Ferrocene clicked polypyrrole derivatives: effect of spacer group on electrochemical properties and post-polymerization functionalization

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ABSTRACT
In this study, novel ferrocene-functionalized N-alkyl substituted pyrrole derivatives, namely 4-ferrocenyl-1-[3-(pyrrol-1-yl)propyl]-1H-1,2,3-triazole (Py3Fc), 4-ferrocenyl-1-[4-(pyrrol-1-yl)butyl]-1H-1,2,3-triazole (Py4Fc), and 4-ferrocenyl-1-[6-(pyrrol-1-yl)hexyl]-1H-1,2,3-triazole (Py6Fc), were synthesized via click reaction and the monomers were characterized by 1H NMR, 13C NMR, FTIR, and HRMS techniques. Redox properties of the monomers were investigated by cyclic voltammetry (CV) studies. Contrary to general literature, both Py4Fc and Py6Fc were electrochemically polymerized without loss in redox activity of ferrocene group. Moreover, click chemistry was utilized in post-polymerization functionalization. For this purpose, three azide-containing polypyrroles, P(Py3N3), P(Py4N3), and P(Py6N3) were electrochemically synthesized and subjected to click reaction in the presence of ethynylferrocene. CV studies of the post-polymerization functionalized polymers revealed quasi-reversible waves, while only P(Py6-post-Fc) showed the characteristic redox behavior of both polypyrrole and ferrocene. Thus, in this study preparation of a conducting homopolymers of pyrrole having covalently bonded ferrocene units was demonstrated and effect of spacer group is investigated.

1. Introduction
Research interest in polypyrrole derivatives have increased significantly due to versatility of their applications.[1–3] Especially, in the field of biosensors polypyrrole derivatives have acquired more importance owing to their biocompatibility, low oxidation potential, high film-forming ability, etc.[4–11] Incorporation of ferrocene to polypyrrole matrices (as mediator) has been particularly examined in order to enhance response of the amperometric biosensors.[12–14] Up to date, several attempts have been made to produce polypyrrole derivatives having covalently attached ferrocene moiety.[15–18] However, synthesis of such polymers through electrochemical polymerization was found to be tricky where in most cases it was only accomplished through electrochemical copolymerization in the presence of similar heterocyclic units not having ferrocene unit.[19]

Since 2001, there has been a growing interest in click chemistry and it has made a significant impact on material and life sciences.[20] Basic features of click reactions could be summarized as simple – selective chemical reaction with high yields, tolerance to variety of solvents, lack significant byproducts. Copper-catalyzed Huisgen 1,3-dipolar cycloaddition of azides and alkynes could be considered as the flagship of click chemistry which has been widely and successfully applied in modification of surfaces,[21–23] synthesis of compounds with complex architecture.[24–26]

Taking advantage of this high yield reaction, we planned to synthesize a series of pyrrole derivatives having ferrocene moiety and investigated their electrochemical polymerization. For this purpose three novel, ferrocene-functionalized N-alkyl substituted polypyrrole derivatives, namely 4-ferrocenyl-1-[3-(pyrrol-1-yl)propyl]-1H-1,2,3-triazole (Py3Fc), 4-ferrocenyl-1-[4-(pyrrol-1-yl)butyl]-1H-1,2,3-triazole (Py4Fc), and 4-ferrocenyl-1-[6-(pyrrol-1-yl)hexyl]-1H-1,2,3-triazole (Py6Fc), were synthesized via ‘click chemistry’ and characterized by 1H NMR, 13C NMR, FTIR, and high-resolution mass spectra (HRMS) techniques. All the monomers were evaluated for their redox behavior and electrochemical syntheses of ferrocene-tethered polypyrroles were investigated. Moreover, we explored the versatility of the ‘click’ chemistry and effect of spacer group to introduce ferrocene through post-polymerization functionalization.
2. Experimental

2.1. General

Except acetonitrile (ACN), dichloromethane (DCM) and propylene carbonate (PC), all the chemicals were purchased from Aldrich or Merck Chemical as reagent grade. Unless otherwise noted, all materials were used as received. Column chromatography was carried out with SiO₂ 60 (particle size 0.040–0.063 mm, 230–400 mesh; Merck) and distilled technical solvents. ACN and DCM were distilled over calcium hydride and kept on 4 Å molecular sieves. Tetra-n-butylammonium hexafluorophosphate (TBAPF₆) and lithium perchlorate (LiClO₄) were electroanalytical grade. Ethynylferrocene [27] was synthesized according to literature.

