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Comparative DSC and oscillatory TMA studies carried out using a variety of cure conditions showed that simply increasing the heating rate did not suppress the residual cure, although in many cases it allowed the glass transition to become apparent before the onset of rapid residual cure. The results indicated that ensuring the reliability of dynamic mechanical measurements of the glass transition temperature is very difficult to accomplish without the knowledge of residual cure properties provided by methods such as DSC.						
14. ABSTRACT Using the dicyanate ester of bisphenol E (also known as Primaset LECy), we have illustrated the difficulties inherent in measurement of the glass transition temperature of this high-temperature thermosetting polymer via dynamic mechanical analysis alone. These difficulties result from the residual cure of samples heated beyond their glass transition temperatures.						
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Effect of In-Situ Cure on Measurement of Glass Transition Temperatures in High-Temperature Thermosetting Polymers

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- Background: Importance of Conversion
- Progress in Conversion Measurement
- Difficulties in Glass Transition Temperature Measurement
- Solutions that Work (and Some that Do Not Work)

Acknowledgement: Air Force Office of Scientific Research; AMG Group Members



Model High-Temperature Thermosetting Polymers: Cyanate Esters





Principal reaction: 3 OCN -> Selectivity: 80% -> 98% $\Delta H_f = -110 \text{ kJ / eq.}$ Conversion kinetics; Auto-catalytic (may be catalyzed) Typical conversion rate: ~50%/hr (max) at 250 °C (uncatalyzed) Extend of conversion: Limited to $T_G = T_{cure} + 30-60 \text{ °C}$

- A single, known reaction predominates.
- Methods for assessing the extent of side reactions, and for minimizing side reactions, are known.
- The structure of even fully cured networks is easily analyzed and described quantitatively.
- Samples are easy to prepare in the laboratory; cure conditions are readily manipulated over a very wide range of rates.

Cyanate Esters: Next-Generation High-Performance Composite Resin





Cyanate Ester Networks: Defined by Composition and Conversion



 In addition to glass transition temperature, many properties, such as density and water uptake, are mainly functions of conversion and monomer type.



Conversion and *T_G* Are Intrinsically Linked in Cyanate Esters



Conversion



- Under non-degrading conditions (T < 300 °C), if conversion changes, then the T_G must also change with it. If the T_G changes, then cure has occurred. The only observed changes should be increases over time!
- A measurement of conversion is a measurement of T_G . Measurements of conversion and T_G may be combined to yield a more precise estimate of both parameters.
- Any measurement that indicates a change in conversion, such as non-reversible heat flow in a DSC experiment, also measures the rate of change of T_G .
- The preceding statements are true for any thermosetting network that forms by a single reaction method and is not subject to degradation during measurement.



0

-0.05

-0.1

-0.15

-0.2

-0.25

-0.3

-0.35

-0.4

100

150

Heat Flow (W / g)

Typical Residual Cure



Even in samples with well-defined exotherms, there is ambiguity ...

At high conversions, the typical approaches become impossible

Temperature (°C)

200

What about this

case?

Looks like it melts.

except that it cannot

and does not melt!

250

300

350

-1st heating

 Current solution: live with absolute error of up to 0.05 in conversion by DSC, corresponding to predicted T_G uncertainty of 25 – 50 °C.



Re-Scanning Shows the Limitations of Typical Baseline Approach



Typical Residual Cure



0 1st heating -0.05 --- 2nd Heating Heat Flow (N / g) -0.15 -0.2 -0.25 -0.1 Enthalpy due to relaxation at the transition, followed by a little residual cure! -0.3 -0.35 -0.4 100 300 150 200 250 350 Temperature (°C)

What about this

case?

... and it works at high conversions

Re-scanning shows us where the baseline is above T_G , how T_G shifts, how T_G is partly hidden, and where instability starts

Better solution: use the re-scan as a baseline (Kessler group).

