



Review

The downward spiral of chronic pain, prescription opioid misuse, and addiction: Cognitive, affective, and neuropsychopharmacologic pathways



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ABSTRACT

Prescription opioid misuse and addiction among chronic pain patients are emerging public health concerns of considerable significance. Estimates suggest that more than 10% of chronic pain patients misuse opioid analgesics, and the number of fatalities related to nonmedical or inappropriate use of prescription opioids is climbing. Because the prevalence and adverse consequences of this threat are increasing, there is a pressing need for research that identifies the biobehavioral risk chain linking chronic pain, opioid analgesia, and addictive behaviors. To that end, the current manuscript draws upon current neuropsychopharmacologic research to provide a conceptual framework of the downward spiral leading to prescription opioid misuse and addiction among chronic pain patients receiving opioid analgesic pharmacotherapy. Addictive use of opioids is described as the outcome of a cycle initiated by chronic pain and negative affect and reinforced by opioidergic-dopaminergic interactions, leading to attentional hypervigilance for pain and drug cues, dysfunctional connectivity between self-referential and cognitive control networks in the brain, and allostatic dysregulation of stress and reward circuitry. Implications for clinical practice are discussed; multimodal, mindfulness-oriented treatment is introduced as a potentially effective approach to disrupting the downward spiral and facilitating recovery from chronic pain and opioid addiction.

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1. Introduction

1.1. The scope of chronic pain and prescription opioid misuse

Misuse of prescription opioid analgesics is an emerging public health concern that confers significant risks for overdose, unsafe drug interactions, and the full panoply of adverse social, legal, and adaptive consequences associated with dependence on any psychoactive drug. Though the prevalence of prescription opioid misuse across the general U.S. population has increased more than threefold over the past two decades (Hall et al., 2008), it is presently unclear to what extent this explosion in prevalence reflects an increase in opioid misuse by persons without a prescription for opioids versus an increase in nonmedical use of opioids by persons prescribed opioids for analgesia. This distinction notwithstanding, nationally representative surveys indicate that prescription opioid misuse is endemic among U.S. adolescents and adults; prescription opioids are now among the most commonly misused drugs in the U.S. (Wilson, 2007). For instance, results of the National Survey on Drug Use and Health (NSDUH) for 18–25 year-olds revealed that 23.8%, 11.1%, and 4.8% reported lifetime, past year, and past month misuse of prescription “pain killers” (Substance Abuse and Mental Health Services Administration, 2011). NSDUH findings further indicated that rates of initiation of nonmedical pain reliever use were second only to those of marijuana, with more than two million persons initiating nonmedical use of opioid analgesics annually. These findings underscore the pervasive availability and misuse of these agents in the U.S. The ready accessibility, prevalent misuse, and euphorogenic effects of prescription opioids would seem to create conditions for widespread dependence on these agents (Mendelson et al., 2008). Recent research does, in fact, suggest that rates of publically funded chemical dependency treatment and emergency department care for prescription opioid use have increased dramatically in recent years and it is estimated that well over a million Americans are currently dependent on prescription opioids (Manubay et al., 2011; Mendelson et al., 2008). As previously underscored, these statistics do not clearly differentiate opioid misusers with and without chronic pain; yet, this is a critically important distinction with serious clinical ramifications.

1.2. Diagnostic classification of opioid use patterns

The issue of prescription opioid misuse is further complicated by the diversity of nosological categories used to classify opioid use patterns and their biopsychosocial consequences. As defined by *Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition* (DSM-IV), *prescription opioid abuse* involves a maladaptive pattern of repeated opioid use that: results in failure to fulfill social, occupational, academic, or familial obligations; continues despite recurrent legal problems related to opioid use; persists in spite of interpersonal problems caused or exacerbated by opioids; and occurs in physically hazardous contexts. In contrast, DSM-IV defines *prescription opioid dependence* as involving physical symptoms of dependence, including tolerance and withdrawal, as well as behavioral symptoms including: taking higher doses than intended; an inability to reduce or stop taking opioids; spending

substantial amounts of time using, obtaining, recovering, or thinking about opioids; and continued use in spite of adverse physical or psychological consequences (Zacny et al., 2003). However, some clinicians believe these behavioral criteria for abuse and dependence are inappropriate for opioid-using chronic pain patients, because patients who take opioids as prescribed may be unable to reduce their opioid use or may continue opioid use despite adverse health consequences due to the intractability of their chronic pain. Instead, many pain and addiction specialists use the American Pain Society (2002) criteria to identify *prescription opioid addiction* among chronic pain patients, including symptoms of impaired control over opioid use, compulsive opioid use, continued use despite harm, and craving for opioids (Sullivan et al., 2008; Wilson, 2007). Addictive tendencies among prescription opioid users may be pre-saged by the presence of *opioid misuse* behaviors such as selling medication or injecting oral formulas (Ives et al., 2006; Sullivan et al., 2008), although less serious forms of opioid misuse like unauthorized dose escalation are relatively common among under-treated chronic pain patients and may not indicate opioid addiction. The presence of more serious misuse behaviors may mark the transition from sanctioned use of opioids to development of opioid use disorders and addiction (Butler et al., 2007).

