GMP CHECKLIST

(Based on WHO Good Manufacturing Practices (GMP) for active pharmaceutical ingredients stated as per Annex 2- WHO Technical report Series(TRS), No. 957, 2010; Good Manufacturing Practice guide for Active Pharmaceutical Ingredients ICH Harmonised Triplicate Guideline stated as per ICH Q9; and GMP requirements as per Directives No. 2001/83/EC latest amended vide Directive 2011/62/EU)

1	Location and surroundings:	Self appraisal to be filled by the manufacturer along with all details (yes or no type reply will not be acceptable)	Observations to be noted by the inspecting team at the time of inspection	Remarks
1.1	How factory building is situated and controlled to avoid risk of contamination from external environment including open sewage, drain, public lavatory or any other factory which produces disagreeable or obnoxious, odors, fumes, excessive soot, dust, and smoke, chemical or biological emissions. <i>Pls specify industries /</i> <i>establishments adjoining</i> <i>manufacturing site.</i>			
2	Building and premises: -			
2.1	 How the building has been designed constructed and maintained to suit the manufacturing operations so as to produce drugs under hygienic conditions. Pls specify nature of construction used in the facility in respect of its maintenance and hygienic conditions. 			
2.2	Whether the building confirm to the conditions laid down in the Factories Act, 1948 Pls attach valid factory certificate/ license issued by the competent authority.			
2.3	 Specify how the premises used for manufacturing operations and testing purpose prevents contaminations and cross contamination is: a) Compatible with other drug manufacturing operations that may be carried out in the same or adjacent area. Pls specify any special criteria for 			

	the product manufacturered a a		
	the product manufacturered. e.g.		
	temperature, humidity, air class		
	requirements maintained for aseptic		
2.4	products, etc.		
2.4	b) Whether adequate working space		
	is provided to allow orderly and		
	logical placement of equipment,		
	materials and movement of		
	personnel so as to avoid risk of mix-		
	up between different categories of		
	drugs and to avoid possibility of the		
	contamination by suitable		
	mechanism.		
	Pls specify space left around the		
	machines. Pls attach equipment lay		
	out, men and material movement,		
	waste movement if applicable.		
2.5	c) Describe the pest, insects, birds		
	and rodents control system followed		
	in the premises.		
	Attach copy of pest / rodent control		
	schedule along with contract		
	agreement if any.		
2.6	d) What measures have been taken		
	to make Interior surface of (walls,		
	floors, and ceilings) smooth and free		
	from cracks, and to permit easy		
	cleaning		
	Specify material of construction and		
	finish for walls, ceiling, floor, coving		
	etc. i.e. whether Epoxy or PU		
	coated, kota / granite stone with		
	epoxy sealed joints, solid / GI /		
	gypsum / cal. Silicate board ceiling		
	with epoxy, PU or any other pre-		
	fabricated panel (GRP, powder		
	coated SS or Aluminum etc.) paint.		
2.7	e) What measures have been taken		
	so that the production and		
	dispensing areas are well lighted and		
	effectively ventilated, with air		
	control facilities.		
	Pls specify the lux level maintained		
	in various parts of the premise.		
2.8	Pls specify the air handling system		
	used in various areas like stores,		
	production, packing, QC areas etc.		
	production, packing, QC areas etc.		

2.9	f) Specify drainage system which		
	prevents back flow and entry of		
	insects and rodents into the		
	premises. Drains should be of		
	adequate size and should be		
	provided with an air break or a		
	suitable device to prevent back-		
	siphonage		
	(pls specify number and location of		
	drains installed)		
2.10	Containment area:		
	Any production activities (including		
	weighing, milling or packaging) of		
	highly toxic non-pharmaceutical		
	materials such as herbicides and		
	pesticides should not be conducted		
	using the buildings and/or		
	equipment being used for the		
	production of APIs. Handling and		
	storage of these highly toxic non-		
	pharmaceutical materials should be		
	separate from APIs.		
3	Water system: -		
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3.1	Whether the unit has validated		
	system for treatment of water drawn		
	from own or any other source to		
	render it potable in accordance with		
	standards specified by BIS or local		
	municipal norms.		
	Pls specify source of raw water and		
	give details of treatment processes,		
	sampling points, distribution and		
	storage system for raw and purified		
	water.		
3.2	How bio burden in purified water		
	controlled / reduced.		
3.3	How water tank are cleaned		
	periodically and records maintained		
	thereof. How water distribution		
	system is sanitized to control		
	microbial contaminations.		
4	Disposal of waste: -		
4.1	Specify the system of disposal of		
	sewage, and effluents (solid, liquid,		
	and gas) from the manufacturing		
	site.		
	(Enclosed the copy of NOC obtained		
	from State Pollution Control Board		
	in this regard).		

