Highly active zinc-finger nucleases by extended modular assembly

Mital S. Bhakta,1,2 Isabelle M. Henry,3 David G. Ousterout,4 Kumitaa Theva Das,1 Sarah H. Lockwood,1 Joshua F. Meckler,1 Mark C. Wallen,1 Artem Zykovich,1 Yawei Yu,5 Heather Leo,5 Lifeng Xu,5 Charles A. Gersbach,4 and David J. Segal1,6

1Genome Center and Department of Biochemistry and Molecular Medicine, University of California, Davis, California 95616, USA; 2Department of Pharmaceutical Sciences, University of Arizona, Tucson, Arizona 85721, USA; 3Department of Plant Biology and Genome Center, University of California, Davis, California 95616, USA; 4Department of Biomedical Engineering and Institute for Genome Science and Policy, Duke University, Durham, North Carolina 27708, USA; 5Department of Microbiology, University of California, Davis, California 95616, USA

Zinc-finger nucleases (ZFNs) are important tools for genome engineering. Despite intense interest by many academic groups, the lack of robust noncommercial methods has hindered their widespread use. The modular assembly (MA) of ZFNs from publicly available one-finger archives provides a rapid method to create proteins that can recognize a very broad spectrum of DNA sequences. However, three- and four-finger arrays often fail to produce active nucleases. Efforts to improve the specificity of the one-finger archives have not increased the success rate above 25%, suggesting that the MA method might be inherently inefficient due to its insensitivity to context-dependent effects. Here we present the first systematic study on the effect of array length on ZFN activity. ZFNs composed of six-finger MA arrays produced mutations at 15 of 21 (71%) targeted loci in human and mouse cells. A novel drop-out linker scheme was used to rapidly assess three- to six-finger combinations, demonstrating that shorter arrays could improve activity in some cases. Analysis of 268 array variants revealed that half of MA ZFNs of any array composition that exceed an ab initio B-score cutoff of 15 were active. These results suggest that, when used appropriately, MA ZFNs are able to target more DNA sequences with higher success rates than other current methods.

[Supplemental material is available for this article.]
To streamline the methodology even further, we demonstrated that the specificity of MA arrays was similar to that of naturally occurring zinc-finger proteins and therefore was unlikely to be the primary cause of their poor performance in ZFNs (Lam et al. 2011). In principle, modules could be re-engineered to have higher affinity. As a more pragmatic solution, longer arrays of modules could be used to increase affinity. Anecdotal evidence supported this concept (Gordley et al. 2009; Guo et al. 2010; Perez-Pinera et al. 2012). However, we showed recently that a caveat to the latter approach is that the addition of modules can, in some cases, reduce the activity of a ZFN, perhaps by allowing subgroups of fingers to bind additional unexpected locations (Shimizu et al. 2011). These considerations led us to perform a systematic investigation of number and composition of modules on ZFN activity.

**Results**

**Longer MA arrays generally increase ZFN activity, but the relationship is not linear**

To systematically examine the relationship between array length and ZFN activity, we designed extended-MAXFNs to eight loci in the human genome near SNPs associated with coronary artery disease. Target sites were chosen that contained spacers of five, six, or seven base pairs between the two zinc-finger binding sites, requiring ZFNs to be constructed with zinc-finger–FokI linkers of TGGS, TGAAAR, and TGGPGAAAAR, respectively (Handel et al. 2009). Unlike CoDA target sites, all sites here contained mixtures of ANN, CNN, GNN, and TNN type triplets (Supplemental Table S1; Supplemental Appendix SI). Arrays composed of three to six fingers were constructed using the Barbas set of zinc-finger modules, which are the most expansive in terms of sequence recognition (Bhakta and Segal 2010). Since the sequential addition of modules is laborious, we developed a “drop-out linker” strategy by which the three-, four-, and five-module arrays could be created from a six-module array using silent restriction sites in coding regions (Fig. 1; Supplemental Fig. S1). In this way, only two six-finger arrays would need to be synthesized or assembled for each heterodimeric ZFN. The two sets of four arrays could then be generated in parallel in one day to enable the rapid testing of all 16 combinations of arrays.

Using a single-strand annealing (SSA) recombination reporter assay in HEK293T cells, we examined the activity of the various combinations of arrays (Fig. 2; Table 1). Generally, ZFNs composed of two three-finger arrays were least active, whereas ZFNs composed of two six-finger arrays were more active (defined as a percentage of the activity of a control ZFN [GZF3-L3 + GZF1-R3]) (Szczepk et al. 2007). However, activity did not correlate linearly with the number of fingers or modules. The optimal lengths for the left and right arrays were often different. For example, increasing the left array of the CS2-1 ZFN showed recently that a caveat to the latter approach is that the addition of modules can, in some cases, reduce the activity of a ZFN, perhaps by allowing subgroups of fingers to bind additional unexpected locations (Shimizu et al. 2011). These considerations led us to perform a systematic investigation of number and composition of modules on ZFN activity.

