

# Container Closure Integrity: Regulations, Test Methods, Application

## Introduction

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# Introduction

- Terms, definitions, and Concepts
- Maximum Allowable Leak Limit (MALL)
- Inherent package integrity
- Package integrity profile

## ***IN SCOPE of USP<1207> - Focus of the course***

Sterile pharmaceutical product packaging (SVP, LVP)

***Examples:***

Vials or bottles closed with elastomeric closures or screw-thread caps

Form-fill-seal plastic or glass ampules

Syringes or cartridges

Flexible bags or pouches.

Packages for some drug/device combination products (e.g., autoinjectors)

## ***OUT OF SCOPE of USP<1207> - methodologies apply***

Packaging systems involved in prep, storage, manufacture

***Examples:*** API, intermediate/final bulk

Sterile diagnostic products or medical devices

Some packages for sterile drug/device combo products

Primary packages with porous barrier materials designed to allow air or gas sterilant passage

**Product:**

- ***Pharmaceutical formulation***
  - Principles apply to containers for API, bulk, intermediates
- ***Packaged headspace***
  - Air or nonreactive gases
  - At specified water vapor content
  - At ambient or sub-ambient pressures

**Package (aka Container-closure):**

- ***Primary package components***
  - In direct product contact (or may be)
- ***Secondary package components critical for ensuring package assembly***
  - *E.g.*, aluminum crimp seal on vial/stopper

**Product-Package:**

- ***The primary package with critical secondary components*** (the container-closure system)
- AND**
- ***The packaged contents*** (the product)

## **Leak:**

A **gap** or **breach** in the container capable of permitting the passage of liquid or gas. Otherwise known as “leak path.”

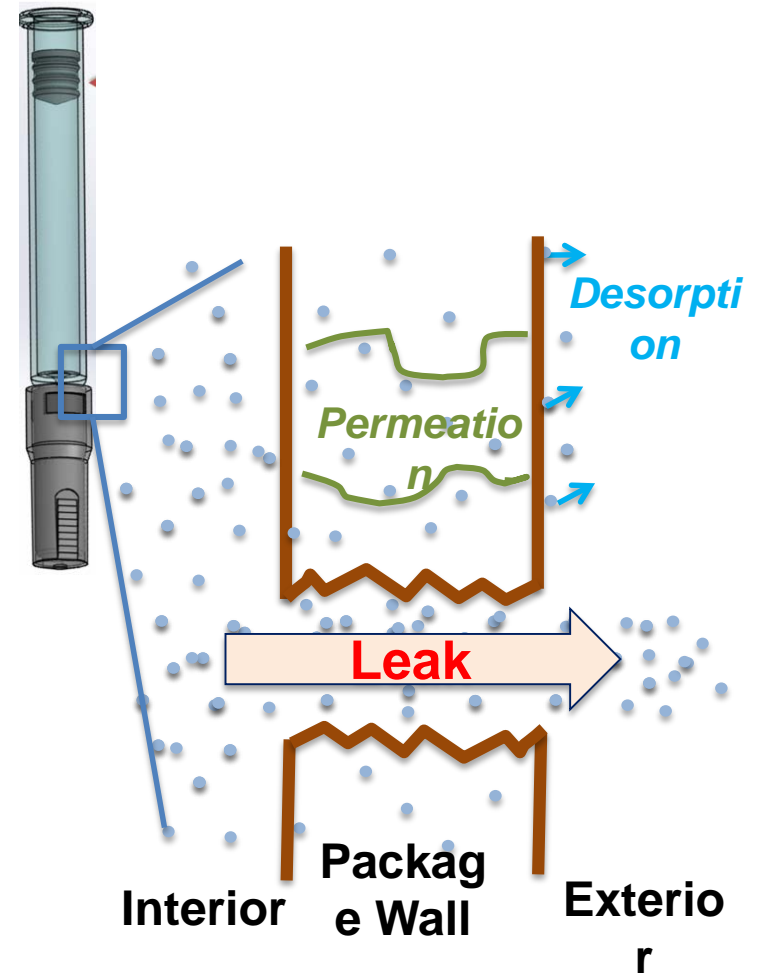
## **Leakage:**

1. The unintentional entry or escape of matter (solid, liquid or gas) through a breach in a package wall or through a gap between package components.
2. The leaking matter itself.

