

# Synthesis, Thermal, and Rheological Evaluation of High C-Yield-Soluble Poly(phenylacetylene) Co-polymer Derivatives as New Carbon Fiber Precursors

Olumide (Matt) Agboola, Joseph Sengeh, Houxiang Li, Wei Zhu, Dr. T.C. Mike Chung\*  
Department of Materials Science and Engineering, The Pennsylvania State University, University Park, PA, 16801

## Introduction

The main precursors in the carbon fiber industry is **polyacrylonitrile (PAN)** and **pitch** which has an ineffective thermal transformation and is costly

Composite overwrapped pressure vessel for 5.6 Kg usable hydrogen

Energy cost (\$/kWh) System cost (\$/vehicle)

	Energy cost (\$/kWh)	System cost (\$/vehicle)
2013 system	\$17	\$3,200
2015 system	\$15	\$2,800
DOE Target	\$10	\$1,900



PAN precursor  $-\text{CH}_2-\text{CH}(\text{CN})-$

Pitch precursor  $\text{C}_{10}\text{H}_8$

**Advantages:**  
➤ Tension during thermal conversion  
➤ Low defects, good alignment, high tensile strength

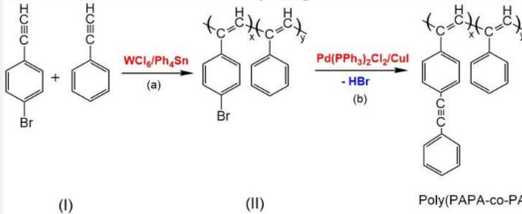
**Disadvantages:**  
➤ High cost, wet-spinning, 50% C-yield

**Advantages:**  
➤ Low cost, melt spinning, 70% C-yield  
➤ High elastic modulus

**Disadvantages:**  
➤ No tension for thermal conversion  
➤ Defects, poor alignment, low tensile strength

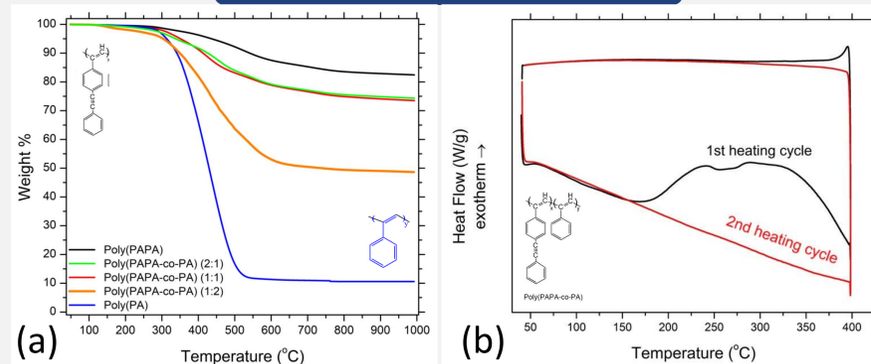
## Synthesis of Precursors

**Schematic 1.** Synthetic route to prepare poly(PAPA-co-PA) via (a) Metathesis (Ziegler-Natta) copolymerization and (b) sonogashira coupling.



**Figure 1.**  $^1\text{H}$  NMR spectra of (a) poly(4BrPa-co-PA) and (b) poly(PAPA-co-PA).  $^{13}\text{C}$  NMR of poly(PAPA-co-PA) (c) all in 1:1.