1.3. Epidemiology of prescription opioid misuse

Parsing apart these conditions, structured psychiatric interviews conducted for the National Epidemiologic Survey on Alcohol and Related Conditions (NESARC) study of more than 43,000 U.S. adults identified 4.7% of respondents as lifetime prescription opioid misusers; whereas 1.4% of respondents met criteria for a DSM-IV prescription opioid use disorder (opioid abuse or dependence; Huang et al., 2006). Men, adults with Axis I and Axis II DSM-IV diagnoses, respondents residing in the West, and young or middle aged adults were at greatest risk for prescription drug misuse and a prescription opioid use disorder. Other studies have identified a history of alcohol or illicit drug misuse, anxiety, depression, and chronic pain as significant risk factors for prescription opioid misuse and dependence (Amari et al., 2011; Pohl and Smith, 2012; Turk et al., 2008). Individuals suffering from chronic pain disorders, who are at risk for becoming physically dependent on opioid analgesics when adhering to their prescribed medication regimen, may be particularly vulnerable to prescription opioid misuse (Butler et al., 2007; Ives et al., 2006; Sullivan et al., 2010).

Though the prevalence of prescription opioid misuse and addiction among chronic pain patients has yet to be firmly established by nationally representative, population-level surveys, chronic pain itself is highly prevalent in modern society; a meta-analysis of 13 studies reported a weighted mean prevalence of 35.5% for chronic pain of any kind and a weighted mean prevalence of 11% for severe chronic pain (Ospina and Harstall, 2002). Many patients with chronic pain have serious medical conditions that require long-term opioid pharmacotherapy, and a subset of these individuals are at significant risk for escalating from appropriate opioid use to misuse and finally to opioid addiction. The best available prevalence estimates for opioid misuse and addiction among chronic pain patients may be derived from research by

Fishbain et al. (2007), who reviewed 67 methodologically rigorous studies including thousands of chronic pain patients and concluded that 3.3%, 11.5%, and 14.5% of these patients, in toto, became addicted to prescription opioids, engaged in opioid misuse behaviors such as unauthorized dose escalation or drug hoarding, and used illicit drugs, respectively. Thus, while prescription opioid addiction among chronic pain patients appears to be relatively rare, opioid misuse in this population is more common. Although relatively few studies have prospectively examined factors related to increased risk for prescription drug misuse among chronic pain patients receiving ongoing opioid pharmacotherapy, recent reports suggest that greater baseline pain intensity (Edwards et al., 2011; Jamison et al., 2009), psychological distress and behavioral problems (Jamison et al., 2010), and status as a cigarette smoker (Novy et al., 2012) predict greater risk for transitions to prescription opioid misuse, abuse and addiction. There is a pressing need for additional studies to better understand the risk factors for progression to opioid misuse and opioid use disorders among chronic pain patients receiving prescription opioid pharmacotherapy (Larance et al., 2011). Further, novel conceptual frameworks are needed to delineate the biobehavioral risk chain linking pain, opioid analgesia, opioid misuse, and addiction.

The purpose of this paper is to describe how the neuropharmacologic properties of prescription opioids interact with cognitive, affective, and physiological factors implicated in chronic pain and addictive behavior. We first review the neurobiology of opioid agents and pain processing in the human nervous system. Next, we present a conceptual model to describe how chronic pain, affective dysregulation, and opioid use interact to potentiate each other and foster opioid addiction. Lastly, we discuss clinical implications of the model for psychological treatment of chronic pain patients and for the prevention of opioid misuse in these populations.

2. Neuropharmacologic effects of acute opioid administration on central nervous system function

2.1. Effects of opioids on neurotransmission

Opioidergic neurotransmission is necessary for the biologic integrity of the human nervous system. Both opioid medication and endogenous opioids (those naturally produced by the body such as beta-endorphin and enkephalins) interact with *mu*, *kappa* and *delta* opioid receptor-proteins on neuronal membranes widely distributed throughout the cerebral cortex, thalamus, hypothalamus, periaqueductal grey, interpeduncular median raphe nuclei, and spinal cord (Arvidsson et al., 1995; Lewis et al., 1983; Mathieu-Kia et al., 2001; Pan and Pasternak, 2011; Traynor and Wood, 1987). While *mu* receptor activation mediates the analgesic properties of selective *mu* opioid agonists like morphine, codeine, heroin, meperidine, methadone, and hydromorphone (Becker et al., 2012; Fields, 2011; Julien, 2007), the abuse liability of prescription opiate medications is thought to be due to the interactive effects of *mu* receptor activation on dopaminergic transmission. Acute administration of *mu* receptor agonists indirectly excites dopamine-secreting neurons in the ventral tegmental area via GABAergic mechanisms (Chiara and North, 1992; Johnson and North, 1992). Indirect action of opioids on dopamine neurotransmission appears to be integral to the reinforcing and rewarding properties of opioids, as well as opioid addiction (Berridge et al., 2009; Le Merrer et al., 2009; Volkow et al., 2011).

2.2. Effects of opioids on neurocognition

Functional magnetic resonance imaging (fMRI) has provided insight into the effects of opioid administration on human

brain function. Single-dose opioid administration in opioid-naïve, healthy individuals activates brain regions involved in processing reward (nucleus accumbens, extended amygdala, orbitalfrontal gyrus, and hippocampus) (Becerra et al., 2006; Wanigasekera et al., 2012), while decreasing brain activation in brain regions involved in cognitive control and attention (dorsolateral prefrontal cortex, anterior cingulate cortex, inferior parietal lobe) (Becerra et al., 2006; Jastrzab et al., 2012; Lee et al., 2012). Opioid-induced hyperactivation of limbic circuitry is similar to that elicited by psychostimulants (Breiter et al., 1997), whereas opioid-induced cortical inhibition is comparable to that elicited by sedative or anesthetic drugs (Fiset et al., 2005). Acute administration of higher concentrations of opioids reduces neural activity in the brain stem and medial thalamic nuclei, which can lead to diminished sensory, motor, and cognitive functioning to the point of unconsciousness (Becerra et al., 2006). Other neuroimaging research, involving single-dose opioid administration in the absence of pain to healthy individuals, revealed activations in orbitofrontal cortex, anterior cingulate cortex, insula, and amygdala that were associated with the time course of subjective reports of dizziness, drowsiness, and euphoria (Leppä et al., 2006). Consistent with prevailing models of drug addiction, these findings suggest that high-risk individuals may start along the path of addiction to prescription opioids after experiencing an initial and transient hyperactivation of reward circuitry coupled with disruption in neural circuitry subserving cognitive function.

