

## Diversification of Transcription Factor Paralogs via Noncanonical Modularity in C2H2 Zinc Finger DNA Binding

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### SUMMARY

A major challenge in obtaining a full molecular description of evolutionary adaptation is to characterize how transcription factor (TF) DNA-binding specificity can change. To identify mechanisms of TF diversification, we performed detailed comparisons of yeast C2H2 ZF proteins with identical canonical recognition residues that are expected to bind the same DNA sequences. Unexpectedly, we found that ZF proteins can adapt to recognize new binding sites in a modular fashion whereby binding to common core sites remains unaffected. We identified two distinct mechanisms, conserved across multiple Ascomycota species, by which this molecular adaptation occurred. Our results suggest a route for TF evolution that alleviates negative pleiotropic effects by modularly gaining new binding sites. These findings expand our current understanding of ZF DNA binding and provide evidence for paralogous ZFs utilizing alternate modes of DNA binding to recognize unique sets of noncanonical binding sites.

### INTRODUCTION

Cross- and intra-species analyses of transcription factor (TF) binding site motifs and chromatin immunoprecipitation profiles have identified considerable binding site turnover from yeast to humans, supporting the prominent role for *cis*-changes in the evolution of regulatory networks (Borneman et al., 2007; Bradley et al., 2010; Gasch et al., 2004; Wilson et al., 2008). However, recent work has highlighted flexibility to changes in TFs themselves (i.e., changes in *trans*) (Baker et al., 2011; Lynch and Wagner, 2008; Nakagawa et al., 2013). A major challenge in obtaining a full molecular description of evolutionary adaptation is to characterize how TFs can change.

In this study, we focused on C2H2 zinc finger (ZF) proteins the largest structural class of TFs in eukaryotes—as a model protein family to investigate novel mechanisms of TF evolution. C2H2 ZF proteins (hereafter referred to as "ZF proteins") bind DNA using arrays of ZF domains, each containing an  $\alpha$  helix and two  $\beta$  strands (Klug, 2010) (Figure 1D). Extensive experimental (Beerli et al., 1998; Choo and Klug, 1997; Enuameh et al., 2013; Persikov et al., 2014; Wolfe et al., 2000) and computational (Benos et al., 2002; Kaplan et al., 2005; Mandel-Gutfreund and Margalit, 1998; Persikov and Singh, 2011; Siggers and Honig, 2007) analyses have established a ZF DNA recognition code in which amino acids at four canonical "recognition" positions (–1, 2, 3, and 6, as in Elrod-Erickson et al., 1996) in each ZF domain mediate DNA base contacts (Figure 1; Figure S1A available online). This stereotyped binding mode has made ZFs an object of intense research for the design of artificial TFs and custom ZF nucleases for site-specific genome editing (Klug, 2010).

Despite the appeal of a simple recognition code based on a few canonical residues, additional features of ZF proteins can affect DNA-binding specificity, including interdomain interactions (Isalan et al., 1997; Liu and Stormo, 2008; Wolfe et al., 1999), the interdomain linker sequence (Handel et al., 2009), ZF docking geometry (Siggers and Honig, 2007), and residues outside the canonical recognition residues (Persikov and Singh, 2011). Residues in the loop between the  $\beta$  strands (i.e.,  $\beta$  turn) can also affect DNA binding affinity and footprinting pattern (Shiraishi et al., 2005). Deviations from a simple recognition code provide an explanation for studies demonstrating that while binding specificity of ZF arrays can target particular sequences, off-target binding will occur (Lam et al., 2011; Ramirez et al., 2008). Binding plasticity is an impediment to DNA binding predictions and design; however, we speculated that it might provide an opportunity to identify mechanisms that perturb or broaden TF-DNAbinding specificity. We reasoned that analysis of binding variability between related ZFs might identify mechanisms for TF diversification and provide insights into TF evolution.

