A comprehensive approach to zinc-finger recombinase customization enables genomic targeting in human cells

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ABSTRACT

Zinc-finger recombinases (ZFRs) represent a potentially powerful class of tools for targeted genetic engineering. These chimeric enzymes are composed of an activated catalytic domain derived from the resolvase/invertase family of serine recombinases and a custom-designed zinc-finger DNA-binding domain. The use of ZFRs, however, has been restricted by sequence requirements imposed by the recombinase catalytic domain. Here, we combine substrate specificity analysis and directed evolution to develop a diverse collection of Gin recombinase catalytic domains capable of recognizing an estimated 3.77 × 10^7 unique DNA sequences. We show that ZFRs assembled from these engineered catalytic domains recombine user-defined DNA targets with high specificity, and that designed ZFRs integrate DNA into targeted endogenous loci in human cells. This study demonstrates the feasibility of generating customized ZFRs and the potential of ZFR technology for a diverse range of applications, including genome engineering, synthetic biology and gene therapy.

INTRODUCTION

Site-specific DNA recombination systems, such as Cre-loxP, FLP-FRT and φC31-att have emerged as powerful tools for genetic engineering (1,2). The enzymes that promote these conservative DNA rearrangements—known as site-specific recombinases—recognize short (30–40 bp) sequences and coordinate DNA cleavage, strand exchange and re-ligation by a mechanism that does not require DNA synthesis or a high-energy cofactor (3). This simplicity has allowed researchers to study gene function with extraordinary spatial and temporal sensitivity. However, the strict sequence requirements imposed by site-specific recombinases have limited their application to cells and organisms that contain artificially introduced recombination sites or pre-existing pseudo-recognition sites. To address this limitation, directed evolution has been used to alter the sequence specificity of several site-specific recombinases towards naturally occurring DNA sequences (4–8). Yet, despite advances (7,8), the widespread adoption of this technology has been hindered by the need for complex mutagenesis and selection strategies (4,7) coupled with the finding that re-engineered recombinase variants routinely demonstrate relaxed substrate specificity (4,6–8).

Zinc-finger recombinases (ZFRs) represent a versatile alternative to conventional site-specific recombination systems (9,10). These chimeric enzymes are composed of an activated catalytic domain derived from the resolvase/invertase family of serine recombinases and a zinc-finger DNA-binding domain, which can be custom-designed to recognize almost any DNA sequence (11–16) (Figure 1A). ZFRs catalyse recombination between specific ZFR target sites (17) that consist of two inverted zinc-finger–binding sites (ZFBS) flanking a central 20-bp core sequence recognized by the recombinase catalytic domain (18) (Figure 1B). In contrast to zinc-finger (19–21) and transcription activator-like (TAL) effector nucleases (22,23), ZFRs function autonomously and can excise and integrate transgenes in human and mouse cells without activating the cellular DNA damage response pathway (9,24–26). However, as with conventional site-specific recombinases, applications of ZFRs have been restricted by sequence requirements imposed by the recombinase catalytic

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domain, which dictate that ZFR target sites contain a 20-bp core derived from a native serine resolvase/invertase recombination site.

To address this problem, we previously described a knowledge-based approach for re-engineering serine recombinase catalytic specificity (27). This strategy, which was based on the saturation mutagenesis of specificity-determining DNA-binding residues, was used to generate recombinase variants that showed >10,000-fold shift in specificity. Significantly, this strategy focused exclusively on amino acid residues located outside the recombinase dimer interface (Supplementary Figure S1). As a result, we found that catalytic domains re-engineered by this method could associate to form ZFR heterodimers, and that designed ZFR pairs could recombine pre-determined DNA sequences with exceptional specificity. Taken together, these results led us to hypothesize that an expanded catalogue of specialized catalytic domains developed by this method could be used for the design of ZFRs with custom specificity. Here, we expand on our previous work by combining substrate specificity analysis and directed evolution to develop a diverse collection of Gin recombinase catalytic domains capable of recognizing an estimated $3.77 \times 10^7$ unique 20-bp core sequences. We show that ZFRs assembled from these re-engineered catalytic domains recombine user-defined sequences with high specificity, and that designed ZFRs integrate DNA into targeted endogenous loci in human cells. To our knowledge, this report describes the first generalized approach for the design of customizable site-specific recombinases and also provides the first demonstration of targeted integration into endogenous human loci by custom-designed site-specific recombinases.

**MATERIALS AND METHODS**

**Plasmids**

The split gene reassembly vector (pBLA) was derived from pBluescriptII SK (−) (Stratagene) and modified to contain a chloramphenicol resistance gene and an interrupted TEM-1 β-lactamase gene under the control of a lac promotor. ZFR target sites were introduced as previously described (8). Briefly, GFPuv (Clontech) was polymerase chain reaction (PCR) amplified with the primers...
GFP–ZFR–XbaI–Fwd and GFP–ZFR–HindIII–Rev and cloned into the SpeI and HindIII restriction sites of pBLA to generate pBLA–ZFR substrates. All primer sequences are provided in Supplementary Table S1.

