Characterization and validation of a pharmacokinetic model for controlled-release oxycodone

JAAP W. MANDEMA¹, ROBERT F. KAIKO², BENJAMIN OSHLACK³,

ROBERT F. REDER² & DONALD R. STANSKI¹

1Stanford University School of Medicine, Department of Anesthesia, Stanford, California, 2The Purdue Frederick Company, Medical Department, Norwalk, Connecticut and 3The Purdue Frederick Research Center, Yonkers, New York, USA

- 1 Oxycodone is a strong opioid agonist that is currently available in immediaterelease (IR) formulations for the treatment of moderate to severe pain. Recently, controlled-release (CR) oxycodone tablets were developed to provide the benefits of twice-a-day dosing to patients treated with oxycodone. The purpose of this investigation was to develop and validate a pharmacokinetic model for CR oxycodone tablets in comparison with IR oxycodone solution.
- 2 Twenty-four normal male volunteers were enrolled in a single-dose, randomized, analytically blinded, two-way crossover study designed to compare the pharmacokinetics of two 10 mg CR oxycodone tablets with 20 mg IR oxycodone oral solution. Pharmacokinetic models describing the oxycodone plasma concentration vs time profiles of CR tablets and IR solution were derived using NONMEM version IV. The predictive performance of the models was assessed by comparison of predicted oxycodone plasma concentrations with actual oxycodone plasma concentrations observed in a separate group of 21 volunteers who received repeated doses of IR and CR oxycodone for 4 days.
- 3 The unit impulse disposition function of oxycodone was best described by a one-compartment model. Absorption rate of the IR solution was best described by a mono-exponential model with a lag time, whereas absorption rate of the CR tablet was best described using a bi-exponential model. The absorption profile of the CR tablets was characterized by a rapid absorption component $(t_{1/2}$ abs=37 min) accounting for 38% of the available dose and a slow
absorption phase $(t_{1/2}$ abs=6.2 b) accounting for 62% of the available dose absorption phase $(t_{1/2}ab = 6.2 h)$ accounting for 62% of the available dose.
Two 10 mg tablets of and CP ayyordana hydrophlarida ware 102.7% Two 10 mg tablets of oral CR oxycodone hydrochloride were 102.7% bioavailable relative to 20 mg of IR oxycodone hydrochloride oral solution. The population model derived after administration of a single dose accurately predicted both the mean and range of oxycodone concentrations observed during 4 days of repeated dosing. The mean prediction error was 2.7% with a coefficient of variation of 54%.
- 4 The absorption characteristics of CR oxycodone tablets should allow effective plasma concentrations of oxycodone to be reached quickly and for effective concentrations to be maintained for a longer period after dosing compared with the IR oral solution. The CR dosage form has pharmacokinetic characteristics that permit 12 hourly dosing.

Keywords oxycodone controlled-release

Correspondence: Dr Jaap W Mandema, Alza Corporation, 950 Page Mill Road, PO Box 10950, CA 94303–0802, USA

A recent survey of anaesthetists in Finland found study. The methods used in this repeated-dose study parenteral oxycodone to be the most popular opioid for have been reported elsewhere [8]. premedication, postoperative pain relief, and sedation in intensive care units [1]. Similar to morphine, oxycodone is a potent opioid agonist with a dose Pharmacokinetic sampling dependent analgesic effect [2], that has proven effective in relieving postsurgical [3] and cancer-related pain Single dose During each treatment period, venous [4]. Side effects that have been observed following blood samples were obtained immediately before study [4]. Side effects that have been observed following blood samples were obtained immediately before study oxycodone administration are similar to those associated $\frac{d}{dx}$ drug was administered (0 h) and at the following oxycodone administration are similar to those associated drug was administered (0 h) and at the following times with other strong opioids.

