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Fluorescence detection of natural RNA using rationally designed “clickable” oligonucleotide probes†

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Herein a reliable approach to the design of effective fluorescent probes for RNA detection is described. The fluorescence signalling of hybridization by internally positioned polyaromatic hydrocarbons and rhodamine dyes was achieved with a low fluorescence background signal, high fluorescence quantum yields at ambient and elevated temperature, high selectivity and signal specificity of the probes when binding to miR-7 and circRNA targets.

RNA targeting is a rapidly evolving field of research and clinical diagnostics which applies synthetic fluorescent probes and modern biophysical techniques to monitor desired targets *in vitro* and *in vivo*.¹ Preparation of fluorescent oligonucleotides with high potential in RNA detection by a simple and rapid method, *e.g.* by fluorescence, is an important aspect of current nucleic acid chemistry. Recently copper(i)-catalyzed azide-alkyne cycloaddition (CuAAC) click chemistry has been successfully applied for nucleic acid labelling, allowing preparation of various oligonucleotides in high yields and purity.^{2,3} Furthermore, attachment of fluorophores by click chemistry to 2'-*O*-propargyl uridine scaffold U^P was proven to yield probes with promising properties in diagnostics (Fig. 1).⁴ However, in order to further utilize “clickable” oligonucleotides in diverse analytical experiments, the fluorescent probes have to meet several additional requirements, such as high photo- and chemical stability, low fluorescence background signal from single stranded molecules, high binding specificity and low limit of target detection.⁵ Ideally, incorporation of fluorescent

dyes should furthermore be simple and allow high flexibility in probe design.

MicroRNAs (miRNAs) are short (~22 nucleotides) noncoding RNA molecules that post-transcriptionally repress the expression of protein-coding genes by binding to 3'-untranslated regions of the target mRNAs.⁶ An increasing amount of evidence shows that miRNAs control a large number of biological processes, including cell differentiation, tumorigenesis and regulation of metabolism. A thoroughly studied miRNA is miRNA-7 (miR-7) which is highly expressed in brain neurons and also an important negative regulator of oncogenes and, hence, functioning as a tumor suppressor factor in oncogenesis.⁷ miR-7 was recently identified to be selectively expressed within the hypothalamus, a part of the brain that controls vital bodily functions.⁷

Another class of recently discovered RNA molecules with important, yet not fully studied, regulatory functions are circular RNAs (circRNAs).^{8,9} Interestingly, it was found that a human circRNA, antisense to the cerebellar degeneration-related protein 1 transcript (CDR1as), is densely bound by

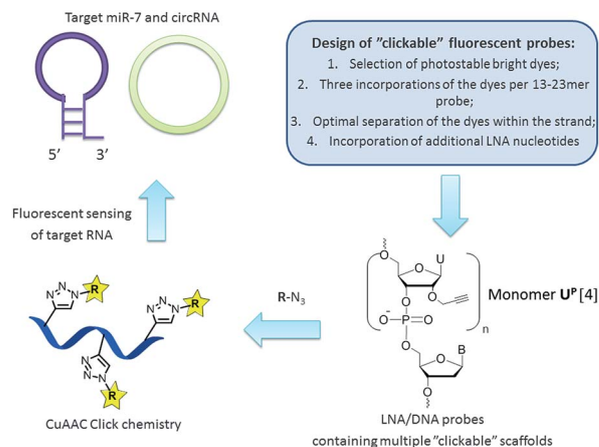


Fig. 1 General strategy for targeting natural RNA molecules (*i.e.* miR-7 and circRNA) by “clickable” fluorescent probes.

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miR-7 molecules in cells and presumably acts as a regulator of miR-7 activity.⁹ Clearly, *in situ* visualization of miR-7 and CDR1as will significantly contribute current understanding of regulatory RNA pathways and may provide further insight into the regulatory role of circRNA.^{6–9} In the present work we aimed at a robust approach to preparation of fluorescent probes with high binding affinity and efficient fluorescence detection of miR-7 and circRNA (Fig. 1). Herein, the design and synthesis of miR-7 and circRNA targeting oligonucleotides containing 2'-O-propargyl uridine scaffolds **U^P**, their fluorescent labelling by CuAAC click reactions, and spectroscopic characterization of the resulting fluorescent probes *in vitro* are reported. The ability to design and rapidly prepare bright selective fluorescent probes for detecting any natural RNA target could strongly support research in this direction.^{6–9}

