A compact, in vivo screen of all 6-mers reveals drivers of tissue-specific expression and guides synthetic regulatory element design

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Abstract

Background: Large-scale annotation efforts have improved our ability to coarsely predict regulatory elements throughout vertebrate genomes. However, it is unclear how complex spatiotemporal patterns of gene expression driven by these elements emerge from the activity of short, transcription factor binding sequences.

Results: We describe a comprehensive promoter extension assay in which the regulatory potential of all 6 base-pair (bp) sequences was tested in the context of a minimal promoter. To enable this large-scale screen, we developed algorithms that use a reverse-complement aware decomposition of the de Bruijn graph to design a library of DNA oligomers incorporating every 6-bp sequence exactly once. Our library multiplexes all 4,096 unique 6-mers into 184 double-stranded 15-bp oligomers, which is sufficiently compact for in vivo testing. We injected each multiplexed construct into zebrafish embryos and scored GFP expression in 15 tissues at two developmental time points. Twenty-seven constructs produced consistent expression patterns, with the majority doing so in only one tissue. Functional sequences are enriched near biologically relevant genes, match motifs for developmental transcription factors, and are required for enhancer activity. By concatenating tissue-specific functional sequences, we generated completely synthetic enhancers for the notochord, epidermis, spinal cord, forebrain and otic lateral line, and show that short regulatory sequences do not always function modularly.

Conclusions: This work introduces a unique in vivo catalog of short, functional regulatory sequences and demonstrates several important principles of regulatory element organization. Furthermore, we provide resources for designing compact, reverse-complement aware k-mer libraries.

Background

The activity of regulatory elements is governed by transcription factors (TFs), which bind to DNA sequences averaging 6 to 9 bp in size [1] and function in combination to achieve diverse gene expression patterns. Knowledge of the vertebrate TF repertoire and individual TF binding site (TFBS) preferences is still far from complete, due to binding-site degeneracy and the challenge of annotating functional TFBSs that may occur over a million base pairs from the transcription start site (TSS) of their target gene. Even less is known about the minimal functional elements that are sufficient to drive tissue-specific expression, and how elaborate patterns result from their interplay. Deciphering the vocabulary and grammar of the vertebrate regulatory code will enable high-resolution mapping of regulatory elements, accurate interpretation of nucleotide variation in regulatory sequences, and the design of regulatory elements to deliver genes to specific tissues for research and therapeutic purposes.

Many top-down approaches are used to predict vertebrate regulatory sequences. For a subset of vertebrate TFs, motifs derived from in vitro measurements of binding affinity [2,3] can be used to identify potential TFBSs in genomic sequences. Distingushing which TFBSs are functional...
is challenging, even with the aid of evolutionary conservation [4,5] and clustering of TFBSs [6,7]. Techniques that combine biochemical assays with massively parallel sequencing, such as ChIP-seq, DNase-seq and formaldehyde-assisted isolation of regulatory elements (FAIRE)-seq, identify potential gene-regulatory elements by measuring TF binding and open chromatin in a specific cell type on a genome-wide scale [8-17]. However, these methods only coarsely locate functional sequences and are predictive in nature, requiring downstream validation. There is evidence that regulatory signals may be short (approximately 44 bp) [18], but canonical regulatory elements identified by these techniques tend to be quite long. For example, the average size of the 846 functional elements currently listed in the VISTA enhancer browser is 1,812 bp [19]. Traditional efforts to bash larger elements for core domains [18] are effective, but are time-consuming and costly.

To complement existing methods for identifying regulatory regions, it will be essential to reverse-engineer regulatory elements to reveal the fundamental units that drive complex expression patterns and the rules that govern their organization. One study that employed such an approach [20] in vitro tested 100-bp synthetic elements, each composed of tandem repeats of multiplexed 10-bp sequences, in six cell lines. Considerable differences in expression levels were observed between cell lines, consistent with the tissue-specificity of native vertebrate regulatory elements [8]. However, in vivo assays are essential to understanding how the vertebrate regulatory logic functions in a physiological environment involving multiple cell types, cellular interactions and temporal stages.

Here, we developed a bottom-up approach to characterize regulatory element vocabulary by systematically cataloging the effect of each 6-bp sequence (6-mer) on a TATA-box-containing 42-bp minimal promoter using zebrafish transgenics. To make this comprehensive screen feasible in vivo, we multiplexed the 6-mers into an ultra-compact library of 184 double-stranded constructs, each 15-bp in length. The activity of each construct was then scored in 15 tissues at 24 and 48 hours post-fertilization (hpf). We identified 27 multiplexed oligomers (15%) that produced distinct, consistent expression patterns. Four such constructs with distinct expression profiles were functionally dissected to identify even shorter 9-bp sequences that match binding motifs of known TF families, suggesting a mechanism by which they achieve tissue-specificity. Removal of these functional sequences from native zebrafish enhancers reduced their activity. Functional sequences were also used to design larger completely synthetic homotypic regulatory modules that drove strong expression to specific tissues. In contrast, heterotypic combinations of synthetic concatemers revealed a non-modular grammar.