Pharmacokinetic studies have revealed that after oral 3, 3.5, 4, 6, 8, 10, 12, 18, 24, 30, and 36 h. Plasma administration, oxycodone is rapidly absorbed to pro-
oxycodone concentrations were analysed by GC/MS administration, oxycodone is rapidly absorbed to pro-
duce an initial peak plasma oxycodone concentration using paltrexone as the internal standard. The quantifiduce an initial peak plasma oxycodone concentration using naltrexone as the internal standard. The quantifi-
in about 2 h [5, 6]. Once peak plasma concentrations cation limit of oxycodone in plasma samples was in about 2 h [5, 6]. Once peak plasma concentrations cation limit of oxycodone in plasma samples was are reached, oxycodone concentrations rapidly decline, 0.2 ng ml^{-1} , with standard curve linearity between are reached, oxycodone concentrations rapidly decline, 0.2 ng ml^{-1} , with standard curve linearity between with an apparent terminal half-life ranging from 3.0 to 0.2 ng ml^{-1} and 100 ng ml⁻¹. Inter-day and intra-da with an apparent terminal half-life ranging from 3.0 to 0.2 ng ml⁻¹ and 100 ng ml⁻¹. Inter-day and intra-day 5.7 h [5, 7]. Because oxycodone is rapidly absorbed precision (expressed as per cent coefficient of variatio 5.7 h [5, 7]. Because oxycodone is rapidly absorbed precision (expressed as per cent coefficient of variation) and quickly eliminated after oral administration, fre-
ranged from 0.4% to 9.9% for nominal standards and quickly eliminated after oral administration, fre-
quent dosing (every 4–6 h) is required to maintain ranging from 0.2 ng ml⁻¹ to 100 ng ml⁻¹. Oxycodone quent dosing (every 4–6 h) is required to maintain ranging from 0.2 ng ml⁻¹ to 100 ng ml⁻¹. Oxycodone plasma concentrations within the therapeutic analgesic analysis was performed by The Purdue Frederick plasma concentrations within the therapeutic analgesic analysis was performed by The Purdue Frederick range.

Recently, controlled-release oxycodone tablets have been developed to extend the duration of action of oral
oxycodone and provide the benefits of 12 hourly dosing.
In the present study, the pharmacokinetics of oxycodone
on days 2–4. In addition, serial blood samples for
con presented. The predictive performance of the models was evaluated by comparing the plasma concentrations predicted by the models with actual plasma concen-
 Pharmacokinetic analysis

blinded, two-way crossover design in 24 healthy, fasting,

male volunteers was used to compare the pharmaco-

kinetics of two 10 mg controlled-release oxycodone

tablets (OxyContin®, The Purdue Frederick Company, dosage f ment phases were separated by a 1 week washout interval. This study was approved by the ethics committee of the contract research organization Hazelton Pharmacokinetic modelling Wisconsin, Inc. All subjects provided written informed The plasma concentration-time profiles were charac- consent before participating in the study.

Repeated dosing Twenty-one healthy volunteers model: received one 10 mg controlled-release oxycodone tablet $C_p = f(t) * g(t)$ (1)
every 12 h and 5 ml of a 5 mg 5 ml⁻¹ immediate-release oxycodone oral solution every 6 h for 4 days in a where C_p is the plasma concentration at time t, $f(t)$ is randomized, analytically blinded, multiple-dose, two-
the absorption model, $g(t)$ is the unit impulse dispositio

Introduction way, crossover study. None of the volunteers who received repeated doses participated in the single-dose

th other strong opioids.

after dosing: 0.25, 0.5, 0.75, 1, 1.25, 1.5, 1.75, 2, 2.25, 2.5,

Pharmacokinetic studies have revealed that after oral 3, 3.5, 4, 6, 8, 10, 12, 18, 24, 30, and 36 h. Plasma Research Center, Yonkers, NY, USA.

