Role and rationale for the use of milnacipran in the management of fibromyalgia

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Abstract: Fibromyalgia (FM) is a complex syndrome characterized by chronic widespread musculoskeletal pain which is often accompanied by multiple other symptoms, including fatigue, sleep disturbances, decreased physical functioning, and dyscognition. Due to these multiple symptoms, as well as high rates of comorbidity with other related disorders, patients with FM often report a reduced quality of life. Although the pathophysiology of FM is not completely understood, patients with FM experience pain differently from the general population, most likely due to dysfunctional pain processing in the central nervous system leading to both hyperalgesia and allodynia. In many patients with FM, this aberrant pain processing, or central sensitization, appears to involve decreased pain inhibition within the spinal tract, which is mediated by descending pathways that utilize serotonin, norepinephrine, and other neurotransmitters. The reduced serotonin and norepinephrine levels observed in patients with FM suggest that medications which increase the levels of these neurotransmitters, such as serotonin and norepinephrine reuptake inhibitors (SNRIs), may have clinically beneficial effects in FM and other chronic pain conditions. Milnacipran is an SNRI that has been approved for the management of FM. In clinical trials, treatment with milnacipran for up to 1 year has been found to improve the pain and other symptoms of FM. Because FM is characterized by multiple symptoms that all contribute to the decreased quality of life and ability to function, the milnacipran pivotal trials implemented responder analyses. These utilized a single composite endpoint to identify the proportion of patients who reported simultaneous and clinically significant improvements in pain, global disease status, and physical function. Other domains assessed during the milnacipran trials include fatigue, multidimensional functioning, mood, sleep quality, and patient-reported dyscognition. This review article provides information intended to help clinicians make informed decisions about the use of milnacipran in the clinical management of patients with FM. It draws primarily on results from 2 of the pivotal clinical trials that formed the basis of approval of milnacipran in the United States by the Food and Drug Administration.

Keywords: fibromyalgia, milnacipran, pain, serotonin and norepinephrine reuptake inhibitors, SNRI

Introduction

Fibromyalgia (FM) is a chronic pain disorder that affects 0.5% to 5% of the general population worldwide, more frequently in women than in men.1-3 The management of this disorder is complicated by the occurrence not only of pain, but of multiple other symptoms including fatigue, stiffness, sleep disturbances, physical dysfunction, and cognitive problems.4-6 It is also complicated by frequent comorbidity with conditions that share clinical and genetic characteristics and possibly a common pathophysiology. These comorbid conditions include low back pain, diabetic peripheral neuropathy, chronic...
fatigue syndrome, irritable bowel syndrome (IBS), migraine headache, interstitial cystitis, multiple chemical sensitivities, and temporomandibular disorder (TMD). The common theme to these conditions appears to be heightened sensitivity to discomfort/stimuli in various regions of the body.

Patients with FM often experience a significantly diminished quality of life, and frequently report an inability to work and feelings of isolation due to their withdrawal from social activities. FM can result in distressing physical, social and psychological consequences. Therefore, the approval of 3 agents over the past few years for the management of FM represents important progress in the ongoing development of evidence-based therapies for FM.

In the late 1990s, Cypress Bioscience, Inc. observed a large, unmet medical need revolving around the satisfactory treatment of chronic pain patients who were being diagnosed with FM. Although the FM diagnosis was somewhat controversial at the time, there was a large group of patients with numerous somatic complaints who were dissatisfied with their current health status. Moreover, physicians who managed FM patients were expressing frustration with the lack of effective treatment options. Various medications were being used off-label to treat these patients, including tricyclic antidepressants (TCAs), selective serotonin reuptake inhibitors (SSRIs), opioids, nonsteroidal anti-inflammatory drugs (NSAIDs), growth hormone, corticosteroids, and sedative hypnotics. While the TCAs have been found to provide some relief from FM symptoms (particularly sleep-related complaints), their use is often limited by safety and tolerability considerations. The SSRIs offer a better tolerability profile, but results from clinical trials in patients with FM have been disappointing. Evidence for the other medications such as opioids and NSAIDs is either lacking or weak.

In response to this unmet need in FM, Cypress began searching for a medication with a suitable neurotransmitter reuptake profile that could be developed as a potential first-line treatment option for patients with FM. One factor contributing to the choice of milnacipran for clinical development was the extensive safety data that already existed for this drug. Although milnacipran was not commercially available in the United States before its approval for the management of FM in 2009, it had been widely used in Europe (since 1997) and in Asia (since 2000) for the treatment of major depressive disorder. Its tolerability in depressed patients had been established in a number of clinical trials and was supported by several million patient-months of postmarketing safety data at the time Cypress licensed the rights for US development. Pierre Fabre Médicament, the original developer of milnacipran, was the licensor and still maintains the global milnacipran safety database.

The scientific rationale behind the development of milnacipran for FM was that, as a serotonin and norepinephrine reuptake inhibitor (SNRI), this drug should have clinically significant analgesic effects. In the central nervous system, both serotonin and norepinephrine have been found to play important roles in pain perception via their involvement in descending antinociceptive pathways. Dysfunction in these descending pathways is thought to result in the hyperalgesic (heightened sensitivity to pain) and allodynic (painful response to nonpainful stimuli) states experienced by patients with FM and other related central sensitization syndromes such as IBS and TMD. The potential benefit of milnacipran in the treatment of FM was further supported by analgesic effects of SNRIs in animal models of pain, as well as findings showing decreased cerebrospinal fluid levels of serotonin and norepinephrine metabolites in patients with FM. Moreover, therapeutic benefits in FM had already been observed with drugs that inhibit the reuptake of both serotonin and norepinephrine, such as the TCA, amitriptyline. However, it should be noted that amitriptyline has not been tested in recent clinical trials using the same rigorous standards currently required by the US Food and Drug Administration (FDA) for approval of drugs for the management of FM. For example, recent FM clinical trials with now-approved medications such as milnacipran had much larger sample sizes and longer treatment durations than the older studies involving amitriptyline. In addition, these more recent studies have consistently implemented multiple efficacy measures, including pain, global improvements, fatigue, mood, and multidimensional functioning in an effort to address the complex, multisymptomatic nature of FM.

