Research Article
Revealing the Role of Social Support on Cognitive Deficits in Fibromyalgia Syndrome

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Despite the relevance of cognitive deficits in fibromyalgia syndrome (FMS) and the attempts to elucidate the influence of the disorder symptoms in the cognitive decline reported by patients, no studies have explored the specific role of social support on cognition in FMS. Social support has been shown to be an essential modulator factor on cognitive performance in other diseases. Sixty-four women with FMS and 32 healthy women participated in the study and completed questionnaires pertaining to anxiety, depression, fatigue, insomnia, clinical pain, and social support, along with a neuropsychological battery assessing verbal memory, organization, strategic and planning abilities, self-regulation, processing speed, attention, and cognitive flexibility. Results showed that FMS patients exhibited lower values in all social support dimensions in comparison with healthy individuals, especially in the socializing dimension. Despite the lower social support observed in FMS, all social support dimensions showed a positive impact on verbal memory, organization and planning abilities, strategic planning, self-regulation, processing speed, attention, and cognitive flexibility in these patients. In fact, social support was associated with greater correct responses and processing speed and minor number of errors in all the neuropsychological battery tests. Socializing was the main predictor of organization and planning abilities, strategic planning, and self-regulation. In sum, results suggest that social support may be a key factor in buffering the cognitive decline observed in FMS. Designing psychoeducation programs and intervention programs directed not only to FMS patients but also relatives, health care workers, and the general population might be essential to improve the social support of FMS patients and positively impact on patient’s cognitive status.

1. Introduction

Fibromyalgia syndrome (FMS) is a chronic pain disorder with a prevalence around 2-4% in the general population, being more frequent in women than in men [1]. FMS may be conceptualized as a widespread and persistent musculoskeletal pain, accompanied by several symptoms such as fatigue, insomnia, morning stiffness, depression, anxiety, and cognitive problems [1–3]. Cognitive impairments, which negatively impact on patient’s life, frequently comprise problems in memory, attention, concentration, language, cognitive flexibility, and processing speed, along with reduced organization and planning abilities, among others, and are considered between the most disabling and worrisome symptoms of the disease [3–8].

Emotional aspects also play a relevant role in FMS. FMS has been associated with high negative affectivity [9–11], pain catastrophizing [12–16], alexithymia [17–19], self-esteem, and self-efficacy deficits [3, 20–23]. Negative affectivity (e.g., anxiety, depression, pain catastrophizing, and alexithymia) increases the intensity and severity of symptoms in FMS, worsening the quality of life of these patients [10, 12, 24, 25]. In addition, these emotional aspects have been associated with lessen cognitive performance in FMS [3, 5, 7, 18, 26–30], indicating the relevance of emotional aspects in cognitive deficit in FMS. In the same line, emotional factors are increasing its relevance due to the transdiagnostic perspective, which is showing greater scientific support and playing a crucial role in clinical management [3, 11, 31, 32]. Moreover, the transdiagnostic perspective
seems to be also crucial for personalized behaviour management, which has shown to be essential for mood regulation as an alternative to medications [33].

Furthermore, other factor that can influence FMS symptoms is social support [34]. Social support may be conceptualized as resources provided to people in need by their social network and may be measured through the individual’s perception of the degree to which interpersonal relationships can fulfill certain social support functions [35]. Social support is part of the social network function of each individual, generally related to the number and/or frequency of contacts with family members, relatives, friends, and colleagues [36]. Social support generally comprises several dimensions such as emotional, instrumental, appraisal (which implies information relevant to self-evaluation), and information, among others [34].

At this regard, some studies have reported a lack of social support in FMS patients [37, 38]. Although studies exploring the social support role in FMS symptoms are scarce, social support seems to contribute to improve mental and physical health in FMS patients [34]. In fact, the positive social interaction subcategory of social support has showed a negative association with the Fibromyalgia Impact (measured with the Fibromyalgia Impact Questionnaire) [34], depression state [34, 39], and alexithymia [40]. Besides, social support has been strongly related to anxiety, burnout, and severity of pain in FMS patients [39]. Similarly, Montoya et al. [41] reported that FMS patients perceived less general pain and thermal pain sensitivity as well as diminished brain activity elicited upon tactile stimulation of a tender point when the significant other was present in comparison with when the patients were alone, confirming the notion of social support as a factor explaining pain processing not only at the subjective behavioural level but also at the central nervous system as a factor explaining pain processing not only at the subjective behavioural level but also at the central nervous system as a factor explaining pain processing not only at the subjective level [33].

2. Materials and Methods

2.1. Participants. In total, 64 women with FMS, recruited from the AFIXA (Fibromyalgia Association of Jaén, Spain), participated in the study. All of them were examined by a rheumatologist and met the 1990 and 2010 American College of Rheumatology criteria for FMS [1, 2]. The control group comprised 32 healthy women. Given the main research objective of the study, analyses were restricted to the FMS group. The control group was only used for comparative purposes. Exclusion criteria for both study groups included the presence of metabolic abnormalities, neurological disorders, drug abuse, and severe somatic (e.g., cancer) or psychiatric (e.g., psychotic) diseases. Healthy individuals were further required not to suffer from any kind of acute or chronic pain. All participants were right handed.

2.2. Instruments and Measures

2.2.1. Psychological Assessment. A semistructured interview was performed to obtain the patients’ clinical history and sociodemographic data. The Structured Clinical Interview for Axis I Disorders of the Diagnostic and Statistical Manual for Mental Disorders (SCID) [55] was applied to assess the presence of possible mental disorders. Furthermore, the following self-report questionnaires were administered (values of Cronbach’s α are reported from the available literature):

(i) State-Trait Anxiety Inventory (STAI) [56, 57]. This 20-item 4-point Likert scale’s instrument allows for the assessment of current and habitual anxiety (e.g., state anxiety and trait anxiety, respectively; score range: 0-60). Cronbach’s α = .93 for the state anxiety and .87 for trait anxiety [57]

(ii) Beck Depression Inventory (BDI) [58, 59]. This 21-item scale was applied to assess depression (4-point Likert scales, scores range: 0-63). Cronbach’s α = .95 [59]

(iii) Fatigue Severity Scale (FSS) [60, 61]. This scale allows assessment of fatigue based on 9 items (7-point Likert scales, score range: 9-63). Cronbach’s α = .88 [61]

(iv) Oviedo Quality of Sleep Questionnaire (OQSQ) [62]. The insomnia subscale of this instrument, comprising of 9 items (5-point Likert scales, score range: 9-45), was used in the study. Cronbach’s α of insomnia = .88 [62]

(v) McGill Pain Questionnaire (MPQ) [63, 64]. This 73-item instrument evaluates the different dimensions of pain. In the current research, the total pain...
experience (score range: 0-167) and current pain intensity (MPQ) were used. Cronbach’s α of total pain = .74 [64]

(vi) SS-B Social Support Scale [65, 66] is a 45-item (4-point Likert scales) questionnaire which allow to obtain information of five modes of supportive behaviours: emotional support, socializing, practical assistance, financial assistance, and advice/guidance. Moreover, this questionnaire is typically completed with respect to family and friends separately (family support and friends’ support, respectively), providing information about the supportive behaviour available from relatives and friends. Cronbach’s α = .82 [65]

2.2.2. Cognitive Assessment

(i) Zoo Map Task (ZMT) from the Behavioural Assessment of the Dysexecutive Syndrome [67, 68] was used to evaluate the planning and organizational abilities. In this test, the participant has to plan a route to visit 6 of 12 possible locations in a zoo. The ZMT has two parts: (1) a more demanding open situation, in which little information is provided that would help to generate an appropriate plan, and (2) a situation that implies simply following a concrete, externally imposed strategy. Execution time and number of errors in each part in addition to the total number of correct responses were used as performance indices.

(ii) Verbal Learning Test (TAVEC) [69] assesses verbal memory function. At the beginning of the test, a list of 16 words (shopping list) is read to the participant five times (list A); the participant has to reproduce as many words as possible directly after each trial (immediate free recall). Immediately after, another list is read once (list B) and then has to be reproduced (interference control condition). Following a 20 min break, the words of list A have to be reproduced again (long-delay recall). Thereafter, a list of 44 words is read, which includes all words of list A, some words of list B, and further distractor words included in neither list A nor list B. The participant has to decide whether or not each of these words is part of list A (recognition). In the analysis, list A (immediate free recall), list B (interference control), short-term free memory, short-term guided memory or with semantic keys, long-term free memory, long-term guided memory or with semantic keys, and recognition correct responses were used as performance parameters. Guided memory refers to the trials in which words are reproduced according to semantic categories (e.g., fruits, clothes or tools).

