OPTIMIR, a novel algorithm for integrating available genomewide genotype data into miRNA sequence alignment analysis.

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Abstract

Next-generation sequencing is an increasingly popular and efficient approach to characterize the full set of microRNAs (miRNAs) present in human biosamples. MiRNAs' detection and quantification still remain a challenge as they can undergo different post transcriptional modifications and might harbor genetic variations (polymiRs) that may impact on the alignment step. We present a novel algorithm, OPTIMIR, that incorporates biological knowledge on miRNA editing and genome-wide genotype data available in the processed samples to improve alignment accuracy.

OPTIMIR was applied to 391 human plasma samples that had been typed with genome-wide genotyping arrays. OPTIMIR was able to detect genotyping errors, suggested the existence of novel miRNAs and highlighted the allelic imbalance expression of polymiRs in heterozygous carriers.

OPTIMIR is written in python, and freely available on the GENMED website (http://www.genmed.fr/index.php/fr/) and on Github (github.com/FlorianThibord/OptimiR).

Introduction

With an average length of 22 nucleotides, microRNAs (miRNAs) belong to a class of small non-coding RNAs known to regulate gene expression by binding messenger RNAs (mRNAs) and interfering with the translational machinery (Filipowicz et al., 2008). MiRNAs are transcribed from primary miRNA sequences (pri-miRNAs) and fold into a hairpin-like structure, which is sequentially processed by two ribonucleases, DROSHA and DICER. The former cleaves the pri-miRNA into a pre-miRNA and the latter completes the miRNA's maturation by cleaving the pre-miRNA near its loop to produce a miRNA duplex composed of 2 mature strands (Kim et al., 2009). Exceptionally, some miRNAs follow a slightly different pathway where only one ribonuclease is needed to complete the maturation (Kim et al., 2016). In any case, only one of the two mature strands is loaded in an effective protein complex called RISC, while the other is degraded (Kawamata and Tomari, 2010). This selection seems mostly driven by the thermodynamic stability of both ends forming the duplex (Meijer et al., 2014).

There is emerging interest in performing miRNA profiling in body fluids or tissues in order to identify novel molecular determinants of human diseases (Mitchell et al., 2008; Pulcrano-Nicolas et al., 2018). Such miRNA profiling can be achieved using either hybridization (microarray), Next Generation Sequencing (NGS) or Real Time-quantitative Polymerase Chain Reaction (RT-qPCR) techniques. With 2,588 known mature miRNAs in humans according to miRBase version 21 (Kozomara and Griffiths-Jones, 2014), RTqPCR would be cumbersome on a genomic scale, but is widely recognized as a gold standard for the validation of few miRNAs. The NGS technology is becoming more popular than microarrays because of its greater detection sensitivity, and higher accuracy in differential expression analysis (Git et al., 2010; Tam et al., 2014). NGS applied to small RNAs revealed a great diversity in the sequences of mature miRNAs originating from the same hairpin. This diversity is mostly attributable to the deletion and addition of nucleotides at the miRNAs' extremities (also known as trimming and tailing events, respectively), due to the activity of terminal nucleotidyl transferases, exoribonucleases, or imprecise cleavage by DROSHA and DICER (Wyman et al., 2011; Neilsen et al., 2012; Ameres and Zamore, 2013). To a lesser extent, the ADAR protein acting on double stranded RNAs and responsible for A-to-I editing is also known to target miRNAs (Nishikura, 2016). These post-transcriptional editing mechanisms have been shown to affect miRNAs' function and stability (Kawahara et al., 2007; Chiang et al., 2010; Burroughs et al., 2010; Katoh et al., 2015).

Lastly, genetic variations have also been shown to contribute to the sequence diversity of miRNAs, and to affect their function and expression (Mencia et al., 2009; Gong et al., 2012; Han and Zheng, 2013; Cammaerts et al., 2015). In the following, we will refer to miRNAs harbouring genetic polymorphisms in their miRNA sequence as "polymiRs". We use the "isomiRs" terminology for miRNAs that underwent post-transcriptional editing events. Of note, polymiRs can also be subject to editing events and thus can also be isomiRs.

The first step in the bioinformatics analysis of miRNA sequencing (miRSeq) data consists in aligning sequenced reads to a reference library of mature miRNAs. This step may be challenging because 1 - the aforementioned variability of isomiRs could lead to imperfect alignments to the reference library; 2 sequenced reads may correspond to (fragments of) other molecules (e.g. other small non coding RNAs like piRNA, tRNA, yRNA, ...), captured during the preparation of the libraries, that might share a high similarity with miRNAs because of their small length and thus might be confused with miRNAs (Chen and Heard, 2013; Heintz-Buschart et al., 2018); 3 - some miRNAs have homologous sequences that are identical or very similar, thus a single read might align ambiguously to multiple reference sequences. In this work, we investigate the impact of the presence of polymorphisms in the sequence of mature miRNAs on their alignment and their expression in the context of miRSeq profiling applied to samples that have also been typed for genome-wide genotype data. A situation we anticipate to become rather common with the rise of increasingly affordable genome-wide association studies (GWAS) and the decreasing cost of next generation exome/genome sequencing techniques. In that context, we developed an original bioinformatics workflow called OPTIMIR, for pOlymorPhisminTegratIon for MIRna data alignment, that integrates genetic information from genotyping arrays or DNA sequencing into the miRSeq data alignment process with the aim of improving the accuracy of polymiRs alignment, while accommodating for isomiRs detection and ambiguously aligned reads. In addition, OPTIMIR allows to assess the association of genotypes on polymiRs with corresponding polymiRs' expression. OPTIMIR was evaluated in the plasma samples of 391 individuals, part of the MARTHA study (Oudot-Mellakh et al., 2012).