## Permeation

*The passage of fluid (e.g., gas) into, through, and out of a nonporous package wall.*

*Permeation (**NOT leakage**) occurs when only a small fraction of molecules is able to move through a barrier by way of any one hole.*



## **Sterile product package integrity or “container closure integrity” (CCI)**

**Definition: The ability of a package to...**

***Keep good stuff in, and  
Keep bad stuff out***

**“A package with integrity”**

***Does not mean***

***the package has passed or is able to pass a  
Microbial ingress test, or product sterility test***

## **Microbial Ingress is a PROBABILISTIC EVENT**

**Difficult to control, predict, measure**

### **FACTORS**

<b>Leak path</b>	<b>size/shape/length/material/blockage</b>
<b>Ingress test parameters</b>	<b>time/pressure/temp</b>
<b>Microorganism</b>	<b>type/size</b>
<b>Liquid tracer</b>	<b>chemistry/concentration</b>
<b>Carrier fluid</b>	<b>viscosity/surface tension/solvent</b>
<b>Visual detection</b>	<b>human variables/inspection conditions</b>
<b>Instrumental detection</b>	<b>instrument/test parameters</b>



## CONSIDER

***IF*** windows keep out birds, ***THEN*** why not detect defective windows by checking homes for birds?



?

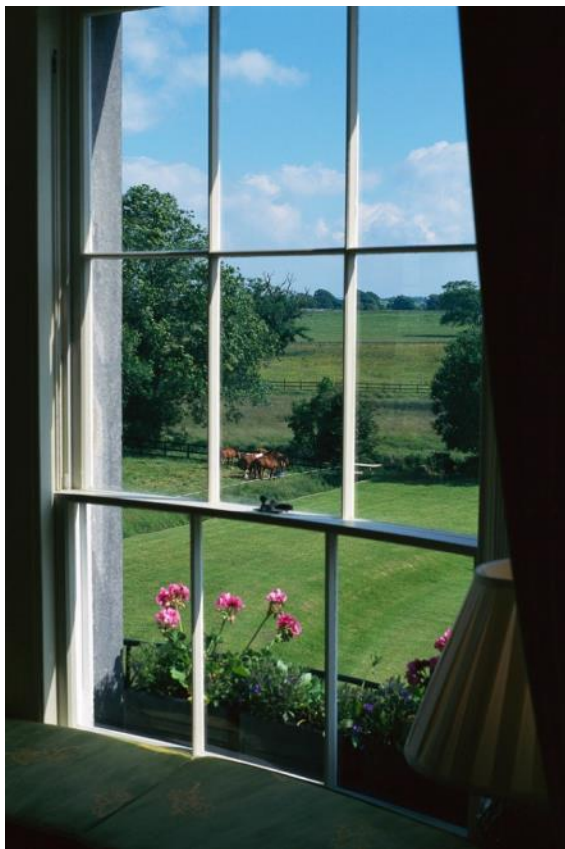


## Package integrity :

- **IS NOT** passing microbial ingress or product sterility tests
- **IS** the absence of a gap/defect that risks product quality
- **IS** the conformance of the package to the maximum allowable leakage limit (i.e., critical leak)
  - Product quality requirements define MALL

