REVIEW ON “GOOD MANUFACTURING PRACTICE FOR PHARMACEUTICAL PRODUCT THROUGH THE I.C.H”

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ABSTRACT
Great manufacturing hones (GMP) may be a portion of quality affirmation which guarantees that items are reliably delivered and controlled to the quality benchmarks fitting to their expecting utilize and as required by the showcasing authorization. GMP rules give least prerequisites for pharmaceutical or a nourishment item producer must meet to guarantee that the items are of tall quality and don’t posture any chance to the buyer or open. To attain the quality objective dependably there must be a comprehensively outlined and accurately executed framework of Quality Affirmation Joining Great Fabricating Hone, and hence Quality Control and Quality Hazard Administration. It ought to be completely archived and its adequacy observed. All parts of the Quality Confirmation frameworks ought to be enough resourced with competent faculty, and appropriate and adequate premises, hardware and offices. There are extra lawful duties for the holder of the fabricating authorize.

KEYWORDS: Great Fabricating Hone for pharmaceutical item, Quality control, Quality confirmation, authorized.

INTRODUCTIONS
The term GMP was presented to direct fabricating and bundling operations within the pharmaceutical industry. The Medication Reviewer of the Division of Wellbeing and Social Security of Britain, in discussion with other interested bodies compiled the direct to GMP too known as the Orange Direct. The primary version of the direct was distributed in 1971, the fabricating of sedate carried out beneath the Medications Act. It was a moderately light volume of 20 pages, and was reissue third impression in 1972, with the expansion of a 2-page reference section on sterile therapeutic items. The color of its cover, it known as the Orange Direct. The moment version (52 pages, counting five reference sections) was distributed in 1977. The third version (110 pages, five reference sections) was distributed in 1983.

The Medications and Healthcare items Administrative organization (MHRA) has distributed modern version of the Orange Direct in 2007. In Joined together States, the primary GMP directions were issued in 1963 and portrayed the GMP to be taken after within the make, bundling, and capacity of wrapped up pharmaceutical items. GMP controls were created by the US FDA and issued the Joined together States CFR Chapter 21 in 1978. The direction was comparable in concept to the Orange Direct, but enforceable by law though the UK direct as an counseling. US congress passed the Government Ani-tempering Act in 1983, making it a wrongdoing to alter with bundled customer items.[1]

This rule is expecting to assist producers actualizing present day quality frameworks and chance administration approaches to meet the prerequisites of quality items to guarantee their aiming reason and to secure the open wellbeing. The necessity in this rule is built up based on the order given to the Authority as stipulated within the Announcement Number 661/2009 for the foundation of Nourishment, Drugs and Healthcare Items in Ethiopia. GMP guarantees that quality is built into the organization and forms included within the make of the items and all those operations ought to be carried out entirely concurring to cGMP.[2]

GOOD MANUFACTURING PRACTICES (GMP) GUIDELINES
Numerous nations have enacted that pharmaceutical and therapeutic gadget companies made their possess GMP rules that compare with their enactment. Essential concepts of GMP rules objective of defending the wellbeing of the persistent as well as creating great quality medication, therapeutic gadgets or dynamic pharmaceutical items. The formalization of GMP commenced within the 1960s and their impact in over 100 nations extending from Afghanistan to Zimbabwe. Cases of these incorporate the following.
a. **Pharmaceutical Inspection Convention (PIC):** Coordinate to GMP for pharmaceutical products: Australia, Austria, Belgium, Canada, Italy, Latvia, Liechtenstein, Denmark, Finland, France, Hungary, Ireland, Malaysia, The Netherlands, Norway, Poland, Portugal, Romania, Singapore, Slovak Republic, Spain, Sweden, Switzerland, and the Joined together Kingdom.

b. **Association of South-East Asia Nations (ASEAN):** Common rules Brunei Darussalam, Indonesia, Lao PDR, Malaysia, Cambodia, Myanmar, Philippines, Singapore, Thailand, and Vietnam.

