VA/DoD CLINICAL PRACTICE GUIDELINE
FOR
MANAGEMENT OF CHRONIC MULTISYMPOTOM ILLNESS

GUIDELINE SUMMARY

Prepared by:
The Management of Chronic Multisymptom Illness Group

With support from:
The Office of Quality, Safety and Value, VA, Washington, DC
and
Quality Performance Assurance Directorate, United States Army MEDCOM

Full guideline available at:

QUALIFYING STATEMENTS
The Department of Veterans Affairs (VA) and The Department of Defense (DoD) guidelines are based upon the best information available at the time of publication. They are designed to provide information and assist decision-making. They are not intended to define a standard of care and should not be construed as one. Neither should they be interpreted as prescribing an exclusive course of management.

Variations in practice will inevitably and appropriately occur when providers take into account the needs of individual patients, available resources, and limitations unique to an institution or type of practice. Every health care professional making use of these guidelines is responsible for evaluating the appropriateness of applying them in the setting of any particular clinical situation.

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INTRODUCTION

Chronic multisymptom illness (CMI) and medically unexplained symptoms are a critical health care issue for the Veterans Health Administration (VHA) and the Department of Defense (DoD). CMI imposes a significant burden of illness, disability, and decreased quality of life on a number of military Service Members, families, and Veterans. Therefore, diagnosis and effective therapy and related management of CMI have great importance for Veterans Affairs (VA) and DoD. After every modern military combat deployment, some Service Members have reported illnesses characterized by multiple chronic symptoms upon their return. Systematic studies have demonstrated that CMI is similar to many historical postwar illnesses. Among these, population-based studies have consistently demonstrated a higher prevalence and severity of symptom reporting in Gulf War Veterans than in non-deployed Veterans or other control groups. While these symptom-based illnesses have been described after military deployments, the experience of CMI is not unique to those who served in the military, to any specific combat era, or to those who were deployed to either combat or non-combat environments.

Although the character of medically unexplained symptoms appears similar after modern wars, at this time there is insufficient evidence to determine if the excess symptoms reported after these deployments share a common precipitating factor or pathophysiology. The authors of this CPG defined a working case definition of chronic multisymptom illness with the goal of enhancing the health care and ultimately improving the health status for all the populations cared for in VA and DoD.

In developing this VA/DoD clinical practice guideline (CPG), the Work Group reviewed randomized clinical trials (RCTs) and systematic reviews on treatments for the symptoms commonly associated with CMI, including studies on related conditions with overlapping symptoms such as fibromyalgia, CFS, and IBS. It is likely that treatments found to be effective for one of these related or comorbid conditions are beneficial for some patients experiencing CMI; however, the generalizability of the findings of the studies of these conditions to CMI has not been definitely established.

While other chronic conditions were not specifically included in the literature review during the development of this CPG, the CMI guideline may be relevant to chronic conditions that manifest with multiple chronic symptoms and functional limitations. Chronic overlapping physical and cognitive symptoms are sometimes attributed to specific events or conditions such as mild traumatic brain injury (mTBI) or post-traumatic stress disorder (PTSD), when instead they may reflect contributions from multiple factors, and thus may be amenable to the recommendations contained in this CPG. Though not specifically studied, this CPG is likely to be a helpful adjunct to the current VA/DoD guidelines for mTBI, PTSD, and major depressive disorder (MDD), especially when patients report multiple chronic symptoms not readily explained by these or other health conditions.

This CPG is intended to provide primary care clinicians with a framework by which to evaluate the individual needs and preferences of patients who may be experiencing chronic multisymptom illness or medically unexplained symptoms, leading to improved clinical outcomes. It is also likely to be used by other health care professionals, including specialty care providers.

The overall expected outcome of successful implementation of this guideline is to:

- Formulate an efficient and effective assessment of the patient’s condition
- Optimize the use of therapy to reduce symptoms and enhance functionality
- Minimize preventable complications and morbidity
- Emphasize the use of personalized, proactive, patient-driven care
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- Emphasize the use of personalized, proactive, patient-driven care
Working Definition of Chronic Multisymptom Illness
Chronic multisymptom illness (CMI) is a label given to a diverse set of disorders including, but not limited to, chronic fatigue syndrome (CFS), fibromyalgia syndrome (FMS), and irritable bowel syndrome (IBS). CMI encompasses military-specific medically unexplained illnesses, such as Gulf War Illness, Gulf War Syndrome, or post-deployment syndrome. The definition of CMI also includes patients without accepted labels, defined by generally accepted criteria, who exhibit persistent or frequently recurring symptoms negatively impacting daily function for a minimum of six months duration from two or more of the following six categories: fatigue, mood and cognition, musculoskeletal (including pain), respiratory, gastrointestinal and neurologic (including headache). Patients with symptoms lasting less than six months, or who experience only one of the listed symptoms, or with a clearly organic-based disease that explains all/most of their symptoms were not covered in this report. Further consideration for inclusion should be given to symptoms affecting the following systems: genitourinary, cardiopulmonary, and sleep.

