MetaLogo: a generator and aligner for multiple sequence logos

Yaowen Chen¹, Zhen He¹, Yahui Men¹, Guohua Dong¹, Shuofeng Hu¹, Xiaomin Ying¹

¹ Center for Computational Biology, Beijing Institute of Basic Medical Sciences, Beijing, China

Corresponding author: Xiaomin Ying, Center for Computational Biology, Beijing Institute of Basic Medical Sciences, Beijing 100850, China. Email: yingxmbio@foxmail.com

Abstract

Sequence logos are used to visually display conservations and variations in short sequences. They can indicate the fixed patterns or conserved motifs in a batch of DNA or protein sequences. However, most of the popular sequence logo generators can only draw a single sequence logo for one group of sequences with the same length, and cannot handle sequences with different lengths or other characteristics. It’s essential to align sequence logos in a group level to reveal the similarities and differences on conservation patterns among groups. To solve these problems, we developed MetaLogo, which can draw sequence logos for sequences from multiple groups in one single plot and align multiple logos to highlight the sequence pattern dynamics across groups, and can also provide statistical figures for sequence conservations and group correlations, thus allowing users to investigate functional motifs in a more delicate and dynamic perspective. We provide users a public MetaLogo web server (http://metalogo.omicsnet.org), a standalone Python package (https://github.com/labomics/MetaLogo), and also a built-in web server available for local deployment. Using MetaLogo, users can draw informative, customized, aesthetic, and publishable sequence logos without any programming experience to present and investigate new knowledges on specific sequence set.

Keywords
Multiple Sequence logo, Logo alignment, Web server
Introduction

Sequence logo was first proposed by Schneider and Stephens in 1990 [1], and has been widely used thousands of times for sequence pattern visualization in the academic field. Each position of a sequence logo is stacked by different amino acids or nucleotides, with the height of each base indicating its degree of conservation at that position. The most commonly used sequence logo generators include Weblogo [2], Seq2Logo [3], ggseqlogo [4], Logomaker [5] and RaacLogo [6] and others, involving web servers, Python and R packages, etc. By showing graphical representations of sequences, these tools greatly accelerate the researchers' exploration of sequence patterns and motifs.

However, a common problem is that most sequence logo tools only support single group of sequences as input, usually with equal length, users need to select the most representative subgroup of sequences for sequence logo when studying a sequence set, which is generally too simplified to represent all. One solution is to perform multiple sequence alignments (MSA) in advance, and use the gapped and aligned sequences as input to construct a single sequence logo, which has been supported by several tools. However, the problem with using MSA results is that it is difficult to indicate whether there are different patterns or motifs among subgroups. Let us take the B cell receptor (BCR) sequences as an example. Complementarity determining region 3 (CDR3) is the most hypervariable region in BCR and it is known that CDR3s of different lengths may have different affinities for certain antigens. Therefore, to discriminate the length of CDR3s when checking the motifs of CDR3s is essential for immune repertoire analysis. In addition to separately studying motifs for sequences of different lengths, we may also need multiple sequence logos for sequences of the same length but from different groups, which could be generated based on sample sources or clustering results. All of the above requires a convenient tool that allows researchers to take multiple sets of sequences as input, draw sequence logos synchronously and align them at the logo level to display pattern dynamics across different groups, so as to understand the sequence characteristics of the sample in a more delicate manner.

Besides CDR3s analysis, other motif-related studies, including transcript factor motif analysis, CRISPR array analysis, evolutionarily conserved sequences analysis and others, all have the same requirements. Note that most of these applications focus on short sequences or limited ranges of sequences, since it is not practical to visually display longish sequence motifs which exceed the visual capacity of one figure.

To solve the problems, we developed MetaLogo, which satisfies the need to allow variable length or multi-group sequence as input and to make sequence logos while performing multiple
logo alignments, and provides researchers basic statistic analysis to reveal the conservation of each group and relationships among groups. All figures are made in an aesthetic, multi-form, and highly customizable way.