### Results

#### Design of an ultra-compact library of multiplexed 6-mers

The number \( n_k \) of unique sequences of length \( k \)-mers is \( 4^k \), and therefore, values in the TFBS-length range of 6 to 9 bp produce an infeasible number of sequences for individual testing. On the other hand, values of \( k \) less than 6 would not be expected to cover a substantial portion of the TFBS binding spectrum. A solution to this problem is to multiplex \( k \)-mers into longer oligomers. However, the length of the oligomers is inversely correlated with the information that can be learned about individual \( k \)-mers and proportional to the follow-up testing (for example, bashing) required for identifying the active \( k \)-mers. Additionally, inefficient multiplexing (Figure 1a), exemplified by concatenating \( k \)-mers or randomly generating oligomers, results in large libraries with biased \( k \)-mer coverage. By developing mathematical and computational techniques for efficiently multiplexing \( k \)-mers into oligomers (Additional file 1), we balanced these criteria at \( k = 6 \) (\( n_k = 4,096 \)) and an oligomer length of 15 bp.

Compact \( k \)-mer libraries have been used in several applications, such as protein-binding microarrays [3], universal DNA tag systems [21], and microfluidic affinity analysis devices [22]. This type of library is constructed from dividing up a de Bruijn sequence, which contains one copy of every \( k \)-mer. This sequence can be generated from a de Bruijn graph (Additional file 2), a mathematical object that has been applied in several biological contexts, particularly read assembly [23]. For \( k = 6 \) and a length of 15 bp, such a library would contain approximately 400 oligomers.

The theory underlying de Bruijn sequences was developed for general alphabets, not DNA. With the exception of self-reverse-complementary palindromes, each \( k \)-mer has a distinct reverse complement that is presumably equivalent to a TF binding double-stranded DNA. By restricting each reverse-complementary pair to a single representative on the forward strand, we reduced the library size by approximately 50%. Our design, which we call a minimal reverse-complementary-covering (MRCC) library of order \( k \) and size \( m \), is a collection of \( m \) DNA sequences in which a single representative for each reverse-complementary pair of \( k \)-mers appears exactly once. To create an MRCC library, we designed an algorithm (Additional file 1) that computes a new type of palindrome-aware edge decomposition of the de Bruijn graph (Figure 1a) and enables us to account for reverse complementarity. In contrast to a de Bruijn sequence, which is obtained by computing a single cyclical path that traverses every edge in the graph exactly once (Additional file 2), an MRCC library is obtained by partitioning the edges into many paths. We note that for odd values of \( k \), the algorithm is similar to a previous heuristic [24] that attempts to compute an analog to the de Bruijn sequence. However, this analog does
Figure 1 In vivo regulatory screen of all 6-mers (a) Left top: naively concatenating three 6-mers creates an oligomer with multiple representatives of ATTGCC (red bars). Center and left: cartoon of a de Bruijn graph. Nodes (colored boxes) represent 6-mers; edges (arrows) represent overlap. A standard de-Bruijn-sequence library is built from one path that traverses each of 4,096 nodes once. Constructed from multiple paths, our MRCC library instead uses one representative for each pair of reverse-complementary 6-mers (green and yellow; self-reverse-complementary palindromes in blue), making it nearly 50% smaller (Additional file 1). Right: 16 of 16,384 edges shown. Our algorithm removes reverse-complementary paths (black, red) between palindrome pairs and then decomposes the remaining graph into reverse-complementary cycles. It allowed us to design an ultra-compact library of DNA sequences with uniform 6-mer coverage. (b) Schematic depicting the sub-cloning of each 15-bp multiplexed oligomer into the E1b-Tol2 vector and subsequent injection into single-cell zebrafish embryos. (c) Violin plots depicting the distribution of the expression patterns of each tissue at 24 hpf. White lines indicate the fractional expression values for the empty vector construct. (d) Scatter plot depicting the method by which we selected consistently expressed multiplexed oligomers whose expression was not significantly correlated to minimal-promoter bias. The vertical dotted line denotes the 40% fractional expression threshold that was used, whereas the horizontal dotted line corresponds to a false discovery rate-adjusted P-value of 0.05. (e) Histogram depicting the tissue specificity of the 22 uncorrelated, consistently expressed constructs at 24 hpf. The full sequence of each construct (with XhoI and BglII flanking sites) is listed below each figure. Both constructs have a 5’ GC-rich region. GFP, green fluorescent protein.
not exist for even values of $k$ larger than 2 (Additional file 1).

Using custom software that we have made available [25], we created an MRCC library of 208 DNA oligomers, each 15 bp, with uniform, single-copy coverage of all 6-mers. By exploiting 6-mers that overlap junctions between oligomers and flanking sequence, a modified version of the algorithm reduced the number to 184 (Additional file 3). This library is about 45% of the minimum size obtainable with previous algorithms that give uniform coverage.

