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Conjugated Ionic Polyacetylenes Spontaneous Polymerization of 2-Ethynyl Pyridine in a Strong Acid

by

Sundar Subramayam and Alexandre Blumstein

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CONJUGATED IONIC POLYACETYLENES. 5: SPONTANEOUS POLYMERIZATION OF 2-ETHYNYLPYRIDINE IN A STRONG ACID¹⁻⁵

Sundar Subramanyam and Alexandre Blumstein[®] Department of Chemistry, Polymer Program University of Massachusetts, Lowell, MA 01854.

Abstract The spontaneous polymerization of the acetylenic bond in 2-ethynylpyridine in concentrated hydrochloric acid resulted in a substituted, extensively conjugated ionic polyacetylene bearing protonated pyridinium side groups. The conjugation length for the polymer was much greater than those obtained by a spontaneous processes via a Menschutkin reaction or by complexation with bromine. Base catalyzed deprotonation of the polymer resulted in a substantial decrease in backbone conjugation. The physical and spectral data prior to and after deprotonation correlate well with calculated values and provides insight into the structural and conformational characteristics of the polymer.

Introduction

In a series of recent papers, we have reported a new family of mono- and disubstituted ionic polyacetylenes that possess extensive backbone conjugation.¹⁻⁵ Such conjugated structures for disubstituted polyacetylenes were hitherto unreported. The polymerization reaction involved activation of the acetylenic triple bond in ethynylpyridine and its derivatives by quaternization via a Menschutkin reaction^{3,4} or by formation of donor-acceptor complexes with bromine.⁵ Figure 1 shows the activated monomers obtained in these reactions. Both methods were, however, limited by relatively low molecular masses (~4000) for the product polymers. The reaction mechanism in these spontaneous processes is an ionic one, since radical inhibitors do

not affect the polymerization. The overall process is similar to that observed for the spontaneous polymerization of 4-vinylpyridine upon quaternization by alkyl halides.^{6,7}

The low molar masses for these ionic polyacetylenes obtained via quaternization may be attributed to relatively slow activation of the triple bond (which results in low concentration of the activated species) compared to rapid propagation and termination steps. Fife et al. have shown that the quaternization of 4-vinylpyridine by ethyl bromide or ethyl tosylate is a rather slow reaction with $t_{1/2} = 1000$ min.⁸ In view of this, an alternative method was sought for a more rapid activation of the monomer. Such methods involve use of a highly reactive quaternizing agent in high concentrations.

In this paper, we report the polymerization of 2-ethynylpyridine by a strong acid. The reaction of 2-ethynylpyridine in concentrated hydrochloric acid resulted in spontaneous polymerization of the acetylenic triple bond giving a highly conjugated, ionic polyacetylene. The proposed structure for the polymer is illustrated in Figure 2.

Experimental

Materials: 2-Ethynylpyridine was obtained from Farchan Laboratories and distilled under vacuum before use. Concentrated hydrochloric acid was obtained from VWR Scientific Company and used as received.

Measurements: Infrared spectra were recorded on a Perkin-Elmer 1600 Series Fourier transform spectrophotometer using KBr plates. UV absorption spectra were obtained at 22° C in DMSO and chloroform on an IBM 9420 visible-ultraviolet spectrophotometer. Thermal analyses were carried out on a DuPont TGA 2950 thermogravimetric analyzer. X-ray diffraction patterns of unoriented samples were obtained on a Rigaku Geigerflex x-ray diffractometer equipped with a Wahrus flat plate camera using Ni-filtered Cu_{α} radiation. Dilute solution viscosity measurements were made in DMSO and chloroform at 25°C using a Cannon Ubbelohde dilution type viscometer. Molecular weight

