



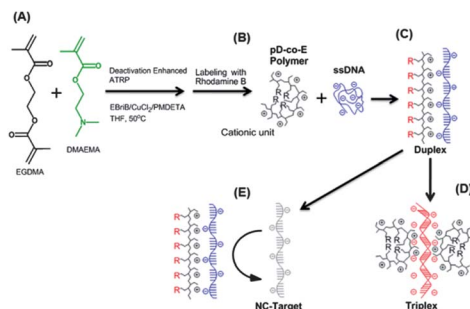
A Fluorescently Labeled, Hyperbranched Polymer Synthesized from DE-ATRP for the Detection of DNA Hybridization

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5 Ahmed Aied, Barry Glynn, Hongliang Cao, Yu Zheng, Hongyun Tai,* Abhay Pandit and Wenxin Wang*

10 A hyperbranched polymer synthesised from DE-ATRP and labelled with Rhodamine B was used to detect DNA hybridisation in serum.



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A fluorescently labeled, hyperbranched polymer synthesized from DE-ATRP for the detection of DNA hybridization†

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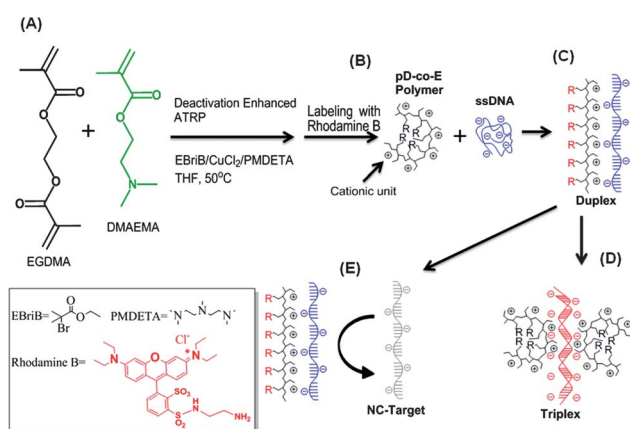
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The early detection of oligonucleotide biomarkers of disease, such as microRNAs, has been established as a fundamental factor in cancer diagnosis. As the levels of these small molecules (microRNAs) in blood have recently been found to be significantly affected in cancer patients, they offer a means of early stage detection of cancer. Towards the goal of creating a novel method of DNA hybridization detection, we report the detection of specific sequences of small oligonucleotides in a model experiment carried out in serum. The results shown here display the versatility of the DE-ATRP method in synthesizing a specific polymer structure capable of changing its physical properties in the presence of double stranded DNA. The polymer was labeled and used to detect single-stranded DNA in serum successfully.

The early detection of cancer represents one of the most promising approaches to reduce the high mortality associated with many cancers. Currently, much interest in the field of diagnostic research is focused on the detection of cancer oligonucleotide biomarkers such as mir-141 in prostate cancer.¹⁻⁴ Rapid and sensitive detection of DNA hybridization reactions is being established, but many of these methods require modification of the complementary probe in a multi-step procedure for optimal detection. In addition, many interesting DNA hybridization transducers have been reported including electrochemical and optical DNA biosensors based on conjugate polymers,⁵⁻⁷ DNA-derivatized nanoparticles,^{8,9} molecular beacons¹⁰ and bioluminescence techniques.¹¹ However, these methods, although functional, require the modification of the polymer or amplifier properties. Even though these methods have distinct advantages over conventional procedures of oligonucleotide detection (RT-PCR, northern blotting, *etc.*¹²), specificity and reproducibility of detection are compromised. In 2004, Dore *et al.* reported that a cationic water-

soluble polythiophene polymer can be used for the detection of nucleic acids at low concentrations.