To generate luciferase reporter plasmids, the Simian vacuolating virus 40 (S(V40) promoter was PCR amplified from pGL3-Prm (Promega) with the primers SV40–ZFR–BglIII–Fwd and SV40–ZFR–HindIII–Rev. PCR products were digested with BglII and HindIII and ligated into the same restriction sites of pGL3-Prm to generate pGL3–ZFR-1, 2, 3...18. The pBPS–ZFR donor plasmids were constructed as previously described (24,27) with the following exception: the ZFR-1, 2 and 3 recombination sites were provided in Supplementary Table S1.

Recombination assays
ZFRs were assembled by PCR as previously described (9,27). PCR products were digested with SacI and XbaI and ligated into the same restrictions sites of pBLA. Ligations were transformed by electroporation into Escherichia coli TOP10F' (Invitrogen). After 1-h recovery in Super Optimal Broth with Catabolite suppression (SOC) medium, cells were incubated with 5 ml of Super broth (SB) medium with 30 μg ml⁻¹ of chloramphenicol and cultured at 37°C. At 16 h, cells were harvested; plasmid DNA was isolated by Mini-prep (Invitrogen); and 200 ng of pBLA was used to transform E. coli TOP10F'. After 1-h recovery in SOC, cells were plated on solid Lysogeny broth (LB) media with 30 μg ml⁻¹ of chloramphenicol or 30 μg ml⁻¹ of chloramphenicol and 100 μg ml⁻¹ of carbenicillin, an ampicillin analogue. Recombination was determined as the number of colonies on LB media containing carbenicillin and chloramphenicol divided by the number of colonies on LB media containing only chloramphenicol. Colony number was determined by automated counting using the GelDoc XR Imaging System (Bio-Rad).

Selections
The ZFR library was constructed by overlap extension PCR as previously described (27). Mutations were introduced into the Gin catalytic domain at positions 120, 123, 136 and 137 with the degenerate codon NNK (N: A, T, C or G and K: G or T), which encodes 120, 123, 127, 136 and 137 with the degenerate codon introduced into the Gin catalytic domain at positions 18. The pBPS–ZFR donor plasmids were constructed as previously described (24,27) and ligated into the Agel and SpeI restriction sites of pBH–Gin–α, β, γ, δ, ε or ζ. Zinc-fingers were assembled by SuperZiF (28) and ligated into the Agel and SpeI restriction sites of pBH–Gin–α, β, γ, δ, ε or ζ to generate pBH–ZFR-L/R-1, 2, 3...18 (L: left ZFR; R: right ZFR) (Supplementary Table S2). ZFR genes were released from pBH by SfiI digestion and ligated into pcDNA 3.1 (Invitrogen) to generate pcDNA–ZFR-L/R-1, 2, 3...18. Correct construction of each ZFR was verified by sequence analysis (Supplementary Table S3).

Luciferase assays
Human embryonic kidney (HEK) 293 and 293T cells (ATCC) were maintained in Dulbecco’s modified Eagle’s medium containing 10% (vol/vol) Fetal Bovine Serum (FBS) and 1% (vol/vol) Antibiotic-Antimycotic (Anti-Anti; Gibco). HEK293T cells were seeded onto 96-well plates at a density of 4 × 10⁴ cells per well and established in a humidified 5% CO₂ atmosphere at 37°C. At 24 h after seeding, cells were transfected with 150 ng of pcDNA–ZFR–L 1–18, 150 ng of pcDNA–ZFR–R 1–18, 2.5 ng of pGL3–ZFR-1, 2, 3...18 and 1 ng of pRL–CMV using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions. At 48 h after transfection, cells were lysed with Passive Lysis Buffer (Promega), and luciferase expression was determined with the Dual-Luciferase Reporter Assay System (Promega) using a Veritas Microplate Luminometer (Turner Biosystems).

Integration assays
HEK293 cells were seeded onto 6-well plates at a density of 5 × 10⁵ cells per well and maintained in serum-containing media in a humidified 5% CO₂ atmosphere at 37°C. At 24 h after seeding, cells were transfected with 1 μg of pcDNA–ZFR–L-1, 2 or 3 and 1 μg of pcDNA–ZFR–R-1, 2 or 3 and 200 ng of pBPS–ZFR-1, 2 or 3 using Lipofectamine 2000 according to the manufacturer’s instructions. At 48 h after transfection, cells were split onto 6-well plates at a density of 5 × 10⁴ cells per well and maintained in serum-containing media with 2 μg ml⁻¹ of puromycin. Cells were harvested on reaching 100% confluence, and genomic DNA was isolated with the Quick Extract DNA Extraction Solution (Epicentre). ZFR targets were PCR amplified with the following primer combinations: ZFR–Target-1, 2 or 3–Fwd and ZFR–Target-1, 2 or 3–Rev (Unmodified target); ZFR–Target-1, 2 or 3–Fwd and CMV–Mid–Prim-1 (Forward integration); and CMV–Mid–Prim-1 and ZFR–Target-1,
2 or 3–Rev (Reverse integration) using the Expand High Fidelity Taq System (Roche). For clonal analysis, at 2 days post-transfection, 1 × 10^6 cells were split onto a 100-mm dish and maintained in serum-containing media with 2 µg ml⁻¹ of puromycin. Individual colonies were isolated with 10-×10-mm open-ended cloning cylinders with sterile silicone grease (Millipore) and expanded in culture. Cells were harvested on reaching 100% confluence, and genomic DNA was isolated and used as template for PCR, as described earlier in the text. For colony counting assays, at 2 days post-transfection, cells were split into 6-well plates at a density of 1 × 10^4 cells per well and maintained in serum-containing media with or without 2 µg ml⁻¹ of puromycin. At 16 days, cells were stained with a 0.2% crystal violet solution, and genome-wide integration rates were determined by counting the number of colonies formed in puromycin-containing media divided by the number of colonies formed in the absence of puromycin. Colony number was determined by automated counting using the GelDoc XR Imaging System (Bio-Rad).

