Fibromyalgia (FM) is a common, potentially disabling, chronic disorder that is defined by widespread pain, often accompanied by fatigue and sleep disturbance, and associated with other symptoms including depression, cognitive dysfunction (e.g. forgetfulness, decreased concentration), irritable bowel syndrome (IBS) and headache (1,2). In the general population, the estimated global prevalence of FM is 2.7% (4.2% female, 1.4% male) (2). In primary care, studies suggest that up to one in 20 patients has FM symptoms (3), and this number is increasing as growing recognition of FM by patients leads to an upsurge in presentation for diagnosis and treatment (4,5). The cause of FM is not known, but research studies suggest genetic predisposition and possible triggering events (6).

Fibromyalgia continues to present a challenge for healthcare professionals (HCPs) (7). The extensive array of symptoms associated with, and gradual evolution of, FM make it difficult to diagnose in primary care settings (7,8), and the condition is often under-diagnosed (5). One study has shown that diagnosis of FM might take more than 2 years, with patients seeing an average of 3.7 different physicians during this time (8). Although the American College of Rheumatology (ACR) has published diagnostic criteria for FM (9,10), these are not widely used in clinical practice, and there remains a knowledge gap among some HCPs, particularly in the primary care setting (7,8,11,12). In addition to diagnostic complexity, therapeutic management might be problematic (13), and there is a lack of prescribing consistency between physicians (14,15). Many patients might not receive treatment, and for those...
who do, repeated therapy switching, polypharmacy and discontinuation are common (16). Some patients may also have unrealistic treatment expectations (17) and difficulty coping with their symptoms, which may contribute to struggles in managing their condition.

The aim of this review was to discuss the current and evolving understanding of FM, provide insights into the challenges around recognition and diagnosis, and recommend improvements in the management and treatment of FM. The review will highlight the role of the primary care physician, and the place of the medical home in FM management.

**Methods**

We reviewed the epidemiology, pathophysiology and management of FM by searching English-language publications in PubMed, and references from relevant articles, published before May 2015. The main search terms were fibromyalgia, epidemiology, pathophysiology, diagnosis, primary care, secondary care, treatment and patient-centred medical home. We selected articles on the basis of quality (robust data published in a peer-reviewed journal that were able to support the conclusions drawn), relevance to the illness and importance in illustrating current management pathways and the potential for future improvements.

**FM overview**

Although the global prevalence of FM is estimated to be 2.7%, epidemiological studies have produced varying results across different countries and continents (2). Until recently, most studies were carried out using the 1990 ACR diagnostic criteria (1), which resulted in notable gender imbalance; using these criteria, the prevalence of FM was 3.4% in females, and 0.5% in males (a ratio of 7 : 1) (18). This might be because the 1990 criteria required pain to be present on palpation of at least 11 of 18 tender points for a diagnosis of FM to be confirmed (Table 1) (1), and males have a higher pressure pain threshold than females (19), making them less likely to meet the 1990 FM criteria (5). A recent analysis using the updated 2010 criteria (9) that do not require a tender point assessment, has provided prevalence estimates of 7.7% in women and 4.9% in men (20), narrowing the gender gap and giving a female:male ratio of 1.6 : 1, which is more similar to that seen in other chronic pain conditions (6).

While many potential mechanisms for FM have been evaluated, recent evidence suggests that dysfunction in central nervous system pain processing mechanisms including central sensitisation or central augmentation of pain contribute to the development of chronic pain in patients with FM (21,22). This results in the ‘volume control’ for pain being turned up (4), and patients experience allodynia (a height-

**Table 1** Differences between the ACR 1990 (1) and the revised ACR 2010 (9) criteria for FM

<table>
<thead>
<tr>
<th>1990</th>
<th>2010</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>History of widespread pain</strong></td>
<td>WPI ≥ 7 and SS ≥ 5 OR WPI 3–6 and SS ≥ 9</td>
</tr>
<tr>
<td><strong>Pain of ≥ 3 months’ duration</strong></td>
<td>Symptoms have been present at a similar level for ≥ 3 months Patient does not have a disorder that would otherwise explain the pain</td>
</tr>
<tr>
<td><strong>Pain in 11 of 18 tender points on digital palpation</strong></td>
<td><strong>WPI score</strong> The number of areas in which patient has had pain over the last week (six lower extremities, six upper extremities, seven axial skeleton) Final score: between 0 and 19</td>
</tr>
<tr>
<td><strong>Definitions</strong></td>
<td><strong>SS score</strong> The sum of severity of fatigue, waking unrefreshed and cognitive symptoms, plus the severity of general somatic symptoms Each symptom is rated on a scale of 0–3, where 0 = no symptoms/problem and 3 = severe symptoms/problems Final score: between 0 and 12</td>
</tr>
<tr>
<td><strong>Widespread pain</strong></td>
<td></td>
</tr>
<tr>
<td>• Pain on left side of body, right side of body, above waist, below waist and axial skeletal pain</td>
<td></td>
</tr>
<tr>
<td><strong>Tender points (all bilateral)</strong></td>
<td></td>
</tr>
<tr>
<td>• Occiput, low cervical, trapezius, supraspinatus, second rib, lateral epicondyle, gluteal, greater trochanter, knee</td>
<td></td>
</tr>
</tbody>
</table>

