Use of *color_seq.pl, color_nums.pl, and color_cols.pl* scripts

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Background and overview

The three scripts *color_seq.pl*, *color_nums.pl*, *and color_cols.pl* are simple command line filters intended to colorize specific value-associated "words" contained within simple ASCII text. This is useful to highlight specific values for quick value identification and pattern visualization within a terminal. The tools take text from a file or standard input then output contents with ANSI escape sequences interjected to color specific characters in a terminal display. The tools also accept various command line options to allow user control over character or token selection, color choice, and other data-dependent options.

This document briefly describes **options** available for these tools and, probably more importantly, provides **example uses**. Given that the point of these tools is to colorize text, examples are accompanied by screen captures illustrating outputs. This documentation is in pdf format to allow these graphics.

The README file, in markdown format, breifely describes the software (mainly dependencies).

color_seq.pl is for coloring DNA sequences. The default behavior is to color what appears to be sequences with a four-color scheme associated with A C G and T. The script contains logic to guess whether a "word" (or token; continuous block of characters) is sequence-containing or not based on the fraction of ACGT characters; Words deemed to *not* be sequence-containing are not colored. Within sequence words, it is possible to limit the range of characters that are colored, with the range being relative to either the beginning or end of the word. By default, entire lines beginning with "#" or ">" are not colored, as these traditionally are associated with comments or the header lines of *fasta* format files. This behavior can be changed via command line switch.

In addition to simply finding and coloring DNA sequences, *color_seq.pl* can be used to highlight attributes often encountered in sequence files. For example, lower case sequence can be ignored (i.e. not colored), or non-ACGT characters within (what appears to be) sequence can be highlighted. This is useful to quickly find IUB characters and any non-sequence characters (both colored differently to stand out).

Another behavior is to color continuous runs of like bases. This yields sparsely colored patterns that aids visual identification of specific sequences / subsequences within larger sets. The length of runs may be specified and the coloring scheme may be inverted (i.e. only non-continuous runs are colored).

Finally, *color_seq.pl* can be used to highlight *window-based* sequence features. With this feature runs of contiguous characters containing, as well as not containing, a specific base are highlighted. This feature supports IUB degenerate codes in addition to the normal ACGT bases. For example, using window-based highlighting on can readily identify runs of GC (IUB code "S"), runs of purines (IUB code "R"), etc. In addition to highlighting windows containing the specified base (or bases corresponding to an IUB code), windows *not* containing runs of the chosen base(s) are also highlighted in a second color.

color_nums.pl is for coloring numbers. The default behavior is to highlight numbers within text using a single color (This is distinct from simply coloring numeral characters). By default, entire lines beginning with "#" are not colored, as these traditionally are associated with comments, but this behavior can be changed via command line switch. It is also possible to only color a subset of number-containing "columns" via command line switch.

Assigning colors based on value allows rapid identification of "high" or "low" values. In addition, coloring a whole table of numbers together can serve as s simple means to visualize patterns in the data.

color_cols.pl is for coloring columns (or rows) so that different columns (e.g. "fields") in a

data table may be readily identified. The default behavior is to cycle through a set of colors for odd number columns and leave even number columns white, such that the cycle repeats every ten columns. Other schemes and cycling frequencies are available (e.g. every 5 or 2 columns), a range of columns to color may be specified, and specific columns may be selected for coloring. By default, entire lines beginning with "#" are not colored, as these traditionally are associated with comments, but this behavior can be changed via command line switch. It is also possible to specify that columns are separated by tabs rather than the default spaces.

Use of *color_seq.pl*

Calling *color_seq.pl* without any arguments (or with "-help" or an unrecognized argument) yields this splash / help screen:

Each command line switch has a brief description of what it does. Given only input and no command switches, default coloring is performed. Figure S1 shows a test-case sequence file displayed via the *cat* command. Figure S2 shows this same content with output from *cat* piped through *color_seq.pl*, which is given a "naked dash" to indicate input is from standard input (*stdin*). Other color schemes may be specified with the <code>-cabi, -cigv</code> and <code>-cgc</code> switches; <code>-corig</code> is for the default scheme. Results of coloring with different schemes is shown in Figure S3, with three separate calls illustrating IGV, ABI and GC-based coloring schemes (top to bottom).