Analyzing the DNA binding of yeast C2H2 ZF proteins, we observed widespread differences among ZFs with identical canonical recognition residues. In addition to high-affinity binding to sites that conform to the canonical ZF recognition rules, we identified binding to noncanonical sites that has been conserved throughout fungal evolution. Unexpectedly, we found that ZF proteins can gain new DNA-binding specificities in a modular fashion whereby binding to common sites is unaffected. Furthermore, we demonstrate that this molecular adaptation occurs via at least two distinct mechanisms conserved across multiple



Ascomycota species. Our results support a model of TF evolution in which the binding to only a subset of DNA binding sites is altered—this allows for evolution of regulatory functions among paralogous TFs while alleviating negative pleiotropic effects.

#### RESULTS

### Yeast C2H2 Proteins Exhibit DNA-Binding Diversity beyond a Simple Recognition Code

To evaluate the DNA binding diversity in a group of related ZFs, we focused on the simplest system available—the ZF proteins from *Saccharomyces cerevisiae* that bind DNA via only two adjacent ZF domains (ZFs) (Figure 1D). We compared the DNA binding of ZF proteins with identical canonical recognition residues; according to the canonical recognition code, these ZF proteins should bind the same DNA sites. We subdivided 28 proteins with two adjacent ZFs into ten "specificity groups" such that proteins in each group have identical recognition residues (Table S1).

High-resolution universal protein binding microarray (uPBM) data were available for 24 proteins in eight specificity groups (Badis et al., 2008; Gordân et al., 2011; Zhu et al., 2009). The uPBM data provide unbiased and comprehensive binding profiles of each ZF protein to all 32,896 possible 8 bp sequences. We quantified the DNA binding similarity between proteins by correlating the binding profiles over the 500 top-scoring 8 bp sequences from each experiment (see Experimental Procedures). Clustering the pairwise comparisons showed clear divisions between proteins within the same specificity groups (Figures 1A and S1B–S1E). These observations demonstrate that, for this model system of two-ZF proteins, mechanisms exist that perturb the DNA-binding specificity from that predicted by a simple model based on canonical recognition residues.

In the S. *cerevisiae* lineage a whole-genome duplication (WGD) event occurred leaving many yeast genes with close paralogs (Wapinski et al., 2007). We found that the DNA-binding specificities for the majority of paralogs (6/8) are highly correlated (e.g., Msn2 and Msn4; Figures 1 and S1G). In contrast, with the exception of Mig proteins and Ygr067c/Yml081w, paralogs that arose prior to the WGD exhibit DNA binding differences. These results suggest that DNA binding differences that deviate from a simple recognition code are the norm, rather than the exception, even for these short C2H2 ZF proteins.

### Msn2-Family Proteins Bind Both Common and TF-Preferred DNA Sequences

To examine in more detail the nature of the binding differences between related ZFs, we focused on the Msn2 specificity group (Msn2/Msn4, Com2, and Rgm1/Usv1). Msn2 and Msn4 proteins are major stress response mediators and bind to the stress response element (STRE) AGGGG in stress response gene promoters (Martínez-Pastor et al., 1996). We compared the binding profiles of paralog representatives and identified the following: (1) "common" sites—high-affinity sites common to both TFs (green points, Figures 1B and 1C)—and (2) "TF-preferred" sites—sites bound preferentially by one TF (orange and magenta points, Figures 1B and 1C). Sequence motifs generated from these distinct sets of sites illustrate the nature of the binding differences (Figure 1D). Common sites recognized by all Msn2 specificity group members match the AGGGG-type STRE reported as an Msn2 and Msn4 target site. Binding to AGGGG is explained by a simple recognition model based on canonical residues and known residue-base preferences (Figures 1D and S1). In contrast, TF-preferred sites differ significantly from the AGGGG common site, with distinct differences at unique base positions throughout the motifs (Figure 1D). These results highlight that TF-preferred sites are recognized in addition to the common sites recognized by all members.

