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Design and synthesis of a new polymer: study of supramolecular structure and functional properties

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I. 1- INTRODUCTION.

Organic molecules for functional applications: solar cells

The efforts on studying a system that could convert light in electrical Energy started in early 1970's in order to reduce the environmental impact of fossil fuels and to offer an alternative to the dangerous nuclear power.

A solar cell is a device that can use the photoelectrical effect (discovered by Baquerel in 1839) to convert light in a lasting current, so it acts as an electrical generator.

In a solar cell the "photogenerated" electrical charges are separated basing on their sign and then collected by two electrodes having different potential. Today's solar cells are realized using various materials and many structures, but solar cells belonging to the first generation, based on p-n junction, are the most efficient and the most diffused in the global market.

In Silicon-based solar cells the n- or p- doping is related to the introduction of an extra-band between the Valence band and the Conducting band belonging to the Silicon.

This treatment is made in order to obtain a constant electrical current in a cell, thanks to the introduction of an Electrical Field.

In case of p- doping, some impurities (e.g. Boron atoms) are introduced in Silicon's crystals; as a consequence, an empty band is introduced in the Energy gap between the Valence band and the Conducting Band of Silicon. After absorbing a photon having the suitable Energy value, electrons can cross the gap between the Silicon's valence band and the new "implanted" band.

The same mechanism is at the basis of n-doping; in this case, an extra band full of electrons is introduced thanks to the implantation of P atoms in Silicon's crystals. In this case electrons can be promoted from dopant's full band to Silicon's conducting band.

Obviously, when an electron moves from a band to another, it leaves a "hole", an empty space that can be filled by another electron. In the recombination process, every free electron can join a hole thus returning a part of its energy to the system in form of heat.

If we consider a Silicon Crystal, when the p-doped and the n-doped zones are joined, the interface between the 2 materials is called p-n junction. An electron transfer from n- to p- due to the majority carriers happens at the interface between the crystals, and obviously a hole transfer from p- to n- happens at the opposite. This behaviour determines at the p-n interface a *depletion zone*, having a double layer of opposites charges and no free charge.

The lighting of the p-n junction generates an exciton. After that, an Electrical field at the depletion zone (called "built in field"), separates the electrons from their holes and force them to the n- zone (the electrons) and the p- zone (the holes).

After that, minority carriers (the electrons in the p-, the holes in the n- material) diffuse to the junction, from where they are swept away, so they accumulate on the other side of the junction, where they become majority carriers (step 1, 2 and 3 in part a in **Figure 1**).



Figure 1. The process of separation, diffusion and accumulation of charge carriers in a simple p-n junction.

This knowledge allowed to develop different varieties of devices based on Silicon; in particular, we can distinguish:

- Amorphous Silicon-based systems
- Polycristalline Silicon-based systems

At today, amorphous Silicon-based systems reached the 15% maximum efficiency in converting light in electrical power, while polycrystalline Silicon-based systems have reached more than 20% maximum efficiency.

Although the good performances in terms of efficiency, maybe one of the most important reason for the lack in widespreading the solar cell's technology is the cost of the solar panels themselves, and the cost of installation, which represents approximatively half of the total prize of a domestical solar implantation.

The research of cheaper materials that can be used in industrial processes is a possible solution to this problem; in alternative a new industrial process to obtain the same device in a cheaper way could be elaborated.

This consideration has led to the development of devices based on Cadmium Telluride (CdTe) and Copper Indium Gallium Selenide (CIGS), which represent the second generation of thin film technologies because they can be manufactured by roll-to-roll processes. These processes have the advantage to be cheaper to manufacture than crystalline silicon and, as a consequence, they lead to lower electricity costs, which are extimated to be around \$1-2/Wp at high productions volumes and efficiency (5-12%).

Since 1980's, thanks to the development of the first Organic Thin-Film Transistor (OTFT's), organic materials (both polymer- and small based molecules based devices)¹⁻³ have attracted scientific interest in the field of electronics.

Organic materials are used as charge carriers in a variety of potential applications, spacing from optoelectronics to electrochemical devices, such as Organic Light Emitting Diodes (OLEDs), sensors, electrochromic devices and Organic Field Effect Transistors (OFETs).

Molecules having a conjugated system are promising materials for photonic applications since their inherent dipole enhances the nonlinear optical response properties of the material.⁴⁻⁵ In fact, the presence of a π - electron system in a molecule allows it to absorb sunlight; after that "photogenerated" charge carriers are created and then they can run across the material.

In general, organic semiconductor can be classified as "intrinsic wide band gap semiconductors" (band gap above 1.4 eV) down to "insulators" (band gap above 3 eV) with a negligibly low intrinsic charge carrier concentration at room temperature in the dark. Chemical, photochemical or electochemical doping can be used to introduce extrinsic charge carriers into organic semiconductors. For example photoinduced electron transfer from a Donor- to an Acceptor-type molecule can be used to introduce free charge carriers.⁶

Unfortunately, low charge mobility hindered the developing of practical applications of these materials. Nevertheless, organic materials are potentially advantageous respect to the inorganic counterparts because of the possibility to develop a low cost synthesis, the possibility to manufacture thin devices by common surface modification methods, or the possibility to create flexible devices that can be produced in large scale at relatively low cost.⁷

In 1991, thanks to the development of the Gratzel's cell,⁸ chemists returned on organic solar cell's development thanks to the concept of Dye Sensitized Solar Cell's (DSSC).

The Gratzel's cell is composed by of two parts, an organic molecule (a metal iridium or platinum complex) is covalently bonded to a inorganic support of TiO_2 . (**Figure 2**) The organic complex acts as light-harvester; it absorbs a photon and after that it starts the formation of an exciton (an anion or cation radical) that can be transferred to the titanium oxide. Obviously, in this red-ox process the organic molecule has to re-generate its oxidation state, so this system needs to take place in a medium (a solution) containing in the most of cases the I^7/I_2 couple.



Figure 2. The Gratzel's cell scheme

Despite the fact that, at today, Gratzel system is the most efficient organicinorganic hybrid system because of the broad spectral range of absorption due to the inorganic semiconductor, this kind of "hybrid" cells has the great disadvantage of needing an electrolytic solution in order to regenerate the oxidation states of the molecules involved in the process. The use of a solution implies some technical difficulties to develop efficient cells because of the risk of leaks in the device, corrosion and the impossibility to realize flexible solar cell.

Moreover, from a performance point of view, Gratzel's cells are far from Silicon cells in terms of performance, in fact the best laboratory cell reaches the 10% maximum efficiency under illumination, with module efficiencies ranging from 3% up to 5%.⁹ Another disadvantage is the request of a broad light absorbing spectra organic molecule in order to act a very efficient light harvesting.

The amount of light that can be absorbed by a single monolayer is poor because the area occupied by a single molecule is much larger than its optical cross section for light capture. Thus, to improve the photovoltaic efficiency it's better to use a porous, nanostructured film of very high surface instead of a flat one. The principle is similar to what that happens in leaves, where the thylakoid vesicles are stacked in order to improve the light harvesting of chlorophyll.¹⁰

Another technology connected to the use of carbon-based molecules is known as Organic Photovoltaics.¹¹ In this case a solution containing the organic molecule is

used to modify a surface thanks to printing or coating techniques. The drying of the solvents allows to deposit the organic molecule on the surface and to obtain a solid state photovoltaic device.¹² This is a valuable approach because of the possibility of using simple, high speed and large scale manufacturing processes.

The key feature to explain how a OPV cell works is the mixture of n-type and p-type charge carriers in the active layer that is used to modify the surface.

Obviously, as organic molecules are the effective actors of charge-carrying, it's not convenient to explain charge transfer in terms of band structure, but in terms of frontier orbitals, so it's better to talk about HOMO (High Occupied Molecular Orbital) and LUMO (Low Unoccupied Molecular Orbital) instead of Valence band and Conducting band.

Organic molecules can withdraw or donate electron or holes; for the sake of simplicity we will talk about Electron-Withdrawing and Electron-Donor molecules; an Electron-Donor (D) molecule has low Ionization Potential and thus a high lying HOMO energy, an Electron-Acceptor molecules (A) have a high Electron Affinity (EA) and, as a consequence, a low lying LUMO Energy. The charge transport happens at the interface between D and A.¹³

The acceptor specie (the n-type molecule) must have a lower LUMO level respect to the donor molecule in order to have a higher probability to verify the charge transfer instead of the return to the ground state.



Figure 3. Different steps during the interaction between a Donor and an Acceptor Organic Molecule involved in a solar cell.

After absorbing a photon, whose energy is larger than the gap between HOMO and LUMO, there thermalization of the holes and electrons near the top of valence and conduction band respectively happens.

In the organic junctions part of this process is similar to that described in Fig.1.

In fact, after the absorption of a photon, we have the formation of excitons that diffuse in the heterojunction (step 2 and 3 in **Fig. 3**), where they dissociate and transfer a electron (or a hole) into the acceptor (or donor) layer (step 4 in **Fig. 3**). This isn't the only pathway that can be followed; the deactivation of the donor exciton (through the Forster energy transfer) leads to the acceptor's excitation; after that, the quenching caused by charge transfer can happen. When the acceptor core is excited, there is a hole transfer moving the hole from the HOMO of the acceptor to the HOMO of the donor can happen. This can lead again to a non emissive charge separated state. Thus, no matter if the donor or the acceptor specie is excited, photoinduced electron (or hole) transfer follows that quenches both the donor and the acceptor fluorescence and leads to a non-emissive photoinduced charge separated state. (**Fig.4**)



Figure 4. Possible patways that can be followed by the couple Donor-Acceptor Molecule after a photon's absorption.

The physical behaviour standing at the base of this phenomenon allows, in principle, a possible all-organic solution to the photovoltaic device development. It could be thought to synthesize organic molecules acting as light harvesters in order to form excitons (charged species having an uncoupled electron).¹⁴

This kind of solution could make chemists able to develop cheaper and more flexible solar cells, but a crude approach to this problem leaves several points to take care.

In fact, in order to convert the most of the light in electrical energy, the molecule acting as light harvester has to absorb the widest visible spectrum as possible, and most of the organic semiconductors investigated today absorb in the visible range, while sun has its maximum photon density at around 700 nm; moreover organic molecules have an intrinsic limited durability because when electrons are excited to higher molecular orbitals, anti-binding states arise and the probability for decomposition of the compound increases.¹⁵ Moreover, an efficient light harvesting process needs not only a Donor-Acceptor interface, but an efficient charge separation and transport. These two features may be guaranteed by the continue

interface between Donor and Acceptor and by a continuous path for the transport of separated charges to the contacts. These are the conditions allowing light harvesting to happen in ultrafast-time scale in order to obtain separated charges that could be collected by the electrodes in form of a current that could be measured. ¹⁶

This is a master point for the Gratzel's cell, but in order to allow a better industrial scale up, scientists have tried to develop systems based on the use of blends¹⁷ of acceptor (n-type dopants) and donor (p-type dopants) molecules.