Testing goals may vary during the product life cycle

## INSTEAD of Checking for Bats.....



***Design and make*** windows that close well based on meaningful, reliable tests

***Test*** for absence of defects that could permit birds

***Monitor*** to ensure control over materials, processes

**“A package with integrity”**

**Means that**

*Gaps/breaches that **COULD** risk product quality are absent*

*i.e., The package meets the*

***MAXIMUM ALLOWABLE LEAKAGE LIMIT (MALL)***

**\*\*\*\***

***What’s the difference?***

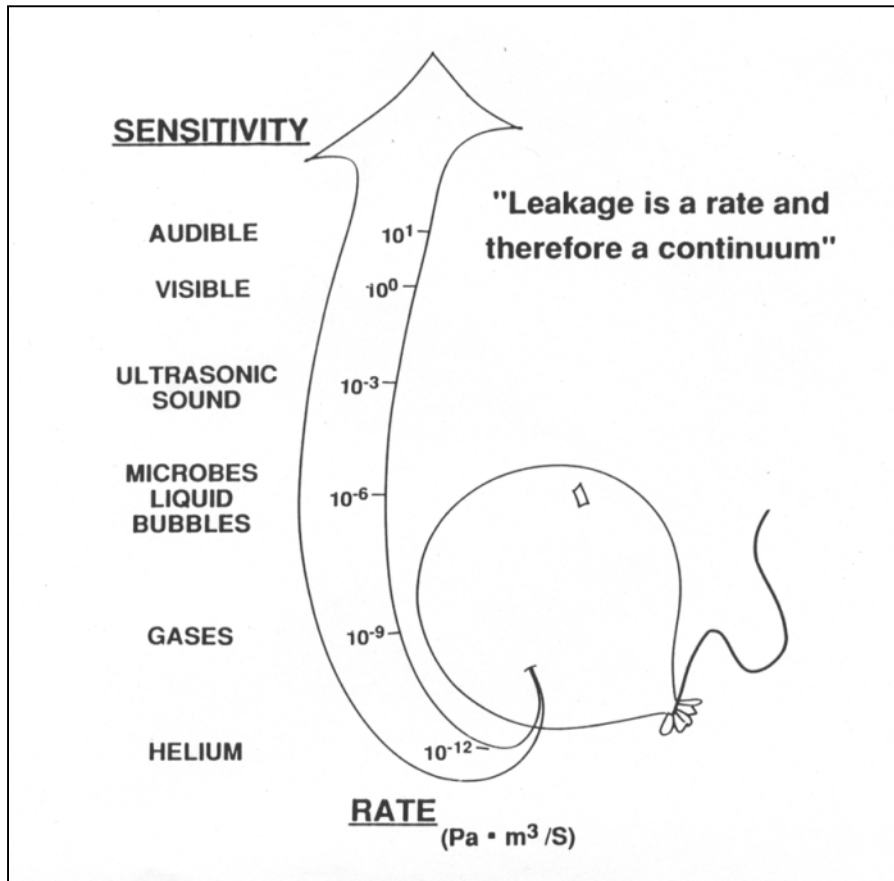
## Maximum Allowable Leakage Limit (MALL)

*is that smallest gap or leak rate that puts  
product quality at risk*

(sometimes called the 'critical leak')



## All physically mated closure systems\* leak to some degree



**Smallest** leaks only allow gas flow

**Larger** leaks may also allow liquid flow

**Largest** leaks may also allow microbial ingress

\*physicochemically bonded seals may only allow permeation

## Sterile product package integrity (CCI)

Category	Leaks of concern	Product quality risks
1	Capable of allowing entry of microorganisms	Failure of product sterility
2	Capable of allowing escape of product dosage form, or entry of external of liquids/solids	Failure of relevant physicochemical quality attributes
3	Capable of allowing change in gas headspace content e.g., escape of nitrogen, loss of vacuum, entry of oxygen, water vapor, or air	Failure of relevant physicochemical quality attributes, And/or hindrance of product access by end-user.

