Evolution of the mating type gene pair and multiple sexes in *Tetrahymena*

Highlights

- The two *Tetrahymena* mating type proteins evolved from a common ancestor.
- Successive replacement waves generated the current diversity of mating type proteins.
- Well defined segments of both mating type proteins show differential lineage sorting.
- Perpetual selfer, *T. shanghaiensis*, has a heterospecific mating type gene pair.

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Evolution of the mating type gene pair and multiple sexes in *Tetrahymena*

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**Summary**

The multiple mating type system of the Ciliate *Tetrahymena thermophila* is a self/non-self recognition system, whose specificity resides in a head-to-head, functionally distinct pair of genes, *MTA* and *MTB*. We have now sequenced and analyzed these mating type genes in nine additional *Tetrahymena* species. We conclude that *MTA* and *MTB* are derived from a common ancestral gene and have co-evolved for at least ~150 Myr. We show that *T. shanghaiensis*, a perpetual selfer (unisexual) species, has a single mating type gene pair, whose *MTA* and *MTB* genes likely have different mating type specificity. We document the recent replacement of a complete different set of mating type specificities for another, illustrating how quickly this can happen. We discuss how varying conditions of reproductive stress could result in evolutionary co-adaptations of *MTA* and *MTB* genes and changes in mating type determination mechanisms.

**Introduction**

Sex is an evolutionary conserved process among organisms, including the Ciliated Protozoa (= Ciliates), a unicellular eukaryotic phylum (Bachtrog et al., 2014). Mating type systems generally ensure sexual self-incompatibility and promote outbreeding. Most eukaryotic species have two mating types or sexes and thus a binary mating system. However, systems with more than two mating types exist in some groups such as ciliates and mushrooms (Phadke and Zufall, 2010; Kues, 2015). Ciliate mating systems are very diverse: they vary in such features as number of mating types, mechanism of mating type determination (MTD), and molecular nature of the mating type proteins. This suggests that many fundamental changes in mating type biology have independently evolved in the major Ciliate clades (Phadke and Zufall, 2010). The *Tetrahymena* genus of ciliates is thus an excellent model system for studying multiple mating type systems and their evolution.

As in other ciliates, cells of most *Tetrahymena* species possess two kinds of nuclei: a diploid, silent germline nucleus (the micronucleus or MIC) and a polyploid, highly expressed somatic nucleus (the macronucleus or MAC). The *Tetrahymena* life cycle consists of two stages: asexual reproduction by binary fission when food is abundant and conjugation triggered by starvation (reviewed in Orias et al., 2011; Orias et al., 2017). Key life cycle features of genetic significance, illustrated in Figure 1, are:

- Only one of the four MIC meiotic products is retained in each conjugant.
- Reciprocal fertilization generates genetically identical, diploid zygote nuclei in each conjugant.
- The zygote nucleus in each conjugant divides twice mitotically, two products are the new MICs, while the other two differentiated into the new MACs.
- During MAC differentiation the five MIC chromosomes undergo programmed site-specific fragmentation, resulting in 180 MAC chromosomes. These acentromeric chromosomes are then amplified (~45 G1 copies).
- The two exconjugant cells from a pair divide, resulting in four cells with genetically identical MICs but independently differentiated MACs, called “karyonides”.

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Phenotypic assortment: when asexual multiplication resumes after conjugation, amitotic division of the polyploid MAC results in the random distribution of daughter chromosome copies at every fission. This allows segregation at all heterozygous loci present in a newly differentiated MAC and ultimately generates whole-genome homozygous MACs.

Sexual progeny are initially sexually immature; in *T. thermophila* they must undergo 40–70 fissions before they reach sexual maturity and can mate again. The diversity of *Tetrahymena* mating type systems was described in detail in a series of papers by Nanney, Elliott, and their collaborators, beginning in the early 1950s (reviewed in (Orias, 1981; Orias et al., 2017)). Among sexual species in this genus, the number of known mating types per species ranges from three to nine. Certain rare species (e.g., *T. shanghaiensis*) are unisexual (selfers), meaning that sexually mature cells in a clonal population mate with one another upon starvation (Chen et al., 1982; Simon et al., 2009) and all their progeny are selfers. Cells of other species (e.g., *T. pyriformis* and *T. vorax*) have lost their MIC (i.e. the germline nucleus) and thus can only reproduce asexually (Gruchy, 1955; Doerder, 2014).

The best characterized species belong to either the “Australis” or the “Borealis” clades, which diverged ~150 Myr ago (Xiong et al., 2019). Earlier studies suggested that *Tetrahymena* MTD patterns can also be classified into two categories that co-branch with the phylogenetic tree. Investigated species in the “Australis” clade exhibit “synclonal MTD”; where the four genetically identical karyonides of a mating pair (the synclone) express the same mating type with a Mendelian inheritance pattern when they reach sexual maturity (Figure S1A). In *T. pigmentosa*, for example, mating types are controlled by three alleles of a single mat locus that show “peck-order” dominance (Simon, 1980). In contrast, investigated species in the “Borealis” clade show “karyonidal MTD”, where mating type is randomly and independently determined in each new MAC. This results in four genetically identical karyonides which often express different mating types (Figure S1B).

The molecular basis of mating type specificity has only been investigated in *T. thermophila* (“Borealis” clade). Mating type is determined by a mating type gene pair (mtGP), a head-to-head arrangement of...
two mating type genes (MTA and MTB) (Cervantes et al., 2013) (Figure 2A). In this study, we investigated the molecular evolution of Tetrahymena mtGPs in a phylogenetically wide range of Tetrahymena species, including two asexual species (pyriformis and vorax), a unisexual obligatory selfer species (shanghaiensis), and the species furthest removed from the "Australis" and "Borealis" clades (paravorax) (Figure 2B). We provide evidence for the evolution of all Tetrahymena mating type proteins from an ancient "Furin-like repeat" protein family, for the coevolution of MTA and MTB genes, for the evolution of a heterotypic MTA-MTB gene pair leading to perpetual selfing, and for the recent replacement of one multiple mating type system with another within a subgroup of the genus Tetrahymena.

**Results**

**Mating type gene pair homologs exist in all but the most distantly related Tetrahymena species examined**

In *T. thermophila*, the MTA and MTB genes shares some similar features but have completely different sequences. The sequence of the terminal exons is highly conserved between MTA genes encoding different mating types (Figure 2A) (nucleotide identity >0.99); the same is true among the MTB alleles. In contrast, the remainder of the MTA and MTB genes and the intergenic region are mating type-specific (nucleotide identity <0.6). The 3′-terminal exons of the MTA and MTB genes both encode five predicted transmembrane helices and a cysteine-rich "Furin-like repeat" domain. These features are diagrammed in Figure 2A; see also (Cervantes et al., 2013; Orias et al., 2017).

We looked for homologs of the *T. thermophila*MAC mating type genes in nine additional species spanning the ~300 Myr old Tetrahymena genus (Xiong et al., 2019) (Table 1). Six species (thermophila, malaccensis, pyriformis, vorax, borealis and canadensis) are in the "Borealis" clade, three (shanghaiensis, americanis, and pigmentosa) are in the "Australis" clade, and one (paravorax) diverged from both clades at the
base of the Tetrahymena genus (Figure 2B). T. pyriformis and vorax cells never mate, they lack an MIC and only reproduce asexually. And to our knowledge, sexual reproduction has not been observed in paravorax.

Altogether, we investigated mating type genes in 19 Tetrahymena strains (Table 2). We found mating type gene homologs in all sexual species, as well as in three asexual species (pyriformis, vorax and paravorax) and in the unisexual strain, shanghaiensis (a “selfer” species, in which starvation of sexual mature cells triggers intraclonal mating). Within each sexual species, we verified that mating only occurs between starved cells of different mating types; no mating was observed between starved cells from different species (for experimental details see Transparent methods).

Our searches revealed a single, mating type-specific mtGP with MTA and MTB homologs in head-to-head orientation for each mating type of every species, with the sole exception of the T. paravorax strain, which has a truncated MTA gene (named MTAL for MTA-like) and lacks an MTB homolog (Figures 2B and S2). Additionally, we determined that all mtGPs of species with sequenced genomes (Xiong et al., 2019), with the single exception of the asexual T. vorax mtGP, have syntenic chromosomal locations (Figure S3).

It has long been known that starvation conditions are required for conjugation in Tetrahymena. Consistent with this, the T. thermophila MTA and MTB genes are highly expressed during starvation, but are essentially silent during vegetative growth (Cervantes et al., 2013). To investigate whether mtGPs of other Tetrahymena species might function similarly in mating, their expression levels were measured during growth and starvation (Figure 2C). In the seven sexual species (including the “selfer” species, T. shanghaiensis), the expression pattern of the mating type genes is identical to that of T. thermophila, consistent with conservation of mtGP function in mating. In the asexual species (pyriformis, vorax, and paravorax), the mating type genes are not induced by starvation, suggesting that they are no longer functional in mating.

**Tetrahymena MTA and MTB genes are likely derived from a common ancestral gene**

All 37 MTA and MTB genes reported in the previous section belong to the same gene superfamily which has a conserved cysteine-rich “Furin-like repeat” domain (Figures 3A and S2). Alignment of the “Furin-like repeat” domains in all MTA and MTB homologs shows that all 14 cysteine residues are highly conserved with one another, although four cysteines are missing in MTAL (Figure 3A). Since cysteine residues play many important roles, such as forming covalent disulfide bonds with each other (Sela and Lifson, 1959; Thornton, 1981), the conservation of these residues may be essential to the secondary structure of the mating type proteins and their function.

