# **Nobel Lecture: ''Synthetic metals'': A novel role for organic polymers**\*

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## **INTRODUCTION**

An organic polymer that possesses the electrical, electronic, magnetic, and optical properties of a metal while retaining the mechanical properties, processibility, etc. commonly associated with a conventional polymer, is termed an *intrinsically conducting polymer* (ICP) more commonly known as a ''synthetic metal.'' Its properties are intrinsic to a ''doped'' form of the polymer. This class of polymer is completely different from ''conducting polymers,'' which are merely a physical mixture of a nonconductive polymer with a conducting material such as a metal or carbon powder distributed throughout the material.

## **THE CONCEPT OF DOPING**

Conjugated organic polymers are either electrical insulators or semiconductors. Those that can have their conductivity increased by several orders of magnitude from the semiconductor regime are generally referred to as *electronic polymers* and have become of very great scientific and technological importance since 1990 because of their use in light-emitting diodes (Nalwa, 1997; Skotheim *et al.*, 1998). *trans*- $(CH)$ <sub>x</sub> and the emeraldine base form of polyaniline are used in Fig. 1 to illustrate the increases in electrical conductivity of many orders of magnitude which can be obtained by doping. The conductivity attainable by an electronic polymer has very recently been increased an infinite number of times by the discovery of superconductivity in regioregular poly(3-hexylthiophene) (Schon *et al.*, 2001). Although this phenomenon was present only in a very thin layer of the polymer in a field-effect transistor (FET) configuration at a very low temperature  $({\sim}2 \text{ K})$ , it represents an historical quantum leap—superconductivity in an organic polymer!

Prior to the discovery of the novel protonic acid doping of polyaniline, during which the number of electrons associated with the polymer chain remain unchanged (MacDiarmid and Epstein, 1989, and references therein), the doping of all conducting polymers had previously been accomplished by redox doping. This involves the partial addition (reduction) or removal (oxidation) of electrons to or from the pi system of the polymer backbone (MacDiarmid and Heeger, 1979, and references therein; Skotheim, 1986; Kanatzidis, 1990).

The concept of doping is the unique, central, underlying, and unifying theme which distinguishes conducting polymers from all other types of polymers (C. K. Chiang *et al.*, 1977, 1978). During the doping process, an organic polymer, either an insulator or a semiconductor having a small conductivity, typically in the range  $10^{-10}$ – $10^{-5}$  S/cm, is converted to a polymer which is in

Conductivity increases with increased doping



FIG. 1. Conductivity of electronic polymers.

<sup>\*</sup>The 2000 Nobel Prize in Chemistry was shared by Alan J. Heeger, Alan G. MacDiarmid, and Hideki Shirakawa. This lecture is the text of Professor MacDiarmid's address on the occasion of the award.

the "metallic" conducting regime  $({\sim}1 - 10^4 \text{ S/cm})$ . The controlled addition of known, usually small  $(\leq 10\%)$ nonstoichiometric quantities of chemical species results in *dramatic* changes in the electronic, electrical, magnetic, optical, and structural properties of the polymer. Doping is reversible to produce the original polymer with little or no degradation of the polymer backbone. Both doping and undoping processes, involving dopant counterions which stabilize the doped state, may be carried out chemically or electrochemically (Kanatzidis, 1990). Transitory doping by methods that introduce no dopant ions are also known (Ziemelis *et al.*, 1991).

By controllably adjusting the doping level, one can easily obtain a conductivity anywhere between that of the nondoped (insulating or semiconducting) and that of the fully doped (highly conducting) form of the polymer. Conducting blends of a (doped) conducting polymer with a conventional polymer (insulator), whose conductivity can be adjusted by varying the relative proportions of each polymer, can be made (Kulkarni *et al.*, 1991). This permits the optimization of the best properties of each type of polymer.

Since the initial discovery in 1977, that polyacetylene  $(CH)_x$ , now commonly known as the prototype conducting polymer, could be *p* or *n* doped either chemically or electrochemically to the metallic state (C. K. Chiang *et al.*, 1977, 1978; Nigrey *et al.*, 1979; Maclnnes *et al.*, 1981), the development of the field of conducting polymers has continued to accelerate at an unexpectedly rapid rate, and a variety of other conducting polymers and their derivatives have been discovered (Skotheim, 1986; Kanatzidis, 1990). This rapid growth rate has been stimulated by the field's fundamental synthetic novelty and importance to a cross-disciplinary section of investigators—chemists, electrochemists, biochemists, experimental and theoretical physicists, and electronic and electrical engineers—and to important technological emerging applications of these materials.

In the doped state, the backbone of a conducting polymer consists of a delocalized pi system. In the undoped state, the polymer may have a conjugated backbone, such as in *trans*- $(CH)$ <sub>x</sub> which is retained in a modified form after doping, or it may have a nonconjugated backbone, as in polyaniline (leuco-emeraldine base form), which becomes truly conjugated only after *p* doping, or a nonconjugated structure, as in the emeraldine base form of polyaniline, which becomes conjugated only after protonic acid doping.

