Certified Solution Standards and Reagents for Therapeutic Drug Monitoring Applications

CERTIFIED SOLUTION STANDARDS & REAGENTS FOR THERAPEUTIC DRUG MONITORING APPLICATIONS

Highly pure, well-characterized, solution based standards or reagents are a good and efficient alternative to the use of neat materials in clinical, toxicology and therapeutic drug monitoring

applications. The accuracy, stability, and consistency of these

- materials is critical to ensure accuracy of results in the analytical laboratory, in clinical applications and medical device performance.
- Certified Solution Standards and Reagents offer a significant advantage over neat reference materials in terms of accuracy, consistency and stability. Long term stability of solution based materials is achievable when appropriate parameters are chosen in the design, preparation, packaging, and storage.
- This poster presents the development and design of Certified Solution Standards and Reagents providing examples of premade solutions that exhibit multi-year stability. Factors critical to the selection of the analytes, diluents, storage and stability are discussed. These include: raw material handling, characterization and potency, certification and qualification of solutions, and homogeneity and stability of the solution. Certified Reference Standard Solutions and Reagents prepared in a diluent that promotes stability and packaged under argon in flame sealed ampoules can be stable for many years.
- Multiple examples are presented. Also presented is a comparison of certified ampouled solutions vs. neat reference materials.
- Certified Solution Standards and Reagents ensure accurate, consistent and reliable results in clinical and toxicology applications and medical device performance.

REFERENCE STANDARDS ARE CRITICAL TO THE QUANTITATION OF THERAPEUTIC DRUGS IN A CLINICAL SETTING

Results are only as accurate as the reference! Accurate quantitative results depend on

 The purity of the reference material The accuracy in the preparation of the solution used

in the analysis

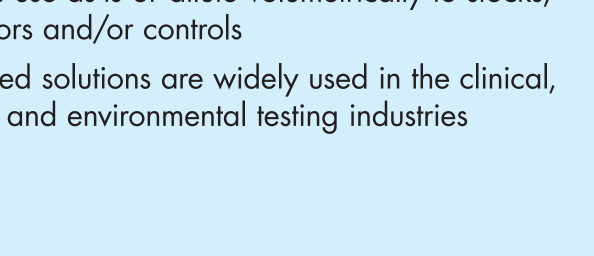
TYPES OF REFERENCE STANDARDS AND HOW THEY ARE USED

Certified neat reference standard Analysts prepare volumetric solutions by weighing the neat materials and diluting just prior to use • This may be a stock solution, calibration curve or control

Ampouled Certified Solutions

 Analysts use as-is or dilute volumetrically to stocks, calibrators and/or controls

 Ampouled solutions are widely used in the clinical, forensic and environmental testing industries



