Databases and ontologies

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DeOri: a database of eukaryotic DNA replication origins

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ABSTRACT

Summary: DNA replication, a central event for cell proliferation, is the basis of biological inheritance. The identification of replication origins helps to reveal the mechanism of the regulation of DNA replication. However, only few eukaryotic replication origins were characterized not long ago; nevertheless, recent genome-wide approaches have boosted the number of mapped replication origins. To gain a comprehensive understanding of the nature of eukaryotic replication origins, we have constructed a Database of Eukaryotic ORIs (DeOri), which contains all the eukaryotic ones identified by genome-wide analyses currently available. A total of 16145 eukaryotic replication origins have been collected from 6 eukaryotic organisms in which genome-wide studies have been performed, the replication-origin numbers being 433, 7489, 1543, 148, 348 and 6184 for humans, mice, Arabidopsis thaliana, Kluyveromyces lactis, Schizosaccharomyces pombe and Drosophila melanogaster, respectively.

Availability: Database of Eukaryotic ORIs (DeOri) can be accessed from http://tubic.tju.edu.cn/deori/

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INTRODUCTION

The initiation of replication is the central event in the cell cycle. In all three domains of life, DNA replication begins at specialized loci termed replication origins. In bacteria, replication initiates from a single replication origin, whereas eukaryotic organisms exploit many replication origins (Costas et al., 2011). In eukaryotes, the origins of DNA replication in the budding yeast Saccharomyces cerevisiae and fission yeast Schizosaccharomyces pombe have been well-characterized (Hayashi et al., 2007; Heichinger et al., 2006; Nieduszynski et al., 2007; Segurado et al., 2003). Recently, a high-resolution genome-wide map of DNA replication origins in Kluyveromyces lactis has also been generated (Liachko et al., 2010). However, only few origins were characterized in the genomes of higher eukaryotes not long ago. Fortunately, recent development of genome-wide approaches has led to a boost in the number of replication origins that have been mapped in eukaryotic genomes, including those in human (Cadoret et al., 2008; Karnani et al., 2010), mouse (Cayrou et al., 2011; Sequeira-Mendes et al., 2009), Arabidopsis thaliana (Costas et al., 2011) and Drosophila melanogaster (Cayrou et al., 2011). The availability of increasing origins throughout the genomes has created opportunities for analysis of the origins on a genome scale. Up to date, the DNA

2 DATABASE CONTENT

All information in DeOri is stored and operated using MySQL relational database management system. One entry in the database corresponds to an origin in a certain organism. The corresponding organism, chromosome, cell type (if any), genomic location, length, GC content, origin sequence and some related links have been displayed. In order to facilitate the comparative genomic analysis and visualize the chromosome context of the origins, the URLs that link to UCSC Genome Browser (Fujita et al., 2011) or NCBI Map Viewer (Sayers et al., 2011) have also been provided. The database access is via a web interface based on PHP script and provides various ways to search for DeOri records, such as DeOri ID, cell type, species and chromosome. In addition, users can also BLAST (Altschul et al., 1997) a query sequence against DeOri to find a homologous one. DeOri will be updated timely and a total of 16 145 eukaryotic DNA replication origins were collected from 6 eukaryotic organisms in genome-wide studies, including 433 human origins, 7489 mouse origins, 1543 A.thaliana origins, 148 K.lactis origins, 348 S.pombe origins and 6184 D.melanogaster origins in the current release (Table 1), which can be accessed from http://tubic.tju.edu.cn/deori/.

Table 1. Contents of DeOri version 1.0

Organism	ORI no.	References
Arabidopsis thaliana	1543	Costas et al., 2011
Drosophila melanogaster	6184	Cayrou et al., 2011
Human_1	283	Cadoret et al., 2008
Human_2	150	Karnani et al., 2010
Kluyveromyces lactis	148	Liachko et al., 2010
Mouse_ES_1	2412	Cayrou et al., 2011
Mouse_ES_2	98	Sequeira-Mendes et al., 2009
Mouse_MEF	2231	Cayrou et al., 2011
Mouse_P19	2748	Cayrou et al., 2011
Schizosaccharomyces pombe	348	Hayashi et al., 2007;
		Heichinger et al., 2006;
		Segurado et al., 2003
	Arabidopsis thaliana Drosophila melanogaster Human_1 Human_2 Kluyveromyces lactis Mouse_ES_1 Mouse_ES_2 Mouse_MEF Mouse_P19	Arabidopsis thaliana 1543 Drosophila melanogaster 6184 Human_1 283 Human_2 150 Kluyveromyces lactis 148 Mouse_ES_1 2412 Mouse_ES_2 98 Mouse_MEF 2231 Mouse_P19 2748

