

Computation of melting temperature of nucleic acid duplexes with **rmelting**

Aravind, J.¹ and Krishna, G. K.²

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1. Division of Germplasm Conservation, ICAR-National Bureau of Plant Genetic Resources, New Delhi.
 2. Division of Crop Physiology, ICAR-Indian Agricultural Research Institute, New Delhi.
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1 Introduction

The R package `rmelting` is an interface to the **MELTING 5** program (Le Novère, 2001; Dumasseau et al., 2012) to compute melting temperatures of nucleic acid duplexes (DNA/DNA, DNA/RNA, RNA/RNA or 2'-O-MeRNA/RNA) along with other thermodynamic parameters such as hybridisation enthalpy and entropy.

Melting temperatures are computed by Nearest-neighbour methods for short sequences or approximative estimation formulae for long sequences. Apart from these, multiple corrections are available to take into account the presence of Cations (Na, Tris, K and Mg) or denaturing agents (DMSO and formamide).



2 Installation

The package can be installed from Bioconductor as follows.

```
if (!"BiocManager" %in% rownames(installed.packages()))
  install.packages("BiocManager")
BiocManager::install("rmelting")
```

The development version can be installed from github as follows.

```
if (!require('devtools')) install.packages('devtools')
devtools::install_github("aravind-j/rmelting")
```

Then the package can be loaded as follows.

```
library(rmelting)
```

3 Basic usage

Melting temperatures are computed in **rmelting** through the core function **melting** which takes a number of arguments (see **?melting**). The following are the essential arguments which are mandatory for computation.

- **sequence**
 - 5' to 3' sequence of one strand of the nucleic acid duplex as a character string. Recognises A, C, G, T, U, I, X_C, X_T, A*, AL, TL, GL and CL (**Table 1**). U and T are not considered identical.

Table 1: Recognized sequences

Code	Type
A	Adenine
C	Cytosine
G	Guanine
T	Thymine
U	Uracil
I	Inosine
X_C	Trans azobenzenes
X_T	Cis azobenzenes
A*	Hydroxyadenine
AL	Locked nucleic acid
TL	"
GL	"
CL	"

- **Comp.sequence**
 - Mandatory if there are mismatches, inosine(s) or hydroxyadenine(s) between the two strands. If not specified, it is computed as the complement of **sequence**. Self-complementarity in **sequence** is detected

even though there may be (are) dangling end(s) and `comp.sequence` is computed.

- `nucleic.acid.conc`
 - In molar concentration (M or mol L⁻¹).
- `Na.conc`, `Mg.conc`, `Tris.conc`, `K.conc`
 - At least one cation (Na, Mg, Tris, K) concentration is mandatory, the other agents(dNTP, DMSO, formamide) are optional.
- `hybridisation.type`
 - The possible options for hybridisation type are as follows (**Table 2**).

Table 2: Hybridisation type options

Option	Sequence	Complementary sequence
<code>dnadna</code>	DNA	DNA
<code>rnarna</code>	RNA	RNA
<code>dnarna</code>	DNA	RNA
<code>rnadna</code>	RNA	DNA
<code>mrnarna</code>	2-o-methyl RNA	RNA
<code>rnamrna</code>	RNA	2-o-methyl RNA

With these arguments, the melting temperature can be computed as follows.

```
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1)
```

```
## [1] 73.35168
```

Only the melting temperature is given as a console output. However, the output can be assigned to an object which contains the details of the environment, options and the thermodynamics results as a list.

```
# Get output as list
out <- melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
               hybridisation.type = "dnadna", Na.conc = 1)
# Environment output
out$Environment
```

```
## $Sequence
## [1] "CAGTGAGACAGCAATGGTCG"
##
## $`Complementary sequence`
## [1] "GTCACCTCTGTCGTTACCAGC"
##
## $`Nucleic acid concentration (M)`
## [1] 2e-06
##
## $`Hybridization type`
```

```
## [1] "dnadna"
##
## $`Na concentration (M)`
## [1] 1
##
## $`Mg concentration (M)`
## [1] 0
##
## $`Tris concentration (M)`
## [1] 0
##
## $`K concentration (M)`
## [1] 0
##
## $`dNTP concentration (M)`
## [1] 0
##
## $`DMSO concentration (%)`
## [1] 0
##
## $`Formamide concentration (M or %)`
## [1] 0
##
## $`Self complementarity`
## [1] FALSE
##
## $`Correction factor`
## [1] 4
##
# Options used
out$options

## $`Approximative formula`
## [1] NA
##
## $`Nearest neighbour model`
## [1] NA
##
## $`GU model`
## [1] NA
##
## $`Single mismatch model`
## [1] NA
##
## $`Tandem mismatch model`
## [1] NA
```

```
##
## `$Single dangling end model`
## [1] NA
##
## `$Double dangling end model`
## [1] NA
##
## `$Long dangling end model`
## [1] NA
##
## `$Internal loop model`
## [1] NA
##
## `$Single bulge loop model`
## [1] NA
##
## `$Long bulge loop model`
## [1] NA
##
## `$CNG repeats model`
## [1] NA
##
## `$Inosine bases model`
## [1] NA
##
## `$Hydroxyadenine bases model`
## [1] NA
##
## `$Azobenzenes model`
## [1] NA
##
## `$Locked nucleic acids model`
## [1] NA
##
## `$Ion correction method`
## [1] NA
##
## `$Na equivalence correction method`
## [1] NA
##
## `$DMSO correction method`
## [1] NA
##
## `$Formamide correction method`
## [1] NA
##
```

```
## $Mode
## [1] NA
# Thermodynamics results
out$results

## $`Enthalpy (cal)`
## [1] -159000
##
## $`Entropy (cal)`
## [1] -430
##
## $`Enthalpy (J)`
## [1] -664620
##
## $`Entropy (J)`
## [1] -1797.4
##
## $`Melting temperature (C)`
## [1] 73.35168
```

The command for the MELTING 5 java version is saved as an attribute in the list `out` and can be retrieved as follows.

```
# Command for MELTING 5
attributes(out)$command

## [1] "-S CAGTGAGACAGCAATGGTCG -H dnadna -P 2e-06 -E Na=1 -T 60"
```

4 Melting temperature computation

Melting temperature is computed by either approximative or nearest neighbour methods according to the length of the oligonucleotide sequences. For longer sequences (longer than the threshold value, the threshold value set by `size.threshold` with the default value 60) approximative method is used, while for others, nearest neighbour method is used.

4.1 Approximative methods

The approximative method for computation can be specified by the argument `method.approx`. The available methods are given in **Table 3**.

Table 3: Details of approximative methods

Formula	Type	Limits/Remarks	Reference
ahs01	DNA	No mismatch	Ahsen et al. (2001)

Formula	Type	Limits/Remarks	Reference
che93	DNA	No mismatch; Na=0, Mg=0.0015, Tris=0.01, K=0.05	Marmur and Doty (1962)
che93corr	DNA	No mismatch; Na=0, Mg=0.0015, Tris=0.01, K=0.05	Marmur and Doty (1962)
schdot	DNA	No mismatch	Wetmur (1991), Marmur and Doty (1962), Chester and Marshak (1993), Schildkraut and Lifson (1965), Wahl et al. (1987), Britten et al. (1974), Hall et al. (1980)
owe69	DNA	No mismatch	Owen et al. (1969), Frank-Kamenetskii (1971), Blake (1996), Blake and Delcourt (1998)
san98	DNA	No mismatch	SantaLucia (1998), Ahsen et al. (2001)
wetdna91*	DNA		Wetmur (1991)
wetrna91*	RNA		Wetmur (1991)
wetdnarna91*	DNA/RNA		Wetmur (1991)

* Default method for computation.

Examples

```

DNA: TCTAATGTGCTGTTAGATGTATCCAGAGATAGCCGAGCATAACTTCAACACACGAGACGTTGATTGGATTTAACCAT
|||||
DNA: AGATTACACGACAATCTACATAGGTCTCTATCGGCTCGTATTTGAAGTTGTGTGCTCTGCAACTAACCTAAATTGGTA

RNA: UUAUAUCUCCGUCAUCUUUAAGCCGUGGAGAGACUGUAGACUUGAACAGGGGUAAGCGGAGGCACGUAGGAUUCACAUC
|||||
RNA: AAUUAGAGGCAGUAGAAAUCGGCACCUCUCUGACAUCUGAACUUGUCCCAUUCGCCUCCGUGCAUCCUAGUGUAG

DNA: TCTAATGTGCTGTTAGATGTATCCAGAGATAGCCGAGCATAACTTCAACACACGAGACGTTGATTGGATTTAACCAT
|||||
RNA: AGAUUACACGACAAUCUACAUAGGUCUCUAUCGGCUCGUAAUUGAAGUUGUGUGCUCUGCAACUAACCUAAAUUGUA

```

```

# Long Nucleotide sequence
DNAseq <- c("TCTAATGTGCTGTTAGATGTATCCAGAGATAGCCGAGCATAACTTCAACACACGAGACGTTGATTGGATTTAACCATAG")
RNAseq <- c("UUAUUCUCCGUCAUCUUUAAGCCGUGGAGAGACUGUAGACUUGAACAGGGGUAAGCGGAGGCACGUAGGAUUCACAUCAU")

# Approximative method - default (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1)

## [1] 87.82455

# Approximative method - wetdna91 (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1,
         method.approx = "wetdna91")

## [1] 87.82455

# Approximative method - ahs01 (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1,
         method.approx = "ahs01")

## [1] 87.325

# Approximative method - che93 (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1,
         method.approx = "che93")

## [1] 77.575

# Approximative method - che93corr (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1,
         method.approx = "che93corr")

## [1] 79.0125

# Approximative method - schdot (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1,
         method.approx = "schdot")

## [1] 89.4625

# Approximative method - owe69 (DNA/DNA)
melting(sequence = DNAseq, nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1,
         method.approx = "owe69")

```

```
## [1] 100.96
```

```
# Approximative method - san98 (DNA/DNA)
melting(sequence = DNaseq, nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1,
        method.approx = "san98")
```

```
## [1] 86.9
```

```
# Approximative method - default (RNA/RNA)
melting(sequence = RNaseq, nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1)
```

```
## [1] 101.1745
```

```
# Approximative method - wetrna91 (RNA/RNA)
melting(sequence = RNaseq, nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1,
        method.approx = "wetrna91")
```

```
## [1] 101.1745
```

```
# Approximative method - wetdnarna91 (DNA/RNA)
melting(sequence = DNaseq, nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnarna", Na.conc = 1)
```

```
## [1] 88.92455
```

```
# Approximative method - wetdnarna91 (DNA/RNA)
melting(sequence = DNaseq, nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnarna", Na.conc = 1,
        method.approx = "wetdnarna91")
```

```
## [1] 88.92455
```

4.2 Nearest neighbour methods

4.2.1 Perfectly matching sequences

The nearest neighbour model for computation in case of perfectly matching sequences can be specified by the argument `method.nn`. The available methods are given in **Table 4**.

Table 4: Details of nearest neighbour methods for perfectly matching sequences

Model	Type	Limits/Remarks	Reference
all197*	DNA		Allawi and SantaLucia (1997)
bre86	DNA		Breslauer et al. (1986)
san04	DNA		SantaLucia and Hicks (2004)
san96	DNA		SantaLucia et al. (1996)

Model	Type	Limits/Remarks	Reference
sug96	DNA		Sugimoto et al. (1996)
tan04	DNA		Tanaka et al. (2004)
fre86	RNA		Freier et al. (1986)
xia98*	RNA		Xia et al. (1998)
sug95*	DNA/ RNA		SantaLucia et al. (1996)
tur06*	2'-O- MeRNA/ RNA	A sodium correction (san04) is automatically applied to convert the entropy (Na = 0.1M) into the entropy (Na = 1M)	Kierzek et al. (2006)

* Default method for computation.

