An axiomatic basis for maximal homology alignment

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ABSTRACT

Maximal homology alignment is a new biologically-relevant approach to DNA sequence alignment that embraces microparalogy. It departs from the current method of gapped alignment, which uses a simple, binary state model of nucleotide position. In gapped alignment nucleotide positions have either no relationship (1-to-None) or else orthological relationship (1-to-1) with nucleotides in other sequences. Maximal homology alignment, however, allows additional states such as 1-to-Many and Many-to-Many, thus modeling both orthological and paralogical relationships, which together comprise the main homology types. Maximal homology alignment collects dispersed microparalogy into the same alignment columns on multiple rows, and thereby generates a two-dimensional representation of a single sequence. Sequence alignment then proceeds as the alignment of two-dimensional topological objects. The operations of producing and aligning two dimensional auto-alignments motivate a need for tests of two dimensional homological integrity that are both necessary and sufficient. Here, I work out and implement basic principles for computationally handling the two dimensions of positional homology, which are inherent to biological sequences due to replication slippage and related errors. I then show that maximal homology alignment is more suited and more informative than gapped alignment for modeling the evolution of genetic sequences, which harbor large numbers of small insertions and deletions originating as local microparalogy. These results show that both conserved and non-conserved genomic sequences are imbued with a signature of replication slippage that is absent from their random permutations.

KEYWORDS
dimensionality; DNA microfoam; Drosophila; enhancer DNAs; gapped alignment (GA); maximal homology alignment (MHA); positional homology; replication slippage; tandem repeats

Sequences of covalently-linked nucleotides and their evolutionary relationships are the basis of gene homology. The smallest possible scale to ascribe such homology is that of individual nucleotide positions (“site positional homology”). Of course this is possible only in the context of homological inference based on multiple nucleotide positions in a sequence.

By studying the function and evolution of regulatory DNA sequences (transcriptional enhancers in Ciona and Drosophila), which are not constrained by rigid, protein-coding, triplet reading frames (Erives et al. 1998; Erives and Levine 2000, 2004; Crocker et al. 2008; Erives 2009; Crocker et al. 2010; Crocker and Erives 2013; Brittain et al. 2014; Stroebele and Erives 2016), I conclude that site positional homology is incorrectly handled by gapped alignment (GA). GA was originally adopted by necessity in order to compare protein sequences of divergent lengths (Bryson and Vogel 1965; Braunitzer 1966; Needleman and Wunsch 1970; Sankoff 1972). In retrospect, GA is not designed to model how regulatory nucleotide sequences evolve for the reasons to be explained here. This issue is exacerbated to great extremes in the regulatory DNAs of the speciose Hawaiian Drosophila (Brittain et al. 2014; O’Grady and DeSalle 2018). Nonetheless, this issue also is seen for the sequences encoding many important regulator proteins enriched in polyglutamine content and other repeats, including the Notch intracellular domain and sub-units of the Mediator complex (Tóth-Petróczy et al. 2008; Fuxreiter et al. 2008; Rice et al. 2015; Stroebele and Erives 2016; Erives 2017). As a related consequence, repeat alleles at these loci are not being accurately genotyped by modern genome assembly despite evidence of profound phenotypic effects (Rice et al. 2015; Chandler et al. 2017; Press and Queitsch 2017; Press et al. 2019).

The premise of gapped alignment is perhaps non-biological because it does not treat one of the most important sources of genetic error that lead to alignment gaps: replication slippage (Strand et al. 1993; Haber and Louis 1998). Replication slippage...
is the result of DNA replication proceeding after a replication fork melts and re-anneals at an incorrect location thereby pro-
ducing short duplicated stretches of sequence related through
microhomology. Other enzymatic process, such as those that
occur during repair or recombination, can generate similar er-
ers (Haber 2000; Symington and Gautier 2011; Malkova and Ira
2013; Kowalczykowski 2015). Nonetheless, current techniques in
computational molecular biology are agnostic as to the source of
insertions and deletions (“indels”) and its homologous relation
to local sequence (Pevzner 2000).

The type of microhomology that is ignored by GA is more
precisely categorized as microparalogy because it is the result of
local duplication. Microparalogy is relevant to the indel prob-
lem of GA, which attempts to identify where null characters
should be inserted such that alignment is restored to homolo-
gous sequences. Thus the description of the goal of GA can be
summarized as the restoration of uniform sequence lengths of
one-dimensional strings by modeling sites as either orthologous
or not, with orthology being used as a proxy for homology of
all types. Although this has been known since the beginnings of
GA, it has not been well communicated or demonstrated how
unfortunate this has been to our understanding of sequence
homology at the smallest scale. Perhaps our present quandary
is the result of not having better methods available to model
microparalogy. This study formally describes an alternative to
GA that is correct, computable, and productive in expected and
unexpected ways.

Figure 1 gives a simple example as to why the search for a
master equation for indel placement (Holmes 2017) is a fruitless
quest and misguided. GA cannot find the best or optimal align-
ment because it does not allow microparalogy to be captured
into the same alignment columns. In Figure 1 a trinucleotide
repeat is variably expanded in two homologous sequences (Fig.
1A). Under GA these different sequences are aligned by the
placement of null characters in any number of possible locations
(Fig. 1B). This is not a contrived case because the majority of
indels are the result of replication slippage, which is exacerbated
within tandem repeats (TRs) (Ananda et al. 2013). Nonetheless,
different homologs do not have to be of different lengths or even
require the insertion of null characters for GA to still be deficient.
The reason is as follows. Even in the gapped alignment of identi-
cial sequences that harbor various TRs, microparalogy remains
de-localized over several columns despite their homological
equivalence. This further highlights that GA is more focused on
restoring length uniformity to character strings than modeling
the evolutionary homology of genetic sequences.

The presence of TRs is mutagenic with increasing repeat
number and prone to elevating the local rates for substitutions
and other errors (Amos 2010; Gemayel et al. 2010; Kelkar et al.
2011; Ananda et al. 2013; Duitama et al. 2014). Heterogeneous
mechanisms likely underlie the instability at TRs, including the
short microsatellite repeats (MSR, 1–9 bp units) and minisatel-
lites (10–50 bp units) (Ellegren 2000a,b; Legendre et al. 2007).
Within transcriptional enhancers some of this repeat content is
built up of ancient repeats of repeats with dynamic instability
(Crocker et al. 2010; Brittain et al. 2014). I refer to these more com-
plex, fractal-like patterns, which defy simple categorization as
direct tandem repeats, as “DNA microfoam”, from a contraction of
“micro-paralogy” and quantum spacetime foam (Wheeler 1957; Hawking 1978), and as an allusion to a common role for
gеometrical changes in dimensionality at the smallest scale.