Figure S1. Test-case sequence file.

```
ryan@verdascend ~/git/comline_color_tools> cat examp.fas | color_seq.pl -
   Comment line for seqs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGCCTGGCTAGGGAAAGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
ryan@verdascend ~/git/comline_color_tools>
ryan@verdascend ~/git/comline_color_tools> cat examp.fas | color_seq.pl - -corg
    Comment line for seqs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGGCCTGGCTAGGGAAAGGGGGGGGGCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC ryan@verdascend ~/git/comline_color_tools> []
```

Figure S2. *color_seq.pl* Default coloring. Test-case sequence 'piped' as input. The command switch '-' is used to indicate input will come from *stdin*. Two sequential calls are shown; the second one with -corg to explicitly request default color scheme.

```
yan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -cigv
   Comment line for seqs ACGT...
>seq_1
TCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
target GGCGGAAATTCTACCCAAGACTCCC
FGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGGCCTGGCTAGGGAAAGGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
ryan@verdascend ~/git/comline_color_tools>
ryan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -cabi
   Comment line for seqs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
ryan@verdascend ~/git/comline_color_tools>
ryan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -cgc
# Comment line for seqs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGG
GTGTGAGTGGAGGGGGCCTGGCTAGGGAAAGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
```

Figure S3. *color_seq.pl* IGV, ABI and GC-based coloring schemes.

Some options can be combined, such as the -1w and -all switches, which, respectively, direct the script to color lowercase characters white (i.e. do not color them) and to include all lines (i.e. do not ignore lines starting with ">" or "#"). The result is shown in Figure S4, which uses the default color scheme. Note that the "ACGT" in the comment on the first line is colored, while the lowercase sequence characters on the third line (and further down) are not. Highlighting non-ACGT within sequence-like strings is showin in Figure S5 with the -nacgt

switch. Normal ACGT bases are not colored, IUB code letters are colored red, and other (non-DNA) characters are colored cyan.

Figure S4. *color_seq.pl* Changing which characters are considered for coloring. The <code>-lw</code> and <code>-all</code> switches result in no color for lower case characters, and all lines being "colorable", respectively.

Figure S5. *color_seq.pl* Highlighting non-ACGT characters within sequence-containing strings.

Highlighting *window-based* sequence features is shown in Figure S6. Contiguous runs of specified base characters, and runs *devoid* of these, are colored within sequence-containing strings, while the remaining sequence characters are not colored. Two parameters define windowing behavior: base type and window size. Base type may be ACGT or any IUB code. Window size dictates the minimal length of runs that qualify for coloring. Runs of selected bases (and non-selected bases) that are at least window size long are colored; Selected base

type(s) are red, and non-selected base type(s) are cyan.

NOTE. Only ACGT bases qualify for selection and lines are considered independently. This means that IUB characters within strings do not count for runs, and runs spanning multiple lines are not colored.... (Maybe in a future version???)

```
yan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -win r
    Comment line for seqs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16 24730001
    >dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGGCCTGGCTAGGGAAAGGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
ryan@verdascend ~/git/comline_color_tools>
ryan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -win r -ws 3
    Comment line for segs ACGT...
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
       >dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGCCTGGCTAGGGAAAGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
```

Figure S6. *color_seq.pl* Coloring windows with like-base runs. Red denotes selected base type(s) and cyan denotes *absence* of selected base type. Top example highlights runs of purine bases (A or G) in red, specified with IUB code "R", using default window size (= 5). Bottom example highlights runs of C, with window size given as 3.

