

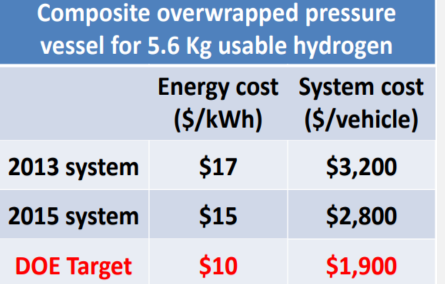
# Synthesis, Thermal, and Rheological Evaluation of High C-Yield-Soluble Poly(phenylacetylene) Co-polymer Derivatives as New Carbon Fiber Precursors

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#### Introduction

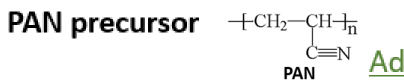
Science

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









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

#### Disadvantages:

- Low cost, melt spinning, 70% C-yield

Pitch precursor

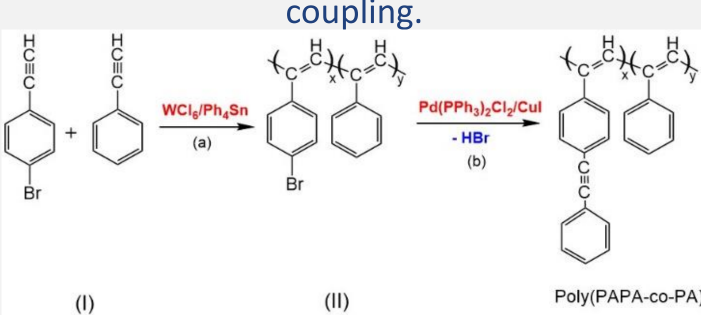
High elastic modulus

#### Disadvantages:

- No tension for thermal conversion
- ➤ High cost, wet-spinning, 50% C-yield ➤ Defects, poor alignment, low tensile

### Synthesis of Precursors

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



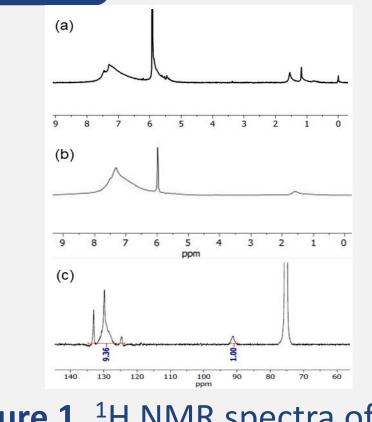
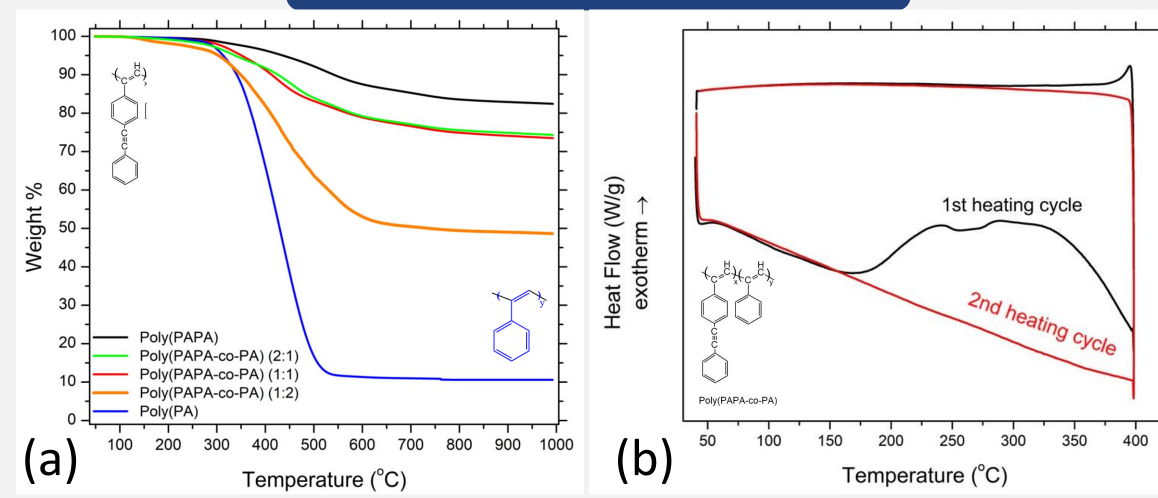


Figure 1. <sup>1</sup>H NMR spectra of (a) poly(4BrPa-co-PA) and (b) poly(PAPA-co-PA). <sup>13</sup>C NMR of poly(PAPA-co-PA) (c) all in 1:1.

