



## A REVIEW ON IMPURITY PROFILING OF PHARMACEUTICALS

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### ABSTRACT

Various regulatory authorities like ICH, USFDA, Canadian Drug and Health Agency are emphasizing on the purity requirements and the identification of impurities in Active Pharmaceutical Ingredient's (API's). Qualification of the impurities is the process of acquiring and evaluating data that establishes biological safety of an individual impurity; thus, revealing the need and scope of impurity profiling of drugs in pharmaceutical research.

Identification of impurities is done by variety of Chromatographic and Spectroscopic techniques, either alone or in combination with other techniques. There are different methods for detecting and characterizing impurities with TLC, HPLC, HPTLC, AAS etc. Conventional Liquid Chromatography, particularly, HPLC has been exploited widely in field of impurity profiling; the wide range of detectors, and stationary phases along with its sensitivity and cost-effective separation have attributed to its varied applications. Among the various Planar Chromatographic Methods; TLC is the most commonly used separation technique, for isolation of impurities; due to its ease of operation and low cost compared to HPLC. An advancement of thin layer chromatography HPTLC, is a well-known technique for the impurity isolation.

Headspace GC is one of the most preferred techniques for identification of residual solvents. The advent of hyphenated techniques has revolutionized impurity profiling, by not only separation but structural identification of impurities as well. Among all hyphenated techniques, the most exploited techniques, for impurity profiling of drugs are LC-MS-MS, LC-NMR, LC-NMR-MS, GC-MS, and LC-MS

**KEYWORDS:** Impurity, Analytical Method Development, Spectrophotometry, Chromatography

### INTRODUCTION

There is an ever increasing interest in impurities present in API's. Recently, not only purity profile but also impurity profile has become essential as per various regulatory requirements. In the pharmaceutical world, an impurity is considered as any other organic material, besides the drug substance, or ingredients, arise out of synthesis or unwanted chemicals that remains with API's. The impurity may be developed either during formulation, or upon aging of both API's and formulated API's in medicines. A good illustration of this definition may be identification of impurity in API's like 1-(1, 2, 3, 5, 6, 7-hexahydro-s-indacen-4-yl)-3-4[- 1- hydroxy-1-methyl-ethyl]-furan-2-sulphonylurea using Multidisciplinary approach [1]. The presence of these unwanted chemicals, even in small amount, may influence the efficacy and safety of the pharmaceutical products. Impurity profiling (i.e., the identity as well as the quantity of impurity in the pharmaceuticals), is now gaining critical attention from regulatory authorities. The different Pharmacopoeias, such as the British Pharmacopoeia (BP), United States Pharmacopoeia (USP), and Indian Pharmacopoeia (IP) are slowly incorporating limits to allowable levels of impurities present in the API's or formulations.

The International Conference on Harmonization of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) has also published guidelines for validation of methods for analysing impurities in new drug substances,

products, residual solvents and microbiological impurities [2-5].

A number of articles [6-8] have stated guidelines and designed approaches for isolation and identification of process-related impurities and degradation products, using Mass spectrometry (MS), Nuclear Magnetic Resonance (NMR), High Performance Liquid Chromatography (HPLC), Fourier Transform Ion Cyclotron Resonance Mass Spectrometry (FTICR-MS), and Tandem Mass Spectrometry for pharmaceutical substances. Present article reveals different impurities found in the API's, methods for identifying them and the possible measures to deal with the interferences caused by them.

### Regulatory Guidelines on Impurities in an Active Pharmaceutical Ingredient

Ethical, economic and competitive reasons as well as those of safety and efficacy support the need to monitor impurities in drug products. However monitoring impurities and controlling these impurities mean different things to different people or to the same people at different times, even those in the pharmaceutical sciences and industry. A unified terminology is necessary to assure that everyone uses the same vocabulary when addressing questions related to impurities. The United States Food and Drug Administration (US FDA) have endorsed the guidance prepared under the guidance of the International Conference of harmonization (ICH). The ICH guideline for impurities in pharmaceuticals was developed with joint efforts



of regulators and industry representatives from the European Union (EU), Japan and United States and it has helped to ensure that different regions have consistent requirements for the data that should be submitted to various regulatory agencies. The guidelines not only aid the sponsors of New Drug Applications (NDA) or Abbreviated New Drug Application (ANDA) with the type of information that should be submitted with their applications, but also assist the FDA reviewers and field investigators in their consistent interpretation and implementation of regulations. The various regulatory guidelines regarding impurities are as follows:

1. ICH guidelines "stability testing of new drug substances and products"- Q1A
2. ICH guidelines "Impurities in New Drug Substances"- Q3A
3. ICH guidelines "Impurities in New Drug Products"- Q3B
4. ICH guidelines "Impurities: Guidelines for residual solvents"- Q3C
5. US-FDA guidelines "NDAs -Impurities in New Drug Substances"
6. US-FDA guidelines "ANDAs – Impurities in New Drug Substances"

Australian regulatory guideline for prescription medicines, Therapeutic Governance Authority (TGA), Australia

### Designation of Impurities

#### A. Common Terms of Impurities [09-15]

Following terms are used by various regulatory bodies and ICH to describe the impurities

- Intermediate
- Penultimate intermediate
- By-products
- Transformation products
- Interaction products
- Related products
- Degradation products
- Intermediate: The compounds produced during synthesis of the desired material or as a part of the route of synthesis.
- Penultimate Intermediate: It is the last compound in the synthesis chain prior to the production of the final desired compound.
- By-products: The compound produced in the reaction other than the required intermediates. They can occur through a variety of side reactions, such as overreaction, incomplete reaction, demonization and rearrangement, unwanted reactions between starting materials or intermediates with chemical reagents or catalysts.
- Transformation Products: They are related to theorized and nontheorized products that can occur in a reaction. They are similar to by-products except that more is known about these reaction products.
- Interaction Products: These products formed either intentionally or unintentionally interaction between various chemicals involved.
- Related Products: These are chemically similar to drug substance and may even possess biological activity.
- Degradation Products: They are formed by the

decomposition of active ingredient or other material of interest by the effect of external factors like heat, light and moisture.

Sr.No.	Impurity type	Impurity source
1	Process-related drug substance	Organic
		Starting material
		Intermediate
		By-product
2	Process-related drug product	Organic or inorganic
		Reagents, catalysts, etc
3	Drug substance or drug product	Organic
		Degradation products
4	Degradation drug product	Organic
		Excipient interaction

### Classification of Impurity [16-18]

#### United States Pharmacopoeia (USP)

According to USP impurities are classified into three sections

1. Impurities in Official Articles
2. Ordinary Impurities
3. Organic Volatile Impurities

### The ICH Terminology

According to ICH guidelines, impurities in drug substance produced by chemical synthesis can be broadly classified into following three categories

1. Organic Impurities (Process and drug-related)
2. Inorganic Impurities (Reagent, ligands, catalysts)
3. Residual Solvents (Volatile solvents)

#### 1. Organic Impurities

These types of impurities arise during the manufacturing process and/or during storage of the drug substance. These include following sub-impurities.

##### • Starting Materials or Intermediate Impurities

These types of impurities occur in almost every API unless a proper care is taken in every step during the multistep synthesis of drug product. Although the end products are always washed with solvents but there are chances of having the residual of unreacted starting materials unless the manufacturers are very careful about the impurities.

##### • By-products

In synthetic organic chemistry, getting a single end product with complete yield is very rare; there is always a chance of having by products along with desired end product.

##### • Degradation Products

Impurities can also be formed by degradation of the end product during manufacturing of bulk drugs. This mainly occurs due to improper storage of formulation.

### Other Types of Organic Impurities [19-20]

#### A. Synthesis Related Impurities

New chemical entity generated during synthetic process from raw material, solvent, intermediate, by- product. During synthesis process, if impurity present in trace or in significant amount in any of substance involved in reaction, that ultimately



result in final product contaminated with one or more unwanted materials. Therefore, synthesis related impurity require utmost care during every step involved in synthesis process to minimize level of impurity that can arise.

### B. Formulation Related Impurities:

Drug substance subjected to variety of conditions that leads to its degradation or other reactions. Solutions and suspensions are prone to degradation due to hydrolysis. Water used in formulation contribute to not only its impurity but also provide situation for hydrolysis and catalysis.

#### Factors Affecting on Formulation Related Impurities

##### A. Environment related

1. Exposed to adverse temperature: Substance which are labile to heat or in tropical temperature lead to degradation of active constitute and formation of impurity occurs. E.g. Vitamins are heat sensitive and its degradation lead to loss in potency.
2. Exposed to light: Photosensitive material when exposed to light / UV light undergo degradation which forms impurity.
3. Humidity: It can be detrimental to bulk powder and formulation containing solid dosage form.

