BRIEFING

Uniformity of Dosage Units, USP 27 page 2396 and page 1587 of PF 28(5) [Sept.–Oct. 2002]. The United States Pharmacopeia is the coordinating pharmacopeia in the efforts toward international harmonization of the specifications provided in this general test chapter. The Stage 5B text for this chapter has been signed-off by the Pharmacopeial Discussion Group members. The text presented herein represents a merger of the Stage 5B and the national USP text. This combined text, which is being published in this issue of PF for information only, not for public comment, is scheduled for publication in USP 28–NF 23, with a scheduled implementation date of 1 April 2006. Earlier implementation by individual companies may be done at their discretion.

Major differences between this current text and that of the STAGE 4 “merged” draft that was published on pages 1595–1599 of PF 28(5) [Sept.–Oct. 2002] are listed below.

1. The text discussing harmonization in the first paragraph of the STAGE 4 draft has been deleted in this current draft because the revised chapter will not officially be harmonized until the EP adopts the global text in the European Pharmacopoeia, Supplement 5.2, official December 2004 and the JP adopts the global text in the JP XV, official April 2006. At that time USP will republish this text with the addition of symbols for the national text throughout and with the explanatory introductory text, “Portions of this general chapter have been harmonized with the corresponding texts of the European Pharmacopoeia or the Japanese Pharmacopoeia. Those portions that are not harmonized are marked with symbols (♦ ♦) to specify this fact."

2. The term “active ingredient” in the STAGE 4 draft has been replaced globally by the preferred term “drug substance”.

3. In the current listing of dosage forms that require testing by Content Uniformity (see item C3 in the text), suspensions or emulsions or gels in unit-dose containers or in soft capsules that are intended for systemic administration only are distinguished from those that are intended for external, cutaneous administration.

4. A new Table 1, linking the various dosage forms to the appropriate Uniformity of Dosage Units test procedure, has been added to the current text.

5. The text listing those dosage forms for which Weight Variation is applicable has been revised for clarity.

6. The definition for the reference value M in Table 2 has been revised to indicate coverage of cases where the target test sample amount (amount of drug substance) at time of manufacture, T, is either (a) less than or equal to 101.5%, or (b) greater than 101.5%.

7. The variable U, an overage factor that was indicated for use in cases where the drug product was formulated with an overage of drug substance to compensate for stability losses during product shelf life, has been deleted from the current text. The addition of such overages is no longer considered to be acceptable.

8. The text regarding the term Acceptance Value, AV, has been expanded for clarity.

9. The text relating to the variables L1 and L2 in Table 2 has been expanded, and the values “0.75M” and “1.25M” have been replaced by formulas that clarify how the computations are made.

10. The first paragraph in the Weight Variation section of the current text contains new text for clarity. This text indicates that the assay results, designated as A, are to be expressed as “% of label claim”.

11. Revised text stating that the drug substance concentration is assumed to be uniform (throughout the batch) has been added to the first paragraph under Weight Variation. For the purpose of this chapter, tablet concentration, unlike tablet content, is assumed to be independent of individual dosage unit weight when Weight Variation is applicable.
Change to read:

(905) UNIFORMITY OF DOSAGE UNITS

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

The uniformity of dosage units can be demonstrated by either of two methods, weight variation or content uniformity. The requirements of this chapter apply both to dosage units containing a single active ingredient and to dosage units containing two or more active ingredients; unless otherwise specified in the individual monograph, they apply individually to each active ingredient in the product.

Content Uniformity requirements may be applied in all cases. The test for Content Uniformity is required for:

1. coated tablets, other than film-coated tablets containing 50 mg or more of an active ingredient that comprises 50% or more (by weight) of one tablet;
2. transdermal systems;
3. suspensions in single-unit containers or in soft capsules;
4. inhalations (other than solutions for inhalation packaged in glass or plastic ampuls, intended for use in nebulizers) packaged in premetered dosage units (For inhalers and premetered dosage units labeled for use with a named inhalation device, also see Aerosols, Metered-Dose Inhalers, and Dry Powder Inhalers (601));
5. solids (including sterile solids) that are packaged in unit-dose containers and that contain active or inactive added substances, except that the test for Weight Variation may be applied in the special situations stated below; and
6. suppositories.

