

eXpress 0.90 BETA

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Generated by Doxygen 1.7.3

Wed Sep 14 2011 16:04:18

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# **Chapter 1**

# **Class Index**

# 1.1 Class Hierarchy

This inheritance list is sorted roughly, but not completely, alphabetically:

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FragMap	1
Fragment	4
FrequencyMatrix	6
MismatchTable	9
Parser	1
BAMParser	5
SAMParser	6
ParseThreadSafety	2
PosWeightTable	3
SeqWeightTable	7
ThreadedMapParser	0
Transcript	1
TranscriptTable	6

2 Class Index

# Chapter 2

# **Class Index**

# 2.1 Class List

Here are the classes, structs, unions and interfaces with brief descriptions:

BAMParser	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	•	5
BiasBoss																													6
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4 Class Index

# **Chapter 3**

# **Class Documentation**

# 3.1 BAMParser Class Reference

#include <mapparser.h>

Inheritance diagram for BAMParser:



## **Public Member Functions**

- BAMParser (BamTools::BamReader \*reader)
- bool next\_fragment (Fragment &f)

## 3.1.1 Detailed Description

The BAMParser class fills Fragment objects by parsing an input file in BAM format.

#### Author

Adam Roberts

## Date

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Definition at line 46 of file mapparser.h.

#### 3.1.2 Constructor & Destructor Documentation

## 3.1.2.1 BAMParser::BAMParser ( BamTools::BamReader \* reader )

BAMParser constructor opens the file

Definition at line 117 of file mapparser.cpp.

## 3.1.3 Member Function Documentation

## **3.1.3.1** bool BAMParser::next\_fragment ( Fragment & f ) [virtual]

a member function that loads all mappings of the next fragment

#### **Parameters**

f the empty Fragment to add mappings to

#### **Returns**

true if more reads remain in the BAM file, false otherwise

Implements Parser.

Definition at line 133 of file mapparser.cpp.

The documentation for this class was generated from the following files:

- · src/mapparser.h
- src/mapparser.cpp

## 3.2 BiasBoss Class Reference

#include <biascorrection.h>

## **Public Member Functions**

- BiasBoss (double alpha)
- void update\_expectations (const Transcript &trans)
- void update\_observed (const FragMap &frag, double mass)
- double get\_transcript\_bias (std::vector< double > &start\_bias, std::vector< double > &end\_bias, const Transcript &trans) const
- std::string to\_string () const
- void append\_output (std::ofstream &outfile) const

## 3.2.1 Detailed Description

The BiasBoss class keeps track of sequence-specific and positional bias. It allows for the bias associated with a given fragment end to be calculated, and for the bias parameters to be updated based on additional observations. All stored and returned values are in log space.

#### Author

Adam Roberts

#### **Date**

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Definition at line 208 of file biascorrection.h.

#### 3.2.2 Constructor & Destructor Documentation

#### 3.2.2.1 BiasBoss::BiasBoss ( double alpha )

**BiasBoss Constructor** 

#### **Parameters**

alpha	a double specifying the strength of the uniform prior (logged pseudo-counts
	for each paramater)

Definition at line 218 of file biascorrection.cpp.

#### 3.2.3 Member Function Documentation

## 3.2.3.1 void BiasBoss::append\_output ( std::ofstream & outfile ) const

a member function that outputs the positional and sequence-specific bias parameter matrices

#### **Parameters**

outfile	the file to append to

# 3.2.3.2 double BiasBoss::get\_transcript\_bias ( std::vector< double > & start\_bias, std::vector< double > & end\_bias, const Transcript & trans ) const

a member function that returns the 5' and 3' bias values at each position in a given transcript based on the current bias parameters

#### **Parameters**

start_bias	a vector containing the logged bias for each 5' start site in the transcript
end_bias	a vector containing the logged bias for each 3' end site in the transcript
trans	the transcript for which to calculate the logged bias

#### Returns

the product of the average 5' and 3' bias (logged)

Definition at line 284 of file biascorrection.cpp.

### 3.2.3.3 string BiasBoss::to\_string ( ) const

a member function that returns a string containing the observed positional nucleotide probabilities (non-logged) in column-major order (A,C,G,T)

#### Returns

the string representation of the observed probabilities

Definition at line 313 of file biascorrection.cpp.

## 3.2.3.4 void BiasBoss::update\_expectations ( const Transcript & trans )

a member function that updates the expectation parameters (sequence-specific and positional) assuming uniform expression of and accross the transcript's sequence

#### **Parameters**

trans the transcript to measure expected counts from
--

Definition at line 225 of file biascorrection.cpp.

#### 3.2.3.5 void BiasBoss::update\_observed ( const FragMap & frag, double mass )

a member function that updates the observed parameters (sequence-specific and positional) given a fragment mapping to a transcript and its logged probabilistic assignment

#### **Parameters**

frag	the fragment mapping
mass	the logged probabality of the mapping, which is the amount to update the
	observed counts by

Definition at line 245 of file biascorrection.cpp.

The documentation for this class was generated from the following files:

· src/biascorrection.h

• src/biascorrection.cpp

## 3.3 FLD Class Reference

#include <fld.h>

## **Public Member Functions**

- FLD (double alpha, size\_t max\_val, size\_t mean, size\_t std\_dev)
- size\_t max\_val () const
- double mean () const
- void add\_val (size\_t len, double mass)
- double pdf (size\_t len) const
- double tot\_mass () const
- std::string to\_string () const
- void append\_output (std::ofstream &outfile) const

## 3.3.1 Detailed Description

The FLD class keeps track of the observed fragment length distribution. It starts with a Gaussian prior with parameters specified by the arguments. A small "Gaussian" kernel is added for each observation. All mass values and probabilities are stored and returned in log space (except in to\_string).