ACR, American College of Rheumatology; FM, fibromyalgia; SS, symptom severity; WPI, Widespread Pain Index.
Barriers to managing FM in primary care

Although our understanding of FM has increased considerably in recent years, the barriers to diagnosis and optimal treatment are many and varied. Globally, there are inconsistencies in the recognition of symptoms, and in the validity of FM as a diagnosis (13). Even where guidelines are available, physicians in different regions may have varying levels of awareness of these guidelines. This, in turn, results in wide variations in the time to diagnosis of FM between geographical regions (ranging from 2.6 to 5 years in the USA, Latin America and Europe) (5,30).

In addition to diagnostic barriers, there are major inconsistencies between treatment practices. There is still some debate over the optimal choice and sequence of treatments for FM (31), and the approval status, availability and reimbursement of therapeutic agents varies between countries (32). Treatment guidelines currently make varying recommendations, possibly because of different criteria used to grade recommendations (33), and there might also be cultural differences regarding patient treatment expectations (e.g. ethnic variance in the level of pain perception) (30). Furthermore, prescribing practices might differ according to whether a patient is seen by a primary care physician or a specialist, on the HCP’s familiarity with treatment guidelines, and on the availability of local resources for disease management.

Finally, the lack of a clear patient pathway and healthcare system for diagnosis and management of FM often results in patients being passed from physician to physician, receiving multiple drugs to treat different symptoms and suffering increased disability (12,30,34). Many primary care physicians still prefer to refer the patient to a specialist (7), particularly when patients have multiple comorbidities that are likely to require a considerable amount of time to investigate and manage. However, the majority of FM cases could be diagnosed and treated in primary care, and a patient-centric multidisciplinary approach to FM in primary care would result in more rapid diagnosis, more effective management, improved outcomes for patients and better use of health resources (4,35).

Unmet needs

Despite improvements in the understanding of the condition, FM remains under-diagnosed and undertreated. A large proportion of physicians, particularly in primary care, report unclear diagnostic criteria, a lack of confidence in using the ACR criteria for diagnosis, insufficient training/skill in diagnosing FM and a lack of knowledge of treatment options (7,11,13). Furthermore, both patients and physicians express dissatisfaction with the delays in reaching a diagnosis and obtaining effective treatment (12). Several surveys of patients with FM have reported dissatisfaction with FM medication and overall treatment (8,16,36). A survey of 800 patients reported that 35% believed that their chronic, widespread pain was not well managed by their current treatment, and 22% were not satisfied with the impact of their treatment on fatigue (8).

Diagnosis of FM

Fibromyalgia is a disease with unique clinical characteristics, making it suitable for diagnosis in the primary care setting. Prompt diagnosis of the disorder is an essential component of successful FM management (18). Studies have shown that a diagnosis of FM is associated with improved satisfaction with health, and a reduction in the utilisation of medical resources and the associated costs (in particular, a reduction in referrals and investigations), relative to patients with FM symptoms who remain undiagnosed (37,38).
Fibromyalgia management for primary care providers

**ACR criteria**
The first ACR criteria for FM, published in 1990 (Table 1) (1,39), were intended mainly for research classification, and were not intended to be used in clinical practice (6). Although commonly cited in the literature, the 1990 ACR criteria were not widely used by primary care physicians, possibly owing to their reliance on tender points and lack of consideration of other symptoms (3). Revised ACR diagnostic criteria, published in 2010 (9), were not meant to replace the 1990 criteria, rather they were an alternative for clinical diagnosis. As the revised criteria do not require a tender point examination (Table 1) (9) and are simple to administer, they might prove to be more practical and user-friendly for primary care physicians.

A further modification of the ACR criteria, in 2011, was intended to simplify them for practical use in epidemiological and clinical studies (10). The 2011 criteria include a 1-page patient self-report symptom survey to determine the locations of pain and the presence/severity of fatigue, sleep disturbances, memory difficulties, headaches, irritable bowel symptoms and mood problems (for further information, Clauw (6) and Wolfe et al. (10)).