**MA ZFNs are active at chromosomal target sites**

To streamline the methodology even further, we demonstrated that we could also omit the SSA assay. Cells were treated directly with the 16 combinations of ZFNs targeting two additional endogenous loci and analyzed using the Surveyor assay, which detects the appearance of mutations due to nonhomologous end joining at the cleavage site. Active MA ZFNs were obtained for both loci (Fig. 3; Supplemental Fig. S2). Consistent with the differential array behaviors observed in the SSA assay, all left arrays of the T2-X6 ZFN supported activity, but only R4 produced an active nuclease.

High-throughput sequencing demonstrated that seven of the 10 L6 + R6 ZFNs (70%) produced indels (Table 1, Exploratory study; Supplemental Fig. S3; Supplemental Appendix III) at 0.5% or greater (a value considered “active” in other ZFN studies) (Gupta et al. 2012). In addition, seven array-length variants that had been found to be more active by SSA or the Surveyor assay were generally found to be more active at their endogenous sites. Some ZFNs remained inactive, perhaps due to an unfavorable chromatin environment at the endogenous locus. Of the 13 ZFNs for which both SSA and indel data were available, the correlation of percent indel activity to percent SSA activity was modest ($R^2 = 0.25$). However, there was perfect correlation between an indel threshold of ≥0.5% and an SSA threshold of ≥8%.

**Development of a scoring system to help predict outcomes**

To apply this “extended-MA” approach as a useful method, we developed a scoring system to rank potential binding sites based on predicted zinc-finger binding affinity. Using the measured affinities of Barbas GNN modules (Segal et al. 1999), previous studies showed that the relative affinity of new 3 × GNN arrays (recognizing sites of the composition 5′-GNNGNNGNN-3′) could be accurately predicted by calculating the value of ΔΔG (Sander et al. 2009). Unfortunately, this method is limited to the prediction of 3 × GNN arrays since similar affinity data do not currently exist for ANN, CNN, or TNN modules. We therefore created an ab initio “B-score” based on the theory that binding energy should be proportional to the number of bivalent hydrogen bonds, such as between G and Arg or between A and Gln or Asn (Fig. 4A; Supplemental Table S2; Supplemental Discussion S1). The B-score correlated well ($R^2 = 0.73–0.86$) (Fig. 4B,C) with the measured affinities of 3 × GNN arrays (Segal et al. 1999; Sander et al. 2009) and

![Figure 1](https://example.com/figure1.png)  
**Figure 1.** The drop-out linker scheme. Sphl, BsmI, and HindIII sites were introduced into the zinc-finger coding region as silent mutations. Digestion with one restriction enzyme followed by ligation allows the full set of arrays to be created in parallel in one day.
moderately well ($R^2 = 0.52$) (Fig. 4D) with the affinities of six-finger arrays of mixed ANN, CNN, or TNN composition (MS Kim et al. 2011; Shimizu et al. 2011). The weak correlation with the latter data set was somewhat expected since it is known that the affinities of arrays longer than three fingers often do not scale linearly. However, the trend predicted by the B-score was more accurate for this set than any currently available method, such as summing the number of GNN modules ($R^2 = 0.12$). Therefore, though not a perfect predictor, the B-score was used as a predictor of relative binding affinity. The SSA scores for 92 ZFN array-length variants were evaluated by receiver operating characteristic (ROC) (Fig. 4E; Supplemental Appendix SI). The Comb.B score was found to be the best classifier for an SSA threshold of $8\%$ (area under the curve [AUC] = 0.77). Importantly, the Comb.B score was more accurate than the number of GNN modules (AUC = 0.66), which was recently suggested to have the best correlation with the activity of $3 \times$ GNN MA ZFNs (Zhu et al. 2011). A Comb.B score of 15 was found to be the best compromise between true and false positives.

**A prospective study based on B-scores**

The most efficient method for most investigators would be to (1) use the B-score to predict sites to which active ZFNs could be prepared; (2) screen just one ZFN combination on such sites for activity; and (3) optionally test all 16 combinations of arrays on active sites to determine the optimal nuclease configuration. ZFNs composed of

