

Provided by the author(s) and University of Galway in accordance with publisher policies. Please cite the published version when available.

Title	Amperometric flow injection analysis of glucose and galactose based on engineered pyranose 2-oxidases and osmium polymers for biosensor applications
Author(s)	Kurbanoglu, Sevinc; Zafar, Muhammed Nadeem; Tasca, Federico; Aslam, Iqra; Spadiut, Oliver; Leech, Dónal; Haltrich, Dietmar; Gorton, Lo
Publication Date	2018-04-25
Publication Information	Kurbanoglu, Sevinc, Zafar, Muhammed Nadeem, Tasca, Federico, Aslam, Iqra, Spadiut, Oliver, Leech, Dónal Haltrich, Dietmar, Gorton, Lo. (2018). Amperometric Flow Injection Analysis of Glucose and Galactose Based on Engineered Pyranose 2-Oxidases and Osmium Polymers for Biosensor Applications. Electroanalysis, 30(7), 1496-1504. doi: doi:10.1002/elan.201800096
Publisher	Wiley
Link to publisher's version	https://doi.org/10.1002/elan.201800096
Item record	http://hdl.handle.net/10379/14964
DOI	http://dx.doi.org/10.1002/elan.201800096

Downloaded 2024-04-28T05:45:39Z

Some rights reserved. For more information, please see the item record link above.



# Amperometric Flow Injection Analysis of Glucose and Galactose Based on Engineered Pyranose 2-Oxidases and Os Polymers for Biosensor Applications

Sevinc Kurbanoglu<sup>a,b</sup>, Muhammed Nadeem Zafar<sup>c\*</sup>, Federico Tasca<sup>d</sup>, Iqra Aslam<sup>e</sup>, Oliver Spadiut<sup>f</sup>, Dónal Leech<sup>g</sup>, Dietmar Haltrich<sup>f</sup>, Lo Gorton<sup>b</sup>

<sup>a</sup>Department of Analytical Chemistry, Ankara University, Tandogan, Ankara, Turkey

<sup>b</sup>Department of Biochemistry and Structural Biology, Lund University, Lund, Sweden

<sup>c</sup>Department of Chemistry, University of Gujrat, Gujrat, Pakistan

<sup>d</sup>Department of Materials Chemistry, University of Santiago of Chile, Santiago, Chile

<sup>e</sup>Department of Biochemistry, Govt. College University Faisalabad, Pakistan

<sup>f</sup>Department of Food Sciences and Technology, University of Natural Resources and Life Sciences, Vienna A-1190, Austria

<sup>g</sup>School of Chemistry & Ryan Institute, National University of Ireland Galway, University Road, Galway, Ireland.

\*Corresponding author: Author to whom correspondence should be addressed; E-Mails: <a href="mailto:nadeem.zafar@uog.edu.pk">nadeem.zafar@uog.edu.pk</a> & <a href="mailto:znadeempk@gmail.com">znadeempk@gmail.com</a> and Lo.Gorton@gmail.com

#### **Abstract**

In the present study, wild type and three mutants of pyranose 2-oxidase (PyOx), which showed improved properties for D-galactose oxidation, were investigated for their oxidising ability when immobilised on graphite electrodes. Four different flexible Os polymers with formal potentials ranging between -0.140 and 0.270 V vs. Ag|AgCl<sub>0.1 M KCl</sub> were applied together with the various forms of PyOx to wire graphite electrodes using polyethylene glycol diglycidyl ether as crosslinking reagent. The pH profiles for the electrodes modified with wild type and all PyOx mutants in combination with Os polymers were investigated with both glucose and galactose respectively, since the PyOx variants showed improved catalytic activity for galactose. All modified electrodes showed highest response in the pH range between 8.5-10 and K<sub>M</sub>, I<sub>max</sub> values for the both substrates glucose and galactose were determined. To prove the catalytic activity, the biosensors were also characterized with cyclic voltammetry. The protein amount 0.26 U was found optimum for PyOx-WT, 0.36 U for PyOx-MT1, 0.41 U for PyOx-MT2 and 0.28 U for PyOx-MT3 and analytical characterization of the enzyme electrodes was performed for glucose and galactose under optimized conditions.

#### **Keywords**

Biosensor, biofuel cell anode, pyranose oxidase, osmium redox polymers, glucose, galactose.

#### 1 Introduction

Pyranose oxidase (PyOx, pyranose:oxygen 2-oxidoreductase, E.C. 1.1.3.10, glucose 2-oxidase) oxidizes glucose as well as other monosaccharides that are found as constituents of hemicelluloses including D-xylose, D-galactose, and L-arabinose, albeit with lower catalytic efficiencies. PyOx is a relatively large flavin adenine dinucleotide (FAD) containing

glycoprotein (ca. 300,000 kDa) produced by white rot fungi, with the FAD covalently attached to the polypeptide backbone [1, 2]. PyOx catalyzes a Ping Pong Bi Bi type reaction that consists of a reductive half reaction in which an aldopyranose substrate reduces the FAD cofactor to yield FADH2 and 2-dehydroaldose, as the result of oxidation of the sugar at position C-2, and the ensuing oxidative half-reaction, which involves the re-oxidation of FADH<sub>2</sub> by an electron acceptor (Figure S1) [3-5]. During this oxidative half-reaction a C-4ahydroperoxyflavin intermediate is formed when oxygen is used, the first evidence of such an intermediate for a flavoprotein oxidase [6]. PyOx can also use 2 e<sup>-</sup>, H<sup>+</sup> acceptors such as various quinones, and 1 e acceptors such as complexed metal ions and radicals as its electron acceptor [7-9]. The ability of PyOx to react with alternative electron acceptors and a range of sugar substrates can be employed in various applications [10-16]. The ability of PyOx to oxidise the sugar at the C-2 position from which follows that PyOx is active on both anomeric forms of sugars, in contrast to glucose oxidase (GOx), the most commonly studied sugaroxidising enzyme in bioelectrochemistry, and in biosensor and biofuel cell applications, which only reacts with B-D-glucose, and most other sugar oxidizing enzymes makes PyOx special but has only to a small extent thus far been a focus in bioelectrochemistry.