Description

MetaLogo provides a public web server (locally deployable), and a stand-alone Python package at the same time to provide researchers with the most convenient service. Users can input files in *Fasta* or *Fastq* format, and specify grouping by length automatically or by group id indicated in sequence names. MetaLogo draws a separate sequence logo for each group, and then performs alignment for multiple sequence logos in a local or global mode, according to users’ choice.

Similarity metric

For each set of sequences, MetaLogo first calculates the information contents of amino acids or nucleotides at each position in bits [7]. In order to align different sequence logos, we need to measure the similarities between bit arrays of positions from different logos. For example, $P$ and $Q$ are bit arrays of positions from two different protein logos and defined as follows:

\[
P = [p_1, p_2, \ldots, p_i, \ldots, p_n],
\]
\[
Q = [q_1, q_2, \ldots, q_i, \ldots, q_n],
\]

where $n$ is the number of amino acids types and item $p_i$ and $q_i$ represent the information contents of the $i^{th}$ amino acid in the two positions, specifically. The arrays are sorted based on a fixed amino acids order.

To measure the similarity between $P$ and $Q$, MetaLogo provides Dot Production (DP) and Cosine Similarity (COS) for users to choose from, which are commonly used as similarity measures and defined as follows:

\[
DP(P,Q) = \sum_i p_i \cdot q_i,
\]
\[
COS(P,Q) = \frac{DP(P,Q)}{\text{Length}(P) \cdot \text{Length}(Q)},
\]

where $\text{Length}(P)$and $\text{Length}(Q)$ represent the length of vector $P$ and $Q$.

Besides bit arrays, we could also use frequency arrays to measure the similarity between positions. For each amino acid in one position, its frequency could be treated as the probability
of one sequence having it in that position. Thus, here we could use similarity measurements designed for probability distributions.

MetaLogo allows users to choose the Jensen–Shannon divergence (JSD) [8] as the similarity measurement. The JSD is a method of measuring the similarity between two probability distributions, and is a symmetrized version of the Kullback–Leibler (KL) divergence [9]. Note in the following context, $P$ and $Q$ represent discrete probability distributions which sum to one. JSD is defined as follows:

$$JSD(P||Q) = \frac{1}{2} D_{KL}(P||M) + \frac{1}{2} D_{KL}(Q||M),$$

where $D_{KL}(P||M) = \sum_i^n p_i \log \frac{p_i}{M_i}$, $D_{KL}(Q||M) = \sum_i^n q_i \log \frac{q_i}{M_i}$, and $M = \frac{1}{2} (P + Q)$.

Bhattacharyya Coefficient (BC) [10] could also be used as a similarity measurement for two statistical samples. Since probability array does not indicate conservation like bit array do, hence MetaLogo provides an entropy (H) [11] adjusted Bhattacharyya Coefficient (EBC) as a choice to measure the probability array similarity, which is defined as follows:

$$EBC(P||Q) = BC(P||Q) \sqrt{ \left(1 - \frac{H(P)}{H_{max}}\right) \left(1 - \frac{H(Q)}{H_{max}}\right)},$$

where $BC(P||Q) = \sum_{i=1}^n \sqrt{p_i q_i}$, $H(P) = -\sum_{i=1}^n P_i \log P_i$; $H(P)$ is the entropy of $P$ and $H_{max}$ is the max entropy for a $n$-dimensional probability vector.

Among these measurements, COS and KL consider both non-conservative and conservative patterns while DP and EBC only value conservative patterns among groups.

### Alignment

The alignment between sequence logos is based on the Needleman–Wunsch algorithm, which is a classic global sequence alignment algorithm. When using MetaLogo, users can choose two alignment modes, one is pairwise alignments between adjacent sequence groups (Figure 1A), and the other is a global logo alignment among all sequence groups (Figure 1B). For global multi-logo alignment, MetaLogo adopts the method of progressive alignment construction [12]. The closest pair of sequence logos are aligned first, and the next logo closest to the aligned sequence logo set is successively added for alignment. Introduced gaps and inserts of each alignment are retained for subsequent alignments until all logos get aligned.