4.0	Whathan provision for discout of	I	
4.2	Whether provision for disposal of		
	bio-medical waste made as per the		
	provisions of the Bio Medical Waste		
	(Management and Handling) Rules		
	1996.		
5	Warehousing Area: -		
5.1	Whether adequate areas have been		
	allocated for warehousing of Raw		
	Materials, intermediates, Packaging		
	Material, products in quarantine,		
	finish products, rejected or returned		
	products.		
	How these areas marked or		
	segregated.		
	Please specify the total area		
5.0	provided for warehousing.		
5.2	How the warehousing areas being		
	maintained to have good storage		
	conditions. Are they clean and dry and maintained within acceptable		
	temperature limits?		
5.3	Specify the storage arrangement		
5.5	provided for materials which		
	sensitive to temperature, humidity		
	and light and how the parameters are		
	monitored.		
	Is cold room or deep freezers		
	required for storage of goods? If yes,		
	how the temperature is monitored.		
5.4	Whether proper racks, bins and		
	platforms have been provided for the		
	storage.		
5.5	Whether receiving and dispatch bays		
	are maintained to protect in coming		
	and out going materials.		
5.6	How incoming materials are treated		
5.0	and cleaned before entry into the		
	plant.		
	Please specify the cleaning system		
	for the outer surface of the		
	container.		
5.7	How quarantined materials are		
	segregated from other materials.		
	How access to quarantined area is		
	restricted.		
5.8	Whether separate sampling area for		
	active Raw Materials and Excipients		
	is provided and maintained.		
	If yes, what is the control on entry of		
	material and men into the sampling		
	area. Whether reverse LAF have		
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	been provided for sampling.		
	Whether log book for sampling		
	booth maintained.		
	If not what provision has been made		
	for sampling so as to prevent		
	contamination, cross contamination		
	and mix-ups at a time of sampling.		
5.9	Specify the arrangements		
	provided to sample the primary		
	packaging materials foils, bottles,		
	etc which are used as such.		
5.10	Pls specify sampling plan used.		
	Which type of sampling tools are		
	used and how they are cleaned, dried		
	and maintained.		
5.11	How containers are cleaned before		
5.11	and after sampling. Who carries out		
	the sampling?		
	(Pls specify whether the sampling is		
5.12	carried out as per the current SOP).		
5.12	What precautions are taken during		
	sampling of photosensitive,		
5.13	hygroscopic materials?		
5.15	What provisions have been made for		
	segregated storage of rejected, recalled or returned materials or		
	products.		
	How is the access to these areas		
	restricted.		
5.14	How highly hazardous, poisonous		
	and explosive materials, narcotics,		
	and psychotropic drugs are handled		
	and stored.		
	How these areas are safe and secure.		
	Is there certification from competent		
	authority for handling of explosives		
	etc. If any. Pls attach the certificate		
	issued by the competent authority.		
5.15	How printed secondary packaging		
	materials are stored in safe, separate		
	and secure manner.		
5.16	Specify the arrangement provided		
	for dispensing of starting materials.		
	What is the control on entry of		
	material and men into the dispensing		
	area? Whether reverse LAF have		
	been provided for dispensing with		
	back ground clean air supply.		
	Whether pressure differential is		
	maintained between the dispensing		
	and adjacent areas.		
	une aujacent areas.		

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5.17	Which type of dispensing tools are		
	used and how they are cleaned, dried		
	and maintained.		
	How containers are cleaned before		
	and after dispensing. Who carries		
	out the dispensing?		
	(Pls specify whether the dispensing		
	is carried out as per the current		
5.10	SOP).		
5.18	How and where sampling of sterile		
- 10	materials carried out.		
5.19	What steps are taken against		
	spillage, breakage and leakage of		
	containers?		
5.20	What provisions have been made to		
	prevent the entry of rodents, insects,		
	birds.		
	Which substance is used for pest		
	control and how it is handled.		
	(Pls specify whether the pest control		
5.01	is carried out as per the SOP).		
5.21	Whether record of master labels is		
	maintained for comparision to		
6	issued labels?		
6	Production Area: -		
6.1	Please specify the design of the		
	manufacturing area which allow uni-		
	flow and logical sequence of		
	operations so as to prevent product		
	contamination/ mix ups.		
	Is there any criss cross of flow of		
	materials and men.		
	Specify the position of IPQC lab in the manufacturing area		
	the manufacturing area.		
	Please specify whether non storage areas used for storage of any		
	material.		
6.2	Whether separate dedicated and self-		
0.2	contained facilities have been		
	provided for the production of		
	sensitive pharmaceutical product		
	like Penicillin, Biological		
	preparation with like micro-		
	organism, Beta lactam, Sex		
	Hormones and Cytotoxic substances.		
	If yes pls explain how and attach		
	copy of plan of premises of each		
	copy of plan of prennises of cach		
	category of drug.		

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6.3	Please specify the provisions of		
	storage of dirty, washed and cleaned		
	equipment parts, tool room, in		
	process storage areas		
	etc. Which provide sequential /		
	logical manner so as to prevent		
	contamination and cross		
	contamination?		
6.4	Please specify how service lines like		
	pipe work, electrical fittings,		
	ventilation openings etc. are		
	identified by colors for nature of		
	supply and direction of the flow.		
	Whether service lines in production		
	areas are through service pendants.		
	If not, how they are placed so as to		
	avoid accumulation of dust.		
7	Ancillary areas: -		
7.1	Please specify the position of rest		
	and refreshment rooms and mention		
	whether they are separate and not		
	leading directly to the manufacturing		
	and warehouse areas.		
7.2	Are there general change rooms in		
	plant?		
	Are toilets, change room separate		
	from mfg. Area? Pls specify number		
	of washing station & toilets		
	provided for number of users.		
	Whether change facilities separated		
	for both sexes.		
	How many sets of protective		
	garments provided for each		
	personnel entering production area.		
	Is there in house general laundry for		
	garment washing / cleaning? If not		
	how garments washing are carried		
	out and monitored		
7.3	Whether maintenance workshop is		
	separate and away from production.		
7.4	Whether animals for production or		
	testing are housed in the facility if so		
	whether areas housing animals are		
	isolated from other areas.		
	Please specify the provision of air		
	conditioned and ventilation system		
	for the animal house.		
	How quarantined, under test and		
	tested animals housed and		
	controlled.		
	How animal carcass are disposed of.		
	Pls attach copy of CPCSEA.		
	tested animals housed and controlled. How animal carcass are disposed of.		