Results

OPTIMIR was composed of three main steps (Materials and Methods). First, miRSeq data are aligned to

a reference library upgraded with sequences integrating alternative alleles of genetic variations. A correction

is then applied for ambiguous and unreliable alignments via a scoring approach. Finally, polymiR alignments

are evaluated to only retain those that are consistent with input genotypes in case these have been provided

by the user.

MiRNA alignment

OPTIMIR was evaluated on 391 miRNA sequencing data files totalizing 7,390,947,662 sequencing

reads. After the pre-alignment step, 2,922,446,965 reads (39.54 % of total reads) remained for alignment.

562,040,494 of these reads (19.23%) were then mapped to mature miRNA reference sequences, of which

10,937,479 (1.95% of mapped reads) aligned ambiguously to two sequences or more. The application of the

OPTIMIR scoring algorithm for alignment disambiguation resulted in a unique solution for 91.6% of these

cross-mapping reads. For 89.5% of reads with multiple alignments, the difference between the two lowest

scores was greater than 2 (Figure 1) which would correspond to alignments that differ from each other by at

least two modifications in the 3' end.

The distribution of alignment scores after disambiguation is shown in Figure 2. Scores ranged from 0 to

76 with 98.0% of alignment scores lower or equal to 9. Beyond this threshold, the curve rapidly increased

indicating that scores with higher values are very sparse and suggesting that such alignments with a very

high number of editing events are likely improper alignments. As a consequence, for the following, we

decided to discard any alignments with a score greater than 9. The resulting distribution of alignment score

for the 550,946,055 remaining reads was given in Figure 3.

IsomiRs distribution

197,808,779 (35.9%) reads perfectly aligned to mature miRNAs, such reads being generally referred to

4

as canonical miRNAs, and received an alignment score of 0. This confirms previous observations suggesting

that a substantial amount of miRNAs are mainly represented by alternative isomiRs (Wu et al., 2018;

Wallaert et al., 2017). The distribution of 3' and 5' ends modifications on mapped reads observed over the

391 samples processed by OPTIMIR is shown in Figure 4.

The most common editing events are on miRNAs' 3' end, with ~34% of trimming, ~17% tailing with

non-templated nucleotides, ~5% tailing with templated nucleotides, and a similar proportion of trimming

events followed by tailing. The latter modification could also be interpreted as nucleotide variation due to

genetic variants or sequencing error. But such events generally occur at a much lesser frequency, suggesting

that a combination of a trimming event due to imprecise cleavage followed by a tailing event is more likely,

especially when we consider the high frequency at which they occur independently.

The 5' end is much less frequently edited, with 94% of reads having no editing on this extremity.

Nevertheless, the most frequently observed modification on 5' ends was trimming that affected 4.4% of all

reads. This may be of biological relevance since such trimming could shift the miRNA binding seed that is

crucial for the miRNA to bind to its mRNA targets.

Alignment on polymiRs

Over all samples, 220,156 reads mapped to 46 polymiRs for a total of 1,786 distinct alignments. As

detailed in Table 1, 19 polymiRs have reads that aligned to an alternative sequence that was introduced in the

upgraded reference library. Two polymiRs (hsa-miR-6796-3p and hsa-miR-1269b) harbor two SNPs, and for

both of them only the reference sequence with both common alleles were found to be expressed. Among the

remaining 44 polymiRs that harbor only one SNP, 15 were expressed with both alleles. It is important to

mention that the allele present in the miRBase reference sequence may not be the most common one (e.g. the

rs2155248 and hsa-miR-1304-3p) which may lead to improperly discard reads if stringent alignment (with

no mismatch allowed) to the original miRBase library is applied.

In total, 721 (0.33% of reads mapped to polymiRs) alignments were found inconsistent with the individual

genotypes among which 507 were unlikely due to sequencing errors. The latter situations involved 10

5

samples and 5 polymiRs that are detailed in Table 2 and further investigated in the next section.

Investigation of inconsistent genotypes

The first case of inconsistent genotype concerns individual PVP28 imputed to be homozygous for the rs12473206-G allele while showing numerous reads mapping to both versions of the hsa-miR-4433b-3p polymiR. Sanger re-sequencing revealed that this individual is in fact heterozygous for this variant which is much more compatible with the number of observed reads at this locus. A rather similar inconsistent genotype was observed for individual SSP20 but lack of available DNA did not allow us to perform Sanger validation. Lack of DNA also prevented us from investigating deeper the inconsistent genotypes observed for rs72631820/hsa-miR-339-3p and rs6771809/hsa-miR-6826-5p.

The inconsistent genotype observed for individual AJA9 at rs6841938 and hsa-miR-1255-5p was challenging. Sanger re-sequencing confirmed that this individual was indeed homozygous for the rs6841938-G allele although it has numerous reads mapping to both versions of hsa-miR-1255-5p. However, the use of BLAST web-service from NCBI (Altschul et al., 1990) revealed that reads aligned on the polymiR sequence containing the rs6841938-A allele perfectly match to the chr1:167,998,699-167,998,720 region but on the opposite strand where hsa-miR-1255-3p (no yet reported) should be located. This observation could be compatible with the presence on this opposite strand of an unreported miRNA locus as it has already been observed for other miRNAs (e.g. hsa-mir-4433a and hsa-mir-4433b). Going back to the discarded alignments revealed that 18 other samples homozygous for rs6841938-G had 1 or 2 reads that mapped to this opposite strand. These alignments had been discarded as one could not have distinguished them from sequencing errors (see *Genotype consistency analysis* section in the Material & Methods).