To address the shortcomings of gapped alignment in not
considering local microparalogy, I developed, formalized, and
implemented “maximal homology alignment” (MHA) to rescue
microhomology that is unavoidably lost under GA. Figure 1C
shows how MHA rescues local microparalogy into the same
alignment columns avoiding any need to insert null characters
dashes). The basic operation of MHA is the “cinch” operation,
which lexically tokenizes a sequence into two paralogously over-
lapping sequences removed to separate rows (Fig. 1D). The cinch
operation thereby cinches, or tightens, the two-dimensional
width of a self-MHA, while increasing its height. Multiple se-
quence alignment (MSA) simply generalizes the cinch operation
to entire disjointed homologs. The robust implementation of
MHA presented here is able to (i) cinch identically perfect TRs,
(ii) cinch imperfect TRs as determined by a substitution model
and scoring system, (iii) cinch fractalized and/or foamy TRs in
which repeats differ in unit size due to variation in number of
sub-repeats, and (iv) resolve most overlapping but conflicting
repeats of the various types.

In implementing a working version of MHA in a program
called maximal, I discovered that necessary and sufficient computa-
tional tests for two dimensional homological integrity can be
derived from a few base principles. The utility of these principles
is that they show which desirable corollaries flow naturally with-
out having to stipulate or test for them outright. One can math-
ematically and computationally test the two-dimensionality of
homology with the tests described here. I then use MHA to
characterize the extent of replication slippage in functional vs.
non-functional biological sequences, and in cis-regulatory vs.
protein-coding sequences. These explorations reveal the extent
to which all biological sequences are imbued with a signature of
persistent replication slippage and the extent to which there is
negative selection to remove these errors from different logical
compartments of the genome. In short the example of biologi-
cal sequence can demonstrate how one dimension of position
becomes two-dimensions informatically.

Materials and Methods

Statistics

Development of the maximal code base included a built-in Fisher-
Yates shuffling option (Fisher and Yates 1953). Fisher-Yates shuf-
fing was used to generate random permutations of natural
sequences to use in millions of auto-MHA s over the course of
software development to identify DNA strings that were prob-
lematic to cinch. Scripts to handle these randomization runs and
to calculate width cinch ratios and related statistics from the
data in the output log files are available with the code base.

The Fisher-Yates option was also used to run the randomiza-
tion trials reported in Table 1 as controls. For these trials, the
software was recompiled fresh each time to stipulate compar-
able sequence lengths suitable for comparisons to the test sequences
(“FY_size”). Additional statistical tests are described in the
footnotes to Table 1.

Software development: maximal

The maximal software is built-around a UNIX-based program
written in C and which has many command line run options
that will be described in a separate publication, which can focus
on the full feature set and be more of a how-to manual. The
project’s code includes visualization and wrapping features to
allow for visualization and measurement of quality control met-
rics, such as the ability to recover an intact 1-D sequence after
A. DNA sequences
>seq. one ATGCAGCAGCAGCAGTAA
>seq. two ATGTCAGCAGCAGTGA

B. GAPPED ALIGNMENT
1 OF 10: seq. one >ATGCAGCAGCAGCAGTAA>
. ||||| ||||| |:
. seq. two >ATGCAGCAGCAGCAGTGA>

5 OF 10: seq. one >ATGCAGCAGCAGCAGTAA>
. ||||| ||||| |||||:
. seq. two >ATGCAGCAGCAGCAGTGA>

10 OF 10: seq. one >ATGCAGCAGCAGCAGTAA>
|| |||||| |||||:
seq. two >AT---GCAGCAGCAGTGA>

C. MAXIMAL HOMOLOGY ALIGNMENT (MHA)
1 OF 1: seq. one >ATGCAG/
....CAG/....CAG/....CAGTAA>
seq. two >ATGCAG/
....CAG/....CAGTGA>
consensus ATGCAGTRA

D. THE CINCH OPERATION OF MHA
Pre-cinch: >ATGCAGCAGCAGCAGTGA> y = 1
x = 123456789........

Post-cinch: >ATGCAG/
....CAG/....CAGTGA> y = 2
x = 123456789........

Figure 1 Gapped vs. maximal homology alignment. (A) The two homologous sequences shown will be aligned by GA (B) and MHA (C) for comparison. In the two homologs, a CAG trinucleotide repeat has variably expanded in number. (B) GA gives 10 possible alignments, differing by where a single parsimonious gap can be inserted in sequence two. There is no formal basis for favoring any one of these many alignments as optimal and any such choice would be artificial. All 10 gapped alignments are deficient in not encoding the multiple positions in microparalogical relationship with other letters in the same sequence and in the other sequence. Thus GA artificially inflates the true number of columns of positional homology four-fold to 12 alignment columns. (C) Shown are the same sequences aligned by MHA, which are read left to right, top to bottom. In contrast to GA, MHA captures the micro-paralogy between CAG repeats into just one set of three columns. There is only one top-scoring alignment and there is no need for inserting null characters at all. Self-MHA treatments often produce 2-D objects that are automatically aligned without comparison to one another. The beginning and end of each sequence are marked with the “>” symbol, which is a convention to make it easier to see where different sequences begin and end in an MHA. A slip symbol ("/”) is used to indicate an inferred replication slip and the continuation of the sequence on the next row. (D) MHA is based on the cinch operation. The cinch operation reduces the 2-D width along the x-axis while increasing its height along the y-axis. Dots are added to MHA sequences only to help visualize each alignment column, but otherwise carry no informational significance.