8.7	Whether separate areas provided for		
	sterility testing within microbiology		
	lab.		
	Whether support areas are under		
	AHU.		
	Whether double door autoclave		
	provided for sterilization of		
	materials.		
8.8			
0.0	Whether entry to the sterility area is		
	through three air lock systems.		
	What is the air class of these testing		
	areas and whether pressure		
	difference is maintained in these		
	areas?		
8.9	Which types of workbenches are		
	provided in these areas for testing?		
	When was the last filter integrity		
	tests performed on HEPA filters		
8.10	How waste (cultures etc) disposed		
	of.		
	Whether in case of antibiotic		
	potency testing, statistical proof of		
	the determination of potency and		
	validity of the test carried out.		
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9 9.1	Personnel: -		
9.1	Whether the manufacturing and		
	testing of drugs is conducted under		
	approved technical staff		
	Names of Technical Staff alongwith		
	qualification & experience		
	For Manufacturing: -		
	For Analysis:		
9.2	Please specify whether head of Q.C.		
	is independent of manufacturing unit		
9.3	Name, qualification and experience		
	of the personnel responsible for		
	Quality Assurance function.		
9.4	Whether responsibilities for		
	production and QC laid down and		
	followed.		
9.5	Whether adequate number of		
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	personnel employed in direct		
	proportion to the work load.		
9.6	What is the firm"s policy on training		
	of personnel at various levels?		
9.7	How is Periodic assessment of the		
	training checked?		
10	Health, clothing and sanitation of		
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10.1	Whether personnel handling Beta lactam antibiotics are tested for		
	penicillin sensitivity before		
	employment.		
10.2	Whether personnel involved in		
	handling of sex hormones, cytotoxic		
	and other portent drugs are		
	periodically examined for adverse		
	effect.		
	(Pls specify whether the current SOP		
	is followed or not).		
10.3	Whether all personnel prior to		
	employment have undergone		
	medical examination including eye examination and all free from		
	Tuberculosis, skin and other communicable or contagious		
	diseases		
10.4	Whether there is a SOP for medical		
	examination.		
10.5	Pls give name and qualification of		
	contracted medical officer for		
10.6	medical examination.		
10.6	Whether investigational reports,		
	films of X rays etc. preserved. Whether records of such medical		
	examination are maintained thereof		
10.7			
10.7	Whether all personnel are trained to ensure high level of personal		
	hygiene.		
	Pls attach training calendar of last		
	two years.		
10.8	Whether proper uniforms and		
10.0	adequate facilities for personal		
	cleanliness are provided.		
	Pls specify nature and type of dress		
	used by the personnel in		
	various areas of operation.		
	How many dress/footwear have been		
	provided to each personnel.		
	Please specify whether cross over		
	bench is in place in the change room		
	and if so whether it rule out the		
	possibility of entering dust particle		
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	Whether for sterile garments in		
	to the clean side. Whether arrangements provided for cleaning of outside dust and dirt from foot Please specify whether hands are disinfected before entering the production area Whether for sterile garments in		

	house clean laundry has been	
	provided.	
11	1	
11	Manufacturing Operations and Controls: -	
11.1	Whether the contents of all vessels	
	and containers used in manufacture	
	and storage is conspicuously labeled	
	with the name of the products. Batch	
	no, Batch Size, and stage of	
	manufacture along with signature of	
	technical staff.	
11.2	Whether the products not prepared	
	under aseptic conditions are free	
	from pathogens like Salmonella,	
	Escherichia coli, Pyocyanea etc.	
11.3	If yes, pls give brief account of	
	measures taken to assure freedom	
	from pathogens.	
11.4	Precautions against mix-up and	
	cross-contamination: -	
11.4.1	Whether proper AHU, pressure	
	differential, segregation, status	
	labeling have been provided to	
	prevent mix-up and cross-	
	contamination in manufacturing area	
11.4.2	Pls specify the areas of dust	
	generation and mechanism involved	
	in controlling the dust.	
11.4.3	Do all the areas have their own	
	independent air locks separately for	
	men and material entry.	
11.4.4	What criteria of pressure differential	
	have been set for production v/s	
	adjoining areas.	
11.4.5	Whether various operations are	
	carried out in segregated areas.	
11.4.6	Whether processing of sensitive	
	drugs like Beta lactum Antibiotics	
	and Sex Hormones is done in	
	segregated areas with independent	
	AHU and proper pressure	
	differentials along with	
	demonstration of effective	
	segregation of these areas with	
11 4 7	records.	
11.4.7	Please specify what measures has	
	been taken to prevent contamination	
	of products with Beta Lactum	
	Antibiotics, Sex harmons and cyto	
	toxic substances	

11.4.8	What measures has been taken to prevent mix-ups during various stages of production.		
11.4.9	Whether equipments use for production are labeled with their current status.		
11.4.10	What is the policy for the use of Recovery material?		
11.4.11	Whether packaging lines are independent and adequately segregated.		
11.4.12	How line clearance is performed. Whether records of line clearance is maintained according to appropriate checklist		
11.4.13	Whether separate coding area has been provided or online coding is performed How coding procedure is controlled.		
11.4.14	Please specify how temperature, humidity and air filtration are controlled in the areas where raw material and/or products are exposed and handled.		
11.4.15	How access of authorized persons to manufacturing areas including packaging is controlled.		
11.4.16	Whether separate gowning provision is follows before entering into the procedure.		
11.4.17	Whether segregated secured areas for recall or rejected materials or for such material which are to be processed or recovered are provided. Please specify the room No. of such areas in the plant.		
11.5	Sanitation in the Manufacturing		
11.5.1	<i>areas:-</i> Specify the cleaning procedure of the manufacturing areas. Whether cleaning procedure is validated. Please specify validation protocol No. of the same.		
11.5.2	Whether the manufacturing areas are used as the general thoroughfare and storage of materials not under process.		

