Caught in the act: the lifetime of synaptic intermediates during the search for homology on DNA

Adam Mani¹, Ido Braslavsky², Rinat Arbel-Goren¹ and Joel Stavans¹,*

¹Department of Physics of Complex Systems, Weizmann Institute of Science, Rehovot 76100, Israel and ²Department of Physics and Astronomy, Ohio University, Athens, OH 45701, USA

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ABSTRACT
Homologous recombination plays pivotal roles in DNA repair and in the generation of genetic diversity. To locate homologous target sequences at which strand exchange can occur within a timescale that a cell’s biology demands, a single-stranded DNA-recombinase complex must search among a large number of sequences on a genome by forming synapses with chromosomal segments of DNA. A key element in the search is the time it takes for the two sequences of DNA to be compared, i.e. the synapse lifetime. Here, we visualize for the first time fluorescently tagged individual synapses formed by RecA, a prokaryotic recombinase, and measure their lifetime as a function of synapse length and differences in sequence between the participating DNAs. Surprisingly, lifetimes can be \( \sim 10 \) s long when the DNAs are fully heterologous, and much longer for partial homology, consistently with ensemble FRET measurements. Synapse lifetime increases rapidly as the length of a region of full homology at either the 3'- or 5'-ends of the invading single-stranded DNA increases above 30 bases. A few mismatches can reduce dramatically the lifetime of synapses formed with nearly homologous DNAs. These results suggest the need for facilitated homology search mechanisms to locate homology successfully within the timescales observed in vivo.

INTRODUCTION
Homologous recombination plays a fundamental role in the generation of genetic diversity, and as a pathway for DNA repair in both prokaryotes and eukaryotes. During horizontal gene transfer processes in bacteria, exogenous pieces of DNA thousands of bases long are imported into a cell and incorporated into the chromosome at specific loci of sufficiently high homology (1). Both the search for homology along a chromosome and the strand exchange process that ensues after homology is found are catalyzed by the RecA protein (2). Since chromosomal DNAs are very long molecules, the number of sequences that must be probed until homology is found can be exceedingly large. Yet, the search for homology is completed within a cell cycle, leading to the consideration of facilitated search processes (3,4), similarly to site-specific proteins searching for their targets (5,6).

The elementary step in the homology search is the formation of a complex (synapse) between a segment of the genomic double-stranded DNA (dsDNA) and a RecA-covered single-stranded tract of DNA (ssDNA). It is within a synapse that the two sequences are fully heterologous, and much longer for partial homology, consistently with ensemble FRET measurements. Synapse lifetime increases rapidly as the length of a region of full homology at either the 3'- or 5'-ends of the invading single-stranded DNA increases above 30 bases. A few mismatches can reduce dramatically the lifetime of synapses formed with nearly homologous DNAs. These results suggest the need for facilitated homology search mechanisms to locate homology successfully within the timescales observed in vivo.

*To whom correspondence should be addressed. Tel: +1 972 8 9342615; Fax: +1 972 8 9344109; Email: joel.stavans@weizmann.ac.il

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The fluorescent signal was imaged with an intensified CCD camera (PentaMAX, Princeton Instruments, Trenton, NJ), and frames were acquired continuously, with an exposure time of 2 s per frame to achieve a good signal-to-noise ratio.

**DNA sequences**

HPLC purified, 5' TAMRA tagged, ssDNA oligonucleotides, as well as 5' biotin modified 100-nt oligonucleotides, were purchased from IDT DNA (Coralville, IA) and from Metabion (Martinsried, Germany). Unmodified oligonucleotides were synthesized and purified by the Biological Services Department of the Weizmann Institute of Science. All ssDNA sequences were checked against secondary structure formation using the Mfold v3.2 software (16).