**Diagnosis of FM in clinical practice**
In clinical practice, FM should be considered in any patient reporting chronic multifocal or diffuse pain (6). FM is also commonly comorbid in patients with rheumatic diseases, including osteoarthritis, rheumatoid arthritis, systemic lupus erythematosus and ankylosing spondylitis (40); in patients with other pain conditions (41); and in those with thyroid dysfunction (42). A suspicion of FM might develop during symptom progression, especially if the patient visits the clinic on multiple occasions reporting chronic pain in various body areas, tiredness and problems with sleeping (5). The presence of some comorbid disorders might also be a key factor in helping to diagnose FM, especially mood disorders, IBS, migraine, pelvic or genitourinary pain and temporomandibular disorder (5). However, the presence of comorbidities increases the complexity of the patient, and is likely to impact on the rapidity of diagnosis. These patients are likely to take more time at the physician’s office and may require collaboration with specialists and other HCPs to produce an accurate diagnosis and optimal management plan (41,43).

Importantly, FM is not a diagnosis of exclusion (5), to be brought out as a last resort after testing for other conditions. The physician can assess the patient’s medical history to determine whether they meet the criteria for FM, and perform a physical examination (evaluation of joints for the presence of inflammation, a neurological examination and an assessment of tenderness or pain threshold by digital palpation) to assess for other potential contributing causes of the symptoms (5). Laboratory tests are usually not necessary to confirm a diagnosis of FM. Basic tests such as blood count and serum chemistries might be of use in guiding the assessment, and a thyroid function test can be used to assess hypothyroidism, which is common and treatable, but detailed serologic studies are not necessary unless an autoimmune or other condition is suspected based on the patient’s history and examination (5,6). If FM is suspected, patient screening can begin by asking the patient to complete self-report measures such as a body pain diagram and assessment of symptoms (5). Once diagnosed, treatment for FM can be initiated immediately, even if a patient requires further tests to clarify some unusual signs or symptoms, or requires referral to a specialist for evaluation of comorbidities (5).

**Treatment of FM**
As the pathogenesis of FM has not been entirely elucidated, this has limited the development of disease-modifying treatments (44). As such, current treatment options focus on symptom-based management to improve function and quality of life. However, it is generally accepted that integration of pharmacological and non-pharmacological treatments will give the best outcome for the patient (6).

**Pharmacological treatments**
Studies have shown that the majority of patients attempt to manage their symptoms themselves before presenting to a physician (8). This might account for the fact that the medications most commonly used by patients with FM include basic analgesics, such as acetaminophen and non-steroidal anti-inflammatory drugs (45), although there is limited evidence that they are effective in FM (46). More concerning, given the potential for misuse and addiction, a commonly prescribed treatment for FM (both before and after diagnosis) is short-acting strong opioids (45,47), despite clinical trial reports indicating that opioids do not reduce pain in FM (44,46,48).

In the USA, three drugs are currently approved for the treatment of FM (32): pregabalin (Pfizer Inc., New York, NY; approved 2007) (49), duloxetine (Eli Lilly and Company, Indianapolis, IN; 2008) (50) and milnacipran (Forest Pharmaceuticals, Inc., St. Louis, MO; 2009) (51). These medications work either to increase the activity of inhibitory neurotransmitters (to ‘turn down the pain volume’) or to reduce the
activity of facilitatory neurotransmitters (which ‘turn up the pain volume’) (6). In contrast, there are currently no medications approved for the treatment of FM in Europe, even though pregabalin, duloxetine and milnacipran have all been approved in Europe for other indications (32). Table 2 summarises the FDA-approved pharmacological treatment options for FM. Titration to the therapeutic dose is recommended to improve patient response. In some patients, starting at a lower dose and titrating more slowly may be necessary to lessen the risk of intolerance and discontinuation of treatment.

Other medications such as amitriptyline, cyclobenzaprine, gabapentin and fluoxetine have demonstrated efficacy in randomised, controlled trials of FM and are commonly used to treat FM, although they are not approved for this indication by the FDA (52–54). The selection of pharmacological agent(s) for the management of FM should be tailored according to a number of factors, including the presence of additional symptoms (e.g. fatigue, sleep disturbances) alongside pain, the presence of comorbidities such as anxiety or rheumatic disease, and the tolerability profile of the therapeutic options (6). Patients with FM often require multiple medications to treat their symptoms and comorbidities, and guidance on possible medication combinations has been previously published (54). It is important to select combination therapies that are not associated with adverse drug–drug interactions.

**Non-pharmacological treatments**
Non-pharmacological treatments should be an integral component of a prescribed treatment plan for patients with FM (31). Patient education, exercise, some forms of cognitive behavioural therapy (CBT), and sleep hygiene are the most-studied non-pharmacological treatments and have demonstrated efficacy in patients with FM (4,6).