The crystal structure of PyOx [17, 18] provides detailed information about the residues interacting with the sugar substrate in the active site. As a consequence, site-directed mutagenesis at these residues allows analysis of structure–function relationships, and also possible improvements by semi-rational protein design. Guided by the crystal structure, saturation mutagenesis was used to exchange all of the active-site residues one by one, and screen for improvements in the catalytic activity with D-galactose and alternative electron acceptors. One of these active-site variants, V546C, was reported [19, 20] to show significantly improved catalytic properties attractive for various biotechnological applications. In homogeneous steady-state characterization, the resulting variants

S113E/T169G/H450G/Q461R/E542K/V546C) show a significant increase in activity when D-galactose is used as electron donor and either 1,4-benzoquinone or the ferricenium ion was used as solution-phase electron acceptor over the wild-type PyOx, thus showing promise for use in enzyme electrodes for determination of galactose, should the enzyme and electron acceptor be co-immobilized on electrodes [19, 20].

There has been intensive research into the use of mixed ligand transition metal complexes co-ordinated to polymers for co-immobilization of biorecognition elements and electron donors/acceptors for application to biosensor [21-23] and biofuel cell [24-31] device development. This research has focused on osmium-based redox polymers as they exhibit several advantages such as lower ionization energy, which leads to a subsequent stabilization of higher oxidation states, lower redox potentials and greater extension of the metal d orbitals, leading to enhanced metal-ligand back bonding and providing increased complex stability [21, 32, 33]. From the first applications of these redox polymers for reagentless mediated biosensing [34, 35], they attract much attention due to their efficient electron shuttling properties combined with the polymeric structure promoting a stable adsorption as well as the possibility for multiple layers of immobilized protein molecules, biological membranes or bacterial cells on the electrode surface [36-38]. So far they have been used for immobilizing many biological catalysts including sugar oxidizing enzymes glucose oxidase, oligosaccharide dehydrogenase, galactose oxidase, PyOx, pyranose dehydrogenase, PQQ and FAD glucose dehydrogenase, cellobiose dehydrogenase, fructose dehydrogenase etc. [39-54], various peroxidases [55-57], multicopper blue oxidases such as laccase and bilirubin oxidase [28, 58-63] and whole bacterial cells [64-67] and biological membranes [68, 69].

Galactose is of considerable importance to the human organism, contributing directly to vital information and control processes in the body [70]. It also functions as a fundamental

and structural substance for cells, cell walls, and intracellular matrix. The quantitative

determination of galactose is thus of great importance in clinical chemistry, food and

fermentation industries [70-72]. In this research, in order to develop enzyme electrodes with

improved sensitivity to galactose, the activity of wild type pyranose oxidase (PyOx-WT) and

that of three different pyranose oxidase mutants (PyOx-MTs) were examined by wiring with

four different redox polymers for use in enzyme electrodes for determination of galactose.

2. Experimental

2.1 Reagents

Glucose and galactose were purchased from Sigma-Aldrich (St. Louis, MO USA).

Polyethyleneglycol diglycidyl ether (PEGDGE) was purchased from Polysciences

(Warrington, PA, USA). Redox polymers 1-3 were provided from the School of Chemistry,

National University of Ireland Galway and redox polymer 0 was synthesized according to a

previously published protocol [74]. The structures of the redox polymers are presented in

Figure S2. The PyOx MTs, with improved activity for galactose, were obtained according to a

previously published protocol [20]. PyOx with active-site variant are as follows:

PyOx MT-1: T169G/H450G/E542K/V546C,

PyOx MT-2: S113E/T169G/H450G/Q461R/V546C,

PyOx MT-3: S113E/T169G/H450G/Q461R/E542K/V546C.

2.2 Instrumentation

Electrochemical studies were performed in an amperometric wall-jet cell connected to

a flow injection analysis (FIA) system [75]. The wall-jet electrochemical flow-through cell

contains three electrodes, a working electrode made of graphite rods (type RW001, 3.05 mm

diameter, Ringdorf Werke GmbH, Bonn, Germany), a reference electrode Ag|AgCl<sub>0.1 M KCl</sub>

5

and a counter electrode made of a platinum wire. An injector (Rheodyne, Cotati, CA, USA), lab balances (Sartorius, Göttingen, Germany), a peristaltic pump (Gilson, Villier-le-Bel, France), pH meter (Metrohm 744, Metrohm Filderstadt, Germany), pipettes (Eppendorf Hamburg, Germany), a potentiostat (Zäta Elektronik, Höör, Sweden), a strip chart recorder (Kipp & Zonen, Delft, The Netherlands) were used in experiments. For cyclic voltammetry studies, an electrochemical analyzer BAS 100A (Bioanalytical Systems, West Lafayette, IN, USA) was used with saturated calomel electrode (SCE) as reference electrode and platinum foil as counter electrode. A 50 mM Tris buffer of pH 8.5 was used as electrolyte and it was thoroughly out-gassed with argon prior to experiments to maintain an inert atmosphere.

# 2.3 Preparation of the PyOx/redox polymer modified electrodes

Graphite rods were cut in approximately 6 cm long pieces and polished on wet emery paper (P2000) to create a flat and smooth surface. After rinsing them thoroughly with Milli-Q water, they were dried at room temperature. Aliquots of PyOx WT or PyOx MTs were drop-cast on the top of the polished end of the graphite electrode using a Hamilton syringe. After about 15-20 min, an aliquot of 2 μL of redox polymer solution (10 mg.mL<sup>-1</sup> in Milli-Q water) was mixed with the remaining drop on the active surface of the electrodes using the micro syringe tip. Finally, 1 μL of a freshly prepared PEGDGE solution (10 mg.mL<sup>-1</sup> in Milli-Q water) was spread on the top of the electrodes and the electrodes were left overnight at 4°C at constant humidity for complete cross-linking reaction. Before using the electrodes in the FIA system, electrodes were intensively rinsed with Milli-Q water in order to remove any weakly attached components. The PyOx modified graphite electrode was fitted into a Teflon holder and inserted into a flow-through amperometric cell of the wall-jet type and kept at a constant distance (ca. 3 mm) from the carrier solution inlet port. The wall-jet cell was connected online to a flow injection analysis system, in which the carrier buffer flow was maintained at a

constant flow rate of 1 mL min<sup>-1</sup> with a peristaltic pump. An injector consisting of an electrically controlled six-port injection valve and a 50 µL injection loop was used. The output signal was recorded on a strip chart recorder. All measurements were performed at room temperature. Every injection was repeated three times and at least 2 electrodes of the same type were tested to check the reproducibility.

