Minimizing off-target signals in RNA fluorescent in situ hybridization

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ABSTRACT

Fluorescent in situ hybridization (FISH) techniques are becoming extremely sensitive, to the point where individual RNA or DNA molecules can be detected with small probes. At this level of sensitivity, the elimination of ‘off-target’ hybridization is of crucial importance, but typical probes used for RNA and DNA FISH contain sequences repeated elsewhere in the genome. We find that very short (e.g. 20 nt) perfect repeated sequences within much longer probes (e.g. 350–1500 nt) can produce significant off-target signals. The extent of noise is surprising given the long length of the probes and the short length of non-specific regions. When we removed the small regions of repeated sequence from either short or long probes, we find that the signal-to-noise ratio is increased by orders of magnitude, putting us in a regime where fluorescent signals can be considered to be a quantitative measure of target transcript numbers. As the majority of genes in complex organisms contain repeated k-mers, we provide genome-wide annotations of k-mer-uniqueness at http://cbio.mskcc.org/~aarvey/repeatmap.

INTRODUCTION

The gene expression profiles of individual cells can be drastically different from that of adjacent cells. This is particularly true in developing or heterogeneous tissues such as embryos (1), proliferative adult epithelia (2) and tumors (3). Visualization of RNA expression patterns in fields of cells is often accomplished with fluorescence in situ hybridization (FISH) using antisense probes. Analysis of cellular patterns of gene expression by FISH has provided insight into prognosis (3) and cell fate (4) of tissues. A challenge for the future is to use FISH in tissues to quantify RNA expression levels on a cell-by-cell basis, which requires high resolution, high sensitivity and high signal-to-noise ratios (1,5–8).

A major hurdle in making RNA FISH methods quantitative has been increasing sensitivity and specificity to the point where genuine target RNA signals can be distinguished from background. One way to produce probes of high specificity has been to produce chemically-synthesized oligonucleotides that are directly labeled with fluorophores, and tiled along regions of RNA sequence (6–8). Although directly-labeled oligo probes are elegant, they have not yet been widely applied, in part due to their expense, and in part due to their relatively low signal strength (6–8). One alternative method for single RNA molecule detection employs long haptenylated riboprobes that are enzymatically synthesized from cDNAs (1,5). Such probes are cheaply and easily produced, and when detected with primary and fluorescently-labeled secondary antibodies, they have higher signal intensities and equivalent resolution when compared to probes that are directly labeled with fluorophores (1,5).

However, tiling probes have a natural advantage with respect to specificity: if a single probe ‘tile’ hybridizes to an off-target transcript, it is unlikely to generate sufficient signal to pass an intensity threshold that is characteristic of genuine RNA transcripts, which contain multiple tiled binding sites. In contrast, a single haptenylated probe, even if fragmented to sizes in the range of hundreds of nucleotides, may yield strong off-target signals due to the amplification conferred by primary and secondary antibodies. One traditional approach to determine background levels of fluorescence, and thus act as a crude estimate of specificity, has been the use of sense

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The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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sequence probes. However, such ‘specificity controls’ are only related to the antisense sequences by complementarity and can thus contain (or lack) repeated sequences that are entirely unrelated to the antisense probes. So while sense probes are sometimes indicative of non-specific ‘sticking’ to the cross-linked cellular matrix, they are not an appropriate control for sequence-based off-target hybridization.

Off-target hybridization signals have often been studied in the context of the specificity of oligonucleotide microarrays. Interestingly, these past studies demonstrated that 25-mer repeated sequences in probes of length 50–70 oligonucleotides could produce non-target ‘noise’ that was about 15% of the signal strength (9,10). However, the conditions and probe length of microarray hybridization and FISH are considerably different, thus leaving the question of specificity of in situ hybridizations unaddressed.

We find that repeated sequences (100% matches) of only 20–25 bp are able to drastically reduce probe specificity in RNA FISH experiments. Some probes with small repeats can even be completely uninformative about the expression pattern of the target gene, whereas others suffer a reduction in signal-to-noise ratio up to an order of magnitude. The removal of small repeated sequences in standard, long FISH probes is an enormous aid in obtaining high quality, high confidence results, and should be a standard step in RNA FISH experiments.

We supply an easy to use program to assay potential probes for repeats and the frequency of repeats in common genomes. We also provide uniqueness annotations for several organisms whose tissues are subjected to RNA FISH.