Examples

```
DNA:CAGTGAGACAGCAATGGTCG
|||||
DNA:GTCACTCTGTCGTTACCAGC
```

```
RNA:CAGUGAGACAGCAAUGGUCG
|||||
RNA:GUCACUCUGUCGUUACCAGC
```

```
DNA:CAGTGAGACAGCAATGGTCG
|||||
RNA:GUCACUCUGUCGUUACCAGC
```

```
# Nearest neighbour method - default (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1)
```

```
## [1] 73.35168
```

```
# Nearest neighbour method - all97 (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
         hybridisation.type = "dnadna", Na.conc = 1, method.nn = "all97")
```

```
## [1] 73.35168
```

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```
# Nearest neighbour method - bre86 (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "bre86")

## [1] 83.2203

# Nearest neighbour method - san04 (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "san04")

## [1] 73.30191

# Nearest neighbour method - san96 (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "san96")

## [1] 75.7102

# Nearest neighbour method - sug96 (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "sug96")

## [1] 78.17556

# Nearest neighbour method - tan04 (DNA/DNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "tan04")

## [1] 71.31413

# Nearest neighbour method - default (RNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGUGAGACAGCAAUGGUCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1)

## [1] 86.77685

# Nearest neighbour method - xia98 (RNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGUGAGACAGCAAUGGUCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1, method.nn = "xia98")

## [1] 86.77685

# Nearest neighbour method - fre86 (RNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGUGAGACAGCAAUGGUCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1, method.nn = "fre86")

## [1] 83.81257

# Nearest neighbour method - default (mRNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGUGAGACAGCAAUGGUCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "mrnarna", Na.conc = 1)
```

```
## [1] 99.01986
# Nearest neighbour method - tur06 (mRNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGUGAGACAGCAAUGGUCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "mrnarna", Na.conc = 1, method.nn = "tur06")
```

```
## [1] 99.01986
# Nearest neighbour method - default (DNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnarna", Na.conc = 1)
```

```
## [1] 66.77049
# Nearest neighbour method - sug95 (DNA/RNA: No Self-Complimentarity)
melting(sequence = "CAGTGAGACAGCAATGGTCG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnarna", Na.conc = 1, method.nn = "sug95")
```

```
## [1] 66.77049
```

Self complementarity for perfect matching sequences or sequences with dangling ends is detected automatically. However it can be enforced by the argument `force.self = TRUE`.

Examples

```
DNA:CATATGGCCATATG
      |||||
DNA:GTATACCGGTATAC
```

```
RNA:AUGUACAU
      |||||
RNA:UACAUGUA
```

```
# Nearest neighbour method - default (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1)
```

```
## [1] 56.00644
# Nearest neighbour method - all97 (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "all97")
```

```
## [1] 56.00644
# Nearest neighbour method - bre86 (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "bre86")
```

```
## [1] 63.44605
```

```
# Nearest neighbour method - san04 (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "san04")

## [1] 57.80792

# Nearest neighbour method - san96 (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "san96")

## [1] 55.0921

# Nearest neighbour method - sug96 (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "sug96")

## [1] 59.06213

# Nearest neighbour method - tan04 (DNA/DNA: Self-Complimentarity)
melting(sequence = "CATATGGCCATATG", nucleic.acid.conc = 2e-06,
        hybridisation.type = "dnadna", Na.conc = 1, method.nn = "tan04")

## [1] 55.65824

# Nearest neighbour method - default (RNA/RNA: Self-Complimentarity)
melting(sequence = "AUGUACAU", nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1)

## [1] 30.27015

# Nearest neighbour method - xia98 (RNA/RNA: Self-Complimentarity)
melting(sequence = "AUGUACAU", nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1, method.nn = "xia98")

## [1] 30.27015

# Nearest neighbour method - fre86 (RNA/RNA: Self-Complimentarity)
melting(sequence = "AUGUACAU", nucleic.acid.conc = 2e-06,
        hybridisation.type = "rnarna", Na.conc = 1, method.nn = "fre86")

## [1] 31.48175
```

4.2.2 GU wobble base pairs effect

The nearest neighbour model for computation in case of sequences with GU wobble base pairs can be specified by the argument `method.GU`. The available methods are given in **Table 5**.

Table 5: Details of methods for sequences with GU wobble base pairs

Model	Type	Limits/Remarks	Reference
tur99	RNA		Mathews et al. (1999)
ser12*	RNA		Chen et al. (2012)

* Default method for computation.

Examples

```
RNA:CCAGCGUCCU
      |||||
RNA:GGTCGCAGGA
```

```
# GU wobble base pairs effect - default (RNA/RNA)
melting(sequence = "CCAGCGUCCU", nucleic.acid.conc = 0.0001,
        hybridisation.type = "rnarna", Na.conc = 1)

## [1] 79.46955

# GU wobble base pairs effect - ser12 (RNA/RNA)
melting(sequence = "CCAGCGUCCU", nucleic.acid.conc = 0.0001,
        hybridisation.type = "rnarna", Na.conc = 1, method.GU = "ser12")

## [1] 79.46955

# GU wobble base pairs effect - tur99 (RNA/RNA)
melting(sequence = "CCAGCGUCCU", nucleic.acid.conc = 0.0001,
        hybridisation.type = "rnarna", Na.conc = 1, method.GU = "tur99")

## [1] 79.46955
```

4.2.3 Single mismatch effect

The nearest neighbour model for computation in case of sequences with a single mismatch can be specified by the argument `method.singleMM`. The available methods are given in **Table 6**.

Table 6: Details of methods for sequences with single mismatch

Model	Type	Limits/Remarks	Reference
allsanpey*	DNA		Allawi and SantaLucia (1997), Allawi and SantaLucia (1998a), Allawi and SantaLucia (1998b), Allawi and SantaLucia (1998c), Peyret et al. (1999)
wat11*	DNA/RNA		Watkins et al. (2011)

Model	Type	Limits/Remarks	Reference
tur06	RNA		Lu et al. (2006)
zno07*	RNA		Davis and Znosko (2007)
zno08	RNA	At least one adjacent GU base pair.	Davis and Znosko (2008)

* Default method for computation.

Examples

```
DNA:CAACTTGATATTAATA
      ||||| |||||
DNA:GTTGAACTCTAATTAT
```

```
RNA:GACAGGCUG
      ||| |||
RNA:CUGUGCGAC
```

```
DNA:CCATAACTACC
      ||| |||||
RNA:GGUAAUGAUGG
```

```
# Single mismatch effect - default (DNA/DNA)
melting(sequence = "CAACTTGATATTAATA", comp.sequence = "GTTGAACTCTAATTAT",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna", Na.conc = 1)
```

```
## [1] 51.97499
```

```
# Single mismatch effect - allsanpey (DNA/DNA)
melting(sequence = "CAACTTGATATTAATA", comp.sequence = "GTTGAACTCTAATTAT",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
         Na.conc = 1, method.singleMM = "allsanpey")
```

```
## [1] 51.97499
```

```
# Single mismatch effect - default (RNA/RNA)
melting(sequence = "GACAGGCUG", comp.sequence = "CUGUGCGAC",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna", Na.conc = 1)
```

```
## [1] 54.40363
```

```
# Single mismatch effect - zno07 (RNA/RNA)
melting(sequence = "GACAGGCUG", comp.sequence = "CUGUGCGAC",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
         Na.conc = 1, method.singleMM = "zno07")
```

```
## [1] 54.40363
# Single mismatch effect - zno08 (RNA/RNA)
melting(sequence = "CAGUACGUC", comp.sequence = "GUCGGGCAG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.singleMM = "zno08")

## [1] 38.26298
# Single mismatch effect - tur06 (RNA/RNA)
melting(sequence = "GACAGGCUG", comp.sequence = "CUGUGCGAC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.singleMM = "tur06")

## [1] 58.27825
# Single mismatch effect - default (DNA/RNA)
melting(sequence = "CCATAACTACC", comp.sequence = "GGUAAUGAUGG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnarna", Na.conc = 1)

## [1] 40.32976
# Single mismatch effect - wat11 (DNA/RNA)
melting(sequence = "CCATAACTACC", comp.sequence = "GGUAAUGAUGG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnarna",
        Na.conc = 1, method.singleMM = "wat11")

## [1] 40.32976
```

4.2.4 Tandem mismatches effect

The nearest neighbour model for computation in case of sequences with tandem mismatches can be specified by the argument `method.tandemMM`. The available methods are given in **Table 7**.

Table 7: Details of methods for sequences with tandem mismatches

Model	Type	Limits/Remarks	Reference
allsanpey*	DNA	Only GT mismatches and TA/TG mismatches.	Allawi and SantaLucia (1997), Allawi and SantaLucia (1998a), Allawi and SantaLucia (1998b), Allawi and SantaLucia (1998c), Peyret et al. (1999)
tur99*	RNA	No adjacent GU or UG base pairs.	Mathews et al. (1999), Lu et al. (2006)

* Default method for computation.

Examples

```

DNA:GACGTTGGAC
    |||  |||
DNA:CTGCGGCCTG

RNA:GAGCGGAG
   ||  ||
RNA:CUCCACUC

# Tandem mismatches effect - default (DNA/DNA)
melting(sequence = "GACGTTGGAC", comp.sequence = "CTGCGGCCTG",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna", Na.conc = 1)

## [1] 50.20175

# Tandem mismatches effect - allsanpey (DNA/DNA)
melting(sequence = "GACGTTGGAC", comp.sequence = "CTGCGGCCTG",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
         Na.conc = 1, method.tandemMM = "allsanpey")

## [1] 50.20175

# Tandem mismatches effect - default (RNA/RNA)
melting(sequence = "GAGCGGAG", comp.sequence = "CUCCACUC",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna", Na.conc = 1)

## [1] 21.07224

# Tandem mismatches effect - tur06 (RNA/RNA)
melting(sequence = "GAGCGGAG", comp.sequence = "CUCCACUC",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
         Na.conc = 1, method.tandemMM = "tur99")

## [1] 21.07224

```

4.2.5 Single dangling end effect

The nearest neighbour model for computation in case of sequences with a single dangling end can be specified by the argument `method.single.dangle`. The available methods are given in **Table 8**.

Table 8: Details of methods for sequences with single dangling end

Model	Type	Limits/Remarks	Reference
bom00*	DNA		Bommarito et al. (2000)

Model	Type	Limits/Remarks	Reference
sugdna02	DNA	Only terminal poly A self complementary sequences.	Ohmichi et al. (2002)
sugrna02	RNA	Only terminal poly A self complementary sequences.	Ohmichi et al. (2002)
ser08*	RNA	Only 3' UA, GU and UG terminal base pairs only 5' UG and GU terminal base pairs.	O'Toole et al. (2006), Miller et al. (2008)

* Default method for computation.

Examples

DNA: -GTAGCTACA

|||||||

DNA: ACATCGATG-

RNA: -GGCGCUG

|||||||

RNA: CCGCGAC

DNA: -GGCGCUG

|||||||

RNA: CCGCGAC

```
# Single dangling end effect - default (DNA/DNA)
melting(sequence = "-GTAGCTACA",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
         Na.conc = 1)
```

```
## [1] 52.58935
```

```
# Single dangling end effect - bom00 (DNA/DNA)
melting(sequence = "-GTAGCTACA",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
         Na.conc = 1, method.single.dangle = "bom00")
```

```
## [1] 52.58935
```

```
# Single dangling end effect - sugdna02 (DNA/DNA)
melting(sequence = "-GTAGCTACA",
```

```

nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
Na.conc = 1, method.single.dangle = "sugdna02")

## [1] 50.78548
# Single dangling end effect - default (RNA/RNA)
melting(sequence = "-GGCGCUG",
nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
Na.conc = 1)

## [1] 65.7647
# Single dangling end effect - ser08 (RNA/RNA)
melting(sequence = "-GGCGCUG",
nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
Na.conc = 1, method.single.dangle = "ser08")

## [1] 65.7647
# Single dangling end effect - sugrna02 (RNA/RNA)
melting(sequence = "-GGCGCUG",
nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
Na.conc = 1, method.single.dangle = "sugrna02")

## [1] 65.7647

```

4.2.6 Double dangling end effect

The nearest neighbour model for computation in case of sequences with a double or secondary dangling ends can be specified by the argument `method.double.dangle`. The available methods are given in **Table 9**.

Table 9: Details of methods for sequences with double dangling ends

Model	Type	Limits/Remarks	Reference
sugdna02*	DNA	Only terminal poly A self complementary sequences.	Ohmichi et al. (2002)
sugrna02	RNA	Only terminal poly A self complementary sequences.	Ohmichi et al. (2002)
ser05	RNA	Depends on the available thermodynamic parameters for single dangling end.	O'Toole et al. (2005)
ser06*	RNA		O'Toole et al. (2006)

* Default method for computation.

Examples

DNA:--ATGCATAA

|||||

DNA:AATACGTA--

RNA:--AUGCAUAA

|||||

RNA:AAUACGUA--

```
# Double dangling end effect - default (DNA/DNA)
melting(sequence = "--ATGCATAA",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
         Na.conc = 1)
```