RESULTS

Specificity profile of the Gin recombinase

To effectively re-engineer serine recombinase catalytic specificity, we first sought to develop a detailed understanding of the factors underlying substrate recognition by this family of enzymes. To accomplish this, we evaluated the ability of an activated mutant of the catalytic domain of the DNA invertase Gin (29) to recombine an extensive set of symmetrically substituted target sites. In nature, the Gin catalytic domain recombines a pseudo-symmetric 20-bp core that consists of two 10-bp half-site regions. Our collection of mutant recombination sites, therefore, contained each possible single-base substitution at positions 10, 9, 8, 7, 6, 5 and 4 and each possible two-base combination at positions 3 and 2 and the dinucleotide core. We determined recombination by split gene reassembly (8), a previously described method that links recombinase activity to antibiotic resistance.

In general, we found that Gin tolerates: (i) 12 of the 16 possible two-base combinations at the dinucleotide core (AA, AT, AC, AG, TA, TT, TC, TG, CA, CT, GA and GT); (ii) 4 of the 16 possible two-base combinations at positions 3 and 2 (CC, CG, GG and TG); (iii) a single A to T substitution within positions 6, 5, or 4; and (iv) all 16 possible single-base combinations at positions 10, 9, 8, and 7 (Figure 2A–D). Furthermore, we found that Gin recombined a target site library containing >10^6 (of a possible 4.29 × 10^9) unique base combinations at positions 10, 9, 8 and 7 within each 20-bp target (Figure 2D). These findings are consistent with observations made from crystal structures of the γδ resolvase (30,31), which indicate that (i) the interactions made by the recombinase dimer across the dinucleotide core are asymmetric and predominately non-specific; (ii) the interactions between an evolutionarily conserved Gly–Arg motif in the recombinase arm region and the DNA minor groove impose a requirement for adenine or thymine at positions 6, 5 and 4; and (iii) there are no sequence-specific interactions between the arm region and the minor groove at positions 10, 9, 8 or 7 (Figure 2E). These results are also consistent with studies that focused on determining the DNA-binding properties of the closely related Hin recombinase (32–34).

Re-engineering Gin recombinase catalytic specificity

Based on the finding that Gin tolerates conservative substitutions at positions 3 and 2 (i.e. CC, CG, GG and TG), we next investigated whether Gin catalytic specificity could be re-engineered to recognize core sequences containing each of the 12 base combinations not tolerated by the native enzyme (Figure 3A). To identify the specific amino acid residues involved in DNA recognition by Gin, we examined the crystal structures of two related serine recombinases, the γδ resolvase (30) and Gin resolvase (35), in complex with their respective DNA targets. Based on these models, we identified five residues that contact DNA at positions 3 and 2: Leu 123, Thr 126, Arg 130, Val 139 and Phe 140 (numbered according to the Gin catalytic domain (Ile 120, Thr 123, Leu 127, Ile 136 and Gly 137) by overlap extension PCR and constructed a library of ZFR mutants by fusing these catalytic domain variants to an unmodified copy of the ‘H1’ zinc-finger protein (ZFP) (9), which recognizes the sequence 5’-GGAGGCGTG-3’. The theoretical size of this library was 3.3 × 10^7 variants.

We cloned the ZFR library into substrate plasmids containing one of five base combinations not tolerated by the native enzyme (GC, GT, CA, AC or TT) and enriched for active ZFRs by split gene reassembly (8) (Figure 3C). After four rounds of selection, we found that the activity of each ZFR population increased >1000-fold on DNA targets containing GC, GT, CA and TT substitutions and >100-fold on a DNA target containing AC substitutions (Figure 3D). We sequenced individual recombinase variants from each population and found that a high level of amino acid diversity was present at positions 120, 123 and 127, and that >80% of selected clones contained Arg at position 136 and Trp or Phe at position 137 (Supplementary Figure S2). These results suggest that positions 120, 123 and 127 play critical roles in the specific recognition of unnatural core sequences, and that positions 136 and 137 are important structural determinants for DNA-binding. We evaluated the ability of each selected enzyme to recombine its target DNA and found that nearly all recombinases showed high activity (>10% recombination) and displayed a >1000-fold shift in specificity towards their intended core sequence (Supplementary Figure S3). As with the parental Gin, we found that several recombinases tolerated conservative substitutions at positions 3 and 2 (i.e. cross-reactivity against GT and CT or AC and AG), indicating that a single re-engineered catalytic domain could be used to target multiple core sites (Supplementary Figure S3).