---

**Table 1. Summary of ZFN activity based on the SSA and genomic assays**

| ZFN    | CS2-1 | CS3-1 | CS5-1 | CS6-1 | CS6-2 | CS7-1 | CS7-2 | CS7-3 | T2-X1 | T2-X6 |
|--------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| Arrays | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 | L6 + R6 |
| % SSA  | 22    | 8     | 0.3   | 6     | 81    | 9     | 27    | n.d.  | n.d.  | n.d.  |
| % Indels | 3.4  | 1.5   | 0.0   | 0.0   | 0.7   | 1.3   | 0.7   | 0.2   | 1.3   |
| Arrays | L5 + R3 | L4 + R4 | L5 + R4 | L5 + R4 | L6 + R5 | L4 + R6 | L6 + R4 | L5 + R4 | n.d. |
| % SSA  | 72    | 0.6   | 3     | 31    | 36    | n.d.  | n.d.  |
| % Indels | 19.5 | 0.1   | 0.0   | 6.6   | 2.0   | 10.9  | 9.4   |
| % SSA  | 18    | 0.5   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   |
| % Indels | 12.9 | 0.5   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   | 0.0   |

(\% SSA) Cleavage activity determined as the percentage of activity compared to the G3-L3 + G1-R3 control nuclease in the single-strand annealing reporter assay in HEK293T cells; (% Indels) cleavage activity determined as the percentage of NHEJ-derived indel mutations found at the chromosomal target site in HEK293T cells (unless indicated); (n.d.) not determined; (l.q.) low quality results not definitive due to low number of quality sequencing reads.

*[^a]Indel frequencies may be overrepresented due to a comparatively low number of reads.*

*[^b]Assay performed in mouse Neuro2a cells.*

*[^c]Assay performed in human T-rex-Dest30-Neo cells.*

---

**Figure 2.** Activity of the CS series of ZFNs determined by a SSA assay. The number of fingers in the left and right arrays for each ZFN are indicated. Note that the CS3-1, 5-1, 6-1, and 6-2 series started with four- or six-finger arrays instead of three-finger arrays to reduce the assembly effort required prior to the invention of the drop-out linker strategy. Based on the data, the missing ZFNs seemed likely to be inactive and were not tested subsequently. (-) SSA reporter only as a negative control. (Ctrl) GZF3-L3 + GZF1-R3 control nuclease to which all activities are normalized. (Error bars) The standard deviation of normalized duplicates from at least two experiments. (*) $P < 0.01$ compared to the negative control based on a one-tailed homoscedastic $t$-test. (Black arrows) ZFNs used in genomic cleavage assays.
The ZFNs made in this study exclusively use canonical TGEKP interfinger linkers, which have been shown to be the optimal linker length for six-finger arrays (Paisach and Pabo 2003; Neuteboom et al. 2006). However, proteins with four or more fingers have the potential to bind to off-target sites using subsets of three fingers (Shimizu et al. 2011). Other studies have suggested that the use of non-canonical TGSQKP linkers between pairs of fingers can improve the specificity and consequently, ZFN activity, presumably by disrupting the binding of three-finger subsets (Moore et al. 2001; Gupta et al. 2012). This strategy has been adapted widely by Sangamo Biosciences and Sigma-Aldrich (Doyon et al. 2008; Perez et al. 2008; Hockemeyer et al. 2009). In contrast to these studies, we observed the use of disruptive linkers dramatically decreased the SSA activity of all configurations of extended-MA ZFNs and did not rescue the activity of any inactive ZFNs (Fig. 5). These data suggest that the use of disruptive linkers is not recommended for extended-MA ZFNs.

Longer MA arrays are not associated with increased cytotoxicity

To better understand the mechanism by which additional fingers affect performance (Supplemental Discussion S2), the affinity and specificity of the three- and five-finger arrays of ZFN CS2-1 were investigated by quantitative electromobility shift assays (Fig. 6A; Supplemental Fig. S4). Consistent with the activity data, CS2-1 L3 did not display detectable binding (KD > 532 nM), whereas L5 bound tightly (KD = 0.08 nM). In contrast, both R3 and R5 bound their target with similar affinity (KD = 0.56 nM). Therefore, the reduction in ZFN activity in comparing CS2-1-R3 to CS2-1-R5 is likely not due to loss of affinity, which is similar to observations we reported previously (Shimizu et al. 2011). The reduction in activity may be due to the ability of subsets of the multiple fingers to bind additional specific or non-specific targets. As one measure of specificity, the proteins were tested for binding to a nontarget DNA (proteins L3 and L5 on the R6 target and vice versa). All proteins failed to shift the nontarget DNA even at the highest protein concentrations used in this assay, suggesting a >1000-fold discrimination between target and non-target. Specificity was further examined using the Bind-n-Seq (Zykovich et al. 2009) target site selection assay (Fig. 6B). Compared to L3, L5 recognized a site that was not only longer but also a closer match with the intended target site. In contrast, the binding site length and composition of R5 improved minimally compared to R3. These results demonstrate that, for the case of CS2-1, ZFN failure was primarily the fault of just one array in the pair. Optimizing the array length improved both the affinity and specificity of the underperforming array.

