significantly add to the cancer risk of the drug substance and impurities could be controlled at acceptable levels for non-mutagenic impurities.

Excipients used in existing marketed products and flavoring agents are excluded from this guideline. Application of this guideline to leachables associated with drug product packaging is not intended, but the safety risk assessment principles outlined in this guideline for limiting potential carcinogenic risk can be used if warranted. The safety risk assessment principles of this guideline can be used if warranted for impurities in excipients that are used for the first time in a drug product and are chemically synthesized.

# 3. GENERAL PRINCIPLES

The focus of this guideline is on DNA reactive substances that have a potential to directly cause DNA damage when present at low levels leading to mutations and therefore, potentially causing cancer. This type of mutagenic carcinogen is usually detected in a bacterial reverse mutation (mutagenicity) assay. Other types of genotoxicants that are non-mutagenic typically have thresholded mechanisms (5-9) and usually do not pose carcinogenic risk in humans at the level ordinarily present as impurities. Therefore to limit a possible human cancer risk associated with the exposure to potentially mutagenic impurities, the bacterial mutagenicity assay is used to assess the mutagenic potential/effect and the need for controls. Structure-based assessments are useful for predicting bacterial mutagenicity outcomes based upon the established knowledge base. There are a variety of approaches to conduct this evaluation including a review of the available literature, and/or computational toxicology assessment.

A Threshold of Toxicological Concern (TTC) concept was developed to define an acceptable intake for any unstudied chemical that will not pose a risk of carcinogenicity or other toxic effects. (10-11) For application of a TTC in the assessment of acceptable limits of mutagenic impurities in drug substances and drug product, a value of 1.5  $\mu$ /day corresponding to a theoretical  $10^{-5}$  excess lifetime risk of cancer, can be justified. The methods upon which the TTC is based are generally considered very conservative since they involve a simple linear extrapolation from the dose giving a 50% tumor incidence (TD<sub>50</sub>) to a 1 in  $10^6$  incidence, using TD<sub>50</sub> data for the most sensitive species and most sensitive site of tumor induction (several "worst case" assumptions). (10) Some structural groups were identified to be of such high potency that intakes even below the TTC would theoretically be associated with a potential for a significant carcinogenic risk. (12-13) This group of high potency mutagenic carcinogens ("cohort of concern") comprises aflatoxin-like-, N-nitroso-, and azoxy compounds.

During clinical development, it is expected that control strategies and approaches will be less developed in earlier phases where overall development experience is limited. This guideline bases acceptable intakes for mutagenic impurities on established risk assessment strategies. Acceptable risk during the early development phase is set at a theoretically calculated level of approximately one additional cancer per million. For later stages in development and marketed products when efficacy has been shown, acceptable increased cancer risk is set at a theoretically calculated level of approximately one in one hundred thousand. These risk levels represent a small theoretical increase in risk when compared to human overall lifetime incidence of developing any type of cancer, which is greater than 1 in 3. (14-15) It is noted that established cancer risk assessments are based on lifetime exposures. Less-than-lifetime exposures both during development and marketing can have higher acceptable intakes of impurities and still maintain comparable risk levels. The use of a numerical cancer risk value (1 in 100,000) and its translation into risk-based doses (TTC) is a highly hypothetical concept that should not

be regarded as a realistic indication of the actual risk. The TTC concept provides an estimate of safe exposures for any mutagenic compound. However, exceeding the TTC is not necessarily associated with an increased cancer risk given the conservative assumptions employed in the derivation of the TTC value. The most likely increase in cancer incidence is actually much less than 1 in 100,000. (13) In addition, in cases where a mutagenic compound is a non-carcinogen in a rodent bioassay, there would be no predicted increase in cancer risk. Based on these considerations, any exposure to an impurity that is later identified as a mutagen is not necessarily associated with an increased cancer risk for patients already exposed to the impurity. A risk assessment would determine whether any further actions would be taken.

Where a potential risk has been identified for an impurity, an appropriate control strategy leveraging process understanding and/or analytical controls should be developed to ensure that mutagenic impurity is at or below the acceptable cancer risk level.

There may be cases when an impurity is also a metabolite of the drug substance. In such cases, the impurity is considered qualified provided that exposure to the metabolite in appropriate nonclinical studies of the drug substance is higher than would be achieved from the impurity in the administered drug substance (ICH Q3A/Q3B).

#### 4. Considerations For Marketed Products

While this guideline is not intended to be applied retrospectively (i.e., to products marketed prior to adoption of this guideline), some types of post-approval changes warrant a reassessment of safety relative to mutagenic impurities. This section is intended to be applied to products marketed prior to, or after, the adoption of this guideline. Section 8.5 (Lifecycle management) contains additional recommendations for products marketed after adoption of this guideline.

# 4.1 Post Approval Changes to the Drug Substance Chemistry, Manufacturing, and Controls

Post approval submissions involving the drug substance chemistry, manufacturing, and controls (changes to the route of synthesis, reagents, solvents, process conditions etc.) should include an evaluation of the potential risk impact associated with mutagenic impurities. Specifically, changes should be evaluated to determine if the change results in any new mutagenic impurities or higher acceptance criteria for existing mutagenic impurities. Reevaluation of impurities not impacted by the change is not required. For example, when only a portion of the manufacturing process is changed, the assessment of risk from mutagenic impurities should be limited to whether any new mutagenic impurities result from the change, whether any mutagenic impurities formed during the affected step are increased, and whether any known mutagenic impurities from upstream steps are increased. Regulatory submissions associated with such changes should include a summary of the assessment and if appropriate an updated control strategy. Changes to site of manufacture would typically not require a reassessment of mutagenic impurity risk.

When a new drug substance supplier is proposed, evidence that drug substance produced by this supplier (using same route of synthesis) has been approved for an existing drug product marketed in the assessor's region is considered to be sufficient evidence of acceptable risk/benefit regarding mutagenic impurities and an assessment per this guideline is not required. If this is not the case, then an assessment per this guideline is expected.