The full length Tetrahymena MTA and MTB genes investigated here contain 5-8 introns. In the sexual species, four introns (#3, #5, #6, and #7) show total conservation of phase and approximate location in the 32 sequenced MTA and MTB genes (Figure 3B and Table S1, details in Figures S2 and S4). Intron #7 precedes the 3’-terminal exon, which encodes the “Furin-like repeat” domain and transmembrane helices of the
MTA, MTAL, and MTB genes. A fifth intron (#2) is conserved among the MTA and MTB genes of all mtGPs, with the exception of *T. canadensis*, which has just one sequenced mtGP (*MTAX*).

The conservation of intron location, in both MTA and MTB, is particularly striking within subgroup alignments (Figure S4) that include at least two species for each subgroup. Astonishingly, 27 (of 38) introns occur at identical codon locations within their gene. In another seven cases, the introns are located at an adjacent codon. In the four remaining cases, the intron is located a few codons further away, but there are clearly insertions/deletions of one or more codons in the immediate neighborhood. The only outlier is intron #2 in the “Bor-Can” MTAs, which has changed phase in *T. borealis MTAZ* and is missing in *T. canadensis* *MTAX*.

The absolute conservation of the “Furin-like repeat” domain, together with the high degree of conservation of intron location and phase in such a large (~1,500 aa) protein, along with chromosomal synteny, are strong evidence that all MTA and MTB genes were derived from a common ancestral gene which existed prior to the divergence of the “Australis” and “Borealis” clades in *Tetrahymena*, ~150 Myr ago (Xiong et al., 2019).

To confirm the evolutionary relationship between MTA and MTB proteins, we generated phylogenetic trees of these proteins from the sexual species. For each protein, we made separate trees for the entire protein (Figure S5), for the C-terminal, transmembrane exon (distal ~1/3 of the protein), and for the rest of the protein (proximal ~2/3 of the protein (Figures 4A and 4B, respectively)). The results show that all three trees have two main branches, such that all MTA proteins fall cleanly into one branch, while all MTB proteins fall cleanly into the other branch. This is consistent with the nearly complete lack of overall sequence similarity observed between the MTA and MTB proteins when they are co-aligned (Figure S2).

Thus the trees support the conclusion that the MTA and MTB genes diverged structurally and functionally from a common ancestral gene early in *Tetrahymena* evolution, prior to the divergence of the “Australis” and “Borealis” clades.

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**Table 2. Relevant strain and mating type information on the *Tetrahymena* strains used in this work**

| Species      | Strain ID*  | Mating typeb |
|--------------|-------------|--------------|
| *thermophila*| SD01580     | II           |
| *thermophila*| SD01653     | III          |
| *thermophila*| SD01582     | IV           |
| *thermophila*| SD01656     | V            |
| *thermophila*| SD01584     | VI           |
| *thermophila*| SD01585     | VII          |
| *Malaccensis*| SD01608     | X            |
| *Pyriformis* | SD00707     | NA           |
| *Vorax*      | SD30421     | NA           |
| *borealis*   | SD01609     | X            |
| *borealis*   | SD19502     | Y            |
| *borealis*   | SD19803     | Z            |
| *canadensis* | SD30770     | X            |
| *shanghaiensis* | SD205039  | Selfer      |
| *pigmentosa* | SD19481     | III          |
| *pigmentosa* | SD20427     | I            |
| *americanis* | SD21194     | X            |
| *americanis* | SD21244     | Y            |
| *paravorax*  | SD205177    | N/D          |

N/A: not applicable; these strains lack a micronucleus and are asexual. N/D: not determined.

* Tetrahymena Stock Center ID numbers.

bWhere the relationship to previously published mating types is undetermined, we have used capital letters X, Y, and Z to avoid confusing the literature.
Figure 3. Highly conserved features of Tetrahymena MTA and MTB genes: FLR repeats and introns

(A) Sequence alignment of the “Furin-like repeat” domains. Cysteine sites are conserved in MTA and MTB proteins; note that four conserved cysteines (darker background) are missing in T. paravorax MTA. Pink letters: partially conserved amino acids. P.tet: Paramecium tetraurelia mtA gene.

(B) Conservation of intron location and phase among MTA and MTB genes. Thick green lines, MTA exons; thick blue lines, MTB exons; thick dark green or dark blue lines, terminal exons. Colored dots, introns: blue, phase 0, inserted between two codons; green, phase 1, inserted between the first and second codon nucleotide; red, phase 2, inserted between the second and third codon nucleotide. Mating type allele shown, for those species with multiple sequenced mating types: T. thermophila: mt II; T. borealis: mt X; T. pigmentosa: mt III; T. americanus: mt X. Green triangle, phase 1 intron that exists in
Interestingly, conserved intron #4 is absent from the MTA genes of all sequenced mating types in the “The-Mal” subgroup. These genes all have another intron (intron #6.5), located between conserved introns #6 and #7. Unexpectedly, the *T. borealis* MTA gene contains an intron at exactly this location (#6.5) and phase (Figure 3B, green triangle and Table S1) but also has intron #4. Conceivably, the MTA genes of the “The-Mal” subgroup and the *T. borealis* MTA gene may share a recent common ancestor.

*Paramecium*, like *Tetrahymena*, belongs to the Ciliate class Oligohymenophorea. The two genera are estimated to have diverged from one another nearly a billion years ago (Xiong et al., 2019). The mating protein (mtA) in *Paramecium tetraurelia* is also a member of the superfamily of genes having a terminal exon containing “Furin-like repeat” domains, which conserves all the cysteines found in *Tetrahymena* and the transmembrane helices (Singh et al., 2014). However, the *Paramecium* gene has only three introns, all at different locations than in the *Tetrahymena* genes. Thus, the *Paramecium* and *Tetrahymena* mating type genes likely had a common ancestor but have undergone extensive independent evolution.

**The *Tetrahymena* mating type proteins exhibit a special type of incomplete lineage sorting**

The topology of the individual MTA and MTB branches of the phylogenetic tree of the two entire proteins (Figure S5) does not exactly match the topology of the species tree (Figure 2B). This finding represents an example of “incomplete lineage sorting”. The discrepancy is limited to the MTA and MTB proteins of *T. malaccensis*, *borealis*, *canadensis*, and *shanghaiensis*.

Interestingly, the topologies of the branches of both the MTA and MTB proteins in the phylogenetic tree for the distal third (C-terminal exon) (Figure 4A) are almost identical to the topology of the species tree (Figure 2B). Indeed, essentially all of the incomplete lineage sorting seen for the whole proteins is accounted for by that in the proximal (N-terminal) roughly two thirds of each protein (Figure 4B). In this context, it is important to note the very different functions of the two segments of the mating type proteins. The distal third (encoded by the C-terminal exon) includes the only predicted intracellular segment of the protein and can thus be inferred to be involved in mating type-non-specific interactions with cell machinery required, for example, for the structural remodeling of the cell in preparation for mating: re-shaping and de-ciliation of the anterior ventral surface where two cells will form a temporary junction lasting many hours (“tip transformation” (Wolfe and Grimes, 1979)). This is the protein segment that has evolved at a rate commensurate with that of other conserved, mating type-unrelated cell proteins. On the other hand, the proximal two thirds of the mating type proteins are predicted to be extracellular and to be the main site of the mating type-specific (positive and negative) interactions that allow the self vs. non-self recognition required to initiate or inhibit mating between two cells. Given that most *Tetrahymena* species possess multiple mating type systems, and that speciation has been accompanied by conservation of some mt protein specificities and evolutionary radiation of others (as described in subsequent sections), the incomplete lineage sorting observed for this protein segment becomes readily understandable. This clear distinction between nearly complete lineage sorting in the distal third and significantly incomplete lineage sorting in the proximal two thirds, seen for both of the two proteins, represents an example of what could be called “composite lineage sorting”.

The individual examples of incomplete lineage sorting detected in this proximal two thirds of the MTA and MTB proteins are addressed in more detail below:

1) *T. malaccensis* MTA and MTB proteins co-branch with *T. thermophila* MTA4 and MTB4, respectively. The two species are very closely related (Figure 2B). This was already reported (Cervantes et al., 2013) and interpreted to mean that these two mtGPs recently evolved from the same mtGP in a common ancestor of the “The-Mal” subgroup.

2) *T. borealis* MTA and MTBZ proteins co-branch with *T. canadensis* MTA and MTBX, respectively, which has a proposed analogous explanation to the previous case. These two species are among the most closely related *Tetrahymena* species pairs known. Their SSUrRNA genes are identical, and their COX1 barcodes show only 4.4% polymorphisms (data not shown). Four percent is the...
COX1 threshold that best corresponds to the ultimate criterion of the *Tetrahymena* species difference, the failure to mate (Doerder, 2019).

The two other discrepancies are more intriguing.

3) The co-branching of *T. borealis* mating type Y with all the *T. thermophila* and *T. malaccensis* mating types is unexpected because these species are in two different phylogenetic subgroups ("Bor-Can" and "The-Mal", respectively). This finding suggests that this particular mtGP has been retained and has changed relatively little since the divergence of the two subgroups. Further supporting this hypothesis, and as described in an earlier section, the *T. borealis* MTAY gene contains an intron at

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**Figure 4. Phylogenetic tree of mating type proteins**

(A) Protein phylogenetic tree based on the C-terminal exon (distal-third). (B) Phylogenetic tree based on the rest of sequence (proximal-two-thirds). The best-fit models were calculated by ProtTest (version 3.4.2) (Darriba et al., 2011). (A) is under JTT + I + G + F model; (B) is under VT + I + G + F model. Numbers at each node, bootstrap values (1000 replicas). Branch length, number of base substitutions per site. Horizontal dashed lines: boundaries between the species clades/subgroups. Red boxes: deviations from the species tree. The MTA and MTB branches are shown opposite to one another to facilitate comparisons between them.
exactly the same location and phase as the additional intron (#6.5) found in all the sequenced MTA genes in the “The-Mal” subgroup and nowhere else (Figure 3B).

