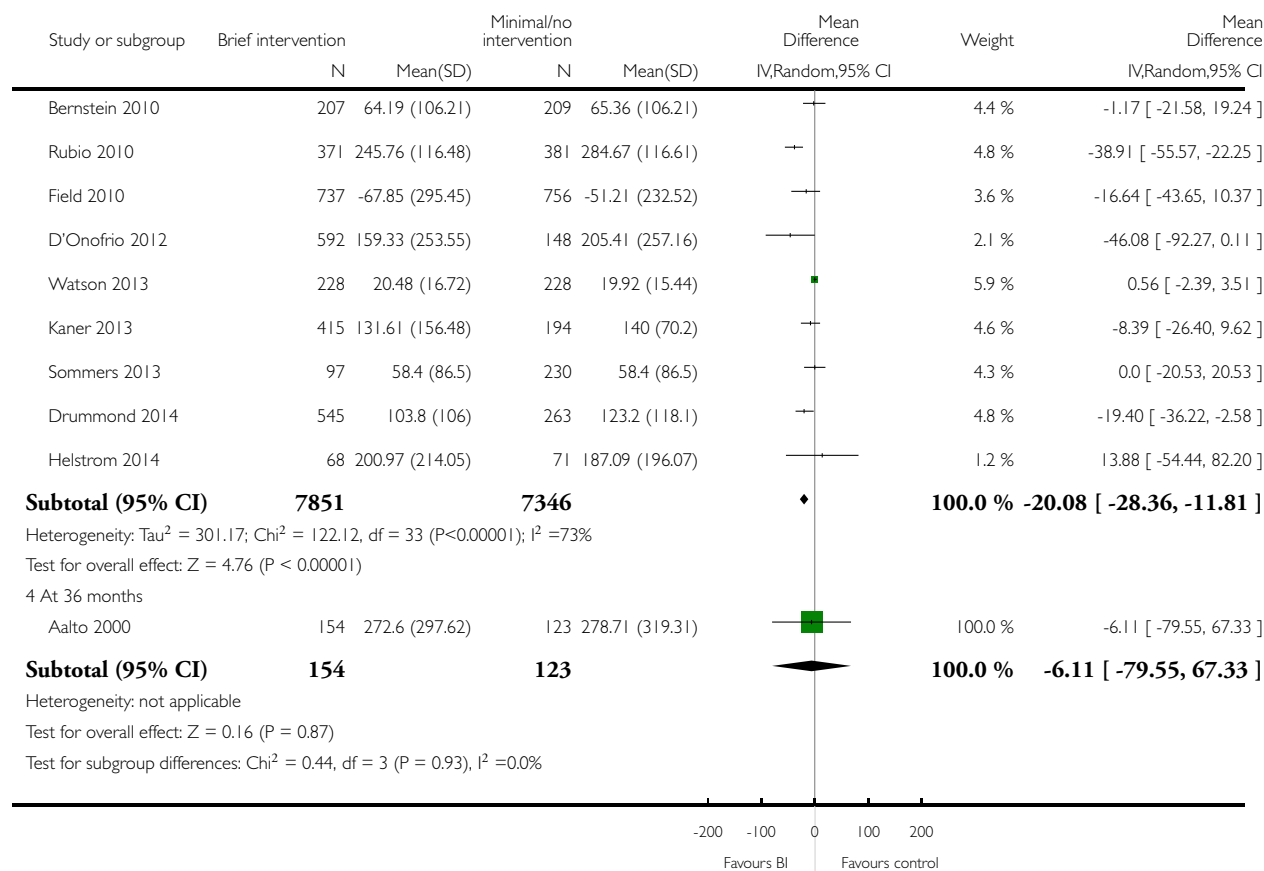
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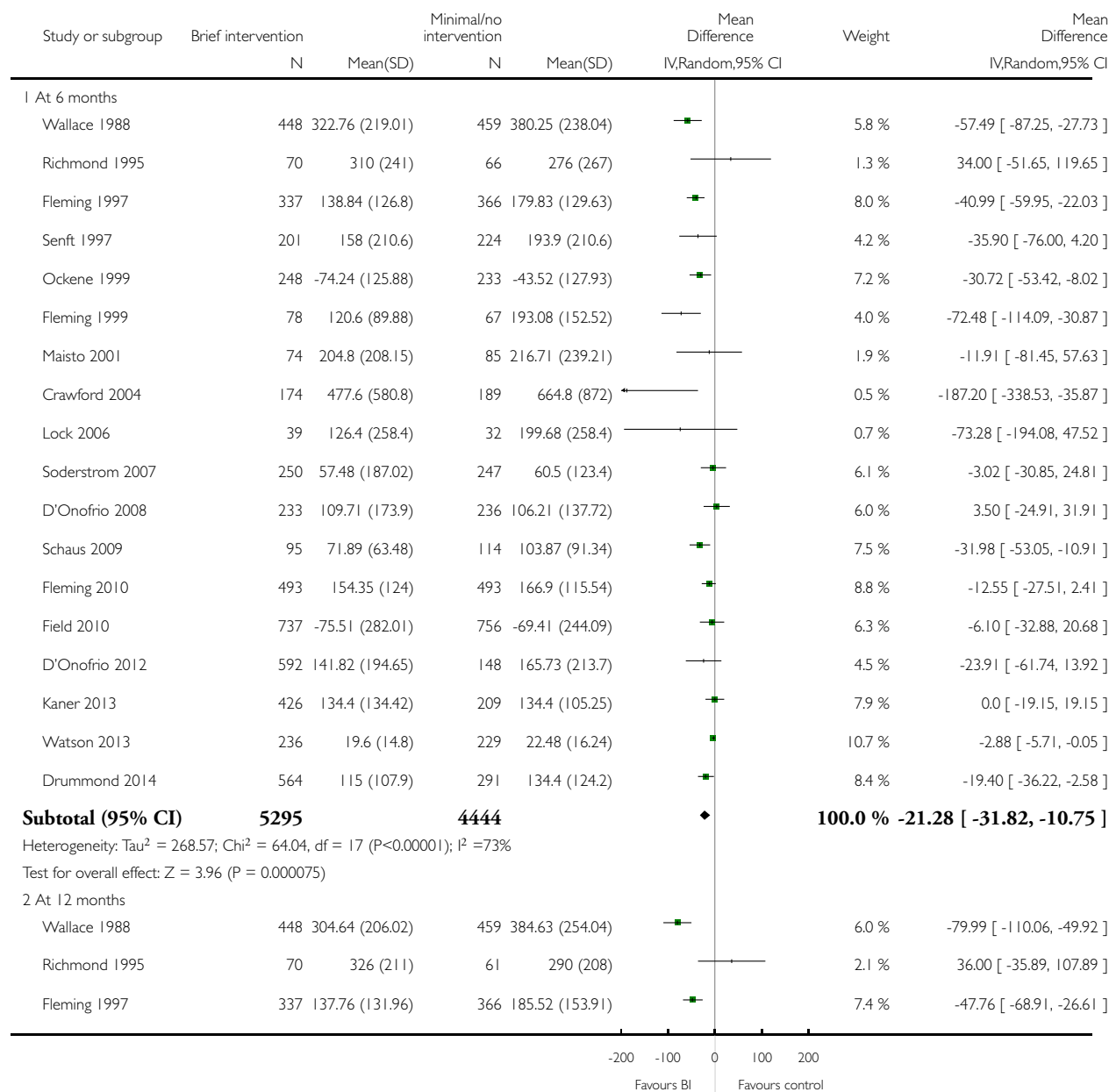


Analysis 1.14. Comparison 1 Brief intervention versus control, Outcome 14 Quantity of drinking (g/week), subgrouped by length of follow-up (6 and 12 months) and restricted to trials with information at both times.

Review: Effectiveness of brief alcohol interventions in primary care populations

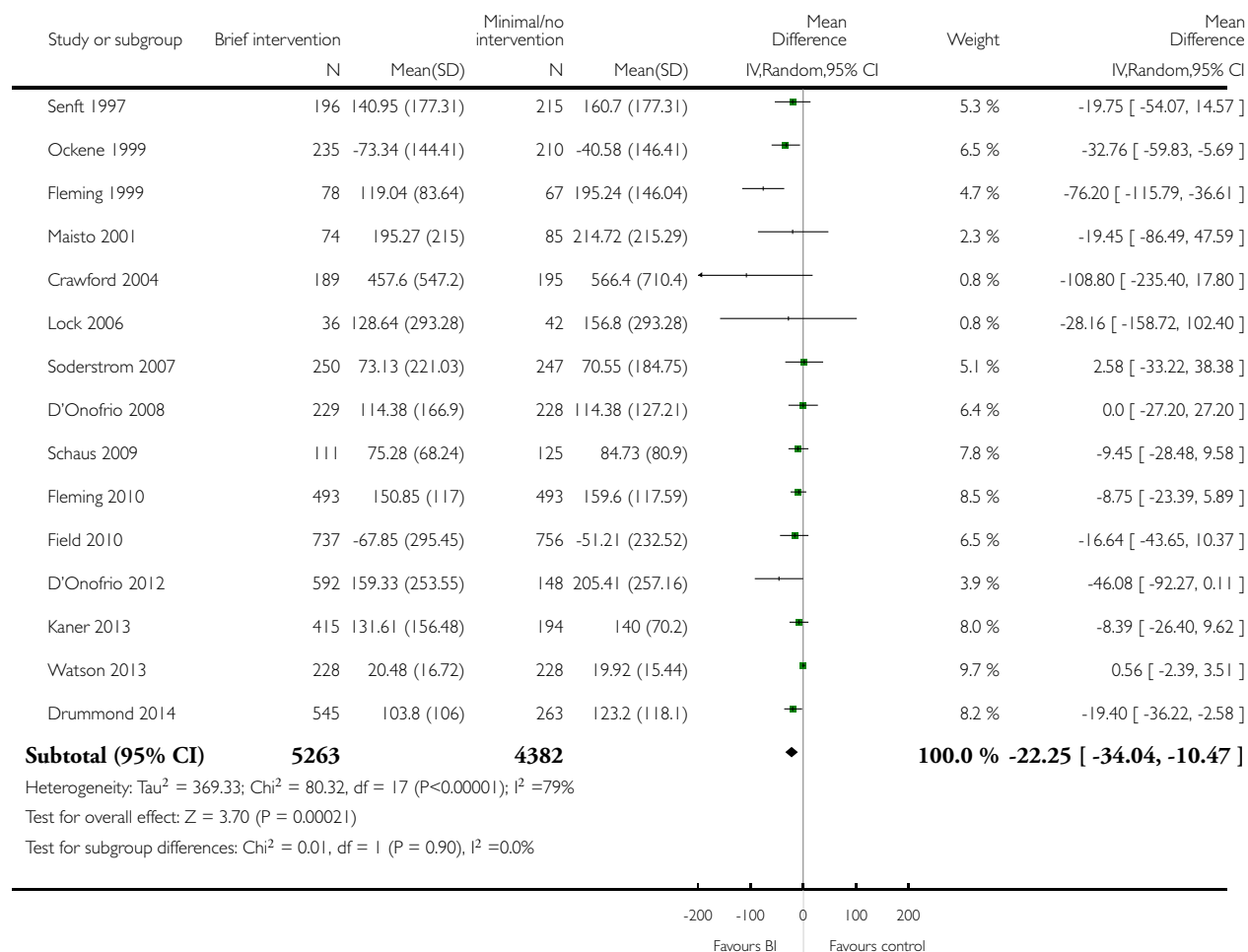
Comparison: 1 Brief intervention versus control

Outcome: 14 Quantity of drinking (g/week), subgrouped by length of follow-up (6 and 12 months) and restricted to trials with information at both times



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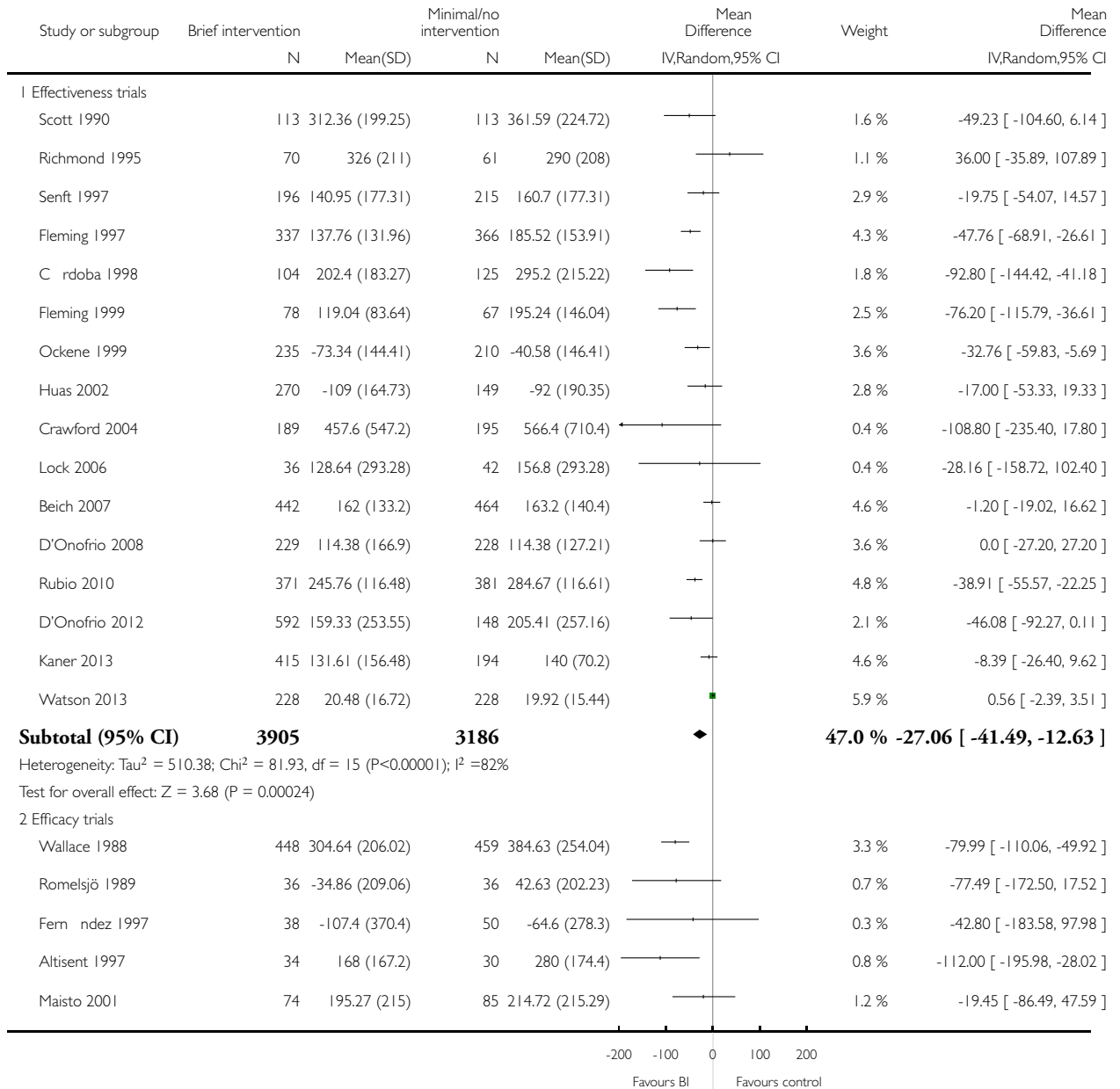


Analysis I.15. Comparison I Brief intervention versus control, Outcome 15 Quantity of drinking (g/week) at 12 months, subgrouped by effectiveness/efficacy.