trations measured after 4 days of repeated dosing.
The peak plasma concentration (C_{max}) and time to C_{max} (t_{max}) were read directly from individual plasma concentration-time curves. Area under the concentration-time **Methods** curve from 0 to 36 h $(AUC(0, 36 h))$ was calculated by the trapezoidal method. The elimination rate constant (λ_z) was calculated by measuring the slope of the linear Study design (λ_z) was calculated by measuring the slope of the linear regression line (log concentration vs time) in the terminal Single dose A randomized, single-dose, analytically elimination phase for each subject. The apparent blinded, two-way crossover design in 24 healthy, fasting,

terized with the following general pharmacokinetic

$$
C_p = f(t) * g(t) \tag{1}
$$

the absorption model, $g(t)$ is the unit impulse disposition

Disposition model The unit impulse disposition function was assumed to be the same for the immediate-release oral solution and the controlled-release tablet and was Model development characterized by a sum of exponentials:

$$
g(t) = \sum_{i=1}^{n} A_i e^{-\alpha_i t}
$$
 (2)

the unit impulse disposition function, respectively. that repeated measures are taken from each individual.

$$
f(t) = k_a e^{-k_a(t - t_{lag})}
$$
 (3)

where k_a is the first order absorption rate constant for to characterize the residual error:
the immediate release and collecting and to is a lag time. the immediate-release oral solution and t_{lag} is a lag time. $\log(Y_{ij}) = \log(Cp_{ij}) + \varepsilon$
This absorption profile is consistent with previous This absorption profile is consistent with previous
studies that compared the pharmacokinetics of immediate-release oral solution with intravenous oxycodone individual predicted by the pharmacokinetic model
[5, 7]. (equat

absorption models (similar to equation 2) to combi-
normally distributed, with mean zero and variance σ^2 . nations of multi-exponential and zero-order absorption models. Absorption of oxycodone from the controlled- Inter-individual variance model The inter-individual release tablets was best described by the following variability in the pharmacokinetic parameters was bi-exponential absorption model: modelled according to an exponential variance model

$$
f(t) = F_{\text{rel}}(f_1 k_{\text{c1}} e^{-k_{\text{c1}}(t - t_{\text{lag}})} + (1 - f_1) k_{\text{c2}} e^{-k_{\text{c2}}(t - t_{\text{lag}})})
$$
\n(4)

where F_{rel} is the relative bioavailability of oxycodone in the controlled-release tablet compared to the immediatethe controlled-release tablet compared to the immediate-
where p_i is the vector of pharmacokinetic parameters of release solution, k_{c1} and k_{c2} are the two first-order the *i*th individual, θ is the vector of population mean apparent absorption rate constants for controlled-release barmacokinetic parameters, and $\exp(\eta_i)$ expresses the oxycodone, and f_1 denotes the fraction of dose absorbed random difference between θ and p_i . Values of oxycodone, and f_1 denotes the fraction of dose absorbed random difference between θ and p_i . Values of η_i are 1 are 1 assumed to be independently multi-variate normally via the absorption process controlled by the rate assumed to be independently multi-variate normally constant k_{-1} . The fraction of the dose absorbed via the distributed, with mean zero and diagonal varianceconstant k_{c1} . The fraction of the dose absorbed via the distributed, with mean zero and diagonal variance-
absorption process controlled by the rate constant k_c covariance matrix Ω with diagonal elements (ω_1^2 is represented by $(1-f_1)$.

release tablet was compared with a nonparametric method in NONMEM. model derived from deconvolution of the plasma concentrations observed after administration of the Model validation To evaluate the predictive performcontrolled-release tablet and the unit impulse disposition ance of the models derived from single-dose adminisfunction derived from the immediate-release solution. tration of immediate-release and controlled-release The nonparametric absorption model was represented oxycodone, the plasma concentrations of oxycodone by a step-wise input model, one step for each interval after repeated administration for 4 days were compared between the following time points: 0, 0.25, 0.5, 1, 1.5, 2, with the concentrations predicted by the models. The 2.5, 3, 4, 6, 8, 10, 12, 18, 24, and 36 h. plasma oxycodone concentrations at 108 h after the

criterion $(P<0.01)$ and visual inspection of the fits. The the analysis because measurable concentrations were difference in −2 times the Log of the Likelihood only observed in a few individuals. Oxycodone plasma (−2LL) between a full and reduced model is asymptoti- concentrations in the remainder of the subjects were

function (i.e. the concentration vs time profile observed difference in number of parameters between the two after a unit intravenous dose), and * denotes convolution. models. A decrease of more than 6.6 in −2LL is significant at the $P < 0.01$ level.

