Research article

A meta-analysis of the efficacy of fibromyalgia treatment according to level of care

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Abstract

Introduction The aim of this paper was to compare the efficacy of the treatments for fibromyalgia currently available in both primary care and specialised settings.

Methods Published reports of randomised controlled trials (RCTs) researching pharmacological and non-pharmacological treatments in patients with fibromyalgia were found in the MEDLINE, EMBASE, the Cochrane Central Register of Controlled Trials and PsychInfo databases. The most recent electronic search was undertaken in June 2006.

Results We identified a total of 594 articles. Based on titles and abstracts, 102 full articles were retrieved, 33 of which met the inclusion criteria. These RCTs assessed 120 treatment interventions in 7789 patients diagnosed with primary fibromyalgia. Of them, 4505 (57.8%) were included in the primary care group of our study and 3284 (42.2%) in the specialised care group. The sample was mostly made up of middle-aged women, who have had fibromyalgia for a mean period of 6 to 10 years. The mean effect size of the efficacy of the 120 treatment interventions in patients with fibromyalgia compared with controls was 0.49 (95% confidence interval [CI] = 0.39 to 0.58; p < 0.001). In the primary care group it was 0.46 (95% CI = 0.33 to 0.58) while in specialised care it was 0.53 (95% CI = 0.38 to 0.69), with no statistical significance in the differences. We analysed the efficacy of treatments by comparing primary and specialised care in the different fibromyalgia groups and there were no significant differences. The variables of the studies that affected the improvements in the efficacy of fibromyalgia treatment were low quality of the studies and a shorter duration of treatment. However, both factors were biased by the heterogeneity of the studies. Other variables that also improved outcome and were not biased by the heterogeneity of the studies, were younger age of the patients and shorter duration of the disorder. On the contrary, gender and type of treatment (pharmacological vs. psychological) did not affect outcome.

Conclusion Based on this meta-analysis and despite the heterogeneity of specialised care studies and of the other limitations described in this article, treating fibromyalgia in specialised care offers no clear advantages.

Introduction

Fibromyalgia is a chronic musculoskeletal pain disorder of unknown aetiology, characterised by widespread pain and muscle tenderness and often accompanied by fatigue, sleep disturbance and depressed mood [1,2]. With an estimated lifetime prevalence of approximately 2% in community samples [3], it accounts for 15% of outpatient rheumatology visits and 5% of primary care visits [4]. The prognosis for symptomatic recovery is generally poor [5]. A wide variety of interventions are used in the management of this disorder, although there is

ACR = American College of Rheumatology; CI = confidence interval; FIQ = fibromyalgia impact questionnaire; RCT = randomized controlled trial; SDM = standardised differences in means.
no clear consensus on the treatment of choice and fibromyalgia remains relatively refractory to treatment.

A number of meta-analyses and reviews have been conducted on the pharmacological [6-8] and non-pharmacological [9,10] treatments available for fibromyalgia. The studies main objectives are to guide clinicians in their everyday practice using evidence-based decisions. However, the aim of our current study is rather different. The high prevalence and clinical impact of fibromyalgia makes it a significant public health problem given its high cost. In Spain and other public health systems, a difficult cost-benefit decision must be taken as to which level of the health care system these patients should be treated in: either in specialised settings, which many patients prefer, or in primary care, which is usually more cost-effective. To our knowledge, there is no published meta-analysis on this subject.

We carried out a systematic review and meta-analysis of all randomised controlled trials (RCTs) of pharmacological and non-pharmacological treatments that are available in standard primary care settings and those that are administered in standard secondary care settings of public health care systems in developed countries for the treatment of fibromyalgia. The aim of this paper is to compare the efficacy of the treatments for fibromyalgia available in both settings using the most important outcomes assessed in this disorder, such as pain, quality of life, depression, etc.

Materials and methods
We followed the QUOROM guidelines for reporting meta-analyses [11].

Database search
Published reports of RCTs researching pharmacological or non-pharmacological treatments in patients with fibromyalgia were found in the following databases: MEDLINE (1966–2006), EMBASE (1988–2006), The Cochrane Central Register of Controlled Trials (the Cochrane Library Issue 2006) and Psychinfo (1987–2006). Search strategy is summarised in the additional data file. The search was performed without language restrictions but was limited to RCTs in humans. The last electronic search was undertaken in June 2006. All primary and review articles, as well as their references, were reviewed independently in duplicate. The authors of the original reports were contacted for additional information where needed.

Selection criteria
Studies were screened for inclusion, by reviewing the title and published abstract, based on the following criteria:

Type of participants
The studies evaluated the treatment or management of fibromyalgia as indicated by the use of recognised diagnostic criteria, such as American College of Rheumatology (ACR) [1].

Despite the concept of primary fibromyalgia (patients in which fibromyalgia can not be explained by other medical disorders) not being accepted by the ACR, most studies on fibromyalgia, and many of the papers included in the meta-analysis, do accept this distinction. Therefore, it has been maintained to increase comparability.

Types of studies
The papers described a randomisation of treatment, placebo control and at least one group receiving an active (pharmacological or non-pharmacological) treatment.

Types of interventions
Treatment can be defined as pharmacological or non-pharmacological, and can be allocated to primary or specialised care. The duration of treatment was at least eight weeks.

