Olmesartan medoxomil-loaded self-nanoemulsifying drug delivery systems: design, in-vitro characterization, and pharmacokinetic assessments in rabbits via LCóMS/MS

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Abstract

Olmesartan medoxomil (OLM) is a lipophilic (log P = 4.31) antihypertensive drug suffering from limited oral bioavailability in humans (26%) due to its low aqueous solubility, uncontrolled enzymatic conversion to the active metabolite (Olmesartan; OL) and efflux by drug resistance pumps. Surmounting such limitations via incorporation of OLM into self-nanoemulsifying drug delivery systems (SNEDDS). Based on OLM-equilibrium solubility studies in various oils, surfactants and couwthcevcpvu."Ecrown Ì "OEO."Vyggp Ì "42."Etgoqrjqt Ì "GN"cpf"rqn{gvj {ngpg"in{eqn} - 400 (PEG) were combined in different ratios to plot ternary phase diagrams. OLMloaded SENDDS were developed and evaluated forparticle size, polydispersity index (PDI), zeta potential, self-emulsification time, morphology, drug released percentages after 5-min (Q5min%), 1 hour (Q1h%) and dissolution efficiency rgtegpvcigu"*FG3j '+0"Vjg"QN"rjctoceqmkpgvkeu"htqo"UPGFFU"*H8+"cpf"DgpkectÌ' tablets were evaluated (LC-MS/MS) in rabbits. Spherical OLM-loaded SNEDDS were developed. The best-achieved SNEDDS (F6) showed short emulsification time (13 s), fine droplet size (60.00nm), low PDI (0.25), negative zeta potential (-14.4mV), promising dissolution parameters; Q5min% (29.78%), Q1h% (66.69%) and DE1h% (47.96%) and enhanced in vivo absorption characteristics; shorter Tmax, higher Cmax and larger AUC(0 6:h; suggesting its potential for the enhancement of the oral absorption of practically insoluble drugs; like OLM.

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