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SUPPLEMENTARY TEXT

Interval-Wise Testing: statistical details.

The IWT is a novel inferential procedure for functional data¹ that performs a global two-sample test on the whole domain of the curves being compared, and simultaneously detects locations where the difference between the two samples of curves is significant. The IWT was developed in order to overcome weaknesses of the two testing procedures^{2,3} previously proposed in the FDA literature to deal with the same inferential problem. These procedures both required an initial discretization step: the Interval Testing Procedure (ITP)² was based on a basis expansion of the curves, while the procedure developed in³ utilized an a priori partition of the curve domain in smaller intervals. Different discretization choices in this initial step can affect test results and conclusions. Notably, despite this issue, the ITP was successfully employed in⁴ to characterize the genomic landscape surrounding endogenous retrovirus locations in human and mouse. The IWT does not require discretization; it operates directly on the original curves, providing more reliable results. Moreover, being a non-parametric permutation test, it can be employed even if the data distribution is skewed, which is the case in our application to IPD values (Fig. S2). Here we used an extended version of the IWT specifically designed for "Omics" data applications⁵. This extension outputs both the locations and the scales that lead to rejecting a null hypothesis. In addition, it allows the user to select among different test statistics that highlight complementary characteristics of the curve distributions.

Let $IPD_{f_i}(t)$ $i = 1, ..., n_f$ be the IPD curves in the n_f motif-containing windows (features), and $IPD_{c,i}(t)$ $i = 1, ..., n_c$ the IPD curves in the n_c motif-free windows (controls). Each curve is defined in the interval I = (-50, 50) (0 representing the center of the motif for motif-containing windows) and comprises 100 values corresponding to the 100 nucleotides where the IPD is measured. Missing IPD measurements are treated as gaps in the curves. We treat $IPD_{f,i}(t)$ $i = 1, ..., n_f$ and $IPD_{c,i}(t)$ $i = 1, ..., n_c$ as two random samples from two independent random functions, and test the null hypothesis H_0^I that the two random functions have the same distribution over the whole interval I, versus the alternative H_1^I that they have different distributions. When we detect significant differences between the two IPD curve distributions (i.e. when we reject the null hypothesis), we aim to identify the portions of the curves (locations) where these differences occurs. Moreover, we want to select the lengths s = |S| (scales) of the subintervals $S = (t_a, t_b) \subseteq I$ where these differences are strong enough to be detected by restricting the null hypothesis to S (indicated as H_0^S).

For each subinterval $S \subseteq I$, we define the mean test statistic as

$$T_{mean}(S) = \frac{1}{|S|} \int_{S} \left(\overline{IPD_{f}}(t) - \overline{IPD_{c}}(t)\right)^{2} dt$$

where

 $\overline{IPD_f}(t) = \frac{1}{n_f} \sum_{i=1}^{n_f} IPD_{f,i}(t) \quad \text{and} \quad \overline{IPD_c}(t) = \frac{1}{n_c} \sum_{i=1}^{n_c} IPD_{ci}(t)$ are the sample means of the IPD curves in the two groups. Similarly, we define the median and the multi-quantile test statistics as

$$T_{median}(S) = \frac{1}{|S|} \int_{S} \left(IPD_{f}^{0.50}(t) - IPD_{c}^{0.50}(t) \right)^{2} dt$$

and

$$T_{multi-quantile}(S) = \sum_{q \in Q} \frac{1}{|S|} \int_{S} \left(IPD_{f}^{q}(t) - IPD_{c}^{q}(t) \right)^{2} dt$$

where, for every $t \in I$, $IPD_f^q(t)$ and $IPD_c^q(t)$ are the quantiles of order q of the IPD curves in the n_f motif-containing windows and n_c motif-free windows, respectively, and Q is a given set of probabilities. Different statistics allow us to focus on different characteristic of the curve distributions. In particular, if the set Q spans a large portion of [0, 1], the multi-quantile statistic is very effective in leveraging information on the whole curve distributions. For example, we can use the quartiles ($Q = \{0.25, 0.50, 0.75\}$) to capture differences in the central part of the distribution, or we can add smaller and larger quantiles ($Q = \{0.05, 0.25, 0.50, 0.75, 0.95\}$) to capture also differences in the tails.

Given a choice of the test statistic T, the first step of the IWT is a functional permutation test for the hypothesis H_0^S versus H_1^S on every subinterval $S = (t_a, t_b) \subseteq I$ and every complementary interval $S = I \setminus (t_a, t_b)$. In particular, we estimate the empirical distribution of the test statistic T under H_0^S conditionally to the data, by evaluating T(S) for all possible permutations of the $n_f + n_c$ observed curves, and we compute the test p-value p^S as the proportion of permutations that lead to a test statistic greater than or equal to the one evaluated on the original data (two-sided test, note that the test statistic is non-negative). The second step of the IWT generates an adjusted p-value curve $\tilde{p}(t)$, defined in each $t \in I$ as

$$\widetilde{p}(t) = \sup_{S \ni t} p^S.$$

This multiple testing correction controls the interval-wise error rate; that is, $\tilde{p}(t)$ controls the probability of rejecting the null hypothesis H_0^S on every interval $S \subseteq I$ where it is true (see details in⁶). Finally, we identify locations with a significant difference in motif vs motif-free windows by selecting all $t \in I$ such that $\tilde{p}(t) \leq \alpha$, where α is the desired significance level.

In order to detect the scales at which the differences in IPD are significant, the extended IWT evaluates multiple scales, generating an adjusted p-value curve $\tilde{p}_s(t)$ for each scale $s \leq |I|$. In particular, for each fixed s, $\tilde{p}_s(t)$ considers only the subintervals $S \subseteq I$ of length $|S| \leq s$ and thus controls the interval-wise error rate on all intervals of length at most s. As a consequence, the extended IWT identifies significant locations for all possible scales s (i.e. the points $t \in I$ such that $\tilde{p}_s(t) \leq \alpha$).

SUPPLEMENTARY TABLES

Table S1. Nucleotides annotated in non-B DNA motifs in the human genome.

The number of nucleotides annotated for each motif type according to the non-B DB⁷ (and according to STR-FM⁸ for STRs). Nucleotides may be annotated as part of one or more motifs.

Motifs	Sequence Definition according to non-B DB	Counts
Direct repeats	10-50 nt repeated within 5 nt spacer	42,300,423
Mirror repeats	10-100 nt mirrored within 100 nt spacer	77,078,820
Inverted repeats	10-100 nt with reverse complement within 100 nt spacer	133,278,477
A-phased repeats	3 or more A-tracts (3-5 As) 10 nt on center each; Spacers between equal sized A-tracts must contain some non As	10,504,652
Z-DNA motis	G followed by Y (C or T) for at least 10 nt; One strand must be alternating Gs	6,700,444
G-quadruplex motifs	4 or more G-tracts (3-7 Gs) separated by 1-7 nt spacers; Preference for short spacers with Cs and/or Ts	10,102,937
STRs	Tandem repeats of 1-4 base pairs per motif	187,657,110

Table S2. Tested non-B DNA motifs.

The last two columns represent the sample size for each motif type on each strand.

Motif	Structure	On both strands	Number of windows with annotation	Number of windows after filtering for overlaps	Number of windows with IPD on reference strand	Number of windows with IPD on reverse strand
A-Phased repeats	slipped- strand	yes	404,289	26,218	26,142	26,143
Direct repeats	slipped- strand	yes	1,501,567	34,778	34,582	34,594
Inverted repeats	cruciform	yes	6,365,102	470,135	468,525	468,520
Mirror repeats	H-DNA	yes	1,895,543	43,053	39,919	39,932
Z-DNA motifs	Z-DNA	yes	412,600	6,229	6,207	6,209
G-quadruplex motifs	G-quad	no	181,230 (+) 180,213 (-)	13,125 (+) 12,971 (-)	13,049(+) 12,876 (-)	13,046 (+) 12,885 (-)

Table S3. Tested STRs.

We studied the motif-specific effect of STRs by collapsing all alignable motifs using the method described in Table S9. Motifs with less than 15 windows having IPD on reference or reverse strand (in gray) were not analyzed. The last two columns represent the sample size for each motif in the two strands.

	Number of windows with	filtering for	Number of windows with IPD on reference	Number of windows with IPD on reverse
Motif	annotations	overlaps	strand	strand
(A) _n	6,727,074	583,681	581,804	581,800
(C) _n	1,263,551	135,124	134,603	134,600
(G) _n	1,263,833	135,109	134,571	134,564
(T) _n	6,758,517	585,904	583,991	584,027
(AC) _n	1,281,488	127,385	126,947	126,947
(AG) _n	1,607,242	166,884	166,312	166,296
(AT) _n	2,107,265	117,575	117,242	117,244
(CG) _n	60,759	6,427	6,378	6,381
(CT) _n	1,608,739	167,349	166,754	166,749
(GT) _n	1,291,081	128,972	128,520	128,525
(AAC) _n	68,259	3,919	3,909	3,909
(AAG) _n	86,740	7,042	7,020	7,019
(AAT) _n	167,160	9,230	9,209	9,209
(ACC) _n	114,798	32,880	32,736	32,739
(ACG) _n	592	18	70	71
(ACT) _n	16,998	1,404	1,402	1,402
(AGC) _n	62,444	7,454	7,421	7,421
(AGG) _n	84,147	7,740	7,706	7,712
(AGT) _n	16,875	1,408	1,405	1,405
(ATC) _n	53,402	3,839	3,829	3,829
(ATG) _n	52,944	3,871	3,858	3,858
(ATT) _n	166,990	9,078	9,050	9,058
(CCG) _n	9,297	413	411	410
(CCT) _n	84,257	7,743	7,702	7,705
(CGG) _n	9,424	427	426	425
(CGT) _n	591	71	71	71