When the test for Content Uniformity is not required, the test for Weight Variation may be applied in any of the following situations:

1. products containing 50 mg or more of an active ingredient comprising 50% or more, by weight, of the dosage unit or, in the case of hard capsules, the capsule contents, except that uniformity of other active ingredients present in lesser proportions is demonstrated by meeting Content Uniformity requirements;
2. liquid-filled soft capsules other than soft capsules containing suspensions;
3. solids (including sterile solids) that are packaged in single-unit containers and contain no added substances, whether active or inactive;
4. solids (including sterile solids) that are packaged in single-unit containers, with or without added substances, whether active or inactive, that have been prepared from true solutions and freeze-dried in the final containers and are labeled to indicate this method of preparation; and
5. solutions for inhalation packaged in glass or plastic ampuls, intended for use in nebulizers, oral solutions, and syrups when these articles are packaged in single-unit containers.
WEIGHT VARIATION

For the determination of dosage-unit uniformity by weight variation, select not fewer than 30 units, and proceed as follows for the dosage form designated. [ NOTE — Specimens other than these test units may be drawn from the same batch for Assay determinations. ]

Uncoated and Film-Coated Tablets — Weigh accurately 10 tablets individually. From the result of the Assay, obtained as directed in the individual monograph, calculate the content of active ingredient in each of the 10 tablets, assuming homogeneous distribution of the active ingredient.

Hard Capsules — Weigh accurately 10 capsules individually, taking care to preserve the identity of each capsule. Remove the contents of each capsule by a suitable means. Weigh accurately 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. From the results of the Assay, obtained as directed in the individual monograph, calculate the content of active ingredient in each of the capsules, assuming homogeneous distribution of the active ingredient.

Soft Capsules — Determine the net weight of the contents of individual capsules as follows. Weigh accurately the 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. From the results of the Assay, obtained as directed in the individual monograph, calculate the content of active ingredient in each of the capsules, assuming homogeneous distribution of the active ingredient.

Solids (Including Sterile Solids) in Single-Unit Containers — Proceed as directed for Hard Capsules, treating each unit as described therein.

Solutions for Inhalation Packaged in Glass or Plastic Ampuls, Intended for Use in Nebulizers — Proceed as directed for Hard Capsules, treating each unit as described therein.

Oral Solutions and Syrups Packaged in Single-Unit Containers — Weigh accurately the amount of liquid that drains in not more than 5 seconds from each of 10 individual containers. If necessary, compute the equivalent volume after determining the apparent density. From the result of the Assay, obtained as directed in the individual monograph, calculate the content of active ingredient in the liquid drained from each of the 10 units.

CONTENT UNIFORMITY

For the determination of dosage-unit uniformity by assay of individual units, select not fewer than 30 units, and proceed as follows for the dosage form designated.

UNCOATED AND COATED TABLETS, HARD AND SOFT CAPSULES, SUPPOSITORIES, TRANSDERMAL SYSTEMS, ORAL SOLUTIONS IN SINGLE-UNIT CONTAINERS, SUSPENSIONS IN SINGLE-UNIT CONTAINERS, SYRUPS IN SINGLE-UNIT CONTAINERS, INHALATIONS PACKAGED IN PREMETERED DOSAGE UNITS, AND SOLIDS (INCLUDING STERILE SOLIDS) IN SINGLE-UNIT CONTAINERS — Assay 10 units individually as directed in the Assay in the individual monograph, unless otherwise specified in the Procedure for content uniformity. For oral solutions, suspensions, and syrups in single-unit containers, conduct the Assay on the amount of well-mixed material that drains from an individual container in not more than 5 seconds, and express the results as delivered dose. Where the amount of active ingredient in a single dosage unit differs from that required in the Assay, adjust the degree of dilution of the
solutions and/or the volume of aliquots so that the concentration of the active ingredients in the final solution is of the same order as that obtained in the Assay procedure; or, in the case of a titrimetric assay, use a titrant of a different concentration, if necessary, so that an adequate volume of titrant is required (see Titrimetry (541)); see also Procedures under Tests and Assays in the General Notices and Requirements. If any such modifications are made in the Assay procedure set forth in the individual monograph, make the appropriate corresponding changes in the calculation formula and titration factor.