Disrupted neural circuitry function in cognitive control networks is evident in a host of studies indicating cognitive and psychomotor impairments such as decrements in working memory, reduced cognitive flexibility, and increased impulsivity in long-term opioid users (Baldacchino et al., 2012; Mintzer et al., 2005; Prosser et al., 2009; Verdejo et al., 2005; Zacny, 1995) and healthy, opioid-naïve individuals (Baldacchino et al., 2012; Chierri et al., 2009; Schneider et al., 1999; Walker and Zacny, 1999; Zacny, 1995). Deactivation in brain regions involved in executive function (e.g., anterior cingulate cortex, dorsolateral prefrontal cortex) following acute, single-dose opioid administration provides further mechanistic insight into the relationship between opioid use and cognitive impairment (Becerra et al., 2006; Jastrzab et al., 2012; Lee et al., 2012; Prosser et al., 2009). Opioid-induced deficits in executive function, compounded with those associated with chronic pain (Kendall et al., 2010; Tassain et al., 2003), may compromise the patient's ability to exert cognitive control needed to cope through non-pharmacologic means, thereby inadvertently promoting dependence on opioids as a means of obtaining relief from pain.

3. Pain and the nociceptive and affective dimensions of opioid-induced analgesia

3.1. Neurophysiology of pain

Pain is a complex, biopsychosocial experience that arises from the interaction between sensory, cognitive, and affective factors. Acute pain is most often induced by noxious stimulation, tissue damage, and/or disease and is a beneficial process that helps to preserve the morphological integrity of the organism by motivating adaptive behavior. Acute pain is associated with activation in a widely distributed network of highly connected brain regions including primary and secondary somatosensory cortices, bilateral insula, thalamus, anterior cingulate cortex, and the prefrontal cortex (Coghill et al., 2003). Chronic pain, on the other hand, may not be clearly linked to an observable physiological or neural pathology. In some cases, the CNS may generate pain in the absence of input from peripheral or visceral nociceptors, as evidenced by conditions like phantom limb pain, irritable bowel syndrome, and chronic pain

due to CNS changes secondary to stroke (Loeser and Melzack, 1999; Melzack, 1999; Spiller et al., 2007). In such cases, the intensity of chronic pain is only weakly or not at all associated with the degree of actual tissue damage (Waddell et al., 1980; Waddell, 1996). With time, pain may become the output of a pattern of familiar sensory inputs triggering cognitive and affective reactions that have previously been tied to the pain experience (Melzack, 1999). In this way, even the anticipation of pain and suffering may result in somatic discomfort or functional disability of a similar or even greater magnitude as the actual painful event (Crombez et al., 1999). Chronic pain may be exacerbated by stress, environmental variables, emotional states, and other psychological factors (Garland, 2012; Loeser and Melzack, 1999; Voscopoulos and Lema, 2010).

3.2. A role for default mode networks in pain processing

Although in some cases chronic pain stems from an initial painful insult to the body, its progression is associated with the interaction between original injury, CNS, and cognitive-affective dysregulation (Apkarian et al., 2005). With regard to the latter factor, recent neuroimaging research has identified a role for self-referential thought processes in chronic pain (Baliki et al., 2006; Loggia et al., 2012; Wasan et al., 2011). In healthy individuals, self-referential thought (i.e., thought pertaining to an individual's autobiographical narrative or life story) is believed to be reflected by oscillating activity in a distinct network of brain regions (posterior cingulate cortex/precuneus, medial prefrontal cortex, and the inferior and lateral temporal cortices) referred to as the default mode network (DMN) (Raichle et al., 2001). Recent findings from resting-state functional connectivity analyses of regional brain signals reveal that chronic pain patients exhibit abnormal functional connections between the default mode network and the anterior insula (Loggia et al., 2012), a region crucially involved in evaluating sensory processes. It has been postulated that the experience of chronic pain is related to abnormal default mode processes where pain experience becomes the primary object of self-referential thought (Zeidan and Coghill, 2012). Dysregulated DMN connectivity may underpin the development of the entrapped sense of identity and autobiographical narrative dominated by pain-related memories and associations so common to chronic pain patients (Hellström, 2001; Morley et al., 2005; Pincus and Morley, 2001). The relationship between DMN and chronic pain is also supported by findings demonstrating the role of the perigenual anterior cingulate cortex, a brain region involved in mediating the cognitive modulation of pain (Vogt, 2005). Recent findings reveal that chronic pain patients' self-reports of pain are negatively associated with connectivity between the perigenual anterior cingulate cortex and the DMN (Loggia et al., 2012; Wasan et al., 2011). That is, individuals who report more intense chronic pain exhibit decreased connectivity between the perigenual anterior cingulate and the DMN, likely reflecting reduced ability to govern pain-related thought processes. Conversely, increased functional connectivity between anterior cingulate and DMN regions may reflect a process for attenuating pain.