## **REDOX DOPING**

All conducting polymers (and most of their derivatives), e.g., poly-(para-phenylene),



poly(phenylenevinylene),

$$
\{\text{supp}\}_{x\in\mathbb{R}^n}\}
$$

polypyrrole,

polythiophene,

polyfuran,

$$
\mathcal{L}^{\mathcal{A}}
$$

$$
\{\langle\!\!\langle\downarrow\rangle\!\!\rangle\}_{\chi}\;,
$$

poly(heteroaromatic vinylenes),

$$
+\langle\overline{\text{R}}\rangle_{\text{G=}\text{G}}+\langle\overline{\text{R}}\rangle_{\text{H}}
$$

(where  $Y=NH$ , NR, S, and O); polyaniline,

etc., undergo either *p* and/or *n* redox doping by chemical and/or electrochemical processes during which the number of electrons associated with the polymer backbone changes (Skotheim, 1986; Kanatzidis, 1990). Selected examples of the different types of doping are presented below.

Chemical and electrochemical <sup>p</sup> doping. *p* doping, i.e., partial oxidation of the  $\pi$  backbone of an organic polymer, was first discovered by treating *trans*-(CH)<sub>x</sub> with an oxidizing agent such as iodine (C. K. Chiang *et al.*, 1977, 1978), viz.,

$$
trans\text{-}[CH]_{x} + 1.5xyI_{2} \to [CH^{+y}(I_{3})_{y}]_{x} \quad (y \le 0.07).
$$

This process was accompanied by an increase in conductivity from  $\sim 10^{-5}$  to  $\sim 10^{3}$  S/cm. If the polymer is stretch oriented fivefold to sixfold before doping, conductivities parallel to the direction of stretching up to  $\sim$ 10<sup>5</sup> S/cm can be obtained (Skotheim, 1986; Kanatzidis, 1990).

Approximately 85% of the positive charge is delocalized over 15 CH units (depicted below, for simplicity, over only five units) to give a positive soliton, viz.,



*p* doping can also be accomplished by electrochemical anodic oxidation by immersing a *trans*- $\text{CH}$ <sub>x</sub> film in, e.g., a solution of  $LiClO<sub>4</sub>$  dissolved in propylene carbonate and attaching it to the positive terminal of a dc power source, the negative terminal being attached to an electrode also immersed in the solution (Nigrey *et al.*, 1979), viz.,

trans-[CH]<sub>x</sub>+(xy)(ClO<sub>4</sub>)<sup>-</sup>  
→[(CH<sup>+y</sup>(ClO<sub>4</sub>)<sub>y</sub><sup>-</sup>]<sub>x</sub>+(xy)
$$
e
$$
<sup>-</sup> (y≤0.1).

Chemical and electrochemical <sup>n</sup> doping. *n* doping, i.e., partial reduction of the backbone pi system of an organic polymer, was also discovered using *trans*-(CH)<sub>x</sub> by treating it with a reducing agent such as liquid sodium amalgam or preferably sodium naphthalide (C. K. Chiang *et al.*, 1977, 1978), viz.,

$$
trans-[CH]x+(xy)Na++(Nphth)-
$$
  
→[Na<sub>y</sub><sup>+</sup>(CH)<sup>-y</sup>]<sub>x</sub>+Nphth (y≤0.1).

The antibonding  $\pi^*$  system is partially populated by this process, which is accompanied by an increase in conductivity of  $\sim 10^3$  S/cm.

*n* doping can also be carried out by electrochemical cathodic reduction (MacInnes *et al.*, 1981) by immersing a *trans*- $(CH)$ <sub>x</sub> film in, for example, a solution of LiClO, dissolved in tetrahydrofuran, and attaching it to the negative terminal of a dc power source, the positive terminal being attached to an electrode also immersed in the solution, viz.,

$$
trans\text{-}[CH]_x + (xy)\text{Li}^+ + (xy)e^-
$$
  
\n
$$
\rightarrow [\text{Li}_y^+(\text{CH})^{-y}]_x \ (y \le 0.1).
$$

In all chemical and electrochemical *p*- and *n*-doping processes discovered for  $(CH)_x$  and for the analogous processes in other conducting polymers, counter ''dopant'' ions are introduced which stabilize the charge on the polymer backbone. In each case, spectroscopic signatures, e.g., those of solitons, polarons, bipolarons, etc., are obtained characteristic of the given charged polymer. However, the concept of doping phenomena extends considerably beyond that given above to ''doping'' processes in which no counter dopant ion is involved, i.e., to doping processes in which transitory ''doped'' species are produced, which have similar spectroscopic signatures to polymers containing dopant ions. Such types of doping can provide information not obtainable by chemical or electrochemical doping. Examples of these types of redox doping, which can be termed *photodoping* and *charge-injection doping*, are given below.

## **DOPING INVOLVING NO DOPANT IONS**

*Photodoping.* When *trans*- $(CH)_x$ , for example, is exposed to radiation of energy greater than its band gap, electrons are promoted across the gap and the polymer undergoes photodoping. Under appropriate experimental conditions, spectroscopic signatures characteristic of, for example, solitons can be observed (Heeger *et al.*, 1988, and references therein), viz.,



The positive and negative solitons are here illustrated diagrammatically for simplicity as residing only on one CH unit; they are actually delocalized over  $\sim$ 15 CH units. They disappear rapidly due to recombination of electrons and holes when irradiation is discontinued. If a potential is applied during irradiation, then the electrons and holes separate and photoconductivity is observed.

