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Notes

To order reprints of this article go to: http://www.bmjjournals.com/cgi/reprintform The difference may be partly accounted for by the factors that influence infectivity of the two viruses. Hepatitis B is much more likely to be transmitted from mother to infant if there is a high concentration of the virus in the mother's blood. This explains the ethnic differences that are observed—for example, the transmission rate is over 70% in Chinese women but less than 10% in white women. This ethnic difference does not seem to apply to hepatitis C infection.

Alcohol intake and obesity are both thought to be associated with more severe hepatitis C, although the exact interaction is unknown. Advanced liver disease, for example, is far worse in people infected with hepatitis C who also have a high alcohol intake than in those with a low intake. About half of patients with hepatitis B infections respond to interferon compared with 15% with hepatitis C. Ongoing trials of interferon and antivirals together may prove more fruitful. Although infection with hepatitis C virus does not necessarily cause abnormal liver function, precirrhotic damage confirmed by biopsy is one reason for starting treatment with interferon.

Abi Berger Science editor, BMJ

Deliberate self harm: systematic review of efficacy of psychosocial and pharmacological treatments in preventing repetition

Keith Hawton, Ella Arensman, Ellen Townsend, Sandy Bremner, Eleanor Feldman, Robert Goldney, David Gunnell, Philip Hazell, Kees van Heeringen, Allan House, David Owens, Isaac Sakinofsky, Lil Träskman-Bendz

Abstract

Objective: To identify and synthesise the findings from all randomised controlled trials that have examined the effectiveness of treatments of patients who have deliberately harmed themselves.

Design: Systematic review of randomised controlled trials of psychosocial and physical treatments. Studies categorised according to type of treatment. When there was more than one investigation in a particular category a summary odds ratio was estimated with the Mantel-Haenszel method.

Setting: Randomised trials available in electronic databases in 1996, in the Cochrane Controlled Trials Register in 1997, and from hand searching of journals to 1997.

Subjects: Patients who had deliberately harmed themselves shortly before entry into the trials with information on repetition of behaviour. The included trials comprised 2452 randomised participants with outcome data.

Main outcome measure: Repetition of self harm. **Results:** 20 trials reported repetition of self harm as an outcome variable, classified into 10 categories. Summary odds ratio (all for comparison with standard aftercare) indicated reduced repetition for problem solving therapy (0.73; 95% confidence interval 0.45 to 1.18) and for provision of an emergency contact card in addition to standard care (0.45; 0.19 to 1.07). The summary odds ratios were 0.83 (0.61 to 1.14) for trials of intensive aftercare plus outreach and 1.19 (0.53 to 2.67) for antidepressant treatment compared with placebo. Significantly reduced rates of further self harm were observed for depot flupenthixol versus placebo in multiple repeaters (0.09; 0.02 to 0.50) and for dialectical behaviour therapy versus standard aftercare (0.24; 0.06 to (0.93)

Conclusion: There remains considerable uncertainty about which forms of psychosocial and physical treatments of patients who harm themselves are most effective. Further larger trials of treatments are needed.

Introduction

Prevention of suicide is now included in health policy initiatives in several countries, and reduction in suicidal behaviour, both fatal and non-fatal, is part of the Health for All targets of the World Health Organisation.¹ In the United Kingdom, reduction in the number of suicides is a central theme in the government's Health of the Nation strategy for England.² There is, however, a considerable lack of information as to which preventive strategies are effective.3 Improvement of outcome after deliberate self harm is an important focus because at least 1% of patients presenting to general hospitals in the United Kingdom after deliberate self harm kill themselves within a year and 3-5% do so within 5-10 years. A history of multiple episodes of deliberate self harm is a particular risk factor.⁴ Higher rates of suicide after deliberate self harm have been reported from other countries.5 6 About half of all people who kill themselves have a history of deliberate self harm, an episode having occurred within the year before death in 20-25%.7 8

It would be difficult to investigate the effectiveness of intervention strategies after deliberate self harm in terms of subsequent actual suicides because extremely large populations of patients would be required. Repetition of deliberate self harm is, however, a reasonable proxy measure because of its strong associations with suicide. It is also in itself an important outcome because it occurs frequently,^{9 10} indicates persistent distress, and results in considerable healthcare costs. Deliberate self harm is common in Europe¹¹ and in other parts of the world,^{12 13} especially in young people. Recent marked increases in rates of deliberate self harm in the United Kingdom,14 15 with a currently estimated 140 000 hospital referrals in England and Wales,10 have highlighted the need for effective aftercare strategies.