Scope of this CPG
Individuals who meet the above descriptive criteria and also meet established criteria for specific symptom-based syndromes (e.g., fibromyalgia, IBS, CFS) may benefit from this CPG. The CPG provides information on potential risk factors for CMI, diagnostic technologies that may be used for screening and assessment of CMI, management of CMI, and pharmacologic and non-pharmacologic therapies for the treatment of CMI. Risk factors that may be associated with predisposing, precipitating, and perpetuating CMI include medical (e.g., obesity), psychological (e.g., abuse history), and occupational/environmental (e.g., chemical exposure). The categories of diagnostic technologies considered under this CPG include biomarkers (biological markers and neuroimaging studies), neuropsychological test batteries, and sleep studies.

Some of the management approaches considered include team-based approaches, core competencies of the treatment team, patient-provider communication styles, the role of occupational and other rehabilitative services, behavioral health services, and patient follow-up practices. Pharmacologic therapies include, among others, antibiotics, antidepressants, and pain medications, while non-pharmacologic therapies included psychological (i.e., hypnosis), physiological (i.e., exercise) and complementary and alternative treatments (i.e., acupuncture, biofeedback, and nutritional supplements).

Evidence Review
The recommendations presented in this CPG are based on a systematic appraisal of the published evidence on the management of Veterans and Service Members with Chronic Multisymptom Illness (CMI). In areas where the evidence is particularly lacking, expert opinion served as the basis for the recommendations. Published evidence was identified through extensive searches of several research databases. Searches were designed to identify unique reviews, trials, and technology assessments. Searches of the World Wide Web were also performed to capture relevant grey literature that has not been indexed to the databases listed previously. The searches covered an extended time period of January 2000 through October 2013.

Evidence Assessment
In order for the clinician to be aware of the evidence base behind the recommendations and the weight that should be given to each recommendation, the recommendations are graded according to the level of confidence with which each recommendation is made. This CPG uses the GRADE methodology to assess the quality of the evidence base and assign a grade for the strength for each recommendation. The GRADE system uses the following four domains to assess the strength of each recommendation:

- Balance of desirable and undesirable outcomes
- Confidence in the quality of the evidence
- Values and preferences
- Other implications, as appropriate, (e.g., Resource use, feasibility, subgroup considerations.)

The strength of a recommendation is defined as the extent to which one can be confident that the desirable effects of an intervention outweigh its undesirable effects and is based on the framework above, which combines the four domains.

The GRADE of a recommendation is based on the following elements:

- Four decision domains used to determine the strength and direction (described above)
- Relative strength (Strong or Weak)
- Direction (For or Against)

The relative strength of the recommendation is based on a binary scale, “Strong” or “Weak.” A strong recommendation indicates that the Work Group is highly confident that desirable outcomes outweigh undesirable outcomes. If the Work Group is less confident of the balance between desirable and undesirable outcomes, the recommendation is graded as weak.

Using these elements, the grade of each recommendation is presented as part of a continuum:

- Strong For (or “We recommend offering this option …”)  
- Weak For (or “We suggest offering this option …”)  
- Weak Against (or “We suggest not offering this option …”)  
- Strong Against (or “We recommend against offering this option …”)

The discussion regarding the quality of evidence and the final grade of recommendations is provided in the full guideline available at: http://www.healthquality.va.gov

Algorithm Format
This clinical practice guideline includes an algorithm, which is designed to maximally facilitate clinical decision-making for the management of CMI. The use of the algorithm format was chosen based on the understanding that such a format can inform diagnostic and therapeutic decision-making, and has the potential to change patterns of resource use. It allows the provider to follow a systematic approach to critical information needed at the major decision points in the clinical process, and includes:

- An ordered sequence of steps of care
- Decisions to be considered
- Actions to be taken

A clinical algorithm diagrams a guideline into a step-by-step decision tree. Standardized symbols are used to display each step in the algorithm, and arrows connect the numbered boxes indicating the order in which the steps should be followed.

This CPG is not intended to serve as a standard of care. Standards of care are determined on the basis of all clinical data available for an individual case and are subject to change as scientific knowledge and technology advances and patterns evolve. This CPG is based on information available at the date of publication, and is intended to provide a general guide to best practices. The guideline can assist care providers, but the use of a CPG must always be considered as a recommendation, within the context of a provider’s clinical judgment, in the care of an individual patient.
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A clinical algorithm diagrams a guideline into a step-by-step decision tree. Standardized symbols are used to display each step in the algorithm, and arrows connect the numbered boxes indicating the order in which the steps should be followed.

- Rounded rectangles represent a clinical state or condition.
- Hexagons represent a decision point in the guideline, formulated as a question that can be answered Yes or No.
- Rectangles represent an action in the process of care.
- Ovals represent a link to another section within the guideline.

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**Algorithm**

**Evaluation and Management**

1. Patient presents with a spectrum of chronic symptoms not explained by other disorders and meeting the criteria for Chronic Multisymptom Illness (CMI) (See Box A)

2. Are unstable or urgent condition(s) present?
   - Y: Refer or treat, as indicated, before continuing in this algorithm
   - N: Conduct a thorough evaluation of symptoms and assess for comorbid conditions (See Box B)

3. Does CMI co-exist with another medical or psychiatric condition that may explain the symptoms?
   - Y: Refer or treat as indicated using appropriate evidence-based clinical practice guidelines
   - N: Continue to next page: Treatment interventions

**Box A: Definition of CMI**

Patients without a formal diagnosis but who exhibit symptoms from two of the following six categories for a minimum of six months duration: fatigue, mood and cognition, musculoskeletal, respiratory, gastrointestinal and neurologic.

