

Management of moderate to severe chronic low back pain with buprenorphine buccal film using novel bioerodible mucoadhesive technology

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Abstract: With a global prevalence of ~9%–12%, low back pain (LBP) is a serious public health issue, associated with high costs for treatment and lost productivity. Chronic LBP (cLBP) involves central sensitization, a neuropathic pain component, and may induce maladaptive coping strategies and depression. Treating cLBP is challenging, and current treatment options are not fully satisfactory. A new BioErodible MucoAdhesive (BEMA[®]) delivery system for buprenorphine has been developed to treat cLBP. The buccal buprenorphine (BBUP) film developed for this product (Belbuca[™]) allows for rapid delivery and titration over a greater range of doses than was previously available with transdermal buprenorphine systems. In clinical studies, BBUP was shown to effectively reduce pain associated with cLBP at 12 weeks with good tolerability. The most frequently reported side effects with the use of BBUP were nausea, constipation, and vomiting. There was no significant effect on the QT interval vs placebo. Chronic pain patients using other opioids can be successfully rotated to BBUP without risk of withdrawal symptoms or inadequate analgesia. The role of BBUP in managing cLBP remains to be determined, but it appears to be a promising new product in the analgesic arsenal in general.

Keywords: buccal, transmucosal, buprenorphine, chronic low back pain, BEMA, drug delivery Belbuca

Introduction

Low back pain (LBP) is one of the top ten causes of years lived with disability (YLD) in every one of the 188 nations surveyed in the Global Burden of Disease Study of 2013 and also the leading cause of YLD in 45 of the 50 developed nations.^{1,2} The metric YLD is the number of incident cases multiplied by its average duration multiplied by a factor known as disability weight or, in some studies, it may be calculated by multiplying prevalent cases by the disability weight.³ The rates of global mortality are not declining as rapidly as the rate of YLDs with the result that health care systems will be increasingly faced with the management of the nonfatal aspects of disease and injury worldwide.⁴

LBP has been defined as pain that lasts at least 1 day in the posterior aspects of the body from the lower margin of the twelfth rib to the lower gluteal folds, which may or may not be accompanied by pain in one or both lower limbs.³ It may be also defined as a pain in the lower back lasting at least 1 day that limits activity with or without pain referred to one or both lower limbs.¹ Global prevalence of LBP is estimated to be ~9%–12%.^{1,3,5} The risk of LBP increases with age⁶ and is most prevalent among persons between the ages of 40 years and 80 years.¹ Most people (24%–80%) who experience one episode of activity-limiting LBP experience recurrent LBP at 1 year.⁶

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The costs of LBP are high, since they include medical expenditures, lost productivity, and expenses related to indemnity payments and litigation.⁷⁻⁹ In 1998, it was estimated that LBP cost the US¹⁰ > \$90 billion and in 2000, the UK estimated direct and indirect costs to be > 11 billion pounds.¹¹ Although it is impossible to state with authority exactly how much LBP costs society, it is clearly one of the costliest health care conditions.^{6,7}

The etiology of LBP can be diverse and may include anatomical or structural problems (including bones, discs, ligaments, nerves, blood vessels, and muscles), osteoporosis, neoplasm, or infection.⁶ Occupational risks for LBP include manual labor, bending, twisting, and being exposed to whole-body vibrations.¹² Other risk factors include low educational levels,¹³ obesity,¹⁴ stress, anxiety, depression,¹⁵⁻¹⁷ and job dissatisfaction,¹⁸ although the latter four factors may be more directly associated with the transition from acute LBP to chronic LBP (cLBP).^{16,19}

Most patients with nonspecific acute LBP have a favorable and unremarkable natural course in that the condition is typically brief and self-limited, with improvements occurring in ~4 weeks.²⁰ Patients with subacute LBP (lasting 4–12 weeks) usually improve as well but not as rapidly or completely. A subset of patients with acute LBP will develop cLBP, defined as LBP lasting ≥12 weeks; cLBP is difficult to treat and usually does not improve substantially over time.²⁰

Treating cLBP is challenging in part because chronic pain involves central sensitization and may include a neuropathic component. Patients with cLBP may also have comorbid conditions and psychosocial factors, such as maladaptive coping strategies and depression.²⁰

The first-line agents generally recommended for the control of cLBP are acetaminophen and nonsteroidal anti-inflammatory drugs.²¹ When such nonopioid pain relievers provide inadequate pain control, physicians may consider opioid analgesics, which carry with them risks for adverse events and abuse liability.^{22,23} Opioid analgesic agents are effective and may be used safely by certain patients under appropriate clinical supervision. Among these agents is a new buccal formulation of buprenorphine that may be particularly well suited for the treatment of cLBP. Buprenorphine is the only opioid agent that can be safely taken by elderly patients with no dosage adjustment²⁴ and addresses both nociceptive and neuropathic pain components.²⁵ Buprenorphine has been clinically evaluated in the treatment of cLBP²⁶⁻³³ and is a well-known and widely used analgesic.

