Out with the old, in with the new: reassessing morpholino knockdowns in light of genome editing technology

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ABSTRACT

Morpholino oligomers have been used widely and for many years in the zebrafish community to transiently knock down the function of target genes. It has often been difficult, however, to reliably discriminate between specific and non-specific effects, and thus generally accepted guidelines to control for morpholino side effects do not exist. In light of recent methodologies to generate mutant lines in virtually any zebrafish gene, we discuss these different approaches with a specific focus on how the first description of a loss-of-function phenotype in zebrafish should be accomplished.

Initially, the genetic analysis of zebrafish development and physiology was dominated by mutants identified in small- and large-scale forward genetic screens (Chakrabarti et al., 1983; Driever et al., 1996; Haffter et al., 1996). Whereas forward genetics was instrumental in establishing zebrafish as an additional vertebrate model system, progress was hampered by the fact that there was no reliable technology to carry out reverse genetics in this model. TILLING (a reverse genetics approach based on high-throughput sequencing of ENU-mutagenized fish) was introduced only in 2002 (Wienholds et al., 2002) and requires considerable up-front investment in logistics and infrastructure. The introduction of morpholinos (MOs) in frogs (Heasman et al., 2000) and zebrafish (Nasevicius and Ekker, 2000) as an antisense reagent to transiently knock down gene function was therefore greeted with considerable excitement, as it appeared to fill a real void in the toolbox. Since its inception, countless studies using this technology have been published, including some using MOs to knock down maternally deposited transcripts to circumvent the generation of maternal-zygotic mutants, and others using caged MOs, which allow for inducible release of these antisense reagents (Shestopalov et al., 2012).

The MO antisense technology is based on nucleic acid bases that are linked to morpholine rings and a non-charged phosphorodiamidate backbone. The rationale for this design was that MOs would not bind electrostatically to protein, hence causing less toxicity, while at the same time being resistant to nucleases (Summerton, 2007). MOs are injected into early zebrafish embryos using standard techniques. Commonly, they are ~25-mers designed to be an exact antisense match against the region surrounding the first translated ATG (to block translation) or against a splice donor or acceptor site (to interfere with precursor mRNA splicing). It quickly became apparent that some MOs could work extremely well, and there are many MO phenotypes that efficiently mimic mutant phenotypes without any noticeable side effects. However, it has also become clear that MOs can lead to artifacts and that for many MOs the phenotypes caused by specific binding to the intended target RNA are difficult to separate from those caused by the non-specific binding to unintended targets (Eisen and Smith, 2008). In fact, a simple calculation suggests that binding to targets other than the intended precursor or mature mRNA is likely. A zebrafish embryo contains ~500 ng of RNA, 2-5% of which is translatable (25 ng) (A. Giraldez, personal communication; see also Davidson, 1986). Assuming that at any given time there are more than 104 different mRNA species present in a cell (Davidson, 1986), and that those transcripts are equally represented among the 25 ng of mRNA, only 2.5 pg of a specific mRNA species is available for targeting. Injections typically deliver ~1 ng of MO, often more. Assuming further that the target mRNA has an average length of 1.25 kb, whereas the MO is a 25-mer, this equates to a 2×104-fold molar excess of MO versus target mRNA. It is therefore most likely that this vast excess of MO will bind other RNA or other macromolecules. This situation would not be such a serious problem if there were reliable ways to distinguish specific from non-specific effects. However, this is not the case, and one can at best only show that MOs affect the target sequence; non-specific effects cannot easily be identified, even when using mRNA rescues (see Del Giacco et al., 2010; Tao et al., 2011). The literature now contains several examples in which developmental delay, defects in organ asymmetry and pericardial edema (among many other ‘phenotypes’) are attributed to knocking down a specific gene, but in which subsequent generation of a mutation in that gene revealed a very different phenotype, and often no phenotype at all. Recent examples include mutations in sox18, nr2fla and prox1a/b, all genes that had been reported to show morphant phenotypes within the lymphatic vasculature, whereas the mutant alleles do not (van Impel et al., 2014).

In several cases it has been possible to circumvent some of the non-specific phenotypes by suppressing p53 activity (Robu et al., 2007), which can reduce the ectopic cell death caused by non-specific MO effects; however, this approach has its own caveats as it effectively generates a phenotype not on a wild-type, but on a p53-deficient, background. Applying such drastic corrective measures to allow a phenotypic analysis raises a number of questions that cannot be easily addressed.

Two surprisingly efficient alternatives for reverse genetics have been recently implemented in zebrafish (Chang et al. 2013; Huang et al., 2011; Sander et al., 2011; Hwang et al., 2013; Zu et al., 2013) and other organisms (Beumer et al., 2008; Tesson et al., 2011; Yang et al., 2013). TALE nucleases and the Crispr/Cas9 system are very efficient at generating mutations. As both techniques have been reviewed extensively (Auer and Del Bene, 2014), we will restrict the discussion here to comparing the principles of the MO and TALEN/Crisp approaches.

First, it should be noted that – like MOs – the implementation of these new technologies can be carried out in virtually any lab. Whereas TILLING really only makes sense for those willing to analyze large numbers of samples and genes, TALENs and Crispr

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do not require a substantial investment and, once established, can be used to generate targeting constructs within 1 (Crispr) to 2 (TALEN) weeks.

Second, as TALENs and Crisprs affect genomic DNA, rather than RNA transcripts, their molecular effect can be determined at the single embryo level (which is more difficult with MOs) to obtain a clear phenotype-genotype correlation. Of course, such an approach requires caution, as these nuclease-injected embryos are most likely to be mosaic for the resulting genomic lesions. Furthermore, TALEN and Crispr constructs can sometimes be efficient enough to generate loss-of-function situations in the actual injected embryos (Dahlem et al., 2012), and so, in a minority of cases, can be used almost like an MO; injection, scoring for phenotypes and confirming that the nuclease works efficiently can be performed within a few days.

Third, the published evidence, although currently limited, suggests that the side effects of these nucleases are often negligible (Hruscha et al., 2013), even though both TALE and Crispr-Cas nucleases can bind and cleave off-target loci (Reyon et al., 2012; Fu et al., 2014). When additional mutations are introduced, they can usually be segregated away from the mutation of interest by one or two outcrosses (as with mutations identified in ENU mutagenesis screens). This specificity is of course a tremendous advantage, and very different from MOs: an MO that binds non-specifically will most likely do so in every injected embryo. Lastly, it is relatively easy to generate multiple mutant alleles in one gene (e.g. by using TALEN pairs that affect different regions of the targeted gene), thus further reducing the chance of being misled by off-target mutations.

Hence, it seems fair to say that within the last year or so, the landscape of reverse genetics in zebrafish has changed, and it has changed for the better. Anyone can now, within a few weeks, generate reagents that can be used for reverse genetic experiments that appear to be of superior reliability and that are less burdened with side effects compared with MOs. Does that mean that we should do away with MOs altogether? Not necessarily: as we pointed out in the first paragraph, there are many MOs that are very useful and that appear to work specifically. We know they work specifically because we can compare them with a mutant phenotype. We would argue that this is the criterion that should be used in most cases: if one can show that an MO phenotype is an exact replicate of a mutant phenotype, then use of this MO is be used in most cases: if one can show that an MO phenotype is an exact replicate of a mutant phenotype, then use of this MO is.

References


Tao, S., Witte, M., Bryson-Richardson, R. J., Currie, P. D., Hogan, B. M. and Schulte-Merker, S. (2011). Zebrafish prox1b mutants develop a lymphatic vasculature, and prox1b does not specifically mark lymphatic endothelial cells. PLoS ONE 6, e28934.


