

STABILITY TESTING OF NEW DRUG SUBSTANCES AND PRODUCTS

guidance in
details in

→ ICH GUIDELINE Q1A(R2)

له في بعض الـ details موجودة بـ Q1B

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product should be
at high quality
safe ← effective

shelf-life ← فعالية

يتم تقديره بناءً

على الـ stability study of :

1. active pharmaceutical ingredient (API)
2. drug product

General Principles

- The purpose of stability testing is:
 - to provide evidence on how the quality of a drug substance or drug product varies with time under the influence of a variety of environmental factors such as temperature, humidity, and light
 - to establish:
 - ① re-test period for the drug substance or
 - ② shelf life for the drug product
 - ③ To recommended storage conditions.

بالسابق صعد رأيي عن حوضتي : 1. stress testing 2. stability testing لازم نفوف الغرف بيهم :

1. stress testing ← بعض الدواء يتغير لـ extremely stressful conditions فبظروف التخزين العادية يتكون درجات حرارة

2. degradation ← عناء أهدر degradation pathway of my product. عمل ليس degradation في درجات حرارة عالية ولا بوجود ماء ربيبي الـ sun light و pH

بالتالي أناسب امره هو ان degradation pathway ← we can avoid it ①

② identify degradation products

↓
شوا impurities يلي راجع تنبع عن كلة ال degradation
سواء ال drug substance or drug product

بينا بال stability testing ← جرفن البعار تحت normal conditions عشان اقدر افهم :

1. shelf-life.

2. storage conditions.

لا لما عت ال stress test نتج عتبي impurities خاينا نسجها X

ولما عت ال stability test مواد ال X واطلع عتبي بالتالي بال stability testing

مش لدرم اتابع ال impurities لانها ما بتنتج في ظروف عادية.

ال re-testing ال drug (API) substance ← يعني بعد انتهاء التاريخ الموضح علينا فانا

ما بكيضا ← لا ممكن أعمل عليها خواصات واشوف اذا هي meet specifications ، اذا محتاجة

كل ال specifications موجودة انا ممكن استقرصها واهنق نول.

ال shelf-life ال drug product ← بعد انتهاء هاي الفترة البعارة عتبي صالح ال product
يعني عازلة.

Aim of the guidelines (Q1A)

The guideline addresses the information to be submitted in registration applications for new molecular entities and associated drug products.

Regulatory authorities
as apart of stability testing in details
information

Stress Testing

Stress testing (drug substance)

- Studies undertaken to elucidate the intrinsic stability of the drug substance. → under sever conditions
- Such testing is part of the development strategy and is normally carried out under more severe conditions than those used for accelerated testing.

Stress testing (drug product)

- Studies undertaken to assess the effect of severe conditions on the drug product.
- Such studies include photostability testing and specific testing on certain products, (e.g., metered dose inhalers, creams, emulsions, refrigerated aqueous liquid products).
① → بالتالي نحدد الـ packaging of our product

Stress Testing

➤ Stress testing of the drug substance can help to:

1. establish the intrinsic stability of the molecule
2. identify the likely degradation products (impurities)
3. establish the degradation pathways → so we can avoid it.
4. validate the stability indicating power of the analytical procedures used.

➤ The nature of the stress testing will depend on the

- ① individual drug substance and the type of drug
- ② product involved.

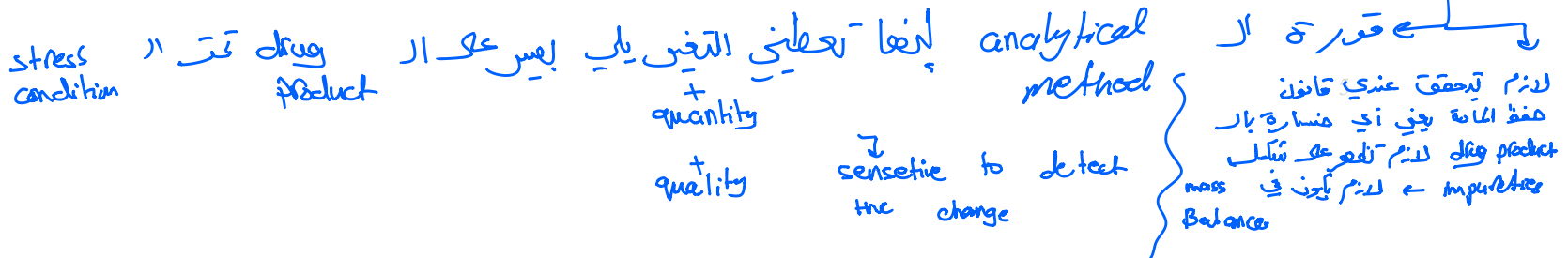
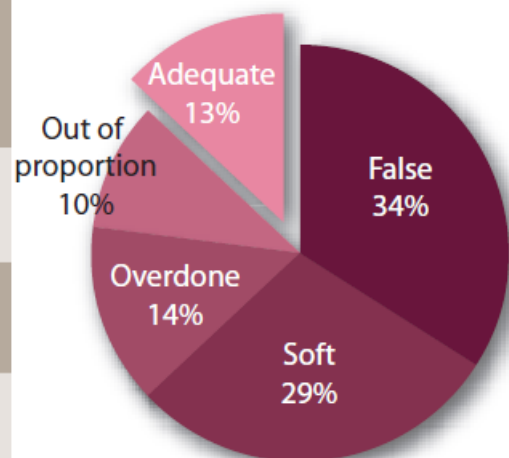