c. **European Economic Community (EEC):** Direct to GMP for restorative items Austria, Belgium, Denmark, Ireland, Italy, Luxembourg, the Netherlands, Finland, France, Germany, Greece, Portugal, Spain, Sweden, and the Joined together Kingdom. In common, GMP has been issued guides to the accomplishment of steady item quality, with elucidation and person varieties being acknowledged. GMP implemented within the Joined together States by the US FDA, beneath Segment 501(B) of the 1938 Nourishment, Sedate, and Corrective Act. The controls utilize the state “current great fabricating practices” (c GMP) and it portrays the guidelines.\(^3\)

**COMPONENTS OF GMP**

GMP requires that the fabricating handle is completely characterized some time recently being started and all the fundamental offices are given. In hone, faculty must be satisfactorily prepared, reasonable premises and hardware utilized, redress materials utilized, endorsed strategies received, appropriate capacity and transport offices accessible, and fitting records made. The basic components of GMP are summarized in Figur. 1.

![Fig. 1: Components of Great Fabricating hone.](image)

Indian arranges M for GMP and prerequisites of premises, plant and equipment for pharmaceutical things. Parcel I joins common prerequisites, Warehousing zone, Era zone, Quality control zone, Work drive, Assistant zone, Prosperity, clothing and sanitation of laborers, Manufacturing operations and controls, Sanitation inside the creating premises, Unrefined materials, Adapt, Documentation and Records, Names and other printed materials, Quality assertion, Self survey and quality survey, Quality control system, Detail, Expert condition records, Squeezing records, Bunch bundling records, Gather planning records, Standard working strategies (SOPs) and records, Reference tests, Reprocessing and recuperations, Spread records, Endorsement and handle endorsement, Thing audits, Complaints and unfavorable reactions and Site-master record. Parcel I-A to parcel I-E takes note nearly the specific prerequisites for create of different things and Parcel I-
QUALITY MANAGEMENT
As it were the holder of a fabricating authorisation must fabricate pharmaceutical items so as to guarantee that they are fit for their aiming utilize, comply with the necessities of the Promoting Authorisation and don't put the client at chance due to lacking security, quality or viability.

The achievement of this quality objective is the obligation of senior administration and requires the support and commitment of staff working completely different offices of the company. To attain the quality objective dependably there must be a comprehensively planned and accurately actualized framework of Quality Confirmation joining Great Fabricating Hone, Quality Control and Quality Chance Administration. It ought to be completely recorded and its adequacy checked. All parts of the Quality Confirmation frameworks ought to be enough resourced with competent work force, and appropriate and adequate premises, hardware and offices. There are extra legitimate duties for the holder of the fabricating authorisation and for the authorized person(s).

The essential concepts of Quality Confirmation, Great Fabricating Hone, Quality Control and Quality Hazard Administration are inter-related. They are portrayed here in arrange to accentuation their connections and their crucial significance to the generation and control of pharmaceutical products.\[^5\]

The holder of a fabricating authorization must fabricate therapeutic items so as to guarantee that they are fit for their expecting utilize ,comply with the necessities of the showcasing authorization and don't put patients at chance due to insufficient security, quality or viability. The achievement of this quality objective is the obligation of senior administration and requires the support and commitment by staff in numerous diverse offices and at all levels inside the company, by the company providers and by the distributors.\[^7\] Within the pharmaceutical industry at expansive, quality administration is more often than not characterized as the angle of administration work that decides and actualizes the “quality policy”, i.e. the generally purposeful and course of an organization with respect to quality, as formally communicated and authorized by beat administration.\[^6\]

QUALITY ASSURANCE
Quality Confirmation may be a wide-ranging concept, which covers all things, which separately or collectively impact the quality of a item. It is the whole add up to of the organized courses of action made with the objective of guaranteeing that therapeutic items are of the quality required for their aiming utilize. Quality Confirmation subsequently consolidates Great Fabricating Hone additionally other components exterior the scope of this Direct. The holder of a fabricating authorization must fabricate restorative items so as to guarantee that they are fit for their aiming utilize, comply with the requirements of the promoting authorization and don't put patients at hazard due to lacking security, quality or adequacy. The fulfillment of this quality objective is the duty of senior administration and requires the cooperation and commitment by staff in numerous distinctive divisions and inside the company, by the company' providers and by the wholesalers. QA may be a wide extending concept, which covers all matte.\[^7\]