(CTG) _n	63,715	7,687	7,660	7,655
(CTT) _n	86,491	7,246	7,229	7,226
(GGT) _n	114,492	32,743	32,576	32,576
(GTT) _n	68,914	3,793	3,779	3,782
(AAAC) _n	41,472	1,579	1,573	1,571
(AAAG) _n	31,680	1,096	1,093	1,093
(AAAT) _n	61,622	2,904	2,891	2,894
(AACC) _n	1,735	122	122	122
(AACG) _n	25	3	3	2
(AACT) _n	453	37	37	37
(AAGC) _n	1,444	107	107	107
(AAGG) _n	11,944	443	440	440
(AAGT) _n	776	74	74	74
(AATC) _n	2,633	246	246	246
(AATG) _n	15,190	1,347	1,345	1,345
(AATT) _n	8,704	62	62	62
(ACAG) _n	2,849	232	232	232
(ACAT) _n	6,599	155	154	154
(ACCC) _n	3,090	144	144	144
(ACCG) _n	23	2	2	2
(ACCT) _n	870	59	59	59
(ACGG) _n	70	2	2	2
(ACTC) _n	2,884	247	246	246
(ACTG) _n	945	98	98	98
(ACTT) _n	749	54	54	54
(AGAT) _n	5,583	104	104	104
(AGCC) _n	2,522	229	229	229
(AGCG) _n	186	10	10	10
(AGCT) _n	673	11	11	11
(AGGC) _n	5,237	325	323	323
(AGGG) _n	10,619	368	367	366
(AGGT) _n	901	67	66	66
(AGTC) _n	916	76	75	75
(AGTG) _n	2,841	253	249	249

(AGTT) _n	403	35	35	35
(ATCC) _n	5,940	217	216	216
(ATCT) _n	5,575	112	112	112
(ATGC) _n	2,277	21	21	21
(ATGG) _n	6,009	179	179	179
(ATGT) _n	6,755	172	171	172
(ATTC) _n	15,055	1,434	1,433	1,431
(ATTG) _n	2,708	242	242	242
(ATTT) _n	62,007	2,933	2,927	2,924
(CCCG) _n	840	18	18	18
(CCCT) _n	10,734	376	375	375
(CCGG) _n	348	6	6	6
(CCGT) _n	44	1	1	1
(CCTG) _n	5,267	341	340	341
(CCTT) _n	11,829	444	444	444
(CGCT) _n	156	10	10	10
(CGGG) _n	804	23	23	23
(CGGT) _n	17	2	2	2
(CGTT) _n	34	2	2	2
(CTGG) _n	2,311	224	223	223
(CTGT) _n	2,787	185	184	184
(CTTG) _n	1,412	120	120	120
(CTTT) _n	32,220	1,136	1,131	1,131
(GGGT) _n	3,260	170	170	170
(GGTT) _n	1,750	154	154	154
(GTTT) _n	41,692	1,533	1,529	1,528

Table S4. Non-B DNA potential (in addition to slipped-strand structures) for microsatellite sequences.

Hairpin (self-complementary)	H-DNA (poly Pur or Poly Pyr)	Z-DNA (Pur-Pyr)
(AT) _n (Ref ⁹ ; a cruciform)	(A) _n (Ref ⁹ ; also form A tract/bent)	(AC) _n (Ref ⁹)
(AAT) _n (predicted from sequence)	(C) _n (Ref ⁹)	(CG) _n (Ref ⁹)
(ACT) _n (predicted from sequence)	(G) _n (Ref ⁹ ; also form A tract/bent)	(GT) _n (Ref ⁹)
(AGC) _n (Ref ¹⁰)	(T) _n (Ref ⁹)	
(AGG) _n (Ref ¹¹)	(AG) _n (Ref ⁹)	
(AGT) _n (predicted from sequence)	(CT) _n (Ref ⁹)	
(ATC) _n (predicted from sequence)	(AAG) _n (Ref ⁹)	
(ATG) _n (predicted from sequence)	(CCT) _n (predicted from sequence)	
(ATT) _n (Ref ¹²)	(CTT) _n (Ref ⁹)	
(CCG) _n (Ref ¹⁰)		
(CGG) _n (Ref ¹⁰)		
(CTG) _n (Ref ¹⁰)		

Table S5. Measures of G-quadruplex stability and structure determined by Circular Dichroism for the ten most common G-quadruplex motifs in the genome.

G1 through G10 indicate, in the order of frequency in the genome, the ten most common G-quadruplex motif types in our annotations (G1 -- the most common, G2 the next most common, etc.). The last column reports the number of occurrences of each motif type after filtering out the ones completely lacking IPD values and the distribution of the mean IPD. Cyan indicates intra-stranded G-quadruplexes, while orange indicates inter-stranded ones. "Intra" -- intramolecular, "bimol" -- bimolecular, "paral" -- parallel structures, "anti" -- antiparallel structures.

Sequence	T _m [°C]	Molecularity	Max delta epsilon	Strand orientation	Mean IPD (5th, 25th, 50th, 75th, 95th quantiles)
G1 GGGGTGGGGGGA GGGGGGAGGG	74.3	intra	248	paral + anti	0.91 1.07 1.19 1.33 1.60 (2,962 occurrences)
G2 GGGAGGGAGGTG GGGGGG	64.8	bimol	298	paral	0.86 0.98 1.06 1.18 1.36 (540 occurrences)
G3 GGGGTCGGGGGA GGGGGGAGGG	74.8	intra	216	paral + anti	0.75 0.84 0.91 0.98 1.14 (440 occurrences)
<mark>G4</mark> GGGGTGGGGGGA GTGGGGAGGG	69.0	intra	209	paral + anti	0.74 0.83 0.90 0.99 1.13 (312 occurrences)
G5 GGGAGGGAGGGA GGGAGGG	69.0	bimol 2 types	300	paral	0.84 0.99 1.15 1.29 1.62 (287 occurrences)
G6 GGGAGGGAGGTG GGGGGGG	68.0	bimol + higher	300	paral	0.81 0.97 1.06 1.16 1.36 (148 occurrences)
G7 GGGTGGAGGGTG GGAGGAGGG	61.5	bimol 2 types	282	paral	0.83 0.92 1.00 1.08 1.28 (262 occurrences)
G8 GGGGTTGGGGGA GGGGGGAGGG	73.2	intra	211	paral + anti	0.78 0.85 0.93 1.01 1.21 (189 occurrences)
G9 GGGGTGGGGGGA GGGGGAGGG	71.9	intra	281	paral + anti	0.93 1.17 1.38 1.66 2.09 (181 occurrences)
G10 GGGGTGGGGGGA GCGGGGAGGG	68.5	intra	216	paral + anti	0.82 0.91 1.00 1.07 1.32 (177 occurrences)

Table S6. Measures of $(GGT)_n$ motif stability and structure determined by Circular Dichroism.

Cyan indicates intra-stranded structures, while orange indicates inter-stranded ones. See other abbreviations explained in the previous table.

Sequence	T _m [°C]	Molecularity	Max delta epsilon	Strand orientation
(GGT) ₄ GGTGGTGGTGGT	48.0	tetra	184	paral + anti
(GGT)₅ GGTGGTGGTGGT GGT	45.2	bimol	138	paral
(GGT)₀ GGTGGTGGTGGT GGTGGT	39.0	bimol + intra	117	paral + anti

Table S7. Sample size (the number of motifs) for computing and testing fold differences in the rates of SMRT sequencing errors.

Motifs	Sample size for sequencing errors
A-phased repeats	10,895
Direct repeats	12,423
Inverted repeats	168,191
Mirror repeats	13,185
Z-DNA motifs	2,764
G-quadruplexes on the reference (G4+)	5,938
G-quadruplexes on the reverse complement (G4-)	5,696

Table S8. STR aligning and collapsing: an example.

The five STRs shown in the table are aligned and collapsed to allow correct motif alignment, and presented as the motif $(ACTT)_n$. A capitalized nucleotide indicates the center of the STR, while bracketed nucleotides show near-central positions chosen to align the motifs.

Motif	STR	Aligned microsatellite
(ACTT) ₂	acttActt	actt[A]ctt
(CTTA) ₃	cttactTactta	cttactT[a]ctta
(TTAC) ₃	ttacttActtac	ttactt[A]cttac
(TACT) ₅	tacttacttaCttacttact	tacttactt[a]Cttacttact
(ACTT) ₄	acttacttActtactt	acttactt[A]cttactt

Table S9. Kinetics and error rates in G-quadruplexes are linked to their divergence and diversity.

Residual IPDs were obtained as differences between observed mean IPDs and the ones predicted according to mononucleotide sequence composition (see Methods). Highly diverged/diverse G-quadruplexes (top 5%) have higher IPD and SMRT mismatch error rates than the ones with low divergence/diversity (bottom 5%). Sample sizes: n=314 (in the top 5% of divergence; the same number of motifs in the bottom 5% of divergence); n=302 (in the top 5% of diversity; the same number of motifs in the bottom 5% of diversity).

	Diverge	ence	Diversity	
	Bottom 5%	Тор 5%	Bottom 5%	Тор 5%
Residuals log mean IPDs	0.1692	0.2331	0.1732	0.2040
t-test p-value	4x1	10-4	0.04	463
Log error rates	0.0226 0.0242		0.0223	0.0235
t-test p-value	0.04	404	0.0	136

SUPPLEMENTARY FIGURES

Figure S1. Window centering of motifs with an even or odd number of nucleotides.

Each box is a nucleotide. The red box/line represent the motif and window centers.

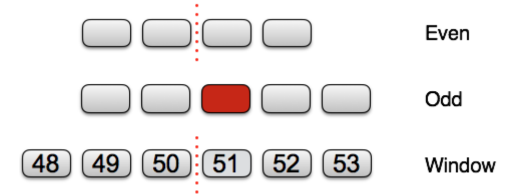
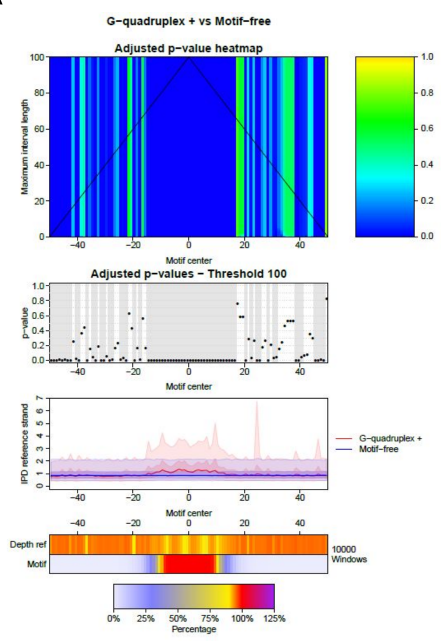


Figure S2. An example of detailed results of Interval-Wise Testing.