These devices were based on simple blends of Fullerenes and P3HTs, but the results were poor respect the theoretical calculations.¹⁸ In fact these molecules have a poor reciprocal solubility, so they tend to segregate themselves after some time, so the exciton's formation can happen only at the interface between the acceptor and donor layer. Obviously this is an aspect limiting the photoconversion, the efficiency and the durability of the cell. To the best of our knowledge, the best results in terms of photoconversion were obtained by Heeger and Guo, recording over the 5,5 % as efficiency.¹⁹⁻²⁰

Maybe the most simple solution to this problem was binding the acceptor molecule to the donor one thanks to the organic synthesis and, in order to rise the solubility of the product molecule, using, for example, the oligomeric species of the P3HT instead of the polymer.²¹⁻²²

This kind of solution is very attractive respect to the use of polymers in terms of tailoring structures and obtain desired features (e.g., energy levels, absorption spectra, charge-carrier mobility, purity), but it is limited by the low reproducibility of the results, as the interactions between the active layer are not easily predictable, moreover, despite the better solubility, the results even in these case are very poor, compared to Silicon based solar cells, because of the difficulty of organic molecules to absorb the whole visible spectrum. To our knowledge, Nierengarten et al. were the first to attempt this approach, binding the PCBM with π -conjugated systems, both vinylene- and acetylene like.²³

Conjugated polymers have been investigated extensively for their applications in optoelectronic devices, such as light emitting diodes, photovoltaic cells, OFET, sensors.²⁴ Their performances in optoelectronic devices is determined mainly by interchain interactions, rather than by intrinsic properties of the conjugated polymer.²⁵ In order to control and take advantage of their interchain behaviour, conjugated polymers or oligomers have been introduced in a rod-coil block copolymer architecture as the rod component.

Many efforts were made to enlarge as much as possible the absorption spectrum of molecules by using pigments.²⁶ Even in this case the pigments can be linked to fullerene or thiophene-based molecules thanks to organic synthesis, but the values obtained and the not so appealing possibilities to make an industrial scale up have limited this approach.

In order to act these applications, electron donating and electron accepting properties of the molecules are of crucial importance.

From a molecular point of view, the target molecule has to be high-conjugated in order to act the charge transfer in a long range between the electrodes. Moreover, depending on the application, a broad absorption spectra in the visible region could be required. Obviously, from a chemical point of view, a focus parameter is the solution processibility in order to do further syntheses.

From a structural point of view it's better to achieve further progress concerning the supra-molecular control of the active layer morphology, i.e., by hydrogen or halogen bonding, exploiding the rigid rod-like structure of monodisperse molecules.

Voltage-current (V-i) measurements are needed to describe the performances of a solar cell. **Fig.5** shows a tipical I-V plot under dark and light conditions. We can obtain many informations about the parameters describing a solar cell.²⁷⁻²⁸

- The open-circuit Voltage (V_{OC}).

It is the Voltage that has to be applied to make the current be equal to zero. It depends by the energy levels' offset of the used materials as well as their interfaces.

- The short circuit current (I_{SC}).

It is the current flowing when the Voltage is equal to zero. This parameter depends by the nanoscale morphology of the organic semiconductor thin film. In a van der Waals crystal, the final nanomorphology depends on film preparation, so experimental parameters as solvent type, the solvent evaporation (crystallization) time, the temperature of the substrate and the deposition method can change the nanomorphology. In BHJ there is an enhanced interfacial area between donor and acceptor phases, but this factor is counterbalanced by a complicated nanomorphology that is difficult to control.

- The maximum power point (P_{MAX}). It correspond to the maximum product between Voltage and current.

- The Fill Factor (FF).

It is the ratio between the P_{MAX} and the maximum Power that could be obtained (the product between I_{SC} and V_{OC}).

$$FF = \prod_{I_{SC}}^{D} \cdot \dots$$

The Fill Factor depends by the charge carriers reaching the electrodes, when the built in field is lowered toward the open circuit voltage. In fact, as just said, there is a competition between charge carrier recombination and transport. Hence, the

product between the lifetimes times the mobility determines the distance *d* that charge carriers can drift under a certain electric field. This product $\mu^*\tau$ has to be maximized.

- The Internal Quantum Efficiency (IQE).

It is the ratio between the P_{MAX} and the short circuit current I_{SC} .

$$IQE = \prod_{I_{SC}}^{D}$$

- The External Quantum Efficiency (EQE). It is the ratio between short circuit current I_{SC} and the forced current I_{F} .

$$EQE = I_{I_F}$$

- The efficiency of the solar cell as a ratio between the P_{MAX} and the incident solar power P_{INC} .

$$\eta = \bigcap_{P_{INC}}^{P}$$



Figure 5. I-V curve showing solar cell's main parameters.

The efficiency, however, it's the best way to compare solar cells between them because it indicates how many incident Energy is really converted in Electrical Energy.

In Organic Photovoltaic (OPV), the short diffusion length of donors (less than 10 nm), which is much shorter than the optical absorption length (100 nm),²⁹ has limited the EQE, and a large fraction of photogenerated charge remains unused for photocurrent generation. Therefore, it is crucial to control the of the morphology and of the mixing state of the Donor-Acceptor heterojunction.

Concluding, several approaches can be applied to improve the efficiency of organic solar cells. These can be reduced to the regulation of two factors:

a- Regulation of the band gap of organic molecules in order to improve electrical parameters as V_{oc} and I_{sc}

b- Regulation of the morphology of the films in order to minimize the losses by recombination of the wrong sign of charges at the wrong electrode.
In this sense, organic molecules are very versatile because the band gap can be regulated by "tailoring" the opportune synthesis.

The nanoscopic organization can be regulated by introducing

- An amphiphilic primary structure like di-block copolymer resulting in a self-organized phase;
- 2- An amine-acid- type hydrogen bonded self organization resulting in a hydrogen bonded polymeric superstructure
- 3- An inorganic (ZnO_x or TiO_x) template nanostructure filled with organic semiconductors
- 4- Liquid crystalline self-organizing columns of donor-acceptor phases.

This strategies require a multi-discipline knowledge in macromolecular chemistry, supramolecular chemistry, physical chemistry, colloid chemistry, photophysics/photochemistry, device physics, nanostructural analysis and thin films technology and, as consequence, great challenges and advancements in the entire field of chemical sciences.

In particular, if we consider a non-covalent approach, supramolecular polymers appear to be interesting because of the possibility of tunable structural properties arising from the dynamic linking between the constituent monomers.

If we consider chemical sensors, it's well known that the aggregation state of the receptor could influence its ability to convert the chemical information in an electrical signal. The same approach could be thought if we consider the organic materials to be used in photovoltaic cells. In case of polymers, we can approximate them as presenting the exciton with a one dimensional diffusional pathway. The excitons therefore undergo a one-dimensional random walk when the polymers are in solution. If the exciton can undergo vectorial transport in a given direction, then much higher amplification factors can be achieved. The situation is more complicated in polymer films. Close proximity of the neighboring polymer chains

facilitates interchain energy migration, and fluorescent polymers often exhibit more planar conformations in thin films and aggregates, which appears to promote exciton diffusion.³⁰ As a results, the research of new methods to aggregate fluorescent polymers into optimal organizations for energy transfer is a great chance for investigations.

II. AIM OF THE RESEARCH

The target to be reached with this project is the development of a new organic molecule acting as organic dopant and, in a second moment, developing a photovoltaic system at the solid state that can be a suitable alternative to the commercial solar cells.

These new devices are based on using thin or ultrathin layers of organic molecules (having a π conjugated electron system) acting as n- and p- dopants. These molecules have been proved to have good electronic properties in solution because of their Molecular Orbitals offset that make them suitable for Organic Photovoltaic applications.

Chemical features as the presence of an appropriate alkyl chain may influence the consequent intermolecular interactions, and thus can can be used even to improve the solubility of the molecule in organic solvent, its homogeneity and finally the cristallinity of the organic layer. In this way we expect to obtain better values of photoresponse respect to the systems used as models.



Figure 6 : Scheme of device's conception. The Silicon surface acts as light harvester. The donor and acceptor molecule act as molecular wires to anode and cathode thanks to Molecular Orbital's tuning, that allows a continue electron transfer between HOMO an LUMO orbitals.

Literature is rich of examples in which organic molecules act as light harvester both as a blend of electron-donor and electron-acceptor molecules layer and a dyad layer (in this case a single molecule has both the acceptor and the donor properties).

At the best of our knowledge, there are few studies regarding organic compounds interacting with Silicon, Stutzmann et al. tried to study the interaction between nanocrystalline Silicon or hydrogenated amorphous Silicon with organic molecules, where it has been observed a little photovoltaic effect.³¹

A photovoltaic system based on inorganic semiconductor could be promising for several reasons. In fact inorganic semiconductor have a much broader absorption spectrum than organic semiconductors, moreover they can be doped to a specific n-or p- type doping level, and (expecially in form of nanocrystals) they can provide a large interface area to the organic counterpart. system based on intrinsic Silicon acting as light harvester that can be implied in the electron transfer process to an acceptor molecule and an acceptor molecule to close the circuit.

The work consists in synthesizing a new molecule, recognizing its band offset and match it to the band offset of Silicon³² in order to improve the charge transfer, thus achieving higher values of photocurrent.

The new molecule that will be studied belongs to the family of PolyPhenyleneEthynylenes (PPEs), in which aryl groups are linked by alkine units and their derivatives. Because of the molecular properties, compounds belonging to this class turned to be suitable candidates for improving the performances of OPV solar cells respect to the poly-phenylenevinilenes and polythiophenes, so in the last years there have been a rise of interest respect to this conjugated polymers.³³

The electronic properties can be modified by introducing appropriate substituents in order to regulate the HOMO-LUMO levels and consequently the band gap between the valence band and the conductive band of the organic molecules.

