CDC Guideline for Prescribing Opioids for Chronic Pain — United States, 2016
Disclosure of Relationship

The Core Expert Group (CEG) members disclose that they have no financial conflicts of interest. Experts disclose the following activities related to the content of this guideline: Pam Archer discloses authorship of the Oklahoma Emergency Department and Urgent Care Clinic Opioid Prescribing Guidelines and the Opioid Prescribing Guidelines for Oklahoma Health Care Providers in the Office Based Setting; Bonnie Burman discloses authorship of the Ohio Guidelines for Prescribing Opioids for the Treatment of Chronic, Non-Terminal Pain; Jane Ballantyne discloses that she has served as a paid consultant to Cohen Milstein Sellers & Toll, PLLC, and has special advisory committee responsibilities on the Food and Drug Administration (FDA) Risk Evaluation and Mitigation Strategies committee; Phillip Coffin discloses that in 2012 he provided expert testimony to the California State Assembly regarding a bill to expand naloxone access and reports that he is the principal investigator on a research study of methamphetamine dependence that receives donated injectable naltrexone from Alkermes, Inc.; Gary Franklin discloses authorship of the AMDG Interagency Guideline on Prescribing Opioids for Pain; Erin Krebs discloses that she represented from Alkermes, Inc.; Gary Franklin discloses authorship of the AMDG Interagency Guideline on Prescribing Opioids for Pain; Erin Krebs discloses that she represented the American College of Physicians at a 2014 Food and Drug Administration meeting on Abuse Deterrent Opioid Formulations; Lewis Nelson discloses his ad-hoc membership on the FDA Drug Safety and Risk Management Advisory Committee; Trupti Patel discloses authorship of the Arizona Opioid Prescribing Guidelines; Robert “Chuck” Rich discloses that he was an author of the 2013 American Academy of Family Physicians position paper on opioids and pain management; Joanna Starrels discloses that she received honoraria from the Betty Ford Institute; Thomas Tape discloses that he was an author of the 2013 American College of Physicians policy position paper on prescription drug abuse. CDC provided 100% of the funding for the supplemental evidence review tasks and meeting support. No foundation or industry support was accepted.

The Opioid Guideline Workgroup (OGW) members disclose that they have no financial conflicts of interest. Experts disclose the following activities related to the content of this guideline: Anne Burns discloses that she participated in a congressional briefing sponsored by Reps. Carter and DeSaulnier on the pharmacist’s role of furnishing Naloxone and that she participates on the National Advisory Board for the Prescription Drug Abuse and Heroin Summit. Chinazo Cunningham discloses that her husband is employed by Quest Diagnostics and Dr. Cunningham was recused from any discussion related to urine drug testing. Traci Green discloses that she was previously employed by Inflexion, a small business that conducts Small Business Innovation Research on behavioral interventions for behavioral health and chronic pain and created several psychometric tools for conducting risk assessment for prescription opioid abuse potential. Dr. Green also discloses that while at the hospital where she is employed, she provided consultation to Purdue Pharma Ltd to design overdose prevention brochures for persons who use diverted prescription opioids non-medically with an emphasis on persons who inject prescription drugs, and not for patients using opioid therapy for pain. Dr. Green was recused from any discussion related to risk assessment tools and patient education materials. Erin Krebs discloses that she served on the CDC Opioid Prescribing Guideline CEG. Christina Porucznik discloses that she served on the Opioid Guideline Workgroup. Greg Terman discloses that he serves as the President of the American Pain Society. Mark Wallace discloses that he served on a Kpharma advisory panel for an abuse-deterrent hydrocodone formulation to treat acute postoperative pain and Dr. Wallace was recused form any discussion related to abuse-deterrent drugs.

The NCIPC Board of Scientific Counselors (BSC) members disclose that they have no financial conflicts of interest. Two BSC members, Traci Green and Christina Porucznik, served on the Opioid Guideline Workgroup. Traci Green discloses that she was previously employed by Inflexion, a small business that conducts Small Business Innovation Research on behavioral interventions for behavioral health and chronic pain and created several psychometric tools for conducting risk assessment for prescription opioid abuse potential. Dr. Green also discloses that while at the hospital where she is employed, she provided consultation to Purdue Pharma Ltd to design overdose prevention brochures for persons who use diverted prescription opioids non-medically with an emphasis on persons who inject prescription drugs, and not for patients using opioid therapy for pain. Dr. Green was recused from any discussion related to risk assessment tools and patient education materials. Christina Porucznik discloses that she served on the CDC Opioid Prescribing Guideline CEG.
This guideline provides recommendations for primary care clinicians who are prescribing opioids for chronic pain outside of active cancer treatment, palliative care, and end-of-life care. The guideline addresses 1) when to initiate or continue opioids for chronic pain; 2) opioid selection, dosage, duration, follow-up, and discontinuation; and 3) assessing risk and addressing harms of opioid use. CDC developed the guideline using the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) framework, and recommendations are made on the basis of a systematic review of the scientific evidence while considering benefits and harms, values and preferences, and resource allocation. CDC obtained input from experts, stakeholders, the public, peer reviewers, and a federally chartered advisory committee. It is important that patients receive appropriate pain treatment with careful consideration of the benefits and risks of treatment options. This guideline is intended to improve communication between clinicians and patients about the risks and benefits of opioid therapy for chronic pain, improve the safety and effectiveness of pain treatment, and reduce the risks associated with long-term opioid therapy, including opioid use disorder, overdose, and death. CDC has provided a checklist for prescribing opioids for chronic pain (http://stacks.cdc.gov/view/cdc/38025) as well as a website (http://www.cdc.gov/drugoverdose/prescribingresources.html) with additional tools to guide clinicians in implementing the recommendations.

Introduction

Background

Opioids are commonly prescribed for pain. An estimated 20% of patients presenting to physician offices with noncancer pain symptoms or pain-related diagnoses (including acute and chronic pain) receive an opioid prescription (1). In 2012, health care providers wrote 259 million prescriptions for opioid pain medication, enough for every adult in the United States to have a bottle of pills (2). Opioid prescriptions per capita increased 7.3% from 2007 to 2012, with opioid prescribing rates increasing more for family practice, general practice, and internal medicine compared with other specialties (3). Rates of opioid prescribing vary greatly across states in ways that cannot be explained by the underlying health status of the population, highlighting the lack of consensus among clinicians on how to use opioid pain medication (2).

Prevention, assessment, and treatment of chronic pain are challenges for health providers and systems. Pain might go unrecognized, and patients, particularly members of racial and ethnic minority groups, women, the elderly, persons with cognitive impairment, and those with cancer and at the end of life, can be at risk for inadequate pain treatment (4). Patients can experience persistent pain that is not well controlled. There are clinical, psychological, and social consequences associated with chronic pain including limitations in complex activities, lost work productivity, reduced quality of life, and stigma, emphasizing the importance of appropriate and compassionate patient care (4). Patients should receive appropriate pain treatment based on a careful consideration of the benefits and risks of treatment options.

Chronic pain has been variably defined but is defined within this guideline as pain that typically lasts >3 months or past the time of normal tissue healing (5). Chronic pain can be the result of an underlying medical disease or condition, injury, medical treatment, inflammation, or an unknown cause (4). Estimates of the prevalence of chronic pain vary, but it is clear that the number of persons experiencing chronic pain in the United States is substantial. The 1999–2002 National Health and Nutrition Examination Survey estimated that 14.6% of adults have current widespread or localized pain lasting at least 3 months (6). Based on a survey conducted during 2001–2003 (7), the overall prevalence of common, predominantly musculoskeletal pain conditions (e.g., arthritis, rheumatism, chronic back or neck problems, and frequent severe headaches) was estimated at 43% among adults in the United States. In 2012, health care providers wrote 259 million prescriptions for opioid pain medication, enough for every adult in the United States to have a bottle of pills (2). Opioid prescriptions per capita increased 7.3% from 2007 to 2012, with opioid prescribing rates increasing more for family practice, general practice, and internal medicine compared with other specialties (3). Rates of opioid prescribing vary greatly across states in ways that cannot be explained by the underlying health status of the population, highlighting the lack of consensus among clinicians on how to use opioid pain medication (2).

Prevention, assessment, and treatment of chronic pain are challenges for health providers and systems. Pain might go unrecognized, and patients, particularly members of racial and ethnic minority groups, women, the elderly, persons with cognitive impairment, and those with cancer and at the end of life, can be at risk for inadequate pain treatment (4). Patients can experience persistent pain that is not well controlled. There are clinical, psychological, and social consequences associated with chronic pain including limitations in complex activities, lost work productivity, reduced quality of life, and stigma, emphasizing the importance of appropriate and compassionate patient care (4). Patients should receive appropriate pain treatment based on a careful consideration of the benefits and risks of treatment options.

Chronic pain has been variably defined but is defined within this guideline as pain that typically lasts >3 months or past the time of normal tissue healing (5). Chronic pain can be the result of an underlying medical disease or condition, injury, medical treatment, inflammation, or an unknown cause (4). Estimates of the prevalence of chronic pain vary, but it is clear that the number of persons experiencing chronic pain in the United States is substantial. The 1999–2002 National Health and Nutrition Examination Survey estimated that 14.6% of adults have current widespread or localized pain lasting at least 3 months (6). Based on a survey conducted during 2001–2003 (7), the overall prevalence of common, predominantly musculoskeletal pain conditions (e.g., arthritis, rheumatism, chronic back or neck problems, and frequent severe headaches) was estimated at 43% among adults in the United States.

Corresponding author: Deborah Dowell, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, CDC. E-mail: gdo7@cdc.gov.
United States, although minimum duration of symptoms was not specified. Most recently, analysis of data from the 2012 National Health Interview Study showed that 11.2% of adults report having daily pain (8). Clinicians should consider the full range of therapeutic options for the treatment of chronic pain. However, it is hard to estimate the number of persons who could potentially benefit from opioid pain medication long term. Evidence supports short-term efficacy of opioids for reducing pain and improving function in noncancer nociceptive and neuropathic pain in randomized clinical trials lasting primarily ≤12 weeks (9,10), and patients receiving opioid therapy for chronic pain report some pain relief when surveyed (11–13). However, few studies have been conducted to rigorously assess the long-term benefits of opioids for chronic pain (pain lasting >3 months) with outcomes examined at least 1 year later (14). On the basis of data available from health systems, researchers estimate that 9.6–11.5 million adults, or approximately 3%–4% of the adult U.S. population, were prescribed long-term opioid therapy in 2005 (15).

Opioid pain medication use presents serious risks, including overdose and opioid use disorder. From 1999 to 2014, more than 165,000 persons died from overdose related to opioid pain medication in the United States (16). In the past decade, while the death rates for the top leading causes of death such as heart disease and cancer have decreased substantially, the death rate associated with opioid pain medication has increased markedly (17). Sales of opioid pain medication have increased in parallel with opioid-related overdose deaths (18). The Drug Abuse Warning Network estimated that >420,000 emergency department visits were related to the misuse or abuse of narcotic pain relievers in 2011, the most recent year for which data are available (19). Although clinical criteria have varied over time, opioid use disorder is a problematic pattern of opioid use leading to clinically significant impairment or distress. This disorder is manifested by specific criteria such as unsuccessful efforts to cut down or control use and use resulting in social problems and a failure to fulfill major role obligations at work, school, or home (20). This diagnosis has also been referred to as “abuse or dependence” and “addiction” in the literature, and is different from tolerance (diminished response to a drug with repeated use) and physical dependence (adaptation to a drug that produces symptoms of withdrawal when the drug is stopped), both of which can exist without a diagnosed disorder. In 2013, on the basis of DSM-IV diagnosis criteria, an estimated 1.9 million persons abused or were dependent on prescription opioid pain medication (21). Having a history of a prescription for an opioid pain medication increases the risk for overdose and opioid use disorder (22–24), highlighting the value of guidance on safer prescribing practices for clinicians. For example, a recent study of patients aged 15–64 years receiving opioids for chronic noncancer pain and followed for up to 13 years revealed that one in 550 patients died from opioid-related overdose at a median of 2.6 years from their first opioid prescription, and one in 32 patients who escalated to opioid dosages >200 morphine milligram equivalents (MME) died from opioid-related overdose (25).

This guideline provides recommendations for the prescribing of opioid pain medication by primary care clinicians for chronic pain (i.e., pain conditions that typically last >3 months or past the time of normal tissue healing) in outpatient settings outside of active cancer treatment, palliative care, and end-of-life care. Although the guideline does not focus broadly on pain management, appropriate use of long-term opioid therapy must be considered within the context of all pain management strategies (including nonopioid pain medications and nonpharmacologic treatments). CDC’s recommendations are made on the basis of a systematic review of the best available evidence, along with input from experts, and further review and deliberation by a federally chartered advisory committee.

The guideline is intended to ensure that clinicians and patients consider safer and more effective treatment, improve patient outcomes such as reduced pain and improved function, and reduce the number of persons who develop opioid use disorder, overdose, or experience other adverse events related to these drugs. Clinical decision making should be based on a relationship between the clinician and patient, and an understanding of the patient’s clinical situation, functioning, and life context. The recommendations in the guideline are voluntary, rather than prescriptive standards. They are based on emerging evidence, including observational studies or randomized clinical trials with notable limitations. Clinicians should consider the circumstances and unique needs of each patient when providing care.

**Rationale**

Primary care clinicians report having concerns about opioid pain medication misuse, find managing patients with chronic pain stressful, express concern about patient addiction, and report insufficient training in prescribing opioids (26). Across specialties, physicians believe that opioid pain medication can be effective in controlling pain, that addiction is a common consequence of prolonged use, and that long-term opioid therapy often is overprescribed for patients with chronic noncancer pain (27). These attitudes and beliefs, combined with increasing trends in opioid-related overdose, underscore the need for better clinician guidance on opioid prescribing. Clinical practice guidelines focused on prescribing can improve clinician knowledge, change prescribing practices (28), and ultimately benefit patient health.
Professional organizations, states, and federal agencies (e.g., the American Pain Society/American Academy of Pain Medicine, 2009; the Washington Agency Medical Directors Group, 2015; and the U.S. Department of Veterans Affairs/Department of Defense, 2010) have developed guidelines for opioid prescribing (29–31). Existing guidelines share some common elements, including dosing thresholds, cautious titration, and risk mitigation strategies such as using risk assessment tools, treatment agreements, and urine drug testing. However, there is considerable variability in the specific recommendations (e.g., range of dosing thresholds of 90 MME/day to 200 MME/day), audience (e.g., primary care clinicians versus specialists), use of evidence (e.g., systematic review, grading of evidence and recommendations, and role of expert opinion), and rigor of methods for addressing conflict of interest (32). Most guidelines, especially those that are not based on evidence from scientific studies published in 2010 or later, also do not reflect the most recent scientific evidence about risks related to opioid dosage.

This CDC guideline offers clarity on recommendations based on the most recent scientific evidence, informed by expert opinion and stakeholder and public input. Scientific research has identified high-risk prescribing practices that have contributed to the overdose epidemic (e.g., high-dose prescribing, overlapping opioid and benzodiazepine prescriptions, and extended-release/long-acting [ER/LA] opioids for acute pain) (24,33,34). Using guidelines to address problematic prescribing has the potential to optimize care and improve patient safety based on evidence-based practice (28), as well as reverse the cycle of opioid pain medication misuse that contributes to the opioid overdose epidemic.

Scope and Audience

This guideline is intended for primary care clinicians (e.g., family physicians and internists) who are treating patients with chronic pain (i.e., pain lasting >3 months or past the time of normal tissue healing) in outpatient settings. Prescriptions by primary care clinicians account for nearly half of all dispensed opioid prescriptions, and the growth in prescribing rates among these clinicians has been above average (3). Primary care clinicians include physicians as well as nurse practitioners and physician assistants. Although the focus is on primary care clinicians, because clinicians work within team-based care, the recommendations refer to and promote integrated pain management and collaborative working relationships with other providers (e.g., behavioral health providers, pharmacists, and pain management specialists). Although the transition from use of opioid therapy for acute pain to use for chronic pain is hard to predict and identify, the guideline is intended to inform clinicians who are considering prescribing opioid pain medication for painful conditions that can or have become chronic.

This guideline is intended to apply to patients aged ≥18 years with chronic pain outside of palliative and end-of-life care. For this guideline, palliative care is defined in a manner consistent with that of the Institute of Medicine as care that provides relief from pain and other symptoms, supports quality of life, and is focused on patients with serious advanced illness. Palliative care can begin early in the course of treatment for any serious illness that requires excellent management of pain or other distressing symptoms (35). End-of-life care is defined as care for persons with a terminal illness or at high risk for dying in the near future in hospice care, hospitals, long-term care settings, or at home. Patients within the scope of this guideline include cancer survivors with chronic pain who have completed cancer treatment, are in clinical remission, and are under cancer surveillance only. The guideline is not intended for patients undergoing active cancer treatment, palliative care, or end-of-life care because of the unique therapeutic goals, ethical considerations, opportunities for medical supervision, and balance of risks and benefits with opioid therapy in such care.

The recommendations address the use of opioid pain medication in certain special populations (e.g., older adults and pregnant women) and in populations with conditions posing special risks (e.g., a history of substance use disorder). The recommendations do not address the use of opioid pain medication in children or adolescents aged <18 years. The available evidence concerning the benefits and harms of long-term opioid therapy in children and adolescents is limited, and few opioid medications provide information on the label regarding safety and effectiveness in pediatric patients. However, observational research shows significant increases in opioid prescriptions for pediatric populations from 2001 to 2010 (36), and a large proportion of adolescents are commonly prescribed opioid pain medications for conditions such as headache and sports injuries (e.g., in one study, 50% of adolescents presenting with headache received a prescription for an opioid pain medication [37,38]). Adolescents who misuse opioid pain medication often misuse medications from their own previous prescriptions (39), with an estimated 20% of adolescents with currently prescribed opioid medications reporting using them intentionally to get high or increase the effects of alcohol or other drugs (40). Use of prescribed opioid pain medication before high school graduation is associated with a 33% increase in the risk of later opioid misuse (41). Misuse of opioid pain medications in adolescence strongly predicts later onset of heroin use (42). Thus, risk of opioid medication use in pediatric populations is of great concern. Additional clinical trial and observational research is needed,
and encouraged, to inform development of future guidelines for this critical population.

The recommendations are not intended to provide guidance on use of opioids as part of medication-assisted treatment for opioid use disorder. Some of the recommendations might be relevant for acute care settings or other specialists, such as emergency physicians or dentists, but use in these settings or by other specialists is not the focus of this guideline. Readers are referred to other sources for prescribing recommendations within acute care settings and in dental practice, such as the American College of Emergency Physicians’ guideline for prescribing of opioids in the emergency department (43); the American Society of Anesthesiologists’ guideline for acute pain management in the perioperative setting (44); the Washington Agency Medical Directors’ Group Interagency Guideline on Prescribing Opioids for Pain, Part II: Prescribing Opioids in the Acute and Subacute Phase (30); and the Pennsylvania Guidelines on the Use of Opioids in Dental Practice (45). In addition, given the challenges of managing the painful complications of sickle cell disease, readers are referred to the NIH National Heart, Lung, and Blood Institute’s Evidence Based Management of Sickle Cell Disease Expert Panel Report for management of sickle cell disease (46).

**Guideline Development Methods**

**Guideline Development Using the Grading of Recommendations Assessment, Development, and Evaluation Method**

CDC developed this guideline using the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) method (http://www.gradeworkinggroup.org). This method specifies the systematic review of scientific evidence and offers a transparent approach to grading quality of evidence and strength of recommendations. The method has been adapted by the CDC Advisory Committee on Immunization Practices (ACIP) (47). CDC has applied the ACIP translation of the GRADE framework in this guideline. Within the ACIP GRADE framework, the body of evidence is categorized in a hierarchy. This hierarchy reflects degree of confidence in the effect of a clinical action on health outcomes. The categories include type 1 evidence (randomized clinical trials or overwhelming evidence from observational studies), type 2 evidence (randomized clinical trials with important limitations, or exceptionally strong evidence from observational studies), type 3 evidence (observational studies or randomized clinical trials with notable limitations), and type 4 evidence (clinical experience and observations, observational studies with important limitations, or randomized clinical trials with several major limitations). Type of evidence is categorized by study design as well as limitations in study design or implementation, imprecision of estimates, variability in findings, indirectness of evidence, publication bias, magnitude of treatment effects, dose-response gradient, and a constellation of plausible biases that could change observations of effects. Type 1 evidence indicates that one can be very confident that the true effect lies close to that of the estimate of the effect; type 2 evidence means that the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different; type 3 evidence means that confidence in the effect estimate is limited and the true effect might be substantially different from the estimate of the effect; and type 4 evidence indicates that one has very little confidence in the effect estimate, and the true effect is likely to be substantially different from the estimate of the effect (47,48). When no studies are present, evidence is considered to be insufficient. The ACIP GRADE framework places recommendations in two categories, Category A and Category B. Four major factors determine the category of the recommendation: the quality of evidence, the balance between desirable and undesirable effects, values and preferences, and resource allocation (cost). Category A recommendations apply to all persons in a specified group and indicate that most patients should receive the recommended course of action. Category B recommendations indicate that there should be individual decision making; different choices will be appropriate for different patients, so clinicians must help patients arrive at a decision consistent with patient values and preferences, and specific clinical situations (47). According to the GRADE methodology, a particular quality of evidence does not necessarily imply a particular strength of recommendation (48–50). Category A recommendations can be made based on type 3 or type 4 evidence when the advantages of a clinical action greatly outweigh the disadvantages based on a consideration of benefits and harms, values and preferences, and costs. Category B recommendations are made when the advantages and disadvantages of a clinical action are more balanced. GRADE methodology is discussed extensively elsewhere (47,51). The U.S. Preventive Services Task Force (USPSTF) follows different methods for developing and categorizing recommendations (http://www.uspreventiveservicestaskforce.org). USPSTF recommendations focus on preventive services and are categorized as A, B, C, D, and I. Under the Affordable Care Act, all “nongrandfathered” health plans (that is, those health plans not in existence prior to March 23, 2010 or those with significant changes to their coverage) and expanded Medicaid plans are required to cover
preventive services recommended by USPSTF with a category A or B rating with no cost sharing. The coverage requirements went into effect September 23, 2010. Similar requirements are in place for vaccinations recommended by ACIP, but do not exist for other recommendations made by CDC, including recommendations within this guideline.

A previously published systematic review sponsored by the Agency for Healthcare Research and Quality (AHRQ) on the effectiveness and risks of long-term opioid treatment of chronic pain (14,52) initially served to directly inform the recommendation statements. This systematic clinical evidence review addressed the effectiveness of long-term opioid therapy for outcomes related to pain, function, and quality of life; the comparative effectiveness of different methods for initiating and titrating opioids; the harms and adverse events associated with opioids; and the accuracy of risk-prediction instruments and effectiveness of risk mitigation strategies on outcomes related to overdose, addiction, abuse, or misuse. For the current guideline development, CDC conducted additional literature searches to update the evidence review to include more recently available publications and to answer an additional clinical question about the effect of opioid therapy for acute pain on long-term use. More details about the literature search strategies and GRADE methods applied are provided in the Clinical Evidence Review (http://stacks.cdc.gov/view/cdc/38026). CDC developed GRADE evidence tables to illustrate the quality of the evidence for each clinical question.

As identified in the AHRQ-sponsored clinical evidence review, the overall evidence base for the effectiveness and risks of long-term opioid therapy is low in quality per the GRADE criteria. Thus, contextual evidence is needed to provide information about the benefits and harms of nonpharmacologic and nonopioid pharmacologic therapy and the epidemiology of opioid pain medication overdose and inform the recommendations. Further, as elucidated by the GRADE Working Group, supplemental information on clinician and patient values and preferences and resource allocation can inform judgments of benefits and harms and be helpful for translating the evidence into recommendations. CDC conducted a contextual evidence review to supplement the clinical evidence review based on systematic searches of the literature. The review focused on the following four areas: effectiveness of nonpharmacologic and nonopioid pharmacologic treatments; benefits and harms related to opioid therapy (including additional studies not included in the clinical evidence review such as studies that evaluated outcomes at any duration or used observational study designs related to specific opioid pain medications, high-dose opioid therapy, co-prescription of opioids with other controlled substances, duration of opioid use, special populations, risk stratification/mitigation approaches, and effectiveness of treatments for addressing potential harms of opioid therapy); clinician and patient values and preferences; and resource allocation. CDC constructed narrative summaries of this contextual evidence and used the information to support the clinical recommendations. More details on methods for the contextual evidence review are provided in the Contextual Evidence Review (http://stacks.cdc.gov/view/cdc/38027).

On the basis of a review of the clinical and contextual evidence (review methods are described in more detail in subsequent sections of this report), CDC drafted recommendation statements focused on determining when to initiate or continue opioids for chronic pain; opioid selection, dosage, duration, follow-up, and discontinuation; and assessing risk and addressing harms of opioid use. To help assure the draft guideline’s integrity and credibility, CDC then began a multistep review process to obtain input from experts, stakeholders, and the public to help refine the recommendations.

**Solicitation of Expert Opinion**

CDC sought the input of experts to assist in reviewing the evidence and providing perspective on how CDC used the evidence to develop the draft recommendations. These experts, referred to as the “Core Expert Group” (CEG) included subject matter experts, representatives of primary care professional societies and state agencies, and an expert in guideline development methodology.* CDC identified subject matter experts with high scientific standing; appropriate academic and clinical training and relevant clinical experience; and proven scientific excellence in opioid prescribing, substance use disorder treatment, and pain management. CDC identified representatives from leading primary care professional organizations to represent the audience for this guideline. Finally, CDC identified state agency officials and representatives based on their experience with state guidelines for opioid prescribing that were developed with multiple agency stakeholders and informed by scientific literature and existing evidence-based guidelines.

Prior to their participation, CDC asked potential experts to reveal possible conflicts of interest such as financial relationships with industry, intellectual preconceptions, or previously stated public positions. Experts could not serve if they had conflicts that might have a direct and predictable effect on the recommendations. CDC excluded experts who had a financial or promotional relationship with a company.

---

* A list of the members appears at the end of this report. The recommendations and all statements included in this guideline are those of CDC and do not necessarily represent the official position of any persons or organizations providing comments on the draft guideline.
that makes a product that might be affected by the guideline. CDC reviewed potential nonfinancial conflicts carefully (e.g., intellectual property, travel, public statements or positions such as congressional testimony) to determine if the activities would have a direct and predictable effect on the recommendations. CDC determined the risk of these types of activities to be minimal for the identified experts. All experts completed a statement certifying that there was no potential or actual conflict of interest. Activities that did not pose a conflict (e.g., participation in Food and Drug Administration [FDA] activities or other guideline efforts) are disclosed.

CDC provided to each expert written summaries of the scientific evidence (both the clinical and contextual evidence reviews conducted for this guideline) and CDC’s draft recommendation statements. Experts provided individual ratings for each draft recommendation statement based on the balance of benefits and harms, evidence strength, certainty of values and preferences, cost, recommendation strength, rationale, importance, clarity, and ease of implementation. CDC hosted an in-person meeting of the experts that was held on June 23–24, 2015, in Atlanta, Georgia, to seek their views on the evidence and draft recommendations and to better understand their premeeting ratings. CDC sought the experts’ individual opinions at the meeting. Although there was widespread agreement on some of the recommendations, there was disagreement on others. Experts did not vote on the recommendations or seek to come to a consensus. Decisions about recommendations to be included in the guideline, and their rationale, were made by CDC. After revising the guideline, CDC sent written copies of it to each of the experts for review and asked for any additional comments; CDC reviewed these written comments and considered them when making further revisions to the draft guideline. The experts have not reviewed the final version of the guideline.

**Federal Partner Engagement**

Given the scope of this guideline and the interest of agencies across the federal government in appropriate pain management, opioid prescribing, and related outcomes, CDC invited its National Institute of Occupational Safety and Health and CDC’s federal partners to observe the expert meeting, provide written comments on the full draft guideline after the meeting, and review the guideline through an agency clearance process; CDC reviewed comments and incorporated changes. Interagency collaboration will be critical for translating these recommendations into clinical practice. Federal partners included representatives from the Substance Abuse and Mental Health Services Administration, the National Institute on Drug Abuse, FDA, the U.S. Department of Veterans Affairs, the U.S. Department of Defense, the Office of the National Coordinator for Health Information Technology, the Centers for Medicare and Medicaid Services, the Health Resources and Services Administration, AHRQ, and the Office of National Drug Control Policy.

**Stakeholder Comment**

Given the importance of the guideline for a wide variety of stakeholders, CDC also invited review from a Stakeholder Review Group (SRG) to provide comment so that CDC could consider modifications that would improve the recommendations’ specificity, applicability, and ease of implementation. The SRG included representatives from professional organizations that represent specialties that commonly prescribe opioids (e.g., pain medicine, physical medicine and rehabilitation), delivery systems within which opioid prescribing occurs (e.g., hospitals), and representation from community organizations with interests in pain management and opioid prescribing.* Representatives from each of the SRG organizations were provided a copy of the guideline for comment. Each of these representatives provided written comments. Once input was received from the full SRG, CDC reviewed all comments and carefully considered them when revising the draft guideline.

**Constituent Engagement**

To obtain initial perspectives from constituents on the recommendation statements, including clinicians and prospective patients, CDC convened a constituent engagement webinar and circulated information about the webinar in advance through announcements to partners. CDC hosted the webinar on September 16 and 17, 2015, provided information about the methodology for developing the guideline, and presented the key recommendations. A fact sheet was posted on the CDC Injury Center website (http://www.cdc.gov/injury) summarizing the guideline development process and clinical practice areas addressed in the guideline; instructions were included on how to submit comments via email. CDC received comments during and for 2 days following the first webinar. Over 1,200 constituent comments were received. Comments were reviewed and carefully considered when revising the draft guideline.

**Peer Review**

Per the final information quality bulletin for peer review (https://www.whitehouse.gov/sites/default/files/omb/memoranda/fy2005/m05-03.pdf), peer review requirements applied to this guideline because it provides influential
scientific information that could have a clear and substantial impact on public- and private-sector decisions. Three experts independently reviewed the guideline to determine the reasonableness and strength of recommendations; the clarity with which scientific uncertainties were clearly identified; and the rationale, importance, clarity, and ease of implementation of the recommendations.* CDC selected peer reviewers based on expertise, diversity of scientific viewpoints, and independence from the guideline development process. CDC assessed and managed potential conflicts of interest using a process similar to the one as described for solicitation of expert opinion. No financial interests were identified in the disclosure and review process, and nonfinancial activities were determined to be of minimal risk; thus, no significant conflict of interest concerns were identified. CDC placed the names of peer reviewers on the CDC and the National Center for Injury Prevention and Control Peer Review Agenda websites that are used to provide information about the peer review of influential documents. CDC reviewed peer review comments and revised the draft guideline accordingly.

**Public Comment**

To obtain comments from the public on the full guideline, CDC published a notice in the Federal Register (80 FR 77351) announcing the availability of the guideline and the supporting clinical and contextual evidence reviews for public comment. The comment period closed January 13, 2016. CDC received more than 4,350 comments from the general public, including patients with chronic pain, clinicians, families who have lost loved ones to overdose, medical associations, professional organizations, academic institutions, state and local governments, and industry. CDC reviewed each of the comments and carefully considered them when revising the draft guideline.

**Federal Advisory Committee Review and Recommendation**

The National Center for Injury Prevention and Control (NCIPC) Board of Scientific Counselors (BSC) is a federal advisory committee that advises and makes recommendations to the Secretary of the Department of Health and Human Services, the Director of CDC, and the Director of NCIPC.* The BSC makes recommendations regarding policies, strategies, objectives, and priorities, and reviews progress toward injury and violence prevention. CDC sought the BSC’s advice on the draft guideline. BSC members are special government employees appointed as CDC advisory committee members; as such, all members completed an OGE Form 450 to disclose relevant interests. BSC members also reported on their disclosures during meetings. Disclosures for the BSC are reported in the guideline.

To assist in guideline review, on December 14, 2015, via Federal Register notice, CDC announced the intent to form an Opioid Guideline Workgroup (OGW) to provide observations on the draft guideline to the BSC. CDC provided the BSC with the draft guideline as well as summaries of comments provided to CDC by stakeholders, constituents, and peer reviewers, and edits made to the draft guideline in response. During an open meeting held on January 7, 2016, the BSC recommended the formation of the OGW. The OGW included a balance of perspectives from audiences directly affected by the guideline, audiences that would be directly involved with implementing the recommendations, and audiences qualified to provide representation. The OGW comprised clinicians, subject matter experts, and a patient representative, with the following perspectives represented: primary care, pain medicine, public health, behavioral health, substance abuse treatment, pharmacy, patients, and research.* Additional sought-after attributes were appropriate academic and clinical training and relevant clinical experience; high scientific standing; and knowledge of the patient, clinician, and caregiver perspectives. In accordance with CDC policy, two BSC committee members also served as OGW members, with one serving as the OGW Chair. The professional credentials and interests of OGW members were carefully reviewed to identify possible conflicts of interest such as financial relationships with industry, intellectual preconceptions, or previously stated public positions. Only OGW members whose interests were determined to be minimal were selected. When an activity was perceived as having the potential to affect a specific aspect of the recommendations, the activity was disclosed, and the OGW member was recused from discussions related to that specific aspect of the recommendations (e.g., urine drug testing and abuse-deterrent formulations). Disclosures for the OGW are reported. CDC and the OGW identified ad-hoc consultants to supplement the workgroup expertise, when needed, in the areas of pediatrics, occupational medicine, obstetrics and gynecology, medical ethics, addiction psychiatry, physical medicine and rehabilitation, guideline development methodology, and the perspective of a family member who lost a loved one to opioid use disorder or overdose.

