

A small targeting domain in Ty1 integrase is sufficient to direct retrotransposon integration upstream of tRNA genes

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Abstract

Integration of transposable elements into the genome is mutagenic. Mechanisms targeting integrations into relatively safe locations, hence minimizing deleterious consequences for cell fitness, have emerged during evolution. In budding yeast, integration of the Ty1 LTR retrotransposon upstream of RNA polymerase III (Pol III)-transcribed genes requires interaction between Ty1 integrase (IN1) and AC40, a subunit common to Pol I and Pol III. Here, we identify the Ty1 targeting domain of IN1 that ensures (i) IN1 binding to Pol I and Pol III through AC40, (ii) IN1 genome-wide recruitment to Pol I- and Pol III-transcribed genes, and (iii) Ty1 integration only at Pol III-transcribed genes, while IN1 recruitment by AC40 is insufficient to target Ty1 integration into Pol I-transcribed genes. Swapping the targeting domains between Ty5 and Ty1 integrases causes Ty5 integration at Pol III-transcribed genes, indicating that the targeting domain of IN1 alone confers Ty1 integration site specificity.

Keywords genome-wide *de novo* insertion sites; integrase genome-wide occupancy; RNA Polymerases I and III; Ty1 retrotransposon

Subject Categories Chromatin, Transcription & Genomics; DNA Replication, Recombination & Repair

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Introduction

Transposable elements (TEs) are mobile repetitive DNA sequences found in the genomes of most organisms (Huang *et al.*, 2012). TEs are mutagenic and represent a threat to genome integrity, inactivating, or altering host gene expression or inducing large chromosomal rearrangements (Bourque, 2009; Levin & Moran, 2011). In humans, more than hundred heritable diseases have been assigned to *de novo*

TE insertions (Hancks & Kazazian, 2016). TEs also play a role in genome evolution by modifying host functions, phenotypes, and gene regulation and can contribute to the long-term adaptation of organisms to different environments (Chuong *et al.*, 2016).

Where TEs integrate in the genome will determine their impact on their host. TE distribution, which is rarely random (Sultana *et al.*, 2017), arises from the balance between two processes. First, selection leads to the elimination of strongly deleterious insertions and the maintenance of beneficial ones (Chuong *et al.*, 2016; Cosby *et al.*, 2019). Second, TEs have repeatedly evolved mechanisms that direct their integration into “safe” locations, where insertions will have minimal adverse effect on the organism’s fitness (Boeke & Devine, 1998; Spaller *et al.*, 2016; Cheung *et al.*, 2018). These regions often consist of non-essential repeated sequences, such as telomeric regions, ribosomal DNA arrays, upstream and downstream regions of transfer RNA genes (*tDNAs*), or non-essential regions upstream of open reading frames (Zou *et al.*, 1996; Penton & Crease, 2004; Fujiwara *et al.*, 2005; Ye *et al.*, 2005; Naito *et al.*, 2009; Guo & Levin, 2010; Pardue & DeBaryshe, 2011; Baller *et al.*, 2012; Mularoni *et al.*, 2012; Spaller *et al.*, 2016; Kling *et al.*, 2018). Preferential targets have been described for the integration of different classes of TEs, including retroelements (Sultana *et al.*, 2017).

Long terminal repeat (LTR) retrotransposons are retroelements related to retroviruses. They replicate by reverse transcription of their mRNA into a double-stranded DNA copy (cDNA), which is imported into the nucleus and integrated into the genome by the LTR retrotransposon integrase (IN). The interaction between IN and cellular tethering factors plays a major role in integration site selection by targeting the integrase/cDNA complexes, aka the intasome, to specific chromosome locations. Tethering factors were first identified for Ty3 and Ty5 in *Saccharomyces cerevisiae*. These LTR retrotransposons integrate at the transcription start site of Pol III-transcribed genes and in subtelomeric regions, respectively (Kirchner *et al.*, 1995; Xie *et al.*, 2001). The Tf1 LTR retrotransposon of *Schyzosaccharomyces pombe* and the MLV retrovirus both target the promoter region of Pol II-transcribed genes (De Rijck *et al.*, 2013;

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Gupta *et al*, 2013; Sharma *et al*, 2013; Hickey *et al*, 2015; Jacobs *et al*, 2015), and the HIV-1 retrovirus targets the gene body of Pol II-transcribed genes (Cherepanov *et al*, 2003; Llano *et al*, 2006). In all cases, tethering factors bind chromatin or have functions related to DNA transcription or replication (Sultana *et al*, 2017).

Ty1, the most active and abundant LTR retrotransposon in *S. cerevisiae* laboratory strain S288C, integrates preferentially within a 1 kb window upstream of Pol III-transcribed genes. It targets nucleosomal DNA near the H2A/H2B interface (Baller *et al*, 2012; Mularoni *et al*, 2012). This integration pattern allows Ty1 to replicate while minimizing disruption to the host genome, as most Pol III-transcribed genes are multicopy *tDNAs* and thus individually non-essential. Furthermore, Ty1 insertion has a limiting impact on *tDNA* expression (Bolton & Boeke, 2003). Targeted integration proximal to *tDNAs* is a strategy that has been adopted several times by TEs to minimize damage to compact genomes (Cheung *et al*, 2018; Kling *et al*, 2018). The integration of Ty1 in these regions requires a functional Pol III promoter in the target gene (Devine & Boeke, 1996) and is influenced by the chromatin-remodeling factor Isw2 and the Bdp1 subunit of TFIIB (Bachman *et al*, 2005).

Recently, we have shown that an interaction between Ty1 IN (IN1) and the AC40 subunit of Pol I and Pol III is a major driver for Ty1 integration upstream of Pol III-transcribed genes (Bridier-Nahmias *et al*, 2015). The study used the *S. pombe* AC40 ortholog (AC40sp) as a loss-of-interaction mutant. The replacement of AC40 by AC40sp severely compromised Ty1 integration upstream of Pol III-transcribed genes, leading to a redistribution of Ty1 insertions in the genome. IN1 binding to other Pol III subunits was also described *in vitro* (Cheung *et al*, 2016). However, it is not clear whether these interactions participate in Ty1 integration site selection.

IN1 has a three-domain organization common to all retroelement integrases: the Zn²⁺ coordinating N-terminal domain (NTD), the catalytic core domain (CCD), and the less conserved C-terminal domain (CTD) (Fig 1A, top) (Wilhelm *et al*, 2005). IN1 C-terminal residues 578–635 are necessary and sufficient to mediate the interaction with AC40 *in vivo* (Bridier-Nahmias *et al*, 2015). This region also contains a bipartite nuclear localization signal (bNLS, residues 596–630; Fig 1A, top) consisting of two Lys-Lys-Arg motifs separated by a 29 amino acid linker (Kenna *et al*, 1998; Moore *et al*, 1998; Lange *et al*, 2011). This raises the question of whether IN1 nuclear import and interaction with AC40 could act in concert during Ty1 replication.

In this study, we identify a short sequence, namely the Ty1 targeting domain, in the bNLS linker of IN1 that directs the interaction with AC40. Single amino acid substitutions in the Ty1 targeting domain do not affect the frequency of Ty1 retrotransposition but impair the recruitment of IN1 to Pol III-transcribed genes. Consequently, these IN1 mutations induce the same changes in the Ty1 integration profile as observed in the AC40sp loss-of-interaction mutant. While IN1 is also recruited to Pol I-transcribed genes through its interaction with AC40, Pol I-transcribed genes are poor targets of Ty1 integration. When the Ty1 targeting domain is used to replace the Ty5 IN sequence responsible for Ty5 integration into heterochromatin, Ty5 integration is redirected to Pol III-transcribed genes. This work therefore confirms the fundamental role of the IN1-AC40 interaction in Ty1 integration site selection and reveals that the targeting domain of Ty1 is necessary and sufficient to confer Ty1 integration preference to another retrotransposon.

Results

Mutations in the IN1 bNLS linker sequence abolish the interaction with AC40

Saccharomyces cerevisiae LTR retrotransposons Ty1, Ty2, and Ty4 have the same integration preferences for regions upstream of Pol III-transcribed genes (Kim *et al*, 1998; Carr *et al*, 2012), and the C-termini of their integrases (IN1, IN2, and IN4, respectively) interact with the Pol III subunit AC40 (Bridier-Nahmias *et al*, 2015). To identify conserved amino acids potentially involved in the AC40 interaction, we aligned the C-terminal sequences of IN1, IN2, and IN4 (Fig 1A, bottom) and observed that IN1 and IN2 are highly similar in this region, whereas IN4 is more divergent. Amino acids at positions 600–601, 609–610, and 617–623 in IN1 were either identical or highly similar in all three INs. We replaced each of these amino acids by alanine, individually or in pairs, in a Gal4-activating domain GAD-IN1_{578–635} fusion protein, checked that protein level was not affected by these mutations (Fig EV1A), and studied the interaction of the mutant fusion proteins with Gal4 binding domain GBD-AC40 using a two-hybrid assay. The interaction between IN1_{578–635} and AC40 was maintained in the presence of mutations LE_{600–601}AA, VS_{609–610}AA, N₆₁₈A, or E₆₂₃A and suppressed by single alanine substitution of K₆₁₇, M₆₁₉, R₆₂₀, S₆₂₁, or L₆₂₂ (Figs 1B and EV1B). GBD-AC40 interacted with GAD-IN1_{578–635} but not with GAD-IN1_{1–578} (Fig 1C) as shown previously (Bridier-Nahmias *et al*, 2015). Since amino acids K₆₁₇-L₆₂₂ are located in the IN1 bNLS linker sequence, we also tested the interaction between GBD-AC40 and GAD fused to the entire bNLS sequence (GAD-IN_{596–630}). Interaction between the two fusion proteins was detected, suggesting that this region of 34 amino acids in IN1 is sufficient for interaction with AC40 (Figs 1C and EV1C and D).

To determine whether amino acids required for the two-hybrid interaction between AC40 and the IN1 C-terminus were also critical for the interaction between the two full-length proteins, we co-expressed AC40-Strep and WT or mutant IN1-C-tag tagged proteins in *E. coli* and performed co-precipitation of the two proteins with the cell extracts. We focused on K₆₁₇, S₆₂₁, and L₆₂₂, given their strict conservation in IN1, IN2, and IN4 (Fig 1A, bottom). In the presence of Strep-Tactin beads, AC40-Strep co-immunoprecipitated with WT IN1 but not with the IN1 K₆₁₇A, S₆₂₁A, and L₆₂₂A mutants (Fig 1D). These data strongly suggest that full-length IN1 and AC40 proteins bind directly to each other, and their interaction depends on residues K₆₁₇, S₆₂₁, and L₆₂₂ located in the IN1 bNLS.