In the alignment process, users need to specify a certain similarity metric we mentioned above, and also the penalty for inserts and gaps. After padding and alignment, MetaLogo can visually highlight the highly similar pairs of positions between groups by connecting them using colorful strips.
MetaLogo supports four different logo layouts, including horizontal, circular, radial, and 3D layouts. As shown in Figure 1 A-E, these diverse layouts are suitable for different scenes specifically. The horizontal layout is the default one, which can deal with most scenarios; the circular layout can more clearly show the conservations across multiple sequence groups; the radial layout is suitable to display sequences with conservative motifs in the middle or at the end of the sequences, rather than at the beginning; the 3D layout makes sequence logos more diverse and aesthetic.

MetaLogo allows customization of most of the operable elements in the figure, including figure size, ticks size, label size, labels, title, grids, margins between items, colors of items and so on. Specific range of sequences could also be chosen to display for long sequences. Users can also choose whether to display axis, ticks, labels, group ids, etc. Multiple formats of figures are supported, including PNG, PDF, SVG, PS and EPS.

Basic statistics

MetaLogo provides users basic statistics of their input data to help users get a more complete understanding of the data. An entropy heatmap is utilized to indicate the conservation of each position in each group. A boxplot is also used to display the entropy distribution of each group. If a global alignment is performed among sequence groups, a clustered heatmap is also provided to show correlations between every pair of groups, which could help users to understand which groups are more similar in conservation pattern and which groups are more divergent.

Package Install and server deployment

Users can directly access our public web server (http://metalogo.omicsnet.org, Figure 1F), or install MetaLogo in Python package locally. Two examples of sequence sets are provided with the codes. One set contains sequences of E. coli transcription factor binding sites [13] (Figure 1 A-E; MetaLogo web server, example 1), the other set contains sequences of CDR3s of verified antibodies detected in BCR repertoires of individuals with COVID-19 [14] (See MetaLogo web server, example 2). A detailed tutorial for MetaLogo is provided online (https://github.com/labomics/MetaLogo/wiki). After the installation, users can run MetaLogo directly in the system terminal or import MetaLogo functions into their own scripts or projects. Users can also deploy MetaLogo as a web service on their local area network through Docker,
which is convenient for people with no programming experience. Relevant parameters could be set for MetaLogo web server, including the number limitation of allowed sequences and the size limitation of uploaded files, etc.

Use case

In the past two years, the COVID-19 pandemic has brought obvious impact on human’s health and social activities all over the world. Many laboratories have been seeking clues from blood of COVID-19 patients to help treat the disease, and several potential therapeutic antibodies have been discovered [15]. Among these exploratory studies, Galson et al. deeply sequenced B cell receptors (BCR) from a cohort of COVID-19 patients and discovered convergent immune signatures towards SARS-CoV-2 [16]. In total four hundred sixty-three convergent clonotypes, or CDR3s, were detected, after matching to earlier published studies. These clonotypes were detected in at least four of the COVID-19 patients, but not present in healthy controls or individuals following seasonal influenza vaccination, which means they might response specifically to SARS-CoV-2. Since they did not focus on motifs of these sequences, we downloaded the CDR3s of these clonotypes as well as IGHV gene annotations and tried to use MetaLogo to reveal the conservation patterns among these sequences.

Figure 2 shows the results of sequence logos and statistical figures from MetaLogo. Figure 2A shows the multi-group sequence logo of all the 463 clonotypes; MetaLogo automatically divided sequences into different groups according to their lengths. It is obvious that the heads and tails of all the groups show conserved pattern while the middle of them seem more diverse. These logos are aligned by using an adjusted multiple sequence alignment algorithm; similar positions between adjacent groups are connected by light red bands. It is worth noting that in longer groups, like Len20, Len19 and Len18 groups, several Ys in a row are present in sequence logos. In addition, there is an obvious GSY motif in the middle of the sequences in Len16 group. Figure 2B shows sequence counts of each group, which shows that about 30% of sequences have a length of 16. Figure 2C shows entropies of each position of each group. The higher the entropy, the less convergent the position. Figure 2D shows entropy distributions of each group. From the entropy perspective, we can see the middle of the sequences are more diverse than both ends, and longer sequences have more higher entropies, but the Len16 group has an apparent lower median entropy that its adjacent groups, which is interesting.