4) The last anomaly is the co-branching of the MTA and MTB proteins of the perpetual selfer *T. shanghaiensis* with *T. pigmentosa* MTA3 and MTB3, respectively; this finding is addressed in the *T. shanghaiensis* section, further below.

A better understanding of the molecular basis of these cases of incomplete lineage sorting will require sequencing additional mating type genes of these and other species in the “Bor-Can” subgroup, as well as additional knowledge of the sequence and organization of the genes encoding these proteins in the germline (micronuclear) mat locus. The latter information is currently available only for *T. thermophila* (Cervantes et al., 2013).

**MTA and MTB genes have coevolved within the different Tetrahymena phylogenetic subgroups**

*T. thermophila* MTA and MTB gene products have non-redundant functions required for mating (Cervantes et al., 2013). Intriguingly, when we compared the two mating type genes in species belonging to different *Tetrahymena* phylogenetic groups, we noticed several cases where recent evolutionary changes in the MTA gene have been mirrored by corresponding changes in the MTB gene. For example, in the “The-Mal” subgroup, the GC content of 3′-terminal exons of both genes is significantly higher than that of other regions of the mtGP (Figures 5A and 5B). This difference is not observed in the other subgroups.

More striking evidence of coevolution is the length of mating type specific region. Previous work in *T. thermophila* (Cervantes et al., 2013) had revealed that the 3′-terminal exons in MTA genes, comprising about 1/3 of each gene, are highly conserved among alleles for the different mating types, while the rest of each gene is mating type-specific; the same is true for MTB alleles (Figure 5C, *T. the*). We examined different mtGPs in species of the “Bor-Can” subgroup (*T. borealis*) and the “Australis” clade (*T. americanis* and *T. pigmentosa*) to see if they also shared a distinct sequence conservation boundary. Sequence conservation plots (Figure 5C) of the MTA and MTB genes of these species show that none of their mating type genes have an abrupt conservation boundary at conserved intron #7. Instead, in the two species of the “Australis” clade, the conserved regions of the MTA and MTB genes are about twice as long as in *T. thermophila* (Figure 5C, *T. ame* and *T. pig*), so that only the 5′-terminal ~1/3 of each gene is unique for each mating type. In further contrast, for the three sequenced mating types of *T. borealis*, essentially the entire length of the MTA and MTB genes is mating type-specific (Figure 5C, *T. bor*).

In an attempt to shed more light on the question of MTA and MTB protein coevolution, we also did an amino acid usage analysis of all the MTA and MTB proteins in the sexual species (Data S1, Figure S6). The results showed some regularities but did not provide clear conclusions. A rigorous answer to this question will likely have to wait for additional experimental investigations and knowledge of the 3D structure of these proteins.

**T. shanghaiensis** cells have one mtGP allele and mate with one another

In contrast to the multiple mating type systems of other *Tetrahymena* species, *T. shanghaiensis* has been reported to be a selfer species by Chen et al. (1982) and Feng et al., 1988, in which sexually mature cells within every *T. shanghaiensis* clone mate with one another upon starvation. By DNA sequencing, we identified a single mtGP in the *T. shanghaiensis* MAC genome, whose MTA and MTB genes are homologous with the respective mating type genes of the other *Tetrahymena* species investigated here (Figures S2 and S4). Furthermore, the *T. shanghaiensis* MTA and MTB homologs have identical expression profiles to those of all the other sexual *Tetrahymena* species during growth and starvation (Figure 2C).

To verify the previously reported observation that the non-assorting selfing trait is transmitted to sexual progeny, we did RNA-Seq experiments and de novo sequence assembly on two additional starved *T. shanghaiensis* populations, obtained as sexual progeny of independent selfing populations (i.e. biological replicates) (Figure S7, frames 5 and 7, highlighted with red stars). Cells in both populations contained transcripts from the MTA and MTB genes of the only previously detected single, genomic mtGP.
conclusion that the basis for the non-assorting selfing of T. shanghaiensis as T. shanghaiensis one mt specificity and one complete genes of different specificity. The non-assorting T. shanghaiensis genetic tools to experimentally modify mating type genes in T. shanghaiensis while this article was under review. These authors report that starved cells expressing one complete Tetrahymena Yao illustrate a new type of molecular basis for selfing in T. shanghaiensis. The above conclusion is supported by the findings of Lin and Yao (2020) in T. thermophila and T. malaccensis, and only in those to species.

To explain the selfing of T. shanghaiensis, we proposed four a priori hypotheses, the first three of which differ with respect to what is encoded in the MAC genome, as illustrated in Figure S8.

1) Two normal mtGPs with different mating type specificities are present in the MAC of every cell; these mtGPs cannot be purified by assortment because the MAC mat locus is homozygous for both genes. MTA and MTB proteins with different mating type specificities are expressed upon starvation and trigger selfing.

2) Two normal mtGPs with different mating type specificities are present in the homozygous MAC genome but only one can be expressed. Frequent gene conversion causes a random mtGP to be expressed in every cell. Thus, even a clonal population will have a mixture of cells expressing different mating type specificities, leading to selfing upon starvation.

3) Only one type of mtGP is present in the homozygous MAC genome, containing MTA and MTB genes of different mating type specificity, i.e. a heterotypic mtGP. The proteins encoded by this mtGP are sufficient to trigger mating upon starvation.

Our finding of a single mtGP in every population is consistent with only the third hypothesis, a heterotypic mtGP.

It could be argued that mating in T. shanghaiensis is controlled by genes other than the mtGP found in the genome. This alternative is unlikely for the following reasons. As already mentioned, the T. shanghaiensis MTA and MTB genes show identical expression pattern during the life cycle as MTA and MTB genes of other Tetrahymena species (Figure 2C). Furthermore, the locations of both genes in the mating type protein phylogenetic tree (Figure 4) and in the amino acid frequency-based clustering (Figure S6) all correspond closely with the location of T. shanghaiensis in the species phylogeny tree (Figure 2B). Therefore, it seems highly probable that the mtGP of T. shanghaiensis functions in mating. Thus the results are entirely compatible with T. shanghaiensis having a heterotypic mtGP which is responsible for the perpetual selfing, as shown in Figure S8C.

The above conclusion is supported by the findings of Lin and Yao (2020) in Tetrahymena thermophila, published while this article was under review. These authors report that starved cells expressing one complete MTA gene of one mt specificity and one complete MTB gene of different mt specificity behave as non-assorting selfers, exactly as T. shanghaiensis, regardless of which pair of different mt specificities are involved. This finding strengthens our conclusion that the basis for the non-assorting selfing of T. shanghaiensis is the possession of MTA and MTB genes of different specificity. The non-assorting T. shanghaiensis selfer and the selfers investigated by Lin & Yao illustrate a new type of molecular basis for selfing in Tetrahymena (see Data S2 for more details about classification of Tetrahymena selfers). Rigorous proof of our hypothesis must await the development of molecular genetic tools to experimentally modify mating type genes in T. shanghaiensis.

Discussion

Proposed steps in the evolution of the Tetrahymena mtGP

Two mating type proteins (MTA and MTB) embody mating type specificity in the multiple mating type system of T. thermophila. These proteins contain ~1,500 amino acids each, and are encoded by adjacent head-to-head genes in the mtGP (Cervantes et al., 2013). The two genes have very similar organization and both belong to the superfamily of FLR domain proteins, but they only share about 6% sequence similarity. In this report, we identified and compared the MAC mating type protein-coding loci in nine
additional Tetrahymena species, encompassing the phylogenetic diversity of the genus. Eight of the additional species were found to have homologs to *T. thermophila* MTA and MTB, while an asexual strain of the most distantly related species, *T. paravorax,* has an MTA homolog but lacks any trace of an MTB homolog.

The findings we have reported here suggest the major steps in the evolution of the *Tetrahymena* mtGP over the last 150 Myr, illustrated in Figure 6:

1) A cysteine-rich “Furin-like repeat” domain protein acquired mating-related activity in an Oligohymenophorean ancestor of *Tetrahymena* and *Paramecium.* A terminal exon, encoding a “Furin-like repeat” domain and five transmembrane helixes, acquires mating-related function, and evolves into distinct mating type genes in *Tetrahymena* and *Paramecium.* Step 2. In *Tetrahymena,* a DNA rearrangement generates a copy of the mating type gene (MTA'). Step 3. The MTA' gene acquires a new mating-related function, becoming the MTB gene. A set of paralogs encoding different mating type specificities (colored arrows) co-evolve from each gene. Intron location and phase are conserved during this differentiation and following events. Step 4. A second translocation causes MTA and MTB to become adjacent genes in opposite orientation, thus generating the head-to-head mtGP. A new wave of co-evolutionary mutational changes generates additional mating type specificities. Step 5. An MTA rearrangement deletes intron #4 (black vertical dotted line) and creates intron #6.5 (marked with triangle) within an MTA gene in a “The-Mal” subgroup ancestor, followed by a new wave of co-evolved changes that generate additional mating type specificities. Note: if the MTA and MTB alleles brought into contiguity at step 4 had encoded different mating type specificities, this evolutionary intermediate initially would have been a perpetual selfer, like present day *T. shanghaiensis.*
4) A subsequent DNA rearrangement brought into contiguity two cognate MTA and MTB genes, in head-to-head orientation, generating the *Tetrahymena* mtGP that we see today.

