Investigating epigenome dynamics the RITE way
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CHAPTER 3

RECOMBINATION-INDUCED TAG EXCHANGE (RITE) CASSETTE SERIES TO MONITOR PROTEIN DYNAMICS IN SACCHAROMYCES CEREVISIAE

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PROTEINS ARE NOT STATIC ENTITIES. THEY ARE HIGHLY MOBILE AND THEIR STEADY STATE LEVELS
ARE ACHIEVED BY A BALANCE BETWEEN ONGOING SYNTHESIS AND DEGRADATION. THE DYNAMIC
PROPERTIES OF A PROTEIN CAN HAVE IMPORTANT CONSEQUENCES FOR ITS FUNCTION. FOR
EXAMPLE, WHEN A PROTEIN IS DEGRADED AND REPLACED BY A NEWLY SYNTHESIZED ONE, POST-
TRANSLATIONAL MODIFICATIONS ARE LOST AND NEED TO BE REINCORPORATED IN THE NEW MOLECULES.
PROTEIN STABILITY AND MOBILITY ARE ALSO RELEVANT FOR DUPLICATION OF MACROMOLECULAR
STRUCTURES OR ORGANELLES, WHICH INVOLVES COORDINATION OF PROTEIN INHERITANCE WITH THE
SYNTHESIS AND ASSEMBLY OF NEWLY SYNTHESIZED PROTEINS. TO MEASURE PROTEIN DYNAMICS
WE RECENTLY DEVELOPED A GENETIC PULSE-CHASE ASSAY CALLED RECOMBINATION-INDUCED
TAG EXCHANGE (RITE). RITE HAS BEEN SUCCESSFULLY USED IN SACCHAROMYCES CEREVISIAE
TO MEASURE TURNOVER AND INHERITANCE OF HISTONE PROTEINS, TO STUDY CHANGES IN POST-
TRANSLATIONAL MODIFICATIONS ON AGING PROTEINS, AND TO VISUALIZE THE SPATIOTEMPORAL
INHERITANCE OF PROTEIN COMPLEXES AND ORGANELLES IN DIVIDING CELLS. HERE WE DESCRIBE
A SERIES OF SUCCESSFUL RITE Cassettes THAT ARE DESIGNED FOR BIOCHEMICAL ANALYSES,
GENOMICS STUDIES, AS WELL AS SINGLE CELL FLUORESCENCE APPLICATIONS. IMPORTANTLY, THE
GENETIC NATURE AND THE STABILITY OF THE TAG-SWITCH OFFER THE UNIQUE POSSIBILITY TO
COMBINE RITE WITH HIGH-THROUGHPUT SCREENING FOR PROTEIN DYNAMICS MUTANTS AND
MECHANISMS. THE RITE Cassettes ARE WIDELY APPLICABLE, MODULAR BY DESIGN, AND CAN
THEREFORE BE EASILY ADAPTED FOR USE IN OTHER CELL TYPES OR ORGANISMS.
INTRODUCTION

Epitope tags provide powerful tools to study protein properties. In general, epitope tags provide a static snapshot of proteins in the cell. However, most proteins are dynamic and this is often an important aspect of their function (Russel et al. 2009; Hinkson and Elias 2011). For example, protein dynamics can influence the mobility and inheritance of proteins, the exchange of subunits of macromolecular complexes, access to otherwise occupied interaction sites of proteins or, when proteins are degraded and replaced by new ones, in resetting post-translational modifications (Hager et al. 2009; Radman-Livaja et al. 2011; Hotz et al. 2012; Menendez-Benito et al. 2013). Over the past few years, several techniques have been developed to measure or visualize protein dynamics. Some of these techniques, such as FRAP, TimeStamp, or derivatives thereof, make use of fluorescent fusion proteins to follow the movement or stability and synthesis of proteins in single cells (Lin and Tsien 2010; Butko et al. 2012). Other methods involve differential labeling of old and new proteins by using SILAC, radioactive labels, or labeling of specific proteins by using SNAP tags or FIASH-ReAsH technology (Jansen et al. 2007; Adams and Tsien 2008; Sweet et al. 2010; Zee et al. 2010; Ray-Gallet et al. 2011). Most of these methods allow detection of old and new proteins but only few methods provide the opportunity to specifically purify old and/or newly synthesized proteins by biochemical methods. This aspect is particularly relevant for the study of chromatin protein dynamics, where affinity purification allows mapping of protein occupancy and dynamics on specific regions of the genome. Several methods have recently been developed to measure chromatin protein dynamics (recently reviewed in Deal and Henikoff 2010a). One is the use of inducible overexpression of a tagged version of the protein of interest in the presence of an endogenously expressed untagged (or differentially tagged) copy (Korber et al. 2004; Dion et al. 2007; Jamai et al. 2007; Kim et al. 2007; Rufiange et al. 2007). Another method (CATCH-IT) involves labeling of newly synthesized proteins by amino acid analogs that can be coupled to biotin and thereby used for selective purification (Deal and Henikoff 2010a; Deal and Henikoff 2010b). In Physarum, the dynamics of histone proteins can be monitored at the single cell level by microinjection of small amounts of labeled histone proteins (Thiriet and Hayes 2005; Ejlassi-Lassallette et al. 2011). We recently developed a versatile and flexible method called Recombination-Induced Tag Exchange (RITE), in which epitope tags on an endogenous protein of interest can be swapped in a conditional manner by an inducible Cre recombinase (Verzijlbergen et al. 2010; De Vos et al. 2011; Radman-Livaja et al. 2011; Hotz et al. 2012).

