

In Vitro Dissolution Testing of Nicotine Release from Smokeless Tobacco Products

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Altria
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Substantial Equivalence (SE) Pathway “905”

Compare Predicate vs. New Product

Demonstrates if the new product has:

- Same characteristics as the predicate
- Product does not raise different questions of public health

US FDA: Abbreviated HPHC List*

| Cigarette Smoke | Cigarette Filler |
|--------------------|---------------------------|
| NNK | NNK |
| NNN | NNN |
| Nicotine | Nicotine |
| Acetaldehyde | Arsenic |
| Crotonaldehyde | Cadmium |
| Formaldehyde | Ammonia |
| Acrolein | |
| Acrylonitrile | |
| Benzene | Smokeless Tobacco |
| 1,3-Butadiene | NNK |
| Isoprene | NNN |
| Toluene | Nicotine (total and free) |
| 4-Aminobiphenyl | Acetaldehyde |
| 1-Aminonaphthalene | Crotonaldehyde |
| 2-Aminonaphthalene | Formaldehyde |
| Ammonia | Arsenic |
| Benzo[a]pyrene | Cadmium |
| Carbon monoxide | Benzo[a]pyrene |

*Reporting Harmful and Potentially Harmful Constituents in Tobacco Products and Tobacco Smoke Under Section 904(a)(3) of the Federal Food, Drug, and Cosmetic Act” (Guidance for the Industry, March 2012).

Substantial Equivalence (SE) for Smokable Tobacco Products

Predicate vs. New Product

- Comparison of HPHCs in cigarette filler and in smoke (route of exposure)
- Smoking machines allow for a comparison of multiple products
- Standardized smoking protocols: ISO, CORESTA & HC
- Not meant to replicate human exposure



Cigarette



Smoking Machine

ISO: International Organization for Standardization CORESTA: Cooperation Center for Scientific Research Relative to Tobacco HC: Health Canada.

Substantial Equivalence (SE) for Smokeless Tobacco Products



Cigarette



Smoking Machine



Loose



Snus



Pouch



Dissolution Testing

In vitro laboratory test method designed to demonstrate how efficiently an active ingredient is extracted out of a solid oral dosage into solution

Applications in Pharmaceutical Industry

- Guide product design
- Quality control testing
- Product to product performance comparison
- Develop *in-vivo/in-vitro* correlation (IVIVC)



Dissolution Methodology Considerations

- Apparatus
- Dissolution Media
- Analytical Method Development & Validation



Dissolution Apparatus

Multiple commercial & non-commercial options available for measuring release of constituents from oral dosage forms