Charge-injection doping. Charge-injection doping is most conveniently carried out using a metal/insulator/ semiconductor (MIS) configuration involving a metal and a conducting polymer separated by a thin layer of a high-dielectric-strength insulator. It was this approach which resulted in the observance of superconductivity in a polythiophene derivative, as described previously. Application of an appropriate potential across the structure

can give rise, for example, to a surface charge layer, the *accumulation layer*, which has been extensively investigated for conducting polymers (Burroughes *et al.*, 1988, 1990; Ziemelis *et al.*, 1991). The resulting charges in the polymer, e.g.,  $(CH)_x$  or poly(3-hexylthiophene), are present without any associated dopant ion. The spectroscopic properties of the charged species so formed can therefore be examined in the absence of dopant ion. Using this approach, spectroscopic studies of  $(CH)_x$  show the signatures characteristic of solitons and the mid-gap absorption band observed in the chemically and electrochemically doped polymer. However, the Coulombic interaction between charge on the chain and dopant ion is a very strong interaction and one that can totally alter the energetics of the system.

**NONREDOX DOPING**



This type of doping differs from redox doping described above in that the number of electrons associated with the polymer backbone does not change during the doping process. The energy levels are rearranged during doping. The emeraldine base form of polyaniline was the first example of the doping of an organic polymer to a highly conducting regime by a process of this type to produce an environmentally stable polysemiquinone radical cation. This was accomplished by treating emeraldine base with aqueous protonic acids and was accompanied by a nine-to-ten order of magnitude increase in conductivity (up to  $\sim 0.3$  S/cm) to produce the protonated emeraldine base (J. C. Chiang and MacDiarmid, 1986; MacDiarmid *et al.*, 1987a; MacDiarmid and Epstein, 1989, and references therein). Protonic acid doping has subsequently been extended to systems such as poly(heteroaromatic vinylenes) (Han and Elsenbaumer, 1989).

## **THE POLYANILINES**

The polyanilines refer to a very important class of electronic/conducting polymers. They can be considered as being derived from a polymer, the base form of which has the generalized composition



and which consists of alternating reduced



and oxidized



repeat units (C. K. Chiang *et al.*, 1977, 1978; J. C. Chiang and MacDiarmid, 1986; MacDiarmid *et al.*, 1987a). The *average* oxidation state  $(1-y)$  can be varied continuously from zero to give the completely reduced polymer



to 0.5 to give the ''half-oxidized'' polymer



to one to give the completely oxidized polymer



The terms "leuco-emeraldine," "emeraldine," and "pernigraniline" refer to the different oxidation states of the polymer where  $(1-y)=0$ , 0.5, and 1, respectively, either in the base form, e.g., emeraldine base, or in the protonated salt form, e.g., emeraldine hydrochloride (C. K. Chiang *et al.*, 1977, 1978; J. C. Chiang and MacDiarmid, 1986; MacDiarmid *et al.*, 1987a). In principle, the imine nitrogen atoms can be protonated in whole or in part to give the corresponding salts, the degree of protonation of the polymeric base depending on its oxidation state and on the *p*H of the aqueous acid. Complete protonation of the imine nitrogen atoms in emeraldine base by aqueous HCl, for example, results in the formation of a delocalized polysemiquinone radical cation (C. K. Chiang *et al.*, 1977, 1978; MacDiarmid *et al.*, 1987a, 1987b) and is accompanied by an increase in conductivity of  $\sim$  10<sup>10</sup>.

The partly protonated emeraldine hydrochloride salt can be synthesized easily by either the chemical or the electrochemical oxidative polymerization of aniline (J. K. Chiang and MacDiarmid, 1986; MacDiarmid *et al.*, 1987a; MacDiarmid and Epstein, 1989). It can be deprotonated by aqueous ammonium hydroxide to give emeraldine base powder (a semiconductor).

## **ALLOWED OXIDATION STATES**

As can be seen from the generalized formula of polyaniline base,



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the polymer could, in principle, exist in a continuum of oxidation states ranging from the completely reduced material in the leuco-emeraldine oxidation state, (1  $(y - y) = 0$ , to the completely oxidized material in the pernigraniline oxidation state,  $(1-y)=1$ . However, we have shown (MacDiarmid and Epstein, 1989) that at least in N-methyl-2-pyrrolidinone (NMP) solution in the range  $(1-y)=0$  to  $(1-y)=0.5$  (emeraldine oxidation state) only two chromophores are present, characteristic of  $(1-y)=0$  and  $(1-y)=0.5$  species and that all intermediate oxidation states consist, at the molecular level, only of *mixtures* of the chromophores characteristic of these two states.

Since most of the properties of polyaniline of interest are concerned with the solid state, we have carried out a series of studies in the solid state which show that the same phenomenon occurs in the  $(1-y)=0$  to  $(1-y)$  $=0.5$  oxidation state range and in the  $(1-y)=0.5$  to  $(1-y)=1$  oxidation state range. Within each of these ranges all intermediate oxidation states consist, at the molecular level, only of mixtures of the chromophores characteristic of the two states defining the beginning and end of each range (Lu *et al.*, 1986; Sun *et al.*, 1990).

## **DOPING**

Polyaniline holds a special position amongst conducting polymers in that its most highly conducting doped form can be reached by two completely different process—protonic acid doping and oxidative doping. Protonic acid doping of emeraldine base units with, for example, 1*M* aqueous HCl results in complete protonation of the imine nitrogen atoms to give the fully protonated emeraldine hydrochloride salt (J. K. Chiang and MacDiarmid, 1986; MacDiarmid *et al.*, 1987a).

As shown in Fig. 2, protonation is accompanied by a 9–10 order of magnitude increase in conductivity reaching a maximum in  $\sim 1$ *M* aqueous HCl.