Descriptive reviews of treatment outcomes in patients who deliberately harm themselves have been

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Correspondence to: Professor Hawton Keith.Hawton@ psychiatry.ox.ac.uk published previously but have not included systematic screening of the literature, quality ratings, and meta-analysis,^{16–18} and have been based on heterogeneous groupings of treatments which do not inform clinical practice.¹⁹

We conducted a systematic review of the worldwide literature regarding treatment studies of patients who deliberately harm themselves. We identified all randomised controlled trials evaluating psychosocial or physical treatments and conducted a meta-analysis to compare the effects of specific treatments on repetition of deliberate self harm with those of control or comparison treatments to identify the most effective interventions.

Methods

Identification of relevant trials-We carried out a literature search using the following electronic databases: Medline 1966 to May 1996; PsycLit 1974 to August 1996; Embase 1980 to November 1996; and the Cochrane Controlled Trials Register (No 4, 1997).²⁰ A wide range of keywords to indicate attempted suicide and a standard search strategy, developed for the Medline database by the Cochrane Collaboration, was used to identify relevant randomised controlled trials. A shorter version of this search strategy was used to perform searches on PsycLit and Embase (details of search strategies available from the authors). In addition, we hand searched 10 journals in the specialty of psychiatry and psychology that had not been searched within the Cochrane Collaboration, including all the English language journals concerned with suicide.

Inclusion criteria—We included studies in the review if they met the following criteria: study participants had to have engaged in deliberate self harm (self poisoning or self injury) shortly before entry into the trial; trials must have reported repetition of deliberate self harm as an outcome measure; and study participants had to have been randomised to treatment and control groups. As long as these three criteria were met we included papers reporting any comparison between different types of treatment, including comparisons with standard (that is, routine) aftercare. When details of standard aftercare were not provided, we attempted to obtain these from the authors.

Grouping of studies-Studies that shared similar treatment strategies were grouped by consensus of the reviewers, blind to the outcome data. The first category (problem solving therapy v standard aftercare) included studies in which participants in the experimental group were offered some form of problem solving therapy which was compared with standard aftercare. Standard aftercare, both here and in other categories, included the usual range of treatment options that were available in routine care at the time in each setting. The second group (intensive intervention plus outreach v standard aftercare) included studies in which the patients in the experimental group had greater access to therapists than in standard care and where efforts were made to keep contact with patients through some form of outreach (for example, home based treatment either as standard or for those patients who defaulted on appointments at a clinic). The third group (emergency card v standard aftercare)

included studies in which patients in the experimental group, in addition to being offered standard aftercare, were given an emergency contact card with which they had 24 hour access to emergency advice from a psychiatrist²¹ or could admit themselves to hospital.²² In only one other group (antidepressant medication v placebo) was there more than one trial. The remainder of the studies are reported singly.

Data extraction and quality assessment-Data were extracted independently by two reviewers. The quality of the papers was rated by two independent reviewers blind to authorship, according to the recommended Cochrane criteria for quality assessment.²³ This rating system is influenced by the finding that the quality of concealment of random allocation can affect the results of trials.24 Studies were assigned a quality score from 1 (poorest quality) to 3 (best quality). Thus, trials rated as inadequately concealed (for example, via alternation or reference to an open random number table) were given a score of 1. Trials that did not give adequate details about how the randomisation procedure was carried out were given a score of 2. Trials that were deemed to have taken adequate measures to conceal allocation (for example, serially numbered, opaque, sealed envelopes; numbered or coded bottles or containers) were given a score of 3. We contacted authors of trials for more information when the concealment of allocation was not clearly reported (that is, when trials were initially in category 2). Blinding of observers was rated according to whether it was absent or unclear, reported but without details, and fully reported.

