Light harvesting by dye linked conducting polymers

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Light Harvesting by Dye Linked Conducting Polymers

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Abstract (max. 2000 char.):
The fact that the fossil fuel is finite and that the detrimental long-term effects of letting CO2 into our atmosphere exist, have created an enormous interest in developing new, cheap, renewable and less polluting energy resources. One of the most obvious abundant sources of energy in the solar system is the sun. Unfortunately the well developed silicon solar cells are very costly to produce. In an attempt to produce cheap and flexible solar cells, plastic solar cells have received a lot of attention in the last decades. There are still a lot of parameters to optimize if the plastic solar cell shall be able to compete with the silicon solar cells. One of the parameters is to ensure a high degree of charge carrier separation. Charge carrier separation can only happen at heterojunctions, which cover for example the interfaces between the polymers and the electrodes or the interface between an n-conductor and a p-conductor. The facts that the charge carrier separation only happens at the heterojunctions limits the thickness of the active layer in solar cells and thereby the effectiveness of the solar cells.

In this project the charge carrier separation is attempted optimized by making plastic solar cells with a molecular heterojunction. The molecular heterojunction has been obtained by synthesizing a three domain super molecular assembly termed NPN. NPN consists of two poly[1-(2,5-dioctyltolanyl)ethynylene] chains (N-domains) coupled to the [10,20-bis(3,5-bis(tert-butylphenyl)-5,15-dibromoporphinato]zinc(II) (P-domain).

It is shown that the N domains in NPN work as effective light harvesting antennas for the P domain and effectively transfer electrically generated excitons in the N domain to the P domain. Unfortunately the P domain does not separate the charge carriers but instead works as a charge carrier trap. This results in a performance of solar cells made of NPN that is much lower than the performance of solar cells made of pure poly[1-(2,5-dioctyltolanyl)-ethynylene], Nn. On the other hand light emitting diodes, LEDs, made of Nn and NPN works very well. The LEDs made of Nn emits greenish blue light while LEDs made of NPN emits light in the near-infrared region.

During the synthesis of Nn and NPN it was found that remnants of the palladium catalysts caused problems in the control of the polymers and further made the resistance in the solar cells and LEDs so low that they did not work. A large effort has been made during the project to develop a method to remove remnants of metal catalysts from organic compounds and in particular polymers so that functional solar cells and LEDs could be made. It was succeeded to find a very effective method to remove remnants of metal catalysts from organic compounds by the discovery of the fact that azothioformamides are capable of dissolving metal nanoparticles by forming electron transfer complexes. Even metal wires of some metals can be dissolved by the azothioformamides within a reasonable time range.

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This thesis presents the main results of my work carried out at The Danish Polymer Centre, Risø National Laboratory and Department of Chemistry, Technical University of Denmark (DTU) in the period from May 2003 to June 2006. The work has been carried out under supervision of Senior Scientist Frederik C. Krebs (Risø) and Associated Professor Pernille Harris (DTU) and was financed by STVF. The thesis is submitted to The Faculty of Chemistry and Biotechnology, Technical University of Denmark, in order to obtain the academic degree of Doctor of Philosophy, Ph.D.

The thesis is divided into three main parts. Part one (chapter 1-3) is introductory chapters, which deals with a short introduction to the different aspects in making solar cells of plastic and gives the status of the research in plastic solar cells. Furthermore the objectives of the Ph.D. project are given and discussed (chapter 1). Chapter 2 and 3 in part one are two theory chapters. The first theory chapter deals with a short introduction to how both inorganic and organic solar cells work (chapter 2), while the second theory chapter describe the Sonogashira cross coupling reaction (chapter 3), which is the main reaction type used in this thesis.

Part two describes the major part of my work, and deals with the synthesis of the monomer unit (chapter 4), problems during synthesis of monodisperse oligomers (chapter 5), the synthesis of the polymers (chapter 6), problems with remnants of the palladium catalysts in the polymers and the development of a method to remove palladium from organic samples (chapter 7), attempts to crystallize mono disperse oligomers (chapter 8), the characterization of the polymers and light emitting devices and solar cells made from the synthesized polymers (chapter 9) and a summery of the main results (chapter 10).

Part three describes my contribution to a project concerning an unusually stable tellurium compound showing pyroelectric properties. The work was carried out in collaboration with people from Copenhagen University.

At the end of the thesis two appendixes are presented. Appendix A describes the experimental work that has not been published and Appendix B contains copies of published work, which I have contributed to. The work carried out by me has altogether contributed to six papers, one patent, one proceeding and one poster.

There are lots of people I would like to thank for their help and interest in my work during the project.

In particular I would like to thank Frederik C. Krebs for his supervision, drive and forever and ever showing optimism, and interest in my work. It has been a true pleasure to work with Frederik and his always high optimism and enthusiasm have been very catching. I would also like to thank Mikkel Jørgensen for his action as a kind of assistant supervisor always ready to answer my questions, when Frederik has been impossible to reach because of his many activities. I am deeply grateful for that.
Special thanks to Ole Hagemann, Jan Alstrup, Holger Spanggaard, Klaus Bechgaard, Kion Norman, Lotte Nielsen, Ole Kristoffersen, Birgit Jensen, Lizette Bruun and Eva Bundgaard. Ole is the most skilled technician in organic chemistry I have ever met, and I have received a lot of benefit from his great knowledge concerning practical organic chemistry. Furthermore he has been a god friend and the fact that our sons have been in the same day nursery has given us a lot of things to talk about. Jan is like Ole a very skilled technician and has helped me producing devices and has always been ready to give guided tours to the production line for plastic solar cells, when I have visits from different High schools in Denmark or Sweden. Holger has been a great laboratory mate and have taught me how to use the fluorescence spectrophotometer. The great collaboration with Holger has lead to two papers, one proceeding and one poster. Klaus is my former head of department and is now full professor at Copenhagen University. Furthermore he has been supervisor for both Frederik and Mikkel, and he has an impressive knowledge of both inorganic and organic chemistry. I would like to thank Klaus for the excellent collaboration in order to remove remnants of palladium and other metal catalyst for our compounds, which have lead to three papers and one patent. Kion is the third senior scientist in the solar cell group at the department, and I will like to thank him for his always good mood, which always spreads happiness around him to the rest of us. His different funny emails have lead to many funny conversations. Thanks for that Kion. Lotte is like Ole and Jan a skilled technician and I will like to thank her for always having time to answer question concerning the department's analytic SEC equipments and to teaching me how to use the MALDI-TOF spectrometer. Ole K. and Birgit is the infrastructure of the department, without them nothing would be possible. They have made it a true pleasure to be organic synthetic chemist in this department by taking care of all the dirty dishes, the purchasing of chemicals, helping with mounting of new equipment etc. Lizette has been a good friend, and we have enjoyed many hard working hours at the spinning lectures in the fitness center, but all her cakes and her ice cream system have resulted in that my shape stays the same. Eva shall have a lot of thanks for always being joyful and being able to coop with all the guy talk at lunch.

I would also thank the rest of the solar cell group for good collaboration and a nice solidarity. Further I would like to thank the rest of the people in the polymer centre for being good colleagues.

From DTU I would like to thank the technicians Lise Lotte Berring and Astrid Schøneberg for helping me to choose the right crystals and to use the SMART system and Pernille Harris for her supervision, and her optimism with regard to growing crystals of the monodisperse oligomers. Further I will like to thank Pernille for teaching me in the use of the SMART and how to use the X-ray software, always having time for me, making sure that the paperwork was done in time. Unfortunately the project did not lead to much structure solving as anticipated in the beginning because I was unable to grow crystals of the monodisperse oligomers, but at least we succeeded in solving the crystal structures of the electron transfer complexes, which lead to one structure paper.

From the University of Roskilde I will like to thank my former supervisor professor Poul Erik Hansen for the many discussions concerning the unusually broad lines in the NMR spectra of the electron transfer complexes and for providing the 600 MHz NMR spectrophotometer to my disposal. Furthermore I will like to thank Bjarke Hansen for the proofreading of this thesis.
Last but not least I will thank my family and particular my wife Bettina for her unusual understanding, when I have been coming home late from work and for her always continuing support.

I therefore dedicate this work to her and my two children Lukas and Laika.
Abstract

The fact that the fossil fuel is finite and that the detrimental long-term effects of letting CO₂ into our atmosphere exist, have created an enormous interest in developing new, cheap, renewable and less polluting energy resources. One of the most obvious abundant sources of energy in the solar system is the sun. Unfortunately the well developed silicon solar cells are very costly to produce. In an attempt to produce cheap and flexible solar cells, plastic solar cells have received a lot of attention in the last decades. There are still a lot of parameters to optimize if the plastic solar cell shall be able to compete with the silicon solar cells. One of the parameters is to ensure a high degree of charge carrier separation. Charge carrier separation can only happen at heterojunctions, which cover for example the interfaces between the polymers and the electrodes or the interface between an n-conductor and a p-conductor. The facts that the charge carrier separation only happens at the heterojunctions limits the thickness of the active layer in solar cells and thereby the effectiveness of the solar cells.

In this project the charge carrier separation is attempted optimized by making plastic solar cells with a molecular heterojunction. The molecular heterojunction has been obtained by synthesizing a three domain super molecular assembly termed NPN. NPN consists of two poly[1-(2,5-dioctyltolanyl)ethynylene] chains (N-domains) coupled to the [10,20-bis(3,5-bis-tert-butylphenyl]-5,15-dibromoporphinato]zinc(II) (P-domain).

！NPN！

The synthesized molecular heterojunction, NPN.

It is shown that the N domains in NPN work as effective light harvesting antennas for the P domain and effectively transfer electrically generated excitons in the N domain to the P domain. Unfortunately the P domain does not separate the charge carriers but instead works as a charge carrier trap. This results in a performance of solar cells made of NPN that is much lower than the performance of solar cells made of pure poly[1-(2,5-dioctyltolanyl)]-ethynylene], Nₙ. On the other hand light emitting diodes, LEDs, made of Nₙ and NPN works very well. The LEDs made of Nₙ emits greenish blue light while LEDs made of NPN emits light in the near-infrared region.

During the synthesis of Nₙ and NPN it was found that remnants of the palladium catalysts caused problems in the control of the polymers and further made the resistance in the solar cells and LEDs so low that they did not work. A large effort has been made during the project to develop a method to remove remnants of metal catalysts from organic compounds and in
particular polymers so that functional solar cells and LEDs could be made. It was succeeded to find a very effective method to remove remnants of metal catalysts from organic compounds by the discovery of the fact that azothioformamides are capable of dissolving metal nanoparticles by forming electron transfer complexes. Even metal wires of some metals can be dissolved by the azothioformamides within a reasonable time range.


I afhandlingen vises det, at N-domænerne i NPN virker som meget effektive lyshøstende antenner for P-domænet og effektivt overfører excitoner fra N-domænerne til P-domænet. Desværre separerer P-domænet ikke ladningsbærererne, menvirker i stedet som en ladningsbærefælde. Dette resulterer i at ydeevnen for solceller lavet af NPN er meget lavere end ydeevnen for solceller lavet af den rene poly[1-(2,5-dioctyltolanyl)ethynylen], Nn. På den anden side virker lysdioder lavet af både NPN og Nn rigtig godt. Lysdioder lavet af Nn udsender blågrønt lys, mens lysdioder lavet af NPN udsender lys i den nær infrarøde region.

Under syntesen af Nn og NPN observerede jeg, at rester af palladiumkatalysatoren gav problemer med at kontrollere polymererne og yderligere blev modstanden i solcellerne og lysdioderne så lav, at de ikke virkede. En stor del af projektet er gået til at udvikle en metode til fjernelse af rester af metalkatalysatoren fra såvel små organiske forbindelser som polymerer. Især har det været vigtigt at få fjernet katalysator rester fra polymererne, så funktionelle solceller og lysdioder har kunnet fremstilles. Den store arbejdssindsats bar frugt, og det lykkedes at finde en meget effektiv metode til fjernelse af rester af metalkatalysatorer.
fra organiske forbindelser ved opdagelsen af azothioformamidernes evne til at opløse metal nanopartikler under dannelse af elektronoverførelses komplekser. Selv metaltråde af visse metaller kan blive opløst af azothioformamiderne inden for en rimelig tidsramme.
## List of Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>DDC</td>
<td>Diethylammonium diethyldithiocarbamate</td>
</tr>
<tr>
<td>$E_F$</td>
<td>Fermi level</td>
</tr>
<tr>
<td>$E_g$</td>
<td>Band gap</td>
</tr>
<tr>
<td>FF</td>
<td>Fill factor</td>
</tr>
<tr>
<td>$I_0$</td>
<td>The dark saturation current</td>
</tr>
<tr>
<td>$I_{dark}$</td>
<td>The dark current</td>
</tr>
<tr>
<td>$I_{max}$</td>
<td>The current at the maximum power point</td>
</tr>
<tr>
<td>IP</td>
<td>Ionization potential</td>
</tr>
<tr>
<td>$I_{ph}$</td>
<td>The photo current</td>
</tr>
<tr>
<td>IPV</td>
<td>Inorganic photovoltaic</td>
</tr>
<tr>
<td>ISC</td>
<td>Inorganic semiconductor</td>
</tr>
<tr>
<td>$I_{SC}$</td>
<td>The short-circuit current</td>
</tr>
<tr>
<td>ITO</td>
<td>Indium tin oxide</td>
</tr>
<tr>
<td>LED</td>
<td>Light emitting diode</td>
</tr>
<tr>
<td>MOPV</td>
<td>Molecular organic photovoltaic</td>
</tr>
<tr>
<td>$M_w$</td>
<td>Molecular weight</td>
</tr>
<tr>
<td>OPV</td>
<td>Organic photovoltaic</td>
</tr>
<tr>
<td>OSC</td>
<td>Organic semiconductor</td>
</tr>
<tr>
<td>PCB14</td>
<td>[6,6]-phenyl C$_{61}$ butyric acid methyl ester</td>
</tr>
<tr>
<td>PD</td>
<td>Polydispersity</td>
</tr>
<tr>
<td>PEDOT:PSS</td>
<td>Poly(ethylene dioxythiophene) doped with polystyrene sulphonic acid</td>
</tr>
<tr>
<td>$P_{in}$</td>
<td>The power input</td>
</tr>
<tr>
<td>POPV</td>
<td>Polymer organic photovoltaic</td>
</tr>
<tr>
<td>$P_{out}$</td>
<td>The power output</td>
</tr>
<tr>
<td>PPE</td>
<td>Poly(phenylene ethynylene)</td>
</tr>
<tr>
<td>PPV</td>
<td>Poly(phenylene vinylene)</td>
</tr>
<tr>
<td>PR-TRMC</td>
<td>Pulse radiolysis time-resolved microwave conductivity</td>
</tr>
<tr>
<td>PV</td>
<td>Photovoltaic</td>
</tr>
<tr>
<td>$R_{serial}$</td>
<td>Serial resistance</td>
</tr>
<tr>
<td>$R_{shunt}$</td>
<td>Shunt resistance</td>
</tr>
<tr>
<td>SEC</td>
<td>Size exclusion chromatography</td>
</tr>
<tr>
<td>TB</td>
<td>Through bond</td>
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### List of Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>TBAF</td>
<td>Tetrabutylammonium fluoride</td>
</tr>
<tr>
<td>TS</td>
<td>Through space</td>
</tr>
<tr>
<td>UPS</td>
<td>Ultra photoelectron spectroscopy</td>
</tr>
<tr>
<td>$V_{\text{max}}$</td>
<td>The voltage at the maximum power point</td>
</tr>
<tr>
<td>$V_{\text{OC}}$</td>
<td>The open-circuit voltage</td>
</tr>
<tr>
<td>$\eta_{\text{eff}}$</td>
<td>The efficiency</td>
</tr>
<tr>
<td>$\Phi$</td>
<td>Workfunction</td>
</tr>
<tr>
<td>$\chi_s$</td>
<td>Electron affinity</td>
</tr>
</tbody>
</table>
The recognition that the supply of fossil fuel is finite and the detrimental long-term effects of letting CO₂ into our atmosphere are the seeds for a great motivation to develop new, cheap, renewable and less polluting energy resources. Obvious alternatives to the fossil fuel today are nuclear power, wind energy, water energy and sun energy. Nuclear power plants deliver very cheap and renewable energy, but unfortunately it is not a potential energy source in many countries for political reasons owed to the fear of security and health risk. The interest in wind energy is growing, and e.g. in Denmark nearly 20 %, of the total annual electricity consumption is covered by windmills.¹ Windmills however are not particularly beautiful, cannot be placed anywhere, and need maintenance. Water energy is cheap and renewable but in general requires rivers, which we do not have in e.g. Denmark. Solar energy is renewable, it is ubiquitous and solar cells can be placed everywhere, even in a satellite, but unfortunately the conventional solar cells on the market are very expensive. Even though the photovoltaic (PV) effect was discovered in 1839 by Edmond Becquerel² and the first crystalline silicon PV device was developed in 1954 at Bell Laboratories³ it has not yet been possible to get the price per watt down, so that it can influence the energy production markets significantly. The reason for the high power prize is that the cells are made of inorganic semiconductors (ISC) such as silicon (Si) and Gallium Arsenide (GaAs), which need very high production temperatures and therefore are expensive to produce. For instance metallic Si is produced by reduction of quartz (SiO₂). The reduction is performed with coke at 4000 ºC and the product is about 98 % pure Si. Further purification is then typically achieved by reaction with hydrogen chloride to form trichlorosilane (SiHCl₃), which is purified by fractional distillation and finally reduced to Si by hydrogen.⁴ More than fifty years of research since the development of the first silicon solar cell has lead to a world PV module production of 560.27 MW in 2002, with a power cost of ~$4/W.⁵ The energy produced by PV therefore counts for less than 0.1 % of the total world energy production.⁶ If the progress of the prizes for the conventional solar cells continue to decrease with the same rate as now, the best estimates of the power cost in 2014 would be 1.4 $/W, which is still too high.⁵

A very promising and interesting alternative to the conventional inorganic solar cells are solar cells based on organic semiconductors (OSC). OSC is a unique class of materials, which already have shown potential applications in several types of optoelectronic devices. Compared with ISC, OSC show some very interesting advantages: They are very cheap in production and can be produced near room temperature, the purification is easy and cheap, the organic photovoltaic (OPV) devices can be made flexible, they are lightweight and can be incorporated in e.g. clothes.⁶,⁷,⁸ It should be noted that the disadvantages of OPV compared to inorganic photovoltaic (IPV) devices are the performance and the lifetime. The power conversion efficiencies for single crystal silicon cells are as high as 24 % and 32.5 % for a multijunction GaAs solar cell, while the best organic solar cells have power conversion efficiencies around 4-5 %.⁷,⁹,¹⁸ Concerning the lifetime the IPV devices are extremely stable and have a lifetime of more than 25 years, while the estimated lifetime of the most stable OPV devices is only a few years.¹⁰ In this context it should be mentioned that the OPV...
devices with longer lifetime are not the same OPV devices with power conversion efficiencies of $\geq 3\%$.

In summary, if either the IPV or the OPV devices shall be able to break through to the energy market and not only be limited to a niche market, they have to fulfil three parameters: Long lifetime, good power conversion efficiency and low cost. Brabec has visualized this by the critical triangle (see Figure 1). As mentioned above and as seen in Figure 1 neither the IPV nor the OPV devices fulfil all three parameters.

![Figure 1. The Brabec critical triangle for photovoltaics. If a Photovoltaic is going to be a success on the energy market, it has to fulfill all three requirements: Lifetime, efficiency and low cost. On the figure the current achievement of IPV and OPV devices is circled. The IPV devices have good power conversion efficiency and a long lifetime, but are very expensive; the OPV devices have very low cost, but moderately low power conversion efficiency and relatively short lifetime.](image)

The basis of this thesis is the OPV devices, which can be divided into two main groups, distinguished by the type of material and fabrication methods: Molecular organic photovoltaics and polymer organic photovoltaics (plastic solar cells). The molecular organic photovoltaics (MOPV) are inspired by the photosynthesis in plants and consist of molecular dyes that are macrocyclic molecules like chlorophyll (Chart 1 show structures of some of the small molecules, which are often used in MOPVs). In MOPV devices the active layer is sublimed under vacuum onto the electrodes. The most efficient solar cells have an active layer, which consists of an organic heterojunction between hole (p-type) and electron (n-type) transporting organic materials. Examples of p-type materials are: Metal-phthalocyanines (MPc), metal-5,10,15,20-tetraphenylporphyrins (MTpp), merocyanine dyes; and n-types are: 3,4,9,10-perylene tetracarboxylic-bis-benzimidazole (PTCBI), metal-5,10,15,20-tetra(4-pyridyl)porphyrin and fullerenes (see Chart 1). The breakthrough of the MOPVs was in 1986, where Tang reported a MOPV device with a power conversion efficiency of 1%. The device was a bilayer, where CuPc was employed as the p-conductor and the n-conductor was PTCBI. Recently, the Princeton group has produced MOPV cells with a power conversion efficiency of 5%. This was achieved by doping CuPc with $C_{60}$ and optimizing the morphology of the heterojunction between $C_{60}$ doped CuPc and PCTB.

The polymer organic photovoltaics (POPV) consist of conjugated polymers sandwiched between two different electrodes. One of the biggest advantages of the POPV devices is that the polymers give the opportunity to prepare devices by solution processing which is very cheap and easy, so it might lead to cheap preparation of large area devices. Solution
processing includes methods like spin coating, screen or inkjet printing and even roll-to-roll processing.\textsuperscript{15}

In Chart 2 examples of different types of conjugated polymer used in POPV devices are shown. Before the POPV devices can become the success many people hope for and before the industry will be interested in producing them, solutions to two common problems are needed. The two problems are the stability and the power conversion efficiencies of the POPV devices.

Regarding the stability or the lifetime of POPV devices, it is not necessary that the devices has the same long lifetime as the IPV devices, because they are much cheaper and can therefore be replaced many times and still be an economically favoured choice compared to the IPV devices. On the other hand before devices for durable applications are interesting, they need a shelf lifetime of several years and an operational lifetime of tens of thousands of hours.\textsuperscript{16}

\textbf{Chart 1.} Examples of some dye molecules used for MOPV devices.
Krebs et al. have recently solved the lifetime problem very elegantly. They have used ester groups as side groups on the polymer backbones. The side groups give the opportunity for solution processing, and can here after be removed by thermal cleaving. In this way they obtain an active layer in the POPV devices, which is very rigid, hard and insoluble. The diffusion of the oxygen and ions from the electrodes through the active layer is slowed down and in that way prevented from destroying the devices. Thus they have obtained devices, which have shelf-lifetimes as well as operational lifetimes of more than 2 years under the exclusion of water and oxygen. Unfortunately the power conversion efficiencies are only 0.167 %.

A big challenge lies in combining this knowledge of lifetime with the knowledge of polymers with good power conversion efficiency. This challenge leads us to the other main problem: The power conversion efficiency.
The power conversion efficiency of the best POPV devices is about 4.4%,\textsuperscript{18} which is still far from the efficiencies of the IPV devices. There are several reasons for the small power conversion efficiencies of the POPV devices. Some obvious reasons are:

- There is not a perfect match between the solar spectrum and the optical absorption bands of the polymers. The maximum spectral flux of the sun (AM1.5) is between 550 and 900 nm (see Figure 2), where the majority of known conjugated polymer absorb below 500 nm. Polymers with low band gap are therefore needed to absorb the red photons.\textsuperscript{5,19,20}

- One of the most fundamental differences between IPV and OPV devices is that in IPV devices light absorption leads directly to the formation of free electron-hole pairs, where the electron and the hole will be driven to the respective electrodes by the built in field.\textsuperscript{21,22} In OPV devices the light absorption leads to formation of excitons, which can be considered as a electrically neutral tightly electrostatically bound electron-hole pair or as a mobile excited state. The binding energy of the excitons is in the order of several tenths of an electron volt,\textsuperscript{7} and a typical value is 0.25 eV.\textsuperscript{23} This means that the built in field generated from the use of electrodes with different workfunctions is insufficient to dissociate the excitons into free charge carriers (electrons and holes),\textsuperscript{7} and the dissociation can therefore only happen at a heterointerface e.g. at the electrodes.

- The low charge carrier mobilities in polymer semiconductors is limiting the thickness of the active layer, because the photo generated excitons have to be transported by diffusion within their lifetime to a heterointerface (e.g. the interface between the electrodes and the active layer) to dissociate into free charge carriers. The diffusion length of an exciton in polymer semiconductors is typically in the order of 10 nm. This means that the exciton can travel 10 nm before it recombines and decays to the ground state.\textsuperscript{5}

- A better understanding of the charge transport across different interfaces, can lead to an optimisation of the cells by reducing e.g. the contact resistance.\textsuperscript{5} Brabec \textit{et al.} have recently shown that the insertion of a thin layer of LiF under e.g. the negative aluminium electrode enhances the efficiency by more than 20 %.\textsuperscript{24}

It can be concluded that lots of areas need to be investigated with an overall performance enhancement of the POPV devices in mind. It is unrealistic to believe that all the mentioned problems can be considered in this thesis. I have therefore decided to limit the topic of this thesis to an optimisation of the power conversion efficiency in POPV devices through a more effective dissociation of the photo generated excitons.

As already mentioned in the list above the only place a photo generated exciton can dissociate in a POPV device consisting of a single conjugated polymer sandwiched between to different electrodes, is at the heterointerface between the electrodes and the polymer. The short diffusion length of the excitons therefore limits the thickness of the polymer layer used to generate free charge carriers and thereby also limits the quantity of photo generated current in the devices. Several attempts have been made to optimize the amount of dissociated excitons, and common for these attempts are the application of doping the polymer with either molecules or polymers having a higher electron affinity and electron transport than the pure polymer. The most successful attempts have employed mixing the polymer with fullerene derivatives (e.g. [6,6]-phenyl C\textsubscript{61} butyric acid methyl ester (PCBM)) and hereby make a bulk heterojunction, by which power conversion efficiencies above 4 % have been achieved.\textsuperscript{18,25,26}

The use of fullerenes results in an ultra fast metastable photoinduced electron transfer through space (TS) from the polymer to the fullerene with a larger electron affinity.\textsuperscript{27,28} As a transfer process through space is involved, the rate constant of the TS electron transfer process depend
on the distance between the donor and the acceptor molecule. To maximize the area of the heterojunction it is required that the PCBM and the polymer are mixed very well. To avoid phase separation or large domain size upon the mixing of the acceptor and the donor molecules, a molecular heterojunction can be designed. In molecular heterojunctions the donor and the acceptor molecules are covalently bound together, which will ensure that the area of the heterojunction is maximized. Nierengarten et al.\textsuperscript{39} Peeters et al.\textsuperscript{30} and van Hal et al.\textsuperscript{31} were among the first who in 1999 and in 2000 investigated charge separation in phenylene vinylene and thiophene oligomers covalently linked to fullerene. They both discovered that the length of the conjugated oligomers determined whether they observed an energy transfer to the fullerene from the oligomers or a charge separation (electron transfer). The critical length of the oligomers was between 3 and 4 units. The longer oligomers favour the electron transfer, while the shorter oligomers favour the energy transfer. Other molecular heterojunctions, where polymers or oligomers have been covalently bound to dyes have been investigated by Krebs et al. and Li et al. Krebs et al. have investigated 4-aza-4-(4′-(poly-2′,5′-dioctyl-4,4′-terphenylene-1-cyanovinylene-2-yl)biphen-4-yl)-8,12-dioxa-4,8,12,12c-tetrahydrodibenzo[cd,mn]pyrenium tetrafluoroborate (JA-assembly)\textsuperscript{32,33} and 10,20,diphenyl-5,15-bis(4-(poly-2′,5′-dioctyl-4,4′-terphenylene-1-cyanovinylene-2-yl)phenyl)porphyrin (JPJ-assembly)\textsuperscript{34} and Li et al. have investigated porphyrins with four monodisperse oligofluorine arms (PF\textsubscript{4} assembly).\textsuperscript{35} Common for the three investigations is the observation of an effective energy transfer from the polymers or the oligomers to the dye molecules. Further Krebs et al. observed a 100-fold increase in magnitude of the short circuit current in PV devices of the JA-assembly compared to the PV devices of the pure J domain (the polyterphenylationcyanovinylene).\textsuperscript{33}

Inspired by the work of Krebs et al.\textsuperscript{1} the aims of the present work are the following:

- To synthesise new molecular heterojunctions consisting of super molecular assemblies, where conjugated polymers are covalently bound to a dye molecule.
- To characterise the structural and electrical properties of the super molecular assemblies and the pure polymers.
- Testing the assemblies and the pure polymers properties as a light harvesting system.
- Testing PV devices made from the synthesised assemblies and polymers.

1.1 The choice of system

To have the best foundation for comparison with results from the literature,\textsuperscript{34,35} it was decided to use porphyrin as the dye. The polymer backbone was chosen to be poly(phenylene ethynylene)s. The reason for the choice of the triple bonds in the polymer backbone was to ensure very stiff polymer chains compared to the PPVs in the study of Krebs et al.\textsuperscript{34} The stiff polymer chain may reduce a potential torsion angle out of plane between the polymer and the dye and a potential torsion angle out of plane between the individual monomer units in the polymer chain (see Figure 3 for illustration of the torsion angles). The reduction of possible distortions of the backbone could open up for an increased conjugation and an increased charge carrier mobility, which hopefully can lead to an increased energy transfer from the polymer chains to the dye molecule.

\textsuperscript{1} The work of Li et al. was not published, when this study was started.
To obtain solubility of the polymer it was decided to incorporate two alkyl groups on the polymer backbone, but only on every second phenyl ring to minimize the distortion effect of the alkyl groups on the backbone. The chosen super molecular assembly is illustrated in Figure 3, where the poly[1-(2,5-dioctyltolanyl)ethynylene] constitute the N-domains and the [10,20-bis(3,5-bis-tert-butylphenyl]-5,15-dibromoporphinato]zinc(II) constitute the P-domain.

Another reason to choose the poly(phenylene ethynylene) as the polymer backbone of the system is that it can be synthesised by the Sonogashira cross-coupling reaction, which in theory gives the possibility to stepwise synthesise monodisperse oligomers. Hence it is possible to investigate the influence of the length of the polymer chains on the properties of the NPN assembly. Furthermore, the combination of the stiff monomer unit and the possibility to synthesise monodisperse oligomers, makes it probable that crystallisation of the oligomers could be achieved allowing for an investigation of the crystal structure of the oligomers.

References

8 http://www.risoe.dk/solarcells/element-ary.htm
The conversion of light into electricity can in principle be divided into the following four steps: Light absorption, generation of an electron hole pair, separation of the electron hole pair into free charge carriers and transport of the free charge carriers to the respective electrodes. In Solar cells (photovoltaic devices), whether it is polymer organic photovoltaics (POPV) or inorganic photovoltaics (IPV), the light absorption takes place in semi conducting materials sandwiched between two electrodes. The absorption of light results in excitation of electrons from the valence bands to the conduction band, but to achieve separation of the electron from the hole an electric field stronger than the columbic forces between the electrons and the holes is needed. It is only at the interface, also called contact junction, between the semiconductors and the electrodes or the interface between two semiconductors with different workfunctions such electrical fields are found. The excitation of electrons and the following charge separation in crystalline inorganic semiconductors is well described by the use of energy band formalism. These processes are in the case of polymeric organic semiconductors not so well described as in the case of the inorganic semiconductors. This is due to both the fact that there has been more research in inorganic semiconductors than organic semiconductors and because the polymeric organic semiconductors miss the three-dimensional crystal lattice. The system of polymeric organic semiconductors is very complex, because of the lack of crystal lattice, which means that the classical energy band formalism cannot be used to describe the solid-state properties satisfactory. In the absence of a better alternative the classical energy formalism is often used as a very rough approximation to give a qualitative description of the fundamental processes in the polymer organic semiconductors. The charge carrier mobilities in the polymeric organic semiconductors are generally several orders of magnitude lower than in the traditional inorganic semiconductors and the light absorption does not lead to generation of free electron hole pairs but instead of excitons as mentioned in the introduction.

In this chapter I will briefly describe how the classical energy band formalisms are used to describe the fundamental processes in POPV. Furthermore, I will describe the device structures and the electrical characteristics of these.

2.1 Inorganic photovoltaic devices

From solid-state physics it is known that all solid materials can be divided roughly into three groups, according to the energy gap size between the two electronic bands: The valence band and the conduction band. The three groups are conductors, semiconductors and insulators. In the case of conductors the two electronic bands are so close that they practically form a continuum. The semiconductors and the insulators are both characterised by an energy gap between the two bands, also called the band gap ($E_g$). The difference between the semiconductors and the insulators is the magnitude of the band gap. Normally it is presumed that materials with a band gap below 2 eV are semiconductors and materials with a band gap above this value are insulators (see Figure 4).
Solid materials can be divided into three groups according to the size of their band gap between the conduction band and valence band. The three groups are conductors (no band gap), semiconductors (band gap lower than 2 eV) and insulators (band gap higher than 2 eV).

The distribution of electrons in the various energy levels is given by the Fermi-Dirac distribution:

\[
f(E) = \frac{1}{1 + \exp \left( \frac{E - E_F}{kT} \right)}
\]

In the Fermi-Dirac distribution \(E_F\) is the Fermi level. From equation (2.1) it can be seen that if \(E=E_F\), then \(f(E)=0.5\), which means the probability that the concerned energy level is occupied is 50%. If \(E<E_F\) the particular energy level is more likely to be occupied than unoccupied and if \(E>E_F\) then the particular energy level is more likely to be unoccupied than occupied. For intrinsic semiconductors with no impurities and at ordinary temperatures the Fermi level lies in the middle of the band gap between the valence and the conduction band (see Figure 5). By doping an intrinsic semiconductor with impurities consisting of e.g. elements from an adjacent group in the periodic table new energy levels are introduced near the band edges. In the case of elements from a higher main group, the impurities have more electrons than can be fitted into the stable structure of the semiconductors. These electrons are therefore more loosely bound and their energy levels (donor levels) are near the bottom of the conduction band. Due to the small energy difference between the energy levels of these electrons and the conduction band, part of these electrons will be excited into the conduction band by the thermal energy. Consequently, the Fermi level in the doped semiconductor will move closer to the conduction band (the Fermi level at low temperature in doped semiconductor will lie approximately in the middle between the energy of the donor levels and energy of the conduction band) (see Figure 5). This type of doped semiconductors is called n-type, because they have excess of negative charge carriers.
In the case of doping a semiconductor with atoms from a lower main group, the impurities have fewer electrons than is needed for the stable structure. Therefore it does not require much energy for electrons from neighbour atoms to occupy these empty positions (acceptor levels) and form covalent bonds. The energy of the acceptor levels is therefore near the valence band edge. At room temperature the acceptor levels will be partly filled by thermal excited electrons from the valence band. Consequently, the Fermi level in this doped semiconductor will move closer to the valence band (the Fermi level at low temperature will approximately be in the middle between energy of the acceptor levels and energy of the valence band) (see Figure 5). This type of doped semiconductor is called p-type, because they have excess of positive charge carriers.

An important parameter for materials is the workfunction, which by definition is the energy difference between the vacuum level and the Fermi level (see Figure 5). The workfunction is denoted $\phi_m$ and $\phi_s$ for the metal and the semiconductors respectively.

### 2.1.1 Schottky junction, ohmic contact and p-n junction

If two materials with different workfunctions are brought into contact the Fermi levels of both materials will coincide at thermal equilibrium. This happens because free carriers in the material with the lower workfunction transfer to the material with the higher workfunction. In this way an electronic double layer is formed at the interface between the two materials, and as a consequence of this, the Fermi level in the material with the lower workfunction will be lowered by an amount equal to the difference between the two workfunctions (relative to the Fermi level in the materials with the high workfunction). If both materials are metals, the electronic double layer, where the charge separation happens, is only few Å thick, due to the large free carrier density (normally $>10^{22}$ per cm$^3$) (see Figure 6).

In the case of semiconductors, where the free carrier density is normally $<10^{18}$ per cm$^3$, free carriers from the bulk area are needed to be transferred to the interface to align the Fermi levels of the two materials. In this case the electronic layer is much thicker (μm-scale), and
this layer is also called the depletion layer or the space charge layer, because it contains net positive charges or negative charges from ionized donors or acceptors.²

**Figure 6.** A band diagram for a metal metal contact.²

If we consider the case where e.g. an n-type semiconductor is brought into ideal contact (in absence of surface states) with a metal with higher workfunction, we will observe that electrons from the semiconductors conduction band are transferred into the metal to line up the Fermi levels. A negative charge is therefore built up at the surface of the metal. In the semiconductor an equal positive charge must be built up, but due to the low free carrier density in the semiconductors compared to the metal, this positive charge is distributed over the entire depletion layer. Consequently a parabolic potential gradient would be built up in the depletion layer and the bands of the semiconductors would be bended as illustrated in Figure 7. This form of energy barrier is called a Schottky barrier.²,⁴

**Figure 7.** A band diagram of the Schottky junction between a metal and a n-type semiconductor ($\phi_n > \phi_s$).²
The magnitude of the band bending in equilibrium, $eV_b^0$, is also called the contact potential and it is given by:

$$eV_b^0 = \phi_m - \phi_s$$  \hspace{1cm} (2.2)

The contact potential is the energy barrier of the flow of electrons from the semiconductor conduction band to the metal. The magnitude of the Schottky barrier for the n-type semiconductors, $\phi_s$, which is the barrier of the flow of electrons from the metal to the semiconductors, is given by:

$$\phi_n = \phi_m - \chi_s$$  \hspace{1cm} (2.3)

where $\chi_s$ is the electron affinity of the semiconductor.

For p-type semiconductors the Schottky barrier, $\phi_p$, is given by:

$$\phi_p = E_g - (\phi_m - \chi_s)$$  \hspace{1cm} (2.4)

In the cases above it was assumed that $\phi_m > \phi_s$ for the n-type semiconductors ($\phi_m < \phi_s$ in the case of the p-type), but in the case where $\phi_m < \phi_s$ for the n-type, electrons will now be transferred from the metal to the semiconductors to line up the Fermi levels. Because of the high free carrier density in the metal and because there is a lot of room for these electrons in the conduction band of the semiconductors, the electronic double layer will be very thin. This layer is so thin that charge carriers easily can tunnel through it from both sides. This type of contact is called an “ohmic contact” (see Figure 8).2,4

Figure 8. A band diagram of the ohmic contact between a metal and a n-type semiconductor ($\phi_m < \phi_s$).2

Focusing on the case where a n-type and a p-type semiconductor is brought into contact, we will observe that electrons from the conduction band in the n-type semiconductor (the semiconductor with the lower workfunction) will diffuse into the valence band of the p-type (the semiconductor with the higher workfunction) until the Fermi levels of the two semiconductors coincide (see Figure 9).
Due to the lower free carrier density in the n-type semiconductor and the low number of unoccupied places in the valence band of the p-type semiconductor a parabolic potential will arise in the depletion layer of both semiconductors.

### 2.1.2 p-n junction solar cells

At thermal equilibrium a built-in potential will be present in the depletion layer as described above and illustrated in Figure 9, and therefore there will be an electrical field in the depletion layer with direction from the n-type to the p-type semiconductor. This means that if an electron from the conduction band of the n-type semiconductor tries to diffuse in to the depletion layer it will be repelled back to the n-type semiconductor by the electrical field. On the other hand a hole in the valence band of the p-type semiconductor will be repelled back to the p-type semiconductor if it tries to diffuse in to the depletion band. At room temperature in dark, electron hole pairs will be generated in both semiconductors, and if they are generated in or near the depletion layer, the electrical field will separate them and the electrons will be accumulated in the n-type semiconductor, while the holes will be accumulated in the p-type semiconductors. This will lead to a net electrical current from the n-type to the p-type semiconductor. The accumulation of electrons and holes in the n-type and p-type semiconductors respectively will lead to a decrease of the band bending. This decrease will allow for electrons from the conduction band of the n-type semiconductors to diffuse into the valence band of the p-type semiconductor, and the opposite is the case for the hole in the valence band of the p-type semiconductor. Hence an electrical current is generated in the opposite direction of the current due to the separation of generated electron hole pairs near the depletion layer. At thermal equilibrium these two currents will outbalance each other. The first type of current will hardly depend on the magnitude of the band bending while the second type of current will depend highly on the magnitude of the band bending. The sum of these two types of current is called the dark current, $I_{\text{dark}}$, and can be expressed by:

$$I_{\text{dark}} = I_0 \exp \left( \frac{e\psi}{kT} \right) - 1$$

In equation (2.5) the first term represents the second type of current while the second term represents the first type of current. $I_0$ is called the dark saturation current and represent the current flowing under high reverse bias. In Figure 10 the $I_{\text{dark}}$ is shown as a function of the
applied bias for a Si solar cell from a garden lamp. It can be seen that p-n junction is showing rectifying behaviour and therefore works like a diode.

Under illumination the generation of electron hole pairs near the depletion layer will increase and give an additional contribution called the photo current, \( I_{ph} \). The total current under illumination is given by the sum of the photo current and the dark current (see Figure 10):

\[
I = I_{ph} + I_{dark} = I_{ph} + I_0 \left[ \exp \left( \frac{eV}{kT} \right) - 1 \right]
\]

Under open-circuit conditions the accumulation of electrons and holes in the n-type and p-type semiconductors respectively will flatten the band bending out so the dark current and the photo current will cancel out. No total net current will be running in the solar cell. The open-circuit photovoltage, \( V_{OC} \), is therefore given by:

\[
V_{OC} = \frac{kT}{e} \ln \left( \frac{I_{ph}}{I_0} + 1 \right)
\]

Under short-circuit conditions the accumulated electrons and holes in the n-type and p-type semiconductors will be transferred through the electrical circuit to the electrodes and no photovoltage will be produced. In this case the band bending is at maximum and therefore the photocurrent will be at maximum. The short-circuit current, \( I_{SC} \), will be equal to the photo current:

\[
I_{SC} = I_{ph}
\]

The diode property of solar cells is described by the fill factor, \( FF \). The \( FF \) is the ratio between the maximum power output from the solar cell (the maximum power point of the \( I/V \) curve in the 4th quadrant) and the maximum theoretical power output if the solar cell was an ideal diode (the product of \( V_{OC} \) and \( I_{SC} \)):

\[
FF = \frac{V_{max} I_{max}}{V_{OC} I_{SC}}
\]

Where \( V_{max} \) and \( I_{max} \) is the voltage and the current respectively at the maximum power point.

One of the most interesting values of a solar cell is the efficiency, which is given as the ratio between the power output, \( P_{out} \), of the cell and the power input, \( P_{in} \), from the sun, \( P_{light} \):

\[
\eta_{eff} = \frac{P_{out}}{P_{in}} = \frac{V_{max} I_{max}}{P_{light}} = \frac{V_{OC} I_{SC} FF}{P_{light}}
\]
2.2 Polymer organic photovoltaics

In contrast to IPV devices the active layer in POPV devices consists of conjugated polymers where the delocalisation of the π-bonding electrons form a “highway” for charge carrier transport along the backbone of the polymer chains. Charge carrier transport also occurs between the different polymer chains by interchain electron transfer. The mobilities are, however, smaller than in the case with inorganic semiconductors, which is caused by the disordering of the polymer chains compared to the crystal lattice observed for the inorganic semiconductors. The π-band, represents the energy levels of the π-bonding and antibonding orbitals denoted π and π* respectively. Those correspond to the valence and conduction band respectively in an inorganic semiconductor. At low temperature the π-band would be completely filled and the π*-band would be empty. Further there would be an energy gap, $E_g$, between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) in the π-band. Conjugated polymers can therefore be considered as semiconductors. The conjugated polymer has, from a synthetic chemistry point of view, the advantage compared to an inorganic semiconductor that the $E_g$ depends on the molecular structure of the polymers. Synthetic chemists have the capability to tune the $E_g$ in new
polymers to a desired level, and thereby determine in which region of the sun light absorption takes places or in the case of light emitting diodes, the colour of the emitted light.\textsuperscript{5} Light absorption in POPV devices, as mentioned in the introduction, does not lead directly to free charge carriers, but instead to neutral excitons (mobile excited states), which to first order are unaffected by electric fields. This means that the generated excitons by diffusion have to be transported to a heterointerface, where a dissociation of the excitons into a free electron in one material and a free hole in the other material can take place. When the excitons are dissociated the free carriers can be transported to their respective electrodes by the built in potential produced by the electrodes with asymmetrical workfunctions. Due to the very low mobility of the excitons in the polymer layer the energy conversion efficiencies for POPV devices made of pure conjugated polymers are very low, typically between $10^{-3}-10^{-1}$ %.\textsuperscript{6,8,9,10}

In order to optimise the POPV devices and to understand their mechanism function, a good model of the devices is useful. The model has to describe how device properties such as the $I_{SC}$, $V_{OC}$, $FF$ and $\eta_{eff}$ depend on the fundamental properties such as the workfunctions, the charge carrier mobilities, the band gap of the polymers, electron affinities etc.

### 2.2.1 The MIM-model

The simplest POPV device consists of a polymer layer sandwiched between two metal electrodes with asymmetrical workfunction. This setup is clearly explained by the MIM-model (Metal-Insulator-Metal) originally developed by Parker for polymer based diodes.\textsuperscript{7} The MIM-model is based on the assumption that the free carrier concentration in the polymers in dark is very low, hence it can be assumed to be an insulator. This has the consequence that the band bending at the metal-polymer interface when applying an electrical field would be very small. The model can therefore be regarded as a rigid band model contra the case of e.g. a Schottky junction (Figure 7). The POPV device can now be viewed in the following four situations: Short circuit condition (zero bias), open circuit condition, under forward bias and under reverse bias.

#### 2.2.1.1 Short-circuit condition

Under short-circuit conditions the Fermi level of the two electrodes will align because of the difference in workfunction, and give rise to a constant electrical field across the polymer layer, which will tilt the polymer bands as illustrated in Figure 11A. Under illumination any generated free charge carriers will experience a driving force from the field such, that electrons will be transported to the electrode with the low workfunction while the holes would be transported to the electrodes with the high workfunction; the devices will exhibit a considerable photocurrent.\textsuperscript{8} In this connection two things have to be pointed out. First, when free charge carriers are generated under illumination, the polymer can no longer be assumed to be an insulator, and there will be some band bending near the metal-polymer interfaces. Second, light absorption in POPV devices, as mentioned, lead to neutral excitons and not to free charge carriers.\textsuperscript{6,10}

#### 2.2.1.2 Open-circuit condition

Under open-circuit condition (also called flat band condition) and illumination electrons and holes will accumulate on the interface at the low and high workfunction electrodes, respectively (see Figure 11B). This accumulation will flatten the band bending so the dark current and the photo current would be equal; this means that no total net current will be
running in the solar cell, as in the case with the IPV devices. The open-circuit photo voltage, $V_{OC}$, is equal to the difference in workfunction of the two electrodes.\(^7\)

**Figure 11.** The MIM-model formulated by Parker\(^7\). A) The short-circuit condition where the Fermi level of the two electrodes aligns and thereby tilts the polymer bands. The built-in potential ensures that the free carrier will be transported to their respective electrodes. B) The open-circuit condition where $V_{OC}$ is determined by the difference in workfunction of the two electrodes. C) Under forward bias electrons and holes can tunnel into the conduction band and the valence band of the polymer respectively, where recombination can lead to electroluminescence (The device is working as a LED). D) Under reverse bias the applied potential tilt the bands of the polymer further and the free carriers are transported to their respective electrodes by the large electrical field (the device works as a photodetector).
2.2.1.3 Forward bias
Under forward bias (the POPV devices work like a diode) the applied electrical field will tilt the polymer bands, and two barriers arise; one caused by the energy difference between the HOMO level for the polymer and the workfunction of the anode (the electrode with the high workfunction) and the other caused by the energy difference between the LUMO level of the polymer and the workfunction of the cathode (the electrode with the low workfunction) (see Figure 11C). When the electrical field increase the barriers will be easier to overcome and eventually the holes from the high workfunction electrode and electrons from the low workfunction electrode will be able to tunnel into the polymer film.\(^7,11\) If the electrons and holes are capable of recombining, electroluminescence can be observed, and a light emitting diode (LED) is made. The \(I/V\) characteristics are determined by the lower of the two barriers, which often is the barrier between the HOMO level of the polymer and the anode, while the device efficiency is determined by the higher barrier, which often is the barrier between the LUMO level of the polymer and the cathode. Changing the anode electrode to e.g. an electrode with a lower workfunction (but not lower than the cathode) will increase the lower barrier height and thereby increase the operating voltage of the diode (the voltage where holes can tunnel into the polymer film), since the polymer bands have to be tilted further before the holes are able to tunnel into the polymer film. Hence the best devices are obtained when the anode workfunction is matched with the HOMO level of the polymer and the anode, while the cathode workfunction is matched with the LUMO level of the polymer. The turn-on voltage of the POPV devices is the voltage required to achieve flat-band condition, which is equal to the band gap of the polymer minus the two barrier offsets.\(^7\)

2.2.1.4 Reverse bias
Under reverse bias the POPV devices exhibit a strong photo response caused by the strong external field, which very efficiently transport the hole to the high workfunction electrode and the electrons to the low workfunction electrode (see Figure 11D). In this way the POPV devices can be used as a photodetector.\(^8\)

2.2.2 Key parameteres
The key parameters to describe the POPV-devices is similar to the IPV devices: \(\eta_{eff}, I_{SC}, V_{OC},\) and \(FF\).

2.2.2.1 The efficiency (\(\eta_{eff}\))
The efficiency, \(\eta_{eff}\), is the most essential parameter for solar cells, which, as in the case of IPV devices, is determined by equation (2.10).

Typically values for single layer POPV devices are between \(10^{-3}\) to \(10^{-1}\) %,\(^8\) while bulk heterojunction POPV devices have shown efficiencies up to \(4.4\) %.\(^{12}\)

2.2.2.2 The open-circuit voltage (\(V_{OC}\))
The open-circuit voltage, \(V_{OC}\), is the maximum voltage the device can produce. The \(V_{OC}\) for POPV devices is higher than for inorganic cells and is typically between 0.4 and 1.5 \(V\).\(^{13}\) In the case of single layer POPV devices that has be described by the MIM-model, the built-in potential can be estimated by the \(V_{OC}\), and therefore the \(V_{OC}\) influences properties such as the
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charge dissociation, charge transport and charge collection in the devices. In the case where the MIM-model is valid, the $V_{OC}$ is estimated as the difference between the workfunction of the two electrodes. In cases where the POPV devices consist of blended polymer and fullerene (bulk heterojunction solar cells) the MIM-model cannot satisfactorily explain the observed $V_{OC}$, which is higher than the difference between the workfunction of the two electrodes. The typical value of $V_{OC}$ for ITO/polymer-fullerene blend/Al devices range from 0.5 V for P3HT and 0.8 V for PPV and up, which is in contrast to the 0.4 V predicted by the MIM-model. The origin of the $V_{OC}$ for the bulk heterojunction solar cells is not fully understood yet, while Brabec et al. have shown that the value of the $V_{OC}$ correlate directly with the acceptor strength of the fullerenes, and depends only very little on the workfunction of the cathode material.

2.2.2.3 The short-circuit current ($I_{SC}$)

The short-circuit current, $I_{SC}$, is the current running in the device under illumination and no applied bias, and therefore it represents the maximum possible current. The magnitude of $I_{SC}$ depends on the efficiency of the charge separation, the mobility and the lifetime of the charge carriers in the active layer and the intensity of the illumination. In order to optimize $I_{SC}$ a fullerene is often blended into the polymer matrix to optimize the interface area, where charge separation can take place. Further it has been shown that the polymer matrix transports the holes and the fullerene molecules transport the electrons. Along the polymer chains the transport of the holes is fast while between the individual chains the transport is nearly 5 times slower. The fullerene molecules transport the electrons by a hopping process between the individual fullerene molecules, and therefore an optimal performance requires that the fullerenes are packed close together. Illumination under AM 1.5 conditions gives an observed magnitude of $I_{SC}$ in the range of 0.2 to 10 mA/cm$^2$.

2.2.2.4 The fill factor ($FF$)

The fill factor, $FF$, which describes the diode properties of the POPV devices, is, like in the case of IPV devices, determined by equation (2.9). $FF$ depends largely on two parameters: The presence of current leakage e.g. due to small holes in the polymer layer and ohmic contacts between the polymer layer and the electrodes. To gain insight in the $FF$ dependence on the two parameters, an equivalent electrical circuit of a POPV device can be drawn. In the equivalent electrical circuit the current leakage is modelled as a small parallel shunt resistance, $R_{Shunt}$ while the ohmic contacts are modelled as a large serial resistance, $R_{Serial}$, which also includes the bulk resistance and the circuit resistance. If the shunt resistance is very low, which means a large current leakage, the $FF$ will be low and the devices will act as an ohmic resistance, which will result in a straight line in the $I/V$ characteristic of the devices. In order to increase the shunt resistance it is very common to spin coat a layer of the hole conductor PEDOT:PSS (Poly(Ethylene DioxyThiophene doped with PolyStyrene Sulphonic acid) on top of the anode electrode. This gives a barrier layer with a conductivity of $\approx 10^{-3}$ S/cm, which increase the shunt resistance and prevents shorts between the two electrodes as a consequence of holes in the polymer layer.
**Figure 12.** The equivalent electrical circuit for a POPV device. $R_{\text{Shunt}}$ is the total shunt in the devices, $R_{\text{Serial}}$ represent the ohmic contacts, the bulk resistivity, the circuit resistivity and $I_{\text{light}}$ represent the photo generated current.