#### 3. Results and discussions

# 3.1 Optimization of applied potentials, comparison of Os polymers towards PyOx catalytic activities and comparison of PyOx-MT towards glucose and galactose catalysis.

The formal potentials (E°') of the Os(II/III) transition for the redox polymers depends on the co-ordination sphere in the metal-complex structure (see Fig. S1). To obtain best response and to facilitate the transfer of electrons from reduced PyOx<sub>FAD</sub> to the redox polymer and then from the redox polymer to electrodes, it is important to establish the E°'-value of each polymer. A 6 mM glucose solution was used as substrate solution and the applied potential was varied stepwise over a potential range and hydrodynamic amperometric response was measured for each applied potential. The E°'-values is estimated as the potential of half-maximum current in the hydrodynamic amperometric response and the applied potentials for all redox polymers was selected to be a potential where maximum current is achieved (between 80 to 140 mV more positive of the formal potential) as presented in Table 1.

To compare the effect of the redox polymer choice on the response to the sugar substrates, four different redox polymers were investigated. The analytical response to glucose obtained from the electrodes modified with the various combinations of PyOx/redox is shown in Figure 1A. The bound FAD of PyOx-WT has an E°'-value of approximately -150 mV [17, 18]. Therefore, redox polymers with an E°' higher than -150 mV are expected to

accept electrons from the reduced enzyme active site and further transfer it to the electrode. As revealed in Fig. 1A the response to glucose for PyOx with redox polymers 0 and 1 is low as a result of the low thermodynamic driving force for electron transfer to these redox polymers in their oxidized states to reoxidize reduced PyOx, whereas redox polymers 2 and 3 deliver current to the electrode as a result of oxidation of glucose by PyOx and electron transfer through the redox polymers to the electrode. When the PyOx-MTs were wired with Os polymer 0 and 1, no detectable responses to glucose were obtained. This may be caused by the unfavourable kinetics of these variants for the substrate glucose (see below). Heller suggested that the E° of the mediator should be about 50 mV more positive than that of the enzyme for optimal performance [76], a suggestion supported by the results in Figure 1A. From Figure 1A, it is clear that for all PyOx variants co-immobilized with redox polymer 3 the highest response is obtained compared to that for all other redox polymers. To compare the catalytic response of PyOx-MTs towards galactose and glucose, the electrodes were therefore prepared by co-immobilizing the PyOx-MTs and redox polymer 3. The results in Figure 1B show that all the mutants have higher catalytic response for galactose compared to that obtained with PyOx-WT, which shows a relatively high response toward glucose. The mutants were engineered to enhance the response for galactose [19, 20] and these results show that the engineering and mutation of the PyOx enzyme were successful in improving response to galactose when the enzymes are co-immobilized with redox polymers as electron acceptors, to provide enzyme electrode biosensors.

### 3.2 Cyclic voltammetry

Cyclic voltammetry (CV) was performed to verify activity of the enzymes immobilized on the graphite electrode. To observe any interaction between the osmium redox polymers and PyOx, CVs were recorded for electrodes made with redox polymer in the presence and absence of PyOx enzyme at a scan rate of 1 mV.s<sup>-1</sup>. The results presented

in Figure 2A show the voltammogram for an electrode modified redox polymer 3 alone compared to that for an electrode modified with redox polymer 3 and PyOx-MT1. It was observed that when redox polymer 3 was co-immobilized with PyOx-MT1, the CV was slightly shifted into a more negative direction and a decrease in the peak current occurs, possibly reflecting interaction between redox polymer and enzyme, as a similar response was observed in our previous studies where pyranose dehydrogenase was co-immobilized with redox polymers [41]. Similar results were also obtained for all of the other osmium polymers (results not shown). The results presented in Figure 2B again show that when PyOx-MT1 is co-immobilized with redox polymer 3, the catalytic activity towards galactose was higher than that for glucose.

## 3.3 pH profile

To study the effect of pH on the activity of enzymes, the pH values of the running buffer (50 mM Tris buffer between pH 7.0 and 8.5 and 50 mM ethanolamine buffer between pH 9.0 and 11.0) was varied. The buffers were thoroughly out-gassed under vacuum before use to prevent the appearance of bubbles in the FIA system. Glucose and galactose solutions (6 mM) were prepared in an appropriate buffer and injected (50 μL) into the carrier stream. The results in Figure 3A again demonstrate that the highest current response was obtained when PyOx-WT was mediated by redox polymer 3, and was so for all pH values. As already shown in Figure 1A, the PyOx-MTs responded better in terms of output current when wired with Os polymer 2 and 3 using galactose as substrate so the pH profiles were only studied with Os polymer 2 and 3. There is a sharp increase in current response around pH 8.5 for PyOx-WT and PyOx-MTs, which increased until pH 10. Even though the pH of 10.0 in 50 mM ethanolamine buffer was found to be the optimum for maximum current response this buffer system could not be used for characterization because exposing the electrode for a long time at such an alkaline pH would lead to denaturation and inactivation of the enzyme and

additionally the sugars may get directly oxidized more easily at high pHs. Therefore, 50 mM, Tris buffer with pH 8.5 was selected as optimal operating medium. Moreover, the experimental data showed that when redox polymer 3 was used as mediator, higher current values could be obtained over the whole studied pH range.

### 3.4 Optimization of enzyme amount

The effect of the amount of enzyme on the electrode response was determined using redox polymer 3. Different amounts of PyOx-WT and PyOx-MTs were used to modify the graphite electrodes along with redox polymer 3. These graphite electrodes were tested when used as working electrodes in the FIA system with 6 mM galactose for PyOx-MTs and 6 mM glucose for PyOx-WT injections (Figure 4). The optimum amount of protein immobilized on the electrode was found to be 3  $\mu$ L (0.26 U) for PyOx-WT, 2  $\mu$ L (0.36 U) for PyOx-MT-1, 4  $\mu$ L (0.41 U) for PyOx-MT-2 and 2  $\mu$ L (0.28 U) for PyOx-MT-3. The optimum enzyme amount was used for preparation of electrodes with PyOx-WT and PyOx-MTs for further studies.