3.3. Descending nociceptive inhibition and opioid analgesia

Thus, the brain does not passively receive nociceptive information from the body, but instead actively regulates nociception by way of interactions between descending pain modulatory system (Heinricher et al., 2009; Melzack and Wall, 1965; Reynolds, 1969) and cortico-cortical networks (Rainville, 2002). The descending pain modulatory system exerts influences on nociceptive input from the spinal cord through a network of cortical, subcortical, and brainstem structures including prefrontal cortex, anterior cingulate cortex, insula, amygdala, hypothalamus,

periaqueductal grey, rostral ventromedial medulla, and dorso-lateral pons (Tracey and Mantyh, 2007). The descending pain modulatory system has been construed as the means by which the CNS inhibits nociceptive signals at the spinal outputs (Heinricher et al., 2009). Endogenous and exogenous opioids relieve pain by targeting the descending pain modulatory system (Besson, 1999), most notably in the periaqueductal grey, a brain region involved in processing the placebo-analgesia (Tracey, 2010). In addition, acute, single-dose administration of opioids in healthy individuals exerts direct analgesic effects by reducing sensory evaluation processes evidenced by reduced activation in brain regions corresponding to processing lower-level afferent processes (i.e., primary and secondary somatosensory cortex, thalamus) (Wagner et al., 2007; Wise et al., 2002), and by modulating neurotransmission in the substantia gelatinosa of the dorsal horn of the spine (Le Bars et al., 1980; Yaksh, 1981, 1987). Moreover, in a preclinical animal model, acute opioid administration elicited dose-dependent modulation of dopaminergic activity in spinal neurons that mediate nociception (Pappas et al., 2011). Among healthy human volunteers, increasing concentrations of opioids across a single, experimental session monotonically decreased sensory processing in primary and secondary somatosensory cortex and the posterior insula, while dramatically attenuating pain processing in brain regions that mediate affective dimension of pain, such as the amygdala (Oertel et al., 2007).

Given the complexity of pain, it is perhaps not surprising that opioid analgesia operates through both neuropharmacologic and psychological mechanisms. In addition to attenuating sensory aspects of pain (i.e., nociception), opioids may alleviate the affective dimensions of pain (e.g., suffering). In that regard, among healthy individuals, analgesia induced through acute opioid administration operates, in part, through modulating activation of the anterior and posterior cingulate cortex (Wagner et al., 2007), anterior insula (Oertel et al., 2007), and the hypothalamus (Becerra et al., 2006), key nodes in the central autonomic network, a series of neural circuits that regulate attention, emotion, and neurovisceral integration (Thayer and Lane, 2009). Opioids, like all drugs of addiction, also stimulate mesolimbic dopamine reward systems (Chiara and North, 1992; Johnson and North, 1992). Opioid-induced dopamine release in the nucleus accumbens associated with positive mood and reward may promote pain management (Center, 2011). Indeed, pleasure and pain have mutually inhibitory effects through complex opioid-dopamine interactions in cortico-limbic-striatal circuitry (Leknes and Tracey, 2008). Through these and other neural targets, opioids change the attentional and affective response to nociceptive information, and in so doing, temper the emotional averseness and unpleasantness of pain (DelleMijn and Vanneste, 1997). To be clear, much of what is known about the psychobiological mechanisms of opioid-induced analgesia comes from studies of healthy individuals exposed to laboratory pain inductions. Yet, the development of co-occurring chronic pain and opioid use disorders over time may alter the neurobiological response to opioids in clinically significant ways, as detailed below.

4. Neurobiological transitions from opioid use to dependence and addiction

4.1. The neurobiological progression from opioid use to dependence

Opioid therapy for chronic pain often provides effective analgesia but confers significant risk for the development of opioid use disorders in a subset of vulnerable individuals (Denisco et al., 2008; Passik, 2009), as described earlier in the introduction of this paper. Prolonged, medically appropriate opioid use produces physical

dependence symptoms via neuroadaptations resulting in tolerance to opioids, withdrawal when opioids are discontinued, and, in some instances, opioid-induced hyperalgesia (Chu et al., 2008). These physical symptoms of opioid dependence are natural responses to extended use of opioids, and do not signal the presence of an opioid use disorder. Thus, there is a difference between physiological dependence on opioids and the pathology of addiction (Shurman et al., 2010). Addiction arises from a usurpation of normal reward learning mechanisms caused by repeated pharmacologic interruption to normal brain function, resulting in neuroadaptations and fundamental alterations to brain structure. Such neuroplastic changes commence with early use of opioids as associations are forged between drug-related stimuli and reward responses (i.e., transient neuroplasticity), and may eventuate in compulsive, addictive habits as these neurobiological changes become entrenched (i.e., stable neuroplasticity) (Kalivas and O'Brien, 2008).

