## COMMENTARY

## **Opioids and Methadone Equivalents for Clinicians**

W. Victor R. Vieweg, M.D.; William F. Carlyle Lipps, Pharm.D.; and Antony Fernandez, M.D.

Settings in which clinicians are likely to prescribe narcotics and opioids include those that manage chronic pain, methadone maintenance programs, and primary care. More commonly, clinicians prescribe psychotropic medications that may potentiate opioid-associated respiratory depression.

The terms *narcotics* and *opioids* are commonly used interchangeably. These substances bind to the  $\mu$  receptors for their euphorigenic, mood-altering, and dependenceproducing properties.<sup>1</sup> These agents also may depress respiration via the  $\mu$  receptors in the pons and medulla oblongata. Positron emission tomography (PET) scan of  $\mu$  receptors shows them to be located in the (1) thalamus (highest concentrations and involved in pain), (2) cerebral cortex (intermediate concentrations), (3) basal ganglia (intermediate concentrations and involved in movement and emotions), and (4) visual cortex (lowest concentrations).<sup>2</sup> PET scanning has also located  $\mu$  receptors in the pons and medulla, but semiquantitative estimates are not available.

The respiratory center is located in the pons and medulla oblongata. This center is a part of the reptilian brain when separating the central nervous system into (1) brain stem (midbrain, pons, and medulla oblongata) and cerebellum (reptilian), (2) limbic system (mammalian), and (3) cortex and neocortex (human) divisions.<sup>3</sup> Because the respiratory center arose in the brain stem more than 200 million years ago with primitive vertebrates, it is hardier than more recent brain structures. Thus, lower doses of narcotics and opioids may provide pain relief as agonists for  $\mu$  receptors. At higher doses, narcotics and opioids may cause respiratory depression.

Clinicians work with patients suffering from both acute and chronic pain. Longterm pain management is more problematic, particularly in the mental health field. Long-acting opioids include methadone, sustained-release preparations of morphine, and fentanyl patches. Of these 3 opioids, clinicians in the mental health field tend to be most familiar with methadone. It is important that clinicians be conversant with methadoneequivalent doses of various opioids that their patients may be taking because of the potential respiratory-depressing effects of nonopioid psychotropic drugs coupled with opioid psychotropic drugs.

There is no clear linear relationship between methadone dose and respiratory depression because of several factors. Subjects receiving long-term opioid treatment usually develop tolerance to the respiratory-depressant effects of these drugs. Respiratory depression may occur when pain is abruptly relieved and the sedative effects of opioids are no longer opposed by the stimulating effects of pain. This is a rare event.

In methadone maintenance programs, effective methadone doses today generally fall in the 60- to 100-mg/day range.<sup>4,5</sup> Methadone doses greater than 40 to 50 mg/day in subjects entering a methadone maintenance program may be associated with respiratory depression.

Received Nov. 22, 2004; accepted March 16, 2005. From the Departments of Psychiatry and Internal Medicine, Medical College of Virginia, Virginia Commonwealth University and Hunter Holmes McGuire Veterans Affairs Medical Center, Richmond, Va.

There was no external funding for this project.

Drs. Vieweg, Lipps, and Fernandez report no financial or other affiliations relevant to this article. Corresponding author and reprints: W. Victor R. Vieweg, M.D., 17 Runswick Drive, Richmond, VA

<sup>23238-5414 (</sup>e-mail: vvieweg@vcu.edu).

Table 1. Commonly U Cytochrome P450 3A and Tissue Levels of M	sed Medications That May Inhibit 4 Metabolism and Increase Blood Aethadone <sup>a</sup>
Fluconazole	Erythromycin
Itraconazole	Clarithromycin
Ketoconazole	Diltiazem
Ritonavir	Verapamil
Nelfinavir	Paroxetine
Amiodarone	Fluoxetine
<sup>a</sup> Based on Lacy et al. <sup>6</sup>	

In terms of drug interactions, methadone is a minor substrate of cytochrome P450 (CYP) 2C8/9, 2C19, and 2D6; it is a major substrate of CYP3A4.<sup>6</sup> Methadone moderately inhibits CYP2D6 and weakly inhibits CYP-3A4.<sup>6</sup> Concomitant administration of drugs inhibiting the CYP3A4 isoenzyme may contribute to methadone overdose by increasing methadone accumulation. Table 1 lists commonly used medications that may inhibit CYP3A4 metabolism.