The RITE system has been developed in budding yeast and is composed of two parts; a tandem-tag cassette that can be integrated behind the gene of interest for conditional C-terminal tagging, and a stably integrated and constitutively expressed hormone-dependent Cre recombinase that allows control of epitope
switching. RITE cassettes encode for one epitope tag (Tag1 or old tag) between two LoxP recombination sites and a second, orphan, epitope tag (Tag 2 or new tag) downstream of the second LoxP site (Fig. 1A). Upon activation of Cre recombinase activity by the simple addition of the hormone estradiol, a tag-switch occurs: the first tag is removed from the genome by recombination between the two LoxP sites and replaced by the second tag. To prevent background recombination, the RITE cassettes contain a selectable marker between the LoxP sites. Note that the LoxP recombination sequence is part of the protein coding sequence in the RITE cassettes, resulting in an in frame tag following the LoxP sites and allowing for switching by recombination.

RITE has several advantages over some of the other methods to measure protein dynamics. The proteins of interest are expressed from their endogenous promoter, avoiding potential problems with overexpression. In addition, a RITE switch does not require specific media changes and is permanent, which allows monitoring of protein dynamics under many different physiological conditions (De Vos et al. 2011; Radman-Livaja et al. 2011; Hotz et al. 2012; Ouellet and Barral 2012; Menendez-Benito et al. 2013). Importantly, old and newly synthesized proteins can be monitored simultaneously. The RITE system is flexible, widely applicable and compatible with switching between different short epitope tags as well as fluorescent tags. Furthermore, RITE allows selective tagging and following one protein of interest in the context of all other unlabeled protein. Finally, RITE allows dynamics measurements in the context of a genetic screen allowing identification of proteins controlling protein turnover (Verzijlbergen et al. 2011; Hotz et al. 2012). Of note, proteins that are subject to very high turnover may not be suitable for analysis by RITE, because Cre-induced recombination of the LoxP sites takes several hours to complete (see below and Verzijlbergen et al. 2010).

Here we present a comprehensive toolbox for the RITE assay. The availability of a RITE cassette series containing diverse biochemical and fluorescent tags allows for selection of tag pairs that are optimal for the protein of interest or for the experimental setup. Furthermore, new RITE cassettes with additional invariant tags are presented that bypass the need for protein-specific antibodies and allow for simultaneous detection of old and new protein (Fig. 1B). The RITE cassettes are modular by design. Therefore, they can be easily adapted to modify tags or to adjust cassettes for use in other cell types or organisms for which inducible Cre recombinases are available.

**MATERIALS AND METHODS**

**Strains and growth conditions**

Yeast and bacteria were cultured under standard conditions (Van Leeuwen and Gottschling 2002). E. coli strain DH5α was used for plasmid preparations. All
Figure 1. Outline of Recombination-Induced Tag Exchange. A) Following integration of a RITE cassette behind the gene of interest (GENE), recombination between LoxP sites is induced by Cre-Recombinase, causing a permanent switch from old Tag 1 to new Tag 2 on the protein of interest. S, spacer; L, LoxP recombination sites; TADH1, ADH1 terminator; HphMX, Hygromycin resistance cassette. B) RITE cassette including an invariant tag (i) upstream of the first LoxP site. The invariant tag is present pre- and post-recombination and can be used for simultaneous detection of the old and new protein of interest.

Yeast strains constructed and used in this study are detailed in Table 1 and were derived from previously published strains (Brachmann et al. 1998; Van Leeuwen et al. 2002; Tong and Boone 2006; Verzijlbergen et al. 2010; De Vos et al. 2011; Verzijlbergen et al. 2011). Strains were grown in YEPD (1% yeast extract, 2% bacto peptone, 2% glucose) in shaking flasks at 30°C. To select for Hygromycin resistance, cells were grown in YEPD containing 200 µg/ml Hygromycin.