Educational materials for patients are widely available on the Internet from many Web sites, including those run by the ACR (http://www.rheumatol-

### Table 2 A comparison of FDA-approved pharmacological medications for FM (pivotal studies) (32,49–51)

<table>
<thead>
<tr>
<th>Drug</th>
<th>FDA approval</th>
<th>Mechanism of action</th>
<th>Efficacy studies</th>
<th>Primary end-points</th>
<th>Dosing</th>
<th>Adverse events*</th>
</tr>
</thead>
</table>
| Pregabalin | 21 June 2007 | Non-selective \(\alpha_2\delta \) ligand | - 14 weeks, randomised, double-blind, placebo-controlled  
- 6 months, randomised, withdrawal | Pain reduction, improvements in PGIC and FIQ | 300–450 mg/day, start at 75 mg bid (might increase to 150 mg bid within 1 week); max dose 225 mg bid | Dizziness, somnolence, dry mouth, oedema, blurred vision, weight gain, abnormal thinking |
| Duloxetine | 16 June 2008 | SNRI                | - 3 months, randomised, double-blind, placebo-controlled  
- 6 months, randomised, double-blind, placebo-controlled | Pain reduction, improvements in PGIC and FIQ | 60 mg/day, start 30 mg/day for 1 week then increase to 60 mg/day | Nausea, dry mouth, somnolence, constipation, decreased appetite, hyperhidrosis |
| Milnacipran| 14 January 2009 | SNRI                | - 3 months, randomised, double-blind, placebo-controlled  
- 6 months, randomised, double-blind, placebo-controlled | Composite end-point that concurrently evaluated improvement in pain (VAS), physical function (SF-36 PCS) and patient global assessment (PGIC) | 100 mg/day, start 12.5 mg/day, increasing incrementally to 50 mg bid in 1 week; maximum dose 100 mg bid | Nausea, constipation, hot flush, hyperhidrosis, vomiting, palpitations, increased heart rate, dry mouth, hypertension |

bid, twice daily; FDA, US Food and Drug Administration; FIQ, Fibromyalgia Impact Questionnaire; FM, fibromyalgia; PGIC, patient global impression of change; SF-36 PCS, Short-Form 36 Physical Component Summary; SNRI, serotonin-norepinephrine re-uptake inhibitor; VAS, visual analogue scale.

*The most commonly reported adverse events are shown. For full details, please refer to the prescribing information for each drug.
FM (4,6,24,31). The non-pharmacological treatment therapy, relaxation training, forms of physical therapy, trigger-point injections, yoga, tai chi, acupuncture, chiropractic, massage therapy, and maintenance of improvements in FM. Referrals to CBT and sleep hygiene specialists should be made based on the facilities available in the local area and affordability for patients.

Complementary and alternative medicine might also be considered, but in general, there are few randomised, controlled trials of these treatments (e.g. yoga, tai chi, acupuncture, chiropractic, massage therapy, trigger-point injections, forms of physical therapy, relaxation training, diet) in patients with FM (4,6,24,31). The non-pharmacological treatment options for FM are summarised in Table 3.

### Strategies to manage FM in primary care
The key to effective management of patients with FM in primary care is an integrated approach to treatment, a coordinated framework of clinical and non-clinical support, multifaceted education and clarity of goals and expectations.

### Physician education
In order for the majority of FM diagnosis and treatment to take place in primary care, non-specialist physicians must have the necessary tools and training to recognise symptoms and feel confident in prescribing treatments. Unfortunately, although most primary care physicians receive some training in basic pain assessment and management, in many cases, it is too brief to be meaningful (11,34). Additional training might be required, either via some form of e-learning, or led by specialists or colleagues with experience in chronic pain, to disseminate information and translate knowledge into skills and actions (11,34).

A lack of knowledge of current diagnostic criteria might be one reason leading to delays in diagnosing FM, but primary care physicians might also be limited by the consultation time available to make a diagnosis, particularly when patients have multiple symptoms that must be evaluated and discussed (8). As patients might initially present with one of the symptoms commonly associated with FM, such as mood symptoms or fatigue, the physician might need to be proactive in enquiring about pain symptoms (48). The development, validation and widespread implementation of tools to simplify symptom assessment could be one way to improve diagnostic accuracy and reduce delays in initiating treatment (11,55).

### Patient education
As with any chronic condition that requires ongoing management, patient education is critical in aiding patient understanding, acceptance and self-management of their condition (4). The primary care physician is uniquely placed to form a strong therapeutic relationship with patients and provide critical ongoing support (48). The use of familiar terminology might help the patient better understand the clinical picture and provide reassurance (4). However, because time for patient education is likely to be limited during a consultation, the use of clinical support staff to provide supplementary information is key, along with details of useful educational sources (books, Web sites, advocacy groups, etc.) (4,48).

In addition to educating patients about FM, it is also recommended that physicians partner with patients to decide on treatments, set goals and manage their expectations of symptom improvement and impact on daily life (4,13,34). Poor communication between patient and physician is likely to lead to frustration and over-reliance on pharmacological interventions with limited benefit; whereas shared decision making and positive interactions might help patients engage with their treatment and actively manage their pain (48). Education around adherence might also be necessary, to encourage the continuation of treatment to allow time for symptomatic improvement (4).