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determination by vapor pressure osmometry was made on a UIC Inc. 070 vapor pressure osmometer in chloroform at 30°. Calibration was done with low molecular weight polystyrene ($M_n = 2000$) with an estimated error of \pm 5%. Fluorescence spectra were obtained on a Perkin Elmer MPF-44B fluorescence spectrophotometer equipped with a PE 150 Xenon source in conjunction with a Stanford Research System Model SR440 DC-300MHz amplifier and a SR400 two channel gated photon counter. All spectra were recorded in DMSO at 22°. Excitation wavelengths of 360 and 265nm respectively were used for the protonated and deprotonated form of polymer VIIIp. Elemental analyses were performed at Atlantic Microlabs, Atlanta, GA. *Poly(2-pyridiniumhydrochloride-2-pyridylacetylene) copolymer VIIIp:* To neat 2-ethynylpyridine (2.06g, 0.02mol) stirred at 0°, concentrated hydrochloric acid (0.71g, 0.02mol) was added dropwise. An exothermic reaction ensued and the reaction mixture acquired a dark color immediately accompanied by a rapid increase in viscosity. The polymer separated out as a black solid mass. It was washed repeatedly

with ether and dried overnight in vacuum at 40°. The dry polymer was obtained as a lustrous black powder (2.0g) that was readily soluble in water and DMSO but insoluble in chloroform.

IR (cm⁻¹) 3414, br (hydrogen bonded ring =N-H stretch), 1617 (backbone C=C stretch), 1605 (aromatic ring C=C stretch); UV, λ_{max} C=C (nm), 530; η_{inh} , DMSO(dL/g) 0.2. Anal. (Calc.), C = 70.88; H = 4.61; N = 11.81; Cl = 12.70, (Found), C = 70.25; H = 4.82; N = 11.80; Cl = 12.75.

Deprotonation of Poly(2-pyridiniumhydrochloride-2-pyridylacetylene) copolymer : The protonated polymer VIIIp (1.0g) was dissolved in 50ml of deionized water and to the stirred solution 100ml of 1M KOH was added slowly. Stirring was continued for 12 hours during which time the deprotonated polymer precipitated out as a dark solid. This was extracted with chloroform and the combined organic layers were dried and the

solvent evaporated under reduced pressure to yield a lustrous black solid. This was dried overnight in vacuum at 40°. The dry polymer (0.5g) was soluble in chloroform and DMSO but insoluble in water.

IR (cm⁻¹) 1634 (C=C), 1585 (aromatic C=C); UV, λ_{max} C=C (nm) 419; η_{inh} , DMSO,(dL/g) 0.2. Anal. (Calc.), C = 79.27; H = 4.82; N = 13.20; CI = 2.71, (Found), C = 75.20; H = 4.67; N = 12.33; CI = 3.01.

Results and Discussion

The polymer obtained by the bulk polymerization of 2-ethynylpyridine in concentrated hydrochloric acid is a black lustrous solid indicative of the highly conjugated nature of the backbone (Fig. 2). This is evident from its infrared spectrum, which reveals a strong absorption band at 1617 cm⁻¹ due to the C=C stretch from the backbone carbons. Furthermore, the \equiv C-H and -C \equiv C- stretching bands at 3290 and 2108 cm⁻¹, respectively, that are present in the monomer, are absent in the product polymer (Fig. 3). The highly charged nature of the polymer is evidenced by the broad, intense band at 3400 cm⁻¹ that arises from the =N-H stretch of the side group pyridinium rings and is indicative of the strong hydrogen bonding effect due to the charged nature of the side groups. This band is characteristic of all polymers reported by us previously possessing similar structures, including disubstituted polyacetylenes.³⁻⁵ The UV-visible spectrum of polymer VIIIp showed a strong absorption centered at 530 nm resulting from the π - π * transition of the conjugated polymer VIIIp. The polymer is soluble in DMSO and in water, but not in chloroform which is characteristic of its ionic nature.

The present polymerization can be compared to that of 4-vinylpyridine in concentrated acids reported by Salamone⁹ and later by Ringsdorf.^{10,11} The former study concluded that an anionic mechanism is unlikely in such strongly protic media and that the polymerization probably proceeds by a step growth process. The latter study

proposed a mechanism wherein the polymerization of the protonated vinylpyridine occurs in an ordered array of the pyridinium halide molecules. The polymerization of 2-ethynylpyridine also occurs in the protonated monomer which activates the acetylenic triple bond. Since the spontaneous polymerization occurs only in the presence of excess unquaternized monomer (no polymerization was observed when monomer was added to excess acid), we conclude that the initiation step occurs via the nucleophilic addition of unquaternized monomer to the triple bond of the quaternized activated species. This is also supported by the fact that the isolated monomer salt failed to undergo polymerization in concentrated HCI. The propagation step, therefore, appears to proceed in a manner analogous to that suggested by Ringsdorf et al., wherein immediate termination via protonation does not occur. The molecular mass of polymer VIIIp is, however, low ($\eta_{inh} = 0.2 dL/g$) suggesting that termination step(s) compete effectively with chain propagation.