***What is the maximum allowable leakage limit (MALL)***

***For categories 1 and 2?***

Prevention of **microbial ingress**

Prevention of **product loss** (liquid or solid) or **external contamination** by liquid or solid matter



## Smallest leak to first allow ingress determination

- Lee Kirsch, et al, PDA J Pharm Sci & Technol, Vol. 51, No. 5, 1997

### • Comparison of orifice helium leak rate vs microbial and liquid tracer ingress

#### – Glass micro-pipettes through wall of stoppered glass vial

- Sized via helium mass spec
- 0.1 to 10 $\mu$ m diameter

#### – Microbial challenge by immersion + liquid tra

- 10<sup>8</sup> to 10<sup>10</sup> *P. diminuta* and *E. coli* cfu/mL
- Tween 80 additive
- Mg ion tracer for liquid path verification
  - Detection by atomic absorption

#### – Challenge conditions

- Airlock elimination procedure
  - Water bath immersion 60°C 2hr, then 25°C 1hr
- 24 hr immersion, ambient pressure

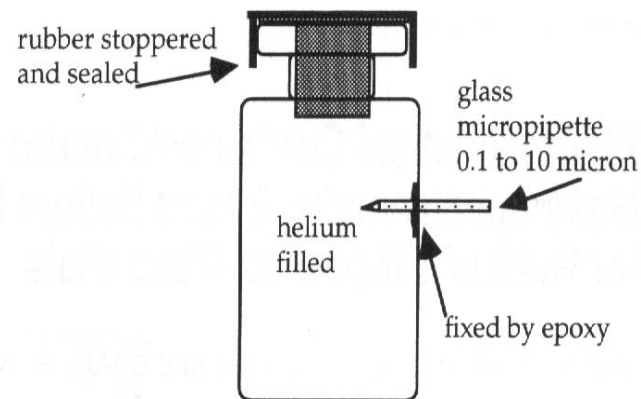


Figure 1—Schematic description of the modified pharmaceutical vials used as test units for the evaluation of mass spectrometry-based helium leak rate measurements.

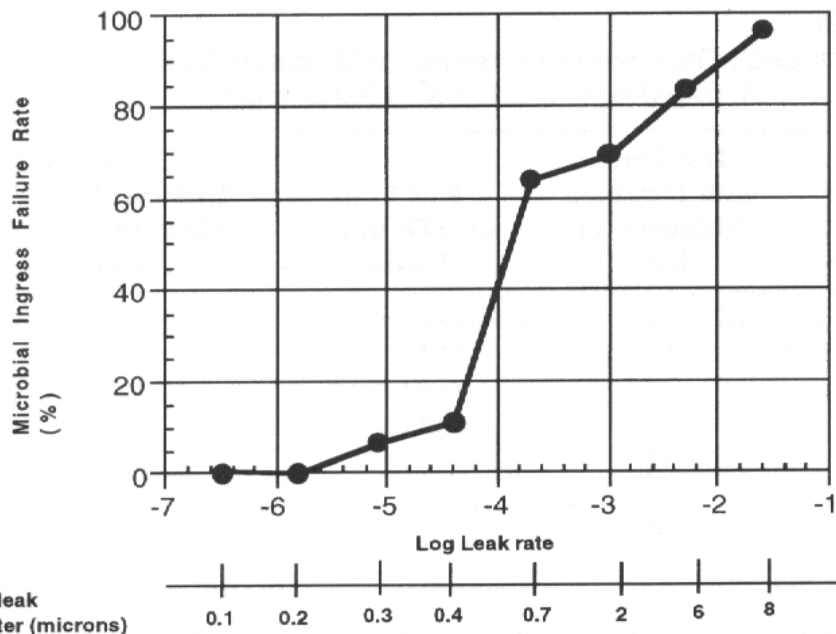
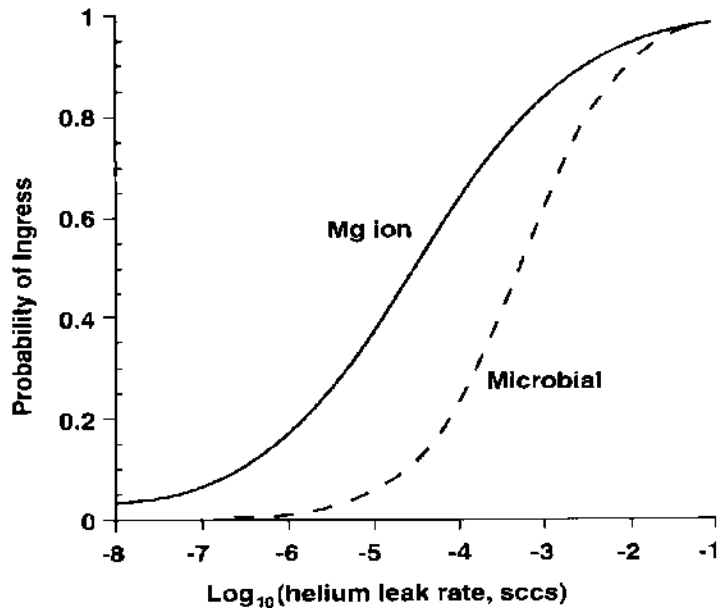