The same doped polymer can be obtained by chemical oxidation (*p* doping) of leuco-emeraldine base (MacDiarmid and Epstein, 1989). This actually involves the oxidation of the  $\sigma/\pi$  system rather than just the  $\pi$  system of the polymer as is usually the case in *p*-type doping. Its reaction with a solution of chlorine in carbon tetrachloride proceeds to give emeraldine hydrochloride:





FIG. 2. Conductivity of emeraldine base as a function of *p*H of HCl dopant solution at it undergoes protonic acid doping:  $\bullet$ and  $\blacksquare$  represent two independent series of experiments (J. C. Chiang and MacDiarmid, 1986; MacDiarmid *et al.*, 1987a).

#### **NANOELECTRONICS**

The basic purpose of this research is to blend the now well-established field of electronic/conducting polymers with the new, emerging field of nanoscience, by electrostatic fabrication (''electrospinning'') to produce "nanoelectronics"—electronic junctions and devices significantly smaller than the diameter of a human hair  $(-50000 \text{ nm})$ . It is commonly accepted that a nanomaterial is defined as one consisting of a substance or structure which exhibits at least one dimension of less than 100 nm  $(0.1 \mu m)$  ("Nanotechnology ...," 1999).

Our objectives were (i) to develop a method by which nanofibers (diameter  $\leq 100 \text{ nm}$ ) of organic polymers could be controllably and reproducibly fabricated such that in one given preparation, all fibers would have a diameter  $<100 \text{ nm}$  *and* (ii) to reproducibly and controllably fabricate, for the first time, nanofibers of electronic polymers (in their semiconducting and metallic regimes) *and/or* their blends in conventional organic polymers for the purpose of ascertaining their applicability in the fabrication of nanoelectronic devices.

We have made substantial progress in achieving these objectives by using a relatively little known, simple, convenient, and inexpensive "electrospinning" method.<sup>1</sup> We have previously reported (Norris *et al.*, 2000; MacDiarmid *et al.*, 2001) fabrication of the first conducting polymer fibers (diameter  $\sim$ 950–2100 nm) of polyaniline doped with *d,l* camphorsulfonic acid (PAn•HCSA) as a blend in polyethylene oxide (PEO). We were surprised to find that an electronic polymer, such as polyaniline, which might have been expected to be more susceptible



FIG. 3. 50 wt % nanofiber blend of PAn•HCSA fabricated from 2 wt % PAn•HCSA and 2 wt % PEO from chloroform solution at 25 000 V (anode/cathode separation, 25 cm). Scale bar: 100 000 nm.

to degradation than most conventional organic polymers, survived, without observable chemical or physical change, following the 25 000 V electrospinning fabrication process in air at room temperature.

Electrospinning. The electrospinning technique involves a simple, rapid, inexpensive, electrostatic, nonmechanical method in which a polymer solution in a variety of different possible common solvents, including water, is placed in a hypodermic syringe or in a glass pipette, at a fixed distance (5–30 cm) from a metal cathode (Doshi and Reneker, 1999; Gibson *et al.*, 1999). The positive (anode) terminal of a variable high-voltage transformer is attached to the metal tip of the hypodermic syringe or to a wire inserted into the polymer solution in the glass pipette, the negative terminal being attached to the metal cathode. The tip of the syringe can be placed vertically over the cathode or at any other convenient angle to it. When the voltage applied between the anode and cathode reaches a critical value,  $\sim$ 14 000 V at a  $\sim$ 20-cm separation, the charge overcomes the surface tension of the deformed drop of the polymer solution on the tip of the syringe and a jet is produced. Since the polymer molecules all bear the same (positive) charge, they repel each other while traveling in air during a few milliseconds from the anode to cathode and become separated (Reneker *et al.*, 2000). At the same time, evaporation of the solvent molecules occurs rapidly. Evaporation of solvent is also enhanced because the similarly charged (positive) solvent molecules repel each other. Under appropriate conditions, dry, meters-long fibers accumulate on the surface of the cathode, resulting in a nonwoven mesh of nano- to micron-diameter fibers, depending on experimental parameters (Fig. 3).

Nanofiber fabrication. Since the submicron fibers (500– 1600 nm) obtained in our initial work (Norris *et al.*, 2000; MacDiarmid, Jones, *et al.*, 2001) were not classifiable as true ''nanofibers,'' our immediate objective was to break the ''nanotechnology barrier'' and to consistently and reproducibly fabricate true nanofibers

<sup>&</sup>lt;sup>1</sup>For background on electrospinning, see Formhals, 1934; Iijima, 1991; Reneker and Chun, 1996; Doshi and Reneker, 1999; Gibson *et al.*, 1999; Norris *et al.*, 2000; Reneker *et al.*, 2000; MacDiarmid, Jones, *et al.*, 2001.



FIG. 4. Electrospun fibers of polystyrene (see text). Scale bar: 1000 nm. The extended length of the fibers is clearly visible.

(diameter  $\leq 100$  nm) of an organic polymer. This was accomplished (see Fig. 4) using an  $\delta$  wt % solution of polystyrene (Mw 212 400) in tetrahydrofuran at a potential of 20 000 V between the anode and cathode, which were separated by 30 cm. The fibers were collected as a mat on an aluminum target and were found to have the following diameter characteristics: average, 43.1 nm; maximum, 55.0 nm; minimum, 26.9 nm. Other studies involving polystyrene gave fibers whose diameters were consistently  $<$ 100 nm; average, 30.5 nm; maximum, 44.8 nm; minimum, 16.0 nm. It might also be noted that the above 16-nm fiber is only  $\sim$ 30 polystyrene molecules wide. It is also of interest to note that a 16-nm fiber such as the one mentioned above lies well within the  $\sim$ 4–30nm-diameter range of multiwalled carbon nanotubes (Iijima, 1991).