A unique characteristic that distinguishes the present polyacetylene from the ones obtained by us via the Menschutkin reaction or complexation with bromine is that unlike the previously described systems wherein the charged side groups consist of either alkylpyridinium or bromopyridinium moieties, it has protonated pyridinium ring substituents. This allows for relatively easy deprotonation of the ionic polyene in basic media giving rise to a largely neutral species. Such a transformation in the previously reported materials would involve drastic conditions and elevated temperatures at which polymer degradation becomes a serious problem.

The analytical data of polymer VIIIp fits remarkably well with a structure wherein only 50% of pyridine rings in the polymer contain quaternary nitrogens, (Table 1). These results imply the incorporation of unquaternized monomeric units in the polymer chain even prior to deprotonation of the pyridinium ring side groups. A possible explanation for this could be the formation a dimeric complex between a protonated and an unprotonated monomeric species via intermolecular hydrogen bonding. This would

then provide sufficient activation for the triple bond in the unquaternized molecule for it to undergo polymerization. Scheme 1 illustrates the proposed reaction mechanism for the polymerization. Such dimer formation has also been reported in partially protonated poly(2-vinylpyridine) between two adjacent pyridine side groups one of which is protonated.¹² A charge transfer complex polymerization mechanism is also possible via the formation of a weakly reactive charge transfer complex biradical between the protonated and unprotonated ethynylpyridine molecules. The low molecular weight of the polymer and lack of temperature dependence of M, however, suggests that the latter mechanism is unlikely.

Treatment of VIIIp with aqueous potassium hydroxide solution gave a compound that possesses approximately 20% of the original charges (based on halide ion content). Table 1 lists the analytical data before and after deprotonation. Assuming that these residual charges are due to charged end groups that cannot undergo neutralization, the DP of the polymer is approximately 12 ethynylpyridine units. Although the deprotonated polyacetylene retained its black color, remarkable changes were observed in its spectral properties. The uv-visible spectrum showed a large blue shift for the π - π * transition of the conjugated polymer backbone compared to the protonated form. The absorption maximum shifted from 530 nm for the protonated form to 419 nm for the deprotonated one. Figure 4 shows the change in the absorption maxima of polymer VIIIp before and after treatment with base. The large hypsochromic shift observed in the present polymer on deprotonation results from reduced conjugation presumably from backbone twist introduced upon deprotonation and is not due to a change in the pyridinium chromophore. This is substantiated by the fact that the uv-vis spectra of protonated poly(2-vinylpyridine) and neutral poly(2-vinylpyridine) do not differ much, displaying absorption maxima at 260 and 250 nm respectively.¹² The backbone twist may be attributed to the absence of electrostatic interactions between the counterions and the pyridinum rings that were present in the protonated form, thereby resulting in the steric

interaction between the side groups becoming a dominating factor. This view is also supported by the fact that the λ_{max} (C=C) for the backbone undergoes a bathochromic shift on reprotonation from 419 to 460nm, trailing to ~ 800nm. The deprotonated polymer is no longer water soluble, but is readily soluble in chloroform due to its considerably diminished ionic character.