Review: Effectiveness of brief alcohol interventions in primary care populations

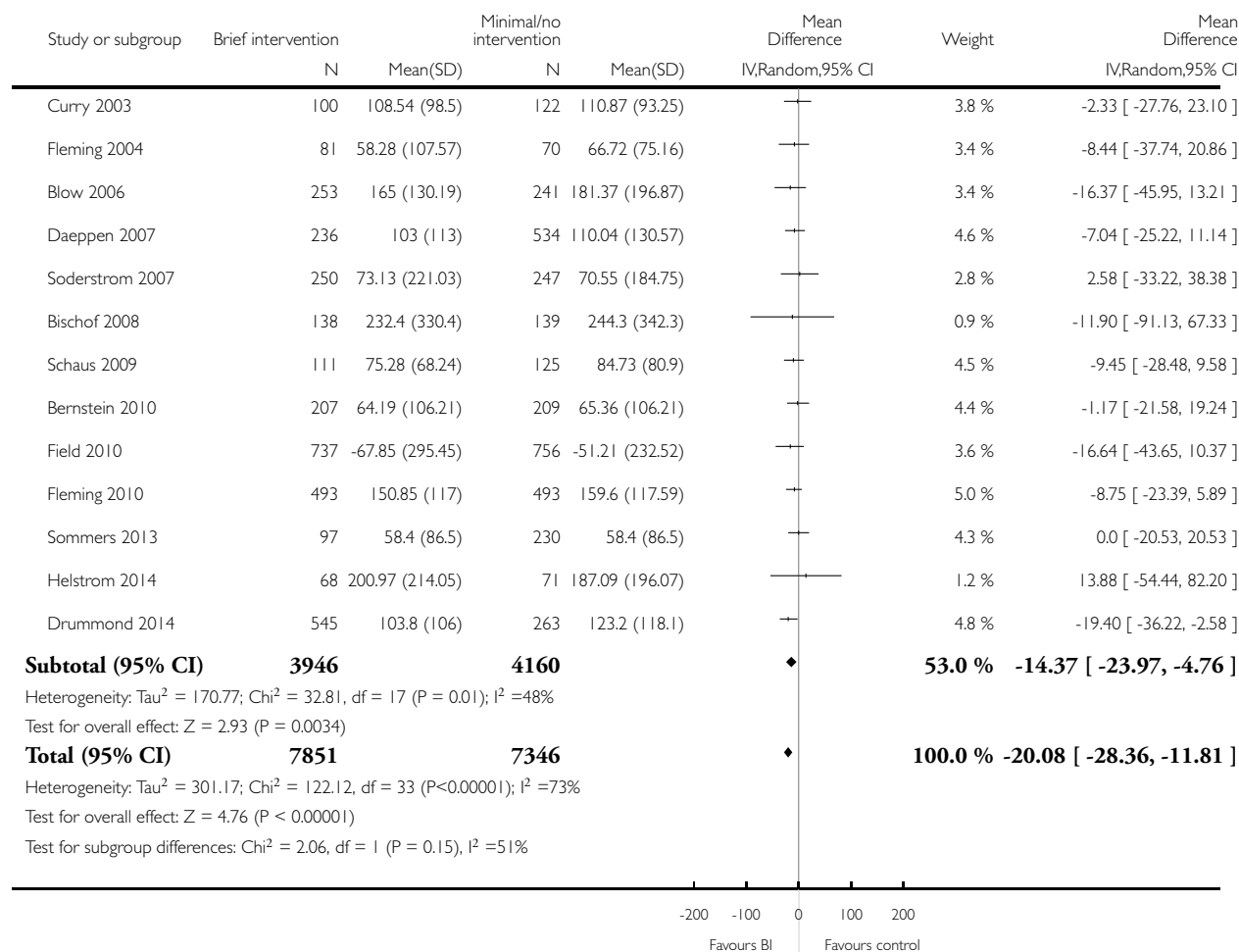
Comparison: I Brief intervention versus control

Outcome: 15 Quantity of drinking (g/week) at 12 months, subgrouped by effectiveness/efficacy



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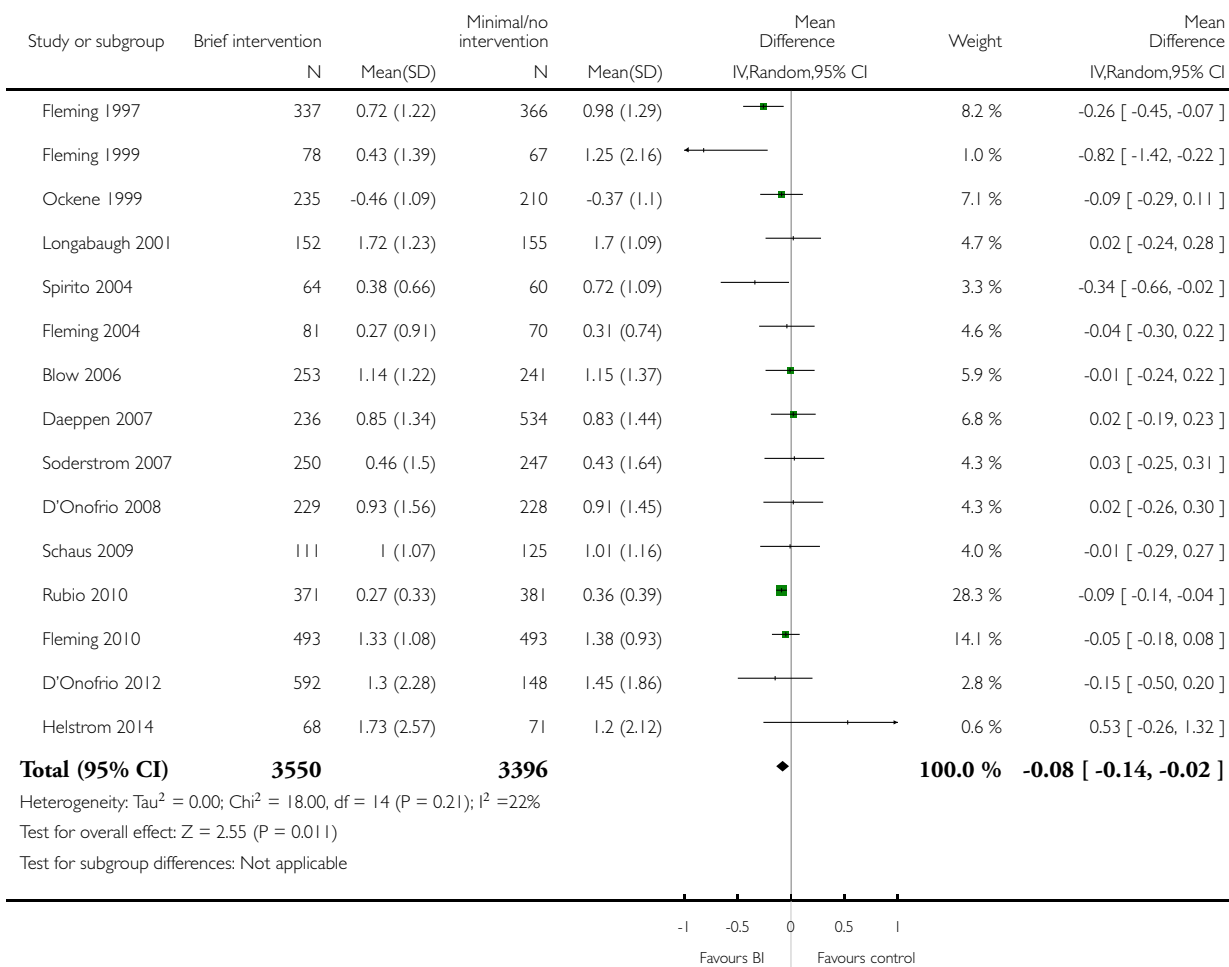


Analysis 1.16. Comparison 1 Brief intervention versus control, Outcome 16 Frequency of binge drinking (number binges/week) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 16 Frequency of binge drinking (number binges/week) at 12 months

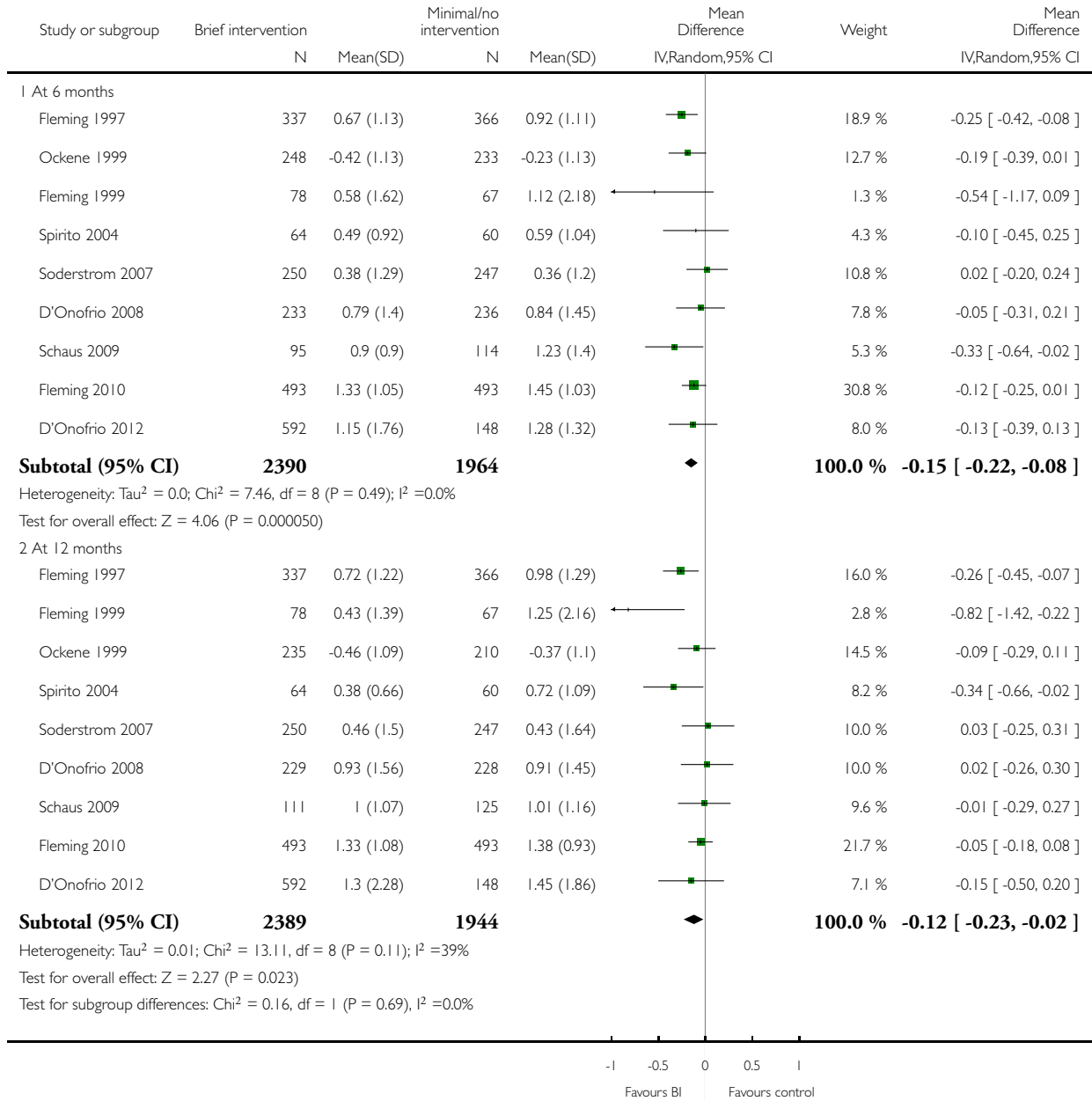


Analysis 1.17. Comparison 1 Brief intervention versus control, Outcome 17 Frequency of drinking (number binges/week), subgrouped by length of follow-up (6 & 12 months) and restricted to trials with information at both times.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 17 Frequency of drinking (number binges/week), subgrouped by length of follow-up (6 & 12 months) and restricted to trials with information at both times

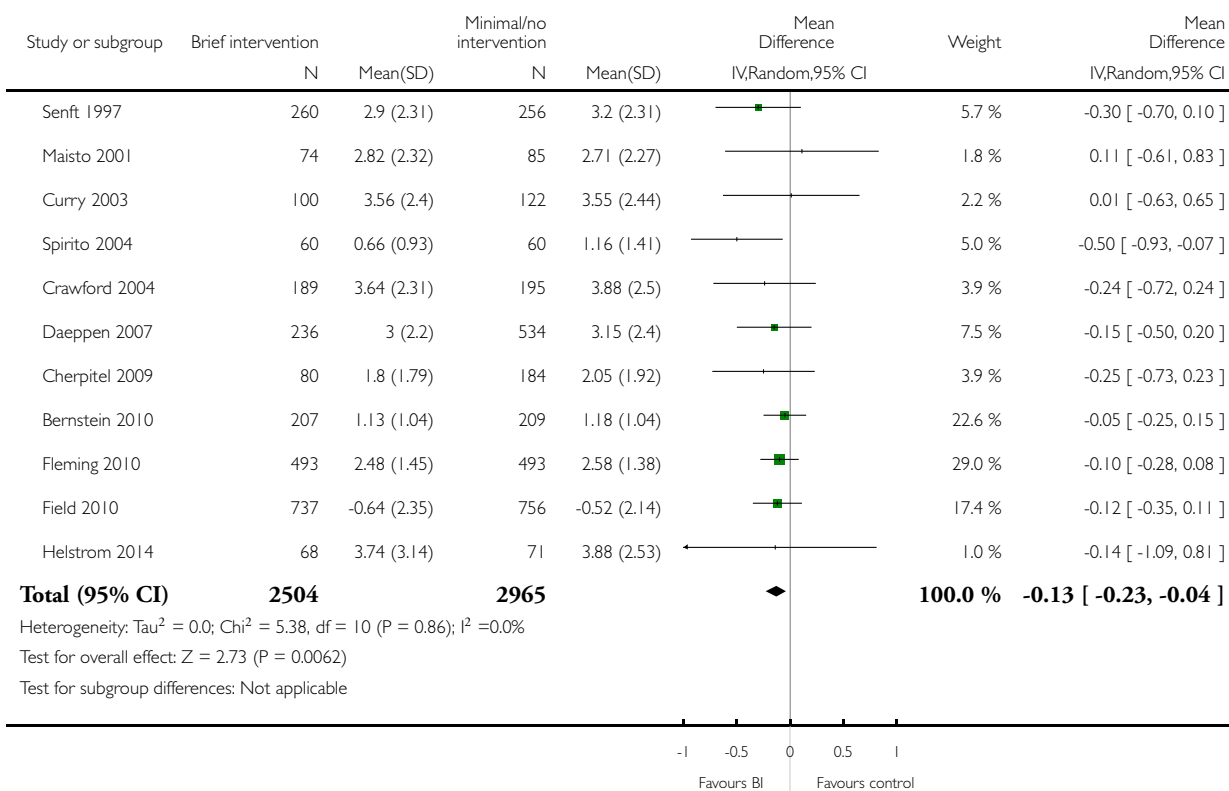


Analysis 1.18. Comparison 1 Brief intervention versus control, Outcome 18 Frequency of drinking (number days drinking/week) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 18 Frequency of drinking (number days drinking/week) at 12 months