Where a special Procedure for content uniformity is specified in the test for Uniformity of dosage units in the individual monograph, make any necessary correction of the results obtained as follows.

1. Prepare a composite specimen of a sufficient number of dosage units to provide the amount of specimen called for in the Assay in the individual monograph plus the amount required for the special Procedure for content uniformity in the monograph by finely powdering tablets or mixing the contents of capsules or oral solutions, syrups, suspensions, or solids in single-unit containers to obtain a homogeneous mixture. If a homogeneous mixture cannot be obtained in this manner, use suitable solvents or other procedures to prepare a solution containing all of the active ingredient, and use appropriate aliquot portions of this solution for the specified procedures.

2. Assay separate, accurately measured portions of the composite specimen of capsules or tablets or suspensions or inhalations or solids in single-unit containers, both (a) as directed in the Assay, and (b) using the special Procedure for content uniformity in the monograph.

3. Calculate the weight of active ingredient equivalent to 1 average dosage unit, by (a) using the results obtained by the Assay procedure, and by (b) using the results obtained by the special procedure.

4. Calculate the correction factor, $F$, by the formula:

$$F = \frac{A}{P}$$

in which $A$ is the weight of active ingredient equivalent to 1 average dosage unit obtained by the Assay procedure; and $P$ is the weight of active ingredient equivalent to 1 average dosage unit obtained by the special procedure. If

$$\frac{100|A - P|}{A}$$

is greater than 10, the use of a correction factor is not valid.

5. A valid correction may be applied only if $F$ is not less than 1.030 nor greater than 1.100, or, not less than 0.900 nor greater than 0.970, and if $F$ is between 0.970 and 1.030 no correction is required.

6. If $F$ lies between 1.030 and 1.100, or between 0.900 and 0.970, calculate the weight of active ingredient in each dosage unit by multiplying each of the weights found using the special procedure by $F$.

**Calculation of the Relative Standard Deviation**

The use of preprogrammed calculators or computers is acceptable. A manual mathematical method is as follows:
\[ s = \text{sample standard deviation}. \]

\[ \text{RSD} = \text{relative standard deviation (the sample standard deviation expressed as a percentage of the mean)}.\]

\[ \text{bar}(X-) = \text{mean of the values obtained from the units tested, expressed as a percentage of the label claim}. \]

\[ n = \text{number of units tested}. \]

\[ * x_1 * x_2 * \cdots * x_n = \text{individual values (} x_i \text{) of the units tested, expressed as a percentage of the label claim}. \]

\[ s = \left[ \frac{\sum (x_i - \bar{X})^2}{n - 1} \right]^{1/2} \]

\[ \text{RSD} = \frac{100s}{\bar{X}}. \]

**Criteria**

Apply the following criteria, unless otherwise specified in the individual monograph.

(A) If the Average of the Limits Specified in the Potency Definition in the Individual Monograph is 100.0 Percent or Less

**COMPRESSED TABLETS (COATED OR UNCOATED), SUPPOSITORYIES, ORAL SOLUTIONS IN SINGLE-UNIT CONTAINERS, SYRUPS IN SINGLE-UNIT CONTAINERS, SUSPENSIONS IN SINGLE-UNIT CONTAINERS, SOLIDS (INCLUDING STERILE SOLIDS) IN SINGLE-UNIT CONTAINERS, AND STERILE SOLIDS FOR PARENTERAL USE** — Unless otherwise specified in the individual monograph, the requirements for dosage uniformity are met if the amount of the active ingredient in each of the 10 dosage units as determined from the Weight Variation or the Content Uniformity method lies within the range of 85.0% to 115.0% of the label claim and the Relative standard deviation is less than or equal to 6.0%.

