Design and Development Challenges of Natural Products Certified Reference Solutions

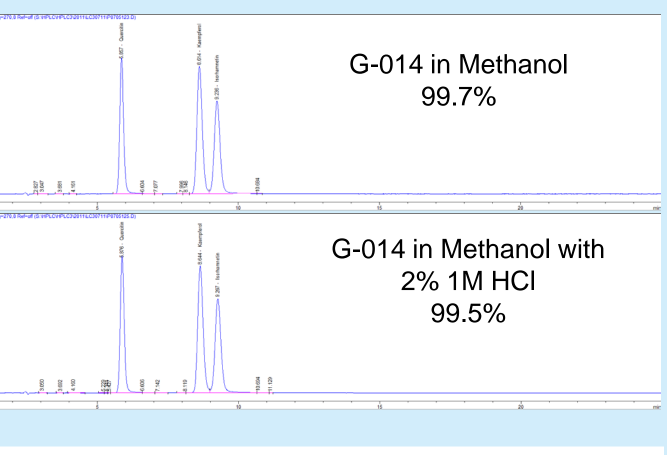
Authors: Tamara Tarbox, Isil Dilek, Derrell Johnson, and Mitzi Rettinger	Pre-Formulation Studies					
Abstract Laboratories often encounter multiple technical challenges in natural products testing of botanicals such as Ginkgo biloba, Ginseng, and Green tea. These phytochemicals offer significant handling issues due to inherent problems with hygroscopicity or sensitivity to air and/or light, which makes preparation of an accurate and consistent reference standard problematic. Weighing accuracy of milligram amounts of phytochemicals can be negatively affected by balance environment and weighing technique. Without consideration of proper diluent selection and controls in weighing, handling and solution preparation, accuracy and stability of phytochemical solution reference standards can be impacted. This poster will discuss these factors and their influence on the design, preparation, and certification of selected multi-component phytochemical solution standards.	 Useful Product Development Tool: Pre-Formulation Study Potential formulations evaluated for: Solubility – several solvent systems investigated at once Compatibility with analysis technique(s) Stability in solvent Soluble formulations are dispensed into ampoules for accelerated stability testing Stability is determined by measuring purity of samples stored at different temperatures such as: sub-freezer, freezer, refrigerator, ambient, and 40°C 					
Introduction	G-015, Ginseng Ginsenosides Mix Pre-Formulation Study Pre-Formulation Study					
 There are several factors critical to production of a high quality natural product reference standard including raw material handling, characterization and potency; certification and qualification of solutions; and homogeneity and stability of the solution. Certified phytochemical reference standard solutions prepared in a diluent that promotes stability and packaged under argon in flame-sealed ampoules can be stable for many years. Accurate results depend on accurate reference standards. Cerilliant addresses the challenges associated with making certified phytochemical reference standards. High purity, well-characterized components High purity diluents and/or stabilizers, compatible with the compound(s) Pre-formulation solubility and accelerated stability studies Environmental controls to ensure integrity of the compounds e.g. glove boxes 	 Three formulations, 100 µg/mL each component: (80:20) Acetonitrile:Water (60:40) Water:Methanol (60:40) Water:Methanol with 5% 1M HCl Acetonitrile-containing formulation: Solubility issues, no accelerated stability Acid-containing formulation: Initial purity value significantly lower At temperatures ≥ refrigerate, rapid degradation Sub-freezer samples did not appear to degrade Pre-Formulation Study Three formulations, 100 µg/mL Acetonitrile Methanol Methanol Methanol with 2% 1M HCl Acetonitrile-containing formulation: Initial purity value significantly lower At temperatures ≥ refrigerate, rapid degradation Sub-freezer samples did not appear to degrade 					
 Preparation using accurate, calibrated, and qualified balances 	Ginseng Ginsenoside Mix, G-015, Pre-Formulation Study Results Summary (Sum of %Peak Area for Eight Components) Ginkgo Biloba Flavonoids Mix, G-014, I Chromatograms for Initial Time					