Unlike the TCAs, milnacipran has no significant direct action on adrenergic, muscarinic, or histaminergic receptors—pharmacologic actions associated with many of the unpleasant side effects of TCAs. Based on this pharmacologic profile, it was postulated that milnacipran would be clinically beneficial to patients with FM, providing improvements in pain with fewer adverse effects. Moreover, it was thought that a dual reuptake inhibitor would have more potential as an analgesic than medications that selectively block the reuptake of serotonin, which have not consistently demonstrated effectiveness in treating FM symptoms. Interestingly, drugs that selectively target noradrenergic activity alone also appear to have limited efficacy in patients with FM. This suggests that both noradrenergic and serotonergic activity are required to produce clinically significant pain relief.
Milnacipran received its approval for the management of FM based on the safety and efficacy results of 2 pivotal trials conducted in the United States. In addition to reviewing the data from these trials, this article describes the various outcome measures that were used to establish the efficacy of milnacipran in patients with FM. The aim of this review article is to highlight the results of these clinical trials in order to provide clinicians with a better understanding of milnacipran as a treatment option in FM.

Pharmacology
Milnacipran is highly water soluble, leading to rapid and wide absorption, with maximum concentration observed within 2 to 4 hours after dosing. The bioavailability of milnacipran is high (approximately 85% to 90%), and absorption is not affected by food intake. Milnacipran undergoes minimal first-pass metabolism, with approximately 55% of the drug excreted unchanged in urine. Its relatively short half-life (approximately 6 to 8 hours) is compatible with the recommended twice-daily dosing. Pharmacokinetic studies indicate that dose adjustment is not necessary based on age, gender, mild-to-moderate renal impairment, or mild-to-moderate hepatic impairment. However, caution should be exercised in patients with moderate renal impairment or severe hepatic impairment. In patients with severe renal impairment, the maintenance dose of milnacipran should be reduced by approximately 50% to 50 mg/day (25 mg twice daily).

The low extent of hepatic metabolism, low protein binding (13%), and minimal effects on cytochrome P450 enzymes indicate a low potential for pharmacokinetic drug-drug interactions. In healthy volunteers, co-administration of carbamazepine, digoxin, and lorazepam did not have clinically meaningful effects on the pharmacokinetics of milnacipran. Switching from fluoxetine to milnacipran also did not have an effect on milnacipran pharmacokinetics. In vitro, milnacipran did not demonstrate any significant affinity for adrenergic, serotonergic, dopaminergic, opiate, histaminergic, muscarinic, benzodiazepine, or gamma-aminobutyric acid receptors. These pharmacologic characteristics of milnacipran may be advantageous for the treatment of patients with FM, many of whom routinely take multiple medications or experience chemical sensitivities.

Clinical efficacy of milnacipran in FM trials
After the completion of a phase 2 trial that demonstrated the potential therapeutic benefits of milnacipran for patients with FM, Cypress Bioscience partnered with Forest Laboratories, Inc. to continue the clinical development of this drug for the management of FM. To date, 3 randomized, double-blind, placebo-controlled, multicenter phase 3 trials have been completed in the United States, and a phase 3 trial sponsored by Pierre Fabre Médicament has been conducted in Europe. This review focuses on data from the first 2 pivotal US trials upon which the FDA approval was based. Most of these data have been previously published or presented at scientific meetings; some of the information included in this review, however, is from data on file (Forest Laboratories, Inc.).

The first US pivotal trial included 888 patients with FM (based on the 1990 American College of Rheumatology [ACR] criteria for FM) and lasted 6 months (Study 1). The second included 1196 patients and lasted 3 months (Study 2). In both trials, patients were randomized to placebo, milnacipran 100 mg/day, or milnacipran 200 mg/day. In Study 1, twice as many patients were randomized to the 200 mg/day group than were randomized to either the 100 mg/day or placebo groups in order to better assess the long-term effects of this higher dose. Patients enrolled in the milnacipran trials were required to discontinue centrally acting pharmacotherapies and nonpharmacologic therapies commonly used to treat FM symptoms. Provision for limited use of rescue medication with hydrocodone was allowed in these 2 trials (dosage: ≤60 mg/day for ≤5 days). Patients were instructed not to take hydrocodone during the 48-hour period prior to scheduled study visits and the 2-week period prior to endpoint data collection. For patients who used rescue medication during the critical time periods, a prespecified data handling provision caused these patients to be analyzed as non-responders to treatment, regardless of their actual pain scores. The percentage of patients taking hydrocodone was similar in all treatment arms.

Patient characteristics
Baseline patient characteristics were similar between the first 2 milnacipran FM pivotal trials. In both trials, the patients were mostly women (>$90%), mostly white (>90%), and had a mean age of approximately 50 years. Based on the body mass index (BMI) criteria issued by the World Health Organization, more than 75% of the patients were either overweight (BMI ≥25 to 30) or obese (BMI ≥30) at baseline. Before initiating treatment, the patients were similarly impaired in terms of baseline pain scores and overall disease activity, as measured by the Fibromyalgia Impact Questionnaire (FIQ). Mean baseline scores on the Short Form-36 Health Survey (SF-36) Physical Component Summary (PCS) indicated that enrolled FM patients had markedly decreased physical functioning compared to US norms.
Patients with current major depressive episodes were excluded from the trials, but approximately 35% of randomized patients in both studies reported a history of depression. Additionally, mean baseline Beck Depression Inventory (BDI) scores in both trials were 14, indicative of mild depressive symptoms.

**Composite responder analyses**

The milnacipran trials were designed to evaluate the effect of milnacipran on multiple symptom domains of FM, including pain, fatigue, global improvement, sleep, and physical and mental functioning. The outcomes evaluated in these studies are consistent with the key symptom domains proposed by the Outcome Measures in Clinical Rheumatology Trials (OMERACT) fibromyalgia working group. One type of efficacy endpoint recognized by OMERACT for its usefulness in FM clinical trials is the composite responder index. This identifies the proportion of individual patients who have simultaneous, clinically meaningful improvements in multiple symptom domains. Composite endpoints have been used in clinical trials for various rheumatologic conditions, including rheumatoid arthritis and osteoarthritis, and provide information about specific patient responses as opposed to group mean changes. Although composite endpoints have some limitations, one strength is that they are inherently conservative estimates of clinical efficacy. Patients who do not meet the criteria for each of the component domains cannot be counted as having responded to treatment, even if they show dramatic improvement for a single domain. In other words, composite endpoints can only be as robust as their weakest constituent component (i.e., the one most difficult to change) and require that the therapy produces clinically significant changes in multiple domains in the same patient. As such, they set a high hurdle for success.

At the time the milnacipran studies were designed, it was not clear whether there was an appropriate instrument for the assessment of physical function as part of a composite responder definition. Therefore, 2 definitions of response were evaluated during the pivotal registration trials: a 2-measure composite responder analysis that required simultaneous improvements in pain and global status; and a more stringent 3-measure composite that required simultaneous improvements in pain, global status, and physical function.