The purpose of this review is to evaluate a new formulation of buprenorphine with BioErodible MucoAdhesive

(BEMA[®]; Endo Pharmaceuticals, Malvern, PA, USA) delivery technology for its potential role in the treatment of cLBP.

Methods

A literature search was conducted for keywords “buccal buprenorphine,” “Belbuca,” and “BEMA buprenorphine” in the PubMed, Cochrane, Embase, and Scopus databases, producing 152 results. Duplicates and articles not in English were removed (n=66), leaving 86 articles. Authors then reviewed the articles for relevance to the subject (buccal buprenorphine [BBUP] using BEMA technology for cLBP). Although no year constraints were employed in the search (to maximize results and not miss anything relevant), authors could also reject articles as outdated. A total of nine results remained (Figure 1).

BEMA technology and buprenorphine

Buprenorphine is a potent opioid that has been described as a partial mu-opioid receptor (MOR) agonist.²⁴ This terminology – a partial agonist – can be misleading, in that buprenorphine often acts like a full agonist in terms of clinical analgesic effect.³⁴ It offers durable analgesia in that it has high affinity for binding to MOR and slow dissociation from the MOR in the central nervous system.³⁵ Unlike other opioids, buprenorphine has been associated with antihyperalgesia³⁶ and has a ceiling effect for both gastrointestinal side effects and respiratory depression.²⁴ Buprenorphine is associated with less abuse liability than many other opioid agents.³⁷ This may be due to the fact that it acts as an agonist at the opioid

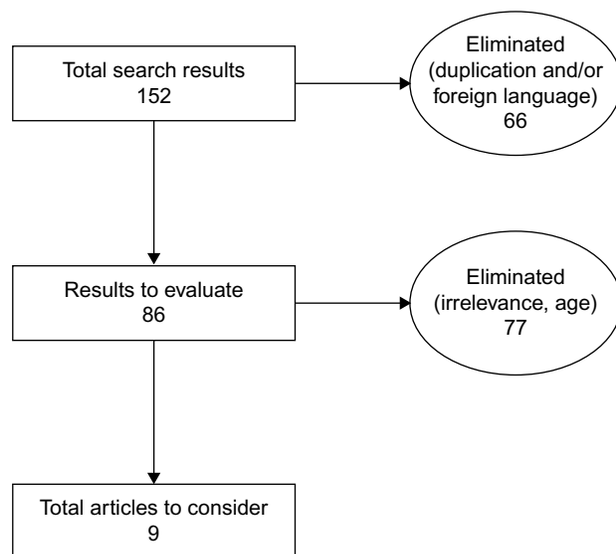


Figure 1 A literature search was conducted using PubMed, Cochrane, Embase, and Scopus databases, resulting in nine articles relevant to the objective.

receptor-like 1 (NOP) receptor, which has been associated with attenuating the rewarding effect common with other opioids.^{25,38}

The relatively poor oral bioavailability of buprenorphine (~15%³⁹) has led to the development of buprenorphine formulations as a transdermal system for pain control and a transmucosal product for opioid maintenance in dependent patients. The 7-day buprenorphine patch for chronic pain management is available in the doses of 5 µg/h, 7.5 µg/h, 10 µg/h, 15 µg/h, and 20 µg/h, with the two lower doses used primarily to initiate and titrate opioid-naïve patients.³⁹ Effective pain control has been demonstrated for buprenorphine patches at the doses of 10 µg/h, 15 µg/h, and 20 µg/h, giving buprenorphine transdermal systems a relatively narrow therapeutic dose range (10–20 µg/h).^{40,41} However, some patients may not achieve adequate analgesia with the doses of 20 µg/h.

In order to enhance and accelerate buprenorphine absorption, a buccal film delivery system was developed (Belbuca™); such a transmucosal film could increase titration flexibility and allow for a greater range of therapeutic doses. A novel and proprietary technology (BEMA) was developed to allow water-soluble polymeric films to adhere to the buccal mucosa and erode in a matter of minutes. Early testing found that this BBUP formulation had an absolute bioavailability of 46%–51% over a 16-fold range of doses.⁴² Since steady-state conditions could be reached within ~3 days of dosing for transdermal products, BBUP offers therapeutic concentrations across a broad range of doses in a shorter time period than transdermal buprenorphine formulations.⁴²

The buccal mucosa is an intensively vascularized tissue with vessels that drain into the jugular vein, although drugs penetrate the epithelium to enter directly into systemic circulation rather than undergoing first-pass hepatic elimination in the gastrointestinal tract. A potential advantage for drug delivery via oral mucosa is accessibility, although saliva (dilution also known as the “washout” effect), oral pH fluctuations, drug taste, and relatively low absorptive surface area compared to the intestines may be viewed as potential drawbacks.⁴³ Two studies were conducted to evaluate the possible effects of ingesting liquids while receiving BBUP 90 µg in 57 healthy subjects.⁴⁴ BBUP was compared to sublingual buprenorphine 8 mg in a study in which they were administered to subjects without liquids, then just prior to hot water, cold water, tepid water, and low-pH and high-pH liquids. While high-pH liquids had no significant effect on systemic exposure to buprenorphine, low-pH liquids decreased the area under the curve (AUC_{inf}) by ~37%. Water regardless of temperature decreased C_{max} and the extent of buprenorphine absorption by 23%–27%. Levels of norbuprenorphine, the metabolite of buprenorphine, were similar in all groups.