Types of outcomes
Outcomes had to be measurable. One of the major problems in fibromyalgia is the wide variety of outcomes. Seven types of outcomes were included: pain, fatigue, quality of life, global function, anxiety/depression, insomnia and tender points. Each of them were assessed with several questionnaires.

Each study was reviewed in duplicate (by EF and JGC) for inclusion with substantial inter-rater agreement (kappa = 0.7). Disagreements were resolved by a consensus agreement. Reviews and abstracts were not considered. The study selection process flowchart is summarised in Figure 1.

Allocation
All studies included were allocated to a level of health care (primary care or specialised care) and category of treatment (pharmacological or non-pharmacological) by a consensus with substantial inter-rater agreement (kappa = 0.91) from a panel of two general practitioners (RM and JM), a psychiatrist (JGC) and a psychologist (EF). A treatment was considered to be available at the primary care level when most general practitioners from most Western national health services were able to provide that treatment without any specific training. Tables 1 and 2 summarise which treatments were allocated to the primary and specialised care groups and to the pharmacological and non-pharmacological treatment groups. We have not included RCTs on acupuncture because of the recent meta-analyses showing that this treatment is not effective [10].

Validity assessment
All included reports were then independently read by two reviewers (EF and JGC) who assessed the validity of the studies using the modified Oxford Scale (Table 3) [12,13]. The minimum score of an included trial was one and the maximum was six. Discrepancies were resolved by discussion or by consulting a third reviewer (RM).
Data abstraction
A data abstraction form was created and the following data were included: number of patients and controls, gender (percentage of women), age (median), diagnosis, time of evolution of the disorder (years), severity of the disorder, level of healthcare (primary care or specialist), kind of treatment (pharmacological or non-pharmacological), duration of treatment, modified Oxford Scale ratings and outcomes (ratings in different used scales of quality of life, pain, depression, anxiety, etc).

Meta-analyses
Both dichotomous and continuous data were extracted. Continuous data were analysed as standardised differences in the means (SDM) with 95% confidence intervals (CI). Where mean values and standard deviations were not reported, the authors of the studies were contacted. If they did not reply and the data were presented graphically, data were extracted from the graphs. If this was not possible, the data were not considered. A random effects model was used by default. Analyses were performed using Comprehensive Meta-analysis, version 2 (Biostat, Englewood, NJ, USA). Data were graphically plotted using forest plots to evaluate treatment effects. Clinical heterogeneity was minimised using stringent diagnostic criteria for fibromyalgia and homogeneous criteria for the treatments and outcomes of the studies included in the meta-analysis.

Results

Literature search and study selection
We first performed our literature search in MEDLINE (374 hits), followed by EMBASE (133 hits), and subsequently in the Cochrane Library (41 hits) and in Psychinfo (34 hits). By checking references, we identified an additional 12 hits, resulting in a total of 594 articles (Figure 1). Based on titles and abstracts, 102 full articles were retrieved, 33 of which met the inclusion criteria [14-46]. These 33 studies are summarised in Table 4.

Of the 69 studies that were excluded 23 were not an RCT; the patient population of 28 was not primary fibromyalgia (but secondary fibromyalgia or allied conditions) or the fibromyalgia criteria used were not ACR criteria [1]; in 16 studies the intervention had a duration shorter than eight weeks or it was so specific that it was not available in standard Western health care systems [47]; and two studies did not use comparable outcome measures. There were seven types of outcomes used in the studies selected from 16 questionnaires or tests summarised in Table 5.
Methodological quality of the included studies

Only 11 of the 33 included studies (33.3%) showed a rating of 5+ on the modified Oxford Scale, that is, a score of high methodological quality. Most of them (nine of 11) were pharmacological studies and the remaining two studies were psychological interventions. Many of them were recent studies, carried out in 2004 and 2005 (six of 11), as can be seen in Tables 4 and 6. The most commonly absent items were an adequate description of the flow of patients and adequate description of double blinding.

Study characteristics

The review selected 33 RCTs that assessed 120 treatment interventions on 7789 patients diagnosed with primary fibromyalgia according to ACR criteria [1]. Of these, 4505 (57.8%) were included in the primary care group and 3284 (42.2%) in the specialised intervention group. The characteristics of the patients included in these studies are summarised in Table 6. The sample was made up of middle-aged women, who had the disorder for between six and 10 years (51.9%), treated mainly with pharmacological approaches (73.3%). The outcome types most frequently assessed were pain (26.6%) and global function (23.4%). Most of the patients were from studies carried out in the USA and Canada (63.6%) and were published after 2000 (61.5%). There were no significant differences in any of the variables studied between control and intervention groups.