The reduced backbone conjugation in the deprotonated form of VIIIp is also supported by its IR spectrum, which shows a corresponding high frequency shift of the backbone C=C stretch from 1617 cm⁻¹ to 1634 cm⁻¹. The aromatic ring C=C band shifted from 1605cm⁻¹ for the protonated form to 1585cm⁻¹ for the deprotonated one, which is identical to that observed for 2-vinylpyridine (1586 cm⁻¹)¹³. The C=N stretch of the pyridine ring occurring at 1458 cm⁻¹ for the ionic form is shifted to 1467 cm⁻¹upon deprotonation, again being similar to that of 2-vinylpyridine (1472cm⁻¹).¹³ The intense band at 3400 cm⁻¹ due to the hydrogen bonded pyridinium ring =N-H stretch in the protonated form disappeared almost completely on deprotonation. Figure 5 shows the IR spectra of polymer VIIIp in its protonated and deprotonated forms. The latter spectrum contains weak but distinct bands at 3292 and 2102 cm⁻¹ from the acetylenic \equiv C-H and C≡C stretching found in the monomer. From the mechanism depicted in Scheme 1, the initiating chain end of polymer VIIIp is an ethynylpyridinium unit linked through the quaternary pyridinium nitrogen. The residual bands may, therefore, be attributed to the end groups in the polymer. This was confirmed by performing an end group analysis based on the acetylenic \equiv C-H stretching band. The relative intensity of this band at 3290 cm⁻¹ for the starting monomer was compared with that for polymer VIIIp after deprotonation. The intensities were normalized using the β -ring bending band of the pyridine ring. This band occurs at 738 and 749 cm⁻¹ for the monomer and polymer respectively, and its intensity is unaffected by the nature of the substituent on the ring i.e. alkynyl or alkeneyl (β-ring bending for 2-vinylpyridine occurs at 745cm⁻¹). Comparison of the infrared absorption ratio $\mathsf{Abs}_{\equiv\mathsf{C}\text{-}\mathsf{H}}\,/\,\mathsf{Abs}_{\beta\text{-ring}}$ for the monomer and deprotonated

polymer (2.30 and 0.24 respectively) yields a value of 9.1% for the end group content, which is in excellent agreement with the value calculated from analytical data (10%), (see Table 1).

TGA analysis of polymer VIIIp in nitrogen atmosphere showed a small, gradual weight loss (~9%) occurring at 150°C, followed by a rapid decomposition at 230° C. The deprotonated form showed an essentially similar behavior, suggesting that the ionic nature of the polymer prior to deprotonation does not contribute significantly to its thermal stability. The decomposition temperatures are similar to those for the ionic polyacetylenes obtained by us via the quaternization method (240°C)^{3,4}, and considerably higher than the ones synthesized by the formation of donor-acceptor complexes with bromine.⁵

A characteristic feature exhibited by these ionic polyacetylenes is a broad low energy band (560nm) in their fluorescence emission spectra. The phenomenon, is attributed to intramolecular excimer formation between the aromatic side group chromophores. It appears to be independent of the polymerization method, since the band was observed for polyacetylenes obtained via both the quaternization and complexation methods.⁶ It has also been reported for protonated poly(4vinylpyridines)¹² and isotactic polystyrene.^{14,15} In these ionic polyacetylenes, such excimer formation is indicative of a highly conjugated backbone with minimal twist, which is a prerequisite for efficient stacking of the pyridine rings. Such stacking seems to be largely due to a strong electrostatic interaction of each counterion with two adjacent pyridinum rings.

The fluorescence emission spectrum of the polymer VIIIp in DMSO, however, displayed two bands centered at 500 and 560 nm respectively, whose intensities are similar. The intensity of the band at 500nm was independent of polymer concentration, while that of the latter decreased slightly with increasing dilution (Figure 6a,b). This phenomenon is attributed to intramolecular excimer formation by the side group

chromophores. It is evident that two distinctly different chromophoric groups of approximately equal number (protonated pyridinium and neutral pyridine rings) are present in the polymer, substantiating the proposed structure (Fig. 2) which is based on its analytical data. The band at 560nm is assigned to the pyridiniur chromophores, since it was also observed for the ionic polyacetylenes obtained by previous methods⁵ wherein more than 98% of the side group pyridine rings were in the quaternized form. The emission at 500nm is attributed to the unquaternized pyridine rings which is expected to occur at a higher frequency. This was verified by the fluorescence spectrum of the deprotonated form, which showed a single emission at 500nm (Fig. 6c). The fluorescence emission due to intramolecular excimer formation of polymer VIIIp was also confirmed by the fact that its emission spectrum at 560nm is essentially the same as its absorbance spectrum. We believe that unlike flexible polymers, these polyacetylenes would not lose its excimer emission at very low temperatures due to their strongly conjugated an rigid backbones that predetermine the conformation of the side group chromophore. The configuration required for energy transfer is, therefore, built up in the polymer chains prior to excitation. Similar observations were made by Ishii et al¹² for protonated poly(2-vinylpyridine), which exhibits a "dimer-like excimer" due to interaction between protonated pyridine rings adjacent to one another. Direct measurements of fluorescence at 77°K were, however, not carried out due to instrument limitation.