**Figure 13.** The $I/V$ characteristic in dark and under illumination (a 1.5 A.M. sun simulator) of a bulkheterojunction POPV device made from P3HT (Poly-3-HexylThiophene), PCBM fullerene and having the ITO and Al electrodes. In this case the area of the solar cell was 10 cm$^2$, $P_{\text{light}} = 100\text{mW/cm}^2$, $V_{\text{OC}} = 0.4$ V, $I_{\text{SC}} = 5.8$ mA, $V_{\text{max}} = 0.15$ V, $I_{\text{max}} = 2.8$ mA, $FF = 0.18$, $\eta_{\text{eff}} = 0.1$ %. It can be seen that the shunt resistivity is very low, which implied that when a large bias is applied an ohmic behaviour is observed (a straight line in the $I/V$ characteristic).
If the serial resistance is too large both the $I_{SC}$ and $I_{max}$ will decrease, which will lower both $\eta_{eff}$ and $FF$. To reduce the resistivity of the ohmic contacts, it has been shown that placing a thin layer (below 20 Å) of LiF between the polymer layer and the Al electrode (cathode) results in a decrease of the serial resistance by a factor of 3-4 and an increase in the forward current, the $FF$ and the $V_{OC}$. The mechanism of the LiF layer is still not fully understood.\textsuperscript{6,16}

Typical values of the $FF$ for POPV devices lie in the range of 0.4 to 0.6.\textsuperscript{13}

2.3 The device construction in this project

In this project both the LED and solar cells device construction are sandwich constructions (see Figure 14) consisting of a single polymer layer. The devices are large area devices, with an active area of 3 cm$^2$. The devices were prepared by spin-coating chloroform solutions of the polymers onto PEDOT:PSS covered ITO glass slides (the cathode (LEDs) or anode (solar cells) electrode). The ITO (Indium doped tin oxide) electrode is the most frequently used cathode (LEDs)/anode (solar cells) electrode, because it is transparent (the transimission of the ITO covered glass slides is $\approx 90\%$ from wavelength at 325 nm and up)\textsuperscript{6} and have a low sheet resistance (5-15 $\Omega$ square\textsuperscript{-1}). The devices were prepared so the polymer films had peak absorbancies of $\approx$0.6 (LEDs) and $\approx$0.4 (solar cells). Ca/Al was used as the cathode (LEDs) or anode (solar cells) electrode, and was evaporated in a vacuum of $< 1 \times 10^{-5}$ mBar. In the case with the solar cells, I have also tested a set of bilayer devices; where the p-layer was the polymers and the n-layer was C$_60$. The bilayer devices were produced by evaporating (vacuum $< 1 \times 10^{-6}$ mBar) a C$_{60}$ layer on top of the polymer before evaporation of the aluminium electrode. The electrical connections were achieved using silver epoxy glue.

Figures 14. Left: A picture of the POPV devices used in this thesis. Right: A schematic draw of the POPV devices used in this thesis. The active area of these devices is 3 cm$^2$, which is considered large area devices.

References

4 Sze, S. M., Physics of Semiconductor Devices, 1981, chapter 5, John Wiley & Sons, USA.
A important reaction type in the synthesis of the electrical conducting polymers in this project is the coupling reaction between terminal acetylenes and sp²-carbon halides. Since 1975¹,⁶,⁷ it has been well known that a σ-bond between sp²- and sp-centers can be formed by a cross-coupling reaction, where e.g. a free alkyne reacts with an alkenyl or aryl halide in the presence of a catalytic amount of a Pd⁰ or a PdⅡ complex. There are several types of these coupling reactions ² that are named after the discoverers. The main difference in these coupling reactions is the reaction conditions such as the choice of catalyst, solvent and reaction temperature. One common coupling reaction is the Sonogashira cross coupling reaction, which has the advantage that it runs under very mild conditions and with a high conversion yield¹,³ (see Figure 15).

![Figure 15. The Sonogashira cross coupling reaction.](image)

In this section the mechanism of the Sonogashira cross coupling reaction will be described for terminal alkynes with sp²-carbon halides. Further emphasis will be on which by-products you can expect, and which reaction conditions that are commonly used to get the optimum reaction rate and conversion.

### 3.1.1 The reaction mechanism

The Sonogashira cross coupling reaction is very complex, as the suggested reaction mechanism in Figure 16 indicates. The reaction however proceeds under mild conditions in the presence of a palladium catalyst and cuprous iodide as a co-catalyst in amine solvents.¹

The reaction mechanism suggested by Sonogashira⁸ is based on his and co-workers discovery of CuI-catalyzed transmetallation of amines⁵. In the mechanism transmetallation occurs in the two catalytic cycles B' and B, by which the terminal alkyne is coordinated (step ii and iii) to palladium to form the Pd-acetylide complexes (compound 1 and 4 in Figure 16). The coordination occurs in two steps. The first step is an acid-base reaction between the terminal alkyne and the CuI, which forms a copper alkyne. The alkyne is a nucleophilic reagent, and in the next step the alkyne makes a nucleophilic attack at the palladium catalyst (PdCl2(PPh3)2). The driving force for the transmetallation is the difference in the electronegativities of the two metals.
The role of the cycle B’ is to form the catalytically active coordinatively unsaturated 14-electron complex (compound 2 in Figure 16), which is formed by reductive elimination (step iii) of the Pd-acetylide complex (compound 1). The term reductive elimination does not have the same meaning as the term reduction in organic chemistry. The reductive elimination covers the loss of two one-electron ligands of cis configuration from the Pd center by combining them into one single elimination product. It means that both the coordination number and the formal oxidation state of Pd is reduced to form Pd(0). In connection to the formation of compound 2, a by-product in the shape of an equivalent amount of the bis-acetylene compound is formed. The fact that the bis-acetylenes compound forms is supported by Dieck and Heck. They have observed that an increased amount of the acetylene leads to higher yields. The coordinatively unsaturated Pd-complex is the catalyst for cycle A. By oxidative addition (step i) the alkenyl or the aryl halide is coordinated to the Pd-complex (compound 2) to form the complex 3. Through the other transmetallation cycle (B) (nucleophilic attack by the copper alkynide), the halide is exchanged by the acetylene (step iii), and the complex 4 is formed. Under a reductive elimination (step iii) the aryl alkyne or the enyne is cleaved from the Pd complex (compound 4), and the coordinative unsaturated Pd-complex (compound 2) is reformed. Without this reductive elimination step, the reaction would stop and the Pd compound would not have any catalytic effect, but react as a stoichiometric compound as in e.g. the Grignard reactions.

Figure 16. The reaction mechanism Sonogashira suggested for the coupling reactions between terminal acetylenes and sp^2-carbon halides. i) is an oxidative addition step, ii and iii) are transmetallation steps and iii and iiii) are reductive elimination steps.
3.1.2 By-products

The Sonogashira reaction is popular because it generally works quite well, and many authors have described many applications for this reaction type. Unfortunately most of the authors never comment on the fact that it is very hard if at all possible to perform the Sonogashira alkynyl-aryl cross coupling without side reactions. Reported by-products are 1,4-diaurylbutadiynes, 1,4-diaurylbuta-2-en-1-yynes and the cyclic trimerization product 1,3,5-triaurylbenzene. It should not be a surprise that the Sonogashira reaction have by-products such as the cyclic trimer because it has been known for more than 50 years that transition metals catalyze the trimerization of alkynes.

The formation of the three types of by-products is not trivial but it looks like the presence of oxygen and the choice of the catalyst system have a decisive effect on which by-product and the amount in which it is formed. Trumbo and Marvel have thus detected between 5 and 20% of the cyclic trimer 1,3,5-triarylbenezene when they have synthesized several poly(phenylacetylenes) with different catalysts in the presence of oxygen. They found that Pd(OAc)$_2$ gave the highest yield of the cyclic trimer and that the lowest yield was obtained by employing PdCl$_2$ as the catalyst. Varlemann observed all three by-products under the synthesis of tolane from iodobenzene and phenylacetylene by using Pd(OAc)$_2$/PPh$_3$/CuI as the catalyst system and NBu$_4$OH as the solvent. When changing the reaction conditions to Pd(OAc)$_2$/PPh$_3$/CuI/TMEDA and NET$_3$ she only observed the 1,4-diarylbutylnyes in a yield of 4%. It should be pointed out that Trumbo & Marvel have performed their syntheses at low temperature while Varlemann has done the synthesis in a steel autoclave at high temperature.

The formation of the cyclic trimer by-product has been confirmed by Gevorgyan et al. which in 1997 and 2001 have shown that substituted benzenes can be constructed from enynes and activated alkynes in a [4 + 2] and from terminal alkynes in a [2 + 2 + 2] fashion in the presence of a Pd(0) catalyst. Both reactions proceed under normal Sonogashira conditions without an inert atmosphere.

Crisp & Flynn observed in 1993 that even when they used Pd(PPh$_3$)$_4$ as the catalyst in a coupling reaction with less reactive electrophiles such as electron rich arylbromides or triflates, they obtained a by-product in the form of 1,4-diaurylbutadiynes, which according to the reaction mechanism for the Sonogashira reaction (see Figure 16) should only be produced if a Pd(II) catalyst was employed. Later they showed that a catalytic amount of both Cu(I) and Pd(0) along with base and oxygen were all required to achieve homocoupling of terminal alkynes. They have thus produced 1,4-diphenylbutadiyne in 82% yield with a reaction time of 1 day from phenylacetylene with triethylamine as solvent and CuI and Pd(PPh$_3$)$_4$ as the catalytic system in presence of atmospheric air. It is therefore extremely important to keep oxygen away from the reaction.

3.1.3 Reaction conditions

Since the discovery of the Sonogashira cross coupling reaction, a lot of different conditions have been reported. The choice of the right condition depends of the reactivity of the selected halide. The reactivity of the halide is given by the following order: Vinyl iodide ≈ vinyl bromide > aryl iodide > vinyl chloride >> aryl bromide.

In order to choose the right condition several parameters have to be considered. The most obvious ones are: The reaction temperature, the choice of base, the choice of the catalyst, the
ratio between the catalyst and the compounds, the presence or the absence of oxygen and the reaction time.

3.1.3.1 The reaction temperature

The choice of the right reaction temperature depends on the reactivity of the halide and the boiling point of the solvent. Generally, the less reactive halide, the higher temperature is necessary. E.g. if the halide is a vinyl iodide or aryl iodide the reaction easily runs at room temperature or even at lower temperature, whereas it seems that aryl bromides normally have to be heated to approximate 353 K\textsuperscript{7-18}. Hundertmark \textit{et al.}\textsuperscript{19} have, however, recently reported a concept (see the Scheme 1), which works for the aryl bromides at room temperature. The catalyst systems consist of palladium with bulky, electron-rich phosphines as ligands. The investigations of Hundertmark \textit{et al.} show that especially P(t-Bu)\textsubscript{3} is a very efficient ligand for the palladium in the Sonogashira reaction at room temperature.

\begin{center}
\begin{tikzpicture}
\node[above] at (0,0) {\textbf{Scheme 1.} Shows the catalytic system for the Sonogashira cross coupling reaction with aryl bromides at r.t. as suggested by Hundertmark \textit{et al}\textsuperscript{19}.}
\end{tikzpicture}
\end{center}

3.1.3.2 The base

The reaction rate of the coupling reaction depends highly on the choice of base\textsuperscript{6}. Alami \textit{et al.}\textsuperscript{18} have made a thorough investigation on, how the choice of base influences the coupling rate. The investigations are made in both the presence and absence of copper halides (see Scheme 2 and Table 1).

\begin{center}
\begin{tikzpicture}
\node[above] at (0,0) {\textbf{Scheme 2.} The coupling reaction in absence of copper halide.}
\end{tikzpicture}
\end{center}
Table 1. The yields and reaction times as function of the base and the presence and the absence of CuI obtained by Alami et al.\textsuperscript{18}

<table>
<thead>
<tr>
<th>Amine</th>
<th>PK\textsubscript{a}\textsuperscript{20}</th>
<th>Absence of CuI</th>
<th>Presence of CuI (10 %)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Time [h]</td>
<td>Isolated yield [%]</td>
<td>Time [min]</td>
</tr>
<tr>
<td>Et\textsubscript{3}N</td>
<td>11.01</td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td>i-Pr\textsubscript{2}NH</td>
<td>10.96</td>
<td>26</td>
<td>2</td>
</tr>
<tr>
<td>Et\textsubscript{2}NH</td>
<td>10.98</td>
<td>24</td>
<td>0</td>
</tr>
<tr>
<td>n-BuNH\textsubscript{2}</td>
<td>10.77</td>
<td>25</td>
<td>93</td>
</tr>
<tr>
<td>Piperidine</td>
<td>11.12</td>
<td>6</td>
<td>96</td>
</tr>
<tr>
<td>Pyrrolidine</td>
<td>11.27</td>
<td>2.5</td>
<td>91</td>
</tr>
</tbody>
</table>

The conclusion that can be drawn from Table 1 is that the two cyclic bases piperidine and pyrrolidine gives the best yield and the highest reaction rate, but in the presence of CuI only the reaction rate differs for the bases.

These results show that the reaction rates are not only described by the base strength as Brandsma et al.\textsuperscript{21} and Dieck and Heck\textsuperscript{6} described. It can be seen that both i-Pr\textsubscript{2}NH and n-BuNH\textsubscript{2} are weaker bases than Et\textsubscript{3}N, and both have higher reaction rates than Et\textsubscript{3}N.

Moreover if we are looking at the pK\textsubscript{a}-values of Et\textsubscript{3}N and e.g. piperidine, which only differ in pK\textsubscript{a}-values by 0.11, it is very hard to believe that this difference would displace the equilibrium in Scheme 3 significantly more towards the right, and thereby produce more of the acetylide. From the reaction mechanism suggested by Sonogashira (Figure 16), the formation of the acetylide is the rate-determining step in the catalytic cycle B.

\[
\text{R} \text{=CH} \quad + \quad \text{Amine} \xrightarrow{\text{Amine}} \quad \text{R} \text{=CH}^{-} \quad + \quad \text{H}^{+} \text{Amine}
\]

\textbf{Scheme 3.} Acid-base equilibrium between an amine and an acetylene compound.

Results from Sanechika et al.\textsuperscript{22} shows that the yield of the reaction depends to a rough approximation on the strength of the bases. The results show that if you choose a weak a base such as pyridine (pK\textsubscript{a} = 5.25)\textsuperscript{20} the reaction will not proceed, which is also expected as seen in the light of the reaction mechanism suggested by Sonogashira.

Why Et\textsubscript{3}N does not work as well as e.g. piperidine, could be that the ethyl groups are a steric hindrance, which slows the abstraction of the proton from the acetylene.

From Table 1 it is seen that the choice of base does not matter, when a copper halide is used as co-catalyst. The reason is probably that the acetylene reacts with Cu\textsuperscript{+} to form the copper acetylde, which can coordinate with the Pd catalyst. It is well known that 1-alkynes react very well with heavy metal ions (Ag\textsuperscript{+}, Cu\textsuperscript{+}) to form insoluble and often unstable salts.\textsuperscript{23}

3.1.3.3 The catalyst

Generally the Pd catalysts may be divided into the following two main groups: The Pd(II) salts and the Pd(0) complexes. There are many different types of Pd catalysts in each group, but commonly used in the Pd(II) group is Pd(PPh\textsubscript{3})\textsubscript{2}Cl\textsubscript{2} and Pd(OAc)\textsubscript{2}, and in the Pd(0) group
it is Pd(PPh$_3$)$_4$ and Pd$_2$(dba)$_3$ (dba=dibenzylideneacetone), which is used in a combination with a copper(I) halide usually CuI.$^{21}$

As described in the reaction mechanism section (chapter 3.1.1) Pd(0) is the active species in the coupling reaction, which is why all the Pd(II) catalysts undergo a reductive elimination to generate the active Pd(0) compound. This reductive elimination will lead to the same amount of the by-product bis-acetylenes as the amount of Pd(II) catalyst.

In general it is very difficult to predict which Pd catalyst will work best in a given reaction, because it is not always the same catalyst, which works best in all types of reactions. Alami et al.$^{18}$ have demonstrated this by an investigation of the activity of four different Pd catalysts on two very similar systems. The two systems are shown in Scheme 4 and Scheme 5, and the results of the investigation are shown in the corresponding tables (Table 2 and Table 3, respectively).

**Scheme 4.** The system used for investigation of the catalyst activities in a coupling reaction between a terminal acetylene and a vinyl iodide.

**Scheme 5.** The system used for investigation of the catalyst activities in a coupling reaction between a terminal acetylene and a vinyl chloride.

**Table 2.** Catalyst activity in a coupling reaction between a terminal acetylene and a vinyl iodide.$^{18}$

<table>
<thead>
<tr>
<th>[Pd]</th>
<th>Time</th>
<th>Isolated yield [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pd(PPh$_3$)$_4$</td>
<td>15 min</td>
<td>93</td>
</tr>
<tr>
<td>PdCl$_2$(PPh$_3$)$_2$</td>
<td>1 h</td>
<td>82</td>
</tr>
<tr>
<td>PdCl$_2$(PhCN)$_2$</td>
<td>22 h</td>
<td>57</td>
</tr>
<tr>
<td>Pd(OAc)$_2$</td>
<td>22 h</td>
<td>55</td>
</tr>
</tbody>
</table>

**Table 3.** Catalyst activity in a coupling reaction between a terminal acetylene and a vinyl chloride.$^{24}$

<table>
<thead>
<tr>
<th>[Pd]</th>
<th>Time</th>
<th>Isolated yield [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pd(PPh$_3$)$_4$</td>
<td>16 h</td>
<td>11</td>
</tr>
<tr>
<td>PdCl$_2$(PPh$_3$)$_2$</td>
<td>20 h</td>
<td>93</td>
</tr>
<tr>
<td>PdCl$_2$(PhCN)$_2$</td>
<td>0.5 h</td>
<td>93</td>
</tr>
</tbody>
</table>
On the basis of the descriptions found in the literature\textsuperscript{1,5,8,18,21,25} of different types of Sonogashira cross coupling reactions it seems in general not to be so important to choose a specific catalyst. This is because all the catalysts give a high yield with a reaction time less than one day, if the halide is an iodide and a copper halide is used as a co-catalyst. It is common to use either Pd(PPh\textsubscript{3})\textsubscript{4} or PdCl\textsubscript{2}(PPh\textsubscript{3})\textsubscript{2} as catalyst, because they generally work very well, are easy to handle and easy to prepare (see appendix A).

It should be emphasized that the use of both a copper halide as the co-catalysts and additionally triphenylphosphine greatly enhance the rate of the coupling reaction, but if too much triphenylphosphine is added it slows down the reaction rate and there is a loss in selectivity.\textsuperscript{9c}

### 3.2 CTS cross coupling

In 1998 Crisp, Turner and Stephens developed a new type of Sonogashira cross coupling reaction\textsuperscript{26} (CTS cross coupling) without copper(I) halide, to try to avoid formation of the homocoupling of terminal alkynes. Instead of the normal co-catalyst CuI they used a combination of ZnCl\textsubscript{2} and NaI as co-catalysts (see Scheme 6).

\[
\begin{align*}
\text{Pd(PPh}_3\text{)}_4 & & \text{ZnCl}_2/\text{NaI} \\
\text{I} & & \text{H} \\
\text{94 \% yield}
\end{align*}
\]

Scheme 6. The CTS cross coupling between an aryl iodide and terminal acetylene. Note that ZnCl\textsubscript{2} is used without drying.

The ZnCl\textsubscript{2}/NaI co-catalyst system works very well for a coupling reaction between an active aryl halide such as an aryl iodide and a terminal alkyne, and no homocoupled alkyne was detected. Even a coupling reaction with less reactive electrophiles such as an electron rich aryl bromide (see Scheme 7) works very well and only a minimum amount of the homocoupled alkyne is detected.

\[
\begin{align*}
\text{MeO} & & \text{Br} & & \text{Bu} & & \text{H} \\
\text{Pd(PPh}_3\text{)}_4 & & \text{ZnCl}_2/\text{NaI} & & \text{Piperidine} \\
\text{86 \% yield}
\end{align*}
\]

Scheme 7. The CTS cross coupling between an aryl bromide and terminal acetylene.

By changing the solvent from piperidine to a combination of DMSO, DBU (1,8-diazabicyclo[5.4.0]undec-7-ene) and NEt\textsubscript{3} and changing ZnCl\textsubscript{2} to Zn dust the yield can be increased to ~100 \% with a minimal amounts of homocoupled alkyne being formed.\textsuperscript{26}

### References

Sonogashira

5 Tsuji, J., Palladium reagents and catalysts, 1995, John Wiley & Sons Ltd., Chichester, chapter 1.
Since the discovery in 1974 of the high electrical conductivity for doped polyacetylene, a large interest in conjugated polymers has followed. Especially the applications for polymer electronic circuits, light-emitting diodes and the polymer solar cells have received considerable attention. Concerning the different types of polymers within the application for POPV devices, the poly(p-phenylenevinylene)s (PPVs) are by far the most investigated polymers. Recently, focus has been towards the low band gap polymers, in order to get a better match with the solar spectrum. Common to most of the conjugated polymers are that they consist of a stiff backbone, which in many cases is linear and can generally be regarded as rigid rods. The poly(phenyleneethynylene)s (PPEs), which are the closet relative to PPVs, are due to the triple bonds among the stiffest polymers. The PPEs have a very high degree of linearity if the arenes are para coupled (180° between the bonds of the arenes). The PPEs can be regarded as nearly perfect rigid rods as long as the chains are under 15 nm long or under 20 PE repeating units long. Above this length the PPE chains will behave as wormlike chains. This high degree of linearity and in addition an extra π bond compared to the PPVs, made me believe that the PPEs would be much more efficient in transferring energy to the dye molecule, than the JA and JPJ systems of Krebs et al.

In defiance of the close relation to the PPVs, the PPEs have only received very little attention. However it has been demonstrated that the PPEs possess fascinating properties such as being: Blue and green Light Emitting Diodes (LEDs), photoluminescent polarizers, TNT chemosensory materials, exhibit three-dimensional energy migration in solid films, and molecular wires.

One of the attractive properties of the PPE is that they give the possibility of using the Sonogashira cross coupling reaction (see chapter 3) for the synthesis of monodisperse oligomers. These can be used as model compounds of the corresponding polymers. Investigations of e.g. the influence of the chain length on the energy transfer properties and perhaps the possibility to growth crystals for structural investigations. Some groups have already reported synthesis of mono disperse oligomers of the PPE type.

As mentioned in the introduction, the monomer unit in this project was chosen to be 2,5-dioctyltolanylalkynylene, see Scheme 8. The octyl groups were included to ensure solubility of the polymers, and are only placed on every second phenyl ring to minimize an eventual torsion angle out of plane between the individual phenyl groups. Further it has been shown that larger alkyl groups such as dodecyl groups on every phenyl groups will give rise to a slight distortion from planarity of the backbone of the polymer in the solid state. In the solid state, which is the form of the polymer in employed devices, it would be preferred to avoid decrease of conjugation length, which a distortion will cause. Of course it is expected that the backbone of the polymers will differ from planarity in solution, since the barrier of the rotation around the triple bond of an alkyne is less than 1 kcal/mol, and the van der Waals forces will ensure that the atoms in the phenyl groups will be as far from each other as possible.
The strategy of the synthesis of the monomer unit is shown in Scheme 8. All detailed information can be found in appendix A (compound 1, 6 and 7) and Appendix B paper 1 (2-5 and 8-10). The synthesis descriptions in these two appendixes represent the normal amount of compounds synthesized at one time. All have been performed several times, until 100 g of the protected monomer unit (9) was synthesized. The synthesis involved the commercially available compounds 1,4-dichlorobenzene and 1-bromo-4-iodobenzene and as such it is a little
tedious due to the 10 steps. The synthesis can be divided into three main parts: A, B and C. In part A the key compound 4-bromo-2,5-dioctylphenylethynylene (5) is synthesized, which involves five steps. The five steps is a Grignard reaction, a simple addition of bromine, a bromine-iodine exchange reaction, a Sonogashira cross coupling reaction and finally the removal of the trimethylsilane protective group.

The first two steps are standard reactions and were running smoothly without any trouble. In the third step, which is the bromine-iodine exchange of compound 2 using n-BuLi in THF at 195 K, it was found to be important to ensure that none of the compound 2 crystallized on the side of the flask during the addition to the n-BuLi solution, when this took place it was not able to react with the n-BuLi, and a mixture of compound 2 and 3 was obtained. This mixture is difficult to separate by recrystallization due to co-crystallization. Otherwise the reaction proceeds well. In the fourth step where compound 3 is cross-coupled with TMSA by a Sonogashira cross coupling reaction to form the 1-(4-bromo-2,5-dioctylphenyl)-2-trimethylsilylacetylene (4), it was found to be important that the coupling reaction took place at low temperature to increase the selectivity between iodo versus bromo substitution. Even at 273 K we obtained ~2 % of 1,4-di(trimethylsilyl)ethynylbenzene (B1) as by-product, which were detected by 1H and 13C NMR, GC-MS and MALDI TOF. B1 was difficult to remove, and was able to react further in some of the following steps to produce by-products such as B5. In this step I also detected 2-4 % of 1,4-bis-trimethylsilylbutadiyne (B2) as by-product, which was expected due to the use of the Pd(PPh3)2Cl2 catalyst (see chapter 3.1.2). B2 was not a problem, because when the TMS group is removed in the second step, the 1,3-butadiyne is removed by evaporation. In the last step the TMS group was removed by K2CO3 and the 4-bromo-2,5-dioctylphenylethynylene (5) was obtained.27

In part B the key compound 1-(4-iodophenyl)-2-trimethylsilylacetylene (7) was synthesized in two steps. The first step was a Sonogashira cross coupling reaction similar to step 4 in part A. In this step it was also important to run the reaction at low temperature to reduce the by-product 1,4-bis(trimethylsilyl)benzene B3. As in part A the by-product B2 was also detected in this step. Fortunately both B2 and B3 could be removed by recrystallization.

In Part C the monomer unit 1-iodo-2,5-dioctyl-4’-ethynyltolane (10) was obtained in three steps. The first step being a Sonogashira cross-coupling reaction between compound 5 and 7 to obtain 1-bromo-2,5-dioctyl-4’-trimethylsilylheptyltolane (8). In this step ~ 3 % 1-bromo-2,5-dioctyl-2’,5’-dioctyl-4-(4’-trimethylsilyl)phenylethynyltolane (B4) was detected as a by-product with 1H and 13C NMR and MALDI TOF, which is a result of insufficient iodo bromo selectivity. Furthermore ~1 % of compound 5 was found, which is a result of incomplete reaction ~1 % of 1,4-dioctyl-2,5-bis(4’-trimethylsilyl)phenyl)-ethynylbenzene (B5) was also detected. The presence of B5 is a result of the difficulty in separating B1 from compound 4. The Rf values in n-heptane of compound 5, 8, B4 and B5 are 0.58, 0.38, 0.25 and 0.08, so it was possible by flash chromatography to separate 8 from the by-products. The removal of by-products in this step was found to be essential in order to control the later polymerization process. As a consequence the yield of pure compound 8 obtained was low. It should be noted that besides the above mentioned by-products, some minor by-products were also detected, but the amount of these was so small that it was impossible to determine the structures.

The second step was a standard bromine-iodine exchange as described in part A, which ran smoothly. The last step consists of removing the TMS group by K2CO3, which also runs
smoothly. This was only done immediately before the monomer unit (10) should be used, to avoid reaction of the reactive terminal ethynylene groups.

Through this synthetic route I successfully produced about 100 g of the protected monomer unit (9). Taught by experience, 100 g of the monomer unit is necessary, if both monodisperse oligomers, polymers, and super molecular structures like NPN is to be synthesized. In Figure 17 and Figure 18 the ¹H- and ¹³C NMR spectra of the protected monomer (9) are shown and in Figure 19 and Figure 20 the ¹H- and ¹³C NMR spectra of the monomer (10) are shown. By comparing the two ¹H spectra it can be seen that 9 is characterized by containing the TMS group showed by a singlet around 0.6 ppm while 10 is characterized by the absence of this peak with a peak at around 3.1 ppm due to the ethynylene proton. By comparing the two ¹³C spectra it can be seen that they only differ in the two carbon atoms in the ethynylene group at which the TMS group in 9 is attached. In compound 9 these two carbon atoms are deshielded by the TMS group and have the chemical shift values of 105.3 and 97.0 ppm, while in 10 the chemical shift values are shifted to 84.0 and 79.6 ppm. NMR therefore offers a fast, easy and effective method to distinguish between these two compounds, and to determine whether a Sonogashira cross coupling reaction has run to completion or not.

![Figure 17. The ¹H-NMR spectrum of 9.](image-url)
Figure 18. The $^{13}$C-NMR spectrum of 9.

Figure 19. The $^1$H-NMR spectrum of the monomer 10.
Figure 20. The $^{13}$C-NMR spectrum of the monomer 10.

References

The Monomer Unit

Monodisperse oligomers are interesting because they serve as excellent models for their corresponding polydisperse polymers. They can give specific information about electronic, photonic, morphological properties, and how long the oligomers have to be before they act like the polydisperse polymers.\(^1\) By synthesizing a series of monodisperse oligomers with different length, the interesting chain dependent properties such as charge carrier mobilities, effective conjugation length, electronic structure and the morphology, can be investigated. Furthermore, characterization of well-defined materials with e.g. X-ray diffraction and NMR spectroscopic techniques are easier than of the polydisperse polymers.\(^1\)

In order to synthesize a series of monodisperse phenyl acetylene terminated oligo(1-(2,5-dioctyltolanyl)ethyne), I was inspired by Godt et al.\(^2\) to use the synthetic route illustrated in Scheme 9.

The chosen synthetic route gives the opportunity to synthesize monodisperse oligomers in a stepwise manner with precise molecular length. The synthetic route consists of two steps, and starts with n=0, which means that in the first step phenyl acetylene is coupled with 10 by a Sonogashira cross coupling reaction to produce the phenyl acetylene terminated monomer. This is done to ensure that the oligomers are not capable of homo coupling in further steps. Godt et al. have not used phenyl acetylene to terminate the monomers, but instead started with the bromo monomer to which the iodo monomer was coupled. In doing it this way, they make use of the bromo iodo selectivity in the Sonogashira cross coupling reaction to ensure that the produced monodisperse oligomers do not homo couple. I'm not convinced that the bromo iodo
selectivity is strong enough to ensure that none of the oligomers are homo coupled; therefore I decided to terminate them with phenyl acetylene. The synthesis of the phenyl acetylene terminated monomer worked well, which was documented by NMR. It was purified by dry column vacuum chromatography using gradient of n-heptane/ethyl acetate as eluent to ensure that the first step did not contain remnants of phenyl acetylene or \( \text{10} \), which in later steps could result in polydisperse oligomers or even polymers. In step two the TMS protecting group of the phenyl acetylene terminated monomer was removed by a base (\( \text{K}_2\text{CO}_3 \) in a 1:1 mixture of methanol and dichloromethane). The reaction times of step 1 and 2 were set to two and one day respectively, to ensure that each step in the synthetic cycle was completed before proceeding to the next step. Further, more catalyst was added in step 1 after one day, to ensure that there was active catalyst present in the reaction mixture at all times. The choice of the catalyst was \( \text{Pd(PPh}_3\text{)}_4 \) because in theory it should not form the 1,4-diarylbutadiynes as a by-product (see chapter 3.1.2).

The idea of the synthetic route was then that it should be carried out by repeating these two steps continuously with only a short aluminium oxide column as the purification method. This way the intention was to e.g. synthesize the trimer in 95 % purity, with only minor impurities such as uncoupled monomer, dimer or possible 1,4-diarylbutadiynes from an oxidative homocoupling of the unprotected ethynyl groups. The 95 % pure trimer should then be purified by preparative Size Exclusion Chromatography (SEC).

![Figure 21](image-url)

**Figure 21.** The SEC trace of what should have been the pure trimer. The molecular weight of the trimer is 1441.27 g/mol, and correspond to the highest peak in the middle of the trace (~16.30 min).

Unfortunately when the raw trimer product was analyzed by SEC (see Figure 21) it contained many by-products, and the purity of the trimer was not in the range of 95 %. In the SEC trace in Figure 21 the high peak in the middle (around 14 min.) corresponds to the trimer. The molecular weight of the trimer is 1441 g/mol, and from SEC it can be seen that it is determined to be 2072 g/mol. The reason for this mismatch is, that in SEC it is not the molecular weight that is determined but the hydrodynamic volume. The hydrodynamic
volume is then converted to mass by comparing with a calibration curve made by well defined polystyrene standards. The polystyrene standards are flexible randomly coiled polymers diametrically opposed to the PPEs that are rigid rods.\textsuperscript{4,5} It is therefore obvious that the SEC method will overestimate the molecular weight of PPEs. Tour et al. have made a calibration curve for $M_n$ values of precisely defined oligo(thiophene ethynylene)s and oligo(phenylene ethynylene)s determined by SEC relative to polystyrene standards.\textsuperscript{1} From the calibration curve it can be seen that the SEC method overestimate the mass of oligo(phenylene ethynylene)s between 50 % and 66 %, when the oligomers has a mass about 1500 g/mol. This result agrees very well with our measurement of the $M_n$ in the case of the trimer, which is 43 % to large. It should be noted that the slope of the calibration curve depends on the actual molecular weight. The dependence of the molecular weight is such that the deviation between the molecular weight determined by SEC and the actual molecular weight increase with the actual molecular weight of the oligomers.

From the SEC trace it can be seen that the product contain both by-products with smaller and larger size. The by-products of lower size are most likely uncoupled monomers and dimers and possibly the 1,4-diarylbutadiynes originating from oxidative dimerization of the unprotected monomer or couplings between the unprotected monomer and the unprotected dimer. The by-products with higher mass most likely include the 1,4-diarylbutadiynes originating from homocoupling of the unprotected dimer or a polymerisation of unreacted monomer units.

In order to reduce the amount of by-products several different attempts were made. For instance different types of catalysts such as Pd(PPh$_3$)$_4$, Pd(PPh$_3$)$_2$Cl$_2$ or Pd$_2$(dba)$_3$, different types of bases such as NEt$_3$ or Piperidine, different types of co-catalysts such as CuI or ZnCl$_2$/NaI (the CTS coupling see chapter 3.2) and changing the reaction time from two days to one week have been attempted but without any significant difference in the amount of by-products formed. Furthermore it was attempted to boil all the solvent under an argon atmosphere before use, to avoid oxygen which can cause the formation of the 1,4-diarylbutadiynes. No significant difference in the amount of formed by-products was observed in spite of all these trials.

Seen in the light of several months work with lot of different attempts to synthesize the trimer, with a poor yield and highly laborious to purify a reasonable amount of the trimer by SEC, I decided to stop further work of making monodisperse phenyl acetylene terminated oligo(1-(2,5-dioctyltolanyl)ethyne) by this type of synthetic route.

Instead I decided to try to synthesize monodisperse oligomers by using the solid phase synthetic route shown in Scheme 10. The route is inspired by Tour et al.\textsuperscript{5} All the experimental details can be found in appendix A.
Monodisperse Oligomers

Scheme 10. The chosen solid phase synthetic route to make monodisperse oligomers.

The advantage of this solid phase synthetic route is, that after the monomer units are attached to the resin, they cannot interact with each other and thus cannot form the 1,4-diarylbutadiynes. This means that if the couplings are 100% head-to-tail coupled, monodisperse oligomers in principle can be obtained without any of the by-products described in chapter 3.1.2. If the couplings proceed in at least 95% yield in each step, the oligomers can be separated by preparative SEC. A coupling success rate of less than 95% is not acceptable, it becomes too expensive to separate the different oligomers from each other.

The first step in the solid phase synthesis is to form the anchor unit, which is achieved by the coupling of the Cs salt to Merrifield's resin (chloromethyl polystyrene). In Solid phase synthesis all coupling and purifying (washes) routines take place in a special flask made for solid phase synthesis (see Figure 22).

Monitoring the yield and progress of the reaction are more complicated in solid phase synthesis than in normal solution synthesis. In solid phase syntheses it is not possible to use TLC and other chromatography techniques because the resin is insoluble in the most common solvents. Spectroscopic techniques as e.g. solid state NMR and FTIR can be used with success especially for short oligomers, but especially solid state NMR is challenging to use and I have not used this method, but instead used FTIR.

The FTIR techniques have the advantage that in these Sonogashira coupling reactions there are two vibrations in a frequency range where normally there are no other absorptions. These two vibrations are the $sp\ C-R$ stretch and the $sp\ C\equiv C-R$ stretch. The frequencies where those vibrations occur depend on the type of radical $R$ attached to the $C$ atoms in the alkynyl groups. Generally, the heavier the $R$ radicals are the higher the frequencies of the vibrations are.
Generally, by this technique it is possible to distinguish between the unprotected alkynyl group, the alkynyl group protected with a trimethylsilyl group and the alkynyl groups in the backbone of the oligomers. In other words, when the cross coupling reaction is complete the vibration of the sp C-H and the C≡C-H stretch disappears. The stretch from the C≡C-R terminals and the backbones C≡C of the oligomers should be left.

![Figure 22. The equipment for the solid phase synthesis.](image)

Tour *et al.*\(^6\) have used FTIR to monitor cross coupling reaction for the protected monomer 1-iodo-3-dodecyl-4-[(trimethylsilyl)ethynyl]benzene. They found the terminal alkynyl carbon-hydrogen and the carbon-carbon stretches at 3311 (strong) and 2109 (weak) cm\(^{-1}\) respectively, while the carbon-carbon stretch of the trimethylsilyl-terminated alkyne occur at 2156 (weak) cm\(^{-1}\).

In my system where the deprotected monomer consists of the 1-iodo-2,5-dioctyl-4'-ethynyltolane, I observed the terminal alkynyl carbon-hydrogen and the carbon-carbon stretches at 3286 (strong) and 2077 (weak) cm\(^{-1}\) respectively, while the carbon-carbon stretch of the trimethylsilyl-terminated alkyne occurs at 2154 (weak) cm\(^{-1}\). The vibration frequencies of the carbon-carbon stretch for the alkyne groups in the backbone occur at 2211 cm\(^{-1}\) (see Figure 23).

Hence by observation of the different alkynyl stretch vibration intensities, FTIR can be used to monitor whether a given coupling reaction is 100 % completed or e.g. only 50 % done.

Both FTIR and solid state NMR have unfortunately the disadvantage that they have a relatively low sensitivity, when they are used for resin analysis.
A more effective method to monitor the coupling reaction is to determine the increase in weight for the resin after each reaction. It is a very slow but very effective method.

The yield after the first coupling reaction can be determined by measuring the increase in weight, \( \Delta W \), of the resin after the reaction and the amount of precipitated silver chloride, \( n_{Cl} \), from the first coupling reaction.

The yield, \( Y \), is given by:

\[
Y = \frac{n_p}{n_{Cl}} \tag{5.1}
\]

where \( n_p \) is the amount of the product in the final resin, and \( n_{Cl} \) is the number of moles of precipitated silver chloride, which is equal to the theoretical moles of free couplings sites.

\( n_p \) can be determined from the following two conditions:

\[
\begin{align*}
1 & \quad n_{Cl} = n_{OH} + n_p \\
2 & \quad \Delta W = m_{OH} + m_p - m_{Cl}
\end{align*} \tag{5.2}
\]

where \( m_{OH} \), \( m_p \) and \( m_{Cl} \) is the mass of the coupled OH (from water in the solvents and atmosphere) and product to the resin and the mass of precipitated chloride ions respectively.

By substitution of condition 1 into condition 2 it can be showed that \( n_p \) is given by:
Monodisperse Oligomers

\[ n_p = \frac{\Delta W - (M_{OH} - M_{Cl})n_{cl}}{M_p - M_{OH}} \]  
(5.3)

where \( M_{OH}, M_p \) and \( M_{Cl} \) is the molar mass of the coupled OH group, the coupled product and the chlorine.

By substitution of equation (5.3) into (5.1) the following expression for the yield is obtained:

\[ Y = \frac{\Delta W - (M_{OH} - M_{Cl})n_{cl}}{(M_p - M_{OH})n_{cl}} \]  
(5.4)

From the experimental details in appendix A it can be seen that in step 1 \( \Delta W = 0.425 \text{ g} \) and that \( n_{cl} = 9.064 \times 10^{-3} \text{ mol} \), which gives a yield of 51 %. This means that 51 % of the free coupling sites are occupied by the anchor unit and all other sites are occupied by OH groups.

Step 2 is a Sonogashira cross coupling reaction where the protected monomer unit (9) is coupled to the anchor unit. The yield in step 2 is given by:

\[ Y = \frac{n_{pl}}{n_p} \]  
(5.5)

Where \( n_{pl} \) is the amount of 9 coupled to the anchor unit. The weight increase after the coupling reaction is:

\[ \Delta W_1 = m_{pl} - m_H \]  
(5.6)

Where \( m_{pl} \) is the weight of 9 (minus the weight of an iodine atom) coupled to the resin, where \( m_H \) is the weight of the hydrogen atoms which are eliminated in the coupling reaction. The \( n_{pl} = n_H \), because the only way a hydrogen atom can be eliminated is when 9 is coupled to the site, therefore is \( n_{pl} \) given by:

\[ n_{pl} = \frac{\Delta W_1}{M_{pl} - M_H} \]  
(5.7)

\( M_{pl} \) and \( M_H \) is the molar mass of 9 (minus the weight of an iodine atom) and a hydrogen atom respectively. By substitution of equation (5.7) and (5.3) into (5.5) following expression for the yield is obtained:

\[ Y = \frac{M_p - M_{OH}}{\Delta W - (M_{OH} - M_{Cl})n_{cl}} \times \frac{\Delta W_1}{M_{pl} - M_H} \]  
(5.8)

From appendix A it can be seen, that \( \Delta W_1 = 0.552 \text{ g} \), when 5.034 g of the resin from step 1 was used. This corresponds to a yield of 50 %. In order to get a higher yield the resin was used again and a new coupling was performed under the same conditions as the first one. In this case the \( \Delta W_1 = 0.075 \text{ g} \), which corresponds to a total \( \Delta W_1 = 0.627 \text{ g} \) and a total yield of 57 %. A third coupling attempt did not lead to any further increase in mass.

In step 3 the TMS protecting groups were removed by treatment with TBAF. In appendix A it can be seen that the mass was reduced by 0.208 g, but should only have been reduced by 0.091 g. I have no explanation for why the mass was reduce so much. I have checked the resin by IR and it shows that it was 100 % deprotected.
In step 4 one more Sonogashira cross coupling reaction was performed to produce the dimer unit. The increase in mass was only 0.464 g in first attempt, 0.068 g in second attempt and 5 mg in third attempt, which give a total mass increase of 0.537 g and corresponds to a yield of 86%.

In step 5 the dimer was cleaved from the resin by treatment with strong base (NaOH) followed by acidification with HCl. The SEC analysis of the dimer showed unfortunately that there was lots of polymer material in the sample, so no attempt was made to purify the dimer.

The solid phase synthesis was attempted three times and every time there were problems with the coupling yield of less than 95%, too much loss in mass during the deprotection steps and formation of polymer materials in the cleaving steps.

On the basis of the unsatisfactory coupling yields and problems with the formation of polymer materials together with the immense time consumption in each step, the solid phase synthetic route was dropped as well.

From these failed trials aimed at synthesizing monodisperse oligomers a remarkable observation can be made. It seems that the coupling reactions continue after the deprotection steps, which is strange because the catalysts should have been removed or was it? If there are remnants of the catalyst present under the deprotection step, where base is also present, then the environment for a Sonogashira cross coupling reaction is present along with the unprotected and uncoupled monomers. This is especially critical in the case of the solution route, whereas in the solid phase synthesis the uncoupled monomers are washed out. The unprotected and uncoupled monomers will be able to couple with each other to produce long oligomers or polymers. Further it is a possibility that the terminated oligomers will homocouple and form 1,4-diarylbutadiynes. By screening the literature I found that there has only been a sporadic interest to ensure no remnants of the Pd catalysts left in the newly synthesized organic compounds, and that this interest has increased significantly during the course of this project.

I suggest therefore that the problems concerning the synthesis of the monodisperse oligomers in high purity is due to the remnants of Pd in the samples under the deprotection steps. This is supported in the literature by Garret and Prasad and Krebs et al.

The topic concerning the remnants of Pd in the synthesis organic compounds will be treated in details in chapter 7.

References
Monodisperse Oligomers


The iodine-terminated homopolymer poly[(2,5-dioctyltolanyl)ethynylene], $N_n$, were synthesized by employing a standard Sonogashira cross-coupling as outlined in Scheme 11. All the experimental details can be seen in paper 1 in appendix B. As shown in Scheme 11 I have used Pd(PPh$_3$)$_4$ as catalyst instead of Pd(PPh$_3$)$_2$Cl$_2$ to avoid or minimize the formation of the 1,4-dibutadiyne by-products (see chapter 3.1.2). In connection to the synthesis of $N_n$, it should be mentioned that attempts to dry the product in a vacuum oven at room temperature gave a darkened product that was no longer soluble in common solvents. This behavior has been observed earlier, and is probably a result of cross linking of the polymer chains. Therefore, I always kept the polymer wet with the appropriate solvent. The molecular weight of the polymer was determined by SEC and the results can be seen in Table 4.

Scheme 11. The synthetic route of synthesizing the iodine-terminated homopolymer poly[(2,5-dioctyltolanyl)ethynylene], $N_n$.

To synthesize the porphyrin-linked homopolymer, NPN, I have tried two different synthetic routes as shown in Scheme 12 and Scheme 13. Both routes should in theory give a super molecular assembly with the NPN structure. The only way the two different synthetic routes differ is in the way the polymer segment is coupled to the porphyrin unit and thereby the distance from the octyl groups to the porphyrin unit is different. For NPN A, where the distance between the octyl groups and the porphyrin unit is shorter than for NPN B, a decrease in the conjugation length will be expected due to steric effects of the octyl groups. In spite of the expected smaller conjugation length of NPN A compared to NPN B I decided to synthesize NPN A first, as I thought that it would be easier to incorporate the diethynylporphyrin, 11, than the dibromoporphyrin, 12, into the polymer chains because compound 11 was expected to have a higher reactivity towards the individual monomers, 10. The polymerization of 10 in presence of 11 proceeded smoothly and a green polymer product was obtained. In order to ensure that only one porphyrin unit was incorporated into the polymer backbone and to investigate the chain length dependence on the energy transfer from the polymer backbone to the porphyrin unit, the crude NPN A product was divided into seven fractions by preparative SEC.
Scheme 12. The unsuccessful synthetic route to synthesize NPN.  

Scheme 13. The successful synthetic route to synthesize NPN.
The molecular weight of the crude NPN A and the seven fractions were determined by SEC and listed in Table 4. If only one porphyrin unit is sandwiched between two polymer chains, the absorption ratio N/P between the polymer, N, and the porphyrin, P, should increase going from the fraction with lowest molecular weight (fraction 7) to the fraction with highest molecular weight (fraction 1) as a consequence of the longer polymer chains in the fraction with highest molecular weights. In Figure 24 the UV-Vis spectra of the seven fractions are shown.

![UV-Vis spectra of seven fractions](image)

**Figure 24.** The normalized (according to $\lambda_{\text{max}}$) UV-Vis spectra of the seven fractions of NPN A in a CHCl$_3$ solution. Very little variation of the N/P absorption ratio is observed.$^2$

From the UV-Vis spectra of the seven fractions it can be seen that the zinc porphyrin molecule, whose UV-Vis spectrum is characterized by a Soret band with maximum at around 460-470 nm and the Q-bands between 600 and 700 nm, was successfully incorporated into the polymer backbone. Furthermore, it can be seen that the N/P absorption ratio, astonishingly enough, is roughly constant, which indicate that it is not only NPN A that is produced but also by-products that consists of more than one porphyrin molecule per polymer chain. The N/P absorption ratio is shown in Table 4. The problem with the synthetic route for NPN A is that the growing chain always exposes an ethynyl group. The free ethynyl groups are known (see chapter 3.1.2) to be capable of undergoing an oxidative homocoupling to form butadiynes, which are the dominating by-products from the Sonogashira cross coupling reaction. The problem with these types of by-products is, that they leave an iodo terminated polymer, which does not represent a termination of chain elongation, instead it allows both monomer and porphyrin units to couple to itself and the polymer chain can continue to grow. The result will be a polymer product with an even distribution of porphyrin molecules, and where the amount of the porphyrin in the polymer backbone largely depends on the initial ratio between the monomer and the porphyrin, and the amount of oxygen present in the reaction mixture. In the light of these arguments it is therefore most likely that the synthetic route in Scheme 12 leads to both the desired NPN structure and a lot of buta-1,3-diyn-1,4-diyl by-products of the type shown in Scheme 14.
Scheme 14. Some of the possible by-products from the Sonogashira cross coupling reaction between 10 and 11. The by-products are due to the homocoupling of ethynyl groups.\textsuperscript{2}

It is possible to get rid of the homocoupling, which is the dominant side reaction by chromatography in small molecules. In the case of polymers the homocoupling will lead to accumulation so that each polymer chain contains several of this type of defect. I have based on the above drawn the conclusion that it is not possible to use the synthetic route in Scheme 12 to synthesize a super molecular assembly like NPN with a high degree of regionregularity and directionality. All further work with NPN A was therefore abandoned.

For NPN B, I assumed that there would only be incorporated one porphyrin molecule in each NPN polymer chain. The reason is that the porphyrin molecule, 12, is bromo terminated in both ends, so that the ethynyl groups on the monomers are coupled to the porphyrin molecule. In this way the growing chain will always be iodo terminated, and therefore not capable of making a homocoupling or a coupling to other porphyrin molecules. On the other hand it is also expected that a considerable amount of the homopolymer would be present, since any homocoupling of the ethynyl groups will lead to homopolymers with iodo termini in both ends of the chain, and therefore are not capable of incorporating the dibromoporphyrin in the chain.

The polymerization of 10 in presence of 12 proceeded smoothly and a green polymer product was obtained. The crude NPN B polymer product was fractionated by SEC into eight fractions (see Figure 25 for the SEC trace) and the molecular weight, $M_w$, was determined to 11546 g/mol with a polydispersity, $PD$, of 3.46. Note that when a SEC analysis was performed on the crude NPN B and on individual fractions the following day, a very remarkable observation was made. The $M_w$ and the $PD$ were significantly increased, which indicate that a cross-linking between the individual chains took place. To make sure that the cross-linking process was complete before further investigation of the polymer, the crude polymer and the fractions were left standing in CHCl\textsubscript{3} solutions in the hood under the ambient atmosphere to slowly evaporate over a period of three days. The evaporated solids were dissolved in boiling CHCl\textsubscript{3} and insoluble parts were filtrated off before they were refractionated by SEC. The new fractions were stable regarding to the $M_w$ and $PD$ and the determined values are listed in Table 4. Fraction 8 was discarded because it consisted of compounds with very low molecular weight. From Table 4 it can be seen that the crude NPN B increase in $M_w$ and $PD$ from 11546 g/mol and 3.46 to 36949 g/mol and 4.09 during the three days, which is surprisingly much.
Figure 25. The SEC trace of the crude NPN B polymer product (right axes) and the 8 fractions (left axes). The UV-signal is monitored at 383 nm, which corresponds to the $\lambda_{\text{max}}$ of the N domain.\textsuperscript{3}

In Figure 26 the UV-Vis spectra of the seven fractions dissolved in CHCl$_3$ are shown. It can be seen that the NPN B polymer shows the desired variation of the N/P ratio between the individual fractions, which indicate that I in principle succeeded to incorporate only one porphyrin molecule in the polymer backbone.

Figure 26. The normalized (according to $\lambda_{\text{max}}$) UV-Vis spectra of the seven fractions of NPN B in a CHCl$_3$ solution. It can be seen how the absorption intensity of the porphyrin ($\lambda = 451$ nm (Soret band))
and \( \lambda = 520-700 \text{ nm (Q band)} \) compared to the absorption intensity of the N domains varies in the individual fraction and thereby varies as a function of the length of the \( N_n \) domains.\(^2\)

Table 4. Data from the SEC and UV-Vis analyse for all three polymers: \( N_n, \) NPN A and NPN B

<table>
<thead>
<tr>
<th>Compound</th>
<th>( M_w ) [g/mol]</th>
<th>PDI</th>
<th>Absorbance ratio</th>
<th>N/P ratio (UV-Vis)</th>
<th>N/P ratio (SEC)</th>
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</thead>
<tbody>
<tr>
<td>( N_n )</td>
<td>92300</td>
<td>2.90</td>
<td>2.5</td>
<td>8.5</td>
<td></td>
</tr>
<tr>
<td>NPN A crude</td>
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<td>2.12</td>
<td>2.5</td>
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<td></td>
</tr>
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<td>Fraction 1</td>
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<td>2.5</td>
<td>8.5</td>
<td></td>
</tr>
<tr>
<td>Fraction 2</td>
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<td>3</td>
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</tr>
<tr>
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<tr>
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<tr>
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<tr>
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<td>2.5</td>
<td>8.5</td>
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<tr>
<td>Fraction 7</td>
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<td>2</td>
<td>6.9</td>
<td></td>
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<tr>
<td>NPN B crude</td>
<td>36949</td>
<td>4.09</td>
<td>3</td>
<td>10.0</td>
<td>84.6</td>
</tr>
<tr>
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<td>6.5</td>
<td>22.2</td>
<td>214.8</td>
</tr>
<tr>
<td>Fraction 2</td>
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<td>5</td>
<td>17.1</td>
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</tr>
<tr>
<td>Fraction 3</td>
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<td>1.28</td>
<td>4.5</td>
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<tr>
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<td>2.5</td>
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<tr>
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<td>2</td>
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<td>7.2</td>
</tr>
<tr>
<td>Fraction 7</td>
<td>1534</td>
<td>1.17</td>
<td>1.5</td>
<td>5.1</td>
<td>1.5</td>
</tr>
</tbody>
</table>

It should be emphasized that neither SEC nor the UV-Vis analysis give information about the homopolymer content as a result of the oxidative homocoupling of the ethynyl groups. Instead the two methods can give an estimate of the length of the N domains and of the degree of cross linking. From the UV-Vis spectra in Figure 26 it is possible to obtain an average N/P ratio in the NPN B polymer. This is done on the basis of the absorbance ratio between the absorption at \( \lambda = 390 \text{ nm} \) and \( \lambda = 451 \text{ nm} \), which represent absorption mainly from the N and P (the Soret band) domains, respectively. The absorbance ratios for the individual fractions are listed in Table 4. Applying the extinction coefficients for the pure homopolymer \( N_n \) and the pure porphyrin molecule 11 an estimate of the N/P ratio can be obtained. The two extinction coefficients were determined to be \( \varepsilon_{390} = 70000 \text{ M}^{-1} \text{ cm}^{-1} \) and \( \varepsilon_{431} = 240000 \text{ M}^{-1} \text{ cm}^{-1} \) respectively.\(^2\) The \( \lambda_{\text{max}} \) of the Soret band in NPN B is red-shifted with 20 nm compared to 11 due to the increased conjugation length, which is the reason for the use of the \( \varepsilon_{431} \) for the absorption at 451 nm in NPN B. The estimated N/P ratio can be seen in Table 4. From the \( M_w \) of the different fractions determined by SEC it is also possible to get an estimate of the N/P ratio, when it is assumed that there is only one porphyrin molecule in each chain and that the molecular weight of the porphyrin unit is 750 g/mol and the weight of the monomer unit is 425 g/mol in the chains. The estimated N/P ratios on basis of the SEC data are also present in Table 4.

