Primer Scheme specifications v3.0.0-alpha

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Abstract

DNA sequencing of tiled amplicon PCR products is an important approach for fast and cost-effective pathogen genome surveillance. Accurate bioinformatic analysis of tiled PCR amplicon sequences depends on knowledge of primer sequences, amplicon layout and coordinates with respect to a reference genome. Analysis and reuse of tiled amplicon sequencing data is currently hindered by the lack of defined file formats for describing primer schemes, a problem highlighted by the proliferation of primer schemes for SARS-CoV-2 genomes during the COVID-19 pandemic. We describe a text-based specification for describing sequencing primers and amplicons with respect to one or more reference chromosomes. This specification formalises an existing widely-used primer scheme interchange format initially adopted by the PrimalScheme primer design tool, but since adopted by a growing ecosystem of community tooling. This specification designates the use of a primer bed file in Browser Extensible Data (BED) format and accompanying reference fasta sequence file in order to define a primer scheme.

Keywords Data standards, Primer Schemes, Amplicon Sequencing

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1. primer.bed file

A primer bed file describes an amplicon sequencing primer scheme in machine and human readable tabular format. Together with an accompanying reference.fasta, its purpose is to encapsulate all of the information needed to i) acquire the primers from suppliers or custom oligonucleotide synthesis, ii) combine the primers correctly to reproduce a pooled primerscheme, and iii) facilitate correct and reproducible bioinformatic analysis of resulting sequencing data. It therefore incorporates both wet lab and analytical elements. This information includes primer sequences, primer pools, coordinates and orientation with respect to a reference sequence, and optionally relative primer concentrations.

1.1. Format overview

primer.bed files are tab-delimited ASCII text files. Each line can either represent a comment line (prefixed with #) or a record line (BedLine), representing a single unique oligonucleotide primer or probe associated with an amplicon. An amplicon comprises at least two primer record lines each describing primers on different strands. A compliant primer bed file contains one or more amplicons.

The format of primer. bed is based on Browser Extensible Data (BED) specification, with each oligonucleotide being treated as a genomic region, enabling compatibility with common BED file tooling.

1.2. Comment Line

Comment lines are minimally parsed, but can optionally contain a scheme-level (key, value) pair. To this end, comment lines containing a single "=" will be split, with the left and right sides representing a scheme-level key and value respectively.

1.3. record line (BedLine) field descriptions

Column	Field name	Туре	Brief description	Restrictions
1	chrom	String	Chromosome name	[A-Za-z0-9_]
2	primerStart	Integer	Primer start position (zero-based, half-open)	Positive integer (u64)
3	primerEnd	Integer	Primer end position (zero-based, half-open)	Positive integer (u64)
4	primerName	String	Primer name	[a-zA- 0-9\-]+_[0-9]+_(LEFT RIGHT PROBE)_[0-9]+
5	pool	Integer	Primer pool	Positive integer (u64)
6	strand	String	Primer strand	[-+]
7	primerSeq	String	The nucleotide sequence in $5' \rightarrow 3'$	ASCII non-whitespace characters
8	primerAttributes	Optional(String)	List of record-level (key, value) pairs separated by `;`. e.g. k1=v1; k2=v2	ASCII non-whitespace characters

Table 1: The column structure and description of a BedLine

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1.3.1. chrom

The name of the corresponding reference sequence chromosome for the primer. This must match a valid sequence ID inside an accompanying reference sequence FASTA file, by convention named reference.fasta.

1.3.2. primerStart

The start position of the primer on the chrom using BED-like zero-based, half-open coordinates.

1.3.3. primerEnd

The non-inclusive end position of the primer on the chrom using BED-like zero-based, half-open coordinates. Must be greater than primerStart.

1.3.4. primerName

The name of the primer in the form "{prefix}_{ampliconNumber}_{direction}_{primerNumber}".

- prefix: Must match regex [a-zA-Z0-9\-]. See best practices
- ampliconNumber: The number of the amplicon for its relevant chrom. Must be a positive integer incrementing from 1.
- direction: The direction of the primer. Must be either LEFT, RIGHT or PROBE.
- primerNumber: The number of the primer. Must be a positive integer incrementing from 1.

1.3.5. pool

The PCR pool the primer belongs to. Must be a positive integer incrementing from 1¹.

1.3.6. strand

The strand of the primer must be either + or -. It must correspond to the direction component of the primerName (see the description of primerName above). LEFT and RIGHT primers must be + and respectively, while PROBE can be either.

1.3.7. primerSeq

The sequence of the primer in the 5' to 3' direction. Unrestricted to contain any non-whitespace ASCII character².

1.3.8. primerAttributes

An **optional** list of a (key, value) pairs used to denote additional arbitrary primer attributes, in the form of pw=1.0; ps=10.0. This is intentionally flexible to allow the storage of additional information. In a primer bed file this can be represented as either an empty 8th column or only 7 columns.

1.3.8.1. Reserved keys

• pw: primerWeight. The concentration of individual primers can be altered to balance amplicon performance. Primer concentration in the PCR should be scaled by primerWeight * [typical PCR conc]. This is restricted to positive floating point numbers (f64 > 0).

1.4. Examples

1.4.1. Simple example

A seven column primer.bed file, with no primerAttributes or comment lines.