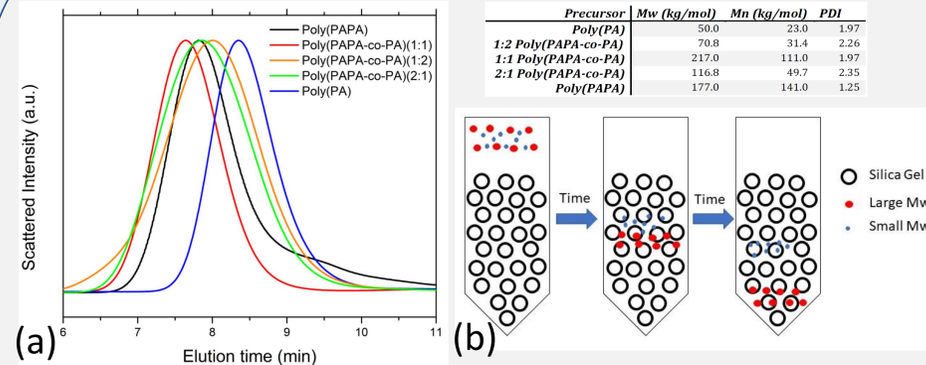
## Thermal Analysis Results



**Figure 2.** (a) Dynamic TGA measurements in  $\text{N}_2$  at  $10\text{ }^\circ\text{C}/\text{min}$  for poly(PA) and its derivatives and (b) DSC trace of poly(PAPA-co-PA) 1:1 also in  $\text{N}_2$  at  $10\text{ }^\circ\text{C}/\text{min}$ .

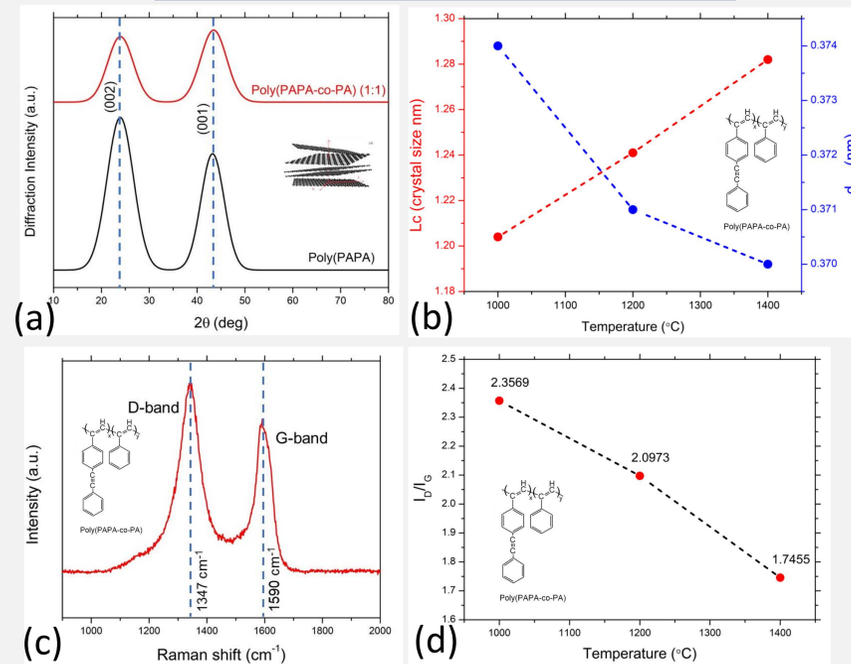
- TGA shows non-linear relation of the composition and the C-yield, which allows for the opportunity to minimize cost and maximize C-yield.
- Exothermic peak observed 1<sup>st</sup> but not 2<sup>nd</sup> cycle, indicating cross-linking.

## Molecular Weight Analysis



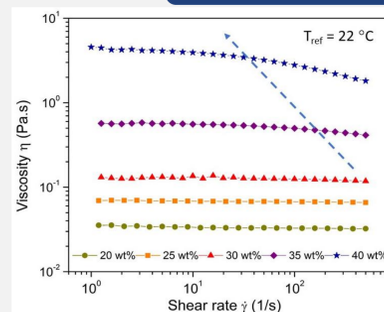
**Figure 3.** (a) Size exclusion chromatography (SEC) measurements of poly(PA) and its derivatives in THF at  $25\text{ }^\circ\text{C}$ . (b) Schematic of SEC methodology.

## Analysis of Carbonized Precursors



**Figure 4.** (a) 1D integrated XRD profiles for poly(PAPA-co-PA) and poly(PAPA) carbonized in argon at  $1400\text{ }^\circ\text{C}$ . (b) Crystal size and d-spacing trends with respect to increasing temperature for poly(PAPA-co-PA). (c) Raman spectra ( $\lambda = 514.5\text{ nm}$ ) of poly(PAPA-co-PA) carbonized in argon at  $1400\text{ }^\circ\text{C}$ . (d) Intensity ratios of the D and G bands at corresponding temperatures for poly(PAPA-co-PA).