Coloring *runs of continuous bases* is shown in Figure S7. The top shows default behavior and the middle shows the inverted case (i.e. all bases *except* runs are colored) with the default run size of 3 bases. The bottom shows runs with size set to 4.

```
ryan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -run
   Comment line for segs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16 24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
<u>ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTTGGTAATAGTGCAGCCCACCC</u>TCTGGTGTGGGGC
GTGTGAGTGGAGGGGGCCTGGCTAGGGAAAGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
ryan@verdascend ~/git/comline_color_tools>
ryan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -run -not
   Comment line for seqs ACGT...
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16_24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCATTTTGCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGCCTGGCTAGGGAAAGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
ryan@verdascend ~/git/comline_color_tools>
ryan@verdascend ~/git/comline_color_tools> color_seq.pl examp.fas -run -rs 4
   Comment line for seqs ACGT...
>seq_1
TTCAAACCCTCATCCCTgtttagggCCGGCAGCGAGGCCTCACCACAGGGACAAGCCCGGCCCATCCCCATC
>target GGCGGAAATTCTACCCAAGACTCCC
TGTGGCGGAAATTCTACCCAAGACTCCCCAAATAATTGGTCACCTGCTCTCCTGGCCCCT
>chr16 24730001
>dirty; has non ACGT and non IUB codes but within a seq block
TCAACAAATAATAGTGGGCNNNNNNNNNNNkkNNxyztattatagcaataaccRYSTMKat
>two
ATCCAATTGTTWTATTAGCGAAATCATCATGSGTTTCA<mark>TTTT</mark>GCTTTGGTAATAGTGCAGCCCACCCTCTGGTGTGGGGC
GTGTGAGTGGAGGGGGCCTGGCTAGGGAAAGGGGGGGGGCCCCCACACTGGGCCCTAAGGTTCAAGGGCTTC
```

Figure S7. *color_seq.pl* Coloring runs of like-bases. Top shows default coloring, middle shows the inverted case (i.e. bases in runs are *not* colored), bottom shows runs with (minimum) size set to 4.

Use of *color_num.pl*

Calling *color_num.pl* without any arguments (or with "-help" or an unrecognized argument) yields this splash / help screen:

Each command line switch has a brief description of what it does. Given only input and no command switches, default coloring is performed. Figure N1 shows the first five lines of three number-containing files displayed via the *head* command. Figure N2 shows this same content with output from *head* piped through *color_num.pl*, which is given a "naked dash" to indicate input is from standard input.

Specific number values can be highlighted. For example, the -rg switch can be used to specify a range of values; Numbers within the range are shown with one color, numbers less than the range are shown in another color, and numbers more than the range are shown in still another color. It is also possible to highlight *only* the within-range values using the -ok switch. Figure N3 shows this behavior, using a chosen range of 69 to 71. There are also options to differentiate only two classes of numbers; Those with values greater than or less than some chosen threshold and the rest.

```
verdascend [~/git/comline_color_seq_num] 193> head -5 *mat
==> test-dat.mat <==
Table with four cols of average sequence values
dp_10
        70.290
                76.299
                        70.290
                                 76.299
dp_20
        69.500
                64.213
                        69.500
                                 64.213
        72.403
                66.563 72.403
                                 66.563
dp_22
==> test-lod.mat <==
Matrix with log odd frquencies for various dinucleotides
RowNames
                AA
                         AC
                                 AG
                                         ΑT
                                                          cc
                                                                  CG
                                                                           CT
                                                                                   GΑ
                                                                                           \mathsf{GC}
p0
        -0.79
                 -0.31
                         -0.65
                                  0.10
                                          0.52
                                                  -0.28
                                                           0.00
                                                                   0.33
                                                                           0.41
                                                                                   -0.71
p1
                         -0.25
                                          0.39
                                                          -0.21
                                                                           -0.43
                                                                                   -0.44
        -0.47
                 0.06
                                  0.58
                                                  -0.89
                                                                  -0.01
p2
         0.58
                                                                  -0.03
                                                                           -0.26
                 -0.53
                         -1.51
                                  0.93
                                          0.98
                                                  -0.59
                                                          -0.75
                                                                                   -1.76
==> test-mix.mat <==
Names len c2CC.3p5.10 c2CY.3p5.10 tmPey0.3p1.10 c2CC.3p3.10 c2CY.3p3.10 win4.1C.3p1.10 win5.1C.
3p1.10 win6.1C.3p1.10 win6.1C.3p3.12
CV10006790 18 0 0 26.002 0 0 1 1 1 1
CV10007783 14 0 0 28.779 0 0 0 0 0 1
CV10007820 16 0 0 25.182 0 0 0 0 0 0
CV10007826 16 0 2 30.266 0 2 2 2 2 3
```