 Accuracy in solvent addition using a gravimetric approach Traceability of all components to container and lot Analysis to verify accuracy, homogeneity, and consistency Appropriate packaging and storage Assessment of shelf life 	Chromatograms for Initial Time Timepoint Storage Condition Sub-freezer Freezer Refrigerate Room Temp. 40°C G-015, 100 µg/mL each Component, in (60:40) Water:Methanol with 5% 1M HCl (% Peak Area) 60:40) Water:Methanol with 5% 1M HCl 60:40) Water:Methanol with 5% 1M HCl					
Neat Material Certification	Initial 88.9 88.9 88.9 88.9 88.9					
 Numerous techniques may be utilized for characterization activities. Testing plans are tailored to each raw material and special consideration is given to the material's intrinsic properties. Purity, potency and impurities Mass balance – orthogonal approach 	1 week89.588.771.920.09.012 weeks89.388.3Not analyzed due to degradation noted at 1 week timepointImage: Component, in (60:40) Water:MethanolImage: Component, in (60:40) Water:MethanolImage: Component, in (60:40) Water:Methanol					
 Multiple techniques for chromatographic purity and residuals Based on ISO Guide 34; used by NIST Chromatographic purity Use of two techniques and different columns – values must agree within 0.5% Purity and related substances – resolution of known impurities Residual impurities Residual water by KF – USP <921> Residual solvent by GC/HS – validated Cerilliant method or USP <467> Residual inorganic content – micro ROI based on USP <281> NMR evaluation, EA and other techniques possible 	(% Peak Area) Initial 97.3 97.3 97.3 97.3 1 week 97.5 97.4 97.1 96.2 96.6 2 weeks 97.4 97.3 97.3 97.4 96.6 4 weeks 96.7 96.2 96.1 96.8 No sample Temperatures Ranges Sub-freezer: -70°C to -80°C Ereezer: -10°C to -25°C G-01					
 Identity – multiple techniques and comparison to literature references 1D and 2D NMR – Proton, Carbon-13, other nuclei GCMS, LCMS, LCMSMS, QTOF Other techniques as needed: EA, optical rotation, DSC, melting point, TGA 	Freezer: -10°C to -25°C Refrigerate: 2°C to 8°C Room Temperature: 15°C to 30°C					
All of the information collected from the neat material characterization is used to assign the purity factor for the neat material using a mass balance equation. $\frac{c_{nrom.}}{Purity(%)} \frac{Residual}{Solvent} \frac{Residual}{Solvent} \frac{Residual}{Solvent} \frac{race}{content(%)} \frac{Partiry}{content(%)} \frac{Partiry}{Pactor for} \frac{Partiry}{Pactor} \frac{Pifference}{from Chrom} \frac{Purity(%)}{Partiry(%)} \frac{Partiry}{Solvent} \frac{Partiry}{Solven} \frac{Partiry}{Solven} $	 Certified Reference Standard Preparation To ensure accurate preparation of the reference standard solution, strict manufacturing controls are used. material properties (such as hygroscopicity, light or oxygen sensitivity) other considerations must be made of the process. Robust manufacturing practices are critical to the accuracy and lot-to-lot consistency of each considerations made for reference standard preparation include: Material and equipment requirements Hygroscopicity, static potential, sensitivity to air or light Room selections, environmental controls – glove box Gravimetric preparation – higher precision than volumetric 					
W3Sobrents: the weight percentage of residual sobrents present in the neat material W3Sobrents: the weight percentage of water present in the neat material W3Sobrents: the weight percentage of inorganic content in the neat material ComPutity: based on the chromatographic purity of the specified primary purity method: either GC or HPLC Cercific Control Analytical Reference Standards Control Control Standards Stand	 Better flexibility in determining batch size Traceability with weigh tapes Dispensing Equipment checks, line purges, segregation Single use volumes flame-sealed under inert atmosphere Sampling plans – allows batch homogeneity verification 					