Statistical methods—Summary odds ratios were calculated with RevMan 3.0 software²⁵ with the Mantel-Haenszel method. Heterogeneity was tested with a χ^2 test.

Results

Twenty studies were identified through the combined search strategies as eligible for inclusion in the study. All reports had been published, although two studies had not been published in full.^{16 26} The full report of one trial was provided by the author in the form of an unpublished manuscript,¹⁶ and a detailed report of the other was obtained from conference proceedings.²⁶ Four studies had been reported in more than one publication.^{16 26-28} One further randomised controlled trial of patients who deliberately harmed themselves was identified, but this did not include repetition of deliberate self harm as an outcome variable.²⁹ We were unable to obtain this information from the authors of this trial.

The trials identified were grouped as described in the methods section. Table 1 summarises the 20 trials included in the review, their groupings, details of participants (sex and the proportion with a history of self harm—"repeaters"), the interventions used, and the quality of concealment scores. Only one trial was specifically of adolescents.²²

The assessment of quality of concealment of allocation (table 1) resulted in 13 trials being given a score of 3 (for adequate concealment), three trials being given a score of 2 (unclear concealment), and four trials being given a score of 1 (inadequate concealment). Blinding of assessors was not stated or

Table 1 Summary of participants, interventions, follow up period, size of trial, and quality of concealment of allocation