# 4.2 Post Approval Changes to the Drug Product Chemistry, Manufacturing, and Controls

Post approval submissions involving the drug product (e.g., change in composition, manufacturing process, dosage form) should include an evaluation of the potential risk associated with any new mutagenic degradants or higher acceptance criteria for existing mutagenic degradants. If appropriate, the regulatory submission would include an updated control strategy. Reevaluation of the drug substance associated with drug products is not required or expected provided there are no changes to the drug substance. Changes to site of manufacture would typically not require a reassessment of mutagenic impurity risk.

# 4.3 Changes to the Clinical Use of Marketed Products

Changes to the clinical use of marketed products that typically may require a reevaluation of the mutagenic impurity limits include a significant increase in clinical dose, an increase in duration of use (in particular when a mutagenic impurity was controlled above the lifetime acceptable intake for a previous indication that may no longer be appropriate for the longer treatment duration associated with the new indication), or for a change in indication from a serious or life threatening condition where higher acceptable intakes were justified (Section 7.5) to an indication for a less serious condition where the existing impurity acceptable intakes may no longer be appropriate. Changes to the clinical use of marketed products associated with new routes of administration or expansion into patient populations that include pregnant women and/or pediatrics typically would not require a reevaluation, assuming no changes in daily dose or duration of treatment.

#### 4.4 Alternative Considerations for Marketed Products

Application of this guideline may be warranted to marketed products if there is specific cause for concern. The existence of impurity structural alerts alone is considered insufficient to trigger follow-up measures, unless it is a structure in the cohort of concern (Section 3). However a specific cause for concern would be new relevant impurity hazard data (classified as Class 1 or 2, Section 6) generated after the overall control strategy and specifications for market authorization were established. This new relevant impurity hazard data should be derived from high-quality scientific studies consistent with relevant regulatory testing guidelines, with data records or reports readily available to marketing application holders. When the applicant becomes aware of this new relevant impurity hazard data, an evaluation should be conducted and if it is concluded by the applicant to affect the acceptable cancer risk/benefit, notification (Section 9) to regulatory authorities with a proposed contemporary control strategy would be warranted.

# 5. DRUG SUBSTANCE AND DRUG PRODUCT IMPURITY ASSESSMENT

Actual and potential impurities that are likely to arise during the synthesis, work-up, and storage of a new drug substance and during manufacturing and storage of a new drug product should be assessed.

The impurity assessment is a two-stage process. Firstly, actual impurities that have been identified should be considered for their mutagenic potential. In parallel, an assessment of potential impurities likely to be present in the final drug substance is carried out to determine if further evaluation of their mutagenic potential is required. The steps as applied to synthetic impurities and degradants are described in Sections 5.1 and 5.2, respectively.

## 5.1 Synthetic Impurities

Actual impurities include those observed in the drug substance above the ICH Q3A reporting thresholds. Identification of actual impurities is expected when the levels exceed the identification thresholds outlined by ICH Q3A. It is acknowledged that some impurities below the identification threshold may also have been identified.

Potential impurities arising from the synthesis of the drug substance could include starting materials, reagents and intermediates, identified impurities in starting materials and intermediates, and reasonably expected reaction by-products based on knowledge of the chemical reactions and conditions involved. Knowledge of the starting material synthesis, in particular the use of mutagenic reagents is an important factor in understanding the potential impurities in the starting materials, especially when there is a reasonable expectation that such impurities may be carried through the synthesis to the drug substance.

All impurities (actual and potential), where the structures are known, should be evaluated for mutagenic potential as described in Section 6.

# 5.2 Degradants

Actual drug substance degradation products include those observed above the ICH Q3A reporting threshold during storage of the drug substance in the proposed long-term storage conditions and primary and secondary packaging. Actual drug product degradation products include those observed above the ICH Q3B reporting threshold during storage of the drug product in the proposed long-term storage conditions and primary and secondary packaging, and also include those impurities that arise during the manufacture of the drug product. Identification of actual degradation products is expected when the levels exceed the identification thresholds outlined by ICH Q3A/Q3B. It is acknowledged that some degradation products below the identification threshold may also have been identified.

Potential degradants in the drug substance and drug product are those that may be reasonably expected to form during long term storage conditions. Potential degradants include those that form above the ICHQ3A/B identification threshold during accelerated stability studies (e.g., 40°C/75% relative humidity for 6 months) and confirmatory photostability studies as described in ICH Q1B, (16) but are yet to be confirmed in the drug substance or drug product in the primary packaging.

Knowledge of relevant degradation pathways can be used to help guide decisions on the selection of potential degradation products to be evaluated for mutagenicity e.g., from degradation chemistry principles, relevant stress testing studies, and development stability studies.

Actual and potential degradants likely to be present in the final drug substance or drug product and where the structure is known should be evaluated for mutagenic potential as described in Section 6.

# 5.3 Considerations for Clinical Development

For products in clinical development, the thresholds outlined in ICHQ3A/B do not apply and it is acknowledged that the thresholds for actual impurities and degradants will typically be higher than those outlined in ICHQ3A/B.

# 6. HAZARD ASSESSMENT ELEMENTS

Hazard assessment involves an initial analysis of actual and potential impurities by conducting database and literature searches for carcinogenicity and bacterial mutagenicity data in order to classify them as Class 1, 2, or 5 according to Table 1. If data for such a classification are not available, an assessment of Structure-Activity Relationships (SAR) that focuses on bacterial mutagenicity predictions should be performed. This could lead to a classification into Class 3, 4, or 5.