Previously we have reported a deactivation enhanced atom transfer polymerization (DE-ATRP) approach, which can suppress the gelation and produce high yield “hyperbranched polymers” by the homopolymerization of multi-vinyl monomers (MVMs) even in a concentrated system.¹³ In a most recent study, we realized that this DE-ATRP reaction not only provides precise control over the molecular weight and polydispersity (PDI) but also provides the potential to kinetically control the molecular architecture in the polymerization of MVMs.¹⁴ In this study, we bring this structure to DNA detection applications through the preparation of a cationic hyperbranched polymer by the copolymerization of 2-(dimethylamino)ethyl methacrylate (DMAEMA) and ethylene glycol dimethacrylate (EGDMA) *via* the DE-ATRP approach (Scheme 1A). We demonstrate the use of a hyperbranched cationic molecule for the simple, fast and specific detection of DNA molecules without the



Scheme 1 (A) Copolymerization of DMAEMA and EGDMA *via* DE-ATRP approach leads to a hyperbranched polymer (B) which is further modified by Rhodamine B. (C) Polymer interaction with negatively charged ssDNA probe leads to an increase in fluorescence due to isolation of polymer molecules (duplex). (D) Upon complementary DNA target hybridization to the ssDNA probe, the polymer forms stronger interactions with the double stranded DNA which condenses into small nanoparticles that quench the rhodamine reducing the fluorescence intensity (triplex). (E) Adding a non-complementary target will have no effect on the fluorescence because it does not hybridize to the duplex.

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† Electronic supplementary information (ESI) available: Description of the material, polymer synthesis and characterization methods, proton nuclear magnetic resonance and polymer/DNA complexation results. See DOI: 10.1039/c2py00516f

1 need for modifications of the DNA probe or target. We hypothesize
2 that a fluorescently labeled hyperbranched polyDMAEMA-co-
3 EGDMA (termed pD-co-E, Scheme 1B) can detect specific sequences
4 of small DNA (18–24 nt) in serum through a change in the fluores-
5 cent signal. More specifically, detection is based on the conforma-
6 tional change of the polymer molecules upon binding to the ssDNA
7 probe, producing a measurable fluorescent signal. Hybridization to
8 the target sequence results in a significant change in fluorescence
9 (Scheme 1D). As with any other cationic polymer, the pD-co-E
10 polymer forms dense 3D-structures in solution quenching the fluo-
11 rescence of rhodamine molecules.

12 Firstly, the formation of the polymer–ssDNA complex (termed
13 duplex, Scheme 1C) changes the structure of the polymer due to the
14 strong ionic interaction between the two molecules, which leads to
15 exposure of the rhodamine fluorophores in the process. The second
16 step involves adding the complementary sequence to the solution
17 containing the duplex. This forms a pD-co-E/ssDNA probe/cDNA
18 complex (termed triplex, Scheme 1D) from which the polymer
19 becomes slightly dissociated. The dissociation of the polymer from
20 the hybridized DNA allows it to regain its original structure. The
21 principle of detection provided herein is therefore based on the
22 amplification of the signal produced when the ssDNA probe binds to
23 the polymer, *via* the change in the polymer's physical properties. One
24 of the advantages of this system is that no isolation step of the duplex
25 or triplex is required in order to obtain a signal from a relatively small
26 sample. The method allowed for detection of specific sequences at the
27 nanogram level in homogeneous solution but the sensitivity was
28 significantly reduced in serum.

29 The hyperbranched polymer was synthesized by DE-ATRP and
30 labeled with ethylenediamine functionalized Rhodamine B (see
31 ESI†). Gel permeation chromatography (GPC) was used to monitor
32 the reaction progression (Fig. S1 in ESI†). The peak area and shift
33 were used to analyze the ratio of monomer conversion and molecular
34 weight ($M_w = 10.6$ kDa, at 60% conversion). The low polydispersity
35 (PDI = 1.47) indicates the formation of monodispersed, branched
36 polymer molecules. The vinyl content, degree of branching and
37 DMAEMA content in the polymer were determined by calculation
38 from the ^1H NMR spectrum (7.1%, 11.8% and 82.1%, respectively)
39 (Fig. S2 and eqn (S1–S6) in ESI†).