\(^2\) Note that I used the extinction coefficient of 11 instead of 12 because it looks more like the porphyrin unit that is incorporated into the polymer backbone of NPN B.
Comparison of the N/P ratio obtained by UV-Vis with the results from the SEC data a mismatch in both the fractions with low and with high molecular weights is observed. Consider first the low molecular range. In this range the N/P ratio is overestimated by the UV-Vis analysis as a result of the assumption that the extinction coefficient for $N_n$ and $P$ is the same in the polymer and for the isolated molecules. In the low molecular range it is therefore the results from the SEC analysis that are most reliable, while keeping in mind that also the results from the SEC are overestimated due to the mismatch between the hydrodynamic volume of the PS standard and the NPN B. From the data in Table 4 it can be seen that fraction 7 most likely cover the molecules where only one monomer is coupled to the porphyrin. In the range of high molecular weight the mismatch is due to both overestimation of $M_w$ by SEC and to cross-linking of the individually NPN B chains. According to the overestimation of $M_w$ by SEC, Tour et al. have shown that the overestimation highly depends on the size of the molecular assemblies in such way that in the molecular weight range from 600 to 4400 g/mol the overestimation is in the range of 20 to 100 %, with the largest overestimation on the largest molecules. But in the case of large polymers with a chain length of more than 20 PPE units, the polymer will no longer act as a perfect rod but begin to act as a wormlike chain, and the overestimation of $M_w$ will therefore probably be lower. The cross-linking between the individual polymer chains also contribute significantly to the mismatch since more than one porphyrin molecule will be present in the molecule, and therefore give a too small N/P ratio in the UV-Vis analysis. During the work up the $M_w$ of the crude NPN B was increased about three times, which indicate that in average three polymer chains have been linked together.

Altogether it can be concluded that I, in spite of a lot of unforeseen difficulties, have succeeded to synthesize a three domain super molecular assembly of the desired NPN type, and have furthermore fractionated them into 7 fractions with different length of N domains.

References

During the failed attempts to synthesize monodisperse oligomers and during the successful synthesis of the \textit{NPN} super molecular assembly some remarkable and inexplicable observations were made. In connection to the oligomers, it was observed that it was impossible to control the stepwise synthesis of monodisperse oligomers, and that the results always were a distribution of oligomers and polymer products. The lack of control (the reason for failing) in the synthesis can most likely be tracked to the deprotection step that it failed. Concerning the \textit{NPN B} polymer, it was observed that the polymer begins to cross-link after work up and purification with SEC. Furthermore it was observed that standard electroactive devices (as described in chapter 2.3), where the active layer consisting of \textit{NPN B} or \textit{Nn}, had a resistance as low as 30 \textit{\Omega} and when a potential was applied the devices short circuit and sparkles from the films as a results of short circuits could be seen. These remarkable observations together with the fact that Krebs \textit{et al.}\textsuperscript{1} at the same time had observed Pd remnants in PPV materials synthesized by the Pd catalyzed Heck reaction indicated that there may also be remnants of Pd present in the PPE polymers even after the purification by SEC. An Inductively Coupled Plasma Sector Field Mass Spectrometry (ICP-SFMS) analysis of 0.5 g of the \textit{Nn} polymer uncovered that it contained 49 ppm (mg/kg) Pd remnants. The real value is actually much higher, because it was impossible to dissolve all the material. Even after three days in aqua regia more than half of the 0.5 g \textit{Nn} sample was left as an orange sticky material.

The remnants of Pd in the samples can explain the three remarkable observations mentioned above. In the case of the oligomers base is present under the deprotection step and together with the remnants of Pd and oxygen from the filtration steps it provides a breeding ground for the homocoupling of the unprotected oligomers. In the case of the cross-linking of the \textit{NPN B} polymers, it has been reported in the literature more than 50 years ago that transition metals catalyze the trimerization of alkynes.\textsuperscript{2} Furthermore in 1997 and 2001 it was shown by Gevorgyan \textit{et al.}\textsuperscript{3,4} that substituted benzenes can be constructed under Sonogashira cross-coupling conditions without an inert atmosphere from enynes and activated alkynes in a [4 + 2] reaction and from terminal alkynes in a [2 + 2 + 2] fashion. In Scheme 15 a possible trimerization process is outlined, showing how two \textit{Nn} chains, which are not coupled to porphyrins, can form an enyne in presence of Pd(0) catalysts. The enyne can then form the trimerization cross-linking product by reaction with a triple bond in either a \textit{Nn} or a \textit{NPN B} chain. In the last case it is obvious that metal in the active layer will lower the resistance and lead to a short circuit of the device.

In order to find a method to remove the Pd remnants from the polymer products and thereby save all the resources that have been put into the work to produce \textit{NPN B} and \textit{Nn}, the literature has been screened thoroughly. It was found that even though transition metals have been used, as catalysts in the synthesis of new organic compounds, only sporadic attention has been put on remnants of the catalysts in the newly synthesized organic compounds. When the
problems of Pd remnants were discovered in this project, it really seemed that removal of e.g. Pd remnants from samples was a large forgotten area in the literature; fortunately the interest has increased during the time of this project.\textsuperscript{1,5,6,7,8,9,10,11,12,13,14,15,16} The reason for this apparent loss of interest concerning the removal e.g. of Pd from organic compounds is probably due to the fact that most impurities present at a level below 0.1\% do not interfere with characterization techniques like NMR, GC/MS, HPLC, IR, UV-Vis and elemental analysis. However, in compounds used for drugs or electronic applications, the impurity level has to be at a much lower level. In defiance of the lack of literature concerning the removal of Pd from samples, it seems to be common knowledge in the pharmaceutical industry that Pd as a catalyst is not allowed in the two last synthetic steps in synthesis of medical compounds due to the difficulties to remove of Pd remnants.\textsuperscript{17} To emphasize that the removal of Pd from organic samples is not a simple problem came from Professor Steven Ley from Cambridge University. He claimed at the Knud Lind Larsen symposium in Copenhagen 2004 that he could perform cross coupling reactions with compounds from Sigma Aldrich without any addition of catalyst, because the remnants of palladium in these compounds were present in quantities sufficient for catalysis. It should be emphasized that it is not a trivial problem to remove a noble transition metal from organic compounds and to ensure that the level is under the oral concentration limits and the parenteral concentrations limit for drugs, which in oral cases are 5 and 15 ppm for Pd and Cu respectively and in the parenteral cases are 0.5 and 1.5 ppm for Pd and Cu respectively.\textsuperscript{14}

\textbf{Scheme 15.} The possible trimerization process of \textbf{NPN B} observed under Sonogashira cross-coupling conditions.\textsuperscript{18}
In this project two model systems of diphenyl acetylene synthesized under normal Sonogashira cross coupling conditions were made (see Scheme 16) one with a Pd catalytic route and one with a Cu catalytic route, to test the ability of different methods to remove just these two metals from the synthesized diphenyl acetylene. It was observed by ICP-SFMS analysis, that even after filtration through a aluminum oxide column and two recrystallization processes in ethanol, the model reaction contained 610 ppm Pd and 1660 ppm Cu respectively (see Table 6). In the literature it is supported that traces of Pd cannot be removed by crystallization\textsuperscript{7,9,10,19} or by chromatography\textsuperscript{9,10} but that methods like distillation\textsuperscript{9} are needed. With distillation the problem is that not all compounds can withstand high temperatures.

\begin{align*}
\text{A) } & \text{I} + \text{Pd(PPh}_3\text{)}_4 (2 \text{ mol\%}) \xleftarrow{\text{Piperidine/THF}} \text{Pd(I)}_2 \text{ (69 \%) } \\
\text{B) } & \text{I} + \text{H} \xrightarrow{\text{CuI (5 mol\%) \text{ } 2 \text{ PPh}_3 \text{ } K_2\text{CO}_3 \text{ } \text{DMF}} } \text{Cu(I)}_2 \text{ (70 \%)}
\end{align*}

\textbf{Scheme 16.} The model system used to test for remnants of Pd and Cu respectively after different purifying methods.\textsuperscript{20}

\section*{7.1 Diethylammonium DiethyldithioCarbamate (DDC)}

Based on of the literature screening it was attempted to use diethylammonium diethyldithiocarbamate, DDC (see Scheme 17a), to remove Pd remnants from the polymers, as described by Jones et al.\textsuperscript{15} in the case of solid-state synthesis. This attempt was done with some success (experimental details and a more thorough discussion about the removal of Pd from the polymers with DDC can be found in paper 1 in appendix B). After the treatment of N\textsubscript{n} and NPN B with DDC, the device resistance was increased from 30-50 Ω to 0.3-2 MΩ and the devices worked without short circuit. The idea with DDC is that it coordinates to Pd like the triple bonds in the alkynes. The coordination is very strong (log } K (\text{Pd[DDC]}_2) = 64.9) and very fast\textsuperscript{21} and palladium will bind to the DDC instead of to the alkyne groups. DDC and the formed Pd(DDC)\textsubscript{2} complex are both soluble in methanol and can be removed by precipitating the polymer mixture containing the DDC and the complex in large amount of methanol. Unfortunately there were some indications that DDC reacted with the polymers. The treatment of the polymers with DDC was performed in following three time intervals: 5 min., 2 hours and 24 hours. With long reaction times dramatic changes in the properties of the polymers were observed. E.g. it was observed that the polymers were difficult to precipitate in methanol, no electroluminescence could be observed, when a potential was applied on standard devices, and significant changes in the UV-Vis spectra were observed (see Figure 27). On the other hand it was impossible to see any changes in NMR and there were no significant changes in the elemental analysis.

The conclusion is, that fast treatment of the polymers with DDC remove the Pd remnants and makes it possible to make standard electroactive devices for LED and POPV application, while a longer treatment time influence the properties of the polymers. However the analysis of the reaction between the polymers and DDC cannot provide a satisfactory explanation for what happens in the reaction between the polymers and DDC.
The Palladium Saga

Figure 27. The UV-Vis spectrum of the Nₙ polymer before and after the treatment with DDC (2 hours treatments time).¹⁸

7.2 The azothioformamides

To gain knowledge on the effect of DDC on the polymers a lot of resources have been invested in this project to find a better alternative to DDC. This was successfully achieved by the discovery of N,N-diethylphenylazothioformamide, L₁, and N,N-bis(tetraethyleneglycolmonomethylether)phenylazothioformamide, L₂. Both L₁ and L₂ possess the property of being capable of dissolving nanoparticles of many different transition metals by forming neutral and intensely colored electron transfer complexes (see paper 3, 4, 6 and 7 in appendix B for more detailed information and Scheme 17b-c for the structures).

L₁ and L₂ belong to the group of compounds called azothioformamides, which were subject to a lot of interest during the sixties and early seventies due to properties such as antifungal activity,²² antibiotic activity,²³ application as pesticides²⁴ and the property to form electron transfer complexes of the type M-N₂S² with a large range of transition metals (see Scheme 18).²⁵ L₁ and L₂ both coordinate faster and stronger to the transitions metals than the alkynes. It was observed that L₁ and L₂ were not only capable of dissolving nanoparticles of the transition metals, which is the form the remnants of the metals typically are in, but also metals in form of wires. Especially in the case of copper the process of disintegration was fast and effective (see Figure 28). Nickel and Platinum wire could also be dissolved by L₁ (it was not tried with L₂) but the process here was much slower, while in the case of palladium wire no disintegration was observed. The missing detection of dissolved palladium is probably due to a surface phenomenon, because it has been observed that L₁ is capable of dissolving palladium black and palladium nanoparticles.
Scheme 17.

Scheme 18. An Electron transfer series of five members.26

The structural differences of L1 compared to L2 makes L1 an excellent tool to remove transition metals from polymers, while L2 is a better choice in the case of removing transition metals from small organic molecules. It should be mentioned that the synthesis of L1 has been reported earlier,25 but I have devised a simple one-pot synthesis of L1 (see paper 3 in appendix B for the synthetic details). The synthesis of L2 has not been reported before, and has been developed for the purpose of removal of Pd remnants from small organic molecules (see paper 4 in appendix B for the synthetic details).
In the case of \textbf{L1} the procedure to remove transition metals from polymers is very simple and is done by a dissolution and precipitation technique. The polymer product is dissolved in a minimal amount of solvent e.g. chloroform. \textbf{L1} is added and the mixture is stirred over night, whereupon the mixture is precipitated in a lot of methanol and the polymer is collected by filtration (both compound \textbf{L1} and the formed metal complexes are soluble in large amounts of methanol). The dissolution and precipitation technique can be performed again without compound \textbf{L1} to be sure that the filtrate does not contain remnants of compound \textbf{L1} or the complexes.

In the case of \textbf{L2} the procedure to remove transition metals from small molecules is nearly the same as removing transition metals from polymer products with \textbf{L1}. Instead of precipitating the mixture in methanol the mixture was filtered through a short column of silica with a polar solvent such as chloroform. The $N,N$-bis(tetraethyleneglycolmonomethylether)amine groups in \textbf{L2} exhibit a strong affinity for silica in polar solvents such as e.g. chloroform. This means that both \textbf{L2} and the complexes with the transition metals are immobile on the columns, whereas small organic molecules will be mobile on the columns (see \textbf{Figure 29}). The pure organic material can therefore be recovered from the filtrate. The $N,N$-bis(tetraethyleneglycolmonomethylether)amine groups also have the property, that they make \textbf{L2} water soluble, and \textbf{L2} can therefore in principle be used to remove transition metals from aqueous solutions. Unfortunately the
distribution coefficient of $L_2$ between water and organic solvents such as chloroform or 1,3-dichlorobenzene is very low, which means that in practice it is impossible to make an extraction of $L_2$ or the complexes from an organic solvent with water. The other way around on the other hand is possible.

### 7.2.1 Analytic tool

A unique feature of both $L_1$ and $L_2$ and probably the derivations of $N,N$-phenylazothioformamide in general is that the formed electron transfer complexes with transition metals unlike the pure ligand has an absorption with a maximum in the range of 800 nm. Because only very few organic materials absorb at this wavelength, they can be used to quantitatively determine the content of the metals in a given organic sample by UV-Vis spectrophotometry. In Figure 30 the UV-Vis spectra of $L_1$ and the four complexes $C_1$, $C_2$, $C_3$ and $C_4$ is shown, and in Table 5 the determined extinction coefficients at $\lambda_{\text{max}}$ for the lowest electronic transition is given. The UV-Vis spectra of $L_2$ and $C_7$ are shown in paper 4 in appendix B. The UV-Vis spectra of $C_5$, $C_6$ and $C_8$ have not been recorded because it turned out that it was very difficult to separate the excess of $L_2$ from the synthesized complexes. Using the combination of $L_1$ and UV-Vis spectrophotometry it is possible in case of the Pd and Pt to determine the content of the metals with a detection limit of 0.1 ppm using 50 mg samples. A detection limit of 1 ppb can be obtained, when using larger samples and concentrating the supernatant.

![UV-Vis spectra](image)

**Figure 30.** The UV-Vis spectra of $L_1$ and the four electron transfer complexes $C_1$, $C_2$, $C_3$ and $C_4$. The spectra are recorded in chloroform at room temperature. The extinction coefficients are listed in Table 5.20
Table 5. The molar extinction coefficients and the $\lambda_{\text{max}}$ values of the first transition for C1, C2, C3 and C4. The values are determined both in chloroform and THF.20

<table>
<thead>
<tr>
<th>Compound</th>
<th>THF $\lambda_{\text{max}}$ [nm]</th>
<th>$\varepsilon$ [M$^{-1}$ cm$^{-1}$]</th>
<th>CHCl$<em>3$ $\lambda</em>{\text{max}}$ [nm]</th>
<th>$\varepsilon$ [M$^{-1}$ cm$^{-1}$]</th>
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</thead>
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<tr>
<td>C1</td>
<td>796</td>
<td>8500</td>
<td>801</td>
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<td>C2</td>
<td>727</td>
<td>26400</td>
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<td>25400</td>
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<td>C3</td>
<td>770</td>
<td>3900</td>
<td>768</td>
<td>3400</td>
</tr>
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<td>C4</td>
<td>830</td>
<td>1800</td>
<td>830</td>
<td>1800</td>
</tr>
</tbody>
</table>

7.2.2 Test of L1 and L2

In order to analyse the ability of L1 to remove transition metals from polymers poly(phenylenevinylene) (PPV), poly(phenyleneethynylenes) (PPE) (Na), and poly(alkylthiophene) (PAT) (all three were synthesised with a Pd catalyst) were tested by L1. In order to analyse the abilities of L2, it was tested on the diphenyl acetylene from the model systems in Scheme 16.

In the case of L2 the Pd and Cu contents in the model system were first determined by ICP-SFMS and by use of the combination of L1 and UV-Vis spectrophotometry. After treatment with L2 and filtration through a pad of silica, the Pd and Cu content in the model system were again determined by ICP-SFMS and by use of L1 and UV-Vis spectrophotometry. The results are listed in Table 6. The two different analytical methods give comparable results and have successfully brought the Pd and Cu contents in the model system under the parenteral limit by use of L2.

In the cases of L1 the content of Pd in the tree polymers was first determined by use of L1 and UV-Vis spectrophotometry. After treatment with L1, the content of Pd was again determined by use of L1 and UV-Vis spectrophotometry. The results are listed in Table 6. It was not possible to use ICP-SFMS to cross check the Pd content determined by L1, because the polymers cannot be fully dissolved in aqua regia.

Also in the cases of the polymers the content of Pd was brought down under the parenteral limit, and more relevantly the resistance of fabricated standard electroactive devices increased by more than a factor 100 and both LED and OPV devices could be prepared after the treatment with L1.

Table 6. The Pd and Cu content in model system (Scheme 16) before and after treatment with L2 and the Pd content before and after treatment with L1 for the polymer samples of the PPV, PPE and PAT type. Furthermore the resistance of standard electroactive devices made of the three polymers before and after the treatment with L1 is presented.20,28

| Content of transition metal [ppm] and device resistance [kΩ] (3 cm$^2$ device area were used) |
|-----------------------------------------------|-----------------------------------------------|
| Model system | Before treatment | Treated with L2 |
|               | UV-vis / ICP-SFMS | UV-vis / ICP-SFMS |
| Pd             | 376 / 610         | <0.1 / 0.5       |
| Cu             | 1074 / 1660       | <0.1 / 0.1       |
| Polymer (Pd)   | Before treatment | Treated with L1 |
|               | UV-vis / resistance | UV-vis / resistance |
| PPV            | 17860 / 0.1       | <0.1 / 30        |
| PPE            | 268 / 0.03        | <0.1 / 45        |
| PAT            | 4073 / 0.06       | <0.1 / 24        |
7.2.3 Structural determination

Before Hazell in 1976 discovered the unusual tetrahedral coordination of the C4 complex it was assumed that all bis-chelate electron transfer complexes had a planar coordination around the metal atom. In 1977 Bechgaard based on powder diffraction postulated that it was not only C4 that had the unusual tetrahedral coordination, but also the C3 complex forms the tetrahedral coordination.

To prove the postulate of Bechgaard and to determine the structures of the other complexes, it was attempted to grow crystals of all the complexes. Because of the short N,N-diethylamine group in L1 compared to the long N,N-bis(tetraethyleneglycolmonomethyl-ether)amine in L2 it was only possible to grow single crystals of the complexes C1, C2, C3 and C4.

Single crystals of C1 and C4 were made by slow evaporation of a dichloromethane solution, while single crystals of C3 were taken from the copper wire in Figure 28. It was very tricky to obtain crystals of C2, because slow evaporation of the most common solvents lead to twin crystals or needle clusters of crystals. However, by applying the sitting-drop vapor diffusion technique a few single crystals were successfully grown. The equipment for the sitting drop technique consists of glass sitting-drop rods placed in a stender dish in glass sealed with vacuum grease (see Figure 31). The condition was room temperature, a reservoir solution of 1 mL Toluene and 0.5 mL THF, a THF solution for the drop and a drop size of 40 μL.

The structure determination was carried out by single-crystal X-ray diffraction at a temperature of 118 K (further information about the experiments and a more detailed structure analysis can be found in paper 6 in appendix B).

From the structure determination it was established that both C3 and C4 crystallize in space group P4_1212 and had the unusual flattened tetrahedral coordination (see Figure 32). This proved that the postulate of Bechgaard was correct. The C1 and C2 complexes on the other hand crystallize in space group C2/c and show the conventional square planar coordination (see Figure 33).
Figure 32. The X-ray structure of C3. The structure of C4 is very similar and can be found in paper 6 in appendix B.26

Figure 33. The X-ray structure of C1. The structure of C2 is very similar and can be found in paper 6 in appendix B.26

Table 7. The key angles [°] for the four complexes. M is the metal ion.

<table>
<thead>
<tr>
<th></th>
<th>C1</th>
<th>C2</th>
<th>C3</th>
<th>C4</th>
</tr>
</thead>
<tbody>
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<td>S-M-S</td>
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<td>180.00(2)</td>
<td>123.62(2)</td>
<td>119.50(3)</td>
</tr>
<tr>
<td>N1-M-N1</td>
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<td>180.00(7)</td>
<td>144.84(7)</td>
<td>144.53(9)</td>
</tr>
<tr>
<td>N1-M-S</td>
<td>82.50(10)</td>
<td>82.11(7)</td>
<td>85.48(4)</td>
<td>85.62(4)</td>
</tr>
<tr>
<td>Plane(^a)</td>
<td>0.00(0.16)</td>
<td>0.00(0.10)</td>
<td>71.26 (3)</td>
<td>75.39(3)</td>
</tr>
<tr>
<td>Plane-phenyl(^b)</td>
<td>51.04(0.13)</td>
<td>52.01(8)</td>
<td>26.55(0.06)</td>
<td>28.08(0.07)</td>
</tr>
</tbody>
</table>

\(^a\) Angle between ligand planes.
\(^b\) Angle between ligand plane and phenyl group.
Table 8. The key distances (bond lengths) [Å] for the four complexes. M is the metal ion.

<table>
<thead>
<tr>
<th></th>
<th>C1</th>
<th>C2</th>
<th>C3</th>
<th>C4</th>
</tr>
</thead>
<tbody>
<tr>
<td>M-S</td>
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<td>C1-S</td>
<td>1.740(4)</td>
<td>1.743(2)</td>
<td>1.7234(15)</td>
<td>1.7226(17)</td>
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<tr>
<td>C1-N2</td>
<td>1.339(5)</td>
<td>1.326(3)</td>
<td>1.3688(18)</td>
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<td>1.339(5)</td>
<td>1.349(3)</td>
<td>1.3226(16)</td>
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<td>M-N1</td>
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</tbody>
</table>

In Table 7 and Table 8 some selected key angle and distances are listed. From the X-ray structure determination it is also possible to get some information about the electronic configuration of the four complexes. This is done on the basis of the work by Blanchard et al. who have demonstrated that the bond length depends on the given oxidation level of the ligand in complexes with 1-phenyl and 1,4-diphenyl substituted S-methylisothiosemicarbazides as the ligands. It is possible from the X-ray data to determine the plausible oxidation level of the metal and the ligand in these four complexes. Blanchard et al. have shown that in the case of a π-radical anion, (L') the N-N bond is rather short (1.34 ±0.01 Å), but if the bond lengths are significantly shorter (1.24 ±0.01 Å), then the ligand is regarded as being in its oxidized level, (L ox). From the data in Table 8 it can be seen that the N-N bond lengths in C1, C2 and C4 complexes all are in the order of 1.34 ± 0.01 Å, and therefore it can be rationalized as M(II) complexes with two π-radical anion ligands antiferromagnetically coupled, [PdII(L•)2], [PtII(L•)2] and [NiII(L•)2].

It is also possible to get information about the delocalization of the π electron density in the complexes. Blanchard et al. have calculated the values of the following bond lengths N=C, C-S, C-N, N-N to 1.294 Å, 1.897 Å, 1.342 Å and 1.343 Å respectively, in the large localized system of (HN-N=C(SH)NH •)1-. Furthermore, the average C-S and C-N distances in the highly delocalized system of dithiocarbamate are 1.70 and 1.34 Å. By comparing these bond lengths with the bond lengths in the five-membered ring of the complexes described in this work, a large delocalization of the π electron density over the five-membered ring is suggested.

Concerning the C3 complex, it can be seen that it has a slightly smaller N-N bond length (1.32 Å) and a slightly larger C-N bond length compared to the other three complexes. This can be rationalized if it is assumed that the C3 complex consists of Cu(I), one π-radical anion ligand and one neutral ligand, [CuI(L•)(L)] as proposed by Bechgaard. However, which of these two configurations, [CuII(L•)2] (antiferromagnetically coupled) or [CuI(L•)(L)], that is the most likely for the C3 (Cu(L)2) complex, cannot be decided on the X-ray data alone.

7.3 Conclusion
An efficient, fast, very mild and potentially low cost (no expensive chemicals) method to remove remnants of transition metals from both polymer and small molecule products to a limit under 0.1 ppm, has been developed. This opens up the opportunity for the industry to also use palladium catalysts in the last synthetic steps of the synthesis of e.g. drugs.
Furthermore, the method gives the possibilities to quantitatively determine the content of transition metals at 1 ppb level in a given organic sample. The discovery of the azothioformamide properties have together with Sigma Aldrich's new but highly expensive metal scavenger products made remnants of Pd in newly synthesised compounds into a saga. There are just no more excuses to have remnants of Pd or other transition metals in your compounds.

It can also be concluded that the coordination of the formed neutral electron complexes depend on the metal and can be either a flattened tetrahedral coordination or a square-planar coordination. Furthermore the formed neutral complexes most likely consist of the metal in oxidization state II and two \( \pi \)-radical anion ligands or of the metal in oxidization state I together with one \( \pi \)-radical anion and one neutral ligand.

References

5. Schmuckler, G. (Technion Research and Development Foundation Ltd.), DE1533131, 1974
13. Private communication with Lizette Bruun (employment at RISØ and former employment at Lundbech A/S, Denmark).
The Palladium Saga

The synthesis of monodisperse oligomers of the PPE type was unsuccessful as mentioned in chapter 5. It has therefore not been possible to make crystals of the PPE and determine the structures by single crystal X-ray diffraction measurement, as I originally set out to do. Instead it has been attempted to grow single crystals of four monodisperse phenylene vinylene oligomers. The oligomers have been synthesized by Jørgensen\textsuperscript{1} and the structures can be seen in Scheme 19.

\begin{equation}
\text{Scheme 19. The four monodisperse phenylene vinylene oligomers synthesized by Jørgensen.}\textsuperscript{1}
\end{equation}
To ensure that they are all crystalline, before any attempts to grow crystals of the four monodisperse oligomers were done, X-ray powder diffraction patterns were recorded of all four oligomers (see Figure 34 and Figure 35).

**Figure 34.** The X-ray powder diffraction pattern of O1 (top) and O2 (bottom).
Figure 35. The X-ray powder diffraction pattern of O3 (top) and O4 (bottom).

From the X-ray diffraction patterns of the oligomers it can be seen that both O2 and O4 are completely amorphous, and no crystals were attempted grown of these two oligomers. O1 and O3 on the other hand contain some level of crystalline order, which is clearer from Figure 36 and Figure 37, where the background subtracted. It looks like the samples of O1 and O3 are micro crystalline.
Figure 36. The X-ray powder diffraction pattern of O1, where the background is subtracted. It can be seen that the oligomer contain some degree of crystalline order. The data is recorded over a period of 20 minutes.

Figure 37. The X-ray powder diffraction pattern of O3, where the background is subtracted. It can be seen that the oligomer contain some degree of crystalline order. The data is recorded over a period of 17 hours.

By comparing the diffraction patterns of O3 and O1 it can be seen that they are very similar regarding the positions of the peaks but not intensity of the peaks. In the pattern of O3 (Figure
peaks one observed at $2\theta = 6.6^\circ, 10.3^\circ, 18.5^\circ, 21.0^\circ, 23.5^\circ, 28.1^\circ$ and $42.2^\circ$, which corresponds to lattice distances (d-spacings) of 13.5, 8.6, 4.8, 4.2, 3.8, 3.2 and 2.1 Å. The quality of the data is however not good enough for unit cell determinations. On the basis of the diffraction pattern of O1 and O3 it was decided to attempt growth of crystals of these two oligomers.

From the literature it was found that some oligo-phenylvinylene has been crystallized before (see Scheme 20) and that it has been done by slow evaporation from solvents such as $n$-hexane, chloroform or chloroform/$n$-hexane mixtures.

I have tried to crystallize O1 and O3 in solvents such as: $n$-hexane, chloroform, toluene, THF, dichloromethane, ethanol, ethyl acetate and a lot of different mixtures of these solvents. Furthermore I have tried to use two different crystallization techniques, which are: Slow evaporation from a flask and the sitting-drop vapor diffusion technique (see figure 31 in chapter 7) but without any luck. Lots of strange and fancy structures were obtained under the different crystallization attempts (see Figure 38 and Figure 39), but none of them gave single crystals or crystals. Only powder patterns could be observed, when the structures were tested in a diffractometer. Not even the structures obtained from O3 (see Figure 39 - right) which in the microscope really looks like crystals with nice sharp edges and shiny surfaces, gave diffraction spots in the diffractometer.

**Scheme 20.** oligo-Phenylvinylene, which have been crystallized previously.
Figure 39. Some of the amorphous/micro crystalline products of O3 obtained by the sitting-drop vapor diffusion techniques.

As a last desperate attempt glass equipment shown in Figure 40 was designed to make it possible to try to crystallize O1 by using stilbene as the solvent. This was inspired by van Hutten et al. who have used a similar method to obtain a single crystal of unsubstituted phenylenevinylene. The idea is that O1 and stilbene (2:100 (w/w)) is placed in the round bottom flask, and the glass equipment is put together. The glass equipment containing the O1 and stilbene is placed in an oven and heated up to 558 K and shaken gently (be careful the equipment is very hot, which the author has learned from bitter experience) until all compounds is in the liquid form and well mixed. The temperature is now lowered down to 518 K and is kept at this temperature until crystals of O1 precipitated (in the case of van Hutten et al. this happened within 2 hours). The precipitated crystals could then be isolated by tipping the equipment (still inside the oven) and filtering of the hot "solvent" through the glass filter. In my case, I waited 6 hours at 518 K for precipitation of crystals of O1 precipitated (in the case of van Hutten et al. this happened within 2 hours). The precipitated crystals could then be isolated by tipping the equipment (still inside the oven) and filtering of the hot "solvent" through the glass filter. In my case, I waited 6 hours at 518 K for precipitation of crystals of O1 precipitated (in the case of van Hutten et al. this happened within 2 hours). I then decided to try to filter the solvent through the filter to test the equipment. After the filtration the equipment was left to cool down and the round bottom flask was attempted to separate from the part containing the filter. It turned out that the two parts was stuck together, and could not be separated without breaking the equipment. No further attempts to crystallize O1 and O3 were performed.

Figure 40. The glass equipment developed for crystallizing O1 from stilbene.
The conclusion is that it was not in my hands possible to grow single crystals or any crystals of the monodisperse oligomers. Why it succeeded for people in the literature to grow crystals of the oligomers in Scheme 20, I do not know. Maybe it is due to fact that the oligomers in Scheme 20 generally are shorter, and in the cases of the long oligomers, there are fewer side groups on the backbone compared to the oligomers I have tried. It is well known that the length and the numbers of the side groups compared to the length of the backbone have a large influence on the packing and crystallizing abilities of molecules.7

References

7 Pedersen, W. B., employed at Risø - personal communication.
In this chapter the characterization of N\textsubscript{n} and the NPN polymers is described. The characterization covers the electronic energy levels and the charge carrier motilities of N\textsubscript{n}, the energy transfer from the N domain to the P domain in the NPN polymer, and tests of LED and POPV devices made of both polymers. All the experimental data concerning the characterizations are presented in paper 1, 2 and 5 in appendix B.

**9.1 The electronic levels of N\textsubscript{n}**

In chapter 2.2.1 it was shown that knowledge of the position of the electronic energy levels in the polymers is important because the position of these levels compared to the levels of the electrodes determines the injection barriers into the valence band and the conduction band for holes and electrons respectively. From Ultra Photoelectron Spectroscopy (UPS) experiments\textsuperscript{1} it is possible to get information on the position of the valence band (E\textsubscript{VB}) and to determine the ionization potential (IP). In the UPS experiment hard UV photons with an energy of 50 eV from a synchrotron source were used. The UV photons interact with the atoms in the polymer surface by the photoelectric effect, causing electrons from one of the filled energy levels in the polymer to be emitted. The kinetic energies of the emitted electrons are given by equation (9.1).

\[
E_{\text{kin}} = h\nu - BE - \phi_s
\]

In equation (9.1) \(h\nu\) is the energy of the UV photons (50 eV), \(BE\) is the binding energy of the emitted electrons, which depends on the given atomic orbital. \(BE\) can be regarded as the ionization energy of a particular shell for a certain atom. \(\phi_s\) is the workfunction of the spectrophotometer. In the UPS experiment a negative bias of \(\sim 9\) V is applied on the sample to ensure that electrons are not trapped by the workfunction of the spectrophotometer.

The UPS can therefore give information on the position of the valence band, while it is not possible to get information on the position of the conduction band because of the missing interaction with the unfilled energy levels.

In Figure 41 the recorded UPS spectrum of N\textsubscript{n} is shown together with an electronic energy band edge diagram derived from the UPS spectrum.\textsuperscript{2}

The Fermi level, \(E_F\), in a thin polymer film placed on a metal substrate has been shown to be determined by the Fermi level of the metal substrate.\textsuperscript{1} From a clean gold substrate the position of the Fermi level, \(E_F\), can be determined by UPS and is given by onset where photoelectrons become detectable. When \(E_F\) is determined it is used as the zero binding energy reference level (per definition \(E_F\) is the zero binding energy) (see Figure 41).
Characterization

Figure 41. Left: The UPS spectrum of an over-layer of N\textsubscript{n} on a gold substrate with an insert (×10) showing the onset of photoelectrons. The difference between the Fermi level, $E_F$, and the onset of the photoelectrons represents the injection barrier for holes from the gold substrate into the valence band of N\textsubscript{n}. Right: The electronic energy band edge diagram of N\textsubscript{n} derived from the UPS spectrum together with a picture of the gold substrate plate (the right corner).\textsuperscript{34}

It is also possible by UPS to determine the workfunction of gold, $\Phi_\text{Au}$, from the clean gold substrate. $\Phi_\text{Au}$ is determined as the ionization potential, which simply is the difference between the energy of the UV photons (50 eV) and the numerical value of secondary electron cutoff at high binding energies, $BE_{\text{max}}$ (see equation (9.2)).\textsuperscript{5}

$$\phi_{\text{Au}} = h\nu - |BE_{\text{max}}| \quad (9.2)$$

The position of the valence band edge for the polymer material is the difference in kinetic energy of the fastest photoelectrons from the gold substrate compared to the fastest electrons in the thin polymer film on top of the gold substrate. In practice this difference, $E_{F_{\text{VB}}}$, is determined as the onset of the photoelectrons towards low binding energies, $BE_{\text{min}}$, with reference to the Fermi level of the gold substrate (see Figure 41 and equation (9.3)). The $E_{F_{\text{VB}}}$ represents the injection barrier for holes from the metal substrate into the valence band of the polymer.\textsuperscript{1,5}

$$E_{F_{\text{VB}}} = BE_{\text{min}} \quad (9.3)$$

The difference between the vacuum level of the polymer and the Fermi level of the metal substrate, $E_{F_{\text{VAC}}}$, is given by equation (9.4):

$$E_{F_{\text{VAC}}} = h\nu - |BE_{\text{max}}| \quad (9.4)$$

The ionization potential, $IP$, for the polymer is the sum of $E_{F_{\text{VB}}}$ and $E_{F_{\text{VAC}}}$ (see equation (9.5)), which is also illustrated in Figure 41. The $IP$ is a material constant and is independent of the alignment of the energy level at the interface.\textsuperscript{1}

$$IP = E_{F_{\text{VB}}} + E_{F_{\text{VAC}}} \quad (9.5)$$

The offset of the vacuum levels for a metal substrate with thin polymer film on top is shifted compared to the vacuum level of the pure metal substrate (see equation (9.6)).
The difference in offsets of the two vacuum levels, $\Delta$, is probably due to the presence of a dipole layer at the surface because of an orientation of molecules near or at the surface. The size and the direction of the dipole moment can determine whether it is easier or more difficult for electrons to escape the surfaces.$^{5,6}$

**Table 9.** The UPS characterization for the $N_n$ polymer on a gold substrate ($\Phi_{Au} = 4.7$ eV). All the units are eV. The UPS data for the equivalent PPV (also with octyl chains on every second benzene ring) are placed for comparison. It should be mentioned that the data of the PPV are recorded on an Ag substrate plate ($\Phi_{Ag} = 4.2$ eV) and not Au.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$E_{F}^{VB}$</th>
<th>$E_{F}^{VAC}$</th>
<th>BE$_{max}$</th>
<th>$\Delta$</th>
<th>IP</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_n$</td>
<td>1.5</td>
<td>4.0</td>
<td>46.0</td>
<td>-0.7</td>
<td>5.5</td>
</tr>
<tr>
<td>PPV$^{7,8}$</td>
<td>0.70</td>
<td>4.65</td>
<td>45.35</td>
<td>0.45</td>
<td>5.35</td>
</tr>
</tbody>
</table>

From the UPS characterization of $N_n$ (see Table 9) it can be seen that the injection barrier for the holes in the metal substrate to the valence band of $N_n$ is 1.5 eV, which means that a bias of at least 1.5 V needs to be applied, if the injection barrier for holes in a potential device has to be overcome. There is also an injection barrier for electrons from the metal substrate to the conduction band of $N_n$, but this cannot be determined by UPS.

By comparing the UPS data for $N_n$ with the UPS data for the corresponding PPV, it can be seen that the largest difference is between the injection barriers of holes, which is twice as large in $N_n$ compared to the PPV. The fact that the IP and the optical band gap (obtained from the UV-Vis spectra) are nearly the same in both polymers, indicates that the difference in value of $E_{F}^{VB}$ and $E_{F}^{VAC}$ reflects that the Fermi level in $N_n$ is placed closer to the conducting band than in the PPV.

### 9.2 Charge carrier mobilities in $N_n$

The charge carrier mobility is an important property for conjugated polymers employed for LED and POPV devices. To ensure a high intensity of electroluminescence in LED devices and a good power conversion in POPV devices it is desirable that the carrier mobility is high, because the higher the carrier mobility the more efficient the devices are.

The charge carrier mobility of $N_n$ as determined by use of the Pulse Radiolysis Time-Resolved Microwave Conductivity technique (PR-TRMC). The technique is well described in the literature$^{7,9}$ and will therefore only be presented very shortly here. PR-TRMC is based on the measurement of the change in the reflected microwave power from a polymer sample influenced by a short pulse of 10 MeV electrons. The high energy electrons create ionisation events whereby electron-hole pairs are formed, leading to a large increase in conductivity. The change in conductivity, $\Delta \sigma$, has been shown by Warman et al. to be directly related to the change in ratio between the change in the reflected microwave power, $\Delta P$, and the reflected microwave power before the electron pulse, $P_r$, see equation (9.7).

$$\left(\frac{\Delta P}{P_r}\right)_{max} = -A \Delta \sigma$$

(9.7)
The parameter \( A \) covers interference effects from the similarity between the dimensions of the sample and the wavelength of the incoming microwaves, and can be calculated from the knowledge of the sample length and the effective dielectric constant.\(^7,9\) Information on the sum of charge carrier mobilities is related to the change in the conductivity, see equation (9.8).

\[
\sum \mu_{\text{min}} = \frac{\Delta \sigma}{e(N_+ + N_-)} \tag{9.8}
\]

\( N \) is the concentration of the carriers and can be determined from dosimetry, see equation (9.9).

\[
N_+ + N_- = \frac{D}{E_p} \tag{9.9}
\]

\( D \) is the measured dose and \( E_p \) is the electron-hole pair formation energy, where a value of 25 eV is normally used.\(^7\)

There is a major advantage and a major disadvantage of this method. The advantage is that you measure the average minimum carrier mobilities and it is done without influence of any electric contacts. The disadvantage is that you can only measure the sum, and it is not possible to distinguish between the electron and the hole mobilities.

From the transient changes in the conductivity the lifetime of the carriers can be determined by a biexponential fit.

In Figure 42 the plot of the electron pulse (duration \( \sim 300 \) ns) and the corresponding dose normalized conductivity transient is shown (normalized after the dose from the pulse).

**Figure 42.** Above: The 300 ns pulse. Below: The dose normalized radiation induced change in conductivity plotted as a function of time after the pulse. The value at the end of pulse conductivity used for the determination of the sum of the mobilities is read 100 ns post-pulse and is \( 5 \times 10^{-8} \) s m\(^2\) J\(^{-1}\).\(^3\)
The determined sum of charge carrier motilities and values of the first half life time are given in Table 10 together with values from different PPVs (see Scheme 21).

**Table 10.** Comparison of minimum sum of carrier mobilities, \( \sum \mu_{\text{min}} \), and the first half life time, \( \tau_{\frac{1}{2}} \), of \( N_n \) and some PPVs.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>( \sum \mu_{\text{min}} ) [m(^2) V(^{-1}) s(^{-1})]</th>
<th>( \tau_{\frac{1}{2}} ) [( \mu )s]</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N_n )</td>
<td>( 1.25 \times 10^{-6} )</td>
<td>10</td>
</tr>
<tr>
<td>MEH-PPV (^a)</td>
<td>( 2.00 \times 10^{-6} )</td>
<td>4</td>
</tr>
<tr>
<td>PPV1 (^a)</td>
<td>( 8.90 \times 10^{-6} )</td>
<td>528</td>
</tr>
<tr>
<td>PPV2 (^a)</td>
<td>( 2.32 \times 10^{-6} )</td>
<td>1</td>
</tr>
<tr>
<td>PPV3 (^a)</td>
<td>( 1.04 \times 10^{-6} )</td>
<td>2</td>
</tr>
<tr>
<td>PPV4 (^a)</td>
<td>( 1.86 \times 10^{-6} )</td>
<td>4</td>
</tr>
</tbody>
</table>

\(^a\) data from the literature.10

From Table 10 it can be seen that the \( \sum \mu_{\text{min}} \) generally of similar magnitude, but the value for the PPV1, which only consists of carbon and hydrogen, is somewhat larger than the others. It seems to be the case that electro negative atoms like fluorine and oxygen decrease the mobilities slightly. The significantly larger value of \( \tau_{\frac{1}{2}} \) in PPV1 has been shown most likely to be due the higher purification level.10 It can be seen that the choice of a more rigid system, \( N_n \) (containing an extra set of \( \pi \) electrons), have not, as hoped, lead to the increased charge carrier mobility compared to the PPVs. At least the charge carrier mobilities and the half-lives are comparable with the values found in PPVs.

**Scheme 21.**

From Table 10 it can be seen that the \( \sum \mu_{\text{min}} \) generally of similar magnitude, but the value for the PPV1, which only consists of carbon and hydrogen, is somewhat larger than the others. It seems to be the case that electro negative atoms like fluorine and oxygen decrease the mobilities slightly. The significantly larger value of \( \tau_{\frac{1}{2}} \) in PPV1 has been shown most likely to be due the higher purification level.10 It can be seen that the choice of a more rigid system, \( N_n \) (containing an extra set of \( \pi \) electrons), have not, as hoped, lead to the increased charge carrier mobility compared to the PPVs. At least the charge carrier mobilities and the half-lives are comparable with the values found in PPVs.

**9.3 UV-Vis characterization**

The UV-Vis spectra of \( N_n \) have been recorded both in solution and in the solid state (see Figure 43), and the narrow band and sharp absorption edges indicated that the polymer has a rigid linear structure as expected. The form of the bands in both solution and in solid state is similar to similar compounds presented in the literature.11 From the UV-Vis spectra it can also be seen that the optical band gap in solution is 3.0 eV and in the thin film it is 2.8 eV. The \( \lambda_{\text{max}} \) of \( N_n \) in solution and solid state is 397 nm and 424 nm respectively, which as expected lies a somewhat above the values of PPEs with electron withdrawing side groups as ester groups and a little under the values of PPEs with electron releasing side groups such as ether
Characterization

groups (see Table 11). It can also be seen that $\lambda_{\text{max}}$ is red shifted about 27 nm in the solid state compared to the solution state, which is an indication of intermolecular interaction in the solid state.

The UV-Vis spectra of the NPN B fractions have been recorded both in solution and in the solid state (see Figure 26 and Figure 44). It can be seen that in both states all the bands are red shifted as a function of increasing chain length, which is a result of increasing conjugation length indicating that the porphyrin is incorporated in the backbone of a conjugated system. The red shift stops at fraction 4, which from the UV-Vis signal corresponds to the hexamer of the $N_n$ chains. This result is in agreement with the results of Jones et al.$^{12}$ who reported studies on oligomers at precise length of the 1,4-phenylene ethynylenes.

![Graph](image)

**Figure 43.** The UV-Vis spectra of $N_n$ in solution and in solid state.

As like in the case of $N_n$ the form of the absorption band for NPN B is characterized by being narrow and having sharp absorption edges indicating that NPN B also has a rigid linear structure. The $\lambda_{\text{max}}$ of NPN B in solution and solid state is 386 nm and 393 nm, respectively. The lower value of $\lambda_{\text{max}}$ compared to $N_n$ could be a result of the porphyrin working as an electron withdrawing group. The 7 nm red shifting of $\lambda_{\text{max}}$ in the solid state compared to the solution state indicate moderate intermolecular interaction in the solid state.

The $\lambda_{\text{max}}$ values for different types of polyphenylene ethynylenes can be found in Table 11.
Characterization

Figure 44. The UV-Vis spectra of thin spin coated film of the NPN B fractions.4

Table 11. $\lambda_{\text{max}}$ values of selected PPEs in solution and in thin film. The $\lambda_{\text{max}}$ values for the solution are found in THF, except in the case of $N_n$ and NPN B where it was determined in CHCl$_3$.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>$\lambda_{\text{max}}$ [nm] solution</th>
<th>$\lambda_{\text{max}}$ [nm] thin film</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_n$</td>
<td>397</td>
<td>424</td>
</tr>
<tr>
<td>NPN B</td>
<td>386</td>
<td>393</td>
</tr>
<tr>
<td>PPE1$^{13}$</td>
<td>388</td>
<td>-</td>
</tr>
<tr>
<td>PPE2$^{11}$</td>
<td>414</td>
<td>452</td>
</tr>
<tr>
<td>PPE3$^{14}$</td>
<td>383</td>
<td>405</td>
</tr>
<tr>
<td>PPE4$^{14}$</td>
<td>408</td>
<td>412</td>
</tr>
</tbody>
</table>
9.4 Fluorescence characterization

The primary reason for the synthesis of NPN B was to make a molecular heterojunction where dissociation of formed excitons, in the case of POPV devices, or a recombination of charges or excitons under emitting of light, in the case of LED devices, could happen. It is possible to make full colour displays of polymer based LEDs, because all three principle colours: Blue, green and red is possible to obtain. Among the principle colours, green is the most developed concerning brightness and efficiency, while there still is relatively few reports concerning LEDs, which emits light in the near infrared (NIR) region. The reports concerning NIR LEDs can be divided into three groups, which are: Low band gap polymers like poly(thiophenes),\textsuperscript{15} polymers doped by rare earth metal ions\textsuperscript{16} and polymer blends where the hosts consist of conjugated polymers with high band gap, which are doped by small red-emitting organic dyes such as porphyrins.\textsuperscript{17} The polymer blends are the most frequently used method to obtain NIR LEDs and the working mechanism is that the host polymers take care of the charge and exciton transport to the dye molecule, where the excitons decays by emission of NIR light. The exciton transport from the host polymer to the dye molecule occurs through space (TS) normally by a Förster energy transfer process.\textsuperscript{18} The Förster energy transfer process is a long-range dipole-dipole coupling process, where the transition dipoles of the excited polymer interact with the transition dipoles of the ground-state of the dye molecule.\textsuperscript{19} The Förster energy transfer process is limited by the fact that it normally only occurs in the distance range of 1 to 50 Å\textsuperscript{20} and that in addition it requires a good spectral overlap between the emission spectra of the polymer host and the absorption spectra of the dye molecule. Good spectral overlap is far from the general observation.\textsubscript{21,22,23,24,25} To overcome the distance range limitation in the polymer blends, the dye molecule can be covalently linked to the polymer as in the case of NPN B, which will ensure that the dye molecules always are well mixed in the polymer host and therefore always are inside the range of 50 Å. The covalent link opens furthermore up for the possibility of energy transfer through bonds (TB),\textsuperscript{26,27,28} which is promoted by an orbital overlap, do not require spectral overlap and is much faster than the TS transfer process.\textsuperscript{19,21,22} Figure 45 illustrate the TB transfer process contra the TS transfer process.

![Energy Transfer Processes](image)

**Figure 45.** The energy transfer processes in polymer blends contra molecules like NPN B containing molecular heterojunction.\textsuperscript{9}

In order to characterize and test the efficiency of the energy transfer processes in molecules like NPN B, emission spectrophotometry have proven to be a very useful tool. All experimental details about the fluorescence analysis can be found in paper 1, 2 and 5 in appendix B.
Solution of $N_n$ and NPN B in chloroform was placed under an UV lamp as a first check to see if $N_n$ and NPN B display fluorescence and to see if there is a difference in their fluorescence, which could indicate an energy transfer from the N domains to the P domain in the NPN B structure. In Figure 46 it can be seen that the displayed fluorescence is deep blue for the two structures, and not so intense in the case of the NPN B, which indicate that an energy transfer from the N domain to the P domain takes place. However the difference is not as significant as anticipated.

![Figure 46. The fluorescence of the $N_n$ (right) and the NPN B (left) molecules in solution when placed under an UV lamp.](image)

In Figure 47 the fluorescence spectra of $N_n$ in solution (CHCl$_3$) and in thin film are shown. The form and position of the band is similar to reported values for similar PPEs.$^{11,13,29,30,31,32,33}$ The fluorescence spectrum of $N_n$ in diluted chloroform solution is characterized by a sharp peak with $\lambda_{\text{max}} = 423$ nm and a broader shoulder with $\lambda = 443$ nm. By comparing with other dialkyl PPEs$^{29}$ it can be seen that the peak maximum is in the range of $\lambda_{\text{max}} = 428$ to 435 nm and the shoulder is in the range of $\lambda = 440$ to 450 nm, which is consistent with my observations. By comparing the fluorescence spectrum of $N_n$ in Figure 47 with the absorption spectrum in Figure 43 it can be seen that no mirror imaging is observed, which indicate that the exited state of $N_n$ have a different geometric arrangement of the nuclei than in the ground state. Further it can be seen that the fluorescence spectrum show more fine structure than the absorption spectrum in solution, which is probably due to the benzene rings in the exited state becoming more co-planar. On model compounds of dialkoxy PPEs$^{11}$ the intensity of the shoulder has been seen to be concentration dependent, which indicate that the shoulder is due to excimer formation. The fluorescence quantum yield for $N_n$ in the chloroform solution is determined to be 0.86 in reference to diphenyl anthracene, DPA. This is somewhat bit low compared to the general value of dialkyl PPEs, which is unity due to the rigid structure, which makes intermolecular dissipation difficult.$^{29}$ The low value is probably due to the content of small amounts of impurities, which are able to quench the fluorescence. This assumption is supported by the fact that the low half-life times in the PR-TRMC analysis of $N_n$ compared to PPV1 also indicate the presence of small amounts of impurities in $N_n$. 

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Figure 47. The absorption spectra of the dibromoporphyrin and the fluorescence spectra of Nₙ in both solid state and in solution. The excitation wavelengths are 397 and 424 nm in the case of solution and solid state respectively.

In the solid state the fluorescence spectrum of Nₙ is strongly red shifted and is characterized by a broad peak with $\lambda_{\text{max}} = 512$ nm and a shoulder at $\lambda = 490$ nm. In the solid state a lack of the mirror symmetry is also observed, and further the broad unstructured emission centered at $\lambda_{\text{max}} = 512$ nm is probably due to excimer formation in the solid state similar to the solution case. The much broader emission band in the solid state compared to the solution indicate that the formation of excimer clusters is more dominating in the solid state compared to solution.

From Figure 47 it can also be seen that there is a good spectral overlap between the absorption of the porphyrin and the emission of the Nₙ polymer in both solution and in solid state. This means that both TS and TB energy transfer processes can be involved in energy transfer from the N to the P domain in NPN B.

In Figure 48 the fluorescence spectrum of the dibromoporphyrin in chloroform is shown. The spectrum is consists of two peaks located at 612 and 662 nm. The quantum yield in reference to DPA was determined to be 0.11.
Figure 48. The fluorescence spectrum of the dibromoporphyrin in chloroform. The excitation wavelength is 426 nm.

Figure 49. Normalized fluorescence spectra of the seven fractions (1-7) of NPN B in degassed chloroform and at a concentration below $10^{-5}$ M. The insert shows the porphyrin emission of fractions 2, 4 and 6. The excitation wavelength is 397 nm.