11.5.3	Whether a routine sanitation		
11.3.3			
	program is in place.		
	Please specify detailed account of		
	sanitation proramme specific to		
11.5.0	various areas, equipment.		
11.5.3	Dose the location facilitate cleaning		
	of equipment as well as the cleaning		
	of the areas in which they are		
11 7 4	installed.		
11.5.4	Whether production area is		
	adequately lit. If yes.		
	Please give lux levels provided in		
	production, visual inspect		
12	Raw Materials: -		
12.1	Whether the hard copies of records		
	of Raw Materials are maintained.		
12.2	Please specify the procedures		
	followed receiving and processing of		
	in-coming materials (Starting		
	materials and packing material).		
12.3	Whether first in / first out or first		
	expiry principal has been adopted.		
12.4	How they are labeled and stored as		
	per their status – Under Test,		
	Approved and Rejected		
12.5	Whether incoming materials are		
	purchased from approved sources.		
12.6	What is the procedure for approving		
	the source for incoming materials.		
12.7	Whether the raw materials are		
	directly purchased from the		
	manufacturers.		
12.8	Whether list of approved vendors is		
	available to the user.		
12.9	How damaged containers are		
	identified recorded and segregated.		
12.10	How damaged containers are		
	identified recorded and segregated.		
12.11	Whether all the containers of each		
	batch of starting materials is		
	sampled for identification test.		
12.12	Whether labels of raw material in		
	the storage area have information		
	like		
	(a) designated name of the product		
	and the internal code reference,		
	where applicable, and analytical		
	reference number;		
	(b) manufacturer's name, address		
	and batch number;		
	(c) the status of the contents (e.g.		

	quarantine, under test, released, approved, rejected); and (d) the manufacturing date, expiry date and re-test date.	
12.13	Whether separate areas are provided for under test, approved and rejected materials.	
12.14	How control on temperature and humidity conditions, wherever necessary, maintained in these storage areas.	
12.15	How the containers from which samples have been drawn labeled.	
12.16	Please specify the procedures by which it is ensured that the raw materials which has been released by the Quality Control Department and which are within their shelf life are going to be used in the product.	
12.17	How materials are stacked in the Stores i.e on Pallets, racks etc.	
13	Equipment: -	
13.1	Whether the equipments are designed aiming to minimize risk of error and permit effective cleaning in order to avoid cross contamination, build up of dust	
13.2	Whether all equipment are provided with log book.	
13.3	Please specify the procedures to clean the equipment after each batch production.	
13.4	Whether validity period for use after the cleaning of equipment is specified.	
13.5	Whether separate area is provided for storage of machine parts etc.	
13.6	Whether balances and other measuring equipments with appropriate range are available in the Raw Material stores & production areas and they are calibrated in accordance with SOP maintained.Specify the calibration schedule of the balances.	

13.7	Diago anagify material of		
15.7	Please specify material of		
	construction of contact parts of the		
13.8	production equipments.		
15.8	Which types of lubricants are used		
	in the equipment.		
	Specify the quality and control		
12.0	reference No. of these lubricants.		
13.9	Specify the procedures to remove		
	defective equipments from		
1.4	production areas.		
14	Documentation and Records: -		
14.1	How the documents are designed,		
	prepared, reviewed and controlled to		
	provide an audit trail.		
	Whether documents are approved		
	signed and dated by appropriate and		
	authorized person.		
	Whether documents are approved		
	signed and dated by appropriate and		
	authorized person.		
	Whether documents specify title,		
	nature and purpose.		
	Whether documents are regularly		
	reviewed and kept up to date. If yes.		
	Please specify review period. Please attached the list of documents		
14.2	maintained by the firmWhether the records are made at the		
14.2	time of each operation in such a way		
	that all significant activities		
	concerning to the production are		
	traceable.		
14.3	Whether data is recorded by		
14.5	electronic data processing system or		
	by other means. If by electronic data		
	processing system then how access		
	is controlled to enter, modify etc. the		
	data.		
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14.4	Whether master formula and		
	detailed operating procedures are		
145	maintained as hard copy.		
14.5	Who is responsible for maintenance of these records.		
15			
15	Labels and Other Printed		
15 1	Materials:		
15.1	Whether the printing is in bright		
	colour and legible on labels and		
15.0	other printed habels (art work) are		
15.2	How printed labels (art work) are		
	approved. Is there any SOP for this		
	if yes please give current SOP No.		

15.0	XX71 ' 1 1 1 1' / ' 1		
15.3	Which colour coding system is used		
	to indicate the status of a product		
	and equipment.		
15.4	How printed packaging materials,		
	product leaflets etc. are stored		
	separately to avoid chances of mix-		
	up.		
15.5	How labels cartons boxes circulars		
	inserts and leaflets are controlled.		
15.6	Whether the samples from the bulk		
	are drawn tested, approved and		
	released prior to packaging and		
	labeling.		
	How carryout the sampling		
15.7	How records of receipt of all		
	labeling and packaging materials are		
	maintained.		
15.8	Whether re-conciliation of used		
15.0	packaging materials is maintained.		
	Whether unused packaging materials		
	return to the store or destroyed.		
15.9	How returned/unused packaging		
15.7	material like foils is controlled so as		
	to prevent contamination and cross-		
	contamination.		
15.10	How the labels of reference standard		
15.10	and culture maintained.		
16			
16.1	Quality Assurance: -		
10.1	Specify the comprehensive quality		
	assurance system maintained by the		
	firm <i>Inter-alia</i> to cover deviation,		
	reporting, investigation and change		
	control.		
	How the products are designed and		
160	developed in accordance with GMP.		
16.2	Please specify the arrangements		
	provided to ensure that correct		
	starting and packaging materials are		
1 6 0	used for manufacture.		
16.3	Please specify the mechanism by		
	which all control like IP QC		
	Calibration, Validation etc. are		
1 < 1	ensured.		
16.4	Please specify the mechanisms to		
	ensure that the finished product has		
	been correctly processed and		
	checked in accordance with the		
	established procedures.		
16.5	Please specify the mechanisms to		
	ensure that Pharmaceuticals		
	products are released for sale by		
	authorization person.		

