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June 12, 2018

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1 Introduction

Methimpute implements a powerful HMM-based binomial test for methylation status calling. Besides improved accuracy over the classical binomial test, the HMM allows imputation of the methylation status of **all cytosines** in the genome. It achieves this by borrowing information from neighboring covered cytosines. The confidence in the methylation status call is reported as well. The HMM can also be used to impute the methylation status for binned data instead of individual cytosines. Furthermore, *methimpute* outputs context-specific conversion rates, which might be used to optimize the experimental procedure.

For the exact workings of *methimpute* we refer the interested reader to our publication at $\frac{https:}{doi.org/10.1186/s12864-018-4641-x}$.

2 Methylation status calling on individual cytosines

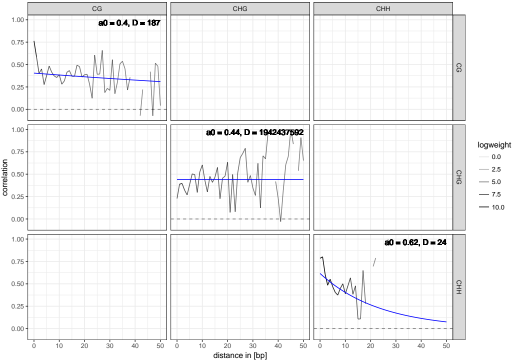
The following examples explain the necessary steps for methylation status calling (and imputation). To keep the calculation time short, it uses only the first 200.000 bp of the Arabidopsis genome. The example consists of three steps: 1) Data import, 2) estimating the distance correlation and 3) methylation status calling. At the end of this example you will see that positions without counts are assigned a methylation status, but the confidence (column "posteriorMax") is generally quite low for those cytosines, whereas it is high for well-covered cytosines (>=0.99).

2.1 Separate-context model

The separate-context model runs a separate HMM for each context. This assumes that only within-context correlations are important, and between-context correlations do not need to be considered.

```
library(methimpute)
# ===== Step 1: Importing the data ===== #
# We load an example file in BSSeeker format that comes with the package
file <- system.file("extdata", "arabidopsis_bsseeker.txt.qz", package="methimpute")</pre>
bsseeker.data <- importBSSeeker(file)</pre>
print(bsseeker.data)
## GRanges object with 110927 ranges and 2 metadata columns:
##
              segnames
                                ranges strand | context counts
##
                 <Rle>
                              <IRanges> <Rle> | <factor> <matrix>
##
          [1]
                  chr1
                               [34, 34]
                                            - 1
                                                        CHG
                                                                 0:4
##
          [2]
                  chr1
                               [80, 80]
                                              - |
                                                        CHH
                                                                 2:9
##
          [3]
                  chr1
                                [84, 84]
                                              + |
                                                        CHH
                                                                 1:1
          [4]
                                [85, 85]
                                                        CHH
##
                  chr1
                                                                 1:1
                                              + |
##
          [5]
                  chr1
                                [86, 86]
                                                        CHH
                                                                 1:1
##
                                                        . . .
                                                                  . . .
                  chr1 [533552, 533552]
                                                                 2:2
                                                         CG
##
     [110923]
                  chr1 [533554, 533554]
     [110924]
                                                         CG
                                                                 2:2
```

```
[110925] chr1 [533595, 533595]
                                         + |
                                                  CHG
                                                           0:1
## [110926] chr1 [533601, 533601]
                                          + |
                                                  CHG
                                                           0:2
##
    [110927] chr1 [533614, 533614]
                                          + |
                                                   CG
                                                           0:2
##
## seqinfo: 1 sequence from an unspecified genome; no seqlengths
# Because most methylation extractor programs report only covered cytosines,
# we need to inflate the data to inlcude all cytosines (including non-covered sites)
fasta.file <- system.file("extdata", "arabidopsis_sequence.fa.gz", package="methimpute")</pre>
cytosine.positions <- extractCytosinesFromFASTA(fasta.file,</pre>
                                             contexts = c('CG','CHG','CHH'))
methylome <- inflateMethylome(bsseeker.data, cytosine.positions)</pre>
print(methylome)
## GRanges object with 199978 ranges and 2 metadata columns:
                            ranges strand | context counts
##
         seqnames
##
               <Rle>
                           <IRanges> <Rle> | <factor> <matrix>
##
         [1]
                chr1
                             [1, 1]
                                        + |
                                                  CHH
##
         [2]
                chr1
                              [2, 2]
                                         + |
                                                  CHH
                                                           0:0
                                         + |
##
         [3]
                chr1
                              [3, 3]
                                                  CHH
                                                           0:0
             chr1
##
         [4]
                             [8, 8]
                                      + |
                                                  CHH
                                                           0:0
                             [9, 9]
##
        [5] chr1
                                        + |
                                                  CHH
                                                          0:0
##
        . . .
                               ... ...
## [199974] chr1 [533554, 533554] - |
                                                          2:2
                                                  CG
##
    [199975] chr1 [533557, 533557] + |
                                                  CHH
                                                          0:0
## [199976] chr1 [533560, 533560] + |
                                                  CG
                                                           0:0
## [199977] chr1 [533561, 533561]
                                         - |
                                                  CG
                                                           0:0
## [199978] chr1 [533565, 533565]
                                         - |
                                                  CHH
                                                           0:0
##
## seqinfo: 1 sequence from an unspecified genome
# ===== Step 2: Obtain correlation parameters ===== #
# The correlation of methylation levels between neighboring cytosines is an important
# parameter for the methylation status calling, so we need to get it first. Keep in mind
# that we only use the first 200.000 bp here, that's why the plot looks a bit meagre.
distcor <- distanceCorrelation(methylome, separate.contexts = TRUE)</pre>
fit <- estimateTransDist(distcor)</pre>
print(fit)
## $transDist
## CG-CG
                  CHG - CHG
                               CHH - CHH
## 1.866779e+02 1.942438e+09 2.374003e+01
##
## $plot
## Warning: Removed 23 rows containing missing values (geom_path).
```

