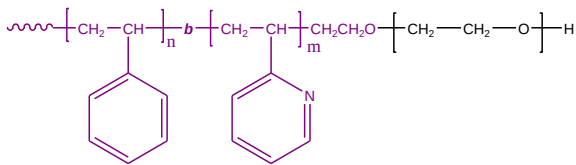


**Sample Name:**

Poly(styrene-b-2-vinyl pyridine-ethylene oxide)

**Sample #:** P18203-S2VPEO**Structure:****Composition:**

Mn x 10 <sup>3</sup> S-b-2VP-b-EO	PDI
166.0-b-304.0-b-45.0	
Calculated from <sup>1</sup> H-NMR	1.2

**Synthesis Procedure:**

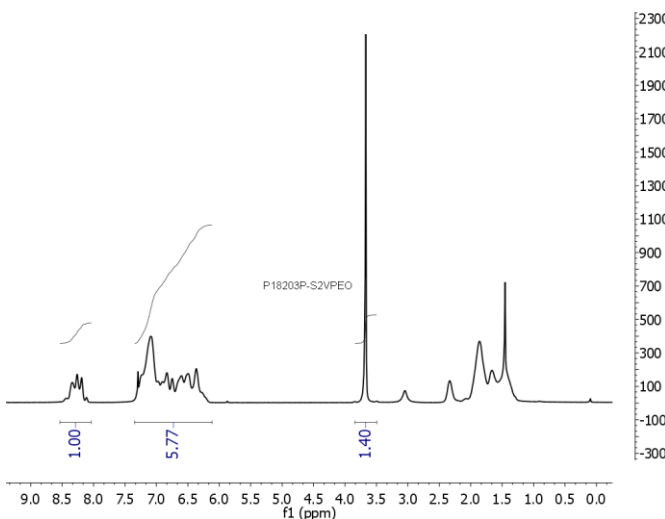
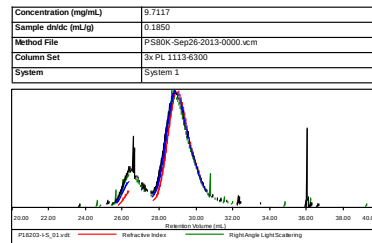
Poly(styrene-b-2-vinyl pyridine-ethylene oxide) triblock copolymer is prepared by living anionic polymerization by successive addition of monomer using cumyl potassium as initiator.

**Characterization:**

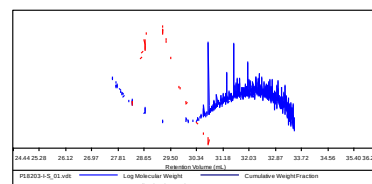
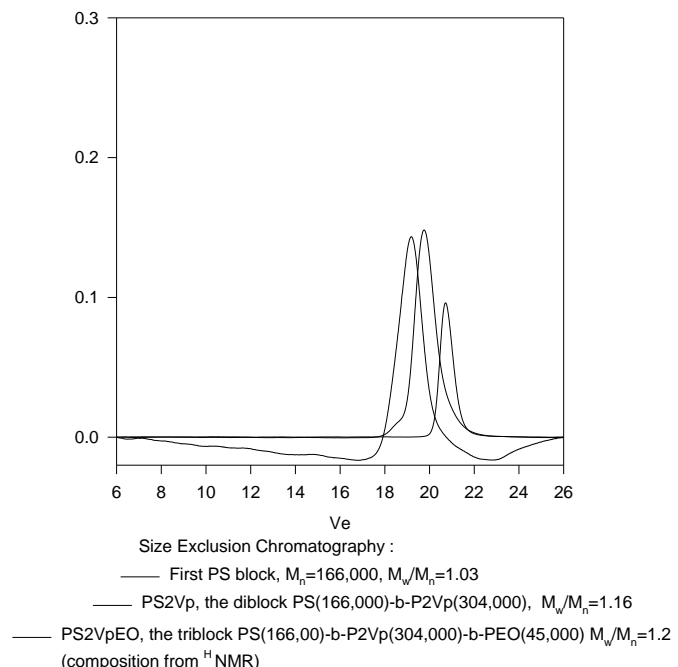
Polymer at different stages of polymerization was analyzed by size exclusion chromatography (SEC). The Block copolymer composition was then calculated from <sup>1</sup>H-NMR spectroscopy .

**Solubility:**

Poly(styrene-b-2 vinylpyridine-b-ethylene oxide) is soluble in THF, toluene, and CHCl<sub>3</sub>.

**<sup>1</sup>H-NMR Spectrum of the polymer S2 VPEO triblock copolymer****Sample ID: P18203-I-S**

Sample	Mn	Mw	Mp	Mw/Mn	IV
P18203-I-S_01.vcd	166,877	170,427	159,807	1.021	0.8232

**SEC for the triblock polymer:  
P18203P-S2VPEO****References:**

1. S. K. Varshney, X. F. Zhong and A. Eisenberg *Macromolecules* **1993**, 26, 701-706.
2. Gohy J.-F., Willet N., Zhang J.-X., Varshney S., Jerome, , pH dependence of the morphology of aqueous micelles formed by poly(styrene)-block-poly(2-vinylpyridine)-block-poly(ethylene oxide) copolymers, e-polymers 2002, 35.
3. Gohy, J.-F., Lohmeijer, B. Varshney S,K, Decamps B., Leroy E., Boileau S., Schubert U. S., Stimuli-responsive aqueous micelles from an ABC metallo-supramolecular triblock copolymer, *Macromolecules* 2002, 35, 9748-9755.
4. Gohy, J.-F., Mores S., Varshney S. K., Jerome, R., Self-organization of water-soluble complexes of a poly(2-vinylpyridinium)-block-poly(ethylene oxide) diblock and a fluorinated anionic surfactant, *Macromolecules* 2003, 36, 2579-2581.
5. Leil L., Gohy J.-F., Willet N., Zhang J.-X., Varshney S., Jerome R., Tuning of the morphology of core-shell-corona aqueous micelles: I. sphere-to-cylinder transition, *Macromolecules* 2004, 37, 1089-1094.