17	Self Inspection and Quality Audit: -		
17.1	Whether the firm has constituted a		
17.1	self inspection team supplemented		
	with a quality audit procedure to		
	evaluate that GMP is being		
	followed. If no. How internal audits		
	are carried out.		
17.2			
17.2	What is the system of monitoring,		
17.0	evaluation of self inspection.		
17.3	How conclusion and recommended		
	correcting actions are followed and		
	adopted.		
17.4	What is the frequency of self-		
	inspection.		
17.5	Is there any proforma for carrying		
	out the self-inspection.		
	Please indicate the date of last self-		
	inspection.		
18	Quality Control System: -		
18.1	Please specify the details of quality		
	control system of the unit.		
18.2	How the reference standards are		
	stored, evaluated and maintained.		
	Please provide list of reference		
	standard and reference impurities		
	procured from the authentic sources.		
18.3	Please specify the procedures of		
	preparation of working standard		
	from the reference standards.		
18.4	Whether SOPs for sampling,		
	inspecting, testing of Raw Materials,		
	Finish products, Packing Materials		
	and for monitoring environmental		
	conditions are available.		
18.5	Whether approved specifications for		
10.5	different materials, products,		
	reagents, solvents including test of		
	identity content, purity and quality		
	available.		
18.6	How reference samples from each		
10.0	batch of the products are maintained.		
18.7	Who releases batch of the products		
10.7	for sale		
18.8	Whether there is check list for		
10.0			
	release of a batch. Please specify current SOP No. for batch release.		
10.0			
18.9	Please specify the sampling		
	procedures from various stages of		
10.10	production.		
18.10	How it is ensured that the sample		
	collected are representative of the		
	whole batch.		

10.11	Discos ana sife the meson during for			
18.11	Please specify the procedures for			
10.10	carrying out the stability studies.			
18.12	Under what condition stability			
	studies of the products are tested.			
	How many stability chambers have			
	been provided.			
18.13	How self life is assigned to a			
	product. Please give current stability			
	protocol No.			
18.14	Whether records of stability studies			
	are maintained.			
18.15	Please attach stability calendar of			
	last year.			
18.16	How complaints are investigated.			
18.17	How instruments are calibrated and			
	at which interval.			
18.18	How testing procedure validated			
	before they are adopted for routine			
	testing.			
18.19	Specify the validation procedure is			
	responsible for validation of			
	procedures.			
18.20	How validation procedures are			
	documented (Please indicate various			
	protocols/ recoding system applied			
	during validation).			
18.21	Whether specifications for raw			
	materials intermediates final			
	products and packaging materials			
	are available.			
18.22	Whether periodic revision of these			
	specifications are carried out.			
	Please specify No. of STPs being			
	maintained by the firm.			
18.23	Which pharmacopoeias in original			
	are available in the plant.			
19	Specifications: -			
19.1	Whether specification of raw			
	material include.			
	(a) the designated name and internal			
	code reference;			
	(b) reference, if any, to a			
	pharmacopoeial monograph;			
	(c) qualitative and quantitative			
	requirements with acceptance limits;			
	(d) name and address of			
	manufacturer or supplier and			
	original manufacturer of the			
	material;			
	(e) specimen of printed material;			
	(f) directions for sampling and			
	testing or reference to procedures;			
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	(g) storage conditions; and		
	(h) Maximum period of storage		
	before re-testing.		
	Whether specification of finished		
	product include		
	(a) the designated name of the		
	product and the code reference;		
	(b) the formula or a reference to the		
	formula and the pharmacopoeial		
	reference;		
	(c) directions for sampling and		
	testing or a reference to procedures;		
	(d) a description of the dosage form		
	and package details;		
	(e) the qualitative and quantitative		
	requirements, with the acceptance		
	limits for release;		
	(f) the storage conditions and		
	precautions, where applicable, and		
	(g) the shelf-life.		
19.2	Whether the container and closures		
	meet the pharmacopial		
	specifications.		
	Whether second hand or used		
	containers and closures used.		
20	Master Formula Records: -		
20.1	How master formula records are		
	prepared, authorized and controlled.		
20.2	Whether head of production, quality		
	control and quality assurance unit		
	endorse this documents. Whether		
	master formula is batch size specific.		
20.3	Whether all products have master		
	formula containing.		
	(a) the name of the product together		
	with product reference code relating		
	to its specifications;		
	(b) the patent or proprietary name of		
	the product along with the generic		
	name, a description of the dosage		
	form, strength, composition of the		
	product and batch size;		
	(c) name, quantity, and reference		
	number of all the starting materials		
	to be used. Mention		
	shall be made of any substance that		
	may "disappear" in the course of		
	processing.		
	(d) a statement of the expected final		
	yield with the acceptable limits, and		
	of relevant intermediate yields,		
	where applicable.		
	(e) a statement of the processing		