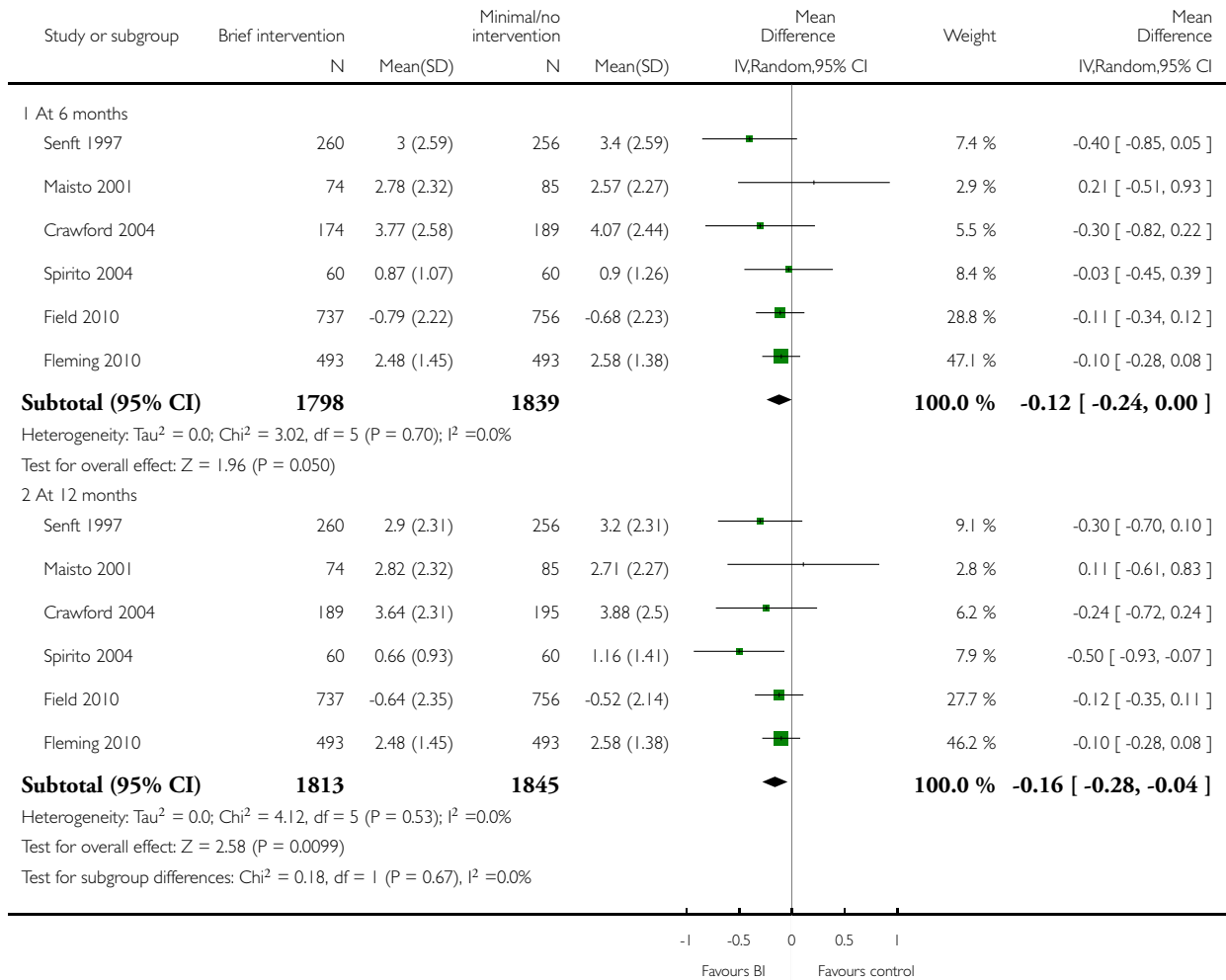


Analysis 1.19. Comparison 1 Brief intervention versus control, Outcome 19 Frequency of drinking (number days drinking/week), subgrouped by length of follow-up (6 & 12 months) and restricted to trials with information at both times.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 19 Frequency of drinking (number days drinking/week), subgrouped by length of follow-up (6 & 12 months) and restricted to trials with information at both times

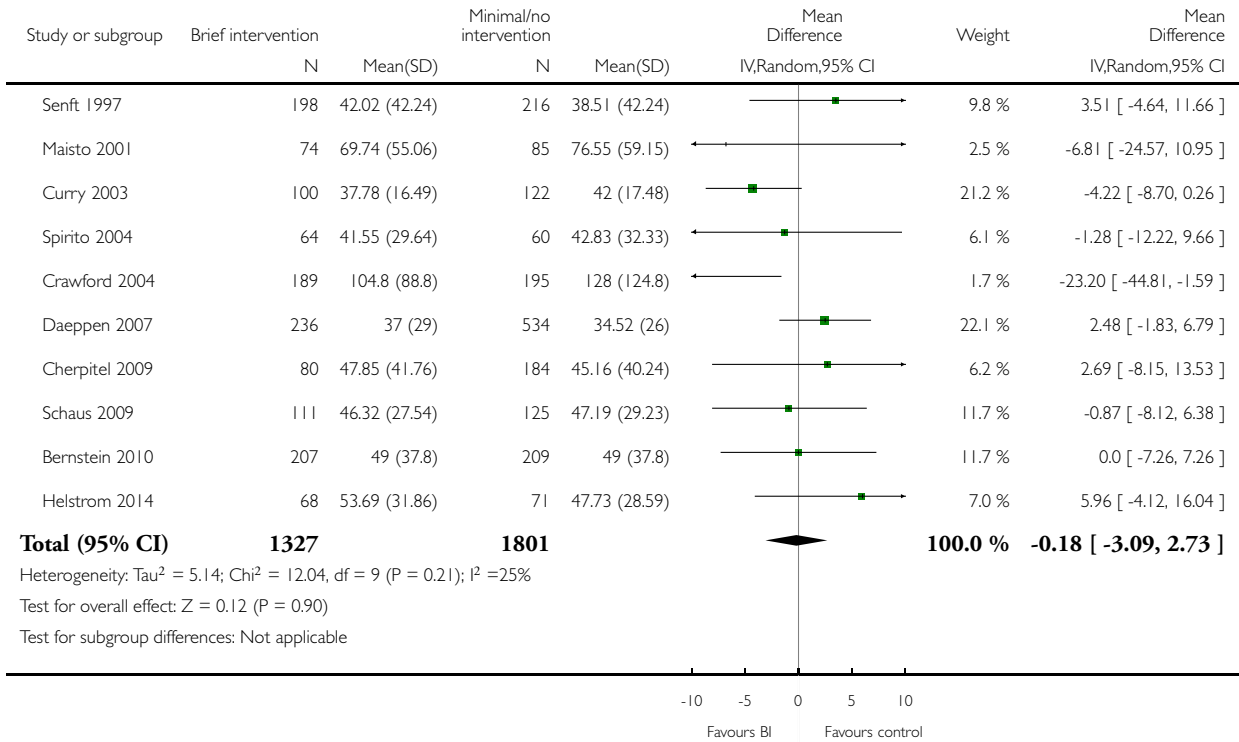


Analysis 1.20. Comparison 1 Brief intervention versus control, Outcome 20 Intensity of drinking (g/drinking day) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 20 Intensity of drinking (g/drinking day) at 12 months

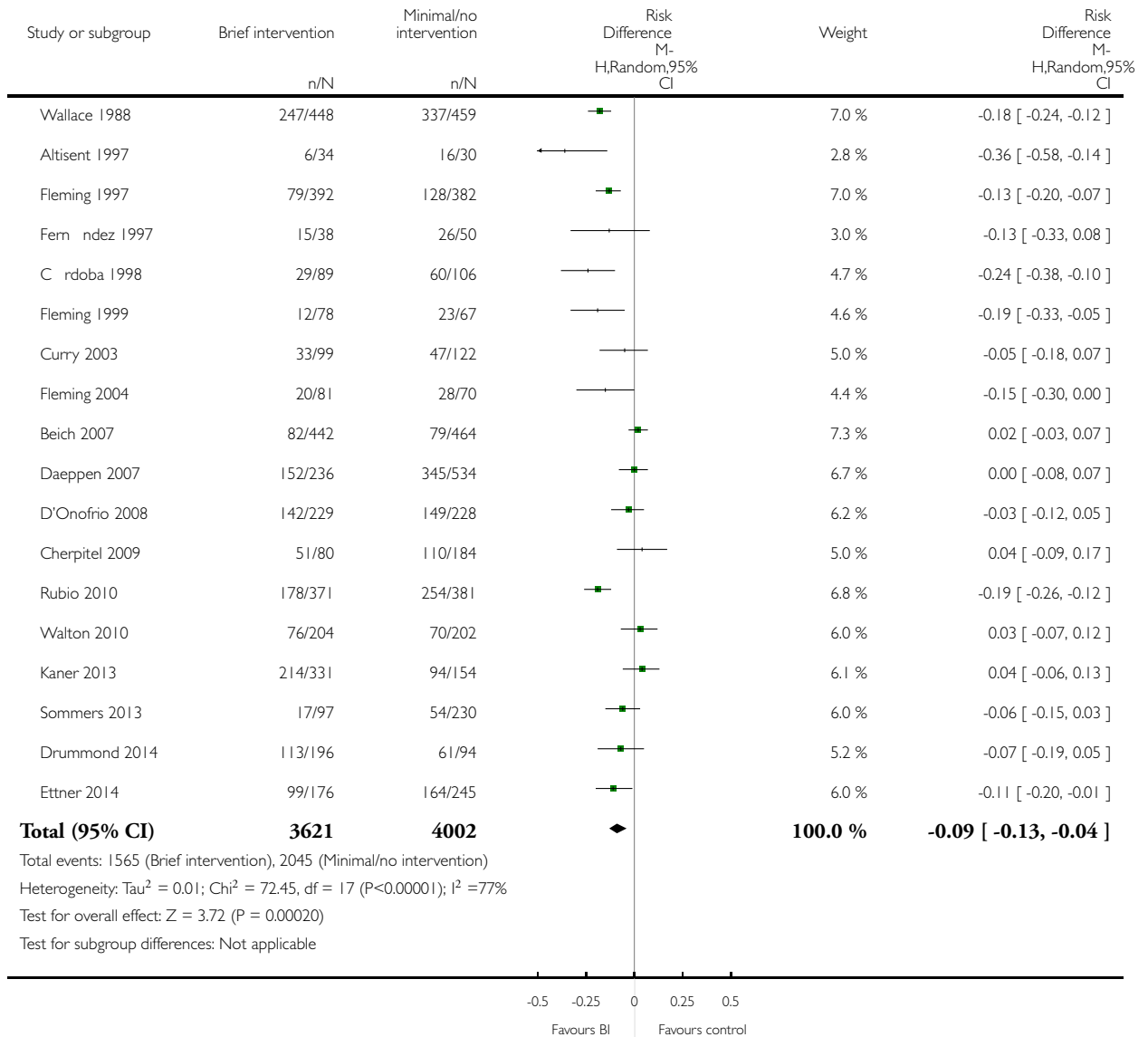


Analysis 1.21. Comparison 1 Brief intervention versus control, Outcome 21 Heavy drinkers at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 21 Heavy drinkers at 12 months

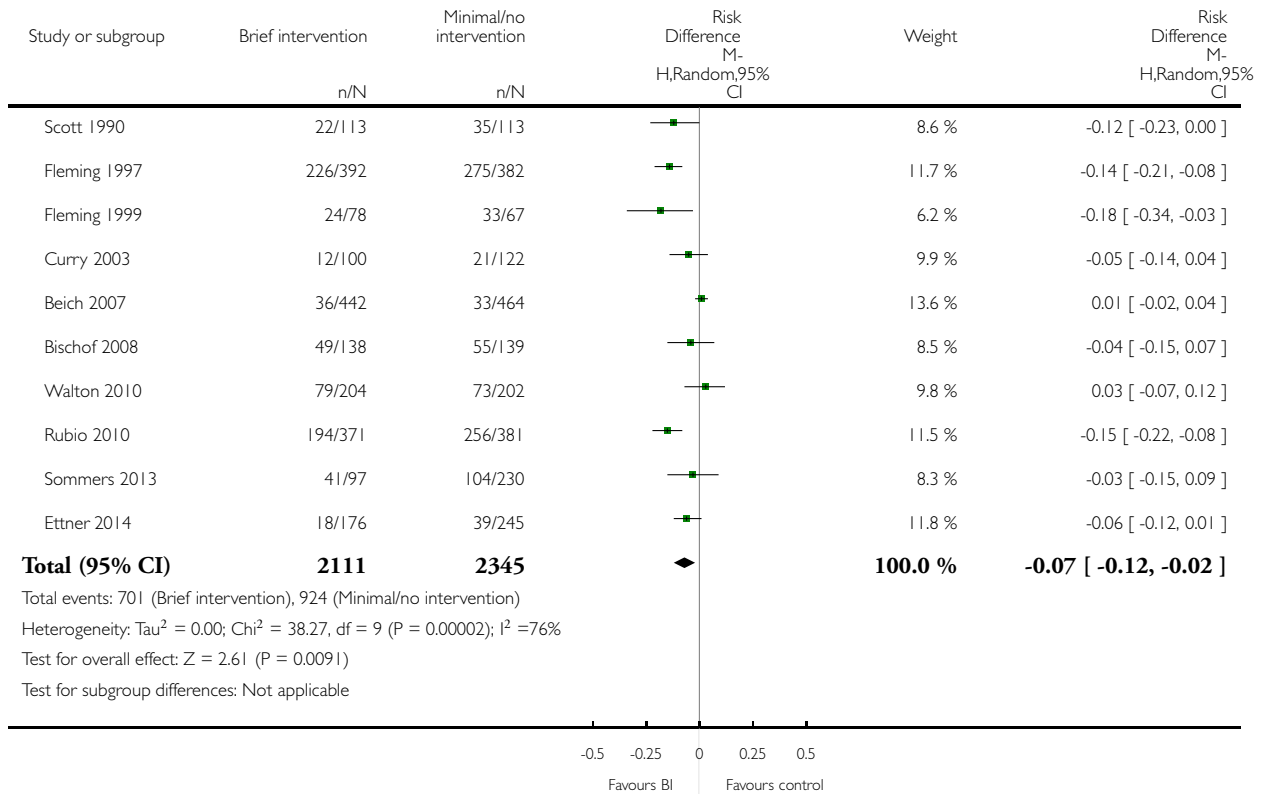


Analysis 1.22. Comparison 1 Brief intervention versus control, Outcome 22 Binge drinkers at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 22 Binge drinkers at 12 months

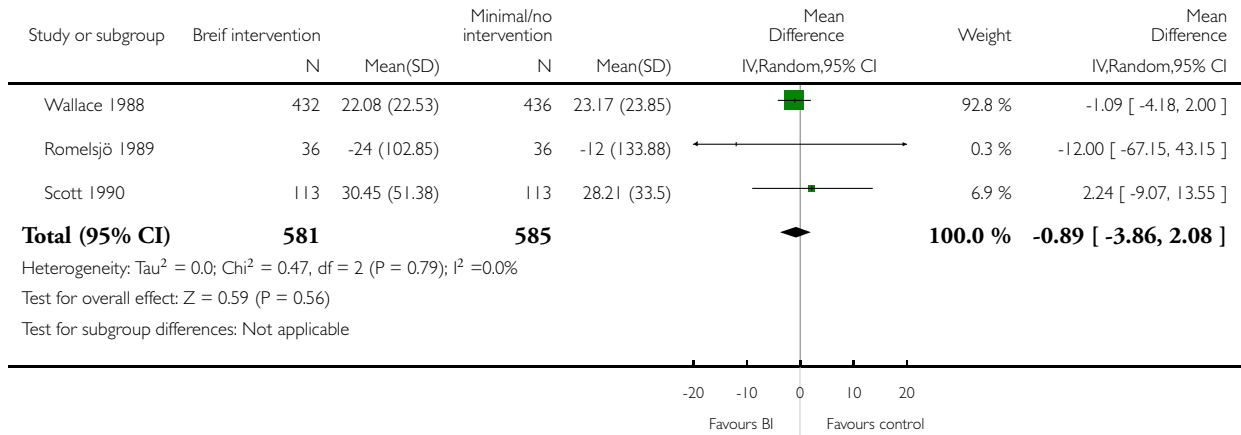


Analysis 1.23. Comparison 1 Brief intervention versus control, Outcome 23 Laboratory markers - GGT (IU/L) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 23 Laboratory markers - GGT (IU/L) at 12 months