Results of IWT using multi-quantile statistic and a random subsample of 10,000 windows for the comparisons **A** G-quadruplex motifs on reference strand vs. motif-free windows. **B** $(AGC)_n$ vs. motif-free windows. The heatmap at the top shows the p-value curves produced by the IWT for every possible scale. The x axis indicates the positions in the 100-bp window. The y axis indicates the scale at which the test is performed, from the 1-bp scale (bottom row of the heatmap, maximum interval length=1) to the maximum possible scale of 100-bp (top row of the heatmap, maximum interval length=100). Blue corresponds to low p-values. The central plot shows the p-value curve at scale 100-bp, with gray areas highlighting significant positions (p-values≤0.05). The plot and heatmap at the bottom show the distribution of IPD values (see caption of Fig. 2A).

Α



(AGC)n vs Motif-free

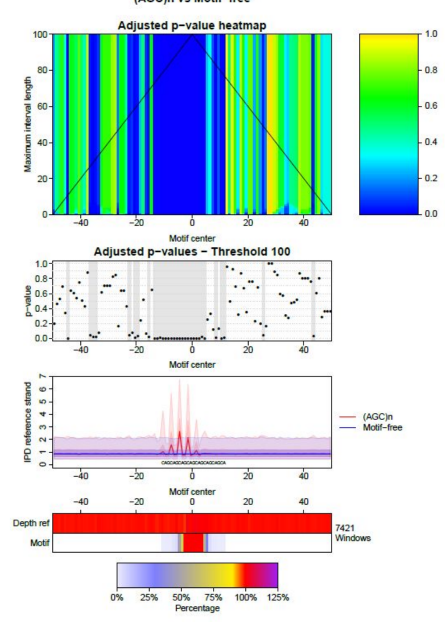
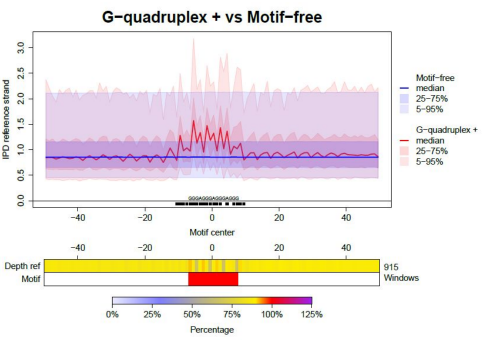
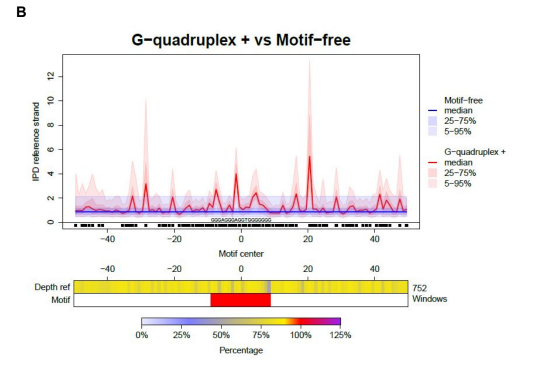


Figure S3. Different shapes of IPD curve distributions among different G-quadruplex motifs.

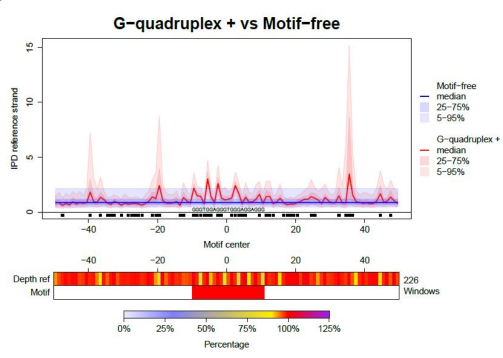
The analysis dividing G4 motifs based on their motifs was performed on the full data set of >300,000 G4 motifs, allowing overlaps between motifs of the same and different types - we do not have enough data to perform such an analysis for our non-overlapping data set of 26,000 motifs. The results still confirm elevated IPDs at G4s demonstrating that filtering for overlapping annotations does not affect our main results. **A** GGGA₃₋₅G₃ motifs only have the central elevation and lack the 3' spike. **B** GGGA₂GGT₁G₇₋₈ and **C** G₃T₁G₂A₁G₃T₁G₃A₁G₂A₁G₃ present only spikes in 5', 3' and overlapping the motif. **D** G₄TN₁G₅A₁G₆A₁G₃, **E** G₄T₁G₆A₁G₅A₁G₃, **F** G₄T₁G₆A₁₋₂G₅A₁G₃, **G** G₄T₁G₆AGN₁G₄A₁G₃, **H** G₄T₁G₆A₁G₅A₁₋₂G₃ and **I** G₄T₁G₆AT₁G₅A₁G₃ all have a central elevation surrounded by spikes as well as the 3' spike. Finally, **J** GGGT₃GGG₁ shows a series of periodic spikes, similar to the pattern observed at many microsatellites. This suggests that the last motif actually folds into a slipped structure and not into a G-quadruplex. See the legend of Fig. 2A.



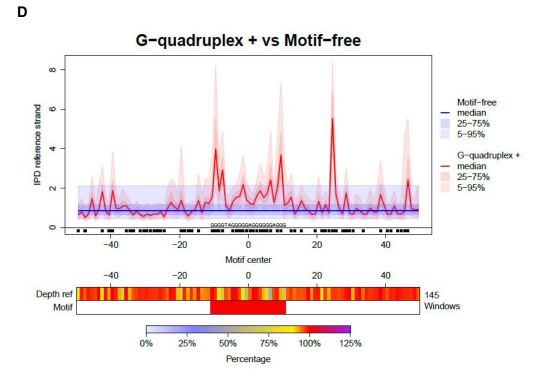




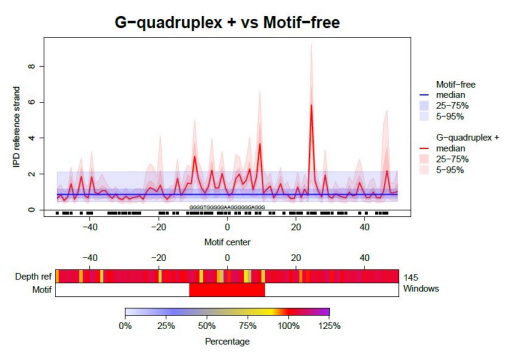
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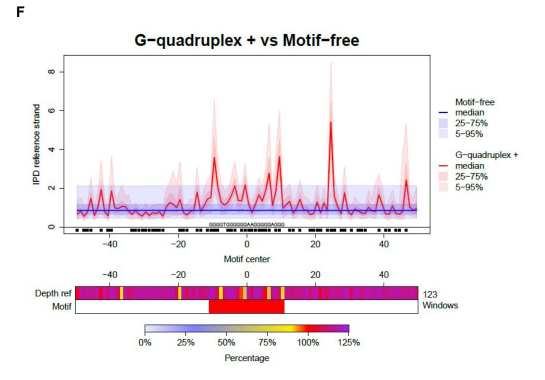


19

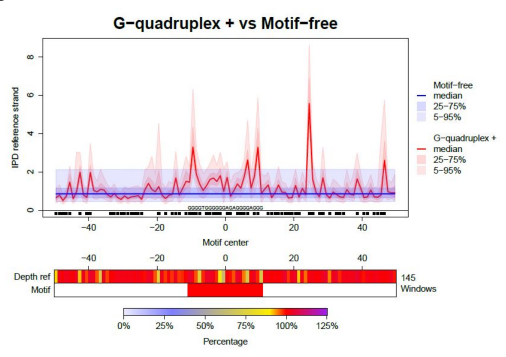


Ε

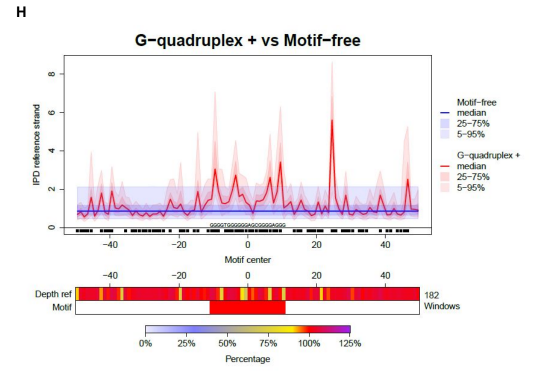




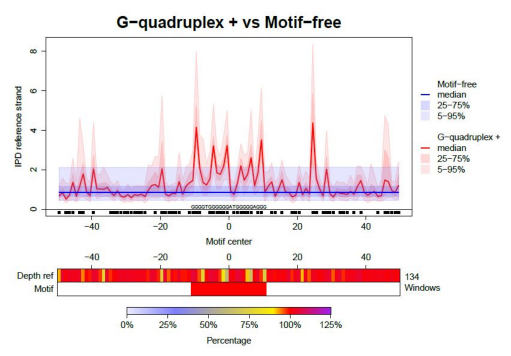
G



21



I



22

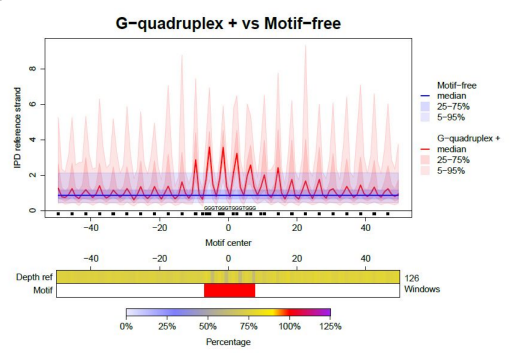
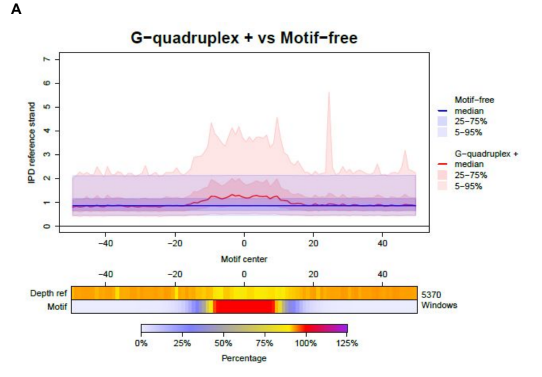


Figure S4. IPD curve distribution for G-quadruplexes identified by in vitro ion concentration manipulations.