	location and the principal equipment		
	to be used.		
	(f) the methods, or reference to the		
	methods, to be used		
	for preparing the critical equipments		
	including cleaning, assembling,		
	calibrating, sterilizing;		
	(g) detailed stepwise processing		
	instructions and the time taken for		
	each step;		
	(h) the instructions for in-process		
	control with their limits;		
	(i) the requirements for storage		
	conditions of the products, including		
	the container, labeling and special		
	storage conditions where applicable;		
	(j) any special precautions to be		
	observed;		
	(k) packing details and specimen		
	labels.		
21	Packaging Records: -		
21.1	Whether authorized packaging		
	instructions for each products, pack		
	size and type are maintained and		
	complied with.		
	Whether following are included in		
	the packaging instructions.		
	(a) Name of the product;		
	(b) the pack size expressed in terms		
	of the weight or volume of the		
	product in the final container;		
	(d) complete list of all		
	the packaging materials required for		
	a standard batch size, including quantities, sizes and types with the		
	code or reference number relating to		
	the specifications of each packaging		
	material.;		
	(e) reproduction of the relevant		
	printed packaging materials and		
	specimens indicating where batch		
	number and expiry date of the		
	product have been applied;		
	(f) special precautions to be		
	observed, including a careful		
	examination of the area and		
	equipment in order to ascertain the		
	line clearance before the operations		
	begin.		
	(g) description of the packaging		
	operation, including any significant		
	subsidiary operations and equipment		
	to be used;		
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	(h) details of in-process controls		
	with instructions for sampling and		
	acceptance; and		
	(i) Re-conciliation after completion		
	of the packing and labeling		
	operation.		
	(j) Whether line clearance records		
	are part of batch packing records.		
22	Batch Processing Records		
	(BPR)		
22.1	Whether BPR are based on current		
	master formula record.		
22.2	How BPR are designed to avoid		
22.2	transcription errors.		
	Whether the Batch Processing		
	Records for each product on the		
	basis of currently approved master		
	formula is being maintained.		
	Whether following information are		
	recorded in BPR		
	(a) the name of the product,		
	(b) the number of the batch being		
	manufactured,		
	(c) dates and time of		
	commencement, significant		
	intermediate stages and completion		
	of production.		
	(d) initials of the operator of		
	different significant steps of		
	production and where appropriate,		
	of the person who checked each of		
	these operations,		
	(e) the batch number and/or		
	analytical control number as well as		
	the quantities of each starting		
	material actually weighed,		
	(f) any relevant processing operation		
	or event and major equipment used,		
	(g) a record of the in-process		
	controls and the initials of the		
	person(s) carrying them out, and the		
	results obtained,		
	(h) the amount of product obtained		
	after different and critical stages of		
	manufacture (yield),		
	(i) comments or explanations for		
	significant deviations from the		
	expected yield limits shall be given,		
	(j) notes on special problems		
	including details, with signed		
	authorization, for any deviation from		
	the Master Formula,		
	(k) Addition of any recovered or		
	(K) Addition of any recovered of		

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	Samples		
	-		
	(i) SOP for training and hygiene for		
	and sanitation		
	(g) SOP for Analytical		
	(f) SOP for equipment assembly and		
	(c) SOP for sampling		
	Equipment		
	(b) SOP for each instrument and		
	material and other materials (b) SOP for each instrument and		
	(b) SOP for each instrument and		
	(b) SOP for each instrument and		
	(d) SOP for batch numbering		
	(e) SOP for testing		
	(f) SOP for equipment assembly and		
	(f) SOP for equipment assembly and		
	validation		
	(g) SOP for Analytical		
	(g) SOP for Analytical		
	apparatus and calibration		
	(h) SOP for maintenance, cleaning		
	(h) SOP for maintenance, cleaning		
	(h) SOP for maintenance, cleaning		
	and sanitation		
	and sanitation		
	and sanitation		
	(i) SOP for training and hygiene for		
	the personal		
	-		
	(j) SOP for retaining reference		
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	(k) SOP for handling, re-processing		
	and recoveries		
	(l) SOP for distribution of the		
	product		
	-		
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	(m) SOP for warehousing of		
	products.		
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	Whether applicable SOPs are		
	1 I		
	available in each area where they are		
	•		
	required.		
	-		
	Whether recording formats are		
	-		
	referred in SOP.		
	Is there SOP for writing on SOP		
	Is there SOP for writing an SOP.		
24			
24	Reference Samples		
24.1	-		
24.1	Specify the procedures for collection		
	of reference samples of active		
	ingredients and finished		
	-		
	formulations and how they are		
	stored and maintained.		
	stored and maintained.		
25	Poprocessing and Pocoveries		
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23	Reprocessing and Recoveries		
25.1	Is appropriate Validation of recoveries and reprocessing done is		

	being performed?	
26	Distribution records	
26.1	Whether pre dispatch inspections are	
20.1	carried out before release.	
26.2	Whether periodic audits of	
	distribution center are carried out to	
	access warehousing practices	
26.3	Whether distribution records are part	
	of the batch record. If not how batch	
	wise distribution record up to retail	
	levels are maintained.	
26.4	Whether instruction for warehousing	
	and stocking of products like LVPs,	
	Heat sensitive etc are available in	
	store.	
26.5	Whether Good Distribution	
	Practices followed	
27	Validation and Process	
	Validation: -	
27.1	Specify the validation policy of the	
	company.	
	Whether validation master plan has	
	been prepared.	
27.2	Whether validation studies of	
	processing, testing and cleaning	
	procedures are conducted as per pre	
	defined protocol.	
27.3	How records and conclusion of such	
	validation studies are prepared and	
AF 1	maintained.	
27.4	Whether master formula is based on	
	approved process validation.	
27.5	Specify how significant changes to	
	the manufacturing process	
	equipments material etc are	
27.6	controlled.	
27.6	Whether DQ,IQ,OQ & PQ are in	
	place for all major equipment and facility.	
27.7	Whether validation records of all	
21.1	utilities and major equipments are	
	available.	
28	Product Recalls: -	
28.1	Specify the product recall system	
20.1	followed by the firm.	
	How promptly recall operation at the	
	level of each distribution channel	
	up-to the retail level can be carried	
	out.	
	Whether there is a SOP for recall of	
	products clearly defining	
	responsibility, procedure, reporting,	

