EBMH Notebook

Generalised anxiety disorder

The *EBMH* Notebook summarises key messages about generalised anxiety disorder, sourced from: *Clin Evid Concise* 2004 (in press); www.clinicalevidence.com. For this review, *Clinical Evidence Concise* searched and appraised material published until June 2003.

DEFINITION

Generalised anxiety disorder (GAD) is defined as excessive worry and tension about every day events and problems on most days, for at least six months, to the point where the person experiences distress or has marked difficulty in performing day to day tasks.¹ It may be characterised by the following symptoms and signs: increased motor tension (fatigability, trembling, restlessness, and muscle tension); autonomic hyperactivity (shortness of breath, rapid heart rate, dry mouth, cold hands, and dizziness); and increased vigilance and scanning (feeling keyed up, increased startling, and impaired concentration), but not panic attacks.¹ One non-systematic review of epidemiological and clinical studies found marked reduction of quality of life and psychosocial functioning in people with anxiety disorders (including GAD).² It also found that people with GAD have low overall life satisfaction and some impairment in ability to fulfil roles, social tasks, or both.²

INCIDENCE/PREVALENCE

One overview of observational studies published in English found that the prevalence of GAD among adults in the community is 1.5-3.0%.3 It found that 3-5% of adults have had GAD in the past year and 4-7% have had GAD during their life. The US National Comorbidity Survey found that over 90% of people diagnosed with GAD had a comorbid diagnosis, including dysthymia (22%), depression (39-69%), somatisation, other anxiety disorders, bipolar disorder, or substance abuse.4 The Harvard Brown Anxiety Research Program also found that only 30/180 (17%) people had GAD alone.5 Subgroup analysis suggested that 46/122 (38%) of people with GAD had comorbid personality disorder.6 A systematic review of the comorbidity of eating disorders and anxiety disorders (search date 2001, two observational studies, 55 people) found a lifetime prevalence of GAD among people with anorexia nervosa of 24% in one study and 31% in the other.7 The lifetime prevalence of GAD in the control group of one of the studies (44 people) was 2%. The reliability of the measures used to diagnose GAD in epidemiological studies is unsatisfactory.89 One US study, with explicit diagnostic criteria (DSM-III-R), estimated that 5% of people will develop GAD at some time during their life.9 A recent cohort study of people with depressive and anxiety disorders found that 49% of people initially diagnosed with GAD retained this diagnosis over two years.¹⁰ The incidence of GAD in men is only half the incidence in women¹¹ and is lower in older people.¹² A non-systematic review (20 observational studies in younger and older adults) suggested that autonomic arousal to stressful tasks is decreased in older people, and that older people become accustomed to stressful tasks more quickly than younger people.13

AETIOLOGY/RISK FACTORS

Generalised anxiety disorder is believed to be associated with an increase in the number of minor stressors, independent of demographic factors,¹⁴¹⁵ but this finding is also common in people with other diagnoses in the clinical population.¹⁰ One non-systematic review (five case control studies) of psychological sequelae to civilian trauma found that rates of GAD reported in four of the five studies were significantly increased compared with a control population (rate ratio 3.3, 95% CI 2.0 to 5.5).¹⁶ One systematic review (search date 1997) of cross sectional studies found that bullying (or peer victimisation) was associated with a significant increase in the incidence of GAD (effect size 0.21).¹⁷ Genetic factors are also implicated. One systematic review (search date not reported, two family studies, 45 index cases, 225 first degree relatives) found a significant association between GAD in the index cases and in their first degree relatives (OR 6.1, 95% CI 2.5 to 14.9).¹⁸ The review also identified three twin studies (13 305 people), which estimated that 32% (95% CI 24% to 39%) of the variance to liability to GAD was explained by genetic factors.

PROGNOSIS

One systematic review found that 25% of adults with GAD will be in full remission after two years, and 38% will have a remission after five years.³ The Harvard-Brown anxiety research program reported five year follow up of 167 people with GAD.¹⁹ In this period, the weighed probability for full remission was 38% and for at least partial remission was 47%: the probability of relapse from full remission was 27% and relapse from partial remission was 39%.

