

Maldives Food and Drug Authority

Ministry of Health

Male', Maldives

Guideline for Product Registration Including Emergency Use Authorization

Medicine and Therapeutic Goods Division, Maldives Food and Drug Authority			Document Created on: 09.08.2	022
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SUMMARY OF CHANGES

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ABBREVIATIONS

ADL	Approved Drugs List
API	Active Pharmaceutical Ingredient
ВЕ	Bioequivalence
ВР	British Pharmacopeia
СЕР	Certification of Suitability
COA	Certificate of Analysis
СОРР	Certificate of Pharmaceutical Product
CTD	Common Technical Document
DP	Drug Product
DS	Drug Substance
EUA	Emergency Use Authorization
FPP	Finished Pharmaceutical Product
GMP	Good Manufacturing Practices
ICH	International Conference on Harmonization
INN	International Non-Proprietary Name
МА	Market Authorization
МАН	Market Authorization Holder
MDI	Metered Dose Inhaler
MFDA	Maldives Food and Drug Authority
MTG	Medicine and Therapeutics Goods Division
NPB	National Pharmaceutical Board
NRA	National Regulatory Authority
PH. EUR	European Pharmacopeia

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PIL	Patient Information Leaflet
PP	Primordial Products
PSUR	Periodic Safety Update Report
RMP	Reference Medicinal Product
SMPC	Summary of Product Characteristics
SRA	Stringent Regulatory Authority
USP	United States Pharmacopeia
WHO	World Health Organisation
NDMA	N-nitroso dimethylamine (NDMA)
EUDRA	European Union Drug Regulatory Authorities

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Definitions

Active Pharmaceutical	a. A substance or mixture of substances intended to be used in the			
Ingredient (API) / Drug	manufacture of a pharmaceutical dosage form and that, when used so,			
Substance	becomes an active ingredient of that pharmaceutical dosage form. Such			
	substances are intended to furnish pharmacological activity or other			
	direct effect in the diagnosis, cure, mitigation, treatment or prevention			
	of disease or to affect the structure and function of the body.			
	b. A substance or compound that is intended to be used in the			
	manufacture a drug product as a pharmacologically active compound			
	(ingredient)			
Applicant	The person or Company who submits a registration application or dossier of			
	a product to the Authority and is responsible for the product information,			
	recall etc., availability.			
Adverse Drug Reaction	A response to a pharmaceutical product that is harmful and unintended and			
	that occurs at doses normally used or tested in humans for prophylaxis,			
	diagnosis, or treatment of disease, or for the modification of physiological			
	function			
Approved Drug List	A list of all medicinal products approved as drug product for use in Maldives.			
Authority	Authority means Maldives Food and Drug Authority (MFDA)			
	A defined quantity of raw material, packaging material, or finished			
Batch	pharmaceutical product processed in a single process or series of processes			
	so that it is expected to be homogeneous. It may sometimes be necessary to			
	divide a batch into a number of sub-batches, which are later brought together			
	to form a final homogeneous batch. In the case of terminal sterilization, the			
	batch size is determined by the capacity of the autoclave. In continuous			
	manufacture, the batch must correspond to a defined fraction of the			
	production, characterized by its intended homogeneity. The batch size may			
	be defined either as a fixed quantity or as the amount produced in a fixed			
	time interval			

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-	1996			
Bioavailability	The extent to which, following administration of a medicine, fraction of the			
a	active form of a drug that reaches systemic circulation unaltered to exert an			
e	effect.			
Bio-equivalence	Two pharmaceutical products are considered bioequivalent if they are			
p	pharmaceutically equivalent to their pharmaceutical alternatives, and their			
b	bio-availabilities (rate and extent of availability), in terms of peak (Cmax and			
Т	Tmax) and total exposure (area under the curve (AUC)) after administration			
C	of the same molar dose under the same conditions. They are similar to such			
a	a degree that their effects can be expected to be essentially the same.			
Certification of Analysis	It is a document that describes the list of tests applied to a particular sample			
v	with the result obtained and the acceptance criteria applied. It indicates			
v	whether the sample complies with the specifications.			
P	An authoritative document showing the results of analysis of a particular			
p	product batch.			
Certification of Suitability	A certificate that certifies compliance of the active pharmaceutical			
ii	ingredients/ Drug substances or pharmaceutical ingredients as per the			
r	monograph of the European Pharmacopoeia (EP)			
Collaborative	Collaborative Registration procedure to accelerate the national registration			
Registration procedure	of prequalified pharmaceutical products and vaccines, or the collaborative			
(CRP)	procedure to accelerate the national registration of products approved by			
S	stringent regulatory authorities (10, 11). The collaborative registration			
p	procedures cover initial registrations and post-registration variations/ post-			
а	approval changes.			
Composition	Composition in relation to a medicinal product means the ingredients of			
v	which it consists, and the proportions, degree of strength, quality, and purity			
C	of those ingredients			

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	The undesired introduction of impurities of a chemical or microbiological	
Contamination	nature, or of foreign matter, into or on to a starting material or intermediate	
	during production, sampling, packaging, or repackaging, storage or transport.	
Container Closure System	A primary container closure system is a packaging component (for example,	
	a vial) that is in, or may come into, direct contact with the final product dosage	
	form, or components that contribute to the container/closure integrity of the	
	primary packaging material for a sterile product.	
	A secondary container closure system is a packaging component (for example,	
	a carton) that is not, and will not be, in direct contact with the dosage form.	
Dosage Form	Formulation of an active ingredient(s) so that it can be administered to a	
	patient in specified quantity, strength, e.g., tablets, capsules, injection	
	solution, syrups, ointments, suppositories, etc.	
Dossier	A detailed compilation of documents generated from the product	
	manufacturer for the purpose of pharmaceutical product registration.	
Drug Product / Finished	Finished Drug Product or drug product or Medicinal Product means a finished	
Drug Product / Medicinal	dosage form that has undergone all stage of manufacturing including	
Product	packaging in its final container and labelling. FPP may contains one or more	
	active pharmaceutical ingredient / drug substance.	
Emergency Use	A risk-based procedure for assessing and listing unlicensed medicines and	
Authorization / Approval	vaccines with the ultimate aim of expediting the availability of these products	
	in the public health emergency.	
Evaluation	Assessment of submitted dossier for product registration based on	
	parameters of safety, efficacy and quality.	
Expiry Date	The date given on the individual container (usually on the label) of a drug	
	product up to and including which the product is expected to remain within	
	specifications, if stored correctly. It is established for each batch by adding	
	the shelf-life period to the date of manufacture. Source: World Health	
	Organization WHO Technical Report Series, No. 863, 1996	
Excipient / Non-active	A substance or compound, other than the API and packaging materials, that	
ingredient	is intended or designated to be used in the manufacture of a FPP. It also	

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	means any component of a finished dosage form that has no therapeutic		
	value.		
Formulation of Medicine	The validated composition of the finished pharmaceutical products i.e active		
	pharmaceutical ingredients/ drug substance and excepient(s).		
	A unique name identifying a particular pharmaceutical substance		
Generic Name	Generic names are officially assigned by international medicines		
Generic Name	nomenclature commissions, and nowadays mostly conform to those assigned		
	by the WHO program on the selection of INNs.		
Contain fortula			
Good Manufacturing	Good Manufacturing Practices is the aspect of quality assurance that ensures		
Practices	that medicinal product(s) are consistently manufactured and controlled to		
	the quality standards appropriate to their intended use and as required by		
	the product specifications.		
Innovator Drug Product	Finished drug (pharmaceutical and biological) products that are first		
	authorized for marketing globally (normally as a patented product) based on		
	parameters of efficacy, safety, and quality.		
International	The shortened scientific name (also known as the generic name) of a		
Nonproprietary Name	pharmaceutical substance assigned by the WHO program on the selection of		
	INNs, the INN is recognized worldwide.		
Label	A printed text attached to or comprising part of a medicine container or		
	package (primary and secondary excluding any outer shipping container),		
	specifying the name, dosage form, composition, batch number,		
	manufacturing date, and expiry date of the contents as well as the name and		
	address of the manufacturing company and/or importer of the product, the		
	product license holder, the permitted retail price, and other relevant		
	information (e.g., recommended storage conditions).		
Manufacturer	A company that carries out any of the operation of manufacturing, packaging,		
	labeling, quality control, final product release and quality assurance of the		
	products.		
Market Authorization/	An official document issued by the competent medicines regulatory authority		
Registration	for the purpose of marketing or free distribution of a product in Maldives		
negisti ation	To the purpose of marketing of free distribution of a product in Maidives		

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	after evaluation for safety, efficacy, and quality. It must set out, inter alia, the		
	name of the product, the pharmaceutical dosage form, the quantitative		
	formula (including excipients) per unit dose (using INNs or national generic		
	names, where they exist), the shelf life and storage conditions, and the		
	packaging characteristics. It specifies the information on which authorization		
	is based (e.g., "The product(s) must conform to all the details provided in your		
	application and as modified in subsequent correspondence"). It also contains		
	the product information approved for health professionals and the public, the		
	sales category, the name and address of the holder of the authorization, and		
	the period of validity of the authorization. Once a product has been given		
	marketing authorization, it is included on a list of authorized products (the		
	register), and is often said to be "registered" or to "have registration." Market		
	authorization may occasionally also be referred to as a license or product		
	license		
	(Please note that the terms market authorizations and registration is used		
	interchangeably in this document.)		
Market Authorization	The duration in which the applicant is allowed to manufacture, import,		
Validity	distribute and market or sell the product in Maldives after being granted		
	Market Authorization.		
Marketing Authorization	The local representative and/or applicant / firm that has the authorization to		
Holder	manufacture / import and/or market a medicinal product in Maldives. It also		
	refers to a person or legal entity allowed to apply for a change to the		
	marketing authorization or registration.		

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Medicinal Products /	Any substance or combination of substances marketed or manufactured to		
Drug Product	be marketed for treating or preventing disease in human beings, or with a		
	view to making a medical diagnosis in human beings, or to restoring,		
	correcting, or modifying physiological functions in human beings. (WHO PQ		
	definition).		
	Any substance or combination of substances presented for treating or		
	preventing disease in human beings. Any substance or combination of		
	substances which may be administered to human beings with a view to		
	making a medical diagnosis or to restoring, correcting, or modifying		
	physiological functions in human beings is likewise considered a medicinal		
	product. (The EU 2001/83/EC Directive)		
Method validation	The documented process by which an analytical procedure (or method)		
	provides a high degree of assurance that a specific process will consistently		
	result in a product that meets its predetermined specifications and quality		
	characteristics.		
National Pharmaceutical	A board assigned by Regulation on National Pharmaceutical Board 2019/R-		
Board	135 to provide technical advice on regulating medicine and medicinal		
	products.		
National Regulatory	Authority responsible for ensuring medicinal products released for public		
Authority	distribution are evaluated properly and meet international standards of		
	safety, efficacy and quality.		
Packaging Material	Any material, including printed material, used in the packaging of a		
	pharmaceutical product, excluding, any outer packaging used for		
	transportation or shipment and packaging materials are referred to as		
	primary or secondary according to whether or not they are intended to be in		
	direct contact with the product.		
Pharmacopeia	A publication issued by an authorized national or international commission /		
	body that specifies quality standards and other properties of pharmaceutical		
	substances and dosage forms		

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Pharmacovigilance	Pharmacovigilance is the science and activity relating to the collection,
	detection, assessment, monitoring, and prevention of adverse effects with
	pharmaceutical and biological drug products.
Post-market surveillance	Set of activities following the market authorization of a drug product including
	maintenance of product authorization and/or registration of variations or
	renewals; regular inspections of manufacturers, wholesalers, distributors,
	and retailers; quality control testing; pharmacovigilance; promotion control;
	public reporting of poor-quality products; handling of market complaints; and
	removal and disposal of non-compliant products.
Primordial Products	These are the medicinal products that have been on Approved Drug List from
	the beginning as approved products but not registered with a full dossier
	submission. These products are indicated with the letters "PP" in ADL.
Prescription Only	Medicines that may only be made available to the consumer through a
Medicines	written order signed by a duly qualified and registered medical prescriber and
	dispensed by a registered pharmacist.
Over-the-counter	Medicines that are generally regarded as safe for the consumer to use by
medicines	following the required label directions and warnings, and which may be
	purchased without a prescription
Pilot scale batch	A batch of an API or FPP manufactured by a procedure fully representative of
	and simulating that to be applied to a full production-scale batch, for
	example, for solid oral dosage forms, a pilot scale is generally, at a minimum,
	one-tenth that of a full production scale for 100,000 tablets or capsules,
	whichever is larger, unless otherwise adequately justified
Primary batch	A batch of an API or FPP used in a stability study, from which stability data are
	submitted in a registration application for the purpose of establishing a re-
	test period or shelf life, as the case may be. A primary batch of an API should
	be at least a pilot-scale batch. For an FPP, two of the three batches should be
	at least pilot-scale batches, and the third batch may be smaller if it is
	representative of the critical manufacturing steps. However, a primary batch
	may be a production batch.

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Reference Regulatory	A national or regional authority or a trusted institution as adopted by the				
Authorities	Maldives Food and Drug Authority for the purpose of reliance registration				
	pathways.				
	Reliance is the act whereby the regulatory authority in one jurisdiction takes				
Reliance	into account and gives significant weight to assessments performed by				
	another regulatory authority or trusted institution, or to any other				
	authoritative information, in reaching its own decision. The relying authority				
	remains independent, responsible and accountable for the decisions taken,				
	even when it relies on the decisions, assessments and information of others.				
	Full reliance means that the authority relies on the entire				
	assessments/inspection and quality control reports performed by another				
	NMRA. Partial reliance means that the authority relies on certain				
	documents/parts of the assessments performed by another NMRA, while for				
	the other part(s) an independent, full assessment of the documentation				
	submitted by the Applicant is conducted.				
	NRA remains independent, responsible, and accountable regarding the				
	decisions taken, even when it relies on the decisions and information of				
	others				
Registered Products	These are medicinal products that are registered and approved with full				
	dossier submission. These products are indicated with the letter "R" in ADL.				
Registration Number	A number assigned to a medicinal product after being given marketing				
	authorization.				
Comparator / Reference	A medicinal product that has been authorized for use on the basis of a full				
Medicinal Product	dossier, including the results of pre-clinical tests and clinical trials. Such				
	products are used as a comparator for the demonstration of the safety,				
	efficacy and quality of a generic drug product seeking marketing				
	authorization.				
Stability	The capacity of drug substance or drug product to remain within specification				
	established to ensure its identity, strength, quality, and purity.				

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Stability study	The evidence on how the quality of a drug substance or drug product varies
	with time under the influence of a variety of environmental factors such as
	temperature, humidity, and light and to establish a re-test period for the drug
	substance or a shelf life for the drug product and recommended storage
	condition.
	Long-term and accelerated (and intermediate) studies undertaken on primary
	and/or commitment batches according to a prescribed stability protocol to
	establish or confirm the re-test period (or shelf life) of an API or the shelf life
	of an FPP
Stability tests (protocol)	A series of tests designed to obtain information on the stability of a
	pharmaceutical product in order to define its shelf life and utilization period
	under specified packaging and storage conditions.
Summary of Product	A regulatory document of a medicinal product as it serves as the basis of
Characteristics	information for healthcare professionals regarding the use of drug products,
	ensuring their safety, efficacy, and quality.
Ongoing Stability	The shirth couried but by the manufacturing or much stick between according
Ongoing Stability	The study carried out by the manufacturer on production batches according
	to a predetermined schedule in order to monitor, confirm, and extend the
	projected re-test period (or shelf life) of the API, or confirm or extend the
	shelf life of the FPP.
Accelrated Stability	Studies designed to increase the rate of chemical degradation and physical
Studies	change of an API or FPP by using exaggerated storage conditions as part of
	the stability testing program. The data thus obtained, in addition to data
	derived from long-term stability studies, may be used to assess long-term
	chemical effects under accelerated conditions and to evaluate the impact of
	short-term excursions outside the label storage conditions, as might occur
Real time/	during shipping. The results of accelerated testing studies are predictive of
Ongoing Stability	tentative shelf life of the drug product.

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Studies	Such studies are designed to simulate the rate of chemical and/or physical				
	degradation of an active ingredient or dosage form or product, under				
	exaggerated storage conditions.				
	Experiments on the physical, chemical, biological, biopharmaceutical, and				
	microbiological characteristics of an API or FPP, during and beyond the				
	expected shelf life and storage periods of samples under the storage				
	conditions expected in the intended market. The results are used to establish				
	the re-test period or the shelf life, to confirm the projected re-test period or				
	shelf life, and to recommend storage conditions.				
Specification	A list of tests, references to analytical procedures, and appropriate				
	acceptance criteria which are numerical limits, ranges, or other criteria for				
	the tests described. It establishes the set of criteria to which any drug				
	substance or any drug product should conform to be considered acceptable				
	for its intended use.				
Storing	The storage of drug products according to the different storage conditions for				
	different drug substances according to their individual requirements.				
Storage condition	The storage condition that guarantees the maintenance of the quality of the				
	product in relation to its safety, efficacy, and acceptability throughout the				
	shelf life, as may be predicted from the stability studies. The described				
	conditions should indicate the temperature or temperature range in degree				
	Celsius, as well as humidity, light, and other relevant conditions.				
Strength	Strength of the medicinal product means the content of the active ingredient				
	expressed quantitatively per dosage unit, per unit of volume or mass or				
	weight, according to the dosage form				
Stringent Regulatory	National Regulatory Authorities that are recognized by WHO as having				
Authority	stringent regulatory practices.				

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Variation	A change to any aspect of a pharmaceutical product safety, efficacy and
	quality including but not limited to any change including but not limited to
	starting material, formulation (API/DS and exipients), method and site of
	manufacture, specifications for the finished product and ingredients,
	container, labeling, product information etc.
Validation	Documented act of proving that any procedure, process, equipment,
	material, activity, or system works correctly and actually leads to the
	expected results.
Variation Application	A variation application is an application for any intended change to already
	approved conditions of an existing registered product which has been
	previously registered as per the criteria laid down by MFDA
PIC/S	The Pharmaceutical Inspection Co-operation Scheme (PIC/S) are
	organizations that play a key role in pharmaceutical regulation and the
	inspection of pharmaceutical manufacturing facilities.
DEG	Diethylene Glycol is a toxic chemical that is used primarily as a solvent,
	antifreeze, and in the manufacturing of plastics and resins. It is a colourless,
	odourless, viscous liquid that is sweet tasting but highly poisonous if ingested,
	inhaled, or absorbed through the skin.
EG	Ethylene Glycol is a colorless, odorless, and sweet-tasting liquid that is widely
	used as an antifreeze, coolant, and in the production of plastics and synthetic
	fibers. It is a common component in engine coolants and de-icing solutions,
	but it is also toxic when ingested.
NCE	New chemical entity is any drug molecule which is not included in the
	Appropved Drug List in any form or any drug molecule which has not been
	imported to the Maldives.
	1

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1 INTRODUCTION

Maldives Food and Drugs Authority (MFDA) is responsible to regulate the medical products in the country under the Health Services Act (29/2015). The Medicine and Therapeutic Goods (MTG) Division of MFDA is mandated to implement a regulatory framework to ensure the accessibility of safe, quality assured and effective medical products for the people of Maldives.

The Health Services Act (29/2015) provides the legal provisions for establishing a national health system and standards for healthcare delivery services. The Health Services Act, Clause 65 (3) states that all medical products that are manufactured, imported, and sold in the country shall be registered by the Maldives Food and Drugs Authority. The Medicine Regulation further explains the procedures implemented for the registration of medicines including pharmaceutical and biological drug products in the country.

This "Guidelines for Medicine Registration including Emergency Use Authorization" will serve as the reference guide for the registration process including the pre-registration and post-marketing quality controls. However, this guideline shall be read in conjunction with the currently applicable laws and regulations together with other relevant legislation applicable to pharmaceutical and biological drug products in the Maldives.

2 OBJECTIVE

This guideline is aimed at supporting applicants for the registration/market authorization of pharmaceutical and biological drug products intended for human use in the Maldives.

3 SCOPE

The scope of this document encompasses the administrative requirements and procedures for submission, evaluation, and approval of registration applications of New Drug Products and Generic Drug Product (pharmaceutical and biological including vaccines), post-registration variations and renewal of registered drugs. It also covers the procedure for Emergency Use Authorization (EUA) applicable under public health emergency situations.

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4 Legal Context

- 4.1 These guidelines shall be read in conjunction with the other applicable legislations on drug product (pharmaceutical and biological products) which include but not limited to:
 - a. Medicine Regulation R-46 (2014)
 - b. Medicine Regulation Amendment R-49 (2016)
 - c. Health Service Act (29/2015)

5 RESPONSIBILITIES AND ACCOUNTABILITY

Pharmaceutical officers of Medicine	Responsible for verifying the documents,
Registration.	accepting the dossiers, evaluating the dossiers,
	submitting the summary of the evaluation to
	the National Board for Pharmaceuticals,
	preparing and issuing product registration
	certificates.
	Is also responsible for effective communication
	with the applicant in a timely manner
Senior pharmacist (Regulation Section)	Responsible for checking and verifying the
	product evaluation documents and to guide the
	pharmaceutical officers on evaluating the
	product.
Pharmaceutical Specialist (Medicine	To approve the medicines, approval from the
Therapeutic Goods Division)	national Pharmaceutical Board
Director General (MFDA)	Final authorization of all the activities related to
	MFDA tasks
National Pharmaceutical Board	For Technical decision for approval of the
	medicine

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6 Medicine registration

- **6.1** All Applications for registration in Maldives are accepted if only the product is categorized as a medicine in the country of origin.
- **6.2** Pathways for Registration / Marketing Authorization.
- **6.3** Registration pathways: Table 1 below shows the registration pathway for medicines and vaccines with the scope of reliance and the criterial for reliance mentioned in the table. Reliance mechanism is applied to make use of the maximum benefits for the available resources.

6.4 Reliance Approach for Registration / Marketing Authorization

- 6.4.1 MFDA recognized the reliance approach in regulatory decision making in line with the WHO guidelines for Good Reliance Practices and as per Guidelines on Good Reliance Practices for Regulation of Medicines. It gives consideration and significant weight to the assessments performed by Reference Regulatory Authorities and WHO Prequalification team. The decisions and other related authoritative information from National Regulatory Authorities and other trusted institutions are also considered while reaching the regulatory decision for enhanced access to safe, efficacious and quality assured products.
- **6.4.2** MFDA applies reliance principles for the products registration that has already undergone full evaluation / assessment by the Reference Regulatory Authorities either in the country of origin or where the product is being exported based on the full dossier assessment performed by these reference regulatory authorities. In such cases, applications will be accepted with abridged data / limited dossier as defined in this guideline.
- **6.4.3** PICs certified manufacturing sites are also included in this reliance process.
- 6.4.4 Reliance pathways are mandatory for all vaccines, biological products, new chemical entities, generic-generics and new fixed dose combinations.
- 6.5 Reference Regulatory Authorities (RRAs) and organizations.
- **6.5.1** MFDA classifies all applications for registration of medicines products into the below mentioned categories as mentioned in the below table 1, based on the product origin and its approval status from other NRAs or other trusted organizations (e.g. WHO).

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- **6.5.2** For this purpose, reference countries are identified based on criteria for stringent regulatory authorities, WHO listed regulatory authorities and those regulatory authorities which have achieved maturity level 3 and above. The documentation requirement is detailed for each reliance pathway in the table 1 below with the assessment duration and validity period.
- 6.5.3 Only the mentioned required documents shall be submitted under each registration pathway.

 If any additional information is required MFDA shall inform the applicant to submit accordingly.
- **6.5.4** Refer to Annex-II for the reference countries for Reliance.