Study	Details of participants	Interventions	No randomised (No lost to follow up or excluded)	Follow up period	Quality of concealment of allocation*
Problem solving thera	by v standard aftercare				
Gibbons et al (UK, 1978) ³⁰	Patients over 17 years who presented to A&E department after deliberate self poisoning; repeaters (\geq 1 attempt) and first timers; 71% female	Experimental (n=200): crisis orientated, time limited, task centred social work at home (problem solving intervention). Control (n=200): routine service—54% GP referral, 33% psychiatric referral, 13% other referral	400	12 months	3
Hawton et al (UK, 1987) ³¹	Patients over 16 years admitted to general hospital for self poisoning; 31% repeaters; 66% female	Experimental (n=41): outpatient problem orientated therapy by non-medical clinicians. Control (n=39): GP care (for example, individual support, marital therapy) after advice from clinician	80	12 months	3
Salkovskis et al (UK, 1990) ³²	Patients aged 16-65 years (mean 27.5) referred by duty psychiatrist after antidepressant self poisoning assessed in A&E department; all repeaters with high risk of further repetition; 50% female	Experimental (n=12): domicillary cognitive behavioural problem solving treatment. Control (n=8): treatment as usual (GP care)	20	12 months	3
McLeavey et al (Ireland, 1994) ³³	Patients aged 15-45 years (mean 24.4) admitted to A&E department after self poisoning; 35.6% repeaters; 74% female	Experimental (n=19): interpersonal problem solving skills training. 34 Control (n=20): brief problem solving therapy 35	39	12 months	1
Intensive care plus out	treach v standard care				
Chowdhury et al (UK, 1973) ³⁶	Patients (all repeaters) admitted to general hospital after deliberate self harm; 57% female	Experimental (n=71): special aftercare—regular outpatient appointments; patients also seen without appointments; home visits to patients who missed appointments; emergency 24 hour telephone access. Control (n=84): normal aftercare—outpatient appointment with psychiatrist and/or social worker; non-attenders not pursued	155	6 months	1
Welu (USA, 1977) ³⁷	Suicide attempters over 16 years brought to A&E department; 60% repeaters; % female not given	Experimental (n=63): special outreach programme—community mental health team contacted patient immediately after discharge; home visit arranged; weekly/twice weekly contact with therapist. Control (n=57): routine care—appointment for evaluation at the community mental health centre next day at request of treating physician	120 (1)	4 months	1
Hawton et al (UK, 1981) ³⁸	Patients ≥16 years (mean 25.3) admitted to general hospital after deliberate self poisoning; 32% repeaters; 70% female	Experimental (n=48): domicillary therapy (brief problem orientated) as often as therapist thought necessary; open telephone access to general hospital service. Control (n=48): outpatient therapy once a week in outpatient clinic in general hospital	96	12 months	3
Allard et al (Canada, 1992) ³⁹	Patients seen in A&E department for suicide attempt; 50% repeaters; 55% female	Experimental (n=76): intensive intervention—schedule of visits was arranged including at least one home visit; therapy provided when needed; reminders (telephone or written) and home visits made if appointments missed. Control (n=74): treatment by another staff team in the same hospital	150 (24)	12 months	3
Van Heeringen et al (Belgium, 1995) ⁴⁰	Patients \ge 15 years treated in A&E department after suicide attempt; 30% repeaters; 43% female	Experimental (n=258): special care—home visits by nurse to patients who did not keep outpatient appointments, reasons for not attending discussed and patient encouraged to attend. Control (n=258): outpatient appointments only; non-compliant patients not visited	516 (125)	12 months	3
Van der Sande et al (Netherlands, 1997) ⁴¹	Patients ≥16 years (mean 36.3) admitted to hospital after suicide attempt; 73% repeaters; 66% female	Experimental (n=140): brief psychiatric unit admission, encouraging patients to contact unit on discharge; outpatient therapy plus 24 hour emergency access to unit. Control (n=134): usual care—25% admitted to hospital, 65% outpatient referral	274	12 months	3
Emergency card <i>v</i> stan	dard aftercare				
Morgan et al (UK, 1993) ²¹	Mean age 30 years; patients admitted after first episode of deliberate self harm; % female not given	Experimental (n=101): standard care plus green card (emergency card indicating that doctor was available and how to contact them). Control (n=111): standard care—for example, referral back to primary healthcare team, psychiatric inpatient admission	112	12 months	3
Cotgrove et al (UK, 1995) ²²	Patients aged 12.2-16.7 years (mean 14.9) admitted after deliberate self harm; % repeaters not given; 85% female	Experimental (n=47): standard care plus green card (emergency card)—green card acted as passport to readmission into paediatric ward in local hospital. Control (n=58): standard follow up treatment from clinic or child psychiatry department	105	12 months	1
Dialectical behaviour t	herapy <i>v</i> standard aftercare				
Linehan et al (USA, 1991) ²⁷	Patients aged 18-45 years who had self harmed within 8 weeks before entering study; all female; all multiple repeaters of self harm	Experimental (n=32): dialectical behaviour therapy (individual and group work) for 1 year; telephone access to therapist. Control (n=31): treatment as usual; 73% individual psychotherapy	63 (24)	12 months	3
Inpatient behaviour the	erapy v inpatient insight orientated therapy				
Liberman and Eckman (USA, 1981) ⁴²	Patients (mean (range) age 29.7 (18-47) years) all repeaters; patients referred by psychiatric emergency service or hospital A&E department after deliberate self harm; 67% female	Experimental (n=12): inpatient treatment with behaviour therapy. Control (n=12): inpatient treatment with insight orientated therapy; both groups received individual and group therapy plus aftercare at community mental health centre or with private therapist	24	24 months	2
Same therapist (contin	uity of care) v different therapist (change of care)				
Torhorst et al (Germany, 1987) ²⁸	Patients referred to toxological department of Technical University Munich after deliberate self poisoning; 48% repeaters; 62% female	Experimental (n=68): continuity of care—therapy with same therapist who assessed patient in hospital after attempt. Control (n=73): change of care—therapy with different therapist than seen at hospital assessment	141 (8)	12 months	2
General hospital admis	ssion <i>v</i> discharge				
(UK, 1990) ⁴³	Patients ≥16 years (mean 30.3) admitted to A&E department for deliberate self harm; 36% repeaters; 63% female	Experimental (n=38): general hospital admission. Control (n=39): discharge from hospital; on discharge both groups advised to contact GP if they needed further help	77	16 weeks	3
Hupenthixol v placebo	Datiente aged 10 60 voors (maan 25 2) ada-titud	Eventimental (n. 19): 20 ma intromusquiar flumenthivel deserve for	27 (7)	6 months	n
1979) ²⁶	after suicidal act; all repeaters; 70% female	6 months. Control (n=19): placebo for 6 months	37 (7)	6 months	3
Hirsch et al (UK, 1982) ¹⁶ B Draper	Patients aged 16-65 years admitted after deliberate	Experimental (n=76): antidepressants—either 30-60 mg mianserin for 6 weeks or 75-150 mg nomifensing for 6 weeks, Control (n=28):	114	12 weeks	3
S Hirsch (personal communication)	given	placebo for 6 weeks	20	6 months	0
(UK, 1983) ⁴⁴	years) admitted to medical ward after deliberate self harm; all repeaters; 66% female	Experimental (1=17), mailsenn 30 mg for 6 months. Control (h=21): placebo	38	o montris	ئ
Torhorst et al	All patients repeaters who had deliberately self	Experimental (n=40); long term therapy—one therapy session a month	80	12	2
(Germany, 1988) ⁴⁵	poisoned; % female not given	for 12 months. Control (n=40): short term therapy—12 weekly therapy sessions for 3 months; all participants had brief crisis intervention (3 days) in hospital		months	-