##### b. Formation of impurities on ageing

Mutual interaction: Interaction between ingredients involved in formulation leads to mutual interaction which causes impurity formation.

##### B. Functional Group Related Impurities

- a) Ester hydrolysis: Drugs like aspirin, benzocaine, cefoxime, cocaine, ethyl paraben undergo ester hydrolysis.
- b) Hydrolysis: Commonly drugs like benzyl penicillin, barbital, and chloramphenicol undergo hydrolysis.
- c) Oxidative degradation: Drugs like hydrocortisone, methotrexate, heterocyclic aromatic ring, nitroso/nitrile derivative.
- d) Photolytic cleavage: Product exposed to light while manufacturing or storage in hospital pending use or by consumer pending use.
- e) Decarboxylation: Some dissolved carboxylic acid such as p-amino salicylic acid loose CO<sub>2</sub> when heated.

### 2. Inorganic Impurities

Inorganic impurities are also obtained from the manufacturing processes which are used in bulk drug formulation. They are normally known and identified.

- a) **Reagent, Ligands and Catalysts:** Rare chances of occurrence of these impurities. If during manufacturing procedure is not followed properly will create a problem.
- b) **Heavy Metals:** Water is generally used in different manufacturing processes which act as the main source of heavy metals, like Ar, Cd, Cr, Na, Mg, Mn, etc., where acidification or acid hydrolysis takes place. By using demineralized water and glass-lined reactors heavy metal impurities can be easily avoided
- c) **Other Materials (Filter Aids, Charcoal):** The filters or filtering aids such as centrifuge bags are routinely used in the bulk drugs manufacturing plants and in many cases, activated carbon is also used which also act as a source of

impurity. Therefore to avoid the contamination, regular monitoring of fibers and black particles in the bulk drugs is essential.

### 3. Residual Solvents

Residual solvents are organic or inorganic liquids used during the manufacturing process. It is very difficult to remove these solvents completely by the work-up process. Some solvent that are known to cause toxicity should be avoided in the production of bulk drugs.

#### Sources of Impurities

From the preceding discussion, it is clear that impurities can originate from several sources; such as; a) Crystallization-related impurities, b) Stereochemistry-related impurities, c) Residual solvents, d) Synthetic intermediates and by-products, e) Formulation-related impurities) Impurities arising during storage, h) Method related impurity, I) Mutual interaction amongst ingredients, h) Functional group-related typical degradation [6].

#### 1. Crystallization-Related Impurities

Based on the realization that the nature of structure adopted by a given compound upon crystallization, could exert a profound effect on the solid-state properties of that system, the pharmaceutical industry is required to take a strong interest in polymorphism and solvatomorphism as per the regulations laid down by the regulatory authorities.

Polymorphism is the term used to indicate crystal system where substances can exist in different crystal packing arrangements, all of which have the same elemental composition. Whereas, when the substance exists in different crystal packing arrangements, with a different elemental composition; the phenomenon is known as Solvatomorphism [6].

#### 2. Stereochemistry-Related Impurities

It is of paramount importance to look for stereochemistry related compounds; that is, those compounds that have similar chemical structure but different spatial orientation, these compounds can be considered as impurities in the APIs. Chiral molecules are frequently called enantiomers. The single enantiomeric form of chiral drug is now considered as an improved chemical entity that may offer a better pharmacological profile and an increased therapeutic index with a more favorable adverse reaction profile. However, the pharmacokinetic profile of levofloxacin (S-isomeric form) and ofloxacin (R-isomeric form) are comparable, suggesting the lack of advantages of single isomer in this regard [21]. The prominent single isomer drugs, which are being marketed, include levofloxacin (S-ofloxacin), 7 evalbuterol (R-albuterol), and esomeprazole (S-omeprazole).

#### 3. Residual solvents

Residual solvents are organic volatile chemicals used during the manufacturing process or generated during the production. Some solvents that are known to cause toxicity should be avoided in the production of bulk drugs. Depending on the possible risk to human health, residual solvents are divided into three classes [4]. Especially, solvents in Class I, viz benzene (2 ppm limit), carbon tetrachloride (4 ppm limit), methylene



chloride (600 ppm), methanol (3000 ppm, pyridine (200 ppm), toluene (890 ppm) should be avoided. In Class II, viz N, N-dimethylformamide (880 ppm), acetonitrile (410 ppm). Class III solvents, viz acetic acid, ethanol, acetone have permitted daily exposure of 50 mg or less per day, as per the ICH guidelines. A selective gas chromatography (GC) method has been developed to determine the purity of acetone, dichloromethane, methanol and toluene. Using this method, the main contaminants of each organic solvent can be quantified. Moreover, the developed method allows the simultaneous determination of ethanol, isopropanol, chloroform, benzene, acetone, dichloromethane, methanol and toluene with propionitrile as the internal standard [22].