The BSC charged the OGW with reviewing the quality of the clinical and contextual evidence reviews and reviewing each of the recommendation statements and accompanying rationales. For each recommendation statement, the OGW considered the quality of the evidence, the balance of benefits and risks, the values and preferences of clinicians and patients, the cost feasibility, and the category designation...
NCIPC announced an open meeting of the NCIPC BSC in the Federal Register on January 11, 2015. The BSC met on January 28, 2016, to discuss the OGW report and deliberate on the draft guideline itself. Members of the public provided comments at this meeting. After discussing the OGW report, deliberating on specific issues about the draft guideline identified at the meeting, and hearing public comment, the BSC voted unanimously: to support the observations made by the OGW; that CDC adopt the guideline recommendations that, according to the workgroup’s report, had unanimous or majority support; and that CDC further consider the guideline recommendations for which the group had mixed opinions. CDC carefully considered the OGW observations, public comments, and BSC recommendations, and revised the guideline in response.

**Summary of the Clinical Evidence Review**

**Primary Clinical Questions**

CDC conducted a clinical systematic review of the scientific evidence to identify the effectiveness, benefits, and harms of long-term opioid therapy for chronic pain, consistent with the GRADE approach \(^{(47,48)}\). Long-term opioid therapy is defined as use of opioids on most days for >3 months. A previously published AHRQ-funded systematic review on the effectiveness and risks of long-term opioid therapy for chronic pain comprehensively addressed four clinical questions \(^{(14,52)}\). CDC, with the assistance of a methodology expert, searched the literature to identify newly published studies on these four original questions. Because long-term opioid use might be affected by use of opioids for acute pain, CDC subsequently developed a fifth clinical question (last in the series below), and in collaboration with a methodologist conducted a systematic review of the scientific evidence to address it. In brief, five clinical questions were addressed:

- The effectiveness of long-term opioid therapy versus placebo, no opioid therapy, or nonopioid therapy for long term (>1 year) outcomes related to pain, function, and quality of life, and how effectiveness varies according to the type/cause of pain, patient demographics, and patient comorbidities (Key Question [KQ] 1).
- The risks of opioids versus placebo or no opioids on abuse, addiction, overdose, and other harms, and how harms vary according to the type/cause of pain, patient demographics, patient comorbidities, and dose (KQ2).
- The comparative effectiveness of opioid dosing strategies (different methods for initiating and titrating opioids; immediate-release versus ER/LA opioids; different ER/LA opioids; immediate-release plus ER/LA opioids versus ER/LA opioids alone; scheduled, continuous versus as-needed dosing; dose escalation versus dose maintenance; opioid rotation versus maintenance; different strategies for treating acute exacerbations of chronic pain; decreasing opioid doses or tapering off versus continuation; and different tapering protocols and strategies) (KQ3).
- The accuracy of instruments for predicting risk for opioid overdose, addiction, abuse, or misuse; the effectiveness of risk mitigation strategies (use of risk prediction instruments); effectiveness of risk mitigation strategies including opioid management plans, patient education, urine drug testing, prescription drug monitoring program (PDMP) data, monitoring instruments, monitoring intervals, pill counts, and abuse-deterrent formulations for reducing risk for opioid overdose, addiction, abuse, or misuse; and the comparative effectiveness of treatment strategies for managing patients with addiction (KQ4).
- The effects of prescribing opioid therapy versus not prescribing opioid therapy for acute pain on long-term use (KQ5).

The review was focused on the effectiveness of long-term opioid therapy on long-term (>1 year) outcomes related to pain, function, and quality of life to ensure that findings are relevant to patients with chronic pain and long-term opioid prescribing. The effectiveness of short-term opioid therapy has already been established \(^{(10)}\). However, opioids have unique effects such as tolerance and physical dependence that might influence assessments of benefit over time. These effects raise questions about whether findings on short-term effectiveness of opioid therapy can be extrapolated to estimate benefits of long-term therapy for chronic pain. Thus, it is important to consider studies that provide data on long-term benefit. For certain opioid-related harms (overdose, fractures, falls, motor vehicle crashes), observational studies were included with outcomes measured at shorter intervals because such outcomes can occur early during opioid therapy, and such harms are not captured well in short-term clinical trials. A detailed listing of the key questions is provided in the Clinical Evidence Review (http://stacks.cdc.gov/view/cdc/38026).
Clinical Evidence Systematic Review Methods

Complete methods and data for the 2014 AHRQ report, upon which this updated systematic review is based, have been published previously (14,52). Study authors developed the protocol using a standardized process (53) with input from experts and the public and registered the protocol in the PROSPERO database (54). For the 2014 AHRQ report, a research librarian searched MEDLINE, the Cochrane Central Register of Controlled Trials, the Cochrane Database of Systematic Reviews, PsycINFO, and CINAHL for English-language articles published January 2008 through August 2014, using search terms for opioid therapy, specific opioids, chronic pain, and comparative study designs. Also included were relevant studies from an earlier review (10) in which searches were conducted without a date restriction, reference lists were reviewed, and ClinicalTrials.gov was searched. CDC updated the AHRQ literature search using the same search strategies as in the original review including studies published before April, 2015. Seven additional studies met inclusion criteria and were added to the review. CDC used the GRADE approach outlined in the ACIP Handbook for Developing Evidence-Based Recommendations (47) to rate the quality of evidence for the full body of evidence (evidence from the 2014 AHRQ review plus the update) for each clinical question. Evidence was categorized into the following types: type 1 (randomized clinical trials or overwhelming evidence from observational studies), type 2 (randomized clinical trials with important limitations, or exceptionally strong evidence from observational studies), type 3 (observational studies, or randomized clinical trials with notable limitations), or type 4 (clinical experience and observations, observational studies with important limitations, or randomized clinical trials with several major limitations). When no studies were present, evidence was considered to be insufficient. Per GRADE methods, type of evidence was categorized by study design as well as a function of limitations in study design or implementation, imprecision of estimates, variability in findings, indirectness of evidence, publication bias, magnitude of treatment effects, dose-response gradient, and constellation of plausible biases that could change effects. Results were synthesized qualitatively, highlighting new evidence identified during the update process. Meta-analysis was not attempted due to the small numbers of studies, variability in study designs and clinical heterogeneity, and methodological shortcomings of the studies. More detailed information about data sources and searches, study selection, data extraction and quality assessment, data synthesis, and update search yield and new evidence for the current review is provided in the Clinical Evidence Review (http://stacks.cdc.gov/view/cdc/38026).

Summary of Findings for Clinical Questions

The main findings of this updated review are consistent with the findings of the 2014 AHRQ report (14). In summary, evidence on long-term opioid therapy for chronic pain outside of end-of-life care remains limited, with insufficient evidence to determine long-term benefits versus no opioid therapy, though evidence suggests risk for serious harms that appears to be dose-dependent. These findings supplement findings from a previous review of the effectiveness of opioids for adults with chronic noncancer pain. In this previous review, based on randomized trials predominantly ≤12 weeks in duration, opioids were found to be moderately effective for pain relief, with small benefits for functional outcomes; although estimates vary, based on uncontrolled studies, a high percentage of patients discontinued long-term opioid use because of lack of efficacy and because of adverse events (10).

The GRADE evidence summary with type of evidence ratings for the five clinical questions for the current evidence review are outlined (Table 1). This summary is based on studies included in the AHRQ 2014 review (35 studies) plus additional studies identified in the updated search (seven studies). Additional details on findings from the original review are provided in the full 2014 AHRQ report (14,52). Full details on the clinical evidence review findings supporting this guideline are provided in the Clinical Evidence Review (http://stacks.cdc.gov/view/cdc/38026).

Effectiveness

For KQ1, no study of opioid therapy versus placebo, no opioid therapy, or nonopioid therapy for chronic pain evaluated long-term (≥1 year) outcomes related to pain, function, or quality of life. Most placebo-controlled randomized clinical trials were ≤6 weeks in duration. Thus, the body of evidence for KQ1 is rated as insufficient (0 studies contributing) (14).

Harms

For KQ2, the body of evidence is rated as type 3 (12 studies contributing; 11 from the original review plus one new study). One fair-quality cohort study found that long-term opioid therapy is associated with increased risk for an opioid abuse or dependence diagnosis (as defined by ICD-9-CM codes) versus no opioid prescription (22). Rates of opioid abuse or dependence diagnosis ranged from 0.7% with lower-dose (≤36 MME) chronic therapy to 6.1% with higher-dose (≥120 MME) chronic therapy, versus 0.004% with no opioids prescribed. Ten fair-quality uncontrolled studies reported estimates of opioid abuse, addiction, and related outcomes (55–65). In primary care settings, prevalence of opioid dependence...
Opioid Dosing Strategies

For KQ3, the body of evidence is rated as type 4 (14 studies contributing; 12 from the original review plus two new studies). For initiation and titration of opioids, the 2014 AHRQ report found insufficient evidence from three fair-quality, open-label trials to determine comparative effectiveness of ER/LA versus immediate-release opioids for titrating patients to stable pain control (75, 76). One new fair-quality cohort study of Veterans Affairs patients found initiation of therapy with an ER/LA opioid associated with greater risk for nonfatal overdose than initiation with an immediate-release opioid, with risk greatest in the first 2 weeks after initiation of treatment (77).

For comparative effectiveness and harms of ER/LA opioids, the 2014 AHRQ report included three randomized, head-to-head trials of various ER/LA opioids that found no clear differences in 1-year outcomes related to pain or function (78–80) but had methodological shortcomings. A fair-quality retrospective cohort study based on national Veterans Health Administration system pharmacy data found that methadone was associated with lower overall risk for all-cause mortality versus morphine (81), and a fair-quality retrospective cohort study based on Oregon Medicaid data found no statistically significant differences between methadone and long-acting morphine in risk for death or overdose symptoms (82). However, a new observational study (83) found methadone associated with increased risk for overdose versus sustained-release morphine among Tennessee Medicaid patients. The observed inconsistency in study findings suggests that risks of methadone might vary in different settings as a function of different monitoring and management protocols, though more research is needed to understand factors associated with safer methadone prescribing.

For dose escalation, the 2014 AHRQ report included one fair-quality randomized trial that found no differences between more liberal dose escalation and maintenance of current doses after 12 months in pain, function, all-cause withdrawals, or withdrawals due to opioid misuse (84). However, the difference in opioid dosages prescribed at the end of the trial was relatively small (mean 52 MME/day with more liberal dosing versus 40 MME/day). Evidence on other comparisons related to opioid dosing strategies (ER/LA versus immediate-release opioids; immediate-release plus ER/LA opioids versus ER/LA opioids alone; scheduled continuous dosing versus as-needed dosing; or opioid rotation versus maintenance of current therapy; long-term effects of strategies for treating acute exacerbations of chronic pain) was not available or too limited to determine effects on long-term clinical outcomes. For example, evidence on the comparative effectiveness of opioid tapering or discontinuation versus maintenance, and of different opioid tapering strategies, was limited to small, poor-quality studies (85–87).

Risk Assessment and Mitigation

For KQ4, the body of evidence is rated as type 3 for the accuracy of risk assessment tools and insufficient for the effectiveness of use of risk assessment tools and mitigation strategies in reducing harms (six studies contributing; four from the original review plus two new studies). The 2014 AHRQ report included four studies (88–91) on the accuracy of risk assessment instruments, administered prior to opioid therapy initiation, for predicting opioid abuse or misuse. Results for the Opioid Risk Tool (ORT) (89–91) were extremely inconsistent; evidence for other risk assessment instruments was very sparse, and studies had serious methodological shortcomings. One additional fair-quality (92) and one poor-quality (93) study identified for this update compared the predictive accuracy of the ORT, the Screener and Opioid Assessment for Patients with Pain–Revised (SOAPP-R), and the Brief Risk Interview.
For the ORT, sensitivity was 0.58 and 0.75 and specificity 0.54 and 0.86; for the SOAPP-R, sensitivity was 0.53 and 0.25 and specificity 0.62 and 0.73; and for the Brief Risk Interview, sensitivity was 0.73 and 0.83 and specificity 0.43 and 0.88. For the ORT, positive likelihood ratios ranged from noninformative (positive likelihood ratio close to 1) to moderately useful (positive likelihood ratio >5). The SOAPP-R was associated with noninformative likelihood ratios (estimates close to 1) in both studies.

No study evaluated the effectiveness of risk mitigation strategies (use of risk assessment instruments, opioid management plans, patient education, urine drug testing, use of PDMP data, use of monitoring instruments, more frequent monitoring intervals, pill counts, or use of abuse-deterrent formulations) for improving outcomes related to overdose, addiction, abuse, or misuse.

Effects of Opioid Therapy for Acute Pain on Long-Term Use

For KQ5, the body of evidence is rated as type 3 (two new studies contributing). Two fair-quality retrospective cohort studies found opioid therapy prescribed for acute pain associated with greater likelihood of long-term use. One study evaluated opioid-naïve patients who had undergone low-risk surgery, such as cataract surgery and varicose vein stripping (94). Use of opioids within 7 days of surgery was associated with increased risk for use at 1 year. The other study found that among patients with a workers’ compensation claim for acute low back pain, compared to patients who did not receive opioids early after injury (defined as use within 15 days following onset of pain), patients who did receive early opioids had an increased likelihood of receiving five or more opioid prescriptions 30–730 days following onset that increased with greater early exposure. Versus no early opioid use, the adjusted OR was 2.08 (95% CI = 1.55–2.78) for 1–140 MME/day and increased to 6.14 (95% confidence interval [CI] = 4.92–7.66) for ≥450 MME/day (95).

Summary of the Contextual Evidence Review

Primary Areas of Focus

Contextual evidence is complementary information that assists in translating the clinical research findings into recommendations. CDC conducted contextual evidence reviews on four topics to supplement the clinical evidence review findings:

- Effectiveness of nonpharmacologic (e.g., cognitive behavioral therapy [CBT], exercise therapy, interventional treatments, and multimodal pain treatment) and nonopioid pharmacologic treatments (e.g., acetaminophen, nonsteroidal anti-inflammatory drugs [NSAIDs], antidepressants, and anticonvulsants), including studies of any duration.
- Benefits and harms of opioid therapy (including additional studies not included in the clinical evidence review, such as studies that were not restricted to patients with chronic pain, evaluated outcomes at any duration, performed ecological analyses, or used observational study designs other than cohort and case-cohort control studies) related to specific opioids, high-dose therapy, co-prescription with other controlled substances, duration of use, special populations, and potential usefulness of risk stratification/mitigation approaches, in addition to effectiveness of treatments associated with addressing potential harms of opioid therapy (opioid use disorder).
- Clinician and patient values and preferences related to opioids and medication risks, benefits, and use.
- Resource allocation including costs and economic efficiency of opioid prescribing and could inform or complement the CDC recommendations under development (e.g., guidelines on nonpharmacologic and nonopioid pharmacologic treatments and guidelines with recommendations related to specific clinician actions such as urine drug testing or opioid tapering protocols).

Contextual Evidence Review Methods

CDC conducted a contextual evidence review to assist in developing the recommendations by providing an assessment of the balance of benefits and harms, values and preferences, and cost, consistent with the GRADE approach. Given the public health urgency for developing opioid prescribing recommendations, a rapid review was required for the contextual evidence review for the current guideline. Rapid reviews are used when there is a need to streamline the systematic review process to obtain evidence quickly (96). Methods used to streamline the process include limiting searches by databases, years, and languages considered, and truncating quality assessment and data abstraction protocols. CDC conducted “rapid reviews” of the contextual evidence on nonpharmacologic and nonopioid pharmacologic treatments, benefits and harms, values and preferences, and resource allocation.

Detailed information about contextual evidence data sources and searches, inclusion criteria, study selection, and
Recommendations and Reports

Summary of Findings for Contextual Areas

Full narrative reviews and tables that summarize key findings from the contextual evidence review are provided in the Contextual Evidence Review (http://stacks.cdc.gov/view/cdc/38027).

Effectiveness of Nonpharmacologic and Nonopioid Pharmacologic Treatments

Several nonpharmacologic and nonopioid pharmacologic treatments have been shown to be effective in managing chronic pain in studies ranging in duration from 2 weeks to 6 months. For example, CBT that trains patients in behavioral techniques and helps patients modify situational factors and cognitive processes that exacerbate pain has small positive effects on disability and catastrophic thinking (97). Exercise therapy can help reduce pain and improve function in chronic low back pain (98), improve function and reduce pain in osteoarthritis of the knee (99) and hip (100), and improve well-being, fibromyalgia symptoms, and physical function in fibromyalgia (101). Multimodal and multidisciplinary therapies (e.g., therapies that combine exercise and related therapies with psychologically based approaches) can help reduce pain and improve function more effectively than single modalities (102,103). Nonopioid pharmacologic approaches used for pain include analgesics such as acetaminophen, NSAIDs, and cyclooxygenase 2 (COX-2) inhibitors; selected anticonvulsants; and selected antidepressants (particularly tricyclics and serotonin and norepinephrine reuptake inhibitors [SNRIs]). Multiple guidelines recommend acetaminophen as first-line pharmacotherapy for osteoarthritis (104–109) or for low back pain (110) but note that it should be avoided in liver failure and that dosage should be reduced in patients with hepatic insufficiency or a history of alcohol abuse (109). Although guidelines also recommend NSAIDs as first-line treatment for osteoarthritis or low back pain (106,110), NSAIDs and COX-2 inhibitors do have risks, including gastrointestinal bleeding or perforation as well as renal and cardiovascular risks (111). FDA has recently strengthened existing label warnings that NSAIDs increase risks for heart attack and stroke, including that these risks might increase with longer use or at higher doses (112). Several guidelines agree that first- and second-line drugs for neuropathic pain include anticonvulsants (gabapentin or pregabalin), tricyclic antidepressants, and SNRIs (113–116). Interventional approaches such as epidural injection for certain conditions (e.g., lumbar radiculopathy) can provide short-term improvement in pain (117–119). Epidural injection has been associated with rare but serious adverse events, including loss of vision, stroke, paralysis, and death (120).

Benefits and Harms of Opioid Therapy

Balance between benefits and harms is a critical factor influencing the strength of clinical recommendations. In particular, CDC considered what is known from the epidemiology research about benefits and harms related to specific opioids and formulations, high dose therapy, co-prescription with other controlled substances, duration of use, special populations, and risk stratification and mitigation approaches. Additional information on benefits and harms of long-term opioid therapy from studies meeting rigorous selection criteria is provided in the clinical evidence review (e.g., see KQ2). CDC also considered the number of persons experiencing chronic pain, numbers potentially benefiting...
from opioids, and numbers affected by opioid-related harms. A review of these data is presented in the background section of this document, with detailed information provided in the Contextual Evidence Review (http://stacks.cdc.gov/view/cdc/38027). Finally, CDC considered the effectiveness of treatments that addressed potential harms of opioid therapy (opioid use disorder).

Regarding specific opioids and formulations, as noted by FDA, there are serious risks of ER/LA opioids, and the indication for this class of medications is for management of pain severe enough to require daily, around-the-clock, long-term opioid treatment in patients for whom other treatment options (e.g., nonopioid analgesics or immediate-release opioids) are ineffective, not tolerated, or would be otherwise inadequate to provide sufficient management of pain (127). Time-scheduled opioid use was associated with substantially higher average daily opioid dosage than as-needed opioid use in one study (122). Methadone has been associated with disproportionate numbers of overdose deaths relative to the frequency with which it is prescribed for pain. Methadone has been found to account for as much as a third of opioid use in treatment programs in the United States; further, methadone was involved in twice as many single-drug deaths as any other prescription opioid (123).

Regarding high-dose therapy, several epidemiologic studies that were excluded from the clinical evidence review because patient samples were not restricted to patients with chronic pain also examined the association between opioid dosage and overdose risk (23,24,124–126). Consistent with the clinical evidence review, the contextual review found that opioid-related overdose risk is dose-dependent, with higher opioid dosages associated with increased overdose risk. Two of these studies (23,24), as well as the two studies in the clinical evidence review (66,67), evaluated similar MME/day dose ranges for association with overdose risk. In these four studies, compared with opioids prescribed at <20 MME/day, the odds of overdose among patients prescribed opioids for chronic nonmalignant pain were between 1.3 (67) and 1.9 (24) for dosages of 20 to <50 MME/day, between 1.9 (67) and 4.6 (24) for dosages of 50 to <100 MME/day, and between 2.0 (67) and 8.9 (66) for dosages of ≥100 MME/day. Compared with dosages of 1–<20 MME/day, absolute risk difference approximation for 50–<100 MME/day was 0.15% for fatal overdose (24) and 1.40% for any overdose (66), and for ≥100 MME/day was 0.25% for fatal overdose (24) and 4.04% for any overdose (66). A recent study of Veterans Health Administration patients with chronic pain found that patients who died of overdoses related to opioids were prescribed higher opioid dosages (mean: 98 MME/day; median: 60 MME/day) than controls (mean: 48 MME/day; median: 25 MME/day) (127). Finally, another recent study of overdose deaths among state residents with and without opioid prescriptions revealed that prescription opioid-related overdose mortality rates rose rapidly up to prescribed doses of 200 MME/day, after which the mortality rates continued to increase but grew more gradually (128). A listing of common opioid medications and their MME equivalents is provided (Table 2).

Regarding coprescription of opioids with benzodiazepines, epidemiologic studies suggest that concurrent use of benzodiazepines and opioids might put patients at greater risk for potentially fatal overdose. Three studies of fatal overdose deaths found evidence of concurrent benzodiazepine use in 31%–61% of decedents (67,128,129). In one of these studies (67), among decedents who received an opioid prescription, those whose deaths were related to opioids were more likely to have obtained opioids from multiple physicians and pharmacies than decedents whose deaths were not related to opioids. Regarding duration of use, patients can experience tolerance and loss of effectiveness of opioids over time (130). Patients who do not experience clinically meaningful pain relief early in treatment (i.e., within 1 month) are unlikely to experience pain relief with longer-term use (131).

Regarding populations potentially at greater risk for harm, risk is greater for patients with sleep apnea or other causes of sleep-disordered breathing, patients with renal or hepatic insufficiency, older adults, pregnant women, patients with depression or other mental health conditions, and patients with alcohol or other substance use disorders. Interpretation of clinical data on the effects of opioids on sleep-disordered breathing is difficult because of the types of study designs and methods employed, and there is no clear consensus regarding association with risk for developing obstructive sleep apnea syndrome (132). However, opioid therapy can decrease respiratory drive, a high percentage of patients on long-term opioid therapy have been reported to have an abnormal apnea-hypopnea index (133), opioid therapy can worsen central sleep apnea in obstructive sleep apnea patients, and it can cause further desaturation in obstructive sleep apnea patients not on continuous positive airway pressure (CPAP) (31). Reduced renal or hepatic function can result in greater peak effect and longer duration of action and reduce the dose at which respiratory depression and overdose occurs (134). Age-related changes in patients aged ≥65 years, such as reduced renal function and medication clearance, even in the absence of renal disease (135), result in a smaller therapeutic window between safe dosages and dosages associated with respiratory depression and overdose. Older adults might also be at increased risk for falls and fractures related to opioids (136–138). Opioids used...
in pregnancy can be associated with additional risks to both mother and fetus. Some studies have shown an association of opioid use in pregnancy with birth defects, including neural tube defects (139,140), congenital heart defects (140), and gastroschisis (140); preterm delivery (141), poor fetal growth (141), and stillbirth (141). Importantly, in some cases, opioid use during pregnancy leads to neonatal opioid withdrawal syndrome (142). Patients with mental health comorbidities and patients with histories of substance use disorders might be at higher risk than other patients for opioid use disorder (62,143,144). Recent analyses found that depressed patients were at higher risk for drug overdose than patients without depression, particularly at higher opioid dosages, although investigators were unable to distinguish unintentional overdose from suicide attempts (145). In case-control and case-cohort studies, substance abuse/dependence was more prevalent among patients experiencing overdose than among patients not experiencing overdose (12% versus 6% [66], 40% versus 10% [24], and 26% versus 9% [23]).

Regarding risk stratification approaches, limited evidence was found regarding benefits and harms. Potential benefits of PDMPs and urine drug testing include the ability to identify patients who might be at higher risk for opioid overdose or opioid use disorder, and help determine which patients will benefit from greater caution and increased monitoring or interventions when risk factors are present. For example, one study found that most fatal overdoses could be identified retrospectively on the basis of two pieces of information, multiple prescribers and high total daily opioid dosage, both important risk factors for overdose (124,146) that are available to prescribers in the PDMP (124). However, limited evaluation of PDMPs at the state level has revealed mixed effects on changes in prescribing and mortality outcomes (28). Potential harms of risk stratification include underestimation of risks of opioid therapy when screening tools are not adequately sensitive, as well as potential overestimation of risk, which could lead to inappropriate clinical decisions.

Regarding risk mitigation approaches, limited evidence was found regarding benefits and harms. Although no studies were found to examine prescribing of naloxone with opioid pain medication in primary care settings, naloxone distribution through community-based programs providing prevention services for substance users has been demonstrated to be associated with decreased risk for opioid overdose death at the community level (147).

Concerns have been raised that prescribing changes such as dose reduction might be associated with unintended negative consequences, such as patients seeking heroin or other illicitly obtained opioids (148) or interference with appropriate pain treatment (149). With the exception of a study noting an association between an abuse-deterrent formulation of OxyContin and heroin use, showing that some patients in qualitative interviews reported switching to another opioid, including heroin, for many reasons, including cost and availability as well as ease of use (150), CDC did not identify studies evaluating these potential outcomes.

Finally, regarding the effectiveness of opioid use disorder treatments, methadone and buprenorphine for opioid use disorder have been found to increase retention in treatment and to decrease illicit opioid use among patients with opioid use disorder involving heroin (151–153). Although findings are mixed, some studies suggest that effectiveness is enhanced when psychosocial treatments (e.g., contingency management, community reinforcement, psychotherapeutic counseling, and family therapy) are used in conjunction with medication-assisted therapy; for example, by reducing opioid misuse and increasing retention during maintenance therapy, and improving compliance after detoxification (154,155).

**Clinician and Patient Values and Preferences**

Clinician and patient values and preferences can inform how benefits and harms of long-term opioid therapy are weighted and estimate the effort and resources required to effectively provide implementation support. Many physicians lack confidence in their ability to prescribe opioids safely (156), to predict (157) or detect (158) prescription drug abuse, and to discuss abuse with their patients (158). Although clinicians have reported favorable beliefs and attitudes about improvements in pain and quality of life attributed to opioids (159), most consider prescription drug abuse to be a “moderate” or “big” problem in their community, and large proportions are “very” concerned about opioid addiction (55%) and death (48%) (160). Clinicians do not consistently use practices intended to decrease the risk for misuse, such as PDMPs (161,162), urine drug testing (163), and opioid treatment agreements (164). This is likely due in part to challenges related to registering for PDMP access and logging into the PDMP (which can interrupt normal clinical workflow if data are not integrated into electronic health record systems) (165), competing clinical demands, perceived inadequate time to discuss the rationale for urine drug testing and to order confirmatory testing, and feeling unprepared to interpret and address results (166).

Many patients do not have an opinion about “opioids” or know what this term means (167). Most are familiar with the term “narcotics.” About a third associated “narcotics” with addiction or abuse, and about half feared “addiction” from long-term “narcotic” use (168). Most patients taking opioids experience side effects (73% of patients taking hydrocodone for noncancer pain [11], 96% of patients taking opioids for chronic pain [12]), and side effects, rather than pain relief,
have been found to explain most of the variation in patients’ preferences related to taking opioids (12). For example, patients taking hydrocodone for noncancer pain commonly reported side effects including dizziness, headache, fatigue, drowsiness, nausea, vomiting, and constipation (11). Patients with chronic pain in focus groups emphasized effectiveness (168). Patients taking high dosages report reliance on opioids despite ambivalence about their benefits (169) and regardless of pain reduction, reported problems, concerns, side effects, or perceived helpfulness (13).

**Resource Allocation**

Resource allocation (cost) is an important consideration in understanding the feasibility of clinical recommendations. CDC searched for evidence on opioid therapy compared with other treatments; costs of misuse, abuse, and overdose from prescription opioids; and costs of specific risk mitigation strategies (e.g., urine drug testing). Yearly direct and indirect costs related to prescription opioids have been estimated (based on studies published since 2010) to be $55.4 billion for nonmedical use of prescription opioids (170); $55.7 billion for abuse, dependence (i.e., opioid use disorder), and misuse of prescription opioids (171); and $20.4 billion for direct and indirect costs related to opioid-related overdose alone (172). In 2012, total expenses for outpatient prescription opioids were estimated at $9.0 billion, an increase of 120% from 2002 (173). Although there are perceptions that opioid therapy for chronic pain is less expensive than more time-intensive nonpharmacologic management approaches, many pain treatments, including acetaminophen, NSAIDs, tricyclic antidepressants, and massage therapy, are associated with lower mean and median annual costs compared with opioid therapy (174). COX-2 inhibitors, SNRIs, anticonvulsants, topical analgesics, physical therapy, and CBT are also associated with lower median annual costs compared with opioid therapy (174). Limited information was found on costs of strategies to decrease risks associated with opioid therapy; however, urine drug testing, including screening and confirmatory tests, has been estimated to cost $211–$363 per test (175).

**Recommendations**

The recommendations are grouped into three areas for consideration:
- Determining when to initiate or continue opioids for chronic pain.
- Opioid selection, dosage, duration, follow-up, and discontinuation.
- Assessing risk and addressing harms of opioid use.

There are 12 recommendations (Box 1). Each recommendation is followed by a rationale for the recommendation, with considerations for implementation noted. In accordance with the ACIP GRADE process, CDC based the recommendations on consideration of the clinical evidence, contextual evidence (including benefits and harms, values and preferences, resource allocation), and expert opinion. For each recommendation statement, CDC notes the recommendation category (A or B) and the type of the evidence (1, 2, 3, or 4) supporting the statement (Box 2). Expert opinion is reflected within each of the recommendation rationales. While there was not an attempt to reach consensus among experts, experts from the Core Expert Group and from the Opioid Guideline Workgroup (”experts”) expressed overall, general support for all recommendations. Where differences in expert opinion emerged for detailed actions within the clinical recommendations or for implementation considerations, CDC notes the differences of opinion in the supporting rationale statements.

Category A recommendations indicate that most patients should receive the recommended course of action; category B recommendations indicate that different choices will be appropriate for different patients, requiring clinicians to help patients arrive at a decision consistent with patient values and preferences and specific clinical situations. Consistent with the ACIP (47) and GRADE process (48), category A recommendations were made, even with type 3 and 4 evidence, when there was broad agreement that the advantages of a clinical action greatly outweighed the disadvantages based on a consideration of benefits and harms, values and preferences, and resource allocation. Category B recommendations were made when there was broad agreement that the advantages and disadvantages of a clinical action were more balanced, but advantages were significant enough to warrant a recommendation. All recommendations are category A recommendations, with the exception of recommendation 10, which is rated as category B. Recommendations were associated with a range of evidence types, from type 2 to type 4.

In summary, the categorization of recommendations was based on the following assessment:
- No evidence shows a long-term benefit of opioids in pain and function versus no opioids for chronic pain with outcomes examined at least 1 year later (with most placebo-controlled randomized trials ≤6 weeks in duration).
- Extensive evidence shows the possible harms of opioids (including opioid use disorder, overdose, and motor vehicle injury).
- Extensive evidence suggests some benefits of nonpharmacologic and nonopioid pharmacologic treatments compared with long-term opioid therapy, with less harm.
**BOX 1. CDC recommendations for prescribing opioids for chronic pain outside of active cancer, palliative, and end-of-life care**

**Determining When to Initiate or Continue Opioids for Chronic Pain**

1. Nonpharmacologic therapy and nonopioid pharmacologic therapy are preferred for chronic pain. Clinicians should consider opioid therapy only if expected benefits for both pain and function are anticipated to outweigh risks to the patient. If opioids are used, they should be combined with nonpharmacologic therapy and nonopioid pharmacologic therapy, as appropriate.

2. Before starting opioid therapy for chronic pain, clinicians should establish treatment goals with all patients, including realistic goals for pain and function, and should consider how therapy will be discontinued if benefits do not outweigh risks. Clinicians should continue opioid therapy only if there is clinically meaningful improvement in pain and function that outweighs risks to patient safety.

3. Before starting and periodically during opioid therapy, clinicians should discuss with patients known risks and realistic benefits of opioid therapy and patient and clinician responsibilities for managing therapy.

**Opioid Selection, Dosage, Duration, Follow-Up, and Discontinuation**

4. When starting opioid therapy for chronic pain, clinicians should prescribe immediate-release opioids instead of extended-release/long-acting (ER/LA) opioids.

5. When opioids are started, clinicians should prescribe the lowest effective dosage. Clinicians should use caution when prescribing opioids at any dosage, should carefully reassess evidence of individual benefits and risks when increasing dosage to ≥50 morphine milligram equivalents (MME)/day, and should avoid increasing dosage to ≥90 MME/day or carefully justify a decision to titrate dosage to ≥90 MME/day.

6. Long-term opioid use often begins with treatment of acute pain. When opioids are used for acute pain, clinicians should prescribe the lowest effective dose of immediate-release opioids and should prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids. Three days or less will often be sufficient; more than seven days will rarely be needed.

7. Clinicians should evaluate benefits and harms with patients within 1 to 4 weeks of starting opioid therapy for chronic pain or of dose escalation. Clinicians should evaluate benefits and harms of continued therapy with patients every 3 months or more frequently. If benefits do not outweigh harms of continued opioid therapy, clinicians should optimize other therapies and work with patients to taper opioids to lower dosages or to taper and discontinue opioids.

**Assessing Risk and Addressing Harms of Opioid Use**

8. Before starting and periodically during continuation of opioid therapy, clinicians should evaluate risk factors for opioid-related harms. Clinicians should incorporate into the management plan strategies to mitigate risk, including considering offering naloxone when factors that increase risk for opioid overdose, such as history of overdose, history of substance use disorder, higher opioid dosages (≥50 MME/day), or concurrent benzodiazepine use, are present.

9. Clinicians should review the patient’s history of controlled substance prescriptions using state prescription drug monitoring program (PDMP) data to determine whether the patient is receiving opioid dosages or dangerous combinations that put him or her at high risk for overdose. Clinicians should review PDMP data when starting opioid therapy for chronic pain and periodically during opioid therapy for chronic pain, ranging from every prescription to every 3 months.