**Box B: Elements of Assessment**

- Obtain medical history, conduct physical examination and psychological assessment.
- Consider diagnostic studies, as indicated*
- Consider additional and/or longer duration encounters.
- For alternative diagnosis only. Avoid any test for which there may be limited additional benefit

**Treatment Interventions**

1. Continue from first page

2. Have symptoms improved to patient satisfaction?
   - Y: Patient presents with pain predominant symptom
     - Consider: trial of SNRIs or TCA
     - Do not use: corticosteroids, antivirals or antibiotics
   - N: Patient presents with gastrointestinal predominant symptom
     - Consider: minimal contact psychological therapies
     - TCA, SSRI or pregabalin may also be considered
     - Use caution when considering stimulants
     - Treat in accordance with recognized evidence-based care for IBS

3. Patient presents with fatigue predominant symptom
   - Consider: trial of SNRIs or TCA
   - Do not use: corticosteroids, antivirals or antibiotics
   - Use caution when considering stimulants
   - Consider acupuncture

4. Patient presents with gastrointestinal predominant symptom
   - Consider: minimal contact psychological therapies
   - TCA, SSRI or pregabalin may also be considered
   - Use caution when considering stimulants
   - Consider acupuncture

5. Patient presents with global CMI
   - Consider trial of SSRIs, SNRIs, or intranasal corticosteroids

6. Have symptoms improved to patient satisfaction?
   - Y: Return to Box 2: Re-evaluate symptoms and management
   - N: Follow-up and re-assess as needed

7. Continue to next page: Treatment interventions

8. Continue from first page
**Algorithm**

**Evaluation and Management**

1. Patient presents with a spectrum of chronic symptoms not explained by other disorders and meeting the criteria for Chronic Multisymptom Illness (CMI) (See Box A).

2. Are unstable or urgent condition(s) present?
   - Y: Refer or treat, as indicated, before continuing in this algorithm.
   - N: Conduct a thorough evaluation of symptoms and assess for comorbid conditions (See Box B) (A, B).

3. Does CMI co-exist with another medical or psychiatric condition that may explain the symptoms?
   - Y: Refer or treat as indicated using appropriate evidence-based clinical practice guidelines.
   - N: Initiate trial of non-pharmacologic interventions:
     - Offer cognitive behavioral therapy.
     - Consider complementary and integrative medicine interventions.
     - Consider: mindfulness based therapy, reattribution, behavioral medical intervention, and/or brief psychodynamic interpersonal psychotherapy. (B)

4. Are unstable or urgent condition(s) present?
   - Y: Continue to next page: Treatment interventions.
   - N: Conduct a thorough evaluation of symptoms and assess for comorbid conditions (See Box B) (A, B).

5. box A: Definition of CMI
   - Patients without a formal diagnosis but who exhibit symptoms from two of the following six categories for a minimum of six months duration: fatigue, mood and cognition, musculoskeletal, respiratory, gastrointestinal, and neurologic.

6. Box B: Elements of Assessment
   - Obtain medical history, conduct physical examination and psychological assessment.
   - Consider diagnostic studies, as indicated.*
   - Consider additional and/or longer duration encounters.

7. Do CMI co-exist with another medical or psychiatric condition that may explain the symptoms?
   - Y: Refer or treat as indicated using appropriate evidence-based clinical practice guidelines.
   - N: Initiate trial of non-pharmacologic interventions:
     - Offer cognitive behavioral therapy.
     - Consider complementary and integrative medicine interventions.
     - Consider: mindfulness based therapy, reattribution, behavioral medical intervention, and/or brief psychodynamic interpersonal psychotherapy.

8. Have symptoms improved to patient satisfaction?
   - Y: Continue from first page.
   - N: Return to Box 2.

9. Patient presents with pain predominant symptom (C)
   - Consider: trial of SNRIs, SNRIs or pregabalin.

10. Patient presents with fatigue predominant symptom (D)
    - Consider: trial of SSRIs or TCA.

11. Patient presents with gastrointestinal predominant symptom (E)
    - Do not use: corticosteroids, antivirals or antibiotics.

12. Patient presents with global CMI (F)
    - Consider a trial of SSRIs, SNRIs, or mintrazapine.

13. Consider non-opioid analgesic or acupuncture.
    - Consider: trial of SNRIs or TCA.
    - Do not use: corticosteroids, antivirals or antibiotics.

    - Use caution when considering stimulants.

