All data taken at Pacific Northwest National Laboratory (PNNL) Operator: Steven W. Sharpe and Robert L. Sams, sw.sharpe@pnl.gov Version 2.0, November, 2000

## Composite spectrum for BENZENE\_50T

Effective burden of composite spectrum: 1 part-per-million-meter (ppm-meter) at 296 K Equivalent concentration x path-length of composite spectrum:  $3.216 \times 10^{-6}$  grams/liter-meter

## Sample Conditions-

- Chemical name and CAS number: Benzene, C<sub>6</sub>H<sub>6</sub>: [71-43-2]
- Physical properties: M.W. 78.11 amu, F.P. –5.5 C, B.P. 80.1 C
- Supplier and stated purity: Aldrich, 99.99%
- Sample class: I (PNNL scale).
- Temperature of sample:  $49.96 \pm 0.02$  C
- Diluent: Sample back filled with ultra high purity nitrogen to 760±5 Torr
- Individual samples at 3.4488, 0.36209, 5.20364, 0.56018, 0.10540, 0.28654, 20.90, 10.3160, 15.55 and 0.19255 Torr. Path length = 19.94 cm. Final data is a composite spectrum.
- Preparation: Multiple freeze-thaw cycles at 77K to remove air.

## **Instrument Parameters-**

- Bruker-66V FTIR, temperature controlled environment, evacuated optics bench
- Modified to include second aperture, between interferometer output and sample cell. This substantially reduces both "ghosting" and warm aperture effects.
- Spectral range: 6,500 to 600 cm<sup>-1</sup> (1.534 to 16.667 microns)
- Instrumental resolution based on maximum interferometer displacement is 0.112 cm<sup>-1</sup>
- Spectral intervals after FFT: 0.06 cm<sup>-1</sup>
- Interferogram zero-fill: 2X
- Apodization: Boxcar
- Phase correction: Mertz
- Beam splitter: Potassium bromide (KBr)
- IR source: Carbide glowbar (22 V)
- Scanner velocity: 60KHz (HeNe crossing frequency)
- Number of interferograms averaged per single channel spectra: 256
- Detector: Mid-band HgCdTe, photoconductive, 77K operation
- Folding limits: 15798 to 0 cm<sup>-1</sup>

## Post Processing and Related Parameters-

- Non-linearity detector correction (Bruker proprietary) applied to interferogram ( =0.85, =530)
- Composite spectrum created from 10 individual absorbance (base-10) spectra via classical least squares fit: Intercept=0, slope is fitted, individual absorbance values weighted by T<sup>2</sup> (transmission squared), all absorbance values 1.6 are given zero weight
- Calculated and estimated errors: Type A = 0.54%, Type B 3%
- Frequency correction (already applied): V(corrected) = V(instrument)\*0.999998+2.164x10<sup>-5</sup>
- Axis units: X=wavenumbers (cm<sup>-1</sup>), Y=Absorbance (base-10)
- Baseline correction via 7<sup>th</sup> order polynomial