As another assay of the specificity of short and long arrays, we examined the cytotoxicity of three inactive and three active ZFNs. ZFNs with poor specificity have more off-target cleavage events, leading to cytotoxicity. CS7-1-L3 + R3 was inactive with three-finger arrays, HIV7533-L6 + R6 was inactive with six-finger arrays (even though the L4 + R3 version was active), and CS2-1-L3 + R5 was inactive with arrays of mixed length (Fig. 6D). Cytotoxicity was assessed as a decrease in the percentage of ZFN-expressing cells on day 5 compared to day 1 (Fig. 6C). None of the inactive ZFNs were observed to be cytotoxic, arguing against the possibility that the long array-ZFNs only appear inactive because they kill the cells expressing them. HIV7533-L4 + R3 was the shortest active combination of arrays in our study; CS7-1-L6 + R6 was active with the longest arrays, and CS2-1-L5 + R3 was active with mixed array lengths. Importantly, the active ZFNs with the shortest and longest arrays were not observed to be cytotoxic, demonstrating that long arrays per se are not problematic. The third ZFN of mixed array length showed a significant level of cytotoxicity. Considering all of the data, it seems likely that such cytotoxicity is due to the specifics of that particular ZFN and not a general deficiency of the methodology. These observations generally support the use of extended MA to create active and nontoxic nucleases.

Discussion

Combining the data from the exploratory and prospective studies, L6 + R6 extended-MA arrays produced active ZFNs at 15 of 21 (71%) of endogenous target sites, greater than the success rates...
recently reported for other MA methods (25%) (S Kim et al. 2011; Zhu et al. 2011) or CoDA (50%) (Sander et al. 2011) and similar to the rate of an engineered two-finger archive (82%) (Gupta et al. 2012). From a practical perspective, this means it might only be necessary to test one or two L6 + R6 ZFNs to different sites to find one that is active. Activity can be improved by using shorter arrays in some cases, and the drop-out linker can facilitate this rapid empirical testing. Shorter arrays can also be useful to maximize the spectrum of sequences that can be targeted, since in some cases the lack of appropriate modules may prevent the construction of two six-finger arrays. Toward that end, the SSA scores of all 268 ZFN array variants examined in this study suggest that any combination of MA arrays would be successful in 38% of cases; but with a combined B-score of \( \Delta G \) they could produce an active ZFN in 52% of cases (Supplemental Table S4), which again would require testing of only one or two ZFNs. Thus, when choosing sites that are not L6 + R6 arrays, the combined B-score is particularly helpful. The accuracy (69%) of using the B-score was found to derive primarily from avoiding true negative unsuccessful short-array target sites (Fig. 7).

Another potential parameter for optimization is the use of long disruptive linkers between sets of two fingers. However, in contrast to previous studies, we observed only deleterious effects from the use of long linkers. It could be that even more empirical testing would be required to realize a benefit, such as systematically evaluating all possible linker substitutions and combinations. It should also be noted that most other studies have used long linkers in the context of two-finger modules rather than one-finger modules. Until better guidance becomes available, long linkers should be avoided for extended-MA ZFNs.
The very comprehensive lexicon of existing one-finger modules enables an extended-MA ZFN site approximately once every 52 bp (Supplemental Discussion S3). To illustrate the impact of the broader sequence recognition, a 60-kb region of DNA (hg18, chr9:22060000-22119999) was searched for potential ZFN sites using websites corresponding to three publicly available engineering methods.

- Our website (http://www.genomecenter.ucdavis.edu/segallab/segallabsoftware) using extended MA found 3474 ZFN sites with a B-score ≥15 (spacers = 5, 6, or 7). The website outputs in BED format, which can be easily uploaded as a custom track in the UCSC GenomeBrowser (http://genome.ucsc.edu/) (Fig. 8A). Of the 3474 ZFNs, 50% (1737) would be expected to be active.

- The ToolGen website (http://toolgen.co.kr/ZFNfinder/sws.cgi) using MA with a “recommended” set of modules (spacers = 5, 6, or 7) found 5401 3 + 3 ZFNs and 1791 4 + 4 ZFNs. 26% of potential cleavage sites could be targeted successfully by 4 + 4 ZFNs but only 9.1% by 3 + 3 ZFNs (Kim et al. 2009). Of the 3610 unique 3 + 3 and 1791 4 + 4 ZFNs, 15% (794) would be expected to be active (not shown).

- The Zinc Finger Consortium ZiFiT website (http://zifit.partners.org/ZiFiT/ChoiceMenu.aspx) using CoDA found 50 sites (spacers = 5, 6, or 7) (Fig. 8A). Of the 50 ZFNs, 50% (25) would be expected to be active (Sander et al. 2011).