Table 1

| Primary care      | Secondary care                  |
|-------------------|---------------------------------|
| Amitriptyline     | Pirlindole                      |
| Tramadol          | Tropisetron                     |
| Milnacipran       | Dehydroepiandrosterone (DHEA)   |
| Moclobemide       | Pramipexole                     |
| Fluoxetine        | Malic acid                      |
| Cyclobenzaprine   | Rehabilitation                  |
| Nortriptyline     | Laser treatment                 |
| Duloxetine        | Hyperbaric oxygen therapy       |
| Pregabalin        | Bright light treatment          |
| Zolpidem          | Aerobic exercise                |
|                   | Exercise                         |
|                   | Stress-reduction treatment       |
|                   | Chiropractic management          |
|                   | Cognitive behavioural therapy    |
|                   | Cognitive educational therapy    |
|                   | Education training               |
|                   | Behavioural insomnia therapy     |
|                   | Music vibration                  |

Table 2

| Pharmacological treatments | Non-pharmacological treatments |
|-----------------------------|--------------------------------|
| Amitriptyline               | Hyperbaric oxygen therapy      |
| Cyclobenzaprine             | Bright light treatment          |
| Dehydroepiandrosterone (DHEA)| Aerobic exercise              |
| Duloxetine                  | Education training             |
| Fluoxetine                  | Behavioural therapy            |
| Malic acid                  | Cognitive behavioural therapy  |
| Milnacipran                 | Cognitive educational therapy   |
| Moclobemide                 | Exercise                        |
| Nortriptyline               | Rehabilitation                  |
| Pirlindole                  | Music vibration                 |
| Pramipexole                 | Chiropractic management         |
| Pregabalin                  | Stress-reduction treatment      |
| Tramadol                    | Behavioural insomnia therapy    |
| Tropisetron                 | Laser                           |
| Zolpidem                    |                                |
The mean effect size of the efficacy of the 120 treatment interventions on patients with fibromyalgia compared with efficacy in controls, regardless of the outcome type assessed or the questionnaire used, was 0.49 (95% CI = 0.39 to 0.58; p < 0.001). This is a medium size effect [62], but it is significant. When we compared the efficacy of these treatments on fibromyalgia, after allocating the treatments to primary or specialised level of care, regardless of the type of outcome assessed or the questionnaire used, mean effect size of efficacy in primary care was 0.46 (95% CI = 0.33 to 0.58) while in specialised care was 0.53 (95% CI = 0.38 to 0.69). These differences are not significant.

When we analysed the efficacy of treatments on the different fibromyalgia outcome types (Table 7), we observed that there is an overlapping of the interval scores comparing primary and specialised care for all outcome types. This means that there are no significant differences. There are insignificant differences favouring secondary or specialised care for tender points (mean = 0.28; 95% CI = 0.12 to 0.68 for primary care; mean = 0.50, 95% CI = 0.0 to 1.0 for specialised care) and pain (mean = 0.48, 95% CI = 0.30 to 0.66 for primary care; mean = 0.73, 95% CI = 0.41 to 1.05 for specialised care). On the other hand, there are insignificant differences in favour of primary care for insomnia (mean = 0.57, 95% CI = 0.15 to 0.99 for primary care; mean = -0.18, 95% CI = -0.62 to 0.27 for specialised care), anxiety/depression (mean = 0.59, 95% CI = 0.10 to 1.08 for primary care; mean = 0.40, 95% CI = 0.12 to 0.67 for specialised care), and fatigue (mean = 0.30, 95% CI = 0.05 to 0.56 for primary care; mean = 0.22, 95% CI = -0.08 to 0.52 for specialised care). For specialized care, there are minimal differences also nonsignificant (surely the cause is higher heterogeneity in these studies). Global function, thought to capture the whole impact of the disease, was quite similar in both levels of care (0.53 in primary care; 0.54 in specialised care). The quality of life outcome could not be compared because there were no studies in primary care assessing this variable.

As an example, we have included the efficacy of the treatments allocated to both levels of care in the outcome of pain in patients with fibromyalgia (Figure 2). This outcome is one of the most important in this disorder and the most thoroughly assessed in the studies reviewed. We can observe that there are insignificant differences favouring specialised care (0.73 for specialised care; 0.48 for primary care). However, in this figure, we can also see that heterogeneity in specialised care treatment is higher than in primary care treatment. In fact, there are two studies [40,46] with outcomes of 3.18 and 2.49, respectively, which are the source of this difference. This higher heterogeneity in specialised care treatments compared with primary care treatments is also found in the remaining types of outcome.

In Table 8 we can see the influence of moderating variables on all of the outcomes assessed (overall efficacy), on the specific outcome Global function and on the Fibromyalgia Impact Questionnaire (FIQ). Obviously, we could have included other outcomes and questionnaires but owing to the great amount of information, we selected these variables because they seem to be the most used in assessing the efficacy of fibromyalgia treatments. The results when the other outcomes or questionnaires are analysed are quite similar.

In Table 8 we can also see that an improvement in the methodological quality of the studies is accompanied by a reduction in size effect in the Global Function outcome (the same is found for FIQ scores or for overall efficacy), owing to lesser heterogeneity. Type of treatment, whether pharmacological or non-pharmacological, did not modify the mean effect size in any of the three variables assessed.