5) A rearrangement in MTA, marked by the loss of intron #4 and the appearance of intron #6.5, occurred in an ancestor of the “The-Mal” subgroup.

It seems very likely that every step was quickly followed by waves of paralogous diversification of MTA and MTB to generate different mating type specificities.

It is possible that steps 2 and 4 occurred at once, i.e., step 2 was a “palindromic duplication”, and thus the head-to-head MTA-MTB contiguity preceded their functional differentiation. However, this seems unlikely because intra-strand gene conversion within the palindrome would have precluded the functional differentiation of the two genes, due to mutual DNA sequence self-correction among the two copies, such as described in metazoan Y chromosome palindromic duplications (Trombetta and Cruciani, 2017).

These findings raise the question, what could have been gained by having MTA and MTB genes immediately adjacent to one another? The tight linkage of mating related genes provides important advantages, such as to “facilitate the coordinated expression” and “cosegregation of the interacting genes” (Uyenoyama, 2005). One additional consequence of MTA-MTB contiguity in *Tetrahymena* is that it minimizes the frequency of selfing among sexual progeny that would otherwise occur as a consequence of independent allelic assortment of the two genes in the MAC. If the MTA and MTB genes were located on different MAC chromosomes, the two genes would assort independently in double heterozygotes. MACs that are pure for non-cognate (heterotypic) MTA and MTB genes would then be frequently generated, ultimately resulting in non-assorting selfers (see Data S3 and Figure S9 for a detailed explanation). Thus, reducing the length and the sequence similarity of the intergenic segment between MTA and MTB genes has the effect of minimizing selfer-generating germline or somatic recombination events. The evolution of the mtGP, with its tight contiguity of the cognate (homotypic) MTA-MTB genes, likely was a major step in the evolution of the cross-breeding genetic economies generally observed in *Tetrahymena* species today.

Finally, the mtGP organization has also proven its versatility by allowing the evolution of additional genetic and molecular mechanisms of MTD capable of adjusting selfing frequency in *Tetrahymena*. These mechanisms can promote outbreeding or inbreeding under conditions of low or high reproductive stress, respectively (see below for details). A better understanding of these mechanisms will come when studies of the MIC organization of the mating type loci of the various species become available.

**True-breeding selfing in *T. shanghaiensis***

Our work has confirmed that *T. shanghaiensis* is a true-breeding selfer and has shown that it contains a single mtGP in its MAC that behaves structurally and functionally like the mtGPs of the other species investigated here (Figures 2, 3, 4, S2, S4, and S6). The simplest explanation for its obligatory selfing behavior is that *T. shanghaiensis* mtGP is heterotypic, i.e. its MTA and MTB genes have different mating type specificity. This conclusion is supported by the finding that *T. thermophila* mutants, which fail to complete MTD and are left in the MAC with intermediates containing a complete MTA and a complete MTB gene but with different mt specificity, are non-assorting selfers (Lin and Yao, 2020) just like wild-type *T. shanghaiensis*. Evolutionarily, the *T. shanghaiensis* chimeric mtGP could have been generated by a simple DNA rearrangement, such as a non-homologous meiotic recombination event between two normal mtGPs of different mating types occurring at the MTA-MTB intergenic region in a heterozygote, resulting in the replacement of either gene with a homolog of different mating type specificity. An alternative way in which the heterotypic mtGP could have arisen is by successive mutations in the two genes of an initially homotypic mtGP that promoted increasingly strong interactions between their encoded proteins ultimately leading to efficient selfing, favored under conditions of high reproductive stress.

As a perpetual selfer, *T. shanghaiensis* can be considered to be unisexual, in the sense used to describe mating in the absence of any intra-species diversity at the mating type locus, as occurs in some fungi (reviewed by Heitman (2015)). Indeed, that author argues that the last eukaryotic common ancestor, was unisexual, i.e., a perpetual selfer using the *Tetrahymena* terminology. The species tree (Figure 2B) shows that *T. shanghaiensis* is surrounded by species with homotypic mtGPs. Thus, it seems most probable that the unis�性 (perpetual selfing) of *T. shanghaiensis* is derived, rather than ancestral in the genus...
Tetrahymena. That does not exclude, however, the possibility of a unisexual common ancestor of the “Bor-ealis” and “Australis” Tetrahymena clades.

The intron #4 and #6.5 rearrangement provide a glimpse into the evolution of a different mating type system

While sexual reorganization events (meiosis, fertilization, MAC differentiation) are highly conserved among Ciliates, a striking variety of mating type systems have evolved within this group (Phadke and Zufall, 2010). For example, in the multiple mating type system of the hypotrich Ciliate Euplotes, the two proteins that embody ligand and receptor function for each mating type specificity are the products of intron-splicing variants of the same gene; both proteins are very small, in the order of 40 amino acids, reminiscent of cytokines of multicellular eukaryotes (Luporini et al., 1986; Miceli et al., 1992). In the binary mating type system of the heterotrich Ciliate Blepharisma, the mating type ligand is not a protein but a small tryptophan-related molecule (Miyake, 1996; Sugiura et al., 2005). This diversity implies that the molecules that embody mating type specificity in Ciliates have independently undergone major successive replacements. However, the lack of evolutionary intermediates makes it extremely challenging to trace how this diversity evolved among major groups.

The results reported here have allowed us to infer a succession of replacement waves, occurring within the genus Tetrahymena, which generated a diversity of mating type proteins from an ancestral “Furin-like repeat” protein (Figure 6). Serendipitously, this work also uncovered a more recent replacement wave in which an MTA allele, generated along the way by the loss of conserved intron #4 and the gain of intron #6.5, is inferred to have de novo replaced the MTA alleles of mtGPs of every mating type specificity among the sequenced MTA genes in the “The-Mal” subgroup. Each of the rearrangements that resulted in the two intron changes likely happened only once, with the final rearrangement presumably resulting in one MTA gene of one particular mating type specificity. This variant MTA gene then had to spread and diversify, to ultimately be able encode every known MTA mating type specificity found today in the “The-Mal” subgroup. It seems reasonable to expect that MTA genes also had to co-evolve, in order to allow all the appropriate positive and negative mating type protein interactions required to promote mating between different mating types and prevent selfing in the multiple mating type system. The generation of “raw material” for the re-evolution of multiple mtGPs of different mating type—all containing the variant introns in the MTA gene—was likely facilitated by two special features: the tandem array organization of multiple mtGPs in the T. thermophila germline (micronuclear) genome (Cervantes et al., 2013), in combination with unequal meiotic crossing over, a capacity which has been well documented in the case of Tetrahymena leucine-rich repeat genes (Xiong et al., 2019).

The mtGP: a durable and effective vehicle for Tetrahymena unicell adaptation to reproductive stress fluctuations

T. thermophila cells, maintained by asexual reproduction in the laboratory for long periods in the absence of mating, eventually become sterile (Simon and Nanney, 1979). This failure was inferred to be due to the random accumulation of deleterious mutations in the MIC. The time-sensitive deterioration of their germline results in the susceptibility of Tetrahymena cells to reproductive stress.

Most Tetrahymena species tend to be primarily outbreeders. As previously discussed (Orias et al., 2017), this represents a balance between mechanisms that promote outbreeding and inbreeding. Features of Tetrahymena biology that promote outbreeding include a long sexual immaturity period, intranuclear coordination during MTD, allelic assortment and, in the “Australis” clade, synclonal MTD. Features capable of promoting inbreeding include multiple mating type systems, selfing, and karyonidal MTD in the “Borealis” clade. This investigation of the mating type genes of a broader set of Tetrahymena species has contributed an additional finding relevant to the balance between inbreeding and outbreeding, namely the rare occurrence of a putative heterotypic mtGP, which ensures obligatory, perpetual selfing in T. shanghaiensis.

Some asexual Tetrahymenas, such as T. pyriformis and T. vorax also studied here, have lost their MIC (so called amicronucleates, amics) and can no longer conjugate. Such an extreme feature avoids the germline deterioration that would affect sexual Tetrahymena cells under conditions of severe sexual reproductive stress. Tetrahymena amics are presumably capable of long-term adaptation to changing environments by virtue of allelic assortment in the polyploid MAC. The number of copies of favorable mutations can
increase by random assortment and come to phenotypic expression, while unfavorable mutations can be eliminated.

A puzzling feature of the asexual strains investigated here is that their mtGP retain open reading frames (at least for the sequenced exons), even though the proteins are no longer needed for mating, and their expression is not induced by starvation. One trait that would delay the emergence of internal in-frame stop codons is the variant Ciliate genetic code, which has a single stop codon, UGA. It is also possible that the mtGP has other useful function(s), unrelated to mating and expression is induced by some condition other than starvation, which keeps it under selection.

The species investigated here were chosen to provide a sample of the diversity of the Tetrahymena genus, which now contains nearly 100 identified species, with no end in sight. As the breeding systems of additional Tetrahymena species are molecularly characterized, our current picture of how they have evolved in reaction to fluctuating levels of reproductive stress will no doubt be enlarged and enriched.

Limitations of the study
In this work, we mainly focused on the MAC mtGP. Even though these results provide some clues of the evolution of the MIC mating type locus, there are still many unknowns. Subsequent studies of MIC mtGPs should provide a more elaborate picture of the evolutionary process. In addition, the most distantly related species, T. paravorax, seems to be asexual, so it will be more informative if we can find and investigate a sexual outgroup species in the future.

Resource availability
Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Wei Miao (miaowei@ihb.ac.cn).

Materials availability
This study did not generate new unique reagents.