40 The ability of pD-co-E to complex (condense) DNA was assessed
41 by UV-Visible spectroscopy (Fig. S4 in ESI†) and agarose gel elec-
42 trophoresis (Fig. S5 in ESI†). The ssDNA probe illustrated in Scheme
43 1 is a model complementary sequence to the mir-141 (a microRNA
44 expressed by prostate cells¹⁵), while the cDNA target has the same
45 sequence as the mir-141 where the thymine groups are replaced by the
46 uracil groups. The non-complementary target (NC-target, negative
47 control), on the other hand, is an ssDNA sequence that is different
48 from both sequences.

49 Different weight ratios of polymer to ssDNA probe were used,
50 with no more than 1 μg of the ssDNA probe for each sample. To
51 ensure that most of the pD-co-E will complex with the ssDNA probe
52 when detecting the cDNA, equal amounts of polymer and ssDNA
53 probe were added (*i.e.* w/w of polymer : ssDNA probe). Using the
54 UV-Visible spectrometer, we were able to detect a decrease in ssDNA
55 concentration after the addition of the polymer and formation of the
56 duplex. When the cDNA target was added, an increase in absorbance
57 was noted; however, adding the non-complementary cDNA target
58 (termed NC-target) to the duplex solution resulted in a higher signal
59 intensity (Fig. S6†). Even though this method can be used to

60 differentiate between the duplex and the triplex in water, it cannot be
61 used in biological solutions because of its non-specificity. Other
62 proteins and DNA molecules will interrupt the signal and give a false
63 reading which makes this method unsuitable for direct detection. This
64 is why we decided to use a fluorescence based method of detection.

65 To detect the DNA hybridization, the polymer was labeled with
66 Rhodamine B. The first set of experiments was carried out in solution
67 where a clear difference in color sharpness between the triplex,
68 duplex + NC-target and duplex alone could be visualized by the
69 naked eye. Fig. 1 shows the contrast between the labeled polymer and
70 the duplex. No color change is observed after adding the non-
71 complementary target, but a change in color is seen when the cDNA
72 (model microRNA) is added.

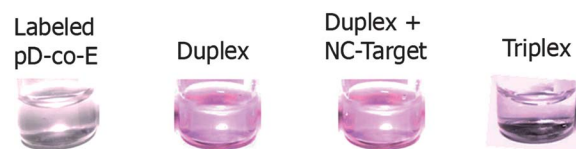
73 Model microRNA target (cDNA) detection was carried out by
74 fluorescence spectroscopy. An LED light source was used for the
75 excitation ($\lambda = 570$ nm) of the rhodamine labeled polymer and
76 emission ($\lambda = 590$ nm) was detected. LEDs are very stable and
77 provide a range of wavelengths with minimal background noise
78 producing a clear signal even at low concentrations of the analyte.¹⁶

79 The fluorescence intensity of four different samples was analyzed
80 at various concentrations (Fig. 2). After the formation of the duplex,
81 an increase in fluorescence over the labeled polymer alone is noted
82 even at a concentration of 8.13×10^{-5} μM of DNA. Adding the
83 cDNA to the duplex solution resulted in a decrease in fluorescence
84 intensity which is statistically significant down to the lowest
85 concentration.

86 The response of the duplex to the NC-target is very clear, indi-
87 cating that there is some sensitivity from the polymer's part towards
88 unspecific targets. However, the change is small and becomes insigni-
89 ficant at lower concentrations. The change is possibly due to the
90 binding of excess polymer with the NC-target, hence the increase in
91 fluorescence in that sample.

92 When the same experiment was carried out in a serum environ-
93 ment (Fig. 3), the fluorescence intensity demonstrated the same
94 changing pattern but at a significantly reduced sensitivity. This is
95 because the experiment was carried out in a complex environment
96 which contains large amounts of proteins and biomolecules that
97 interfere with the signal. The results are reproducible even at
98 a concentration of 3.25×10^{-4} mM.