## 3.5 Analytical performance

The calibration curves for galactose and glucose with redox polymer 3 immobilized with PyOx-MT1, PyOx-MT2 and PyOx-MT3 on graphite electrodes were recorded in Tris buffer at pH 8.5. The enzyme electrodes were placed in the cell, and various concentrations of galactose and glucose dissolved in Tris buffer (pH 8.5) were injected into the flow system. The results of the calibration curves obtained using the biosensors based on redox polymer 3 immobilized with PyOx-MT1, PyOx-MT2 and PyOx-MT3 are presented in Figure 5 and Table 2 showed the linear ranges of the calibration curves for all types of the enzymemodified electrode. The linear ranges for PyOx-MT1 was 0.1–8 mM galactose whereas PyOx-MT2 and PyOx-MT3 showed similar linear range (0.1–10 mM galactose). The detection limits for galactose were 2.9, 3.2 and 3.4 μM for PyOx-MT1, PyOx-MT2 and

PyOx-MT3 respectively. The detection limits of PyOx-MT1, PyOx-MT2 and PyOx-MT3 for glucose were 4.5, 9.0 and 8.5 μM respectively which were higher than that for galactose. The results reveal that all the mutants not only showed improved catalytic resoponse for galactose but also lower detection limits. All developed biosensors showed good reproducibility of the analytical response.

The parameter,  $K_M$  is a measure of the affinity of an enzyme for its substrate, which is inversely proportional to the affinity, and  $I_{max}$  is the maximum rate of the enzyme reaction.  $K_M$  can also altered due to diffusional limitations of substrate or electron transport to the electrode surface.  $K_M$  and  $I_{max}$  can be calculated for immobilized enzymes by fitting the experimental data into an electrochemical Michaelis-Menten plot (current vs. substrate concentration) [77]. The parameters  $K_M$  and  $I_{max}$  were determined for glucose and galactose from the calibration curves and reported in Table 2. The results show that there was no significant changes in the  $K_M$  values of PyOx-WT and PyOx-MTs for glucose. In contrast, the  $K_M$  values of PyOx-MTs are significantly lower for galactose, compared to that for PyOx-WT which indicates that the substrate affinity of the mutants for galactose is better than that of PyOx-WT.

### 4. Conclusions

In homogeneous steady-state characterization, the resulting mutants (T169G/H450G/E542K /V546C/S113E/T169G/H450G/Q461R/V546C and S113E/T169G/H450G/Q461R/E542K/ V546C) showed a significant increase in activity when D-galactose was used as electron donor and Os polymers were used as electron acceptor. The catalytic efficiency increased up to 30-fold. The PyOx-MTs, which showed improved activity for D-galactose as a substrate were used for biosensing of galactose. They were coimmobilized with different redox polymers with formal potentials ranging between -0.140 and 0.270 V. The PyOx-WT and the three PyOx-MTs were characterized by their pH profile, substrate specificity and enzyme loading experiments. The parameters of the biosensing system,  $I_{max}$  and  $K_{M}$ , for glucose and galactose were also determined. With the results it can be concluded that these optimized PyOx-based biosensors mediated using osmium based redox polymers can be used as bioanodes for biosensor and biofuel cell studies.

# Acknowledgements

The authors thank the following agencies for financial support: **MN Zafar** The Higher Education Commission of Pakistan for financial support (grant number: 6515/Punjab/NRPU/R&D/HEC/2016), **LG** The Swedish Research Council (grant number 2014-5908), **DL** and **LG** The European Commission (grant number: FP7-PEOPLE-2013-ITN-607793).

#### References

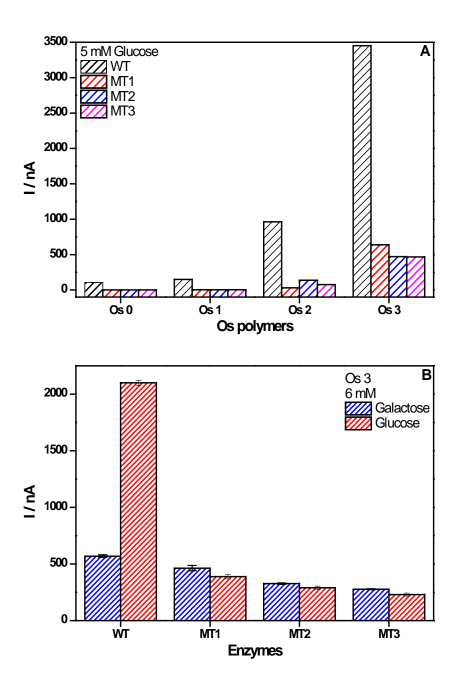
- [1] G. Daniel, J. Volc, E. Kubatova, Pyranose oxidase, a major source of H<sub>2</sub>O<sub>2</sub> during wood degradation by *Phanerochaete-chrysosporium*, *Trametes-versicolor*, and *Oudemansiella-mucida*, Appl. Environ. Microbiol., 60 (1994) 2524-2532.
- [2] P. Halada, C. Leitner, P. Sedmera, D. Haltrich, J. Volc, Identification of the covalent flavin adenine dinucleotide-binding region in pyranose 2-oxidase from *Trametes multicolor*, Anal. Biochem., 314 (2003) 235-242.
- [3] S. Freimund, A. Huwig, F. Giffhorn, S. Kopper, Rare keto-aldoses from enzymatic oxidation: Substrates and oxidation products of pyranose 2-oxidase, Chem. Eur. J., 4 (1998) 2442-2455.
- [4] M.J. Artolozaga, E. Kubatova, J. Volc, H.M. Kalisz, Pyranose 2-oxidase from *Phanerochaete chrysosporium* Further biochemical characterisation, Appl. Microbiol. Biotechnol., 47 (1997) 508-514.
- [5] S. Ghisla, V. Massey, Mechanisms of flavoprotein-catalyzed reactions, Eur. J. Biochem., 181 (1989) 1-17.
- [6] J. Sucharitakul, M. Prongjit, D. Haltrich, P. Chaiyen, Detection of a C4a-hydroperoxyflavin intermediate in the reaction of a flavoprotein oxidase, Biochemistry, 47 (2008) 8485-8490.
- [7] C. Leitner, J. Volc, D. Haltrich, Purification and characterization of pyranose oxidase from the white rot fungus *Trametes multicolor*, Appl. Environ. Microbiol., 67 (2001) 3636-3644.
- [8] F.W. Janssen, H.W. Ruelius, Pyranose oxidase from *Polyporus obtusus*, Meth. Enzymol., 41 (1975) 170-173.
- [9] U. Baminger, R. Ludwig, C. Galhaup, C. Leitner, K.D. Kulbe, D. Haltrich, Continuous enzymatic regeneration of redox mediators used in biotransformation reactions employing flavoproteins, J. Mol. Catal. B, 11 (2001) 541-550.
- [10] F.W. Janssen, H.W. Ruelius, Carbohydrate oxidase a novel enzyme from *Polyporus obtusus* .2. Specificity and characterization of reaction products, Biochim. Biophys. Acta, 167 (1968) 501-&.
- [11] L. Olsson, C.F. Mandenius, Determination of monosaccharides in cellulosic hydrolyzates using immobilized pyranose oxidase in a continuous amperometric analyzer, Anal. Chem., 62 (1990) 2688-2691.
- [12] V. Pacheco, A. Karmali, Chromatographic behaviour of glucose 1- and 2-oxidases from fungal strains on immobilized metal chelates, J. Ind. Microbiol. Biotechnol., 21 (1998) 57-64.
- [13] F. Giffhorn, S. Kopper, A. Huwig, S. Freimund, Rare sugars and sugar-based synthons by chemo-enzymatic synthesis, Enzyme Microb. Technol., 27 (2000) 734-742.
- [14] S. Timur, Y. Yigzaw, L. Gorton, Electrical wiring of pyranose oxidase with osmium redox polymers, Sens. Actuat. B-Chem., 113 (2006) 684-691.
- [15] B. Weigel, B. Hitzmann, G. Kretzmer, K. Schügerl, A. Huwig, F. Giffhorn, Analysis of various sugars by means of immobilized enzyme coupled flow injection analysis, J. Biotechnol., 50 (1996) 93-106.
- [16] N. Namba, F. Watanabe, M. Tokuda, M. Mino, E. Furuya, A new method of quantitating serum and urinary levels of 1,5-anhydroglucitol in insulin-dependent diabetes-mellitus, Diabetes Res. Clinical Pr., 24 (1994) 55-61.
- [17] B.M. Hallberg, C. Leitner, D. Haltrich, C. Divne, Crystal structure of the 270 kDa homotetrameric lignin-degrading enzyme pyranose 2-oxidase, J. Mol. Biol., 341 (2004) 781-796.