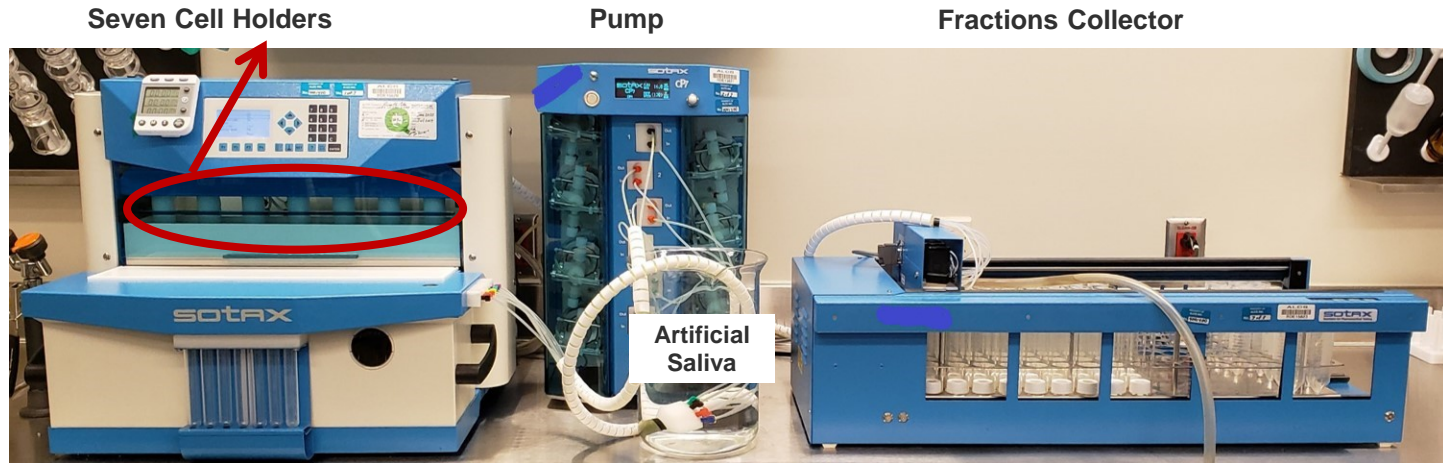
Example: U.S. Pharmacopeia Apparatus

| Name | Apparatus Type |
|-------|------------------------|
| USP-1 | Basket |
| USP-2 | Paddle |
| USP-3 | Reciprocating cylinder |
| USP-4 | Flow-through cell |
| USP-5 | Paddle over disk |
| USP-6 | Cylinder |
| USP-7 | Reciprocating holder |



ALCS Dissolution Method

SOTAX CE7 Smart Flow-Through Cell Apparatus (USP-4)