To further investigate recombinase specificity, we determined the recombination profiles of five Gin variants (hereafter designated Gin β, γ, δ, ε and ζ)
shown to recognize 9 of the 12 possible two-base combinations at positions 3 and 2 not tolerated by the parental enzyme (GC, TC, GT, CT, GA, CA, AG, AC and TT) (Table 1). We found that Ginβ, γ and ζ recombined their intended core sequences with activity and specificity near that of the parental enzyme (hereafter referred to as Ginα), and that Ginγ, δ and ζ were able to recombine their intended core sequences with specificity exceeding that of Ginα (Figure 3E). Each recombinase displayed a >1000-fold preference for adenine or thymine at positions 6, 5 and 4 and showed no base preference at positions 10, 9, 8 and 7 (Supplementary Figure S4). These results indicate that mutagenesis of the DNA-binding arm allows for reprogramming of recombinase specificity at positions 3 and 2 without compromising recognition elsewhere. We were unable to select for Gin variants capable of tolerating AA, AT or TA substitutions at positions 3 and 2. One possibility for this result is that DNA targets containing >4 consecutive A–T base pairs might exhibit bent DNA conformations that interfere with recombinase binding and/or catalysis.

**Engineering ZFRs to recombine user-defined sequences**

We next investigated whether ZFRs composed of the re-engineered catalytic domains could recombine pre-determined sequences. To test this possibility, we searched the human genome (GRCh37 primary reference assembly) for potential ZFR target sites using a 44-bp consensus recombination site predicted to occur approximately once every 7.44 × 10^6 bp of random DNA (Figure 4A). This ZFR consensus target site, which was derived from the core sequence profiles of the selected Gin variants, includes ~3.77 × 10^7 (of a possible
1.0955 \times 10^{12}) unique 20-bp core combinations predicted to be tolerated by the 21 possible catalytic domain combinations and conservatively excludes low-affinity or unavailable 5’-CNN-3’ and 5’-TNN-3’ triplets within each ZFBS. Using ZFP specificity as the primary determinant for selection (36), we identified 18 possible ZFR target sites across eight human chromosomes (Chromosome 1, 2, 4, 6, 7, 11, 13 and X) at non-protein coding loci. On average, each 20-bp core showed \sim 46\% sequence identity to the core sequence recognized by the native Gin catalytic domain (Figure 4B). We constructed each corresponding ZFR by modular assembly (28) (‘Materials and Methods’ section).

To determine whether each ZFR pair could recombine its intended DNA target, we developed a transient reporter assay that correlates ZFR-mediated recombination to reduced luciferase expression (Figure 4A and Supplementary Figure S5). To accomplish this, we introduced ZFR target sites upstream and downstream an SV40 promoter that drives expression of a luciferase reporter gene. HEK293T cells were co-transfected with expression vectors for each ZFR pair and the corresponding reporter plasmid. Luciferase expression was measured 48 h after transfection. Of the 18 ZFR pairs analysed, 38\% (7 of 18) reduced luciferase expression by \> 75-fold and 22\% (4 of 18) decreased luciferase expression by \> 140-fold (Figure 4B). In comparison, GinC4, a positive ZFR control designed to target the core sequence recognized by the native Gin catalytic domain, reduced

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<th>Table 1. Catalytic domain substitutions and intended DNA targets</th>
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a Wild-type DNA target.
b The c catalytic domain also contains the substitutions E117L and L118S.
c The c catalytic domain also contains the substitutions M124S, R131I and P141R.

Figure 3. Re-engineering Gin recombinase catalytic specificity. (A) The canonical 20-bp core recognized by the Gin catalytic domain. Positions 3 and 2 are boxed. (B) (Top) Structure of the γδ resolvase in complex with DNA (PDB ID: 1GDT). Arm region residues selected for mutagenesis are shown as magenta sticks. (Bottom) Sequence alignment of the γδ resolvase and Gin recombinase catalytic domains. Conserved residues are shaded orange. Black arrows indicate arm region positions selected for mutagenesis. (C) Schematic representation of the split gene reassembly selection system. Expression of active ZFR variants leads to restoration of the β-lactamase reading frame and host-cell resistance to ampicillin. Solid lines indicate the locations and identity of the ZFR target sites. Positions 3 and 2 are underlined. (D) Selection of Gin mutants that recombine core sites containing GC, GT, CA, TT and AC base combinations at positions 3 and 2. Asterisks indicate selection steps in which incubation time was decreased from 16 h to 6 h (‘Materials and Methods’ section). (E) Recombination specificity of the selected catalytic domains (β, γ, δ, ε and ζ, wild-type Gin indicated by a) for each possible two-base combination at positions 3 and 2. Intended DNA targets are underlined. Recombination was determined by split gene reassembly and performed in triplicate.
luciferase expression by 107-fold. Overall, we found that 50% (9 of 18) of the evaluated ZFR pairs decreased luciferase expression by >20-fold. The remaining ZFR pairs, however, had a negligible affect on luciferase expression. Importantly, virtually every catalytic domain that displayed significant activity in bacterial cells (>20% recombination) was successfully used to recombine at least one naturally occurring sequence in mammalian cells.

