Guided self-instructions for people with chronic fatigue syndrome: randomised controlled trial

Hans Knoop, Jos W. M. van der Meer and Gijs Bleijenberg

Summary
A minimal intervention, based on cognitive–behavioural therapy for chronic fatigue syndrome and consisting of self-instructions combined with email contact, was tested in a randomised controlled trial (ISRCTN22793439). A total of 171 patients participated in the trial. 85 were allocated to the intervention condition and 86 to the waiting-list condition. All patients met the Centers for Disease Control and Prevention criteria for chronic fatigue syndrome. An intention-to-treat analysis showed a significant decrease in fatigue and disability after self-instruction. The level of disability was negatively correlated with treatment outcome. Guided self-instructions are an effective treatment for patients with relatively less severe chronic fatigue syndrome.

Declaration of interest
None.

Method

Patients referred for CBT to our tertiary care facility were eligible to enter the study if they were ≥18 years old; spoke and read Dutch; met the 1994 US Center for Disease Control and Prevention criteria for chronic fatigue syndrome; were not engaged in a legal procedure concerning disability-related financial benefits; scored ≥35 on the Checklist Individual Strength (CIS); fatigue severity sub-scale; had a total score of >700 on the Sickness Impact Profile–8 (SIP8); and had given written informed consent. The local ethics committee approved the study.

After baseline assessment performed by research assistants, patients were offered CBT. If they agreed, they were placed on a waiting list for a period of 6–12 months depending on available treatment capacity. Patients were informed about the study and if they gave informed consent, were randomly assigned to either the guided self-instructions or the waiting-list condition. Allocation to group was carried out by a therapist using cards containing information about chronic fatigue syndrome and weekly assignments. The programme took at least 16 weeks, but often more if patients formulated long-term goals such as returning to work. Patients were asked to email (or telephone if they did not have email) at least once every 2 weeks to report their progress. A cognitive–behavioural therapist, trained in regular CBT for chronic fatigue syndrome, responded to this email or call. If patients did not respond every 2 weeks, a reminder was sent by email or patients were telephoned.

The CIS sub-scale ‘fatigue severity’ was used to measure the level of fatigue over the past 2 weeks. Scores ranged from 8 (no fatigue) to 56 (severely fatigued). The weighted total score on eight sub-scales of the SIP8 (SIP8 total score) was used to assess functional disability in all domains of functioning. Physical disabilities were measured with the physical functioning sub-scale of the 36-item Short Form Health Survey (SF–36). Scores ranged from 0 (maximum physical limitations) to 100 (ability to do vigorous activity).

Clinically significant improvement was defined as a reliable change index >1.96 and a score of <35 on the CIS sub-scale fatigue severity at second assessment. This score is within two standard deviations of the mean for healthy adults. A score of <35 would reflect a significant reduction of fatigue. We assumed that in the waiting-list condition 10% of the patients would have a fatigue score of <35. Power calculation showed that 98 patients in each condition were needed to detect a difference of 15% in the proportion of patients with a fatigue score within normal limits, assuming a significance of 5%, power of 90% and a drop-out rate of 20%.

Data analyses were performed using SPSS (version 14) for Windows. Significance was assumed at P<0.05. To test whether there was a difference between the two conditions on outcome measures, ANCOVA was used with the score on the second assessment as the dependent variable, baseline score as the covariate and condition as the fixed factor. To test whether the proportion of patients with a clinically significant improvement differed between conditions, logistic regression with condition and baseline CIS fatigue as predictors was used. To test whether the treatment effect was moderated by patient characteristics, ANCOVA for CIS fatigue on the second assessment was repeated with condition (standardised; Z-score with mean=0, s.d.=1), baseline CIS fatigue (standardised), baseline SIP8 total score, condition × baseline CIS score, and condition × baseline SIP8 score as predictors. All comparisons were performed on the basis of intention to treat. For missing data, the last observation was carried forward.

Results

During the study it became clear that the number of patients lost at the second assessment was lower than expected (about 5%).
The power analysis was repeated but now using an estimated drop-out rate of 5%, indicating that a sample size of 85 in each condition sufficed. The inclusion of patients stopped when this sample size was reached (Fig.1).