The framework of QA fitting for the fabricate of pharmaceutical items ought to guarantee that:

a. Pharmaceutical items are outlined and created in a way that takes account of the prerequisites of GMP and other related codes such as those of great research facility hone (GLP) and great clinical hone (GCP).
Generation and control operations are clearly indicated in a composed frame and GMP necessities are embraced. Courses of action are made for the make, supply and utilize of the proper beginning and bundling materials. All essential controls on beginning materials, middle of the road items, and bulk items and other in process controls, calibrations, and validations are carried out. The wrapped up item is accurately prepared and checked, concurrently to the characterized methods, pharmaceutical items are not sold or provided some time recently the authorized people have certified that each generation clump has been delivered and controlled in agreement with the prerequisites of the promoting authorization and any other controls significant to the generation, control and discharge of pharmaceutical items. Palatable courses of action exist to guarantee, as distant as conceivable, that the pharmaceutical items are put away by the producer, disseminated, and along these lines taken care of so that quality is kept up all through their shelf-life; Deviations are detailed, examined and recorded. Standard assessments of the quality of pharmaceutical items ought to be conducted with the objective of confirming the consistency of the method and guaranteeing its ceaseless advancement.[9]

Good manufacturing practice - the general/current state

Pharmaceutical quality system
This run the show depicts a comprehensive appear for an practicality quality system of helpful things, based on the concepts of ISO quality and its utilization all through all stages of the life cycle of the thing. This run the show will improvement the mechanical improvement and invigorating of the interface between pharmaceutical change and creating works out. The run the show applies to supporting the headway and create of substances of Pharmaceutical Industry, Energetic Pharmaceutical Settling and therapeutic things, tallying biotechnology and natural things all through the life cycle of the thing.

Quality affirmation can be a wide concept that joins all things that only or collectively impact the quality of a thing, that’s, organization of the quality of rough materials, things and other components, organizations related to era, and organization, era and evaluation shapes. It is associated in pre-production to confirm what will be made meets judgments and necessities conjointly while manufacturing era. Two measures included in quality affirmation are: fit for purpose where the thing got to be sensible for the aiming reason; and “right to start with time” botches need to be unequivocally arranged of. In orchestra to accomplish quality, there must be a system of comprehensive quality confirmation and actualized it precisely. This last issue consolidate the organization of GMP, quality control and quality chance.[10-14]

Personnel
Concurring to EC GMP, the administration of an venture ought to decide and give satisfactory and fitting assets such as human assets, budgetary, materials, offices and gear to actualize and keep up the Quality Administration Framework and move forward viability. Compelling coordination and administration of human assets are key variables within the appropriate working of any venture. To this conclusion, endeavor administration has obligations and duties in staff enrollment as well as the appointment of errands.[15,16]

Production
Generation operations must clearly take after the methods. They must comply with the standards of GMP in arrange to get quality items and be in understanding with the significant fabricating and MA. Generation ought to be performed and directed by competent individuals. All dealing with of materials and items, such as gathering and isolate, testing, capacity, labeling, apportioning, preparing, bundling and dissemination ought to be exhausted understanding with composed strategies or informational and where fundamental, recorded.[17-19]

Quality control
Quality control is concerned with testing, details and testing as well as the organization, documentation and discharge methods which guarantee that the desired and relevant tests are carried out, which materials are not discharged for utilize, nor items discharged for deal or supply, until their quality has been judged palatable. Quality control isn't confined to research facility operations, but must be included in all choices that will concern the quality of the item. The autonomy of quality control from generation is considered principal to the palatable operation of quality control.[20-23]

