

Status: Currently Official on 12-Feb-2025  
 Official Date: Official as of 01-Aug-2023  
 Document Type: General Chapter  
 DocId: GUID-BA3755E4-77AA-4DEB-8FE2-4FC78C587E9E\_3\_en-US  
 DOI: [https://doi.org/10.31003/USPNF\\_M99694\\_03\\_01](https://doi.org/10.31003/USPNF_M99694_03_01)  
 DOI Ref: fdg70

© 2025 USPC  
 Do not distribute

## ⟨905⟩ UNIFORMITY OF DOSAGE UNITS

**Change to read:**

This general chapter is harmonized with the corresponding texts of the *European Pharmacopoeia* and the *Japanese Pharmacopoeia*. Portions of the general chapter text that are national *USP* text, and are not part of the harmonized text, are marked with symbols (♣) to specify this fact.

[NOTE—In this chapter, *unit* and *dosage unit* are synonymous.]

To ensure the consistency of dosage units, each unit in a batch should have a drug substance content within a narrow range around the label claim. Dosage units are defined as dosage forms containing a single dose or a part of a dose of drug substance in each unit. The uniformity of dosage units specification is not intended to apply to ▲solutions,▲ (USP 1-Aug-2023) suspensions, emulsions, or gels in unit-dose containers intended for ▲local action following▲ (USP 1-Aug-2023) external, cutaneous administration.

The term “uniformity of dosage unit” is defined as the degree of uniformity in the amount of the drug substance among dosage units. Therefore, the requirements of this chapter apply to each drug substance being comprised in dosage units containing one or more drug substances, unless otherwise specified elsewhere in this Pharmacopoeia.

The uniformity of dosage units can be demonstrated by either of two methods, *Content Uniformity* or ♣*Weight Variation* (see [Table 1](#)). The test for *Content Uniformity* of preparations presented in dosage units is based on the assay of the individual content of drug substance(s) in a number of dosage units to determine whether the individual content is within the limits set. The *Content Uniformity* method may be applied in all cases.

The test for ♣*Weight Variation* is applicable for the following dosage forms:

(W1)	Solutions enclosed in unit-dose containers and into soft capsules;
(W2)	Solids (including powders, granules, and sterile solids) that are packaged in single-unit containers and contain no active or inactive added substances;
(W3)	Solids (including sterile solids) that are packaged in single-unit containers, with or without active or inactive added substances, that have been prepared from true solutions and freeze-dried in the final containers and are labeled to indicate this method of preparation; and
(W4)	Hard capsules, uncoated tablets, or film-coated tablets, containing 25 mg or more of a drug substance comprising 25% or more, by weight, of the dosage unit or, in the case of hard capsules, the capsule contents, except that uniformity of other drug substances present in lesser proportions is demonstrated by meeting the requirements for <i>Content Uniformity</i> .

The test for *Content Uniformity* is required for all dosage forms not meeting the above conditions for the ♣*Weight Variation* test.<sup>1</sup>

**Table 1. Application of Content Uniformity (CU) and Weight Variation (WV) Tests for Dosage Forms**

Dosage Form	Type	Subtype	Dose & Ratio of Drug Substance	
			≥25 mg and ≥25%	<25 mg or <25%
Tablets	Uncoated	—	WV	CU
	Coated	Film	WV	CU

Dosage Form	Type	Subtype	Dose & Ratio of Drug Substance	
			≥25 mg and ≥25%	<25 mg or <25%
		Others	CU	CU
Capsules	Hard	—	WV	CU
	Soft	Suspension, emulsion, or gel	CU	CU
		Solutions	WV	WV
Solids in single-unit containers	Single component	—	WV	WV
	Multiple components	Solution freeze-dried in final container	WV	WV
		Others	CU	CU
Solutions in unit-dose containers and into soft capsules,	—	—	WV	WV
Others <sup>a</sup> (USP 1-Aug-2023)	—	—	CU	CU

<sup>a</sup> Dosage forms not addressed by the other categories in this table including but not limited to suppositories, transdermal systems (patches), and semisolid preparations applied cutaneously and intended for systemic distribution of the drug substance.