To evaluate the magnitude of the specificity differences, we determined equilibrium binding constants ( $K_d$ ) for select DNA sites by electrophoretic mobility shift assay (EMSA) (Figures 1E and S2). Binding experiments for Com2, Usv1, and Msn2 demonstrated high affinity (i.e., lower nanomolar) binding to the common and their "preferred" sites. In contrast, binding to the preferred sites of the other proteins was significantly lower (e.g., Com2 bound 10.7-fold more weakly to the Usv1-preferred site than its own). These results demonstrate that the PBM data correspond well with traditional equilibrium binding affinities, as has been shown in previous studies (Siggers et al., 2011). Furthermore, the data show that binding affinities to the common and TF-preferred sites are of comparable magnitude.

To determine whether the TF-preferred sites are functionally relevant, we tested for enrichment of the TF-preferred sites in genomic regions bound in vivo. TF-preferred sites specific to the Usv1/Rgm1 paralogs, and not bound by other Msn2-group proteins, are significantly enriched ( $p < 1 \times 10^{-5}$ , Fisher's exact test) in regions bound by Rgm1 during growth in complete medium (Wang et al., 2011), supporting that the TF-preferred sites are utilized in vivo (see Experimental Procedures).

# Msn2-Family DNA-Binding Differences Are Conserved throughout Fungal Evolution

We next examined whether the DNA binding preferences were conserved in orthologs from other species or whether they occurred only in *S. cerevisiae*. We performed uPBM experiments for 18 Msn2-family orthologs from five other Ascomycota fungi, including three species that diverged before the WGD (*Candida albicans, Kluyveromyces waltii,* and *Kluyveromyces lactis*) and two that diverged afterward (*Saccharomyces castellii* and *Candida glabrata*) (Table S2). Clustering the binding profiles revealed that binding by the orthologs from the five other fungal species fell into the same three specificity groups that we previously identified (Figure S1F). Thus, the TF-preferred binding identified for the Msn2 family orthologs has been conserved since the last common ancestor of *C. albicans* and *S. cerevisiae*, approximately 300 million years ago, and therefore likely represents a selected and functionally relevant deviation from the canonical binding mode.

## **Com2-Preferred Binding Is Mediated in a Modular Fashion and Requires an N-Terminal Basic Motif**

We sought to determine the mechanism by which the Com2 protein recognizes the Com2-preferred sites. We identified a conserved RGRK motif N-terminal to ZF1 in the Com2 orthologs that is not present in the other Msn2-family members (Figures 1F and S3). Strikingly, mutating the RGRK motif to RGEE (Com2 RK $\rightarrow$ EE; Figure S3; Table S2) selectively abrogated the Com2-preferred binding behavior (compare Figures 2A and 2B) but

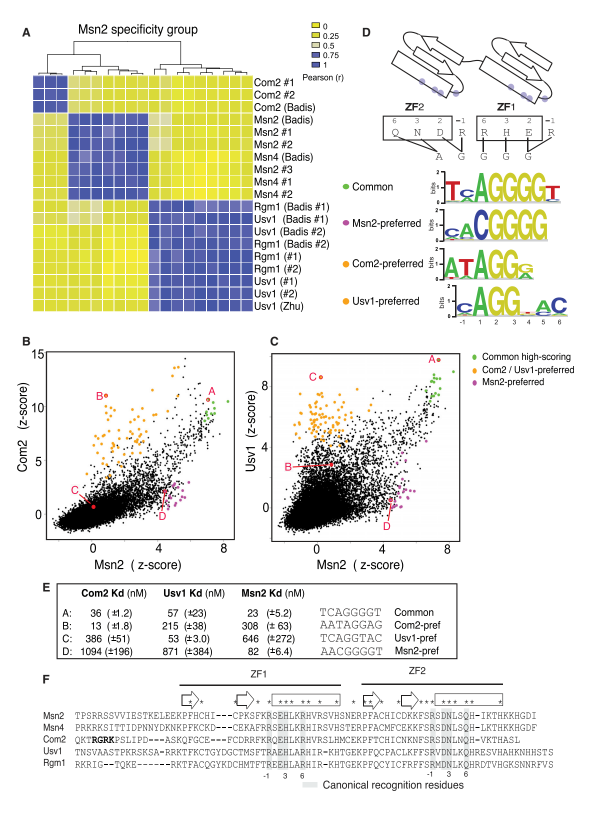


Figure 1. Common and TF-Preferred DNA-Binding Specificities of Msn2-Family Members