Table II: Type of degradation observed with a fixed set of fast and severe stress conditions.

Category	Explanation
Soft	No significant degradation and therefore no relevant degradation products observed
False	Fair amount of degradation (<15%), however no relevant degradation product(s) observed
Adequate	Fair amount of degradation (<15%) and at least one or all relevant degradation product(s) observed
Out of proportion	Between 15 and 100% degradation and at least one relevant degradation product observed
Overdone	Between 15 and 100% degradation, however, no relevant degradation products are observed



Klick et al. Pharm Technol. 2005 Feb;29(2):48-66.

Stress Testing

➤ Stress testing is likely to be carried out on a **single batch** of the drug substance.

➤ It should evaluate :

1. the effect of temperatures (in 10°C increments → (e.g., 50°C, 60°C, etc.) above that for accelerated testing),

ضعيف ١٠ درجات
حرارة بعد ١٠
accelerating testing
condition = 40°C

2. the effect of humidity (e.g., 75% RH or greater) where appropriate,

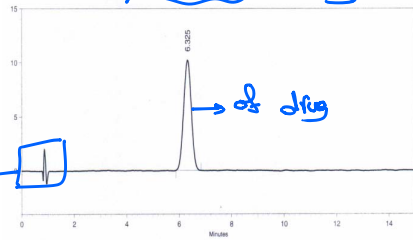
3. Oxidation

4. photolysis.

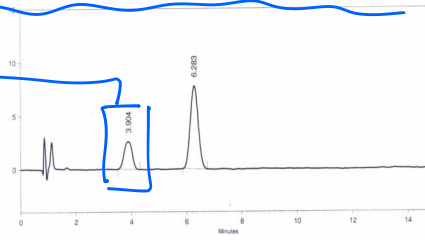
5. the susceptibility of the drug substance to hydrolysis across a wide range of pH values when in solution or suspension.

↓
acidic or basic pH
↓
mainly for suspension & solution.

standard (drug) . (Reference)

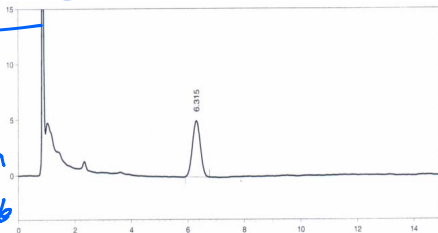


Degraded under acidic conditions



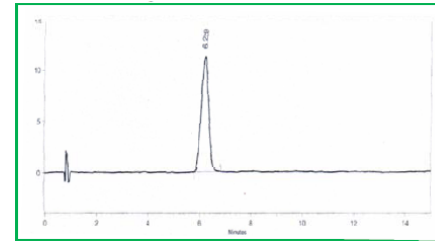
degradation product
 drug 1 peak
 not stable under acidic conditions.

Degraded under basic conditions



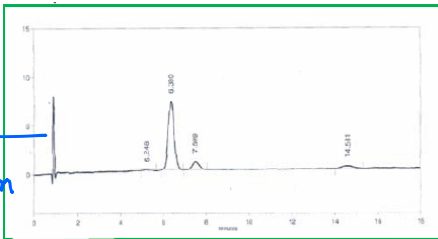
there are new peaks
 ↓
 degradation
 ↓
 not stable under basic conditions.

under thermal degradation conditions



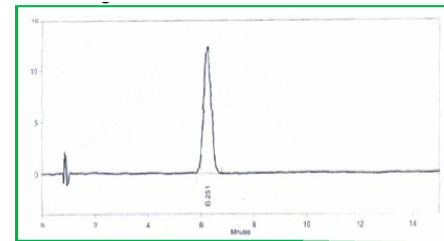
→ stable
 ↓
 no change
 ↓
 not thermolabile

Degraded under peroxide (H2O2) conditions



There are degradation

Exposed to direct sunlight



→ stable
 ↓
 not photosensitive
 لا تتأثر بالضوء
 الالكترية فاني تفسر

Chromatogram of Olopatadine HCl

الظروف التي تتأثر بها الدواء عند التحلل في ظروف حمضية ، قاعدية ، أكسدة ، حرارة ، ضوء
 ← نوع ال degradation product

Stress Testing

جزء أساسي في عملية التسجيل

- Results from these studies will form an integral part of the information provided to regulatory authorities. **Registration**
- It may not be necessary to examine specifically for certain degradation products appearing in stress testing if it has been demonstrated that they are not formed under accelerated or long term storage conditions.
- **bracketing** is defined as "the design of a stability schedule such that only samples on the extremes of certain design factors, e.g., strength, package size, are tested at all time points as in a full design."
- **Matrixing** is "the design of a stability schedule such that a selected subset of the total number of possible samples for all factor combinations is tested at a specified time point. At a subsequent time point, another subset of samples for all factor combinations is tested."