A The IPD profile for G4+ on the reference strand (computed on 5,370 windows) is very similar to the one obtained considering all G4+ motifs (13,049 windows; see top panel of Fig. 2A). **B** The IPD profile for G4- on the reference strand (computed on 5,463 windows) is very similar to the mirror image of the one obtained considering all G4+ motifs on the reverse complement strand (13,046 windows; see bottom panel of Fig. 2A). No statistical test was performed. Additional details on various elements of these graphical representations can be found in the legend of Fig. 2A.



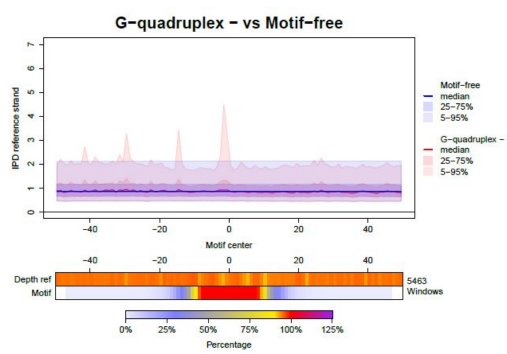
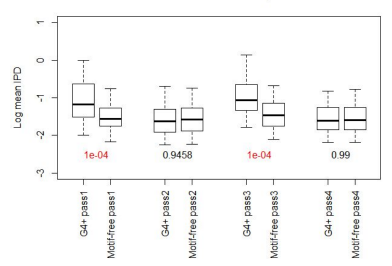


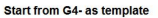
Figure S5. G-quadruplex structure is stable after multiple passes of sequencing of the circular template.

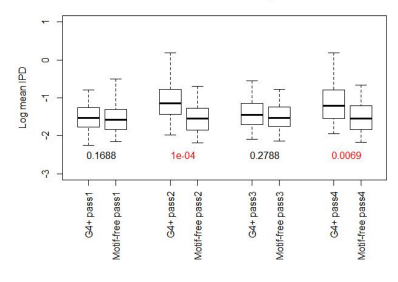
For every G4+ motif occurrence and matching motif-free region, we considered one molecule sequenced by exactly 4 passes (before polymerase drops, it uses G4+ as a template exactly twice), extracted the raw IPD information (using time between incorporation of consecutive bases in seconds) and computed the mean IPD. For each pass, we tested for differences between the mean IPD in G4+ and motif-free regions (two-sided test, multi-quantile statistic). We also tested for differences in mean IPDs between the first, and the second, the third, or the last (the 4th) pass in motif-free passes, finding no significance. **A** Molecules starting from G4+ as a template (142 molecules) versus motif-free passes. **B** Molecules starting from G4- as a template (115 molecules) versus motif-free passes. **C** Different motif-free passes. Boxplot whiskers mark the 5th and 95th quantiles. White: not significant (p-value>0.05). Red (Blue): significant with mean IPD higher (lower) in G4+ than motif-free regions. The analysis was performed on subsampled PacBio data with average depth of 12x.

Α

Start from G4+ as template







С

Motif-free regions

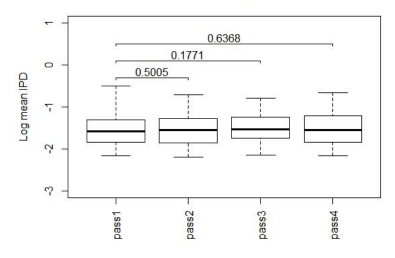
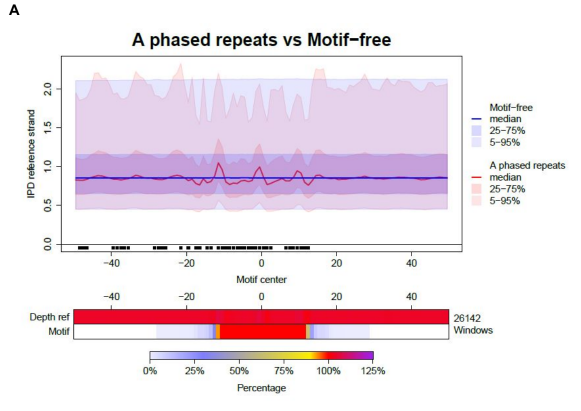


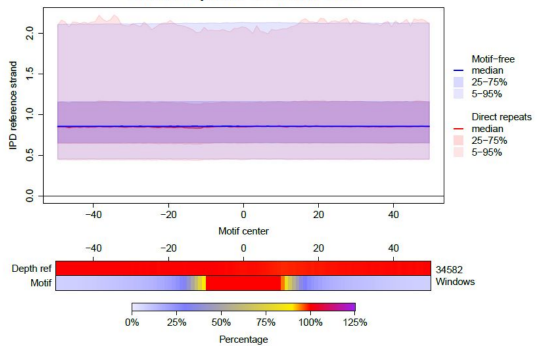
Figure S6. Effect of different non-B DNA motifs on IPDs.

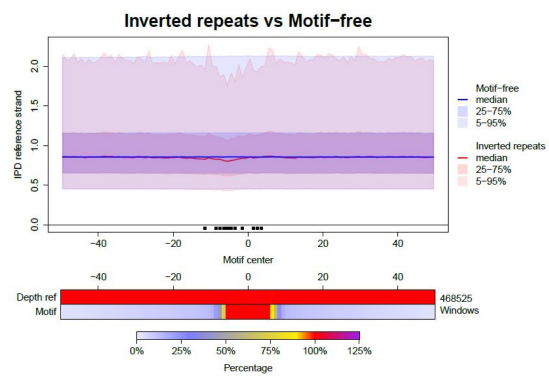
A A-phased repeats depress the IPD distribution. **B** Direct Repeats do not significantly change the IPD distribution. **C** Inverted Repeats depress the IPD distribution slightly. **D** Mirror Repeats slightly depress the IPD distribution. **E** Z-DNA motifs slightly increase the IPD distribution in both strands. See the legend of Fig. 2A for details.



В

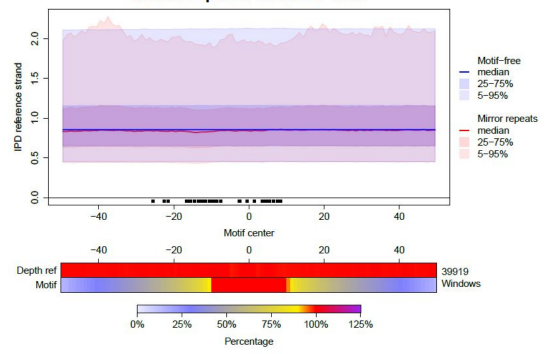
Direct repeats vs Motif-free





D

Mirror repeats vs Motif-free



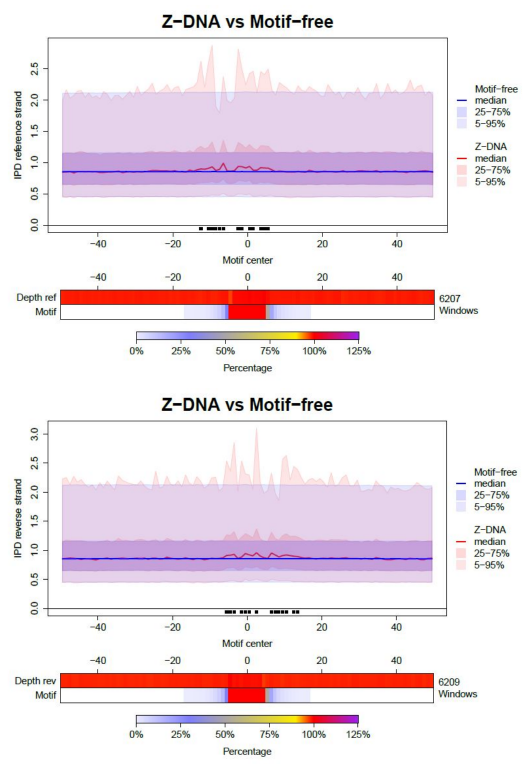
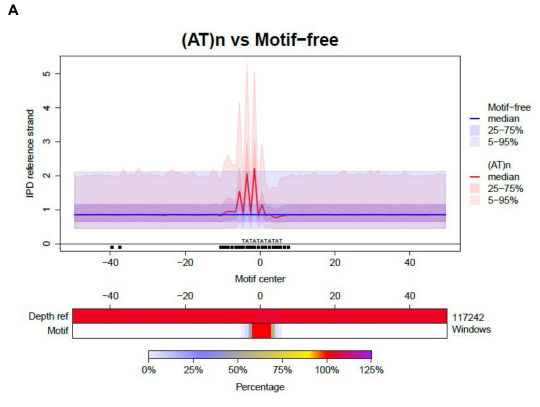


Figure S7. The effect of STRs that can form hairpins on polymerization kinetics.

 \mathbf{A} (AT)_n. \mathbf{B} (AAT)_n. \mathbf{C} (ACT)_n. \mathbf{D} (AGG)_n. \mathbf{E} (AGT)_n. \mathbf{F} (ATC)_n. \mathbf{G} (ATG)_n. \mathbf{H} (ATT)_n. See the legend of Fig. 2A.