### Setting treatment goals
It is important for patients with FM to understand the limitations of current treatments for their condition, and to acknowledge that therapy might restore and maintain quality of life and considerably reduce pain, but will seldom remove pain completely (17,48). As many aspects of daily life might be affected by FM, a key step is to identify which are most important to the patient and develop a treat-
ment plan based on prioritising the areas that affect them most (4). While some patients might simply want a reduction in pain, others might prefer to focus on obtaining restorative sleep, or reducing fatigue levels to improve work or family relationships (17). These goals should be established early after diagnosis, to provide structure and guidance for future consultations and treatment decisions, but it is important that they be realistic, specific and easily tracked to provide a measure of treatment benefit (4).

### Integrated multimodal treatment

A comprehensive treatment plan should include non-pharmacological treatments, pharmacological therapies and active patient coping strategies. As FM is associated with a constellation of symptoms, no single treatment can be expected to target every one of them. The treatment approach must be flexible to incorporate changes as the condition progresses, and it is likely to require the collaboration of a number of HCPs, particularly for the treatment of some

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Regimen</th>
<th>Reported outcomes</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patient education</td>
<td>Provide core information about diagnosis, treatment and prognosis; manage expectations</td>
<td>Can improve symptoms and functionality; might reduce disability levels</td>
<td>• Can be carried out as part of normal consultations</td>
<td>• Might need to be repeated during each consultation or require separate educational sessions • Might be time-consuming • Might require additional support staff to help provide education</td>
</tr>
<tr>
<td>Exercise</td>
<td>Start low, go slow: build up to moderate activity over time</td>
<td>Can improve physical function, quality of life and reduce symptoms of pain and depression</td>
<td>• Easily incorporated into daily routine • Even small increases in activity have been shown to be of value</td>
<td>• Might cause worsening of symptoms if exercise programme is begun too rapidly • Access to exercise facilities might be limited • Might require consultation with other HCPs (e.g. physical therapists)</td>
</tr>
<tr>
<td>CBT</td>
<td>Face-to-face counselling, online self-help courses, books, CDs, FM Web sites</td>
<td>Provides knowledge about FM and coping strategies. Can provide sustained improvements in FM symptoms, and reduce impact on daily life</td>
<td>• Effective in one-on-one settings, small groups and via the Internet • Internet-based programmes provide convenience for patients</td>
<td>• Most effective when combined with other treatments • Access to mental health providers might be limited and might be costly</td>
</tr>
<tr>
<td>Sleep hygiene</td>
<td>Optimise sleep environment and prioritise relaxing sleep routine</td>
<td>Can improve pain scores and mental well-being</td>
<td>• Easily incorporated into daily routine</td>
<td>• Patient might be resistant to changes in routine (e.g. avoiding coffee at night, not watching television in bed)</td>
</tr>
<tr>
<td>CAM therapies</td>
<td>Various: examples include tai chi, yoga, massage, diet, balneotherapy and acupuncture</td>
<td>Can increase patient self-sufficiency and improve pain/functioning</td>
<td>• Limited evidence for efficacy</td>
<td>• Most CAM therapies have not been rigorously studied • Limited access to some of these treatments in some communities • Might be costly</td>
</tr>
</tbody>
</table>

CAM, complementary and alternative medicine; CBT, cognitive behavioural therapy; FM, fibromyalgia; HCP, healthcare professional.

Table 3. A comparison of non-pharmacological therapies for FM (4,6)

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comorbidities (4). Patients can be encouraged to identify and maintain active coping strategies, in an attempt to reduce disability (34). Comorbidities, such as severe depression or marked psychosocial stressors, might necessitate referral to a mental health specialist, while medical comorbidities might require additional treatment from a range of specialists such as rheumatologists, gastroenterologists and sleep specialists. The primary care physician plays an important role in coordinating specialists and ancillary HCPs to provide continuity of care for the patient.

**Tracking progress**

Surveys of HCPs have reported that many primary care physicians report a lack of knowledge of treatment options and monitoring tools (11). This is a key limitation, because it is only by tracking symptom presence and severity that the impact of treatment can be evaluated. There are several scales and questionnaires available that have been developed to evaluate the different symptoms of FM, and these might be useful to provide an initial health status, and a marker from which progress can be tracked (4,35). However, such tools need to be reliable, validated in patients with FM, rapid to administer and easy to interpret, to be globally accepted and used routinely in the clinic.