2.2. Equipment

An Ivium stat potentiostat/galvanostat was used to supply potential during electrochemical synthesis and cyclic voltammetry (CV). A Basi MF-2062 Non-aqueous Reference Electrode Kit was used as the Ag/Ag⁺ reference electrode in ACN with 0.01 M AgNO₃ and 0.1 M TBAPF₆. Spectroelectrochemistry studies of the polymer films were performed in a UV-cuvette with three-electrodes placed in the sample compartment of a Thermo Evolution Array UV–vis spectrophotometer. For these studies Pt and Ag wires were used as the counter and reference electrodes, respectively. NMR spectra were recorded with a Bruker Avance DPX-400 spectrometer at 400 MHz for 1H NMR, to literature.

2.3. General procedure for synthesis of N-substituted pyrroles (Py-N-Br)

N-substituted pyrroles were synthesized through typical alkaline-mediated coupling reaction between pyrrole and dibromoalkanes. As a first step, pyrrole (20 mmol) and KOH (25 mmol) were mixed at 0 °C for 3 h. Later, a solution of 60 mmol dibromoalkane (1,3-dibromopropane, 1,4-dibromo-1-butene, or 1,6-dibromo-1-hexene) in DMF (40 mL) was added dropwise to this mixture. The resulting solution was stirred for 12 h at 0 °C and then it was extracted with Et₂O/H₂O, dried MgSO₄, and purified by SiO₂ chromatography (20:1 hexane/ethyl acetate). For Py3Br (Yield: 51%): 1H NMR (300 MHz, CDCl₃) (δ/ppm): 6.71 (d, J = 1.93 Hz, 2H), 6.20 (t, J = 2.0 Hz, 2H), 4.11 (t, J = 6.38 Hz, 2H), 3.34 (t, J = 6.19 Hz, 2H), 2.29 (m, 2H) (Figure S1). 13C NMR (75 MHz, CDCl₃) (δ/ppm): 120.68, 108.46, 47.08, 34.23, 30.32 (Figure S2). For Py4Br (Yield: 54%): 1H NMR (400 MHz, CDCl₃) (δ/ppm): 6.57 (t, J = 2.08 Hz, 2H), 6.07 (t, J = 2.09 Hz, 2H), 3.85 (t, J = 6.75 Hz, 2H), 3.30 (t, J = 6.44 Hz, 2H), 1.86 (m, 2H), 1.76 (m, 2H) (Figure S5). 13C NMR (101 MHz, CDCl₃) (δ/ppm): 119.39, 107.17, 47.67, 31.95, 29.02, 28.78 (Figure S6). For Py6Br (Yield: 43%): 1H NMR (300 MHz, CDCl₃) (δ/ppm): 6.71 (t, J = 2.68 Hz, 2H), 6.21 (t, J = 2.76 Hz, 2H), 3.94 (t, J = 7.06 Hz, 2H), 3.45 (t, J = 6.74 Hz, 2H), 1.87 (m, 4H), 1.52 (m, 4H) (Figure S9). 13C NMR (75 MHz, CDCl₃) (δ/ppm): 120.54, 108.04, 49.51, 33.93, 32.75, 31.55, 27.86, 26.03 (Figure S10).

