Biofunctionalisation of p-doped silicon with cytochrome c553 minimises charge recombination and enhances photovoltaic performance of the all-solid-state photosystem I-based biophotocathode†

Julian David Janna Olmos, a,b Philippe Becquet, c Dominik Gront, d Jarostaw Sar, e Andrzej Dąbrowski, e Grzegorz Gawlik, e Marian Teodorczyk, e Dorota Pawlak e and Joanna Kargul a,**

Surface-directed passivation of p-doped silicon (Si) substrate was achieved by its biofunctionalisation with hexahistidine (His6)–tagged cytochrome c553 (cyt c553), a soluble electroactive photosynthetic protein responsible for electron donation to photooxidised photosystem I (PSI). Five distinct variants of cyt c553 were genetically engineered by introducing the specific linker peptides of 0–19 amino acids (AA) in length between the cyt c553 holoprotein and a C-terminal His6-tag, the latter being the affinity ‘anchor’ used for the specific immobilisation of this protein on the semiconductor surface. Calculation of 2D Gibbs free energy maps for the five cyt c553 variants showed a significantly higher number of thermodynamically feasible conformations of immobilised cyt c variants containing longer linker peptides. Here we show that the distinct cyt c553–based Si bioelectrodes display some characteristics of the p–n-type diodes, albeit varying in the level of dark saturation current J0 considered as the charge recombination parameter. These combined bioinformatic and electrochemical analyses indicate that the cyt c553 variants with longer linker peptides, up to 19AA in length, allow for more structural flexibility of immobilised cyt c553 in terms of both, orientation and distance of the haem group with respect to the Si surface, and promote the efficient biopassivation of the semiconductor substrate. Incorporation of the specifically immobilised 19AA cyt c553 variant into the all-solid-state biophotocathodes containing light harvesting PSI module enhanced biophotovoltaic performance of the PSI biophotocathode compared to the analogous device devoid of cyt c553.

1. Introduction

To date, the most technologically advanced photovoltaic devices are those based on silicon (Si). By taking advantage of the current state of the art of Si-based photovoltaics we are presently able to generate electricity from sunlight relatively cheaply. Currently, more than 90% of the global photovoltaics market is based on Si technology and assuming a growth rate for this technology of 40% p.a., Si solar cells carry the great potential to contribute significantly towards meeting the ever increasing energy demand of mankind. As Si is the second most abundant and evenly distributed element in the earth’s crust there is no risk of its shortage in the foreseeable future. To date, Si-based solar cells reach the highest reported power conversion efficiencies among single-junction solar cells. In fact, the second highest most efficient single-junction solar cell is the black Si solar cell with a reported efficiency of 27.6 ± 1.2\%.

One of the key factors for improving the power conversion efficiency of solar cells is to minimise wasteful processes of charge recombination by optimisation of the nanostructuring of the working modules and their interfaces. To this end, semiconductor surface passivation is of the particular importance, as it often results in significant minimisation of surface charge recombination, leading to the better overall performance of semiconductor-based solar cells. To date, several passivation approaches have been applied mainly based on elaborate chemical and electric field effects. Recently, a new generation of passivated Si solar cells with an efficiency of 22.1\%...
has been reported by Savin et al. who employed atomic layer deposition of Al2O3 on the surface of the nanostructured black Si to significantly decrease surface charge recombination by extending minority carrier lifetimes to the millisecond timescale. More recently, it has been shown that the key strategy for the enhancement of the power conversion efficiency is based on including two passivation layers; a top passivation layer on top of the photoactive substrate and a bottom passivation layer between Si and the photoactive substrate. Such an approach resulted in an increase of power conversion efficiency to over 26%.

Functionalisation of semiconductor electrode materials with natural light harvesting proteins such as photosystem I (PSI) is a particularly attractive concept, as such a ‘green’ biohybrid solar technology utilises the combination of relatively cheap and stable semiconducting material, such as Si, together with the highly efficient light-harvesting/charge separating natural pigment/protein complex which absorbs well in the blue-red far red part of the electromagnetic spectrum, and operates with the impressive internal quantum yield close to unity. However, the efficiency of the reported to date biohybrid PSI-based photoelectrodes is still limited to values below 1% mainly due to the recombination of photo-excited electron–hole pairs at the interface of the electrode modules, rather than by the PSI photochemical activity or stability. Other obstacles to obtain high power conversion efficiency of PSI-based biophotoelectrodes are poor electronic contact between the photoactive module and the electrode, as well as insufficient and heterogeneous PSI loading onto the electrode. Nevertheless, it has been recently shown that high energy photoactivated electrons that are generated in PSI upon light absorption can be recovered in biophotoelectrodes even faster than in natural photosynthesis itself, highlighting the great potential of PSI-based biohybrid solar technology once it is optimised.

Previously metal electrodes have commonly been used in PSI-based photobioelectrodes; however, application of semiconducting electrode materials allowed for a significant improvement of photocurrent and photovoltage outputs. Recently, the p-doped Si was found to be an excellent platform for electron donation to the photo-oxidised PSI due to well-matched valence and conduction bands for Si (0.5 V and -0.6 V vs. SHE, respectively), the P700 primary electron donor and the terminal electron acceptor, the iron–sulphur cluster in (0.43 V and -0.58 V vs. NHE, respectively).

We report that increasing the distance and structural flexibility of the haem group of immobilised cyt c553 (cyt c553) protein, the mobile redox-active electron donor of PSI, upon its specific immobilisation on the p-doped Si electrode leads to efficient passivation of the semiconductor surface, as measured by minimisation of the dark saturation current (I0) also known as the semiconductor surface recombination parameter. Importantly, incorporation of the most structurally flexible 19AA peptide linker cyt c553 variant as the conductive layer into the all-solid-state PSI-based biophotoelectrodes significantly enhanced the power conversion efficiency (20-fold) compared to the analogous device devoid of cyt c553, demonstrating feasibility of this molecular approach for improving the overall performance of PSI-based biophotoelectrodes.