**MATERIALS AND METHODS**

**RNA probe preparation**

*Sce* probe templates were prepared by PCR of genomic *Drosophila melanogaster* DNA and cloned into a PCRII vector (Invitrogen). The *Scr* 3’UTR unique probe template is 359-bp long (GC content 39%), and was generated using PCR primers 5′-ATATGGATCCGCGCGTCGATCAACCAACATCC and 5′-AGTCATACTAGTAA GGACCCTCGAGCATTCG. The *Scr* 3’UTR non-unique probe template is 357-bp long (GC content 65%) and was generated using primers 5′-GACAGGATCCAGTGTTATCAGTCGAGG and 5′-CGTCATCTCAGGACAGCAGAACAAGTTCG. The *abd-A* probe templates were prepared by PCR from cDNA. The *abd-A* unique probe template is 1.47 kb long (GC content 43%) and was generated using primers 5′-TCGCATCGGATCC and 5′-GACAGATGGTAA GGCCCTCGAGCATTCG. The *abd-A* non-unique template is 1.65 bp long (GC content 44%) and was generated using the primers 5′-GAGAGCATGAAATCAGGGCC and 5′-GCGATCG ATTGAAAGGCCCT. Hatpientylated RNA probes were synthesized as described previously (1), and diluted 70-fold in hybridization buffer before use.

**In situ hybridization and staining**

In *Scr* FISH experiments, *D. melanogaster* embryos were collected 3–4 h after egg lay so that most embryos were at stages 6–7. In *abd-A* FISH experiments, embryos were collected 4–8 h after egg lay to obtain stage 11 embryos. Fixation of the embryos and simultaneous detection of RNAs in FISH experiments were performed as described previously (1). RNA probe hybridization was carried out overnight at 55°C in hybridization buffer. For the quantitative experiments shown in Figure 4, after several washes in PBT and a blocking step in 1.5% western blocking reagent/PBT for 30 min, embryos were incubated overnight at 4°C with primary sheep anti-DIG antibodies (1:400 dilution, Roche), and then for two h at room temperature with secondary donkey anti-sheep antibodies (1:500 dilution) conjugated to Alexa647 fluorophores (Invitrogen). Nuclei were stained with DAPI (4′,6-diamidino-2-phenylindole, dihydrochloride, Invitrogen), which was added to the secondary antibody solution. Embryos were mounted in Prolong Antifade (Invitrogen), and imaged 5–10 days after they were mounted.

**Image acquisition and analysis**

Fluorescently labeled embryos were imaged using a Leica TCS SP2 AOBS confocal scanning microscope (Leica Microsystems, Germany) with a 40× oil immersion lens for overviews of embryonic hybridization signals, and also with a 63× oil-immersion lens with a 2.3 optical zoom for high-resolution visualization of regions of interest. A 633-nm laser was used to excite Alexa 647 fluorophores. Photomultiplier tube gain was adjusted to non-saturated intensity levels. Image stacks of 6-μm depth were taken at z-step intervals of 0.25 μm. Images were corrected for Z-axis chromatic aberration. 63× magnification image stacks (2.3 zoom) were deconvolved utilizing AutoDeblur software (MedicaCybernetics), and further processed in Volocity (Improvision, Perkin Elmer), a 3D rendering and image analysis program. In Volocity, high-resolution punctate object counts in small regions were done as follows. For *Scr* FISH images, objects in a cylindrical volume with an x–y diameter of 15 μm and z-depth of 6 μm were counted. For *abd-A* images, objects in a volume that had an x–y diameter of 20 μm and z-depth of 6 μm were counted. These volumes included approximately 8–10 cell volumes. To define punctate object numbers, first, we set minimal pixel intensity thresholds to 8–10 (of a maximum of 255 in 8-bit images); second, we excluded objects smaller than 0.065 μm³ (eight voxels); and third, objects with two or more intense centroids were separated and counted as multiple objects.

**Determining k-mer uniqueness**

We designed an algorithm for determining the k-mer-uniqueness of large sets of sequences. While we focus here on *D. melanogaster*, our tool is applicable for any organism with a sequenced genome. Preexisting algorithms, such as those similar to BLAST (11,12),
uncompressed suffix arrays (13), and variants of the compressed suffix array, including the Burrows-Wheeler Transform (14–17), are not optimized to determine the location and frequencies of repeated sequences in potential FISH probes.