2.4. General procedure for synthesis of azido pyrroles

About 1 mmol Py-N-Br and 10 mmol of Na₃N were mixed at 50 °C for 6 h in 10 mL of DMF. After cooling to the room temperature, the reaction mixture was partitioned between water and DCM, and the aqueous phase was washed with DCM. The combined organic extracts were washed with concentrated NaHCO₃, brine, and dried with MgSO₄. The crude product was purified by chromatography (10:1 hexane/ethyl acetate). For Py3N₃ (Yield: 82%): 1H NMR (300 MHz, CDCl₃) (δ/ppm): 6.68 (t, J = 2.54 Hz, 2H), 6.19 (t, J = 2.6 Hz, 2H), 4.02 (t, J = 6.65 Hz, 2H), 3.28 (t, J = 6.42 Hz, 2H), 2.03 (m, 2H) (Figure S3). 13C NMR (75 MHz, CDCl₃) (δ/ppm): 120.60, 108.44, 48.29, 46.21, 30.73 (Figure S4). For Py4N₃ (Yield: 84%): 1H NMR (400 MHz, CDCl₃) (δ/ppm): 6.69 (t, J = 2.06 Hz, 2H), 6.19 (t, J = 2.08 Hz, 2H), 3.95 (t, J = 6.9 Hz, 2H), 3.30 (t, J = 6.73 Hz, 2H), 1.88 (m, 2H), 1.60 ppm (m, 2H) (Figure S7). 13C NMR (101 MHz, CDCl₃) (δ/ppm): 120.46, 108.20, 51.04, 49.07, 28.76, 26.24 (Figure S8). For Py6N₃ (Yield: 73%): 1H NMR (300 MHz, CDCl₃) (δ/ppm): 6.70 (t, J = 2.04 Hz, 2H), 6.20 (t, J = 2.08 Hz, 2H), 3.93 (t, J = 7.06 Hz, 2H), 3.30 (t, J = 6.82 Hz, 2H), 1.84 (m, 2H), 1.65 (m, 2H), 1.43 (m, 2H) (Figure S11). 13C NMR (75 MHz, CDCl₃) (δ/ppm): 120.50, 108.00, 51.37, 49.47, 31.54, 28.82, 26.37 (Figure S12).

2.5. General procedure for click chemistry

Py₃N₃, Py₄N₃ or Py₆N₃ (0.5 mmol) and ethynylferrocene (0.5 mmol) were dissolved in H₂O/THF (1:1, 10 mL) aqueous CuSO₄·5H₂O (0.42 mL, 1.0 M), and sodium ascorbate (0.86 mL, 1.0 M) were added and the reaction mixture was stirred at room temperature for 3 h. The resulting mixture was diluted with DCM (100 mL) and the organic phase was separated. The aqueous phase was extracted with DCM (2 × 50 mL). The combined organic phases were washed with brine and dried over MgSO₄ and filtered. After removal of the solvent under reduced pressure, the residue was subjected to purification by chromatography on silica gel.
2.6. Synthesis of polymers

The electrochemical polymerizations were carried out in a single compartment cell containing 0.01 M monomer and 0.1 M TBAPF$_6$ in 10:1 DCM/ACN or in pure DCM. All polymers were synthesized during potentiodynamic cycling between −0.5 and 1.2 V. Platinum wire or ITO-coated glass can be used as working electrodes. The counter and reference electrodes were a platinum wire and Ag/Ag$^+$, respectively. After the polymerization, the polymer was rinsed with fresh ACN and characterized by electrochemical and spectral methods.

2.7. Post-polymerization functionalization of P(Py-N$_3$) derivatives ferrocene

In order to anchor ferrocene to azide-containing polypyrroles (P(Py-N$_3$)) a classical click reaction was performed, where P(Py-N$_3$) coated ITO electrodes were soaked in 5 mL ethanol/water (1:1) solution containing 0.1 M ethynylferrocene, 1.0 M CuSO$_4$ and 1.0 M sodium ascorbate at room temperature for 24 h. Subsequently, the electrodes were thoroughly washed with methanol and distilled water, dried under vacuum to remove adsorbed ethynylferrocene. Later on these polymers were subjected to CV and FTIR analyses.

3. Results and discussion

3.1. Synthesis of pyrrole derivatives

In order to explore the effect of spacer group on electrochemical polymerization of ferrocene clicked pyrroles and modification of polypyrrole-coated electrodes, three different N-substituted pyrrole derivatives were synthesized via alkaline-mediated coupling reaction between pyrrole and dibromoalkanes in the presence of KOH. Later, the resultant bromine containing pyrroles were subjected to excessive NaN$_3$ to yield azido-pyrroles (Py3N$_3$, Py4N$_3$, Py6N$_3$). As the final step, copper(I)-catalyzed azide-alkyne Huisgen 1,3-dipolar cycloaddition reaction between ethynylferrocene and azido-pyrroles were carried out under common ‘click’ conditions to afford ferrocene-tethered pyrroles (Py3Fc, Py4Fc, Py6Fc) (Scheme 1). Chemical structures of all monomers were thoroughly proved by $^1$H NMR, $^{13}$C NMR, FTIR spectra, and HRMS analysis.