## CONVERSION TABLE

We provide an opioid conversion table (Appendix 1)<sup>7-10</sup> for commonly used opioid preparations to help clinicians better understand the relationship between these agents and methadone. Conversion must take into consideration clinical issues that affect translation of equivalents to and from methadone.

Concomitant drugs affecting the metabolism of the nonmethadone drug may not similarly affect the metabolism of methadone and vice versa during and after conversion.<sup>11</sup> Similarly, organ disease, particularly liver disease, may confound the conversion process. Also, the nonmethadone drug may have a different effect than methadone on such psychiatric conditions as anxiety and depression. In a perfect world, the clinician would have serial plasma levels of the nonmethadone drug before conversion and then serial plasma levels of methadone during and after conversion. However, even here there may not be a linear relationship between plasma level and clinical effect. Methadone plasma levels are most useful when the patient's current state does not agree with the expected state of methadone treatment. Ultimately, clinicians must depend on the patient's report and their own assessment and experience when making medical judgments. This principle most certainly applies when converting from a nonmethadone narcotic to methadone.

In *The American Psychiatric Publishing Textbook of Substance Abuse Treatment*,<sup>12</sup> Gunderson and Stimmel dedicate a section of their chapter on treating pain in drugaddicted patients to the conversion from one opioid to another. They emphasize the importance of using equivalence doses. Because of incomplete cross-tolerance, Gunderson and Stimmel<sup>12</sup> recommend dividing the calculated equivalent dose in half followed by adjusting that dose upward for the first 24 hours of conversion to optimize analgesia. For patients dependent on methadone in a methadone treatment program, prescription of an opioid analgesic in excess of baseline requirements is recommended-initially prescribe two thirds of the calculated total dose and adjust upward as needed. When changing from other opioids to methadone for pain control, clinicians should start methadone at a dose much lower than the equi-analgesic dose-down to one tenth the calculated dose-and then monitor the patient closely over the ensuing 3 to 6 days.

*Drug names:* amiodarone (Cordarone, Pacerone, and others), clarithromycin (Biaxin and others), diltiazem (Cardizem, Tiazac, and others), erythromycin (Eryc, PCE, and others), fluconazole (Diflucan and others), fluoxetine (Prozac and others), itraconazole (Sporanox and others), ketoconazole (Ketozole, Nizoral, and others), nelfinavir (Viracept), paroxetine (Paxil and others), ritonavir (Norvir), verapamil (Calan, Isoptin, and others).

## REFERENCES

- Jaffe JH, Jaffe AB. Neurobiology of opioids. In: Galanter M, Kleber HD, eds. Textbook of Substance Abuse Treatment. 3rd ed. Washington, DC: American Psychiatric Publishing, Inc; 2004:17–30
- Ravert HT, Bencherif B, Madar I, et al. PET imaging of opioid receptors in pain: progress and new directions. Curr Pharm Des 2004;10:759–768
- Carter R. Mapping the Mind. Berkeley, Calif: University of California Press; 1998
- Strain EC, Bigelow GE, Liebson IA, et al. Moderate- vs high-dose methadone in the treatment of opioid dependence: a randomized trial. JAMA 1999;281:1000–1005
- Schottenfeld RS. Opioids. Maintenance Treatment. In: Galanter M, Kleber HD, eds. Textbook of Substance Abuse Treatment. 3rd ed. Washington, DC: American Psychiatric Publishing, Inc; 2004:291–304
- Lacy CF, Armstrong LL, Goldman MP, et al. Drug Information Handbook 2004–2005. Hudson, Ohio: Lexi-Comp; 2004
- DuBe JE, Koo PJ. Pain. In: Young LY, Koda-Kimble MA, eds. The Clinical Use of Drugs. Vancouver, Wash: Applied Therapeutics; 1988: 9-1–9-40
- MS Contin (morphine sulfate-controlled-release) [package insert]. Stamford, Conn: Purdue Frederick; 1995
- Lichtor JL, Sevarino FB, Joshi GP, et al. The relative potency of oral transmucosal fentanyl citrate compared with intravenous morphine in the treatment of moderate to severe postoperative pain. Anesth Analg 1999;89:732–738
- ASHP Drug Information 2004. Bethesda, Md: American Society of Health-System Pharmacists, Inc; 2004
- Payte JT, Zweben JE. Opioid maintenance therapies. In: Graham AW, Schultz TK, Wilford BB, eds. Principles of Addiction Medicine. 2nd ed. Chevy Case, Md: American Society of Addiction Medicine, Inc; 1998: 557–570
- Gunderson EW, Stimmel B. Treatment of pain in drug-addicted persons. In: Galanter M, Kleber HD, eds. Textbook of Substance Abuse Treatment. 3rd ed. Washington, DC: American Psychiatric Publishing, Inc; 2004:563–573