حكما فوق
بانه اذا

degradation
product

stress
test

تفحص في ال

وما تفحص في ال
stability
test

بالتالي عند ذلك

افحص بعد

صليته بال

stability
testing

size معين لـ

1. drug product

2. drug substances.

3 different patches

stability
test

* في ال

Drug product → in final Backiging → لا تملك واحد ال shelf-life
 بالمعدلية بال Final Backiging

1. number of batches should be → 3 patches
2. size of the patch.

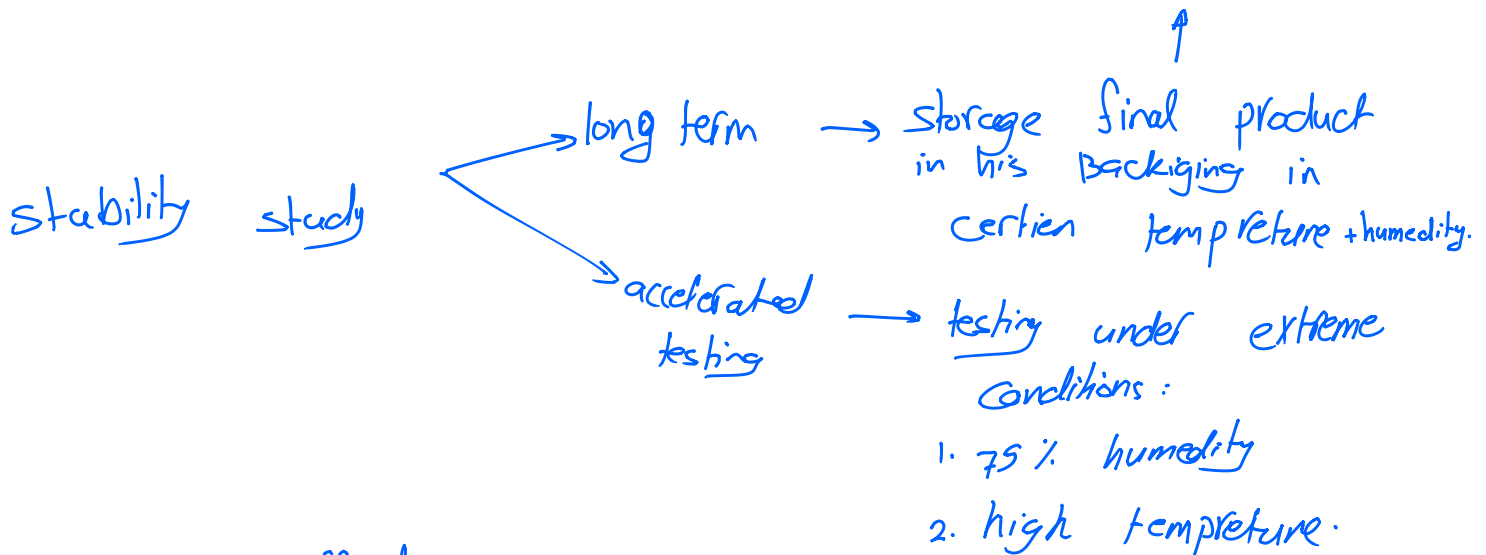
لا نعني drug يعني فيه different strength ← كل strength لازم
 ندرج stability ← عدد كبير من العينات يحتاج تحليل + يحتاج وقت + need money
 test

يعني عندي دواء في فيه ٣ عيارات 250, 300, 500 ← كل عيار لازم
 يجرى stability ← ببلاها 250 + 500 ← stable فسيلاي بنفاز 200
 انعاما تكون stable ← مفعلة الوقت وتكلفة عالية بالتالي سه حونا باستي اسسه

1. bracketing

2. matrixing

تفضل لفترة طويلة ← بعد testing ← time interval معين.



different time ← drug ← ل بعد تحليل
 ↓
 time = Zero, 6 months.