Based on AUC_t and AUC_{inf}, the relative bioavailability of BBUP compared to sublingual buprenorphine was 187% and 192%, respectively.⁴⁴

A number of buccal mucosa delivery systems for a variety of pharmaceuticals are currently in development or evaluation. The therapeutic areas for these novel products include pain, allergy, diabetes, and nicotine dependence.⁴⁵ BBUP with BEMA technology has been approved for the treatment of chronic pain severe enough to require daily, around-the-clock, long-term opioid therapy for which alternative treatment options are inadequate.⁴⁶

Pharmacokinetics

BBUP is available as buccal films in the doses of 75 µg, 150 µg, 300 µg, 450 µg, 600 µg, 750 µg, and 900 µg. It is metabolized by N-dealkylation, mainly via the cytochrome 3A4 enzyme, and glucuronidation and is eliminated in the feces (~70%) and urine (~30%). The T_{max} is 2.5–3.0 hours with steady-state concentrations reached prior to the sixth dose, and it has a half-life of 27.6 hours.⁴⁷ The half-life for sublingual and transdermal buprenorphine formulations ranges from 20 hours to 73 hours.⁴⁸ The mean plasma elimination half-life of BBUP is 27.6±11.2 hours.⁴⁹ With the 7-day transdermal buprenorphine 20 µg/h patch, there is a gradual increase in the plasma concentration of buprenorphine over the first 2 days, plateauing at ~300 pg/mL at 48 hours. The mean plasma concentration at 24 hours is 143.5 pg/mL, with C_{max} 318.6 pg/mL, and the mean AUC is 41,792.1 pg h/mL.⁵⁰ Sublingual buprenorphine, on the other hand, showed a plasma profile of 400 µg every 8 hours with large peak-to-trough differences over 24 hours. Note that buprenorphine absorption may be affected by the location of the site of application (Table 1).⁵⁰

Clinical efficacy

In a randomized, multicenter, double-blind, placebo-controlled study of 749 opioid-naïve patients with moderate to severe cLBP, patients were titrated to a dose of BBUP in the range of 150–450 µg every 12 hours that they could tolerate well and which provided adequate pain control for at least 14 days.⁵² At that point, patients were randomized to continue BBUP (n=229) or be treated with placebo (n=232) and evaluated for the change in daily average pain intensity scores (numeric 11-point scale) from baseline to week 12 of treatment. The mean daily pain intensity score at baseline was 7.15±1.05. Pain was reduced markedly at drug titration with the mean daily pain intensity score of 2.81±1.07. Once patients were randomized, placebo patients had an increase in the pain score of 1.59±2.04 compared to BBUP

Table 1 Pharmacokinetic parameters across various buprenorphine products indicated for pain control

	C_{max} (ng/mL)	AUC_{0-t} (h ng/mL)	AUC_{0-inf} (h ng/mL)	T_{max} (hours), median (range)	Absolute bioavailability (%)
Transdermal 7 d/5 µg/h	0.176	–	12.087	26	15
Transdermal 7 d/10 µg/h	0.191	27.543	27.035	26	15
Transdermal 7 d/20 µg/h	0.471	–	54.294	26	15
BBUP (Belbuca) 75 µg	0.17±0.30	0.46±0.22	0.63±0.24	3.0 (1.5–4.0)	46–65
BBUP (Belbuca) 300 µg	0.47±0.47	2.00±0.68	2.3±0.68	2.5 (0.5–4.0)	46–65
BBUP (Belbuca) 1,200 µg	1.43±0.45	9.6±2.9	10.5±3.32	3.0 (1.0–4.0)	46–65

Note: “–” data not available. Data from Endo⁴⁹ and Purdue Pharma⁵¹ package inserts.

Abbreviation: BBUP, buccal buprenorphine.

where pain levels decreased by a mean of 0.94 ± 1.85 . The difference between BBUP and placebo patients at 12 weeks was significant (-0.67 difference, 95% CI, range -1.07 to -0.26 , $P=0.0012$). Moreover, the proportion of patients who achieved a $\geq 30\%$ reduction in pain was significantly larger in the BBUP group than in the placebo group (63% vs 47%, respectively, $P=0.0012$) (Figure 2). The most common adverse events reported in the double-blind treatment phase of the study for BBUP were nausea, constipation, and vomiting (10%, 4%, and 4%, respectively), while the most common adverse events in the placebo group were nausea, upper respiratory tract infection, headache, and diarrhea (7%, 4%, 3%, and 3%, respectively).⁵² There are several limitations of this study that must be noted. First, the double-blind phase of this study lasted 12 weeks, and longer term results cannot be inferred from these findings. Second, this study included a very specific patient population (opioid-naïve adults with cLBP for ≥ 6 months), and patients had to achieve adequate analgesia with BBUP prior to entering the study. Third, this study excluded patients with certain serious comorbid conditions, such as sleep apnea and unstable cardiac disease, and therefore, results may not be generalizable to the population of cLBP patients at large. Finally, this study used a placebo control rather than an active comparator control (Figure 2).