```
## [1] 44.88615
```

```
# Double dangling end effect - sugdna02 (DNA/DNA)
melting(sequence = "--ATGCATAA",
         nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
         Na.conc = 1, method.double.dangle = "sugdna02")
```

```
## [1] 44.88615
```

```
# Double dangling end effect - default (RNA/RNA)
melting(sequence = "--AUGCAUAA",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
         Na.conc = 1)
```

```
## [1] 42.79724
```

```
# Double dangling end effect - ser06 (RNA/RNA)
melting(sequence = "--AUGCAUAA",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
         Na.conc = 1, method.double.dangle = "ser06")
```

```
## [1] 42.79724
```

```
# Double dangling end effect - sugrna02 (RNA/RNA)
melting(sequence = "--AUGCAUAA",
         nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
         Na.conc = 1, method.double.dangle = "sugrna02")
```

```
## [1] 41.82788
```

```
# Double dangling end effect - ser05 (RNA/RNA)
melting(sequence = "--AUGCAUAA",
```

```
nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
Na.conc = 1, method.double.dangle = "ser05")
```

```
## [1] 42.78815
```

4.2.7 Long dangling end effect

The nearest neighbour model for computation in case of sequences with a double or secondary dangling ends can be specified by the argument `method.long.dangle`. The available methods are given in **Table 10**.

Table 10: Details of methods for sequences with long dangling ends

Model	Type	Limits/Remarks	Reference
sugdna02*	DNA	Only terminal poly A self complementary sequences.	Ohmichi et al. (2002)
sugrna02*	RNA	Only terminal poly A self complementary sequences.	Ohmichi et al. (2002)

* Default method for computation.

Examples

DNA:----GCATATGCAAAA

|||||||

DNA:AAAACGTATACG----

RNA:AAAAGCAUAUGC----

|||||||

RNA:----CGUAUACGAAAA

```
# Long dangling end effect - default (DNA/DNA)
melting(sequence = "----GCATATGCAAAA",
        nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
        Na.conc = 1)
```

```
## [1] 55.69854
```

```
# Long dangling end effect - sugdna02 (DNA/DNA)
melting(sequence = "----GCATATGCAAAA",
        nucleic.acid.conc = 0.0004, hybridisation.type = "dnadna",
        Na.conc = 1, method.long.dangle = "sugdna02")
```

```
## [1] 55.69854
```

```
# Long dangling end effect - default (RNA/RNA)
melting(sequence = "AAAAGCAUAUGC----",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1)
```

```
## [1] 57.21314
```

```
# Long dangling end effect - sugrna02 (RNA/RNA)
melting(sequence = "AAAAGCAUAUGC----",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.long.dangle = "sugrna02")
```

```
## [1] 57.21314
```

4.2.8 Internal loop effect

The nearest neighbour model for computation in case of sequences with an internal loop (more than two adjacent mismatches) can be specified by the argument `method.internal.loop`. The available methods are given in **Table 11**.

Table 11: Details of methods for sequences with internal loops

Model	Type	Limits/Remarks	Reference
san04*	DNA	Missing asymmetry penalty. Not tested with experimental results.	SantaLucia and Hicks (2004)
tur06	RNA	Not tested with experimental results.	Lu et al. (2006)
zno07*	RNA	Only for 1x2 loop.	Badhwar et al. (2007)

* Default method for computation.

Examples

```
DNA:GCGATTGGCACTTTGGTGAAC
      |||||      |||||
DNA:CGCTACATATGAAACCACTTG

RNA:GACAC-GCUG
      ||||  |||
RNA:CUGUAUCGAC
```

```
# Internal loop effect - default (DNA/DNA)
melting(sequence = "GCGATTGGCACTTTGGTGAAC", comp.sequence = "CGCTACATATGAAACCACTTG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)
```

```
## [1] 84.09052
```

```
# Internal loop effect - san04 (DNA/DNA)
melting(sequence = "GCGATTGGCACTTTGGTGAAC", comp.sequence = "CGCTACATATGAAACCACTTG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.internal.loop = "san04")
```

```
## [1] 84.09052
```

```
# Internal loop effect - default (RNA/RNA)
melting(sequence = "GACAC-GCUG", comp.sequence = "CUGUAUCGAC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1)
```

```
## [1] 45.98713
```

```
# Internal loop effect - zno07 (RNA/RNA)
melting(sequence = "GACAC-GCUG", comp.sequence = "CUGUAUCGAC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.internal.loop = "zno07")
```

```
## [1] 40.49012
```

```
# Internal loop effect - tur06 (RNA/RNA)
melting(sequence = "GACAC-GCUG", comp.sequence = "CUGUAUCGAC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.internal.loop = "tur06")
```

```
## [1] 45.98713
```

4.2.9 Single bulge loop effect

The nearest neighbour model for computation in case of sequences with a single bulge loop can be specified by the argument `method.single.bulge.loop`. The available methods are given in **Table 12**.

Table 12: Details of methods for sequences with single bulge loop

Model	Type	Limits/Remarks	Reference
tan04*	DNA		Tan and Chen (2007)
san04	DNA	Missing closing AT penalty.	SantaLucia and Hicks (2004)

Model	Type	Limits/Remarks	Reference
ser07	RNA	Less reliable results. Some missing parameters.	Blose et al. (2007)
tur06*	RNA		Lu et al. (2006)

* Default method for computation.

Examples

DNA:TCGATTAGCGACACAGG

||||| |||||

DNA:AGCTAATC-CTGTGTCC

RNA:GACUCUGUC

|||| |||

RNA:CUGA-ACAG

Single bulge loop effect - default (DNA/DNA)

```
melting(sequence = "TCGATTAGCGACACAGG", comp.sequence = "AGCTAATC-CTGTGTCC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)
```

```
## [1] 71.12754
```

Single bulge loop effect - tan04 (DNA/DNA)

```
melting(sequence = "TCGATTAGCGACACAGG", comp.sequence = "AGCTAATC-CTGTGTCC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.single.bulge.loop = "tan04")
```

```
## [1] 71.12754
```

Single bulge loop effect - san04 (DNA/DNA)

```
melting(sequence = "TCGATTAGCGACACAGG", comp.sequence = "AGCTAATC-CTGTGTCC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.single.bulge.loop = "san04")
```

```
## [1] 62.0496
```

Single bulge loop effect - default (RNA/RNA)

```
melting(sequence = "GACUCUGUC", comp.sequence = "CUGA-ACAG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1)
```

```
## [1] 39.47787
```

```
# Single bulge loop effect - tur06 (RNA/RNA)
melting(sequence = "GACUCUGUC", comp.sequence = "CUGA-ACAG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.single.bulge.loop = "tur06")
```

```
## [1] 39.47787
```

```
# Single bulge loop effect - ser07 (RNA/RNA)
melting(sequence = "GACUCUGUC", comp.sequence = "CUGA-ACAG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.single.bulge.loop = "ser07")
```

```
## [1] 31.42849
```

4.2.10 Long bulge loop effect

The nearest neighbour model for computation in case of sequences with long bulge loop can be specified by the argument `method.long.bulge.loop`. The available methods are given in **Table 13**.

Table 13: Details of methods for sequences with long bulge loop

Model	Type	Limits/Remarks	Reference
san04*	DNA	Missing closing AT penalty.	SantaLucia and Hicks (2004)
tur06*	RNA	Not tested with experimental results.	Mathews et al. (1999), Lu et al. (2006)

* Default method for computation.

Examples

```
DNA: ATATGACGCCACAGCG
      ||||  |||||
DNA: TATAC---GGTGTGCGC

RNA: AUAUGACGCCACAGCG
      ||||  |||||
RNA: UAUAC---GGUGUCGC
```

```
# Long bulge loop effect - default (DNA/DNA)
melting(sequence = "ATATGACGCCACAGCG", comp.sequence = "TATAC---GGTGTGCGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)
```

```
## [1] 51.7104
```

```
# Long bulge loop effect - san04 (DNA/DNA)
```

```
melting(sequence = "ATATGACGCCACAGCG", comp.sequence = "TATAC---GGTGTCGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.long.bulge.loop = "san04")
```

```
## [1] 51.7104
```

```
# Long bulge loop effect - default (RNA/RNA)
```

```
melting(sequence = "AUAUGACGCCACAGCG", comp.sequence = "UAUAC---GGUGUCGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1)
```

```
## [1] 66.0497
```

```
# Long bulge loop effect - tur06 (RNA/RNA)
```

```
melting(sequence = "AUAUGACGCCACAGCG", comp.sequence = "UAUAC---GGUGUCGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.long.bulge.loop = "tur06")
```

```
## [1] 66.0497
```

4.2.11 CNG repeats effect

The nearest neighbour model for computation in case of sequences with CNG repeats can be specified by the argument `method.CNG`. The available methods are given in **Table 14**.

Table 14: Details of methods for sequences with CNG repeats

Model	Type	Limits/Remarks	Reference
bro05*	RNA	Self complementary sequences. 2 to 7 CNG repeats.	Broda et al. (2005)

* Default method for computation.

Examples

```
RNA:GCGGCGGCGGC
```

```
|||||||
```

```
RNA:CGCCGCCGCCG
```

```
# CNG repeats effect - default (RNA/RNA)
```

```
melting(sequence = "GCGGCGGCGGC",
```

```
nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
Na.conc = 1)

## [1] 94.25719

# CNG repeats effect - bro05 (RNA/RNA)
melting(sequence = "GCGGCGGCGGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.CNG = "bro05")

## [1] 94.25719
```

4.2.12 Inosine bases effect

The nearest neighbour model for computation in case of sequences with inosine bases (I) can be specified by the argument `method.inosine`. The available methods are given in **Table 15**.

Table 15: Details of methods for sequences with inosine bases

Model	Type	Limits/Remarks	Reference
san05*	DNA	Missing parameters for tandem base pairs containing inosine bases.	Watkins and SantaLucia (2005)
zno07*	RNA	Only IU base pairs.	Wright et al. (2007)

* Default method for computation.

Examples

```
DNA:CCGICTGTIGCG
    ||| ||| |||
DNA:GGCCGACACCGC

RNA:GCAICGC
   ||| |||
RNA:CGUUGCG

# Inosine bases effect - default (DNA/DNA)
melting(sequence = "CCGICTGTIGCG", comp.sequence = "GGCCGACACCGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)

## [1] 65.36853
```

```
# Inosine bases effect - san05 (DNA/DNA)
melting(sequence = "CCGICTGTIGCG", comp.sequence = "GGCCGACACCGC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.inosine = "san05")
```

```
## [1] 65.36853
```

```
# Inosine bases effect - default (RNA/RNA)
melting(sequence = "GCAICGC", comp.sequence = "CGUUGCG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1)
```

```
## [1] 46.75042
```

```
# Inosine bases effect - zno07 (RNA/RNA)
melting(sequence = "GCAICGC", comp.sequence = "CGUUGCG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "rnarna",
        Na.conc = 1, method.inosine = "zno07")
```

```
## [1] 46.75042
```

4.2.13 Hydroxyadenine bases effect

The nearest neighbour model for computation in case of sequences with hydroxyadenine bases can be specified by the argument `method.hydroxyadenine`. The available methods are given in **Table 16**.

Table 16: Details of methods for sequences with hydroxyadenine bases

Model	Type	Limits/Remarks	Reference
sug01*	DNA	Only 5' GA*C 3' and 5' TA*A 3' contexts.	Kawakami et al. (2001)

* Default method for computation.

Examples

```
      *
DNA:AGAAATGACACGGTG
    |||||
DNA:TCTTTACCGTGCCAC
```

```
# Hydroxyadenine bases effect - default (DNA/DNA)
melting(sequence = "AGAAATGA*ACCGTG", comp.sequence = "TCTTTACCGTGCCAC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)
```

```
## [1] 68.46041
# Hydroxyadenine bases effect - sug01 (DNA/DNA)
melting(sequence = "AGAAATGA*CACGGTG", comp.sequence = "TCTTTACCGTGCCAC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.hydroxyadenine = "sug01")
```

```
## [1] 68.46041
```

4.2.14 Azobenzenes effect

The nearest neighbour model for computation in case of sequences with azobenzenes (X_T for trans azobenzenes and X_C for cis azobenzenes) can be specified by the argument `method.azobenzenes`. The available methods are given in **Table 17**.

Table 17: Details of methods for sequences with azobenzenes

Model	Type	Limits/Remarks	Reference
asa05*	DNA	Less reliable results when the number of cis azobenzene increases.	Asanuma et al. (2005)

* Default method for computation.

Examples