Non-AC40 binding IN1 mutants do not affect Ty1 integration frequency

Mutations in the IN1 bNLS that induce a substantial or complete loss of IN1 nuclear accumulation reduce the frequency of Ty1 retro-mobility, as seen when the two Lys-Lys-Arg (KKR) motifs are mutated, either individually or simultaneously, but also for mutations of specific acidic residues in the linker region (Kenna *et al*, 1998; Moore *et al*, 1998; Lange *et al*, 2011). To investigate whether the amino acids we identified in the bNLS linker sequence as being necessary for the AC40 interaction were also required for Ty1 nuclear import and retromobility, we introduced K₆₁₇A, S₆₂₁A, or

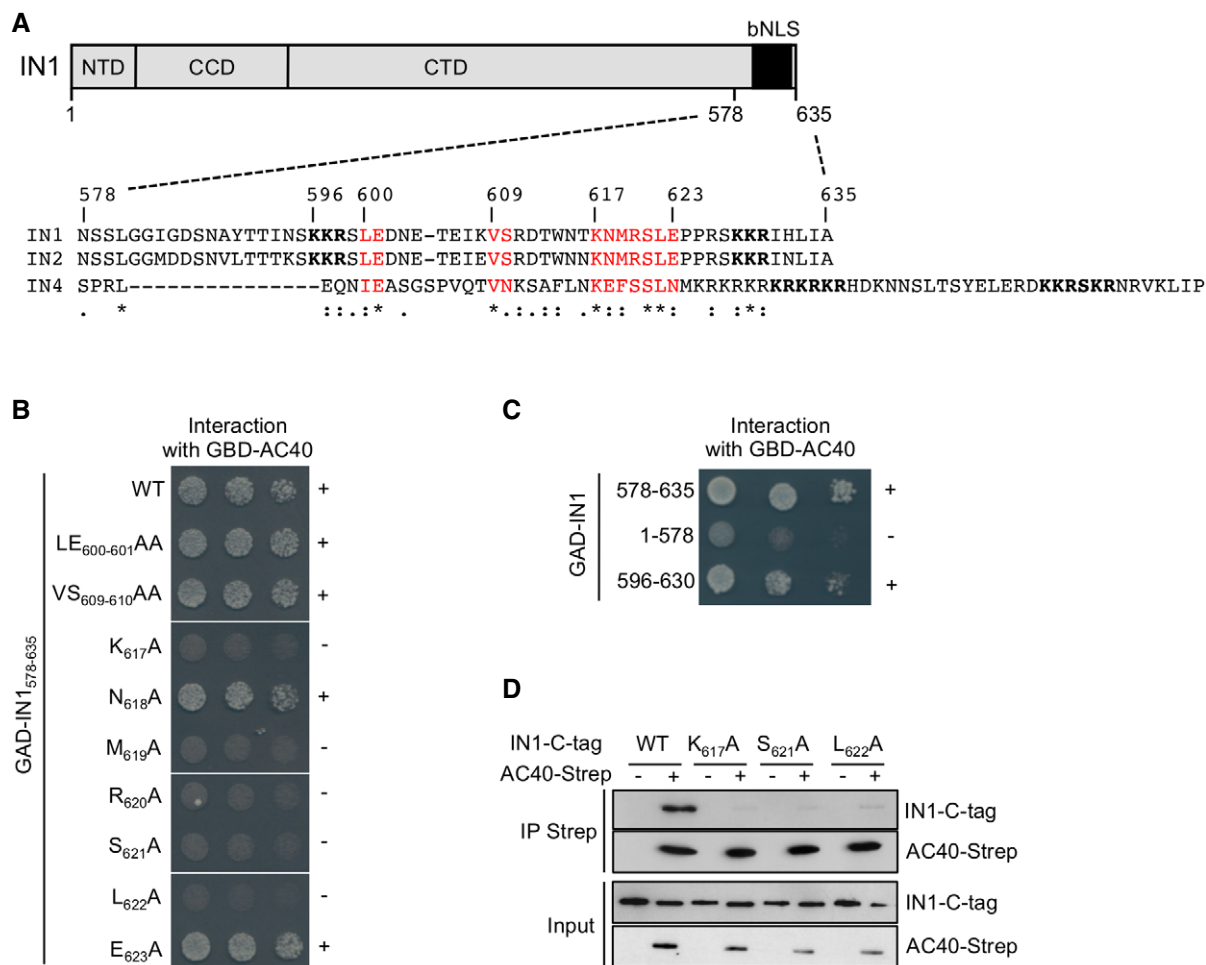


Figure 1. Mutations in the IN1 bNLS linker sequence abolish the interaction with AC40.

A *Top*. The Ty1 integrase (IN1) showing N-terminal and catalytic core domains (NTD and CCD) and the bipartite NLS at the C-terminus (CTD). *Bottom*. Alignment of amino acid sequences of Ty1, Ty2, and Ty4 integrase C-termini (IN1, IN2, and IN4, respectively). In bold: Basic amino acids required for NLS function. In red: Amino acids in the bNLS linker relatively conserved between the three integrases. *, identity; :, high similarity; ., low similarity; -, gap in sequence.

B Two-hybrid interaction between GBD-AC40 and WT or mutant GAD-IN1₅₇₈₋₆₃₅. Alanine substitutions in IN1₅₇₈₋₆₃₅ are indicated. Cells were plated in twofold serial dilutions on DO-Leu-Trp-His plates. No growth or protein expression defects were detected (Fig EV1A and B). +, interaction; -, no interaction.

C Two-hybrid interaction between GBD-AC40 and different IN1 regions fused to GAD, as indicated. Cells were plated in ten-fold serial dilutions on DO-Leu-Trp-His plates. No growth or protein expression defects were detected (Fig EV1C and D). +, interaction; -, no interaction.

D *In vitro* interaction between AC40 and IN1 proteins co-expressed in *E. coli*. Co-precipitation of WT or mutated (K₆₁₇A, S₆₂₁A, or L₆₂₂A) IN1-C-tag using AC40-Twin-Strep-tag as bait from bacterial protein extracts co-expressing (+) or not (-) these proteins. Expected sizes are 41 kDa for AC40-Twin-Strep-tag and 100 kDa for IN1-C-tag (WT and mutants).

Source data are available online for this figure.

L₆₂₂A mutations into a GFP₂-bNLS fusion protein previously used to assess IN1 bNLS function (McLane *et al*, 2008). In contrast to the IN1-bNLSmut construct (S₅₉₆KKR₅₉₈-AAA and S₆₂₈KKR₆₃₀-AAA), the three IN1-bNLS single mutants were still able to target GFP₂ into the nucleus (Fig 2A). Thus, amino acids required for the IN1-AC40 interaction are dispensable for NLS function.

The same mutations were introduced individually into a Ty1 element expressed from the *GAL1* promoter in a 2-micron plasmid and containing the retromobility indicator gene *his3AI*, allowing detection of Ty1-*HIS3* insertion events as His⁺ prototroph cells (Curcio & Garfinkel, 1991). To determine the frequency of Ty1*his3AI* integration in the genome, we expressed

this plasmid in a *spt3-101* null *rad52Δ* mutant strain, deficient in endogenous Ty1 expression and homologous recombination. *SPT3* is required for Ty1 transcription, and its absence prevents the trans-complementation of the mutant IN1 by WT IN1 from endogenous Ty1 elements (Curcio & Garfinkel, 1992). *RAD52* deletion precludes insertion of the Ty1-*HIS3* cDNA by homologous recombination with preexisting genomic Ty1 copies, a preferred pathway when IN1-dependent integration is defective (Sharon *et al*, 1994). The frequency of His⁺ cells was similar between strains expressing WT or K₆₁₇A, M₆₁₉A, R₆₂₀A, S₆₂₁A, or L₆₂₂A mutant Ty1*his3AI* (Fig 2B). In contrast, mutations that inactivate IN1 nuclear import (Moore *et al*, 1998) or catalytic

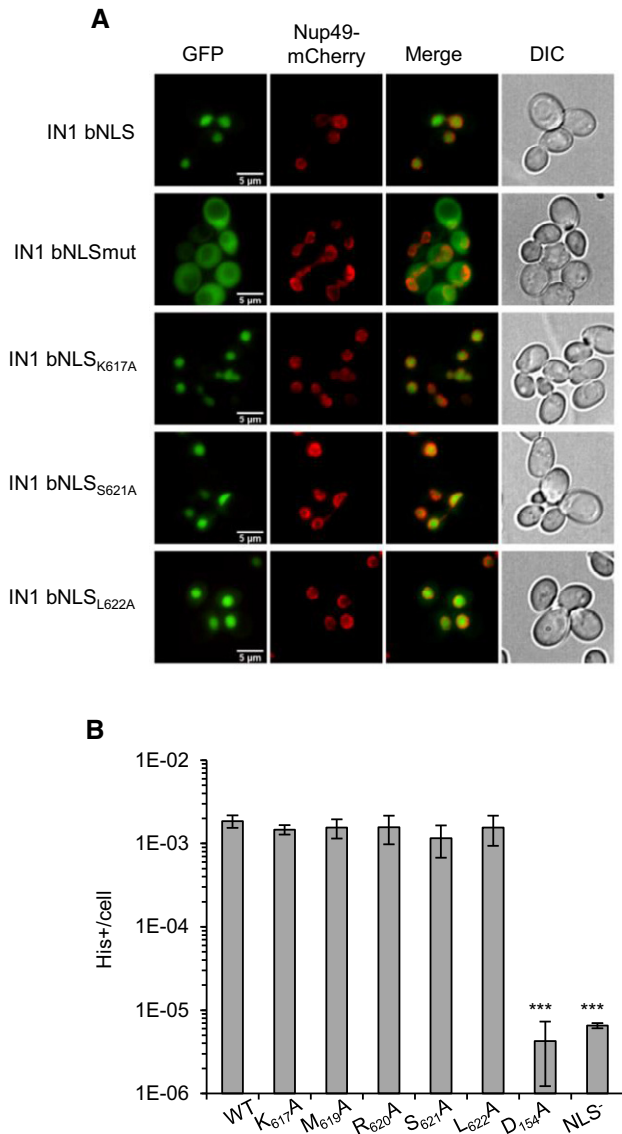


Figure 2. Non-AC40 binding IN1 mutants do not affect Ty1 integration frequency.

A Localization of GFP₂-IN1 bNLS variants in yeast cells by direct fluorescence microscopy. Nup49-mCherry signal was used to visualize the location of the nuclear envelope. Corresponding DIC images are shown. Scale bar is 5 μ m.

B Retrotransposition frequency (log scale) of pGAL1-Ty1his3AI bearing the indicated substitutions of conserved residues in a *spt3-101 rad52 Δ* strain. IN1 catalytic core domain mutant D₁₅₄A is defective for integration and NLS mutant (KKR₆₂₈₋₆₃₀GGT) for nuclear import. Values are mean \pm SD, *n* = 4 experiments, each performed with four independent colonies. ****P* \leq 0.001, otherwise not significant (Student *t*-test).

Source data are available online for this figure.

activity (Wilhelm & Wilhelm, 2005) caused a substantial decrease in the frequency of His⁺ cells compared to WT (Fig 2B, mutants NLS⁻ and D₁₅₄A, respectively).

Thus, single amino acid mutations in IN1 that strongly affect the interaction with AC40 do not impair Ty1 retrotransposition. These

data confirm that the IN1-AC40 interaction is not required for Ty1 overall integration frequency (Bridier-Nahmias *et al*, 2015).