**Multiplexed oligomer screen identifies functional units driving tissue-specific expression**

Each of the 184 multiplexed 15-bp oligomers was synthesized and cloned upstream of the 42-bp adenovirus type 5 E1b [26] minimal promoter and green fluorescent protein (GFP) in a Tol2 retrotransposon-based [27] zebrafish reporter vector [28]. Each construct was injected into single-cell zebrafish embryos, and GFP expression was monitored in 15 tissues at 24 and 48 hpf (Figure 1b). These time points coincide with the establishment of the basic vertebrate brain plan and the development of key structures. In addition, embryos were injected with an empty vector control (identical flanking sequence but no insert) to assess the contribution of the minimal promoter. A substantial portion of the MRCC library (70/184 constructs, approximately 38%, at 24 hpf) drove consistent (in >40% of embryos) expression patterns, though our design was unguided by a priori knowledge and only a small amount of information could be encoded in each oligomer. A potential explanation for this is that our constructs, like regulatory regions marked selectively by histone 3 lysine 4 mono-methylation (H3K4me1), have a higher GC content than the zebrafish genome (average 43.1% for H3K4me1 and 50% for our multiplexed oligomers, versus 36.4% genome-wide). An alternative explanation is that while the oligomers are very short, the design of our MRCC library encodes a maximum amount of sequence diversity. It is thus likely that the MRCC library contains more active sequences than random genomic sequence of the same size. Our results are consistent with a much smaller percentage of the genome being active in regulation.

We observed a diverse range of expression at both time points (Figure 1c; Additional files 2 and 3). However, there was a clear bias towards amplifying the background bias of the minimal promoter (Figure 1c; Additional file 2). By removing expression patterns that were significantly correlated with background expression (Spearman rank correlation coefficient, false discovery rate (FDR) $q < 0.05$), we identified 22 multiplexed constructs that drove GFP expression consistently at 24 hpf to at least one tissue (Figure 1d; Additional file 3). An additional five constructs produced uncorrelated and consistent expression solely at the 48 hpf time point (Additional file 2).

Tissue specificity is a common feature of regulatory elements [29]. We thus examined the tissue distribution of constructs expressed consistently in one or more tissue. Eleven out of the 22 constructs (50%) that produced consistent and uncorrelated expression at 24 hpf did so in a single tissue (Figure 1e). In contrast, several correlated constructs exhibited very broad expression patterns. For example, 1EF03 and 3EF09 produced the strongest observed signal in ten tissues across the two time points, likely due to a GC-rich sequence embedded at the 5’ end of each oligomer (Figure 1f).

We chose a subset of the consistently expressed, tissue-specific, uncorrelated constructs to pinpoint functional sequences within the constructs and the zebrafish genome: 2CD12/epidermis, 2GH08/brain, 2GH12/notochord, and 3CD05/spinal cord (Figure 2). 2GH08/brain was unique in this set in that it expressed consistently in more than one tissue: bilaterally in rostral forebrain neurons at both time points and in ganglia of the otic lateral line at 24 hpf. For comparison, representative negative embryos injected with the empty vector control are depicted in Additional file 2. We confirmed that the expression pattern of these four constructs was due to genomic integration by producing F1 transgenics. In each case, F1 GFP expression resembled that of F0 embryos (Additional file 2). To address the dependence of each sequence on the E1b minimal promoter, we swapped it with another TATA-box containing 31-bp element commonly used in luciferase reporter assays (see Materials and methods). Three of the four constructs had essentially identical patterns, suggesting that the observed tissue-specific activity is guided by the 15-bp sequence (Additional file 2). 2GH12/notochord was the outlier, suggesting that a sequence within the E1b minimal promoter could be needed for its tissue specificity.

**Multiplexed oligomers contain shorter, functional sequences**

A previously reported machine-learning algorithm assigned weights to 6-mers based on their association with putative forebrain enhancers [6], as defined by the binding of the E1A binding protein p300 (EP300) in mouse forebrain at embryonic day 11.5. We used these forebrain enhancer weights to score each of the 184 multiplexed constructs based on the combined weight of all 6-mers it contains. Higher scores indicate constructs with sequence patterns that are positively associated with EP300 binding in embryonic mouse forebrain. Using a logistic regression model, we compared the scores of constructs that had 24 hpf forebrain expression in the top 10% of our study with the scores of the remainder of the constructs. This analysis identified a significant association between our top forebrain constructs and 6-mers identified by Lee et al. (odds ratio $= 1.17, P = 0.008$; Wald test of logistic regression coefficient), despite differences in methods, organism, and
developmental stage between the two studies. We found an even stronger association using the top 5% of the forebrain constructs (odds ratio = 1.23, $P = 0.008$; Wald test).