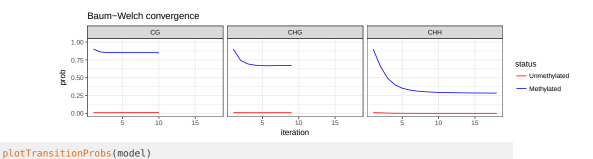


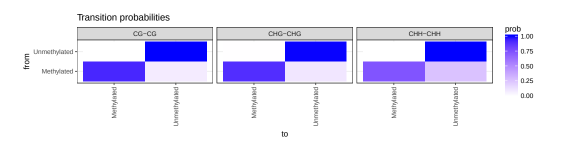
```
# ===== Step 3: Methylation status calling (and imputation) ===== #
model <- callMethylationSeparate(data = methylome, transDist = fit$transDist,</pre>
                                  verbosity = 0)
# The confidence in the methylation status call is given in the column "posteriorMax".
# For further analysis one could split the results into high-confidence
# (posteriorMax >= 0.98) and low-confidence calls (posteriorMax < 0.98) for instance.
print(model)
## GRanges object with 199978 ranges and 9 metadata columns:
##
              seqnames
                                  ranges strand | context counts distance
##
                  <Rle>
                               <IRanges> <Rle> | <factor> <matrix> <numeric>
##
          [1]
                  chr1
                                  [1, 1]
                                               + |
                                                        CHH
                                                                  0:0
                                                                            Inf
##
          [2]
                  chr1
                                  [2, 2]
                                                         CHH
                                                                  0:0
                                                                               0
                                  [3, 3]
##
          [3]
                  chr1
                                                         CHH
                                                                  0:0
                                                                               0
          [4]
                                  [8, 8]
##
                   chr1
                                                         CHH
                                                                  0:0
                                                                               4
##
          [5]
                   chr1
                                  [9, 9]
                                                         CHH
                                                                  0:0
                                                                               0
##
                   . . .
          . . .
                                      . . .
                                                         . . .
                                                                  . . .
##
     [199974]
                  chr1 [533554, 533554]
                                                         CG
                                                                  2:2
                  chr1 [533557, 533557]
##
     [199975]
                                                         CHH
                                                                  0:0
                                                                               8
##
     [199976]
                  chr1 [533560, 533560]
                                                         CG
                                                                  0:0
                                                                               5
##
     [199977]
                  chr1 [533561, 533561]
                                                         CG
                                                                  0:0
                                                                               0
##
     [199978]
                   chr1 [533565, 533565]
                                               - |
                                                         CHH
                                                                  0:0
##
              transitionContext posteriorMax posteriorMeth posteriorUnmeth
##
                        <factor>
                                    <numeric>
                                                   <numeric>
                                                                    <numeric>
##
          [1]
                            <NA>
                                    0.5004034
                                                   0.5004034
                                                                    0.4995966
                                                                    0.6301245
##
          [2]
                         CHH - CHH
                                    0.6301245
                                                   0.3698755
                                    0.7265858
##
          [3]
                         CHH - CHH
                                                   0.2734142
                                                                    0.7265858
##
          [4]
                         CHH-CHH
                                    0.7516365
                                                   0.2483635
                                                                    0.7516365
          [5]
                         CHH-CHH
                                    0.8163712
                                                   0.1836288
                                                                    0.8163712
```

```
##
##
     [199974]
                          CG-CG
                                   0.9999755
                                                  0.9999755
                                                               2.451116e-05
##
     [199975]
                        CHH-CHH
                                   0.7742228
                                                  0.2257772
                                                               7.742228e-01
##
     [199976]
                          CG-CG
                                   0.9057913
                                                  0.9057913
                                                               9.420875e-02
##
     [199977]
                          CG-CG
                                   0.8304996
                                                  0.8304996
                                                               1.695004e-01
##
     [199978]
                        CHH-CHH
                                   0.7480418
                                                  0.2519582
                                                               7.480418e-01
##
                    status rc.meth.lvl
##
                  <factor>
                            <numeric>
##
          [1]
               Methylated 0.14214505
##
          [2] Unmethylated 0.10509472
##
          [3] Unmethylated 0.07771417
##
          [4] Unmethylated 0.07060353
##
          [5] Unmethylated 0.05222859
##
##
     [199974]
                Methylated 0.85151435
##
     [199975] Unmethylated 0.06419243
                Methylated 0.77221618
##
     [199976]
##
                Methylated 0.70882454
     [199977]
     [199978] Unmethylated 0.07162390
##
##
##
     seqinfo: 1 sequence from an unspecified genome
# Bisulfite conversion rates can be obtained with
   model$params$emissionParams$Unmethylated
##
            prob
## CG 0.9904123
## CHG 0.9907659
## CHH 0.9998944
```