AC40 recruits IN1 at Pol I and Pol III-transcribed genes

In addition to AC40, other Pol III subunits have been suggested to mediate the interaction between IN1 and Pol III (Cheung *et al*, 2016). Pol III is a stable complex, and immunoprecipitation of a specific subunit results in the co-precipitation of the whole complex (Oficjalska-Pham *et al*, 2006; Bhalla *et al*, 2019). To determine whether IN1 remains associated with Pol III in the absence of the IN1-AC40 interaction, we immunoprecipitated Pol III in yeast cells expressing hemagglutinin (HA)-tagged C160, the largest Pol III subunit, and WT or mutant IN1 fused to streptavidin (IN1-Strep). WT IN1 was associated with Pol III but not the K₆₁₇A, S₆₂₁A, or L₆₂₂A IN1 mutants (Fig 3A). Therefore, the interaction with AC40 is necessary for IN1 binding to Pol III *in vivo*. Since an interaction was detected *in vitro* between IN1 and the C34 and C53 proteins (Cheung *et al*, 2016), we asked whether these associations could be detected *in vivo* apart from the Pol III complex. This was not the case as both C53 and C34 co-immunoprecipitated with WT IN1 but not with the IN1 mutant K₆₁₇A (Fig EV2A and B). These data indicate that *in vivo*, the interaction of C34 and C53 with IN1 occurs primarily when Pol III is associated with IN1. Altogether, these data show that AC40 is determinant for the binding of IN1 to the Pol III complex.

To establish whether AC40 plays a major role in IN1 recruitment to Pol III-transcribed genes, we developed IN1 chromatin immunoprecipitation (ChIP) experiments to assay the effect of the K₆₁₇A, S₆₂₁A, and L₆₂₂A mutations on recruitment. WT IN1 and the various mutants were tagged at their N-terminus using a 3xHA epitope tag and expressed from a tetracycline-off promoter. Quantitative PCR revealed significant enrichment of ectopic WT HA-IN1 at all tested Pol III-transcript loci, compared to background level measured on the *GAL1* gene promoter (Fig 3B). In contrast, HA-IN1 mutants that did not interact with AC40 were barely detected at these loci. Thus, recruitment of IN1 to Pol III-transcribed loci depends on its interaction with AC40.

To assess whether the genome-wide occupancy of WT IN1 correlates with Ty1 integration site preferences, we performed ChIP sequencing (ChIP-seq) using the same HA-IN1 constructs, with a strain expressing IN1-Strep as control. Analysis of reads mapping to unique sites revealed a strong association of WT HA-IN1 with most nuclear *tDNAs* and the Pol III-transcribed genes *SNR6*, *SNR52*, *SCR1*, *RPR1*, and *RDN5*. Very weak or no HA-IN1 binding was observed for *RNA170* and *ZOD1*, previously shown to have low level of Pol III occupancy (Moqtaderi & Struhl, 2004) (Fig 3C and Table EV1). Three *tDNAs*, *tK(CUU)C*, *tM(CAU)C*, and *tD(GUC)N*, were not recovered: These genes are either absent or transcriptionally inactive in the laboratory strain we used (Kumar & Bhargava, 2013; Patterson *et al*, 2019). HA-IN1 was absent from most Pol II-transcribed genes (Table EV1) or present at a much lower level than at Pol III-transcribed genes (Fig 3C). Low HA-IN1 occupancy at Pol II-transcribed genes may be an artifact of ChIP-seq due to the level of expression of these genes (Teytelman *et al*, 2013). The genome-wide distribution of ectopic WT IN1 revealed a strong bias for Pol III-transcribed genes, confirming that the interaction of IN1 with the Pol III is the main driver for targeted integration of Ty1.

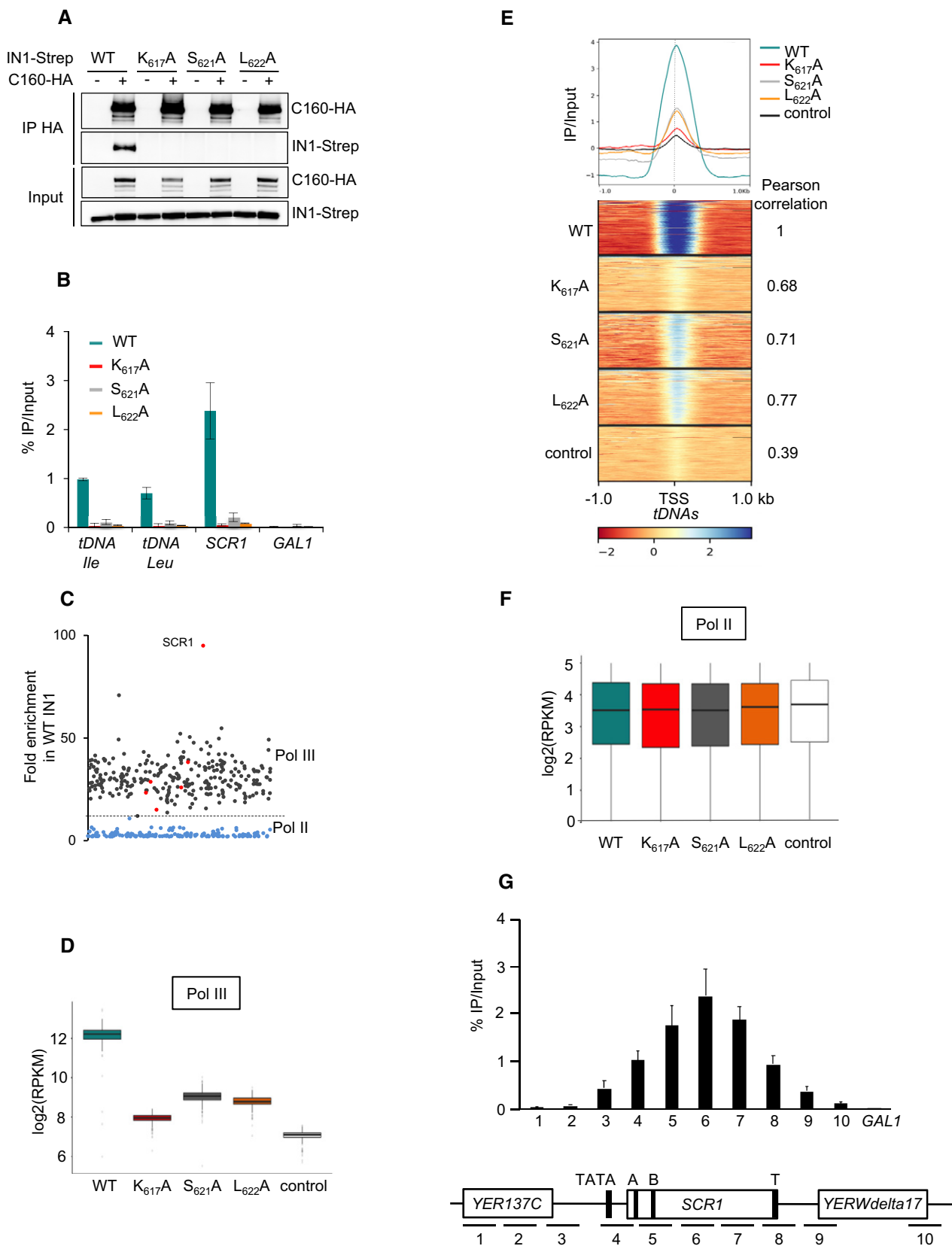


Figure 3.

Figure 3. AC40 recruits IN1 at Pol III-transcribed genes.

- A Co-immunoprecipitation of ectopic IN1 using C160-HA as bait, from yeast protein extracts expressing WT or mutated IN1-Strep (K_{617A} , S_{621A} , or L_{622A}) from the *GAL1* promoter in the presence of galactose. Expected sizes are 160 kDa for C160-HA and 100 kDa for IN1-Strep (WT and mutants).
- B Quantitative ChIP analysis of HA-IN1 enrichment at Pol III-transcribed genes. Immunoprecipitated DNA from yeast cells producing ectopic IN1 is expressed as a value relative to that of the input. Pol III-transcribed genes: *tDNA-Ile* and *tDNA-Leu* families (16 and 22 genes, respectively) and the unique *SCR1* gene. *GAL1* ORF serves as a control. Data represent means \pm SD ($n \geq 3$).
- C Fold enrichment of WT HA-IN1 over input at all the genes where HA-IN1 has been detected by ChIP-seq analysis. Each dot represents a gene. Gray, *tDNAs*; red, other Pol III-transcribed genes; blue, Pol II-transcribed genes. The dashed line indicates the separation between Pol II and Pol III-transcribed genes.
- D WT and mutant IN1 association with all Pol III-transcribed genes. Values obtained from ChIP-seq analysis have been normalized in log₂ RPKM (reads per kilobase per million mapped reads). Control is anti-HA immunoprecipitation of chromatin extracts in cells expressing IN1-Strep. Immunopurified DNA from 3–4 independent experiments was combined in one replicate. Data are presented as boxplots showing the median, the first quartile, and the third quartile. Error bars show minimum and maximum values. Regions used for calculation are gene bodies.
- E Genome-wide occupancy profiles (top) and heatmaps (bottom) of WT and HA-IN1 mutants over a 1 kb window upstream and downstream of all *tDNAs*. Pearson correlation to WT values corresponding to Fig EV2D is indicated.
- F WT and mutant IN1 association to all Pol II-transcribed genes, respectively. Control, replicates and data presentation, as described for panel D. Regions used for calculation are gene bodies.
- G Recruitment of IN1 along the *SCR1* gene locus. *Top*. ChIP-qPCR analysis as described for panel Fig 3B. Data represent means \pm SD ($n = 3$). *Bottom*. Schematic representation of the *SCR1* locus with DNA amplicon positions. TATA, A and B boxes and transcription terminator (T) are indicated.
- Source data are available online for this figure.

ChIP-seq analysis of the HA-IN1 mutants that compromise the IN1-AC40 interaction (K_{617A} , S_{621A} , and L_{622A}) revealed their occupancy was substantially reduced at all Pol III-transcribed genes, as indicated by the lower number of reads corresponding to Pol III-transcribed genes with the three mutants, compare to WT HA-IN1 (Fig 3D), and by a metagene analysis comparing WT and mutant HA-IN1 binding on all the 275 nuclear *tDNAs* (Fig 3E). No effect of these mutants was observed at Pol II-transcribed genes (Fig 3F). Quantification by pair-wise Pearson correlation between WT and either K_{617A} , S_{621A} , or L_{622A} IN1 confirmed the apparent stronger decrease in Pol III-transcribed gene occupancy of K_{617A} HA-IN1 compared to the other mutants (Figs 3E and EV2C and D). The metagene analysis indicated a sharp peak around the transcription start site (TSS), which does not coincide with Ty1 integration sites, normally located upstream of Pol III-transcribed genes (Baller *et al*, 2012; Mularoni *et al*, 2012; Bridier-Nahmias *et al*, 2015). Since *tDNAs* are too small to allow spatial resolution of Pol III localization, we performed ChIP analysis on *SCR1*, the longest Pol III-transcribed gene, to determine whether IN1 is associated with the polymerase over the entire gene. WT HA-IN1 was detected all along the *SCR1* gene with maximum binding in the coding region (Fig 3G). This pattern is similar to that observed with Pol III (Ghavi-Helm *et al*, 2008), confirming that the interaction with Pol III is determinant for IN1 recruitment to the chromatin. HA-IN1 occupancy was not completely suppressed in the three mutants, suggesting that the K_{617A} , S_{621A} , or L_{622A} mutants may have a residual level of interaction with AC40 that would not be detected in experiments where IN1 is overexpressed (*i.e.*, in two-hybrid or *in vitro* Co-IP assays). Alternatively, additional protein–protein interactions, like those previously identified with other Pol III subunits, could contribute to the recruitment of Ty1 integration complex at Pol III-transcribed genes in the absence of the IN1-AC40 interaction (Cheung *et al*, 2016).