Electronic polymer fibers. By using a previously observed method for producing polyaniline fibers (Reneker and Chun, 1996), we have prepared highly conducting sulfuric acid-doped polyaniline fibers (diameters: average, 139 nm; maximum, 275 nm; minimum, 96 nm) by placing a  $\sim$ 20 wt % solution of polyaniline in 98% sulfuric acid in a glass pipette with the tip  $\sim$ 3 cm above the surface of a copper cathode immersed in pure water at 5000 V potential difference. The fibers collect in or on the surface of the water. The conductivity of a single fiber was  $\sim$ 0.1 S/cm, as expected, since partial fiber de-doping oc-



FIG. 6. Polystryrene fibers collected on a bent copper wire (magnification  $33\times$ ) and subsequently coated with a thin layer of polypyrrole by *in situ* deposition from aqueous solution. Scale bar: 1 mm.

curred in the water cathode. The diameter and length of the fibers appear (Fig. 5) to be sensitive to the nature of the polyaniline used. No great difficulty is foreseen in producing fibers <100 nm diameter.

It is relatively easy to prepare conducting blends of PAn•HCSA in a variety of different conventional polymers such as polyethylene oxide, polystyrene, polyacrylonitrile, etc. For example,  $\sim 20$  wt % blends of PAn•HCSA in polystyrene (Mw 114 200) are obtained by electrospinning a chloroform solution; fiber diameter characteristics: average, 85.8 nm; maximum, 100.0 nm; minimum 72.0 nm. These fibers are sufficiently electrically conductive that their scanning electron microscopies may be recorded without the necessity of applying a gold coating.

Separate, individual nanofibers can be collected and examined if so desired. An appropriate substrate—glass slide, silicon wafer, or loop of copper wire, etc.—is held between the anode and cathode at a position close to the cathode for a few seconds to collect individual fibers (see Fig. 6).

Current/voltage (*I*/*V*) curves are given in Fig. 7 for a single 419-nm-diameter fiber (fiber 1) and for a  $\sim 600$ nm-diameter fiber (fiber 2) of a blend of 50 wt %



FIG. 5. 100% polyaniline fiber with an average diameter of 139 nm.



FIG. 7. Current/voltage curves of 50 wt % PAn•HCSA/PEO blend nanofiber.



FIG. 8. Conducting polypyrrole coated polyacrylonitrile nanofibers. Scale bar: 1000 nm.

PAn•HCSA and polyethylene oxide collected on a silicon wafer coated with a thin layer of  $SiO<sub>2</sub>$ . Two gold electrodes separated by  $60.3 \mu m$  are deposited on the fiber after its deposition on the substrate.

Nanofibers as substrates. The large surface-to-volume ratio offered by nanofibers makes them excellent, potentially useful substrates for the fabrication of coaxial nanofibers consisting of superimposed layers of different materials. Catalysts and electronically active materials can be deposited on them by chemical, electrochemical, solvent, chemical vapor, or other means, for use in nanoelectronic junctions and devices.

We have found, for example, that polyacrylonitrile nanofibers can be easily and evenly coated with a 20– 25-nm layer of conducting polypyrrole (Fig. 8) by immersion in an aqueous solution of polymerizing polypyrrole (Gregory *et al.*, 1989; Huang *et al.*, 1997). Analogously, we have found that electroless deposition of metals can also be performed. Polyacrylonitrile fibers, for example, can be evenly coated with gold by electroless deposition (Sullivan and Kohl, 1995).

Carbon nanofibers. As previously reported (Chun *et al.*, 1999; Hohnholz and MacDiarmid, 2001), polyacrylonitrile fibers may be thermally converted to carbon nanofibers with some shrinkage. We have similarly converted polyacrylonitrile nanofibers to carbon nanofibers.

In summary, electronic polymers have been used for the past 20 years to produce rectifying diodes by Schottky and  $p/n$  junctions, transistors, light-emitting devices, photovoltaic cells, rechargeable batteries, etc. (Nalwa, 1997). Now, the ability to fabricate nanofibers of electronic polymers that are only a few molecules thick suggests the emergence of a field of nanoelectronics whereby the electronic properties of such nanofibers can be exploited for technological purposes.

## **LINE PATTERNING OF CONDUCTING POLYMERS**

One of the exciting challenges of the first part of this century will be the development of low-cost disposable plastic/paper electronic devices (Okusaki and Osada, 1993; Dodabalapur *et al.*, 1998, and references therein;



FIG. 9. 625-pixel polymer dispersed liquid crystal (PDLC) display.

Drury *et al.*, 1998, and references therein). Conventional inorganic conductors, such as metals, and semiconductors, such as silicon, commonly require multiple etching and lithographic steps in fabricating them for use in electronic devices. The number of processing steps and chemical etching steps involved limit the minimum price and therefore their applicability in disposable electronics. On the other hand, conducting polymers combine many advantages of plastics, e.g., flexibility and processing from solution, with the additional advantage of conductivity either in the metallic or semiconducting regimes; however, the lack of simple methods to obtain inexpensive conductive polymer shapes/patterns limit many applications. We here describe a novel, simple, and cheap method for preparing patterns of conducting polymers by a process which we term ''line patterning.''