To evaluate ZFR specificity, we separately co-transfected HEK293T cells with expression plasmids for the nine most active ZFRs with each non-cognate reporter plasmid. Every ZFR pair demonstrated high specificity for its intended DNA target, and 77% (7 of 9) of the evaluated ZFRs showed an overall recombination specificity nearly identical to that of the positive control, GinC4 (Figure 4C). To establish that reduced luciferase expression was the product of the intended ZFR heterodimer and not the byproduct of recombination-competent ZFR homodimers, we measured the contribution of each ZFR pair to recombination at least one naturally occurring sequence in mammalian cells.

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Engineered ZFRs target integration into the human genome

We next evaluated whether ZFRs could integrate DNA into endogenous loci in human cells. To accomplish this, we co-transfected HEK293 cells with ZFR expression vectors and a corresponding DNA donor plasmid that contained a specific ZFR target site and a puromycin-resistance gene under the control of an SV40 promoter (24) (Figure 5A). For this analysis, we used ZFR pairs 1, 2 and 3, which were designed to target non-protein coding loci on human chromosomes 4, X and 4, respectively (Figure 5A). At 2 days post-transfection, we incubated cells with puromycin-containing media and measured genome-wide integration rates by determining the number of puromycin-resistant (puro R) colonies. We found that (i) co-transfection of the donor plasmid and the corresponding ZFR pair led to a >12-fold increase in puroR colonies in comparison with transfection with donor plasmid only, and (ii) co-transfection with both ZFRs led to a 6- to 9-fold increase in puroR colonies in comparison with transfection with individual ZFR monomers (Figure 5B). The overall integration rates for ZFR pairs 1, 2 and 3 were determined to be 0.14 ± 0.06%, 0.24 ± 0.02% and 0.31 ± 0.1%, respectively. By comparison, the genome-wide integration rate of our internal ZFR positive control, GinC4, towards a pre-introduced target site (24,25) was previously determined to be 1%. To evaluate whether each ZFR pair correctly targeted integration, we isolated genomic DNA from puroR populations and amplified the targeted loci by PCR. The PCR products...

![Figure 4. ZFRs recombine user-defined sequences in mammalian cells.](http://nar.oxfordjournals.org/)

**Figure 4.** ZFRs recombine user-defined sequences in mammalian cells. (A) Schematic representation of the luciferase reporter system used to evaluate ZFR activity in mammalian cells. ZFR target sites flank an SV40 promoter that drives luciferase expression. Solid lines denote the 44-bp consensus target sequence used to identify potential ZFR target sites. The consensus ZFR target site consists of two-inverted 12-bp ZFBS flanking a central 20-bp core sequence recognized by the ZFR catalytic domain. Underlined bases indicate zinc-finger targets and positions 3 and 2. (B) Fold-reduction of luciferase expression in HEK293T cells co-transfected with designed ZFR pairs and their cognate reporter plasmid. Fold-reduction was normalized to transfection with empty vector and reporter plasmid. Renilla luciferase expression was used to normalize for transfection efficiency and cell number. The sequence identity and chromosomal location of each ZFR target site and the catalytic domain composition of each ZFR pair are shown. Underlined bases indicate positions 3 and 2. Standard errors were calculated from three independent experiments. ZFR amino acid sequences are provided in Supplementary Table S3. (C) Specificity of ZFR pairs. Fold-reduction of luciferase expression was measured for ZFR pairs 1 through 9 and GinC4 for each non-cognate reporter plasmid. Recombination was normalized to the fold-reduction of each ZFR pair with its cognate reporter plasmid. Assays were performed in triplicate.
corresponding to integration in the forward and reverse orientation were observed at the loci targeted by ZFR pairs 1 and 2 (Figure 5C). ZFR pair 3 was found to target integration only in the reverse orientation. The reason for this bias remains unclear, but it could be explained by preferential formation of a particular synaptic complex topology (37). To determine the overall specificity of ZFR-mediated integration, we isolated genomic DNA from clonal cell populations and evaluated plasmid insertion by PCR. This analysis revealed targeting specificities of 14.2% (5 of 35 clones), 8.3% (1 of 12 clones) and 9.1% (1 of 11 clones) for ZFR pairs 1, 2 and 3, respectively (Supplementary Figure S7). Sequence analysis of each PCR product confirmed ZFR-mediated integration (Figure 5D); however, we observed mutations within the donor plasmid nearby the anticipated junctions for each ZFR pair. The mechanism underlying how these mutations were introduced remains unknown. Taken together, these results indicate that ZFRs can be designed to integrate DNA into endogenous loci. Finally, we note that the ZFR-1 ‘left’ monomer was found to target integration into the ZFR-1 locus in the absence of the corresponding ‘right’ ZFR monomer (Figure 5C). This result is consistent with the luciferase reporter studies described earlier in the text (Supplementary Figure S6) and indicates that recombination-competent ZFR homodimers have the capacity to mediate off-target integration. The comprehensive evaluation of off-target integration events and the development of optimized obligate heterodimeric ZFR architectures should lead to the design of ZFRs that show greater targeting efficiency and specificity.