Opioid-induced durable changes in brain structure and function referred to as neuroplasticity, like the neuroadaptations stimulated by other drugs of addiction, may stem from a cascade of neurobiological events governed by dopaminergic activity in cortico-limbic-striatal circuits (Kalivas and O'Brien, 2008). Acute administration of opioids results in dopamine release from the ventral tegmental area to the prefrontal cortex and nucleus accumbens (Center, 2011; Shippenberg et al., 1993). This dopaminergically mediated process facilitates the learning of drug reward; when opioids result in pain relief, this signals the reward value of opioid-related stimuli (Becker et al., 2012), increasing their incentive salience (Robinson and Berridge, 2008). In other words, the individual learns that opioids and opioid-related cues (e.g., the sight of a pill bottle, a prescription slip, or even the pharmacy where the opioids are dispensed) are motivationally salient, and should be sought after as a means of maintaining a positive hedonic tone. As chronic pain patients engage in recurrent opioid use or misuse, opioid cues may become salient through the pharmacologic reward induced by opioid consumption (positive reinforcement), as well as through the analgesic effects of the drug that remove or allay the aversive experience of pain (negative reinforcement) (Fields, 2004). The incentive salience of opioid cues may increase over time as opioid dependence is established via the process of sensitization of the mesolimbic dopamine system (Robinson and Berridge, 2008). Such reward learning may be coupled with a strong drive towards opioids or a “wanting” that is independent of any “liking” or conscious preference for the drug (Berridge et al., 2009). In fact, many chronic pain patients express a strong dislike of opioids due to their side effects or the stigma associated with opioid use, yet the compulsion to take opioids increases despite this sentiment of disdain.

4.2. Establishment of opioid misuse habits

Over time, this transient neuroplasticity becomes more stable as individuals begin to form unconscious appetitive habits, mediated by structural and functional changes in cortico-limbic-striatal circuits (Yin and Knowlton, 2006). Modified striatal circuitry may govern the implicit neurocognitive operations that direct and impel addictive behaviors and craving states characteristic of disordered forms of substance use, including prescription opioid use disorders (Stacy and Wiers, 2010). Repeated substance use is thought to establish automatic drug-use action schemas (i.e., memory systems that compel and coordinate consumption of the substance through automatized sequences of stimulus-bound, context-dependent behavior, including the biasing of attention towards substance-relevant stimuli) (Garland et al., 2011a; Pierce and Vanderschuren, 2010; Tiffany, 1990). Insofar as cues associated with past drug use are motivationally salient for habitual drug

users, they are able to consequently capture attention, which in turn amplifies their motivational salience (Franken, 2003). This phenomenon, known as addiction attentional bias, is associated with craving (Field et al., 2009), increases drug use (Field and Eastwood, 2005), and predicts relapse (Garland et al., 2012b) among persons with alcohol and illicit drug dependence. A recent study from our laboratory demonstrated that chronic pain patients who met diagnostic criteria for opioid dependence exhibited a significant attentional bias towards prescription opioid cues (e.g., photographs of a pills or pill bottles) (Garland et al., 2012c). Moreover, opioid attentional bias was positively correlated with opioid craving. When external (e.g., the sight of a pill bottle) or internal (e.g., pain, stress) cues associated with past opioid use capture attention, they may trigger dopamine release and activate habitual drug use routines subserved by neuroplastic alterations to cortico-limbic-striatal circuits (Everitt and Robbins, 2005; Kalivas and Volkow, 2005).

4.3. Dysregulation of reward processing and default mode activity in opioid addiction

As addictive automaticity becomes entrenched, neural circuits mediating associations between drug cues and learned appetitive responses are strengthened, while non-drug-related learning is diminished (Hyman, 2005; Kalivas and O'Brien, 2008; Kalivas and Volkow, 2005). Thus, recurrent opioid use may lead to insensitivity in the dopamine system toward natural rewards, like food, sex, or social affiliation (Volkow et al., 2011). Indeed, decreased responsiveness to natural reinforcers has been observed among opiate dependent individuals (Lubman et al., 2008) and is robustly predictive of future opiate consumption (Lubman et al., 2009). Thus, the addictive cycle becomes more and more insidious as both the pain threshold and the ability to experience natural pleasure decreases. Indeed, increased sensitization to pain, when coupled with tolerance to the analgesic effects of opioids, can result in increased opioid craving (Ren et al., 2009) and consumption (Martell et al., 2007). Furthermore, pain itself may decrease responsiveness to natural rewards, resulting in blunted dopaminergic responses to naturally rewarding stimuli and anhedonia (Becker et al., 2012; Leknes and Tracey, 2008). As a result of increasing hyperalgesia and anhedonia, the normative affective balance is tipped towards negativity, further propelling the downward spiral of pain and opioid addiction.

This downward spiral may also crucially involve maladaptive patterns of self-referential thought and aberrant default mode processes as the healthy sense of identity transmogrifies into a self-concept entrapped by pain and entangled with dependence on opioid analgesia. Recent findings provide evidence for dysfunctional default mode-related neural function in opiate addicts, characterized by strong functional connectivity between the nucleus accumbens and prefrontal cortex coupled with attenuated functional connectivity between the prefrontal cortex and anterior cingulate cortex (Ma et al., 2010), a brain structure whose cognitive control functions are significantly impaired in opiate addicts (Yücel et al., 2007). Similarly, chronic pain patients reporting greater pain exhibit decreased connectivity between the default mode network and the anterior cingulate cortex (Loggia et al., 2012). Speculatively, similar changes in self-referential, executive, and reward circuitry might underlie the transition from opioid use to misuse and addiction, as the patient's sense of self and autobiographical narrative becomes more entrenched in the downward spiral of seeking relief from pain through opioids. It is likely that interactions between sensory, cognitive, and self-referential processes fuel the progression toward addictive use of opioids among chronic pain patients.