2. Results and discussion

Optimisation of the electron transfer between an appropriate semiconductor substrate and PSI photoactive module is essential for improving the performance of such biophotoelectrodes. However, most of the PSI biosolar systems reported to date have relied on mass transfer through the use of artificial electron mediators (ref. 21, 23 and 24 and references therein), resulting in the significantly decreased performance of such devices. To overcome this limitation, our approach was to rationally engineer efficient direct electron transfer (DET) at the interface between the conductor surface and the (photo)electroactive PSI module by using the specifically oriented conductive layer of redox active cyt c553 in the all-solid-state device configuration. The 9 kDa cyt c553 protein functions in vivo as the natural electron donor to the photooxidised P700+ reaction centre of PSI upon the highly specific protein–protein interaction. The domain-specific interaction between cyt c553 and the PSI complex has been recently applied in the plasmonic nanostructures of silver nanowires, where it led to the spectacular enhancement of optical properties of PSI as well as improvement of photovoltaic output in various types of biophotoelectrodes.

2.1 Linker peptide design and structural modelling of the cyt c553 variants

The electroactive protein used in our study, that constituted the biological conductive layer of the PSI-based biophotoelectrode, is cyt c553 from an extremophilic red microalga Cyanidoschyzon merolae, whereas the electrocompatible semiconductor substrate is heavily p-doped Si. The cyt c553 protein is a structural and functional homologue of cyt c6 which is a soluble redox active protein serving in vivo as the native electron donor to the photo-oxidised P700+ reaction centre of PSI. Application of the highly robust redox active proteins as modules of biophotoelectrodes, such as the photosynthetic proteins from C. merolae used in this study, carries the great potential to improve the long-term performance and stability of such novel, mediatorless all-solid-state biodevices due to inherently increased stability of extremophilic biological components across a wide range of extreme conditions.

The nuclear genome of C. merolae contains a single gene petJ which encodes cyt c553 protein that is characterised spectroscopically by the specific x peak at 553 nm in its reduced state. The X-ray analysis of the red algal cyt c553 protein showed that it is a Class I c-type cyt in which the redox-active haem prosthetic group covalently binds to Cys34 and Cys37. The central Fe atom of the haem group displays octahedral coordination with His37 and Met8a axial ligands.

In the process of exploring alternative routes for the improvement of electron transfer at the solid-state interface of photovoltaic devices, it is important to design a rational approach for prediction of the optimal conformation of the immobilised redox active components that would allow for the efficient DET at the interfaces between the working modules. To this end, four distinct linker peptides of 5–19 amino acids (AA)
in length were designed between the cyt c553 holoprotein and a C-terminal His6-tag (see Table 1), the latter used to anchor the cyt protein to the Ni–NTA-functionalised p-doped Si electrode surface. The control His6-tagged cyt c553 variant, termed 0AA, constituted the His6-tagged cyt c553 holoprotein without any additional peptide linker. Overall, the rationale in the design of the four peptide linker variants of cyt c was to choose AA residues in the peptide sequence that would allow for a variable degree of conformational freedom of motion of the immobilised cyt c553. The 4 peptide linkers were designed based on their rigidity versus flexibility, as well as using a specific sequence known to naturally form a semi-β-helix in the case of the 19-AA linker peptide (see Fig. 1). The inclusion of hydrophobic (Ala) and highly hydrophobic (Leu and Val) AA residues aimed to induce rigidity to the specifically immobilised cyt c553.

The following AA variant characteristics could be predicted based on the physico-chemical properties of the respective AA: (i), 0AA: most restrictive conformation allowing for no rotation; (ii), 5AA: a minor degree of rotation and movement; (iii), 10AA: a small degree of rotation or movement; (iv), 12AA: more rotation or movement, and (v), 19AA: L-shaped hinged linker, with a semi-β-helical structure predicted to allow for conformational flexibility of the immobilised cyt c553 protein. The 5AA linker peptide is composed mainly of small, polar amino acids allowing for limited structural flexibility, particularly at the more charged C-terminal end of the sequence. The 10AA linker peptide comprises mainly polar Gly/Ser repeats and a charged Glu residue allowing for more structural flexibility of the immobilised cyt c553 variant compared to the 5AA variant. The 12AA linker corresponds to the original linker peptide encoded within the multiple cloning site of the petJ gene expression vector that has been genetically engineered between the cyt c553 holoprotein and a C-terminal His6-tag. It comprises polar Gly/Ser residues, non-polar Val/Ala/Leu residues, and negatively charged Asp/Glu residues. Finally, the 19AA cyt variant was designed to introduce a semi-helical ‘hinged’ linker peptide to the immobilised cyt c553 holoprotein. For this variant, we defined the Phi/Psi angles of each residue to be equal to −62/−41, which corresponds to the average values found experimentally for β-helices which are non-geometrically ideal.²⁸

### 2.2 Mapping of the Gibbs free energy of the cyt c553 variants and estimation of their structural flexibility

To obtain an insight into the thermodynamic properties of the distinct variants of cyt c553 upon their immobilisation on the electrode surface, we calculated a theoretical probability of acceptable conformation that each peptide linker can adopt.

### Table 1  Physico-chemical properties of the linker peptide sequences introduced within cyt c553 structure

| Cyt c553 variant | AA sequence of linker peptides | Linker length (AA no.) | Molecular weight (Da) | Theoretical pI |
|-----------------|--------------------------------|------------------------|-----------------------|----------------|
| 0AA             | Holoprotein-HHHHHHH             | N/A                    | N/A                   | N/A            |
| 5AA             | Holoprotein-GSGLE-HHHHHH        | 5                      | 461.4                 | 4.00           |
| 10AA            | Holoprotein-GSGSGSGLLE-HHHHHH   | 10                     | 836.8                 | 4.00           |
| 12AA            | Holoprotein-GSSVDKLAAALE-HHHHHH | 12                     | 1160.2                | 4.37           |
| 19AA (semi β-helix) | Holoprotein-AEAAAKEAAAKEAAKALE-HHHHHH | 19                     | 1814.0                | 4.95           |

Fig. 1  Modelling of the His6-tagged peptide linkers into the crystal structure of cyt c6 from Synechococcus sp. PCC 7002. The cyt c6 structure (PDB: 4EIC)²⁴ is the highest resolution type I c-type cytochrome (a close homologue of cyt c553 from C. merolae) crystal structure available to date and was used to model in the five distinct peptide linkers used in this study. The haem group of cyt c6 with its central Fe atom is displayed in red.