The normalized fluorescence spectra of the seven fractions of NPN B in chloroform are shown in Figure 49. The excitation wavelength is set to $\lambda_{\text{max}}$ of N$_n$ (397 nm), and from Figure 47 and Figure 48 it can be seen that the emission below 600 nm is mainly due to the N domain and emission above 600 nm is due to the P domain. It can be seen that the emission is
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slightly red shifted as a function of the chain length as expected, but generally there is very little variation in the spectra as a function of the chain length with fraction 6 and 7 as the exceptions probably due to the content of relatively short chain segments. The most interesting and most worrying observation is that there seems to be a very limited energy transfer from the N domain to the P domain in solution. The overall quantum yield from NPN B is around 0.6 and is mainly caused by the N domain. From the measured fluorescence emission and from the assumption that the ratio between the fluorescence rate and the nonradiative decay rates are the same in the NPN B as in the pure porphyrin and \( \text{N}_n \) compounds a photo balance can be calculated (see Figure 50).

![Figure 50](image)

**Figure 50.** The photo balance of the NPN B structure in diluted degassed chloroform solution. The absorption is set to 1000 photons and the fluorescence emission was measured, while the energy transfer and nonradiative decay is calculated from the fluorescence quantum yields of the pure porphyrin and \( \text{N}_n \) compounds.\(^{34}\)

The only positive result of this photo balance is, that the energy transfer is slightly better than the 10,20-diphenyl-5,15-bis(4-(poly-2',5'-dioctyl-4,4''-terphenylene-1-cyanovinylene-2-yl)-phenyl)porphyrin (JPJ) three domain structure of Krebs et al (see Scheme 22 and Figure 51).\(^{35}\)

![Scheme 22](image)

**Scheme 22.** The JPJ three domain structure synthesized by Krebs et al.\(^{35}\)
One last unexpected piece of information can be drawn from Figure 49 and Figure 50, where it can be seen that the emission of the P domain increases, when the chain length of the N domain decreases. The reason is probably that because of the cross linking of the NPN B under the purification by SEC (see chapter 6), the N domains are not perfectly conjugated, but consist of a number of segments. Excitons can therefore be trapped in the segments in the N domain. Thus by increasing the length of the N domain the probability of an exciton trapped on a segment that is not in the vicinity of the P domain increase, and therefore the efficiency of the energy transfer will decrease.

In Figure 52 the fluorescence spectra in the solid state of the pure dibromoporphyrin, Nₐ and fraction 2 to 7 of NPN B are presented. The fluorescence spectra of the fractions of NPN B are all characterized by a very intense and sharp red emission peak with λ_max about 695 nm, a weak and broader shoulder about 750 nm and a weak and broad emission in the range of 400 to 550 nm. This emission in the red area is due to the P domain, while the emission in the range of 400 to 550 nm is due to the N domain.

By comparing the fluorescence spectra of the NPN B fractions with the fluorescence spectra of the pure domains, it can be seen that the emission of the N domain is nearly quenched 100%, which indicate a very effective energy transfer from the N domains to the P domain. The effectiveness emphasizes the fact that PPEs normally have a high quantum yield and is often near unity. Only in fraction 2 and 3 (the two fractions with the longest N domains) a weak emission of the N domain can be observed. The weak emission from the N domains in fraction 2 and 3 are probably due to the excitons trapped in segments far from the P domains like in the solution cases due to cross-linking. A larger out of plane torsion angle would reduce the conjugation length and thereby also reduce TB energy transfer. This could explain why the energy transfer is much more efficient in the solid state compared to solution (see Figure 49 and Figure 52), which probably is due to the torsion angle between the N and P domains are much more out of plane in solution compared to solid state as a results of steric hindrance. This is supported of the fact that it is known that porphyrin units preferentially are stacked in solid state. The much higher energy transfer in solid state compared to
solution is reported for similar systems in the literature,\textsuperscript{22,35,36} and indicate that TB energy transfer has a large contribution to the overall energy transfer in these domain structures.

![Fluorescence spectra](image)

**Figure 52.** The fluorescence spectra in solid state of the dibromoporphyrin, \(N_n\) and fraction 2 to 7 of NPN B.\textsuperscript{4} The excitation wavelength is 424 nm in the case of \(N_n\) and NPN B and 436 nm in the case of the dibromoporphyrin.

### 9.5 LED devices

Single-layer LED devices with the configuration as shown in Figure 14 in chapter 2 have been made of \(N_n\) and NPN B (further details can be found in paper 2 and 5 in appendix B). Both types of devices had low turn on voltages of 3.3 V. As a comparison the turn on voltages of phenylene ethynylene pentamers,\textsuperscript{32} 2,5-dialkoxy substituted poly\((p\)-phenylene ethynylene\) derivates,\textsuperscript{37} derivates of poly(thiophenes),\textsuperscript{15} 2,5-alkoxy substituted poly(phenylenes)\textsuperscript{17} and other poly\((p\)-phenyleneethynlenes)\textsuperscript{38} are reported to be 16 V, 8.5-15 V, 1.4-7 V, 4V and 5.5 V, respectively. The devices of \(N_n\) and NPN B were during electroluminescence measurements operated at 5.5 V with current densities in the range of 3-8 mA cm\(^{-2}\). The measured electroluminescence spectra are shown in Figure 53.

The electroluminescence of \(N_n\) is greenish blue light and consists of a broad peak, which is red shifted \(\sim 100\) nm compared to the fluorescence spectrum (Figure 52). The large red shift indicates that the emission from the N domain is due to a relaxed state. Furthermore, no fine structure is observed as in the case of the fluorescence spectrum. In Figure 54 the band diagram of \(N_n\) is shown. It can be seen that there is a large mismatch in energy for the hole injection, which maybe is a limitation factor for the effectiveness of the device.
The electroluminescence spectra of $N_n$ and NPN B.

The electroluminescence of NPN B could not be detected visually but measurements revealed that the emission is in the NIR region and that it is very similar to the solid state fluorescence spectrum of NPN B. I interpret this in such a way that the N domains work as very effective antennas, which are capable of transporting electrically generated excitons in the N domain to the P domain where in a photon is emitted as if it was ordinary fluorescence.

The band diagram of $N_n$. The HOMO and LUMO levels are determined by UPS and UV-Vis spectroscopy and the values of the electrodes are from the literature.
9.6 POPV devices

Despite a large and still increasing interest towards the unique photophysical and semiconducting properties of conjugated polymers, I have not been able to find one publication concerning the characteristics of POPV devices where the active layer consist of only a PPE derivative. However, PPE-C$_{60}$ solar cells have been reported. Furthermore, there are only very few publications concerning the POPV devices with an active layer consisting of polymers in where the monomer is a co-monomer between phenyl ethynylene and electron acceptor compound such as benzothiadiazole (BT). The missing interest concerning PPEs as the active layer in POPV devices can either mean that no one have discovered the unique properties of PPEs or more likely that the PPEs do not work very well as the active layer in POPV devices. The last is the most probable, which next paragraph will reveal.

POPV devices of $N_n$ and NPN B have been made as described in chapter 2.3 and dark and photo I/V characteristics have been measured (see paper 5 in appendix B for experimental details). The measured I/V characteristics are present in Figure 55 and Figure 56 as linear and logarithmic plots respectively. Data about the fill factor (FF), the power conversion efficiency ($\eta_{\text{eff}}$), the short-circuit current ($I_{\text{SC}}$), the open circuit voltage ($V_{\text{OC}}$) and the maximum power output ($P_{\text{out}}$) have been obtained as described in chapter 2.2.2 and are presented in Table 12.

From Figure 55 and Figure 56 it can be seen that I succeeded in making a POPV device of $N_n$, which shown good diode behavior with a rectification ratio of $\sim 35$ at bias of $\pm 1$ V. Furthermore only a small ohmic contribution is observed at high negative bias, which means that the shunt resistance is large and only a small current leakage is present. Unfortunately the power conversion efficiency of $N_n$ under illumination of white light is lower than the typical value of 0.001 % for single layer POPV devices. Lower values of $I_{\text{SC}}$ and $V_{\text{OC}}$ than usual for single layer POPV devices are also observed. The lower value of $V_{\text{OC}}$ compared to the difference in workfunctions of the two electrodes is probably related to the fast degradation of the polymer caused by oxygen. This explanation is supported by the fact that the devices were made under a normal atmosphere containing air, were not encapsulated and had a very short life times (see Figure 57). The low $I_{\text{SC}}$ values are probably caused by two things. First, the devices made in this project have an active area of 3 cm$^2$, which is much larger than the typical active area of POPV devices reported in the literature, which is a few mm$^2$. Research at Risø, Denmark, have shown that it seems to be the general case that there is a large serial resistance present in large area POPV devices, which decrease both $I_{\text{SC}}$ and $I_{\text{max}}$ and naturally also both $\eta_{\text{eff}}$ and FF. Second, molecular oxygen might have attacked the very sensitive triple bonds in the polymer backbone and thereby have broken some of the conjugation. This will cause some of the charge carriers to be trapped in isolated segments between the broken conjugations, and thereby lead to an increase in the serial resistance.

In conclusion the low value for $\eta_{\text{eff}}$ in $N_n$ is most likely due to the larger mismatch with the solar spectrum compared to PPVs or PATs and to the fact that the sensitive triple bonds very easily are attacked by molecular oxygen or perhaps by the negative metal electrode. The above reasons are probably also the reason, why no one else has reported characterization of POPV devices made of single layer PPEs.
Figure 55. The linear dark and photo I/V characteristic of bilayer devices of C₆₀ with Nₙ and NPN B respectively and blended devices with PCBM and Nₙ.⁴

Figure 56. The logarithmic dark and photo I/V characteristic of bilayer device of C₆₀ with NPN B and a blended device of PCBM and Nₙ respectively.
Figure 57. The absolute value of $I_{SC}$ as function of the time for the N$_n$/PCBM device.

Table 12. The summarized I/V characteristics of the POPV devices of N$_n$ and NPN B. The solar simulator have generated AM 1.5 light ($\sim$100 mW cm$^{-2}$) and the active area of the devices was 3 cm$^2$.

<table>
<thead>
<tr>
<th>POPV device</th>
<th>$I_{sc}$ [$\mu$A cm$^{-2}$]</th>
<th>$V_{oc}$ [V]</th>
<th>$P_{out}$ [$\mu$W cm$^{-2}$]</th>
<th>FF [%]</th>
<th>$\eta_{eff}$ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>N$_n$</td>
<td>2.78</td>
<td>0.19</td>
<td>$1.4 \times 10^{-1}$</td>
<td>26</td>
<td>$1 \times 10^{-4}$</td>
</tr>
<tr>
<td>N$<em>n$/C$</em>{60}$</td>
<td>11.3</td>
<td>0.20</td>
<td>$7.2 \times 10^{-1}$</td>
<td>32</td>
<td>$7 \times 10^{-4}$</td>
</tr>
<tr>
<td>N$_n$/PCBM</td>
<td>18.3</td>
<td>0.47</td>
<td>2.0</td>
<td>23</td>
<td>$2 \times 10^{-3}$</td>
</tr>
<tr>
<td>NPN B</td>
<td>0.16</td>
<td>0.01</td>
<td>$5.9 \times 10^{-4}$</td>
<td>37</td>
<td>$6 \times 10^{-7}$</td>
</tr>
<tr>
<td>NPN B/C$_{60}$</td>
<td>6.88</td>
<td>0.14</td>
<td>$2.8 \times 10^{-1}$</td>
<td>29</td>
<td>$3 \times 10^{-4}$</td>
</tr>
</tbody>
</table>

The low performance of POPV devices made of N$_n$ can be tolerated as long as a significant improvement in performance is observed in the POPV device made of NPN B. Unfortunately it can be seen from Figure 55, Figure 56 and Table 12 that performance of the NPN B POPV devices are even worse than the N$_n$ POPV devices. Even the diode behavior is poorer than the N$_n$ devices. From both the fluorescence and electroluminescence characteristic it is known, that the N domains work as very effective antennas for the P domain. Taking this knowledge into account the poor performance must be due to the fact that either the dissociation of excitons at the P domain is very poor, which means that the molecular heterojunction is not working as intended, or the P domain has difficulties to deliver electrons from dissociated excitons to the electrode or in the case of the bilayer cell to the C$_{60}$ layer. In both cases the excitons or the free charge carriers will recombine after the end of their lifetime and e.g. emitting light or generating heat. Both explanations indicate why the NPN B POPV devices
are much poorer than the Nn POPV devices, and why NPN B works as an effective LED. Similar observations have been made by Krebs & Spanggaard\textsuperscript{45} for a three domain super molecular system, TPT, consisting of the same zinc porphyrin as the P domain and oligo(3-hexylthiophenes) as the T domains. It should be mentioned that Collin et al.\textsuperscript{46} have shown that ruthenium(II) bisterpyridyl complexes covalently linked to zinc porphyrins effectively quenches the fluorescence of the porphyrin, and that this quenching is due to a rapid intramolecular electron transfer process. However both Krebs and Spanggaards and my observation concerning the TPT and NPN B systems show, that more complex systems are needed if effective molecular heterojunctions using porphyrin compounds should be obtained. One possible problem of e.g. the NPN B system is that if excitons are dissociated in the P domain, no good electron conductor is attached to the P domain, and the electrons can have problems reaching the electrodes. A good hole conductor such as a PPV derivative and a good electron conductor such as a fullerene covalently linked to e.g. the ruthenium complex systems of Collin et al. could be a suggestion for a more complex system. This system should make use of the system of Collin et al., which has been proposed to produce free charge carriers, combined with good conductors that are capable of leading the produced free holes and electrons effectively away from the porphyrin unit. The chemistry of this system is however much more complicated, which in the end could be the main problem of these systems. Only time will show.

References

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In this project the synthesis of a new derivative of a poly(phenylene ethynylene), $N_n$, together with the synthesis of a new three domain super molecular structure consisting of a zinc porphyrin linked conjugated polymer, $NPN B$, is presented. During the synthesis problems concerning the control of the nature of the polymer were observed, and it was concluded that the problems were caused primarily by two factors: Oxidative dimerization of the ethynylene groups and remnants of Pd catalyst. The problem concerning the remnants of the Pd catalyst has been solved in an elegant and non expensive manner by dissolving the remnants of the Pd catalysts with the $N,N$-diethylphenylazothioformamide through formation of electron transfer complexes. The electron transfer complexes show either an unusually distorted tetrahedral coordination for electron transfer complexes or a more conventional square planar coordination, depending on the metal. The $N,N$-diethylphenylazothioformamide is not limited to dissolve Pd, as it is shown that $N,N$-diethylphenylazothioformamide also has the ability to dissolve other transition metals like e.g. Pt, Ni and Cu. The uniqueness of $N,N$-diethylphenylazothioformamide is furthermore that it gives the possibility to quantitatively analyze the content of a given transition metal by UV-Vis spectroscopy, with a detection limit as low as 10 ppb. To extend the tool box for removal of remnants of transition metals from a given sample, the development of $N,N$-bis[tetra(ethylene glycol)monomethylether]phenylazothioformamide is presented. The $N,N$-bis[tetra(ethylene glycol)monomethylether]-phenylazothioformamide is specially designed to remove transition metals from samples containing small molecules. The long ethylene glycol monomethylether chains give the ligand and the formed electron transfer complexes a very high affinity for silica gel in even polar solvents like chloroform.

Both LED and POPV devices of $N_n$ and $NPN B$ have been prepared and characterized. It was observed that LED devices based on $N_n$ emit greenish blue light while LED devices based on $NPN B$ emit light in the near-infrared region. It is demonstrated that the N domains in $NPN B$ work as very effective antennas for the P domain, and effectively transfer electrically generated excitons in the N domain to the P domain. The performance of the POPV devices based on $N_n$ is however much lower than normally observed for single layer polymer devices sandwiched between two electrodes. It is concluded that the low performance is due to a mismatch with the solar spectrum and the attack of molecular oxygen on the very sensitive triple bonds in the backbone of the polymer. The POPV devices based on the $NPN B$ polymer exhibit even poorer performance than devices based on $N_n$. It is concluded that the poor performance of the POPV devices based on $NPN B$ is due to either a missing dissociation of the excitons in the P domain or problems of delivering free electrons from dissociated excitons in the P domain to the electrodes. The overall conclusion is therefore that a more complex system is needed if a molecular heterojunction is to work efficiently.
In this chapter I will summarize my contribution to the paper: "Synthesis, Structure and Properties of 4,7-dimethoxybenzo[c]tellurophene, a Molecular Pyroelectric Material" (paper 7 in appendix B). My contribution consisted of the structural characterization as a function of the temperature by X-ray diffraction and the calculation of the pyroelectric coefficient as function of the temperature. The work was done in collaboration with Michael Pittelkow, Theis K. Reenberg, Magnus J. Magnussen, Theis I. Søling and Jørn B. Christensen from University of Copenhagen.

11.1 Introduction

One of the more remarkable aspects in solid-state chemistry is the connection between the symmetry properties and the electrical properties. The polar axis is responsible for properties such as ferroelectricity and pyroelectricity, and if the point group is non-centrosymmetric properties such as piezoelectricity is a possible. Materials with one of these properties have very interesting applications such as capacitors, infrared radiation detectors and transducers for converting mechanical energy to electrical energy (and vice versa), respectively.

Ferroelectrics have an extremely large permittivity and can retain some residual electrical polarization after an applied potential has been removed. The size of the polarization depends on the size of the applied potential, but has a saturation polarization, $P_s$, at high field strength. A further property of the ferroelectrics is that applying a potential in the opposite direction can reverse the polarization.

Piezoelectrics crystals develop electric charges on the opposite crystal faces, when the crystals are applied a mechanical stress. The polarization, $P$, and the stress, $\sigma$, are proportional and the proportionality coefficient is called the piezoelectric coefficient, $d^2$:

$$ P = d \sigma $$

(11.1)

This property has made piezoelectrics useful, and e.g. physicists at Roskilde University have specialized in making rheometric instruments of Piezoelectric materials to measure both bulk and shear modulus of supercooled liquids.

Pyroelectrics are materials where in a change in the spontaneous polarization, $P$, is observed in connection to a change in temperature, $T$:

$$ \Delta P = p \Delta T $$

(11.2)

where $p$ is the pyroelectric coefficient.

Crystals that exhibit one of the above properties have to be non-centrosymmetric. It should be mentioned that it is still a large unsolved problem to predict and control the crystallization of organic molecules to obtain desired symmetry. Furthermore, it should also be mentioned that only few organic compounds crystallize in a point group, that has a polar axis. Therefore it
is always interesting when a new organic molecule is found to crystallize in a non-centrosymmetric polar point group, and investigations for possible properties like the ones mentioned above should be carried out.

In the present work the unusually stable tellurium compound: 4,7-dimethoxybenzo-[c]tellurophene has been synthesized (see Figure 58). Single crystal diffraction measurements showed that the compound crystallizes in a non-centrosymmetric polar space group, which means that the compound probably exhibit one or more of the three above mentioned properties. Test for ferroelectricity demands large crystals, which not was easy to grow. Concerning the piezoelectric properties, the crystals are very fragile, which means that even if the compound has piezoelectric properties, it cannot be used as transducers because it cannot withstand the mechanical strain. We have therefore decided not to test for piezoelectric properties. From DSC measurements an exothermic phase transition is observed at about 353 K (see supporting information of paper 7 in appendix B). This phase transition opens up for the possibility of the rare primary pyroelectric effect (will be described in following paragraph), if the compound exhibit pyroelectricity.

Before testing the pyroelectric effects, a more clarifying explanation of the pyroelectricity is needed.

11.2 Pyroelectricity
A material is pyroelectric when it shows spontaneous change in polarization when the temperature is changed. The requirement of the polar axis and the non-centrosymmetric symmetry cause crystals that exhibit pyroelectric effects to belong to one of the following 10 out of the 32 possible point groups: Triclinic 1, monoclinic m and 2, orthorhombic mm2, trigonal 3 and 3m, tetragonal 4 and 4mm and hexagonal 6 and 6mm. In point group 1 the polar axis is not determined by symmetry and the vector description of the polar axis therefore has three components for a rectangular coordinate system. In point group m the polar axis lies in the mirror plane and therefore it has only two components. In the other eight point groups the polar axis lies parallel to the unique rotational axis. The unique rotational axis in these point groups corresponds to one of the major crystallographic axes. The vector description of the polar axis therefore has only one component.
The magnitude of the pyroelectric effect is related to the pyroelectric coefficient, which can be divided into three contributions named the primary, secondary and tertiary pyroelectric coefficients. The primary pyroelectric coefficient describes the pyroelectric effect when the shape and volume of the crystal are fixed and the crystal is exposed to a temperature change. It therefore describes the strain-free cases, \( p^S \). The primary pyroelectric effect covers the change in polarization due to abrupt atomic or molecular movements within the crystal as a function of the temperature. The effect is often observed in connection to or near a phase transition where a partial or full cancellation of the dipole moments occur as a consequence of e.g. symmetry changes from polar to non-polar point group.\(^7,^9\) The secondary effect describes the pyroelectric effect when the shape and volume of the crystal are free to change under a uniform change in temperature, which is equivalent to a stress-free case, \( p^T \). This means that the secondary effect describes the change in polarization due to thermal expansion. In practice reported pyroelectric effects on crystals is mostly on the \( p^T \) because this condition is more easily obtained in practice. The tertiary pyroelectric effect is used when crystals experience non-uniform temperature change.\(^8\) By ensuring a uniform temperature change in the crystals, the contribution from tertiary effects to the overall pyroelectric effect is eliminated, and the effect will therefore not be commented anymore.

Since the pyroelectric effect is an equilibrium property it is possible to define the pyroelectric coefficient by use of thermodynamics:\(^*\)

\[
p^T_n = p^S_n + e^e_{n\mu} \alpha^E_{\mu}
\]

(11.3)

where \( p, n, T, S, e, T, \mu, a \) and \( E \) represent the pyroelectric coefficient, the polar axis, constant stress, constant strain, piezoelectric stress coefficient, constant temperature, tensor components, thermal expansion constant and constant electric field. The first term in equation (11.3) represents the primary pyroelectric effect while the second term represents the secondary pyroelectric effect.\(^8,^9\)

### 11.3 4,7-dimethoxybenzo[c]tellurophene

The X-ray structure of 4,7-dimethoxybenzo[c]tellurophene has been solved at 100 K. The compound crystallizes in the polar non-centrosymmetric space group C2, where the polar axis and the C2 axis lie along the crystallographic b axis. There are two molecules in the asymmetric unit cell. The molecules are planar and from the crystal packing (see Figure 59) it can be seen that all the molecules are arranged around the C2 axis in such a way that the plane of the molecules is tilted with an angle of 48\(^\circ\) with respect to the C2 axis.

The crystal packing of the 4,7-dimethoxybenzo[c]tellurophene cause a permanent polarization in the crystal, and in order to investigate the pyroelectric properties and to determine whether the pyroelectric effect is primary or secondary I have investigated the X-ray structure as a function of the temperature. X-ray data have been recorded at the temperatures: 100, 150, 200, 250, 300, 323, 343 and 373 K. The temperature range up to the phase transition at 353 K allowed me to investigate the secondary pyroelectric effect, and the high temperature on the phase transition can give information about a possible primary pyroelectric effect. X-ray data

\(^*\) The definition of the pyroelectric coefficient by thermodynamics is very lengthy and complicated and it is out with of the scope of this thesis to show this definition, there is therefore referred to the literature\(^8\) for a comprehensive description of the definition.
were collected successfully in the temperature range up to 343 K but it turns out that when the temperature was increased to 373 K the crystal blackened and its edges became rounded. Furthermore no diffraction was observed when I attempted to record X-ray data. The temperature was then decreased to 343 K but still no diffraction was observed. This indicates that the crystal is possibly not an equilibrium structure, and further polymorphism is not likely to be present since all diffraction disappears after the heating to the high temperature. In the case of the data collection at 343 K I observed that the intensity of the diffraction began to decrease as a function of time, which results in virtually no diffraction observed after 2 hours. Furthermore the crystals very slowly blackened at this temperature also. It therefore seems plausible that the 4,7-dimethoxybenzoc]tellurophene is not stable at high temperature and looses tellurium, which is collected on the surface of the crystal.

Figure 59. The crystal packing of 4,7-dimethoxybenzoc]tellurophene at 100 K. Note how the arrangement of the molecules causes a net dipole moment along the crystallographic b axis.7

The solution of the X-ray structure of 4,7-dimethoxybenzoc]tellurophene was unproblematic at all temperatures except at 323 K, where it was necessary to employ constraints on the molecular geometry as a consequence of the instability of 4,7-dimethoxybenzo[c]tellurophene at high temperature. The constraints consisted of forcing the benzene ring to approach a hexagonal geometry.

In the case of a possible primary pyroelectric effect, it can be concluded that the phase transition at 353 K observed in DSC does not represent a primary pyroelectric effect since; the crystal seems to break down at this transition. Concerning a possible primary pyroelectric effect in the temperature range from 100 K to 323 K, a structural change has to be observed as function of the temperature. The structural changes have to be of the kind that affects the polarization, which only can happen if the molecules move in such a way that the angle between the dipole of the molecules and the C2 axis changes. In Figure 60 the angles between the molecular plane of both molecules in the asymmetric unit cell determined by the least square fit and the C2 axis have been plotted as function of the temperature. It can be seen that the changes in angle are not significant. I therefore conclude that the contribution from a possible primary pyroelectric effect is negligible.
Figure 60. The angle between the plane core of 4,7-dimethoxybenzo[c]tellurophene and the C2 axis as function of the temperature. The large standard deviation on the data at 323 K is due to the instability at high temperature.

Since secondary pyroelectric effects are solely due to the thermal expansion of the material as a function of temperature, the secondary pyroelectric effect can be estimated by observing the volume of the crystal as a function of the temperature. In Figure 61 the crystallographic b axis...
and the volume is plotted as a function of the temperature. It can be seen that there is a change in size of both components as a function of the temperature, which a secondary pyroelectric effect.

Since the space group is C2, the molecular dipole, $\mu$, is aligned along the crystallographic b axis, and the polarization, $P$, is given in equation (11.4).

$$P(T) = \frac{Z}{V(T)} \begin{pmatrix} 0 \\ \mu \\ 0 \end{pmatrix}$$ (11.4)

where $Z$, $T$, and $V$ are the molecular entities per unit cell, the temperature and the volume as function of the temperature. Since the pyroelectric coefficient at constant stress, $P^T$, is given as the change in polarization with respect to the change in temperature, the following expression for the secondary pyroelectric coefficient is obtained:

$$\frac{dp(T)}{dT} = P^T = \frac{Z}{aT^2 + 2bT + \frac{b^2}{a}} \mu \cos(\theta)$$ (11.5)

where $aT^2 + 2bT + \frac{b^2}{a}$ and $\theta$ are the derivatives of the linear thermal expansion with respect to the temperature and the angle between the plane molecular core and the C2 axis.

![Graph](image)

**Figure 62.** The calculated secondary pyroelectric coefficient as a function of the temperature. In the plot both $p_n^T(T)$ observed and the $p_n^T(T)$ mean are shown. The $p_n^T(T)$ calculated is the secondary pyroelectric effect calculated on basis of equation (11.5) by use of the calculated dipole moments (see paper 7), the thermal expansion from Figure 61 and the angle between the plane molecular core and the C2 axis from Figure 60. The $p_n^T(T)$ mean is calculated almost as the $p_n^T(T)$ observed but by use of the mean value of the calculated dipole moments and the mean value of the angle between the plane molecular core and the C2 axis.  

On basis of equation (11.5) and the molecular dipole moment, which can be calculated on basis of the molecular coordinates obtained from the X-ray crystal structures (see paper 7 for
further details), it is possible to calculate the secondary pyroelectric coefficient as a function of the temperature. In Figure 62 the calculated secondary pyroelectric coefficient is plotted as a function of the temperature.

By comparing with other pyroelectric materials it can be seen that the pyroelectric coefficient is comparable in magnitude with known values for other organic compounds such as phosphangulene, which have a coefficient of $3.2 \times 10^{-6} \text{ C m}^{-2} \text{ K}^{-1}$. The magnitude of the pyroelectric coefficient is however nearly a factor 1000 smaller than some of the best inorganic pyroelectric materials such as La:SBN.9

References
Appendix A

12.1 Pd(PPh₃)₄ catalyst

The tetrakispladium(0) was prepared as described in the literature. The reaction runs as mentioned below:

\[
2 \text{PdCl}_2 + 8\text{P(C}_6\text{H}_5\text{)}_3 + 5 \text{NH}_2\text{NH}_2 \text{H}_2\text{O} \rightarrow 2 \text{Pd}[\text{P(C}_6\text{H}_5\text{)}_3]_4 + 4 \text{NH}_2\text{NH}_2 \text{HCl} + \text{N}_2 + 5 \text{H}_2\text{O}
\]

In a 500 mL round-bottom three-necked flask palladium dichloride (0.014 mol, 2.53 g) and triphenylphosphine (0.070 mol, 18.69 g) were suspended in dimethyl sulfoxide (170 mL). The flask was equipped with an egg shaped magnetic stirring bar, rubber septum, thermometer and a condenser connected to an argon system with provision made for pressure relief though a silicon oil bubbler. The greenish yellow suspension was heated under stirring until complete solution occurred, which was about at 413 K. The heating was then stopped; and the solution was rapidly stirred until the temperature of the solution fell to 403 K. Hydrazine monohydrate (0.056 mol, 2.73 mL) was then added through the septum by a plastic syringe over 1 minute. It is important to cool the solution down to avoid that the hydrazine should decompose in an uncontrolled manner, but the temperature of the solution must not be lower than 403 K, because PdCl₂(PPh₃)₂ will then precipitate. After the addition of the hydrazine, the solution was cooled down to room temperature and the yellow solution was filtered through a glass filter funnel under an argon atmosphere and washed with 100 mL ethanol and 200 mL diethyl ether. Passing a steam of argon over the funnel for 1 hour dried the product. The yield was 16.00 g (99 %).

12.2 PdCl₂(PPh₃)₂ catalyst

The PdCl₂(PPh₃)₂ catalyst was prepared as described in the literature. The reaction is as follow:

\[
PdCl_2 + 2\text{P(C}_6\text{H}_5\text{)}_3 \xrightarrow{\text{DMSO}} PdCl_2(\text{PPh}_3)_2
\]

In a 500 mL round-bottom three-necked flask equipped with an egg shaped magnetic stirring bar, thermometer and condenser DMSO (400 mL) was added and flushed with argon for 15 min. Palladium dichloride (0.029 mol, 5.16 g) and triphenylphosphine (0.073 mol, 19.25 g) were added under an argon atmosphere. The suspension was heated to 413 K while stirring. The solution changed colour from cloudy yellow to clear red. The heating and stirring were stopped and the solution was allowed to cool down slowly, while yellow crystals precipitated. The solution was filtered through a glass filter funnel and the crystals were washed with 1 L diethyl ether. The crystals were dried in a vacuum oven. The yield was 19.82 g (97 %).
12.3 2,5-dioctylbenzene (1)

2,5-dioctylbenzene was prepared by a normal Grignard reaction\(^2\).

The reaction runs as showed below:

\[
\begin{align*}
\text{Mg (s)} & \quad + \quad \text{C}_8\text{H}_{17}\text{Br} \quad \rightarrow \quad \text{C}_8\text{H}_{17}\text{MgBr} \\
2 \text{C}_8\text{H}_{17}\text{MgBr} & \quad + \quad \text{Cl} \quad \rightarrow \quad \text{C}_8\text{H}_{17}\text{Cl}_2 \\
\text{C}_8\text{H}_{17}\text{Cl}_2 & \quad + \quad \text{C}_8\text{H}_{17}\text{MgBr} \quad \rightarrow \quad \text{C}_8\text{H}_{17}\text{MgBr} \\
\text{C}_8\text{H}_{17}\text{MgBr} & \quad + \quad \text{NiCl}_2\text{dppp} \quad \rightarrow \quad \text{C}_8\text{H}_{17}\text{C}_8\text{H}_{17} \\
\text{C}_8\text{H}_{17}\text{C}_8\text{H}_{17} & \quad + \quad \text{NiCl}_2\text{dppp} \quad \rightarrow \quad \text{C}_8\text{H}_{17}\text{C}_8\text{H}_{17}
\end{align*}
\]

Exact Mass: 145.97
Mol. Wt.: 147.00
C, 49.02; H, 2.74; Cl, 48.23

Exact Mass: 302.30
Mol. Wt.: 302.54
C, 87.34; H, 12.66

In a 2000 mL three-necked round-bottom flask filled with argon, Mg (1.06 mol, 25.71 g, 6 % excess) was covered with Et\(_2\)O (150 mL). Et\(_2\)O (350 mL) was added drop wise in a mixture with the n-octylbromide (0.99 mol, 191.61 g) under argon atmosphere and over a time period of 2.5 hour. Even though the reaction is exothermic, it was necessary to add some crystals of iodine and heat the solution with a heat gun before the reaction started. After the addition of the halide the solution was stirred in 1 hour under argon atmosphere. Hereafter the catalyst (1.01 g of NiCl\(_2\)dppp) and a solution of 1,4-dichlorobenzene (0.50 mol, 73.57 g) in Et\(_2\)O (500 mL) were added to the solution, and the color of the mixture shifted from gray to light green. The solution of dichlorobenzene in Et\(_2\)O was flushed with argon in 15 minutes before the addition. After the addition the mixture was stirred and refluxed overnight. Next day a lot of solid matter (MgBrCl) had precipitated, so the solution seemed nearly solid, and the color of the solution was shifted from green to brown. The solution was poured into 1.5 L ice water, and the phases were separated by 2 L Et\(_2\)O. Note that Mg reacts very violently with water. The organic phase was washed three times with water, and dried with MgSO\(_4\). After drying, the organic phase was filtered and evaporated to dryness to provide 127.55 g of oil. Yield 84 %. \(^1\)H and \(^13\)C NMR identified the product as 1,4-dioctylbenzene, and showed that it was clean.

\(^1\)H NMR (250 MHz, CDCl\(_3\), 300 K, TMS): \(\delta = 0.87\) (6 H, double t), 1.26 (20 H, m), 1.58 (4 H, double q), 2.54 (4 H, double t), 7.05 (4 H, s).

\(^13\)C NMR (63 MHz, CDCl\(_3\), 300 K, TMS): \(\delta = 14.13, 22.78, 29.41, 29.53, 29.64, 31.69, 32.03, 35.71, 128.26, 140.03\).
12.4 1-(4-bromophenyl)-2-trimethylsilylacetylene (6)

1-(4-bromophenyl)-2-trimethylsilylacetylene was prepared as described in the literature with a few exceptions.

The reaction runs as mentioned below:

\[ \text{BrI} + \text{TMS} \xrightarrow{\text{Pd(PPh}_3\text{)}_2\text{Cl}_2, \text{CuI, N(CH}_2\text{CH}_3)_3} \text{HN(CH}_2\text{CH}_3)_3\text{I} \]

\[ \text{C}_6\text{H}_4\text{BrI} \quad \text{Mol. Wt.: 282.90} \]
\[ \text{C}_5\text{H}_{10}\text{Si} \quad \text{Mol. Wt.: 98.22} \]
\[ \text{C}_{11}\text{H}_{13}\text{BrSi} \quad \text{Exact Mass: 252.00} \]
\[ \text{Mol. Wt.: 253.21} \]

C, 52.18; H, 5.17; Br, 31.56; Si, 11.09

A 500 mL round-bottom flask was dried with a heat gun and filled with argon. Then a mixture of 1-bromo-4-iodobenzene (82.7 mmol, 23.39 g) and trimethylsilylacetylene (82.7 mmol, 8.12 g) was added together with triethylamine (200 mL). The solution was placed in an ice bath and flushed with argon for 30 minutes, before the catalyst Pd(PPh\(_3\))\(_2\)Cl\(_2\) (1.16 g, \(\approx 2\) mol \%) and the co-catalyst CuI (1.55 g, \(\approx 10\) mol \%) were added under an argon atmosphere. The mixture was stirred overnight under an argon atmosphere. The colour of the mixture was dark before the addition of the catalyst. After the addition of the catalysts the colour became brownish and cloudy.

Next day GC/MS showed that the reaction had stopped and there was still about 9 % of 1-bromo-4-iodobenzene left in the mixture. 9 mole % of trimethylsilylacetylene (7.4 mmol, 0.73 g) was therefore added together with a little more catalyst (Pd(PPh\(_3\))\(_2\)Cl\(_2\) 0.11 g, \(\approx 2\) mol \%) and CuI (0.14 g, \(\approx 10\) mol \%), and the mixture was stirred under an argon atmosphere for another 4 hours. At this point GC/MS showed that all of the 1-bromo-4-iodobenzene was converted to 1-(4-bromophenyl)-2-trimethylsilylacetylene. Diethyl ether (150 mL) was added, and the suspension was filtered through a glass filter-funnel (size 3) (triethylammoniumhydrogen iodide was removed). The filtrate was collected and evaporated to dryness by rotary evaporation. The liquid residue (yellow oil) was dissolved in n-heptane (300 mL) and then purified on an aluminium oxide column. The n-heptane was removed by evaporation, and the product was further purified by recrystallization in 50 mL methanol and washed with cold methanol. 1-(4-bromophenyl)-2-trimethylsilylacetylene was obtained as a white powder in 67 % yield (14.09 g).

\(^1\)H NMR (250 MHz, CDCl\(_3\), 300 K, TMS): \(\delta = 0.24\) (9 H, s), 7.30 (2H, double t (aa’bb’para-system), \(^3J = 8.6\) Hz, \(^4J = 2.0\) Hz), 7.49 (2H, double t (aa’bb’para-system), \(^3J = 8.6\) Hz, \(^4J = 2.0\) Hz).

\(^13\)C NMR (63 MHz, CDCl\(_3\), 300 K, TMS): \(\delta = 0.00\), 95.6, 103.9, 122.1, 122.7, 131.5, 133.4.

Anal. Calcd. for C\(_{11}\)H\(_{13}\)BrSi: C: 52.18, H: 5.17, N: 0.00. Found: C: 52.03, H: 4.98, N:0.00.
12.5 1-(4-iodophenyl)-2-trimethylsilylacetylene (7)

The reaction:

\[
\text{Br} + \text{THF} + \text{n-BuLi} \rightarrow \text{Li} + \text{I}_2
\]

C\text{\(_{11}\)H\text{\(_{13}\)BrSi}}

Exact Mass: 252.0
Mol. Wt.: 253.21
C, 52.18; H, 5.17; Br, 31.56; Si, 11.09

In a 500 mL conical flask 1-(4-bromophenyl)-2-trimethylsilylacetylene (1.35 \times 10^{-1} \text{ mol}, 34.12 g) was dissolved in dry THF (200 mL) under an argon atmosphere. Meanwhile 350 mL THF was cooled to 195 K in a 2000 mL three-necked round-bottom flask. After the THF had reached the temperature of 195 K, n-BuLi (130 mL of a 1.1 M solution in hexane (0.143 mol, \approx 10 \% excess) was dissolved in the THF under an argon atmosphere. The dissolution process is very exothermic and the temperature of the solution increased to 233 K. The solution was cooled to 195 K, whereupon the solution of 1-(4-bromophenyl)-2-trimethylsilylacetylene was added drop wise over a period of 2 hours and under an argon atmosphere, so the temperature did not exceed 223 K. After the temperature again was cooled down to 195 K (after about 30 minutes), I\text{\(_2\)} (0.15 mol (\approx 10 \% excess), 37.69 g) was added and the temperature of the mixture was allowed to rise to room temperature. The mixture was then treated with petroleum ether (200 mL), and washed with a saturated solution of Na\text{\(_2\)}S\text{\(_2\)}O\text{\(_3\)} \cdot 5H\text{\(_2\)O} in water (4-250 mL). The aqueous phase was then washed with diethyl ether (100 mL). The organic phases was collected and dried with MgSO\text{\(_4\)}, whereupon the solvent was removed by evaporation. The yield was 39.42 g of yellow oil. The product was recrystallized in methanol (200 mL) and washed with cold methanol to provide 29.98 g of a white powder. Yield: 74 \%.

\(^1\text{H NMR (250 MHz, CDCl\text{\(_3\)}, 300 K, TMS):} \delta = 0.24 (9 \text{ H, s}), 7.20 (2\text{H, double t (aa’bb’para-system)}, \(3J = 8.4 \text{ Hz}, 4J = 1.9 \text{ Hz})\), 7.6 (2\text{H, double t (aa’bb’para-system)}, \(3J = 8.4 \text{ Hz, } 4J = 1.8 \text{ Hz})\).

\(^{13}\text{C NMR (63 MHz, CDCl\text{\(_3\)}, 300 K, TMS):} \delta = -0.1, 94.4, 95.9, 104.0, 122.6, 133.4, 137.4\).
12.6 4-(2-trimethylsilylethyl)benzoic acid

\[
\begin{align*}
\text{C}_7\text{H}_4\text{IO}_2 &+ \text{TMS} \xrightarrow{\text{PdCl}_2(\text{PPh}_3)_2, \text{CuI}, \text{Et}_3\text{N}, \text{THF}} \text{C}_{12}\text{H}_{14}\text{O}_2\text{Si} \\
\text{Exact Mass: 247.93} & \quad \text{Exact Mass: 218.08} \\
\text{Mol. Wt.: 248.02} & \quad \text{Mol. Wt.: 218.32} \\
\end{align*}
\]

In a 500 mL round-bottom flask the iodobenzoic acid (0.097 mol, 24.07 g) was dissolved in 150 mL triethylamine and 250 mL THF. The solution was flushed with argon in 30 minutes before the catalyst PdCl₂(PPh₃)₂ (1.00 g ≈ 1.5 mol%) and the co-catalyst CuI (1.50 g ≈ 10 mol%) were added under an argon atmosphere. Immediately after, TMSA (0.107 mol, 15.09 mL ≈ 10% excess with respect to iodobenzoic acid) was added. The solution was stirred overnight at RT. Before the addition of the catalysts, the color of the solution was clear, but it shifted quickly to green when adding the catalysts. After the addition of TMSA the color shifted to black.

Next day a lot of grayish black solid matter had precipitated. Diethyl ether (150 mL) was added and the suspension was filtered through a glass filter-funnel. The filtrate was collected and evaporated to dryness by rotary evaporation. The liquid residue (yellow oil) was dissolved in dichloromethane (100 mL) and tried purified on an aluminum oxide (neutral) column. Unfortunately, the product interacts very strongly with the aluminum oxide, and it was impossible to liberate the product from the column with normal organic solvents. The product was successively removed from the column by glacial acetic acid (500 mL). The acetic acid was removed by rotary evaporation, and the yield was 17.52 g (83%). It was attempted to further purify the product by recrystallization in ethanol, but it failed because the product was too soluble in ethanol. The recrystallization also failed in both heptane and acetyl acetate because the product was too soluble in these solvents. Another attempt to purify the product was made by dissolving the product in water and THF followed by addition of ½ eqv. Cs₂CO₃ (12.44 g) to precipitate the Cs salt. The solution was then filtered. The solvents were then removed by evaporation (they foam very much). NMR shows that the compound was not clean.

The protecting group (TMS) was removed by K₂CO₃. The product was dissolved in 150 mL methanol, and after flushing with argon for 30 minutes 10 eqv. K₂CO₃ (14 g) was added, and the solution was stirred for 2 hours under argon atmosphere. After 2 hours NMR showed that the TMS group had been removed. The solvent was removed by evaporation and the product was dissolved in 500 mL 10 % HCl and 500 mL chloroform to form the acid. A lot of copper colored solid matter was precipitated, when the HCl was added. The solid matter was collected and extracted by soxhlet in 2 L chloroform over 1 week to give clean 4-ethynylbenzoic acid.

\(^1\text{H NMR (250 MHz, DMSO, 300 K, TMS): } \delta = 4.43 (1\text{H, s}), \quad 7.61 (2\text{H, dm (aa’bb’para-system)}), \quad 3^J = 8.49 \text{ Hz}, \quad 4^J = 3.49 \text{ Hz}), \quad 7.96 (2\text{H, dm (aa’bb’para-system)}), \quad 3^J = 8.57 \text{ Hz}, \quad 4^J = 3.55 \text{ Hz}), \quad 13.15 (1\text{H, s}).}
$^{13}$C NMR (63 MHz, DMSO, 300 K, TMS): $\delta$ = 82.68, 83.40, 125.96, 129.40, 130.86, 131.82, 166.56.

Anal. Calcd. for C$_{11}$H$_{13}$BrSi: C: 73.97, H: 4.14, N: 0.00. Found: C: 73.61, H: 3.98, N: 0.00.

12.7 The Cs salt

The Cs salt of the 4-ethynylbenzoic acid was produced by dissolving 4-ethynylbenzoic acid (10.2 mmol, 1.487 g) and $\frac{1}{2}$ eqv. Cs$_2$CO$_3$ (1.660 g) in 200 mL ethanol (96 %) and letting it reflux for 2 hours. The solvent was then removed by evaporation, and the salt was then dried in a vacuum oven at 333 K over night. Yield is 2.824 g (98 %).

12.8 Coupling of the Cs salt to the resin

The coupling is made with 10 mmol Cs salt (2.824 g), 10.012 g resin and 1 mmol CsI (0.263 g). All three compounds were mixed in a special flask made for solid phase synthesis, and about 150 mL dried DMF (freshly distilled) was added. The suspension was flushed with argon for 30 minutes and the flask was closed and shaken for one week.

After a week the suspension was washed with 200 mL DMF, then it was shaken by 150 mL DMF for one hour, and then washed with the following series of liquids: 150 mL water, 150 mL ethanol, 150 mL acetone, 150 mL THF and 150 mL diethyl ether.

The resin was then dried in a vacuum oven and weighed. The yield was 10.437 g, which corresponds to $\Delta W = 0.425$ g.

All the solvents from the washes were collected and evaporated. The residue was then treated with 65 % nitric acid and then treated with a little excess of silver nitrate. The precipitated silver chloride was collected, dried in the vacuum oven and then weighed. The yield was 1.299 g.

12.9 Coupling of the iodomonomer to the anchor unit

Here the general coupling of an iodomonomer to an anchor unit or to other monomers is described. The description is made on the basis of the coupling of the iodomonomer to the anchor unit.

The resin (5.034 g) and 9 (5 mmol, 3.173 g) were mixed in the flask and 150 mL NEt$_3$ was added. The suspension was flushed with argon for 30 minutes before CuI (10 mol%, 0.094 g) and Pd(PPh$_3$)$_4$ (2 mol%, 0.116 g) were added and the flask was closed and shaken for one week at room temperature.

After a week the suspension was washed with 200 mL DMF, then it was shaked by 150 mL DMF for one hour, and then washed with the following series of liquids: 150 mL water, 150 mL ethanol, 150 mL acetone, 150 mL THF and 150 mL diethyl ether.

The resin was then dried in a vacuum oven and weighed. The yield was 5.586 g, which corresponds to $\Delta W = 0.552$ g.
12.10 The deprotection of the anchored unit
Here the general deprotection of the anchored unit is described. The description is made on the basis of the deprotection of the anchored monomer unit.

The resin (5.661 g) was mixed in the flask with 150 mL THF and flushed with argon for 30 minutes before TBAF (tetrabutylammonium fluoride) (~10 eqv. (large excess), ~3 g) was added and the flask was closed and shaken for two days at room temperature.

The suspension was washed with 200 mL DMF, and then it was shaken by 150 mL DMF for one hour. After this it was washed with the following series of liquids: 150 mL water, 150 mL ethanol, 150 mL acetone, 150 mL THF and 150 mL diethyl ether.

The resin was dried in a vacuum oven and weighted. The yield was 5.453 g, which corresponds to \(\Delta W = 0.208 \) g.

12.11 Cleaving of the anchored unit
NaOH (~20 eqv. ~1 g) was dissolved in 50 mL methanol before 50 mL THF were added. The mixture was flushed with argon in 30 min. before the resin was added. The suspension was stirred and refluxed under an argon atmosphere over night. Next day the temperature of the solution was lowered to RT and 3M hydrochloric acid was added drop wise to acidy the suspension. The suspension was filtered and the organic phase was separated and washed with water three times (50 mL) and dried with magnesium sulphate. The solution was filtered and tested by SEC. SEC unfortunately showed that the product mainly consisted of polymers. Further workup was therefore abandoned.

References
Appendix B contains copies of all the publications pertaining to this work.

13.1 The list of publications

Paper one:

Paper two:

Paper three:

Paper four:

Paper five:
Appendix B

**Paper six:**


**Paper seven:**


**Poster:**


**Patent:**

Synthesis, Light Harvesting, and Energy Transfer Properties of a Zinc Porphyrin Linked Poly(phenyleneethynylene)

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ABSTRACT: The synthesis of a zinc porphyrin linked conjugated homopolymer is presented. The synthetic strategy employed a directional polymerization of 1-iodo-2,5-diocylo[4,4'-terphenyl]-10,20-bis(5,5-bis-tert-butylphenyl)-5,15-diethynylporphinatozinc(II) (12) or 10,20-bis(5,5-bis-tert-butylphenyl)-5,15-dibromoazoporphinatozinc(II) (16). In the case of 12 the reaction was found to give a polymer product with a constant ratio of zinc porphyrin to incorporated monomer units regardless of the molecular weight. In the case of 16, however, the ratio of zinc porphyrin to incorporated monomer units decreased with increasing molecular weight of the polymer product. The polymer products were separated into fractions using preparative SEC. Studies of the energy transfer properties from the polymer to the zinc porphyrin revealed that there was little energy transfer in solution but quantitative energy transfer in the solid state. The polymer products proved difficult to handle in terms of device construction where a thin polymer film is sandwiched between two electrodes. We found this to be a consequence of the palladium catalysis employed during the polymerization. We finally discuss the issues concerning the problems with remnants from the Pd catalyst in the polymer samples and their removal.

Introduction

Since the discovery in 1974 of doped polyacetylene showing electrical conductivity,† there has been a large interest in conjugated polymers and their application for e.g., light-emitting diodes,‡ polymer electronic circuits,§ and polymer solar cells.¶

Recently, the synthesis of a three-domain (JPJ) and a five-domain (FPFJ) structure were reported.** The three-domain structure consisted of two homopolymer blocks (J domains) with a porphyrin dye molecule (P domain) in the middle. In this three-domain structure, the J domains were poly(Z,5'-dicarbonyl-4,4'-terphenyl-5,5'-poxy-1,1'-biphenyl], and the distribution of the individual incorporated monomer units was a broad distribution. The P domain was a 5,10,15,20-tetrapheny1porphyrin derivative, and there was only one porphyrin molecule in each JPJ unit. In contrast to JPJ the PPFJ consisted of four oligoquinone (F domains) linked to a porphyrin dye molecule (P domain). A very interesting property observed for these three-domain and five-domain structures was that both the J and the F domain served as a light-harvesting antenna for the P domain. We decided to investigate and evaluate the use of poly(phenyleneethynylene) as the backbone of the light-harvesting polymer chains for the porphyrin dyes and have two alkyl groups on every second phenyl ring to obtain solubility and reduce the distortion of the backbone.¶

In this paper we present a synthesis of a conjugated homopolymer of the poly(phenyleneethynylene) type. We discuss two different synthetic strategies toward linking the conjugated homopolymer blocks via a zinc porphyrin to give the three-domain structure NPN, as shown in Figure 1.

We present studies of the optical properties of the compounds with a focus on the band gap, the quantum efficiency of the fluorescence, and the lifetimes. These studies were performed on the homopolymer, poly(Z,5'-diocylo[4,4'-terphenyl]-10,20-bis(5,5-bis-tert-butylphenyl)-5,15-diethynylporphinatozinc(II) (NPN), on the NPN structure with a broad distribution and on fractions with a narrow distribution prepared by preparative size exclusion chromatography (SEC). Finally, we discuss the problems concerning the use of Pd as a catalyst for the polymerization due to the catalyst remnants contained in the polymer product and provide a solution to this problem in the context of this work.

Experimental Section

1H and 13C NMR spectra were recorded on a 200 MHz Bruker NMR spectrometer at 300 K unless otherwise stated. All spectra were recorded in CDCl3, with TMS as an internal reference. Compound 8 and 1,4-diocylo[4,4'-terphenyl] were prepared according to the procedures described in the literature. Compounds 9–12 were prepared according to the procedures reported by Plater et al. with the exception that compound 12 was purified by preparative SEC before deprotection of the TMS groups. Detailed descriptions of synthetic procedures and analytical data for compounds 1, 2, 3, 4, 6, 7, and 8 can be found in the Supporting Information. Electroactive device preparation is described in detail in the Supporting Information.

Liquid Chromatography Methods. The apparatus for the SEC consists of a preparative system with a 3 mL injection loop. The detector is a UV–vis diode array with a spectral resolution of 1 nm. A 16-port switch valve allows for separation into fractions. Two different column systems were used. They consisted of DVB linked polystyrene gel columns comprising a guard column and two gel columns. One system had a 100 Å and a 1000 Å column in succession, and the other had a 500 Å and a 1000 Å column in succession. The eluent was chloroform, and the flow rate was 20 mL/min with a pressure of 95 bar. The molecular weight was calculated on the basis of polystyrene standards having molecular weights of 515 000, 127 000, 68 000, 36 500, 19 900, 19 900, 32 500, and 13000. In all cases the polydispersity of the standards was smaller than 1.007.

Photophysical Methods. The UV–vis solution spectra were recorded in dry CHCl3 degassed with argon. Emission spectra and lifetimes were recorded in degassed CHCl3. The absorption maximum was kept below 0.07. All emission spectra are corrected. Quantum yields where measured rela-
Appendix B

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**Zn Porphyrin Linked Poly(phenyleneethynylene) 1181**

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**Figure 1.** Three-domain structure of the NPN dye linked homopolymer. Adhesive to DPA. The UV-vis solid-state spectra were recorded on spin-coated films of the polymers. Poly(2,5-diocytotiolylthiophenylene) (N₂). A 250 mL round-bottom flask was dried using a heat gun and flushed with argon. Compound 8 (1.36 mmol, 0.750 g) was placed in the flask and dissolved in 40 mL of piperidine. The solution was flushed with argon for 30 min before the catalyst Pd(PPh₃)₄ (30 mg, 0.02 mol%) and the cocatalyst CuCl (25 mg, 0.10 mol%) were added under an argon atmosphere. After the addition of the catalyst the mixture was refluxed overnight and was poured into 500 mL of methanol. The following day a green precipitate formed that was filtered on a glass filter-funnel. The filter cake was washed with both cold methanol and cold diethyl ether and was stirred in the air to give 1.021 g of polymer. **H** NMR δ: 0.73–1.03 (m, 190 H), 1.03–1.91 (m, 740 H), 2.15 (m, 8 H), 2.32 (m, 9 H), 2.56 (m, 13 H), 2.82 (broad s, 7 H), 6.31–6.61 (m, 14 H), 6.79–7.16 (m, 286 H), 9.91 (m, 4 H), 9.79 (m, 4 H).