10. When prescribing opioids for chronic pain, clinicians should use urine drug testing before starting opioid therapy and consider urine drug testing at least annually to assess for prescribed medications as well as other controlled prescription drugs and illicit drugs.

11. Clinicians should avoid prescribing opioid pain medication and benzodiazepines concurrently whenever possible.

12. Clinicians should offer or arrange evidence-based treatment (usually medication-assisted treatment with buprenorphine or methadone in combination with behavioral therapies) for patients with opioid use disorder.

*All recommendations are category A (apply to all patients outside of active cancer treatment, palliative care, and end-of-life care) except recommendation 10 (designated category B, with individual decision making required); see full guideline for evidence ratings.*
**Recommendation Categories**

- **Category A recommendation**: Applies to all persons; most patients should receive the recommended course of action.
- **Category B recommendation**: Individual decision making needed; different choices will be appropriate for different patients. Clinicians help patients arrive at a decision consistent with patient values and preferences and specific clinical situations.

**Evidence Type**

- **Type 1 evidence**: Randomized clinical trials or overwhelming evidence from observational studies.
- **Type 2 evidence**: Randomized clinical trials with important limitations, or exceptionally strong evidence from observational studies.
- **Type 3 evidence**: Observational studies or randomized clinical trials with notable limitations.
- **Type 4 evidence**: Clinical experience and observations, observational studies with important limitations, or randomized clinical trials with several major limitations.

---

**Determining When to Initiate or Continue Opioids for Chronic Pain**

1. **Nonpharmacologic therapy and nonopioid pharmacologic therapy are preferred for chronic pain. Clinicians should consider opioid therapy only if expected benefits for both pain and function are anticipated to outweigh risks to the patient. If opioids are used, they should be combined with nonpharmacologic therapy and nonopioid pharmacologic therapy, as appropriate (recommendation category: A, evidence type: 3).**

Patients with pain should receive treatment that provides the greatest benefits relative to risks. The contextual evidence review found that many nonpharmacologic therapies, including physical therapy, weight loss for knee osteoarthritis, psychological therapies such as CBT, and certain interventional procedures can ameliorate chronic pain. There is high-quality evidence that exercise therapy (a prominent modality in physical therapy) for hip (100) or knee (99) osteoarthritis reduces pain and improves function immediately after treatment and that the improvements are sustained for at least 2–6 months. Previous guidelines have strongly recommended aerobic, aquatic, and/or resistance exercises for patients with osteoarthritis of the knee or hip (176). Exercise therapy also can help reduce pain and improve function in low back pain and can improve global well-being and physical function in fibromyalgia (98,101). Multimodal therapies and multidisciplinary biopsychosocial rehabilitation-combining approaches (e.g., psychological therapies with exercise) can reduce long-term pain and disability compared with usual care and compared with physical treatments (e.g., exercise) alone. Multimodal therapies are not always available or reimbursed by insurance and can be time-consuming and costly for patients. Intervventional approaches such as arthrocentesis and intraarticular glucocorticoid injection for pain associated with rheumatoid arthritis (117) or osteoarthritis (118) and subacromial corticosteroid injection for rotator cuff disease (119) can provide short-term improvement in pain and function. Evidence is insufficient to determine the extent to which repeated glucocorticoid injection increases potential risks such as articular cartilage changes (in osteoarthritis) and sepsis (118). Serious adverse events are rare but have been reported with epidural injection (120).

Several nonopioid pharmacologic therapies (including acetaminophen, NSAIDs, and selected antidepressants and anticonvulsants) are effective for chronic pain. In particular, acetaminophen and NSAIDs can be useful for arthritis and low back pain. Selected anticonvulsants such as pregabalin and gabapentin can improve pain in diabetic neuropathy and post-herpetic neuralgia (contextual evidence review). Pregabalin, gabapentin, and carbamazepine are FDA-approved for treatment of certain neuropathic pain conditions, and pregabalin is FDA approved for fibromyalgia management. In patients with or without depression, tricyclic antidepressants and SNRIs provide effective analgesia for neuropathic pain conditions including diabetic neuropathy and post-herpetic neuralgia, often at lower dosages and with a shorter time to onset of effect than for treatment of depression (see contextual evidence review). Tricyclics and SNRIs can also relieve fibromyalgia symptoms. The SNRI duloxetine is FDA-approved for the treatment of diabetic neuropathy and fibromyalgia. Because patients with chronic pain often suffer from concurrent depression (144), and depression can exacerbate physical symptoms including pain (177), patients with co-occurring pain and depression are especially likely to benefit from antidepressant medication (see Recommendation 8). Nonopioid pharmacologic therapies
are not generally associated with substance use disorder, and the numbers of fatal overdoses associated with nonopioid medications are a fraction of those associated with opioid medications (contextual evidence review). For example, acetaminophen, NSAIDs, and opioid pain medication were involved in 881, 228, and 16,651 pharmaceutical overdose deaths in the United States in 2010 (178). However, nonopioid pharmacologic therapies are associated with certain risks, particularly in older patients, pregnant patients, and patients with certain co-morbidities such as cardiovascular, renal, gastrointestinal, and liver disease (see contextual evidence review). For example, acetaminophen can be hepatotoxic at dosages of >3–4 grams/day and at lower dosages in patients with chronic alcohol use or liver disease (see contextual evidence review). NSAID use has been associated with gastritis, peptic ulcer disease, cardiovascular events (111,112), and fluid retention, and most NSAIDs (choline magnesium trilisate and selective COX-2 inhibitors are exceptions) interfere with platelet aggregation (179). Clinicians should review FDA-approved labeling including boxed warnings before initiating treatment with any pharmacologic therapy.

Although opioids can reduce pain during short-term use, the clinical evidence review found insufficient evidence to determine whether pain relief is sustained and whether function or quality of life improves with long-term opioid therapy (KQ1). While benefits for pain relief, function, and quality of life with long-term opioid use for chronic pain are uncertain, risks associated with long-term opioid use are clearer and significant. Based on the clinical evidence review, long-term opioid use for chronic pain is associated with serious risks including increased risk for opioid use disorder, overdose, myocardial infarction, and motor vehicle injury (KQ2). At a population level, more than 165,000 persons in the United States have died from opioid pain-medication-related overdoses since 1999 (see Contextual Evidence Review).

Integrated pain management requires coordination of medical, psychological, and social aspects of health care and includes primary care, mental health care, and specialist services when needed (180). Nonpharmacologic physical and psychological treatments such as exercise and CBT are approaches that encourage active patient participation in the care plan, address the effects of pain in the patient’s life, and can result in sustained improvements in pain and function without apparent risks. Despite this, these therapies are not always or fully covered by insurance, and access and cost can be barriers for patients. For many patients, aspects of these approaches can be used even when there is limited access to specialty care. For example, previous guidelines have strongly recommended aerobic, aquatic, and/or resistance exercises for patients with osteoarthritis of the knee or hip (176) and maintenance of activity for patients with low back pain (110). A randomized trial found no difference in reduced chronic low back pain intensity, frequency or disability between patients assigned to relatively low-cost group aerobics and individual physiotherapy or muscle reconditioning sessions (181). Low-cost options to integrate exercise include brisk walking in public spaces or use of public recreation facilities for group exercise. CBT addresses psychosocial contributors to pain and improves function (97). Primary care clinicians can integrate elements of a cognitive behavioral approach into their practice by encouraging patients to take an active role in the care plan, by supporting patients in engaging in beneficial but potentially anxiety-provoking activities, such as exercise (179), or by providing education in relaxation techniques and coping strategies. In many locations, there are free or low-cost patient support, self-help, and educational community-based programs that can provide stress reduction and other mental health benefits. Patients with more entrenched anxiety or fear related to pain, or other significant psychological distress, can be referred for formal therapy with a mental health specialist (e.g., psychologist, psychiatrist, clinical social worker). Multimodal therapies should be considered for patients not responding to single-modality therapy, and combinations should be tailored depending on patient needs, cost, and convenience.

To guide patient-specific selection of therapy, clinicians should evaluate patients and establish or confirm the diagnosis. Detailed recommendations on diagnosis are provided in other guidelines (110,179), but evaluation should generally include a focused history, including history and characteristics of pain and potentially contributing factors (e.g., function, psychosocial stressors, sleep) and physical exam, with imaging or other diagnostic testing only if indicated (e.g., if severe or progressive neurologic deficits are present or if serious underlying conditions are suspected) (110,179). For complex pain syndromes, pain specialty consultation can be considered to assist with diagnosis as well as management. Diagnosis can help identify disease-specific interventions to reverse or ameliorate pain; for example, improving glucose control to prevent progression of diabetic neuropathy; immune-modulating agents for rheumatoid arthritis; physical or occupational therapy to address posture, muscle weakness, or repetitive occupational motions that contribute to musculoskeletal pain; or surgical intervention to relieve mechanical/compressive pain (179). The underlying mechanism for most pain syndromes can be categorized as neuropathic (e.g., diabetic neuropathy, postherpetic neuralgia, fibromyalgia), or nociceptive (e.g., osteoarthritis, muscular back pain). The diagnosis and pathophysiologic mechanism of pain have implications for symptomatic pain treatment with medication. For example, evidence is limited or insufficient
for improved pain or function with long-term use of opioids for several chronic pain conditions for which opioids are commonly prescribed, such as low back pain (182), headache (183), and fibromyalgia (184). Although NSAIDs can be used for exacerbations of nociceptive pain, other medications (e.g., tricyclics, selected anticonvulsants, or transdermal lidocaine) generally are recommended for neuropathic pain. In addition, improvement of neuropathic pain can begin weeks or longer after symptomatic treatment is initiated (179). Medications should be used only after assessment and determination that expected benefits outweigh risks given patient-specific factors. For example, clinicians should consider falls risk when selecting and dosing potentially sedating medications such as tricyclics, anticonvulsants, or opioids, and should weigh risks and benefits of use, dose, and duration of NSAIDs when treating older adults as well as patients with hypertension, renal insufficiency, or heart failure, or those with risk for peptic ulcer disease or cardiovascular disease. Some guidelines recommend topical NSAIDs for localized osteoarthritis (e.g., knee osteoarthritis) over oral NSAIDs in patients aged ≥75 years to minimize systemic effects (176).

Experts agreed that opioids should not be considered first-line or routine therapy for chronic pain (i.e., pain continuing or expected to continue >3 months or past the time of normal tissue healing) outside of active cancer, palliative, and end-of-life care, given small to moderate short-term benefits, uncertain long-term benefits, and potential for serious harms; although evidence on long-term benefits of nonopioid therapies is also limited, these therapies are also associated with short-term benefits, and risks are much lower. This does not mean that patients should be required to sequentially “fail” nonpharmacologic and nonopioid pharmacologic therapy before proceeding to opioid therapy. Rather, expected benefits specific to the clinical context should be weighed against risks before initiating therapy. In some clinical contexts (e.g., headache or fibromyalgia), expected benefits of initiating opioids are unlikely to outweigh risks regardless of previous nonpharmacologic and nonopioid pharmacologic therapies used. In other situations (e.g., serious illness in a patient with poor prognosis for return to previous level of function, contraindications to other therapies, and clinician and patient agreement that the overriding goal is patient comfort), opioids might be appropriate regardless of previous therapies used. In addition, when opioid pain medication is used, it is more likely to be effective if integrated with nonpharmacologic therapy. Nonpharmacologic approaches such as exercise and CBT should be used to reduce pain and improve function in patients with chronic pain. Nonopioid pharmacologic therapy should be used when benefits outweigh risks and should be combined with nonpharmacologic therapy to reduce pain and improve function. If opioids are used, they should be combined with nonpharmacologic therapy and nonopioid pharmacologic therapy, as appropriate, to provide greater benefits to patients in improving pain and function.

2. Before starting opioid therapy for chronic pain, clinicians should establish treatment goals with all patients, including realistic goals for pain and function, and should consider how opioid therapy will be discontinued if benefits do not outweigh risks. Clinicians should continue opioid therapy only if there is clinically meaningful improvement in pain and function that outweighs risks to patient safety (recommendation category: A, evidence type: 4).

The clinical evidence review found insufficient evidence to determine long-term benefits of opioid therapy for chronic pain and found an increased risk for serious harms related to long-term opioid therapy that appears to be dose-dependent. In addition, studies on currently available risk assessment instruments were sparse and showed inconsistent results (KQ4). The clinical evidence review for the current guideline considered studies with outcomes examined at ≥1 year that compared opioid use versus nonuse or placebo. Studies of opioid therapy for chronic pain that did not have a nonopioid control group have found that although many patients discontinue opioid therapy for chronic noncancer pain due to adverse effects or insufficient pain relief, there is weak evidence that patients who are able to continue opioid therapy for at least 6 months can experience clinically significant pain relief and insufficient evidence that function or quality of life improves (185). These findings suggest that it is very difficult for clinicians to predict whether benefits of opioids for chronic pain will outweigh risks of ongoing treatment for individual patients. Opioid therapy should not be initiated without consideration of an “exit strategy” to be used if the therapy is unsuccessful.

Experts agreed that before opioid therapy is initiated for chronic pain outside of active cancer, palliative, and end-of-life care, clinicians should determine how effectiveness will be evaluated and should establish treatment goals with patients. Because the line between acute pain and initial chronic pain is not always clear, it might be difficult for clinicians to determine when they are initiating opioids for chronic pain rather than treating acute pain. Pain lasting longer than 3 months or past the time of normal tissue healing (which could be substantially shorter than 3 months, depending on the condition) is generally no longer considered acute. However, establishing treatment goals with a patient who has already received opioid therapy for 3 months would defer this discussion well past the point of
initiation of opioid therapy for chronic pain. Clinicians often write prescriptions for long-term use in 30-day increments, and opioid prescriptions written for ≥30 days are likely to represent initiation or continuation of long-term opioid therapy. Before writing an opioid prescription for ≥30 days, clinicians should establish treatment goals with patients. Clinicians seeing new patients already receiving opioids should establish treatment goals for continued opioid therapy. Although the clinical evidence review did not find studies evaluating the effectiveness of written agreements or treatment plans (KQ4), clinicians and patients who set a plan in advance will clarify expectations regarding how opioids will be prescribed and monitored, as well as situations in which opioids will be discontinued or doses tapered (e.g., if treatment goals are not met, opioids are no longer needed, or adverse events put the patient at risk) to improve patient safety.

Experts thought that goals should include improvement in both pain relief and function (and therefore in quality of life). However, there are some clinical circumstances under which reductions in pain without improvement in physical function might be a more realistic goal (e.g., diseases typically associated with progressive functional impairment or catastrophic injuries such as spinal cord trauma). Experts noted that function can include emotional and social as well as physical dimensions. In addition, experts emphasized that mood has important interactions with pain and function. Experts agreed that clinicians may use validated instruments such as the three-item “Pain average, interference with Enjoyment of life, and interference with General activity” (PEG) Assessment Scale (186) to track patient outcomes. Clinically meaningful improvement has been defined as a 30% improvement in scores for both pain and function (187). Monitoring progress toward patient-centered functional goals (e.g., walking the dog or walking around the block, returning to part-time work, attending family sports or recreational activities) can also contribute to the assessment of functional improvement. Clinicians should use these goals in assessing benefits of opioid therapy for individual patients and in weighing benefits against risks of continued opioid therapy (see Recommendation 7, including recommended intervals for follow-up). Because depression, anxiety, and other psychological co-morbidities often coexist with and can interfere with resolution of pain, clinicians should use validated instruments to assess for these conditions (see Recommendation 8) and ensure that treatment for these conditions is optimized. If patients receiving opioid therapy for chronic pain do not experience meaningful improvements in both pain and function compared with prior to initiation of opioid therapy, clinicians should consider working with patients to taper and discontinue opioids (see Recommendation 7) and should use nonpharmacologic and nonopioid pharmacologic approaches to pain management (see Recommendation 1).

3. **Before starting and periodically during opioid therapy, clinicians should discuss with patients known risks and realistic benefits of opioid therapy and patient and clinician responsibilities for managing therapy** (recommendation category: A, evidence type: 3).

The clinical evidence review did not find studies evaluating effectiveness of patient education or opioid treatment plans as risk-mitigation strategies (KQ4). However, the contextual evidence review found that many patients lack information about opioids and identified concerns that some clinicians miss opportunities to effectively communicate about safety. Given the substantial evidence gaps on opioids, uncertain benefits of long-term use, and potential for serious harms, patient education and discussion before starting opioid therapy are critical so that patient preferences and values can be understood and used to inform clinical decisions. Experts agreed that essential elements to communicate to patients before starting and periodically during opioid therapy include realistic expected benefits, common and serious harms, and expectations for clinician and patient responsibilities to mitigate risks of opioid therapy.

Clinicians should involve patients in decisions about whether to start or continue opioid therapy. Given potentially serious risks of long-term opioid therapy, clinicians should ensure that patients are aware of potential benefits of, harms of, and alternatives to opioids before starting or continuing opioid therapy. Clinicians are encouraged to have open and honest discussions with patients to inform mutual decisions about whether to start or continue opioid therapy. Important considerations include the following:

- **Be explicit and realistic about expected benefits of opioids**, explaining that while opioids can reduce pain during short-term use, there is no good evidence that opioids improve pain or function with long-term use, and that complete relief of pain is unlikely (clinical evidence review, KQ1).
- **Emphasize improvement in function as a primary goal and that function can improve even when pain is still present.**
- **Advise patients about serious adverse effects of opioids**, including potentially fatal respiratory depression and development of a potentially serious lifelong opioid use disorder that can cause distress and inability to fulfill major role obligations.
- **Advise patients about common effects of opioids**, such as constipation, dry mouth, nausea, vomiting, drowsiness, confusion, tolerance, physical dependence, and withdrawal symptoms when stopping opioids. To prevent constipation associated with opioid use, advise patients to increase...
Opioid Selection, Dosage, Duration, Follow-Up, and Discontinuation


ER/LA opioids include methadone, transdermal fentanyl, and extended-release versions of opioids such as oxycodone, oxymorphone, hydrocodone, and morphine. The clinical evidence review found a fair-quality study showing a higher risk for overdose among patients initiating treatment with ER/LA opioids than among those initiating treatment with immediate-release opioids (77). The clinical evidence review did not find evidence that continuous, time-scheduled use of ER/LA opioids is more effective or safer than intermittent use of immediate-release opioids or that time-scheduled use of ER/LA opioids reduces risks for opioid misuse or addiction (KQ3).

In 2014, the FDA modified the labeling for ER/LA opioid pain medications, noting serious risks and recommending that ER/LA opioids be reserved for “management of pain severe enough to require daily, around-the-clock, long-term opioid treatment” when “alternative treatment options (e.g., nonopioid analgesics or immediate-release opioids) are ineffective, not tolerated, or would be otherwise inadequate to provide sufficient management of pain” and not used as “as needed” pain relievers (121). FDA has also noted that some ER/LA opioids are only appropriate for opioid-tolerant patients, defined as patients who have received certain dosages of opioids (e.g., 60 mg daily of oral morphine, 30 mg daily of oral oxycodone, or equianalgesic dosages of other opioids) for at least 1 week (189). Time-scheduled opioid use can be associated with greater total average daily opioid dosage compared with intermittent, as-needed opioid use (contextual evidence review). In addition, experts indicated that there was not enough evidence to determine the safety of using immediate-release opioids for breakthrough pain when ER/LA opioids are used for chronic pain outside of active cancer pain, palliative care, or end-of-life care, and that this practice might be associated with dose escalation.

Abuse-deterrent technologies have been employed to prevent manipulation intended to defeat extended-release properties of ER/LA opioids and to prevent opioid use by unintended routes of administration, such as injection of oral opioids. As indicated in FDA guidance for industry on evaluation and labeling of abuse-deterrent opioids (190), although abuse-deterrent technologies are expected to make manipulation of opioids more difficult or less rewarding, they do not prevent

hydration and fiber intake and to maintain or increase physical activity. Stool softeners or laxatives might be needed.
• Discuss effects that opioids might have on ability to safely operate a vehicle, particularly when opioids are initiated, when dosages are increased, or when other central nervous system depressants, such as benzodiazepines or alcohol, are used concurrently.
• Discuss increased risks for opioid use disorder, respiratory depression, and death at higher dosages, along with the importance of taking only the amount of opioids prescribed, i.e., not taking more opioids or taking them more often.
• Review increased risks for respiratory depression when opioids are taken with benzodiazepines, other sedatives, alcohol, illicit drugs such as heroin, or other opioids.
• Discuss risks to household members and other individuals if opioids are intentionally or unintentionally shared with others for whom they are not prescribed, including the possibility that others might experience overdose at the same or at lower dosage than prescribed for the patient, and that young children are susceptible to unintentional ingestion. Discuss storage of opioids in a secure, preferably locked location and options for safe disposal of unused opioids (188).
• Discuss the importance of periodic reassessment to ensure that opioids are helping to meet patient goals and to allow opportunities for opioid discontinuation and consideration of additional nonpharmacologic or nonopioid pharmacologic treatment options if opioids are not effective or are harmful.
• Discuss planned use of precautions to reduce risks, including use of prescription drug monitoring program information (see Recommendation 9) and urine drug testing (see Recommendation 10). Consider including discussion of naloxone use for overdose reversal (see Recommendation 8).
• Consider whether cognitive limitations might interfere with management of opioid therapy (for older adults in particular) and, if so, determine whether a caregiver can responsibly co-manage medication therapy. Discuss the importance of reassessing safer medication use with both the patient and caregiver.

Given the possibility that benefits of opioid therapy might diminish or that risks might become more prominent over time, it is important that clinicians review expected benefits and risks of continued opioid therapy with patients periodically, at least every 3 months (see Recommendation 7).
opioid abuse through oral intake, the most common route of opioid abuse, and can still be abused by nonoral routes. The “abuse-deterrent” label does not indicate that there is no risk for abuse. No studies were found in the clinical evidence review assessing the effectiveness of abuse-deterrent technologies as a risk mitigation strategy for deterring or preventing abuse. In addition, abuse-deterrent technologies do not prevent unintentional overdose through oral intake. Experts agreed that recommendations could not be offered at this time related to use of abuse-deterrent formulations.

In comparing different ER/LA formulations, the clinical evidence review found inconsistent results for overdose risk with methadone versus other ER/LA opioids used for chronic pain (KQ3). The contextual evidence review found that methadone has been associated with disproportionate numbers of overdose deaths relative to the frequency with which it is prescribed for chronic pain. In addition, methadone is associated with cardiac arrhythmias along with QT prolongation on the electrocardiogram, and it has complicated pharmacokinetics and pharmacodynamics, including a long and variable half-life and peak respiratory depressant effect occurring later and lasting longer than peak analgesic effect. Experts noted that the pharmacodynamics of methadone are subject to more inter-individual variability than other opioids. In regard to other ER/LA opioid formulations, experts noted that the absorption and pharmacodynamics of transdermal fentanyl are complex, with gradually increasing serum concentration during the first part of the 72-hour dosing interval, as well as variable absorption based on factors such as external heat. In addition, the dosing of transdermal fentanyl in mcg/hour, which is not typical for a drug used by outpatients, can be confusing. Experts thought that these complexities might increase the risk for fatal overdose when methadone or transdermal fentanyl is prescribed to a patient who has not used it previously or by clinicians who are not familiar with its effects.

Experts agreed that for patients not already receiving opioids, clinicians should not initiate opioid treatment with ER/LA opioids and should not prescribe ER/LA opioids for intermittent use. ER/LA opioids should be reserved for severe, continuous pain and should be considered only for patients who have received immediate-release opioids daily for at least 1 week. When changing to an ER/LA opioid for a patient previously receiving a different immediate-release opioid, clinicians should consult product labeling and reduce total daily dosage to account for incomplete opioid cross-tolerance. Clinicians should use additional caution with ER/LA opioids and consider a longer dosing interval when prescribing to patients with renal or hepatic dysfunction because decreased clearance of drugs among these patients can lead to accumulation of drugs to toxic levels and persistence in the body for longer durations. Although there might be situations in which clinicians need to prescribe immediate-release and ER/LA opioids together (e.g., transitioning patients from ER/LA opioids to immediate-release opioids by temporarily using lower dosages of both), in general, avoiding the use of immediate-release opioids in combination with ER/LA opioids is preferable, given potentially increased risk and diminishing returns of such an approach for chronic pain.

When an ER/LA opioid is prescribed, using one with predictable pharmacokinetics and pharmacodynamics is preferred to minimize unintentional overdose risk. In particular, unusual characteristics of methadone and of transdermal fentanyl make safe prescribing of these medications for pain especially challenging.

- **Methadone should not be the first choice for an ER/LA opioid.** Only clinicians who are familiar with methadone's unique risk profile and who are prepared to educate and closely monitor their patients, including risk assessment for QT prolongation and consideration of electrocardiographic monitoring, should consider prescribing methadone for pain. A clinical practice guideline that contains further guidance regarding methadone prescribing for pain has been published previously (191).

- **Because dosing effects of transdermal fentanyl are often misunderstood by both clinicians and patients, only clinicians who are familiar with the dosing and absorption properties of transdermal fentanyl and are prepared to educate their patients about its use should consider prescribing it.**

5. **When opioids are started, clinicians should prescribe the lowest effective dosage.** Clinicians should use caution when prescribing opioids at any dosage, should carefully reassess evidence of individual benefits and risks when considering increasing dosage to ≥50 morphine milligram equivalents (MME)/day, and should avoid increasing dosage to ≥90 MME/day or carefully justify a decision to titrate dosage to ≥90 MME/day (recommendation category: A, evidence type: 3).

Benefits of high-dose opioids for chronic pain are not established. The clinical evidence review found only one study (84) addressing effectiveness of dose titration for outcomes related to pain control, function, and quality of life (KQ3). This randomized trial found no difference in pain or function between a more liberal opioid dose escalation strategy and maintenance of current dosage. (These groups were prescribed average dosages of 52 and 40 MME/day, respectively, at the end of the trial.) At the same time, risks for serious harms...
related to opioid therapy increase at higher opioid dosage. The clinical evidence review found that higher opioid dosages are associated with increased risks for motor vehicle injury, opioid use disorder, and overdose (KQ2). The clinical and contextual evidence reviews found that opioid overdose risk increases in a dose-response manner, that dosages of 50–<100 MME/day have been found to increase risks for opioid overdose by factors of 1.9 to 4.6 compared with dosages of 1–<20 MME/day, and that dosages ≥100 MME/day are associated with increased risks of overdose 2.0–8.9 times the risk at 1–<20 MME/day. In a national sample of Veterans Health Administration patients with chronic pain who were prescribed opioids, mean prescribed opioid dosage among patients who died from opioid overdose was 98 MME (median 60 MME) compared with mean prescribed opioid dosage of 48 MME (median 25 MME) among patients not experiencing fatal overdose (127).

The contextual evidence review found that although there is not a single dosage threshold below which overdose risk is eliminated, holding dosages <50 MME/day would likely reduce risk among a large proportion of patients who would experience fatal overdose at higher prescribed dosages. Experts agreed that lower dosages of opioids reduce the risk for overdose, but that a single dosage threshold for safe opioid use could not be identified. Experts noted that daily opioid dosages close to or greater than 100 MME/day are associated with significant risks, that dosages <50 MME/day are safer than dosages of 50–100 MME/day, and that dosages <20 MME/day are safer than dosages of 20–50 MME/day. One expert thought that a specific dosage at which the benefit/risk ratio of opioid therapy decreases could not be identified. Most experts agreed that, in general, increasing dosages to 50 or more MME/day increases overdose risk without necessarily adding benefits for pain control or function and that clinicians should carefully reassess evidence of individual benefits and risks when considering increasing opioid dosages to ≥50 MME/day. Most experts also agreed that opioid dosages should not be increased to ≥90 MME/day without careful justification based on diagnosis and on individualized assessment of benefits and risks.

When opioids are used for chronic pain outside of active cancer, palliative, and end-of-life care, clinicians should start opioids at the lowest possible effective dosage (the lowest starting dosage on product labeling for patients not already taking opioids and according to product labeling guidance regarding tolerance for patients already taking opioids). Clinicians should use additional caution when initiating opioids for patients aged ≥65 years and for patients with renal or hepatic insufficiency because decreased clearance of drugs in these patients can result in accumulation of drugs to toxic levels. Clinicians should use caution when increasing opioid dosages and increase dosage by the smallest practical amount because overdose risk increases with increases in opioid dosage. Although there is limited evidence to recommend specific intervals for dosage titration, a previous guideline recommended waiting at least five half-lives before increasing dosage and waiting at least a week before increasing dosage of methadone to make sure that full effects of the previous dosage are evident (31). Clinicians should re-evaluate patients after increasing dosage for changes in pain, function, and risk for harm (see Recommendation 7). Before increasing total opioid dosage to ≥50 MME/day, clinicians should reassess whether opioid treatment is meeting the patient’s treatment goals (see Recommendation 2). If a patient’s opioid dosage for all sources of opioids combined reaches or exceeds 50 MME/day, clinicians should implement additional precautions, including increased frequency of follow-up (see Recommendation 7) and considering offering naloxone and overdose prevention education to both patients and the patients’ household members (see Recommendation 8). Clinicians should avoid increasing opioid dosages to ≥90 MME/day or should carefully justify a decision to increase dosage to ≥90 MME/day based on individualized assessment of benefits and risks and weighing factors such as diagnosis, incremental benefits for pain and function relative to harms as dosages approach 90 MME/day, other treatments and effectiveness, and recommendations based on consultation with pain specialists. If patients do not experience improvement in pain and function at ≥90 MME/day, or if there are escalating dosage requirements, clinicians should discuss other approaches to pain management with the patient, consider working with patients to taper opioids to a lower dosage or to taper and discontinue opioids (see Recommendation 7), and consider consulting a pain specialist. Some states require clinicians to implement clinical protocols at specific dosage levels. For example, before increasing long-term opioid therapy dosage to >120 MME/day, clinicians in Washington state must obtain consultation from a pain specialist who agrees that this is indicated and appropriate (30). Clinicians should be aware of rules related to MME thresholds and associated clinical protocols established by their states.

Established patients already taking high dosages of opioids, as well as patients transferring from other clinicians, might consider the possibility of opioid dosage reduction to be anxiety-provoking, and tapering opioids can be especially challenging after years on high dosages because of physical and psychological dependence. However, these patients should be offered the opportunity to re-evaluate their continued use of opioids at high dosages in light of recent evidence regarding the association of opioid dosage and overdose risk. Clinicians should explain in a nonjudgmental manner to patients already taking high opioid dosages (≥90 MME/day) that there is
now an established body of scientific evidence showing that overdose risk is increased at higher opioid dosages. Clinicians should empathically review benefits and risks of continued high-dosage opioid therapy and should offer to work with the patient to taper opioids to safer dosages. For patients who agree to taper opioids to lower dosages, clinicians should collaborate with the patient on a tapering plan (see Recommendation 7). Experts noted that patients tapering opioids after taking them for years might require very slow opioid tapers as well as pauses in the taper to allow gradual accommodation to lower opioid dosages. Clinicians should remain alert to signs of anxiety, depression, and opioid use disorder (see Recommendations 8 and 12) that might be unmasked by an opioid taper and arrange for management of these co-morbidities. For patients agreeing to taper to lower opioid dosages as well as for those remaining on high opioid dosages, clinicians should establish goals with the patient for continued opioid therapy (see Recommendation 2), maximize pain treatment with nonpharmacologic and nonopioid pharmacologic treatments as appropriate (see Recommendation 1), and consider consulting a pain specialist as needed to assist with pain management.

6. Long-term opioid use often begins with treatment of acute pain. When opioids are used for acute pain, clinicians should prescribe the lowest effective dose of immediate-release opioids and should prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids. Three days or less will often be sufficient; more than seven days will rarely be needed (recommendation category: A, evidence type: 4).

The clinical evidence review found that opioid use for acute pain (i.e., pain with abrupt onset and caused by an injury or other process that is not ongoing) is associated with long-term opioid use, and that a greater amount of early opioid exposure is associated with greater risk for long-term use (KQ5). Several guidelines on opioid prescribing for acute pain from emergency departments (192–194) and other settings (195,196) have recommended prescribing ≤3 days of opioids in most cases, whereas others have recommended ≤7 days (197) or ≤14 days (30). Because physical dependence on opioids is an expected physiologic response in patients exposed to opioids for more than a few days (contextual evidence review), limiting days of opioids prescribed also should minimize the need to taper opioids to prevent distressing or unpleasant withdrawal symptoms. Experts noted that more than a few days of exposure to opioids significantly increases hazards, that each day of unnecessary opioid use increases likelihood of physical dependence without adding benefit, and that prescriptions with fewer days’ supply will minimize the number of pills available for unintentional or intentional diversion.

Experts agreed that when opioids are needed for acute pain, clinicians should prescribe opioids at the lowest effective dose and for no longer than the expected duration of pain severe enough to require opioids to minimize unintentional initiation of long-term opioid use. The lowest effective dose can be determined using product labeling as a starting point with calibration as needed based on the severity of pain and on other clinical factors such as renal or hepatic insufficiency (see Recommendation 8). Experts thought, based on clinical experience regarding anticipated duration of pain severe enough to require an opioid, that in most cases of acute pain not related to surgery or trauma, a ≤3 days’ supply of opioids will be sufficient. For example, in one study of the course of acute low back pain (not associated with malignancies, infections, spondylarthropathies, fractures, or neurological signs) in a primary care setting, there was a large decrease in pain until the fourth day after treatment with paracetamol, with smaller decreases thereafter (198). Some experts thought that because some types of acute pain might require more than 3 days of opioid treatment, it would be appropriate to recommend a range of ≤3–5 days or ≤3–7 days when opioids are needed. Some experts thought that a range including 7 days was too long given the expected course of severe acute pain for most acute pain syndromes seen in primary care.