The models were fitted to the data using the nonlinear regression program NONMEM version IV $[9, 10]$. Data were analysed using population analysis techwhere A_i and α_i are the coefficients and exponents of inques. The population analysis technique recognizes the unit impulse disposition function recognizes that repeated measures are taken from each individual The population pharmacokinetic model incorporates Absorption model The absorption model was assumed
to be different for the two preparations. It was assumed
that the absorption after administration of the immedi-
ate-release oral solution is rapid and could be described
b

Residual variance model The following model was used
to characterize the residual error:

$$
\log (Y_{ij}) = \log (C p_{ij}) + \varepsilon_{ij} \tag{5}
$$

Various absorption models were evaluated for the residual departure of $log(Y_{ij})$ from $log(Cp_{ij})$ is represented controlled-release tablet, ranging from multi-exponential by ε_{ij} . Values of ε_{ij} are assumed to be indep by ε_{ij} . Values of ε

> with the assumption that the pharmacokinetic parameters are log-normally distributed:

$$
p_i = \theta \cdot \exp(\eta_i) \tag{6}
$$

i
: assumed to be independently multi-variate normally absorption process controlled by the rate constant k_{c2} covariance matrix Ω with diagonal elements $(\omega_1^2, \dots, \omega_m^2)$. The values of the population parameters θ , σ^2 , The parametric absorption model for the co The parametric absorption model for the controlled- and Ω were estimated using the so-called first order

Model selection was based on the Log Likelihood start of treatment (last sample) were not included in cally χ^2 distributed with degrees of freedom equal to the below the quantification limit of the assay, and inclusion estimate of the mean prediction error. 76 kg (range 63–91), respectively.

The prediction error was calculated to evaluate how well the population pharmacokinetic models predicted the population mean response after 4 days of continuous administration. The log prediction error for a specific Pharmacokinetic analysis measurement was defined as:

$$
LPE_{ij} = \log(Y_{ij}) - \log(Cp) \tag{7}
$$

individual and C_p is the population mean concentration predicted by the pharmacokinetic model. The population release solution (89.5–115.9%), the two formulations mean response was obtained from a Monte Carlo were considered equally bioavailable (Table 1). The simulation of 2500 subjects using the parameter estimates AUC(0, 36 h) was not significantly different ($P > 0.05$) (mean and variance) of the population model. C_p for for the two dosage forms. However, differences in mean each time point was calculated directly from the C_{max} , t_{max} and apparent $t_{\frac{1}{2},z}$ were significant (P= simulated data as the logarithmic mean of all simulated 0.0001). Figure 1 illustrates the mean oxycodone concentrations. An estimate of the bias of the model plasma concentration vs time curve for the two dosage predictions was given by the average LPE. The variance forms. As would be expected with a controlled-release predictions was given by the average LPE. The variance forms. As would be expected with a controlled-release
of LPE indicates the variability of the measured formulation, the peak oxycodone plasma concentration of *LPE* indicates the variability of the measured formulation, the peak oxycodone plasma concentration concentration concentration concentration concentrations around the population mean prediction was lower after the con concentrations around the population mean prediction. The definition of prediction error according to equation the immediate-release solution and plasma levels 7 was preferred over the percentage prediction error. declined less rapidly after the peak plasma concen-7 was preferred over the percentage prediction error, declined less rapidly $(Y - C_n)$ 100/ C_n , because the individual plasma con- $(Y_{ii}-Cp_{ii})100/Cp_{ii}$, because the individual plasma concentration measurements and predictions tend to follow a logarithmic distribution rather than a normal distribution.

The Monte Carlo simulation of 2500 subjects using the parameter estimates (mean and variance) of the population model was also used to evaluate how well the population model predicted the range of observed concentrations after repeated dosing for 4 days.

