



Direct Visualization of Horizontal Gene Transfer

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RecA family member, have been identified (12). Because DMC1B is expressed in vegetatively growing trophozoites and cysts whereas DMC1A is cyst-specific, we postulate that DMC1B may have a Rad51-like function and be involved in somatic DNA damage repair. Similarly, we suspect that although Mnd1 is involved in meiosis-specific recombination in other organisms, it may have a more general role in *Giardia*.

To determine when HOP1::GFP begins to be localized to the cyst nuclei, we monitored encystation over 24 hours with an antibody to cyst wall protein (CWP), which is transported via encystation-specific vesicles (ESVs) to the exterior of the cell and deposited to form the cyst wall (14). Three hours after induction of encystation, CWP was detected in ESVs in the cytoplasm (Fig. 4A), and the number of ESVs continued to increase through the first 12 hours (Fig. 4B). After 15 hours, cells were filled with ESVs, and HOP1::GFP was first detected in the two nuclei of the encysting trophozoite (Fig. 4C). By 24 hours (Fig. 4D), mature cysts with four nuclei and a defined wall were present, and HOP1::GFP was localized to all four nuclei. Thus, HOP1, which is involved in binding double-strand breaks during meiosis in yeast (15), is first detected in the nuclei in *Giardia* during encystation and persists indefinitely in cysts.

Karyogamy during encystation, if accompanied by homologous recombination and/or gene conversion driven by the HMGs, could provide a mechanism by which *Giardia* maintains low levels of allelic heterozygosity. This parasexual process, which we call diplomixis, appears to be unique to *Giardia*, although we predict that it occurs in other members of the order Diplomonadida. Unlike automixis, diplomixis is not accompanied by meiotic genome reduction and the subsequent fusion of gametes from the same parent, as is found in the sexual or parasexual life cycle of other organisms (16). It is also possible that rare meiotic events occur in the wild, as suggested by a recent *Giardia* population study (17). Understanding the functions of the HMGs and the behavior of chromosomes as nuclei fuse will be essential to test for the occurrence of homologous recombination. A deeper understanding of the roles of these genes, as well as others involved in karyogamy, may shed light on the evolution of meiosis and provide new targets for drug treatments.

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Supporting Online Material

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Materials and Methods

Figs. S1 and S2

References

Movies S1 to S6

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Direct Visualization of Horizontal Gene Transfer

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Conjugation allows bacteria to acquire genes for antibiotic resistance, novel virulence attributes, and alternative metabolic pathways. Using a fluorescent protein fusion, SeqA-YFP, we have visualized this process in real time and in single cells of *Escherichia coli*. We found that the F pilus mediates DNA transfer at considerable cell-to-cell distances. Integration of transferred DNA by recombination occurred in up to 96% of recipients; in the remaining cells, the transferred DNA was fully degraded by the RecBCD helicase/nuclease. The acquired integrated DNA was tracked through successive replication rounds and was found to occasionally split and segregate with different chromosomes, leading to the inheritance of different gene clusters within the cell lineage. The incidence of DNA splitting corresponds to about one crossover per cell generation.

Together with transformation and phage-mediated transduction, conjugation is a key mechanism for horizontal gene transfer in bacteria (1). The first evidence for sex by conjugation in *E. coli* was provided by Lederberg, who obtained prototrophic progeny by mixing two different auxotrophic parents (2). Since then, the phenomenon of horizontal gene transfer has been shown to be responsible for widespread transfer among bacterial populations of genes conferring antibiotic resistance, metabolic functions, and virulence determinants.

Conjugational DNA transfer is driven by the F plasmid unidirectionally from an F⁺ donor cell to an F⁻ recipient cell. The F plasmid contains all the genes required for conjugation (e.g., mediat-

ing the contact between donor and recipient cells) and for regulation of DNA mobilization and its unidirectional transfer (3). At low frequencies, the F plasmid can integrate into the chromosome of the host cell, giving rise to an Hfr (high frequency of recombination) strain (4). Chromosomal genes of the Hfr bacterium can be mobilized and transferred to a recipient. In some cases, F can excise from the chromosome of Hfr, creating an F' molecule that carries chromosomal genes as well as the conjugation genes (5). Both Hfr and F' can serve as DNA vehicles in horizontal gene transfer between bacteria.