(A) Hierarchical clustering of pairwise binding profile comparisons for the Msn2-specificity-group proteins. Comparisons were performed for published data sets (Badis et al., 2008; Zhu et al., 2009) and duplicate PBMs from this study.

did not affect binding to the common sites. Therefore, an intact RGRK motif is required for the Com2-preferred binding. An N-terminal truncated version of Com2 (Com2  $\Delta$ N-term, Figure S3) missing the entire N terminus flanking ZF1 produced nearly identical results to the Com2 RK  $\rightarrow$  EE mutant (Figure S2B).

To determine if the Com2 RK $\rightarrow$ EE adversely affected the overall binding affinity, in addition to altering the binding specificity, we determined equilibrium binding constants (K<sub>d</sub>s) for the mutant protein to the select DNA sequences analyzed previously (Figures 2B and 2K). We found that the Com2 RK $\rightarrow$  EE mutant maintains its high affinity to the common site sequence (15 nM) but has an 18.5-fold reduction in affinity for the Com2-preferred site (13 nm to 240 nM; Figure 2K). These results demonstrate that binding to the Com2-preferred sites is a completely modular activity that can be specifically removed without affecting binding to the common site. Furthermore, the modular nature of binding suggests that Com2 uses two distinct binding modes to recognize different sequences.

## The Com2 N-Terminal Basic Region Enhances Binding to AT-Rich Sites

A general feature of RXR and RXXR peptide motifs is to select for AT-rich DNA sequences via DNA minor groove contacts (Rohs et al., 2009). Com2-preferred sites show a strong preference for an AT dinucleotide at positions -2 and -1 (Figure 1D). To test if Com2-preferred binding, mediated by the RGRK motif, operates by selective stabilization to sites with AT sequences 5' to the AGGGG core, we compared the binding of Com2 and the Com2 RK $\rightarrow$ EE mutant to three different "core" sequences (AGGGG, AGGAG, and AGGGT) with either a 5' TC dinucleotide (seen in the common sites) or a 5' AT dinucleotide (see in Com2preferred sites). As predicted, the Com2 N-terminal region enhanced the binding affinity to all sequences with the AT dinucleotide at positions -2 and -1 but did not affect the relative preference for the AGGGG, AGGAG, or AGGGT core sequences (Figure 2C). These results, in conjunction with the K<sub>d</sub> values (Figure 2K), demonstrate that the Com2 N-terminal region enhances binding to sites with an AT dinucleotide 5' to the common AGGGG core.

## Noncanonical Residues in Usv1 ZF1 Are Involved in Binding to Usv1-Preferred Sites

We next examined the mechanism of Usv1/Rgm1-preferred binding. In contrast to Com2, removal of the N- and C-terminal sequence outside the ZF domains (Table S2) had no effect on Usv1 binding (Figure 2D); therefore, Usv1-preferred binding operates by a different mechanism. We focused on residues within ZF1, which is predicted to interact with DNA bases at positions 4,

5, and 6 that varied in the Usv1-preferred sites (Figure 1D). In ZF1, we identified four residues (-5, -2, 5, and 8 canonical numbering) conserved in Usv1, Rgm1, and their orthologs but not in the other Msn2-group members (Table S2; Usv1 4-Res). Mutating these four residues to their Msn2 counterparts significantly and selectively weakened binding to Usv1-preferred sites, while not affecting binding to common sites (Figures 2E and 2F). These results demonstrate that residues in ZF1, distinct from the canonical recognition residues, are involved in the selective binding to Usv1-preferred sites.