Results

Subjects

Of the 24 volunteers enrolled in the single-dose study, 23 completed both study phases and were included in the bioavailability analysis. One subject wished to Figure 1 Mean plasma oxycodone concentration vs time withdraw from the study and was discontinued before profiles observed after a single dose of two 10 mg controlledthe second study phase. The mean age of subjects release oxycodone tablets $\ddot{\bullet}$ or 20 mg of immediate-release enrolled in the study was 30 years (range $21-45$). Mean oxycodone solution (\circ).

of these samples would have resulted in a biased height and weight were 179 cm (range 162–191) and

Relative to the immediate-release solution, controlled-
release oxycodone was 102.7% bioavailable. Based on
the 90% confidence interval for the AUC(0, 36 h) where Y_{ij} is the jth measured concentration in the *i*th the 90% confidence interval for the AUC(0, 36 h) individual and C_p is the population mean concentration ratio's of the controlled-release tablet and immediate

Table 1 Single-dose pharmacokinetics of immediate-release (IR) and controlled-release (CR) oxycodone

Parameter	Mean [*] \pm s.d. (n = 23)				
	20 mg CR oxycodone	20 mg IR oxycodone	$($ %) CR/IR	P†	90% confidence interval
AUC(0, 36 h) $(ng \, ml^{-1} h)$	$199.7 + 65.3$	$194.4 + 23.4$	102.7	NS	$89.5 - 115.9$
C_{max} (ng ml ⁻¹)	$18.6 + 6.1$	$41.6 + 13.2$	44.8	0.0001	$32.5 - 57.0$
$t_{\rm max}$ (h):	$2.62 + 1.07$	$1.30 + 0.63$	200.8	0.0001	169.8–232.6
Apparent $t_{\frac{1}{2},z}$ (h)	$7.99 + 2.96$	$3.21 + 0.87$	249.15	0.0001	$216 - 310.7$

*Arithmetic mean. †ANOVA test of significant difference (statistically significant = P < 0.05). \ddagger The range of t_{max} values was 1 to 6 h for CR oxycodone and 0.2 to 2.5 h for IR oxycodone.

order absorption model (equation 3) and a mono- the population analysis as well as the actual concenexponential unit impulse disposition function (equa- trations measured after administration of single doses tion 2) best described the concentrations observed after of controlled-release and immediate-release oxycodone. administration of the immediate-release oral solution. Figure 2 displays the fit for the first 8 h, highlighting the This is consistent with the mean concentration profile absorption phase. Figure 3 shows the fit over the entire observed in Figure 1. For the final analysis, the single-dose study period, highlighting the terminal disposition model was parameterized in CL/F and V_d/F , where CL denotes clearance, V_d is volume of distribution The cumulative fraction of drug absorbed from the details and F is the hispanilability of the and controlled places tablet following administration of a at steady-state, and F is the bioavailability of the oral controlled-release tablet following administration of a solution. The actual bioavailability of the immediate-
single dose is shown in Figure 4. The thick solid line release oral solution was not determined in this study represents the median fraction absorbed obtained from because an intravenous reference dose was not a Monte Carlo simulation of 2500 individuals using the evaluated. population mean pharmacokinetic parameters and their

Controlled-release Various absorption models were
evaluated for the controlled-release tablet, but only the
model that best described the data (as determined by
the Log Likelihood criterion and visual inspection) is
presen release tablets was best described by the bi-exponential absorption model described by equation 4. This model was preferred over a mono-exponential, zero-order drug input, or mono-exponential combined with zero-order drug input. The controlled-release data were best described using the same lag time as for the immediaterelease oral solution.

Results of the population analysis are presented in Table 2. The inclusion of separate residual variance terms for the immediate-release solution and controlledrelease tablets in the analysis did not significantly improve the fit of the models.