**Diethylammonium Diethylthiocarbamate (DDC).** DDC was prepared by the method described in the literature for sodium methylthiocarbamate. Carbon disulfide (10.15 mol, 11.42 g) was added dropwise to a solution of diethylamine (0.4 mol, 29.25 g) in diethyl ether (350 mL), and after a while small colorless crystals were formed. The material was collected on a filter and washed with copious amounts of diethyl ether and dried in a vacuum oven. The yield was 12.20 (68%); mp 81–82 °C. **H** NMR δ: 1.24 (t, 6 H, J = 7.3 Hz), 1.37 (t, 6 H, J = 7 Hz), 2.15 (q, 4 H, J = 7.3 Hz), 4.10 (q, 4 H, J = 7 Hz), 8.27 (broad s, 2 H). **13C** NMR δ: 12.09, 13.05, 41.70, 48.54. Anal. Caled for C₆H₁₂S₂N: C, 45.60; H, 9.97; N, 12.60; S, 28.83. Found: C, 48.73; H, 10.17; N, 12.54; S, 28.37.

**General Procedure for the Removal of Pd from Polymer Samples Using DDC.** The polymer was dissolved in the minimum amount of chloroform. DDC (5 equiv relative to the Pd employed in the catalyst) was added, and the solution was stirred for 5 min. The solution was poured into methanol (10 time the volume of chloroform) and the polymer precipitated. The precipitate was filtered and washed with methanol and diethyl ether.

**Results and Discussion**

**Monomer Synthesis.** The synthesis of the monomer unit is shown in Scheme 1. The synthesis involves the readily available compounds 1,4-dibromo-2,5-diocytolbenzene (1) and 1-(4-isodophenyl)-2-trimethylsilylethynylene (TMSA). The first step (Scheme 1A) is a bromine–iodine exchange of compound 1 using n-BuLi in THF at -78 °C. The bromoiodo compound (2) was cross-coupled with TMSA by Pd/Cu catalysis in the following step to form the 1-(4-bromo-2,5-diocytolphenyl)-2-trimethylsilylethynylene (3). In this step it was found important that the coupling reaction took place at low temperature to increase the selectivity between iodo vs bromo substitution. Even at 273 K we obtained ~2% of
Appendix B

Scheme 1. Synthetic Procedure Leading to the Monomer Unit 1-Iodo-2,5-dioctyl-4'-trimethylsilylthynyltolane (7)*

- **A)**
- **B)**
- **C)**

* (a) (1) n-BuLi, THF, (2) Lg, (3) TMSA, Pd(PPh3)4Cl2, CuI, EqsN, (c) K2CO3, MeOH, CH3Cl. Compounds B1-B4 were side products isolated from the reaction mixtures in step b.

1,4-di(trimethylsilylethylene)-2,5-dioctylbenzene (B1) and 2-4% of 1,4-di(trimethylsilylbutadiyne) (B2) as byproducts which were detected by 1H and 13C NMR, GC-MS, and MALDI TOF. The TMS group was removed by K2CO3, and the 4-bromo-2,5-dioctylphenylethylenylene (4) was obtained. The monomer unit 1-iodo-2,5-dioctyl-4'-trimethylsilylthynyltolane (7) was then obtained by an additional two-step reaction. The first step being a Sonogashira cross coupling reaction between compound 4 and 1-(4-iodophenyl)-2-trimethylsilylacetylene (5) (Scheme 1B) to obtain 1-bromo-2,5-dioctyl-4'-trimethylsilylthynyltolane (6). The final step was a standard bromine-iodine exchange.

The Sonogashira reaction for this type of molecules has been described by others13 and generally works quite well. Most authors, however, never comment on the fact that the Sonogashira alkylnyl-aryl cross-coupling cannot be performed without significant side reactions. Known byproducts are 1,4-diarylbisynes, 1,4 diarylbuto 2-en-1-ynes, and the trimerization product 1,3,5-triarylbenzene.18-20 Of those byproducts we have only detected butadiyne, and this was in connection with the synthesis of compound 3. With 1H and 13C NMR and MALDI TOF we instead detected 3% 1-bromo-2,5-dioctyl-4'-4'-trimethylsilylthynylphenyl-ethynyltolane (B3) as a byproduct during the synthesis of compound 6. Furthermore, we found ~1% of compound 4. We believe this to be a result of incomplete reaction. We also detected ~1% of 1,4-dioctyl-2,5-bis-((4'-trimethylsilylthynylphenylethynyl)benzene (B4). The presence of B4 is a result of the difficulty in separating B1 from compound 3. The Rf values in n-heptane of compounds 4, 6, B3, and B4 are 0.58, 0.38, 0.25, and 0.08, so it was possible by flash chromatography to separate 6 from the byproducts. The removal of byproducts in this step was found to be essential in order to control the later polymerization process. As a consequence, compound 6 was obtained in low yield. It should be noted that, besides the above-mentioned byproducts, we have detected some minor byproducts, but the amount of these was so small that we could not determine the structures.

Porphyrin Synthesis. The porphyrin subunit should have a structure that allowed for attachment of the polymer blocks in the 5,15-positions. We chose the synthesis scheme presented by Plater et al.11 due to the possibility of large scale synthesis. The reported procedure allowed for the synthesis of the porphyrin (9) (see Scheme 4) on a 100 g scale of sufficient purity for the synthesis of the zinc dibromoporphyrin (10).

It was possible to obtain 10 in high purity, which we ascribe to the fortuitous low solubility of 10 making the aleviation of byproducts relatively easy. Plater et al. also reported the synthesis of biethynylporphyrin (12) through Sonogashira coupling of TMSA onto 10 followed by removal of the TMS groups. The synthesis of 12, however, proved very difficult in our hands, and while a product was obtained the analytical data showed that it consisted of oligomers and other impurities. SEC revealed many different components as shown in Figure 2. We decided to purify the crude product by preparative SEC and isolated the purified fraction of 11. The similarity between the 1H NMR spectra for the crude fraction and the purified product was striking, and it was virtually impossible to establish the extent of impurities from the 1H NMR spectra alone, so a technique like SEC is absolutely needed.

We propose that the large molecular weight impurities are oligomers of 12 to account for the similarities in the 1H NMR spectra and for the fact that the UV spectra of the fractions were very similar and contained the signatures of a zinc porphyrin. The purified fraction containing mainly 11 could be deprotected followed by recrystallization from chloroform, and the product was filtered using a 0.45 μm filter to give pure 12. Similar
Zn Porphyrin Linked Poly(phenyleneethynylene) 1183

Scheme 2. Synthetic Procedure Leading to the Homopolymer Poly[(2,5-dioctyltolanyl)Ethynylene] \([N_n]^{*}\)

Active monomer

* (a) K₂CO₃, MeOH, CH₂Cl₂; (b) Pd(PPPh₃)₄, CuI, piperidine.

| Table 1. Data from the SEC and UV–Vis Analysis for All Three Polymers | 
|------------------------|---------------------|
| compound               | \(M_w\) (g mol⁻¹) | PDI | absorbance ratio | N/P ratio | cross-linkage \((\text{UV–vis } M_c)\) |
| \(N_n\)                | 92 300             | 2.90 | 2.5             | 8.5       | 10.0 |
| NPN A crude            | 19 299             | 2.12 | 2.5             | 8.5       | 10.0 |
| fraction 1             | 66 020             | 1.18 | 2.5             | 8.5       | 10.0 |
| fraction 2             | 41 255             | 1.16 | 3.5             | 10.0      | 12.0 |
| fraction 3             | 22 175             | 1.15 | 2.5             | 8.5       | 10.0 |
| fraction 4             | 19 058             | 1.18 | 2.5             | 8.5       | 10.0 |
| fraction 5             | 8 372              | 1.11 | 2.5             | 8.5       | 10.0 |
| fraction 6             | 4 727              | 1.09 | 2.5             | 8.5       | 10.0 |
| fraction 7             | 2 901              | 1.00 | 6.9             | 10.0      | 12.0 |
| NPN B crude            | 36 949             | 4.09 | 10.0            | 3.2' (n.a.) | 12.0 |
| fraction 1             | 95 275             | 1.10 | 6.5             | 22.2      | 9.1  |
| fraction 2             | 60 144             | 1.42 | 5.5             | 17.1      | 7.5  |
| fraction 3             | 31 612             | 1.28 | 4.5             | 15.4      | 4.3  |
| fraction 4             | 14 225             | 1.19 | 3.5             | 12.0      | 2.4  |
| fraction 5             | 9 120              | 1.24 | 2.5             | 8.5       | 2.1  |
| fraction 6             | 4 074              | 1.30 | 2.5             | 6.9       | 1.1  |
| fraction 7             | 1 534              | 1.17 | 1.5             | 5.1       | 1.1  |

* The absorbance ratio is the ratio of the absorbance from the \(N_n\) polymer (at 391 nm) and the porphyrin (at 461 nm). The N/P ratio listed in this table is the ratio between the N and P units based on their individual extinction coefficients. (see the section Method B: Polymerization in the Presence of Dilbereoporphyrin for the results). Molecular weights of 425 and 750 g mol⁻¹ were used respectively for the monomer and the porphyrin constituents in the polymer. Estimated from the increase in molecular weight from 11 546 to 36 949 g mol⁻¹ on standing. A value for cross-linking of 0.5 is obtained since the extinction coefficient for N is underestimated at low molecular weights. A value <1 has no physical meaning in terms of cross-linkage; it has therefore not been shown.

problems have been described earlier by Tomizaki et al., where a seemingly pure product in terms of porphyrin content (NMR and UV–vis data) turned out to be a mixture of molecules with different size (i.e., different molecular weights) when SEC was used. This could be oligomers or higher porphyrin components (gel-like molecules).

Homopolymer Synthesis. Scheme 2 shows the synthetic procedure of the conjugated homopolymer. Employing Sonogashira cross-coupling conditions following standard methods, the iodine-terminated homopolymer poly[(2,5-dioctyltolanyl)ethynylene] \([N_n]\) was obtained. We have used Pd(PPPh₃)₄ as catalyst instead of Pd(PPPh₃)Cl₂ to minimize the formation of the butadiyne byproducts, where two terminal acetylenes are coupled when reducing Pd(II) to Pd(0). Pd(0) is most probably the active species in the catalytic cycle for the Sonogashira cross-coupling reaction. The molecular weight \(M_w\) of the polymer \(N_n\) was determined using SEC, and the results are listed in Table 1. Measurements on electroteactive devices prepared using \(N_n\) showed that it contained residual palladium, which were confirmed by inductively coupled plasma sector field mass spectrometry (ICP-SFMS) measurements. The content of Pd in the \(N_n\) polymer was determined to 49.3 mg kg⁻¹. The Pd was removed by using DDC (see the section Problems with Pd).

Porphyrin Linked Homopolymer Synthesis. We attempted two different strategies to synthesize a molecule with the general structure NPN, as shown in the lower part of Figure 1. While the two different procedures shown in Scheme 4 both accord with the general structure depicted above, they differ in the way the polymer segments are coupled to the porphyrin nucleus. The only difference is the distance from the porphyrin to the first phenyl ring bearing octyl substituents. The proximity of the octyl substituents in the case of NPN A reduces the degree of conjugation between the polymer segment and the porphyrin nucleus due to steric effects.

Therefore, there would be a difference between the electronic properties of the NPN A polymer and the NPN B polymer. This was confirmed by the differences in the UV–vis spectra for the two products.

Method A: Polymerization in the Presence of Diethynylporphyrin. Our first choice of porphyrin derivative was the diethynylporphyrin 12 since we desired a reactivity of the porphyrin derivative that was comparable or larger than the individual monomers to ensure attachment of the porphyrin molecules to the growing polymer chains during the polymerization reaction.

The polymerization reaction of the monomer 8 in the presence of the deprotected diethynylporphyrin 12 proceeded smoothly, giving a polymer product. The evidence for zinc porphyrin content in the polymer was found by the UV–vis spectrum of the product during SEC analysis. We employed fractionation of the crude polymer product using preparative SEC and obtained seven fractions. The UV–vis spectra of the fractions are shown in Figure 3, and while they show a gradual red shift of the absorption maximum for the polymer component with increasing molecular weight (as expected) the porphyrin content was roughly constant. The ratio of the N and P absorption and the molecular weight \(M_w\) determined by SEC in the seven fractions are listed in Table 1. The ratio between the porphyrin and polymer showed little variation and did not depend on the molecular weight as expected, if only one porphyrin molecule had been incorporated into each NPN molecule. These results could indicate that not only the NPN A polymer was produced but also a large amount of the buta-1,3-diyne-1,4-diyl byproduct derived from oxidative dimerization (see Scheme 5).

To account for this finding, we assume that a side reaction involving the ethynyl groups of either the growing polymer chain or the porphyrin takes place. In the case of the polymerization starting from the diethynylporphyrin the growing polymer chain will always expose an ethynyl group and thus be susceptible to oxidative coupling of the ethynyl group to give a butadiyne, which in principle gives a homopolymer.
Scheme 3. Two Different Synthetic Producers Leading to the Three-Domain Structure NPN*  

A) 

\[ \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \]

A \rightarrow \text{Zn} \quad \text{N} \quad \text{N} \quad \text{Cp}^* \text{Cp}^{17}

\[ \text{N} \quad \text{N} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \]

B) 

\[ \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \]

B \rightarrow \text{Br} \quad \text{Br} \quad \text{N} \quad \text{N}

\[ \text{N} \quad \text{N} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \quad \text{Cp}^* \text{Cp}^{17} \]

* (a) Pd(PPh₃)₄, CuI, Et₃N, THF.

The segment with a defect in the growth direction such that both polymer termini present iodine atoms. Such an event, however, does not represent a termination of chain elongation as a monomer or another diethynylporphyrin can react at the iodinated chain end hence continue chain growth. The result is a polymer product with an even distribution of porphyrin molecules with the amount of porphyrin determined largely by the ratio of polymer to monomer in the initial reaction mixture. In conclusion, the synthetic scheme
for the incorporation of a single porphyrin molecule between two polymer blocks based on a diethynylporphyrin does not work because of the dimerization side reaction that always takes place during the Sonogashira reaction.

From a synthetic point of view, a small molecule product is generally not affected by a small amount of dimerization (of the order of 1-10%) as means of purification are available (i.e., distillation, crystallization). When employing the Sonogashira reaction for polymerization, however, the defects introduced by the side reaction are accumulated in the polymer product and make the applicability of the Sonogashira reaction for polymerization questionable, if a product with a high degree of regioregularity or directionality is required.

Figure 3. UV–vis spectra of the seven fractions of the NPN A polymer formed during the polymerization of the monomer 8 in the presence of the diethynyl porphyrin 12. The spectra are normalized according to $I_{max}$ of the polymer. It can be seen how the absorption of the porphyrin ($i = 451$ nm) does not vary significantly in intensity as a function of the length of the NPN polymers.

Method B: Polymerization in the Presence of Dibromoporphyrin. In this method we used dibromoporphyrin, 10, as the source of porphyrin during the polymerization reaction. Seen in light of the findings above, we would expect only one porphyrin molecule to be incorporated in each polymer chain assuming that 10 does not lead to homocoupled byproducts. We would also expect a considerable amount of homopolymer since any dimerization of a pair of ethynyl groups will lead to homopolymers with halogen termini in both ends of the molecule that does not allow for the incorporation of the dibromoporphyrin. From the SEC trace of the crude product it was clear that the synthetic scheme worked in terms of a decreasing porphyrin content with increasing molecular weight. The SEC trace also confirmed the presence of porphyrin in molecules of high molecular weight but revealed nothing about the homopolymer content.

The crude polymer product had a molecular weight ($M_w$) of 11 546 g/mol and a polydispersity of 3.456. Fractionation of the crude product into seven fractions
using preparative SEC as above revealed as expected a variation of the porphyrin content in the individual fractions. When SEC analysis was performed on the individual fractions the following day, it was surprising that the molecular weight of the fraction had increased, as if cross-linking of the products took place. The fractions were then left to stand and obtained for the hood in air for 3 days to allow for completion of the cross-linking process. The resulting solids were boiled in chloroform that dissolved only a part of the polymer product. This finding is consistent with earlier reports on the insolubility of poly(phenyleneethynylene)s once having been subjected to complete drying. The soluble fraction was then refraccionated using preparative SEC, and this gave stable molecular weight fractions. It should be noted that the molecular weight of the crude polymer was increased to 36 949 g/mol with a polydispersity of 4.09 during the 3 days. The fractions show the desired variation of the porphyrin content between the individual fractions (see Figure 4). The repurified fractions were used for the emission spectrosopy studies. The UV–vis solution spectra of the seven fractions were recorded, and from these spectra it can be seen that the absorption maximum, λ_{max}, as a function of the polymer chain length was red-shifted 9 nm from fraction 7 through fraction 1. From Figure 4 it is observed that the red shift of λ_{max} already stops at fraction 5, which corresponds to an approximately a decain of the N_6 in NPN B. These results are in agreement with the results of Jones et al., who reported studies on oligomers at a precise length of 1,4-phenylene ethynylene. Bredias et al. have shown that the band gap as a function of n^{-1} (for large n > 5) has a linear relationship. The average ratio of N and P in the NPN polymer is obtained on the basis of the measured λ_{max} at 390 nm (absorption mainly from N_6) and 451 nm (absorption mainly from P). The ratio for the N_6 and P absorption in the fraction are listed in Table 1 together with the molecular weight (M_6) determined by SEC. From the extinction coefficient for N_6 and P (we used the ε_{295} in 12 for the absorption at 455 nm) an estimate for the ratio between N and P can be obtained. The two extinction coefficients were determined to be ε_{295} = 70 000 M^{-1} cm^{-1} and ε_{295} = 240 000 M^{-1} cm^{-1}. This indicates a N/P ratio of 23.2:1, 17.1:1, 15.4:1, 12.1:1, 8.5:1, 6.4:1, and 5.1:1 for fractions 1 to 7, respectively (see Figure 4).

A comparison of the molecular size obtained by UV–vis with the results from the SEC analysis shows a mismatch in both high and low molecular weights. In the low molecular weight range the mismatch is a result of the estimate that the extinction coefficient of both N_6 and P is the same in the polymer and for the isolated molecules. In the high molecular weight range the mismatch is due to cross-linking since more than one porphyrin molecule would be present in the cross-linked supernaules. This means that the molecular weight (M_6) determined by SEC should be given by M_6 = x(N/P) M_{monomer} + M_{porphyrin}, which is in good accordance with our observations (see Table 1). In the equation x is the number of cross-link. N/P is the ratio between the N_6 monomer and the porphyrin obtained by the UV–vis measurements, and M_{monomer} and M_{porphyrin} are the molecular weights monomer and porphyrin, respectively. Further, SEC has a tendency to overestimate the molecular weight of linear rigid molecules when using polystyrene standards. SEC probes the hydrodynamic volume of the polymer, and since polystyrene can be assumed globular in chloroform solution and the polymer systems studied remain linear, this will contribute to the overestimation as the molecular weight increases. This effect has not been taken into account in Table 1. The average of the cross-linking, x, in Table 1 has a value of 3.8 for the seven fractions, which is in good accordance with the value of 3.2 obtained for the molecular weight increase of the homopolymer on standing (not corrected for the weight of the individual fractions).

Problems with Pd. The problems concerning the removal of Pd catalyst from samples is a large forgotten area in the literature, which only very few authors have reported. Earlier work with palladium-catalyzed polymerization using the Heck reaction for the polymerization has unfortunately shown that the removal of palladium catalyst remnants can be problematic. It is probably difficult to envisage complete removal of catalyst remnants in the case of polymeric products. With this paper and two recent ones we hope to start a discussion on this topic. It is very important to be able to remove all of the Pd catalyst, when e.g. a photovoltic or a light-emitting device of conjugated polymers is produced, because remnants from the Pd catalyst in the form of nanoparticles can lead to a short circuit of the device.

From our experience with the palladium-catalyzed Heck reaction, we would expect some remnants of palladium in both the N_6 and the NPN (both A and B) polymer. The remnants of Pd in the N_6 were confirmed by ICP-SFMS and determined to be 49.3 mg/kg. We had hoped that the remnants of palladium did not influence the electrical and optical properties of the polymers too much. Unfortunately, when we made a standard electroactive device of the polymers produced by sandwiching the polymer between two electrodes (see Supporting Information) the resistance was as low as 30 Ω.

In connection to removing Pd from our polymers, we have attempted to use diethylammonium diethylthiocarbamate (DDC) with some success as reported by Jones et al. for solid-state synthesis. The reason for choosing DDC is that it coordinates very strongly to Pd (log K (Pd[DDC]) = 64.9) and more strongly than the alkyne groups in the polymers. Furthermore, Briscoe and Humphries have showed that the coordination of DDC to Pd is very fast and that the coordination process is complete after 5 min at room temperature. There were several indications that DDC coordinated to Pd as
expected and thereby removed Pd from the polymers. The indications were that the solution shifted color from yellow to dark brown very quickly (in a few seconds) upon addition of DDC. Furthermore, the precipitated polymer product after the treatment with DDC had a lighter yellow color than the crude polymer product; the resistance through a standard electroactive device geometry (produced by sandwiching the polymer between two electrodes) increased from 30–50 Ω to 0.3–2 MΩ after the treatment with DDC. Our results indicate that DDC very effectively removed or passivated Pd from polymer samples. Unfortunately, there were also some indications that DDC reacted with the polymers.

We observed that treatment with DDC, for periods of time longer than 5 min (we attempted 2 and 24 h), led to a dramatic change in some of the properties of the polymers. The polymer became difficult to precipitate from in methanol, and no electroluminescence could be observed when a potential was applied. On the other hand, we had success in removing Pd from the polymer samples without the above-mentioned changes in properties for the polymers, when we used approximately 5 times excess of DDC, compared to the amount of Pd used in the catalyst, and a reaction time between 3 and 5 min. From the UV–vis spectra of the N₈ in the solid state (see Supporting Information) it can be seen that the optical properties have changed dramatically, when the polymer was treated with DDC for more than 5 min. It should be emphasized that there is little change in the UV–vis spectrum of the polymer, which have been treated with DDC for 5 min compared with the crude polymer. It was not possible to observe the dithiocarbamoyl groups in the ¹H NMR spectra recorded on the DDC treated N₈ polymer product. By comparing the fluorescence spectra of the N₈ polymer in solid state before and after a long DDC treatment time, it can be seen that there were significant changes (see Supporting Information). The changes cannot however explain why the electroactive devices after long DDC treatment did not show electroluminescence. The quantum yield of fluorescence is approximately constant; there is however a small blue shift of the emission maximum upon DDC treatment and a shoulder at 415 nm that becomes suppressed.

An elementary analysis of the N₈ polymer after a 2 h DDC treatment time shows that the polymer contains C, 84.8%; H, 9.1%; N, 0.5%; and S, <0.1% compared with C, 88.4%; H, 9.5%; N, <0.3%; and S, <0.1% for the untreated N₈ polymer. The elementary analysis indicates that it is not the DDC group itself that is added to the polymer, since no S could be detected in the polymer. However, we detected a little increase in the contents of N corresponding to one N atom for every six incorporated monomer units. From the elementary analysis we get an indication that it is an N-containing compound that is attached to the polymer by treatment with DDC. While we cannot provide a solid explanation for the reaction between DDC and the polymer, our photophysical and electroactive device measurements show that DDC treatment changes the properties of the polymer. The NPN B polymer has also been treated with DDC with success; electroluminescent devices of the N₈ itself and of NPN B polymer were prepared, and the nature of the electroluminescence has been characterized. To decide whether the apparent cross-linkage of the NPN B polymer products could be a result of the content of Pd in the polymers, we performed a search in the literature for a useful model study and found that it has been known for more than 50 years that transition metals catalyze the trimerization of alkenes. In 1997 and 2001, it was further shown by Gervogyan et al.that substituted benzenes can be constructed from enynes and activated alkynes in a [4 + 2] and from
terminal alkyne in a $[2 + 2 + 2]$ fashion in the presence of a Pd(0) catalyst. Both reactions proceeded under normal Sonogashira conditions without an inert atmosphere. By comparing these model studies of the trimerization of alkyne with the known byproducts from a Sonogashira cross-coupling reaction, it is most likely that the apparent cross-linkage of our polymer products are formed by a trimerization process. A probable trimerization process could be that two homopolymers $N_k$ with terminal alkyne (that are not coupled to the porphyrin) are forming an enyme in the presence of the Pd(0) catalysts. This enzyme can then react with one of the triple bonds in both $N_k$ and NPN B molecules to form the trimerization cross-linkage (see Scheme 6). This can be used to explain how the molecular weight of a polymer fraction can increase when standing and ultimately form an insoluble product with a high degree of cross-linking. To avoid this trimerization cross-linkage of the polymer, it is very important to remove all the catalyst during precipitation of the polymers.

**Fluorescence Measurements.** We have performed fluorescence measurements on the $N_k$ and the NPN B polymers. In Figure 5 normalized fluorescence spectra of the seven fractions of NPN B are shown. In all cases the excitation wavelength was set to the maximum absorption of the polymer (around 388 nm). The emission below 600 nm is from the N domain, whereas the porphyrin unit emits around 660 nm. The polymer emission from the seven fractions shows little variation. However, fractions 6 and 7, with the shortest N domains, differ somewhat from the remaining fractions: both spectra are more intense at 440 nm. In addition, the spectrum of fraction 7 is blue-shifted 8 nm compared to the remaining spectra, probably due to the relatively short chain segments (oligomers). Another notable feature is that the polymer emission extends more toward long wavelengths as the N domains get larger as expected. The most interesting difference is that the porphyrin emission seems to gradually decrease as the N domain gets shorter (see inset in Figure 5). We interpret this as a consequence of the fact that the whole polymer chain is probably not perfectly conjugated. Hence, the chain will consist of a number of segments. Thus, with increasing polymer length the more likely it becomes that an exciton gets trapped on a segment that is not in the porphyrin vicinity, and the energy transfer efficiency decreases. Since the N domain fluorescence in the spectral area where the P domain absorbs, a Förster type of energy transfer may be involved. Using the same methodology as described earlier, photon balances could be derived for the individual fractions.

**Figure 6.** Typical photon balance of the NPN B assembly in dilute degassed chloroform solution. The absorption is arbitrarily set to 1000 photons, and the fluorescence emission was measured. Energy transfer as well as nonradiative decay is calculated from the fluorescence quantum yields of the pure N and P domains. The relative porphyrin emission increases as the N chain length decreases.

**Figure 7.** Normalized fluorescence emission from spin-coated films of the $N_k$ polymer ($M_w = 92,000$ g mol$^{-1}$) and the NPN B assembly. The polymer emission is almost completely quenched by the porphyrin.

The fluorescence quantum yields for the pure N domain and pure P domain were determined to be 87% and 10.5%, respectively, using DPA as reference. Assuming that the ratio between the fluorescence rate and the nonradiative decay rates are similar in the NPN B assembly as in the model compounds, a photon balance can be calculated from the measured fluorescence emission. This is shown in Figure 6. The overall quantum yield for the NPN B assembly was in general around 60%, which mainly was caused by the N domain emission. The fluorescence from the P domain was limited compared with the N domain. The relative emission of the P domain was found to increase 5-fold going from fraction 1 to 7. This means that in solution (or in a diluted matrix) the polymer length may be crucial in multidomain light-harvesting systems.

Solid-state studies of NPN B shows that whereas energy transfer is limited in solution, it becomes the main deactivation route of the N domain in the solid phase (see Figure 7); thus, N emission is nearly completely quenched by the porphyrin that emit strongly. Thus, in the solid state the chain length is much less important than in solution with regards to efficient energy transfer. The more efficient energy transfer in the solid state seems to be a general rule for this kind of polymer-dye systems. The N and P emission are red-shifted by approximately 65 nm compared to the solution spectra. Also, the relative intensity of the N domain and P domain peaks are changed.

The lifetime decay of the pure P domain could be fitted with a single-exponential function, with a decay of 2.4 ns in degassed chloroform. The lifetime decay of the N domain needed a linear combination of two
exponential functions to obtain a reasonable fit in degassed chloroform: 259 ps (86%) and 1.6 ps (14%). In the spin-coated film the measured lifetime of the N-doped polymer is significantly larger: 1.4 ns (53%) and 2.7 ns (47%). This could be caused by the hampering of some of the excitons to recombine due to the surface roughness. A side effect is that the exciton has more time to be transferred to a porphyrin unit, and this could be one of the reasons why more efficient energy transfer is observed. Interestingly, the fluorescence lifetime of the porphyrin unit was significantly shorter in the solid state than in solution. The lifetime decay was complex, and a good fit could not be obtained. This is not unexpected since a number of interactions and different local environments are present in the solid NPB matrix. The average lifetime was approximately 300 ps.

Conclusions

We have presented the synthesis of a new zinc porphyrin linked conjugated polymer. We experienced problems in controlling the nature of polymer product and found that it was a result of many factors: oxidative dimerization and cross-linking of the ethynylene groups, and palladium remaining from the catalyst. We thoroughly studied products and the reaction by two different synthetic approaches and demonstrated that a polymer product with one porphyrin molecule inserted in each polymer chain could be obtained by choosing the right synthetic approach. We separated the polymer material into fractions using preparative SEC and studied both the cross-linking of the individual fractions and further to finally perform thorough photophysical studies. We showed the light-harvesting properties of the three-dimension structure as a function of the chain length of the polymer. This new zinc porphyrin linked homopolymer shows a very effective energy transfer from the polymer to the porphyrin in solid state. Finally, we have identified some problems concerning remnants of Pd from the catalyst in the polymer product, and devise a procedure for the removal of the Pd remnants such that standard electroactive devices could be constructed.

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Supporting Information Available: General analytical techniques, synthetic procedures, characterization, UV–vis, and fluorescence spectra of the polymer and the DDC-treated polymer, and electroactive device preparation. This material is available free of charge via the Internet at http://pubs.acs.org.

References and Notes

Supplementary information

Thin layer chromatography (TLC) was carried out using Merck Silica (Kieselgel) 60F_{254} on aluminium sheets and the eluent was n-heptane/1,2-dichloroethane in ratio 5:1 unless otherwise stated. To purify the compounds both flash, short column and preparative liquid chromatography (SEC) was carried out. The column material for the flash column chromatography was Merck Kieselgel 60 (particle size 0.015-0.040 mm) and for the short column chromatography both Aluminium oxid aktiv, neutr., type 507c, korrngr. ~ 150 mesh, 58 Å and Kieselgel, korrnr. 70-230 mesh, 60 Å (both from Aldrich) were used. GC/MS was used to detect compounds with a mass up to 700 g/mol, and MALDI-TOF-MS with a DHB matrix and positive ion detection was used to detect compounds with a larger mass. Both TLC, GC/MS and MALDI-TOF-MS were used to monitor the reactions. All reagents were commercially available and reagent grade.

1,4-dibromo-2,5-dioctylbenzene (1): The following describes a modified procedure compared to the literature. 1,4-dibromo-2,5-dioctylbenzene was prepared from 1,4-dioctylbenzene by a simple addition reaction of bromine. 1,4-dioctylbenzene (0.42 mol, 127.55 g) was mixed with 0.3 g iodine (acts as a catalyst). The bromine (0.99 mol, 50.8 mL) was added drop wise over a period of 2.5 hour. The reaction is exothermic, and the mixture became warm during the addition of the bromine. The hydrogen bromide gas that evolved was collected in a water trap. After the addition of the bromine, the mixture was stirred overnight. The following day the product had crystallized and was dissolved in dichloromethane (900 mL) and washed with water (4 x 200 mL) and finally with 1M Na_{2}S_{2}O_{3}(aq). The light yellow organic solution was dried using MgSO_{4}, filtered and evaporated to give a light yellow oil (161.0 g). The product was purified by recrystallization from ethanol (2.5 L). Yield: 120.0 g (62%) as colourless crystals. M. p. 321-322 K. \(^{1}\)H NMR (250 MHz, CDCl\textsubscript{3}, 300K, TMS) \(\delta\): 0.88 (doublet of t, 6H), 1.21-1.42 (m, 20H), 1.57 (m, 4H), 2.63 (doublet of t, 4H), 7.35 (s, 2H); \(^{13}\)C NMR (62.9 MHz, CDCl\textsubscript{3}, 300K, TMS) \(\delta\): 14.80, 23.37, 29.92, 30.05, 30.07, 30.51, 32.58, 36.23, 123.77, 134.44, 142.03. Anal. Calcd. for C\textsubscript{22}H\textsubscript{36}Br\textsubscript{2}: C: 57.40, H: 7.88. Found: C: 57.02, H: 7.86.

General procedure for the bromine-iodine-exchange (2 & 7): This procedure is modified compared to the literature. THF was cooled to 195 K in a 1-litre three-necked round-bottom flask. After the THF had reached the temperature of 195 K, n-BuLi (typically 10 % excess) was added. The solution was cooled back to 195 K and a solution of the bromo-compound in dry THF was added dropwise over a period of 2.5 hour, at such a rate that the temperature did not exceed 223 K. After the addition the temperature was allowed to reach 195 K, and I\textsubscript{2} (10 % excess) was added. The temperature of the mixture was now allowed to reach room temperature and the solution was stirred for 1.5 hour. Light petroleum was added and the mixture was washed with a 1 M solution of Na\textsubscript{2}S\textsubscript{2}O\textsubscript{3}(aq) (3 x 500mL). The organic phase was separated, dried with MgSO\textsubscript{4}, and the solvent was removed by evaporation. The product was purified further by recrystallization in ethanol or methanol.

1-bromo-2,5-dioctyl-4-iodobenzene (2): The reaction of compound 1 (0.163 mol, 75.04 g) in THF (200 mL) with n-BuLi (160 mL of a 1.1 M solution in hexane, 0.11 mol) and I\textsubscript{2} (0.22 mol, 55.85 g) in THF (300 mL) gave, after recrystallization from ethanol (4 L), 65.0 g (78%) of colourless crystals. M. p. 318.5-319 K. \(^{1}\)H NMR (250 MHz, CDCl\textsubscript{3}, 300K, TMS) \(\delta\): 0.88 (doublet of t, 6H), 1.17-1.44 (m, 20H), 1.44-1.65 (m, 4H), 2.61 (doublet of t, 4H), 7.32 (s,
1H), 7.61 (s, 1H); $^{13}$C NMR (62.9 MHz, CDCl$_3$, 300K, TMS) $\delta$: 14.80, 23.36, 29.92, 30.00, 30.06, 30.54, 30.84, 32.57, 36.01, 40.79, 99.51, 125.17, 133.48, 140.99, 142.23, 145.37. Anal. Calcd. for C$_{22}$H$_{36}$BrI: C: 52.08, H: 7.15. Found: C: 52.67, H: 7.23.

1-iodo-2,5-dioctyl-4'-trimethylsilylethynyltolane (7): The reaction of compound 6 (3.87 $\times$ 10$^{-1}$ mol, 22.38 g) in THF (100 mL) with n-BuLi (38.7 mL of a 1.1 M solution in hexane, 0.043 mol) and I$_2$ (0.045 mol, 11.31 g) in THF (200 mL) gave 22.0 g (91%) of light red oil. The product was used without further purification. $^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.25 (s, 9H), 0.77-0.93 (doublet of t, 6H), 1.15-1.47 (m, 20H), 1.47-1.72 (m, 4H), 2.56-2.78 (m, 4H), 7.29 (s, 1H), 7.42 (s, 4H). $^{13}$C NMR (62.9 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.61, 14.80, 23.35, 29.95, 30.05, 30.12, 30.22, 30.90, 31.34, 32.58, 34.53, 34.53, 40.90, 90.46, 93.80, 97.02, 101.89, 105.33, 123.17, 123.69, 124.12, 131.88, 132.62, 132.93, 140.22, 143.51, 144.79.

General procedure for Sonogashira cross-coupling (3 and 6). The procedure for the coupling reaction between terminal acetylenes and sp$^2$-carbon halides is the same as described in the literature$^{3,4}$ with a few modifications. A 500 mL round-bottom flask was dried using a heat gun and flushed with argon. Then a mixture of the terminal acetylene (5% excess) and the sp$^2$-carbon halide was dissolved in triethylamine (2.5 L per mole halide). After the reagents had dissolved, the solution was flushed with argon for 30 minutes, before the catalyst Pd(PPh$_3$)$_2$Cl$_2$ ($\approx$ 2 mol %) and the co-catalyst CuI ($\approx$ 10 mol %) were added under an argon atmosphere. After the addition of the catalyst the mixture was stirred under an argon atmosphere for 9 hours. GC/MS and TLC were used to monitor the reaction. After the 9 hours more catalyst and acetylene were added and the reaction was continued until the TLC showed that the reaction was done. This is important because a mixture of e.g. 1-bromo-4-iodobenzene and 1-(4-bromophenyl)-2-trimethylsilylacetylene were found difficult to separate. When the reaction was complete diethyl ether was added (the same volume as the triethylamine), and the suspension was filtered through a glass filter funnel (triethylaminehydrogeniodide was removed). The filtrate was collected and evaporated to dryness by a rotary evaporation. The liquid residue (usually an oil) was dissolved in n-heptane and then purified on an aluminium oxide column with n-heptane as the eluent. The n-heptane was removed by evaporation to give the product.

1-(4-bromo-2,5-dioctylphenyl)-2-trimethylsilylacetylene (3). Compound 3 was prepared from Compound 2 (3.96 $\times$ 10$^{-2}$ mol, 20.07g) and trimethylsilylacetylene (4.20 $\times$ 10$^{-2}$ mol, 4.12 g). The reaction was performed in an ice bath to increase the selectivity between the iodo and the bromo places. The reaction was completed after 9 hours. The yield was 17.95 g (95%) of clear light yellow oil. $^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.25 (s, 9H), 0.80-095 (dt, 6H), 1.15-1.42 (m, 20H), 1.47-1.68 (m, 4H), 2.61 (doublet of t, 4H), 7.26 (s, 1H), 7.33 (s, 1H); $^{13}$C NMR (62.9 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.65, 14.79, 23.37, 29.93, 30.00, 30.10, 30.16, 30.29, 30.58, 31.25, 32.59, 34.80, 36.26, 98.93, 104.01, 122.39, 125.45, 133.43, 134.33, 140.03, 145.35. Anal. Calcd. for C$_{27}$H$_{45}$BrSi: C: 67.89, H: 9.5. Found: C: 67.88, H: 9.58.

1-bromo-2,5-dioctyl-4'-trimethylsilylthynyltolane (6). Compound 6 was prepared from compound 4 (2.57 $\times$ 10$^{-3}$ mol, 1.04 g) and compound 5 (2.60 $\times$ 10$^{-3}$ mol, 0.78 g). The product was purified on a flash column with n-heptane as eluent. The yield was 0.50 g (34%) of a light yellow oil. The low yield is caused by the importance in this step to separate the by-product from the main product. From TLC it was known that there were four products, with following R$_f$ values: 0.58, 0.38, 0.25 and 0.08. By $^1$H and $^{13}$C NMR and MALDI TOFF it were
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determined that the first product mainly consisted of compound 4, the second was compound 6, the third consisted mainly of 1-bromo-2,5-dioctyl-2',5'-dioctyl-4-(4''-trimethylsilyl-ethynylphenyl)ethynyltolane (B2), and the fourth of 1,4-dioctyl-2,5-bis((4''-trimethylsilyl-ethynylphenyl)ethynyl)benzene (B3).

$^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.27 (s, 9H), 0.83-0.97 (doublet of t, 6H), 1.17-1.47 (m, 20H), 1.53-1.75 (m, 4H), 2.61-2.84 (m, 4H), 7.33 (s, 1H), 7.39 (s, 1H), 7.44 (s, 4H); $^1$C NMR (62.9 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.61, 14.78, 23.36, 29.93, 29.95, 30.09, 30.11, 30.14, 30.20, 30.56, 31.29, 32.58, 34.73, 36.29, 90.37, 93.57, 97.02, 105.35, 122.26, 123.70, 124.17, 125.55, 131.86, 132.63, 133.57, 134.13, 140.23, 144.86. Anal. Calcd. for C$_{35}$H$_{49}$BrSi: C: 72.76, H: 8.55. Found: C: 73.14, H: 8.63.

General procedure for the deprotection of the trimethylsilyl group (4 and 8). The deprotection of the trimethylsilyl group was performed as described in the literature with minor modifications. In a round-bottom flask the silylated compound was dissolved in a 1:1 dichloromethane:methanol mixture (8 L per mole) and flushed with argon in 30 minutes before potassium carbonate ($\approx$ 10 equiv) was added under an argon atmosphere. The mixture was stirred at RT and followed by NMR and GC/MS. While shorter reaction times have been reported, the typical reaction time was 1-2 days. At completion the solvent was removed by evaporation. The excess potassium carbonate was eliminated by filtration. The products were used without further purification.

4-bromo-2,5-dioctylphenylacetylene (4). Compound 4 was prepared from compound 3 (6.67 $\cdot$ 10$^{-2}$ mol, 31.87 g). Yield 26.60 (98 %) g as a red oil. $^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.78-0.95 (doublet of t, 6H), 1.16-1.48 (m, 20H), 1.50-1.70 (m, 4H), 2.57-2.78 (m, 4H), 3.25 (s, 1H), 7.30 (s, 1H), 7.37 (s, 1H); $^1$C NMR (62.9 MHz, CDCl$_3$, 300K, TMS) $\delta$: 14.79, 23.37, 29.92, 30.07, 30.09, 30.51, 31.14, 32.58, 34.45, 36.23, 81.55, 82.54, 121.37, 125.81, 133.47, 134.82.

1-iodo-2,5-dioctyl-4'-ethynyltolane (8). Compound 8 was prepared from compound 7 (1.68 $\cdot$ 10$^{-3}$ mol, 1.05 g). The yield was 0.750 g (81 %) of a yellow oil. $^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 0.79-0.96 (doublet of t, 6H), 1.14-1.45 (m, 20H), 1.50-1.77 (m, 4H), 2.60-2.79 (m, 4H), 3.18 (s, 1H), 7.30 (s, 1H), 7.46 (s, 4H), 7.68 (s, 1H); $^1$C NMR (62.9 MHz, CDCl$_3$, 300K, TMS) $\delta$: 14.78, 23.38, 29.71, 29.95, 30.06, 30.13, 30.22, 30.92, 31.34, 32.59, 34.54, 40.92, 79.60, 83.97, 90.55, 93.60, 101.95, 122.69, 123.14, 124.60, 131.99, 132.81, 132.96, 140.27, 143.57, 144.85.

Photophysical results from the DDC treatment
Figure S1: A comparison of the UV-vis spectra for the $N_n$ polymer in solid state as a function of the DDC treatment time. It should be emphasized that for a treatment time in the order of 5 minutes there were no significant changes in the UV-vis spectrum.
Figure S2: The comparison of the fluorescence spectra of the Nₐ polymer before and after a long DDC treatment time in spincoated films.

Standard electroactive device fabrication

The substrates were 10 Ω square⁻¹ ITO slides that had been etched in one end by immersion in a 338 K aqueous solution of HCl(aq) 20% and HNO₃(aq) 5% for typically 60 seconds. The etched substrates were cleaned by immersion in isopropanol and subjected to ultrasound for 10 min. After being blown dry in a stream of dry argon, a layer of PEDOT:PSS was applied by spincoating an aqueous solution of PEDOT:PSS (1.3% wt.) containing sorbitol (2% wt.). The substrates were placed in an oven at 373 °C and ramped up to a temperature of 453 K during 1 h and kept at 453 K for 1 h. Polymer solutions in chloroform at a concentration of 2-10 mg mL⁻¹ were micro-filtered and spin coated at 1500 rpm onto the substrates. The typical film thicknesses were 50-100 nm as measured by a DEKTAK 3030 and the typical absorbance of the films were 0.25-1.0 absorbance units. The active area of the device was 3 cm² (20 mm x 15 mm). The aluminium electrode was evaporated onto the device at a pressure of < 5 × 10⁻⁶ mBar. After application of the electrodes the samples were removed from the evaporator chamber, electrical connections were made using conducting silver epoxy glue.
Figure S3: A schematic illustration of the electroactive device geometry (not drawn to scale).

References


Dye linked conjugated homopolymers: using conjugated polymer electroluminescence to optically pump porphyrin-dye emission

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Abstract

Zinc-porphyrin dye molecules were incorporated into the backbone of a conjugated polymer material by a method, which allowed for the incorporation of only one zinc-porphyrin dye molecule into the backbone of each conjugated polymer molecule. The electronic properties of the homopolymer were established using ultraviolet photoelectron spectroscopy (UPS) for the determination of the electronic energy levels and the injection barrier for holes into the valence band. Pulse radiolysis time resolved microwave conductivity (PR-TRMC) was used to determine the sum of charge carrier mobilities. Electroluminescent devices of the homopolymer itself and of the zinc-porphyrin containing polymer were prepared and the nature of the electroluminescence was characterized. The homopolymer segments were found to optically pump the emission of the zinc-porphyrin dye moieties. The homopolymer exhibits blue-green emission and the zinc-porphyrin linked homopolymers emit near-infrared/infrared light. This was demonstrated to be due to electroluminescence pumping of the zinc-porphyrin moieties that were covalently linked to homopolymer material. When only one zinc-porphyrin dye was incorporated into the backbone of each homopolymer molecule complete suppression of homopolymer electroluminescence was observed with exclusive near-infrared emission.

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Keywords: Dye-linked conjugated homopolymers; Zinc-porphyrin dye containing conjugated polymers; Zinc-porphyrin linked polyphenyleneethylenes; Near-infrared/infrared electroluminescence; Electroluminescence pumping

1. Introduction

Electrical conductivity of doped polyacetylene was shown in 1974 [11]. Since then there has been an increasing activity within the field of conjugated polymers and their application, for e.g. light emitting diodes (LED) [2], polymer electronic circuits [3] and polymer solar cells [4].

The light emitting diodes got their breakthrough, when Burroughes et al. [2] first published a paper about the preparation and light emitting properties of a green LED made from poly(p-phenylenevinylene). After this pioneering report there has been a growing interest in polymer-based LED devices [4–11]. Some of the reasons why there is a large interest in this relatively new discovery compared to conventional inorganic semiconducting devices is due to the potential for low cost, high flexibility and ease of processing [5] of the polymer-based LEDs. Furthermore, it is relatively easy to tune the colour of the light emission by chemical modification [6,8,9].

Reports on polymer-based LEDs with colours from blue to near-infrared (NIR) can be found in the literature [9]. Especially the NIR emitting polymer-based LEDs have attracted a lot of interest in the last couple of years, because they open the possibility for the application of the polymer-based LEDs in telecommunications devices [7,12,13]. The reports about these polymer-based NIR LEDs are still relatively few, and most of these reports are concerned with polymers containing rare earth metal ions as dopants [13], low band gap polymers [9] and blends of polymers and small dye molecules [11]. In the later case the polymer matrix works as the donor and transfers the energy by a Förster-type energy transfer process to the dye molecules (typically an organometallic complex) [11]. The Förster energy transfer process is energy transfer by long-range dipole–dipole coupling. A conceivable way to optimise the energy transfer from the polymer matrix to the dye molecule could be to covalently link the dye molecule to the backbone.
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of the polymer. In this way the energy transfer could still happen via the long-range dipole-dipole coupling, but also through the bonds. Recent studies on the light harvesting and energy transfer properties of dye-terminated conjugated polymers and dye-linked conjugated homopolymers have shown that the light energy harvested by the conjugated polymer segment can be transferred to the dye molecule and either be re-emitted as pumped fluorescence or can increase the photovoltaic response when applied as the active material in a polymer photovoltaic set-up [14, 15].

In this study we demonstrate the principle of transfer of electroluminescence to a dye molecule that is covalently linked to the conjugated polymer backbone and demonstrate how a blue-green emitting polymer can transfer energy to a zinc-pteroporphyrin dye molecule and give almost exclusively near-infrared light emission. The dye-linked conjugated homopolymer we used in these studies have the three-domain structure, NPN, that can be seen in Fig. 1. The N and P domains consist of poly(2,5-dioctyl-tolanyl)-ethylenylene and [10,20-Bis-(3,5-bis-tert-butylphenyl)-5,15-dihromophosphinato]zinc(II), respectively.

We also report the electronic properties of the homopolymer by determination of the electronic energy levels, the injection barrier for holes into the valence band, the ionisation potential as determined by ultraviolet photoelectron spectroscopy (UPS) and the sum of charge carrier mobilities for the N polymer as determined by pulse radiolysis time resolved microwave conductivity (PR-TRMC).

2. Experimental

The polymer materials N and NPN were prepared and purified from the contaminant palladium nanoparticles as described previously [16]. Ultraviolet photoelectron spectroscopy [17] was performed using 50 eV photons in a set-up and by a procedure described in the literature [18]. Carrier mobilities of the homopolymer were measured using the PR-TRMC technique as described in the literature [19] using a set-up described earlier [20]. We employed 10 MeV electrons, pulse lengths of ~300 ns and typical pulse doses of 60 Gy. Electroluminescent devices were prepared by spincoating chloroform solutions onto PEDOT:PSS covered ITO slides giving films with peak absorbencies of ~0.6. The metal electrodes were evaporated at a vacuum of ~1 × 10\(^{-3}\) mbar and consisted of a calcium layer (~30 nm) covered by an aluminium layer (~100 nm). The electrical connections were obtained using silver epoxy. Electroluminescence spectra were recorded in a FLS920 from Edinburgh Instruments. The LED devices had turn on voltages of 3.3 V and were during electroluminescence measurements operated at 5.5 V where they drew a typical current of 3–8 mA cm\(^{-2}\).

3. Results and discussion

3.1. Synthesis

Recently we have reported a synthesis of a new homopolymer and a new zinc-pteroporphyrin linked homopolymer [16]. The homopolymer is the iodine terminated poly(2,5-dioctyl-tolanyl)-ethylenylene (N) (see Scheme 1), and the zinc-pteroporphyrin linked homopolymer NPN is the homopolymer N linked to the pteroporphyrin [10,20 Bis (3,5 bis-tert-butylphenyl)-5,15-dibromophosphinato]zinc(II) (2) (see Scheme 2). N was synthesized by a normal Sonogashira cross-coupling reaction from the 1-iodo-2,5-dioctyl-4'-ethynylolane (1).

The synthetic procedure to NPN allowed for the incorporation of only one zinc-pteroporphyrin dye moiety into the conjugated polymer backbone. Once a pteroporphyrin molecule is incorporated the terminal groups are halogen that does not allow further incorporation dibromozincophosphinates.

Both N and NPN were purified from the contaminant palladium nanoparticles by treatment with diethyldithiocarbamate (DDC) as described previously [16].

3.2. Ultraviolet photoelectron spectroscopy (UPS)

When preparing electroluminescent devices from conjugated polymer materials an important parameter is the position of the electronic energy levels [17] since the matching of the energy levels to the electrode ultimately
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Scheme 2. The synthetic produce leading to the three-domain structure NPN. (b) Fe(PPPh3)3, CuI, Et3N, THF.

Fig. 2. The UPS spectrum for N with an inset (×10) showing the onset of photoelectrons. The distance from the Fermi level, $E_F$, to the onset of photoelectrons (the valence band edge in the material) represent the injection barrier for holes from the substrate into the valence band of the material (left). The electronic band structure derived from the UPS spectrum is also shown (right).

determines the injection barrier for hole carrier injection into the valence band and electron injection into the conduction band during operation. The UPS experiment employs hard UV photons from a synchrotron source. We employed UV photons with an energy of 30 eV that upon interaction with the surface of the polymer material emit a photoelectron from one of the filled energy levels in the material. For this reason UPS can only reveal information on the filled states in the material under study i.e. the valence band. A second parameter that is obtained is the ionisation potential for the material. The UPS spectrum is shown in Fig. 2.

On the right in Fig. 1 the electronic band structure is shown. The distance from the Fermi level to the valence band edge, $E_{V}$, was found to be 1.5 eV and represent the injection barrier for holes into the valence band of the polymer material. The parameters derived from the UPS spectrum are summarized in Table 1.

In terms of electroluminescent devices based on N an injection barrier of 1.5 eV implies that a bias of at least 1.5 V will have to be applied to overcome the injection barrier for holes in the device. There is equally an injection barrier for electrons into the conduction band, but this cannot be measured by UPS as only filled levels are probed by this method. The sum of the hole and electron injection barriers define the value at which electroluminescence can be observed in principle. There are, however, other factors that come into play since electroluminescent light emission through formation of excitons requires a balanced injection of holes and electrons into the polymer in the device.

3.3. Pulse radiolysis time resolved microwave conductivity (PR-TRMC)

A pre-requisite for function of electroluminescent devices is that the conjugated polymer material is able to transport charge carriers. The charge carrier mobility hinges on many physical parameters such as the organization of the molecules, interface phenomena, impurity levels.