If 1 unit is outside the range of 85.0% to 115.0% of label claim and no unit is outside the range of 75.0% to 125.0% of label claim, or if the Relative standard deviation is greater than 6.0%, or if both conditions prevail, test 20 additional units. The requirements are met if not more than 1 unit of the 30 is
outside the range of 85.0% to 115.0% of label claim and no unit is outside the range of 75.0% to
125.0% of label claim and the \textit{Relative-standard-deviation} of the 30 dosage units does not exceed 7.8%.

CAPSULES, TRANSDERMAL SYSTEMS, INHALATIONS PACKAGED IN PREMETERED DOSAGE UNITS, AND
MOLDED TABLETS — Unless otherwise specified in the individual monograph, the requirements for
dosage uniformity are met if the amount of the active ingredient in not less than 9 of the 10 dosage
units as determined from the \textit{Weight Variation} or the \textit{Content Uniformity} method lies within the range of
85.0% to 115.0% of label claim and no unit is outside the range of 75.0% to 125.0% of label claim and the
\textit{Relative-standard-deviation} of the 10 dosage units is less than or equal to 6.0%.

If 2 or 3 dosage units are outside the range of 85.0% to 115.0% of label claim, but not outside the
range of 75.0% to 125.0% of label claim, or if the \textit{Relative-standard-deviation} is greater than 6.0% or if
both conditions prevail, test 20 additional units. The requirements are met if not more than 3 units of
the 30 are outside the range of 85.0% to 115.0% of label claim and no unit is outside the range of
75.0% to 125.0% of label claim, and the \textit{Relative-standard-deviation} of the 30 dosage units does not
exceed 7.8%.

(B) If the Average of the Limits Specified in the Potency Definition in the Individual Monograph is
Greater than 100.0 Percent —

1. If the average value of the dosage units tested is 100.0 percent or less, the requirements are
   as in (A).

2. If the average value of the dosage units tested is greater than or equal to the average of the
   limits specified in the potency definition in the individual monograph, the requirements are as in
   (A), except that the words "label claim" are replaced by the words "label claim multiplied by
   the average of the limits specified in the potency definition in the monograph divided by 100".

3. If the average value of the dosage units tested is between 100 percent and the average of the
   limits specified in the potency definition in the individual monograph, the requirements are as in
   (A), except that the words "label claim" are replaced by the words "label claim multiplied by
   the average value of the dosage units tested (expressed as a percent of label claim) divided by
   100".
Add the following:

(905) **UNIFORMITY OF DOSAGE UNITS**

[ 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 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 in the individual monograph.

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* is based on the assay of the individual content of drug substance(s) in a number of individual 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 *Content Uniformity* is required for those dosage forms described in (C1)–(C6) below:

| (C1) | coated tablets, other than film-coated tablets containing 25 mg or more of a drug substance that comprises 25% or more (by weight) of one tablet; |
| (C2) | transdermal systems; |
| (C3) | suspensions or emulsions or gels in unit-dose containers or in soft capsules that are intended for systemic administration only (not for those drug products that are intended for external, cutaneous administration); |
| (C4) | inhalations (other than solutions for inhalation packaged in glass or plastic ampul and intended for use in nebulizers) packaged in premetered dosage units. For inhalers and premetered dosage units labeled for use with a named inhalation device, also see *Aerosols, Metered-Dose Inhalers, and Dry Powder Inhalers* (601); |
| (C5) | solids (including sterile solids) that are packaged in single-unit containers and that contain active or inactive added substances, except that the test for *Weight Variation* may be applied in the special situations stated in (W2) and (W3) below; and |
| (C6) | suppositories. |

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

| (W1) | solutions for inhalation that are packaged in glass or plastic ampul and intended for use in nebulizers, and oral solutions packaged in unit-dose containers and into soft capsules; |
| (W2) | solids (including sterile solids) that are packaged in single-unit containers and contain no added substances, whether active or inactive; |
| (W3) | solids (including sterile solids) that are packaged in single-unit containers, with or without added substances, whether active or inactive, 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 *Content Uniformity* requirements. |

The test for *Content Uniformity* is required for all dosage forms not meeting the above conditions for the *Weight Variation* test. Alternatively, products listed in item (W4) above that do not meet the 25 mg/25% threshold limit may be tested for uniformity of dosage units by *Weight 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. 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.