#### 4. Synthetic Intermediates and By-Products

Impurities in pharmaceutical compounds or a new chemical entity (NCE) can originate during the synthetic process from raw materials, intermediates and/or by-products. For example, impurity profiling of ecstasy tablets by GC-MS, and MDMA samples, produced impurities in intermediates via reductive amination route [23].

#### 5. Formulation-Related Impurities

Many impurities in a drug product can originate from excipients used to formulate a drug substance. In addition, a drug substance is subjected to a variety of conditions in the process of formulation that can cause its degradation or have other undesirable reactions. If the source is from an excipient, variability from lot to lot may make a marginal product, unacceptable for reliability. Solutions and suspensions are inherently prone to degradation due to hydrolysis or solvolysis Fluocinonide Topical Solution USP, 0.05%, in 60-mL bottles, was recalled in the United States because of degradation/impurities leading to sub-potency [24]. In general, liquid dosage forms are susceptible to both degradation and microbiological contamination. In this regard, water content, pH of the solution/suspension, compatibility of anions and cations, mutual interactions of ingredients, and the primary container are critical factors Microbiological growth resulting from the growth of bacteria, fungi, and yeast in a humid and warm environment may result in unsuitability of an oral liquid product for safe human consumption. Microbial contamination may occur during the shelf life and subsequent consumer-use of a multiple-dose product, either due to inappropriate use of a multiple-dose product, either due to inappropriate use of certain preservatives in the preparations, or because of the semi-permeable nature of primary containers [25].

#### 6. Impurities Arising During Storage

A number of impurities can originate during storage or shipment of drug products. It is essential to carry out stability studies to predict, evaluate, and ensure drug product safety [6].

#### 7. Method Related Impurity

A known impurity, 1-(2, 6-dichlorophenyl) indolin-2-one is formed in the production of a parenteral dosage form of diclofenac sodium, if it is terminally sterilized by autoclave. The conditions of the autoclave method (i.e., 123 + 2 °C) enforce the intramolecular cyclic reaction of diclofenac sodium forming an indolinone derivative and sodium hydroxide. The formation of this impurity has been found to depend on initial pH

of the formulation [26]

#### 8. Mutual interaction amongst ingredients

Most vitamins are very labile and on aging they create a problem of instability in different dosage forms, especially in liquid dosage forms. Degradation of vitamins does not give toxic impurities; however, potency of active ingredients drops below Pharmacopeial specifications.

Because of mutual interaction, the presence of nicotinamide in a formulation containing four vitamins (nicotinamide, pyridoxine, riboflavin, and thiamine) can cause the degradation of thiamine to a sub-standard level within a one-year shelf life of vitamin B-complex injections. [27] The marketed samples of vitamin B-complex injections were found to have a pH range of 2.8 - 4.0. A custom-made formulation with simple distilled-water and a typical formulated vehicle including disodium edetate and benzyl alcohol were investigated, and similar mutual interactions causing degradation were observed.

#### 9. Functional group-related typical degradation

Ester hydrolysis can be explained with a few drugs viz aspirin, benzocaine, cefotaxime, ethyl paraben, and cefpodoxime proxetil [28].

Hydrolysis is the common phenomenon for ester type of drugs, especially in liquid dosage forms viz benzylpenicillin, oxazepam and lincomycin.

Oxidative degradation of drugs like hydrocortisone, methotrexate, hydroxyl group directly bonded to an aromatic ring (viz phenol derivatives such as catecholamines and morphine), conjugated dienes (viz vitamin A and unsaturated free fatty acids), heterocyclic aromatic rings, nitroso and nitrite derivatives, and aldehydes (especially flavorings) are all susceptible to oxidative degradation.

In maziopredone, the hydrolytic and oxidative degradation pathway in 0.1 mol L<sup>-1</sup> hydrochloric acid and sodium hydroxide at 80°C were studied [29].

Photolytic cleavage includes example of pharmaceutical products that are exposed to light while being manufactured as solid or solution, packaged, or when being stored in pharmacy shops or hospitals for use by consumers.