Fig. 2  Acceptance rates of different cyt c553 variants after Monte Carlo simulation test. The acceptance rate was calculated as the ratio of the number of energetically permissive moves to the total number of moves using the standard Metropolis criterion.²⁴ The acceptance rate calculation was excluded for the 0AA cyt variant as there is no peptide linker peptide between the two points of this simulation: namely the C-terminus of the holoprotein sequence and the first His residue of the His6-tag. Values of the acceptance rates are displayed above the error bars of each cyt c variant.
under random Brownian motion. Firstly, we applied a Monte Carlo simulation to calculate the acceptance rate for each cyt c<sub>553</sub> variant, corresponding to the ratio of a total number of energetically permissive moves divided by the total number of all the attempted moves. During the simulation, Phi and Psi dihedral angles of the linker were altered randomly from a uniform distribution whilst including the Si plane in the simulations (see Materials and Methods). The acceptance rate was used in this context as a measure of the thermodynamic ‘freedom of motion’ of each cyt c<sub>553</sub> variant upon its attachment to the p-doped Si surface via a His<sub>6</sub>-tag. It is worth noting that the acceptance rate largely depends on the degree of flexibility of the different AA side chains present within the linker peptides, implying that longer linker peptides would have larger probabilities of adopting different three-dimensional conformations, compared to the shorter counterparts.

Fig. 2 shows that the highest acceptance rate (0.6851 ± 0.04), corresponding to the most restricted free motion of the haem with respect to the Si surface, was calculated for the shortest (5AA) cyt c<sub>553</sub> variant, whereas the lowest acceptance rate value (0.3868 ± 0.02), corresponding to the least restricted free motion of the haem with respect to the Si surface, was attributed to the cyt c<sub>553</sub> variant containing a semi-α-helix (19AA) linker peptide within its C-terminal domain. The acceptance rate calculation was not applicable to the 0AA linker variant as no peptide was present between the C-terminus of the cyt c<sub>553</sub> protein and the affinity His<sub>6</sub>-tag.

We then calculated the 2D Gibbs free energy maps for each cyt c<sub>553</sub> linker peptide variant as a function of the absolute distance of the haem group of cyt c (i.e., its central Fe atom) and the relative planar angle between the haem and a flat electrode surface (see Fig. 3). Fig. 4 shows that the presence of the longer peptide linkers within the cyt c structure evokes a significantly larger number of thermodynamically favourable conformations of the anchored cyt c (in terms of probability of energetically favourable alignment of the haem group to the electrode surface) after a given move, and consequently, a lower acceptance rate (see Fig. 2), as per Monte Carlo criteria. On the other hand, a short peptide linker is more likely to adopt a thermodynamically unfavourable conformation, since very few combinations of dihedral angles are possible within the whole linker length. Therefore, a lower probability of proximal alignment of the haem group to the Si surface exists (see Fig. 4).

To assess whether the calculated Gibbs free energy levels and conformational flexibility of the experimentally designed peptide linkers can be extended to much larger populations of the similar-length peptides, we conducted simulations on a high number of randomly de novo generated peptide linkers of 5AA, 7AA, 10AA, 12AA and 19AA in length (see ESI Fig. 1†). The aim of this in silico study was to investigate how far the cyt c<sub>553</sub> protein can depart from the electrode for a given length of
a peptide linker, independently of its actual AA sequence. To this end, 600 linker sequences were randomly generated for each given peptide length, with each AA residue equally probable. For this simulation, we employed the same set of randomly chosen movers and applied one of them at each iteration on a random residue of the linker. The de novo designed 7AA linker population provided a new class of peptide linkers that were omitted in the original experimental design.

The energy levels of experimentally and in silico designed linkers are similar, indicating energetically favourable properties of the experimental linkers compared to the whole random peptide dataset (see ESI Fig. 1†). Significantly, these simulations also revealed the existence of peptide linkers that exhibit more suitable levels of free energy, which potentially could be beneficial for the highly efficient DET between the semiconductor substrate and P700 reaction centre of PSI. This hypothesis will be verified in the future study.

2.3 Purification and biochemical characterisation of cytochrome c553 variants

We have cloned and heterologously overexpressed the transgenes encoding all the five variants of cyt c553 in E. coli using the tightly catabolically controlled co-expression system in conjunction with the constitutive expression of the haem maturation cassette to ensure synthesis of the fully redox-active cyt c holoproteins. All the five His6-tagged cyt c553 variants were purified to homogeneity using Ni²⁺-NTA immobilised metal affinity chromatography, as shown in Fig. 5. The predominant pools of each cyt c553 variant were obtained in the reduced state, with ~20% of the total cyt yield obtained in the oxidised state (see Fig. 5B). The total yield of redox active cyt c553 variants was estimated in the range of 1–1.5 mg L⁻¹. We confirmed the full redox activity of all the purified cyt c553 variants by measuring the redox difference absorption spectra, in which the disappearance of 553 nm and 521 nm haem-specific peaks was clearly observed upon chemical oxidation (see Fig. 5C and ESI Fig. 2†).

2.4 Construction and characterisation of the cyt c553/p-doped Si bioelectrodes

Five distinct bioelectrodes were constructed by immobilisation of the purified His6-tagged cyt c553 variants on a p-doped Si substrate. The electrode surface was chemically functionalised with a Ni²⁺-NTA self-assembled monolayer (SAM) to facilitate anchoring of cyt c553 on the electrode surface (see ESI Fig. 3†). We introduced the Ag and Au/Cr back and front contacts (200 and 50 nm, respectively) to perform J–V measurements, as schematically depicted in Fig. 6A. We have chosen p-doped Si as the electrode material as this semiconductor has been shown to...
serve as an excellent substrate for cyt c,
serve as an excellent substrate for cyt c, 
serve as an excellent substrate for cyt c, due to the mid-point potential of this type of cyt (+210 mV) being close to the estimated Fermi level of the conduction band of p-doped Si (+300 mV vs. NHE) with the level of p-doping used in our study (see Fig. 6B for the energy band diagram). The thickness of a single layer of cyt c was estimated as 3 nm, taking account of the dimension of the close homologue of this protein, cyt c53, determined by X-ray crystallography (PDB: 4EIC).14

One of the major questions we set out to address in this study was to evaluate the effect of redox active cyt c53 layer on the inherent charge transfer within the p-doped Si substrate. To this end, we assessed the effect of the distinct cyt c53 variants on the electron diffusion and recombination processes at the surface of the biofunctionalised Si electrode by two independent approaches: J–V measurements in the dark (see Fig. 7) and electrochemically, by two-electrode all-solid-state linear sweep voltammetry (Fig. 8). All the bioelectrodes displayed some diode-like behaviour. Notably, the dark saturation current ($J_0$) of heavily p-doped Si, which is the recombination parameter within a given semiconductor in its solid-state configuration,19 was lower upon biofunctionalisation of the p-doped Si substrate with all the cyt c53 variants compared to the p-doped Si-SAM control (see Fig. 7, 8 and Table 2), albeit to a different degree depending on the peptide variant used. On one hand, the presence of short peptide linker cyt c53 variants on the Si surface resulted in the higher $J_0$ values compared to variants with longer peptide linkers (see Table 2).

