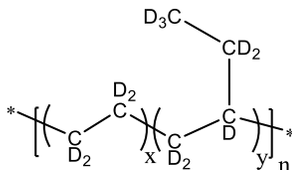


Sample Name:

Deuterated Poly(ethylene-d₄-co-butylene-d₈)

Sample #: **P42213A-dPE**

Structure:



Composition:

$M_n \times 10^3$ (g/mol)	M_w/M_n
109.0	1.02

Thermal properties:

Melting point, T_m	Crystallization point, T_{cr}
44 °C	37 °C

Synthesis procedure:

Deuterated poly(ethylene-co-butylene) was obtained by deuteration of poly(1,4-butadiene-d₆), which was synthesized by living anionic polymerization of butadiene-d₆ in non-polar solvent.

Characterization:

Deuterium NMR spectroscopy was used to confirm the structure of polybutadiene-d₆ rich in 1,4-addition.

The complete deuteration of the product was confirmed by FT-IR spectroscopy analysis by disappearance of alkene double bond (C=C at 971 cm⁻¹).

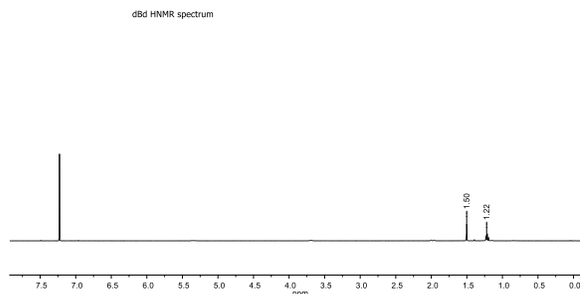
The molecular weight and polydispersity index were obtained by size exclusion chromatography (SEC) of poly(1,4-butadiene-d₆) precursor using THF as an eluent; and the molecular weight of polyethylene-d₄ was calculated accordingly.

Thermal analysis was performed on TA Instruments Q100 differential scanning calorimeter (DSC) under a nitrogen atmosphere at a scan rate 10 °C/min.

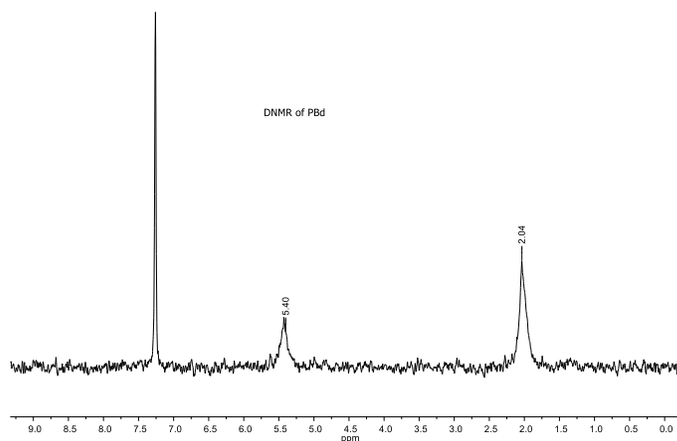
Solubility:

The product is soluble in hot toluene and xylene. The obtained solution has light ivory color; this coloration is due to the presence of trace amount (we expect <5–6 ppm) of the Wilkinson catalyst used in synthesis (and which is hard to remove from the final product).

H NMR spectrum of dPBd precursor:



D NMR spectrum of PBd precursor:

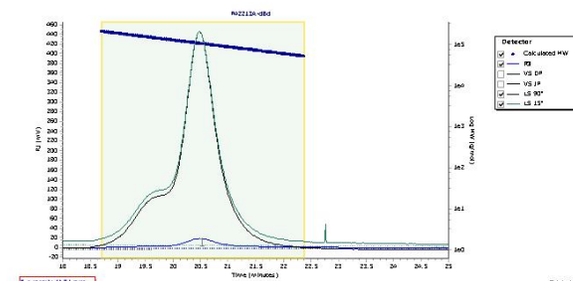


SEC chromatogram of dPBd precursor:

Agilent GPC/SEC Software

P42213A-dBd

Chromatogram Plot

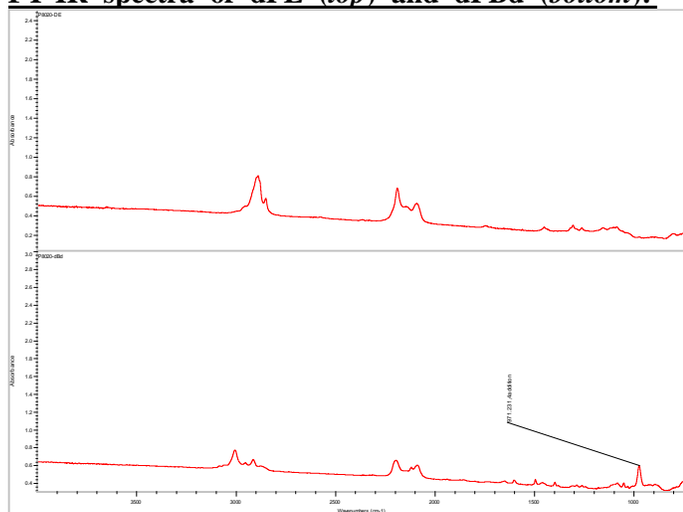


Molecular Weight Averages

Peak	Mp (g/mol)	Mn (g/mol)	Mw (g/mol)	Mz (g/mol)	Mz-1 (g/mol)	Mv (g/mol)	PD
Peak 1	108216	105047	109417	113920	118770	111987	1.042

Processing Parameters

FT-IR spectra of dPE (top) and dPBd (bottom):



DSC thermogram:

heating (bottom) and cooling (top) scans at 10 °C/min.