The contact between mating cells is mediated by a tube-like structure known as the F pilus (3). DNA is transferred from the donor to the recip-

ient in single-stranded form and converted to duplex DNA by the synthesis of the complementary strand in the recipient cell. Once the conjugational transfer ceases, double-stranded donor DNA is either circularized (in the case of F' transfer) or, in the case of Hfr transfer, incorporated into the recipient chromosome by RecA-dependent homologous recombination or degraded by RecBCD exonuclease (3, 6).

Many aspects of the mechanism and consequences of conjugation remain unresolved, including the role of the F pilus in DNA transfer during conjugation, the fate of the transferred DNA, the global frequency of the horizontal gene transfer (versus the frequency of inheritance of individual genetic markers), and the pattern of inheritance of donor DNA present in the initial transconjugant cell. To address these questions, we have developed an experimental system that enables us to distinguish the transferred donor DNA from both donor and recipient DNA, and to visualize DNA transfer and recombination by means of fluorescence microscopy in real time,

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at the level of individual living cells. This tool allowed us to quantify the ongoing transfer of DNA during conjugation and to acquire time-lapse movies that follow the fate of the newly acquired DNA in individual cells through any number of cell divisions.

SeqA is a negative regulator of initiation of *E. coli* replication, with a high affinity for DNA that has been hemimethylated (i.e., only one strand in the duplex DNA is methylated) by Dam methylase at 5' GATC sequences (7–9). We mated Dam methylation-proficient donor (Hfr or F⁺) cells with methylation-deficient recipient cells, producing the SeqA-YFP (yellow fluorescent protein) fusion protein, which enabled us to specifically and permanently label only the transferred DNA. Once methylated donor DNA is transferred to the Dam⁻ recipient as a single strand and converted to a DNA duplex by the synthesis of the complementary nonmethylated strand, the acquired DNA is expected to remain permanently hemimethylated and therefore bound by SeqA-YFP. Indeed, we have previously shown the binding of SeqA to conjugally transferred DNA by immunofluorescence on fixed cells (10).

The SeqA-YFP translational fusion was constructed and placed in the chromosome under the control of the native *seqA* promoter, replacing the resident *seqA* gene (11). As expected, in Dam⁺ cells SeqA-YFP localizes to DNA replication forks to form compact foci (Fig. 1A). As DNA methylation lags considerably behind DNA synthesis, the newly replicated DNA is transiently hemimethylated DNA and therefore transiently bound by SeqA-YFP (Fig. 1A) (12–14). In contrast, in Dam⁻ cells SeqA-YFP fluorescence is diffuse (Fig. 1B). The number of SeqA-YFP foci in Dam⁺ cells varied from one to two in minimal medium, and up to eight in LB (Luria Bertani) medium, as expected from the occurrence of multiple replication forks in rich medium (13, 15). Foci were positioned at the half- or quarter-length positions in the cell. Both the number and position of SeqA-YFP foci corresponded to those previously reported for SeqA (13, 14), which suggests that SeqA-YFP is functional in terms of DNA binding specificity.

To investigate the localization of SeqA-YFP protein during conjugation, we placed a mixture of *dam*⁺ donors and *dam*⁻ *seqA-yfp* recipients on a nutrient-containing agarose slab in a sealed-cavity microscope slide and observed it with time-lapse fluorescence microscopy. About 5 min after mixing the parental cells, distinct SeqA-YFP foci started to appear in recipient cells (Fig. 1, D and E). During the next 30 to 40 min, almost all recipients in the vicinity of donors acquired intense fluorescent foci of the SeqA-YFP due to the sequestration of previously diffused SeqA-YFP by the transferred DNA (Fig. 1E and movie S1). A single recipient cell could receive DNA more than once, as shown by the presence of independent, well-separated foci arising at different time points at different positions in the cell (fig. S1 and movie S1).