Ergometrine, nifedipine, nitroprusside, riboflavin and phenothiazines are very liable to photo-oxidation. In susceptible compounds, photochemical energy creates free radical intermediates, which can perpetuate chain reactions. Most compounds will degrade as solutions when exposed to high-energy UV exposures. Fluroquinolone antibiotics are also found to be susceptible to photolytic cleavage.

In ciprofloxacin eye drop preparation (0.3%), sunlight induces photocleavage reaction producing ethylenediamine analog of ciprofloxacin [30].

Decarboxylation of some dissolved carboxylic acids, such as p-aminosalicylic acid; shows the loss of carbon dioxide from the carboxyl group when heated. An example of decarboxylation is



the photoreaction of rifloxacin.

As seen earlier, impurities in drug products can come from the drug or from excipients or can be brought into the system through an inprocess step by contact with the packaging material.

For most drugs, the reactive species consist of;

- Water- that can hydrolyze some drugs or affect the dosage form performance
- Small electrophiles- like aldehydes and carboxylic acid derivatives
- Peroxides- that can oxidize some drugs
- Metals- which can catalyze oxidation of drugs and the degradation pathway
- Leachable or Extractables- can come from glass, rubber stoppers, and plastic packaging materials. Metal oxides such as Na<sub>2</sub>O, SiO<sub>2</sub>, CaO, MgO, are the major components leached/extracted from glass. Generally most synthetic materials contain leachable oligomers/monomers, vulcanizing agents, accelerators, plasticizers, and antioxidants [31]. Some examples of leachable/extractables from synthetic materials include styrene from polystyrene, diethylhexylphthalate (DEHP, plasticizer in PVC), dioctyltin isooctylmercaptoacetate (stabilizer for PVC), zinc stearate (stabilizer in PVC and polypropylene), [32] 2- mercaptobenzothiazole (accelerator in rubber stopper), and furfural from rayon. These impurities are needed to be analyzed by using different analytical methods.

### Analytical Method Development

New drug development requires meaningful and reliable analytical data to be produced at various stages of the development [33-35].

- a) Sample set selection for analytical method development
- b) Screening of Chromatographic conditions and Phases, typically using the linear- solvent-strength model of gradient elution
- c) Optimization of the method to fine-tune parameters related to ruggedness and robustness. The above three methods are briefly discussed in Table 2.

The impurities can be identified predominantly by following methods;

- Reference standard method
- Spectroscopic method
- Separation method
- Isolation method
- Characterization method
- Reference standard method

The key objective of this is to provide clarity to the overall life cycle, qualification and governance of reference standards used in development and control of new drugs. Reference standards serve as the basis of evaluation of both process and product performance and are the benchmarks for assessment of drug safety for patient consumption. These standards are needed, not only for the active ingredients in dosage forms but also for impurities, degradation products, starting materials, process intermediates, and excipients.

- Spectroscopic methods

The UV, IR, MS, NMR and Raman spectroscopic methods are routinely being used for characterizing impurities [6].

- Separation methods

The Capillary electrophoresis (CE), Chiral Separations, Gas Chromatography (GC), Supercritical Fluid Chromatography (SFC), TLC, HPTLC, HPLC are regularly being used for separation of impurities and degradation products [6]

- Isolation methods

It is often necessary to isolate impurities. but if the instrumental methods are used isolation of impurities is avoided is directly characterizes the impurities. Generally, chromatographic and non-chromatographic techniques are used for isolation of impurities prior its characterization. The term 'chromatographic reactor' refers to the use of an analytical-scale column as both a flow-through reactor, and simultaneously, as separation medium for the reactant(s) and product(s). By using an HPLC, chromatographic reactor approach, the solution- phase hydrolysis kinetics of the Aprepitant (Emend<sup>TM</sup>) prodrug, fosaprepitant dimeglumine, were investigated. In loratidine, impurity found was ofloratidine, other examples include celecoxib [36], and amikacin [37]. A list of methods that can be used for isolation of impurities is given below.