```
      C   C   C   C   C
DNA:CTXTTAAXGAAGXGAGAXTATAXCC
   || |||| |||| |||| |||| ||
DNA:GA AATT CTTC CTCT ATAT GG

# Azobenzenes effect - default (DNA/DNA)
melting(sequence = "CTX_CTTAAX_CGAAGX_CGAGAX_CTATAX_CCC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)
```

```
## [1] 47.85385

# Azobenzenes effect - asa05 (DNA/DNA)
melting(sequence = "CTX_CTTAAX_CGAAGX_CGAGAX_CTATAX_CCC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.azobenzenes = "asa05")
```

```
## [1] 47.85385
```

4.2.15 Single Locked nucleic acid effect

The nearest neighbour model for computation in case of sequences with single locked nucleic acids can be specified by the argument `method.locked`. The available methods are given in **Table 18**.

Table 18: Details of methods for sequences with single locked nucleic acids

Model	Type	Limits/Remarks	Reference
mct04	DNA		McTigue et al. (2004)
owc11*	DNA		Owczarzy et al. (2011)

* Default method for computation.

Examples

```

      L
DNA:CCATTGCTACC
      |||||
DNA:GGTAACGATGG

# Single locked nucleic acids effect - default (DNA/DNA)
melting(sequence = "CCATTGCTACC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)

## [1] 63.48299

# Single locked nucleic acids effect - mct04 (DNA/DNA)
melting(sequence = "CCATTGCTACC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.locked = "mct04")

## [1] 63.61426

# Single locked nucleic acids effect - owc11 (DNA/DNA)
melting(sequence = "CCATTGCTACC",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.locked = "owc11")

## [1] 63.48299

```

4.3 Consecutive Locked nucleic acids effect

The nearest neighbour model for computation in case of sequences with consecutive locked nucleic acids can be specified by the argument `method.consecutive.locked`. The available methods are given in **Table 19**.

Table 19: Details of methods for sequences with single locked nucleic acids

Model	Type	Limits/Remarks	Reference
owc11*	DNA		Owczarzy et al. (2011)

* Default method for computation.

Examples

```

LL
DNA:GACC
||||
DNA:CTGG

# Consecutive locked nucleic acids effect - default (DNA/DNA)
melting(sequence = "GALCLC",
         nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
         Na.conc = 1)

## [1] 12.94323

# Consecutive locked nucleic acids effect - owc11 (DNA/DNA)
melting(sequence = "GALCLC",
         nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
         Na.conc = 1, method.consecutive.locked = "owc11")

## [1] 12.94323

```

4.4 Consecutive Locked nucleic acids with a single mismatch effect

The nearest neighbour model for computation in case of sequences with consecutive locked nucleic acids with single mismatch can be specified by the argument `method.consecutive.locked.singleMM`. The available methods are given in **Table 20**.

Table 20: Details of methods for sequences with single locked nucleic acids

Model	Type	Limits/Remarks	Reference
owc11*	DNA		Owczarzy et al. (2011)

* Default method for computation.

Examples

```

      LLL
DNA:GACGC
  || ||
DNA:CTTCG

# Consecutive locked nucleic acids effect - default (DNA/DNA)
melting(sequence = "GALCLGLC", comp.sequence = "CTTCG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1)

## [1] 0.2520424

# Consecutive locked nucleic acids effect - owc11 (DNA/DNA)
melting(sequence = "GALCLGLC", comp.sequence = "CTTCG",
        nucleic.acid.conc = 0.0001, hybridisation.type = "dnadna",
        Na.conc = 1, method.consecutive.locked.singleMM = "owc11")

## [1] 0.2520424

```

5 Corrections

Once the melting temperature is computed, a correction is applied to it according to the concentration of nucleic acids, cations and/or denaturing agents.

5.1 Nucleic acid concentration

For self complementary sequences (auto detected or specified by `force.self`) it is 1. Otherwise it is 4 if the both strands are present in equivalent amount and 1 if one strand is in excess.

5.2 Ion corrections

Melting temperature is computed initially for $[\text{Na}^+] = 1 \text{ M}$, after which a correction for the presence of cations ($[\text{Na}^+]$, $[\text{K}^+]$, $[\text{Tris}^+]$ and $[\text{Mg}^+]$) is applied either directly on the computed melting temperature or on the computed entropy.

The correction methods for cation concentration can be specified by the argument `correction.ion`.

5.2.1 Sodium corrections

The available correction methods for sodium concentration are given in **Table 21**.

Table 21: Details of the corrections for sodium concentration

Correction	Type	Limits/Remarks	Reference
<code>ahs01</code>	DNA	$Na > 0$.	Ahsen et al. (2001)
<code>kam71</code>	DNA	$Na > 0$; $Na \geq 0.069$; $Na \leq 1.02$.	Frank-Kamenetskii (1971)
<code>marschdot</code>	DNA	$Na \geq 0.069$; $Na \leq 1.02$.	Marmur and Doty (1962), Blake and Delcourt (1998)
<code>owc1904</code>	DNA	$Na > 0$. (equation 19)	Owczarzy et al. (2004)
<code>owc2004</code>	DNA	$Na > 0$. (equation 20)	Owczarzy et al. (2004)
<code>owc2104</code>	DNA	$Na > 0$. (equation 21)	Owczarzy et al. (2004)
<code>owc2204*</code>	DNA	$Na > 0$. (equation 22)	Owczarzy et al. (2004)
<code>san96</code>	DNA	$Na \geq 0.1$.	SantaLucia et al. (1996)
<code>san04</code>	DNA	$Na \geq 0.05$; $Na \leq 1.1$; Oligonucleotides inferior to 16 bases.	SantaLucia and Hicks (2004), SantaLucia (1998)
<code>schlif</code>	DNA	$Na \geq 0.07$; $Na \leq 0.12$.	Schildkraut and Lifson (1965)
<code>tanna06</code>	DNA	$Na \geq 0.001$; $Na \leq 1$.	Tan and Chen (2006)
<code>tanna07*</code>	RNA or 2'-O-MeRNA/RNA	$Na \geq 0.003$; $Na \leq 1$.	Tan and Chen (2007)
<code>wet91</code>	RNA, DNA and RNA/DNA	$Na > 0$.	Wetmur (1991)

* Default method for computation.

```
# Na correction - default (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
         Na.conc = 0.069)
```

```
## [1] 56.70492
```

```
# Na correction - owc2204 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
```

```

Na.conc = 0.069, correction.ion = "owc2204")

## [1] 56.70492
# Na correction - ahs01 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "ahs01")

## [1] 54.1569
# Na correction - kam71 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "kam71")

## [1] 51.72963
# Na correction - marschdot (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "marschdot")

## [1] 49.18075
# Na correction - owc1904 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "owc1904")

## [1] 56.18571
# Na correction - owc2004 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "owc2004")

## [1] 56.67553
# Na correction - owc2104 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "owc2104")

## [1] 56.63967
# Na correction - san96 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "san96")

```

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```
## [1] 53.01651
```

```
# Na correction - san04 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "san04")
```

```
## [1] 54.15157
```

```
# Na correction - schlif (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "schlif")
```

```
## [1] 48.25579
```

```
# Na correction - tanna06 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "tanna06")
```

```
## [1] 55.26711
```

```
# Na correction - wet91 (DNA/DNA)
melting(sequence = "CCAGCCAGTCTCTCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, correction.ion = "wet91")
```

```
## [1] 51.74573
```

```
# Na correction - default (RNA/RNA)
melting(sequence = "CCAGCCAGUCUCUCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Na.conc = 0.069)
```

```
## [1] 75.1552
```

```
# Na correction - tanna07 (RNA/RNA)
melting(sequence = "CCAGCCAGUCUCUCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Na.conc = 0.069, correction.ion = "tanna07")
```

```
## [1] 75.1552
```

```
# Na correction - wet91 (RNA/RNA)
melting(sequence = "CCAGCCAGUCUCUCC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Na.conc = 0.069, correction.ion = "wet91")
```

```
## [1] 69.55572
```

```

# Na correction - default (mRNA/RNA)
melting(sequence = "UACGCGUCAAUAACGCUA",
         nucleic.acid.conc = 0.000002, hybridisation.type = "mrnarna",
         Na.conc = 0.069)

## [1] 81.57763

# Na correction - tanna07 (mRNA/RNA)
melting(sequence = "UACGCGUCAAUAACGCUA",
         nucleic.acid.conc = 0.000002, hybridisation.type = "mrnarna",
         Na.conc = 0.069, correction.ion = "tanna07")

## [1] 81.57763

# Na correction - default (DNA/RNA)
melting(sequence = "CCAGCCAGTCTCTCC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnarna",
         Na.conc = 0.069)

## [1] 62.08869

# Na correction - wet91 (DNA/RNA)
melting(sequence = "CCAGCCAGTCTCTCC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnarna",
         Na.conc = 0.069, correction.ion = "wet91")

## [1] 62.08869

```

5.2.2 Magnesium corrections

The available correction methods for magnesium concentration are given in **Table 22**.

Table 22: Details of the corrections for magnesium concentration

Correction	Type	Limits/Remarks	Reference
owcmg08*	DNA	Mg $\geq$ 0.0005; Mg $\leq$ 0.6.	Owczarzy et al. (2008)
tanmg06	DNA	Mg $\geq$ 0.0001; Mg $\leq$ 1; Oligomer length superior to 6 base pairs.	Tan and Chen (2006)
tanmg07*	RNA or 2'-O- MeRNA/RNA	Mg $\geq$ 0.1; Mg $\leq$ 0.3.	Tan and Chen (2007)

* Default method for computation.

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```
# Mg correction - default (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Mg.conc = 0.0015)
```

```
## [1] 65.52043
```

```
# Mg correction - owcmg08 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Mg.conc = 0.0015, correction.ion = "owcmg08")
```

```
## [1] 65.52043
```

```
# Mg correction - tanmg06 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Mg.conc = 0.0015, correction.ion = "tanmg06")
```

```
## [1] 64.88082
```

```
# Mg correction - default (RNA/RNA)
melting(sequence = "CAGCCUCGUCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Mg.conc = 0.0015)
```

```
## [1] 82.0796
```

```
# Mg correction - tanmg07 (RNA/RNA)
melting(sequence = "CAGCCUCGUCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Mg.conc = 0.0015, correction.ion = "tanmg07")
```

```
## [1] 82.0796
```

```
# Mg correction - default (mRNA/RNA)
melting(sequence = "UACGCGUCAUAACGCUA",
        nucleic.acid.conc = 0.000002, hybridisation.type = "mrnarna",
        Mg.conc = 0.0015)
```

```
## [1] 90.06842
```

```
# Mg correction - tanmg07 (mRNA/RNA)
melting(sequence = "UACGCGUCAUAACGCUA",
        nucleic.acid.conc = 0.000002, hybridisation.type = "mrnarna",
        Mg.conc = 0.0015, correction.ion = "tanmg07")
```

```
## [1] 90.06842
```

5.2.3 Mixed Sodium and Magnesium corrections

The available correction methods for mixed sodium magnesium concentration are given in **Table 23**.

Table 23: Details of the corrections for mixed sodium and magnesium concentration

Correction	Type	Limits/Remarks	Reference
owcmix08*	DNA	Mg $\geq$ 0.0005; Mg $\leq$ 0.6; Na+K+Tris/2 $\geq$ 0.	Owczarzy et al. (2008)
tanmix07	DNA, RNA or 2'-O- MeRNA/RNA	Mg $\geq$ 0.1; Mg $\leq$ 0.3; Na+K+Tris/2 $\geq$ 0.1; Na+K+Tris/2 $\leq$ 0.3.	Tan and Chen (2007)

* Default method for computation.