Building a Baseline from Re-Scanning



Typical Residual Cure





What about this

case?

Magnitudes of step change should match (roughly), shift in step change should fit diBenedetto equation ... and the exact same algorithm works well with samples at high conversion

Even better solution: use the re-scan + account for shift in T_G & instability (our idea).



Estimating Uncertainties



 Uses – range of possible offset values (light red); range of possible instability corrections (orange); estimated error in slope of extrapolation in re-scan using 0.05 W/g expected size of step change (purple). These errors are assumed to be uncorrelated. Assessment of New Baseline Method

Old Way: Re-scan with range of possible offsets = baseline

New Way



- Uncatalyzed LECy with cure at 170 °C / 24 hr + post-cure from none to 240 °C for 1 hr.
- Old way predicts glass transition temp at full cure = 282 °C; new method predicts 312 °C, experiments show it is at least 304 °C.



LECy, catalyzed with 2 phr nonylphenol + 160 ppm Cu as Cu(II)Acac Cured at 150 °C for 12 hours



 Oscillatory TMA done in accordance with typical procedure shows three peaks for this single-component system, one peak corresponds to an increase of stiffness.





• Excess heat release above 175 °C indicates that the sample is undergoing cure.



LECy, catalyzed with 2 phr nonylphenol + 160 ppm Cu as Cu(II)Acac Cured at 150 °C for 12 hours



 In the shaded region, the sample is undergoing cure, which means that the T_G is changing while an attempt is being made to measure it!







 The darker green lines indicate the uncertainty limits for T_G. Main sources of uncertainty are DSC to OTMA correspondence, and effect of degradation. Note that T_G increases faster than the scan temperature.



Why Not "Outrun" T_G ?





- Residual cure tends to show the universality associated with activated rate processes, which means that a change in scan rate and a shift in temperature are functionally equivalent.
- As a result, the T_G instability number tends to be insensitive to heating rate. A faster heating rate only provides a head start in the race against cure.



Even Very Rapid Scans Do Not Always Prevent *In-Situ* Cure (1)



Uncatalyzed LECy (ageing study), cured at 210 °C for 24 hrs



- In the 2010 sample, 10 °C / min. was a little too slow to prevent in-situ cure (the true T_G at about 240 °C) is visible as the "shoulder".
- In the 2012 sample, the true T_G is no longer discernible.



Even Very Rapid Scans Do Not Always Prevent *In-Situ* Cure (2)



Uncatalyzed LECy (unaged), cured at 210 °C for 24 hrs 2014 – < 1 yr stored at 5 °C



- The DSC scan shows clearly that *in-situ* cure begins immediately after passing the true T_G at about 240 °C.
- Although the loss peak in the OTMA captures the true T_G, both the storage modulus and tan delta measurements are affected by *in-situ* cure. Thus, even 50 °C/min. is only just fast enough for the OTMA scan.



Uncatalyzed Samples; After Heating 210 °C for 24 hrs



- No cure means the T_G does not change, so the cannot have ever been in the "flat" part of the curve.
- Only a change as drastic as passing through T_G would cause such a rapid initiation of cure.
- T_G values inferred in this manner are reproducible to within 5 °C; slower heating rates work well for samples with lower values of T_G at full cure.





- In polycyanurate networks (and any other thermoset polymer network for which a single, path-independent cure mechanism exists), conversion and the glass transition temperature are intrinsically linked.
- With careful consideration of the physical properties of these networks, DSC methods can be utilized to measure conversions to within 1-2% at >80% conversion.
- Dynamic mechanical methods (especially at slow heating rates) will not measure the glass transition temperature of undercured networks correctly, due to *in-situ* cure.
- In many cases, *in-situ* cure and the resulting measurement issues cannot be avoided simply by choosing a more rapid heating rate.
- DSC methods (especially at rapid heating rates) appear to be bestsuited for assessing "as cured" glass transition temperatures.

QUESTIONS?