- Solid-phase extraction methods
- Liquid-liquid extraction methods
- Accelerated solvent extraction methods
- Supercritical fluid extraction
- Column chromatography
- Flash chromatography
- TLC
- GC
- HPLC
- HPTLC
- Capillary electrophoresis (CE)
- Supercritical fluid chromatography (SFC)
- Characterization methods

Highly sophisticated instrumentation, such as MS attached to a GC or HPLC, are inevitable tools in the identification of minor components (drugs, impurities, degradation products, metabolites) in various matrices. For characterization of impurities, different techniques are used; which are as follows;

#### 1. NMR

The ability of NMR to provide information regarding the specific bonding structure and stereochemistry of molecules of pharmaceutical interest has made it a powerful analytical instrument for structural elucidation. The ability of NMR- based diffusion coefficient determination to distinguish between monomeric and dimeric substances was validated using a standard mixture of authentic materials containing both monomers and dimers [38]. Unfortunately, NMR has traditionally been used as a less sensitive method compared to other analytical techniques. Conventional sample requirements for NMR are on the order of 10 mg, as compared with MS, which requires less than 1 mg.

#### 2. MS

It has an increasingly significant impact on the pharmaceutical



development process over the past several decades. Advances in the design and efficiency of the interfaces, that directly connect separation techniques with Mass Spectrometers have afforded new opportunities for monitoring, characterizing, and quantification of drug-related substances in active pharmaceutical ingredients and pharmaceutical formulations.

If single method fails to provide the necessary selectivity, orthogonal coupling of chromatographic techniques such as HPLC-TLC and HPLC-CE, or coupling of chromatographic separations with information rich spectroscopic methods such as HPLC-MS or HPLC-NMR may need to be contemplated, but hopefully only as a development tool rather than a tool for routine QC use.

**Hyphenated Methods**

- LC-MS-MS
- HPLC-DAD-MS
- HPLC-DAD-NMR-MS
- GC-MS
- LC-MS

An example of reverse-phase LC-MS analysis in gradient elution with two distinct soft ionization techniques is the Atmospheric pressure ionization with electrospray source (API-ESI) and the chemical ionization of d- allethrine [39].

The popularity of LC-MS-MS systems for complex mixture analysis of thermally labile and biologically relevant molecules, viz mosapride, is largely attributed to the “soft” nature of atmospheric pressure chemical ionization (APCI), and atmospheric pressure ionization (APPI), [40].

HPLC-DAD-MS (HPLC coupled with a diode array UV detector and a mass spectrometer, and such other techniques are

**Goals of impurity investigations**

Sr. No.	Process-Related Impurities	Degradation-Related Impurities
1	Identify significant impurities	Identify potential degradation product through stress testing and actual degradation products through stability studies.
2	Determine origin of impurities and method for elimination or reduction	Understand degradation pathway and methods to minimize degradation
3	Establish a control system for impurities involving Processing/manufacturing conditions Suitable analytical methods/ specifications	Establish a control system for impurities involving: 1.Processing/manufacturing conditions Suitable analytical methods/ specifications 2.Long term storage conditions including packaging 3.Formulation.

**CONCLUSION**

Nowadays, it is mandatory requirements in various pharmacopoeias to know the impurities present in API's. Isolation and characterization of impurities is required for acquiring and evaluating data that establishes biological safety which reveals the need and scope of impurity profiling of drugs in pharmaceutical research. To isolate and quantify the impurities, various instrumental analytical techniques are routinely been used.

almost routinely used NMR has now been added to this combination to provide HPLC-DAD-NMR-MS capabilities in a commercial instrument.

In GC-MS of methamphetamine and in LC-MS of risperidone, and cetirizine tablets a number of other chromatographic and spectroscopic configurations are found to be perfectly suitable for initial characterization of the impurities [41-44].

The goal for investigation of impurities is outlined in Table 3. A common goal for investigation of both process and product degradation-related impurities is to determine which of the many potential impurities are, in fact, produced in the manufacturing process and which occur under a given set of storage conditions

**Limits for Impurities**

According to ICH guidelines on impurities in new drug products, identification of impurities below 0.1% level, is not considered to be necessary, unless potential impurities are expected to be unusually potent or toxic. According to ICH, the maximum daily dose qualification threshold to be considered is as follows as shown in table no.2-5;< 2g/day0.1 % or 1 mg per day intake (whichever is lower) >2g/day 0.05%

**APPLICATIONS**

Numerous applications have been sought in the areas of drug designing and in monitoring quality, stability, and safety of pharmaceutical compounds, whether produced synthetically, extracted from natural products or produced by recombinant methods. The applications include alkaloids, amines, amino acids, analgesics, antibacterials, anticonvulsants, antidepressant, tranquilizers, antineoplastic agents, local anesthetics, macromolecules, steroids, miscellaneous.

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