(iii) Revised Strategy Application Test (R-SAT) contains three paper-and-pencil activities, all of which include visual, motor, and linguistic abilities, specifically, figure tracing, sentence, assigned a weighted value, summed, divided by the total possible score for each category, and multiplied by 100. R-SAT allows to measure the strategic planning and self-regulation [70]. The task includes three simple activities, e.g., figure tracing, sentence copying, and object numbering. Activities are presented in two different stacks of 120 items each one. Items differ in terms of their size (large, small) and time requirements (brief, medium, long). A large item scores 0 points, and a small item scores 100 points, where participants are instructed to get as many points as possible. In addition, items in which a face is displayed have to be avoided. The items are inter-mixed; nevertheless, the number of brief items decreases progressively within both stacks. As the execution time of the task is restricted to 10 min, the most efficient strategy is to complete brief items instead of longer ones. Therefore, the predisposition to complete items in the presented sequence has to be overcome. R-SAT further provokes an unstructured environment in the laboratory in which environmental cues and internal habits oppose the most efficient strategy, thus reproducing the real-life situations. At the end of the task, participants are asked about the strategy which, according to their appraisal, was optimal to get the maximal number of points [70]. Participants also have to mark in a separate sheet when they think a minute has been spent (control marks) without using any watch. Performance was indexed by the number of correct answers (brief items), errors (long items and faces), and control marks.

(iv) Trail Making Test (TMT) [71, 72] evaluates processing speed, attention, and cognitive flexibility. The test, in which visual targets (numbers, letters) are presented on sheets of paper, includes the following tasks, all of which have to be executed as fast as possible: (1) visual scanning (cross out all number 3s on a page with different numbers), (2) number sequence (connect the numbers 1 to 15 in sequential order), (3) letter sequence (connect the letters A to P in alphabetic order), (4) switching (connect numbers and letters in alternating order, e.g., 1, A, 2, and B), and (5) motor speed (trace a predefined path). In addition to execution time, the following kinds of errors were recorded: For condition 1: (1) omissions (when the participant fails to mark any 3) and (2) commissions (when the person marks a letter or a number other than 3); for the rest of conditions (2, 3, and 4): (3) sequence (connection of correct item with an incorrect one), (4) set loss (connection of items of different categories) and (5) time out (exceeding the time limit of 250 s).

2.3. Procedure. The study was conducted in one session, divided in two parts conducted on the same day. During the first part, a clinical psychologist took the patients’ clinical
3. Results and Discussion

Table 1 displays the sociodemographic and clinical data of both study groups. FMS patients displayed higher values for all clinical and emotional variables in comparison with healthy individuals (all ps < .0001). In addition, FMS patients displayed lower values for all social support variables compared to healthy individuals (all ps < .05, except financial assistance (SSB) (p = .050) and friends’ support (SSB) (p = .363)).

2.4. Statistical Analysis. In order to determine the optimal sample size based on expected effect sizes, the G*Power 3.1.7 program was used [73]. Assuming an effect size of .75 and an alpha level of .05 and a beta error of 20% as a basis, a sample size from 21 participants per group appeared optimal. Comparisons between FMS patients and healthy individuals in clinical and demographic variables were performed using F-tests and χ²-tests. Group differences in cognitive performance were analysed by means of multivariate analysis of variance (MANOVA). Age, years of education, and body mass index were entered as covariates in this analysis (MANCOVA). A second MANCOVA was performed with the purpose to determine the possible role of emotional (i.e., state and trait anxiety and depression) and clinical variables (i.e., insomnia and fatigue) in cognitive performance. Effect sizes are indicated by adjusted eta squared (η²p). Associations between social support questionnaire dimension scores and neuropsychological test performance were evaluated in two steps, both restricted to the FMS group (N = 64). Firstly, at an exploratory level, Pearson correlations were computed. Secondly, multiple regression analyses were performed. Two blocks of variables were used as predictors in the analyses: (1) to control for the effects of age, body mass index, and years of education, these variables were entered simultaneously (enter method); (2) to determine social support predictive power for cognitive performance, the dimensions of the social support scales that showed significant correlations with the different neuropsychological parameters, in the exploratory analysis, were included (stepwise method) together with total and intensity of clinical pain (MPQ). The inclusion of clinical pain as a possible predictor lies in its relevance as one of the main explanatory mechanisms of cognitive deficits in FMS [7, 8, 26, 74, 75]. The SPSS software (version 22.0) was employed for data analysis (IBM Corporation, Armonk, NY).

3.1. Group Differences in Cognitive Performance. Table 2 shows neuropsychological test scores of FMS patients and healthy individuals and statistics of the univariate group comparisons. The MANOVA for the neuropsychological battery scores showed a multivariate group effect (F [26, 69] = 3.39, p < .0001, η²p = .56). Moreover, this multivariate group effect (F [26, 66] = 2.96, p < .0001, η²p = .54) remains significant in the MANCOVA (using as covariables the age, BMI, and years of education). Additionally, the second MANCOVA (including as covariable state and trait anxiety, depression, fatigue, and insomnia) showed that the multivariate group effect (F [26, 64] = 1.71, p = .042, η²p = .41) remains significant. On purpose, also at the multivariate level, state anxiety (F [26, 64] = 1.41, p = .132, η²p = .37) and insomnia (F [26, 64] = .91, p = .599, η²p = .27) did not show any significant effect. By contrast, trait anxiety (F [26, 64] = 2.18, p = .066, η²p = .47), depression (F [26, 64] = 3.77, p < .0001, η²p = .61), and fatigue (F [26, 64] = 1.90, p = .020, η²p = .44) exhibited a significant multivariate effect.

3.2. Correlations between Social Support and Cognitive Performance in FMS Patients. Table 3 displays correlations between social support dimensions and cognitive performance in FMS patients.

All SSB dimensions were positively associated with organization and planning abilities and strategic planning and self-regulation (total correct responses of ZMT and correct responses: short items of R-SAT). In addition, emotional support (SSB) and friends’ support (SSB) were positively associated with verbal memory (list A: immediate free recall of TAVEC). Emotional support (SSB) was also positively associated with verbal memory (short-term guided memory of TAVEC). Practical assistance (SSB), socializing (SSB), and friends’ support (SSB) were positively associated with strategic planning and self-regulation (control marks of R-SAT).

All SSB dimensions were also negatively associated with errors in organization and planning abilities and strategic planning and self-regulation tasks (error versions 1 and 2 of ZMT and error long items of R-SAT). Moreover, emotional support (SSB), practical assistance (SSB), socializing (SSB), financial assistance (SSB), and advice/guidance (SSB) were negatively associated with errors in strategic planning and self-regulation (error face items of R-SAT). Emotional support (SSB) was also negatively associated with processing time, attention, and cognitive flexibility (execution time 2: number sequence of TMT). Family support (SSB) was negatively associated with the time spending in organization and planning abilities (execution time version 2: ZMT). Practical assistance (SSB), socializing (SSB), financial assistance (SSB), and family support (SSB) were negatively associated with processing speed, attention, and cognitive flexibility (sequence errors of TMT).

3.3. Results of Multiple Regression Analysis. Table 4 shows the significant results of the multiple regression analyses
Regarding the second models, current pain intensity (MPQ) was the main predictor of organization and planning abilities (total correct responses of ZMT and error version 2 of ZMT). Furthermore, advice/guidance (SSB) was the main predictor of errors in organization and planning abilities (error version 2 ZMT). Besides, practical assistance (SSB) was the main predictor of strategic planning and self-regulation and verbal memory (correct responses: short items of R-SAT, error face items of R-SAT, and list A: immediate free recall of TAVEC). Emotional support (SSB) was the main predictor of processing speed, attention, and cognitive flexibility (execution time 2: number sequence of TMT).

