HAYATI Journal of Biosciences 22 (2015) 181-185

Contents lists available at ScienceDirect



# HAYATI Journal of Biosciences

journal homepage: http://www.journals.elsevier.com/ hayati-journal-of-biosciences

# Original research article

# Putative DNA-dependent RNA polymerase in Mitochondrial Plasmid of *Paramecium caudatum* Stock GT704





# Trina Ekawati Tallei

Department of Biology, Faculty of Mathematics and Natural Sciences, University of Sam Ratulangi, Manado, Indonesia.

#### ARTICLE INFO

Article history: Received 13 November 2015 Received in revised form 13 January 2016 Accepted 13 January 2016 Available online 3 February 2016

KEYWORDS: ciliate, DNA-dependent RNA polymerase, linear mitochondrial plasmid, Paramecium caudatum

### ABSTRACT

Mitochondria of *Paramecium caudatum* stock GT704 has a set of four kinds of linear plasmids with sizes of 8.2, 4.1, 2.8 and 1.4 kb. The plasmids of 8.2 and 2.8 kb exist as dimers consisting of 4.1- and 1.4-kb monomers, respectively. The plasmid 2.8 kb, designated as pGT704-2.8, contains an open reading frame encodes for putative DNA-dependent RNA polymerase (RNAP). This study reveals that this RNAP belongs to superfamily of DNA/RNA polymerase and family of T7/T3 single chain RNA polymerase and those of mitochondrial plasmid of fungi belonging to Basidiomycota and Ascomycota. It is suggested that RNAP of pGT704-2.8 can perform transcription without transcription factor as promoter recognition. Given that only two motifs were found, it could not be ascertained whether this RNAP has a full function independently or integrated with mtDNA in carrying out its function.

Copyright © 2016 Institut Pertanian Bogor. Production and hosting by Elsevier B.V. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

#### 1. Introduction

Plasmids are extrachromosomal genetic elements found in various organisms. Two types of plasmid structures known so far, namely the double-stranded closed circular plasmid which commonly found in cells of prokaryotes and some eukaryotes, and single- and double-stranded linear plasmids found in fungi, yeast, and ciliates (Tallei *et al.* 2002; Hausner 2003).

Linear plasmid has terminal inverted repeat at both ends and covalently-bound terminal protein (TP) at 5' end which is similar to some viral genome structure (Andrade *et al.* 2009). Two open reading frames (ORFs) of these plasmids encode for DNA and RNA polymerases which are required for their replication (Griffiths 1995; Oeser and Tudzynski 1989; Chan *et al.* 1991). RNA polymerase (RNAP) in the linear plasmids are DNA-directed RNAP (DNAdependent RNAP) and abbreviated as RNAP, which serves to catalyze the synthesis of a DNA chain using DNA as a template. The RNAP characteristic is similar to those of the bacteriophage and yeast mitochondria. The 5' end, besides being involved in replication, also has a role in the integration of the plasmid into mitochondrial genome (Griffiths 1995; Andrade *et al.* 2009; Andrade *et al.* 2013). Mitochondrial plasmid of *Moniliophthora perniciosa* can integrate fully into the mitochondrial genome (Andrade *et al.* 

E-mail address: trina\_tallei@unsrat.ac.id.

Peer review under responsibility of Institut Pertanian Bogor.

2009). The plasmid is likely to have a function in the aging process (senescence) (Chan *et al.* 1991; van Diepeningen *et al.* 2008).

*Paramecium* is the first example in which an mtplasmid was discovered among ciliates. *Paramecium caudatum* stock GT704 has type-II DNA with two sets of mtplasmids. The 8.2 and 2.8 kb DNAs are dimers of the 4.1 and 1.4 kb DNA, respectively (Tallei *et al.* 2002; Endoh *et al.* 1994). The plasmid 2.8 kb (designated as pGT704-2.8) has an ORF for RNAP. This study aimed at characterizing RNAP encoded by pGT704-2.8 and analyzing its relationship with RNAPs from other organisms to predict its function in mitochondria.

### 2. Material and Methods

#### 2.1. Stock and culture

From previous research conducted by Tallei *et al.* (2002), *P. caudatum* stock GT704 used in this study contains plasmid type II. Plasmid type II always has four types of plasmid DNA with sizes of 8.2, 4.1, 2.8, and 1.4 kb. Plasmid DNA 8.2 and 2.8 kb are always in the form of dimer, each of monomer molecule of 4.1 and 1.4 kb, respectively. The cells of *P. caudatum* were grown at 25 °C in fresh lettuce juice medium inoculated with non-pathogenic strains of *Kleibsiella pneumoniae* 1 day before use.