**Sample preparation for single molecule assays**

All chemicals were purchased from Sigma-Aldrich unless specified. All dsDNA constructs were attached to coverslips via their biotinylated tails, using previously described methods (17), but injecting dsDNA at 25 nM in EB3 buffer (50-mM Tris–HCl, 20-mM MgCl₂) into the chamber. To form the duplexes, biotin-modified 100-mers (sequence a1, see Supplementary Data), with either of the non-tagged oligomers 21a2, 50a2 and 70a, were mixed, at a molar ratio of 2:3, respectively, in 10-mM Tris–HCl, 100-mM NaCl, 10-mM MgCl₂ at pH 7.5, and were hybridized by cooling from 90°C to 20°C over 5 h. The suffix a2 was added to denote strands fully complementary to a portion of a1 beginning at the 3'-end of a1, so that the a1 and a2 hybridization results in a duplex having a biotin-modified single-stranded overhang through which the duplex is attached to the glass surface (Figure 1A). In the changing synapse length experiment, constructs were prepared by hybridization of a1 with 21a2, 50a2 or 70a2. All the rest of the experiments were carried out with a duplex formed by a1 and 50a2. For experiments testing synapse lifetime as a function of the length of homologous tract at the 3'-end of the incoming strand, the incoming strands HOM0, HOM15, HOM26, HOM35 and HOM50 were used. For experiments with a varying length of homologous tract at the 5'-end, the incoming strands HOM0, 5HOM26, 5HOM36 and HOM50 were used. For experiments testing the effects of point mutations on synapse lifetime, the incoming strands MM1 and MM3 were used. The sequences that were used are in the Supplementary Data.

**Sample preparation for FRET assays**

dsDNA was prepared by hybridization of the tagged oligomers Phi50 and Phi60 (see Supplementary Data). The competitor dsDNA was prepared by hybridization of the oligomers COM5 and COMAS. The complementarity strands were mixed in a 1:1 molar ratio in 10-mM HEPES–KOH (pH 7.5), 100-mM NaCl, 10-mM MgCl₂, and were hybridized by cooling from 90°C to 20°C over 5 h. The concentrations of the invading ssDNAs (PhiH50) were 2-fold those of tagged duplexes. FRET measurements were carried out in a reaction buffer containing

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**MATERIALS AND METHODS**

**Experimental setups**

An objective-type total internal reflection microscope (TIRF) was constructed using published methods (15). Evanescent illumination was achieved using a high numerical aperture objective off-axis as illustrated by the arrow, illuminates light produced by a narrow laser beam (entering a high numerical aperture objective) as illustrated by the arrow, illuminates only a ~200 nm thick layer above the glass. (B) Snapshot of synaptic intermediate complexes, ~25 μM across. (C) Fluorescence versus time of a single synapse formed with a fully heterologous, 50-bp-long ssDNA–RecA complex, or (D), with a fully homologous ssDNA–RecA complex. Traces and snapshot represent raw, unfiltered data.

Figure 1. Experimental scheme for observing individual synapses and sample results. (A) Fluorescently-tagged ssDNA–RecA complexes are observed only when forming synaptic intermediates with dsDNA anchored on a glass surface. The exponentially decaying evanescent light produced by a narrow laser beam (entering a high numerical aperture objective off-axis as illustrated by the arrow), illuminates only a ~200 nm thick layer above the glass. (B) Snapshot of synaptic intermediate complexes, ~25 μM across. (C) Fluorescence versus time of a single synapse formed with a fully heterologous, 50-bp-long ssDNA–RecA complex, or (D), with a fully homologous ssDNA–RecA complex. Traces and snapshot represent raw, unfiltered data.
20 mM HEPES–KOH (pH 7.5), 3.3 mM ATP and 20 mM MgCl₂. In these assays, 200 nM ssDNA and 3.5 μM E. coli RecA (New England Biolabs, Ipswich, MA, USA) were pre-incubated for 5 min at 37°C in the reaction buffer. Following pre-incubation, 100 nM of dsDNA tagged at both strands and the indicated ratio of competitor dsDNA were added and then injected into the measuring chamber as described previously (14). Experiments were carried out at 28°C. The sequences that were used are in the Supplementary Data.

Reaction conditions for single molecule assays

ssDNA (1–5 nM) having the same length as the short strand in the duplex, tagged with TAMRA at its 5'-end and bearing the sequence relevant to each of the experiments, was mixed with 1.5 μM RecA and 2 mM ATP in EB3 buffer containing an oxygen scavenging system [2.5 mM protocatechue acid, 30 mM protocatechu 3,4-dioxygenase (18), 1 mM Trolox (19)]. The mixture was then pre-incubated for 10 min at 37°C, in order to allow the pre-synaptic complexes to form. The mixture was then injected into the reaction cell. Using the oxygen scavenging system, the characteristic bleaching time of the fluorophores was ~15 min.