Definition at line 21 of file fld.h.

## 3.3.2 Constructor & Destructor Documentation

## 3.3.2.1 FLD::FLD ( double alpha, size\_t max\_val, size\_t mean, size\_t std\_dev )

## **FLD** Constructor

#### **Parameters**

alpha double that sets the average pseudo-counts (logged)			
max_val	an integer that sets the maximum allowable FragMap length		
mean	a size_t for the mean of the prior gaussian dist		
std_dev	a size_t for the std dev of the prior gaussian dist		

Definition at line 21 of file fld.cpp.

#### 3.3.3 Member Function Documentation

## 3.3.3.1 void FLD::add\_val ( size\_t len, double mass )

a member function that updates the distribution based on a new FragMap observation

#### **Parameters**

len	an integer for the observed FragMap length
mass	a double for the mass (logged) of the observed FragMap

Definition at line 46 of file fld.cpp.

#### 3.3.3.2 void FLD::append\_output ( std::ofstream & outfile ) const

a member function that appends the FLD parameters to the end of the given file

#### **Parameters**

outfile the file to append to

#### 3.3.3.3 size\_t FLD::max\_val() const

a member function that returns the maximum allowed FragMap length

### Returns

max allowed FragMap length

Definition at line 41 of file fld.cpp.

## 3.3.3.4 double FLD::mean ( ) const

a member function that returns the mean FragMap length

#### Returns

mean observed FragMap length

Definition at line 74 of file fld.cpp.

#### 3.3.3.5 double FLD::pdf ( size\_t len ) const

a member function that returns the (logged) probability of a given FragMap length

## **Parameters**

len an integer for the FragMap length to return the probability of

#### Returns

(logged) probability of observing the given FragMap length

Definition at line 62 of file fld.cpp.

## 3.3.3.6 string FLD::to\_string ( ) const

a member function that returns a string containing the current distribution

#### Returns

space-separated string of probabilities ordered from length 0 to max\_val (non-logged)

Definition at line 79 of file fld.cpp.

## 3.3.3.7 double FLD::tot\_mass ( ) const

a member function that returns the (logged) number of observed fragmaps (including pseudo-counts)

## Returns

number of observed fragments

Definition at line 69 of file fld.cpp.

The documentation for this class was generated from the following files:

- src/fld.h
- src/fld.cpp

# 3.4 FragMap Struct Reference

#include <fragments.h>

## **Public Member Functions**

- int length () const
- PairStatus pair\_status () const

#### **Public Attributes**

- std::string name
- TransID trans\_id
- Transcript \* mapped\_trans

- std::string seq\_1
- std::string seq\_r
- int left
- int right
- int mate\_1
- bool left\_first

## 3.4.1 Detailed Description

The FragMap struct stores the information for a single (multi-)mapping of a fragment.

#### **Author**

Adam Roberts

#### **Date**

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Definition at line 36 of file fragments.h.

#### 3.4.2 Member Function Documentation

```
3.4.2.1 int FragMap::length ( ) const [inline]
```

a member function returning the length of the fragment according to this mapping note, that this result will be invalid if the fragment is single-end

## Returns

int length of fragment mapping

Definition at line 94 of file fragments.h.

### 3.4.2.2 PairStatus FragMap::pair\_status ( ) const [inline]

a member function returning whether the mapping is PAIRED, LEFT\_ONLY, or RIGHT\_ONLY LEFT\_ONLY denotes that the single read is not reverse complemented => its left end is the left fragment end RIGHT\_ONLY denotes that the single read is reverse complemented => its right end is the right fragment end

#### Returns

PairStatus the pair status of the mapping

Definition at line 105 of file fragments.h.

#### 3.4.3 Member Data Documentation

#### 3.4.3.1 int FragMap::left

a public int containing the 0-based leftmost coordinate mapped to in the transcript valid only if PairStatus is PAIRED or LEFT\_ONLY

Definition at line 67 of file fragments.h.

### 3.4.3.2 bool FragMap::left\_first

a public bool specifying that the "right" (second according to SAM flag) is reverse complemented when true and the "left" (first according to SAM flag) is reverse complemented when false in other words, the "left" read is truly left of the "right" read in transcript coordinate space when true

Definition at line 87 of file fragments.h.

## 3.4.3.3 Transcript\* FragMap::mapped\_trans

a public pointer to the transcript mapped to

Definition at line 51 of file fragments.h.

## 3.4.3.4 int FragMap::mate\_l

a public int containing the left position for the mate of the first read read in from the SAM file 0 if single-end fragment this is temporarily used to help find the mate, but is not important later on

Definition at line 80 of file fragments.h.

## 3.4.3.5 std::string FragMap::name

a public string for the SAM "Query Template Name" (fragment name)

Definition at line 41 of file fragments.h.

#### 3.4.3.6 int FragMap::right

a public int containing the position following the 0-based rightmost coordinate mapped to in the transcript valid only if PairStatus is PAIRED or RIGHT\_ONLY

Definition at line 73 of file fragments.h.

## 3.4.3.7 std::string FragMap::seq\_l

a public string containing the "left" read sequence (first according to SAM flag)

Definition at line 56 of file fragments.h.

# 3.4.3.8 std::string $FragMap::seq_r$

a public string containing the "right" read sequence (second according to SAM flag) Definition at line 61 of file fragments.h.

## 3.4.3.9 TransID FragMap::trans\_id

a public TransID for the transcript mapped to

Definition at line 46 of file fragments.h.