4. Discussion

The main objective of the current research was to analyse the effect of social support on the cognitive performance in FMS (including verbal memory, organization, strategic and planning abilities, self-regulation, processing speed, attention, and cognitive flexibility). The present results reaffirm the higher values of clinical and emotional symptoms (e.g., clinical pain, insomnia, fatigue, depression, and anxiety) in FMS compared to healthy individuals [3, 4, 6–8, 18]. Furthermore, the cognitive impairments in FMS (especially in verbal memory, organization, strategic and planning abilities, self-regulation, processing speed, attention, and cognitive flexibility) are confirmed in line with the available scientific evidence [3–8].
Table 2: Mean (±SD) of neuropsychological test scores of FMS patients (N = 64) and healthy individuals (N = 32) and statistics of the univariate group comparisons.

|                          | FMS patients   | Healthy individuals | F [4,66] | p     | \(n^2\) |
|--------------------------|----------------|---------------------|----------|-------|--------|
| **ZMT**                  |                |                     |          |       |        |
| Total correct responses  | 12.13 ± 2.84   | 13.38 ± 2.49        | 3.43     | .067  | .04    |
| Execution time version 1 | 239.33 ± 127.90| 179.06 ± 82.45      | 3.87     | .052  | .04    |
| Execution time version 2 | 159.34 ± 107.42| 90.13 ± 50.32       | 7.91     | .006  | .08    |
| Error version 1          | 2.89 ± 1.96    | 1.94 ± 1.16         | 4.90     | .029  | .05    |
| Error version 2          | .88 ± 1.06     | 2.22 ± 10.56        | .80      | .373  | .01    |
| **TAVEC**                |                |                     |          |       |        |
| List A (immediate free recall) | 37.50 ± 9.98  | 46.31 ± 10.52       | 11.85    | .001  | .12    |
| List B (interference control) | 3.53 ± 1.89   | 4.53 ± 2.00         | 3.82     | .054  | .04    |
| Short-term free memory   | 7.91 ± 3.43    | 10.44 ± 2.97        | 9.25     | .003  | .09    |
| Long-term guided memory  | 9.59 ± 2.73    | 10.84 ± 3.03        | 2.30     | .133  | .03    |
| Long-term free memory    | 8.80 ± 2.82    | 10.53 ± 3.20        | 4.96     | .028  | .05    |
| Recognition correct responses | 9.28 ± 6.06  | 15.16 ± 1.51        | 24.95    | <.0001 | .22    |
| **R-SAT**                |                |                     |          |       |        |
| Correct responses (short items) | 44.22 ± 9.11 | 50.34 ± 7.50        | 13.52    | <.0001 | .13    |
| Error long items         | 4.95 ± 7.41    | 2.03 ± 2.40         | 4.97     | .028  | .05    |
| Error face items         | 1.61 ± 2.10    | .44 ± .91           | 8.31     | .005  | .08    |
| Control marks            | 4.84 ± 2.26    | 6.41 ± 2.28         | 10.00    | .002  | .10    |
| **TMT**                  |                |                     |          |       |        |
| Execution time 1 (visual scanning) | 80.03 ± 108.79| 41.88 ± 13.31       | 3.42     | .068  | .04    |
| Execution time 2 (number sequence) | 98.55 ± 63.96 | 60.31 ± 25.71       | 7.42     | .008  | .08    |
| Execution time 3 (letter sequence) | 105.34 ± 70.65| 66.91 ± 30.18       | 5.61     | .020  | .06    |
| Execution time 4 (switching) | 231.05 ± 130.98| 119.16 ± 51.34     | 18.24    | <.0001 | .17    |
| Execution time 5 (motor speed) | 140.91 ± 58.38| 98.09 ± 49.73       | 7.77     | .006  | .08    |
| Omission errors          | .22 ± .68      | .06 ± .25           | 1.50     | .224  | .02    |
| Commission errors        | .03 ± .18      | .00 ± .00           | 1.76     | .188  | .02    |
| Sequence errors          | 2.28 ± 3.09    | .91 ± 1.51          | 4.25     | .042  | .05    |
| Set loss errors          | 3.05 ± 5.65    | .31 ± .69           | 6.53     | .012  | .07    |
| Time out errors          | 10.91 ± 10.80  | .78 ± 2.06          | 22.05    | <.0001 | .20    |

Note: ZMT: Zoo Map Test; TAVEC: Verbal Learning Test; R-SAT: Revised Strategy Application Test; TMT: Trail Making Test. All execution times are indicated in s. *p < .05. **p < .01.

Results of the second MANCOVA further reveal that group differences in cognitive performance seem to be independent of emotional and clinical symptoms [76, 77]. At this regard, Roldán-Tapia et al. [76] pointed out that cognitive impairment in FMS patients could not be explained by the collateral effects of such pathologies, because cognitive profiles were different and appeared from the onset of the disease notion also supported by the research of Simos et al. [77].

Refer to social support, FMS patients displayed lower values for all social support variables compared to healthy individuals [37, 38], especially in the socializing area. Nevertheless, there was no any significant difference in financial assistance and friends’ support between both groups. Although more studies are required to firm clear conclusions, some research has suggested that FMS patients seem to be more likely to include their physicians as intimate members of their social networks and less willing to take initiative in meeting new people than patients with rheumatoid arthritis (RA) [78]. This may explain the lower observed social support values associated with the socializing sphere in FMS patients. This finding also reinforces the significant role of health professionals in FMS [79]. Another explanation might relapse in the fact that FMS patients usually perceive little social support at work due to the lack of social knowledge and awareness on the disease [80]. This lack of social support reduces personal relationships at work. This is influenced by the misunderstanding about the lower effectiveness of FMS patients at work [80]. At the same time, the problems in the professional field and the need to stop working in some cases because of the illness symptoms might worsen the social isolation. It is well known that staying in the workplace prevents FMS patients from social isolation and reduces the negative impact of the disease on their quality of life [81]. Therefore, it is necessary to design programs...
|                          | Emotional support (SSB) | Practical assistance (SSB) | Socializing (SSB) | Financial assistance (SSB) | Advice/guidance (SSB) | Family support (SSB) | Friends' support (SSB) |
|--------------------------|-------------------------|---------------------------|------------------|---------------------------|----------------------|---------------------|-----------------------|
| **ZMT**                  |                         |                           |                  |                           |                      |                     |                       |
| Total correct responses  | .367**                  | .387**                    | .421**           | .396**                    | .368**               | .360**              | .282**                |
| Execution time version 1 | .050                    | -.009                     | .046             | .039                      | .063                 | .012                | .118                  |
| Execution time version 2 | -.124                   | -.203                     | -.192            | -.181                     | -.157               | -.257*              | -.047                 |
| Error version 1          | -.385**                 | -.419**                   | -.438**          | -.434**                   | -.371**              | -.326**             | -.364**               |
| Error version 2          | -.373**                 | -.385**                   | -.425**          | -.395**                   | -.377**              | -.331**             | -.299**               |
| **TAVEC**                |                         |                           |                  |                           |                      |                     |                       |
| List A (immediate free recall) | .274*                  | .101                      | .143             | .129                      | .158                | -.023               | .269*                 |
| List B (interference control) | .138                   | .079                      | .068             | .087                      | .083                | .045                | .071                  |
| Short-term free memory   | .223                    | .138                      | .163             | .162                      | .131                | .094                | .203                  |
| Short-term guided memory | .247*                   | .150                      | .155             | .181                      | .107                | .016                | .163                  |
| Long-term free memory    | .186                    | .189                      | .197             | .194                      | .136                | .049                | .232                  |
| Long-term guided memory  | .162                    | .068                      | .111             | .090                      | .027                | -.064               | .124                  |
| Recognition correct responses | .134                   | .118                      | .094             | .137                      | .159                | .202                | .095                  |
| **R-SAT**                |                         |                           |                  |                           |                      |                     |                       |
| Correct responses (short items) | .401**                  | .404**                    | .460**           | .411**                    | .419**              | .312*               | .370**                |
| Error long items         | -.362**                 | -.343**                   | -.376**          | -.391**                   | -.366**             | -.312*              | -.307*                |
| Error face items         | -.348**                 | -.318*                    | -.344**          | -.384**                   | -.338**             | -.241               | -.235                 |
| Control marks            | .094                    | .249*                     | .252*            | .237                      | .225                | .139                | .258*                 |
| **TMT**                  |                         |                           |                  |                           |                      |                     |                       |
| Execution time 1 (visual scanning) | .003                   | .190                      | .217             | .149                      | .072                | .023                | .124                  |
| Execution time 2 (number sequence) | -.257*                  | -.217                     | -.224            | -.223                     | -.189               | -.188               | -.131                 |
| Execution time 3 (letter sequence) | -.230                   | -.165                     | -.171            | -.182                     | -.153               | -.110               | -.141                 |
| Execution time 4 (switching) | -.220                   | -.226                     | -.233            | -.200                     | -.204               | -.156               | -.139                 |
| Execution time 5 (motor speed) | -.067                   | -.166                     | -.130            | -.113                     | -.126               | -.166               | -.066                 |
| Omission errors          | -.190                   | -.262*                    | -.251*           | -.252*                    | -.204               | -.356**             | -.120                 |
| Commission errors        | -.098                   | -.122                     | -.129            | -.118                     | -.115               | .029                | -.197                 |
| Sequence errors          | -.254*                  | -.262*                    | -.306*           | -.293*                    | -.270*              | -.116               | -.308*                |
| Set loss errors          | -.174                   | -.175                     | -.201            | -.217                     | -.186               | -.169               | -.122                 |
| Time out errors          | -.134                   | -.227                     | -.232            | -.169                     | -.177               | -.194               | -.104                 |

Note: ZMT: Zoo Map Test; TAVEC: Verbal Learning Test; R-SAT: Revised Strategy Application Test; TMT: Trail Making Test. *p < .05. **p < .01.
to increase the training and sensitization of relatives and friends of FMS patients, as well as of health providers and general population [79].