Manufacture of medicinal products
Manufacture of solid and semi-solid medicinal products
Since this sort of restorative items is especially vulnerable to microbial contaminants and other contaminants amid fabricating, it is vital to take after preventive methods and it ought to be a need for the producer MA holder.[24,25]

Manufacture of sterile medicinal products
The make of sterile items requires extraordinary necessities in arrange to play down dangers of microbiological defilement, and of particulate and pyrogen defilement, being profoundly subordinate on information, preparing and demeanor of the work force included. This sort of make must entirely take after strategies and arrangement forms, carefully built up and approved, since the quality affirmation, is of specific significance. The make of sterile items ought to happen in clean ranges whose get to must be accomplished through airtight chambers for faculty and/or hardware and materials.[26-31]
Manufacture of biological active substances and medicinal products for human use

The strategies utilized within the make of natural dynamic substances and organic restorative items for human utilize are basic components in forming the fitting administrative control, since the fabricate of these includes certain particular, contemplations emerging from the nature of items and fabricating forms, being fundamental take a few extraordinary safeguards. Not at all like routine therapeutic items, which are regularly created and controlled utilizing reproducible chemical and physical strategies, organic items are made through strategies that include organic forms and materials, such as development cells or extraction of fabric from living life forms. These natural forms may show characteristic changeability and consequently, that the run and nature of the by-products may be variable. [32-34]

Manufacture of radiopharmaceuticals

Radiopharmaceuticals must be fabricated and taken care of with a few uncommon care. The administrative methods fundamental for the control of radiopharmaceuticals are decided in expansive portion by the sources of these items and the generation strategies. The level of hazard depends basically on the sorts of radiation radiated and the half-lives of radioactive isotopes. It is vital to pay extraordinary consideration to the cross defilement, the contaminant of radioactive isotopes and to squander transfer. Due to its brief half-life, a few radiopharmaceuticals are discharged and managed to the patients after their generation, some time recently completing all quality control tests. The EC rule is connected to fabricating strategies utilized by mechanical producers, Atomic Centers/Institutes and positron emanation tomography (PET) Centers for the generation and quality control of the taking after sorts of items: Radiopharmaceuticals, PET radiopharmaceuticals, Radioactive Antecedents for radiopharmaceutical generation and finally Radionuclide Generators.

The Universal Nuclear Vitality Organization is the office that directs the transport of radiopharmaceuticals and the necessities for assurance against radiation. [35,36]

Medicinal products derived from human blood or plasma

The necessities for the collection, testing and quality control of restorative items determined from human blood or plasma are characterized by a framework of quality confirmation, based on the presence of a national structure which is free of producers, complying with the standards and rules of GMP. Those items are considered to be organic restorative items due to their particular characteristics and the beginning materials incorporate organic substances, such as cells or liquids (counting blood or plasma) of human root. The arrangements of CE apply to restorative items inferred from human blood or plasma, fractionated in or imported into the EU/EEA. Besides, apply to the beginning fabric (e.g., human plasma) for these items and for steady subordinates of human blood or human plasma (e.g., egg whites) joined into restorative gadgets. In any case, it does not apply to blood components expecting for transfusion. [37,38]

Manufacture of herbal medicinal products

The methods and procedures utilized within the make and quality control of home grown drugs are regularly considerably distinctive from those utilized for customary therapeutic items. The home grown substance ought to be of appropriate quality. The supporting information ought to be given to the producer of the home grown preparation/herbal therapeutic item. A steady quality confirmation of home grown substances requires more point by point data on its agrarian production.