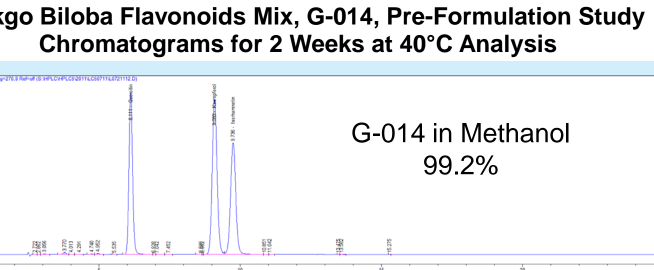


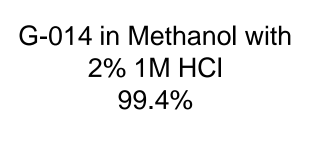
inkgo Biloba Flavonoids Mix ulation Study

- ormulations, 100 µg/mL each component: etonitrile
- thanol
- thanol with 2% 1M HCI
- itrile-containing formulation:
- lubility issues, no accelerated stability nol formulations:
- nimal differences in purity values over 2 weeks d formulation had higher purity at 40°C
- other purity values higher in Methanol without use additives

o Biloba Flavonoids Mix, G-014, Pre-Formulation Study **Chromatograms for Initial Timepoint Analysis**







15 20 mic

acturing controls are used. In addition to specific siderations must be made regarding each step lot-to-lot consistency of each product.

Weighing Accuracy



Hygroscopic materials are handled in glove box – inert atmosphere, relative humidity $\leq 5\%$

e environment and weighing technique nificantly influence reference accuracy

Analytical Verification and Certification

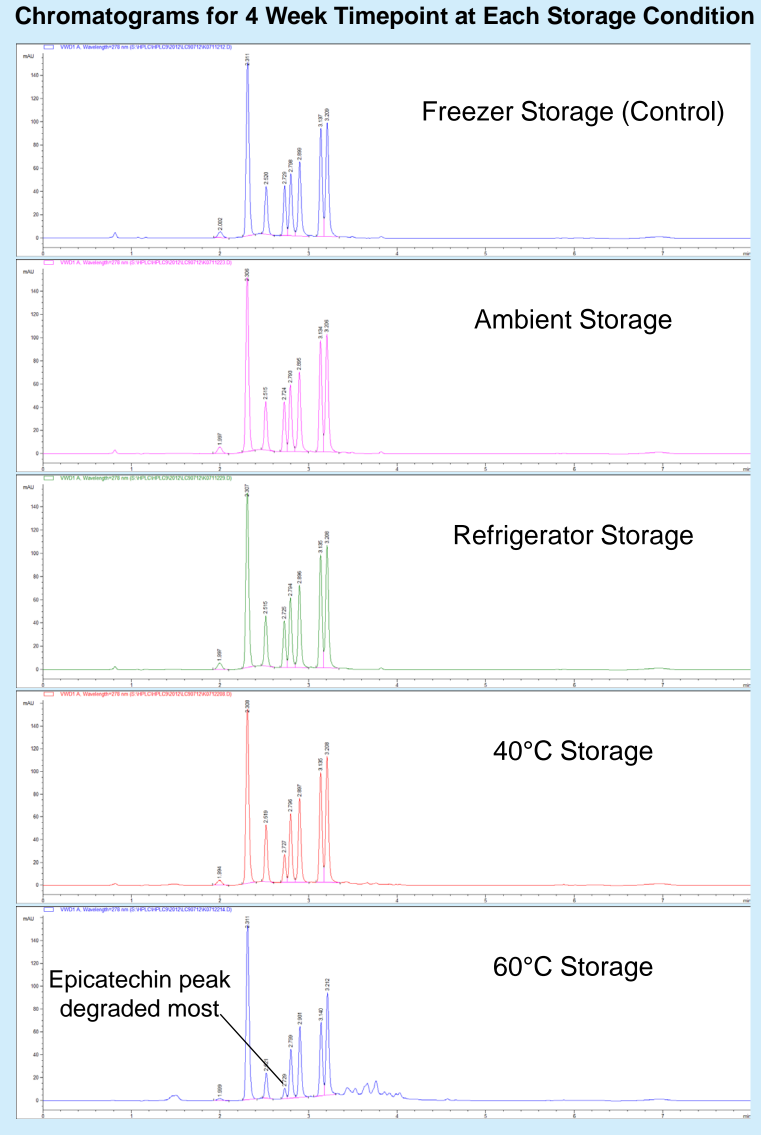
Analytical verification and certification, or release testing, comprises accuracy, consistency, homogeneity, and purity testing of the solution via HPLC/UV under various columns and run conditions. Accuracy of the prepared concentration is verified by comparison to a calibration curve or standard – a primary source or certified second source. This comparison of multiple independent preparations also includes previous lots where available which ensures lot-to-lot consistency of reference standard preparations. Homogeneity is confirmed across each batch of reference solution prepared. The solution purity is evaluated for consistency with the neat material to rule out degradation and contamination.