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[&]quot;Existing schemes/literature use refer to 'pool 1 and pool 2'. Therefore 1-based indexing is expected"

² This is intentionally unrestricted (rather than IUPAC-only) to allow Primer Modification. Such as /56-FAM/ {primerSeq} to represent 5' 6-FAM fluorescent dye labelled probe"

```
MN908947.3 100 131 example_1_LEFT_1 1 + CTCTTGTAGATCTGTTCTCTAAACGAACTTT
MN908947.3 419 447 example_1_RIGHT_1 1 - AAAACGCCTTTTTCAACTTCTACTAAGC
{\tt MN908947.3} \quad {\tt 344\ 366\ example\_2\_LEFT\_1} \quad {\tt 2\ +\ TCGTACGTGGCTTTGGAGACTC}
MN908947.3 707 732 example 2 RIGHT 1 2 - TCTTCATAAGGATCAGTGCCAAGCT
```

1.4.2. Complex example

An eight column primer.bed file. With primerAttributes defined, and comment lines providing a chrom alias and explaining the gc primerAttributes.

```
# example scheme
# gc=fraction gc
# MN908947.3=sars-cov-2
MN908947.3 100 131 example_1_LEFT_1 1 + CTCTTGTAGATCTGTTCTCTAAACGAACTTT pw=1.4;gc=0.35
MN908947.3 419 447 example_1_RIGHT_1 1 - AAAACGCCTTTTTCAACTTCTAAGC pw=1.4;gc=0.36
MN908947.3 344 366 example_2_LEFT_1 2 + TCGTACGTGGCTTTGGAGACTC pw=1;gc=0.55
MN908947.3 707 732 example 2 RIGHT 1 2 - TCTTCATAAGGATCAGTGCCAAGCT pw=1;qc=0.44
```

1.4.3. qPCR example

An eight column primer. bed file. Showing a fictional qPCR assay. The specific dyes and quenchers are (optionally) included in the comments lines.

```
# example multiplexed-qPCR assay
# gc=fraction gc
# /3BHQ 1/=Black Hole Quencher 1
# /56-FAM/=FAM
# /5HEX/=HEX
target1 \quad 2010 \ 2030 \ iad3\_1\_LEFT\_1 \quad 1 \ + \ AAAGGTCAGTCAACCCGTTC \ pw=1
 target1 \quad 2035 \quad 2060 \quad iad3\_1\_PROBE\_1 \quad 1 \quad - \quad /56-FAM/GCGTTGTTCAATTGCCTTGCTGATT/3BHQ\_1/ \quad pw=19.1 \quad 2035 \quad 2060 \quad iad3\_1\_PROBE\_1 \quad 1 \quad - \quad /56-FAM/GCGTTGTTCAATTGCCTTGCTGATT/3BHQ\_1/ \quad pw=19.1 \quad 2035 \quad 2060 \quad 
 target1 2903 2923 iad3_1_RIGHT_1 1 - TCGGGCCACCGCGTATGAAG pw=1
target2 5167 5187 rfw1_1_LEFT_1 1 + TCGTAGCATGGACTCGATGA pw=1
target2 5271 5296 rfw1_1_PROBE_1 1 + /5HEX/TGATCCGCGTTTACTGTTCGACGCG/3BHQ_1/ pw=20.2
 target2 5301 5321 rfw1_1_RIGHT_1 1 - GTTTACCAAGGAACCATCCA pw=1
```

1.5. Best Practices

primer.bed contain information about how to replicate the primer pools used in multiplexed PCR. They do not contain information about the PCR protocol, input material, or sequencing method and analysis. Therefore, additional information is needed for true reproducibility.

1.5.1. Other metadata standards

To explicitly differentiate different versions of primer.bed, this spec is designed to fit into larger metadata standards, such as primal-page with PrimalScheme Labs or primaschema with pha4ge <u>primer-schemes</u>

1.5.2. Other tooling

primalbedtools is a python package that carries out schema validation and conversion, and common operations on primer.bed files.

1.5.3. primerName: prefix

The primerName: prefix should be as unique as possible (for example a short uuid, 359ba5) and different for each chrom and each scheme generation run.

• Using primerName: prefix like scheme or sars-cov-2 might seen easier, however, will result in a freezer / LIMS full of similar names leading to pooling mistakes.

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1.5.4. comment line

The comment line's key=value pattern undergoes limited validation, and therefore should be not be critical for tooling. A suitable use case might be to document custom primerAttributes. Another use is providing aliases for different chrom.

2. reference.fasta file

A reference, fasta file contains the DNA sequences of all the primary-reference genomes, used in primer scheme generation. Its purpose is to provide a reference genome and coordinate system for use in reference-based assembly and consensus generation.

2.1. Format overview

reference. fasta files are typical . fasta format files, with text representing the nucleotide sequence of the reference. Each genome starts with a header line (starting with >) that denotes the id of the genome, followed by lines of nucleotide data.

All chrom fields of the record lines must have a corresponding id in the reference.fasta.

2.2. Examples

2.2.1. Single fasta

```
>MN908947.3
ATTAAAGGTTTATACCTTCCCA...
```

The corresponding primer. bed file should contain the chrom MN908947.3

2.2.2. Multi fasta

```
>MN908947.3
ATTAAAGGTTTATACCTTCCCA...
>NC 006432.1
CGGACACACAAAAAGAAAGAAA...
```

The corresponding primer.bed file should contain the chrom MN908947.3 and NC_006432.1

2.3. Best practices

As the reference. fasta is often used for referenced-based assembly, using high quality genome with few Ns or ambiguous bases is advisable. Using RNA sequences in the reference.fasta is not recommended, as DNA is expected.

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