```
# Mixed Na & Mg correction - default (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015)

## [1] 65.83371

# Mixed Na & Mg correction - owcmix08 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015, correction.ion = "owcmix08")

## [1] 65.83371

# Mixed Na & Mg correction - tanmix07 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015, correction.ion = "tanmix07")

## [1] 63.21723

# Mixed Na & Mg correction - default (RNA/RNA)
melting(sequence = "CAGCCUCGUCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Na.conc = 0.069, Mg.conc = 0.0015)

## [1] 79.40119
```

```
# Mixed Na & Mg correction - tanmix07 (RNA/RNA)
melting(sequence = "CAGCCUGUCGACGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "rnarna",
        Na.conc = 0.069, Mg.conc = 0.0015, correction.ion = "tanmix07")
```

```
## [1] 79.40119
```

```
# Mixed Na & Mg correction - default (mRNA/RNA)
melting(sequence = "UACGCGUCAUAACGCUA",
        nucleic.acid.conc = 0.000002, hybridisation.type = "mrnarna",
        Na.conc = 0.069, Mg.conc = 0.0015)
```

```
## [1] 96.46186
```

```
# Mixed Na & Mg correction - tanmix07 (mRNA/RNA)
melting(sequence = "UACGCGUCAUAACGCUA",
        nucleic.acid.conc = 0.000002, hybridisation.type = "mrnarna",
        Na.conc = 0.069, Mg.conc = 0.0015, correction.ion = "tanmix07")
```

```
## [1] 96.46186
```

The ion correction by Owczarzy et al. (2008) is used by default according to the $\frac{[\text{Mg}^{2+}]^{0.5}}{[\text{Mon}^+]}$ ratio, where $[\text{Mon}^+] = [\text{Na}^+] + [\text{Tris}^+] + [\text{K}^+]$.

If,

- $[\text{K}^+] = 0$, default sodium correction is used;
- $\text{Ratio} < 0.22$, default sodium correction is used;
- $0.22 \leq \text{Ratio} < 6$, default mixed Na and Mg correction is used and
- $\text{Ratio} \geq 6$, default magnesium correction is used.

Note that $[\text{Tris}^+]$ is about half of the total tris buffer concentration.

5.2.4 Sodium equivalent concentration methods

The available correction methods for mixed sodium magnesium concentration are given in **Table 24**.

Table 24: Details of the methods for computation of sodium equivalent concentration in the presence of other ions

Correction	Type	Limits/Remarks	Reference
ahs01*	DNA		Ahsen et al. (2001)
mit96	DNA		Mitsuhashi (1996)
pey00	DNA		Peyret (2000)

* Default method for computation.

```

# Na equivalent concentration method - default (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015)

## [1] 65.83371

# Na equivalent concentration method - ahs01 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015, method.Naeq = "ahs01")

## [1] 65.83371

# Na equivalent concentration method - mit96 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015, method.Naeq = "mit96")

## [1] 65.83371

# Na equivalent concentration method - pey00 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 0.069, Mg.conc = 0.0015, method.Naeq = "pey00")

## [1] 65.83371

```

5.3 Denaturing agent corrections

These include melting temperature corrections for concentration of formamide and DMSO.

5.3.1 DMSO corrections

The available correction methods for DMSO concentration are given in **Table 25**.

Table 25: Details of the corrections for DMSO concentration

Correction	Type	Limits/Remarks	Reference
ahs01*	DNA	Not tested with experimental results.	Ahsen et al. (2001)
cu176	DNA	Not tested with experimental results.	Cullen and Bick (1976)

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Correction	Type	Limits/Remarks	Reference
esc80	DNA	Not tested with experimental results.	Escara and Hutton (1980)
mus81	DNA	Not tested with experimental results.	Musielski et al. (1981)

* Default method for computation.

```
# DMSO correction - default (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
         Na.conc = 1, DMSO.conc = 10)
```

```
## [1] 65.40154
```

```
# DMSO correction - ahs01 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
         Na.conc = 1, DMSO.conc = 10, correction.DMSO = "ahs01")
```

```
## [1] 65.40154
```

```
# DMSO correction - cul76 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
         Na.conc = 1, DMSO.conc = 10, correction.DMSO = "cul76")
```

```
## [1] 67.90154
```

```
# DMSO correction - esc80 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
         Na.conc = 1, DMSO.conc = 10, correction.DMSO = "esc80")
```

```
## [1] 66.15154
```

```
# DMSO correction - mus80 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
         nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
         Na.conc = 1, DMSO.conc = 10, correction.DMSO = "mus81")
```

```
## [1] 66.90154
```

5.3.2 Formamide corrections

The available correction methods for formamide concentration are given in **Table 26**.

Table 26: Details of the corrections for formamide concentration

Correction	Type	Limits/Remarks	Reference
bla96*	DNA	With formamide concentration in mol/L.	Blake (1996)
lincorr	DNA	With a % of formamide volume.	McConaughy et al. (1969), Record (1967), Casey and Davidson (1977), Hutton (1977)

* Default method for computation.

```
# Formamide correction - default (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 1, formamide.conc = 0.06)

## [1] 72.74867

# Formamide correction - bla96 (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 1, formamide.conc = 0.06, correction.formamide = "bla96")

## [1] 72.74867

# Formamide correction - lincorr (DNA/DNA)
melting(sequence = "CAGCCTCGTCGCAGC",
        nucleic.acid.conc = 0.000002, hybridisation.type = "dnadna",
        Na.conc = 1, formamide.conc = 10, correction.formamide = "lincorr")

## [1] 66.40154
```

6 Equivalent options in MELTING 5

The options in MELTING 5 command line equivalent to the arguments in `rmelting` are given in **Table 27**.

Table 27: Arguments in `rmelting` and their equivalent options in MELTING 5 command line.

<code>rmelting</code>	MELTING 5 (command line)
<code>sequence</code>	<code>-S</code>
<code>comp.sequence</code>	<code>-C</code>
<code>nucleic.acid.conc</code>	<code>-P</code>
<code>hybridisation.type</code>	<code>-H</code>
<code>Na.conc</code>	<code>-E</code>
<code>Mg.conc</code>	<code>-E</code>
<code>Tris.conc</code>	<code>-E</code>
<code>K.conc</code>	<code>-E</code>
<code>dNTP.conc</code>	<code>-E</code>
<code>DMSO.conc</code>	<code>-E</code>
<code>formamide.conc</code>	<code>-E</code>
<code>size.threshold</code>	<code>-T</code>
<code>self</code>	<code>-self</code>
<code>correction.factor</code>	<code>-F</code>
<code>method.approx</code>	<code>-am</code>
<code>method.nn</code>	<code>-nn</code>
<code>method.GU</code>	<code>-GU</code>
<code>method.singleMM</code>	<code>-sinMM</code>
<code>method.tandemMM</code>	<code>-tanMM</code>
<code>method.single.dangle</code>	<code>-sinDE</code>
<code>method.double.dangle</code>	<code>-secDE</code>
<code>method.long.dangle</code>	<code>-lonDE</code>
<code>method.internal.loop</code>	<code>-intLP</code>
<code>method.single.bulge.loop</code>	<code>-sinBU</code>
<code>method.long.bulge.loop</code>	<code>-lonBU</code>
<code>method.CNG</code>	<code>-CNG</code>
<code>method.inosine</code>	<code>-ino</code>
<code>method.hydroxyadenine</code>	<code>-ha</code>
<code>method.azobenzenes</code>	<code>-azo</code>
<code>method.locked</code>	<code>-lck</code>
<code>method.consecutive.locked</code>	<code>-tanLck</code>
<code>method.consecutive.locked.singleMM</code>	<code>-sinMMLck</code>
<code>correction.ion</code>	<code>-ion</code>
<code>method.Naeq</code>	<code>-naeq</code>

7 Batch computation

Melting temperature for multiple nucleic acid duplexes can be computed using the `meltingBatch` function.

```
sequence <- c("CAAAAAG", "CAAAAAG", "TTTATAATAAA", "CCATCGCTACC",
              "CAAAACAAAG", "CCATTGCTACC", "CAAAAAAAG", "GTGAAC", "AAAAA",
              "CAACTTGATATTATTA", "CAAATAAAG", "GCGAGC", "GGGACC",
```

```

"CAAAGAAAG", "CTGACAAGTGTC", "GCGAAAAGCG")

meltingBatch(sequence, nucleic.acid.conc = 0.0004,
              hybridisation.type = "dnadna", Na.conc = 1)

```

```

##      Environment.Sequence Environment.Complementary sequence
## [1,] "CAAAAAG"          "GTTTTTC"
## [2,] "CAAAAAG"          "GTTTTTC"
## [3,] "TTTATAATAAA"      "AAAATATTATT"
## [4,] "CCATCGCTACC"      "GGTAGCGATGG"
## [5,] "CAAACAAAG"        "GTTTGTTTC"
## [6,] "CCATTGCTACC"      "GGTAACGATGG"
## [7,] "CAAAAAAAG"        "GTTTTTTTC"
## [8,] "GTGAAC"           "CACTTG"
## [9,] "AAAAAAA"          "TTTTTTT"
## [10,] "CAACTTGATATTATTA" "GTTGAACTATAATAAT"
## [11,] "CAAATAAAG"        "GTTTATTTC"
## [12,] "GCGAGC"          "CGCTCG"
## [13,] "GGGACC"           "CCCTGG"
## [14,] "CAAAGAAAG"        "GTTTCTTTC"
## [15,] "CTGACAAGTGTC"     "GACTGTTTCACAG"
## [16,] "GCGAAAAGCG"       "CGCTTTTCGC"
##      Environment.Nucleic acid concentration (M) Environment.Hybridization type
## [1,] "4e-04"              "dnadna"
## [2,] "4e-04"              "dnadna"
## [3,] "4e-04"              "dnadna"
## [4,] "4e-04"              "dnadna"
## [5,] "4e-04"              "dnadna"
## [6,] "4e-04"              "dnadna"
## [7,] "4e-04"              "dnadna"
## [8,] "4e-04"              "dnadna"
## [9,] "4e-04"              "dnadna"
## [10,] "4e-04"             "dnadna"
## [11,] "4e-04"             "dnadna"
## [12,] "4e-04"             "dnadna"
## [13,] "4e-04"             "dnadna"
## [14,] "4e-04"             "dnadna"
## [15,] "4e-04"             "dnadna"
## [16,] "4e-04"             "dnadna"
##      Environment.Na concentration (M) Environment.Mg concentration (M)
## [1,] "1"                   "0"
## [2,] "1"                   "0"
## [3,] "1"                   "0"
## [4,] "1"                   "0"
## [5,] "1"                   "0"

```

```
## [6,] "1" "0"
## [7,] "1" "0"
## [8,] "1" "0"
## [9,] "1" "0"
## [10,] "1" "0"
## [11,] "1" "0"
## [12,] "1" "0"
## [13,] "1" "0"
## [14,] "1" "0"
## [15,] "1" "0"
## [16,] "1" "0"
## Environment.Tris concentration (M) Environment.K concentration (M)
## [1,] "0" "0"
## [2,] "0" "0"
## [3,] "0" "0"
## [4,] "0" "0"
## [5,] "0" "0"
## [6,] "0" "0"
## [7,] "0" "0"
## [8,] "0" "0"
## [9,] "0" "0"
## [10,] "0" "0"
## [11,] "0" "0"
## [12,] "0" "0"
## [13,] "0" "0"
## [14,] "0" "0"
## [15,] "0" "0"
## [16,] "0" "0"
## Environment.dNTP concentration (M) Environment.DMSO concentration (%)
## [1,] "0" "0"
## [2,] "0" "0"
## [3,] "0" "0"
## [4,] "0" "0"
## [5,] "0" "0"
## [6,] "0" "0"
## [7,] "0" "0"
## [8,] "0" "0"
## [9,] "0" "0"
## [10,] "0" "0"
## [11,] "0" "0"
## [12,] "0" "0"
## [13,] "0" "0"
## [14,] "0" "0"
## [15,] "0" "0"
## [16,] "0" "0"
## Environment.Formamide concentration (M or %)
```

```
## [1,] "0"
## [2,] "0"
## [3,] "0"
## [4,] "0"
## [5,] "0"
## [6,] "0"
## [7,] "0"
## [8,] "0"
## [9,] "0"
## [10,] "0"
## [11,] "0"
## [12,] "0"
## [13,] "0"
## [14,] "0"
## [15,] "0"
## [16,] "0"
##      Environment.Self complementarity Environment.Correction factor
## [1,] "FALSE"                                "4"
## [2,] "FALSE"                                "4"
## [3,] "FALSE"                                "4"
## [4,] "FALSE"                                "4"
## [5,] "FALSE"                                "4"
## [6,] "FALSE"                                "4"
## [7,] "FALSE"                                "4"
## [8,] "FALSE"                                "4"
## [9,] "FALSE"                                "4"
## [10,] "FALSE"                               "4"
## [11,] "FALSE"                               "4"
## [12,] "FALSE"                               "4"
## [13,] "FALSE"                               "4"
## [14,] "FALSE"                               "4"
## [15,] "FALSE"                               "4"
## [16,] "FALSE"                               "4"
##      Options.Approximative formula Options.Nearest neighbour model
## [1,] NA                                NA
## [2,] NA                                NA
## [3,] NA                                NA
## [4,] NA                                NA
## [5,] NA                                NA
## [6,] NA                                NA
## [7,] NA                                NA
## [8,] NA                                NA
## [9,] NA                                NA
## [10,] NA                               NA
## [11,] NA                               NA
## [12,] NA                               NA
```