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

<table>
<thead>
<tr>
<th>Dosage Form</th>
<th>Type</th>
<th>Subtype</th>
<th>Dose &amp; Ratio of Drug Substance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tablets</td>
<td>Uncoated</td>
<td>WV</td>
<td>≥25 mg &amp; ≥25%</td>
</tr>
<tr>
<td></td>
<td>Coated</td>
<td>WV</td>
<td>&lt;25 mg &amp; &lt;25%</td>
</tr>
<tr>
<td>Capsules</td>
<td>Hard</td>
<td>WV</td>
<td>CU</td>
</tr>
<tr>
<td></td>
<td>Soft</td>
<td>CU</td>
<td>CU</td>
</tr>
<tr>
<td>Solids in single-unit</td>
<td>Single</td>
<td>WV</td>
<td>CU</td>
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<td>containers</td>
<td>component</td>
<td>WV</td>
<td>CU</td>
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<tr>
<td></td>
<td>Multiple</td>
<td>WV</td>
<td>WV</td>
</tr>
<tr>
<td></td>
<td>components</td>
<td>WV</td>
<td>WV</td>
</tr>
<tr>
<td></td>
<td>Others</td>
<td>WV</td>
<td>WV</td>
</tr>
<tr>
<td>Suspension, emulsion, or gel</td>
<td></td>
<td>CV</td>
<td>CV</td>
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<tr>
<td>for systemic use only</td>
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<td></td>
<td></td>
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<tr>
<td>only packaged in single-unit</td>
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<td></td>
<td></td>
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<tr>
<td>containers</td>
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<tr>
<td>Solutions for inhalation</td>
<td></td>
<td>WV</td>
<td>WV</td>
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<tr>
<td>packaged in glass or plastic</td>
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<tr>
<td>ampuls and intended for use</td>
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<tr>
<td>in nebulizers, and oral</td>
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<tr>
<td>solutions packaged in unit-</td>
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<tr>
<td>dose containers and into soft</td>
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<tr>
<td>capsules</td>
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<tr>
<td>Inhalations (other than</td>
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<td>CV</td>
<td>CV</td>
</tr>
<tr>
<td>solutions for inhalation</td>
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<tr>
<td>packaged in glass or plastic</td>
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<tr>
<td>ampuls and intended for use</td>
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<tr>
<td>in nebulizers) packaged in</td>
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<tr>
<td>premetered dosage units</td>
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<tr>
<td>Transdermal systems</td>
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<td>CU</td>
<td>CU</td>
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<tr>
<td>Suppositories</td>
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<td>CU</td>
<td>CU</td>
</tr>
<tr>
<td>Others</td>
<td></td>
<td>CU</td>
<td>CU</td>
</tr>
</tbody>
</table>
**CONTENT UNIFORMITY**

Select not less than 30 units, and proceed as follows for the dosage form designated. Where the amount of drug substance in a single dosage unit differs from that required in the Assay, adjust the degree of dilution of the solutions and/or the volume of aliquots so that the concentration of the drug substances in the final solution is of the same order as that obtained in the Assay procedure; or, in the case of a titrimetric assay, use a titrant of a different concentration, if necessary, so that an adequate volume of titrant is required (see Titrimetry (541)); see also Procedures under Tests and Assays in the General Notices and Requirements. If any such modifications are made in the Assay procedure set forth in the individual monograph, make the appropriate corresponding changes in the calculation formula and titration factor.

Where a special Procedure for content uniformity is specified in the test for Uniformity of dosage units in the individual monograph, make any necessary correction of the results obtained as follows.

1. Prepare a composite specimen of a sufficient number of dosage units to provide the amount of specimen called for in the Assay in the individual monograph plus the amount required for the special Procedure for content uniformity in the monograph by finely powdering tablets or mixing the contents of capsules or oral solutions, suspensions, emulsions, gels, or solids in single-unit containers to obtain a homogeneous mixture. If a homogeneous mixture cannot be obtained in this manner, use suitable solvents or other procedures to prepare a solution containing all of the drug substance, and use appropriate aliquot portions of this solution for the specified procedures.