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	Table 1: Medicine Registration Pathways with requirements							
Ge	Registration	Pathways	plications	All Applications shall have this following information: Letter of Appointment as in Annex I of this document Letter from the manufacture to MFDA. (for all new registrations) Artwork, 360 angle picture of the product with package insert. Completed checklist as per registration pathway Requirements Assessment duration from validity the date of receiving the				
					submission fee			
		on approval by reference NRAs	in a Reference NRA	 Evidence of approval in the country of manufacture. Evidence of approval in one of the reference countries as mentioned in Annex II of this document. (For the above 2 points, a link should be provided to trace the evidence or registration or marketing authorization certificate.) Product information for reliance as in Annex III Stability study report covering applicable climatic zone as mentioned in section F 2.1, 	days	5 years with no variations except changes in artwork		
		1.2 GMP	1.2.1	F12, (2.1) and F3 • Evidence of PIC/S-GMP	Calendar 40	5 years and no		

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verification	Manufacturing	compliance for the site where	days	variations
	site certified	the finished dosage form is		except changes
	by PIC/S	manufactured and batch		in artwork
	member NRA	release takes place1.		
		Evidence of approval by the		
		NRA of the country where the		
		finished dosage form is		
		manufactured, and batch		
		release takes place. For this, a		
		link should be provided to		
		trace the evidence or		
		registration or marketing		
		authorization certificate.		
		Verifiable declaration of		
		approval by at least 3 other		
		NRAs. For this, a link should be		
		provided to trace the evidence		
		or registration or marketing		
		authorization certificate(NRA		
		website shall verify that the		
		product is registered there		
		even though CoPP is provided)		
		Product information for		
		reliance as in Annex III		
		Stability study report covering		
		applicable climatic zone as		

¹ PICs certified manufacturing sites from the EU and USA can be verified using the following links. Alternatively, GMP certificates from PIC/S authorities can also be provided:

- 1. http://eudragmdp.ema.europa.eu/inspections/displayWelcome.do;jsessionid=VSj9l6duvYdglizmjw5ojlLTiRKH5-EwlbnUqabF2Ks1lh8NuukA!-1996855337
- 2. https://datadashboard.fda.gov/ora/index.htm

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			mentioned in section F 2.1,			
			F12, (2.1) and F3			
1.	.2.2	•	Evidence that a WHO	Calendar	45	5 years and no
M	/lanufacturing		prequalified medicine or	days		variations
sit	ite of a WHO		vaccine is manufactured and			except changes
pr	requalified		the batch released on the			in artwork
pr	roducts		same site			
			https://extranet.who.int/preq			
			ual/medicines/prequalified/fin			
			ished-pharmaceutical-products			
		•	Evidence of approval by the			
			NRA of the country where the			
			finished dosage form is			
			manufactured, and batch			
			release takes place. For this, a			
			link should be provided to			
			trace the evidence or			
			registration or marketing			
			authorization certificate.			
			Verifiable declaration of			
			approval by at least 3 other			
			NRAs. For this, a link should be			
			provided to trace the evidence			
			or registration or marketing			
			authorization certificate. (NRA			
			website shall verify that the			
			product is registered there			
			even though CoPP is provided)			
		•	Product information for			
			reliance as in Annex III			
		•	Stability study report covering			

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		applicable climatic zone as		
		mentioned in section F 2.1,		
		F12, (2.1) and F3		
1.3	1.3.1CRP-PQ	Letter of manufacturer	Working 90	5 years and no
Collaborativ		showing interest in going for	days	variations
е		this pathway. Once letter is		except changes
registration		received from the		in artwork
procedure		manufacturer the information		
(CRP)		of the dossier will be retrieved		.
		from the relevant		
		authorities/organization by		
		MFDA to process the		
		registration of the product.		
		Shall follow the requirements		
		same as in 1.1.1 of this table		
	1.3.2 CRP-SRA	Letter of manufacturer	Working 90	5 years and no
		showing interest in going for	days	variations
		this pathway. Once letter is		except changes
		received from the		in artwork
		manufacturer the information		
		of the dossier will be retrieved		
		from the relevant		
		authorities/organization by		
		MFDA to process the		
		registration of the product		
		Shall follow the requirements		
		same as in 1.1.1 of this table		

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1.Full	1.1 Full dossier	•	• Product full dossier with the	Working	150	5 years +	all
procedure			document requirement in	days +	any	variations	
			section 11.2	clock	stop		
			Declaration of regulatory	required	for		
			status in other countries.	external			
		•	Full ICH M2 and M3 dossier	expertise			
			parts.				
		•	Product information as per the				
			information mentioned in				
			section 12.3 B and				
			Manufacturer information as				
			per section C2 and C3				
		•	Samples				
1.Notifica	These are for low-risk	•	Evidence of approval by the	Working	120	5 years +	all
tion	medicines like vitamins and		NRA of the country where the	days		variations	
	vitamin preparation that are		finished dosage form is				
	categorized as medicines.		manufactured, and batch				
	Those that falls under		release takes place. For this, a				
	nutraceuticals shall follow the		link should be provided to				
	registration procedure for		trace the evidence or				
	nutraceuticals.		registration or marketing				
			authorization certificate.				
		•	Verifiable declaration of				
			regulatory status in at least 3				
			other NRAs. For this, a link				
			should be provided to trace				
			the evidence or registration or				
			marketing authorization				
			certificate.				
		•	Stability study report covering				
			applicable climatic zone as				

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	mentioned in section F 2.1,
	F12, (2.1) and F3
Products	Copy of appeal, procurement,
for	or request issued by the
exclusive	concerned programme.
use in	Declaration of regulatory
public	status in other countries;
health or	Proof of approval and
disease	marketing in one or more
control	reference country.
program	Detailed description of the
mes	product approved in the
	mentioned reference country
	OR evidence of PIC/S-GMP
	compliance for the site where
	the finished dosage form is
	manufactured and batch
	release takes place AND
	evidence of approval by the
	NRA of the country where the
	finished dosage form is
	manufactured and batch
	release takes place;
	Stability study report covering
	applicable climatic zone as
	mentioned in section F 2.1,
	F12, (2.1) and F3
Donations	Copy of appeal or request for
	donation issued by a national
	institution.
	Copy of marketing

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6.6 Import requirements

- **6.6.1** All applications once registered and approved shall fulfill the import requirements as assigned by the guidelines of MFDA.
- **6.6.2** This includes and is not limited to:
 - a. Tested reports on the DEG/EG for all cold and cough preparation and other preparations as indicated by MFDA.
 - b. NDMA reports for Ranitidine and its formulations
 - c. Batch certificates for all IV fluids and biological preparations
 - d. Export license for all Controlled medicines

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7 Types of Application

7.1 New Drug Application / New Chemical Entity Application

- **7.1.1** The registration of a new active pharmaceutical ingredient that has not been previously approved by the authority and present in the most recent Approved Drug list (ADL), either as a single ingredient drug or as part of a combination product.
- **7.1.2** These applications will be subject to a high level of scrutiny in terms of efficacy, safety and their contribution to therapeutic improvement.
- **7.1.3** Only reliance pathways are accepted for these applications

7.2 Generic Drug Application (Generic –Generic and branded -Generic)

- **7.2.1** The registration of a medicine that has the same active ingredient as the innovator or patented medicine, including dosage and having the same safety, efficacy, stability and quality requirements.
- **7.2.2** Majority of the applications received falls under this category.
- **7.2.3** Only Reliance pathways are accepted for Generic-generic application.
- **7.2.4** Reliance pathways are encouraged for Branded generics
- 7.3 Registration of products that has already been registered by another party.
- **7.3.1** These are the products that has already undergone full evaluation by assessing the documents submitted as per the criteria defined, by another party.
- **7.3.2** Application for this shall be limited as already the product has been fully evaluated.
- **7.3.3** The Applicant shall ensure that exact same product is submitted and also shall submit the Product information for reliance as in Annex III by completing the information in points 2 to 13, 16 to 19, 21, 24 to 25 and 27.

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- **7.3.4** The applicant shall ensure the safety, quality and efficacy of the product and shall submit batches of tested reports as per the official monogram published from a WHO pre-qualified laboratory for each batch imported. The applicant shall provide evidence documents that the tests done are from WHO pre-qualified laboratories
- 7.3.5 Refer to the list of WHO pre-qualified laboratories from: https://extranet.who.int/prequal/medicines/quality-control-laboratories,

7.4 Re-registration/renewal Application

- **7.4.1** This is the application for a product that has been registered previously under the criteria set by the authority for medicine registration.
- **7.4.2** All these registered products will be indicated with the letter "R", in the Approved Drug List. The validity period for the registered product under the previous criteria is 5 years.
- **7.4.3** Before the expiry of the validity the client shall submit the application for re registration as per the current criteria.
- **7.4.4** Any product registered and approved under the previous criteria/s shall be considered as **new** registrations and shall follow the new criteria.
- 7.4.5 Re-registration shall also be considered based on the import of the product during the past 5 years.
- **7.4.6** Application for renewal of registered product shall be submitted at least 30 days before the expiry date of the current MA validity along with the processing fee.
- **7.4.7** The general procedure for the renewal of the reregistration is the same as the initial registration.
- **7.4.8** If the registration of the product is expired and the applicant did not apply for renewal, then product will be removed from ADL within 1 month after the expiry. If the applicant wants to register it again the application will be treated as a new application as per the criteria mentioned in this guideline.
- **7.4.9** Requirements for re-registration: Table 2 (below)

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Table 2: Re-registration Requirements

Registration			Re-registration Requirements	Assessment duration from the date of receiving the submission fee	Registration validity
A. Reliance	1.1Reliance on approval by reference NRAs	1.1.1 Approved in a Reference NRA	 Updated Product information for reliance as in Annex III Post approval changes or variations if any. Evidence of Reference country approval in consistence with the initial registration requirement. Any change in the reference NRA registration shall be provided with justification. Samples Compiled Import data for the past 5 years 	Calendar 20 days	5 years and no variations except changes in artwork
	1.2 GMP verification	1.2.1 Manufacturing site certified by PIC/S member NRA	 Evidence of PIC/S-GMP compliance for the site where the finished dosage form is manufactured Updated Product information for reliance as in Annex III Post approval changes or variations in any. Evidence of approval by the NRA of the country where the 	Calendar 20 days	5 years and no variations except changes in artwork

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			finished dosage form is		
			manufactured		
			Verifiable approvals by at least		
		•			
			3 other NRAs. Which shall be		
			inconsistence with the initial		
			registration. Any change in the		
			reference NRA registrations		
			shall be provided with		
			justification.		
		•	NRA website shall verify that		
			the product is registered there		
			even though CoPP is provided.		
		•	Samples		
		•	Compiled Import data for the		
			past 5 years.		
	1.2.2	•	•Updated Product information	Calendar 20 days	5 years and no
	Manufacturing		for reliance as in Annex III		variations
	site of a WHO	•	Post approval changes or		except changes
	prequalified		variations in any.		in artwork
	products	•	Evidence of approval by the		
			NRA of the country where the		
			finished dosage form is		
			manufactured		
		•	Verifiable approvals by at least		
			3 other NRAs, which shall be		
			inconsistence with the initial		
			registration. Any change in the		
			reference NRA registrations		
			shall be provided with		
			justification.		
		•	Samples		

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			Compiled Import data for the
			past 5 years.
	1.3	1.3.1CRP-PQ	Updated Product information
	Collaborative	1.3.1CRF-FQ	for reliance as in Annex III
	registration		Post approval changes or
	procedure		variations in any.
	(CRP)		• Samples
	(CIVI)		Compiled Import data for the
			past 5 years.
		1.3.2 CRP-SRA	Updated Product information
		1.3.2 CNF-3NA	for reliance as in Annex III
			Post approval changes or
			variations in any.
			Samples
			Compiled Import data for the
			past 5 years
B. Full	1.Full	1.1 Full dossier	Product full dossier with the Working 90 days 5 years + all
Dossier	procedure		document requirement in + any clock stop variations
			section 11.2 required for
			Declaration of regulatory external
			status in other countries. expertise
			Full ICH M2 and M3 dossier
			parts.
			Updated Product information
			as per the information
)	mentioned in section 12.3 B
			and Manufacturer information
			as per section C2 and C3
			• Samples
			Compiled Import data for the
			past 5 years

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1.Notification	These are for	• Evidence of approval by the Working 60 days 5 years + al
	low-risk	NRA of the country where the variations
	medicines like	finished dosage form is
	vitamins and	manufactured and batch
	vitamin	release takes place. For this, a
	preparation that	link should be provided to
	are categorized	trace the evidence or
	as medicines.	registration or marketing
		authorization certificate.
		Verifiable declaration of
		regulatory status in at least 3
		other NRAs. For this, a link
		should be provided to trace the
		evidence or registration or
		marketing authorization
		certificate.
		Stability study report covering
		applicable climatic zone as
		mentioned in section F 2.1,
		F12, (2.1) and F3
		• Samples
		Compiled Import data for the
		past 5 years

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7.5 Variation Application

- **7.5.1** A variation is a change in the dossier of a product that has already been registered and granted Market Authorization (MA) under the criteria set by the authority for medicine registration.
- **7.5.2** These changes can include a change in label, shelf-life, excipients and stability data.
- **7.5.3** Any variation to a product that is listed as a primordial product (PP) or Preauthorization required product (PA) in ADL or a product which has not undergone a registration process will not be considered as a variation application.
- **7.5.4** Variation to a registered medicine can be considered as a new application based on the formulation change, change in the strength of the active ingredient and excipients or any other change as the authority may decide based on the product.
- **7.5.5** For details on variations and requirements refer to clause 15 of this document.

8 Conditional Marketing Approval.

- 8.1 Conditional marketing approval is a fast-track procedure for availability of new medicine and vaccines with a positive benefit-risk balance and has the potential to address unmet medical needs in the country. This program intends to provide a time limited approval to new drugs and vaccines for serious or life-threatening diseases that have no alternative treatments, based on the conditional or special approval in the reference authorities with limited clinical data or less comprehensive clinical data.
- **8.2** The conditional marketing approval can be converted to full registration based on the submission and review of full data and full registration status in reference countries.
- **8.3** The National Pharmaceutical Board (NPB) considers application under conditional marketing approval pathway with the intent to provides patients and healthcare professional with faster access to new drug products.

8.4 Application submission and Review Process.

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- **8.4.1** Applicants must justify applicability of conditional marketing approval pathway in the cover letter of registration application, providing brief justification for fulfilling the eligibility criteria for consideration under this non-routine pathway along with the approval status in other regulatory authorities.
- **8.4.2** The MFDA evaluators will review the application, validate and assesses the application and gives a positive or negative opinion on the application within 120 days, taking into account the urgency and the public health need. Application may be rejected if the benefit-risk ratio is negative.
- **8.4.3** The applicant must fulfill the specific obligations and conditions attached to the authorization for submission of remaining data on completion of studies.
- **8.4.4** The MFDA will review the conditional authorization at least once a year, based on the data submitted by the applicant, and decide whether to renew, vary, suspend, or revoke it.

9 Emergency Use Authorization (EUA)

- **9.1** Emergency Use Authorization (EUA) is grant of conditional registration for a medical product on priority basis during declared health emergency situations.
- 9.2 As the routine registration regulatory process require complete clinical trials as per international harmonized requirements (like ICH guidelines) which cannot be followed in emergency health situation, thus EUA enables expedited authorization of an unapproved medical product and riskbenefit analysis depicts that its use will be helpful in reducing fatalities / mortality due to instant health emergency.
- **9.3** The procedure for granting EUA has been adopted by Reference Regulatory Authorities and also recommended by the World Health Organization.
- 9.4 To obtain EUA, a manufacturer / importer shall submit a request to MFDA with evidence from clinical trials or other adequate and well-controlled clinical investigations or approval of any Reference Regulatory Authority that suggests that the product can be effective in preventing, diagnosing, or treating the serious or life-threatening disease or condition.

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- **9.5** After reviewing such evidence and being satisfied, MFDA may authorize the conditional use of the product under EUA once it is proven that the known and potential benefits outweigh the known and potential risks.
- **9.6** EUA is a temporary authorization and EUA holder shall continue to collect evidence of the safety, efficacy, and quality of the product along with updated status by Reference Regulatory Authorities and submit such data to MFDA for review and appropriate decision.

9.7 Criteria for consideration of Emergency Use Authorization application

- **9.7.1** Criteria for consideration of Emergency Use Authorization is as follows:
- 9.7.1.1 A declaration of a public health emergency by the relevant authorities of Maldives such as pandemics or natural disasters or rare diseases with no approved treatment options or life-threatening diseases with limited treatment options.
- 9.7.1.2 Medical products already registered by MFDA either cannot be used in prevailing health emergencies in the country or very limited alternative medical products are available for diagnosis, prevention, or treatment of disease.
- 9.7.1.3 If any Reference Regulatory Authority has registered any medicine for treatment and use in prevailing health emergency which has not been yet registered by MFDA, then MFDA will process case for priority consideration of registration of such medicine and will convene special meetings of National Pharmaceutical Board for priority decision.
- 9.7.1.4 If any Reference Regulatory Authority has not yet registered any medical products for use in prevailing health emergency but granted Emergency Use Authorization (at least completed or on-going Phase III that clearly demonstrates the safety and efficacy of the product), then these guidelines will enable MFDA to authorize the use of unapproved medical either through verification or Abridged pathway (as the case may be) by following Guidelines for Good Reliance Practices for Regulation of Medicines. Moreover, the applicant has provided sufficient data of demonstrated appropriate efficacy and safety in preliminary trials and risk-benefit analysis allows use of medicinal product with certain conditions.
- 9.7.1.5 Applicant of EUA (Manufacturer/importer) has adequate plan for monitoring the safety and efficacy of the product i.e. Risk Assessment Plan and Risk Management Plan.

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9.8 Procedure for Processing of EUA Applications.

- **9.8.1** The initiation of the EUA by MFDA is based on the notification of emergency declaration by relevant authorities of Maldives.
- **9.8.2** Once the relevant authority notifies the nature of health emergency, MFDA will nominate a focal person for coordinating EUA procedures.
- **9.8.3** Establishment of a dedicated hotline and email address for manufacturers/importers to facilitate EUA submissions.
- 9.8.4 Focal point will be coordinating all activities regarding EUA application including coordination with stakeholders, Ministry of Public Health, and other government agencies to implement EUA procedures, submission of application by the company, prioritizing product dossier by MFDA, application assessment by Medicine Therapeutic Goods Division, convening of National Pharmaceutical Board and priority decision.
- 9.8.5 Medicine Therapeutic Goods Division, MFDA shall evaluate all EUA applications and will convey shortcomings (if any) to applicant on priority. After reply of the applicant, Medicine Therapeutic Goods Division will prepare agenda of National Pharmaceutical Board meeting and will mention all details of application, assessment report and any other relevant information in agenda for the consideration of National Pharmaceutical Board.
- **9.8.6** The National Pharmaceutical Board may co-opt relevant experts (if needed) like experts in pharmaceutical and vaccine regulation and manufacturing, medical experts, epidemiologists, clinical and pharmacovigilance experts etc.
- 9.8.7 National Pharmaceutical Board may exempt any requirement in registration application which in its opinion cannot be fulfilled in such health emergency situations and it has no or very limited impact or can be confirmed by MFDA through some other means like online data or communication with another NRA via email etc.
- **9.8.8** An EUA will only be granted if MFDA, after complete review, finds that the product meets all the applicable criteria of safety, efficacy, and quality. In the event of approval, letter will be issued immediately.

9.9 Process Timeline

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9.9.1 Applications for pharmaceutical products during health emergency fulfilling conditions get priority review in 07 working days after complete application received.

9.10 EUA Validity

9.10.1 Emergency Use Authorization will be reviewed by the National Pharmaceutical Board every three months based on available safety, efficacy and quality data. Thereafter, the applicant will be required to submit the applicable data as per the routine registration pathway for conversion to full registration status and renewal accordingly.

9.11 Data Requirement for EUA

- **9.11.1** The minimum requirement for registration of drug products, biological and vaccines will be as specified in the Guidelines for regulation requirement for product registration and approval of vaccine in emergency (Doc Number: MTG/RE-LA/STD-TE 003) or as determined by National Pharmaceutical Board on case-to-case basis.
- **9.11.2** All Applications for registration in Maldives are accepted if only the product is categorized as a medicine or vaccine in the country of origin.

10 Pre-Application Process for product registration.

10.1 Applicants for Product Registration.

- **10.1.1** For registration of a product in Maldives, the manufacturing company shall have a local representative, or a locally incorporated company authorized by the manufacturer or Marketing Authorization Holder in the country of origin, who will be responsible for all the communications to the Authority. The local representative or the applicant shall have a valid medicine import license as per the criteria defined by the Authority.
- **10.1.2** The local representative and the applicant can be the same, but local representative or applicant shall be a Maldivian National.

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- **10.1.3** It is the responsibility of the assigned local representative to furnish all the information required for product registration and all supporting documents as defined by the Authority and ensure that these documents are legitimate and valid.
- **10.1.4** Applicant/local representative shall verify that all the required documents are submitted by using the checklist provided in Annex-V or Annex VI to facilitate the acceptance of the documents for registration.
- **10.1.5** This Application Checklists mentioned in 10.1.4, shall be used to ensure the submission of a complete application.
- **10.1.6** Please note that not all documents mentioned in the checklist are mandatory and the required documents is dependent on the type pf application and the Pathway the product is applying under.
- 10.1.7 All documents required shall be submitted in coloured softcopy via Dhirithi portal. However, MFDA reserves the rights to request for the original or certified true copy of submitted documents if there is any doubt that a submitted scanned document is not an accurate reflection of the original document.
- **10.1.8** The initial acceptance of the application after screening does not ensure that all information provided are within the acceptance criteria. MFDA has the right to requests for additional documents or changes to the information/documents during evaluation.
- **10.1.9** This check list shall be filled and uploaded in excel format to Dhirithi Portal with relevant documents based on the registration pathway. (This form can be retrieved in Dhirithi Portal under publication).
- **10.1.10** If any section within the criteria of the form left unfilled, or/ and a mandatory document and the required information is not submitted, MFDA will reject the application.
- 10.1.11 The Authority may request additional information not described in this document that is deemed necessary to ensure the safety, efficacy and quality of the product. This will be informed by the Authority as a written request to the applicant or the local representative.

10.2 Responsibilities of Applicants.

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- **10.2.1** The manufacturer in the country of origin, shall designate a local representative by issuing an authorization letter to MFDA, indicating that all responsibilities in communicating on behalf of the manufacturer shall be done by the local representative to the Authority which includes supplying all the relevant information for product registration.
- **10.2.2** The designated or assigned local representative can then take responsibility as the applicant in supplying the required information to the authority for the registration procedure.
- **10.2.3** The key responsibilities of applicants are as follows:
 - 1. The applicant shall be registered as an authorized medicine importer under the Authority as per the criteria of medicine regulation.
 - 2. The person or an authorized representative established in Maldives, shall take full responsibility of the medicine that they supply to the market, which includes, informing the authority of any variations in the product after issuance of MA, recalling the registered medicine if required, and providing PSUR's if needed. It is the responsibility of the applicant or MAH of the product to ensure that the medicine complies with the specification as approved by MFDA throughout the supply chain. The evidence of these shall be documented with the applicant and shared with the authority when needed.
 - 3. The applicant shall have an established system for reporting and handling adverse drug reactions and for these focal points shall be identified and documented. The focal point shall closely liaise with the authority and shall provide the needed information to the authority. This system shall include market safety information of the drug product as well.
 - 4. The Applicant or the MAH shall have a list of suppliers and manufacturers through which the medicines are imported. It is the responsibility of the Applicant to validate and verify the authenticity of the suppliers and notify MFDA when required.
 - 5. It's the applicant's responsibility to provide the required information for registration and re-registration with in the required time lines.
 - 6. The re-registration shall be submitted 30 days before the expiry of the product

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7. It's the applicant's responsibility to ensure that the exact same product from the exact same site as registered, is imported to Maldives

11 Pre-Approval Process

11.1 General Considerations

- **11.1.1** All documents submitted for registration purpose shall be in English language and shall be signed and endorsed.
- **11.1.2** All electronic documents submitted shall be signed and endorsed unless such documents can be verified by the regulatory authorities.
- **11.1.3** The documents required for the registration of a product differ according to the Registration pathways as described in Table 1 of this document.
- **11.1.4** The acceptance criteria of each document are indicated below in Clause 11.2 for full dossier. Any application with missing documents or documents that do not meet the criteria set will be rejected.
- **11.1.5** The reason for the rejection will be indicated in the Dhirithi portal for the applicant to see.
- **11.1.6** A separate application is required for each drug product i.e., products containing the same ingredients but made to different specifications in terms of strength, content of API/DS, dosage form, description and pack size etc.
- 11.1.7 For tablets and capsules if it undergoes the same production and packing process under the same quality assurance system, it will be considered as a one drug product regardless of its pack size. Example: A having 12 tablets per strip and 10 tablets per strip will be considered as one product if the product undergoes the same process and has the same labelling information on the product.
- **11.1.8** MFDA shall reject the applications which do not fulfill the criteria as per applicable guidelines.

11.2 Data Requirement and Acceptance Criteria for full Dossier Preparation.

11.2.1 The table below contains a list of documents and required data in a full dossier along with the information that shall be included in each document for it to be accepted by the Authority.

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- **11.2.2** As all applications do not require a full dossier, to determine the type of dossier application and the required documents, refer to Table 1 of this document to determine product application category and registration pathways.
- **11.2.3** The codes and document title stated under column-I and II respectively in the following table are consistent as they appear on the Dhirithi portal for ease of interpretation.