بالعادة ما يتناول أكثر من 6 أشهر وفي باليف كان عينة
 ٣ أشهر أو شهرين

استقرار لا middle strength في صا 500 في 500 - 250 - 500
 يعني اذا ال 250 وال 500
 stable بالاتي ال 300
 تكون stable دافي داعر
 اعل عليها stability test

أقواس (عنى ريف هين)

Bracketing Example:

• **Drug Product:** Paracetamol tablets
Factors:

1 2 3
 • **Strength:** 250 mg (low), 300 mg (middle), 500 mg (high)

• **Package size:** 10 tablets (small) and 100 tablets (large)

لصون لازم ادرس الاثنين في بي اذا في عدد بالهن بينهم ما يدركه

Assumes that only the **extreme strengths** (250 mg and 500 mg) and the **extreme package sizes** (10 tablets and 100 tablets) are tested, with the middle case excluded.

drug هو جود في (علبة زي ال 100) (ansophar)
 انه الموجود بين حبات الدواء بتختلف باختلاف عدد الحبات في صون برفو على 9 أشهر هذا المثال ناقص

Strength	Package Size	<u>0</u> Month	<u>6</u> Months	<u>12</u> Months	<u>18</u> Months	<u>24</u> Months	<u>36</u> Months
<u>250</u> mg	10 tablets	✓	✓	✓	✓	✓	✓
<u>250</u> mg	100 tablets	✓	✓	✓	✓	✓	✓
<u>300</u> mg	50 tablets	Excluded	Excluded	Excluded	Excluded	Excluded	Excluded
<u>500</u> mg	<u>10</u> tablets	✓	✓	✓	✓	✓	✓
<u>500</u> mg	<u>100</u> tablets	✓	✓	✓	✓	✓	✓

وبأثره
 ال stability
 testing according specifications
 صون حابر صوم
 testing at all time point.

قادرست ال strength
time point

Matrixing Example

• **Drug Product:** Paracetamol tablets

Factors:

• **Strength:** 250 mg (low), 300 mg (middle), 500 mg (high)

• **Package size:** 10 tablets (small) and 100 tablets (large)

➤ Involves selecting a **subset** of all possible samples (strengths, package sizes, and time points). Testing is rotated among subsets over the study period.

1. strength
2. size
3. time point

Strength	Package Size	0 Month	<u>6</u> Months	<u>12</u> Months	18 Months	24 Months	36 Months
250 mg	<u>10</u> tablets	✓	✓ <u> </u>		✓ <u> </u>		✓
250 mg	100 tablets	✓		✓		✓	
300 mg	50 tablets	✓			✓		✓
500 mg	<u>10</u> tablets	✓	✓ <u> </u>		✓		✓
500 mg	100 tablets	✓		✓		✓	

یک بولیم
test
منت بعد 6 أشهر
فا بولیم
بعد 12 أشهر
↓
Rotation
of
testing
over time
period.
فا نجلس الوقت
بالا وأنا دارس
المستحضرات
نفس عدد
المرات

Selection of Batches

Pilot scale batch

Stability test. production batch
تكون المبر بال حجم في ال

- A batch of a drug substance or drug product manufactured by a procedure fully representative of and simulating that to be applied to a full production scale batch.
- For solid oral dosage forms, a pilot scale is generally, at a minimum, one-tenth (1/10) that of a full production scale or 100,000 tablets or capsules, whichever is the larger.

Production batch

- A batch of a drug substance or drug product manufactured at production scale by using production

equipment in a production facility as specified in the application.

Storage Conditions: Drug substance

الدفعة التي يستخدمها للإثبات ملاحظة المفتوح عند التسجيل

Primary batch

- A batch of a drug substance or drug product used in a formal stability study, from which stability data are submitted in a registration application for the purpose of establishing a re-test period or shelf life, respectively.
- A primary batch:
 - 3 batch at least pilot
 - For a drug substance: should be at least a pilot scale batch.
 - For a drug product: two of the three batches should be at least pilot scale batch, and the third batch can be smaller if it is representative with regard to the critical manufacturing steps.
- However, a primary batch may be a production batch.

حرفه انوار Primary Batch تكون pilot حداقل minimum يعني يكون استخدام Production Batch عادي جداً.

Selection of Batches

Drug product

Drug substance

<u>3</u> (preferably from <u>different batches</u> of drug substance)	<u>3</u>	Number of <u>batches</u>
Two: <u>at least pilot</u> One: <u>smaller scale</u> , if justified	<u>Minimum pilot</u>	<u>Scale</u>
same <u>formulation and package</u> as proposed for marketing	<u>method of manufacture</u> <u>simulates production scale</u>	
on each <u>individual strength and</u> <u>container size unless bracketing or</u> <u>matrixing is applied</u>		
<u>container closure system proposed</u> <u>for marketing (including any</u> <u>secondary packaging and</u> <u>container label)</u>	same as or simulates the <u>packaging proposed for</u> <u>storage and distribution</u> <u>1</u> <u>2</u>	

+ primary

Specifications

Drug product

Drug substance

Release and shelf life
1 2

Release

Type

Stability studies should include testing of those attributes of the drug substance that are susceptible to change during storage and are likely to influence quality, safety, and/or efficacy.