The documentation for this struct was generated from the following file:

• src/fragments.h

## 3.5 Fragment Class Reference

```
#include <fragments.h>
```

## **Public Member Functions**

- ∼Fragment ()
- bool add\_map\_end (FragMap \*f)
- const std::string name () const
- const size\_t num\_maps () const
- const std::vector< FragMap \* > & maps () const

## 3.5.1 Detailed Description

The Fragment class stores information for all multi-mappings of a single fragment. By design, only paired-end mappings of paired-end reads will be accepted. All mappings of single-end reads will be accepted.

### Author

Adam Roberts

## **Date**

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Definition at line 123 of file fragments.h.

#### 3.5.2 Constructor & Destructor Documentation

#### 3.5.2.1 Fragment::~Fragment ( )

Fragment destructor deletes all FragMap objects pointed to by the Fragment Definition at line 14 of file fragments.cpp.

#### 3.5.3 Member Function Documentation

## 3.5.3.1 bool Fragment::add\_map\_end ( FragMap \* f)

a member function that adds a new FragMap (single read at this point) to the Fragment if it is the first FragMap, it sets the Fragment name and is added to \_open\_mates, if the fragment is not paired, it is added to \_frag\_maps, otherwise, add\_open\_mate is called

#### **Parameters**

```
f the FragMap to be added
```

Definition at line 27 of file fragments.cpp.

```
3.5.3.2 const std::vector<FragMap*>& Fragment::maps ( ) const [inline]
```

a member function that returns FragMap multi-mappings of the fragment

#### Returns

a vector containing pointers to the FragMap multi-mappings

Definition at line 181 of file fragments.h.

```
3.5.3.3 const std::string Fragment::name ( ) const [inline]
```

a member function that returns the SAM "Query Template Name" (fragment name)

### Returns

the string SAM "Query Template Name" (fragment name)

Definition at line 169 of file fragments.h.

```
3.5.3.4 const size_t Fragment::num_maps() const [inline]
```

a member function that returns the number of multi-mappings for the fragment

#### Returns

number of multi-mappings for fragment

Definition at line 175 of file fragments.h.

The documentation for this class was generated from the following files:

- · src/fragments.h
- src/fragments.cpp

## 3.6 FrequencyMatrix Class Reference

#include <frequencymatrix.h>

#### **Public Member Functions**

- FrequencyMatrix ()
- FrequencyMatrix (size\_t m, size\_t n, double alpha)
- double operator() (size\_t i, size\_t j) const
- double operator() (size\_t k) const
- void increment (size\_t i, size\_t j, double incr\_amt)
- void increment (size\_t k, double incr\_amt)
- double arr (size\_t k) const
- double row (size\_t i) const

## 3.6.1 Detailed Description

The FrequencyMatrix class keeps track of the frequency parameters in order to allow for constant-time probability look-ups and updates. The table is rectangular to allow for multiple distributions to be stored in one FrequencyMatrix. Rows are distributions. All values are stored and returned in log space.

#### Author

Adam Roberts

#### **Date**

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Definition at line 24 of file frequencymatrix.h.

## 3.6.2 Constructor & Destructor Documentation

## $\textbf{3.6.2.1} \quad \textbf{FrequencyMatrix::} \textbf{FrequencyMatrix} \textbf{( )} \quad \texttt{[inline]}$

dummy constructor

Definition at line 51 of file frequencymatrix.h.

#### 3.6.2.2 FrequencyMatrix::FrequencyMatrix ( size\_t m, size\_t n, double alpha )

FrequencyMatrix constructor initializes the matrix based on the log of the given pseudocounts

#### **Parameters**

m	a size_t specifying the number of distributions (rows)
n	a size_t specifying the number of values in each distribution (columns)
alpha	a double specifying the intial psuedo-counts (un-logged)

Definition at line 16 of file frequencymatrix.cpp.

## 3.6.3 Member Function Documentation

## **3.6.3.1** double FrequencyMatrix::arr(size\_t k)const [inline]

a member function that returns the raw value stored at a given position of the flattened matrix

#### **Parameters**

,	· ·
lz	the array position
Λ.	the array position

#### Returns

a double specifying the raw value stored at the given position of the flattened matrix

Definition at line 96 of file frequencymatrix.h.

## 3.6.3.2 void FrequencyMatrix::increment ( size\_t i, size\_t j, double incr\_amt )

a member function to increase the mass of a given position in the matrix

#### **Parameters**

i	the distribution (row)
j	the value (column)
incr_amt	the logged amount to increase the mass by

Definition at line 37 of file frequencymatrix.cpp.

## 3.6.3.3 void FrequencyMatrix::increment ( size\_t k, double incr\_amt )

a member function to increase the mass of a given position in the flattened matrix

#### **Parameters**

k the array position
----------------------

<i>incr amt</i> the logged amount to increase the mass b	inc	cr	amt	the	logged	amount	to	increase	the	mass	b	v
--	-----	----	-----	-----	--------	--------	----	----------	-----	------	---	---

Definition at line 45 of file frequencymatrix.cpp.

## 3.6.3.4 double FrequencyMatrix::operator() ( size\_t k ) const

a member function to extract the logged probability of a given position in the flattened matrix

#### **Parameters**

1.	the amore position
K	the array position
	T T

## Returns

a double specifying the logged probability of the given position in the flattened matrix

Definition at line 31 of file frequencymatrix.cpp.

## 3.6.3.5 double FrequencyMatrix::operator() ( size\_t i, size\_t j ) const

a member function to extract the logged probability of a given position in the matrix

#### **Parameters**

i	the distribution (row)
j	the value (column)

## Returns

a double specifying the logged probability of the given value in the given distribution

Definition at line 24 of file frequencymatrix.cpp.