Line patterning uses the different responses of a substrate and insulating lines which have been printed on it by a conventional copying or printing process towards a fluid (or vapor) to which they are both simultaneously exposed. The substrate and printed lines react differently or at different rates with the fluid (or vapor) to which they have been exposed. This results in a nonuniform deposition on the substrate as compared to the printed lines. If the fluid contains a conducting polymer, which remains as a film after evaporation of the solvent, a pattern of conducting polymer results. A pattern is first designed on a computer and is then printed on, for example, an overhead transparency using a standard, nonmodified office laser printer.

The printed (insulating) lines can be easily removed, if necessary, in a few seconds by ultrasonic treatment in toluene, dissolving the printed lines and leaving a clean pattern of deposited material on the substrate whose shape was originally defined by the now nonexistent printed lines. Line patterning has the following advantages: no photolithography is involved; no printing of



FIG. 10. A simple electronic circuit (''pushbutton'' switch).



FIG. 11. "Pushbutton" switch.

conducting polymer is involved; it uses only, for example, a standard office laser printer, which is not modified in any way; commercially available flexible, transparent plastic or paper substrates can be used; solutions of commercially available conducting or nonconducting polymers can be used from which the polymers may be deposited on substrates; it is inexpensive; rapid development of customized patterns (within hours) from a computer-designed pattern to product is routine.

We have exploited, for example, the observation that a commercial dispersion of poly-3,4-ethylenedioxythiophene  $(PEDOT)^2$  wets a commercial plastic overhead transparency, but not the lines printed on it by a standard office laser printer. A coating of PEDOT can be applied by a roller and, after evaporation of the solvent, the printed lines can be easily and cleanly removed by sonication, leaving only the conducting polymer on the transparency.

Two electrodes were prepared in this way, each containing 25 lines/inch. A drop of a standard commercial polymer dispersed liquid crystal (PDLC) display mixture<sup>3</sup> containing an optical adhesive and  $15-\mu m$ spacer spheres was placed on the center of each electrode. The second electrode was placed on top at an angle of 90 $^{\circ}$  to the first. This resulted in a 25 $\times$ 25, i.e., 625 pixels/sq. inch matrix (Fig. 9). Exposure to UV light for a few minutes resulted in polymerization of the mixture to bind the two electrodes together and to produce a free-standing working PDLC display device. When an electrode pattern of 100 lines/inch was used a working 10 000 pixel/sq. inch display was produced.

We have devised a novel way of separating conducting polymer circuits from each other by making use of the height ( $\sim$ 4–5  $\mu$ m) of the printed toner lines, using a standard office printer, above the substrate, e.g., on an overhead transparency. This is illustrated (Figs. 10 and 11) by a ''pushbutton'' switch to open and close a simple electrical circuit. A combination of two patterned transparencies where the two adjacent conductive areas are electrically separated from each other by two  $\sim$ 4–5- $\mu$ m



• Connection to power supply

FIG. 12. Three-dimensional connections.

nonconductive printed toner lines is obtained by placing the printed lines on top of each other as shown in Figs. 11 and 12. Depression of the areas labeled ''PRESS'' causes the upper transparency to bend. This electrically connects the conducting PEDOT surfaces. When released, the transparency film returns back to its original position, thus breaking the electrical circuit.

The two-dimensional conducting polymer circuits may be readily converted to three-dimensional circuits by two different methods, as shown in Fig. 12, simply by (i) stapling two two-dimensional circuits together using a common office desk stapler, so that the metal stapler joins together electrically the conducting polymer areas on two different substrates *or* (ii) making a pinhole through the sheets, as shown, before applying the PEDOT solution. Some of the solution enters the pinhole and joins together electrically the conducting surfaces on the two different circuits.

We have recently observed a curious field effect which thin films of PEDOT exhibit when exposed to a positive gate potential in an FET configuration as shown in Fig. 13. A source/drain electrode and a gate electrode are prepared by line patterning and are covered by a thin layer of PEDOT as described above. A drop of the optical adhesive containing spacer spheres, described above, is placed on the source/drain electrode upon which the gate electrode is then placed at 90°. The two electrodes are manually squeezed together and the optical adhesive is polymerized by exposure to UV light, as was done for the PDLC display described previously. Several thousand of these interconnected transistor-type devices could be readily fabricated per square inch by the line-patterning process, if it were considered desirable. The free-standing, flexible device shown in Figs. 13 and 14 is produced.

The device exhibits the same general reversible features commonly associated with a field-effect transistor (FET), as shown in Fig. 15.



FET-type device

FIG. 13. Field-effect doped ''PEDOT.''

<sup>&</sup>lt;sup>2</sup>"Baytron P," made by Bayer Corporation.

<sup>&</sup>lt;sup>3</sup>Made by Merck Corporation, Germany, "Li crystal E7" and ''NOA-65 optical adhesive'' (Norland Products, New Hampshire, USA).



FIG. 14. FET-type device.

The doped "metallic" PEDOT film  $(\sigma \sim 2 \text{ S/cm at }$ room temperature) would not be expected to show a change in conductivity by exposure to a field of this type under the configuration used. We believe this effect presents an entirely new method for ascertaining the nature of highly doped ''metallic'' conducting polymers. It has been postulated that a doped conducting polymer consists of metallic "islands" surrounded by low-conducting "beaches," as shown in Fig. 16. We postulate that in the effect we have observed only the low-frequency conducting ''beaches'' and not the metallic ''islands'' respond to the applied field. The application of an electric field changes the conductivity of the ''beaches'' and hence the extent of electrical percolation in the source/ drain PEDOT electrode between the metallic ''islands,'' hence changing the bulk conductivity of the material. The response time for our device is much slower than for a conventional field-effect transistor. We therefore conjecture that the chief changes in conductivity are probably caused by slow diffusion of the dopant anions under influence of the applied field. On removal of the field the system reverts to its original state.