**Using electronic records**

The use of computers and technology is now ubiquitous throughout society, and health care is no exception. In recent years, HCPs have moved towards keeping electronic records, providing an opportunity to integrate FM management, improve outcomes and reduce costs and unnecessary testing (35). Electronic records can improve access to patient information across multiple specialties that might be involved in care decisions, provide information to guide prescribing decisions according to current recommendations, reduce medication errors and possibly aid in identifying undiagnosed patients (35). In a recent retrospective analysis, it was shown that a potential diagnosis of FM was associated with more frequent emergency room visits, outpatient visits, and hospitalisations and higher medication use. The authors concluded that all of these variables could be identified from electronic medical records, suggesting that routine data collection and input could have a direct application to FM diagnosis and care management (56).

For HCPs, identification of patients undergoing multiple exploratory tests might aid in focusing resources, to break the cycle of long-term medical spending. Online or application-based tools could also expedite administration and interpretation of monitoring scales, to rapidly gain a clear picture of symptom control and therapeutic outcome (35).

**The medical home for management of FM**

It is possible to transform primary care into a system in which medical practices can be improved to provide team-based care and data-driven integrated delivery, using the concept of the patient-centred medical home (PCMH). In the PCMH, decision making is guided by evidence-based medicine and decision-support tools. Patients are active partners in their treatment and information technology is utilised to support education, communication, data collection and performance measurement (57,58). The principles of the medical home were developed by key organisations, including the American College of Physicians and the American Academy of Family Physicians. The aim of a PCMH was to provide comprehensive primary care for all ages and throughout all stages of life, by coordinating and integrating care (chronic, acute, preventative and end-of-life) across all elements of the healthcare system, to improve efficiency and effectiveness (Figure 1) (57,58).

While the PCMH may not be feasible in all practices (owing to an absence or scarcity of resources) or in all countries (due to the widely varying healthcare systems between nations), it can provide a vision for the future management of FM and other chronic conditions by demonstrating how integration and coordination of doctors, hospitals, pharmacies and community resources can improve patient experience and outcomes while potentially reducing waste and inefficiency (59,60). The changing landscape of health management across the US and elsewhere (60–64) provides an opportunity for many HCPs and practices to implement a chronic care framework for FM management, similar to that already in use for diabetes (4). Results to date indicate that the PCMH is a viable mechanism to qualitatively improve diabetes management, while potentially reducing the costs of long-term care (65–68). The PCMH concept has also been successfully implemented in the field of mental health, resulting in reduced rates of hospitalisations, fewer specialty care visits and increased primary care consultations for patients with conditions such as post-traumatic stress disorder (69,70). However, of all the patients treated in primary care, those with chronic pain are most in need of practice reform (71). The first steps towards improving FM care have already been taken, with recent publications from the USA and the UK laying the groundwork for a focused and supported management pathway for patients with FM and chronic pain (4,34,48). It is hoped that by addressing the
current challenges and suggesting potential areas for restructuring, proposals for PCMH implementation and FM management in primary care can be implemented rapidly and smoothly into current practices.

Implementing a medical home for FM

Personnel
One obvious factor affecting the adoption of any new primary care framework is practice size. Small practices, with just one or two physicians, are unlikely to have either the personnel or the systems to be able to fully implement the PCMH concept (72). However, the current trend in the USA is towards larger practice sizes, since they might enjoy economies of scale, whereby several physicians can share support staff (72,73).

In the PCMH model, a typical primary care office is likely to require two to four support staff for each physician (74,75). In an office with four to six full-time employees, this is likely to mean two full-time physicians and several part-time support staff in various ratios. Support staff commonly includes registered nurses, physician assistants, nurse practitioners and medical assistants (see Appendix 1), as well as a pharmacist, who might be shared between several practices (74,75). For FM, and other chronic pain conditions, registered nurses or health coaches are likely to be a key among these team members, enabling patients to understand their condition, and instructing them in the mechanisms and benefits of self-management (76–78). Since patients with FM commonly have psychiatric comorbidities, behavioural health workers might also be a necessary adjunct to the team, alongside care coordinators, a largely clerical role, but pivotal to ensuring referrals are made and followed up (76–78).

The aim of the PCMH is to engage multiple HCPs in providing hands-on management to assist patients in navigating the care system. This requires a team-based approach, to spread the load, maximise efficiency and make the best use of each team member’s professional skills (79). One of the key ingredients of a successful PCMH is effective leadership within the practice, both to facilitate the transition and to serve as the patient’s primary care provider (62). Depending on state law, this leadership might come from a physician or from a nurse practitioner (78–80). In either case, the individual must be able to meld diverse personalities with widely differing levels of training into a cohesive team, all members of which are functioning at the highest level and contributing to the health of their patients (81,82). Conversely, one potential obstacle to overcome might be a reluc-
Fibromyalgia management for primary care providers

Improved HCP and increased job satisfaction, because they have an improved HCP–patient relationship, and are better able to focus on the more complex aspects of care (76). Medical assistants and nursing staff report improved job satisfaction from the increased responsibility and feeling more involved in patient care (76). Furthermore, PCMH reform can help to improve primary care attitudes towards patients with chronic pain, by providing incentives and increasing opportunities for specialised education and training (71).

