Reumatismo, 2012: 64 (1): 27-34

Preliminary identification of key clinical domains for outcome evaluation in fibromyalgia using the Delphi method: the Italian experience

F. Salaffi¹, A. Ciapetti¹, P. Sarzi Puttini², F. Atzeni², C. Iannuccelli³, M. Di Franco³, M. Cazzola⁴, L. Bazzichi⁵

¹Clinica Reumatologica, Università Politecnica delle Marche, Ancona; ²Unità di Reumatologia, Azienda Ospedaliera Polo Universitario "L. Sacco", Milano; ³Dipartimento di Medicina Interna, Reumatologia, Università "La Sapienza" di Roma; ⁴Unità Operativa Medicina Riabilitativa, "Ospedale di Circolo", Busto Arsizio, Presidio di Saronno, Varese; Dipartimento di Medicina Interna Divisione di Reumatologia, Ospedale "S. Chiara", Università di Pisa, Italy

SUMMARY

Objective: Fibromyalgia (FM) is a complex syndrome that, in Italy, affects at least 2% of the adult population. It is characterized by chronic widespread musculoskeletal pain often accompanied by multiple other symptoms. The aim of this study was to identify a set of clinical domains for FM considered relevant by both clinicians and patients using a consensus process.

Methods: Consensus was achieved using the Delphi method based on questionnaires and systematic, controlled opinion feedback. The Delphi exercise involved a panel of 252 rheumatologists and 86 patients with FM as defined by the American College of Rheumatology criteria. All of the patients and clinicians were asked to rank the relative different domains of FM in order of priority. The content validity index (CVI) was used to establish the percentage agreement. The importance of each item was ranked on a 0-3 Likert scale. The frequency, mean relevance scores, and frequency importance product were also calculated.

Results: The Delphi exercise showed that the domains ranked highest by patients were similar to those of the clinicians, with the exception of tender point intensity (considered relevant by the clinicians but not by the patients) and environmental sensitivity (considered important by the patients but not by the clinicians). A final 8-item model was developed which was considered to demonstrate adequate validity.

Conclusions: The Delphi exercises identified and ranked relevant key clinical domains that need to be assessed in FM research. On the basis of these results, a new patient-reported composite outcome index can be developed and used in clinical trials.

Key words: fibromyalgia, Delphi method, pain scales, health-related quality of life, patient-reported outcomes.

Reumatismo, 2012; 64 (1): 28-35

INTRODUCTION

libromyalgia (FM) is a complex syndrome which, in Italy, affects at least 2% of the adult population (1). It is characterized by widespread chronic pain, often associated with other symptoms such as fatigue, disturbed sleep patterns, functional limitations and cognitive impairment (2, 3). The presence of these symptoms and any comorbidity (4) significantly compromises the quality of life of FM patients (5-8). Defining the impact of symptoms on patients with FM can be a useful tool to evaluate patient activity and and social function, and response to treatment in research and daily clinical practice (9-14).

The group of experts in fibromyalgia of the OMERACT (Outcome Measures in Rheumatology) group, using working party and the Delphi methods, has recently examined the possibility of patients identifying principle clinical domains to be used as an evaluation tool in clinical trials (2, 15). Beginning with an initial list of 40 potential domains, the experts gave marks in terms of importance of each domain. In the same way, a similar selection was made by dif-

Corresponding author: Dott. Fausto Salaffi Clinica Reumatologica Università Politecnica delle Marche Ospedale "C. Urbani" Via dei Colli 52 - 60035 Jesi, Ancona, Italy E-mail: fsalaff@tin.it

ferent groups of patients with FM from different centers. The results from both the groups of experts and of patients were the same. The list of principal domains selected included: pain, overall state of health, fatigue, quality of life (QoL), multidimensional function, sleep pattern and depression. The so-called secondary domains included physical function, tender points, cognitive impairment, (lack of concentration, disorganized thoughts, etc.) and anxiety (15). This working method allowed us to sustain construct validity of the potential domains, while feasibility and the selective ability of the specific instruments to be used in its application were the subject of a separate systematic randomized controlled trial (RCT) (12, 16).