Validated stability-indicating analytical procedures should be applied.

Any differences between the release and shelf life acceptance criteria for antimicrobial preservative content should be supported by a validated correlation of chemical content and preservative effectiveness

Ex:

preservative release → 95 - 105
⇒ shelf-life → 85 - 95

اذا data بس لازم يكون عندي
still 85 لو توكيزه نزل ل antimicrobial effect
قاد ريفيل

15
3. challenge test
→ 1. microbial test
2. preservative effective test

Testing Frequency

Drug product

Drug substance

long term studies

1st year: every 3 months

2nd year: every 6 months

After the 2nd year: annually through the proposed re-test period/shelf life.

→ 0, 3, 6, 9, 12 (months)

→ 18, 24 (months)

→ every year.

long term studies

a minimum of three time points, including the initial and final time points (e.g., 0, 3, and 6 months),

accelerated storage condition

Storage Conditions: Drug substance

→ should stick specifications
shelf-life. or release → slow ↓

- In general, a drug substance should be evaluated under storage conditions that test its thermal stability and, if applicable, its sensitivity to moisture. ①

any deviation from specifications ↓
failed.

- The long term testing should cover a minimum of 12 months' duration on at least three primary batches at the time of submission.
لـ صون تجاز 12 شهر عشان أقدر اسجل ادوا.

- Testing should be continued for a period of time sufficient to cover the proposed re-test period.

→ لازم
تعمل
stability data
بعد التقييم

- Data from the accelerated storage condition and, if appropriate, from the intermediate storage condition can be used to evaluate the effect of short term excursions outside the label storage conditions (such as might occur during shipping).
in 6 months under harsh conditions

long term stability study
اذا ربيت على ك أسعر بالكي لدرم اسنن اد

stable or not.
drug تبي
لـ عشان أقدر اسجل اذا اد

Storage Conditions : Drug substance

General case

Study	Storage condition	<u>Minimum time period covered by data at submission</u>
<u>Long term*</u>	<u>$25^{\circ}\text{C} \pm 2^{\circ}\text{C}/60\% \text{ RH} \pm 5\% \text{ RH}$</u> or <u>$30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$</u>	<u>12 months</u>
<u>Intermediate**</u>	<u>$30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$</u>	<u>6 months</u>
<u>Accelerated</u>	<u>$40^{\circ}\text{C} \pm 2^{\circ}\text{C}/75\% \text{ RH} \pm 5\% \text{ RH}$</u>	<u>6 months</u>

يقوم على ال
Climatic
zone
اللب
المنطقة

*It is up to the applicant to decide whether long term stability studies are performed at $25 \pm 2^{\circ}\text{C}/60\% \text{ RH} \pm 5\% \text{ RH}$ or $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$.

****If $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$ is the long-term condition, there is no intermediate condition.**

18

← إذا ال accelerated فمع فيها مشكلة بآباء ال intermediate عشان اتقن شوي + support of product
accelerate study. فمعها إذا ال

Storage Conditions : Drug substance

General case

- If long-term studies are conducted at $25 \pm 2^{\circ}\text{C}/60\% \pm 5\% \text{ RH}$ and “**significant change**” occurs at any time during 6 months’ testing at the accelerated storage condition, additional testing at the intermediate storage condition should be conducted and evaluated against significant change criteria.
- “**Significant change**” for a drug substance is defined as failure to meet its specification.
- Testing at the intermediate storage condition should include all tests, unless otherwise justified.
- The initial application should include a minimum of 6 months’ data from a 12-month study at the intermediate storage condition.

Storage Conditions : Drug substance

Drug substances intended for storage in a refrigerator

Study	Storage condition	Minimum time period covered by data at submission
Long term	<u>$5^{\circ}\text{C} \pm 3^{\circ}\text{C}$</u>	<u>12 months</u>
Accelerated	<u>$25^{\circ}\text{C} \pm 2^{\circ}\text{C}/60\% \text{ RH} \pm 5\% \text{ RH}$</u>	<u>6 months</u>

→ stability of accelerated failed between 3-6 months → لازم لأجل re-testing

- If significant change occurs between 3 and 6 months' testing at the accelerated storage condition, **the proposed re-test period should be based on the real time data** available at the long term storage condition.

لأنه تكون على الـ long term data

- If significant change occurs within the first 3 months' testing at the accelerated storage condition, a discussion should be provided to address the effect of short term

our product under question.

← في حالي فسيكون ←

Storage Conditions : Drug substance

~~Drug substances~~ intended for storage in a freezer

Study	<u>Storage condition</u>	Minimum time period covered by data at submission
<u>Long term</u>	- <u>20°C ± 5°C</u>	<u>12 months</u>

- For drug substances intended for storage in a freezer, the re-test period should be based on the real time data obtained at the long term storage condition.
- testing on a single batch at an elevated temperature (e.g., 5°C ± 3°C or 25°C ± 2°C) for an appropriate time period should be conducted to address the effect of short term excursions outside the proposed label storage condition, e.g., during shipping or handling.