## Usv1 Binding Preferences Can Be Engineered onto Msn2 in a Modular Fashion

To better understand the modularity and possible evolutionary path to the observed TF-preferred binding, we set out to engineer the Com2- and Usv1-preferred binding activity onto Msn2. Adding the Com2 N-terminal region onto Msn2 (Table S2; Msn2 ZFs/Com2 N-term) did not result in Com2-preferred binding for Msn2 (data not shown), suggesting that additional amino acid positions within the ZF domains are required to stabilize or permit the Com2-preferred binding mode. In contrast, an extensive set of mutations across the Msn2 ZFs-changing the Msn2 residues to their Usv1 counterparts (Table S2; Msn2 mut ZFs)-led to strong Usv1-preferred binding for Msn2 (compare Figures 1C and 2G). When mutations were restricted to ZF1 and the interdomain linker region, less pronounced but clearly present Usv1-preferred binding remained (Figure 2H). Therefore, while full Usv1-preferred binding requires residues from ZF2, partial Usv1-preferred binding can be obtained with residues from only ZF1 and the linker region. Further restrictions to the mutated residues in Msn2 largely abrogated the Usv1preferred binding (Figures 2I and 2J). These results demonstrate the following: (1) Usv1-preferred binding can be added in a modular fashion without affecting binding to the common sites, and (2) full Usv1-preferred binding requires residues distributed across both ZF domains and the inter-ZF linker. These results demonstrate a second mechanism to expand C2H2 TF binding in a modular fashion.

## Adr1 Specificity Group Proteins Exhibit Binding Similarities with Msn2 Group Proteins

Our initial survey of ZF DNA binding profiles identified binding variability within multiple ZF specificity groups (Figures S1B–S1E; Table S1). We examined the nature of the binding variability in the other large specificity group, hereafter referred to as the Adr1 group (Table S1, top row) (Ypr022c, Rsf2, Yml081w/Ygr067c, and Adr1). As seen for the Msn2 group, comparisons of Adr1 versus Ygr067c (Figure 3A) and Ypr022c versus

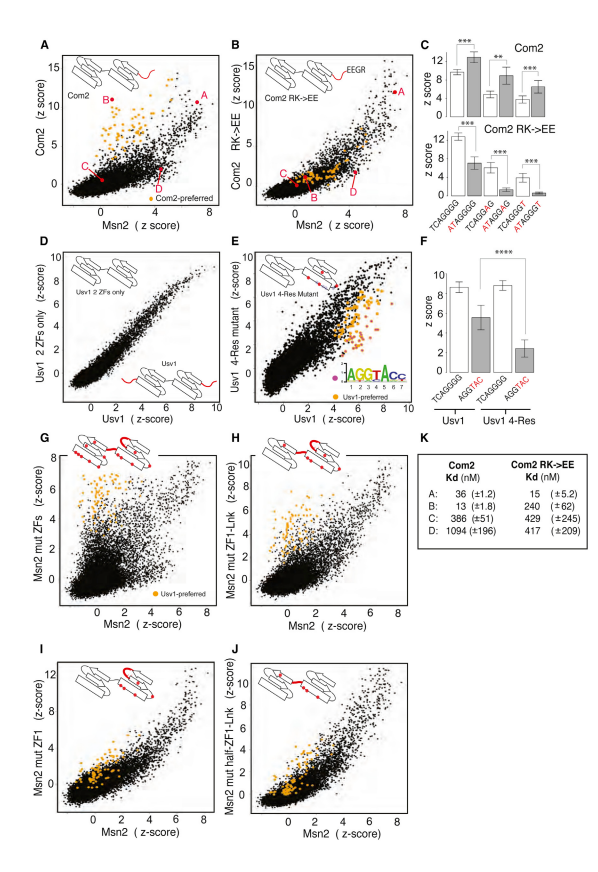
<sup>(</sup>B) Comparison of Msn2 and Com2 binding to all possible (32,896) 8 bp sequences. Z scores are transformed 8-mer median signal intensities (see Experimental Procedures). TF-preferred sites bound preferentially by Msn2 or Com2 are in magenta or orange, respectively. Common sites bound significantly (PBM E score > 0.48) by both proteins are highlighted. Four 8 bp sequences (labeled A, B, C, and D) assayed for binding in EMSAs are in red.