replication origin database for budding yeast (Nieduszynski et al., 2007) has been built, whereas the database of other eukaryotic origins has not been available. In order to gain a comprehensive understanding of the nature of eukaryotic replication origins, we have constructed a Database of Eukaryotic ORIs (DeOri), which contains all the eukaryotic DNA replication origins identified by genome-wide analyses currently available.

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3 DATA RETRIEVAL

The replication origins stored in DeOri include those identified in the genomes of human (Cadoret et al., 2008; Karnani et al., 2010), mouse (Cayrou et al., 2011; Sequeira-Mendes et al., 2009), A.thaliana (Costas et al., 2011), D.melanogaster (Cayrou et al., 2011), K.lactis (Liachko et al., 2010) and S.pombe (Hayashi et al., 2007; Heichinger et al., 2006; Segurado et al., 2003). In the human genome, 283 replication origins have been systematically mapped using the HeLa S3 suspension cell line (Cadoret et al., 2008) and in an independent study, 150 new origins have been identified in adherent HeLa cells in the ENCODE area (Karnani et al., 2010). In mouse, identification of preferential sites of DNA replication initiation in 0.4% of the mouse genome has resulted in 97 new ORIs (Sequeira-Mendes et al., 2009). By performing a genome-wide analysis in both *Drosophila* and mouse cell lines, up to 2748 ORIs on mouse Chromosome 11 (P19 cells) and 6184 ORIs in the Drosophila genome have been characterized (Cayrou et al., 2011). In A.thaliana, by high-throughput sequencing of newly synthesized DNA, ~1500 putative have been identified at the genome-wide level (Costas et al., 2011). In K.lactis, a total of 148 ARSs have been identified by a predict-and-verify approach (Liachko et al., 2010). In the S.pombe genome, 385 ORIs have been predicted initially from AT content calculation (Segurado et al., 2003). Subsequent experiments have also confirmed most of the prediction (Hayashi et al., 2007; Heichinger et al., 2006). In addition, the whole genome sequences of human (hg17), mouse (mm8, mm9), *D.melanogaster* (dm2) were downloaded from http://hgdownload.cse.ucsc.edu/downloads.html, and the whole genome sequences of A.thaliana, S.pombe and K.lactis were downloaded from the NCBI FTP server (ftp://ftp.ncbi.nih.gov/genomes/). Based on the position information provided in the literatures or by personal communication and the corresponding genome sequences, we have obtained the sequences and other information of the origins. It should be noted that the positions of the origins in A.thaliana and S.pombe have been relocated and mapped to the newest releases of genome sequences from GenBank in order to remove the sequencing errors in the older versions, and the positions of the origins in Mouse_ES_2 have also been relocated and mapped to the genome sequences of mouse (mm9) in order to make the UCSC Genome Browser available.

4 CONCLUSION

Consequently, we have constructed a database DeOri, which contains all the eukaryotic DNA replication origins identified by genome-wide analyses currently available. With the availability of the origins newly identified by genome-wide analyses, we will update DeOri constantly to include more entries and integrate more information for each entry. This database will facilitate the comparative genomic analysis of replication origins, and provide some insight into the nature of replication origins on a genome scale. One of the applications is to predict replication origins based on homologous sequence search against the origins of closely related species in DeOri. For example, if query sequences in the rice genome

using BLAST have homologous origins of *A.thaliana* in DeOri, it is likely that the query sequences are also served as origins in the rice genome. Another application is to find some principles for specific organisms by the genome-wide characterization of the origins in DeOri, which will be useful to develop new algorithms to predict replication origins. For example, we can use some tools for motif discovery to find the conserved elements within the origins of a specific organism.

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Conflict of Interest: none declared.

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