Scheme 1. Synthetic route to monomers.
DESIGNED MONOMERS AND POLYMERS

3.2. Electrochemical properties of ferrocene-anchored pyrroles

The presence of low potential reversible redox signals (around $E_{1/2} = 0.2 \text{ V}$) in all monomers is consistent with the electrochemical properties of ferrocene, which is known to exhibit reversible redox process (around $E_{1/2} = 0.25 \text{ V}$, Figure 1(a)). Both Py3Fc and Py4Fc revealed a single reversible redox couple in TBAPF$_6$/ACN system without any significant current increase at the higher potential range and without formation of polymer on the electrode surface.

On the other hand, it was possible to graft the conductive films of Py4Fc on Pt electrode in ACN/DCM (1:10) solvent mixture. As seen in Figure 1(b) Py4Fc revealed a reversible redox couple ($E_{1/2} = 0.27 \text{ V}$) which is similar to pure ferrocene (Figure 1(a)) in the same media. Amplification of the polarization potential beyond 1.0 V resulted in a steady increase in the current density indicating formation of radical cation of pyrrole unit.[29,30] Such behavior is in accordance with literature [31–33] and Py4N$_3$ as seen in Figure 1(c).

Starting from the second scan (Figure 1(d)), a new quasi
reversible redox couple was observed in the higher potential range ($E_{\text{pox}} = 0.51 \text{ V}$) which is due to doping–dedoping process of polypyrrole backbone. Moreover, increase in the electrode area was evident by the steady increase in the current density with increasing number of cycles for both redox processes. The presence of polypyrrole-based capacitative response assured the formation of a polymer without significant defects. Formation of electroactive, ferrocene containing P(Py4Fc) film was further supported by voltammogram (Figure 1(e)) which were conducted in monomer-free TBAPF$_6$/ACN/DCM media. Moreover, a direct correlation between the current intensities and scan rate was observed for both ferrocene ($R^2_{\text{anodic}} = 0.987$, $R^2_{\text{cathodic}} = 0.984$) and polymer backbone ($R^2_{\text{anodic}} = 0.999$, $R^2_{\text{cathodic}} = 0.999$) related processes in TBAPF$_6$/ACN/DCM electrolyte system (Figure S20). Similar type of results was achieved for the studies conducted in the same polymerization media with ITO electrode, which is suitable for post-polymerization studies. However, it was interesting to see the effect of solvent on the relative current intensities of ferrocene and polymer backbone-related redox processes for the polymers that were synthesized on ITO under identical conditions. As we could see the ratio of the former to the later process is higher than one for the polymers characterized in DCM/ACN system (Figure 2(a)), which is lower than one in ACN (Figure 2(b)). Such effect presumably originates from the difference in the solubility parameters of the solvents, leading to a variation in degree of swelling and availability of the active sites. In order to explore the effect of solvent on the redox behavior of Py4Fc, further studies were conducted in TBAPF$_6$/DCM system. As in the case of ACN/DCM mixture system, in DCM Py4Fc revealed a well-defined reversible redox process at around 0.3 V and an irreversible oxidation at beyond 0.9 V, where the former process is attributed to its ferrocene moiety and the latter signified the formation of radical cation of the monomer (Figure S18). However, contrary to ACN/DCM solvent system, upon continuous cycling the color of the solution in the vicinity of the working electrode changed and a homogeneous, adherent thin film was not achieved. This again could be related to the higher solvation power of DCM which prevents precipitation of polymer on electrode and leads to leakage of the oligomers in to the polymerization media.

Contrary to Py4Fc, despite varying the polarization potential, the electrode type (Pt, ITO), the supporting electrolyte (LiClO$_4$, TBAPF$_6$) and the solvent (DCM, ACN, PC) it was not possible to electrochemically polymerize Py3Fc. Following the first scan, Py3Fc revealed ill-defined redox behavior presumably due to interaction of ferrocenium ion (Fc$^+$) with radical cation of pyrrole, whereby inhibiting the oxidative coupling of radical cations (Figure S17).[16,34]