# Thermal Analysis Results



**Figure 2.** (a) Dynamic TGA measurements in N<sub>2</sub> at 10 °C/min for poly(PA) and its derivatives and (b) DSC trace of poly(PAPA-co-PA) 1:1 also in N<sub>2</sub> at 10 °C/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.
- $\triangleright$  Exothermic peak observed 1<sup>st</sup> but not 2<sup>nd</sup> cycle, indicating cross-linking.

# Molecular Weight Analysis

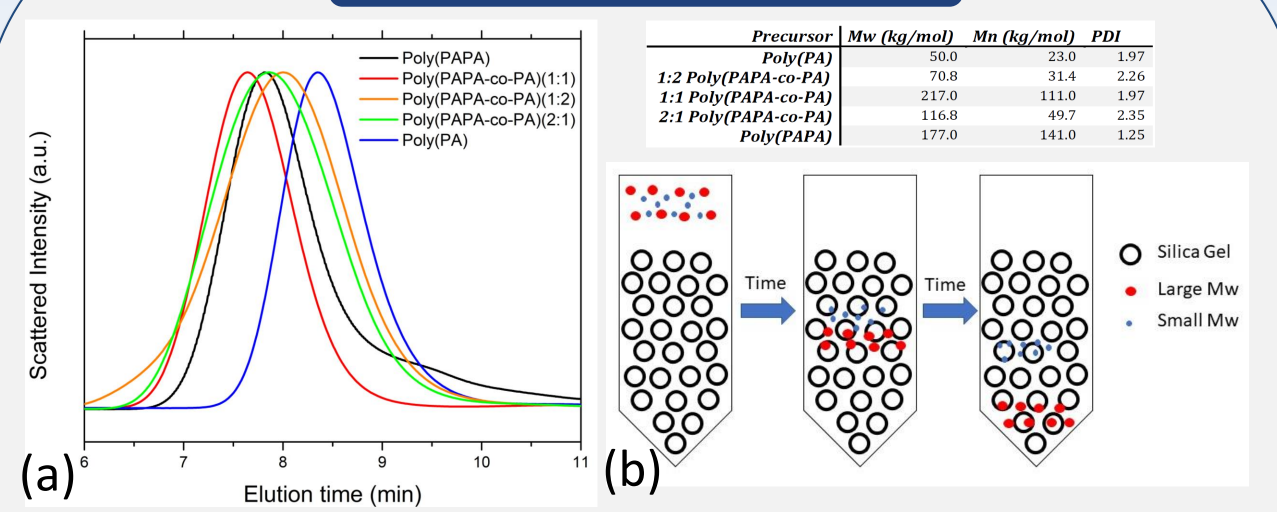


Figure 3. (a) Size exclusion chromatography (SEC) measurements of poly(PA) and its derivatives in THF at 25 °C. (b) Schematic of SEC methodology.