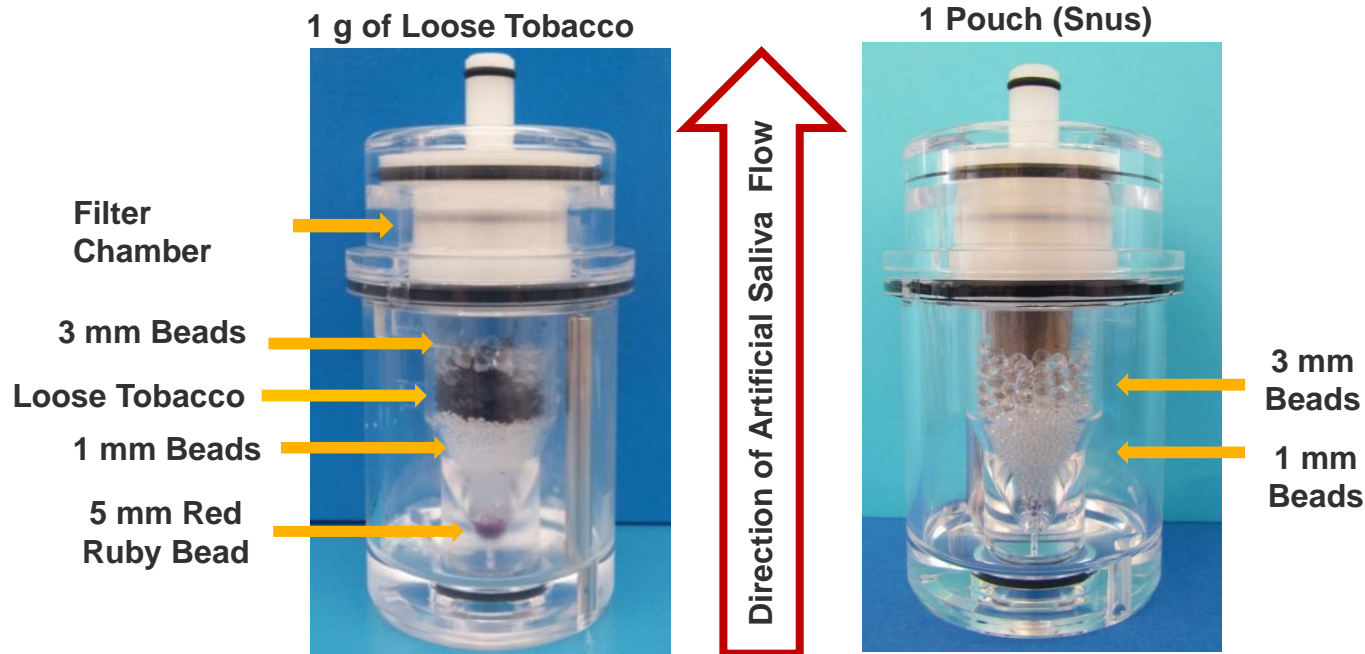


- Flow rate 4.0 mL/min
- Temperature 37°C



ALCS Dissolution Method

Flow-Through Cell in USP-4 Apparatus



ALCS Dissolution Method

Artificial Saliva Composition*

| Ingredient | Per 1000 mL |
|--|---------------------|
| Magnesium Chloride Hexahydrate ($\text{MgCl} \cdot 6\text{H}_2\text{O}$) | 0.17 g |
| Potassium Hydrogen Phosphate anhydrous ($\text{K}_2\text{HPO}_4 \cdot \text{H}_2\text{O}$) | 0.68 g |
| Sodium Chloride (NaCl) | 0.33 g |
| Potassium Chloride (KCl) | 0.75 g |
| Calcium chloride dihydrate ($\text{CaCl} \cdot 2\text{H}_2\text{O}$) | 0.15 g |
| Potassium Carbonate (K_2CO_3) | 0.53 g |
| Type I Water (De-ionized) | 1000 mL |
| Hydrochloric acid | To pH 6.8 ± 0.1 |

*German Institute for Standardization (DIN) Recipe is based upon German standard DIN v53160-1, "Determination of the Colour Release of Articles of Daily Use, Part1: Resistance to Artificial Saliva", section 4.2, October 2002.



ALCS Dissolution Method

Fraction Collection Protocol

| Fraction Number | Fraction Collection Time (min) | Fraction Collection Duration (min) | Volume Collected (mL) |
|-----------------|--------------------------------|------------------------------------|-----------------------|
| 1 | 4 | 4 | 16 |
| 2 | 8 | 4 | 16 |
| 3 | 12 | 4 | 16 |
| 4 | 16 | 4 | 16 |
| 5 | 20 | 4 | 16 |
| 6 | 30 | 10 | 40 |
| 7 | 40 | 10 | 40 |
| 8 | 50 | 10 | 40 |
| 9 | 60 | 10 | 40 |

- Flow rate 4.0 mL/min
- Temperature 37°C



Analytical Method: Determination of Nicotine in Dissolution Fractions

Sample Preparation in Autosampler Vials

1. 0.1 mL of the dissolution fraction
2. 0.8 mL of artificial saliva
3. 0.1 mL of ISTD (Ethyl Benzoate, 1 mg/mL)
4. Cap, vortex and analyze by UPLC-PDA*

ISTD: Internal Standard

UPLC-PDA: Ultra-high Performance Liquid Chromatography Photodiode Array

Analytical Method: Determination of Nicotine in Dissolution Fractions

Instrument: Waters Acquity I-Class UPLC system coupled to Photodiode Array Detector

Column: BEH C18, 2.1 x 100 mm, 1.7 μm , Waters Corporation

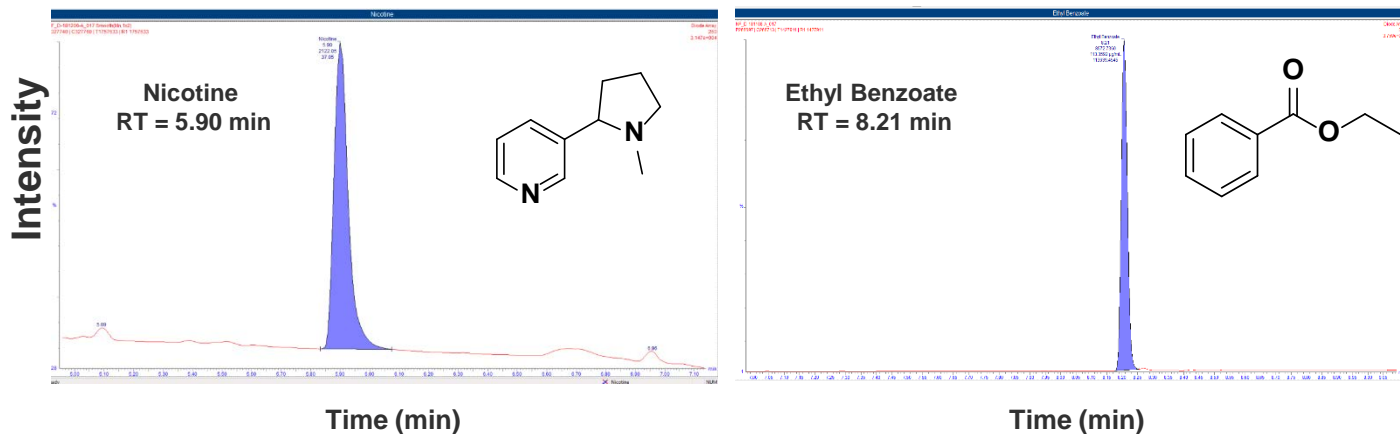
Guard column: BEH C18 VanGuard, 2.1 x 5 mm, 1.7 μm , Waters Corporation