AC40 is common to both Pol I and Pol III, suggesting that IN1 may also interact with Pol I. Consistently, we found that Pol I is associated with IN1 *in vivo*. Pull-down of A190-TAP, the largest subunit of Pol I, retained WT IN1-HBH, whereas no association was detected with the IN1 mutants (Fig 4A). The *RDN1* locus is composed of 100–200 tandem repeats of the 35S-precursor *rDNA* (*RDN37*), transcribed by Pol I, and the 5S *rDNA* (*RDN5*), transcribed

by Pol III (Dammann *et al*, 1993). Analysis of ChIP-seq reads mapping at multiple positions revealed WT HA-IN1 at both genes (Fig 4B). IN1 occupancy may be overestimated, as reads corresponding to all repeats are aggregated on the two copies that are represented in the *S. cerevisiae* reference genome (<https://www.yeastgenome.org>). However, IN1 mutants compromising the IN1-AC40 interaction— S_{621A} , L_{622A} , and particularly K_{617A} —reduced HA-IN1 occupancy at both loci. Since ChIP-seq data result in a poor resolution of the *RDN1* locus, we also performed conventional ChIP experiments that showed a clear enrichment of IN1 at both Pol I- and Pol III-transcribed genes but not at the intergenic NTS1 and NTS2 regions (Fig 4C), confirming that the presence of IN1 correlates with that of the two polymerases. WT HA-IN1 occupancy appeared lower at *RDN5* than at other Pol III-transcribed genes (compare Figs 3B and 4C), probably because Pol III is less associated with *RDN5* (Moqtaderi & Struhl, 2004). On the other hand, IN1 binding to the 18S locus transcribed by Pol I decreased in the presence of the K_{617A} , S_{621A} , or L_{622A} mutations (Fig 4D). Altogether, these results support a role of AC40 in the binding of IN1 to the *RDN1* repeats.

Finally, the reduced association of HA-IN1 mutants with Pol I- and Pol III-transcribed genes did not result in a significant increase of IN1 occupancy at other specific loci and particularly at subtelomeres (Table EV1). Collectively, these results indicate that the IN1-AC40 interaction is necessary for the interaction between IN1 and Pol III and thus its recruitment at Pol III-transcribed genes. IN1 also interacts with Pol I via AC40 and is present at Pol I-transcribed genes.

Non-AC40 binding Ty1 mutants have altered integration profiles

To investigate the integration profile of Ty1 mutants that have an impaired IN1-AC40 interaction, libraries of His⁺ selected *de novo* Ty1 insertion events were generated in cells expressing WT or mutant Ty1*his3AI* elements from the *GAL1* promoter (Barkova *et al*, 2018). We used an *spt3-101* null *rad52Δ* mutant strain to avoid both trans-complementation of the mutant IN1 by endogenous WT IN1 and Rad52-dependent recombination events. Initially, we performed qualitative PCR to monitor Ty1 insertion events at the *SUF16* tRNA gene and the *SEO1* subtelomeric gene. These genes were identified

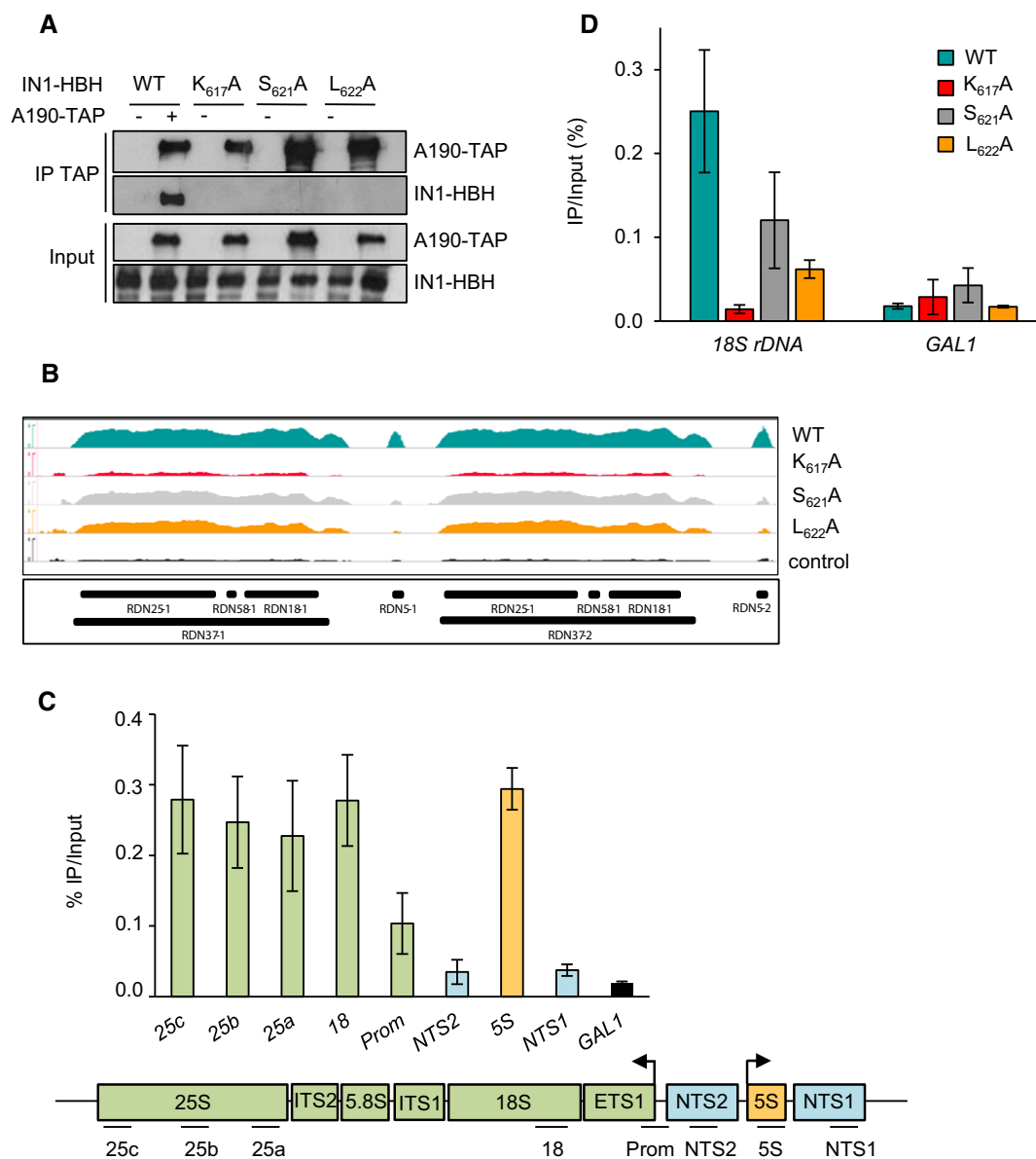


Figure 4. AC40 recruits IN1 at the RDN1 repeats.

A Co-immunoprecipitation of ectopic IN1 using TAP-tagged-A190 as bait from yeast protein extracts expressing WT or the indicated IN1-HBH mutants from a pTet-Off promoter in the absence of doxycycline. Expected sizes are 204 kDa for A190-HA and 82 kDa for IN1-Strep (WT and mutants).

B Genome browser visualization of HA-IN1 occupancy at the RDN1 locus. Occupancy of WT HA-IN1 and K₆₁₇A, S₆₂₁A, and L₆₂₂A HA-IN1 mutants is represented in each panel. Control, as described for panel 3D.

C Top. IN1 is recruited at Pol I-transcribed genes. ChIP-qPCR analysis as described for panel 3B. Data represent means \pm SD ($n = 3$). Bottom. Schematic representation of the RDN1 locus with DNA amplicon positions.

D Recruitment of WT and mutant HA-IN1 (K₆₁₇A, S₆₂₁A, or L₆₂₂A) at the 18S rDNA locus transcribed by Pol I. ChIP-qPCR analysis as described for panel 3B. Data represent means \pm SD ($n = 3$). GAL1 ORF serves as a control.

Source data are available online for this figure.

as hot spots of Ty1 integration in WT and AC40sp loss-of-interaction mutant, respectively (Bridier-Nahmias *et al*, 2015). In independent cultures expressing WT Ty1his3AI, we observed multiple bands upstream of the *SUF16* tRNA gene, characteristic of Ty1-HIS3 insertion in the three nucleosomes upstream of tDNA genes (Bachman *et al*, 2005; Dakshinamurthy *et al*, 2010) (Fig 5A). This profile was

significantly different for Ty1his3AI harboring K₆₁₇A, S₆₂₁A, or L₆₂₂A mutations in IN1, with many fewer integration events upstream of *SUF16*, and increased insertion at *SEO1*, compared to WT Ty1his3AI (Fig 5A). This observation suggests that IN1 mutations at K₆₁₇, S₆₂₁, and L₆₂₂ have the same effect on Ty1 integration site targeting as the AC40sp loss-of-interaction mutant.