*1=poorest quality; 3=best quality.

 Table 2
 Summary of outcome data on repetition of deliberate self harm

	Proportion (%) of participants who repeated behaviour during follow up		
Category, trial	Experimental	Control	
Problem solving therapy v standard aftercare			
Gibbons et al (1978) ³⁰	27/200 (13.5)	29/200 (14.5)	
Hawton et al (1987) ³¹	3/41 (7.3) (1*)	6/39 (15.4) (0*)	
Salkovskis et al (1990) ³²	3/12 (25.0)	4/8 (50.0)	
McLeavey et al (1994) ³³	2/19 (10.5)	5/20 (25.0)	
Intensive care plus outreach v standard care			
Chowdhury et al (1973) ³⁶	17/71 (23.9)	19/84 (22.6)	
Welu (1977) ³⁷	3/62 (4.8)	9/57 (15.8)	
Hawton et al (1981) ³⁸	5/48 (10.4) (0*)	7/48 (14.6) (0*)	
Allard et al (1992) ³⁹	22/63 (34.9) (3*)	19/63 (30.2) (1*)	
Van Heeringen et al (1995) ⁴⁰	21/196 (10.7) (6*)	34/195 (17.4) (7*)	
Van der Sande et al (1997) ⁴¹	24/140 (17.1)	20/134 (14.9)	
Emergency card v standard aftercare			
Morgan et al (1993) ²¹	5/101 (5.0) (0*)	12/111 (10.8) (0*)	
Cotgrove et al (1995) ²²	3/47 (6.4)	7/58 (12.1)	
Dialectical behaviour therapy v standard afte	rcare		
Linehan et al (1991) ²⁷ †	5/19 (26.3)	12/20 (60.0)	
Inpatient behaviour therapy v inpatient insigh	t orientated therapy		
Liberman and Eckman (1981) ⁴²	2/12 (16.7)	3/12 (25.0)	
Same therapist (continuity of care) v differen	t therapist (change of care)		
Torhorst et al (1987) ²⁸	12/68 (17.6)	4/73 (5.5)	
General hospital admission v discharge			
Waterhouse and Platt (1990) ⁴³	3/38 (7.9)	4/39 (10.3)	
Flupenthixol v placebo			
Montgomery et al (1979) ²⁶	3/14 (21.4)	12/16 (75.0)	
Antidepressants v placebo			
Hirsch et al (1982), ¹⁶ R Draper, S Hirsch (personal communication)	16/76 (21.1) (0*)	5/38 (13.2) (0*)	
Montgomery et al (1983) ⁴⁴	8/17 (47.1)	12/21 (57.1)	
Long term therapy v short term therapy			
Torhorst et al (1988) ⁴⁵	9/40 (22.5)	9/40 (22.5)	
*Reported suicides.			

†Data supplied by authors.

was unclear in 12 trials, was indicated but no further details given in seven trials, and was fully reported in one trial.

A total of 2641 patients were randomised in the 20 trials, and outcome data regarding repetition of deliberate self harm during follow up were available for 2452. The results of the individual studies in terms of repetition of deliberate self harm during follow up are shown in table 2. Also shown is the number of suicides, when these were reported.