On the contrary, shorter length of treatment favours differences that increase size effect. However, these differences can be explained by higher heterogeneity in the studies with shorter treatments, as can be seen on Figure 2 and not by a decrease in the therapeutic effect in longer treatments. With regard to mean participant age, we can observe higher improvement in all outcomes assessed in younger patients. However, the number of studies that evaluate the period of age extremes (young and older people) and assess overall efficacy is low. There are no differences in outcome in relation to

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Table 3

Modified Oxford Scale. Validity score (0 to 7)

| Variable                  | Score |
|---------------------------|-------|
| Randomisation             | 0 None|
|                           | 1 Mentioned |
|                           | 2 Described and adequate |
| Concealment of allocation | 0 None |
|                           | 1 Yes |
| Double blinding           | 0 None |
|                           | 1 Mentioned |
|                           | 2 Described and adequate |
| Flow of patients          | 0 None |
|                           | 1 Described but incomplete |
|                           | 2 Described and adequate |
Table 4

Characteristics of the 33 selected randomised controlled trials and the patients studied in them

| N  | Randomized controlled trial | Year | Country | Treatment | Level of care | % of women | Mean age | Length of treatment (years) | Simple size | Oxford scoring (quality) | Duration of disease at baseline (years) | Instruments used (outcome assessed) |
|----|-----------------------------|------|---------|-----------|---------------|------------|---------|----------------------------|-------------|------------------------|--------------------------------------|-----------------------------------|
| 1  | Carette                     | 94   | Canada  | Amitryptiline, Cyclobenzaprine | Primary care | 93.8       | 44.4    | 24                         | 208         | 3                      | 7.7                                  | Mc Gill-BPI (p) SIP (gf)            |
| 2  | Russell                     | 94   | USA     | Malic acid | Specialised care | 90         | 49.5    | 8                          | 24          | 3                      | 3                                    | VAS (p) TPI (tp)                    |
| 3  | Wolfe                       | 94   | USA     | Fluoxetine | Primary care | 100        | 50.4    | 3                          | 42          | 5                      | 13                                   | TPI (tp) BDI (ad)                   |
| 4  | Carette                     | 95   | Canada  | Amitryptiline | Primary care | 95.5       | 43.8    | 8                          | 22          | 2                      | 6.9                                  | VAS (p) VAS (lg) VAS (gf)           |
| 5  | Chesky                      | 95   | USA     | Music vibration | Specialised care | 92.6       | 48.8    | 30 minutes                 | 26          | 3                      | 11                                   | VAS (p) TPI (tp)                    |
| 6  | Goldenberg                  | 96   | USA     | Fluoxetine, Amitryptiline | Primary care | 90.3       | 43      | 6                          | 31          | 5                      | 5.7                                  | VAS (p) FIQ (GF) BDI (ad) VAS (lg) VAS (gf) VAS (lf) TPI (tp) |
| 7  | Ginsberg                    | 96   | Belgium | Amitryptiline | Primary care | 82.5       | 46      | 8                          | 46          | 2                      | 32                                   | VAS (p) VAS (lg) TPI (lp) VAS (lf) VAS (gf) NTP (tp) |
| 8  | Moldofsky                   | 96   | Canada  | Zolpidem | Primary care | 95        | 42     | 2.5                        | 19          | 4                      | 4                                    | NTP (lp) PGI (i)                    |
| 9  | Vlayen                      | 96   | Holland | Cognitive behavioural therapy Education training | Specialised care | 87        | 44      | 6                          | 131         | 5                      | 10                                   | BDI (ad)                           |
| 10 | Wigers                      | 96   | Norway  | Aerobic exercise, Stress-reduction treatment | Specialised care | 92        | 44      | 14                         | 48          | 3                      | 10                                   | VAS (p) VAS (lg) VAS (lf)           |
| 11 | Pearl                       | 96   | Canada  | Bright light treatment | Specialised care | 100        | 38      | 10                         | 14          | 2                      | 5                                    | VAS (p) VAS (lg) VAS (lf)           |
| 12 | Kelli                       | 97   | Canada  | Chiropractic treatment | Specialised care | -         | 49      | 4                          | 19          | 4                      | 8                                    | VAS (p) NTP (lp)                    |
| 13 | Hannonen                    | 98   | Finland | Moclobemide, Amitryptiline | Primary care | 100       | 49      | 12                         | 130         | 5                      | 11.2                                 | NTP (lp) VAS (p) VAS (lg) VAS (lf) |
| 14 | Yavuzer                     | 98   | Turkey  | Moclobemide | Primary care | 58        | 33      | 6                          | 60          | 1                      | 1                                    | TPI (tp)                           |
| 15 | Ginsberg                    | 98   | Belgium | Parindole | Specialised care | 85        | 40      | 4                          | 61          | 4                      | 2.9                                  | VAS (p) TPI (lp) VAS (gf)           |
| 16 | Russell                     | 99   | USA     | Tramadol | Primary care | 94        | 49      | 6                          | 69          | 4                      | 4.7                                  | VAS (p) FIQ (gf) NTP (lp)           |
| 17 | Heymann                     | 01   | Brazil  | Amitryptiline, Nortryptline | Primary care | 100       | 50      | 8                          | 118         | 4                      | 4                                    | FIQ (gf) NTP (lp)                    |
| 18 | Färber                      | 01   | Germany | Tropisetron | Specialised care | 92        | 48      | 1.5                        | 403         | 3                      | 11                                   | VAS (p) NTP (lp)                    |
| 19 | Gowans                      | 01   | Canada  | Exercise | Specialised care | 88        | 47      | 23                         | 50          | 3                      | 9                                    | FIQ (gf) BDI (ad) STAI (ad) NTP (lp) |
| 20 | Mannerkorpi                 | 01   | Sweden  | Education training | Specialised care | 100       | 46      | 24                         | 58          | 4                      | 8.7                                  | FIQ (gf) QOLS (lg)                  |
| 21 | Gür                         | 02   | Turkey  | Laser, Amitryptiline | Primary care (Amitryptiline) Specialised care (laser) | 80        | 30      | 8                          | 75          | 3                      | 4.6                                  | HADS (ad) FIQ (gf)                  |
gender in any of the three variables evaluated. Finally, the duration of the disorder influences the outcome: a shorter evolution of the disease is associated with higher improvement in any outcome. Again, the number of these kinds of studies is low and heterogeneity is greater, so interpretation of the results is more subjective.