To extend our analysis to the entire genome, we characterized Ty1-HIS3 *de novo* insertion event libraries using high-throughput sequencing. We could discriminate Ty1-HIS3 *de novo* insertions from endogenous elements using six nucleotides in the 3' LTR that

were specific to the Ty1his3AI element (Baller *et al*, 2012). By comparing the Z-score of Ty1 insertions on four non-overlapping features (Fig 5B), we confirmed that WT Ty1 insertions occurred mainly in a 1 kb window upstream of most Pol III-transcribed genes

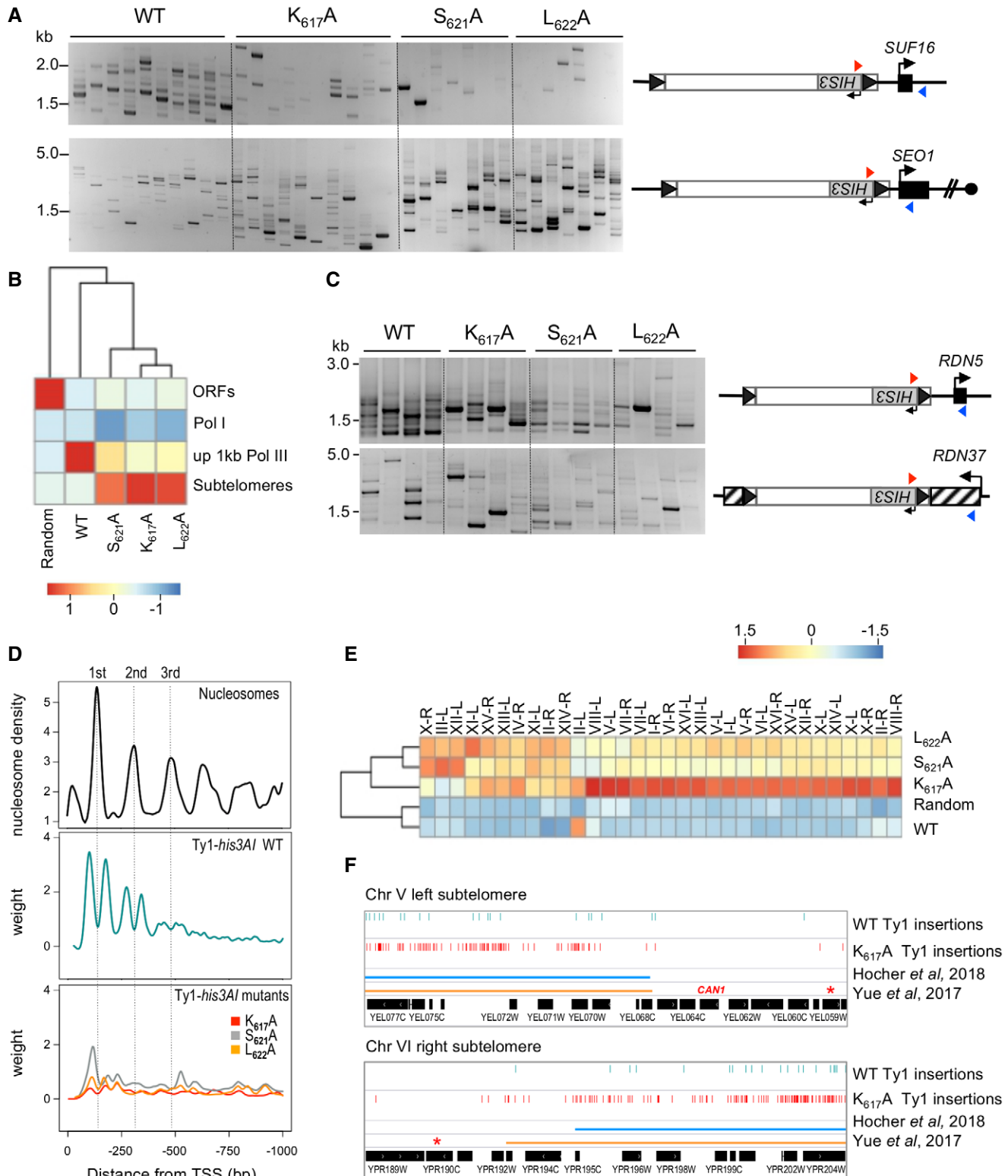


Figure 5.

Figure 5. Non-AC40 binding Ty1 mutants have altered integration profiles.

- A Detection of *de novo* Ty1 insertions upstream of the *SUF16* and *SEO1* genes by PCR using a primer in *HIS3* (red triangle) and a primer in the locus of interest (blue triangle). Ty1 retrotransposition was induced in cells transformed with plasmids expressing WT or mutant (IN1 K_{617A} , S_{621A} , and L_{622A}) Ty1 $his3AI$ from the *GAL1* promoter. Total genomic DNA was extracted from His⁺ cells obtained from independent cultures.
- B Genome-wide Ty1 insertion frequencies at each genomic feature are clustered in a heatmap. Score is computed in column Z-score. ORFs, all Pol II-transcribed genes except genes at subtelomeres; Pol I, one *RDN37* copy; up 1 kb Pol III, 1 kb upstream of all Pol III-transcribed genes; subtelomeres, genomic coordinates corresponding to chromatin covered by Sir2 and Sir3, when they are co-overexpressed (Hocher et al, 2018); Random, 100,000 random Ty1 computed insertions in the genome.
- C Detection of *de novo* Ty1 insertions upstream of *RDN5* and in *RDN37* genes by PCR as described for panel 5A. Four total genomic DNA preparations from panel A were randomly tested by PCR.
- D Ty1 insertion profile upstream of *tDNAs*. Total genomic DNA extracted in (B) was prepared for Ty1 *de novo* integration event sequencing. Ty1 insertions are computed in a 1 kb window upstream of all the 275 nuclear *tDNAs* (position 0 in the graph). Each position is divided by the number of insertions at this position (weight). The Smoothing curves indicate the general trend. Nucleosome center positions for the three-first nucleosomes upstream of all *tDNAs* are from (Brogaard et al, 2012).
- E Ty1 insertion frequencies for each left and right subtelomere of chromosomes are clustered in a heatmap. Score is computed in row Z-score. Random, as described for panel B. High level of WT Ty1-*HIS3* integration in II-L is due to the presence of the tRNA gene *tF(GAA)B*.
- F Genome browser visualization of WT and IN1 K_{617A} mutant Ty1-*HIS3* insertions into chromosome V left and chromosome VI right subtelomeres compared to the subtelomere boundaries defined by Hocher et al (2018) and Yue et al (2017), respectively. Red stars indicate the first essential gene of each subtelomere. The *CAN1* gene is indicated on chromosome V.

Source data are available online for this figure.

(Table EV2), the only exceptions being *tDNAs* that were absent or not transcribed in our strain. There were significantly fewer Ty1-*HIS3* insertions at Pol III-transcribed genes with the three IN1 mutants (Fig 5B). However, the preference for Pol III-transcribed genes was not completely lost, as insertions in these regions were higher than expected if Ty1 targeting was random, supporting the idea that the interaction with Pol III is not fully abolished with these mutants. Using the sequencing reads mapping at multiple positions in the genome, we also detected Ty1-*HIS3 de novo* insertion events at *RDN5* and *RDN37* genes in the *RDN1* repeats and in the adjacent *RDN5* variants (Fig EV3A). To determine whether these insertions were or not an artifact of aggregating integrations occurring in all *RDN1* repeats in the only two repeats of the reference genome, we compared the percentage of Ty1-*HIS3* insertions to random insertions generated *in silico* in two artificial yeast genomes composed of 20 and 60 *RDN1* repeats. We relied on two studies to estimate the actual number of actively transcribed *RDN1* repeats. In Ide et al, a yeast strain with only 20 *RDN1* copies is viable and displays no growth defect, suggesting that the level of Pol I transcription in WT cells may correspond to only 20 actively transcribed copies (Ide et al, 2010), while Merz et al estimate that, among the 150–200 *rDNA* repeats present in the genome, about 30–50%, i.e. approximately 60 copies, are actively transcribed (Merz et al, 2008). With this approach, we observed a 7- to 20-fold enrichment of WT Ty1-*HIS3* insertions in a 1-kb window upstream of *RDN5* locus compared to the two situations where these insertions would be randomly distributed (Fig EV3B). In the presence of the K_{617A} , S_{621A} , or L_{622A} mutations, the percentage of insertions decreased but was still higher than the two random situations (Fig EV3B). A similar decrease was observed by qualitative PCR (Fig 5C). This could be explained by the residual recruitment of IN1 mutants still observed by ChIP-seq (Fig 4B). This confirms that the Pol III-transcribed gene *RDN5* is a hotspot of Ty1 insertions, which depends on the interaction between IN1 and AC40. In contrast, the percentage of WT Ty1-*HIS3* insertion events at *RDN37* was close to the random situations, lower than at *RDN5* and not decreased by the K_{617A} , S_{621A} , or L_{622A} mutations (Fig 5C and EV3B). These results indicate that Ty1 integration is not preferentially targeted to *RDN37* and that the recruitment of IN1 at Pol I-transcribed genes is not sufficient for subsequent Ty1 integration.

WT Ty1-*HIS3* insertions displayed a periodic profile in the region of the three nucleosomes located upstream of Pol III-transcribed genes, with two insertion sites per nucleosome, as seen previously (Fig 5D) (Baller et al, 2012; Mularoni et al, 2012; Bridier-Nahmias et al, 2015). This profile was modified with the three mutants, with the first site of the first nucleosome being less affected with the S_{621A} mutant, suggesting that close proximity to the *tDNA* is a determinant of integration. The K_{617A} mutant, which had the lowest HA-IN1 occupancy at Pol III-transcribed genes (Fig 3D and E), displayed the largest decrease in integration events at Pol III-transcribed genes (Fig 5B and D). This correlation suggests that the strength of the IN1-AC40 interaction influences integration efficiency at Pol III-transcribed genes. Concomitant with the decrease in integration at Pol III-transcribed genes, the three IN1 mutants showed an increase in integration events at the end of each chromosome arm, a phenotype that was most pronounced for the K_{617A} mutant (Fig 5B and E). These results are consistent with the redistribution of Ty1 insertions in these regions observed in the AC40sp loss-of-interaction mutant (Bridier-Nahmias et al, 2015). Ty1 *de novo* insertions at the ends of chromosomes were mostly located in regions defined as subtelomeres, based on heterochromatin specificities or loss of synteny between different *Saccharomyces* strains (Yue et al, 2017; Hocher et al, 2018), suggesting that these subtelomeric regions harbor determinants allowing Ty1 targeting (Fig 5F). In contrast, no increase in Ty1-*HIS3 de novo* insertion events was detected in Pol II-transcribed genes for the three mutants (Fig 5B), as exemplified for the *CAN1* gene (Fig 5F).

Altogether, these results further support a major role for the IN1-AC40 interaction in Ty1 integration targeting at Pol III-transcribed genes. They also confirm that, when this interaction is compromised, Ty1 insertions are not random but principally occur in subtelomeres. Based on these data, we have named the IN1 sequence that contains the amino acids required for the interaction with AC40, the Ty1 targeting domain of IN1 (TD1).

IN1 targeting domain directs Ty5 integration at Pol III-transcribed genes

To determine whether TD1 is sufficient to confer Ty1 integration site preferences, we transferred the sequence into the Ty5

retrotransposon, which preferentially integrates into heterochromatin at yeast silent mating loci (*HMR* and *HML*) and near telomeres (Zou *et al*, 1996). Ty5 selectivity relies on an interaction between a hexapeptide (TD5, targeting domain of Ty5) in the C-terminus of IN5 and the heterochromatin protein Sir4 (Gai & Voytas, 1998; Xie *et al*, 2001). Exchange of TD5 for the IN1 bNLS containing TD1 in IN5, expressed in a two-hybrid vector, resulted in IN5 interacting with AC40, but not with Sir4 (Figs 6A and EV4A–D). WT IN5, IN5 lacking TD5 (IN5 $_{\Delta$ TD5), and IN5 $_{\Delta$ TD5+bNLS harboring the L₆₂₂A mutation in TD1, all failed to interact with AC40, demonstrating that the interaction was strictly dependent on TD1.