Code	Document Title	Acceptance Criteria				
		·				
В	Pharmaceutical	The pharmaceutical information shall be supplied by the				
	Information	manufacture with signed and endorsed including the following				
	Sheet	information:				
		a. API information: Shall include the API information as per one of the				
		following criteria:				
		i. Confirmation of API Prequalification document (CPQ) as				
		specified in Annex-IX				
		ii. Certificate of suitability of the European				
		Pharmacopoeia (CEP) as specified in Annex-X.				
		iii. Technical Information on the active pharmaceutical				
		ingredient/s as specified in Annex-XI				
		b. Brand name, Trade name or Product name: Shall provide this				
		information in the format as provided by the manufacturer				
		i. The product name shall be entered according to the				
		submitted product label and shall be same with the				
		product name mentioned in submitted Certificate of				
		Pharmaceutical product (CoPP).				
		ii. The strength of the active pharmaceutical ingredient /				
		Drug substance shall generally be included as part of the				
		product name to allow differentiation between different				
		products containing the same active pharmaceutical				
		ingredient / Drug substance.				
		c. International Nonproprietary Name (INN) or the Active				
		Pharmaceutical Ingredient (API) or Generic name: Shall provide the				
		. Harmaceatical ingredient (All I) of deficite fiame, shall provide the				

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		information with the details as mentioned below:
		i. The name and amount of active pharmaceutical ingredient(s) /
		Drug substance(s) present in the formulation and in the form of
		salts or chelates shall be clearly stated. Example: Each film
		coated tablet contains Calcium carbonate equivalent to
		elemental calcium 500mg.
		ii. If more than one active ingredient is present in the preparation,
		it shall be separated by a + between each active ingredient.
		Example.: Calcium carbonate 500mg + Docosahexaenoic Acid
		150mg+vitamin D3 200IU
		d. Non active increasions of European Chall provide these details of
		d. Non-active ingredient or Excipient: Shall provide these details as
		mentioned below:
		i. All non-active ingredients and all proprietary ingredients (e.g.,
		colorants, flavoring agents, etc.) used in the product shall be
		mentioned with the composition with the grade of the
		excipients.
		ii. In case of cough and cold preparations and paracetamol
		preparations, the exact grade of the excipients shall be
		mentioned with supporting documents.
		iii. For all the pediatric oral formulations including cough, cold, and
		paracetamol formulation the certificate of analysis (COA) shall
		be submitted for all the excipients used, specifically if glycerin
		or glycerol or propylene glycol or high risk excipients for DEG/EG
		is used, verifying that it does not contain the impurities
		diethylene glycol (DEG) and ethylene glycol (EG).Refer to Annex
		IV for the list of Excipients of high risk of contamination with DEG/EG
		iv. Manufacture validation protocols of the excipients shall be
		submitted specifically those that are at a risk for diethylene
		glycol (DEG) and ethylene glycol (EG) contamination. For such
		8.755. (= = 5, 2 3). 5 (= 5) 55 (= 5) 55 (= 5) 55

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Code	Document Title	Acceptance Criteria
		excipients each container of the excipients shall be tested for purity and validity and evidence documents shall be submitted. v. The test mentioned in point 4 of the clause shall be as per the
		official monogram for purity.
		vi. Registration holder will ensure that the manufacturer will
		perform impurity testing as identified by the manufacturer of
		innovator drug product like N-Nitroso dimethylamine (NDMA),
		N-Nitroso diethylamine (NDEA) in valsartan, metformin etc.
		e. Pharmacopeia standard / Formulation of the product: Shall provide
		the details as mentioned below
		i. All Active pharmaceutical ingredient(s) / drug substance(s) and
		all excipients in the product shall be listed with their
		Pharmacopeia standard i.e., British pharmacopeia (BP), US
		pharmacopeia (USP) or Indian pharmacopeia (IP) or any pharmacopeia of stringent regulatory authorities.
		ii. If there is no pharmacopeial formulation as mentioned in point
		1), method validation report of the in-house method shall be
		provided which has to be endorsed by a third party. The third
		party can be an accredited laboratory, regulatory authority or
		any other external or internal assessment body as nominated
		by the Authority.
		f. Pharmaceutical Dosage Form: Pharmaceutical dosage form is
		defined as the physical form of the drug product which is intended
		for administration to the patients:
		i. Dosage form shall be as specific as possible with respect to the
		product's actual dosage form. Example: Film-coated Tablet
		instead of Tablet, sustained release tablets instead of tablets.
		ii. In certain cases, the dosage form may also include information
		about the container closure system. Example: pre-filled syringe,
		spray pump and pressurized container.

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		g. Strength: Strength is defined as the amount of active pharmaceutical ingredient(s) / Drug substance(s) in the dosage form. Strength shall be provided for all APIs/DS including if the product is a combination drug. Each strength shall be separated with a "+" Example.: 500mg + 250mg
		h. Volume of the preparation: Applicable for liquid and semi solid dosage forms like oral liquids, injectables, creams and ointments etc and the volume shall be clearly mentioned as follows:
		i. Volume is not applicable for Tablet and capsules
		ii. For semi-solid dosage forms (i.e., ointments, pastes, cream,
		gels), liquid dosage forms (i.e., suspensions, syrup, liquid for
		injection), powders, suppositories and MDI's, volume shall
		be indicated as per product label.
		i. Product description, Container type and Pack sizes: shall
		provide this information:
		i. Description of primary packaging shall be defined with the
		pack size. Example blister pack of 12 tablets
		ii. Description of secondary packaging shall be defined with
		the pack size. Example 12 tablet blister pack of 10 blisters
		equal to total 120 tablets in 1 box
		iii. Length, width, height of primary and secondary packaging
		shall be provided in detail.
		iv. Weight of drug product which is submitted for registration
		and shall also indicate the deviation level for the weight.
		v. Odor description of the product submitted for registration
		shall be submitted.

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		 vi. A visual description of the product shall be submitted including the shape, size, color any engraving or any other detail of the product. vii. Preparations whose primary packing is plastic or if the preparation comes in direct contact with the plastic 		
		container shall provide studies on such containers to demonstrate the safety of the material used to the preparation.		
		j. Route(s) of Administration:		
		All routes of administration proposed for the product shall be		
		included and specified accordingly.		
		k. Indication or Use of the product:		
		The intended use or the indication of the product shall be clear		
		specified. Example: Used for upper respiratory infections		
		I. Therapeutic Class according to WHO ATC Index shall be		
		indicated with:		
		i. The WHO ATC code		
		ii. WHO ATC classification		
		iii. Shall be provided for each therapeutic indication proposed for a product.		
		m. Storage conditions of the product shall be provided:		
		i. The condition in which the drug product shall be stored and		
		kept shall be clearly specified. Example temperature,		
		humidity etc. Of the product storage shall be specified		
		ii. Non-numeral statements such as "Store in a cool dry place"is not encouraged.		
		n. Shelf life of the product: The shelf life is the period between the		
		execution of the preparation and its expiry date. Product Shelf		
		life shall be specified in months.		

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		 Dispensing Category in country of origin shall be specified. If the product is a prescription only medicine (POM), over the counter (OTC), Hospital use etc shall be defined as per the registration of the product in the country of origin 		
C2	Manufacturer responsible for lot release of the Finished Product	 i. Full address of the manufacturer(s) shall be provided with site and country of origin including the phone number, fax, e-mail. ii. Contact details of the manufacturer(s) shall be provided that can be reachable if needed. iii. This contact shall provide further information and verification if needed by the authority and shall be responsive to the queries sent by the authority. 		
СЗ	Manufacturer responsible for packaging / final batch release of the Finished Product, if different.	 i. Full address of the manufacturer shall be provided with site and country of origin including the phone number, fax, e-mail. ii. Contact details of the manufacturer shall be provided that can be reachable if needed. iii. This contact shall provide further information and verification if needed by the authority and shall be responsive to the queries sent by the authority. 		
C4	Manufacturing License	A manufacturing license is a permit issued by the regulatory authority of the country of origin to manufacture drug product. i. The manufacturing license shall be at least 6 months valid at the time of submission. ii. The manufacturing license shall contain date of issue, expiry, identity of issuing authority, the activities or the products covered under the license and full manufacturing site address. iii. The manufacturing license shall be self-attested and notarized copy.		
C61	Valid GMP certificate	Good Manufacturing Practices (GMP, also referred to as 'cGMP' or		

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		'current Good Manufacturing Practice') is the aspect of quality		
		assurance that ensures that medicinal products are consistently		
		produced and controlled to the quality standards appropriate to		
		their intended use and as required by the product specification.		
		GMP defines quality measures for both production and quality		
		control and defines general measures to ensure that processes		
		necessary for production and testing are clearly defined, validated,		
		reviewed, and documented, and that the personnel, premises and		
		materials are suitable for the manufacturing of pharmaceuticals		
		and biologicals including vaccines. GMP also has legal components,		
		covering responsibilities for distribution, contract manufacturing		
		and testing, and responses to product defects and complaints.		
		Following documents related to GMP need to be provided.		
		i. Proof of GMP compliance (valid GMP certificate) shall be		
		submitted for all the sites involved in any step of		
		manufacturing of the product		
		ii. A color scanned copy of the original or certified true copy of		
		GMP certification document issued by the relevant drug		
		regulatory agency shall be submitted, certifying that the		
		manufacturer concerned complies with current applicable		
		GMP standard.		
		iii. GMP Certificate shall have the following information; date of		
		issue, identity of issuing authority or agency approving GMP		
		certificate, validity of the GMP, manufacturing site address		
		and dosage forms of productions.		
		iv. GMP Certificate shall have the validity of 6 months at the time		
		of submission.		
		v. If the GMP does not have the 6-month validity, the applicant		
		shall submit proof document requesting to renew the GMP		

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Code	Document Title	from the country of origin and hence these applications shall be put on hold till the new GMP is submitted. Once the new GMP certificate is submitted, registration of the product will be issued. vi. If the validity period or expiry date is not stated on the GMP Certificate, the applicant shall supply supporting documents to confirm the validity period of the GMP certificate. vii. Commitment letters of GMP renewal are not accepted. viii. The names and addresses of manufacturer(s)/repacked(s)/batch releaser(s) shall be consistent with the information provided in the GMP certificate
		 ix. The specific dosage form applied for registration shall be mentioned in the GMP x. The applicant shall submit with valid GMP certificate, the most recent GMP inspection report or a summary of the inspection report endorsed by the inspection authority.
С7	Proof of Validation of the Manufacturing method	As per ICH recommendations, copies of the validation process of Manufacturing method shall be provided including: i. Short description of the process with a summary of the critical processing steps or critical process parameters to be monitored during validation. ii. API /DS validation report iii. Excipients validation report iv. Finished product specification report v. Finished product specification report, specifically tested for Diethylene Glycol and Ethylene Glycol impurities in oral cough and cold preparations vi. Details of analytical methods

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		vii. In-process controls proposed with acceptance criteria
		viii. Additional testing intended to be carried out (e.g., with
		proposed acceptance criteria and analytical validation as
		appropriate)
		ix. Sampling plan - where, when and how the samples are taken
		x. Details of methods for recording and evaluation of results
		xi. Proposed timeframe.
		xii. Any variation from the validation protocol shall be documented
		with appropriate justification
		xiii. Following completion of the validation, a report containing the
		following information signed by the authorized person shall be
		provided:
Co	Canadaud Rakek Sina	 Batch analytical data Certificates of analysis Batch production records Report on any unusual findings, modifications or changes found necessary with appropriate rationale Report of the validation studies shall be submitted with a conclusive statement of the results, comments on any deviations observed, including recommending changes to correct deficiencies. xiv. Refer to ICH Quality Guidelines Q7 Good Manufacturing Practice Guide for Active Pharmaceutical Ingredients Section 12 for further reference.
C8	Standard Batch size Quantity	 Shall submit the information specifying Label claim, Batch size, Quantity of all active ingredients and excipients per batch and per dosage form at relevant stages of manufacture, Overages
		and other adjustments with justification
		ii. For multiple batch sizes, the batch formula for each batch size is
		to be provided
С9	Technical Specification and source of all	Technical specification of all excipients and API(s) shall be provided

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	material(s)	indicating the pharmacopeial specification followed:		
		i. Specification provided shall be consistent with label claim		
		ii. Certificate of analysis (CoA) of all ingredients (APIs/DS and		
		excipients) shall be provided along with specifications		
		iii. The quality of the ingredients used in the production of the drug		
		substance (or drug product) shall meet standards appropriate		
		for their intended use.		
		iv. The quality of the excipients used in the drug product		
		formulation (and in some cases, in the drug substance), as well		
		as the container/closure systems, shall meet pharmacopeia		
		standards, where applicable and suitable acceptance criteria		
		shall be established for the non-pharmacopeial excipients.		
		v. Information on measures taken to ensure the quality and		
		control of these materials shall be provided.		
		vi. Source(s) of all excipients and API/DS shall be listed and the		
		origin or source of the API/DS and excipients shall be approved		
		by the manufacturer and this document shall be provided.		
		vii. Shall submit documents stating that all excipients used are of		
		pharmaceutical grade or grade approved for manufacturing the		
		pharmaceutical product.		
		viii. A signed `statement shall be provided by the manufacturer		
		indicating that all excipients and API(s)/DS are obtained through		
		approved vendor(s) in the country of origin.		
		ix. The manufacturer shall submit document evidence on how		
		vendor assessment is done for API/DS and excipients.		
C10	Brief profile of	i. A brief description of the manufacturer, when it was established		
	Manufacturer(s)	and the products approved / manufactured shall be submitted.		
		ii. Shall Provide a short description of accreditations, achievements		
		and standards practices of the manufacturer.		

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		iii.	Shall Provide list of products manufactured and specify those currently marketed in the country of origin.
		iv.	Shall provide a list of products manufactured and exported to
			other countries, specifying which products are exported to which
			country.
	-	_	
CM1	Company profile	i.	Shall include a detailed profile of the company including but not
			limited to history, accreditations, and standards practices and
			international/national levels of recognition achieved, company
			information, staff, organizational chart, equipment used, quality
			control procedures used, etc for all new companies/applying first
			time to MFDA.
		ii.	A detailed Company profile is not required for companies that
			already have a drug product(s) registered in Maldives.
		iii.	For all new sites involved in any step of manufacturing also
			require company profile documentation.
		iv.	Company profile is also required in instances of a major change
			brought to a company that has previously registered a product in Maldives
		v.	Shall also state whether the company is manufacturing under
			loan license or not. If so, shall include all details of loan
			manufacturer including regulatory details of approving NRA.
C11.0	Manufacturing plant		
	layout and machinery	i.	Shall include a list of equipment which is relevant to the product
	involved		under application along with details of water treatment, HVAC
			and waste disposal systems.
		ii.	Manufacturing plant lay-out shall be clear, legible and relevant
			to the product under application
C11.1	Manufacturing and	i.	Process flow chart of the whole process of the manufacturing of
			the product shall be provided identifying the critical control

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C12	Packaging process List of personnel, their responsibilities and qualifications	points at every stage ii. Manufacturing monograph shall be provided iii. An executed Batch Manufacturing Record for the product under application shall be provided 1)Name, qualification and experience (in years) of the authorized key personnel shall be provided including: • Head of Quality assurance, Quality Control, Storage and		
		production etc., where possible shall provide signatures of the personnel All such Information shall be up to date.		
C13	Letter from Manufacturer to MFDA	The manufacture shall submit a letter to the authority identifying the responsible local representation for the product and this letter shall contain and not limited to: i. Manufacturer details ii. The name and address of local representative authorized to apply for product registration on the manufacturer's behalf iii. Product details iv. Name, designation and signature of the authorizing personnel of the manufacture		
C14	Regulatory decisions taken on this Finished Product from any drug regulatory authorities	 i. A formal, signed statement from the manufacturer and/or MAH is required stating that no regulatory actions such as recalls, bans or alerts have been issued for any batches of the product under applications by any National Regulatory Authorities including that of the country of origin. ii. If any actions as such have been taken by any National Regulatory Authorities regarding product quality, safety and/or efficacy, shall please provide full details with the endorsed 		

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		statement of how the issue was resolved.		
D3	Product (CoPP) Product (CoPP)	 i. CoPP shall be in the format of the WHO Certification Scheme on the Quality of Pharmaceutical Products. ii. A color scanned copy of the original or certified true copy of CoPP certification document issued by the relevant drug regulatory authority of the country of origin shall be submitted which necessarily does not require to be country specific iii. CoPP shall have the following information: Date of issue Expiry date Product name Label claim Excipients (preferred) Name and address of Manufacturer Registration status in exporting country Market availability of product in the exporting country Name and address of issuing authority iv. CoPP Certificate shall have validity of 6 months from the time of submission of the application v. If the certificate is nearing its expiry, evidence of application or under process letter for renewal issued by the same licensing authority shall be submitted along with the current CoPP. vi. If the expiry of the CoPP is not mentioned in the certificate, evidence document shall be submitted for assurance of the validity 		
D4	Registration status of	A list of countries in which the product is registered, including the		
	Finished Product in countries other than	country of origin, shall be provided along with registration number and date of issue. For this purpose, preferably a weblink shall be		
	countries other than	provided with the documents for further verification.		
D5	Proof of registration of	i. This document shall be mandatory when applying under the		
	the Finished Product in	criteria as mentioned in table 1 of this document.		

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	Reference regulatory authority/ies	 ii. This document shall have the same product as that of the product under application iii. For ease of application, a registration certificate of the product or preferably a weblink shall be provided for verification. If the documents are not in English an official, signed and endorsed translation shall be provided. 		
E2	Copy of the Finished Product specification	 i. The finished product specification report shall be based on a reference to an official monograph and if an in-house method is used it shall be endorsed by a third party. The third party can be an accredited laboratory, regulatory authority or any other external or internal assessment body notified by the authority. ii. As per ICH recommendations, finished product specification shall include the following information: * a. Description: a qualitative statement about the state (e.g., solid, liquid), shape and color of the drug substance b. Identification: Identification tests shall be specific for all API(s)/DS. c. Assay: A specific, stability-indicating assay to determine strength (content) in % shall be included for all active pharmaceutical Ingredient(s)/DS. d. Impurities and related substance: Acceptance limits shall be stated for specified degradation products, which may include both identified and unidentified degradation products as appropriate. e. Water content. f. Dissolution. g. Uniformity of dosage units. h. Microbial limits. i. All tests shall specify the pharmacopeial standard used j. Shall include reference pharmacopeial standard used for finished pharmaceutical product (FPP) *Please note the tests can vary based on the type of dosage form and additional tests/criteria shall be included in the specification when the tests are relevant to the quality of the drug substance. 		

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		Refer to ICH Quality Guidelines Q6A-Q6B Specifications.
E5	Certificate of Analysis	i. This certificate shall be a notarized true copy
	for batch release/	ii. This certificate shall contain the following information:
	Certificate of Analysis	 Name and address of the certifying/notarizing authority
	of Finished Product	Batch details
	(CoA)	iii. Batch analyses data from a minimum of 2 batches shall be
		submitted for the product submitted for registration.
		iv. Shall include a conclusion specifying that the product is in
		compliance.
		v. CoA should include result data, reference range and
		pharmacopeial references for each test parameter. For non-
		pharmacopeial test parameter, analysis of the samples will be
		performed using the analytical method and specifications as per
		validated method of analysis.
F. 2.1	Real-Time Stability	1) Stability is the ability of a drug product to retain its chemical,
	Data	physical, microbiological and biopharmaceutical properties within
		specified limits throughout its shelf-life.
		2) Stability tests are a series of tests designed to obtain information
		on the stability of a drug product in order to define its shelf-life and
		utilization period under specified packaging and storage conditions.
		3) Real-time (long-term) stability studies refers to experiments on the
		physical, chemical, biological, biopharmaceutical and
		microbiological characteristics of a drug, during and beyond the
		expected shelf-life and storage periods of samples under the
		storage conditions expected in the intended market. The results are
		used to establish the shelf-life, to confirm the projected shelf life,
		and to recommend storage conditions.
		4) For registration of the product, the authority shall require the

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		manufacturer to submit information on the stability of the product derived from tests on the final dosage form in its final container and packaging. The data submitted is obtained from both accelerated
		and real-time studies. Published and/or recently obtained experimental supporting stability data may also be submitted, e.g.
		on the stability of active ingredients and related formulations.
		5) Both Real time and accelerated stability data shall be submitted separately.
		6) The objective of stability testing is:
		6.1 To select adequate (from the viewpoint of stability) formulations and container closure systems. This is done during the
		development stage of the product via accelerated stability testing
		6.2 To determine shelf-life and storage conditions of the product. This
		testing is done during the development of the product and for the
		registration dossier via both Quality assurances in general, including quality control
		6.3 To substantiate the claimed shelf-life of the product. This is done
		for Registration dossier and is done via real time stability testing.
		6.4 To verify that no changes have been introduced in the formulation or manufacturing process that can adversely affect the stability of
		the product. This is done for quality assurance of the product
		process including quality control and this is done via both Quality assurance in general, including quality control
		7) Where the product is to be diluted or reconstituted before being
		administered to the patient (e.g. a powder for injection or a
		concentrate for oral suspension), "in use" stability data must be
		submitted to support the recommended storage time and conditions for those dosage forms.

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		8) The design of the stability testing program shall take into account	
		the intended market and the climatic conditions in the area in which	
		the drug products will be used. As per ICH zones, following climatic	
		zones are distinguished for the purpose of worldwide stability	
		testing, as follows:	
		 Zone I: Temperate. Zone II: Subtropical, with possible high humidity. Zone III: Hot/dry. Zone IV A: Hot humid / Tropical Zone IV B: Hot/Higher humidity 	
		9) In a stability study, the effect on the product in question of	
		variations in temperature, time, humidity, light intensity and partial	
		vapors pressure (in special cases) are investigated. The effective or	
		mean kinetic temperature therefore reflects the actual situation	
		better than the measured mean temperature; a product kept for 1	
		month at 20°C and 1 month at 40°C will differ from one kept for 2	
		months at 30°C. Moreover, the storage conditions are often such	
		that the temperature is higher than the average meteorological	
		data for a country would indicate.	
		10) For registration purposes, test samples of products containing fairly	
		stable active ingredients shall be taken from two different	
		production batches, in contrast, samples shall be taken from three	
		batches of products containing easily degradable active ingredients	
		or substances on which limited stability data are available. The	
		batches to be sampled shall be representative of the manufacturing	
		process, whether pilot plant or full production scale. Where	
		possible, the batches to be tested should be manufactured from	
		different batches of active ingredients.	
		11) Detailed information on the batches shall be included in the test	
		records, namely the packaging of the drug product, the batch	
		number, the date of manufacture, the batch size, etc.	

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		12) For products containing new APIs/DS, data from stability studies
		shall be provided on at least three primary batches. Two of the three
		batches shall be at least pilot-scale batches and the third batch can
		be smaller (one tenth of full production scale or 100 000 tablets or
		capsules, whichever is the larger, if justified.
		13) For products containing existing APIs (e.g., generics), data shall be
		provided on not less than two batches
		14) Long term or Real-time stability data shall be provided for the
		duration of the proposed shelf life of the product with storage
		condition of 25 °C \pm 2 °C/60% RH \pm 5% RH or 30 °C \pm 2 °C/65% RH \pm
		5% RH or 30 °C ± 2 °C/75% RH ± 5% RH.
		15) Accelerated Stability Data shall be provided for minimum 6
		months with storage condition of 40 °C \pm 2 °C/75% RH \pm 5% RH
		16) For products intended for storage in a refrigerator (2-8°C):
		Long term stability data shall be provided for the duration of
		proposed shelf life with storage condition of 5°C ± 3°C
		Accelerated Stability Data shall be provided for minimum 6
		months with storage condition of 25 °C ± 2 °C or 30 °C ± 2 °C
		17) For products intended for storage in a freezer:
		Long term stability data shall be provided for the duration of
		proposed shelf life with storage condition of –20 °C ± 5 °C
		Accelerated Stability Data shall be provided for minimum 6
		months with storage condition of 5 $^{\circ}$ C \pm 3 $^{\circ}$ C.
		18) The testing shall cover, as appropriate, the physical, chemical,
		biological and microbiological attributes, preservative content and
		functionality tests (e.g., Appearance, Average weight,
		Disintegrating time, pH, Dissolution time, Relative substance,
		Microbial Limit test and Assay)
		19) Analytical procedures shall be fully validated including:
		The orientation of the product during storage, i.e., upright, on

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E12	Accelerated Stability	the side or inverted, where relevant. Date started and end date (Manufactured date/ Expired date) Signature of quality control Packaging of the product It is mandatory to include a conclusion statement in both real time and accelerated stability data indicating that the stability data is acceptable for the specific climate zone. Stability data submitted to this authority for registration purpose must not be more than 5 years after the completion of the study. However, stability studies can be accepted beyond this period if there is no change in the shelf life of the product. For this the applicant shall provide a declaration from the manufacturer. **The results shall be presented in an appropriate format such as tabular, graphical, or narrative description.	
F12 (2.1)	Data Data	 Accelerated stability testing refers to studies designed to increase the rate of chemical degradation and physical change of a drug by using exaggerated storage conditions as part of the formal stability testing programme. The data thus obtained, in addition to those derived from real-time stability studies, may be used to assess longer-term chemical effects under non-accelerated conditions and to evaluate the impact of short-term excursions outside the label storage conditions, as might occur during shipping. The results of accelerated testing studies are not always predictive of physical changes. Accelerated stability tests provide a means of comparing alternative formulations, packaging materials, and/or manufacturing processes in short-term experiments. As soon as the final formulation and 	

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F3	Stability Report and statement	manufacturing process have been established, the manufacturer carries out a series of accelerated stability tests which will enable the stability of the drug product to be predicted and its shelf-life and storage conditions determined. Real-time studies must be started at the same time for confirmation purposes. Suitable measures should be taken to establish the utilization period for preparations in multidose containers, especially for topical use. 1) A brief summary of stability report shall be established and shall be submitted with the dossier giving details of the design of the study, as well as the results and conclusions. The stability of a given product, and therefore the proposed shelf-life and storage conditions, must be determined on the basis of these results 2) An official statement issued by the manufacturer that all stability tests are performed of the same formula, manufactured at the same site(s) and packed in the same packing material as the product shall	
G1 (1.1)	In vivo Bioequivalence Study Required for all immediate release oral solid dosage forms only (i.e., tablets, capsules)	be provided with the dossier. This statement shall be signed and endorsed by the manufacturer. The Bio equivalence study is the comparative analysis between the innovator or comparator or reference drug product with that of the product submit for registration to assure that the product submitted for registration can show the same efficacy as that of the innovator or comparator or reference drug product.	