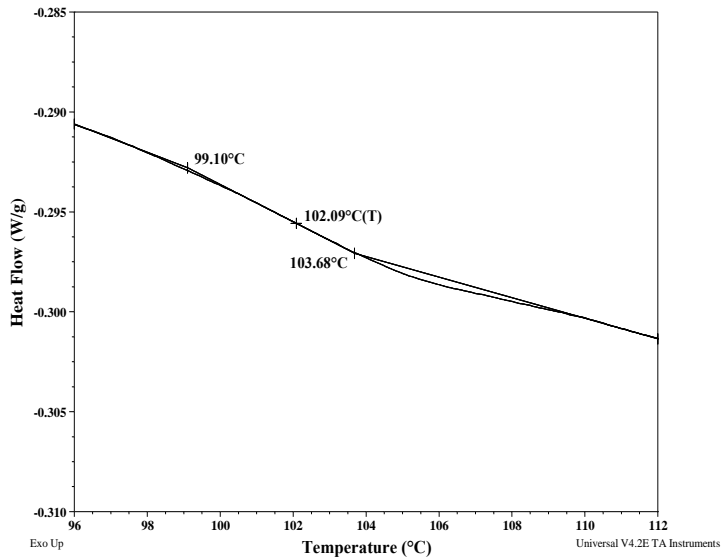
## Thermal Analysis of the sample S2VPEO

Thermal analysis of the samples was carried out on a TA Q100 differential scanning calorimeter at a heating rate of 10°C/min. The midpoint of the slope change of the heat flow plot of the second heating scan was considered as the glass transition temperature ( $T_g$ ).

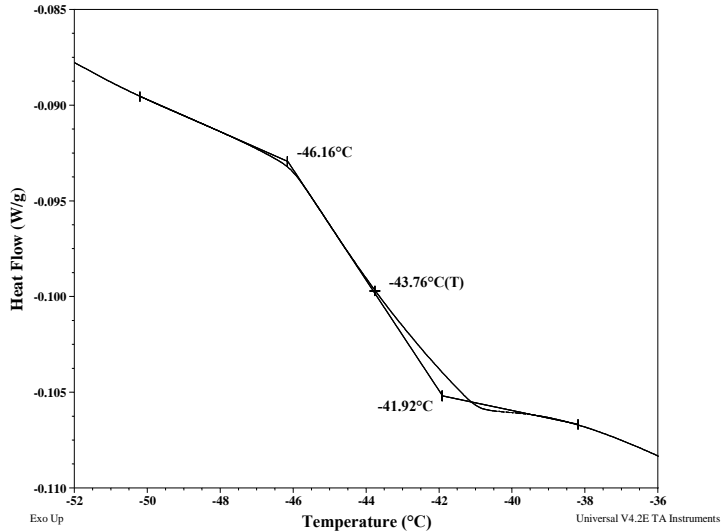
### Thermal analysis results at a glance

<b>For PS block:</b> $T_g$ : 102°C	<b>For 2VP block:</b> $T_g$ : Not distinct	
<b>For PEO block</b>		
$T_g$ : -44°C	$T_m$ : 61°C	$T_c$ : 34°C

### Thermogram for PS block:



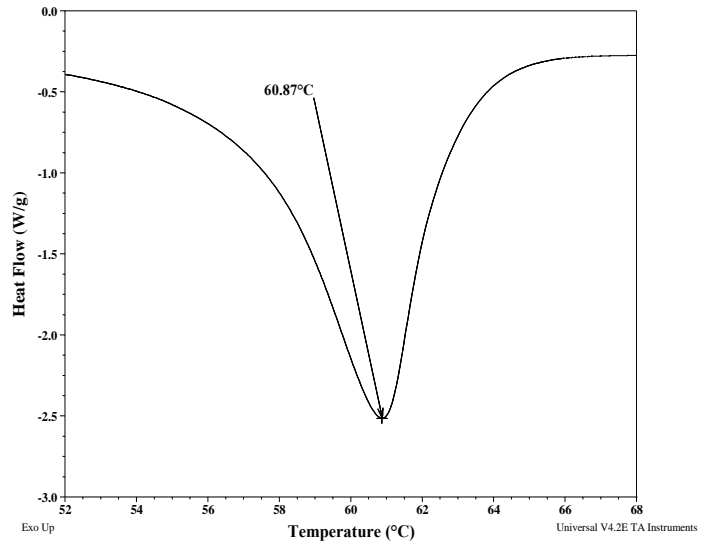
### Thermogram for PEO block:



## Melting and crystallization curve for the sample

The melting temperature ( $T_m$ ) was taken as the maximum of the endothermic peak where as the crystallization temperature ( $T_c$ ) was considered as the minimum of the exothermic peak.

### Melting curve for PEO block



### Crystallization curve For PEO block