We enumerate all $k$-mers of a genome into a list, sort this list, count repeated sequences that are 100% matches elsewhere in the genome, and output a database of all repeated elements along with the number of times they appear in the genome. We then search this new list in logarithmic time by using the standard bisection algorithm. The above algorithm requires $O(n \log n)$ time and $O(n)$ space, where $n$ is the size of the genome. Many computers do not have $O(n)$ space in memory to store large genomes and querying parts of the genome from disk would require orders of magnitude more time. Thus we sample the genome into m bins and perform a disk-driven radix sort where each bin is sorted in memory. While this may take longer than if all data were in memory, it is more tractable given the memory constraints of current desktop computers. We sample 1% of the genome’s $k$-mers and determine approximate frequencies. Using the frequencies, we create a binary tree with m leaves that partitions $k$-mers such that each leaf has approximately the same number of $k$-mers. This allows us to output - lists $A_1$, $A_2$, ..., $A_m$ of nearly identical lengths, such that all $k$-mers in list $A_i$ precede all the $k$-mers in list $A_j$ if $i < j$. Each of these smaller lists is sorted and then combined to form a final sorted list. This algorithm takes time $O(m n/m \log n/m) = O(n \log n/m)$ time and more importantly $O(n/m)$ space. Since we break the problem into $m$ sub-problems, we can run our algorithm on a single desktop (since memory requirements are reduced) or across a cluster of computers (since each bin can be sorted in parallel). For large genomes we typically use $m = 1024$.

Note that our method exploits the fact that $k$-mer repeat locations are not considered. Our algorithm also exploits the fact that we are only interested in repeated sequences. Thus, all unique sequences can be discarded from the final database. The resulting output set of $k$-mers is typically 1–3 orders of magnitude smaller than the original genome, depending on organism and value of $k$. We could create the repeat $k$-mer database for the D. melanogaster genome in roughly 1 h of computation time (even faster on a parallel machine) with a memory footprint of less than 150 MB. The human genome is more demanding, but could be completed in less than 3 h on an 8-processor machine with a memory footprint of less than 1 GB. For all organisms and all values of $k$ that we have explored, the operator queries using the RepeatMap window that determine the position and number of repeated sequences in a potential 5-kb probe require less than a second, faster than any other method. Graphical outputs of the program for two example D. melanogaster probe sequences are shown in Figure 2.

Documentation, open source software, and UCSC Genome Browser tracks are available at http://cbio.mskcc.org/~aarvey/repeatmap/. This site also provides access to genome-wide searchable repeat $k$-mer libraries for fly (D. melanogaster), mouse (Mus musculus), and human. We also provide open-source software so that users can generate searchable repeat $k$-mer libraries from the genome of any other organism.

RESULTS

We hypothesized that a principal cause for background signals in RNA FISH protocols is hybridization to small (e.g. 20–30 bp) complementary repeat sequences in off-target transcripts. Such repeats are common in cellular RNAs. For example, of the 22797 different stable RNAs that are produced by the D. melanogaster genome and represented in cDNA libraries (18), 6995 (31%) have 20 mers that are repeated more than five times elsewhere in the genome, and 3318 (15%) have 20 mers that are repeated more than 100 times (Figure 1). However, almost all of these full-length RNAs have lengthy sub-regions that lack repeats larger than 20 nt (Figure 1).

To test the noise provided by small repeated sequences, we designed pairs of haptenylated probes against mRNA from two different genes, so that each gene had a ‘unique’ and a ‘non-unique’ probe (Figure 2). The detection and mapping of the repeats was accomplished with the program RepeatMap, a rapid algorithm for identifying the number of small genomic repeat sequences in potential probes (‘Materials and Methods’ section). This program is open source, and can be downloaded at http://cbio.mskcc.org/~aarvey/repeatmap/downloads.html. Simple graphical documentation to assist in using the program is also available at this site. The program searches genomic repeat libraries that are maintained on the repeatmap.ucsd.edu server.