Our Gibbs free energy modelling and calculation of movement acceptance rates (see Fig. 2 and 4) showed that the short linker peptide cyt c variants display limited structural flexibility upon their anchoring with the Si surface. It is therefore plausible that their redox active haem groups cannot properly align with the semiconductor surface to promote efficient reversible redox reactions, i.e., Fe$^{3+}$/Fe$^{2+}$ interconversion within the haem group. We identified two distinct populations of the electrodes biofunctionalised with the 0AA cyt c53 variant (see Table 2, Fig. 7 and 8), in line with identifying two distinct conformational populations of this variant (see Fig. 4). The existence of such distinct electrode populations for the 0AA variant can be due to the presence of favourable and unfavourable orientation of the haem group in terms of reversible redox reactions and minimisation of surface charge recombination.

On the other hand, the haem groups present in the longer linker peptide variants of cyt c53 are likely to be more capable of the favourable alignment with the semiconductor surface due to higher structural flexibility of the peptide linkers compared to their shorter counterparts (see Fig. 4). Consequently, of all the cyt c53/p-doped Si bioelectrodes analysed in this study, the device constructed with the 19AA peptide linker variant displays the lowest degree of p-doped Si surface charge recombination, as measured by the lowest dark $J_0$ parameter (see Table 2), most likely due to the largest number of the thermodynamically favourable conformations of 19AA cyt c53 variant upon its immobilisation on the electrode surface, as well as the highest probability of the haem orientation favouring minimisation of the Si surface charge recombination.

The precise mechanism of electron transfer between the semiconductor electrode and the haem group of cyt c is still not fully understood. While some quantum mechanical calculations suggest that the haem group must be parallel to the electrode surface in order to promote the efficient DET,41,42 also described as “superexchange”,43 other studies suggest the
possibility of electron transfer when the haem group is perpendicular to the surface of the electrode.\(^{44}\) Introduction of the 19AA and to a smaller degree, the 12AA linker peptides into the cyt c\(_{553}\) structure allows for a larger number of thermodynamically favourable conformations of immobilised cyt c with various types of the haem group alignment with respect to the plane of the electrode, as evidenced by both, our Gibbs free energy maps (see Fig. 4) and lower \(J_0\) values (see Table 2, Fig. 7 and 8). An alternative mechanistic explanation could be provided in the context of molecular orbital theory and electronic excitation of the haem group in the solid-state configuration. As short, rigid AA linkers yield more restricted conformations, there is less time for the haem group to relax electronically to the lower energy state (or energy well) characteristic for steady-state charge carrier hopping,\(^{45}\) as observed in solid-state electronic devices.\(^{46}\) This hypothesis would explain why the \(J_0\) parameter for the 5AA cyt c\(_{553}\) variant biodiode is much higher than those with longer (see Table 2), more flexible variants, whereby the haem group would be expected to relax longer, being in a further distance from the p-doped Si surface.

Although a number of studies have attempted to understand the intermolecular kinetics dominating the complex processes of electron transfer between the active site of the redox active protein and the semiconductor surface,\(^{47-50}\) to date only several reports have employed p-doped Si as the semiconductor substrate together with cyt c protein either to investigate the level of conductance\(^{41,46,48,51}\) or the electrochemical properties of cyt c on p-doped Si.\(^{39}\) Nevertheless, none of these studies have addressed the electrochemical effects of cyt c on the inherent semiconductor solid-state surface junction properties. Amdursky and colleagues\(^{46}\) showed that the specific electronic

![Diagram](image)

**Fig. 8** Linear sweep (two-electrode) voltammograms of the all-solid-state cyt c\(_{553}/\)Si bioelectrodes. The cyclic voltammograms were obtained in a two-electrode measuring setup in the dark, at a scanning rate of 10 mV s\(^{-1}\). The potential was cycled from −500 mV to 500 mV. Note that the Ni–NTA SAM also contributes to the redox activity of the solid-state construct. The absolute value of the dark saturation current (mA) is inversely dependent on the length of the linker peptide, with the highest current for the 0AA linker variant and the lowest current for the 19AA cyt c\(_{553}\) variant electrode.

![Diagram](image)

**Fig. 9** Diagrammatic representation of PSI-LHCl containing devices. (A) PSI-LHCl/p-doped Si nanodevice (third (III) generation biophotovoltaic device). The thickness of each layer is indicated on the right and was estimated by sputtering detection. The components are not to scale. (B) Diagrammatic representation of the PSI-LHCl/cyt c\(_{553}\) p-doped Si nanodevice (fourth (IV) generation biophotovoltaic device). The thickness of each layer indicated on the right was estimated by sputtering detection. The components are not to scale. All Si layers were functionalised and contained a Ni–NTA SAM, as depicted schematically in ESI Fig. 3.f

| Bioelectrode type | \(J_0^e\) (mA cm\(^{-2}\)) |
|------------------|-----------------|
| Control with SAM\(^b\) | 1.2 \times 10^{-1} |
| 0AA\(^e\) | 4.5 \times 10^{-2} |
| 0AA\(^d\) | 2.8 \times 10^{-3} |
| 5AA | 6.9 \times 10^{-2} |
| 10AA | 2.3 \times 10^{-2} |
| 12AA | 3.5 \times 10^{-2} |
| 19AA | 7.5 \times 10^{-3} |

\(J_0^e\), dark saturation current (inherent recombination parameter) of each solid-state device. The \(J_0\) values were obtained from eqn (1) (assuming that at 300 K, \(kT/q = 25.8\) mV) and were also estimated graphically at the y-intercept in ESI Fig. 4. \(^e\) For a scheme depicting SAM synthesis refer to ESI Fig. 3. \(^f\) Value was obtained for two 0AA linker variant electrodes which displayed similar \(J-V\) curves (see Fig. 7). \(^e\) Value was obtained for two 0AA linker variant electrodes which displayed similar \(J-V\) curves (see Fig. 7). \(^d\) Values were calculated for the six different bioelectrodes that were \(J-V\) characterised at 300 K.