```
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.GU model Options.Single mismatch model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Tandem mismatch model Options.Single dangling end model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Double dangling end model Options.Long dangling end model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
```

```
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Internal loop model Options.Single bulge loop model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Long bulge loop model Options.CNG repeats model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Inosine bases model Options.Hydroxyadenine bases model
## [1,] NA NA
## [2,] NA NA
```

```
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Azobenzenes model Options.Locked nucleic acids model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Ion correction method Options.Na equivalence correction method
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
```

```

## [15,] NA NA
## [16,] NA NA
## Options.DMSO correction method Options.Formamide correction method
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## Options.Mode Results.Enthalpy (cal) Results.Entropy (cal)
## [1,] NA "-47700" "-138.1"
## [2,] NA "-55600" "-160.3"
## [3,] NA "-78800" "-229.7"
## [4,] NA "-83500" "-227"
## [5,] NA "-64600" "-183.2"
## [6,] NA "-81100" "-222.5"
## [7,] NA "-63500" "-182.5"
## [8,] NA "-41200" "-117.5"
## [9,] NA "-50700" "-147.2"
## [10,] NA "-113800" "-323.6"
## [11,] NA "-62100" "-179.8"
## [12,] NA "-46000" "-124.8"
## [13,] NA "-40400" "-109.9"
## [14,] NA "-63700" "-181.3"
## [15,] NA "-90400" "-249.5"
## [16,] NA "-80300" "-218.6"
## Results.Enthalpy (J) Results.Entropy (J) Results.Melting temperature (C)
## [1,] "-199386" "-577.258" "31.7814953255144"
## [2,] "-232408" "-670.054" "38.1103863719918"
## [3,] "-329384" "-960.146" "44.5553259145469"
## [4,] "-349030" "-948.86" "67.2098590318261"
## [5,] "-270028" "-765.776" "47.400072116762"
## [6,] "-338998" "-930.05" "63.6040550863501"
## [7,] "-265430" "-762.85" "43.0400604037136"
## [8,] "-172216" "-491.15" "30.1735038367475"
## [9,] "-211926" "-615.296" "33.1415226764116"

```

```
## [10,] "-475684"          "-1352.648"          "59.6680431282422"
## [11,] "-259578"          "-751.564"           "40.2828264688437"
## [12,] "-192280"          "-521.664"           "48.2393469411973"
## [13,] "-168872"          "-459.382"           "41.9123740666287"
## [14,] "-266266"          "-757.834"           "45.9425910944819"
## [15,] "-377872"          "-1042.91"           "64.379329012421"
## [16,] "-335654"          "-913.748"           "65.7707030297917"
##      Message
## [1,] NA
## [2,] NA
## [3,] NA
## [4,] NA
## [5,] NA
## [6,] NA
## [7,] NA
## [8,] NA
## [9,] NA
## [10,] NA
## [11,] NA
## [12,] NA
## [13,] NA
## [14,] NA
## [15,] NA
## [16,] NA
```

Complementary sequences are computed by default, but need to be specified in case of mismatches, inosine(s) or hydroxyadenine(s) between the two strands.

```
seq <- c("GCAUACG", "CAGUAGGUC", "CGCUCGC", "GAGUGGAG", "GACAGGCUC",
         "CAGUACGUC", "GACAUCUG", "GACCACUG", "CAGAAUGUC", "GCGUCGC",
         "CGUCCGG", "GACUCUCUG", "CAGCUGGUC", "GACUAGCUC", "CUCUGCUC",
         "GCGUCCG", "GUCCGCG", "CGAUCAC", "GACUACCUG", "GACGAUCUG")

comp.seq <- c("CGUUUGC", "GUCGGCCAG", "GCGUGCG", "CUCUUCUC", "CUGUGCGAC",
              "GUCGGGCAG", "CUGUUGGAC", "CUGGGGGAC", "GUCUGGCAG", "CGCUGCG",
              "GCUGGCC", "CUGAUAGAC", "GUCGUUCAG", "CUGAGCGAC", "GAGUUGAG",
              "CGCUGGC", "CUGGCGC", "GCUUGUG", "CUGAGGGAC", "CUGCCAGAC")

meltingBatch(sequence = seq, comp.seq = comp.seq, nucleic.acid.conc = 0.0004,
              hybridisation.type = "rnarna", Na.conc = 1,
              method.singleMM = "tur06")
```

```
##      Environment.Sequence Environment.Complementary sequence
## [1,] "GCAUACG"          "CGUUUGC"
## [2,] "CAGUAGGUC"        "GUCGGCCAG"
## [3,] "CGCUCGC"          "GCGUGCG"
## [4,] "GAGUGGAG"         "CUCUUCUC"
```

```

## [5,] "GACAGGCUG"      "CUGUGCGAC"
## [6,] "CAGUACGUC"      "GUCGGGCAG"
## [7,] "GACAUCCUG"      "CUGUUGGAC"
## [8,] "GACCACCUG"      "CUGGGGGAC"
## [9,] "CAGAAUGUC"      "GUCUGGCAG"
## [10,] "GCGUCGC"       "CGCUGCG"
## [11,] "CGUCCGG"       "GCUGGCC"
## [12,] "GACUCUCUG"     "CUGAUAGAC"
## [13,] "CAGCUGGUC"     "GUCGUUCAG"
## [14,] "GACUAGCUG"     "CUGAGCGAC"
## [15,] "CUCUGCUC"      "GAGUUGAG"
## [16,] "GCGUCCG"       "CGCUGGC"
## [17,] "GUCCGCG"       "CUGGCGC"
## [18,] "CGAUCAC"       "GCUUGUG"
## [19,] "GACUACCUG"     "CUGAGGGAC"
## [20,] "GACGAUCUG"     "CUGCCAGAC"
##      Environment.Nucleic acid concentration (M) Environment.Hybridization type
## [1,] "4e-04"                                     "rna"
## [2,] "4e-04"                                     "rna"
## [3,] "4e-04"                                     "rna"
## [4,] "4e-04"                                     "rna"
## [5,] "4e-04"                                     "rna"
## [6,] "4e-04"                                     "rna"
## [7,] "4e-04"                                     "rna"
## [8,] "4e-04"                                     "rna"
## [9,] "4e-04"                                     "rna"
## [10,] "4e-04"                                    "rna"
## [11,] "4e-04"                                    "rna"
## [12,] "4e-04"                                    "rna"
## [13,] "4e-04"                                    "rna"
## [14,] "4e-04"                                    "rna"
## [15,] "4e-04"                                    "rna"
## [16,] "4e-04"                                    "rna"
## [17,] "4e-04"                                    "rna"
## [18,] "4e-04"                                    "rna"
## [19,] "4e-04"                                    "rna"
## [20,] "4e-04"                                    "rna"
##      Environment.Na concentration (M) Environment.Mg concentration (M)
## [1,] "1"                                     "0"
## [2,] "1"                                     "0"
## [3,] "1"                                     "0"
## [4,] "1"                                     "0"
## [5,] "1"                                     "0"
## [6,] "1"                                     "0"
## [7,] "1"                                     "0"
## [8,] "1"                                     "0"

```

```
## [9,] "1" "0"
## [10,] "1" "0"
## [11,] "1" "0"
## [12,] "1" "0"
## [13,] "1" "0"
## [14,] "1" "0"
## [15,] "1" "0"
## [16,] "1" "0"
## [17,] "1" "0"
## [18,] "1" "0"
## [19,] "1" "0"
## [20,] "1" "0"
## Environment.Tris concentration (M) Environment.K concentration (M)
## [1,] "0" "0"
## [2,] "0" "0"
## [3,] "0" "0"
## [4,] "0" "0"
## [5,] "0" "0"
## [6,] "0" "0"
## [7,] "0" "0"
## [8,] "0" "0"
## [9,] "0" "0"
## [10,] "0" "0"
## [11,] "0" "0"
## [12,] "0" "0"
## [13,] "0" "0"
## [14,] "0" "0"
## [15,] "0" "0"
## [16,] "0" "0"
## [17,] "0" "0"
## [18,] "0" "0"
## [19,] "0" "0"
## [20,] "0" "0"
## Environment.dNTP concentration (M) Environment.DMSO concentration (%)
## [1,] "0" "0"
## [2,] "0" "0"
## [3,] "0" "0"
## [4,] "0" "0"
## [5,] "0" "0"
## [6,] "0" "0"
## [7,] "0" "0"
## [8,] "0" "0"
## [9,] "0" "0"
## [10,] "0" "0"
## [11,] "0" "0"
## [12,] "0" "0"
```

```
## [13,] "0" "0"
## [14,] "0" "0"
## [15,] "0" "0"
## [16,] "0" "0"
## [17,] "0" "0"
## [18,] "0" "0"
## [19,] "0" "0"
## [20,] "0" "0"
##      Environment.Formamide concentration (M or %)
## [1,] "0"
## [2,] "0"
## [3,] "0"
## [4,] "0"
## [5,] "0"
## [6,] "0"
## [7,] "0"
## [8,] "0"
## [9,] "0"
## [10,] "0"
## [11,] "0"
## [12,] "0"
## [13,] "0"
## [14,] "0"
## [15,] "0"
## [16,] "0"
## [17,] "0"
## [18,] "0"
## [19,] "0"
## [20,] "0"
##      Environment.Self complementarity Environment.Correction factor
## [1,] "FALSE" "4"
## [2,] "FALSE" "4"
## [3,] "FALSE" "4"
## [4,] "FALSE" "4"
## [5,] "FALSE" "4"
## [6,] "FALSE" "4"
## [7,] "FALSE" "4"
## [8,] "FALSE" "4"
## [9,] "FALSE" "4"
## [10,] "FALSE" "4"
## [11,] "FALSE" "4"
## [12,] "FALSE" "4"
## [13,] "FALSE" "4"
## [14,] "FALSE" "4"
## [15,] "FALSE" "4"
## [16,] "FALSE" "4"
```

```
## [17,] "FALSE" "4"
## [18,] "FALSE" "4"
## [19,] "FALSE" "4"
## [20,] "FALSE" "4"
## Options.Approximative formula Options.Nearest neighbour model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.GU model Options.Single mismatch model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
```

```

##      Options.Tandem mismatch model Options.Single dangling end model
## [1,] NA                          NA
## [2,] NA                          NA
## [3,] NA                          NA
## [4,] NA                          NA
## [5,] NA                          NA
## [6,] NA                          NA
## [7,] NA                          NA
## [8,] NA                          NA
## [9,] NA                          NA
## [10,] NA                         NA
## [11,] NA                         NA
## [12,] NA                         NA
## [13,] NA                         NA
## [14,] NA                         NA
## [15,] NA                         NA
## [16,] NA                         NA
## [17,] NA                         NA
## [18,] NA                         NA
## [19,] NA                         NA
## [20,] NA                         NA
##      Options.Double dangling end model Options.Long dangling end model
## [1,] NA                          NA
## [2,] NA                          NA
## [3,] NA                          NA
## [4,] NA                          NA
## [5,] NA                          NA
## [6,] NA                          NA
## [7,] NA                          NA
## [8,] NA                          NA
## [9,] NA                          NA
## [10,] NA                         NA
## [11,] NA                         NA
## [12,] NA                         NA
## [13,] NA                         NA
## [14,] NA                         NA
## [15,] NA                         NA
## [16,] NA                         NA
## [17,] NA                         NA
## [18,] NA                         NA
## [19,] NA                         NA
## [20,] NA                         NA
##      Options.Internal loop model Options.Single bulge loop model
## [1,] NA                          NA
## [2,] NA                          NA
## [3,] NA                          NA

```

```
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.Long bulge loop model Options.CNG repeats model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.Inosine bases model Options.Hydroxyadenine bases model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
```

```
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.Azobenzenes model Options.Locked nucleic acids model
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.Ion correction method Options.Na equivalence correction method
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
```

--- Computation of melting temperature of nucleic acid duplexes with `rmelting`