Acute pain can often be managed without opioids. It is important to evaluate the patient for reversible causes of pain, for underlying etiologies with potentially serious sequelae, and to determine appropriate treatment. When the diagnosis and severity of nontraumatic, nonsurgical acute pain are reasonably assumed to warrant the use of opioids, clinicians should prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids, often 3 days or less, unless circumstances clearly warrant additional opioid therapy. More than 7 days will rarely be needed. Opioid treatment for post-surgical pain is outside the scope of this guideline but has been addressed elsewhere (30). Clinicians should not prescribe additional opioids to patients “just in case” pain continues longer than expected. Clinicians should re-evaluate the subset of patients who experience severe acute pain that continues longer than the expected duration to confirm or revise the initial diagnosis and to adjust management accordingly. Given longer half-lives and longer duration of effects (e.g., respiratory depression) with ER/LA opioids such as methadone, fentanyl patches, or extended release versions of opioids such as oxycodone, oxymorphone, or morphine, clinicians should not prescribe ER/LA opioids for the treatment of acute pain.
7. Clinicians should evaluate benefits and harms with patients within 1 to 4 weeks of starting opioid therapy for chronic pain or of dose escalation. Clinicians should evaluate benefits and harms of continued therapy with patients every 3 months or more frequently. If benefits do not outweigh harms of continued opioid therapy, clinicians should optimize other therapies and work with patients to taper opioids to lower dosages or to taper and discontinue opioids (recommendation category: A, evidence type: 4).

Although the clinical evidence review did not find studies evaluating the effectiveness of more frequent monitoring intervals (KQ4), it did find that continuing opioid therapy for 3 months substantially increases risk for opioid use disorder (KQ2); therefore, follow-up earlier than 3 months might be necessary to provide the greatest opportunity to prevent the development of opioid use disorder. In addition, risk for overdose associated with ER/LA opioids might be particularly high during the first 2 weeks of treatment (KQ3). The contextual evidence review found that patients who do not have pain relief with opioids at 1 month are unlikely to experience pain relief with opioids at 6 months. Although evidence is insufficient to determine at what point within the first 3 months of opioid therapy the risks for opioid use disorder increase, reassessment of pain and function within 1 month of initiating opioids provides an opportunity to minimize risks of long-term opioid use by discontinuing opioids among patients not receiving a clear benefit from these medications. Experts noted that risks for opioid overdose are greatest during the first 3–7 days after opioid initiation or increase in dosage, particularly when methadone or transdermal fentanyl are prescribed; that follow-up within 3 days is appropriate when initiating or increasing the dosage of methadone; and that follow-up within 1 week might be appropriate when initiating or increasing the dosage of other ER/LA opioids.

Clinicians should evaluate patients to assess benefits and harms of opioids within 1 to 4 weeks of starting long-term opioid therapy or of dose escalation. Clinicians should consider follow-up intervals within the lower end of this range when ER/LA opioids are started or increased or when total daily opioid dosage is ≥50 MME/day. Shorter follow-up intervals (within 3 days) should be strongly considered when starting or increasing the dosage of methadone. At follow-up, clinicians should assess benefits in function, pain control, and quality of life using tools such as the three-item “Pain average, interference with Enjoyment of life, and interference with General activity” (PEG) Assessment Scale (186) and/or asking patients about progress toward functional goals that have meaning for them (see Recommendation 2). Clinicians should also ask patients about common adverse effects such as constipation and drowsiness (see Recommendation 3), as well as asking about and assessing for effects that might be early warning signs for more serious problems such as overdose (e.g., sedation or slurred speech) or opioid use disorder (e.g., craving, wanting to take opioids in greater quantities or more frequently than prescribed, or difficulty controlling use). Clinicians should ask patients about their preferences for continuing opioids, given their effects on pain and function relative to any adverse effects experienced.

Because of potential changes in the balance of benefits and risks of opioid therapy over time, clinicians should regularly reassess all patients receiving long-term opioid therapy, including patients who are new to the clinician but on long-term opioid therapy, at least every 3 months. At reassessment, clinicians should determine whether opioids continue to meet treatment goals, including sustained improvement in pain and function, whether the patient has experienced common or serious adverse events or early warning signs of serious adverse events, signs of opioid use disorder (e.g., difficulty controlling use, work or family problems related to opioid use), whether benefits of opioids continue to outweigh risks, and whether opioid dosage can be reduced or opioids can be discontinued. Ideally, these reassessments would take place in person and be conducted by the prescribing clinician. In practice contexts where virtual visits are part of standard care (e.g., in remote areas where distance or other issues make follow-up visits challenging), follow-up assessments that allow the clinician to communicate with and observe the patient through video and audio could be conducted, with in-person visits occurring at least once per year. Clinicians should re-evaluate patients who are exposed to greater risk of opioid use disorder or overdose (e.g., patients with depression or other mental health conditions, a history of substance use disorder, a history of overdose, taking ≥50 MME/day, or taking other central nervous system depressants with opioids) more frequently than every 3 months. If clinically meaningful improvements in pain and function are not sustained, if patients are taking high-risk regimens (e.g., dosages ≥50 MME/day or opioids combined with benzodiazepines) without evidence of benefit, if patients believe benefits no longer outweigh risks or if they request dosage reduction or discontinuation, or if patients experience overdose or other serious adverse events (e.g., an event leading to hospitalization or disability) or warning signs of serious adverse events, clinicians should work with patients to reduce opioid dosage or to discontinue opioids when possible. Clinicians should maximize pain treatment with nonpharmacologic and nonopioid pharmacologic treatments as appropriate (see Recommendation 1) and consider consulting a pain specialist as needed to assist with pain management.
Considerations for Tapering Opioids

Although the clinical evidence review did not find high-quality studies comparing the effectiveness of different tapering protocols for use when opioid dosage is reduced or opioids are discontinued (KQ3), tapers reducing weekly dosage by 10%–50% of the original dosage have been recommended by other clinical guidelines (199), and a rapid taper over 2–3 weeks has been recommended in the case of a severe adverse event such as overdose (30). Experts noted that taps slower than 10% per week (e.g., 10% per month) also might be appropriate and better tolerated than more rapid taps, particularly when patients have been taking opioids for longer durations (e.g., for years). Opioid withdrawal during pregnancy has been associated with spontaneous abortion and premature labor.

When opioids are reduced or discontinued, a taper slow enough to minimize symptoms and signs of opioid withdrawal (e.g., drug craving, anxiety, insomnia, abdominal pain, vomiting, diarrhea, diaphoresis, mydriasis, tremor, tachycardia, or piloerection) should be used. A decrease of 10% of the original dose per week is a reasonable starting point; experts agreed that tapering plans may be individualized based on patient goals and concerns. Experts noted that at times, taps might have to be paused and restarted again when the patient is ready and might have to be slowed once patients reach low dosages. Tapers may be considered successful as long as the patient is making progress. Once the smallest available dose is reached, the interval between doses can be extended. Opioids may be stopped when taken less frequently than once a day. More rapid taps might be needed for patient safety under certain circumstances (e.g., for patients who have experienced overdose on their current dosage). Ultrarapid detoxification under anesthesia is associated with substantial risks, including death, and should not be used (200). Clinicians should access appropriate expertise if considering tapering opioids during pregnancy because of possible risk to the pregnant patient and to the fetus if the patient goes into withdrawal. Patients who are not taking opioids (including patients who are diverting all opioids they obtain) do not require taps. Clinicians should discuss with patients undergoing tapering the increased risk for overdose on abrupt return to a previously prescribed higher dose. Primary care clinicians should collaborate with mental health providers and with other specialists as needed to optimize nonopioid pain management (see Recommendation 1), as well as psychosocial support for anxiety related to the taper. More detailed guidance on tapering, including management of withdrawal symptoms has been published previously (30,201). If a patient exhibits signs of opioid use disorder, clinicians should offer or arrange for treatment of opioid use disorder (see Recommendation 12) and consider offering naloxone for overdose prevention (see Recommendation 8).

Assessing Risk and Addressing Harms of Opioid Use

8. Before starting and periodically during continuation of opioid therapy, clinicians should evaluate risk factors for opioid-related harms. Clinicians should incorporate into the management plan strategies to mitigate risk, including considering offering naloxone when factors that increase risk for opioid overdose, such as history of overdose, history of substance use disorder, higher opioid dosages (≥50 MME/day), or concurrent benzodiazepine use, are present (recommendation category: A, evidence type: 4).

The clinical evidence review found insufficient evidence to determine how harms of opioids differ depending on patient demographics or patient comorbidities (KQ2). However, based on the contextual evidence review and expert opinion, certain risk factors are likely to increase susceptibility to opioid-associated harms and warrant incorporation of additional strategies into the management plan to mitigate risk. Clinicians should assess these risk factors periodically, with frequency varying by risk factor and patient characteristics. For example, factors that vary more frequently over time, such as alcohol use, require more frequent follow up. In addition, clinicians should consider offering naloxone, re-evaluating patients more frequently (see Recommendation 7), and referring to pain and/or behavioral health specialists when factors that increase risk for harm, such as history of overdose, history of substance use disorder, higher dosages of opioids (≥50 MME/day), and concurrent use of benzodiazepines with opioids, are present.

Patients with Sleep-Disordered Breathing, Including Sleep Apnea

Risk factors for sleep-disordered breathing include congestive heart failure, and obesity. Experts noted that careful monitoring and cautious dose titration should be used if opioids are prescribed for patients with mild sleep-disordered breathing. Clinicians should avoid prescribing opioids to patients with moderate or severe sleep-disordered breathing whenever possible to minimize risks for opioid overdose (contextual evidence review).

Pregnant Women

Opioids used in pregnancy might be associated with additional risks to both mother and fetus. Some studies have shown an association of opioid use in pregnancy with stillbirth, poor fetal growth, pre-term delivery, and birth defects (contextual evidence review). Importantly, in some cases, opioid use during pregnancy leads to neonatal opioid withdrawal syndrome. Clinicians and patients together should carefully weigh risks and benefits when making decisions.
about whether to initiate opioid therapy for chronic pain during pregnancy. In addition, before initiating opioid therapy for chronic pain for reproductive-age women, clinicians should discuss family planning and how long-term opioid use might affect any future pregnancy. For pregnant women already receiving opioids, clinicians should access appropriate expertise if considering tapering opioids because of possible risk to the pregnant patient and to the fetus if the patient goes into withdrawal (see Recommendation 7). For pregnant women with opioid use disorder, medication-assisted therapy with buprenorphine or methadone has been associated with improved maternal outcomes and should be offered (202) (see Recommendation 12). Clinicians caring for pregnant women receiving opioids for pain or receiving buprenorphine or methadone for opioid use disorder should arrange for delivery at a facility prepared to monitor, evaluate for, and treat neonatal opioid withdrawal syndrome. In instances when travel to such a facility would present an undue burden on the pregnant woman, it is appropriate to deliver locally, monitor and evaluate the newborn for neonatal opioid withdrawal syndrome, and transfer the newborn for additional treatment if needed. Neonatal toxicity and death have been reported in breast-feeding infants whose mothers are taking codeine (contextual evidence review); previous guidelines have recommended that codeine be avoided whenever possible among mothers who are breast feeding and, if used, should be limited to the lowest possible dose and to a 4-day supply (203).

**Patients with Renal or Hepatic Insufficiency**

Clinicians should use additional caution and increased monitoring (see Recommendation 7) to minimize risks of opioids prescribed for patients with renal or hepatic insufficiency, given their decreased ability to process and excrete drugs, susceptibility to accumulation of opioids, and reduced therapeutic window between safe dosages and dosages associated with respiratory depression and overdose (contextual evidence review; see Recommendations 4, 5, and 7).

**Patients Aged ≥65 Years**

Inadequate pain treatment among persons aged ≥65 years has been documented (204). Pain management for older patients can be challenging given increased risks of both nonopioid pharmacologic therapies (see Recommendation 1) and opioid therapy in this population. Given reduced renal function and medication clearance even in the absence of renal disease, patients aged ≥65 years might have increased susceptibility to accumulation of opioids and a smaller therapeutic window between safe dosages and dosages associated with respiratory depression and overdose (contextual evidence review). Some older adults suffer from cognitive impairment, which can increase risk for medication errors and make opioid-related confusion more dangerous. In addition, older adults are more likely than younger adults to experience co-morbid medical conditions and more likely to receive multiple medications, some of which might interact with opioids (such as benzodiazepines). Clinicians should use additional caution and increased monitoring (see Recommendations 4, 5, and 7) to minimize risks of opioids prescribed for patients aged ≥65 years. Experts suggested that clinicians educate older adults receiving opioids to avoid risky medication-related behaviors such as obtaining controlled medications from multiple prescribers and saving unused medications. Clinicians should also implement interventions to mitigate common risks of opioid therapy among older adults, such as exercise or bowel regimens to prevent constipation, risk assessment for falls, and patient monitoring for cognitive impairment.

**Patients with Mental Health Conditions**

Because psychological distress frequently interferes with improvement of pain and function in patients with chronic pain, using validated instruments such as the Generalized Anxiety Disorder (GAD)-7 and the Patient Health Questionnaire (PHQ)-9 or the PHQ-4 to assess for anxiety, post-traumatic stress disorder, and/or depression (205), might help clinicians improve overall pain treatment outcomes. Experts noted that clinicians should use additional caution and increased monitoring (see Recommendation 7) to lessen the increased risk for opioid use disorder among patients with mental health conditions (including depression, anxiety disorders, and PTSD), as well as increased risk for drug overdose among patients with depression. Previous guidelines have noted that opioid therapy should not be initiated during acute psychiatric instability or uncontrolled suicide risk, and that clinicians should consider behavioral health specialist consultation for any patient with a history of suicide attempt or psychiatric disorder (31). In addition, patients with anxiety disorders and other mental health conditions are more likely to receive benzodiazepines, which can exacerbate opioid-induced respiratory depression and increase risk for overdose (see Recommendation 11). Clinicians should ensure that treatment for depression and other mental health conditions is optimized, consulting with behavioral health specialists when needed. Treatment for depression can improve pain symptoms as well as depression and might decrease overdose risk (contextual evidence review). For treatment of chronic pain in patients with depression, clinicians should strongly consider using tricyclic or SNRI antidepressants for analgesic as well as antidepressant effects if these medications are not otherwise contraindicated (see Recommendation 1).
Patients with Substance Use Disorder

Illicit drugs and alcohol are listed as contributory factors on a substantial proportion of death certificates for opioid-related overdose deaths (contextual evidence review). Previous guidelines have recommended screening or risk assessment tools to identify patients at higher risk for misuse or abuse of opioids. However, the clinical evidence review found that currently available risk-stratification tools (e.g., Opioid Risk Tool, Screener and Opioid Assessment for Patients with Pain Version 1, SOAPP-R, and Brief Risk Interview) show insufficient accuracy for classification of patients as at low or high risk for abuse or misuse (KQ4). Clinicians should always exercise caution when considering or prescribing opioids for any patient with chronic pain outside of active cancer, palliative, and end-of-life care and should not overestimate the ability of these tools to rule out risks from long-term opioid therapy.

Clinicians should ask patients about their drug and alcohol use. Single screening questions can be used (206). For example, the question “How many times in the past year have you used an illegal drug or used a prescription medication for nonmedical reasons?” (with an answer of one or more considered positive) was found in a primary care setting to be 100% sensitive and 73.5% specific for the detection of a drug use disorder compared with a standardized diagnostic interview (207). Validated screening tools such as the Drug Abuse Screening Test (DAST) (208) and the Alcohol Use Disorders Identification Test (AUDIT) (209) can also be used. Clinicians should use PDMP data (see Recommendation 9) and drug testing (see Recommendation 10) as appropriate to assess for concurrent substance use that might place patients at higher risk for opioid use disorder and overdose. Clinicians should also provide specific counseling on increased risks for overdose when opioids are combined with other drugs or alcohol (see Recommendation 3) and ensure that patients receive effective treatment for substance use disorders when needed (see Recommendation 12).

The clinical evidence review found insufficient evidence to determine how harms of opioids differ depending on past or current substance use disorder (KQ2), although a history of substance use disorder was associated with misuse. Similarly, based on contextual evidence, patients with drug or alcohol use disorders are likely to experience greater risks for opioid use disorder and overdose than persons without these conditions. If clinicians consider opioid therapy for chronic pain outside of active cancer, palliative, and end-of-life care for patients with drug or alcohol use disorders, they should discuss increased risks for opioid use disorder and overdose with patients, carefully consider whether benefits of opioids outweigh increased risks, and incorporate strategies to mitigate risk into the management plan, such as considering offering naloxone (see Offering Naloxone to Patients When Factors That Increase Risk for Opioid-Related Harms Are Present) and increasing frequency of monitoring (see Recommendation 7) when opioids are prescribed. Because pain management in patients with substance use disorder can be complex, clinicians should consider consulting substance use disorder specialists and pain specialists regarding pain management for persons with active or recent past history of substance abuse. Experts also noted that clinicians should communicate with patients’ substance use disorder treatment providers if opioids are prescribed.

Patients with Prior Nonfatal Overdose

Although studies were not identified that directly addressed the risk for overdose among patients with prior nonfatal overdose who are prescribed opioids, based on clinical experience, experts thought that prior nonfatal overdose would substantially increase risk for future nonfatal or fatal opioid overdose. If patients experience nonfatal opioid overdose, clinicians should work with them to reduce opioid dosage and to discontinue opioids when possible (see Recommendation 7). If clinicians continue opioid therapy for chronic pain outside of active cancer, palliative, and end-of-life care in patients with prior opioid overdose, they should discuss increased risks for overdose with patients, carefully consider whether benefits of opioids outweigh substantial risks, and incorporate strategies to mitigate risk into the management plan, such as considering offering naloxone (see Offering Naloxone to Patients When Factors That Increase Risk for Opioid-Related Harms Are Present) and increasing frequency of monitoring (see Recommendation 7) when opioids are prescribed.

Offering Naloxone to Patients When Factors That Increase Risk for Opioid-Related Harms Are Present

Naloxone is an opioid antagonist that can reverse severe respiratory depression; its administration by lay persons, such as friends and family of persons who experience opioid overdose, can save lives. Naloxone precipitates acute withdrawal among patients physically dependent on opioids. Serious adverse effects, such as pulmonary edema, cardiovascular instability, and seizures, have been reported but are rare at doses consistent with labeled use for opioid overdose (210). The contextual evidence review did not find any studies on effectiveness of prescribing naloxone for overdose prevention among patients prescribed opioids for chronic pain. However, there is evidence for effectiveness of naloxone provision in preventing opioid-related overdose death at the community level through community-based distribution (e.g., through overdose education and naloxone distribution programs in community service agencies) to persons at risk for overdose.
(mostly due to illicit opiate use), and it is plausible that effectiveness would be observed when naloxone is provided in the clinical setting as well. Experts agreed that it is preferable not to initiate opioid treatment when factors that increase risk for opioid-related harms are present. Opinions diverged about the likelihood of naloxone being useful to patients and the circumstances under which it should be offered. However, most experts agreed that clinicians should consider offering naloxone when prescribing opioids to patients at increased risk for overdose, including patients with a history of overdose, patients with a history of substance use disorder, patients taking benzodiazepines with opioids (see Recommendation 11), patients at risk for returning to a high dose to which they are no longer tolerant (e.g., patients recently released from prison), and patients taking higher dosages of opioids (≥50 MME/day). Practices should provide education on overdose prevention and naloxone use to patients receiving naloxone prescriptions and to members of their households. Experts noted that naloxone co-prescribing can be facilitated by clinics or practices with resources to provide naloxone training and by collaborative practice models with pharmacists. Resources for prescribing naloxone in primary care settings can be found through Prescribe to Prevent at http://prescribetoprevent.org.

9. Clinicians should review the patient's history of controlled substance prescriptions using state prescription drug monitoring program (PDMP) data to determine whether the patient is receiving opioid dosages or dangerous combinations that put him or her at high risk for overdose. Clinicians should review PDMP data when starting opioid therapy for chronic pain and periodically during opioid therapy for chronic pain, ranging from every prescription to every 3 months (recommendation category: A, evidence type: 4).

PDMPs are state-based databases that collect information on controlled prescription drugs dispensed by pharmacies in most states and, in select states, by dispensing physicians as well. In addition, some clinicians employed by the federal government, including some clinicians in the Indian Health Care Delivery System, are not licensed in the states where they practice, and do not have access to PDMP data. Certain states require clinicians to review PDMP data prior to writing each opioid prescription (see state-level PDMP-related policies on the National Alliance for Model State Drug Laws website at http://www.namsdl.org/prescription-monitoring-programs.cfm). The clinical evidence review did not find studies evaluating the effectiveness of PDMPs on outcomes related to overdose, addiction, abuse, or misuse (KQ4). However, even though evidence is limited on the effectiveness of PDMP implementation at the state level on prescribing and mortality outcomes (28), the contextual evidence review found that most fatal overdoses were associated with patients receiving opioids from multiple prescribers and/or with patients receiving high total daily opioid dosages; information on both of these risk factors for overdose are available to prescribers in the PDMP. PDMP data also can be helpful when patient medication history is not otherwise available (e.g., for patients from other locales) and when patients transition care to a new clinician. The contextual evidence review also found that PDMP information could be used in a way that is harmful to patients. For example, it has been used to dismiss patients from clinician practices (211), which might adversely affect patient safety.

The contextual review found variation in state policies that affect timeliness of PDMP data (and therefore benefits of reviewing PDMP data) as well as time and workload for clinicians in accessing PDMP data. In states that permit delegating access to other members of the health care team, workload for prescribers can be reduced. These differences might result in a different balance of benefits to clinician workload in different states. Experts agreed that PDMPs are useful tools that should be consulted when starting a patient on opioid therapy and periodically during long-term opioid therapy. However, experts disagreed on how frequently clinicians should check the PDMP during long-term opioid therapy, given PDMP access issues and the lag time in reporting in some states. Most experts agreed that PDMP data should be reviewed every 3 months or more frequently during long-term opioid therapy. A minority of experts noted that, given the current burden of accessing PDMP data in some states and the lack of evidence surrounding the most effective interval for PDMP review to improve patient outcomes, annual review of PDMP data during long-term opioid therapy would be reasonable when factors that increase risk for opioid-related harms are not present.

Clinicians should review PDMP data for opioids and other controlled medications patients might have received from additional prescribers to determine whether a patient is receiving high total opioid dosages or dangerous combinations (e.g., opioids combined with benzodiazepines) that put him or her at high risk for overdose. Ideally, PDMP data should be reviewed before every opioid prescription. This is recommended in all states with well-functioning PDMPs and where PDMP access policies make this practicable (e.g., clinician and delegate access permitted), but it is not currently possible in states without functional PDMPs or in those that do not permit certain prescribers to access them. As vendors and practices facilitate integration of PDMP information into regular clinical workflow (e.g., data made available in electronic health records), clinicians’ ease of access in reviewing PDMP data is expected to improve.
In addition, improved timeliness of PDMP data will improve their value in identifying patient risks.

If patients are found to have high opioid dosages, dangerous combinations of medications, or multiple controlled substance prescriptions written by different clinicians, several actions can be taken to augment clinicians’ abilities to improve patient safety:

- Clinicians should discuss information from the PDMP with their patient and confirm that the patient is aware of the additional prescriptions. Occasionally, PDMP information can be incorrect (e.g., if the wrong name or birthdate has been entered, the patient uses a nickname or maiden name, or another person has used the patient’s identity to obtain prescriptions).
- Clinicians should discuss safety concerns, including increased risk for respiratory depression and overdose, with patients found to be receiving opioids from more than one prescriber or receiving medications that increase risk when combined with opioids (e.g., benzodiazepines) and consider offering naloxone (see Recommendation 8).
- Clinicians should avoid prescribing opioids and benzodiazepines concurrently whenever possible. Clinicians should communicate with others managing the patient to discuss the patient’s needs, prioritize patient goals, weigh risks of concurrent benzodiazepine and opioid exposure, and coordinate care (see Recommendation 11).
- Clinicians should calculate the total MME/day for concurrent opioid prescriptions to help assess the patient’s overdose risk (see Recommendation 5). If patients are found to be receiving high total daily dosages of opioids, clinicians should discuss their safety concerns with the patient, consider tapering to a safer dosage (see Recommendations 5 and 7), and consider offering naloxone (see Recommendation 8).
- Clinicians should discuss safety concerns with other clinicians who are prescribing controlled substances for their patient. Ideally clinicians should first discuss concerns with their patient and inform him or her that they plan to coordinate care with the patient’s other prescribers to improve the patient’s safety.
- Clinicians should consider the possibility of a substance use disorder and discuss concerns with their patient (see Recommendation 12).
- If clinicians suspect their patient might be sharing or selling opioids and not taking them, clinicians should consider urine drug testing to assist in determining whether opioids can be discontinued without causing withdrawal (see Recommendations 7 and 10). A negative drug test for prescribed opioids might indicate the patient is not taking prescribed opioids, although clinicians should consider other possible reasons for this test result (see Recommendation 10).

Experts agreed that clinicians should not dismiss patients from their practice on the basis of PDMP information. Doing so can adversely affect patient safety, could represent patient abandonment, and could result in missed opportunities to provide potentially lifesaving information (e.g., about risks of opioids and overdose prevention) and interventions (e.g., safer prescriptions, nonopioid pain treatment [see Recommendation 1], naloxone [see Recommendation 8], and effective treatment for substance use disorder [see Recommendation 12]).

10. When prescribing opioids for chronic pain, clinicians should use urine drug testing before starting opioid therapy and consider urine drug testing at least annually to assess for prescribed medications as well as other controlled prescription drugs and illicit drugs (recommendation category: B, evidence type: 4).

Concurrent use of opioid pain medications with other opioid pain medications, benzodiazepines, or heroin can increase patients’ risk for overdose. Urine drug tests can provide information about drug use that is not reported by the patient. In addition, urine drug tests can assist clinicians in identifying when patients are not taking opioids prescribed for them, which might in some cases indicate diversion or other clinically important issues such as difficulties with adverse effects. Urine drug tests do not provide accurate information about how much or what dose of opioids or other drugs a patient took. The clinical evidence review did not find studies evaluating the effectiveness of urine drug screening for risk mitigation during opioid prescribing for pain (KQ4). The contextual evidence review found that urine drug testing can provide useful information about patients assumed not to be using unreported drugs. Urine drug testing results can be subject to misinterpretation and might sometimes be associated with practices that might harm patients (e.g., stigmatization, inappropriate termination from care). Routine use of urine drug tests with standardized policies at the practice or clinic level might destigmatize their use. Although random drug testing also might destigmatize urine drug testing, experts thought that truly random testing was not feasible in clinical practice. Some clinics obtain a urine specimen at every visit, but only send it for testing on a random schedule. Experts noted that in addition to direct costs of urine drug testing, which often are not covered fully by insurance and can be a burden for patients, clinician time is needed to interpret, confirm, and communicate results.

Experts agreed that prior to starting opioids for chronic pain and periodically during opioid therapy, clinicians should
use urine drug testing to assess for prescribed opioids as well as other controlled substances and illicit drugs that increase risk for overdose when combined with opioids, including nonprescribed opioids, benzodiazepines, and heroin. There was some difference of opinion among experts as to whether this recommendation should apply to all patients, or whether this recommendation should entail individual decision making with different choices for different patients based on values, preferences, and clinical situations. While experts agreed that clinicians should use urine drug testing before initiating opioid therapy for chronic pain, they disagreed on how frequently urine drug testing should be conducted during long-term opioid therapy. Most experts agreed that urine drug testing at least annually for all patients was reasonable. Some experts noted that this interval might be too long in some cases and too short in others, and that the follow-up interval should be left to the discretion of the clinician. Previous guidelines have recommended more frequent urine drug testing in patients thought to be at higher risk for substance use disorder (30). However, experts thought that predicting risk prior to urine drug testing is challenging and that currently available tools do not allow clinicians to reliably identify patients who are at low risk for substance use disorder.

In most situations, initial urine drug testing can be performed with a relatively inexpensive immunoassay panel for commonly prescribed opioids and illicit drugs. Patients prescribed less commonly used opioids might require specific testing for those agents. The use of confirmatory testing adds substantial costs and should be based on the need to detect specific opioids that cannot be identified on standard immunoassays or on the presence of unexpected urine drug test results. Clinicians should be familiar with the drugs included in urine drug testing panels used in their practice and should understand how to interpret results for these drugs. For example, a positive “opiates” immunoassay detects morphine, which might reflect patient use of morphine, codeine, or heroin, but this immunoassay does not detect synthetic opioids (e.g., fentanyl or methadone) and might not detect semisynthetic opioids (e.g., oxycodone). However, many laboratories use an oxycodone immunoassay that detects oxycodone and oxymorphone. In some cases, positive results for specific opioids might reflect metabolites from opioids the patient is taking and might not mean the patient is taking the specific opioid for which the test was positive. For example, hydromorphone is a metabolite of hydrocodone, and oxymorphone is a metabolite of oxycodone. Detailed guidance on interpretation of urine drug test results, including which tests to order and expected results, drug detection time in urine, drug metabolism, and other considerations has been published previously (30). Clinicians should not test for substances for which results would not affect patient management or for which implications for patient management are unclear. For example, experts noted that there might be uncertainty about the clinical implications of a positive urine drug test for tetrahyrdocannabinol (THC). In addition, restricting confirmatory testing to situations and substances for which results can reasonably be expected to affect patient management can reduce costs of urine drug testing, given the substantial costs associated with confirmatory testing methods. Before ordering urine drug testing, clinicians should have a plan for responding to unexpected results. Clinicians should explain to patients that urine drug testing is intended to improve their safety and should also explain expected results (e.g., presence of prescribed medication and absence of drugs, including illicit drugs, not reported by the patient). Clinicians should ask patients about use of prescribed and other drugs and ask whether there might be unexpected results. This will provide an opportunity for patients to provide information about changes in their use of prescribed opioids or other drugs. Clinicians should discuss unexpected results with the local laboratory or toxicologist and with the patient. Discussion with patients prior to specific confirmatory testing can sometimes yield a candid explanation of why a particular substance is present or absent and obviate the need for expensive confirmatory testing on that visit. For example, a patient might explain that the test is negative for prescribed opioids because she felt opioids were no longer helping and discontinued them. If unexpected results are not explained, a confirmatory test using a method selective enough to differentiate specific opioids and metabolites (e.g., gas or liquid chromatography/mass spectrometry) might be warranted to clarify the situation.

Clinicians should use unexpected results to improve patient safety (e.g., change in pain management strategy [see Recommendation 1], tapering or discontinuation of opioids [see Recommendation 7], more frequent re-evaluation [see Recommendation 7], offering naloxone [see Recommendation 8], or referral for treatment for substance use disorder [see Recommendation 12], all as appropriate). If tests for prescribed opioids are repeatedly negative, confirming that the patient is not taking the prescribed opioid, clinicians can discontinue the prescription without a taper. Clinicians should not dismiss patients from care based on a urine drug test result because this could constitute patient abandonment and could have adverse consequences for patient safety, potentially including the patient obtaining opioids from alternative sources and the clinician missing opportunities to facilitate treatment for substance use disorder.

11. Clinicians should avoid prescribing opioid pain medication and benzodiazepines concurrently
Recommendations and Reports

...whenever possible (recommendation category: A, evidence type: 3).

Benzodiazepines and opioids both cause central nervous system depression and can decrease respiratory drive. Concurrent use is likely to put patients at greater risk for potentially fatal overdose. The clinical evidence review did not address risks of benzodiazepine co-prescription among patients prescribed opioids. However, the contextual evidence review found evidence in epidemiologic series of concurrent benzodiazepine use in large proportions of opioid-related overdose deaths, and a case-cohort study found concurrent benzodiazepine prescription with opioid prescription to be associated with a near quadrupling of risk for overdose death compared with opioid prescription alone (212). Experts agreed that although there are circumstances when it might be appropriate to prescribe opioids to a patient receiving benzodiazepines (e.g., severe acute pain in a patient taking long-term, stable low-dose benzodiazepine therapy), clinicians should avoid prescribing opioids and benzodiazepines concurrently whenever possible. In addition, given that other central nervous system depressants (e.g., muscle relaxants, hypnotics) can potentiate central nervous system depression associated with opioids, clinicians should consider whether benefits outweigh risks of concurrent use of these drugs. Clinicians should check the PDMP for concurrent controlled medications prescribed by other clinicians (see Recommendation 9) and should consider involving pharmacists and pain specialists as part of the management team when opioids are co-prescribed with other central nervous system depressants. Because of greater risks of benzodiazepine withdrawal relative to opioid withdrawal, and because tapering opioids can be associated with anxiety, when patients receiving both benzodiazepines and opioids require tapering to reduce risk for fatal respiratory depression, it might be safer and more practical to taper opioids first (see Recommendation 7). Clinicians should taper benzodiazepines gradually if discontinued because abrupt withdrawal can be associated with rebound anxiety, hallucinations, seizures, delirium tremens, and, in rare cases, death (contextual evidence review). A commonly used tapering schedule that has been used safely and with moderate success is a reduction of the benzodiazepine dose by 25% every 1–2 weeks (213,214). CBT increases tapering success rates and might be particularly helpful for patients struggling with a benzodiazepine taper (213). If benzodiazepines prescribed for anxiety are tapered or discontinued, or if patients receiving opioids require treatment for anxiety, evidence-based psychotherapies (e.g., CBT) and/or specific anti-depressants or other nonbenzodiazepine medications approved for anxiety should be offered. Experts emphasized that clinicians should communicate with mental health professionals managing the patient to discuss the patient’s needs, prioritize patient goals, weigh risks of concurrent benzodiazepine and opioid exposure, and coordinate care.

12. Clinicians should offer or arrange evidence-based treatment (usually medication-assisted treatment with buprenorphine or methadone in combination with behavioral therapies) for patients with opioid use disorder (recommendation category: A, evidence type: 2).

Opioid use disorder (previously classified as opioid abuse or opioid dependence) is defined in the Diagnostic and Statistical Manual of Mental Disorders, 5th edition (DSM-5) as a problematic pattern of opioid use leading to clinically significant impairment or distress, manifested by at least two defined criteria occurring within a year (http://pcssmat.org/wp-content/uploads/2014/02/5B-DSM-5-Opioid-Use-Disorder-Diagnostic-Criteria.pdf) (20).