Construction of RITE cassettes
All plasmids constructed in this study were derived from the previously described pFvL100 (Verzijlbergen et al. 2010) and are listed in Fig. 2A. pFvL106 contains a cassette switching from HA to T7 (HA→T7), but the HphMX selection marker (the Hygromycin B phospho transferase gene under control of the AgTEF1 promoter and terminator) is replaced by a URA3 selection marker. pFvL118 and pFvL119 (HA→T7) are nearly identical, except for small sequence differences around the HphMX gene (Verzijlbergen et al. 2010). For pFvL160, oligos were created containing either 2xT7 with BsrGI overhangs or 2xHA with SpeI overhangs. Single stranded oligos (Table 2) of matching tags were incubated in 40 mM Tris pH8 and 100 mM NaCl
<table>
<thead>
<tr>
<th>Name</th>
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<th>Switch parent</th>
<th>Source*</th>
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<td>NKI2036**</td>
<td>MATα his3Δ1 leu2Δ0 lys2Δ0 met15Δ0 ura3Δ0 hhf1-hht1Δ::LEU2</td>
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<td>KV</td>
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<td>NKI2036; bar1Δ::HisG his3Δ1::HIS3-P_{Tdh1}^::CRE-EBD78 hht2::HHT2-LoxP-Δ_{Adh1}^::HphMX-LoxP-T7</td>
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<td>KV</td>
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<td>NKI2148 This study</td>
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<td>BY4733</td>
<td>MATα his3Δ200 leu2Δ0 met15Δ0 trp1Δ63 ura3Δ0</td>
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<td>NKI8054</td>
<td>NKI2001; htz1::HTZ1-LoxP-2xT7-T_{Adh1}^-HphMX-LoxP-HA-6xHis</td>
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<td>This study</td>
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<td>NKI8055</td>
<td>NKI2001; htz1::HTZ1-LoxP-2xFLAG-T_{Adh1}^-HphMX-LoxP-HA-6xHis</td>
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<td>NKI4044</td>
<td>BY4733; pgk1::PGK1-V5-LoxP-HA-yEGFP-T_{Adh1}^-HphMX-LoxP-T7-mRFP</td>
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<td>NKI4044post</td>
<td>NKI4044; pgk1::PGK1-LoxP-T7-mRFP</td>
<td></td>
<td>NKI4044 This study</td>
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<td>Y7092</td>
<td>MATα can1Δ::P_{ade2}^::Sp-his5 lyp1Δ his3Δ1 leu2Δ0 ura3Δ0 met15Δ0</td>
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<td>AT</td>
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<td>SPC42G</td>
<td>Y7092; lyp1Δ::NATMX-P_{Tdh1}^::CRE-EBD78 spc42::SPC42-LoxP-T7-T_{Adh1}^-HphMX-LoxP-HA-GFP</td>
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<td>SPC42G This study</td>
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<tr>
<td>SPC42Gpost</td>
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<td>SPC42R</td>
<td>Y7092; lyp1Δ::NATMX-P_{Tdh1}^::CRE-EBD78 spc42::SPC42-LoxP-3xH A-T_{Adh1}^-HphMX-LoxP-3xT7-mRFP</td>
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<td>SPC42Rpost</td>
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<td>SPC42R This study</td>
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*Sources: KV, Verzijlbergen et al 2010; CB, Brachmann et al 1995; AT, Tong et al, 2006; FvL, van Leeuwen et al., 2002, see Materials and Methods.

**NKI2036 was derived from crosses between BY4742, BY4727 and UCC1369 (CB and FvL).
in a thermal cycler set at 94°C for 3 minutes, then gradually cooling down to 15°C. The double-stranded fragments were purified over a microspin G-50 column (Amersham Biosciences). Oligos were sequentially cloned into pFvL119 digested with the appropriate restriction enzymes. pTW073 was constructed by amplifying a fragment of pFvL118 with primers containing a V5 tag extension and BsrGI restriction sites. Subsequently, this fragment and pFvL118 were digested with BsrGI and ligated together, creating an HA→V5 cassette. A 3-step-PCR approach was used to construct pTW081 and pTW087. Fragments containing either HA flanked by MluI and SphI restriction sites, or T7 flanked by Clal and Nsil restriction sites were amplified from pFvL118. These fragments and pFvL118 were digested with the respective restriction enzymes and ligated to construct a cassette with swapped tags (pTW081; T7→HA; (Verzijlbergen et al. 2010)). For pTW087 a fragment containing HA was amplified using primers with a tail of 6xHis, creating an HA-6xHis fragment flanked by Clal and Nsil restriction sites. The fragment and pFvL118 were digested with Clal and Nsil and ligated to construct an HA-6xHis→T7 cassette (pTW087, (De Vos et al. 2011). An HA-6xHis fragment flanked by BgIII and BsrGI restriction sites was amplified by PCR from pTW087. The fragment and pTW081 were digested with these enzymes and ligated, constructing pTW094 (T7→HA-6xHis). For pMT001 (HA-6xHis→V5), plasmids pTW087 and pTW073 were digested with Nsil and MluI. The fragment of pTW087 containing spacer-LoxP-HA-6xHis- TADH1-HphMX was ligated into the fragment of pTW073 containing LoxP-V5. For pMT004-006 a fragment of double stranded DNA containing V5, 2xT7, or 2xFLAG with KpnI

Table 2. Oligos used for plasmid construction

<table>
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<th>Sequence</th>
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<tr>
<td>CLTSB19a</td>
<td>GTACTATGGCTTCTATGACAGGAGGTCAACAGATGGGAGGAATGGCCTCAATGACCCGGTGCCAGCAATGGGAT</td>
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<td>CLTSB19b</td>
<td>GTACATCCCATTTCGTCGGCCACCGGTCATTGAGGCCATTCCTCCCCTCTGTTGACCTCTGTCAATGAGCCTA</td>
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<td>CLTSB17a</td>
<td>CTAATCGTACTTCATGACACCGGCTCATTGAGGATTACGCTATCCTCGATAGTACGCTCTCAGTCAGTCC</td>
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<tr>
<td>CLTSB17b</td>
<td>CTAATCGTACTTCATGACACCGGCTCATTGAGGATTACGCTATCCTCGATAGTACGCTCTCAGTCAGTCC</td>
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<tr>
<td>KpnI-V5-SpeI Fwd</td>
<td>CGGTAAGCGCTATCCATACCCCTCCTCCCTCCTCGCACTGTTCTAGATATTACTGATCAGGATGAGTGAT</td>
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<td>KpnI-V5-SpeI Rev</td>
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<td>KpnI-2xFLAG-SpeI Fwd</td>
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<td>KpnI-2xFLAG-SpeI Rev</td>
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<td>KpnI-2xT7-SpeI Rev</td>
<td>CTAGTCATCTGTCACTCGTCTTGAAATACGCTCTCGAATAGTACGCTCTCGAATAGTACGCTCTCGAATAGTACGCT</td>
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and SpeI overhangs were made by oligo annealing (Table 2). Oligos were ligated into pFvL119 digested with *Kpn*I and SpeI. The tags of the resulting plasmids were combined with the HA-6xHis tag of pTW094 using Ascl and *SpI* restriction sites to generate pMT004 (V5→HA-6xHis), pMT005 (2xT7→HA-6xHis), and pMT006 (2xFLAG→HA-6xHis). pKV005 was constructed by a 3-step PCR on pKV001 (a pFvL119 derivative containing additional restriction sites), introducing an invariant MYC tag between the spacer and LoxP site. The resulting cassette is MYCi-HA→T7.