For most conjugated polymers, when the conformation of a chain passes from a "twisted" into a relatively planar one, the π electron delocalization along the backbone would become easier, and hence the spectra would red shift to a certain extent. ³⁴ This result can be reached through the introduction of suitable

ramifications and the use of spacers allowing both π delocalization and steric rigidity.³⁵

In addition, the rigid acetylenic spacer helps to minimize neighboring aromatic-ring interactions in this series should result in a bathocromic shift in the UV-vis absorption spectra maxima with corresponding reduced bandgap energy when compared to polythiofene or polyalkylthiophene analogues.³⁶

The presence of triple bonds as spacers can even regulate the interactions between different polymeric chains. It has been found that the intrachain effect is extremely strong for PPEs, owing to the relatively low rotational barrier of the aryl alkine single bonds along the backbone, which is extimated at less than 1 Kcalmol⁻¹. On the other hand the backbone planarization will give rise to a closer interchain contact, that will give rise to a closer interchain contact, which in turn facilitates the formation of the aggregates in the most of circumstances.³⁷

Although these possible advantages, at the best of our knowledge, there isn't any example in literature attesting the effective use of PPEs polymers in working OPV devices.

In order to analyze the behaviour of the new molecule, as well as to develop the new photovoltaic device's architecture, the first step was testing a model device based on well known organic molecules acting as photovoltaic materials. At today, molecules belonging to the Fullerene family and to the *p*-thiofenes are the most used in this field of the research.

The most used donor molecules belong to the Poly-3-Exyl Thiofene (P3HT) or the poly-p-phenylene vinylene (PPV) family because of the possibility to form well-ordered structures at the solid state. These polymers allow us to achieve many advantages, in fact the chainlike structure leads to strong coupling of the electronic states to conformational excitations peculiar of the one dimensional systems, moreover the relatively weak interchain binding allows diffusion of dopants molecules into the structure (between chains) while the strong interchain C-C bonds keeps the integrity of the polymer.

Because of their good solubility, good processability and environmental stability we choose P3HT as reference molecule to test the behaviour of the new synthesized compound and the developing of the device having a new architecture.

As a model for exploring the properties of this device [6,6]-phenyl-C61- butyrric acid methyl esther (PCBM) will be used as n-type dopant because of its ability to accept up to six electron in its conduction band. ³⁸ In turn, poly-3-hexylthiofene (P3HT), one of the most representive conjugated polimer donor material will be used as p-dopant.³⁹

[6,6]-phenyl-C61- butiric acid methyl esther (PCBM), is a well known n-type molecule. It can be synthesized starting from fullerene C_{60} by using 1-phenyl-1-(3-(methoxycarbonyl)propyl)diazomethane⁴⁰, the phenyl butyric methyl esther moiety make it soluble in common organic solvents. PCBM maintains one of the most attractive features of C_{60} , which is its long electron diffusion lenght, that is longer than Perilene carboxy diimide derivatives. This property is due to the rapid intersystem crossing of the singlet excited state to the long lived triplet excited state, allowing ample time for its diffusion.⁴¹ The absorption spectrum of C_{60} is reported in **Fig. 7**.



Fig. 7. Absorption Spctra for PC61BM (green line) and several other molecules used for photovoltaic application

As we can see, the negligible absorption in the visible and infrared spectrum is attributed to a high degree of symmetry, making the lowest-energy transitions formally dipole forbidden. It has been proved that reducing the symmetry, by replacing PCBM with a less symmetrical or elliptical fullerene (as $PC_{71}BM$) these transiction become allowed and a dramatic increase in light harvesting is expected. Therefore, the PCE in many solar cell devices has been enhanced by using C_{70} derivatives as the acceptor material. Unfortunately, C_{70} derivatives are more difficult and expensive to separate, so C_{60} ones are still used.

This feature, mixed to the stability of blends with P3HT, the good alignment between HOMO and LUMO orbitals and the fact that at today PCBM is commercially available should encourage the use of these blends in commercial device; unfortunately PCBM high price and the natural tendence of fullerenes to photooxidation is a real limit to a large scale use. Moreover, lower-lying LUMO Energy level in PCBM reduces the V_{OC} , which is determined between the HOMO level of the donor material and LUMO level of the acceptor material.

III- RESULTS AND DISCUSSION.

III-1 SYNTHETIC ROUTE TO THE n-TYPE ORGANIC MOLECULE

In recent years there have been an increasing interest in synthesizing conjugated polymers and in studying their photoluminescence and electroluminescence properties.⁴²

These molecules can sum the processability of organic materials and the electronic and photophysical properties of semiconductors, so they are attractive candidates as active layers for electroluminescence devices.

Polymers belonging to the *p*-phenylenethinylene family are particularly attractive because of their flexible molecular orbitals energetic, known to be tunable *via* an appropriate skeletal functionalization. Moreover, the advaiability of efficient synthetic protocols⁴³ allows the effective π -coniugation length of these shape persistent rod-like structures to be easily varied by controlling the number of arylacetylenic repeating units.

From the structural point of view, the alkyne linkages are more accommodating than alkenes to steric and conformational constraints because the *quasi-cylindrical* electronic symmetry.⁴⁴

Moreover, the use of acetylenic spacers into a poly-aryl backbone offer us several distinct advantages. In fact triple bonds can act as rigid conjugative spacers linking two benzene repeating units along the 4,4' positions on the same plane.

Recently, *p*-acetylenes and their ring-substituted derivatives have been investigated in order to improve their processability, the long term stability, the charge transport properties, their crystallinity as well as the possibility to use them in a BHJ device after spin coating them on a suitable support.⁴⁵

It was found that alkoxy chains, grafted as side groups on the backbone of conjugated polymers, do not only enhance their solubility and subsequently their processability into thin films for various applications, but can also lead to dramatic changes in optical, electronic and transport properties as well the thermal behaviour

that conjugated polymers show in the solid state. This effect was demonstrated to depend by size, dimensions and positions of the alkoxy chain.⁴⁶

The chromic behaviour of PPEs depends by the environment surrounding the molecules.

It has been demonstrated that, when turning from dilute solution to solid films, the absorption and emission spectra of PPEs always have a large bathochromic shift relative to other conjugated polymers, such as polyfluorenes and poly(p-phenylene vinylene)s.⁴⁷

In previous studies, the bathochromic shifts of PPEs were attributed either to the aggregate or to the back bone planarization. The aggregate is an interchain behaviour, which involves intimate π - π interaction of two or more chromophores in the ground state by extending the delocalization of π electrons over those chromophores. It usually leads to bathochromic shifts or new peaks in the spectra. In contrast, the backbone planarization is an intrachain behaviour, which is associated with the conformational transformation of a single chain.

In past years, the use of rod-coil block copolimers as photoactive material in BHJ's has been interpreted as a possible way to overcome the drawbacks regarding the polymer-fullerene blends: a poorly controlled D/A domain size distribution and the inherent morphological instability. Rod-coil block copolymers are indeed well known to self assemble through microphase separation into highly ordered nanostructures that are thermodynamically stable and exhibit spatial periodicities on the 1-10 nm scale, that make them useful if we compare it to the diffusion length of an exciton.⁴⁸ In this work we propose a synthesis for a p-phenylethinylene bearing a calix[5]arene moiety. In literature⁴⁹ polymers having a well known receptor in their ramifications are known in the field of sensing, but there aren't many examples of their application in OPV.

This approach would allow us to keep the basic charge carrier properties of pphenylenevinylene because there is still a conjugated wire in which the photogenerated exciton could go across, but the calix[5]arene, acting as a receptor,

could improve the crystallinity and even the photochemical behaviour of the whole system.

The final molecule is a "block" polymer, so the synthesis of two blocks is here described.

The synthesis of **Monomer 1** (**Scheme 1**) required to obtain preliminarily the 5,11,17,23,29-pentat-butyl-31-monohydroxy-32,33,34,35-(4-methylpenthyloxy) calix[5]arene (**4**). The procedure was reported in literature⁵⁰ and required to start from the 5,11,17,23,29-penta-*t*-butyl-31,32,33,34,35-pentahydroxy calix[5]arene (**1**). One of the –OH groups was preliminary protected with a benzyl group (**2**); subsequently a S_N reaction with 1-bromo,4-methyl-pentane in order to introduce the alkyl chains in the lower rim have been carried out to the 5,11,17,23,29-penta*t*-butyl-31,32,33,34,35-(4-methylpenthyloxy)calix[5]arene (**3**). A final hydrolysis of the benzyl group allowed us to obtain the free -OH of (**4**).

Monomer 1 was synthesized starting from 1-4 diodo, 2-5 dihydroxy benzene. A S_N reaction using 1-Tosyl, 6-hydroxy hexane reported in literature allows us to introduce a 1-hydroxyhexyl substituent in 2-5 position (**5**). The OH group of the hexyl chain can be removed for simple chlorination using SOCl₂ (**6**). The final block can be obtained by reacting (**6**) with calix[5] arene having a free OH group in the lower rim.

Monomer 2 was synthesized using a procedure known in literature⁵¹ starting from 1-4 diodo, 2-5 dihydroxy benzene (**Scheme 2**). A S_N reaction using 1-bromo,4methylpentane allowed us to introduce a 4-methylpenthyl substituent in 2-5 position. The lodine atoms in benzene rings were used to introduce a trimethylethynilsilylgroup thanks to a Sonogashira coupling with ethyniltrimethylsilane. A simple hydrolysis step allowed us to remove the trimethylsilyl- groups and obtain a benzene derivative having free acetylenic hydrogens (**12**).

The final step (**Scheme 3**) consists in polymerizing the two blocks; it is based on a coupling between the Monomer 1, bearing the calix[5]arene moiety and the Monomer 2 having the acetylenic protons. The reaction is catalyzed by Cul and

Pd(PPh₃)4. The molecule we used as stopper for the polymerization is the iodobenzene; the time at which we added the stopper influenced the length of the final molecular wire. In a first case the stopper was added after 2 days and gave us a final mixture having a 20% of polymer. In a second time the stopper was added after two weeks of reaction between the two blocks. In this case we obtained a final mixture having 80% of polymer, that has been characterized by ¹HNMR and GPC. ¹H and ¹³C NMR spectra of **PC[5]** showed the complete absence of structural defects. In fact, the integration data related to the proton signals of the polymer shows that the two structural units derived from monomers **9** and **12** are present in a 1:1 ratio. GPC analysis allowed us to say that there are in average 16 repeating units for every polymeric chain.