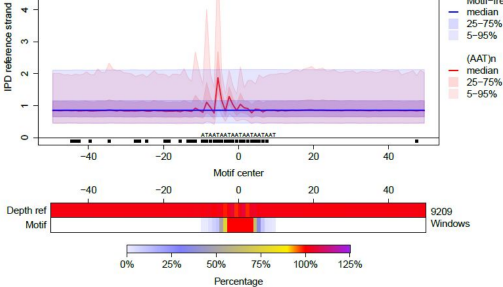
В

5

4

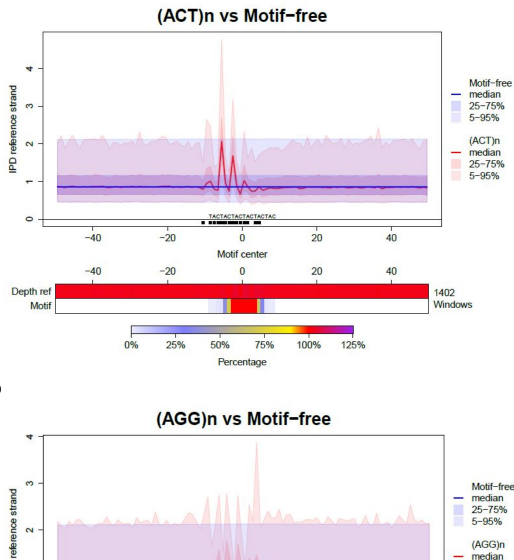


(AAT)n vs Motif-free



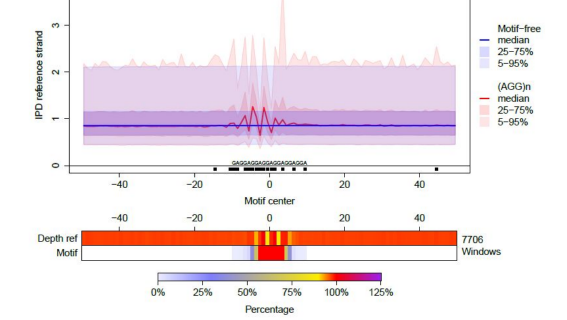
Motif-free

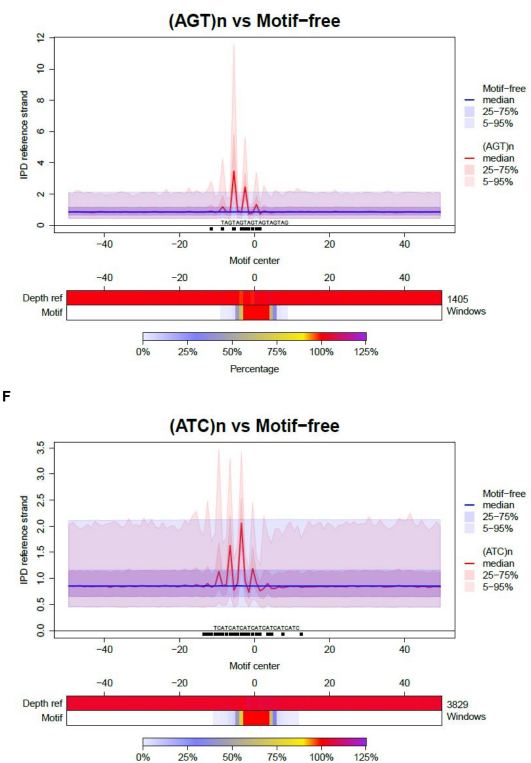
median



D

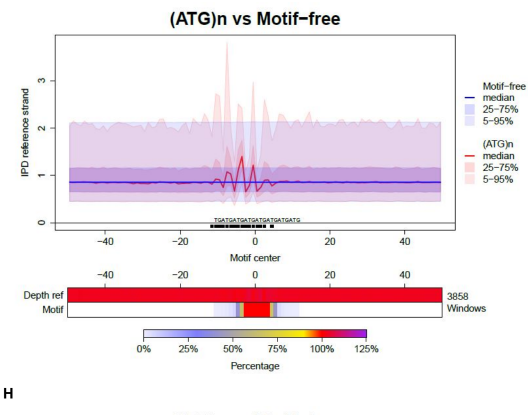
32





Percentage

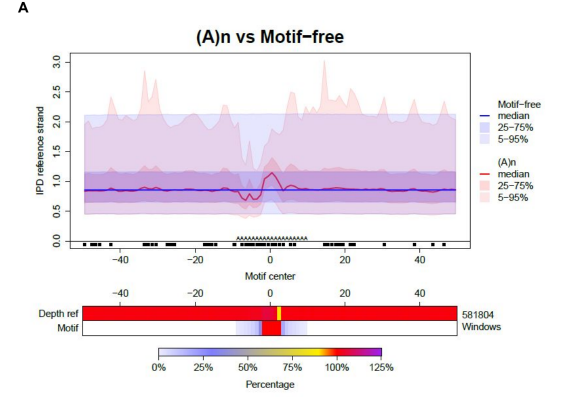
Е

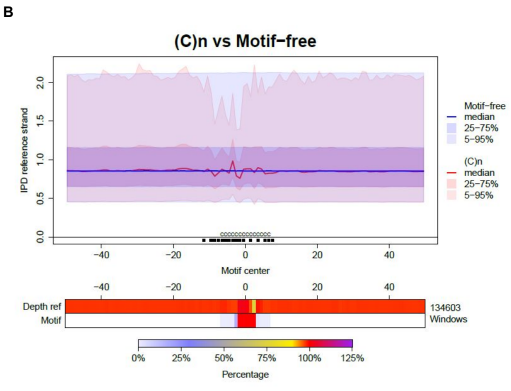


(ATT)n vs Motif-free 3 Motif-free IPD reference strand median 25-75% 5-95% 2 (ATT)n median 25-75% 5-95% _ -TTATTATTATTATTATTATT 0 t -40 -20 Ó 20 40 Motif center -40 -20 0 20 40 Depth ref 9050 Motif Windows 0% 100% 25% 50% 75% 125% Percentage

Figure S8. The effect of homopolymers and STRs that can form H-DNA on polymerization kinetics.

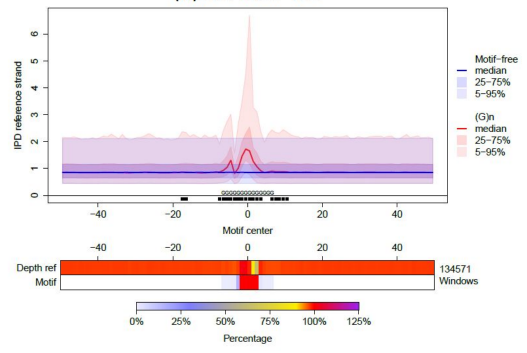
A (A)_n. **B** (C)_n. **C** (G)_n. **D** (T)_n. **E** (A)_n with different lengths. **F** (T)_n with different lengths. **G** (AG)_n. **H** (CT)_n. **I** (CCT)_n. See the legend of Fig. 2A for details about panels A-D and G-J. Panels E-F show the summary of the IWT results (see caption of Fig. 2E for details) for the comparisons of motif-containing vs. motif-free windows, with motif-containing windows grouped by the number of nucleotides in the motif (excluding lengths with fewer than 10 windows). We did not perform the analysis for (C)_n and (G)_n of different lengths because they are too short (their length ranges from 5 to 14 nucleotides, but only ~0.4% of them, 611 (C)_n and 570 (G)_n, have length >7 nt). The relationship between mean IPD in the 100-bp windows (on a logarithmic scale) and motif length was also analyzed for all non-B DNA motifs using boxplots (results not shown).



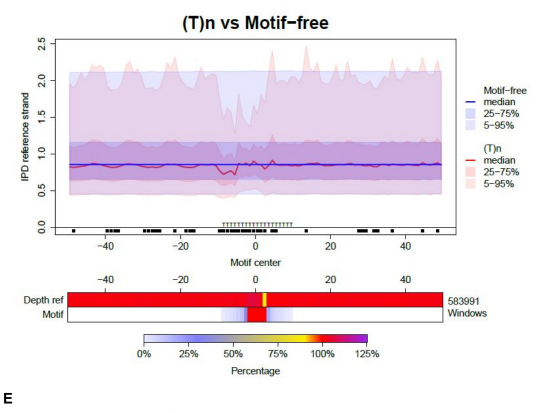


С

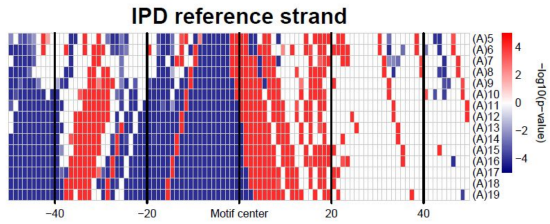
(G)n vs Motif-free

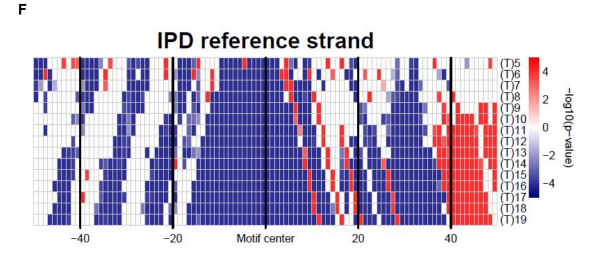


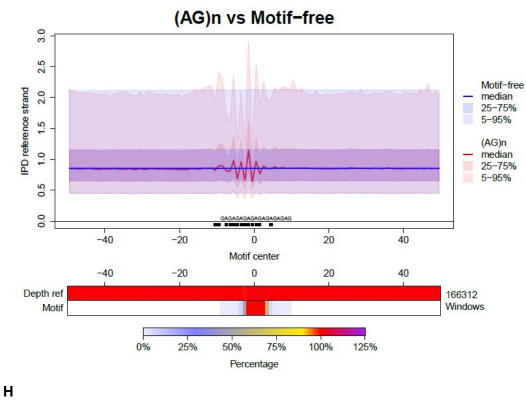
36



D

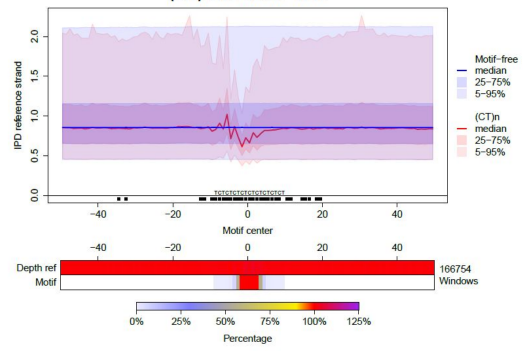






|

(CT)n vs Motif-free



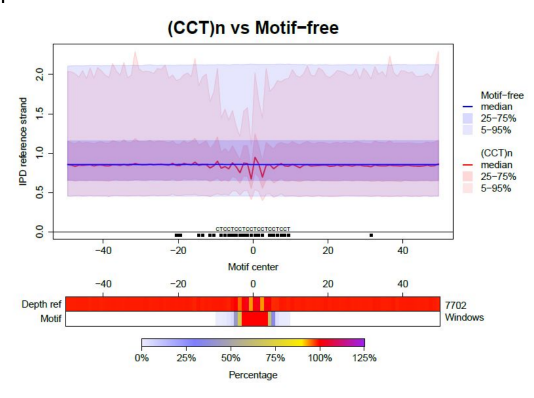
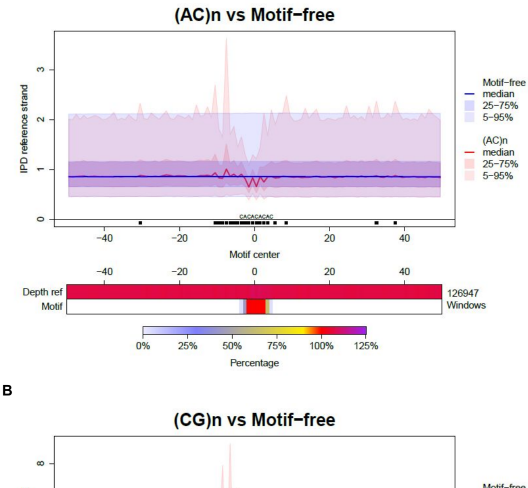
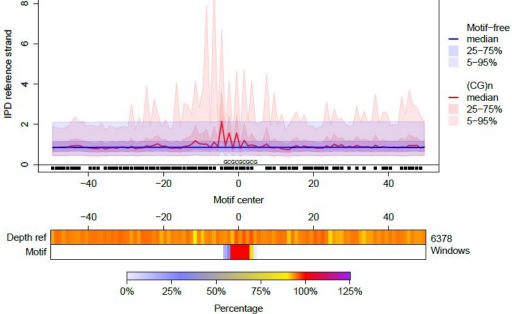