Table 1: Impurities Classification with Respect to Mutagenic and Carcinogenic Potential and Resulting Control Actions (according to Ref. 17 with modifications)

Class	Definition	Proposed action for control (details in Section 7)
1	Known mutagenic carcinogens	Control at or below compound- specific acceptable limit
2	Known mutagens with unknown carcinogenic potential (bacterial mutagenicity positive*, no rodent carcinogenicity data)	Control at or below acceptable limits (generic or adjusted TTC)
3	Alerting structure, unrelated to the structure of the drug substance; no mutagenicity data.	Control at or below acceptable limits (generic or adjusted TTC) or do bacterial mutagenicity assay; If non-mutagenic = Class 5 If mutagenic = Class 2
4	Alerting structure, same alert in drug substance which has been tested and is non-mutagenic	Treat as non-mutagenic impurity
5	No structural alerts, or alerting structure with sufficient data to demonstrate lack of mutagenicity	Treat as non-mutagenic impurity

<sup>\*</sup>Or other relevant positive mutagenicity data indicative of DNA-reactivity related induction of gene mutations (e.g., positive findings in *in vivo* gene mutation studies)

A computational toxicology assessment should be performed using (Q)SAR methodologies that predict the outcome of a bacterial mutagenicity assay. Two (Q)SAR prediction methodologies that complement each other should be applied. One methodology should be expert rule-based and the second methodology should be statistical-based. (Q)SAR models utilizing these prediction methodologies should follow the validation principles set forth by the Organisation for Economic Co-operation and Development (OECD). (18)

The outcome of any computer system-based analysis should be reviewed with the use of expert knowledge in order to provide additional supportive evidence on relevance of any positive or negative prediction and to elucidate underlying reasons in case of conflicting results.

The absence of structural alerts from two complementary (Q)SAR methodologies (expert rule-based and statistical) is sufficient to conclude that the impurity is of no concern, and no further testing is required (Class 5 in Table 1).

To follow up on a structural alert (Class 3 in Table 1), a bacterial mutagenicity assay can be applied. An appropriately conducted negative bacterial mutagenicity assay (Note 2) would overrule any structure-based concern, and no further genotoxicity assessments would be required (Note 1). These impurities (Class 5 in Table 1) should be considered as a non-mutagenic impurity. A positive bacterial mutagenicity result would warrant further hazard assessment and/or control measures (Class 2 in Table 1). Alternatively adequate control measures in the case of a positive structural alert alone could be applied in place of bacterial mutagenicity testing.

An impurity with a structural alert that is shared with the drug substance (e.g., same structural alert in the same position and environment in the impurity and the drug substance) can be considered as non-mutagenic (Class 4 in Table 1) if the testing of the drug substance in the bacterial mutagenicity assay was negative.

Further hazard assessment of an impurity with a positive bacterial mutagenicity result (Class 2 in Table 1) may be appropriate for instance, when levels of the impurity cannot be controlled at an appropriate acceptable limit. In order to understand the relevance of the bacterial mutagenicity assay result under *in vivo* conditions, it is recommended that the impurity is tested in an *in vivo* gene mutation assay. The selection of other *in vivo* genotoxicity assays should be scientifically justified based on knowledge of the mechanism of action of the impurity and its organ site of contact (Note 3). *In vivo* studies should be designed taking into consideration existing guidance as per ICH S2(R1). (19) Negative results in the appropriate *in vivo* assay may support setting impurity limits in excess of the acceptable limits.

## 7. RISK CHARACTERIZATION

As a result of hazard assessment described in Section 6, each impurity will be assigned to one of the five classes in Table 1. For impurities belonging into Classes 1, 2, and 3 (Class 3 only if presence of a structural alert is not followed up in a bacterial mutagenicity assay), the principles of risk characterization used to derive acceptable intakes are described in this section.

#### 7.1 Generic TTC-based Acceptable Intakes

A TTC-based acceptable intake of a mutagenic impurity of 1.5 µg per person per day is considered to be associated with a negligible risk (theoretical excess cancer risk of <1 in 100,000 over a lifetime of exposure) and can in general be used for most pharmaceuticals as a default to derive an acceptable limit for control. This generic approach would usually be used for mutagenic impurities present in pharmaceuticals for long-term treatment (> 10 years) and where no carcinogenicity data are available (Classes 2 and 3).

# 7.2 Acceptable Intakes Based on Compound-Specific Risk Assessments

# 7.2.1 Mutagenic Impurities with Positive Carcinogenicity Data (Class 1 in Table 1)

Compound-specific risk assessments to derive acceptable intakes should be applied instead of the TTC-based acceptable intakes where sufficient carcinogenicity data exist. For a known mutagenic carcinogen, a compound-specific acceptable intake can be calculated based on carcinogenic potency and linear extrapolation as a default approach. Alternatively, other established risk assessment practices such as those used by international regulatory bodies may be applied either to calculate acceptable intakes or to use already existing values published by regulatory bodies (Note 4).

Compound-specific calculations for acceptable intakes can be applied case-by-case for impurities which are chemically similar to a known carcinogen compound class (class-specific acceptable intakes) provided that a rationale for chemical similarity and supporting data can be demonstrated (Note 5).

# 7.2.2 Mutagenic Impurities with Evidence for a Practical Threshold

The existence of mechanisms leading to a dose response that is non-linear or has a practical threshold is increasingly recognized, not only for compounds that interact with non-DNA targets but also for DNA-reactive compounds, whose effects may be modulated by, for example, rapid detoxification before coming into contact with DNA, or by effective repair of induced damage. The regulatory approach to such compounds can be based on the identification of a critical No-Observed Effect Level (NOEL) and use of uncertainty factors (ICH Q3C(R5)) (20) when data are available (Note 6).

The acceptable intakes derived from compound-specific risk assessments can be adjusted for shorter term use in the same proportions as defined in the following sections (Section 7.3.1 and 7.3.2).