To be classified as 2-measure composite responders, individual patients were required to experience ≥30% improvement from baseline in pain, as measured by Visual Analog Scale (VAS) 24-hour recall scores (recorded on a patient experience diary [PED], as described below), and a rating of “Much Improved” or “Very Much Improved” on the Patient Global Impression of Change (PGIC). Patients who experienced a ≥6-point improvement from baseline in their SF-36 PCS score, in addition to meeting the pain and PGIC criteria, were classified as 3-measure composite responders. All of the definitions of improvement were based on changes that met or exceeded established minimal clinically important differences for those measures.

All statistical analyses were conducted on the intent-to-treat (ITT) population. Several imputation methods were used to handle missing data in these registration trials. Baseline observation carried forward (BOCF), a conservative approach requested by the FDA for the approval of milnacipran for the management of FM, was the primary imputation method for analyzing the composite responder endpoints. In BOCF, a patient who does not successfully complete the final trial visit—irrespective of the reason—is declared a nonresponder, regardless of the actual data recorded up to that point. Sensitivity analyses were also conducted using last observation carried forward (LOCF) and observed cases (OC). LOCF is a widely employed imputation method that uses the last recorded value for patients with missing data, while OC is a straightforward method that only analyzes recorded observations and does not impute missing data. As an estimate of outcomes among patients who comply with and tolerate treatment, OC analyses of the milnacipran data may provide practicing physicians with useful information. OC results are therefore highlighted in this summary.

Patients who were treated with milnacipran experienced multidimensional improvements in pain, global status, and physical function. After 3 months of treatment in each of the pivotal studies, a significantly higher proportion of patients in the milnacipran groups met the 2-measure and 3-measure composite responder criteria, as compared with the placebo group ($P < 0.01$, both doses vs placebo; OC) (Figure 1). For the more stringent 3-measure composite analysis, response rates among milnacipran-treated patients were approximately twice the rates found in placebo-treated patients. Results after 6 months of treatment were similar to those found at the 3-month endpoint. At 6 months, response rates for the 2-measure composite responder analysis were 43.8%, 45.2%, and 27.9% for milnacipran 100 mg/day, 200 mg/day, and placebo, respectively ($P < 0.05$, both doses vs placebo; OC).

**Pain**

Improvement in pain was included as part of the composite responder analyses because chronic widespread pain is
central to the definition of FM and is rated by both patients and physicians as the most important core domain to be assessed in FM clinical trials. In addition to being included as one component of the primary composite endpoints, pain was evaluated separately in the milnacipran trials using various secondary outcome measures, given the primacy of this symptom in the experience of patients with FM. Pain data was collected on electronic PEDs that prompted patients to record their 24-hour recall pain, weekly recall pain, and current level of pain (“real-time”) by marking VAS scales displayed on these hand-held electronic diaries. The PEDs, which were customized for use in the milnacipran trials, provided patients with a more accurate tool to report on their pain experiences. In post hoc analyses of the milnacipran pivotal trials, these electronic PEDs were found to be more discriminatory and sensitive than paper-based pain assessments. This was probably due to the minimization of recall bias and the ability to capture data in the patients’ home environment. Use of these electronic diaries also helped to satisfy the FDA’s recent rigorous approach to the use of patient-reported outcomes in registration trials. At the time of application for FDA approval, over 1 million pain data points had been collected from patients enrolled in the milnacipran FM trials. The PED pain data were supplemented by paper VAS pain assessments captured from patients at each study visit.

Milnacipran has proven to be effective in reducing FM pain. Compared with placebo, milnacipran was associated with significant improvements in PED and paper-based VAS pain measures. Significant sustained pain reductions were observed as early as 1 week after stable-dose treatment with milnacipran (P < 0.05 vs placebo), and maximal pain relief was reached by 9 weeks of treatment (Figure 2). The pain component of the composite responder analysis (ie, ≥30% improvement from baseline PED VAS 24-hour recall pain score) represents a clinically meaningful improvement in FM pain. A significantly higher proportion of patients experienced ≥30% improvements in pain with milnacipran than with placebo in Study 1 (52.8%, 100 mg/day; 56.2%, 200 mg/day; placebo, 40.2%; P < 0.05, both doses vs placebo; OC) and Study 2 (52.3%, 100 mg/day; 54.8%, 200 mg/day; 38.4%, placebo; P < 0.01, both doses vs placebo; OC).

A post hoc analysis of these 2 pivotal trials was conducted to determine the proportion of days in which patients experienced a ≥30% improvement in pain. Over a 3-month treatment period, patients treated with milnacipran vs placebo had a significantly higher percentage of days with clinically meaningful pain relief (45%, 100 mg/day; 46%, 200 mg/day; 33%, placebo; P < 0.0001, both doses vs placebo; OC). Significant differences between milnacipran and placebo in the proportion of pain relief days were also detected using a more stringent measure of ≥50% improvement from baseline in PED VAS 24-hour recall pain scores (27%, 100 mg/day; 29%, 200 mg/day; 18%, placebo; P < 0.0001, both doses vs placebo; OC).
“Patient global” is one of the core domains identified by OMERACT as being essential for inclusion in FM clinical trial designs. While most FM patients have symptoms involving multiple domains, not all patients rate all symptoms as being of similar importance. Therefore, a measure that implicitly evaluates those domains of most importance to an individual patient has practical utility and face validity. In the milnacipran studies, patient global was incorporated into the composite responder endpoints using the PGIC, a simple instrument that asks patients to rate their overall improvement by completing a single statement (“Since the start of the study, overall, my fibromyalgia is”) using a Likert scale ranging from 1 (“Very Much Improved”) to 7 (“Very Much Worse”). After 3 months of treatment, approximately one-half of the patients treated with milnacipran had marked global improvements (ie, PGIC score \( \leq 2 \)) compared with approximately one-third of the placebo-treated patients (49.4%, 100 mg/day; 52.8%, 200 mg/day; 33.8%, placebo; OC; pooled data on file).

In analyses conducted to evaluate which symptom domains in the milnacipran trials were associated with global well-being, pain was the strongest independent factor correlating with PGIC scores among patients who reported any global improvements (PGIC score \( \leq 3 \)). Vitality, sleep, dyscognition, and physical function were also significantly and independently associated with PGIC improvements. These findings are similar to those recently reported by OMERACT. Using the PGIC as a surrogate of overall improvement, OMERACT researchers examined data from 10 clinical trials of FM and found that measures of pain, fatigue, multidimensional function, physical function, and stiffness had the highest correlation with PGIC ratings. These types of correlations suggest that overall changes in patient global well-being reflect changes in multiple FM symptoms, underscoring the need for therapies that have multidimensional clinical benefits.