Storage Conditions : Drug substance

Drug substances intended for storage below -20°C → Ex: Vaccines

- Drug substances intended for storage below -20°C
should be treated on a case-by-case basis.

Stability Commitment : Drug substance

- When available long term stability data on primary batches do not cover the proposed re-test period granted at the time of approval, a commitment should be made to continue the stability studies post approval in order to firmly establish the re-test period.
- Where the submission includes long term stability data on three production batches covering the proposed re-test period, a post approval commitment is considered unnecessary.

بعد مدة من الدراسة في اسجل المستحسن تبقي كذا لعدم تغطيه
رغم انكل post-approval stability studies
stability test

Stability Commitment : Drug substance

3 حالات في القصة :-

-Otherwise, one of the following commitments should be made:

12 months

1. If the submission includes data from stability studies on at least three production batches, a commitment should be made to continue these studies through the proposed re-test period.

2. If the submission includes data from stability studies on fewer than three production batches, a commitment should be made to continue these studies through the proposed re-test period and to place additional production batches, to a total of at least three, on long term stability studies through the proposed re-test period.

نقطة
أول
دراسة
على ال
Batch
الاول

3. If the submission does not include stability data on production batches, a commitment should be made to²⁴

to place the first 3 production batches on long term stability studies through the proposed re-test period.

Storage Conditions: Drug product

- In general, a drug product should be evaluated under storage conditions (with appropriate tolerances) that test its thermal stability and, if applicable, its sensitivity to moisture or potential for solvent loss.₃ ²
- Stability testing of the drug product after constitution or dilution, if applicable, should be conducted to provide information for the labeling on the preparation, storage condition, and in-use period of the constituted or diluted product (In-use stability).

↓
drug product should be diluted for use
such as : powder for reconstitution

Storage Conditions : Drug product

- **In-use stability testing** should be performed on primary batches as part of the formal stability studies at initial and final time points. → time 12 months. ↓
time zero
- If full shelf life long term data will not be available before submission, **in-use stability testing** should be performed at 12 months or the last time point for which data will be available.
- In general, this testing need not be repeated on commitment batches.
استقرار كل Batch من قبل
Stability
- WHO guidelines requires a **minimum of two batches, at least pilot-scale batches,** to be subjected to the test.

In-Use Stability (WHO guidelines)

Aim:

The purpose of in-use stability testing is to provide information for the labelling on the preparation, storage conditions and utilization period of multidose products after opening, reconstitution or dilution of a solution, e.g. an antibiotic injection supplied as a powder for reconstitution.

In-Use Stability (WHO guidelines)

Number and type of batches:

- A minimum of two batches, at least pilot-scale batches, should be subjected to the test.
- At least one of these batches should be chosen towards the end of its shelf-life.
- If such results are not available, one batch should be tested at the final point of the submitted stability studies.

Storage Conditions : Drug product

-General case

Study	Storage condition	Minimum time period covered by data at submission
<u>Long term</u> *	$25^{\circ}\text{C} \pm 2^{\circ}\text{C}/60\% \text{ RH} \pm 5\% \text{ RH}$ or $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$	12 months
<u>Intermediate</u> **	$30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$	6 months
<u>Accelerated</u>	$40^{\circ}\text{C} \pm 2^{\circ}\text{C}/75\% \text{ RH} \pm 5\% \text{ RH}$	6 months

*It is up to the applicant to decide whether long term stability studies are performed at $25 \pm 2^{\circ}\text{C}/60\% \text{ RH} \pm 5\% \text{ RH}$ or $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$.

**If $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH} \pm 5\% \text{ RH}$ is the long-term condition, there is no intermediate condition.

Storage Conditions : Drug product

-General case

- If long-term studies are conducted at $25 \pm 2^\circ\text{C}/60\% \pm 5\% \text{ RH}$ and “significant change” occurs at any time during 6 months’ testing at the accelerated storage condition, additional testing at the intermediate storage condition should be conducted and evaluated against significant change criteria. 1. impurities
2. change in assay.
3. dissolution
↓
shouldn't have any deviation from the reference
- The initial application should include a minimum of 6 months’ data from a 12-month study at the intermediate storage condition.