# 7.3 Acceptable Intakes in Relation to LTL Exposure

The TTC-based acceptable intake of 1.5  $\mu$ g/day is considered to be protective for a lifetime of daily exposure. To address Less-Than-Lifetime (LTL) exposures to mutagenic impurities in pharmaceuticals, an approach is applied in which the acceptable cumulative lifetime dose (1.5  $\mu$ g/day x 25,550 days = 38.3 mg) is uniformly distributed over the total number of exposure days during LTL exposure. (21) This would allow higher daily intake of mutagenic impurities than would be the case for lifetime exposure and still maintain comparable risk levels for daily and non-daily treatment regimens. In the case of intermittent (non-daily) dosing, the acceptable intake will be capped by the total cumulative dose or the maximum acceptable intake (i.e., 120  $\mu$ g/day), whichever is lower. Table 2 illustrates the acceptable intakes for LTL to lifetime exposures for clinical development and marketing.

•			•
			>10
$\leq 1$	>1 - 12	>1 - 10	years to
month	months	years	lifetime
120	20	10	1.5
	month	month months	month months years

Table 2: Acceptable Intakes for an Individual Impurity

# 7.3.1 Clinical Development

[µg/day]

Using this LTL concept, acceptable intakes of mutagenic impurities are recommended for limited treatment periods during clinical development of up to 1 month, 1 to 12 months and more than one year up to completion of Phase III clinical trials (Table 2). These adjusted acceptable intake values maintain a 10<sup>-6</sup> risk level in early clinical development when benefit has not yet been established and a 10<sup>-5</sup> risk level for later stages in development (Note 7).

An alternative approach to the strict use of an adjusted acceptable intake for any mutagenic impurity could be applied for Phase I clinical trials of up to 14 days. Only impurities that are known mutagenic carcinogens (Class 1) and known mutagens of unknown carcinogenic potential (Class 2), as well as impurities in the cohort of concern

chemical class, should be controlled (see Section 8) to acceptable limits as described in Section 7. All other impurities would be treated as non-mutagenic impurities. This includes impurities which contain structural alerts (Class 3), which alone would not trigger action for an assessment for this limited Phase I duration.

#### 7.3.2 Marketed Products

Standard risk assessments of known carcinogens operate under the assumption that cancer risk increases as a function of cumulative dose. Thus, cancer risk of a continuous low dose over a lifetime would be equivalent to the cancer risk associated with an identical cumulative exposure averaged over a shorter duration or lifetime average daily dose. This assumption has been advocated by other regulatory agencies (22) and proposed elsewhere. (21)

For marketed product treatments with cumulative intakes of less than 10 years (continuous or total of intermittent treatments), the acceptable intake can be adjusted to  $\leq 10~\mu g/day$ . For marketed products with much shorter treatment duration indications, the acceptable intake values of Table 2 can be applied. The proposed intakes would all comply with the principle of not exceeding a  $10^{-5}$  cancer risk level (Note 7).

## 7.4 Acceptable Intakes for Multiple Mutagenic Impurities

The TTC-based acceptable intakes should be applied to each individual impurity. When there are multiple mutagenic impurities specified on the drug substance specification, total mutagenic impurities should be limited as described in Table 3 for clinical development and marketed products:

Table 3: A	Acceptable	Intakes fo	or Total	<b>Impurities</b>
------------	------------	------------	----------	-------------------

Duration of treatment	≤ 1 month	>1 - 12 months	>1 - 10 years	>10 years to lifetime
Daily intake [µg/day]	120	60	10 (30*)	5

<sup>\*</sup>For clinical development up to 3 years. Similar principles could be applied to marketed products with justification.

Only impurities that are specified on the drug substance specification contribute to the calculation for total. Degradants which form in the drug product would be controlled individually and a total limit would not be applied. The above approach is supported by a detailed analysis of the effect of combining multiple impurities that are in similar or different chemical classes and by the conservative assumptions incorporated into the TTC, and the low likelihood of synergistic carcinogenic effects at very low mutagenic impurity levels. (23)

# 7.5 Exceptions and Flexibility in Approaches

- Higher acceptable intakes may be justified when human exposure to the impurity will be much greater from other sources e.g., food, or endogenous metabolism (e.g., formaldehyde).
- Case-by-case exceptions to the use of the appropriate acceptable intake can be justified in cases of severe disease, reduced life expectancy, late onset but chronic disease, or with limited therapeutic alternatives.

• A disproportionally high number of members of some structural classes of mutagens, i.e., aflatoxin-like-, N-nitroso-, and azoxy structures, of which some may occur as impurities in pharmaceuticals, display extremely high carcinogenic potency. Acceptable intakes for these high-potency carcinogens would likely be significantly lower than the acceptable intakes defined in this guideline. While the principles of this guideline can be used, a case-by-case approach using e.g., carcinogenicity data from closely related structures, if available, usually needs to be developed to justify acceptable intakes for pharmaceutical development and marketed products.

The above risk approaches are applicable to all routes of administration and no corrections to acceptable intakes are generally warranted. Exceptions to consider may include situations where data justifies route-specific concerns that need to be evaluated case-by-case. These approaches are also applicable to all patient populations based upon the conservative nature of the risk approaches being applied.

#### 8. CONTROL

A control strategy is a planned set of controls, derived from current product and process understanding that assures process performance and product quality (ICH Q10). (24) A control strategy can include, but is not limited to, the following:

- Controls on material attributes (including raw materials, starting materials, intermediates, reagents, solvents, primary packaging materials);
- Facility and equipment operating conditions;
- Controls implicit in the design of the manufacturing process;
- In-process controls (including in-process tests and process parameters);
- Controls on drug substance and drug product (e.g., release testing).

When an impurity has been characterized as mutagenic, it is important to develop a control strategy that assures that the level of this impurity in the drug substance and drug product is below the acceptable limit. A thorough knowledge of the chemistry associated with the drug substance manufacturing process, the drug product manufacturing process, along with an understanding of the overall stability of the drug substance and drug product is fundamental to developing the appropriate controls. Developing a strategy to mitigate mutagenic impurities in the drug product is consistent with risk management processes identified in ICH Q9. (25) A control strategy that is based on product and process understanding and utilisation of risk management principles will lead to a combination of process design and control and appropriate analytical testing, which can also provide an opportunity to shift controls upstream and minimize the need for end-product testing.