### CONTENT UNIFORMITY

Select not fewer than 30 units, and proceed as follows for the dosage form designated.

Where different procedures are used for assay of the preparation and for the *Content Uniformity* test, it may be necessary to establish a correction factor to be applied to the results of the latter.

#### Solid Dosage Forms

Assay 10 units individually using an appropriate analytical method. Calculate the acceptance value (see [Table 2](#)).

#### Liquid or Semi-Solid Dosage Forms

Assay 10 units individually using an appropriate analytical method. Carry out the assay on the amount of well-mixed material that is removed from an individual container in conditions of normal use, and express the results as delivered dose. Calculate the acceptance value (see [Table 2](#)).

#### Calculation of Acceptance Value

Calculate the acceptance value by the formula:

$$|M - \bar{X}| + ks$$

in which the terms are as defined in [Table 2](#).

**Table 2**

Variable	Definition	Conditions	Value
$\bar{X}$	Mean of individual contents ( $x_1, x_2, \dots, x_n$ ), expressed as a percentage of the label claim	—	—
$x_1, x_2, \dots, x_n$	Individual contents of the units tested, expressed as a percentage of the label claim	—	—

Variable	Definition	Conditions	Value
n	Sample size (number of units in a sample)	—	—
k	Acceptability constant	If n = 10, then k =	2.4
		If n = 30, then k =	2.0
s	Sample standard deviation	—	$\left[ \frac{\sum_{i=1}^n (x_i - \bar{X})^2}{n-1} \right]^{\frac{1}{2}}$
RSD	Relative standard deviation (the sample standard deviation expressed as a percentage of the mean)	—	$100s/\bar{X}$
M (case 1) to be applied when $T \leq 101.5$	Reference value	If $98.5\% \leq \bar{X} \leq 101.5\%$ , then	$M = \bar{X}$ (AV = ks)
		If $\bar{X} < 98.5\%$ , then	$M = 98.5\%$ (AV = $98.5 - \bar{X} + ks$ )
		If $\bar{X} > 101.5\%$ , then	$M = 101.5\%$ (AV = $\bar{X} - 101.5 + ks$ )
M (case 2) to be applied when $T > 101.5$	Reference value	If $98.5 \leq \bar{X} \leq T$ , then	$M = \bar{X}$ (AV = ks)
		If $\bar{X} < 98.5\%$ , then	$M = 98.5\%$ (AV = $98.5 - \bar{X} + ks$ )
		If $\bar{X} > T$ , then	$M = T\%$ (AV = $\bar{X} - T + ks$ )
Acceptance value (AV)	—	—	General formula: $ M - \bar{X}  + ks$ (Calculations are specified above for the different cases.)
L1	Maximum allowed acceptance value	—	L1 = 15.0 unless otherwise specified
L2	Maximum allowed range for deviation of each dosage unit tested from the calculated value of M	On the low side, no dosage unit result can be less than $[1 - (0.01)(L2)]M$ , while on the high side, no dosage unit result can be greater than $[1 + (0.01)(L2)]M$ . (This is based on an L2 value of 25.0.)	L2 = 25.0 unless otherwise specified
T	Target content per dosage unit at the time of manufacture, expressed as a percentage of the label claim. Unless otherwise stated, T is 100.0%, or T is the manufacturer's approved target content per dosage unit.	—	—

**WEIGHT VARIATION**

Carry out an assay for the drug substance(s) on a representative sample of the batch using an appropriate analytical method. This value is result A, expressed as percentage of label claim (see *Calculation of Acceptance Value*). Assume that the concentration (weight of drug substance per weight of dosage unit) is uniform. Select not fewer than 30 dosage units, and proceed as follows for the dosage form designated.

**Uncoated or Film-Coated Tablets**

Accurately weigh 10 tablets individually. Calculate the content, expressed as percentage of label claim, of each tablet from the weight of the individual tablet and the result of the Assay. Calculate the acceptance value.

**Hard Capsules**

Accurately weigh 10 capsules individually, taking care to preserve the identity of each capsule. Remove the contents of each capsule by a suitable means. Accurately weigh the emptied shells individually, and calculate for each capsule the net weight of its contents by subtracting the weight of the shell from the respective gross weight. Calculate the drug substance content of each capsule from the net weight of the individual capsule content, and the result of the Assay. Calculate the acceptance value.