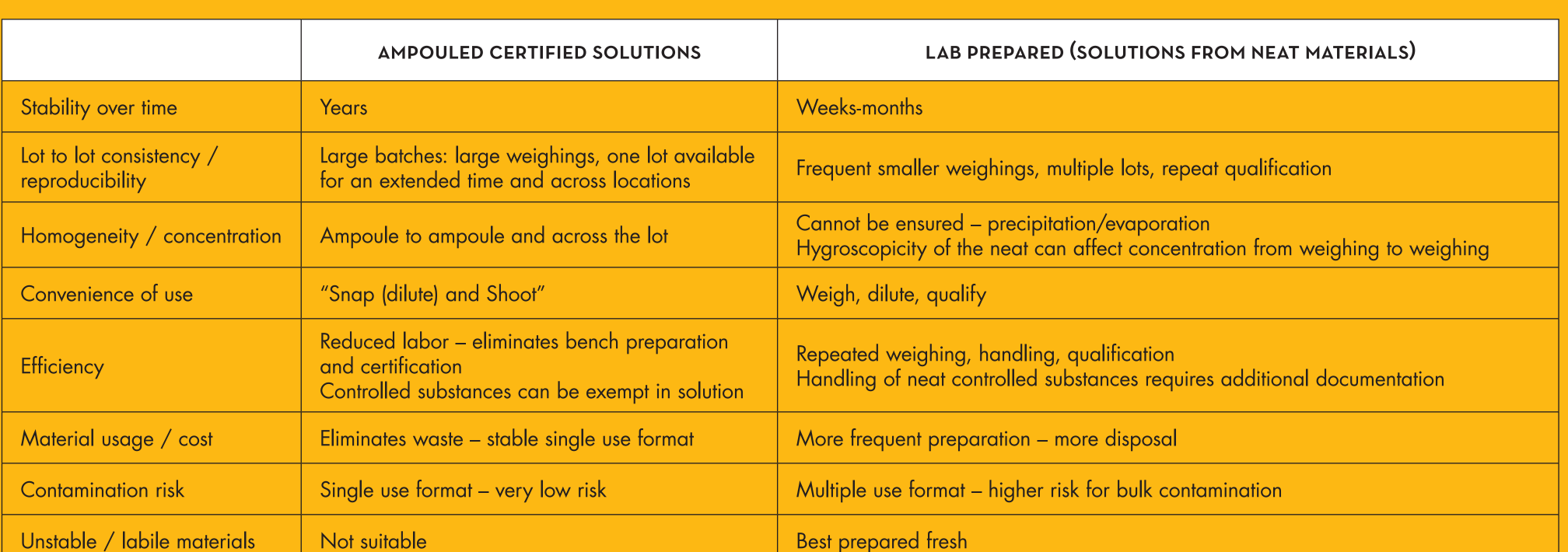
CHARACTERISTICS OF GOOD **REFERENCE STANDARDS**

Thoroughly characterized components

- Use of accurate, calibrated pipettes & balances Analyzed to verify accuracy & consistency
- Traceability of all components
- High purity diluents and/or stabilizers, compatible with the compound(s)

quality Ampouled Certified Solution Standards = Good Reference Standards

AMPOULED CERTIFIED SOLUTIONS – A BETTER ALTERNATIVE IN MOST CASES



PREPARATION OF CERTIFIED SOLUTIONS

NEAT MATERIAL

Complete characterization of the neat material is critical to the accuracy of the solution

Certification should include: Purity and impurities

- Residual water, solvent, inorganic content (KF, headspace, micro-ash/ROI)
- Chromatographic purity, resolution of impurities (LC, GC) - Quantitation by assay (LC, GC, titration)

Considerations

- Are vendor certified values complete, accurate and reliable?
- Reliability/repeatability of method?

Verification of identity (NMR, FTIR, MS)

- Is the compound more stable as salt vs. free base?
- Is the compound stable in the diluent?
- Is there an adjustment for salt form?
- Does the vendor provide uncertainty on the purity factor (potency)?

PREPARATION OF CERTIFIED SOLUTIONS

PURITY FACTOR (POTENCY) - SOURCES OF

- Chromatographic purity homogeneity of the neat and analytical
- Residual solvent sample preparation, weighing, and analytical
- analyst technique, and instrument tolerances
- Purity Factor = [(100 wt% residual solvent -
- Chromatographic Purity/100]

PREPARATION OF CERTIFIED SOLUTIONS

DILUENT/SOLVENT CONSIDERATIONS

Solubility

- Precipitation can occur over time or at reduced storage temperatures

- Solvent interferences in the chromatogram: UV cut-off; baseline
- Non-polar solvents not ideal with reverse phase HPLC

Solvent stability

THF/ethers form peroxides

Compound stability in the solvent

- stable in acetonitrile long term
- Oxazepam decomposes over time in methanol

PREPARATION OF CERTIFIED SOLUTIONS

SOLUTION PREPARATION

be handled in a glove box

Neat Material Handling – Air or moisture sensitive compounds handled in inert atmosphere - Use appropriate PPE - toxic and highly labile compounds must