Table 2 Pharmacokinetic parameter estimates for a single 20 mg dose of immediate-release oxycodone oral solution and two 10 mg controlled-release oxycodone tablets

	Population analysis			
Parameter*	Mean	Variance		
CL/F (1 h ⁻¹)	110(5)	0.045(0.014)		
$V_{d}/F(1)$	593 (32)	0.061(0.017)		
k_a (h ⁻¹)	4.19(0.58)	1.12(0.38)		
$\boldsymbol{F}_{\textrm{rel}}$	1.02(0.09)	0.15(0.04)		
k_{c1} (h ⁻¹)	1.11(0.21)	0.25(0.14)		
k_{c2} (h ⁻¹)	0.110(0.009)	0.23(0.09)		
f_{1}	0.38(0.02)	0.053(0.043)		
	0.206(0.007)	0.023(0.012)		
t_{lag} (h) σ^2	0.064(0.005)			

, 105% for k_a , 38% for F_{rel} , 50% for k_a 48% for k_{c2} , 23% for f_1 approximately equal to the percentage of residual variability a one-compartment disposition model following which is 25% for the population analysis. and administration of the controlled-release tablets.

Pharmacokinetic model **Figures** 2 and 3 show oxycodone concentrations predicted by pharmacokinetic models using the typical Immediate-release The combination of a simple, first-values of the pharmacokinetic parameters derived from elimination phase.

inter-individual variances. The interval in which 80% of

*Based on 912 concentration measurements from 23
Figure 2 Fit of concentrations predicted by the the actual plasma
harmacokinetic models (solid line) to the actual plasma individuals. †The square root of the variance is

approximately equal to the percentage of inter-individual

variability in the pharmacokinetic parameters which is 21%

for CL, 25% for V_d , 105% for k_a , 38% for F_{rel} , function with a one-compartment disposition model following
administration of the immediate-release oral solution. (b) fit of the model linking the bi-exponential absorption function with

Figure 3 Fit of concentrations predicted by the pharmacokinetic models (solid line) to the actual plasma concentrations $(•)$ during the first 36 h following a single dose: (a) fit of the model linking the monoexponential absorption function with a one-compartment disposition model following administration of the immediaterelease oral solution. (b) fit of the model linking the bi-exponential absorption function with a one-compartment disposition model to the actual concentrations following administration of the controlled-release tablets.

controlled-release tablet. The solid line in this figure population mean oxycodone concentrations (solid lines) simulation of 2500 individuals using the mean release oxycodone solution every 6 h (a) or one 10 mg indicate the interval between which 80% of the simulated 2500 subjects using the parameter estimates from the data lies (between the 10th and 90th percentiles). population analysis following a single dose.

tion pattern derived using the non-parametric absorption model (deconvolution data not shown). The nonparametric absorption model, using 11 more parameters, resulted in only a minor improvement of the fit $(6.1 \text{ points decrease in } -2LL)$, confirming that the derived bi-exponential absorption model is a good description of the absorption profile of the controlledrelease tablet.

Predictive performance

The mean and variance of the prediction errors were −0.0061 and 0.291, respectively, for the immediate release solution and 0.0335 and 0.287 for the controlled release tablet. The exponent of the mean LPE indicates that compared with the actual mean plasma concentrations, predicted concentrations were on average 0.6% higher for the immediate-release solution and 3.3% lower for the controlled-release tablets.

Figure 5 shows the comparison of the mean oxyco-

Figure 5 Comparison of the observed population mean Figure 4 Cumulative amount of drug absorbed from the oxycodone concentrations (symbols) and the predicted represents the median absorption profile obtained from a during 4 days of repeated dosing with 5 mg of immediatepharmacokinetic parameter estimates with associated controlled-release oxycodone tablet every 12 h (b). Model variances from the population analysis. The thin lines predictions were obtained by a Monte Carlo simulation of