**Soft Capsules**

Accurately weigh 10 intact capsules individually to obtain their gross weights, taking care to preserve the identity of each capsule. Then cut open the capsules by means of a suitable clean, dry cutting instrument such as scissors or a sharp open blade, and remove the contents by washing with a suitable solvent. Allow the occluded solvent to evaporate from the shells at room temperature over a period of about 30 minutes, taking precautions to avoid uptake or loss of moisture. Weigh the individual shells, and calculate the net contents. Calculate the drug substance content in each capsule from the weight of product removed from the individual capsules and the result of the Assay. Calculate the acceptance value.

**Solid Dosage Forms Other Than Tablets and Capsules**

Proceed as directed for *Hard Capsules*, treating each unit as described therein. Calculate the acceptance value.

**Liquid Dosage Forms**

Accurately weigh the amount of liquid that is removed from each of 10 individual containers in conditions of normal use. If necessary, compute the equivalent volume after determining the density. Calculate the drug substance content in each container from the mass of product removed from the individual containers and the result of the Assay. Calculate the acceptance value.

**Calculation of Acceptance Value**

Calculate the acceptance value as shown in *Content Uniformity*, except that the individual contents of the units are replaced with the individual estimated contents defined below.

$x_1, x_2, \dots, x_n$  = individual estimated contents of the units tested, where  $x_i = w_i \times A/\bar{W}$

$w_1, w_2, \dots, w_n$  = individual weights of the units tested

A = content of drug substance (% of label claim) obtained using an appropriate analytical method

$\bar{W}$  = mean of individual weights ( $w_1, w_2, \dots, w_n$ )

**CRITERIA**

Apply the following criteria, unless otherwise specified.

**Solid, Semi-Solid, and Liquid Dosage Forms**

The requirements for dosage uniformity are met if the acceptance value of the first 10 dosage units is less than or equal to L1%. If the acceptance value is > L1%, test the next 20 units, and calculate the acceptance value. The requirements are met if the final acceptance value of the 30 dosage units is  $\leq$  L1%, and no individual content of any dosage unit is less than  $[1 - (0.01)(L2)]M$  nor more than  $[1 + (0.01)(L2)]M$  as specified in the *Calculation of Acceptance Value* under *Content Uniformity* or under *Weight Variation*. Unless otherwise specified, L1 is 15.0 and L2 is 25.0.

<sup>1</sup> European Pharmacopoeia and Japanese Pharmacopoeia text not accepted by the United States Pharmacopeia: Alternatively, products listed in item (4) above that do not meet the 25 mg/25% threshold limit may be tested for uniformity of dosage units by Mass Variation instead of the Content Uniformity test if the concentration relative standard deviation (RSD) of the drug substance in the final dosage units is not more than 2%, based on process validation data and development data, and if there has been regulatory approval of such a change. The

concentration RSD is the RSD of the concentration per dosage unit (w/w or w/v), where concentration per dosage unit equals the assay result per dosage unit divided by the individual dosage unit weight. See the RSD formula in [Table 2](#).

**Auxiliary Information** - Please [check for your question in the FAQs](#) before contacting USP.

Topic/Question	Contact	Expert Committee
<905> UNIFORMITY OF DOSAGE UNITS	<a href="#">Ravi Dasari</a> Statistician	GCDF2020 General Chapters - Dosage Forms 2020

**Most Recently Appeared In:**

Pharmacopeial Forum: Volume No. PF 35(3)

**Current DocID:** [GUID-BA3755E4-77AA-4DEB-8FE2-4FC78C587E9E\\_3\\_en-US](#)

**Previous DocID:** [GUID-BA3755E4-77AA-4DEB-8FE2-4FC78C587E9E\\_2\\_en-US](#)

**DOI:** [https://doi.org/10.31003/USPNF\\_M99694\\_03\\_01](https://doi.org/10.31003/USPNF_M99694_03_01)

**DOI ref:** [fdg70](#)

OFFICIAL