The last inconsistent genotypes were observed at rs3817551/hsa-miR-7107-3p for five independent individuals. For 4 of them with available DNA, Sanger re-sequencing confirmed the initial homozygous genotype for rs3817551-G allele while these individuals were found to express only the hsa-miR-7107-3p version with the T allele. On average, these inconsistent alignments received an alignment score of 1.41, and none of the reads involved mapped to other sequences. Concerning the samples that had reads aligned on this same reference with a consistent genotype, the average alignment score was very close to 1.47. This score indicates that reads share a high sequence similarity with the hsa-miR-7107-3p, and there is no significant difference between the group with an inconsistent genotype, and the group with a consistent one. BLAST analysis did not enable us to identify any homologous sequence that could explain these observations that

still remain to be further investigated in order to be sure that associated reads do well originate from hsa-

miR-7107-3p.

Analysis of polymiRs allele specific expression

As shown in Table 1, 29 polymiRs with a SNP in their mature sequence were found to be expressed in

individuals heterozygotes for this SNP. While one could anticipate that, in heterozygous individuals for such

SNP, the polymiRs could have balanced expression (as measured by read counts) of both the reference and

alternative sequences, this was hardly observed. Indeed, we observed a strong preference for either the

reference or the alternative version of the polymiR in heterozygotes.

Figure 5 shows the alignments of 4 polymiRs involving heterozygous variant according to the vcf

genotype. The dotted line represents a theoretical balanced expression between reference and alternative

sequences. We can see that for hsa-miR-1255b-5p/rs6841938 and hsa-miR-5189-3p/rs35613341 situations

are close to the y-axis, which are situations were only the reference is expressed. The polymiR with the

expression closest to allelic balance for many samples is hsa-miR-4433b-3p but, even then, the average rate

of alternative reads across all 147 samples is of 0.8 (see Supplementary Table 4 for details on polymiRs

involved in heterozygous situations).

As an illustration, 65 MARTHA individuals were heterozygous for the rs2925980 variant associated with

polymiR hsa-miR-7854-3p. These genotypes could be considered as reliable as this SNP was directly typed

on the array. Among these 65 individuals, 54 expressed only the alternative version of polymiR hsa-miR-

7854-3p and the remaining 11 expressed both polymiR versions with a mean ratio of 0.96 in favor of the

alternative allele.

Finally, we used the RNAfold program (Lorenz et al., 2011) to predict the secondary structure of the 29

pri-miRNA (i.e. hairpins) induced by the presence of a SNP in the polymiR sequence (Supplementary Figure

S1). Most genetic variations create either a new bulge, or a wobble pairing, or have no impact on the

secondary structure. A notable exception relates to rs35613341 located on the hsa-miR-5189-3p where the G

allele completely changed the secondary structure of the hairpin (see Supplementary Figure S1-u) making it

difficult to access for the DICER and DROSHA machinery. In MARTHA samples, 161 individuals were

heterozygous and 34 homozygous for the rs35613341-G allele. None of these individuals were found to

express the alternative sequence of polymiR hsa-miR-5189-3p which could support the hypothesis that the

rs35613341-G allele impacts the maturation of this miRNA.

Lastly, by the completion of the OPTIMIR pipeline on MARTHA samples, 7.45% of sequenced reads

were aligned. This value had to be compared with 7.68% and 8.24% obtained by 2 other recent pipelines for

miRSeq data, sRNAnalyzer (Wu et al., 2017) and miRge (Baras et al., 2015), respectively. These

discrepancies could be explained by the less stringency of these programs allowing up to 2 mismatches.

The files generated by OPTIMIR include: 1/ global abundances of miRNAs (counts of isomiRs and

polymiRs are merged with the reference mature sequences' counts); 2/ specific abundances for each polymiR

sequence; 3/ specific abundances for each isomiR sequence; 4/ alignments that are inconsistent with

provided genotypes; 5/ two annotation files containing details on templated nucleotides and alignments that

could not be disambiguated.

Discussion

In this work, we propose a novel algorithm, OPTIMIR, for aligning miRNA sequences obtained from

next generation sequencing and we applied it to plasma samples of 391 individuals from the MARTHA

study. Borrowing some ideas from other alignment pipelines such as the addition of new sequences to the

reference library corresponding to allelic versions of polymiRs(Baras et al., 2014; Russell et al., 2018) and a

scoring strategy for handling cross-mapping reads (Urgese et al., 2016) or for discarding unlikely isomiRs

(Bofill-De Ros et al., 2018), OPTIMIR has two features that make it unique. First, OPTIMIR is based on a

scoring strategy that incorporate biological knowledge on miRNA editing to identify the most likely

alignment in presence of cross-mapping reads. Second, OPTIMIR allows the user to provide genotype

information, in particular data obtained from genome wide genotyping arrays, to improve alignment

accuracy. This option revealed several interesting observations when OPTIMIR was applied to MARTHA

plasma samples.

First, it allowed to identify improperly imputed genotypes despite overall good imputation quality.