These enhanced capabilities should also allow extended MA to target 91% of 28,527 unique protein-coding transcripts in the zebrafish genome (Ensemble Zv9 database), with an average of 118 sites per transcript, and 99% of 27,251 unique protein-coding transcripts in Arabidopsis (The Arabidopsis Information Resource 10 release) with an average of 26 sites per transcript (Fig. 8B). In comparison, CoDA was reported to target only 81% and 63% of these coding sequences, respectively, with an average of 4.4 and 2.5 sites per transcript (Sander et al. 2011).

The extended-MA method therefore provides dramatically improved genome engineering capabilities compared to other publicly available ZFN assembly methods. These results demonstrate that, when used appropriately, modular assembly can outperform methods that were developed to address its deficiencies (Cathomen and Joung 2008; Maeder et al. 2008; Sander et al. 2011). We believe that these results clarify a common misconception in the field that modular assembly is fundamentally inefficient. We note, however, that transcription activator-like effector nucleases (TALENs) appear to have an even greater success rate and target an even broader spectrum of DNA sequences than is currently possible with extended MA (Bogdanove and Voytas 2011; Reyon et al. 2012).

Robust assembly methods have already enabled widespread adaptation of TALENs for genome engineering applications in contrast to the limited ZFN methods that inspired this study.

Methods

Construction of zinc-finger arrays

In most cases, coding regions were synthesized by BioBasic, Inc. Arrays of three, four, five, and six fingers were created using the drop-out linker strategy described in Figure 1 and Supplemental Figure S1, sequential cloning of additional fingers by PCR, or using

---

**Figure 6.** An analysis of affinity, specificity, and cytotoxicity. (A) EMSA was used to determine the affinity of CS2-1 three- and five-finger arrays for their specific (cognate) target as well as a nontarget (e.g., the L3 array on the R3 binding site). A large ratio of nontarget:specific binding is an indicator of good specificity. (B) In vitro binding specificity was also determined using the Bind-n-Seq target site selection assay (Zykovich et al. 2009). The binding preference of CS2-1-L3 appears to differ from the intended target site at three positions (red boxes). (C) Cytotoxicity was assessed as a decrease in the percentage of ZFN-expressing cells on day 5 compared to day 1. To follow only those cells expressing nuclease, a GFP expression vector was cotransfected with the indicated ZFN expression vectors. (GFP) Cells cotransfected with GFP and empty ZFN vector as a positive control. (GZF3-wt) A nuclease known to be cytotoxic (Szczepk et al. 2007) as a negative control. (Error bars) The standard deviation of normalized duplicates from at least two experiments. (*) P < 0.00001 compared to the GFP-only positive control based on a one-tailed homoscedastic t-test. (D) Cleavage activity is a summary of the SSA data from other figures and is shown here for reference.
the SuperZiF system (Gonzalez et al. 2010). Additional fingers were digested with XhoI/AgeI and cloned between the XhoI/XmaI sites of these vectors so as to preserve canonical TGEKP linkers between all fingers. The sequences of all zinc-finger coding regions are provided in Supplemental Appendices SI and SII. Most heterodimeric ZFNs were expressed in vector pPGK-FokI and contained the obligate heterodimer modifications DD + RR (Szczepek et al. 2007). CS6-2, T2-X1, and T2-X6 were expressed in pCMV-FokI (DA + RV) (Szczepek et al. 2007). Dys5, Neo2, Neo3, DZF17, DZF24, DZF34, and DZF35 were expressed in pCMV-FokI (wt) and did not contain obligate heterodimer modifications. ZFNs used on target sites with 5-, 6-, or 7-bp spacers used zinc-finger–FokI linkers TGGS, TGAAAR, and TGPGAAAR, respectively (Handel et al. 2009).

Single-strand annealing (SSA) recombination reporter assay

The SSA assay is a plasmid-based reporter assay for detecting ZFN-induced repair of a split luciferase gene. A ZFN-induced double-strand break between homologous regions of the split luciferase will allow the SSA repair pathway to reconstruct an active luciferase gene. Construction of the SSA luciferase reporter plasmid, pSSA Rep 3-1, has been described previously (Szczepek et al. 2007; Bhakta and Segal 2010). The pSSA Rep3-1 and control GZF3-L3 + GZF1-R3 ZFN plasmids are available from Addgene (ID 5091–5094). Briefly, ZFN binding sites were introduced into the left and/or right arms of a split firefly luciferase gene by PCR and cloned into the BgIII/EcoRI sites of the vector. All primers used for SSA construction are listed in Supplemental Table S5. In poly-l-lysine treated, 24-well plates, HEK293T cells at 80% confluency in DMEM supplemented with 10% fetal calf serum and 1 unit/mL of penicillin and 1 μg/mL of streptomycin were cotransfected with 100 ng of each ZFN monomer expression plasmid, 25 ng of pRL-TK- Renilla Luciferase (as a transfection control), and 25 ng of SSA reporter plasmid using Lipofectamine 2000 (Invitrogen). Cells were harvested 48 h post-transfection by removing media, washing with 500 μL of 1× DPBS followed by lysis in 100 μL of 1× Passive Lysis Buffer (Promega) with 1× Complete protease inhibitors (454 Life Sciences [Roche]). Clarified cell lysis supernatants (20 μL) were used to determine the luciferase activity using DualGlo or BrightGlo reagents (Promega) in a Veritas microplate luminometer (Turner Biosystems). All experiments were performed in duplicates and repeated on at least two different days.

