Solid Dosage Forms: A Detailed Research on Non-conforming Product Quality

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ABSTRACT

Introduction: Any defects in pharmaceutical products lead to minor, major, or critical deviation and if such a product is released, it also leads to product recalls and legal actions. In such cases, the concern is about patient’s safety rather than the company’s economy and fame. This study aimed to investigate the various quality defects which have occurred in marketed tablets. Materials and Methods: The investigation was carried out which involved identification of defects, categorizing the type, root causes, suggesting remedial measures and documenting the clinical significance of defective dosage forms of tablets. Dosage forms included were film-coated tablets, uncoated tablets, uncoated caplet shaped tablets and dispersible tablets which are prescribed for their specific therapeutic activity. Results: This research work has revealed quality issues of minor, major and critical types. These defects were unique in nature and appeared to be stray incidents which were not available even in standard textbooks or literature. Conclusion: Medicines being the silver sword to treat an ailment in a patient, it should not involve any quality issues leading to a significant impact on the safety and efficacy of the prescribed drug. It is essential to embed an attentive and dedicated quality assurance (QA) team and adhere to all the regulations laid by regulatory agencies in the pharmaceutical setup thereby ensuring that high-quality medicines are supplied and the safety of the patient is shielded.

Key words: Tablets, Solid dosage forms, Tablet defects, Patient perception, Investigation, Quality Assurance.

INTRODUCTION

The drug is defined as a substance recognized by official pharmacopoeia / In house (IH) which, is intended for its use in the diagnosis, cure, mitigation, treatment, or prevention of disease. Rarely drug is given in its pure chemical form. To ease the drug administration by a human being, it is essential to convert it into physical form in which drug is dispensed known as dosage form.1,2 The dosage form is a package of Active Pharmaceutical Ingredient (API) along with selective non medicinal compounds known as excipients. These non-medicinal compounds may be used to flavor, solubilize, color, preserve, dilute, emulsify, suspend and thicken medicinal agents into efficacious and appealing dosage forms.3 Besides dispensing the drug into physical form, the dosage form is needed for the following additional reasons:

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Different Types of Packaging

Based on the level of contact of formulation with the container

1. **Primary packaging**: This packaging material is in direct contact with the formulation. Hence it is mandatory to ensure that packaging material doesn't interact with the drug. E.g. blister packs, strip packs, containers of liquid dosage forms, etc.

2. **Secondary packaging**: This packaging type is in contact with primary packaging, keeping multiple units of products in place during transportation. E.g. 1-ply, 2-ply and 3-ply corrugated boxes.

3. **Tertiary packaging**: This packaging material conceals the above two units of packed products and is in direct contact with secondary packaging. E.g. shrink wraps, plain boxes, cardboards.

Usually, the secondary package is a composite of primary packaging material with the product and Patient information leaflet (PIL). PIL is a document added by the manufacturer which gives information about clinical indications of the drug, type of dosage form, route of administration, storage condition, adverse drug reactions, contraindications and details of the manufacturer.

Packaging types used for tablets

a. **Blister packaging**: WHO defines a blister pack as a multi-dose container consisting of two layers, of which one is shaped to contain the individual doses and is heat-sealed with other layer which may be aluminum, paper, or PET (Polyethylene terephthalate).

b. **Strip packaging**: As per WHO strip pack is defined as A multi-dose container consisting of two layers, usually provided with perforations, suitable for containing single doses of solid or semi-solid preparations. The two layers are made of heat-sealable paper alloy, aluminum films, glassine.

Both above-mentioned packaging types involve sealing of individual doses, which requires breaking or tearing of individual compartment before administration.

The major difference between the above two types is, the former has cavities on thermoplastic which are preformed and the tablet is dropped before closing the cavity. Whereas in later, the tablet is dropped in between the two layers in which it has to be embedded and sealed.

Packaging materials used for tablets

1. **Polyvinyl Chloride (PVC)**: PVC is highly moisture resistant and is available in different gauges. It can be transparent or can be made opaque or can be tinted.
in different colors to block specific wavelengths of light. It is the most commonly used blister material because of its affordability and its characteristics like flexibility, thermoforming and rigidity.\textsuperscript{11}

2. Polychlorotrifluoroethylene (PCTFE) laminations: It is thermoplastic manufactured by modification of polyethylene (PE). It is fixed to PVC by the aid of adhesive.

3. Aluminum: The various combination of packs is formed by combining. E.g. Alu-Alu, Aluminum-paper, aluminum-PET. Aluminum is widely used in strip packs and as lid material in the blister pack.