For the 18 multiplexed oligomers with forebrain expression in the top 10% of our study, an average of only 8 of 14 distinct 6-mers in each construct have positive weights. This suggests that multiple 6-mers in each construct are dispensable for forebrain expression. To functionally explore this hypothesis in multiple tissues, we performed a functional dissection experiment on the four selected tissue-specific constructs. For each, we removed 6 bp to leave the left (L9), middle (M9), and right (R9) 9 bp of each 15-bp insert (Figure 3a). In all but one case (2GH12/notochord), a single 9-bp sequence was sufficient to largely reproduce the expression pattern of the original multiplexed construct (Figure 3b; Additional file 4).
We used Tomtom [30] to identify JASPAR [31] and UniPROBE [32] position weight matrices (PWMs) that were similar to the functional 9-bp (or 15-bp, in the case of 2GH12) sequences identified through the functional dissection experiments. Although TF binding specificities are thought to be conserved, the PWMs in these databases are not derived from zebrafish TF binding specificities, and thus are not ideal. However, in each case, the top hit (Figure 3b) suggested TFs or TF families that plausibly interact with these sequences. The M9 region of 2CD12/epidermis contains a GATA motif on the reverse strand, matching the UniPROBE PWM for Gata5 (P = 0.001, Tomtom empirical P-value [30]) but likely corresponding to Gata3 in zebrafish, which binds to a very similar motif in vitro (Additional file 2) and is a regulator of epidermal cell lineage [33]. Likewise, the L9 region of 3CD05/spinal cord matches the UniPROBE PWM for Hoxa10 (P = 0.0005), although this motif likely is recognized by a more medially expressed hox in zebrafish, such as Hoxa9a. Mammalian Hoxa10 and Hoxa9 binding motifs are extremely similar, and hoxa9a expression precisely mirrors the activity of 3CD05/spinal cord (Additional file 2). The L9 region of 2GH08/brain strongly matches the JASPAR PWM for Esrb (P = 0.001), the zebrafish homolog of which is expressed in forebrain and hindbrain nuclei, recapitulating the expression pattern driven by the 15-bp multiplexed oligomer (Additional file 2). 2GH12/notochord strongly matches the UniPROBE PWM for Foxa2 (P = 0.0009), which is expressed in the zebrafish notochord at the Prim-5 stage (approximately 24 hpf) [34]. The longer length of this motif provides a clue as to why 2GH12 could not be functionally dissected. In summary, the functional dissection experiments suggest the expression patterns of the four tissue-specific multiplexed oligomers are largely due to the action of shorter, 6- to 9-bp sequences.

Design of synthetic regulatory elements from smaller functional units
Regulatory sequences often contain clusters of homotypic TFBSs, with an average of five per cluster [35]. To determine whether this principle could aid in the design of synthetic regulatory elements, we generated concatemers of each of the four tissue-specific sequences, with five copies of each 15-bp insert, and tested them in zebrafish. 5x2CD12/epidermis, 5x2GH12/notochord and 5x3CD05/spinal cord produced more intense and specific expression patterns than their single-copy counterparts (Figure 4a; Additional file 2). In contrast, concatenation of 2GH08/brain abolished specific expression in the forebrain and otic lateral line entirely (Figure 4a). Combined with the functional dissection result, these data suggested that the XhoI-flanking sequence is necessary for proper function of the 2GH08 sequence. In a second round of concatemer design, we concatenated repeats of the functional 9-bp sequence from the functional dissection experiments (L9) along with the 6-bp XhoI flanking region. The construct XhoI-2GH08(L9) produced consistent expression (67% of embryos expressing) in rostral forebrain and otic lateral line in a more intense and specific pattern than 2GH08/brain (Figure 4b; Additional file 2). These results demonstrate that the strength of a regulatory element can be tuned by simple repeats of short, key regulatory sequences in vivo.

Large, distal-acting enhancers (average length 1,529 bp) have been shown to autonomously drive diverse expression [36]. However, the internal circuitry of individual enhancers is much more complex, requiring the precise spacing and ordering of regulatory sequences [37-40]. To assess the syntactic rules that apply at the scale of our synthetic regulatory elements, we assayed tandem combinations of two or three concatemers. A tandem combination of brain (5xXhoI-2GH08(L9)) and skin (5x2CD12) concatemers abolished expression in skin and forebrain but not the otic lateral line. A tandem combination of brain and notochord (5x2GH12) concatemers, as well as a tandem brain, skin and notochord triplet abolished regulatory element activity in those tissues (Figure 5a,c). In contrast, injecting pairs of concatemers as separate constructs yielded consistent, combinatorial expression of the two patterns (Figure 5b, c). In each tandem combination, the XhoI-2GH08(L9)/brain concatemer was the most distant from the minimal promoter, while the intervening sequence and the distance to the minimal promoter varied. The fact that two intervening sequences abolished the activity of the brain concatemer (brain+notochord and brain+skin +notochord), but another (brain+skin) only limited its spatial extent, suggests the local context of a regulatory element is essential to its function. These data are consistent with the idea that large, autonomous enhancers contain insulator elements and/or signals mediating long-range interactions [37] that are absent in our synthetic regulatory elements.

Short, tissue-specific oligomers are essential components of native regulatory elements
To explore the native function of the multiplexed sequences in the zebrafish genome, we identified construct alignments in H3K4me1-positive, H3K4me3-negative predicted-enhancer regions [12] and then employed GREAT [41] to identify significantly enriched gene ontology terms for each construct. For each of the four 15-bp constructs, we identified one or more significantly enriched biological processes that were relevant to the tissue expression pattern (binomial test, FDR q < 0.05; Figure 6a; Additional file 5). We further aligned 9-bp sequences identified in the functional dissection experiments and performed the same
analysis in GREAT, identifying several additional enriched terms that matched the corresponding expression pattern of the functional 9-bp sequence (Figure 6b; Additional file 6).