Electromobility shift assay (EMSA)

The DNA binding domains of CS2-1-L3, -L5, -R3, and -R5 were subcloned into the prokaryotic expression vector pMal-c2 (New England Biolabs) that had been modified to contain a C-terminal His6 tag. Proteins were expressed in BL21 (DE3) E. coli (Invitrogen) for 6 h at 37°C, purified over amyllose resin in Zinc Buffer A (ZBA: 100 mM Tris, pH 7.5, 90 mM KCl, 1 mM MgCl2, and 100 mM ZnCl2), and eluted in ZBA, 10 mM maltose, and 5 mM DTT. Complementary pairs of 5’-biotin labeled forward and 5’-poly-T reverse oligonucleotides were annealed to obtain double-stranded target DNAs. The sequences of all oligonucleotide targets are listed in Supplemental Table S6. Protein-DNA binding reactions were performed for 1 h at room temperature in ZBA, 150 mM KCl, 5 mM DTT, 0.1 mg/mL BSA, 0.05% NP-40, 35 pM target DNA, and purified ZFPs at the indicated concentrations. Gel electrophoresis was performed on 10% TBE-polyacrylamide gel in 0.5× TBE buffer at 4°C. The LightShift Chemiluminescent EMSA Kit (Pierce) was used according to the manufacturer’s protocol. After blotting on a Biodyne B nylon membrane (Pierce) for 1 h at 100 V at 4°C, the DNA was cross-linked by a UV cross-linker (Stratagene) for 4 min. Equilibrium binding constants (Kd) were calculated from protein titration experiments imaged on X-ray film. All experiments were performed in at least two biological repeats.

Figure 7. The utility of the B-score for extended-MA. Distribution of combined B-scores for successful (red bars) and unsuccessful (blue bars) ZFNs based on the 268 array combinations in this study. The optimal cutoff of 15 is indicated (green line).

Figure 8. Spectrum of sequences targetable using extended MA. (A) An image from the UCSC Genome Browser shows potential extended-MA and CoDA ZFNs sites found within a 60-kb region at the human 9p21 locus (hg18, chr9:22060000-22119999). Each vertical black bar represents a target site. (B) A comparison of the transcript targeting capabilities reported for CoDA and calculated from the data in this study for extended MA.
duplicates and repeated on two different days. The reported values represent the results of at least two experiments, with a standard deviation of ±50%.

Binding site specificity assay using massively parallel sequencing (Bind-n-Seq)

Bind-n-Seq was performed essentially as described (Zykovich et al. 2009). Briefly, the coding regions CS2-1-L3, -L5, -R3, and -R5 were subcloned into pDest-GST-MBP, expressed in BL21 (DE3) E. coli (Invitrogen), and purified over amyllose resin in ZBA, 10 mM maltose, 5 mM DTT. The maltose was removed by overnight dialysis in 2 L of ZBA, 5 mM DTT. Bar-coded 93-mer double-stranded oligonucleotide targets containing Illumina primer binding sites and a 21-nt random region were incubated with various concentrations of proteins and salt (50 mM protein, 1 mM KCl; 50 mM protein, 50 mM KCl; 50 mM protein, 100 mM KCl; 5 mM protein, 100 mM KCl; and 410 mM [CS2-1-L3, L5] or 200 mM [CS2-1-R3, R5] protein, 100 mM KCl), Bound complexes were precipitated using amyllose resin and enriched by six wash steps in the corresponding salt buffer. Eluted DNA was sequenced on an Illumina GAII sequencer. Motifs were determined using a custom motif finder that will be described elsewhere.