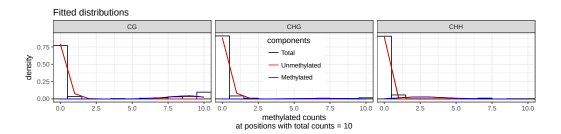
You can also check several properties of the fitted Hidden Markov Model, such as convergence or transition probabilities, and check how well the fitted distributions describe the data.

plotConvergence(model)





```
plotHistogram(model, total.counts = 10)
```

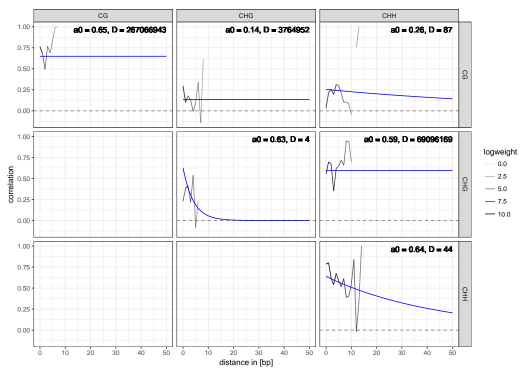


2.2 Interacting-context model

The interacting-context model runs a single HMM for all contexts. This takes into account the within-context and between-context correlations and should be more accurate than the separate-context model if sufficient data is available. However, we have observed that in low coverage settings too much information from well covered contexts is diffusing into the low covered contexts (e.g. CHH and CHG will look like CG with very low coverage). In this case, please use the separate-context model in section 2.1.

```
library(methimpute)
# ===== Step 1: Importing the data ===== #
# We load an example file in BSSeeker format that comes with the package
file <- system.file("extdata", "arabidopsis_bsseeker.txt.qz", package="methimpute")</pre>
bsseeker.data <- importBSSeeker(file)</pre>
print(bsseeker.data)
## GRanges object with 110927 ranges and 2 metadata columns:
##
                                   ranges strand | context
               segnames
                                                               counts
##
                  <Rle>
                                <IRanges> <Rle> | <factor> <matrix>
                   chr1
                                [34, 34]
##
          [1]
                                                         CHG
                                                                   0:4
                                               - |
          [2]
##
                                 [80, 80]
                                                         CHH
                                                                   2:9
                   chr1
                                               - 1
##
          [3]
                                 [84, 84]
                                                         CHH
                                                                   1:1
                   chr1
                                               + |
##
          [4]
                   chr1
                                 [85, 85]
                                               + |
                                                         CHH
                                                                   1:1
##
                                 [86, 86]
                                                         CHH
                                                                   1:1
          [5]
                   chr1
##
                                                         . . .
                                                                   . . .
##
     [110923]
                   chr1 [533552, 533552]
                                                          CG
                                                                   2:2
                   chr1 [533554, 533554]
                                                                   2:2
##
     [110924]
                                                          CG
                   chr1 [533595, 533595]
##
     [110925]
                                                         CHG
                                                                   0:1
                   chr1 [533601, 533601]
##
     [110926]
                                                         CHG
                                                                   0:2
##
     [110927]
                   chr1 [533614, 533614]
                                                          CG
                                                                   0:2
##
     seqinfo: 1 sequence from an unspecified genome; no seqlengths
# Because most methylation extractor programs report only covered cytosines,
# we need to inflate the data to inlcude all cytosines (including non-covered sites)
fasta.file <- system.file("extdata", "arabidopsis_sequence.fa.gz", package="methimpute")</pre>
cytosine.positions <- extractCytosinesFromFASTA(fasta.file,</pre>
                                                   contexts = c('CG','CHG','CHH'))
methylome <- inflateMethylome(bsseeker.data, cytosine.positions)</pre>
```

```
print(methylome)
## GRanges object with 199978 ranges and 2 metadata columns:
      seqnames ranges strand | context counts
##
              <Rle>
                          <IRanges> <Rle> | <factor> <matrix>
                                                       0:0
                                                       0:0
                                                       0:0
                                                       0:0
                                                       0:0
                                                       2:2
                                                       0:0
                                                       0:0
## [199977] chr1 [533561, 533561] - |
                                               CG
                                                       0:0
## [199978] chr1 [533565, 533565] - | CHH
                                                       0:0
##
    -----
## seqinfo: 1 sequence from an unspecified genome
# ===== Step 2: Obtain correlation parameters ===== #
# The correlation of methylation levels between neighboring cytosines is an important
# parameter for the methylation status calling, so we need to get it first. Keep in mind
# that we only use the first 200.000 bp here, that's why the plot looks a bit meagre.
distcor <- distanceCorrelation(methylome)</pre>
fit <- estimateTransDist(distcor)</pre>
print(fit)
## $transDist
## CG-CG
                  CG-CHG
                                         CHG - CHG
                              CG - CHH
                                                     CHG-CHH
                                                                 CHH-CHH
## 2.670669e+08 3.764952e+06 8.663764e+01 4.124081e+00 6.909617e+07 4.409969e+01
##
## $plot
## Warning: Removed 24 rows containing missing values (geom_path).
```