<table>
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<th>Table 1</th>
<th>Data from the photoelectron spectra for the polymer I on a gold substrate ($\Phi_{Au} = 4.7$ eV)</th>
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<td><strong>Compound</strong></td>
<td><strong>$E_{V}^B$</strong></td>
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<td>N</td>
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and domain boundaries. An excellent method that allows for the determination of the charge carrier mobility in weakly conducting or semiconducting materials is the PR-TRMC method that is a contactless technique. The technique relies on the transmission of 26–40 GHz microwaves through a sample of the polymer material placed in a shorted waveguide. The reflected microwave signal is then monitored as 10 MeV electrons are passed through the waveguide and the sample. Upon passage some of the high energy electrons creates ionisation events whereby hole-electron pairs are formed. This radiation doping then leads to a large increase in the conductivity that can be measured as a transient change in the microwave conductivity signal. Using fast data collection the transient can be recorded and information on the charge carrier lifetime and on the magnitude of the maximum trap free sum of charge carrier mobilities can be obtained. A major advantage is that no electrical contacts to the sample are needed. This is often the largest problem with other methods of charge carrier mobility measurements techniques such as time of flight methods (TOF) [21] and field effect methods (FEM) [22]. A disadvantage of the PR-TRMC method is that the geometry of the experiment is very far from the device geometry in contrast to the TOF and FEM methods that are highly device oriented. The PR-TRMC method, however, gives the chemist looking for new materials a good overall measure of the material performance without the device-dependent parameters. A plot of the electron pulse and the corresponding microwave conductivity transient is shown in Fig. 3. The electron pulse has a duration of ~300 ns.

From the radiation dose in the pulse the dose normalized conductivity transient can be derived as shown in the lower part of Fig. 3. This gives a measure of the sum of charge carrier mobilities provided that the carrier pair formation energy is known. A value of 25 eV is typically employed as a value for the pair formation energy [23]. A value for the sum of carrier mobilities using 25 eV for the pair formation energy and an end-of-pulse conductivity of $5 \times 10^{-6} \text{s m}^{-1}$ (taken at 100 ns post-pulse)
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gives a sum of charge carrier mobilities, Σμmax = 1.25 × 10^-4 m^2 V^{-1} s^{-1} and a first half-life, \( t_{1/2} \sim 10 \mu s \). The value for Σμmax has not been corrected for the survival probability of the charge carrier pair nor for the differential dose distribution between the conjugated backbone and the alkyl sidechains. This value for the carrier mobility is comparable when compared to values determined earlier for similar compounds [24]. It must be borne in mind that the experimental charge carrier mobilities in an actual device may be much lower due to trapping of carriers at domain boundaries and by impurities.

3.4. Electroluminescence

The electroluminescence of NPN and N along with the fluorescence spectra of spincoted films are shown in Fig. 4. When examining the electroluminescence of the pure N-polymer it is interesting to note that the electroluminescence of the N-domain is red shifted ~100 nm compared to the fluorescence spectrum. The electroluminescence is a broad peak whereas the fluorescence exhibits fine structure. The large red shift of the N-domain indicate emission from a relaxed state. The first thing to notice when examining the electroluminescence of NPN is that it is very similar to the solid-state fluorescence spectrum of NPN. We interpret this as the efficient transfer of an electrically generated excitation in the N-domain to the P-domain that then emit a photon as if it was ordinary fluorescence. Electroluminescence is often distinguished by a considerable redshift in emission maximum from its fluorescence maximum as indeed observed here. In the solid state the N-domain effectively transfers its harvested energy to the porphyrin that fluorescence around 700 nm. Comparing the electroluminescence with the fluorescence it is seen that the N-domain also effectively transfers electroluminescence to the P-domain where virtually no emission from the N-domain can be observed.

The porphyrin emission from electroluminescence devices is a broad peak with a maximum that is slightly blue shifted when compared to the fluorescence emission maximum from thin spincoted films that is a much sharper peak.

4. Conclusion

In this paper, we have presented a characterization of the recently synthesized homopolymer N, with respect to its electronic energy levels and carrier mobilities. We have demonstrated how light emitting devices based on N exhibit blue-green emission. Introduction of a zinc-porphyrin dye into the conjugated backbone of the polymer and subsequent preparation of light emitting devices were shown to emit near-infrared light. We demonstrated that the optical pumping of the zinc-porphyrin moiety arises by excitonic energy transfer from electrically generated excitons in the N-domain to the P-domain.

Acknowledgements

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References

Appendix B

Removal of Palladium Nanoparticles from Polymer Materials

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Palladium catalysis in synthetic organic chemistry has been welcomed as a versatile tool and has made it possible to make complex molecules with many sensitive groups that can be synthesized only under the mild conditions offered by palladium catalysis. The use of palladium in synthetic organic chemistry has led to the development of an arsenal of reaction types that were not possible (or very difficult) without palladium catalysis. Most well known coupling reactions are Heck, Stille, Suzuki, Negishi, and Sonogashira, and while many more exist they are variations over the general theme. An often neglected fact is the formation of palladium nanoparticles during the chemical reactions when the catalyst degrades. When the product of the chemical reaction can be purified by distillation (or for small molecules crystallized), it is often possible to obtain a palladium-free product. For large molecules and polymers, however, these means of purification are not available, and a contamination of the product with palladium should always be assumed. The extent to which palladium nanoparticles are withheld in the product depends on how strong the affinity for palladium is. As an example, palladium nanoparticles bind well to conjugated polymers, leading to a product with palladium that cannot be completely removed using current purification techniques. While the contaminant palladium nanoparticles/catalyst to a large extent can be removed, the crude product typically contains palladium nanoparticles/catalyst. The palladium contamination (0.0001–1% w/w) often goes by unnoticed or undetected, as it does not interfere with common analytical techniques (elemental analysis, NMR, etc.). When, however, the electrical properties of the product are important, even the smallest contamination has a detrimental effect on thin film device performance. We recently discovered the problem for a polyphenylenevinylene) product prepared by two different routes: an elaborated palladium-free route and a one-step palladium route based on the Heck reaction. The polymer products were identical in most physical–chemical aspects (NMR, UV–vis absorption, etc.), but when electroluminescent devices were made using the polymer product from the palladium route, they were not functional whereas the non-palladium product gave functional devices. The lack of functionality was observed as a very low device resistance and the absence of electroluminescence. We later demonstrated the same problem for poly[phenylenevinylene] prepared by a palladium route following Sonogashira conditions. We attempted to remove the palladium contamination using

$N,N$-diethylthiocarbamate (2) and found that we were able to remove the palladium to a level where device fabrication was possible. The shortcoming of 2, however, was that prolonged reaction times altered the photophysical properties of the poly[phenylenevinylene] product to an extent where the desired electroluminescence became suppressed. The device resistance, however, remained high.

The efficient chemical removal of palladium nanoparticles/catalyst is not a trivial task if the desired organic product is to be left unaffected, and while 2 provided a partial solution and has been employed successfully for removal of palladium during solid-phase synthesis, a new procedure was highly desirable. Various approaches to the removal of heavy metals using $N$-acyclicstene, polystyrene-based thiouronium salts, or silicate particles bearing pendant alkyl groups with terminal thiol functionalities have been reported. While they bind palladium efficiently, they do not work well for polymer products since a large proportion of the polymer product is absorbed by the silica or polymer media. A solubilizing agent would thus be more appropriate. Azothiophoramide have been reported to dissolve metals like zinc and cadmium in the form of their amalgam and Ni(II) compounds through Ni(CO) and Ni(PG)-

Scheme 1

![Scheme 1](image1)

Scheme 2

![Scheme 2](image2)

1 A patent application covering this invention has been filed.
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Figure 1. UV–vis spectrum of 1 and 3 in chloroform at room temperature showing their extinction coefficients. Compound 3 shows a moderately strong absorption with a maximum at 801 nm where compound 1 does not absorb, making this technique applicable for the analytical determination of residual palladium content in materials where palladium catalysis has been employed.

Azothioformamide skeleton can be envisaged that can encompass different solubility requirements of the derived palladium complexes, making its removal easy in terms of solubility/insolubility. Having demonstrated the successful reaction of 1 with metallic palladium nanoparticles/catalyst it remained to evaluate possible side reactions of 1 with the conjugated polymer product. The most likely reaction between an unsaturation such as vinylene or ethynylenylene groups is an electrocyclic condensation reaction of the Diels–Alder type. We performed model reactions between 1 and stilbene or tolane in both the presence and absence of palladium. We were not able to demonstrate detectable amounts of the corresponding electrocyclic condensation products (see Supporting Information). The major problem with such side reactions during polymer synthesis is that they are accumulated, and their alleviation is difficult to envisage. The removal of palladium using 1 is thus possible. While the treatment can be performed at elevated temperature, it works best at room temperature and was normally complete within 1 h. The progress of the palladium dissolution could be conveniently followed using simple UV–vis spectroscopy. This has the advantage that the complex exhibits a moderately strong absorption in a wavelength region normally not explored for saturated polymers and rarely for conjugated polymers and organic materials (ε\text{Pd(THP)} \approx 8500 \text{ M}^{-1} \text{ cm}^{-1}, \varepsilon\text{Pd(Thiolamine)} = 10 \times 300 \text{ M}^{-1} \text{ cm}^{-1}).

The method presented here can thus be used for preparative work where alleviation of residual palladium catalyst is of interest and for analytical purposes where the palladium content can be determined by simple treatment with 1 followed by UV–vis analysis. In the case where there is an overlap, a correct analytical determination of the palladium content is still possible provided that the UV–vis spectrum of the sample is known and can be accurately subtracted.

We have shown the applicability of 1 for the removal of residual palladium catalyst through 3 that in our case is a complex readily soluble in common organic solvents including methanol commonly used to precipitate polymer materials. Using the method of analysis presented here, it is possible to determine the palladium content with a detection limit of 0.1 ppm (w/w) using 50 mg sample. Employing larger samples and concentration of the supernatant the detection limit can be lowered to 1 ppm (w/w). The results are summarized in Table 1.

Table 1. Palladium Content before and after Treatment with 1 for Polymer Samples of the Poly(phenylenevinylene), Poly(phenylethyleneylene), and Poly(alkylthiophene) Type and the Device Resistance According to the Device Geometry (See Supporting Information)

<table>
<thead>
<tr>
<th>polymer</th>
<th>Pd content before/after (ppm)</th>
<th>resistance before/after (kΩ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPV</td>
<td>17860/0.1</td>
<td>0.10/39</td>
</tr>
<tr>
<td>PPE</td>
<td>2565/0.1</td>
<td>0.02/45</td>
</tr>
<tr>
<td>PAT</td>
<td>4073/0.1</td>
<td>0.00/24</td>
</tr>
</tbody>
</table>

The polymers employed for the demonstration were prepared by typical palladium routes (see Supporting Information) and cover a broad range of conjugated polymers: poly(phenylenevinylene) (PPV), poly(phenylethyleneylene) (PPE), and poly(alkylthiophene) (PAT). While the palladium content depends on the amount of catalyst used and on the conditions of the experiment, the levels observed before treatment are typical. Removal using 1 is efficient, and this is reflected in the device resistance and device function. Electroluminescent and photovoltaic devices could be prepared after palladium removal using this method.

In conclusion, our report describes the efficient removal of the palladium nanoparticle/catalyst contained in polymer materials by a palladium route. We further demonstrate the possibility of a quantitative determination of the palladium nanoparticles/catalyst content at the 1 ppm (w/w) level using simple UV–vis spectrophotometry. It should be emphasized that the method is general and that it can be applied to any sample, small molecule or macromolecular. The only requirement is solubility of the sample.

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Supporting Information Available: Details of the one-pot synthesis to 1, preparation of 3, general procedure for the removal of palladium remnants from conjugated polymer materials, analysis of the palladium content in a conjugated polymer sample at the 0.1 ppm level, the device geometry, and UV–vis in THF. This material is available free of charge via the Internet at http://pubs.acs.org.

References and Notes


MA0478S55T
Supporting Information

Removal of palladium nanoparticles from polymer materials
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General information on the properties

Compound 3 was found to exhibit both thermochromic and solvatochromic properties. The UV-vis data reported here thus refer to compound 3 dissolved in THF or chloroform at 25 °C. Compound 1 does not fully dissolve metallic palladium on realistic timescales. It does however dissolve palladium nanoparticles formed during catalysis completely in a short time interval (typically 60 min) as established by ultracentrifugation and following the absorption at 800 nm. Shaking at room temperature overnight was often employed with success. Heating during reaction of 1 with palladium nanoparticles could be employed but room temperature was found to be better. We tested compound 3 as a catalyst in a standard Suzuki reaction between phenylboronic acid and 4-iodoanisole and found no methoxybiphenyl product. We thus conclude that compound 3 is a poor source of palladium during catalysis and that its catalytic activity in palladium catalysed reactions is poor or absent. To ensure that compound 1 does not have any undesirable effect on the polymers, we have evaluated the reactivity of compound 1 and an unsaturation such as vinylene or ethynylene groups. Because 1 has the same unsaturation as 1,4-butadiene, we found that the most likely side reaction would be an electrocyclic condensation reaction of the Diels-Alder type. To test for this type of side reactions, we have made model reactions between 1 and both stilbene and tolane. We did not detect the possible electrocyclic condensation products between respectively stilbene and tolane as shown in scheme S1.

\[ \begin{align*}
\text{Scheme S1. The possible electrocyclic condensation products between 1 and stilbene or tolane.}
\end{align*} \]

The synthetic conditions for the reactions between compound 1 and stilbene or tolane were as follows:

**Compound 1 and stilbene:** In a 5 mL conical flask compound 1 (2.1 mmol, 0.5g) and stilbene (2.1 mmol, 0.38 g) were mixed and dissolved in xylene (3 mL) and flushed with argon for 5 minutes. The solution was then refluxed for 7 days. Every day a sample was drawn to test for the presence of 4. The
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sample was tested with NMR, MALDI-TOF and GC/MS and even after 7 days it was not possible to detect 4 in the solution. This experiment was repeated several times with both more and less solvent volume and with other types of solvent such as THF, NEt₃ and toluene. It was not possible to detect 4 with the given analysis methods in any of the experiments.

**Compound 1 and tolane:** In a 5 mL conical flask 1 (2.1 mmol, 0.5 g) and tolane (2.1 mmol, 0.37 g) were mixed and dissolved in xylene (3 mL) and flushed with argon for 5 minutes. The solution was then refluxed for 7 days. Every day a sample was drawn to test for 5. The sample was tested with NMR, MALDI-TOF and GC/MS. After 7 days it was not possible to detect 5 in the solution. This experiment was repeated several times with both more and less solvent volumes and with other types of solvent such as THF, NEt₃ and toluene. It was not possible to detect 5 with the given analysis methods in any of the experiments.

**Synthesis of compound 1.**

In this section we present the one-pot synthesis of N,N-diethylphenylazothioformamide (1), the synthesis of the Trans-(di(N,N-diethyl-(2-phenyldiazenyl)thioformamide-κS,κN²))palladium (3) and how to use compound 1 to remove Pd from polymer products and to analyse the amount of Pd in the polymer concerned.

![Scheme S2. The synthetic one pot procedure leading to the N,N-diethylphenylazothioformamide.](image)

**N,N-diethylphenylazothioformamide (1);** IUPAC name: N,N-diethyl-(2-phenyldiazenyl)thioformamide (Scheme S2): Ethanol (750 mL) was placed in a 2-litre three necked round bottom flask. The ethanol was degassed with argon for 30 minutes. Phenylhydrazine (39.36 mL, 0.40 mol) was added and the solution stirred under argon with a mechanical stirring. CS₂ (27.60 mL, 0.45 mol) was added drop wise during 15 minutes. A thick colourless precipitate formed. After the mixture had been stirred for an additional 30 minutes, KOH (27 g) in EtOH (200 mL) (≈15% excess) was added. The precipitate dissolved and the colour changed to orange. The solution was stirred for an additional 30 minutes, where a colourless precipitate formed. CH₃I (27 mL ≈15% excess) was added quickly and nearly all the precipitate redissolved. The solution was stirred for an additional 30 minutes. The colour of the solution changed first to red and then to a cloudy very light yellow colour. The solvent was removed by evaporation and diethylamine (300 mL) was added. The solution was then refluxed for 3 days and followed by drawing small samples and subjecting them to ¹H NMR analysis. The reaction was stopped when ¹H NMR showed complete conversion to 4,4-diethyl-1-phenyl-thiosemicarbazide. The solution was cooled to r.t. and atmospheric oxygen was bubbled rapidly through the solution through a glass frit for 24 hours. The colour of the solution changed quickly to dark red. The solvent was removed by evaporation and the liquid residue (a dark red oil) was dissolved in ethyl acetate (300 mL). The organic phase was washed with brine (3 x 250 mL) and dried with MgSO₄. After drying, the organic phase was filtered and evaporated to dryness to give 87.81 g of dark red oil. Recrystallization twice from
n-heptane:ethyl acetate/20:1 (3.15 L) gave 1 in 60% yield (53.23 g) as orange crystals. M.p. 57.5 °C, DSC.: onset: 54.8 °C, peak: 57.5 °C, $\Delta H = 20175.51$ J/mol.

**Synthesis of 3 from metallic palladium black**

\[ 
\text{N=N-S} \xrightarrow{\text{Pd(0)}} \text{N-S-Pd-N-N} 
\]

**Scheme S3:** The synthetic procedure leading to the Trans-(di(N,N-diethyl-(2-phenyldiazenyl)thioformamide-$\kappa S$,$\kappa N^2$))palladium (3) complex.

Palladium black contains of nanoparticles but also of larger particles that were not so reactive towards 1 and therefore a longer reaction time was needed.

**Trans-(di(N,N-diethyl-(2-phenyldiazenyl)thioformamide-$\kappa S$,$\kappa N^2$))palladium (3) (Scheme S3):** THF (100 mL) was placed in a 250 mL conical flask and degassed with argon. N,N-diethylphenylazothioformamide (1) (1.00 g, 4.21 mmol) and Palladium black (Aldrich) (0.25 g, 2.32 mmol) was added and the solution is refluxed under an argon atmosphere for 7 days. The solution was filtered through a thick (4cm) layer of Celite on a sintered funnel. The filtrate was collected and evaporated to dryness. The residue was purified by dry column vacuum chromatography with n-heptane/1,2-dichloroethane in the ratio 10:1 as the eluent. The eluent was removed by slow evaporation to provide dark green crystals. The yield was 164 mg (14 %) of pure compound 3.

**Synthesis of 3 from palladium nanoparticles prepared in-situ (Scheme S4):**

\[ 
\text{Pd(OAc)}_2 \xrightarrow{A)} \text{H}_{2}N-\text{N}^\oplus \xrightarrow{B)} \text{N=N-S} \xrightarrow{\text{Pd(0)}} \text{N-S-Pd-N-N} 
\]

**Scheme S4:** An alternative procedure to synthesize the Trans-(di(N,N-diethyl-(2-phenyldiazenyl)thioformamide-$\kappa S$,$\kappa N^2$))palladium (3) complex.

Palladium (II) acetate (268 mg, 1.19 mmol) was dissolved in DMSO (50 mL) and flushed with argon. Under an argon atmosphere phenylhydrazine (74 mg, 0.69 mmol) was added. The colour of the solution shifted quickly from brownish orange to black. After stirring for 10 minutes compound 1 (680 mg, 2.87 mmol) was added and the solution was refluxed under an argon atmosphere for $1\frac{1}{2}$ hour. The solution was then poured into 600 mL water, and was let standing in the hood without an argon
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atmosphere for 1½ day. The colour of the mixture had changed from very dark brown to dark greenish blue. The mixture was washed four times with 400 mL n-heptane/ethyl acetate mixture (1:1). The entire organic fractions were collected and dried with magnesium sulphate. After filtration the solvent was removed by evaporation. The residue was then purified by dry column vacuum chromatography with n-heptane/ethyl acetate as the eluent in the ratio 10:1. The eluent was removed by evaporation to provide dark green powder. The yield was 480 mg (73 %) of compound 3. An analytical sample for determination of the molar extinction coefficient was obtained by an additional dry column chromatography.

$^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS, the spectrum is presented in figure S1) $\delta$: 1.218 (t ($^3$J = 6.9 Hz), 12 H), 4.177 (broad s, 8 H), 6.431 (broad s, 2 H), 6.909 (broad s, 4 H), 7.575 (broad s, 4H). $^{13}$C NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 16.309, 44.819, 123.692, 131.381, 175.869. MALDI-TOF MS [M+H$^+$] = 549 (the isotopic pattern was confirmed); Anal. Calcd. for C$_{22}$H$_{30}$N$_6$Pd$_2$: C:48.13, H:5.51, N:15.31, S:11.68. Found: C:48.05, H:5.65, N:14.41, S:11.50.

UV-vis (CHCl$_3$ @ 25°C): $\lambda_{\text{max, chloroform}}$ = 801 nm, $\varepsilon_{\text{chloroform}}$ = 10300 M$^{-1}$ cm$^{-1}$

UV-vis in (THF @ 25°C): $\lambda_{\text{max, THF}}$ = 797 nm, $\varepsilon_{\text{THF}}$ = 8500 M$^{-1}$ cm$^{-1}$ see Fig. S2

Figure S1. The $^1$H-NMR spectrum of compound 3. The spectrum was measured in CDCl$_3$ at 300 K.
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General procedure for the removal of Pd from polymer samples by N,N-diethylphenylazothioformamide (1): The polymer was dissolved in THF (no more than necessary, typically 1 ml/20 mg sample) under an argon atmosphere and compound 1 (10 times excess with respect to the amount of Pd in the catalyst used) was added and the solution was stirred between 30 minutes and 1 hour. The mixture can also be refluxed. The polymer was precipitated with methanol (typically 10 times the amount of THF), filtered and washed with methanol. The polymer was redissolved in THF under an argon atmosphere and precipitated with methanol a second time, filtered and washed with methanol. The polymer was then dried in a vacuum oven.

General procedure to analyze the contents of Pd in a polymer sample by N,N-diethylphenylazothioformamide (1): The polymer sample was very precisely weighted out and dissolved in THF (typically 20 mg/ml) under an argon atmosphere. Compound 1 (10 times excess with respect to the amount of Pd in the catalyst) was added and the solution is stirred between 30 minutes and 1 hour. The solvent was evaporated and the solid was redissolved in a precisely known volume of solvent and a UV-vis spectrum of the solution was recorded.

Examples of Pd-catalysed polymerisation to conjugated polymers, Pd analysis and device preparation.

Three types of polymer, PPE\(^1\), PPV\(^4\) and PAT\(^5\) were used in the investigations.

Example 1: Poly[1-(2,5-dioctyl-tolanyl)-ethynylene] (PPE)\(^1\). A 100 mL round-bottom flask was dried using a heat gun and flushed with argon. 1-iodo-2,5-dioctyl-4'-ethynyltolane (1.36mmol, 0.750 g) was dissolved in piperidine (40 mL) and flushed with argon for 30 minutes, before the catalyst Pd(PPh\(_3\))\(_4\) (≈ 2 mol %) and the co-catalyst CuI (≈ 10 mol %) were added under an argon atmosphere. The solution was stirred under an argon atmosphere over night at room temperature. A yellow
precipitate quickly formed. The colour of the solution was yellow during the reaction. The following
day the reaction mixture had solidified. Dry THF (50 mL) was added and the solution was heated to
near reflux temperature. The precipitate dissolved and the solution was stirred for another 5 hours. The
solution was then poured into 300 mL methanol and a yellow precipitate formed. The suspension was
filtered through a glass filter-funnel. The filter cake was washed with both cold methanol, cold diethyl
ether and was sucked as dry as possible. The wet product was kept for further use. Attempts to dry the
product in a vacuum at RT gave a darkened product that was no longer soluble in common solvents.
This behaviour has been observed\(^2\) earlier. The total yield based on a completely dried sample was
0.35 g (95%). The wet product that was kept weighed 0.6 g and contained 40% solvent.\(^1\)H NMR \(\delta:\)
0.76-0.95 (m, 6H), 1.13-1.50 (m, 20 H), 1.50-1.83 (m, 4H), 2.70-2.90 (m, 4H), 7.31-7.43 (broad s),
7.43-7.60 (broad s) (7.31-7.60, 6H).

**Pd-analysis of the PPE polymer**  The PPE polymer (0.1028 g) was dissolved in 3 mL THF under an
argon atmosphere. Compound 1 (0.035 g) was added and the solution was stirred under an argon
atmosphere in 1 hour. The solution was diluted to a volume of 5 mL and the UV-vis spectrum was
recorded. This gave an absorbance of 0.441 which corresponds to a concentration in \(3\) of 51.9 \(\mu\)M.
The amount of Pd in the polymer was thus 0.259 \(\mu\)mol ~ 27.60 \(\mu\)g Pd in 0.1028 g of polymer. This is
equivalent to 268 mg Pd kg\(^{-1}\) or 268 ppm. The result of the Pd determination in the polymer is listed in
table 1 in the article.

**Example 2: Poly-1,2\,'''-(2,5-dioctyl-1,4-phenylene-1',2'-vinylene-1''',4'''-phenylene-1''',2'''',
vinylene) PPV.\(^4\)** 1,4-Dibromo-2,5-dioctylbenzene (1 g, 2.17 mmol), 1,4-divinylbenzene (0.28 g, 2.17
mmol), triethylamine (10 mL) were mixed in dry THF (50 mL) and the mixture was degassed with
argon for 5 min. (PPh\(_3\))\(_2\)PdCl\(_2\) (250 mg) was added and the mixture was heated to reflux. After 36 h
the mixture was poured into methanol to precipitate the bright yellow product. It was then filtered and
dried to give a yellow solid in 85% yield (0.79g). Analytical data as in ref. 4. a mixture of palladium
nanoparticles and polymer could be obtained by dissolution of the product in THF followed by
ultracentrifugation.

**Pd-analysis of the PPV polymer**  The PPV polymer (0.0577 g) was dissolved in 5 mL THF under an
argon atmosphere. Compound 1 (0.035 g) was added and the solution was stirred under an argon
atmosphere in 1 hour. The solution is diluted to 50 mL. From this sample the UV-vis spectrum is
recorded. This gave an absorbance of 0.247 corresponding to a concentration in \(3\) of 29 \(\mu\)M. The dilution factor of the 50 mL solution was 6.67 corresponding to a total volume of 333.3 mL. The amount of Pd in the polymer was thus 9.7 \(\mu\)mol ~ 1030 \(\mu\)g Pd in 0.0577 g of polymer. This is equivalent to 17860 mg Pd kg\(^{-1}\) or 17860 ppm. The result
of the Pd determination in the polymer is listed in table 1 in the article.

**Example 3: Poly-(3-hexylthiophene) PAT.\(^5\)**  DMSO (30mL) and DMF (30mL) was degassed using
argon and (PPh\(_3\))\(_2\)PdCl\(_2\) (90mg, 0.128 mmol, catalyst) was added and the mixture was heated to 75 \({^\circ}\C with stirring. 2-Bromo-3-hexyl-5-trimethylstannylthiophene (1.75g, 4.26 mmol) was added drop wise
from a syringe. The colour of the mixture quickly changed to clear yellow and after 1 min. it became
cloudy and orange and after another minute it became red/brown and a material started to separate.
After 3 days under argon the mixture was a light yellow clear solution with a black precipitate. The
mixture was filtered and the solid washed with methanol (5 x 50mL). Drying gave a black solid 0.69g
(97%).

**Pd-analysis of the PAT polymer**  The PAT polymer (0.035 g) was dissolved in 5 mL THF under an
argon atmosphere. Compound 1 (0.030 g) was added and the solution was stirred under an argon
atmosphere in 1 hour. The solution is diluted to 50 mL. From this sample the UV-vis spectrum is
recorded. This gave an absorbance of 0.228, which gives a concentration in \(3\) of 26.8 \(\mu\)M. The amount of Pd in the polymer was thus 1.34 \(\mu\)mol ~ 143 \(\mu\)g Pd in 0.035 g of polymer. This is equivalent to
4073 mg Pd kg$^{-1}$ or 4073 ppm. The result of the Pd determination in the polymer is listed in table 1 in the article.

**Standard electroactive device for fabrication:** The substrates were 10 Ω square$^{-1}$ ITO slides that had been etched in one end by immersion in a 65 °C aqueous solution of HCl(aq) 20% and HNO$_3$(aq) 5% for typically 60 seconds. The device geometry is shown in figure S3.

![Figure S3. A schematic illustration of the electroactive device geometry (not drawn to scale).](image)

The etched substrates were cleaned by immersion in isopropanol and subjected to ultrasound for 10 min. After being blown dry in a stream of dry argon a layer of PEDOT:PSS was applied by spincoating an aqueous solution of PEDOT:PSS (1.3% wt.) containing sorbitol (2% wt.). The substrates were placed in an oven at 100 °C and ramped up to a temperature of 180 °C during 1h and kept at 180 °C for 1h. After cooling to room temperature the polymer solutions (in chlorobenzene at a concentration of 2-10 mg mL$^{-1}$) were micro-filtered and spin coated at 1500 rpm onto the substrates. The typical film thicknesses were 50-100 nm as measured by a DEKTAK 3030 and the typical absorbance of the films were 0.25-1.0 absorbance units. The geometry of the devices is shown in figure 1. The aluminium electrode was evaporated onto the device at a pressure of $< 5 \times 10^{-6}$ mBar. After application of the electrodes the samples were removed from the evaporator chamber, electrical connections were made using conducting silver epoxy glue. The active area of the device was 3 cm$^2$ (20 mm x 15 mm). The resistance was measured using a Keithley 2400 Sourcemeter. Typical experimental variation between devices was $< 10\%$. 

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References


Effective Removal and Quantitative Analysis of Pd, Cu, Ni, and Pt Catalysts from Small-Molecule Products

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Abstract: We demonstrate the ability of N,N-diethylphenylazothioformamide and N,N-bis[6-(ethylyene glycol)monomethyl ether]phenylazothioformamide ligands to dissolve the four transition metals: palladium, copper, nickel, and platinum. The method allows for the quantitative analysis of the transition metals content in their bulk metallic forms and as nanoparticles produced during their use as catalysts in Sonogashira coupling reactions.

Key words: palladium, copper, nickel, cross-coupling reaction, catalysis

For more than 100 years organic chemists have been attracted to transition metals, due to the vast number of possibilities that are offered in the field of organic synthesis, e.g., the selective activation of C–H, C–C and C–X bonds (X = good leaving groups such as halides). There has, however, been only sporadic interest in ensuring that the amount of transition metals remaining in the newly synthesized organic compounds are at the ppb level.1–12 Organic chemists rarely consider impurities at a level below 0.1%, because impurities at that level normally do not interfere with characterization techniques such as NMR, GC-MS, HPLC, IR, UV-Vis and elemental analysis. The impurity level in organic compounds should, however, be much smaller than 0.1%, if the compounds are to be used in pharmaceutical or electronic applications.1–3 In particular, removal of trace amounts of palladium from drugs is a significant problem for the pharmaceutical industry; an aspect that is often neglected in the literature.2,5 In this respect it should be mentioned that concentration limits of, for example, Pd and Cu in orally administered drugs should be below 5 ppm and 15 ppm respectively, while the parental concentration limits should be 1/10 of the limit for the oral doses.2

We have recently demonstrated that palladium impurities of between 268 ppm and 17860 ppm in derivatives of polyphenylene vinylene (PPV), poly(alklyliylene) (PAT) and poly(phenylene ethynylene) (PPE) have a dramatic effect on the electrical properties of thin film devices made from these polymers.15–18 We observed that, as the resistance of the devices (OLEDs) approached short circuit, no electroluminescence could be observed. In this case we found that the palladium remnants could not be removed by normal purification methods such as size-exclusion chromatography (SEC), but that Pd nanoparticles could be dissolved and thereby removed from the polymers by treatment with N,N-diethylphenylazothioformamide (1) (Figure 1a). The contamination level of palladium in these polymers, after such treatment, was lowered to below 0.1 ppm and the resultant thin polymer film devices had high resistance and were electroluminescent.

Even though palladium is by far the most commonly used transition-metal catalyst19 (used in e.g., Heck,20 Sonogashira,21 Stille,22 Suzuki23 and Negishi24 cross-coupling reactions), it would be a great advantage if compound 1 also worked for other transition metals. In this paper we focus on four transition metals: copper, nickel, palladium and platinum – and demonstrate that 1 also has the ability to dissolve these metals while forming the coordination compounds shown in Figure 1b. Furthermore, we present the synthesis of N,N-bis[6-(ethylene glycol)monomethyl ether]phenylazothioformamide (2) (Figure 1a) designed to extend the use of these ligands to small-molecule samples and demonstrate that compound 2 can also dissolve the transition metals by forming the coordination compounds shown in Figure 1a.

The synthetic route to 2 is shown in Scheme 1 and is described in the experimental section. The most interesting difference between 2 and 1 is that the N,N-bis[6-(ethylyene glycol)monomethyl ether]amine group in 2 exhibits a strong affinity for silica in polar solvents such as CHCl3. When the pure ligand 2 or the complexes 7–10 are filtered through silica gel, with CHCl3 as eluent, the materials are immobilised on the columns. This property makes it possible to remove transition-metal catalysts from samples of small organic molecules in a very easy manner; simple dissolution of the compound in CHCl3, together with 2, followed by stirring overnight and subsequent filtration through a pad of silica gel. The organic material can then be recovered from the filtrate. While 2 is water-soluble and, in principle, can be used for removal of metals from aqueous solutions, the distribution coefficient of 2 between water and organic solvents such as CHCl3 or 1,2-dichlorobenzene is rather low. This means that, in practice, it is impossible to carry out an extraction of 2, or of its complexes, from an organic solvent into water. Compound 2 can, however, be used to remove catalyst reman...
The model reaction chosen was of the Sonogashira cross-coupling type, since this reaction can be performed either with Pd or Cu alone as the catalyst, or with Pd as the catalyst and Cu as a co-catalyst. It has also been reported that Ni works as a catalyst for this type of reaction, but that reaction did not work satisfactorily in our hands. The two transition metals Pd and Cu do not behave

Scheme 1 The synthetic route used for the preparation of compound 2.

identically in the Sonogashira reactions, as a consequence we have used different solvent-base pairs for the two model systems (Scheme 2). More details of the model system are to be found in the literature (Pd\textsuperscript{17} and Cu\textsuperscript{21}) and in the experimental section.

As described earlier,\textsuperscript{14} the use of 1 to remove palladium, also makes possible the analysis of the palladium content in the samples by UV-Vis spectroscopy, with a detection limit as low as 10 ppb. We demonstrate here that both 1 and 2 have the ability to remove a range of transition metals from an organic sample and furthermore 1 can be used to quantify that content. Note that 1 is designed to remove transition metals from large molecules through precipitation from excess methanol, a solvent in which both 1 and its metal complexes are soluble. Compound 2, on the other hand, is designed to remove transition metals from small organic molecules, which can not be precipitated but where filtration through a short silica column works well.

From the first type of experiments we observed that 1 very effectively dissolves a copper metal wire. Figure 2 shows the results after the copper wire had been immersed in a concentrated THF solution of 1 for just one week. Other metals are not dissolved as quickly. Visual inspection of nickel wire under the same conditions showed that, while some of the metal was dissolved, the extent of erosion was significantly less pronounced than that of the copper wire. We could only detect the dissolution of the platinum wire by a UV-Vis analysis of the solution. No solubilisation of the palladium wire was detected; however, we have demonstrated that 1 dissolves both palladium nanoparticles and palladium black, so the inability of this reagent to dissolve this metal seems to be due to a surface phenomenon.

From this set of experiments we can conclude that the reactivity of 1 towards the four transition metals increases in the order Pd < Pt < Ni < Cu. Visual inspection of the experiments with compound 2 showed that this compound does not seem to be as effective as 1 in dissolving the metals. This might be due to the steric hindrance of the long tetra(ethylene glycol) monomethyl ether groups, but the trend appears to be the same.

To be able to make a quantitative analysis of the transition-metal content in newly synthesised organic compounds as described earlier for palladium,\textsuperscript{14} we have determined the molar extinction coefficients at \(\lambda_{\text{max}}\) for the four complexes with the ligand 1 in both CHCl\textsubscript{3} and THF. The values and the respective \(\lambda_{\text{max}}\) are listed in Table 1.

The UV-Vis spectra for the four complexes are shown in Figure 3. It can be seen that the \(\lambda_{\text{max}}\) increased in the order Pt < Cu < Pd < Ni and the extinction coefficients in the series Ni < Cu < Pd < Pt.

We have also synthesised complexes with 2 as the ligand, however, it turned out that it was very difficult to separate the excess of 2 from the complex of 2 and the transition metal. By examining the UV-vis spectrum of the Cu complex (9) in water and CHCl\textsubscript{3} (Figure 4), it can be seen that the spectrum in water is characterized by a broad peak with maximum at 657 nm. This peak is not present in the spectrum of the complex in CHCl\textsubscript{3}. The explanation to this peak is probably that the complex is easily oxidized to the CuL\textsubscript{2}\textsuperscript{2+} complex in aqueous solution. This explanation is supported by the literature\textsuperscript{23} where the spectrum of the CuL\textsubscript{2}\textsuperscript{2+} (L = 1) is also characterized by this broad peak.

The content of Pd and Cu in the final product of the model reaction was analysed with 1 and, as can be seen in

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**Figure 2** Copper wire after one week in a concentrated THF solution of 1. Notice how thin the copper wire is in the middle compared to the ends, and how the crystals of the cis-[d1]Ni(N,N-diethyl(2-phenylisobenzofuran-2-y)imino)benzofuran-2-y]copper (5) are growing on the wire.

**Table 1** Molar Extinction Coefficients and the \(\lambda_{\text{max}}\) Values for the Four Complexes

<table>
<thead>
<tr>
<th>Compd</th>
<th>M</th>
<th>THF (\lambda_{\text{max}}) (nm)</th>
<th>e (M\textsuperscript{-1} cm\textsuperscript{2})</th>
<th>CHCl\textsubscript{3} (\lambda_{\text{max}}) (nm)</th>
<th>e (M\textsuperscript{-1} cm\textsuperscript{2})</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Pd</td>
<td>790</td>
<td>8500</td>
<td>811</td>
<td>10300</td>
</tr>
<tr>
<td>4</td>
<td>Pt</td>
<td>727</td>
<td>26400</td>
<td>731</td>
<td>25400</td>
</tr>
<tr>
<td>5</td>
<td>Cu</td>
<td>770</td>
<td>39000</td>
<td>768</td>
<td>3400</td>
</tr>
<tr>
<td>6</td>
<td>Ni</td>
<td>830</td>
<td>1800</td>
<td>830</td>
<td>1800</td>
</tr>
</tbody>
</table>

In order to validate the Pd and Cu content after treatment with 2, we also analysed the samples with inductively coupled plasma sector field mass spectrometry (ICP-SFMS). As the results show (Table 2), both analytical methods gave comparable results for transition-metal content both before and after treatment with 2. Further, it should be emphasised that both analytical methods determined the content of the transition metals after treatment with 2 to be below the parenteral concentration limit for drugs.

Table 2  Transition-Metal Content in the Products of the Model Reaction before and after Treatment with 2 as Determined by UV-Vis Spectroscopy with 1 and ICP-SFMS

<table>
<thead>
<tr>
<th>Metal</th>
<th>UV-Vis (ppm) / ICP-SFMS (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pd</td>
<td>Before treatment: 376/610</td>
</tr>
<tr>
<td>Cu</td>
<td>Before treatment: 1074/1660</td>
</tr>
</tbody>
</table>

In conclusion, we have demonstrated the ability of 1 and 2 to dissolve four transition metals Pd, Pt, Cu and Ni, in a contaminated organic product prepared via metal catalysis. We demonstrated the use of 1 for the analysis and quantitative determination of the content of the four metals in a given organic sample by UV-Vis spectrophotometry. Furthermore we have demonstrated that, unlike 1, the high affinity of 2 for silica enables its use in the facile removal of any of these transition metals from small organic compounds simply by filtration through a short silica pad. It should be emphasised that this method is fast, easy, potentially low-cost (no expensive chemicals) and can be applied to any organic sample, as long the sample is soluble.

$^1$H and $^{13}$C NMR spectra were recorded on a 250 MHz NMR spectrometer at 300 K unless otherwise stated. All spectra were recorded in CDCl$_3$, with TMS as an internal reference. Compound 1 was prepared according to the procedures described in the literature.
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Scheme 3 Modified synthesis of N,N-bis[2(ethylene glycol) monomethyl ether]amine, using NaH instead of KOH. 

N,N-Bis[2(ethylene glycol) monomethyl ether]amine

The modified literature procedure used in the synthesis of N,N-bis[2(ethylene glycol) monomethyl ether]amine is illustrated in Scheme 3. We found that changes to the procedure were necessary since, in our hands, the use of 12 equivalents of KOH in step 4 led to a yield of only 25%, whereas a yield of 81% was obtained when NaH was used as a base. All other details were the same as described in the literature.

N,N-Bis[2(ethylene glycol) monomethyl ether]phenylazothioformanide

IUPAC name: N,N-bis[2(ethylene glycol) monomethyl ether]-2-phenylazothioformanide.

A 500-mL three-neck, round-bottom flask, equipped with a mechanical stirrer, under argon was charged with EtOH (75 mL) and degassed with argon for 30 min. Phenyl hydrazine (5.41 g, 49.2 mL, 50.0 mmol) was added and stirred under argon before Cs$_2$(3.81 g, 3.02 mL, 50.0 mmol) was added dropwise, forming a thick colourless precipitate. The mixture was stirred for 30 min then KOH (3.00 g, 59.9 mmol (<10% excess) in EtOH (20 mL)) was added. The resulting orange solution was stirred for an additional 30 min, during which a colourless precipitate formed. CH$_2$Cl$_2$ (7.45 mL, 32.5 mL, 55% (<5% excess)) was added quickly and nearly all of the precipitate dissolved. The solution was stirred for an additional 30 min, during which the colour of the solution changed first to red and then to a very light, cloudy yellow. The solution was filtered under an argon atmosphere, and the solvent was removed by evaporation to give a light-yellow solid (10.0 g mass balance). 4.66 g of this compound was transferred to a 250-mL round-bottom flask, equipped with a mechanical stirrer, under argon was charged with EtOH (75 mL) and degassed with argon for 30 min before use, and refluxed under argon for 5 h. The reaction was stopped when $^1$H NMR showed complete conversion to 4,4-bis[2(ethylene glycol) monomethyl ether]-1-phenylthiosemicarbazide.

The solution was cooled to r.t. and atmospheric O$_2$ was bubbled rapidly through the solution through a glass stirrer for 24 h. The colour of the solution changed quickly to dark red. The solvent was removed by evaporation to give a dark-red oil that was purified by dry column vacuum chromatography with a gradient of n-heptane-EtOAc (containing 1 to 5% of Et$_3$N) as the eluent. Yield: 2.25 g (18%); dark-orange oil; $R_f$ = 0.13 (yellow band) (2 drops Et$_3$N per 10 mL EtOAc).

$^1$H NMR (200 MHz, CDC$_3$) δ = 3.37 (s, 6 H), 3.49-3.66 (m, 26 H), 3.87 (t, $J$ = 5.7 Hz, 2 H), 3.95 (t, $J$ = 5.3 Hz, 2 H), 4.28 (t, $J$ = 5.3 Hz, 2 H), 7.47-7.58 (m, 3 H), 7.64-7.90 (m, 2 H).

$^{13}$C NMR (250 MHz, CDC$_3$) δ = 51.9, 54.3, 59.0, 60.8, 69.3, 70.4, 70.5, 70.6, 70.7, 70.7, 70.7, 72.0, 123.7, 123.9, 123.9, 151.9, 195.1.

Anal. Calc'd for C$_{29}$H$_{45}$N$_3$O$_3$: C, 55.03; H, 7.94; N, 7.70. Found: C, 55.34; H, 8.06; N, 7.61.

trans-Di[2(2,6-bisazine)-2(ethylene glycol) monomethyl ether]-2-phenylazothioformanide-(5,5,6,7,7,8,8,8-d$_8$)-copper(0) (9)

A 2-cm-long copper rod was placed in a solution of 2 (400 mg, 0.733 mmol) in THF (1 mL). After a week the rod was removed, the solution was evaporated and the residue was dissolved in CHCl$_3$. Purification by size-exclusion chromatography gave a brownish-red oil (152 mg).

MALDI-TOF: m/z [M + H$^+$] calcd: 1153.49; found: 1153.35 (the isotopic pattern was confirmed).


It was not possible to obtain NMR data because Cu is paramagnetic.

Pd-catalysed Synthesis of Diphenyl Acetylene

A 250-mL Erlemeier flask equipped with a magnetic stirring bar was charged with iodobenzene (12.2 g, 6.70 mL, 60.0 mmol), pyridine (50 mL) and THF (20 mL). The mixture was flushed with argon for 30 min before Pd(PPh$_3$)$_2$Cl$_2$ (2 mol%, 1.145 g, 0.99 mmol) was added under argon atmosphere. Phenylation (6.64 g, 7.14 mL, 65.0 mmol) was then added and the mixture was stirred at r.t. under an argon atmosphere. After 2 d, the solvent was removed by evaporation and the solid residue was dissolved in CHCl$_3$ and filtered through a short aluminium oxide column. The solvent was removed by evaporation and the crude product was recrystallized from EtOH (100 mL) to give diphenyl acetylene (7.25 g, 69%).

Cu-catalysed Synthesis of Diphenyl Acetylene

A 250-mL round-bottom flask, under argon, equipped with a magnetic stirring bar and a condenser was charged with dry DMF (110 mL) (flushed with argon for 30 min before use), iodobenzene (12.2 g, 6.70 mL, 60.0 mmol), K$_2$CO$_3$ (=140 mol%, 11.6 g, 84.1 mol), PPh$_3$ (=10 mol%, 1.60 g, 6.10 mmol), Cu(II) 5 mol%, 0.560 g, 29.9 mmol) and phenylacetylene (6.64 g, 7.14 mL, 65.0 mmol) and the mixture was stirred at 110°C for 3 days. The solvent was removed by evaporation and the solid residue was dissolved in CHCl$_3$, and filtered through a short aluminium oxide column. The solvent was removed by evaporation and the crude product was recrystallized from EtOAc to give diphenyl acetylene (7.49 g, 70%).

Quantitative Analysis of Transition Metals in an Organic Sample Using 1: General Procedure

The diphenyl acetylene was very precisely weighed out and dissolved in CHCl$_3$, (typically 500 mg/mL) under an argon atmosphere. Compound (1 tenfold excess with respect to the amount of Pd) was added and the solution was stirred overnight. The solution was diluted to a precisely known volume with CHCl$_3$, and a UV-Vis spectrum was recorded.

Removal of Transition Metals from an Organic Sample by 2: General Procedure

The diphenyl acetylene was dissolved in a minimum volume of CHCl$_3$, (typically 2 mL/500 mg sample) under an argon atmosphere and compound 2 (tenfold excess with respect to the amount of Pd) was added and the solution was stirred overnight. The solution was
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diluted with CHCl$_3$ (10 mL) and filtered through a short pad of silica gel. The silica gel was washed with CHCl$_3$ (100 mL) and the solvent was then removed by evaporation. The purified diphenyl acetylene was then ready to be tested for transition metals using the method described above.

Acknowledgment

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References

Design of porphyrin-polyphenyleneethynylene light harvesting systems

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ABSTRACT

The design of light harvesting systems based on zinc-porphyrin linked polyphenyleneethynylene systems is demonstrated through two different synthetic procedures. The use of the Sonogashira cross-coupling reaction was found to be problematic in several aspects and it was only possible for one of the synthetic strategies to incorporate a single porphyrin molecule in each polymer chain. The polymer materials were separated into fractions using preparative SEC and subjected to photophysical studies in both the solution and in the solid state. In the solid state light harvesting by the pendent polyphenyleneethynylene and energy transfer to the zinc-porphyrin was found to be high whereas it was low in solution. This observation was confirmed when making electroactive devices. I/V characteristics were measured on fabricated solar cells of the polymers. Material properties for the native polyphenyleneethynylene were determined using ultra violet photoelectron spectroscopy on thin films and carrier mobility studies were performed using pulse radiolysis time resolved microwave conductivity.

Keywords: Dye-linked conjugated homopolymers, zinc-porphyrin linked polyphenyleneethylenes, near-infrared electroluminescence, energy transfer through bonds, organic based photovoltaics, organic light emitting diodes.

1. INTRODUCTION

The finite supply of fossil fuels is a great motivation to develop new and readily less polluting energy resources. One alternative energy resource to the fossil fuel is the Sun. Unfortunately the current cost of energy (~54/W) produced by conventional inorganic photovoltaics (PV) or solar cells fabricated form crystalline silicon is still too high to compete with the fossil fuel or wind energy.\textsuperscript{1} Lots of work have been afforded to find more inexpensive materials there can replace silicon in the PVs. Since the beginning of the 90s a huge interest for application of organic based photovoltaics (OPV)\textsuperscript{2} where the silicon is replaced by e.g. polymer has been build up. Like OPVs, polymer based organic light emitting diodes (OLEDs) have also attracted large scientific and industrial attention due to their very promising applications e.g. flexible large-area flat-panel displays.\textsuperscript{3} Some of the biggest advantages of polymer based OPVS and OLEDs in comparison to traditional inorganic semi-conducting materials are first and foremost the low cost of the materials and the much easier fabrication methods. The traditional inorganic semi conducting components are normally prepared by a series of complicated vacuum techniques, while polymer based OPVs and OLEDs can be prepared by techniques like spin coating, screen or ink jet printing.\textsuperscript{4} Furthermore, the possibilities to tune the colour of the light emission of the polymer based OLEDs by chemical modifications are nearly unlimited and relatively easy.\textsuperscript{4,5} The OLEDs can be used for full colour displays because they are able to emit all three principal colours: Blue, green and red. Among these colours green is the most developed concerning brightness and efficiency. The reports on polymer based OLEDs which emits in the near infrared (NIR) region are still relatively few, and include mostly low band gap polymers like poly(thiophenes),\textsuperscript{6} polymers doped by rare earth metal ions,\textsuperscript{7} and polymer blends where the hosts are a high band gap conjugated polymer doped with small red-emitting organic dyes such as porphyrins.\textsuperscript{8} The latter case is the most frequently used and the idea in this case are that the conjugated polymer host both work as charge transporter and take care of the transfer of the excitons formed within itself to the dye molecule. In the dye molecule a decay of the excitations happens while emitting NIR light. The transfer of the excitons to the dye molecule typically occurs through space (TS) normally by a Förster energy transfer process,\textsuperscript{9} which is a long-range dipole-dipole coupling process.
between the transition dipoles of the excited polymer and ground-state dye molecule. The rate constant of the Förster energy transfer process, $k_F(r)$, is proportional to $r^{-6}$, where $r$ is the distance between the donor and the acceptor. Hence the Förster energy transfer process occurs normally only over a range of 1 to 50 Å. The problems with the efficiency of the NIR light-emitting polymer blends are that the energy transfer efficiency among other things depend on how close the donor and acceptor are to each other, which means that it is very important that the polymer and the dye molecules are mixed very thoroughly. Furthermore, it requires a good spectral overlap between the emission and the absorption spectra for the polymer and the dye molecules respectively, which is far from the general observation.

Recently, new NIR light-emitting materials have been produced by linking the dye molecules to the backbones of conjugated polymers by covalent bonds. In this way, it is ensured that the dye molecules are placed close and in a regular fashion with respect to the polymers. The TS energy transfer is therefore optimized, and an eventual aggregate of the dye molecules, which can lead to self-quenching of their fluorescence, is prevented. Finally, the covalent link between the polymer and the dye molecule also opens up for the possibility of energy transfer through bonds (TB) (see Fig. 1). The TB energy transfer is promoted by an orbital overlap, and is much faster than the TS transfer process. The rate of the energy transfer TB can be tuned, if the dye molecule forms a part of the conjugated polymer backbone.

In relation to light-emitting materials where the dye molecule is covalently linked to the polymer backbone we have recently synthesized, a covalently linked three-domain structure of the type outlined in Fig. 1. The three-domain structure, NPN, consisted of two poly[1-(2,5-dioctyl-tolanyl)-ethylenylene] chains (N domains) and a dye molecule in the shape of [10,20-Bis-(3,5-bis-tert-butylphenyl)-5,15-dibromophenanthro[2,1] (P domain) (See Scheme 1). In solid state this structure shows a very interesting property. The N domains serve as light-harvesting antenna for the P domain, which then emits the energy in the NIR region. The same property has been reported for two similar domain structures.

In this paper we will briefly describe the synthesis of NPN and some of its properties. Furthermore we will focus on how the length of the N domains influences the efficiency of the energy transfer process in the NPN. To control the length of the N domain, we applied two different methods. The first method employed monodisperse phenylacetylene terminated oligo(1-(2,5-dioctyltolanyl)ethylene) of precise molecular length (See Scheme 3 for further details). The oligomers were then coupled to the porphyrin dye molecule by the Sonogashira cross coupling reaction. The second method employed the use of Size Exclusion Chromatography (SEC) to make narrow fractions with different molecular weight of NPN, whereby different lengths of the N domains were obtained.

2. EXPERIMENTAL

The compounds polyphenyleneethylenylene (N), NPN and 1-iodo-2,5-dioctyl-4’-trimethylsilylenyltolane (protected monomer unit) were prepared and purified from the contaminated palladium nanoparticles as described previously. Ultraviolet photoelectron spectroscopy (UPS) and measurements of carrier mobilities of the homopolymer by using the pulse radiolysis time resolved microwave conductivity (PR-TRMC) technique were reported previously.
LED and solar cell devices were prepared by spincoating chloroform solutions onto PEDOT:PSS covered ITO slides (the hole-injecting (LEDs) or collecting (solar cells) electrode). The prepared films had peak absorbencies of -0.6 (LEDs) and -0.4 (solar cells). Ca/Al was used as the electron-injecting (LEDs) or collecting (solar cells) electrode, and was evaporated in a vacuum of $< 1 \times 10^{-3}$ mbar. In the case with the solar cells, sets of bilayer devices were also made. In the bilayer devices a layer of Ca$\text{}_6$ was evaporated (vacuum $< 1 \times 10^{-5}$ mbar) on top of the polymer before evaporation of the aluminium electrode. The electrical connections were achieved using silver epoxy. Electroluminescence and fluorescence spectra were recorded in a FLS920 from Edinburgh Instruments. I/V characteristics were measured in dark and under a solar simulator generating AM 1.5 light (~100mW/cm$^2$) with a Keithley 2400 sourcemeter. The apparatus for the SEC was from Knauer and consists of a preparative system with a 3 mL injection loop. The detector was a UV-vis diode array with a resolution of 1 nm. A 16-port switch valve allows for separation of fractions. Two different column systems were used. They consisted of DVB linked polystyrene gel columns comprising a guard column and two gel columns. One system had a 100 Å and a 1000 Å column in succession, and the other had a 500 Å and a 10000 Å column in succession. The eluent was chloroform and the flow rate was 20 mL/min with a pressure of 75 MPa. The molecular weight was calculated on the basis of polystyrene standards having molecular weights of: 515000, 127000, 68000, 36300, 19000, 10900, 3250 and 1360. In all cases the polydispersities of the standards were smaller than 1.007.