```
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.DMSO correction method Options.Formamide correction method
## [1,] NA NA
## [2,] NA NA
## [3,] NA NA
## [4,] NA NA
## [5,] NA NA
## [6,] NA NA
## [7,] NA NA
## [8,] NA NA
## [9,] NA NA
## [10,] NA NA
## [11,] NA NA
## [12,] NA NA
## [13,] NA NA
## [14,] NA NA
## [15,] NA NA
## [16,] NA NA
## [17,] NA NA
## [18,] NA NA
## [19,] NA NA
## [20,] NA NA
## Options.Mode Results.Enthalpy (cal) Results.Entropy (cal)
## [1,] NA "-47650" "-138.8"
## [2,] NA "-71130" "-200.5"
## [3,] NA "-57930" "-164.1"
## [4,] NA "-60570" "-176.6"
## [5,] NA "-79870" "-219.9"
## [6,] NA "-68380" "-194.5"
## [7,] NA "-73880" "-208.3"
## [8,] NA "-78430" "-218.3"
## [9,] NA "-59650" "-171.5"
## [10,] NA "-61330" "-173.8"
## [11,] NA "-58350" "-165.4"
## [12,] NA "-64570" "-184.7"
## [13,] NA "-70970" "-200.6"
## [14,] NA "-72010" "-203"
## [15,] NA "-58820" "-171"
```

```

## [16,] NA          "-59840"          "-169.6"
## [17,] NA          "-59840"          "-169.6"
## [18,] NA          "-50210"          "-148.3"
## [19,] NA          "-70520"          "-198.8"
## [20,] NA          "-69730"          "-198.2"
##      Results.Enthalpy (J) Results.Entropy (J) Results.Melting temperature (C)
## [1,] "-199177"          "-580.184"          "30.1048299398322"
## [2,] "-297323.4"        "-838.09"          "51.8989532242754"
## [3,] "-242147.4"        "-685.938"          "44.3989325444856"
## [4,] "-253182.6"        "-738.188"          "37.5791954133529"
## [5,] "-333856.6"        "-919.182"          "62.1162425798375"
## [6,] "-285828.4"        "-813.01"          "48.141439592185"
## [7,] "-308818.4"        "-870.694"          "52.845957204839"
## [8,] "-327837.4"        "-912.494"          "58.2977096620104"
## [9,] "-249337"          "-716.87"          "41.08087522322"
## [10,] "-256359.4"        "-726.484"          "46.0633145887674"
## [11,] "-243903"          "-691.372"          "44.4380466975271"
## [12,] "-269902.6"        "-772.046"          "44.8840464672343"
## [13,] "-296654.6"        "-838.508"          "51.0196486690585"
## [14,] "-301001.8"        "-848.54"          "52.203376709706"
## [15,] "-245867.6"        "-714.78"          "37.5268181873443"
## [16,] "-250131.2"        "-708.928"          "45.2688421309843"
## [17,] "-250131.2"        "-708.928"          "45.2688421309843"
## [18,] "-209877.8"        "-619.894"          "28.1788644808993"
## [19,] "-294773.6"        "-830.984"          "51.6345164549562"
## [20,] "-291471.4"        "-828.476"          "48.8860141674642"
##      Message
## [1,] NA
## [2,] NA
## [3,] NA
## [4,] NA
## [5,] NA
## [6,] NA
## [7,] NA
## [8,] NA
## [9,] NA
## [10,] NA
## [11,] NA
## [12,] NA
## [13,] NA
## [14,] NA
## [15,] NA
## [16,] NA
## [17,] NA
## [18,] NA
## [19,] NA

```

```
## [20,] NA
```

8 Further reading

Further details about algorithm, formulae and methods are available in the [MELTING 5 documentation](#).

9 Citing `rmelting`

```
## To cite the R package 'rmelting' in publications use:
##
## Aravind, J. and Krishna, G. K. (2023).  rmelting: R Interface to
## MELTING 5. R package version 1.17.0,
## https://aravind-j.github.io/rmelting/.
##
## A BibTeX entry for LaTeX users is
##
## @Manual{,
##   title = {rmelting: R Interface to MELTING 5},
##   author = {J. Aravind and G. K. Krishna},
##   year = {2023},
##   note = {R package version 1.17.0},
##   note = {https://aravind-j.github.io/rmelting/},
## }
##
## This free and open-source software implements academic research by the
## authors and co-workers. If you use it, please support the project by
## citing the package.
```

10 Session Info

```
sessionInfo()
```

```
## R version 4.3.0 RC (2023-04-18 r84287)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 22.04.2 LTS
##
## Matrix products: default
## BLAS: /home/biocbuild/bbs-3.18-bioc/R/lib/libRblas.so
## LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
##
## locale:
##  [1] LC_CTYPE=en_US.UTF-8          LC_NUMERIC=C
##  [3] LC_TIME=en_GB                 LC_COLLATE=C
```