Related to social support and cognitive performance in FMS patients, all studied SBB dimensions (e.g., emotional support, practical assistance, socializing, financial assistance, advice/guidance, family support, and friends’ support) were positively related to a better cognitive performance (higher levels of correct responses) in the organization and planning abilities (ZMT) and strategic planning and self-regulation tasks (R-SAT). Similarly, practical assistance, socializing, and friends’ support (SSB) were positively linked to higher strategic planning and self-regulation performance (R-SAT). Moreover, emotional support and friends’ support were positively associated with a better performance in the verbal memory domain (TAVEC).

In addition, the social support was not only positively related with the number of correct responses of neuropsychological test but also was associated with a reduction of the processing speed. In fact, family support was negatively associated with execution time version 2 of ZMT, which measures organization and planning abilities, whereas emotional support was also negatively linked to execution time 2 (number sequence) of TMT which assesses processing speed, attention, and cognitive flexibility.

Regarding the number of errors, social support also might be involved in the reduction of these one. In this sense, practical assistance, socializing, financial assistance, and advice/guidance were negatively associated with errors in organization and planning abilities tasks and strategic planning and self-regulation tasks (errors of both versions in ZMT and error long items of R-SAT, respectively). Additionally, emotional support, practical assistance, socializing, financial assistance, and advice/guidance were negatively related to error face items of R-SAT, which measure strategic planning and self-regulation performance. And all dimensions of the social support scale were negatively associated with different errors of TMT, which assesses processing speed, attention, and cognitive flexibility.

| Table 4: Significant results of the second block (step-wise method) of the multiple regression analysis for the prediction of neuropsychological test scores by clinical pain and social support variables in FMS patients (N = 64). |
|------------------|---------------------------------|--------|------|-----|
| M               | Predictors                       | β      | Δr²  | t   |
| Total correct responses | Socializing (SSB)                            | .43    | .18  | 4.20| <.001|
|                  | Socializing (SSB)                            | .43    | .08  | 4.52| <.001|
|                  | Current pain intensity (MPQ)                       | -.30   | -.06 | -3.06| .003|
| Execution time version 2 | Current pain intensity (MPQ)                       | .30    | .09  | 2.65| .010|
|                  | Socializing (SSB)                            | -.40   | -.15 | -3.77| <.001|
| ZMT              | Socializing (SSB)                            | -1.37  | -.05 | -3.07| .003|
| Error version 1  | Advice/guidance (SSB)                        | 1.00   | 2.24 | .029|
|                  | Socializing (SSB)                            | -.39   | -.15 | -3.50| .001|
| Error version 2  | Socializing (SSB)                            | -.39   | -.07 | -3.68| .001|
|                  | Current pain intensity (MPQ)                       | .28    | 2.49 | .015|
| TAVEC            | Friends’ support (SSB)                         | .29    | .08  | 2.53| .014|
| List A (immediate free recall) | Friends’ support (SSB)                         | .74    | 3.79 | <.001|
|                  | Practical assistance (SSB)                       | -.55   | -.29 | <.001|
| Short-term guided memory | Current pain intensity (MPQ)                       | -.34   | -.11 | -2.89| .005|
| Correct responses (short items) | Socializing (SSB)                            | .47    | .22  | 4.38| <.001|
|                  | Socializing (SSB)                            | 1.44   | 2.97 | .004|
| R-SAT            | Practical assistance (SSB)                       | -.99   | -.05 | -2.05| .045|
| Errors (long items) | Financial assistance (SSB)                       | -.38   | -.14 | -3.15| .003|
|                  | Financial assistance (SSB)                       | -.39   | -.15 | -3.21| .002|
| Errors (face items) | Financial assistance (SSB)                       | -1.67  | -.08 | -3.10| .003|
|                  | Practical assistance (SSB)                       | 1.31   | 2.43 | .018|
| Control marks    | Friends’ support (SSB)                         | .26    | .07  | 2.09| .041|
| Execution time 2 (number sequence) | Current pain intensity (MPQ)                       | .34    | .10  | 3.21| .002|
| TMT              | Current pain intensity (MPQ)                       | .36    | .05  | 3.53| .001|
| Sequence errors  | Emotional support (SSB)                         | -.23   | -.23 | .030|
|                  | Friends’ support (SSB)                         | -.31   | -.10 | -2.57| .013|
| Omissions errors | Family support (SSB)                          | -.36   | -.12 | -2.89| .005|

Note: Model (M), standardized β, change in r² (Δr²), t, and p are indicated. Results of the first block, which served to control for the effects of age, education, and BMI, are not reported. ZMT: Zoo Map Test; TAVEC: Verbal Learning Test; R-SAT: Revised Strategy Application Test; TMT: Trail Making Test.
Moreover, socializing was the main predictor of organization and planning abilities and strategic planning and self-regulation (total correct responses of ZMT and R-SAT and errors of both versions in the ZMT test). One possible explanation can be the positive effect of social support in pain processing at the subjective behavioural level and at the central nervous system level of FMS patients [41]. In addition, an analgesic effect of social support has been reported in healthy population, even without verbal or physical contact [42]. Similarly, it is possible that socializing might have an analgesic effect on pain (one of the main explanatory factors of cognitive impairments in FMS [8, 74], which indirectly improve the cognitive performance of patients). Otherwise, this analgesic effect of social support in pain processing might be also favouring the cognitive processing areas. It is well known that pain is an attention-demanding stimulus that recruits brain areas also relevant for cognitive processing [8, 74, 82]. It would be interesting to assess the possible mediating role of social support in the relation between pain and cognitive performance in FMS.

Our research reveals the positive effect of social support especially in verbal memory, organization and planning abilities, strategic planning, self-regulation, processing speed, attention, and cognitive flexibility. Results are in line with previous studies suggesting a positive association between social support and health outcomes in FMS patients [34, 39–41].

Additionally, emotional support showed to be the social support’s variable that account for the majority of associations with the cognitive parameters, suggesting, among all SBB dimensions, a special role of it in FMS cognition. This finding is according to previous studies that pointed out that social support, and particularly emotional support, was associated with decreases in health care use within a primary care setting [83]. Unfortunately, although the previous mentioned research is very meaningful, no cognitive aspects were evaluated, being this study the first one on this issue. Therefore, the role of social support on cognitive performance in FMS can be considered a research gap that needs to be overcome to better understand the disease and provide FMS patients a more holistic and personalized treatment. In addition, emotional support would need also to be promoted based on the transdiagnostic perspective which insist on the relevance of emotional and social aspect of the illnesses apart from treat on the disease symptoms [31, 32]. Furthermore, based on previous findings [33], it might be hypothesized the relevance of improving social support, especially emotional support, in order to personalized FMS treatment.