FIG. 15. Field-effect transistor characteristics.

Preliminary studies suggest that the effect is also present in polyaniline; if so, it may possibly be found in many other conducting polymers and would therefore represent a general phenomenon characteristic of all conducting polymers, at least within certain ranges of doping.

#### **SUMMARY**

- Polyacetylene,  $(CH)_x$ , the simplest organic polymer, can be reversibly doped to the metallic regime by partial oxidation or reduction either chemically or electrochemically.
- Polyaniline can be doped to the metallic regime by a simple acid/base protonation.
- A large number of electronic conductive polymers are now known.
- A variety of technological applications of electronic conductive polymers, present and projected, are apparent.

#### **Percolation Field Effect in Doped Conducting Polymers**

Metallic "islands" separated by "beaches" of non- or lowly-conducting (semiconducting) polymer\*



- The field changes:
	- conductivity of the semiconducting "beaches" but not the metallic "islands".
	- ... and hence changes the extent of electrical percolation between the metallic "islands".
	- ... and therefore changes the bulk conductivity of the material.

FIG. 16. Percolation field effect in doped conducting polymers.

## **ACKNOWLEDGMENTS**

This Nobel Prize has world-wide implications since it shows the ever-increasing importance of interdisciplinary research—in this case collaborative research between a polymer chemist, Hideki Shirakawa, Alan Heeger, a physicist, and myself, an organometallic chemist. Each of us had the task of learning the specialized scientific language of the other in order to collectively focus on one specific scientific challenge, an example of the whole being greater than the sum of its parts.

The prize is also recognition of the good fortune that Alan, Hideki, and I had in having each other as such excellent colleagues and also in having such creative colleagues in each of our respective individual research groups—the work stemming from a research group cannot be better than the persons carrying it out. It is a recognition of them and their work and also the work of countless others world wide during the past 24 years who put the "flesh on the skeleton" of work carried out by us in the 1970s at Penn. If it were not for them there would be no prize today in the field.

Research in an experimental science (and also in many other fields) cannot be accomplished without financial support for stipends, apparatus, supplies, and the like. A funding organization and project officers within such an organization have tremendous control over the future of science and technology in any given country. In this respect Dr. Kenneth J. Wynne, my contracting officer at the U.S. Office of Naval Research for many years, before his recent retirement, had the scientific intuition and foresight to fund our first work on conducting polymers—the first funding of work of this type anywhere in the world. He funded it because of its scientific interest. The fact that it now has great technological potential was not a consideration at that time.

"Of what use is a beautiful poem?" It gives intellectual stimulation and enjoyment. Similarly with research. If it has some practical use, that is merely ''icing on the cake!''

## **Early Studies**

*Polyacetylene*, (CH)*<sup>x</sup>* : Alan J. Heeger (*formerly* Physics Department, Univ. of Penn), Hideki Shirakawa (Tsukuba University) and many undergraduate, graduate students and postdoctoral fellows. *Financial Support:* Principally, U.S. Office of Naval Research;<sup>4</sup> University of Pennsylvania Materials Science Laboratory.

*Polyaniline*: Arthur J. Epstein (Physics Department, Ohio State University) and many undergraduate, graduate students and postdoctoral fellows. *Financial Support*: Principally, U.S. Office of Naval Research;<sup>4</sup> University of Pennsylvania Materials Science Laboratory.

#### **Recent Studies**

*Nanofibers (''Electrospinning'')*: I. D. Norris, J. Gao, F. K. Ko, W. E. Jones, Jr., A. T. Johnson, Jr. *Financial Support*: U.S. Office of Naval Research;<sup>4</sup> Army Research Office–Multidisciplinary University Research Initiative.

*Line Patterning*: D. Hohnholz, H. Okuzaki. *Financial Support*: Subcontract, Kent Displays, Inc. (ONR-SBIR Program); U.S. Office of Naval Research;<sup>4</sup> Fellowship from Ministry of Education, Science, Culture and Sports, Japan.

