Supersensitive Multifluorophore RNA-FISH for Early Virus Detection and Flow-FISH by Using Click Chemistry


The reliable detection of transcription events through the quantification of the corresponding mRNA is of paramount importance for the diagnostics of infections and diseases. The quantification and localization analysis of the transcripts of a particular gene allows disease states to be characterized more directly compared to an analysis on the transcriptome wide level. This is particularly needed for the early detection of virus infections as now required for emergent viral diseases, e.g., Covid-19. In situ mRNA analysis, however, is a formidable challenge and currently performed with sets of single-fluorophore-containing oligonucleotide probes that hybridize to the mRNA in question. Often a large number of probe strands (>30) are required to get a reliable signal. The more oligonucleotide probes are used, however, the higher the potential off-target binding effects that create background noise. Here, we used click chemistry and alkyne-modified DNA oligonucleotides to prepare multiple-fluorophore-containing probes. We found that these multiple-dye probes allow reliable detection and direct visualization of mRNA with only a very small number (5–10) of probe strands. The new method enabled the in situ detection of viral transcripts as early as 4 hours after infection.

Introduction

Gene expression varies significantly between individual cells and it is strongly altered in disease states. Viral infections for example lead to early transcription of virus-specific genes that could be exploited for an early diagnosis and characterization of the infection. In general, basically all malfunctioning processes in cells induce transcriptional changes.[1,2] These go in hand with altered levels of messenger RNAs (mRNAs). In extreme cases disease related mRNA may not be present at all in the normal state. In most cases however, the levels of specific mRNA will be changed, which requires reliable methods to quantify mRNA transcripts. Detection and quantification of a specific mRNA is thus highly desirable from a diagnostic point of view. Particularly informative are methods that allow the quantification of mRNA levels with spatial resolution. Currently, however, intracellular localization and quantification of mRNA faces a number of challenges that hinder routine use. The most common way to detect mRNA (or other RNAs) in cells is fluorescence in situ hybridization (RNA-FISH).[3] The method reveals localization patterns of individual RNA transcripts in cells or tissues and as such, RNA-FISH is the method of choice for quantitative single-cell transcriptomic studies.[4–8] The currently available technology behind RNA-FISH technologies is based on multiple (up to 50) individual anti-sense single-stranded (ss) DNA probes, which are approximately 22 nucleotides long. Each probe oligonucleotide carries a single fluorophore, which is typically introduced as its activated NHS ester to an amino group present at the 3’-end of the probe.[9,10] The pooled fluorescent ssDNA probes are finally added to fixed and permeabilized cells for hybridization with the target RNA. The large number of probe strands in such experiments is needed to create a sufficiently strong fluorescence signal. However, generally the larger the number of probe oligonucleotides that are used, the larger is often also off-target staining, which obscures the signal-to-noise ratio. A solution to the problem is deconvolution software that is able to increase the specific signal.[10] From a chemical point of view reduction of the number of probe strands is desirable and this has led to efforts to modify the probe oligonucleotides with, for example, LNA to increase binding. Importantly, mRNA analysis based on flow-cytometry is so far very challenging with contemporary RNA-FISH.

Here we report a small FISH-probe set for mRNA, where every probe contains three fluorophores instead of just one. These multichromophore probes were conveniently prepared...
using the Cu\(^{1+}\)-catalyzed azide-alkyne click reaction.\[^{[10–19]}\] In order to avoid stacking of the fluorophores on top of each other, which might induce self-quenching, we chose a fluorophores with two additional sulfonate groups, which provide two negative charges per fluorophore. This is supposed to minimize the interaction with the fluorophores with each other and with the negatively charged DNA. Indeed, with this design a small number of probe strands (5–10) was found to be sufficient for the visualization of RNA transcripts. The new probes design allowed not only transcript quantification and localization by microscopy, but it also enabled transcript analysis using flow-cytometry.