Nonetheless, despite the role of social support in FMS needs to continue being studied, our results may be explained based on previous research in other populations, which point out that social support is a significant factor involved in the maintenance or enhancement of mental health and cognitive functioning in general (e.g., elderly people [43–49], caregivers [50], and academic performance [51–53, 84]). In this regard, it has been noted that social relationships, especially social activities and networks, have a protective effect against greater cognitive decline in older adults [85, 86]. Cotton et al. [87] explored the neural correlates of social support in older adults and reported the existence of a gray matter network related to social support (including prefrontal, hippocampal, amygdala, cingulate, and thalamic regions), which was in turn associated with memory and executive function. Cacioppo and Hawkley [88] further showed that this gray matter network associated with tangible social support was composed by regions previously linked to memory, executive function, aging, and dementia. Authors concluded that more longitudinal research of the interrelationships between social support, brain structure, and cognition was needed and advised of the importance of strengthen social support as a new path toward improving cognition in aging [88]. Moreover, research and interventions in this field would help to better understand the contribution of poor psychosocial functioning (e.g., social support) and unsatisfactory relationships in the origin and maintenance of chronic pain [40], especially in FMS.

Likewise, social isolation (e.g., loneliness) has been proposed as a risk factor for and may contribute to poorer overall cognitive performance, faster cognitive decline, poorer executive functioning, and higher negativity and depressive cognition [88]. Loneliness has exhibited a mediating role between social isolation and subjective cognitive impairment in older people [89–92], including older immigrants [89]. The association between loneliness and subjective cognitive deficits has been well-established [93, 94]. On purpose, social support, indeed, is known to play a critical role in the detection and treatment of mild cognitive impairment [95–97].

Loneliness has been proposed as a risk factor of dementias, especially Alzheimer disease [96–99]. In general, higher levels of social engagement have been related to greater levels of cognition across the lifespan, association that seems to be more significant in populations at risk of cognitive impairment [100]. The research in other populations confirmed the protective role of social support in cognitive performance, although the exact nature of this association remains unclear [46, 85, 86, 89, 93, 94, 100]. Previous research also highlights the need of promoting psychosocial interventions and related public health strategies to enhance neurocognitive health, increasing specific forms of social support [94, 100–102], such as supportive listening [102]. In fact, social support group interventions in people with dementia and mild cognitive impairment exhibited psychological benefits, specifically, a reduction of depression and an improvement of the quality of life and the self-esteem [103]. Considering the prevalence and negative impact of cognitive deficits in FMS, it is worth keep exploring the effect of social support at this regard.

The main limitation of our study was its cross-sectional design, which does not allow for the establishment of causal associations. Longitudinal studies are required to better understand the association between social support and cognitive impairment in FMS. It also would be interesting to compare FMS patients with other clinical samples (e.g., RA patients), to determine if the observed association between social support and better cognitive performance is or not specific of FMS but can be extended to other pain conditions. Strengths of the study included the novelty and clinical relevance of the results, the statistical control of ...
sociodemographic variables, and the determination of the sample through the G*Power program to ensure the statistical power of the analysis. Moreover, social support in FMS has been evaluated conforming the recommendation of not only to consider the amount of support reported by the patients but also the type of support they received [40].

Due to the relevance of cognitive deficits in FMS [3–8, 104], a routine screening for cognitive impairments in these patients should be included in both diagnosis and treatment [105]. Moreover, taking the biopsychosocial perspective into account, it must be also mandatory to explore and promote the social support network of these patients in order to prevent the worsening of cognitive performance and subsequent health-related quality of life. It is necessary to continue researching the mechanism through the social support benefit the cognitive performance in chronic pain patients, including FMS, and confirming the protective role of social support in this kind of disorders. It might be a promising future line of research and clinical practice to improve the quality of life of these patients.

Furthermore, the design of psychoeducation programs and intervention programs to improve the social support of FMS cannot only positively influence on patient’s status but also in the health system as proposed by previous studies [106, 107]. In fact, some researches showed that one’s social network was positively linked to health status and negatively related to health care use, reducing prescriptions, laboratory tests, and visits to a nurse, nurse practitioner, and/or physicians’ assistant [107]. Nonetheless, it is important to highlight that for establishing an effective social support network for FMS patients, it is necessary to provide relatives and friends with disease information for a better understanding. Therefore, it is important to also involve them in the psychoeducation programs [108]. At this regard, Kool et al. [109] analysed if social support and invalidation (lack of understanding and discounting by others) were differently related to physical and mental health. They studied 1455 patients with different chronic pain diseases, such as FMS, RA, ankylosing spondylitis, and osteoarthritis [109]. Their results confirmed that both invalidation and social support were linked to patients’ mental health, but only invalidation was significantly related to patients’ physical health, suggesting the relevance to include social support and invalidation in programs to improve health of patients with rheumatic diseases [109]. Besides, the therapeutic adherence in some FMS programs seems to be associated with some factors such as lack of motivation and lack of social support, among others [110]; therefore, enhancing the social support of these patients might have a positive impact in their treatments.

5. Conclusions

This is the first study exploring the influence of social support in cognitive performance of FMS patients. FMS displayed lower values in all social support variables compared to healthy individuals, especially in the socializing area, which was the main predictor of organization and planning abilities, and strategic planning and self-regulation of these patients (total correct responses of ZMT and R-SAT and errors of both versions in the ZMT test). All dimensions of social support exhibited a positive impact on verbal memory, organization and planning abilities, strategic planning, self-regulation, processing speed, attention, and cognitive flexibility. Social support dimensions not only positively impact on the number of correct responses and processing speed of neuropsychological test but also seem to reduce the number of errors. Improving the social networks of FMS patients might help to ameliorate their health status and cognitive performance while simultaneously reducing health care utilization. It is vital to involve not only FMS patients but also their relatives and friends along with health care workers and general population, in order to improve the FMS knowledge and sensitize people to the importance of social support in this disease.

Data Availability

The data presented in this study are available on request from the corresponding author.

Ethical Approval

The procedure followed the general criterion of the local ethics committee, based on the Helsinki Declaration principles, and was approved by the Bioethics Committee of the University of Jaén.

Conflicts of Interest

All the authors declare that they have no conflict of interest derived from the outcomes of this study.

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References

[1] F. Wolfe, D. J. Clauw, M. A. Fitzcharles et al., “The American College of Rheumatology preliminary diagnostic criteria for fibromyalgia and measurement of symptom severity,” Arthritis Care & Research, vol. 62, no. 5, pp. 600–610, 2010.
[2] F. Wolfe, H. A. Smythe, M. B. Yunus et al., “The American College of Rheumatology 1990 criteria for the classification of fibromyalgia,” Arthritis and Rheumatism, vol. 33, no. 2, pp. 160–172, 1990.
[3] C. M. Galvez-Sánchez, G. A. Reyes Del Paso, and S. Duschek, “Cognitive impairments in fibromyalgia syndrome: associations with positive and negative affect, alexithymia, pain catastrophizing and self-esteem,” Frontiers in Psychology, vol. 9, p. 377, 2018.
[4] C. M. Galvez-Sánchez, P. de la Coba, J. M. Colmenero, G. A. Reyes Del Paso, and S. Duschek, “Attentional function in
fibromyalgia and rheumatoid arthritis,” *PLoS One*, vol. 16, no. 1, article e0246128, 2021.

[5] O. Gelonch, M. Garolera, J. Valls, L. Rosselló, and J. Pifarré, “Cognitive complaints in women with fibromyalgia: are they due to depression or to objective cognitive dysfunction?,” *Journal of Clinical and Experimental Neuropsychology*, vol. 39, no. 10, pp. 1013–1025, 2017.

[6] C. Muñoz Ladrón de Guevara, M. J. Fernández-Serrano, G. A. Reyes del Paso, and S. Duschek, “Executive function impairments in fibromyalgia syndrome: relevance of clinical variables and body mass index,” *PLoS One*, vol. 13, no. 4, article e0196329, 2018.

[7] C. I. Montoro, S. Duschek, C. Muñoz Ladrón de Guevara, M. J. Fernández-Serrano, and G. A. Reyes del Paso, “Aberrant cerebral blood flow responses during cognition: implications for the understanding of cognitive deficits in fibromyalgia,” *Neuropsychology*, vol. 29, no. 2, pp. 173–182, 2015.

[8] G. A. Reyes del Paso, Á. Pulgar, S. Duschek, and S. Garrido, “Cognitive impairment in fibromyalgia syndrome: the impact of cardiovascular regulation, pain, emotional disorders and medication,” *European Journal of Pain*, vol. 16, no. 3, pp. 421–429, 2012.

[9] D. Watson, L. A. Clark, and A. Tellegen, “Development and validation of brief measures of positive and negative affect: the PANAS scales,” *Journal of Personality and Social Psychology*, vol. 54, no. 6, pp. 1063–1070, 1988.