**Functioning**

Given the impact of pain on daily functioning, patient-reported quality of life measures have been increasingly utilized in pain clinical trials and are now considered to be an essential area of assessment for the approval of chronic pain medications. OMERACT has similarly recognized the importance of using quality of life measures to assess the efficacy of FM therapies. One such measure is the SF-36, a validated health status scale that has been used in over 70 studies involving FM clinical patients. The SF-36 includes 36 items assessing 8 domains (physical functioning, bodily pain, role limitations due to physical problems, role limitations due to emotional problems, general health, mental health, energy/vitality, and social functioning) from which the PCS and Mental Component Summary (MCS) scores are derived.

In a recent review comparing the health status profile of individuals with FM with that of the general population and patients with other health conditions, FM patients were found to have poorer scores on all 8 SF-36 domains than did patients with hypertension, recent acute myocardial infarction, type II...
Fatigue

Fatigue is a common complaint among patients with FM, usually described as being physically or emotionally draining.\(^5^,\)\(^6\) Like pain, fatigue is a constant presence that fluctuates in intensity throughout the day, affecting a patient’s ability to perform daily tasks, to function in the work force, and to enjoy social or recreational activities.\(^4,\)\(^6,\)\(^5\)\(^9\) Therefore, medications that reduce fatigue are important to both patients with FM and treating physicians.

To assess the efficacy of milnacipran on fatigue, the pivotal trials utilized the Multidimensional Fatigue Inventory (MFI), a 20-item self-report instrument that measures several dimensions of fatigue: general fatigue, physical fatigue, mental fatigue, reduced motivation, and reduced activity.\(^6\)\(^0\) The MFI has been used to assess fatigue levels in various chronic conditions such as rheumatoid arthritis,\(^6\)\(^1\) ankylosing spondylitis,\(^6\)\(^2\) and Sjögren’s syndrome.\(^6\)\(^3\) In the milnacipran pivotal trials, significant improvements in fatigue were observed with milnacipran 200 mg/day compared with placebo, as evidenced by MFI total and all subscale scores (Table 1).\(^6\)\(^4\) Patients treated with milnacipran 100 mg/day compared with placebo showed significant improvements in MFI total scores and subscale scores of general fatigue, physical fatigue, and reduced motivation.

### Cognitive impairment

The cognitive functions most affected in patients with FM are working memory, episodic memory, and semantic memory; other impairments include attentional control, susceptibility to distraction, and loss of vocabulary.\(^6\)\(^5\) Patients who have cognitive dysfunction often describe their experiences as feeling disorganized, having difficulty planning, and being unable to remember words.\(^5\)\(^9\) Like pain and fatigue, this “fibro-fog” affects daily functioning, particularly with regard to a patient’s ability to work and drive.\(^5\)\(^9\)

The milnacipran clinical trials assessed self-reported cognitive function in FM patients by using the Multiple Ability Self-Report Questionnaire (MASQ). The MASQ assesses 5 cognitive domains: language, verbal memory, visuo-perceptual ability, visual memory, and attention.\(^6\)\(^6\) Significant improvements in the MASQ total score were

### Table 1 Additional significant outcomes in fibromyalgia patients receiving 3 months of milnacipran treatment

| Symptom domain | Outcome measures | Treatment groups* |
|----------------|------------------|-------------------|
| Function       | SF-36 PCS        | 100, 200          |
|                | SF-36 MCS        | 200               |
|                | SF-36 bodily pain| 100, 200          |
|                | SF-36 physical functioning | 100, 200 |
|                | SF-36 energy/vitality | 100, 200 |
|                | SF-36 role limitations-physical | 200 |
|                | SF-36 role limitations-emotional | 200 |
|                | SF-36 general health | 200 |
|                | SF-36 mental health | 200 |
|                | SF-36 social functioning | 200 |
| Fatigue        | MFI total        | 100, 200          |
|                | MFI general fatigue | 100, 200 |
|                | MFI physical fatigue | 100, 200 |
|                | MFI mental fatigue | 200 |
|                | MFI reduced motivation | 100, 200 |
|                | MFI reduced activity | 200 |
| Cognition      | MASQ total       | 200               |
|                | MASQ attention   | 100, 200          |
|                | MASQ verbal memory | 100, 200 |

*Treatment groups in pooled analysis of 2 pivotal trials\(^25,\)\(^26\) demonstrating significant least square mean differences from placebo in change from baseline score (\(P < 0.05\), OC).

**Abbreviations:** 100, milnacipran 100 mg/day; 200, milnacipran 200 mg/day; MASQ, Multiple Ability Self-Report Questionnaire; MCS, Mental Component Summary; MFI, Multidimensional Fatigue Inventory; OC, observed cases; PCS, Physical Component Summary; SF-36, Short Form-36 Health Survey.
observed in patients treated with milnacipran 200 mg/day compared with placebo (Table 1, pooled data on file). Similarly, significant results with both doses of milnacipran were found for the MASQ verbal memory and attention domains, both of which are reflective of FM cognitive deficits. These results indicate that milnacipran improves cognitive function in patients with FM, particularly in the domains (eg, memory and attention) most affected by this disorder.

Long-term studies
Until recently, FM clinical trials have tended to be short in duration, generally lasting 3 months or less. Given the chronic nature of FM, experts in the field have recommended that longer clinical trials be conducted in order to evaluate whether the benefits of FM treatments persist over time.\(^3\) To this end, 2 randomized, double-blind extension studies of milnacipran have been conducted in the United States: a 28-week extension study to the 6-month pivotal trial (\(N = 449\))\(^5\) and an extension study adding up to 39 weeks to the 3-month pivotal trial (\(N = 384\)).\(^6\) Additionally, a 12-month extension of the 3-month European milnacipran trial has been completed.\(^7\) In the US studies, patients who received milnacipran 200 mg/day during the lead-in pivotal trials, were continued on the same dosage in a blinded fashion during the extension studies. Patients who received placebo or milnacipran 100 mg/day were re-randomized to milnacipran 100 mg/day or 200 mg/day.