Figure N1. Test-case number-containing files (first five lines for each of three).

```
verdascend [~/git/comline_color_seq_num] 194> head -5 *mat | color_nums.pl -
==> test-dat.mat <==
Table with four cols of average sequence values
dp_10
        70.290
                76.299
                        70.290
                                 76.299
dp_20
        69.500
                64.213
                        69.500
                                 64.213
dp_22
        72.403
                66.563
                        72.403
                                66.563
==> test-lod.mat <==
Matrix with log odd frquencies for various dinucleotides
RowNames
                AA
                        AC
                                 AG
                                                 CA
                                                                                           GC
                                         ΑT
                                                          cc
                                                                  CG
                                                                          CT
                                                                                   GΑ
                                                  -0.28
                        -0.65
                                          0.52
                                                                                   -0.71
        -0.79
                -0.31
                                                          0.00
                                                                   0.33
                                                                           0.41
p0
                                  0.10
p1
        -0.47
                 0.06
                         -0.25
                                          0.39
                                                  -0.89
                                                          -0.21
                                                                  -0.01
                                                                          -0.43
                                                                                   -0.44
                                  0.58
         0.58
                -0.53
                         -1.51
                                  0.93
                                                  -0.59
                                                                                   -1.76
p2
                                          0.98
                                                          -0.75
                                                                  -0.03
                                                                          -0.26
==> test-mix.mat <==
Names len c2CC.3p5.10 c2CY.3p5.10 tmPey0.3p1.10 c2CC.3p3.10 c2CY.3p3.10 win4.1C.3p1.10 win5.1C.
3p1.10 win6.1C.3p1.10 win6.1C.3p3.12
CV10006790 18 0 0 26.002 0 0 1 1 1 1
CV10007783 14 0 0 28.779 0 0 0 0 0 1
CV10007820 16 0 0 25.182 0 0 0 0 0 0
CV10007826 16 0 2 30.266 0 2 2 2 2 3
```

Figure N2. *color_nums.pl* Test-case number-containing files with default coloring.

```
verdascend [~/git/comline_color_seq_num] 206> color_nums.pl test-dat.mat -rg 69 71
Table with four cols of average sequence values
dp_10
        70.290
                        70.290
dp 20
        69.500
                64.213
                        69.500
                                64.213
dp 22
                66.563
                                66.563
dp 61
                        69.403
dp_72
        69.403
dp 87
        70.166
                        70.166
        67.750
                64.194
dp_91
                       67.750
                                64.194
dp_104
        69.838
                70.310
                       69.838
                                70.310
dp_143
        66.662
                63.678
                       66.662
                                63.678
dp_147
        68.924
                        68.924
dp_175
        70.184
                        70.184
   214
       67.335 65.891
                       67.335
                               65.891
dp_260 66.187
                59.892
                                59.892
dp_292 69.324 60.051 69.324 60.051
verdascend [~/git/comline color seg num] 207> color nums.pl test-dat.mat -rg 69 71 -ok
Table with four cols of average sequence values
dp_10
        70.290
                76.299
                        70.290
                                76.299
dp_20
        69.500
                64.213
                        69.500
                                64.213
dp_22
        72.403
                66.563
                        72.403
                                66.563
dp_61
        71.595
                74.616
                        71.595
                                74.616
dp_72
        69.403
                72.568
                        69.403
                                72.568
dp_87
        70.166
                82.582
                        70.166
                                82.582
dp_91
        67.750
                64.194
                       67.750
                                64.194
dp_104
       69.838
                70.310
                       69.838
                                70.310
   143
        66.662
                63.678
                        66.662
dp_147
        68.924
                74.292
                       68.924
                                74.292
dp_175
       70.184
                75.051
                        70.184
                                75.051
dp 214 71.625 72.597
                       71.625
                                72.597
      67.335 65.891
                       67.335
                                65.891
dp_260 66.187 59.892 66.187
                                59.892
dp_292 69.324 60.051 69.324 60.051
```

Figure N3. *color nums.pl* Specifying a range of values to highlight (e.g. 69 to 71).