## **3.6.3.6** double FrequencyMatrix::row ( size\_t i ) const [inline]

a member function that returns the raw row sum

#### **Parameters**

i	the distribution (row)

#### Returns

a double specifying the raw row sum for the given distribution

Definition at line 103 of file frequencymatrix.h.

The documentation for this class was generated from the following files:

- src/frequencymatrix.h
- src/frequencymatrix.cpp

## 3.7 MismatchTable Class Reference

#include <mismatchmodel.h>

## **Public Member Functions**

- MismatchTable (double alpha)
- void activate (bool active=true)
- double log\_likelihood (const FragMap &f) const
- void update (const FragMap &, double mass)
- std::string to\_string () const
- void append\_output (std::ofstream &outfile) const

#### 3.7.1 Detailed Description

The MismatchTable class is used to store and update mismatch (error) parameters using a first-order Markov model based on nucleotide and position in a ride and to return likelihoods of mismatches in given reads. All values are stored and returned in log space.

#### **Author**

Adam Roberts

## Date

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Definition at line 24 of file mismatchmodel.h.

## 3.7.2 Constructor & Destructor Documentation

## 3.7.2.1 MismatchTable::MismatchTable ( double alpha )

MismatchTable constructor initializes the model parameters using the specified (non-logged) pseudo-counts.

#### **Parameters**

alpha	a double containing the non-logged pseudo-counts for parameter initializa-
	tion

Definition at line 18 of file mismatchmodel.cpp.

#### 3.7.3 Member Function Documentation

## 3.7.3.1 void MismatchTable::activate ( bool active = true ) [inline]

member function that 'activates' the table to allow its values to be used in calculating log-likelihoods when it is sufficiently burned-in

#### **Parameters**

active a boolean specifying whether to activate (true) or deactivate (false)

Definition at line 59 of file mismatchmodel.h.

#### 3.7.3.2 void MismatchTable::append\_output ( std::ofstream & outfile ) const

a member function that outputs the final model parameters in a tab-separated file the file has 1 row for each read position and the parameters are in columns indexed as (ref, prev, obs) in base 4 with A,C,G,T encoded as 0,1,2,3.

#### **Parameters**

file	stream to append to

Definition at line 85 of file biascorrection.cpp.

## 3.7.3.3 double MismatchTable::log\_likelihood ( const FragMap & f ) const

member function returns the log likeihood of mismatches in the mapping given the current error model paramaters

#### **Parameters**

f the fragment mapping to calculate the log likelihood for

#### Returns

the log likelihood of the mapping based on mismatches

Definition at line 25 of file mismatchmodel.cpp.

## 3.7.3.4 string MismatchTable::to\_string ( ) const

member function that returns a string containing a collapsed confusion matrix based on the model parameters for the first read

#### Returns

a space-separated string for the flattened, collapsed confusion matrix in row-major

format (observed value as rows)

Definition at line 113 of file mismatchmodel.cpp.

## 3.7.3.5 void MismatchTable::update ( const FragMap & f, double mass )

member function that updates the error model parameters based on a mapping and its (logged) mass

#### **Parameters**

f	the fragment mapping
mass	the logged mass to increase the parameters by

Definition at line 72 of file mismatchmodel.cpp.

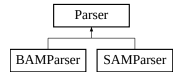
The documentation for this class was generated from the following files:

- src/mismatchmodel.h
- src/biascorrection.cpp
- src/fld.cpp
- src/mismatchmodel.cpp

## 3.8 Parser Class Reference

#include <mapparser.h>

Inheritance diagram for Parser:



## **Public Member Functions**

• virtual bool next\_fragment (Fragment &f)=0

## 3.8.1 Detailed Description

The Parser class is an abstract class that can be a SAMParser or BAMParser.

#### Author

Adam Roberts

#### Date

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Definition at line 27 of file mapparser.h.

#### 3.8.2 Member Function Documentation

## **3.8.2.1 virtual bool Parser::next\_fragment ( Fragment & f )** [pure virtual]

a member function that loads all mappings of the next fragment

#### **Parameters**

f the empty Fragment to add mappings to

#### Returns

true if more reads remain in the SAM/BAM file/stream, false otherwise

Implemented in BAMParser, and SAMParser.

The documentation for this class was generated from the following file:

· src/mapparser.h

# 3.9 ParseThreadSafety Struct Reference

```
#include <mapparser.h>
```

## **Public Attributes**

- Fragment \* next\_frag
- boost::mutex proc\_lk
- boost::mutex parse\_lk

## 3.9.1 Detailed Description

The ParseThreadSafety struct stores objects to allow for parsing to safely occur on a separate thread from processing.

## Author

Adam Roberts

#### Date

2011 Artistic License 2.0

Definition at line 126 of file mapparser.h.

#### 3.9.2 Member Data Documentation

## 3.9.2.1 Fragment\* ParseThreadSafety::next\_frag

a pointer to the next Fragment to be processed by the main thread Definition at line 131 of file mapparser.h.

#### 3.9.2.2 boost::mutex ParseThreadSafety::parse\_lk

a mutex to lock the parsing thread when the next\_frag pointer should be not modified Definition at line 141 of file mapparser.h.

### 3.9.2.3 boost::mutex ParseThreadSafety::proc\_lk

a mutex to lock the main (processing) thread when next\_frag has not yet been updated Definition at line 136 of file mapparser.h.