Figure 2 Overlapping repeats require special cinching for maintaining 2-D homological integrity. Shown is a 24 bp sequence containing two tandem repeat (TR) patterns that overlap. Both TRs are perfect in that the repeating units are exactly identical. (MHA in the maximal program also handles imperfect TRs.) The first repeat is a TR with a unit \( k = 3 \) size that repeats twice after the first unit \( r = 2 \). When the two additional repeats are cinched under MHA, they end up being tucked underneath the first unit \( k \)-mer. Thus, 6 bp are tucked away to produce a 2-D auto-alignment that is 18 bp in width. If the second TR pattern of \( k = 8 \) is also cinched without consideration of the previously cinched upstream TR, then one would get a “bad slip”, which is characterized by alignment columns with non-homological calling (yellow shaded columns). For example, the \( T \) in the second position of the sequence erroneously gets called as being paralogous to the letter at position 16, which happens to be an A. Likewise, but less obvious, the \( G \) in position three gets erroneously called as being paralogous to the \( G \) in position 17. While these letters are identical, they should not be in the same column because there is no basis for inferring a paralogous relationship between them. The correct way of restoring local microparalogy into the same alignment columns is shown at the bottom. Here, the \( k = 3 \) repeat pattern is “cyclicalized” as a repeat of CAG to a repeat of AGC in exchange for reducing the repeat number from \( r = 2 \) to \( r = 1 \). This allows the second \( k = 8 \) TR pattern to be cinched in a compatible manner for a final 2-D width of 12 bp. This final cinching represents a width cinch ratio (WCR) of \( 12/24 = 0.500 \). This also serves as an example of repeats of repeats, potentially created by one initial repeat’s instability.
2-D manipulations. The C code base is supplemented by UNIX-shell scripts to do a variety of tasks, such as run maximal on Fisher-Yates shuffled versions of endogenous sequences from several different Drosophila species with different genome sizes, or to do simple pair-wise MHA comparisons. Future versions will scale the software to handle larger DNA sequences with dynamic memory allocation, and to conduct more complicated multiple sequence alignments of 2-D MHA structures. Also future versions will improve upon speed as the current version is laden with useful software development and run reporting options.

**Organization of maximal repository files.** The software code base is organized by a surfing theme. Executable script files have a base name starting with “surfboard-“. Data sets are stored in the sub-directory called “waves/”. The exon sequences used in this study are stored with an example base name as follows “waves/animals/seal_vnd_ex3_Dmela.txt”. The enhancer sequences are stored with an example base name as follows: “waves/animals/shark_vnd_NEE_Dmela.txt”. Tricky randomly-shuffled strings that were once puzzles and that aided code development are named with information related to their origin in a sub-directory called “waves/tubespit/”. Any remaining tricky strings are stored in “waves/chowder/”. The present version of the software is also set up to automatically save any new tricky strings in a file called “waves/tubespit/”. Any remaining tricky strings in “waves/chowder/”. The present version of the software is also set up to automatically save any new tricky strings in a file called “waves/tubespit/”. Any remaining tricky strings in “waves/chowder/”. The present version of the software is also set up to automatically save any new tricky strings in a file called “waves/tubespit/”. Any remaining tricky strings in “waves/chowder/”. The present version of the software is also set up to automatically save any new tricky strings in a file called “Surf_wavereport.log”. Additional output files are produced for other options but these will be discussed in a publication that is more focused on the software and its usage.

**Two dimensional array strategy.** The following describes the initial two dimensional array aspects of maximal, which is now supplemented with structure type definitions defined by the MHA axiomatic formalism described further below.

Path traversal: The maximal implementation of MHA begins similarly to the dynamic programming strategy used in global (Needleman and Wunsch 1970) and local (Smith and Waterman 1981) alignment with the construction of a path box, which when traversed describes the cinching required to produce local self-alignment. This path box is filled with scoring values according to a substitution matrix. Although the current maximal software recognizes the difference between DNA, RNA, protein, and alphabetic (non-biological) strings, a substitution model is only coded for DNA at the moment. I use a nucleotide substitution matrix that gives 1/2 the maximum identity score to transition positions for every two-dimensional point, and they are inefficient to crawl over while looking for sequence. More importantly, two dimensional arrays do not take advantage of the axiomatic basis of MHA that is described here.

C code library functions for MHA structure types. Initial development versions of maximal exclusively used two dimensional arrays because these objects were simple to code, easy to print computationally, and because an elegant axiomatic basis for MHA was not at first appreciated. There are several downsides to using two dimensional arrays for MHA computation: they use a lot of memory, they do not store the one-dimensional positions for every two-dimensional point, and they are inefficient to overwrite while looking for sequence. More importantly, two dimensional arrays do not take advantage of the axiomatic basis of MHA that is described here.