To establish whether the interaction with AC40 is sufficient to target IN5 $_{\Delta$ TD5+bNLS to Pol III-transcribed genes, we performed ChIP-seq in strains ectopically expressing HA-tagged IN5, IN5 $_{\Delta$ TD5, and IN5 $_{\Delta$ TD5+bNLS. Metagenome analysis of the tagged proteins binding to *tDNAs* revealed a clear enrichment of IN5 $_{\Delta$ TD5+bNLS at these loci, not detected for IN5 and IN5 $_{\Delta$ TD5 (Fig 6B). IN5 $_{\Delta$ TD5+bNLS was also present at the other Pol III-transcribed and Pol I-transcribed genes (Figs 6C and EV4E). Overall, IN5 $_{\Delta$ TD5+bNLS genome occupancy profile was very similar to that of IN1 (Pearson correlation of R = 0.9 Fig EV4F and Table EV3), confirming that the TD1-AC40 interaction is sufficient for recruitment at Pol III-transcribed genes. We did not detect IN5 enrichment at subtelomeric regions bound by Sir4 (Zill *et al*, 2010), nor at *HML* and *HMR*, which are Ty5 integration sites, which may be due to the weak association between Ty5 integration sites and Sir4 occupancy (Baller *et al*, 2011) and may reflect a loose and dynamic interaction between IN5 and Sir4 proteins difficult to detect by ChIP.

To explore the impact of the TD1-TD5 exchange on Ty5 integration site selectivity, we introduced IN5 $_{\Delta$ TD5+bNLS into a functional Ty5*his3AI* reporter expressed from the *GAL1* promoter. This replacement caused a 10-fold decrease in the frequency of Ty5 retrotransposition but did not inactivate the element. His⁺ colonies represented *bona fide* integration events, as a similar colony frequency was observed in the absence of homologous recombination (Fig EV4G). His⁺ selected *de novo* insertion events were generated for WT and mutant Ty5*his3AI*, and the insertion profiles were monitored at specific loci by qualitative PCR (Fig 6D). We used an *spt3-101* null strain that does not express endogenous Ty1 elements, to avoid interference between insertions of endogenous Ty1 and the mutant Ty5 at Pol III-transcribed genes. The Ty1*his3AI* profile displayed multiple insertion events at the Pol III-transcribed *SCR1* gene and glycine *tDNAs*, whereas no insertion was seen at *HML* and *HMR*. The WT Ty5 profile displayed multiple insertion events at *HML* and *HMR*, whereas no insertions were recovered at *SCR1* and very few were seen at the glycine *tDNAs*. In contrast, Ty5 $_{\Delta$ TD5+bNLS did not integrate at *HML* and *HMR*, whereas multiple insertion events were detected at the Pol III-reporter genes. The position of nucleosomes over the *SCR1* and glycine *tDNAs* suggests that the difference in Ty1 and Ty5 $_{\Delta$ TD5+bNLS banding patterns at Pol III-transcribed genes might be due to Ty1, but not Ty5, preferentially integrating into nucleosomes. Thus, the Ty1 preference for nucleosomes might not depend on interaction with AC40.

Together, these data indicate that TD1 is sufficient to direct the integration of the Ty5 retrotransposon upstream of Pol III-transcribed genes.

Discussion

Here, we identify the Ty1 targeting domain of IN1 (TD1) and show that it plays a critical role in the interaction with the Ty1 tethering factor AC40. We demonstrate that the interaction with AC40 orchestrates the recruitment of IN1 over the genome and that TD1 can function as an independent module that targets the integration of another related retrotransposon upstream of the typical Ty1 Pol III-transcribed target genes.

We provide several lines of evidence supporting a major role for AC40 in recruiting Ty1 to both Pol III and Pol I-transcribed genes. First, IN1 interacts directly with AC40 in the absence of other yeast proteins. Second, mutations in TD1 that reduce the interaction with AC40 abolish IN1 association with both Pol I and Pol III transcription complexes. Third, IN1 is present at both Pol I- and Pol III-transcribed genes and this presence depends on the IN1-AC40 interaction. Fourth, Ty5 integrase harboring TD1 interacts with AC40 and is recruited to Pol I- and Pol III-transcribed genes. Direct interactions were previously observed *in vitro* between IN1 and the C31, C34, and C53 Pol III-specific subunits but their precise role in the recruitment of the IN1 complex *in vivo* was not determined (Cheung *et al*, 2016). Here, we show that *in vivo*, the interaction of IN1 with C53 and C34 within Pol III depends on the IN1-AC40 interaction. The redistribution of Ty1 integration into subtelomeres, seen in the absence of the IN1-AC40 interaction ((Bridier-Nahmias *et al*, 2015) and this study), was not observed in a *rpc53 Δ 2-280* mutant, which decreases Ty1 integration at the *SUF16* tRNA gene (Cheung *et al*, 2016). This suggests that either C53 secures IN1 binding to Pol III once IN1 has been recruited by AC40 or helps Ty1 integration at a step downstream of IN1 recruitment. Further studies will be necessary to address the role of the Pol III complex, and especially of C31, C34, and C53, in Ty1 integration.

Upon maturation of the Gag-Pol protein in the VLPs, IN1 is associated with Ty1 cDNA. Under our ChIP experimental conditions, HA-IN1 was ectopically expressed and was therefore probably absent of the VLPs. Moreover, HA-IN1 was probably not associated with Ty1 cDNA, since cells were grown at 30°C, a temperature that restricts Ty1 replication resulting in a reduced level of Ty1 cDNA (Lawler *et al*, 2002; Garfinkel *et al*, 2005). Therefore, our results suggest that the cDNA might not be necessary for IN1 recruitment at Pol III-transcribed genes. A recent study reached a similar conclusion for the recruitment of Ty3 integrase at tRNA genes (Patterson *et al*, 2019).

Previous genome-wide mapping of Ty1 insertion sites did not reveal a clear pattern of Ty1 insertion into Pol I-transcribed genes at the *RDN1* locus (Baller *et al*, 2012; Mularoni *et al*, 2012; Bridier-Nahmias *et al*, 2015), likely due to the highly repetitive nature of the *RDN1* repeats (Bridier-Nahmias *et al*, 2015). Here, we show that Ty1 integration in the *RDN1* repeats occurs mostly at Pol III-transcribed genes and at nearly random levels in Pol I-transcribed genes. A similar pattern has already been described but was based on a small number of Ty1 insertions and was shown not to be the consequence of mitotic instability of Ty1 insertions in *RDN1* (Bryk *et al*, 1997). Importantly, the recruitment of IN1 by AC40 appears to be insufficient for Ty1 integration into Pol I-transcribed genes. Consistently, the loss of IN1-AC40 interaction did not further decrease the number of insertion events. These observations suggest that Ty1 integration into Pol I-transcribed genes may require additional co-

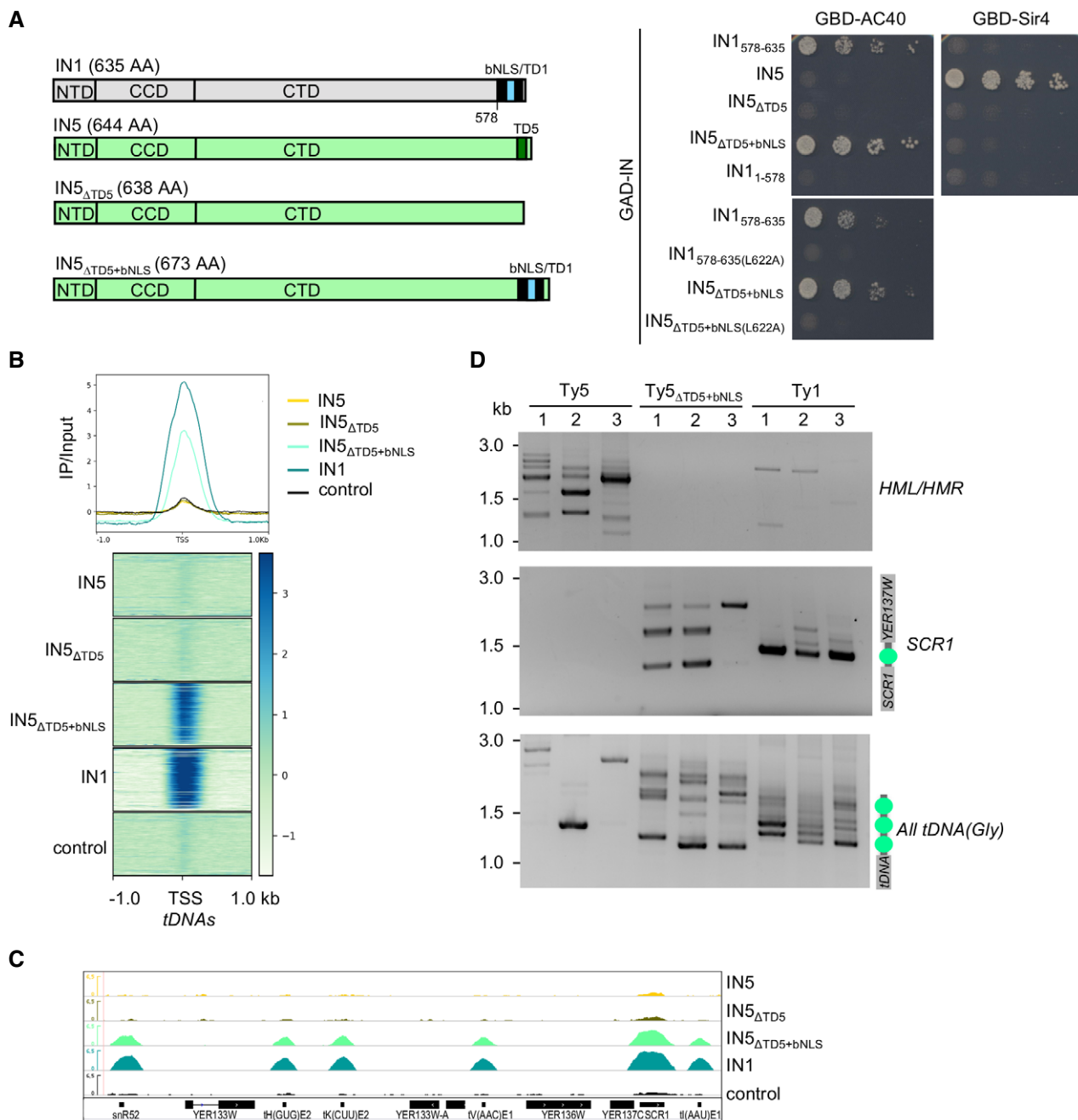


Figure 6. IN1 targeting domain directs Ty5 integration at Pol III-transcribed genes.

A Two-hybrid interaction between GBD-AC40 (left) or GBD-Sir4 (right) and different GAD-IN5 or GAD-IN1 constructions. Cells were plated in fivefold serial dilutions on DO-Leu-Trp-His plates. No growth or protein expression defects were detected (Fig EV4A–D).

B Genome-wide occupancy profiles (top) and heatmaps (bottom) of IN5, WT, and indicated mutants, ± 1 kb upstream and downstream of all *tDNAs*.