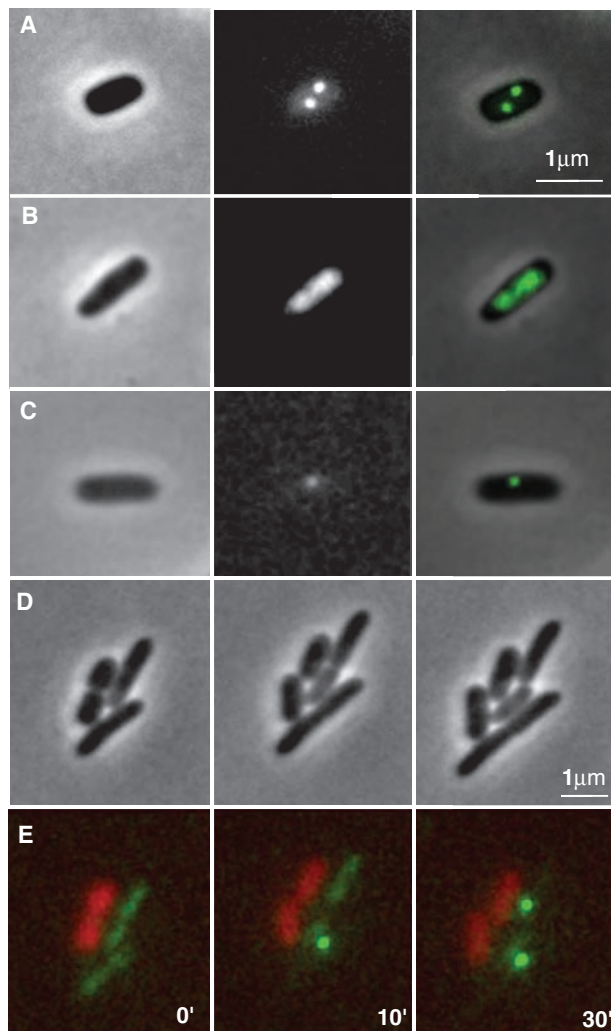
As expected, when *dam*⁻ donors were used in conjugation, no SeqA-YFP foci were observed in the *dam*⁻ recipients. Also, no SeqA-YFP foci were seen when donors lacking the structural pilus protein (TraA) were used and were therefore transfer-deficient (16). These results confirm that SeqA-YFP foci formation reflects the entry of a methylated donor DNA strand into the *dam*⁻ recipient during conjugation. Although the conjugal transfer of the extrachromosomal R plasmid was previously observed by fluorescence of a specific locus on the plasmid (17), our system allows visualization of transfer of any extrachromosomal or chromosomal DNA in a sequence-independent fashion.

To visualize the expression of conjugally transferred DNA in the recipient cells, we constructed the ANB35 donor strain of *E. coli*, which carries the tetracycline (Tet) promoter-driven monomeric red fluorescent protein (*mrfp1*) gene (18) and *tetR* (repressor) gene (11). In the absence of induction, donor cells exhibit weak red fluorescence because of the leakiness of the Tet promoter (movie S2). When ANB35 donors were mated with *dam*⁻ *seqA-yfp* recipients, we first observed the appearance of the SeqA-YFP

focus, denoting DNA transfer, followed by the red fluorescence in the recipient cell, showing *mrfp1* gene expression, about 2 hours after the transfer of the *mrfp1* gene. This expression of the *mrfp1* gene was detected in 25 of 307 mated recipients (8.1%). Not all successful transfers would have included the *mrfp1* gene, because conjugational transfer ceases at random points, and the probability of transferring a specific marker decreases exponentially with its distance from origin of transfer *oriT* (19). The recipient cell eventually became intensely red relative to the fluorescence of the donor (movie S2). Upon entry of the donor DNA into the recipient cell lacking the TetR repressor, the transcription of *mrfp1* occurred before sufficient TetR repressor was expressed and functional. The recipient cell therefore became intensely red before repression started. This phenomenon is known as induction of gene expression by conjugation, or zygotic induction (20).