The current C code has transitioned from exclusive use of two dimensional arrays to also using structure type definitions, which can store one-dimensional positions alongside two dimensional coordinates. The new structure type definitions allow for efficient testing of 2-D homological integrity (see The Principle of Continuity). A library of special functions (“check_tela”, “push_tela”, “get_1Dz”, “update_tela” and others) have been written and are in use (named for “tela” Spanish for a thin fabric, from Latin for thin web-like membrane). Future software versions will continue to expand the use of these data types and their special functionality for self-validation. For clarity, the structure coord array is called stringy[i] in maximal’s main() function but received and encoded as tela[i] in the library because the library functions can get calls from other structure arrays of the same coord type.
implementation of MHA, as well as earlier versions of the code base. Below, pseudo-code is provided as a sketch of how the main axioms of maximal homology alignment are implemented in structure type definitions of the C programming language. In this prototype, transition mutations are marked in the structure member stringy[i].t and derive from the equivalence symbols stored in the structure member stringy[n].e. Other linked structure members store x and y coordinates and letter characters. In this structure type definition, the structure array index n is the one-dimensional position, which as such requires no separate structure member to store. The "_tela" library functions are called with a simple pointer to the special structure coord type array as defined below.

```c
/* DECLARE A COORD-TYPE STRUCTURE ARRAY */
struct coord stringy[MAXROW];

/* PRINCIPLE OF CONTINUITY */
/* PRINCIPLE OF EQUIVALENCE */
/* DECLARE A STRUCTURE TYPE NAMED COORD */
struct coord {
    int x; /* COLUMN COORDINATE */
    int y; /* ROW COORDINATE */
    char c; /* CHARACTER SYMBOL: A,C,G,T */
    char t; /* CALLED TRANSITIONS */
    char e; /* EQUIVALENCE CLASS: R,Y */
}

/* DEFINE A STRUCTURE TYPE NAMED COORD */
int define_coord {
    int x; /* COLUMN COORDINATE */
    int y; /* ROW COORDINATE */
    char c; /* CHARACTER SYMBOL: A,C,G,T */
    char t; /* CALLED TRANSITIONS */
    char e; /* EQUIVALENCE CLASS: R,Y */
}

/* PRINCIPLE OF CONTINUITY */
/* PRINCIPLE OF EQUIVALENCE */

Data Availability

The maximal program, a working implementation of maximal homology alignment and application of the axioms developed here, and its code base are all available on github (http://github.com/microfoam/maximal.git) under a GNU public license. Also, available at the maximal github repository, is a large set of sequences identified as being initially refractory to cinching.

Results

A need for tests of 2-D homological integrity

Before defining the axiomatic principles underlying two-dimensional positional homology and its implementation for MHA, I describe the types of computational problems that are solved in an MHA program called maximal (see Figure 2). Many of these problems were not obvious until implementation of MHA was tackled. The types of operations that one has to do often demand that the resulting 2-D objects be checked for two dimensional homological "integrity". A desire to check for this integrity then led to the identification and simplification of the axiomatic logic of 2-D alignments.

Over slips and bad slips. Over slips are caused when a TR is cinched that overlaps a previously cinched TR. This results from a strategy of cinching sequence in a moving window that proceeds left to right (5' to 3'). Over slips are particularly problematic when the earlier cinch effectively moves the upstream letters of the second cinch to be downstream of the second cinches start along the x-axis. In the maximal program these are known as "bad slips" and are subsequently undone. Bad slips are undone in the maximal program because the second cinch will always be larger than the first cinch due to the nature of the search. Over slips that do not need to be undone are simply ones that are compatible with the new cinch provided the new x starting coordinate is properly adjusted.

Cyclization. Cyclization refers to the operation of shifting the starting frame of a cyclizable TR, which must necessarily have a minimum size of 2k + 1 where k is the TR's unit size. A cyclizable sequence would be something like 5'-CAGCAGC. In this sequence a repeating triplet can be cinched either as CAG at position four, or else as AGC at position five. In this example positions four and five are the starts of the second intact TR unit and where a new row would begin (i.e., a translational increment along the y-axis instead of along the x-axis). For this reason repeat number in this MHA implementation begins the count for the number of repeats only with a second unit repeat (r = 1, and the first unit could be considered repeat zero). This convention also avoids having to consider that every possible lexical tokenization of a sequence is a repeat of one unit.

In the case where there are no overlapping TRs, maximal simply takes the first frame. In the case of a bad slip, sometimes they can be cyclized rather than undone, a preferred operation in regions of microfoam. In maximal this is handled by the "cyclelize" module, which currently cyclizes k-mers of size two, which are the most common.

Microfoam. As TR instability increases with repeat number, many areas of repeats show signatures of generating repeats of sub-strings of the founding repeats, which is referred to as DNA microfoam here. The combination of the two overlapping repeats of Figure 2 is an example of the simplest type of the resulting DNA microfoam. Nonetheless, sometimes there is a base pattern in which the unit size is different only because of a variable repeat number of an inner constituent repeat. Importantly, different unit repeat sizes are frequently normalized in size in the (x-axis) consensus of a 2-D self-alignment. Thus repeats of repeats can often be cinched by looking for tandem repeats in the consensus row and cinching accordingly if the operation would maintain 2-D integrity. In maximal, this is handled by the cinch-d module, which cinches de novo repeats of repeats.

In short the possible operations of cinching both TRs and imperfect TRs have to deal with over slips, bad slips, cyclizations, and microfoam. Choosing which operations to do necessitates having available efficient tests of the validity of a proposed cinching operation. The following base axioms of MHA satisfy this need.

The Principle of Continuity

The principle of continuity states how two adjacent letters m and n = m + 1 from a one-dimensional biological sequence are related in the two dimensions that are required to model positional homology. I reserve the use of m and n to describe different one-dimensional positions, and reserve the use of x and y coordinates to describe the two-dimensional points of each one-dimensional position: m → (x_m,y_m) and n → (x_n,y_n). For clarity, I also reserve the use of the word "position" when referring to the unique one-dimensional location, and reserve the word "point" when referring to its (preferably unique) two-dimensional homological location, which is given in paired coordinates.
To describe the underlying principles of MHA, I will refer to biological homology mapping functions that will return either the $x$ or $y$ coordinates from the two-dimensional representation given the one dimensional position $n$. Two other mapping functions will also be defined. Computationally these mapping functions correspond to memory pointers to the data indexed and arrayed by the the separate members of the structure type array objects (see Materials and Methods). These mapping functions are discrete, non-linear functions and can be considered a type of pathological mathematical object (see Discussion: MHA widths are not additive). These mapping functions will be represented by blackboard type uppercase symbols ($\mathbb{X}$, $\mathbb{Y}$, $\mathbb{Z}$, and $\mathbb{C}$).