To further explore the characteristics of sequences, we selected sequences from the Len14, Len15 and Len16 groups with annotated V genes IGHV 3-30, IGHV 3-30-3 and IGHV 3-33,
which are the most frequently annotated genes in this dataset. From Figure 2E, it reveals that not all sequence with 16 length have the GSY motif. It is indicated that the GSY motif comes from the IGHV 3-30 and IGHV 3-30-3 genes, and more enriched in sequence with 16 length than other lengths. Figure 2F shows the clustering result of groups, groups Len16-V3-30 and Len16-V3-30-3 are clearly clustered together, which shows the specificity of combination of length 16 and annotation IGHV 3-30 (or 3-30-3).

According to the above results from MetaLogo, we can conclude that sequences with 16-length and IGHV 3-30 (or 3-30-3) IGHV gene annotation have a clear motif GSY in the middle of the sequences and are distinct from sequences from other groups. We checked the published potential antibodies against SARS-CoV-2 from OAS (Observed Antibody Space) [17] database and found many antibodies contain the motif GSY in their CDR3 sequences, which is consist with our results but reminds the necessity of in-depth study of this phenomenon. The results also indicate the capacity of MetaLogo to discover valuable knowledge among sequences in a convenient way.

Conclusion

MetaLogo is a new generator for aesthetic, customized and informative sequence logos. Unlike existing tools, MetaLogo can draw multiple sequence logos for sequences from different groups in one figure, and perform alignment of sequence logos, as well as basic statistical analysis, to reveal the pattern dynamics across groups. MetaLogo provides a free web server for public use, as well as a stand-alone Python package and a docker web service for local deployment. We will value the suggestions and comments from users, and continue to maintain code updates and upgrades to continuously contribute to the community.

To be noted, MetaLogo does not take phylogeny into account when making and aligning sequence logos, which means MetaLogo-produced sequence logos can only tell the patterns of the sampled sequences provided by users, rather than of the whole ecological background. Since it is more sophisticated to precisely estimate evolutionary conservation, we will consider add a new feature in the future version of MetaLogo to address that problem.

Key points

- MetaLogo is a new sequence logo generator for variable-length sequences or multi-group sequences;
- MetaLogo performs pairwise and global sequence logos alignment to highlight the sequence pattern dynamics across different sequence groups.
MetaLogo provides basic statistical analysis to additionally reveal the sequence conservation convergence and divergences among sequence groups.

MetaLogo provides public web server, deployable local web server with docker, as well as stand-alone Python package for making highly customized sequence logos.

**Funding**

This work was supported by National Science and Technology Major Project grant [2018ZX10201001] and by the National Natural Science Foundation of China grant [31970567].

**Conflicts of interest**

The authors have declared no competing interests.

**Acknowledgments**

We thank colleagues in our lab including Pu Liu, Chao Feng, Sijing An and Runyan Liu for their careful reviews and feedbacks on the MetaLogo web server.

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Figures

Figure 1. Layouts, alignment modes and web server interface provided by MetaLogo.  
A. Horizontal layout of MetaLogo under pairwise alignment mode. Conserved positions between sequence logos are indicated by grey bands. B. Horizontal layout of MetaLogo under global alignment mode. Sequence logos were filled with paddings according to a global logo alignment. Conserved positions among sequence logos are indicated by grey bands. C. Circular layout of MetaLogo under global alignment mode. D. Radial layout of MetaLogo under global alignment mode. E. 3D layout of MetaLogo under global alignment mode. F. Web server interface provided by MetaLogo.

Figure 2. Sequence logos and analysis result from MetaLogo on the BCR clonotypes.  
A. Sequence logos for all the BCR clonotypes. Sequences are divided into different groups by lengths. B. Sequence counts of each length-divided group. C. Entropy heatmap of positions in all groups. Xs represents gaps. D. The entropy distribution of positions in each group. E. Sequence logos for sequences with 14, 15, and 16 lengths, as well as IGHV v3-33, V3-30, V3-30-3 annotations. F. Clustering result of sequence logo groups to reveal the relationships among them.