Challenges

While the PCMH is an appealing proposition in terms of benefit to patients and HCPs, there are also several challenges associated with the concept, which need to be carefully considered prior to initiating practice reform. Significant time and expense may be needed to meet the required criteria and benchmarks (58–60,83), which may tax the resources of small practices and solo practitioners. It may be necessary to hire additional staff to meet the management and administrative demands of PCMH operations, upgrade and maintain IT infrastructure, and establish the type of electronic records network necessary to fulfil PCMH technology and access requirements (58,62). Geographical location may also be an issue, because a small rural practice without adequate local specialists, non-physician HCPs or supportive community resources may be limited in its ability to meet collaborative care standards (83).

However, physicians should not be discouraged from implementing at least some aspects of the PCMH, and should seek advice from experienced healthcare advisors who will be able to assess the ability of each practice to meet the PCMH requirements or develop other viable options that may be better suited to the needs and capabilities of any given practice. Furthermore, financial support, training and technical aid may be available to assist in the transition process towards PCMH recognition (59,64,83).

Best practice

For a patient such as Susan, getting a diagnosis of FM often takes several years, many examinations and procedures, and multiple visits to various doctors. However, implementation of a medical home is an opportunity to reduce the timescale between presentation and diagnosis, and revise the scenario to limit unnecessary tests and referrals. FM is a clinical diagnosis that can be appropriately made by primary care physicians based on the clinical characteristics of the disorder. Faster symptom recognition and diagnosis might be possible, to enable earlier treatment initiation. The PCMH has been shown to improve outcomes in diabetes and mental health; thus, it should be viable to adapt the model for FM and chronic pain.

Case study: Susan King

| Current symptoms: In addition to widespread pain, Susan reports regular sleepless nights, resulting in feeling unrefreshed and tired for most of the day. She feels that she is not ‘clear-headed’ and is unable to concentrate on regular tasks at times. Her fatigue means that she is so exhausted after work that she is unable to interact with her husband and daughter, or take part in normal social activities. Susan is also conscious that since she is sedentary at her job, she should make time for physical exercise. However, although she previously participated in regular aerobic exercise, she has not exercised in the past 9 months due to always feeling tired. With further enquiry, Susan remembers that during her childhood, her mother also had similar complaints |
| Case study: Susan King |

Patient: Susan King is a white female aged 45 years, married, with one child (a girl, currently 15 years of age)

Medical history: Susan has a history of migraines that started in adolescence. Susan also had some depressive episodes while in college but did not seek treatment and was never formally diagnosed. Just over 3 years ago, she was promoted to a more stressful position at work. Around the same time she began to suffer from widespread pain and symptoms of irritable bowel syndrome. These symptoms resulted in Susan having to take time off from work because of pain and fatigue. Depressive symptoms also recurred a couple of years ago, subsequent to the promotion and following several months of unexplained pain

Given her symptoms, Susan is most likely to present to her primary care doctor several times over a few weeks or months. The primary care physician is therefore ideally placed to observe and record these seemingly disparate and generalised symptoms (pain, depression, fatigue, IBS), and to suspect that FM could be the underlying cause that links them together. In addition to more education in chronic pain, the development of FM- or pain-specific tools that could be easily used during an office consultation would further assist the primary care physician in making the diagnosis of FM. Several such screening/diagnostic tools...
are currently under evaluation for use in primary care, including the Fibromyalgia Diagnostic Screen (55,84) and the FibroDetect® tool (85). Both appear to have good sensitivity and specificity, and may facilitate the identification of patients with FM in the primary care setting, although further validation in diverse settings is required.

**Case study: Susan King**

**Diagnosis**
What: medical history, physical examination, basic laboratory tests. Who: primary care physician, nurse practitioner or physician assistant. Results discussed with team, and diagnosis relayed to patient by primary care physician

With the primary care physician as PCMH ‘team captain’, he/she makes the diagnosis and manages effective treatment, and other members of the team act on their roles in ongoing care. Physician assistants and nurse practitioners might carry out tests to evaluate the patient’s symptoms and will liaise with the primary care physician to develop a management plan. A clinical pharmacist advises on treatment guidelines and local availability of medication, and allows for remote dispensing. Registered nurses and health coaches help patients to take control of their situation and coach them on self-management techniques. Care coordinators and medical assistants ensure that required tests are carried out, that results are entered into an electronic health record system that allows access by all stakeholders, and that any referrals deemed necessary are coordinated with the relevant hospital or specialist.

