Revealing details: whole mount microRNA in situ hybridization protocol for zebrafish embryos and adult tissues

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Biology Open 1, 566–569
doi: 10.1242/bio.2012810

Summary
Non-coding microRNA (miRNA) molecules bind their target mRNAs and thereby modulate the amount of protein produced. To understand the significance of a potential miRNA-mRNA interaction, temporal and spatial information on miRNA and mRNA expression is essential. Here, we provide a detailed protocol for miRNA whole mount in situ hybridization. We introduce the use of Morpholino based oligos as antisense probes for miRNA detection, in addition to the current “gold standard” locked nucleic acid (LNA) probes. Furthermore we have modified existing miRNA in situ hybridization protocols thereby improving both sensitivity and resolution of miRNA visualization in whole zebrafish embryos and adult tissues.

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Key words: MicroRNA, in situ hybridization, Zebrafish

Introduction
Since the discovery of the first regulating small RNAs (later classified as microRNAs), lin-4 and let-7, in the nematode Caenorhabditis elegans (Lee et al., 1993; Reinhart et al., 2000), ongoing high throughput cloning approaches have led to the discovery of hundreds of miRNAs. MiRNAs inhibit protein expression by binding mainly to 3'UTR sequences of messenger RNA molecules (reviewed by Bartel, 2009). Functional studies thus far have implicated miRNAs to function in various processes during embryonic development and disease (reviewed by Hagen and Lai, 2008). Since miRNAs do not translate into protein, the identification of the tissues and cell types in which specific miRNAs are expressed relies heavily on in situ hybridization detection techniques. To date, LNA probes complementary to mature miRNA sequences are the most commonly used oligos for miRNA detection from various species for in situ hybridization as well as for probing miRNAs on microarray platforms (Ason et al., 2006; Darnell et al., 2006; Kloosterman et al., 2006; Oberschützer et al., 2007; Mishima et al., 2009; Pase et al., 2009; Pena et al., 2009; Goljanek-Whysall et al., 2011; Preis et al., 2011). Nevertheless, alternatives for LNA based probes have been reported (Soo et al., 2011).

Current miRNA in situ hybridization protocols provide limited spatial resolution of low-abundant miRNAs. Diffusion of small miRNAs into the tissue after formaldehyde fixation has been suggested to cause these limitations in spatial resolution (Pena et al., 2009). We have applied 1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide (EDC) fixation on zebrafish (Danio rerio) embryos and adult zebrafish tissues. EDC crosslinks the 5’-phosphate of miRNAs, which is formed during miRNA maturation, to amino groups in the protein matrix. Here, we present a miRNA whole mount in situ hybridization method allowing a sensitive detection with high spatial resolution of miRNAs expression in both zebrafish embryos and adult tissues.

Materials and Methods
A step-by-step protocol with a summary of required materials, details on reagent preparation and additional procedures are available at http://www.hubrecht.eu/research/bakkers/protocols.html and we strongly recommend that these data are consulted in combination with the information provided below.

Embryonic and adult tissue preparation
Paraformaldehyde fixation
Zebrafish embryos were staged live and subsequently fixed in PFA1 (4% paraformaldehyde, 4% sucrose in PBS), overnight (o/n) at 4°C in 4 ml glass vials on a 3D rocker.

Adult zebrafish tissues were dissected and fixed in PFA1 for 48 hours at 4°C, while on a 3D rocker.

1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide fixation
PFA1 fixed embryos and adult tissues were washed three times in PBS containing 0.01% Tween-20 (PBS-T) at room temperature (RT), for 10 min on a 3D rocker to remove residual PFA1. Subsequently, all samples were washed three times in fresh 1-methylimidazole buffer (1-MIB) (1% 1-methylimidazole, 300 mM NaCl in water pH 8.0 (HCl)) for 15 min while on a 3D rocker at RT. Next, embryos were fixed in fresh 0.16 M 1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide (EDC) diluted in 1-MIB (pH 8.0). Upon addition of EDC fixative embryos were first fixed for 2 hours at RT followed by o/n fixation at 4°C, while on a 3D rocker.

10 min (embryos) or 20 min (adult tissues) per solution. The last step (100% water pH 8.0) was then removed under a gentle flow of water.