![Image](image)
The conductance to the haem group of bovine cyt c is directly related to the distance between the haem group and the electrode surface, in line with our observations of this study. However, the same authors focused mainly on defining the physico-chemical parameters governing the Fe–Si conductivity, rather than p-doped Si biopassivation by cyt c, nor did they display any charge transfer data from the p-doped Si control, in contrast to our study. Interestingly, although two of the aforementioned studies investigated the level of conductance rather than the overall effect of redox active proteins on the inherent charge transfer properties of the semiconductor substrate, their \( J-V \) curves show a striking similarity to our dark \( J-V \) data (Fig. 7 and 8), indicating that indeed, the cyt c553 may evoke similar redox processes at the surface of the semiconductor, promoting “superexchange” at the solid-state interface. Moreover, Amdursky and colleagues ascertain that distinctly high current density values (in the mA cm\(^{-2}\) range) may be attributed to properly linked cyt c holoprotein with a fully operational redox active haem in the vicinity of the p-Si surface, in line with our data. The same authors demonstrated the direct

![Photochronoamperometric determination of open-circuit potential (\(V_{oc}\)) within PSI/p-doped Si (Panel A) and PSI/19AA cyt c553/p-doped Si biophotovoltaic devices (Panel B). After averaging the three independent devices, an increase in 333 µV is observed for the 19AA cyt variant-containing biophotoelectrode (Panel B) compared to the biophotoelectrode containing the PSI module alone (Panel A). The average \( V_{oc} \) increment for the biophotoelectrode containing PSI-LHCI sample is 1000 µV (shown in Panel A), while the average \( V_{oc} \) increment for the biophotoelectrode containing PSI-LHCI in addition to the 19AA linker variant is 1333 µV (shown in Panel B). Blue and red arrows symbolise light on and off, respectively. Potential (µV) values on the y-axis have been rescaled for the ease of \( \Delta V_{oc} \) estimation.](Fig. 10)

![Photochronoamperometric determination of short circuit current (\(J_{sc}\)) within the best performing PSI/p-doped Si and PSI/19AA cyt c553/p-doped Si biophotovoltaic devices. No overpotential was applied to determine \( J_{sc} \) parameter. Blue and red arrows symbolise light on and off, respectively. Current density values on the y-axis have been rescaled for clarity.](Fig. 11)
proportionality between haem group distance from the Si surface and an increased series resistance, similary to the $J-V$ data in the present study. Interestingly, such correlation between the level of passivation and the increased resistivity seems to be a general feature also in the case of chemically passivated semiconductor substrates, including Si.

2.5 Construction and photovoltaic characterisation of the PSI/cyt $c_{553}$/p-doped Si biophotocathodes

Our $J-V$ and in silico data strongly suggest that the 19AA linker variant of cyt $c_{553}$ is likely to be the best candidate for biopassivation of the p-doped Si substrate, as this variant significantly minimises the $J_0$ parameter within the p-doped Si substrate, over and above the other cyt variants used in this study. We therefore hypothesised that this feature introduced with the specific immobilisation of the 19AA cyt $c_{553}$ variant onto the p-doped Si substrate may lead to the overall improvement of the PSI-based biosolar cell's performance. Thus, the additional charge-separation event upon photoactivation of the PSI module on top of the cyt $c_{553}$/p-doped Si electrode would promote the enhanced DET, resulting in the desired downhill energy transfer and enhanced power conversion efficiency.

To verify this hypothesis, we constructed and characterised the photovoltaic properties of two types of nanodevices encompassing both 19AA cyt $c_{553}$ variant and PSI biological redox active modules (see Fig. 9 for schematic depiction of the PSI-based biophotocathodes), and compared these characteristics with the control devices built exclusively with PSI photoactive layer on top of the p-doped Si substrate, and devoid of cyt $c_{553}$. In order to assess the biophotovoltaic performance of the biophotoelectrode, we determined the open-circuit potential ($V_{oc}$) and the short-circuit current ($J_{sc}$) parameters by photocurrent-voltage measurement (see Fig. 10, respectively).

Table 3 Comparison of PV performance of all-solid-state mediatorless PSI/p-doped Si biophotovoltaic devices

| Device configuration | $J_{sc}$ [mA cm$^{-2}$] | $V_{oc}$ [V] | FF | $\eta$ [%] | References |
|----------------------|--------------------------|--------------|----|-----------|------------|
| p-Si/19AA/PSI        | 1,550$^b$                | 0.001333$^b$ | 0.3$^c$ | 0.0006$^c$ | This work  |
| p-Si/PSI             | 100$^b$                  | 0.001$^b$    | 0.3$^c$ | 0.00003$^c$ | This work  |
| p-Si/PSI/ZnO/ITO     | 127                      | 0.214        | 0.28 | 0.0077    | 19         |
| PAni/PSI/TiO$_2$/SnO$_2$ | 72                      | 0.299        | 0.42 | 0.0091    | 15         |

$^{a} J_{sc}$: Short circuit current, the current density when the voltage across the device is 0 V (see Fig. 11). $V_{oc}$: open circuit voltage or open circuit potential (the photovoltage generated when no bias is present across the circuit) (see Fig. 10). FF: fill factor. $\eta$: external power conversion efficiency of the biophotovoltaic device. The performance of the listed solid state biosolar cells was evaluated at standard white light illumination of 100 mW cm$^{-2}$ (AM 1.5), as described in Materials and Methods. $^b$ Values were estimated for selected electrodes with the best performance. Significant variabiliy was observed between devices due to fragile construction (brittle nature of the p-Si). $^c$ Calculation of fill factor and efficiency was performed with the open circuit voltage for the estimated values presented in the table assuming an ideality factor of 1 ($n = 1$) and a temperature of 300 K, as specified in Materials and Methods.

3. Concluding remarks

In summary, we provide the first evidence that incorporation of a redox-active robust biological component, the extremophilic cyt $c_{553}$ protein, allows for passivation of the p-doped Si substrate by minimisation of the inherent dark saturation current ($J_0$). By introducing the specific linker peptides within the cyt $c_{553}$ structure, we achieved minimisation of surface charge recombination within the p-doped Si substrate by optimisation of the alignment of the haem group with respect to the semiconductor surface. We show that the cyt $c_{553}$ variant characterised by the largest structural flexibility upon its specific immobilisation on the semiconductor surface allowed for a significantly improved biophotovoltaic performance of the all-solid-state PSI-based biophotocathode. Thus, our molecular approach has paved the way for developing of an alternative, cheap and facile route for significant reduction of the inherent minority charge recombination on the semiconductor surface. As such, it carries the great potential for future biotechnological applications, such as the development of viable ‘green’ solar-to-fuel devices.