Appendix 1 appears on page 88.

Appendix 1. O <sub>I</sub>	vioid Conversion Dat	a: Equi-Analge:	sic Dosing G	uide Equivalenc	cy Table				
Chemical Name	Trade Name (examples)	FDA Schedule	Onset	Duration	Half-Life of Parent Drug	Typical Dose Ranges	Example Daily Dosing Regimen	Equivalent Morphine Dose (po) (mg/d) to Example	Equivalent Methadone Dose (po) (mg/d) to Example
Fentanyl patch	Duragesic	=	12–24 h	60–72 h	7 h	25–300 µg (transdermal) q 72 h	50 µg q 72 h	150-200	60-80
Fentanyl transmucosal	Actiq	=	5–15 min	4–8 h (dependent on blood levels)	17 h	200–1600 Jug q 8–12 h (only for chronic opiate users)	400 µg q 8 h	45-60	18–24
Hydromorphone oral	Dilaudid	=	15–30 min	4–6 h	2–4 h	2–8 mg po q 6–8 h	4 mg q 6 h	64	26
Meperidine oral	Demerol	=	10–15 min	2–4 h	3-4 h; liver disease patients = $7-11$ h; active metabolite = 15-30 h	50–100 mg po q 4–6 h	100 mg q 6 h	40	16
Methadone oral	Dolophine, Methadose	=	30–60 min	4–8 h > 8 h (chronic)	15–29 h	10–100 mg q 6–8 h	20 mg q 8 h	150	60
Morphine oral	MSIR, Roxanol	=	15-60 min	3–6 h	2–4 h	5-20 mg q 4-6 h (as adjunct)	10 mg q 4 h	60	24
Morphine oral (long-acting)	Avinza, Kadian, MS Contin, Oramorph	=	2–3 h (Avinza = 30 min)	8-14 h (Avinza = 18-24 h)	2–4 h	15–100 mg q 8–12 h	60 mg q 12 h	120	48
Oxycodone oral	OxyFast, Percocet, Percodan, Roxicet, Roxicodone, Tylox	=	15-30 min	4–6 h	3-4 h	5-15 mg q 4-6 h (as adjunct)	10 mg q 6 h	80	32
Oxycodone oral (long-acting)	OxyContin	=	1–2 h	6–10 h	3–4 h	10 mg–80 mg q 12 h	40 mg q 12 h	160	64
Hydrocodone oral	Anexsia, Lorcet, Lortab, Norco, Vicodin, Vicoprofen, Zydone	Ξ	30–60 min	4–8 h	3–4.5 h	5–15 mg q 3–8 h	10 mg q 6 h	No data (in practice, 40, approximately)	No data (in practice,16, approximately)
Codeine oral	Tylenol #3 (combination)	II (alone), III, V in combinations	30–60 min	4–8 h	3–4 h	15-60 mg q 4-6 h	30 mg q 6 h	20	œ
Propoxyphene oral	Darvocet, Darvon	N	30–60 min	4–6 h	3–15 h	50–100 mg q 4–6 h	100 mg q 6 h	60	24
Abbreviation: FI [Vieweg WVR, I	DA = U.S. Food and Dr. Jpps WFC, Fernandez	ig Administratior A. Opioids and m	ı. ethadone equiv	valents for clinicia	ans. Prim Care Compan	ion J Clin Psychiatry 2005;	7:86–88]		

BibCCAYRICAMP26065 PEIro Payetta Po 200573) JATE PRESS, INC. O COPYRIGHT 2005 PHYSICIANS POSTGRADUATE PRESS, INC.

88