Summary odds ratios

The summary odds ratios for each of the treatment categories that included more than one study are shown in the figure, which also indicates the total number of patients within each category.

Problem solving therapy versus standard aftercare—All four studies reported reduced repetition of deliberate self harm in patients in the experimental groups. The summary odds ratio of 0.73 (95% confidence interval 0.45 to 1.18), however, was not significant. It should be noted that the effect size was smallest in the largest trial³⁰ in this category. The trial which involved two forms of problem solving was included here because the control treatment of brief problem solving was standard treatment by the time this study was conducted.³³ Omission of this trial, which was also the only one in this category not given the highest rating of quality of concealment of allocation, made little difference to the summary odds ratio (0.78; 0.47 to 1.29). Intensive intervention plus outreach versus standard aftercare—There was no consistent direction of effect among studies in this group (see figure). Inclusion of only those trials with the highest quality of concealment of allocation did not greatly alter the summary odds ratio (0.86; 0.60 to 1.23).

Emergency card versus standard aftercare—In both studies in this comparison there was a tendency towards less repetition of self harm in the experimental group, but the summary odds ratio 0.45 (0.19 to 1.07) was not significant. The odds ratio was similar when only the results of the trial with the highest quality rating were analysed (0.43; 0.15 to 1.27).

Dialectical behaviour therapy versus standard aftercare—In this study²⁷ there was a significantly lower rate of repetition of self harm during follow up in patients who received dialectical behaviour therapy (see table 2; 0.24; 0.06 to 0.93; number needed to treat = 3). This comparison, however, was restricted to a subgroup of randomly assigned patients which was smaller than that which entered the original trial.⁴⁶

Inpatient behaviour therapy versus inpatient insight orientated therapy—The small sample size of the single study in this comparison (see table 2) precluded meaningful conclusions from the odds ratio analysis (0.60; 0.08 to 4.45).

Same therapist versus different therapist—The repetition rate in the group of patients who received aftercare from the same person who assessed them in hospital after their initial episode of deliberate self harm was significantly higher than that of patients who had a change of clinician (see table 2; 3.70; 1.13 to 12.09). The authors reported, however, that despite randomisation there were several imbalances between the experimental and control groups, resulting in a greater prevalence of risk factors for repetition in the experimental group. It is of note that continuity of therapist resulted in 48/68 (71%) patients attending at least one outpatient treatment session compared with 34/73 (47%) patients in the control group (2.75; 1.37 to 5.52).

General hospital admission versus discharge—The odds ratio from the one study in this category (see table 2) did not indicate a beneficial effect of general hospital admission after deliberate self harm (0.75; 0.16 to 3.60). Only 15% of patients referred, however, were eligible for inclusion in the study as only those attempters at low risk and without immediate medical or psychiatric needs could be considered for discharge without treatment. The follow up period was relatively short.

Flupenthixol versus placebo—There was a significant reduction in repetition of deliberate self harm in patients receiving flupenthixol (see table 2; 0.09; 0.02 to 0.50; number needed to treat = 2). The trial was relatively small and all the patients were repeaters.

Antidepressants versus placebo—The summary odds ratio for the two studies in this category (see figure) indicated no apparent benefit regarding repetition of deliberate self harm for patients treated with mianserin or nomifensine compared with placebo (1.19; 0.53 to 2.67).

Long term therapy versus short term therapy—There was no indication that long term therapy was more effective in terms of preventing repetition than short term therapy for patients with a history of self harm (see table 2; 1.0; 0.35 to 2.86).