| UPLC Parameter | Setting |
|-------------------------|------------------------------|
| Run Time | 12 min |
| Injection Volume | 10 μL |
| Autosampler Temperature | 5 $^{\circ}\text{C}$ |
| Column Temperature | 45 $^{\circ}\text{C}$ |
| Mobile Phase A | 10 mM Ammonium Acetate pH 10 |
| Mobile Phase B | Acetonitrile |
| Pump Program | Gradient Elution |
| Flow Rate | 0.5 mL /min |
| Compound | PDA Setting |
| Nicotine | 260 nm |
| Ethyl benzoate | 260 nm |



Analytical Method: Determination of Nicotine in Dissolution Fractions

UPLC-PDA Chromatograms of Nicotine and Ethyl Benzoate (ISTD)



ISTD: Internal Standard

UPLC-PDA: Ultra-high Performance Liquid Chromatography Photodiode Array

Analytical Method: Determination of Nicotine in Dissolution Fractions

Method Validation

| Parameter | Outcome |
|---|---|
| Calibration (0.5-100 µg/mL) | <ul style="list-style-type: none">▪ $R^2 > 0.998$ on all days▪ %RCR < 10% |
| Accuracy - 3 fortification levels in triplicate for each product type | 96.2% - 102% |
| Repeatability (Intra-day precision, n = 6) | < 2.0% |
| Intermediate Precision (Inter-days precision, n = 18) | < 6.0% |
| Specificity | No interferences observed at the retention time of nicotine or IS |
| LOQ | 0.5 µg/mL |
| Stability - Dissolution samples and final extracts | Stable for up to 15 days when stored in amber glass vials at 0-4°C (refrigerated) |

Analysis of Products for Regulatory Submissions

Predicate vs. New Product

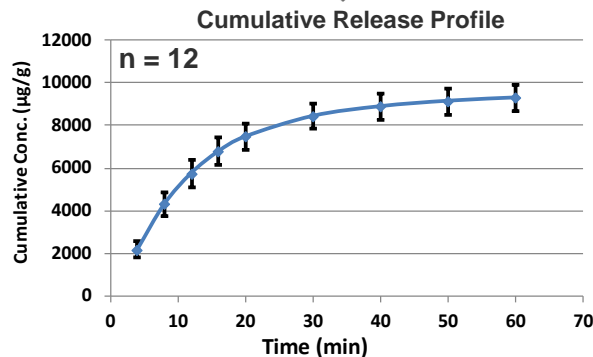
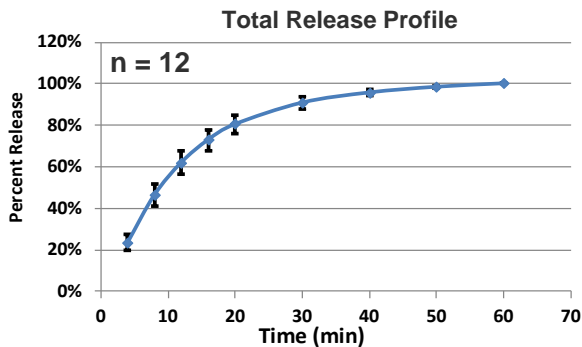
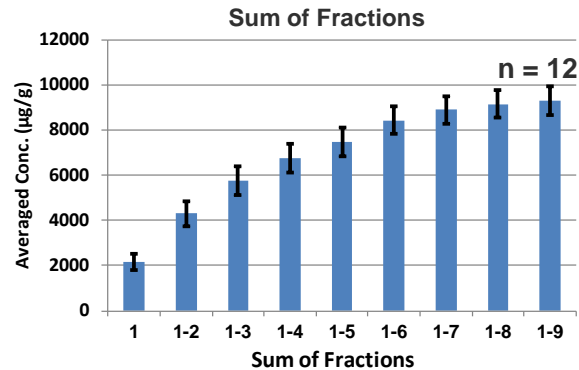
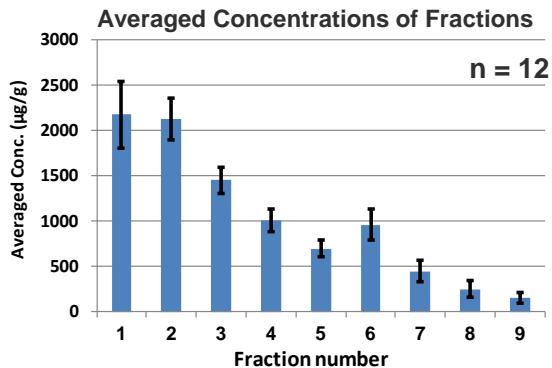
12 replicates per product*