Scheme 1: Representation of the synthetic route to Monomer 1 (9)







Scheme 2: Representation of the synthetic route to Monomer 2 (12)



Scheme 3: Synthesis of PC[5]

III-2 NMR EVIDENCE FOR POLYMER ASSEMBLY

As we stated in the chapter concerning the aim of this research, one of our target is to evaluate how the increase of the structural order of the polymeric system can influence the performance of the whole molecule in a photovoltaic device. For this reason we used the well known ability of the p-tertbutilcalix[5]arenes having a fixed cone-like arrangement to selectively form 1:1 inclusion complexes with linear alkyldiammonium ions⁵² and their ability to encapsulate long chain α,ω -alcanediyldiammonium ions in which every ditopic guest of appropriate length coordinates to a pair of calix[5]arene units.⁵³

Specifically, we intend to use the reversible incapsulation of 1,10decanediyldiammonium ion to form a network between two polymeric chains that could interact each other thanks to two calix[5]arenes complexing the ammonium group of a suitable salt.

After the verification of the stoichiometry of the complexation we intend to make a confront between the pristine polymer and its supramolecular adduct to verify if the complex keeps on existing even at the solid state and what is the supramolecular network's contribution in a diode simulating the final device.

We performed a ¹HNMR titration using the **PC[5]** as a Host and the 1,10decanediyldiammonium picrate salt as a Guest. This technique proved to be useful to study the association-dissociation of the supramolecular complex because both host and guest posses diagnostic probe signals undergoing (in slow exchange regime in the NMR scale) substantial and distinctive chemical shift changes upon self assembly as a function of the symmetry elements present within a given assembled specie.

As we can see from **Spectrum 1** and **Spectrum 2**, after the addiction of 0.1 eq of diammonium picrate salt, inclusion signals at negative values of ppm appear. These signals are due to the formation of a "capsule" between two calix[5]arenes belonging to two different polymeric wires and the diammonium ion.

When 0.5 equiv of 1,10- decanediyldiammonium are added to a 1 mM (per repeat unit for the polymer) CDCl₃/CD₃OH (4/1, v/v) solution of **PC[5]**, reaching the 2:1 host:guest ratio, the spectrum showed signals that were compatible to a polycapsular complex (from now on **PC[5]**_{net}), as demonstrated by the presence in the high field region (-2.0 to 0.3 ppm) of a single set of peaks for the α - to ϵ -CH₂ and the symmetry-related α^{1} - to ϵ^{1} -CH₂ groups of the guest , shielded by the π -rich cavities of the two molecules arranged in a polycapsular fashion.

Resonance peaks compatible with the presence of either free guest

 $(\alpha$ -CH₂ = α^1 - CH₂) or 1:2 bis-endo cavity (from now on **PC[5]**_{cap}) assembly are barely observed in the diagnostic δ =2.82 region. Upon further addition of 0.5 equiv of Guest to the pre-formed **PC[5]**_{net} assembly, a second, more spread set of signals for the included guest becomes evident in the high field region of the spectra (δ = -1.9 to 0.4 region), as a result of the 1:2 bis-endo-cavity assembly. This conclusion is also confirmed by the presence of two equally intense (AX signals(4.3-4.5 and 3.3-3.8 ppm respectively) relatively to the calix[5]arene methylenes and by the triplet detected at δ =2.82 ppm, which is assigned to the α^1 -CH₂ protons of the guest and it is judged a diagnostic evidence for the 1:2 Host/Guest complex formation.

Further addiction of 1.5 equiv of Guest to the above solution, so as reach the 1:2.5 Host/Guest ratio, mainly led to the formation of the latter architecture with the presence of the signals belonging to the α - to α^{1} -CH₂ groups of the alkyl chain in the -2.0 to 0.3 ppm high field region.

We can summarize the results of this experiment saying that complex formation induces symmetrization and desymmetrization of the guest. When the two methylene groups (α and α^1) are both cavity included, they are equivalent, and as result of the shielding of the aromatic con of the calixarenes, so resonate at high fields (δ =-1.30 ppm) as a broad singlet, whereas they appears as two distinct resounances at very different field strengths when only half of the dication is included in the calixarene moieties (i.e. **PC[5]**_{cap}). In this case the included α -CH₂

signal resonates at -1.27 ppm, whereas the α^1 -CH₂, not included, appears as a triplet at 2.82 ppm.

These peaks suggest us that in solution we have the formation of different ratio complexes between PC[5] and the dialkylammonium ion, as described in **Fig. 8 a** and **8 b**.



Fig 8 a: Model we use to explain the formation of the **PC[5]**_{net} network using the calixarenes (pink vessels) and the ammonium ions (blue circles).



Fig. 8 b: Model we use to explain the formation of a PC[5]_{cap} stoichometry complex between the polymer and the diammonium picrate ion.



Spectrum 1: ¹H-NMR signals obtained during the NMR titration a) **PC[5]**, b)H/G= 1:0,1, c) H/G= 1:0,2; d) H/G= 1:0.3; e) H/G= 1:0.4; f) H/G= 1:0.5; g) H/G= 1:1; h) H/G= 1:2 i) H/G= 1:2.5



Spectrum 2: Detail oof the ¹H-NMR signals obtained during the NMR titration a) **PC[5]**, b)H/G= 1:0,1, c) H/G= 1:0,2; d) H/G= 1:0.3; e) H/G= 1:0.4; f) H/G= 1:0.5; g) H/G= 1:1; h) H/G= 1:2 i) H/G= 1:2.5. The ¹H-NMR signals related to the Hidrogens of the diylammonium ions show the evolution of the complex stoichiometry during the NMR titration

III-3 UV-vis AND FLUORESCENCE SPECTRA OF THE NEW POLYMER

The properties of organic semiconductors not only depend by the molecular structure, but are also strongly influenced by supramolecular ordering and aggregation phenomena, both in solution and in the solid state. Since the PPEs show a propensity to form aggregates is solution of some solvents or solvent mixtures⁵⁴, with strong influence on the properties, the study of the assembly of polymeric chains is a mandatory step to tune the optoelectronic properties of the conductive polymer-based device.

The first step was evaluating the Uv-vis absorption spectra of the polymer, so a cuvette containing a solution of polymer PC[5] 5 x 10^{-6} M in a mixture CH₃Cl/ CH₃OH (4/1, v,v) was prepared and measured. (see **Fig. 9a**)



Fig. 9a : Absorption spectra of PC[5] (green line)and C10 diammonium picrate (blu line) salt in CH_3CI/CH_3OH (4/1 v/v)



Fig. 9b : Fluorescence spectra of PC[5] in CH₃Cl/ CH₃OH 4/1 mixture

Fluorescence measurements (Fig. **9b**) on the same solution showed in the range 400-700 nm only one detectable band at 480 nm, confirming us that this polymer belonging to the aryl acetylenes family is a blue-green emitter.

After that a Host-Guest titration using the polymer as host and the C_{10} diammonium picrate as guest has been carried using a CHCl₃/CH₃OH (4/1 v/v) as solvent was performed.

Fluorescence titration experiments were performed in order to determine the fluorescence behavior for the assemblies **PC[5]**_{net} (2:1 H/G stoichiometry) and **PC[5]**_{cap} (1:2 H/G stoichiometry) by addiction of different aliquots of C10 diammonium picrate salt (0.02-20.0 equiv). Interestingly, we noticed in the range of 0.02-0.5 equiv of C10 diammonium picrate salt added that the fluorescence emission increases respect with to the free Polymer and after that remains nearly constant.

On the other hand, upon further addiction of 0.5 equiv of C10 diammonium picrate salt, a progressive quenching of emission is observed. This behavior is related to the formation of the **PC[5]**_{cap} complex .

This trend confirms the formation of two different host-guest species. The initial increasing in the 0.02-0.5 equiv range of diammonium salt added suggests the formation of a more ordered specie with respect to the starting polymer, while the decrease of emission after the addiction of 0.5 equiv indicates the formation of a less ordered species. (See **Fig. 10**)



Figure 10: Normalized emissions of PC[5] ($5.0*10^{-6}$ M) with different equivalents of C102Pic (0, 0.02, 0.04, 0.06, 0.1, 0.2, 0.3, 0.4, 0.6, 0.7, 0.8 and 0.9) in CHCl₃/CH₃OH (4/1,v/v); λ_{exc} = 440 nm, λ_{em} = 480 nm.

Binding constants were determined following the emission variation at 480 nm, using HypSpec (v1.1.33), a software designed to extract equilibrium constant from potentiometric and/or spectrofotometric data. HypSpec starts with an assumed complex formation scheme and uses a least-squares approach to derive the spectra of the complexes and the stability constant. All emission curves were normalized to eliminate the dilution effect.
The constant value for the initial association was extimated to be $\log K_{net}$ = 9.88± 0.85.

The binding constant related to this specie was extimated to be log K_{cap} = 7.08± 0.15.

Non covalent polymers have the specific ability of varying their properties in response to an environment modification. This property makes them different respect with the covalently bound polymers. In this respect, the couple PC[5]/decanediyldiammonium ion is an extremely useful precursor for network formation because the 1,10-decanediyldiammonium promptly responds to external stimuli (e.g. acid/base treatment), activating and deactivating the self-assembly process.

In particular, the diammonium ions can be easily deprotonated to the free diaminic forms by treatment with a mild organic base such as Triethylamine (Et₃N). The amino derivative can be transformed in the diammonium salt upon successive treatment with a TriFluoroacetic Acid (TFA) solution.

In order to verify the possibility to go back and forth on the formation of the supramolecular complex we treated **PC[5]**_{cap} and **PC[5]**_{net} solutions with a Base (Tri-Ethylenamine, ET₃N) and an Acid (Tri-Fluoro Acetic Acid, TFA).



Fig 11 : Normalized change emission of **PC[5]**_{net} (squares) and PC[5]_{cap} (rcircles) during the cyclic addiction of 1 equiv of base (Et₃N) and acid (TFA) in CHCl₃/CH₃OH (4/1 v/v); λ_{ex} =440 nm, λ_{em} =480 nm. Black and red lines refer to **PC[5]**_{net} and **PC[5]**_{cap} respectively

We noticed that cyclic additions of Base and acid to the **PC[5]**_{net} solution don't modify significantly the intensity of the emission of the supramolecular network (**see Fig. 11**, black line)

If we compare the values of Fluorescence Intensity for the **PC[5]**_{net} and **PC[5]**_{cap} ratio complexes, we can notice that the larger differences in every Acid-Base treatment are observed in the **PC[5]**_{cap}. Moreover, only in **PC[5]**_{cap} we noted a real back and forth behavior after treating with acid and base This experiment proved that the supramolecular network of **PC[5]**_{net} is a well founded construction, in which the guest molecules are strongly embedded in the calix[5]arene cavities and the basic conditions cannot provide the network's disassembly.