# Analysis of Carbonized Precursors

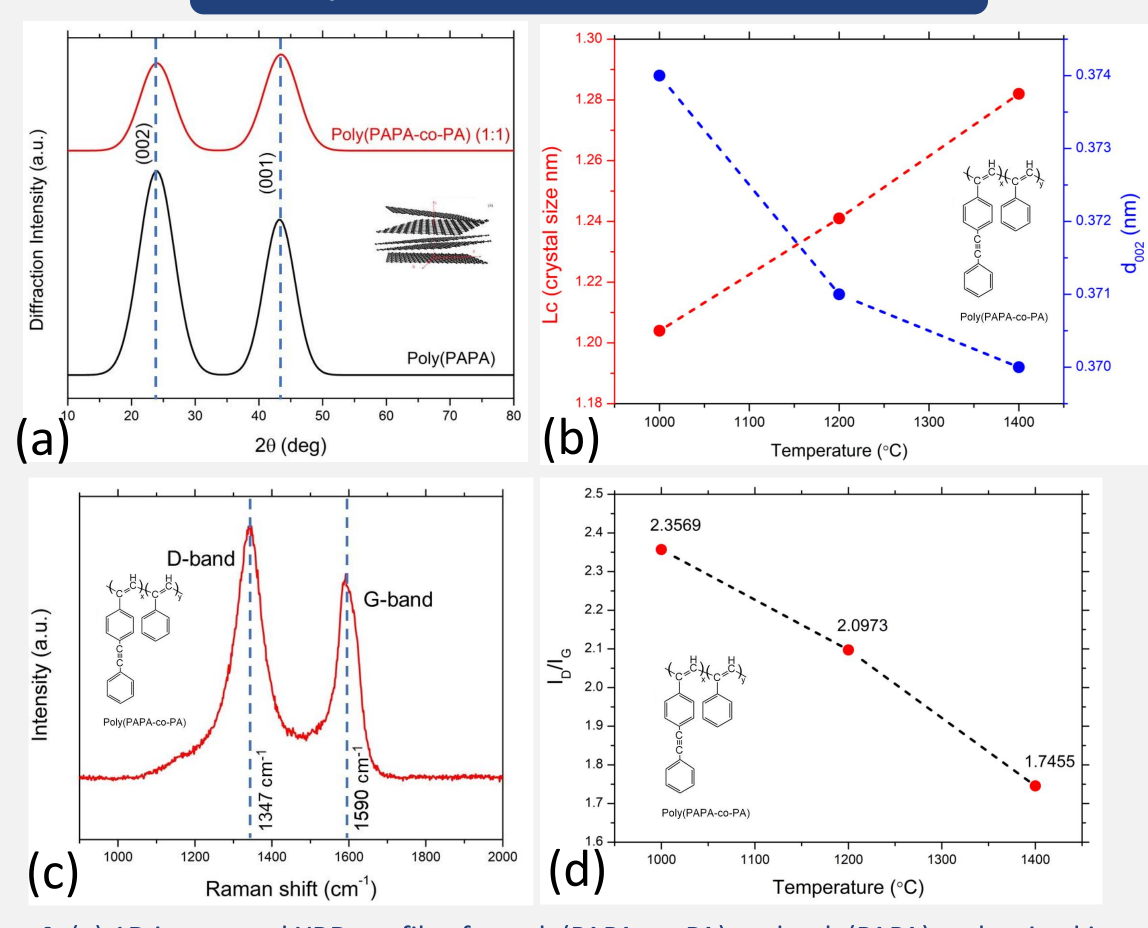
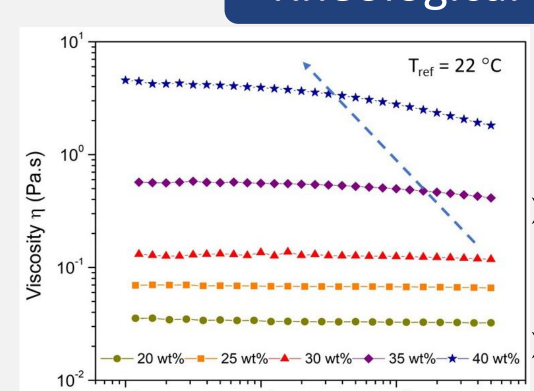


Figure 4. (a) 1D integrated XRD profiles for poly(PAPA-co-PA) and poly(PAPA) carbonized in argon at 1400 °C. (b) Crystal size and d-spacing trends with respect to increasing temperature for poly(PAPAco-PA).(c) Raman spectra ( $\lambda$  = 514.5 nm) of poly(PAPA-co-PA) carbonized in argon at 1400 °C.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

# Rheological Characteristics

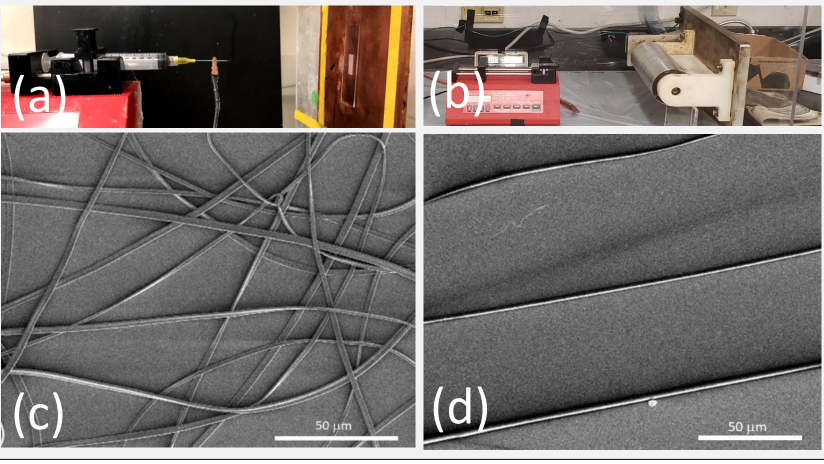


Shear rate  $\dot{\gamma}$  (1/s)

