Applications Note



Sequence analysis

FlexiDot: highly customizable, ambiguity-aware dotplots for visual sequence analyses

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Abstract

Summary: FlexiDot is a cross-platform dotplot suite generating high quality self, pairwise and all-against-all visualizations. To improve dotplot suitability for comparison of consensus and errorprone sequences, FlexiDot harbors routines for strict and relaxed handling of ambiguities and substitutions. Our shading modules facilitate dotplot interpretation and motif identification by adding information on sequence annotations and sequence similarities. Combined with collage-like outputs, FlexiDot supports simultaneous visual screening of large sequence sets, enabling dotplot use for routine analyses.

Availability and implementation: FlexiDot is implemented in Python 2.7. Software and documentation are freely available at http://github.com/molbio-dresden/flexidot.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

First described five decades ago (Gibbs and McIntyre, 1970), dotplots remain effective tools for sequence exploration, still conveying key messages in current research (Hosaka et al., 2017). Dotplots allow characterization of complex or repetitive sequences, visual detection of DNA motifs and identification of modular similarities between sequences.

Despite advances in dotplot algorithms and availability of different software tools, essential features are missing and, if available, scattered across various tools (Table 1, Supplementary Material). This prompted us to combine established functionalities (e.g. allagainst-all modes) with new features for dotplot improvement (base ambiguity handling, shading, integration of annotations), while retaining usability and customizability.

2 Features and implementation

FlexiDot is a multi-purpose dotplot suite for publication-ready dotplots, handling self, pairwise and all-against-all comparisons with individual and combined visualizations (Fig. 1A-C, see Supplementary Material for details). We want to highlight that (i) our mismatch and ambiguity handling enables analyses of degenerate consensus sequences and error-prone long reads (Fig. 1B), and that (ii) our sequence similarity and annotation-based shadings for self and all-against-all representations (Fig. 1A and C, respectively) convey descriptive information to facilitate sequence interpretation.

The FlexiDot algorithm identifies matches, transforms them into diagonals and creates clear vector images (pdf, svg) or standard raster graphics (png). Less stringent matching is possible by addressing ambiguous residues specifically or by allowing a defined number of substitutions. A tabular output with lengths of the longest match (longest common subsequence, LCS) of all sequence pairs is provided.

FlexiDot integrates highly customizable shadings: (i) Self dotplot regions can be highlighted according to their sequence annotation provided as general feature file (Fig. 1A). (ii) All-against-all comparisons can be shaded according to the LCS length in forward, reverse or both directions (Fig. 1C). (iii) The user can provide a matrix with numerical values (e.g. identities) to guide shading. Matrix values can be displayed in the dotplot.

FlexiDot uses Python 2.7 with numpy, matplotlib, biopython, regex, colormap and colour libraries. It is operated from the command line under Windows, Linux, and Mac. Input sequences are either specified as single or multi-fasta, or automatically detected in the working directory.

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Table 1. Feature list of commonly used dotplot tools

Tools	Ambiguity handling	Ambiguity Annotation All-against-all Batch Interactive handling shading mode analyses GUI	ll-against-all mode	Batch I analyses		Input: DNA-DNA I	Input: JNA-protein pr	Input: otein-protein	Input: Input: Input: Multiple output Reverse Self/pairwise Similarity Strict/relaxed DNA-DNA DNA-protein protein-protein formats complement collages shading matching	Reverse complement	Self/pairwise collages	Similarity shading	Strict/relaxed matching	Citation
FlexiDot	+	+	+	+	1	+	1	+	+	+	+	+	+/+	here
Dotmatcher	ı	ı	ı	+	ı	+	I	+	+	I	I	ı	+/-	Rice et al. (2000)
Dotter	I	I	+	I	+	+	+	+	I	+	I	ı	+/+	Sonnhammer and
														Durbin (1995)
Dottup	ı	ı	1	+	ı	+	I	+	+	I	I	ı	-/+	Rice et al. (2000)
Gepard	I	+	+	I	+	+	I	+	I	+	I	ı		Krumsiek et al. (2007)
PolyDot	ı	ı	+	+	ı	+	I	+	+	I	ı	ı	_/+	Rice <i>et al.</i> (2000)
YASS webserver	+	ı	I	ı	+	+	I	+	ı	+	I	ı	+/+	Noé et al. (2005)

GUI, Graphical user interface.

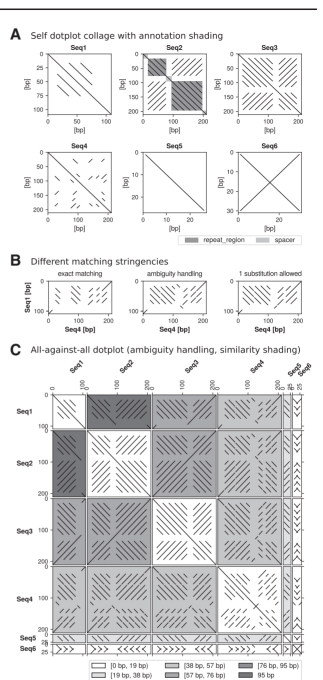


Fig. 1. Visual sequence comparison by FlexiDot with window size 10 using six artificial test sequences. (A) Self dotplot collage. The Seq2 dotplot is shaded with custom annotations. (B) Influence of ambiguity and mismatch handling on pairwise dotplots. (C) All-against-all dotplot of the six sequences with ambiguity handling and similarity shading

3 Application

As demonstrated for a variety of use cases in the Supplementary Material, FlexiDot creates publication-ready figures for complex sequences. This facilitates:

- evaluation of tandem repeat higher order structures of error-prone long reads, e.g. as seen in Sevim *et al.* (2016) and Symonova *et al.* (2017),
- combined depiction of sequence structure and functional annotations,
- identification of conserved motifs in related sequences,

- gene or repeat comparisons using degenerated consensus sequences (Schwichtenberg et al., 2016; Weber et al., 2013),
- analysis of terminal or internal inverted or direct repeats, e.g. for transposable element annotation (Hosaka et al., 2017).

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