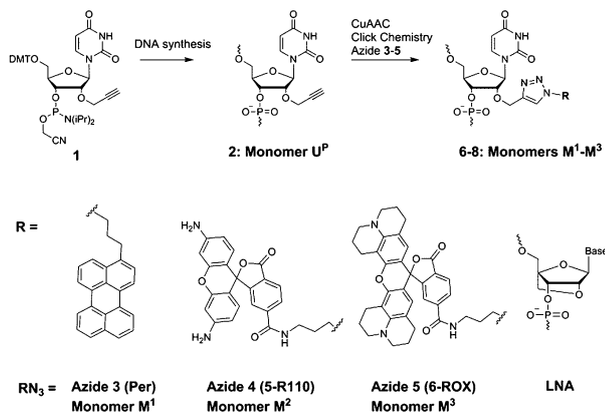
First, we selected perylene and two rhodamine derivatives, 5-R110 and 6-ROX, as fluorophores with interesting optical properties to be internally attached to the scaffold **U^P** within new probes (Fig. 1 and Scheme 1).^{4,10,11} Second, three incorporations of the dyes within 13–21mer oligonucleotides were previously demonstrated to be optimal for spectral properties of the probes.⁴ Therefore commercially available phosphoramidite reagent **1** was triply incorporated into the 23mer miR-7 and circRNA targeting sequences resulting in oligonucleotides **ON1**, **ON2** (Fig. 1 and Table 1; ESI, Table S1†).^{6–9} Third, the “clickable” scaffolds **U^P** within **ON1**, **ON2** were separated by 1–2 nucleotides in order to achieve optimal fluorescence emitted from the probes upon target binding.⁴ As a final design aspect, additional LNA (locked nucleic acids; Scheme 1) monomers were incorporated into **ON1**, **ON2** to enhance binding affinity and specificity of the probes to complementary targets.¹¹ Finally, reference probes **ON3**, **ON4** were prepared in order to evaluate influence of the modified monomers **U^P** and **M¹–M³** on target binding (Table 1).

CuAAC click conjugation of **ON1**, **ON2** with azides 3–5 was performed either under microwave conditions for 15 min at 60 °C (azide 3), or at ambient temperature for 24 h (azides 4 and 5), giving the desired products **P1–P6** in high purity and yields

of 74–83% (ESI†). Microwave conditions of the click reactions were applied for incorporation of monomer **M¹** in order to improve solubility of the azide 3. The products were characterized by ion-exchange (IE) HPLC and MALDI MS (ESI, Table S1†). Incorporation of the fluorescent monomers into **ON1**, **ON2** was further confirmed by characteristic bands in the visible region of the corresponding absorbance spectra (Table 2; ESI, Table S4 and Fig. S1†). Notably, hybridization had an influence on the shape and intensity of absorbance curves which implied interaction of the dyes with microenvironment, *e.g.* dye aggregation and stacking interactions between the dyes and nucleobases.^{4,12,13}

Hybridization of the prepared probes with target DNA and RNA strands was studied using T_m and circular dichroism (CD) experiments in a medium salt phosphate buffer (Table 1, ESI, Fig. S2, S3 and Table S3†). For all the modified probes S-shape of the melting curves and the characteristic CD signal of an A/B type duplex confirmed successful adoption of the dyes within the duplexes (ESI, Fig. S2 and S3†).¹¹ Additional LNA nucleotides improved binding affinities of the modified probes in spite of 5–9 °C destabilizing effect of monomer **U^P** (Table 1, T_m for **ON1–ON4** compared to unmodified references). Perylene-labelled probes **P1**, **P2** showed high binding affinity towards complementary DNA/RNA (T_m 64–68 °C), which is most likely caused by additional stacking interactions provided by the polyaromatic hydrocarbons within the double stranded complexes.¹⁰ Furthermore, rhodamine-modified probes **P3–P6** showed superior binding affinity towards RNA in comparison to DNA targets, indicating at better adoption of the bulky rhodamine fluorophores within the corresponding DNA–RNA hybrids (T_m 55–57 °C *vs.* 59–62 °C for the duplexes with DNA and RNA targets, respectively). However, incorporation of the fluorophores within monomers **M²**, **M³** had a negative effect on T_m values compared to **U^P**-modified and LNA/DNA references (T_m decrease \sim 5 °C). All the probes discriminated a single mismatch in the central positions of DNA and RNA targets by -3.5 to 8 °C and -1 to 8 °C, respectively (ESI, Fig. S3†). Notably, the T_m values were comparable to those for 5'- and internally labeled LNA/DNA fluorescent probes obtained from commercial suppliers (ESI, Tables S2 and S3†). Finally, since all the duplexes having single-nucleotide mismatches were formed at the temperature above 19 °C, all of them were subjected to the fluorescence studies described below.