- Requires two runs for each product (7 cell holders per unit)
 1. 6 Reps + 1 IQCS (CRP 2.1, American style loose moist snuff)
 2. 6 Reps + 1 reagent blank
- 9 fractions collected for each sample replicate

**Guidance for Industry: Dissolution Testing of Immediate Release Solid Oral Dosage Forms, FDA Center for Drug Evaluation and Research (CDER), 1997.*

- **IQCS:** Internal Quality Control Sample

Calculations for Cumulative Release Profile



Error Bars ± 1 S.D.



CORESTA Smokeless Tobacco Reference Products (CRPs)

CRP 1.1 (Swedish style snus pouch)



CRP 2.1 (American style loose moist snuff)



CRP 4 (loose-leaf chewing tobacco)



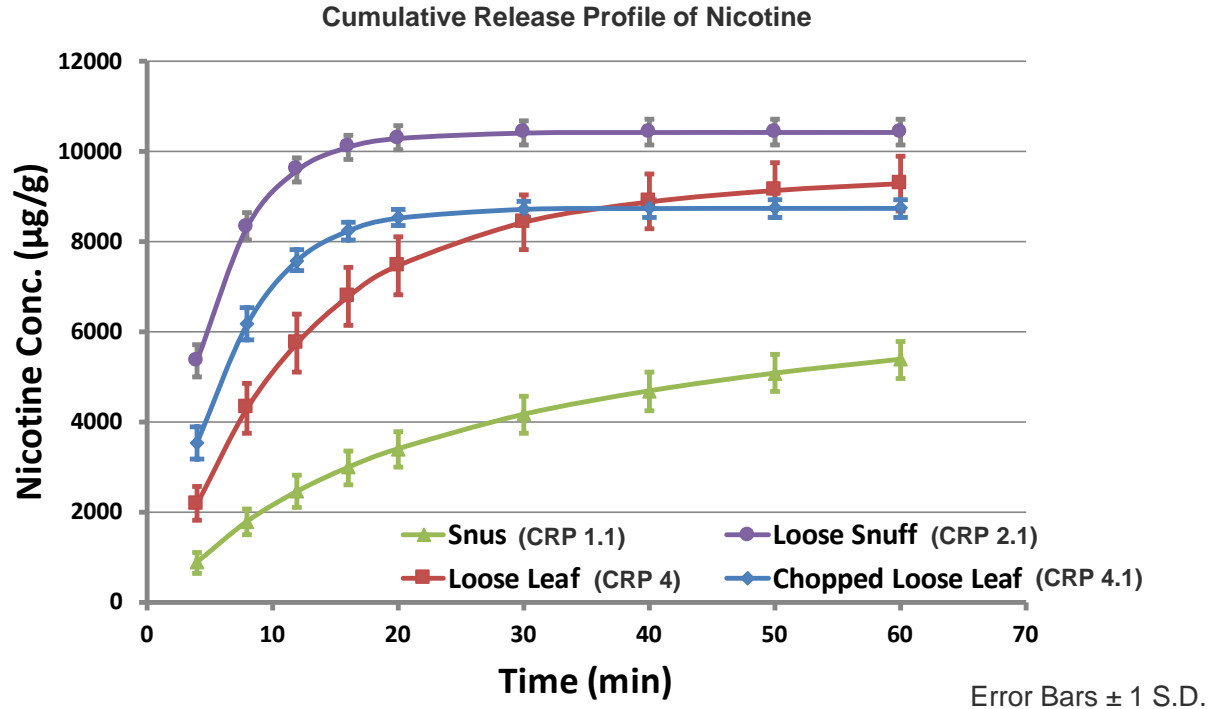
CRP 4.1 (chopped loose-leaf chewing tobacco)