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The x-ray diffractogram of an unoriented sample of VIIIp indicates an amorphous scattering with a broad ring centered at d = 11.8Å, possibly due to a layered structure. Similar diffraction patterns were also displayed by poly(2-ethylpyridiniumacetylene-iodide) obtained via the quaternization method and by poly[(2-bromopyridiniumy!)-acetylene-bromide] synthesized by the complexation method.⁵ In both cases, the broad small angle ring occurred at d = 11.8Å, which suggests that this layered structure is not dependent on the nature of the atom/group attached to the pyridine nitrogen located in

a

the 2- position with respect to the polymer backbone. This reflection may be related to the ordering of the side group pyridinium rings which would be the case if the polymer chain adopted a *trans-transoidal* conformation. The effect of *cis-trans* stereoregularity in substituted polyacetylenes on the x-ray diffraction patterns has been reported in a recent study. ¹⁶ It was found that for poly(*tert*-butylacetylene), the d-spacing increases with increasing trans content from 8.5Å to 10.5Å. The diffraction pattern of deprotonated VIIIp, however, does not exhibit this characteristic ring. This is probably due to destruction of the layered arrangement caused by the increased backbone twist resulting from deprotonation. Figure 7 shows the x-ray diffractograms of the polymer prior to and after deprotonation.

The conjugation lengths based on both IR and uv-visible spectra for VIIIp is, by far, the highest observed for these ionic polyacetylenes; the spontaneous polymerization of 2-ethynylpyridine via quaternization and halogen complexation methods resulted in polymers with absorption maxima between 370 and 490 nm for the former method^{3,4} and 470 nm for the latter.⁶ The extended conjugation observed in these systems in spite of their substituted backbones is surprising, and is in contrast to substituted polyacetylenes synthesized using coordination catalysts. Results obtained by us recently from modelling studies on these ionic systems using dynamic simulation methods¹⁷ indicate that both the *trans-transoidal* and the *cis- transoidal* conformations possess similar energetics and are, therefore equally probable. The energy minimized conformers for ten repeat units are shown in Figure 8. The trans-cissoidal and the ciscissoidal conformers are substantially higher in energy. It is also evident from the study that the counter ions in these charged polyacetylenes interact strongly with adjacent pyridinium rings, thereby preventing backbone :wisting. This is reflected by a large electrostatic energy contribution that is responsible for the stability of such conjugated conformers in these systems. This unique feature also distinguishes them from other substituted, uncharged polyacetylenes that do not possess such extended conjugation

of the backbone. Our modelling study also predicts that the removal of counterions in these polymers (i.e. the uncharged state) would result in a greater backbone twist due to steric factors introduced by the substituents in the absence of the ionic interactions, leading to reduced conjugation.

A literature survey shows that the conjugation number '*n*' for a polyene may be estimated from the uv-visible spectra of the conjugated segment present in the polymer. This method has been applied by Dai and White¹⁸ who have correlated the absorption maxima of conjugated segments obtained by dehydrohalogenation of iodine doped 1,4-polyisoprene with the conjugation number '*n*'. Plotting '*n*' versus the square of the wavelength for each absorption, they obtained a straight line indicating that the Lewis-Calvin equation, $\lambda^2_{max} = kn$, is obeyed fairly well.¹⁹ This relation, which has been verified for low molecular polyenes (n= 3-10)²⁰, was shown by these authors to be valid for '*n*' ranging from 4 to 11. The conjugated sequences calculated from a single value of k (k was calculated to be 17,532) gave a good correlation of expected and found values.¹⁸ The study also concluded that it was possible to predict the distribution of conjugated sequences insulated from one another from their characteristic absorption maxima for '*n*' values of up to 11.