To determine if these sequences contribute to zebrafish regulatory modules, we picked ten 2GH12/notochord and ten 2GH08(L9)/brain alignments that overlap 24-hpf whole-embryo H3K4me1+/me3- ChIP-seq peaks.

Figure 3 Functional dissection of multiplexed sequences reveals key motifs (a) We tested three deletion mutants for each of the four tissue-specific multiplexed sequences, using the first (L9, peach), middle (M9, green) and last (R9, blue) 9 bp of each sequence cloned between the same flanking sequences. (b) With one exception (2GH12/notochord), a 9-bp sequence was sufficient to drive the expression pattern of the original multiplexed construct. Percentages of fish expressing in the appropriate tissue at 24 hpf are shown, with the value for the functional 9-mer highlighted in black. The top-matching position weight-matrix identified by Tomtom [30] is listed for each construct.
Figure 4 Design of synthetic enhancers from short multiplexed sequences. (a) We generated concatemers containing five copies of each of the six multiplexed sequences. Three of these (5x2CD12/epidermis, 5x2GH12/notochord, 5x3CD05/spinal cord) drove more intense expression in the same tissue as the original 15-bp sequence. The fourth construct, 5x2GH08/brain, did not produce any detectable expression, suggesting that the XhoI-flanking sequence may be important. White arrows mark the epidermis, notochord, and spinal cord, respectively. (b) A second version of the 2GH08/brain concatemer contained five tandem repeats of the XhoI flanking region with the 2GH08(L9) functional sequence that was identified by the 9-bp dissection experiments. This arrangement produced a strong regulatory element that drives consistent expression to the forebrain and otic lateral line at 24 and 48 hpf. White arrows mark the forebrain and otic lateral line patterns. All pictures are of 24 hpf embryos.
Combination of synthetic regulatory elements underscores the importance of context (a) Left: a tandem combination of the 5xXhoI-2GH08(L9)/brain and 5x2GH12/notochord concatemers failed to produce any enhancer activity above background. Right: a tandem combination of the 5xXhoI-2GH08(L9)/brain and 5x2CD12/epidermis concatemers resulted in consistent otic lateral line expression. Expression in the forebrain and epidermis was detected inconsistently in a handful of embryos (such as the one shown), but not as intensely as the original concatemer. (b) Injection of a mixture of the 5xXhoI-2GH08(L9)/brain and 5x2GH12/notochord (left) or 5xXhoI-2GH08(L9)/brain and 5x2CD12/epidermis constructs (right) resulted in the expected combinatorial pattern (white arrows). (c) Percentage of embryos expressing in the epidermis, forebrain and notochord for the 5x concatemers individually, as combinations or as a mixture of two constructs. Tandem arrangement of concatemers in the same regulatory element largely failed to reproduce the expression patterns driven by concatemers individually or in a mixture, with the exception of the brain-epidermis combination, which preserved otic lateral line expression.
We cloned approximately 1,000-bp fragments, including the alignment and overlapping H3K4me1 signal, and tested them for enhancer activity in zebrafish. Three 2GH12- and five 2GH08(L9)-containing fragments drove consistent (>40% of fish expressing) expression to notochord and forebrain, respectively (Figure 6c-j; Additional file 7). A random selection of ten H3K4me1+/me3- regions identified two strong forebrain enhancers and two weak enhancers, driving expression consistently yet nonspecifically to the notochord and forebrain, respectively (Additional files 2 and 7). Together, these results indicate that these multiplexed sequences could be used in conjunction with other genome-wide datasets to improve the identification of functional, tissue-specific enhancers.

To determine the relative strength with which our synthetic elements can drive expression, we performed quantitative PCR on pools of 25 GFP-positive embryos injected with the original 15-bp insert, the 5x concatemer, or a strong native enhancer (Nc2/Fb2) corresponding to 2GH08/brain or 2GH12/notochord (Additional file 2). In each case, the native enhancer produced higher GFP expression than both the 15-bp insert (3.0- and 10.4-fold for brain and notochord, respectively) and the 5x concatemer (2.5- to 5.0-fold, respectively), suggesting that synthetic 15-mers and 5x concatamers are not as strong as native enhancers.

We next performed deletion mutagenesis on each positive 1,000-bp enhancer, removing the 2GH12/notochord- and 2GH08(L9)/brain-aligned regions to determine if they are required to drive the specific expression patterns observed. The deletions reduced the specific expression of the enhancer in seven out of eight cases and abolished it for one of three positive 2GH12- and two of five positive 2GH08(L9)-containing enhancers (Figure 6k,l; Additional file 7). These results demonstrate that the multiplexed regulatory sequences identified by our screen are necessary for the proper spatiotemporal expression of specific enhancers in the developing zebrafish.