WHAT ARE THE EFFECTS OF TREATMENTS? Likely to be beneficial Buspirone

Randomised controlled trials (RCTs) have found that buspirone improves symptoms compared with placebo over 4–9 weeks. RCTs found no significant difference in symptoms over 6–8 weeks between buspirone and antidepressants, diazepam, or hydroxyzine, but the studies may have lacked power to detect clinically important differences among treatments.

Certain antidepressants (imipramine, opipramol, paroxetine, and venlafaxine)

Randomised controlled trials have found that antidepressants (imipramine, opipramol, paroxetine, and venlafaxine) improve symptoms over 4–28 weeks compared with placebo. RCTs found no significant difference among these antidepressants or between antidepressants and benzodiazepines or buspirone. RCTs and observational studies have found that antidepressants are associated with sedation, dizziness, nausea, falls, and sexual dysfunction.

Cognitive behavioural therapy

Two systematic reviews and two subsequent RCTs have found that cognitive behavioural therapy (using a combination of interventions, such as exposure, relaxation, and cognitive restructuring) improves anxiety and depression over 4-12 weeks compared with waiting list control, anxiety management alone, relaxation alone, or non-directive psychotherapy. Three subsequent RCTs, two in people aged ≥ 60 years, found no significant difference in symptoms at 13 weeks, six months, or 24 months between cognitive therapy and applied relaxation.

Hydroxyzine

Three RCTs comparing hydroxyzine versus placebo found different results. Two RCTs found that, compared with placebo, hydroxyzine improved symptoms of anxiety at four or 12 weeks, but a third RCT found no significant difference in the proportion of people with improved symptoms of anxiety at five weeks. One of the RCTs found that hydroxyzine increased somnolence and headaches compared with placebo. One RCT found no significant difference between hydroxyzine and bromazepam in the proportion of people who responded after six weeks. Another RCT found no significant difference between hydroxyzine and buspirone in the proportion of people who responded after four weeks.

Trade off between benefits and harms

Benzodiazepines

One systematic review and one subsequent RCT found that benzodiazepines reduced symptoms over 2-9 weeks compared with placebo. RCTs found no significant difference in symptoms over 3-8 weeks between alprazolam and bromazepam or mexazolam, or between benzodiazepines and buspirone, hydroxyzine, abecarnil, or antidepressants. RCTs and observational studies found that benzodiazepines increased the risk of dependence, sedation, industrial accidents, and road traffic accidents and that, if used in late pregnancy or while breast feeding, benzodiazepines may cause adverse effects in neonates. RCTs found no significant difference in symptoms over 3-8 weeks between alprazolam and bromazepam or mexazolam, or between benzodiazepines and buspirone, hydroxyzine, abecarnil, or antidepressants. One systematic review of poor quality RCTs provided insufficient evidence to assess long term treatment with benzodiazepines.

Kava

One systematic review in people with anxiety disorders, including generalised anxiety disorder, found that kava reduced symptoms of anxiety over 1-24 weeks compared with placebo. It is unclear whether results of the review are generalisable to people with generalised anxiety disorder. Observational evidence suggests that kava may be associated with hepatotoxicity.

Trifluoperazine

One large RCT found that trifluoperazine reduced anxiety after four weeks compared with placebo, but caused more drowsiness, extrapyramidal reactions, and other movement disorders.

Unknown effectiveness

Abecarnil

One RCT found limited evidence that low dose abecarnil improved symptoms compared with placebo. Another RCT found no significant difference in symptoms at six weeks between abecarnil and placebo or diazepam. Both RCTs found that abecarnil increased drowsiness compared with placebo.

Applied relaxation

We found no RCTs comparing applied relaxation versus placebo or no treatment. Three RCTs found no significant difference in symptoms at 13 weeks, six months, or 24 months between applied relaxation and cognitive behavioural therapy.

β blockers

We found no RCTs on the effects of β blockers in people with generalised anxiety disorder.