Storage Conditions : Drug product

-General case

affecting on shelf-life. →

التركيبات
95 - 105 %

“Significant change” for a drug product is defined as:

1. A 5% change in assay from its initial value; or failure to meet the acceptance criteria for potency when using biological or immunological procedures;
2. Any degradation product's exceeding its acceptance criterion;
3. Failure to meet the acceptance criteria for appearance, physical attributes, and functionality test (e.g., color, phase separation, resuspendibility, caking, hardness, dose delivery per actuation); however, some changes in physical attributes (e.g., softening of suppositories, melting of creams) may be expected under accelerated conditions;

↓
for suppositories or creams

4. Failure to meet the acceptance criteria for pH. ^{→ solution or suspension.}

5. Failure to meet acceptance criteria for dissolution for 12 dosage units.

Storage Conditions : Drug product

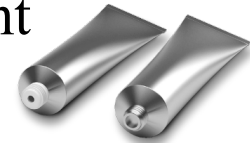
مقاومة للرطوبة أو يتلف أو يتدخل أو الغازات. moisture

Drug products packaged in impermeable containers

Impermeable containers: Containers that provide a permanent barrier to the passage of gases or solvents, e.g., sealed aluminum tubes for semi-solids, sealed glass ampoules for solutions. moisture من ار sensitivity test حاجب للرطوبة



Sensitivity to moisture or potential for solvent loss is **not a concern for drug products packaged in impermeable containers** that provide a permanent barrier to passage of moisture or solvent.



Thus, stability studies for products stored in impermeable containers can be conducted under any controlled or ambient humidity condition.

Storage Conditions : Drug product

- *Drug products packaged in semi-permeable containers*

Semi-permeable containers:

Containers that allow the passage of solvent, usually water, while preventing solute loss.

The mechanism for solvent transport occurs by absorption into one container surface, diffusion² through the bulk of the container material, and ~~the other surface.~~ *and deposition from other surface.*

In such cases, the concentration of drug



(assay) may increase with time.

Examples of semi-permeable containers include:

- *plastic bags and semi-rigid, low-density polyethylene (LDPE) pouches for large volume parenterals (LVPs)*

LDPE ampoules, bottles and vials.

Storage Conditions : Drug product

-Drug products packaged in semi-permeable containers

- Aqueous-based products packaged in semi-permeable containers should be evaluated for potential water loss in addition to physical, chemical, biological, and microbiological stability.
- This evaluation can be carried out under conditions of low relative humidity (RH%).
- Ultimately, it should be demonstrated that aqueous-based drug products stored in semi-permeable containers can withstand low relative humidity environments.
- Other comparable approaches can be developed and reported for non-aqueous, solvent-based products.

Storage Conditions : Drug product

↓ RH changes.

-Drug products packaged in semi-permeable containers

Study	Storage condition	Minimum time period covered by data at submission
<u>Long term*</u>	$25^{\circ}\text{C} \pm 2^{\circ}\text{C}/\underline{40\% \text{ RH}} \pm 5\% \text{ RH}$ or $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/\underline{35\% \text{ RH}} \pm 5\% \text{ RH}$	12 months
<u>Intermediate**</u>	$\underline{30^{\circ}\text{C} \pm 2^{\circ}\text{C}/65\% \text{ RH}} \pm 5\% \text{ RH}$	6 months
<u>Accelerated</u>	$40^{\circ}\text{C} \pm 2^{\circ}\text{C}/\text{not more than (NMT)} 25\% \text{ RH}$	6 months

*It is up to the applicant to decide whether long term stability studies are performed at $25 \pm 2^{\circ}\text{C}/40\% \text{ RH} \pm 5\% \text{ RH}$ or $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/35\% \text{ RH} \pm 5\% \text{ RH}$.

**If $30^{\circ}\text{C} \pm 2^{\circ}\text{C}/35\% \text{ RH} \pm 5\% \text{ RH}$ is the long-term

Storage Conditions : Drug product

Drug products intended for storage in a refrigerator

Study	Storage condition	Minimum time period covered by data at submission
Long term	5°C ± 3°C	12 months
Accelerated	25°C ± 2°C/60% RH ± 5% RH	6 months

- If significant change occurs between 3 and 6 months' testing at the accelerated storage condition, the proposed shelf life should be based on the real time data available at the long term storage condition.
- If significant change occurs within the first 3 months' testing at the accelerated storage condition, a discussion should be provided to address the effect of short term

Storage Conditions : Drug product

Drug products intended for storage in a freezer

Study	Storage condition	Minimum time period covered by data at submission
Long term	- 20°C ± 5°C	12 months

- For drug products intended for storage in a freezer, **the proposed shelf life should be based on the real time data** obtained at the long term storage condition.
- testing on a single batch at an elevated temperature (e.g., 5°C ± 3°C or 25°C ± 2°C) for an appropriate time period should be conducted **to address the effect of short term excursions outside the proposed label** storage condition, e.g., during shipping or handling.