===== Step 3: Methylation status calling (and imputation) ===== # model <- callMethylation(data = methylome, transDist = fit\$transDist)</pre> Iteration log(P) dlog(P) Time in sec ## 0 -inf 0 1 -40631.538451 0 ## inf 2 -26304.340662 14327.197788 0 ## ## 3 -24210.680437 2093.660225 ## 4 -23635.136896 575.543541 ## -23374.140217 260.996679 -23224.427468 149.712749 ## 6 ## 7 -23134.096595 90.330873 1 ## 8 -23079.925956 54.170639 1 9 -23047.280535 32.645421 1 ## ## 10 -23027.416404 19.864131 ## 11 -23015.215509 12.200895 ## 12 -23007.665767 7.549743 ## 13 -23002.971021 4.694745 ## 14 -23000.044357 2.926664 1 ## 15 -22998.218617 1.825740 1 ## 16 -22997.079103 1.139514 1 ## 17 -22996.365832 0.713271 1 ## HMM: Convergence reached! # The confidence in the methylation status call is given in the column "posteriorMax". # For further analysis one could split the results into high-confidence # (posteriorMax >= 0.98) and low-confidence calls (posteriorMax < 0.98) for instance. print(model) ## GRanges object with 199978 ranges and 9 metadata columns:

```
segnames
                            ranges strand | context counts distance
              <Rle>
                         <IRanges> <Rle> | <factor> <matrix> <numeric>
##
        [1]
             chr1
                          [1, 1]
                                    + | CHH
                                                      0:0 Inf
##
        [2]
            chr1
                           [2, 2]
                                     + |
                                              CHH
                                                      0:0
                                                                 0
                                    + | CHH
+ | CHH
+ | CHH
        [3] chr1
                           [3, 3]
##
                                                      0:0
                                                                 0
        [4] chr1
                           [8, 8]
                                                      0:0
                                                               4
##
             chr1
                                                               0
##
        [5]
                           [9, 9]
                                                      0:0
    [199974] chrl [533554, 533554] - | CG
[199975] chrl [533557, 533557] + | CHH
        ... ...
974] chr1 [533554, 533554]
##
                                                      . . .
                                                               . . .
                                                           0 2
##
                                                      2:2
                                              CHH
##
                                                      0:0
                                    + | CG
                                                               2
    [199976] chr1 [533560, 533560]
                                                      0:0
##
           chr1 [533561, 533561] - | CG
chr1 [533565, 533565] - | CHH
                                              CG
                                                               0
##
                                                      0:0
    [199977]
##
    [199978]
                                                      0:0
                                                               3
##
           transitionContext posteriorMax posteriorMeth posteriorUnmeth
##
                   <factor> <numeric> <numeric>
                                                       <numeric>
##
        [1]
                     <NA>
                              0.6606448 0.3393552
                                                       0.6606448
                              0.7068376 0.2931624
##
        [2]
                   CHH - CHH
                                                      0.7068376
                              0.7471539 0.2528461
                    CHH - CHH
##
        [3]
                                                      0.7471539
                   CHH - CHH
                              0.7624977 0.2375023
        [4]
                                                      0.7624977
##
                              0.7963517 0.2036483
                   CHH - CHH
                                                      0.7963517
##
        [5]
                              ...
##
        . . .
                      . . .
                                          . . .
                                                            . . .
                                       0.9999923
##
    [199974]
                     CG-CG
                              0.9999923
                                                     7.668704e-06
                   CG-CHH
                              0.5029823 0.4970177
##
    [199975]
                                                     5.029823e-01
    [199976]
                   CHH - CG
##
                              0.5294277 0.4705723
                                                     5.294277e-01
##
                     CG-CG 0.5595388 0.4404612 5.595388e-01
    [199977]
##
    [199978]
                   CG-CHH
                              0.7689333 0.2310667
                                                     7.689333e-01
##
                 status rc.meth.lvl
##
               <factor> <numeric>
     [1] Unmethylated 0.11618208
##
##
      [2] Unmethylated 0.10047755
      [3] Unmethylated 0.08677090
##
       [4] Unmethylated 0.08155434
##
        [5] Unmethylated 0.07004477
##
##
        . . .
             . . . .
    [199974] Methylated 0.8238826
##
    [199975] Unmethylated 0.1697838
##
##
    [199976] Unmethylated 0.3906849
##
    [199977] Unmethylated 0.3660465
##
    [199978] Unmethylated 0.0793664
##
## seqinfo: 1 sequence from an unspecified genome
# Bisulfite conversion rates can be obtained with
1 - model$params$emissionParams$Unmethylated
          prob
## CG 0.9943607
## CHG 0.9979215
## CHH 0.9991911
```

