The Browser Extensible Data (BED) format

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1 Specification

Browser Extensible Data (BED) is a whitespace-delimited file format, where each **file** consists of one or more **lines**.¹ Each **line** describes discrete genomic **features** by physical start and end position on a linear **chromosome**. The file extension for the BED format is .bed.

1.1 Typographic conventions

This document uses the following typographic conventions:

Style	Meaning	Examples	
Bold	Terms defined in subsections 1.2–1.3	chromosome file	
Sans serif	Names of field s	chrom chromStart chromEnd	
Fixed-width	Literals or regexes ²	.bed grep [[:alnum:]]+ ATCG	

1.2 Terminology and concepts

0-start, half-open coordinate system: A coordinate system where the first base starts at position 0, and the start of the interval is included but the end is not. For example, for a sequence of bases ACTGCG, the bases given by the interval [2, 4) are TG.

BEDn: A file with the first n fields of the BED format. For example, **BED3** means a file with only the first 3 fields; **BED12** means a file with all 12 fields.

BEDn+: A file that has n fields of the BED format, followed by any number of fields of custom data defined by a user.

BEDn+m: A file that has a custom tab-delimited format starting with the first n fields of the BED format, followed by m fields of custom data defined by a user. For example, **BED6**+4 means a file with the first 6 fields of the BED format, followed by 4 user-defined fields.

block: Linear subfeatures within a feature. Usually used to designate exons.

[&]quot;Frequently Asked Questions: Data File Formats." University of California, Santa Cruz (UCSC) Genome Browser FAQ, https://genome.ucsc.edu/FAQ/FAQformat.html

POSIX/IEEE 1003.1–2017 Extended Regular Expressions, for the "C" locale. IEEE Standard for Information Technology—Portable Operating System Interface (POSIX) Base Specifications, IEEE 1003.1–2017, 2017

chromosome: A sequence of nucleobases with a name. In this specification, "chromosome" may also describe a named scaffold that does not fit the biological definition of a chromosome. Often, chromosomes are numbered starting from 1. There are also often sex chromosomes such as W, X, Y, and Z, mitochondrial chromosomes such as M, and possibly scaffolds from an unknown chromosome, often labeled Un. The name of each chromosome is often prefixed with chr. Examples of chromosome names include chr1, 21, chrX, chrM, chrUn, chr19_KI270914v1_alt, and chrUn_KI270435v1.

feature: A linear region of a **chromosome** with specified properties. For example, a **file**'s **feature**s might all be peaks called from ChIP-seq data, or transcript.

field: Data stored as non-tab text. All fields are 7-bit US ASCII.

file: Sequence of one or more **lines**.

line: String terminated by a line separator, in one of the following classes. Either a data line, a comment line, or a blank line. Discussed more fully in subsection 1.3

line separator: Either carriage return, line feed, or carriage return followed by line feed.

1.3 Lines

1.3.1 Data lines

Data lines contain **feature** information. A data line is composed of **field**s separated by whitespace. The whitespace must match the regex [[:space:]]+³.

1.3.2 Comment lines and blank lines

Both comment lines and blank lines provide no feature data.

Comment lines start with # with no whitespace beforehand. A # appearing anywhere else in a line is treated as **feature** data, not a comment.

Blank lines consist entirely of whitespace. Both comment and blank lines may appear as any line in a file, at the beginning, middle, or end of the file. They may appear in any quantity.

1.4 BED fields

Each **data line** contains between 3 and 12 whitespace-delimited **fields**. The first 3 **fields** are mandatory, and the last 9 **fields** are optional. In optional **fields**, the order is binding—if 1 **field** is filled, then all previous **fields** must also be filled. However, **BED10** is not allowed.⁴

In a BED file, each data line must have the same number of fields. The positions in BED fields are all described in the **0-based**, half-open coordinate system.

³ [[:space:]] includes the following characters: space, form-feed, newline, carriage-return, tab, and vertical-tab

⁴ Knowing only the number of **block**s has almost no use cases. **BED11** is allowed, however, because there are use cases for having blocks with known starting positions but unspecified ends.

Col	Field	Type	Regex or range	Brief description
1	chrom	String	[[:alnum:]_]{1,255} ⁵	Chromosome name
2	${\sf chromStart}$	Int	$[0, 2^{32} - 1]$	Feature start position
3	${\sf chromEnd}$	Int	$[0, 2^{32} - 1]$	Feature end position
4	name	String	[^\t]{0,255}	Feature description
5	score	Int	[0, 1000]	A numerical value
6	strand	String	[-+.]	Feature strand
7	thickStart	Int	$[0, 2^{32} - 1]$	Thick start position
8	thickEnd	Int	$[0, 2^{32} - 1]$	Thick end position
9	itemRgb	$_{\mathrm{Int,Int,Int}}$	$([0, 255], [0, 255], [0, 255]) \mid 0$	Display color
10	blockCount	Int	$[0,chromEnd-chromStart]^6$	Number of block s
11	blockSizes	$\operatorname{List}[\operatorname{Int}]$	([[:digit:]]+,){blockCount-1}[[:digit:]]+,? ⁷	Block sizes
12	${\sf blockStarts}$	List[Int]	([[:digit:]]+,){blockCount-1}[[:digit:]]+,?	Block start positions