For pKV006, a V5-fragment with *Eco*47III restriction sites was amplified by PCR. The fragment and pKV001 were digested by *Eco*47III and ligated, constructing a cassette with an invariant V5 tag between the spacer and LoxP site (V5i-HA→T7). For pKV014, mRFP was amplified by PCR with primers containing *Bsr*GI and *Hind*III restriction sites at the ends. This fragment and pFvL160 were digested with *Bsr*GI and *Hind*III and ligated, resulting in a cassette with a fluorescent tag post-recombination (3xHA→3xT7-mRFP). For pKV015, yEGFP and mRFP were amplified with primers containing restriction sites for SpeI, and for *Bsr*GI and *Hind*III, respectively. These fragments and pKV006 were digested with the appropriate enzymes and ligated, resulting in a cassette that combines epitope tags with fluorescent tags, pKV015 (V5i-HA-yEGFP→T7-mRFP, (Verzijlbergen et al. 2010)). For pVM013, pTW081 and pVM012 were digested with *Bsr*GI and *Hind*III. The small fragment of pVM012 containing yEGFP was ligated into the large fragment of pTW081 (T7→HA), resulting in a T7→HA-yEGFP cassette (Menendez-Benito et al. 2013).

**PCR-mediated gene tagging**

To target RITE cassettes to the gene of interest, the cassette was amplified using integration primers that contain 40 bp of sequence of the gene of interest for homologous recombination (Fig. 2B). The forward primer (F1) sequence of the cassette is 5’-GGT GGA TCT GGT GGA TCT-3’. For pFvL106 to pKV006 the reverse primer used to amplify the cassette is 5’-AGGGAACAAAAGCTTGCATG-3’ (R1), which anneals 54 bp downstream of the cassette. For pKV014 to pKV016 the reverse primer sequence of the cassette is 5’-TCAAGGGCGCCGGTGGAGTGGCG-3’ (R3). This primer is mRFP specific and anneals at the end of mRFP to introduce a stop codon (underlined sequence), which is missing in the mRFP sequence. The reverse primer used for pVM013 is 5’-TGATTACGCAAAGCTCG-3’ (R2), which anneals further downstream than R1 (90 bp downstream of cassette) and which can be used when the sequence of R1 is absent. For one strain in this study (SPC42G), a reverse primer was used that anneals even further downstream of the cassette. When designing primers to target RITE to the gene of interest, it is important that the reading frame of the ORF and the fused cassette is maintained. The PCR products were transformed into strain NKI2176, NKI2036 or Y7092 using standard transformation protocols (Gietz and Schiestl 2007). After transformation, cells were plated onto YEPD plates and incubated overnight at 30°C; the following day plates
Figure 2. RITE cassettes and Cre recombinase vectors. A) Available RITE cassettes. The cassettes can be amplified with the primers indicated (see panel B). The sequence of the forward primer F1 is 5'-GGT GGA TCT GGT GGA TCT-3' (corresponding to the reading frame of the spacer sequence); the sequences of the reverse primers are: R1 5'-AGGGAACAAAAGCTTGCATG-3'; R2 5'-TGATTACGCCAAGCTCG-3'; R3 5'-TCA GGCGCCGGTGGAGTGGCG-3'. Note that primer R3 contains a STOP codon (underlined), since the mRFP sequence in pKV14-16 lacks a stop codon. B) Gene targeting of RITE cassettes by homologous recombination. The RITE cassettes can be PCR-amplified with a forward and reverse primer that have 20 bp homology with the cassette and a tail of at least 40 bp homology with the 3'end of the gene of interest and 3' UTR, respectively. C) Constructs to integrate the Cre recombinase expression vector in the yeast genome by homologous recombination. Unique restriction sites can be used to digest the plasmid and integrate the linear fragment by single cross-over (see Materials and Methods).
were replica-plated onto selection plates (YEPD containing 200 µg/ml Hygromycin). Integration of the cassettes was checked by colony PCR.

**Cre recombinase vectors**

Two plasmids were constructed to integrate Cre recombinase in the yeast genome. pTW040 was constructed by cloning a pTDH3-Cre-EBD78 fragment digested with PvuII into pRS303 digested with SmaI. For the pTDH3-Cre-EBD78 fragment, pTDH3 was cloned with ApaI and BspEI upstream of Cre-EBD78, where the pGAL had been replaced by a multiple cloning site. pSS146 was constructed by cloning the pTDH3-Cre-EBD78 fragment of pTW040 into pRS306 with EcoRI and NotI. Both plasmids contain unique restriction sites that can be used to integrate Cre-Recombinase at the HIS3 locus, the URA3 locus or the CYC1 terminator (Fig. 2C).

