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# Ride the wavelet: A multiscale analysis of genomic contexts flanking small insertions and deletions

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Recent studies have revealed that insertions and deletions (indels) are more different in their formation than previously assumed. What remains enigmatic is how the local DNA sequence context contributes to these differences. To investigate the relative impact of various molecular mechanisms to indel formation, we analyzed sequence contexts of indels in the non protein- or RNA-coding, nonrepetitive (NCNR) portion of the human genome. We considered small ( $\leq 30$ -bp) indels occurring in the human lineage since its divergence from chimpanzee and used wavelet techniques to study, simultaneously for multiple scales, the spatial patterns of short sequence motifs associated with indel mutagenesis. In particular, we focused on motifs associated with DNA polymerase activity, topoisomerase cleavage, double-strand breaks (DSBs), and their repair. We came to the following conclusions. First, many motifs are characterized by unique enrichment profiles in the vicinity of indels vs. indel-free portions of the genome, verifying the importance of sequence context in indel mutagenesis. Second, only limited similarity in motif frequency profiles is evident flanking insertions vs. deletions, confirming differences in their mutagenesis. Third, substantial similarity in frequency profiles exists between pairs of individual motifs flanking insertions (and separately deletions), suggesting “cooperation” among motifs, and thus molecular mechanisms, during indel formation. Fourth, the wavelet analyses demonstrate that all these patterns are highly dependent on scale (the size of an interval considered). Finally, our results depict a model of indel mutagenesis comprising both replication and recombination (via repair of paused replication forks and site-specific recombination).

[Supplemental material is available online at [www.genome.org](http://www.genome.org) and at [http://bx.psu.edu/makova\\_lab/](http://bx.psu.edu/makova_lab/).]

Indels cause multiple human genetic diseases (Cooper et al. 2006) and are a source of natural intra- and interspecific genetic variation (e.g., Mills et al. 2006; Clark et al. 2007). Recent studies highlight the differences in rates of small ( $\leq 100$ -bp) insertions vs. deletions (e.g., Chen et al. 2007; Messer and Arndt 2007) and suggest that these two types of mutations might at least in part be caused by distinct molecular mechanisms (Kvikstad et al. 2007). In contrast to earlier reports attributing all indels to replication slippage errors (Ball et al. 2005; Messer and Arndt 2007), we have previously demonstrated the significance of recombination to the formation of small indels, in particular insertions (Kvikstad et al. 2007). Several analyses of protein coding regions have demonstrated sequence context dissimilarities between insertions and deletions, such as differences in local base composition and distinct hotspots (Halangoda et al. 2001; Kondrashov and Rogozin 2004; Ball et al. 2005), which again indicate differences in their mutagenesis. Recently, Tanay and Siggia (2008) analyzed the base composition of sequences flanking insertions and deletions and developed models of genome-wide indel propensity that differ between the two mutation types.

Despite these exciting advances, many questions still remain. For example, it is not known to what extent natural selection could have affected signatures of hotspots and motifs in the vicinity of indels identified in genic sequences (e.g., Kondrashov and Rogozin 2004; Ball et al. 2005). Heterogeneity in base composition and substitution rate along the genome may have confounded some

results (e.g., Halangoda et al. 2001; Tanay and Siggia 2008). Additionally, just as combinations of specific binding sites act jointly to promote transcription (for review, see Ji and Wong 2006), the simultaneous presence of multiple motifs could provide important clues to indel mutagenesis and requires investigation. Finally, determining the scale at which a hotspot or a motif can be detected and implicated in a particular molecular process constitutes an active area of research (e.g., Berry et al. 2006). Thus, analyzing sequence contexts flanking a genome-wide set of indels located outside of genes with a multiscale methodology is likely to aid in discerning the biological mechanisms underlying these mutations.

Here we utilize wavelet techniques to study sequences flanking small indels in the NCNR portion of the human genome. Wavelet transformations, traditionally used in the study of time series, allow one to re-express a signal preserving both its global trends and its local fluctuations (Percival and Walden 2006). For wavelet analysis of DNA, the nucleotide sequence is regarded as “time,” and features mapped along the sequence represent the “signal.” A wavelet transformation is multiscale since a scale (or window size) is not chosen a priori; rather, a signal is analyzed across multiple scales simultaneously. Exploiting this fact, we expand on previous presence/absence enrichment tests (Abeyasinghe et al. 2003; Kondrashov and Rogozin 2004; Ball et al. 2005) and use wavelet transformations for the detection of spatial patterns (i.e., local fluctuations in occurrence) among sequence motifs flanking small indels. To our knowledge, this represents the first application of wavelet techniques to the entire human genome.

Specifically, we address the following questions. First, are regions flanking indels similar or different in terms of their motif content when compared to the indel-free portion of the genome? Second, can we discern significance in the spatial patterns of

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motifs in proximity to indels? Third, do motifs show similar or different spatial patterns when flanking insertions vs. deletions? And finally, is it possible to detect colocation of different motifs, indicative of the simultaneous action of different mechanisms in the generation of indels?

## Results

Small indels occurring in the human lineage since its divergence from chimpanzee were identified in the NCNR portion of the genome (see Methods; Supplemental material). A total of 65,353 and 135,989 human-specific insertions and deletions, respectively, were detected in the human–chimpanzee–macaque three-way MULTIZ alignments (Rhesus Macaque Genome Sequencing and Analysis Consortium 2007), as described by Kvikstad et al. (2007). An excess of deletions over insertions is consistent with other studies (Chen et al. 2007; Kvikstad et al. 2007; Messer and Arndt 2007; Tanay and Siggia 2008).

To implement our analyses, we created five nonoverlapping human “subgenomes” (sets of sequences): four subgenomes corresponded to the regions flanking insertion or deletion breakpoints, separately, and in 5′ or 3′ position along the chromosome (in the reference genome), separately. The 5′ and 3′ subgenomes were considered separately because asymmetries in processing of stalled replication forks and double-strand break (DSB) repair have been noted for motifs in specific strand orientations (Kosmider and Wells 2006; Pollard et al. 2007); both processes potentially contribute to indel formation (see below). Additionally, a fifth (“control”) subgenome was built from the NCNR indel- and microsatellite-free portion of the human genome (see Methods).

Each subgenome was screened for the presence of motifs known to promote recombination, induce replication pausing and/or frameshift, and affect DNA stability, as well as several insertion/deletion hotspots (Table 1; Abeyasinghe et al. 2003; Ball et al. 2005). We detected matches to each motif, its complement, mirror image (reverse), and reverse complement. As a result, a total of 126 motifs grouped into eight classes were sought. For each motif, we computed a total frequency profile in each subgenome by summing motif counts across all insertion/deletion events in contiguous 10-bp increments (see Methods). Total frequency profiles thus “average out” potential genome landscape effects. Moreover, no significant base composition differences were detected between indel and control subgenomes ( $P$ -values of 0.701, 0.671, 0.134, 0.127 for deletions 5′, deletions 3′, insertions 5′, and insertions 3′, respectively; Wilcoxon two-sided nonparametric tests). Hence, genome-wide heterogeneity in base composition is unlikely to affect motif detection in each subgenome.