Figure 3(a) represents the cyclic voltammogram of Py6Fc in TBAPF$_6$/DCM. As we could see Py6Fc revealed two well-defined reversible redox process with peak potentials of $E_{\text{pox}} = 0.21 \text{ V}$ and $E_{\text{pox}} = 0.35 \text{ V}$. Following several potentiodynamic switches between −0.5 and 1.0 V, an electroactive, well-adhered polymer film[35] readily deposited on ITO electrode, which showed linear increase in current densities with increase in scan rate (Figure 3(b), Figure S21). Contrary to other ferrocene-containing pyrroles in the literature [19,36,37] both Py4Fc and Py6Fc were readily polymerizable without requiring the presence of a comonomer for polymerization to take place. Moreover, since the ferrocene unit is separated by the relatively long spacer groups, for both polymers it was possible to see the individual redox processes stemming from the polypyrrole and ferrocene units, which is not valid for most of the former studies.[18,38,39] However, despite their well-defined quasi-reversible redox activity on ITO electrodes, none of the ferrocene-containing polypyrroles (P(Py4Fc), P(Py6Fc)) displayed significant spectral (color) variations during repetitive redox switches between neutral and oxidized states. Optical band gap of (P(Py4Fc)) was estimated as 2.75 eV, which was calculated through the lower energy absorption edge of the spectrum.
Figure 4. Cyclic voltammogram of (a) Py3N3, (b) Py4N3 and (c) Py6N3 at 100 mV/s⁻¹ in DCM/TBA PF₆, where Pt wire, Ag/Ag⁺ and ITO were used as the counter electrode, reference electrode and working electrode, respectively.

Figure 5. Spectroelectrochemistry of (a) P(Py3N3), (b) P(Py4N3) and (c) P(Py6N3) in DCM/TBA PF₆ at applied potentials between −0.5 and 1.1 V (a: −0.5 V, b: −0.2 V, c: 0.0 V, d: 0.1 V, e: 0.2 V, f: 0.3 V, g: 0.4 V, h: 0.5 V, i: 0.6 V, j: 0.7 V, k: 0.8 V, l: 0.9 V, m: 1.0 V, n: 1.1 V).
purpose, we have investigated the redox properties of all Py-N-N₃ (Py₃N₃, Py₄N₃, Py₆N₃) derivatives. As seen Figure 4 all the monomers revealed an irreversible oxidation beyond 0.85 V, which is typical for N-substituted pyrroles. Formation of azide-containing polypyrroles (P(Py-N-N₃)) were followed by the gradual increase in the current upon repetitive cycling and by the growth of a new redox process emerging at around 0.5 V due to the doping–dedoping of polypyrrole backbone. Once the polymer films were formed they were subjected to CV studies in a fresh, monomer-free electrolyte solution at various scan rates (Figure S19). All the polymers revealed the typical surface confined redox behavior where a linear dependence between the current and scan rate is observed (Figure S22). Moreover, existence of the strong peak at around 2090 cm⁻¹ in FTIR spectra of all P(Py-N-N₃) derivatives indicated the unaltered presence of azide groups during polymerization.

The UV–vis absorption spectra of P(Py₆-post-Fc) (Figure 5) showed two absorption bands at 330–365 and 470–525 nm at applied potentials between −0.5 and 1.1 V. The former broad absorption band is typical for π–π* transitions, which dominates the optical behavior of the polymer at neutral state, and they are in good agreement with previous studies.[40] The optical gap band of P(Py₃N₃),

3.3. Post-polymerization functionalization of azide-containing pyrroles

Being one of the major advantages of the click chemistry, we performed post-polymerization functionalization of the polypyrrole derivatives through azide groups. For this purpose, we have investigated the redox properties of all Py-N-N₃ (Py₃N₃, Py₄N₃, Py₆N₃) derivatives. As seen Figure 4 all the monomers revealed an irreversible oxidation beyond 0.85 V, which is typical for N-substituted pyrroles. Formation of azide-containing polypyrroles (P(Py-N-N₃)) were followed by the gradual increase in the current upon repetitive cycling and by the growth of a new redox process emerging at around 0.5 V due to the doping–dedoping of polypyrrole backbone. Once the polymer films were formed they were subjected to CV studies in a fresh, monomer-free electrolyte solution at various scan rates (Figure S19). All the polymers revealed the typical surface confined redox behavior where a linear dependence between the current and scan rate is observed (Figure S22). Moreover, existence of the strong peak at around 2090 cm⁻¹ in FTIR spectra of all P(Py-N-N₃) derivatives indicated the unaltered presence of azide groups during polymerization.