In a randomized, placebo-controlled, double-blind withdrawal study, opioid-experienced patients with moderate to severe cLBP were randomized to BBUP or placebo.⁵³ To enter the study, patients had to be previously taking $30\text{--}160$ mg/d of morphine equivalents, and the doses were then tapered to ≤ 30 mg/d. At that point, patients underwent an open-label titration of BBUP (range $150\text{--}900$ µg every 12 hours) that would provide adequate pain control and would be well tolerated. After 14 days, patients were randomized in a double-blind fashion to receive either BBUP or placebo, and the primary endpoint was the mean of the average daily pain intensity score change from baseline to week 12. Prior to titration, the mean pain score was 6.7 ± 1.30 . Following

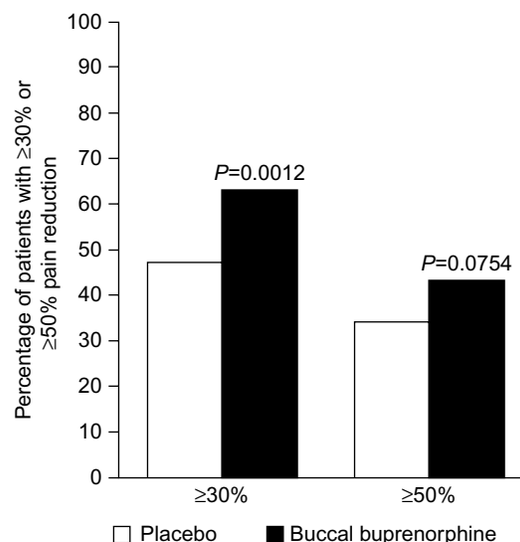


Figure 2 Significantly more patients achieved $\geq 30\%$ pain reduction in the BBUP (63%) vs placebo (47%) groups, $P=0.0012$.

Note: Significance is indicated by the asterisk. The black bars indicate the proportion of patients who achieved at $\geq 30\%$ and $\geq 50\%$ pain intensity with BBUP. The white bars indicate the placebo rates. Data from Rauck et al.⁵²

Abbreviation: BBUP, buccal buprenorphine.

titration, the average baseline pain score had dropped to 2.8 ± 1.02 . Over the course of the study, pain scores increased to 1.92 ± 1.87 for the placebo patients when compared with 0.88 ± 1.79 for the BBUP patients ($P<0.00001$). Significantly more patients who achieved pain reductions by $\geq 30\%$ and $\geq 50\%$ were in the BBUP group compared to the placebo group (64% vs 31%, $P<0.0001$ and 40% vs 17%, $P<0.0001$, respectively).⁵³ The same limitations apply to this study: a specific and tightly defined patient population, placebo rather than active comparator as control, exclusion of patients with serious comorbidities, and adequate analgesia and tolerability of BBUP prior to entering the study. Again, this study had a double-blind phase of 12 weeks, and results cannot be generalized to longer time periods.

Safety

There have been concerns that buprenorphine, even at therapeutic doses, prolongs the QT interval on electrocardiography,³⁹

which has raised concern about prescribing buprenorphine to patients taking antiarrhythmic agents. Moreover, prolonged QT intervals could be problematic in any number of cardiac conditions, including, but not limited to, unstable atrial fibrillation and symptomatic bradycardia. A thorough QT study evaluated the effect of buprenorphine as a buccal soluble film on cardiac repolarization in 58 healthy subjects.⁵⁴ In this study, subjects received BBUP with naltrexone to mitigate opioid-associated effects that might potentially confound results. A separate group receiving only naltrexone was evaluated in the study. Subjects were randomized to one of the four groups receiving 3 mg of BBUP with naltrexone (50 mg of naltrexone was administered 12 hours prior to the first dose of buprenorphine); 50 mg naltrexone only on a placebo film; placebo (placebo on placebo film); and open-label moxifloxacin 400 mg. Except for the moxifloxacin group, all treatments were administered in a double-blind fashion. The study was designed as a four-period, crossover, single-dose evaluation. During each period, blood was drawn for pharmacokinetic evaluation, and 12-lead electrocardiography assessments were conducted. This study found that even supratherapeutic doses of BBUP administered with naltrexone did not cause any clinically significant QT interval prolongation. While there was a slight shortening of the changes from the baseline PR interval in all treatment periods, it appeared to be smaller in the BBUP and naltrexone groups than in the naltrexone alone and placebo groups. Naltrexone by itself did not appear to exert any relevant effect on the QT interval. Changes from the baseline QRS-interval were very small (≤ 1 ms at all time points).⁵⁴