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REFERENCES

- American Psychiatric Association. Diagnostic and statistical manual of mental disorders, 4th ed. Washington, DC: American Psychiatric Association, 1991
- Mendlowicz MV, Stein MB. Quality of life in individuals with anxiety disorders. Am J Psychiatry 2000;157:669–82.
- Kessler RD Wittchen HU. Patterns and correlates of generalized anxiety disorder in community samples. J Clin Psychiatry 2002;63:4–10.
- Stein D. Comorbidity in generalised anxiety disorder: impact and implications. J Clin Psychiatry 2001;62:29–34.
- 5 Goldenberg IM, White K, Yonkers K, et al. The infrequency of "pure culture" diagnoses among the anxiety disorders. J Clin Psych 1996;57:528-33.
- 6 Dyck IR, Phillips KA, Warshaw MG, et al. Patterns of personality pathology in potients with generalized anxiety disorder, panic disorder with and without agoraphobia, and social phobia. *J Personal Disord* 2001;**15**:60–71.
- Godart NT, Flament MF, Perdereau F, et al. Comorbidity between eating disorders and anxiety disorders: a review. Int J Eat Disord 2002;32:253-70.
- usuruers unu anxiety aisoraers: a review. Int J Eat Disord 2002;32:253–70.
 Search date 2001; primary source Medline.
 Judd LL, Kessler RC, Paulus MP, et al. Comorbidity as a fundamental feature of generalised anxiety disorders: results from the National Comorbidity Study (NCS). Acta Psychiatry Scand 1998;98:6–11.
 Andrew G, Patria LC, Comorbidity Study (NCS).
- Andrews G, Peters L, Guzman AM, et al. A comparison of two structured diagnostic interviews: CIDI and SCAN. Aust N Z J Psychiatry 1995:29:124-32.
- 10 Kessler RC, McGonagle KA, Zhao S, et al. Lifetime and 12-month prevalence of DSM-III-R psychiatric disorders in the United States: results from the national comorbidity survey. Arch Gen Psychiatry 1994;**51**:8–19. Seivewright N, Tyrer P, Ferguson B, *et al.* Longitudinal study of the influence of
- 11 life events and personality status on diagnostic change in three neurotic disorders. *Depression Anx* 2000;11:105–13.
- 12 Pigott T. Gender differences in the epidemiology and treatment of anxiety disorders. J Clin Psychiatry 1999;60:4-15.
- 13 Jorm AF. Does old age reduce the risk of anxiety and depression? A review of epidemiological studies across the adult life span. *Psychol Med* 2000;**30**:11–22.
- 14 Lau AW, Edelstein BA, Larkin KT. Psychophysiological arousal in older adults: a critical review. *Clin Psychol Rev* 2001;**21**:609–30.
- 15 Brantley PJ, Mehan DJ Jr, Ames SC, et al. Minor stressors and generalised anxiety disorders among low income patients attending primary care clinics. J Nerv Ment Dis 1999;187:435–40.
- 16 Brown ES, Fulton MK, Wilkeson A, et al. The psychiatric sequelae of civilian trauma. Comp Psychiatry 2000;41:19-23.
- 17 Hawker DSJ, Boulton MJ. Twenty years' research on peer victimisation and Hawker DSJ, bouton MJ. Iwenty years research on peer vicrimisation and psychosocial maladjustment: a meta-analytic review of cross-sectional studies. *J Child Psychol Psychiatr* 200;**4**1:441–5. Search date 1997; primary sources Psychlit, Social Science Citation Index, OCLC Firstsearch, hand searches of relevant journals, bibliographies, reviews, reference lists of relevant articles, and book chapters, and personal contact with authors.
 Hettema JM, Neale MC, Kendler KS. A review and meta-analysis of the representation of the product of the produ
- genetic enjidemiology of anxiety disorders. *Am J Psychiatry* 2001;**158**:1568–78. Search date not reported; primary source Medline.
- Yonkers KA, Dyck IR, Warshaw M, et al. Factors predicting the clinical course of generalised anxiety disorder. Br J Psychiatry 2000;176:544–9. 19

³⁴EBMH Notebook

Meta-analyses and megatrials: neither is the infallible, universal standard

Now owadays, most would agree that we need evidence from randomised control trials (RCTs) to evaluate the effectiveness of a health intervention. It used to be that we did not have enough RCTs in mental health; the irony today is that at times it seems we have too many of them, especially when they draw conflicting conclusions.

A natural solution is to seek "stronger" evidence. Metaanalysis might provide that evidence but, alas, meta-analyses sometimes do not agree among themselves either.¹ Another possible solution is a bigger and better trial, a megatrial (also known as the large, simple trial). Unfortunately megatrials and meta-analyses do not always agree either: one group has claimed that—taking megatrials as the gold standardmeta-analyses drew wrong conclusions 35% of the time²; another group estimated the degree of disagreement to be between 10% and 23%.³ Megatrials sometimes do not agree with each other either, and discrepancies among megatrials are just as large as those between meta-analyses and megatrials.⁴

These discrepancies reinforce a conclusion that the days of dogmatic advocacy of the methodological hierarchy of evidence are over.⁵

Here, I will take three examples to illustrate that we will always need "good common sense", coupled with content expertise and an understanding of methodology, to weigh the available evidence relevant to a mental health problem.