Dehydration
Embryos and adult tissues were dehydrated through a series of methanol (MeOH) diluted in PBS-T (25% MeOH, 50% MeOH, 75% MeOH, 100% MeOH) at RT for 10 min (embryos) or 20 min (adult tissues) per solution. The last step (100% water pH 8.0) was then removed under a gentle flow of water.
MeOH) was repeated once, after which embryos and adult tissues were stored at −20°C until use. However, when in situ hybridization procedure 1 was initiated on the same day after dehydration, embryos and adult tissues were first incubated in the final 100% MeOH solution at RT for at least two hours to ensure full dehydration of the embryos.

All dehydration washes were performed on a 3D rocker. Embryos remained in 4 ml glass screw top vials and adult tissues in 10 ml plastic screw top tubes.

**MicroRNA in situ hybridization**

**Rehydration**

Dehydrated embryos stored in 100% MeOH at −20°C were divided per sample into 1.5 ml Eppendorf tubes (<30 embryos). Adult tissues were divided into 2 ml Eppendorf tubes (<3 hearts, <3 fins). Samples were re-hydrated through a series of MeOH in PBS-T (75% MeOH, 50% MeOH, 25% MeOH). Samples were washed (embryos 10 min, tissue 20 min) in each solution, while on a 3D rocker at RT. Afterwards samples were placed in PBS-T and washed four times in PBS-T while on a 3D rocker for 10 min per wash at RT (embryos and adult tissue).

**Protease K treatment**

After the re-hydration all samples were treated with protease K to facilitate infiltration of the probes into the tissue. Protease K treatment should be optimised for the embryonic stage, type of tissue used and when using a new batch of enzyme. We have listed an overview of the concentrations, incubation times and temperatures we used in supplementary material Table S1. Embryos were incubated in 1.5 ml tubes containing 500 μl protease K (10 μg/ml) diluted in PBS-T. Adult tissues were incubated in 1 ml protease K solution in 2 ml tubes. Samples should not move during the protease K treatment.

After protease K treatment embryos and adult tissues were re-fixed in 4% PFA2 (4% paraformaldehyde in PBS, pH 7.4) for 10 min (embryos and adult tissue) at RT. Following re-fixation samples were washed five times for 10 min per wash in PBS-T at RT.

**Hybridization**

Next, PBS-T was replaced by hybridization buffer (HYB+, 50% deionized formamide, 5× saline sodium citrate (SSC), 0.1% Tween-20, 9.2 mM citric acid) supplemented with 50 μg/ml Heparin and 0.5 mg/ml RNA (HYB+) ([500 μl HYB+ in 1.5 ml Eppendorf tube] and 1 ml HYB+ for adult fins (2 ml tube)). All samples were placed in heat blocks or water baths to pre-hybrdize for minimally 2 hours at the corresponding hybridization temperature (supplementary material Table S2). Subsequently, samples were incubated o/n with probes diluted in HYB+ at the corresponding hybridization temperature. For embryos in 1.5 ml tubes and adult hearts in 2 ml tubes we use 500 μl of probe dilution. For caudal fins in 2 ml tubes we used 1 ml of probe dilution.

After o/n probe hybridization, samples were washed once in pre-heated HYB- for 10 min at the hybridization temperature. Afterwards, all samples are taken through a series of pre-heated HYB- diluted in 2× SSC-T (SSC containing 0.1% Tween-20), first 75% HYB-→2×SSC-T, then 50% HYB-→2×SSC-T and finally 25% HYB-→2×SSC-T. For each step during this series, embryos and adult tissues were incubated for 15 min at the probe hybridization temperature. Afterwards, all samples were incubated 15 min in pre-heated 2×SSC-T at the hybridization temperature followed by two 30 min incubations in pre-heated 0.2×SSC-T at the hybridization temperature. All samples were subsequently taken through a graded series of 0.2×SSC-T diluted in PBS-T, first 75% 0.2×SSC-T/ PBS-T, then 50% 0.2×SSC-T/PBS-T and finally 25% 0.2×SSC-T/PBS-T, all at RT. All samples were subsequently washed twice in PBS-T at RT for 10 min before being transferred to blocking buffer (BB, 2% sheep serum, 2 mg/ml BSA in PBS-T) for minimally 1 hour at RT while on a 3D rocker. After blocking, all samples were incubated o/n in BB containing either pre-incubated anti-digoxigenin (Roche, 1:5000) or anti-fluorescein (Roche, 1:5000) antibodies while on a 3D rocker for 10 min at RT. Subsequently samples were re-fixed through a series of methanol (MeOH) diluted in PBS-T (25% MeOH, 50% MeOH, 75% MeOH, 100% MeOH) at RT while on a 3D rocker for 10 min (embryos) or 20 min (adult tissues) per solution. Dehydration was followed by two washes in 100% MeOH for at least 30 min each to ensure all residual PBS-T was removed. Finally the MeOH was replaced by Murray’s solution (benzylbenzoate/benzylalkohol 2:1). The Murray’s solution cleans the tissue and can be used for long-term storage of the samples at 4°C in the dark in polypropylene tubes. Imaging of the miRNA expression pattern was performed in Murray’s solution after an o/n incubation at 4°C.