Figure S9. The effect of STRs that can form Z-DNA on polymerization kinetics. A $(AC)_n$. B $(CG)_n$. C $(GT)_n$. See the legend of Fig. 2A. A





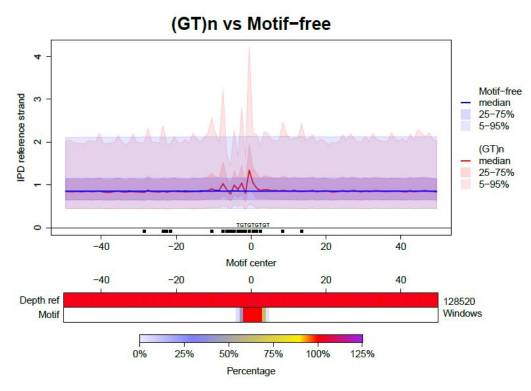
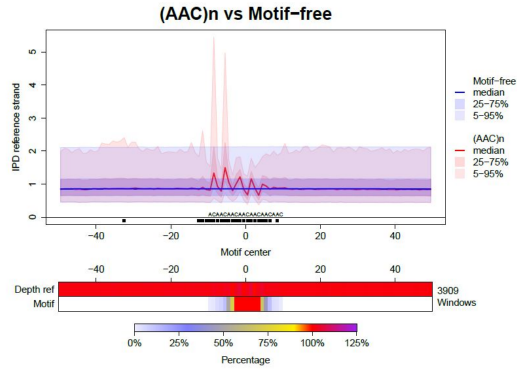
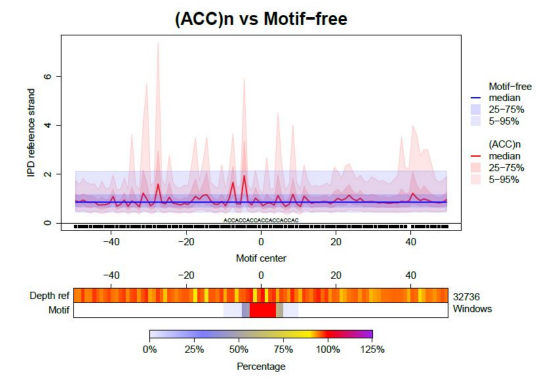


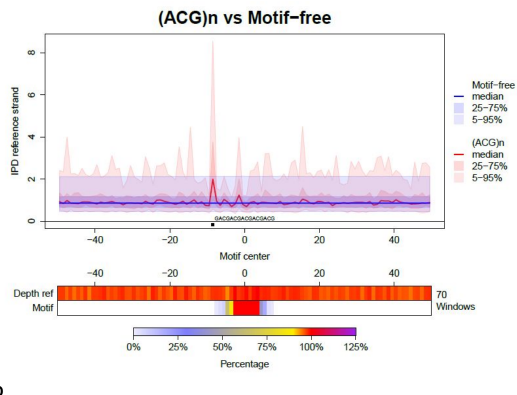
Figure S10. The effect of STRs on polymerization kinetics.

 $\boldsymbol{A} (AAC)_n. \ \boldsymbol{B} (ACC)_n. \ \boldsymbol{C} (ACG)_n. \ \boldsymbol{D} (CGT)_n. \ \boldsymbol{E} (GGT)_n. \ \boldsymbol{F} (GTT)_n. \ See the legend of Fig. 2A.$ Α



В

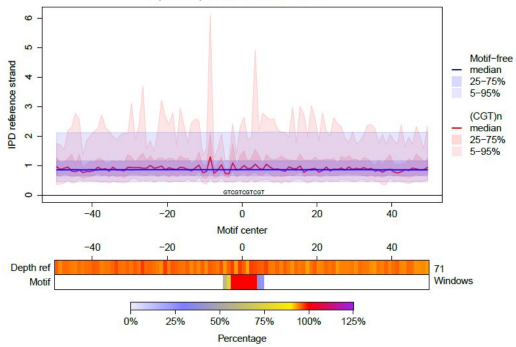


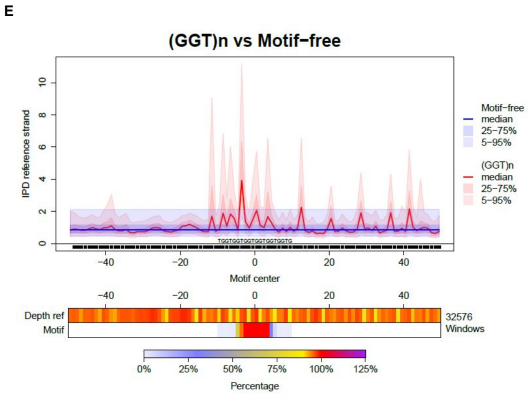


D

С

(CGT)n vs Motif-free



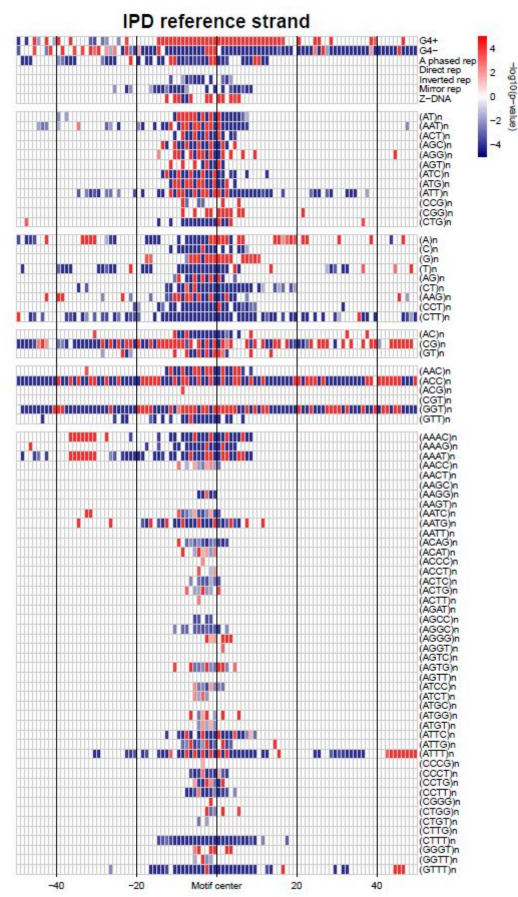


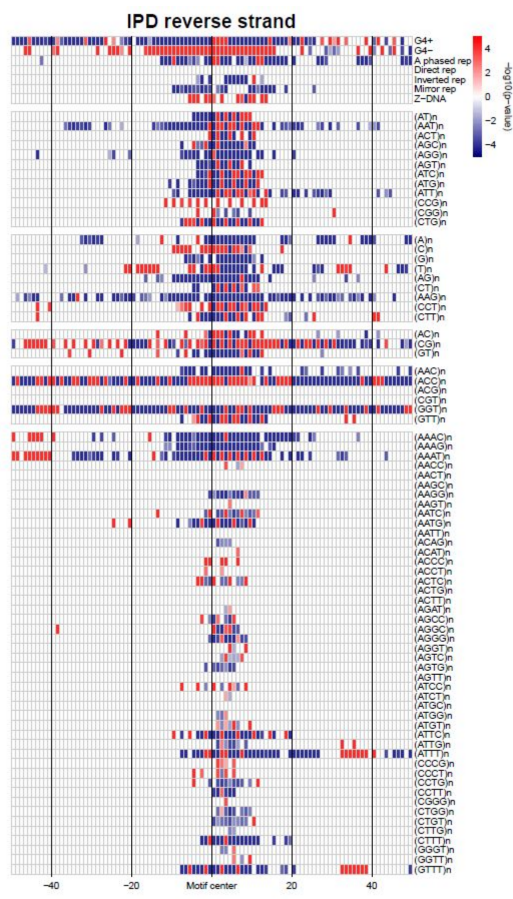
F

(GTT)n vs Motif-free 2.0 Motif-free IPD reference strand median 1.5 25-75% 5-95% 1.0 (GTT)n median 25-75% 5-95% 25-75% 0.5 0.0 TGTTGTTGTTGTTGTTGTTG -40 -20 20 40 Ó Motif center -40 -20 0 20 40 Depth ref 3779 Windows Motif 0% 75% 100% 25% 50% 125% Percentage

Figure S11. Summary of Interval-Wise Testing results for differences in IPDs.