To better highlight specific numbers, subsets of (potentially) number-containing tokens may be ignored. Only values found in a range of columns can be colored, thus allowing some number-containing columns to be easily ignored. Column selection is done with the <code>-col</code> switch, which is associated with two number to specify start and end columns. Use of the <code>-not</code> switch inverts column selection criteria, so that numbers within selected columns are *not* colored but numbers in other columns may be colored. The <code>-iz</code> switch allows zero values to be ignored. Figure N4 shows the result of combining all three above mentioned options, in this case selecting all columns *except* column 5, specifying a coloring range from 2 to 3, and ignoring zeros.

```
verdascend [~/git/comline_color_seq_num] 222> color_nums.pl test-mix.mat -col 5 5 -not -iz -rg 2 3
Names len c2CC.3p5.10 c2CY.3p5.10 tmPey0.3p1.10 c2CC.3p3.10 c2CY.3p3.10 win4.1C.3p1.10 win5.1C.3p1.10
win6.1C.3p1.10 win6.1C.3p3.12
CV10006790
              18 0 0 26.002 0 0 1 1 1 1
CV10007783 14
                0 0 28.779 0 0 0 0 0 1
                0 0 25.182 0 0 0 0 0 0
CV10007820
CV10007826
                0 2 30.266 0
                                2
CV10008635
                0 0 40.532 0 0 3 3 3 2
             14 0 0 38.379 0 0 1 1 1 1
14 0 1 41.133 0 1 1 1 1 1
CV10008675
CV10008677
                0 0 20.602 0 0 2 2 2 0
CV10011080
CV10011088 14 3 3 40.661 3
CV10012722 17 0 0 32.071 0 0 2 2 2 1
CV10018478 15 0 1 39.820 0 1 1 1 1 2
CV10020570 17 0 1 23.102 0 1 1 1 1 2
CV10020577 15 0 0 26.405 0 0 1 1 1 2
CV10021253 17 2 3 31.539 2 3 3 3 3 3
```

Figure N4. *color_nums.pl* Selecting specific columns, ignoring zero values and coloring by value range. All columns except column 5 are selected via the combination of switches -col 5 -not and zeros are ignored (i.e. not colored) because of the -iz switch. Finally, a color range of 2 to 3 is specified for differential value-based color assignment (i.e. in range values are yellow; under range values are blue; over range values are red).

Options for predefined ranges include the -nr switch for 0 to 1, and the -n2 switch for -1 to 1. Figure N5 shows an example.

verdas	verdascend [~/git/comline_color_seq_num] 233> color_nums.pl test-lod.mat -n2												
	Matrix with log odd frquencies for various dinucleotides												
RowNam	es	AA	AC	AG	AT	CA	CC	CG	CT	GA	GC		
p0	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71			
p1	-0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44			
p2	0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76			
p3	1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36			
p4	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70			
p5	1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07			
p6	1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13			
p7	1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81			
p8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56			
p9	0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15			
p10	0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38			
p11	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36			

Figure N5. *color nums.pl* Predefined range coloring, from -1 to 1.