III- 4 ELECTROCHEMICAL MEASUREMENTS

The electrochemical properties of the **PC[5]** were determined by Cyclic Voltammetry (CV).

The CV was performed with a solution of tetrabutylammonium perchlorate in acetonitrile at scan rates of 50 and 20 mV/s at room temperature under a Nitrogen flux. A platinum electrode (0.05 mm²) covered with a polymeric film was used as working electrode. Pt was even used as counter electrode and working electrode.

From cyclic voltammetry characterization we can estimate the HOMO and LUMO Energy values and, as a consequence, the Band gap of the molecule. The LUMO Energy value can be estimated from the cathodic onset potential, obviously the HOMO Energy can be obtained by measuring the anodic onset potential.

HOMO and LUMO Energy level were calculated with respect to the appropriate ferrocene/ferrocenium oxidation or reduction. The HOMO and LUMO Energy levels were calculated thanks to an empirical formula:

HOMO: -e(4.8 V+ E_{ons-an})

LUMO: -e(4.8 V+ E_{ons-cat})

Where 4.8 V is the potential of the referral ferrocene/ferrocenium respect to the Standard Hydrogen Electrode (SHE) in vacuum, E_{ons-an} is the onset anodic potential value and $E_{ons-cat}$ is the onset catodic potential value. This value was corrected respect to the Ferrocenium/Ferrocene redox couple used as standard in the solvent that was used for the measurement (CH₂Cl₂) according to the literature.⁵⁶ Mean values obtained for HOMO and LUMO levels are -5,85 eV and --3,98 eV respectively. These values suggest a potential use as PCBM sostitute (n-dopant)

III-5 DEVELOPING THE SIMULATING DEVICE

After studying the polymer's behaviour in solution, our purpose was to perform measurements on simulating diodes in order to confirm the possibility of developing a device based on an interface between Silicon and organic molecules.

HOMO-LUMO band alignment (See Fig.12) suggests us a mechanism in which:

- When light interacts with an interface p-Si/n-type molecule (as PCBM or our polymer) Si absorbs light, holes and electrons separate and electrons can be transferred to the organic molecule.
- When light interacts with an interface n-Si/p-type molecule (as P3HT), the organic molecule absorbs light, after that electrons go to Silicon.

Thus, the global behaviour can be schematized in the **Fig.13**, in which Silicon "pumps" electrons from its conducting band to the LUMO orbital of the n-type molecule, while holes go from Silicon's Valence Band to the HOMO of the p-type molecule.





Fig.12 : Band Gap alignment for a) p-Si/PCBM, b) p-Si/Polymer, c) n-Si/P3HT. Color Code. Red line: Conducting Band (or LUMO orbital), Blue line: Valence Band (or HOMO orbital), Dotted line: Fermi Level. Note: 1) Alignment has been made respect to the Fermi Level. 2) For organic molecules Fermi Level is considered lying in the middle between HOMO and LUMO Levels



Fig. 13: Schematic representation of the final device. Colour code: black square: n-.type molecule (PCBM or polymer), blue square: Silicon, red square: n-type molecule (P3HT), green circle: electron, white circle, hole.

In this sense, electrical measurements performed on a solid state device could are expected to offer a wider knowledge on the possible behaviour's variation of the organic molecule when it passes from the solution to solid.

Based on the voltammetry measurements executed on our molecule and the well known behaviour of the PCBM, P3HT as n-type and p-type dopants respectively, we dealt with the preparation of interfaces Silicon-organic molecule.

In order to simplify electrical measurements, we used Silicon wafers retro-implanted with Boron (for the p-type Silicon) and Phosphorus (for the n-type Silicon) (from ST-Microelectronics).

We used spin coating as procedure to modify Silicon's surfaces.

This technique is based on depositing an excess of solution on the desired substrate. Because of the centrifugal force, due to the rotation of the sample, the solution covers the entire surface, while the excess is pushed out of the sample.

This method allows to deposit layers in the order of nanometers.

Tecnically, spin-coating is divided in four step (Fig.14):

- a) Solution deposition on surface
- b) Spin rate raising and solution distribution
- c) Constant rate spinning of substrate. The layer thickness is controlled by solution viscosity
- d) Substrate keeps on spinning and film thickness is controlled by the evaporation rate of the solvent

This approach was repeated for different spinning rates (500, 1000, 1500 rpm) to obtain different layer thicknesses



Fig. 14: Representation of the spin-coatin process

In order to study the interaction between pSi/n-type molecule, p-doped silicon surfaces were spin-coated with PCBM thin films; for this reason solutions of PCBM in choloroform having different concentrations (1.5; 3; 6; 12 mg/ml) were prepared

to obtain layers having different thickness. Solutions were stirred for 4 h to ensure the complete solubilization of PCBM before starting the spin-coating process.

Before the deposition, p-doped silicon is etched with a 5% HF solution, to eliminate the native oxide on silicon surface. After that, surfaces were rinsed with MilliQ water to eliminate the eventual HF in excess and then dried with a N_2 flux to avoid the instant silicon oxide formation.

The same procedure, obviously performed using n-doped Silicon wafers as substrate and solutions of P3HT having different concentrations, was repeated to prepare diodes based on the interaction between n-Si/p-type molecule.

After performing the measurements on the samples modified with organic molecules acting as referring, we executed the same procedure using chloroform solutions the pristine polymer **PC[5]**, the **PC[5]**_{net} (branched with a 2:1 stoichiometry) and the **PC[5]**_{cap} (branched with a 1:2 stoichiometry).

III- 5a AFM MEASUREMENTS ON PCBM AND POLYMER MODIFIED SURFACES

Atomic Force Microscopy (AFM) analysis have been carried out using commercial probes for Silicon tapping (Digital) having crown tips and a radius of curvature of 10 nm at the apex. During the analysis, the "cantilever" (length: 125 μ m, elastic constant 20-100 N/m) fluctuated at its resonance frequency (circa 300 KHz). Images have been obtained with 1-400 μ m² scans and collecting 512x512 points for scan at a rate of 0.5-2 Hz.

AFM analysis of deposited ultra-thin films of PCBM (6,7,10 nm) show extremely flat films without any morphological variation on varying the thickness. (**Fig. 15**) This technique demonstrates that the spin coating is useful to obtain ultra-thin and

highly homogeneous films of organic molecules, even when they are single molecules instead of polymers.



PCBM su Si 500 rpm pcbm_si_500rpm.002

15 a



PCBM su Si 1000 rpm pcbm_si_1000rpm.002







Fig. 15 : AFM images (10µm x 10µm) of a) 10 nm PCBM layer (deposited at 500 rpm) b) 7 nm PCBM layer (deposited at 1000 rpm), c) 6 nm PCBM layer (deposited at 1500 rpm).

Because of the high roughness of the samples, the measurement of the layer thickness for spin coated samples of polymer, PC[5]_{net} and PC[5]_{cap} was not possible; however for the AFM analysis we used samples prepared by drop-casting to the morphologies of the films obtained for the polymer and the polymer/picrate complexes at different ratios.

The results were correlated to the information we obtained from the in-solution techniques (NMR, UV-vis, Fluorescence) about the interactions between the different polymeric wires in absence and in presence of different amounts of C10 diammonium picrate spacer.

AFM analysis gave us the following images (see **Fig. 16**), from which we can deduce that:

- a solution containing the polymer without any spacer (Fig. 16 a), gives isolated aggregates with size ranging from few nanometers to few hundreds of nanometers.
- a solution containing PC[5]_{net} (Fig. 16 b), the Host-guest interaction between the calix[5]arene moiety and the ammonium ion leads to a nanostructure in solution. These interactions lead to the formation of a homogeneous layer at the solid state having almost uniform thickness (4.68 ± 018 nm) as measured from the cross section. However, we can recognize the presence of isolated molecules of polymer, but this is not the dominant system at the solid state.
- a solution containing PC[5]_{cap} (Fig. 16 c), AFM analysis shows a less homogeneous coverage of the surface, in which we can find both microscopic aggregates and a few residual network fragments as suggested by the steps in the cross section. This image confirms the data obtained from the H-NMR titration in excess of titrating agent (the ammonium picrate salt) telling us that the interaction between different polymeric chains based on the capsule formation between calix[5]arene moiety and C10 diammonium picrate is not completely broken.



16 b



16 c

Fig.16: AFM images ($5\mu m \times 5\mu m$) of a) spin coated PC[5], b)PC[5]_{net}, c)PC[5]_{cap}

III- 5b XRR MEASUREMENTS ON ORGANIC LAYERS

XRR reflectivity curves give us information about the layer density, thickness and roughness. In particular, here the experimental XRR curves for each deposition condition are reported and the curve fitting. Curve fitting is based on iterative of even more refined values of density, thickness and roughness of the deposited layer. In particular, it has been observed that density is independent of deposition conditions and assumes values of 1.5 g/cm³. Film thickness instead, depends on the deposition conditions, as can be seen by the reduction in Kiessling fringes' spacing, allowing to deduce a raising of thickness, depending on concentration and spin rate. XRR reflectivity values for the various films are reported in the **Table 1**

PCBM Solutions		
Concentration	Revolutions per	Thickness
	minute	
1.65E-3 M	1500	6.1±0.5 nm
1.65E-3 M	1000	7.3±0.5 nm
1.65E-3 M	500	10.2±1 nm
3.29E-3 M	1500	15.3±1.5 nm
3.29E-3 M	1000	18.0±2 nm
3.29E-3 M	500	23.2±2.5 nm
6.59E-3 M	1500	28.5±2.5 nm
6.59E-3 M	1000	32.7±4 nm
6.59E-3 M	500	46.2±5 nm

1.32E-2 M	1500	61.8±5 nm	
1.32E-2 M	500	98.0±7 nm	
P3HT Solutions			
1.5E-3 M	1500	15.0±0.1 nm	
1.5E-3 M	1000	18.0±0.5 nm	
1.5E-3 M	500	25.0±0.1 nm	

Table 1: Different thickness values of PCBM and P3HT films

III 6- Electrical measurements

Electrical measurements were performed at Electrical Measurement laboratory of the IMM-CNR of Catania. Because of the difficulty in depositing a metal contact on the organic layers by using the sputter deposition technique, measurements have been performed using an Au tip as front contact and the retro-implanted side of the Silicon layer as back contact as depicted in **Fig. 17**.