**Principle 1 (Continuity).** Given one-dimensional molecular positions $m$ and $n = m + 1$, and $\mathbb{X}$ and $\mathbb{Y}$ biological homology mapping functions, which return two-dimensional $x$ and $y$ coordinates respectively from one-dimensional positions, then one and only one of the following two cases can and must be true:

- **Case one (non-homology):**
  \[
  \mathbb{X}[n] = \mathbb{X}[m + 1] = \mathbb{X}[m] + 1
  \]
- **Case two (micro-paralogy):**
  \[
  \mathbb{Y}[n] = \mathbb{Y}[m + 1] = \mathbb{Y}[m] + 1
  \]

In short, adjacent letters increase in position incrementally by single steps in either the $x$ direction or the $y$ direction, but never in both directions at once. Furthermore, when a position is incremented in the $y$ direction ($\mathbb{Y}[m] \to \mathbb{Y}[m + 1] = \mathbb{Y}[m] + 1$), then the $x$ coordinate must decrease or at least not increase ($\mathbb{X}[m] \to \mathbb{X}[m + 1] \leq \mathbb{X}[m]$). The second principle, the principle of paralogical equivalence, explains the condition under which this can occur.

The Principle of Continuity immediately implies three useful corollaries: Corollary 1.1: Mutual exclusivity of coordinate points, Corollary 1.2: Existence of the $\mathbb{Z}$ mapping function, and Corollary 1.3: Guarantee of $y$-monotonicity.

**Corollary 1.1 (Mutual exclusivity of coordinate points).** Given one-dimensional molecular positions $m$ and $n$ such that $n > m$ (or simply $n \neq m$), then $n$ and $m$ cannot share the same set of $x$ and $y$ homological coordinates.

*Proof.* For $n > m$, let $d = n - m$. For $n \neq m$, let $n$ be the larger of the two positions, and $m$ be the smaller. From the principle of equivalence, it must follow that $\max(\mathbb{X}[i] - \mathbb{X}[i - 1], \mathbb{Y}[i] - \mathbb{Y}[i - 1]) = 1$. The reason for this is due to equations (1) and (3), which guarantee discrete one step movements in one axis from one letter to the next, and equations (2) and (4), which guarantee that when movement occurs in one axis, translational movement along the other axis is either zero or in the negative direction. Thus, if $d = 0$, then the $x$ and $y$ coordinates of $n$ could not have changed from those of position $m$, and it must be that $n = m$ (i.e., $n$ and $m$ cannot be different).

**Corollary 1.2 (Existence of the $\mathbb{Z}$ mapping function).** By discretely counting the maximum of homologous $\mathbb{X}$ and $\mathbb{Y}$ increments relative to molecular increments, we can define a mapping function $\mathbb{Z}$ that corresponds to one dimensional position. More succinctly,
given a one-dimensional position, \( n \), we can construct an equivalent homological mapping function based on two-dimensional coordinates:

\[
\mathbf{Z}[n] = \sum_{i=0}^{\infty} \max(X[i] - X[i-1], Y[i] - Y[i-1]) = n.
\]  

(5)

Proof. From equations (1) + (2) under case one of non-homology, we have that \( \max(1, 0) = 1 \). Alternatively from equations (3) and (4) under case two of micro-paralogy, we have that \( \max(-k, 1) = 1 \), where \( -k < 0 \) and \( k \) is the \( k \)-mer size of the unit repeat, which is tucked underneath on the next row with a corresponding slip in the \( x \) axis. So by keeping a running count in this way we get a mapping function \( \mathbf{Z} \) that behaves like a one-dimensional discrete position index but is based on two-dimensional biological homology (in conjunction with the Principle of Paralogical Equivalence).

With this definition of \( \mathbf{Z} \), we get a mapping function that counts two-dimensional homological position discretely, stepwise, and monotonically. This is something that we have taken for granted with nucleotide position, which always increases in discrete single steps and is monotonic in the \( 5' \) to \( 3' \) direction. This corollary only holds if we respect the two mutually-exclusive cases described by the Principle of Continuity.

Corollary 1.3 (Guarantee of \( y \)-monotonicity). Given one-dimensional molecular coordinates \( m \) and \( n \) such that \( n > m \), then

\[
\mathbf{Y}[n] \geq \mathbf{Y}[m].
\]  

(6)

This last corollary should be shocking in its lonesomeness: there can never be any such guarantee of monotonicity for the \( x \)-axis. This result can be understood as follows. When reading DNA sequence left to right (\( 5' \) to \( 3' \)) one can eventually discover that a sequence eventually completes a second unit of a tandem repeat, resulting in a slipping of the \( x \)-axis position and a loss of monotonicity.

In practice, such as in the maximal implementation of MHA, TR unit sizes are limited by a hemi-diagonal sparse matrix approach with a bandwidth of 200 bp (in the maximal cinch-t module; see Materials and Methods). This is a reasonable cut-off because the majority of TRs and more extensive microsatellite repeats (MSRs) are composed of units that are less than 100 bp and most of this is less than 10 bp (Ananda et al. 2013). In other words \( y \)-monotonicity is restored only at a larger scale where a moving average of \( x \)-positions are taken for a window of a certain size.

The Principle of Paralogical Equivalence

By definition there are no orthological relations in an auto- or self-MHA, so internal homological relationships within a single sequence are all paralogical. The essential principle of equivalence states that two letters can occupy the same alignment column (share the same \( x \) position) if and only if there is an assumption of paralogical equivalence (homology of position).

Principle 2 (Paralogical Equivalence). Given one-dimensional molecular coordinates \( m \) and \( n > m \), and the mapping function \( \mathbf{C} \), which returns the character letter from a one-dimensional position, if \( \mathbf{X}[n] = \mathbf{X}[m] \), then

\[
\mathbf{C}[n] \begin{cases} 
= \mathbf{C}[m] \text{ with probability } P_1, \\
\neq \mathbf{C}[m] \text{ but } \neq \mathbf{C}[n] \text{ with probability } P_2 < P_1, \\
\neq \mathbf{C}[m] \text{ with probability } P_3 < P_2
\end{cases}
\]  

(7)

such that \( P_1 + P_2 + P_3 = 1 \).

The principle of paralogical equivalence means that two one-dimensional positions can share the same \( x \) coordinate if and only if the positions are paralogous positions in the two sequences located at \( \mathbf{Y}[m] \) and \( \mathbf{Y}[n] \). This means that first and foremost the two one-dimensional positions share identical \( x \)-coordinates. Second, it means that this is due to membership in a paralogous string, which must satisfy some simple probabilistic criteria that led to the calling of micro-paralogy in the first place. As such it is highly likely that the letters are identical (\( P_1 \)) or else members of a preferred mutational equivalence class (\( P_2 \) transitions: purine \( \leftrightarrow \) purine and pyrimidine \( \leftrightarrow \) pyrimidine). Nonetheless, the letters may be different and belong to an improbable mutational class (\( P_3 \) transversions: purine \( \leftrightarrow \) pyrimidine), while being embedded in local micro-paralogy.

This principle states explicitly that we are establishing a primary for the determination of orthology for a site position (an empty slot wherein one can place a letter) after considering the identity or equivalence or non-equivalence of letters in a column and nearby columns. Nonetheless, determination of site orthology is of course highly dependent on sequence identity of neighboring letters. This principle may be a truism, but it is one that is required for axiomatic completeness.

A proposition for a “Principle of Precedence”

The principle of precedence is a proposition for relating and connecting maximal homology alignment to gapped alignment.