Use of *color_cols.pl*

Calling *color_col.pl* without any arguments (or with "-help" or an unrecognized argument) yields this splash / help screen (which includes colors to illustrate cycling patterns):

```
Usage: <infile> ['-' for stdin] [...options]
  <infile> Text file (e.g. data with 'word' tokens)
  -m # Mark col # (1-based index on tokens)
  -s # Step; Mark every #'th col (starting from 0)
  -o # Offset for starting steps (default 0)
  -g # Group columns / rows # at a time (default 1)
  -cr # Colums in range # to # colored (1-based)
  -nc Invert column range qualifications
  -rr # # Rows in range # to # colored (1-based)
  -nr Invert row range qualifications
  -tab Separate input columns by tab (default is space)
  -sep X Separate input columns by X string
  -2c Two color scheme: Cycle 1 2
  -5c Five color scheme: Cycle 1 2
  -5c Five color scheme: Cycle 1 2 3 4 5 6 7 8 9 10
  -10r Ten rainbow scheme: Cycle 1 2 3 4 5 6 7 8 9 10
  -fg X Set foreground color to X [RYGBCMW] (default red)
  -bg X Set background color to X [RYGBCMW] (default white)
  -iv Invert foreground / background (i.e. for -mark or -step)
  -all Color all lines; Default ignores comment '#'
  -row Apply coloring to *Rows* not columns
  -prd Previous line differences (per char)
```

```
Separate input columns by X string
-sep X
-2c
          Two color scheme:
                               Cycle 1 2
-5c
          Five color scheme:
                                Cvcle 1 2 3 4 5
-10c
          Ten color scheme:
                                Cycle 1 2 3 4 5 6 7 8 9 10
          Ten rainbow scheme: Cycle 1 2 3 4 5 6 7 8 9 10
-10r
-fg X
          Set foreground color to X [RYGBCMW] (default red)
           Set background color to X [RYGBCMW] (default white)
           Invest foreground / background (i
```

Figure C1. *color_cols.pl* Color cycle portion of splash / help screen.

Each command line switch has a brief description of what it does. Default behavior is to color columns by counting non-whitespace "words" and applying the chosen color scheme.

Examples of several color schemes using the test file "test-lod.mat" are shown Figure C2.

ryan@verdascen	d ~/git/	comline_	color_to	ols> colo	or_cols.	pl test-	lod.mat		
Matrix with lo									
RowNames	AA	AC	AG	AT	CA	CC	CG	CT	GA
p0 -0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71
p1 -0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44
p2 0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76
p1 -0.47 p2 0.58 p3 1.71 p4 1.31 p5 1.73 p6 1.01 p7 1.97 p8 0.21 p9 0.31	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36
p4 1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70
p5 1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07
p6 1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13
p7 1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81
p8 0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56
	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15
p10 0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38
p11 -0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36
ryan@verdascen									
ryan@verdascen							lod.mat	-5c	
Matrix with lo	g odd fr	quencies	for var	ious dinu	ıcleotid	es			
RowNames	AA	AC	AG	AT	CA	CC	CG	CT	GA
p0 -0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71
p1 -0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44
p2 0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76
p3 1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36
p4 1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70
p5 1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07
p6 1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13
p7 1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81
p1 -0.47 p2 0.58 p3 1.71 p4 1.31 p5 1.73 p6 1.01 p7 1.97 p8 0.21 p9 0.31	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56
	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15
p10 0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38
p11 -0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36

Figure C2. color_cols.pl Coloring schemes: top default ("-10r"); bottom "-5c".

Selection of subsets of columns / rows is possible by specifying ranges -cr and -rr respectively. Indexes are one-based, and column and row subsets may be specified together. Inversion of range selection (i.e. color columns / rows *outside* of specified ranges) is accomplished with -nc and -nr respectively. Examples of subset selection are shown in Figure C3 where we also explicitly set the background color (to 'm' = magenta) so that subset selection is more clear.

Columns (or rows) can be marked singly or periodically, or processed as multi-token (or multi-row) groups. The mark $_{-m}$ x option will color column (or row) X with the foreground color. Using step $_{-s}$ x one can mark every X'th col (or row), starting at the first X; Combined with offset $_{-s}$ Y the starting position of the stepping can be changed to Y as shown in Figure C4. Grouping columns (or rows) N at a time is done with $_{-g}$ N as shown in Figure C5.