Statistical heterogeneity has been assessed by inconsistency [63]; in our study this is 75%, which is considered to be highly inconsistent. In these cases, the use of random effects analysis is recommended, which we did. A funnel plot between standard error and mean standardised difference, a quality measure to assess publication bias, indicates that most studies are distributed around the central line and are placed in the middle of the graph. There are some small sample studies scattered on the right and on the lower part of the graph that imbalance the weight towards positive values.

**Discussion**

There have been studies assessing multi-modal treatments in primary care [64] and trying to improve the efficacy of primary care treatments for patients with fibromyalgia through better communication [65]. However, to the best of our knowledge this is the first meta-analysis on the efficacy of the treatment of fibromyalgia according to level of care. The clinical and economical relevance of this disorder makes this a key question of research in free, universal health systems in which general practitioners are the gateway to the system. Prevalent and chronic disorders such as fibromyalgia are a huge cost to the health care system [3] and it is necessary to demonstrate whether treatment in a specialised care setting improves the outcome compared with its routine management in a primary care setting.

Only 33 studies from 594 papers examined met the inclusion criteria of our study. These 33 RCTs assessed 120 treatment interventions on patients diagnosed with primary fibromyalgia, 4505 (57.8%) of whom were allocated to primary care and 3284 (42.2%) to specialised care. The sample was made up of middle-aged women, with an average duration of the disorder of six to 10 years, mainly treated with pharmacological approaches. Most of the studies were carried out in the USA.
and Canada and were published after 2000. Owing to the
great variety of outcomes and questionnaires used to assess
the patients, we have summarised the results of the most fre-
quently used in the studies revised: global function, pain and
FIQ. The quality of the studies was rather low with only one-
third of them rating 5+ on the Oxford Scale.

The studies by Yildiz and colleagues [40] and Edinger and col-
leagues [46] could be considered as "outliers" because the
treatments assessed in both studies were much more effica-
cious than the other treatments allocated to specialised care,
but the size sample in both studies was small and the duration
of treatment somewhat short. However, we have not ruled out
these two studies from the meta-analysis for the following
reasons:

• These studies fulfil the stringent selection criteria of the meta-
analysis. Methodological quality was rated independently and
this variable is not an exclusion criteria.

• We expected this meta-analysis to show great heterogeneity
owing to the different kinds of treatments included. We can
not eliminate these studies merely as a result of their hetero-
genesity, since they are as valuable as the other studies
included. We have used a random effects model for the
analysis.

• Both studies assess non-pharmacological treatments and
both were allocated to specialised care. To exclude them
could bias the study towards pharmacological treatments and
primary care.

• We recalculated the meta-analysis excluding these two stud-
ies and the results were the same: there were no significant
differences in the efficacy of the treatments for fibromyalgia
when comparing primary care and specialised care.

Our meta-analysis demonstrates that there are no differences
in the overall outcome of fibromyalgia regardless of the level of
care in which the patient is treated. This article only summa-
rises some outcomes and questionnaires, but we have not
found differences favouring either specialised or primary care
for any of the seven outcomes or the many questionnaires
assessed. In the case of quality of life, the two levels of care
could not be compared. We consider that the external validity
of these data is high because the selection criteria of the stud-
ies allow it to be generalised to most western health services.
However, with respect to internal validity, this data should be

Table 5

| Type of outcome | Questionnaires                                      |
|-----------------|----------------------------------------------------|
| Pain (p)        | McGill Pain Questionnaire [48]                     |
|                 | Brief Pain Inventory [49]                          |
|                 | Visual Analogue Scale [50]                         |
| Quality of life (ql) | SF-36 [51]                        |
|                 | Quality of Life Scale (QOLS) [52]                  |
| Anxiety and depression (ad) | Beck Depression Inventory [53]                |
|                 | Hospital Anxiety and Depression Scale [54]       |
|                 | Hamilton Depression Scale [55]                    |
|                 | State-trait Anxiety Inventory [56]                 |
| Insomnia (i)    | Visual Analog Scale [50]                           |
|                 | Patient Global Impression [57]                     |
| Tender points (tp) | Tender Points Index [58]                        |
|                 | Number of Tender Points according to American College of Rheumatology criteria [1] |
| Fatigue (f)     | Visual Analog Scale [50]                           |
|                 | Multi-dimensional Assessment of Fatigue [59]       |
| Global Function (gf) | Visual Analog Scale [50]                        |
|                 | Fibromyalgia Impact Questionnaire [60]             |
|                 | Clinical Global Impression of Severity [57]        |
|                 | Sickness Impact Profile (SIP) [61]                 |
Table 6