Second, it suggested the existence of a new miRNA not indexed in the miRBase v21 database that would be

located on the opposite strand of the hsa-miR-1255b-5p. Thirdly, it suggested that reads aligned to hsa-miR-7107-3p are likely false alignments and would more likely come from other non-coding fragments that share sequence similarity with hsa-miR-7107-3p. These last two hypotheses would need to be further validated but this is out of the scope of the bioinformatics workflow described in the current work. Even more interesting was the study of polymiRs' expressions in heterozygous individuals for SNPs on these polymiRs. OPTIMIR clearly showed that plasma allelic expression of polymiRs is unbalanced for most polymiRs. One allelic version of a polymiR is much more expressed than the other and this is not necessarily the one carrying the most common allele. This observation is consistent with previous works showing that SNPs in (pri-) miRNA sequences can influence miRNA expression through their impact on the DROSHA/DICER RNAses' machinery (Duan et al., 2007; Cammaerts et al., 2015). Epigenetic mechanisms could also explain the allelic imbalance expression of miRNAs (Morales et al., 2017).

Several limitations shall however be acknowledged. OPTIMIR requires to fix two parameters, a weight W_3 for penalizing 5' end editing event (set to 4 in the current application) and a score threshold (set to 9 here) to discard unreliable alignments. The former tends to have little impact on the general findings (see Supplementary Table 2) while the second may be study-specific and may depend on the number of studied samples in the study and the kind of tissue analyzed. These parameters can be easily modified by the user. There is so far no gold standard program for miRNA alignment analysis but our preliminary study suggests that OPTIMIR aligns slightly less reads than two other recently proposed software, sRNAnalyzer(Wu et al., 2017) and miRge(Baras et al., 2015). This is likely due to the higher number of mismatches allowed by the laters for aligning reads while OPTIMIR tends to be more stringent. Without extensive investigations including experimental validations, it is not possible to really appreciate which alignments are the correct ones.

Finally, several improvements could be considered such as: 1/ the integration of A-to-I editing events in the definition of our reference library and of our scoring strategy, even if we anticipate that it might be difficult to distinguish these rare events from sequencing errors and 2/ the extension of the OPTIMIR workflow to analyze other small coding RNAs (e.g. piRNA and tRNA, rRNA, snRNA or yRNA derived fragments) that are generally sequenced together with miRNAs in a miRseq profiling. Applications to other

tissue from the samples processed for miR-Seq data deserve to be conducted to generalize the findings

observed in the current plasma samples.

Materials & Methods

The MARTHA dataset

The MARseilleTHrombosisAssocation study is a collection of patients with venous thrombosis (VT)

recruited at the La Timone Hospital (Marseille, France) between 1994 and 2005 and aimed at identifying

novel molecular determinants for VT and its associated endophenotypes(Oudot-Mellakh et al., 2012; Dick et

al., 2014).

For the present study, 391 MARTHA participants with available plasma samples were processed for

plasma miRNA profiling through miRSeq. These individuals had been previously typed for genome-wide

genotyping arrays and imputed for single nucleotide polymorphisms (SNPs) available in the 1000G reference

database (Germain et al., 2015).

MiRNA extraction and preparation followed the same protocol as the one previously described in (Roux

et al., 2018). Briefly, from 400µL of plasma, total RNA was first extracted using the miRNeasy

serum/plasma kit for Qiagen. miRNA libraries were then prepared using the NEBNext Multiplex Small RNA

Library Prep Set for adapter ligation and PCR, with adapter sequences

GATCGGAAGAGCACACGTCTGAACTCCAGTCAC (3' adapter) and

CGACAGGTTCAGAGTCTACAGTCCGACGATC (5' adapter) followed by a size selection using

AMPure XP beads. Pools of equal quantity of 24 purified libraries were constructed and tagged with

different indexes. Pools were then sequenced using a 75bp single-end strategy on an Illumina NextSeq500

10

instrument.

The OPTIMIR workflow

Alignment

<u>Pre-alignment data processing</u> - 3' adapters were removed using cutadapt (Martin, 2011) with a base quality filter set to 28. Remaining reads with sequence length between 15 and 27, which generally correspond to miRNA sequences, were then kept for alignment. Note that identical reads were collapsed together to decrease computational burden associated with processing n times n identical reads.

Definition of an alignment reference library - Read alignment generally starts by the selection of a reference library to which reads shall be aligned. The miRBase 21 database (Kozomara and Griffiths-Jones, 2014) containing known human mature miRNAs is usually adopted for miRSeq data. We first upgrade this reference library by adding new sequences corresponding to the alternate forms of polymiRs showing genetic polymorphisms in their mature sequence as previously proposed (Baras et al., 2014; Russell et al., 2018). In case a polymiR contains more than one polymorphism, new sequences corresponding to all possible haplotypes are generated. These variants, that are provided by the OPTIMIR's user in a vcf format file (Danecek et al., 2011), are mapped to miRNAs using miRBase miRNA coordinates file (i.e. positions of miRNAs on the human reference genome). The generation of new sequences is automated via a standalone python script provided with the OPTIMIR pipeline.

For the current application to MARTHA GWAS data, we identified 88 single nucleotide polymorphisms (SNPs) for which we have a reliable genotype data defined as SNPs with imputation $r^2 > 0.8$. Note that some SNPs may map to 2 distinct miRNAs if the latter are transcribed from opposite strands. Some miRNAs may also have more than one SNP in their sequence. In our application, 5 SNPs mapped to 2 distinct miRNAs and 3 miRNAs contained more than one SNP. As a result, the reference library was upgraded with 96 new alternative sequences corresponding to all possible haplotypes derived from the 90 identified polymiRs.