<https://www.coresta.org/coresta-smokeless-tobacco-reference-products>

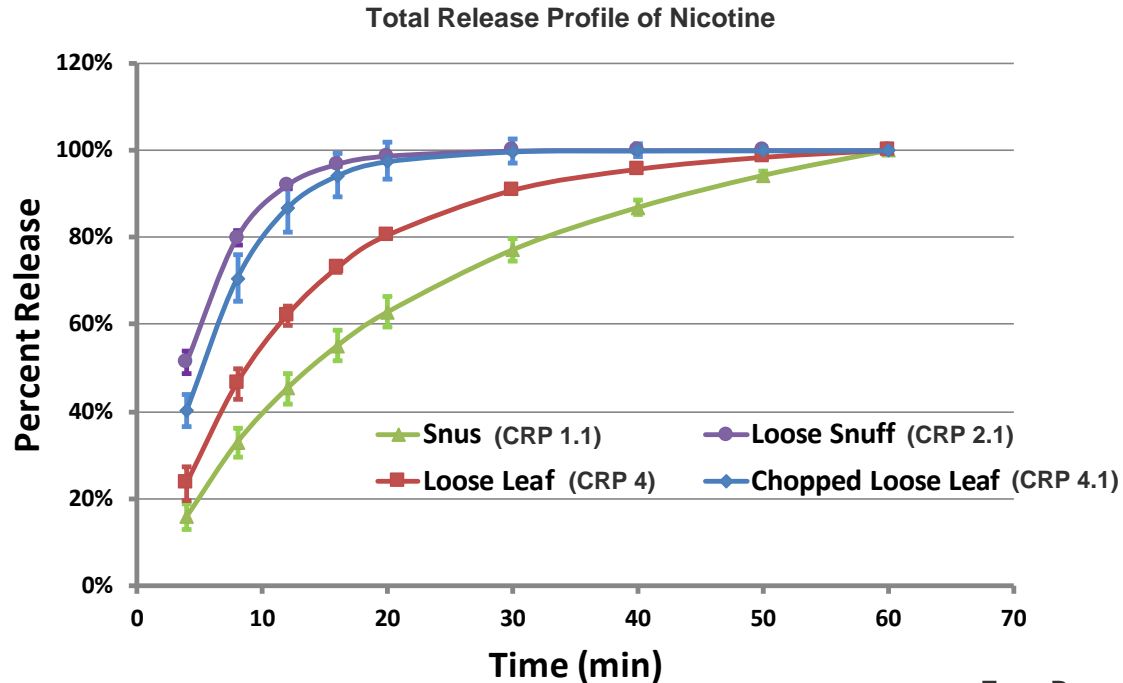
Results

CORESTA Smokeless Tobacco Reference Products (CRPs)



Results

CORESTA Smokeless Tobacco Reference Products (CRPs)



Error Bars \pm 1 S.D.