Characteristics of the patients included in the meta-analysis

|                          | Total | %   | Control group | %   | Intervention group | %   |
|--------------------------|-------|-----|---------------|-----|--------------------|-----|
| **Level of care**        |       |     |               |     |                    |     |
| Primary care             | 4505  | 57.8| 2127          | 57.6| 2378               | 58.1|
| Specialised care         | 3284  | 42.2| 1567          | 42.4| 1717               | 41.9|
| Overall                  | 7789  | 100.0| 3694          | 100.0| 4095               | 100.0|
| **Kind of treatment**    |       |     |               |     |                    |     |
| Pharmacological          | 5706  | 73.3| 2684          | 72.7| 3022               | 73.8|
| Non-pharmacological      | 2083  | 26.7| 1010          | 27.3| 1073               | 26.2|
| Overall                  | 7789  | 100.0| 3694          | 100.0| 4095               | 100.0|
| **Outcome assessed**     |       |     |               |     |                    |     |
| Anxiety/depression       | 1195  | 15.3| 570           | 15.4| 625                | 15.3|
| Quality of life          | 115   | 1.5 | 51            | 1.4 | 64                 | 1.6 |
| Pain                     | 2074  | 26.6| 980           | 26.5| 1094               | 26.7|
| Fatigue                  | 650   | 8.3 | 320           | 8.7 | 330                | 8.1 |
| Tender points            | 1533  | 19.7| 738           | 20.0| 795                | 19.4|
| Insomnia                 | 397   | 5.1 | 196           | 5.3 | 201                | 4.9 |
| Global function          | 1825  | 23.4| 839           | 22.7| 986                | 24.1|
| Overall                  | 7789  | 100.0| 3694          | 100.0| 4095               | 100.0|
| **Methodological quality**|      |     |               |     |                    |     |
| 1 to 2                   | 595   | 7.6 | 289           | 7.8 | 306                | 7.5 |
| 3 to 4                   | 3396  | 43.6| 1577          | 42.7| 1819               | 44.4|
| 5 to 6                   | 3798  | 48.8| 1828          | 49.5| 1970               | 48.1|
| Overall                  | 7789  | 100.0| 3694          | 100.0| 4095               | 100.0|
| **Length of treatment (weeks)** | |     |               |     |                    |     |
| 0 to 8                   | 3644  | 46.8| 1750          | 47.4| 1894               | 46.3|
| 09 to 16                 | 3467  | 44.5| 1678          | 45.4| 1789               | 43.7|
| 17 to 24                 | 678   | 8.7 | 266           | 7.2 | 412                | 10.1|
| Overall                  | 7789  | 100.0| 3694          | 100.0| 4095               | 100.0|
| **Age**                  |       |     |               |     |                    |     |
| 30 to 39                 | 253   | 3.2 | 125           | 3.4 | 128                | 3.1 |
| 40 to 49                 | 6662  | 85.5| 3140          | 85.0| 3522               | 86.0|
| 50 to 59                 | 874   | 11.2| 429           | 11.6| 445                | 10.9|
| Overall                  | 7789  | 100.0| 3694          | 100.0| 4095               | 100.0|
| **Percentage of women**  |       |     |               |     |                    |     |
| < 80                     | 151   | 1.9 | 73            | 2.0 | 78                 | 1.9 |
| 80 to 89                 | 2258  | 29.0| 1111          | 30.1| 1147               | 28.0|
analysed cautiously because statistical heterogeneity was important for specialised care studies whereas primary care studies show great homogeneity. In any case, the study points to moderate efficacy of any of the treatments described for fibromyalgia and similar efficacy in both primary and specialised levels of care.

Two of the variables that improve treatment efficacy in fibromyalgia are low quality of the studies and shorter duration of

Table 6 (Continued)

Characteristics of the patients included in the meta-analysis

| Duration of the disorder (years) | 90 to 99 | 100 | Overall |
|---------------------------------|---------|-----|---------|
| 0 to 5                          | 2874    | 1297| 7789    |
| 6 to 10                         | 1213    | 1293| 7889    |
| 11 to 15                        | 3694    | 4095| 8789    |

| Country                        | Percentage |
|--------------------------------|------------|
| Germany                        | 5.3        |
| Belgium                        | 5.9        |
| Brasil                         | 3.6        |
| Canada                         | 21.2       |
| Finland                        | 6.3        |
| Holland                        | 2.2        |
| Norway                         | 2.5        |
| Sweden                         | 3.3        |
| Switzerland                    | 3.5        |
| Turkey                         | 3.8        |
| USA                            | 42.4       |

| Year of publication | 1994 | 1995 | 1996 | 1997 | 1998 | 2000 | 2001 | 2002 | 2003 | 2004 | 2005 | Overall |
|---------------------|------|------|------|------|------|------|------|------|------|------|------|---------|
| Percentage          | 8.0  | 2.3  | 16.8 | 0.5  | 9.3  | 2.7  | 11.9 | 10.0 | 5.0  | 15.9 | 17.7 | 100.0   |
| Percentage          | 6.3  | 2.3  | 12.2 | 0.5  | 9.4  | 2.8  | 12.5 | 10.1 | 5.3  | 16.8 | 16.8 | 100.0   |
| Percentage          | 386  | 92   | 669  | 20   | 375  | 105  | 466  | 408  | 196  | 620  | 758  | 4095    |

(Continued)
treatment, although both of these are biased by the heterogeneity of the studies. Other variables that also improve outcome, which are not biased by the heterogeneity of the studies, are younger patient age and shorter duration of the disorder. In elderly patients, treatment efficacy shows no significant difference when compared with the control group. However, gender and type of treatment (pharmacological vs.