The clinical evidence review found prevalence of opioid dependence (using DSM-IV diagnosis criteria) in primary care settings among patients with chronic pain on opioid therapy to be 3%–26% (KQ2). As found in the contextual evidence review and supported by moderate quality evidence, opioid agonist or partial agonist treatment with methadone maintenance therapy or buprenorphine has been shown to be more effective in preventing relapse among patients with opioid use disorder (151–153). Some studies suggest that using behavioral therapies in combination with these treatments can reduce opioid misuse and increase retention during maintenance therapy and improve compliance after detoxification (154,155); behavioral therapies are also recommended by clinical practice guidelines (215). The cited studies primarily evaluated patients with a history of illicit opioid use, rather than prescription opioid use for chronic pain. Recent studies among patients with prescription opioid dependence (based on DSM-IV criteria) have found maintenance therapy with buprenorphine and buprenorphine-naloxone effective in preventing relapse (216,217). Treatment need in a community is often not met by capacity to provide buprenorphine or methadone maintenance therapy (218), and patient cost can be a barrier to buprenorphine treatment because insurance coverage of buprenorphine for opioid use disorder is often limited (219). Oral or long-acting injectable formulations of naltrexone can also be used as medication-assisted treatment for opioid use disorder in nonpregnant adults, particularly for highly motivated persons (220,221). Experts agreed that clinicians prescribing opioids should identify treatment resources for opioid use disorder in the community and should work together to ensure sufficient treatment capacity for opioid use disorder at the practice level.
If clinicians suspect opioid use disorder based on patient concerns or behaviors or on findings in prescription drug monitoring program data (see Recommendation 9) or from urine drug testing (see Recommendation 10), they should discuss their concern with their patient and provide an opportunity for the patient to disclose related concerns or problems. Clinicians should assess for the presence of opioid use disorder using DSM-5 criteria (20). Alternatively, clinicians can arrange for a substance use disorder treatment specialist to assess for the presence of opioid use disorder. For patients meeting criteria for opioid use disorder, clinicians should offer or arrange for patients to receive evidence-based treatment, usually medication-assisted treatment with buprenorphine or methadone maintenance therapy in combination with behavioral therapies. Oral or long-acting injectable naltrexone, a long-acting opioid antagonist, can also be used in non-pregnant adults. Naltrexone blocks the effects of opioids if they are used but requires adherence to daily oral therapy or monthly injections. For pregnant women with opioid use disorder, medication-assisted therapy with buprenorphine (without naloxone) or methadone has been associated with improved maternal outcomes and should be offered (see Recommendation 8). Clinicians should also consider offering naloxone for overdose prevention to patients with opioid use disorder (see Recommendation 8). For patients with problematic opioid use that does not meet criteria for opioid use disorder, experts noted that clinicians can offer to taper and discontinue opioids (see Recommendation 7). For patients who choose to but are unable to taper, clinicians may reassess for opioid use disorder and offer opioid agonist therapy if criteria are met.

Physicians not already certified to provide buprenorphine in an office-based setting can undergo training to receive a waiver from the Substance Abuse and Mental Health Services Administration (SAMHSA) that allows them to prescribe buprenorphine to treat patients with opioid use disorder. Physicians prescribing opioids in communities without sufficient treatment capacity for opioid use disorder should strongly consider obtaining this waiver. Information about qualifications and the process to obtain a waiver are available from SAMHSA (222). Clinicians do not need a waiver to offer naltrexone for opioid use disorder as part of their practice.

Additional guidance has been published previously (215) on induction, use, and monitoring of buprenorphine treatment (see Part 5) and naltrexone treatment (see Part 6) for opioid use disorder and on goals, components of, and types of effective psychosocial treatment that are recommended in conjunction with pharmacological treatment of opioid use disorder (see Part 7). Clinicians unable to provide treatment themselves should arrange for patients with opioid use disorder to receive care from a substance use disorder treatment specialist, such as an office-based buprenorphine or naltrexone treatment provider, or from an opioid treatment program certified by SAMHSA to provide supervised medication-assisted treatment for patients with opioid use disorder. Clinicians should assist patients in finding qualified treatment providers and should arrange for patients to follow up with these providers, as well as arranging for ongoing coordination of care. Clinicians should not dismiss patients from their practice because of a substance use disorder because this can adversely affect patient safety and could represent patient abandonment. Identification of substance use disorder represents an opportunity for a clinician to initiate potentially life-saving interventions, and it is important for the clinician to collaborate with the patient regarding their safety to increase the likelihood of successful treatment. In addition, although identification of an opioid use disorder can alter the expected benefits and risks of opioid therapy for pain, patients with co-occurring pain and substance use disorder require ongoing pain management that maximizes benefits relative to risks. Clinicians should continue to use nonpharmacologic and nonopioid pharmacologic pain treatments as appropriate (see Recommendation 1) and consider consulting a pain specialist as needed to provide optimal pain management.

Resources to help with arranging for treatment include SAMHSA’s buprenorphine physician locator (http://buprenorphine.samhsa.gov/bwns_locator); SAMHSA’s Opioid Treatment Program Directory (http://dpt2.samhsa.gov/treatment/directory.aspx); SAMHSA’s Provider Clinical Support System for Opioid Therapies (http://pcss-o.org), which offers extensive experience in the treatment of substance use disorders and specifically of opioid use disorder, as well as expertise on the interface of pain and opioid misuse; and SAMHSA’s Provider’s Clinical Support System for Medication-Assisted Treatment (http://pcssmat.org), which offers expert physician mentors to answer questions about assessment for and treatment of substance use disorders.

Conclusions and Future Directions

Clinical guidelines represent one strategy for improving prescribing practices and health outcomes. Efforts are required to disseminate the guideline and achieve widespread adoption and implementation of the recommendations in clinical settings. CDC will translate this guideline into user-friendly materials for distribution and use by health systems, medical professional societies, insurers, public health departments, health information technology developers, and clinicians and engage in dissemination efforts. CDC has provided a
checklist for prescribing opioids for chronic pain (http://stacks.cdc.gov/view/cdc/38025), additional resources such as fact sheets (http://www.cdc.gov/drugoverdose/prescribing/resources.html), and will provide a mobile application to guide clinicians in implementing the recommendations. CDC will also work with partners to support clinician education on pain management options, opioid therapy, and risk mitigation strategies (e.g., urine drug testing). Activities such as development of clinical decision support in electronic health records to assist clinicians’ treatment decisions at the point of care; identification of mechanisms that insurers and pharmacy benefit plan managers can use to promote safer prescribing within plans; and development of clinical quality improvement measures and initiatives to improve prescribing and patient care within health systems have promise for increasing guideline adoption and improving practice. In addition, policy initiatives that address barriers to implementation of the guidelines, such as increasing accessibility of PDMP data within and across states, e-prescribing, and availability of clinicians who can offer medication-assisted treatment for opioid use disorder, are strategies to consider to enhance implementation of the recommended practices. CDC will work with federal partners and payers to evaluate strategies such as payment reform and health care delivery models that could improve patient health and safety. For example, strategies might include strengthened coverage for nonpharmacologic treatments, appropriate urine drug testing, and medication-assisted treatment; reimbursable time for patient counseling; and payment models that improve access to interdisciplinary, coordinated care.

As highlighted in the forthcoming report on the National Pain Strategy, an overarching federal effort that outlines a comprehensive population-level health strategy for addressing pain as a public health problem, clinical guidelines complement other strategies aimed at preventing illnesses and injuries that lead to pain. A draft of the National Pain Strategy has been published previously (180). These strategies include strengthening the evidence base for pain prevention and treatment strategies, reducing disparities in pain treatment, improving service delivery and reimbursement, supporting professional education and training, and providing public education. It is important that overall improvements be made in developing the workforce to address pain management in general, in addition to opioid prescribing specifically. This guideline also complements other federal efforts focused on addressing the opioid overdose epidemic including prescriber training and education, improving access to treatment for opioid use disorder, safe storage and disposal programs, utilization management mechanisms, naloxone distribution programs, law enforcement and supply reduction efforts, prescription drug monitoring program improvements, and support for community coalitions and state prevention programs.

This guideline provides recommendations that are based on the best available evidence that was interpreted and informed by expert opinion. The clinical scientific evidence informing the recommendations is low in quality. To inform future guideline development, more research is necessary to fill in critical evidence gaps. The evidence reviews forming the basis of this guideline clearly illustrate that there is much yet to be learned about the effectiveness, safety, and economic efficiency of long-term opioid therapy. As highlighted by an expert panel in a recent workshop sponsored by the National Institutes of Health on the role of opioid pain medications in the treatment of chronic pain, “evidence is insufficient for every clinical decision that a provider needs to make about the use of opioids for chronic pain” (223). The National Institutes of Health panel recommended that research is needed to improve our understanding of which types of pain, specific diseases, and patients are most likely to be associated with benefit and harm from opioid pain medications; evaluate multidisciplinary pain interventions; estimate cost-benefit; develop and validate tools for identification of patient risk and outcomes; assess the effectiveness and harms of opioid pain medications with alternative study designs; and investigate risk identification and mitigation strategies and their effects on patient and public health outcomes. It is also important to obtain data to inform the cost feasibility and cost-effectiveness of recommended actions, such as use of nonpharmacologic therapy and urine drug testing. Research that contributes to safer and more effective pain treatment can be implemented across public health entities and federal agencies (4). Additional research can inform the development of future guidelines for special populations that could not be adequately addressed in this guideline, such as children and adolescents, where evidence and guidance is needed but currently lacking. CDC is committed to working with partners to identify the highest priority research areas to build the evidence base. Yet, given that chronic pain is recognized as a significant public health problem, the risks associated with long-term opioid therapy, the availability of effective nonpharmacological and nonopioid pharmacologic treatment options for pain, and the potential for improvement in the quality of health care with the implementation of recommended practices, a guideline for prescribing is warranted with the evidence that is currently available. The balance between the benefits and the risks of long-term opioid therapy for chronic pain based on both clinical and contextual evidence is strong enough to support the issuance of category A recommendations in most cases.
CDC will revisit this guideline as new evidence becomes available to determine when evidence gaps have been sufficiently closed to warrant an update of the guideline. Until this research is conducted, clinical practice guidelines will have to be based on the best available evidence and expert opinion. This guideline is intended to improve communication between clinicians and patients about the risks and benefits of opioid therapy for chronic pain, improve the safety and effectiveness of pain treatment, and reduce the risks associated with long-term opioid therapy, including opioid use disorder, overdose, and death. CDC is committed to evaluating the guideline to identify the impact of the recommendations on clinician and patient outcomes, both intended and unintended, and revising the recommendations in future updates when warranted.

Acknowledgments

Members of the Core Expert Group; the Core Expert Group facilitator: Don Teater, MD; members of the Stakeholder Review Group; peer reviewers; the Opioid Guideline Workgroup, consultants, and the NCIPC Board of Scientific Counselors; federal partners: Richard Kronick, PhD, Deborah G. Perfetto, PharmD, Agency for Healthcare Research and Quality; Jeffrey A. Kelman, MD, Diane L. McNally, Centers for Medicare & Medicaid Services; Jonathan Woodson, MD, David Smith, MD, Jack Smith, MD, Christopher Spevak, MD, Department of Defense; Stephen M. Ostroff, MD, Christopher M. Jones, PharmD, Food and Drug Administration; Jim Macrae, MA, MPP, Alexander F. Ross, ScD, Health Resources and Services Administration; Nora Volkow, MD, David Thomas, PhD, National Institute of Drug Abuse; John Howard, MD, Douglas Trout, MD, National Institute for Occupational Safety and Health; Karen B. DeSalvo, MD, Jennifer Frazier, MPH, Office of the National Coordinator, Michael Botticelli, MEd, Cecelia McNamara Spitznas, PhD, Office of National Drug Control Policy; Kana Enomoto, MA, Jinhee Lee, PharmD, Substance Abuse and Mental Health Services Administration; Robert McDonald, MBA, Jack M. Rosenberg, MD, Veterans Administration; members of the public who provided comment during the webinar; Douglas McDonald, PhD, Brandy Wyant, MPH, Kenneth Carlson, Amy Berninger, MPH, Abi Associates; Thomas Frieden, MD, Anne Schuchat, MD, Ileana Arias, PhD, CDC Office of the Director, Debra Houry, MD, National Center for Injury Prevention and Control, Amy Peples, MPA, National Center for Injury Prevention and Control, Arlene Greenspan, DrPH, National Center for Injury Prevention and Control, Grant Baldwin, PhD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Rita Noonan, PhD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Julie Gilchrist, MD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Terry Davis, EdD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Wes Sargent, EdD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Brian Manns, PharmD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Lisa Garbarino, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Donovan Newton, MPA, Division of Analysis, Research and Practice Integration, National Center for Injury Prevention and Control, Joann Kang, JD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Noah Aleshire, JD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Jennifer VanderVeur, JD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, LeShaundra Scott, MPH, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Sarah Lewis, MPH, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Helen Kingery, MPH, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Kristen Sanderson, MPH, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, Kate Fox, MPP, National Center for Injury Prevention and Control, Leslie Dorigo, MA, National Center for Injury Prevention and Control, Erin Connelly, MPA, National Center for Injury Prevention and Control, Sara Patterson, MA, National Center for Injury Prevention and Control, Mark Biagioni, MPA, National Center for Injury Prevention and Control, and Leonard J. Paulozzi, MD, Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, CDC.

References


Recommendations and Reports


<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of evidence</th>
<th>Other factors</th>
<th>Estimates of effect/findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Effectiveness and comparative effectiveness (KQ1)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Effectiveness of long-term opioid therapy versus placebo or no opioid therapy for long-term (≥1 year) outcomes</td>
<td>None</td>
<td>—†</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Pain, function, and quality of life</td>
<td>None</td>
<td>—†</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Harms and adverse events (KQ2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Risks of opioids versus placebo or no opioids on opioid abuse, addiction, and related outcomes; overdose; and other harms</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abuse or addiction</td>
<td>1 cohort study (n = 568,640)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>One retrospective cohort study found long-term use of prescribed opioids associated with an increased risk of abuse or dependence diagnosis versus no opioid use (adjusted OR ranged from 14.9 to 122.5, depending on dose).</td>
</tr>
<tr>
<td>Abuse or addiction</td>
<td>10 uncontrolled studies (n = 3,780)</td>
<td>Very serious limitations</td>
<td>Very serious inconsistency</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>In primary care settings, prevalence of opioid abuse ranged from 0.6% to 8% and prevalence of dependence from 3% to 26%. In pain clinic settings, prevalence of misuse ranged from 8% to 16% and addiction from 2% to 14%. Prevalence of aberrant drug-related behaviors ranged from 6% to 37%.</td>
</tr>
<tr>
<td>Overdose</td>
<td>1 cohort study (n = 9,940)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Current opioid use associated with increased risk of any overdose events (adjusted HR 5.2, 95% CI = 2.1–12) and serious overdose events (adjusted HR 8.4, 95% CI = 2.5–28) versus current nonuse.</td>
</tr>
<tr>
<td>Fractures</td>
<td>1 cohort study (n = 2,341) and 1 case–control study (n = 21,739 case patients)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Opioid use associated with increased risk of fracture in 1 cohort study (adjusted HR 1.28, 95% CI = 0.99–1.64) and 1 case-control study (adjusted OR 1.27, 95% CI = 1.21–1.33).</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>1 cohort study (n = 426,124) and 1 case–control study (n = 11,693 case patients)</td>
<td>No limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Current opioid use associated with increased risk of myocardial infarction versus nonuse (adjusted OR 1.28, 95% CI = 1.19–1.37 and incidence rate ratio 2.66, 95% CI = 2.30–3.08).</td>
</tr>
<tr>
<td>Endocrinologic harms</td>
<td>1 cross-sectional study (n = 11,327)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Long-term opioid use associated with increased risk for use of medications for erectile dysfunction or testosterone replacement versus nonuse (adjusted OR 1.5, 95% CI = 1.1–1.9).</td>
</tr>
<tr>
<td>How do harms vary depending on the opioid dose used?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abuse or addiction</td>
<td>1 cohort study (n = 568,640)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>One retrospective cohort study found higher doses of long-term opioid therapy associated with increased risk of opioid abuse or dependence than lower doses. Compared to no opioid prescription, the adjusted odds ratios were 15 (95% CI = 10–21) for 1 to 36 MME/day, 29 (95% CI = 20–41) for 36 to 120 MME/day, and 122 (95% CI = 73–205) for ≥120 MME/day.</td>
</tr>
<tr>
<td>Overdose</td>
<td>1 cohort study (n = 9,940) and 1 case–control study (n = 593 case patients in primary analysis)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>Magnitude of effect, dose response relationship</td>
<td>Versus 1 to &lt;20 MME/day, one cohort study found an adjusted HR for an overdose event of 1.44 (95% CI = 0.57–3.62) for 20 to &lt;50 MME/day that increased to 8.87 (95% CI = 3.99–19.72) at ≥100 MME/day; one case-control study found an adjusted OR for an opioid-related death of 1.32 (95% CI = 0.94–1.84) for 20 to 49 MME/day that increased to 2.88 (95% CI = 1.79–4.63) at ≥200 MME/day.</td>
</tr>
<tr>
<td>Fractures</td>
<td>1 cohort study (n = 2,341)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Risk of fracture increased from an adjusted HR of 1.20 (95% CI = 0.92–1.56) at 1 to &lt;20 MME/day to 2.00 (95% CI = 1.24–3.24) at ≥30 MME/day; the trend was of borderline statistical significance.</td>
</tr>
</tbody>
</table>

See table footnotes on page 47.
### TABLE 1. (Continued) Grading of Recommendations Assessment, Development and Evaluation (GRADE) clinical evidence review ratings of the evidence for the key clinical questions regarding effectiveness and risks of long-term opioid therapy for chronic pain

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of evidence</th>
<th>Other factors</th>
<th>Estimates of effect/findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myocardial infarction</td>
<td>1 cohort study (n = 426,124)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Relative to a cumulative dose of 0 to 1,350 MME during a 90-day period, the incidence rate ratio for myocardial infarction for 1350 to &lt;2700 MME was 1.21 (95% CI = 1.02–1.45), for 2,700 to &lt;8,100 MME was 1.42 (95% CI = 1.21–1.67), for 8,100 to &lt;18,000 MME was 1.89 (95% CI = 1.54–2.33), and for ≥18,000 MME was 1.73 (95% CI = 1.32–2.26).</td>
</tr>
<tr>
<td>Motor vehicle crash injuries</td>
<td>1 case–control study (n = 5,300 case patients)</td>
<td>No limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>No association between opioid dose and risk of motor vehicle crash injuries even though opioid doses &gt;20 MME/day were associated with increased odds of road trauma among drivers.</td>
</tr>
<tr>
<td>Endocrinologic harms</td>
<td>1 cross-sectional study (n = 11,327) New for update: 1 additional cross-sectional study (n = 1,585)</td>
<td>Serious limitations</td>
<td>Consistent</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Relative to 0 to &lt;20 MME/day, the adjusted OR for ≥120 MME/day for use of medications for erectile dysfunction or testosterone replacement was 1.6 (95% CI = 1.0–2.4). One new cross-sectional study found higher-dose long-term opioid therapy associated with increased risk of androgen deficiency among men receiving immediate-release opioids (adjusted OR per 10 MME/day 1.16, 95% CI = 1.09–1.23), but the dose response was very weak among men receiving ER/LA opioids.</td>
</tr>
<tr>
<td>Dosing strategies (KQ3)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Comparative effectiveness of different methods for initiating opioid therapy and titrating doses</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain</td>
<td>3 randomized trials (n = 93)</td>
<td>Serious limitations</td>
<td>Serious inconsistency</td>
<td>Very serious imprecision</td>
<td>4</td>
<td>None identified</td>
<td>Trials on effects of titration with immediate-release versus ER/LA opioids reported inconsistent results and had additional differences between treatment arms in dosing protocols (titrated versus fixed dosing) and doses of opioids used.</td>
</tr>
<tr>
<td>Overdose</td>
<td>New for update: 1 cohort study (n = 840,606)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One new cross-sectional study found initiation of therapy with an ER/LA opioid associated with increased risk of overdose versus initiation with an immediate-release opioid (adjusted HR 2.33, 95% CI = 1.26–4.32).</td>
</tr>
<tr>
<td><strong>Comparative effectiveness of different ER/LA opioids</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain and function</td>
<td>3 randomized trials (n = 1,850) 1 cohort study (n = 108,492) New for update: 1 cohort study (n = 38,756)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>No differences</td>
</tr>
<tr>
<td>All-cause mortality</td>
<td>New for update: 1 cohort study (n = 38,756)</td>
<td>Serious limitations</td>
<td>Serious inconsistency</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One cohort study found methadone to be associated with lower all-cause mortality risk than sustained-release morphine in a propensity-adjusted analysis (adjusted HR 0.56, 95% CI = 0.51–0.62) and one cohort study among Tennessee Medicaid patients found methadone to be associated with higher risk of all-cause mortality than sustained-release morphine (adjusted HR 1.46, 95% CI = 1.17–1.73).</td>
</tr>
<tr>
<td>Abuse and related outcomes</td>
<td>1 cohort study (n = 5,684)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Serious imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One cohort study found some differences between ER/LA opioids in rates of adverse outcomes related to abuse, but outcomes were nonspecific for opioid-related adverse events, precluding reliable conclusions.</td>
</tr>
<tr>
<td>ER/LA versus immediate-release opioids</td>
<td>New for update: 1 cross-sectional study (n = 1,585)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One cross-sectional study found ER/LA opioids associated with increased risk of androgen deficiency versus immediate-release opioids (adjusted OR 3.39, 95% CI = 2.39–4.77).</td>
</tr>
</tbody>
</table>

See table footnotes on page 47.
TABLE 1. (Continued) Grading of Recommendations Assessment, Development and Evaluation (GRADE) clinical evidence review ratings of the evidence for the key clinical questions regarding effectiveness and risks of long-term opioid therapy for chronic pain

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of evidence</th>
<th>Other factors</th>
<th>Estimates of effect/findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dose escalation versus dose maintenance or use of dose thresholds</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>No difference between more liberal dose escalation versus maintenance of current doses in pain, function, or risk of withdrawal due to opioid misuse, but there was limited separation in opioid doses between groups (52 versus 40 MME/day at the end of the trial).</td>
</tr>
<tr>
<td>Pain, function, or withdrawal due to opioid misuse</td>
<td>1 randomized trial</td>
<td>Serious</td>
<td>Unknown</td>
<td>Very serious</td>
<td>3</td>
<td>None identified</td>
<td></td>
</tr>
<tr>
<td>(n = 140)</td>
<td>(1 study)</td>
<td>limitations</td>
<td></td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immediate-release versus ER/LA opioids; immediate-release plus ER/LA opioids versus ER/LA opioids alone; scheduled and continuous versus as-needed dosing of opioids; or opioid rotation versus maintenance of current therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Abrupt cessation of morphine was associated with increased pain and decreased function compared with continuation of morphine.</td>
</tr>
<tr>
<td>Pain, function, quality of life, and outcomes related to abuse</td>
<td>None</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Effects of decreasing or tapering opioid doses versus continuation of opioid therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain and function</td>
<td>1 randomized trial</td>
<td>Very serious</td>
<td>Unknown</td>
<td>Very serious</td>
<td>4</td>
<td>None identified</td>
<td>Abrupt cessation of morphine was associated with increased pain and decreased function compared with continuation of morphine.</td>
</tr>
<tr>
<td>(n = 10)</td>
<td>(1 study)</td>
<td>limitations</td>
<td></td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Comparative effectiveness of different tapering protocols and strategies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>No clear differences between different methods for opioid discontinuation or tapering in likelihood of opioid abstinence after 3–6 months</td>
</tr>
<tr>
<td>Opioid abstinence</td>
<td>2 nonrandomized trials</td>
<td>Very serious</td>
<td>No inconsistency</td>
<td>Very serious</td>
<td>4</td>
<td>None identified</td>
<td>No clear differences between different methods for opioid discontinuation or tapering in likelihood of opioid abstinence after 3–6 months</td>
</tr>
<tr>
<td>(n = 150)</td>
<td>(n = 150)</td>
<td>limitations</td>
<td></td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Risk assessment and risk mitigation strategies (KQ4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diagnostic accuracy of instruments for predicting risk for opioid overdose, addiction, abuse, or misuse among patients with chronic pain being considered for long-term opioid therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Opioid risk tool</td>
<td>3 studies of diagnostic accuracy</td>
<td>Serious</td>
<td>Very serious</td>
<td>Serious</td>
<td>4</td>
<td>None identified</td>
<td>Based on a cutoff score of &gt;4 (or unspecified), five studies (two fair-quality, three poor-quality) reported sensitivity that ranged from 0.20 to 0.99 and specificity that ranged from 0.16 to 0.88.</td>
</tr>
<tr>
<td>(n = 496)</td>
<td>(n = 496)</td>
<td>limitations</td>
<td>inconsistency</td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Screener and Opioid Assessment for Patients with Pain, Version 1</td>
<td>New for update: 2 studies of diagnostic</td>
<td>Very serious</td>
<td>No inconsistency</td>
<td>Serious</td>
<td>3</td>
<td>None identified</td>
<td>Based on a cutoff score of &gt;3 or unspecified, sensitivity was 0.68 and specificity was 0.38 in one study, for a positive likelihood ratio of 1.11 and a negative likelihood ratio of 0.83. Based on a cutoff score of &gt;6, sensitivity was 0.73 in one study.</td>
</tr>
<tr>
<td>(n = 320)</td>
<td>accuracy (n = 320)</td>
<td>limitations</td>
<td></td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Screener and Opioid Assessment for Patients with Pain-Revised</td>
<td>New for update: 2 studies of diagnostic</td>
<td>Very serious</td>
<td>No inconsistency</td>
<td>Serious</td>
<td>3</td>
<td>None identified</td>
<td>Based on a cutoff score of &gt;3 or unspecified, sensitivity was 0.25 and 0.53 and specificity was 0.62 and 0.73 in two studies, for likelihood ratios close to 1. Based on a &quot;high risk&quot; assessment, sensitivity was 0.73 and 0.83 and specificity was 0.43 and 0.88 in two studies, for positive likelihood ratios of 1.28 and 7.18 and negative likelihood ratios of 0.63 and 0.19.</td>
</tr>
<tr>
<td>(n = 320)</td>
<td>accuracy (n = 320)</td>
<td>limitations</td>
<td></td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Brief Risk Interview</td>
<td>New for update: 2 studies of diagnostic</td>
<td>Very serious</td>
<td>No inconsistency</td>
<td>Serious</td>
<td>3</td>
<td>None identified</td>
<td>Based on a &quot;high risk&quot; assessment, sensitivity was 0.73 and 0.83 and specificity was 0.43 and 0.88 in two studies, for positive likelihood ratios of 1.28 and 7.18 and negative likelihood ratios of 0.63 and 0.19.</td>
</tr>
<tr>
<td>(n = 320)</td>
<td>accuracy (n = 320)</td>
<td>limitations</td>
<td></td>
<td>imprecision</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

See table footnotes on page 47.
TABLE 1. (Continued) Grading of Recommendations Assessment, Development and Evaluation (GRADE) clinical evidence review ratings of the evidence for the key clinical questions regarding effectiveness and risks of long-term opioid therapy for chronic pain

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of evidence</th>
<th>Other factors</th>
<th>Estimates of effect/findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effectiveness of risk prediction instruments on outcomes related to overdose, addiction, abuse, or misuse in patients with chronic pain</td>
<td>Outcomes related to abuse</td>
<td>None</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Effectiveness of risk mitigation strategies, including opioid management plans, patient education, urine drug screening, use of prescription drug monitoring program data, use of monitoring instruments, more frequent monitoring intervals, pill counts, and use of abuse-deterrent formulations, on outcomes related to overdose, addiction, abuse, or misuse</td>
<td>Outcomes related to abuse</td>
<td>None</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Effectiveness of risk prediction instruments on outcomes related to overdose, addiction, abuse, or misuse in patients with chronic pain</td>
<td>Outcomes related to abuse</td>
<td>None</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Effectiveness of risk mitigation strategies, including opioid management plans, patient education, urine drug screening, use of prescription drug monitoring program data, use of monitoring instruments, more frequent monitoring intervals, pill counts, and use of abuse-deterrent formulations, on outcomes related to overdose, addiction, abuse, or misuse</td>
<td>Outcomes related to abuse</td>
<td>None</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Comparative effectiveness of treatment strategies for managing patients with addiction to prescription opioids</td>
<td>Outcomes related to abuse</td>
<td>None</td>
<td>—</td>
<td>—</td>
<td>Insufficient</td>
<td>—</td>
<td>No evidence</td>
</tr>
<tr>
<td>Effects of opioid therapy for acute pain on long-term use (KQ5)</td>
<td>Long-term opioid use</td>
<td>New for update: 2 cohort studies (n = 399,852)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
</tr>
</tbody>
</table>

Abbreviations: CI = confidence interval; ER/LA = extended release/long-acting; HR = hazard ratio; MME = morphine milligram equivalents; OR = odds ratio.

* Ratings were made per GRADE quality assessment criteria; “no limitations” indicates that limitations assessed through the GRADE method were not identified.
† Not applicable as no evidence was available for rating.
TABLE 2. Morphine milligram equivalent (MME) doses for commonly prescribed opioids

<table>
<thead>
<tr>
<th>Opioid</th>
<th>Conversion factor*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Codeine</td>
<td>0.15</td>
</tr>
<tr>
<td>Fentanyl transdermal (in mcg/hr)</td>
<td>2.4</td>
</tr>
<tr>
<td>Hydrocodone</td>
<td>1</td>
</tr>
<tr>
<td>Hydromorphone</td>
<td>4</td>
</tr>
<tr>
<td>Methadone 1–20 mg/day</td>
<td>4</td>
</tr>
<tr>
<td>Methadone 21–40 mg/day</td>
<td>8</td>
</tr>
<tr>
<td>Methadone 41–60 mg/day</td>
<td>10</td>
</tr>
<tr>
<td>Methadone ≥61–80 mg/day</td>
<td>12</td>
</tr>
<tr>
<td>Methadone 1–20 mg/day</td>
<td>4</td>
</tr>
<tr>
<td>Morphine</td>
<td>1</td>
</tr>
<tr>
<td>Oxycodone</td>
<td>1.5</td>
</tr>
<tr>
<td>Oxymorphone</td>
<td>3</td>
</tr>
<tr>
<td>Tapentadol†</td>
<td>0.4</td>
</tr>
</tbody>
</table>

* Multiply the dose for each opioid by the conversion factor to determine the dose in MMEs. For example, tablets containing hydrocodone 5 mg and acetaminophen 300 mg taken four times a day would contain a total of 20 mg of hydrocodone daily, equivalent to 20 MME daily; extended-release tablets containing oxycodone 10 mg and taken twice a day would contain a total of 20 mg of oxycodone daily, equivalent to 30 MME daily. The following cautions should be noted: 1) All doses are in mg/day except for fentanyl, which is mcg/hr. 2) Equianalgesic dose conversions are only estimates and cannot account for individual variability in genetics and pharmacokinetics. 3) Do not use the calculated dose in MMEs to determine the doses to use when converting opioid to another; when converting opioids the new opioid is typically dosed at substantially lower than the calculated MME dose to avoid accidental overdose due to incomplete cross-tolerance and individual variability in opioid pharmacokinetics. 4) Use particular caution with methadone dose conversions because the conversion factor increases at higher doses. 5) Use particular caution with fentanyl since it is dosed in mcg/hr instead of mg/day, and its absorption is affected by heat and other factors.

† Tapentadol is a mu receptor agonist and norepinephrine reuptake inhibitor. MMEs are based on degree of mu-receptor agonist activity, but it is unknown if this drug is associated with overdose in the same dose-dependent manner as observed with medications that are solely mu receptor agonists.

Steering Committee and Core Expert Group Members

**Steering Committee:** Deborah Dowell, MD, Tamara M. Haegerich, PhD; Division of Unintentional Injury Prevention, National Center for Injury Prevention and Control, CDC; Roger Chou, MD; on detail to CDC under contract.

**Core Expert Group Members:** Pam Archer, MPH, Oklahoma State Department of Health; Jane Ballantyne, MD; University of Washington (retired); Amy Bohnert, PhD; University of Michigan; Bonnie Burman, ScD; Ohio Department on Aging; Roger Chou, MD; on detail to CDC under contract; Phillip Coffin, MD, San Francisco Department of Public Health; Gary Franklin, MPH, Washington State Department of Labor and Industries/University of Washington; Erin Krebs, MDH; Minneapolis VA Health Care System/University of Minnesota; Michel Mutter, MD, Tennessee Department of Health; Lewis Nelson, MD; New York University School of Medicine; Trupti Patel, MD; Arizona Department of Health Services; Christina A. Porucznik, PhD, University of Utah; Robert “Chuck” Rich, MD, FAAFP, American Academy of Family Physicians; Joanna Starrels, MD, Albert Einstein College of Medicine of Yeshiva University; Michael Steinman, MD, Society of General Internal Medicine; Thomas Tape, MD, American College of Physicians; Judith Turner, PhD, University of Washington.

Stakeholder Review Group

John Markman, MD, American Academy of Neurology; Bob Tuillman, PhD, American Academy of Pain Management; Edward C. Covington, MD, American Academy of Pain Medicine; Roger F. Suchyta, MD, FAAP, American Academy of Pediatrics; Kavitha V. Neerukonda, JD, American Academy of Physical Medicine and Rehabilitation; Mark Fleury, PhD, American Cancer Society Cancer Action Network; Penney Cowan, American Chronic Pain Association; David Juurlink, BPharm, MD, PhD, American College of Medical Toxicology; Gerald “Jerry” E Joseph, Jr, MD, American College of Obstetrics and Gynecology; Bruce Ferrell, MD, AGSF; M. Carrington Reid, MD, PhD, American Geriatrics Society; Ashley Thompson, American Hospital Association; Barry D. Dickinson, PhD, American Medical Association; Gregory Terman MD, PhD, American Pain Society; Beth Haynes, MPPA, American Society of Addiction Medicine; Asokumar Buvanendran, MD, American Society of Anesthesiologists; Robert M. Plovnick; MD, American Society of Hematology; Sanford M. Silverman, MD, American Society of Interventional Pain Physicians; Andrew Kolodny, MD, Physicians for Responsible Opioid Prescribing.