Data and code availability
The accession number for the data reported in this paper is GEO: PRJNAS10545.

Methods
All methods can be found in the accompanying Transparent methods supplemental file.

Supplemental information
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101950.

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Author contributions
G.Y. designed and performed most of the experiments, analyses, and wrote the manuscript. W.Y. contributed to genome/transcriptome assembling. X.H. amplified and sequenced introns and intergenic regions of mtGPs which only have RNA-Seq data. K.C. contributed to mating type test and strain storage. J.X. and E.O. contributed to data analyses. E.H. and E.O. reviewed the manuscript. W.M. contributed to the experimental design, and reviewed the manuscript.
Declaration of interests

The authors declare no competing interests.

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Supplemental Information

Evolution of the mating type
gene pair and multiple
sexes in *Tetrahymena*

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Supplemental Information

Figure S1. Mating type determination in *Tetrahymena*. Related to Figure 1. (A) Synclonal MTD. The four karyonides from a pair are usually determined by Mendelian genetics to express the same mating type (shown as the same color) which could restrict inbreeding. All possible outcomes (genotypes and mating types) are illustrated in the sample cross shown. (B) Karyonoidal MTD. The four karyonides from a pair express independently determined mating types, unrelated to the mating type of either parent or to those of the other karyonides from same pair. Thus, karyonoidal MTD could favor inbreeding. For more information, please see Phadke and Zufall (2010) and Orias et al. (2017).

Figure S2. Sequence alignment and conservation of intron location and phase among all sequenced MTA, MTB and MTAL *Tetrahymena* proteins. Related to Figure 3B. Colored dots, introns: blue, phase 0, inserted between two codons; green, phase 1, inserted between the 1st and 2nd codon nucleotide; red, phase 2, inserted between the 2nd and 3rd codon nucleotide). Black lines separate MTA, MTB and MTAL proteins. “Dark red background”, identical site (Global score = 1); “Pink letter on white background”, high similarity (Global score over 0.7); “Black letter on white background”, low similarity (Global score below 0.7). Similarity at each site was based on GONNET matrices. “Pink background”, conserved cysteine residue in all proteins except MTAL.

Figure S3. mtGP synteny in *Tetrahymena* species. Related to Figure 2A. Note that a) the MTAL gene of *T. paravorax* is syntenic with all the other MTA genes; but there is no MTB gene homolog; b) the synteny of *T. americanis* mtGP could not be tested due to lack of a sequenced genome and c) *T. vorax* is the only species whose mtGP not syntenic with others.

Figure S4. Strong conservation of MTA and MTB intron location and phase within each of the three subgroups. Related to Figure 3B. Intron colors as in Fig. S2. Only relevant segments of the multiple alignments are shown in this figure.

Figure S5. Phylogenetic tree of mating type proteins based on full length MTA and MTB sequences. Related to Figure 4. The phylogenetic tree was constructed under WAG+I+G+F model. Numbers at each node, bootstrap values (1000 replicas). Branch length, number of base substitutions per site. Red boxes: deviations from the species tree. The MTA and MTBs branches are shown opposite to one another to facilitate comparisons between them.

Figure S6. MTA and MTB proteins show different amino acid usage frequencies among different subgroups. Related to Figure 5. (A), (B) and (C) Protein clustering based on Euclidean distance calculated by amino acid usage frequency. (A) is based on full-length sequences; (B) is based on C-terminal exon (Distal-third) encoded sequences; (C) is based on the rest of sequence (Proximal-two-thirds). Color bar, amino acid usage frequency. Min and Max, minimum and maximum value of each column, respectively. (D) Principal Components Analysis (PCA) of amino acid usage frequency (based on full-length sequences). Data points for mating type proteins of each subgroup are shown in the same colors as in panel A. *T. shanghaiensis* data points are highlighted for reasons explained in the last Results section.
Figure S7. Pedigree of two *T. shanghaiensis* populations whose mtGPs were independently sequenced as biological replicates. Related to Figure 6. Key to panel labels. S, starvation. G, growth. Note that selfing leads to the same conjugation events described in Figure 1, including the generation of sexual progeny with new MICs and MACs. Green dashed oval in panel 4: mature progeny cell, originally derived from the pair within the green dashed circle in panel 3, which was cloned and allowed to starve and self again (panel 7). Red stars: two selfing cultures which represent independently sequenced biological replicates.

Figure S8. Possible mechanisms responsible for obligatory selfing in *T. shanghaiensis*. Related to Figure 6. Key to symbols: orange and blue, two different mating type specificities; divergent thick arrows, MAC mtGP; sticks attached to cell membrane, mating type proteins: thick, MTA; thin, MTB. (A) mtGPs with different mating type specificities are present in the MAC of every cell; these mtGPs cannot be purified by assortment because the MAC *mat* locus is homozygous for both alleles. MTA and MTB proteins with different mating type specificities are expressed upon starvation and trigger selfing. (B) mtGPs with two different mating type specificities are present in the homozygous MAC genome but only one can be expressed. Frequent gene conversion causes a random mtGP to be expressed in every cell. Thus, even a clonal population will have a mixture of cells expressing different mating type specificities, leading to selfing upon starvation. (C) Only one type of mtGP is present in the homozygous MAC genome, containing *MTA* and *MTB* genes of different mating type specificity, i.e. heterotypic mtGP. The proteins encoded by this mtGP are sufficient to trigger mating upon starvation.

Figure S9. **MTA** and **MTB** genes could have been on the same or different MAC chromosomes during mtGP evolution. Related to Figure 6. (A). The *MTA* and *MTB* genes are adjacent on the same MAC chromosome, the arrangement we find today. (B) and (C). Possible types of *MTA* and *MTB* gene organization at stage 2 of mtGP evolution in Figure 6. The *MTA* and *MTB* genes are on different (B) or same (C) MAC chromosomes. Key to symbols. Left-pointing triangles, *MTA* genes; right-pointing triangles, *MTB* genes; blue and red, different mating type specificities (e.g., mt I and II). Key to stages. MAC initially heterozygous for I and II mating type specificities: 1, before assortment; 2, after assortment is completed; 3, behavior of sexually mature terminal assortants after starvation (selfing or no selfing). Under case A, only a single, homotypic *MTA* and *MTB* gene pair remains in the MAC of terminal assortants (stage 2) and no selfing is expected. Under case B, only a single *MTA* allele and a single *MTB* allele remain in the MAC of terminal assortants. The MACs of 50% of the terminal assortants will be pure for heterotypic *MTA* and *MTB* alleles, and those cells will be non-assorting selfers (see Text S2). Under case C, two main types of terminal assortants are expected with equal frequencies having homotypic *MTA* and *MTB* gene combinations, respectively. Rare MAC recombination could occasionally generate a heterotypic mtGP (bottom chromosome in panel C1); terminal assortants pure for this chromosome would be non-assorting selfers. (See Text S3 for more details).

Table S1. Conservation of intron phase and location among functional *Tetrahymena* mating type genes. Related to Figure 3.

Table S2. Number of independent mating tests done within and between sexual *Tetrahymena* strains used. Related to Table 2.
Table S3. Collection sites and other information on previously unreported strains. Related to Table 2.

Data S1. Evolution of amino acid usage among the mating type genes of sexual *Tetrahymena* species. Related to Figure 5.

Data S2. Classification of *Tetrahymena* selfers. Related to Figure 6.

Data S3. Evolution of *MTA-MTB* contiguity after functional differentiation would have discouraged selfing. Related to Figure 6.
Figure S1

A  Synclonal MTD

mt I (mat-1/mat-3)

mt II (mat-2/mat-3)

or

mt II (mat-1/mat-2 or mat-1/mat-3)

or

mt III (mat-2/mat-3)

or

mt III (mat-3/mat-3)

B  Karyonidal MTD

mt I (mat-1/mat-1)

mt II (mat-1/mat-1)

mt III (mat-1/mat-1)

or

mt III (mat-1/mat-1)

or

.....

or

many others
Figure S3

T. the  
T. mal  
T. pyr  
T. bor  
T. can  
T. sha  
T. pig  
T. par
Figure S6

The Distance - Mal
0.90
0.95

"Pig" Mal" subgroup

The Hydrophobic - Mal subgroup

PC1 (62.8%)

PC2 (12.2%)

The-Mal vs. Bor-Can
The-Mal vs. Pig-Ame
Bor-Can vs. Pig-Ame

The-Mal
Bor-Can
Pig-Ame

PC1 (62.8%)

PC2 (12.2%)

The-Mal
Bor-Can
Pig-Ame

The-Mal vs. Bor-Can
The-Mal vs. Pig-Ame
Bor-Can vs. Pig-Ame

The-Mal
Bor-Can
Pig-Ame

The-Mal
Bor-Can
Pig-Ame
**Figure S9**

**A**
- **1.** Non-assorting selfer
- **2.** Homotypic MAC
- **3.** No selfing

**B**
- **1.** Non-assorting selfer
- **2.** Homotypic MAC
- **3.** Non-assorting selfer

**C**
- **1.** Non-assorting selfer
- **2.** Homotypic MAC
- **3.** No selfing
Table S1. Conservation of intron phase and location among functional *Tetrahymena* mating type genes. Related to Figure 3.