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		provided that the reference product used in study has established bioavailability comparative to the innovator drug product
		2.The study report shall contain the following information:
		 Information about the reference and test products,
		such as the product name, strength, dosage form, batch
		number, manufacturing site, batch size of the test
		product, etc. The reference product shall have the
		exact same strength and in the exact same formulation
		the product submitted for registration.
		Certificates of Analysis of the reference and test
		products used in the BE study, including the batch size
		of the test product and manufacturing/expiry date of
		both products (where applicable)
		Bioanalytical study report summary and description of
		the bioanalytical method validation
		3.A complete bioequivalence study report including all appendices and
		data and conclusive statement of the end results shall be provided.
		4. A signed statement confirming that the test product used in the BE
		study is the same formulation and is manufactured by the same
		process as the product submitted for registration shall be provided
	V	5. Bio waiver requests:
		The biopharmaceutics classification system (BCS)
		classified APIs into the following groups based on level
		of solubility and permeability.
		BCS class I HIGH solubility and HIGH permeability
		BCS class II LOW solubility and HIGH permeability

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		BCS class III HIGH solubility and LOW permeability
		 BCS class IV LOW solubility and LOW permeability
		 As per criteria set by WHO, products containing API's
		belonging to BCS class I and BCS class III qualify for a BCS-
		based biowaiver request, provided that the following
		criteria is met and surrogate information is submitted.
		6. Criteria for biowaiver**
		 The API shall belong to BCS class I, or BCS class III.
		 The product shall not be a narrow therapeutic index
		(NTI) drug.
		**Please note that biowaivers submitted for pharmaceutical products that do not fit the criteria described will not be accepted even if a comparative dissolution profile is provided.
		7. Data to support requests for biowaiver.
		Data supporting high solubility of product
		 Data supporting high permeability of product
		8. A satisfactory dissolution study with reference product as per
		criteria described below:
		Demonstrate that the excipients used are well-
		established and do not alter the pharmacokinetics of
	\cup	API.
		9. Bio waiver request for multiple strength product
		 If the test product used in the BE study is of a different
		strength from that proposed for registration, a signed
		statement confirming that the test product used in the
		BE study has the same qualitative composition and
		quantitatively proportional composition and is
		manufactured by the same process as that proposed
		for registration shall be submitted.

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		 10. Additional justifications and relevant supporting documents for bio waiver requests shall be submitted if requested by the authority. 11. A statement signed and endorsed by the manufacturer shall be submitted stating the conclusion of the bioequivalence study indicating that the product applied for registration is bioequivalent to the innovator or the reference product
G1	In vitro Dissolution	
(1.2)	Test	The dissolution profiles shall be submitted by following the method described in the monograph of the relevant pharmacopoeia.
		2. If in-house method is used it shall be endorsed and validated by a third party. The third party can be an accredited laboratory, regulatory authority or any other external or internal assessment body as nominated by the Authority.
		3. The following data shall be submitted:4. Information about the reference/innovator/comparator drug
		product and test products, such as the product name, strength,
		dosage form, batch number, manufacturing site, batch size of the
		test product, etc.
		5. The dissolution apparatus, media, results and the conditions at
		which it is operated shall be specified and in accordance with an
		established pharmacopeia dissolution test guideline. (e.g. European
		Pharmacopoeia (Ph. Eur.), United States Pharmacopeia (USP) etc.).
		6. The reference/innovator/comparator drug product shall have the
		exact same strength and in the exact same formulation the product
		submitted for registration.
		A complete Dissolution study report including all appendices
		and data and conclusive statement of the end results shall be
		provided.
		A statement signed and endorsed by the manufacturer shall be

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		submitted stating the conclusion of the dissolution study indicating that the product applied for registration is in compliant with the requirements stated.
Н5	Product Label/Packing	1. Product label shall contain the following information;
	insert	 Brand name, Generic name, strength and dosage form Full manufacturing site address of the product. Exemptions: In case the manufacturer is not defined in the label, a specific code (bar code or QR code) shall be in the label to trace the manufacture name and address, and this shall be same as with the submitted document. Direction for use Special precaution if applicable Shelf life Storage condition Shall be submitted in English.
		2. The draft artwork of the outer carton labels shall be in the actual format, design and colour that are to be printed.3. Separate labels shall be submitted for each pack size of the product.
		4. Packing insert/Patient leaflet criteria: Packing insert/ Patient leaflet/
		SmPC shall be clear, concise and shall contain the following information:
		Name of the product: The brand name shall be followed by both the strength and the pharmaceutical form.
		The International Nonproprietary Name (INN) or the usual common name of the API/DS shall be used when referring to
		properties of the active substance(s) rather than the brand name.
		• Strength: The strength shall be the relevant quantity for
		identification and use of the product and shall be
		consistent throughout other sections of the packing insert/patient leaflet

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Code	Document Title	Acceptance Criteria
		Pharmaceutical form/Dosage form: The dosage form
		of a product shall be described by a single full standard
		term according to the relevant pharmacopeia used.
		• Composition: Full details of the qualitative and
		quantitative composition in terms of the APIs/DS shall
		be provided as a separate subheading qualitatively and
		quantitatively.
		• The active substance shall be written in its
		recommended INN, accompanied by its salt or hydrate
		form if relevant.
		• The quantity of the active substance shall be expressed
		per dosage unit and in an internationally recognized
		standard term.
		• Indication: The indication(s) shall be stated clearly and
		concisely and shall define the target disease or
		condition distinguishing between treatment
		(symptomatic, curative or modifying the evolution or
		progression of the disease), prevention (primary or
		secondary) and diagnostic indication.
		Where required, it shall define the target population
		especially when restrictions to the patient populations
		apply.
		 It shall be stated in which age groups the product is indicated, specifying the age limits
		 Dosage: The dosage shall be clearly specified for each
		method/route of administration and for each
		indication, as appropriate.
		Dosage adjustments or other posology related
		information in specific patient groups shall be stated
		where necessary, in well-defined sub-sections, e.g.

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Code	Document Title	Acceptance Criteria				
		elderly population, renal impairment and other				
		relevant special population				
		 If the product is indicated in the pediatric population, 				
		dosage and administration recommendations shall be				
		given for each of the relevant subsets				
		 If there is no indication for the product in some or all 				
		subsets of the pediatric population, no dosage				
		recommendation can be made, but available				
		information shall be provided using the following				
		standard statements such as "No data are available" or				
		"The safety and efficacy of the product in children aged				
		x to y has not been established"				
		• Administration: The route of administration and				
		concise relevant instruction for correct administration				
		and use shall be given here. Information on instructions				
		for preparation or reconstitution shall be provided				
		Any specific recommendation for use related to the				
		dosage form shall be explained e.g., Tablet shall not be				
		crushed due to "xyz"				
		Contraindications: All situations and circumstances				
		where the drug product shall not be given for safety				
		reasons shall be clearly defined and explained				
		Special warnings and precautions for use: Special				
		patient groups that are at increased risk or are the only				
		groups at risk of experiencing product or product class-				
		related adverse reactions				
		Any special precautions related to the administration or				
		use of the drug product by the healthcare professionals,				
		the patient or caregivers				
		Any need for specific clinical or laboratory monitoring				

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Code	Document Title	Acceptance Criteria	
Code	Document Title	for use during pregnancy and lactation on the basis of increasing human experience in exposed pregnancies which eventually supersede the animal data. • If there is no data available at all, then this shall be clearly stated. • Side effects/Undesirable effects: This section shall include all adverse reactions from clinical trials, safety studies and spontaneous reporting for which, after thorough assessment, a causal relationship between the medicinal product and the adverse event is at least a reasonable possibility. This section shall be regularly reviewed and, if necessary, updated with the aim to ensure appropriate information to health care professionals on the safety profile of the product. • Overdose: Describe acute symptoms and signs of different dose levels of the medicinal product based on all available information including accidental intake,	
		different dose levels of the medicinal product based on all available information including accidental intake, mistakes and suicide attempts by patients. • Taking into account all relevant evidence, describe the	
		management of an overdose e.g. in relation to monitoring or use of specific agonists/antagonists, antidotes (no dosage recommendations) or methods to increase elimination of the medicinal product such as dialysis.	
		 Pharmacodynamics/Pharmacokinetic properties: Pharmacokinetic/Pharmacokinetic properties of the active substance(s) relevant for the advised dose, strength and the pharmaceutical formulation marketed shall be given in this section. Shelf life 	

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Code	Document Title	Acceptance Criteria
		 Special precautions for storage Date of publication/revision Innovator drug products shall follow the above information as per approval of SRA. Generic drug products shall follow and maintain the consistency in above information with the innovator drug products as approved by SRAs. Please note that the information listed above is not meant to be exhaustive and any information that is vital in the safe and effective administration of the product by healthcare professionals shall be included.
11	Cost and Retail price	 Cost price (USD) shall be provided specifying the quantity. E.g., per tablet or per 5ml composition. Proposed price for retail in Maldives (USD) be provided specifying the quantity. Eg. per tablet or per 5ml composition The price structure shall include the name of the product.

11.3 Additional Requirements for Non-Pharmacopeial Products for full dossiers.

11.3.1 For in-house methods, other than official methods as per the official pharmacopeias, in house method validation report shall be submitted which shall have the parameters or validation characteristics for testing methods as under:

	Testing Methods		Validation Parameters
1)	Identification	a.	Specificity
2)	Assay (content and dissolution measurement	b.	Accuracy
	only)	c.	Precision (repeatability, intermediate)
3)	Impurities (quantitative & limit test)	d.	Linearity & Range
,	3) Impurities (quantitative & limit test)	e.	Detection Limit
		f.	Quantitation Limit
		g.	Robustness

- **11.3.2** In addition to method validation report, in-house method shall also provide documentary evidence for the identification of sources and quantitation of potential errors, determine if the method is acceptable for intended use and establish proof that a method can be used for decision-making.
- **11.3.3** The in-house method validation report shall be endorsed by a third party. The third party can be an accredited laboratory, regulatory authority or any other external or internal assessment body as approved by NRA.

11.4 Submission of Product Samples

- **11.4.1** Applicants are required to submit the product samples to the MFDA for all new products in the quantities as described below for each type for full dossier applications. However, MFDA can request more quantities if required for analysis or as notified from time to time for various dosage forms.
- **11.4.2** Samples are not required for reliance pathways registration. However, samples are required for re-registration for products under reliance pathways.

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Туре	Description	Quantity
Liquid dosage forms	These include solutions, syrups,	03 units shall be submitted.
	elixirs, suspensions, emulsions etc.,	(e.g. 03 bottles of liquids.)
Solid dosage forms	These include tablets, capsules, ,	60 units shall be submitted
	lozenges, etc.	(e.g. 60 tablets, 60 capsules).
Semisolid dosage	These include creams, ointments,	
forms	gels, pastes, and suppositories.	
Parental Preparations	These include ampules, vials,	03 units shall be submitted
	infusions, etc.	(e.g. 03 vials or ampoules of
		injections)
Inhalational products	These include inhalers, nebulizers	
Others	Transdermal patches,	

- **11.4.3** Request for sample import shall be applied online through Dhirithi portal. MFDA considers requests for grant approval within 20 working days. Samples shall only be imported once sample authorization approval has been issued.
- 11.4.4 All imported sample (full quantity imported) shall be handed over to MFDA once it is cleared from the ports and these samples shall be handed over to the product registration unit. The Sample submission sheet. (Annex VII) shall be submitted via e mail (medicine.registration@health.gov.mv) before handing over the samples.
- **11.4.5** Samples shall be submitted to MFDA within 72 hours of the clearance from ports and it shall be submitted on Mondays and Thursdays from 10.00hours to 12.00hours.
- **11.4.6** Samples shall tally with the documents submitted for registration, otherwise the application shall be rejected.
- **11.4.7** In case an application is rejected, the samples shall be kept in MFDA for 60 days from the date of rejection and then they will be disposed of as per recommended method.

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- **11.4.8** National Health Laboratory (NHL) is the designated national laboratory for testing pharmaceuticals. Samples that are submitted with the dossiers are also tested from NHL depending on the testing capacity of the laboratory for that specific product.
- **11.4.9** Upon testing, if the sample fails in any of the qualitative analysis parameters, the product will be rejected.
- **11.4.10** The product will also be rejected if the quantitative analysis fails.
- **11.4.11** The reason of rejection of the application will be informed and updated via the dhirithi portal.

12 Application Submission and Review Process

- 12.1 Product registration application submission.
- **12.1.1** All product registration Applications shall be submitted Online via Dhirithi portal 'https://dhirithi.egov.mv'.
- **12.1.2** The applicant shall first register as a user in Dhirithi portal using the form available on the MOH website, under, https://health.gov.mv/dv/downloads/dhirithi-portal-user-registration-form. The form is also available in Dhrithi portal under "Publications". If the applicant is an authorized medicine importer, they would already be registered as a Dhirithi user and hence can directly apply for medicine registration.
- **12.1.3** Once the applicant is registered in the Dhirithi portal, the applicant can then select "Pharmaceutical" and then "medicine registration" and submit the dossier.
- **12.1.4** To ensure all mandatory documents are submitted, the applicant shall refer to
- Table 1 of this document to select the registration pathway and identify the required documents relevant to that pathway
- 2. Application checklists for reliance in Annex V
- 3. Application checklists for full dossier in Annex VI

12.2 Application submission fee

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- 12.2.1 If all the requirements are complete, the application/dossier will be accepted and a submission fee of 100 MVR (hundred Maldivian Rufiyaa) shall be paid via Bandeyri Pay (https://bp.finance.gov.mv/), within 5 working days from the time of dossier acceptance to Dhirithi portal. If the payment is not made within the given 5 days, the dossier will be rejected.
- **12.2.2** This submission fee is non-refundable.
- **12.2.3** Once the payment is made the evaluation process of the application/ dossiers will be initiated with regards to safety, quality and efficacy of the product. The time lines for the process is indicated in Table 1 of this document.

12.3 Pre-Screening of Dossier

- **12.3.1** Once the application/dossier is submitted, it will be checked for document completion and legibility. If all the requirements as per the acceptance criteria are fulfilled, then only the dossier will be accepted.
- **12.3.2** MFDA shall have the right to reject incomplete dossiers and hence it's the applicant's responsibility to ensure that all are in accordance with the requirements as mentioned.
- **12.3.3** Applications/Dossiers that require further clarification will be put to "Need clarification" status on Dhirithi portal. The clarification requested by the Authority shall be resolved within 10 working days by the applicant, otherwise, the dossier will be rejected.
- **12.3.4** In the case of a rejection, the reason for the rejection will be specified.
- **12.3.5** The application/dossier evaluation process may be prolonged due to the time taken by the applicant to respond to the Authority's request to provide additional information or further clarification.

12.4 Assessment of Application

12.4.1 Pharmaceutical officers of Medicine Registration act as assessors and verify the required documents, accept the applications/dossiers, and evaluating the applications/dossiers as per approved criteria. In case further clarification is required, these officers are also responsible for effective communication with the applicant in a timely manner.

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- **12.4.2** A summary of the evaluation is generated in the approved format by the Assessors and submitted before the National Pharmaceutical Board for Pharmaceuticals for decision-making.
- **12.4.3** The Director, Pharmaceuticals (regulation Section) is responsible for cross-review by checking and verifying the product evaluation documents submitted in the dossiers, and to guide the pharmaceutical officers on evaluating the product.

12.5 Decision by the National Pharmaceutical Board (NPB)

- **12.5.1** Upon successful evaluation of the applications/dossier, the documents are submitted to the National Pharmaceutical Board (NPB) for approval or rejection. The status will read as "Pending committee decision" at this stage on Dhirithi portal.
- 12.5.2 If recommended to approve the product by the NPB, the applicant will be notified to pay a registration fee of 300 MVR via Bandeyri Pay within 5 working days. If the payment is not made within the given five days, the client will be informed via email. If the payment is not made within 10 working days of the notification, the application will be rejected. The applicant must process this dossier as a new application again.
- **12.5.3** If recommended to reject the product by the NPB, the dossier will be rejected, and the applicant will be notified via Dhirithi portal indicating the reason for rejection.
- **12.5.4** The Deputy Director General (Medicine Therapeutic Goods Division) and Director General MFDA is responsible to finally approve and authorize the product after it's been approved by the National Pharmaceutical board.

12.6 Scientific Expert committee for Medicine Registration

- **12.6.1** The Authority shall establish an expert Committee to obtain scientific evidence and provide technical assistance to the National Pharmaceutical Board if required.
- **12.6.2** The main function of this committee is to obtain scientific evidence in determining the necessary measures and decisions regarding the safety of drugs and to recommend a decision based on this information after conducting a risk benefit assessment. In addition, the purpose of this committee is to:
 - Advice on taking the decision on the registration of Pharmaceuticals Products and Medical Devices.

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- 2. Advice on taking all necessary steps to strengthen the inspection system.
- 3. Advice on conducting a risk benefit assessment to change the category of the drug and plan based on the results of this assessment.
- 3. Provide advice and guidance on adverse drug reactions, development and strengthening of the management system.
- 4. Advice on making decisions regarding the safety of vaccines and biologicals.
- Scientific evidence and risk benefit assessment required for approval of a new drug.
- 6. Guide and advice on establish public awareness system on drug safety and to participate in awareness programs
- 7. Provide Guidance to take necessary steps to strengthen the regulatory system of drugs.
- 10. Technical advice in the preparation of drug related rules and regulations

12.7 Members of the Scientific Expert committee

- 12.7.1 The committee shall consist of a member representing the Pharmaceutical Board.
- **12.7.2** Technical members from the pharmaceutical or pharmaceutical sector shall be represented as members on the committee.
- **12.7.3** Committee meetings shall be held at least four times a year. The number of members of this committee is 05 (five). The quorum is 03(three).
- **12.7.4** The members of the committee shall be independent of the activities of the pharmaceutical industry (manufacturing, importing and trading of pharmaceuticals).
- **12.7.5** The Maldives Food and Drug Authority will be responsible for the secretariat of the committee.
- **12.7.6** An expert/experts may be invited to the meetings of this committee for advice on drug related matters.

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12.7.7 A person in the field may be invited to attend a committee meeting to discuss a matter relating to alternative medicine as described in (a) of this Article. Veterinarians or a veterinary expert can be invited to the meeting to discuss the medication used for animals.



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12.8 Issuance of Registration Certificate

- **12.8.1** After the registration fee has been paid, the Authority will issue a Certificate of Registration of a Drug Product and Agreement to the applicant within 20 working days.
- **12.8.2** The applicant will be notified via email to report to MFDA 2nd floor for the agreement signing and certificate issuance within this period. The applicant shall bear the responsibility of attending the signing and failure to attend within 15 working days of notification will result in cancellation of the agreement.
- **12.8.3** The product can only be imported, distributed and sold in the country once it has been registered and added to the Approved Drug List (ADL).

12.9 Validity of Registration

12.9.1 The validity period of the registration is mentioned in the table 1 of this document.

12.10 Classification of Medicine Registration

12.10.1 Registration will be issued under the following classifications which determine the level of access control.

Classification	Remark
Restricted for Hospital and Institutional (HI) use only	Medicinal products restricted to special expertise and Health facilities and clinics with registered medical practitioners. These products cannot be kept for sale in pharmacies. These products can only be imported by designated parties.
Restricted for Hospital use only (HO)	These medicinal products can only be imported and registered by designated parties. These products cannot be kept for sale in pharmacies. These products are restricted to special expertise for

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Classification	Remark
	hospitals only
Controlled Drug C	These medicinal products are controlled and can only be imported by designated parties. Within this class, Narcotics cannot be kept in pharmacies for sale. Controlled Drugs include Narcotics and Psychotropic drugs (Internationally and Nationally Controlled).
Over The Counter Medicine (OTC)	These medicinal products can be sold without prescriptions.
Prescription Only Medicines (POM)	These medicinal products can only be prescribed by a registered medical practitioner. This product can only be sold with a valid prescription.

12.11 Cancellation, Suspension, Revocation of Registration

- **12.11.1** The Maldives Food and Drug Authority reserves the right to reject, cancel or suspend the registration of any product if:
 - a. There are deficiencies in safety, quality, or efficacy of the product.
 - b. Failure to comply with conditions of registration.
 - c. If the product has quality issues from Market surveillance.
 - d. If the product is removed from Approved Drug List due to any reason including safety and efficacy.
 - e. The information provided at the time of the submission of the application is later found to be false or insufficient.
 - f. If it is substantiated that the formulation has serious side effects and any of Reference Regulatory Authority or WHO or other national and international agency prohibited used of such formulation.
 - g. If it is found that the manufacturer is not in compliance with Good Manufacturing Practices (GMP), or for any other reasons, that the manufacturer has repeated violations like manufacturing of sub-standard drug products.
 - h. If any adverse regulatory action is taken against the manufacturer abroad by the regulatory authority of country of origin.
 - i. If the MAH fail to inform the MFDA of any serious adverse reactions of the registered product upon receipt of such reports.
 - j. If any post-registration variation has been done including the composition, label, packaging, manufacturing method, drug product specifications, indication or any other particulars of the product has been changed without the approval of the MFDA.
 - k. If foreign manufacturer of the registered product has decided to withdraw and not to sell the product.
 - I. Any of the conditions of registration of the product have been contravened.
 - m. Any report on adverse drug reactions of a serious nature has been received from national or international sources.

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n. For any other matters as specified by the National Pharmaceutical Board at the time of cancellation.

12.12 Temporary and Permanent Ban of a Product and/or Manufacturer

- **12.12.1** A company or manufacturer can be temporarily or permanently banned if repeated incidents of quality failure are identified from their products.
- **12.12.2** If the company or manufacturer is permanently banned, the company will not be allowed to enter the Maldives market until a period of a minimum 5 years has passed. After 5 years from the date of the ban, they can apply for registration with the current criteria only. Reliance pathways is encouraged for such products.
- **12.12.3** The final decision for the product approval will be made based on the technical advice from National Pharmaceutical Board
- **12.12.4** If a company or manufacturer is temporarily banned, the products of that company can only enter back into the market as a new applicant under the current procedure. All products of that company shall be registered as per the criteria established by the MFDA.
- **12.12.5** Due to a quality issue, if a product is recalled from the market or product application rejected during registration or re-registration, these products can only be submitted with additional documents in addition to the documents required in the registration pathways. The additional documents shall include and is not limited to:
 - a. Product tested reports from a WHO prequalified laboratory
 - b. Manufacturer's declaration of the quality, safety and efficacy of the product.
 - c. Product assessment report from a Stringent NRA
- **12.12.6** If the products fail repeatedly (more than 2 times) as mentioned in point 12.12.5, the product shall be rejected, and it shall not be again submitted for a period of 5 years.

13 Post Registration Variations

13.1 Classification of Variations

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- **13.1.1** Following are the basis of classification of post-registration variations.
 - a. An administrative change such as a change of company name and/or address.
 - b. A change to the characteristics of drug product that can affect its quality like change of storage condition.
 - c. A change to the safety, efficacy or pharmacovigilance information of the product.
- **13.1.2** Types of variations for a registered drug product are classified as Minor and Major.

13.2 Minor Variations (MiV)

13.2.1 Variation to a registered finished product in terms of changes which has minimal or insignificant impact on the aspects of safety, efficacy and quality. Minor variations are further divided into following sub types: -

1) Minor Variations – Notification (MiV-N)

- a. Minor Variations Notification (MiV-N) have little or no impact on the safety, efficacy and quality of registered drug product e.g. administrative modifications.
- b. MiV-N procedures are classed a 'do-and-tell' procedure, means registration holder should implement the change and intimate / notify to MFDA by fulfilling the conditions and supporting documents.
- c. Applicant should ensure this notification/intimation must reach relevant section of MFDA within 30 days of implementation of change for MFDA's record and it can be considered accepted if an objection is not issued by the MFDA within 30 working days of the date of submission of variation application.

1) Minor Variations – Prior Approval (MiV-PA)

- a. If the change is more significant than (MiV-N) change but it does not fall under major variation category, then it is considered as MiV-PA change.
- b. These changes need prior approval from MFDA before implementation. (e.g. Change in brand/proprietary name, title of firm, etc.).
- c. Registration holder / MA holder is required to submit a variation application for the proposed change to relevant section of MFDA along with supporting documents and fulfill the conditions as described in these guidelines.

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d. If the application fulfills the prescribed criteria, Medicine Therapeutic Goods Division will process the case for approval of the proposed change by Head of the Division.