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**Table 7**

| Global function | Pain | Tender points | Quality of life | Anxiety/depression | Insomnia | Fatigue |
|-----------------|------|---------------|----------------|--------------------|----------|---------|
| Low-Up Limit    | Std diff in means | Low-Up Limit | Std diff in means | Low-Up Limit | Std diff in means | Low-Up Limit | Std diff in means | Low-Up Limit | Std diff in means | Low-Up Limit | Std diff in means | Low-Up Limit |
| 0.30 0.76 0.48 0.30 0.66 0.28 0.12 0.68 0.59 0.10 1.08 0.57 0.15 0.99 0.30 0.05 0.56 | 0.32 0.77 0.73 0.41 1.05 0.50 0.00 1.00 1.22 0.49 1.95 0.40 0.12 0.67 - 0.18 0.62 0.22 - 0.08 0.52 | 0.38 0.69 0.54 0.38 0.70 0.37 0.05 0.68 1.22 0.49 1.95 0.44 0.20 ## 0.22 - 0.52 0.27 0.07 0.46 |

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**Figure 2**

Efficacy of the treatments allocated to both levels of care according to the type of pain in patients with fibromyalgia.
Table 8

Influence of moderating variables on all the outcomes assessed, on the Fibromyalgia Impact Questionnaire and on the Global Function

|                          | All the outcome assessed | Fibromyalgia Impact Questionnaire | Global function |
|--------------------------|-------------------------|----------------------------------|-----------------|
|                          | Std dif in means | Lower | Upper | p-value | Std dif in means | Lower | Upper | p-value | Std dif in means | Lower | Upper | p-value |
| Methodological quality   |                        |       |       |         |                        |       |       |         |                        |       |       |         |
| 1 to 2                   | 1.21                   | 0.74  | 1.69  | 0.000   | 1.40                   | 0.75  | 2.04  | 0.000   |                        |       |       |         |
| 3 to 4                   | 0.57                   | 0.42  | 0.72  | 0.000   | 0.66                   | 0.37  | 0.95  | 0.000   | 0.58                   | 0.33  | 0.83  | 0.000   |
| 5 to 6                   | 0.23                   | 0.14  | 0.31  | 0.000   | 0.36                   | 0.18  | 0.55  | 0.000   | 0.37                   | 0.22  | 0.51  | 0.000   |
| Overall                  | 0.33                   | 0.26  | 0.41  | 0.000   | 0.45                   | 0.29  | 0.60  | 0.000   | 0.46                   | 0.33  | 0.58  | 0.000   |
| Kind of treatment        |                        |       |       |         |                        |       |       |         |                        |       |       |         |
| Pharmacological          | 0.42                   | 0.32  | 0.53  | 0.000   | 0.59                   | 0.29  | 0.89  | 0.000   | 0.54                   | 0.33  | 0.74  | 0.000   |
| Non-pharmacological      | 0.63                   | 0.43  | 0.83  | 0.000   | 0.52                   | 0.26  | 0.79  | 0.000   | 0.52                   | 0.26  | 0.79  | 0.000   |
| Overall                  | 0.47                   | 0.37  | 0.56  | 0.000   | 0.55                   | 0.35  | 0.75  | 0.000   | 0.53                   | 0.37  | 0.70  | 0.000   |
| Length of treatment (weeks) |                     |       |       |         |                        |       |       |         |                        |       |       |         |
| 0 to 8                   | 0.73                   | 0.57  | 0.89  | 0.000   | 0.83                   | 0.35  | 1.30  | 0.001   | 0.82                   | 0.50  | 1.14  | 0.000   |
| 9 to 16                  | 0.20                   | 0.11  | 0.28  | 0.000   | 0.35                   | 0.20  | 0.49  | 0.000   | 0.33                   | 0.20  | 0.46  | 0.000   |
| 17 to 24                 | 0.36                   | 0.14  | 0.58  | 0.001   | 0.73                   | 0.29  | 1.16  | 0.001   | 0.35                   | -0.01 | 0.71  | 0.055   |
| Overall                  | 0.31                   | 0.24  | 0.38  | 0.000   | 0.42                   | 0.29  | 0.55  | 0.000   | 0.40                   | 0.28  | 0.51  | 0.000   |
| Age                      |                        |       |       |         |                        |       |       |         |                        |       |       |         |
| 30 to 39                 | 1.41                   | 0.95  | 1.88  | 0.000   | 1.41                   | 0.97  | 1.85  | 0.000   | 1.41                   | 0.97  | 1.85  | 0.000   |
| 40 to 49                 | 0.44                   | 0.35  | 0.54  | 0.000   | 0.37                   | 0.25  | 0.50  | 0.000   | 0.39                   | 0.28  | 0.51  | 0.000   |
| 50 to 59                 | 0.50                   | 0.12  | 0.88  | 0.010   | 0.99                   | -0.22 | 2.20  | 0.108   | 0.99                   | -0.22 | 2.20  | 0.108   |
| Overall                  | 0.48                   | 0.39  | 0.58  | 0.000   | 0.46                   | 0.34  | 0.58  | 0.000   | 0.47                   | 0.35  | 0.58  | 0.000   |
| Women (%)                |                        |       |       |         |                        |       |       |         |                        |       |       |         |
| < 80                     | 3.22                   | 0.68  | 5.76  | 0.013   | 0.96                   | 0.30  | 1.62  | 0.004   | 0.85                   | 0.46  | 1.24  | 0.000   |
| 80 to 89                 | 0.70                   | 0.49  | 0.91  | 0.000   | 0.96                   | 0.30  | 1.62  | 0.004   | 0.85                   | 0.46  | 1.24  | 0.000   |
| 90 to 99                 | 0.33                   | 0.23  | 0.44  | 0.000   | 0.37                   | 0.13  | 0.61  | 0.002   | 0.31                   | 0.14  | 0.47  | 0.000   |
| 100                      | 0.40                   | 0.25  | 0.55  | 0.000   | 0.50                   | 0.23  | 0.78  | 0.000   | 0.50                   | 0.23  | 0.78  | 0.000   |
| Overall                  | 0.41                   | 0.33  | 0.49  | 0.000   | 0.46                   | 0.29  | 0.64  | 0.000   | 0.42                   | 0.28  | 0.55  | 0.000   |
| Duration of disorder (years) |                     |       |       |         |                        |       |       |         |                        |       |       |         |
| 0 to 5                   | 0.85                   | 0.57  | 1.13  | 0.000   | 0.57                   | 0.21  | 0.93  | 0.002   | 0.66                   | 0.34  | 0.98  | 0.000   |
| 6 to 10                  | 0.36                   | 0.27  | 0.45  | 0.000   | 0.45                   | 0.26  | 0.64  | 0.000   | 0.38                   | 0.22  | 0.54  | 0.000   |
| 11 to 15                 | 0.16                   | 0.02  | 0.29  | 0.023   | 0.49                   | 0.33  | 0.65  | 0.000   | 0.45                   | 0.31  | 0.58  | 0.000   |
| Overall                  | 0.36                   | 0.29  | 0.43  | 0.000   | 0.49                   | 0.33  | 0.65  | 0.000   | 0.45                   | 0.31  | 0.58  | 0.000   |
| Country                  |                        |       |       |         |                        |       |       |         |                        |       |       |         |
| Germany                  | 0.36                   | 0.17  | 0.56  | 0.000   | 0.36                   | 0.17  | 0.56  | 0.000   | 0.36                   | 0.17  | 0.56  | 0.000   |
psychological) are not related to outcome. These results insist on the importance of an early diagnosis of the disorder, a fact that is usually related to a younger age of the patient.