Storage Conditions : Drug-product

-Drug products intended for storage below -20°C

- Drug products intended for storage below -20°C **should be treated on a case-by-case basis.**

Storage Condition : Drug product

TABLE 2 Storage Conditions for Stability Evaluation of Drug Products

Stability Study Type	Stability Storage Conditions	Minimum Time Period Covered by Data at Submission (months)
<i>Marketed Drug Product Intended for <u>Room Temperature Storage Conditions</u></i>		
Long term	25°C ± 2°C, 60% RH ± 5%	12
	RH or 30°C ± 2°C, 65% RH ± 5% RH	12
Intermediate	30°C ± 2°C, 65% RH ± 5% RH	6
Accelerated	40°C ± 2°C, 75% RH ± 5% RH	6
<i>Marketed Drug Product Packaged in <u>Semipermeable Containers</u></i>		
Long term	25°C ± 2°C, 40% RH ± 5% RH or 30°C ± 2°C, 35% RH ± 5% RH	12
Intermediate	30°C ± 2°C, 65% RH ± 5% RH	6
Accelerated	40°C ± 2°C, no more than 25% RH	6
<i>Marketed Drug Product Intended for <u>Storage in Refrigerator</u></i>		
Long term	5°C ± 3°C	12
Accelerated	25°C ± 2°C, 60% RH ± 5% RH	6
<i>Marketed API Intended for <u>Storage in Freezer</u></i>		
Long term	-20°C ± 5°C	12

Summary
for all
Requirements

Storage Conditions : Climatic

- 1 • Climatic Zone I. *Temperate climate*, includes Canada, New Zealand, northern Europe, Russia, United Kingdom
- 2 • Climatic Zone II. *Subtropical and Mediterranean climate*, includes Japan, southern Europe, USA, southern Africa, parts of South America
- 3 • Climatic Zone III. *Hot and dry climate*, includes Argentina, Australia, Botswana, Middle East, northern Africa
- 4 • Climatic Zone IV. *Hot and humid climate*, includes Brazil, much of central Africa including Ghana and Nigeria, Indonesia, Nicaragua, the Philippines, Malaysia
 - o IV-A: Hot and humid climate
 - o IV-B: Hot and very humid climate

Storage Conditions : Climatic Zones

Table 49.2 Long-term test conditions for the various climatic zones, as defined by the World Health Organization (2009)

Climatic zone	Definition	Long-term test conditions	
		Temperature (°C)	Relative humidity (% R.H)
I	Temperate climate	21	45
II	Subtropical and Mediterranean climate	25	60
III	Hot and dry climate	30	35
IVA	Hot and humid climate	30	65
IVB	Hot and very humid climate	30	75

Storage Conditions : Drug product

Climatic Zones III and IV: ICH Q1F

- ICH Q1 A (R2) adopted conditions corresponding to the ICH members (Zone I and II).
- For **other countries in climatic Zone III/IV** 30°C/65% RH was defined as the long-term storage condition in ICH Q1F.
- However, based on new calculations and discussions, some countries in **Climatic Zone IV** have expressed their wish to include a larger safety margin for medicinal products to be marketed in their region than foreseen in ICH Q1F.
- As a consequence, several countries and regions have revised their own stability testing guidelines, defining up to **30°C/75 % RH** as the long-term storage conditions for hot and humid regions

Stability Commitment

- When available long term stability data on primary batches do not cover the proposed shelf life granted at the time of approval, **a commitment should be made to continue the stability studies post approval** in order to firmly establish the re-test period.
- Where the submission includes long term stability data on three production batches covering the proposed shelf life , **a post approval commitment is considered unnecessary.**

Stability Commitment

Otherwise, one of the following commitments should be made:

1. If the submission includes data from stability studies on at least three production batches, **a commitment should be made to continue these studies through the proposed shelf life** .
2. If the submission includes data from stability studies on fewer than three production batches, **a commitment should be made to continue these studies through the proposed shelf life and to place additional production batches, to a total of at least three**, on long term stability studies through the proposed shelf life .
3. If the submission does not include stability data on production batches, **a commitment should be made to**

EVALUATION FOR STABILITY DATA

ICH GUIDELINE Q1E

General Principles

- The purpose of a stability study is to establish, based on testing a minimum of three batches of the drug substance or product a retest period or shelf life and label storage instructions applicable to all future batches manufactured and packaged under similar circumstances.
- The degree of variability of individual batches affects the confidence that a future production batch will remain within acceptance criteria throughout its retest period or shelf life.

General Principles

- it is important that the drug product be formulated with the intent to provide 100 percent of the labeled amount of the drug substance at the time of batch release.
- If assay at the time of release for stability batches is higher than 100 percent → the shelf life proposed in the application can be overestimated
- If the assay value of a batch is lower than 100 percent of label claim at the time of batch release, it might fall below the lower acceptance criterion before the end of the proposed shelf life.

General Principles

- The stability information should include, as appropriate, results from the physical, chemical, biological, and microbiological tests, including those related to particular attributes of the dosage form (for example, dissolution rate for solid oral dosage forms).
- The adequacy of the mass balance should be assessed.
- Factors that can cause an apparent lack of mass balance should be considered, including, for example:
 - 1) ➤ the mechanisms of degradation
 - 2) ➤ the stability-indicating capability of the analytical procedures
 - 3) ➤ inherent variability of the analytical procedures.