Use of the so-called patient reported outcomes (PRO) is the most accepted method of evaluating disease status, especially in conditions characterized by chronic pain (17, 18). This method is highly appropriate for use in FM (10) given that most of the clinical domains considered essential for evaluation of the disease, and defined as such by the experts of the OMERACT group (2, 15, 19) and of the Food and Drug Administration (FDA) (20), are patient centered.

In line with methods used by the OMER-ACT group (15, 20), this study was carried out using the Delphi method, and involved 252 rheumatologists and 86 patients with FM. The principle aim of the study was to reach a unanimous consensus on the list of health domains considered to be high priority both in a research context and in daily clinical practice, and considered to be useful for disease evaluation.

■ MATERIALS AND METHODS

Expert working parties

The items were generated in two successive phases. First, we carried out a literature review of clinical studies in FM in order to identify the principle outcome measures used. Clinical trials published up to September 2010 were downloaded from the internet using MEDLINE, CINAHL

and EMBASE search engines. A strategy of specific research was developed for each database using Cochrane filter for RCT and including articles published in English and in Italian between 1990 and 2009. The key words used were: "Fibromyalgia", "Chronic pain syndrome", "Widespread pain", "State of Health", "Multidisciplinary", "Patient care team", "Treatment of fibromyalgia" or "Management of fibomyalgia" and "Trial".

The search strategy produced 2,055 articles, of which 204 were selected on the basis of titles, summaries and key words, from which a total of 39 papers were subsequently selected. The list of items was re-examined for importance of content by a multidisciplinary task force of experts in FM: 25 rheumatologists, 4 orthopedic surgeons, 3 neurologists, 3 algologists, 2 psychologists, 6 physiotherapists and 3 occupational therapists. Each doctor was invited to decide which specific method they considered most appropriate to measure each clinical domain and to mark each item selected according to feasibility. In order to reduce the number of clinical domains. items were excluded from the list if:

- a) they were related to gender;
- b) they required use of special equipment;
- c) they used terminology which was ambiguous or difficult to understand;
- d) they presented alternatives to other items, duplicated them and/or were similar to them. The final list was made up of a pool of 58 symptoms/domains (see Appendix).

Final item selection

The number of items was reduced in order to avoid symptoms and/domains which were superfluous or repetetive. This was achieved by keeping 8-10 symptoms/domains considered to be the most important and representative of patients' general state of health. In the next phase, 252 rheumatologists, randomly selected from the list of 1,200 members of the Società Italiana di Reumatologia (Italian Society of Rheumatology) were interviewed through the internet (http://www.reumatologia.it).

Each rheumatologist was invited to make

their own independent selection, without any assistance, of the symptoms/domains they considered to be the most important and representative from the list of 58 symptoms/domains which had been identified by the task force. Response was 70.4% (252 of 538 contacted) representing a third of all rheumatologists interviewed. The content validity index (CVI) was used to establish the proportion/percentage of agreement between the experts. Lynn (21) recommended using a relevance evaluation scale for each item providing ordinal level data based on the Likert scale from 0 to 4 (4=highly relevant, extremely important; 3=very relevant, very important; 2=not very relevant, not very important; 1=not relevant, unimportant). Once scoring was completed, the CVI only included elements with a mark of 3 or 4, while all descriptors with a mark of 1 and 2 were not considered. The CVI formula was: CVI or percentage of agreement = the number of experts in agreement on items receiving scores of 3 or 4 divided by the total number of experts. The items were considered to have adequate content validity when agreement was 70% or over, and to be questionable or unacceptable when agreement among experts was 60-70% or less than 60%, respectively.