[10] P. H. Finan, A. J. Zautra, and M. C. Davis, “Daily affect relations in fibromyalgia patients reveal positive affective disturbance,” *Psychosomatic Medicine*, vol. 71, no. 4, pp. 474–482, 2009.

[11] K. Malin and G. O. Littlejohn, “Stress modulates key psychological processes and characteristic symptoms in females with fibromyalgia,” *Clinical and Experimental Rheumatology*, vol. 31, 6 Supplement 79, pp. S64–S71, 2013.

[12] C. M. Galvez-Sánchez, C. I. Montoro, S. Duschek, and G. A. Reyes del Paso, “Pain catastrophizing mediates the negative influence of pain and trait-anxiety on health-related quality of life in fibromyalgia,” *Quality of Life Research: An International Journal of Quality of Life Aspects of Treatment, Care and Rehabilitation*, vol. 29, no. 7, pp. 1871–1881, 2020.

[13] C. P. Van Wilgen, M. W. van Ittersum, A. A. Kaptein, and M. van Wiehe, “Illness perceptions in patients with fibromyalgia and their relationship to quality of life and catastrophizing,” *Arthritis and Rheumatism*, vol. 58, no. 11, pp. 3618–3626, 2008.

[14] R. R. Edwards, C. O. Bingham, J. Batson, and J. A. Haythornthwaite, “Catastrophizing and pain in arthritis, fibromyalgia, and other rheumatic diseases,” *Arthritis and Rheumatism*, vol. 55, no. 2, pp. 325–332, 2006.

[15] C. Toth, S. Brady, and M. Hatfield, “The importance of catastrophizing for successful pharmacological treatment of peripheral neuropathic pain,” *Journal of Pain Research*, vol. 7, pp. 327–338, 2014.

[16] B. Rodero, J. García-Campayo, B. Casanueva, and Y. Buriel, “Tratamientos no farmacológicos en fibromialgia: una revisión actual,” *Revista de Psicopatología Psicologia Clinica*, vol. 14, no. 3, pp. 137–151, 2009.

[17] L. Castelli, V. Tesio, F. Colonna et al., “Alexithymia and psychological distress in fibromyalgia: prevalence and relation with quality of life,” *Clinical and Experimental Rheumatology*, vol. 30, 6 Supplement 74, pp. 70–77, 2012.

[18] C. I. Montoro, G. A. Reyes del Paso, and S. y Duschek, “Alexithymia in fibromyalgia syndrome,” *Personality and Individual Differences*, vol. 102, pp. 170–179, 2016.

[19] M. P. Martinez, A. Sánchez, E. Miró, M. J. Lami, G. Prados, and A. Morales, “Relationships between physical symptoms, emotional distress, and pain appraisal in fibromyalgia: the moderator effect of alexithymia,” *The Journal of Psychology*, vol. 149, pp. 115–140, 2014.

[20] H. J. Michielsen, B. Van Houdenhove, I. Leirs, A. Vandenbroeck, and P. Onghena, “Depression, attribution style and self-esteem in chronic fatigue syndrome and fibromyalgia patients: is there a link?,” *Clinical Rheumatology*, vol. 25, no. 2, pp. 183–188, 2006.

[21] C. Peñacoba Puente, L. Velasco Furlong, C. Écija Gallardo, M. Cigarrán Méndez, D. Bedmar Cruz, and C. Fernández-de-las-Peñas, “Self-efficacy and affect as mediators between pain dimensions and emotional symptoms and functional limitation in women with fibromyalgia,” *Pain Management Nursing*, vol. 16, no. 1, pp. 60–68, 2015.

[22] M. Garaigordobil and L. Govilard, “Fibromyalgia: discapacidad funcional, autoestima y perfil de personalidad,” *Información Psicológica*, vol. 106, pp. 4–16, 2013.

[23] E. Miró, M. P. Martinez, A. I. Sánchez, G. Prados, and A. Medina, “When is pain related to emotional distress and daily functioning in fibromyalgia syndrome? The mediating roles of self-efficacy and sleep quality,” *British Journal of Health Psychology*, vol. 16, no. 4, pp. 799–814, 2011.

[24] A. J. Zautra, R. Fasman, J. W. Reich et al., “Fibromyalgia: evidence for deficits in positive affect regulation,” *Psychosomatic Medicine*, vol. 67, no. 1, pp. 147–155, 2005.

[25] C. M. Galvez-Sánchez, C. I. Montoro, S. Duschek, and G. A. Reyes Del Paso, “Depression and trait-anxiety mediate the influence of clinical pain on health-related quality of life in fibromyalgia,” *Journal of Affective Disorders*, vol. 265, pp. 486–495, 2020.

[26] D. Munguía-Izquierdo, A. Legaz-Arrese, D. Moliner-Urdiales, and J. Reverter-Masia, “Neuropsychological performance in patients with fibromyalgia syndrome: relation to pain and anxiety,” *Psicothema*, vol. 20, no. 3, pp. 427–431, 2008.

[27] R. Schütze, C. Rees, M. Preece, and M. Schütze, “Low mindfulness predicts pain catastrophizing in a fear-avoidance model of chronic pain,” *Pain*, vol. 148, no. 1, pp. 120–127, 2010.

[28] O. Gelonch, M. Garolera, J. Valls, L. Rosselló, and J. Pifarré, “Executive function in fibromyalgia: comparing subjective and objective measures,” *Comprehensive Psychiatry*, vol. 66, pp. 113–122, 2016.

[29] K. V. Keefer, G. J. Taylor, J. Parker, and R. M. Bagby, “Taxometric analysis of the Toronto structured interview for alexithymia: further evidence that alexithymia is a dimensional construct,” *Assessment*, vol. 26, no. 3, pp. 364–374, 2019.

[30] A. L. Hassett, L. E. Simonelli, D. C. Radvanski, S. Buyske, S. V. Savage, and L. H. Sigal, “The relationship between affect balance style and clinical outcomes in fibromyalgia,” *Arthritis and Rheumatism*, vol. 59, no. 6, pp. 833–840, 2008.

[31] B. Faustino, “Transdiagnostic perspective on psychological inflexibility and emotional dysregulation,” *Behavioural and Cognitive Psychotherapy*, vol. 49, no. 2, pp. 233–246, 2021.

[32] P. Amici, “Intolerance of uncertainty: from transdiagnostic model to clinical management,” *Psychiatria Danubina*, vol. 33, Supplement 9, pp. 22–25, 2021.
[33] D. M. Davydov, C. M. Galvez-Sanchez, C. I. Montoro, C. M. L. de Guevara, and G. A. Reyes del Paso, "Personalized behavior management as a replacement for medications for pain control and mood regulation," Scientific Reports, vol. 11, no. 1, p. 20297, 2021.

[34] R. Freitas, S. C. Andrade, M. Spyrides, M. Micussi, and M. Souza, "Impacto do apoio social sobre os sintomas de mulheres brasileiras com fibromialgia," Revista Brasileira de Reumatologia, vol. 57, no. 3, pp. 197–203, 2017.

[35] A. J. Oliveira, C. S. Lopes, A. de Leon et al., "Social support and leisure-time physical activity: longitudinal evidence from the Brazilian Pró-Saúde cohort study," The International Journal of Behavioral Nutrition and Physical Activity, vol. 8, no. 1, p. 77, 2011.

[36] J. E. Hupecey, "Clarifying the social support theory-research linkage," Journal of Advanced Nursing, vol. 27, no. 6, pp. 1231–1241, 1998.

[37] L. M. Arnold, L. J. Crofe, P. J. Mease et al., "Patient perspectives on the impact of fibromyalgia," Patient Education and Counseling, vol. 73, no. 1, pp. 114–120, 2008.

[38] A. L. Bernard, A. Prince, and P. Edsall, "Quality of life issues for fibromyalgia patients," Arthritis Care and Research, vol. 13, no. 1, pp. 42–50, 2000.

[39] N. Gündüz, A. Üşen, and E. Aydin Atar, "The impact of perceived social support on anxiety, depression and severity of pain and burnout among Turkish females with fibromyalgia," Archives of Rheumatology, vol. 34, no. 2, pp. 186–195, 2018.

[40] M. Di Tella, V. Tesio, A. Ghiggia et al., "Coping strategies and perceived social support in fibromyalgia syndrome: relationship with alexithymia," Scandinavian Journal of Psychology, vol. 59, no. 2, pp. 167–176, 2018.