Results from the extension studies indicate that milnacipran effectively improves the pain and other symptoms of FM for at least 1 year.\(^6,8\) The clinical benefits of milnacipran were maintained in patients who received milnacipran in both the lead-in and extension studies (ie, 12 months of continuous milnacipran treatment). In these patients, efficacy outcomes at the end of the extension study (eg, painVAS scores, FIQ total and physical function scores, PGIC scores) were similar to those observed at the end of the lead-in study. A subgroup of patients (\(n = 100\)) re-randomized from placebo to milnacipran 200 mg/day after 6 months of placebo in the double-blind lead-in trial experienced additional improvements in pain, FIQ, and PGIC scores during the extension study period.\(^6\) By the end of 12 months, this subgroup of patients had achieved the same relative degree of pain and global improvement reported by those treated continuously with milnacipran for the entire 12 months. Results from another subgroup of patients who were re-randomized from 100 mg/day to 200 mg/day (\(n = 92\)) indicate that some patients may benefit from increasing the milnacipran dosage.\(^6\) In these patients, additional improvements in pain and other outcomes were observed during the extension study period during which they received the higher dosage. Data from patients receiving milnacipran for up to 3 years in the US trials will be available after the conclusion of a currently ongoing study.

Safety and tolerability
The safety and tolerability information presented here is from patients with FM who were enrolled in the phase 2 clinical trial\(^3\) and the 2 pivotal registration trials.\(^25,26\) Together, these trials included 2209 patients, 1557 of whom were treated with milnacipran and 652 of whom received placebo.

Discontinuation due to adverse events (AEs) occurred in 23.0% and 26.0% of patients receiving milnacipran 100 mg/day and 200 mg/day, respectively, compared with 12.1% of placebo-treated patients.\(^7\) Nausea and palpitations were the only AEs resulting in discontinuation in ≥2% of milnacipran patients and at an incidence greater than placebo. Treatment-emergent AEs (TEAEs) in the milnacipran trials were generally mild to moderate in severity, with nausea being the most common TEAE in all treatment groups. The placebo-corrected rates of nausea for milnacipran 100 mg/day and 200 mg/day were 14.9% and 19.7%, respectively.\(^7\) Nausea typically occurred early in treatment and was generally manageable by recommending that medication be taken with food, incorporating gradual dose escalation, and providing patient counseling. While the clinical trials were designed to have relatively inflexible dose escalation phases lasting 2 weeks, recent anecdotal reports from commercial usage suggest that a slower, more flexible approach to dose increases may be beneficial.

Headache was the second most common TEAE, occurring in 18.6%, 17.2%, and 13.7% of all patients receiving milnacipran 100 mg/day, 200 mg/day, and placebo, respectively.\(^7\) TEAEs that occurred in ≥5% of all milnacipran-treated patients at an incidence at least twice that of placebo were constipation, hot flush, hyperhidrosis (sweating), palpitations, vomiting, increased heart rate, dry mouth, and hypertension. Most of the aforementioned TEAEs would appear to be directly related to the effect of increased norepinephrine levels in the periphery. No new safety concerns were observed with prolonged exposure to milnacipran in the extension studies.\(^5,8,9\)

As with other drugs in this class, cardiovascular effects have been reported with milnacipran. In the FM clinical trials, treatment with milnacipran 100 mg/day was associated with mean increases in blood pressure up to 3.1 mmHg, as well as mean increases in pulse rate of 7 to 8 bpm.\(^20\) Potentially clinically significant increases in supine systolic blood pressure (≥180 mmHg with an increase of ≥20 mmHg)
occurred rarely (≤0.2%) in patients from all treatment groups.71 Potentially clinically significant increases in supine diastolic blood pressure (≥110 mmHg with an increase of ≥15 mmHg) occurred in 0.9% of patients receiving milnacipran, compared with 0.3% of patients receiving placebo. Potentially clinically significant increases in supine pulse rate (≥120 bpm with an increase of ≥20 bpm) occurred in 0.7% and 0% of patients receiving milnacipran and placebo, respectively (pooled data on file). Results from a recently published study indicate that at suprathapeutic doses, milnacipran does not significantly affect cardiac repolarization or contribute to QTc prolongation.72

Milnacipran was not associated with weight gain, which can occur with other medications used to treat FM.73 During the course of therapy, patients who received milnacipran tended to lose weight. At 3 and 6 months, the proportion of patients with clinically significant weight loss (≥5% of baseline body weight) was significantly higher with milnacipran vs placebo (P < 0.01, both doses vs placebo; both endpoints).74 Similar nausea rates in milnacipran-treated patients who lost or gained weight indicate that this weight loss was unrelated to nausea. The lack of weight gain has been observed for up to 12 months of milnacipran treatment.68 In contrast to the use of SSRIs for FM,74 sexual side effects were reported in <1.0% of milnacipran-treated patients enrolled in the placebo-controlled FM trials (data on file). These findings of lack of sexual side effects were also supported by the lack of a significant difference between milnacipran and placebo groups on the Arizona Sexual Experience Scale.25 In the small population of males with FM (n = 87) included in the placebo-controlled trials, genitourinary AEs were reported in at least 2% of male patients treated with milnacipran and occurred at a rate greater than in placebo-treated male patients.28 This observation is consistent with the mechanism of milnacipran in which the increased peripheral norepinephrine level causes an increase in muscle tone, including in the urethra.

**Conclusion: place in therapy**

The complexity and heterogeneity of FM limits the utility of single-instrument outcomes in determining therapeutic efficacy. The multifaceted approach used to evaluate efficacy in the milnacipran clinical trials reflects ongoing discussions by groups such as OMERACT regarding the key domains and assessment tools needed to adequately evaluate FM outcomes.42–44 In the milnacipran clinical trials, the implementation of 2 composite responder indices allowed investigators to identify the proportion of individual patients who experienced simultaneous improvements in multiple symptom domains. As the development of pharmacotherapies for FM and other chronic pain syndromes progresses, it is expected that other clinical studies, such as the recently reported trials with sodium oxybate,75,76 will continue to use such outcome measures in order to better address the multisymptomatic nature of these disorders.

Results of the responder analyses in the milnacipran trials indicate that the therapeutic benefits of this medication extend beyond its analgesic effects.25,26,34,35 Based on the OC analyses of patients who were compliant with treatment in the clinical trial program, the rate of response with milnacipran treatment was approximately 50% on the 2-measure composite endpoint (clinically meaningful reduction in pain plus “much improved” or “very much improved” global status). These encouraging results suggest that if patients with FM are counseled on the importance of medication compliance and are adequately informed about the side effects of milnacipran, which are generally mild and transient, approximately half of these patients will achieve clinically significant and meaningful improvement in multiple domains.