In this same study, 67.2% (n=39) of patients reported at least one treatment-emergent adverse event.⁵⁴ This rate was highest among the buprenorphine plus naltrexone patients (53.7%) compared to patients who took naltrexone alone (30.8%), placebo patients (5.9%), or moxifloxacin patients (15.7%). In the BBUP and naltrexone groups, the most frequently reported adverse events were nausea (27.8%), dizziness (14.8%), vomiting (13.0%), and headache (7.5%); these events were relatively uncommon in the naltrexone alone, placebo, and moxifloxacin groups (<10% for each). Most of the adverse events were deemed to be mild, no serious adverse events were observed, and no subject withdrew from the study because of adverse events.⁵⁴

In a 12-week study of BBUP for cLBP (n=511), the most commonly reported adverse events during the double-blind phase of the study were nausea and vomiting (8% and 6%, respectively) for the BBUP patients and drug withdrawal syndrome and nausea (10% and 7%, respectively) for the placebo patients.⁵³

Opioid rotation to BBUP

Opioid rotation can be an important pharmacological strategy to allow patients with suboptimal results on one opioid to potentially achieve better results (improved analgesia, greater tolerability, or both) with another opioid agent.⁵⁵ Patients receiving a dose equivalent to 80–220 mg of oral morphine sulfate (n=35) were transitioned to BBUP at 50% of their full dose in a randomized, double-blind, double-dummy, active-control two-period crossover study.⁵⁶ The primary endpoint of this study was a composite: either a maximum Clinical Opiate Withdrawal Scale score of ≥ 13 (moderate withdrawal) or the use of rescue medication to manage pain. All patients in the study had chronic pain and were confirmed as opioid dependent by naloxone challenge. It was found that these patients could be switched to BBUP without increased risk of withdrawal or exposure to inadequate analgesia.

Discussion

Chronic pain is a debilitating condition that decreases productivity, limits function, and reduces the quality of life. More than 100 million Americans suffer chronic painful conditions, with cLBP a leading cause, and many of these patients may be considered for opioid therapy.⁵⁷ Unrelieved chronic pain is epidemic in the US, and many cLBP and other chronic pain patients are not highly passionate advocates for better pain control. This is in part the nature of the disorder: with the passage of time, chronic pain causes individuals to withdraw from society, to retreat from activities, and to cope with life rather than engage. Opioids are effective pain relievers, and many patients benefit from their appropriate use. However, opioids carry with them a well-recognized abuse liability, and for this reason, many clinicians hesitate to prescribe them except for patients with the most severe forms of chronic noncancer pain. Indeed, there are respected experts who challenge the use of opioids in the setting of chronic nonmalignant pain altogether.^{58,59} Recent guidelines from the Centers for Disease Control and Prevention recommend limiting the use of opioids in patient populations such as patients with cLBP.⁶⁰

Pain experts are left to weigh the evidence in evaluating various pain control strategies and to make prescribing choices based on the needs and lifestyles of individual patients. The perfect analgesic agent does not yet exist. We have a pharmacological armamentarium of effective pain relief products, all of which carry some degree of risk. It is in this light that buprenorphine should be reconsidered. According to the World Health Organization, buprenorphine

is considered as a strong opioid,⁶¹ and it possesses a unique pharmacology and limited abuse liability.²⁴ It may be used without dose adjustment in elderly patients (a large subset of the chronic pain population) and those with renal dysfunction.²⁵ Buprenorphine administered transdermally is a convenient way of providing round-the-clock analgesia with good patient adherence.

The role of buprenorphine for pain control has been limited by concerns that it may be difficult to titrate adequately. Despite the apparent advantages of buprenorphine,²⁵ it is often considered as an “older drug” with limited utility. New indications for chronic pain, our knowledge of the drug and its unique pharmacology,²⁴ and the BBUP product make it worthy of a second look.

Transmucosal products are easy to administer and convenient for the clinical team. Patient acceptance of transmucosal products has not been well studied, although another transmucosal product (fentanyl) is on the market. The advantages of a convenient, rapid-onset opioid pain reliever with limited abuse liability makes the advent of this new BBUP particularly promising.

As with any opioid pain reliever, patients should be informed about the potential risks of opioid therapy, notably the risk of certain opioid-associated side effects and abuse liability. Opioid therapy should be closely supervised with patients and physicians setting goals for treatment that might include pain control, increased function, and specific personal goals such as being able to sit through a movie comfortably or being able to go out to dinner with the family. While pain assessments using validated metrics are important clinical tools, personal goals can have more relevance and be more motivational to the patient and his or her family than decreasing pain intensity numbers.

There are only a few studies in the literature addressing BBUP to date, and these studies have certain limitations. The authors could not find a BBUP study in the literature that compared BBUP with an active comparator. Furthermore, the studies described in our article evaluated BBUP over a 12-week time period (double-blind phase) in very refined patient populations who had achieved adequate analgesia and good tolerability with BBUP before entering the study. Thus, prudence is warranted when considering this new drug. Nevertheless, studies so far indicate that BBUP offers clinical efficacy, safety, tolerability, and versatility to help address cLBP. As such it must be considered as a promising and important new product for clinicians and their patients suffering from moderate to severe cLBP.