2) Major Variations (MaV)

- a. Variation to a registered pharmaceutical finished product that may affect significantly and/or directly the aspects of quality, safety and efficacy and it does not fall within the definition of minor variation are considered as major variations.
- b. Registration holder needs to seek prior approval for major variations before they are made. Registration holder MA holder is required to submit an application for proposed variation to Medicine Therapeutic Goods Division, MFDA along with supporting documents and fulfill the conditions as described in these guidelines.

Details of Major variations (MaV-1)

MaV-1	Change and/or additional indication/dosing regimen/patient					
	population/inclusion of clinical information extending the usage of the					
	product					
Conditions to be	1. Product labeling refers to Package Insert (PI), Patient Information Leaflet					
fulfilled	(PIL), unit carton label, inner label and/or blister strips.					
	2. As a subsequent change due to revision of Summary of Product					
	Characteristics (SmPC) or equivalent document (USPI)					
Documents to be	1. Approved product labeling.					
submitted	2. Proposed product labeling, a clean and annotated version highlighting the					
	changes made.					
	3. Approved PI/SmPC/PIL from an approved reference regulatory agency or the					
	country of origin containing the proposed changes (where applicable).					
	4. Justifications for the changes proposed.					
	5. Approval letters from reference countries or country of origin which have					
	approved the proposed indication or dosing regimen (where applicable).					
	6. Clinical expert reports and/or clinical trial reports (where applicable).					
	7. Clinical documents as per Common Technical Dossier (CTD) format (where					
	applicable).					

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MaV-2			Change of content of product labeling				
Conditions	to	be	1. Product labeling refers to Package Insert (PI), Patient Information Leaflet				
fulfilled			(PIL), unit carton label, inner label and/or blister strips.				
			2. The change is not a minor variation and not within the scope of MaV-1.				
			3. As a subsequent change due to revision of Summary of Product				
			Characteristics (SmPC) or equivalent document (USPI).				
Documents	to	be	1. Approved product labeling.				
submitted			2. Proposed product labeling, a clean and annotated version highlighting the				
			changes made.				
			3. Approved PI/SmPC/PIL from an approved reference regulatory agency or the				
			country of origin containing the proposed changes (where applicable).				
			4. Justifications for the changes proposed and supporting clinical documents				
			when applicable.				
MaV-3			Change of batch size of sterile drug product				
Conditions	to	be	1. The change does not affect the consistency of production.				
fulfilled			2. The product formulation remains unchanged.				
			3. Release and shelf-life specifications of drug product remain unchanged.				
			4. Process validation scheme and/or report is available or validation of the				
			manufacturing process has been successfully carried out according to protocol				
			with at least three batches appropriate to the proposed batch size.				
Documents	to	be	1.Comparative tabulated format of approved and proposed batch				
submitted			manufacturing formula.				
			2. Validation scheme and/or report of the manufacturing process				
			3. Release and shelf-life specifications of the drug product.				
	7		4. Certificate of analysis and/or batch analysis data (in a comparative tabulated				
			format) of drug product of at least two production batches manufactured				
			according to approved and proposed batch sizes.				
			5. Stability data				
MaV-4			Change of batch size of non-sterile drug product				
Conditions	to	be	1. The change does not affect consistency of production.				

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fulfilled			2. The product formulation remains unchanged.
			3. Release and shelf-life specifications of drug product remain unchanged.
			4. Process validation scheme and/or report is available or validation of the
			manufacturing process has been successfully carried out according to protocol
			with at least three batches.
Documents	to	be	1. Comparative dissolution profile data of at least one pilot/production batch
submitted			of the drug product manufactured in the approved and proposed batch size for
			oral solid dosage forms (where applicable).
			2. Comparative tabulated format of approved and proposed batch
			manufacturing formula.
			3. Validation scheme and/or report of the manufacturing process 4. Release
			and shelf-life specifications of the drug product.
			5. Certificate of analysis and/or batch analysis data (in a comparative tabulated
			format) of drug product on a minimum of one production batch manufactured
			according to approved and proposed batch sizes and letter of undertaking to
			submit batch analysis data on the next one full production batch.
			6. Stability data and Stability Study of Drug Product and report if any results
			fall outside shelf-life specifications (with proposed action).
MaV-5			Major change in the manufacturing process for drug product
Conditions	to	be	1. The change does not cause a negative impact on the quality, safety and
fulfilled			efficacy of the drug product.
			2. The manufacturing site remains unchanged. If there is a change in
			manufacturing site.
Documents	to	be	1. Description of the proposed manufacturing process and technical
submitted			justification for the change.
			2. Comparative dissolution profile data of at least one pilot/production batch
			of the drug product manufactured in the approved and proposed
			manufacturing process for oral solid dosage forms.
			manufacturing process for oral solid dosage forms. 3. Validation scheme and/or report of the proposed manufacturing process

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			of proposed release and shelf-life specifications that supports that the
			proposed process must lead to an identical or better product regarding all
			aspects of quality, safety and efficacy.
			5. Certificate of analysis and/or batch analysis data (in a comparative tabulated
			format) of drug product for a minimum of one production batch manufactured
			according to approved and proposed processes.
			6. Stability data or Stability Study of Drug Product report if any results fall
			outside shelf-life specifications (with proposed action).
MaV-6			Qualitative or quantitative change of excipient
			a) For immediate release oral dosage forms
			b) For modified release oral dosage forms
			c) For other critical dosage forms such as sterile preparations.
Conditions	to	be	1. Change will need to comply with the finished product specifications for
fulfilled			example release and shelf-life specifications of the drug product remain
			unchanged, excluding product description except for update of product
			description with respect to appearance/odour/taste as a consequence of the
			change (where applicable).
			2. Replacement of an excipient with a comparable excipient of the same
			functional characteristics.
			3. The dissolution profile of the proposed product is comparable to that of the
			approved product.
			4. Process validation scheme and/or report is available or validation of the
			manufacturing process has been successfully carried out according to protocol
			with at least three batches of the proposed product formula
Documents	to	be	1. Revised drafts of the package insert and labeling incorporating the proposed
submitted			variation.
			2. A declaration that the proposed excipient does not interfere with the drug
			product release and shelf-life specifications test method.
			3. Justification for the change must be given by appropriate development of
			pharmaceutics.
			4. Comparative tabulated format of the approved and proposed product

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formulation with calculated changes highlighted (please state changes in the percentage of the proposed excipient out of the total target dosage form weight (where applicable). 5. Comparative dissolution profile data of at least one pilot/production batch of the drug product manufactured in the approved and proposed formulation for oral solid dosage forms 6. Revised batch manufacturing formula. 7. Validation scheme and/or report of the manufacturing process 8. Specifications of the proposed excipient. 9.. For proposed excipients made of ruminants source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform Encephalopathy (BSE)-free cert issued from relevant authority of the issuing country and/or documentary evidence from the supplier (where applicable). 10. Drug product release and shelf-life specifications. 11. Certificate of analysis and/or batch analysis data (in a comparative tabulated format) of drug product on at least two productions (or one production batch and two pilot batches) according to approved and proposed product formula. 12. Stability data or Stability Study of Drug Product and report if any results fall outside shelf-life specifications (with proposed 13. For quantitative and qualitative changes in preservative, results of Preservative Effectiveness Test (PET) at lowest specified preservative level (where applicable). MaV-7 Quantitative change in coating of tablets and/or size of capsule shell for modified release oral dosage form Conditions to be 1. The dissolution profile of the proposed product is comparable to that of the fulfilled approved product. 2. The release and shelf-life specifications of the drug product remain unchanged except for the weight and/or size 3. For quantitative change in coating of tablets or weight and/or size of capsule shell for immediate release oral solid dosage forms 1. Revised draft of product label incorporating the proposed change **Documents** to be

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submitted	2. A declaration that the change does not interfere with the drug product				
	release and shelf-life specifications test method.				
	3. Comparative dissolution profile data of at least one pilot/production batch				
	of the drug product manufactured in the approved and proposed composition				
	for oral solid dosage forms				
	4. Approved and proposed product and batch manufacturing formula.				
	5. Revised release and shelf-life specifications of the drug product. 6. Stability				
	data or Stability Study of Drug Product and report if any results fall outside				
	shelf-life specifications (with proposed action).				
MaV-8	Change in primary packaging material for sterile product				
	a) Qualitative and quantitative composition and/or				
	b) Type of container and/or				
	c) Inclusion of primary packaging material				
Conditions to be	1. Release and shelf-life specifications of the drug product remain				
fulfilled	unchanged.				
	2. For change in the primary packaging material for non-sterile drug				
	product				
Documents to be	1. Revised drafts of the package insert and labeling incorporating the proposed				
submitted	variation.				
	2. Appropriate scientific data on proposed packaging (comparative data on				
	permeability, e.g. moisture, O2, CO2).				
	3. Proof must be provided that no interaction between the content and the				
	packaging material occurs				
	4. Validation scheme and/or report of the manufacturing and sterilization				
	process				
	5. Comparative tabulated format of specifications of the approved and				
	proposed primary packaging material.				
	6. Stability data or Stability Study of Drug Product and report if any results fall				
	autoide chalf life annoificetions (with mannon dection)				
'	outside shelf-life specifications (with proposed action).				

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			dimension of container or closure for sterile solid and liquid drug product			
Conditions	to	be	1. The proposed pack size is consistent with the dosage regimen and duration			
fulfilled			of use as approved in the package insert.			
			2. The packaging material remains unchanged.			
			3. Release and shelf-life specifications of the drug product are not affected			
			except pack size/fill volume specification.			
			4. Change or addition of pack size/fill volume and/or change of shape or			
			dimension of container or closure for non-sterile drug product.			
Documents	to	be	1. Revised drafts of the package insert and labeling incorporating the proposed			
submitted			variation.			
			2. Justification that the proposed pack size is consistent with the dosage			
			regimen and duration of use as approved in the package insert.			
			3. Validation data of the manufacturing process, sterilization and container			
			closure system (where applicable).			
			4. Stability data or Stability Study of Drug Product and report if any results fall			
			outside shelf-life specifications (with proposed action).			
MaV-10			Inclusion or replacement of the solvent/diluent for the drug product			
Conditions	to	be	1. The proposed change does not result in any change in the dosage form,			
fulfilled			regimen, indication, method of administration of the product.			
			2. For deletion of the solvent/diluent.			
			3. For change of shelf-life and/or storage condition of the drug product after			
			first opening and/or after dilution/reconstitution.			
Documents	to	be	1. Revised drafts of the package insert and labeling incorporating the proposed			
submitted			variation.			
			2. Documentary evidence to certify the manufacturing site of diluents/solvents			
			complies with current applicable GMP standards.			
			3. Batch numbering system.			
			4. A letter of authorization from product owner to authorize the manufacturing			
			site to manufacture and package the solvent/diluent.			
			5. A declaration from the marketing authorization holder that the release and			
			shelf-life specifications of drug product are not affected.			

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			6. In addition to section P for the solvent/diluent and reconstitution stability
			data, section S is also required (where applicable).
MaV-11			Extension of shelf-life of the drug product
			a) As a package for sale and/or
			b) After first opening and/or
			c) After dilution/reconstitution
Conditions	to	be	1. For (a) & (b) - The studies must show conformance to the approved shelf-life
fulfilled			specification.
			2. For (c)—The studies must show conformance to the approved shelf-life
			specification for the reconstituted product.
Documents	to	be	1. Revised drafts of the package insert and labeling incorporating the proposed
submitted			variation (where applicable).
			2. Technical justification for the proposed change (where applicable).
			3. A letter of commitment from product owner or marketing authorization
			holder to inform users of the relevant change (where applicable).
			4. Results of appropriate long term stability studies covering the duration of
			proposed shelf-life of the product in the authorized packaging material
MaV-12			Change of storage conditions of the drug product (Lowering from the
			approved storage condition) a) As a package for sale and/or b) After first
			opening and/or c) After dilution/reconstitution
Conditions	to	be	1. For (a) & (b) - The studies must show conformance to the approved shelf-life
fulfilled			specification.
			2. For (c) — The studies must show conformance to the approved shelf-life
			specification for the reconstituted product.
Documents	to	be	1. Revised drafts of the package insert and labeling incorporating the proposed
submitted			variation.
			2. Technical justification for the proposed change.
			3. Results of appropriate long term stability studies covering the duration of
			approved shelf-life (at proposed storage condition) of the product and in the

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			authorized packaging material				
			a) as a package for sale and/or				
			b) After first opening and/or c) after the dilution/reconstitution				
			Stability Study of Drug Product, results of microbiological testing should be				
			included.				
MaV-13			Major change in the manufacturing process of the drug substance				
Conditions	to	be	1. The synthetic route is different with potential change in qualitative and/or				
fulfilled			quantitative impurity profile which would require further qualifications in				
			safety studies.				
			2. Manufacturing process of drug substance does not use any materials of				
			human/animal origin for which assessment is required of viral safety, unless				
			otherwise justified.				
			3. Physicochemical characteristics and other relevant properties of drug				
			substances remain unchanged.				
			4. Stability performance of drug substance remain unchanged.				
			5. If there are changes to the specification of drug substance.				
Documents	to	be	1. Relevant CTD section, or both the open and closed part of the Drug Master				
submitted			File with the Letter of Access or equivalent audit document/certification from				
			reference country which is deemed appropriate by the Drug Regulatory				
			Authority.				
			2. Comparative tabulated format of the approved and proposed processes with				
			changes highlighted.				
			3. For sterile drug substance, process validation report (where applicable).				
			4. A letter of declaration from marketing authorization holder stating that no				
			new impurities have been introduced at or above the accepted threshold for				
			qualification of impurities or that there is no increase in the levels of impurities,				
			which require further safety studies.				
			5. A letter of declaration from the marketing authorization holder stating that				
			the specifications of the drug substance have not changed.				
			6. Certificate of analysis and/or batch analysis data (in a comparative tabulated				
			format) for at least two pilot batches of the drug substance from the approved				
			and proposed process.				

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		7. A declaration from the marketing authorization holder that the relevant					
		stability studies of the drug product.					
		8. Certificate of analysis and/or batch analysis data (in a comparative tabulated					
		format) of drug product of at least two batches (pilot/production scale)					
		manufactured with the drug substance according to the approved and					
		proposed processes.					
MaV-14		Change or addition in the source of Active Pharmaceutical Ingredient / Drug					
		Substance or Half-Finished Products i.e. Pellets / Granules / Ready to Fill Bulk					
		etc.					
Conditions to	be	Specifications of drug substances remain unchanged.					
fulfilled							
Documents to	be	a. a. Real time and accelerated stability studies of DS / Half finished					
submitted		products (pellets / granules / ready to fill bulk) conducted by					
		manufacturer of DS / half finished product as per conditions of zone IV-A or zone IV-B on 3 commercial scale batches.					
		b. Comparative tabulated format of the approved and proposed drug					
		substance manufacture information (where applicable).					
		 Certificate of analysis and/or batch analysis data (in a comparative tabulated format) for at least two pilot batches of the drug substance 					
		from the approved and proposed manufacturing sites.					
		 Documents confirming that the proposed source has valid permission for manufacturing of DS / pellets / granules / ready to fill bulk by the 					
		regulatory authority of country of origin.					
		e. Copy of registration letter and last renewal status.					
		f. A letter of commitment from marketing authorization/ registration					
		holder to conduct long term and accelerated stability studies for the drug product manufactured with the drug substance from the proposed					
		manufacturing site, and report if any results fall outside shelf-life					
		specifications (with proposed action) or when requested.					
MaV-15		Change in Prescribing Information (PI) and labelling related to changes					
1110 1-13		in Indications, Contraindications, dosage etc.					
Conditions to	h.c						
Conditions to	be	These changes have already been approved by any reference authority					
fulfilled		and implemented by innovator drug product					
Documents to	be	a. Justification of proposed changes and difference between existing and					
submitted		proposed information in tabulated form. b. Reference of prescribing information approved by Reference					
		Regulatory Authorities and innovator product.					
		c. Copy of registration letter and last renewal status.					

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			d. Copy of approval from regulatory agency / authority from country of origin for innovator's drug product.					
			Copy of label outer pack in case of changes in indication/ dose/					
			administration etc.					
MaV-16			Change of specifications or method of analysis of finished drug product					
Conditions	to	be	a. Proposed specifications or method of analysis are not					
fulfilled			included in any pharmacopeia except in cases where proposed manufacturer specifications are more stringent than pharmacopeial specifications.					
			 For change in specifications, method of analysis will remain same or with minor change. 					
			c. The change is not necessitated by failure to meet					
			specifications resulting from unexpected events arising during					
			manufacture, or because of stability concerns.					
			d. There is no legal case / proceeding is pending at any forum /					
			court of law concerning with the proposed change					
Documents	to	be	a. Justification for the proposed change,					
submitted			 b. Comparative table of current and proposed specifications or method of analysis. 					
			c. Validated method of analysis in case of change of method of analysis.					
			d. Copy of registration letter and last renewal status.					
			e. Certificate of analysis of at least one batch and comparative summary of results, in tabular format, for one batch using current and proposed					
			procedures.					
			f. Undertaking that:					
			 No case is pending at any forum / court of law regarding this product. 					
			ii. In case of any quality complaint/ OOS result observed by the marketing authorization holder as a result of this change, the same will be reported to MFDA and all the stock will be recalled from the market immediately.					

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Any variation application shall be submitted through the online portal by submitting the Variation Applications of a Pharmaceutical Product as in Annex VIII.

If the application fulfills the prescribed criteria, Medicine Therapeutic Goods Division will process the case for consideration

13.3 Exclusion criteria for Post Registration Variations

- **13.3.1** The following cases of variations / changes shall not be considered as post-registration variations and require new product registration.
- 13.3.2 Change of Active Pharmaceutical Ingredient / Drug Substance
 - a. Change of the API to a different API including change in the salt or isomer form of API.
 - b. Inclusion or exclusion of an API to a multicomponent product.
 - c. Change in the strength of one or more APIs.
- **13.3.3** Change of Pharmaceutical Form /Dosage Form
 - a. Change in release profile of drug product like change from an immediate-release product to a slow-or delayed release dosage form and vice versa.
 - b. Change from a liquid to a powder for reconstitution, or vice versa.
- **13.3.4** Change in the route of administration.
- **13.3.5** Additional volume of already registered injectable drug products.
- 13.4 Procedure for submission of Post Registration Variations.
- **13.4.1** Variation applications can be submitted through Dhrithi portal.
- **13.4.2** The relevant information once received is evaluated and submitted to the technical committee of the authority for final approval or rejection. Once approved, the information is updated, and the applicant is informed via email or through Dhirithi portal.
- **13.4.3** Depending on the type of variation application it will take 10 to 20 working days for approval of the variation.

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14 Details of Variations, Applicable conditions and Required Documents.

14.1 Applicable Conditions

14.1.1 For each variation, attempts have been made to identify conditions or circumstance that need to be fulfilled for submitting variation application. For all changes, it remains the responsibility of the applicant to provide all necessary documents to demonstrate that the change does not adversely affect the quality, safety and efficacy of the drug product.

14.2 Required Documents

14.2.1 The list of documents required to be submitted along with application is identified for each variation in this guideline. However, this list is not exhaustive and further documentation if required may be asked from the applicant by MFDA. Regardless of the documents specified, applicants shall ensure provision of all relevant information to support the applied variation. Alternative approaches to the principles and practices described in this document may be acceptable provided that such variations / changes / practices or proposed alternatives are being supported with adequate scientific justification.

14.3 Details of Minor Variation-Notification (MiV-N change)

Minor Variation-Notification (MiV-N)				
MiV-N 1	Change in the name of Active Pharmaceutical Ingredient / Drug Substance, while the drug substance remains the same molecule(s)			
Conditions to be	a. No change in Active Pharmaceutical Ingredient/Drug substance (s).			
fulfilled				
Documents to be	a. Proof of acceptance by WHO or copy of the latest version of			
submitted	International Non-proprietary Names (INN) list mentioning proposed name of API/DS.			
MiV-N 2	Change of the name and/or address (e.g., street name) of a manufacturer of the			
	drug substance (API)			
Conditions to be	a. The manufacturing site of the drug substance remains unchanged.			
fulfilled	b. No other changes, except for the change of the name and/or address			
Turrinea	of a manufacturer of the drug substance			
Documents to be	a. Updated information of the manufacturer of the drug substance.			
submitted	b. Official document/evidence when required			
MiV-N 3	Change of manufacturing company`s owner			

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Conditions to be	a. The manufacturing and batch release site of the drug product remains			
fulfilled	the same.			
Documents to be	b. No other changes, except for change in the owner of manufacturera. Declaration on the transfer of ownership.			
	b. Official letter about change of old owner to new owner.			
submitted	b. Official letter about change of old owner to flew owner.			
MiV-N 4	Change in any part of the (primary) packaging material not in contact with the finished product formulation such as color of flip-off caps, color code rings on ampoules, change of needle shield (different plastic used).			
Conditions to be	The change is not related to primary or secondary packaging material			
fulfilled	and shall not affects the delivery, use, safety or stability of the finished drug product			
Documents to	a. Reason / justification for proposed change.			
be submitted	 b. Amendment of the relevant section(s) of the dossier, including revised product labeling as appropriate. 			
MiV-N 5	Withdrawal/deletion of the alternative manufacturer(s) for drug substance			
_	and/or drug product			
Conditions to be fulfilled	An alternative manufacturer is registered			
Documents to be submitted	Reason for withdrawal/deletion			
MiV-N 6	Minor change in the manufacturing process of an immediate release solid oral dosage form, semi solid or oral solutions			
Conditions to be	a. The change includes following:			
fulfilled	i. Change from non-automated or non-mechanical equipment to automated			
	or mechanical equipment to move ingredients.			
	ii. Change to alternative equipment of the same design and operating			
	principles of the same or of a different capacity.			
	iii. Process changes including changes such as mixing times and operating			
	speeds within application/validation ranges.			
5	b.No change in qualitative and quantitative impurity profile or in physio- chemical properties.			
	c. The manufacturing principle for individual manufacturing steps remain unchanged, e.g., there are no changes in the processing intermediates and manufacturing solvent(s) used in the process.			

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	d.The proposed process must be controlled by relevant in-process controls
	used in the approved process and no changes (widening or deletion of
	limits) are required for these controls.
	e. The specifications of the finished product and/or process intermediates
	remain unchanged.
	f. The proposed process must lead to an identical product regarding all
	aspects of quality, safety and efficacy
Documents to be	a. Amendment of the relevant section(s) of the dossier, as appropriate,
submitted	including a direct comparison of the approved and proposed processes.
	b.Copy of approved drug product specifications.
	c. Certificate of analysis and/or batch analysis data (in a comparative
	tabulated format) on a minimum of one batch manufactured to both the
	approved and the proposed process.
	d.A declaration/ undertaking from registration holder that:
	i. Batch analysis data on the next two full production batches shall be made
	available upon request and reported by the marketing authorization
	holder if outside specification (with proposed action).
	ii. The relevant stability studies of the drug product shall be started and that
	the relevant stability studies shall be finalized; data shall be provided only
	if outside specification (with proposed action).
MiV-N 7	Change of release and shelf-life specifications of the drug product, and/or
	drug substance, and/or excipient, following the inclusion in the
	compendium / pharmacopeia
Conditions to be	Drug products / Drug substances / excipients which are now included in the
fulfilled	compendia / pharmacopeia and applicant intends to adopt these
	specifications
Documents to be	a. Tabulation of the current and revised release and shelf-life specifications of

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submitted	the drug product, with changes highlighted.					
	b. Revised release and shelf-life specifications.					
	c. Copy of the relevant monograph from the compendium					
MiV-N 8	Change of imprints, bossing or other markings on tablets or printing on					
	capsules including addition or change of ink used for product marking					
Conditions to be	a. New markings do not cause confusion with other tablets or capsules.					
fulfilled	b.Any ink proposed for use must be edible.					
	c. Release and shelf-life specifications of the drug product remain					
	unchanged except appearance.					
Documents to be	a. Details and specifications of the proposed new ink (where applicable)					
submitted	b.Detailed drawing or written description of the current and proposed					
	imprint/bossing/markings.					
	c. Revised draft of package inserts and labeling incorporating the proposed					
	variation (where applicable).					
	d.Release and shelf-life specifications of drug product with new product					
	description.					
MiV-N 9	Addition or replacement of measuring device for oral liquid dosage forms					
	etc					
Conditions to be	a. Size and accuracy of the proposed measuring device must be compatible					
fulfilled	with the approved posology.					
	b. The new device is compatible with the drug product					
Documents to be	a. Justification for the proposed change.					
submitted	b Deviced duests of the medians income and labeling incomenting the					
	b. Revised draft of the package inserts and labeling incorporating the					
	proposed variations (where applicable).					
	c. Description of the device (where applicable)					
	d. Composition of device material and the material should be of					

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	pharmacopeial / pharmaceutical grade.
e.	Justification that size and accuracy of the device are adequate for the posology as approved in the product labeling.