**Antisense probes**

All LNA probes (Exiqon) were diluted in milli-Q water to a stock concentration of 10 μM and stored at −80°C. The double Digoxigenin labelled (3′ and 5′) miR-23 LNA probe (Exiqon) was diluted 700 times in hybridization buffer as a working dilution. All other LNA oligonucleotides were labelled with digoxigenin-11-dUTP (Roche) at the 3′ end using a terminal transferase kit (Roche) and purified using MicroSpin G-25 Columns (Amersham Biosciences). Single 3′ end labelled LNA probes were diluted 200 times in hybridization buffer (HYB+) as a working dilution.

Carboxyfluorescein (CF)-labelled Morpholino oligos (MO) (Gene Tools) were diluted in milli-Q water to a stock concentration of 1 mM and stored at −20°C. All CF-labelled MO probes were diluted 50,000 times in HYB+ as a working dilution. All LNA and MO probe dilutions were re-used up to 10 times.

**Morpholino injections**

MiR-23 MO (Gene Tools) was diluted and injected as described previously (Lagendijk et al., 2011). Sequence of the CF-labelled miR-124 MO used for miR-124 knock-down was as follows: CF-labelled miR-124 MO: 5′-tgccattcagcggctttac-3′

For injection, the CF-labelled miR-124 MO was diluted to 0.2 mM, of which 0.1 μl was injected at the one cell stage.

**Antibody staining and sectioning**

Tropomyosin antibody staining and subsequent plastic sectioning of miR-23 in situ stained embryos was performed as described previously (Lagendijk et al., 2011).

**Results and Discussion**

**Morpholino oligomers detect miRNA expression by whole mount in situ hybridisation**

While LNA oligos are widely used as antisense probes to detect miRNA expression, their use as knock-down reagent in zebralish is less versatile (Kloosterman et al., 2007). Morpholino oligos (MO) have been shown to bind miRNAs with high affinity in vivo resulting in efficient knock-down of miRNA expression in zebralish embryos (Kloosterman et al., 2007), however their use as antisense probes had not been reported. Therefore we used carboxyfluorescein(CF)-labelled MOs, overlapping with the mature miRNA sequences of different miRNAs, as miRNA probes. In situ hybridisation on 4 day old zebralish larvae using these CF-labelled MOs as probes and anti-fluorescein antibodies for signal amplification and detection resulted in specific staining patterns (Fig. 1A–D). The observed staining patterns were indistinguishable from previously published patterns using LNA probes (Wienholds et al., 2005; Kloosterman et al., 2006). Next we investigated whether a CF-labelled MO could still function as a
knock-down reagent. Therefore we performed miR-124 in situ hybridization on embryos previously injected by CF-labelled MO complementary to miR-124. As a result we observed a loss of mature miR-124 expression (supplementary material Fig. S1) demonstrating that CF-labelled MOs targeting miRNAs can still function as miRNA knock-down reagents. To summarize, we conclude that CF-labelled MOs can be used for both miRNA detection and knock-down purposes. Since many researchers already have Morpholino oligos in-hand for their knockdown experiments, it would be both convenient and economical to be able to use these same oligos for both knockdown and in situ hybridization.

Whole mount EDC fixation improves sensitivity and spatial resolution in miRNA detection

Both LNA- and MO-based probes allow detection of robust miRNA expression in whole mount zebrafish embryos of late developmental stages (Fig. 1) (Wienholds et al., 2005; Kloosterman et al., 2006). However, their use in detecting low-abundance miRNA expression, such as during early stages of embryo development, can be hampered by the appearance of non-specific background signals. Fixation of paraffin sections in 1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide (EDC) has been reported to prevent miRNA diffusion by crosslinking miRNA 5′ ends with amino groups in the protein matrix and thereby improving spatial resolution (Pena et al., 2009). To investigate whether EDC fixation could improve detection and spatial resolution of miRNA expression signals in whole mount in situ hybridization we compared fixation of zebrafish embryos in PFA with PFA+EDC. We used both digoxigenin (DIG) labelled LNA based probes as well as CF-labelled MO based probes to detect miRNAs at various developmental stages. In all cases we observed that embryos fixed in PFA+EDC showed a significant improvement in spatial resolution of the miRNA expression pattern when compared to PFA fixation alone (Fig. 2A–R). Non-specific background staining was very low in PFA+EDC fixed embryos, allowing the detection of specific expression patterns not observed with PFA fixation alone (Fig. 2, arrowheads). Together, these results demonstrate that the combination of both EDC and PFA fixation results in a robust improvement in spatial resolution and sensitivity of miRNA expression detection by whole mount in situ hybridization.