5. Pain, negative emotion, and opioid addiction

5.1. Emotional modulation of pain

The aversive nature of pain elicits a powerful emotional reaction that feeds back to modulate pain perception. Pain is often accompanied by feelings of anger, sadness and fear depending on how the pain is cognitively appraised. Persistent negative evaluations of noxious sensory events can lead to pain catastrophizing, where the individual interprets uncomfortable or even innocuous somatic sensations as indicating the presence of a serious or mortal threat and consequently underestimate their ability to cope with those respective sensations. These experiences illustrate the importance of cognitive appraisal processes in the construction of the subjective experience of pain. Catastrophic appraisals are largely mediated by fear and produce an intensified pain or allodynia which can result in greater functional disability and reduced quality of life (Crombez et al., 1999; Turner et al., 2002).

Brain regions involved in the processing negative affect are also associated with processing the subjective experience of pain. These areas include executive-level brain regions such as the anterior cingulate and prefrontal cortex and sensory and self-evaluatory regions such as the anterior insula (Craig, 2003; Wiech and Tracey, 2009). Thus, when individuals experience pain-related negative emotions, the heightened neural processing of threat produces a pre-cognitive mental set that primes the individual for an exacerbated subjective experience of pain (De Wied and Verbaten, 2001; Kirwilliam and Derbyshire, 2008; Ploner et al., 2010) and subsequently increases the likelihood that innocuous somatic sensations will be interpreted as painful (Bogaerts et al., 2009; Panerai, 2011; Strigo et al., 2008).

The fear of pain, a clinical feature of chronic pain patients, is associated with hypervigilance for and sustained attention to pain-related stimuli (Haggman et al., 2010; Keogh et al., 2001). Thus, negative affect biases attention toward pain, which then increases its aversive quality. Activation of the neural circuitry subserving negative affect, stress and pain disrupts the function of brain circuits that govern cognitive control (i.e., circuits governed by regions of prefrontal cortex), which may reduce the ability to regulate pain using higher order cognitive coping strategies like reappraisal or viewing the pain as controllable (Arnsten, 2009; Lawrence et al., 2011). Furthermore, stress and negative affect intensify sympathetic nervous system activity associated with pain, manifested in increased anxiety, heart rate, skin conductance responses, and muscle tension that is perceived as painful muscle spasms (Flor et al., 1985; Lundberg et al., 1999; Tousignant-Laflamme and Marchand, 2006). Thus, negative affective states may result from chronic pain and feed back into biobehavioral processes that amplify pain perception (Loggia et al., 2012; Ploner et al., 2010) and exacerbate suffering. Concomitantly, negative affect predicts increased opioid craving among long-term opioid users (Wasan et al., 2012) and is associated with initiation and continued misuse of prescribed opioids (Sullivan et al., 2006). Opioid misusers may use opioids to self-medicate (Khantzian, 1997) the negative affective states and autonomic arousal that cause, co-occur with, or result from pain (Jänig and Habler, 2000; Martenson et al., 2009). In turn, self-medication of negative emotions with opioids may promote the development of opioid addiction.

5.2. Allostasis and the downward spiral of chronic pain and opioid addiction

Negative emotions may contribute to the development of addiction through the progressive dysregulation of stress and reward circuitry in the extended amygdala and altered neurotransmitter function characterized by increased recruitment of other brain

arousal-stress systems (Koob and Le Moal, 2008; Koob, 2009; Smith and Aston-Jones, 2008). These systems are thought to mediate not only stress but also antireward—that is, a homeostatic process whereby brain systems maintain reward within a given limit or set point (Koob and Le Moal, 1997, 2001, 2008). According to Koob and Le Moal's allostatic model of addiction (1997, 2001), use of substances to compensate for dysphoria results in an opponent process that leads to neurobiological sensitization in the extended amygdala and brain arousal-stress systems to threat and harm, while sensitivity to reward is decreased. Thus, in addition, homeostasis is broken, leading to a chronic deviation of brain systems from their normal operating level, termed allostasis. This shift in reward threshold, induced by activation of brain antireward systems, may then elicit increased consumption of psychoactive substances as a means of achieving a hedonic equilibrium. Ironically, this attempt to reach a hedonic state comes with a cost: the continued use of substances further increases the reward set point in the brain, making the individual increasingly insensitive to naturally rewarding experiences while becoming increasingly sensitive to punishment, pain, and other aversive states (Koob and Le Moal, 2008). Thus, allostatic load incurred from prior injury and enhanced by the pharmacological effects of psychoactive substances promotes neuroadaptation to drug effects, modulates sensitization to rewards and punishment, and intensifies negative mood states and pain. In addition to hyperalgesia, prolonged misuse of opioids to medicate negative affective states and induce euphoria may result in *hyperkatifeia*, an increased sensitivity to emotional distress (Shurman et al., 2010). This cyclic process then elicits further substance use as a means of countering dysphoric mood (see Fig. 1). Recurrent use of opioids to maintain positive emotional functioning in the face of stress and pain may result in a shift in the hedonic set point of the brain such that the individual becomes increasingly dependent on opioids to maintain a sense of psychological well-being. In turn, the sense of self may become more entangled with narratives of pain and opioid use.