4. Cellulose polymers: Are the main components of paper-based packs. Based on the concentration pulp they are used as lid material for Aluminum or PVC blister packs

These are the materials used in the packaging of tablets. However, the manufacturer cannot use material of his choice to prepare containers or packages for any dosage form. The list of materials published by the FDA (Food and Drug Administration) which are “Generally regarded as safe (GRAS)” have to be used in packaging material. If the manufacturer intends to use a material that is out of the GRAS list, he has to conduct tests for the intended material and file a New Drug Application to FDA.\textsuperscript{10,12-14}

Defects in Tablet manufacturing

Figure 3 gives the classification of Defects based on their occurrence in coated and uncoated tablets. The tablet manufacturing process involves several unit operations from which defects in tablets arise. Though there are significant innovative solutions that have come up in the manufacturing process there is no decrease in cases of tablet defects.\textsuperscript{15,16} Both to the manufacturer (e.g. simplicity and economy of the preparation, stability, and convenience in packing, shipping and dispensing

Few major defects have been discussed below:

1. Capping
It is identified when there is partial or complete separation of the top or bottom crowns of tablets

Causes:
- Deep concave punches and poorly finished dies
- Over lubrication or very low moisture content of granules
- High compression pressure
- Deformational properties of ingredients
- Incorrect set up of the tableting machine

Remedies:
- Pre-compression and decreasing the compression pressure
- Ensure proper setup of tableting machine.\textsuperscript{14,16}

2. Cracking
In Uncoated tablets: Cracking can be described as small cracks that occur in upper and lower surfaces of tablets.

Causes:
- Due to usage of large size and dry granules
- Due to entrapment of air in granules during punching

Remedies:
- Reduce the size of granules, incorporate fines along and maintain the standard amount of moisture in granules
- Use tapered die.

In Coated tablets: Cracking is a defect in which cracks occur along with the crown of tablet or splits along the edges of a tablet (Splitting).

Causes:
- Due to internal stress developed within film after tablet dries
- Usage of high molecular weight polymer for tablet coating

Remedies:
- Implementation of post compaction relaxation phenomena after tablet punching
- Addition of plasticizer into coating material thereby increasing the flexibility of tablet coat.\textsuperscript{14,16,17}

3. Orange peel effect: It is a surface-based defect. The coated tablet looks rough and non-glossy.

Causes:
- Premature drying of atomized droplets or spraying of a viscous coating solution
- Tablets subjected to a rapid drying process

Remedies:
- Decreasing the viscosity of coating solution by adding other solvents
- Subject tablets to mild drying conditions.\textsuperscript{16,17}

Our mankind involves day to day activities which involve sectors like architecture, agriculture, fashion, transport, telecommunication and pharma. Defects in none of the sectors are tolerated. In the same lines, defects in any pharmaceutical product may be fatal. Although Pharmaceutical sector is constrained by stringent regulations laid by Regulatory agencies, (United States Food and Drug Administration, European Medicines Agency, Central Drugs Standard Control Organization, Therapeutics Goods Administration and Medicines
and Healthcare Regulatory Agency) defects have been occurring which may have a clinical effect on the patient and economical significance on the manufacturer.

In this article, we present our case studies which involve research on product defects or quality issues. Our research work involved a unique in-house mechanism that started from the identification of a product, procurement of products and its detailed study. The following are the case studies/defects and based on the detailed investigation we have identified the category of complaint, probable root causes, remediation and clinical significance of complaint. Table 1. Gives an overview of dosage forms and investigated defects.

**CASE STUDY 1 – Investigation of broken film coated tablets**

**Dosage form:** Film coated tablets.

**Generic Name:** Flavonoids.\(^{18}\)
Defects: Broken film coated tablets are found within intact blister pocket (Figure 4).

Category of complaint: Major

Probable root causes:
- Low hardness of tablets.³
- Improper machine setting during packaging.
- Improperly designed packaging change parts having a very low clearance of the pocket.
- Selection of inadequate packing material – film former or lidding foil leading to high moisture uptake.²⁰
- Inadequate training to operators involved in packing.

Table 1: Overview of Investigated defective products.