99 However, in order to detect the endogenous microRNA that is
100 upregulated in prostate cancer patients (present at several hundred
101 thousand copies per microlitre in blood¹⁷), the detection sensitivity
102 and specificity have to be optimized for the detection of lower
103 concentrations. We speculate that by using a more sensitive fluo-
104 rescence spectrometer, and optimizing the polymer molecular weight
105 and charge, we will see an increase in the sensitivity of the system. The
106 detection specificity to a single mismatch can be achieved by immo-
107 bilizing the polymer onto a silica or glass surface by which removal of



108 **Fig. 1** Photographs of solutions (1.4×10^{-7} M, polymer concentration)
109 containing the rhodamine labeled polymer. These images display the
110 color changing capability of the labeled polymer when forming the
111 duplex and triplex.

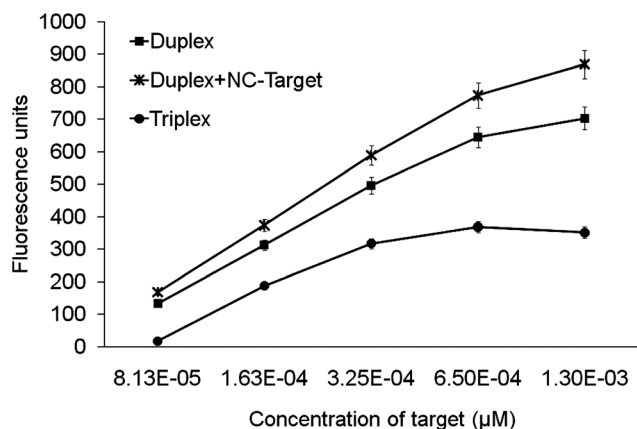


Fig. 2 Fluorescence intensity plots for the duplex, duplex + NC-target and triplex at five different concentrations. When the NC-target is added, there is a slight increase in fluorescence; however, upon adding the complementary sequence (triplex) there is a significant reduction in fluorescence further proving the change in the labeled polymer's behavior upon binding to double stranded DNA. In contrast, when the NC-target is added, the fluorescence intensity remains high ($n = 5$) (\pm S.D.) (for statistical significance mentioned in text $p < 0.05$).

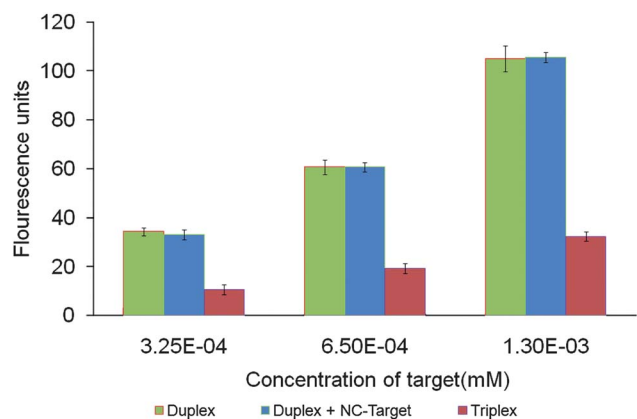


Fig. 3 Bar chart showing fluorescence intensity for the pD-co-E, duplex, triplex, and duplex/NC-target in serum. It is clear that even in the serum environment reproducible results are obtained allowing for easy differentiation between the complementary target and NC-target at various concentrations. No statistical significance between samples was obtained below a concentration of 3.25×10^{-4} mM ($n = 5$) (\pm S.D.) ($p < 0.05$).

unbound or unhybridized DNA can bring about specific sequence detection.

In conclusion, we have demonstrated successful synthesis of hyperbranched polymer using the DE-ATRP method. This hyperbranched pD-co-E polymer has shown exceptional physical properties demonstrated by its capability to form condensing particles that can quench the fluorescence of rhodamine molecules in

a homogeneous solution and in serum. The capability of the polymer to form large complexes with double stranded DNA is what gives it the ability to quench the rhodamine molecules. The polymer and the method of detection developed herein are unique and demonstrate the applicability of the newly developed cationic, hyperbranched polymer in bioanalytical applications and specifically for the detection of DNA hybridization.

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