Opioid Guideline Workgroup

**Chair:** Christina Porucznik, PhD, MSPH

**Workgroup Members:** Anne Burns, RPh; Penney Cowan; Chinazo Cunningham, MD, MS; Katherine Galluzzi, DO; Traci Green, PhD, MSC; Mitchell Katz, MD; Erin Krebs, MD, MPH; Gregory Terman, MD, PhD; Mark Wallace, MD. **Workgroup Consultants:** Roger Chou, MD; Edward Covington, MD; Diana Eppolito; Michael Greene, MD; Steven Stason, DO.

Peer Reviewers

Jeanmarie Perrone, MD, University of Pennsylvania; Matthew Bair, MD, Indiana University School of Medicine; David Tauben, MD, University of Washington.

NCIPC Board of Scientific Counselors

Chair: Stephen Hargarten, MD, MPH; Members: John Allegriante, PhD; Joan Marie Duwee, MD, Samuel Forjuoh, MD, MPH, DrPH, FGCP; Gerard Gioia, PhD; Deborah Gorman-Smith, PhD; Traci Green, PhD; Sherry Lynne Hanby, PhD; Robert Johnson, MD; Angela Mickalide, PhD, MCHES; Sherry Molock, PhD; Christina Porucznik, PhD, MSPH; Jay Silverman, PhD; Maria Tesa, PhD; Shelly Timmons, MD, PhD, FACS, FAANS; Ex Officio Members: Melissa Brodowski, PhD; Dawn Castillo, MPH; Wilson Compton, MD, MPE; Elizabeth Edgerton, MD, MPH; Thomas Feucht, PhD; Meredith Fox, PhD; Holly Hedegaard, MD, MSPH; John Howard, MD; Lyndon Joseph, PhD; Jinho Lee, PharmD; Iris Malby-Hernandez, MD, MPH; Valeri Maholmes, PhD; Angela Moore Parmley, PhD; Thomas Schroeder, MS.
In the report, “CDC Guideline for Prescribing Opioids for Chronic Pain — United States, 2016,” three errors occurred. On page 1, the last sentence of the Summary should read, “CDC has provided a checklist for prescribing opioids for chronic pain (http://stacks.cdc.gov/view/cdc/38025) as well as a website (http://www.cdc.gov/drugoverdose/prescribing/resources.html) with additional tools to guide clinicians in implementing the recommendations.” On page 8, the first sentence of the first full paragraph should read, “NCIPC announced an open meeting of the NCIPC BSC in the Federal Register on January 11, 2016.” On page 49, in the fourth line of the Stakeholder Review Group, the affiliation for Gerald “Jerry” F. Joseph should read, “American College of Obstetricians and Gynecologists.”


Use of trade names and commercial sources is for identification only and does not imply endorsement by the U.S. Department of Health and Human Services.

References to non-CDC sites on the Internet are provided as a service to MMWR readers and do not constitute or imply endorsement of these organizations or their programs by CDC or the U.S. Department of Health and Human Services. CDC is not responsible for the content of pages found at these sites. URL addresses listed in MMWR were current as of the date of publication.

All HTML versions of MMWR articles are generated from final proofs through an automated process. This conversion might result in character translation or format errors in the HTML version. Users are referred to the electronic PDF.
version ([https://www.cdc.gov/mmwr](https://www.cdc.gov/mmwr)) and/or the original MMWR paper copy for printable versions of official text, figures, and tables.

Questions or messages regarding errors in formatting should be addressed to mmwrq@cdc.gov.
Guideline Information for Patients

Safer, More Effective Pain Management

Living with chronic pain can be challenging. It is essential that you and your doctor discuss treatment options with all of the risks and benefits carefully considered. Some medications, such as prescription opioids, can help relieve pain in the short term but also come with serious risks and potential complications—and must be prescribed and used carefully.

The new CDC Guideline for Prescribing Opioids for Chronic Pain(https://www.cdc.gov/drugoverdose/prescribing/guideline.html) helps inform providers’ ability to offer safer, more effective pain management and supports clinical decision making about prescribing opioids.

What are opioids?

Opioids are natural or synthetic chemicals that reduce feelings of pain. Common prescription opioid pain relievers include:

- Hydrocodone (Vicodin)
- Oxycodone (OxyContin)
- Oxymorphone (Opana)
- Methadone
- Fentanyl
Are opioids safe?

Prescription opioids can help with some types of pain in the short term but have serious risks. They can be an important part of treatment in some circumstances and can effectively relieve suffering for patients with active cancer or others in hospice or palliative care, but studies are not available to indicate whether opioids control chronic pain well when used long-term. Before taking opioid medication for your chronic pain:

- Discuss pain treatment options, including ones that do not involve prescription drugs.
- Tell your doctor about past or current drug and alcohol use.
- Discuss all of the risks and benefits of taking prescription opioids.

What are the risks from opioids?

Patients taking prescription opioids are at risk for unintentional overdose or death and can become addicted. From 1999 to 2014, more than 165,000 persons died from overdose related to prescription opioids in the United States. Up to 1 out of 4 people receiving long-term opioid therapy in a primary care setting struggles with addiction. In addition to the serious risks of addiction and overdose, the use of prescription opioid pain relievers can have a number of side effects, even when taken as directed:

- Tolerance—meaning you might need to take more of the medication for the same pain relief
- Physical dependence—meaning you have symptoms of withdrawal when the medication is stopped
- Increased sensitivity to pain
- Constipation
- Nausea, vomiting, and dry mouth
- Sleepiness and dizziness
- Confusion
- Depression
- Low levels of testosterone that can result in lower sex drive, energy, and strength
- Itching and sweating

Remember, your doctor is a partner in your pain treatment plan. It’s important to talk about any and all side effects and concerns to make sure you’re getting the safest and most effective care.
What is the new opioid prescribing guideline and how will it affect me?

CDC developed the new Guideline for Prescribing Opioids for Chronic Pain to help primary care providers make informed prescribing decisions and improve patient care for those who suffer from chronic pain (pain lasting more than 3 months) in outpatient settings. The guideline is not intended for patients who are in active cancer treatment, palliative care, or end-of-life care.

If you have chronic pain and are prescribed opioids as part of your treatment, your doctor should monitor you regularly. This might include extra assessments, a pain treatment plan, more frequent office visits, and urine testing. Prescription opioids can be very dangerous if not used properly. Make sure to follow all of your doctor’s recommendations.

If you are prescribed opioids

- Use them only as instructed by your doctor. Never take opioids in greater amounts or more often than prescribed.
- Avoid these other drugs while taking this medication:
  - Alcohol
  - Benzodiazepines (such as Xanax and Valium), unless specifically advised by your doctor
  - Muscle relaxants (such as Soma or Flexeril), unless specifically advised by your doctor
  - Hypnotics (such as Ambien or Lunesta), unless specifically advised by your doctor
  - Other prescription opioid pain relievers
- Work with your doctor to create a plan on how to manage your pain, and consider non-opioid options.
- Follow up regularly with your doctor.
- Talk to your doctor about any and all side effects and concerns.
- Store opioid pain relievers in a safe place and out of reach of others.
  - Help prevent misuse and abuse by not selling or sharing prescription opioid pain relievers. Never use another person’s prescription opioids.
  - Find your community drug take-back program or your pharmacy mail-back program to safely dispose of unused prescription opioids pain relievers.

Know your options

Talk to your doctor about ways to manage your pain that don’t involve prescription opioids. Some of these options may actually work better and have fewer risks and side effects. Options may include:

- Acetaminophen (Tylenol®) or ibuprofen (Advil®)
- Cognitive behavioral therapy
- Physical therapy and exercise
- Medications for depression or for seizures
- Interventional therapies (injections)
Where can I get help?

If you or someone close to you needs help for substance abuse problems, talk to your doctor or call SAMHSA’s National Helpline at 1-800-662-HELP or go to SAMHSA’s [SAMHSA’s Behavioral Health Treatment Services Locator](https://www.samhsa.gov/find-treatment). If you have questions about any medicines, call the U.S. Department of Health and Human Services Poison Help Hotline at 1-800-222-1222.

Tools and resources for patients:

- [Pregnancy and Opioid Pain Medications](https://www.cdc.gov/drugoverdose/pdf/pregnancy_opioid_pain_factsheet-a.pdf) [PDF - 1 MB]
  Women who take opioid pain relievers should be aware of the possible risks during pregnancy.


Guideline Information for Providers

Safe Prescribing Saves Lives

Chronic pain is common, multidimensional, and individualized, and treatment can be challenging for healthcare providers as well as patients. In response to the critical need for consistent and current opioid prescribing guidelines, the CDC released the new Guideline for Prescribing Opioids for Chronic Pain(https://www.cdc.gov/drugoverdose/prescribing/guideline.html).

Since 1999, opioid prescriptions have quadrupled, and over 183,000 people have died from prescription opioids.¹ ² These new recommendations focus on clinical practice and provide evidence and guidance to improve how these drugs are prescribed—and ultimately improve patient care.

What is the purpose of the new guideline?

The guideline helps providers make informed decisions about pain treatment for patients 18 and older in primary care settings. The recommendations focus on the use of opioids in treating chronic pain—pain lasting longer than three months or past the time of normal tissue healing. The guideline is not intended for patients who are in active cancer treatment, palliative care, or end-of-life care.

Opioids pose a risk to all patients. The guideline encourages providers to implement best practices for responsible prescribing.

Use nonopioid therapies

Use nonpharmacologic therapies (such as exercise and cognitive behavioral therapy) and nonopioid pharmacologic therapies (such as anti-inflammatories) for chronic pain. Don't use opioids routinely for chronic pain. When opioids are used, combine them with nonpharmacologic or nonopioid pharmacologic therapy, as appropriate, to provide greater benefits.

Start low and go slow

When opioids are used, prescribe the lowest possible effective dosage and start with immediate-release opioids instead of extended-release/long-acting opioids. Only provide the quantity needed for the expected duration of pain.
Follow-up

Regularly monitor patients to make sure opioids are improving pain and function without causing harm. If benefits do not outweigh harms, optimize other therapies and work with patients to taper or reduce dosage and discontinue, if needed.

What’s included in the guideline?

The guideline addresses patient-centered clinical practices including conducting thorough assessments, considering all possible treatments, closely monitoring risks, and safely discontinuing opioids. The three main focus areas in the guideline include:

1. Determining when to initiate or continue opioids for chronic pain
   - Selection of non-pharmacologic therapy, non-opioid pharmacologic therapy, opioid therapy
   - Establishment of treatment goals
   - Discussion of risks and benefits of therapy with patients

2. Opioid selection, dosage, duration, follow-up and discontinuation
   - Selection of immediate-release or extended-release and long-acting opioids
   - Dosage considerations
   - Duration of treatment
   - Considerations for follow-up and discontinuation of opioid therapy

3. Assessing risk and addressing harms of opioid use
   - Evaluation of risk factors for opioid-related harms and ways to mitigate/reduce patient risk
   - Review of prescription drug monitoring program (PDMP) data
   - Use of urine drug testing
   - Considerations for co-prescribing benzodiazepines
   - Arrangement of treatment for opioid use disorder
What’s new in the CDC Guideline?

Dosage Recommendations

The dosage recommendations for exercising caution are lower than older opioid prescribing guidelines. Higher doses of opioids are associated with higher risk of overdose and death—even relatively low doses (20-50 morphine milligram equivalents (MME) per day) increase risk.

Assessing Risks and Harms

Previous guidelines focused safety precautions on “high risk patients,” however, opioids pose risk to all patients, and currently available tools cannot rule out risk for abuse or other serious harm. The CDC guideline provides recommendations on providing safer care for all patients. The guideline also encourages use of recent technological advances, such as state prescription drug monitoring programs.

Monitoring and Discontinuing

The guideline provides more specific recommendations compared to previous guidelines on monitoring and discontinuing opioids when risks and harms outweigh benefits.
What else is CDC doing?

The new prescribing guideline is just one of the strategies to reduce the number of people who suffer from opioid use disorder or overdose related to these drugs. Other efforts include:

- Enhancing and maximizing the use of PDMPs
- Helping states scale up effective programs through the Prevention for States program
- Conducting policy evaluations
- Developing and implementing Rapid Response Projects
- Improving data quality and tracking trends to monitor the epidemic
WHY GUIDELINES FOR PRIMARY CARE PROVIDERS?

Primary care providers account for approximately 50% of prescription opioids dispensed.

Nearly 2 million Americans, aged 12 or older, either abused or were dependent on prescription opioids in 2014.

- An estimated 11% of adults experience daily pain.
- Millions of Americans are treated with prescription opioids for chronic pain.
- Primary care providers are concerned about patient addiction and report insufficient training in prescribing opioids.

MYTH VS TRUTH

1. Opioids are effective long-term treatments for chronic pain.
   - While evidence supports short-term effectiveness of opioids, there is insufficient evidence that opioids control chronic pain effectively over the long term, and there is evidence that other treatments can be effective with less harm.

2. There is no unsafe dose of opioids as long as opioids are titrated slowly.
   - Daily opioid dosages close to or greater than 90 MME/day are associated with significant risks, and lower dosages are safer.

3. The risk of addiction is minimal.
   - Up to one quarter of patients receiving prescription opioids long-term in a primary care setting struggle with addiction. Certain risk factors increase susceptibility to opioid-associated harms: history of overdose, history of substance use disorder, higher opioid dosages, or concurrent benzodiazepine use.

WHAT CAN PROVIDERS DO?

First, do no harm. Long-term opioid use has uncertain benefits but known, serious risks. CDC’s Guideline for Prescribing Opioids for Chronic Pain will support informed clinical decision making, improved communication between patients and providers, and appropriate prescribing.

PRACTICES AND ACTIONS

- **USE NONOPIOID TREATMENT**
  - Opioids are not first-line or routine therapy for chronic pain (Recommendation #1)
  - In a systematic review, opioids did not differ from nonopioid medication in pain reduction, and nonopioid medications were better tolerated, with greater improvements in physical function.

- **REVIEW PDMP**
  - Check prescription drug monitoring program data for high dosages and prescriptions from other providers (Recommendation #9)
  - A study showed patients with one or more risk factors (4 or more prescribers, 4 or more pharmacies, or dosage >100 MME/day) accounted for 55% of all overdose deaths.

- **OFFER TREATMENT FOR OPIOID USE DISORDER**
  - Offer or arrange evidence-based treatment (e.g. medication-assisted treatment and behavioral therapies) for patients with opioid use disorder (Recommendation #12)
  - A study showed patients prescribed high dosages of opioids long-term (>90 days) had 122 times the risk of opioid use disorder compared to patients not prescribed opioids.

- **START LOW AND GO SLOW**
  - When opioids are started, prescribe them at the lowest effective dose (Recommendation #5)
  - Studies show that high dosages (>100 MME/day) are associated with 2 to 9 times the risk of overdose compared to <20 MME/day.

- **AVOID CONCURRENT PRESCRIBING**
  - Avoid prescribing opioids and benzodiazepines concurrently whenever possible (Recommendation #11)
  - One study found concurrent prescribing to be associated with a near quadrupling of risk for overdose death compared with opioid prescription alone.

LEARN MORE | www.cdc.gov/drugoverdose/prescribing/guideline.html
CDC Guideline for Prescribing Opioids for Chronic Pain—United States, 2016

Deborah Dowell, MD, MPH; Tamara M. Haegerich, PhD; Roger Chou, MD

IMPORTANT PRIMARY CARE CLINICIANS FIND MANAGING CHRONIC PAIN CHALLENGING. EVIDENCE OF LONG-TERM EFICACY OF OPIOIDS FOR CHRONIC PAIN IS LIMITED. OPIOID USE IS ASSOCIATED WITH SERIOUS RISKS, INCLUDING OPIOID USE DISORDER AND OVERDOSE.

OBJECTIVE TO PROVIDE RECOMMENDATIONS ABOUT OPIOID PRESCRIBING FOR PRIMARY CARE CLINICIANS TREATING ADULT PATIENTS WITH CHRONIC PAIN OUTSIDE OF ACTIVE CANCER TREATMENT, PALLIATIVE CARE, AND END-OF-LIFE CARE.

PROCESS THE CENTERS FOR DISEASE CONTROL AND PREVENTION (CDC) UPDATED A 2014 SYSTEMATIC REVIEW ON EFFECTIVENESS AND RISKS OF OPIOIDS AND CONDUCTED A SUPPLEMENTAL REVIEW ON BENEFITS AND HARMs, VALUES AND PREFERENCES, AND COSTS. CDC USED THE GRADING OF RECOMMENDATIONS ASSESSMENT, DEVELOPMENT, AND EVALUATION (GRADE) FRAMEWORK TO ASSESS EVIDENCE TYPE AND DETERMINE THE RECOMMENDATION CATEGORY.

EVIDENCE SYNTHESIS EVIDENCE CONSISTED OF OBSERVATIONAL STUDIES OR RANDOMIZED CLINICAL TRIALS WITH NOTABLE LIMITATIONS, CHARACTERIZED AS LOW QUALITY USING GRADE METHODOLOGY. META-ANALYSIS WAS NOT ATTEMPTED DUE TO THE LIMITED NUMBER OF STUDIES, VARIABILITY IN STUDY DESIGNS AND CLINICAL HETEROGENEITY, AND METHODOLOGICAL SHORTCOMINGS OF STUDIES. NO STUDY EVALUATED LONG-TERM (>1 YEAR) BENEFIT OF OPIOIDS FOR CHRONIC PAIN. OPIOIDS WERE ASSOCIATED WITH INCREASED RISKS, INCLUDING OPIOID USE DISORDER, OVERDOSE, AND DEATH, WITH DOSE-DEPENDENT EFFECTS.

RECOMMENDATIONS THERE ARE 12 RECOMMENDATIONS. OF PRIMARY IMPORTANCE, NONOPIOID THERAPY IS PREFERRED FOR TREATMENT OF CHRONIC PAIN. OPIOIDS SHOULD BE USED ONLY WHEN BENEFITS FOR PAIN AND FUNCTION ARE EXPECTED TO OWEIGH RISKS. BEFORE STARTING OPIOIDS, CLINICIANS SHOULD ESTABLISH TREATMENT GOALS WITH PATIENTS AND CONSIDER HOW OPIOIDS WILL BE DISCONTINUED IF BENEFITS DO NOT OWEIGH RISKS. WHEN OPIOIDS ARE USED, CLINICIANS SHOULD PRESCRIBE THE LOWEST EFFECTIVE DOSAGE, CAREFULLY REASSESS BENEFITS AND RISKS WHEN CONSIDERING INCREASING DOSAGE TO 50 MORPHINE MILLIGRAM EQUIVALENTS OR MORE PER DAY, AND AVOID CONCURRENT OPIOIDS AND BENZODIAZEPINES WHENEVER POSSIBLE. CLINICIANS SHOULD EVALUATE BENEFITS AND HARMs OF CONTINUED OPIOID THERAPY WITH PATIENTS EVERY 3 MONTHS OR MORE FREQUENTLY AND REVIEW PRESCRIPTION DRUG MONITORING PROGRAM DATA, WHEN AVAILABLE, FOR HIGH-RISK COMBINATIONS OR DOSAGES. FOR PATIENTS WITH OPIOID USE DISORDER, CLINICIANS SHOULD OFFER OR ARRANGE EVIDENCE-BASED TREATMENT, SUCH AS MEDICATION-ASSISTED TREATMENT WITH BUPRENORPHINE OR METHADONE.

CONCLUSIONS AND RELEVANCE THE GUIDELINE IS INTENDED TO IMPROVE COMMUNICATION ABOUT BENEFITS AND RISKS OF OPIOIDS FOR CHRONIC PAIN, IMPROVE SAFETY AND EFFECTIVENESS OF PAIN TREATMENT, AND REDUCE RISKS ASSOCIATED WITH LONG-TERM OPIOID THERAPY.

Published online March 15, 2016.
T
he number of people experiencing chronic pain is substantial, with US prevalence estimated at 11.2% of the adult population. Patients should receive appropriate pain treatment based on a careful consideration of the benefits and risks of treatment options. Opioids are commonly prescribed for pain, with approximately 3% to 4% of the adult US population prescribed long-term opioid therapy. Evidence supports short-term efficacy of opioids in randomized clinical trials lasting primarily 12 weeks or less, and patients receiving opioid therapy for chronic pain report some pain relief when surveyed. However, few studies have been conducted to rigorously assess the long-term benefits of opioids for chronic pain (pain lasting >3 months) with outcomes examined at least 1 year later. Opioid pain medication uses present serious risks. From 1999 to 2014, more than 165,000 persons died of overdose related to opioid pain medication in the United States. In 2013 alone, an estimated 1.9 million persons abused or were dependent on prescription opioid pain medication. Primary care clinicians report concern about opioid pain medication misuse, find managing patients with chronic pain stressful, express concern about patient addiction, and report insufficient training in prescribing opioids. The "CDC Guideline for Prescribing Opioids for Chronic Pain—United States, 2016," is intended for primary care clinicians (eg, family physicians, internists, nurse practitioners, and physician assistants) who are treating patients with chronic pain (ie, pain conditions that typically last >3 months or past the time of normal tissue healing) in outpatient settings. The guideline is intended to apply to patients 18 years and older with chronic pain outside of active cancer treatment, palliative care, and end-of-life care. Some of the recommendations might be relevant for acute care settings or other specialists, such as emergency physicians or dentists, but use in these settings or by other specialists is not the focus of the guideline.

The guideline is intended to improve communication between clinicians and patients about the risks and benefits of opioid therapy for chronic pain, improve the safety and effectiveness of pain treatment, and reduce the risks associated with long-term opioid therapy, including opioid use disorder, overdose, and death. Clinical decision making should be based on a relationship between the clinician and patient and an understanding of the patient’s clinical situation, functioning, and life context. The recommendations in the guideline are voluntary, rather than prescriptive standards. They are based on emerging evidence, including observational studies or randomized clinical trials with notable limitations. Clinicians should consider the circumstances and unique needs of each patient when providing care. This Special Communication details evidence reviewed by and official recommendations issued by the Centers for Disease Control and Prevention (CDC) and provides key highlights from a more extensive guideline; the full guideline with detailed information on disclosures and conflict of interest protocols, methods, scientific findings, and recommendation rationales can be found in the Morbidity and Mortality Weekly Report (MMWR).

Guideline Development Process

Grading of Recommendations Assessment, Development, and Evaluation Method

CDC used the CDC Advisory Committee on Immunization Practices (ACIP) translation of the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) method for guideline development. Within the ACIP GRADE framework, the quality of a body of evidence was graded, and the recommendations were developed and placed into categories (A or B) based on the quality of evidence, balance of benefits and harms, values and preferences, and resource allocation (Box 1).

CDC obtained input from experts, stakeholders, the public, peer reviewers, and a federally chartered advisory committee in the development process. CDC drafted a set of recommendations and invited subject matter experts, primary care professional society representatives, and state agency representatives (Core Expert Group, listed at the end of the article) to provide individual perspectives on how CDC used the evidence to develop the recommendations. CDC asked experts to undergo a rigorous process to assess and manage possible conflicts of interest; full details on protocols and disclosures are reported in the MMWR. CDC also engaged partners from 10 federal agencies and a Stakeholder Review Group of 18 organizations (listed at the end of the article) to provide comment. CDC convened a constituent engagement webinar to obtain additional perspectives from constituents on the key recommendations. To obtain comments from the public on the full guideline, CDC published a notice in the Federal Register (80 FR 77351) announcing the availability of the guideline and the supporting clinical and contextual evidence reviews for public comment. Per the final information quality bulletin for peer review (https://www.whitehouse.gov/sites/default/files/omb/memoranda/fy2005/m05-03.pdf), the guideline was peer reviewed because it provides influential scientific...
information. In addition, the National Center for Injury Prevention and Control Board of Scientific Counselors (BSC), a federal advisory committee, established an Opioid Guideline Workgroup (OGW) to review the guideline (members of the BSC and OGW are listed at the end of the article). The OGW issued a report of observations to the BSC. At an in-person meeting, the BSC considered the OGW report, deliberated on the draft guideline itself, and offered an additional opportunity for public comment. The BSC voted unanimously to support the observations made by the OGW; that CDC adopt the guideline recommendations that, according to the workgroup’s report, had unanimous or majority support; and that CDC further consider the guideline recommendations for which the group had mixed opinions. At each stage, CDC reviewed and carefully considered comments and revised the guideline.

**Clinical Evidence Review**

To inform the guideline development process, CDC updated a systematic review sponsored by the Agency for Healthcare Research and Quality (AHRQ) on the effectiveness and risks of long-term opioid treatment of chronic pain. This addressed clinical questions about effectiveness of long-term opioid therapy for outcomes at least 1 year later related to pain, function, and quality of life. The effectiveness of short-term opioid therapy has been established previously. In randomized clinical trials 12 weeks or shorter in duration, opioids were moderately effective for pain relief, with small benefits for functional outcomes; although estimates varied, based on uncontrolled studies, a high percentage of patients discontinued long-term opioid use because of lack of efficacy and because of adverse events. Opioids have unique effects such as tolerance and physical dependence that might influence assessments of benefit over time. These effects raise questions about whether findings on short-term effectiveness of opioid therapy can be extrapolated to estimate benefits of long-term therapy for chronic pain. Thus, it is important to consider studies that provide data on long-term benefit. For opioid-related harms (overdose, fractures, falls, motor vehicle crashes), studies were included with outcomes measured at shorter intervals because such outcomes can occur early during opioid therapy.

The review also considered evidence related to initiation and titration, harms and adverse events, and risk mitigation. CDC updated the review with more recent studies. Because long-term opioid use may be affected by use of opioids for acute pain, CDC added a clinical question on the effects of prescribing opioids for acute pain on long-term use (Box 2).

CDC updated the systematic literature search using search terms for opioid therapy, specific opioids, chronic pain, and comparative study designs; assessed the overall strength of each body of evidence using methods developed by the GRADE Working Group; and qualitatively synthesized results. Complete methods and data for the clinical evidence review, including information about data sources and searches, study selection, data extraction and quality assessment, data synthesis, and update search yield and new evidence may be found in the MMWR and associated online appendixes.

The updated review revealed that evidence on long-term opioid therapy for chronic pain outside of end-of-life care remains limited, with insufficient evidence to determine long-term benefits, although evidence suggests risk of serious harms that is dose-dependent. Table 1 provides a summary of the evidence and the quality ratings assigned. Full details on methodology and findings are available in the 2014 AHRQ report and the MMWR report. The body of evidence for each clinical question was categorized as evidence type 3 or 4 (observational studies or randomized clinical trials with notable limitations or clinical experience and observation). We highlight important findings from the review for each key question (KQ) below.

**KQ1: Effectiveness and Comparative Effectiveness**

No study of opioid therapy vs placebo, no opioid therapy, or non-opioid therapy for chronic pain evaluated long-term (>1 year) outcomes related to pain, function, or quality of life. Most placebo-controlled randomized clinical trials were 6 weeks or shorter in duration.

**KQ2: Harms and Adverse Events**

Long-term opioid therapy was associated with problematic patterns of opioid use leading to clinically significant impairment or distress. Varying terminology has been used to reflect this pattern, including “addiction” (more informally), “opioid abuse and opioid dependence” (per Diagnosis and Statistical Manual of Mental Disorders [Fourth Edition] [DSM-IV] or International Classification of Diseases, Ninth Revision, Clinical Modification [ICD-9-CM]), and “opioid use disorder” (per DSM-5). Such disorders are manifested by similar criteria, including unsuccessful efforts to reduce or control use and use resulting in social problems and a failure to fulfill major role obligations at work, school, or home. Disorders are different from tolerance (diminished response to a drug with repeated use) and physical dependence (adaptation to a drug that produces symptoms of withdrawal when the drug is stopped), both of which can exist without a diagnosed disorder.

Long-term opioid therapy was associated with an increased risk of an opioid abuse or dependence diagnosis (as defined by ICD-9-CM codes) vs no opioid prescription. In primary care settings, prevalence of opioid dependence (using DSM-IV criteria) ranged from 3% to 26%. Factors associated with increased risk of misuse included history of substance use disorder, younger age, major depression, and use of psychotropic medications. Opioid use was associated with a dose-dependent increased risk of fatal and nonfatal overdose and road trauma.

**KQ3: Dosing Strategies**

Initiation of therapy with an extended-release/long-acting (ER/LA) opioid was associated with greater risk of nonfatal overdose than initiation with an immediate-release opioid in 1 study, with risk greatest in the first 2 weeks after initiation of treatment. Three studies of various ER/LA opioids found no clear differences related to pain or function; there were mixed findings regarding the differences between methadone and morphine in overall risk for nonfatal or fatal overdose, suggesting that risks of methadone might vary in different settings. One study found no differences between more liberal dose escalation and maintenance of current doses after 12 months; evidence on other comparisons related to opioid dosing strategies was too limited to determine effects on outcomes.

**KQ4: Risk Assessment and Risk Mitigation Strategies**

Evidence on the accuracy of risk assessment instruments for predicting opioid abuse or misuse was inconsistent for the Opioid Risk
Box 2. Key Questions for the Clinical Evidence Review

Key Question 1. Effectiveness and Comparative Effectiveness
   a. In patients with chronic pain, what is the effectiveness of long-term opioid therapy vs placebo or no opioid therapy for long-term (>1 year) outcomes related to pain, function, and quality of life?
   b. How does effectiveness vary depending on: (1) the specific type or cause of pain (eg, neuropathic, musculoskeletal [including low back pain], fibromyalgia, sickle cell disease, inflammatory pain, and headache disorders); (2) patient demographics (eg, age, race, ethnicity, gender); and (3) patient comorbidities (including past or current alcohol or substance use disorders, mental health disorders, medical comorbidities, and high risk for addiction)?
   c. In patients with chronic pain, what is the comparative effectiveness of opioids vs nonopioid therapies (pharmacologic or nonpharmacologic) on outcomes related to pain, function, and quality of life?
   d. In patients with chronic pain, what is the comparative effectiveness of immediate-release plus nonopioid interventions (pharmacologic or nonpharmacologic) vs opioids or nonopioid interventions alone on outcomes related to pain, function, quality of life, and doses of opioids used?

Key Question 2. Harms and Adverse Events
   a. In patients with chronic pain, what are the risks of opioids vs placebo or no opioid on (1) opioid abuse, addiction, and related outcomes; (2) overdose; and (3) other harms, including gastrointestinal-related harms, falls, fractures, motor vehicle crashes, endocrinologic harms, infections, cardiovascular events, cognitive harms, and psychological harms (eg, depression)?
   b. How do harms vary depending on (1) the specific type or cause of pain (eg, neuropathic, musculoskeletal [including back pain], fibromyalgia, sickle cell disease, inflammatory pain, headache disorders); (2) patient demographics; (3) patient comorbidities (including past or current substance use disorder or at high risk for addiction); and (4) the dose of opioids used?

Key Question 3. Dosing Strategies
   a. In patients with chronic pain, what is the comparative effectiveness of different methods for initiating and titrating opioids on outcomes related to pain, function, and quality of life; risk of overdose, addiction, abuse, or misuse; and doses of opioids used?
   b. In patients with chronic pain, what is the comparative effectiveness of immediate-release vs extended-release/long-acting (ER/LA) opioids on outcomes related to pain, function, and quality of life; risk of overdose, addiction, abuse, or misuse; and doses of opioids used?
   c. In patients with chronic pain, what is the comparative effectiveness of different ER/LA opioids on outcomes related to pain, function, and quality of life; risk of overdose, addiction, abuse, or misuse; and doses of opioids used?
   d. In patients with chronic pain, what is the comparative effectiveness of immediate-release plus ER/LA opioids vs ER/LA opioids alone on outcomes related to pain, function, and quality of life; risk of overdose, addiction, abuse, or misuse; and doses of opioids used?

Key Question 4. Risk Assessment and Risk Mitigation Strategies
   a. In patients with chronic pain being considered for long-term opioid therapy, what is the accuracy of instruments for predicting risk of opioid overdose, addiction, abuse, or misuse?
   b. In patients with chronic pain, what is the effectiveness of use of risk prediction instruments on outcomes related to overdose, addiction, abuse, or misuse?
   c. In patients with chronic pain prescribed long-term opioid therapy, what is the effectiveness of risk mitigation strategies, including (1) opioid management plans, (2) patient education, (3) urine drug screening, (4) use of prescription drug monitoring program data, (5) use of monitoring instruments, (6) more frequent monitoring intervals, (7) pill counts, and (8) use of abuse-deterrent formulations on outcomes related to overdose, addiction, abuse, or misuse?
   d. What is the comparative effectiveness of different treatment strategies for managing patients with addiction to prescription opioids on outcomes related to overdose, abuse, misuse, pain, function, and quality of life?

Key Question 5. Effect of Opioid Therapy for Acute Pain on Long-term Use
   a. In patients with acute pain, what are the effects of prescribing opioid therapy vs not prescribing opioid therapy for acute pain on long-term opioid use?

Key questions 1-4 were developed for the Agency for Healthcare Research and Quality review.7

Tool41-43 and limited for other risk assessment instruments.41,44,45

No study evaluated the effectiveness of risk mitigation strategies.