3 Methylation status calling on binned data

The following examples explain the necessary steps for methylation status calling (and imputation) on binned data, such as commonly used 100bp bins. To keep the calculation time short, it uses only the first 200.000 bp of the Arabidopsis genome. The example consists of four steps: 1) Data import, 2) binning and 3) methylation status calling.

```
library(methimpute)
# ===== Step 1: Importing the data ===== #
# We load an example file in BSSeeker format that comes with the package
file <- system.file("extdata", "arabidopsis_bsseeker.txt.gz", package="methimpute")</pre>
bsseeker.data <- importBSSeeker(file)</pre>
print(bsseeker.data)
## GRanges object with 110927 ranges and 2 metadata columns:
         seqnames ranges strand | context counts
##

      <Rle>
      <IRanges>
      <Rle> | <Tactor>

      [1] chr1
      [34, 34]
      - | CHG

      [2] chr1
      [80, 80]
      - | CHH

      [3] chr1
      [84, 84]
      + | CHH

      [4] chr1
      [85, 85]
      + | CHH

      [5] chr1
      [86, 86]
      + | CHH

      ...
      ...
      ...

##
                 <Rle>
                               <IRanges> <Rle> | <factor> <matrix>
##
##
                                                                    2:9
##
                                                                   1 · 1
                                                                   1:1
##
##
                                                         CHH
                                                                  1:1
##
                                                                   . . .
## [110923] chr1 [533552, 533552]
                                                          CG
                                                                   2:2
                                             - |
##
    [110924] chr1 [533554, 533554]
                                                          CG
                                                                   2:2
## [110925] chr1 [533595, 533595]
                                              + |
                                                                    0:1
                                                          CHG
    [110926] chr1 [533601, 533601]
##
                                                + |
                                                          CHG
                                                                    0:2
##
     [110927]
                  chr1 [533614, 533614]
                                                           CG
                                                                    0:2
##
     seqinfo: 1 sequence from an unspecified genome; no seqlengths
# Because most methylation extractor programs report only covered cytosines,
# we need to inflate the data to inlcude all cytosines (including non-covered sites)
fasta.file <- system.file("extdata", "arabidopsis_sequence.fa.gz", package="methimpute")</pre>
cytosine.positions <- extractCytosinesFromFASTA(fasta.file,</pre>
                                                   contexts = c('CG','CHG','CHH'))
methylome <- inflateMethylome(bsseeker.data, cytosine.positions)</pre>
print(methylome)
## GRanges object with 199978 ranges and 2 metadata columns:
          seqnames ranges strand | context counts
##
                 <Rle>
                              <IRanges> <Rle> | <factor> <matrix>
               chr1
##
          [1]
                                 [1, 1] + |
                                                         CHH
                                                                    0:0
          [2] chr1
##
                                 [2, 2]
                                                + |
                                                          CHH
                                                                    0:0
##
          [3] chr1
                                 [3, 3] + |
                                                          CHH
                                                                    0 • 0
##
          [4] chr1
                                 [8, 8] + |
                                                          CHH
                                                                    0:0
##
          [5] chr1
                                  [9, 9] + |
                                                          CHH
                                                                    0:0
##
                   . . .
                                                          . . .
                                                                    . . .
          . . .
    [199974] chr1 [533554, 533554]
                                                          CG
                                                                    2:2
##
     [199975] chr1 [533557, 533557]
                                                + |
                                                          CHH
                                                                    0:0
                                                + |
    [199976] chr1 [533560, 533560]
                                                                    0:0
##
                                                          CG
     [199977]
                  chr1 [533561, 533561]
##
                                                - |
                                                           CG
                                                                    0:0
     [199978] chr1 [533565, 533565]
                                                          CHH
                                                                    0:0
```

```
## ----
## seqinfo: 1 sequence from an unspecified genome
# ===== Step 2: Binning into 100bp bins ===== #
binned Methylome <- binMethylome (methylome, binsize = 100, contexts = c('total', 'CG'))
print(binnedMethylome$CG)
## GRanges object with 5335 ranges and 3 metadata columns:
##
      seqnames ranges strand | cytosines context counts