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Dosage form</th>
<th>Generic name</th>
<th>Defect</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Film coated tablets</td>
<td>Flavanoids</td>
<td>Broken film coated tablets within intact blister pocket</td>
</tr>
<tr>
<td>2</td>
<td>Uncoated tablets</td>
<td>Levocetrizine hydrochloride</td>
<td>Empty blister pocket</td>
</tr>
<tr>
<td>3</td>
<td>Uncoated tablets</td>
<td>Glibenclamide and Metformin hydrochloride</td>
<td>Presence of Hair follicle within two intact blister pockets</td>
</tr>
<tr>
<td>4</td>
<td>Film coated tablets</td>
<td>Desloratadine</td>
<td>Soiled and powders adhered to tablets</td>
</tr>
<tr>
<td>5</td>
<td>Uncoated caplet shaped tablets</td>
<td>Paracetamol and Aceclofenac sodium</td>
<td>Spoiled and fragile tablet</td>
</tr>
<tr>
<td>6</td>
<td>Dispersible tablets</td>
<td>Lactic acid bacillus, Folic acid and Vitamin B₁₂</td>
<td>Empty blister sheet</td>
</tr>
</tbody>
</table>

Therapeutic category: Phlebotropic/Venoactive drug.³⁹
Remediation

- To check whether the tablet meets the required hardness specifications. Improve tablet formulation leading to improved hardness if found low.

- Review the film former and lidding foil quality with respect to its Water Vapor Transmission Rate (WVTR) and propensity of the lidding foil to have pin holes. Use film former with better barrier properties and higher thickness of lidding foil.

- Increase the pocket depth by getting new dies for the film formation.

- Recalibrate recipe of the pocket formation steps by increasing film-forming temperature and vacuum.

- Impart training to packaging personnel to adhere to the prescribed machine settings and calibrating the temperature and vacuum sensors.

Clinical Significance

Reduced patient compliance. Altered therapeutic efficacy due to the degradation of API. Reduced bioavailability due to altered dissolution. The stability of active ingredient may be affected due to exposure of the product to low gastric pH.

CASE STUDY 2 – Investigation of empty blister pocket

Dosage form: Uncoated tablets

Generic Name: Levocetirizine hydrochloride tablets

Therapeutic category: Selective histamine H1 receptor antagonist, with anti-inflammatory and potential anti-angiogenic activities

Defects: Empty pocket found in intact blister (Figure. 5).

Category of complaint: Minor

Probable root causes

- Inadequate in-process quality checks during primary packing
- Camera based inspection was not carried out

Remediation

- Implementation of automatic check weigher to reject underweight blisters.
- Camera based visual check to reject blisters with empty pockets.
- Cleaning of tablet delivery chutes to the blister to ensure uniform flow of tablets to the blisters.

Clinical Significance: Patient acceptance decreases and the patient may discontinue purchasing the same brand again.

CASE STUDY 3 – Investigation of the tablet with hair follicle

Dosage form: Uncoated tablets

Generic Name: Glibenclamide and Metformin hydrochloride tablets

Therapeutic category: Antidiabetic

Defects: Hair follicle found within TWO intact blister pockets (Figure 6a & 6b).

Category of complaint: Critical

Probable root causes

- Staff not wearing hand gloves.
- Inadequate training on gowning procedure
- Inadequate environmental control.
- Use of poor quality of film former and/or quality check.

Remediation

- Strict adherence to GMP (Good manufacturing practice) systems.
Ensure isolation of personnel during blister packaging operation from the equipment.

- Refresh training on proper gowning procedures for personnel.
- Ensure that the area where blister packaging operation is taking place is under positive pressure compared to the personnel area.
- Review/inspect the quality of the film-forming foil so that the same is free of any extraneous materials.
- Ensure check on the quality of incoming materials before the release of the packaging material.
- Ensure that the storage area of the film formers is having same degree of personnel control as the manufacturing area.
- Reassess vendor qualification to ensure that the film supplying vendor is having the same degree of environmental control as a drug product manufacturer/formulator.

### Clinical Significance

Reduced patient compliance. The presence of hair follicle in the tablets may affect the therapeutic efficacy of the drug due to contamination (hair follicle, dandruff, etc.), psychological issues, or non-adherence.

#### CASE STUDY 4 – Investigation of soiled film coated tablet

**Dosage form:** Film coated tablets  
**Generic Name:** Desloratadine tablets  
**Therapeutic category:** Antihistamine

**Defects**

- a) The film coated tablets are soiled.
- b) Powders adhered to film coated tablets. (Figure 7)

**Category of complaint: Major**

**Probable root causes**

- Damaged film coated tablet got packed in the blister.
- Improper inspection activity.
- Lower hardness of the tablet leading to highly friable tablets.
- Very high coating solution flow rate leading to part of the tablet getting disintegrated.
- Improper / inadequate coating.