**Results and Discussion**

The new procedure based on click chemistry is illustrated in Figure 1. As a test-system for the mRNA-FISH we used a HEK293T cell line transfected with a plasmid containing the gene coding for the enhanced green fluorescent protein (eGFP). We synthesized ten DNA probe oligonucleotides targeting specific areas of the eGFP-mRNA transcript, with each one containing 22 nucleotides (Figure S1 in the Supporting Information). Into each of the ten DNA probe strands we inserted three C8-alkyne-dU building blocks at former dT positions (Figure 1). The new procedure based on click chemistry is illustrated in Figure 1. As a test-system for the mRNA-FISH we used a HEK293T cell line transfected with a plasmid containing the gene coding for the enhanced green fluorescent protein (eGFP). We synthesized ten DNA probe oligonucleotides targeting specific areas of the eGFP-mRNA transcript, with each one containing 22 nucleotides (Figure S1 in the Supporting Information). Into each of the ten DNA probe strands we inserted three C8-alkyne-dU building blocks at former dT positions (Figure 1). The so obtained three Eterneon containing DNA probes (10 × 3) were finally purified by a simple ethanol precipitation. Due to the high efficiency of the click reaction, no further HPLC purification of the probes was required. For the probes prepared for the

![Figure 1. Depiction of RNA-FISH and schematic representation of the probe synthesis. a) Synthetic oligonucleotides with C8-alkyne-dU modifications in various positions were individually conjugated with a fluorescent dye azide. After reaction, the oligonucleotides were mixed to a probe set. b) The probe set was hybridized to the mRNA. After in situ hybridization, the mRNA molecules can be detected by flow-cytometry and/or microscopy. c) Depiction of the probes labeled with click chemistry hybridized to the target mRNA.

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![Figure 2. mRNA of eGFP-expressing HEK cells labeled with 10 × 3 and with 30 × 1 probe sets. a) FISH microscopy images of the 10 × 3 and 30 × 1 set of probes (scale bar: 10 μm) b) Comparison of the signal and background intensities. c) Signal and background obtained with CuAAC-labeled probes when using 3 and 10 oligos.

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We next experimented with different number of probe strands and dye loading and found that the most reproducible data were indeed obtained with ten oligonucleotide probe strands containing each three fluorophores. In our hands this probe design provided in all investigated cases the best signal-to-noise ratios (Figures 2c and 56).

We next investigated if we could further reduce the background signal when we performed the click reaction after in situ hybridization as shown in (Figure S7). For this study, we used a set of 10 probe strands with 3 alkyne units, hybridized them with the cellular mRNA and performed the CuAAC-reaction subsequently in situ with a TAMRA azide. While the signal-to-noise ratio indeed improved, we noted that we had to perform extensive washing in order to remove additional dye, which is typically used in large excess. This makes this procedure a little more accurate but cumbersome to perform.

We next investigated if the performance of the 10 × 3 probe design allows to detect mRNA even using flow-cytometry in a mixed cell population. For the experiment, we mixed HEK293T cells with and without eGFP-expression in a ratio 20:1 (95 % HEK293T + 5 % HEK293T-eGFP). Then, flow-cytometric measurements of the mixed cell populations were performed at \( \lambda_{\text{ex}} = 488 \text{ nm/} \lambda_{\text{em}} = 530 \text{ nm} \) for the detection of the eGFP protein. The correct ratio of the mixed cell population was nicely reproduced (Figure 3a). We then used the flow-FISH protocol described by Arrigucci et al., which involves trypsinization and resuspension of cells. The cells in suspension were permeabilized, fixed and subsequently incubated with the probe sets. Again, we performed the study with the new 10 × 3 set in comparison to a classical 30 × 1 design. When we measured at \( \lambda_{\text{ex}} = 633 \text{ nm/} \lambda_{\text{em}} = 660 \text{ nm} \), which is suitable for both the Eternoon-Red 645 azide and the Quasar 670 dye, in the absence of hybridized probes, a single population was observed (Figure S8i). The upper plot of Figure 3b shows the hybridization experiment using the classical probe set at 0.05 ng/μL. Here, only a single population containing both GFP-positive and GFP-negative cells was detected at 660 nm. When the 10 × 3 probe set was used however at the same concentration (lower plot), the GFP-positive population nicely separated from the GFP-negative cells. When the separated population was gated (in blue), the exact proportion of cells expressing the eGFP was observed for the two different detection wavelengths. The same result with the classical 30 × 1 probe set was only obtained, when the concentration was increased fourfold to 0.2 ngDNA/μL (Figure S8ii). These flow-FISH data show again that the 10 × 3 probe allows to perform flow-FISH. Although an exact comparison between the 10 × 3 and the 30 × 1 probe sets is not possible, because of the differences in the fluorophore, the connectivity of the fluorophore to the probe and the quality of the strands, we believe that the data support the idea that flow-FISH is possible with our new probe design that has a limited number of probe oligonucleotides each one carrying three fluorophores.