Patient working parties

The aim of this phase of the study was to reach unanimous agreement, in terms of priority and importance of the clinical domains, among patients diagnosed with FM according to the criteria of the American College of Rheumatology (ACR) (22). Rheumatologists involved in the study

excluded diagnoses other than FM. Other exclusion criteria were: comorbidities such as to impede full participation in study procedures (terminal stage, such as, for example, advanced stage renal disease, cardiac insufficiency or neoplasms), alcohol abuse, significant cognitive impairment, or psychiatric symptoms which could have compromised efficient completion of the questionnaire.

After obtaining local ethics committee approval for the study, a group of 86 patients with FM were invited to take part and gave their signed informed consent. These patients were from three different rheumatology centers: the centers of northern, central and southern Italy. The patient group was considered to provide an adequate representative sample of the disease in terms of heterogeneity of the symptoms presented at the moment of inclusion in the study.

A rheumatologist experienced in developing measurement strategies presented each patient with the list of the 42 domains that were considered to be the most important in relation to FM. During the interview, each patient was asked to put the various domains in the order of priority (mean importance, MI) giving each one a score on the Likert scale from 1 to 3 (1=not relevant, not important; 2=not very relevant, not very important; 3=very relevant, very important). Mean values were then calculated for each item which had obtained a score of 2. Analysis considered frequency of each symptom, and the prevalence and domains which satisfied 60% or over of prevalence criteria. The frequency importance product (FIP) was, therefore, generated for each

Table I - Domains classified according to importance by clinicians.

Domain	Frequency	Mean importance (MI)	Frequency importance product (FIP)
1. Pain	79.0	2.85	225.2
2. Fatigue	78.6	2.65	208.3
3. Disturbed sleep patterns	74.6	2.70	201.4
4. Health-related quality of life (HRQL)	73.8	2.60	191.9
5. Depression	70.2	2.60	182.5
6. Anxiety	71.8	2.50	179.5
7. Memory lapse and lack of concentration	69.1	2.40	165.8
8. Tender points	60.2	2.20	126.4

item, multiplying the frequency by the MI assigned to each domain. For example, the FIP relating to the domain *pain*, estimated by clinicians to be the most frequent and with the highest range, was 225.2 (FIP *pain* = frequency 79% x MI 2.85 = 225.2) (Table I). The FIP represented, therefore, a composite, two-dimensional measurement particularly useful in establishing the extent of importance of a symptom or domain.

■ RESULTS

Use of the Delphi method among clinicians

Using the Delphi method, 252 rheumatologists evaluated the importance of each item. The items considered important (pain, tiredness, disturbed sleep patterns, quality of life, physical function, depression, cognitive impairment, anxiety and tender points) which reached a 60.2-79.0% agreement among clinicians were included in the final list. Table I shows the frequency, relevance and FIP of each item. Clinicians considered pain (FIP = 225.5), tiredness (FIP = 208.3), disturbed sleep patterns (FIP = 201.4) and quality of life (FIP = 191.9) to be the most important items.

Use of the Delphi method among patients The study included 86 patients: 70 women and 16 men, mean age 43.9 years, range 22-68 years, female:male ratio 5:1, comparable and similar to other studies in the literature. The largest proportion of patients (58%) had been diagnosed with FM 1-4 years before the study began. Seventyone percent of patients reported the onset of symptoms to have been over five years before, 35% did not work because of the disease, while 42% classified their state of health as "poor". Their level of education was generally low: 60.1% had only received primary school education (6-11 years of age) and only 19.9% had gone to upper secondary school (14-19 years of age). Most patients (77.9%) were married and lived with their family while 34% were housewives. Forty-one of the 86 subjects (52.6%) reported one or more comorbidity, in particular, cardiovascular disease (29.2%), and respiratory (14.1%) and metabolic (10.5%) problems.

Overall, classification of the domains selected by patients was the same as that proposed by the clinicians. *Pain* was the most important symptom (FIP = 282.2), while patients considered *tiredness*, *disturbance of sleep pattern*, *overall state of health*, *quality of life*, *depression*, *anxiety* and *cognitive impairment* (memory lapse and lack of concentration) to be equally important. *Tender points* were considered to be important by clinicians but not by patients. On the other hand, *excessive sensitivity to external stimuli* (sounds, lights,

Tabella II - Domains classified according to importance by patients.