C Genome browser visualization of different HA-IN occupancy for chromosome V (chrV:431129..443275). Occupancy of the indicated integrase is represented in each panel. Control, as panel 3D. The region contains *tH(GUG)E2*, *tK(CUU)E2*, *tV(AAC)E1*, *tI(AAU)E1*, *SNR52*, and *SCR1*, all transcribed by Pol III.

D Detection of Ty5, Ty5 Δ TD5+bNLS or Ty1 *de novo* integration events at *HMR* and *HML* loci, *SCR1*, and upstream of all glycine *tDNAs*, by PCR using a primer in *HIS3* and a primer in the locus of interest. Retrotransposition was induced in cells transformed with WT or mutated pGAL1-Ty5his3AI (Ty5 or Ty5 Δ TD5+bNLS) and pGAL1-Ty1his3AI (Ty1). Total genomic DNA was extracted from His⁺ cells obtained from independent cultures. Nucleosome position (green circles) is indicated on the right of the panels (Broggaard et al, 2012).

Source data are available online for this figure.

factors or a particular chromatin structure, like the well-positioned nucleosomes present upstream of Pol III-transcribed genes.

The IN1 mutants that disrupt the interaction with AC40 induce the same redistribution of Ty1 insertions at chromosome ends as seen in a AC40sp loss-of-interaction mutant (Bridier-Nahmias *et al*, 2015). The insertion sites of these Ty1 mutants are scattered throughout subtelomeric regions. Ty5 insertions also occur throughout subtelomeric regions (Baller *et al*, 2011). This scattered dispersion may explain why we have failed to detect IN1 and IN5 by ChIP-seq in these regions. High-resolution mapping of DNA binding sites will be required to address this point (Hafner *et al*, 2018; Meers *et al*, 2019).

To date, the specific retargeting of TE integration sites when the main targeting is compromised has only been observed for Ty1. When the interaction between HIV, MLV, and Ty5 INs and their primary tethering factors (LEDGF/p75, BET proteins, and Sir4, respectively) is altered, the integration of these retroelements at their usual targets decreases substantially and becomes random (Gai & Voytas, 1998; Schrijvers *et al*, 2012; Wang *et al*, 2012; De Rijck *et al*, 2013; Sharma *et al*, 2013). Chromosome ends are preferential targets of several TE families in different organisms (Casacuberta, 2017). In *S. cerevisiae*, subtelomeres are devoid of essential genes, are rich in stress-responsive genes, and evolve rapidly in response to stress (Snoek *et al*, 2014) or domestication (Yue *et al*, 2017). Targeting of Ty1 integration may have evolved to provide a balance between integration into “safe” genomic regions, i.e., *tDNAs*, and integration into fast-evolving regions when adaptation is necessary, i.e., subtelomeres. Accordingly, we propose that the IN1-AC40 interaction may be regulated by environmental stress. The observations that Ty5 targeted integration requires phosphorylation of the IN5 targeting domain, which is reduced by stress (Dai *et al*, 2007), and nutrient starvation regulates the Ty1 replication cycle (Morillon *et al*, 2000; Todeschini *et al*, 2005) both lend support this hypothesis.

The pattern of integration of the Ty1 IN1 mutants largely overlaps in subtelomeric domains (Yue *et al*, 2017; Hocher *et al*, 2018). This correlation suggests that specific feature(s) in subtelomeres attract or facilitate Ty1 integration in the absence of the IN1-AC40 interaction. As we failed to detect an interaction between IN1 and Sir4, Ty1 integration into subtelomeres probably involves a mechanism different from that of Ty5. In a *cis*-targeting model, IN1 is tethered to subtelomeres through a weak interaction with a subtelomeric specific co-factor, such that the subtelomeric interaction will only be favored when binding to AC40 is compromised. The dispersed nature of Ty1 insertion sites suggests that the co-factor could be distributed across subtelomeres, like a histone mark specifically enriched in these regions (Hocher *et al*, 2018). Alternatively, IN1 co-factor could be present at a limited number of subtelomeric sites, and after recruitment, the Ty1 intasome would scan for a chromatin environment favorable for integration. Given Ty1 targets stable nucleosomes upstream of Pol III-transcribed genes (Baller *et al*, 2012; Mularoni *et al*, 2012), nucleosome stability could also be a determinant for integration at subtelomeres. A similar two-step targeting model has been proposed to explain the absence of correlation between Sir4 binding sites and Ty5 integration sites in subtelomeres (Baller *et al*, 2011). In a *trans*-targeting model, the proximity of the subtelomeres with the nuclear pores (Zimmer & Fabre, 2011), through which the Ty1 intasome transits, would

facilitate Ty1 integration in these regions, especially when the interaction with AC40 is compromised. Consistent with this hypothesis, mutations in several components of the nuclear pore alter Ty1 integration preferences (Manhas *et al*, 2018). HIV integration also occurs preferentially in chromatin proximal to the nuclear periphery (Di Primio *et al*, 2013; Lelek *et al*, 2015; Marini *et al*, 2015).

This work demonstrates that the IN1 targeting domain functions as an independent module to target integration at Pol III-transcribed genes: The addition of this sequence to Ty5 IN is sufficient to direct Ty5 integration to these loci. Entry of the Ty1 intasome into the nucleus involves the classical import machinery and requires an interaction between importin- α and the IN1 bNLS (McLane *et al*, 2008). The IN1 bNLS consists of two regions of basic amino acids, which are essential for IN1 nuclear import, separated by a linker sequence that also contributes, although to a lesser extent, to import (Kenna *et al*, 1998; Moore *et al*, 1998; Lange *et al*, 2011). The residues that are required for IN1 interaction with AC40 and Ty1 integration upstream of Pol III-transcribed genes cluster in a short peptide, namely TD1, within the linker. The IN1 linker has been proposed to induce a conformation that facilitates interaction of the two basic amino acid-rich regions with two NLS-binding pockets present in importin- α (Kosugi *et al*, 2009; Lange *et al*, 2011). The conformation of the linker when the basic amino acid-rich regions are bound to importin- α could expose TD1 recognized by AC40. Mutations in TD1 substantially reduce the interaction with AC40 but not Ty1 retrotransposition frequency or IN1 nuclear import (this study). Thus, IN1 bNLS and TD1 have independent functions in nuclear import and integration targeting, respectively.

Retroviral vectors have been used in gene therapy to correct various monogenic disorders. However, these vectors rely on the properties of HIV and MLV integrases, whose preferences for either transcribed genes (i.e., HIV) or promoters (i.e., MLV) make them potentially harmful for the genome (Anguela & High, 2019; Goswami *et al*, 2019). Chimeras between IN from retroviruses and retroelements and chromatin-binding proteins have already been constructed to channel integration to new positions in the genome (Gao *et al*, 2008; Ferris *et al*, 2010). As Ty1 integration upstream of Pol III-transcribed genes preserves gene integrity (Bolton & Boeke, 2003) and given that Pol III transcription and structure, including the presence of AC40 and nucleosome positioning at Pol III-transcribed genes (Helbo *et al*, 2017), are conserved between yeast and humans, adapting Ty1 integration targeting to retroviral integrases could overcome the risk of insertional mutagenesis associated with current MLV and HIV retroviral-based vectors, allowing the development of safer retrovirus-based vectors for use in human gene transfer technologies.

Materials and Methods

Growth media, yeast strains, and plasmids construction

Saccharomyces cerevisiae strains used in this study were grown using standard methods and are listed in Table EV4. All plasmids and primers used in this study are reported in Tables EV5 and EV6. Plasmids were constructed using standard molecular biology procedures (details can be obtained on request). Mutations were introduced in plasmids with Q5[®] Site-directed mutagenesis (NEB). All

the constructs were validated by DNA sequencing (Eurofins Genomics).

Two-hybrid assays

Two-hybrid assays were performed in strain PJ69-4A which contains the *HIS3* reporter to detect positive interactions (James et al, 1996). Cultures of transformants were grown overnight at 30°C in synthetic complete medium lacking leucine and tryptophan (SC-LEU-TRP) to maintain plasmid selection. Serial dilutions of aliquots of 1 optical density at 600 nm (OD_{600}), washed in 1 ml of H_2O , were plated on SC-LEU-TRP (growth control) or SC-LEU-TRP-HIS (interaction), starting from 10^{-1} . Plates were incubated 4 days at 20°C. Each two-hybrid assay is a representative example of at least two biological replicates.

Transposition assays

To estimate the frequency of retrotransposition in strains transformed by p*GAL1-Ty1his3AI*, p*GAL1-Ty5his3AI*, or their derivatives, four independent transformants of each strain were grown to saturation 2 days at 30°C in liquid SC-URA containing 2% raffinose. Each culture was diluted thousand-fold in liquid SC-URA containing 2% galactose and grown 5 days to saturation at 20°C, which is the optimal temperature for Ty1 retrotransposition. Aliquots of cultures were plated on YEPD (100 μ l at 10^{-5}) and SC-HIS (Ty1: 100 μ l at 10^{-2} , Ty5: 1 ml). Plates were incubated for 3 days at 30°C and colonies counted to determine the fraction of [HIS^+] prototroph. Retrotransposition frequencies were defined as the mean of at least four experiments, and each one performed with four independent transformants.

Direct fluorescence microscopy

BY4742 WT cells were transformed with pMET-GFP₂-IN1 NLS constructs and pUN100-Nup49-mCherry. Cells were grown overnight in SC-URA-LEU medium at 30°C. Cells were then washed twice in H_2O and resuspended in SC-URA-LEU-MET to induce GFP₂-IN1 NLS expression for 6 h at 30°C. Live cell imaging was done using a widefield microscopy system featuring a Nikon Ti-E body equipped with the Perfect Focus System and a 100 \times oil immersion objective. We used an Andor Neo sCMOS camera, which features a large field of view of $276 \times 233 \mu$ m at a pixel size of 108 nm. We acquired 3D z-stacks consisting of 15 frames with z-steps of 300 nm. We used a dual-band filter set (eGFP, mRFP) and for each z position acquired two color channels consecutively with an exposure time of 300 ms. The complete imaging system including camera, piezo, LEDs (SpectraX) was controlled by the Andor IQ2 software.

PCR assays for detection of Ty1 and Ty5 integration events

After retrotransposition induction as described above, total genomic DNA was extracted from yeast cultures grown at 20°C for 5 days according to classical procedures (Barkova et al, 2018). Double-strand DNA (dsDNA) concentration was determined using Qubit™ fluorometric quantification (Thermo Fisher Scientific). *De novo* Ty1 insertions upstream of *tG(GCC)C (SUF16)* were amplified with PCR

primers O-AL27 and O-AB91 and at the *SEO1* subtelomeric gene with PCR primers O-AL27 and O-AL10. *De novo* Ty5 insertions at the *HML* and *HMR* loci, the *SCR1* gene, and all glycine *tDNAs* were amplified with O-AL27 combined with O-CG27, O-AL64, or O-AB115, respectively. PCR reaction consisted of 30 ng of dsDNA (or 75 ng for detection of insertions at *SEO1*), 5 μ l Buffer 5 \times , 0.5 μ l dNTP 10 mM, 0.625 μ l of each primer at 20 μ M, 0.25 μ l of Phusion DNA Polymerase (Thermo Scientific) in a 25 μ l final volume. Amplification was performed with the following cycling conditions in ProFlex™ PCR System (Life Technologies) cycler: 98°C 2 min, 30 \times [98°C 10 s, 60°C 30 s, 72°C 1 min], 72°C 5 min, and hold 4°C. PCR products were separated on a 1.5% agarose gel.