In order to show that the new probe oligonucleotides are able to report transcription of a relevant endogenous gene by flow-cytometry, we next tested if the above method would be suitable for the detection of the ABL1-transcript. This time, we used a slightly larger 15 × 3 probe set. In order to again obtain comparative information we compared our design with a reported detection that used in this case 48 single labelled oligonucleotides (Semrau et al.). The results are depicted in Figure 3c. While the 15 × 3 probe set with only 0.05 ng DNA/μL provides a clear shift in fluorescence compared to the negative control, the 48 × 1 probe set was under these conditions unable to provide a specific signal separation at this concentration.

We finally explored if the new (10 × 3) probe design enables imaging of RNA targets that are hard to image with conventional FISH probes. Such targets are characterized by extensive secondary structures that provides only few accessible sites for probe hybridization. We tested our probes by targeting a ~1 kb RNA viral transcript of the rotavirus A (RVA) gene segment-7. The idea was to test whether our set of probes detects these transcripts shortly after virus infection when the concentration of the transcript is expected to be very low. This specific target was chosen because of its extensive secondary structure that precludes hybridization of multiple probes, posing additional
challenges for conventional FISH.\textsuperscript{[21]} We fixed rotavirus-infected cells, 4 hours post infection and incubated them with the (10×3) DNA probe set targeting the RNA regions that were predicted to have less pronounced secondary structures.\textsuperscript{[20]} To facilitate the detection of virus-infected cells, we took advantage of a stable cell line that expresses the rotavirus non-structural protein (NSPS)\textsuperscript{[26]} fused to eGFP (see Methods in the Supporting Information).

After 4 hours post infection, RNA transcripts could be readily detected (Figure 4, top panel, red signal). No transcript specific signal was observed in mock-infected cells (Figure 4, bottom panel).

In parallel, we also analyzed eGFP-NSP5-marked foci in RVA-infected cells to identify cells at a more advanced stage of infection (Figure 4, top panel, green signal). As expected, these cells showed higher levels of the RNA transcript fully consistent with an increase of the amount of viral RNA transcripts over the course of the viral infection.

Importantly, the increased sensitivity of the new 10×3 detection approach uncovered a number of interesting insights. First, the data show a large variation in the amount of transcripts in different cells at the onset of infection. These variations likely reflect differences in the transcriptional activities of different rotavirus particles.\textsuperscript{[20]} Furthermore, the transcripts do not seem to accumulate in a particular cellular compartment or in virus-induced organelles during early infection. These results thus show that the new 10×3 probe design is a significant step forward because it provides new biological insights.

Conclusions

We show here that by decreasing the number of probes and increasing the number of fluorophores on oligonucleotide probes using click chemistry provides fluorescent probe strands that allow efficient detection of RNA transcripts in live cells. The probes have a superb sensitivity and allow detection of transcripts that due to high secondary structure content provide only a limited number of binding sites. Self-quenching seems to play only a minor role, possibly because we used fluorophores that carry two negative charges each. The new probe design is so sensitive that it even allows flow RNA-FISH to be established for demanding flow cytometry applications. These results pave the way for the detection of small highly structure RNA transcripts and transcripts with low abundance. A potential application could be the monitoring of leukemia therapy by flow-FISH to prevent dangerous relapse cases or the very early detection of virus infections now needed for the detection of the Corona virus to reduce the diagnostic gap and prevent uncontrolled propagation of the disease.

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Conflict of Interest

The authors declare no conflict of interest.

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![Figure 4. Rotavirus gene segment-7 RNA transcripts imaged with 10×3 probes. Upper panel: RNA transcripts (red) in virus-infected cells stably expressing rotavirus non-structural protein NSPS fused to eGFP (green) detected 4 hours post infection. Lower panel: Mock-infected MA104 cells stably expressing NSPS-eGFP. Scale bars: 20 μm.](image-url)

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