#### 8.1 Control of Process Related Impurities

There are 4 potential approaches to development of a control strategy for drug substance:

## Option 1

Include a test for the impurity in the drug substance specification with an acceptance criterion at or below the acceptable limit using an appropriate analytical procedure. It is considered possible to apply periodic (verification) testing per ICH Q6A. (26)

## Option 2

Include a test for the impurity in the specification for a raw material, starting material or intermediate, or as an in-process control, with an acceptance criterion at or below the acceptable limit using an appropriate analytical procedure.

## Option 3

Include a test for the impurity in the specification for a raw material, starting material or intermediate, or as an in-process control, with an acceptance criterion above the acceptable limit using an appropriate analytical procedure coupled with demonstrated understanding of fate and purge and associated process controls that assure the level in the drug substance is below the acceptable limit without the need for any additional testing.

# Option 4

Understanding of process parameters and impact on residual impurity levels (including fate and purge knowledge) with sufficient confidence that the level of the impurity in the drug substance will be below the acceptable limit such that no analytical testing is needed for this impurity.

# 8.2 Discussion of Control Approaches

A control strategy that relies on process controls in lieu of analytical testing (Option 4) can be appropriate if the process chemistry and process parameters that impact levels of mutagenic impurities are understood and the risk of an impurity residing in the final drug substance or drug product above the acceptable limit is determined to be negligible. Elements of a scientific risk assessment/chemistry rationale should include an assessment of various factors that influence the fate and purge of an impurity including chemical reactivity, solubility, volatility, ionizability and any physical process steps designed to remove impurities. This option is especially useful for those impurities that are inherently unstable (e.g., thionyl chloride that reacts rapidly and completely with water) or for those impurities that are introduced early in the synthesis and are effectively purged.

For Option 4 approaches where justification based on scientific principles alone is not considered sufficient, as well as for Option 3 approaches, analytical data to support the control approach is expected. This could include as appropriate information on the structural changes to the impurity caused by downstream chemistry ("fate"), analytical data on pilot scale batches, and in some cases, laboratory scale studies with intentional addition of the impurity ("spiking studies"). In these cases, it is important to demonstrate that the fate/purge argument for the impurity is robust and will consistently assure a negligible probability of an impurity residing in the final drug substance above the acceptable limit. Where the purge factor is based on developmental data, it is important to address the expected scale-dependence or independence. In the case that the small scale model used in the development stage is considered to not represent the commercial scale, confirmation of suitable control in pilot scale and/or initial commercial batches is necessary. The need for data from pilot/commercial batches is influenced by the magnitude of the purge factor calculated from laboratory or pilot scale data, point of entry of the impurity, and knowledge of downstream process purge points.

If Options 3 and 4 cannot be justified, then a test for the impurity on the specification for a raw material, starting material or intermediate, or as an in-process control (Option 2) for drug substance (Option 1) at the acceptable limit should be included. For impurities introduced in the last synthetic step, an Option 1 control approach would be expected unless otherwise justified.

The application of "As Low As Reasonably Practicable" (ALARP) is not necessary if the level of the mutagenic impurity is below acceptable limits. Similarly, it is not necessary to demonstrate that alternate routes of synthesis have been explored.

In cases where control efforts cannot reduce the level of the mutagenic impurity to below the acceptable limit and levels are as low as reasonably practical, a higher limit may be justified based on a risk/benefit analysis.

## 8.3 Considerations for Periodic Testing

The above options include situations where a test is recommended to be included in the specification, but where routine measurement for release of every batch may not be necessary. This approach, referred to as periodic or skip testing in ICH Q6A could also be called "Periodic Verification Testing." This approach may be appropriate when it can be demonstrated that processing subsequent to impurity formation/introduction clears It should be noted that allowance of Periodic Verification Testing is contingent upon use of a process that is under a state of control (i.e., produces a quality product that consistently meets specifications and conforms to an appropriately established facility, equipment, processing, and operational control regimen). If upon testing, the drug substance or drug product fails an established specification, the drug producer should immediately revert to full testing (i.e., testing of every batch for the attribute specified) until the cause of the failure has been conclusively determined, corrective action has been implemented, and the process is again documented to be in a state of control. As noted in ICH Q6A, regulatory authorities should be notified of a periodic verification test failure to evaluate the risk/benefit of previously released batches that were not tested.

## 8.4 Control of Degradants

For a potential degradant that has been characterized as mutagenic, it is important to understand if the degradation pathway is relevant to the drug substance and drug product manufacturing processes and/or their proposed packaging and storage conditions. A well-designed accelerated stability study (e.g., 40°C/75% relative humidity, 6 months) in the proposed packaging, with appropriate analytical procedures is recommended to determine the relevance of the potential degradation product. Alternatively, well designed kinetically equivalent shorter term stability studies at higher temperatures in the proposed commercial package may be used to determine the relevance of the degradation pathway prior to initiating longer term stability studies. This type of study would be especially useful to understand the relevance of those potential degradants that are based on knowledge of potential degradation pathways but not yet observed in the product.

Based on the result of these accelerated studies, if it is anticipated that the degradant will form at levels approaching the acceptable limit under the proposed packaging and storage conditions, then efforts to control formation of the degradant is expected. The extent of degradation can often be lowered through formulation development and/or packaging designed to protect from moisture, light, or oxygen. Monitoring for the drug substance or drug product degradant in long term primary stability studies at the proposed storage conditions (in the proposed commercial pack) will generally be expected in these cases. The determination of the need for a specification for the mutagenic degradant will generally depend on the results from these stability studies.