**DISCUSSION**

Targeted genome engineering is driving progress in new areas of research in gene therapy, synthetic biology and basic science. Although improvements in the design and assembly of zinc-finger and TAL effector nucleases have
been central to this revolution, the development of new methods that do not rely on DNA double-strand breaks and thus, do not carry the risk of non-homologous end joining-mediated mutagenesis, are necessary to improve the safety of genome engineering. ZFRs capable of autonomously catalysing recombination between DNA targets represent one such alternative. Yet, despite their promise, the use of ZFRs has been limited by the strict sequence requirements imposed by the ZFR catalytic domain. In the present study, we have addressed this problem by combining substrate specificity analysis and directed evolution to establish a user-friendly toolbox of modified serine recombinase catalytic domains suitable for the design of ZFRs with custom specificity. Guided by an extensive evaluation of serine recombinase catalytic specificity, we have developed a collection of re-engineered Gin recombinase catalytic domains that recognize an estimated $3.77 \times 10^7$ unique 20-bp core sequences. We have shown that ZFRs assembled from these re-engineered catalytic domains recombine user-defined sequences with high specificity and that designed ZFRs integrate DNA into pre-determined endogenous loci in human cells. Although previous studies have shown that site-specific recombinases, such as the φC31 integrase, can mediate integration into the human (38) and mouse genomes (39), these efforts were based on the presence of pseudo-recognition sites tolerated by the native enzyme (40), did not require catalytic reprogramming, and thus did not allow for targeting of user-defined sequences. To our knowledge, this report describes the first general approach for the design of site-specific recombinases with customizable specificity and also provides the first demonstration of targeted integration into endogenous human loci by customized site-specific recombinases.

Based on our current archive of >45 pre-selected zinc-finger modules, we estimate that ZFRs can now be designed to recognize between 5000 and 20 000 unique 44-bp DNA sequences in the human genome (Supplementary Note). This corresponds to approximately one potential ZFR target site for every 160 000–620 000 bp of random sequence and represents a substantial improvement in targeting capacity compared with conventional site-specific recombinases, which typically require complex evolution-ary methods for reprogramming (4,7). Currently, the requirement for adenine by the Gin recombinase within positions 6, 5 and 4 represents the only major sequence restriction with the strategy described. To alleviate this constraint, structurally and functionally related serine recombinase variants (18) with broad or complementary sequence requirements at these positions could be subjected to the types of directed evolution described in this study. This approach may effectively expand the targeting repertoire of this custom-designed site-specific recombinase family. Additional improvements in the targeting capacity of this technology could be envisioned with the incorporation of alternate DNA-binding domains; in particular, we anticipate that the re-engineered catalytic domains described herein should be compatible with recently described TAL effector recombinases (41). Application of more sophisticated and high-throughput methods for specificity profiling (42) should lead to more effective use of the evolved catalytic domains and may also improve ZFR activity. Finally, although the efficiency of ZFR-mediated integration is lower than that achieved by zinc-finger (43,44) or TAL effector (22) nuclease-based approaches, we anticipate that optimization of the ZFR architecture will lead to reduced off-target integration events and higher targeting efficiency. Additional studies aimed at evaluating whether ZFR activity is cell type (25) or chromatin structure dependent (45) may also help establish limitations and clarify opportunities for ZFR targeting. In conclusion, we have developed a diverse collection of re-engineered Gin recombinase catalytic domains suitable for the design of ZFRs with custom specificity. We have shown that ZFRs can be assembled to recombine user-defined DNA targets, and that designed ZFRs integrate DNA into endogenous genomic loci. This work illustrates the potential of ZFRs for a wide range of applications, including genome engineering, synthetic biology and gene therapy.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1–3, Supplementary Figures 1–7 and Supplementary Note.