As the cycle spirals further downward, the experience of pain may cue habitual drug use routines, even when they have been rendered ineffectual due to increased tolerance to opioid analgesia. Although chronic use of opioids may produce limited analgesic effects, once the habit of opioid misuse has been established, it may continue despite countervailing health, social, and legal reasons to adhere to the prescribed medication regimen. Thus, pain comes to trigger the automatic reaction (i.e. taking medication at higher than prescribed doses), and the conscious decision to take the medication is no longer the dominant factor in opioid use. Unconscious habits replace conscious consideration and deliberate evaluation of the level of pain as the primary determinate of opioid dosing. Addictive opioid use may capitalize on non-drug striatal habit mechanisms, which, once established, operate in the absence of pharmacologic reward or drug-induced disruption of normal control (Graybiel, 2008). In other words, whether or not opioid use works to relieve pain (or even produces hyperalgesia), it may persist as an ingrained habit triggered by sensations of pain and external drug-related cues which were originally imbued with motivational salience during the acquisition of the habit of opioid use (Wood and Neal, 2007). This rigid, scripted response to contextual conditions, known as mindlessness (Langer, 1992), may fuel the downward spiral of chronic pain, suffering, and opioid addiction.

When access to opioids is restricted despite the presence of pain and addictive cues, opioid dependent individuals may experience an intensely aversive state, characterized by dysphoria, anxiety, and craving coupled with somatic symptoms of withdrawal that drives an obsessive or intrusive preoccupation with obtaining opioids (Shurman et al., 2010). These experiences promote addictive use of opioids as dependent individuals feel impelled to avoid the discomfort, dysphoria and psychic distress that accompany withdrawal. Compulsive consumption of opioids to satisfy craving and

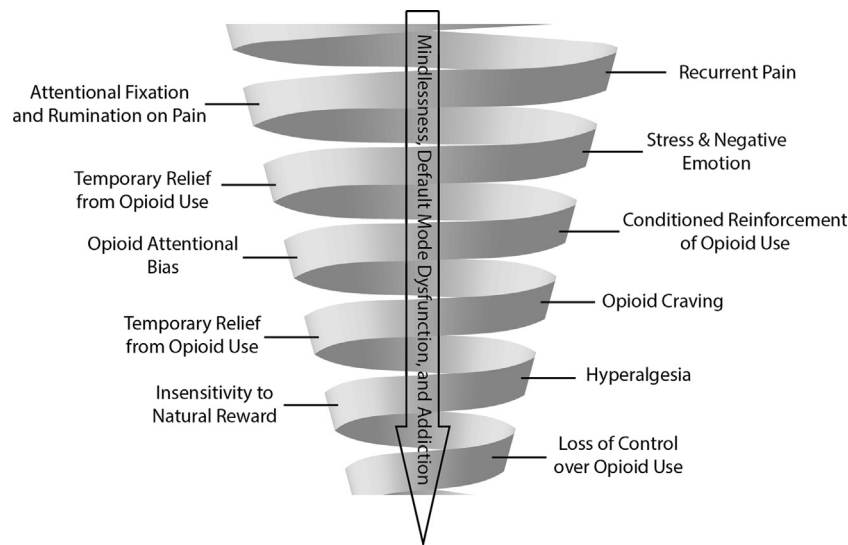


Fig. 1. The downward spiral of chronic pain and opioid addiction.

In brief, the problem of co-occurring chronic pain and opioid addiction involves a cycle of behavioral escalation in which nociception triggers pain hypervigilance and catastrophizing, amplifying pain with emotional anguish. Among affectively dysregulated individuals, recurrent self-administration of opioids in response to pain and negative emotions results in associative learning processes that bias attention towards opioid-related cues (e.g., a sight of a pill bottle), strengthening the automatic habit of opioid use despite tolerance to opioid analgesia. Chronic pain and prolonged opioid misuse causes allostatic changes to stress and reward circuitry in the brain, increasing sensitivity to pain and decreasing the pleasure derived from healthful objects and events. As functional connectivity between the default mode network and other neural circuits changes over time, the sense of self may become entwined with pain-laden narratives and entrapped by a compulsive drive for relief. This downward spiral may ultimately result in mindless, uncontrolled opioid use and addiction.

allay withdrawal irrespective of the presence of actual injury or tissue damage may be seen as a hallmark of addiction among chronic pain patients.

6. Clinical implications

In light of the complex, insidious processes outlined in this paper, multimodal interventions are needed to target the manifold links in the risk chain between chronic pain and prescription opioid addiction. Novel therapies that can facilitate attentional regulation of opioid cue-reactivity and enhance positive emotion while ameliorating pain may be efficacious means of addressing this pernicious and prevalent social problem. In that regard, mindfulness-based therapies, which are held to strengthen attentional control and temper emotional reactivity (Hölzel et al., 2011; Vago and Silbersweig, 2012), may be especially promising.

We hypothesize that mindfulness training may interrupt the downward spiral linking pain to addictive use of opioids in the following ways. Moving down the spiral (see Fig. 1) from top to bottom, mindfulness training may first reduce the severity and unpleasantness of chronic pain (Chiesa and Serretti, 2011; Gaylord et al., 2011; Rosenzweig et al., 2010; Zeidan et al., 2011) by attenuating emotionally aversive appraisals of pain sensations and decreasing pain sensitivity (Garland et al., 2011b; Zeidan et al., 2012). Second, due to its facilitative effects on attentional re-orienting capacity (Jha et al., 2007) mindfulness may reduce attentional fixation on and hypervigilance for pain (Vago and Nakamura, 2011; Garland and Howard, 2013). Third, because mindfulness involves adopting a nonjudgmental stance towards emotional experience (Hölzel et al., 2011), mindfulness may decrease negative emotional reactivity (Froeliger et al., 2012b), thereby reducing pain catastrophizing (Garland et al., 2011b) and preventing the need to self-medicate stress and negative emotions with opioids. Fourth, mindfulness may lessen addiction attentional bias (Garland et al., 2012a, 2010), and, as such, reduce biased processing of prescription opioid-related cues. Fifth, mindfulness