| Tetrahymena Mating Type Genes | Introns and their phases |
|-------------------------------|--------------------------|
| Species | MT | Gene | 1 | 2 | 3 | 4 | 5 | 6 | 6.5 | 7 |
| thermophila | II | A | - | 1 | 0 | - | 1 | 0 | 1 | 0 |
| thermophila | III | A | 0 | 1 | 0 | - | 1 | 0 | 1 | 0 |
| thermophila | IV | A | - | 1 | 0 | - | 1 | 0 | 1 | 0 |
| thermophila | V | A | 0 | 1 | 0 | - | 1 | 0 | 1 | 0 |
| thermophila | VI | A | 0 | 1 | 0 | - | 1 | 0 | 1 | 0 |
| thermophila | VII | A | 0 | 1 | 0 | - | 1 | 0 | 1 | 0 |
| malaccensis | X | A | 0 | 1 | 0 | - | 1 | 0 | 1 | 0 |
| borealis | X | A | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| borealis | Y | A | 0 | 1 | 0 | 0 | 1 | 0 | 1 | 0 |
| borealis | Z | A | - | 2 | 0 | 1 | 1 | 0 | - | 0 |
| canadensis | X | A | - | - | 0 | 1 | 1 | 0 | - | 0 |
| shanghaiensis | Selfer | A | 0 | 1 | 0 | 1 | 1 | 0 | - | 0 |
| americanis | X | A | 0 | 1 | 0 | 1 | 1 | 0 | - | 0 |
| americanis | Y | A | 0 | 1 | 0 | 1 | 1 | 0 | - | 0 |
| pigmentosa | I | A | 0 | 1 | 0 | 1 | 1 | 0 | - | 0 |
| pigmentosa | III | A | 0 | 1 | 0 | 1 | 1 | 0 | - | 0 |
| thermophila | II | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| thermophila | III | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| thermophila | IV | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| thermophila | V | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| thermophila | VI | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| thermophila | VII | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| malaccensis | V | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| borealis | X | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| borealis | Y | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| borealis | Z | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| canadensis | X | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| shanghaiensis | Selfer | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| americanis | X | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| americanis | Y | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| pigmentosa | I | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |
| pigmentosa | III | B | - | 1 | 0 | 1 | 1 | 0 | - | 0 |

Key to cell colors: blue, green and pink: phase 0, 1 and 2 intron, respectively; yellow: missing conserved intron.
Table S2. Number of independent mating tests done within and between sexual *Tetrahymena* strains used. Related to Table 2.

| the mt II | the mt III | the mt IV | the mt V | the mt VI | the mt VII | X | bor the mt X | bor mt Y | bor mt Z | can the mt X | pig the mt I | pig the mt III | ame the mt X | ame the mt Y | Mating type |
|----------|------------|----------|----------|----------|-----------|---|-------------|----------|-----------|-------------|--------------|--------------|-------------|-------------|-------------|
| >>4 neg  | >>4 pos    | >>4 pos  | >>4 pos  | >>4 pos  | 4 neg     | 4 neg | 4 neg       | 4 neg    | 4 neg     | 4 neg       | 4 neg        | 4 neg        | 4 neg       | 4 neg       | >>4 neg     | the mt II   |
| >>4 neg  | >>4 pos    | >>4 pos  | >>4 pos  | >>4 pos  | 4 neg     | 4 neg | 4 neg       | 4 neg    | 4 neg     | 4 neg       | 4 neg        | 4 neg        | 4 neg       | 4 neg       | >>4 neg     | the mt III  |
| >>4 neg  | >>4 pos    | >>4 pos  | >>4 pos  | >>4 pos  | 4 neg     | 4 neg | 4 neg       | 4 neg    | 4 neg     | 4 neg       | 4 neg        | 4 neg        | 4 neg       | 4 neg       | >>4 neg     | the mt IV   |
| >>4 neg  | >>4 pos    | >>4 pos  | >>4 pos  | >>4 pos  | 4 neg     | 4 neg | 4 neg       | 4 neg    | 4 neg     | 4 neg       | 4 neg        | 4 neg        | 4 neg       | 4 neg       | >>4 neg     | the mt V    |
| >>4 neg  | >>4 pos    | >>4 pos  | >>4 pos  | >>4 pos  | 4 neg     | 4 neg | 4 neg       | 4 neg    | 4 neg     | 4 neg       | 4 neg        | 4 neg        | 4 neg       | 4 neg       | >>4 neg     | the mt VI   |
| >>4 neg  | >>4 pos    | >>4 pos  | >>4 pos  | >>4 pos  | 4 neg     | 4 neg | 4 neg       | 4 neg    | 4 neg     | 4 neg       | 4 neg        | 4 neg        | 4 neg       | 4 neg       | >>4 neg     | the mt VII  |

Key to entries:
- the *T. thermophila*
- mal *T. malaccensis*
- bor *T. borealis*
- can *T. canadensis*
- pig *T. pigmentosa*
- ame *T. americanis*
- pos Mating observed
- neg No mating observed

- >> 4 Many independent tests, including for many experiments unrelated to this project.
- >3 More that 4 independent tests, including for experiments unrelated to this project.
Table S3. Collection sites and other information on previously unreported strains. Related to Table 2.

| Species       | Mating type | Strain  | Collection site       | Type  | State | Latitude    | Longitude  |
|---------------|-------------|---------|-----------------------|-------|-------|-------------|------------|
| *T. americanis* | X           | SD21194 | LakeBarkley1          | lake  | KY    | 37.05093    | -88.15287  |
| *T. americanis* | Y           | SD21244 | ShantyHollowLake      | lake  | KY    | 37.14453    | -86.38432  |
| *T. borealis*  | Y           | SD19502 | KinzuaBayElijah       | lake  | PA    | 41.81892    | -78.94685  |
| *T. borealis*  | Z           | SD19803 | LittleSalmonCr165     | stream| PA    | 41.50955    | -79.15158  |
| *T. pigmentosa*| III         | SD19481 | FishCrSloughAshlanč   | stream| WI    | 46.58583    | -90.932    |
| *T. pigmentosa*| I           | SD20427 | HalfMoonPondBrook     | stream| NH    | 43.17025    | -72.08968  |
Data S1. Evolution of amino acid usage among the mating type genes of sexual *Tetrahymena* species.

In main text we have reported evidence of co-variation in *MTA* and *MTB* genes that could reflect changes in putative requirements for functional protein interactions between the two gene products (Orias et al., 2017). To take a deeper look, we examined amino acid usage in the MTA and MTB proteins of sexual species. Hydrophobicity, for example, can influence protein flexibility during protein-protein interaction (Radivojac et al., 2004). By using either the hierarchical or k-means method, we clustered the amino acid usage frequencies of the MTA and MTB proteins separately for a) the entire lengths of the proteins, b) the proximal ~2/3 of the proteins (from N-terminus to intron 7), and c) the distal ~1/3 (C-terminal exon) of the proteins. The results are shown in the 3 panels of Figure S6. Comparison of the three clustering trees shows that they are different from one another and that none of them have the same topology as either the species phylogenetic tree (Figure 2C) or the mating type protein trees for each of the three segments (Figure 4).

The following unusual topologies were common to all three amino acid usage clustering trees.

1) The main branches of the amino acid clustering trees tend to be different species phylogenetic groups rather the MTA vs. MTB main branching found in the protein phylogenetic tree.

2) Amino acid usage the “Bor-Can” subclade of the borealis clade clusters with the “Australis” clade rather than with the other (“The-Mal”) subgroup of the “Borealis” clade.

Although we don’t have a clear rationale for these differences, we can understand some of the variables influencing the branching pattern of particular amino acid usage clustering trees for the distal 1/3 of the MTA and MTB proteins.

a) For *T. thermophila*: the terminal half of the terminal exon (distal ~1/6) of both *MTA* and *MTB* genes are identical because the tandem array of MIC mtGPs contain only one copy of the *MTA* distal ~1/6 (at the “head” of the array), and only one copy of the *MTA* distal ~1/6 (at the “tail” of the array). Even for the rest of the C-terminal exon, the copies present in the six genes are >95% identical to one another (Cervantes et al., 2013). Thus, within the *T. thermophila* branch of that distal 1/3 clustering tree, all the MTA proteins must cluster together in one subcluster, while all the MTB proteins must cluster together in the other subcluster, as they indeed do.

b) For the “Australis” clade: although the mtGP composition of their MIC mating type locus has not yet been described, we show here that the two sequenced MTA proteins in each species investigated (*pigmentosa* and *americanis*) show high sequence conservation with one another. The same is true for MTB proteins. Interestingly, for the “Australis” clade, the high degree of amino acid sequence conservation seen in the distal ~1/3 extends to the “middle third”. This has to influence the branching pattern seen for the proximal 2/3 clustering tree in a different way for “Australis” species than for the rest of the species.
c) For *T. borealis*, there is essentially no sequence conservation in the distal 1/3 of either MTA or MTB proteins. This disorganizes the branching patterns seen for the three MTAs and MTB proteins relative to the other groups.

A “Principal Components Analysis” (PCA) of the amino acid usage frequency for each full-length mating type protein of every *Tetrahymena* sexual species (Figure S6D) shows a similar clustering pattern as that obtained by either the hierarchical or k-means method of the full-length proteins (Figure S6A).

Although we can make some sense of the amino acid usage clustering we observe, it is clear that to fully understand the details of the clustering we must wait until we have an understanding of the 3-D structure of these proteins and experimental analyses of their interactions.

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Data S2. Classification of Tetrahymena selfers.

While trying to distinguish various types of selfers encountered in this work, in the context of our current molecular knowledge about Tetrahymena mating types, it became useful to devise a more descriptive categorization of Tetrahymena selfers. Our proposed classification, summarized in Table SD1 below, is based on a combination of the genetic behavior of selfers during asexual reproduction and after sexual reorganization upon selfing. We consider it a descriptive classification because more than one molecular mechanism may be found to generate the same type of selfer genetic behavior.

