Dossier Requirement

Evaluation	NDA	ANDA	OTC Monograph Drug Application
Reference Drug	Not required	Required	Compiled with Monograph
Safety Efficacy	Pharm/ToxPK/PD/BA/BEClinical trials	Bioequivalence (BE) as a surrogate to clinical trial	Not required
Quality	 Chemistry, Manufacturing and Controls(CMC) PIC/S GMP GLP, GCP 		
Labeling	Labeling (direction of use)		



Registration Procedures of OTC Drugs