**Remediation**

- Review the manufacturing process to ensure that minimum tablet hardness is ensured before starting film coating operation.
- Weight check of individual blister before cartoning.
- Friability of the tablet to be ensured at less than 1%.  
- Monitor coating solution flow rate and bed temperature during coating operation.
- 100 % inspection of coated tablets for rejecting such defective tablets.

**Clinical Significance:** Patient acceptance may decrease. The soiled tablet may alter therapeutic efficacy due to the degradation of API. Reduced bioavailability due to altered dissolution. Patients may consume the contaminated tablet unknowingly leading to adverse events.

#### CASE STUDY 5 – Investigation of blister containing spoiled tablets

**Dosage form:** Uncoated caplet shaped tablets  
**Generic Name:** Uncoated tablets containing paracetamol and aceclofenac sodium  
**Therapeutic category:** Anti Pyretic and Anti-inflammatory agent

**Defects:**

- a) A spoiled uncoated tablet found inside intact amber color blister pack.
- b) The patient complained of a tablet getting broken while taking it out from blister and bad appearance (Figure 8a & 8b).

**Category of complaint: Critical**

**Probable root causes**

- Improperly dried granules leading to high/low moisture content.
- Manufacturing process error – inadequate granulation.
- Change of grade of API and/or binder.
- Inadequate compression setting.

**Discussion**

The probable root cause is inadequate granulation and/or mixing. The moisture content has to be very tightly controlled for paracetamol tablets to achieve required tablet hardness. Low/high moisture content of the tablet can lead to loss of hardness. There can also change of grade of API/binder as the tablets were discolored. Paracetamol supports microbial growth. A high-water content can lead to the promotion of microbial growth which in turn can cause discoloration. Paracetamol is a highly incompressible API. The use of a pre-compression stage before the compression stage can alleviate the problem to a certain extent.

**Remediation**

- Review the manufacturing process for the batch and the grade of API/excipient used and loss on drying (LOD) of the tablets. Implement better manufacturing controls especially during granulation and drying. Review the grade of the API used.
- Adherence to GMP to ensure adequate cleaning before manufacturing to assure that basic cleanliness...
of the equipment is followed to prevent microbial growth.\textsuperscript{30}  

- Evaluate the compression technique with a pre-compression facility to reduce challenges due to the incompressible nature of the API.

**Clinical significance:** Altered therapeutic efficacy due to the degradation of API. Reduced bioavailability due to improper granulation and spoiled medicine.\textsuperscript{42}

**CASE STUDY 6 – Investigation of empty blister sheet**

**Dosage form:** Dispersible tablets\textsuperscript{37}

**Generic Name:** Lactic acid bacillus,\textsuperscript{43} Folic acid\textsuperscript{44} and preconceptional intake of dietary folic acid (FA and Vitamin B\textsubscript{12})\textsuperscript{45}  

**Therapeutic category:** Anti-diarrhoeal, habitual constipation, flatulence and megaloblastic anemia\textsuperscript{46}  

**Defects:** Absence of tablets in blister pockets (Figure 9).

**Category of complaint:** Minor

**Probable root causes:**
- Blockage of discharge chute into the blister pockets\textsuperscript{37}
- Empty hopper processed along with packaging
- Inadequate QC (Quality control) check before loading into secondary packaging material

**Remediation**
- Proper steps need to be implemented after line clearance.
- Camera to be installed to identify such blisters and 100% visual inspection of primary packing can also be performed.
- Adequate checks need to be taken before packing into secondary packaging material

**Clinical Significance**

This will not have any significant impact on the patient. However, the patient may purchase and witness this defect after going to the house while taking the dose. This will be a problem for geriatric patients, as they have to visit the pharmacy shop again for replacing this blister.

To summarize, apart from listed defects or routine defects in tablet manufacturing as quoted in this article, we can positively assume that various types of quality issues can be encountered in marketed brands as well. Surprisingly, we encountered defects like broken and soiled film coated tablets, spoiled tablets, hair follicle with tablets and empty pocket, or empty blister. The observations were of a minor, major and critical types of defects. These may be stray incidents, but all the defects should be an eye-opener for manufacturing firms and are learning for improvising the manufacturing systems.