There were no statistical differences in composite responder rates between milnacipran 100 mg/day and 200 mg/day, although these trials were not powered to detect such differences. Moreover, with the exception of nausea, the AE profile was similar between the dosages. In clinical settings, however, some patients may benefit from the higher dose, as evidenced by results from the extension studies in which patients who were re-randomized to the higher dose for an additional 6 to 9 months of treatment experienced additional improvements in pain and multidimensional function. Thus, although milnacipran 100 mg/day is the recommended starting dose, some FM patients can be escalated to the higher dose on an as-needed basis and if tolerability allows. Due to the potential for cardiovascular side effects, blood pressure and pulse rate should be measured before starting therapy with milnacipran and periodically monitored in all patients throughout the course of treatment. If baseline hypertension or tachycardia is detected, this should be treated and controlled prior to starting therapy with any drug that impacts noradrenergic activity.

One aim of the milnacipran clinical development program was to differentiate the analgesic effects of milnacipran from possible antidepressant effects. Although milnacipran was found to relieve pain in both depressed and nondepressed FM patients during the phase 2 trial,33 a change was made in the phase 3 trial design to exclude patients with current major depressive episodes, so that the analgesic and
other therapeutic effects could be more clearly assessed in the absence of a potentially confounding effect on depression. In the phase 3 trials, improvements in pain and other efficacy outcomes were robust in nondepressed FM populations,25,26,34,35 with an analysis of Study 1 showing that milnacipran resulted in global improvements regardless of baseline depressive symptom severity.77

Standardized guidelines for the management of FM have not been established, although it has been recommended that drugs with strong clinical evidence be used as first-line therapies in patients with moderate to severe pain.67 Strong evidence has been found for all of the FDA-approved medications for FM (milnacipran, duloxetine, pregabalin),12 although no direct comparisons between these medications can be made in the absence of head-to-head clinical trials. Making comparisons based on available clinical trial results is also not possible due to the use of different study designs and primary and secondary outcome measures in the trials.78 However, it has been suggested that SNRIs, such as milnacipran and duloxetine, may be tried in patients with comorbid mood disturbances and/or physical function deficits whereas pregabalin, an alpha-2-delta ligand, may be more appropriate for patients with prominent sleep disturbances or anxiety.26,67 For patients who have a partial response to monotherapies, combination treatment with medications having different mechanisms of action may be beneficial.67,79 The pending results from a multicenter, randomized, controlled pilot study investigating the addition of milnacipran to pregabalin treatment are expected to help clarify the potential benefits of combination therapy.80 Such research may soon be complemented by findings from genetic and pharmacogenetic studies,81 which are being conducted with the goal of allowing more personalized approaches to FM treatment in the future.

Although the approval of milnacipran and other medications represents progress in the treatment of FM, none of these treatments has proven to be completely effective in treating chronic pain conditions. Additional therapeutic approaches including combination and multimodal therapies will likely be needed. Development of personalized approaches, perhaps coupled with agents that modulate other aspects of the CNS pain processing system, will be needed as we strive to improve the treatment of chronic pain conditions.

Disclosures
Drs Kranzler and Gendreau are officers of and shareholders in Cypress Bioscience, Inc., one of the companies involved in the development of milnacipran for fibromyalgia.

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References
1. Wolfe F, Ross K, Anderson J, Russell IJ, Hebert L. The prevalence and characteristics of fibromyalgia in the general population. Arthritis Rheum. 1995;38(1):19–28.
2. White KP, Harth M. Classification, epidemiology, and natural history of fibromyalgia. Curr Pain Headache Rep. 2001;5(4):320–329.
3. Branco JC, Bannwarth B, Failde I, et al. Prevalence of fibromyalgia: a survey in five European countries. Semin Arthritis Rheum. 2009.
4. Mease P. Fibromyalgia syndrome: review of clinical presentation, pathogenesis, outcome measures, and treatment. J Rheumatol. 2005;32 Suppl 75:6–21.
5. Bennett RM, Jones J, Turk DC, Russell IJ, Matallana L. An internet survey of 2,596 people with fibromyalgia. BMC Musculoskeletal Disord. 2007;8:27.
6. Bennett RM. Clinical manifestations and diagnosis of fibromyalgia. Rheum Dis Clin North Am. 2009;35(2):215–232.
7. Aaron LA, Buchwald D. Chronic diffuse musculoskeletal pain, fibromyalgia and co-morbid unexplained clinical conditions. Best Pract Res Clin Rheumatol. 2003;17(4):563–574.
8. Crooks VA. Exploring the altered daily geographies and lifeworlds of women living with fibromyalgia syndrome: a mixed-method approach. Soc Sci Med. 2007;64(5):577–588.
9. Clauw DJ. Fibromyalgia: update on mechanisms and management. J Clin Rheumatol. 2007;13(2):102–109.
10. Nishishinya B, Urrutia G, Walitt B, et al. Amitriptyline in the treatment of fibromyalgia: a systematic review of its efficacy. Rheumatology (Oxford). 2008;47(12):1741–1746.
11. Clauw DJ. Pharmacotherapy for patients with fibromyalgia. J Clin Psychiatry. 2008;69 Suppl 2:25–29.
12. Mease P, Choy EH. Pharmacotherapy of fibromyalgia. Rheum Dis Clin North Am. 2009;35(2):359–372.
13. Goldenberg DL, Burckhardt C, Crofford L. Management of fibromyalgia syndrome. JAMA. 2004;292(19):2388–2395.
14. Nakagawa A, Watanabe N, Omori IM, et al. Milnacipran versus other antidepressive agents for depression. Cochrane Database Syst Rev. 2009(3):CD006529.
15. Millan MJ. Descending control of pain. Prog Neurobiol. 2002;66(6):355–474.
16. Stahl SM. Fibromyalgia – pathways and neurotransmitters. Hum Psychopharmacol. 2009;24 Suppl 1:S11–S17.
17. Ablin K, Clauw DJ. From fibrositis to functional somatic syndromes to a bell-shaped curve of pain and sensory sensitivity: evolution to a clinical construct. Rheum Dis Clin North Am. 2009;35(2):233–251.
18. Mochizuki D. Serotonin and noradrenaline reuptake inhibitors in animal models of pain. Hum Psychopharmacol. 2004;19 Suppl 1:S15–S19.
19. Russell IJ, Vaeroy H, Javors M, Nyberg F. Cerebrospinal fluid biogenic amine metabolites in fibromyalgia/fibrositis syndrome and rheumatoid arthritis. Arthritis Rheum. 1992;35(5):550–556.
20. Arnold LM, Keck PE Jr, Welge JA. Antidepressant treatment of fibromyalgia. A meta-analysis and review. Psychosomatics. 2000;41(2):104–113.
21. O’Malley PG, Belden E, Tomkins G, Santoro J, Kroenke K, Jackson JL. Treatment of fibromyalgia with antidepressants: a meta-analysis. J Gen Intern Med. 2000;15(9):659–666.
22. Rao SG. Current progress in the pharmacological therapy of fibromyalgia. Expert Opin Investig Drugs. 2009;18(10):1479–1493.
23. Cates M, Boggs AA, Feldman J. Major depressive disorder. In: Chisholm-Burns MA, Wells BG, Schwinghammer TL, Malone PM, Kolesar JM, eds. Pharmacotherapy Principles and Practice. New York: McGraw-Hill; 2008:569–584.