	Definition	Strengths	Weaknesses
Applies to both meta-analyses and megatrials		 Can ascertain moderate but worthwhile treatment benefits (small effect on major outcomes, such as death or disablement). This characteristic is important because nowadays we can seldom expect a large treatment gain by breakthrough technology. Samples and results are heterogeneous not only in meta-analysis but also in one megatrial. Ironically, however, despite cries for "tailor made" medicine, it is usually the overall results of the meta-analysis or megatrial and not post hoc subgroup results that are more generalisable. 	they are not big enough to tell us much about
Meta-analysis	Combination of data from several independently performed single or multicentre trials with the purpose of assessing effects on endpoints for which the individual trials are usually non-informative due to lack of statistical power.	 Provides the most reliable treatment estimate in the absence of a definitive trial. Although quite labour intensive, less expensive to conduct than a megatrial. Can be seen as exploratory and hypothesis generating for the planning of a definitive large trial. 	 Biases and flaws of individual trials are incorporate and new sources of bias may be incorporated (publication bias, prematurely terminated studies, small studies) In addition to publication bias of trials, publication bias of outcomes is huge (not all identified trials report on the same primary outcomes in the same way). Harms are even less often uniformly assessed thar primary endpoints, so that harm assessment is less precise than benefit assessment. Different statistical techniques can result in conflictin results, based on the same data.
Megatrial	Very large randomised controlled trials, usually recruiting thousands of subjects and usually multicentred. Recruitment criteria are very broad, protocols are maximally simplified, and endpoints are unambiguous, such as death. Also often referred to as a 'large, simple trial'. Typical examples are seen in cardiovascular medicine.	 Can provide accurate estimates of pragmatic effectiveness and side effects in the real world. Is designed from the beginning and conducted throughout to give precise measurement of treatment effects and side effects in question. 	 Large sample size required, and hence very expensive—for example, 100 million US dollars for GUSTO-I. Simplification of recruitment and data collection increases the risks of protocol deviation, poor data quality, misclassification, and non-trial use of trial treatments, all of which create a bias towards the n hypothesis. The control condition is sometimes defined as "treatment as usual" but this is often not standardised. Megatrials can be properly designed only after ma smaller trials have clarified the characteristics of the intervention in guestion.

Well conducted systematic reviews including megatrials usually offer the best guide to overall treatment effect. For example, in the case of risperidone versus typical antipsychotics for schizophrenia, a very large multinational, multicentre RCT (n = 1362) found no statistically significant difference between these two drugs (RR of no response = 0.94: 95% CI 0.79 to 1.11).⁶ A subsequent Cochrane review that included an additional 1006 subjects did show, in contrast, a significant and important random effects RR of 0.84 (95% CI 0.76 to 0.92) in favour of risperidone. There was no indication of heterogeneity across trials (p = 0.63).⁷ It appears that one of the largest trials to date in mental health6 was still underpowered to detect a small yet important difference.

When available studies for meta-analysis are limited in number, sample size, or quality of methodology, we are in a more difficult position. Another Cochrane review concluded that lithium therapy is an efficacious maintenance treatment for bipolar disorder.⁸ Combining three studies (total n = 412), this review found a statistically significant and clinically meaningful reduction in relapse for patients with bipolar disorder on lithium compared to placebo (random effects RR = 0.60, 95% CI 0.41 to 0.87). Heterogeneity among the included RCTs was not statistically significant (p = 0.13) but substantive $(I^2 = 51.6\%)$. Although two older studies found lithium to be superior to placebo, the most recent study failed to find a statistically significant difference between the two arms (RR = 0.71, 95% CI 0.39 to 1.31).⁹ One reasonable conclusion was that the latest study was underpowered and was in fact in concordance with previous studies. Considerable debate ensued after publication of this pivotal study and there was ongoing debate on the methodological adequacy of the older trials with lithium. The superficial interpretation of the more recent study as "negative" seemed to support claims against accepted wisdom in modern psychiatry. This clinical and scientific chagrin abated somewhat when the same group of researchers published a similarly planned maintenance RCT and found a significant reduction in relapse on lithium in comparison with placebo.10 Closer reading of their report reveals, however, that lithium reduced relapse over 12 months only at the expense of increasing dropouts due to adverse events; survival on the medication without relapse or dropout was no different on lithium or on placebo. Only 22% and 16%, respectively, of those starting on lithium or placebo remained on the same drug without relapse until the study termination up to 18 months. The value of lithium appears small at best.