Principle 3 (Precedence). In aligning one-dimensional biological homologs, alignment proceeds first by auto-MHA (self-alignment), then by step-wise ordered multiple sequence alignment (MSA). During MSA additional cinching of divergent paralogy is first attempted. Relaxation of inferred micro-paralogy is attempted after that. Third and last the insertion of null characters (dashes) can be taken to finish the alignment if necessary. The insertion of null characters is reserved for columns containing unique non-paralogical sequence.

Justification: The purpose of maximal homology alignment is not to provide a master equation for indel placement in a gapped alignment. One could envision conducting gapped alignment and then considering local micro-paralogy at indels. However, the nature of micro-paralogy demands that all local sequence be considered, including sequences somewhat removed from the placement of a gap. This follows from the absence of a monotonicity guarantee in the \( x \)-axis unlike the situation for the \( y \)-axis (see Guarantee of \( y \)-monotonicity). Stated differently, the positions of gaps in a gapped alignment are not and can never be precise measurements of micro-paralogy.

It is expected that the cinching of inferred micro-paralogy might correspond to a false-positive cinching. Setting aside the possibility of determining the case of convergent evolution versus true homology, a false-positive cinch can occur when adjacent sequences independently evolve to be identical or nearly identical sequences. Under MHA, these sequences might be cinched as a type of false-positive micro-paralogy. Later, when conducting multiple sequence alignment, the position of false positive micro-paralogy might correspond to a place one would normally add null characters. However, because MHA is being employed, one has recourse to first relaxing the cinch and returning the cinched letters to the would-be gap. Thus for pair-wise or MSA, over-cinching is inherently not an issue in MHA and can be reserved as a primary source of fill characters, relegating null characters as a second, backstop source. For single or auto-MHA, the maximal program is equipped with an optional “relax-2D” function, which relaxes homopolymeric runs that did
not aid in the cinching of fractalized TRs of different unit sizes. This is sometimes preferable for display purposes of MHAs of aesthetically-pleasing proportions.

**MHA dispenses with the gap problem**

Figure 4 shows the process of pairwise MHA in comparison to current methodology. To demonstrate the power of MHA, I have chosen to do pairwise MHA for the edge of a conserved enhancer where the homology of sequence becomes difficult to discern in distantly-related Drosophila species. I am using the sequences for D. melanogaster and D. erecta, which began diverging from a common ancestor in the melanogaster subgroup about 10 million years ago (Powell 1997). The enhancer edge corresponds to the upstream flanking region of the NEE of the ventral nervous system defective (vnd) locus (see Fig. 5). For comparison standard global alignment-type GA and a genome browser reference alignment are shown to demonstrate the issues with various types of GA (Fig. 4A and 4B). Self-MHAs of each sequence with normal non-agressive MHA parameters are almost auto-aligned, while already capturing microparalogy into the same columns (yellow highlighted sequences in Fig. 4C). Under the proposed principle of precedence, true pair-wise alignment first attempts to restore alignment by capturing (cinching) divergent microparalogy. Divergent microparalogy would be sequence that failed to meet a threshold alignment score during self-MHA, but which is evidently microparalogical when its homologous sequence from another species is able to surpass threshold in comparison. This first source of alignment (divergent microparalogy) is sufficient to restore perfect alignment and avoids resorting to the relaxation of false-positive microparalogy and last the insertion of null characters. Figure 4 could rightly be called the first real DNA alignment under MHA. Unlike GA this pairwise MHA readily shows the location of microparalogy, which are sites more susceptible to instability and evolutionary divergence (see yellow highlighted columns in Fig. 4). This MHA also shows that substitutions are readily polarized without the aide of a third outgroup homolog (see cyan highlighted letters in Fig. 4). This is possible because these sites are all places where multiple (≥2) microparalogical sequences exist for comparison.

**2-D widths of genetic sequences vs. random permutations**

One can define a width cinch ratio (WCR) as the ratio of the final MHA cinch width (the 2-D width) to the starting sequence length (i.e., the 1-D width). A priori it is not clear how biological versus non-biological sequences would compare in their WCR values. On the one hand, biological sequences might be expected to have lower WCRs relative to randomly shuffled versions of the same sequences. This would be the case if the natural sequences are imbued with a deep signature of persistent replication slippage. On the other hand, it is possible that this signature is under intense negative selection. Tandem repeats and the fractal-like repeats of repeats referred to here as DNA microfoam are susceptible to instability and could be a target of persistent negative selection. If this latter case is likely, then we should be able to test this hypothesis by comparing functional biological sequences to non-functional biological sequences.

Table 1 shows the WCRs for a set of natural sequences (protein-coding exons or CDS’s, and developmental enhancers) from eight different species of Drosophila. These species are disparately related within the Sophophora subgenus of Drosophila and care was taken to remove closely-related sister species (sequences and scripts to run these tests are available with the

**Figure 4** MHA does not typically need gaps to align distant homologs of different lengths. **(A)** Example GA using global alignment (Needleman and Wunsch 1970). Sequences correspond to the upstream edges of the vnd NEE in Drosophila species that began diverging 10 Mya. This region is where it becomes difficult to recognize homology. Differences (highlighted in magenta) are unpolared in the absence of out-group sequences, an issue that is often not manifested in MHA when it occurs in microparalogy. **(B)** Example of the same window of sequence in a genome browser view from the perspective of the reference species D. melanogaster. Sequences present only in D. erecta cannot be shown in relation to the reference genome, and is an issue that is not manifested in MHA. **(C)** Normal self-MHA preparation of the enhancer edges is almost sufficient to bring the divergent sequences into alignment. More aggressive self-MHA preparation can produce perfect auto-alignment but is sometimes problematic at DNA microfoam “knots”. **(D)** Pair-wise alignment of the auto-MHAs. This alignment is the result of cinching divergent microparalogy in comparison to the homology (see text). The cyan highlighted sites are polarized substitutions based on the pair-wise alignment and later confirmed by consulting out-groups. These regions of microparalogy (yellow highlighting) are not identified in GA.
Figure 5 The vnd locus of Drosophila melanogaster with the three regions (red boxes) used in comparing functional vs. non-functional, and regulatory vs. protein-coding sequences. A 500 bp non-conserved intronic fragment was used as a proxy for a non-functional sequence relative to the conserved neurogenic ectodermal enhancer (NEE), which is located just downstream in the same intron. Specifically, a 640 bp core window of highly conserved sequence was used for the NEE. For comparison, a 956 bp protein-coding exon constituting the entire CDS of exon 3 was used. This region encodes the highly conserved DNA-binding homeodomain and NK2-specific domain of the Vnd transcription factor. Shown below are plots of the conservation across different Drosophila and Anopheles species based on the Multiz alignment blocks (Blanchette et al. 2004). Table 1 details how conservation in each of the three windows is correlated with larger (less pronounced) width cinch ratios (WCRs). Nonetheless, all three boxed regions are enriched in signatures of replication slippage relative to randomly shuffled sequences derived from each box.