Neat Material Weighing

- Use a qualified balance appropriate for the amount to
- Calibrated with NIST traceable weights
- 5-place, 6-place, 7-place?
- Qualified to <0.1% relative error per USP
- Use weighing techniques that minimize error Tongs vs. gloved hands
- Balance equilibration
- Material acclimated to conditions
- movement, temperature)
- Contamination weighing operations should be isolated from other operations

PREPARATION OF CERTIFIED SOLUTIONS

• Balance selection: Balances should be qualified and assigned

• Weighing techniques: Handling of samples during weighing can

have significant impact on weighing accuracy and uncertainty

Certified Solutions prepared in bulk with larger weighing have

reduced weighing error compared to smaller weighings typical

6-place 5-place 4-place

1 mg 3 mg 20 mg 125 mg

minimum weighings specific to installed conditions and lab

BALANCE SELECTION AND WEIGHING

when neat materials are weighed at the bench

0.00075%

USP specified minimum relative error of NMT 0.1%

methods outlined in the "ISO Guide to the Expression of Uncertainty in Measurement

**Minimum weights calculated by Mettler during balance qualification to achieve

and includes contributions from standard deviation, sensitivity, and linearity.

TECHNIQUE ARE CRITICAL TO

WEIGHING ACCURACY

specified weighing techniques

Larger weighings are more accurate

UNCERTAINTY

- Residual water sample weighing, influence of ambient moisture,
- Inorganic content sample weighing and homogeneity of the sample
- wt% residual water wt% residual inorganics) x
- Assay value reference standard uncertainty, sample preparation, and analytical uncertainty

- Does the target compound dissolve at the required concentration?

Compatibility with analysis

- Water not compatible with GC
- Protic solvents cocaine degrades in methanol over time but is

Purity, identity and traceability

PREPARATION OF CERTIFIED SOLUTIONS

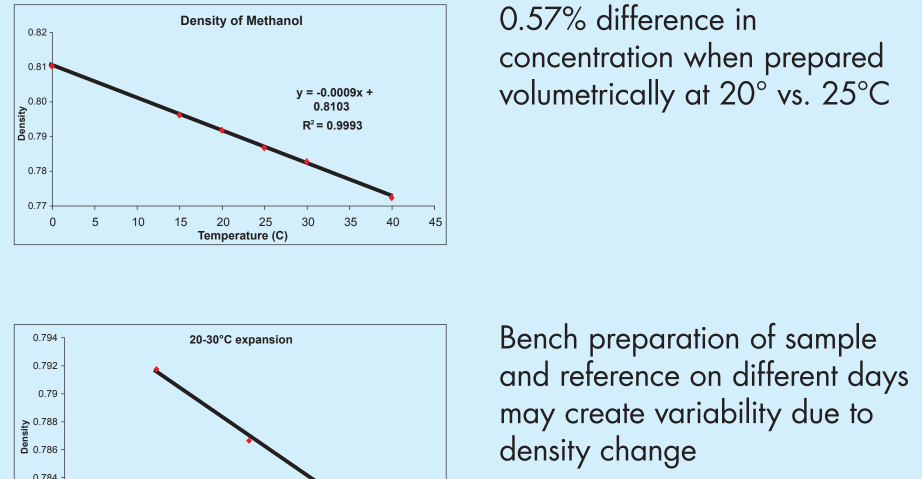
SOLVENT ADDITION

Gravimetric is more accurate than volumetric addition

METHOD	BATCH SIZE		
METHOD	10 mL	100 mL	1000 mL
Volumetric flask standard error Source: ASTM E288-03, Standard specification for laboratory glassware, 2003	0.20%	0.08%	0.03%
Analytical balance uncertainty Balance Type Typical values per Mettler Toledo	5 Place 0.001%	5 Place 0.0001%	1 Place 0.009%
Values established by Cerilliant based on typical values by Mettler and Cerilliant weighing SOPs	0.0036%	0.00125%	0.009%