These rules apply to all home grown beginning materials: Restorative plants, home grown substances or home grown preparations. [39,40]

GOOD MANUFACTURING PRACTICE FOR MEDICINAL PRODUCTS (GMP)

Great Fabricating Hone is that portion of Quality Affirmation which guarantees that Restorative items are reliably delivered and controlled to the quality guidelines suitable to their expecting utilize and as required by the promoting authorization or item detail. Great Fabricating Hone is concerned with both generation and quality control. GMP is pointed essentially at lessening the dangers inalienable in any pharmaceutical generation. [41]

The fundamental prerequisites of GMP are that:

- all fabricating forms are clearly characterized, efficiently surveyed within the light of involvement and appeared to be able of reliably manufacturing medicinal items of the desired quality and complying with their details;
- critical steps of fabricating forms and noteworthy changes to the method are approved;
- all essential offices for GMP are given counting:
  a. fittingly qualified and prepared work force;
  b. satisfactory premises and space;
  c. reasonable gear and administrations;
  d. redress materials, holders and names;
  e. affirmed methods and enlightening;
  f. reasonable capacity and transport;
- iv. Enlightening and methods are composed in an guidelines shape in clear and unambiguous dialect, particularly pertinent to the offices given;
- vi. Records are made, physically and/or by recording disobedient, amid fabricate which illustrate that all the steps required by the characterized strategies and enlightening were in reality taken which the amount and quality of the item was as expected9. Any noteworthy deviations are completely recorded and explored;
- vii. Records of fabricate counting dissemination which empower the total history of a clump to be followed, are held in a comprehensible and available shape;
viii. The dispersion (wholesaling) of the items minimizes any hazard to their quality; ix. A framework is accessible to review any clump of item, from deal or supply; x. complaints almost showcased items are inspected, the causes of quality absconds examined and fitting measures taken in regard of the imperfect items and to avoid re-occurrence.[42]

Sanitation and Hygiene

Tall level or sanitation and cleanliness ought to be practiced in each viewpoint of fabricating pharmaceutical items. The scope of sanitation and cleanliness covers work force, premises, hardware and device, generation materials and holders and anything that seem gotten to be a source of defilement to the item. Potential sources of defilement ought to be dispensed with through an coordinates comprehensive program of sanitation and cleanliness.[43]

Each individual entering the fabricating regions ought to wear defensive pieces of clothing suitable to the operations to be carried out.

Individual cleanliness strategies counting necessity of utilizing defensive clothing ought to apply to all people entering generation zones, whether they are transitory or full-time representatives or non-employees on company property, e.g. contractor’s representatives, guests, senior administration and auditors.

To guarantee assurance of the item from contaminations as well as the security of the staff, they ought to wear clean body-coverings suitable to the obligations they perform, counting suitable hair covering. Dirty regalia and dirtied cleaning cloths (in case reusable) ought to be put away in isolated closed holders until appropriately laundered.

Point by point cleanliness programs ought to be set up and adjusted to the distinctive needs inside the fabricating range. They ought to incorporate strategies relating to the wellbeing, cleanliness hones and clothing of staff. These methods ought to be understood and taken after in a really strict way by each individual whose obligations take him into the generation and control ranges. Cleanliness programs ought to be advanced by administration and broadly talked about amid preparing sessions.

All staff ought to get restorative examination upon enrollment. It must be the manufacturer's duty that there are informational guaranteeing that wellbeing conditions that can be of significance to the quality of items come to the manufacturer's information. After the primary therapeutic examination, examinations ought to be carried out occasionally for the work and individual wellbeing. Visual auditors ought to moreover experience occasional eye examination.

The ranges, surfaces, and hardware in and on which items are made must be kept clean. Earth, and the organisms that it can harbor, must not get into or on items. Disinfectants can be inactivated by earth. Soil (especially sleek or oily movies and protein like matter) can too secure microorganisms against the activity of disinfectants. So, some time recently sanitization, it is critical to to begin with clean surfaces. Where net sums of earth are show, it may be essential to to begin with evacuate most of it by scouring. At that point surfaces may be cleaned by the application of a cleaning specialist, taken after by rising.[44]

Validation

Approval is characterized as the setting up of recorded prove which gives a tall degree of affirmation that a arranged prepare will reliably perform concuring to the expecting indicated results. Approval thinks about ought to strengthen GMP and be conducted in agreement with characterized strategies. Comes about and conclusions ought to be recorded. When any unused fabricating equation or strategy of planning is embraced, steps ought to be taken to illustrate its reasonableness for schedule handling.[45]