- [18] M. Kujawa, H. Ebner, C. Leitner, B.M. Hallberg, M. Prongjit, J. Sucharitakul, R. Ludwig, U. Rudsander, C. Peterbauer, P. Chaiyen, D. Haltrich, C. Divne, Structural basis for substrate binding and regioselective oxidation of monosaccharides at C3 by pyranose 2-oxidase, J. Biol. Chem., 281 (2006) 35104-35115.
- [19] O. Spadiut, D. Brugger, V. Coman, D. Haltrich, L. Gorton, Engineered pyranose 2-oxidase: Efficiently turning sugars into electrical energy, Electroanalysis, 22 (2010) 813-820.
- [20] O. Spadiut, I. Pisanelli, T. Maischberger, C. Peterbauer, L. Gorton, P. Chaiyen, D. Haltrich, Engineering of pyranose 2-oxidase: Improvement for biofuel cell and food applications through semi-rational protein design, J. Biotechnol., 139 (2009) 250-257.
- [21] P. Kavanagh, D. Leech, Redox polymer and probe DNA tethered to gold electrodes for enzyme-amplified amperometric detection of DNA hybridization, Anal. Chem., 78 (2006) 2710-2716.
- [22] S. Boland, F. Barrière, D. Leech, Designing stable redox-active surfaces: Chemical attachment of an osmium complex to glassy carbon electrodes prefunctionalized by electrochemical reduction of an in situ-generated aryldiazonium cation, Langmuir, 24 (2008) 6351-6358.
- [23] M. Pellissier, F. Barrière, A.J. Downard, D. Leech, Improved stability of redox enzyme layers on glassy carbon electrodes via covalent grafting, Electrochem. Commun., 10 (2008) 835-838.
- [24] F. Gao, Y. Yan, L. Su, L. Wang, L. Mao, An enzymatic glucose/O<sub>2</sub> biofuel cell: Preparation, characterization and performance in serum, Electrochem. Commun., 9 (2007) 989-996.
- [25] N. Mano, F. Mao, A. Heller, Characteristics of a miniature compartment-less glucose-O<sub>2</sub> biofuel cell and its operation in a living plant, J. Am. Chem. Soc., 125 (2003) 6588-6594.
- [26] N. Mano, F. Mao, A. Heller, A miniature membrane-less biofuel cell operating at +0.60 V under physiological conditions, ChemBioChem, 5 (2004) 1703-1705.
- [27] E. Katz, I. Willner, A.B. Kotlyar, A non-compartmentalized glucose vertical bar  $O_2$  biofuel cell by bioengineered electrode surfaces, J. Electroanal. Chem., 479 (1999) 64-68.
- [28] D. Leech, P. Kavanagh, W. Schuhmann, Enzymatic fuel cells: Recent progress, Electrochim. Acta, 84 (2012) 223-234.
- [29] M. Falk, M. Alcalde, P.N. Bartlett, A.L. De Lacey, L. Gorton, C. Gutierrez-Sanchez, R. Haddad, J. Kilburn, D. Leech, R. Ludwig, E. Magner, D.M. Mate, P. Ò Conghaile, R. Ortiz, M. Pita, S. Pöller, T. Ruzgas, U. Salaj-Kosla, W. Schuhmann, F. Sebelius, M. Shao, L. Stoica, C. Sygmund, J. Tilly, M.D. Toscano, J. Vivekananthan, E. Wright, S. Shleev, Self-powered wireless carbohydrate/oxygen sensitive biodevice based on radio signal transmission, Plos One, 9 (2014).
- [30] M. Shao, M.N. Zafar, M. Falk, R. Ludwig, C. Sygmund, C.K. Peterbauer, D.A. Guschin, D. MacAodha, P. Ò Conghaile, D. Leech, M.D. Toscano, S. Shleev, W. Schuhmann, L. Gorton, Optimization of a membraneless glucose/oxygen enzymatic fuel cell based on a bioanode with high coulombic efficiency and current density, ChemPhysChem, 14 (2013) 2260-2269.
- [31] P. O Conghaile, M. Falk, D. MacAodha, M.E. Yakovleva, C. Gonaus, C.K. Peterbauer, L. Gorton, S. Shleev, D. Leech, Fully enzymatic membraneless glucoseloxygen fuel cell that provides 0.275 mA cm(-2) in 5 mM glucose, operates in human physiological solutions, and powers transmission of sensing data, Anal. Chem., 88 (2016) 2156-2163.
- [32] T. Maddanimath, Y.B. Khollam, A. Aslam, I.S. Mulla, K. Vijayamohanan, Self-assembled monolayers of diphenyl disulphide: a novel cathode material for rechargeable lithium batteries, J. Power Sources, 124 (2003) 133-142.