Figure 2—The correlation of microbial failure rate (%) and the mean logarithm of the absolute leak rate and nominal leak diameter for modified SVPs. The absolute leak rate (standard cubic centimeters per second) was determined by mass spectrometry-based helium leak rate detection. Microbial failure was measured by microbial ingress after 24 hour immersion in a bath (37°C) containing  $10^8$  to  $10^{10}$  *P. diminuta* and *E. coli* organisms/mL and a 13 day, 35°C incubation.

- **Microbial ingress risk dropped dramatically at**
  - **Log -3.8 sccs ( $< \sim 1\mu\text{m}$ )**
- **Low risk of ingress ( $< 0.10$ ) at helium leak rate of**
  - **$6 \times 10^{-6}$  mbarL/s**

**Figure 1: Logistical regression models describing the probability of microbial or liquid tracer (Mg ion) as a function of the logarithm of the helium leak rates. Curves were generated using Equation 1 and parameters estimated with the logistical regression platform in the software JMP (10).**



- **Microbial ingress requires liquid flow**
  - Increased liquid flow equals increased microbial ingress risk
- **Liquid flow  $\neq$  microbial ingress**

## MALL as a function of leak path morphology and test conditions

Study Author	Challenge medium	Challenge microbe	Challenge path	Challenge conditions	Microbial ingress first observed
Kirsch JPDA '97-'99	Liquid	<i>P. diminuta</i> <i>E. coli</i>	Glass micro-pipette thru vial wall	Airlock elimination step + 24 hr ambient	0.3 µm orifice
Burrell JPDA 2000	Liquid	<i>E. Coli</i>	Poly-coated glass micro-tube thru stopper	ISO closure reseal: 30 min 22" Hg + 30 min ambient	10 µm ID tube
Morrical JPDA 2007	Liquid	<i>Serratia marcescens</i>	Metal plate micro-hole in stopper	-0.4 bar 1 hr +0.4 bar 1 hr	4 µm orifice
Morrical JPDA 2007	Liquid	<i>Serratia marcescens</i>	Copper wire between stopper/vial	-0.4 bar 1 hr +0.4 bar 1 hr	20 µm OD wire
Keller <i>J Applied Pkgs Res</i> 2006	Aerosol	<i>P. Fragi</i>	Nickel micro-tube in 3mL vial	Varied: -20 kPa to +20 kPa 4 to 37°C	5 µm ID tube

Kirsch reported smallest leak (nominal hole size) that first demonstrated:

- **microbial ingress:** **0.2 - 0.3  $\mu\text{m}$**
- **aqueous liquid passage:** **0.1  $\mu\text{m}$ \***

\*Absolute cut-off was not defined as smaller leaks were not evaluated

- **Liquid presence** in the leak path was **required**, but **did not guarantee** microbial ingress
- **Airborne microbial ingress** only possible with larger leaks

## MALL size of “**Real leaks**” is **undefined**

- Real leak paths are not holes, tubes, pipettes
- Natural defects are long, complex, irregular channels
- Defects consist of actual package materials
- Air pockets, debris, product may block leak flow or microbial ingress