Three are three main limitations to this research:

- Heterogeneity of the disorder: there are considered to be several subgroups of patients with fibromyalgia [66], so each of them could have different treatment of choice and, consequently, the results of this study could be different for every subgroup.

- The variability of outcomes used in fibromyalgia: it is difficult to obtain a single outcome summarising the efficacy of the treatment.

- Allocation of treatments to a level of care is a subjective matter: some treatments are new and we cannot foresee which level they will be used at, while others that can theoretically be used in primary care are scarcely utilised at this level. Decisions on which level these should be allocated to had been attained by agreement among four clinicians belonging to different levels and several specialities. Another criticism could be that all the treatments allocated to primary care could also be used at a specialist level, so the comparison should have been primary care treatments vs. all fibromyalgia treatments. However, we did not use this approach because our aim was to assess what was the added value in using specialised treatments for increasing efficacy of fibromyalgia treatment.

The health system operating in Spain is quite similar to the UK and Dutch health systems and is also comparable to other European and Western health systems, with some modifications. The Spanish health system is available to all Spaniards and citizens of the EU and is completely free at all levels. For this reason, there are no economic limitations to accessibility to the system in Spain, and only geographical limitations may exist (inhabitants of small and isolated mountain villages would have more difficulty accessing large hospitals). In this sense, the results of our study could be useful for most Western health systems. Obviously, the results of the meta-analysis are more difficult to extrapolate to countries such as the USA where primary care is not the entrance gate to the system. However, the most important conclusion that can be extrapolated from the study is that family doctors can see similar improvements in their patients with their treatment than specialised clinics in the management of patients with fibromyalgia.

Conclusion

Based on this meta-analysis and despite the heterogeneity of specialised care studies, and of the other limitations described, there is no clear indicator for treating fibromyalgia in specialised care or for the development of specialised units for the treatment of this disorder. According to this study, the same moderate efficacy can be obtained in primary care settings with the routine treatments available at this level. Obviously, any new treatment could upset the balance of this comparison and, whatever the case, new research studies that are more focused on cost-efficacy analysis are necessary to confirm this data.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

JGC is the principal researcher and developed the original idea for the study. The study design was further developed by JM and RM. EFG and MS carried out the electronic search and reviewed the studies. EA and JM developed the statistical
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