If it is anticipated that formulation development and packaging design options are unable to control mutagenic degradant levels to less than the acceptable limit and levels are as low as reasonably practicable, a higher limit can be justified based on a risk/benefit analysis.

## 8.5 Lifecycle Management

This section is intended to apply to those products approved after the issuance of this guideline.

The quality system elements and management responsibilities described in ICH Q10 are intended to encourage the use of science-based and risk-based approaches at each lifecycle stage, thereby promoting continual improvement across the entire product lifecycle. Product and process knowledge should be managed from development through the commercial life of the product up to and including product discontinuation.

The development and improvement of a drug substance or drug product manufacturing process usually continues over its lifecycle. Manufacturing process performance, including the effectiveness of the control strategy, should be periodically evaluated. Knowledge gained from commercial manufacturing can be used to further improve process understanding and process performance and to adjust the control strategy.

Any proposed change to the manufacturing process should be evaluated for the impact on the quality of drug substance and drug product. This evaluation should be based on understanding of the manufacturing process and should determine if appropriate testing to analyze the impact of the proposed changes is required. Additionally, improvements in analytical procedures may lead to identification of an existing impurity or a new impurity. In those cases the new structure would be assessed for mutagenicity as described in this guideline.

Throughout the lifecycle of the product, it will be important to reassess if testing is needed when intended or unintended changes occur in the process. This applies when there is no routine monitoring at the acceptable limit (Option 3 or Option 4 control approaches), or when applying periodic rather than batch-by-batch testing. The appropriate testing to analyze the impact of the proposed change could include, but is not limited to, an assessment of current and potential new impurities and an assessment of the test procedures' abilities to detect any new impurities. This testing should be performed at an appropriate point in the manufacturing process.

In some cases, the use of statistical process control and trending of process measurements that are important for an Option 3 or Option 4 approach can be useful for continued suitability and capability of processes to provide adequate control on the impurity.

All changes should be subject to internal change management processes as part of the quality system. (24) Changes to information filed and approved in a dossier should be reported to regulatory authorities in accordance with regional regulations and guidelines.

# 8.6 Considerations for Clinical Development

It is recognized that product and process knowledge increases over the course of development and therefore it is expected that data to support control strategies in the clinical development trial phases will be less than at the marketing registration phase. A risk-based approach based on process chemistry fundamentals is encouraged to prioritize analytical efforts on those impurities with the highest likelihood of being present in the drug substance or drug product. Analytical data may not be needed to support early clinical development when the likelihood of an impurity being present is low, but in a similar situation analytical data may be needed to support the control approach for the marketing application. It is also recognized that commercial formulation design occurs later in clinical development and therefore efforts associated with drug product degradants will be limited in the earlier phases.

# 9. DOCUMENTATION

Information relevant to the application of this guideline should be provided at the following stages:

# 9.1 Clinical Development Trial Applications

- It is expected that the number of structures assessed for mutagenicity, and the collection of analytical data will both increase throughout the clinical development period.
- For Phase I clinical trials of 14 days or less, a summary of efforts to mitigate risks of mutagenic impurities focused on Class 1 and 2 impurities and those in the cohort of concern as outlined in Section 7 should be included.
- For other clinical development trials including Phase I studies of longer than 14 days, a list of the structures assessed by (Q)SAR should be included, and any Class 1, 2 or 3 actual and potential impurities should be described along with plans for control. The *in silico* (Q)SAR systems used to perform the assessments should be stated.
- Chemistry arguments may be appropriate instead of analytical data for potential impurities that present a low likelihood of being present as described in Section 8.6.

## 9.2 Common Technical Document (Marketing Application)

- For all actual and potential process related impurities and degradants where assessments according to this guideline are conducted, the mutagenic impurity classification and rationale for this classification should be provided:
  - o This would include the results and description of *in silico* (Q)SAR systems used, and as appropriate, supporting information to arrive at the overall conclusion for Class 4 and 5 impurities.
  - When bacterial mutagenicity assays were performed on impurities, all results and the study reports should be provided for any bacterial mutagenicitynegative impurities.
- Justification for the proposed specification and the approach to control should be provided (e.g., ICH Q11 example 5b). (27) For example, this information could include the acceptable intake, the location and sensitivity of relevant routine monitoring. For Option 3 and Option 4 control approaches,, a summary of knowledge of the purge factor, and identification of factors providing control (e.g., process steps, solubility in wash solutions, etc.) is important.

#### NOTES

- Note 1 The ICH M7 Guideline recommendations provide a state-of-the-art approach for assessing the potential of impurities to induce point mutations and ensure that such impurities are controlled to safe levels so that below or above the qualification threshold no further qualification for mutagenic potential is required. This includes the initial use of (Q)SAR tools to predict bacterial mutagenicity. In cases where the amount of the impurity exceeds 1 mg daily dose for chronic administration, evaluation of genotoxic potential as recommended in ICH Q3A/B could be considered.
- Note 2 To assess the mutagenic potential of impurities, a single bacterial mutagenicity assay can be carried out with a fully adequate protocol according to ICH S2(R1) and OECD 471 guidelines. The assays are expected to be performed in

compliance with Good Laboratory Practices (GLP) regulations; however, it is noted that the test article may not be prepared or analyzed in compliance with GLP regulations. Lack of full GLP compliance does not necessarily mean that the data cannot be used to support clinical trials and marketing authorizations. Such deviations should be described in the study report. In some cases, the selection of bacterial tester strains may be limited to those proven to be sensitive to an alert. For degradants that are not feasible to isolate or synthesize or when compound quantity is limited, it may not be possible to achieve the highest test concentrations recommended for an ICH-compliant bacterial mutagenicity assay according to the current testing guidelines. In this case, bacterial mutagenicity testing could be carried out using a miniaturized assay format with proven high concordance to the ICH-compliant assay to enable testing at higher concentrations with justification. Confidence in detection of mutagens requires testing concentrations at levels >250 µg/plate.