Conclusion

Buprenorphine is a potent and important opioid analgesic with a unique pharmacology. Until recently, buprenorphine's

availability to chronic pain patients was limited to transdermal patch systems in only three doses. BBUP offers a novel transmucosal technology that may enhance and accelerate buprenorphine absorption in the body. BBUP allows for greater dosing versatility (an important consideration for some cLBP patients). Buprenorphine is associated with a lower abuse liability than many other strong opioids, a ceiling effect for respiratory depression, and no requirement to adjust doses for the elderly or renally impaired patients. QT interval prolongation was not evident with BBUP even at supratherapeutic doses. Thus, BBUP may be an important new addition to help treat patients with moderate to severe pain associated with cLBP.

Disclosure

Dr. Pergolizzi is a speaker for ENDO, DEPOMED, Grünenthal, Purdue Pharma, and Mundipharma; he is a researcher for Purdue Pharma, Kaleo, and Collegium; and a consultant for Adapt, ENDO, Grünenthal, Purdue Pharma, and Insys Therapeutics. Dr. Pergolizzi received no funding related to the production of this manuscript. Dr. Raffa is a speaker, consultant, and/or basic science investigator for several pharmaceutical companies involved in analgesics research but receives no royalty (cash or otherwise) from the sale of any product. Drs. Fleischer, Taylor, and Zampogna are research fellows for NEMA Research Inc., a clinical research organization which provides clinical, regulatory, and product development services to a number of pharmaceutical companies.

References

1. Hoy D, Bain C, Williams G, et al. A systematic review of the global prevalence of low back pain. *Arthritis Rheum.* 2012;64(6):2028–2037.
2. GBD 2013 DALYs and HALE Collaborators, Murray CJ, Barber RM, et al. Global, regional, and national disability-adjusted life years (DALYs) for 306 diseases and injuries and healthy life expectancy (HALE) for 188 countries, 1990–2013: quantifying the epidemiological transition. *Lancet.* 2015;386(10009):2145–2191.
3. March L, Smith EU, Hoy DG, et al. Burden of disability due to musculoskeletal (MSK) disorders. *Best Pract Res Clin Rheumatol.* 2014;28(3):353–366.
4. Global Burden of Disease Study 2013 Collaborators. Global, regional, and national incidence, prevalence, and years lived with disability for 301 acute and chronic diseases and injuries in 188 countries, 1990–2013: a systematic analysis for the Global Burden of Disease study 2013. *Lancet.* 2015;386(9995):743–800.
5. Hoy DG, Smith E, Cross M, et al. The global burden of musculoskeletal conditions for 2010: an overview of methods. *Ann Rheum Dis.* 2014;73(6):982–989.
6. Hoy D, Brooks P, Blyth F, Buchbinder R. The epidemiology of low back pain. *Best Pract Res Clin Rheumatol.* 2010;24(6):769–781.
7. Walker B, Muller R, Grant W. Low back pain in Australian adults: the economic burden. *Asia Pac J Public Health.* 2003;15(2):79–87.
8. Buchbinder R, Blyth FM, March LM, Brooks P, Woolf AD, Hoy DG. Placing the global burden of low back pain in context. *Best Pract Res Clin Rheumatol.* 2013;27(5):575–589.
9. Kent PM, Keating JL. The epidemiology of low back pain in primary care. *Chiropr Osteopat.* 2005;13:13.

