Full Paper LaTeX2 ε Template for Genome Informatics 2001

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Abstract

This document is a LaTeX2 ε template file for preparing a full paper. Please read the instructions carefully and prepare your manuscript.

Keywords: aberrant splicing, database, point mutation, scanning model

1 Introduction

The page limit for full paper is TEN. In case the number of pages still exceeds this limit after your efforts, please contact the Editor-in-Chief (Satoru Miyano: giw@ims.u-tokyo.ac.jp) for advice.

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2 Method and Results

Of course you can create more sections with any section titles.

2.1 Tables

Table 1 is an example of a table.

Table 1: Bioinformatics conf	ferences in 2001.	
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Conference	Date	Site
PSB	January 3-7	Hawaii
RECOMB	April $22-25$	$\operatorname{Montréal}$
ISMB	July 21-25	$\operatorname{Copenhagen}$
GIW	December 17-19	Tokyo

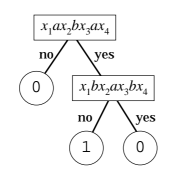


Figure 1: This is a sample figure.



Figure 2: This is a color sample figure.

2.2 Figures

When you use figures (Fig. 1, Fig. 2) with a special style file, please provide it.

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Here are examples of citation [1, 2, 3, 4, 6]. Reference style and citation should strictly follow this template.

- 1. References should be arranged in the alphabetical order of authors and all references should be cited in the paper.
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 - (a) Journal: author(s), title, journal name, vol(issue):startpage-endpage, year.
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3 Discussions

Authors are recommended to use spell checker program and grammar check program for improving your final version.

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