done plasma concentrations observed during the 4 days Safety of repeated dosing and the population mean concentrations predicted by the Monte Carlo simulations using Seven subjects reported a total of 31 adverse experiences the parameters from the population analysis (all means after taking controlled-release oxycodone, and 12 subare logarithmic means). Figure 6 shows the range of jects reported a total of 55 adverse experiences after concentrations predicted by the Monte Carlo simu- receiving the immediate-release oxycodone solution lations in comparison with the actual observed concen- (Table 3). Most adverse experiences involved the gastrotrations. The thin lines cover the region in which 90% intestinal tract or central nervous system. None of the of the simulated data lies (5th–95th percentile). The subjects required medical treatment for an adverse population model predicts the distribution of observed experience. There were no medically significant changes concentrations quite accurately. The simulated 5th in results of physical examinations or abnormal clinical percentile actually includes 9.7% of the observed laboratory findings attributed to study medication. concentrations for the immediate-release oral solution, There were no medically significant changes in vital whereas the simulated 95th percentile includes 93.8% of signs observed during the study. the observed concentrations (84% of the observed concentrations fall within the two lines). The values for the controlled-release tablets are 6.8% and 98.2%, respectively (92% of the observed concentrations fall Discussion within the two lines).

by fitting the pharmacokinetic models to the concen- done administered as a controlled-release tablet and tration data observed after repeated administration. compared it with immediate-release oxycodone oral Except for the apparent absorption rate constant of the solution. A simple one-compartment model with firstimmediate-release solution, all the pharmacokinetic order absorption best described the oxycodone plasma parameters obtained after 4 days of repeated adminis- concentration vs time profile of the immediate-release tration were similar to the parameters obtained after oral solution. Absorption of oxycodone from the immedisingle-dose administration. This is consistent with the ate-release solution was rapid, resulting in a high initial ability of the pharmacokinetic model derived after a peak concentration. After the peak oxycodone concensingle dose to predict plasma concentrations during 4 tration was reached, plasma concentrations declined days of repeated dosing. The model fitting also revealed rapidly. The pharmacokinetic profile of immediate-release that the variability of the observed concentrations oxycodone obtained in this study was consistent with around the predicted concentrations was similar after those that have been previously reported [11, 5]. single-dose and repeated-dose administration, 47% and In the present study, oxycodone clearance (CL/F) 53%, respectively. following administration of the immediate-release

Pharmacokinetic parameter estimates were derived We have characterized the pharmacokinetics of oxyco-

Number of subjects* (number of reports) Body system Adverse experience Controlled-release Immediate-release Body Headache $4 (9)$ $3 (12)$ Cardiovascular Pallor 0 1 (2) Syncope $1 (1)$ 1 (1) 1 (1) Vasodilatation $1 (2)$ 0 (0) Digestive Nausea $3(6)$ 6 (12) Vomiting $1 (1)$ $3 (6)$ Nervous Dizziness 6 (10) 11 (18) Paresthesia 1 (1) 0 Somnolence 0 1 (1) Respiratory Hiccup 0 1 (1) Skin Sweat $1 (1)$ $2 (2)$ Total $7(31)$ 12 (55)

Table 3 Adverse experiences following a single dose of controlledrelease oxycodone tablets or immediate-release oral solution

*13 subjects reported one or more adverse experiences. †All adverse events were considered possibly or probably related to study medication.

Figure 6 Actual (\bullet) and predicted oxycodone concentrations (solid lines) during the last day of repeated dosing with 5 mg of immediate-release oxycodone solution every 6 h (a) or one 10 mg controlled-release oxycodone tablet every 12 h (b). The thick line is the population mean response obtained from a Monte Carlo simulation of 2500 subjects using parameter estimates from the population analysis following a single dose. The thin lines cover the region in which 90% of the simulated data lies (5th–95th percentile).

reported by Pöyhiä et al. $[11]$ and Leow et al. $[5]$. model not only characterizes the population mean However, the differences in oxycodone clearance between response, but also characterizes the variability in plasma this study and previous studies were found to be of no drug concentrations between individuals by estimating clinical significance when the data were adjusted for the inter-individual variability in the pharmacokinetic body weight $(1.1-1.4 \lg^{-1} h^{-1})$. parameters. The characterization of the distribution of

release tablets was equivalent to that of the immediate- is crucial for the design of dosage regimens that will release oral solution. This is supported by the fact that ensure that most of the patients receive a safe and both the nonparametric analysis and the parametric efficacious treatment. analysis produced similar estimates of bioavailability for The variability in plasma concentrations for both controlled-release oxycodone relative to immediate- immediate-release and controlled-release oxycodone was release oxycodone (Tables 1 and 2). within the range expected for oral opioids. The average