Reads alignment process - For read alignment, we opted for the bowtie2 software (Langmead and Salzberg, 2012) that can handle trimming and tailing events at the reads' extremities via its local alignment mode which has been shown to be efficient for miRSeq data alignment (Ziemann et al., 2016). Only reads with a sequence of at least 17 consecutive nucleotides (defined as the alignment seed) that matches with the reference library without any mismatch are kept in the analysis (see Supplementary Table 1 for details concerning the choice of the seed value and its consequences on isomiR detection). For miRNAs that

underwent tailing events, or trimming events followed by tailing events, additional bases exceeding or

differing from the reference are soft-clipped and do not participate in the alignment. Reads were allowed to

align to multiple reference sequences in order to take into account the different mature miRNAs with similar

sequences from which they could originate. Finally, we did not allow reverse complement alignment as small

RNAs were first ligated with different 5' and 3' adapters before single-end sequencing, which implies that

RNA strands were sequenced in only one direction.

Resolution of ambigous alignments

To handle multiple ambiguous alignments, OPTIMIR integrated a scoring algorithm aimed at identifying the

most plausible alignment(s) while discarding likely erroneous ones. Of note, beforehand, for reads mapping

to a mature miRNA that can be produced by two different pri-miRNAs (e.g. hsa-miR-1255b-5p can originate

from hsa-mir-1255b-1 or hsa-mir-1255b-2 located on chromosome 4 and 1, respectively), we used the

information on templated tailed nucleotides (i.e. nucleotides in the pri-miRNA sequences (also available in

the miRBase 21) that surround the mature miRNA sequence) to deduce from which locus these reads might

come from. This information, that has no impact on the alignment per se, is stored in an output file (named

expressed_hairpins.annot).

Each alignment was assigned a score based on the number of trimming and tailing events that could make a

given read perfectly match with a mature miRNA sequence. Since trimming and tailing are frequently

observed in the 3' end of miRNAs but rare in the 5' end (Neilsen et al., 2012; Wu et al., 2018), a more

penalizing weight was applied on events observed on the 5' end. Alignments with a lower score would be

considered as more reliable as they would correspond to a miRNA with less editing events. The alignment

score is calculated as follow:

Alignment score = $N_5 x W_5 + N_3$

where N_5 and N_3 represent the number of editing events observed on the 5' and 3' extremities of the read,

respectively. W_5 is the weight applied to the events observed on the 5' end. Several W_5 values were tested

and their impact on the alignment results are shown in Supplementary Table 2. Finally, for our application,

W₅ was set to 4 in order to resolve as many ambiguous alignments as possible without penalizing too much 5'

events compared to 3' events as they represent ~60% of the editing events. Templated tailed nucleotides do

not count as editing events as they tend to validate the parentage of a read to its reference. These templated

nucleotides are most likely the result of imprecise cleavage by DROSHA and DICER. However, they might

occasionally result from the action of the terminal nucleotidyl transferase that adds the same nucleotides as

those surrounding the original sequence, in which case they cannot be distinguished.

By the end of the scoring algorithm, the alignment with the lowest score was retained. In case of an

ambiguous read with n possible alignments having the same score, all alignments are kept and assigned to a

weight of 1/n, and corresponding alignments are listed in an output file (named

remaining_ambiguous.annot).

Of note, bowtie2 also integrates an alignment score. However, this scoring is general and does not integrate

the biological knowledge on editing events specific to miRNAs. The OPTIMIR scoring algorithm also

differs from the one recently proposed in IsomiR-SEA pipeline (Urgese et al., 2016) which is based on the

number of observed mismatches and the difference in size between a given read and the reference mature

miRNAs.

Genotype consistency analysis

In case users provide genetic information for individuals that have been miRSeq profiled, the last step of the

OPTIMIR workflow is to provide a comparison analysis of the genotype data provided by the user (in a

standard vcf format) and the genotype data that could be inferred from the sequenced reads aligned to

polymiRs. For an individual whose reads aligned onto a polymiR sequence that harbors the alternate allele of

a SNP, consistency will be called if this individual is either heterozygous or homozygous for this allele in the

provided vcf genotype file. Inconsistent alignments are discarded but saved in an output file (named

inconsistents.sam).

Indeed, it may occur that some reads align to a polymiR sequence that harbors a given allele in an individual

that is not expected to carry it. This could be due to sequencing errors reproducing genetic variations.

However, for a given individual and a given polymiR, if this event is observed for a large number of reads

(e.g. more than 1% of the reads mapping to the polymiR), another explanation must be looked for. For

instance, this could occur when reads originate from sequenced fragments of other small non-coding RNAs

that share a high similarity with a polymiR. Such situations are detailed in a separate output (named

13

consistency_table.annot).

Care is needed to call inconsistency when a polymiR may have homologous mature sequences and one of them is polymorphic. For example, the mature miRNA hsa-miR-1255b-5p can originate either from the pri-miRNA hsa-miR-1255b-1 located on chromosome 4 or from hsa-mir-1255b-2 located on chromosome 1. However, only the chromosome 4 copy contains a variant. If reads with the alternate allele can easily be deduced to originate from hsa-miR-1255b-1, reads with the reference allele can come from both chromosome 1 and 4 copies. As a consequence, an homozygous carriers of the alternate allele can still have reads mapping to chromosome 1 copies and such reads shall not be considered as inconsistent. We have listed in Supplementary Table 3 all mature miRNAs that have multiple pri-miRNAs sequences and tagged those that are polymorphic.

