# MiRduplexSVM - Help file

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## <u>MiRduplexSVM - miRNA: miRNA\* duplex prediction.</u>

This tool predicts the most likely miRNA duplex on a given miRNA hairpin and offers two modes of operation.

- 1. You can upload a text file (.txt) containing your sequences in fasta format, including the fasta file header.
- 2. You can paste your sequences (including the fasta file header) directly into the provided textbox, which is the default operation.

The output is presented as a three-column tab-delimited table. The first column lists the sequence IDs as specified by the user, while the second and third columns display the most probable matures from the 5' and the 3' strands of the hairpin, respectively. Entries for which the RNAfold output is not a hairpin or consists of multiple stems are filtered out, and "Multibranch - No prediction" is returned as an output for the respective lines.

If you wish to receive your results via email, you can provide your email address. Thank you for using *MiRduplexSVM duplex prediction*.

*Input Example:* 

>MI0000060

AGGUUGAGGUAGGUUGUAUAGUUUAGAAUUACAUCAAGGGAGAUAACUGUACAGCCUCCUAGCUUUCCU>MI0000065

CCUAGGAAGAGGUAGUAGGUUGCAUAGUUUUAGGGCAGGGAUUUUGCCCACAAGGAGGUAACUAUACGACCUGCUGCCUUUC UUAGG

>MI0003603

UCUUAUCAAUGAGGUAGACCAUGGGUUCUCAUUGUAAUAGUGUAGAAUGUUGGUUAACUGUGGACUCCCUGGCUCUGUCUCA AAUCUACUGAUUC

>MI0000070

GUCAGCAGUGCCUUAGCAGCACGUAAAUAUUGGCGUUAAGAUUCUAAAAUUAUCUCCAGUAUUAACUGUGCUGAAGUAA GGUUGAC

>MI0003681

>MI0000071

GUCAGAAUAAUGUCAAAGUGCUUACAGUGCAGGUAGUGAUAUGUGCAUCUACUGCAGUGAAGGCACUUGUAGCAUUAUGGUG

Output example:

sequenceID 5strandMature 3strandMature

MI0000060	UGAGGUAGUAGGUUGUAUAGUU	CUAUACAAUCUACUGUCUUUC
MI0000061	UGAGGUAGUAGGUUGUAUAGUU	CUGUACAGCCUCCUAGCUUUCC
MI0000065	AGAGGUAGUAGGUUGCAUAGUU	CUAUACGACCUGCUGCCUUUC
MI0003603	Multibranch - No predicti	on
MI0000070	UAGCAGCACGUAAAUAUUGGCG	CCAGUAUUAACUGUGCUGCUGA
MI0003681	UGGAGAAGCGUGGACCGGUCC	CAGGUUCUCACCCUCUCUAGG
MI0000071	CAAAGUGCUUACAGUGCAGGUAG	ACUGCAGUGAAGGCACUUGUAG

## MiRduplexSVM: miRNA Evaluator - Missing duplex finder.

This tool evaluates a mature miRNA using MiRduplexSVM's score and predicts the mature molecule located on the opposite strand. It has two modes of operation.

You can either upload two text files (.txt) with your sequences in fasta format, including the fasta file header. One file should contain the miRNA precursor sequences; the other should contain the respective mature miRNA sequences. Alternatively, you can paste your sequences in fasta format (include fasta file header) in the appropriate text boxes in the online form.

The output is a four-column tab-delimited table. The first column specifies the sequences' IDs as provided by the user. The second and third columns show the matures from the 5' and 3' strands of the hairpin. One mature corresponds to the one provided by the user, and the other one is the program's prediction. Entries for which the RNAfold output is not a hairpin or consists of multiple stems are filtered out, and "Multibranch - No prediction" is returned as an output for the respective line.

If you wish to receive your results via email, you can provide your email address. Thank you for using *MiRduplexSVM*: *Evaluator – Missing duplex finder*.