![Figure 1](https://example.com/figure1.png)

Figure 1. A large fraction of RNA FISH probes contain short non-unique regions. The Y-axis shows the fraction of D. melanogaster RNA-coding sequences that contain genomic repeated sequences of size 20 nt (20-mers) or greater. Sequences have been grouped into full-length mRNA-coding (cDNA) sequences, or subsets of such coding sequences that are of lengths 2000, 1000 or 300 nt, respectively. The X-axis shows the number of 20-mer repeats that are present at a given probe sequence length. For example, about 15% of full-length cDNA sequences have 20-mer or larger repeats that are found more than 100 times in the genome.
We tested two antisense probes for transcripts from the Sex combs reduced (Scr) and abd-A genes, with the positions of probe regions denoted beneath. The shaded boxes represent the open reading frames of the respective mRNAs. Below the non-unique probes is shown the graphical output from the RepeatMap program, set to detect repeated sequences of 20 nt. The Y-axis log10 scale shows the number of times a k-mer repeat (k = 20) is found elsewhere in the D. melanogaster genome. The X-axis plot shows the nucleotide position of the repeat within the probe sequence.

**Figure 2.** Scr and abd-A probes with and without 20-mer repeats. The lines at the top represent mature mRNA sequences for the Scr (A) and abd-A (B) genes, with the positions of probe regions denoted beneath. The shaded boxes represent the open reading frames of the respective mRNAs. Below the non-unique probes is shown the graphical output from the RepeatMap program, set to detect repeated sequences of 20 nt. The Y-axis log10 scale shows the number of times a k-mer repeat (k = 20) is found elsewhere in the D. melanogaster genome. The X-axis plot shows the nucleotide position of the repeat within the probe sequence.

Unique and non-unique probes

We tested two antisense probes for transcripts from the Hox gene Sex combs reduced, (Scr). The probes were each ~350 bp, and mapped adjacent to each other in the 3'UTR (Figure 2A). One of the probes contains no 20-mer perfect matches to sequence elsewhere in the D. melanogaster genome, whereas the other has regions that are repeated elsewhere in the genome. Specifically, the non-unique probe contains a sequence complementary to a 20–24-nt repeat region found at 67 other sites in the D. melanogaster genome, and 51 of these repeats map in the sense strand of protein- or UTR-coding sequences of other mRNAs (Supplementary Table S1). Of the 24 off-target genes with homology to the non-unique Scr probe whose expression patterns have been determined by FISH, 24 are expressed in embryos at the same stages as abd-A, either in patterns or ubiquitously throughout all embryonic cells (Supplementary Table S1). The unique probe was nearly identical to the non-unique probe, except that the region complementary to the repeat at the 5'-end of the sequence was not included (Figure 2B).

In early D. melanogaster embryos, Scr transcription is limited to the primordia of the labial and first thoracic segments (19). We used FISH to detect Scr transcripts in stage 6 embryos, where transcription is limited to a stripe of cells just posterior to the cephalic furrow. Figure 3A and B show images of whole mount embryos obtained from the unique and non-unique Scr probes. At low resolution, both unique and non-unique probes detected Scr transcripts at embryonic stage 6 posterior to the cephalic furrow, but the unique probe signal was strikingly better at revealing the canonical Scr transcription domain. In stage 11 D. melanogaster embryos, the abd-A expression pattern is limited to the anterior abdominal segments 1–7 (20). At low resolution, the unfragmented unique and non-unique probes detected the pattern of abd-A transcripts at stage 11 in the abdomen, with the unique probe signal showing less background signal than the non-unique probe outside the abdomen (Figure 3C–F).
Quantitation of Unique and Non-unique Scr and abd-A FISH probes

An even more dramatic illustration of the difference between unique and non-unique probe signals is seen in high-resolution images with the two 350-bp Scr probes (Figure 4A). The unique Scr probe yielded intense punctuate signals in labial/first thoracic cells, where Scr transcripts are normally expressed, and this probe yielded only a few signals of similar size and intensity in anterior head cells that are outside the Scr expression region. The non-unique Scr probe yielded similar numbers of intense punctuate signals in both labial/first thoracic cells and anterior head cells (Figure 4A). Quantitative analysis of the number of punctuate signals detected by the Scr probes shows that the unique probe is much more specific (Figure 4A and Supplementary Figure S1). The unique Scr probe was closely examined on nine high-resolution images in regions of constant volume, each encompassing 8–10 cells (Figure 4A). The unique probe yielded an average number of 334 punctate signals inside groups of cells that were within the Scr transcription domain, but only eight punctate signals inside groups of cells outside the Scr transcriptional domain. Staining with the non-unique Scr probe on 10 high-resolution images yielded an average number of 941 punctate signals in groups of cells inside the Scr transcription domain, and 858 punctate signals in groups of cells outside the Scr transcription domain. Representative images are shown in Figure 4A. The unique Scr probe had a signal-to-noise ratio of 42 to 1, while the non-unique probe signal-to-noise ratio was only 1.1 to 1 (Figure 4A). Thus we conclude for the non-unique probe, that at least half of the signals within the Scr expression domain, and nearly all the signals outside the Scr expression domain, represented off-target hybridizations. Our previous studies indicate that most of the punctate signals, whether on-target, or off-target, represent individual cytoplasmic RNA transcripts (5).