[41] P. Montoya, W. Larbig, C. Braun, H. Preisel, and N. Birbaumer, "Influence of social support and emotional context on pain processing and magnetic brain responses in fibromyalgia," Arthritis and Rheumatism, vol. 50, no. 12, pp. 4035–4044, 2004.

[42] S. Duschek, L. Nassauer, C. I. Montoro, A. Bair, and P. Montoya, "Dispositional empathy is associated with experimental pain reduction during provision of social support by romantic partners," Scandinavian Journal of Psychology, vol. 40, no. 1, pp. 205–209, 2019.

[43] T. R. de Brito and S. C. I. Pavarini, "The relationship between social support and functional capacity in elderly persons with cognitive alterations," Revista Latino-Americana de Enfermagem, vol. 20, no. 4, pp. 677–684, 2012.

[44] R. C. Sims, M. Hoseny, S. A. Levy, K. E. Whitfield, L. I. Katznel, and S. R. Waldstein, "Distinct functions of social support and cognitive function among older adults," Experimental Aging Research, vol. 40, no. 1, pp. 40–59, 2014.

[45] R. Pais, L. Ruano, C. Moreira, S. Fraga, O. P. Carvalho, and H. Barros, "Social support and cognitive impairment: results from a Portuguese 4-year prospective study," International Journal of Environmental Research and Public Health, vol. 18, no. 16, p. 8841, 2021.

[46] S. Ge, B. Wu, D. E. Bailey Jr., and X. Dong, "Social support, social strain, and cognitive function among community-dwelling U.S. Chinese older adults," The Journals of Gerontology. Series A, Biological Sciences and Medical Sciences, vol. 72, Supplement_1, pp. S16–S21, 2017.

[47] I. Kawachi and L. F. Berkman, "Social ties and mental health," Journal of Urban Health, vol. 78, no. 3, pp. 458–467, 2001.

[48] S. C. Pillemer and R. Holtzer, "The differential relationships of dimensions of perceived social support with cognitive function among older adults," Aging Mental Health, vol. 20, no. 7, pp. 727–735, 2016.

[49] A. López and M. Calero, "Predictors of cognitive decline in the elderly," Revista Española de Geriatría y Gerontología, vol. 44, no. 4, pp. 220–224, 2009.

[50] S. C. I. Pavarini, A. C. Ottaviani, A. G. Bregola et al., "Association between perceived social support and better cognitive performance among caregivers and non-caregivers," Revista Brasileira de Enfermamento, vol. 74, article e20200329, Supplement 2, 2021.

[51] O. Fernández-Lasarte, E. Ramos-Díaz, E. G. Palacios, and A. Rodríguez-Fernández, "Estudio comparativo entre educación superior y media: efecto del apoyo social percibido, el autoconcepto y la inteligencia emocional en el rendimiento académico," Educación XXI, vol. 22, no. 2, 2019.

[52] M. Richardson, C. Abraham, and R. Bond, "Psychological correlates of university students' academic performance: a systematic review and meta-analysis," Psychological Bulletin, vol. 138, no. 2, pp. 353–387, 2012.

[53] Y. López-Angulo, M. V. Pérez-Villalobos, R. C. Coborrendón, and A. E. Díaz-Mujica, "Apoyo social, sexo y área del conocimiento en el rendimiento académico autopercepcion de estudiantes universitarios chilenos," Formación Universitaria, vol. 13, no. 3, pp. 11–18, 2020.

[54] J. Granero-Molina, T. M. Matarin Jiménez, C. Ramos Rodríguez, J. M. Hernández-Padilla, A. M. Castro-Sánchez, and C. Fernández-Sola, "Social support for female sexual dysfunction in fibromyalgia," Clinical Nursing Research, vol. 27, no. 3, pp. 296–314, 2018.

[55] M. First, R. L. Spitzer, M. Gibbon, and J. B. W. Williams, "Entrevista Clínica Estructurada Para los Trastornos del Eje I del DSM-IV: SCID," Versión Clínica, Madsson, Barcelona, 1999.

[56] C. D. Spielberger, R. Gorsuch, and R. Lushene, Manual for the state-trait anxiety inventory, Consulting Psychologist, 1970.

[57] C. D. Spielberger, R. L. Gorsuch, and R. E. Lushene, Manual del Cuestionario de Ansiedad Estado/Rasgo (STAI), TEA Ediciones, 1982.

[58] A. T. Beck, C. H. Ward, M. Mendelson, J. Mock, and J. Erbaugh, "An inventory for measuring depression," Archives of General Psychiatry, vol. 4, no. 6, pp. 561–571, 1961.

[59] C. Vázquez and J. Sanz, "Fiabilidad y validez de la versión española del Inventario para la Depresión de Beck de 1978 en pacientes con trastornos psicológicos," Clínica y Salud, vol. 1, pp. 59–81, 1999.

[60] L. B. Krupp, N. G. LaRocca, J. Muir-Nash, and A. D. Steinberg, "The fatigue severity scale: Application to patients with multiple sclerosis and systemic lupus erythematosus," Archives of Neurology, vol. 46, no. 10, pp. 1121–1123, 1989.

[61] A. Bulbena, G. E. Berrios, and P. Fernández de Larrinoa, Medición clínica en psiquiatría y psicología, Masson, 2000.

[62] J. Bobes, M. P. González, P. A. Sáiz, M. T. Bascarán, C. Iglesias, and J. M. Fernández, "Propiedades psicométricas del cuestionario Oviedo de Sueño," Psicothema, vol. 1, pp. 107–112, 2000.

[63] R. Melzac, "The McGill pain questionnaire: major properties and scoring methods," Pain, vol. 1, no. 3, pp. 277–299, 1975.
[64] C. Lázaro, F. Bosch, R. Torrubia, and J. E. Baños, “The development of a Spanish questionnaire for assessing pain: preliminary data concerning reliability and validity,” *European Journal of Psychological Assessment*, vol. 10, no. 145-151, p. 53, 1994.

[65] A. Vaux, S. Riedel, and D. Stewart, “Modes of social support: the social support behaviors (SS-B) scale,” *American Journal of Community Psychology*, vol. 15, no. 2, pp. 209–232, 1987.

[66] B. Sandin and P. y Chorot, *Escala de Apoyo Social SS-B de Vaux, Riedel y Stewart*, UNED, 1990.

[67] B. A. Wilson, N. Alderman, P. W. Burgess, H. Emslie, and J. J. Evans, *Behavioural Assessment of the Dysexecutive Syndrome*, Thames Valley Test Company, 1996.

[68] M. L. Vargas, J. C. Sanz, and J. J. Marin, “Behavioral assessment of the dysexecutive syndrome battery (BADS) in schizophrenia: a pilot study in the Spanish population,” *Cognitive and Behavioral Neurology*, vol. 22, no. 2, pp. 95–100, 2009.

[69] M. J. Benedet, M. A. Alejandre, and A. Pamos, *Test de Aprendizaje Verbal España-Complutense: manual*, TEA Ediciones, 1998.

[70] S. Birnboim, “Strategy application test: discriminate validity studies,” *Revue Canadienne D’ergotherapie*, vol. 71, no. 1, pp. 47–55, 2004.

[71] D. C. Delis, E. Kaplan, and J. H. Kramer, *Delis-Kaplan Executive Function System (D-KEFS)*, The Psychological Corporation, 2001.

[72] J. J. Ibor, “The trail making tests A+B,” *Schizophrenia Research*, vol. 78, pp. 147–156, 2005.

[73] F. Faul, E. Erdfelder, A. Buchner, and A. G. Lang, “Statistical power analyses using G * Power 3.1: tests for correlation and regression analyses.” *Behavior Research Methods*, vol. 41, no. 4, pp. 1149–1160, 2009.

[74] S. Weiss, A. Winkelmann, and S. Duschek, “Recognition of facially expressed emotions in patients with fibromyalgia syndrome,” *Behavioral Medicine*, vol. 39, no. 4, pp. 146–154, 2013.

[75] S. Duschek, N. S. Werner, A. Winkelmann, and S. Wankner, “Implicit memory function in fibromyalgia syndrome,” *Behavioral Medicine*, vol. 39, no. 1, pp. 11–16, 2013.

[76] L. Roldán-Tapia, R. Cánovas-López, J. Cimadevilla, and M. Valverde, “Cognition and perception deficits in fibromyalgia and rheumatoid arthritis,” *Reumatología Clinica*, vol. 3, no. 3, pp. 101–109, 2007.