10. Luo X, Pietrobon R, Sun SX, Liu GG, Hey L. Estimates and patterns of direct health care expenditures among individuals with back pain in the United States. *Spine (Phila Pa 1976)*. 2004;29(1):79–86.
11. Maniadakis N, Gray A. The economic burden of back pain in the UK. *Pain Manag Nurs*. 2000;84(1):95–103.
12. Hoogendoorn WE, van Poppel MN, Bongers PM, Koes BW, Bouter LM. Systematic review of psychosocial factors at work and private life as risk factors for back pain. *Spine (Phila Pa 1976)*. 2000;25(16):2114–2125.
13. Dionne CE, Von Korff M, Koepsell TD, Deyo RA, Barlow WE, Checkoway H. Formal education and back pain: a review. *J Epidemiol Community Health*. 2001;55(7):455–468.
14. Webb R, Brammah T, Lunt M, Urwin M, Allison T, Symmons D. Prevalence and predictors of intense, chronic, and disabling neck and back pain in the UK general population. *Spine (Phila Pa 1976)*. 2003;28(11):1195–1202.
15. Rubin DI. Epidemiology and risk factors for spine pain. *Neurol Clin*. 2007;25(2):353–371.
16. Linton S. A review of psychological risk factors in back and neck pain. *Spine*. 2000;25(9):1148–1156.
17. Carroll LJ, Cassidy JD, Côté P. Depression as a risk factor for onset of an episode of troublesome neck and low back pain. *Pain*. 2004;107(1–2):134–139.
18. van Tulder M, Koes B, Bombardier C. Low back pain. *Best Pract Res Clin Rheumatol*. 2002;16(5):761–775.
19. Pincus T, Burton AK, Vogel S, Field AP. A systematic review of psychological factors as predictors of chronicity/disability in prospective cohorts of low back pain. *Spine (Phila Pa 1976)*. 2002;27(5):E109–E120.
20. Chou R, McCarberg B. Managing acute back pain patients to avoid the transition to chronic pain. *Pain Manag*. 2011;1(1):69–79.
21. Chou R, Qaseem A, Snow V, et al. Diagnosis and treatment of low back pain: a joint clinical practice guideline from the American College of Physicians and the American Pain Society. *Ann Intern Med*. 2007;147(7):478–491.
22. Chou R, Fanciullo GJ, Fine PG, et al. Clinical guidelines for the use of chronic opioid therapy in chronic noncancer pain. *J Pain*. 2009;10(2):113–130.
23. Chou R. 2009 clinical guidelines from the American Pain Society and the American Academy of Pain Medicine on the use of chronic opioid therapy in chronic noncancer pain: what are the key messages for clinical practice? *Pol Arch Med Wewn*. 2009;119(7–8):469–477.
24. Pergolizzi J, Aloisi AM, Dahan A, et al. Current knowledge of buprenorphine and its unique pharmacological profile. *Pain Pract*. 2010;10(5):428–450.
25. Davis MP. Twelve reasons for considering buprenorphine as a frontline analgesic in the management of pain. *J Support Oncol*. 2012;10(6):209–219.
26. Chaparro LE, Furlan AD, Deshpande A, Mailis-Gagnon A, Atlas S, Turk DC. Opioids compared to placebo or other treatments for chronic low-back pain. *Cochrane Database Syst Rev*. 2013;(8):CD004959.
27. Gatti A, Dauri M, Leonardi F, et al. Transdermal buprenorphine in non-oncological moderate-to-severe chronic pain. *Clin Drug Investig*. 2010;30(suppl 2):31–38.
28. Gordon A, Rashiq S, Moulin DE, et al. Buprenorphine transdermal system for opioid therapy in patients with chronic low back pain. *Pain Res Manag*. 2010;15(3):169–178.
29. Gordon A, Callaghan D, Spink D, et al. Buprenorphine transdermal system in adults with chronic low back pain: a randomized, double-blind, placebo-controlled crossover study, followed by an open-label extension phase. *Clin Ther*. 2010;32(5):844–860.
30. Miller K, Yaras A, Wen W, et al. Buprenorphine transdermal system and quality of life in opioid-experienced patients with chronic low back pain. *Expert Opin Pharmacother*. 2013;14(3):269–277.
31. Miller K, Yaras A, Wen W, et al. The impact of buprenorphine transdermal delivery system on activities of daily living among patients with chronic low back pain: an application of the international classification of functioning, disability and health. *Clin J Pain*. 2014;30(12):1015–1022.
32. Petzke F, Welsch P, Schaefer R, Sommer C, Hauser W. Opioid bei chronischem Kreuzschmerz. Systematische Übersicht und Metaanalyse der Wirksamkeit, Verträglichkeit und Sicherheit in randomisierten, placebokontrollierten Studien über mindestens 4 Wochen [Opioids in chronic low back pain. A systematic review and meta-analysis of efficacy, tolerability and safety in randomized placebo-controlled studies of at least 4 weeks duration]. *Schmerz*. 2014;29(1):60–72. German.
33. Pota V, Barbarisi M, Sansone P, et al. Combination therapy with transdermal buprenorphine and pregabalin for chronic low back pain. *Pain Manag*. 2012;2(1):23–31.
34. Raffa RB, Ding Z. Examination of the preclinical antinociceptive efficacy of buprenorphine and its designation as full- or partial-agonist. *Acute Pain*. 2007;9(3):145–152.
35. Wolff RF, Reid K, di Nisio M, et al. Systematic review of adverse events of buprenorphine patch versus fentanyl patch in patients with chronic moderate-to-severe pain. *Pain Manag*. 2012;2(4):351–362.
36. Koppert W, Ihmsen H, Körber N, et al. Different profiles of buprenorphine-induced analgesia and antihyperalgesia in a human pain model. *Pain*. 2005;118(1–2):15–22.
37. Lofwall MR, Walsh SL. A review of buprenorphine diversion and misuse: the current evidence base and experiences from around the world. *J Addict Med*. 2014;8(5):315–326.
38. Huang P, Kehner GB, Cowan A, Liu-Chen LY. Comparison of pharmacological activities of buprenorphine and norbuprenorphine: norbuprenorphine is a potent opioid agonist. *J Pharmacol Exp Ther*. 2001;297(2):688–695.
39. Drugs.com [webpage on the Internet]. Butrans Skin Patch. 2016. Available from: <http://www.drugs.com/pro/butrans-patch.html>. Accessed March 19, 2016.
40. Steiner D, Munera C, Hale M, Ripa S, Landau C. Efficacy and safety of buprenorphine transdermal system (BTDS) for chronic moderate to severe low back pain: a randomized, double-blind study. *J Pain*. 2011;12(11):1163–1173.
41. Steiner DJ, Sitar S, Wen W, et al. Efficacy and safety of the seven-day buprenorphine transdermal system in opioid-naïve patients with moderate to severe chronic low back pain: an enriched, randomized, double-blind, placebo-controlled study. *J Pain Symptom Manage*. 2011;42(6):903–917.
42. Bai SA, Xiang Q, Finn A. Evaluation of the pharmacokinetics of single- and multiple-dose buprenorphine buccal film in healthy volunteers. *Clin Ther*. 2016;38(2):358–369.
43. Pather SI, Rathbone MJ, Senel S. Current status and the future of buccal drug delivery systems. *Expert Opin Drug Deliv*. 2008;5(5):531–542.
44. Priestley T, Xiang S, Vasisht N, Cheruvu N. Factors impacting the relative bioavailability of a new formulation of buprenorphine. *J Pain Palliat Care Pharmacother*. 2016;17(4 suppl 1):S888.
45. Silva BM, Borges AF, Silva C, Coelho JF, Simões S. Mucoadhesive oral films: the potential for unmet needs. *Int J Pharm*. 2015;494(1):537–551.
46. Han D [webpage on the Internet]. Belbuca Now Available for Managing Severe, Chronic Pain. 2016. Available from: <http://www.empr.com/news/belbuca-now-available-for-managing-severe-chronic-pain/article/478192/>. Accessed March 19, 2016.
47. Buprenorphine buccal film (Belbuca) for chronic pain. *Med Lett Drugs Ther*. 2016;58(1492):47–48.
48. Khanna IK, Pillarsetti S. Buprenorphine – an attractive opioid with underutilized potential in treatment of chronic pain. *J Pain Res*. 2015;8:859–870.
49. Endo [webpage on the Internet]. BELBUCA-Buprenorphine Hydrochloride Film. Highlights of Prescribing Information. 2015. Available from: http://www.endo.com/File%20Library/Products/Prescribing%20Information/BELBUCA_prescribing_information.html. Accessed June 9, 2016.
50. Plosker GL. Buprenorphine 5, 10 and 20 mug/h transdermal patch: a review of its use in the management of chronic non-malignant pain. *Drugs*. 2011;71(18):2491–2509.
51. Purdue Pharma [webpage on the Internet]. BUTRANS-Buprenorphine Patch, Extended Release. 2014. Available from: <http://app.purduepharma.com/xmlpublishing/pi.aspx?id=b>. Accessed June 9, 2016.
52. Rauck RL, Potts J, Xiang Q, Tzanis E, Finn A. Efficacy and tolerability of buccal buprenorphine in opioid-naïve patients with moderate to severe chronic low back pain. *Postgrad Med*. 2016;128(1):1–11.