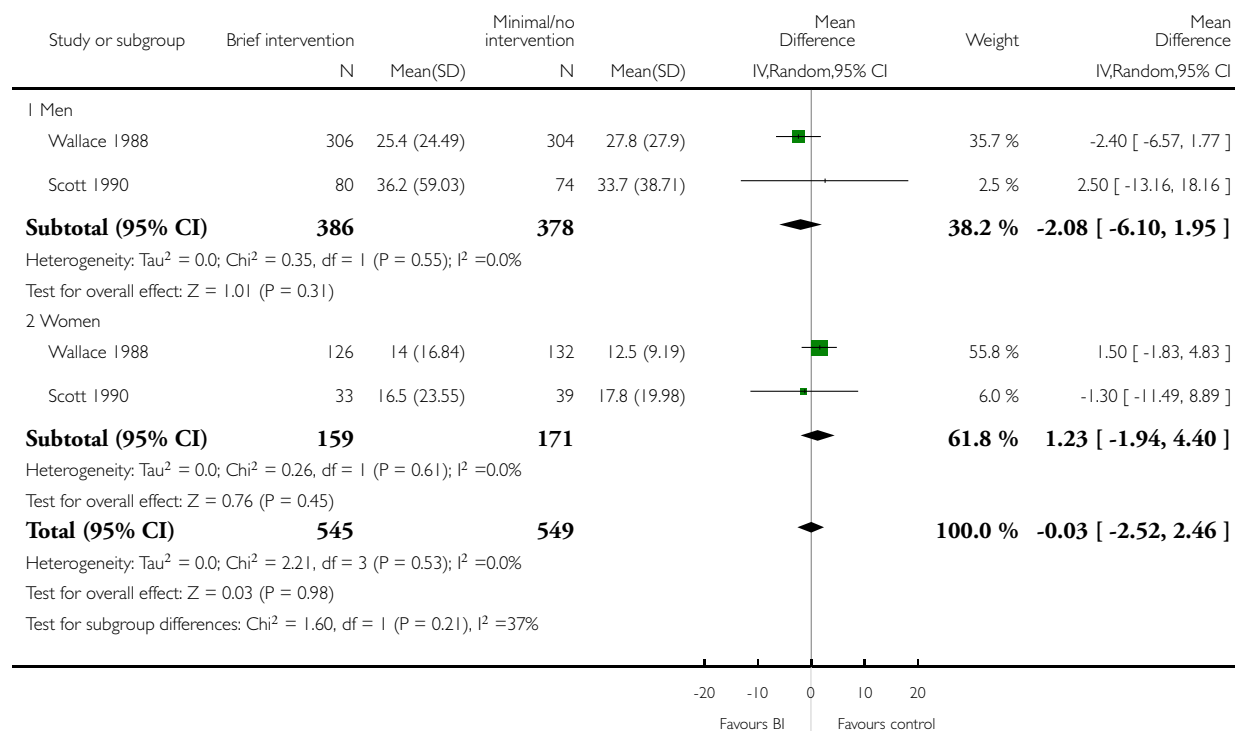


Analysis 1.24. Comparison 1 Brief intervention versus control, Outcome 24 Laboratory markers - GGT (IU/L) at 12 months, subgrouped by gender.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 1 Brief intervention versus control

Outcome: 24 Laboratory markers - GGT (IU/L) at 12 months, subgrouped by gender

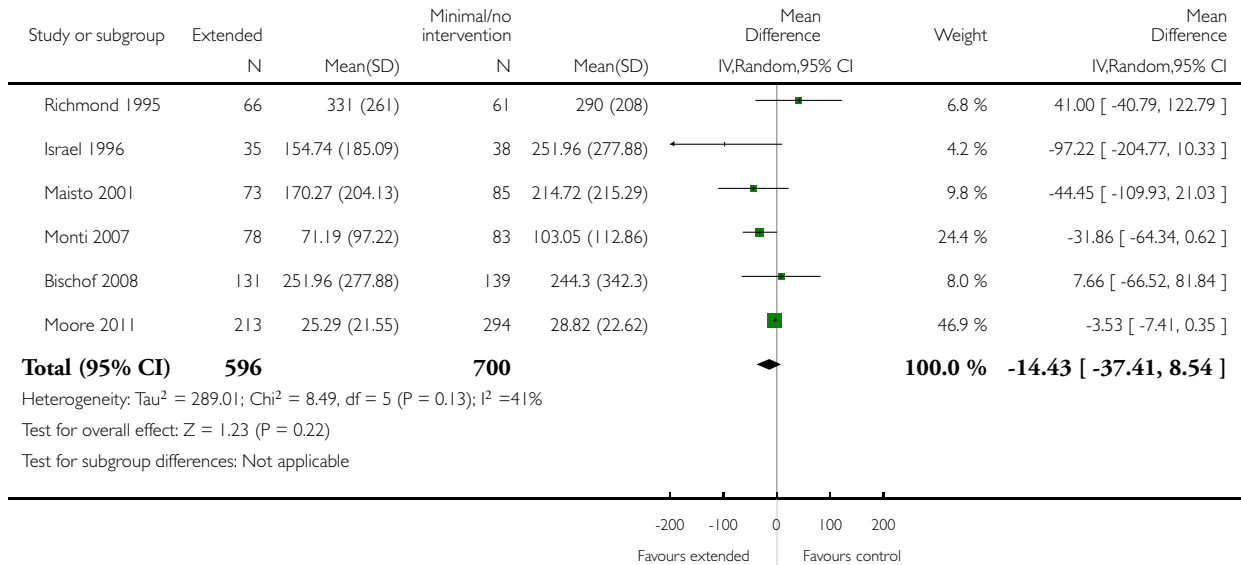


Analysis 2.1. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 1 Quantity of drinking (g/week) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 1 Quantity of drinking (g/week) at 12 months

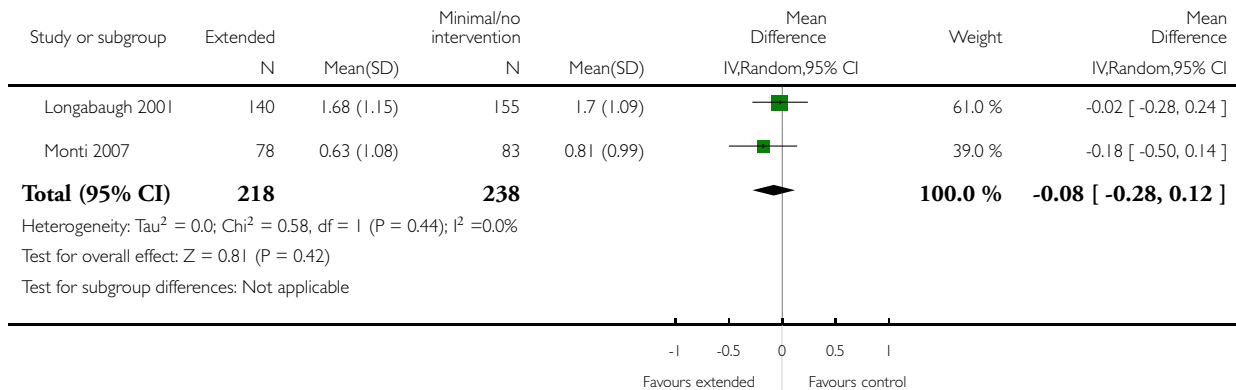


**Analysis 2.2. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 2
Frequency of binge drinking (number binges/week) at 12 months.**

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 2 Frequency of binge drinking (number binges/week) at 12 months

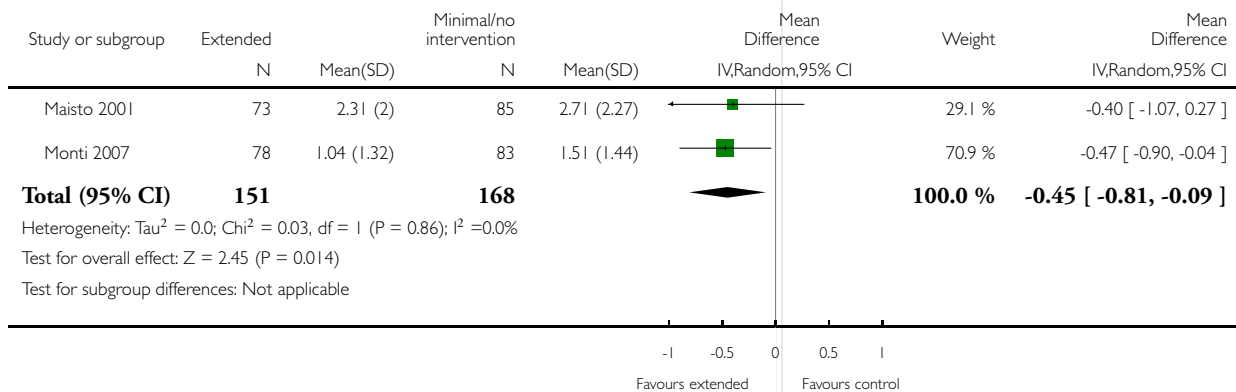


**Analysis 2.3. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 3
Frequency of drinking (number days drinking/week) at 12 months.**

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 3 Frequency of drinking (number days drinking/week) at 12 months

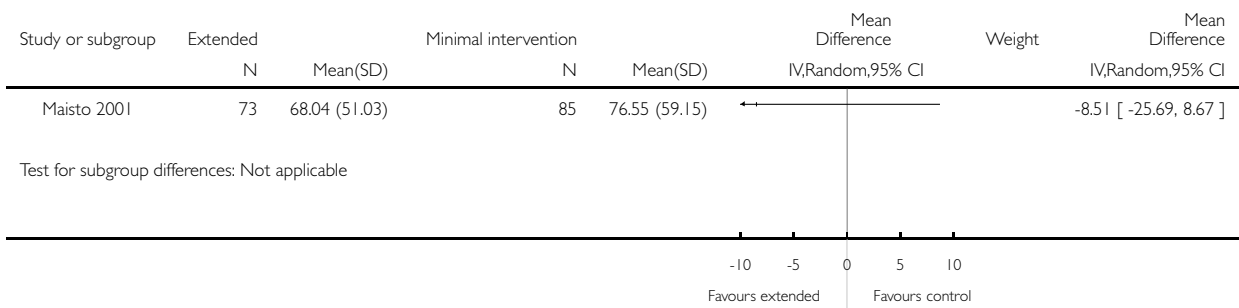


Analysis 2.4. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 4 Intensity of drinking (g/drinking day) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 4 Intensity of drinking (g/drinking day) at 12 months

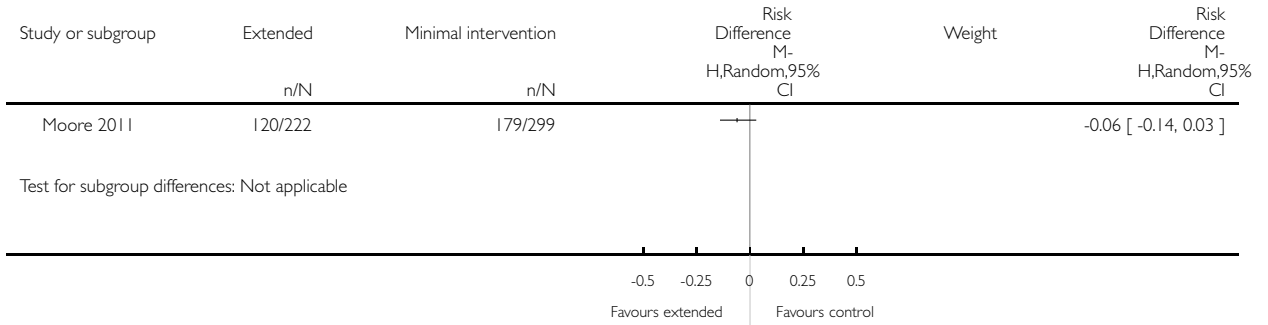


Analysis 2.5. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 5 Heavy drinkers at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 5 Heavy drinkers at 12 months

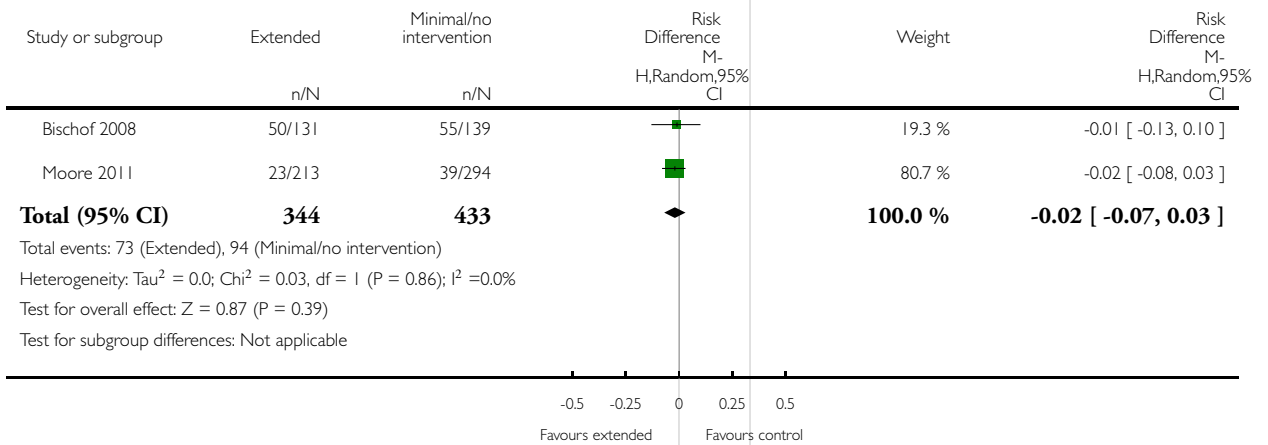


Analysis 2.6. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 6 Binge drinkers at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 6 Binge drinkers at 12 months

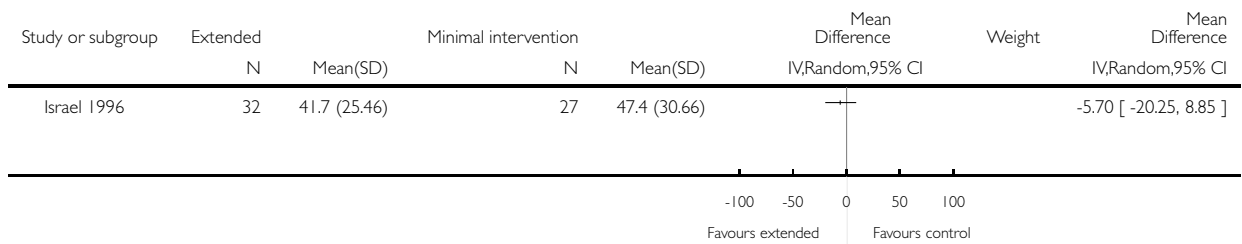


Analysis 2.7. Comparison 2 Extended intervention versus minimal or no intervention, Outcome 7 Laboratory markers - GGT (IU/L) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 2 Extended intervention versus minimal or no intervention

Outcome: 7 Laboratory markers - GGT (IU/L) at 12 months

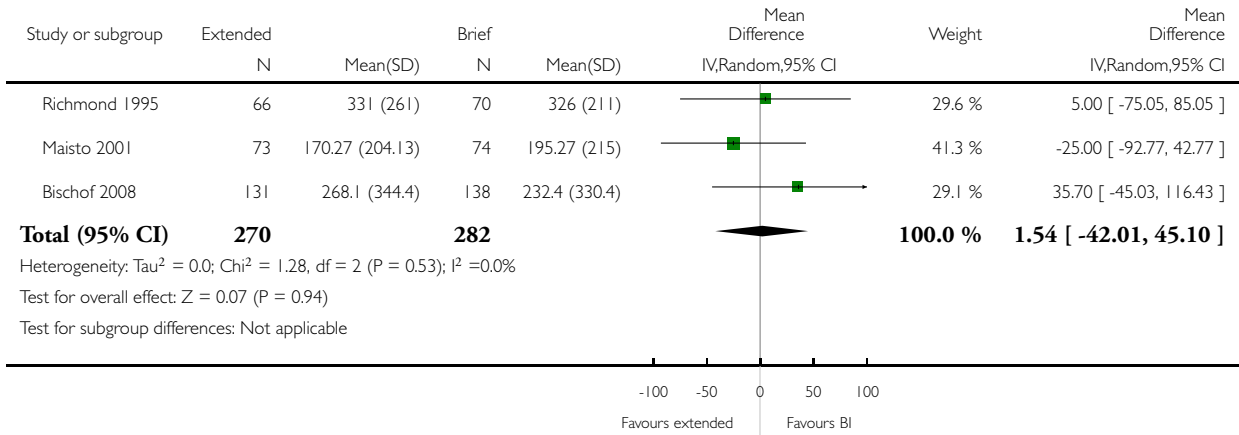


Analysis 3.1. Comparison 3 Extended versus brief intervention, Outcome 1 Quantity of drinking (g/week) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 3 Extended versus brief intervention