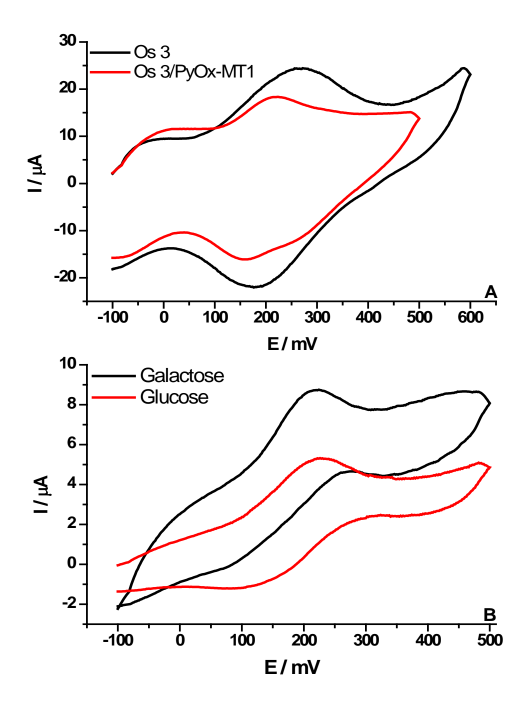
- [33] C.H. Park, Y.K. Sun, D.W. Kim, Blended polymer electrolytes based on poly(lithium 4-styrene sulfonate) for the rechargeable lithium polymer batteries, Electrochim. Acta, 50 (2004) 375-378.
- [34] Y. Degani, A. Heller, Electrical communication between redox centers of glucose-oxidase and electrodes via electrostatically and covalently bound redox polymers, J. Am. Chem. Soc., 111 (1989) 2357-2358.
- [35] A. Heller, Electrical connection of enzyme redox centers to electrodes, J. Phys. Chem., 96 (1992) 3579-3587.
- [36] T.J. Ohara, R. Rajagopalan, A. Heller, Glucose electrodes based on cross-linked Os(bpy)<sub>2</sub> (+/2+) complexed poly(l-vinylimidazole) films, Anal. Chem., 65 (1993) 3512-3517.
- [37] T.J. Ohara, R. Rajagopalan, A. Heller, Wired enzyme electrodes for amperometric determination of glucose or lactate in the presence of interfering substances, Anal. Chem., 66 (1994) 2451-2457.
- [38] M. Kirthiga, L. Rajendran, C. Fernandez, Theoretical treatment of diffusion and kinetics of osmium redox polymer mediated glucose oxidase enzyme electrodes: Analytical expression of current density for varying potential, Electrochim. Acta, 230 (2017) 89-97.
- [39] M. Shao, M.N. Zafar, C. Sygmund, D.A. Guschin, R. Ludwig, C.K. Peterbauer, W. Schuhmann, L. Gorton, Mutual enhancement of the current density and the coulombic efficiency for a bioanode by entrapping bi-enzymes with Os-complex modified electrodeposition paints, Biosens. Bioelectron., 40 (2013) 308-314.
- [40] M.N. Zafar, N. Beden, D. Leech, C. Sygmund, R. Ludwig, L. Gorton, Characterization of different FAD-dependent glucose dehydrogenases for possible use in glucose-based biosensors and biofuel cells, Anal. Bioanal. Chem., 402 (2012) 2069-2077.
- [41] M.N. Zafar, F. Tasca, S. Boland, M. Kujawa, I. Patel, C.K. Peterbauer, D. Leech, L. Gorton, Wiring of pyranose dehydrogenase with osmium polymers of different redox potentials, Bioelectrochemistry, 80 (2010) 38-42.
- [42] M.N. Zafar, X. Wang, C. Sygmund, R. Ludwig, D. Leech, L. Gorton, Electron-transfer studies with a new flavin adenine dinucleotide dependent glucose dehydrogenase and osmium polymers of different redox potentials, Anal. Chem., 84 (2012) 334-341.
- [43] F. Tasca, S. Timur, R. Ludwig, D. Haltrich, J. Volc, R. Antiochia, L. Gorton, Amperometric biosensors for detection of sugars based on the electrical wiring of different pyranose oxidases and pyranose dehydrogenases with osmium redox polymer on graphite electrodes, Electroanalysis, 19 (2007) 294-302.
- [44] R. Antiochia, F. Mazzei, L. Gorton, D. Leech, G. Favero, Composite material based on macroporous polyaniline and osmium redox complex for biosensor development, Electroanalysis, 26 (2014) 1623-1630.
- [45] R. Antiochia, L. Gorton, Development of a carbon nanotube paste electrode osmium polymer-mediated biosensor for determination of glucose in alcoholic beverages, Biosens. Bioelectron., 22 (2007) 2611-2617.
- [46] R. Antiochia, L. Gorton, A new osmium-polymer modified screen-printed graphene electrode for fructose detection, Sens. Actuat. B-Chem., 195 (2014) 287-293.
- [47] R. Antiochia, G. Vinci, L. Gorton, Rapid and direct determination of fructose in food: A new osmium-polymer mediated biosensor, Food Chem, 140 (2013) 742-747.
- [48] R. Ludwig, W. Harreither, F. Tasca, L. Gorton, Cellobiose dehydrogenase: A versatile catalyst for electrochemical applications, ChemPhysChem, 11 (2010) 2674-2697.
- [49] F. Tasca, L. Gorton, W. Harreither, D. Haltrich, R. Ludwig, G. Nöll, Comparison of direct and mediated electron transfer for cellobiose dehydrogenase from *Phanerochaete soridida*, Anal. Chem., 81 (2009) 2791-2798.