Data acquisition and analysis of single-molecule data

Frames acquired by the ICCD camera were captured by WinView32 software (Princeton Instruments, Trenton, NJ). The intensity of the molecules was tracked by a version of the ‘IDL Particle Tracking’ software (http://www.physics.emory.edu/~weeks/IDL; 20) adapted to MatLab (Mathworks, Natick, MA, USA) by D. Blair and E. Dufresne (http://physics.georgetown.edu/matlab). Briefly, the images were first spatially filtered with a bandpass filter to eliminate pixel noise and spatial variations in the illumination over the field of view. Then, the tracking procedure was used to generate intensity versus time information of each identified DNA site. The events were then counted and put into a histogram of lifetimes. The process was tested by marking specific molecules manually and acquiring their time traces. In compiling histograms of lifetimes, events lasting for only one frame (‘Materials and Methods’ section). In these experiments, the incoming and the short, non-anchored strand on the duplex had the same length (Figure 1A), and therefore the lifetime of synaptic complexes of smaller length for which these two strands were not in full registry—i.e. overlapping only along part of their full length—was also probed. It is noteworthy that the lifetimes of events of long duration are of the order of ~10 s irrespective of synapse length, in agreement with the estimates of the off rate of the synapse formation reaction (14) and FRET assay results presented.

RESULTS

In this study, we have measured the lifetime of synaptic complexes by visualizing them directly, at the single-molecule level. This technique offers a number of significant advantages over FRET (14) in measuring synapse lifetimes. The estimate based on FRET is indirect, as it is inferred from the retardation effected by a heterologous DNA competitor on a strand-exchange reaction. Moreover, the FRET measurements do not separate between the on rate of the reaction (synapse formation by molecular encounters in solution) and the off rate (reflecting the synapse lifetime). While the FRET estimate was made under the assumption that the rate of molecular encounters is much faster than the inverse lifetime (14), the single molecule study proves directly that the synapse lifetime is a rate limiting step. In order to measure synaptic complexes for as long as they last, surface anchoring and TIRF, as well as conditions enabling a long lifespan for the fluorophores were chosen.

In order to measure synapse lifetime at the single-molecule level, we used the scheme depicted in Figure 1A. The capture of tagged ssDNA–RecA complexes from solution by anchored duplexes resulted in the appearance of bright spots (Figure 1B). The spots were monitored over time until their disappearance (Supplementary videos 1 and 2 online show typical movies in the heterologous and homologous cases, respectively), and the duration of such events, which we refer to as the synapse lifetime was recorded. Typical intensity versus time traces of events corresponding to ssDNAs heterologous and homologous to the target duplexes are shown in Figure 1C and D, respectively.

The lifetime of heterologous synapses and its dependence on synapse length

We measured the lifetime of synapses formed between highly heterologous sequences for three synapse lengths: 21, 50 and 70 bp. The corresponding histograms, shown in Figure 2, have been normalized to unit area after neglecting events lasting only one frame (‘Materials and Methods’ section). In these experiments, the incoming and the short, non-anchored strand on the duplex had the same length (Figure 1A), and therefore the lifetime of synaptic complexes of smaller length for which these two strands were not in full registry—i.e. overlapping only along part of their full length—was also probed. It is noteworthy that the lifetimes of events of long duration are of the order of ~10 s irrespective of synapse length, in agreement with the estimates of the off rate of the synapse formation reaction (14) and FRET assay results presented.

![Figure 2. Histograms of lifetimes of individual, fully heterologous synapses versus synapse length. Three different synapse lengths were tested: 21, 50 and 70 bp (sequences 21SR, 50SR and 70SR, respectively; Supplementary Data). Each histogram (total area normalized to 1) comprises a few thousand events, obtained in at least two independent experimental runs.](https://academic.oup.com/nar/article-abstract/38/6/2036/3112554)
below, and contradicting assertions that scanning non-specific sites must be extremely fast (10).