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Tables

<u>Table 1: Summary information at detected polymiRs.</u>

Table 1. Sulli	illuly illioi	mation a	t detectes	a porymi	113.		Misses	han af	
polymiR	rsID		samples exp R with genot		Number of reads aligned on polymiR sequence integrating the allele		Number of inconsistent reads removed on the sequence integrating the allele		Inconsistent alignments to investigate (2)
		0/0	0/1	1/1	reference	alternative	reference	alternative	. ,
hsa-miR-4433b-3p	rs12473206	223 (/230)	147 (/147)	14 (/14)	63 762	96 915	4	475	2
hsa-miR-1255b-5p	rs6841938	244 (/323)	48 (/66)	2 (/2)	29642	66	0	52	1
hsa-miR-1304-3p	rs2155248	0 (/0)	5 (/7)	256 (/384)	146	11 928	4	0	0
hsa-miR-339-3p	rs72631820	213 (/386)	3 (/5)	0 (/0)	6 293	0	0	21	1
hsa-miR-7854-3p	rs2925980	22 (/159)	65 (/167)	35 (/65)	392	2 015	1	4	0
hsa-miR-4745-5p	rs10422347	86 (/300)	14 (/81)	0 (/10)	2 330	15	0	0	0
hsa-miR-5189-3p	rs35613341	42 (/196)	20 (/161)	0 (/34)	1 315	0	2	1	0
hsa-miR-7107-3p	rs3817551	20 (/157)	33 (/177)	0 (/57)	1 104	1	122	0	5
hsa-miR-4741	rs7227168	40 (/293)	4 (/93)	0 (/5)	695	0	0	0	0
hsa-miR-548l	rs13447640	30 (/373)	0 (/18)	0 (/0)	613	0	0	0	0
hsa-miR-3620-5p	rs2070960	47 (/348)	4 (/42)	0 (/1)	459	0	0	0	0
hsa-miR-6826-5p	rs6771809	11 (/286)	11 (/92)	2 (/13)	199	243	0	21	1
hsa-miR-4638-5p	rs146528803	20 (/369)	1 (/22)	0 (/0)	313	0	0	0	0
hsa-miR-3622a-5p	rs66683138	6 (/231)	4 (/138)	0 (/22)	241	24	0	0	0
hsa-miR-4707-3p	rs2273626	4 (/86)	6 (/195)	0 (/110)	162	0	0	0	0
hsa-miR-1269a	rs73239138	5 (/236)	2 (/131)	0 (/24)	107	31.5 (**)	0	0	0
hsa-miR-4781-3p	rs74085143	8 (/366)	0 (/25)	0 (/0)	135	0	0	0	0
hsa-miR-6763-3p	rs3751304	1 (/39)	2 (/153)	2 (/199)	86	29	0	0	0
hsa-miR-4804-5p	rs266435	0 (/7)	3 (/91)	7 (/293)	0	114	1	0	0
hsa-miR-4999-5p	rs72996752	5 (/189)	1 (/163)	0 (/39)	86	26	0	1	0
hsa-miR-6839-5p	rs7804972	1 (/153)	2 (/196)	0 (/42)	56	0	1	0	0
hsa-miR-4459	rs73112689	3 (/245)	1 (/133)	0 (/13)	51	0	0	0	0
hsa-miR-585-3p	rs62376935	0 (/330)	2 (/52)	0 (/9)	2	48	0	0	0
hsa-miR-3928-5p	rs5997893	1 (/38)	1 (/157)	1 (/196)	47	9	0	0	0
hsa-miR-548al	rs515924	0 (/307)	1 (/80)	1 (/4)	0	37	0	0	0
hsa-miR-3117-3p	rs12402181	1 (/288)	0 (/97)	0 (/6)	35	0	0	0	0
hsa-miR-3130-3p	rs2241347	2 (/254)	3 (/119)	0 (/18)	35	0	0	0	0
hsa-miR-4532	rs73177830	2 (/344)	0 (/44)	0 (/3)	34	0	0	0	0
hsa-miR-302c-3p	rs199971565	1 (/371)	0 (/20)	0 (/0)	30	0	0	0	0
hsa-miR-4772-5p	rs62154973	3 (/339)	0 (/51)	0 (/1)	25	0	0	0	0
hsa-miR-1343-5p	rs2986407	0 (/20)	0 (/111)	2 (/260)	0	24	0	0	0
hsa-miR-548ap-5p	rs4414449	2 (/47)	11 (/180)	1 (/164)	16	4	4	0	0
hsa-miR-6885-5p	rs78293125	1 (/341)	0 (/47)	0 (/3)	15	0	0	0	0
hsa-miR-548h-3p	rs73235381	3 (/379)	0 (/11)	0 (/1)	14	0	0	0	0
hsa-miR-4433a-5p	rs12473206	1 (/230)	2 (/147)	0 (/14)	12	0	0	0	0