TEMPERATURE VS. DENSITY

Change in density with temperature can affect volumetric preparation of a solution but can be controlled by gravimetric addition of solvent



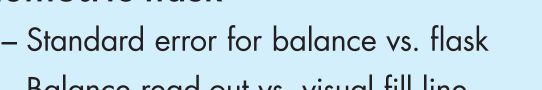
Source: Handbook of Thermophysical and Thermochemical Data, CRC Press

Add solvent by weight

15 20 25 30 35 temperature (°C)

- Target solvent weight calculated from target
- volume by adjusting for density Actual solvent weight can be calculated back into volume to report concentration in mg/mL

Weighing has lower error than volumetric flask



- Balance read-out vs. visual fill line Weighing tapes provide traceability

Control of lot to lot consistency Control of temperature/density variable

- Subjectivity of volumetric preparation technique

Use of high quality, qualified, balance has

lower error than Class-A volumetric flask

PREPARATION OF CERTIFIED SOLUTIONS

ISPENSING & PACKAGING

 Solution standards prepared in bulk can be dispensed into single use volumes and flame sealed under inert atmosphere

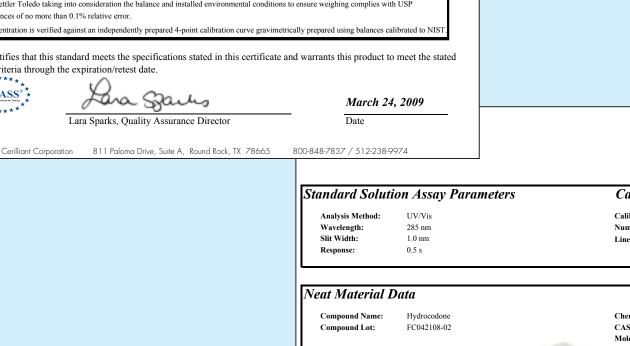
- Rigorous process controls ensure Consistency of volume dispensed - Homogeneity from vial to vial and across the lot
- No contamination No degradation