4. Materials and methods

4.1 Monte Carlo simulations

A Monte Carlo object was used to assess the probability of free motion of the various peptide linkers attached to cyt c. This object accepts or rejects a modified protein using standard Metropolis criterion. An acceptance rate is defined by PyRosetta® software and it corresponds to the number of accepted moves divided by the total number of moves attempted. The Monte Carlo test used a score function with the

$\text{References}$
default Rosetta full-atom energy terms and weights (see http://www.pyrosetta.org). A set of the following movers, provided by the standard PyRosetta® distribution, was employed for the simulation: (1) SmallMover: modifies the Phi or Psi angle of a randomly chosen residue, by a small random angle. It was also used under the Metropolis acceptance criterion;15 (2) ShearMover: modifies the Phi and Psi angle of a randomly chosen residue, by a small random angle. This was also used under the Metropolis acceptance criterion;15 (3) SequenceMover: applies a sequence of several movers in succession. In our case, this mover contained the two previous movers. (4) RandomMover: applies a randomly chosen mover within a given list. For this simulation, we applied the SmallMover, the ShearMover, or RandomMover: applies a randomly chosen mover within a list. For this simulation, we applied the SmallMover, the ShearMover, or both of them (SequenceMover); (5) RepeatMover: applies a mover \( n \) times, \( n \) being given by the user. The RandomMover simulation was repeated several times on each cyt c variant. The same set of randomly chosen movers were applied, one per each iteration, on randomly generated AA residues of the fixed-length linkers of 5AA, 7AA, 10AA, 12AA and 19AA in length. The standard full-atom scoring function was used without hydrogen bonds, solvation and electrostatics energy components. For each of the linkers, the energy levels lower than 0.5% of the maximum energy value were extracted and, within this range, the maximum and minimum lengths in \( \AA \) as well as the maximum and minimum energies were stored and plotted in the graphs in ESI Fig. 1. The same simulation was performed on the experimentally designed linkers.

4.2 Bioinformatics calculations of the Gibbs free energy and modelling of solid-state cytochrome c-based bioelectrode

The Monte Carlo simulation protocol described above was used to characterise Gibbs free energy as a function of haem orientation relative to the electrode, as described by two internal coordinates. The distance between Fe atom and the flat electrode surface and the angle between haem plane and flat electrode surface (see Fig. 3) were used for this purpose. All the simulations were performed using the crystal structure of cyt \( \text{c}_6 \) from Synechococcus sp. PCC 7002 (PDB: 4EIC)24 with an appropriate linker added. The geometry was calculated for each modelled conformation with the BioShell package.25

4.3 Cloning and expression of \( \text{petJ} \) gene in \( E. \) coli

Genomic DNA from \( C. \) merolae was used as a template for polymerase chain reaction to amplify the \( \text{petJ} \) gene (annotated CMV193C; http://merolae.biol.s.u-tokyo.ac.jp/db/) encoding the cyt \( \text{c}_{553} \) protein. The original verified sequence of \( \text{petJ} \) was cloned into pET28b(+) vector (Invitrogen®) together with four distinct DNA sequences corresponding to the four distinct AA linkers (see Table 1 for AA sequences) between the \( \text{petJ} \) sequence and the C-terminal His\(_6\)-tag. The entire constructs were excised with Ncol and PstI from pET28b(+) vectors and subcloned into a pBAD/HisA expression vector (Invitrogen®) using the same restriction enzymes, then transformed into the \( E. \) coli strain DH5\( \alpha \). The correctness of the petJ::pBAD/HisA construct was confirmed by DNA sequencing. A specialised constructively expressed vector comprising a multiple gene cassette encoding the cyt c type haem group maturation enzymes24 (a gift from Prof. A. Szczepaniak, Wrocław Univ., Poland), was co-transformed with the petJ::pBAD/HisA recombinant vector into the expression host strain of \( E. \) coli, TOP10. Expression cultures inoculated from the single recombinant colonies were supplemented with 30 \( \mu \)g mL\(^{-1}\) chloramphenicol, 100 \( \mu \)g mL\(^{-1}\) ampicillin, and grown overnight. The overnight saturated pre-cultures (1 : 100 volume ratio) were then grown to OD\(_{600}\) 0.4. The cultures were then supplemented with 100 mg L\(^{-1}\) Fe\(_2\)SO\(_4\) and 17 mg L\(^{-1}\) \( \delta \)-aminolevulinic acid, and \( \text{petJ} \) gene expression was induced with 0.2% arabinose and allowed to proceed for 20 hours at 37 °C, with shaking at 160 rpm. The cells were harvested at 4000 \( \times \) \( g \) for 15 min. at 15 °C and then chilled on ice for 20 min.