- [50] F. Tasca, L. Gorton, M. Kujawa, I. Patel, W. Harreither, C.K. Peterbauer, R. Ludwig, G. Nöll, Increasing the coulombic efficiency of glucose biofuel cell anodes by combination of redox enzymes, Biosens. Bioelectron., 25 (2010) 1710-1716.
- [51] M. Tessema, E. Csöregi, T. Ruzgas, G. Kenausis, T. Solomon, L. Gorton, Oligosaccharide dehydrogenase-modified graphite electrodes for the amperometric determination of sugars in a flow injection system, Anal. Chem., 69 (1997) 4039-4044.
- [52] M. Tessema, T. Larsson, T. Buttler, E. Csöregi, T. Ruzgas, M. Nordling, S.E. Lindquist, G. Pettersson, L. Gorton, Simultaneous amperometric determination of some mono-, di-, and oligosaccharides in flow injection and liquid chromatography using two working enzyme electrodes with different selectivity, Anal. Chim. Acta, 349 (1997) 179-188.
- [53] Y. Ling, M. Hämmerle, A.J.J. Olsthoorn, W. Schuhmann, H.L. Schmidt, J.A. Duine, A. Heller, High-current density wired quinoprotein glucose-dehydrogenase electrode, Anal. Chem., 65 (1993) 238-241.
- [54] W. Schuhmann, T.J. Ohara, H.L. Schmidt, A. Heller, Electron-transfer between glucose-oxidase and electrodes via redox mediators bound with flexible chains to the enzyme surface, J. Am. Chem. Soc., 113 (1991) 1394-1397.
- [55] P. Bollella, L. Medici, M. Tessema, A.A. Poloznikov, D.M. Hushpulian, V.I. Tishkov, R. Andreu, D. Leech, N. Megersa, M. Marcaccio, L. Gorton, R. Antiochia, Highly sensitive, stable and selective hydrogen peroxide amperometric biosensors based on peroxidases from different sources wired by Os-polymer: A comparative study, Solid State Ionics, 314 (2018) 178-186.
- [56] M. Vreeke, R. Maidan, A. Heller, Hydrogen-peroxide and β-nicotinamide adenine-dinucleotide sensing amperometric electrodes based on electrical connection of horseradish-peroxidase redox centers to electrodes through a 3-dimensional electron relaying polymer network, Anal. Chem., 64 (1992) 3084-3090.
- [57] A. Belay, A. Collins, T. Ruzgas, P.T. Kissinger, L. Gorton, E. Csöregi, Redox hydrogel based bienzyme electrode for L-glutamate monitoring, J. Pharmaceut. Biomed. Anal., 19 (1999) 93-105.
- [58] F. Trudeau, F. Daigle, D. Leech, Reagentless mediated laccase electrode for the detection of enzyme modulators, Anal. Chem., 69 (1997) 882-886.
- [59] F. Barrière, Y. Ferry, D. Rochefort, D. Leech, Targetting redox polymers as mediators for laccase oxygen reduction in a membrane-less biofuel cell, Electrochem. Commun., 6 (2004) 237-241.
- [60] N. Mano, H.H. Kim, Y.C. Zhang, A. Heller, An oxygen cathode operating in a physiological solution, J. Am. Chem. Soc., 124 (2002) 6480-6486.
- [61] N. Mano, V. Soukharev, A. Heller, A laccase-wiring redox hydrogel for efficient catalysis of  $O_2$  electroreduction, J. Phys. Chem. B, 110 (2006) 11180-11187.
- [62] V. Soukharev, N. Mano, A. Heller, A four-electron  $O_2$ -electroreduction biocatalyst superior to platinum and a biofuel cell operating at 0.88 V, J. Am. Chem. Soc., 126 (2004) 8368-8369.
- [63] A. Heller, Electron-conducting redox hydrogels: design, characteristics and synthesis, Curr. Opin. Chem. Biol., 10 (2006) 664-672.
- [64] G. Pankratova, L. Gorton, Electrochemical communication between living cells and conductive surfaces, Curr. Opin. Electrochem., 5 (2017) 193-202.
- [65] Y. Yuan, H. Shin, C. Kang, S. Kim, Wiring microbial biofilms to the electrode by osmium redox polymer for the performance enhancement of microbial fuel cells, Bioelectrochemistry, 108 (2016) 8-12.
- [66] S. Aslan, P. O Conghaile, D. Leech, L. Gorton, S. Timur, U. Anik, Development of an osmium redox polymer mediated bioanode and examination of its performance in *Gluconobacter oxydans* based microbial fuel cell, Electroanalysis, 29 (2017) 1651-1657.