### 2.2. Purification of mitochondria

The cell suspension was filtered with gauze to remove dirt, centrifuged, and washed twice in a Dryl solution (Tsukii *et al.* 1994). Mitochondrial purification was performed following the method

http://dx.doi.org/10.1016/j.hjb.2016.01.002

<sup>1978-3019/</sup>Copyright © 2016 Institut Pertanian Bogor. Production and hosting by Elsevier B.V. This is an open access article under the CC BY-NC-ND license (http:// creativecommons.org/licenses/by-nc-nd/4.0/).

of Goddard and Cummings (1975) with slight modification. The centrifuged cells were resuspended in two times volume of ice cold mannitol solution (0.44 M mannitol, 1 mM KCl, 1 mM Tris-HCl pH 7.5, and 0.25% bovine serum albumin). Cells of *P. caudatum* were crushed using a homogenizer until almost all the cells ruptured. Homogenized cells were resuspended in ice cold mannitol and centrifuged at 600 g for 6 minutes at 0 °C. Supernatant was centrifuged at 5,000 g for 6 minutes. Mitochondrial pellets were resuspended in 1 mL of mannitol solution containing 30 U of DNAse I (Takara Shuzo Co., Japan) and 5 mM MgCl<sub>2</sub>. Mitochondrial suspension was incubated on ice for 25 minutes and added with mannitol solution containing 10 mM EDTA, followed by centrifugation at 5,000 g for 6 minutes. Pellets were used for pure mitochondrial fraction.

#### 2.3. Extraction of mitochondrial DNA and plasmids

Mitochondria were lysed in 0.1 M Tris-HCl (pH 8.0), 50 mM EDTA, 1% (w/v) sodium dodecyl sulfate and 0.1 mg/mL proteinase K for 50 °C for 3 hours. Lysate was extracted twice with phenol (same volume) and treated with proteinase K (Tsukii *et al.* 1994).

#### 2.4. DNA sequencing

Nucleotide sequencing of pGT-704-2.8 kb was conducted at Genetics Laboratory, University of Kanazawa, Japan.

# 3. Results

Mtplasmids 8.2 and 2.8 kb are dimers of monomers 4.1 and 1.4 kb, respectively. Sequence analyzed was dimer 2.8 kb (pGT704-2.8) (GenBank accession: FJ222255). Nucleotide Basic Local Alignment Search Tool result shows that pGT704-2.8 encodes for putative DNA-directed RNAP. This RNAP has sequence identity with RNAP encoded by several mtplasmids found in fungi, namely pHC of *Hebeloma circinans* (75%), pLP of *Pleurotus ostreatus* (74%), pFv of *Flammulina velutipes* (70%), pEM of *Agaricus bitorquis* (59%), pAL2-1 of *Podospora olymera* (55%), kal of *Gelasinospora sp.* (54%), kalilo of *Neurospora intermedia* (54%), and pCLK of *Claviceps purpurea* (53%).

Nucleotide sequence of pGT704-2.8 was translated using special code for mitochondria. The following is the amino acid sequence of putative DNA-directed RNAP:

## MNHRDKYQEYENIYDYDEKSSI**NSLEA**NNLHKIDALLVKHVLEVLD VIT**IHDCF**GTRIKDVAKLIDVVNIYYQRYTSNDNYSIFILX

The sequences are in conserved domain of RNAP (Figure 1). The figure shows that RNAP of pGT704-2.8 is a part of single chain RNAP

family and resembles RNAP of T-even phages group (T3/T7). Basic Local Alignment Search Tool protein search reveals that this putative protein has identity with RNAP of other mtplasmids found in *Hebeloma circinans* (41%), *Pleurotus ostreatus* (40%), *Agaricus bitorquis* (39%), *Flammulina velutipes* (39%), *Claviceps purpurea* (32%), *Gelasinospora* sp. (30%), *Neurospora intermedia* (29%), *Podospora anserina* (30%), and RNAP of Phage T3 (53%).

Figure 2 shows the alignment of conserved domain of several fungal RNAPs which has high identity with RNAP found in pGT704-2.8. Figure 3 shows motifs resulted using Multiple EM for Motif Elicitation (MEME) software developed by Bailey and Elkan (1994). Based on this MEME motif, RNAP of pGT704-2.8 does not have solid motif on block X so it cannot be produced by the program, while the block XI can be generated. Figure 4 shows phylogenetic tree reconstructed using Geneious version 8.0.4. RNAP of pGT704-2.8 is in monophyletic group (clade 1) with RNAP of mtplasmids of Hebeloma circinans, Pleurotus ostreatus, and Agaricus bitorquis, all of them belong to division Basidiomycota which have two linear plasmids in their mitochondria. Flammulina velutipes also belongs to division Basidiomycota although it is in the same monophyletic group (clade 2) together with Claviceps purpurea, Podospora anserina, Gelasinospora sp., and Neurospora intermedia which belong to division Ascomycota that have one linear plasmid in their mitochondria.