24. Abeles M, Soltar BM, Pillinger MH, Abeles AM. Update on fibromyalgia therapy. Am J Med. 2008;121(7):555–561.

25. Mease PJ, Clauw DJ, Gendreau RM, et al. The efficacy and safety of milnacipran for treatment of fibromyalgia. a randomized, double-blind, placebo-controlled trial [published correction appears in J Rheumatol 2009;36(3):661]. J Rheumatol. 2009;36(2):398–409.

26. Clauw DJ, Mease P, Palmer RH, Gendreau RM, Wang Y. Milnacipran for the treatment of fibromyalgia in adults: a 15-week, multicenter, randomized, double-blind, placebo-controlled, multiple-dose clinical trial [published correction appears in Clin Ther 2009;31(2):446]. Clin Ther. 2008;30(11):1988–2004.

27. Puozzo C, Panconi E, Deprez D. Pharmacology and pharmacokinetics of milnacipran. Int Clin Psychopharmacol. 2002;17 Suppl 1:S25–35.

28. Forest Laboratories, Inc. Savella package insert, July 2009. Available from www.savella.com. Accessed December 8, 2009.

29. Puozzo C, Pozet N, Deprez D, Baillie P, Ung HL, Zech P. Pharmacokinetics of milnacipran in renal impairment. Eur J Drug Metab Pharmacokinet. 1998;23(2):280–286.

30. Puozzo C, Albin H, Vincon G, Deprez D, Raymond JM, Amouretti M. Pharmacokinetics of milnacipran in liver impairment. Eur J Drug Metab Pharmacokinet. 1998;23(2):273–279.

31. Puozzo C, Lens S, Reb C, et al. Lack of interaction of milnacipran with the cytochrome p450 enzymes frequently involved in the metabolism of antidepressants. Clin Pharmacokinet. 2005;44(9):977–988.

32. Paris BL, Ogilvie BW, Scheinkoenig JA, Ndikum-Moffor F, Gibson R, Kolesar JM, et al. The Multicenter Coordinator Study Group. A European multicenter randomized double-blind placebo-controlled monotherapy clinical trial of milnacipran in the treatment of fibromyalgia. J Rheumatol. 2005;32(10):1975–1985.

33. Arnold LM, Gendreau RM, Spera A, Gendreau J, Wang Y. Milnacipran 100 mg/day in the treatment of fibromyalgia: a randomized, double-blind, placebo-controlled trial [abstract]. Arthritis Rheum. 2009;60 Suppl 10:S57.

34. Branco JC, Zachrisson O, Perrot S, Mainguy Y, on behalf of The IMMPACT recommendations. J Pain. 2008;9(2):105–121.

35. Bennett RM, Palmer RH, Wang Y, Hufford MR. Day-to-day pain relief in fibromyalgia patients: results from 2 clinical trials [abstract]. J Pain. 2008;9(4 Suppl 2):P16.

36. Mease P, Palmer RH, Wang Y, Hufford MR. Day-to-day pain relief in fibromyalgia patients: results from 2 clinical trials [abstract]. J Pain. 2008;9(4 Suppl 2):P16.

37. Bennett RM, Palmer RH, Wang Y, Hufford MR. Day-to-day pain relief in fibromyalgia patients: results from 2 clinical trials [abstract]. J Pain. 2008;9(4 Suppl 2):P16.

38. Werner P, Clauw DJ, Arnold LM, et al. Efficacy and safety of milnacipran with the cytochrome p450 isoenzymes frequently involved in the metabolism of antidepressants. Clin Pharmacokinet. 2005;44(9):977–988.

39. Bennett RM, Schein J, Kosinski MR, Hewitt DJ, Jordan DM, et al. Efficacy and safety of milnacipran with the cytochrome p450 isoenzymes frequently involved in the metabolism of antidepressants. Clin Pharmacokinet. 2005;44(9):977–988.

40. Bennett RM, Arnold LM, Bennett R, et al. Fibromyalgia syndrome. J Rheumatol. 2007;34(6):1415–1425.

41. Mease P, Arnold LM, Choy EH, et al. Fibromyalgia syndrome module at OMERACT 9: domain construct. J Rheumatol. 2009;36(10):2318–2329.

42. Bennett RM, Palmer RH, Wang Y, Hufford MR. Day-to-day pain relief in fibromyalgia patients: results from 2 clinical trials [abstract]. J Pain. 2008;9(4 Suppl 2):P16.

43. Bennett RM, Arnold LM, Bennett R, et al. Fibromyalgia syndrome. J Rheumatol. 2007;34(6):1415–1425.

44. Mease P, Arnold LM, Bennett R, et al. Fibromyalgia syndrome. J Rheumatol. 2007;34(6):1415–1425.

45. Schwieterman WD. Issues in the design of new clinical trials for rheumatoid arthritis therapeutics. Nat Clin Pract Rheumatol. 2008;4(12):641–648.

46. Bingham CO 3rd, Bird SR, Smugger SS, Xu X, Tershakovec AM. Respondor analysis and correlation of outcome measures: pooled results from two identical studies comparing etoricoxib, celecoxib, and placebo in osteoarthritis. Osteoarthritis Cartilage. 2008;16(11):1289–1293.

47. Ferrera-Gonzalez I, Permanyer-Miralda G, Busse JW, et al. Methodologic discussions for using and interpreting composite endpoints are limited, but still identify major concerns. J Clin Epidemiol. 2007;60(7):651–657.

48. 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(2):105–121.

49. Farrar JT, Young JP Jr, LaMoreaux L, Werth JL, Poole RM. Clinical importance of changes in chronic pain intensity measured on an 11-point numerical pain rating scale. Pain. 2001;94(2):149–158.

50. Strand V, Singh JA. Improved health-related quality of life with effective disease-modifying antirheumatic drugs: evidence from randomized controlled trials. Am J Manag Care. 2008;14(4):234–254.

51. US Food and Drug Administration. Drug Approval Package, Savella (milnacipran HCl tablets). Available from www.accessdata.fda.gov/ drugatfda_docs/nDA/2009/022256s000TOC.pdf. Accessed December 8, 2009.