Outcome: 1 Quantity of drinking (g/week) at 12 months

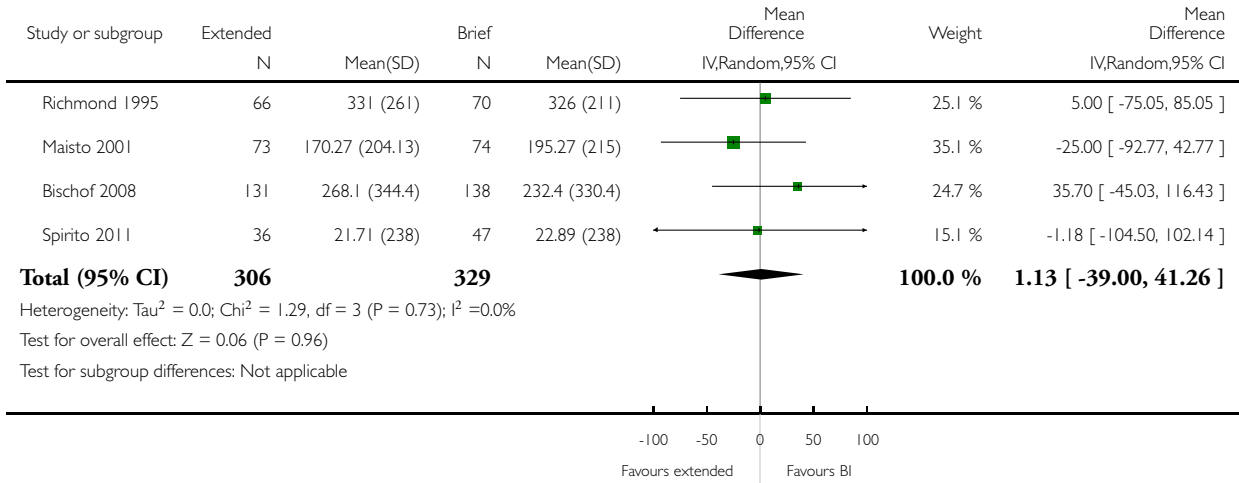


Analysis 3.2. Comparison 3 Extended versus brief intervention, Outcome 2 Quantity of drinking (g/week) at 12 months, with imputation of unknown standard deviations.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 3 Extended versus brief intervention

Outcome: 2 Quantity of drinking (g/week) at 12 months, with imputation of unknown standard deviations

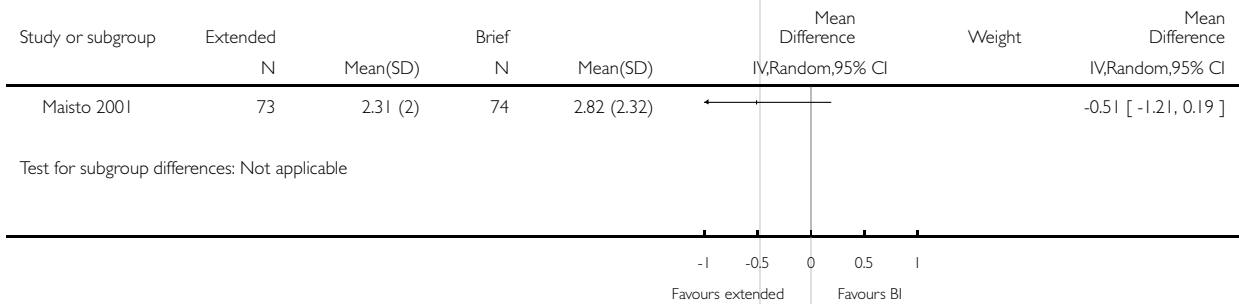


Analysis 3.3. Comparison 3 Extended versus brief intervention, Outcome 3 Frequency of drinking (number days drinking/week) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 3 Extended versus brief intervention

Outcome: 3 Frequency of drinking (number days drinking/week) at 12 months

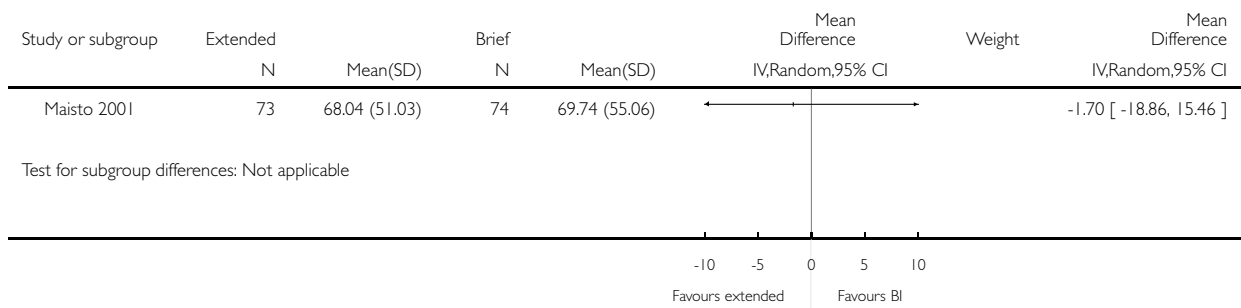


Analysis 3.4. Comparison 3 Extended versus brief intervention, Outcome 4 Intensity of drinking (g/drinking day).

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 3 Extended versus brief intervention

Outcome: 4 Intensity of drinking (g/drinking day)

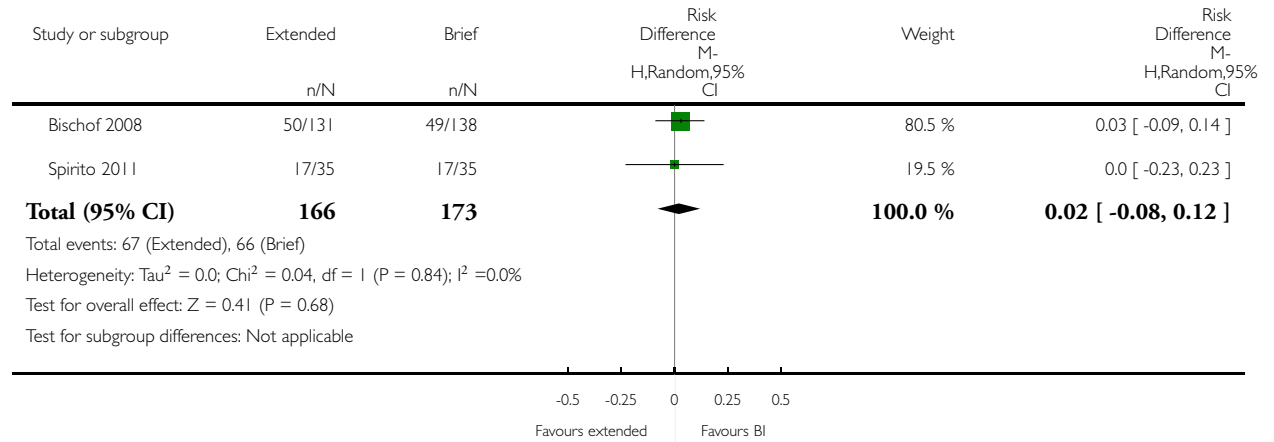


Analysis 3.5. Comparison 3 Extended versus brief intervention, Outcome 5 Binge drinkers at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 3 Extended versus brief intervention

Outcome: 5 Binge drinkers at 12 months

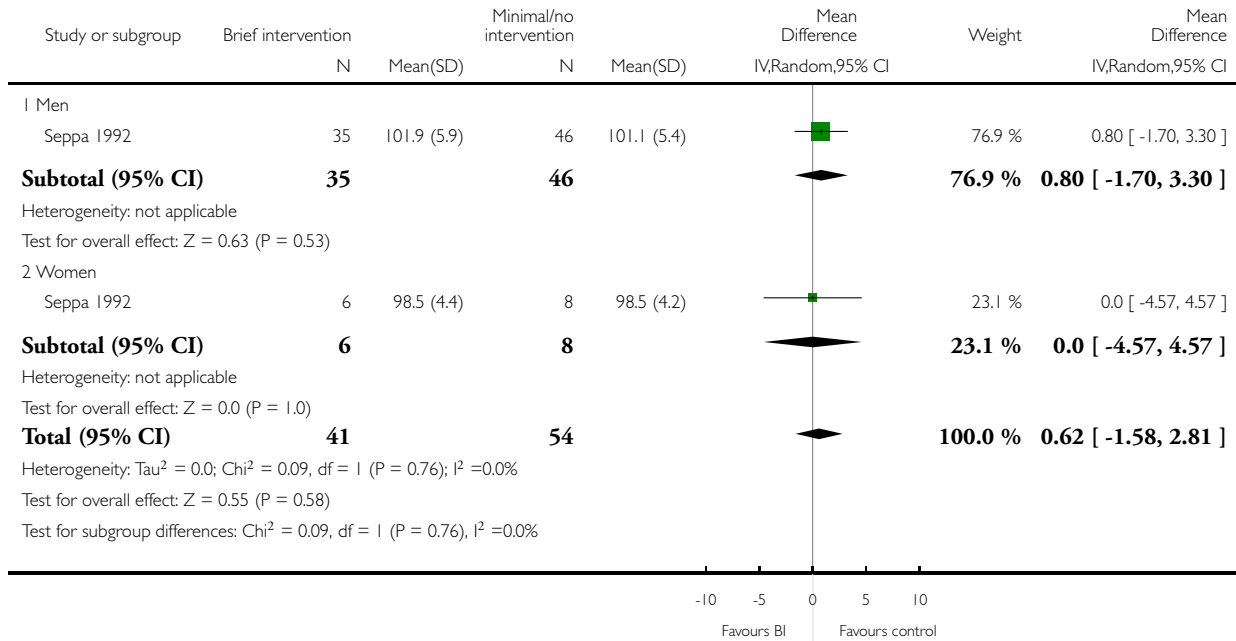


Analysis 3.6. Comparison 3 Extended versus brief intervention, Outcome 6 Laboratory markers - MCV (fl) at 12 months.

Review: Effectiveness of brief alcohol interventions in primary care populations

Comparison: 3 Extended versus brief intervention

Outcome: 6 Laboratory markers - MCV (fl) at 12 months



ADDITIONAL TABLES

Table 1. Effectiveness - efficacy scale

Scale item	Score	Meaning
Patients and problems	2	Clinically representative people initially present with a typically wide range of problems via self-referral or invitation for a health check
	1	Mixed: e.g. routine patients but paid for participation in study, or patients prescreened then invited research representative subjects may be paid patients
	0	Researcher-solicited volunteers (e.g. via advertisement) or referrals from specialist services
Practice context	2	Clinically representative is a community-based setting in which a range of clinical services are usually provided to patients

Table 1. Effectiveness - efficacy scale (Continued)

	1	Mixed
	0	Research representative is a setting in which the research function clearly dominates any clinical one (e.g. clinic at a university or hospital)
Practitioners and therapists	2	Clinically representative practitioners are practising doctors, nurses and qualified therapists who earn their main living by providing health services in primary care
	1	Qualified clinician but specifically recruited for the study
	0	Research representative practitioners are non-clinicians, or clinicians in training, who are contracted to deliver interventions for the purposes of the study
Intervention content	2	Clinically representative intervention fits with current practice in terms of timing, content or style (e.g. for primary care 5 to 15 minutes for a GP; 20 to 30 minutes for a nurse or initial screening accompanied by a return visit for brief intervention; for emergency settings motivational interviewing-style intervention fits in here, e.g. 45 minutes)
	0	Research representative treatment would not normally occur in routine practice e.g. unusually long consultations
Therapeutic flexibility	1	Clinically representative: allows professional judgement in how an intervention is delivered e.g. freedom to focus on particular issues according to patient need
	½	Flexible protocol (tailored to participants)
	0	Research representative: strict adherence to a prescribed protocol or script that does not allow for variability in practice
Pre-therapy training	1	Clinically representative training in intervention procedures occurs according to typical CPD/CME procedures (e.g. outreach visits, seminars, one-off training days) Full day off-site (for emergency care staff)
	½	Full day off-site (for primary care staff)
	0	Research representative training is unusually intensive or requiring of atypical levels of interest or motivation, e.g. prolonged or intensive courses, formal qualification
Intervention support	1	Clinically representative support occurs within standard practice resources (e.g. colleague assistance with screening, IT flagging, provision of general (uncustomised) manual. Note for emergency care settings some procedures are clinically representative (e.g. taking bloods and doing various tests)
	0	Research representative support would not typically be available (e.g. researcher help to flag notes, extra staff for period of the trial)
Intervention monitoring	1	Clinically representative monitoring of intervention delivery does not interfere with practitioners' behaviour or their relationships with patients

Table 1. Effectiveness - efficacy scale (Continued)

	0	Research representative monitoring would be direct observation of therapist behaviour or ongoing/immediate feedback to practitioners after each session
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Abbreviations: CPD: continuing professional development; CME: continuing medical education; GP: general practitioner; IT: information technology

Table 2. Conversion factors for alcohol consumption and efficacy scores of trials

Trial	Reported units	Conversion factor	Source of conversion	Efficacy score	Treatment exposure ¹
Aalto 2000	g/week	1	NA	10.5	45, 105
Altisent 1997	Units/week	8	Altisent 1997	8.5	20
Babor 2006	Drinks/week	14	Babor 2006	11	4
Bazargan-Hejazi 2005	Alcohol consumption	NR	NA	9	17.5
Beckham 2007	Drinks/day	11.671 x 7	Miller 1991	4.5	52.5
Beich 2007	Drinks/week	12	Beich 2007	11	10
Bernstein 2010	Drinks/week	11.671	Miller 1991	6.5	32.5
Bischof 2008	g/day	7	NA	6.5	60
Blow 2006	Drinks/week	13	Blow 2009*	7.5	7.5
Cherpitel 2009	Drinks/week	11.671	Miller 1991	9	17.5
Córdoba 1998	Units/week	8	Córdoba 1998	11	15
Crawford 2004	Units/week	8	Miller 1991	10.5	30
Crawford 2010	Alcohol consumption	NR	NA	8.5	30
Curry 2003	Drinks/week	11.671	Miller 1991	8.5	22.5
D'Amico 2008	Basic statistics	NR	NA	7.5	25
D'Onofrio 2008	Drinks/week	11.671	Miller 1991	10.5	6.7
D'Onofrio 2012	Drinks/week	11.671	Miller 1991	10.5	7;17

Table 2. Conversion factors for alcohol consumption and efficacy scores of trials (Continued)

Daepfen 2007	Drinks/week	10	Daepfen 2007	7	15
Dent 2008	Alcohol consumption	NR	NA	11.5	5; 45
Désy 2010	Drinks/week	11.671	Miller 1991	10.5	7.5
Díez 2002	Units/week	8	Díez 2002	10.5	10
Drummond 2009	Drinks previous 180 days	8 x (7/180)	Drummond 2009	10.5	40; 200
Drummond 2014	Drinks/day	8 x 7	Drummond 2009	8	26
Ettner 2014	Drinks/week	11.671	Miller 1991	11	108
Fernández 1997	Units/week	10	Miller 1991	7.5	10
Field 2010	Drinks/week	11.671	Miller 1991	7.5	17.5
Fleming 1997	Drinks/week	12	Fleming 1997	10.5	40
Fleming 1999	Drinks/week	12	Fleming 1999	9	40
Fleming 2004	Drinks/month	11.671 x (12/52)	Miller 1991	4.5	40
Fleming 2010	Drinks in previous 28 days	11.671 x (7/28)	Miller 1991	8.5	42.5
Heather 1987	Units/month	8 x (12/52)	Heather 1987	8.5	7.5
Helstrom 2014	Drinks/day	11.671 x 7	Miller 1991	7	37.5
Huas 2002	Units/week	10	Heather 2006	10	7.5
Israel 1996	Drinks/month	13.456 x (12/52)	Miller 1991	7.5	150
Kaner 2013	Drinks/day	8 x 7	Kaner 2013	11	26
Kunz 2004	Drinks/week	11.671	Miller 1991	6	37.5
L'Engle 2014	Alcohol consumption	NR	NA	8	180
Lane 2008	Alcohol consumption	NR	NA	10.5	7.5
Lock 2006	Drinks/week	8	Miller 1991	12	7.5