Fluorescence detection of hybridization with complementary miRNA, circRNA and their cDNA analogues was studied by *in vitro* experiments at ambient (19 °C) and elevated temperature (37 °C). The latter was done in order to evaluate potential of **P1–P6** for *in vivo* applications. The *in vitro* assay using **P1–P6** was characterized by series of photophysical and diagnostic parameters: discrimination values (D), fluorescence quantum yields (Φ_f), fluorescence brightness (FB) and limit of target detection (LOD) values (Table 2; ESI, Table S4 and Fig. S4†).¹⁰ In order to efficiently detect a target *in vivo*, D value of ≥ 10 is preferred. Perylene-labelled probes **P1**, **P2** displayed high D values (up to 10.6), accompanied by high fluorescence quantum yields and high FBs at both 19 °C and 37 °C. The probes **P3**, **P4** showed the highest Φ_f and FB values (up to 0.99 and 138,



Scheme 1 Modified monomers applied in this study; postsynthetic CuAAC click chemistry generating fluorescent miR-7 and circRNA oligonucleotides. LNA = locked nucleic acid.



Table 1 Thermal denaturation temperatures of the modified duplexes in a medium salt phosphate buffer^a

#	Sequence 5' → 3'	Target	T _m , °C vs. complementary	
			DNA	RNA
ON1	ACAACAAA ^L ATC ^L ACU ^P AGU ^P CU ^P TCC ^L A	miR-7	62.0	66.5
ON2	GAAGAC ^L GTG ^L GAU ^P TU ^P TCU ^P GGA ^L AGA	circRNA	62.0	66.0
ON3	ACAACAAA ^L ATC ^L ACTAGTCTTCC ^L A	miR-7	67.0	66.0
ON4	GAAGAC ^L GTG ^L GATTTTCTGGA ^L AGA	circRNA	70.0	72.0
P1	ACAACAAA ^L ATC ^L ACM ¹ AGM ¹ CM ¹ TCC ^L A	miR-7	68.0	68.0
P2	GAAGAC ^L GTG ^L GAM ¹ TM ¹ TCM ¹ GGA ^L AGA	circRNA	68.0	64.0
P3	ACAACAAA ^L ATC ^L ACM ² AGM ² CM ² TCC ^L A	miR-7	57.0	62.0
P4	GAAGAC ^L GTG ^L GAM ² TM ² TCM ² GGA ^L AGA	circRNA	57.0	61.0
P5	ACAACAAA ^L ATC ^L ACM ³ AGM ³ CM ³ TCC ^L A	miR-7	55.0	60.0
P6	GAAGAC ^L GTG ^L GAM ³ TM ³ TCM ³ GGA ^L AGA	circRNA	57.0	59.0

^a T_m of unmodified duplexes formed by miR-7 and circRNA: 61.0 °C and 60.0 °C with complementary DNA, and 57 °C and 55.5 °C with complementary RNA, respectively; LNA monomers are marked with an L in superscript.

Table 2 Spectral, photophysical properties and diagnostic characteristics of the complexes between the probes and RNA targets in a medium salt buffer^a

#	λ _{max} ^{abs} (nm)	λ ^{ex} (nm)	λ _{max} ^{fl} (nm)	D	Φ _f		FB		LOD, nM	
					19 °C	37 °C	19 °C	37 °C	19 °C	37 °C
P1	423, 451	423	450	10.4	0.58	0.45	27	23	10	10
P2	423, 451	423	488	5.6	0.47	0.40	34	32	10	10
P3	480, 506	480	526	2.1	0.90	0.96	102	104	<5	<5
P4	482, 506	480	528	6.9	0.90	0.99	133	138	<5	<5
P5	549, 584	580	605	1.0	0.23	0.23	32	32	10	10
P6	546, 585	580	607	1.1	0.18	0.18	30	30	10	10

^a λ_{max}^{abs}, λ^{ex}, λ_{max}^{fl}, Φ_f, FB and LOD are maxima of absorbance, excitation wavelength, fluorescence maxima, fluorescence quantum yield, FB (fluorescence brightness) = Φ_f × ε_{max}/1000, and limit of target detection values. Φ_f values are measured by a relative method (ESI) using perylene and oxazine 170 perchlorate as fluorescence standards. D is determined at 19 °C as a ratio of fluorescence intensities at fluorescence maximum of double-stranded complex to the corresponding single-stranded probe (452 nm for **P1**, **P2**, 530 nm (**P3**, **P4**), and 605 nm (**P5**, **P6**)). LODs were determined by series of target titration experiments described in ESI.