In an attempt to verify the applicability of the Lewis-Calvin equation for these ionic polyacetylenes, the uv-vis absorption maxima of their conjugated backbones were correlated with the conjugation number n. Figure 9 shows a plot of λ_{max}^2 (C=C) versus n obtained for polyenes with n = 3 - 10 that were obtained in the previous study.²⁰ Extrapolation from this plot for polymer VIIIp using its λ_{max} (C=C) absorption maximum gave a conjugation length n of 16. If the proposed reaction mechanism for the polymerization that involves a nucleophilic addition to the acetylenic bond in the propagation step occurs in a trans fashion (which is commonly observed for additions across carbon-carbon multiple bonds), the product polymer would possess a fully extended, *trans-transoidal* conformation through the entire length of the chain. Such a

conformation is also predicted to be a highly stable one for these systems by our modelling studies.¹⁷ If this is true, the conjugation number n for these polymers should be equal to their degree of polymerization, DP. The validity of this assumption was verified for polymer VIIIp by comparing its calculated molecular weight (assuming n = DP) from the uv-vis spectra with the experimentally determined number average molecular weight (M_n) for the deprotonated form obtained by vapor pressure osmometry (Figure 10a). The agreement between the calculated and experimental values within the error of the experiment (\pm 5%) is particularly good. This suggests that these ionic systems do possess a fully extended, trans-transoidal conformation through almost the entire length of the polymer chain. The equation was also found to be applicable for poly(2-n-octadecylpyridiniumacetylenebromide) which was obtained by the quaternization method by making a similar comparison with its molecular weight obtained by direct determination using VPO (Fig. 10b). The degree of polymerization, DP of 12 and the molecular weight (M) of 1240 calculated from the end group analysis for deprotonated VIIIp (based on its analytical data and IR spectrum) is also in excellent agreement experimentally determined one (DP = 13, M_n = 1310 by VPO). This is shown in Table 2.

The validity of the Lewis-Calvin relation for the ionic polyacetylenes implies that the properties of the conjugated polymer backbones in these systems resemble those of the corresponding dimethyl- polyenes for which the equation was deduced. Furthermore, it is evident from this study that the range of *n* over which it is applicable may be extended to an *n* value of up to 16. Obviously, the expression cannot be expected to be valid for high values of *n*, since it is known that bond alternation effects in long polyenes cause the band gap [and λ_{max} (C=C)] to asymtotically approach a constant value. Based on a band gap energy of 2.4eV for fully *trans*-polyacetylene, the λ_{max} (C=C) saturates at ~625nm.²¹ A recent study of the non- linear optical (NLO) susceptibility of third harmonic generation (THG) of *trans*-polyacetylene indicates a finite

conjugation length comprising of <120 carbon atoms (n - 60).²² Hence the plot of λ_{max} (or λ_{max}^2) versus n results in marked deviation from linearity at high values of n, eventually saturating around $\lambda_{max} - 625$ nm, as is evident from Figure 9.

Since the molecular weight calculated for polymer VIIIp based on its extrapolated n value agrees well with the experimentally determined one, any deviation from linearity for up to n = 16 should be minimal. The most significant point that emerges from this correlation for the present systems is that the conjugation number n is close to the degree of polymerization, DP, within the limits of n for which it is valid. It provides an effective method for estimating their molecular mass from the uv-vis absorption maxima of their conjugated backbones, which is particularly helpful in view of their charged nature and strongly colored solutions which make determination of M by standard methods difficult. It must be emphasized, however, that this method may not be applicable for higher values of n due to the nonlinearity of the curve in Figure 9.

In summary, a novel method for the synthesis of substituted, extensively conjugated ionic polyacetylenes has been described. This synthetic route complements the previously described methods for generating such structures by activation of the acetylenic bond in ethynylpyridines via quaternization with alkyl halides and sulfonates, and by complex formation with bromine. A comparison of the uv-vis spectral properties of the polymers obtained by the various routes indicates that the present method involving protonation of the pyridine nitrogen results in greater conjugation in the polymer backbone (also indicating a higher degree of polymerization, DP) and a narrower distribution of molecular masses. We attribute this to the activation step (via protonation) being relatively fast compared to those via quaternization by alkyl halides or methanesulfonates and by complex formation with bromine.