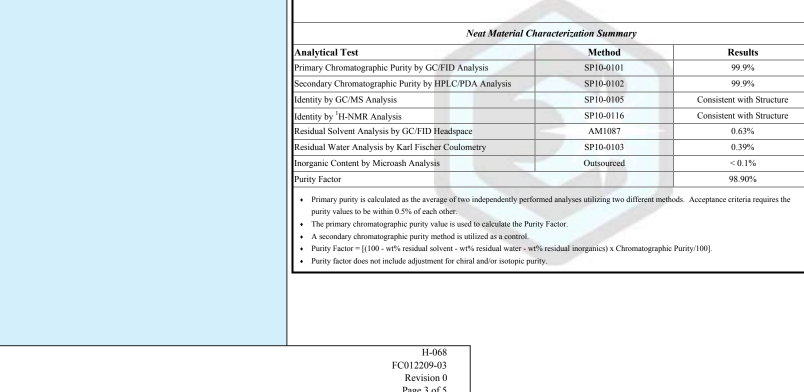


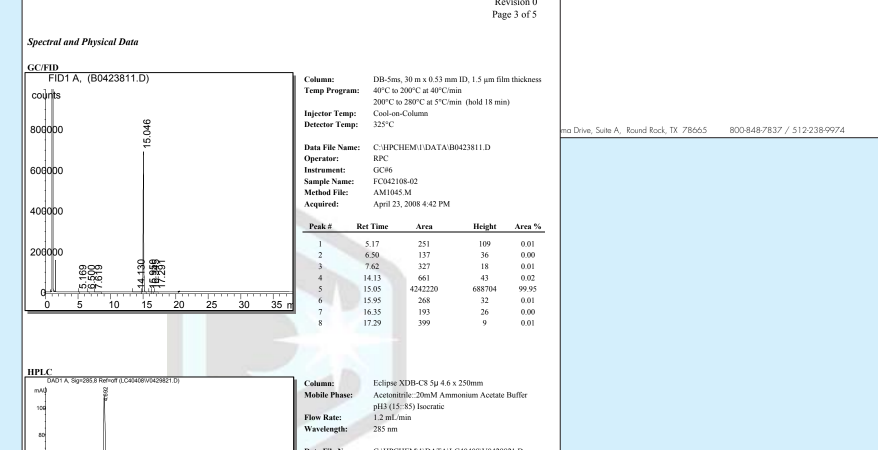


CERTIFICATION OF THE









Method File: RMA013.M Acquired: April 29, 2008 4:54 PM

Peak# Ret Time Area Height Area %

 3.43
 1.01
 0.10
 0.00

 3.88
 0.26
 0.05
 0.01

 4.69
 1751.35
 109.44
 99.9

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0 2.5 5 7.5 10 12.5 15 17.5

- Comparison to a primary source or certified second source curve/calibration standard
- Comparison of multiple independent preparations Consistency
- Lot-to-lot consistency verified by comparing to the previous lot
- Homogeneity
 Across the batch of ampoules/vials
- Test for contamination and degradation Traceability
- For all raw materials and calibrations Established through comparison to older lots

Ampouled solutions provide long-term stability

- Comparison to a primary source or certified second source Sealed amber ampoules protect from light and air
- Inert gas purge displaces oxygen

Silanization can reduce absorption to glass

- Sealed ampoules prevent concentration changes due to evaporation, absorption of moisture, and degradation Solution stability is assessed to determine shelf life
- Established by assay comparison to fresh solution Retest dates assigned to evaluate stability at set intervals Expiration dates established over time
- Sealed ampouled solutions can exhibit excellent long term stability (>5 years for some)

SOLUTION STABILITY EXAMPLES

6-ACETYLMORPHIN

NORTRIPTYLINE HCI

SOLUTION LOT NUMBE

Number of Calibration Points: 4

Solution Purity Analysis Method: HPLC

FE072108-01

FN071607-01

Number of Points: .

Previous Lot

Linearity (r):

Catalog Product:

Temp program:..

Injector Temp:

• No loss of chromatographic purity from originally established neat material value (within 0.5%)

COMPOUND/SOLVENT	AGE OF STABILITY SAMPLE	% DIFFERENCE IN CONCENTRATION BETWEEN STABILITY LOT AND NEW LOT
6-Acetylmorphine /acetonitrile	5.5 years	-2.3%
Nortriptyline HCl / methanol	5 years	-2.9%
Codeine / methanol	5.5 years	-0.7%
Haloperidol / methanol	6 years	-0.4%
Fentanyl /methanol	5 years	-1.3%

• Concentration meets original acceptance criteria – reported differences may be reflective of method variability

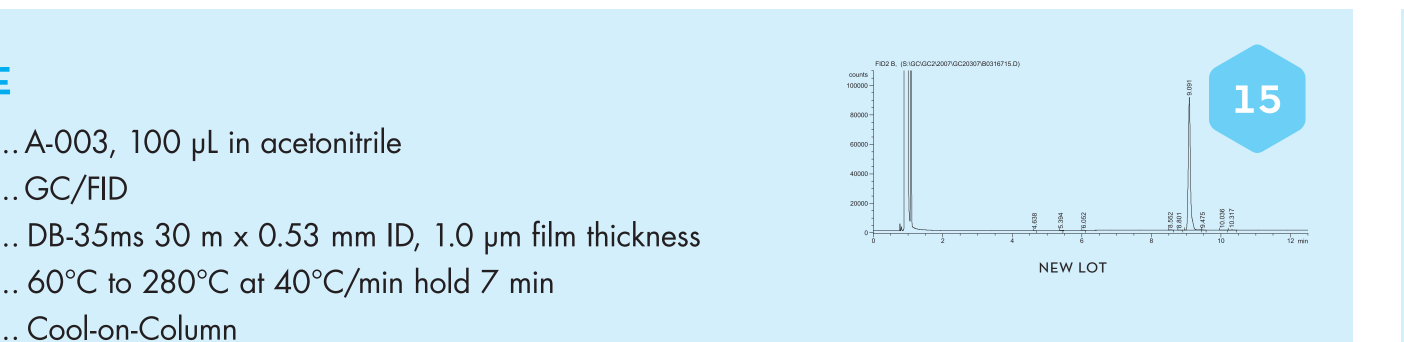
Mobile Phase:

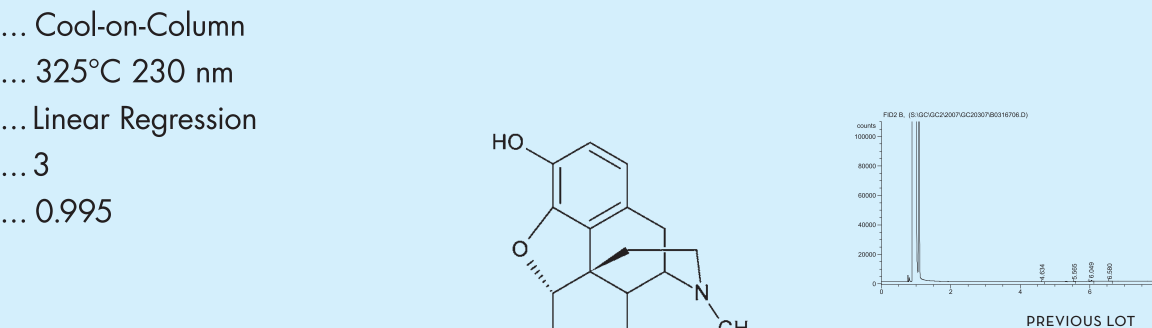
Flow Rate:

Wavelength

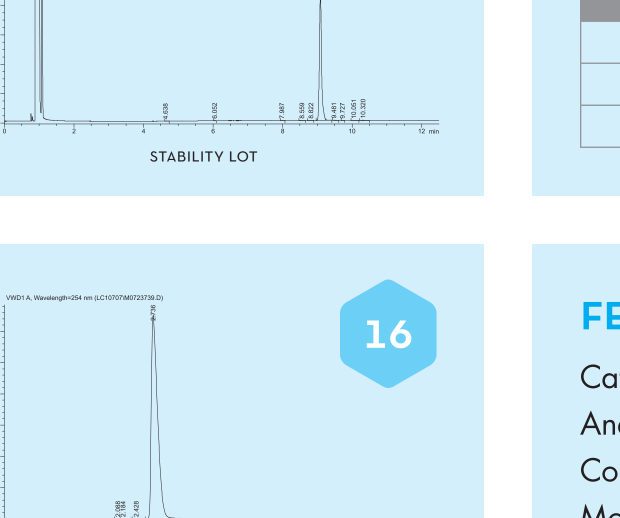
Linearity (r):

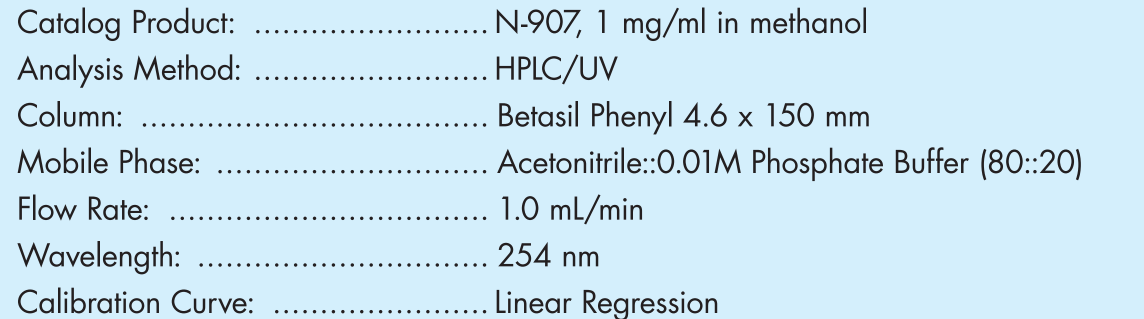
Number of Points:





			0		
					FID2 B, (S:\GC\GC2\2007\GC20307\B0316724.D) counts]
SOLUTION	LOT NUMBER	MANUFACTURE DATE	% CONC. DIFF FROM NEW PREP	SOLUTION PURITY	100000 - 800000
New Lot	FC022707-01A	3 / 2007	-	99.4%	40000
Previous Lot	FC040405-02B	9 / 2005	2.0	99.2%	20000 –
Stability Lot	34265-11B	7 / 2001	-2.3	99.5%	250
					STABILITY LOT





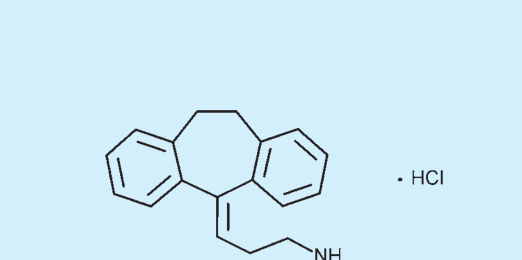
.. C-006, 1 mg/ml in methanol

. Betasil Phenyl 4.6 x 150 mm

8 / 2006

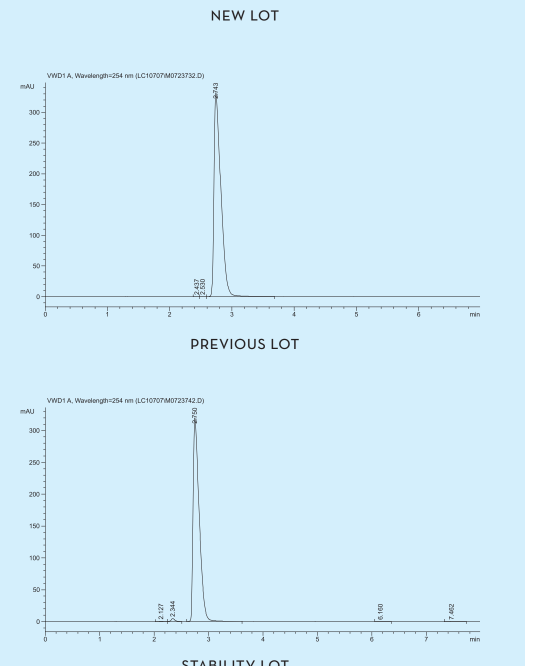
Acetonitrile::0.01M Phosphate Buffer (70::30)

.....Linear Regression

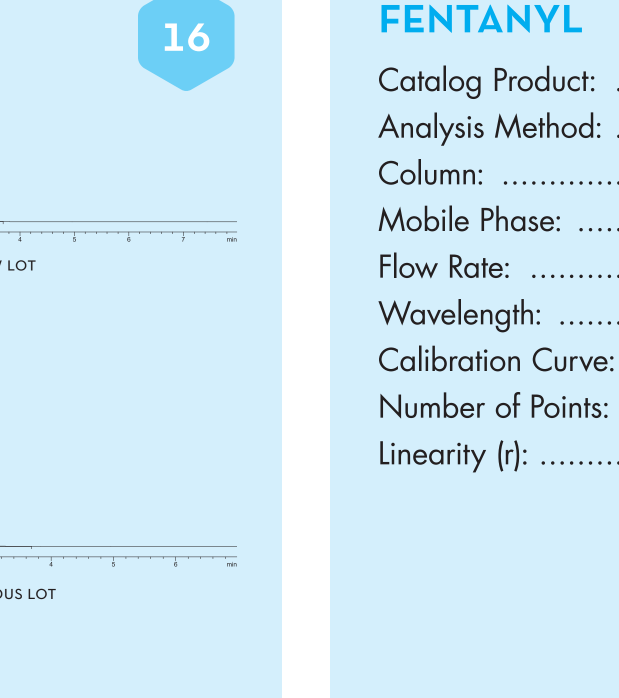


		CH ₃	mAU 300 - 250 -
MANUFACTURE DATE	% CONC. DIFF FROM NEW PREP	SOLUTION PURITY	200 -
7 / 2007	-	99.9%	100 -
7 / 2006	-0.7	99.9%	50-