The documentation for this struct was generated from the following file:

• src/mapparser.h

# 3.10 PosWeightTable Class Reference

#include <biascorrection.h>

## **Public Member Functions**

- PosWeightTable (const std::vector < size\_t > &len\_bins, const std::vector < double > &pos\_bins, double alpha)
- const std::vector< size\_t > & len\_bins () const
- const std::vector< double > & pos\_bins () const
- void increment\_expected (size\_t len, double pos)
- void increment\_expected (size\_t l, size\_t p)
- void increment\_observed (size\_t len, double pos, double normalized\_mass)
- void increment\_observed (size\_t l, size\_t p, double normalized\_mass)
- double get\_weight (size\_t len, double pos) const
- double get weight (size t l, size t p) const
- void append\_output (std::ofstream &outfile) const

## 3.10.1 Detailed Description

The PosWeightTable class keeps track of fractional position bias parameters in log space. It allows for the bias associated with a given fractional position to be calculated, and for the bias parameters to be updated based on additional fragment observations.

#### **Author**

Adam Roberts

#### **Date**

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Definition at line 101 of file biascorrection.h.

#### 3.10.2 Constructor & Destructor Documentation

3.10.2.1 PosWeightTable::PosWeightTable ( const std::vector< size\_t > & len\_bins, const std::vector< double > & pos\_bins, double alpha )

PosWeightTable Constructor

#### **Parameters**

len_bins	a vector of unsigned integers specifying the bin ranges for transcript lengths
pos_bins	a vector of doubles specifying the bin ranges for fractional positions
alpha	a double specifying the strength of the uniform prior (logged pseudo-counts
	for each paramater)

Definition at line 123 of file biascorrection.cpp.

## 3.10.3 Member Function Documentation

## 3.10.3.1 void PosWeightTable::append\_output ( std::ofstream & outfile ) const

a member function that outputs the fractional position probabilities in matrix format with length bins as rows and fractional position bins as columns

## **Parameters**

outfile	the file to append to

## 3.10.3.2 double PosWeightTable::get\_weight ( size\_t len, double pos ) const

a member function that return the bias weight (logged) of a fractional transcript position

#### **Parameters**

len	the transcript length
pos	the fractional transcript position

#### Returns

the logged bias weight for the fractional transcript position

Definition at line 156 of file biascorrection.cpp.

## 3.10.3.3 double PosWeightTable::get\_weight ( size\_t I, size\_t p ) const

a member function that return the bias weight (logged) of a fractional transcript position bin

#### **Parameters**

l	the transcript length bin
p	the fractional transcript position bin

#### Returns

the logged bias weight for the fractional transcript position

Definition at line 55 of file biascorrection.cpp.

## 3.10.3.4 void PosWeightTable::increment\_expected ( size\_t I, size\_t p )

a member function that increments the expected counts for the given fractional position bin by 1 (logged)

#### **Parameters**

l	the transcript length bin
p	the fractional transcript position bin

Definition at line 137 of file biascorrection.cpp.

## 3.10.3.5 void PosWeightTable::increment\_expected ( size\_t len, double pos )

a member function that increments the expected counts for the given fractional position by  $1 \ (logged)$ 

## Parameters

len	the transcript length
pos	the fractional transcript position

Definition at line 130 of file biascorrection.cpp.

# 3.10.3.6 void PosWeightTable::increment\_observed ( size\_t len, double pos, double normalized\_mass )

a member function that increments the observed counts for the given fragment position by some mass (logged)

## **Parameters**

len	the transcript length
pos	the fractional transcript position
	the mass (logged probabilistic assignment) of the fragment normalized by
normalized	its estimated expression
mass	

Definition at line 143 of file biascorrection.cpp.

# 3.10.3.7 void PosWeightTable::increment\_observed ( size\_t l, size\_t p, double normalized\_mass )

a member function that increments the observed counts for the given fragment position bin by some mass (logged)

## **Parameters**

l	the transcript length bin
p	the fractional transcript position bin
	the mass (logged probabilistic assignment) of the fragment normalized by
normalized	its estimated expression
mass	

Definition at line 44 of file biascorrection.cpp.

The documentation for this class was generated from the following files:

- · src/biascorrection.h
- src/biascorrection.cpp

## 3.11 SAMParser Class Reference

#include <mapparser.h>

Inheritance diagram for SAMParser:



## **Public Member Functions**

- SAMParser (std::istream \*in)
- bool next\_fragment (Fragment &f)

## 3.11.1 Detailed Description

The SAMParser class fills Fragment objects by parsing an input in SAM format. The input may come from a file or stdin.

## Author

Adam Roberts

#### Date

2011 Artistic License 2.0

Definition at line 85 of file mapparser.h.

#### 3.11.2 Constructor & Destructor Documentation

## 3.11.2.1 SAMParser::SAMParser ( std::istream \* in )

SAMParser constructor removes the header, and parses the first line

Definition at line 183 of file mapparser.cpp.

#### 3.11.3 Member Function Documentation

#### 3.11.3.1 bool SAMParser::next\_fragment ( Fragment & f ) [virtual]

a member function that loads all mappings of the next fragment when the next fragment is reached, the current alignment is left in the \_frag\_buff for the next call

#### **Parameters**

f the empty Fragment to add mappings to

#### Returns

true if more reads remain in the SAM file, false otherwise

Implements Parser.

Definition at line 211 of file mapparser.cpp.

The documentation for this class was generated from the following files:

- src/mapparser.h
- · src/mapparser.cpp

## 3.12 SeqWeightTable Class Reference

#include <biascorrection.h>

## **Public Member Functions**

- SeqWeightTable (size\_t window\_size, double alpha)
- void increment\_expected (char c)
- void increment\_observed (std::string &seq, double normalized\_mass)
- double get\_weight (const std::string &seq, int i) const
- std::string to\_string () const
- void append\_output (std::ofstream &outfile) const

## 3.12.1 Detailed Description

The SeqWeightTable class keeps track of sequence-specific bias parameters. It allows for the bias associated with a given sequence to be calculated, and for the bias parameters to be updated based on additional observations. All values stored in log space.