Note 3 Tests to Investigate the *in vivo* Relevance of *in vitro* Mutagens (Positive Bacterial Mutagenicity)

In vivo test	Mechanistic data to justify choice of test as fit-for-purpose
Transgenic mutation assays	For any bacterial mutagenicity positive. Justify selection of assay tissue/organ
Pig-a assay (blood)	• For directly acting mutagens (bacterial mutagenicity positive without S9)*
Micronucleus test (blood or bone marrow)	• For directly acting mutagens (bacterial mutagenicity positive without S9) and compounds known to be clastogenic*
Rat liver Unscheduled DNA Synthesis (UDS) test	<ul> <li>In particular for bacterial mutagenicity positive with S9 only</li> <li>Responsible liver metabolite known         <ul> <li>to be generated in test species used</li> <li>to induce bulky adducts</li> </ul> </li> </ul>
Comet assay	<ul> <li>Justification needed (chemical class specific mode of action to form alkaline labile sites or single-strand breaks as preceding DNA damage that can potentially lead to mutations</li> <li>Justify selection of assay tissue/organ</li> </ul>
Others	With convincing justification

<sup>\*</sup>For indirect acting mutagens (requiring metabolic activation), justification needed for sufficient exposure to metabolite(s)

#### Note 4 Example of linear extrapolation from the TD<sub>50</sub>

It is possible to calculate a compound-specific acceptable intake based on rodent carcinogenicity potency data such as  $TD_{50}$  values (doses giving a 50% tumor incidence equivalent to a cancer risk probability level of 1:2). Linear extrapolation to a probability of 1 in 100,000 (i.e., the accepted lifetime risk level used) is achieved by simply dividing the  $TD_{50}$  by 50,000. This procedure is similar to that employed for derivation of the TTC.

## Calculation example: Ethylene oxide

TD<sub>50</sub> values for ethylene oxide according to the Carcinogenic Potency Database (29) are 21.3 mg/kg body weight/day (rat) and 63.7 mg/kg body weight/day

(mouse). For the calculation of an acceptable intake, the lower (i.e., more conservative) value of the rat is used.

To derive a dose to cause tumors in 1 in 100,000 animals, divide by 50,000:

 $21.3 \text{ mg/kg} \div 50,000 = 0.42 \,\mu\text{g/kg}$ 

To derive a total human daily dose:

 $0.42 \mu g/kg/day \times 50 kg body weight = 21.3 \mu g/person/day$ 

Hence, a daily life-long intake of 21.3  $\mu$ g ethylene oxide would correspond to a theoretical cancer risk of  $10^{-5}$  and therefore be an acceptable intake when present as an impurity in a drug substance.

# Alternative methods and published regulatory limits for cancer risk assessment

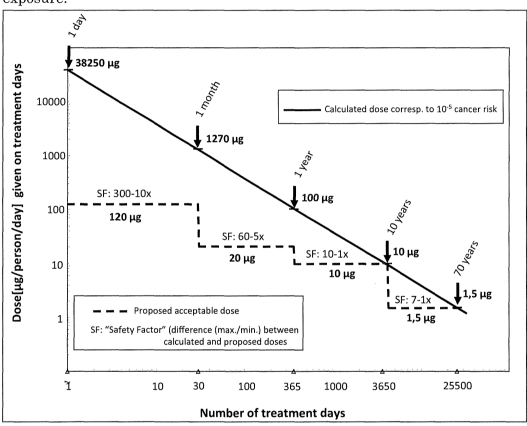
As an alternative of using the most conservative TD<sub>50</sub> value from rodent carcinogenicity studies irrespective of its relevance to humans, an in-depth toxicological expert assessment of the available carcinogenicity data can be done in order to initially identify the findings (species, organ etc) with highest relevance to human risk assessment as a basis for deriving a reference point for linear extrapolation. Also, in order to better take into account directly the shape of the dose-response curve, a benchmark dose such as a benchmark dose lower confidence limit 10% (BMDL10, an estimate of the lowest dose which is 95% certain to cause no more than a 10% cancer incidence in rodents) may be used instead of TD<sub>50</sub> values as a numerical index for carcinogenic potency. Linear extrapolation to a probability of 1 in 100,000 (i.e., the accepted lifetime risk level used) is then achieved by simply dividing the BMDL10 by 10,000.

Compound-specific acceptable intakes can also be derived from published recommended values from internationally recognized bodies such as World Health Organization (WHO, International Program on Chemical Safety [IPCS] Cancer Risk Assessment Programme) (30) and others using the appropriate 10<sup>-5</sup> lifetime risk level. In general, a regulatory limit that is applied should be based on the most current and scientifically supported data and/or methodology.

- Note 5 A compound-specific calculation of acceptable intakes for mutagenic impurities may be applied for mutagenic impurities (without carcinogenicity data) which are structurally similar to a chemically-defined class of known carcinogen. For example, factors that are associated with the carcinogenic potency of alkyl halides have been identified (31) and can be used to modify the safe acceptable intake of monofunctional alkyl halides, a group of alkyl halides commonly used Compared to multifunctional alkyl halides drug synthesis. monofunctional compounds are much less potent carcinogens with TD50 values ranging from 36 to 1810 mg/kg/day (n=15; epichlorohydrin with two distinctly different functional groups is excluded). (31) A TD<sub>50</sub> value of 36 mg/kg/day can thus be used as a still very conservative class-specific potency reference point for calculation of acceptable intakes for monofunctional alkyl halides. potency level is at least ten-fold lower than the TD<sub>50</sub> of 1.25 mg/kg/day corresponding to the default lifetime TTC (1.5 µg/day) and therefore justifies lifetime and less-than-lifetime daily intakes for monofunctional alkyl halides ten times the default ones.
- Note 6 Some published data give reliable experimental evidence for (practical) thresholds in the dose response for compounds that are positive for bacterial mutagenicity. This includes examples of thresholds in error-free repair capacity

of the mutagenic DNA-ethylating agent ethyl methanesulfonate (EMS) (32) or similarly for methylating agents. (33) Thresholds involving metabolic detoxification processes also appear to exist for 1, 3-butadiene. (34) Further, a threshold for oxidative DNA damage associated with the buildup of hemosiderin has been shown for p-chloroaniline hydrochloride. (35) Aside of mechanistic considerations supporting an experimentally observed threshold, it is important that a proper statistical analysis supports this assumption as well. (36)