A. Selfing behavior during asexual reproduction.

1) Assorting selfers. When sexual progeny become sexually mature, a mixture of mating type alleles is often found in the MAC; if allowed to starve, subcultures of these cell lines will self. Upon further asexual propagation, the mating type alleles ultimately assort, so cells become pure for a single MAC mating type allele. At this point, they, and their subsequent vegetative descendants no longer self and instead express a defined mating type. Examples of such selfers are the “classical” selfers analyzed in T. thermophila by Allen and Nanney (1958).

2) Non-assorting selfers. Upon reaching sexual maturity, progeny cells and their vegetative descendants always self upon starvation. No vegetative assortants expressing a single mating type are obtained. Examples are the “suicidal” Tetrahymena selfers described by Nanney (1953), the persistent selfers reported by Lin and Yao (2020) and the T. shanghaiensis perpetual selfers investigated in this study.

B. Perpetuation of selfing after sexual reorganization and maturity.

Two possibilities are known:

1) Sexually non-true-breeding selfers. The sexual progeny of selfing pairs are not themselves selfers when they reach sexual maturity. Examples are the selfers investigated by Nanney and Allen listed under A1 above, where the old MAC, responsible for selfing, is destroyed upon conjugation. Persistent selfers obtained by post-conjugation Ku80 silencing (Lin and Yao, 2020) should also fall in this category.

2) Sexually true-breeding selfers. The sexual progeny of non-assorting selfers are themselves always vegetatively non-assorting selfers. The T. shanghaiensis selfers used in this
study are an example.

Based on the above classifications, the ordinary *T. thermophila* selfers analyzed by Allen and Nanney (1958), are classified as vegetatively assorting, sexually non-true-breeding selfers. In contrast, the *T. shanghaiensis* strain used here is a vegetatively non-assorting, sexually true-breeding selfer.

Other types of selfers may well be discovered in nature or in the lab.

**Table SD1. Proposed classification and examples *Tetrahymena* selfers. Related to Figure S7.**

| Sexual Behavior       | Asexual Behavior                                                                 |
|-----------------------|---------------------------------------------------------------------------------|
|                       | **Vegetatively assorting**                                                      |
|                       | **Vegetatively non-assorting**                                                  |
| Sexually non-true-breeding | *T. thermophila*  
|                        | “classical” selfers (Allen and Nanney, 1958)                                    |
| Sexually true-breeding | *T. shanghaiensis* selfers  
|                        | (this article) and newly described *T. thermophila* persistent selfers (Lin and Yao, 2020) |

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NANNEY, D. L. 1953. Nucleo-cytoplasmic interaction during conjugation in *Tetrahymena*. *The Biological Bulletin*, 105, 133-148.
Data S3. Evolution of MTA-MTB contiguity after functional differentiation would have discouraged selfing.

The putative events that led to the evolution of the Tetrahymena mtGP have been presented (Figure 6) and discussed in main text. We have argued that MTA and MTB were not tightly linked to one another at the time they evolved their distinct and complementary functions in co-stimulation and pair formation. Otherwise, frequent gene conversion within the palindromic duplication would have acted to maintain the similarity of the two genes and would have precluded the evolution of their functional differences.

The possible non-adjacency of MTA and MTB, once the two genes had evolved functional differences and replaced an older mating type system, is interesting with regard to selfing. The frequency of selfing after allelic assortment is complete would have depended on the location of the two genes in a common ancestor of the “Borealis” and “Australis” clades, as illustrated in Figure S9. If MTA and MTB resided on different MAC chromosomes when they evolved their functional differences, the probability of generating MAC genotypes capable of resulting in non-assorting selfers would have been high (Figure S9B) compared to what we see now (Figure S9A).

The opposite is true if MTA and MTB had evolved at locations that ended up on the same MAC chromosome Figure S9, panel C). In this case, the probability of selfing would be very low because intrachromosomal MAC recombination is rare between genes on the same MAC chromosome (Longcor et al., 1996), likely related to infrequent DNA damage repair events occurring between the two genes during asexual reproduction. The heterotypic MAC chromosome shown at the bottom of Figure S9, panel C1 is an example of such rare MAC intrachromosomal recombination. If the actual number of chromosome types in such MAC had been 22 MTA1-MTB1 copies, 22 MTA2-MTB2 copies and 1 MTA1-MTB2 copy, then, after complete assortment of this clone, the percentage of assortants would have been 49% mt I, 49% mt II and 2% non-assorting selfers, respectively.

Recombination events (MIC or MAC) between MTA and MTB genes in the current mtGPs are extremely rare. No cases have been reported in the extensive laboratory work with T. thermophila, but such an event is a possible basis for the naturally occurring heterotypic T. shanghaensis mtGP described in this article. The low probability of recombination is explained by the main features of the contemporary mtGP: the side by side location of the genes and the short length and mating type-specific sequence diversification of the intergenic region.

Reference

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Transparent Methods

Biological methods

The *Tetrahymena* species whose genomes we sequenced were identical to those in our recent comparative genomics report (Xiong et al., 2019). *T. americanis* and *T. pigmentosa* strains were provided by Dr. Paul Doerder (Cleveland State University, USA). All cells were grown in SPP medium (1% Proteose Peptone, 0.2% glucose, 0.1% yeast extract and 0.003% Sequestrene).

Mating type differences between strains of the same species were tested by a pairing assay. To induce starvation, cells were washed twice with 10 mM Tris-Cl (pH 7.4) and then resuspended in the same solution and incubated for at least 12 h, at which this time no obvious food vacuoles could be observed. For the mating tests, equal numbers of fully starved cells of two different strains were mixed at a final cell density of approximately $2 \times 10^5$ cells/ml (final volume ~1 ml). If any pairs formed (usually over 80%), and no pairs were found in the starved, unmixed controls, then the two strains were considered to be of different mating type. Every mating test was repeated at least four times (see Table S2 for more details).

Identification of mating type homologs in published *Tetrahymena* genomes

A total of ten *Tetrahymena* genomes were chosen for our analyses: *T. thermophila*, *T. malaccensis*, *T. pyriformis*, *T. vorax*, *T. borealis*, *T. canadensis*, *T. pigmentosa*, *T. americanis*, *T. shanghaiensis*, and *T. paravorax* (Table 1). Candidate mating type gene homologs in genome sequences were determined by performing a local tBLASTn (version 2.2.25) (Altschul et al., 1997) search against the *T. thermophila* MTA and MTB genes (Cervantes et al., 2013). To confirm intron/exon structures, RNA-Seq data was mapped to each genome with TopHat (version 2.0.9) and visually checked using IGV (version 2.4.6). This resulted in a correction of previously predicted (Xiong et al., 2019) intron/exon structures for *T. canadensis* and *T. shanghaiensis*. To check whether mating type genes were at the same MAC genome location as in *T. thermophila*, chromosome synteny analysis of the surrounding regions was done based on the *Tetrahymena* comparative genomics database (Yang et al., 2019) (http://ciliate.ihb.ac.cn/).

Identification of mating type homologs in strains without sequenced genomes
Complete mating type homologs, in strains without sequenced genomes, were first identified by RNA sequencing; PCR was used to fill any gaps. This method was used for two strains of *T. borealis, T. americanis,* and *T. pigmentosa* (Table 1). Strain details are shown in Table S3. Total RNA was extracted from each strain at the starvation stage using the RNeasy Protect Cell Mini Kit (Qiagen), as described (*TetraFGD*) (Xiong et al., 2011). Poly-A tailed mRNA was then enriched using Sera-Mag magnetic oligo (dT) beads. Illumina sequencing libraries were constructed and paired-end (150 bp × 2) sequencing was done for all samples using Illumina Hiseq4000 sequencer. After adaptor trimming by Trim-Galore (version 0.4.0) (Wu et al., 2011), clean reads were *de novo* assembled using Trinity (version trinityrnaseq_r20140717) (Grabherr et al., 2011). The resulting transcriptomes were subsequently searched with *T. thermophila* MTA and MTB gene sequence using tBLASTn (version 2.2.25). Usually only incomplete transcripts were assembled; PCR amplification and Sanger sequencing were used to link these fragments and obtain the intergenic regions of the mating type genes. PCR primers are listed in Table S4. To confirm the intron/exon structures, clean reads were mapped back to the sequence with TopHat (version 2.0.9) (Kim and Salzberg, 2011) and visually checked by IGV (Integrative Genome Viewer; version 2.4.6) (Robinson et al., 2011, Thorvaldsdóttir et al., 2013).

The sequences of MAC mating type loci generated in this study have been deposited in GenBank (Table S5). Genome data for all strains except *T. pigmentosa* and *T. americanis* (which do not yet have complete genome assemblies) can be accessed through Tetrahymena comparative genomics database (http://ciliate.ihb.ac.cn/); additional sequencing data has been submitted under accession numbers PRJNA510545.

**Estimation of gene expression levels**

To estimate gene expression levels, RNA sequencing reads were mapped to the MAC mating type loci with TopHat (version 2.0.9) and the FPKM value was obtained using Cufflinks (version 2.1.1) (Trapnell et al., 2012).

**Sequence analyses**

To calculate GC content, MAC Mating type loci were cut every 50 bp using a sliding-window approach. Multiple sequence alignment was done using ClustalW (Larkin et al., 2007) and the results were visualized in ESPrint 3 (Robert and Gouet, 2014). Positional sequence...
similarity was calculated based on the alignment results and were averaged using a 20 aa sliding-window. To construct phylogenetic tree, protein sequences were aligned by ClustalW, and sites with over 50% gaps were deleted. Then the best-fit model was calculated by ProtTest (version 3.4.2) (Darriba et al., 2011), and the phylogenetic tree was generated by IQ-TREE (version 1.6.12) (Nguyen et al., 2015). Principal Components Analysis and protein clustering according to the percentage of amino acids were done by MeV (version 4.9.0) (Euclidean distance was used) (Saeed et al., 2003). ANOVA and t-test were done using GraphPad (version 8.0.2).