## **REFERENCES**

- Burroughes, J. H., D. D. C. Bradley, A. R. Brown, R. N. Marks, K. MacKay, R. H. Friend, P. L. Burns, and A. B. Holmes, 1990, Nature (London) **347**, 539.
- Burroughes, J. H., C. A. Jones, and R. H. Friend, 1988, Nature (London) **335**, 137.
- Chiang, C. K., M. A. Druy, S. C. Gau, A. J. Heeger, E. J. Louis, and A. G. MacDiarmid, 1978, J. Am. Chem. Soc. **100**, 1013.
- Chiang, C. K., C. R. Fincher, Jr., Y. W. Park, A. J. Heeger, H. Shirakawa, E. J. Louis, and A. G. MacDiarmid, 1977, Phys. Rev. Lett. **39**, 1098.
- Chiang, J. C., and A. G. MacDiarmid, 1986, Synth. Met. **13**, 193.
- Chun, I., D. H. Reneker, H. Fong, X. Fang, J. Deitzel, N. B. Tan, and K. Kearns, 1999, Adv. Mater. **31**, 36.
- Dodabalapur, A., Z. Bao, A. Makhija, J. G. Laquindanum, V. R. Raju, Y. Feng, H. E. Katz, and J. Rogers, 1998, Appl. Phys. Lett. **73**, 142.
- Doshi, J., and D. H. Reneker, 1995, J. Electrost. **35**, 151.
- Drury, C. J., C. M. J. Mutsaers, C. M. Hart, M. Matters, and D. M. de Leeuw, 1998, Appl. Phys. Lett. **73**, 108.
- Formhals, A., 1934, U.S. Patent No. 1,975,504.
- Gibson, P. W., H. L. Schreuder-Gibson, and D. Riven, 1999, AIChE J. **45**, 190.
- Gregory, R. V., W. C. Kimbrell, and H. H. Kuhn, 1989, Synth. Met. **28**, C823.
- Han, C. C., and R. L. Elsenbaumer, 1989, Synth. Met. **30**, 123.
- Heeger, A. J., S. Kivelson, J. R. Schrieffer, and W.-P. Su, 1988, Rev. Mod. Phys. **60**, 781.
- Hohnholz, D., and A. G. MacDiarmid, 2001, Synth. Met. (in press).
- Huang, Z., P.-C. Wang, A. G. MacDiarmid, Y. Xia, and G. M. Whitesides, 1997, Langmuir **13**, 6480.
- Iijima, S., 1991, Nature (London) **354**, 56.
- Kanatzidis, M. G., 1990, Chem. Eng. News 3 December, **68**, 36. Kulkarni, V. G., W. R. Mathew, J. C. Campbell, C. J. Dinkins, and P. J. Durbin, 1991, in *Proceedings of 49th ANTEC Conference*, Montreal, Canada, 5–9 May 1991 (Society of Plastic Engineers and Plastic Engineering), p. 663.
- Lu, F. L., F. Wudl, M. Nowak, and A. J. Heeger, 1986, J. Am. Chem. Soc. **108**, 8311.
- MacDiarmid, A. G., J.-C. Chiang, A. F. Richter, and A. J. Epstein, 1987a, Synth. Met. **18**, 285.
- MacDiarmid, A. G., J.-C. Chiang, A. F. Richter, N. L. D. Somasiri, and A. J. Epstein, 1987b, in *Conducting Polymers*, edited by L. Alcacér (Reidel, Dordrecht), p. 105.
- MacDiarmid, A. G., and A. J. Epstein, 1989, Faraday Discuss. Chem. Soc. **88**, 317.

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- MacDiarmid, A. G., and A. J. Epstein, 1990, in *Science and Applications of Conducting Polymers*, edited by W. R. Salaneck, D. T. Clark, and E. J. Samuelsen (Hilger, Bristol, UK), p. 117.
- MacDiarmid, A. G., and A. J. Heeger, 1979, Synth. Met. **1**, 101.
- MacDiarmid, A. G., W. E. Jones, Jr., I. D. Norris, J. Gao, A. T. Johnson, Jr., N. J. Pinto, J. Hone, B. Han, F. K. Ko, H. Okuzaki, and M. Llagune, 2001, Synth. Met. (in press).
- Maclnnes, D., Jr., M. A. Druy, P. J. Nigrey, D. P. Nairns, A. G. MacDiarmid, and A. J. Heeger, 1981, J. Chem. Soc. Chem. Commun. **1981**, 317.
- Nalwa, H. S., 1997, Ed., *Handbook of Organic Conductive Materials and Polymers* (Wiley, New York).
- ''Nanotechnology—a revolution in the making—vision for R&D in the next decade," March 10, 1999, Interagency Working Group on Nanoscience, Engineering, and Technology.
- Nigrey, P. J., A. G. MacDiarmid, and A. J. Heeger, 1979, J. Chem. Soc. Chem. Commun. **1979**, 594.
- Norris, I. D., M. M. Shaker, F. K. Ko, and A. G. MacDiarmid, 2000, Synth. Met. **114**, 109.
- Okusaki, H., and Y. Osada, 1993, J. Intell. Mater. Syst. Struct. **4**, 50.
- Reneker, D. H., and I. Chun, 1996, Nanotechnology **7**, 216.
- Reneker, D. H., A. L. Yarin, H. Fong, and S. Koombhongse, 2000, J. Appl. Phys. **87**, 4531.
- Schon, J. H., A. Dodabalapur, Z. Bao, C. Kloc, O. Schenker, and B. Batlogg, 2001, Nature (London) **410**, 189.
- Shacklette, L. W., N. F. Colaneri, V. G. Kulkarni, and B. Wessling, 1991, in *Proceedings 49th ANTEC Conference*, Montreal, Canada, 5–9 May (Society of Plastic Engineers and Plastic Engineering), p. 665.
- Skotheim, T. A., R. L. Elsenbaumer, and J. F. Reynolds, 1998, Eds., *Handbook of Conducting Polymers*, 2nd Ed. (Dekker, New York).
- Sullivan, A. M., and P. A. Kohl, 1995, J. Electrochem. Soc. **142**, 2250.
- Sun, Y., A. G. MacDiarmid, and A. J. Epstein, 1990, J. Chem. Soc. Chem. Commun. **1990**, 529.
- Ziemelis, K. E., A. T. Hussain, D. D. C. Bradley, R. H. Friend, J. Rilhe, and G. Wegner, 1991, Phys. Rev. Lett. **66**, 2231.