If the distinction between 5′ vs. 3′ position with respect to an indel breakpoint were unimportant, the total frequency profiles of any given motif should be symmetric 5′ vs. 3′, showing high correlation (e.g., as measured nonparametrically by Kendall tau). In contrast with this expectation, in our data set many motifs display asymmetrical frequency behaviors 5′ and 3′ to indel breakpoints (for one example motif, topoisomerase cleavage site 4; see Supplemental Fig. S1). Indeed, the 5′ vs. 3′ correlations of total frequency profiles are relatively weak for all motifs considered here (Kendall tau coefficients are all  $<0.5$ —computed separately for insertions′ and deletions′ flanks), suggesting differences in motif occurrences on the two sides of breakpoints. Similarly, if indel mutations were equally likely to occur on the two DNA strands, then any given motif involved in indel mutagenesis and its reverse complement should show high 5′ vs. 3′ correlation (i.e., similarity

between a motif’s occurrence and its reverse complement’s occurrence on the opposite strand). Conversely, the 5′ vs. 3′ correlations of total frequency profiles for each motif (e.g., 5′) and its reverse complement (e.g., 3′) are relatively weak (Kendall tau coefficients are all  $<0.4$ —again, computed separately for insertions′ and deletions′ flanks).

We assessed the significance of 5′ vs. 3′ positional differences with a permutation scheme based on randomly reassigning 5′ and 3′ flank labels (Supplemental Fig. S1; Supplemental Table S1; see Methods). Interestingly, nearly twice as many motifs exhibit significant positional asymmetry when flanking deletions than insertions (Supplemental Table S1), and the flank closest to the indel (first 10 bp from the breakpoint) displays extreme positional asymmetry for as many as 25% of the motifs analyzed here (Supplemental Table S1; Supplemental Fig. S1)—no such bias was observed in the control subgenome. Significance of motif positional differences prompted us to consider frequency profiles separately for 5′ and 3′ subgenomes.

### Simple (nonwavelet) motif overrepresentation

We commenced with an analysis of global trends using a simple (nonwavelet-based) procedure to detect motif overrepresentation in indel flanking sequences, because overrepresentation of particular motifs in the vicinity of indels suggests their importance to indel formation. For a given motif, its total occurrences were computed for each of the four indel-related subgenomes and contrasted with the control subgenome using a resampling scheme, separately at various fixed distances from indel breakpoint (Methods; Fig. 1; Supplemental Table S2). Importantly, a comparison of motif total occurrences aggregated across all distances (for each subgenome) reveals no striking differences (Supplemental Table S3), demonstrating the importance of scale for the discriminatory power of simple presence/absence overrepresentation testing.

DNA polymerase (pol) pause/frameshift hotspots, topoisomerase cleavage sites, and motifs promoting site-specific recombination are significantly overrepresented in sequences flanking insertions and deletions alike (Fig. 1). However, there are noticeable differences between insertions and deletions in the proportions of overrepresented motifs by class and by distance from the breakpoint. For instance, at all but the largest (320 and 640 bp) distances, DNA pol pause/frameshift hotspots represent a larger proportion of the overrepresented motifs proximal to insertions than to deletions. In contrast, at even the smallest (10 bp) distance, a higher proportion of topoisomerase cleavage sites are overrepresented in proximity to deletions than insertions. Among them, the example motif topoisomerase cleavage site 4 displays overrepresentation near deletions in 5′ orientation at all distances (Supplemental Table S2).

Our analysis of the NCNR portion of the genome indicates overrepresentation of V(D)J recombination signals, X-like sites, and translin binding targets in the vicinity of indels (Fig. 1), while no such trends were observed in previous reports focusing on indels located in genes (Ball et al. 2005)—confirming ascertainment bias when considering genic sequences. Surprisingly, both insertion hotspots and deletion hotspots (Steinmetz et al. 1987; Krawczak and Cooper 1991; Chuzhanova et al. 2002; Kondrashov and Rogozin 2004) display overrepresentation around both insertions and deletions, depending on the distance analyzed (Fig. 1). Deletion hotspots in particular are overrepresented flanking insertions and deletions at the smallest distance considered (10 bp).

**Table 1.** DNA sequence motifs

Motif class	No.	Motif name	Reference	Sequence
Deletion hotspots	1	"Deletion hotspot 1" <sup>a</sup>	Cooper and Krawczak (1993)	M <sup>b</sup> KRRGT
	2	"Deletion hotspot 2"	Kondrashov and Rogozin (2004)	YYTG
	3	Murine MHC hotspot	Steinmetz et al. (1987)	CAGR
DNA pol pause/ frameshift hotspots	1	Alpha pause site core sequence	Abeyasinghe et al. (2003); Ball et al. (2005)	GAG
	2	Alpha pause site core sequence	Abeyasinghe et al. (2003); Ball et al. (2005)	ACG
	3	Alpha pause site core sequence	Abeyasinghe et al. (2003); Ball et al. (2005)	GCS
	4	Polymerase arrest site	Abeyasinghe et al. (2003); Ball et al. (2005)	WGGAG
	5	Alpha frameshift	Abeyasinghe et al. (2003); Ball et al. (2005)	TCCCCC
	6	Alpha frameshift	Abeyasinghe et al. (2003); Ball et al. (2005)	CTGGCG
	7	Beta frameshift	Abeyasinghe et al. (2003); Ball et al. (2005)	ACCCWR
	8	Beta frameshift	Abeyasinghe et al. (2003); Ball et al. (2005)	TTTT
	9	Alpha/beta frameshift	Abeyasinghe et al. (2003); Ball et al. (2005)	TGGNGT
	10	Alpha/beta frameshift	Abeyasinghe et al. (2003); Ball et al. (2005)	ACCCCA
Indel hotspot	1	"Indel hotspot"	Chuzhanova (2002)	GTAAGT
	1	"Insertion hotspot 1"	Kondrashov and Rogozin (2004)	ATMMGCC
Insertion hotspots	2	"Insertion hotspot 2"	Kondrashov and Rogozin (2004)	TACCRG
	1	Topoisomerase I consensus cleavage site	Abeyasinghe et al. (2003); Ball et al. (2005)	CAT
Topoisomerase cleavage sites	2	Topoisomerase I consensus cleavage site	Abeyasinghe et al. (2003); Ball et al. (2005)	CTY
	3	Topoisomerase I consensus cleavage site	Abeyasinghe et al. (2003); Ball et al. (2005)	GTY
	4	Topoisomerase I consensus cleavage site	Abeyasinghe et al. (2003); Ball et al. (2005)	RAK
	5	Topoisomerase I consensus cleavage site	Abeyasinghe et al. (2003); Ball et al. (2005)	YCCTT
	6	Topoisomerase I consensus cleavage site	Abeyasinghe et al. (2003); Ball et al. (2005)	YTA
	Translin targets	1	"Translin target 1"	Abeyasinghe et al. (2003); Ball et al. (2005)
2		"Translin target 2"	Abeyasinghe et al. (2003); Ball et al. (2005)	GCCCWSSW
V(D)J recombination signals	1	Heptamer	Abeyasinghe et al. (2003); Ball et al. (2005)	CACAGTG
	2	Nonamer	Abeyasinghe et al. (2003); Ball et al. (2005)	ACAAAAACC
X-like sites	1	Fragile X breakpoint cluster	Abeyasinghe et al. (2003); Ball et al. (2005)	CCG
	2	X element	Abeyasinghe et al. (2003); Ball et al. (2005)	GCTGGTGG
	3	Hypervariable minisatellite core sequence	Abeyasinghe et al. (2003); Ball et al. (2005)	GGCAGGANG
	4	minisatellite X-like element	Abeyasinghe et al. (2003); Ball et al. (2005)	GCWGGWGG
	7	Human hypervariable minisatellite core sequence	Abeyasinghe et al. (2003); Ball et al. (2005)	GGAGGTGGGCAGGARG <sup>c</sup>
	8	Human hypervariable minisatellite recombination sequence	Abeyasinghe et al. (2003); Ball et al. (2005)	AGAGGTGGGCAGGTGG <sup>c</sup>

<sup>a</sup>Motif names in quotation marks were given for reference in this work only.