**Detection of recombination by Southern blot**

For Southern blotting 5x10⁸ cells were spun and frozen at −80°C. A histone H3 (HHT2)-specific probe was made by PCR amplification using primers HHT2_HindIII_for (GAATCTTTCTGTGACGCTTGG) and HHT2_HindIII_rev (GGGGGAAGAACAGTTGGAAGG), resulting in a 650-bp amplicon covering the region 576,144–576,794 of chromosome XIV. When used on genomic DNA that was digested with HindIII, the three bands recognized are specific for before the switch (3000 bp, pre), after the switch (931 bp, post), or as an internal control (1538 bp, control). Radioactive Southern blotting was performed using 50 μCi of ³²P-dCTP; incubation was done overnight at 65°C.

**Protein detection by immunoblot and antibodies**

For immunoblotting, strains were grown to mid-log phase (OD₆₆₀ 0.6-0.9). Samples of 2x10⁸ cells were harvested and washed with 10 mM Tris pH 8, 1 mM EDTA (TE) containing 0.2 mM PMSF. Cell pellets were stored at -80°C until further processing, but at least thirty minutes. Whole cell extracts were prepared in SUMEB (1% SDS, 8 M Urea, 10 mM MOPS pH 6.8, 10 mM EDTA, 0.01% bromophenol blue) containing protease inhibitors (1 mM PMSF, 1 mM DTT, 5 mM benzamidine, 1 µg/ml pepstatin, 1 µg/ml leupeptin) by glass bead disruption in a multivortex. The resulting lysate was incubated for 10 minutes at 65°C and subsequently clarified by centrifuging 5 minutes at 14000 rpm. Prior to immunoblotting, 4-10 µl of lysate was separated on a polyacrylamide gel (16% for histone H3 and H2B, 10% for Pgk1, Spc42, and Sir2). Separated proteins were transferred to a 0.45 µm nitrocellulose membrane for one (H3 and H2B) or two (Pgk1, Spc42, and Sir2) hours at 0.1 A. Membranes were blocked with PBS containing 2% or 5% Nutrilon (Nutricia) for one hour, first antibody incubations were done either for two hours at room temperature or overnight at 4°C in TBST with 2% Nutrilon. After washing three times in TBS containing 10% Tween-20 (TBST), secondary antibody incubation was performed in TBST with 2%
Nutrilon and LI-COR Odyssey IRDye 800CW antibody at 1:10,000 for 45 minutes at room temperature in the dark followed by two times 10-minute wash in TBST and one time 10-minute wash in PBS. Membranes were scanned using a LI-COR Odyssey IR Imager (Biosciences) and analyzed using the Odyssey LI-COR software package version 3.0. Antibodies used in this study are Pgk1 (A-6457, Invitrogen), Sir2 (Sc-6666, Santa Cruz), histone H2B (39238, Active Motif), HA (12CA5), T7 (A190-117A, Bethyl), Flag (M2 F3165, Sigma), V5 (R960-25, Invitrogen), histone H3 and LoxP (Verzijlbergen et al. 2010), and GFP and mRFP (Rocha et al. 2009).

Chromatin Immuno Precipitation

For ChIP, cells were grown to mid-log phase in YEPD with 200 µg/ml Hygromycin for pre-switch strains, or YEPD for post-switch strains. Samples of 1-3x10⁹ cells were taken, fixed for 10 minutes in 1% formaldehyde and washed with cold TBS. Pellets were stored in 12 ml flat bottom-tubes or 2 ml screw-cap tubes at -80°C until further processing. Cells in 12 ml tubes were disrupted in 300 µl breaking buffer (100 mM Tris pH 7.9, 20% glycerol, protease inhibitor cocktail EDTA-free (Roche)) with 400 µl glass beads in a multivortex for 20 minutes at 4°C. Cells in screw-cap tubes were disrupted in 200 µl breaking buffer with 500 µl glass beads in a bead beater in a 4°C cold block for 2 minutes. Lysis was at least 70%, as determined by microscopy. Lysates were transferred to 1.5 or 2 ml tubes and 1 ml FA buffer (50 mM HEPES-KOH pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% Na-deoxycholate, protease inhibitor cocktail EDTA-free) was added. The mixture was centrifuged for 1 minute at 14000 rpm at 4°C, and the pellet was washed once more with FA buffer. The pellet was resuspended in 450 µl FA, divided over two 1.5 ml tubes and sonicated for 6-7 minutes in a Bioruptor (Diagenode) with 30 second on-off cycles on high power. Lysates were cleared by centrifugation for 5 minutes at 4°C at 14000 rpm. Supernatant containing chromatin was transferred to 1.5 ml tube and 1 ml FA was added to samples of the 12 ml tubes. Screw-cap tube chromatin samples were treated with Micrococcal Nuclease (MN) to generate mononucleosomes. For these samples, 800 µl final buffer (15 mM Tris pH 7.4, 50 mM NaCl, 1.5 mM CaCl₂, 5 mM β-mercaptoethanol, 5 mM MgCl₂) was added. Samples were incubated with 30 units of MN (Fermentas) at 37°C for 20 minutes. The reaction was stopped by adding EGTA and EDTA to a final concentration of 10 mM and placing tubes on ice. The chromatin solution was centrifuged for 15 minutes at 14000 rpm at 4°C; the supernatant was transferred to a new 1.5 ml tube and stored at -20°C. Magnetic dynabeads coupled with Protein G (Dynal) were incubated in PBS containing 5 mg/ml BSA with antibody for at least four hours at 4°C. 200 µl soluble chromatin was added to prepared dynabeads and incubated rotating overnight at 4°C. Then 1 ml of FA buffer was added and samples were incubated rotating for 5 minutes at RT. The samples were washed twice with each of the buffers FA, FA-HS (50 mM HEPES-KOH pH 7.5, 500 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% Na-deoxycholate),
RIPA (10 mM Tris pH 8, 250 mM LiCl, 0.5% NP-40, 0.5% Na-deoxycholate, 1 mM EDTA). Finally, the samples were washed once with TE (10 mM Tris pH 8, 1 mM EDTA). 100 µl elution buffer (50 mM Tris pH 8, 10 mM EDTA, 1% SDS) was added to the samples and incubated for 10 minutes at 65°C. Subsequently the samples were centrifuged 1 minute at 14000 rpm and 80 µl supernatant was collected. 70 µl TE was added to samples and cross links were reversed in 0.625 mg/ml ProtK and 3 µg/ml RNaseA incubated for 1 hour at 50°C and subsequently overnight at 65°C. For input samples, 40 µl chromatin solution was combined with 60 µl elution buffer and 70 µl TE, and treated in the same manner as IP samples to reverse cross links. DNA was purified by using the High Pure PCR Product Purification Kit (Roche). Alternatively, DNA was extracted by using Chelex-100 resin (Bio-Rad) (Walsh et al. 1991; Nelson et al. 2009).