We note that to accumulate significant statistics in the 21 bp synapse experiment a concentration of ssDNA–RecA complexes up to 30 times higher than in the 50- and 70 bp experiments had to be used (few events were observed when the concentration was similar to that used in experiments with longer synapses). The integration time of our camera (~2 s) limits us to observing only long events. For a given concentration of incoming ssDNA–RecA complexes, viewing fewer events for the 21-bp case may imply a decreased binding rate, as well as a shift of the lifetime distribution towards shorter times. In conclusion, synapse lifetime increases with length, although few events with lifetimes beyond ~200 s are seen.

In control experiments, the absence of RecA yielded approximately two events, in comparison to thousands in its presence. To assess the degree of non-specific binding of ssDNA–RecA complexes to the surface, experiments in the absence of anchored duplexes were carried out. We observed events of a wide range of durations, but their number was ~6% compared to experiments in the presence of duplexes. Taken together, these measurements demonstrate that even in the case of heterology, in which synapse lifetimes should be the shortest, lifetimes can still be seconds long.

**Measurement of average lifetime of heterologous synapses by ensemble FRET**

In order to ascertain that the second-long events observed in the single-molecule experiments are representative of the average synapse lifetime, we have extended the FRET measurements in Sagi et al. (14), probing events down to the millisecond timescale. Briefly, 50 bp dsDNA targets tagged at the same end with donor and acceptor moieties were combined with non-tagged fully homologous 50 nt ssDNA–RecA complexes in solution, in the presence of various concentrations of competing, fully heterologous non-tagged dsDNA (Figure 3A). The strand exchange reaction between the targets and the ssDNA–RecA complexes was monitored over time by following the decrease in the measured FRET efficiency, and the time by which this reaction was slowed down by the presence of the heterologous competitor dsDNA was measured. Under the assumption that the rate of the strand exchange reaction is limited by synapse lifetime and not by the rate of molecular encounters, it was proposed in (14) that the increase in the fraction of targets having undergone strand exchange, deduced from the FRET efficiency measurements, should be characterized by a rise time given approximately by

\[
\tau = \tau_0 + \frac{n_c}{n_0} \tau_s
\]

where \( \tau \) is the rise time in the presence of competitor and \( \tau_0 \) in its absence, \( n_c/n_0 \) is the molar ratio between competitor and target dsDNA concentrations and \( \tau_s \) is the average synapse lifetime. Note that the linearity of the rise time with competitor concentration implied by Equation (1) was not previously checked experimentally (14). Figure 3B shows plots of the fraction of strand exchange versus time for the reaction, in the presence of different heterologous competitor concentrations. The decrease in the saturation value with increasing competitor concentration (Figure 3B) is due to the reduction in the effective steady-state concentration of ssDNA by the competitor dsDNA. The balance is between the strand exchange and its opposite reaction, in which tagged ssDNA strands replace non-tagged ones in homologous duplexes. Solid lines represent exponential fits to the data. (C) Rise time of the curves in B as function of competitor concentration. The straight line is a linear fit to the data. Error bars represent the uncertainties derived from the fits in B.
The effects of sequence homology at the ends of an invading strand on synapse lifetime

The point of initial pairing between the pre-synaptic complex and the duplex may occur randomly along the length of DNAs involved (21). If local homology is found, a strand exchange reaction proceeds outwards from that point (see ‘Discussion’ section). In addition, it has been shown that there is a strong dependence of the end product of the recombination reaction on the different ends of an invading strand (14). We therefore investigated how synapse lifetimes change when a segment of full homology is included at either the 3’- or 5’-end of the invading strand. Lifetime measurements were carried out on 50-bp-long synapses formed with invading ssDNAs whose sequences were chosen so that the first x bases at the 3’ end were fully homologous with the target dsDNA, while the rest (50- x bases) shared no homology with the target. As Figure 4A clearly shows, events of long duration become more frequent as x increases (note that the last column in all the histograms includes also all events with lifetimes longer than 800 s). Similar results were observed when 26- or 36-base-long segments of homology were present at the 5’-end of the invading ssDNA strand (see Supplementary Figure S1). A control experiment in which the fluorophore was located at the opposite end of the invading strand yielded similar results. We conjecture that the enhanced stability of synapses formed with a region of homology at the incoming strand is associated with the stability of a nascent duplex.