Table 2. Conversion factors for alcohol consumption and efficacy scores of trials (Continued)

Longabaugh 2001	Alcohol consumption	NR	NA	6	50
Maisto 2001	Drinks/month	11.671 x (12/52)	Miller 1991	5	13.5; 65
McDevitt-Murphy 2014	Drinks/week	11.671	Miller 1991	5.5	60
McIntosh 1997	Drinks/month	13.456 x (12/52)	Miller 1991	10.5	60
Mello 2008	Alcohol consumption	NR	NA	7	45
Mertens 2014	Alcohol consumption	NR	NA	7.5	10
Monti 2007	Drinks/week	11.671	Miller 1991	7.5	85
Moore 2011	Standard drinks/month	11.671 x (12/52)	Miller 1991	9.5	83
Noknoy 2010	Drinks/week	10	Furtwaengler 2013	11	45
Ockene 1999	Drinks/week	12.8	Ockene 1999	10	7.5
Richmond 1995	Drinks/week	10	Richmond 1995	9.5	5, 57.5
Rodríguez 2003	Alcohol consumption	NR	NA	10.5	17.5
Romelsjö 1989	g/day	1 x 7	NA	4.5	7.5
Rubio 2010	Drinks/week	12.8	Rubio 2010	9.5	32.5
Schaus 2009	Drinks/week	11.671	Miller 1991	8	40
Scott 1990	Units/week	8	Miller 1991	11	10
Segatto 2011	Alcohol consumption	NR	NA	6	45
Senft 1997	Drinks/3 months	11.671 x (4/52)	Miller 1991	9	16
Seppa 1992	Alcohol consumption	NR	NA	8.5	15
Soderstrom 2007	Drinks/last 90 days	11.671 x (7/90)	Miller 1991	6.5	32.5

Table 2. Conversion factors for alcohol consumption and efficacy scores of trials (Continued)

Sommers 2006	Alcohol consumption	NR	NA	7	40
Sommers 2013	Drinks/week	11.671	Miller 1991	7.5	40
Spirito 2004	Standard drinks/month	11.671 x (12/52)	Miller 1991	7	40
Spirito 2011	Alcohol consumption	NR	NA	7.5	52.5
Tait 2004	alcohol consumption	NR	NA	8	37.5
Tomson 1998	g/week	1	NA	8.5	37.5
Wallace 1988	Units/week	8	Miller 1991	9.5	7.5
Walton 2010	Alcohol consumption	NR	NA	7	37
Watson 2013	Drinks/week	8	Watson 2013	9.5	20
Woolard 2013	Alcohol consumption	NR	NA	5.5	72.5

* Blow 2009 is a report of [Blow 2006](#).

¹ Treatment exposure was calculated in minutes. Where two values appear, these are the durations of different intervention arms. Abbreviations: NA: not applicable; NR: not reported.

APPENDICES

Appendix I. Cochrane Library (Wiley) search strategy

<http://www.thecochranelibrary.com/>

Searched 19 June 2012, 6 January 2015, 25 September 2017

Databases searched: CENTRAL, CDSR, DARE, HTA, NHS-EED

#1 MeSH descriptor: [General Practice] explode all trees

#2 MeSH descriptor: [Primary Health Care] this term only

#3 MeSH descriptor: [Community Health Services] this term only

#4 MeSH descriptor: [Emergency Service, Hospital] explode all trees

#5 MeSH descriptor: [Physicians, Family] this term only

#6 MeSH descriptor: [Physicians, Primary Care] this term only

#7 (general pract* or family pract* or GP or physician* or emergency department or emergency room or "accident and emergency" or community):ti,ab

#8 (primary near/2 care):ti,ab
 #9 (shared next care):ti,ab
 #10 #1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9
 #11 MeSH descriptor: [Alcohol-Related Disorders] explode all trees
 #12 MeSH descriptor: [Alcohol Drinking] this term only
 #13 (alcohol near/3 (drink* or intoxicat* or abus* or misus* or risk* or consum* or withdraw* or detox* or treat* or therap* or excess* or reduc* or cessation or intervention)):ti,ab
 #14 (drink* near/3 (excess or heavy or heavily or hazard* or binge or harmful or problem*)):ti,ab
 #15 (“alcohol use” or alcoholic*):ti,ab
 #16 #11 or #12 or #13 or #14 or #15
 #17 MeSH descriptor: [Counseling] this term only
 #18 MeSH descriptor: [Psychotherapy, Brief] this term only
 #19 MeSH descriptor: [Interview, Psychological] this term only
 #20 MeSH descriptor: [Behavior Therapy] this term only
 #21 MeSH descriptor: [Cognitive Therapy] this term only
 #22 MeSH descriptor: [Motivation] this term only
 #23 (brief near/3 (intervention* or therap* or interview* or advice)):ti,ab
 #24 (minimal near/3 (intervention* or therap* or interview* or advice)):ti,ab
 #25 (early near/3 (intervention* or therap* or interview* or advice)):ti,ab
 #26 (motivat near/3 (intervention* or therap* or interview* or advice)):ti,ab
 #27 (counselling or counseling or advice):ti,ab
 #28 #17 or #18 or #19 or #20 or #21 or #22 or #23 or #24 or #25 or #26 or #27
 #29 #10 and #16 and #28
 #30 “brief alcohol intervention*”:ti,ab
 #31 #29 or #30

Appendix 2. Cochrane Drugs & Alcohol Group Specialised Register search strategy

Searched 23rd May 2012, subsequently searched via CENTRAL

In the fields “diagnosis” or in title or abstract : alcohol* AND In all fields: counsel* or advice or screen* or ((behaviour* or brief* or minimal or early or motivat*) AND (intervention* or therap* or interview*)) AND In all fields: “general practice” or GP or “primary care” or “primary health” or “family practice” or or emergency or physician* or ((family or community or district or practice*) and (doctor or practitioner* or health* or nurse))

Appendix 3. CENTRAL search strategy

Database(s): EBM Reviews - Cochrane Central Register of Controlled Trials June 2012

Searched 26 June 2012 (superseded by search in [Appendix 1](#))

1. exp General Practice/
2. Primary Health Care/
3. Community Health Services/
4. exp Emergency Service, Hospital/
5. Physicians, Primary Care/ or Physicians, Family/
6. General Practitioners/
7. (general pract\$ or family pract\$ or GP or physician\$ or emergency department\$ or emergency room\$ or trauma care or “accident and emergency” or community).tw.
8. ((family or community or practice*) adj (medic* or doctor* or physician*)).tw.
9. (primary adj2 care).tw.
10. shared care.tw.
11. or/1-10
12. exp Alcohol-Related Disorders/
13. Alcohol Drinking/

14. (alcohol adj3 (drink\$ or use\$ or abus\$ or misus\$ or risk\$ or consum\$ or withdraw\$ or intoxicat\$ or detox\$ or treat\$ or therap\$ or excess\$ or reduc\$ or cessation or intervention)).tw.
15. (drink\$ adj3 (excess or heavy or heavily or hazard\$ or binge or harmful or problem\$)).tw.
16. ("alcohol use" or alcoholic\$).tw.
17. or/12-16
18. Counseling/
19. Psychotherapy, Brief/
20. Interview, Psychological/
21. Behavior Therapy/
22. Cognitive Therapy/
23. Motivation/
24. ((brief or minimal or early or motivat\$) adj3 (intervention\$ or therap\$ or interview\$ or advice)).tw.
25. (counselling or counseling or advice).tw.
26. or/18-25
27. and/11,17,26

Appendix 4. MEDLINE search strategy

Database(s): Ovid MEDLINE(R) In-Process & Other Non-Indexed Citations and Ovid MEDLINE(R) 1946 to September week 2 2017

Searched 21 September 2017 - thesaurus headings updated compared to previous search (see below)

1. exp General Practice/
2. Primary Health Care/
3. Community Health Services/
4. exp Emergency Service, Hospital/
5. Physicians, Primary Care/ or Physicians, Family/
6. General Practitioners/
7. (general pract\$ or family pract\$ or GP or physician\$ or emergency department\$ or emergency room\$ or trauma care or "accident and emergency" or community).tw.
8. ((family or community or practice\$) adj (medic\$ or doctor\$ or physician\$ or health\$ or nurs\$)).tw.
9. (primary adj2 care).tw.
10. shared care.tw.
11. or/1-10
12. exp Alcohol-Related Disorders/
13. Alcohol Drinking/
14. (alcohol adj3 (drink\$ or intoxicat\$ or use\$ or abus\$ or misus\$ or risk\$ or consum\$ or withdraw\$ or detox\$ or treat\$ or therap\$ or excess\$ or reduc\$ or cessation or intervention\$)).tw.
15. (drink\$ adj3 (excess or heavy or heavily or harm or harmful or hazard\$ or binge or harmful or problem\$)).tw.
16. ("alcohol use" or alcoholic\$).tw.
17. or/12-16
18. Counseling/
19. Psychotherapy, Brief/
20. Interview, Psychological/
21. Behavior Therapy/
22. Cognitive Therapy/
23. Motivation/
24. ((brief or minimal or early or motivat\$) adj3 (intervention\$ or therap\$ or interview\$ or advice)).tw.
25. (counselling or counseling or advice).tw.
26. or/18-25
27. and/11,17,26
28. brief alcohol intervention\$.tw.
29. or/27-28

30. randomized controlled trial.pt.
31. controlled clinical trial.pt.
32. randomi\$.ab.
33. placebo.ab.
34. drug therapy.fs.
35. randomly.ab.
36. trial.ab.
37. groups.ab.
38. 30 or 31 or 32 or 33 or 34 or 35 or 36 or 37
39. (animals not (humans and animals)).sh.
40. 38 not 39
41. 29 and 40
42. limit 41 to yr="2005 -Current"

The following search strategy was used to search MEDLINE (OVID) in 2005

1. family practice/
2. family pract\$.tw.
3. general practice.sh.
4. general pract\$.tw.
5. primary health care/
6. primary care/
7. community health services/
8. Community Care/
9. shared care.mp.
10. Patient Care/ or patient care team.mp.
11. family medicine/
12. family physician/
13. family phys\$.tw.
14. exp alcohol/
15. alcohol\$.tw.
16. 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13
17. 14 or 15
18. 16 and 17
19. alcohol reduction.mp.
20. brief intervention.mp.
21. early intervention.mp.
22. minimal intervention.mp.
23. alcohol therapy.mp.
24. harm reduction,.mp.
25. screening.mp.
26. (counseling or counselling).mp.
27. controlled drinking.mp.
28. (brief counseling or brief counselling).mp.
29. physician based intervention.mp.
30. general practitioner intervention.mp.
31. secondary prevention.mp.
32. general practitioner's advice.mp.
33. brief physician-delivered counseling.mp.
34. brief nurse-delivered counseling.mp.
35. identification.mp.
36. intervention.mp.
37. 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 29 or 30 or 31 or 32 or 33 or 34 or 35 or 36
38. (alcohol or alcohol consumption).mp.
39. 37 and 38

40. 39 and 18
41. randomized controlled trial.mp.
42. controlled clinical trial.mp.
43. randomized controlled trials.mp.
44. random allocation.mp.
45. double blind method.mp.
46. single blind method.mp.
47. or/41-46
48. (animal not human).mp.
49. 47 not 48
50. clinical trial.mp.
51. exp clinical trials/
52. (clin\$ adj2 trial\$).ti,ab.
53. ((singl\$ or doubl\$ or trebl\$ or tripl\$) adj2 (blind\$ or mask\$)).ti,ab.
54. placebos.mp.
55. placebo\$.ti,ab.
56. random\$.ti,ab.
57. research design.mp.
58. or/50-57
59. 58 not 48
60. 59 not 49
61. comparative study.mp.
62. exp evaluation studies/
63. follow up studies.mp.
64. prospective studies.mp.
65. (control\$ or prospectiv\$ or volunteer\$).ti,ab.
66. or/61-65
67. 66 not 48
68. 66 not (49 or 60)
69. 49 or 60 or 68
70. 69 and 40

Appendix 5. Embase search strategy

Database(s): Embase 1980 to 2017 week 38

Searched 21 September 2017

1. random\$.tw.
2. clinical trial\$.mp.
3. exp treatment outcome/
4. or/1-3
5. general practice/
6. exp primary health care/
7. community care/
8. emergency health service/
9. general practitioner/
10. (general pract\$ or family pract\$ or GP or physician\$ or emergency department\$ or emergency room\$ or trauma care or "accident and emergency" or community).tw.
11. ((family or community or practice\$) adj (medic\$ or doctor\$ or physician\$ or health\$ or nurs\$)).tw.
12. (primary adj2 care).tw.
13. shared care.tw.
14. or/5-13
15. alcoholism/

16. alcohol intoxication/
17. alcohol abuse/
18. drinking behavior/
19. (alcohol adj3 (drink\$ or intoxicat\$ or use\$ or abus\$ or misus\$ or risk\$ or consum\$ or withdraw\$ or detox\$ or treat\$ or therap\$ or excess\$ or reduc\$ or cessation or intervention)).tw.
20. (drink\$ adj3 (excess or heavy or heavily or hazard\$ or binge or harmful or problem\$)).tw.
21. (“alcohol use” or alcoholic\$).tw.
22. or/15-21
23. counseling/
24. psychotherapy/
25. psychologic test/
26. behavior therapy/
27. cognitive therapy/
28. motivation/
29. ((brief or minimal or early or motivat\$) adj3 (intervention\$ or therap\$ or interview\$ or advice)).tw.
30. (counselling or counseling or advice).tw.
31. or/23-30
32. 14 and 22 and 31
33. brief alcohol intervention\$.tw.
34. or/32-33
35. 4 and 34

Appendix 6. CINAHL search strategy

CINAHL (EBSCO) 1981-2017

Searched 25 September 2017

S31 S18 AND S30

S30 S19 or S20 or S21 or S22 or S23 or S24 or S25 or S26 or S27 or S28 or S29

S29 TX allocat* random*

S28 (MH “Quantitative Studies”)

S27 (MH “Placebos”)

S26 TX placebo*

S25 TX random* allocat*

S24 (MH “Random Assignment”)

S23 TX randomi* control* trial*

S22 TX ((singl* n1 blind*) or (singl* n1 mask*))

S21 TX clinic* N1 trial*

S20 PT Clinical trial

S19 (MH “Clinical Trials+”)

S18 S16 OR S17

S17 TI (“brief alcohol intervention*”) OR AB (“brief alcohol intervention*”)

S16 S5 and S10 and S15

S15 S11 or S12 or S13 or S14

S14 TI (counseling OR counselling OR advice) OR AB (counseling OR counselling OR advice)

S13 TI ((brief N3 intervention*) OR (brief N3 therap*) OR (brief N3 interview*) OR (minimal N3 intervention*) OR (minimal N3 therap*) OR (minimal N3 interview*) OR (early N3 intervention*) OR (early N3 therap*) OR (early N3 interview*) OR (motivat* N3 intervention*) OR (motivat* N3 therap*) OR (motivat* N3 interview*)) OR AB ((brief N3 intervention*) OR (brief N3 therap*) OR (brief N3 interview*) OR (minimal N3 intervention*) OR (minimal N3 therap*) OR (minimal N3 interview*) OR (early N3 intervention*) OR (early N3 therap*) OR (early N3 interview*) OR (motivat* N3 intervention*) OR (motivat* N3 therap*) OR (motivat* N3 interview*))

S12 (MH “Behavior Therapy+”)