When a systematic review is of inferior quality, we are in an even more difficult position. A meta-analysis of alprazolam for anxiety disorders involving 8878 randomised patients claimed to have confirmed its efficacy.¹¹ Alprazolam may indeed be better than placebo in reducing panic and associated anxiety over 8-12 weeks but we need no more than a well designed, well analysed study of 154 patients to convincingly disqualify alprazolam as drug of choice for anxiety disorders. Aprazolam alone was not as good as exposure therapy alone for the acute phase of treatment, and the addition of alprazolam to exposure therapy resulted in even worse outcomes at follow up than exposure alone.¹²

Having observed these illustrative cases and having appreciated that a thorough critical reading of a comprehensive meta-analysis is a formidable task, we in the Department of Psychiatry at Nagoya City University tend to examine meta-analysis as a navigator for sound evidence on a clinical topic. Looking at the whole map of available trials in the metaview of the Cochrane Library, we often choose to critically appraise and learn from the best—the largest, the most recent, the best known, the closest to the overall mean, whatever-trial in detail. We find that such practice often brings more insight to the bedside the next day than critically appraising the meta-analysis itself.

The strengths and weaknesses of meta-analyses and megatrials are shown in table 1. We can never arrive at infallible truth because, firstly, that is simply not the nature of scientific knowledge13 and, secondly, in clinical medicine we are dealing with complex, ever changing units of analysis that are people with illnesses.

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- Jadad AR, Cook DJ, Browman GP. A guide to interpreting discordant systematic reviews. Can Med Assoc J 1997;156:1411-16.
- 2 LeLorier J, Gregoire G, Benhaddad A, et al. Discrepancies between metaanalyses and subsequent large randomized, controlled trials. N Engl J Med 1997:337:536-42.
- Ioanidis JPA, Cappelleri JC, Lau J. Issues in comparisons between meta-analyses and large trials. JAMA 1998;**279**:1089–93.
 Furukawa TA, Streiner DL, Hori S. Discrepancies among megatrials. J Clin
- Epidemiol 2000;53:1193-9.
- 5 Glasziou P, Vandenbroucke J, Chalmers I. Assessing the quality of research. BMJ 2004;328:39-41.
- 6 Peuskens J. Risperidone in the treatment of patients with chronic schizophrenia: a multi-national, multi-centre, double-blind, parallel-group study versus haloperidol. Risperidone Study Group. Br J Psychiatry 1995;166:712-26; discussion 727-33.
- 7 Hunter RH, Joy CB, Kennedy E, et al. Risperidone versus typical antipsychotic medication for schizophrenia. Cochrane Database Syst Rev 2003;(2):CD000440.
- 8 Burgess S, Geddes J, Hawton K, et al. Lithium for maintenance treatment of mood disorders. Cochrane Database Syst Rev 2001;(3):CD003013.
- 9 Bowden CL, Calabrese JR, McElroy SL, et al. A randomized, placebo controlled 12-month trial of divalproex and lithium in treatment of outpatients with bipolar I disorder. Divalproex Maintenance Study Group. Arch Gen Psychiatry 2000;57:481-9.
- Bowden CL, Calabrese JR, Sachs G, et al. Lamictal 606 Study Group. A placebo-controlled 18-month trial of lamotrigine and lithium maintenance treatment in recently manic or hypomanic patients with bipolar I disorder. Arch Gen Psychiatry 2003;**60**:392–400.
- 11 Jonas JM, Cohon MS. A comparison of the safety and efficacy of alprazolam versus other agents in the treatment of anxiety, panic, and depression: a review of the literature. J Clin Psychiatry 1993;54(Suppl):25–45; discussion 46-8.
- 12 Marks IM, Swinson RP, Basoglu M, et al. Alprazolam and exposure alone and combined in panic disorder with agoraphobia: a controlled study in London and Toronto. Br J Psychiatry 1993;**162**:776–87.
- 13 Popper K. The Logic of Scientific Discovery. 15th edn. London: Routledge, 2002.