The F pilus is a tubular organelle, extruding from the surface of the donor cell, that is required for conjugation (21). Some authors have argued that the F pilus serves only to bring the mating cells in direct contact, and that DNA is trans-

Fig. 1. Localization of SeqA-YFP in DNA replication and in conjugation. (A) Dam-proficient cells. (B) Dam-deficient cells. (C) Recipient with SeqA-YFP focus after conjugation with F donor. Left, phase contrast image; center, fluorescence image; right, overlay between the phase contrast image and the fluorescence image represented in green. (D and E) Real-time conjugation [(D), phase contrast; (E), fluorescence overlay] of donors (red cells) with recipient (green cells) as followed by time-lapse microscopy at 0, 10, and 30 min after plating on nutrient-agarose cavity slide.



ferred subsequently through a pore in the membrane (22, 23). Others have shown that DNA can be transferred from the donor to the recipient even when mating cells were apparently physically separated (24). In three experiments, we observed SeqA-YFP foci appearing in 43 of 753 recipients that were not in direct cell wall contact; the maximal observed distance to the closest donor was 12 μm (Fig. 2 and movie S3). No distant transfer of DNA was observed in the presence of 0.01% SDS (which depolymerizes F pili) or when using the transfer-deficient *traA*⁻ donors (which lack the pili). Clearly, the appearance of distant SeqA-YFP foci depends on conjugation and on the presence of F pili. Thus, our results offer evidence supporting the idea that the F pilus, in addition to establishing the contact between mating cells, serves as a channel for single-stranded DNA transfer during conjugation.

In conjugations with recombination-deficient recipients (*recA*⁻), transferred DNA is known to be degraded by the RecBCD exonuclease (6, 25). RecBCD is a highly processive enzyme degrading single- and double-stranded linear DNA (26, 27). The kinetics of degradation of conjugally transferred DNA have so far only been measured in large cell populations (6, 25) and not at the level of single cells.

In our experiments we found that the SeqA-YFP focus fluorescence intensity in the recipient is proportional to the time of active conjugation, that is, to the length of transferred DNA (fig. S2). The gradual disappearance of the fluorescence of

SeqA-YFP foci that we observed in recombination-deficient cells (movie S4) probably reflected DNA degradation mediated by the RecBCD helicase/nuclease complex. Indeed, Hfr DNA transferred into *recA*⁻*recD*⁻ recipients was largely left unchanged, and no degradation was seen in 459 of 471 recipients [i.e., $97 \pm 0.24\%$ (SEM) in three experiments] 4 hours after interruption of mating. By contrast, in *recA*⁻ recipients, transferred DNA was rapidly broken down in 313 of 343 recipients (i.e., $91.3 \pm 4.1\%$ in three experiments). DNA degradation over time in most cases was linear, yet individual cells exhibited variable degradation rates (fig. S3A). Occasionally (for 9 of 135 foci in two experiments) we saw a delay in the onset of degradation (fig. S3B). Interestingly, a subpopulation of SeqA-YFP foci ($8.7 \pm 4.1\%$ for 343 tracked foci) exhibited an initial decrease in fluorescence due to degradation of DNA but eventually reached a stable fluorescence level, even 4 hours after the interruption of mating (fig. S3C). This contrasts with the findings that no genuine genetic recombinants can be obtained in crosses with *recA*⁻ recipients (28). In these cases, the nonreplicative Hfr DNA somehow must be end-protected—for example, by circularization via microhomologies, palindromic end structures, or protein binding.

The fate of transferred DNA in recombination-proficient (*recA*⁺) recipients was followed after interruption of mating (11). In *E. coli*, all free linear DNA is rapidly degraded by the RecBCD exonuclease (27); therefore, SeqA-YFP foci that remain stable in *recA*⁺ recipients for at least 4

hours were considered as recombined DNA. Transferred DNA followed two fates in the *recA*⁺ recipients: In 11 of 336 mated recipients ($3.3 \pm 0.83\%$ in three experiments), SeqA-YFP foci disappeared, indicating that the donor DNA was degraded (fig. S4). However, in the majority ($96.7 \pm 0.83\%$), the fluorescence intensity remained constant or slowly increased for the higher-intensity foci. This is coherent with the binding of newly synthesized SeqA-YFP to available GATC sequences of large conjugated fragments, whereas small fragments are readily saturated by available SeqA-YFP (fig. S4).