Although patients with FM can be very challenging to diagnose and treat, there is good evidence to suggest that interventions meeting PCMH criteria are associated with an overall improvement in patient satisfaction and perceptions of care (63). By putting the patient at the centre of care, the PCMH allows patients to manage their own lives (86), and gives them strategies to help themselves (87), rather than viewing themselves as invalids reliant upon HCPs to ‘cure’ them. Currently, patients with FM are inclined to try to use specialists as primary care providers, whereas the PCMH would reduce this problem, introducing specialist consultations only when needed. However, to achieve this, appropriate self-management tools are necessary, and the development of suitable Web sites and community resources will be a key element.

**Conclusions**

The management pathway for FM and chronic pain is currently often lengthy and complex, involving repeated clinic visits, unnecessary referrals and costly tests. The medical home, a patient-centred management framework which has been successfully implemented in other chronic diseases, might provide the key to reducing diagnosis time and improving patient outcomes. The PCMH sets up a health delivery model within the practice via the provision of a primary care team incorporating professionals with a range of skills and training, all functioning at the highest level for maximum efficiency and working together for the benefit of the patient. A multifaceted approach to treatment, including patient education and non-pharmacological and pharmacological therapies, is a key, but prioritising symptoms, tracking progress and managing patient expectations are equally important. Effective approaches to helping practices adopt the
medical home and tailor it to the needs of the patient with chronic pain will be important. Although there remain several barriers to overcome, implementation of a PCMH for chronic pain would allow FM to be successfully managed in the primary care setting.

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Author contributions

All authors contributed to the article conception, critical revision of each draft and approval of the final version.

References


50 Cymbalta (duloxetine delayed-release capsules) for oral use [package insert]. Indianapolis, IN: Lilly USA, LLC, 2012.


# Appendix 1: Healthcare provider definitions.

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<thead>
<tr>
<th>Job title</th>
<th>Responsibilities</th>
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<tbody>
<tr>
<td>Behavioural health worker</td>
<td>Support staff worker who provides psychological therapeutic support to patients with behavioural health issues and psychological disorders; generally requires a qualification in psychology, social work, counselling or nursing</td>
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<tr>
<td>Care coordinator</td>
<td>Liaises between patients and other healthcare professionals; ensures patients understand their medical condition and treatment, locates community resources and coordinates patient care services and referrals</td>
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<tr>
<td>Dietician</td>
<td>An expert in human nutrition and the regulation of diet; advises people on what to eat to achieve health-related goals</td>
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<tr>
<td>Health coach</td>
<td>An individual trained to assist patients by promoting coping behaviours, goal setting and overcoming negativity; generally requires a qualification in exercise science, nutrition, health care or wellness. Similar processes may also be performed by a psychotherapist</td>
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<tr>
<td>Healthcare professional (HCP)</td>
<td>Any individual trained to provide healthcare services; may include physicians, nurses, therapists and support workers</td>
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<tr>
<td>Medical assistant</td>
<td>A healthcare professional supporting physicians and other healthcare providers; they perform routine tasks and procedures such as measuring vital signs, collecting biological specimens, completing electronic medical records and scheduling appointments. Qualifications and requirements for certification vary between jurisdictions</td>
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<tr>
<td>Nurse practitioner</td>
<td>An advanced practice registered nurse who has been trained to diagnose and manage acute illness and chronic conditions. A nurse practitioner may serve as a primary care provider; in the USA, depending upon which state they work in, nurse practitioners may or may not be required to practice under the supervision of a physician</td>
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<td>Pharmacist</td>
<td>Healthcare professional who understands the mechanisms and actions of drugs, side effects, drug interactions and monitoring requirements; they provide pharmaceutical information and oversee the dispensation of prescription medication as well as non-prescription or over-the-counter drugs. A further education qualification is required</td>
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<tr>
<td>Physical therapist</td>
<td>Rehabilitation professional who manages patients with health conditions that limit their ability to move and perform functional activities</td>
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<td>Physician assistant</td>
<td>A healthcare professional who is licenced to practice medicine as part of a team with physicians and other providers; may be known as a physician associate in the UK. A physician assistant may conduct physical exams, order tests, diagnose and treat illnesses and perform medical procedures under the supervision of another physician</td>
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<tr>
<td>Primary care physician</td>
<td>A physician who provides the first point of contact for a patient and continuing care of medical conditions; may be known as a general practitioner in English-speaking countries outside of the USA</td>
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<tr>
<td>Primary care provider</td>
<td>A healthcare professional providing day-to-day health care in a primary care setting; may be a primary care physician, nurse practitioner or physician assistant</td>
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<tr>
<td>Psychiatrist</td>
<td>A physician specialising in the diagnosis and treatment of mental disorders</td>
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<td>Registered nurse</td>
<td>A nurse who has undergone training and met the requirements to obtain a nursing licence</td>
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<tr>
<td>Specialist</td>
<td>A physician or surgeon who has completed further medical education and training in a specific branch of medical practice</td>
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