Dissolution Profiles Comparison

Approach based on FDA guidance from CDER*

- Determine nicotine release profiles for predicate and new product (12 replicates per product)
- Use mean dissolution values from both curves at each time point to calculate difference factor (f_1) and similarity factor (f_2)

f_1 values up to 15 (0-15) and f_2 values of 50 or greater (50-100)
ensure similarity or equivalence between two products

**Guidance for Industry: Dissolution Testing of Immediate Release Solid Oral Dosage Forms, FDA Center for Drug Evaluation and Research (CDER), 1997.*



Results

Product to Product Comparison using f_1 and f_2

| Compared Products | f_1 | f_2 | Equivalency |
|--------------------|-------|-------|-------------|
| CRP 2.1 vs CRP 4.1 | 11.3 | 52.7 | Yes |
| CRP 2.1 vs CRP 4 | 27.5 | 30.0 | No |
| CRP 2.1 vs CRP 1.1 | 39.2 | 21.9 | No |
| CRP 4.1 vs CRP 4 | 23.1 | 35.3 | No |
| CRP 4.1 vs CRP 1.1 | 36.1 | 24.8 | No |
| CRP 4 vs CRP 1.1 | 20.2 | 42.2 | No |

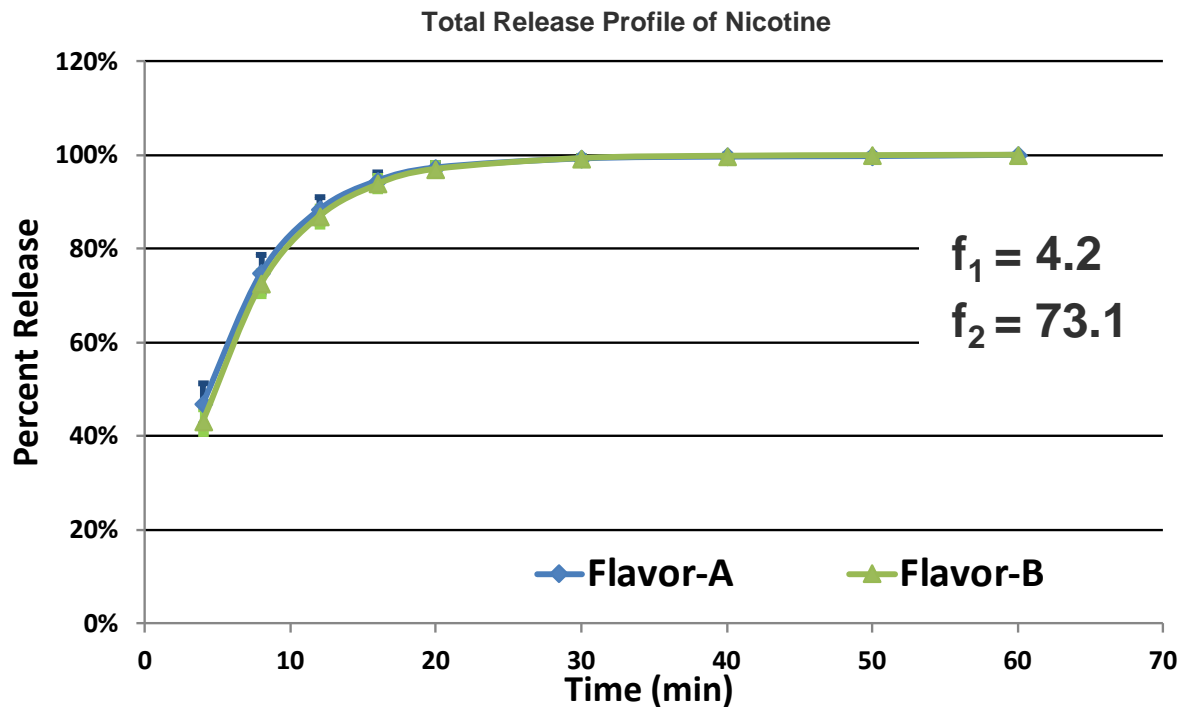
CRP 1.1 (Swedish style snus pouch)
CRP 2.1 (American style loose moist snuff)
CRP 4 (loose-leaf chewing tobacco)
CRP 4.1 (chopped loose-leaf chewing tobacco).