All three methods are superior for polymerization of ethynylpyridine (and its derivatives) compared to polymers obtained by use of coordination catalysts. The polymerization of 2-ethynylpyridine with WCl₆ and MoCl₅ were found to yield low

molecular weight products ($\eta_{inh} = 0.04-0.1 dL/g$) and considerably lower backbone conjugation.²³ The absorption maxima for the unsaturated backbone for these compounds exhibited only broad shoulders around 350 nm, which is indicative of a greater backbone twist in these systems due to the pyridine ring substituents. This phenomenon is characteristic of most substituted polyacetylenes synthesized by using coordination catalysts²⁴, wherein a *trans-transoidal* conformation is precluded by steric factors. The extraordinary structural features of the ionic polyacetylenes obtained by us may be largely attributed to the mechanism of polymerization and the charged nature of the side groups in the resulting polymers. Confirmation of these structural characteristics by solution NMR was complicated by considerable line broadening effects arising from aggregation of the highly charged polymer. Solid state NMR using CPMAS methods on both the protonated and deprotonated polymers are being investigated to substantiate the proposed structures arrived at from spectral data.

The family of ionic polyacetylenes provide, what we believe, the first reported examples of substituted analogs of fully *trans*-polyacetylene. The preparative methods outlined by us for these ionic systems provides a novel approach for the synthesis of all-*trans* substituted ionic polyacetylenes which has, to our knowledge, not been achieved. These results may be particularly important in synthetic polyene chemistry since it offers the possibility of obtaining polyacetylenes possessing the stability and processability that is characteristic of substitution, combined with electrical properties of unsubstituted polyacetylene due to their extended conjugation. The significant increase in electrical conductivity of these ionic polyenes on introduction of dopants is indicative of this. ^{4,25,26} Furthermore, their highly charged nature confers on them the unique properties of polyelectrolytes that may have potential for applications. The inherent limitations in these reactions that preclude the formation of high molar masses is, however, an important factor that must be overcome in order for these materials to be suited for such

applications. Modification of the synthetic approach is, therefore, necessary in order to circumvent this limitation. This aspect is currently under investigation.

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- Figure 1. Activated 2-ethynylpyridine monomers obtained by (a) Menschutkin reaction and (b) complexation reaction with bromine
- Figure 2.
- Figure 3. Infrared spectra of (a) 2-ethynylpyridine and (b) polymer VIIIp
- Figure 4. UV-visible spectra of (a) monomer and polymer VIIIp in the (b) deprotonated and (c) protonated forms
- Figure 5. Infrared spectra of polymer VIIIp in the (a) protonated and (b) deprotonated forms
- Scheme 1.
- Figure 6. Fluorescence emission spectra of VIIIp. (a) 6.5×10^{-3} (b) 1.3×10^{-4} g/L and (c) deprotonated form (6.5×10^{-3} g/L)
- Figure 7. X-ray diffractograms of VIIIp in the (a) protonated and b) deprotonated forms
- Figure 8. Energy minimized conformers of poly(4-methylpyridiniumacetylenebromide) in the (a) *trans-transoidal* and (b) *cis-transoidal* conformations (from Ref. 16)
- **Figure 9.** Plot of λ_{max}^2 (C=C) absorption maxima versus conjugation number *n* for polyenes. (Δ) from Ref 17 (\blacktriangle) this work
- Figure 10. Plot of VPO data for (a) deprotonated VIIIp and (b) poly(2-n-octadecylpyridiniumacetylenebromide)

	Mol. formula		С	н	N	CI
Prior to	[C ₁₄ H ₁₁ CIN] _n	Calc. %	70.88	4.61	11.81	12.70
deprotonation		Found %	70.25	4.82	11.80	12.75
Treated with	[C ₁₄ H ₅ (0.1)CIN] _n ⁺	Calc. %	79.27	4.82	13.20	2.71
KOH		Found %	75.20	4.67	12.33	3.01

Table 1. Elemental composition of poly(2-pyridiniumhydrochloride-2-pyridyl acetylene) copolymer before and after depronation by base

[†] Assuming 10% residual chloride ions

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Table 2.	Calculated and	experimentally	determined	conjugation	numbers	n and DP
	for ionic polya	cetylenes				

Polymer	λ _{max} (C=C) (nm)	n, DP (calc.)	n, DP (det.)	
	530	16 ^a 12 ^b	13 ^c	
H n CieH37 Br	370	8 ^a	6 ^c	

^a From Lewis-Calvin equation $\lambda_{max}^2 = kn$, (Fig. 9)

b From end group analysis, assuming n = DP

c From VPO measurement

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Absorbance

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+

Py := unquaternized monomer

a

b

a

FIGURE &

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n

FIGURE 10