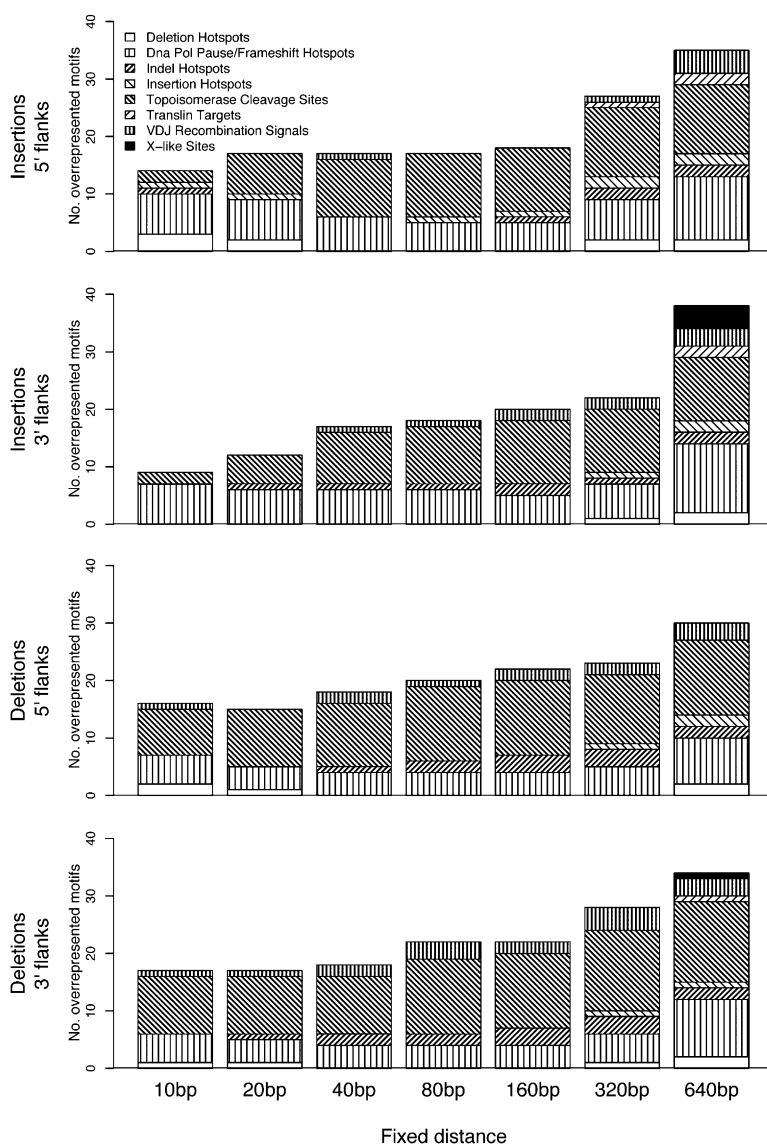
<sup>b</sup>IUPAC ambiguities: R = A/G, Y = C/T, K = G/T, M = A/C, S = G/C, W = A/T, N=A/C/G/T.

<sup>c</sup>Motifs longer than 10 nucleotides (unit for wavelet transform) were sought for simple enrichment test only, at scales of 20-bp and larger.

### Multiscale wavelet analyses

In addition to overrepresentation, the spatial pattern in a motif's occurrence can also convey information suggestive of a role in indel mutagenesis. For a given scale, the motif could have similar

global trends in indel flanks and controls (i.e., no overrepresentation), but a very different profile of local fluctuations (i.e., spatial pattern), or vice versa. In the following, we utilize wavelet transformations to investigate motifs' spatial patterns, at multiple scales simultaneously.



**Figure 1.** Distribution of overrepresented motifs (nonwavelet based) across motif classes (patterned areas) and distances from the breakpoint (bars), separately for the 5' and 3' flanks of insertions and deletions. Patterned areas are proportional to the number of significant motifs per class (significance is assessed resampling from the control subgenome, and subject to a false discovery rate [FDR] correction; FDR controlled at 5%).

### Motif enrichment profiles

For a given motif, we created four enrichment profiles, each computed as the difference between its total frequency profile (the sum of motif counts across all insertion/deletion events in contiguous 10-bp increments, see above) from an indel-related subgenome and its corresponding profile from the control subgenome (Fig. 2A; see Methods). A multiscale analysis using wavelet-transformed enrichment profiles and their second moments was conducted to assess significance of the magnitude of each enrichment profile (see Methods). This approach is illustrated for the example motif topoisomerase cleavage site 4, which displays a significant enrichment profile flanking deletions in 5' position (Fig. 2A; Supplemental S2).

Results summarized by motif class (Fig. 3; for individual motifs, see Supplemental Table S4) reveal that a greater number of

motifs have significant enrichment profiles flanking indels at intermediate to large (160–640 bp) than at small scales. Nevertheless, DNA pol pause/frameshift motifs exhibit significant enrichment profiles flanking deletions at the finest (10–20 bp) scales (only 5' with respect to breakpoint); for insertions, they become significant at intermediate (80 bp) scales.

Interestingly, there are motif classes (e.g., insertion and deletion hotspots, topoisomerase cleavage sites) that display both significant global trends as measured by nonwavelet overrepresentation (Fig. 1) and significant local fluctuations captured by enrichment profiles (Fig. 3). However, some motif classes display significance in only one of these two measures. For instance, despite lacking significant overrepresentation 5' to deletions (Fig. 1), translin target binding sites exhibit significant enrichment profiles in this position flanking deletions at a 160-bp scale (and never for insertions) (Fig. 3). Conversely, neither indel hotspots nor VDJ recombination signals exhibit significant enrichment profiles (Fig. 3), although they are significantly overrepresented proximal to both insertions and deletions (Fig. 1). These results corroborate differences in the information carried by total occurrences versus spatial patterns.