O:		1-4-	/1 <i>i</i>	-1 4-	-1 /-	-11	1	- 1 - d	h	2 7		
	verdascend							t-Lod.mat	-bg m	-CF 3 /		
RowNar		AA	AC	AG Val	AT	CA	CC	CG	СТ	GA	GC	
p0	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	dc	
p1	-0.47	0.06	-0.25	0.58	0.32	-0.89	-0.21	-0.01	-0.43	-0.44		
p2	0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76		
p3	1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36		
p.4	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70		
p4 p5	1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07		
p6	1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13		
p7	1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81		
p8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56		
p9	0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15		
p10	0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38		
p11	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36		
	verdascend											
	/erdascend					olor col	s.pl tes	t-lod.mat	-ba m	-cr 3 7	-rr 5 1	0
	with log								-3			
RowNar		AA	. AC	AG	AT	CA	CC	CG	CT	GA	GC	
p0	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71		
p1	-0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44		
p2	0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76		
p3	1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36		
p4	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70		
p5	1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07		
p6	1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13		
p7	1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81		
p8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56		
р9	0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15		
p10	0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38		
p11	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36		
	/erdascend											
ryan@v	/erdascend	~/git/	comline_c	color_to	ols> ./c	olor_col	s.pl tes	t-lod.mat	-bg m	-сг 3 7	-rr 5 1	0 -nc -nr
	with log								CT			
RowNar		AA	AC 6F	AG	AT F2	CA	CC	CG	CT	GA 71	GC	
p0 p1	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71		
p1 p2	-0.47 0.58	0.06 -0.53	-0.25 -1.51	0.58 0.93	0.39 0.98	-0.89 -0.59	-0.21 -0.75	-0.01 -0.03	-0.43 -0.26	-0.44 -1.76		
р2 р3	1.71	-0.52	0.25	0.86	0.95	-0.43	-0.73	-1.04	-0.25	-1.76		
p4	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70		
р4 р5	1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.70		
p6	1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13		
p7	1.97	0.29	1.98	1.74	0.22	-2.43	-0.24	-0.50	0.62	0.81		
D8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56		
p7 p8 p9	0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15		
p10	0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38		
p11	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36		
	0.01		0.20	0.10	0.00	0	O . L I	0.0.	0.00	0.50		

Figure C3. *color_cols.pl* Column / row subset selection. Top shows selection of columns 3 through 7; Middle shows selection of columns 3-7 *and* rows 5-10; Bottom shows inverted selection of columns 3-7 *and* rows 5-10. In all cases, the background color is explicitly specified as magenta (-bg m).

ryan@verdascend	~/git/	comline_c	olor_to	ols> ./co	olor_cols	s.pl test	t-lod.ma	t -m 3		
Matrix with log										
RowNames	AA	AC	AG	AT	CA	CC	CG	CT	GA	GC
p0 -0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	
p1 -0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44	
p2 0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76	
p3 1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36	
p4 1.31 p5 1.73	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70	
p5 1.73 p6 1.01	0.02 0.31	1.79 1.62	1.62 1.43	0.55 1.49	-2.00 -2.45	-0.44 -0.24	-0.60 -0.38	1.17 1.34	-1.07 -0.13	
p7 1.97	0.29	1.98	1.74	0.22	-2.45	-0.24	-0.50	0.62	0.81	
p8 0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56	
p9 0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15	
p10 0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38	
p11 -0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36	
ryan@verdascend										
ryan@verdascend					olor col	s.pl tes	t-lod.ma	t -s 3		
Matrix with log										
RowNames	AA	AC	AG	AT	CA	CC	CG	CT	GA	GC
p0 -0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	
p1 -0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44	
p2 0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76	
p3 1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36	
p4 1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70	
p5 1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07	
p6 1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13	
p7 1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81	
p8 0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56	
p9 0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15	
p10 0.24 p11 -0.04	0.32 0.21	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38	
<pre>p11 -0.04 ryan@verdascend</pre>		0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36	
ryan@verdascend					alor cole	nl test	t-lod ma	+ -c 3 -	. 1	
Matrix with log	odd fr	quencies	for var	ious din	icleotide	o.hr res	L- LOU . Ma	L -5 J -	0 1	
RowNames	AA	AC	AG	AT	CA	CC	CG	СТ	GA	GC
p0 -0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	
p1 -0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44	
p2 0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76	
p3 1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36	
p4 1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70	
p5 1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07	
p6 1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13	
p7 1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81	
p8 0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56	
p9 0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15	
p10 0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38	
p11 -0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36	