Both in vivo and in vitro studies have revealed the existence of a minimum length of homology (~30 bp), known as the minimum efficient processing segment or MEPS, below which homologous recombination is inefficient (14,22). In order to determine whether our data bear any signature of the MEPS, we assume a minimal model in which observed events are divided into those for which the incoming strand is in full registry with the displaced strand and those for which it is not. There are about 200 configurations out of registry compared to one in registry for 50-base-long strands, considering both parallel and anti-parallel alignments. Within the model, we assume that out-of-registry configurations, all of which are highly heterologous, are characterized by a typical synapse lifetime \( \tau_{\text{het}} \), which we determine by a single exponential fit to the \( x=0 \) histogram, yielding \( \tau_{\text{het}} = 10.45 \pm 0.13 \) s. Assuming a typical synapse lifetime \( \tau_{\text{hom}} \) for in-registry events, we make fits of the normalized histograms for \( x \neq 0 \) with a sum of two exponentials:

\[
\text{Counts} = A_{\text{het}} e^{-t/\tau_{\text{het}}} + A_{\text{hom}} e^{-t/\tau_{\text{hom}}} \tag{2}
\]

with \( A_{\text{het}}, A_{\text{hom}} \) and \( \tau_{\text{hom}} \) as free parameters and \( \tau_{\text{het}} \) as fixed. As shown in Figure 4B, \( \tau_{\text{hom}} \) increases rapidly with \( x \) from ~30 bp, in full agreement with results in vivo and in vitro FRET measurements (14,22), and at \( x = 50 \) the histogram has becomes massively populated at long times. In this particular case, the fluorescence signal represents mostly events in which strand exchange has been completed and not synapses involving the tagged ssDNA–RecA and the anchored duplex. The duration of events in our measurements is limited from above both by photobleaching and by the time of observation, which cut long events short. We note that \( \tau_{\text{hom}} \) does not represent the average synapse lifetime due to these effects, as well as due to the insensitivity of our measurements to events whose duration is shorter than the integration time. Two interesting features of the fits of the data with Equation (2) are noteworthy: the ratio of weights of the two exponentials, \( A_{\text{hom}}/A_{\text{het}} = 0.040 \pm 0.005 \), is nearly constant for all \( x \neq 0 \), and its value agrees within an order of magnitude with the ratio of multiplicities of in-registry to out-of-registry configurations (note that of all possible out-of-registry configurations, only those with high enough overlap contribute significantly, as the experiments with 21-bp-long heterologous
synapses demonstrate). Allowing $\tau_{\text{het}}$ to vary as a free parameter in the double-exponential fits yields values that differ from 10.45s by not more than $\sim$10%, demonstrating that the separation of timescales between heterologous and partially homologous events is justified.

**Full homology is crucial for a stable nascent duplex**

The efficiency of strand exchange can be exquisitely sensitive to the presence of even a single mismatch, and few clustered mismatches reduce significantly the efficiency of strand exchange (14). Here, we tested the hypothesis that these effects should also manifest themselves as an inhibition of the formation of long-lived synapses. The histograms of synapse lifetimes displayed in Figure 5 illustrate the effects of one and three clustered mismatches together with data from fully homologous and highly heterologous sequences, also shown for comparison. Fitting the data with Equation (2) indeed shows that one mismatch reduces $\tau_{\text{hom}}$ from $\sim$221 s in the fully homologous case to $\sim$145 s, while a cluster of three mismatches separated from each other by less than three bases, starting at position 25, behaves nearly as if no homology were present after the cluster (compare with $x = 15$ and 26, Figure 4A). Hence, full homology during the initial stages of the process is a crucial determinant in promoting efficient strand exchange.

**DISCUSSION**

The present study demonstrates that the lifetime of fully heterologous synapses tens of base pairs long can be $\sim$10 s or longer, posing a severe challenge in a cellular context. This result has been obtained independently by interrogating molecular ensembles as well as single synapse. Our direct single molecule assay allows us to monitor synapses while they last, and to isolate the measurement of their lifetimes from any kinetics of molecular encounters, demonstrating that homology readout can be an intrinsically long process, independent of side reactions such as free RecA oligomerization (3).