The rudimental preparation of the sample, doesn't provide quantitative values to talk about as well for I/V measurements as for the Sheet Resistance values that have been obtained using the "four probe measurement" approach, but a remark about the qualitative behaviour of the samples analyzed can surely be done.



Fig.17 : Scheme of the simulating diode contacts

I-V characteristics of Si/PCBM, Si/P3HT, Si/Polymer, Si/PC[5]_{net} and Si/PC[5]_{cap} diodes have been measured in order to evaluate the diode behaviour on varying the interface between silicon and the different kind of organic molecule, but also to

study the diode's behaviour on varying the thickness and the structure of the deposited organic layer.

III 6a – Sheet Resistance measurements of PCBM and different polymer layers

Four probe measurements (see **Fig. 18** for the scheme) have been performed by scanning samples in a range of current between 10^{-12} A and 10^{-3} A; values in which the behavior of the samples was linear have been considered.



Fig.18 : Four probes measurement's scheme

Measurements on different layers where obtained spin coating different solutions of of PCBM Silicon having native oxide (3 nm) and subjected to HF (5% solution), thus obtaining V/I values reported in **Fig 19**. We can see that while decreasing the thickness of the organic layer we have a decrease of resistivity. This can be explained considering that when charges separate at the interfaces, they can go across multiple paths until they are collected by the electrodes. The thinner is the layer, the lower thew number of multiple paths, thus decreasing the resistance of the samples.



Fig.19 : V/I values of different PCBM layers

PC[5], PC[5]_{net} and PC[5]_{cap} solutions were deposited on a thermal grown SiO₂ substrate (300 nm) supplied by ST Microelectronics. R_s show an insulating behavior of the polymer when interfaced with the oxide. (**Fig 20**)



Fig.20 : Sheet resistance values for different layers of PC, PC[5]_{net} (1:1) and PC[5]_{cap} measured on a 300 nm layer of thermal grown Silicon

III 6b – I/V measurements on different layers of PCBM and P3HT

Because of the interesting behavior of the ultra thin layer in terms of sheet resistance, we performed an I/V measurement to study the interaction p-Silicon/PCBM on 6 nm PCBM layer obtained by spin coating a solution of PCBM in CHCl₃ having 1.5 mg/ml at 1500 rpm

In the **Fig.22** we can notice a rectifying behaviour of the diode showing that the exciton can be formed at the interface between p-Silicon and PCBM. Higher values in terms of I_{sc} can be noticed respect to the control of pristine silicon, but a negligible Voc can be observed



Fig.21 : I/V characteristic for the 6 nm PCBM layer interfaced with p-Silicon.

Since we found of interest the behaviour of the lowest organic layer interface with Silicon, we repeated the experience using a 15 nm (deposited at 1500 rpm) layer of P3HT interfaced with n-Silicon. Even in this case we found a clear photovoltaic effect, with slightly higher values of I_{sc} for the organic layer respect to the Silicon Control, but a negligible V_{oc} can be seen (**Fig. 22**).



Fig.22 : I/V characteristic for the 15 nm P3HT layer interfaced with n-Silicon.

III 6c – I/V measurements on different layers of Polymer, PC[5]_{net} and PC[5] _{cap} deposited on n-Silicon and p-Silicon

After controlling the behaviour of the simulating diode formed using PCBM and P3HT interacting with p- and n- Silicon respectively, the experience was repeated using the polymer in its pristine form and its "complexed" forms **PC[5]**_{net} and

PC[5] cap.

Solutions of 1.5 x 10^{-5} M containing the polymer, **PC[5]**_{net} (2:1 complex) and **PC[5]**_{cap} (1:2 complex) were spin coated at different spin rates (1500, 1000 and 500 rpm respectively) on both n-Si and p-Si substrates. After that the behaviour of the diode on dark and light was evaluated.

When considering the different organic layers interacting with n-Si (see Fig. 23), we noticed under dark conditions an insulating behaviour of both polymer and $PC[5]_{net}$ and $PC[5]_{cap}$. The I_{sc} was in the same order of magnitude of the Silicon used as referral.

Measurements performed under light underlined the insulating behaviour of the polymer and its complexes, thus confirming the n-behaviour of the polymer we deduced after the voltammetry measurements performed in solution.

While the measurements performed on diodes obtained spinning the pristine polymer at different spinning rates (red lines) show similar values despite the layer's thickness, the same thing cannot be deduced for the different complexes.

PC[5]_{net} (2:1 stoichiometry complex) spin coated at 500 rpm shows lower values of Isc respect to the other samples deposited using 1000 and 1500 rpm, having a similar behaviour.

This trend can be noticed both under dark and light conditions.

Complex **PC[5]** _{cap} deposited at 500 rpm seems to conduct more current respect to the **PC[5]**_{net} (2 orders of magnitude higher). Moreover, even in this case we can

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recognize a behaviour depending by the thickness of the layer; in fact the organic layer deposited at 500 rpm seems less conductive respect to the ones deposited at 1000 at 1500 rpm respectively.



23 a



23 b

Fig.23 : I/V showing the behavior of different layers of **PC[5]** (red lines), **PC[5]**_{net}(green lines) and **PC[5]**_{cap} (purple lines) deposited on n-type Silicon. a) Dark, b) Light

A clear difference can be noticed when considering the different organic layers interacting with p-Si (see **Fig. 24**). A rectifying behaviour under both dark and light conditions can be noticed for all kind of samples Under dark all samples seem more conductive than the Silicon used as referral, expecially the different layers of 2:1 complex (purple lines) showing similar behaviour and an Isc 3 orders of magnitude higher than the control of Silicon.

Under illumination the rectifying behaviour is confirmed for all samples . Even if the **PC[5]**_{cap} complex samples show the same behaviour at different thickness, the same thing cannot be said for the other samples. The pristine polymer (red lines) shows lower lsc values for the sample deposited at 500 rpm respect to the 1000 and 1500 deposited ones that seems more similar.

Samples obtained using $PC[5]_{net}$ (green lines) show higher values of V_{oc} respect to the Silicon used as reference, moreover the I_{sc} measured under reverse voltage seems to depend by the thickness because we can notice lower values for the sample spinned at 500 rpm; this value increases passing to the samples deposited at 1000 and 1500 rpm respectively.

If we consider the Voc values, we can see that samples having **PC[5]**_{net} have the higher values, expecially the ones deposited at 1000 and 500 rpm

From these measurements we can deduce that the best performances are shown by the **PC[5]**_{cap}.

The higher lsc values respect to the Silicon referring confirm us that the **PC[5]** and its adducts have a suitable band offset for photovoltaic applications, but a quantitative result can be obtained only having a good metal contact.

The low Voc values for all samples obtained with the new organic molecule and its adducts suggests us the possibility of an improvement in terms of Si/material interface because the interface limits the V_{oc} .







24 b

Fig.24 : I/V showing the behavior of different layers of **PC[5]** (red lines), **PC[5]**_{net}(green lines) and **PC[5]**_{cap} (purple lines) deposited on p-type Silicon. a) Dark, b) Light

IV. EXPERIMENTAL

Synthesis of 5,11,17,23,29-pentat-butyl-31-benzyloxy-32,33,34,35-tetra-hydroxy-calix[5]arene (2)

A solution of 5,11,17,23,29-penta-*t*-butyl-31,32,33,34,35-pentahydroxy-calix[5]arene (5.0000 g, 0.0062 mol) has been mixed with KHCO₃ (1.2336 g, 0.0123 mol) and Benzylchloride (0.2625 g, 0.0021mol) in CH₃CN (50 ml) and then allowed to stir at reflux for 24 h.

The reaction was controlled with TLC (1:1). The product has been purified using a chromatographic column (SiO₂, Hexane/Toluene 1:2, v/v as eluent). The fraction corresponding to the product (lower Rf) was collected and the solvent removed, obtaining a white solid.

¹H-NMR(CDCl₃): δ (ppm) 7.70 (d, J=6.8 Hz, 2H, ArH), 7.50 (t, J=7.8 Hz, 2H), 7.43 (t, J=7.8 Hz, 1H), 7.19 (s, 4H), 7.17(q, J=2.5 Hz, 4H) 7.14(s, 2H), 5.29 (s, ArOCH₂), 4.41-4.10 (d, J=13.5 Hz, 4H), 3.45 and 3.42(AX system, J=13.5 Hz, 6H), 1.28-1.23-1.09 (s,2:2:1,45H)



Synthesis of 5,11,17,23,29-penta*t*-butyl-31-benzyloxy-32,33,34,35-tetra(4-methylpentyloxy) calix[5]arene (3)

A suspension of the product (**2**) (1.3075g, 0.00144.mol), 1-bromo-4-methyl pentane (2.857 g, 0.0173 mol) and K_2CO_3 (2.4039 g, 0.0173 mol) in CH₃CN (20 ml) has been stirred at reflux for one day.

The reaction was controlled by using TLC Hexane/AcEt 20/1. The pale white solution was rotovaporated. As the solvent was removed, the residue was extracted using CH_2Cl_2 . The organic phase was washed with a solution of HCl 0.1M, then it was dried on Na_2SO_4 .. (80% yeald)

¹H-NMR(CDCl₃): δ (ppm) 7.58 (d, J=6.8 Hz, 2H, ArH), 7.37 (t, J=7.8 Hz, 2H), 7.10(s, 2H), 7.20-7.04 (ABq, J=2.5 Hz, 4H), 6.72-6.54 (ABq, J=2.5 Hz, 4H), 5.29 (s, ArOCH₂), 4.57-3.22 (AX, J=14.0, ArCH₂Ar, 4H), 4.56-3.31 (AX, J=14.0, ArCH₂Ar, 2H), 4.51-3.25 (AX, J=14.5, ArCH₂Ar, 4H), 3.66 and 3.55 (m, 4H), 1.88-1.72 (m, 8H), 1.67-1.56 (m,8H), 1.30-1.24 (m, 4H) 1.20,1.14,0.85 (s, 2:1:2, 45 H), 0.95,0.94, 0.86 (d, 2:1:1, J=.6.5Hz, 24 H),



Synthesis of 5,11,17,23,29-penta*t*-butyl-31-monohydroxy-32,33,34,35-(4methylpentyloxy) calix[5]arene (4)

A suspension of the product (3) (1.3057g, 0.00145 mol) and 5% Pd/C (0.1640 g) in AcOEt (120 ml) has been stirred in H₂ atmosphere for 2 days.

The reaction was controlled by using TLC Hexane/AcEt 20/1. The reaction was filtered in a bed of celite and then the filtrate was evaporated to dryness to give a solid. Recrystallization from CH_3OH/CH_2Cl_2 gave the product (**4**).