15. Do not use acupuncture.
    - Treat in accordance with recognized evidence-based care for IBS.

16. Have symptoms improved to patient satisfaction?
    - Y: Continue from first page.
    - N: Return to Box 2.

17. Follow-up and reassess as needed.

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* For alternative diagnosis only. Avoid any test for which there may be limited additional benefit.
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**A. Conduct a Thorough Evaluation of Symptoms**

1. The guideline panel recommends that all patients receive a thorough evaluation of symptoms based on clinical judgment. [Strong For]

2. This guideline panel recommends against the use of any test for which there may be limited additional benefit to confirm the diagnosis of CMI. Testing for rare exposures or biologic effects should only be done in the presence of supportive history or physical findings. [Strong Against]

3. This guideline panel suggests discussing risk factors using principles of health risk communication within a therapeutic patient-provider alliance for those patients who wish to further understand factors that could contribute to their condition. [Weak For]

**B. Management Strategies for Care of Patients with CMI**

4. The guideline panel recommends using a collaborative, team-based approach, including a behavioral health specialist, for the primary care management of patients with CMI. [Strong For]

5. The guideline panel recommends that the healthcare team use shared decision-making principles to develop a comprehensive and personalized treatment plan in the care and management of patients with CMI. [Strong For]

6. The guideline panel suggests that all providers involved in the care of patients with CMI enhance their knowledge of the following critical domains:
   a. Communication skills (e.g., active listening, risk communication/perception)
   b. Empathy skills
   c. Working with interdisciplinary teams
   d. The biopsychosocial model
   e. Risk factors for CMI and analogous conditions
   f. Military cultural competency
   g. Deployment related exposures [Weak For]

**C. Therapeutic Interventions for Global CMI**

7. The guideline panel suggests incorporating appropriate elements of physical activity as part of a comprehensive and integrated treatment plan for patients with CMI. [Strong For]

8. The guideline panel recommends offering cognitive behavioral therapy, delivered by trained professionals, for patients with CMI. [Strong For]

9. The guideline panel recommends considering mindfulness-based therapy, reattribution, behavioral medical intervention, and/or brief psychodynamic interpersonal psychotherapy, delivered by trained professionals, for patients with CMI. [Weak For]

10. The guideline panel recommends considering complementary and integrated medicine interventions as a component of personalized, proactive patient-driven care in the management of patients with CMI. [Weak For]

11. The guideline panel suggests considering a trial of selective serotonin reuptake inhibitor (SSRI), serotonin-norepinephrine reuptake inhibitor (SNRI), or mirtazapine for the treatment of clinical symptoms of CMI. [Weak For]

12. The guideline panel suggests against the use of doxycycline for the treatment of patients with clinical symptoms of pain-predominant CMI. [Weak Against]

13. The guideline panel recommends against the long-term use of opioid medications for the management of patients with CMI. [Strong Against]

**D. Therapeutic Interventions for Pain-Predominant CMI**

14. The guideline panel recommends considering acupuncture as part of the management of patients with pain-predominant symptoms of CMI. [Weak For]

15. The guideline panel recommends considering non-steroidal anti-inflammatory drugs (NSAIDs) for treating certain peripheral pain symptoms associated with CMI, though they do not necessarily lead to global beneficial effect. [Weak For]

16. The guideline panel suggests considering tramadol for treating certain peripheral pain symptoms associated with CMI that fail to respond to other non-opioid analgesic medications or non-pharmacologic approaches. [Weak For]

17. The guideline panel suggests a trial of serotonin-norepinephrine reuptake inhibitor (SNRI) for the treatment of patients with clinical symptoms of pain-predominant CMI. [Weak For]

18. The guideline panel suggests considering a trial of tricyclic antidepressants (TCA), selective serotonin reuptake inhibitor (SSRI), or pregabalin (PGB) for the treatment of patients with clinical symptoms of pain-predominant CMI. [Weak For]

**E. Therapeutic Interventions for Fatigue-Predominant CMI**

19. The guideline panel recommends considering acupuncture as part of the management of patients with fatigue-predominant symptoms of CMI. [Weak For]

20. The guideline panel suggests considering a trial of serotonin-norepinephrine reuptake inhibitor (SNRI) or tricyclic antidepressants (TCA) for patients with clinical symptoms of fatigue-predominant CMI. [Weak For]

21. The guideline panel suggests against the use of pharmacologic agents for sleep disturbances in CMI. [Weak Against]

22. The guideline panel suggests against the use of stimulants for the treatment of fatigue predominant CMI. [Weak Against]

23. The guideline panel recommends against the use of immunotherapy for the treatment of the symptoms of fatigue predominant CMI. [Strong Against]

24. The guideline panel recommends against the empiric use of antivirals or antibiotics for fatigue predominant symptoms of CMI. [Strong Against]

**F. Therapeutic Interventions for Gastrointestinal-Predominant CMI**

25. The guideline panel suggests treating patients with CMI and predominantly gastrointestinal symptoms, in accordance with recognized evidence-based care for IBS. [Weak For]

26. The guideline panel recommends considering minimal contact psychological therapies for treatment of GI predominant CMI. [Weak For]

27. The guideline panel suggests against the use of acupuncture for treatment of patients with gastrointestinal-predominant symptoms of CMI. [Weak Against]

Strength of Recommendation is indicated in [ ]
A. Conduct a Thorough Evaluation of Symptoms