KQ5: Effect of Opioid Therapy for Acute Pain on Long-term Use

Studies examining patients who underwent low-risk surgery or experienced low back pain from injury revealed that opioid therapy prescribed for acute pain was associated with greater likelihood of long-term use.46,47 Compared with no early opioid use for acute low back pain, the adjusted odds ratio for receiving 5 or more opioid prescriptions from 30 to 730 days after onset was 2.08 (95% CI, 1.55-2.78) for 1 to 140 morphine milligram equivalents (MME) per day and increased to 6.14 (95% CI, 4.92-7.66) for 450 MME or more per day.47
Table 1. GRADE Ratings of the Evidence for the Key Clinical Questions

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of Evidence</th>
<th>Other Factors</th>
<th>Estimates of Effect or Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Effectiveness and Comparative Effectiveness (Key Question 1)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Effectiveness of long-term opioid therapy vs placebo or no opioid therapy for long-term (≥1 y) outcomes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain, function, and quality of life</td>
<td>None</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Insufficient</td>
<td>NA</td>
<td>No evidence.</td>
</tr>
<tr>
<td>Harms and Adverse Events (Key Question 2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Risks of opioids vs placebo or no opioids on opioid abuse, addiction, and related outcomes; overdose; and other harms</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abuse or addiction</td>
<td>1 cohort study (n = 568640)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>One retrospective cohort study found long-term use of prescribed opioids was associated with an increased risk of abuse or dependence diagnosis vs no opioid use (adjusted OR range, 14.9-122.5, depending on dose).</td>
</tr>
<tr>
<td>Abuse or addiction</td>
<td>10 uncontrolled studies (n = 3780)</td>
<td>Very serious limitations</td>
<td>Very serious inconsistency</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>In primary care settings, prevalence of opioid abuse ranged from 0.6%-8%; prevalence of dependence, 3%-26%. In pain clinic settings, prevalence of misuse, 8%-16%, and addiction, 2%-14%. Prevalence of aberrant drug-related behaviors, 6%-37%.</td>
</tr>
<tr>
<td>Overdose</td>
<td>1 cohort study (n = 9940)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Current opioid use associated with increased risk of any overdose events, adjusted HR, 5.2 (95% CI, 2.1-12), and serious overdose events, adjusted HR, 8.4 (95% CI, 2.5-28) vs current nonuse.</td>
</tr>
<tr>
<td>Fractures</td>
<td>1 cohort study (n = 2341)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Opioid use associated with increased risk of fracture in 1 cohort study, adjusted HR, 1.28 (95% CI, 0.99-1.64), and 1 case-control study, adjusted OR, 1.27 (95% CI, 1.21-1.33).</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>1 cohort study (n = 426 124)</td>
<td>No limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Current opioid use associated with increased risk of myocardial infarction vs nonuse, adjusted OR, 1.28 (95% CI, 1.19-1.37) and IRR, 2.66 (95% CI, 2.30-3.08).</td>
</tr>
<tr>
<td>Endocrinologic harms</td>
<td>1 cross-sectional study (n = 11327)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Long-term opioid use associated with increased risk for use of medications for erectile dysfunction or testosterone replacement vs nonuse, adjusted OR, 1.5 (95% CI, 1.1-1.9).</td>
</tr>
<tr>
<td>How do harms vary depending on the opioid dose used?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abuse or addiction</td>
<td>1 cohort study (n = 568640)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>One retrospective cohort study found higher doses of long-term opioid therapy associated with increased risk of opioid abuse or dependence than lower doses. Compared with no opioid prescription, the adjusted ORs were 15 (95% CI, 10-21) for 1-36 MME/day, 29 (95% CI, 20-40) for 36-120 MME/day, and 122 (95% CI, 73-205) for ≥120 MME/day.</td>
</tr>
</tbody>
</table>
Table 1. GRADE Ratings of the Evidence for the Key Clinical Questionsa (continued)

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of Evidenceb</th>
<th>Other Factors</th>
<th>Estimates of Effect or Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Overdose</td>
<td>1 cohort study (n = 9940) and 1 case-control study (n = 593 case patients in primary analysis)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>Magnitude of effect, dose-response relationship</td>
<td>Compared with 1–20 MME/d, 1 cohort study found an adjusted HR for an overdose event of 1.44 (95% CI, 0.57–3.62) for 20–&lt;50 MME/d that increased to 8.87 (95% CI, 3.99–19.72) at ≥100 MME/d; 1 case-control study found an adjusted OR for an opioid-related death of 1.32 (95% CI, 0.34–1.19) for 20–49 MME/d that increased to 2.88 (95% CI, 1.79–4.63) at ≥200 MME/d.</td>
</tr>
<tr>
<td>Fractures</td>
<td>1 cohort study (n = 2341)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Risk of fracture increased from an adjusted HR of 1.20 (95% CI, 0.92–1.56) at 1–&lt;20 MME/d to 2.00 (95% CI, 1.24–3.24) at ≥50 MME/d; the trend was of borderline statistical significance.</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>1 cohort study (n = 426 124)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Relative to a cumulative dose of 0–1350 MME during a 90-d period, the IRR for myocardial infarction for 1350–&lt;2700 MME was 1.21 (95% CI, 1.02–1.45); for 2700–&lt;8100 MME, 1.42 (95% CI, 1.21–1.67); for 8100–18 000 MME, 1.89 (95% CI, 1.54–2.33); and for ≥18 000 MME, 1.73 (95% CI, 1.32–2.26).</td>
</tr>
<tr>
<td>Motor vehicle crash injuries</td>
<td>1 case-control study (n = 5300 case patients)</td>
<td>No limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>No association between opioid dose and risk of motor vehicle crash injuries even though opioid dosages ≥20 MME/d were associated with increased odds of road trauma among drivers.</td>
</tr>
<tr>
<td>Endocrinologic harms</td>
<td>1 cross-sectional study (n = 11 327); new for update: 1 additional cross-sectional study (n = 1585)</td>
<td>Serious limitations</td>
<td>Consistent</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Relative to 0–20 MME/d, the adjusted OR for ≥120 MME/d for use of medications for erectile dysfunction or testosterone replacement was 1.6 (95% CI, 1.0–2.4). One new cross-sectional study found higher-dose long-term opioid therapy associated with increased risk of androgen deficiency among men receiving immediate-release opioids, adjusted OR per 10 MME/d, 1.16 (95% CI, 1.09–1.23), but the dose response was very weak among men receiving ER/LA opioids.</td>
</tr>
</tbody>
</table>

Dosing Strategies (Key Question 3)

Comparative effectiveness of different methods for initiating opioid therapy and titrating doses

| Pain                        | 3 randomized trials (n = 93)                                        | Serious limitations | Serious inconsistency | Very serious imprecision | 4 | None identified | Trials on effects of titration with immediate-release vs ER/LA opioids reported inconsistent results and had additional differences between trial groups in dosing protocols (titrated vs fixed doses) and doses of opioids used. |
| Overdose                    | New for update: 1 cohort study (n = 840 606)                        | Serious limitations | Unknown (1 study)    | No imprecision           | 4 | None identified | One new cross-sectional study found initiation of therapy with an ER/LA opioid associated with increased risk of overdose vs initiation with an immediate-release opioid, adjusted HR, 2.33 (95% CI, 1.26–4.32). |

(continued)
<table>
<thead>
<tr>
<th>Outcome</th>
<th>Studies</th>
<th>Limitations</th>
<th>Inconsistency</th>
<th>Imprecision</th>
<th>Type of Evidence</th>
<th>Other Factors</th>
<th>Estimates of Effect or Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Comparative effectiveness of different ER/LA opioids</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain and function</td>
<td>3 randomized trials (n = 1850)</td>
<td>Serious limitations</td>
<td>No inconsistency</td>
<td>No imprecision</td>
<td>3</td>
<td>None identified</td>
<td>No differences.</td>
</tr>
<tr>
<td>All-cause mortality</td>
<td>1 cohort study (n = 108,492); new for update: 1 cohort study (n = 38,756)</td>
<td>Serious limitations</td>
<td>Serious inconsistency</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One cohort study found methadone to be associated with lower all-cause mortality risk than sustained-release morphine in a propensity-adjusted analysis, adjusted HR, 0.56 (95% CI, 0.51-0.62). One cohort study among Tennessee Medicaid patients found methadone to be associated with higher risk of all-cause mortality than sustained-release morphine, adjusted HR, 1.46 (95% CI, 1.17-1.73).</td>
</tr>
<tr>
<td>Abuse and related outcomes</td>
<td>1 cohort study (n = 5,684)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Serious imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One cohort study found some differences between ER/LA opioids in rates of adverse outcomes related to abuse, but outcomes were nonspecific for opioid-related adverse events, precluding reliable conclusions.</td>
</tr>
<tr>
<td>ER/LA vs immediate-release opioids</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Endocrinologic harms</td>
<td>New for update: 1 cross-sectional study (n = 1585)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>No imprecision</td>
<td>4</td>
<td>None identified</td>
<td>One cross-sectional study found ER/LA opioids associated with increased risk of androgen deficiency vs immediate-release opioids, adjusted OR, 3.39 (95% CI, 2.39-4.77).</td>
</tr>
<tr>
<td>Dose escalation vs dose maintenance or use of dose thresholds</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Immediate-release vs ER/LA opioids, immediate-release plus ER/LA opioids vs ER/LA opioids alone, scheduled and continuous vs as-needed dosing of opioids, or opioid rotation vs maintenance of current therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain, function, quality of life, and outcomes related to abuse</td>
<td>None</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>Insufficient</td>
<td>NA</td>
<td>No evidence.</td>
</tr>
<tr>
<td>Effects of decreasing or tapering opioid doses vs continuation of opioid therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain and function</td>
<td>1 randomized trial (n = 140)</td>
<td>Serious limitations</td>
<td>Unknown (1 study)</td>
<td>Very serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>No difference between more liberal dose escalation vs maintenance of current doses in pain, function, or risk of withdrawal due to opioid misuse, but there was limited separation in opioid doses between groups (52 vs 40 MME/d at the end of the trial).</td>
</tr>
<tr>
<td>Comparative effectiveness of different tapering protocols and strategies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Opioid abstinence</td>
<td>2 nonrandomized trials (n = 150)</td>
<td>Very serious limitations</td>
<td>No inconsistency</td>
<td>Very serious imprecision</td>
<td>4</td>
<td>None identified</td>
<td>No clear differences between different methods for opioid discontinuation or tapering in likelihood of opioid abstinence after 3-6 mo.</td>
</tr>
<tr>
<td>Outcome</td>
<td>Studies</td>
<td>Limitations</td>
<td>Inconsistency</td>
<td>Imprecision</td>
<td>Type of Evidence</td>
<td>Other Factors</td>
<td>Estimates of Effect or Findings</td>
</tr>
<tr>
<td>---------</td>
<td>---------</td>
<td>-------------</td>
<td>---------------</td>
<td>-------------</td>
<td>-----------------</td>
<td>--------------</td>
<td>--------------------------------</td>
</tr>
<tr>
<td>Risk Assessment and Risk Mitigation Strategies (Key Question 4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diagnostic accuracy of instruments for predicting risk for opioid overdose, addiction, abuse, or misuse among patients with chronic pain being considered for long-term opioid therapy</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Opioid Risk Tool</td>
<td>3 studies of diagnostic accuracy (n = 496); new for update: 2 studies of diagnostic accuracy (n = 320)</td>
<td>Serious limitations</td>
<td>Very serious inconsistency</td>
<td>Serious imprecision</td>
<td>4</td>
<td>None identified</td>
<td>Based on a cutoff score of &gt;4 (or unspecified), 5 studies (2 fair-quality, 3 poor-quality) reported sensitivity that ranged from 0.20-0.99 and specificity that ranged from 0.16-0.88.</td>
</tr>
<tr>
<td>Screener and opioid assessment for patients with pain, version 1</td>
<td>2 studies of diagnostic accuracy (n = 203)</td>
<td>Very serious limitations</td>
<td>No inconsistency</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Based on a cutoff score of ≥8, sensitivity was 0.68 and specificity was 0.38 in 1 study, for a positive likelihood ratio of 1.11 and a negative likelihood ratio of 0.83. Based on a cutoff score of &gt;6, sensitivity was 0.73 in 1 study.</td>
</tr>
<tr>
<td>Screener and opioid assessment for patients with pain: revised</td>
<td>New for update: 2 studies of diagnostic accuracy (n = 320)</td>
<td>Very serious limitations</td>
<td>No inconsistency</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Based on a cutoff score of &gt;3 or unspecified, sensitivity was 0.25 and 0.53 and specificity was 0.62 and 0.73 in 2 studies, for likelihood ratios close to 1.</td>
</tr>
<tr>
<td>Brief risk interview</td>
<td>New for update: 2 studies of diagnostic accuracy (n = 320)</td>
<td>Very serious limitations</td>
<td>No inconsistency</td>
<td>Serious imprecision</td>
<td>3</td>
<td>None identified</td>
<td>Based on a &quot;high-risk&quot; assessment, sensitivity was 0.73 and 0.83 and specificity was 0.43 and 0.88 in 2 studies, for positive likelihood ratios of 1.28 and 7.18 and negative likelihood ratios of 0.63 and 0.19.</td>
</tr>
<tr>
<td>Effectiveness of risk prediction instruments on outcomes related to overdose, addiction, abuse, or misuse in patients with chronic pain</td>
<td></td>
<td>None</td>
<td>NA</td>
<td>NA</td>
<td>Insufficient</td>
<td>NA</td>
<td>No evidence.</td>
</tr>
<tr>
<td>Effectiveness of risk mitigation strategies, including opioid management plans, patient education, urine drug screening, use of prescription drug monitoring program data, use of monitoring instruments, more frequent monitoring intervals, pill counts, and use of abuse-deterrent formulations, on outcomes related to overdose, addiction, abuse, or misuse</td>
<td></td>
<td>None</td>
<td>NA</td>
<td>NA</td>
<td>Insufficient</td>
<td>NA</td>
<td>No evidence.</td>
</tr>
<tr>
<td>Comparative effectiveness of treatment strategies for managing patients with addiction to prescription opioids</td>
<td></td>
<td>None</td>
<td>NA</td>
<td>NA</td>
<td>Insufficient</td>
<td>NA</td>
<td>No evidence.</td>
</tr>
</tbody>
</table>
Contextual Evidence Review

CDC conducted a contextual evidence review to assist in developing the recommendations by providing an assessment of the balance of benefits and harms, values and preferences, and cost, consistent with the GRADE approach (Box 3). Rapid review methods were used to streamline the process and obtain evidence quickly (e.g., by limiting database searches and summarizing study quality based on author reports rather than applying objective quality rating protocols). Full details on methodology, including data sources and searches, inclusion criteria, study selection, and data extraction and synthesis, and findings are available in the MMWR report.11 In this article, we summarize benefits and harms of nonopioid therapies found in the clinical literature and harms of opioid therapy, including additional studies not included in the clinical evidence review (e.g., studies not restricted to patients with chronic pain).

Several nonpharmacologic and nonopioid pharmacologic treatments were found to be effective for chronic pain in studies ranging in duration from 2 weeks to 6 months48-66 (Table 3). For example, cognitive behavioral therapy (CBT) had small positive effects on disability and catastrophic thinking.66 Exercise therapy reduced pain and improved function in chronic low back pain54; improved function and reduced pain in osteoarthritis of the knee53 and hip52; and improved well-being, fibromyalgia symptoms, and physical function in fibromyalgia.48 Multimodal and multidisciplinary therapies helped reduce pain and improve function more effectively than single modalities.55,67 Multiple guidelines recommended acetaminophen as first-line pharmacotherapy for osteoarthritis68-73 or for low back pain74 and nonsteroidal anti-inflammatory drugs (NSAIDs) as first-line treatment for osteoarthritis or low back pain70,74; first- and second-line drugs for neuropathic pain include anticonvulsants (gabapentin or pregabalin), tricyclic antidepressants, and serotonin-norepinephrine reuptake inhibitors (SNRIs).75-78 Nonsteroidal anti-inflammatory drugs have been associated with hepatic, gastrointestinal, renal, and cardiovascular risks.63,73,79

Opioid-related overdose risk was dose-dependent, with higher opioid dosages associated with increased overdose risk (Table 2).19-27 Compared with dosages of 1 to <20 MME per day, dosages of 50 to <100 MME per day were found to increase risks for opioid overdose by factors of 1.920 to 4.6,22 with absolute risk difference approximation of 0.15% for fatal overdose22 and 1.40% for any overdose;19 dosages of 100 MME or more per day were found to increase risks for opioid overdose by factors of 2.020 to 8.919 relative to dosages of 1 to <20 MME per day, with absolute risk difference approximation 0.25% for fatal overdose24 and 4.04% for any overdose.79 Veterans Health Administration patients with chronic pain who died of overdoses related to opioids were prescribed higher mean opioid dosages (98 MME/d) than controls (48 MME/d),27 above 200 MME per day, mortality rates continue to increase more gradually.23 (See Table 4 and Box 4 for a list of common opioid medications and their MME equivalents.)

Other findings included disproportionate numbers of overdose deaths associated with methadone80; fatal overdose risk associated with co-prescription of opioids and benzodiazepines82,83; and risks associated with sleep-disordered breathing,82,83 reduced renal or hepatic function,84 older age,85-88 pregnancy,89-92 mental health comorbidities, and history of substance use disorder.88,93,94 Indirect evidence was found for potential utility of
risk stratification and mitigation strategies for identifying risky opioid-taking behaviors and prescribing practices, such as checking prescription drug monitoring program (PDMP) data\textsuperscript{95} and urine drug testing.\textsuperscript{96} as well as co-prescription of naloxone.\textsuperscript{97} In addition, methadone and buprenorphine for opioid use disorder were found to increase retention in treatment and to decrease illicit opioid use among patients with opioid use disorder, and some studies suggest that effectiveness is enhanced when psychosocial treatments are used in conjunction with medication-assisted therapy.\textsuperscript{98-102}

**Recommendations**

The guideline includes 12 recommendations (Box 5). GRADE recommendation categories were based on the following assessment:

- No evidence shows a long-term benefit of opioids in pain and function vs no opioids for chronic pain with outcomes examined at least 1 year later (with most placebo-controlled randomized clinical trials ≤6 weeks in duration).
- Extensive evidence shows the possible harms of opioids (including opioid use disorder, overdose, and motor vehicle injury).
- Extensive evidence suggests some benefits of nonpharmacologic and nonopioid pharmacologic therapy, with less harm.

**Determining When to Initiate or Continue Opioids for Chronic Pain**

1. Nonpharmacologic therapy and nonopioid pharmacologic therapy are preferred for chronic pain. Clinicians should consider opioid therapy only if expected benefits for both pain and function are anticipated to outweigh risks to the patient. If opioids are used, they should be combined with nonpharmacologic therapy.
and nonopioid pharmacologic therapy, as appropriate. (Recommendation category: A; evidence type: 3)

Nonpharmacologic therapy (such as exercise therapy and CBT) should be used to reduce pain and improve function in patients with chronic pain. Aspects of these approaches can be used even when there is limited access to specialty care. For example, primary care clinicians can encourage patients to take an active role in the care plan and support patients in engaging in exercise. Nonopioid pharmacologic therapy (such as NSAIDs, acetaminophen, anticonvulsants, and SNRIs) should be used when benefits outweigh risks and should be combined with nonpharmacologic therapy. Opioids should not be considered first-line or routine therapy for chronic pain outside of active cancer, palliative, and end-of-life care, given small to moderate short-term benefits, uncertain long-term benefits, and potential for serious harms; although evidence on long-term benefits of nonopioid therapies is also limited, these therapies are also associated with short-term benefits, and risks are much lower. This does not mean that patients should be required to sequentially “fail” nonpharmacologic and nonopioid pharmacologic therapy before proceeding to opioid therapy. Rather, expected benefits specific to the clinical context should be weighed against risks before initiating therapy. In some clinical contexts (eg, headache, fibromyalgia), expected benefits of initiating opioids are unlikely to outweigh risks regardless of previous nonpharmacologic and nonopioid pharmacologic therapies used. In other situations (eg, serious illness in a patient with poor prognosis for return to previous level of function, contraindications to other therapies, and clinician and patient agreement that the overriding goal is patient comfort), opioids might be appropriate regardless of previous therapies used. If opioids are used, they should be combined with nonpharmacologic therapy and nonopioid pharmacologic therapy, as appropriate, to provide greater benefits to patients.

2. Before starting opioid therapy for chronic pain, clinicians should establish treatment goals with all patients, including realistic goals for pain and function, and consider how opioid therapy will be discontinued if benefits do not outweigh risks. Clinicians should continue opioid therapy only if there is clinically meaningful improvement in pain and function that outweighs risks to patient safety. (Recommendation category: A; evidence type: 4)

Before opioid therapy is initiated for chronic pain, clinicians should determine how effectiveness will be evaluated and should establish treatment goals with patients. Clinicians seeing new patients already receiving opioids should establish treatment goals for continued treatment. Goals should include improvement in both pain relief and function. However, there are some clinical circumstances under which reductions in pain without improvement in physical function might be a more realistic goal (eg, diseases typically associated with progressive functional impairment or catastrophic injuries such as spinal cord trauma). Experts noted that function can include emotional and social as well as physical dimensions. In addition, experts emphasized that mood has important interactions with pain and function. Clinicians may use validated instruments such as the 3-item “Pain average, interference with Enjoyment of life, and interference with General activity” (PEG) Assessment Scale to track patient outcomes. Clinically meaningful improvement has been defined as a 30% improvement in scores for both pain and function. Because depression, anxiety, and other psychological comorbidities often coexist with and can interfere with resolution of pain, clinicians should use validated instruments to assess for these conditions and ensure that treatment for these conditions is optimized.

3. Before starting and periodically during opioid therapy, clinicians should discuss with patients known risks and realistic benefits of opioid therapy and patient and clinician responsibilities for managing therapy. (Recommendation category: A; evidence type: 3)

Clinicians should ensure that patients are aware of potential benefits of, harms of, and alternatives to opioids before starting or continuing opioid therapy. Clinicians are encouraged to have open and honest discussions with patients to inform mutual decisions about whether to start or continue opioid therapy. Important considerations include the following:

- Be explicit and realistic about expected benefits of opioids, explaining that while opioids can reduce pain during short-term use, there is no good evidence that opioids improve pain or function with long-term use and that complete relief of pain is unlikely.
- Emphasize improvement in function as a primary goal and that function can improve even when pain is still present.
- Advise patients about serious adverse effects of opioids, including potentially fatal respiratory depression and development of a potentially serious lifelong opioid use disorder.
- Advise patients about common effects of opioids, such as constipation, dry mouth, nausea, vomiting, drowsiness, confusion, tolerance, physical dependence, and withdrawal symptoms when stopping opioids.
- Discuss effects that opioids may have on ability to safely operate a vehicle, particularly when opioids are initiated, when dosages are increased, or when other central nervous system depressants, such as benzodiazepines or alcohol, are used concurrently.
Table 3. Effectiveness and Harms of Nonpharmacologic and Nonopioid Pharmacologic Treatments

<table>
<thead>
<tr>
<th>Source</th>
<th>Topic or Intervention</th>
<th>Participants or Population</th>
<th>Primary Outcomes</th>
<th>Key Findings</th>
<th>Study Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Busch et al. 2007</td>
<td>Exercise training vs untreated control or no exercise intervention</td>
<td>Systematic review of 33 RCTs with fibromyalgia patients</td>
<td>Global well-being, selected signs and symptoms, and physical function</td>
<td>Exercise training improves global well-being and physical function; supervised aerobic exercise training has beneficial effects on physical capacity and fibromyalgia symptoms.</td>
<td>Four studies were classified as high quality, 15 as moderate quality, and 14 as low quality</td>
</tr>
<tr>
<td>Chaparro et al. 2014</td>
<td>Noninjectable opioids vs placebo or other treatments</td>
<td>Systematic review of 15 RCTs with patients with chronic low back pain</td>
<td>Pain</td>
<td>One trial found tramadol similar to celecoxib for pain relief. Two trials did not find a difference between opioids and antidepressants for pain or function.</td>
<td>Low- to moderate-quality evidence</td>
</tr>
<tr>
<td>Collins et al. 2000</td>
<td>Antidepressants vs placebo; anticonvulsants vs placebo</td>
<td>Systematic review of 19 RCTs for diabetic neuropathy or postherpetic neuralgia</td>
<td>Pain</td>
<td>For diabetic neuropathy, the NNT for ≥50% pain relief was 3.4 for antidepressants (12 trials, 10 evaluated TCAs and 3 SSRIIs) and 2.7 for anticonvulsants (3 trials). For postherpetic neuralgia, the NNT was 2.1 for antidepressants (3 studies evaluating TCAs) and 3.2 for anticonvulsants (1 study evaluating gabapentin).</td>
<td>The mean and median quality score for included studies was 4 on a scale of 1-5</td>
</tr>
<tr>
<td>Fransen et al. 2015</td>
<td>Exercise vs nonexercise group (active or no treatment)</td>
<td>Systematic review of 54 RCTs or quasi-randomized trials for knee osteoarthritis</td>
<td>Reduced joint pain or improved physical function and quality of life</td>
<td>Exercise reduced pain, improved function, and improved quality of life immediately after treatment; in studies providing posttreatment follow-up data, improved pain and function were sustained for 2-6 mo.</td>
<td>High-quality evidence for reduced pain and improved quality of life and moderate-quality evidence for improved function</td>
</tr>
<tr>
<td>Fransen et al. 2014</td>
<td>Exercise vs nonexercise group (active or no treatment)</td>
<td>Systematic review of 10 RCTs or quasi-randomized trials for hip osteoarthritis</td>
<td>Reduced joint pain and improved physical function and quality of life</td>
<td>Exercise reduced pain and improved function immediately after treatment; in studies providing posttreatment follow-up data, improved pain and function were sustained for at least 3-6 mo.</td>
<td>High-quality evidence for reduced pain and improved function</td>
</tr>
<tr>
<td>Häuser et al. 2014</td>
<td>Duloxetine vs placebo; milnacipran vs placebo</td>
<td>Systematic review of 10 RCTs for fibromyalgia patients</td>
<td>Benefits and harms</td>
<td>Duloxetine and milnacipran reduced pain by a small amount compared with placebo.</td>
<td>Risk of bias in included studies was low</td>
</tr>
<tr>
<td>Hayden et al. 2005</td>
<td>Exercise therapy vs no treatment, other conservative treatments</td>
<td>Systematic review consisting of 61 RCTs for low back pain</td>
<td>Pain, function</td>
<td>Exercise therapy reduces pain and improves function with small magnitudes of effect. Effectiveness of exercise therapy appears to be greater in populations visiting a health care provider compared with the general population.</td>
<td>Only a small number of studies rated as high quality; potential publication bias</td>
</tr>
<tr>
<td>Lee et al. 2014</td>
<td>CIM therapies vs single self-care CIM, non–self-care CIM, usual care/no treatment, other multimodal program, or other control</td>
<td>Systematic review of 26 RCTs for management of chronic pain</td>
<td>Pain symptoms</td>
<td>Integrative multimodal therapies resulted in positive, but sometimes mixed, effects on pain symptoms compared with active controls or single self-care modalities. More studies are needed to make strong conclusions about effectiveness.</td>
<td>Large majority of poor quality, including weaknesses in randomization and allocation concealment</td>
</tr>
<tr>
<td>Lunn et al. 2014</td>
<td>Duloxetine vs placebo or other controls</td>
<td>Systematic review of 18 RCTs for neuropathic pain, chronic pain conditions without identified cause, or fibromyalgia</td>
<td>Benefits and harms of duloxetine</td>
<td>Duloxetine at 60 mg and 120 mg daily, but not lower dosages, were effective in reducing pain in diabetic peripheral neuropathy pain and in fibromyalgia.</td>
<td>Moderate-quality evidence for diabetic neuropathy; lower-quality evidence for fibromyalgia; some risk of bias</td>
</tr>
<tr>
<td>Moore et al. 2009</td>
<td>Pregabalin vs placebo or any active control</td>
<td>Systematic review of 25 double-blind RCTs for postherpetic neuralgia, painful diabetic neuropathy, central neuropathic pain, and fibromyalgia</td>
<td>Analgesic efficacy and associated adverse events</td>
<td>Pregabalin was effective in patients with postherpetic neuralgia, diabetic neuropathy, central neuropathic pain, and fibromyalgia at doses of 300 mg, 450 mg, and 600 mg (but not at 150 mg) daily. NNTs were generally ≤6 for moderate benefit in postherpetic neuralgia and diabetic neuropathy but &gt;7 for fibromyalgia.</td>
<td>Studies all had Oxford quality scores based on randomization, blinding, and reporting of dropout ≥3 (out of maximum of 5)</td>
</tr>
<tr>
<td>Moore et al. 2014</td>
<td>Gabapentin vs placebo</td>
<td>Systematic review of 37 RCTs for neuropathic pain or fibromyalgia</td>
<td>Analgesic efficacy and adverse effects</td>
<td>Gabapentin was significantly more effective than placebo in reducing pain in diabetic neuropathy and postherpetic neuralgia. Evidence was insufficient for other conditions.</td>
<td>“Second-tier” evidence (some risk of bias, but adequate numbers in the trials)</td>
</tr>
</tbody>
</table>

(continued)
<table>
<thead>
<tr>
<th>Source</th>
<th>Topic or Intervention</th>
<th>Participants or Population</th>
<th>Primary Outcomes</th>
<th>Key Findings</th>
<th>Study Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roe13</td>
<td>NSAIDs and COX-2 inhibitors vs control</td>
<td>Systematic review of 65 RCTs for nonspecific low back pain</td>
<td>Acute low back pain</td>
<td>NSAIDs are more effective than placebo for acute and chronic low back pain without sciatica, but have more adverse effects. NSAIDs are not more effective than acetaminophen but had more adverse effects. No type of NSAIDs, including COX-2 inhibitors, was found to be more effective than other NSAIDs.</td>
<td>Mixed high- and low-quality studies</td>
</tr>
<tr>
<td>Saarto et al. 66 2010</td>
<td>Antidepressants vs placebo or other controls</td>
<td>Systematic review of 61 RCTs for neuropathic pain</td>
<td>Pain</td>
<td>TCAs and venlafaxine have low NNTs (3.6 and 3.1, respectively) for at least moderate pain relief. Study quality limited by insufficient reporting detail</td>
<td>Moderate-quality studies</td>
</tr>
<tr>
<td>Salerno et al. 61 2002</td>
<td>Antidepressants vs placebo</td>
<td>Systematic review of 9 RCTs for chronic back pain</td>
<td>Back pain</td>
<td>Antidepressants were associated with small but significant improvement in pain severity; improvements in function were not significant. Most (6) studies evaluated TCAs.</td>
<td>Moderate-quality studies</td>
</tr>
<tr>
<td>Staiger et al. 60 2010</td>
<td>Antidepressants vs placebo</td>
<td>Systematic review of 7 RCTs in patients with chronic low back pain</td>
<td>Back pain</td>
<td>Four of 5 studies evaluating TCA and tetracyclic antidepressants found significant improvement in chronic low back pain. Other antidepressants studied (2 studies evaluating SSRIs and 1 evaluating trazodone) did not show significant pain improvement.</td>
<td>Mixed quality (quality scores ranged from 11-19 out of 22)</td>
</tr>
<tr>
<td>Trelle et al. 62 2008</td>
<td>NSAIDs vs other NSAIDs or placebo</td>
<td>Meta-analysis of 31 RCTs comparing any NSAID with other NSAID or placebo for any medical condition</td>
<td>Myocardial infarction, stroke, cardiovascular death, death from any cause</td>
<td>Compared with placebo, NSAIDs were associated with increased risk of myocardial infarction, stroke, and cardiovascular death. Study quality limited by small numbers of participants and outcomes of limited clinical utility or both.</td>
<td>Generally high</td>
</tr>
<tr>
<td>Welsch et al. 64 2015</td>
<td>Opioids (including tramadol) vs nonopioids (including acetaminophen, NSAIDs/COX-2 inhibitors, mexiteline, anticonvulsants, antidepressants, and muscle relaxants)</td>
<td>Systematic review of 10 RCTs in patients with neuropathic pain, low back pain, or osteoarthritis</td>
<td>Efficacy (including various pain measures), tolerability, and safety</td>
<td>There was no significant difference between opioids and nonopioid analogics in pain reduction; nonopioids were superior to opioids in improving physical function and were better tolerated. When patients from tramadol trials (n randomized = 2788) were removed from results of the review, results for pain and function for patients receiving opioids (morphine) compared with alternative drugs (n randomized = 223) had wide, overlapping confidence intervals. Improved tolerability for alternative drugs vs morphine remained significant.</td>
<td>One study had a high, 2 studies a moderate, and 7 studies a low study quality</td>
</tr>
<tr>
<td>Wiffen et al. 65 2014</td>
<td>Carbamazepine vs placebo or other active control</td>
<td>Systematic review consisting of 10 RCTs in adults with chronic neuropathic pain or fibromyalgia</td>
<td>Pain relief</td>
<td>Carbamazepine provided better pain relief than placebo for trigeminal neuralgia, diabetic neuropathy, and poststroke pain for ≤4 weeks. Dizziness and drowsiness were commonly reported with carbamazepine. In 4 studies, 65% of patients receiving carbamazepine vs 27% receiving placebo experienced ≥1 adverse event. In 8 studies, 3% of patients receiving carbamazepine withdrew because of adverse events (vs 0% taking placebo).</td>
<td>Third-tier evidence (trials involving small numbers of participants, considered likely to be biased, with outcomes of limited clinical utility, or both)</td>
</tr>
<tr>
<td>Williams et al. 66 2012</td>
<td>Cognitive behavioral therapy or behavioral therapy</td>
<td>Systematic review of 42 RCTs for patients with nonmalignant chronic pain except headache</td>
<td>Pain, disability, mood, and catastrophic thinking</td>
<td>Cognitive behavioral therapy was found to have small to moderate effects on pain, disability, mood, and catastrophic thinking immediately after treatment when compared with usual treatment or deferred cognitive behavioral therapy, but only effects on mood persisted at follow-up. Behavioral therapy had a positive effect on mood immediately after treatment.</td>
<td>Mean quality of study design, 15.8 out of 26 (SD 4.3; range, 9-24 out of 26)</td>
</tr>
</tbody>
</table>

Abbreviations: CIM, complementary and integrative multimodal; COX-2, cyclooxygenase 2; NNT, number needed to treat; NSAID, nonsteroidal anti-inflammatory drug; RCTs, randomized clinical trials; SSRIs, selective serotonin reuptake inhibitors; TCA, tricyclic antidepressants.