2. Assay separate, accurately measured portions of the composite specimen of capsules or tablets or suspensions or inhalations or solids in single-unit containers, both (a) as directed in the Assay, and (b) using the special Procedure for content uniformity in the monograph.

3. Calculate the weight of drug substance equivalent to 1 average dosage unit, by (a) using the results obtained by the Assay procedure, and by (b) using the results obtained by the special procedure.

4. Calculate the correction factor, $F$, by the formula:

$$ F = \frac{W}{P}, $$

in which $W$ is the weight of drug substance equivalent to 1 average dosage unit obtained by the Assay procedure, and $P$ is the weight of drug substance equivalent to 1 average dosage unit obtained by the special procedure. If

$$ \frac{100|W - P|}{W} $$

is greater than 10, the use of a correction factor is not valid.

5. The correction factor is to be applied only if $F$ is not less than 1.030 nor greater than 1.100, or not less than 0.900 nor greater than 0.970. If $F$ is between 0.970 and 1.030, no correction is required.

6. If $F$ lies between 1.030 and 1.100, or between 0.900 and 0.970, calculate the weight of drug substance in each dosage unit by multiplying each of the weights found using the special procedure by $F$.

**Uncoated, Coated, or Molded Tablets, Capsules, Oral Solutions in Single-Unit Containers, Oral Suspensions or Oral Emulsions or Oral Gels in Single-Unit Containers, and Solids (including Sterile Solids) in Single-Unit Containers**— Assay 10 units individually as directed in the Assay in the individual monograph.
monograph, unless otherwise specified in the *Procedure for content uniformity* in the individual monograph. Calculate the acceptance value as directed below.

For oral solutions, oral suspensions, oral emulsions, or oral gels in single-unit containers, conduct the *Assay* on the amount of well-mixed material that drains from an individual container in not more than 5 seconds, or for highly viscous products, conduct the *Assay* on the amount of well-mixed material that is obtained by quantitatively removing the contents from an individual container, and express the results as the delivered dose.

**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>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Conditions</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\bar{X})</td>
<td>Mean of individual contents ((\chi_1, \chi_2, \ldots, \chi_n)), expressed as a percentage of the label claim</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(\chi_1, \chi_2, \ldots, \chi_n)</td>
<td>Individual contents of the units tested, expressed as a percentage of the label claim</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(n)</td>
<td>Sample size (number of units in a sample)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(k)</td>
<td>Acceptability constant</td>
<td>If (n = 10), then (k = 2.4)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>If (n = 30), then (k = 2.0)</td>
<td></td>
</tr>
<tr>
<td>(s)</td>
<td>Sample standard deviation</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\[
\sqrt{n - 1} \left( \sum_{i=1}^{n} (\chi_i - \bar{X})^2 \right) \right]^{1/2}
\]
<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Conditions</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>RSD</td>
<td>Relative standard deviation (the sample standard deviation expressed as a percentage of the mean)</td>
<td></td>
<td><img src="http://www.usppf.com/pf/pub/index.html" alt="Image" /></td>
</tr>
<tr>
<td>M (case 1) to be applied when T (\leq) 101.5</td>
<td>Reference value</td>
<td>If 98.5% (\leq) (\overline{X}) (\leq) 101.5%, then</td>
<td>(M = \overline{X}) ((AV = ks))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If (\overline{X}) &lt; 98.5%, then</td>
<td>(M = 98.5%) ((AV = 98.5 - \overline{X} + ks))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If (\overline{X}) &gt; 101.5%, then</td>
<td>(M = 101.5%) ((AV = \overline{X} - 101.5 + ks))</td>
</tr>
<tr>
<td>M (case 2) to be applied when T &gt; 101.5</td>
<td>Reference value</td>
<td>If 98.5 (\leq) (\overline{X}) (\leq) (T), then</td>
<td>(M = \overline{X}) ((AV = ks))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If (\overline{X}) &lt; 98.5%, then</td>
<td>(M = 98.5%) ((AV = 98.5 - \overline{X} + ks))</td>
</tr>
<tr>
<td></td>
<td></td>
<td>If (\overline{X}) &gt; (T), then</td>
<td>(M = T%) ((AV = \overline{X} - T + ks))</td>
</tr>
<tr>
<td>Acceptance value (AV)</td>
<td></td>
<td></td>
<td>(M - \overline{X} + ks)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(Calculations are specified above for the different cases.)</td>
</tr>
<tr>
<td>L1</td>
<td>Maximum allowed acceptance value</td>
<td></td>
<td>(L1 = 15.0) unless otherwise specified in the individual monograph</td>
</tr>
<tr>
<td>L2</td>
<td>Maximum allowed range for deviation of each dosage unit tested from the calculated value of M</td>
<td>On the low side, no dosage unit result can be less than ((1 - L2<em>0.01)M), while on the high side no dosage unit result can be greater than ((1 + L2</em>0.01)M). (This is based on an L2 value of 25.0.)</td>
<td>(L2 = 25.0) unless otherwise specified in the individual monograph</td>
</tr>
</tbody>
</table>
### Table