                         <IRanges> <Rle> | <integer> <factor> <matrix>
##
##
       [1]
              chr1
                       [ 1, 100] * | 0
                                                         CG
                                                                7:19
       [2] chr1
##
                       [101, 200]
                                       * |
                                                  6
                                                          CG
                                                                41:62
                                   * | 6

* | 0

* | 2

* | 1

....

* | 0

* | 14

* | 10
      [3] chr1 [201, 300]
[4] chr1 [301, 400]
[5] chr1 [401, 500]
##
                                                          CG
                                                                3:58
##
                                                          CG
                                                                 4:43
##
                                                          CG
                                                                 0:19
##
               . . .
                              . . .
                                                          . . .
                                                                  . . .
       . . .
    [5331] chr1 [533001, 533100] chr1 [533101, 533200]
##
                                                          CG
                                                                 0:55
##
                                                          CG
                                                                3:171
    [5333] chr1 [533201, 533300]
##
                                                          CG
                                                                 0:16
    [5334] chr1 [533301, 533400]
                                                                 0:35
##
                                                 2
                                                          CG
                                       * |
##
    [5335]
              chr1 [533401, 533500]
                                                   8
                                                          CG
                                                                 1:44
                                        *
##
## seqinfo: 1 sequence from an unspecified genome
# ===== Step 3: Methylation status calling (and imputation) ===== #
binnedModel <- callMethylation(data = binnedMethylome$CG)</pre>
## Iteration
                         log(P)
                                           dlog(P)
                                                     Time in sec
                           -inf
                  -27231.884280
##
          1
                                              inf
          2
                  -16567.652577
                                    10664.231703
##
                  -14617.459829
##
          3
                                     1950.192748
                  -13264.059065
                                     1353.400764
##
          4
##
          5
                  -12272.683299
                                       991.375766
##
          6
                  -11836.442261
                                       436.241038
                                      190.496922
##
          7
                  -11645.945339
          8
##
                  -11556.833035
                                        89.112304
                  -11524.535678
          9
                                       32.297356
##
         10
                  -11514.865975
##
                                        9.669703
                                                              0
         11
                  -11512.066277
                                        2.799698
##
                  -11511.303950
          12
                                        0.762327
## HMM: Convergence reached!
print(binnedModel)
## GRanges object with 5335 ranges and 10 metadata columns:
##
           seqnames
                            ranges strand | cytosines context counts
              <Rle>
##
                          <IRanges> <Rle> | <integer> <factor> <matrix>
##
       [1]
              chr1
                       [ 1, 100]
                                    * | 0
                                                          CG
                                                                7:19
                                  * | ...

* | 1

...

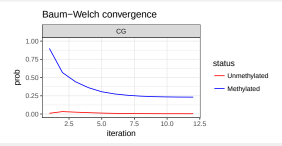
* | 0

* | 7
                       [101, 200]
##
       [2] chr1
                                                           CG
                                                                41:62
##
              chr1
                       [201, 300]
                                                 0
                                                          CG
                                                                3:58
       [3]
                        [301, 400]
                                                  2
##
       [4]
              chr1
                                                          CG
                                                                4:43
       [5]
                        [401, 500]
##
              chr1
                                                          CG
                                                                 0:19
##
               . . .
                                                                  . . .
##
    [5331]
              chr1 [533001, 533100]
                                                          CG
                                                                 0:55
    [5332]
              chr1 [533101, 533200]
                                                          CG
                                                                3:171
##
    [5333]
              chr1 [533201, 533300]
                                                          CG
                                                                 0:16
```