	TOTAL	1062	400	324	108574,5	111540,5	140	581	10
1130-111111-12090	rs7210937	1 (/326)	0 (/60)	0 (/5)	31.3()	0()	0	U	0
hsa-miR-1269b	rs12451747	0 (/82)	1 (/171)	0 (/138)	31.5 (**)	0 (*)	0	0	0
113а 11111 07 90-5р	rs3745198	3 (/131)	0 (/190)	0 (/70)	30	0()	0	0	0
hsa-miR-6796-3p	rs3745199	3 (/131)	0 (/193)	0 (/67)	56	0 (*)	0		
hsa-miR-6863	rs12708966	1 (/385)	0 (/6)	0 (/0)	1	0	0	0	0
hsa-miR-548h-5p	rs9913045	0 (/143)	0 (/197)	0 (/51)	0	0	1	0	0
hsa-miR-4520-3p	rs8078913	0 (/87)	1 (/198)	0 (/106)	1	0	0	0	0
hsa-miR-4302	rs11048315	1 (/278)	2 (/102)	0 (/11)	1	2	0	6	0
hsa-miR-6805-3p	rs56312243	1 (/337)	0 (/50)	0 (/4)	5	0	0	0	0
hsa-miR-1229-5p	rs2291418	2 (/363)	0 (/27)	0 (/1)	7	0	0	0	0
hsa-miR-4520-5p	rs8078913	0 (/87)	0 (/198)	1 (/106)	0	9	0	0	0
hsa-miR-4482-5p	rs45596840	1 (/167)	0 (/179)	0 (/45)	10	0	0	0	0
hsa-miR-6887-5p	rs1688017	1 (/152)	0 (/193)	0 (/46)	10	0	0	0	0

⁽¹⁾ Between parenthesis is the total number of samples carrying the variant with homozygous reference allele (0/0), heterozygous (0/1) or homozygous alternative allele (1/1)

Table 2: Inconsistent genotypes unlikely due to sequencing errors

polymiR	rsID	Alleles: Reference, Alternative	MAF	R ²	SAMPLE	Genotype from vcf	Counts Reference	Counts Alternative	SANGER validation
hao miD 4422h 2n	rs12473206	C,G	0.23	0.99	PVP28	C/C	84	333	G/C
hsa-miR-4433b-3p	1512473206	C,G	0.23	0.99	PVPZO	C/C	04	333	G/C
hsa-miR-4433b-3p	rs12473206	C,G	0.23	0.99	SSP20	C/C	135	8	N.A.
hsa-miR-339-3p	rs72631820	T,C	0.009	0.98	CED24	T/T	139	7	N.A.
hsa-miR-6826-5p	rs6771809	T,C	0.14	0.98	DTD19	T/T	1	20	N.A.
hsa-miR-1255b-5p	rs6841938	G,A	0.09	0.96	AJA9	G/G	92	28	G/G
hsa-miR-7107-3p	rs3817551	T,G	0.36	0.96	MSM20	G/G	27	0	N.A.
hsa-miR-7107-3p	rs3817551	T,G	0.36	0.96	MCC20	G/G	15	0	G/G
hsa-miR-7107-3p	rs3817551	T,G	0.36	0.96	NFN22	G/G	23	0	G/G
hsa-miR-7107-3p	rs3817551	T,G	0.36	0.96	NHN19	G/G	25	0	G/G
hsa-miR-7107-3p	rs3817551	T,G	0.36	0.96	GLG20	G/G	17	0	G/G

MAF: Minor Allele Frequency R²: imputation quality criteria

Figure Legends

Figure 1: Percentiles of the score difference of ambiguous alignment.

This figure represent in percentiles the distribution of the score difference between the two best alignments of each cross-mapping reads. 8.4% of cross-mapping reads have a score difference of 0, which means they aligned on different sequences with the same score, thus they cannot be disambiguated.

Figure 2: Percentiles of the alignment score distribution.

The alignment score distribution after cross-mapping reads disambiguation represented in percentiles. 98% of alignments have a score lower or equal to 9. The remaining 2 % of alignments have a score ranging from 10 to 76, which are categorized as unreliable isomiRs because of the unlikely number of editing events they would have underwent. About 42% of alignments have a score equal to 0, meaning the reads involved are identical to the reference sequence, or have templated additions.

⁽²⁾ Alignments with a high number of inconsistent reads (higher than 1% of the sum of reads gathered by the polymiR sequences), which is incompatible with sequencing errors replicating a genetic variant.

^(*) For multi variant polymiRs, reads did not mapp to any of the sequence integrating alternative haplotypes

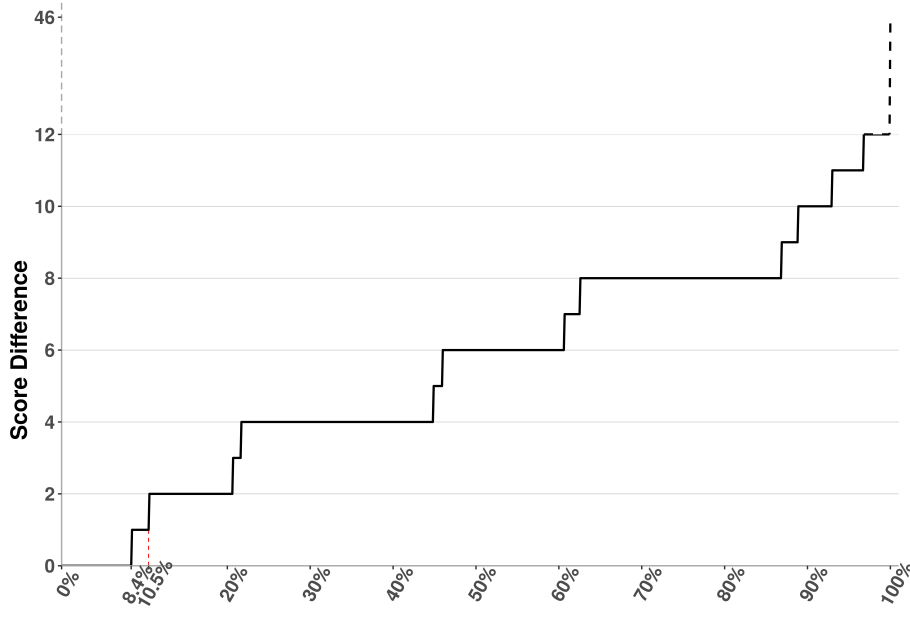
^(**) The sequence of hsa-miR-1269a integrating the rs73239138-A allele is identical to the hsa-miR-1269b sequence. Hence each of the 63 reads that cross-mapped on both sequence received a count of ½ on each sequence.