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### Table SE1. Primers used to fill introns and intergenic region sequence.

| Species          | Mating type | Primer       | Sequence (5’→3’)               |
|------------------|-------------|--------------|---------------------------------|
| **T. borealis**  | Y           | BorII-1183-F | TTACTAGCAACGAGCATTGATG          |
|                  |             | BorII-3109-F | AACCTAAAGGCCAAATTTTAAGAGAAG     |
|                  |             | BorII-2889-F | TAAAGGTTAATCGAGGTGGTGAATG       |
|                  |             | BorII-2889-Fs* | TGCGCAATCTCAGAGATAAGA          |
|                  |             | BorII-4346-F | TTAATAGGTTACCATAAAGCGG          |
|                  |             | BorII-4940-Rs* | TAACTTTTCACGCAGATTT            |
|                  |             | BorII-4940-R | CATCAGAAGAATCTCGTGAGGATG       |
|                  |             | BorII-6393-Rs* | GTGAAGGATAGTGAGATGGAAG         |
|                  |             | BorII-6217-R | AGAGTTAAGGAATCGAGGGATG         |
|                  |             | BorII-6217Fs* | GTATCTTATCTAAGCGATCAA          |
|                  |             | BorII-8515-R | AATAGGTTAGGCTGATGAGTAC         |
|                  |             | BorII-9760-R | TAGCAGCTAAATACAGATAGAAG         |
|                  | Z           | BorIII-1201-F | CATTGCATTGCTTATGCTTCTTAG        |
|                  |             | BorIII-3289-F | CAAATGCTTTATGAGCAAAGATG         |
|                  |             | BorIII-3027-F | TTTGTATTCCAATCTTGTAGCCG         |
|                  |             | BorIII-4846-R | GAGAATATCCGATATGAGGAAG         |
|                  |             | BorIII-4332-F | ATTTAGGATCGTTATGAGTGATG         |
|                  |             | BorIII-6428-R | TTAGAGTTAAGTACTGAGGCTAAC        |
|                  |             | BorIII-5500-F | TGTTACCCCTTCAGGTATTTTTT         |
|                  |             | BorIII-5500-Fs* | TTTGATGAAATGCTTGGCCCT          |
|                  |             | BorIII-7565-R | ATAGATAATTCTGCTTAGCCTTAG        |
|                  |             | BorIII-9070-Rs* | TACATTTCTTATTGGCTTC             |
|                  |             | BorIII-9070-R | TGCCCTTATGCTATTGTTGC           |
| **T. pigmentosa** | I           | Pig-1325-F   | TATCTCCCACTACATCCAGAAACAAG      |
|                  |             | Pig-3194-R   | CATCTGCTCTGATGATAACAAAGC       |
|                  |             | Pig-2861-F   | TCAATGCTCGTCATATTTAACT         |
|                  |             | Pig-2861-Fs* | ACCTGATAATTCGATTGACT           |
|                  |             | PigI-5069-Rs* | TCCTTTGTCAAAACGCA            |
|                  |             | PigI-5069-R  | GGAATATGAAATGCAAATCAGAG        |
|                  |             | PigI-4572-F   | CTGAATTTAAATGCTTTTGT           |
|                  |             | PigI-4572-Fs1* | TCCTAGTAGGATGACTCCTCAA        |
|                  |             | PigI-4572-Fs2* | ACTATGAGATACATGTTGT            |
|                  |             | PigI-4572-Fs3* | AGTTTAGCCTTTTCCTACGAGATG       |
|                  |             | Pig-6984-F   | TCCCTGACAATATCCGAAAC           |
|                  |             | Pig-8757-R   | TGTAGGATGGAGGGTTTGT            |
|                  | III         | Pig-1325-F   | TATCTCCCACTACATCCAGAAACAAG     |
|                  |             | Pig-3194-R   | ATCTGCTCTGATGATAATCAACG       |
|                  |             | Pig-2861-F   | TCAATGCTCGTCATATTTAACT         |
|                  |             | Pig-2861-Fs* | CTCAATCTCAGGGAAACTTAT          |
|   | Sequences                        |
|----|---------------------------------|
|   | PigIII-4214-F                  | TTCACCAATATTTAGTTATTAGCTG |
|   | PigIII-4214-Fs*                | ACTCAAATGAAGGATATGAAAAAC |
|   | PigIII-6856-R                  | TAAATGCTTGAGGAATAGTAATG  |
|   | PigIII-5750-F                  | AATCTGCCCTAAAGCAATGC    |
|   | PigIII-6986-R                  | GGTTCGGATAATTGTCAGG     |
|   | Pig-6984-F                     | TCCCTGACAATTATCCGAAAC   |
|   | Pig-8757-R                     | TGTAGGATTGGAGGAGGTTTG   |
| X | Ame-749-F                      | CCTAATGCTCCGATTTCTGC    |
|   | Ame-2196-R                     | GAAGTTATCCTGCCACTCAAAC  |
|   | Ame-2074-F                     | GATGATTAAGTGAACGTACTCAAGA|
|   | Amel-3341-R                    | GCCATGGTGTTGTTTCTTAGG   |
|   | Amel-3136-F                    | ATTTGCCTCACTAAATCAGATAT |
|   | Amel-3136-Fs*                  | CTCTAACTAAAACATTGGACTATCTT|
|   | Amel-4413-R                    | TATGCATTTAAACTAAGGGGTG  |
|   | Amel-4072-F                    | ACTACATTCTACACATACTCTTA |
|   | Amel-5013-R                    | TTCCTGATTTCAGATTTTG     |
|   | Amel-4877-F                    | CTATGGATGTGATGGAGACCTGT |
|   | Amel-5968-R                    | TTTATTTAACAGATGGGACAG   |
|   | Amel-5522-F                    | AGACCTGGATGCCCATAAGTATAG|
|   | Amel-6475-R                    | AGATGGAGCTGGCTTAGGT     |
|   | Amel-6358-F                    | AAGCATAGCCTCTGAAGATAAA  |
|   | Amel-8111-R                    | GCATTATTATTTATGGCATCTAC |
| Y | AmeII-4085-Rs*                 | TGAAGGCCCAGATTAATAGAAAT |
|   | AmeII-4085-R                   | GTGTTGGAATGACCCCTCTAT   |
|   | AmeII-3767-F                   | TCGAAATCTTCCTTCGATTCAG  |
|   | AmeII-3767-Fs*                 | CCACTACATTCTAATCCTACTTG |
|   | AmeII-4887-R                   | TTAAGAGAGTGGCTCATACACCC |
|   | AmeII-4723-F                   | TGAGAATAAGAGATGGAGCAG   |
|   | AmeII-5968-R                   | TTAATTTAACAGATGGAGCAG   |
|   | AmeII-5707-F                   | AATTAAAGCTTTTTGTAACGTT  |
|   | Ame-6475-R                     | AGATGGAGCTGCGTTAGGGT    |
|   | Ame-6358-F                     | AAGCATAGCCTCGAAGATAAA   |
|   | Ame-8111-R                     | GCATTATTTATTTATGGCATCTAC|

* Primers only used for sequencing.
| Species            | Strain       | Mating type | Sequence source          | Accession number | Reference(s)                              |
|--------------------|--------------|-------------|--------------------------|------------------|-------------------------------------------|
| *T. thermophila*   | SB4208       | II          | PCR & Transcriptome       | KC405255         | Cervantes et al, 2013                      |
|                    | SB4213       | III         | PCR & Transcriptome       | KC405261         | Cervantes et al, 2013                      |
|                    | SB4214       | IV          | PCR & Transcriptome       | KC405259         | Cervantes et al, 2013                      |
|                    | SB4218       | V           | PCR & Transcriptome       | KC405256         | Cervantes et al, 2013                      |
|                    | SB4220       | VI          | Genome                   | KC405258         | Eisen et al, 2006; Cervantes et al, 2013  |
|                    | SB4223       | VII         | PCR & Transcriptome       | KC405260         | Cervantes et al, 2013                      |
| *T. malaccensis*   | SD01608      | -           | Genome                   | MK315120         | Xiong et al, 2019                          |
| *T. pyriformis*    | GL           | -           | Genome                   | MK315121         | Xiong et al, 2019                          |
| *T. vorax*         | SD30421      | -           | Genome                   | MK315122         | Xiong et al, 2019                          |
| *T. borealis*      | SD01609      | X           | Genome                   | MK315123         | Xiong et al, 2019                          |
|                    | SD19502      | Y           | Transcriptome & PCR      | MK315124         | This report                               |
|                    | SD19803      | Z           | Transcriptome & PCR      | MK315125         | This report                               |
| *T. canadensis*    | SD30770      | -           | Genome                   | MK315126         | Xiong et al, 2019                          |
| *T. shanghaiensis* | SD205039     | -           | Genome                   | MK315127         | Xiong et al, 2019                          |
| *T. pigmentosa*    | SD19481      | X           | Transcriptome & PCR      | MK315128         | This report                               |
|                    | SD20427      | Y           | Transcriptome & PCR      | MK315129         | This report                               |
| *T. americanis*    | SD21194      | X           | Transcriptome & PCR      | MK315130         | This report                               |
|                    | SD21244      | Y           | Transcriptome & PCR      | MK315131         | This report                               |
| *T. paravorax*     | SD205177     | -           | Genome                   | MK315132         | Xiong et al, 2019                          |