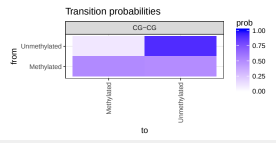
##	[5334]	chr1 [533	3301, 533400]	*	2 CG	0:35
##	[5335]	chr1 [533	3401, 533500]	*	8 CG	1:44
##		distance tra	ansitionContext	posteriorMax	posteriorMeth	posteriorUnmeth
##		<numeric></numeric>	<factor></factor>	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>
##	[1]	Inf	<na></na>	1.0000000	1.00000000	3.278506e-109
##	[2]	0	CG-CG	1.0000000	1.00000000	6.634096e-70
##	[3]	0	CG-CG	0.6422146	0.35778544	6.422146e-01
##	[4]	0	CG-CG	0.9742299	0.97422989	2.577011e-02
##	[5]	0	CG-CG	0.9649472	0.03505281	9.649472e-01
##						
##	[5331]	Θ	CG-CG	1.0000000	4.023375e-08	1.0000000
##	[5332]	Θ	CG-CG	1.0000000	1.362313e-15	1.0000000
##	[5333]	Θ	CG-CG	0.9990175	9.824911e-04	0.9990175
##	[5334]	0	CG-CG	0.9999928	7.221815e-06	0.9999928
##	[5335]	0	CG-CG	0.9999085	9.153180e-05	0.9999085
##			rc.meth.lvl			
##			<numeric></numeric>			
##	[1]	•	0.23143841			
##	[2]	•	0.23143841			
##		Unmethylated				
##	[4]	•	0.22558103			
##		Unmethylated	0.01211214			
##						
##	-	Unmethylated				
##		Unmethylated				
##		Unmethylated				
##		Unmethylated				
##		Unmethylated	0.0041656/3			
##			6			
##	seqinfo	o: I sequence	from an unspec	itied genome		

You can also check several properties of the fitted Hidden Markov Model, such as convergence or transition probabilities, and check how well the fitted distributions describe the data. This last point is important because the binomial distributions that the HMM uses were originally meant to describe individual cytosines and not bins. However, we have observed that they still capture the bimodal distributions of methylation levels in binned data quite well. Note that the histogram for our example looks quite sparse due to the very low number of bins that were used.