#### **Author**

Adam Roberts

#### **Date**

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Definition at line 29 of file biascorrection.h.

### 3.12.2 Constructor & Destructor Documentation

#### 3.12.2.1 SeqWeightTable::SeqWeightTable ( size\_t window\_size, double alpha )

SeqWeightTable Constructor

## **Parameters**

window_size	an unsigned integer specifying the size of the bias window surrounding frag- ment ends
alpha	a double specifying the strength of the uniform prior (logged pseudo-counts
	for each paramater)

Definition at line 29 of file biascorrection.cpp.

### 3.12.3 Member Function Documentation

## 3.12.3.1 void SeqWeightTable::append\_output ( std::ofstream & outfile ) const

a member function that outputs the positional nucleotide probabilities in matrix format with nucleotides (A,C,G,T) as rows and window position as columns

#### **Parameters**

outfile	the file to append to

## 3.12.3.2 double SeqWeightTable::get\_weight ( const std::string & seq, int i ) const

a member function that calculates the bias weight (logged) of a bias window

#### **Parameters**

seq	the transcript sequence the fragment maps to
i	the fragment end point (the central point of the bias window)

#### Returns

the bias weight for the bias window which is the product of the individual nucleotide bias weights

## 3.12.3.3 void SeqWeightTable::increment\_expected ( char c )

a member function that increments the expected counts for the given nucleotide by 1 (logged)

### **Parameters**

c a char representing a nucleotide that has been observed in the transcriptome

Definition at line 34 of file biascorrection.cpp.

# 3.12.3.4 void SeqWeightTable::increment\_observed ( std::string & seq, double normalized\_mass

a member function that increments the observed counts for the given fragment sequence by some mass (logged)

### **Parameters**

	seq	a string of nucleotides in the bias window for the sequenced fragment end
		the mass (logged probabilistic assignment) of the fragment normalized by
norm	nalized	its estimated expression
	mass	

## 3.12.3.5 string SeqWeightTable::to\_string ( ) const

a member function that returns a string containing the positional nucleotide probabilities in column-major order (A,C,G,T)

#### **Returns**

the string representation of the positional nucleotide probabilities

Definition at line 68 of file biascorrection.cpp.

The documentation for this class was generated from the following files:

- · src/biascorrection.h
- src/biascorrection.cpp

# 3.13 ThreadedMapParser Class Reference

```
#include <mapparser.h>
```

#### **Public Member Functions**

- ThreadedMapParser (std::string input\_file)
- void threaded\_parse (ParseThreadSafety \*thread\_safety, TranscriptTable \*trans\_table)

## 3.13.1 Detailed Description

The ThreadedMapParser class is meant to be run on as a separate thread from the main processing. Once started, this thread will read input from a file or stream in SAM/BAM format, parse, and collect read alignments into fragment alignments, and fragment alignments into fragments, which are placed on a buffer for the processing thread. Once the processing thread copies the fragment address from the buffer, the parser is unlocked to load the next fragment. The process stops when EOF is reached

## Author

Adam Roberts

#### **Date**

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Definition at line 154 of file mapparser.h.

## 3.13.2 Constructor & Destructor Documentation

## 3.13.2.1 ThreadedMapParser::ThreadedMapParser ( std::string input\_file )

ThreadedMapParser constructor determines what format the input is in and initializes the correct parser.

Definition at line 51 of file mapparser.cpp.

#### 3.13.3 Member Function Documentation

## 3.13.3.1 void ThreadedMapParser::threaded\_parse ( ParseThreadSafety \* thread\_safety, TranscriptTable \* trans\_table )

a member function that drives the parse thread when all valid mappings of a fragment have been parsed, its mapped transcripts are found and the information is passed in a Fragment object to the processing thread through the ParseThreadSafety struct

#### **Parameters**

thread	a pointer to the struct containing shared locks and data with the processing
safety	thread
trans_table	a pointer to the table of Transcript objects to lookup the mapped transcripts

Definition at line 80 of file mapparser.cpp.

The documentation for this class was generated from the following files:

- src/mapparser.h
- src/mapparser.cpp

## 3.14 Transcript Class Reference

#include <transcripts.h>

## **Public Member Functions**

- Transcript (const std::string &name, const std::string &seq, double alpha, const FLD \*fld, const BiasBoss \*bias\_table, const MismatchTable \*mismatch\_table)
- const std::string & name () const
- TransID id () const
- const std::string & seq () const
- size\_t length () const
- double mass () const
- double var () const
- size\_t tot\_counts () const
- size\_t uniq\_counts () const
- void add\_mass (double p, double mass)
- void incr\_bundle\_counts (size\_t incr\_amt=1)
- size\_t bundle\_counts ()
- double log\_likelihood (const FragMap &frag) const
- double effective\_length () const
- double est\_effective\_length () const
- double unbiased\_effective\_length () const
- void update\_transcript\_bias ()

## 3.14.1 Detailed Description

The Transcript class is used to store objects for the transcripts being mapped to. Besides storing basic information about the object (id, length), it also stores a mass based on the number of fragments mapping to the object. To help with updating this number, it returns the likelihood that a given fragment originated from it. These values are stored and returned in log space.

#### Author

Adam Roberts

## **Date**

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Definition at line 57 of file transcripts.h.