[77] P. Simos, G. Kitstaki, G. Dimitraki et al., “Cognitive deficits early in the course of rheumatoid arthritis,” *Journal of Clinical and Experimental Neuropsychology*, vol. 38, no. 7, pp. 820–829, 2016.

[78] P. H. Bolwijn, M. H. S. van Santen-Hoeufft, H. M. J. Baars, and S. van der Linden, “Social network characteristics in fibromyalgia or rheumatoid arthritis,” *Arthritis Care and Research*, vol. 7, no. 1, pp. 46–49, 1994.

[79] C. Martins, S. Campos, R. Martins, T. Moreira, C. Chaves, and M. Vieira, “Perception and impact of pain on patients with fibromyalgia,” *Atención Primaria*, vol. 1, no. 48, pp. 224–228, 2016.

[80] E. Briones-Vozmediano, E. Ronda-Pérez, and C. Vives-Cases, “Fibromyalgia patients perceptions of the impact of the disease in the workplace,” *Atención Primaria*, vol. 47, no. 4, pp. 205–212, 2015.

[81] M. Vicente-Herrero, M. J. Terradillos-Garcia, L. M. Capdevila-Garcia, M. V. Ramirez-ñiguez de la Torre, and A. A. López-Gonzalez, “Fibromyalgia and laboral activities on the Spain legislation,” *Revista Médica del Instituto Mexicano del Seguro Social*, vol. 49, no. 5, pp. 511–516, 2011.

[82] G. A. Reyes Del Paso, C. I. Montoro, and S. Duschek, “Reaction time, cerebral blood flow, and heart rate responses in fibromyalgia: evidence of alterations in attentional control,” *Journal of Clinical and Experimental Neuropsychology*, vol. 37, no. 4, pp. 414–428, 2015.

[83] W. E. Broadhead, S. H. Gehrlich, F. V. deGruy, and B. H. Kaplan, “Functional versus structural social support and health care utilization in a family medicine outpatient practice,” *Journal of Aging Science*, vol. 76, no. 7, pp. 221–233, 1989.

[84] L. Feldman, L. Goncalves, G. C. Chacon Puignau, J. Zaragoza, N. Bagés, and J. De Paulo, “Relaciones entre estrés académico, apoyo social, salud mental y rendimiento académico en estudiantes universitarios venezolanos,” *Universitas Psychologica*, vol. 7, no. 3, pp. 739–752, 2008.

[85] Z. Fan, X. Lv, L. Tu, M. Zhang, X. Yu, and H. Wang, “Reduced social activities and networks, but not social support, are associated with cognitive decline among older Chinese adults: a prospective study,” *Social Science & Medicine*, vol. 289, no. 1989, p. 114423, 2021.

[86] M. E. Kelly, H. Duff, S. Kelly et al., “The impact of social activities, social networks, social support and social relationships on the cognitive functioning of healthy older adults: a systematic review,” *Systematic Reviews*, vol. 6, no. 1, p. 259, 2017.

[87] K. Cotton, J. Verghese, and H. M. Blumen, “Grave matter volume covariance networks, social support, and cognition in older adults,” *The Journals of Gerontology. Series B, Psychological Sciences and Social Sciences*, vol. 75, no. 6, pp. 1219–1229, 2020.

[88] J. T. Caciquoppo and L. C. Hawkley, “Perceived social isolation and cognition,” *Trends in Cognitive Sciences*, vol. 13, no. 10, pp. 447–454, 2009.

[89] Y. Jang, E. Y. Choi, N. S. Park, D. A. Chiriboga, L. Duan, and M. T. Kim, “Cognitive health risks posed by social isolation and loneliness in older Korean Americans,” *BMC Geriatrics*, vol. 21, no. 1, p. 123, 2021.

[90] A. Hajek, S. K. Riedel-Heller, and H. H. König, “Perceived social isolation and cognitive functioning. Longitudinal findings based on the German ageing survey,” *International Journal of Geriatric Psychiatry*, vol. 35, no. 3, pp. 276–281, 2020.

[91] S. Okamoto and E. Kobayashi, “Social isolation and cognitive functioning: a quasi-experimental approach,” *The Journals of Gerontology. Series B, Psychological Sciences and Social Sciences*, vol. 76, no. 7, pp. 1441–1451, 2021.

[92] I. Evans, D. J. Llewellyn, F. E. Matthews, R. T. Woods, C. Brayne, and L. Clare, “Social isolation, cognitive reserve, and cognition in older people with depression and anxiety,” *Aging & Mental Health*, vol. 23, no. 12, pp. 1691–1700, 2019.

[93] J. Bellor and A. Wagner, “Disentangling loneliness: differential effects of subjective loneliness, network quality, network size, and living alone on physical, mental, and cognitive health,” *Journal of Aging and Health*, vol. 30, no. 4, pp. 521–539, 2018.

[94] P. M. Carrasco, D. P. Crespo, E. Pedrero-Pérez, and M. Montenegro-Peña, “Subjective cognitive decline: mental health, loneliness, pain and quality of life: population study,” *Journal of Aging Science*, vol. 8, no. 1, p. 218, 2020.
Y. Lu, C. Liu, S. Fawkes, J. Ma, Y. Liu, and D. Yu, “Inequality in social support associated with mild cognitive impairment: a cross-sectional study of older (≥60 years) residents in Shanghai, China,” *Frontiers in Public Health*, vol. 9, article 706322, 2021.

H. Gardener, B. Levin, J. DeRosa et al., “Social connectivity is related to mild cognitive impairment and dementia,” *Journal of Alzheimer’s Disease*, vol. 84, no. 4, pp. 1811–1820, 2021.

N. J. Donovan, O. I. Okereke, P. Vannini et al., “Association of higher cortical amyloid burden with loneliness in cognitively normal older adults,” *JAMA Psychiatry*, vol. 73, no. 12, pp. 1230–1237, 2016.

R. S. Wilson, K. R. Krueger, S. E. Arnold et al., “Loneliness and risk of Alzheimer disease,” *Archives of General Psychiatry*, vol. 64, no. 2, pp. 234–240, 2007.

Y. H. Hsiao, C. H. Chang, and P. W. Gean, “Impact of social relationships on Alzheimer’s memory impairment: mechanistic studies,” *Journal of Biomedical Science*, vol. 25, no. 1, p. 3, 2018.

D. H. Kang, L. Boss, and L. Clowitis, “Social support and cognition,” *Western Journal of Nursing Research*, vol. 38, no. 12, pp. 1639–1659, 2016.

J. Salinas, A. O’Donnell, D. J. Kojis et al., “Association of social support with brain volume and cognition,” *JAMA Network Open*, vol. 4, no. 8, article e2121122, 2021.

A. J. Gow, J. Corley, J. M. Starr, and I. J. Deary, “Which social network or support factors are associated with cognitive abilities in old age?,” *Gerontology*, vol. 59, no. 5, pp. 454–463, 2013.

P. Leung, M. Orrell, and V. Orgeta, “Social support group interventions in people with dementia and mild cognitive impairment: a systematic review of the literature,” *International Journal of Geriatric Psychiatry*, vol. 30, no. 1, pp. 1–9, 2015.

C. M. Galvez-Sánchez, C. Muñoz Ladrón de Guevara, C. I. Montoro, M. J. Fernández-Serrano, S. Duschek, and G. A. Reyes del Paso, “Cognitive deficits in fibromyalgia syndrome are associated with pain responses to low intensity pressure stimulation,” *PLoS One*, vol. 13, no. 8, article e0201488, 2018.

A. Feliu-Soler, X. Borràs, M. T. Peñarrubia-Maria et al., “Cost-utility and biological underpinnings of mindfulness-based stress reduction (MBSR) versus a psychoeducational programme (FibroQoL) for fibromyalgia: a 12-month randomised controlled trial (EUDAIMON study),” *BMC Complementary and Alternative Medicine*, vol. 16, no. 1, p. 81, 2016.

K. Oliver, T. A. Cronan, H. R. Walen, and M. Tomita, “Effects of social support and education on health care costs for patients with fibromyalgia,” *The Journal of Rheumatology*, vol. 28, no. 12, pp. 2711–2719, 2001.

G. Reig-Garcia, C. Bosch-Farré, R. Suñer-Soler et al., “The impact of a peer social support network from the perspective of women with fibromyalgia: a qualitative study,” *International Journal of Environmental Research and Public Health*, vol. 18, no. 23, p. 12801, 2021.