52. Little RJA, Rubin DB. Statistical Analysis With Missing Data. 2nd ed. Hoboken, N.J.: Wiley; 2002.

53. Williams DA, Gendreau RM, Clauw DJ. Electronic diaries have superior discrimination compared to paper-based pain assessment in individuals with fibromyalgia [abstract]. Arthritis Rheum. 2007;56 Suppl 9:S607.

54. Williams DAG, R M, Clauw DJ. A comparison between electronic diaries and paper-based pain assessment in individuals with fibromyalgia [abstract]. J Pain. 2008;9(4 Suppl 2):P16.

55. Mease P, Palmer RH, Wang Y, Hufford MR. Day-to-day pain relief in fibromyalgia patients: results from 2 clinical trials [abstract]. J Pain. 2008;9(4 Suppl 2):P16.

56. Geisser ME, Clauw DJ, Strand V, Gendreau RM, Palmer R, Williams DA. Contributions of change in clinical status parameters to Patient Global Impression of Change (PGIC) scores among person with fibromyalgia treated with milnacipran. Pain. 2010 Mar 22. [Epub ahead of print].

57. Gatgel RJ, Theodore BR. Evidence-based outcomes in pain research and clinical practice. Pain Pract. 2008;8(6):452–460.

58. Hoffman DL, Dukes EM. The health status burden of people with fibromyalgia: a review of studies that assessed health status with the SF-36 or the SF-12. Int J Clin Pract. 2008;62(1):115–126.

59. Arnold LM, Crofford LJ, Mease PJ, et al. Patient perspectives on the impact of fibromyalgia. Patient Educ Couns. 2008;73(1):114–120.

60. Snets EM, Garssen B, Bonke B, De Haes JC. The Multidimensional Fatigue Inventory (MFI) psychometric qualities of an instrument to assess fatigue. J Psychosom Res. 1995;39(3):315–325.

61. Hitchet S, Heir M, Kirwan JR. Measuring fatigue in rheumatoid arthritis: a systematic review of scales in use. Arthritis Rheum. 2007;57(3):429–439.

62. van Tubergen A, Coenen J, Landewe R, et al. Assessment of fatigue in patients with ankylosing spondylitis: a psychometric analysis. Arthritis Rheum. 2002;47(1):8–16.

63. d’Elsa HF, Rehnhberg E, Kivist G, Ericsson A, Konttinen Y, Mannerkorpi K. Fatigue and blood pressure in primary Sjogren’s syndrome. Scand J Rheumatol. 2008;37(4):284–292.

64. Clauw DJ, Palmer RH, Hufford MR, Zablocki R, Wang Y. Milnacipran improves fatigue in patients with fibromyalgia: results from two randomized, double-blind, placebo-controlled trials [poster presenta
tion]. Presented at the 72nd Annual American College of Rheumatology and 43rd Annual Association of Rheumatology Health Professionals Meeting. San Francisco, CA; 2008.
65. Glass JM. Fibromyalgia and cognition. J Clin Psychiatry. 2008;69 Suppl 2:20–24.
66. Seidenberg M, Haltiner A, Taylor MA, Hermann BB, Wyler A. Development and validation of a Multiple Ability Self-Report Questionnaire. J Clin Exp Neuropsychol. 1994;16(1):93–104.
67. Arnold LM. Biology and therapy of fibromyalgia. New therapies in fibromyalgia. Arthritis Res Ther. 2006;8(4):212.
68. Goldenberg DL, Clauw DJ, Palmer RH, Mease P, Chen W, Gendreau RM. Durability of therapeutic response to milnacipran treatment for fibromyalgia. Results of a randomized, double-blind, monotherapy 6-month extension study. Pain Med. 2010;11(2):180–194.
69. Ferrera R, Palmer R, Chen W, Gendreau R. Improvements in fibromyalgia symptoms are sustained for 1 year with milnacipran treatment: results from 2 double-blind, dose-controlled extension studies [abstract]. J Pain. 2009;10(4 Suppl 1):S60.
70. Branco JC, Cherin P, Späth M, Mainguy Y. Long-term therapeutic response to milnacipran treatment for fibromyalgia. A European 1-year extension study following a 3-month study [abstract]. Arthritis Rheum. 2009;60 Suppl 10:S529.
71. Gendreau J, Palmer RH, Thacker K. Milnacipran is safe and well tolerated in the treatment of fibromyalgia [poster presentation]. Presented at the 27th Annual Scientific Meeting of the American Pain Society; May 8–10, 2008; Tampa, Florida.
72. Periclou A, Palmer RH, Zheng H, Lindamood C. Effects of milnacipran on cardiac repolarization in healthy participants. J Clin Pharmacol. 2010;50(4):422–433.
73. Pfizer Inc. Lyrica package insert, April 2009. Available from www.lyrica.com. Accessed December 8, 2009.
74. Ferguson JM. SSRI antidepressant medications: adverse effects and tolerability. Prim Care Companion J Clin Psychiatry. 2010;3(1):22–27.
75. Russell JJ, Perkins AT, Michalek JE. Sodium oxybate relieves pain and improves function in fibromyalgia syndrome: a randomized, double-blind, placebo-controlled, multicenter clinical trial. Arthritis Rheum. 2009;60(1):299–309.
76. Russell JJ, Alvarez-Horine S, Zheng Y, Guinta DR, Holman AJ, Swick T. Effect of sodium oxybate on pain, PGIc, and composite scores in fibromyalgia – results from a phase 3 controlled trial [abstract]. Arthritis Rheum. 2009;60 Suppl 10:S528.
77. Gendreau MR, Clauw DJ, Palmer RH, et al. Efficacy of milnacipran in the treatment of fibromyalgia among patients with varying degrees of depressed mood [poster presentation]. Presented at the 161st Annual Meeting of the American Psychiatric Association; May 3–8, 2008; Washington, DC, USA.
78. Boomershine CS. First pregabalin and now duloxetine for fibromyalgia syndrome: closer to a brave new world? Nat Clin Pract Rheumatol. 2008;4(12):636–637.
79. Mease PJ, Seymour K. Fibromyalgia: should the treatment paradigm be monotherapy or combination pharmacotherapy? Curr Pain Headache Rep. 2008;12(6):399–405.
80. US National Institutes of Health, Clinical Trials Registry. Available from www.clinicaltrials.gov. Accessed December 8, 2009.
81. Buskila D, Sarzi-Puttini P, Ablin JN. The genetics of fibromyalgia syndrome. Pharmacogenomics. 2007;8(1):67–74.