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REFERENCES

Supporting Information:

A comprehensive approach to zinc-finger recombinase customization enables genomic targeting in human cells

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Supplementary Figure S1. Recombinase DNA-binding residues are located outside the dimer interface. The γδ resolvase in complex with target DNA. Catalytic domain dimer is colored cyan. DNA is colored grey. Arm region residues are shown as red sticks. Residues at the dimer interface are shown as purple sticks (PDB ID: 1GDT). Because the residues essential for specific DNA-binding are located outside the dimer interface, recombinase heterodimers with unique, asymmetric catalytic specificity can be generated.
Supplementary Figure S2. Sequence analysis of selected recombinases. Pie charts showing the percentage of amino acid substitutions at each targeted arm position. After the 4th round of selection, >20 clones were sequenced from each library. Sequence analysis of clones that recombine TT are described elsewhere (1).
Supplementary Figure S3. Core specificity of isolated catalytic domains. After 4 rounds of selection, the ability of selected catalytic domains to recombine core sequences with substitutions at positions 3 and 2 was evaluated. Assigned DNA targets are underlined. Recombinase mutations are shown. Asterisks indicate catalytic domains selected for further analysis. Wild-type base combination at positions 3 and 2 is CC. Recombination was determined by split gene reassembly (2) and performed in triplicate. Catalytic domains that recombine TT substitutions are described elsewhere (1).
Supplementary Figure S4. Position specificity of selected catalytic domains. Recombination assays between the $\alpha$, $\beta$, $\gamma$, $\delta$ and $\zeta$ catalytic domains and symmetrically substituted target sites. Recombination was measured on a library DNA targets that contained (a) >4,000 random strong base (S: G or C) substitutions at positions 6, 5 and 4 and (b) >10$^6$ (of a possible 4.29 x 10$^9$) unique base combinations at positions 10, 9, 8 and 7 (N: A, T, C or G). Recombination was measured by split gene reassembly (2) ($n = 3$).
Supplementary Figure S5. Reduced luciferase expression is dependent on the presence of ZFRs with complementary zinc-finger DNA-binding and recombinase catalytic domains. HEK293T cells were co-transfected with sub-optimal amounts of pGL3-PH4.20G reporter plasmid (25 ng) and increasing amounts of pcDNA-GinPH4 (fully complementary), pcDNA-Tn3PH4 (not fully complementary; mismatched catalytic domain) or empty pcDNA (negative). Renilla luciferase expression was used to normalize for transfection efficiency and cell numbers.
Supplementary Figure S6. ZFR homodimer activity. HEK293T cells were co-transfected with 150 ng ZFR-L or 150 ng ZFR-R with 2.5 ng of corresponding pGL3 ZFR reporter plasmid. Recombination was normalized to co-transfection with 150 ng ZFR-L and 150 ng ZFR-R with 2.5 ng pGL3 ZFR reporter plasmid.
**Supplementary Figure S7. Clonal analysis of ZFR-modified cells.** PCR primer combinations amplified either unmodified genomic target or integrated plasmid in the forward or reverse orientation.
Supplementary Table S1. Primers used in this study.

> GFP-ZFR-20G-XbaI-Fwd
TTAATTAAGAGTCTAGAGGAGGCGTGTTCAAAAACCATGTTTACAGCAGCCTCCAG
ATCTAGGAGGAATTTAAAAATGAG

> GFP-ZFR-20G-HindIII-Rev
ACTGACCTAGAGAGATTGGAGCGTGCTGTTACAAACCATGTTTTTGACAGCCCTCCC
TGCAATTTATTTGTACAGTTCATC

> SV40-ZFR-1-BglII-Fwd
TTAATTAAGAGAGATCGTGATCGAGTAGACAGAAACCCAGGTCCACCTTTCTTACTTGTGCTG
CTGGCGCATCTGCATCTCAATTAGTCAGC

> SV40-ZFR-1-HindIII-Rev
ACTGACCTAGAGAGATTGGAGCGTGCTGTTACAAACCATGTTTTTGACAGCCCTCCC
TGCAATTTATTTGTACAGTTCATC

> CMV-PstI-ZFR-1 Rev
CACCACCACCGCATCCGCAGCAAGTAGAAGAAAACCTTTGTCTGTATCTGCATCA
GCAATTTGATAAGCCAGCTCCAAA

> 5' Gin-HBS-Koz
CACCACCACCGCATCCGCAGCAAGTAGAAGAAAACCTTTGTCTGTATCTGCATCA
GCAATTTGATAAGCCAGCTCCAAA

> 3' Gin-Agel-Rev
CACCACCACCGCATCCGCAGCAAGTAGAAGAAAACCTTTGTCTGTATCTGCATCA
GCAATTTGATAAGCCAGCTCCAAA

> ZFR-Target-1-Fwd
GTTCTGTCCAGATCCACTAG

> ZFR-Target-1-Rev
GCATGTGTCCAGATGCATAGG

> ZFR-Target-2-Fwd
CACCTTCTCAGGATAAG

> ZFR-Target-2-Rev
GTTGCCCTGTATTCCCTGG

> ZFR-Target-3-Fwd
AATGAAGTTCCCTGGCACTTC

> ZFR-Target-3-Rev
CTGAAGGGTTTTAAGTGCAAGAAG
> CMV-Mid Prim-1
TGACGTCAGTACGGTTAATGG

ZFR targets are underlined.
**Supplementary Table S2.** The amino acid sequences of the catalytic domains described in this study.