Co-precipitation experiments

For experiments performed with proteins expressed in *E. coli*, BL21 (DE3) Rosetta (Novagen) bacterial cells were transformed with pet28-6xHis-IN1-C-tag (C-tag is a four amino acid peptide tag (Djender et al, 2014)) and pACYC-AC40-Twin-Strep-tag, and streaked on LB agar plates supplemented with ampicillin, chloramphenicol, and 1% glucose. Transformants were inoculated in the same medium overnight at 30°C, diluted to 0.05 OD_{600} in fresh 100 ml of LB medium with ampicillin and chloramphenicol without glucose and grown at 30°C to an $OD_{600} = 0.5$. Cells were transferred at 24°C for 30 min, and protein expression was induced by adding 0.5 mM isopropyl 1-thio- β -D-galactopyranoside (IPTG) and growing cells for 3 h at 24°C. Cells were harvested at 1,100 g for 15 min at room temperature, washed with water, and resuspended in 1 ml of ice-cold extraction buffer (20 mM Tris-HCl pH 7.5, 300 mM NaCl, 10% glycerol, 0.1% NP40) before adding 1 μ g/ml of lysozyme. The extract was kept 10 min on ice and then lysed by sonication (Q700 sonicator, Qsonica) on ice for 5 cycles (output amplitude 10%, 5 s ON, 40 s OFF). The lysate was centrifuged at 21,000 g at 4°C for 30 min, and the supernatant was collected on a separate tube. A 800 μ l of protein extract was incubated with 40 μ l of Magstrep XT beads (Iba-lifescience) for 3 h at 4°C on a wheel and washed twice with 1 ml of extraction buffer. Bound proteins were eluted with 50 μ l of 1 \times Laemmli sample buffer, separated by SDS-PAGE, and analyzed by Western blot using CaptureSelect™ Biotin Anti-C-tag conjugate (1/5,000 dilution, Thermo Fisher) and anti-Strep-Tactin-HRP conjugate (1/5,000 dilution, Iba-lifescience). Experiments were reproduced at least 3 times with independent cultures.

Co-IP experiments in yeast cells between Pol III (C160-HA) and IN1-Strep were performed as followed. MW4415 or LV33 yeast strains expressing HA-tagged or untagged C160 respectively were transformed with a 2 micron plasmid expressing WT or K_{617A}, S_{621A}, or L_{622A} IN1 mutants from an inducible *GAL1* promoter. Overnight cell cultures were grown to saturation at 30°C in SC-URA to maintain plasmid selection, were diluted at $OD_{600} = 0.001$ in 100 ml of SC-URA and containing 2% galactose, and grown at 30°C. At $OD_{600} = 1$, cultures were harvested by centrifugation at 4°C. Cell pellets were resuspended in 500 μ l of IP extraction buffer (50 mM HEPES-KOH pH 7.5, 300 mM NaCl, 1 mM EDTA, 0.05% NP40, 0.5 mM DTT, 5% glycerol) supplemented with protease inhibitors (Thermo Fisher), and lysed with 0.25 ml of glass beads using a vortex (Disruptor Genie®, VWR) for 30 min at 4°C. The whole protein extract (except an aliquot kept for the input control) was incubated with 50 μ l of Dynabeads Pan Mouse IgG (Thermo Fisher)

coated with anti-HA antibody (12CA5, Roche) 1 h at 4°C on a wheel. Samples were washed three times with IP extraction buffer. Immunoprecipitated proteins were eluted from beads by boiling samples for 5 min at 95°C with 25 µl of 2× SDS sample buffer, and the entire eluted fraction was analyzed by Western blot. Total lanes correspond to 1/20 dilution of the total input engaged in the IP. Proteins were detected with thousand-fold diluted primary antibodies (anti-HA antibody (12CA5, Roche) and anti-Strep (Qiagen)) and revealed with ECL (Thermo Fisher) and Fusion FX camera (Vilbert-Lourmat). Experiments were reproduced twice with independent cultures.

Co-IP experiments in yeast cells between Pol I (A190-TAP) and ectopically expressed IN1-HBH (HBH is an histidine biotin tag (Tagwerker *et al*, 2006)) were performed as described for Pol III but with some minor modifications. All the cultures (MGD353-13D or MGD353-13D A190-TAP transformed by pCM185-IN1-HBH) were grown in SC-TRP containing 2% glucose. Immunoprecipitation was performed with Dynabeads Pan Mouse IgG. Proteins were detected with primary antibodies anti-TAP (1/5000 dilution, Invitrogen), and anti-Streptavidin-HRP (1/15,000 dilution, Pierce). Experiments were reproduced three times with independent cultures.

Co-IP experiments in yeast cells between ectopically expressed IN1-Strep and C34 or C53 were done as described for Pol III, except that IP was performed using Dynabeads Pan Mouse IgG (Thermo Fisher) coated with anti-Strep (Qiagen). C34 and C53 proteins were detected with primary antibodies (Huet *et al*, 1985), at respectively 1/3,000 and 1/10,000 dilutions.

Chromatin immunoprecipitation and chromatin immunoprecipitation sequencing

Yeast strain LV1689 was transformed with pCM185 expressing 3xHA-IN1-C-tag WT or harboring IN1 K₆₁₇A, S₆₂₁A, or L₆₂₂A mutations. ChIP was performed as previously described (Harismendy *et al*, 2003), with minor modifications. Briefly, log-phase cultures (50 ml) in SC-TRP to maintain plasmid selection were cross-linked with 1% formaldehyde for 5 min at room temperature. Cells were pelleted by centrifugation and lysed with glass beads using an orbital shaker (IKA; VRX basic Vibrax; 500 g, 40 min, 4°C). The cell lysate was drawn off the beads and then spun for 20 min at 24,000 g in a microcentrifuge at 4°C. The cell pellet was resuspended in sonication buffer (50 mM HEPES-KOH pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% N-dodecylcholate, 1 mM PMSF, O-Complete protease inhibitor (Roche)), placed on a rotative wheel for 1 h at 4°C, and sonicated for 5 cycles (40 s ON at high level and 20 s OFF) in a Bioruptor (Diagenode; Denville, NJ, USA). After an additional 1-h incubation at 4°C on a rotative wheel, the solubilized chromatin was recovered by centrifugation for 15 min at 7,700 g at 4°C. The preparation of magnetic beads, immunoprecipitation with anti-HA 12CA5 antibodies, elution from beads, and reversal of cross-linking were performed as described (Harismendy *et al*, 2003). Immunoprecipitated DNA was purified using a QIAquick PCR Purification Kit (Qiagen). The purified DNA samples were analyzed by quantitative real-time PCR using the SYBR[®] Green PCR master Mix kit and an ABI PRISM 7500 (Applied Biosystems). The results were normalized with the input DNA PCR signals and indicated by relative IP in the graphs. For ChIP-seq analysis, the immunopurified

DNA from 3 to 4 independent experiments was combined after validation by quantitative real-time PCR of *SCR1* and *GAL1* genes. DNA sequencing of 40-nucleotide tags was performed on a GA-IIx, Hi-Seq, or Next-Seq sequencer using the procedures recommended by the manufacturer (Illumina). Input DNA and DNA from ChIP of a strain expressing IN-Strep were used as negative controls.

ChIP-seq data analysis

Raw sequence reads quality was controlled with FastQC software tool (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>). Reads were trimmed from the adapter sequence and matched to reference genome (SacCer3) using Bowtie2 (Langmead & Salzberg, 2012), with default parameters. Peak calling was performed with MACS2 (Zhang *et al*, 2008) by comparing ChIP to the corresponding input. Enrichment profile and heatmaps of normalized Reads per Kilo Millions of Mapped reads (RPKM) ratio of IP/input in Log₂ were plotted with Deeptools (Ramírez *et al*, 2016). Bigwig files were generated for genome browser visualizations. All figures were prepared using R packages (<http://www.R-project.org>). Graphics were generated using ggplot2 (Wickham, 2016).

Construction of *de novo* Ty1 insertion libraries for high-throughput sequencing

LV174 strain (*spt3-101 rad52Δ*) was transformed with pGTy1-*his3AI*-SCUF WT (Baller *et al*, 2012) or pGTy1-*his3AI*-SCUF harboring mutations in IN1 sequences (IN1 K₆₁₇A, S₆₂₁A, L₆₂₂A). For each transformant, total genomic DNA was extracted from 70,000 to 100,000 His⁺ colonies recovered from seven to ten independent cultures grown at 20°C in the presence of galactose as described in (Barkova *et al*, 2018). Fasteris prepared and sequenced Illumina libraries.

Ty1 integration data analysis

Pipeline described in the ChIP-seq Data Analysis section was used for quality check and trimming. To detect *de novo* Ty1 insertions, only reads containing the SCUF sequence and ending with Ty1 LTR sequence were considered. Clean reads were then matched to reference genome (SacCer3) using BWA short aligner (Li & Durbin, 2009) for paired-end reads. PCR duplicates were removed with MarkDuplicates (Picard tools, <http://broadinstitute.github.io/picard/>), and ambiguous reads ending by Ty1 LTR were discarded. Only properly aligned paired reads (one start and end coordinates) were retained for further analysis (WT 64 002 paired reads; K₆₁₇A, 48 845; S₆₂₁A, 5 052; L₆₂₂A, 11 654). Ty1 *de novo* integrations were assigned to the corresponding genomic features, and analyses were performed using in-house R (<http://www.R-project.org>) pipelines. To compare the associations of integration profiles (both *in vivo* and *in silico*) with selected genomic features, we used a statistical approach in which the enrichment of each association with a specific feature was expressed as a z-score, which reflects the number of standard deviations from the mean value of the compared features. With this approach, the differences in the total number of insertions recovered in each library do not impact the analysis. Z-score is calculated as follow: $z = (x - \mu) / ((x) / \text{number of insertions (specific$

feature)/number of total insertions, μ ; mean value (all compared features), (σ ; standard deviation (all compared features). Graphics were generated using ggplot2 (Wickham, 2016).

Data availability

The datasets produced in this study are available in the following databases:

- Chip-Seq data: Array Express E-MTAB-9038 (<https://www.ebi.ac.uk/arrayexpress/experiments/E-MTAB-9038/>);
- Ty1 *de novo* insertion data: Sequence Read Archive PRJNA597319 (<https://www.ncbi.nlm.nih.gov/bioproject/PRJNA597319/>).

Expanded View for this article is available online.

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Author contributions

AA-L, AB, CC, CG, HF, IA, NP, and RM performed the experiments. AA-L performed computational analysis. AA-L, AB, CC, JA, and PL analyzed data and prepared figures. AA-L, AB, JA, and PL wrote the manuscript. JA and PL conceived and supervised the study and secured funding.

Conflict of interest

The authors declare that they have no conflict of interest.

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