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P(Py4N₃), P(Py6N₃) are calculated as 2.37, 2.61, 2.67 eV, respectively. As the oxidation proceeds, the later transitions in the spectrum became dominant indicating the generation of charge carrier bands inside the band gap. As typical of N-substituted polypyrrole derivatives, these polymers also could be oxidized up to a certain level by means of simple electrochemical oxidation. Contrary to P(Py4N₃), P(Py6N₃), these polymer films displayed reversible color transition between brownish yellow and blue upon switching.

Later on, these polymers were subjected to a simple click reaction with ethynylferrocene in 1.0 M CuSO₄, 1.0 M sodium ascorbate ethanol/water (1:1) media for 24 h. [41,42] After washing thoroughly with methanol (to eliminate the effect of adsorbed alkyl) the polypyrrole films (P(Py-N-post-Fc)) were characterized by electrochemical methods and FTIR. Figures 6(a)–(c) represents the CV of P(Py3-post-Fc), P(Py4-post-Fc) and P(Py6-post-Fc) and their azido polypyrrole (P(Py3N₃), P(Py4N₃), P(Py6N₃)) analogs, which were treated under click conditions without the presence of ethynylferrocene, respectively. As we could see, the former two (P(Py3-post-Fc), P(Py4-post-Fc)) displayed very similar redox behavior to their azido derivatives. P(Py6-post-Fc), on the other hand, revealed two quasi-reversible redox couples at around $E_{p_{1/2}} = 0.37$ and $E_{p_{1/3}} = 0.53$ V as in the case of P(Py6Fc). The former redox couple at lower potential range could be attributed to ferrocene units and later couple could be related to doping of polypyrrole main chain (Figure 6(d)). The higher reversibility and better peak separation in the CV of P(Py6Fc) could stem from the presence higher amount of ferrocene (one for each pyrrole unit) in the bulk of the polymer, whereas in post-polymerization functionalization mainly anchoring on the surface of the film is expected. Thus, current intensity and reversibility of the redox couples (especially those related to ferrocene units) are less defined in case of P(Py6-post-Fc) than P(Py6Fc).

Moreover, when compared with FTIR spectrum of P(Py6N₃), P(Py6-post-Fc) revealed a significant reduction in the strong IR signal at 2090 cm⁻¹ (azide stretching) and presence of new peaks at 480 cm⁻¹ due to covalent bonding of ferrocene unit (Fe/C bond vibration). Thus, CV and FTIR studies afforded solid evidence of post-polymerization functionalization only for P(Py6N₃) electrode, where effective tethering of ferrocene seems to depend on length of spacer group. The optoelectronic properties of P(Py6-post-Fc) was investigated in 0.1 M LiClO₄/ACN solution (Figure 7). While amplifying the potential from 0.0 to 1.0 V, intensity of the absorption peak at 355 nm (due to $n-\pi^*$ transition) decreased and simultaneously a new the absorption peak around 510 nm intensified due to formation of charge carrier bands. These results are in accordance with P(Py6N₃), which underlines the fact that contrary to P(Py6Fc), P(Py6-post-Fc) can be reversibly doped and de-doped with a well-defined electrochromic response. Hence, through utilization of post-polymerization functionalization it was possible to achieve an electrochromic, ferrocene-containing polypyrrole derivative.

4. Conclusion

This study demonstrates a detailed investigation on synthesis of ferrocene-anchored polypyrroles through alternative approach. Our methodology utilizes the versatility of the click chemistry and considers the effect of spacer group between pyrrole and ferrocene units both during monomer synthesis and post-polymerization functionalization stages. Our studies indicated the importance of the spacer group, which played a key role in electrochemical polymerization process by providing the opportunity for the essential pyrrole-based radical cation coupling step to take place. Contrary to literature studies, where polymerization is mainly achieved by electrochemical copolymerization, with this approach it was possible to synthesize homopolymers of pyrrole derivatives having covalently bonded ferrocene units. Furthermore, three different electrochemically synthesized azide-containing polypyrroles were subjected to click reaction in the presence of ethynylferrocene, where effective clicking (anchoring) was only achieved in case of longest spacer group. Future studies are on the way for utilization of these ferrocene-containing polymers in biosensor applications.

Disclosure statement

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