1. The guideline panel recommends that all patients receive a thorough evaluation of symptoms based on clinical judgment. [Strong For]
2. This guideline panel recommends against the use of any test for which there may be limited additional benefit to confirm the diagnosis of CMI. Testing for rare exposures or biologic effects should only be done in the presence of supportive history or physical findings. [Strong Against]
3. This guideline panel suggests discussing risk factors using principles of health risk communication within a therapeutic patient-provider alliance for those patients who wish to further understand factors that could contribute to their condition. [Weak For]

B. Management Strategies for Care of Patients with CMI

4. The guideline panel recommends using a collaborative, team-based approach, including a behavioral health specialist, for the primary care management of patients with CMI. [Strong For]
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   d. The biopsychosocial model
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   [Weak For]

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7. The guideline panel suggests incorporating appropriate elements of physical activity as part of a comprehensive and integrated treatment plan for patients with CMI. [Strong For]
8. The guideline panel recommends offering cognitive behavioral therapy, delivered by trained professionals, for patients with CMI. [Strong For]
9. The guideline panel recommends considering mindfulness-based therapy, reattribution, behavioral medical intervention, and/or brief psychodynamic interpersonal psychotherapy, delivered by trained professionals, for patients with CMI. [Weak For]
10. The guideline panel recommends considering complementary and integrated medicine interventions as a component of personalized, proactive patient-driven care in the management of patients with CMI. [Weak For]
11. The guideline panel suggests considering a trial of selective serotonin reuptake inhibitor (SSRI), serotonin–norepinephrine reuptake inhibitor (SNRI), or mirtazapine for the treatment of clinical symptoms of CMI. [Weak For]
12. The guideline panel suggests against the use of doxycycline for the treatment of patients with clinical symptoms of pain-predominant CMI. [Weak Against]
13. The guideline panel recommends against the long-term use of opioid medications for the management of patients with CMI. [Strong Against]

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14. The guideline panel recommends considering acupuncture as part of the management of patients with pain-predominant symptoms of CMI. [Weak For]
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21. The guideline panel suggests against the use of pharmacologic agents for sleep disturbances in CMI. [Weak Against]
22. The guideline panel suggests against the use of stimulants for the treatment of fatigue predominant CMI. [Weak Against]
23. The guideline panel recommends against the use of immunotherapy for the treatment of the symptoms of fatigue predominant CMI. [Strong Against]
24. The guideline panel recommends against the empiric use of antivirals or antibiotics for fatigue predominant symptoms of CMI. [Strong Against]

F. Therapeutic Interventions for Gastrointestinal-Predominant CMI

25. The guideline panel suggests treating patients with CMI and predominantly gastrointestinal symptoms, in accordance with recognized evidence-based care for IBS. [Weak For]
26. The guideline panel recommends considering minimal contact psychological therapies for treatment of GI predominant CMI. [Weak For]
27. The guideline panel suggests against the use of acupuncture for treatment of patients with gastrointestinal-predominant symptoms of CMI. [Weak Against]
In addition, the following other resources may be helpful:

1. War Related Illness and Injury Study Center – a National VA Post-Deployment Health Resource which provides post deployment health expertise to Veterans and their healthcare providers through clinical programs, research, education, and risk communication. Find out more here: [http://www.warrelatedillness.va.gov/](http://www.warrelatedillness.va.gov/)

2. Department of Veterans Affairs Office of Public Health – an office within the Veterans Health Administration which serves as the leader and authority in public health. Learn more about it here: [http://www.publichealth.va.gov/](http://www.publichealth.va.gov/)

3. Deployment Health Clinical Center – a site designed to provide a gateway to information on deployment health and healthcare for healthcare providers, service members, veterans, and their families. Check it out here: [http://wwwdeployment.mil/](http://wwwdeployment.mil/)

### ADDITIONAL RESOURCES

For more information, refer to the VA/DoD Evidence-based Clinical Practice Guideline for the Management of Chronic Multisymptom Illness, found at: [http://www.healthquality.va.gov/](http://www.healthquality.va.gov/)

### Table 1. Symptom Efficacy of Selected Pharmacotherapies

<table>
<thead>
<tr>
<th>Agent</th>
<th>Predominant Symptom</th>
</tr>
</thead>
<tbody>
<tr>
<td>Escitalopram</td>
<td>Global</td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>Global *</td>
</tr>
<tr>
<td>Sertraline</td>
<td>Global *</td>
</tr>
<tr>
<td>Venlafaxine</td>
<td>Global *</td>
</tr>
<tr>
<td>Venlafaxine Extended-release</td>
<td>Global *</td>
</tr>
<tr>
<td>Mirtazapine</td>
<td>Global</td>
</tr>
<tr>
<td>Duloxetine</td>
<td>Pain, Fatigue</td>
</tr>
<tr>
<td>Milnacipran</td>
<td>Pain, Fatigue</td>
</tr>
<tr>
<td>Amtriptyline</td>
<td>Pain, Fatigue</td>
</tr>
<tr>
<td>Pregabalin</td>
<td>Pain</td>
</tr>
<tr>
<td>Paroxetine Controlled-release</td>
<td>Pain</td>
</tr>
<tr>
<td>Citalopram</td>
<td>Pain</td>
</tr>
</tbody>
</table>

* Equivocal efficacy; not compared with placebo.