This direct, physical, measure of recombination shows substantially higher recombination frequencies than those measured genetically as the acquisition of specific genetic markers (ranging from 10 to 30% depending on specific genetic marker, donor strain, and mating conditions) (29, 30). Our results show that conjugational recombination is an extremely efficient process when donors and recipients are essentially genetically identical strains.

Remarkably, SeqA-YFP foci frequently split within single cell lineages (Fig. 3 and movie S5). The sum of fluorescence intensities of newly formed foci was not statistically different from the intensity of the ancestral focus (*t* test, $P = 0.62$, $n = 20$); that is, the splitting events were conservative, showing no evidence of significant DNA degradation (Fig. 3). Most likely, the splitting of SeqA-YFP foci reflects the physical breakage of the initial transferred DNA strand into two DNA fragments (11). This is reminiscent of reciprocal crossovers (sister chromatid exchange) because the splitting frequency is proportional to the DNA size and occurs only in recombination-proficient cells (11). Extrapolation of the splitting frequency of the acquired DNA to the entire genome suggests that there are about 1.4 crossovers (sister or cousin molecule exchange) per cell generation.

This work allowed us to visualize and quantify the DNA of any sequence as it is being transferred from one individual cell to another, and to watch its stable genomic acquisition via genetic recombination (horizontal gene transfer) in real time. The observations also implicated F pilus-mediated conjugation at cell-to-cell distances of up to 12 μm and permitted estimation of the frequency of intragenomic crossover events, or sister-chromatid exchanges. This experimental system can now be applied to monitor horizontal gene transfer by indefinitely following the fate of DNA acquired in intra- and inter-species crosses.

Fig. 2. Representative example of conjugational DNA transfer without visible cell contact between donors and recipients. Recipients and donors were plated on the nutrient-agarose slab without previously being in contact and were observed by time-lapse fluorescence microscopy. (A) Phase contrast images. (B) Overlay of fluorescence images of recipients (green cells) and donors (red cells) at 0, 5, and 10 min after plating on LB-agarose in a cavity slide.

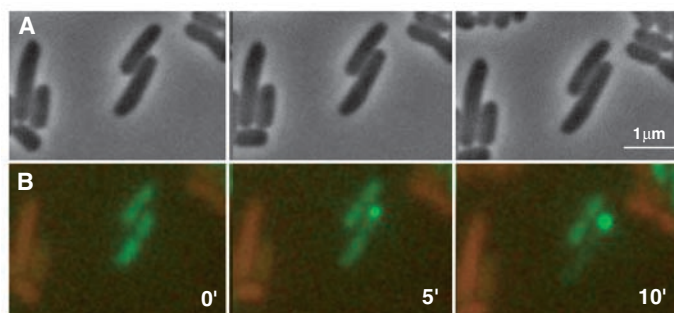
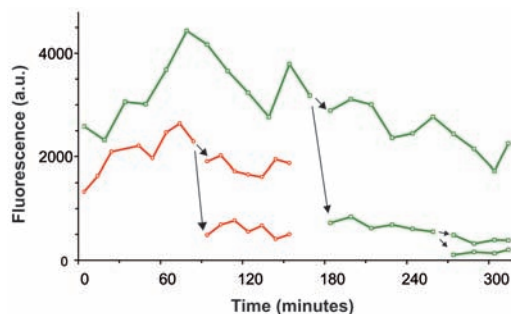


Fig. 3. Two representative cases of the SeqA-YFP focus splitting, differentiated by color. SeqA-YFP foci were acquired during 30 min of mating on a nitrocellulose filter with a recombination-proficient recipient. Time 0 represents the interruption of mating by nalidixic acid and transfer under the microscope for observation. Times of splitting are indicated by arrows; a.u., arbitrary units. Fluorescence 0 is the average background SeqA-YFP fluorescence level of nonconjugating recipients. Each point represents an integral (total) fluorescence of SeqA-YFP focus.