S11 (MH “Counseling”) OR (MH “Motivational Interviewing”)

S10 S6 or S7 or S8 or S9
 S9 TI (alcohol N2 use*) OR AB (alcohol N2 use*)
 S8 TI (drink* N3 (heavy OR heavily OR hazard* OR binge OR harmful)) OR AB (drink* N3 (heavy OR heavily OR hazard* OR binge OR harmful))
 S7 TI (alcohol N3 (drink* OR abus* OR misus* OR risk* OR consum* OR withdraw* OR intoxicat* OR detox* OR treat* OR therap* OR excess* OR reduc* OR cessation OR intervention)) OR AB (alcohol N3 (drink* OR abus* OR misus* OR risk* OR consum* OR withdraw* OR intoxicat* OR detox* OR treat* OR therap* OR excess* OR reduc* OR cessation OR intervention))
 S6 (MH "Alcohol-Related Disorders") OR (MH "Alcohol Abuse") OR (MH "Alcoholic Intoxication") OR (MH "Alcoholism") OR (MH "Alcohol Drinking") OR (MH "Alcohol Rehabilitation Programs")
 S5 S1 or S2 or S3 or S4
 S4 TI (shared care) OR AB (shared care)
 S3 TI (primary N2 care) OR AB (primary N2 care)
 S2 TI (general pract* OR family pract* OR GP OR physician* OR emergency department OR emergency room OR "accident and emergency" OR community) OR AB (general pract* OR family pract* OR GP OR physician* OR emergency department* OR emergency room* OR trauma care OR "accident and emergency" OR community)
 S1 (MH "Family Practice") OR (MH "Primary Health Care") OR (MH "Community Health Services") OR (MH "Rehabilitation, Community-Based") OR (MH "Emergency Service+") OR (MH "Physicians, Emergency") OR (MH "Physicians, Family")

Appendix 7. PsycINFO search strategy

Database(s): PsycINFO 1806 to September week 3 2017

Searched 21 September 2017

1. primary health care/
2. general practitioners/
3. emergency services/
4. family physicians/
5. (general pract\$ or family pract\$ or GP or physician\$ or emergency department\$ or emergency room\$ or trauma care or "accident and emergency" or community).ti,ab.
6. ((family or community or practice*) adj (medic* or doctor* or physician*)).tw.
7. (primary adj2 care).ti,ab.
8. shared care.ti,ab.
9. or/1-8
10. exp alcohol intoxication/
11. exp alcohol abuse/
12. alcohol rehabilitation/
13. alcohol drinking patterns/
14. (alcohol adj3 (drink\$ or intoxicat\$ or use\$ or abus\$ or misus\$ or risk\$ or consum\$ or withdraw\$ or detox\$ or treat\$ or therap\$ or excess\$ or reduc\$ or cessation or intervention)).ti,ab.
15. (drink\$ adj3 (excess or heavy or heavily or hazard\$ or binge or harmful or problem\$)).ti,ab.
16. ("alcohol use" or alcoholic\$).ti,ab.
17. or/10-16
18. counseling/ or rehabilitation counseling/
19. brief psychotherapy/
20. motivational interviewing/
21. cognitive behavior therapy/ or behavior therapy/
22. ((brief or minimal or early or motivat\$) adj3 (intervention\$ or therap\$ or interview\$)).ti,ab.
23. (counselling or counseling or advice).ti,ab.
24. or/18-23
25. 9 and 17 and 24
26. brief alcohol intervention\$.tw.
27. or/25-26
28. control\$.tw.

29. random\$.tw.
30. exp Treatment/
31. or/28-30
32. 27 and 31

Appendix 8. Science Citation Index, Social Science Citation Index, Conference Proceedings Citation Index-Science, Emerging Sources Citation Index search strategy

SCI-EXPANDED/SSCI (Web of Science) 1970-2017

CPCI-S (Web of Science) 1990-2017

ESCI (Web of Science) 2015-2017

Searched 25 September 2017

15 #14 AND #13

14 TS=randomi* OR TS=randomly OR TS=placebo* OR TS=trial*

13 #11 AND #8 AND #4

12 #11 AND #8 AND #4

11 #10 OR #9

10 TS=(counselling or counseling or advice)

9 TS=((brief NEAR/3 intervention*) OR (brief NEAR/3 therap*) OR (brief NEAR/3 interview*) OR (minimal NEAR/3 intervention*) OR (minimal NEAR/3 therap*) OR (minimal NEAR/3 interview*) OR (early NEAR/3 intervention*) OR (early NEAR/3 therap*) OR (early NEAR/3 interview*) OR (motivat* NEAR/3 intervention*) OR (motivat* NEAR/3 therap*) OR (motivat* NEAR/3 interview*))

8 #7 OR #6 OR #5

7 TS="alcohol use" OR TS=alcoholic*

6 TS=((drink NEAR/3 excess) OR (drink NEAR/3 heavily) OR (drink* NEAR/3 heavy) OR (drink* NEAR/3 hazard*) OR (drink* NEAR/3 binge) OR (drink* NEAR/3 harmful) OR (drink* NEAR/3 problem*))

5 TS=((alcohol NEAR/3 drink*) OR (alcohol NEAR/3 intoxicat*) OR (alcohol NEAR/3 abus*) OR (alcohol NEAR/3 misus*) OR (alcohol NEAR/3 risk*) OR (alcohol NEAR/3 consum*) OR (alcohol NEAR/3 withdraw*) OR (alcohol NEAR/3 detox*) OR (alcohol NEAR/3 treat*) OR (alcohol NEAR/3 therap*) OR (alcohol NEAR/3 excess*) OR (alcohol NEAR/3 reduc*) OR (alcohol NEAR/3 cessation) OR (alcohol NEAR/3 intervention))

4 #3 OR #2 OR #1

3 TS="shared care"

2 TS=("primary care" OR "primary health care")

1 TS=("general pract*" OR "family pract*" OR GP OR physician* OR "emergency department" OR "accident and emergency" OR community)

Appendix 9. Clinicaltrials.gov search strategy

<http://clinicaltrials.gov/>

Searched 25 September 2017

Advanced search

Search terms: alcohol

Interventions: brief

Appendix 10. Risk of bias assessment criteria

Item	Judgment	Description
1. Random sequence generation (selection bias)	Low risk	The investigators describe a random component in the sequence generation process such as: random number table, computer random number generator, coin tossing, shuffling cards or envelopes, throwing dice, drawing of lots, minimisation
	High risk	The investigators describe a non-random component in the sequence generation process such as: odd or even date of birth; date (or day) of admission; hospital or clinic record number; alternation; judgement of the clinician; results of a laboratory test or a series of tests; availability of the intervention
	Unclear risk	Insufficient information to permit judgement of low or high risk
2. Allocation concealment (selection bias)	Low risk	Investigators enrolling participants could not foresee assignment because one of the following, or an equivalent method, was used to conceal allocation: central allocation (including telephone, web-based, and pharmacy-controlled, randomisation); sequentially numbered, opaque, sealed envelopes. Very few studies which used envelopes to conceal allocation described them as serially numbered, opaque <i>and</i> sealed, but some studies nevertheless reported enough to suggest adequate allocation concealment. We used all the text describing allocation concealment to inform our judgement of this item
	High risk	Investigators enrolling participants could possibly foresee assignments because one of the following method was used: open random allocation schedule (e.g. a list of random numbers); assignment envelopes without appropriate safeguards (e.g. if envelopes were unsealed or nonopaque or not sequentially numbered); alternation or rotation; date of birth; case record number; any other explicitly unconcealed procedure
	Unclear risk	Insufficient information to permit judgement of low or high risk
3. Blinding of providers (performance bias)	Low risk	No blinding or incomplete blinding, but the review authors judge that the outcome is not likely to be influenced by lack of blinding Blinding of providers and unlikely that the blinding could have been broken If the study was a cluster randomised study where the arms were physically separated and the intervention providers only had contact with a single arm If the pre-intervention steps (e.g. screening, randomisation, baseline assessment) were carried out by an independent person (e.g. researcher, research or practice nurse), and the intervention provider only had contact with the intervention arm, so that there was no chance of 'contamination' in the control arm

(Continued)

	High risk	No blinding or incomplete blinding, and the outcome is likely to be influenced by lack of blinding; Blinding of key study participants and personnel attempted, but likely that the blinding could have been broken, and the outcome is likely to be influenced by lack of blinding
	Unclear risk	Insufficient information to permit judgement of low or high risk
4. Blinding of participants (performance bias)	Low risk	Blinding of participants and unlikely that the blinding could have been broken At screening or assessment, participants were unaware that the trial specifically focused on alcohol consumption, e.g. tools encompassed wider health or lifestyle questions
	High risk	No blinding or incomplete blinding, and the outcome is likely to be influenced by lack of blinding; Blinding of key study participants and personnel attempted, but likely that the blinding could have been broken, and the outcome is likely to be influenced by lack of blinding Studies received “high risk” as default unless there was an explicit attempt to blind participants
	Unclear risk	Insufficient information to permit judgement of low or high risk
5. Blinding of outcome assessors (detection bias)	Low risk	No blinding of outcome assessment, but the review authors judge that the outcome measurement is not likely to be influenced by lack of blinding Blinding of outcome assessors and unlikely that the blinding could have been broken
	High risk	No blinding or incomplete blinding
	Unclear risk	Outcome collection was not automated and insufficient information is provided to assess blinding
6. Incomplete outcome data (attrition bias) For all outcomes except retention in treatment or drop out	Low risk	No missing outcome data; Reasons for missing outcome data unlikely to be related to true outcome (for survival data, censoring unlikely to be introducing bias); Missing outcome data balanced in numbers across intervention groups, with similar reasons for missing data across groups; For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk not enough to have a clinically relevant impact on the intervention effect estimate; For continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes not enough to have a clinically relevant impact on observed effect size; Missing data have been imputed using appropriate methods All randomised patients are reported/analysed in the group they were allocated to by randomisation irrespective of non-compliance and co-

(Continued)

		interventions (intention to treat)
	High risk	Reason for missing outcome data likely to be related to true outcome, with either imbalance in numbers or reasons for missing data across intervention groups; For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk enough to induce clinically relevant bias in intervention effect estimate; For continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes enough to induce clinically relevant bias in observed effect size; 'As-treated' analysis done with substantial departure of the intervention received from that assigned at randomisation
	Unclear risk	Insufficient information to permit judgement of low or high risk (e.g. number randomised not stated, no reasons for missing data provided; number of drop out not reported for each group)
7. Selective reporting (reporting bias)	Low risk	The study protocol is available and all of the study's pre-specified (primary and secondary) outcomes that are of interest in the review have been reported in the pre-specified way; The study protocol is not available but it is clear that the published reports include all expected outcomes, including those that were pre-specified (convincing text of this nature may be uncommon)
	High risk	Not all of the study's pre-specified primary outcomes have been reported; One or more primary outcomes is reported using measurements, analysis methods or subsets of the data (e.g. subscales) that were not pre-specified; One or more reported primary outcomes were not pre-specified (unless clear justification for their reporting is provided, such as an unexpected adverse effect); One or more outcomes of interest in the review are reported incompletely so that they cannot be entered in a meta-analysis; The study report fails to include results for a key outcome that would be expected to have been reported for such a study
	Unclear risk	Insufficient information to permit judgement of low or high risk

FEEDBACK

Anders Beich, 27 May 2009

Summary

The effect of brief interventions for hazardous drinking among primary care patients has been investigated in several controlled trials and meta-analyses. In most trials accomplished during the last 25 years intervention groups, mostly men, have subsequently reported lower alcohol consumption than their matching controls at follow-up. Therefore, it is no surprise that Kaner et al. in their meta-analysis [1] conclude that brief alcohol intervention in primary care contexts results in significant reductions in weekly consumption at least for men.

The crucial point is whether these results represent effectiveness rather than efficacy [2], because if so then health care professionals should be able to contribute to less hazardous drinking habits among their patients by adopting this technology. One question to address could then be how the technology fits within the context of primary care and how these results could be used out there where health professionals meet patients on a continuous basis.

The Cochrane-analysis [1] aims to examine to which degree the included trials have reported results that are externally valid and it is claimed that "The lack of evidence of any difference in outcomes between efficacy and effectiveness trials suggests that the current literature is relevant to routine primary care and we feel that the current body of brief alcohol intervention research is applicable to clinical practice. I find this part of the analysis/interpretation to be problematic and the methods they use for reaching this conclusion to be inappropriate. The strength of their argument is obviously weak and their way of reasoning is deceptive. Just because two objects have the same colour you can hardly conclude, that this suggests they can be used for a specific purpose, and feelings should in my opinion be kept out of the argumentation.

I find the analysis and the discussion of the meta-analysis results to have several serious weaknesses:

1) Which drinkers are we dealing with?

The heterogeneity among trial groups in relation to baseline consumption is substantial among the included studies. Average consumption ranges from about 17 drinks/week to 54, that is, from slightly hazardous drinking to obviously harmful consumption and possible dependency. The risk reduction by reducing from 54 to 49 drinks per week can hardly be comparable to a reduction from 17 to 12 drinks per week and the report lacks a qualified discussion of this issue.

2) Study quality analysis as an invocation

Although a quality analysis of the included studies is carried out, the results are not allowed to affect the calculation of average intervention effects: Poor quality studies are given the same weight as high quality studies. I am aware that this is often the case in meta-analyses, but when results rely on self-reports I believe this issue is at least worth a discussion, which leads to the next weakness

3) The "Garbage in" "garbage out" problem

The analysis is not concerned with the quality and quality control of data (self-reported drinking). Self-reported drinking has been shown to be influenced by situational and contextual matters and the impossibility of blinding of study subjects may constitute a major source of bias in several of the included studies and this is not discussed in the report. A sensitivity analysis focusing on effects in studies that had other more objective data available could have been attempted or at least discussed.

4) Biased effectiveness perspective

One of the aims of the work is to investigate to which degree the included trials can be characterized as effectiveness trials and thereby relevant to routine primary care. This is done by analysing to which degree the setting can be characterized as a clinical rather than a research setting (reasons for encounter, the advisor/interventionist and the character of the clinic, as well as support, help and supervision provided in clinical trials).

The analysis avoids to analyse to which degree the trial groups can be characterized as clinical rather than research groups, that is, no attempts were made to find out how representative the included individuals were in relation to how many were actually screened hazardous drinking positive in the first place. Because screening and brief intervention recommendations come together [3] brief intervention seems to involve some kind of screening as it did the included trials it should at least have been clarified how refined the trial groups were compared to the groups that screened positive at baseline.

An example of the problem: A heavy weight study in the present meta-analysis is a study by Fleming et al.[4], who screened 17,695 patients, found 2,925 (17%) to be hazardous drinkers, of which 774 were included in the trial after all. That is, 2,151 hazardous drinkers were lost before the trial started for reasons poorly described. It is indeed possible that the most interested (/ready /motivated) drinkers were over represented in the trial groups. This may in more than one way lead to biased results and it is not justified to claim that the results do also count for the majority of hazardous drinkers who were excluded in this study.

This issue is not at all new to most of the Cochrane-group though they manage to maintain absolute silence about the lost-subjects-before-randomization issue and a meta-analysis on exactly that issue in 2003[5], one that caused them to react very strongly at the time and later on in public accuse the authors of the analysis for causing the death of hazardous drinkers because these discussions might delay implementation efforts that they were in charge of.

5) Conflict of interests

The trustworthiness of the interpretation by Kaner et al. is in my opinion severely weakened by the fact that the authors claim no conflict of interests when more of them are well known for their involvement in implementation activities regarding brief interventions for decades. It is an open question what should be included as conflict of interests, but when more of the authors, the first author inclusive, have had implementation of these brief intervention technologies as a main employment for years, it seems unlikely that this would not affect which questions they wish to answer and the basic choices they make in the process of a meta-analysis of the effectiveness of the very technology.