-2.9 99.9%



STABILITY LOT



NEW LOT NEW LOT PREVIOUS LOT	2.088				
PREVIOUS LOT	1 2	3 4	5 6	7 min	
PREVIOUS LOT ***********************************		NEW LOT			
PREVIOUS LOT ***********************************					
PREVIOUS LOT ***********************************	enath=254 nm (LC10707\M07237;	32 D)			
PREVIOUS LOT ingth=254 nm (LC10707IM0723742.D) 2	ngin=294 nm (LU-10/0/1W0/237.	8-743			
ongth=254 nm (LC10707IM0723742.D)	1 2		5	6 min	
angth=254 nm (LC10707M0723742.D)		PREVIOUS LOT			
2.344 2.344 6.160					
2.127 2.344 6.160	ength=254 nm (LC10707\M072374	(42.D)			
1 2 3 4 5 6 7 min		£750	6.160	7.462	
		3 4		7 min	

SOLUTION LOT NUMBER MANUFACTURE DATE SOLUTION PURITY 99.8% 11 / 2000 STABILITY LOT .. F-002, 100 µg/mL in methanol . Betasil Phenyl 4.6 x 150 mm . Acetonitrile::0.01M Phosphate Buffer (70::30) . 1.0 mL/min Wavelength .. Linear Regression

.. H-030, 1 mg/ml in methanol

. Betasil Phenyl 4.6 x 150 mm

. 1.0 mL/min

.. Linear Regression

.. 245 nm

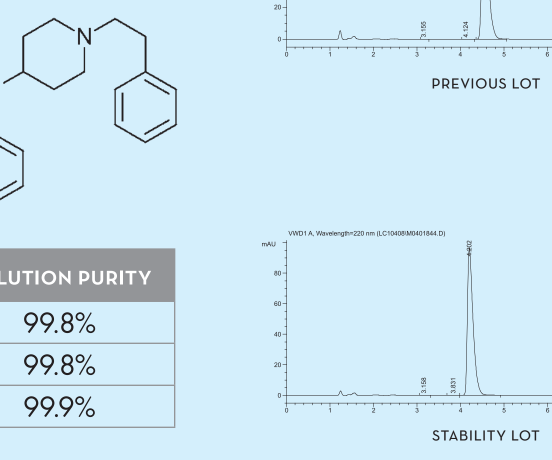
.. Acetonitrile::0.02M Ammonium Acetate, pH=3

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science, smarter.

			H ₃ C	
SOLUTION	LOT NUMBER	MANUFACTURE DATE	% CONC. DIFF FROM NEW PREP	SOLUTION PURITY
New Lot	FE022508-02	2 / 2008	-	99.8%
Provious Let	25215 25B	3 / 2006	0.5	00.8%

1 / 2003



AMPOULED CERTIFIED SOLUTIONS ARE EFFICIENT, ACCURATE & CONSISTENT

Single use format produced in large lots

- Low risk of contamination More efficient use of material
- Improved consistency and accuracy
- Single lot used over longer periods of time and across locations Reduces labor and time for routine standard preparation at the bench
- Sealed containers and inert environment protect against evaporation and degradation
- Solution stability established through testing • DEA exemptions for solutions of controlled substances available

High quality Certified Solution Standards and Reagents are an excellent alternative to the use of neat materials for clinical and toxicology applications

Linearity (r): ..

Mobile Phase:

Flow Rate:

Isil Dilek PhD, Kevin Gates, Rebecca Johnson, Sherri Pogue, Mitzi Rettinger, Uma Sreenivasan PhD

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