The inclusion of a region of homology at either the 3’ or 5’ end of the incoming strand leads in general to the appearance of events with longer lifetimes, hinting at an enhancement of synaptic stability. The enhancement is, however, mild, in spite of the fact that the region of full homology can extend up to $\sim$30 bases, suggesting the synaptic complex is still unstable and dissociates within few tens of seconds. We conjecture that the enhanced stability of synapses formed with a region of homology at the ends of the incoming strand is associated with the stability of a nascent duplex, and these considerations determine the MEPS lengthscale (14,22).

RecA-mediated strand exchange is a process that consists of several steps. Our experiments show that the early step of synapse formation is not sensitive to whether a homologous tract is located at the 3’- or 5’-end of the invading strand. However, the final result of the process, namely the yield of a heteroduplex and a fully displaced strand, has been found to be much more sensitive to mismatches near the 3’-end of the invading strand rather than near the 5’-end (14). Consistent with this, it has also been reported that a homologous 3’-end is more invasive than a homologous 5’ one (23–25). This has been attributed to the higher likelihood of the 3’-end to be covered by RecA polymerization, which proceeds in the 5’- to 3’-direction (26,27). Other experiments have reported a higher invasiveness of the 5’-end (28). Our single molecule experiments, conducted under conditions in which the high concentration of RecA most likely ensured the complete coverage of the ssDNA, did not reveal a difference in invasiveness between the 3’- and 5’-ends.

The directionality of RecA-mediated homologous recombination has been extensively discussed in the literature. Initial pairing can occur at random points along homologous regions of the participating DNAs and extends in both directions from the point of initial pairing (21). The initial pairing involves local strand exchange (29). While our results do not rule out nascent duplex formation occurring randomly along the homologous tract on the ssDNA, we believe it is more likely that invasion occurs preferably from the incoming strand’s ends, due to the short length of the DNA used [for an extensive discussion see (29)].

Consistent with the nascent duplex interpretation, the presence of a single base mismatch in an otherwise fully homologous invading strand has a noticeable effect suppressing events of long duration. In addition, an incoming strand with three closely clustered mismatches so that adjacent ones were separated by no more than three bases (the site covered by each RecA monomer) reduced
significantly synapse lifetimes. Thus, synapse lifetime is highly sensitive to differences in sequence and is an important element in the comparison of sequences during the search for homology.

It has previously been reported that RecA-mediated strand exchange can traverse substitutional heterologies when using non-hydrolyzable ATP analog, ATPγS, instead of ATP (30). We found in our single-molecule experiments that ATPγS induced an irreversible aggregation of labeled oligomers on the non-labeled dsDNA anchored on the surface (data not shown) and individual synapses could not be resolved. Similar effects were previously observed in ensemble experiments in solution (14).

Synapse lifetimes are observed to increase when the synapse length is increased between 21 and 70 bp in the case of highly heterologous synapses. It is noteworthy that in the 21-long synapse case events of long enough duration to be detected were rare unless the concentration of pre-synaptic complexes was increased significantly. This suggests that heterologous synapses of this length are highly unstable.

Taken together, our data led us to suggest that facilitated/parallel search mechanisms may be essential to locate homology along chromosomes: the combination of long synapse lifetimes together with homology search by random collisions as commonly held (11,12) may take prohibitively long times. While sliding appears not to occur (3), the simultaneous formation of synapses between a RecA-covered ssDNA and a number of chromosomal segments, intersegmental transfer and inch-worming may be crucial mechanisms to overcome slow homology readout (4). It remains a challenge to study if these or other mechanisms are operative in the cell. One cannot, however, dismiss the possibility that in vivo, not all the genome is available for scanning during the search for homology, due to packing considerations and the association of genomic DNA with other proteins (31), and that homology may only be found in a subpopulation of the cells.

The present experiments pave the way for carrying out similar investigations into the behavior of eukaryotic recombinases. In contrast to RecA, homology search processes in eukaryotic organisms may be assisted not only by recombinases, but by additional factors (12,32,33), although evidence exist that a Rad51–ssDNA nucleofilament is sufficient to capture nucleosomal homology during double-strand break repair by recombination (34). Further studies will elucidate whether eukaryotic and prokaryotic organisms employ the same search strategies. Be that as it may, synapse lifetimes are a fundamental ingredient that must be taken into account in the construction of models of homology search. Long synapse lifetimes will provide strong constraints on such models.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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