Surveyor assay

HEK293T cells (5 × 10^5) were seeded in six-well plates. The next day, cells were transfected at 90% confluence with 1 μg of each ZFN using Lipofectamine 2000 (Invitrogen). Cells were harvested 72 h post-transfection and genomic DNA was extracted using a Macherey-Nagel tissue prep kit or Qiagen Puregene Core kit A according to manufacturer’s instructions. The ZFN target site region was amplified from 100 ng of genomic DNA (2 min at 95°C; 15 sec at 95°C, 30 sec at 58°C, 1 min at 72°C, 35 cycles; 5 min at 72°C). The sequences of all primers are listed in Supplemental Table S7. PCR products (3 μL) were diluted in 1× Taq Pro Buffer (Denville), 50 mM KCl. The ampiclon mixture was heat denatured to allow for wild-type and mutant alleles to reanneal and form heteroduplexes (5 min at 95°C; 95°C to 85°C at -2°C/sec; 85°C to 25°C at -0.1°C/sec). The heteroduplex product was digested at the mismatch locus with Surveyor nuclease (Transgenomic) in 1× Enhancer Solution at 20 min at 42°C. Digestion was completed by adding 1 μL of stop solution and the product was resolved by running on 10% TBE-polyacrylamide gel. Gels were stained with ethidium bromide and visualized using a UV imager. Band intensities were quantified using ImageJ as described previously (Guschin et al. 2010).

Mutation analysis at endogenous loci using massively parallel sequencing

Genomic DNA from ZFN-treated cells was isolated as in the Surveyor assay. Genomic regions flanking the ZFN sites were amplified with primers containing Illumina sequencing ends and adapter sequence, followed by a 5-bp barcode and genomic priming site. The sequences of all primers are listed in Supplemental Table S8. Specific fragments containing the binding sites were amplified from genomic DNA (200 ng) 72 h post-transfection using Phusion polymerase (New England Biolabs) in a 100 μL reaction (2 min at 95°C; 15 sec at 95°C, 30 sec at 57°C or 60°C, 30 sec at 72°C, 31 cycles; 5 min at 72°C). The ampiclons were purified and separated on 1% agarose gel. The band of interest was extracted and eluted in Elution Buffer (Qiagen). Samples were checked for quality on a BioAnalyzer and sequenced using 100-bp or 150-bp paired-end reads on an Illumina HiSeq sequencer, 160-bp paired-end on Illumina MiSeq sequencer, or 85-bp paired-end reads on an Illumina GAII sequencer. Analysis was performed using a custom bioinformatics pipeline. In short, reads were sorted according to their respective barcodes and trimmed for sequence quality, presence of adapter sequence, and minimum length. Pairs of quality-filtered reads were surveyed for the presence of their specific genomic priming sites. Read pairs that did not contain perfect matches to both primer sequences were discarded. Pairs of sequences were compared to each other to search for overlap between the ends of the reads. When overlap was found, both sequences were combined into a single DNA sequence. Each of these resulting sequences was compared to the sequence and length of the wild-type (WT) target site sequence and the number of instances of each particular sequence was recorded.

Cytotoxicity assay

Toxicity was assessed by measuring the ratio of transfected cells on day 5 compared to day 1, as described by (Cornu and Cathomen 2010). HEK293T cells (5 × 10^5) were seeded in six-well plates in 2 mL of DMEM/BSA/PenStrep. Cells were transfected using Lipofectamine-2000 and 1 μg of pPNLK-GFP and 1 μg of each ZFN or PGK-empty (3 μg of total DNA). On day 1 (24 h post-transfection), cells were harvested for flow cytometry. Cells were washed with 1 mL of DPBS. Cells were released with 300 μL of trypsin then harvested in an additional 900 μL of DMEM complete medium. A portion of this (350 μL) was applied to a new six-well plate + 2 mL DMEM complete as the Day 5 sample. The remaining cells were washed with DPBS, resuspended in 300 μL of DPBS, and transferred to tubes for flow cytometry on ice. Cells (50,000) were scanned for GFP using a BD Biosciences LSRII. On day 5, cells were similarly harvested and scanned. All experiments were performed in at least duplicates and repeated on two different days.

Data access

Illumina massively parallel sequencing data are available at the NCBI BioProject archive (www.ncbi.nlm.nih.gov/bioproject) under accession number PRJNA179355. A website implementing the B-score and drop-out linker strategies is available at http://www.genomecenter.ucdavis.edu/segallab/segallabsoftware.

Acknowledgments

We thank Zhen Jia, Alice Adriaenssens, Sara Venturini, Lauren E. Cosby, and Matthew T. Brown for their technical contributions to this work. Funding was provided by the W.M. Keck Foundation and the UC Davis Clinical and Translational Sciences Center through NIH grant R0024146. I.M.H. was supported by NSF grant DBI 0733857. D.G.O. and C.A.G. were supported by an NIH Director’s New Innovator Award (DP2OD008586), the Hartwell Foundation, and a Basil O’Connor Starter Scholar Award from the March of Dimes. Author contributions: D.J.S. and C.A.G. conceived of the method; M.S.B., L.X., C.A.G., and D.J.S. designed research; M.S.B., D.G.O., K.T.D., S.H.L., J.E.M., M.C.W., Y.Y., and H.L. performed experiments; I.M.H., A.Z., and D.J.S. performed bioinformatics analyses; M.S.B., L.X., C.A.G., and D.J.S. wrote the manuscript.