1.5 Coordinates

- 1. **chrom**: The name of the **chromosome** or scaffold where the **feature** is present. Limiting only to word characters only, instead of all non-whitespace characters, makes BED **files** more portable to varying environments which may make different assumptions about allowed characters. The name must be between 1 and 255 characters long, inclusive.
- 2. chromStart: Start position of the **feature** on the **chromosome** or scaffold. **chromStart** must be an integer greater than or equal to 0 and less than the total number of bases of the **chromosome** to which it belongs. If the size of the **chromosome** is unknown, then **chromStart** must be less than or equal to $2^{32} 1$, which is the maximum size of an unsigned 32-bit integer.
- 3. chromEnd: End position of the **feature** on the **chromosome** or scaffold. chromEnd must be an integer greater than or equal to the value of **chromStart** and less than or equal to the total number of bases in the **chromosome** to which it belongs. If the size of the **chromosome** is unknown, then **chromEnd** must be less than or equal to $2^{32} 1$, the maximum size of an unsigned 32-bit integer.

1.6 Simple attributes

- 4. name: String that describes the **feature**. The name must be 0 to 255 non-tab characters. The name must not be empty or contain whitespace, unless all fields in file are delimited exclusively using single tab characters. A visual representation of the BED format may display the name next to the **feature**.
- 5. score: Integer between 0 and 1000, inclusive. If the **feature** has no score information, then 0 should be used as a default value. A visual representation of the BED format may shade features differently depending on their score.
- 6. strand: Strand that the **feature** appears on. The strand may either refer to the + (sense or coding) strand or the (antisense or complementary) strand. If the **feature** has no strand information or unknown strand, then a dot (.) must be used.

⁵ [[:alnum:]_] is equivalent to the regex [A-Za-z0-9_]. It is also equivalent to the Perl extension [[:word:]].

⁶ chromEnd-chromStart is the maximum number of **block**s that may exist without overlaps.

For example, if blockCount = 4, then the allowed regex would be ([[:digit:]]+,){3}[[:digit:]]+,?

1.7 Display attributes

- 7. thickStart: Start position at which the **feature** is visualized with a thicker or accented display. This value must be an integer between chromStart and chromEnd, inclusive. There is no specified default value for thickStart.
- 8. thickEnd: End position at which the **feature** is visualized with a thicker or accented display. This value must be an integer greater than or equal to thickStart and less than or equal to chromEnd, inclusive. In BED files with fewer than 7 fields, the whole **feature** has thick display. In BED7+ files, to achieve the same effect, set thickStart equal to chromStart and thickEnd equal to chromEnd. If this field is not specified but thickStart is, then the entire **feature** has thick display. There is no specified default value for thickEnd.
- 9. itemRgb: A triple of integers that determines the color of this **feature** when visualized. The triple is three integers separated by commas. Each integer is between 0 and 255, inclusive. To make a **feature** black, itemRgb should be a single 0 rather than a triplet.

1.8 Blocks

- 10. blockCount: Number of blocks in the feature. blockCount must be an integer greater than 0. blockCount is mandatory in BED10+ files. Null or empty blockCount are not allowed, because blockSizes and blockStarts rely on blockCount. A visual representation of the BED format may have blocks appear thicker than the rest of the feature.
- 11. blockSizes: Comma-separated list of length blockCount containing the size of each block. There must be no spaces before or after commas. There may be a trailing comma after the last element of the list. blockSizes is mandatory in BED11+ files. Null or empty blockSizes is not allowed, because blockStarts cannot be verified without blockSizes.
- 12. blockStarts: Comma-separated list of length blockCount containing each block's start position, relative to chromStart. There must not be spaces before or after the commas. There may be a trailing comma after the last element of the list. Each element in blockStarts is paired with the corresponding element in blockSizes. Each blockStarts element must be an integer between 0 and chromEnd chromStart, inclusive. For each couple *i* of (blockStarts_i, blockSizes_i), the quantity chromStart + blockStarts_i + blockSizes_i must be less or equal to chromEnd. These conditions enforce that each block is contained within the feature. The first block must start at chromStart and the last block must end at chromEnd. Moreover, the blocks must not overlap. The list must be sorted in ascending order. blockStarts is mandatory in BED12 files. Null or empty blockStarts is not allowed.

1.9 User-defined fields

In custom BED files with user-defined fields, each field must be a single value or a list of values. A list of values must be comma-separated. Each field's type must be one of Integer, Flag, Float, Character, or String.