Choosing the critical leak size (rate) that will ensure product sterility and prevent product formulation loss is a ***SCIENCE AND RISK BASED DECISION***

- In general, for **nonporous rigid packages** such as
  - Parenteral vials, bottles
  - Syringes, cartridges
  - Form fill seal glass/plastic ampoules
  - Drug/Device package systems (e.g., autoinjectors)

Helium leakages rate of **< 6 E-6 mbarL/s**  
(leakage through an orifice of about 0.1 to 0.3  $\mu\text{m}$ )  
have a **low risk** of microbial ingress or liquid product loss.

***Adopting this MALL for such product-packages may eliminate the need for microbial ingress or liquid challenge studies as a function of leak size.***

## Ingress or product loss risk is not as well defined

**For other package systems** such as Flexible polymeric packages

**For leak types/morphologies** more complex or lengthy

**For products more likely to leak** such as cosolvent systems

**The MALL is UNIQUE for each product-package**

***A SCIENCE AND RISK BASED DECISION***

**Determine the risk of microbial ingress or liquid passage as a function of defect size/type.**

## ***What is the maximum allowable leakage limit (MALL) for Category 3?***

Prevention of **change in gas headspace content** that risks product quality, and/or risks ease of product access

e.g., N<sub>2</sub> escape; vacuum loss; entry of O<sub>2</sub>, H<sub>2</sub>O vapor, or air

**The MALL is UNIQUE for each product-package**

***A SCIENCE BASED DECISION***

***Consider***

Headspace quality requirements: Initial and at expiry

Package headspace volume

Package permeation

Product-package storage, distribution environment



## ***What is the “in-use” maximum allowable leakage limit (MALL) for multiple dose product packages?***

An in-use sub-category of categories 1, 2, 3.

e.g., Multiple dose vials or cartridges

Prevention of product loss or microbial ingress between and during dosage access

**The MALL is UNIQUE for each product-package.**

***A SCIENCE AND RISK BASED DECISION***

***Determine***

Attempts of product access – quantity and mode

Risk of microbial ingress and/or product loss

## The MALL is based on product quality requirements

1. **Prevention of microbial ingress** to ensure product sterility
2. **Prevention of product formulation loss** and **product formulation contamination by external solids/liquids** to ensure conformance to relevant physicochemical product quality attributes.
3. **Prevention of headspace content change** to ensure conformance to relevant physicochemical product quality attributes, and to assure product access.

**Establishing the MALL is a science-based  
and often a risk-based decision**



# Inherent Package Integrity

***The leakage rate (or the equivalent leak size) of a well-assembled package using no-defect components.***

***Best-case leak tightness***, given anticipated variables:

Material composition, dimension, processing, and assembly.  
Final product storage, distribution and use.

Determined during product-package **R&D, validation**

**Acceptable inherent package integrity conforms to the specific product-package MALL**

## Package integrity profile

- **Ongoing database** – *Product life-cycle leak and seal quality tests' results*
- Offers a risk management tool of package integrity assurance
- **Demonstrates integrity as a function of ongoing, operative variations**
  - Package component design/material
  - Package assembly
  - Package and package component processing
  - Package storage, distribution, stability

A package with integrity is one with an **absence of gaps/breaches** in packages that **COULD** risk product quality by allowing solid/liquid contaminant ingress, product formulation loss, and in some cases, headspace change.

i.e., Meets the **Maximum Allowable Leakage Limit**

**Reporting leak size/rate** can be done a variety of ways.

- **Key is to be clear, noting methodology**
- **Units of measure should be relevant to the MALL**

**Microbial ingress/liquid tracer tests** are probabilistic methods that cannot solely be relied upon for package integrity assurance.

*Tests may miss harmful leak paths*

Develop/validate **CC system** having **inherent package integrity** that meets the product **MALL specification**

Use ongoing **product package integrity profile data** to monitor for and minimize integrity failure risks