References

Bae KH, Kwon YD, Shin HC, Hwang MS, Byu EH, Park KS, Yang HY, Lee DK, Lee Y, Park J, et al. 2003. Human zinc fingers as building blocks in the construction of artificial transcription factors. Nat Biotechnol 21: 275–280. Bhakta MS, Segal DJ. 2010. The generation of zinc finger proteins by modular assembly. Methods Mol Biol 649: 3–30.
Bogdanove AJ, Voytas DF. 2011. TAL effectors: Customizable proteins for DNA targeting. Science 333: 1843–1846.

Cathomen T, Joung JK. 2008. Zinc-finger nucleases: The next generation emerges. Mol Ther 16: 1200–1207.

Cornu TI, Cathomen T. 2010. Quantification of zinc finger nuclease-associated toxicity. Methods Mol Biol 649: 237–245.

Doyon Y, McCammon JM, Miller JC, Faraj E, Ngo C, Katibah GE, Amora R, Hocking TD, Zhang L, Rebar EJ, et al. 2008. Heritable targeted gene disruption in zebrafish using designed zinc-finger nucleases. Nat Biotechnol 26: 702–708.

Gonzalez-B, Schwimmer LJ, Fuller RP, Ye Y, Asawapommongkol L, Barbas CF III. 2010. Modular system for the construction of zinc-finger libraries and proteins. Nat Protoc 5: 791–810.

Gordley RM, Gersbach CA, Barbas CF III. 2009. Synthesis of programmable integrases. Proc Natl Acad Sci 106: 5053–5058.

Guo J, Gaj T, Barbas CF III. 2010. Directed evolution of an enhanced and highly efficient FokI cleavage domain for zinc finger nucleases. J Mol Biol 400: 96–107.

Gupta A, Christensen RG, Rayla AL, Lakshmanan A, Stormo GD, Wolfe SA. 2012. An optimized two-finger archive for ZFN-mediated gene targeting. Nat Methods 9: 588–590.

Hockemeyer D, Soldner F, Beard C, Gao Q, Mitalipova M, DeKelver RC, Handel EM, Alwin S, Cathomen T. 2009. Expanding or restricting the target site repertoire of zinc-finger nucleases: The inter-domain linker as a major determinant of target site selectivity. Mol Ther 17: 104–111.

Kim HJ, Lee HJ, Kim H, Cho SW, Kim JS. 2009. Targeted genome editing in human cells with zinc finger nucleases constructed via modular assembly. Genome Res 19: 1279–1288.

Kim S, Lee MJ, Kim H, Kang M, Kim JS. 2011. Preassembled zinc finger arrays for rapid construction of zinc finger nucleases. Nat Methods 8: 7.

Kim MS, Stybaeva G, Lee YJ, Revzin A, Segal DJ. 2011. A zinc finger protein array for the visual detection of specific DNA sequences for diagnostic applications. Nucleic Acids Res 39: e29.

Lam KN, van Bakel H, Cote AG, van der Ven A, Hughes TR. 2011. Sequence specificity is obtained from the majority of modular C2H2 zinc-finger arrays. Nucleic Acids Res 39: 4680–4690.

Mackay JP, Segal DJ, ed. 2010. Engineered zinc finger proteins: Methods and protocols. Springer Science and Business Media, New York.

Moore M, Klug A, Choo Y. 2001. Improved DNA binding specificity from polyzinc finger peptides by using strings of two-finger units. Proc Natl Acad Sci 98: 1437–1441.

Neuteboom LW, Lindhout BI, Saman IL, Hooykaas PJ, van der Zaal BJ. 2006. Effects of different zinc finger transcription factors on genomic targets. Biochim Biophys Acta Mol Cells 339: 263–270.

Pinski E, Pabo CO. 2003. Constraints for zinc finger linker design as inferred from X-ray crystal structure of tandem Zif268-DNA complexes. J Mol Biol 330: 1–7.

Perez EE, Wang J, Miller JC, Jouvenot Y, Kim KA, Liu O, Wang N, Lee G, Bartsch K, VV, Lee YL, et al. 2008. Establishment of HIV-1 resistance in CD4+ T cells by genome editing using zinc-finger nucleases. Nat Biotechnol 26: 808–816.

Ramirez CL, Foley JE, Wright DA, Muller-Lerch E, Rahman SH, Cornu TI, Winfrey RJ, Sander JD, Fu F, Townsend JA, et al. 2008. Unexpected failure rates for modular assembly of engineered zinc fingers. Nat Methods 5: 374–375.