Discussion

The results of this systematic review indicate that currently there is insufficient evidence on which to make firm recommendations about the most effective forms of treatment for patients who have recently deliberately harmed themselves. This is a serious situation given the size of the problem of deliberate self harm throughout the world^{10–13} and the importance of dealing with the problem to prevent suicide.³

In nearly all trials the subjects were recruited after attendance at a general hospital because of deliberate self harm. Some trials included only patients who had poisoned themselves, who constitute most of the patients who deliberately harm themselves¹⁰; others included patients who had poisoned themselves and those who had injured themselves, whereas some did not specify the method of self harm. Most of the studies focused on patients who could be treated as outpatients. Patients who, for example, required psychiatric hospital inpatient care because of severe mental illness or serious risk of suicide, or both, were excluded, but these comprise the minority of patients who harm themselves and who present to general hospitals.¹⁰ The studies examined are therefore of relevance to a large proportion of patients who deliberately harm themselves who will be treated in the community. Most patients in the studies had a history of episodes of self harm, and in seven trials the whole sample consisted of such patients.^{26 27 32 36 42 44} Only one study included only patients with no history of self harm.²¹ In view of the considerable problem of deliberate self harm in adolescents in many countries^{11 47} it is surprising that only one trial focused on this specific clinical population.²²

Shortcomings of trials

The comparison intervention for most of the studies of psychosocial intervention was standard care. In some studies details of this care were not provided, particularly in terms of treatment content. Future studies in which standard care is included should define precisely the nature of the treatment patients received. The dependent variable studied in this review—namely, repetition of self harm—was not consistently defined and measured in a standard way across all studies. In most studies repetition was based on hospital referral for further deliberate self harm, whereas in some studies interviews with patients and other informants also identified episodes of self harm which did not result in hospital referral.

The main problem with nearly all trials in this study is that they included far too few subjects to have the statistical power to detect clinically meaningful differences in rates of repetition of deliberate self harm between experimental and control treatments, if such differences existed. The number needed is a function of both the expected rate of repetition (that is, that in the control group) and the size of the difference. If the predicted rate were 10% in the experimental group versus 15% in the control, with α set at 0.05 and β set at 0.2, 1560 subjects would be required in each treatment group, whereas if the rates were 20% and 30%, 293 subjects would be required in each group.48 Even when the results from similar trials were synthesised with meta-analytical techniques there were insufficient numbers of patients to detect such differences. The



Summary odds ratios for repetition of deliberate self harm during follow up in treatment categories for which there was more than one trial

only significant findings have come from smaller studies, which may reflect publication bias.⁴⁹

Further research indicated

Promising results were found for problem solving therapy, which is a brief and reasonably easily taught form of treatment.⁵⁰ Although a clinically insignificant difference was found in the biggest trial,³⁰ a larger trial of this treatment is indicated. There were also trends favouring provision of an emergency access card in addition to standard aftercare but again a larger trial is required, including specific attention to what part the card might play as only a small minority of patients actually used the facility provided by possession of the card.^{21 22}

A larger replicative trial is required of the promising single study of dialectical behaviour therapy (which is similar to cognitive behaviour therapy) in women with borderline personality disorder who have a history of multiple episodes of deliberate self harm.²⁷ As the intervention is intensive,⁵¹ development and evaluation of a shorter form of this treatment more suited to provision of general psychiatric services and investigation of its efficacy in men are needed.

The positive result of depot neuroleptic medication in a single small study of patients who repeatedly harm themselves²⁶ suggests that this treatment should be subjected to further evaluation in a larger study,

Key messages

- A systematic review of the effectiveness of psychosocial and drug treatments of patients who deliberately harm themselves identified 20 randomised controlled trials in which repetition of self harm was reported as an outcome
- Promising results were found for problem solving therapy, provision of a card to allow patients to make emergency contact with services, depot flupenthixol for recurrent self harm, and long term psychological therapy for female patients with borderline personality disorder and recurrent self harm
- Assertive outreach can help to keep patients in treatment
- Nearly all the trials included too few patients to detect clinically significant differences in repetition of self harm, and even synthesis of results by meta-analysis did not have the power to detect such differences
- There is an urgent need for large trials of promising therapies for this substantial clinical population

although reluctance of patients to accept depot medication, side effects, and other practical and ethical implications, may limit its applicability. There was little indication that intensive intervention plus outreach was effective. In one relatively large study in this group, which evaluated community follow up of patients who did not attend outpatient appointments,⁴⁰ there was a significant increase in outpatient attendance from 42.5% before the home visit (39.8% in the control group) to 51.2% after the visit (odds ratio 1.58; 1.15 to 2.33) and a near significant difference in repetition of deliberate self harm of 10.7% compared with 17.4% (0.57; 0.32 to 1.02). Home treatment was also found substantially to increase the rate of take up of treatment in another trial in this review.38 Assertive outreach for poorly compliant patients may, therefore, be a necessary component in maximising the delivery of any treatment that is shown to be effective.