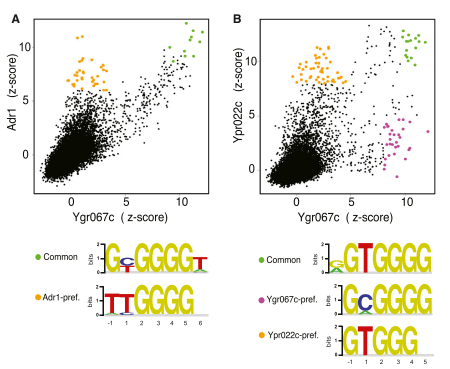
<sup>(</sup>C) Comparison for Msn2 and Usv1 (details as in [B], except that Usv1-preferred sites are in orange).

<sup>(</sup>D) Binding schematic for Msn2-family proteins and binding site motifs for TF-preferred and common sites. Canonical recognition residues are indicated (gray dots). Proposed interaction map for canonical residues and DNA bases of STRE is shown (see also Figure S1).

<sup>(</sup>E) Dissociation binding constants (mean and standard deviation) for Com2, Usv1, and Msn2 to selected DNA sequences are listed.

<sup>(</sup>F) Protein sequence alignment for Msn2 specificity group proteins. Highlighted are conserved residues (\*), canonical ZF DNA-contacting residues (gray bars, canonical numbering scheme), Com2 N-terminal RGRK motif (bold), and ZF secondary structure elements: β strands (empty arrows) and α helices (empty boxes).





Ygr067c (Figure 3B) revealed both common and TF-preferred sites. Furthermore, additional features of the TF-preferred bind-

ing in the Adr1 group resemble those seen for the Msn2 group,

suggesting similar mechanisms might be operating. For

example, both the Com2- and Adr1-preferred sites are distin-

guished by the preference for a Thy at position -1 (Figures 1

and 3). Provocatively, studies have shown that an N-terminal

proximal accessory region (PAR) in Adr1 is necessary for high-af-

finity binding to the 5'-TTGGAG UAS1 element that resembles

the Adr1-preferred sites (Schaufler and Klevit, 2003; Thukral

et al., 1991). This suggests that the N-terminal PAR domain in

Adr1 may mediate Adr1-preferred binding in a manner analo-

gous to the Com2 N-terminal region described here. Future

studies should clarify whether the TF-preferred binding in the

Adr1-group is similarly modular, and whether the mechanisms

used are identical to those of Com2 and Usv1, or if they repre-

(A and B) Pairwise binding profile comparisons for Adr1 and Ypr022c relative to Ygr067c. Z scores are as in Figures 1B and 1C. Common and TFpreferred sites are highlighted. Binding motifs are shown for highlighted TF-preferred and common sites (base numbering as in Figure 1D, with base congruence defined by ZF binding schematic, see Figure S1).

## DISCUSSION

In this work, we sought to uncover new mechanisms by which related TFs can diversify their DNA binding. Analyzing a model system of two-ZF proteins in yeast, we identified two distinct mechanisms by which ZF proteins can gain binding specificity in a modular fashion. We propose that the modular binding of these proteins comes from using multiple modes of DNA binding (Figure 4; discussed more below). Analysis of orthologs from other Ascomycota species demonstrated that this

modular diversification strategy has been conserved since the divergence of *C. albicans* and *S. cerevisiae* and is likely a functionally important adaptation. Modular evolution of TF binding specificity, in which protein changes mediate binding to new sites while not affecting the binding to a core set of common sites, provides an elegant solution to the problem of widespread negative pleiotropic effects—modularity allows a TF to gain novel regulatory functions while avoiding potential negative consequences from loss of regulation from the core sites.

The ability of individual TFs to bind DNA via multiple binding modes has been reported for a number TFs (reviewed in Siggers and Gordân, 2014) and likely represents a general mechanism for TF binding diversification. A recent study has described multiple binding modes for forkhead TFs and provided evidence that the distinct binding modes have arisen repeatedly and independently in the course of forkhead evolution (Nakagawa et al., 2013). Interestingly, the DNA sequences bound by distinct

#### Figure 2. Binding Specificities of ZF Mutants

sent additional ways to diversify ZF binding.