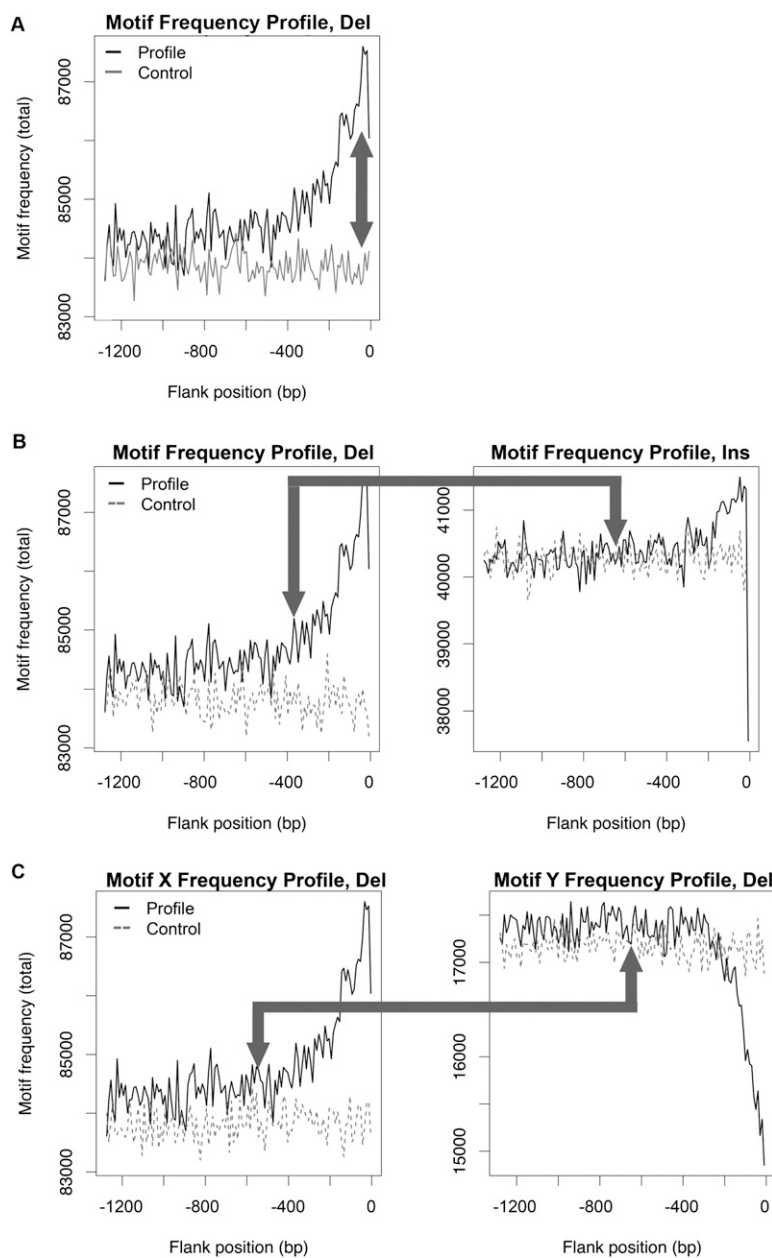
Potentially, indels of different sizes (e.g., 1-bp events) or evolutionary “ages” (times of occurrence, e.g., events polymorphic in human populations) could show distinct motif enrichment profiles (see Supplemental material; Supplemental Tables S5–S8) suggestive of distinct mutagenic mechanisms. However, preliminary investigation of data from chromosome 1 indicates that the majority of motifs with significant enrichment profiles around 1 bp or polymorphic indels represent subsets of the motifs identified considering all events. Thus, variation

due to indel size and/or evolutionary “age” is unlikely to significantly affect our conclusions.

### Similarity between motif frequency profiles flanking insertions and deletions

Next, we investigated whether a motif presents similar spatial patterns flanking both insertions and deletions, suggesting shared mechanisms of origin for these two mutation types. To measure similarity, we used the coefficients of wavelet-transformed frequency profiles to compute multiscale correlations between insertion and deletion behavior (a motif with itself) in the 5' flank, and separately, in the 3' flank (Fig. 2B; Supplemental Fig. S3; Methods).

Unexpectedly, we discover significant similarity in spatial patterns (significantly positive Kendall tau correlation) around insertions and deletions for only a limited number of motifs at



**Figure 2.** Multiscale wavelet analyses (for details, see text; Supplemental Figs. S2–S4): total frequency profiles illustrating three types of comparisons (arrows) using topoisomerase cleavage site 4 as an example motif. (A) Indel vs. control: enrichment profiles. The spatial occurrence pattern characterizing a motif is investigated forming an enrichment profile, i.e., the difference between the motif's total frequency profile in an indel-related subgenome (e.g., the 5' flank of deletions; black line) and that in the corresponding control subgenome (gray line). (B) Insertions vs. deletions: similarity between profiles. This is investigated by comparing the motif's total frequency profile in a deletion-related subgenome (e.g., the 5' flank; black line; left) with that in the corresponding insertion-related subgenome (black line; right). Control profiles (dashed lines) are provided for visual reference only. (C) Motif X vs. motif Y: colocation in profiles. Along with topoisomerase cleavage site 4 (X), here we consider DNA pol pause/frameshift hotspot 1 (Y). Colocation is investigated by comparing the total frequency profiles of X vs. Y in an indel-related subgenome (e.g., the 5' flank of deletions; black lines for X on the left, and Y on the right). Control profiles (dashed lines) are provided for visual reference only.

small scales ( $\leq 80$ -bp): 13 and 12 in 5' and 3' orientations, respectively (Fig. 4, two motifs are significantly correlated in both orientations while only six motifs are among the top 25% most "abundant" motifs identified; Supplemental Table S3). DNA pol

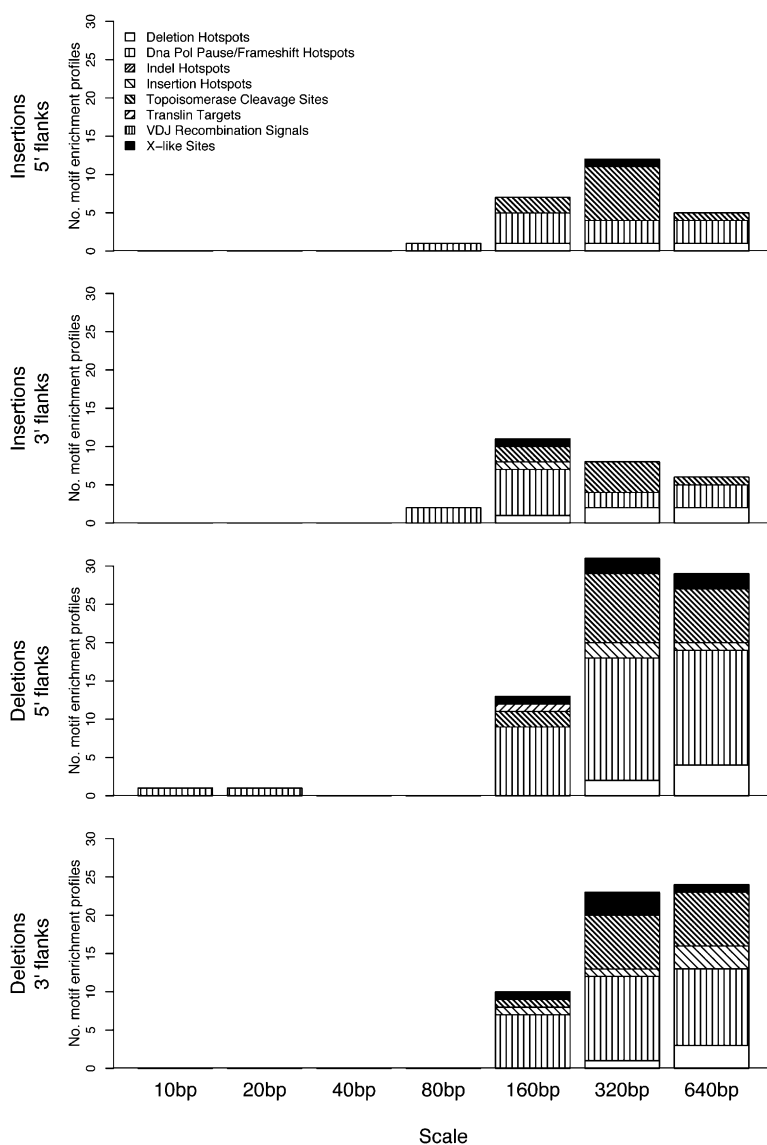
pause/frameshift hotspots and topoisomerase cleavage sites possess significant similarity between insertions and deletions, as do several of the insertion and deletion hotspots (Fig. 4), confirming that the latter motifs fail to discriminate the two mutation types (see above). V(D)J recombination signals and X-like sites also have significant similarity between insertions and deletions, but only 5' from the breakpoint (Fig. 4A). Such a small number of motifs with significant similarity in spatial patterns around insertions and deletions is surprising given that many motifs were previously thought to play similar roles during generation of both mutation types (Ball et al. 2005). Moreover, three motifs show significant negative correlations in spatial patterns around insertions vs. deletions, including the example topoisomerase cleavage site 4 (Fig. 2B; Supplemental Fig. S3), suggesting that these motifs may differentiate between the two mutation types (Fig. 4).

