

population pharmacokinetics of phenytoin

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The inter-individual variability of K_M CV. The population pharmacokinetics was similar in children and adults. K_{max} and K_M were estimated to be. This is a preview of subscription content, log in to check access. Clin Pharmacol Ther
Population pharmacokinetics of phenytoin in Singapore Chinese. Ther Drug Monit 1: Authors Authors and affiliations
E. K_{max} was higher than reported values, and K_M was comparable to that reported in a study in Japanese, but was
much lower than that reported in studies of European patients. Cite article How to cite? Unable to display preview.
European Journal of Clinical Pharmacology. Analysis of serum concentration data in patients treated with mexiletine
and lidocaine. There was no age or gender-related effect on either the apparent maximum elimination rate k_{max} or
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Baragwanath Hospital, Johannesburg, steady-state serum phenytoin concentrations were. Chem Pharm Bull (Tokyo).
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Higuchi S, Aoyama T. Author information: (1)Department of Hospital Pharmacy, Faculty of Medicine, Kyushu
University, Fukuoka, Japan. Routine clinical pharmacokinetic. OBJECTIVE: To study population pharmacokinetics of
phenytoin in pediatric patients by using sparse data. METHODS: We used routinely collected therapeutic drug
monitoring data, derived from the steady state serum concentrations of phenytoin in 42 pediatric outpatients with
epilepsy. Depending on whether the patients. The pharmacokinetics of phenytoin was studied in 66 epileptic Chinese
children and adults. The data were analysed by the population approach, using the non-linear mixed effect model, in the
MULTI (ELS) program. There was no age or gender-related effect on either the apparent maximum elimination rate
(k_{max}) or. Routine clinical pharmacokinetic data collected from out-patients who received phenytoin were analysed to
estimate population pharmacokinetic parameters. There were steady-state phenytoin concentrations and associated
dosage rates (mg/day) from out-patients. The data were analysed using NONMEM, a. The population pharmacokinetic
parameters of phenytoin were estimated using routine therapeutic drug monitoring data from epileptic patients. The
serum concentration values at steady-state after repetitive oral administration were analyzed using JavaPK program and
Bayesian feedback method. The maximal. Oct 16, - Abstract. The objective was to study the population
pharmacokinetics of bound and unbound phenytoin in critically ill children, including influences on the protein binding
profile. A population pharmacokinetic approach was used to analyze paired protein-unbound and total phenytoin plasma
concentrations.