Capability of frameworks and gear is subsequently a portion of handle of approval. It may be a necessity of nourishment and sedate, pharmaceutical directing organizations like FDA’s rules. Since a wide variety of methods, forms, and exercises have to be approved, the field of approval is separated into a number of subsections counting the taking after:
• Gear approval
• Offices approval
• HVAC framework approval
• Cleaning approval
• Prepare Approval
• Expository strategy approval
• Computer framework approval
• Bundling approval

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• Expository strategy approval
• Computer framework approval
• Bundling approval

Premises

Premises and gear must be found, planned, developed, adjusted and kept up to suit the operations to be carried out. Their format and plan must point to limit the chance of blunders and allow compelling cleaning and upkeep in arrange to maintain a strategic distance from cross-contamination.

Premises must be found, developed, adjusted, planned, and kept up to suit the operations to be carried out. The format and plan of premises must point to play down the hazard of mistakes and allow successful cleaning and upkeep in arrange to maintain a strategic distance from cross defilement, build-up of clean or earth, and in common, any antagonistic impact on the quality of products15. The choices of materials of development for fabricating offices are various. A few cases are displayed hence.[46]
CONCLUSION

GMP could be a generation and testing hone that makes a difference to guarantee in built quality item. Numerous nations have enacted that pharmaceutical companies must take after GMP strategies, and have made their claim GMP rules that compare with their enactment. Fundamental concepts of all of these rules stay more or less comparable to the extreme objectives of shielding the wellbeing of the understanding as well as creating great quality medications.

Quality objective can be accomplished as it were through cautious arranging and usage of QA framework and down to earth usage of GMP. The compelling usage of GMP requires broad care and information around the diverse components of GMP that ought to be consolidated frame the initiation of the fabricating building and item improvement till the production.

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a. Dividers: Dividers in fabricating regions, bundling ranges and hallways ought to be of mortar wrap up on high-quality concrete pieces or gypsum board. The wrap up ought to be smooth, ordinarily with finish or epoxy paint. They ought to be launder able and able to stand up to rehashed applications of cleaning and sanitizing operators.

b. Floors: Floor covering ought to be chosen for strength as well as for clean capacity and resistance to the chemicals with which it is likely to come into contact. Epoxy flooring gives a strong and promptly cleanable surface.

c. Ceilings: Suspended ceilings may be given in office zones, toilets, research facilities and cafeterias. They more often than not comprise of lay-in acoustical boards of non delicate, non friable, non asbestos and non combustible fabric. Fabricating zones require a smooth wrap up, regularly of consistent mortar or gypsum board. All ceiling installations such as light fittings, discuss outlets and returns ought to be planned to guarantee ease of cleaning and to play down the potential for amassing of clean.

d. Administrations: Within the building plan, arrangements must be made for channels, steam, power, water and other administrations to permit for ease of support. Get to ought to, in a perfect world, be conceivable without disturbance of action inside the real rooms given with the administrations. Entryways and window frames ought to all have a difficult, smooth, impenetrable wrap up, and ought to near firmly. Window and entryway outlines ought to be fitted flush, at slightest on sides confronting internal to handling ranges. Entryways, but crisis exits, ought to not open specifically from generation regions to the exterior world. Any crisis exit entryways ought to be kept closed and fixed, and outlined so as to be open able as it were when crisis requests. [47]

Equipment

Fabricating hardware ought to be able of creating materials, items and intermediates that are expecting which conform to the desired or indicated quality characteristics. The gear must be planned and built so that it is conceivable to clean it altogether. Surfaces that come into contact with items ought to have cleaned wraps up, smooth with no breaks, hole, troublesome corners, uneven joints, dead-legs, projections, or unpleasant welds to harbor defilement or make cleaning troublesome. Hardware must too be competent of withstanding rehashed, exhaustive cleaning. Follows of past item, at levels that may well be acceptable in other businesses, are completely unsatisfactory within the fabricate of solutions. Between clumps all fabricating gear must be completely cleaned and cleaned or sterilized. [48]


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