High-resolution images of signals obtained in abdominal cells versus thoracic cells with unique and non-unique abd-A probes allowed a quantitative measure of the difference in specificity between these two 1.5-kb probes (Figure 4B,C). The unique (unfragmented) probe detected an average of 412 punctate signals in regions of constant size (∼10 cells) in abdominal segment 7 (n = 8), but an average of only six punctate signals in thoracic regions of the same size (n = 6). The non-unique abd-A probe detected an average of 535 punctate signals in similar sized regions in abdominal segment 7 (n = 7), and an average of 141 such signals in thoracic regions of the same size (n = 5). The unique abd-A probe had a signal-to-noise ratio of 70:1, while the non-unique probe ratio was 4:1 (Figure 4B).

We wished to test whether fragmentation of probes would influence specificity or sensitivity under our conditions. Many previous FISH protocols use tiled or fragmented nucleic acid probes to enhance penetration of probes into a fixed tissue (1,6,7,21–23). Under some conditions, fragmentation could increase sensitivity by 3- to 10-fold (21,22). However, new fixation and treatment methods appear to be less dependent on probe size (5), perhaps rendering fragmentation unnecessary. Furthermore, smaller fragment size may be more permissive to off-target hybridization, since short regions of
similarity would have larger contribution to the hybridization free energy than short regions in longer probes. To test this, we carbonate-fragmented abd-A probes and processed images in a manner identical as for the unfragmented abd-A probes. Interestingly, the use of fragmented probes resulted in a slightly higher level of off-target FISH signals outside the region of expression for abd-A (Figure 4C).

To quantitatively interpret high-resolution fluorescence signal, we need to ensure that we are not sacrificing sensitivity in exchange for specificity. That is, even though we improve the signal-to-noise ratio, it is possible that we are also reducing signal sensitivity. Potential loss of sensitivity is an argument as to why large probes and fragmentation may still be necessary. However, we find that no sensitivity is lost (Supplementary Figure S1). If the signal is interpreted as noise + signal where noise is estimated from signal outside the expression region, then it becomes clear that any additional ‘signal’ is really a summation of the additional noise (Supplementary Figure S1). This suggests there is no advantage, at least in D. melanogaster embryos, to fragmentation of relatively large synthesized RNA probes, as long as the probes are less than 2 kb in length. The same conclusion was reached by Lecuyer et al. (24).

**DISCUSSION**

We have studied the effects of short repeated sequences on FISH signals under conditions of very high sensitivity. Our results indicate that avoiding even short repetitive sequences, even in long RNA probes, is essential to ensure specificity when doing experiments at the resolution and sensitivity that allow detection and quantification of individual RNA molecules in fixed tissues (5). If single molecule FISH studies are to be used in a quantitative setting, the signal-to-noise ratio needs to be much higher than those provided by probes with even small repeats, which are common in full-length mRNA-coding sequences (Figure 1).

To assist in the detection of small repeats in potential FISH probes, we provide a software program, RepeatMap (see ‘Materials and Methods’ section) that will search repeat sequence databases and display the location and number of repeated sequences. We also provide all code to generate easily searchable databases of repeats in any genome or sets of genomes. Repeat annotations are likely to assist FISH in the unambiguous detections of a particular strain of bacteria, virus, or parasite in genetically diverse populations. There are also other uses of sequence uniqueness annotations, including sequencing experiments (where short reads need to be mapped back to unique regions), other hybridization methods (e.g. DNA FISH, and microarrays), and discovery of functionally relevant sequences [e.g. RNAi targets (12)].

Our results also suggest that the practice of fragmenting long haptenylated probes may be unnecessary for whole-mount *Drosophila* embryos. It would be interesting to explore the consequences of fragmentation (or lack thereof) in tissues from other organisms and fixed in other manners. We would also like to explore the utility of our methods to tissues where sensitivity is still quite challenging, such as paraffin-embedded histological sections.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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