14.4 Details of Min	nor Variation- Prior Approval (MiV-PA change)			
MiV-PA 1	Change of the name/title or address (e.g., street name/number) of the manufacturer of drug product			
Conditions to be	a. The manufacturing site remains the same.			
fulfilled	b. There is no legal case / proceeding is pending at any forum / court of law concerning with the proposed change			
	c. No other changes, except for the change of the name and/or address of			
	a manufacturer of the drug product.			
	d. Ownership of the company is unchanged.			
Documents to be submitted	 For imported drug products, official letter from related NRA or municipality (for address only) or original legalized CoPP as per WHO format for new manufacturer's name, or original legalized GMP 			
	certificate of new manufacturing site with free sale certificate from regulatory body of country of origin or any legalized document of concerned regulatory authority confirming the change of name of Manufacturer without change in manufacturing site			
	b. For local manufactured drug products, MFDA's approval letter for proposed variation.			
	c. Copy of registration letter and last renewal status			
	d. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable).			
MiV-PA 2	Change in the name / title and address of Registration Holder or MAH in			
	exporting country (for finished imported products)			
Conditions to be fulfilled	a. Registration holder / MAH and manufacturer should be separate entities.			
Tullilled	b. The change in address refers to only documentary change in address and the manufacturing site remains the same.			
	c. The name change refers to the renaming of a company or organization.			
	d. The change shall not include transfer of marketing authorization to another company.			
	e. There is no legal case / proceeding is pending at any forum / court of law concerning with the proposed change.			
	f. No other changes, except for the change of the name / Title of Registration Holder or MAH of the drug product.			

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Documents to be submitted MiV-PA 3	 a. Original legalized CoPP as per WHO format for new registration holder / MAH and/or address, or free sale certificate from regulatory body of country of origin or any legalized document of concerned regulatory authority confirming the change of name of Marketing Authorization Holder without change in manufacturing site. b. Evidence of the contract between registration / MA holder and manufacturer (with changed / new name), as the manufacturer and product license/registration holder are different entities. c. Copy of registration letter and last renewal status d. Revised notarized sole agency agreement with new registration holder. e. Revised draft of the package inserts and labeling incorporating the proposed variation (where applicable). f. An undertaking that the formulation, API source and Specifications, manufacturing process, release and shelf-life specifications have not changed. Change of name and address of importer 				
Conditions to be	a. The manufacturer including batch release site of drug product				
	remains the same.				
fulfilled	remains the same.				
Documents to be	a. MFDA's Drug import License with new address.				
submitted	b. Revised drafts of the package insert and labeling incorporating the				
Judillitted	proposed variation (where applicable).				
	c. Copy of registration letter and last renewal status				
	d. An undertaking that the firm that no case is pending at any forum / court of				
	law regarding previous name and/or address				
NA:V DA A	Change in Common of Books Change to sixting () aboltion (Bation) before ation				
MiV-PA 4	Changes in Summary of Product Characteristics/Labelling/Patient Information Leaflet				
Conditions to be	a. The changes shall be in accordance of the innovator drug products as				
fulfilled	approved by any of the Reference Authority				
Documents to be	a. Previously approved product labelling (SmPC, PIL etc) if any.				
submitted	b. Tabulated comparison in existing and proposed SmPC/PIL highlighting the				
	changes made.				
	c. Copy of registration letter and last renewal status				
	d. Copy of approved SmPC/PIL from any of the Reference Authority.				
	e. Latest version of SmPC/PIL of the Innovator product approved from any of the Reference Authority.				
MiV-PA 5	Change of drug product name				
Conditions to be	a. There shall be no change to the product specifications including formulation,				
fulfilled	release, and shelf-life specifications, manufacturing process etc except				

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	I	
		change of product name.
	L	No litigation shall be pending at any forward / sourt of law conservation with
	b.	No litigation shall be pending at any forum / court of law concerning with
		the proposed change.
	c.	The registration / MA holder will check the suitability of proposed names
		to ensure that no resemblance or phonetic matching with already
		registered products as per LASA so that the proposed names should not
		be liable to cause confusion in print, handwriting or speech with the
		(Proprietary / brand) name of another registered product.
	d.	The proposed name shall not suggest greater safety or efficacy than
		supported by clinical data, convey misleading therapeutic use or imply
		superiority over another similar product or show the presence of
		substance(s) present or not present in the product.
		substance(s) present of not present in the product.
Documents to be	a.	Justification for the proposed change
submitted	b.	Information regarding previous approvals of change of brand name
Sasimetea		since registration of drug product.
	C.	Details (batch number, date of manufacture, quantity and stock position) regarding last batch manufactured / imported.
	d.	Copy of registration letter and last renewal status
	e.	An undertaking that the proposed names do not resemble with already
		registered brands and in case of resemblance /similarity with already
		registered drug, the applicant will be liable to change immediately.
		Moreover, no case is pending at any forum / court of law regarding this
	f.	matter. Line extension Legalized CoPP or FSC in case of imported drug products.
	g.	Revised draft package insert and labeling incorporating the
	8.	proposed variation.
	h.	Official letter from product owner or marketing authorization holder
		authorizing the change of product name and committing to inform users
		of the relevant changes (where applicable).
MiV-PA 6	Change	e of importer/ MA holder in Maldives
Conditions to be	a.	The manufacturing site remains unchanged.
fulfilled	b.	No litigation shall be pending at any forum / court of law concerning with
		the proposed change.
		Present importer has not imported registered drug product for more than
	C.	Present importer has not imported registered drug product for more than 3 years.
		5 years.

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Documents to be	a. Notarized authority letter/sole agent letter (original) from marketing
submitted	authorization holder/ manufacturer (if both are separate) in name of new
	importer.
	b. Copy of registration letter and last renewal status
	 Revised drafts of the package insert and labeling incorporating the proposed variation.
	d. Legalized Certificate of Pharmaceutical Product (CoPP) or other relevant
	documents as defined by MFDA for new registration
MiV-PA 7	Change in Market Authorization Holder (MAH) in exporting country
Conditions to be	a. The manufacturing site remains unchanged.
fulfilled	
	b. The registration holder in Maldives shall remain the same.
	And the last of the state of th
	c. Undertaking from new MA holder in exporting country that no
	litigation is pending at any forum / court of law concerning with the
	proposed change
Documents to be	a. Notarized authority letter/sole agent letter (original) from new
submitted	Market Authorization Holder (MAH) abroad.
Subillitteu	ivial ket Authorization Holder (IviAH) abroau.
	b. Approval of new MAH from regulatory body of exporting country
	or Legalized Certificate of Pharmaceutical Product (CoPP) mentioning
	new Market Authorization Holder (MAH).
	c. Copy of registration letter and last renewal status
MiV-PA 8	Addition or replacement of alternative site for primary packaging (direct
	contact with drug product) for non-sterile product
Conditions to be	a. No other changes except for the addition or replacement of alternative
fulfilled	site for primary packaging (direct contact with drug product)
Documents to be	a. Justification for the proposed change.
submitted	b. For imported products, proof that the proposed site is legally authorized
	for the packaging activity of the drug product concerned such as CoPP (legalized) which covers GMP certification.
	C. For locally manufactured drug products, MFDA's regulatory approval for
	the proposed variation.
	d. Copy of registration letter and last renewal status.
	e. Validation scheme and/or report of the manufacturing process to the
	proposed change of alternative site for primary packaging (where
	applicable).

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	f. Revised drafts of the package insert and labeling incorporating the
	proposed variation (where applicable).
	g. Holding time studies testing of bulk pack during storage and
	transportation between the bulk production site to primary packager
	(where applicable).
	h. A letter of commitment from marketing authorization holder to conduct
	long term and accelerated stability studies for the first three batches of
	drug product packed at the proposed site, and report if any results fall
	outside shelf-life specifications (with proposed action) or when requested.
MiV-PA 9	Change of batch size of drug product
Conditions to be	a. The change does not affect consistency of production.
fulfilled	b. The product formulation remains unchanged.
Turrinea	c. Shelf-life specifications of drug product remain unchanged.
	d. This is applicable to change of batch size up to 10-fold compared to
	the approved batch size.
	e. The manufacturing process shall remain unchanged
Documents to be	a. Justification for the proposed change
submitted	b. Comparative tabulated format of approved and proposed batch size and
Jabiilitteu	batch manufacturing formula.
	c. Validation scheme and/or report of the manufacturing process of the
	proposed batch size.
	d. Copy of registration letter and last renewal status.
	e. Specifications of the drug product
	f. Revised section of registration application form (where applicable).
	g. Release and shelf-life specifications of the drug product.
	h. Certificate of analysis and/or batch analysis data (in a comparative
	tabulated format) of drug product of at least two production batches
	manufactured according to approved and proposed batch sizes.
	i. For oral solid dosage forms, comparative dissolution profile for at least
_	one production batch (where applicable).
	j. Appropriate real time and accelerated stability data to support proposed
	variation.
MiV-PA 10	Quantitative change in coating of tablets and/or size of capsule shell
Conditions to be	a. The dissolution profile of the proposed product is comparable to that of
fulfilled	the approved product.
Tullilleu	b. Specifications of the drug product remain unchanged except for the
	weight and/or size (where applicable).
Documents to be	a. Justification for the proposed change.
submitted	h Commonative tobulated formers of annual and annual an
	b. Comparative tabulated format of approved and proposed product and
	batch manufacturing formula.
	c. Comparative dissolution profile data of at least one production batch of the
	drug product manufactured in the approved and proposed composition
	for oral solid dosage forms.

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	 d. Revised draft of product label incorporating the proposed change (where applicable).
	e. For modified release oral products, stability data of the drug product and to report if any results fall outside shelf-life specifications (with proposed action)
	f. Copy of registration letter and last renewal status.
	g. Specifications of drug product.
	h. A declaration/ undertaking that:
	i. The change does not interfere with the drug product specifications test
	method.
	ii. The relevant stability studies of the drug product have been started and
	shall be reported if any results fall outside specifications (with proposed
	action).
MiV-PA 11	Change of dimensions and/or shape of tablets, capsules, suppositories
	or Pessaries
Candidiana ta ba	
Conditions to be	There will be no qualitative or quantitative change in API.
fulfilled	
Documents to be	a. Justification for the proposed change.
submitted	b. Revised drafts of the package insert and labeling incorporating the
	proposed variation (where applicable).
	c. Copy of registration letter and last renewal status.
	d. Detailed drawing or written description of the approved and proposed appearance.
	e. Comparative dissolution profile data of at least one production batch of
	the drug product manufactured in the approved and proposed
	dimensions/shape for oral solid dosage forms.
	f. For scored tablets, data on test of uniformity of the subdivided parts of
	tablets at release as conformed to compendia requirement.
	g. Specifications of the drug product with proposed dimension and/or shape.
MiV-PA 12	Change of secondary packaging materials
Conditions to be	The proposed packaging material must be at least equivalent to the
fulfilled	approved material in respect of its relevant properties
Documents to be	a. Justification for the proposed change.
1	

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submitted	b. Tabulated differences between existing and proposed information.			
	c. Copy of registration letter and last renewal status			
	d. An undertaking that:			
	 i. The proposed color scheme / label has no resemble with already registered Products. In case of resemblance, new label will be changed immediately. ii. Proposed change has not impact on the shelf life of the product. iii. No case is pending at any forum / court of law regarding this matter. 			
MiV-PA 13	Change in the design or color scheme of packaging material			
Conditions to be	a. The proposed packaging design/color scheme must not resemble to			
fulfilled	already registered product.			
	b. Packaging material shall remain same			
Documents to be	a. Justification for the proposed change.			
submitted	b. Tabulated comparison of differences between existing and proposed			
	design. c. Regulatory approval of change from country of export in case of imported			
	drug.			
	d. Copy of registration letter and last renewal status			
	e. An undertaking/ declaration that:i. No case is pending at any forum / court of law regarding this matter.			
	ii. All information related to the product like dosage, administration,			
	indication and direction for use etc. on the label are in line with the			
	registration / marketing authorization.			
	iii. The proposed label complies all provisions of relevant rules and regulations.			
MiV-PA 14	Change of the coloring agent /capsule shell color of the product			
Conditions to be	a. Same functional characteristics / specifications including no change in			
fulfilled	dissolution profile for solid oral dosage forms.			
	b. The proposed coloring agents /capsule shell are of pharmaceutical grade.c. The specifications of the drug product remain unchanged, except for			
	the update of product description with respect to			
	appearance/odor/taste as a consequence of the change (where			
Documents to be	applicable). a. Justification for the proposed change.			
	b. Revised drafts of the package insert and labeling incorporating the			
submitted	proposed variation (where applicable).			
	c. Revised product formulation and batch manufacturing formula.			
	d. Copy of registration letter and last renewal status.			

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	e. Tabulated comparison of qualitative and quantitative information of the approved and proposed coloring agent /capsule shell color.				
	f. For proposed excipients made of ruminant's source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform Encephalopathy (BSE)-free certificate issued from relevant authority of the issuing country and/or documentary evidence from the supplier (where applicable).				
	g. Revised specifications of the drug product.				
	h. Certificate of Analysis of proposed coloring agent /capsule shell (where applicable).				
	i. A declaration/ undertaking that:				
	i. The proposed coloring agent/capsule shell color does not interfere with				
	the drug product specifications test method.				
	ii. A letter of commitment from marketing authorization holder to inform				
	users of the relevant change (where applicable).				
	iii. Stability study has been started and report if any results fall outside drug				
	product specifications (with proposed action).				
MiV-PA 15	Addition/ change of flavoring agent of the product e.g oral liquid/ dry				
	powder suspension/sachet				
Conditions to be	a. Same functional characteristics, no change in dissolution profile for solid				
fulfilled	oral dosage forms.				
	oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use.				
	oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for				
	oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use.				
	oral dosage forms.b. The proposed flavoring agents must be of pharmaceutical use.c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a				
fulfilled Documents to be	 oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). 				
fulfilled	 oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). 				
fulfilled Documents to be	 oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. 				
fulfilled Documents to be	 oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the 				
fulfilled Documents to be	 oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. 				
fulfilled Documents to be	 oral dosage forms. b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status 				
fulfilled Documents to be	 b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status f. For proposed excipients made of ruminant's source, Transmitting Animal 				
fulfilled Documents to be	 b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status f. For proposed excipients made of ruminant's source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform 				
fulfilled Documents to be	 b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status f. For proposed excipients made of ruminant's source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform Encephalopathy (BSE)-free certificate issued from relevant authority of 				
fulfilled Documents to be	 b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status f. For proposed excipients made of ruminant's source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform 				
fulfilled Documents to be	 b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status f. For proposed excipients made of ruminant's source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform Encephalopathy (BSE)-free certificate issued from relevant authority of the issuing country and/or documentary evidence from the supplier 				
fulfilled Documents to be	 b. The proposed flavoring agents must be of pharmaceutical use. c. The specifications of the drug product remain unchanged, except for the update of product description with respect to flavor/taste as a consequence of the change (where applicable). a. Justification for the proposed change. b. Revised drafts of the package insert and labeling incorporating the proposed variation (where applicable). c. Revised product formulation and batch manufacturing formula. d. Tabulated comparison of qualitative and quantitative information of the approved and proposed flavoring agent. e. Copy of registration letter and last renewal status f. For proposed excipients made of ruminant's source, Transmitting Animal Spongiform Encephalopathy (TSE)-free certificate or Bovine Spongiform Encephalopathy (BSE)-free certificate issued from relevant authority of the issuing country and/or documentary evidence from the supplier (where applicable). 				

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	i. The proposed flavoring agent does not interfere with the drug product
	specifications test method.
	specifications test metriou.
	ii Aletter of commitment from marketing authorization holder to inform
	ii. A letter of commitment from marketing authorization holder to inform
	users of the relevant change (where applicable).
	iii. Stability study has been started and report if any results fall outside drug
	product specifications (with proposed action).
MiV-PA 16	Change of shape or dimension of container closure system
Conditions to be	a. The change only concerns the same packaging type and material.
	b. The proposed pack size is consistent with the dosage regimen and duration
fulfilled	of use as approved in the package insert.
	c. Change in the dimension of the primary packaging (where applicable).
	d. Specifications of the drug product remain unchanged.
Documents to be	
Documents to be	a. Justification for the proposed changes in container closure system.
submitted	b. Information on the proposed container-closure system (e.g. description,
	materials of construction, and specifications).
	c. Copy of registration letter and last renewal status.
	d. Revised drafts of the package insert and labeling incorporating the
	proposed variation (where applicable).
	e. Re-validation studies for manufacturing process and sterilization process
	performed in case of sterile products which are terminally sterilized.
	f. Revised Sections of registration applications (where applicable).
	g. A declaration/undertaking that the relevant stability studies of the drug
	product have been started and that the relevant stability studies shall be
	finalized; data shall be provided only if outside specification (with
	proposed action).
	h. The proposed label complies all provisions of relevant rules and
	regulations.
	i. An undertaking that:
	i. Other specifications of the product would remain the same.
	ii. There is no change in the qualitative & quantitative composition of the
	product and manufacturer will conduct product development,
	accelerated and real time stability studies, validation of manufacturing
	process and method of analysis before sale of drug.
	iii. In the case of changes to the thickness of a packaging component or for
	sterile FPPs: stability data (as per conditions of zone IV-A), where
	applicable, results of photo-stability studies will be conducted on 03 lab
	scale batches or developmental scale batches.
	iv. In the case of a change in the headspace or a change in the
	surface/volume ratio for non-sterile FPPs, a commitment for the above
	studies to ensure appropriate delivery.
	v. In case of any quality complaint/ OOS result observed by the marketing

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	authorization holder as a result of this change, the same will be reported to MFDA and all the stock will be recalled from the market immediately.					
MiV-PA 17	Replacement of the company or party responsible for batch release					
Conditions to be	a. Only applicable for batch release.					
fulfilled	b. Method transfer from the currently approved to the proposed site or test					
	laboratory has been successfully completed.					
	c. The manufacturer of the drug product remains the same					
Documents to be	a. Justification for the proposed change.					
submitted	b. Official letter from manufacturing company that the batch release					
	responsible is changed.					
	c. Proof that the proposed site is appropriately authorized (accredited by the					
	NRA) to be responsible for batch release, such as a valid GMP certificate or					
	CoPP which covers the GMP certification for imported products and MFDA approval for locally manufactured products.					
	d. Copy of registration letter and last renewal status.					
	e. Document for method transfer to the proposed site along with validation					
	of method and mock testing on already manufactured batches,					
	f. Revised drafts of the package insert and labeling incorporating the					
	proposed variation (where applicable).					
MiV-PA 18	Change of in-process controls applied during the manufacture of the drug					
	product (including tightening and addition of new in- process test)					
Conditions to be	a. Release and shelf-life specifications of drug product remain unchanged.					
fulfilled						
	b.The change does not result from unexpected events arising during					
	manufacture (e.g., new unqualified impurity, change in total impurity					
	limit)					
Documents to be	a. Justification for the proposed change.					
submitted	b. Copy of registration letter and last renewal status.					
submitted	c. A description of the analytical methodology and summary of validation					
	data must be provided for all new analytical methods (where applicable).					
	d. Revised in-process specifications together with justification and relevant					
	process validation data.					
	e. Comparative tabulated format change of the in-process controls.					
	f. Certificate of analysis and comparative batch analysis data of drug product					
	of at least two production/pilot batches					
MiV-PA 19						
	Change in the test procedure of the drug product (including replacement					
	Change in the test procedure of the drug product (including replacement or addition of a test procedure)					
Conditions to be						
	or addition of a test procedure)					
Conditions to be fulfilled	or addition of a test procedure) a. Drug product specifications are not adversely affected unless the					

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	a The decree to the latest health and the first of the control of				
	c. The change should not be the result of unexpected events arising during				
Documents to be	manufacture or because of stability concerns				
	a. Justification for the proposed change.				
submitted	b. Copy of registration letter and last renewal status.				
	c. Description of the analytical methodology.				
	d. Appropriate verification/validation data and comparative analytical results between the currently approved and proposed test.				
	e. Comparative tabulated format of the currently approved and proposed				
	release and shelf-life specifications of the drug product.				
	f. Certificate of analysis and batch analysis data of the finished product of				
	two production batches when made available				
MiV-PA 20	Change of release and shelf-life specifications of the drug product				
	a) Specification limits are tightened				
	b) Addition of new test parameter and limits				
Conditions to be	a. Applicable to non-compendial / non-pharmacopeial methods.				
fulfilled	b. The change should not be the result of unexpected events arising during				
Tannica	manufacture or because of stability concerns.				
	c. The test methods remain the same or changes in the test methods are				
	minor.				
	d. If there are changes to the test procedure, then relevant conditions and				
	documents will be required.				
Documents to be	a. Specification limits are tightened				
submitted	 Tabulated comparison of the current and revised release and shelf-life specifications of the drug product with changes highlighted. 				
	ii. Certificate of analysis and comparative batch analysis data of the drug				
	product for all tests in the new specification of at least two batches.				
	b. Addition of new test parameter and limits				
	i. Justification for the proposed change				
	ii. Description of new test parameter and limits along with validated				
	method and summary of analytical validation data for non-compendial				
	method.				
	iii. Stability data and report if any results fall outside shelf-life				
	specifications (with proposed action) (where applicable).				
MiV-PA 21	Standardization of formulation in accordance with the Innovator's Drug Product/ Reference Authorities and Pharmacopeias				
Conditions to be	Existing formulation shall remain the same				
fulfilled					
Documents to be	a. Justification for the proposed change.				
submitted	b. Copy of registration letter and last renewal status.				
-	c. Document in support of proposed correction/evidence of approval status				
	by Reference Authorities/ innovator drug product and/ or				
	Pharmacopeias.				
	d. Undertaking that the provided information/ documents is true/ correct.				
MiV-PA 22	Reduction or removal of overage				

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Conditions to be	a. Change of previously approved manufacturing overages of drug						
	substance only.						
fulfilled	b. Release and shelf-life specification of drug product remain unchanged.						
Documents to be	a. Justification for the proposed change.						
au busittad	b. Tabulated comparison of currently approved and proposed batch						
submitted	manufacturing formula.						
	c. Certificate of analysis of 2 batches of finished drug product.						
	d. Stability data and report, if any result fall outside shelf-life specification.						
MiV-PA 23	Change in source of empty hard capsule						
Conditions to be	a. No change in formulation and manufacturing process of drug product.						
fulfilled	b. No applicable to change from hard capsule to soft gel.						
rannica	c. Formulation including excipients will remain unchanged						
	d. Release and shelf-life specification of drug product remain unchanged.						
Documents to be	a. Comparative dissolution profile data of one batch representative of						
submitted	pilot/production batch of the drug product using hard capsule between						
	two sources (where applicable).						
	b. Certificate of analysis of empty hard capsule of the proposed source.						
	c. Specification and composition of empty hard capsule of new source						
	(including origin i.e. synthetic, vegetable or animal source).						
MiV-PA 24	d. Stability data and report, if any result fall outside shelf-life specification. Addition or removal of score / break line on tablet						
Conditions to be	a. Innovator drug product has same score / break line on tablet.						
fulfilled	b. Release and shelf-life specifications of the drug product rema						
December 1	unchanged except appearance.						
Documents to be	a. Justification of the proposed change (including change in dosage						
submitted	regimen)						
	b. Details and specifications of the proposed change.						
	c. Detailed drawing or written description of the current and proposed						
	score / break line on tablet.						
	Score / Break line on tablet.						
	d. Revised draft of package inserts and labeling incorporating the proposed						
	variation (where applicable).						
	e. Release and shelf-life specifications of drug product with new product						
	description.						
	f. Certificate of analysis of two production /pilot scale batches.						

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14.5 General consideration for variations.

- **14.5.1** Variations that are not listed in the table will be reviewed and processed by the Authority using a risk-based approach and under the recommended guidance of WHO "Guidance on variations to a pregualified product dossier".
- **14.5.2** The list of documentation in the table is meant for guidance purposes and it shall be noted that the Authority reserves the right to request further information not explicitly described in the guideline.
- **14.5.3** For all changes it remains the responsibility of the applicant to provide all necessary documents to demonstrate that the change does not have a negative impact on the safety, efficacy or quality of the finished pharmaceutical product.

14.6 Variations which will be not acceptable

a. Incomplete or Insufficient Data

- > Reason: Lack of adequate scientific evidence to support the proposed change.
- > Examples:
 - Missing validation data for new manufacturing processes.
 - Absence of stability studies for a new formulation.
 - Inadequate clinical data for an expanded indication.

b. Non-Compliance with Regulatory Guidelines

- Reason: Proposed variations that violate regulatory standards or fall outside the allowed scope.
- **Examples**:
 - Changes submitted in an incorrect variation category (e.g., Type IA instead of Type II in the EU).
 - Non-compliance with quality, safety, or efficacy guidelines.
 - Missing mandatory documents like GMP certifications.

c. Significant Alteration of Product Identity

- Reason: Changes that fundamentally alter the pharmaceutical's approved identity.
- > Examples:
 - Changing the active ingredient or its salt/ester form.