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Conditions</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T$</td>
<td>Target test sample amount at time of manufacture. For purposes of this Pharmacopeia, unless otherwise specified in the individual monograph, $T$ is 100.0%, and for manufacturing purposes, $T$ is the manufacturer's approved target test amount value at the time of manufacture.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$M = \bar{X} \ (AV = ks)$

### Suppositories, Transdermal Systems, and Inhalations Packaged in Premetered Dosage Units—

NOTE— Acceptance value calculations are not required for these dosage forms. ] Assay 10 units individually as directed in the Assay in the individual monograph, unless otherwise specified in the Procedure for content uniformity.

### WEIGHT VARIATION

Select not less than 30 dosage units, and proceed as follows for the dosage form designated. The result of the Assay, obtained as directed in the individual monograph, is designated as result $A$, expressed as % of label claim (see Calculation of the Acceptance Value). Assume that the concentration (weight of drug substance per weight of dosage unit) is uniform. [ NOTE— Specimens other than these test units may be drawn from the same batch for assay determinations. ]

### Uncoated or Film-Coated Tablets—

Accurately weigh 10 tablets individually. Calculate the drug substance content, expressed as % 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, expressed as % of label claim, 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, expressed as % of label claim, in each capsule from the net weight of product removed from the individual capsules and the result of the Assay. Calculate the acceptance value.
Solids (Including Sterile Solids) in Single-Unit Containers— Proceed as directed for *Hard Capsules*, treating each unit as described therein. Calculate the acceptance value.

Oral Solutions Packaged in Single-Unit Containers— Accurately weigh the amount of liquid that drains in not more than 5 seconds from each of 10 individual containers. If necessary, compute the equivalent volume after determining the density. Calculate the drug substance content, expressed as % of label claim, in the liquid drained from each unit from the net weight of the individual container content 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.

\[
\chi_1, \chi_2, \ldots, \chi_n = \text{the individual estimated contents of the units tested, where}
\]

\[
\chi_i = \frac{w_i \times A}{W},
\]

\[
W = \text{individual weights of the units tested,}
\]

\[
A = \text{content of drug substance (% of label claim) determined as described in the Assay, and}
\]

\[
\bar{W} = \text{mean of individual weights (} w_1, w_2, \ldots, w_n).\]

Solutions for Inhalation Packaged in Glass or Plastic Ampuls and Intended for Use in Nebulizers— [NOTE— Acceptance value calculations are not required for these dosage forms. ] Accurately weigh 10 containers individually, taking care to preserve the identity of each container. Remove the contents of each container by a suitable means. Accurately weigh the emptied containers individually, and calculate for each container the net weight of its contents by subtracting the weight of the container from the respective gross weight. From the results of the Assay, obtained as directed in the individual monograph, calculate the drug substance content, expressed as % of label claim, in each of the containers.