```
## [5] LC_MONETARY=en_US.UTF-8      LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=en_US.UTF-8         LC_NAME=en_US.UTF-8
## [9] LC_ADDRESS=en_US.UTF-8       LC_TELEPHONE=en_US.UTF-8
## [11] LC_MEASUREMENT=en_US.UTF-8   LC_IDENTIFICATION=en_US.UTF-8
##
## time zone: America/New_York
## tzcode source: system (glibc)
##
## attached base packages:
## [1] stats      graphics  grDevices  utils      datasets  methods    base
##
## other attached packages:
## [1] rmelting_1.17.0 readxl_1.4.2
##
## loaded via a namespace (and not attached):
## [1] digest_0.6.31   utf8_1.2.3      fastmap_1.1.1   xfun_0.39
## [5] cellranger_1.1.0 magrittr_2.0.3   glue_1.6.2      tibble_3.2.1
## [9] knitr_1.42      pkgconfig_2.0.3  htmltools_0.5.5 rJava_1.0-6
## [13] rmarkdown_2.21  lifecycle_1.0.3  Rdpack_2.4      cli_3.6.1
## [17] pander_0.6.5    fansi_1.0.4      vctrs_0.6.2     compiler_4.3.0
## [21] rbibutils_2.2.13 tools_4.3.0      pillar_1.9.0    evaluate_0.20
## [25] Rcpp_1.0.10     yaml_2.3.7       rlang_1.1.0
```

References

- Ahsen, N. von, Wittwer, C. T., and Schütz, E. (2001). Oligonucleotide melting temperatures under PCR conditions: Nearest-neighbor corrections for Mg²⁺, deoxynucleotide triphosphate, and dimethyl sulfoxide concentrations with comparison to alternative empirical formulas. *Clinical Chemistry* 47, 1956–1961. Available at: <http://clinchem.aaccjnls.org/content/47/11/1956>.
- Allawi, H. T., and SantaLucia, J. (1997). Thermodynamics and NMR of internal G · T mismatches in dna. *Biochemistry* 36, 10581–10594. doi:[10.1021/bi962590c](https://doi.org/10.1021/bi962590c).
- Allawi, H. T., and SantaLucia, J. (1998a). Nearest neighbor thermodynamic parameters for internal G · A mismatches in DNA. *Biochemistry* 37, 2170–2179. doi:[10.1021/bi9724873](https://doi.org/10.1021/bi9724873).
- Allawi, H. T., and SantaLucia, J. (1998b). Nearest-neighbor thermodynamics of internal A · C mismatches in dna: Sequence dependence and pH effects. *Biochemistry* 37, 9435–9444. doi:[10.1021/bi9803729](https://doi.org/10.1021/bi9803729).
- Allawi, H. T., and SantaLucia, J. (1998c). Thermodynamics of internal C · T mismatches in DNA. *Nucleic Acids Research* 26, 2694–2701. doi:[10.1093/nar/26.11.2694](https://doi.org/10.1093/nar/26.11.2694).
- Asanuma, H., Matsunaga, D., and Komiyama, M. (2005). Clear-cut photo-

regulation of the formation and dissociation of the DNA duplex by modified oligonucleotide involving multiple azobenzenes. *Nucleic Acids Symposium Series*, 35–36. doi:[10.1093/nass/49.1.35](https://doi.org/10.1093/nass/49.1.35).

Badhwar, J., Karri, S., Cass, C. K., Wunderlich, E. L., and Znosko, B. M. (2007). Thermodynamic characterization of RNA duplexes containing naturally occurring 1×2 nucleotide internal loops. *Biochemistry* 46, 14715–14724. doi:[10.1021/bi701024w](https://doi.org/10.1021/bi701024w).

Blake, R. D. (1996). “Denaturation of DNA,” in *Encyclopedia of molecular biology and molecular medicine*, ed. R. A. Meyers (Weinheim, Germany: VCH Verlagsgesellschaft), 1–19.

Blake, R. D., and Delcourt, S. G. (1998). Thermal stability of DNA. *Nucleic Acids Research* 26, 3323–3332. doi:[10.1093/nar/26.14.3323](https://doi.org/10.1093/nar/26.14.3323).

Blose, J. M., Manni, M. L., Klappec, K. A., Stranger-Jones, Y., Zyra, A. C., Sim, V., et al. (2007). Non-nearest-neighbor dependence of stability for RNA bulge loops based on the complete set of group i single nucleotide bulge loops. *Biochemistry* 46, 15123–15135. doi:[10.1021/bi700736f](https://doi.org/10.1021/bi700736f).

Bommarito, S., Peyret, N., and SantaLucia, J. (2000). Thermodynamic parameters for DNA sequences with dangling ends. *Nucleic Acids Research* 28, 1929–1934. doi:[10.1093/nar/28.9.1929](https://doi.org/10.1093/nar/28.9.1929).

Breslauer, K. J., Frank, R., Blöcker, H., and Marky, L. A. (1986). Predicting DNA duplex stability from the base sequence. *Proceedings of the National Academy of Sciences* 83, 3746. doi:[10.1073/pnas.83.11.3746](https://doi.org/10.1073/pnas.83.11.3746).

Britten, R. J., Graham, D. E., and Neufeld, B. R. (1974). Analysis of repeating DNA sequences by reassociation. *Methods in Enzymology* 29, 363–418. doi:[10.1016/0076-6879\(74\)29033-5](https://doi.org/10.1016/0076-6879(74)29033-5).

Broda, M., Kierzek, E., Gdaniec, Z., Kulinski, T., and Kierzek, R. (2005). Thermodynamic stability of RNA structures formed by CNG trinucleotide repeats. Implication for prediction of RNA structure. *Biochemistry* 44, 10873–10882. doi:[10.1021/bi0502339](https://doi.org/10.1021/bi0502339).

Casey, J., and Davidson, N. (1977). Rates of formation and thermal stabilities of RNA:DNA and DNA:DNA duplexes at high concentrations of formamide. *Nucleic Acids Research* 4, 1539–1552. doi:[10.1093/nar/4.5.1539](https://doi.org/10.1093/nar/4.5.1539).

Chen, J. L., Dishler, A. L., Kennedy, S. D., Yildirim, I., Liu, B., Turner, D. H., et al. (2012). Testing the nearest neighbor model for canonical rna base pairs: Revision of GU parameters. *Biochemistry* 51, 3508–3522. doi:[10.1021/bi3002709](https://doi.org/10.1021/bi3002709).

Chester, N., and Marshak, D. R. (1993). Dimethyl sulfoxide-mediated primer Tm reduction: A method for analyzing the role of renaturation temperature in the polymerase chain reaction. *Analytical Biochemistry* 209, 284–290. doi:[10.1006/abio.1993.1121](https://doi.org/10.1006/abio.1993.1121).

- Cullen, B. R., and Bick, M. D. (1976). Thermal denaturation of DNA from bromodeoxyuridine substituted cells. *Nucleic Acids Research* 3, 49–62. doi:[10.1093/nar/3.1.49](https://doi.org/10.1093/nar/3.1.49).
- Davis, A. R., and Znosko, B. M. (2007). Thermodynamic characterization of single mismatches found in naturally occurring RNA. *Biochemistry* 46, 13425–13436. doi:[10.1021/bi701311c](https://doi.org/10.1021/bi701311c).
- Davis, A. R., and Znosko, B. M. (2008). Thermodynamic characterization of naturally occurring RNA single mismatches with G-U nearest neighbors. *Biochemistry* 47, 10178–10187. doi:[10.1021/bi800471z](https://doi.org/10.1021/bi800471z).
- Dumousseau, M., Rodriguez, N., Juty, N., and Le Novère, N. (2012). MELTING, a flexible platform to predict the melting temperatures of nucleic acids. *BMC Bioinformatics* 13, 101. doi:[10.1186/1471-2105-13-101](https://doi.org/10.1186/1471-2105-13-101).
- Escara, J. F., and Hutton, J. R. (1980). Thermal stability and renaturation of DNA in dimethyl sulfoxide solutions: Acceleration of the renaturation rate. *Biopolymers* 19, 1315–1327. doi:[10.1002/bip.1980.360190708](https://doi.org/10.1002/bip.1980.360190708).
- Frank-Kamenetskii, M. D. (1971). Simplification of the empirical relationship between melting temperature of DNA, its GC content and concentration of sodium ions in solution. *Biopolymers* 10, 2623–2624. doi:[10.1002/bip.360101223](https://doi.org/10.1002/bip.360101223).
- Freier, S. M., Kierzek, R., Jaeger, J. A., Sugimoto, N., Caruthers, M. H., Neilson, T., et al. (1986). Improved free-energy parameters for predictions of RNA duplex stability. *Proceedings of the National Academy of Sciences* 83, 9373. doi:[10.1073/pnas.83.24.9373](https://doi.org/10.1073/pnas.83.24.9373).
- Hall, T. J., Grula, J. W., Davidson, E. H., and Britten, R. J. (1980). Evolution of sea urchin non-repetitive DNA. *Journal of Molecular Evolution* 16, 95–110. doi:[10.1007/BF01731580](https://doi.org/10.1007/BF01731580).
- Hutton, J. R. (1977). Renaturation kinetics and thermal stability of DNA in aqueous solutions of formamide and urea. *Nucleic Acids Research* 4, 3537–3555. doi:[10.1093/nar/4.10.3537](https://doi.org/10.1093/nar/4.10.3537).
- Kawakami, J., Kamiya, H., Yasuda, K., Fujiki, H., Kasai, H., and Sugimoto, N. (2001). Thermodynamic stability of base pairs between 2-hydroxyadenine and incoming nucleotides as a determinant of nucleotide incorporation specificity during replication. *Nucleic Acids Research* 29, 3289–3296. doi:[10.1093/nar/29.16.3289](https://doi.org/10.1093/nar/29.16.3289).
- Kierzek, E., Mathews, D. H., Ciesielska, A., Turner, D. H., and Kierzek, R. (2006). Nearest neighbor parameters for Watson-Crick complementary heteroduplexes formed between 2'-O-methyl RNA and RNA oligonucleotides. *Nucleic Acids Research* 34, 3609–3614. doi:[10.1093/nar/gkl232](https://doi.org/10.1093/nar/gkl232).
- Le Novère, N. (2001). MELTING, computing the melting temperature of nucleic acid duplex. *Bioinformatics* 17, 1226–1227. doi:[10.1093/bioinformatics/17.12.1226](https://doi.org/10.1093/bioinformatics/17.12.1226).
- Lu, Z. J., Turner, D. H., and Mathews, D. H. (2006). A set of nearest neighbor parameters for predicting the enthalpy change of RNA secondary structure

- formation. *Nucleic Acids Research* 34, 4912–4924. doi:[10.1093/nar/gkl472](https://doi.org/10.1093/nar/gkl472).
- Marmur, J., and Doty, P. (1962). Determination of the base composition of deoxyribonucleic acid from its thermal denaturation temperature. *Journal of Molecular Biology* 5, 109–118. doi:[10.1016/S0022-2836\(62\)80066-7](https://doi.org/10.1016/S0022-2836(62)80066-7).
- Mathews, D. H., Sabina, J., Zuker, M., and Turner, D. H. (1999). Expanded sequence dependence of thermodynamic parameters improves prediction of RNA secondary structure. *Journal of Molecular Biology* 288, 911–940. doi:[10.1006/jmbi.1999.2700](https://doi.org/10.1006/jmbi.1999.2700).
- McConaughy, B. L., Laird, C., and McCarthy, B. J. (1969). Nucleic acid reassociation in formamide. *Biochemistry* 8, 3289–3295. doi:[10.1021/bi00836a024](https://doi.org/10.1021/bi00836a024).
- McTigue, P. M., Peterson, R. J., and Kahn, J. D. (2004). Sequence-dependent thermodynamic parameters for locked nucleic acid (LNA)-DNA duplex formation. *Biochemistry* 43, 5388–5405. doi:[10.1021/bi035976d](https://doi.org/10.1021/bi035976d).
- Miller, S., Jones, L. E., Giovannitti, K., Piper, D., and Serra, M. J. (2008). Thermodynamic analysis of 5' and 3' single- and 3' double-nucleotide overhangs neighboring wobble terminal base pairs. *Nucleic Acids Research* 36, 5652–5659. doi:[10.1093/nar/gkn525](https://doi.org/10.1093/nar/gkn525).
- Mitsuhashi, M. (1996). Technical report: Part 1. Basic requirements for designing optimal oligonucleotide probe sequences. *Journal of Clinical Laboratory Analysis* 10, 277–284. doi:[10/cw9bn6](https://doi.org/10/cw9bn6).
- Musielski, H., Mann, W., Laue, R., and Michel, S. (1981). Influence of dimethylsulfoxide on transcription by bacteriophage T3-induced RNA polymerase. *Zeitschrift für allgemeine Mikrobiologie* 21, 447–456. doi:[10.1002/jobm.19810210606](https://doi.org/10.1002/jobm.19810210606).
- Ohmichi, T., Nakano, S.-i., Miyoshi, D., and Sugimoto, N. (2002). Long RNA dangling end has large energetic contribution to duplex stability. *Journal of the American Chemical Society* 124, 10367–10372. doi:[10.1021/ja0255406](https://doi.org/10.1021/ja0255406).
- O'Toole, A. S., Miller, S., Haines, N., Zink, M. C., and Serra, M. J. (2006). Comprehensive thermodynamic analysis of 3' double-nucleotide overhangs neighboring Watson-Crick terminal base pairs. *Nucleic Acids Research* 34, 3338–3344. doi:[10.1093/nar/gkl428](https://doi.org/10.1093/nar/gkl428).
- O'Toole, A. S., Miller, S., and Serra, M. J. (2005). Stability of 3' double nucleotide overhangs that model the 3' ends of siRNA. *RNA* 11, 512–516. doi:[10.1261/rna.7254905](https://doi.org/10.1261/rna.7254905).
- Owczarzy, R., Moreira, B. G., You, Y., Behlke, M. A., and Walder, J. A. (2008). Predicting stability of DNA duplexes in solutions containing magnesium and monovalent cations. *Biochemistry* 47, 5336–5353. doi:[10.1021/bi702363u](https://doi.org/10.1021/bi702363u).
- Owczarzy, R., You, Y., Groth, C. L., and Tataurov, A. V. (2011). Stability and mismatch discrimination of locked nucleic acid–DNA duplexes. *Biochemistry* 50, 9352–9367. doi:[10.1021/bi200904e](https://doi.org/10.1021/bi200904e).

- Owczarzy, R., You, Y., Moreira, B. G., Manthey, J. A., Huang, L., Behlke, M. A., et al. (2004). Effects of sodium ions on DNA duplex oligomers: Improved predictions of melting temperatures. *Biochemistry* 43, 3537–3554. doi:[10.1021/bi034621r](https://doi.org/10.1021/bi034621r).
- Owen, R., Hill, L., and Lapage, S. (1969). Determination of DNA base compositions from melting profiles in dilute buffers. *Biopolymers* 7, 503–516. doi:[10.1002/bip.1969.360070408](https://doi.org/10.1002/bip.1969.360070408).
- Peyret, N. (2000). Prediction of nucleic acid hybridization: Parameters and algorithms. Available at: <http://elibrary.wayne.edu/record=2760965>.
- Peyret, N., Seneviratne, P. A., Allawi, H. T., and SantaLucia, J. (1999). Nearest-Neighbor Thermodynamics and NMR of DNA Sequences with Internal A · A, C · C, G · G, and T · T Mismatches. *Biochemistry* 38, 3468–3477. doi:[10.1021/bi9825091](https://doi.org/10.1021/bi9825091).
- Record, M. T. (1967). Electrostatic effects on polynucleotide transitions. I. Behavior at neutral pH. *Biopolymers* 5, 975–992. doi:[10.1002/bip.1967.360051010](https://doi.org/10.1002/bip.1967.360051010).
- SantaLucia, J. (1998). A unified view of polymer, dumbbell, and oligonucleotide DNA nearest-neighbor thermodynamics. *Proceedings of the National Academy of Sciences* 95, 1460. doi:[10.1073/pnas.95.4.1460](https://doi.org/10.1073/pnas.95.4.1460).
- SantaLucia, J., and Hicks, D. (2004). The thermodynamics of DNA structural motifs. *Annual Review of Biophysics and Biomolecular Structure* 33, 415–440. doi:[10.1146/annurev.biophys.32.110601.141800](https://doi.org/10.1146/annurev.biophys.32.110601.141800).
- SantaLucia, John, Allawi, H. T., and Seneviratne, P. A. (1996). Improved nearest-neighbor parameters for predicting DNA duplex stability. *Biochemistry* 35, 3555–3562. doi:[10.1021/bi951907q](https://doi.org/10.1021/bi951907q).
- Schildkraut, C., and Lifson, S. (1965). Dependence of the melting temperature of DNA on salt concentration. *Biopolymers* 3, 195–208. doi:[10.1002/bip.360030207](https://doi.org/10.1002/bip.360030207).
- Sugimoto, N., Nakano, S., Yoneyama, M., and Honda, K. (1996). Improved thermodynamic parameters and helix initiation factor to predict stability of DNA duplexes. *Nucleic Acids Research* 24, 4501–4505. doi:[10.1093/nar/24.22.4501](https://doi.org/10.1093/nar/24.22.4501).
- Tan, Z.-J., and Chen, S.-J. (2006). Nucleic acid helix stability: Effects of salt concentration, cation valence and size, and chain length. *Biophysical Journal* 90, 1175–1190. doi:[10.1529/biophysj.105.070904](https://doi.org/10.1529/biophysj.105.070904).
- Tan, Z.-J., and Chen, S.-J. (2007). RNA helix stability in mixed Na(+)/Mg(2+) solution. *Biophysical Journal* 92, 3615–3632. doi:[10.1529/biophysj.106.100388](https://doi.org/10.1529/biophysj.106.100388).
- Tanaka, F., Kameda, A., Yamamoto, M., and Ohuchi, A. (2004). Thermodynamic parameters based on a nearest-neighbor model for DNA sequences with a single-bulge loop. *Biochemistry* 43, 7143–7150. doi:[10.1021/bi036188r](https://doi.org/10.1021/bi036188r).
- Wahl, G. M., Barger, S. L., and Kimmel, A. R. (1987). Molecular hybridization of immobilized nucleic acids: Theoretical concepts and practical considerations.

Methods in Enzymology 152, 399–407. doi:[10.1016/0076-6879\(87\)52046-8](https://doi.org/10.1016/0076-6879(87)52046-8).

Watkins, N. E., Kennelly, W. J., Tsay, M. J., Tuin, A., Swenson, L., Lee, H.-R., et al. (2011). Thermodynamic contributions of single internal rA · dA, rC · dC, rG · dG and rU · dT mismatches in RNA/DNA duplexes. *Nucleic Acids Research* 39, 1894–1902. doi:[10/cdm4jh](https://doi.org/10/cdm4jh).

Watkins, N. E., and SantaLucia, J. (2005). Nearest-neighbor thermodynamics of deoxyinosine pairs in DNA duplexes. *Nucleic Acids Research* 33, 6258–6267. doi:[10.1093/nar/gki918](https://doi.org/10.1093/nar/gki918).

Wetmur, J. G. (1991). DNA probes: Applications of the principles of nucleic acid hybridization. *Critical Reviews in Biochemistry and Molecular Biology* 26, 227–259. doi:[10.3109/10409239109114069](https://doi.org/10.3109/10409239109114069).

Wright, D. J., Rice, J. L., Yanker, D. M., and Znosko, B. M. (2007). Nearest neighbor parameters for inosine · uridine pairs in RNA duplexes. *Biochemistry* 46, 4625–4634. doi:[10.1021/bi0616910](https://doi.org/10.1021/bi0616910).

Xia, T., SantaLucia, J., Burkard, M. E., Kierzek, R., Schroeder, S. J., Jiao, X., et al. (1998). Thermodynamic parameters for an expanded nearest-neighbor model for formation of RNA duplexes with Watson-Crick base pairs. *Biochemistry* 37, 14719–14735. doi:[10.1021/bi9809425](https://doi.org/10.1021/bi9809425).