*Input Example - precursors sequences:* 

>MI0000064

GCAUCCGGGUUGAGGUAGGUUGUAUGGUUUAGAGUUACACCCUGGGAGUUAACUGUACAACCUUCUAGCUUUCCUUGGA

>MI0000238

GUGAAUUAGGUAGUUUCAUGUUGGGCCUGGGUUUCUGAACACAACAUUAAACCACCCGAUUCAC

>MI0003603

UCUUAUCAAUGAGGUAGACCAUGGGUUCUCAUUGUAAUAGUGUAGAAUGUUGGUUAACUGUGGACUCCCUGGCUCUGUCUCA AAUCUACUGAUUC

>MI0000240

UCAUUGGUCCAGAGGGGAGAUAGGUUCCUGUGAUUUUUCCUUCUUCUCUAUAGAAUAAAUGA

>MI0003681

>MI0000265

AGAUUAGAGUGGCUGUGGUCUAGUGCUGUGGAAGACUAGUGAUUUUGUUGUUCUGAUGUACUACGACAACAAGUCACAGC CGGCCUCAUAGCGCAGACUCCCUUCGAC

>MI0000291

AAUAUUCUGUAUGACUGUGCUACUUCAA

*Input Example - mature sequences:* 

>MI0000064

UGAGGUAGUAGGUUGUAUGGUU

>MI0000238

UAGGUAGUUUCAUGUUGUUGGG

>MI0003603

AGACCAUGGGUUCUCAUUGU

>MI0000240

GGUCCAGAGGGGAGAUAGGUUC

>MI0003681

GGCAGGUUCUCACCCUCUCUAGG

>MI0000265

UGGAAGACUAGUGAUUUUGUUGU

>MI0000291

AUGACCUAUGAAUUGACAGAC

#### Output example:

sequenceID	5strandMature	3strandMature
MI0000064	UGAGGUAGUAGGUUGUAUGGUU	CUGUACAACCUUCUAGCUUUCC
MI0000238	UAGGUAGUUUCAUGUUGUUGGG	CAACAACAUUAAACCACCCGAU
MI0003603	Multibranch - No predict:	ion
MI0000240	GGUCCAGAGGGGAGAUAGGUUC	UCCUUCUUCUAUAGAAUAAA
MI0003681	UGGAGAAGCGUGGACCGGUCCG	GGCAGGUUCUCACCCUCUCUAGG
MI0000265	UGGAAGACUAGUGAUUUUGUUGU	CAACAAGUCACAGCCGGCCUCA
MI0000291	AUGACCUAUGAAUUGACAGAC	UCUGUCAUUUCUUUAGGCCAAU

## <u>MiRduplexSVM - miRNA:miRNA\* Duplex Evaluator.</u>

This tool evaluates a miRNA:miRNA\* duplex by assigning MiRduplexSVM's score. It has two modes of operation.

You can either upload three text files (.txt) with your sequences in fasta format, including the fasta file header. The first file should contain the miRNA precursor sequences; the second should contain the mature miRNA sequences produced by the 3' strand, and the third file the mature miRNA sequences produced by the 5' strand. Alternatively, you can paste your sequences in fasta format (include fasta file header) in the appropriate text boxes in the online form.

The output is a four-column tab-delimited table. The first column specifies the sequences' IDs as provided by the user. The second and third columns show the matures from the 5' and 3' strands of the hairpin, and the fourth column represents MiRduplexSVM's score. Entries for which the RNAfold output is not a hairpin or consists of multiple stems are filtered out, and "Multibranch - No prediction" is returned as an output for the respective line.

If you wish to receive your results via email, you can provide your email address. Thank you for using *MiRduplexSVM – miRNA:miRNA\* Duplex Evaluator*.

*Input Example - precursors sequences:* 

>MI0000063

CGGGGUGAGGUAGUAGGUUGUGGUUUCAGGGCAGUGAUGUUGCCCCUCGGAAGAUAACUAUACAACCUACUGCCUUCCCU

>MI0000286

ACCCGGCAGUGCCUCCAGGCGCAGGGCAGCCCCUGCCCACCGCACACUGCGCUGCCCCAGACCCACUGUGCGUGUGACAGCGGCUGAUCUGUGCCUGGGCAGCGCGACCC

>MI0000447

UGAGCUGUUGGAUUCGGGGCCGUAGCACUGUCUGAGAGGUUUACAUUUCUCACAGUGAACCGGUCUCUUUUUUCAGCUGCUUC >MI0000450

ACAAUGCUUUGCUAGAGCUGGUAAAAUGGAACCAAAUCGCCUCUUCAAUGGAUUUGGUCCCCUUCAACCAGCUGUAGCUAUGCUUGA

>MI0000451

GGGAGCCAAAUGCUUUGCUAGAGCUGGUAAAAUGGAACCAAAUCGACUGUCCAAUGGAUUUGGUCCCCUUCAACCAGCUGUA GCUGUGCAUUGAUGGCGCCG

>MI0000474

#### *Input Example - mature sequences from 3' strand:*

>MI0000063

CUAUACAACCUACUGCCUUCCC

>MI0000286

CUGUGCGUGUGACAGCGGCUGA

>MI0000447

UCACAGUGAACCGGUCUCUUU

>MI0000450

UUUGGUCCCCUUCAACCAGCUG

>MI0000451

UUUGGUCCCCUUCAACCAGCUG

>MI0000474

CCUGUGGGCCACCUAGUCACCAA

### *Input Example - mature sequences from 5' strand:*

>MI0000063

UGAGGUAGUAGGUUGUGUGGUU

>MI0000286

AGCCCCUGCCCACCGCACACUG

>MI0000447

CGGGGCCGUAGCACUGUCUGAGA

>MI0000450

AGCUGGUAAAAUGGAACCAAAU

>MI0000451

AGCUGGUAAAAUGGAACCAAAU

>MI0000474

UGUGACUGGUUGACCAGAGGGG

#### Output example:

sequenceID	5strandMature	3strandMature	MiRduplexSVM-score
MI0000063	UGAGGUAGUAGGUUGUGUGGUU	CUAUACAACCUACUGCCUUCCC	9.780586e-01
MI0000286	AGCCCCUGCCCACCGCACACUG	CUGUGCGUGUGACAGCGGCUGA	1.175949e+00

MI0000447	CGGGGCCGUAGCACUGUCUGAGA	UCACAGUGAACCGGUCUCUUU	1.003558e+00
MI0000450	AGCUGGUAAAAUGGAACCAAAU	UUUGGUCCCCUUCAACCAGCUG	1.380121e+00
MI0000451	AGCUGGUAAAAUGGAACCAAAU	UUUGGUCCCCUUCAACCAGCUG	1.121841e+00
MI0000474	UGUGACUGGUUGACCAGAGGGG	CCUGUGGGCCACCUAGUCACCAA	-6.557550e-02