# 4. Discussion

Linear mtplasmids have often been found in fungi and plants. The first discovery of this kind of plasmid in animal kingdom was in *P. caudatum* (protozoa: ciliates) (Tallei *et al.* 2002). Furthermore, the linear mtplasmid was found in *Oxytricha trifallax* (protozoa: ciliates) by Swart *et al.* (2012). This plasmid has a size of approximately 5 kb and has 251 bp sequence with identity 82% with its mitochondrial genome (mtDNA), so it is suggested that this mtplasmid had been integrated in the mtDNA.

Circular mtplasmid usually has one ORF encodes for one DNA polymerase or one reverse transcriptase, while linear mtplasmid generally has two ORFs encode for putative DNA polymerase and RNAP, in which their amino acid motifs have homology to viral polymerases (Griffiths 1995). In linear plasmids, DNA polymerase and RNAP each is encoded by one single plasmid, or encoded by different plasmid if there are more than one plasmids in mitochondria (Cermakian *et al.* 1997).

According to Griffiths (1995), the characteristics of the linear plasmid among others, have the ORF that encodes a viral type of DNA polymerase and RNAP similar to RNAP of bacteriophage or



Figure 1. Conserved domain of RNA polymerase (RNAP) of pGT704-2.8 based on conserved domain search developed by Marchler-Bauer *et al.* (2011) showing that this RNAP belongs to DNA-dependent RNAP, a family of single chain RNAP and resembles T3/T7-like RNAP.

		Х			XI			
Paramecium caudatum	EYENIYDYDE	KSSINSLEAD	NLHKIDALLV	KHVLEVL-DV	ITIHDCFGTR	IKDVAKLIDV	V-NIYYQRYT	SNDNYSIFI
Pleurotus ostreatus	DYSSIKTGLM	PNFIHSLDAS	NIHMLINLII	QLQIKN-MNL	YTIHDCFATD	YKNIALLEIL	IKKSFTDIYF	NKK-YLDFI
Hebeloma circinas	DYPKIKLGLM	PNFIHSLDAS	NIHLLISNIN	KYNLKD-LNL	YTIHDCFASD	YKNIAIIELL	VKHSFIELYF	QVD-YLEAI
Agaricus bitorquis	DVLSTKRSFM	PNFIHSLDAS	NVHLLLNSV-	SYKN—LPV	YTVHDCFAST	ANNMFKLEKL	VKNAFINIYF	NDEGYLLKL
Gelasinospora sp.	DSRREVPAII	PNIIHSLDAA	HLPMLI	DSWDS Y	ILEIHDCFGT	HPNDMFKLAE	QVRECFILLY	SKND-FLSK
Neurospora intermedia	DSRREVQAII	PNIIHSLDAS	HLTMII	DSWDS Y	ILFIHECFGT	HPNDMYKLAE	QVRECFILLY	SKND-FLNK
Flammulina velutipes	NTFKQELAII	PNVIHSLDAS	HLYEIILETR	KNNINN	ILAVHDCFGT	HPNKMGDLIH	AVRKTFIIQY	TDLN-FLNK
Claviceps purpurea	DKVKQQIALM	PNLIHSLDAA	SL-IMLYFAF	NKSIGSGVVN	FYSVHDCYGV	TAKYIDLLIS	QLRAVYIELY	SKDG-YIAH
Podospora anserina	NNSKQITAFM	PNLVHSLDAA	SLTLLLDFYF	KESID-VKNI	YTIHDCFAVP	ANKMECLISL	LKLTYIKLYS	DDK-YLLKL
Enterobacteria Ph T3	DAHKQESGIA	PNIVHSQDGS	HLRMTVVYAH	EKYGIES	FAIIHDSFGT	IPADAGKLFK	AVRETMVITY	ENND-VLAD

Figure 2. Amino acid sequences alignment of pGT704-2.8 RNA polymerase (RNAP) with other fungal RNAPs showing two conserved motifs at blocks X and XI.



Figure 3. Motifs X and XI generated using MEME motif.