Note 7 Establishing less-than-lifetime acceptable intakes for mutagenic impurities in pharmaceuticals has precedent in the establishment of the staged TTC limits for clinical development. (17) The calculation of less-than-lifetime Acceptable Intakes (AI) is predicated on the principle of Haber's rule, a fundamental concept in toxicology where concentration (C) x time (T) = a constant (k). Therefore, the carcinogenic effect is based on both dose and duration of exposure.



<u>Figure 1:</u> Illustration of calculated daily dose of a mutagenic impurity corresponding to a theoretical 1:100,000 cancer risk as a function of duration of treatment in comparison to the acceptable intake levels as recommended in Section 7.3.

The solid line in Figure 1 represents the linear relationship between the amount of daily intake of a mutagenic impurity corresponding to a  $10^{-5}$  cancer risk and the number of treatment days. The calculation is based on the TTC level as applied in this guideline for life-long treatment i.e., 1.5 µg per person per day using the formula:

Less-than-lifetime AI =  $\underline{1.5 \mu g \times (365 \text{ days} \times 70 \text{ years lifetime} = 25,550)}$ Total number of treatment days

The calculated daily intake levels would thus be  $1.5~\mu g$  for treatment duration of 70 years, 10  $\mu g$  for 10 years, 100  $\mu g$  for 1 year, 1270  $\mu g$  for 1 month and

approximately 38.3 mg as a single dose, all resulting in the same cumulative intake and therefore theoretically in the same cancer risk (1 in 100,000).

The dashed step-shaped curve represents the actual daily intake levels adjusted to less-than-lifetime exposure as recommended in Section 7 of this guideline for products in clinical development and marketed products. These proposed levels are in general significantly lower than the calculated values thus providing safety factors that increases with shorter treatment durations.

The proposed accepted daily intakes are also in compliance with a 10-6 cancer risk level if treatment durations are not longer than 6 months\* and are therefore applicable in early clinical trials with volunteers/patients where benefit has not yet been established. In this case the safety factors as shown in the upper graph would be reduced by a factor of 10.

\*At 6 months the calculated dose at a  $10^{-6}$  risk level would be  $20~\mu g$  which is identical to the recommended accepted dose i.e., there is no extra safety factor; at longer duration the theoretical  $10^{-6}$  risk level would be exceeded.

#### GLOSSARY

## Acceptable intake:

In the context of this guideline, an intake level that is without appreciable cancer risk.

# Acceptable limit:

Maximum acceptable concentration of an impurity in a drug substance or drug product derived from the acceptable intake and the daily dose of the drug.

# Acceptance criterion:

Numerical limits, ranges, or other suitable measures for acceptance of the results of analytical procedures.

#### BMDL10:

The lower 95% confidence interval of a Benchmark-dose representing a 10% response (e.g., tumor response upon lifetime exposure), i.e., the lower 95% confidence interval of a BMD10. BMD10 is the Benchmark-dose (BMD) associated with a 10% response adjusted for background.

## Control strategy:

A planned set of controls, derived from current product and process understanding that ensures process performance and product quality. The controls can include parameters and attributes related to drug substance and drug product materials and components, facility and equipment operating conditions, in-process controls, finished product specifications, and the associated methods and frequency of monitoring and control.

#### Cumulative intake:

The total intake of a substance that a person is exposed to over time.

#### Degradant:

Degradation product as defined in ICH Q3B.

#### DNA-reactive:

Substances that have a potential to induce direct DNA damage through chemical reaction with DNA.

#### Expert knowledge:

In the context of this guideline, expert knowledge can be generalized as a review of preexisting data and the use of any other relevant information to evaluate the accuracy of an *in silico* model prediction for mutagenicity.

#### Genotoxicity:

A broad term that refers to any deleterious change in the genetic material regardless of the mechanism by which the change is induced.

#### In-process control:

Checks performed during production to monitor and, if appropriate, to adjust the process and/or to ensure that the intermediate or drug substance conforms to its specifications.

# Mutagenic impurity:

An impurity that has been demonstrated to be mutagenic in an appropriate mutagenicity test model, e.g., bacterial mutagenicity assay.

# **NOEL:**

Abbreviation for No-Observed-Effect-Dose (level): The highest dose of substance at which there are no biologically significant increases in frequency or severity of any effects in the exposed humans or animals.

# Periodic (verification) testing:

Also known as periodic or skip testing in ICH Q6A.

# (Q)SAR and SAR:

In the context of this guideline, refers to the relationship between the molecular (sub) structure of a compound and its mutagenic activity using (Quantitative) Structure-Activity Relationships derived from experimental data.

## Purge factor:

Purge reflects the ability of a process to reduce the level of an impurity, and the purge factor is defined as the level of an impurity at an upstream point in a process divided by the level of an impurity at a downstream point in a process. Purge factors may be measured or predicted.

## Statistical process control:

Application of statistical methodology and procedures to analyze the inherent variability of a process.

#### Structural alert:

In the context of this guideline, a chemical grouping or molecular (sub) structure which is associated with mutagenicity.

#### $TD_{50}$ :

The dose-rate in mg/kg body weight/day which, if administered chronically for the standard lifespan of the species, will halve the probability of remaining tumorless throughout that period.

#### Threshold:

Categorically, a dose of a substance or exposure concentration below which a stated effect is not observed or expected to occur.

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