> *Gin* α
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRALKRLQKGDTLVWKLDR
RLGRSMKHLISLVLGELRERGINFRSLTDSIDTSSPMGRFF
VMGALAEMELEGIERMA
AARNKGRG
> *Gin* β
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRALKRLQKGDTLVWKLDR
RLGRSMKHLISLVLGELRERGINFRSLTDSIDTSSPMGRFF
VMGALAEMELEGIERMA
AARNKGRG
> *Gin* γ
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRALKRLQKGDTLVWKLDR
RLGRSMKHLISLVLGELRERGINFRSLTDSIDTSSPMGRFF
VMGALAEMELEGIERMA
AARNKGRG
> *Gin* δ
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRALKRLQKGDTLVWKLDR
RLGRSMKHLISLVLGELRERGINFRSLTDSIDTSSPMGRFF
VMGALAEMELEGIERMA
AARNKGRG
> *Gin* ε
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRALKRLQKGDTLVWKLDR
RLGRSMKHLISLVLGELRERGINFRSLTDSIDTSSPMGRFF
VMGALAEMELEGIERMA
AARNKGRG
> *Gin* ζ
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRALKRLQKGDTLVWKLDR
RLGRSMKHLISLVLGELRERGINFRSLTDSIDTSSPMGRFF
VMGALAEMELEGIERMA
AARNKGRG

Targeted arm region positions are highlighted red. Random substitutions are highlighted blue. The hyperactivating H106Y mutation is highlighted green.
**Supplementary Table S3.** The amino acid sequences of the ZFRs described in this study.

>**ZFR-1 Left**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSTSGLNVRHQRHTGKEKYPKCEPCGKSFSQSGDLR
HQRHTGKEKYPKCEPCGKSFS

>**ZFR-1 Right**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSHRTLTNHQRHTGKEKYPKCEPCGKSFSQSGDLRR
HQRHTGKEKYPKCEPCGKSFS

>**ZFR-2 Left**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSQSGDLRRHQRHTGKEKYPKCEPCGKSFSQRAH
LERHQRHTGKEKYPKCEPCGKSFS

>**ZFR-2 Right**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSQSGDLRRHQRHTGKEKYPKCEPCGKSFSQRAH
LERHQRHTGKEKYPKCEPCGKSFS

>**ZFR-3 Left**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSQSGDLRRHQRHTGKEKYPKCEPCGKSFSQRAH
LERHQRHTGKEKYPKCEPCGKSFS

>**ZFR-3 Right**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSQSGDLRRHQRHTGKEKYPKCEPCGKSFSQRAH
LERHQRHTGKEKYPKCEPCGKSFS

>**ZFR-4 Left**
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDRPGLKRLKRLQGDLTVVWKL
RLGRSMKHLISLVGELRERGINFRSLTDSTDDSPMGRFFYVMGALAEREMELIFERMACLA
AARNKGRICRPPKSTGKEKYPKCEPCGKSFSQSGDLRRHQRHTGKEKYPKCEPCGKSFSQRAH
LERHQRHTGKEKYPKCEPCGKSFS
>ZFR-4 Right
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ

>ZFR-5 Left
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ

>ZFR-5 Right
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ

>ZFR-6 Left
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ

>ZFR-6 Right
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ

>ZFR-7 Left
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ

>ZFR-7 Right
MLIGYVRVSTNDQNTDLQRNALVCAGCEQIFEDKLSGTRTDPRGKLRLQKGDTLVVWKLD
RLGRSMKHLISLVLGELREGINFRSLTDSDTSPPMSGRFFFYVMGALAEMERELIPEQMAGTA
AARNKGRVRCPKPSGTGKEPKYPECGKSFSQRAHLERHQVRTHTGEPKYPECGKSFSQRAHLERHQRT
TGKKTSGQAGQ
Arm region mutations are highlighted red. Specificity-determining α-helical zinc-finger residues are highlighted green.
Supplementary Notes

The total number of 20-bp core sequences that can be targeted by each possible combination of Gin catalytic domain heterodimers was calculated as the product of the consensus sequence: 5’-NNNAAABNWWVTNNTNNN-3’, where N: A, T, C or G, B: T, C or G, V: A, C or G and W: A or T. This value was determined to be 3.77 x 10^7.

To determine the frequency of a 44-bp ZFR target site occurring randomly in the human genome, we first divided the total number of possible 44-bp ZFR targets sites based on our archive of 45 pre-selected zinc-finger modules (3-5) (6.33 x 10^{20}) with the total number of unique base combinations possible across this region (3.094 x 10^{26}). This relative frequency (1.606 x 10^{6}) was then multiplied by one haploid human genome (3.1 x 10^9 bp). The total number of ZFR target sites was thus determined to be 4,980.

We next estimated the overall frequency by dividing one haploid human genome (3.1 x 10^9 bp) with the total number of possible ZFR targets (4,980). By this calculation, one potential ZFR target site was predicted to occur approximately once every 620,000 bp of random sequence.

We note that by expanding the sequence constraints at the dinucleotide core to include all base combinations except those that are SS (S: C or G), the total number of ‘targetable’ 20-bp core sequences can be increased to ~1.1 x 10^8. Using this value, we estimate that one potential ZFR target site is predicted to occur in the human genome approximately once every 160,000 bp of random sequence.
Supplementary References