³⁶Other articles noted

• he journals that are reviewed and the criteria for selecting articles from these journals for inclusion in *Evidence-Based Mental Health* are set out in the purpose and procedure in each issue. These articles met inclusion criteria for *Evidence-Based Mental Health*, but were not abstracted due to lack of space.

THERAPEUTICS

Bearman SK. Evaluation of an intervention targeting both depressive and bulimic pathology: a randomized prevention trial. *Behavior Therapy* 2003;**34**:277–93.

Fitzgerald PB, Brown TL, Marston NA, *et al.* Transcranial magnetic stimulation in the treatment of depression: a double-blind, placebo-controlled trial. *Arch Gen Psychiatry* 2003;**60**:1002–8.

Guthrie E, Kapur N, Mackway-Jones K, *et al.* Predictors of outcome following brief psychodynamic-interpersonal therapy for deliberate self-poisoning. *Aust N Z J Psychiatry* 2003;**37**:532–6.

Harvey PD, Napolitano JA, Mao L, *et al*. Comparative effects of risperidone and olanzapine on cognition in elderly patients with schizophrenia or schizoaffective disorder. *Int J Geriatr Psychiatry* 2003;**18**:820–9.

Krishnan KR, Charles HC, Doraiswamy PM, *et al.* Randomized, placebo-controlled trial of the effects of donepezil on neuronal markers and hippocampal volumes in Alzheimer's disease. *Am J Psychiatry* 2003;**160**: 2003–11.

Lenox-Smith AJ, Reynolds A. A double-blind, randomised, placebo controlled study of venlafaxine XL in patients with generalised anxiety disorder in primary care. *Br J Gen Pract* 2003;**53**:772–7.

O'Donnell C, Donohoe G, Sharkey L, *et al*. Compliance therapy: a randomised controlled trial in schizophrenia. *BMJ* 2003;**327**:834.

Rapaport MH, Schneider LS, Dunner DL, *et al*. Efficacy of controlled-release paroxetine in the treatment of late-life depression. *J Clin Psychiatry* 2003;**64**:1065–74.

Rentz TO, Powers MB, Smits JA, *et al*. Active-imaginal exposure: examination of a new behavioral treatment for cynophobia (dog phobia). *Behav Res Ther* 2003;**41**: 1337–53.

Rosenheck RA, Resnick SG, Morrissey JP. Closing service system gaps for homeless clients with a dual diagnosis: integrated teams and interagency cooperation. *J Ment Health Policy Econ* 2003;**6**:77–87.

Snowden M, Sato K, Roy-Byrne P. Assessment and treatment of nursing home residents with depression or behavioral symptoms associated with dementia: a review of the literature. *J Am Geriatr Soc* 2003;**51**:1305–17.

van der Wurff FB, Stek ML, Hoogendijk WJ, *et al.* The efficacy and safety of ECT in depressed older adults: a literature review. *Int J Geriatr Psychiatry* 2003;**18**:894–904.

AETIOLOGY

Fergusson DM. Major depression and cigarette smoking: results of a 21-year longitudinal study. *Psychol Med* 2003;**33**:1357–68.

Gilman SE. Socio-economic status, family disruption and residential stability in childhood: relation to onset, recurrence and remission of major depression. *Psychol Med* 2003;**33**:1341–56.

Goodwin FK, Fireman B, Simon GE, *et al.* Suicide risk in bipolar disorder during treatment with lithium and dival-proex. *JAMA* 2003;**290**:1467–73.

Ruschena D, Mullen PE, Palmer S, *et al*. Choking deaths: the role of antipsychotic medication. *Br J Psychiatry* 2003;**183**:446–50.

DIAGNOSIS

Lipton RB, Katz MJ, Kuslansky G, *et al*. Screening for dementia by telephone using the memory impairment screen. *J Am Geriatr Soc* 2003;**51**:1382–90.

PROGNOSIS

Bradshaw Z, Slade P. The effects of induced abortion on emotional experiences and relationships: A critical review of the literature. *Clin Psychol Rev* 2003;**23**:929–58.