Figure 4. Phylogenetic tree reconstructed using Geneious version 8.0.4 with Unweighted Pair Group Method with Arithmetic Mean (UPGMA) method without outgroup. Alignment type is global alignment with free end gaps and cost matrix Blosum 62. Jukes-Cantor was used for genetic distance model. Clade 1 has two linear plasmids and clade 2 has one linear plasmid, except *Flammulina velutipes* which has two linear plasmids. The scale bar corresponds to 0.1 amino acid substitutions per site.

yeast mitochondria. This type of RNAP is also named DNAdependent RNAP, which is a single polypeptide and does not require any other polymerase for promoter recognition. DNAdependent RNAP catalyzes the transcription of DNA into RNA by using four ribonucleotide triphosphate as its substrate (Youn *et al.*) 2001). RNAP coding gene is probably the gene that undergoes fragmentation because of plasmid integration into mtDNA through recombination. Integrated plasmid sequences have degenerated over time in the absence of selection pressure (Hausner 2003). Plasmid RNAP is presumably bifunctional that recognizes

sequences at the end terminal of plasmid and promoter sequence (Vickery 2011).

Putative RNAP encoded by pGT704-2.8 is homolog to RNAP of Teven bacteriophages and of mtplasmids from divisions Basidiomycota and Ascomycota. Analysis using QuickPhyre (Kelley and Sternberg 2015) shows that RNAP of pGT704-2.8 is located on DNA/RNAP superfamily and on T7 RNAP family with 92.1% confidence by the single highest scoring template. This is supported by Figure 1 which shows conserved domain of RNAP of pGT704-2.8 is based on conserved domain search, suggesting that this RNAP belongs to DNA-dependent RNAP, a family of single chain RNAP and resembles T3/T7-like RNAP. In this group of RNAP, a number of conserved motifs have been identified (Chan et al. 1991). Cermakian et al. (1997) states that motif X (INSLEA: Ile-Asn-Ser-Leu-Glu-Ala in pGT704-2.8) resides in the nucleotides 783-788 while motif XI (IHDCF: Ile-His-Asp-Cys-Phe) resides in the nucleotides 810–814. Considering the length of pGT704-2.8 as 2.8 indicates that the ORF is located in the center. Motif INSLEA is a conserved motif at block X according to Oeser (1988). Motif proposed by Oeser and Tudzynski (1989) for block XI is bHDCf. This matches motif IHDCF found in pGT704-2.8 (Figure 2).

In well characterized RNAP of bacteriophage T7, these motifs were found in a proteolytic fragment which does not have 172 *N*-terminal amino acid but still retains a catalytic activity. According to Chan *et al.* (1991), the presence of these motifs (motifs I–XII) indicates that the RNAP is functional. The discovery of only two motifs found in pGT704-2.8 is because the sequencing process is done only partially and randomly. The most conserved part of a protein usually is an important part of the function of the protein. This section is often surrounded by a less conserved structure that modifies the function of the protein. The changes to these additional structures often result in minor changes to the character of protein, for example, the differences in substrate specificity, although the function of the protein is not changed (Černý *et al.* 2014).

Conserved domain usually is a functional domain, so most likely proteins that are within one family that has the same motif shares the same function as well. Sequence alignment and phylogenetic analysis are aimed at inferring gene function. If the percentage of identity between two compared proteins is >35% (>40% in case of long sequence alignment), it is alleged that the two proteins are homologous, because it has the same structure and function. Homology will decrease when it is in a twilight zone, 20%–35% identity of protein sequence (Rost 1999). Hence, RNAP of pGT704-2.8 can be said to be homologous to the mitochondrial RNAP from plasmid contained in *Hebeloma circinans* (41%), *Pleurotus ostreatus* (40%), *Agaricus bitorquis* (39%), *Flammulina velutipes* (39%), and RNAP of enterobacteria phage T3 (53%).

In the phylogenetic tree (Figure 4), RNAP of pGT704-2.8 is clustered together in a group of RNAP of fungal mtplasmid of Hebeloma circinans, Pleurotus ostreatus, and Agaricus bitorquis which belong to division Basidiomycota with two linear plasmids (clade 1). This is supported by the finding from our previous research (Tallei et al. 2002) that pGT704 has two sets of linear plasmids. Flammulina velutipes also has two linear plasmids and belongs to division Basidiomycota although not clustered in clade 1. Claviceps purpurea, Podospora anserina, Gelasinospora sp. and Neurospora intermedia belong to division Ascomycota with one linear plasmid. Mitochondrial plasmid pMLPw from Pleurotus ostreatus has 5'-end TP. This plasmid has two ORFs, one of them encodes for protein product homolog to RNAP of yeast mitochondria and bacteriophages (Kim et al. 2000). Mtplasmids found in fungi and plants have 5'-end TP and terminal inverted repeat, and encodes for RNAP (Meinhard et al. 1990). Previous research conducted by Tsukii *et al.* (1994) revealed that mtplasmid GT704 has covalently bonded protein at 5'-end. Based on this information, it is assumed that pGT704-2.8 shares similar structure.