For additional drug information see Table 2

### Table 2. Pharmacologic Agents for Chronic Multisymptom Illness

<table>
<thead>
<tr>
<th>Agent</th>
<th>Dosage in Adults</th>
<th>Notable Adverse Effects</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Escitalopram</strong></td>
<td>10–20 mg/d</td>
<td>Headache</td>
<td>Saucidal ideation</td>
</tr>
<tr>
<td></td>
<td>Titrated up from 10 mg/d to 20 mg/d after 1 month.</td>
<td>Naussea</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Adequate Trial: 12 weeks</td>
<td>Naussea</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td><strong>Fluoxetine</strong></td>
<td>10–80 mg/d</td>
<td>Headache</td>
<td>Suicidal ideation</td>
</tr>
<tr>
<td></td>
<td>Titrated up from 10 mg/d by 10 mg/d at intervals of at least 1 week.</td>
<td>Nausea</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Adequate trial: 6–12 weeks</td>
<td>Headache</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td></td>
<td>Hepatic impairment: Use lower doses or less frequent dosing</td>
<td>Nausea</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td><strong>Sertraline</strong></td>
<td>25–350 mg/d</td>
<td>Nausea</td>
<td>Sexual dysfunction</td>
</tr>
<tr>
<td></td>
<td>Titrated up from 25 mg/d by 10 mg/d at intervals of at least 1 week.</td>
<td>Somnolence</td>
<td>Suicidal ideation</td>
</tr>
<tr>
<td></td>
<td>Adequate Trial: 12 weeks</td>
<td>Nausea</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td></td>
<td>Sexual dysfunction</td>
<td>Headache</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Headache</td>
<td>Somnolence</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td><strong>Venlafaxine</strong> XR</td>
<td>75–225 mg/d</td>
<td>Nausea</td>
<td>Sexual dysfunction</td>
</tr>
<tr>
<td></td>
<td>Titrated up by 75 mg/d at intervals of at least 1 week.</td>
<td>Somnolence</td>
<td>Suicidal ideation</td>
</tr>
<tr>
<td></td>
<td>Adequate Trial: 12 weeks</td>
<td>Nausea</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td></td>
<td>Somnolence</td>
<td>Dry mouth</td>
<td>Discontinuation syndrome</td>
</tr>
<tr>
<td></td>
<td>Headache</td>
<td>Dry mouth</td>
<td>Discontinuation syndrome</td>
</tr>
<tr>
<td></td>
<td>Fatigue</td>
<td>Increased blood pressure</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Dizziness</td>
<td>Headache</td>
<td>Discontinuation syndrome</td>
</tr>
<tr>
<td><strong>Mirtazapine</strong></td>
<td>15–60 mg/d</td>
<td>Somnolence</td>
<td>Increased cholesterol</td>
</tr>
<tr>
<td></td>
<td>Titrated up from 15 mg/d by 15 mg/d at intervals of at least 1–2 weeks</td>
<td>Dizziness</td>
<td>Neutropenia</td>
</tr>
<tr>
<td></td>
<td>Maximum: 60 mg/d</td>
<td>Headache</td>
<td>Suicidal ideation</td>
</tr>
<tr>
<td></td>
<td>Adequate Trial: 12 weeks</td>
<td>Naussea</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td></td>
<td>Somnolence</td>
<td>Dry mouth</td>
<td>Saucidal ideation</td>
</tr>
<tr>
<td></td>
<td>Insomnia</td>
<td>Increased appetite</td>
<td>Discontinuation syndrome</td>
</tr>
<tr>
<td></td>
<td>Weight gain</td>
<td>Constipation</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td><strong>Duloxetine</strong></td>
<td>60–120 mg/d</td>
<td>Nausea</td>
<td>Increased cholesterol</td>
</tr>
<tr>
<td></td>
<td>Titrated up from 20–30 mg/d by 20–10 mg/d over 2 weeks</td>
<td>Headache</td>
<td>Neutropenia</td>
</tr>
<tr>
<td></td>
<td>Adequate trial: 12 weeks</td>
<td>Naussea</td>
<td>Suicidal ideation</td>
</tr>
<tr>
<td></td>
<td>Do not ordinarily use in patients with hepatic insufficiency.</td>
<td>Dry mouth</td>
<td>Serotonin syndrome</td>
</tr>
<tr>
<td></td>
<td>Not recommended in patients with severe renal impairment (eGFR &lt;30 ml/min)</td>
<td>Somnolence</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Urinary retention</td>
<td>Constipation</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Headache</td>
<td>Constipation</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Dry mouth</td>
<td>Constipation</td>
<td>QRS-lengthening</td>
</tr>
<tr>
<td></td>
<td>Constipation</td>
<td>Insomnia</td>
<td>QRS-lengthening</td>
</tr>
</tbody>
</table>