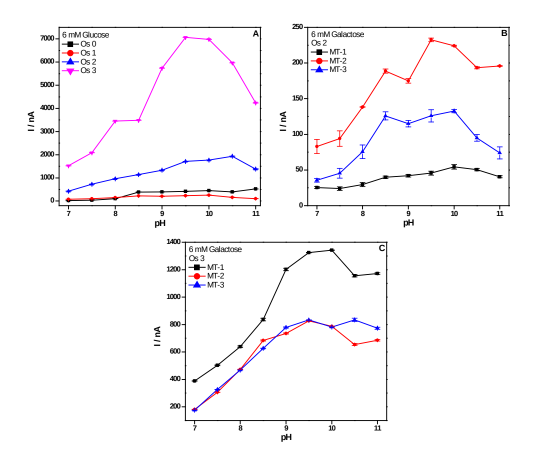
- [67] S. Aslan, P. Ò Conghaile, D. Leech, L. Gorton, S. Timur, U. Anik, Development of a bioanode for microbial fuel cells based on the combination of a MWCNT-Au-Pt hybrid nanomaterial, an osmium redox polymer and *Gluconobacter oxydans* DSM 2343 Cells, Chem. Select, 2 (2017) 12034-12040.
- [68] G. Pankratova, D. Pankratov, K. Hasan, H.-E. Akerlund, P.-A. Albertsson, D. Leech, S. Shleev, L. Gorton, Supercapacitive photo-bioanodes and biosolar cells: A novel approach for solar energy harnessing, Adv. Energy Mater., 7 (2017).
- [69] H. Hamidi, K. Hasan, S.C. Emek, Y. Dilgin, H.-E. Åkerlund, P.-A. Albertsson, D. Leech, L. Gorton, Photocurrent generation from thylakoid membranes on osmium-redox-polymer-modified electrodes, ChemSusChem, 8 (2015) 990-993.
- [70] H.J. Danneel, M. Ullrich, F. Giffhorn, Goal-oriented screening method for carbohydrate oxidases produced by filamentous fungi, Enzyme Microb. Technol., 14 (1992) 898-903.
- [71] O. Spadiut, T.-T. Nguyen, D. Haltrich, Thermostable variants of pyranose 2-oxidase showing altered substrate selectivity for glucose and galactose, J. Agric. Food Chem., 58 (2010) 3465-3471.
- [72] P. Kanyong, F.D. Krampa, Y. Aniweh, G.A. Awandare, Enzyme-based amperometric galactose biosensors: a review, Microchim. Acta, 184 (2017) 3663-3671.
- [73] S. Zhang, C. Li, G. Zhou, G. Che, J. You, Y. Suo, Determination of the carbohydrates from *Notopterygium forbesii* Boiss by HPLC with fluorescence detection, Carbohyd. Polym., 97 (2013) 794-799.
- [74] F. Mao, N. Mano, A. Heller, Long tethers binding redox centers to polymer backbones enhance electron transport in enzyme "wiring" hydrogels, J. Am. Chem. Soc., 125 (2003) 4951-4957.
- [75] R. Appelqvist, G. Markovarga, L. Gorton, A. Torstensson, G. Johansson, Enzymatic determination of glucose in a flow system by catalytic-oxidation of the nicotinamide coenzyme at a modified electrode, Anal. Chim. Acta, 169 (1985) 237-247.
- [76] A. Heller, Miniature biofuel cells, Phys. Chem. Chem. Phys., 6 (2004) 209-216.
- [77] L. Michaelis, M.L. Menten, K.A. Johnson, R.S. Goody, The original Michaelis constant: translation of the 1913 Michaelis-Menten paper, Biochemistry, 50 (2011) 8264-8269.



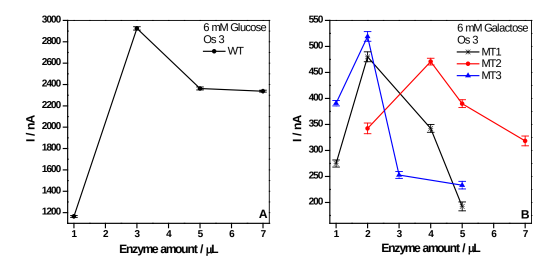
**Figure 1.** Comparison of Os polymers towards PyOx-WT and PyOx-MTs catalytic activities (A) and comparison of PyOx-MTs for glucose and galactose catalysis (B). Experiments were performed in 50 mM Tris buffer at pH 8.



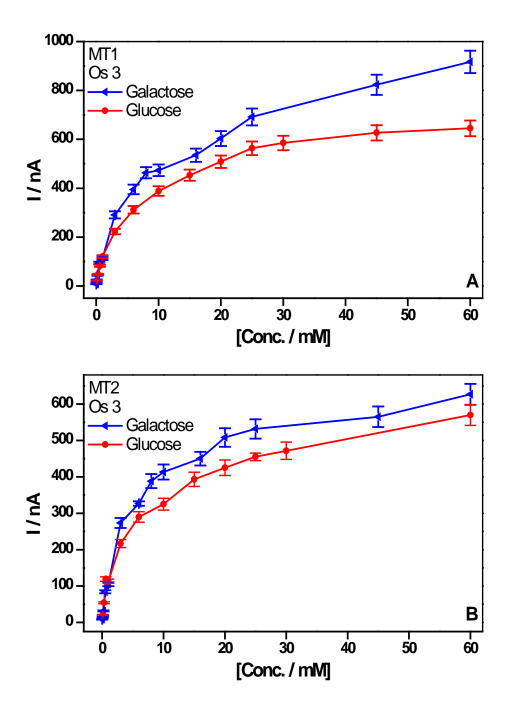
**Figure 2.** Cyclic voltammograms of (A) Os polymer 3 modified electrode (black line) and PyOx-MT1/Os polymer 3 modified electrode (red line) in the absence of substrate and (B) PyOX-MT1/Os polymer modified electrodes in the presence of galactose (black line) and glucose (red line). Experiments were performed in 50 mM Tris buffer at pH 8.5 at scan rate of 1 mV s<sup>-1</sup>.



**Figure 3.** pH profiles for (A) PyOx WT mediated with redox polymer 0, 1, 2, 3 (B) PyOx MTs wired with Os polymer 2 and (C) PyOx MTs wired with Os polymer 3



**Figure 4.** Effect of enzyme loading on the catalysis of (A) glucose on PyOx WT/Os 3 modifies electrodes and (B) galactose on PyOx MTs/Os 3 modifies electrodes.



**Figure 5.** Calibration curves of PyOx MTs wired with Os polymer 3 on graphite electrodes, obtained by measuring MET using D-glucose and D-galactose in 50 mM Tris buffer at pH 8.5 (A) PyOx MT-2 (B) PyOx MT-3

Table 1. Redox polymer formal and applied potentials vs.  $Ag|AgCl_{0.1\,M\;KCl}$ 

Os polymers	Formal Potential/(mV)	Applied Potential/(mV)
Os 0	-185	-80.0
Os 1	-140	0.00
Os 2	180	320
Os 3	270	350

 $\textbf{Table 2.} \ I_{max}, \ K_M, \ LOD, \ LOQ \ and \ linear \ ranges \ for \ PyOx \ enzymes \ using \ galactose \ and \ glucose \ as \ substrates.$ 

Enzyme	$I_{max}/(nA)$	K <sub>M</sub> /(mM)	LOD/(µM)	LOQ/(µM)	Linear range/(mM)		
			Galactos	se			
PyOx-MT1	1249.8	17.39	2.9	9.0	0.1–08		
PyOx-MT2	667.3	5.51	3.2	9.8	0.1–10		
PyOx-MT3	1031.6	6.42	3.4	10	0.1–10		
PyOx-WT	819.9	31.75					
		Glucose					
PyOx-MT1	719.3	7.53	4.5	14	0.1–10		
PyOx-MT2	613.3	7.03	9.0	27	0.1–15		
PyOx-MT3	559.5	8.64	8.5	30	0.1–15		
PyOx-WT	1916.6	7.68					