Discussion

In this study, we made several observations on the role of TFBS-sized sequences in gene regulation in vivo. First, we identified a considerable number of very short DNA sequences that are drivers of consistent, tissue-specific expression. A previous report carried out in various cell lines [20] suggested, as a rationale for screening concatameric sequences, that such short sequences could not drive observable levels of transcription. Our results suggest...
that this is not the case, and that there exists an intermediate stage between neutral sequence and fully-fledged regulatory elements. We also showed that expression levels can be amplified in vivo through homotypic concatenation, a principle that has previously been demonstrated primarily in vitro transient transfections [38,42,43]. Together, these findings suggest a potential mechanism through which short sequences could gain a toehold as minimal regulatory elements and evolve into more complex clusters.

A limitation of this approach is that each multiplexed construct contains multiple k-mers, requiring further bashing to determine functional sequences. However, such functional dissection is negligible compared to what would be required for a typical enhancer of more than 1,000 bp. Furthermore, multiplexing 6-mers into 15-bp oligomers identified sequences such as 2GH12, which was the only construct to drive notochord-specific expression and could not be broken down into smaller, functional units. This result, in particular, points to the potential benefits of testing higher order k-mers, ideally without multiplexing.

We additionally demonstrate that short k-mers (9 to 15 bp) are important for the proper spatiotemporal expression of a subset of distal-acting regulatory elements. Thus, even though our multiplexed sequences were generated synthetically, they harbor endogenous signals that confer tissue-specificity. Using a straightforward TFBS analysis tool and publicly available data, we identified potential interacting proteins for all four tissue-specific signals. However, these matches were not perfect, highlighting the need for more in vivo testing of short sequences.

Another contribution of this work is a mathematical and computational framework for the design of ultra-compact DNA libraries with single-copy representation of pairs of reverse-complementary k-mers. We show here that the reverse complementarity of DNA can be exploited to construct an MRCC library, which is about half the size of the analogous de Bruijn sequence library and yet still maintains uniform, minimal coverage of k-mers. Traditional de Bruijn sequences have proven to be extremely useful for biological applications, including protein-binding microarrays [3], universal DNA tag systems [21] and microfluidic affinity analysis devices [22]. Our formal analysis of the structure of the de Bruijn graph induced by the reverse complementarity of DNA should aid in the advancement of these and other specialized techniques that use k-mer libraries [20,44]. In addition, the use of programmable microarrays to generate barcoded libraries of sequences for functional testing would provide an enticing platform for applying MRCC libraries. These ‘massively parallel reporter assays’ (MPRAs) have so far only been used to test variants of an individual promoter or enhancer [45–47], but could be used to screen k-mer libraries on a larger scale.

Extending such methodologies to higher k may aid in the discovery of drivers for tissues such as the eye, otic vesicle, and fins, which were poorly represented in the 6-mer screen. For these tissues, it is entirely possible that 6-mers are simply not sufficient to drive expression. Although our screen covered 100% of 6-mers, it only captures 32% of all 7-mers, 9% of all 8-mers, and a mere 2% of the 262,144 unique 9-mers. Such methods could be combined with transgenic lines with secondary markers for specific tissues of interest to improve annotations.

Finally, as evidenced by a homotypic concatemer whose expression is affected by downstream sequences, we show that local context affects enhancer function. Our findings are consistent with previous reports that show that the function of regulatory elements is sensitive to changes of spacing and order [37–40], but could also imply that our synthetic enhancers are missing key, yet undiscovered, signals typical of endogenous enhancers. While deciphering the rules of syntax by testing individual combinations is technically challenging, this process could also be accelerated by testing rearrangements of a known enhancer in parallel, using MPRAs. One such example is the recent use of MPRA technology to characterize the regulatory logic of promoters in yeast [43]. Such large-scale functional dissections will aid in the design of synthetic regulatory elements that can drive molecules to a specified level of expression in any combination of tissues and time points.

Conclusions
Our unique catalog of short, functional sequences is a good starting point in the search for the basic functional units contributing to regulatory element activity. These sequences could be used to improve genome annotation and to interpret the effects of non-coding genetic variants. Another application is in the design of synthetic regulatory elements that can drive expression of molecules to specific tissues and time points. Using only a small subset of the ‘words’ in our 6-mer regulatory catalog, we demonstrate a simple type of ‘sentence’ that obeys the syntactical rule of amplification through repetition. Attempts to construct more elaborate sentences revealed a non-modular grammar at the scale of TFBS-sized elements, highlighting the need for functional assays with higher throughput.

Materials and methods
Ultra-compact 6-mer library design
Although we chose k = 6, our methods are applicable in principle to larger values of k and libraries of varying size. We restricted the design to include just one k-mer out of each reverse-complementary pair of k-mers (kRC-pair). For even values of k, a kRC-pair may consist of a
single self-reverse-complementary palindrome, rather than two sequences. For an RC-pair $X$ and collection $S$ of $m$ DNA sequences, $\chi(X,S)$ denotes the total number of times that the $k$-mers in $X$ appear as substrings in all sequences of $S$. For example, if $X = \{\text{TATTTTAATTTA}, \text{TAAATGGAATTTAAA}\}$ and $S = \{\text{TATTTTAATTTAGGG, TAAATGGAATTTAAA}\}$, then $\chi(X,S) = 4$. We introduced the concept of a MRCC library of order $k$ and size $m$, which, formally, is a collection $S$ of $m$ DNA sequences such that for every $k$RC-pair $X$, $\chi(X,S) = 1$.