There was no evidence that antidepressants were generally effective in preventing repetition of behaviour in patients who deliberately harm themselves. One of the drugs investigated, however, is no longer available (nomifensine) and the other (mianserin) is now little used. This review does not give any indication of whether other antidepressants could be of benefit.

Repetition of deliberate self harm has been the sole outcome variable investigated in this review. It will be important to determine whether there is evidence of benefits with regard to other outcomes (for example, depression or problem resolution), although the data for these factors, when reported, were often inadequate for meta-analysis. The groups of patients were often heterogeneous in term of sex, age, and presenting problems. Further work in this specialty should examine the efficacy of interventions according to such factors.

Conclusion

At present, evidence is lacking to indicate the most effective forms of treatment for patients who deliberately harm themselves. This is a serious situation given the size of the population at risk and the risks of subsequent self harm, including suicide. Large trials are required of the interventions shown in small trials to be of possible benefit. There is also a need for development of further treatment approaches informed by current knowledge

about the psychosocial and biological characteristics of these patients and the socioeconomic and sociocultural context of the behaviour.

This study was carried out under the auspices of the Depression, Anxiety and Neurosis Review Group of the Cochrane Collaboration. Updated versions of the review will appear in the Cochrane Controlled Trials Register in the future. We thank Rachel Churchill and Drs Douglas Altman, Clive Adams, John Geddes, and Henry McQuay for advice. We also thank the several authors who supplied us with unpublished information regarding their trials.

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An epidemiological needs assessment of carotid endarterectomy in an English health region. Is the need being met?

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Abstract

Objective: To compare the level of provision of carotid endarterectomy (an intervention of proved efficacy for prevention of stroke in patients with symptomatic high grade carotid artery stenosis) with estimates of need.

Design: Comparison of regional, district, and age-sex specific operation rates derived from hospital episode statistics with estimates of need based on demographic and epidemiological data; interviews with regional vascular surgeons and a joint provider-purchaser workshop to discuss implications. Setting: Former Wessex Regional Health Authority, 1991-2 to 1995-6.

Subjects: All residents covered by Wessex region treated for carotid artery reconstruction.

Main outcome measures: Regional, district, and age-sex operation rates as three year average 1993-6 (use) compared with respective estimates of need for carotid endarterectomy among those who presented with symptomatic carotid disease-transient ischaemic attack or minor stroke.

Results: The operation rate more than doubled between 1991-2 and 1995-6, from 35 to 89 per million population, compared with an estimated level of need in the region's general population of 153 per million population (transient ischaemic attack 77, minor stroke 76). The ratio of use to need was 0.47

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(95% confidence interval 0.4 to 0.54); district ratios were 0.28 (0.19-0.38) to 0.81 (0.62 to 1.06). The annual use:need ratio rose over the three years 1993-6 from 0.38 to 0.59. Use:need ratios were lower in elderly and female patients. Providers were keen to develop guidelines for referral and to increase access to diagnostic facilities; purchasers were more reluctant, given the limited impact of this intervention on the incidence of stroke and the relatively high cost of the operation.

Conclusion: Although treatment rates increased in Wessex there is still unmet need. Further research is needed to determine the referral pathways of patients with symptomatic carotid disease for diagnosis and operation and to evaluate strategies to improve access to diagnostic facilities.

Introduction

Stroke is the third commonest cause of death in the United Kingdom and the commonest cause of physical disability.¹ Each year 64 000 deaths (12% of all deaths) are attributed to stroke.1 Treating and managing patients who have had a stroke is estimated to consume about 4.0% of NHS resources.2

Carotid endarterectomy (the removal of atherosclerotic lesions from the inner wall of the common carotid artery extending up to the internal carotid artery) has a small but significant part to play in

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