may decrease craving (Bowen et al., 2009) by attenuating bottom-up reactivity to drug-related stimuli (Westbrook et al., 2011) and decoupling negative affect from the urge to use substances (Witkiewitz and Bowen, 2011). Sixth, mindfulness may enhance positive affect and hedonic processing (Geschwind et al., 2011), undoing the insensitivity to natural reward that fuels increasing dependence on opioids. Seventh, mindfulness may decrease habit behavior (Wenk-Sormaz, 2005) and reduce rigid adherence to scripted cognitive responses (Greenberg et al., 2012), resulting in enhanced cognitive control of compulsive and habitual opioid use. Lastly, mindfulness may aid in regulating pain-primed mental states and self-referential processes that have been found to exacerbate and prolong the subjective experience of chronic pain (Loggia et al., 2012). In that regard, studies suggest that mindfulness practice is associated with decreased self-referential activations in cortical midline brain structures (Farb et al., 2007), reduced default mode processing (Brewer et al., 2011) and increased functional connectivity between default mode networks and attentional networks (Froeliger et al., 2012a,b). Through these therapeutic pathways, mindfulness-based interventions might help to disrupt and reverse the cycle of maladaptive pain coping habits and aberrant drug-seeking behaviors that comprise the downward spiral of pain and opioid addiction.

Though mindfulness-based approaches are relatively novel means of addressing co-occurring pain and opioid misuse, cognitive-behavioral therapy (CBT) has been used for more than three decades to treat chronic pain syndromes. As early as 1976, Fordyce advanced a behavioral model of pain management focused on operant conditioning, reinforcement of functional behaviors, and progressive relaxation (Fordyce, 1976). Modern CBT, which combines behavioral techniques with challenging of maladaptive thoughts contributing to pain catastrophizing and addictive behavior, have been shown to enhance coping with pain (Morley et al., 2008) and reduce substance misuse (Magill and Ray, 2009). A systematic review and meta-analysis conducted by Williams et al. (2012) indicated that CBT produced small to medium effect size improvements in pain, disability, and mood.

In addition, research on positive psychology suggests that savoring pleasant experiences can induce motivation and increase positive emotions (Quoidback et al., 2010), which in turn are associated with decreased pain hypervigilance and fear of pain (Crombez et al., 2012). A host of basic biobehavioral research studies support the presence of positive affect analgesia in acute pain. In healthy subjects, presentation of positive affective photographs produced greater pain tolerance on a cold pressor task than neutral and negative affective photos (De Wied and Verbaten, 2001). Positive affect analgesia has also been observed following induction of positive emotions through other means including: humorous (Weisenberg et al., 1998) and romantic films (Zillman et al., 1996), laughter (Cogan et al., 1987), and pleasant music (Silvestrini et al., 2011). Moreover, induction of positive affect through therapeutic suggestion (Zelman et al., 1991) and guided imagery (Bruehl et al., 1993) has also been shown to reduce pain. It is unknown to what extent positive affect analgesia may influence chronic pain.

Multimodal interventions that capitalize on these combined therapeutic approaches might be especially efficacious. In one of the first, well-controlled studies of a psychological treatment for chronic pain patients at high risk for misusing opioids, Jamison et al. (2010) conducted a randomized controlled trial of a multimodal intervention that combined cognitive-behavioral (e.g., coping with urges, problem solving) and motivational elements (e.g., maintaining abstinence motivations, balancing fleeting versus durable satisfactions) to decrease prescription opioid misuse and increase medication compliance. Significantly fewer high-risk individuals randomly assigned to this experimental intervention misused opioids at a six-month follow-up than those assigned to a standard medical care condition. Though most were satisfied with the study treatment, less than one-third of participants felt that it improved their pain.

To address this unmet need, Mindfulness-Oriented Recovery Enhancement (MORE; Garland, 2013), unites complementary aspects of mindfulness training, CBT, and principles from positive psychology into an integrative approach to treating co-occurring chronic pain and prescription opioid misuse. Recent findings from a randomized controlled trial demonstrate that MORE led to significant reductions in pain attentional bias coupled with decreased emotional reactivity and increased perceived control over pain in a sample of chronic pain patients receiving long-term opioid therapy, many of whom were opioid misusers (Garland and Howard, 2013). New clinical outcome data from this trial suggest that MORE may significantly reduce pain severity and functional interference while decreasing opioid misuse and craving (Garland et al., 2013). Whether mindfulness training or other forms of behavioral intervention can disrupt the downward spiral of chronic pain and opioid addiction is an open question to be explored in future clinical research.

7. Conclusion

In summary, we theorize that the problem of co-occurring chronic pain and opioid addiction involves a cycle of behavioral escalation where nociception and stress trigger hypervigilance and catastrophizing, amplifying pain and provoking recurrent self-medication with opioids, which in turn biases attention towards opioid-related cues that come to elicit the habit of drug use despite ever diminishing analgesia. Uncontrolled use of opioids coupled with chronic pain dysregulates reward processing in the brain, depriving the individual from experiencing pleasure from objects and events in the natural environment. As the sense of self becomes ensnared in pain-laden narratives and entrapped by a compulsive drive for relief, this downward spiral may result in a loss of control

over opioid use and increasing reliance on opioids to maintain an ever-diminishing sense of well-being.

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