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- Switching the product's intended route of administration (e.g., from oral to injectable).
- Modifications that lead to a completely different dosage form or strength not covered by the initial approval.

d. Risk to Safety or Efficacy

- > Reason: Variations introducing risks that cannot be mitigated through testing or data.
- > Examples:
 - Substitution of critical excipients leading to altered pharmacokinetics or reduced efficacy.
 - Changes in the manufacturing process introducing impurities or affecting bioavailability.
 - Introduction of a new indication with insufficient clinical evidence.

e. Unapproved or Poorly Documented Manufacturing Facilities

- > Reason: Use of facilities that do not meet GMP requirements or lack regulatory approval.
- > Examples:
 - Changing of manufacturing site
 - Use of unqualified third-party contract manufacturers.

f. Variations Impacting Intellectual Property (IP) or Exclusivity

- Reason: Changes violating IP laws or infringing on patent rights.
- > Examples:
 - Proposing a variation that copies a patented formulation or process.
 - Unauthorized use of proprietary excipients or technologies.

g. Variations Conflicting with Regulatory Agreements

- Reason: Deviations from commitments made during the original approval process.
- > Examples:
 - Alterations to post-marketing commitments (e.g., agreed pharmacovigilance plans).
 - Failing to maintain previously approved manufacturing standards.

h. Lack of Scientific or Regulatory Justification

Reason: Failure to provide a clear rationale for the proposed change.

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Examples:

- Introducing changes without demonstrating their necessity or benefit.
- Modifications that are cosmetic or unnecessary (e.g., minor labeling tweaks not improving readability).

i. Variations Impacting Product Supply Chain

- Reason: Changes disrupting the integrity or traceability of the product.
- > Examples:
 - Switching primary packaging to materials not validated for stability.
 - Introducing a distribution channel that violates cold chain requirements.

j. Timing or Scope of Variation Not Permitted

- Reason: Regulatory rules or agreements may restrict certain changes during specific periods or for specific products.
- > Examples:
 - Proposing changes during ongoing product recalls or safety investigations.
 - Submitting variations for products under temporary approval conditions (e.g., emergency use authorizations).

k. Regulatory Considerations:

- > Pre-Consultation: Consult with regulatory authorities before submitting complex or non-standard variations.
- > Impact Analysis: Conduct thorough risk assessments to understand the variation's impact on safety, efficacy, and quality.
- Alternatives: If a variation is likely to be rejected, consider alternative approaches, such as filing a new marketing authorization.

15 Special Permissions and Exemptions

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- 15.1 MFDA grants special permission for import of unregistered drugs to hospital to ensure access of drugs for the treatment of patients. These exemptions are granted in the form of Preauthorization in reference to the Guideline for Pre-Authorization Approval of Medicines (MTG-RE-PA/GLN-TE 010) and are exclusively granted to hospitals for use of medicines in their own facilities only.
- 15.2 These permissions are subjected to following points:
 - a. For hospital use medicines that are essential and low in volume and quantity due to difficulties in acquiring the required documentation for registration, exemption approvals will be given to hospitals to import that specific product.
 - b. The approvals will be given under a set of conditions by means of a signed agreement between MFDA and the importing hospital.

16 Clinician's Request for Approval of New Medicine

16.1 This form is introduced to the doctors to request to add new chemicals, new dosage form or new strengths, to the Approved Drug List upon the requirement of the patients, and to make sure to maintain the uninterrupted availability of the medicines.

16.2 Required Documents

- a. A completed "Clinician's Request for Approval of New Medicine" signed by the requesting doctor and approved by head of the applicants' organization/Health facility.
- b. Additional information and the picture shots of the product.
- c. Research Paper of the product

16.3 Process of clinician form

- **16.3.1** If the application is for a new chemical, new strength and new dosage form, and the form is filled in completely and provides the required document the application will be accepted.
- **16.3.2** The evaluation will be carried out within the next 45 (Forty-five) working days.
- **16.3.3** Once the product has been approved by NPB the product will be added to ADL and inform the client within 7 working days.

16.4 Rejection of Form

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16.4.1 If the application is incomplete or if the application is form for an existing medicine in the ADL, the application will be rejected and informed to the applicant during working 7 days via email.

17 Post Marketing Surveillance

- **17.1** Once the product is registered, imported and introduced to the market, the product shall be on surveillance as to ensure that the same product registered is in the market and if the product is safe, of good quality and efficacy in accordance with the applied documents for registration.
- 17.2 Once registered, as part of the post market surveillance, samples will be collected from the market and tested from National Health Laboratory NHL as well as the assigned laboratory from abroad and these results will be published.

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18 References

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Letter of Appointment



Maldives Food and Drug Authority

Ministry of Health and Family

Male`, Republic of Maldives

Statement by the Local Applicant for submission of Application

1) I have received and accepted the entire dossier from <u>Company name and address</u> for the product Name of the product, Brand/ Generic/ Dosage Form/ Strength.

This dossier includes all data in support of the original documents as per the format of MFDA.

- 2) I hereby agree that I have sole responsibility for the mentioned product including obtaining approvals for any subsequent product variation and maintenance of the product registration.
- 3) I declare that information submitted in this application is correct and complete. I authorize the Maldives Food and Drug Authority to obtain information from any institution previously or currently associated with my company. If any information supplied by me is considered to be false, incomplete or misleading in any aspect, Maldives Food and Drug Authority has the right to take action as it believes necessary including the disclosure of the information to any person or body the Maldives Food and Drug Authority considers has a legitimate interest in receiving it and I consent to such disclosure. I understand the Maldives Food and Drug Authority reserves the right to vary or revoke any decision made on the basis of untrue, incomplete or misleading information. Moreover, I will co-

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operate with any person representing the Maldives Food and Drug Authority, by providing additional information or making the manufacturing premises available for inspection as required.

4) I also acknowledge the responsibility in the event of pharmacovigilance issues or quality defects

associated with the product that may occur after the registration.

5) The information provided to the Maldives Food and Drug Authority contain confidential

information that can hinder our business and hence this information shall be kept confidential and

shall not be disclosed to any third party without our consent.

6) I shall take the responsibility for updating any information relevant to the product/application and

will take the initiative to inform MFDA in a timely manner any change in product information during

the course of evaluation, and after product registration, especially if the information pertains to

rejection/withdrawal and will provide, additional data on product efficacy and safety or current Good

Manufacturing Practice (cGMP) compliance of the manufacturers (and repackers, if applicable).

7) I will also supply relevant information in case where the manufacturing facility is sold, merged or

changed to another.

8) As the local agent for marketing the product, I shall take full responsibility for assuring the quality,

safety and efficacy of this product throughout the supply chain.

Applicant representative information (whom MFDA will contact);

Name:

Phone number

Email:

Signed:

Full Name:

Identity Card Number:

Full Address:

Status of the signatory:

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(To be signed by the managing director/president/CEO or an equivalent person who has overall responsibility for the company or organization)

Official company stamp:

Fax Number/Telephone Number:

E mail contact details:

Date:

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Reference countries list for Reliance

#	Country	Authority	Reliance Scope	Reliance Criteria
1	Australia	Therapeutic Goods Administration	Medicines and Vaccines	A stringent regulatory authority (SRA)
2	Austria	Austrian Agency for Health and Food Safety (AGES)	Medicines and Vaccines	A stringent regulatory authority (SRA)
3	Belgium	Federal Agency for Medicines and Health Products (FAMHP)	Medicines and Vaccines	A stringent regulatory authority (SRA)
4	Bulgaria	Bulgarian Drug Agency	Medicines and Vaccines	A stringent regulatory authority (SRA)
5	Canada	Health Canada	Medicines and Vaccines	A stringent regulatory authority (SRA)
6	Croatia	Agency for Medicinal Products and Medical Devices of Croatia (HALMED)	Medicines and Vaccines	A stringent regulatory authority (SRA)
7	Cyprus	Ministry of Health — Pharmaceutical Services	Medicines and Vaccines	A stringent regulatory authority (SRA)
8	Czech Republic	State Institute for Drug Control (SUKL)	Medicines and Vaccines	A stringent regulatory authority (SRA)
9	Denmark	Danish Medicines Agency	Medicines and Vaccines	A stringent regulatory authority (SRA)
10	Estonia	State Agency of Medicines (Ravimiamet)	Medicines and Vaccines	A stringent regulatory authority (SRA)
11	Finland	Finnish Medicines Agency (Fimea)	Medicines and Vaccines	A stringent regulatory authority (SRA)

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12	France	National Agency for the Safety of	Medicines and	A stringent regulatory
		Medicine and Health Products	Vaccines	authority (SRA)
		(ANSM)		
13	Ghana	Food And Drugs Authority (FDA)	Medicines and	Maturity Level 3, based
			Vaccines	on WHO GBT
				Assessment
14	Germany	Federal Institute for Drugs and	Medicines and	A stringent regulatory
		Medical Devices	Vaccines	authority (SRA)
15	Greece	National Organization for	Medicines and	A stringent regulatory
		Medicines	Vaccines	authority (SRA)
16	Hungary	National Institute of Pharmacy and	Medicines and	A stringent regulatory
10	Trungary	Nutrition (OGYEI)		
		Nutrition (OGYEI)	Vaccines	authority (SRA)
17	Iceland	Icelandic Medicines Agency	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
18	Indonesia	BADAN POM (Agency for Drug and	Medicines and	Maturity Level 3, based
		Food Control, or Indonesian FDA)	Vaccines	on WHO GBT
				Assessment
19	Ireland	Health Products Regulatory	Medicines and	A stringent regulatory
		Authority	Vaccines	authority (SRA)
20	Italy	Italian Medicines Agency (AIFA)	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
21	Japan	Ministry of Health, Labour and	Medicines and	A stringent regulatory
21	Japan	Welfare/Pharmaceuticals and		
			Vaccines	authority (SRA)
		Medical Devices Agency		
22	Latvia	State Agency of Medicines	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
23	Liechtenstein	Office of Health / Department of	Medicines and	A stringent regulatory
		Pharmaceuticals	Vaccines	authority (SRA)

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24	Lithuania	State Medicines Control Agency	Medicines and	A stringent regulatory
		(VVKT)	Vaccines	authority (SRA)
25	Luxembourg	Ministry of Health	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
26	Malta	Medicines Authority	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
27	Netherlands	Health and Youth Care Inspectorate	Medicines and	A stringent regulatory
		(IGZ)	Vaccines	authority (SRA)
28	Nigeria	National Agency for Food and Drug	Medicines and	Maturity Level 3, based
		administration and control	Vaccines	on WHO GBT
		(NAFDAC)		Assessment
29	Norway	Norwegian Medicines Agency	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
30	Poland	Chief Pharmaceutical Inspectorate	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
31	Portugal	National Authority of Medicines	Medicines and	A stringent regulatory
		and Health Products (Infarmed)	Vaccines	authority (SRA)
32	Republic of Korea	Ministry of Food and Drug Safety	Medicines and	Maturity Level 4, based
		(MFDS)intlpharm@korea.kr	Vaccines	on WHO GBT
				Assessment
33	Romania	National Agency for Medicines and	Medicines and	A stringent regulatory
		Medical Devices	Vaccines	authority (SRA)
34	Saudi Arabia	Saudi Food and Drug Authority	Medicines and	Maturity Level 4, based
		(SFDA)	Vaccines	on WHO GBT
				Assessment
35	Singapore	Health Sciences Authority (HSA)	Medicines and	Maturity Level 4, based
			Vaccines	on WHO GBT
				Assessment

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36	Slovakia	State Institute for Drug Control	Medicines and	A stringent regulatory
		(SIDC)	Vaccines	authority (SRA)
37	Slovenia	Agency for Medicinal Products and	Medicines and	A stringent regulatory
		Medical Devices (JAZMP)	Vaccines	authority (SRA)
38	Spain	Spanish Agency of Medicines and	Medicines and	A stringent regulatory
		Medical Devices (AEMPS)	Vaccines	authority (SRA)
39	Sweden	Medical Products Agency	Medicines and	A stringent regulatory
			Vaccines	authority (SRA)
40	Switzerland	Swiss Agency for Therapeutic	Medicines and	A stringent regulatory
		Products (Swissmedic)	Vaccines	authority (SRA)
41	Tanzania	Tanzania medicines and Medical	Medicines and	Maturity Level 3, based
		Devices Authority (TMDA)	Vaccines	on WHO GBT
				Assessment
42	Türkiye	Turkish medicines and Medical	Medicines and	Maturity Level 3, based
		Devices Authority (TITCK)	Vaccines	on WHO GBT
				Assessment
43	United Kingdom	Medicines and Healthcare products	Medicines and	A stringent regulatory
		Regulatory Agency (MHRA)	Vaccines	authority (SRA)
44	United States of	Food and Drug Administration	Medicines and	A stringent regulatory
	America		Vaccines	authority (SRA)
45	Zimbabwe	Medicines Control Authority of	Medicines and	Maturity Level 3, based
		Zimbabwe (MCAZ)	Vaccines	on WHO GBT
				Assessment
46	China	National Medicines Products	Vaccines only	Maturity Level 3, based
		Administration (NMPA)		on WHO GBT
				Assessment
47	India	Central Drugs Standard Control	Vaccines only	Maturity Level 3, based
		Organization (CDSCO)		on WHO GBT
				Assessment

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48	Thailand	Food and Drug Administration	Vaccines only	Maturity Level 3, based	
				on WHO GBT	
				Assessment	

- A stringent regulatory authority (SRA) is a national drug regulation authority which the World Health Organization (WHO)
 considers to apply stringent standards for quality, safety, and efficacy in its process of regulatory review of drugs and vaccines for
 marketing authorization
- Maturity level 3: Meaning "stable, well-functioning and integrated regulatory systems"
- Maturity Level 4: Meaning "regulatory systems operating at advanced level of performance and continuous improvement

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Product information required for Reliance

Min	Minimum information required for reliance for Market Authorization					
The	following information sho	ould be accessible from the	reference NRA (publicly, or s	hared by the NRA,		
e.g.	upon request): ingredient	(API)/drug substance				
	Evidence of approval from the reference organization with sufficient details (in one or several					
1.	documents, e.g. evidence of MA, public¹ evaluation reports or CoPP). This shall be submitted as					
	a separate attachm	ent.				
2.	Proprietary product	name /Brand Name:				
3.	International Nonp	roprietary Name (INN) of the	e active pharmaceutical /Gen	eric Name:		
4	Composition:					
	Component and	Function of the	Quantity per unit	%		
	quality standard	component in	(mg)			
		the formulation				
	Total					
5.	Strength:					
6.	Pharmaceutical form	m (Dosage form):				
7.	Indication or Use of	the product:				
8.	Storage conditions:					
9.	Shelf life of the pro	duct:				
10.	Attach Stability sum	nmary and conclusions (inclu	uding the storage statement a	and shelf-life)		
11.	Dispensing Categor	y:				
12.	Pack size / Volume:					
13.	WHO ATCC code:					

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	Date of approval by the reference organization: Date of approval with Link to verify
14.	
15.	Marketing authorization number from the reference organization (if applicable):
16.	Name(s) and complete address(es) (including specific unit/blocks) of the manufacturer(s) of the
	finished pharmaceutical product(s) [FPP(s)] or biological drug products(s) (DP(s)), including the
	final product release if different from the manufacturer:
17.	Attach Detailed description of the primary and secondary packing including pack size or volume
18.	Attach Product information for healthcare professionals (SMPC)
19.	Provide a valid GMP certificate or incorporate the information in this sheet as per the
	mentioned information below, if providing a certificate, it must be attached separately.
	cGMP certificate information, including:
	o certificate number
	o applicable standards
	o site name
	o site address
	o issue date and validity
	o Scope: product
	o Scope: Manufacturing operations
20.	Attach Public ² evaluation reports (for new medicines ³ only)
In ac	ddition, the following information is also required but would generally be accessed through the
appli	icant as it may be considered to be confidential:
21.	Attach Declaration of sameness of the product as that of the product registered and approved
	in the reference NRA
22.	Attach, as far as possible, the full unredacted assessment reports from the reference
	authority/institution
23.	Name and complete address (including specific unit/blocks) of the API/drug substance
	manufacturer(s):
24.	Attach detailed Description (visual appearance) of the finished pharmaceutical product
	(eg:color, texture etc of tablet, syrup etc)
25.	Attach Specifications for the finished product
26.	Attach Product information for patients including product artwork
27.	Cost and propose retail price in USD:

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1 Unredacted assessment reports are also evidence of approval but are generally not accessible publicly. The sharing of such document would usually require a Memorandum of Understanding and/or a Confidential Disclosure Agreement

2 Unredacted assessment reports would be ideal but they are generally not accessible publicly. The sharing of such document would usually require a Memorandum of Understanding and/or a Confidential Disclosure Agreement.

3 In the context of this document, new medicines/vaccines refer to either a substance, association of substances, pharmaceutical form, dosage, indication, or posology which has never been granted a marketing authorization in the country.

References:

Members can refer to the Template/ checklist for countries to conduct "Recognition" OR "Verification of the product sameness for MA/RI": Appendix 2, Verification for product submitted under the WHO collaborative procedure (page 259), https://apps.who.int/iris/bitstream/handle/10665/312316/9789241210287-eng.pdf?sequence=1&isAllowed=y#page=249&zoom=auto,-182,680

Model certificate of a Certificate of a pharmaceutical product: https://www.who.int/teams/regulation-prequalification/regulation-andsafety/regulatory-convergence-networks/certification-scheme/modelcertificate-of-a-pharmaceutical-product

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Excipient at risk for DEG and EG

These are the list of excipients added to oral liquid preparations which are at risk for DEG/EG contamination and Adulteration

- 1. Glycerin or Glycerol
- 2. Saccharide and Polysaccharide solutions
- 3. Sorbitol solutions.
 - Noncrystallizing sorbitol solution
 - Sorbitol sorbitan solution
 - Maltitol solution
 - Liquid products of hydrogenated starch hydrolysate
- 4. Propylene glycol
- 5. Ethoxylated excipients
 - Polyethylene glycol 200
 - Polyethylene glycol 300
 - Polyethylene glycol 400
 - Polyethylene glycol 600
 - Polyethylene glycol 1000
 - Polysorbate 20
 - Polysorbate 40
 - Polysorbate 60
 - Polysorbate 80
 - Polyethylene glycol monomethyl ether 350
 - Polyethylene glycol monomethyl ether 550
 - Polyoxyl 35 castor oil
 - Polyoxyl 15 hydroxystearate
 - Polyoxyl 20 cetostearyl ether
 - Polyoxyl 8 stearate; Octoxynol 9
 - Nonoxynol 9

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Checklist for Reliance Applications



Maldives Food and Drug Authority Ministry of Health

Male`

Republic of Maldives

Rec.No: MTG/RE-ES/Re-0005/2024 - 000

1. Registration Pathways		
Reliance Option 1.1.1. Reliance on MA (approved by reference NRAs)	5 years	
Reliance Option1.2.1 GMP verification (manufacturing site certified by PIC/S member NRA) Abridge	5 years	
Reliance Option 1.2.2 WHO prequalified products/manufacturers	5 years	

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Reliance Option 1.3.1 Collaboraive registration procedure (CRP-PQ)	5 years	
Reliance Option 1.3.2 Collaboraive registration procedure (CRP-SRA)	5 years	
Notification: These are for low-risk medicines like vitamins and vitamin preparation that are categorized as medicines.	5 years	

2. Registration type (to be filled by MFDA)

Date of Submission	
Date of Acceptance	
A) New Registration	
B) Re-registration	

3. Product Evaluation Summary

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	Client submitted information	Evaluation Remarks	Check verified	and
Α.	General Information for all application	ns		
Letter of appointment from the local agent				
Letter from manufacturer to MFDA				
Artwork, 360 angle picture of the product with package insert.				
B. Information required for Rel	iance: Option 1.1.1. Reliance on MA (approved by reference NRAs	5)	
1) Evidence of approval from the country of manufacture (A link should be provided to trace the evidence or registration or marketing authorization certificate.)				
2) Evidence of approval and marketing in one reference country as mentioned in Annex II. (A link should be provided to trace the evidence or registration or marketing authorization certificate.)				
3)Product information for reliance as in Annex III:				

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2.1 Evidence of approval from the reference organization with			
3.1 Evidence of approval from the reference organization with			
sufficient details (in one or several documents, e.g. evidence of MA			
public1 evaluation reports or CoPP). This shall be submitted as a			
separate attachment. (THIS IS SAME AS IN POINT 2 OF THIS CLAUSE			
3.2 Proprietary product name /Brand Name:			
3.3 International Nonproprietary Name (INN) of the active			
pharmaceutical /Generic Name:			
3.4 Compositions including all active and non-active ingredients:			
	Function of the component in the		.,
Component and quality standard		Quantity per unit (mg) %	%
Component and quality standard	formulation	Z [F (/-
component and quanty standard	formulation	Z	
Component and quality standard	formulation	Zaman J. Parama (v. G)	
component and quanty standard	formulation		
component and quanty standard	formulation		
Component and quanty standard	formulation		
Component and quanty standard	formulation		
Component and quanty standard	formulation		
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Component and quanty standard	formulation		

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	Tatal	
	Total	
2 F Strongth		
3.5 Strength:		
3.6 Pharmaceutical form (Dosage form):		
()		
3.7 Indication or Use of the product:		
3.8 Storage conditions:		
5.0 Storage conditions.		
3.9 Shelf life of the product:		
2.40 \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\		
3.10 WHO ATCC code:		
3.11 Stability summary and conclusions (including the storage		
statement and shelf-life) This to be attached		
3.12 Dispensing Category:		
3.12 Dispersing category.		
3.13 Pack size / Volume:		
3.14 Date of approval by the reference organization: Date of		
3.14 Date of approval by the reference organization. Date of		
approval with Link to verify		

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3.15 Marketing authorization number from the reference organization (if applicable):		
3.16 Name(s) and complete address(es) (including specific unit/blocks) of the manufacturer(s) of the finished pharmaceutical product(s) [FPP(s)] or biological drug products(s) (DP(s)), including		
the final product release if different from the manufacturer:		
3.17 Detailed description of the primary and secondary packing including pack size or volume to be attached		
3.18 Attach Product information for healthcare professionals (SMPC)		
3.19 Provide a valid GMP certificate or incorporate the information in this sheet as per the mentioned information below, if providing a		
certificate, it must be attached separately. cGMP certificates		
information, including:		
3.19.1 certificate number		
3.19.2 applicable standards		
3.19.3 site name		
3.19.4 site address		
3.19.5 issue date and validity		

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3.19.6 Scope: products		
3.19.7 Scope: Manufacturing operations		
3.19.8 Authorizating organization:		
3.20 Attach Public assessment reports (for new medicines only)		
3.21 Attach Declaration of sameness of the product as that of the		
product registered and approved in the reference NRA		
3.22Attach, as far as possible, the full unredacted assessment		
reports from the reference authority/institution or a link to verify		
this information.		
3.23 Name and complete address (including specific unit/blocks) of		
the API/drug substance manufacturer(s):		
3.24 Attach detailed Description (visual appearance) of the finished		
pharmaceutical product (eg:color , texture etc of tablet, syrup etc)		
3.25 Attach Specifications for the finished product		
3.26 Attach Product information for patients.		
3.27 Cost and propose retail price in USD:		
4) Stability study report covering applicable climatic zone as		
mentioned in section F 2.1, F12, (2.1) and F3		

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B. Information required for Relia	nce: Option 1.2.1 Manufacturing site	certified by PIC/S member N	IRA
1) Submit the evidence of PIC/S-GMP compliance for the site			
where the finished dosage form is manufactured and batch			
release takes place.			
2)Submit Evidence of approval by the NRA of the country where			
the finished dosage form is manufactured, and batch release takes			
place. For this, a link should be provided to trace the evidence or			
registration or marketing authorization certificate			
3) Submit verifiable declaration of approval by at least 3 other			
NRAs. For this, a link should be provided to trace the evidence or			
registration or marketing authorization certificate. (NRA website			
shall verify that the product is registered there even though CoPP			
is provided)			
4)Product information for reliance as in Annex III:			
4) Found in or marion for remarke as in Almex in .			
4.1Evidence of approval from the reference organization with			
sufficient details (in one or several documents, e.g. evidence of MA,			
public1 evaluation reports or CoPP). This shall be submitted as a			
separate attachment. (THIS IS SAME AS IN POINT 2 OF THIS CLAUSE)			
4.2Dropriotony product name /Dropd Name			
4.2Proprietary product name /Brand Name:			

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4.3International Nonproprietary Name (INN) of the active			
pharmaceutical /Generic Name:			
pharmaceuticary deficite Name:			
4.4 Compositions:			
4.4 compositions.			
	- · · · · · · · · · · · · · · · · · · ·		
Component and quality standard	Function of the component in the	Quantity per unit (mg) %	%
, , , , , , , , , , , , , , , , , , ,	formulation	Quantity per anni (mg)	,-
Total			
. 544			
4.5 Strength:			
4.6 Pharmaceutical form (Dosage form):			
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4.7 Indication or Use of the product:		
4.8 Storage conditions:		
4.9 Shelf life of the product:		
4.10 WHO ATCC code:		
4.11 Stability summary and conclusions (including the storage		
statement and shelf-life) This to be attached		
4.12 Dispensing Category:		
4.13 Pack size / Volume:		
4.14 Date of approval by the reference organization: Date of		
approval with Link to verify		
4.15 Marketing authorization number from the reference		
organization (if applicable):		
4.16 Name(s) and complete address(es) (including specific		
unit/blocks) of the manufacturer(s) of the finished pharmaceutical		
product(s) [FPP(s)] or biological drug products(s) (DP(s)), including		
the final product release if different from the manufacturer:		
4.17 Detailed description of the primary and secondary packing		
including pack size or volume to be attached		