Release Testing Data for Natural Products Reference Standards	Catalog Number	Name	Number of Components	Accuracy: % Difference from Curve	Homogeneity: %RSD of Samples	Solution Purity (Sum of Components)
	G-013	Ginkgo Biloba Terpene Lactones Mix	5	-0.76 to 0.73	0.15 to 1.40	94.9%
	G-014	Ginkgo Biloba Flavonoids Mix	3	-0.86 to -0.21	0.26 to 0.33	99.9%
	G-015	Ginseng Ginsenosides Mix	8	-1.30 to 0.73	0.78 to 2.50	98.1%
	G-016	Green Tea Catechin Mix	8	-3.41 to -0.29	0.50 to 1.33	99.7%

Stability Testing

- Solution stability or expiration (shelf established through long-term stabilit
- Solution purity/concentration eva regular intervals – up to five yea expiration dates
- Properly designed & prepared sc standards stable for years vs. we which is typical for most lab prep
- Most accurate information, but re to complete
- Accelerated stability studies are a useful tool in estimating shelf life:
 - Comprehensive, formalized approach needed to predict long-term stability
 - Predictive stability study performed for G-016 using Arrhenius equation

Predictive Stability Study of G-016



Conclusions

Certified reference solutions offer significant advantages over neat reference materials, especially with regard to ease of use and handling of sensitive materials. Despite the challenges associated with handling and characterizing natural product materials, the use of a variety of development tools enables the formulation of accurate, consistent, and stable reference solutions.

Through the use of fully characterized neat materials, a mass balance approach to determining purity factor, pre-formulation studies, careful material handling and manufacturing controls, and reference standard verification against appropriate calibrators, natural product reference standards have successfully been developed.

life) is ity studies: aluated at ars to assign	Real-Time Stability of Anthocyanins							
	Catalog Number	Name	Neat Material Purity (%)	Solution Standard Purity (%) Release Testing	Solution Standard Purity (%) 4 years			
	C-069	Cyanidin-3-glucoside	94.6	95.2	93.1			
olution veeks or months parations requires years	C-070	Cyanidin-3-galactoside	94.6	93.4	82.3			
	P-057	Petunidin-3-glucoside	91.9	91.1	92.1			
	P-058	Petunidin-3-galactoside	92.4	90.2	88.6			

Time points

Storage Temperature

Rate (k) Plots

Arrhenius Plot

Prediction from Model:

Degradation Rate Curves for Four Storage Conditions* Freezer Storage (Control) Ref Amb **Ambient Storage** ▲ 40°C • 60°C 25 30 15 20 Timepoint (Days) **Refrigerator Storage** Arrhenius Plot for Epicatechin in G-016 Determined from Degradation at Four Temperatures Arrhenius equation: 40°C Storage • $\mathbf{k} = \mathbf{A}\mathbf{e}^{-(\mathsf{E}a/\mathsf{R}\mathsf{T})}$ E y = -6127.7x + 19.02260°C Storage $R^2 = 0.8667$ • 0.0029 0.0030 0.0031 0.0032 0.0033 0.0034 0.0035 0.0036 0.0037 1/T *Time points and storage conditions the same as the predictive study

Green Tea Catechin Mix, G-016, Epicatechin Peak

Predictive Stability Study of G-016

3, 7, 14, 21, 28 days

Freezer (as control), Refrigerator, Ambient, 40°C, 60°C

Degradation vs. Time for each Temperature (T)

Rate (1/k) vs. Temperature (1/T) for Epicatechin

 \leq 5% degradation in 4.4 years of Freezer Storage