To verify whether the observed staining pattern in PFA+EDC fixed embryos resulted from specific miRNA expression, we performed a similar miRNA in situ hybridization analysis on embryos in which production of the mature miRNA was blocked. Indeed all miR-23 signals were lost in embryos injected with an antisense MO targeting miR-23 (supplementary material Fig. S2a,b). In addition, we tested whether it was possible to perform immunohistochemistry on embryos that had been processed for PFA+EDC fixation and in situ hybridization. Using an α-tropomyosin antibody we observed robust levels of tropomyosin in the myocardium of PFA+EDC fixed embryos processed for in situ hybridization prior to the immunohistochemistry procedure (supplementary material Fig. S2c).

EDC fixation reveals specific miRNA expression in adult zebrafish tissues

MiRNA expression studies in adult tissues are mostly performed by in situ hybridization on paraffin embedded sections (Obernosterer et al., 2006; Obernosterer et al., 2007; Pena et al., 2009). Here we investigated whether EDC fixation would
probes can be used as an alternative for LNA based probes. This
embryos and adult tissues. First we have shown that MO based
in situ miRNAs during tissue regeneration and disease.
hybridization for the detection of miRNA expression in adult
in situ with a probe (Fig. 3G). The application of whole mount
hearts was not detected in control hearts that were not incubated
fixed tissues (Fig. 3A–F). The specific staining pattern in adult
PFA+EDC fixed tissues, which was reduced or absent in PFA
caudal fins and observed a specific staining pattern for miR-23 in
m.

Fig. 3. Whole mount miRNA in situ hybridization on adult tissues.
(A–D) MiR-23 is expressed in the bulbus arteriosus (b) of zebrafish adult hearts
fixed in PFA (A,C) and PFA+EDC (B–D). PFA+EDC fixed tissues revealed a
more detailed miR-23 expression pattern with a ring-like expression (arrowhead
in B) at the inflow area of the atrium (a) and in coronary arteries (arrowheads in D)
overlying the ventricle (v). (E,F) Robust miR-23 expression was detected in
joint structures of PFA+EDC fixed caudal fins (arrowheads in F) versus more
diffuse miR-23 expression in caudal fins fixed in PFA only (E). (G) Negative
control heart was fixed in PFA+EDC but not incubated with a probe. Scale bars represent 100 μm.

allow detection of miRNA expression by whole mount in situ
hybridization on adult tissues. We used adult zebrafish hearts and
caudal fins and observed a specific staining pattern for miR-23 in
PFA+EDC fixed tissues, which was reduced or absent in PFA
fixed tissues (Fig. 3A–F). The specific staining pattern in adult
hearts was not detected in control hearts that were not incubated
with a probe (Fig. 3G). The application of whole mount in situ
hybridization for the detection of miRNA expression in adult
tissues will be a valuable new tool to address the function of
miRNAs during tissue regeneration and disease.

In conclusion we have optimised a method for whole mount in situ
hybridization to detect non-coding miRNAs in zebrafish
embryos and adult tissues. First we have shown that MO based
probes can be used as an alternative for LNA based probes. This
allows the use of a single CF-labelled MO as an antisense probe for
in situ hybridization and as a knock-down reagent. Second we
have applied EDC fixation to improve sensitivity and spatial
resolution of the whole mount in situ hybridisation method.

Acknowledgements
We thank Wigard Kloosterman and members of the Bakker’s
laboratory for their suggestions during this work, and Gilbert
Weidinger and Leonie Huitema for their judgment on some of the
expression patterns. A.K.L. was supported by a Concordia fellowship
by stichting vrienden van het Hubrecht. Research in J.B.’s laboratory
was supported by the Netherlands Organization for Scientific
Research (NWO/ALW) grant 864.08.009.

Competing Interests
Jon D. Moulton is employed by Gene Tools LLC and stands to profit
by increased demand for Morpholinino oligos. The other authors do
not declare any competing interests.

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