53. Gimbel J, Spierings EL, Katz N, Xiang Q, Tzannis E, Finn A. Efficacy and tolerability of BEMA buprenorphine in opioid-experienced patients with moderate-to-severe chronic low back pain: primary results from a phase 3, enriched-enrollment, randomized withdrawal study. *J Pain*. 2015; 16(4):S85.
54. Darpo B, Zhou M, Bai SA, Ferber G, Xiang Q, Finn A. Differentiating the effect of an opioid agonist on cardiac repolarization from micro-receptor-mediated, indirect effects on the QT interval: a randomized, 3-way crossover study in healthy subjects. *Clin Ther*. 2016;38(2):315–326.
55. Nalamachu SR. Opioid rotation in clinical practice. *Adv Ther*. 2012;29(10): 849–863.
56. Webster L, Gruener D, Kirby T, Xiang Q, Tzannis E, Finn A. Evaluation of the tolerability of switching patients on chronic full mu-opioid agonist therapy to buccal buprenorphine. *Pain Med*. Epub 25 February, 2016.
57. Trang T, Al-Hasani R, Salvemini D, Salter MW, Gutstein H, Cahill CM. Pain and poppies: the good, the bad, and the ugly of opioid analgesics. *J Neurosci*. 2015;35(41):13879–13888.
58. Manchikanti L, Atluri S, Hansen H, et al. Opioids in chronic noncancer pain: have we reached a boiling point yet? *Pain Physician*. 2014;17(1): E1–E10.
59. Manchikanti L, Benyamin R, Datta S, Vallejo R, Smith H. Opioids in chronic noncancer pain. *Expert Rev Neurother*. 2010;10(5): 775–789.
60. Dowell D, Haegerich TM, Chou R. CDC guideline for prescribing opioids for chronic pain – United States, 2016. *MMWR Recomm Rep*. 2016; 65(1):1–49.
61. World Health Organization [webpage on the Internet]. WHO's Pain Ladder for Adults. 1988. Available from: <http://www.who.int/cancer/palliative/painladder/en/>. Accessed May 7, 2013.

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