Domain	Items	Frequency	Mean importance (MI)	Frequency importance product (FIP)
1. Pain	Pain or physical distress; painful joints; pain on palpation	97.3	2.9	282.2
2. Fatigue	Tiredness; low energy levels	93.6	2.7	252.7
3. Quality of sleep	Difficulty in sleeping; insomnia; frequent waking during the night	90.1	2.6	234.3
4. Multidimensional function	Difficulting in moving around, in walking or doing physical exercises; problems in carrying out daily routine activities, work and study, and impact on daily routine	89.8	2.6	233.5
5. Depression	Feeling sad, unmotivated, pessimistic, isolated, listless	81.5	2.4	195.6
6. Sensitivity to external stimuli	Sensitivity to sounds, light, smells and/or the cold	78.3	2.4	187.9
7. Anxiety	Feeling frustrated, worried, scared	76.1	2.3	175.0
8. Cognitive Memory lapse or difficulties in remembering things; loss of impairment memory; lack of concentration		74.7	2.1	156.9

smells) were considered to be important by patients but not by clinicians. Patients did not take into consideration the domain health-related quality of life while the rheumatologists did. In spite of this, the impact of the disease on physical function was, however, considered to be important. In particular, reference was made to difficulty in moving around, walking or doing physical exercises, problems in carrying out daily routine activities, work and study, and the impact of the disease on daily life. The 8 items reached a prevalence of at least 70%, while the mean importance (MI) of marks given varied between 2.1 (cognitive impairment) and 2.9 (pain). The FIP of the 8 items was between 156.9 and 282.2. Clinicians summarized the last domain as cognitive impairment while patients described it in different terms such as, for example, "losing attention", "lack of concentration", "memory lapse" and "disorganized thoughts". The domains with the highest marks are shown in Table II.

DISCUSSION

In their definition of the classification of the 8 items considered to be important during the course of FM, a high percentage of doctors and patients generally gave pain, tiredness and disturbed sleep patterns the highest marks of importance. (23) This, therefore, confirms the importance of these domains, and suggests they should be included in evaluation criteria for this disease (19). It is interesting to note that these items represent the three domains which make up the "Fibromialgia Activity Score (FAS)" (24) self-evaluation scale, and as such are in agreement with recommendations proposed by the OMERACT group of experts in FM, the "Syndrome Workshop" (15, 19) and the IMMPACT (Initiative on Methods, Measurements and Pain Assessment in Clinical Trial) group (25). The other five domains selected by clinicians, including depression and anxiety, state of health-quality of life, memory lapse and lack of concentration, and tender points, are generally considered to be

important also by patients (15, 19). However, with respect to clinicians, patients also consider excessive sensitivity to external stimuli to be important. The most important functional items, which were underlined by the patients, include effects of the disease on personal day to day planning, achieving routine objectives and completing daily tasks, even though they did not usually consider health-related state of health to be important. According to the OMERACT group of experts (15, 19). clinicians should consider health-related quality of life to be very important in order to adequately evaluate the much wider multidimensional aspect of function rather than simply physical function. It is interesting to note that while rigidity was considered to be an important domain by the OMERACT group of experts (2,.15, 19). by systematic revision (12,26) and in investigations conducted via internet (27). this was not among the domains selected by patients who took part in the study, who did not consider it to be important. In contrast, it is not surprising that pain was the domain selected most, often being the first symptom reported by the patient. However, even though it was classified in first place as an indicator of disease severity, pain is considered the principle criterion by 79% of the doctors interviewed. It should also be noted that an objective clinical evaluation made through investigation into tender points was reported by only 60.2% of clinicians and that all other criteria selected are to be considered "patient-centered". The variety of criteria available for FM, and the lack of a unanimous consensus agreement, underlines the need to validate an adequate means to measure disease severity. A recent study (8) which involved 788 patients with FM, defined pain and tiredness to be the most important symptoms in 54% of participants and to be the most characteristic signs of disease, while fatigue was chosen as the most important item by 28% of those interviewed. Tiredness, a common symptom among patients with FM (2, 15, 27) which often represents one of the most worrying problems, is a subjective experience which is described as "extreme and persistent mental and/or physical weakness or tiredness". In spite of the fact that the causes of tiredness are little known, it has been increasingly used in recent years as a measurement of outcome in clinical studies (28). Since there is still no theoretical model, it remains difficult to define tiredness in FM. Along with pain and tiredness, disturbed sleep patterns is constantly referred to as important by patients and doctors (27, 29-33). These play a primary role in the disease course. Estimates of percentage of patients with FM affected by disturbed sleep patterns vary from 74 to 95-99% (34). It has also been shown that disturbed sleep patterns have a significant impact on amplifying pain and on the consequent negative impact on function and psychological-emotional well-being (32-34). It is not surprising, therefore, that both patients and clinicians underline the importance of improving the quality of sleep, together with treating pain.