## 3.14.2 Constructor & Destructor Documentation

3.14.2.1 Transcript::Transcript ( const std::string & name, const std::string & seq, double alpha, const FLD \* fld, const BiasBoss \* bias\_table, const MismatchTable \* mismatch\_table )

**Transcript** Constructor

#### **Parameters**

name	a string that stores the transcript name
seq	a string that stores the transcript sequence
alpha	a double that specifies the intial pseudo-counts (non-logged)
fld	a pointer to the global Fragment Length Distribution (FLD) object
bias_table	a pointer to the global BiasBoss object
mismatch	a pointer to the global MismatchTable object
table	

Definition at line 24 of file transcripts.cpp.

## 3.14.3 Member Function Documentation

## 3.14.3.1 void Transcript::add\_mass ( double p, double mass )

a member function that increases the expected fragment counts and variance by a given (logged) fragment mass

#### **Parameters**

p	a double for the (logged) probability that the fragment was generated by	
	this transcript	
mass	a double specifying the (logged) mass of the fragment being mapped	

Definition at line 41 of file transcripts.cpp.

## 3.14.3.2 size\_t Transcript::bundle\_counts() [inline]

a member function that returns the counts mapping to the bundle this transcript is in the total bundle counts is the sum of this value for all transcripts in the bundle

#### Returns

a portion of the counts mapping to the bundle this transcript is in

Definition at line 231 of file transcripts.h.

## 3.14.3.3 double Transcript::effective\_length ( ) const

a member function that calcualtes and returns the effective length of the transcript (non-logged)

#### Returns

the effective length of the transcript calculated as  $\tilde{l} = \sum_{l=1}^{L(t)} \sum_{i=1}^{L(t)} D(l) b_5[i] * b_3[i+l]$ 

Definition at line 110 of file transcripts.cpp.

## 3.14.3.4 double Transcript::est\_effective\_length ( ) const

a member function that calcualtes and returns the estimated effective length of the transcript (non-logged) using the avg bias

## Returns

the estimated effective length of the transcript calculated as  $\tilde{l}=b\bar{las}\sum_{l=1}^{L(t)}D(l)(L(t)-l+1)$ 

Definition at line 91 of file transcripts.cpp.

## 3.14.3.5 TransID Transcript::id ( ) const [inline]

a member function that returns the transcript id

## Returns

TransID transcript ID

Definition at line 172 of file transcripts.h.

#### 3.14.3.6 void Transcript::incr\_bundle\_counts ( size\_t incr\_amt = 1 ) [inline]

a member function that increases the counts mapping to the bundle this transcript is in the total bundle counts is the sum of this value for all transcripts in the bundle

#### **Parameters**

```
incr_amt a size_t to increase the counts by
```

Definition at line 221 of file transcripts.h.

```
3.14.3.7 size_t Transcript::length ( ) const [inline]
```

a member function that returns the transcript length

#### Returns

transcript length

Definition at line 183 of file transcripts.h.

## 3.14.3.8 double Transcript::log\_likelihood ( const FragMap & frag ) const

a member function that returns (a value proportional to) the log likelihood the given fragment originated from this transcript

#### **Parameters**

frag | a FragMap to return the likelihood of being originated from this transcript

## Returns

(a value proportional to) the log likelihood the given fragment originated from this transcript

Definition at line 57 of file transcripts.cpp.

```
3.14.3.9 double Transcript::mass ( ) const [inline]
```

a member function that returns the current (logged) fragment mass

## Returns

logged mass

Definition at line 189 of file transcripts.h.

## 3.14.3.10 const std::string& Transcript::name ( ) const [inline]

a member function that returns the transcript name

#### Returns

string containing transcript name

Definition at line 166 of file transcripts.h.

## 3.14.3.11 const std::string& Transcript::seq ( ) const [inline]

a member function that returns the transcript sequence

#### Returns

string containing transcript sequence

Definition at line 177 of file transcripts.h.

```
3.14.3.12 size_t Transcript::tot_counts() const [inline]
```

a member function that returns the current count of fragments mapped to this transcript (uniquely or ambiguously)

#### Returns

total fragment count

Definition at line 201 of file transcripts.h.

## 3.14.3.13 double Transcript::unbiased\_effective\_length ( ) const

a member function that calcualtes and returns the effective length of the transcript (non-logged) ignoring bias and using the prior FLD distribution

#### Returns

the effective length of the transcript calculated as  $\tilde{l} = \sum_{l=1}^{L(t)} D_{prior}(l) (L(t) - l + 1)$ 

Definition at line 105 of file transcripts.cpp.

```
3.14.3.14 size_t Transcript::uniq_counts ( ) const [inline]
```

a member function that returns the current count of fragments uniquely mapped to this transcript

## Returns

unique fragment count

Definition at line 207 of file transcripts.h.

#### 3.14.3.15 void Transcript::update\_transcript\_bias ( )

a member function that causes the transcript bias to be re-calculated by the \_bias\_table based on curent parameters

Definition at line 134 of file transcripts.cpp.

```
3.14.3.16 double Transcript::var ( ) const [inline]
```

a member function that returns the current (logged) variance

#### **Returns**

logged mass variance

Definition at line 195 of file transcripts.h.