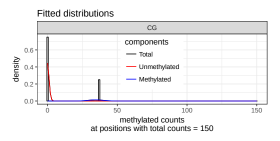
plotConvergence(binnedModel)



plotTransitionProbs(binnedModel)



plotHistogram(binnedModel, total.counts = 150)



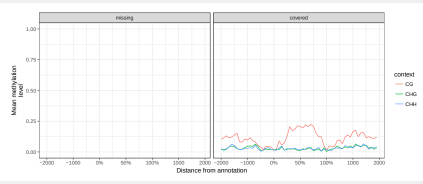
4 Description of columns in the output

- context The sequence context of the cytosine.
- counts Counts for methylated and total number of reads at each position.
- distance The distance in base-pairs from the previous to the current cytosine.
- transitionContext Transition context in the form "previous-current".
- posteriorMax Maximum posterior value of the methylation status call, can be interpreted as the confidence in the call.
- posteriorMeth Posterior value of the "methylated" component.
- posteriorUnmeth Posterior value of the "unmethylated" component.
- status Methylation status.
- **rc.meth.lvl** Recalibrated methylation level, calculated from the posteriors and the fitted parameters (see ?methimputeBinomialHMM for details).

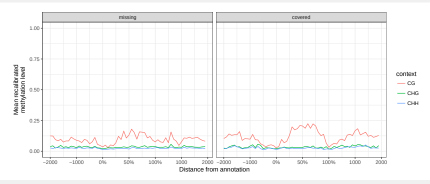
5 Plots and enrichment analysis

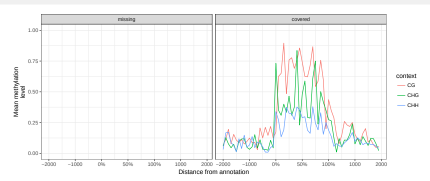
This package also offers plotting functions for a simple enrichment analysis. Let's say we are interested in the methylation level around genes and transposable elements. We would also like to see how the imputation works on cytosines with missing data compared to covered cytosines.

```
# Define categories to distinguish missing from covered cytosines
model$data$category <- factor('covered', levels=c('missing', 'covered'))
model$data$category[model$data$counts[,'total']>=1] <- 'covered'
model$data$category[model$data$counts[,'total']==0] <- 'missing'</pre>
```

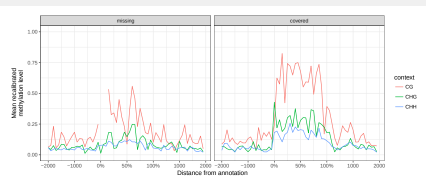


\$rc.meth.lvl





##
\$rc.meth.lvl



6 Export results

You can export the results as TSV file with the following columns:

 chromosome, position, strand, context, counts.methylated, counts.total, posteriorMax, posteriorMeth, posteriorUnmeth, status, rc.meth.lvl

exportMethylome(model, filename = tempfile())

Please see section 4 for a description of the columns.

7 Session Info

toLatex(sessionInfo())

- R version 3.4.3 (2017-11-30), x86_64-apple-darwin15.6.0
- Locale: C/UTF-8/C/C/C/C
- Running under: macOS High Sierra 10.13.4
- Matrix products: default
- $\hbox{\bf BLAS: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib}$
- LAPACK: /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: BiocGenerics 0.24.0, GenomeInfoDb 1.14.0, GenomicRanges 1.30.3, IRanges 2.12.0, S4Vectors 0.16.0, ggplot2 2.2.1, methimpute 1.3.1
- Loaded via a namespace (and not attached): BiocStyle 2.6.1, Biostrings 2.46.0, GenomeInfoDbData 1.0.0, RCurl 1.95-4.10, Rcpp 0.12.16, XVector 0.18.0, backports 1.1.2, bitops 1.0-6, colorspace 1.3-2, compiler 3.4.3, data.table 1.10.4-3, digest 0.6.15, evaluate 0.10.1, grid 3.4.3, gtable 0.2.0, highr 0.6, htmltools 0.3.6, knitr 1.19, labeling 0.3, lazyeval 0.2.1, magrittr 1.5, minpack.lm 1.2-1, munsell 0.4.3, pillar 1.1.0, plyr 1.8.4, reshape2 1.4.3, rlang 0.1.6, rmarkdown 1.8, rprojroot 1.3-2, scales 0.5.0, stringi 1.1.6, stringr 1.2.0, tibble 1.4.2, tools 3.4.3, yaml 2.1.16, zlibbioc 1.24.0