### 3. RESULTS/DISCUSION

#### 3.1 The synthesis of NPN

NPN was synthesized by two different synthetic routes, which are illustrated in scheme 1 and 2. It was concluded that only the synthetic route illustrated in scheme 1 lead to the three domain structure shown in Fig. 1, where only one porphyrin unit is covalent linked in a sandwich between two polymer chains.

![Scheme 1](image)

Scheme 1: NPN was synthesized by a Sonogashira cross coupling reaction.\textsuperscript{17}
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Scheme 2: The unsuccessful synthesis of NPN by a Sonogashira cross coupling reaction.\textsuperscript{17}

The problem with the synthetic route illustrated in Scheme 2 is, that the growing chain always expose an ethynyl group, which is capable of undergoing an oxidative coupling to give a butadiyne byproduct. It means that the synthetic route in addition to the desired NPN structure also give a lot of buta-1,3-diyne-1,4-diyl byproducts derived from undesired couplings such as ...NP-PN... and ...NPN-NPN...

![Figure 2: The electronic band structure derived from the UPS spectrum. EFVFB = 1.5 eV; E_VAC = 4.0 eV; B_{max} = 46.0 eV; \Delta = -0.70 eV; IP = 5.50 eV and \phi_{a0} = 4.7 eV. In the right corner of the figure the substrate plate is shown.\textsuperscript{18}]

The native polyphenyleneethynylene (N) was synthesized by a Sonogashira cross coupling reaction to give information of the carrier mobility, the electronic energy levels and the injection barrier for holes into the valence band of the N domains. The sum of charge carrier mobility was determined by PR-TRMC\textsuperscript{24,25} and the results were \( \Sigma_{\text{sum}} = 1.25 \times 10^{-6} \) m\(^2\) V\(^{-1}\) s\(^{-1}\) and a first half life of \( \tau_0 \approx 10 \) μs. The value for \( \Sigma_{\text{sum}} \) has not been corrected for the survival probability charge carrier pair or for the differential dose distribution between the conjugated backbone and the alkyl sidechains.\textsuperscript{18} The carrier mobility was not as high as hoped, but at least it was comparable to the values determined earlier for similar compounds.\textsuperscript{26} The electronic energy levels and the injection barrier were determined by UPS,\textsuperscript{27} which employs hard UV radiation from a synchrotron source. The results of the UPS measurements are listed in Fig. 2.
3.2 Synthesis of oligomers

To make monodisperse phenylacetylene terminated oligo(1-(2,5-dioctyloxylanyl)ethyne) we have employed the synthetic route illustrated in scheme 3. The first step was to couple phenylacetylene with the protected iodo-monomer unit by a Sonogashira cross coupling reaction, which progressed nicely. The second step was removal of the protecting group TMS by a base (e.g. K₂CO₃) followed by a new coupling reaction with the protected iodo-monomer unit to yield the protected dimer. These steps were repeated until the required molecular length was obtained.

Scheme 3: The stepwise route to synthesize monodisperse phenylacetylene terminated oligo(1-(2,5-dioctyloxylanyl)ethyne) of precise molecular length. 1) Pd(PPh₃)₄ (2 mol %), Cul (10 mol %) and THF at r.t. 2) K₂CO₃ (10 equiv.) and CH₂Cl₂/MeOH (1:1) at r.t.

The idea was that this cycle should be carried out without any other purification than filtration and a short aluminium oxide column providing e.g. the trimer in 95 % purity with only minor impurities in the form of the dimer, monomer or 1,4-diarylbutadiynes from a oxidative dimerization of the unprotected ethynyl groups. Preparative SEC could then be used to purify the trimer. To be sure that each step in the synthetic cycle was completed before changing to another step, we kept each step running for 2 days. Furthermore we added more catalyst to the coupling reaction after one day, to be sure that there was still active catalyst present in the reaction mixture.

![UV-signal vs. Time](image_url)

Figure 3: The SEC trace of the product, which should have been the pure trimer. The molecule weight of the trimer is 1441.27 g/mol, and correspond to the high peak in the middle of the trace.

Unfortunately the raw trimer product contained many byproducts (see the SEC trace in Fig. 3). In the SEC trace (Fig. 3) the high peak in the middle corresponds to the trimer. It can be seen that there are byproducts at lower mass, which most likely includes the monomer, dimer and 1,4-diarylbutadiynes originated from oxidative dimerization of the unprotected monomer or a combination between the monomer and the dimer. Furthermore there are a lot of byproducts with higher mass, which most likely includes 1,4-diarylbutadiynes originated from oxidative dimerization of the unprotected dimer and trimer or a combination of those, and a polymerisation of unreacted iodo-monomer units. The molecular weight for the trimer derived from SEC is 2072 g/mol even though the molecular weight is only 1441 g/mol. The reason is that it is not the molecular weight that is determined in SEC, but the hydrodynamic volume, which is compared to polystyrene
Appendix B

standards. It means when a rigid rod polymer is compared with the randomly coiled polystyrene standards the estimation of the molecular weight is too large. James Tour has made a calibration curve for $M_n$ values of precisely defined oligothiophene ethynylene)s and oligo(phenylene ethynylene)s determined by SEC relatively to polystyrene standards. The calibration curve shows that the SEC method overestimate the mass between 50 and 66 % for oligomers with a mass of 1500 g/mol. This result agrees very well with our measurement of the $M_n$ in the case of the trimer, which is 43 % to large. It should be mentioned that the slope of the calibration curve is not constant, but depends on the actual molecular weight. It means that deviation between the molecular weight determined by SEC and the actual molecular weight increase with the actual molecular weight.

In the light of the poor yield of the trimer and the high expenses to purify a reasonable amount of the trimer by SEC, we decided to interrupt further work of making monodisperse phenylacetylene terminated oligo($1$-($2,5$-diocetyltoyl)ethyne). Instead we decided simply to use SEC to control the length of the N domains.

3.3 Controlling the length of N domains by SEC

Preparative SEC was used to divide the NPN polymer into 8 fractions (see the SEC trace in Fig. 4), and the molecular weight was determined relatively to polystyrene standards. The SEC data can be seen in table 1. Fraction 8 consisted of compounds with very low molecular weight, and was therefore discarded.

Table 1: The SEC data of the NPN polymer and the fractions. Note that the molecular weight is determined relatively to polystyrene standards, and the derivation of the SEC determined molecular weights proportional to the actual molecular weight are greatest in the region of high molecular weight.13

<table>
<thead>
<tr>
<th>Compound</th>
<th>$M_n$ [g/mol]</th>
<th>PDI</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPN crude</td>
<td>36949</td>
<td>4.09</td>
</tr>
<tr>
<td>Fraction 1</td>
<td>92275</td>
<td>1.07</td>
</tr>
<tr>
<td>Fraction 2</td>
<td>60246</td>
<td>1.21</td>
</tr>
<tr>
<td>Fraction 3</td>
<td>31612</td>
<td>1.28</td>
</tr>
<tr>
<td>Fraction 4</td>
<td>14225</td>
<td>1.19</td>
</tr>
<tr>
<td>Fraction 5</td>
<td>9239</td>
<td>1.24</td>
</tr>
<tr>
<td>Fraction 6</td>
<td>4074</td>
<td>1.39</td>
</tr>
<tr>
<td>Fraction 7</td>
<td>1534</td>
<td>1.17</td>
</tr>
</tbody>
</table>

Figure 4: The SEC trace of the NPN polymer (right axes) and the 8 fractions (left axes). The UV-signal is monitored at 383 nm, which corresponds to the absorption of the N domain.
Fraction 1 was not used because the amount of this fraction was very small and furthermore it was not possible to dissolve it again after evaporation, probably due to cross-linking of the polymer chains caused by remnants of the palladium catalysts.

3.4 Fluorescence and UV-vis measurements in solid state

In Fig. 5, we present the absorption spectra of the fractions 2 to 7 of the NPN polymer. It can be seen that the Soret band with maximum at around 460-470 nm together with the Q-bands between 600 and 700 nm of the porphyrin decrease in intensity when we are going from the fractions with low molecular weight to the fractions with high molecular weight (fraction 7 to 2). This indicates that we successfully have made NPN polymers with only one porphyrin linked into the polymer backbone, and that we with SEC are able to divide the NPN polymer up into fractions according to the length of the N domains.

From Fig. 5 it is also observed that all bands are red shifted with the increased length of the N domains. This indicates that the porphyrin is a part of a conjugated system. It can further be observed that the red shift is stopped at fraction 5, which corresponds to approximately the decamer of the N domains. These results are in agreement with the results of Jones et al. who reported studies on oligomers at a precise length of 1,4-phenylene ethynlenes.

In Fig. 6, the absorption and the fluorescence spectra of the porphyrin and the native N polymer are presented. It can be seen that there is a full spectral overlap between the absorption of the porphyrin Q-bands and the emission of the native N polymer. Further there is a partial spectral overlap between the absorption of the porphyrin Soret band and the emission of the N polymer. This opens up for the possibilities of a TS energy transfer process.

The fluorescence spectra of the fractions (Fig. 7) are all characterized by a very intense and sharp red emission peak at around 695 nm and a weak shoulder at around 750 nm. Furthermore it contains a weak broad emission in the range of 400 to 550 nm from the N domains. By comparing the fluorescence spectra of the NPN fractions with the fluorescence spectrum of the native N polymer, it can be seen, that there is a very effective energy transfer from the N domains to the P domain. Actually it is only in fraction 2 and 3, which corresponds to long N domains, where there is a weak emission from the N domains. When the length of the N domain exceeds the exciton diffusion length the probability of energy transfer to the P domain diminishes and some emission from N is observed.

![Figure 5: The UV-vis spectra of spin-coated thin films of the different NPN fractions. Note that the content of the porphyrin varies in the different fractions as expected.](image)

By comparing the fluorescence spectra of the NPN fraction in thin film with the one in solution from the literature, it can be seen that the energy transfer is significantly more effective in the solid state. The difference of the NPN in solid state compared with NPN in solution is probably that the torsion angle between the N and P domains are much larger in solution as a results of a steric hindrance. This is indicated by the fact that the Soret and the Q-bands are red shifted.

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¹ Recently experiments have shown that the N domains in the NPN polymer can be cross-bonded by remnants of the palladium catalyst and oxygen. This would cause NPN strings with more than one porphyrin unit to be incorporated into the polymer backbone. 

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around 20 and 80 nm respectively in the solid state compared to the solution, whereas the maximum of the N domains absorption only are red shifted around 10 nm. The less red shift in solution compared to solid state is probably due to a decrease in the conjugation length of the system caused by a torsion angle between the P and N domain out of plane. A torsion angle would also reduce the TB energy transfer to the P domain. In solid state it is shown that the porphyrin units are preferentially stacked, which forces P into the same plane as the N domains, and thereby allowing a more efficient TB energy transfer. The energy transfer dependence on the torsion angle indicates that it primarily occurs by TB and not by TS. TS energy transfer should not depend on the torsion angle.

![Absorption spectra](image1)

Figure 6: The absorption spectra of the porphyrin and the fluorescence spectra of the native N polymer in solid state.

![Fluorescence spectra](image2)

Figure 7: The fluorescence spectra of the dibromoporphyrin, the native N polymer and some of the 8 fractions of the NPN polymer in solid state.

### 3.5 Electroluminescence measurements

The LED devices for the electroluminescence measurements had turn on voltages of 3.3 V and were during electroluminescence measurements operated at 5.5 V, where they drew a typical current of 3-8 mA cm$^{-2}$.

In Fig. 8 the electroluminescence spectra of the native N-polymer and the crude NPN polymer are shown. By comparing the electroluminescence spectrum of the pure N-polymer (Fig. 8) with its fluorescence spectrum (Fig. 7) it can be seen that the electroluminescence of the N-domain is red shifted ~100 nm compared to the fluorescence...
spectrum. The large red shift of the N-domain indicates emission from a relaxed state. In contrast to this it can be seen that the electroluminescence of NPN is very similar to the fluorescence spectrum of the NPN fractions. We interpret this as the N domains work as very effective antennas that transport an electrically generated exciton in the N-domain to the P-domain, which is then emitted as a photon.

It should be mentioned that when we first made these LEDs, they all short-circuited. We discovered that it was remnants of the palladium from the catalysis that short-circuited the devices. We removed the palladium remnants with diethylammonium diethyldithiocarbamate with success. Later we have discovered that N,N-diethyl-(2-phenyldiazanyl)-thioformamide (see scheme 4) remove palladium nanoparticles from polymer products much more gently and efficiently.31

![Electroluminescence spectra](image)

Figure 8: The electroluminescence spectra of the native N polymer and the NPN crude polymer. Note that in the NPN a complete suppression of the N domains electroluminescence occurs and only exclusive near-infrared emission are observed.16

![Scheme 4](image)

Scheme 4: We have discovered that N,N-diethyl-(2-phenyldiazanyl)thioformamide can dissolve palladium nanoparticles by forming a complex.31

3.6 I/V characteristics of solar cell devices

The I/V characteristics of the solar cells are summarized in Table 2. In Fig. 9 the I/V curve for the bilayer N and NPN devices are shown. Especially the bilayer N cell shows a good diode behavior with a rectification ratio of ~35 at ± 1V (in dark). From the I/V characteristics three observations can be made. Firstly, the best solar cells are obtained with the native N polymer as the active layer and C80 as the electron acceptor layer. From the fluorescence and electroluminescence data we know that the N domains served as effective antennas for the P domain. The poor power conversion efficiencies for the devices with the NPN polymer must therefore be a result of the P domain that is having difficulty in delivering electrons to the electrode or to C80 in the case of the bilayer cell. Secondly, under the I/V measurements we observed that the life times of these devices were very short, which is probably due to attack of molecular oxygen on the very sensitive triple bonds in the polymer backbone. Thirdly, the overall performance of the solar cells is very poor.
Table 2: Summarized I/V characteristics for the solar cells of the N and NPN polymers. $I_s$ is the short-circuit current, $V_o$ is the open-circuit voltage, $P_{max}$ is the maximum power, FF is the fill factor, $\eta$ is the power conversion efficiency, the solar simulator generate AM 1.5 light (~100mW/cm$^2$), and the active area was 3 cm$^2$.

<table>
<thead>
<tr>
<th>Solar cells</th>
<th>$I_s$ [mA/cm$^2$]</th>
<th>$V_o$ [V]</th>
<th>$P_{max}$ [µW/cm$^2$]</th>
<th>FF [%]</th>
<th>$\eta$ [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>2.78</td>
<td>0.19</td>
<td>$1.4 \times 10^4$</td>
<td>26</td>
<td>$1 \times 10^4$</td>
</tr>
<tr>
<td>N/C$_{60}$</td>
<td>11.3</td>
<td>0.20</td>
<td>$7.2 \times 10^4$</td>
<td>32</td>
<td>$7 \times 10^4$</td>
</tr>
<tr>
<td>NPN</td>
<td>0.16</td>
<td>0.01</td>
<td>$5.9 \times 10^4$</td>
<td>37</td>
<td>$6 \times 10^7$</td>
</tr>
<tr>
<td>NPN/C$_{60}$</td>
<td>6.88</td>
<td>0.14</td>
<td>$2.8 \times 10^4$</td>
<td>29</td>
<td>$3 \times 10^4$</td>
</tr>
</tbody>
</table>

Figure 9: Linear I/V characteristics of the bilayer N and NPN devices.

4. CONCLUSION

We have presented the design of a light harvesting system based on a zinc-porphyrin linked polyphenyleneethynylene system (NPN). The system was prepared using the Sonogashira cross coupling technique, which turned out to be problematic. The problems consisted of e.g. oxidative dimerisation and cross linking of the ethynylene groups and palladium remnants from the catalyst, and we abandoned our attempt to synthesize monodisperse phenylacetylene terminated oligo(1-(2,5-diocytotolany)ethyne) at a precise molecular length.

The NPN showed a very effective energy transfer from the polymer to the porphyrin unit in solid state, while in solution the efficiency was decreased probably as a consequence of an out of plane torsion angle between the P and N domains.

There were several indications that the main energy transfer process was of the TB type. In NPN LED devices the emission of the polyphenyleneethynylene (N domains) are completely suppressed and only exclusive near-infrared emission from the porphyrin unit was observed.

From the I/V characteristics of the solar cells in the dark a good diode behavior were observed. The overall performance of the solar cells, especially for the NPN cells, was however much worse than expected.

ACKNOWLEDGEMENT

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A structural study of four complexes of the M–N$_2$S$_2$ type derived from the ligand N,N-diethylphenylazonothioformamide and the metals: Palladium, Platinum, Copper and Nickel.

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Abstract

The four electron transfer complexes: Trans-(di(N,N-diethyl- (2-phenylazinonyl)thioformamide-κS,κN))nickel, trans-(di(N,N-diethyl-(2-phenylazinonyl)thioformamide-κS,κN))copper, trans-(di(N,N-diethyl-(2-phenylazinonyl)thioformamide-κS,κN))palladium and trans-(di(N,N-diethyl-(2-phenylazinonyl)thioformamide-κS,κN))platinum have been crystallized, and their structures have been determined at low temperature.

All the complexes are of the M–N$_2$S$_2$ type. The crystals of both the nickel and the copper complex belong to the tetragonal system with the symmetry described by the space group P4$_1$2$_1$2. The molecular symmetry around the central metal atom is tetrahedral, which is unusual for the M–N$_2$S$_2$ electron transfer complexes. The crystals of the palladium and platinum complexes on the other hand belong to the monoclinic system with the symmetry described by the space group C2/c. The molecular symmetry around these metal atoms is square planar. It is demonstrated that the electron density in the ligand planes has a high degree of delocalization. Furthermore, unusually large line broadening of the $^1$H-NMR spectra was observed and investigated as a function of temperature for the palladium complex.

Comment

Thiosemicarbazides have been known for more than 100 years (Busch & Ridder, 1904), they can be transformed to an azothioformamide in a two electron oxidation process (see Fig. 1) (Pyl et al., 1963). During the sixties and the early seventies research concerning the azothioformamides progressed caused by the discovery of antifungal activity (Pluijgers et al., 1968), antibiotic activity (Kosower & Miyadera, 1972), and potential use as pesticides (Sasse & Hammann, 1971). They also form electron transfer complexes with many kind of metals analogous to Metal dithiol-
enes (Davidson et al., 1963). Especially Holm and Bechgaard have investigated the structural
and electronic properties of electron transfer complexes series of the type M—NiS2₂ (see Fig. 2)
(Holm et al., 1967; Forbes et al., 1971; Jensen et al., 1972; Bechgaard, 1973; Bechgaard, 1974;
Bechgaard, 1977).

Recently, we have discovered that azothioformamides such as N,N-diethylphenylazothioform-
amide (L) (see Fig. 3) and N,N-bis(tetraethyleneglycolmonomethyl)etherphenylazothioform-
amide have the ability to dissolve palladium and copper nanoparticles (Nielsen et al., 2005;
Nielsen et al., 2006), produced under catalytic reactions e.g. Sonogashira Cross Coupling
reactions (Sonogashira et al., 1975), by forming intensely coloured electron transfer complexes,
Pd(L)₂⁺ and Cu(L)₂⁺ (see Fig. 3), belonging to the series shown in Fig. 2.

We have demonstrated that this finding can be utilized to quantitatively analyze and remove
remnants of catalyst from organic samples (Nielsen et al., 2005; Nielsen et al., 2006). Remnants
of metal catalysts in organic compounds present a huge problem in the pharmaceutical industry
(Garret & Prasad, 2004). It is therefore promising that the azothioformamides may offer a cheap
and easy way to solve this problem. We therefore believe that the downward tendency of interest
for the azothioformamides during the eighties and nineties will reverse.

In the context of the M(L)₂⁺ compounds as electron-transfer complexes, we present here a
structural study of the Pd(L)₂⁺, Pt(L)₂⁺, Ni(L)₂⁺ and Cu(L)₂⁺ complexes. Before Hazells (1976)
surprising finding of the tetrahedral structure of the Ni(L)₂⁺ complex, it was assumed that all bis-
chelate electron transfer complexes showed a planar coordination around the central metal atom
(Bechgaard, 1974; Eisenberg, 1970). Based on powder diffraction, the Cu(L)₂⁺ complex was later
assumed to be tetrahedral like the Ni(L)₂⁺ complex (Bechgaard, 1977). In this study we reconfirm
both tetrahedral structures, and we also identify the more conventional (square) planar structure
of Pd(L)₂⁺ and Pt(L)₂⁺ that is usually found for Me-bis-dithiolene-like compounds (Eisenberg,
1970). Forbes et al. (1971) have suggested this structure for the N,N-dimethylphenylazothioform-
amide Platinum complexes.

For all four M(L)₂⁺ complexes the coordination site in L is the sulfur S and the nitrogen N₁
(see Fig. 2), so that each metal atom is participating in two five-membered rings.

To quantitatively describe the structural conformations of the M(L)₂⁺ complexes, and conclude
whether the coordination should be characterized as square-planar or flattened tetrahedral, we
have listed the relevant bond distances and angles in Table 1 to 4. These include the coordination
distance to the central metal atom and the corresponding angles involving the central metal
atom. To describe the overall geometry we have determined the dihedral angle between the
two ligand planes and the dihedral angle between the plane of the phenyl ring and the plane
of the ligand. To shed light on the electronic configuration of the ligand and the metal we
have listed all the bond lengths in the five membered chelate rings. From Table 1 and 2 it
is seen that both the Ni and Cu complexes adapt the flattened tetrahedral coordination, as
already discovered by Hazell (1976) for the Ni complex. The structures of these two complexes
are shown in Fig. 5. They belong to the tetragonal space group P4₁2₁2. The in-ring N₁–Me–
S angles are 85.61(4)°/85.48(4)°, for Ni and Cu respectively, which is below the 109° in a
regular tetrahedron and the N—Me—N angle of 144.55(9)°/144.84(7)°, which is larger than the 109° in a tetrahedral coordination. Furthermore, the dihedral angle between the two planes, defined by the five atoms in the two five-membered ligand rings, is 75.39(3)°/71.26(3)°, which is smaller than the expected 90° for perfect tetrahedral coordination. It should be mentioned that Hazell determined the dihedral angle in the Ni complex to 70.4°. The difference could be due to the different data collection temperatures (298 K versus 118.5 K in the present study). The dihedral angles between the plane of the phenyl ring and the plane of the ligand (the five-membered ring) are 28.08 (7)°/26.55 (6)°, respectively, which means that the phenyl ring is twisted out of plane compared to the five-membered ring. The smaller dihedral angles in the Cu(L)₂⁺ complex compared to the Ni(L)₂⁺ complex means that the Cu(L)₂⁺ complex has an even more flattened tetrahedral coordination than the Ni(L)₂⁺ complex. The smaller angle between the phenyl group and the ligand plane combined with the more flattened coordination imply that the distance between the two C11 atoms (the two closely spaced atoms in the phenyl groups) are longer than in the Ni(L)₂⁺ complex, 4.522 Å to 4.405 Å respectively. This indicates that the intermolecular packing forces are stronger in the Ni(L)₂⁺ compared to the Cu(L)₂⁺ complex. The structure determination of the neutral Pd and Pt complexes show that both adapt the more conventional square planar structure with the metal atom placed on a crystallographic inversion centre (see Fig. 6). This is in agreement with predictions made by Forbes et al. (1971) based on a combination of electrochemical, electron paramagnetic resonance and electronic spectral studies. As seen in Table 1 the key angles N—C—C and C—N—S are both 180°, which by symmetry are characteristic for square planar coordination. The in-ring N1—Me—S angles are 82.49 (9)/82.11 (7)°, for Pd and Pt respectively. Furthermore, the dihedral angle between the two ligand planes is exactly 0°. The angles between the phenyl group and the ligand plane are 51.04 (13)°/52.01 (8)° for the Pd(L)₂⁺ and the Pt(L)₂⁺ complexes respectively.

Blanchard et al. (2005) have demonstrated that X-ray crystallography can be used as a tool to assign the given oxidation level of the ligand. They have shown that for complexes with 1-phenyl and 1,4-diphenyl substituted S-methylisothiosemicarbazides as the ligands, the bond lengths of the N—N and C—N bonds are characteristic for the oxidation level of the ligand. E.g. if the N—N bond is rather short (1.34 ±0.01 Å) the ligand can be regarded as a π-radical anion, (L)⁻, but if the bond lengths are significantly shorter (1.24 ±0.01 Å), then the ligand can be regarded as being on its oxidized level, (L⁰). From our data (see Table 1–4) it can be seen that the Ni(L)₂⁺, Pd(L)₂⁺ and Pt(L)₂⁺ complexes all have N—N bond lengths of 1.34 ± 0.01 Å, and therefore can be rationalized as M(II) complexes with two π-radical anion ligands antiferromagnetically coupled, [Ni(II)(L)₂], [Pd(II)(L)₂] and [Pt(II)(L)₂]. Blanchard et al. have calculated the value of the following bond lengths N=C, C—S, C—N, N—N in (HN—N=C(SH)NH)⁻ to 1.294 Å, 1.897 Å, 1.342 Å and 1.343 Å respectively. Furthermore, the average S—C and C—N distances in dithiocarbamate structures are 1.70 and 1.34 Å (Eisenberg, 1970). By comparing these bond lengths with the bond lengths in the five-membered ring in the ligand described in this work, we find an indication for a large delocalization of the π electron density over the five-membered ring. The Cu(L)₂⁺ complex has a slightly smaller N—N bond length (1.32 Å) and a slightly larger C—N bond compared with
the other three complexes. This can be rationalized if one assumes that the Cu(L)$_2$$^0$ complex consists of Cu$^+$, one $\pi$-radical anion ligand and one neutral ligand, [Cu$^+(L \cdot)$] as proposed by Bechgaard (1977). Which of these two configurations, [Cu$^{1+}(L \cdot )_{\text{2}}$] (antiferromagically coupled) or [Cu$^+(L \cdot )_2$], that is the most reasonable for the Cu(L)$_2$$^0$ complex, can not be decided on the X-ray data alone. The M—N and M—S bond lengths are very similar to bond lengths of the same type observed in complexes such as Ni(S$_2$CN(C$_2$H$_5$)$_2$)$_2$, Pd(S$_2$CC$_2$H$_5$)$_2$ (Eisenberg, 1979), Pd(ESDT)PrNH$_2$Cl (Faraglia et al., 2001), Pd complexes with a-iminophenolate derivatives as the ligand (Kokatam et al., 2005), and Ni complexes with 1-phenyl and 1,4-diphenyl substituted S-methylisothiosemicarbazides as the ligand (Blanchard et al., 2005).

The Cu(L)$_2$$^0$ complex is strongly paramagnetic and it is not possible to record a $^1$H-NMR spectrum as reported earlier, while the other three complexes all give rather broad lines for the nuclei close to the metal atom (Nielsen et al., 2006). These broad lines for the Ni(L)$_2$$^0$, Pd(L)$_2$$^0$ and Pt(L)$_2$$^0$ complexes are particularly striking since Blanchard et al. (2005) did not observe broad lines for the nickel complexes with the 1-phenyl and 1,4-diphenyl substituted S-methylisothiosemicarbazides and neither did Faraglia (2001) for the Pd(ESDT)PrNH$_2$Cl complex.

We have recorded $^1$H-NMR spectra as a function of the temperature (from 450 K to 210 K) for Pd(L)$_2$$^0$ (which is the complex with the largest line broadening) to investigate whether an equilibrium exist between two conformations, e.g. a cis-trans equilibrium. The $^1$H-NMR spectra are seen in Fig. 7 (temperature range from 300 K to 210 K), together with the spectrum of L at 300 K, and in Fig. 8 (temperature range of 440 K to 300 K). The position of the lines for the protons close to Pd move as a function of the temperature. The experiment shows a clear manifest of coalescence indicative of some kind of equilibrium. However, the lines do not split up into two sets of lines upon cooling in the applied temperature range; neither do they get any sharper at higher temperatures. Based on this experiment it is therefore not possible to decide what type of equilibrium there is present, but a cis-trans equilibrium is a plausible guess.

In conclusion we have presented a detailed structural analysis of the four M—N$_2$S$_2$ complexes: Ni(L)$_2$$^0$, Cu(L)$_2$$^0$, Pd(L)$_2$$^0$ and Pt(L)$_2$$^0$, where the ligand L is N,N-diethylphenylazothioformamide. All four complexes are the central members in five different five membered electron transfer systems. We reconfirm the tetrahedral coordination of the Ni(L)$_2$$^0$ complex at 118.5 K, which was discovered by Hazell (1976) at 298 K and establish the tetrahedral coordination of the Cu(L)$_2$$^0$ complex envisaged by Bechgaard (1977). The square planar coordination of the Pd(L)$_2$$^0$ and Pt(L)$_2$$^0$ complexes envisaged by Forbes et al. (1971) are also established. We suggest that there is a high degree of delocalization of the $\pi$ electron density in the ligand planes. We further demonstrate that an equilibrium most likely is the origin for the unusually broad lines in the $^1$H-NMR spectra of the complexes.
Appendix B

PREVIEW (FM)

Experimental

The ligand L and the three metal complexes: Pd(L)$_2$$^0$, Pt(L)$_2$$^0$, Ni(L)$_2$$^0$ were synthesised as described in the literature (Jensen et al., 1972). The Cu(L)$_2$$^0$ complex was prepared by immersion of a copper wire in a concentrated thf solution of L. L dissolves the metal after which single crystals of Cu(L)$_2$$^0$ grow on the wire. The crystals can be used directly for X-ray diffraction measurements (see Fig. 4) (Nielsen et al., 2006). Crystals of the Pd(L)$_2$$^0$ and Ni(L)$_2$$^0$ complexes were made by a slow evaporation from a CH$_2$Cl$_2$ solution in a round bottom flask equipped with a glass funnel. The crystals of Pt(L)$_2$$^0$ were very tricky to grow, since slow evaporation from many different solvents led to twin crystals or needle clusters of crystals, which looked like a Christmas tree. By applying the sitting-drop vapor diffusion technique, known from protein crystallography we succeeded to grow single crystals. The equipment for the sitting-drop technique consists of glass sitting-drop rods placed in a stender dish in glass sealed with vacuum grease. The reservoir solution consists of 1 ml toluene and 0.5 ml thf, while the solvent for the drop was thf. The size of the drops was 40 $\mu$L.

Single-crystal X-ray diffraction data were collected using a Bruker Smart diffractometer with CCD area detector.

$^1$H-NMR spectra of the Pd(L)$_2$$^0$ complex and pure L were recorded at a 250 MHz Bruker NMR spectrometer in CDCl$_3$ with TMS as an internal reference. The $^1$H-spectrum of the Pd(L)$_2$$^0$ was investigated as a function of the temperature (from 300 K to 210 K in CDCl$_3$ and from 300 K to 450 K in 1,2-dichlorbenzol-d$_4$).

Compound NiL$_2$$^0$

Crystal data
C$_{22}$H$_{30}$N$_6$NiS$_2$

$M_r = 501.35$

Tetragonal

$P4_12_12$

$\alpha = 10.2852$ (3) $\AA$

$c = 22.2862$ (13) $\AA$

$V = 2357.55$ (17) $\AA^3$

$Z = 4$

$D_x = 1.413$ Mg m$^{-3}$

$D_m$ not measured

Mo $K\alpha$ radiation

$\lambda = 0.71073$ $\AA$

Cell parameters from 12044 reflections

$\theta = 2.605-31.004^\circ$

$\mu = 1.022$ mm$^{-1}$

$T = 118$ (2) K

Octahedral

Brown

$0.50 \times 0.30 \times 0.05$ mm
PREVIEW (FM)

Data collection
CCD area detector diffractometer
Omega scans
Absorption correction:
  empirical SADABS, ver2.03 (Sheldrick, 2001)
  
  \( T_{\text{min}} = 0.74, \ T_{\text{max}} = 0.96 \)

31608 measured reflections
3630 independent reflections
3436 reflections with
  \( >2\sigma(I) \)

\( R_{\text{int}} = 0.0430 \)

Refinement
Refinement on \( F^2 \)
\( R[F^2 > 2\sigma(F^2)] = 0.0347 \)
\( wR(F^2) = 0.0799 \)
\( S = 1.174 \)
3630 reflections
141 parameters
H-atom parameters constrained
\( w = 1/[\sigma^2(F_h^2) + (0.0407P)^2 + 0.3442P] \)
  where \( P = (F_h^2 + 2F_c^2)/3 \)

(\( \Delta/\sigma \))_{\text{max}} = 0.001
\( \Delta\rho_{\text{max}} = 0.606 \ \text{e} \ \text{Å}^{-3} \)
\( \Delta\rho_{\text{min}} = -0.262 \ \text{e} \ \text{Å}^{-3} \)

Extinction correction: none

Scattering factors from \textit{International Tables for Crystallography} (Vol. C)

Flack parameter = 0.021 (12)

<table>
<thead>
<tr>
<th>Bond</th>
<th>Distance (Å)</th>
<th>Error (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni—N1^1</td>
<td>1.8728 (14)</td>
<td>N1—N2</td>
</tr>
<tr>
<td>Ni—S^1</td>
<td>2.2085 (4)</td>
<td>N2—C1</td>
</tr>
<tr>
<td>S—C1</td>
<td>1.7213 (17)</td>
<td></td>
</tr>
<tr>
<td>N1^1—Ni—N1</td>
<td>144.55 (9)</td>
<td>S^1—Ni—S</td>
</tr>
<tr>
<td>N1^1—Ni—S^1</td>
<td>85.61 (4)</td>
<td></td>
</tr>
</tbody>
</table>

Symmetry codes: (i) \( y, x, -z \).

Table 1. \textit{Selected geometric parameters} (Å, °) for NiL_2O
Compound CuL₂O

Crystal data
C₂₂H₉₅Cu₆S₂
M_r = 506.18
Tetragonal
P₄₁₂₁₂
a = 10.2643 (4) Å
b = 22.3519 (13) Å
V = 2354.90 (19) Å³
Z = 4
D_x = 1.428 Mg m⁻³
D_m not measured

Mo Kα radiation
λ = 0.71073 Å
Cell parameters from 10241 reflections
θ = 2.695–27.956°
μ = 1.127 mm⁻¹
T = 118 K
Octahedral
Black
0.40 × 0.35 × 0.15 mm

Data collection
CCD area detector diffractometer
Omega scans
Absorption correction:
empirical SADABS,ver2.03 (Sheldrick,2001)
θ_min = 0.6614, θ_max = 0.8492
16807 measured reflections
2832 independent reflections
2788 reflections with
>2σ(I)
R(int) = 0.0266
θ_max = 28.01°
h = −13 → 12
k = −13 → 13
l = −28 → 28
every 0 reflections
frequency: 0 min
intensity decay: none

Refinement
Refinement on F²
R[F² > 2σ(F²)] = 0.0237
wR(F²) = 0.0702
S = 1.097
2832 reflections
141 parameters

H-atom parameters constrained
w = 1/[σ²(F²) + (0.0000P)² + 1.079P]
where P = (P² + 2F²)/3

(Δ/σ)_{max} = 0.000
Δρ_{max} = 0.361 e Å⁻³
Δρ_{min} = −0.175 e Å⁻³
Extinction correction: none
Scattering factors from International Tables for Crystallography (Vol. C)

Flack parameter = 0.012 (9)

Table 2. Selected geometric parameters (Å, °) for CuL₂O

<table>
<thead>
<tr>
<th>Bond</th>
<th>Distance (Å)</th>
<th>Error (Å)</th>
<th>Angle (°)</th>
<th>Error (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cu—N1</td>
<td>1.9219</td>
<td>0.0012</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cu—S</td>
<td>2.2756</td>
<td>0.0022</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S—C1</td>
<td>1.7220</td>
<td>0.0005</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N1—Cu—N1</td>
<td>144.83</td>
<td>0.02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N1—Cu—S1</td>
<td>85.46</td>
<td>0.02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C1—N2</td>
<td>1.3706</td>
<td>0.0018</td>
<td></td>
<td></td>
</tr>
<tr>
<td>N1—N2</td>
<td>1.3230</td>
<td>0.0020</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S1—Cu—S</td>
<td>123.60</td>
<td>0.20</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
PREVIEW (FM)

Symmetry codes: (i) y, x, -z.

**Compound PdL₂O**

**Crystal data**

C₂₂H₂₀N₆Pd₂S₂  
Mᵣ = 549.04  
Monoclinic  
\( C₂/c \)  
\( a = 16.0360 \ (16) \ \text{Å} \)  
\( b = 12.4564 \ (13) \ \text{Å} \)  
\( c = 12.3311 \ (12) \ \text{Å} \)  
\( \beta = 100.952 \ (2)^° \)  
\( V = 2418.3 \ (4) \ \text{Å}^³ \)  
\( Z = 4 \)  
\( D_x = 1.508 \ \text{Mg} \ \text{m}^{-³} \)  
\( D_m \) not measured

**Mο Kα radiation**  
\( \lambda = 0.71073 \ \text{Å} \)  
\( \theta = 2.520-27.821° \)  
\( \mu = 0.962 \ \text{mm}^{-1} \)  
\( T = 118 \ \text{K} \)  
Plate  
Green  
0.30 \times 0.25 \times 0.01 \ \text{mm}

**Data collection**

CCD area detector diffractometer  
\( \theta_{\text{max}} = 28.00° \)  
Omega scans  
\( h = -19 \rightarrow 21 \)  
\( k = -16 \rightarrow 16 \)  
\( l = -9 \rightarrow 16 \)  
8430 measured reflections  
2847 independent reflections  
2431 reflections with  
>2sigma(I)  
\( R_{\text{min}} = 0.0427 \)

**Refinement**

Refinement on \( F^2 \)  
\( R[F^2 > 2\sigma(F^2)] = 0.0489 \)  
\( wR(F^2) = 0.1071 \)  
\( S = 1.156 \)  
2847 reflections  
149 parameters  
H-atom parameters constrained

\( w=1/\left[\sigma^2(F_o^2) + (0.0364P)^2 + 13.3379P\right] \)

where  
\( P = (F_o^2 + 2F_c^2)/3 \)

(\( \Delta/\sigma \))\(_{\text{max}} \) = 0.000  
\( \Delta\rho_{\text{max}} = 1.476 \ \text{e} \ \text{Å}^{-³} \)  
\( \Delta\rho_{\text{min}} = -2.281 \ \text{e} \ \text{Å}^{-³} \)  
Extinction correction: none  
Scattering factors from *International Tables for Crystallography* (Vol. C)

**Table 3. Selected geometric parameters (Å, °) for PdL₂O**

<table>
<thead>
<tr>
<th>Bond</th>
<th>Distance (Å)</th>
<th>Angle (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pd—N₁</td>
<td>1.993 (3)</td>
<td>N₁−N₂</td>
</tr>
<tr>
<td>Pd—S</td>
<td>2.2930 (10)</td>
<td>N₂−C₁</td>
</tr>
<tr>
<td>S—Cl</td>
<td>1.741 (4)</td>
<td></td>
</tr>
</tbody>
</table>
Appendix B

PREVIEW (FM)

N1—Pd—N1 180.0
N1—Pd—S 82.49 (9)
Symmetry codes: (i) $\frac{1}{2} - x, \frac{1}{2} - y, 1 - z$.

Compound PtL₂O

Crystal data

$C_{25}H_{30}N_8PtS_2$

$M_r = 637.73$

Monoclinic

$C_2/c$

$a = 15.9317$ (7) Å

$b = 12.5300$ (5) Å

$c = 12.3798$ (5) Å

$\beta = 100.7550$ (10)$^\circ$

$V = 2427.90$ (17) Å³

$Z = 4$

$D_e = 1.745$ Mg m⁻³

$D_m$ not measured

Data collection

CCD area detector diffractometer

Omega scans

Absorption correction:

emperical SADABS ver2.03 (Sheldrick, 2001)

$T_{\text{min}} = 0.255$, $T_{\text{max}} = 0.408$

14253 measured reflections

2937 independent reflections

2632 reflections with

$>2\sigma(I)$

$R_{int} = 0.1204$

Refinement

Refinement on $F^2$

$R[F^2 > 2\sigma(F^2)] = 0.0365$

$wR(F^2) = 0.0977$

$S = 1.088$

2937 reflections

144 parameters

H-atom parameters constrained

Mo $K\alpha$ radiation

$\lambda = 0.71073$ Å

Cell parameters from 9821 reflections

$\theta = 2.516$–27.962$^\circ$

$\mu = 5.973$ mm⁻¹

$T = 118$ K

Plate

Dark green

0.25 × 0.20 × 0.15 mm

$\theta_{\text{max}} = 27.98^\circ$

$h = -21 \rightarrow 21$

$k = -16 \rightarrow 16$

$t = -16 \rightarrow 16$

every 0 reflections

frequency: 0 min

intensity decay: none

$w = 1/\sigma^2(P^2) + (0.0604P)^2 + 0.0000P$

where $P = (P^2 + 2F^2)/3$

$(\Delta/\sigma)_{\text{max}} = 0.000$

$\Delta\rho_{\text{max}} = 3.079$ e Å⁻³

$\Delta\rho_{\text{min}} = -4.143$ e Å⁻³

Extinction correction: none

Scattering factors from International Tables for Crystallography (Vol. C)
Table 4. Selected geometric parameters (Å, °) for PtL₂O

<table>
<thead>
<tr>
<th>Bond</th>
<th>Distance (Å)</th>
<th>Bond</th>
<th>Distance (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt1—N1</td>
<td>1.964 (2)</td>
<td>N1—N2</td>
<td>1.349 (3)</td>
</tr>
<tr>
<td>Pt1—S1</td>
<td>2.2932 (6)</td>
<td>N2—C1</td>
<td>1.326 (3)</td>
</tr>
<tr>
<td>S1—C1</td>
<td>1.743 (2)</td>
<td>N1—Pt1—N1</td>
<td>180.0</td>
</tr>
<tr>
<td>S1—Pt1—S1</td>
<td>82.11 (7)</td>
<td>S1—Pt1—S1³</td>
<td>180.000 (17)</td>
</tr>
</tbody>
</table>

Symmetry codes: (i) $\frac{1}{2} - x, \frac{1}{2} - y, 1 - z$.

For NiL₂O, data collection: Bruker SMART (Bruker AXS, 1998); cell refinement: Bruker SMART (Bruker AXS, 1998); data reduction: Bruker SHELXTL (Sheldrick, 2001); program(s) used to solve structure: Bruker SHELXTL (Sheldrick, 2001); program(s) used to refine structure: Bruker SHELXTL (Sheldrick, 2001); molecular graphics: Bruker SHELXTL (Sheldrick, 2001); software used to prepare material for publication: WinGX publication routines (Farrugia, 1999).

For CuL₂O, data collection: Bruker SMART; cell refinement: Bruker SMART; data reduction: Bruker SHELXTL; program(s) used to solve structure: Bruker SHELXTL; program(s) used to refine structure: Bruker SHELXTL; molecular graphics: Bruker SHELXTL; software used to prepare material for publication: WinGX publication routines (Farrugia, 1999).

For PdL₂O, data collection: Bruker SMART; cell refinement: Bruker SMART; data reduction: Bruker SHELXTL; program(s) used to solve structure: Bruker SHELXTL; program(s) used to refine structure: Bruker SHELXTL; molecular graphics: Bruker SHELXTL; software used to prepare material for publication: WinGX publication routines (Farrugia, 1999).

For PtL₂O, data collection: Bruker SMART; cell refinement: Bruker SMART; data reduction: Bruker SHELXTL; program(s) used to solve structure: Bruker SHELXTL; program(s) used to refine structure: Bruker SHELXTL; molecular graphics: Bruker SHELXTL; software used to prepare material for publication: WinGX publication routines (Farrugia, 1999).

Acknowledgements This work was supported by the Danish Technical Research Council (STVF 26–02–0174, STVF 2058–03–0016), the Danish Strategic Research Council (DSF 2104–04–0030 and DSF 2104–05–0052).
Supplementary data for this paper are available from the IUCr electronic archives (Reference: PREVIEW). Services for accessing these data are described at the back of the journal.

References


Fig. 1. Oxidation of thiourea (a) to azothioformamide (b).

Fig. 2. Electron transfer series containing five members.

Fig. 3. The structure of [i]N,N[/i]-diethylphenylazothioformamide (L) and the corresponding electron transfer complexes of the type M—N₂S₂⁺.

Fig. 4. Picture of a crystal of the Cu(L)₂⁺ complex growing on a copper wire. The crystal consumes metallic copper when growing. The wire has been immersed in a concentrated thf solution of L.
PREVIEW (FM)

Fig. 5. The X-ray structures of the Ni(L)_{2}\degree complex. The structures are viewed normal to (100).
The structure of the Cu(L)_{2}\degree complex is similar.

Fig. 6. The X-ray structures of the Pd(L)_{2}\degree complex. It can be seen that the coordination about
the metal atom is square planar. The structure of the Ir(II)_{2}\degree complex is similar.

Fig. 7. The ¹H-NMR of the Pd(L)_{2}\degree complex as function of the temperature. At the bottom the
temperature is 210 K and at the top 300 K (step of 10 K). Furthermore, the spectrum of pure L
is shown at the top.

Fig. 8. The ¹H-NMR of the Pd(L)_{2}\degree complex as a function of the temperature. At the bottom the
temperature is 300 K and in the top 450 K (step of 50 K).

\[ \text{Reactions:} \]

\[ \text{M-N}_2S_2^{2+} + 2e^- = M-N_2S_2^+ \]
\[ \text{M-N}_2S_2^+ + e^- = M-N_2S_2^0 \]
\[ \text{M-N}_2S_2^0 + e^- = M-N_2S_2^- \]
\[ \text{M-N}_2S_2^- + e^- = M-N_2S_2^{2-} \]

Figure 1.

Figure 2.
Appendix B

Figure 6.

Figure 7.

Figure 8.
Communications

Heterocycles

 DOI: 10.1002/anie.200601298


Michael Pittelkow, Theo K. Reenberg, Kim T. Nielsen, Magnus I. Magnussen, Theis L. Selling, Frederik C. Krebs, and Jørn B. Christensen

The formation of molecular crystals that possess a polar axis in the solid state is largely an unexplored area, and no generic understanding is available to this day. Materials with a polar axis (a permanent dipole moment in the solid state) potentially possess ferroelectric, piezoelectric, and pyroelectric properties. Pyroelectric solids are used as key material in a variety of components, which range from microphones and infrared detectors to frequency-doubling crystals in laser technology, just to mention a few. Examples of the design and synthesis of organic materials with pyroelectric properties are scarce because of the difficulties in predicting the crystalline properties of matter. An organic material must crystallize in one of the ten polar point groups to exhibit pyroelectric properties. Although most organic molecules possess a permanent dipole moment, few crystallize in a polar space group. Generally, derivatives based on meta-disubstituted benzene have proven to be useful as they have a tendency to crystallize in one of these ten required polar point groups and, thus, exhibit a pyroelectric effect. Apart from this rather vague design rule, little is known about the structural features that promote crystallization in a polar space group and it is only on the odd and unexpected occasion that novel pyroelectric materials appear.

Herein, we describe the synthesis, structure, and properties of an unusually stable Te-containing heterocyclic compound that crystallizes in a polar space group and possesses pyroelectric properties: 4,7-dimethoxybenzof[c]tellurophene (9, Scheme 1). The highly reactive α-quinoid heterocycles benzof[c]furran, benzof[c]thiophene, and benzof[c]selenophene have been the object of considerable interest both from a synthetic and a theoretical perspective. A particularly interesting application of these types of compounds is their role as precursors for low-band-gap conducting polymers. Cava and co-workers first reported benzof[c]tellurophene (Scheme 1), whose properties constitute a largely untouched area of tellurium-based heterocyclic chemistry largely as a result of the lack of synthetic procedures and the inherent instability of the parent heterocycle.

The treatment of 7 with elemental Te and Na in methoxyethanol yields the orange 4,7-dimethoxy-1,3-dihydrobenzof[c]tellurophene diiodide (8) in 64% yield (Scheme 2). Attempts to eliminate HI from 8 to yield 9 by

Scheme 1. Benzof[c]tellurophenes prepared by Cava and co-workers.

Scheme 2. Synthesis of 4,7-dimethoxybenzof[c]tellurophene (9).

R X:

A: Na, Te, methoxyethanol (64%); B: Et,N, benzene; C: CF3CO2Ag, benzene (99%); D: Et,N, benzene (72%).

using Et2N in benzene gave a 2:1 mixture of the desired product 9 (major product) and 1,3-dihydro-4,7-dimethoxybenzof[c]tellurophene (8; minor product; isolated after removal of 9 by reaction with maleic anhydride), as seen by 1H NMR spectroscopic analysis. As we were not able to separate the two products, 8 was converted into the bistrifluoracetate H by treatment with CF3CO2Ag in anhydrous benzene. Elimination from 11 to the desired product 9 was achieved by using Et2N in benzene. Compound 9 was purified by column chromatography on silica gel in 72% yield. The benzof[c]tellurophene (9) was stable in the solid form at room temperature and ambient atmosphere for (at least) several months in the dark (no darkening of the product was observed). It was also stable in solution with benzene and heptane, but it decomposed in CHCl3 after a few hours to give
Appendix B

a gray precipitate (probably elemental Te). Elemental analysis, cyclic voltammetry, and \(^{3}H\) and \(^{125}\)CNMR, UV/Vis, and IR spectroscopy (see the Supporting Information) were used to characterize 9. Cyclic voltammetry of 9 showed that the heterocycle was oxidized irreversibly and had an oxidation potential of 0.65 eV in CHCl\(_3\) with nBu4PF\(_6\) as the electrolyte and Pt versus the standard calomel electrode as the reference. However, we did not observe any polymerization of the monomer during the oxidation process.

Final proof of the structure of 9 came from the single-crystal X-ray structure (Figure 1). Needle-shaped yellow crystals were grown by slow evaporation of a solution of 9 in benzene at room temperature in the dark.

Figure 1. Asymmetric unit cell of the X-ray structure of 9 at 100 K.

The structure of 9 is as expected, planar, and shows similarity in bond lengths and angles with the tellurophene\(^{[17]}\) and bitellurophene\(^{[18]}\). These data also agree well with microwave spectroscopic data recorded for tellurophene\(^{[19]}\). The crystal packing of 9 exhibits few similarities with the only reported structure of a benzo[\(c\)]tellurophene\(^{[20]}\).

There are two molecules in the asymmetric unit (Figure 1). At first glance, these seem to be located randomly in space. More information is evident in the crystal packing, in which a C\(_{2}\) axis materializes and Te–Te interactions are the dominant force (Figures 2 and 3). In fact, the Te–Te distance is 4.06–4.10 Å, which is approximately 0.2 Å shorter than the van der Waals distance of Te (4.30 Å). The benzo[\(c\)]tellurophene cores are tilted with an angle of approximately 48° with respect to the C\(_{2}\) axis and are arranged two by two. The vector of the molecular dipole has a significant component in the direction of the polar axis (b axis). The vector sums of the molecular dipoles cancel in the a–c plane, whereas the vector sum gives a net dipole moment along the b axis, which results in a permanent polarization in the solid state. The distance between two Te atoms in benzo[\(c\)]tellurophene molecules that are stacked on top of each other is approximately 5.4 Å, which is too large to be accounted for by the van der Waals radius of Te. However, the distance that separates two layers of benzo[\(c\)]tellurophenes can be attributed to \(\pi\)–\(\pi\) interactions (ca. 3.5–3.6 Å) between the aromatic cores\(^{[21]}\). The unit cell is shown in Figure 2 and shows the molecular tilt with the b axis. Four molecules form a tilted square with twofold symmetry, with a distance between all Te atoms in the tilted square of 4.06–4.10 Å. This type of structure with a short Te–Te interaction and a polar space group has also been seen for the structure of 9-(9'-H-fluoren-9'-ylidene)-9H-telluroxanthene and could be a more general property of organometallum compounds\(^{[22]}\).

Figure 2. The crystal packing of 9 at 100 K. It can be seen how all the molecules are pointing in the same direction.
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Communications

The fact that all the benz[c]ellulorophene molecules point in the same direction in space introduces a permanent polarization into the crystal, thus prompting us to investigate the pyroelectric properties. The pyroelectric effect can be divided into three contributions: primary, secondary, and tertiary effects. The primary effect has origin in abrupt atomic or molecular movements within the crystal as a function of the temperature that leads to changes in the polarization in the material. This behavior is often observed in connection with phase transitions in which the spontaneous polarization either appears cancels, or changes radically; for example, if the symmetry changes from polar to nonpolar. The secondary effect is due to the change in polarization as a function of the thermal expansion (change of the volume as a function of the temperature) of the unit cell. The tertiary effect is coupled to the piezoelectric properties that pyroelectric materials invariably have. By ensuring homogeneous heating of the material and careful design of the detector geometry the tertiary effect can be eliminated from the experiment.

We investigated the X-ray structure as a function of the temperature to determine whether the observed pyroelectric effect is a primary or secondary effect. Therefore, the X-ray diffraction data of 9 were recorded at 100, 150, 200, 250, 300, 323, 343, and 373 K. An exothermic phase transition was observed at about 353 K, as shown by differential scanning calorimetry (DSC; see the Supporting Information). The X-ray data could be collected at 343 K, but the crystal blackened and its edges became rounded upon heating to 373 K. No diffraction could be observed when the temperature was lowered back down to 343 K. This behavior indicates that the observed crystal structure may not be an equilibrium structure. We, however, rule out the possibility of polymorphism as all diffraction disappears after the heating step to 373 K. We observed a decrease in the diffraction intensity as a function of time when collecting data at 343 K, and no diffraction was observed after 2 h. It, therefore, seems plausible that 9 loses tellurium at high temperature.