The $k$-dimensional de Bruijn graph (Figure 1a; Additional file 2) has $n_k$ vertices, each labeled by a unique $k$-mer, and $n_{k+1}$ directed edges, each labeled by a unique $(k+1)$-mer. The edge labeled by $x$ is directed from the vertex labeled by the first $k$ bases of $x$ to the vertex labeled by the last $k$ bases of $x$. Every path, i.e., sequence of edges, corresponds to a DNA sequence: the label of the first edge gives the first $k+1$ bases of the sequence, and for $i > 2$, the final base in the label of the $i$-th edge gives the $(k+i)$-th base in the sequence. The reverse complement of a path $t$ is also a path: it consists of the reverse complements of the edges in $t$, traversed in reverse order. The $(k-1)$-dimensional de Bruijn graph is used to efficiently generate a de Bruijn sequence of order $k$ via the computation of an Eulerian cycle, that is, a single cyclical path that traverses every edge in the graph exactly once (Additional file 2). There is an equivalent path that traverses every node exactly once in the more visually intuitive $k$-dimensional de Bruijn graph, described in Figure 1a.

To create an MRCC library, we designed an algorithm (Additional file 1) that partitions the edges of the de Bruijn graph such that there are at most two palindromes per path (Additional file 2). Our algorithm has two main stages. Stage 1, which is only executed for even values of $k$ ($k > 2$), iterates over pairs of palindromes, removing from the graph two reverse-complementary paths between each pair. The paths correspond to reverse-complementary sequences, one of which is added to the library. Stage 2 partitions the remaining edges into two reverse-complementary cycles. The sequence that corresponds to one of those cycles is then divided into the remaining number needed of consecutive oligomers such that the final $k$-bases of each sequence overlap the first $k$-bases of the next (Additional file 2). The overlap is required to maintain $k$RC-pair coverage. These oligomers complete the library.

The algorithm is defined for any integers $k$ and $m$ such that $k > 0$ and $\frac{1}{2}p_k < m < \frac{1}{2}(n_k - \frac{1}{2}k)p_k$ (for $k = 2$, the range is $0 < m < \frac{1}{3}n_k$). For $k = 6$, $m$ must be between 32 and 1,952, which meets our constraints of our construct design, which required $m$ to be a feasible number for in vivo testing.

Stage 2 of our algorithm, which is an adaptation of the classic Eulerian-cycle algorithm by Hierholzer [48], is similar to a heuristic algorithm sketched by Mintseris and Eisen [24] that attempts to partition all edges of the $(k-1)$-dimensional de Bruijn graph into just two reverse-complementary cycles. For even values of $k$ larger than 2, it is not possible to partition the edges this way; every MRCC library has size $m > 1$ (Additional files 1 and 2). For odd values of $k$ and $k = 2$, this edge partitioning does exist (Additional file 2); one of the two cycles is equivalent to an MRCC sequence.

If the initial bases of an oligomer insert match the final bases of the left flanking sequence, then those bases can be eliminated from the oligomer to reduce its length without sacrificing its $k$RC-pair coverage. The same is true for the final bases of the oligomer and the right flanking sequence. We created a modified version of the algorithm that uses heuristics during cycle-finding and sequence-splitting to produce oligomers with these characteristics. This technique further reduced library size and the number of $k$-mers repeated across the junctions between oligomers and flanking sequences.

Software implementation
We implemented the original algorithm and used it to produce an initial construct design for $k = 6$ that contains $m = 208$ oligomers, each of length 15 bp. The number $N_k$ of $k$RC-pairs is $N_k = p_k + (\frac{1}{2})(n_k - p_k) = (\frac{1}{2})(n_k + p_k)$, where $n_k = 4^k$ is the number of $k$-mers and $p_k = 4^{k/2}$ is the number of palindromes. For $k = 6$, $N_k = 2,080$, $n_k = 4,096$, and $p_k = 64$, which implies that the initial design is optimally compact, excluding flanking sequences. With our implementation of the flanking-sequence-based, modified version of the algorithm, we reduced the number of oligomers by more than 10%, from 208 to 184, still each 15 bp.

Due to Perl’s facility in handling sequences, all implementations are in Perl. They can be adapted for better time and space efficiency, if needed for larger values of $k$, using a compiled programming language, such as C, and specific data structures (Additional file 1). All code is available from [25].

Plasmids and constructs
For the primary screen and deletion series, each oligomer (Additional file 3) and its reverse complement were synthesized (Bioneer Inc., Daejeon, Republic of Korea) with 5’ and 3’ overhangs, annealed and then cloned between XhoI and BglII restriction sites of the zebrafish enhancer vector E1b-Tol2 [28]. For the endogenous enhancer screen, 1,000-bp regions were amplified from zebrafish genomic DNA using CACC flanked primers, and then cloned into pENTR using the TOPO Cloning kit (Life Technologies, Carlsbad, CA, USA). Cloned enhancers were subsequently transferred to the E1b-Tol2 vector using the LR Clonase II cloning enzyme (Life Technologies). For the concatemeric constructs, 150- to 225-bp
regions were synthesized and cloned into E1b-Tol2 (Biomatik USA, LLC, Wilmington, DE, USA).