Quantitative PCR
Quantitative real-time-PCR (qPCR) was performed with SYBRgreen master mix (Applied Biosystems or Roche) according to the manufacturer’s manual. IP samples were diluted 10 times, and input samples were diluted 100 times before analyzing by qPCR on a 7500 Fast Real-Time PCR system (Applied Biosystems) or LightCycler 480 II (Roche). qPCR primers are shown in Table 3.

Microscopy
Microscopy samples were fixed with 4% formaldehyde, stained with 1 µg/ml Hoechst 33342 (Invitrogen) and mounted with Vectashield solution (Vector Laboratories) onto ConA-coated cover slides. Samples were analyzed on a Leica SP5 confocal system using a 405 nm laser to excite Hoechst, 488 nm for yEGFP and 561 nm for mRFP.

RESULTS AND DISCUSSION
RITE cassette and Cre vector construction
RITE cassettes containing different combinations of small epitope tags or fluorescent tags were generated by combining tag modules of previously described RITE

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>PTC6_Qfor</td>
<td>ATCGGGGCAATTAAGCATC</td>
</tr>
<tr>
<td>PTC6_Qrev</td>
<td>CCCGTAACAAGTCAGCTTC</td>
</tr>
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</tr>
<tr>
<td>RSA4_Qrev</td>
<td>CACAATATACAGATGCTCCACCTGT</td>
</tr>
<tr>
<td>SPA2_Qfor</td>
<td>ATCAAGAAGAGGGGTTCGACAAG</td>
</tr>
<tr>
<td>SPA2_Qrev</td>
<td>CATCGGCTCCGTTAATGG</td>
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</tbody>
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cassettes, or with new epitope tags as described in Materials and Methods. The RITE cassettes are shown in Fig. 2A. Many of the RITE cassettes contain unique restriction sites between the different elements, which facilitates modification of the constructs for other applications. Some of the RITE cassettes contain an invariant tag for simultaneous detection of old and new protein using commercially available antibodies. The cassettes can be integrated by homologous recombination behind any gene of interest (Fig. 2B).

Two vectors are available for integration of the hormone-dependent Cre recombinase (Fig. 2C). Unique restriction sites in the Cre vectors allow integration of the constructs by single crossover at the CYC1 terminator region, or at the HIS3 or URA3 regions. These options and the efficiency of integration depend on the auxotrophic alleles present in the target strain. For example, targeting of the HIS3-Cre cassette (pTW040) to the commonly used his3Δ200 allele (see http://wiki.yeastgenome.org/index.php/Strains) is inefficient due to the relatively short region of homology on one end. The advantage of targeting to this locus is that the integrated Cre is less prone to be lost by pop-out recombination. When other strategies are used it is recommended to maintain selection for the integrated Cre vector to select against pop-out events. Cre recombinase is expressed constitutively under control of the TDH3 promoter and CYC1 terminator and fused to the human Estrogen Binding Domain (EBD), which makes the nuclear activity of Cre dependent on the hormone β-estradiol (Logie and Stewart 1995). Estradiol releases the EBD from cytosolic heat shock proteins and allows entry of the EBD-Cre chimeric protein into the nucleus for recombination (Logie and Stewart 1995). This allows timed control of the RITE tag-switching. Here we use a derivative of Cre-EBD (Cre-EBD78) that is tightly dependent on the addition of β-estradiol in budding yeast (Lindstrom and Gottschling 2009; Dymond et al. 2011). Background recombination before induction of the tag-switch and recombination efficiency after induction can be determined by Southern blot analysis or by plating cells on non-selective media and then replica-plating the colonies to media containing Hygromycin (see Fig. 3 and Verzijlbergen et al. 2010). In a typical experiment, the average background recombination is 10% or lower, whereas the Cre-induced recombination efficiency is 95% or higher (see Figure 3 and Verzijlbergen et al. 2010). The completion of a recombination-induced tag-switch in a population of cells generally takes a few hours (Fig. 3). This time window should be taken into consideration when RITE is applied to study the dynamics of proteins with a very high turnover rate. Of note, the rate of induced versus background recombination can vary between strains, tagged genes, cell cycle stages, and experimental conditions. We occasionally encountered strains that showed high background recombination or low levels of induced recombination (data not shown), the reasons of which are unknown. We also found low levels of induced recombination in synthetic media containing mono sodium glutamate instead of ammonium sulphate as the nitrogen source (data not shown).
Immunoblot detection of RITE tags