Given the multidimensional characteristic of FM and the reported validity and usefulness of PRO in evaluating and monitoring these patients, we have recently developed the fibromyalgia assessment status (FAS) questionnaire which combines self-evaluation of the degree of tiredness and pain, and quality of sleep (on the basis of sites listed in the self-assessment pain scale. SAPS) into a single measurement, with scores ranging from 0 to 10 (23). This index includes the first 3 items considered to be important by both clinicians and patients, and can be used to evaluate disease severity and define response to treatment. In conclusion, this study using the Delphi method of evaluation has shown that the domains considered to be the most important by patients are similar to those indicated by clinicians, with the exception of tender points (which were not considered to be important by patients). The final model of 8 items was shown to be have adequate construct validity.

REFERENCES

 Salaffi F, De Angelis R, Grassi W. MArche Pain Prevalence; INvestigation Group (MAP-

- PING) study: Prevalence of musculoskeletal conditions in an Italian population sample: results of a regional community-based study. I. The MAPPING study. Clin Exp Rheumatol. 2005; 23: 819-28.
- Mease P. Fibromyalgia syndrome: review of clinical presentation, pathogenesis, outcome measures, and treatment. J Rheumatol. 2005; 75: 6-21.
- 3. Rehm SE, Koroschetz J, Gockel U, et al. A cross-sectional survey of 3035 patients with fibromyalgia: subgroups of patients with typical comorbidities and sensory symptom profiles. Rheumatology. 2010; 49: 1146-52.
- McBeth J, Symmons DP, Silman AJ, et al. Musculoskeletal pain is associated with a long-term increased risk of cancer and cardiovascular-related mortality. Rheumatology. 2009; 48: 74-7.
- Mease P, Arnold LM, Bennett R, et al. Fibromyalgia syndrome. J Rheumatol. 2007; 34: 1415-25.
- Salaffi F, Sarzi-Puttini P, Girolimetti R, et al. Health related quality of life in fibromyalgia patients: a comparison with rheumatoid arthritis patients and the general population using the SF-36 health survey. Clin Exp Rheumatol. 2009; 27: S67-74.
- Salaffi F, De Angelis R, Stancati A, Grassi W. MArche Pain; Prevalence INvestigation Group (MAPPING) study. Health-related quality of life in multiple musculoskeletal conditions: a cross sectional population based epidemiological study. II. The MAP-PING study. Clin Exp Rheumatol. 2005; 23: 829-39.
- 8. van Wilgen CP, van Ittersum MW, Kaptein AA, van Wijhe M. Illness perceptions in patients with fibromyalgia and their relationship to quality of life and catastrophizing. Arthritis Rheum. 2008; 58: 3618-26.
- Bennett RM, Jon Russell J, Cappelleri JC, et al. Identification of symptom and functional domains that fibromyalgia patients would like to see improved: a cluster analysis. BMC Musculoskeletal Disorders. 2010; 11: 134.
- Scascighini L, Toma V, Dober-Spielmann S, Sprott H. Multidisciplinary treatment for chronic pain: a systematic review of interventions and outcomes. Rheumatology. 2008;47:670-8.
- Salaffi F, Sarzi-Puttini P, Ciapetti A, Atzeni F. Assessment instruments for patients with fibromyalgia: properties, applications and interpretation. Clin Exp Rheumatol. 2009; 27: S92-105.
- Carville SF, Choy EHS. Systematic review of discriminating power of outcome measures used in clinical trials of fibromyalgia. J Rheumatol. 2008; 35: 2094-105.
- 13. Sarzi-Puttini P, Buskila D, Carrabba M, et al. Treatment strategy in fibromyalgia syn-