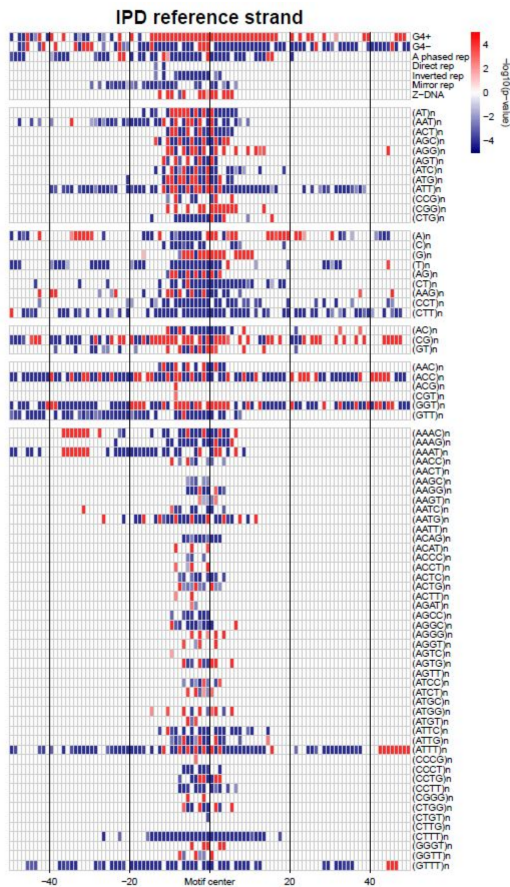
A Reference strand, multi-quantile statistic. **B** Reverse complement strand, multi-quantile statistic. **C** Reference strand, mean statistic. **D** Reverse complement strand, mean statistic. **E** Reference strand, median statistic. **F** Reverse complement strand, median statistic. See the legend of Fig. 2E for details.

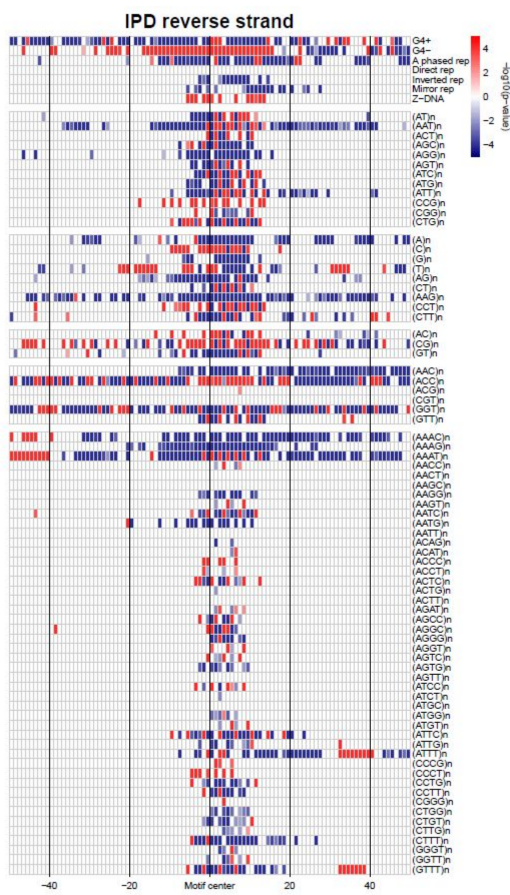


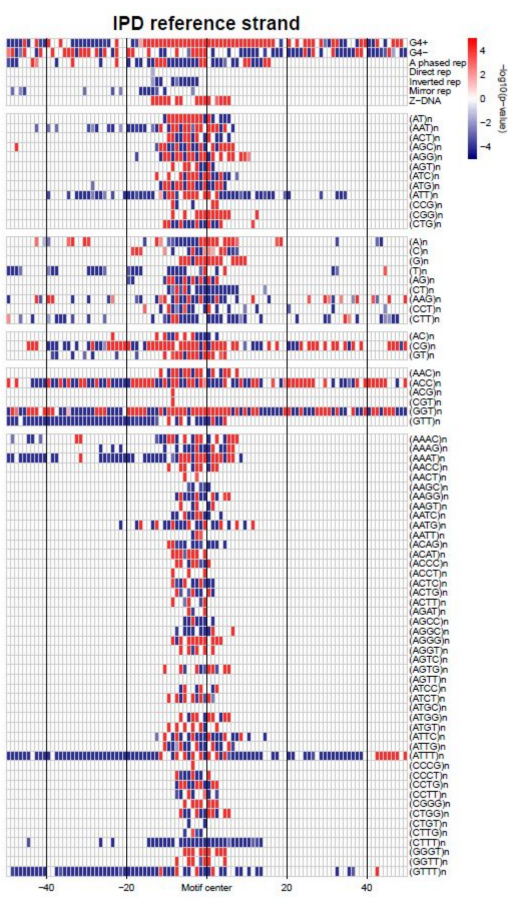




С







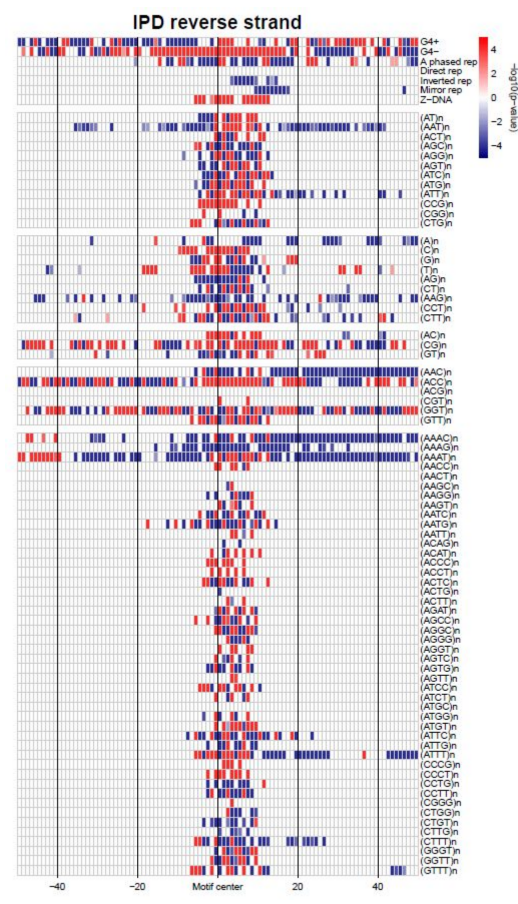
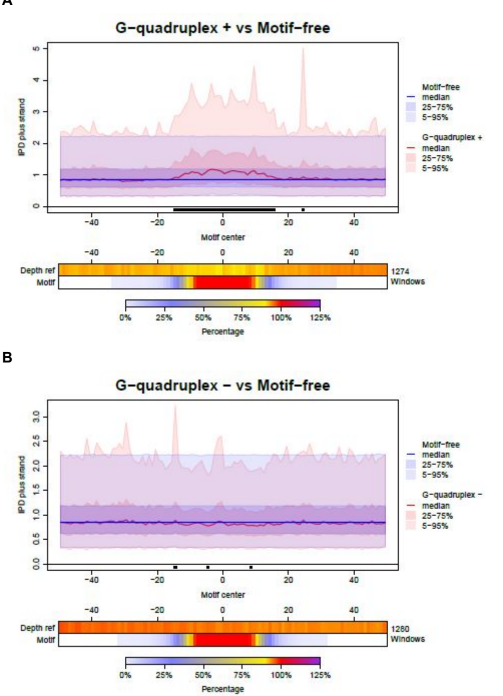
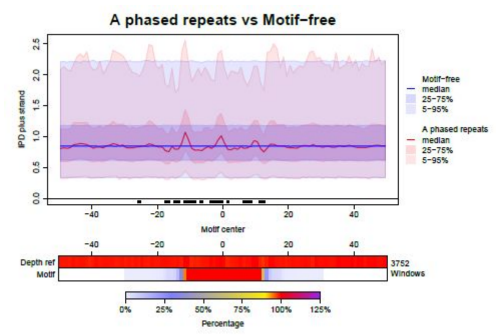


Figure S12. Variation in IPD remains in PCR-amplified sequences.

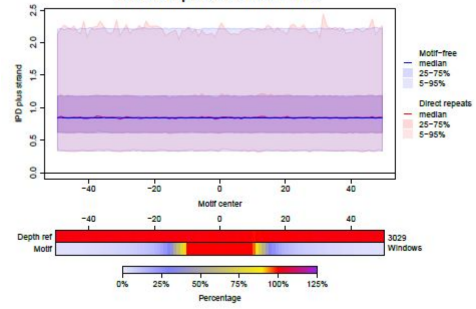
The chromosome 21 from Sumatran orangutan was flow-sorted from a cell line using a previously described protocol¹³. Subsequently, the flow-sorted material was used as a template for WGA performed with the REPLI-g Single Cell Kit (Qiagen). After de-branching¹⁴, the whole-genome amplified material was sequenced on 4 SMRT cells of the RSII instrument. Non-B DNA annotations of orangutan were obtained from the non-B DB ⁷. **A** G+ motifs. **B** G- motifs. **C** A-phased repeats. **D** Direct repeats. **E** Inverted repeats. **F** Mirror repeats. **G** Z-DNA motifs. See the legend of Fig. 2A for details.

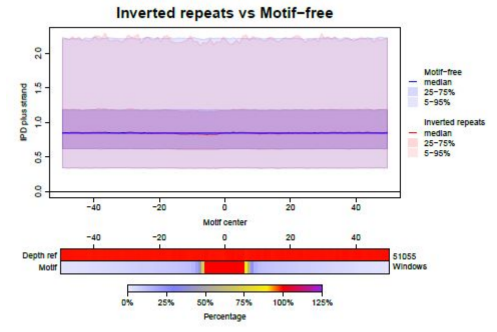




D

Direct repeats vs Motif-free

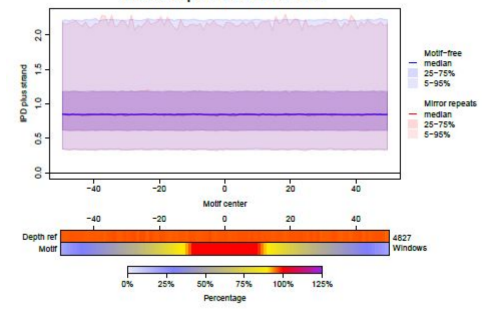






Ε

Mirror repeats vs Motif-free



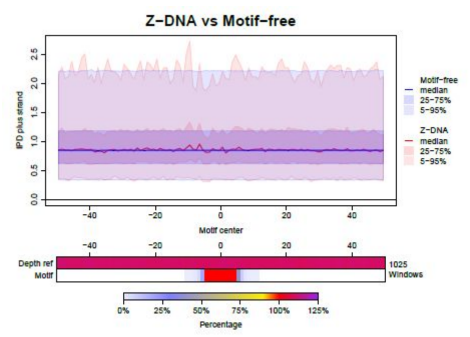
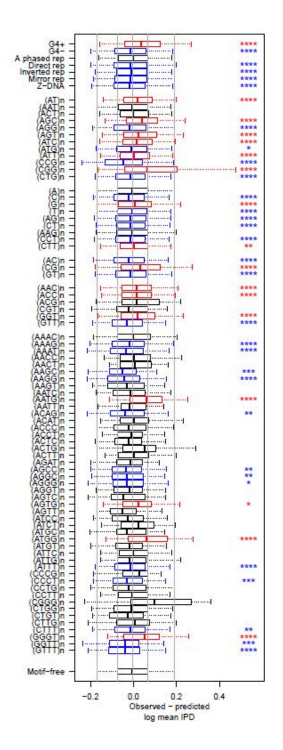


Figure S13. A comparison between observed and predicted mean IPD.