#### *Similarity in frequency profiles between pairs of motifs (colocation)*

Last, we screened for potential signatures of simultaneous mutagenic action for pairs of motifs, as indicated by colocation around indels. The coefficients of wavelet-transformed frequency profiles for two motifs, say X and Y (Fig. 2C; Supplemental Fig. S4), were used to compute multiscale Kendall tau correlations between their spatial occurrences, separately for 5' and 3' flanks, and for insertions and deletions. Significantly positive correlations suggest "cooperation." Figure 2C (for details, see Supplemental Fig. S4) compares the frequency profiles for our example motif topoisomerase cleavage site 4 and a DNA pol pause/frameshift hotspot, both considered in the 5' flank of deletions. At small scales, these two motifs show significantly positive correlation in spatial occurrence patterns, whereas at larger scales their occurrences are anti-correlated (although this is not significant) (Supplemental Fig. S4). This fine-scale colocation would remain undetected if correlation were assessed at larger scales only. As there are 7875 pairs of motifs (and thus 7875 tests for each of the four subgenomes; see Methods), we present summary results for motifs grouped by class and across all scales

(Table 2; for scale-specific results, see Supplemental Table S9).

We detected 185 (261) and 220 (289) significantly correlated pairs of motifs in 5' and 3' positions from insertion (deletion) breakpoint. Despite similar overall numbers, insertions



**Figure 3.** Distribution of motifs with significant enrichment profiles (wavelet-based) across motif classes (patterned areas) and scales (bars), separately for the 5' and 3' flanks of insertions and deletions. Patterned areas are proportional to the number of significant motifs per class (magnitude is measured taking second moments after wavelet transform of each enrichment profile, and significance is assessed through a random permutation scheme, and subject to a FDR correction; FDR controlled at 5%).

and deletions display noticeably different trends (Table 2). For instance, translin targets positively correlate with more motifs when flanking deletions than insertions in 3' positions, while these motifs are more frequently observed to anti-correlate in pairwise comparisons flanking insertions (translin targets are found in relatively similar abundance flanking both mutation types) (Supplemental Table S3). Interestingly, indel hotspots positively correlate with a greater number of motif classes (including V(D)J recombination signals and DNA pol pause/frameshift hotspots) when proximal to insertions than deletions.

In addition to the differences described above, there are colocation behaviors that are shared between insertions and deletions. Among them, the deletion hotspots positively correlate with similar motif classes 5' and 3' to both insertions and dele-

tions (Table 2), confirming that these motifs do not distinguish the two types of mutations (see also Figs. 1, 3, 4). Similarly, different topoisomerase cleavage sites, DNA pol pause/frameshift hotspots, and X-like sites (that are among the most abundant motifs) (Supplemental Table S3) positively correlate within and among these motif classes at multiple scales when flanking both insertions and deletions (Table 2; Supplemental Table S9).

## Discussion

Our novel wavelet-based analyses of motifs associated with DNA pol activity, topoisomerase cleavage, DSBs, and their repair emphasize differences between insertions and deletions in NCNR DNA. Additionally, we find motifs possessing patterns of overrepresentation, spatial occurrence, and colocation—analyzed here for the first time—that frequently differ in the proximity of insertions versus deletions, and at different scales. Taken together, these results provide clues to the relative contributions of different mechanisms associated with indel mutagenesis.

## Importance of replication

Consistent with previous reports (Ball et al. 2005), we observe that DNA pol pausing and/or frameshift motifs are important to indel formation. The analyses of overrepresentation, enrichment profiles, and similarity in spatial patterns of these motifs flanking insertions and deletions (Figs. 1, 3, 4) all suggest that replication is critical for indel mutagenesis (Fig. 5). Note that the DNA pol hotspots analyzed here correspond to eukaryotic polymerases alpha and beta (Table 1; Moon et al. 2007; McCulloch and Kunkel 2008). In particular, the former polymerase primarily synthesizes primer extension for Okazaki fragments on the lagging strand (McCulloch and Kunkel 2008), whereas the latter functions in DNA synthesis during base excision repair (Sweasy et al. 2006; Moon et al. 2007). The significance of DNA pol alpha motifs at small scales (e.g., Fig. 4; Supplemental Tables S2–S4) indicates that lesions might be more likely to escape repair on the lagging strand (Fig. 5) by bypassing replication checkpoint response (Tourriere and Pasero 2007; McCulloch and Kunkel 2008). Notably, we demonstrate that frequency profiles of motifs involved in pausing/frameshift are highly correlated with those of topoisomerase cleavage sites and other motif classes (translin binding targets, X-like sites and V(D)J recombination signals; see discussion below) (Table 2), suggesting that indel formation can be affected by the combined action of replication and other processes.

**A**

Deletion Hotspot 1	0.089 (0.35)	0.008 (0.31)	<b>0.73 (&gt;16)</b>
Deletion Hotspot 1_rc	<b>0.39 (&gt;16)</b>	0.1 (0.35)	0.6 (1.01)
Dna Pol Pause/Frameshift Hotspot 2_r	0.2 (0.58)	-0.04 (0.31)	<b>0.68 (1.55)</b>
Dna Pol Pause/Frameshift Hotspot 3	0.17 (0.55)	0.22 (0.47)	<b>0.8 (&gt;16)</b>
Dna Pol Pause/Frameshift Hotspot 3_rc	0.21 (0.58)	0.1 (0.35)	<b>0.68 (1.47)</b>
Insertion Hotspot 1_c	0.025 (0.31)	0.067 (0.35)	<b>0.78 (&gt;16)</b>
Topoisomerase Cleavage Site 2_rc	<b>0.39 (1.42)</b>	0.23 (0.49)	0.43 (0.67)
Topoisomerase Cleavage Site 4	0.17 (0.55)	0.02 (0.31)	<b>-0.77 (1.55)</b>
Topoisomerase Cleavage Site 6_c	<b>0.34 (1.42)</b>	0.15 (0.35)	0.18 (0.41)
VDJ Recombination Signal 1_rc	-0.089 (0.35)	-0.16 (0.35)	<b>0.78 (1.72)</b>
X-like Site 1_c	0.20 (0.58)	0.065 (0.33)	<b>0.72 (1.72)</b>
X-like Site 1_r	0.31 (1.04)	0.3 (0.59)	<b>0.8 (&gt;16)</b>
X-like Site 4_c	0.15 (0.5)	-0.47 (0.94)	<b>-0.67 (1.47)</b>
	20bp	40bp	80bp

**B**

Deletion Hotspot 3_rc	0.001 (0.31)	0.24 (0.51)	<b>0.73 (&gt;16)</b>
Dna Pol Pause/Frameshift Hotspot 1_rc	0.18 (0.55)	0.27 (0.58)	<b>0.83 (&gt;16)</b>
Dna Pol Pause/Frameshift Hotspot 2_rc	-0.12 (0.42)	0.028 (0.3)	<b>0.73 (1.49)</b>
Dna Pol Pause/Frameshift Hotspot 3_r	<b>0.37 (&gt;16)</b>	0.43 (1.02)	<b>0.70 (1.49)</b>
Dna Pol Pause/Frameshift Hotspot 5_c	-0.025 (0.32)	0.053 (0.34)	<b>-0.72 (1.37)</b>
Dna Pol Pause/Frameshift Hotspot 8	-0.046 (0.33)	<b>0.55 (1.55)</b>	0.37 (0.71)
Dna Pol Pause/Frameshift Hotspot 8_r	-0.046 (0.33)	<b>0.55 (&gt;16)</b>	0.37 (0.73)
Insertion Hotspot 1_c	-0.35 (0.82)	-0.057 (0.35)	<b>0.72 (&gt;16)</b>
Topoisomerase Cleavage Site 1_c	-0.073 (0.35)	<b>0.54 (&gt;16)</b>	0.35 (0.71)
Topoisomerase Cleavage Site 2	0.21 (0.6)	0.12 (0.4)	<b>0.78 (&gt;16)</b>
Topoisomerase Cleavage Site 2_rc	0.29 (0.64)	<b>0.58 (&gt;16)</b>	0.1 (0.38)
Topoisomerase Cleavage Site 6	0.084 (0.39)	0.012 (0.3)	<b>0.77 (1.64)</b>
	20bp	40bp	80bp