(F) Comparison of Usv1 and Usv1 4-Res mutant binding to two binding sequences: (i) TCAGGGG common site (Figure 1D) and (ii) AGGTAC – a Usv1-preferred site (Figure 1D) that was bound poorly by Usv1 4-Res mutant. Scores are mean Z scores for the eight different 8-mers containing TCAGGGG (columns 1 and 3) and for the 48 different 8-mers containing AGGTAC (columns 2 and 4) (error bars = 1 SD). (\*\*\*\*)  $p < 10^{-15}$ , unpaired t test.

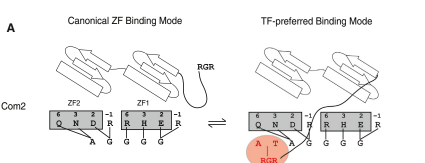
(G–J) Binding specificities of Msn2 wild-type and mutant proteins. Binding profile comparison for Msn2 mutants relative to Msn2. Usv1-preferred sites (as in Figure 1D) are highlighted (orange).

(K) Dissociation binding constants (mean and SD) for Com2 (as in Figure 1E, for comparison) and Com2 RK $\rightarrow$  EE mutant to select DNA sequences are listed. Probe sequences are as in Figure 1E.

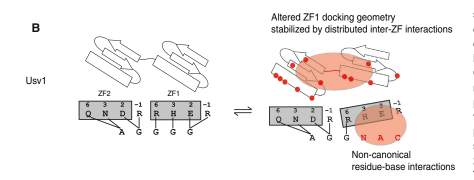
<sup>(</sup>A and B) Binding profile comparisons (as in Figure 1) for Com2 and Com2 RK  $\rightarrow$  EE mutant relative to Msn2. Com2-preferred sites (as in Figure 1B) are highlighted (orange). Four 8 bp sequences (labeled A, B, C and D) assayed in EMSAs are in red.

<sup>(</sup>C) Comparison of Com2 and Com2 RK  $\rightarrow$  EE mutant binding to select binding sites. DNA base differences from the common site TCAGGGG (Figure 1D) are in red. Scores are mean Z scores for the eight different 8-mers containing the 7-mer sites shown (error bars = 1 SD). (\*\*) p < 10<sup>-3</sup>; (\*\*\*) p < 10<sup>-4</sup>, unpaired Student's t test. (D and E) Binding profile comparisons for Usv1 two-ZF and 4-Res mutants shown relative to Usv1. Usv1-preferred sites (as in Figure 1D) are highlighted (orange); the subset of sites preferentially bound by Usv1 relative to Usv1 4-Res mutant are highlighted (magenta). Approximate positions of canonical recognition residues (gray dots) and mutated residues (red dots) are illustrated in ZF cartoons.

## Molecular Cell Diversification of Zinc Finger DNA Binding



AT-stabilization by RGR motif via DNA minor groove contacts



## Figure 4. Models for TF-Specific Binding Modes

(A and B) Binding schematics depict key features of the canonical ZF binding mode and the TFspecific binding modes that facilitate the binding to the Msn2-family common and TF-preferred sequences. Models are presented for (A) Com2 and (B) Usv1 (same model applies for Rgm1). Key features proposed for the TF-specific binding modes are highlighted. Residues mutated in Msn2 and Usv1 constructs (Figure 2) that affected DNA binding are highlighted (red dots).

selectively abrogated), and do not involve changes in ZF number or canonical residue identity. One mechanism involves a basic RGRK motif found N-terminal to Com2 that stabilizes the binding to ATrich DNA sites (Figures 1 and 2). We propose that the RGRK motif works similarly to other RXR motifs in which Arg residues project into the DNA minor groove and stabilize AT-rich sequences (Rohs et al., 2009) (Figure 4A). The *D. melanogaster* Trl (GAGA) ZF protein also uses a basic region N-terminal to the ZF to mediate minor