¹H-NMR (CDCl₃) δ (ppm): 7.17 (s, ArH, 2H), 7.09-7.07 (ABq, J=2.5 Hz, 4H), 6.73-6.58 (ABq, J=2.5 Hz, 4H), 4.53-3.31 (AX, J=14.0, ArCH₂Ar, 4H), 4.49-3.26 (AX, J=14.0, ArCH₂Ar, 2H), 4.39-3.38 (AX, J=14.5, ArCH₂Ar, 4H), 3.92 and 3.81 (m, 4H), 3.66 and 3.45 (m, 4H), 1.99-1.86 (m, 8H), 1.69-1.60 (m,8H), 1.31-1.22 (m, 4H), 1.34,1.18,0.78 (s, 2:1:1, 45 H), 0.99,0.98, 0.95 (d, 2:1:1, J=.6.5Hz, 24 H),



Synthesis of 1,4-bis (4-methylpentyloxy)-2,5-diiodobenzene (10)

A solution of 2,5-diiodohydroquinone (0.4708 g, 0.0012 mol) and anhydrous K_2CO_3 (0.5613 g, 0.004 mol) in dry CH₃CN (15 ml) was stirred under a N₂ atmosphere. After 20 minutes 1-bromo-4-methyl pentane (0.4105 g, 0.002 mol) was added dropwise, then the solution was heated to reflux for 36 h.

As the solvent was removed, the residue was extracted using CH_2CI_2 . The organic phase was washed with a solution of HCI 0.1N, then it was dried on Na_2SO_4 . The crude product has been purified using a chromatographic column (SiO₂, Hexane/EtOAc 15:1 v/v as eluent) to obtain the desired product as a white solid after precipitation from CH_2CI_2 with CH_3OH .

¹HNMR (500 MHz, CDCl₃): 7.16 (s, 2H, ArH), 3.91 (t, J=7.0 Hz, 4H, -O*CH2*CH₂), 1.82-1.79 (m, 4H) 1.64-1.61 (m, 2H), 1.40-1.35 (m, 4H), 0.95-0.92 (m, 12H) ¹³CNMR-APT (125.7 MHz, CHCl₃): δ 152.9, 122.8, 86.3, 70.7, 35.1, 27.7, 27.0, 22.6





Synthesis of 1,4-bis[(Trimethylsilyl)ethynyl]-2,5-bis-(4methylpentyloxy)

benzene (11)

In a round bottomed flask compound (**10**) (0.395 g, $7.45*10^{-4}$ mol), trimethylsilylacetylene (0.146 g, $1.49*10^{-3}$ mol) Pd(PPh₃)Cl₂ (0.0264 g, $3.7*10^{-5}$ mol), Cul (0.0008 g, $0.42*10^{-5}$ mol) were mixed in 15 ml of diisopropylamine and allowed to stir at reflux in Nitrogen atmosphere for 16 h.

After cooling, toluene (10 ml) was added and the ammonium iodide (white precipitate) was filtered off. The solution was passed through a plug of silica gel using Toluene as solvent. The evaporation of solvent led to a yellow oil which crystallized upon standing. Recrystallization from hexane yielded to white needles (0.3240 g, 92%).

¹HNMR (500 MHz, CDCl₃): δ = 6.89 (s, 2H), 3.93 (t, J=6.0 Hz, 4H), 1.82-1.76 (m, 4H), 1.64-1.57 (m, 2H), 1.41-1.36 (m, 4H, -OCH₂CH₂CH₂CH-), 0.92 (d, J= 7.0 Hz, 12H), 0.25 (s, 18H) ppm

¹³CNMR-APT(125.7 MHz, CDCl₃): δ= 154.0, 117.2, 113.9, 101.0, 100.0, 69.7, 35.2, 27.8, 27.2, 22.6, -0.02 ppm

Elemental analysis calcd (%) for $C_{28}H_{46}O_2Si_2$: C 71.43, H 9.85; found: C 71.16, H 9.88.




Synthesis of 1,4-Diethynil-2,5 bis(4-methylpenthyloxy) benzene (12)

Compound (**11**) (200.0 mg, 0.424 mmol) was dissolved in THF (10 ml), after that CH_3OH (5 ml) and aqueous 20% KOH solution (300µl) were added. The mixture was allowed to stir for 4 h, then purified by removing the solvent. The yellow solid was purified for recrystallization from Hexane with charcoal, giving pale yellow crystals (111 mg, 80%).

¹HNMR (500 MHz, CDCl₃): δ= 6.95 (s, 2H) 3.95 (t, J= 6.5 Hz, 4H), 3.33 (s, 2H), 1.83-1.77 (m, 4H), 1.64-1.57 (m, 2H), 1.37-1.32 (m, 4H), 0.92 (d, J= 6.5 Hz, 12H) ppm.

¹³CNMR-APT (125.7 MHz, CDCl₃): δ=154.1, 117.8, 113.3, 82.4, 82.3, 70.0, 35.1, 27.7, 27.0, 22.5 ppm.

Elemental analysis calcd (%) for C₂₂H₃₀O₂: C 80.94, H 9.26; found: C 81.16, H 9.84.





Synthesis of 1-4Bis[6-chlorohexyloxy]-2,5-Diiodobenzene(8)

A solution of **7** (536 mg, 0.953 mmol) in 15 mL of CH_2Cl_2 was stirred in a ice bath for 20 min. $SOCl_2$ (454 mg, 3.813 mmol) and Et_3N (386 mg, 3.813 mmol) were added and the mixture was allowed to stir for 3 h. The organic phase was washed with H_2O and then dried over anhydrous Na_2SO_4 . The solvent was evaporated at reduced pressure and the crude product was purified by CC (eluent: hexane/CH₂Cl₂ 3:1) to afford the desired product that was precipitated in hexane to give 350 mg of **8** (61% yield).

¹H-NMR (500 MHz, CDCl₃): δ= 7.17(s, 2H), 3.94(t, J= 6.5Hz, 4H), 3.55(t, J= 6.5 Hz,

4H), 1.86-1.79(m, 8H), 1.57-1.50(m, 8H) ppm

¹³C-NMR (125.7 MHz, CDCl₃): δ= 152.8, 122.8, 86.3, 70.1, 45.0,32.5, 29.0, 26.5, 25.4 ppm

Elemental analysis calcd (%) for $C_{18}H_{26}I_2CI_2O_2$: C 36.09, H 4.37; found: C 35.88, H 4.41.





Synthesis of bis calix[5]arene (9)

In a round- bottomed flask the calix[5]arene (**4**) (0.2457 g, $2.15*10^{-4}$ mol) was mixed with anhydrous K₂CO₃ (0.300 g, $2.15*10^{-3}$ mol) and 1-4Bis [6-chlorohexyloxy] 2,5-Diiodobenzene (**8**) (0.0600 g, $9.77*10^{-5}$ mol).

The reagents were dissolved in Acetonitrile (20 ml). The mixture was allowed to stir at reflux in Nitrogen atmosphere.

After 48 h the reaction was stopped and the mixture was separated using a CC (Eluent: Hexane/CH₂Cl₂ 3:1, v/v). The fraction corresponding to the lower Rf is the expected product, so after collecting the fraction, the solvent was removed and finally the product (**9**) was obtained by precipitation from CH_2Cl_2 using CH_3OH .

¹H-NMR(CDCl₃): δ (ppm) 7.17(s, 2H), 7.04,7.02, 7.01 (pseudo-s, 1:2:2, 20H, ArH), 3.94(t, J=6.0 Hz, 4H), 3.66-3.55 (m, 20H), 4.54 and 3.25(AX system, J=13.5 Hz, 20H), 1.93-1.81 (m, 24H), 1.66-1.58 (m, 8H), 1.28-1.36 (m, 24H), 1.05-1.03-1.01 (pseudo-s,1:2:2, 90H), 0.95 (d, J=7.0 Hz, 48H) ppm.

¹³C NMR (125.7 MHz, CDCl₃): δ = 152.9, 152.7, 151.8, 145.5, 144.8, 144.5, 134.0, 133.9, 133.8, 133.4, 132.8, 126.8, 126.0, 125.6, 125.4, 125.3, 124.8, 86.3, 75.0, 74.2, 74.0, 73.7, 35.23, 35.19, 34.0, 33.8, 31.7, 31.4, 31.0, 30.9, 30.4, 29.7, 29.4, 29.3, 28.4, 28.3, 28.0, 26.5, 26.0, 22.86, 22.83, 22.7 ppm

MALDI –TOF (m/z): 2821 (MH⁺), 2843 (MNa⁺)

Elemental analysis calculated (%) for $C_{176}H_{260}I_2O_{16}$: C 74.91; H 9.29, found, C 74.36; H 9.48







Synthesis of polymer PC[5]

In a round bottomed flask the bis calix[5]arene (**9**) (80.0 mg, 0.0283 mmol) was mixed with the 1,4-diethynil-2,5-diodo benzene (10.2 mg, 0.0312 mmol), $Pd(PPh_3)_4$ (2.1 mg, 0.0018 mmol) and Cul (0.22 mg, 0.00116 mmol). A mixture of Toluene (7 ml) and Diisopropylamine (3 ml) was used as solvent, and the mixture was allowed to stir at 70 °C for 10 days.

Ammonium iodide salts were formed immediately after the start of the reaction and the mixture became highly fluorescent. Iodobenzene (6.7 mg, 0.034 mmol) was added and the mixture was allowed to stir again at reflux for four days.

After a total reaction time of 14 days, the reaction mixture was cooled to room temperature and added dropwise to rapidly stirred methanol (30 ml). After stirring for 2 h, the precipitate was collected and washed with hot methanol and hot acetonitrile. After drying overnight at room temperature, PC[5] was purified by means of a short column of silica gel using $CH_2Cl_2/MeOH$ 6:1 to afford a dark red solid (56 mg, 69%).