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Materials and Methods

SOM Text

Figs. S1 to S4

Movies S1 to S5

References

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Neurokinin 1 Receptor Antagonism as a Possible Therapy for Alcoholism

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Alcohol dependence is a major public health challenge in need of new treatments. As alcoholism evolves, stress systems in the brain play an increasing role in motivating continued alcohol use and relapse. We investigated the role of the neurokinin 1 receptor (NK1R), a mediator of behavioral stress responses, in alcohol dependence and treatment. In preclinical studies, mice genetically deficient in NK1R showed a marked decrease in voluntary alcohol consumption and had an increased sensitivity to the sedative effects of alcohol. In a randomized controlled experimental study, we treated recently detoxified alcoholic inpatients with an NK1R antagonist (LY686017; $n = 25$) or placebo ($n = 25$). LY686017 suppressed spontaneous alcohol cravings, improved overall well-being, blunted cravings induced by a challenge procedure, and attenuated concomitant cortisol responses. Brain functional magnetic resonance imaging responses to affective stimuli likewise suggested beneficial LY686017 effects. Thus, as assessed by these surrogate markers of efficacy, NK1R antagonism warrants further investigation as a treatment in alcoholism.

Alcohol use accounts for 4% of global disease burden (1). Alcohol dependence, or alcoholism, is characterized by a chronic relapsing course, in which alcohol-associated cues and stress are known relapse triggers (2–6). Recent research suggests that neural systems mediating behavioral stress responses may offer useful targets for pharmacotherapy of alcohol-

ism. In animal models, excessive alcohol consumption that results from a history of alcohol dependence is accompanied by increased behavioral sensitivity to stress (7). Up-regulated corticotropin-releasing hormone (CRH) signaling in extrahypothalamic brain sites contributes to these dependence-induced changes, but other stress-related neurotransmitters may also play a role.

One such neurotransmitter is substance P (SP), which together with its preferred neurokinin 1 receptor (NK1R) is highly expressed in brain areas involved in stress responses and drug reward, including the hypothalamus, amygdala, and nucleus accumbens. In rodents, psychological stressors induce release of SP in the amygdala, whereas genetic deletion or pharmacological blockade of NK1R inhibits the associated behavioral responses (8). Furthermore,

genetic deletion of NK1Rs causes a loss of conditioned place preference for opiates and opiate self-administration (9, 10). In humans, the NK1 antagonist GR205171 reduces symptoms of social anxiety and suppresses brain responses to the Trier Social Stress Test (TSST) (11). Together, these findings suggest that blockade of NK1Rs might modulate stress- and reward-related processes of importance for excessive alcohol use and relapse. To our knowledge, no data are presently available to address this hypothesis.

We first explored preclinically whether inactivation of NK1R might modulate stress- and reward-related processes that impact alcohol use. We chose a genetic inactivation strategy, because available NK1R antagonists have limited activity in rats and mice, because of insufficient NK1R amino acid homology between humans and these rodent species (8). We evaluated NK1R null-mutant mice for voluntary alcohol consumption, alcohol sensitivity, and alcohol metabolism (12). NK1R null mice (13) were back-crossed into a C57BL/6 background for 10 generations to ensure that there was adequate voluntary alcohol consumption in control animals (14). We used a two-bottle free-choice model with increasing alcohol concentration, and alcohol was continuously available. Wild-type littermates (+/+) ultimately consumed in excess of 10 g alcohol/kg of body weight per day at the end of an escalation procedure in which alcohol concentration was gradually increased from 3 to 15% over 60 days.

Alcohol consumption by NK1R^{-/-} mice was markedly lower than that by wild-type controls (Fig. 1A). The difference was most prominent at higher alcohol concentrations, at which consumption motivated by pharmacological alcohol effects dominates over intake for taste, calories, or other nonpharmacological effects (14). Alcohol consumption by heterozygous (+/-) mice was similar to wild-type controls, highlighting the necessity for near-complete inactivation of NK1Rs to suppress alcohol consumption. Sev-

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