4.4 Cytochrome \( \text{c}_{553} \) purification

Recombinant \( E. \) coli cells expressing the \( \text{petJ} \) gene were resuspended in a buffer containing 20 mM imidazole, 100 mM standard phosphate buffer (PB) (pH 7.5), 25% glycerol (w/v), 30 \( \mu \)M PMSF and 20 \( \mu \)g of DNase I per 50 mL of suspension or a protease inhibitor cocktail (Roche®, 1 tablet per 50 mL of resuspension buffer). Cells were ruptured by vigorous agitation with 0.1 mm glass beads in a Bead Beater (BioSpec®) using 18 cycles of 45 s and interim 4 min. cooling-off periods. Cell lysate was separated from the beads by vacuum filtration. Cell debris was removed by ultracentrifugation for 45 min at 100 000 \( \times \) \( g \) at 4 °C. After centrifugation samples were filtered using 0.2 \( \mu \)m disposable filtering devices (Millipore®) and applied onto an HPLC preparative Ni–NTA GE Healthcare HisTrapTM® HPLC/FPLC column (1 mL bed volume) at a flow rate of 0.5 mL min\(^{-1}\) to ensure efficient binding of the His\(_6\)-tagged cyt \( \text{c}_{553} \) variants. The column was then washed for 30 min at a flow rate of 0.5 mL min\(^{-1}\) with 50 mM imidazole and then the bound proteins were eluted with a linear gradient of 50–420 mM imidazole at a flow rate of 0.5 mL min\(^{-1}\). The column was further washed with 500 mM imidazole for 30 min to remove any residual proteins. Elution of cyt \( \text{c}_{553} \)-containing fractions was monitored at 416 nm, 521 nm and 553 nm, corresponding to the absorption peaks of reduced cyt \( \text{c}_{553} \). Cyt-containing fractions were pooled and concentrated using VIVASPIN-2 devices (3000 MWCO, Sartorius-Intec®, Poland), then resuspended in a buffer containing 100 mM PB (phosphate buffer, 100 mM Na\(_2\)HPO\(_4\), 18 mM KH\(_2\)PO\(_4\), pH 7.5) and 25% glycerol (w/v). The total protein concentration was determined using the Bradford assay23 using bovine serum albumin as a standard. Purity of the cyt \( \text{c}_{553} \) protein was verified on 16% Tris–Tricine SDS-PAGE gels using a standard protocol.26 The cyt \( \text{c}_{553} \) concentration was determined by redox difference absorption spectroscopy using extinction coefficient of 24.1 mM\(^{-1}\) cm\(^{-1}\) as described below (refer to ESI Fig. 2† for exemplary redox difference spectra).
4.5 SDS-PAGE gel electrophoresis and His₆-tag detection

SDS-PAGE was performed using a Tris–Tricine system.²⁶ Protein bands of IMAC pooled fractions (normally 3–5 µg protein per lane or equal volume, see below) were resolved on a 16% polyacrylamide gel at a constant voltage of 30 V for 30 min. After 30 min., the voltage was increased to 200 V for 45–60 min. Proteins were visualised with Coomassie Brilliant Blue R-250 staining or electro-transferred onto a polyvinylidene difluoride membrane (Millipore®) for detection of His₆-tagged proteins. Blots were probed with the HisProbe-HRP® conjugate (Thermo Fisher Scientific®, USA) and visualised by chemiluminescence, according to the manufacturer’s protocol. Prior to SDS-PAGE, samples were resuspended in 5 × SDS-PAGE sample buffer (0.225 M Tris–HCl, pH 6.7, 50% glycerol, 5% SDS, 0.05% bromophenol blue, and 0.25 M DTT), vortexed vigorously and boiled at 95 °C for 8 min with continuous shaking. Samples were centrifuged at 14 100 × g for 5 min and the supernatant was carefully loaded onto the SDS-PAGE gels. The volume was adjusted to 30 µg total protein per lane.

4.6 Redox difference absorption spectroscopy

UV-VIS spectra of all cyt c₅₅₃ variants were recorded using a Shimadzu UV 1800 spectrophotometer at RT. The cyt c₅₅₃ protein was titrated to the appropriate concentration (5–20 µg mL⁻¹ cyt c₅₅₃ in 100 mM standard phosphate buffer, pH 7.5) and the first RT absorbance spectrum was measured. To ensure the complete reduction of cyt, a few grains of sodium dithionite were added to the cuvette. After the grains were fully dissolved, another spectrum was measured, and if necessary more grains of sodium dithionite were added (grain by grain) until the fully reduced state of cyt was achieved. Subsequently, a grain of FeCN was added to oxidise the cyt c₅₅₃ protein. A spectrum was measured, in which flattening of the 521 nm and 553 nm peaks was observed. A grain of FeCN was added repeatedly until cyt c₅₅₃ was fully oxidised displaying a shift in the blue region from 416 nm to 406 nm. The cyt c protein was subsequently re-reduced with addition of approximately two times the amount of sodium dithionite upon which the reappearance of the 416, 521, 553 nm peaks occurred. Delta Aₕₜₜ (difference between maximally reduced and maximally oxidised species) was used to calculate the exact concentration of redox active cyt c₅₅₃ using the molar extinction coefficient of 24.1 mM⁻¹ cm⁻¹.²⁷ For all the cyt c₅₅₃ preparations, the absolute amounts of protein determined via the Bradford assay and by redox difference absorption spectroscopy were comparable. Refer to ESI Fig. 2f for exemplary redox difference spectra of the 4 cyt c₅₅₃ variants used in this study excluding the 10AA linker variant spectra which is displayed in Fig. 5C.

4.7 Construction of the cyt c₅₅₃ bioelectrodes and dark current characterisation

Heavily p-doped Si wafers were obtained from Department of Silicon Technology in the Institute of Electronic Materials Technology (ITME, Warsaw, Poland). Silicon wafers had a thickness of 500 ± 25 µm and a resistivity of 0.001–0.005 Ω cm equivalent to a boron doping density of 2.13 × 10¹⁹–1.3 × 10²⁰ cm⁻³. The wafers were pre-treated by submerging in SWIPE® for 24 hours. After removal of the detergent, the wafers were treated with hot, concentrated sulfuric acid (96%), a solution of ammonia in hydrogen peroxide and water (H₂O : H₂O₂ : NH₄OH / 1 : 1 : 0.05), and finally in diluted hydrofluoric acid in water (HF : H₂O / 1 : 10). Subsequently a 200 nm Ag layer was deposited through chemical evaporation on the back side of the wafers. Rapid Thermal Processing (RTP) was performed to improve metal/semiconductor contact. Subsequently, the wafers were again treated with diluted hydrofluoric acid (as above) and then functionalised with Ni-NTA according to a protocol from⁸ (See ESI Fig. 3f). After functionalisation, cyt c₅₅₃ variants were deposited using vacuum-assisted spin coating to generate a thin layer of cyt chelated to the Ni–NTA moiety. For this, a solution of 500 µL of 125 µM cyt c₅₅₃ variant was employed for deposition via vacuum-assisted spin coating. After deposition cyt c₅₅₃ was initially dried with a nitrogen stream and then left for further drying in nitrogen cabinet for 24 h. A transparent 5 nm layer of an Au–Cr alloy was then deposited to ensure the proper metal/cyt c₅₅₃ electrical contact. Several contact points were grown by chemical evaporation of this Au–Cr alloy employing laser-cut masks and the rest of the surface was sealed by a transparent 50 nm SiO₂ layer. Four to six biocathodes were cut into the precise square shapes and measured to determine exact area (3.24 cm²) of the biocathodes for each single variant electrode. An extra 50 nm layer of the Au–Cr alloy was chemically deposited on top of the bottom Ag contact to avoid any unwanted electrochemical Ag oxidation/reaction upon further manipulation for all-solid-state (mediatorless) electrochemical experiments.