Of 84 patients in the self-instructions condition, 55 (66%) emailed, 5 (6%) exclusively used the telephone and 10 (12%) did both. Fourteen patients (16%) had no contact with a therapist: 1 patient completed the programme by herself, the remaining 13 did not start. There was no significant difference in mean time passed in months between the two assessments for the guided self-instruction (10.5 months, s.d.=4.0) and the waiting-list condition (9.7 months, s.d.=3.6, t=3.6, d.f.=157, P=0.018). Patients from the intervention condition were significantly less fatigued (intervention mean=38.9 v. waiting-list mean=46.6), reported fewer disabilities (mean=1079 v. mean=1319), scored significantly higher on the SF–36 physical functioning sub-scale (mean=65.9 v. mean=60.2) and more often showed a clinically significant improvement in fatigue (27% v. 7%) at second assessment (online Tables DS1–3).

Of the interaction terms, only the condition × SIP8 total score interaction effect was significant (B=3.533, t=2.250, P=0.026), indicating that the treatment effect is more than halved for patients with an SIP8 score of 1 standard deviation above the mean.

### Discussion

It was already known that individual CBT is an effective treatment for chronic fatigue syndrome. The results also suggest that more severely disabled patients benefit less from the self-instructions and could perhaps be referred for face-to-face CBT instead.

As we did not use a control condition we cannot be sure that the specific elements in the minimal intervention condition were responsible for the reduction of fatigue and disabilities. However, two randomised controlled trials that compared the effect of CBT for chronic fatigue syndrome with a placebo or non-specific condition both showed CBT to have a superior effect. Furthermore, Cho et al showed that the placebo response of patients with chronic fatigue syndrome to psychological interventions is lower than in other medical conditions.

### Acknowledgements

Trial register number: ISRCTN27293439. Ethical approval was obtained from the human ethics committee of the Radboud University Nijmegen Medical Centre. We thank the patients who participated in the study and Thea Berends, Annemarie Gerritsen, Gerrie van de Heijdt, Henriette Vermeiren and Hein Voskamp for treating the patients in the guided self-instruction condition. Carel Knopp, Tiny Fasotti and Lianne Vermeer for their assistance in data collection, and Roger Borders for statistical advice.

### References


Data supplement

### Table DS1  Characteristics between groups at baseline

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Self-instructions (n=84)</th>
<th>Waiting list (n=85)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years: mean (s.d.)</td>
<td>37.6 (10.0)</td>
<td>38.5 (10.6)</td>
</tr>
<tr>
<td>Duration of complaints, months: median (min., max.)</td>
<td>72 (12, 420)</td>
<td>96 (12, 420)</td>
</tr>
<tr>
<td>Male/female</td>
<td>15/69</td>
<td>20/65</td>
</tr>
</tbody>
</table>

### Table DS2  Change in outcome measures between baseline and second assessment

<table>
<thead>
<tr>
<th>Outcome measure</th>
<th>Baseline Mean (s.d.)</th>
<th>Second assessment Mean (s.d.)</th>
<th>Baseline Mean (s.d.)</th>
<th>Second assessment Mean (s.d.)</th>
<th>Difference Mean (95% CI)</th>
<th>F(1,166)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIS fatigue severity</td>
<td>49.1 (5.2)</td>
<td>38.9 (12.1)</td>
<td>49.9 (5.6)</td>
<td>46.4 (8.7)</td>
<td>-6.7 [-9.7 to -3.6]</td>
<td>20.61</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>SIP8 total score</td>
<td>1659 (648)</td>
<td>1079 (690)</td>
<td>1515 (545)</td>
<td>1319 (619)</td>
<td>-384 [-543 to -225]</td>
<td>19.77</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>SF–36 physical functional</td>
<td>52.3 (20.4)</td>
<td>65.9 (23.2)</td>
<td>54.1 (21.1)</td>
<td>60.2 (23.7)</td>
<td>7.5 [1.8 to 13.1]</td>
<td>6.56</td>
<td>0.011</td>
</tr>
</tbody>
</table>

CIS, Checklist Individual Strength; SIP8, Sickness Impact Profile; SF–36, 36-item Short Form Health Survey.

### Table DS3  Change in CIS fatigue severity between groups

<table>
<thead>
<tr>
<th>Outcome measure</th>
<th>Self-instructions (n=84)</th>
<th>Waiting list (n=85)</th>
<th>OR (95% CI)</th>
<th>Wald</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIS fatigue severity</td>
<td>23/84 (27, 18–37)</td>
<td>6/85 (7, 2–13)</td>
<td>4.9 (1.9–12.9)</td>
<td>10.35</td>
<td>0.001</td>
</tr>
</tbody>
</table>

CIS, Checklist Individual Strength.  
a. CIS <35 and reliable change index of >1.96.
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