Figure C4. *color_cols.pl* Column / row marking and stepping (cycling). Top shows marking of column 3; Center shows stepping every 3 (with default start of zero); Bottom shows stepping every 3 but with an offset of 1.

ryan@v	erdascend	~/git/	comline_c	olor_to	ols> ./co	olor_col	s.pl tes	t-lod.mat	t -g 2		
	with log										
RowNam		AA	AC	AG	AT	CA	CC	CG	CT	GA	GC
P0 P1 P2 P3 P4 P5 P6	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	
p1	-0.47	0.06	-0.25 -1.51	0.58 0.93	0.39 0.98	-0.89 -0.59	-0.21 -0.75	-0.01	-0.43	-0.44 -1.76	
p2	0.58 1.71	-0.53 -0.52	0.25	0.86	0.98	-0.39	-0.75	-0.03 -1.04	-0.26 -0.25	-1.76	
p3	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70	
0.5	1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07	
p.6	1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13	
p7	1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81	
D8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56	
p8 p9 p10	0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15	
p10	0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38	
p11	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36	
ryan@v	erdascend										
	erdascend							t-lod.mat	t -g 2 -	row	
Matrix	with log	odd fr				ucleotide					
RowNam		AA	AC	AG	AT	CA	CC	CG	CT	GA	GC
p0	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	
p1	-0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44	
p2	0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76	
р3	1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36	
p4	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70	
p5 p6	1.73	0.02	1.79	1.62	0.55 1.49	-2.00	-0.44	-0.60	1.17 1.34	-1.07	
р6 р7	1.01 1.97	0.31	1.62 1.98	1.43 1.74	0.22	-2.45 -2.05	-0.24 -0.87	-0.38 -0.50	0.62	-0.13 0.81	
p8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56	
p9	0.31	0.34	0.31	0.78	0.46	-1.16	-0.68	0.04	0.19	0.15	
p10	0.24	0.32	-0.03	0.72	0.12	-0.67	-0.41	-0.09	0.02	-1.38	
D11	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.36	
	erdascend									0100	
ryan@v	erdascend	~/git/	comline c	olor too	ols> ./co	lor_cols	s.pl tes	t-lod.mai	t -g 2 -i	m 3 -гоw	
	with log										
RowNam	es	AA	AC	AG	AT	CA	CC	CG	CT	GA	GC
p0	-0.79	-0.31	-0.65	0.10	0.52	-0.28	0.00	0.33	0.41	-0.71	
p1	-0.47	0.06	-0.25	0.58	0.39	-0.89	-0.21	-0.01	-0.43	-0.44	
p2 p3 p4	0.58	-0.53	-1.51	0.93	0.98	-0.59	-0.75	-0.03	-0.26	-1.76	
р3	1.71	-0.52	0.25	0.86	0.95	-0.43	-0.68	-1.04	-0.25	-1.36	
p4	1.31	0.00	1.43	1.26	1.93	-1.25	-0.20	-0.87	0.48	-1.70	
p5	1.73	0.02	1.79	1.62	0.55	-2.00	-0.44	-0.60	1.17	-1.07	
p6	1.01	0.31	1.62	1.43	1.49	-2.45	-0.24	-0.38	1.34	-0.13	
p7	1.97	0.29	1.98	1.74	0.22	-2.05	-0.87	-0.50	0.62	0.81	
p8	0.21	0.45	1.27	1.94	0.25	-1.55	-0.83	-0.55	0.64	0.56	
p9 p10	0.31 0.24	0.34	0.31	0.78	0.46 0.12	-1.16 -0.67	-0.68 -0.41	0.04	0.19 0.02	0.15 -1.38	
p10			-0.03 0.29	0.72 0.10		-0.67	0.21	-0.09		0.36	
DII	-0.04	0.21	0.29	0.10	0.00	-0.43	0.21	-0.07	-0.60	0.30	

Figure C5. $color_cols.pl$ Column / row grouping. Top shows grouping of columns by 2 (with default column color scheme); Center shows grouping of rows by 2; Bottom shows grouping and marking combined. The third group (-m 3) of two rows (-g 3) is marked.