4.8 J–V measurements of cyt c₅₅₃/p-doped Si bioelectrodes

All the J–V data was acquired using a custom-made setup comprising a PC with an interface card (Texas Instruments®, an AGILENT 34401A digital multimeter, the AGILENT 6634B system DC power supply and a home-made interface card (Department of Optoelectronics, ITME). Data was processed using ITME-customised LAB VIEW 8.6 software. Data was averaged using four independently prepared bioelectrodes using the selected area of 3.24 cm² which was employed as a standard area for all the electrodes constructed in this study. The entire protocol was repeated twice to ensure reproducibility of the J–V results. The J–V values were averaged for three independent bioelectrodes using the data from three most similar J–V curves from a total set of four measurements. All the J₀ values were calculated from eqn (1) below, and were also determined from the J–V curves presented on a semi-logarithmic scale, where the J₀ value corresponds to the y-intercept (see ESI Fig. 4f).⁴⁶

\[
J = J_0 e^{(qV/nkT)}
\]  
(1)

where: \(J\), the net current flowing through the diode; \(J_0\), dark saturation current (diode leakage current in the dark); \(V\), applied voltage across the terminals of the diode; \(q\), absolute
value of electron charge; $n$, ideality factor; $k$, Boltzmann’s constant; $T$, absolute temperature (K).

### 4.9 All-solid-state two-electrode cyclic voltammetry of cyt c$_{553}$/p-doped Si bioelectrodes

The CVs were obtained in a two-electrode configuration at a scan rate of 10 mV s$^{-1}$ using an SP-300 Bio-Logic Potentiostat. The potential was cycled from $-500$ mV to 500 mV. The bioelectrodes were connected by gold-plated Kelvin clamp. The top contact of the bioelectrode was connected as a working electrode and the counter and reference electrodes were connected to the bottom contact.

### 4.10 Construction of PSI/p-doped Si and PSI/cyt c$_{553}$/p-doped Si devices

The p-doped Si wafers were prepared essentially as described in Section 4.7, with or without chemical modification, where appropriate. After biofunctionalisation with cyt c$_{553}$, the Si wafers were dried in a semi-anaerobic nitrogen box for 24 h. A 500 mL suspension of the C. merolae PSI-LHCI complex at 1 mg mL$^{-1}$ Chla was overlaid on the pre-treated Si wafers and the protocol described in Section 4.7 was followed for the vacuum-assisted spin coating of the protein sample, with the exception that instead of a 5 nm of Au layer, a 5 nm of Ag was employed. For a flow-chart depicting preparation of all-solid-state PSI/cyt c$_{553}$/p-doped Si biophotovoltaic devices refer to ESI Fig. 5.$^\dagger$

### 4.11 Photochronoamperometry

All photoelectrochemical measurements on PSI-based devices were performed with a CH Instruments CHI660a electrochemical workstation (CH instruments, Inc.). The working electrode clip was attached to the Ag bottom contact layer, and the counter and reference electrodes were clipped on the top Ag layer contact as depicted in Fig. 9. The light source employed was an Industrial Solar Cell Testing Light Source (150/300 W) (Solar Light®) and the intensity output was 100 mW cm$^{-2}$ for all the photocurrent measurements, reasessed independently with a quantum metre (HD 2302.01) (DeltaOhm®) to a value of 2200 μmol photons per m² per s. Samples were positioned in the same distance from the light source for all the measurements, with frequent verification of constant light intensity. Photochronoamperometric data was collected with no bias between the reference and working electrodes for measurement of photo-driven open-circuit voltage changes. Short circuit current was measured when the voltage across the p-n-junction was 0 V.

The following equations were employed for determining fill factor (FF) and external quantum efficiency ($\eta$) of the PSI-containing biophotocatalysts (see Table 3):

$$FF = \left(\frac{V_{oc} - \ln(V_{oc} + 0.72)}{V_{oc} + 1}\right)$$

$$\eta = \left(\frac{V_{oc} \times J_{sc} \times FF}{P_{in}}\right) \times 100\%$$

where: $P_{in}$ power input was determined to be 0.324 W (with AM 1.5 irradiation) into an 18 × 18 mm$^2$ biophotocatalyst surface used throughout this study. Normalised $V_{oc}$ was employed for all calculations and thermal equilibrium was assumed, with the ideality factor of 1 and the temperature 300 K.

### Conflicts of interest

There are no conflicts to declare.

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### References

1. N. Armaroli and V. Balzani, Chem.–Eur. J., 2016, 22, 32–57.
2. G. Kavlak, J. McNerney, R. L. Jaffe and J. E. Traneck, Energy Environ. Sci., 2015, 8, 1651–1659.
3. M. A. Green, K. Emery, Y. Hishikawa, W. Warta and E. D. Dunlop, Prog. Photovoltaics, 2015, 23, 1–9.
4. K. Yoshikawa, H. Kawasaki, W. Yoshida, T. Irie, K. Konishi, K. Nakano, T. Uto, D. Adachi, M. Kanematsu, H. Uzu and K. Yamamoto, Nat. Energy, 2017, 2, 17032.
5. J. Schmidt, F. Werner, B. Veith, D. Zielke, S. Steingrube, P. P. Altermatt, S. Gatz, T. Dullweber and R. Brendel, Energy Procedia, 2012, 15, 30–39.
6. T. Tian, R. Gao, Q. Zhang, S. Zhang, Y. Li, J. Lan, X. Qu and G. Cao, J. Phys. Chem. C, 2012, 116, 18655–18662.
7. J. Tian, Q. Zhang, E. Uchaker, R. Gao, X. Qu, S. Zhang and G. Cao, Energy Environ. Sci., 2013, 6, 3542.
8. H. Savin, P. Repo, G. von Gastrow, P. Ortega, E. Calle, M. Garin and R. Alcubilla, Nat. Nanotechnol., 2015, 10, 1–6.
9. S. W. Hogewoning, E. Wiensjes, P. Douwstra, G. Trouwborst, W. van Ieperen, R. Croce and J. Harbinson, Plant Cell, 2012, 24, 1921–1935.
10. T. Kothe, S. Poeller, F. Zhao, P. Fortgang, M. Roegner, W. Schuhmann and N. Plumere, Chem.–Eur. J., 2014, 20, 11029–11034.