¹H-NMR (500 MHz, CDCl₃) δ: 7.17 (s,2H), 6.96-6.91 (m, 22H), 3.98-3.92 (m, 4H), 3.67-3.58 (m, 24H), 4.55 and 3.25 (AX system, J=14.0 Hz, 20H), 1.92-1.84 (m, 28 H),1.65-1.59 (m, 10 H) 1.36-1.30 (m, 28 H), 1.05,1.04,1.03 (s, 2:1:2, 90 H) 0.94 (d, J=6.5 Hz, 60 H) ppm

¹³CNMR-APT (125.7 CDCl₃): δ= 152.6, 144.9, 133.4, 133.26, 133.17, 125.7, 125.6,122.5, 116.4, 86.2, 74.5, 35.1, 34.6, 33.4, 30.2, 30.1, 29.7, 29.3, 29.2, 28.6, 28.2, 28.1, 27.7, 27.4, 26.9, 26.3, 25.9, 22.8, 22.4 ppm

GPC (THF, 35° , polystyrene standards as calibrant): M_w =61800, M_n =46100, PDI=1.34 (degree of polymerization [DP]~16)

Elemental analysis calcd (%) for C₁₉₇H₂₈₄O₁₄: C 82.26, H 9.95; found: C 81.93, H 10.26.







NMR Titration polymer/diammonium picrate

A solution 1.17 x 10^{-3} M in CDCl₃ containing the Host (the polymer) has been prepared. Progressive addictions of a solution 3.54 x 10^{-3} M of diammonium picrate (guest) have been executed, obtaining the Host/Guest ratios: 0.1, 0.2, 0.3, 0.40, 2.00, 2.5. The NMR spectrum have been memorized after every addiction

Entry	[H]/[G]	V _{host} (µI)	V _{guest} (µI)	V _{снсіз} (µI)	[Host]	[Guest]
1		600	0	600	1.17x10 ⁻³	0
3	0.10	600	20	620	1.13x10⁻³	1.14x10 ⁻⁴
5	0.20	600	40	640	1.10x10 ⁻³	2.21x10 ⁻⁴
7	0.30	600	60	660	1.06x10 ⁻³	3.22x10 ⁻⁴
9	0.40	600	80	680	1.03x10 ⁻³	4.16x10 ⁻⁴
11	0.50	600	100	700	1.00x10 ⁻³	5.07x10 ⁻⁴
16	1.00	600	200	800	8.78x10 ⁻⁴	8.85x10 ⁻⁴
19	2.00	600	400	1000	7.02x10 ⁻⁴	1.42x10 ⁻³
20	2.50	600	500	1100	6.38x10 ⁻⁴	1.61x10 ⁻³

Fluorescence Titration polymer/diammonium picrate

Two mother solutions of PC[5] and C10*2Pic 1.17×10^{-4} M in dry CHCl3/CH3OH were prepared. From these, different solutions with different ratio Host/Guest were prepared and fluorescence spectra (λ_{exc} =440 nm) were recorded.

Entry	Vtot	[Diammonium	۱/۱ _o
		Picrate]	
1	2000	0	1
2	2010	4,97512E-7	0,98251
3	2020	9,90099E-7	0,95866
4	2030	1,47783E-6	0,94595
5	2040	1,96078E-6	0,938
6	2050	2,43902E-6	0,92687
7	2060	2,91262E-6	0,91415
8	2070	3,38164E-6	0,91415
9	2080	3,84615E-6	0,89984
10	2090	4,30622E-6	0,89189
11	2100	4,7619E-6	0,88712
12	2120	5,66038E-6	0,86328
13	2140	6,54206E-6	0,84102
14	2160	7,40741E-6	0,8283
15	2180	8,25688E-6	0,80445
16	2200	9,09091E-6	0,78537
17	2250	1,11111E-5	0,74563
18	2300	1,30435E-5	0,70588
19	2350	1,48936E-5	0,6725

20	2400	1,66667E-5	0,6407
21	2500	2E-5	0,58506
22	2600	2,30769E	0,50874
23	2700	2,59259E	0,46105
24	2800	2,85714E	0,41335
25	2900	3,10345E	0,38156
26	3000	3,33333E	0,31797
27	3200	3,75E	0,27027
28	3500	4,28571E	0,17488

Acid-Base Effect on Fluorescence of PC[5]net

Different solutions, respectively containing polymer $(1.0 \times 10^{-4} \text{ M})$, diammonium picrate $(2 \times 10^{-3} \text{ M})$, Et₃N (4×10^{-3}) and Tri-Fluoro Acetic Acid (TFA) (4×10^{-3}) have been prepared. After mixing the polymer and the diammonium picrate to form **PC[5]**_{net}, different amounts of Et₃N and TFA have been done. After every addiction the fluorescence spectra have been recorded.

PC[5]r	net			
Entry	VEt₃N	VTFA	V _{τοτ} (μl)	Intensity
	(µI)	(µl)		(a.u.)
1	0	0	2000	663
2	5	0	2005	1
3	0	5	2010	-3
4	5	0	2015	-1
5	0	5	2020	-3
6	5	0	2025	-4
7	0	5	2030	-3



Acid-Base Effect on Fluorescence of PC[5]cap

Different solutions, respectively containing polymer $(1 \times 10^{-4} \text{ M})$, diammonium picrate $(2 \times 10^{-3} \text{ M})$, Et₃N (4×10^{-3}) and Tri-Fluoro Acetic Acid (TFA) (4×10^{-3}) have been prepared. After mixing the polymer and the diammonium picrate to form **PC[5]**_{cap}, different amounts of Et₃N and TFA have been done. After every addiction the fluorescence spectra have been recorded.

PC[5] _{cap}							
Entry	VEt₃N	VTFA	V _{τοτ} (μl)	I-I _o			
	(µI)	(µl)					
1	0	0	2000	0			
2	5	0	2005	15			
3	0	5	2010	-5			
4	5	0	2015	16			
5	0	5	2020	-9			
6	5	0	2025	11			
7	0	5	2030	-6			



Electrochemical measurements

Voltammetric characterization have been done in Tetrabutylammonium perchlorate

(TBAP) 0.1M solution in CH_3CN .

The electrochemical HOMO and LUMO energies were calculated from the oxidation

and reduction onset values and are shown in Table

CV Poly	mer (reduc	tion)					
Entry	Scan rate	Epc (mV)	Epa(mV)	Efc(mV)	LUMO	Mean	S.d
	(mV/s)						
1	50	-1604	-1266	-745	-4.06	-4.04	0.02
2	50	-1599	-1276	-739	-4.06		
3	50	-1620	-1250	-801	-4.00		
4	20	-1555	-1175	-760	-4.04		
5	20	-1495	-1171	-773	-4.03		
CV Polyı	ner (oxida	tion)		·		·	
Entry	Scan rate	Epc (mV)	Epa(mV)	Efc(mV)	НОМО	Mean	s.d
	(mV/s)						
1	50	+365	+552	+718	-5.52	-5.52	0
2	50	+350	+550	+720	-5.52	1	

General procedure for spin coating of different organic solutions.

Different solutions containing PCBM, P3HT and PPEcalix[5] in Chloroform were spin coated at different values of revolutions per minute (1500, 1000, 500 r.p.m) on n-Si or p-Si substrates. Samples were studied by XRR and AFM analysis



25 a



25 b



Fig. 10 : XRR graphics of PCBM layers a) 6 nm, b) 7 nm, c) 10 nm



26 a



26 b



Fig. 26 : XRR graphics of PCBM layers a) 15 nm, b) 18 nm, c) 23 nm



27 a



27 b



27 c

Fig. 27 : XRR graphics of PCBM layers a) 28 nm, b) 32 nm, c) 46 nm





Fig. 28 : XRR graphics of PCBM layers a) 61, b) 98 nm









Fig. 29 : XRR graphics of P3HT's layers a) 15 nm, b) 18 nm, c) 25 nm

V-CONCLUSIONS

A new aryl-acetylene polymer bearing calix[5]arene moieties as branches has been designed, synthesized and characterized with ¹H-NMR, GPC, Uv-vis and Fluorescence.

In order to evaluate the potential application of this polymer as a material for photovoltaic devices, it has been carried out a ¹HNMR and fluorescence titration with a 1,10-decanediyldiammonium picrate salt.

The ¹HNMR titration confirmed that there is an interaction between the calix[5]arene cavity and the diammonium salt; in particular we noticed that the different amounts of 1,10 decanediyldiammonium ion can regulate the supramolecular structure, allowing us to obtain two structures: PC[5]_{net} and PC[5]_{cap.}

Fluorescence titration demonstrated the reversibility of the assembly process of the two supramolecular species. Moreover, this experiment demonstrated the stability of the polycapsular polymer network upon the successive addictions of Et₃N and TFA.

Cyclic Voltammetry measurements have been carried out in solution in order to evaluate the values of the polymer's molecular orbital.

This measurements indicated that the HOMO-LUMO Energy values (-5.52 eV for the HOMO, -4.04 eV for the LUMO) make the PC[5] a suitable n-dopant in hybrid inorganic-organic solar cells.

In order to evaluate the possibility of using a new architecture for photovoltaic devices involving both inorganic (crystalline Silicon) and organic materials, spin coating of different solutions of organic molecules have been carried out.

Surface analysis (XRD and AFM) demonstrate that homogeneous layers having different thickness of organic materials can be obtained.

Finally, electrical measurements demonstrated that the polymer is suitable to assemble a diode, which, under illumination, is able to rectify the current in the correct sense for p-doped Silicon/PCBM interfaces and n-doped Silicon/P3HT interfaces.

This approach suggested the use of the same architecture for samples having the new polymer as n-type dopant.

I/V measurements showed that PC[5], PC[5]_{net} and PC[5]_{cap} can interact with p-type Silicon, with a rectifying effect, thus confirming the Voltammetry measurements suggesting this molecule as n-type dopant, in absence of a good metal contact Isc values cannot be determined. Qualitatively we found a good behaviour in terms of Isc for all the samples base on PC[5]_{cap}. This can be related to the supramolecular architecture allowing a better charge transfer to the probe electrode. The lower V_{oc} values respect to the probe molecule (PCBM) suggested the possibility of improving the Si/organic material interface.

We verified the possibility to create a supramolecular network based on the interaction between the calix[5]arene moieties and a suitable spacer in order to obtain a 3D network that could guarantee a better pathway for the charges that have to be collected after the exciton formation. Interestingly, after the electrical measurements on the simulating diodes, we saw that the better response was for the samples obtained by depositing on Si the PC[5]_{cap}, the less ordered specie, instead of the PC[5]_{net}, the more ordered one. This can be related not to the order of the organic layer, but simply to the fact that the PC[5]_{cap} is less fluorescent specie; so, although the less ordered pathway, the Energy provided by lighting the device can be involved in the Charge Transfer process instead of the Energy transfer. An higher charge transfer means a higher number of excitons can be formed, thus allowing higher and more reproducible I_{sc} values for the PC[5] layers depositing on Si.

VI- BIBLIOGRAPHY

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