**Patient perception about quality complaints**

A drug is not only a chemical compound formed from the right blend of different elements but also trust, motivation builder for the consumer (patient). The drug is purchased with the faith that it would cure the patient of the ailment, or it would lead to better immunity. Like all manufacturing products, drugs also are subject to errors due to poor raw material quality, machine errors, or human errors. These errors can be summarized as medication errors and can have effects ranging from harmless and temporary to dangerous and permanent on the customer’s mind.\textsuperscript{48,49} Little information is available on precise costs of medication errors. This study estimated the cost of medication errors reported by clinical pharmacists using a modified societal perspective. METHODS: Information on 779 medication errors was collected in the Medication Error Detection, Amelioration and Prevention (MEDAP).

The errors can be like an empty pocket in the blister and the consumer may point it out to the pharmacist and tell them to be careful, he may end up purchasing the drug again. But if the errors are of the type of broken tablets, or a hair follicle found in the tablet, then the impact may be high. The consumer may consume the tablet, which could have either been exposed to nature or may be contaminated. We need to understand the consumer attitude towards pharma products here. Research shows that influencers, reliability, awareness, corporate image and promotion are the five factors responsible for the purchase of OTC (Over the counter) pharmaceutical products.\textsuperscript{50} Awareness, reliability and corporate image form the attitude of the customer towards a brand. Post consumption, if there is a dissatisfaction/discomfort to the consumer, then a negative attitude is formed, leading to a change in consumer behavior towards that brand. This change will negatively affect the future consumption of the same drug. It is understood that attitudes do not change over time and they also bring consistency in the behavior. Moreover, this attitude is formed due to the cognitive component, i.e., the consumer’s knowledge and experience, hence it may have a long-lasting impact on the consumer.

This attitude leads to a formation of perception called perceptual defense. Consumer perceptions are usually based either on prior personal experiences or reports of the previous experiences of others. The consumer may abandon the purchase of other drugs from the same brand/manufacturer. The consumer may start negative word of mouthing for the drug and/or manufacturer, which can lead to a dip in sales.\textsuperscript{51,52} The consumer may complain to the pharmacist and the doctor, leading to a
lower prescription of the drug by the doctor and further less/no purchase by the pharmacist. This fall in sales can also lead to legal hurdles to the company, which will result in drug withdrawals, penalty, blacklisting of the company, loss of goodwill and financial losses also. Also, the impact of mistakes may be fatal with a geriatric patient as their immunity levels are less

A recent market report by KPMG points out that the consumer healthcare industry is growing fast and expected to touch 18 to 20 USD billion. If pharmaceutical companies wish to encash this segment, then they will have to bring in zero-defect drugs for the consumers.

**CONCLUSION**

Regardless of the therapeutic use of medicine, none of the patients prefers a defective product to be purchased or expects any quality issue. The pharmaceutical industry deals with the life of human beings; the occurrence of defects in medicines is least anticipated by anyone working in the healthcare system. Tablets are the widely used dosage forms due to their affordability, stability and other user-friendly considerations. To prevent defects, the utmost care has to be taken during its genesis which starts from the procurement of API until it goes into primary packaging material. Just release of regulations by international agencies like US-FDA, WHO, ICH, EMA, etc. is not enough to ensure that quality and safety medicines are being manufactured and served to patients.

It is important to implement Good Manufacturing Practices and abide by rules and regulations laid by them. Hence, to sum up, self-inspection, external audits, timely revision of Standard operating procedures, training to personnel, management of deviations and implementing Corrective and Preventive actions (CAPAs) have to be followed to prevent defects in medicines to ensure patients safety and achieve better therapeutic compliance. Authors also anticipate the investigation strategy followed in this article can be implemented in post marketing surveillance phase of clinical trials by randomly sampling the products from various parts of the country stored at extreme conditions (cold, hot, humid etc.) and comparing against reserve samples. This study also helps in developing and inculcating quality, ethics and GMP right from the academic setup, in addition the pharmaceutical companies may adopt this methodology for investigating the market complaints or defects obtained thereby adding value and quality to the medicinal product.

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**CONFLICT OF INTEREST**

The authors declare no conflict of interest.

**ABBREVIATIONS**


**REFERENCES**


SUMMARY

The defects reported in this research work were captured by using the In-House mechanism. Products were procured, defects were identified and went through a process of investigation which categorized defects into minor, major and critical complaints. The defects identified are unique enough that they are not found in any standard textbooks and literature. The key message given through this article to the Pharmaceutical industry is that it is necessary for the implementation and practice of rules and regulation laid by regulatory agencies which ensure the quality of drugs and safety of patients.

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