**Figure 4.** Significant Kendall tau correlations of total motif frequency profiles around insertions vs. deletions, separately for various scales, and for 5' (A) and 3' (B) flanks. Positive and negative correlations are indicated by sign, with bolded cells corresponding to those significant after FDR correction (FDR controlled at 5%;  $-\log_{10}$  of adjusted *P*-values are reported in parentheses, with ">16" signifying a *P*-value of  $<10^{-16}$ ). Motif names are given suffixes to indicate reverse (r), complement (c), and reverse complement (rc). Scales below 20 and above 80 bp are not reported because no motifs had significant correlations at such scales after FDR correction.

### Interplay of recombination and replication in indel formation

The significance of topoisomerase cleavage sites in our analysis suggests that recombination-mediated repair of paused replication forks can contribute to indel mutagenesis (Fig. 5). We found that topoisomerase cleavage sites display overrepresentation, enrichment profiles, and positive profile correlations (Figs. 1, 3, 4) flanking both insertions and deletions. Since the type I class of topoisomerases (represented by the motifs analyzed here) break and rejoin gapped single-stranded DNA (ssDNA) during replica-

tion (Hyrien 2000; Wang 2002; Tourriere and Pasero 2007), these results again support the importance of replication for indel formation (Fig. 5). Moreover, colocation detected between frequency profiles of DNA pol pause/frameshift hotspots and topoisomerase cleavage sites at intermediate (80–160 bp) scales (Supplemental Table S4) may reflect checkpoint mediated restart of paused forks contributing to indel mutations (Fig. 5; Tourriere and Pasero 2007).

### Site-specific recombination and indel formation

Our analyses of the NCNR genome indicate that translin targets, V(D)J recombination signals, and X-like sites (the motifs associated with site-specific recombination and genome instability) might be significant for small indel mutagenesis. First, translin may be involved in the formation of deletions. Enrichment profiles flanking deletions, but not insertions, are observed for translin targets (Fig. 3). These are binding sites for recognition of the translin protein to ssDNA at staggered breaks in the first steps of their repair (Kasai et al. 1997; Erdemir et al. 2002a,b; Sengupta and Rao 2002). Previously, overrepresentation of translin targets implicated NHEJ in mediating chromosomal rearrangements such as translocations and large deletions (Abeysinghe et al. 2003; Gajicka et al. 2006a,b) and microsatellite/telomere repeat expansions/contractions (Jacob et al. 2004). Our results suggest that translin may mediate small deletion formation as well. Additionally, we demonstrate for the first time that translin targets have frequency profiles positively correlated with other motifs (e.g., DNA pol pause/frameshift hotspots and topoisomerase cleavage sites) 3' to deletion breakpoints (Table 2). Translin is thought to act in cellular response to DNA damage via the translin-associated factor X (TRAX) that interacts with the nuclear matrix protein C1D (Erdemir et al. 2002a,b; Cho et al. 2004). Since C1D regulates a number of cellular functions, including replication, NHEJ, homologous recombination, and topoisomerase activity (Erdemir et al. 2002a,b; Cho et al. 2004; Yang et al. 2004), our results suggest that recruitment of DSB repair, possibly mediated by TRAX and C1D, can contribute to indel (particularly deletion) formation (Fig. 5).

Second, V(D)J recombination may lead to both insertions and deletions. V(D)J recombination signals are overrepresented flanking insertions and deletions at almost all distances from the breakpoint (Fig. 1), and have similar frequency profiles 5' to both mutation types (Fig. 4A). As these motifs correspond to the heptamer and nonamer signals required for creating the DSBs and NHEJ of immune receptor molecules (Lu et al. 2007), our results suggest illegitimate recruitment of V(D)J recombination or NHEJ leading to indel mutation (Fig. 5).

Finally, our results indicate that X-like sites may promote insertions and deletions, but in different ways (Fig. 5). We observe that X-like sites are enriched in the vicinity of indels (Fig. 2), particularly in the 3' position to insertion breakpoints. X-like sites include the X element and other conserved sequences known to induce recombination (Table 1; Seitz and Kowalczykowski 2006). In agreement with this, an X-like consensus sequence previously hypothesized to promote insertions and deletions via V(D)J recombination (Wyatt et al. 1992) is significant to insertion formation (for the X-like motif 4, see Fig. 4A; Supplemental Tables S2, S4). Interestingly, the Fragile X breakpoint cluster motif displays significant enrichment profiles at multiple scales flanking deletions but not insertions (Supplemental Table S4). The Fragile X breakpoint cluster motif is known to adopt many non-B

**Table 2.** Number of significant pairwise correlations (Kendall tau's) in frequency profiles of different motifs, flanking insertions (shaded) and deletions (unshaded), grouped by class**(A) 5' Flanking regions**

	Deletion hotspots	DNA pol pause/frameshift hotspots	Indel hotspots	Insertion hotspots	Topoisomerase cleavage sites	Translin targets	VDJ signals	X-like sites	
Insertions	Deletion hotspots $N = 12$	3 (0) 6 (0)	15 (3)	0 (1)	1 (0)	9 (3)	2 (1)	NS	2 (0)
	DNA pol pause/frameshift hotspots $N = 38$	7 (1)	53 (9) 43 (2)	0 (2)	5 (4)	28 (19)	NS	2 (1)	26 (1)
	Indel hotspots $N = 4$	NS <sup>a</sup>	NS	NS	NS	NS	NS	0 (1)	NS
	Insertion hotspots $N = 8$	0 (1)	1 (7)	NS	1 (0)	NS	1 (0)	NS	1 (1)
	Topoisomerase cleavage sites $N = 24$	7 (3)	12 (8)	2 (1)	2 (0)	37 (8) 34 (2)	0 (2)	1 (1)	7 (5)
	Translin targets $N = 8$	NS	1 (2)	NS	NS	NS	NS	NS	1 (0)
	VDJ signals $N = 8$	0 (2)	2 (1)	NS	1 (0)	NS	1 (1)	1 (0)	NS
	X-like sites $N = 24$	2 (0)	22 (3)	NS	0 (1)	2 (0)	NS	NS	3 (0) 4 (1)

**(B) 3' Flanking regions**

	Deletion hotspots	DNA pol pause/frameshift hotspots	Indel hotspots	Insertion hotspots	Topoisomerase cleavage sites	Translin targets	VDJ signals	X-like sites	
Insertions	Deletion hotspots $N = 12$	6 (0) 6 (0)	11 (4)	NS	1 (0)	14 (6)	1 (0)	1 (1)	8 (0)
	DNA pol pause/frameshift hotspots $N = 38$	5 (1)	53 (3) 39 (9)	0 (1)	3 (1)	23 (24)	4 (0)	1 (3)	35 (1)
	Indel hotspots $N = 4$	NS <sup>a</sup>	2 (1)	NS	1 (0)	NS	0 (1)	NS	NS
	Insertion hotspots $N = 8$	1 (0)	1 (1)	1 (0)	NS	2 (2)	NS	1 (1)	1 (0)
	Topoisomerase cleavage sites $N = 24$	14 (2)	19 (11)	2 (0)	1 (1)	34 (8) 36 (3)	7 (2)	1 (0)	4 (10)
	Translin targets $N = 8$	0 (2)	2 (3)	NS	NS	1 (0)	NS	NS	2 (0)
	VDJ signals $N = 8$	NS	1 (3)	1 (0)	2 (0)	2 (1)	1 (0) NS	NS	NS
	X-like sites $N = 24$	3 (0)	24 (4)	NS	NS	7 (1)	NS	1 (0) NS	7 (0) 5 (0)

Results per cell correspond to the number of significant positively (negatively) correlated motif pairs in total for all scales. N corresponds to number of motifs per class. Correlation results for a motif class with itself (diagonal) are restricted to correlations between pairs of different motifs in the same class (see Methods).