respectively), also at elevated temperature. High FBs resulted in low LOD values for **P1–P4** (<5–10 nM), which is beneficial for diagnostic applications of the probes.⁵ However, low D values make them not applicable for diagnostic settings. Finally, **P5**, **P6** showed similar FB and LOD values compared to those observed for **P1**, **P2**, and lower D values between single-stranded conjugates and corresponding complexes with miR-7 and circRNA (Table 2). The observed quenching of fluorescence within single-stranded **P1–P4** most likely results from aggregation of the hydrophobic dyes surrounded by aqueous media, since both extinction coefficient and ratio of visible absorbance bands I/II of **P1–P4** vary for the single strands and duplexes (ESI, Fig. S1†).¹³ In turn, as suggested by altered ratio of absorbance bands and increased extinction, hybridization results in positioning of the dyes in a less hydrophilic environment, resulting in the fluorescent signal.

Next, none of the commercially available 5'- or internally labeled LNA/DNA probes **R1–R8** used in this study showed sensitivity of fluorescence to single-nucleotide mismatch in the RNA/cDNA targets (D 0.6–1.5; ESI, Fig. S5;† **R1–R8** contain fluorophores with similar optical properties to **M¹–M³**). On the contrary, “clickable” probes **P1–P6** showed moderate

fluorescence mismatch discrimination in cDNA targets (*i.e.* only numerous positions were discriminated; data not shown), and efficient discrimination in most of the examined miR-7 and circRNA targets by quenching of fluorescence (*e.g.* D 0.1–0.7 for **MT1–MT8** and probes **P1**, **P3**, **P5**; ESI, Tables S4, S5 and Fig. S5†). Overall, four duplexes out of forty eight studied showed low mismatch discrimination (D 0.9–1.6 by **P4**: **MT1/MT3/MT8** and **P2**: **MT7**; ESI, Table S4†). This implies that the performed internal incorporation of monomers **M¹–M³** with expanded aromatic π-electron systems results in the probes which are in general potent to sense minor changes in micro-environment such as single-nucleotide mismatches.

A critical challenge for detecting single-nucleotide polymorphisms (SNPs) in natural RNA targets is that most often a mutant genotype is present at a very low concentration with respect to the wild-type variant.¹⁴ Therefore, as the final aspect, biologically relevant mixtures of a wild-type and mismatched circRNA targets were analysed using the selected bright probe **P4** and circRNA target **MT13** containing A → G mismatch in the central position (Fig. 2; ESI, Tables S3 and S7†). Fluorescence of fully matched duplex **P4–MT13** consistently decreased upon increasing concentration of the



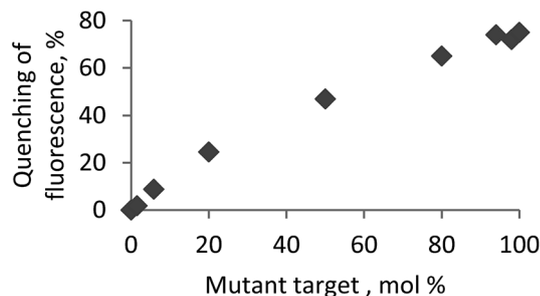


Fig. 2 Genotyping mutant circRNA MT13 in the mixture with a wild-type target circRNA using the probe P4. Fluorescence assay was performed at 19 °C using excitation wavelength of 480 nm and recording fluorescence intensity at 528 nm. Total concentration of the probe and RNA targets was 1.0 μ M. circRNA: 5'-r(UCU UCC AGA AAA UCC ACG UCU UC); MT13: 5'-r(UCU UCC AGA AGA UCC ACG UCU UC); mismatched nucleotide (replacement A \rightarrow G) is underlined.

mismatched complex which allows accurate estimation of the mismatch abundance in the corresponding analyte. Finally, high base specificity of fluorescence signal is an additional advantage of the prepared probes in genotyping of natural RNA.¹⁴

Conclusions

In this work the rationally designed “clickable” probes are demonstrated to be promising tools in RNA detection by fluorescence. The background fluorescence of single strands was significantly reduced, while upon formation of complexes with target miR-7 and circRNA emission increased giving remarkably high quantum yields (Φ_f up to 0.99) and, therefore, low limit of detection values (<5 nM in solution). Importantly, this was achieved by internal modification of the LNA/DNA oligonucleotides providing additional thermal stability and selectivity of the probes towards the target. Accompanied by simple design and robust CuAAC synthetic route, simple purification and high yields of the products, effective target binding, low single-strand background signal (D up to 10.6) and consistent specific fluorescence, the “clickable” probes could become a reliable platform for detection of natural RNA.

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