forkhead modes are completely different (i.e., 5'-GTAAACAA versus 5'-GACGC), whereas in this study we find that different binding modes exhibit different preferences over only a portion of the binding site (Figures 1 and 3), suggesting different mechanisms are operating. Future studies using similar approaches will be highly informative to delineate both the mechanisms and the evolutionary history of TF diversification by the gain and loss of alternate binding modes. In addition to altering a TF's target sites, alternate binding modes provide a mechanism for DNA allostery whereby DNA sequence differences can affect protein structure and result in differences in cofactor recruitment and transcriptional activity (reviewed in Siggers and Gordân, 2014). Our data support a model in which the ZF proteins adopt different conformations based on the DNA sequence. Future studies are needed to investigate whether the alternate binding modes and target sequences identified for the Msn2-family might relate to differential cofactor recruitment and transcriptional activity.

C2H2 ZFs have been studied as a paradigm of modularity in DNA binding and as a powerful scaffold for designing synthetic ZFs (Klug, 2010). For protein engineers, the appeal of this family has been the ability to model and manipulate DNA-binding specificity by altering a small set of canonical residues. While it is widely appreciated that residues outside of the canonical recognition positions affect DNA binding, the mechanisms by which these residues alter binding remain unclear (Lam et al., 2011; Persikov et al., 2014; Persikov and Singh, 2011; Ramirez et al., 2008). Here we present two mechanisms for altering ZF DNAbinding specificity that operate via stabilization of alternate binding modes. Importantly, these alternate modes bind DNA with comparable affinity, operate in a modular fashion (i.e., can be groove interactions; however, its role in binding specificity remains unknown (Omichinski et al., 1997). Analysis of existing ZF structures suggests that the RGRK motif in Com2 could reach and interact with the AT base pairs (as in Figure 4) via the minor groove. The second mechanism we identified operates via a distributed set of amino acids throughout the two ZFs and inter-ZF linker of Usv1 (Figures 1 and 2). We propose a model in which an altered binding mode, or docking geometry, of ZF1 is stabilized by interdomain residue-residue contacts (Figure 4B). In this altered binding mode, alternate DNA-base contacts are made, either by the residues at the canonical recognition positions or by alternate, noncanonical recognition residues.

Our results that ZF proteins can switch between alternate binding modes to recognize different DNA sites (Figure 4) has implications for predicting and designing ZF DNA binding. First, the ability to selectively abrogate binding to a subset of the DNA sequences presents a conceptual complication to representing the DNA-binding specificity by single binding models such as a position weight matrix (PWM) (Enuameh et al., 2013; Gupta et al., 2014; Jolma et al., 2013; Zhao and Stormo, 2011). Second, the ability to switch between different binding modes and bind different DNA sites could lead to off-target binding in a ZF design experiment-one might optimize residues to bind select target sites only to find that an alternate mode permitted binding to undesired sites. The existence of multiple binding modes further motivates the utility of selection assays and the inability to simply "stich-together" ZF domains of characterized specificity (Beerli et al., 1998; Choo and Klug, 1997; Enuameh et al., 2013; Klug, 2010; Persikov et al., 2014; Wolfe et al., 2000) and suggests that studies aimed at finding ways to inhibit alternate binding modes may minimize off-target binding of synthetic ZF proteins.

#### **EXPERIMENTAL PROCEDURES**

GST-tagged proteins were expressed and purified from bacteria or made by in vitro transcription translation (IVT); all ZF constructs are listed in Table S3. PBM experiments and analysis were carried out as previously described (Berger and Bulyk, 2006, 2009); all PBM data are provided in Tables S3 and S4. Genome analyses were performed using custom Perl scripts (available on request), and clustering was performed using the R statistical package. (See Supplemental Experimental Procedures for full details.)

#### SUPPLEMENTAL INFORMATION

Supplemental Information includes three figures, four tables, and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.molcel.2014.06.019.

#### **AUTHOR CONTRIBUTIONS**

T.S. and J.R. cloned TF ZF domains; B.B. performed protein expression, purification, and EMSA binding assays; T.S., J.R., and B.B. performed PBM experiments; T.S. performed data analysis; T.S. and M.L.B. designed the study and the experiments; and T.S. and M.L.B. wrote the manuscript.

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