<sup>a</sup>NS indicates no significant correlations at any scales after FDR correction (FDR adjusted  $P < 0.05$ ).

DNA structures (Bacolla et al. 2006). Indeed, the recombination-mediated repair of DSBs at this cluster has been reported to differ in induced mutations when present on leading vs. lagging strands (Kosmider and Wells 2006). This motif also displayed significant positional asymmetry (Supplemental Table S1). Thus, recombination-mediated repair of secondary structures at this X-site may lead to more deletions than insertions and in an orientation-dependent manner.

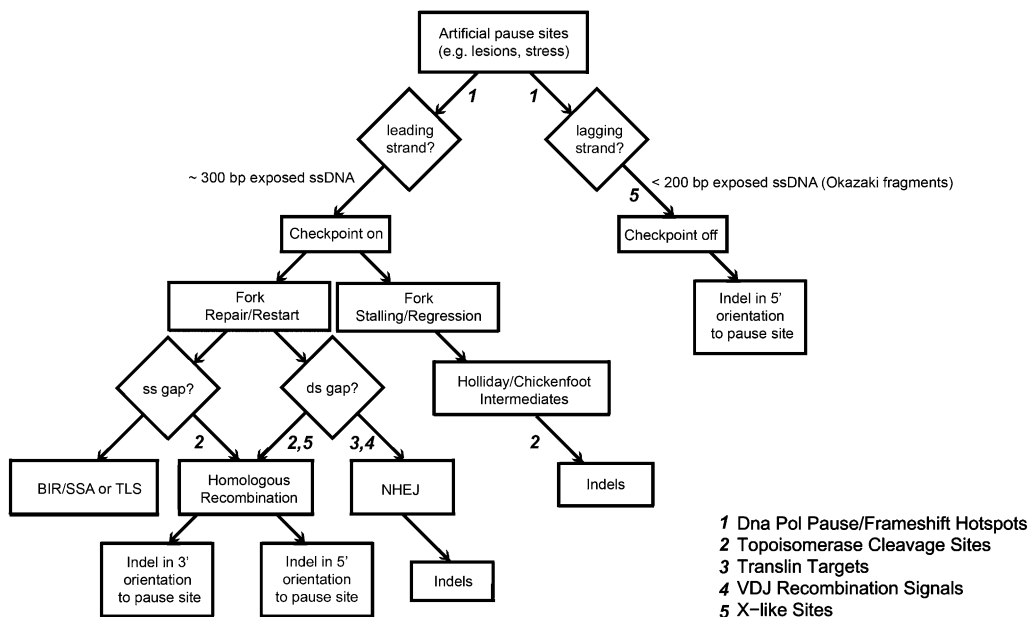
### Chromatin organization and indel formation

From our wavelet-based analyses, the 80-bp scale emerges as particularly relevant given the significance of spatial occurrence patterns at this scale for many motif classes, including DNA pol pause/frameshift hotspots, topoisomerase cleavage sites, V(D)J recombination signals, and X-like sites (e.g., Fig. 4; Supplemental

Table S4). Notably, 80 bp corresponds to approximately half the length of DNA involved in nucleosome positioning (~150 bp) (Luger et al. 1997; Kornberg and Lorch 1999; Segal et al. 2006; Gupta et al. 2008). Since wavelets capture fluctuations in signals representing changes from “peaks” (e.g., nucleosome spacer regions) to “valleys” (e.g., nucleosome-occupied regions), our results corroborate the importance of chromatin structure in the regulation of molecular processes (e.g., Wang 2002; Nightingale et al. 2007) and, as recently noted in medaka, the potential formation of indels (Sasaki et al. 2009).

### Hotspots: Lessons from noncoding DNA

Our results for hotspots reveal similarities in the frequencies of these motifs in insertion- and deletion-flanking regions. Both insertion and deletion hotspots show overrepresentation



**Figure 5.** Replication and recombination pathways proposed to contribute to insertion and deletion mutagenesis (numbers correspond to motif classes indicating involvement as evidenced by results; for details, see Discussion). DNA replication can pause when encountering natural pause sites or artificially due to lesions, stress, and/or specific sequence cues. Depending on the strand, single-stranded DNA (ssDNA) is exposed and replication checkpoint can be triggered or bypassed. Cells respond to replication pausing by repair or restart of the forks, depending on the type of lesion created at the paused site. Double-strand gaps are repaired via homologous recombination or nonhomologous end joining (NHEJ), whereas single-stranded gaps can be repaired via homologous recombination, break-induced replication (BIR) followed by single-strand annealing (SSA), or translesion synthesis (TLS). Our results confirm that NHEJ repair of double-stranded gaps and homologous recombination (of double- and/or single-stranded gaps) could contribute to indel mutagenesis. Stalling of the fork can result in regression and Holliday junction/chickenfoot intermediates, which can be repaired and thus cause indel formation. Checkpoint bypass can also lead to indels.

(Fig. 1), enrichment profiles (Fig. 3), and interesting correlations in frequency profiles (Fig. 4) around both mutation types. Therefore, the classification of hotspots as insertion- or deletion-specific does not hold when analyzing the NCNR genome. Indel (as opposed to insertion or deletion) hotspots behave differently: Their frequency profiles correlate with those of several biological classes of motifs (topoisomerase cleavage sites, V(D)J recombination signals, and DNA pol pause/frameshift hotspots) around insertions, but not deletions (Table 2). Thus, indel hotspots might act in conjunction with other sequences to facilitate the formation of more insertions than deletions, during the processes noted above.

### 5' vs. 3' positional asymmetry

We noted differences in motif occurrence patterns between the 5' and 3' indel-related subgenomes, suggesting the possibility for asymmetrical interactions between DNA flanking sequences and processes leading to indel formation. In contrast to the control subgenome, several motifs exhibit significantly different frequency profiles 5' and 3' of indels (particularly deletions) (Supplemental Fig. S1; Supplemental Table S1). The extreme discrepancy in positional behavior for the flanks closest to the event implies direct involvement of the motif in promoting mutagenesis, particularly for deletions (~85% of small insertions are tandem duplications) (Messer and Arndt 2007). Uncoupling of leading vs. lagging strand synthesis during the processing of stalled replication forks could potentially account for such asymmetry, since as many as a third of human replication forks experience arrest (Conti et al. 2007). Furthermore, the processes of transcription-mediated mutation (Green et al. 2003), homologous

recombination, and chromatin assembly have inferred strand asymmetries based on footprints of their interactions with DNA sequences (Touchon and Rocha 2008). Indeed, processing of stalled replication forks and repair of DSBs at motifs involved in Fragile X Syndrome and Friedreich ataxia, respectively, exhibit specific leading versus lagging strand orientation effects on the subsequent mutational spectra (Kosmider and Wells 2006; Pollard et al. 2007); the same may hold true for motifs analyzed here that likely promote indels.