Equivalent:
f1 values up to 15 (0-15)
f2 values of 50 or greater (50-100)



Results

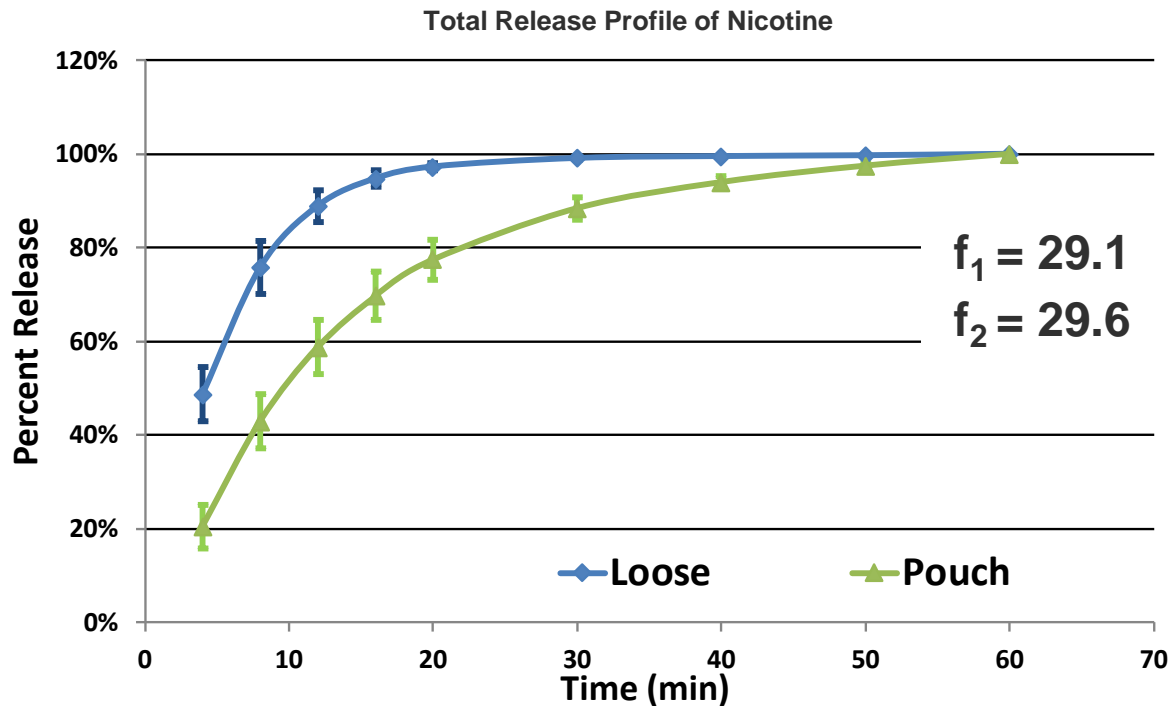
Commercial Products with Different Flavors



Error Bars ± 1 S.D.

Results

Loose vs Pouched Commercial Products



Error Bars \pm 1 S.D.

Summary

- An efficient method for the dissolution testing and quantitation of nicotine was validated for a variety of smokeless tobacco products using a USP-4 flow-through cell apparatus and UPLC-PDA
- Percent of total release profiles, and f_1 and f_2 calculations were used to evaluate the similarity and differences between the smokeless tobacco products
- The nicotine release profiles are dependent on the form and cut of the smokeless tobacco products
- Validated dissolution methodologies can be an important tool for smokeless tobacco product assessments and product-to-product comparisons



This presentation may be accessed @
www.altria.com/ALCS-Science



Results

f_1 and f_2 Calculations

$$f_1 = \left\{ \left[\sum_{t=1}^n |R - T| \right] / \left[\sum_{t=1}^n R \right] \right\} \times 100$$

$$f_2 = 50 \cdot \log \left[\frac{100}{\sqrt{1 + \frac{\sum_{t=1}^n [R_t - T_t]^2}{n}}} \right]$$

R_t and T_t are the cumulative percentage dissolved at each of the selected n time points of the two products.

f_1 values up to 15 (0-15) and f_2 values of 50 or greater (50-100) ensure similarity or equivalence between two products.