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



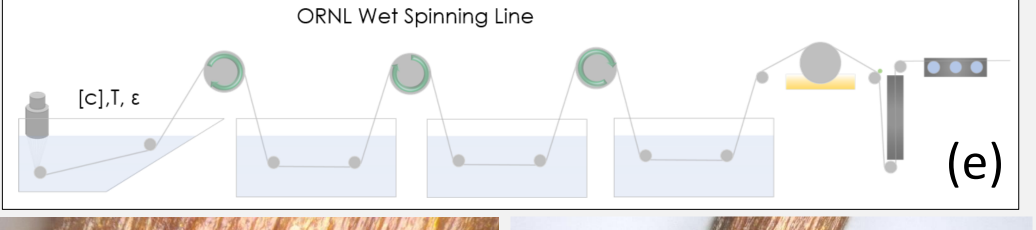
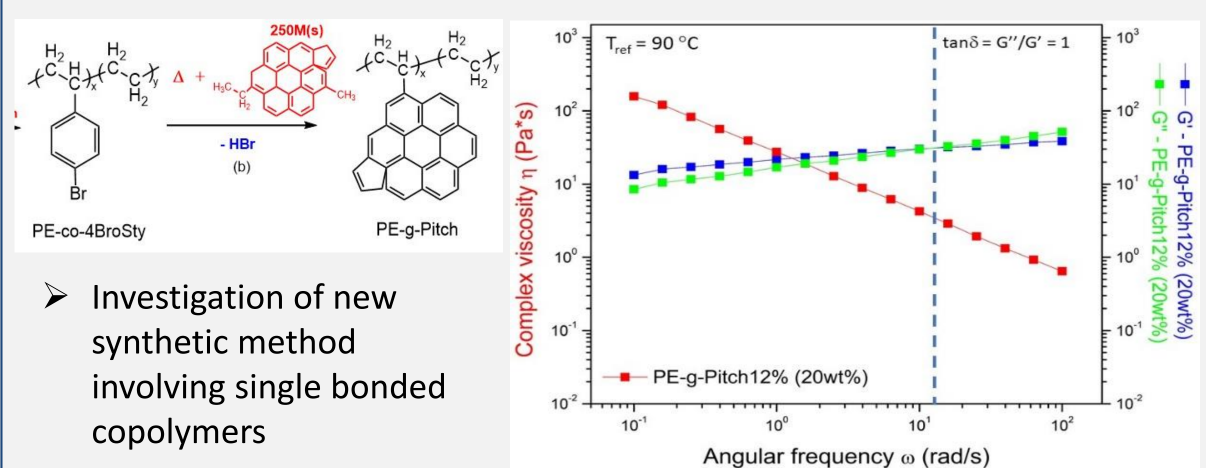




Figure 5. Depiction of electrospinning apparatus for (a) disordered and (b) aligned microfibers. SEM micrographs of iridium coated, (b) disordered (collected on flat glass slide) and (c) aligned poly(PAPA-co-PA) ~10 micron fibers from 35 wt% THF solution, 10 mL/hr flow rate, 15 kV, a distance of 30 cm, 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 wetspun poly(PAPA-co-PA) (~100 micron) fibers.

#### Conclusions / Future Work

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



# Acknowledgements & References

The author would like to thank the US Department of Energy for the financial support. Thank you to the Dr. Ralph Colby group for training on the AERES-RFS3 rheometer. 1. Weise, B. A.; Wirth, K. G.; Völkel, L.; Morgenstern, M.; Seide, G.; Carbon N. Y. 2019, 144, 351–361. 2. Schwartz M.; Encyclopedia of materials, parts, and finishes. 2nd ed. Boca Raton, Florida: CRC Press; 2002. 3. Cato Anthony D.; Edie DD; Carbon, **2003**, 41 (7), 1411-1417

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