Predictions of mean IPD values in motif-containing windows are obtained from a compositional regression model fitted considering dinucleotide sequence composition on motif-free windows. **A** Reference strand. **B** Reverse complement strand. Bonferroni-corrected t-test p-values for differences: ≤ 0.0001 (*****', ≤ 0.001 (***', ≤ 0.05 (*'). Black: non-significant (corrected p-value > 0.05); red/blue: significant, with observed mean IPDs higher/lower than composition-based predictions. Boxplot whiskers: 5th and 95th quantiles of the differences.



Α

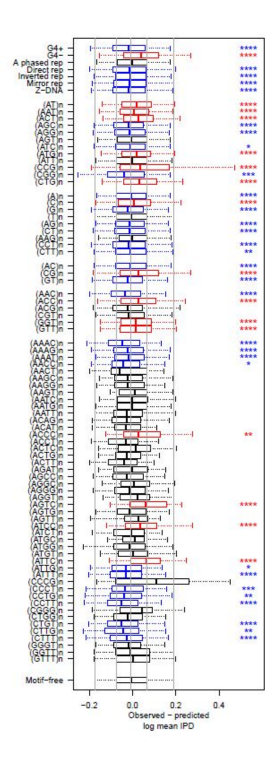


Figure S14. The relationship between IPD and sequence composition.

Plot of the mean IPD in each motif-free window in relation to sequence composition (percentage of A, T, G and C in the window). The red clouds indicate observed IPDs, while the blue clouds correspond to the compositional regression model with the mean IPD as response and the single nucleotide sequence composition as the predictor. The top right of each panel reports the correlation between the percentage of each nucleotide and the mean IPD in motif-free windows.

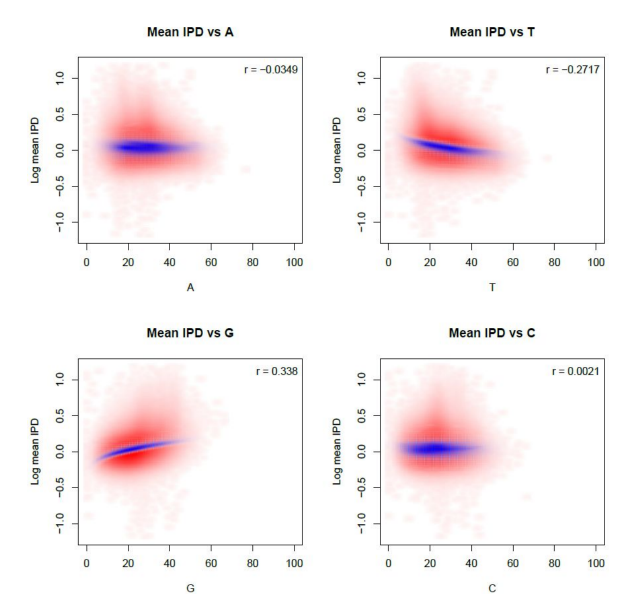
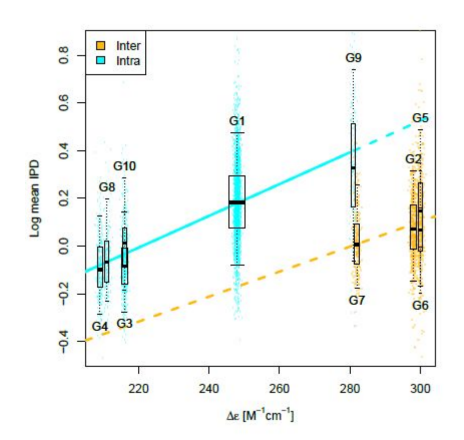


Figure S15. G-quadruplex thermostability and molecularity as predictors of polymerization kinetics.

G1 through G10 indicate, in order, the ten most common G-quadruplex motif types in our annotations (G1 the most common, G2 the next most common, etc.; Table S5). For each motif type we measured delta epsilon and T_m once, while we computed an average IPD for each occurrence of the motif in the genome, thus thousands of motifs were analyzed (Table S5). The average IPD value was then regressed against **A** circular dichroism (delta epsilon), or **B** melting temperature (T_m), considering intraand intermolecular G4s together and using molecularity (intra/inter-strandedness) as a binary predictor (dashed lines; solid lines represent the model obtained using only intramolecular G4s). R-squared 28.4% for delta epsilon (molecularity significantly changes the slope, but not the intercept, of the line), 6.7% for T_m (molecularity significantly changes both the slope and the intercept of the line). Yellow: intermolecular G-quadruplexes. Boxplot whiskers mark the 5th and 95th quantiles. **A**



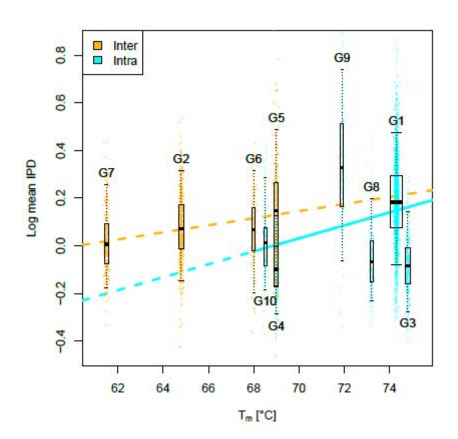
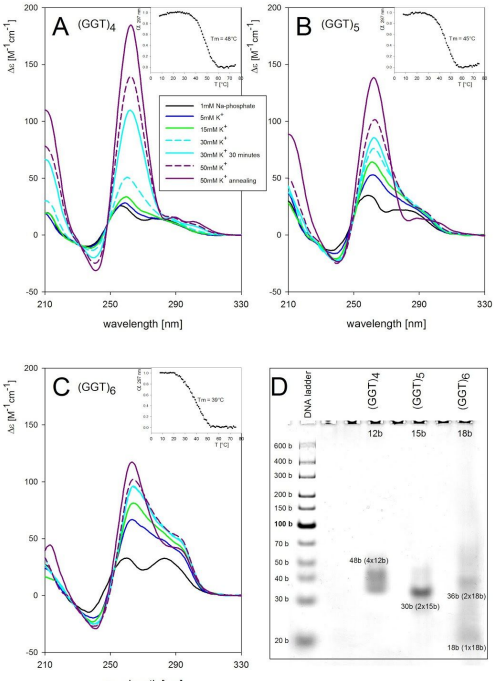


Figure S16. CD spectra, thermal denaturation and PAGE.

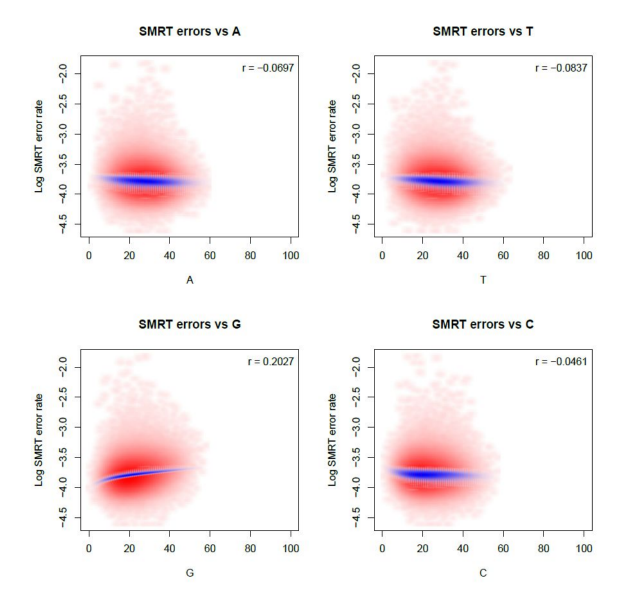
A $(GGT)_4$. **B** $(GGT)_5$. **C** $(GGT)_6$. CD spectra of all three oligonucleotides were measured at various potassium concentrations and kinetics (after 30 minutes period after K+ addition or after slow annealing). Insert figures show thermal denaturation curves and T_m . (D) Native 16% PAGE (10mM K-phosphate+35mM KCI, pH 7.0, stained by Stains All) shows tetramolecular quadruplex in (GGT)₄, bimolecular quadruplex in (GGT)₅ and bi- and monomolecular quadruplex in (GGT)₆. Samples in the PAGE were slowly annealed for 2 hours before loading onto the gel.

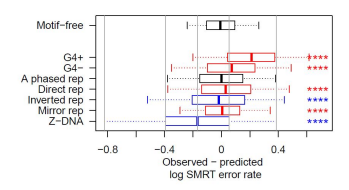


wavelength [nm]

Figure S17. The relationship between error rate and sequence composition.

A. Plot of SMRT error rate in each motif-free window in relation to sequence composition (percentage of A, T, G and C in the window). The red clouds indicate observed SMRT mismatch rates, while the blue clouds correspond to the compositional regression model with SMRT mismatch rate as response and the single nucleotide sequence composition as the predictor. The top right of each panel reports the correlation between the percentage of each nucleotide and SMRT error rate in motif-free windows. **B** Comparison between SMRT mismatch rate observed, and the one predicted using a compositional regression model fitted considering mononucleotide sequence composition on motif-free windows. Bonferroni-corrected t-test p-values for differences: ≤ 0.0001 (*****', ≤ 0.001 (***', ≤ 0.01 (***', ≤ 0.05 (*'. Black: non-significant (corrected p-value > 0.05); red/blue: significant, with observed error rates higher/lower than composition-based predictions. Boxplot whiskers: 5th and 95th quantiles of the differences. **A**.





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 ATT
 Trinucleotide Repeats Adopts Stable Hairpin Structures Induced by Minor Groove
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