CRITERIA

Apply the following criteria, unless otherwise specified in the individual monograph.

Uncoated, Coated, or Molded Tablets, Capsules, Oral Solutions in Single-Unit Containers, Oral Suspensions or Oral Emulsions or Oral Gels in Single-Unit Containers, and Solids (Including Sterile Solids) in Single-Unit Containers— 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 greater than \( L1 \)% 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 less than or equal to \( L1 \)% and no individual content of any dosage unit is less than \((1 - L2*0.01)M\) nor more than \((1 + L2*0.01)M\) as specified in the Calculation of Acceptance Value under *Content Uniformity* or under *Weight Variation*. Unless otherwise specified in the individual monograph, \( L1 \) is 15.0 and \( L2 \) is 25.0.

Suppositories—

*Limit A* (if the average of the limits specified in the potency definition in the individual monograph is 100.0 percent or less) — Unless otherwise specified in the individual monograph, the requirements for
Dosage uniformity are met if the amount of the drug substance in each of the 10 dosage units as determined from the Content Uniformity method lies within the range of 85.0% to 115.0% of the label claim, and the RSD is less than or equal to 6.0%.

If 1 unit is outside the range of 85.0% to 115.0% of label claim, and no unit is outside the range of 75.0% to 125.0% of label claim, or if the RSD is greater than 6.0%, or if both conditions prevail, test 20 additional units. The requirements are met if not more than 1 unit of the 30 is outside the range of 85.0% to 115.0% of label claim and the RSD of the 30 dosage units does not exceed 7.8%.

Limit B (if the average of the limits specified in the potency definition in the individual monograph is greater than 100.0 percent)—

1. If the average value of the dosage units tested is greater than or equal to the average of the limits specified in the potency definition in the individual monograph, the requirements are as specified under Limit A, except that the words “label claim” are replaced by the words “label claim multiplied by the average of the limits specified in the potency definition in the monograph divided by 100”.

2. If the average value of the dosage units tested is between 100 percent and the average of the limits specified in the potency definition in the individual monograph, the requirements are as specified under Limit A, except that the words “label claim” are replaced by the words “label claim multiplied by the average value of the dosage units tested (expressed as a percent of label claim) divided by 100”.

Transdermal Systems and Inhalations Packaged in Premetered Dosage Units—

Limit A (if the average of the limits specified in the potency definition in the individual monograph is 100.0 percent or less) — Unless otherwise specified in the individual monograph, the requirements for dosage uniformity are met if the amount of the drug substance in not less than 9 of the 10 dosage units as determined from the Content Uniformity method (or, in the case of solutions for inhalation packaged in glass or plastic ampuls and intended for use in nebulizers, from either the Content Uniformity or the Weight Variation method) lies within the range of 85.0% to 115.0% of label claim, and no unit is outside the range of 75.0% to 125.0% of label claim, and the RSD of the 10 dosage units is less than or equal to 6.0%.

If 2 or 3 dosage units are outside the range of 85.0% to 115.0% of label claim, but not outside the range of 75.0% to 125.0% of label claim, or if the RSD is greater than 6.0% or if both conditions prevail, test 20 additional units. The requirements are met if not more than 3 units of the 30 are outside the range of 85.0% to 115.0% of label claim and no unit is outside the range of 75.0% to 125.0% of label claim, and the RSD of the 30 dosage units does not exceed 7.8%.

Limit B (if the average of the limits specified in the potency definition in the individual monograph is greater than 100.0 percent)—

1. If the average value of the dosage units tested is greater than or equal to the average of the limits specified in the potency definition in the individual monograph, the requirements are as specified under Limit A, except that the words “label claim” are replaced by the words “label claim multiplied by the average of the limits specified in the potency definition in the monograph divided by 100”.

2. If the average value of the dosage units tested is between 100 percent and the average of the limits specified in the potency definition in the individual monograph, the requirements are as specified under Limit A, except that the words “label claim” are replaced by the words “label claim multiplied by the average value of the dosage units tested (expressed as a percent of label claim) divided by 100”.

(Official April 1, 2006)