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4.18 Attach Product information for healthcare professionals		
(SMPC)		
4.19 Provide a valid GMP certificate or incorporate the information		
in this sheet as per the mentioned information below, if providing a		
certificate, it must be attached separately. cGMP certificates		
information, including:		
4.19.1 certificate number		
4.19.2 applicable standards		
4.15.2 applicable standards		
4.19.3 Site name		
4.19.4 Site address		
4.19.5 Issue date and validity		
4.19.6 Scope: products		
4.19.7 Scope: Manufacturing operations		
4.15.7 Scope. Manufacturing operations		
4.20 Attach Public evaluation reports (for new medicines only)		
4.21 Attach Declaration of sameness of the product as that of the		
product registered and approved in the reference NRA		
4.22Attach, as far as possible, the full unredacted assessment		
reports from the reference authority/institution		
reports from the reference authority/filstitution		

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4.23 Name and complete address (including specific unit/blocks) of			
the API/drug substance manufacturer(s):			
4.24 Attach detailed Description (visual appearance) of the finished			
pharmaceutical product (eg:color , texture etc of tablet, syrup etc)			
4.25 Attack Considerations for the finished made to			
4.25 Attach Specifications for the finished product			
4.26 Attach Product information for patients including product			
artwork			
4.27 Cost and propose retail price:			
F\Ctability study report soussing applicable alimetic con-			
5)Stability study report covering applicable climatic zone as			
montioned in coction E 2.1 E12 /2.1) and E2			
mentioned in section F 2.1, F12, (2.1) and F3			
	aco: Ontion 1 2 2 Manufacturing cita	of a WHO prognalified produ	icto
	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Reliant 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site https://extranet.who.int/prequal/medicines/prequalified/finished-	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Reliant 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site https://extranet.who.int/prequal/medicines/prequalified/finished-pharmaceutical-products	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site https://extranet.who.int/prequal/medicines/prequalified/finished-pharmaceutical-products 2) Submit Evidence of approval by the NRA of the country where	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site https://extranet.who.int/prequal/medicines/prequalified/finished-pharmaceutical-products 2) Submit Evidence of approval by the NRA of the country where the finished dosage form is manufactured, and batch release takes	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site https://extranet.who.int/prequal/medicines/prequalified/finished-pharmaceutical-products 2) Submit Evidence of approval by the NRA of the country where the finished dosage form is manufactured, and batch release takes place. For this, a link should be provided to trace the evidence or	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts
B. Information required for Relian 1) Submit Evidence that a WHO prequalified medicine or vaccine is manufactured and the batch released on the same site https://extranet.who.int/prequal/medicines/prequalified/finished-pharmaceutical-products 2) Submit Evidence of approval by the NRA of the country where the finished dosage form is manufactured, and batch release takes	nce: Option 1.2.2 Manufacturing site	of a WHO prequalified produ	ucts

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3) Submit Verifiable declaration of approval by at least 3 other			
NRAs. For this, a link should be provided to trace the evidence or			
registration or marketing authorization certificate. (NRA website			
shall verify that the product is registered there even though CoPP is			
provided)			
4)Product information for reliance as in Annex III:			
4.1Evidence of approval from the reference organization with			
sufficient details (in one or several documents, e.g. evidence of MA,			
public1 evaluation reports or CoPP). This shall be submitted as a			
separate attachment. (THIS IS SAME AS IN POINT 2 OF THIS CLAUSE)			
4.2Proprietary product name /Brand Name:			
4.3International Nonproprietary Name (INN) of the active			
pharmaceutical /Generic Name:			
4.4 Compositions:			
	Function of the component in the		
Component and quality standard	formulation	Quantity per unit (mg) %	%
	formulation		
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Total		
4.5 Strength:		
4.6 Pharmaceutical form (Dosage form):		
4.7 Indication or Use of the product:		
4.8 Storage conditions:		
4.9 Shelf life of the product:		
4.10 WHO ATCC code:		
4.11 Stability summary and conclusions (including the storage statement and shelf-life) This to be attached		
4.12 Dispensing Category:		
4.13 Pack size / Volume:		
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4.14 Date of approval by the reference organization: Date of approval with Link to verify		
4.15 Marketing authorization number from the reference		
organization (if applicable):		
4.16 Name(s) and complete address(es) (including specific		
unit/blocks) of the manufacturer(s) of the finished pharmaceutical		
product(s) [FPP(s)] or biological drug products(s) (DP(s)), including		
the final product release if different from the manufacturer:		
4.17 Detailed description of the primary and secondary packing		
including pack size or volume to be attached		
4.18 Attach Product information for healthcare professionals		
(SMPC)		
4.19Provide a valid GMP certificate or incorporate the information		
in this sheet as per the mentioned information below, if providing a		
certificate, it must be attached separately. cGMP certificates		
information, including:		
4.19.1 certificate number		
4.19.2 applicable standards		
4.19.3 Site name		

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4.19.4 Site address		
4.19.5 Issue date and validity		
4.19.6 Scope: products		
4.19.7 Scope: Manufacturing operations		
4.20 Attach Public evaluation reports (for new medicines only)		
4.21 Attach Declaration of sameness of the product as that of the		
product registered and approved in the reference NRA		
4.22Attach, as far as possible, the full unredacted assessment		
reports from the reference authority/institution		
4.23 Name and complete address (including specific unit/blocks) of		
the API/drug substance manufacturer(s):		
4.24 Attach detailed Description (visual appearance) of the finished		
pharmaceutical product (eg:color , texture etc of tablet, syrup etc)		
4.25 Attach Specifications for the finished product		
4.26 Attach Product information for patients including product		
artwork		
4.27 Cost and propose retail price:		

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B. Information required for Reliance:	e: Option 1.3.1 Collaborative registration procedure (CRP) - prequalified (PQ)		
1)Letter of manufacturer showing interest in going for this pathway.			
Evidence of approval from the country of manufacture (A link should be provided to trace the evidence or registration or marketing authorization certificate.)			
3) Evidence of approval and marketing in one reference country as mentioned in Annex II. (A link should be provided to trace the evidence or registration or marketing authorization certificate.)			
4)Product information for reliance as in Annex III :			
4.1 Evidence of approval from the reference organization with sufficient details (in one or several documents, e.g. evidence of MA, public1 evaluation reports or CoPP). This shall be submitted as a separate attachment. (THIS IS SAME AS IN POINT 2 OF THIS CLAUSE)			
4.2Proprietary product name /Brand Name:			
4.3International Nonproprietary Name (INN) of the active pharmaceutical /Generic Name:			

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4.4 Compositions:			
Component and quality standard	Function of the component in the formulation	Quantity per unit (mg) %	%
Total			
4.5 Strength:			
4.6 Pharmaceutical form (Dosage form):			
4.7 Indication or Use of the product:			

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4.8 Storage conditions:		
4.9 Shelf life of the product:		
4.10 WHO ATCC code:		
4.11 Stability summary and conclusions (including the storage		
statement and shelf-life) This to be attached		
4.12 Dispensing Category:		
4.13 Pack size / Volume:		
4.14 Date of approval by the reference organization: Date of		
approval with Link to verify		
4.15 Marketing authorization number from the reference		
organization (if applicable):		
4.16 Name(s) and complete address(es) (including specific		
unit/blocks) of the manufacturer(s) of the finished pharmaceutical		
product(s) [FPP(s)] or biological drug products(s) (DP(s)), including		
the final product release if different from the manufacturer:		
4.17 Detailed description of the primary and secondary packing		
including pack size or volume to be attached		
4.18 Attach Product information for healthcare professionals		
(SMPC)		

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4.19 Provide a valid GMP certificate or incorporate the information		
in this sheet as per the mentioned information below, if providing a		
certificate, it must be attached separately. cGMP certificates		
information, including:		
4.19.1 certificate number		
4.19.2 applicable standards		
4.19.3 site name		
4.19.4 site address		
4.19.5 issue date and validity		
4.19.6 Scope: products		
4.19.0 Scope. products		
4.19.7 Scope: Manufacturing operations		
4.20 Attach Public evaluation reports (for new medicines only)		
4.21 Attach Declaration of sameness of the product as that of the		
product registered and approved in the reference NRA		
4.22Attach, as far as possible, the full unredacted assessment		
reports from the reference authority/institution		
4.22 Name and complete address (including specific unit /hlasta) of		
4.23 Name and complete address (including specific unit/blocks) of		
the API/drug substance manufacturer(s):		

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4.24 Attach detailed Description (visual appearance) of the finished			
pharmaceutical product (eg:color , texture etc of tablet, syrup etc)			
4.25 Attach Specifications for the finished product			
4.26 Attach Product information for patients including product			
artwork			
4.27 Cost and propose retail price:			
5) Stability study report covering applicable climatic zone as			
mentioned in section F 2.1, F12, (2.1) and F3			
	C. Notification		
Evidence of approval by the NRA of the country where the			
finished dosage form is manufactured and batch release takes			
place. For this, a link should be provided to trace the evidence or registration or marketing authorization certificate.			
2) Verifiable declaration of regulatory status in at least 3 other			
NRAs. For this, a link should be provided to trace the evidence			
or registration or marketing authorization certificate.			
3) Stability study report covering applicable climatic zone as mentioned in section F 2.1, F12, (2.1) and F3			
4) Product information for reliance as in Annex III			
If the product is already available in ADL provide the product			
number			
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Copy Letter:

Cost price in USD:					
Retail price in USD:					
		FOR MT	G TO USE		
Prepared by:				Checked and Verified by:	
Pharmaceutical Officer				Senior Pharmacist	
Date:				Date:	
TECHNICAL DECISION OF THE NATIONAL P	PHARMACEUTICAL BOARD	D:			
APPROVED					
REJECTED					
NPB Chairman's Signature:					
Medicine and Therapeutic Goods Division, Maldives Food a	and Drug Authority	Document Created on: 09.08.20	22		
	for Product Registration Includ				
Doc Hame dudeline	roddoc megistration illelaa				

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Issued Date: 17.12.2024

Copy Letter:

Name:	
Date:	
Approved by:	Authorized by:
Pharmaceutical Specialist	Deputy Director General, Health Laboratory Services
Date:	Date:
-	

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Checklist for full dossier Applications

	•
	Rec.No: MTG/RE-ES/Re-0005/2024 - 000
Maldives Food and Drug Authority	
Ministry of Health	
Male`	
Republic of Maldives	
1. Registration Pathway	
Full procedure 1.1 Full dossier	5 years
2. Registration type	

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Date of submission	-	-	-
Date of Acceptance	-	<u>-</u>	-
A) New Registration			
B) Re-registration			
3. Product Evaluation Summary -	- -		
	Client submitted information	Evaluation Remarks	Check and verified
A) Legal status		-	-
Letter of appointment as the local agent if the applicant is local	Statement by Applicant (local) Submitted		
Letter from manufacturer to MFDA			

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B) General Product Information		
a) API information under any of the mentioned criteria		
b) Product name		
c) International Nonproprietary Name (INN) or the Active Pharmaceutical Ingredient (API) or Generic name including e. Pharmacopeia standard / Formulation of the product		
d) Non-active ingredient or Excipient including e. Pharmacopeia standard / Formulation of the product		
iii. For all the pediatric oral formulations including cough, cold and paracetamol formulation the certificate of analysis (COA) shall be submitted for all the excipients used, specifically if glycerin or glycerol or propylene glycol is used, verifying that it does not contain the impurities diethylene glycol (DEG) and ethylene glycol (EG).		

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iv. Manufacture validation protocols of the excipients		
shall be submitted specifically those that are at a risk for		
diethylene glycol (DEG) and ethylene glycol (EG)		
contamination. For such excipients each container of the		
excipients shall be tested for purity and validity and		
evidence documents shall be submitted		
v. The test mentioned above shall be as per the official		
monogram for purity.		
vi. Registration holder will ensure that the manufacturer		
-		
will perform impurity testing as identified by the		
manufacturer of innovator drug product like N-Nitroso		
dimethylamine (NDMA), N-Nitroso dimethylamine (NDEA)		
in valsartan, metformin etc.		
e. ii) ii. If there is no pharmacopeial formulation as		
mentioned in point c and d method validation report of		
the in-house method shall be provided which has to be		
endorsed by a third party. The third party can be an		
accredited laboratory, regulatory authority or any other		
external or internal assessment body as nominated by the		
·		
Authority.		
f) Dosage form of the product		

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g) Strength(s) of the product		
h) Volume of the preparation		
i. Product description, Container type and Pack sizes		
j. Route(s) of Administration		
k. Indication or Use of the product		
I. Therapeutic Class	ATC classification and code: Contraindication: Precaution:	
m. Storage conditions		
n. Shelf life of the product		
o. Dispensing Category		

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C2) Manufacturer responsible for lot release of the		
finished dosage form		
C3) Manufacturer responsible for packaging of the		
finished product, if different		
C4) Manufacturing License number		
Sample /artwork, Full package picture provided?		
Sample / artwork, i an package pictare provided.		
Documents as per registration pathway		
Whether the Full ICH M2 and M3 dossier parts provided?		
4. Is the formulation acceptable?		
Is the packaging acceptable? primary container, and		
packaging material label, box, blister/ strip foil		
Product information Literature/ Product package insert		
acceptable?		

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5. Is quality acceptable?		
API Validation report		
If applicable bioavailability studies against a benchmark product provided?		
Proof of Validation of the manufacturing method provided?		
Standard batch size quantity with label claim, Batch size, quantity of all active ingredients and excipients provided?		
Technical specifications and sources of all raw material(s) with pharmacopeia specification provided?		
Brief profile of manufacturer with products manufactured provided?		
Company profile (for newly registering manufacturers) provided?		
Manufacturing plant layout and machinery involved provided?		
Manufacturing and packaging process Provided?		

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List of personnel, their responsibilities and qualifications			
(Name, qualification and experience (in years) of the			
authorized key personnel)			
Regulatory decisions taken on this finished product from			
any regulatory authorities (A formal, signed statement			
from the manufacturer and/or MAH)			
Copy of the finished product specification (based on a			
reference to an official monogram and if an in-house			
method is used, it shall be endorsed by a third party)			
	Real-time:		
Shelf life validated by stability studies for climatic zone IV			
	Accelerated:		
Whether the stability report statement Provided?			
6. Documents for quality verification	L		
Certificate of Analysis for batch release, Certificate of			
Analysis of Finished product (CoA)			
1		I	

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Certificate of a Pharmaceutical product (CoPP)		
Registration status of Finished product in countries other		
than country of origin and country of origin		
Proof of registration of the finished product in a Category 1 country (for CAT 2.1)		
Valid GMP certificate attesting to the status of the		
manufacturer as to competency, of personnel, equipment		
and facilities.		
7. Other factors considered		
For cough/cold, paracetamol preparations: -Tests for		
DEG/EG submitted for all excipients and source validation		
of excipients submitted		
Can the medicine be used with the facilities and		
professional expertise available		
If the product sample is testable at NHL, does the product		
pass?		
pass:		

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number A) Cost price in USD: Price of the product in USD B) Retail price in USD:	A) Cost price in USD:	Is the medicine available in ADL, If provide the Product			
A) Cost price in USD: Price of the product in USD	A) Cost price in USD: Price of the product in USD				
Price of the product in USD	Price of the product in USD	Humber			
Price of the product in USD	Price of the product in USD				
			A) Cost price in USD:		
B) Retail price in USD:	B) Retail price in USD:	Price of the product in USD			
			B) Retail price in USD:		
			, .		
				1	1

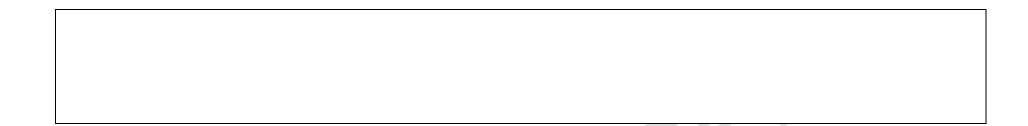
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FOR MTG TO USE	
Prepared by:	Checked and verified by:
Pharmaceutical Officer	Senior Pharmacist
Date:	Date:
TECHNICAL DECISION OF THE NATIONAL	
PHARMACEUTICAL BOARD:	
APPROVED	

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REJECTED	
Chairman's Signature:	
Name:	
Date:	
Approved by:	Authorized by:
Pharmaceutical Specialist	Deputy Director General, Health Laboratory
	Services
Date:	Date:

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MTG/RE-VA/Re 0125/2024

Registration of a Pharmaceutical Product Format for Sample submission 2024

N 0	Bra nd Na me	Generic Name (Inclu ding formulation if given eg. BP, IP, USP)	Bat ch No.	Stren gth	Dos age for m	Full Name of the Manufacture r (as in the product label)	Coun try of origin	Ex. Da te	Quanti ty	Remarks (Importer details)

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	For office use only							
Received by	Signature	Date						

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MTG/RE-SU/Re 0124/2024



Maldives Food and Drug Authority Ministry of Health Male` Republic of Maldives

Variation Applications of a Pharmaceutical Product

Product Details	
Brand Name:	
Generic Name:	
Strength:	

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Dosage Form:	
Product No:	
Registration No:	
Registered Date:	
Expiry Date:	
Date of submission:	

Importer Details: (write the product importer details)

Type of Variation

(insert Variation number)(Refer to the guideline for the variation number)), Mention if it is a change in artwork, label,manufacturing address, Product specification, change in packing insert or any.... (Also highlight the specific change from the document before and after. For this purpose no need to submit the full document if its large only the document name and the highlighted information changed (Before and after.)

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			Check
As Per Registered Product	Proposed Change In Detail	Variation Remarks	and
			verified
1.Brick Red colored capsule shaped	Brick Red colored capsule shaped biconvex coated		
biconvex film coated tablets having lip	tablets and plain on both side.(please highlight		
break line on one side and plain on other	the specific change),		
side. (please highlight the specific change)			
2.If it is a change in product label, specific	If it is a change in product label mention the		
information of product label as per	specific change in product label (in this case		
registered product must be mentioned.(in	proposed product label must be provided by		
this case previously submited label must	highlighting specific change) (Description of the		
be provided by highlighting specific	updated artwork or label)		
change) (Description of the original			
artwork or label)(Description of the			
original artwork or label)			

Medicine and Therapeutic Goods Division, Maldives Food and Drug Authority			Document Created on: 09.08.20)22
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	FOR MTG TO USE	
Checked by:	Checked and verified by:	Approved by:
	Mohamed Fazeen	Aishath Mohamed
Pharmaceutical Officer	Senior Pharmacist	Pharmaceutical Specialist
Date:	Date:	Date:
Sign:	Sign:	Sign:

Medicine and Therapeutic Goods Division, Maldives Food and Drug Authority		Document Created on: 09.08.2	022	
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CONFIRMATION OF API PREQUALIFICATION DOCUMENT

A complete copy of the Confirmation of API Prequalification document shall be provided, and it shall contain the following information.

- General properties discussions on any additional applicable physicochemical and other relevant API properties that are not controlled by the API manufacturer's specifications e.g., solubilities and polymorphs.
- **Sterility**-If the sterility of the FPP is based upon the sterile manufacture of the API then data on the sterilization process together with full validation data shall be provided.
- Specification the specifications of the FPP manufacturer including all tests and limits of the
 API manufacturer's specifications and any additional tests and acceptance criteria that are
 not controlled by the API manufacturer's specifications such as polymorphs and/or particle
 size distribution.
- **Batch analysis** results from two batches of at least pilot scale, demonstrating compliance with the FPP manufacturer's API specifications.
- Reference standards or materials information on the FPP manufacturer's reference standards.
- Stability data to support the retest period if either the proposed retest period is longer or the proposed storage conditions are at a lower temperature or humidity to that of the Prequalified API.

CERTIFICATES OF SUITABILITY (CEP)

CEP stands for certification of the suitability of European Pharmacopoeia monographs/Certificate of Pharmacopoeia.

The CEP is a document that is used to demonstrate the purity of a given API produced by a given manufacturer is suitably controlled by the relevant monograph(s) of the European Pharmacopoeia. By demonstrating grant a CEP for a given API, the suppliers of the API can prove such suitability to their pharmaceutical industry clients and Regulatory authority.

Certificate of Suitability of the European Pharmacopoeia (CEP) A complete copy of the CEP (including any annexes) shall be provided. The declaration of access for the CEP shall be duly filled out by the CEP holder on behalf of the FPP manufacturer or applicant who refers to the CEP.

In addition, a written commitment shall be included that the applicant will inform MFDA in the event that the CEP is withdrawn. It shall also be acknowledged by the applicant that withdrawal of the CEP would require additional consideration of the API data requirements to support the application. The written commitment shall accompany the copy of the CEP.

Along with the CEP, the applicant shall supply the following information

- **General properties** discussions on any additional applicable physicochemical and other relevant API properties that are not controlled by the CEP and Ph.Eur. monograph, e.g. solubilities and polymorphs
- Specification the specifications of the FPP manufacturer including all tests and limits of the CEP and Ph.Eur. monograph and any additional tests and acceptance criteria that are not controlled in the CEP and Ph.Eur. monograph, such as polymorphs and/or particle size distribution.

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- **Analytical procedures and validation** for any methods used by the FPP manufacturer in addition to those in the CEP and Ph.Eur. monograph.
- **Batch analysis** results from two batches of at least pilot scale, demonstrating compliance with the FPP manufacturer"s API specifications.
- Reference standards or materials information on the FPP manufacturer's reference standards.
- **Container closure system** specifications including descriptions and identification of primary packaging components. Exception: where the CEP specifies a container closure system and the applicant declares to use the same container closure system.
- **Stability** exception: where the CEP specifies a re-test period that is the same as or of longer duration, and storage conditions which are the same or higher temperature and humidity as proposed by the applicant. In the case of sterile APIs, data on the sterilization process of the API, including validation data, shall be included.

Annex-XI

TECHNICAL INFORMATION ON THE ACTIVE PHARMACEUTICAL INGREDIENT

The documentation shall also contain the following information:

1. General information:

- a) International Non-Proprietary Name.
- b) Chemical name
- c) Synonyms with complete reference
- d) Molecular and structural formulas
- e) Molecular weight
- f) Physical form
- g) Melting or boiling point
- h) Solubility
- i) Loss on drying
- j) Physical characteristics (crystalline, amorphous, particle size, solvation, etc.)
- k) pka and pH
- I) Preservation measures
- m) Organoleptic properties

2. API manufacturing process:

- a) Manufacturer(s): name, full address, company responsible for each manufacturing process step and quality
- b) control (including contracted companies, third parties).
- c) Description of the production process, including materials, equipment and operating conditions (for example,

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- d) temperature, pressure, pH, time ranges, stirring speed, etc.); and of the in-process controls.
- e) Identification of the critical steps including the respective tests and acceptance criteria
- f) Production process flowchart indicating the formation of intermediates and possible impurities, including the
- g) clarification of the respective chemical structures.
- h) Indication of the raw materials, solvents, catalysts, etc...
- i) Indicate the production scale and yield.
- j) Specifications of the raw materials and packaging materials.

3. Characterization:

Physicochemical tests allowing elucidation of the API structure:

- a) Analyses of an industrial batch evidencing the functional groups, the chemical structure and the molecular formula expected for the API.
- b) Possible Isomers.
- c) Polymorphism, describing the characteristics of the polymorph used and of others related to the active pharmaceutical ingredient.

4. Impurity profile:

- a) Description of the potential impurities, resulting from the synthesis, with a brief description and indicating the origin.
- b) Organic Impurities (of the process and related substances): raw materials (starting), related products,
- c) intermediate products, degradation products, reagents and catalysts.
- d) Inorganic Impurities: reagents and catalysts, heavy metals, inorganic salts.
- e) Residual solvents.

5. Quality Control of the API:

- a) Appearance
- b) Identification
- c) Assay
- d) Impurities (organic, inorganic and residual solvents)
- e) Physicochemical properties (pH, melting point, etc.).
- f) Particle size distribution.
- g) Polymorphism, including the adopted analytical methodology and results of the tests intended to determine the probable polymorphs of the ingredient.
- h) For chiral ingredients, data on the stereoisomer content.
- i) Water determination
- j) Microbiological limits: sterility, endotoxins (if applicable).
- k) Specific optical rotation (if applicable

6. Description of the analytical methodology:

- a) Validation of analytical methodology according to the current specific technical regulation for the validation of analytical and bioanalytical methods when the pharmacopeial methodology is not used.
- b) In case of pharmacopeial methodology, the company shall submit the method covalidation.

7. Packaging Material:

a) Description and specification of the primary packaging

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- b) Stability and Photostability Report
- c) photostability studies shall be conducted in compliance with the specific technical regulation /standard

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