*Conditional risk of QTc prolongation†

**Adverse Events and Comments**

- **Insomnia**: Occurs in the early weeks of treatment. Can be managed with sleep hygiene or a sedative hypnotic.
- **Dry mouth**: Can be managed with frequent fluids or sugarless gum.
- **Dizziness**: Can be managed with slow dosage increases and avoidance of alcoholic beverages.
- **Headache**: Can be managed with acetaminophen or nonsteroidal anti-inflammatory drugs (NSAIDs).
- **Nausea**: Can be managed with antiemetics or dosage reduction.
- **Fatigue**: Can be managed with increased sleep or rest.
- **Constipation**: Can be managed with dietary fiber or bulk laxatives.
- **Weight gain**: Can be managed with dietary changes or increased physical activity.
- **Increased appetite**: Can be managed with dietary changes or increased physical activity.
- **Dry mouth**: Can be managed with frequent fluids or sugarless gum.
- **Dizziness**: Can be managed with slow dosage increases and avoidance of alcoholic beverages.
- **Headache**: Can be managed with acetaminophen or nonsteroidal anti-inflammatory drugs (NSAIDs).
- **Nausea**: Can be managed with antiemetics or dosage reduction.
- **Somnolence**: Can be managed with sleep hygiene or a sedative hypnotic.
- **Angina**: Can be managed with beta-blockers or other antianginal medications.
- **Sexual dysfunction**: Can be managed with sildenafil or other phosphodiesterase 5 inhibitors.
- **Discontinuation**: Can be managed with gradual dosage tapering to avoid withdrawal symptoms.
- **Hepatotoxicity**: Can be managed with discontinuation of the medication.
- **Suicidal ideation**: Can be managed with increased social support and mental health evaluation.
- **Neutropenia**: Can be managed with discontinuation of the medication.
- **Increased cholesterol**: Can be managed with statin therapy.
- **Low dose**: Can be managed with increasing the dose or switching to a different medication.
- **QRS-lengthening**: Can be managed with discontinuation of the medication.
- **Serotonin syndrome**: Can be managed with discontinuation of the medication.
- **Increased pupillary light response**: Can be managed with discontinuation of the medication.
- **Increased appetite**: Can be managed with dietary changes or increased physical activity.
- **Increased blood pressure**: Can be managed with beta-blockers or other antihypertensive medications.
- **Skin rash**: Can be managed with discontinuation of the medication.
- **Blurred vision**: Can be managed with discontinuation of the medication.
- **Increased serum cholesterol**: Can be managed with statin therapy.
- **Anorexia**: Can be managed with dietary changes or increased physical activity.
- **Diarrhea**: Can be managed with dietary changes or increased physical activity.
- **Rhabdomyolysis**: Can be managed with discontinuation of the medication.
- **Transaminase elevation**: Can be managed with discontinuation of the medication.
- **Liver failure**: Can be managed with discontinuation of the medication.
In addition, the following other resources may be helpful:

1. War Related Illness and Injury Study Center – a National VA Post-Deployment Health Resource which provides post deployment health expertise to Veterans and their healthcare providers through clinical programs, research, education, and risk communication. Find out more here: http://www.warrelatedillness.va.gov/

2. Department of Veterans Affairs Office of Public Health – an office within the Veterans Health Administration which serves as the leader and authority in public health. Learn more about it here: http://www.publichealth.va.gov/

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For additional drug information see Table 2

### Table 2. Pharmacologic Agents for Chronic Multisymptom Illness

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<tr>
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<th>Dosage in Adults</th>
<th>Notable Adverse Effects</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Escitalopram</td>
<td>10–20 mg/d; Titrated up from 10 mg/d to 20 mg/d after 1 month; Adequate Trial: 12 weeks</td>
<td>• Headache • Nausea • Naïve hypoglycemics • Insomnia • Sexual dysfunction</td>
<td>• Suicide ideation • QTC prolongation • Serotonin syndrome</td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>10–80 mg/d; Titrated up from 10 mg/d to 10 mg/d at intervals of at least 1 week; Adequate trial: 6–12 weeks</td>
<td>• Nausea • Headache • Insomnia • Nervousness • Anxiety • Somnolence • Asthenia</td>
<td>• Diarrhea • Anorexia • Suicide ideation • Serotonin syndrome • QTC prolongation</td>
</tr>
<tr>
<td>Sertraline</td>
<td>25–350 mg/d; Titrated up from 25 mg/d by 50 mg/d at intervals of at least 1 week; Adequate Trial: 12 weeks</td>
<td>• Nausea • Somnolence • Dry mouth • Constipation • Dizziness</td>
<td>• Sexual dysfunctions • Suicide ideation • Serotonin syndrome • QTC prolongation</td>
</tr>
<tr>
<td>Venlafaxine</td>
<td>37.5–225 mg/d; Titrated up from 37.5 mg/d by 37.5–75 mg/d at intervals of at least 1 week; Adequate trial: 12 weeks</td>
<td>• Nausea • Headache • Fatigue • Dizziness • Constipation • Tremor • Dry mouth • Increased blood pressure</td>
<td>• Sexual dysfunctions • Suicide ideation • Serotonin syndrome • QTC prolongation • Discontinuation syndrome</td>
</tr>
<tr>
<td>Venlafaxine XR</td>
<td>75–225 mg/d; Titrated up by 75 mg/d at intervals of at least 1 week; Adequate Trial: 12 weeks</td>
<td>• Nausea • Headache • Fatigue • Dizziness • Constipation • Tremor • Dry mouth • Increased blood pressure</td>
<td>• Sexual dysfunctions • Suicide ideation • Serotonin syndrome • QTC prolongation • Discontinuation syndrome</td>
</tr>
<tr>
<td>Mirtazapine</td>
<td>15–60 mg/d; Titrated up from 15 mg/d by 15 mg/d at intervals of at least 1–2 weeks; Maximum: 60 mg/d; Adequate Trial: 12 weeks</td>
<td>• Somnolence • Dizziness • Dry mouth • Increased appetite • Weight gain • Constipation</td>
<td>• Increased cholesterols • Neutropenia • Suicide ideation • Serotonin syndrome</td>
</tr>
<tr>
<td>Duloxetine</td>
<td>60–120 mg/d; Titrated up from 30–30 mg/d by 20–30 mg/d over 2 weeks; Adequate trial: 12 weeks</td>
<td>• Nausea • Headache • Dry mouth • Fatigue • Somnolence • Constipation • Insomnia</td>
<td>• Urinary retention • Serotonin syndrome • Suicide ideation • Hepatotoxicity</td>
</tr>
</tbody>
</table>