The documentation for this class was generated from the following files:

- · src/transcripts.h
- src/transcripts.cpp

## 3.15 TranscriptTable Class Reference

```
#include <transcripts.h>
```

#### **Public Member Functions**

- TranscriptTable (const std::string &trans\_fasta\_file, double alpha, const FLD \*fld, BiasBoss \*bias\_table, const MismatchTable \*mismatch\_table)
- ~TranscriptTable ()
- Transcript \* get\_trans (TransID id)
- size\_t size () const
- void update\_covar (TransID trans1, TransID trans2, double covar)
- double get\_covar (TransID trans1, TransID trans2)
- size\_t covar\_size () const
- TransID get\_trans\_rep (TransID trans)
- TransID merge\_bundles (TransID rep1, TransID rep2)
- size\_t num\_bundles ()
- void output\_bundles (std::string output\_dir)
- void threaded\_bias\_update ()
- void output\_results (std::string output\_dir, size\_t tot\_counts, bool output\_varcov)

## 3.15.1 Detailed Description

The TranscriptTable class is used to keep track of the Transcript objects for a run. The constructor parses a fasta file to generate the Transcript objects and store them in a map that allows them to be looked up based on their string id.

#### Author

Adam Roberts

## Date

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Definition at line 285 of file transcripts.h.

## 3.15.2 Constructor & Destructor Documentation

```
3.15.2.1 TranscriptTable::TranscriptTable ( const std::string & trans_fasta_file, double alpha, const FLD * fld, BiasBoss * bias_table, const MismatchTable * mismatch_table )
```

TranscriptTable Constructor

## **Parameters**

trans	a string storing the path to the fasta file from which to load transcripts	
fasta_file		
alpha	a double that specifies the intial pseudo-counts for each bp of the transcripts	
	(non-logged)	
fld	a pointer to the global Fragment Length Distribution (FLD) object	
bias_table	a pointer to the global BiasBoss object	
mismatch	a pointer to the global MismatchTable object	
table		

Definition at line 142 of file transcripts.cpp.

## 3.15.2.2 TranscriptTable::~TranscriptTable ( )

TranscriptTable Destructor deletes all of the transcript objects in the table Definition at line 199 of file transcripts.cpp.

## 3.15.3 Member Function Documentation

## 3.15.3.1 size\_t TranscriptTable::covar\_size( ) const [inline]

a member function that returns the number of pairs of transcripts with non-zero covariance

#### Returns

the number of transcript pairs with non-zero covariance

Definition at line 368 of file transcripts.h.

## 3.15.3.2 double TranscriptTable::get\_covar ( TransID trans1, TransID trans2 )

a member function that returns the covariance between two transcripts these returned value will be the log of the negative of the true value

#### **Parameters**

trans1	one of the transcripts in the pair	
trans2	the other transcript in the pair	

#### **Returns**

the log of the negative of the pair's covariance

Definition at line 248 of file transcripts.cpp.

## 3.15.3.3 Transcript \* TranscriptTable::get\_trans ( TransID id )

a member function that returns a pointer to the transcript with the given id

## **Parameters**

Γ	. 1	
	ıa	of the transcript queried
	101	or the transcript queries

## Returns

pointer to the transcript wit the given id

Definition at line 227 of file transcripts.cpp.

## 3.15.3.4 TransID TranscriptTable::get\_trans\_rep ( TransID trans )

a member function that returns the bundle representative of the given transcript in the partitioning

## **Parameters**

trans	the TransID of the transcript whose representative is requested

#### Returns

the TransID of the representative for the bundle the given transcript is in

Definition at line 262 of file transcripts.cpp.

## 3.15.3.5 TransID TranscriptTable::merge\_bundles ( TransID rep1, TransID rep2 )

a member function that merges the bundles represented by the two given transcripts

## **Parameters**

rep1	the TransID of the first bundle representative
rep2	the TransID of the second bundle representative

#### Returns

the TransID of the representative for the new merged bundle

Definition at line 268 of file transcripts.cpp.

## 3.15.3.6 size\_t TranscriptTable::num\_bundles ( )

a member function that returns the number of bundles in the partition

#### Returns

the number of bundles in the partition

Definition at line 277 of file transcripts.cpp.

## 3.15.3.7 void TranscriptTable::output\_bundles ( std::string output\_dir )

a member function that outputs the bundles of the partition in a tab-delimited file called 'bundles.tab' in the given output directory each line contains a space-separated list of transcripts in a single bundle

## **Parameters**

output	<i>t_dir</i> the directory to outpu	t the bundle file to

Definition at line 288 of file transcripts.cpp.

## 3.15.3.8 void TranscriptTable::output\_results ( std::string output\_dir, size\_t tot\_counts, bool output\_varcov )

a member function that outputs the final expression data in a file called 'results.xprs' and (optionally) the variance-covariance matrix in 'varcov.xprs' in the given output directory

## **Parameters**

output_dir	the directory to output the expression file to	
tot_counts	the total number of observed mapped fragments	
output	boolean specifying whether to also output the variance-covariance matrix	
varcov		

Definition at line 368 of file transcripts.cpp.

## 3.15.3.9 size\_t TranscriptTable::size ( ) const [inline]

a member function that returns the number of transcripts in the table

## Returns

number of transcripts in the table

Definition at line 344 of file transcripts.h.

## 3.15.3.10 void TranscriptTable::threaded\_bias\_update( )

a member function for driving a thread that continuously updates the transcript bias values

Definition at line 308 of file transcripts.cpp.

## 3.15.3.11 void TranscriptTable::update\_covar ( TransID trans1, TransID trans2, double covar )

a member function that increases the covariance between two transcripts by the specified (logged) amount these values are stored positive even though they are negative

## **Parameters**

trans1	one of the transcripts in the pair	
trans2	the other transcript in the pair	
covar	a double specifying the (logged) amount to increase the pair's covariance	
	by	

Definition at line 235 of file transcripts.cpp.

The documentation for this class was generated from the following files:

- src/transcripts.h
- src/transcripts.cpp

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