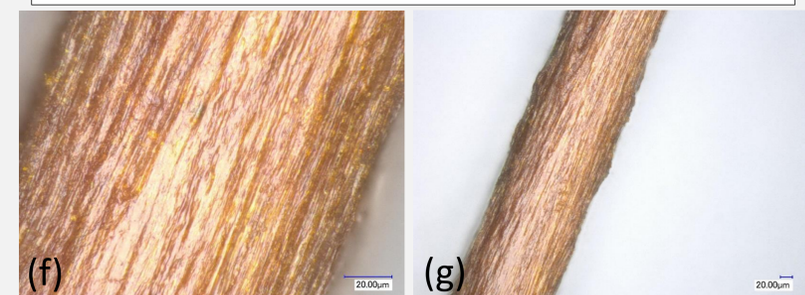
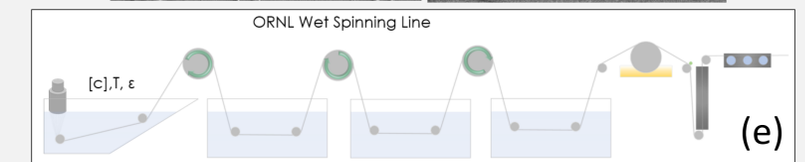
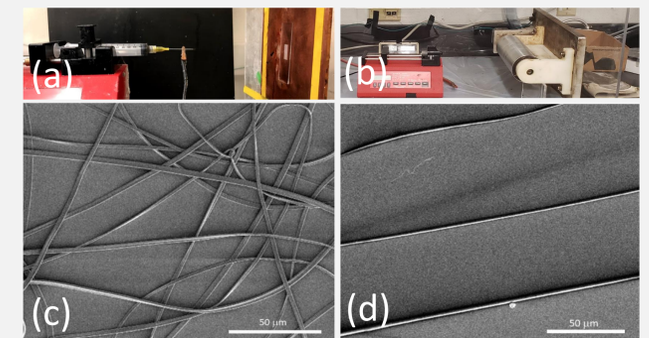
## Rheological Characteristics



**Figure 5.** Concentration dependent flow sweep rheological behavior of poly(PAPA-co-PA) (1:1) in toluene at  $22\text{ }^\circ\text{C}$ .

- Arrow indicates the on-set of shear thinning, which is a requirement of wet or electrospinning
- Within desired zero-shear viscosity range for solutions w/ shear-thinning.

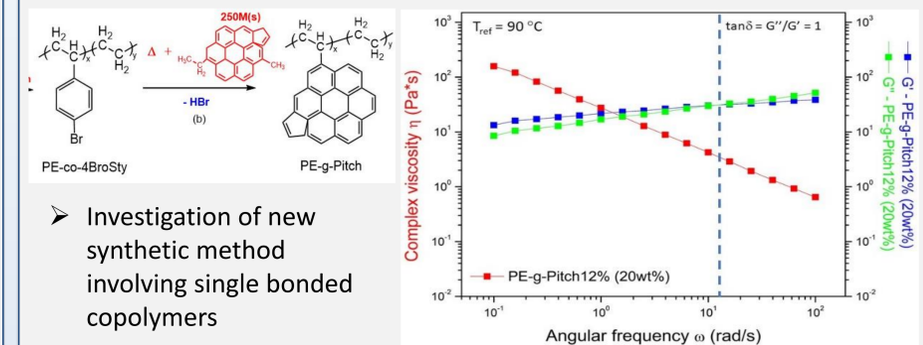
## Microfiber Processing and Morphology



**Figure 5.** Depiction of electrospinning apparatus for (a) disordered and (b) aligned microfibers. SEM micrographs of iridium coated (c) disordered (collected on flat glass slide) and (d) aligned poly(PAPA-co-PA)  $\sim 10$  micron fibers from 35 wt% THF solution, 10 mL/hr flow rate, 15 kV, a distance of 30 cm, and onto a 15 cm diameter cylindrical drum at 10 RPM. (e) Schematic of ORNL wet-spinning apparatus. (f) and (g) Show optical micrographs of wet-spun poly(PAPA-co-PA) ( $\sim 100$  micron) fibers.

## Conclusions / Future Work

- Reduction of fiber diameter and Mechanical properties of carbonized fibers are currently being tested and processed.



- Investigation of new synthetic method involving single bonded copolymers

## Acknowledgements & References

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