Zebrafish injections
Constructs were injected with Tol2 by standard techniques [27,49]. An average of 60 embryos were injected for the primary screen of 184 oligomers, with the secondary requirement that at least 25 embryos survived to 48 hpf. The expression patterns of the four tissue-specific constructs chosen for follow-up experiments were monitored in at least 100 embryos at each time point. For the alternative minimal promoter screen, we replaced the E1b minimal promoter with the 31-bp minimal promoter region from the pGL4.23 luciferase reporter vector (Promega, Madison, WI, USA).

GFP expression was recorded in 13 tissues at 24 hpf and 15 tissues at 48 hpf according to established ZFIN nomenclature [50]. At least two images (one lateral, one dorsal or ventral) were acquired for each construct, with more images taken for constructs showing consistent expression patterns. A complete collection of screen data and a collection of image data for consistently expressing constructs are available from our website [51].

Correlation analysis
For each multiplexed construct, fractional expression values were determined by dividing the number of embryos expressing in a given tissue by the total number surviving at each time point. The same values were obtained for an “empty vector” control construct lacking an oligomer insert. Correlation between these two profiles was then determined for each construct and time point using the Spearman’s rank correlation coefficient. Constructs with an FDR-adjusted P-value <0.05 were considered correlated to the empty vector.

Support-vector machine forebrain enhancer weights
Using the 6-mer weights assigned by a previously reported support vector machine (SVM) that predicts EP300-binding in forebrain [6], we computed a score for each construct by summing the weights of the 6-mers in the construct (15-bp oligomer insert with 2-bp flanking sequence on each end). We binarized our expression data, assigning 1 to a construct if its forebrain expression value was in the top 10% (or 5%) of all constructs at 24 hpf, and 0 otherwise. Those data were used to fit a logistic regression model using R [52], with construct score as the predictor variable. We tested for statistical significance of the association between the score and the probability of forebrain expression in the top 10% (5%) using a Wald test.

Transcription factor binding site analysis
Nine or 15-bp (in the case of 2GH12) functional sequences were compared to known vertebrate JASPAR [31] and UniPROBE [32] PWMs using Sandelin-Wasserman similarity [53] in Tomtom [30]. The most statistically significant PWM match to each construct was identified.

Alignments to the zebrafish genome
Fifteen-base-pair inserts were aligned to the zebrafish genome (build Zv9) using bowtie v0.12.730 [54], allowing three mismatches and masking repeats (UCSC Genome Browser [55] RepeatMasker track). For the shorter, functional sequences identified by the functional dissection experiments, we aligned 11-mers (9 bp + 2 bp of flanking sequence), with one mismatch allowed.

Enrichment analyses
GC content was computed from the Zv9 assembly, with repeats masked. Enrichment analyses were performed using Fisher’s exact test. P-values were Bonferroni adjusted.

We predicted enhancer regions using previously reported histone methylation data [12] available from the UCSC Genome Browser [55]. We used the UCSC table browser to intersect 15- or 11-bp alignments with predicted enhancers, and fed the results into GREAT [41] using the default settings, with an FDR-adjusted binomial P-value threshold of 0.05.

Quantitative PCR
Twenty-five GFP expressing embryos were pooled for each injected construct (as well as an uninjected control pool) and dounce homogenized in TRIZol reagent (Life Technologies). Total RNA was extracted according to the manufacturer’s instructions, and subsequently cleaned up using the RNeasy mini kit (Qiagen, Hilden, Germany). To reduce genomic DNA, we performed an on-column digestion with the RNase-free DNase set (Qiagen). Total RNA (1.5 µg) was processed into cDNA using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA, USA). For each reaction, we performed a control without the reverse transcriptase enzyme (no RT) to quantify residual genomic GFP. We performed Taqman assays for GFP or beta-actin 1 (actb1) using the Taqman Universal PCR Master Mix, No AmpErase UNG reagent (Applied Biosystems), in triplicate. GFP expression was first normalized to the no RT conditions to single out GFP RNA, then to actb1 as a loading control.

Additional material

Additional file 1: Supplemental note. Two results that justify and give a mathematical proof of correctness of our formal algorithm (in box), which we used to construct the oligomer library.

Additional file 2: Supplemental figures. Seven supplemental figures and legends.
Additional file 3: Supplemental Table 1. Multiplexed sequence screen data for 184 constructs tested using zebrafish transgenic assays.

Additional file 4: Supplemental Table 2. Embryo counts from the functional dissection experiments.

Additional file 5: Supplemental Table 3. Enriched gene ontology terms for 15-bp alignments of the four tissue-specific multiplexed oligomers.

Additional file 6: Supplemental Table 4. Enriched gene ontology terms for 9-bp (+2 flanking bp) alignments of the three tissue-specific multiplexed oligomers.

Additional file 7: Supplemental Table 5. Genomic position and embryo counts for the thirty approximately 1,000-bp regions tested for enhancer activity.

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