To demonstrate the functionality of the short biochemical epitope tags in the new RITE cassettes, the respective RITE cassettes were PCR amplified and targeted to the histone H3 gene HHT2 (Fig. 2B). To avoid interference of non-tagged histone H3, the other gene encoding H3 (HHT1) was deleted. We analyzed strains prior to the tag-switch as well as strains that had undergone a permanent tag switch. Immunoblot analysis of whole cell extracts showed that both tags (pre-switch shows old Tag 1, post-switch shows new Tag 2) can be detected using tag-specific antibodies (Figs. 4A and B). The old Tag 1 was detected before the switch, while the new Tag 2 was detected after the switch. Due to background recombination, low levels of the

Figure 3: Cre-recombination kinetics. A) Southern blot analysis of H3-HA→T7 (strain NKI2048). Recombination was induced in log-phase or nutrient-starved cells and analyzed using an H3-specific probe on HindIII-digested DNA. The three bands recognized are specific for pre-switch (pre), post-switch (post), or an internal control present in both (control). B) Plating assay of starved Spc42-3xHA→3xT7-RFP cells (strain SPC42R) following induction of Cre-recombination. Cells were plated on YEPD media and then replica plated to YEPD media containing Hygromycin. C) Quantification of recombination levels as measured by plating assay of H3-V5→HA-6xHIS (strain NKI8050) and strain SPC42R in log phase and during starvation.
new Tag 2 were detected before the switch in some of the samples. RITE was also applied to Pgk1 (3-phosphoglycerate kinase), a housekeeping protein. In this case, the RITE cassette contained combinations of short epitope tags and fluorescent tags (pKV015). The old and new tags could be detected on immunoblots before and after the switch, respectively (Fig. 4C). In addition, the RITE cassette also harbors an invariant V5 tag, which should be present pre-switch as well as post-switch. Pgk1 containing the invariant V5 tag could be detected in both samples (Fig. 4C).

Detection of RITE tags by ChIP

One of the advantages of RITE compared to other methods for measuring protein dynamics is the possibility to apply it to affinity purification. We determined how the various tags in the RITE cassettes perform in ChIP experiments. For this purpose, we tagged histone H3 (HHT2, in the absence of HHT1) and histone variant H2A.Z (HTZ1) with different RITE cassettes and performed ChIP-qPCR experiments on pre-switch strains (see Materials and Methods). Three loci were examined (primer sequences are listed in Table 3), representing high (PTC6, RSA4) and low (SPA2) levels of steady state H2A.Z and similar levels of H3 (Kobor et al. 2004; Albert et al. 2007). Indeed, using the RITE tags, H2A.Z levels relative to H3 levels differed according to the previously shown steady state levels (Fig. 5). Note that although the trends were similar, the relative ChIP efficiency (H2A.Z/H3) was not the same for the different RITE tags (Fig. 5). Apparently, the short epitope tags do not work equally well for each protein. Fortunately, the RITE assay is very flexible. The expanded RITE cassette series that we describe here affords the selection of epitope tag pairs that work efficiently for the protein of interest. Furthermore, since the RITE cassettes are modular by design, other tag sequences can be readily incorporated.

Combining RITE with fluorescence microscopy

The addition of fluorescent markers to the RITE technology adds the possibility of spatiotemporal monitoring of proteins of interest (Verzijlbergen et al. 2010; Hotz et al. 2012; Menendez-Benito et al. 2013). We previously described the HA-yEGFP→T7-mRFP and 3xHA-yEGFP→3xT7-mRFP RITE cassettes (see Fig. 2A) (Verzijlbergen et al. 2010; Menendez-Benito et al. 2013). An mCherry→GFP RITE-like cassette has been described by the Barral lab (Hotz et al. 2012). Two additional RITE cassettes can be used to switch from a small epitope tag to a larger fluorescent tag (Fig. 2A). They can be used for conditional knock-ins when larger tags cause growth defects. As the protein of interest is only fluorescent after the switch, these RITE cassettes are particularly suited for monitoring synthesis and the spatiotemporal behavior of the new protein by microscopy. Fig. 6 shows Spc42, a component of the yeast spindle pole body, tagged with these RITE cassettes (Figs. 6A and D). The spindle pole body could be observed after the tag switch as a bright dot next to the nucleus (Fig. 6B and E), whereas before the tag switch only background signal was observed. Expression of the proteins was
Figure 4. Immunoblot analysis of RITE-tagged histone H3 and Pgk1. A) Histone H3 (HHT2) was tagged with different RITE cassettes. Before tag switch (pre) detects the old Tag 1. A wild-type strain (-, untagged H3) was used as a negative control. H2B was used as a loading control. B) After tag switch (post) detects the new Tag 2. Post-switch strains are the recombined counterparts of the pre-switch strains shown in panel A. Strains used: 1. NKI8001; 2. NKI2148; 3. NKI2158; 4. NKI2178; 5. NKI2220; 6. NKI8051; 7. NKI8056; 8. NKI8050; 9. NKI8052; 10. NKI8001; 11. NKI8085; 12. NKI4138; 13. NKI8086; 14. NKI8037; 15. NKI8089; 16. NKI8058; 17. NKI8087; 18. NKI8088. C) Pgk1 was tagged with an HA-yEGFP→T7-mRFP RITE cassette containing an invariant epitope tag V5 (V5i). Sir2 was used as a loading control.
confirmed by immunoblot analysis (Fig. 6C and 6F). Of note, such color switch tags can be combined with microscopy- or flow cytometry-based genetic screens to identify proteins that control protein or organelle dynamics (Hotz et al. 2012).