* Equivocal efficacy; not compared with placebo.

For additional drug information see Table 2.
<table>
<thead>
<tr>
<th>Agent</th>
<th>Dosage in Adults</th>
<th>Notable Adverse Effects</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Milnacipran</td>
<td>100 mg/d (100-200 mg/d) in 2 divided doses; Titrate up from 12.5 mg by 12.5–50 mg/d per week over 3–4 weeks Adequate trial: 12 weeks Dose in patients with severe renal impairment (5–29 ml/min): 50–100 mg/d in 2 divided doses Do not ordinarily use in patients with substantial alcohol use or chronic liver disease. Not recommended in patients with end-stage renal disease.</td>
<td>• Nausea • Headache • Constipation • Insomnia • Dizziness • Hot flush • Somnolence • Suicidal ideation • Increased blood pressure and heart rate • Urinary retention • Hepatotoxicity • Withdrawal symptoms</td>
<td>Contraindicated with MAOIs and within 14 days of starting or stopping MAOIs MAOIs contraindicated within 5 days of discontinuing milnacipran Contraindicated in patients with uncontrolled narrow-angle glaucoma</td>
</tr>
<tr>
<td>Amitriptyline</td>
<td>10-50 mg daily Adequate trial: 6-8 weeks Use lower doses in the elderly</td>
<td>• Dry mouth • Fatigue • Sedation • Vannagal reaction • Orthostatic hypotension</td>
<td>Contraindicated with MAOIs and within 14 days of starting or stopping MAOIs Contraindicated with cisapride Avoid use with QTc prolonging drugs, anticholinergics Use with caution in patients with cardio- or cerebrovascular disease</td>
</tr>
<tr>
<td>Pregabalin</td>
<td>300–450 mg/d divided BID-TID; Starting at 150 mg/d and increasing by 100 mg/d every week Adequate trial: 8 weeks Adjust dose based on renal function</td>
<td>• Dizziness • Somnolence • Headache • Weight gain • Angioedema • Suicidal ideation • Peripheral edema • Withdrawal symptoms • Blurred vision; visual field changes</td>
<td>Dose of 600 mg/d was studied but showed no additional benefit and increased the risk of adverse events</td>
</tr>
<tr>
<td>Paroxetine CR</td>
<td>62.5 mg/d (12.5–75 mg/d), Starting at 25 mg/d and increasing by 100 mg/d at intervals of at least 1 week Adequate trial: 12 weeks Severe renal impairment (CrCl &lt;30 ml/min) or severe hepatic impairment: Use lower starting dose Elderly 12.5–50 mg/d</td>
<td>• Drowsiness • Nausea • Insomnia • Headache • Ejaculatory disorder • Dizziness • Decreased libido • Diaphoresis • Weakness • Constipation • Diarrhea • Dry mouth • Akathisia • Suicidal ideation • Somnolecne</td>
<td>Contraindicated with MAOIs and within 14 days of starting or stopping MAOIs Most sedating SSRI Potent anticholinergic effects</td>
</tr>
<tr>
<td>Citalopram</td>
<td>20–40 mg/d; Titrate up at intervals of at least 1 week Adequate trial: 8–16 weeks Elderly (&gt;60 y) and Hepatic impairment: Max 20 mg/d</td>
<td>• Drowsiness • Nausea • Dry mouth • Somnolence • Insomnia • Hyperhidrosis</td>
<td>Contraindicated with MAOIs and within 14 days of starting or stopping MAOIs Avoid using citalopram with other QTc prolonging drugs</td>
</tr>
</tbody>
</table>

Refer to current Product Information for additional prescribing information.

For relative usage and timing of therapies refer to the full clinical practice guideline for management of CMI at: [http://www.healthquality.va.gov/](http://www.healthquality.va.gov/)† Associated with risk of torsade de pointes in the presence of other risk factors for QTc prolongation (e.g. high dose, hypokalemia, hypomagnesemia, drug interaction or congenital long QT).