The structural refinement of 9 was problematic at all temperatures, except at 323 K, for which we employed constraints on the molecular geometry by forcing the benzene ring to approach a hexagonal geometry.

If the pyroelectric effect is a primary effect, a structural change as a function of the temperature, which affects the polarization, has to be observed. A change in the polarization can only happen if the angle between the $C_2$ axis and the plane of the molecule changes. We have plotted the angle between the least-squares-fitted plane and the $C_2$ axis as a function of the temperature for one of the molecules in the asymmetric unit (the same results were obtained for the other molecule in the asymmetric unit; see the Supporting Information). No significant changes were observed which means that a primary pyroelectric effect can be excluded.

In Figure 4, the polar axes of the unit cell and the cell volume is plotted as a function of the temperature, thus clearly indicating a secondary pyroelectric effect.

The secondary pyroelectric effect is solely due to the thermal expansion of the material with temperature. Compound 9 expands when heated, and as the dipoles in the unit cell stay constant the polarization decreases, thus giving the secondary pyroelectric effect as [Eq. (1)]

$$\frac{dP}{dT} = p_e(T) = \frac{Z}{a^2} \frac{1}{a^2 + b^2} \frac{d\cos(\theta)}{T}$$

where $P$, $a^2 + b^2 / \mu$, $T$, $Z$, $\mu$, and $\theta$ are the polarization, the derivatives of the linear thermal expansion with respect to the temperature, the temperature, the molecular entities per unit cell, the molecular dipole moment, and the angle between the molecular plane and the $C_2$ axis, respectively. The dipole moment of the molecules can be enhanced in the solid state, as reported previously, but this accounts for only minor changes in the overall magnitude of the pyroelectric effect.

The dipole moments of the single molecules and the unit cells were calculated at the B3LYP/3-21G level of theory using the Gaussian 98 suite of programs. The molecular coordinates from the X-ray crystal structures were used as input, and the results are summarized in Table 1. We observe that the dipole moment stays relatively constant with temperature, except for when the temperature was raised to near the phase transition.

<table>
<thead>
<tr>
<th>T [K]</th>
<th>$dP$</th>
<th>$\mu$ [D]</th>
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<tbody>
<tr>
<td>100</td>
<td>2.17</td>
<td>15.96</td>
</tr>
<tr>
<td>150</td>
<td>2.16</td>
<td>14.10</td>
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<tr>
<td>200</td>
<td>2.19</td>
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<tr>
<td>250</td>
<td>2.16</td>
<td>13.47</td>
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<tr>
<td>300</td>
<td>2.18</td>
<td>12.89</td>
</tr>
<tr>
<td>323</td>
<td>2.29</td>
<td>15.96</td>
</tr>
</tbody>
</table>

The BHLYP/3-21G. [6] Single molecule. [6] Component along the b axis in a single unit cell, the value is equal to the total dipole moment of one unit cell.

On the basis of the equation for the thermal expansion (Figure 4) and the calculated dipole moment (Table 1), it becomes possible to calculate the pyroelectric effect as a function of the temperature (Figure 5).


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Appendix B

From the point of view of application, primary pyroelectric materials have the advantage that large effects are seen near the phase transition, and therefore very sensitive detector systems can be designed. Their disadvantage is that the operation of the detector system is limited to temperatures near the phase transition. The application of the secondary pyroelectric effect in a detector system has the advantage that it can work over a large temperature range, and organic systems generally have high merit factors.29

We are puzzled by the stability of 9 relative to the other known 1,3-unsubstituted benz[e][1]phenophenes, which decompose upon isolation.31-34 We speculate that the stability of 9 is due to the steric effect that the methoxy groups in the 4- and 7-positions impose on the 1- and 3-positions of the heterocycle. This steric effect prevents these reactive positions from polymerizing, and we consider this to be a per effect.35

To summarize, we have presented the synthesis and characterization of a stable benz[e][1]phenophene derivative, including the first crystal structure of this type of heterocycle. In addition, we observe an unusual stabilizing effect of the electron-donating methoxy substituents in the 4- and 7- positions. The pyroelectric properties of 9 have been investigated, and a secondary effect was found. This discovery suggests that tellurium-containing heterocycles could be attractive in materials science.

Experimental Section

All synthetic procedures and characterization data are given in the Supporting Information. Atomic coordinates and further crystallographic details have been deposited with the Cambridge Crystallographic Data Centre, University Chemical Laboratory, Lensfield Road, Cambridge CB2 1EW, England. CCDC 299607-299612 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

Received: April 3, 2006
Published online: July 21, 2006

Keywords: density functional calculations · heterocycles · molecular pyroelectrics · X-ray diffraction

[8] For a general review about the synthesis of benz[e][naphtho-

References

Communications

Supporting Material
Synthesis, Structure and Properties of 4,7-Dimethoxybenzo[c]tellurophene, a Molecular Pyroelectric Material

Michael Pittelkow, Theis K. Reenberg, Kim T. Nielsen, Magnus J. Magnussen, Theis I. Sølling, Frederik C. Krebs and Jørn B. Christensen

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CV of compound 9……………………………………………………………...S7
DSC of compound 9…………………………………………………………….S8
Angle between mean planes vs. temperature……………………………………S9

Synthetic procedures.

2,2-Diiodo-4,7-dimethoxy-2,3-dihydro-benzo[c]tellurophene (8)
Bis-bromide (3.0 g, 9.26 mmol) (7), Te (1.18 g, 9.26 mmol) and NaI (11.1 g, 74 mmol) were suspended in 2-methoxyethanol (45 mL) and heated gently to reflux for 2 hours. The reaction mixture was cooled to room temperature and water (50 mL) was added. The orange precipitate was filtered off and washed with water (2×50 mL) and ether (50 mL). The product was dried in vacuum (2 mmHg) at 30°C. This yields the desired product as an orange solid; yield 3.25 g, 64 %. 1H NMR (CDCl3, 300 MHz) δ 6.89 (s, 2H), 4.57 (s, 4H), 3.78 (s, 6H). 13C NMR (CDCl3, 75 MHz) δ 152.4, 130.1, 110.2,
55.6, 44.0. Mp 230-233°C (decomposition). Anal. Caled for C_{10}H_{12}O_{2}TeI_{2}: C, 22.01; H, 2.22. Found: C, 22.12; H, 1.96. MS (DI) m/z 294 (M-2I^+).

4.7-Dimethoxy benzol[c]tellurophene (9)
Bis-trifluoroacetate (11) (1.32 g, 2.55 mmol) was dissolved in benzene (80 mL), Et_{3}N (2.58 g, 3.55 mL, 25.5 mmol) was added and the mixture was heated gently to reflux for 1 hour. Water (200 mL) was added and the phases separated. The organic phase was dried over Na_{2}SO_{4} and evaporated to dryness. The crude was redissolved in CH_{2}Cl_{2} and purified by dry column vacuum chromatography on silica (heptane/EtOAc mixtures) to yield the benzol[c]tellurophene as a yellow solid. Yield: 0.532 g, 72 %.

1H NMR (CDCl_{3}, 300 MHz) δ 9.50 (t, 2H, J_{Te-H} = 50.1 Hz), 5.96 (s, 2H), 3.89 (s, 6H). 13C NMR (CDCl_{3}, 75 MHz) δ 148.6, 143.6, 120.1, 97.7, 55.1. Mp 124-126°C (decomposition). IR (KBr, cm^{-1}) 3082 (m), 2982 (m), 2932 (m), 2827 (m), 1621 (s), 1502 (s), 1453 (m), 1445 (m), 1416 (m), 1354 (s), 1256 (s), 1227 (m), 1176 (m), 1135 (m), 1123 (m), 1058 (s), 959 (m), 800 (m), 737 (m), 713 (m). Anal. Caled for C_{10}H_{10}O_{2}Te: C, 41.45; H, 3.48. Found: C, 42.55; H, 3.65. MS (FAB^+) m/z: 291.96 (M+H^+).

4,7-Dimethoxy-1,3-dihydro-benzo[c]tellurophene (10)
Diodide (8) (1.47 g, 2.69 mmol) was suspended in dry benzene (150 mL) and Et_{3}N (3 mL) was added and the reaction mixture was heated gently to reflux for 1 hour. After cooling the reaction mixture to room temperature it was washed with water (2 × 100 mL), dried (MgSO_{4}) and evaporated to dryness. The resulting yellow solid compound and maleic anhydride (528 mg, 5.39 mmol, 2 eq) was suspended in anhydrous benzene (50 mL) and heated to reflux for 2 hours. The reaction mixture was then evaporated to dryness and purified directly by dry column vacuum chromatography (heptane to EtOAc with 5 % increments). This yields 10 as an off-white solid: 100 mg, 13%. Also a small amount of the highly symmetrical naphthalene compound 12 was isolated and characterized.

10: 1H NMR (CDCl_{3}) δ 6.55 (s, 2H), 4.42 (s, 4H), 3.72 (s, 6H). 13C NMR (CDCl_{3}) δ 152.1, 135.4, 108.8, 55.9, 5.7. GC-MS m/z 294 (130Te). Mp. 91-92°C. Anal. Caled for C_{10}H_{12}O_{2}Te: C, 41.16; H, 4.15. Found: C, 41.25; H, 4.01. IR (KBr, cm^{-1}) 2990, 2916, 2827, 1639, 1593, 1557, 1476, 1431, 1410, 1252, 1070, 943, 798, 774, 717.

12: 1H NMR (CDCl_{3}) δ 8.88 (s, 2H), 6.99 (s, 2H), 4.01 (s, 6H). 13C NMR (CDCl_{3}) δ 163.8, 151.1, 129.1, 125.8, 122.7, 108.5, 56.3. GC-MS m/z 258.

4,7-Dimethoxy-2-(2,2,2-trifluoro-acetoxy)-2,3-dihydro-benzo[c]tellurophen-2-yl ester (11)
Diiodide (8) (500 mg, 0.916 mmol) and silvertrifluoroacetate (405 mg, 1.833 mmol, 2 eq) was suspended in dry benzene (20 mL) and stirred for 2 hours at room temperature. The reaction mixture was filtered and the filtrate evaporated to dryness to yield the title compound as an off-white solid. Yield 470 mg, 99 %. 1H NMR (CDCl_{3}, 300 MHz) δ 6.13 (s, 2H), 4.21 (s, 4H), 3.10 (s, 6H). 13C NMR (CDCl_{3}, 75 MHz) δ 161.9, 152.8, 116.7, 112.9, 110.5, 54.8, 45.2. Mp. 174-175°C (decomposition). Anal. Caled for C_{14}H_{12}O_{6}F_{6}Te: C, 32.47; H, 2.34. Found: C, 32.69; H, 1.88. MS (DI) m/z 520 (M^+).
Appendix B

Uv-vis of compound 9 in CH$_2$Cl$_2$

![Uv-vis spectrum of compound 9 in CH$_2$Cl$_2$](image)

**Absorbance / a.u.**

**wavelength / nm**

4,7-Dimethoxy benzo[c]tellurophene

259, 315.5, 383.5

-0.5 0.0 0.5 1.0 1.5 2.0 2.5 3.0 3.5 4.0

200 250 300 350 400 450 500

CV of compound 9

![Cyclic voltamogram of compound 9](image)

The cyclic voltamogram of 9. The heterocycle was oxidized irreversibly and has an oxidation potential of 0.65 eV in CH$_2$Cl$_2$ solution using n-Bu$_4$PF$_6$ as the electrolyte and Pt vs. standard calomel electrode as the reference.
DSC of compound 9

![DSC for 4,7-dimethoxybenzolo[c]elluphene](image)

Angle between least squares plane in x-structure of 9 and the c2 axis vs. temperature

The angle between the least square plane of 9 and the C2 axes, as a function of the temperature. It can be seen that the angles remain virtually constant over the investigated temperature range.

![Angle vs. Temperature](image)
Design of porphyrin-polyphenyleneethynylene light harvesting systems

Kim T. Nielsen, Holger Spanggaard, Frederik Krebs

Polymer Solar Cell Initiative, The Danish Polymer Centre, Risø National Laboratory, PO Box 49, DK-4000 Roskilde, Denmark.

Department of Chemistry, Technical University of Denmark DTU-207, DK-2800 Kgs. Lyngby, Denmark.

Abstract

A zinc-porphyrin linked polyphenyleneethynylene system were synthesized. It was demonstrated that the polyphenyleneethynylene chains works as light harvesting antennas and efficiently could transfer the energy to the porphyrin molecules.

Goals

The Goals are divided into two main parts:

1) To synthesize a three-domain (NPN) structure consisting of two polyphenyleneethynlenes as the N domains and a porphyrin dye molecule as the P domain (see figure 1 and 4).

2) To evaluate the properties of the N domains as light harvesting antennas for the P domain.

Introduction

Normally in OPV devices the exciton diffusion length is on the order of 10 nm, which means that the active layers in these devices are very thin. The idea of this work is to synthesize a three-domain structure, NPN (see figure 1 and 4), which allows a charge separation in the interface between the domains inside the active layer. This opens the possibility to make the active layer thicker and thereby increase the power conversion efficiency.

N-Domain

P-Domain

Figure 1: The three-domain structure NPN.

It was our intention to determine the optimal length of the N domains by making mono-disperse polyphenylene ethynylene terminated oligo(3,5-dioctylphenyleneethynylene) at a precise molecular weight length by the aromatic diazobenzene synthesis illustrated in figure 2.

Figure 2: The successful Synthesis of mono-disperse N domains.

Unfortunately this failed in our hands caused by the formation of the 1,4-diazabutadiynes in each coupling step (see figure 3). These

Experimental

Figure 3: The SEC trace of the product which should have been the pure solvent. The molecular weight of the solvent is 141.27 g/mol.

Figure 4: NPN matrix synthesized by a nanoparticulate zinc coupling method, where NPN is used as catalyst.

Figure 5: Student Van der Waals well model for monomeric NPN and the P domain. This is supported by surface plasmon resonance where the strongest absorption is observed in the gap of the metal.

Conclusion

We present here the synthesis of a new zincporphyrin linked conjugated polymer. We found that Pd remnants from the catalyst result in cross linking of the ethynylene groups which cause problems when preparing the film devices. We found that an acetaldehyde side ligand efficiently removes Pd nanoparticles from polymer products.

We demonstrated an excitonic energy transfer from electrically generated excitons in the N-domain to the P-domain.

Acknowledgements

This work was performed at the Danish National Research Foundation Center of Polymer Nanodevices (NPD).

References


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Your Ref.: PJS/ALH/P10083GB 27 September 2004

PATENT APPLICATION NUMBER 0421335.1

The Patent Office confirms receipt of a request for grant of a patent, details of which have been recorded as follows:

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The application number included in the heading above should be quoted on all correspondence with The Patent Office.

Any queries on this receipt should be addressed to Janine Geran, tel. 01633 814570. All other enquiries should be directed to Central Enquiry Unit, tel. 0845 9 500 505.

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An Executive Agency of the Department of Trade and Industry
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1 The Patents Act 1977 requires the Comptroller to publish the application, including the description and claims, after the expiry of eighteen months from the earliest declared priority date or, where no earlier priority is claimed, from the date of application provided the application has not been withdrawn before completion of the preparations for its publication. The applicant may request publication to take place before the expiry of this period.

2 A request for a preliminary examination and search (with the prescribed fee) must be made within twelve months of the earliest declared priority date or, where no earlier priority is claimed, from the date of the application.

3 Failure to make the request mentioned in 2 above within the twelve month period will result in the application being treated as withdrawn. (This period can be extended by one month upon filing the prescribed fee, or possibly for a longer period at the Comptroller's discretion also upon filing the prescribed fee - in either case together with Form 52/77).

4 One or more claims and an abstract must also be filed. In the case of an application containing no declared priority date, the period allowed is twelve months from the date of filing the application. For an application which does claim an earlier priority it is twelve months from the earliest declared priority date or one month from the date of the application, whichever is the later.

5 Failure to file documents mentioned in 4 above within the prescribed periods may result in the application being treated as withdrawn.

The information above does not apply to international applications which enter the national phase under Section 89 of the Patents Act 1977. Enquiries on such applications should be addressed to the Patent Office. Tel 01633 814586.
# Request for grant of a patent

(See the notes on the back of this form. You can also get an explanatory leaflet from the Patent Office to help you fill in this form)

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<tr>
<td>3. Full name, address and postcode of the or of each applicant (underline all surnames)</td>
<td>RISØ NATIONAL LABORATORY Building 101 Frederiksborgvej 399 DK-4000 Roskilde Denmark</td>
</tr>
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<td>Patents ADP number (if you know it)</td>
<td></td>
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<tr>
<td>If the applicant is a corporate body, give the country/state of its incorporation</td>
<td>Denmark</td>
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<tr>
<td>4. Title of the invention</td>
<td>COMPLEXATION OF PALLADIUM</td>
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<tr>
<td>5. Name of your agent (if you have one)</td>
<td>W. H. Beck, Greener &amp; Co.</td>
</tr>
<tr>
<td>Address for service* in the United Kingdom to which all correspondence should be sent (including the postcode)</td>
<td>W. H. Beck, Greener &amp; Co. 7 Stone Buildings Lincoln's Inn London WC2A 3SZ</td>
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Patents Form 1/77
Appendix B

**Patents Form 1/77**

9. **Accompanying documents:** A patent application must include a description of the invention. Not counting duplicates, please enter the number of pages of each item accompanying this form:

<table>
<thead>
<tr>
<th>Description</th>
<th>23</th>
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<tr>
<td>Drawing(s)</td>
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10. **Priority documents**

| Translations of priority documents |

| Statement of inventorship and right to grant of a patent (Patent Form 7/77) |

| Request for a preliminary examination and search (Patent Form 9/77) |

| Request for a substantive examination (Patent Form 10/77) |

**Any other documents (please specify)**

11. **We request the grant of a patent on the basis of this application.**

**Signature(s):**

W. H. Bexley, Greene & Co.

**Date:** 24.09.04

12. **Name, daytime telephone number and e-mail address, if any, of person to contact in the United Kingdom:**

Mrs. Anna L. Hatt - (020) 7693 5600

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d) If you have answered YES in part 8, a Patents Form 7/77 will need to be filed.

e) Once you have filed in the form you must remember to sign and date it.

f) Part 7 should only be completed when a divisional application is being made under section 15(4), or when an application is being made under section 8(3), 12(6) or 37(6) following an entitlement dispute. By completing part 7 you are requesting that this application takes the same filing date as an earlier UK application. If you want the new application to have the same priority date(s) as the earlier UK application, you should also complete part 6 with the priority details.
COMPLEXATION OF PALLADIUM

FIELD OF THE INVENTION

The present invention relates to applications of the complexation of palladium atoms or ions by azothioformamide ligands, to a method of synthesizing such ligands, to purified polymers obtainable using such ligands and to electroactive devices comprising such polymers.

BACKGROUND OF THE INVENTION

There is a need to separate palladium, in particular palladium catalyst, from reaction mixtures and to determine the quantity of palladium present in such mixtures.

Palladium is a common catalyst. Examples of reactions using palladium catalysts are the Suzuki, Stille, Sonogashira and Heck cross-coupling reactions.

Within the field of medicinal chemistry, it is very important to be able to remove catalyst remnants efficiently from drug products.

Since the discovery in 1974 of doped polyacetylene showing electrical conductivity there has been considerable interest in conjugated polymers and their applications, for example in light emitting diodes, polymer electronic circuits and polymer solar cells. Many different conjugated polymers, for example derivatives of poly-phenylene-ethynylene and poly-phenylvinylene, can conveniently be synthesized using palladium catalysts.

A disadvantage with the use of palladium catalysts when synthesizing conjugated polymers is that palladium coordinates very well to both double and triple bonds and is therefore difficult to remove from products containing such
Appendix B

P10083US 28.09.04

bonds. For polymer products the difficulty in removing the palladium catalyst is increased because polymer molecules tend to become entangled around palladium nanoparticles, making separation by physical means such as centrifugation impossible. Work with palladium catalysed polymerisation using the Heck cross-coupling reaction showed that the removal of palladium catalyst remnants is very difficult.

Using present techniques, it is difficult to achieve complete removal of catalyst remnants in the case of polymeric products.

The importance of being able to remove all of the palladium catalyst is extremely high when the conjugated polymer is to be used in an electrical device, for example a photovoltaic or a light-emitting device, because remaining palladium nanoparticles can lead to a short circuit of the device.

The present inventors have recently showed that a standard electroactive device produced by sandwiching a polymer (chosen from iodine terminated poly[1-(2,5-dioctyltolanyl)ethylene], zinc porphyrin linked poly[1-(2,5-dioctyltolanyl)ethylenes], derivatives of polyp- phenylenevinylene and poly-2,5-(3-alkylthiophenes)) between two electrodes contained remnants of palladium catalyst and had a resistance as low as 30 Ω for the chosen device geometry (2.1 cm x 1.7 cm = 3.57 cm²). Furthermore, when a potential was applied it led to a short circuit in some parts of the device.

Previous approaches to separating metal from a mixture have included the use of ligands which complex the metal atoms or ions.
Appendix B

US 4584628 discloses removal of spent catalytic metal from a support by oxidation and complexation with a ligand which is a 1,1-dithiolate, dithiocarbamate or dithiocarbamate salt. The metal may be palladium.

US 4855400 discloses purification of polyketone polymers using a complexing agent for palladium. The complexing agents include phosphines.

EP 1203777 discloses a process for removal of iron and rhodium or platinum group metals from hydrogenated nitrile rubber using an ion-exchange resin having thiourea functional groups.

The present inventors have, with some success, attempted to use diethylammonium diethylidithiocarbamate (DDC) in solution to remove palladium from conjugated polymers as reported by Jones et al. (Jones, L.; Schumm, J. S. and Tour, J. M., J. Org. Chem., 1997, 62, 1388-1410) for solid state synthesis. The use of this compound in complexing metals is also disclosed in US 4584628.

The resistance through the standard electroactive device prepared as described above increased from 30 Ω to 0.3-2 MΩ after treatment with DDC. However, there were indications that DDC reacted with the polymers, possibly with the triple bonds. It was observed that treatment with DDC for periods of time longer than 5 min led to a dramatic change in some of the properties of the polymers. The polymers became soluble in methanol, and no electroluminescence could be observed when a potential was applied. Furthermore, DDC is a powerful nucleophilic reagent, which means that it cannot be used for polymers containing good leaving groups.

Various approaches to the removal of heavy metals using N-acetylcyesteine, polystyrene based thionaurium salts or silica
particles bearing pendant alkyl groups with terminal thiol functionalities have been reported. While they bind palladium efficiently they do not work well for polymer products since a large proportion of the polymer product is absorbed by the silica or polymer media.

There therefore remains a need for effective methods of separating palladium from reaction mixtures.

SUMMARY OF THE INVENTION

Surprisingly, the present inventors found that using azothioformamide complexing agents provided good results.

In a first aspect, the present invention provides a method of separating palladium from a reaction mixture comprising treating the mixture with an azothioformamide complexing agent to form a palladium complex and separating the palladium complex from other components of the reaction mixture.

Optionally, the palladium complex is separated by dissolving the palladium complex in a solvent to form a solution followed by phase separation. Suitable separation techniques which can be applied include precipitation, crystallization, centrifugation, filtration, decanting, adsorption, extraction, run-off of a lower liquid phase or suction of an upper liquid phase. Settling may be necessary before separation takes place.

Alternatively, the complexing agent may be an azothioformamide dimer which forms a solid (polymeric) palladium complex, and the solid palladium complex is separated by phase separation. In this case, the other components of the reaction mixture may be dissolved in a solvent to form a solution before the phase separation. Suitable solvents include ethers, esters, alcohols, ketones,
alkanes, aromatics, amides and sulfoxides. Examples include pentane, hexane, heptane, methanol, ethanol, propanol, isopropanol, butanol, water, dimethylsulfoxide, N,N-dimethylformamide, tetrahydrofuran, chloroform, dichloromethane, ethyl acetate, acetone, methyl ethyl ketone, benzene, toluene, xylene, dimethyl acetamide and N-methylpyrrolidone.

Preferably, the palladium is in oxidation state 0, more preferably in elemental form, but it may also be in other oxidation states. It is desirable for the palladium to be in the form of small particles (for example nanoparticles) since these react quickly with the azothioformamide complexing agent. Suitable nanoparticles have a diameter of 1 to 100 nm, preferably 1 to 20 nm, more preferably 1 to 10 nm, for example 1 to 5 nm.

Preferably, the other components of the reaction mixture comprise conjugated polymer or oligomer. Preferred conjugated polymers are optionally substituted poly(phenyleneethynylene) (PPE) poly(phenylene vinylene) (PPV), poly(thiophene), poly(alkylthiophene) (PAT), poly(pyridylvinylenne), poly(vinylcarbazole), polyfluorene, polyphenylene and polyarylene. Preferred conjugated oligomers are optionally substituted oligo(phenyleneethynylene) (PPE) oligo(phenylene vinylene) (PPV), oligo(thiophene), oligo(alkylthiophene) (PAT), oligo(pyridylvinylenne), oligo(vinylcarbazole), oligofluorene, oligophenylene and oligoarylene.

Preferably, the conjugated polymer or oligomer has a molecular weight of 200 to 2000000 Da.

Preferably, the complexing agent is an azothioformamide of Formula A:
wherein \( R, R', \) and \( R'' \) are each independently hydrogen or optionally substituted alkyl, cycloalkyl, alkenyl, aryl, ether, ester or amide, or a polymer attached via a linker group or an azothioformamide moiety attached via a linker group.

Suitably, the alkyl groups are \( C_1 \) to \( C_{18} \) alkyl groups. For example, the alkyl groups may be methyl or ethyl groups. Suitably, the cycloalkyl groups are \( C_5 \) to \( C_{18} \) cycloalkyl groups. For example, the cycloalkyl groups may be cyclopentyl or cyclohexyl groups. Suitably, the alkenyl groups are \( C_2 \) to \( C_{18} \) alkenyl groups. For example, the alkenyl groups may be ethenyl groups. Suitably, the aryl groups are phenyl groups. Suitably, the ether, ester or amide groups are \( C_2 \) to \( C_{18} \) ether, ester or amide groups.

Suitable linker groups are optionally substituted alkylenne, alkenylene, aryl, ether, ester or amide.

Suitably, the alkylenne groups are \( C_1 \) to \( C_{18} \) alkylenne groups. For example, the alkyl groups may be methylene or ethylene groups. Suitably, the alkenylene groups are \( C_2 \) to \( C_{18} \) alkenylene groups. For example, the alkenylene groups may be ethylene groups. Suitably, the aryl groups are phenyl groups. Suitably, the ether, ester or amide groups are \( C_2 \) to \( C_{18} \) ether, ester or amide groups.

Where \( R, R' \) and/or \( R'' \) are substituted, fluorine is a preferred substituent.
It is particularly preferred for R to be phenyl.
In preferred compounds of Formula A, R, R’, and R” are as follows:

<table>
<thead>
<tr>
<th>R</th>
<th>R’</th>
<th>R”</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ph</td>
<td>Et</td>
<td>Et (N,N-diethylphenylazothioformamide)</td>
</tr>
<tr>
<td>Ph</td>
<td>-(CH₂)₄- (with R”)</td>
<td></td>
</tr>
<tr>
<td>p-CH₃C₆H₄</td>
<td>tBu</td>
<td>H</td>
</tr>
<tr>
<td>Ph</td>
<td>tBu</td>
<td>H</td>
</tr>
<tr>
<td>tBu</td>
<td>tPr</td>
<td>H</td>
</tr>
</tbody>
</table>

In another preferred embodiment, at least one of R, R’, and R” is a polymer selected from polystyrene, polycrylate, polyethylene, polypropylene, PET, polyarylene, polyalkene, polyester, polyether or polythioether attached via a linker group.

Suitable compounds of Formula A are disclosed in DE 2145418 (where they are disclosed as acaricides and fungicides) and in Jensen et al. (1972) (where they are disclosed as complexing agents).

A unique feature of the derivatives of N,N-phenylazothioformamide is that the complex with palladium, unlike the pure ligand, has an absorption maximum at 801 nm in chloroform and 797 nm in THF (see Figs. 1 and 2). Because only very few conjugated polymers absorb at this wavelength, UV-visible spectroscopy can be used to monitor the dissolution and removal of palladium. This may be done in a quantitative manner if required.
In a second aspect, the present invention provides a method of determining palladium content in a mixture, comprising treating the mixture with an azothioformamide complexing agent to form a palladium complex, dissolving the palladium complex in a solvent to form a solution, and measuring the absorbance of the palladium complex solution.

Optionally, phase separation is carried out to separate the palladium complex solution from other components of the mixture before measuring absorbance. However, this is not necessary.

Preferably, absorbance is measured at a wavelength in the range of 770 nm to 830 nm, more preferably at 797 nm or 801 nm.

The high extinction coefficient of the palladium complex in chloroform at 801 nm (about 10000 M⁻¹ cm⁻¹) coupled with a sensitivity of 1 mAU for a typical UV-vis spectrophotometer gives a detection limit of 500 ppb using 20 mg mL⁻¹ of sample solution. Concentration of the analyte solution easily allows for a detection limit of 1 ppb when using 5g of sample.

Any preferred feature described in connection with the first aspect of the invention can be used in connection with the second aspect of the invention. In particular, the method is particularly useful in determining palladium content in a mixture comprising conjugated polymer.

Synthetic routes are known to compounds of Formula A. Jensen et al. discloses formation of N,N-diethylphenylazothioformamide (Formula A) by quinone oxidation of 4,4-diethyl-1-phenyl-thiosemicarbazide, and discloses oxidation of the complexed thiosemicarbazide using iodine.
In a third aspect, the present invention provides a method of synthesising an azothioformamide of Formula A, comprising the steps of:

a) contacting a hydrazine derivative with carbon disulphide
b) contacting the reaction mixture with an amine
c) exposing the reaction mixture to elemental oxygen.

This method can conveniently be conducted as a one-pot reaction. The method is general and the substituents can be chosen to obtain the desired solubility of the metal complex in polar or non-polar solvents.

Preferably, the hydrazine derivative is phenylhydrazine.

Preferably, the amine is a secondary amine, for example diethylamine.

Optionally, the amine is a linear or cyclic diamine, preferably a secondary diamine. In this way, azothioformamide dimer ligands may be synthesized. Suitable diamines are EtNH(CH$_2$)$_3$NH$_2$Et and piperazine.

Preferably, in step c) the reaction is exposed to air.

Preferably, a further step:
d) contacting the reaction mixture with a base and an alkyl halide is carried out between steps a) and b).

Preferably, the alkyl halide is a methyl halide and/or a an alkyl iodide. Suitable alkyl halides are methyl iodide, methyl bromide, methyl chloride or ethyl iodide, ethyl bromide or ethyl chloride. The use of a methyl halide means that during reaction with the amine gaseous methanethiol is formed as a by-product and escapes from the reaction mixture. The use of an alkyl iodide is preferred because iodide is a good leaving group.
Preferably, the base is a metal hydroxide, for example potassium hydroxide.

In a fourth aspect, the present invention provides an optionally substituted polyphenyleneethynylene (PPE) polymer having a palladium content of less than 250 ppm. Preferably, the palladium content is less than 10 ppm, more preferably less than 1 ppm. PPE polymers can only be synthesised by a palladium catalysed route, and as explained above no effective purification technique has previously been known.

Thus, PPE polymers having a low palladium have not previously been made.

In a fifth aspect, the present invention provides an electroactive device comprising such a polymer.

DESCRIPTION OF THE DRAWINGS

The present invention will be further described with reference to the non-limiting Examples, as illustrated in the Figures, in which:

Figure 1 shows the UV-vis spectra of N,N-diethylphenylazothioformamide (compound 1) and the corresponding palladium complex (compound 3) in chloroform. The spectrum is normalized to the maximum absorption, which corresponds to a molar extinction coefficient of 17100 and 29000 \( \text{(M cm)}^{-1} \) for N,N-diethylphenylazothioformamide and the complex respectively.

Figure 2 shows the UV-vis spectra of compounds 1 and 3 in THF. Compound 3 shows a moderately strong absorption with a maximum at 797 nm where compound 1 does not absorb. The maximum absorption corresponds to a molar
extinction coefficient of 8500 (M cm)^{-1} for the complex.

Figure 3 shows the ^1H-NMR spectrum of compound 3. The spectrum is measured in CDCl₃ at 300 K.

Figure 4 is a schematic illustration of the electroactive device.

DETAILED DESCRIPTION OF THE INVENTION

N,N-diethylphenylazothioformamide (1) Synthesis

Ethanol (750 mL) was placed in a 2 L three-necked round bottom flask. The ethanol was degassed with argon for 30 minutes. Phenylhydrazine (39.36 mL, 0.40 mol) was added and the solution stirred under argon with mechanical stirring. CS₂ (27.4 mL, 0.45 mol) was added dropwise over 15 minutes. A thick colourless precipitate formed. After the mixture had been stirred for an additional 30 minutes, KOH (27 g) in EtOH (200 mL) (~15% excess) was added. The precipitate dissolved and the colour changed to orange. The solution was stirred for an additional 30 minutes, where a colourless precipitate formed. CH₃I (27 mL ~15% excess) was added
quickly and nearly all the precipitate redissolved. The solution was stirred for an additional 30 minutes. The colour of the solution changed first to red and then to a cloudy very light yellow colour. The solvent was removed by evaporation and diethylamine (300 mL) was added. The solution was then refluxed for 3 days and followed by drawing small samples and subjecting them to $^1$H NMR analysis. The reaction was stopped when $^1$H NMR showed complete conversion to 4,4-diethyl-1-phenyl-thiosemicarbazide. The solution was cooled to r.t. and atmospheric oxygen was bubbled rapidly through the solution through a glass frit for 24 hours. The colour of the solution changed quickly to dark red. The solvent was now removed by evaporation and the liquid residue (a dark red oil) was dissolved in ethyl acetate (300 mL). The organic phase was then washed with brine (3 x 250mL) and dried with MgSO$_4$. After drying, the organic phase was filtered and evaporated to dryness to give 87.81 g of dark red oil. Recrystallization twice from n-heptane:acetylacetate/20:1 (3.15L) gave 1 in 60% yield (53.23 g) as orange crystals. M.p. 57.5 °C, DSC.: onset: 54.8 °C, peak: 57.5 °C, ΔH = 20175.51 J/mol.

Compound 1 is stable in air and readily soluble in most organic solvents such as methanol, chloroform, light petroleum etc.

Compound 1 dissolves palladium metal efficiently (see below). This can be directly monitored as a change in colour from the light orange colour of 1 to a dark brown green colour.

Having demonstrated the successful reaction of 1 with metallic palladium it remained to evaluate possible side reactions of 1 with conjugated polymer. The most likely
reaction between an unsaturated group such as a vinylene or ethynylene group is an electrocyclic Diels-Alder type condensation reaction, because 1 has the same unsaturation as 1,4-butadiene. Model reactions were performed between 1 and stilbene or tolane in both the presence and absence of palladium. No corresponding electrocyclic condensation products 4 and 5 were detected after prolonged reflux under argon in both the presence and absence of palladium catalyst.

The synthetic conditions of the reactions of the two models between 1 and stilbene or tolane is as follows:

**Compound 1 and stilbene:** In a 5 mL Erlenmeyer flask compound 1 (2.1 mmol, 0.5g) and stilbene (2.1 mmol, 0.38 g) were dissolved in xylene (3 mL) and flushed with argon for 5 minutes. The solution was then refluxed for 7 days. Every day a sample was drawn to test for 4. The sample was tested with NMR, MALDI-TOF and GC/MS and even after 7 days it was not possible to detect 4 in the solution. This experiment was repeated several times with both more and less solvent and with other solvents including THF, triethylamine and
toluene. It was not possible to detect 4 with the given analysis methods in any of the experiments.

**Compound 1 and tolane:** In a 5 mL Erlenmeyer flask 1 (2.1 mmol, 0.5g) and tolane (2.1 mmol, 0.37 g) were dissolved in xylene (3 mL) and flushed with argon for 5 minutes. The solution was then refluxed for 7 days. Every day a sample was drawn to test for 5. The sample was tested with NMR, MALDI-TOF and GC/MS and even after 7 days it was not possible to detect 5 in the solution. This experiment was repeated several times with both more and less solvent and with other solvents including THF, triethylamine and toluene. It was not possible to detect 5 with the given analysis methods in any of the experiments.

**Bis(N,N-diethylphenylazothioformamide-κ^2S,N)palladium-0 (3)**

**Synthesis**

\[
\text{THF (100mL) was placed in a 250 mL conical flask and degassed with argon. } N,N\text{-diethylphenylazothioformamide (1) (1.00 g, 4.21 mmol) and Palladium black (Aldrich) (0.25 g, 2.32 mmol) was added and the solution was refluxed under an argon atmosphere for 7 days. The solution was then filtered through a thick (4cm) layer of Celite on a sintered funnel.}\]
The filtrate was collected and evaporated to dryness. The residue was then purified by dry column vacuum chromatography with n-heptane/1,2-dichloroethane in the ratio 10:1 as the eluent. The eluent was removed by slow evaporation to provide dark green crystals. The yield was 164 mg (14%) of pure compound 3.

$^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS)$\delta$: 1.218 (t (J = 6.9 Hz), 12 H), 4.177 (broad s, 8 H), 5.431 (broad s, 2 H), 6.909 (broad s, 4 H), 7.575 (broad s, 4H). $^{13}$C NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 16.309, 44.819, 123.692, 131.381, 175.869. MALDI-TOF MS [M+H$^+$] = 549 (the isotopic pattern was confirmed); Anal. Calcd. for C$_{22}$H$_{29}$N$_6$PdS$_2$: C:48.13, H:5.51, N:15.31, S:11.68. Found: C:48.05, H:5.65, N:14.41, S:11.50.

**Synthesis of 3 from palladium nanoparticles prepared in-situ**

![Chemical structure](image.png)

Palladium (II) acetate (268 mg, 1.19 mmol) was dissolved in DMSO (50 mL) and flushed with argon. Under an argon atmosphere phenylhydrazine (74 mg, 0.69 mmol) was added. The colour of the solution shifted quickly from brownish orange to black. After stirring for 10 minutes compound 1 (680 mg, 2.87 mmol) was added and the solution was refluxed under an argon atmosphere for 1½ hour. The solution was then poured into 600 mL water, and was left standing in the hood without an argon atmosphere for 1½ day.
The colour of the mixture had then changed from very dark brown to dark greenish blue. The mixture was washed four times with 400 mL n-heptane/ethyl acetate mixture (1:1). All organic fractions were collected and dried with magnesium sulphate. After filtration the solvent was removed by evaporation. The residue was purified by dry column vacuum chromatography with n-heptane/ethyl acetate as the eluent in a ratio of 10:1. The eluent was removed by evaporation to provide dark green powder. The yield was 480 mg (73 %) of compound 3. An analytical sample for determination of the molar extinction coefficient was obtained by additional dry column chromatography.

$^1$H NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 1.218 (t ($^3$J = 6.9 Hz), 12 H), 4.177 (broad s, 8 H), 6.431 (broad s, 2 H), 6.909 (broad s, 4 H), 7.575 (broad s, 4H). $^{13}$C NMR (250 MHz, CDCl$_3$, 300K, TMS) $\delta$: 16.309, 44.819, 123.692, 131.381, 175.869. MALDI-TOF MS [M+H$^+$] = 549 (the isotopic pattern was confirmed); Anal. Calcd. for C$_{32}$H$_{30}$N$_4$Pd$_2$: C:48.13, H:5.51, N:15.31, S:11.68. Found: C:48.05, H:5.65, N:14.41, S:11.50.

**Removal of Palladium from Conjugated Polymer**

\[
Pd(0) + R_1-N=N-S \rightarrow R_2 + N-N-S \rightarrow R_3
\]

Insoluble

Soluble

Polymer synthesised by the method of Krebs et al. (Krebs, F. C.; Nyberg, R. B. and Jørgensen, M., Chem. Mater., 2004, 16, 1313-1318) is dissolved in minimum THF (typically 1 mL/20 mg sample) under an argon atmosphere. 10 times excess of compound 1, compared with the palladium, is
added and the solution is stirred for between 30 minutes and 1 hour, optionally with reflux. The polymer is precipitated in excess methanol (typically 10 times the amount of THF), filtered and washed with more methanol. The polymer is redissolved in THF under an argon atmosphere and precipitated in methanol, filtered and washed with more methanol. The clean polymer is then dried in a vacuum oven.

**Analysis of Palladium content in a Polymer Sample**

The polymer sample is very precisely weighed out and dissolved in THF (typically 20 mg/ml) under an argon atmosphere. Compound 1 (10 times excess with respect to the amount of Pd in the catalyst) is added and the solution is stirred for between 30 minutes and 1 hour.

The solvent is evaporated and the residue is dissolved in a precisely known volume of solvent. A UV-vis spectrum of the solution is recorded. The complex (compound 3) has an absorption maximum at 801 nm in chloroform with an extinction coefficient of 10300 M\(^{-1}\) cm\(^{-1}\). From the UV-vis spectrum the concentration of Pd in the polymer can be calculated.

There is a significant different difference in the UV-vis spectrum extinction coefficient and wavelength of maximum absorption in THF and chloroform.

**UV-vis (CHCl\(_3\) @ 25 °C):**

\[ \lambda_{\text{max, chloroform}} = 801 \text{ nm} \quad \varepsilon_{\text{chloroform}} = 10300 \text{ M}^{-1} \text{ cm}^{-1} \]

**UV-vis (THF @ 25 °C):**

\[ \lambda_{\text{max, THF}} = 797 \text{ nm} \quad \varepsilon_{\text{THF}} = 8500 \text{ M}^{-1} \text{ cm}^{-1} \]
Examples of Palladium-catalysed Polymerisation to form Conjugated polymers

Example 1: Poly[1-(2,5-dioctyl-tolanyl)-ethylenylene] (PPE)

A 100 mL round-bottom flask was dried using a heat gun and flushed with argon. 1-iodo-2,5-dioctyl-4'-ethynyltolane (1.36x10^{-3} Mol, 0.750 g) was dissolved in piperidine (40 mL) and flushed with argon for 30 minutes, before the catalyst Pd(PPh₃)₄ (= 2 mol %) and the co-catalyst CuI (= 10 mol %) were added under an argon atmosphere. The solution was stirred under an argon atmosphere overnight at room temperature. A yellow precipitate quickly formed. The colour of the solution was yellow during the reaction. The following day the reaction mixture had solidified. Dry THF (50 mL) was added and the solution was heated to near reflux temperature. The precipitate dissolved and the solution was stirred for another 5 hours. The solution was then poured into 300 mL methanol and a yellow precipitate formed. The suspension was filtered through a glass filter-funnel. The filter cake was washed with both cold methanol, cold diethyl ether and was sucked as dry as possible. The wet product was kept for further use. Attempts to dry the product in a vacuum at RT gave a darkened product that was no longer soluble in common solvents. The total yield based on a completely dried sample was 0.35 g (95%). The wet product that was kept weighed 0.6 g and contained 40% solvent. H NMR δ: 0.76-0.95 (m, 6H), 1.13-1.50 (m, 20 H), 1.50-1.83 (m, 4H), 2.70-2.90 (m, 4H), 7.31-7.43 (broad s), 7.43-7.60 (broad s) (7.31-7.60, 6H).
Appendix B

Pd-analysis of the PPE polymer: The PPE polymer (0.1028 g) was dissolved in 3 mL THF under an argon atmosphere. Compound 1 (0.035 g) was added and the solution was stirred under an argon atmosphere for 1 hour. The solution was diluted to a volume of 5 mL and the UV-vis spectrum was recorded. This gave an absorbance of 0.441 corresponding to a concentration of 3 of 51.9 μM. The amount of Pd in the polymer was thus 0.259 μmol ~ 27.60 μg Pd in 0.1028 g of polymer. This is equivalent to 268 mg Pd kg⁻¹ or 268 ppm.

Example 2: Poly-1,2′′′-(2,5-dioctyl-1,4-phenylene-1′,2′-vinylene-1′′,4′′-phenylene-1′′′,2′′′-vinylene) PPV

1,4-Dibromo-2,5-dioctylbenzene (1 g, 2.17 mmol), 1,4-divinylbenzene (0.28 g, 2.17 mmol) and triethylamine (10 mL) were mixed in dry THF (50 mL) and the mixture was degassed with argon for 5 minutes. (PPh₃)₂PdCl₂ (250 mg) was added and the mixture was heated to reflux. After 36 h the mixture was poured into methanol to precipitate the bright yellow product. The mixture was then filtered and dried to give a yellow solid in 85% yield (0.79 g).

Pd-analysis of the PPV polymer: The PPV polymer (0.0577 g) was dissolved in 5 mL THF under an argon atmosphere. Compound 1 (0.035 g) was added and the solution was stirred under an argon atmosphere for 1 hour. The solution was diluted to 50 mL and a sample of 3 mL was drawn and diluted to 20 mL. From this sample the UV-vis spectrum was recorded. This gave an absorbance of 0.247 which corresponds to a concentration of 3 of 29 μM. The dilution factor of the 50 mL solution was 6.67 corresponding to a total volume of 333.3 mL. The amount of Pd in the polymer was thus 9.7 μmol.
Appendix B

- 1030 µg Pd in 0.0577 g of polymer. This is equivalent to 17860 mg Pd kg⁻¹ or 17860 ppm.

Example 3: Poly- (3-hexylthiophene) PAT.

DMSO (30mL) and DMF (30mL) were degassed using argon and (PPh₃)₂PdCl₂ (90mg, 0.128 mmol, catalyst) was added and the mixture was heated to 75 °C with stirring. 2-Bromo-3-hexyl-5-trimethylstannylthiophene (1.75g, 4.26 mmol) was added dropwise from a syringe. The colour of the mixture quickly changed to clear yellow and after 1 minute it became cloudy and orange and after another minute it became red/brown and a material started to separate. After 3 days under argon the mixture was a light yellow clear solution with a black precipitate. The mixture was filtered and the solid washed with methanol (5 x 50mL). Drying gave a black solid 0.69g (97%).

Pd-analysis of the PAT polymer: The PAT polymer (0.035 g) was dissolved in 5 mL THF under an argon atmosphere.

Compound 1 (0.030 g) was added and the solution was stirred under an argon atmosphere for 1 hour. The solution was diluted to 50 mL. From this sample the UV-vis spectrum was recorded. This gave an absorbance of 0.228 which corresponds to a concentration of 3 of 26.8 µM. The amount of Pd in the polymer was thus 1.34 µmol ~ 143 µg Pd in 0.035 g of polymer. This is equivalent to 4073 mg Pd kg⁻¹ or 4073 ppm.

Standard electroactive device for fabrication

The substrates are 10 Ω square² ITO slides which have been etched at one end by immersion in a 65 °C aqueous solution of HCl(aq) 20% and HNO₃(aq) 5% for typically 60
seconds. The etched substrates are cleaned by immersion in isopropanol and subjected to ultrasound for 10 min. After being blown dry in a stream of dry argon a layer of PEDOT:PSS is applied by spin-coating an aqueous solution of PEDOT:PSS (1.3% wt.) containing sorbitol (2% wt.). The substrates are placed in an oven at 100 °C and the temperature is ramped up to 180 °C over 1h and kept at 180 °C for 1h. After cooling to room temperature the polymer solutions (in chlorobenzene at a concentration of 2-10 mg mL⁻¹) are micro-filtered and spin coated at 1500 rpm onto the substrates. The typical film thicknesses are 50-100 nm as measured by a DEKTAK 3030 and the typical absorbance of the films is 0.25-1.0 absorbance units. The geometry of the devices is shown in Fig. 4.

The aluminium electrode is evaporated onto the device at a pressure of < 5·10⁻⁶ mBar. After application of the electrodes the samples are removed from the evaporator chamber and electrical connections are made using conducting silver epoxy glue. The active area of the device is 3 cm² (20 mm x 15 mm).

The resistance is measured using a Keithley 2400 Sourcemeter. Typical experimental variation between devices is < 10%.
### Appendix B

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Pd content before/after (ppm)</th>
<th>Resistance before/after (kΩ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPV</td>
<td>17860/&gt;0.1</td>
<td>0.10/30</td>
</tr>
<tr>
<td>PPE</td>
<td>268/&gt;0.1</td>
<td>0.03/45</td>
</tr>
<tr>
<td>PAT</td>
<td>4073/&gt;0.1</td>
<td>0.06/24</td>
</tr>
</tbody>
</table>

Pd(0) + \[R_1-N=N-N(R_2)\_N^3-N=N-R_1\] \[\rightarrow\] \[\text{Soluble or insoluble depending on R-groups}\]

**Synthesis of azothioformamide dimers**

1. \[\text{1)} H \rightarrow H \] \[\text{2)} \text{atmospheric oxygen}\]
2. \[\text{1)} \text{atmospheric oxygen}\] \[\text{2)} \text{atmospheric oxygen}\]

The materials were prepared by a procedure analogous to the one described for N,N-diethylphenylazothioformamide (1).

The preferred embodiment of the invention has several advantages.
The derivatives of N,N-phenylazothioformamide coordinate very strongly to palladium like DDC, but unlike DDC they can dissolve metallic palladium. The derivatives of N,N-phenylazothioformamide have the advantage compared to DDC that they do not react with the polymer, but coordinate to and dissolve metallic palladium remnants in the product mixture.

This means that using the N,N-phenylazothioformamide complexing agent, conjugated polymers can be synthesized by a convenient palladium catalysed route and can be purified sufficiently to be used in electroactive devices.

Also, the derivatives of N,N-phenylazothioformamide also have the advantage that they form a complex with palladium which has a distinctive UV absorption maximum in a region where conjugated polymers do not absorb. Thus, the concentration of palladium in a sample of conjugated polymer can be determined without the need to separate the palladium from the conjugated polymer.
Claims

1. A method of separating palladium from a reaction mixture comprising treating the mixture with an azothioformamide complexing agent to form a palladium complex and separating the palladium complex from other components of the reaction mixture.

2. A method as claimed in Claim 1, wherein the palladium complex is separated by dissolving the palladium complex in a solvent to form a solution followed by phase separation.

3. A method as claimed in Claim 1, wherein the complexing agent is an azothioformamide dimer which forms solid palladium complex, and wherein the solid palladium complex is separated by phase separation.

4. A method as claimed in Claim 3, wherein the other components of the reaction mixture are dissolved in a solvent to form a solution before the phase separation.

5. A method as claimed in Claim 2, wherein the solvent is selected from pentane, hexane, heptane, methanol, ethanol, propanol, isopropanol, butanol, water, dimethylsulfoxide, N,N-dimethylformamide, tetrahydrofuran, chloroform, dichloromethane, ethyl acetate, acetone, methylethylketone, benzene, toluene, xylene, dimethylacetamide and N-methylpyrrolidone.
6. A method as claimed in Claim 1, wherein the other components of the reaction mixture comprise conjugated polymer or conjugated oligomer.

7. A method as claimed in Claim 1, wherein the palladium is in oxidation state 0.

8. A method as claimed in Claim 7, wherein the palladium is in elemental form.

9. A method as claimed in Claim 1, wherein the complexing agent is an azothioformamide of Formula A:

\[
\begin{array}{c}
\text{R-}N\equiv S\equiv N-R' \\
\text{R''}
\end{array}
\]

Formula A

wherein R, R' and R'' are each independently hydrogen or optionally substituted alkyl, cycloalkyl, alkenyl, aryl, ether, ester or amide, or a polymer attached via a linker group or an azothioformamide moiety attached via a linker group.

10. A method as claimed in Claim 9, wherein at least one of R, R' and R'' is a polymer selected from polystyrene, polyacrylate, polyethylene, polypropylene, PET, polyarylene, polyalkene, polyester, polyether or polythioether attached via a linker group.

11. A method as claimed in Claim 9, wherein R is phenyl.
12. A method of determining palladium content in a mixture, comprising treating the mixture with an azothioformamide complexing agent to form a palladium complex, dissolving the palladium complex in a solvent to form a solution, and measuring the absorbance of the palladium complex solution.

13. A method as claimed in Claim 12, wherein the palladium complex solution is separated from other phases of the mixture before measuring absorbance.

14. A method as claimed in Claim 12, wherein absorbance is measured at a wavelength of 770 to 830 nm.

15. A method as claimed in Claim 14, wherein absorbance is measured at a wavelength of 801 nm.

16. A method of synthesising an azothioformamide of Formula A, comprising the steps of:
   a) contacting a hydrazine derivative with carbon disulphide
   b) contacting the reaction mixture with an amine
   c) exposing the reaction mixture to oxygen.

17. A method as claimed in Claim 16, wherein the amine is a diamine.

18. A method as claimed in Claim 16, wherein in step c) the reaction is exposed to air.

19. A method as claimed in Claim 17, wherein a further step:
d) contacting the reaction mixture with a base and an alkyl halide
   is carried out between steps a) and b).

20. An optionally substituted polyphenyleneethynylene (PPE) polymer having a palladium content of less than 250 ppm.

21. An optionally substituted PPE polymer as claimed in Claim 20, having a palladium content of less than 10 ppm.

22. An optionally substituted PPE polymer as claimed in Claim 21, having a palladium content of less than 1 ppm.

23. An electroactive device comprising a polymer as claimed in Claim 20.
ABSTRACT

COMPLEXATION OF PALLADIUM

A method of separating palladium from a reaction mixture comprises treating the mixture with an azothioformamide complexing agent to form a palladium complex and separating the palladium complex from other components of the reaction mixture.

A method of determining palladium content in a mixture comprises treating the mixture with an azothioformamide complexing agent to form a palladium complex, dissolving the palladium complex in a solvent to form a solution, and measuring the absorbance of the palladium complex solution.

A method of synthesising an azothioformamide comprises the steps of:

a) contacting a hydrazine derivative with carbon disulphide

b) contacting the reaction mixture with an amine
c) exposing the reaction mixture to oxygen.
Fig. 1

Fig. 2
Fig. 3
Fig. 4
Risø’s research is aimed at solving concrete problems in the society.

Research targets are set through continuous dialogue with business, the political system and researchers.

The effects of our research are sustainable energy supply and new technology for the health sector.