	re-conciliation etc.		
29	Complaints and Adverse Reactions:		
29.1	Specify the review system for		
	complaints concerning the quality of		
20.2	products.		
29.2	How records of complaint maintained.		
29.3			
29.5	Whether reports of serious complaints with comments and		
	documents immediately sent to		
	Licensing Authority		
29.4	Is there any criteria for action to be		
27.4	taken on the basis of nature of		
	complaint.		
30	Site Master file: -		
30.1	Whether all the relevant information		
50.1	have been included in the site master		
	file.		
30.2	Whether quality policy has been		
	included in the site master file.		
	Please attach the current version		
30.3	Is there a master plan (Master		
	validation plan) covering:		
30.4	Resources and those responsible for		
	its implementation.		
30.5	Identification of the systems and		
	processes to be validated		
30.6	Documentation and standard		
	operating procedures (SOPs), Work		
	Instructions and Standards		
	(applicable national and		
	international standards)		
30.7	Validation list: facilities, processes		
	(e.g. aseptic filling), products		
30.8	Key approval criteria		
30.9	Protocol format		
30.10	Each validation activity, including		
	re-validation and reasonable		
	unforeseen events (power failures,		
	system crash and recovery, filter		
	integrity failurer. Please attach		
20.11	validation calendar.		
30.11	Pls specify whether the critical		
	processes validated Prospectively, retrospectively or concurrently.		
30.12	Whether validation of following		
30.12	performed and documented:		
	Analytical methods, Production and		
	assay equipment, Sterile production		
	processes, Non-sterile production		
	processes, Ron-sterne production processes, Cleaning procedures,		
	Critical support systems (purified		
	Cinical support systems (purified		

	water water for injections sin		
	water, water for injections, air,		
20.12	vapor, etc.), Facilities		
30.13	Please list reasons considered		
	important for validation or re-		
	validation.		
30.14	In case electronic data processing		
	systems are used, are these		
	validated?		
	Please specify whether periodical		
	challenge tests performed on the		
	system to verify reliability.		
30.15	Are the validation studies performed		
	according to pre-defined protocols?		
	Is a written report summarized,		
	results and conclusions prepared and		
	maintained? Is the validity of the		
	critical processes and procedures		
	established based on a validation		
	study?		
30.16	Are criteria established to assess the		
	changes originating a revalidation?		
	Are trend analyses performed to		
	assess the need to re-validate in		
	order to assure the processes and		
	procedures continue to obtain the		
	desired results?		
31	WATER SYSTEM		
	PURIFIED WATER		
	WATER FOR INJECTIONS		
31.1	Please specify whether waster		
	system qualification (IQ, OQ and		
	PQ) has been carried out as per		
	protocol and repots have been		
	prepared and maintained.		
31.2	Whether IQ protocol include at least		
51.2	facility review, equipment		
	specification vs. design, welding		
	roughness testing on pipelines,		
	absence of dead points / section in		
	the pipelines, pipe and tank		
	passivation, drawings, SOP for		
	operations, cleaning, sanitation,		
	maintenance and calibration of		
	gadgets. Whether its report includes		
	Conclusion / Summary, description		
	of the performed assay, Data tables,		
	Results, Conclusions, Protocol		
	reference, Revision and approval		
21.2	signatures.		
31.3	Whether OQ protocol include at		
	least System production capacity		
	(L/min), Flow type and water rate,		
	Valve operation, Alarm system		

	operation and Controls operation?		
31.4	Whether its report includes		
51.4	Conclusion / Summary, description		
	• • •		
	of the performed assay, Data tables,		
	Results, Conclusions, Protocol		
	reference, Revision and approval		
01.5	signatures.		
31.5	Please specify the water whether		
	Phase 1, Phase 2 and Phase 3 studies		
	carried out in at PQ stages?		
31.5.1	Phase 1 : Whether the operations		
	parameters, cleaning and sanitation		
	procedures & frequencies defined.		
	Whether daily sampling records for		
	every pretreatment point and usage		
	point for a period of 2 to 4 weeks		
	maintained and SOP's prepared.		
31.5.2	PHASE 2 : Whether daily sampling		
	records for every pretreatment point		
	and usage point for a period of 4 to 5		
	weeks after Phase 1 maintained and		
	reviewed.		
31.5.3	PHASE 3 : Whether weekly		
	sampling records available of every		
	usage point for a one-year period.		
	In the case of water for injections		
	systems, are the daily sampling		
	records of at least one usage point		
	available, with all the usage points		
	sampled weekly?		
	Whether results of these records		
	summarized to show suitability.		
	Are there personnel training		
	records?		
32	EQUIPMENT		
32.1	Are the equipment installation		
52.1	Qualification (IQ) protocols contains		
	followings: Introduction, Installation		
	description, Responsibilities,		
	Performed tests/assays,		
	Qualification acceptance criteria and		
	Data recording and reporting?		
32.2	Whether report contains Summary,		
32.2	Description of performed		
	tests/assays, Obtained data tables,		
	Results, Conclusions, Installation		
	diagrams, Revision and approval		
20.0	signatures.		
32.3	Whether the equipment operation		
	qualification (OQ) protocols		
	contains following: Introduction,		
	Equipment description, Description		
	of the equipment operation steps		