maximal code repository.) In comparison to these sets of natural sequences (exons or enhancers), which have average WCRs of 0.401 and 0.434, respectively, randomly permuted sequences produced by Fisher-Yates shuffling of their D. melanogaster counterpart (exon or enhancer as appropriate) have much larger WCRs. The Drosophila exon set 1a is about −1.5 standard deviations below the average of 1000 permutations of the D. melanogaster vnd exon. Similarly, the Drosophila enhancer set 1b is about −1.0 standard deviations below the average of 1000 permutations of the D. melanogaster vnd enhancer.

To understand whether there is a basis for negative selection purifying functional sequences of replication slippage, I compared the upstream, non-conserved intronic region to the neurogenic ectodermal enhancer (NEE) of the vnd gene of D. melanogaster. Unlike the larger enhancer set 1b from several species, which encompasses the NEE and potentially extensive non-functional flanking sequences from different species with differently sized genomes, for this computational test I made the most contrasting comparison possible. I took the core 640 bp window of the NEE that is the most conserved across insects and compared it to the 500 bp window just upstream that is the least conserved (see Fig. 5). This 500 bp window is the largest obtainable relatively non-conserved stretch I could get without getting into the upstream exons or the downstream NEE. Table 1 shows that the non-functional intronic window has a WCR value (0.338) that is about 11% tighter than the core NEE (0.447), which occupies the same intron. In comparison, the protein-coding CDS of the third and terminal exon from the same locus, has a WCR that is smaller but more comparable to that of the functional (intronic) enhancer.

I also performed multiple trials of Fisher-Yates permutations based on each of the three windows, separately (x1000 each). Each of these permutations is thus matched by sequence composition and length to their biological counterparts. Comparison to the non-biological shuffling experiments shows that, the biological windows tested from the vnd locus are 0.32 (CDS), 0.52 (NEE), and 3.30 standard deviations below the average of their matched trials. This suggests that while all of the windows are imbued with some degree of replication slippage signature relative to matched random permutations, the degree is correlated to the (relative absence of) evolutionary conservation and/or constraint for that window.

These results also underscore the effect on WCR ratios in the small D. melanogaster genome relative to other species in the genus (Gregory and Johnston 2008). For example, the D. melanogaster vnd exon 3 has a WCR of 0.433, while the Sophophora average is 0.401, representing a five-fold increase in the number of standard deviations below the average from random shuffling (see Table 1). Thus, MHA cinching occurs to a much greater extent in the orthologous sequences from other Drosophila species, which tend to have larger genomes.

Discussion

The MHA Principle of Continuity and its corollaries together with the Principle of Paralogical Equivalence constitute the analogs to properties that we take for granted as being inherent to one-dimensional sequence, but which are not tenable in the two dimensions of biological sequence homology (see Fig. 3). It is an unchallenged fallacy that our goal is to build 1-to-1 alignments of molecular sequence for diverse tasks in computational molecular genetics. It is not, or at least should not be so. Our goal is instead to build alignments of positional homology, which must encompass both orthology and paralogy to be biologically relevant. The base axioms of maximal homology alignment are offered here as a foundation for biologically-relevant sequence alignment in way that is not possible in GA. Furthermore, by explicitly attempting to detail MHA on an axiomatic basis, I hope to highlight pathological aspects, which mark a departure from more familiar mathematical or topological objects (see section below: MHA widths are not additive).

In the proposed Principle of Precedence, I argue for the use of “cinch-able” microparalogy and “relax-able pseudo-paralogy” as the primary alignment operations, while relegating the insertion of non-biological null characters (dashes) as a secondary, backstop operation. As shown in this study, MHA can reduce the need to use any fill characters (both pseudo-paralogy and null characters) as it rescues the replication slippage errors that are primarily responsible for gaps under GA (Fig. 4). This initial assessment and proposal will have to be tested with more extensive use of MHA and various implementations of MHA.

By working out axiomatic principles of two-dimensional sequence homology, I identify computational tests that are necessary and sufficient for validating two dimensional homological representations and their underlying cinching operations (for example, see C pseudo-code in Materials and Methods, or functional C code in the repository for the maximal software project). For example, we would expect that different nucleotides in a DNA sequence should occupy different two-dimensional points in a 2-D auto-alignment. Initially, I thought this would be its own principle to be required and tested, a principle of exclusivity. But later, I saw that it was a only corollary of the principle of continuity along with the principle of y-monotonicity. So rather than have to test these three properties separately, one
only has to test the main principle of continuity with the aid of the principle of paralogical equivalence.

In this first study, I used the principles to complete the first working version of maximal and to show that biological sequences are imbued with a signature of replication slippage that is under negative/purifying selection with increasing functional constraint (Fig. 5 and Table 1). Future versions of maximal will become much speedier because various operations in two dimensional arrays will be increasingly replaced in all places by the more abstract operations on the described C structure types that allow axiomatic testing (see Materials and Methods). Currently this is done in the early cinch-t module, and its first application immediately reduced the maximal failure rate. The current version of maximal already uses both methods (two dimensional arrays and structure arrays) for historical reasons and for software development tests. For example, special programming functions to check homological integrity and to translate 2-D coordinates to 1-D positions were written and applied.

**MHA widths are not additive**

The corollaries stemming from Principle 1 all apply to static MHAs, and so y-monotonicity and the absence of \( x \)-monotonicity hold only for static MHAs. In the maximal implementation of MHA, we have the case of cinch-t operations reversing bad slips when considering new sequence from a moving window for initial cinching. This is trivial example of an MHA addition operation, in which we are adding new un-cinched sequence to an existing (or growing) MHA. What follows is an important note about additivity of MHAs in general.