Each type is defined as:

Type	Definition	
Integer	32-bit signed integer	
Float	32-bit floating point defined by IEEE-754-1985 standard	
Flag	0 or 1, representing False or True	
Character	A single ASCII character	
String	One or more ASCII characters	

2 Examples

2.1 Example BED6 file from the UCSC Genome Browser FAQ⁸

```
chr7
      127471196
                 127472363
                             Pos1
chr7
      127472363
                 127473530
                             Pos2
      127473530
                 127474697
                             Pos3
chr7
      127474697
chr7
                 127475864
                             Pos4
                                   0
                             Neg1
chr7
      127475864
                 127477031
                                   0
      127477031
chr7
                 127478198
                             Neg2
                                   0
chr7
      127478198
                 127479365
                             Neg3
                                   0
      127479365
                 127480532
                                    0
chr7
                             Pos5
chr7
      127480532
                 127481699
                             Neg4
```

2.2 Example BED12 file from the UCSC Genome Browser FAQ

```
chr22 1000 5000 cloneA 960 + 1000 5000 0 2 567,488, 0,3512 chr22 2000 6000 cloneB 900 - 2000 6000 0 2 433,399, 0,3601
```

The **block**s in this example satisfy the required constraints. The first **block** starts at **chromStart** since the first **blockStarts** element is 0. The last **block** ends at **chromEnd** since the last **block** starts at 3512 with size 488, and 3512 + 488 = chromEnd - chromStart.

3 Recommended practice for the BED format

3.1 Mandatory fields

• chrom: The name of each **chromosome** should also match the names from a reference genome, if applicable. For example, in the human genome, the chromosomes may be named **chr1** to **chr22**, **chrX**, **chrY**, and **chrM**. Names should be consistent within a **file**. For example, one should not use both 17 and **chr17** to represent the same **chromosome** in the same **file**.

3.2 Optional fields

• name: If a feature has no name, then a dot (.) should be used. Names should avoid using the space character even if the file is exclusively delimited with single tab characters because parsers may interpret a space as a delimiter.

^{8 &}quot;Frequently Asked Questions: Data File Formats." UCSC Genome Browser FAQ, https://genome.ucsc.edu/FAQ/FAQformat.html

- itemRgb: Eight or fewer colors should be used as too many colors may slow down visualizations and are difficult for humans to distinguish.⁹
- blockSizes and blockStarts: The length of the list of blocks should equal to blockCount. If either of these lists are longer, then their trailing items are ignored.

3.3 Sorting

BED files should be sorted by chrom, then by chromStart numerically, and finally by chromEnd numerically. chrom may be sorted using any scheme (such as lexicographic or numeric order), but all lines with the same chrom value should occur consecutively. For example, the lexicographic order of chr1, chr10, chr11, chr12, ..., chr2, chr20, chr21, ..., chr3, ..., chrX, chrY, chrM is an acceptable sorting. The numeric order of chr1, chr2, ..., chr21, chr22, chrM, chrX, chrY is also acceptable. Regardless of the chromosome sorting scheme, lines for two features on the same chromosome should not have any lines for features on other chromosomes between them.

3.4 Whitespace

Though lines may use any kind of whitespace as a delimiter between **fields**, a single tab (\t) should be used. This is because almost all tools support tabs while some tools do not support other kinds of whitespace. Also, whitespace within the name field may be used only if the **field** delimiter is tab throughout the **file**.

3.5 Large BED files

If a **file** intended for visualization is over 50 MiB in size, the **file** should be converted to **bigBed** format, which is an indexed binary format. ¹⁰ The **bedToBigBed** program may perform this conversion. ¹¹

4 UCSC track files

Track files are files that contain additional information intended for a visualization tool such as the UCSC Genome Browser. Track files contain browser lines and track lines that precede lines from a file format supported by the Genome Browser. Track files are not valid BED files — valid BED files must not have any browser or track lines. To distinguish between BED files and track files, track files should use the file extension .track.

5 Acronyms

ASCII American Standard Code for Information Interchange

BED Browser Extensible Data

⁹ "Frequently Asked Questions: Data File Formats." UCSC Genome Browser FAQ, https://genome.ucsc.edu/FAQ/FAQformat.html

¹⁰ Kent, W. James et al. (2010) "BigWig and BigBed: enabling browsing of large distributed datasets." *Bioinformatics* 26(17):2204–2207. https://doi.org/10.1093/bioinformatics/btq351

^{11 &}quot;bigBed Track Format." UCSC Genome Browser FAQ, https://genome.ucsc.edu/goldenPath/help/bigBed.html

¹² Haeussler, Maximilian et al. (2019) "The University of California, Santa Cruz Genome Browser database: 2019 update." Nucleic Acids Research 47(D1):D853-D858. https://doi.org/10.1093/nar/gky1095

[&]quot;Displaying your own annotations in the Genome Browser." UCSC Genome Browser FAQ, https://genome.ucsc.edu/goldenPath/help/customTrack.html#lines

GA4GH Global Alliance for Genomics and Health

regex regular expression

UCSC University of California, Santa Cruz

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