- drome: where are we now? Semin Arthritis Rheum. 2008; 37: 353-65.
- Carville SF, Arendt-Nielsen S, Bliddal H, et al. EULAR evidence-based recommendations for the management of fibromyalgia syndrome. Ann Rheum Dis. 2008; 67: 536-41.
- Mease PJ, Arnold LM, Crofford LJ, et al. Identifying the clinical domains of fibromyalgia: contributions from clinician and patient Delphi exercises. Arthritis Rheum. 2008; 59: 952-60.
- Choy EH, Arnold LM, Clauw DJ, et al. Content and criterion validity of the preliminary core dataset for clinical trials in fibromyalgia syndrome. J Rheumatol. 2009; 36:.2330-4.
- 17. Salaffi F, Migliore A, Scarpellini M, et al. Psychometric properties of an index of three patient reported outcome (PRO) measures, termed the CLinical ARthritis Activity (PRO-CLARA) in patients with rheumatoid arthritis. The NEW INDICES study. Clin Exp Rheumatol. 2010; 28: 186-200.
- Pincus T, Chung C, Segurado OG, et al. An index of patient reported outcomes (PRO-Index) discriminates effectively between active and control treatment in 4 clinical trials of adalimumab in rheumatoid arthritis. J Rheumatol. 2006; 33: 2146-52.
- Mease P, Arnold LM, Choy EH, et al. OMER-ACT Fibromyalgia Working Group. Fibromyalgia syndrome module at OMERACT 9: domain construct. J Rheumatol. 2009; 36: 2318-29.
- 20. US Department of Health and Human Services, Food and Drug Administration, Center for Drug Evaluation and Research (CDER), Center for Biologics Evaluation and Research (CBER), and Center for Devices and Radiological Health (CDRH). Draft guidance for industry: patient-reported out come measures. Use in medical product development to support labeling claims. 2006. Available from: http://www.fda.gov/cder/guidance/5460dft.pdf
- Lynn MR. Determination and quantification of content validity. Nursing Res. 1986; 35: 382-5.
- 22. Wolfe F, Smythe HA, Yunus MB, et al. The American College of Rheumatology 1990 criteria for the classification of fibromyalgia:

- report of the Multicenter Criteria Committee. Arthritis Rheum. 1990: 33: 160-7.
- Silverman S, Sadosky A, Evans C, et al. Toward characterization and definition of fibromyalgia severity. BMC Musculoskeletal Disorders. 2010; 11: 66.
- 24. Salaffi F, Sarzi-Puttini P, Girolimetti R, et al. Development and validation of the self-administered Fibromyalgia Assessment Status: a disease-specific composite measure for evaluating treatment effect. Arthritis Res Ther. 2009; 11: R125.
- Dworkin RH, Turk DC, Wyrwich KW, et al. Interpreting the clinical importance of treatment outcomes in chronic pain clinical trials: IMMPACT recommendations. J Pain. 2008; 9: 105-21.
- Choy EH, Mease PJ. Key symptom domains to be assessed in fibromyalgia (outcome measures in rheumatoid arthritis clinical trials). Rheum Dis Clin North Am. 2009; 35: 329-37.
- Bennett RM, Jones J, Turk DC, Matallana L. An internet survey of 2,596 people with fibromyalgia. BMC Musculoskelet Disord. 2007; 8: 27.
- 28. Norheim KB, Jonsson G, Omdal R. Biological mechanisms of chronic fatigue. Rheumatology. 2011; 50: 1009-18.
- Arnold LM, Crofford LJ, Mease PJ, et al. Patient perspectives on the impact of fibromyalgia. Patient Educ Couns. 2008; 73: 114-20.
- Hallberg LRM, Carlsson SG. Coping with fibromyalgia. A qualitative study. Scand J Caring Sci. 2000; 14: 29-36.
- 31. Cunningham MM, Jillings C. Individuals' descriptions of living with fibromyalgia. Clin Nurs Res. 2006; 15: 258-73.
- 32. Martin S, Chandran A, Zografos L, Zzateva G. Evaluation of the impact of fibromyalgia on patients' sleep and the content validity of two sleep scales. Health Qual Life Outcomes. 2009; 10: 64.
- 33. Bigatti SM, Hernandez AM, Cronan TA, Rand KL. Sleep disturbances in fibromyalgia syndrome: Relationship to pain and depression. Arthritis Rheum. 2008; 59: 961-7.
- 34. Theadom A, Cropley M, Humphrey KL. Exploring the role of sleep and coping in quality of life in fibromyalgia. J Psychosom Res. 2007; 62: 145-51.



Appendix								
Domains	presented i	in the	Delphi	study				

- 1. Low back pain
- 2. Irritable bowel syndrome
- 3. Restless leg syndrome
- 4. Tinnitus
- 5. Cognitive impairment
- 6. Morning rigidity
- 7. Fatique
- 8. Widespread chronic pain
- 9. Tender point
- 10. Multidimensional function
- 11. Memory lapse
- 12. Tingling extremities
- 13. Constipation
- 14. Dysmenorrhea
- 15. Sjogren's syndrome
- 16. Ravnaud's phenonemon
- 17. Physical weakness
- 18. Widespread paresthesias
- 19. Temporomandibular joint dysfunction
- 20. Losing attention or lack of concentration (e.g. difficulties in concentrating and following train of thought, "fibro-fog")
- 21. Difficulty in moving around, walking or doing physical exercises
- 22. Unable to do physical exercise
- Depression (e.g. dissatisfaction, sadness, resignation or selfeffacement, unmotivated)
- 24. Unmotivated
- 25. Disorganized thoughts (difficulty in expressing oneself, in prompt verbal response or in making personal programs and plans)
- 26. Numbness or tingling in the fingers and toes
- 27. Cephalea or migraine
- 28. Sensitivity to environmental factors (e.g. smells, sounds, lights, changes in temperature)
- 29. Skin sensitivity

- 30. Involuntary movement
- 31. Vertigo
- 32. Impact on sleep (e.g. difficulty in falling asleep, insomnia, frequent waking during the night)
- 33. Limiting/compromising normal daily routine activities
- 34. Impact on social life (e.g. less time for hobbies or free time, problems in traveling)
- 35. Problems taking medication (e.g. side effects or intolerance)
- Impact on family life (e.g. relationship problems with partner or children)
- 37. Impact on work/study
- 38. Lack of or low sex drive
- 39. Economic impact (e.g., cost of medication, health insurance)
- 40. Problems driving (e.g. pain or fatigue or memory lapse)
- 41. Anger or irritability
- 42. Frustration (e.g. due to function limitations, due to social relationships)
- 43. Anxiety (e.g. nervousness)
- 44. Panic attacks
- 45. Chest pain
- 46. Abdominal pain or cramps
- 47. Urinary incontinence
- 48. Strangury, dysuria, difficulties in urinating
- 49. Shortness of breath
- 50. Weight loss (<10 kg)
- 51. Loss of appetite
- 52. Loss of smell, taste or flavor
- 53. Difficulties in urinating
- 54. Gynecological problems (women)
- 55. Muscle pain, pain or cramps
- 56. Cold hands
- 57. Swollen hands
- 58. Swelling of other joints