* All the studies in this table were included in the contextual evidence review.
• Consider whether cognitive limitations might interfere with management of opioid therapy (for older adults in particular), and if so, determine whether a caregiver can responsibly co-manage medication therapy. Discuss the importance of reassessing safer medication use with both the patient and caregiver.

Opioid Selection, Dosage, Duration, Follow-up, and Discontinuation

4. When starting opioid therapy for chronic pain, clinicians should prescribe immediate-release opioids instead of extended-release/long-acting (ER/LA) opioids. (Recommendation category: A; evidence type: 4)

Clinicians should not initiate opioid treatment with ER/LA opioids and should not prescribe ER/LA opioids for intermittent use. In general, avoiding the use of immediate-release opioids in combination with ER/LA opioids is preferable.

When an ER/LA opioid is prescribed, using a product with predictable pharmacokinetics and pharmacodynamics is preferred to minimize unintentional overdose risk.

• Methadone should not be the first choice for an ER/LA opioid. Only clinicians who are familiar with methadone’s unique risk profile and who are prepared to educate and closely monitor their patients—including risk assessment for QT prolongation and consideration of electrocardiographic monitoring—should consider prescribing methadone for pain.

• Because dosing effects of transdermal fentanyl are often misunderstood by both clinicians and patients, only clinicians who are familiar with the dosing and absorption properties of transdermal fentanyl and are prepared to educate their patients about its use should consider prescribing it.

5. When opioids are started, clinicians should prescribe the lowest effective dosage. Clinicians should use caution when prescribing opioids at any dosage, should carefully reassess evidence of individual benefits and risks when considering increasing dosage to 50 morphine milligram equivalents (MME) or more per day, and should avoid increasing dosage to 90 MME or more per day or carefully justify a decision to titrate dosage to 90 MME or more per day. (Recommendation category: A; evidence type: 3)

Clinicians should start opioids at the lowest effective dosage, use caution when increasing opioid dosages, and increase dosage by the smallest practical amount. Before increasing total opioid dosage to 50 MME or more per day, clinicians should reassess whether

Table 4. Morphine Milligram Equivalent Doses for Commonly Prescribed Opioids

<table>
<thead>
<tr>
<th>Opioid</th>
<th>Conversion Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Codeine</td>
<td>0.15</td>
</tr>
<tr>
<td>Fentanyl transdermal, μg/h</td>
<td>2.4</td>
</tr>
<tr>
<td>Hydrocodone</td>
<td>1</td>
</tr>
<tr>
<td>Hydromorphone</td>
<td>4</td>
</tr>
<tr>
<td>Methadone, mg/d</td>
<td></td>
</tr>
<tr>
<td>1-20 mg/d</td>
<td>4</td>
</tr>
<tr>
<td>21-40 mg/d</td>
<td>8</td>
</tr>
<tr>
<td>41-60 mg/d</td>
<td>10</td>
</tr>
<tr>
<td>≥61-80 mg/d</td>
<td>12</td>
</tr>
<tr>
<td>Morphine</td>
<td>1</td>
</tr>
<tr>
<td>Oxycodone</td>
<td>1.5</td>
</tr>
<tr>
<td>Oxymorphone</td>
<td>3</td>
</tr>
<tr>
<td>Tapentadolc</td>
<td>0.4</td>
</tr>
<tr>
<td>*All doses are in mg/d except for fentanyl, which is μg/h. Multiply the daily dosage for each opioid by the conversion factor to determine the dose in morphine milligram equivalents (MME). For example, tablets containing hydrocodone, 5 mg, and acetaminophen, 300 mg, taken 4 times a day would contain a total of 20 mg of hydrocodone daily, equivalent to 20 MME daily; extended-release tablets containing oxycodone, 10 mg, and taken twice a day would contain a total of 20 mg of oxycodone daily, equivalent to 30 MME daily.</td>
<td></td>
</tr>
<tr>
<td>* Tapentadol is a μ-receptor agonist and norepinephrine reuptake inhibitor. Morphine milligram equivalents are based on degree of μ-receptor agonist activity, but it is unknown if this drug is associated with overdose in the same dose-dependent manner as observed with medications that are solely μ-receptor agonists.</td>
<td></td>
</tr>
</tbody>
</table>

Box 4. Cautions About Calculating Morphine Milligram Equivalent Doses

• Equianalgesic dose conversions are only estimates and cannot account for individual variability in genetics and pharmacokinetics.

• Do not use the calculated dose in morphine milligram equivalents (MME) to determine the doses to use when converting one opioid to another; when converting opioids, the new opioid is typically dosed at substantially lower than the calculated MME dose to avoid accidental overdose due to incomplete cross-tolerance and individual variability in opioid pharmacokinetics.

• Use particular caution with methadone dose conversions because the conversion factor increases at higher doses.

• Use particular caution with fentanyl because it is dosed in μg/h instead of mg/d, and its absorption is affected by heat and other factors.

Copyright 2016 American Medical Association. All rights reserved.
Clinical Review & Education Special Communication

Opioid Selection, Dosage, Duration, Follow-up, and Discontinuation

4. When starting opioid therapy for chronic pain, clinicians should prescribe immediate-release opioids instead of extended-release/long-acting (ER/LA) opioids.

5. When opioids are started, clinicians should discuss the lowest effective dosage. Clinicians should use caution when prescribing opioids at any dosage, should carefully reassess evidence of individual benefits and risks when increasing dosage to 50 morphine milligram equivalents (MME) or more per day, and should avoid increasing dosage to 90 MME or more per day or carefully justify a decision to titrate dosage to 90 MME or more per day.

6. Long-term opioid use often begins with treatment of acute pain. When opioids are used for acute pain, clinicians should prescribe the lowest effective dose of immediate-release opioids and should prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids. Three days or less will often be sufficient; more than 7 days will rarely be needed.

7. Clinicians should evaluate benefits and harms with patients within 1 to 4 weeks of starting opioid therapy for chronic pain or of dose escalation. Clinicians should evaluate benefits and harms of continued therapy with patients every 3 months or more frequently. If benefits do not outweigh harms of continued opioid therapy, clinicians should optimize therapies and work with patients to taper opioids to lower dosages or to taper and discontinue opioids.

Assessing Risk and Addressing Harms of Opioid Use

8. Before starting and periodically during continuation of opioid therapy, clinicians should evaluate risk factors for opioid-related harms. Clinicians should incorporate into the management plan strategies to mitigate risk, including considering offering naloxone when factors that increase risk for opioid overdose, such as history of overdose, history of substance use disorder, higher opioid dosages (≥50 MME/d), or concurrent benzodiazepine use are present.

9. Clinicians should review the patient’s history of controlled substance prescriptions using state prescription drug monitoring program (PDMP) data to determine whether the patient is receiving opioid dosages or dangerous combinations that put him or her at high risk for overdose. Clinicians should review PDMP data when starting opioid therapy for chronic pain and periodically during opioid therapy for chronic pain, ranging from every prescription to every 3 months.

10. When prescribing opioids for chronic pain, clinicians should use urine drug testing before starting opioid therapy and consider urine drug testing at least annually to assess for prescribed medications as well as other controlled prescription drugs and illicit drugs.

11. Clinicians should avoid prescribing opioid pain medication and benzodiazepines concurrently whenever possible.

12. Clinicians should offer or arrange evidence-based treatment (usually medication-assisted treatment with buprenorphine or methadone in combination with behavioral therapies) for patients with opioid use disorder.

All recommendations are category A (apply to all patients outside of active cancer treatment, palliative care, and end-of-life care) except recommendation 10 (designated category B, with individual decision making required); detailed ratings of the evidence supporting the recommendations are provided in the full guideline publication.

Established patients already prescribed high dosages of opioids (≥90 MME/d), including patients transferring from other clinicians, should be offered the opportunity to reevaluate their continued use of opioids at high dosages in light of recent evidence regarding the association of opioid dosage and overdose risk. For patients who agree to taper opioids to lower dosages, clinicians should collaborate with the patient on a tapering plan.

6. Long-term opioid use often begins with treatment of acute pain. When opioids are used for acute pain, clinicians should prescribe the lowest effective dose of immediate-release opioids and should prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids. Three days or less will often be sufficient; more than 7 days will rarely be needed. (Recommendation category: A; evidence type: 4)

Acute pain can often be managed without opioids. When diagnosis and severity of nontraumatic, nonsurgical pain are reasonably assumed to warrant the use of opioids, clinicians should...
prescribe no greater quantity than needed for the expected duration of pain severe enough to require opioids, often 3 days or less, unless circumstances clearly warrant additional opioid therapy. More than 7 days will rarely be needed. Postsurgical pain is outside the scope of this guideline but has been addressed elsewhere. Clinicians should not prescribe additional opioids to patients “just in case” pain continues longer than expected. Clinicians should reevaluate the subset of patients who experience severe acute pain that continues longer than the expected duration to confirm or revise the initial diagnosis and to adjust management accordingly. Clinicians should not prescribe ER/LA opioids for the treatment of acute pain.

7. Clinicians should evaluate benefits and harms with patients within 1 to 4 weeks of starting opioid therapy for chronic pain or of dose escalation. Clinicians should evaluate benefits and harms of continued therapy with patients every 3 months or more frequently. If benefits do not outweigh harms of continued opioid therapy, clinicians should optimize other therapies and work with patients to taper opioids to lower dosages or to taper and discontinue opioids. (Recommendation category: A; evidence type: 4)

Clinicians should evaluate patients to assess benefits and harms of opioids within 1 to 4 weeks of starting long-term opioid therapy or of dose escalation, consider follow-up intervals within the lower end of this range when ER/LA opioids are started or increased or when total daily opioid dosage is 50 MME per day or greater, and strongly consider shorter follow-up intervals (within 3 days) when starting or increasing the dosage of methadone. Clinicians should regularly reassess all patients receiving long-term opioid therapy, including patients who are new to the clinician but taking long-term therapy, at least every 3 months and reevaluate patients exposed to greater risk of opioid use disorder or overdose (eg, patients with depression or other mental health conditions, history of substance use disorder or overdose, taking ≥50 MME/d, taking other central nervous system depressants) more frequently.

At follow-up, clinicians should determine whether opioids continue to meet treatment goals, including sustained improvement in pain and function, whether the patient has experienced common or serious adverse events or has early warning signs of serious adverse events such as overdose (eg, sedation, slurred speech) or opioid use disorder (eg, difficulty controlling use), whether benefits of opioids continue to outweigh risks, and whether opioid dosage can be reduced or opioids can be discontinued.

Clinicians should work with patients to reduce opioid dosage or to discontinue opioids when possible if clinically meaningful improvements in pain and function are not sustained, if patients are taking high-risk regimens (eg, dosages ≥50 MME/d or opioids combined with benzodiazepines) without evidence of benefit, if patients believe benefits no longer outweigh risks or request dosage reduction or discontinuation, or if patients experience overdose or other serious adverse events or warning signs of serious adverse events.

When opioids are reduced or discontinued, a taper slow enough to minimize symptoms and signs of opioid withdrawal should be used. A decrease of 10% of the original dose per week is a reasonable starting point; tapering plans may be individualized based on patient goals and concerns. Slower tapers (eg, 10% per month) might be appropriate and better tolerated, particularly when patients have been taking opioids for years. More rapid tapers might be needed for patients who have overdosed on their current dosage. Clinicians should access appropriate expertise if considering tapering opioids during pregnancy because of possible risk to the pregnant patient and to the fetus if the patient goes into withdrawal. Primary care clinicians should collaborate with mental health clinicians and with other specialists as needed to optimize nonopioid pain management, as well as psychosocial support for anxiety related to the taper.

Assessing Risk and Addressing Harms of Opioid Use

8. Before starting and periodically during continuation of opioid therapy, clinicians should evaluate risk factors for opioid-related harms. Clinicians should incorporate into the management plan strategies to mitigate risk, including considering offering naloxone when factors that increase risk for opioid overdose, such as history of overdose, history of substance use disorder, higher opioid dosages (≥50 MME/d), or concurrent benzodiazepine use, are present. (Recommendation category: A; evidence type: 4)

Certain risk factors can increase susceptibility to opioid-associated harms. Clinicians should avoid prescribing opioids to patients with moderate or severe sleep-disordered breathing whenever possible. During pregnancy, clinicians and patients together should carefully weigh risks and benefits when making decisions about whether to initiate opioid therapy. Clinicians caring for pregnant women receiving opioids should arrange for delivery at a facility prepared to evaluate and treat neonatal opioid withdrawal syndrome. Clinicians should use additional caution and increased monitoring to minimize risks of opioids prescribed for patients with renal or hepatic insufficiency, patients 65 years and older, and patients with anxiety or depression. Clinicians should ensure that treatment for depression and other mental health conditions is optimized, consulting with behavioral health specialists when needed. If clinicians consider opioid therapy for patients with drug or alcohol use disorders or for patients with prior nonfatal overdose, they should discuss increased risks for opioid use disorder and overdose with patients, carefully consider whether benefits of opioids outweigh increased risks, and increase frequency of monitoring opioid therapy.

Clinicians should consider offering naloxone when prescribing opioids to patients at increased risk of overdose, including patients with a history of overdose, patients with a history of substance use disorder, patients taking benzodiazepines with opioids, patients at risk of returning to a high dose to which they are no longer tolerant (eg, patients recently released from prison), and patients taking higher dosages of opioids (≥50 MME/d). Practices should provide education on overdose prevention and naloxone use to patients receiving naloxone prescriptions and to members of their households.

9. Clinicians should review the patient’s history of controlled substance prescriptions using state prescription drug monitoring program (PDMP) data to determine whether the patient is receiving opioid dosages or dangerous combinations that put him or her at high risk for overdose. Clinicians should review PDMP data when starting opioid therapy for chronic pain and periodically during opioid therapy for chronic pain, ranging from every prescription to every 3 months. (Recommendation category: A; evidence type: 4)
Clinicians should review PDMP data for opioids and other controlled medications patients might have received from additional prescribers to determine whether a patient is receiving high total opioid dosages or dangerous combinations (eg, opioids combined with benzodiazepines) that put him or her at high risk for overdose. Ideally, PDMP data should be reviewed before every opioid prescription. This is recommended in all states with well-functioning PDMPs and where PDMP access policies make this practicable (eg, clinician and delegate access permitted), but it is not currently possible in states without functional PDMPs or in those that do not permit certain prescribers to access them.

If patients are found to have high opioid dosages, dangerous combinations of medications, or multiple controlled substance prescriptions written by different clinicians, several actions can be taken to augment clinicians’ abilities to improve patient safety:

- Clinicians should discuss information from the PDMP with their patient and confirm that the patient is aware of the additional prescriptions.
- Clinicians should discuss safety concerns, including increased risk for respiratory depression and overdose, with patients found to be receiving opioids from more than 1 prescriber or receiving medications that increase risk when combined with opioids (eg, benzodiazepines).
- Clinicians should avoid prescribing opioids and benzodiazepines concurrently whenever possible. Clinicians should communicate with others managing the patient to discuss the patient’s needs, prioritize patient goals, weigh risks of concurrent benzodiazepine and opioid exposure, and coordinate care.
- Clinicians should calculate the total MME/d for concurrent opioid prescriptions. If patients are found to be receiving high total daily dosages of opioids, clinicians should discuss their safety concerns with the patient, consider tapering to a safer dosage, and consider offering naloxone.
- Clinicians should discuss safety concerns with other clinicians who are prescribing controlled substances for their patient.
- Clinicians should consider the possibility of a substance use disorder and discuss concerns with their patient.
- If clinicians suspect their patient might be sharing or selling opioids and not taking them, clinicians should consider urine drug testing to assist in determining whether opioids can be discontinued without causing withdrawal. A negative drug test for prescribed opioids might indicate the patient is not taking prescribed opioids, although clinicians should consider other possible reasons for this test result.

Clinicians should not dismiss patients from their practice on the basis of PDMP information. Doing so could result in missed opportunities to provide potentially lifesaving information and interventions.

**10. When prescribing opioids for chronic pain, clinicians should use urine drug testing before starting opioid therapy and consider urine drug testing at least annually to assess for prescribed medications as well as other controlled prescription drugs and illicit drugs.** (Recommendation category: B; evidence type: 4)

Prior to starting opioids for chronic pain and periodically during opioid therapy, clinicians should use urine drug testing to assess for prescribed opioids as well as other controlled substances and illicit drugs that increase risk for overdose when combined with opioids, including nonprescribed opioids, benzodiazepines, and heroin.

In most situations, initial urine drug testing can be performed with a relatively inexpensive immunoassay panel for commonly prescribed opioids and illicit drugs. Patients prescribed less commonly used opioids might require specific testing for those agents. The use of confirmatory testing adds substantial costs and should be based on the need to detect specific opioids that cannot be identified on standard immunoassays or on the presence of unexpected urine drug test results. In addition, clinicians should not test for substances for which results would not affect patient management or for which implications for patient management are unclear. Clinicians should be familiar with the drugs included in urine drug testing panels used in their practice and should understand how to interpret results for these drugs.

Before ordering urine drug testing, clinicians should explain to patients that testing is intended to improve their safety, should explain expected results (eg, presence of prescribed medication and absence of drugs, including illicit drugs, not reported by the patient), and should ask patients whether there might be unexpected results. Clinicians should discuss unexpected results with the local laboratory or toxicologist and with the patient. Discussion with patients prior to specific confirmatory testing can sometimes yield a candid explanation of why a particular substance is present or absent and obviate the need for expensive confirmatory testing on that visit. If unexpected results are not explained, a confirmatory test using a method selective enough to differentiate specific opioids and metabolites (eg, gas or liquid chromatography/mass spectrometry) might be warranted to clarify the situation.

Clinicians should not dismiss patients from care based on a urine drug test result. This could have adverse consequences for patient safety, including missed opportunities to facilitate treatment for substance use disorder.

**11. Clinicians should avoid prescribing opioid pain medication and benzodiazepines concurrently whenever possible.** (Recommendation category: A; evidence type: 3)

Although there are circumstances when it might be appropriate to prescribe opioids to a patient receiving benzodiazepines (eg, severe acute pain in a patient taking long-term, stable low-dose benzodiazepine therapy), clinicians should avoid prescribing opioids and benzodiazepines concurrently whenever possible. In addition, given that other central nervous system depressants (eg, muscle relaxants, hypnotics) can potentiate central nervous system depression associated with opioids, clinicians should consider whether benefits outweigh risks of concurrent use of these drugs. Clinicians should check the PDMP for concurrent controlled medications prescribed by other clinicians and should consider involving pharmacists and pain specialists as part of the management team when opioids are co-prescribed with other central nervous system depressants. When patients require tapering of benzodiazepines or opioids to reduce risk of fatal respiratory depression, it might be safer and more practical to taper opioids first. Clinicians should taper benzodiazepines gradually if discontinued because abrupt withdrawal can be associated with rebound anxiety, hallucinations, seizures, delirium tremens, and, in rare cases, death. If benzodiazepines prescribed for anxiety are tapered or discontinued, evidence-based psychotherapies (eg, CBT) and specific antidepressants or other nonbenzodiazepine medications approved for anxiety should be offered. Clini-
Clinicians should communicate with mental health professionals managing the patient to discuss the patient's needs, prioritize patient goals, weigh risks of concurrent benzodiazepine and opioid exposure, and coordinate care.

12. Clinicians should offer or arrange evidence-based treatment (usually medication-assisted treatment with buprenorphine or methadone in combination with behavioral therapies) for patients with opioid use disorder. (Recommendation category: A; evidence type: 2)

If clinicians suspect opioid use disorder based on patient concerns or behaviors or on findings in PDMP data or from urine drug testing, they should discuss their concerns with their patient and provide an opportunity for the patient to disclose related concerns or problems. Clinicians should assess for opioid use disorder using DSM-5 criteria. Clinicians should offer or arrange for patients with opioid use disorder to receive evidence-based treatment (usually medication-assisted treatment with buprenorphine or methadone maintenance therapy in combination with behavioral therapies). Oral or long-acting injectable naltrexone can also be used in nonpregnant adults. For pregnant women with opioid use disorder, medication-assisted therapy with buprenorphine (without naloxone) or methadone has been associated with improved maternal outcomes and should be offered.

Physicians prescribing opioids in communities without sufficient treatment capacity for opioid use disorder should strongly consider obtaining a waiver from the Substance Abuse and Mental Health Services Administration (SAMHSA) that allows them to prescribe buprenorphine to treat patients with opioid use disorder. Clinicians do not need a waiver to offer naltrexone for opioid use disorder as part of their practice. Clinicians unable to provide treatment themselves should arrange for patients with opioid use disorder to receive care from a substance use disorder treatment specialist, such as an office-based clinician who prescribes buprenorphine or naltrexone treatment, or from an opioid treatment program certified by SAMHSA to provide supervised medication-assisted treatment for patients with opioid use disorder.

Discussion

The evidence review focused on 5 key questions (Box 2) that have resulted in 12 recommendations (Box 5) in 3 areas: determining when to initiate or continue opioids for chronic pain; opioid selection, dosage, duration, follow-up, and discontinuation; and assessing risk and addressing harms of opioid use. The objective of these recommendations is to provide information about opioid prescribing for primary care clinicians treating adult patients with chronic pain.

Of primary importance, nonopioid therapy is preferred for treatment of chronic pain. Opioids should be used only when benefits for pain and function are expected to outweigh risks. Before starting opioids, clinicians should establish treatment goals with patients and consider how opioids will be discontinued if benefits do not outweigh risks. When opioids are used, clinicians should prescribe the lowest effective dosage, carefully reassess benefits and risks when considering increasing dosage to 50 MME or more per day, and avoid concurrent opioids and benzodiazepines whenever possible. Clinicians should evaluate benefits and harms of continued opioid therapy with patients every 3 months or more frequently and review prescription drug monitoring program data, when available, for high-risk combinations or dosages. For patients with opioid use disorder, clinicians should offer or arrange evidence-based treatment, such as medication-assisted treatment with buprenorphine or methadone.

Clinical guidelines complement other strategies such as strengthening the evidence base for pain prevention and treatment, reducing disparities in pain treatment, improving service delivery and reimbursement, and supporting professional and public education. To aid the application of the guideline in clinical practice, CDC is translating the guideline into user-friendly materials, such as a checklist decision aid (eFigure in the Supplement), fact sheets (available at http://www.cdc.gov/drugoverdose/prescribing/resources.html), and a mobile application. CDC will also work with partners to support clinician education on pain management options, opioid therapy, and risk mitigation strategies. Efforts that might enhance implementation of recommended practices include development of quality improvement measures, implementing clinical decision support, and integrating initiatives to promote safer prescribing within insurance plans. In addition, policy initiatives that address barriers to implementation of the guideline, such as increasing accessibility of PDMP data, e-prescribing, and availability of clinicians who can offer medication-assisted treatment for opioid use disorder are strategies to consider to enhance implementation of the recommended practices. CDC will work with federal partners and payers to evaluate strategies such as payment reform and health care delivery models that could improve patient health and safety. For example, strategies might include strengthened coverage for nonpharmacologic treatments, appropriate urine drug testing, and medication-assisted treatment; reimbursable time for patient counseling; and payment models that improve access to interdisciplinary, coordinated care.

The CDC guideline provides recommendations that are based on best available evidence, interpreted and informed by expert opinion. Evidence informing the recommendations is based on observational studies or randomized clinical trials with notable limitations, as well as clinical experience and observations, characterized as low in quality under GRADE methodology. As highlighted by a National Institutes of Health expert panel, “evidence is insufficient for every clinical decision that a provider needs to make about the use of opioids for chronic pain.” The expert panel recommended that research is needed to improve current understanding of which types of pain, specific diseases, and patients are most likely to be associated with benefit and harm from opioid pain medications; evaluate and estimate cost-benefit of multidisciplinary pain interventions; develop and validate tools for identification of patient risk and outcomes; assess the effectiveness and harms of opioid pain medications with alternative study designs; and investigate risk identification and mitigation strategies and their effects on patient and public health outcomes.

To inform future guideline development, more research is needed to fill critical evidence gaps. Yet given that chronic pain is a significant public health problem, the risks associated with long-term opioid therapy, the availability of effective alternative treatment options for pain, and the potential for improvement in the quality of health care with the implementation of recommended practices, a guideline for prescribing is warranted with currently available evidence. The balance between benefits and harms of long-
term opioid therapy for chronic pain based on both clinical and contextual evidence is sufficiently clear to support the issuance of category A recommendations in most cases.

Conclusions

The guideline is intended to improve communication between clinicians and patients about the risks and benefits of opioid therapy for chronic pain, improve the safety and effectiveness of pain treatment, and reduce the risks associated with long-term opioid therapy, including opioid use disorder, overdose, and death. CDC is committed to evaluating the guideline to identify effects on clinician and patient outcomes, both intended and unintended, and will revisit the guideline to determine if evidence gaps have been sufficiently addressed to warrant an update of the guideline and revise the recommendations in future updates when warranted.

REFERENCES


ARTICLE INFORMATION

Published Online: March 15, 2016. doi:10.1001/jama.2016.3464.

Author Contributions: Drs Dowell and Chou had full access to all of the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Dowell, Haegerich.

Acquisition, analysis, or interpretation of data: Dowell, Haegerich, Chou.

Drafting of the manuscript: Dowell, Haegerich, Chou.

Critical revision of the manuscript for important intellectual content: Dowell, Haegerich, Chou.

Statistical analysis: Chou.

Administrative, technical, or material support: Haegerich.

Study supervision: Dowell, Haegerich.

Conflict of Interest Disclosures: All authors have completed and submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Drs Dowell and Haegerich are employees of the Centers for Disease Control and Prevention. Dr Chou was supported under contract through a deal at CDC. No other disclosures were reported.

Funding/Support: The Centers for Disease Control and Prevention (CDC) supported the development of the guideline. Dr Chou’s activities were supported through a short-term deal under contract at CDC (515PA-I505457B). The clinical evidence review was updated based on a previously published 2014 report funded by the Agency for Healthcare Research and Quality (AHRQ) under contract to the Pacific Northwest Evidence-based Practice Center (contract 290-2012-00014-I). Abt Associates collected, managed, analyzed, and interpreted data in the contextual evidence review, funded through a contract (290-2013-M-53890; task order 260-2015-F-62036) supported by CDC.

Role of the Funder/Sponsor: CDC conducted the full guideline development process, directing the design and conduct of the systematic reviews; collection, management, analysis, and interpretation of the data; and preparation, review, and approval for submission of the manuscript for publication. CDC staff members were responsible for the overall design and conduct of the guideline and preparation, review, and approval of the manuscript.

Core Expert Group: Pam Archer, MPH, Oklahoma State Department of Health; Jane Ballantyne, MD, University of Washington (retired); Amy Bohnert, MHS, PhD, University of Michigan; Ronnie Burman, ScD, Ohio Department on Aging; Roger Chou, on deal to CDC under contract; Phillipp Coffin, MD, MIA, University of California, San Francisco; Gary Franklin, MD, MPH, Washington State Department of Labor and Industries/University of Washington; Erin Krebs, MD, MPH, Minneapolis VA Health Care System/University of Minnesota; Mitchel Mutter, MD, Tennessee Department of Health; Lewis Nelson, MD, New York University School of Medicine; Trupti Patel, MD, Arizona Department of Health Services; Christina A. Porucznik, PhD, MSPH, University of Utah; Robert “Chuck” Rich, MD, American Academy of Family Physicians; Joanna Starrels, MD, MS, Albert Einstein College of Medicine of Yeshiva University; Michael Steinman, MD, Society of General Internal Medicine; Thomas Tape, MD, American College of Physicians; Judith Turner, PhD, University of Washington. No compensation was received for serving as a member of the Core Expert Group. Disclosures may be found in the primary MMWR publication.

Stakeholder Review Group: American Academy of Neurology, John Markman, MD; American Academy of Pain Management, Bob Twillman, PhD; American Academy of Pain Medicine, Edward C. Covington, MD; American Academy of Pediatrics, Roger F. Suchyta, MD; American Academy of Physical Medicine and Rehabilitation, Kavitha V. Neerukonda, JD, MHA; American Cancer Society Cancer Action Network, Mark Fleury, PhD; American Chronic Pain Association, Penney Cowan; American College of Medical Toxicology, David Jaurinlink, BPharm, MD, PhD; American College of Obstetricians and Gynecologists, Gerald “Jerry” F. Joseph Jr, MD; American Geriatrics Society, Bruce Ferrell, MD, and M. Carrington Reid, MD, PhD; American Hospital Association, Ashley Thompson; American Medical Association, Barry D. Dickinson, PhD; American Pain Society, Gregory Terman, MD, PhD; American Society of Addiction Medicine, Beth Haynes, MPPA; American Society of Anesthesiologists, Asokumar Buvanendran, MD; American Society of Hematology, Robert M. Plovnick, MD, MS; American Society of Interventional Pain Physicians, Sanford M. Silverman, MD; Physicians for Responsible Opioid Prescribing, Andrew Kolody, MD. No compensation was received for serving as a member of the Stakeholder Review Group.

Opioid Guideline Workgroup: Chair: Christina Porucznik, PhD, MSPH; Workgroup members: Anne Burns, RPh; Penney Cowan; Chizao Cunningham, MD, MS; Katherine Galluzzo, DO; Traci Green, PhD, MSC; Mitchell Katz, MD; Erin Krebs, MD, MPH; Gregory Terman, MD, PhD; Mark Wallace, MD. Workgroup consultants: Roger Chou, MD; Edward Covington; Diana Eppolito; Michael Greene, MD; Steven Stanos, DO. Disclosures may be found in the primary MMWR publication.

National Center for Injury Prevention and Control Board of Scientific Counselors: Chair: Stephen Hargarten, MD, MPH; Members: John Allegrante, PhD; Joan Marie Duwwe, MD, Samuel Forjou, MD, MPH, DrPH, FCCP; Gerard Gioia, PhD; Deborah Gorman-Smith, PhD; Traci Green, PhD; Sherry Lynne Hamby, PhD; Robert Johnson, MD; Angela Mickalide, PhD, MCHES; Sherry Molock, PhD; Christina Porucznik, PhD, MSPH; Jay Silverman, PhD; Maria Testa, PhD; Shelly Timmons, MD, PhD, Ex officio members: Melissa Brodowski, PhD; Dawn Castillo, MPH; Wilson Compton, MD, MPE, Elizabeth Edgerton, MD, MPH; Thomas Feucht, PhD; Meredith Fox, PhD; Holly Hedegaard, MD, MSPH; John Howard, MD; Lyndon Joseph, PhD; Jinhee Lee, PharmD; Iris Mabry-Hernandez, MD, MPH; Valeri Mahalomes, PhD; Angela Moore Parmyler, PhD; Thomas Schroeder, MS. Disclosures may be found in the primary MMWR publication.

Additional Contributions: We acknowledge Jeanmarie Perrone, MD, Matthew Bair, MD, and David Taube, MD, for conducting initial peer reviews of the guideline for the CDC prior to journal submission; peer reviewers were not compensated for their contributions. We acknowledge Don Teater, MD, for facilitating the Core Expert Group. We acknowledge the work that the medical writers, editors, and reviewers from Ariande Labs provided to produce the checklist for prescribing opioids for chronic pain.

American Academy of Pain Medicine and Rehabilitation, Kavitha V. Neerukonda, JD, MHA; American Cancer Society Cancer Action Network, Mark Fleury, PhD; American Chronic Pain Association, Penney Cowan; American College of Medical Toxicology, David Jaurinlink, BPharm, MD, PhD; American College of Obstetricians and Gynecologists, Gerald “Jerry” F. Joseph Jr, MD; American Geriatrics Society, Bruce Ferrell, MD, and M. Carrington Reid, MD, PhD; American Hospital Association, Ashley Thompson; American Medical Association, Barry D. Dickinson, PhD; American Pain Society, Gregory Terman, MD, PhD; American Society of Addiction Medicine, Beth Haynes, MPPA; American Society of Anesthesiologists, Asokumar Buvanendran, MD; American Society of Hematology, Robert M. Plovnick, MD, MS; American Society of Interventional Pain Physicians, Sanford M. Silverman, MD; Physicians for Responsible Opioid Prescribing, Andrew Kolody, MD. No compensation was received for serving as a member of the Stakeholder Review Group.

Opioid Guideline Workgroup: Chair: Christina Porucznik, PhD, MSPH; Workgroup members: Anne Burns, RPh; Penney Cowan; Chizao Cunningham, MD, MS; Katherine Galluzzo, DO; Traci Green, PhD, MSC; Mitchell Katz, MD; Erin Krebs, MD, MPH; Gregory Terman, MD, PhD; Mark Wallace, MD. Workgroup consultants: Roger Chou, MD; Edward Covington; Diana Eppolito; Michael Greene, MD; Steven Stanos, DO. Disclosures may be found in the primary MMWR publication.

National Center for Injury Prevention and Control Board of Scientific Counselors: Chair: Stephen Hargarten, MD, MPH; Members: John Allegrante, PhD; Joan Marie Duwwe, MD, Samuel Forjou, MD, MPH, DrPH, FCCP; Gerard Gioia, PhD; Deborah Gorman-Smith, PhD; Traci Green, PhD; Sherry Lynne Hamby, PhD; Robert Johnson, MD; Angela Mickalide, PhD, MCHES; Sherry Molock, PhD; Christina Porucznik, PhD, MSPH; Jay Silverman, PhD; Maria Testa, PhD; Shelly Timmons, MD, PhD, Ex officio members: Melissa Brodowski, PhD; Dawn Castillo, MPH; Wilson Compton, MD, MPE, Elizabeth Edgerton, MD, MPH; Thomas Feucht, PhD; Meredith Fox, PhD; Holly Hedegaard, MD, MSPH; John Howard, MD; Lyndon Joseph, PhD; Jinhee Lee, PharmD; Iris Mabry-Hernandez, MD, MPH; Valeri Mahalomes, PhD; Angela Moore Parmyler, PhD; Thomas Schroeder, MS. Disclosures may be found in the primary MMWR publication.


jama.com