Figure 3: Percentiles of the alignment score distribution after discarding alignments with a score higher than 9.

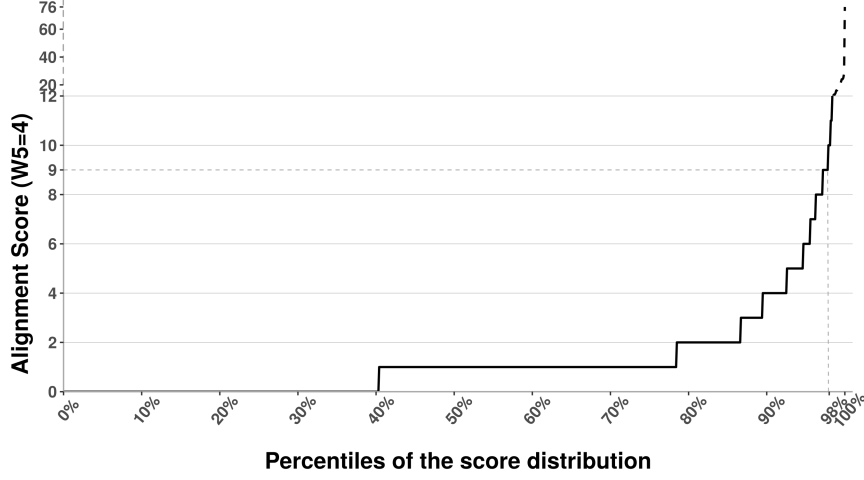
Figure 4: IsomiRsprofiling.

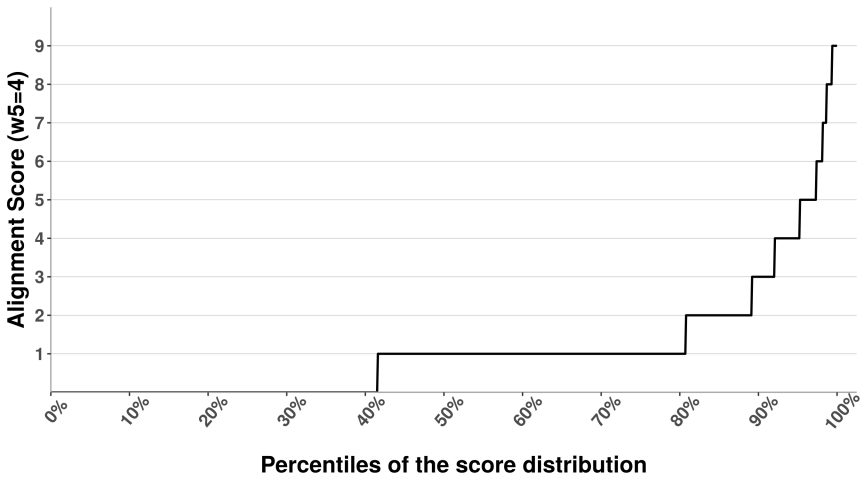
Pie charts representing the proportion of aligned reads corresponding to isomiRs. While ~60% of reads aligned have a 3' end that differs from the reference sequence, only ~6% have a different 5' end. (Canonical: No difference from the reference sequence; Trim: Trimming event; NTA: Non Templated Addition; Trim+NTA: Trimming followed by NTA; TA: Templated Addition)

Figure 5: Examples of allele imbalance expressions observed at 4 polymiRs



Distribution percentiles: Score difference of ambiguous alignments





TOTAL - Read counts = 550946055



Canonicals vs isomiRs

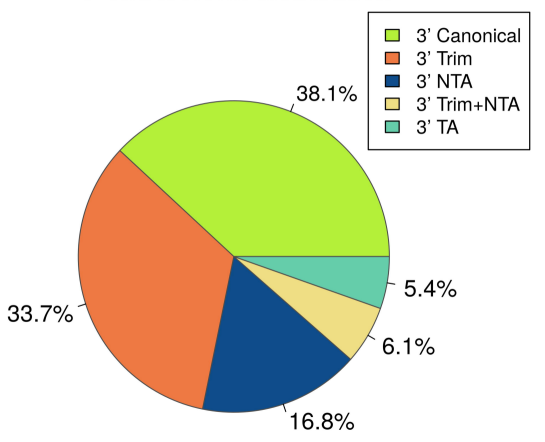
proportions

64.1%

,35.9%

Canonicals

IsomiRs



5' end isoform distribution