When the authors state that they have made inquiries on non-published studies I find it strange that they do not mention a Danish pragmatic randomised trial [7] that was not yet published when they ended their literature search, but I have personally presented the results for at least three of the authors more than once, results indicating that brief intervention had no effect on men and might be worse than no intervention for women when it was put out in real life circumstances and all recommendations were followed. Also, other results from this Danish group of independent researchers indicating that screening based brief intervention is not compatible with the work of the general practitioner [8] and that for the documentation so far rest on highly refined trial groups [5] are carefully ignored in this Cochrane-paper.

This meta-analysis tells us that some hazardous drinking male patients in primary health care can benefit from a brief intervention. But we can not say anything about the proportion, how much they will reduce their drinking or how we should find them from this analysis. I find the approach of this meta-analysis to be selective and biased and the purpose of it to be doubtful.

I propose that The Cochrane Collaboration encourage future author groups to declare all possible conflict of interests, not just obviously commercial ones.

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1. Kaner EF, Beyer F, Dickinson HO et al. Effectiveness of brief alcohol interventions in primary care populations. *Cochrane Database Syst Rev* 2007:CD004148.
2. Flay BR. Efficacy and effectiveness trials (and other phases of research) in the development of health promotion programs. *Prev Med* 1986;15:451-74.
3. Babor TF, Higgins-Biddle JC. *Brief Intervention for Hazardous and Harmful Drinking - A Manual for Use in Primary Care*. Geneva: World Health Organization, 2001: 1-29
4. Fleming MF, Barry KL, Manwell LB et al. Brief physician advice for problem alcohol drinkers - A randomised controlled trial in community-based primary care practices. *JAMA* 1997;277:1039-45.
5. Beich A, Thorsen T, Rollnick S. Screening in brief intervention trials targeting excessive drinkers in general practice: systematic review and meta-analysis. *BMJ* 2003;327:536-40.
6. Kaner EF, Lock CA, McAvoy BR, Heather N, Gilvray E. A RCT of three training and support strategies to encourage implementation of screening and brief alcohol intervention by general practitioners. *Br J Gen Pract* 1999;49:699-703.
7. Beich A, Gannik D, Saelan H et al. Screening and Brief Intervention Targeting Risky Drinkers in Danish General Practice A Pragmatic Controlled Trial. *Alcohol Alcohol* 2007.
8. Beich A, Gannik D, Malterud K. Screening and brief intervention for excessive alcohol use: qualitative interview study of the experiences of general practitioners. *BMJ* 2002;325:870-3.

Reply

We are pleased that the commentator confirms the validity of our primary review finding that brief interventions are effective at reducing excessive alcohol consumption, particularly in men. We welcome critique of our approach in exploring the efficacy-effectiveness question indeed we pointed out in our discussion some weaknesses that we felt were inherent in our approach. Despite this, we felt it was worth trying to unravel the efficacy-effectiveness issue and believe our work has made a start, even if this is an imperfect approach. This said, we feel there is merit in trying to clarify important parameters such as whether if the interventions are delivered in typical practices by practitioners who deliver primary care as their main occupation and if the interventions are delivered to routinely presenting patients. Furthermore, we also tried to assess whether brief interventions were delivered within normal consultation times and if preparatory training is what most practitioners are likely to be able access. These are just 4 of the 8 parameters we assessed trials on in our attempt to establish the internal or external validity of the trial design.

1) We have described the characteristics of subjects in the brief intervention trials and we state both narratively and statistically in the review that there is a great deal of heterogeneity in the field.

2) The extensive sensitivity analysis that we carried out deals with the range of well-accepted quality criteria for research trials. Studies that were lacking on various quality criteria were omitted from the numerous sensitivity analyses that we reported and the key findings of the review were not substantially altered by this process.

3) This review was based on a range of outcome measures reported in 29 trials, some of which were self-reported by patients and some of which were more objectively measured. Thomas Greenfield and Lorraine Midanik have published a number of papers on the issue of self-report as a measure of alcohol use and have found this to be a valid approach. Furthermore, Babor and colleagues have confirmed that self-reported data if collected in the right way are sufficiently valid for research and less intrusive for patients [Babor T, Steinberg K, Anton RF, Del Boca F. Talk is cheap: measuring drinking outcomes in clinical trials. *Journal of Studies on Alcohol* 2000;61:55-63]. In addition, whilst self-reported drinking may not be a gold-standard measure of alcohol consumption, this approach was employed in both the control and treatment arms of the trials and so this issue would not essentially alter the outcome findings of the review. Lastly, we conducted a meta-analysis which considered the outcomes of individually randomised trials compared to cluster randomised trials (analysis 01.08) and found no statistically significant difference in outcome. Thus we do not think that the self-reported outcomes or the lack of blinding in individually randomised trials substantively affects the findings of our review. Finally, two included trials reported both self-reported outcomes and laboratory markers (GGT) of alcohol intake; both studies were consistent in finding that the intervention was more effective than control on the basis of both self-reported and objective outcomes.

4) We are aware that more individuals are screened than enrolled into the treatment or control conditions of the trials, this is indeed the purpose of screening. The number of patients that need to be screened in order to identify patients suitable for intervention is an important implementation issue for practice (and practitioners) which has been specifically investigated elsewhere (reference 5 below) but it was not central to the aims of our effectiveness review.

In our review, we specifically report the loss to follow-up of patients post intervention which was greater in brief intervention conditions compared to control conditions. We cite this as a weakness in the brief intervention evidence-base. However, the loss of patients between screening and enrolment into the trial was not relevant to the aim of our review which was to ascertain the relative effectiveness of brief intervention versus control conditions in reducing excessive drinking. Since allocation at enrolment was randomly determined, it is clear that post-screening attrition would be equally experienced in both control and brief intervention conditions. Thus this issue would not substantively alter the findings of our review.

The efficiency of screening prior to brief intervention was covered in a BMJ review in 2003 and there was a considerable debate on this issue at the time (in which 2 out of the nine authors of the current review participated). Since the majority of the co-authors were not involved in that debate, it is not accurate or helpful to state that most of the Cochrane group have maintained a silence on the discussion. As stated above, the issue as to whether screening is an efficient means of identifying excessive drinkers in primary care was not relevant to our key aim of evaluating the effectiveness of brief intervention at reducing heavy drinking.

5) Of the nine co-authors of the current review, three have been involved in both implementation research and outcome evaluation trials in the field of brief alcohol interventions. Indeed, two of these individuals were involved in conducting three of the null-effect trials that contributed to the review (Heather et al. 1987, Richmond et al. 1995 and Lock et al. 2006). The statement that implementation of brief intervention technologies has been the main employment for years of any of the review authors is inaccurate. In addition, we do not agree with the proposition that involvement in implementation research is in of itself favourable towards brief intervention (for an example see reference 5) and the findings of null-effect outcome evaluation research highlighted above substantiates this view. Finally, all the meta-analysis conducted in the Cochrane review was conducted by two research scientists who have never previously worked in the brief intervention field the respective contribution of all the review co-authors to each aspect of the review process is clearly outlined on the published review (page 44). Thus we feel confident about our statement concerning conflict of interest regarding the review.

The current review clearly reported that its search strategy extended to 2006. Thus it obviously would not have included reference 7 which was published in 2007. Indeed the initial review analysis was completed in November 2006 (the group was awarded a small grant by the Cochrane Collaboration to complete the review by this time). However, we re-ran our analysis in February 2007 after correspondence from a trial author (Curry 2003) whose work had been excluded from the review in error. The published version of reference 7 states that it was published (following amendment made after peer-review) in July 2007. Thus the peer-reviewed version of this work was not available for this review. However, this work will, of course, be included in any future update of the Cochrane review. Nevertheless, since the current review clearly contains both null-effect and positive-effect trials of brief intervention, it is clear that we have not selectively included only work with positive outcomes. The Cochrane review meta-analysis clearly reports both the average reduction in the percentage of heavy drinkers (Analysis 1.21) and the average weekly reduction in drinking after brief alcohol intervention (Analysis 1.15) along with a wide range of other relevant outcome measures.

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1. Kaner EF, Beyer F, Dickinson HO et al. Effectiveness of brief alcohol interventions in primary care populations. *Cochrane Database Syst Rev* 2007:CD004148.
2. Flay BR. Efficacy and effectiveness trials (and other phases of research) in the development of health promotion programs. *Prev Med* 1986;15:451-74.
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8. Beich A, Gannik D, Malterud K. Screening and brief intervention for excessive alcohol use: qualitative interview study of the experiences of general practitioners. *BMJ* 2002;325:870-3.

Contributors

Feedback from Anders Beich

Reply from Kaner EF, Dickinson HO, Beyer FR, Campbell F, Schlesinger C, Heather N, Saunders JB, Burnand B, Pienaar ED

WHAT'S NEW

Last assessed as up-to-date: 25 September 2017.

Date	Event	Description
25 September 2017	New search has been performed	<ol style="list-style-type: none"> 1. The search was updated in September 2017 and 42 new trials (24,057 participants) were included, bringing the total to 69 included studies (33,642 participants). Two trials in the original review were excluded from this update 2. The review has been substantially revised in accordance with Cochrane guidance, including the implementation of GRADE assessment, creation of 'Summary of findings' tables for the main comparisons, and further subgroup analyses
25 September 2017	New citation required and conclusions have changed	<ol style="list-style-type: none"> 1. The addition of 42 new studies changed conclusions with respect to the effect for women, and enabled further subgroup analyses 2. The review authorship changed.

HISTORY

Protocol first published: Issue 2, 2003

Review first published: Issue 2, 2007

Date	Event	Description
28 July 2009	Feedback has been incorporated	We have replied to feedback from Dr Anders Beich
19 June 2008	Amended	corrected figures, estimates and add minor changes
29 April 2008	Amended	little changes
29 April 2008	New search has been performed	Add one study but conclusions are not changed
15 February 2007	New citation required and conclusions have changed	Substantive amendment

CONTRIBUTIONS OF AUTHORS

Kaner conceived, designed and co-ordinated the review. Kaner and Campbell wrote the original protocol with advice from Heather and Saunders to develop the efficacy/effectiveness scale, from Dickinson on statistical issues and from Pienaar on other issues. Pienaar, Campbell and Kaner conducted the searches. Beyer conducted the updated searches. Campbell and Beyer managed the reference databases. Kaner, Schlesinger, Bertholet, Daeppen, Bernand and Pienaar sifted the references. Kaner, Schlesinger, Pienaar, Campbell, Beyer and Dickinson extracted data. Dickinson and Muirhead performed statistical analysis. Dickinson, Beyer, Campbell and Kaner wrote the review. Muirhead, Campbell, Kaner and Beyer updated the text for this version. All authors read and commented on the updated review.

DECLARATIONS OF INTEREST

Nicolas Bertholet is salaried by Lausanne University Hospital, a public institution; he has received grants from the Swiss National Science Foundation, the Swiss Foundation for Alcohol Research, and the Department of Community Medicine and Health from the Lausanne University Hospital. He has received no personal support from industry sources such as pharmaceutical, alcohol and tobacco companies and holds no personal stock. Dr Bertholet is an author of a previous systematic review on the subject ([Bertholet 2005](#)), and has participated in numerous studies on screening and brief intervention (not limited to the scope of this review), including as a primary investigator of a randomised trial of an electronic screening and brief intervention.

Bernard Burnand, MD, MPH, is Chief Physician at the Institute of Social and Preventive medicine, Lausanne University Hospital. He is salaried full time by the this public institution (Canton of Vaud, Switzerland) and a full professor at the Faculty of Biology and Medicine, University of Lausanne. Bernard Burnand is also Director of Cochrane Switzerland (Cochrane associated centre). His research is supported by Lausanne University Hospital and funding from public granting institutions. ORCID ID: orcid.org/0000-0002-5678-6044

Fiona Campbell has no conflicts of interest to declare.

Jean-Bernard Daeppen received personal fees from Lundbeck SAS for lectures and advice and was involved in the [Daeppen 2007](#) trial, which is an included trial in this systematic review.

Elizabeth Pienaar is in the full-time employ of the South African Medical Research Council and has no conflicts of interest to declare.

John B Saunders has no conflicts of interest to declare.

Eileen Kaner, Fiona Beyer and Colin Muirhead are authors on a related Cochrane Review on interventions for reducing hazardous and harmful alcohol consumption ([Kaner 2017](#)).

Eileen Kaner is an investigator on the ongoing SIPS Junior trial (NIHR programme grant number NIHR RP-PG-0609-10162). She was also involved in the SIPS trial ([Kaner 2013](#)) and the [Lock 2006](#) trial which are included studies in this systematic review.

SOURCES OF SUPPORT

Internal sources

- Small Grant from the University of Newcastle upon Tyne, UK.

External sources

- Department of Health Primary Care Career Scientist Award, UK.
- Cochrane Collaboration, UK.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

Authorship

Dr Heather Dickinson, Dr Carla Schlesinger and Professor Nick Heather were unable to contribute to this review update and are no longer authors. We would like to acknowledge their earlier input. Dr Colin Muirhead, Dr Nicolas Bertholet and Professor Jean-Bernard Daeppen have become authors.

Objective

The objective in the original review contained the sentence “To assess whether outcomes differ between trials in research settings and those in routine clinical settings”, reflecting the fact that there was a strong focus on our efficacy effectiveness tool, and we reported every analysis with trials subgrouped by efficacy or effectiveness. This update does not contain such a strong focus on efficacy/effectiveness because validated tools have superseded ours. The review includes a single subgroup analysis focused on efficacy/effectiveness so no longer warrants a statement in the objective.

Included studies

When compared to the broader range of trials available to include in this review update, two previously included studies did not really fit and we decided it would be more robust to exclude them from the review. Participants in the [Gentilello 1999](#) trial were inpatients and the trial is included in a Cochrane Review of brief interventions in patients admitted to hospital. [Chang 1997](#) compared brief intervention to alcohol treatment referral, which is not an analogous control group with the included trials.

Outcomes

Although not specified in the protocol, we reported when participants were categorised as 'heavy drinker' or 'binge drinker' at follow up.

Meta-analyses

All analyses comprise 12 month outcomes, except where otherwise stated.

Some of the studies added for this update provided data for a new comparison (extended versus minimal or no intervention) and, on reflection, some studies from the previous version of the review also fitted into this comparison and were added.

Subgroup analyses

1. Emergency versus general practice settings

Since publication of the previous version of this review, the effectiveness of brief interventions has been tested more commonly in emergency departments. It is plausible that patients in emergency departments may have a different response to a brief intervention, particularly if alcohol consumption has contributed to their visit. Therefore, we carried out a subgroup analysis of emergency versus general practice settings.

2. Theoretical basis or modality of intervention

Another change over time is that more brief interventions have incorporated counselling or motivational interviewing techniques. Again, it is plausible this would have an impact on the effectiveness of the intervention.

3. Control group content

As described in the [Discussion](#), more recent studies exploring the screening and assessment reactivity effect have been published since our 2007 review appeared. We decided to explore effects in a subgroup analysis according to the intervention received by control group participants: explicitly alcohol-related content versus not.

4. Trials included in the previous version of this review

Given the modifications made since our 2007 review was published, we also performed subgroup analysis according to whether trials appeared in the previous version or the updated version of the review.

5. Efficacy/effectiveness score

Meta-analyses were dichotomised according to trials effectiveness/efficacy scores to assess whether these were relevant to clinical settings in the 2007 review. However, little difference was detected between trials categorised as efficacy versus effectiveness trials; hence, we conducted a single subgroup analysis only for this update.

6. Loss to follow-up

We planned analyses linked of losses to follow up for both this update and the 2007 review. However, because of differing definitions of loss to follow up, and despite many trials presenting results adjusted for missing data, it was not always clear if allowance had been made for loss to follow up. It was decided that a sensitivity analysis of losses to follow-up would not be practicable. However, an analysis that excluded trials at high risk of bias due to incomplete outcome data gave similar results to those from the primary meta-analysis.

Additional tables

'Summary of Findings' tables and risk of bias tables have been added to the review.

INDEX TERMS

Medical Subject Headings (MeSH)

Alcohol Drinking [*therapy]; Alcoholism [*therapy]; Emergencies; Family Practice; Randomized Controlled Trials as Topic

MeSH check words

Humans