## Methods

### Indel identification and motif detection

Small ( $\leq 30$ -bp) indels occurring in the human lineage since its divergence from chimpanzee were identified from the human-chimpanzee-macaque (hg18-panTro2-rheMac2) three-way MULTIZ alignments (Blanchette et al. 2004), following the methods described by Kvikstad et al. (2007), which include rigorous filtering to remove false positives likely attributable to issues of sequence, assembly and alignment accuracy (for details, see Supplemental material). Indels were further excluded if they intersected with human microsatellites, as annotated by Kelkar et al. (2008). To reduce potential effects of natural selection (see also Supplemental material), we restricted our analysis to indels occurring in the "noncoding" portion of the human genome by excluding "known genes" plus their 5 kb upstream and downstream regions. We determined that only 0.02% of the resulting sequences overlapped with annotated RNA-coding genes (Blankenberg et al. 2007), less than the genome-wide annotation (Karolchik et al. 2008). Thus, to the best of our ability, we isolated

the nongenic portion of the genome. Furthermore, we excluded repetitive elements annotated by RepeatMasker (<http://www.repeatmasker.org/>) in any of the three species to produce the resulting NCNR genome.

Experimentally determined consensus hotspots or recognition sites for various classes of motifs were obtained from Abeysinghe et al. (2003) and Ball et al. (2005) (Table 1). Computationally predicted indel, insertion, and deletion mutation hotspots, separately, were defined as according to the method of Steinmetz et al. (1987), Krawczak and Cooper (1991), Chuzhanova et al. (2002), and Kondrashov and Rogozin (2004). Perl scripts were developed to screen nucleotide sequences for perfect matches to a given set of input motifs, including match to each motif, its complement, reverse (i.e., mirror image), and reverse complement, separately.

### Motif total frequency profiles

An ordered series was created for each of the four indel-related subgenomes by breaking the sequence upstream (downstream) of each insertion (deletion) into 128 consecutive, nonoverlapping increments of 10 bp. Each 10-bp increment was screened for motif occurrences as described above (motifs overlapping increment endpoints were not scored), and increments were then pooled across all insertions (deletions) for each position relative to the insertion (deletion) breakpoint. Total numbers of occurrences for each motif in each increment were obtained to construct each total frequency profile.

A subset of motifs (Supplemental Table S1) displayed different frequency profiles upstream vs. downstream of indel breakpoints (e.g., Supplemental Fig. S1). To assess the significance of these asymmetries, we reconstructed each motif's total frequency profile after randomly permuting 5' and 3' position labels—this was repeated 100,000 times to create “null” distributions for the profiles, which allowed us to detect significant asymmetries, especially in the first 10-bp increments (the ones closest to the indel event) (Supplemental Table S1).

The control subgenome corresponding to the NCNR indel-free (i.e., excluding 1280 bp 5' and 3' to all indels) and microsatellite-free portion of the human genome was obtained using Galaxy (Blankenberg et al. 2007). To construct control (total) frequency profiles for each motif, we randomly sampled segments of 1280 bp from the control subgenome in equal number to the sequences comprising the insertion (deletion) subgenomes, respectively. These segments were then broken into 128 contiguous 10-bp increments and screened for motif occurrences, applying the procedure used for indel-related subgenomes.

### Simple motif overrepresentation

Overrepresentation of motifs upstream (5') and downstream (3') of insertions and deletions, separately, was assessed in comparison to their occurrence in the control subgenome. This was performed at various fixed distances (10, 20, 40, 80, 160, 320, and 640 bp) from the indel breakpoints. For each distance, a number of sequences equal to that in the corresponding indel-related subgenome was randomly sampled from the control subgenome and scanned to obtain frequencies for each motif. This was repeated 1000 times to create empirical *P*-values for each motif, each event type (insertion, deletion), each flank (5', 3'), and each scale (Supplemental Table S2). These *P*-values were then adjusted for multiple testing according to the method of Benjamini and Hochberg (1995), as to control the FDR. Significance was reported in all cases with an adjusted *P*-value < 0.05 (i.e., capping the FDR at 5%).

### Multiscale analyses using wavelets

Three multiscale analyses were conducted using wavelet transforms (see Supplemental material). In each, the relevant input signal was decomposed using the Haar wavelet basis function to construct a discrete wavelet transformation (similar results were obtained using other wavelet filters; data not shown). All analyses were performed using the wavelet libraries (“waveslim,” “wave-thresh”) available in the R statistical package (R Development Core Team 2005).

#### Motif enrichment profiles

To compare a motif's spatial patterns in 5' and 3' indel-flanking regions to those in control regions, we formed four enrichment profiles for each motif as differences between its total frequency profiles in the four indel-related subgenomes and the corresponding frequency profiles derived from the control subgenome (Fig. 2A). Next, we obtained wavelet decompositions of the enrichment profiles and analyzed the resulting coefficients' second raw moments (squared deviations from zero) on a multiscale basis (Supplemental Fig. S2). These second moments measured the size of the difference between motifs occurrence patterns in indel flanks vs. control regions. Using a random permutation scheme (see Supplemental material), we tested the null hypothesis that this difference is equal to zero (no difference between indel-related subgenome and control), at various scales. This involved 1000 permutations of the frequency profiles' ordered series before computing differences (enrichment profile) and their wavelet decomposition, each followed by the multiscale analysis of second raw moments (Supplemental Fig. S2; Supplemental Table S4). Multiple testing correction was performed as described above for simple motif overrepresentation.

#### Shared profiles: Similarity between motif frequency profiles flanking insertions and deletions

To investigate the similarity of a motif occurrence patterns around insertions and deletions, we sought positive correlations between its total frequency profiles in the 5' and 3' flanks (separately) of the two types of events (Fig. 2B). For each comparison, we obtained the wavelet decompositions of the two frequency profiles in question, and analyzed their Kendall tau (Kendall 1938) rank-based correlations on a multiscale basis (Supplemental Fig. S3). These provided a robust measure of association between motif occurrence patterns in indel flanks and one that was more conservative than either the Pearson correlation or the Spearman's Rho (Colwell and Gillett 1982), resulting in higher *P*-values and broader null bands when assessing significance through permutations (data not shown). Similar to above, we used a random permutation scheme to test the null hypothesis of no association, at various scales. Here, permutations of the frequency profiles' ordered series were performed, and wavelet decompositions were computed, followed by multiscale analyses of Kendall tau correlations (Supplemental Fig. S3). Multiple testing correction was performed as described above for simple motif overrepresentation.

#### Colocation: Similarity in frequency profiles between pairs of motifs

To investigate the similarity of occurrence patterns between two different motifs (colocation), we sought positive correlations between their total frequency profiles in the flanks of insertions and (separately) deletions, 5' and (separately) 3' (Fig. 2C). For each comparison (7875 in total), we obtained the wavelet decompositions of the frequency profiles in question, and once again analyzed the Kendall tau rank-based correlations on a multiscale

basis. Once more, testing for no association was performed with 1000 permutations of the frequency profiles ordered series before computing wavelet decompositions, each followed by the multi-scale analysis of Kendall tau correlations (Supplemental Fig. S4; Supplemental Table S9). Multiple testing correction was performed as described above for simple motif overrepresentation.

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