The absence of \( x \)-monotonicity indicates that maximal homology alignment is a pathological mathematical object in which two dimensional width is not additive. The reasons for this have to do with 2-D width being a measurement of span along the \( x \)-axis, and with the nature of the absence of \( x \)-monotonicity. First, adding new sequence to a growing MHA might represent adding sequence that can be tucked under as additional microparalogy without having to change the existing rows. Second, adding new sequence might reveal a new completed repeat pattern at the edge, resulting in a smaller width for the existing MHA. Third, adding new sequence might reveal either a new set of cinching operation spanning the splice and/or a more optimal set of cinching operations that supersedes a previous set of operations. Thus, we can summarize the significance of MHA additivity with another mathematical statement.

**Corollary 3.1 (Non-additivity of homological widths).** Given two separate sequences \( a \) and \( b \), and defining \( a + b \) to be the new longer sequence formed by linking \( a \) and \( b \), then

\[
w_a + w_b \leq w_{a+b},
\]

where \( w_s \) is the two-dimensional (homological) width of sequence \( s \) of length \( n \). In terms of the homological mapping function \( \mathcal{X} \), the above relation is equivalent to \( \mathcal{X}[a_n] + \mathcal{X}[b_n] \leq \mathcal{X}[a_{n+b}] \). Therefore, MHA widths are not additive.

In the list of reasons leading to a new summed width that is less than the added widths, if \( w_a \) is the width of the existing/growing MHA and \( w_a \) is the width of the sequence being added to the \( 3' \) end of \( a \), then \( w_{a1} \to w_{a2} < w_{a1} \) in the first and third examples, and \( w_{a1} \to w_{a2} < w_{a1} \) in the second and third examples.

In this study I focused on constructing and making practical use of an axiomatic basis for MHA from biological considerations alone. Nonetheless, it will be interesting to study whether there are applicable theoretical treatments of low-dimensional topological objects, such as from mathematical knot theory [e.g., Blair and Tomova (2013)] or from models of quantum gravity in physics [e.g., Carlip (2017)]. If so there may exist additional mathematical tools that can be adapted for the alignment of two dimensional biological sequence homology.

**The evolutionary context of transversions**

There are many, obvious, possible ramifications of MHA from more accurate genome assembly and improved GWAS results to better understanding of enhancer evolution. In closing I will simply pose an unusual but interesting MHA-motivated question concerning the context of transversion mutations. Transitions are

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**Table 1 Width cinch ratios of biological vs. shuffled, and functional vs. non-functional sequences**

| Set | Description | \( N \) | WCR | \( \sigma \) | \( (x - \mu)/\sigma \) |
|-----|-------------|-----|-----|-----|----------------|
| 1a  | Exon three (HD-encoding) of \( vnd \) from eight species (962 bp avg.) | 8   | 0.401 | 0.015 | -1.45 |
| 1b  | Enhancers from six species (\( vnd \) NEEs and \( nab \) DWME, 996 bp avg.) | 7   | 0.434 | 0.021 | -0.98 |
| 2   | Conserved CDS from \( D. \ mel. \ vnd \) exon three (956 bp) | 1   | 0.426 | –     | -0.32 |
| 2R  | Fisher-Yates shuffled set #2 (956 bp each) | 1000 | 0.433 | 0.022 | –     |
| 3   | Conserved intronic enhancer from \( D. \ mel. \ vnd \) intron one (640 bp) | 1   | 0.447 | –     | -0.52 |
| 3R  | Fisher-Yates shuffled set #3 (640 bp each) | 1000 | 0.462 | 0.029 | –     |
| 4   | Non-conserved intronic region from \( D. \ mel. \ vnd \) intron one (500 bp) | 1   | 0.338 | –     | -3.30 |
| 4R  | Fisher-Yates shuffled set #4 (500 bp each) | 1000 | 0.434 | 0.029 | –     |

\( R = \) Fisher-Yates shuffled and sampled to the indicated length times the \# of indicated replicates (\( N \)).

\( a \) Trials of Fisher-Yates shuffled sets are ordered and numbered by increasing WCR.

\( d \) Post-cinch-d Width cinch ratio (WCR) (or average WCR for shuffled sets) using maximal version 3.64.

\( e \) Standard deviation

\( f \) Based on the standard deviation (\( \sigma \)) of 1000 random permutations of same sequence (sets 2–4) or based on the 1000 random permutations of the \( D. \ melanogaster \) exon (set 1a) or enhancer (set 1b).

\( g \) This is the exon encoding the conserved homeodomain and NKX2-specific sequence.

| Set | Description | \( N \) | WCR | \( \sigma \) | \( (x - \mu)/\sigma \) |
|-----|-------------|-----|-----|-----|----------------|
| 2   | Conserved CDS from \( D. \ mel. \ vnd \) exon three (956 bp) | 1000 | 0.433 | 0.022 | –     |
| 3   | Conserved intronic enhancer from \( D. \ mel. \ vnd \) intron one (640 bp) | 1000 | 0.462 | 0.029 | –     |
| 4   | Non-conserved intronic region from \( D. \ mel. \ vnd \) intron one (500 bp) | 1000 | 0.434 | 0.029 | –     |

\( R = \) Fisher-Yates shuffled and sampled to the indicated length times the \# of indicated replicates (\( N \)).

\( a \) Trials of Fisher-Yates shuffled sets are ordered and numbered by increasing WCR.

\( d \) Post-cinch-d Width cinch ratio (WCR) (or average WCR for shuffled sets) using maximal version 3.64.

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\( g \) This is the exon encoding the conserved homeodomain and NKX2-specific sequence.
These sequences are also nested for compaction (a built-in plated by canonical complementarity with local microparalogy. In the context of the MHA representation, it could indicate that the tetranucleotide sequence with one transition mismatch. In the letters). This transversion triggers the cinching of an imperfect 

\[ C \rightarrow \] of the two sequences. This is the result of a single transversion mutation in \( D. mauritiana \) relative to other closely-related species: \( C \rightarrow G \) (bold underlined \( G \) letter). Due to this substitution the auto-MHA process for the derived \( D. mauritiana \) sequence cinches up differently in the middle third of the alignment. This is due to the CTGA sequence forming an imperfect repeat with the upstream tetra-nucleotide CCGA. This might suggest that the apparent transversion may have been mistemplated and if so would actually represent microparalogy. As such the substitution is not so much a transversion in the sense that ionizing radiation or chemically-induced modification directly caused for a different letter to appear in this position. Rather the mutation would represent canonical complementary templating via a stretch of local sequence. Thus, MHA may allow for the in-depth study of the sequence context surrounding transversion mutations and the extent to which transversions are correlated to adjacent microparalogical content. It will then be informative to see whether this is more common during DNA replication, meiotic recombination, and/or various repair pathways.

**Figure 6** A transversion substitution potentially mistemplated by canonical complementarity with local microparalogy. Shown are the raw maximal homology alignments produced by the *maximal* software. These sequences correspond to the upstream edges of the beginnings of the neurogenic ectodermal enhancers (NEEs) from *Drosophila simulans* (top sequence) and *Drosophila mauritiana* (bottom sequence), which correspond to two closely related species that diverged \(< 250 \text{kya. These sequences are also nested for compaction (a built-in feature of maximal)}\). The MHA segment spanning the asterisked lines, inclusive of the lines, is the only segment where the cinching is different between the two sequences. This is the result of a single transversion mutation in *D. mauritiana* relative to other closely-related species: \( C \rightarrow G \) (bold underlined letters). This transversion triggers the cinching of an imperfect tetranucleotide sequence with one transition mismatch. In the context of the MHA representation, it could indicate that the transversion substitution may correspond to a transition substitution that was mis-templated from adjacent microparalogy.

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\[ ATCGC/ \]

\[ ---CT/ \]

\[ TCGC/ \]

\[ CTA/ \]

\[ AGCTC/ \]

\[ ..........CGACTGA/ \]

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\[ ..........|.....A/ \]

\[ D. simul. \]

\[ D. mauritiana \]