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	(SOP's), Responsibilities,		
	Qualification acceptance criteria,		
	Data recording and reporting.		
	Whether report contains Summary,		
	Description of performed		
	tests/assays, Obtained data tables,		
	Results, Conclusions, Revision and		
	approval signatures.		
32.4	Whether equipment performance		
	qualification (PQ) protocols contains		
	followings: Introduction,		
	Responsibilities, Performed assays,		
	Qualification acceptance criteria,		
	Data recording and reporting.		
32.5	Whether report contains Summary,		
	Description of performed		
	tests/assays, Obtained data tables,		
	Results, Conclusions, Revision and		
	approval signatures.		
32.6	Whether Preventive Maintenance		
	Schedule of the equipments is		
	followed and records available?		
33	Analytical Method Validation		
33.1	Please specify whether following		
	Characteristics are considered		
	during validation of analytical		
	methods:		
	— specificity		
	— linearity		
	— range		
	— accuracy		
	— precision		
	— detection limit		
	— quantitation limit		
	— Robustness.		
33.2	Whether Paharmocopial methods are		
	also validated. If yes, how.		
33.3	Whether system suitable testing is		
	included in testing protocols e.g.		
	HPLC, GC etc.		
33.4	Whether the procedure covers all		
	aspects of impurity profiling		
	required		
33.5	Whether procedure covers all		
	aspects of Organic Volatile		
	Impurities detection and		
	quantification		
34	CLEANING		
34.1	Is a validation performed to confirm		
	cleaning effectiveness?		
34.2	Does the protocol define the		
1	selection criteria for products or		

	groups of products subject to		
24.2	cleaning validation?		
34.3	Is data produced supporting the		
	conclusion that residues were		
24.4	removed to an acceptable level?		
34.4	Please specify whether the		
	validation is implemented to verify		
	cleaning of:		
	Surfaces in contact with the product,		
	After a change in product, Between		
	shift batches.		
34.5	Please specify whether the		
	Validation Strategy include		
	contamination risks, equipment		
	storage time, the need to store		
	equipment dry and sterilize and free		
	of pyrogens if necessary?		
34.6	Whether the cleaning Validation		
	Protocol include:		
	a. Interval between the end of		
	production and the beginning of the		
	cleaning SOP's.		
	b. Cleaning SOP's to be used.		
	c. Any monitoring equipment to be		
	used.		
	d. Number of consecutive cleaning		
	cycles performed?		
	e. Clearly defined sampling points.		
34.7	Whether Quality Control responsible		
	of the sampling for cleaning		
	verification?		
34.8	Whether personnel engaged in		
	cleaning, sampling etc. trained.		
34.9	Please specify whether acceptance		
	limits been set for cleaning		
	verification and are based on		
	following criteria:		
	a. Visually clean.		
	b. 10 ppm in another product		
	c. 0.1% of the therapeutic dose?		
34.10	Please specify whether detergent		
	residues investigated and		
	degradation products verified during		
	validation.		
34.11	Whether validation records include		
	Recovery study data, Analytical		
	methods including Detection Limits		
	and Quantification Limits,		
	Acceptance Criteria, Signatures of		
	the Quality Assurance Manager,		
	employee in charge of cleaning and		
	the verification from Production and		
	Quality Control.		
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35	Air Handling System		
35.1	Please specify whether following		
	parameters have been qualified:		
	— temperature		
	— relative humidity		
	— supply air quantities for all		
	diffusers		
	— return air or exhaust air quantities		
	— room air change rates		
	— room pressures (pressure		
	differentials)		
	— room airflow patterns		
	- unidirectional flow velocities		
	- containment system velocities		
	—filter penetration tests (HEPA)		
	— room particle counts		
	— room clean-up rates		
	— microbiological air and surface		
	counts where appropriate		
	— operation of de-dusting		
	— warning/alarm systems where		
	applicable.		
35.2	Whether strategic tests like Particle		
	count, air pressure differential, air		
	flow volume, air flow velocity etc.		
	included in Air Handling System		
26	qualification.		
36	Media fill test		
36.1	Whether medial fill tests carried out		
	twice in a year during normal		
36.2	working conditions.		
36.3	Pls give date of last such test.		
30.3	How many units are filled and tested.		
36.4	What is the criterion for		
50.4	qualification of this test?		
36.5	In case of failure of media fill test,		
50.5	what precautions or actions are		
	taken.		
37	Product Information		
37.1	Name of product		
37.2	Whether validated master formula is		
57.2	available?		
37.3	Whether specific SOP for product		
	processing is available?		
37.4	Comments on the above SOP		
37.5	Process Validation performed for the		
	product covers all aspects and the		
	approach is Risk Based		
37.6	No. of Batches Produced		
37.7	Stability studies		
51.1	(i) Accelerated		
		1	

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	(ii) Real Time		
	(iii) Whether the expiry date		
	assigned on the basis of stability		
	study?		
37.8	Whether trend analysis was carried		
	out and interpretation thereof?		
37.9	Whether Annual product review		
	(APR) is carried out? Whether the		
	following parameters considered in		
	the Annual product review?		
	1 critical in-process control and		
	critical API test results		
	2 all batches that failed to meet		
	established specification(s)		
	3 all critical deviations or non-		
	conformances and related		
	investigations		
	4 any changes carried out to the		
	processes or analytical methods		
	5 results of the stability monitoring		
	programme		
	6 quality-related returns, complaints		
	and recalls and adequacy of		
	corrective actions		
37.10	Is there any complaint received for		
	the product and If any, whether the		
	investigation report along with ATR		
	is maintained?		