**Detecting protein dynamics**

RITE enables following protein dynamics under many physiological conditions. Here, we show the application of one of the new RITE cassettes, which switches from V5 to HA-6xHis and was applied to histone H3 (HHT2, in the absence of HHT1) to measure chromatin protein dynamics (Fig. 7A). Cells were arrested by starvation and a pre-switch sample was taken before inducing Cre recombinase (Fig. 7B). After 16 hours of recombination, a post-switch sample was taken before the cells were released in fresh media. Two additional samples were taken during the release, at 2 hours, before cell division, and at 4 hours, when most of the cells had divided once. During the release, the amount of old Tag 1 (V5) decreased, while new Tag 2 (HA-6xHis) increased, as expected (Fig. 7C). In the pre- and post-switch samples we detected V5-, but hardly any HA-tagged protein. The small amount of HA-tagged protein found in the post-switch sample is consistent with previous results showing that in starved cells there is a low level of histone exchange (Verzijlbergen et al. 2010). Using
an antibody against the spacer-LoxP peptide sequence we could detect old Tag 1 and new Tag 2 simultaneously on a single blot due to different mobility on SDS-PAGE gels.

Next, we performed ChIP on these samples and examined occupancy of old and new histone H3 at the PTC6 gene and the SPA2 gene (Table 3). For both genes we detected a decrease in V5-tagged protein (old H3) and an increase in HA-tagged protein (new H3) when switched cells were analyzed four hours after release into fresh media (Fig. 7D and E). The level of histone H3 exchange, i.e. the ratio of new H3 over old H3 (new/old), increased over time (Fig. 7F). Since the cells are released into log phase and undergo DNA replication, the exchange measured here was mostly caused by replication-coupled histone deposition, and to a lesser extent by replication-independent histone exchange, the latter of which can vary from gene to gene (Dion et al. 2007; Jamai et al. 2007; Rufiange et al. 2007; Deal et al. 2010; Verzijlbergen et al. 2010).

Figure 6. Analysis of fluorescent knock-in RITE cassettes. A) Spc42, a subunit of the spindle pole body, was tagged with a 3xHA→3xT7-mRFP RITE cassette (strain Spc42R). B) Fluorescent microscopy of strain Spc42R. In the pre-switch sample, no mRFP was detected; in the post-switch sample Spc42-mRFP was clearly visible. C) Immunoblot analysis of the Spc42R samples described in panel B. Pgk1 was used as loading control. D-F) as in panels A-C using Spc42 tagged with a 1xT7→1xHA-yEGFP RITE cassette (strain Spc42G). Spc42 containing a single HA or T7 tag could not be detected on blots using the tag-specific antibodies. However, an antibody against spacer-LoxP could visualize old and new Spc42. Scale bars, 2 μm.
Figure 7. Immunoblot and ChIP analysis of histone H3 dynamics. A) Histone H3 was tagged with a V5→HA-6xHis RITE cassette in strain NKI8050. B) Outline of experimental set up. C) Immunoblot analysis of cells arrested by starvation (pre), switched for 16 h (post), and subsequently released into fresh media for 2 hours (no cell division) or 4 hours (one cell division). An antibody against spacer-LoxP detects both old and new H3. D-E) H3-V5 and H3-HA-6xHis ChIP analysis of samples described in panel B to measure occupancy (IP/input) of old and new H3 at PTC6 (around TSS) and SPA2 (coding sequence) (primer sequences are listed in Table 3). F) Exchange of histone H3 determined as the ChIP ratio of new H3 over old H3 (HA/V5). Average of three biological replicates +/- SEM.
CONCLUSION

RITE is a versatile method that can be used to study protein dynamics by different downstream applications on any protein of interest that tolerates C-terminal tags. The GFP tagged yeast library shows the many examples where C-terminal tagging is allowed without significant effects on cell viability or other vital functions (Ghaemmaghami et al. 2003; Huh et al. 2003). The short epitope tags can be used for immunoblot and affinity purification protocols such as ChIP. Thereby, RITE can be combined with proteomics methods, genomics methods, or DNA-based high-throughput screens (Verzijlbergen et al. 2010; De Vos et al. 2011; Radman-Livaja et al. 2011; Verzijlbergen et al. 2011). The fluorescent tags in the RITE cassettes can be applied to measure protein dynamics in single cells by live imaging (Verzijlbergen et al. 2010; Hotz et al. 2012; Menendez-Benito et al. 2013) and further expanded towards high-throughput genetic screening. The suite of RITE cassettes enables a flexible design for many applications. Furthermore, the RITE cassettes are modular by design and can therefore be easily adapted for use in other cell types or organisms. The RITE plasmids and sequence information will be available through EUROSCARF (Frankfurt, http://web.uni-frankfurt.de/fb15/mikro/euroscarf/index.html).

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