was best described using a bi-exponential absorption around the mean were approximately 47% after adminismodel, with a rapid initial absorption component tration of a single dose and 53% after 4 days of repeated $(t_4$ abs = 37 min) accounting for 38% of the available dose followed by a slow absorption phase $(t_4$ abs = 6.2 h) between the two formulations. Population analysis $\frac{1}{2}$ absent the two formulations. Population analysis $\frac{1}{2}$ between the two formulations. Population an accounting for 62% of the available dose. indicated that about 70% of the variability in plasma

phase of controlled-release oxycodone was smaller than individual variability in pharmacokinetic parameters. the elimination rate constant of oxycodone observed The greatest variance occurred in those parameters from the immediate-release oral solution data. describing the absorption processes. The Monte Carlo Consequently, a slower terminal decline in oxycodone simulations showed that the population model derived plasma concentrations was observed with the controlled- from the single dose data accurately predicts the release tablets compared with the immediate-release distribution of oxycodone plasma concentrations during solution. This should allow for a prolonged duration of repeated dosing for four days. The fact that the analgesic activity. At the same time, the initial rapid population model is able to predict both the mean and absorption phase for controlled-release oxycodone pro- range of oxycodone plasma concentrations during 4 duced a rapid rise in oxycodone plasma concentrations days of consecutive dosing clearly supports the validity that should ensure a rapid onset of action. and usefulness of the proposed population model and

Lower peak oxycodone concentrations after dosing estimated population parameters. with controlled-release oxycodone were associated with Comparable plasma concentration profiles were fewer side effects compared with dosing with immediate- obtained with 10 mg controlled-release tablets every release solution at the same total dose. However, these 12 h and 5 mg immediate-release solution every 6 h, differences were not statistically significant in the data suggesting that either formulation could be used for we examined, and whether or not higher peak plasma long-term pain management. However, there are fewer concentrations observed with immediate-release oxyco- peak/trough oscillations and less fluctuation in plasma done were the cause of the increased number of side concentrations during steady-state dosing with the effects can not be determined. controlled-release tablets. The controlled-release formu-

model to characterize the plasma oxycodone profile administration. after administration of the controlled-release tablet is In conclusion, the pharmacokinetics of a newly confirmed by the good fit to the data (Figures 2 and 3) developed controlled-release oxycodone tablet have been and the close agreement between the bi-exponential characterized and compared with an immediate-release absorption model and the step-wise input profile oral solution. The absorption characteristics of the determined nonparametrically by deconvolution. The controlled-release tablet should allow effective plasma validity of the proposed pharmacokinetic models is also concentrations of oxycodone to be reached quickly, and confirmed by the ability of the models to predict the for effective concentrations to be maintained for a longer plasma concentrations in a group of different subjects period after dosing compared with an immediate-release during 4 days of consecutive dosing with the immediate- oral formulation, thus allowing dosing every 12 h. release oral solution (5 mg every 6 h) and controlledrelease tablets (10 mg every 12 h). The population model This study was supported by The Purdue Frederick Company, was able to predict the population mean response in Norwalk, CT, USA. Results included in this paper have b was able to predict the population mean response in Norwalk, CT, USA. Results included in this paper have been
these new individuals accurately Indeed considering presented previously at the annual meetings of the American these new individuals accurately. Indeed, considering
the inter-individual variability observed after adminis-
tration of each of the dosage forms, the predictive and Society, November 1994. performance of the proposed models is very good, with a bias of less than 3.5% in predictions of mean References oxycodone concentrations based on parameter estimates from the population analysis. Figure 5 shows the close \qquad 1 Pöyhiä R. Opioids in anaesthesia: A questionnaire survey relationship between the observed and predicted mean in Finland. Eur J Anaesthesiol 1994; 11: 221–230.

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The absorption profile of the controlled-release tablet coefficients of variation of the plasma concentrations dosing. There was no difference in the overall variability The absorption rate constant for the slow absorption oxycodone concentrations could be attributed to inter-

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