The similarity between mitochondrial linear plasmid with some phages and viruses implies that they may have originated from a common ancestor, namely virus (Kempken *et al.* 1992; Griffith 1995). Even though RNAP of pGT704-2.8 has a high identity with enterobacteria phage T3 (53%), but it has a big gap. A gap is a space that is introduced to the process of alignment to compensate for insertions and deletions. RNAP of T7 has *N*-terminal domain that participates in interaction of RNAP at promoter's specific sites and the binding to the adjacent RNA by changing conformation in the area. At element-1 of AT-rich (adenine -thymine-rich) DNA recognition area (residues 93–101), specificity loop (residues 739–770) and intercalating  $\beta$ -hairpin including Val237 (residues 230–245) hold responsibility for promoter recognition, binding, and melting (Draculic *et al.* 2014).

The same thing occurred to the function of putative replication protein O of *M. perniciosa*. Although there is lack of information, but with the high identity of its structure and sequence with other RNAP mtplasmids, it was assumed that implication of this enzyme was in the mechanism and expression of mitochondrial genome (Andrade *et al.* 2009). The enzyme has a length of 766 amino acids and belongs to family of single chain RNAP that is common to virus and cellular organelles (Andrade *et al.* 2013). The gene of this protein is expressed in almost all phases of fungal life cycle (Andrade *et al.* 2013).

Replication of bacteriophage T7 DNA is initiated by the binding of T7 RNAP to origin of replication site and production of short RNA molecule that functions as primer (Shi 2013). Carboxyl terminus of phage's enzyme is thought to be involved in promoter recognition, while its catalytic activity is located at terminal end of protein at the downstream region of amino acids 172 (T7) and 173 (T3). Amino-terminal end of T7 RNAP is proposed to be involved in stability of the polymerase-promoter complex (Masters *et al.* 1987). RNAP of bacteriophages T7/T3 has a simple structure, namely single protein subunit capable of performing transcription cycle without any other protein transcription factor. The rate of synthesis is higher than RNAP in bacteria (Tunitskaya and Kochetkov 2002).

Genes of RNAP, besides having a very important function in the process of transcription and replication, their sequences evolve relatively rapid, so it can be used as a marker in phylogenetic studies. The gene can also be used in the study of DNA barcoding (Singh *et al.* 2012; Krawczyk and Sawicki 2013; Enan and Ahamed 2014).

This study infers that DNA-dependent RNAP of mtplasmid of *P. caudatum* pGT704-2.8 belongs to the family of single chain RNAP and homolog to RNAP of T-even bacteriophages group, as well as RNAP of mtplasmid from divisions Basidiomycota and Ascomycota. Its function is to perform the transcription without any other protein transcription factor, because the protein is involved in promoter recognition and catalyzing transcription of DNA into RNA using four type of ribonucleotide triphosphate as it substrate. It is also presumed that this RNAP is involved in mtplasmid replication. Considering only the two conserved motifs found in this protein, it could not be ascertained whether the protein has a full function independently or integrated with mtDNA in carrying out its functions.

## **Conflict of interest**

The author declares that there is no conflict of interest attached to this publication.

#### Acknowledgments

The author thank the Association of International Education of Japan for providing grant to conduct the research at Laboratory of Genetics, Faculty of Science, Kanazawa University, Japan.

### References

- Andrade B, Souza C, Góes-Neto A. 2013. Molecular docking between the RNA polymerase of the *Moniliophthora perniciosa* mitochondrial plasmid and rifampicin produces a highly stable complex. *Theor Biol Med Model* 10:15. http:// dx.doi.org/10.1186/1742-4682-10-15.
- Andrade BS, Taranto AG, Góes-Neto A, Duarte AA. 2009. Comparative modeling of DNA and RNA polymerases from *Moniliophthora perniciosa* mitochondrial plasmid. *Theor Biol Med Model* 6:22. http://dx.doi.org/10.1186/1742-4682-6-22.
- Bailey TL, Elkan C. 1994. Fitting a mixture model by expectation maximization to discover motifs in biopolymers. In: Proceedings of the Second International Conference on Intelligent Systems for Molecular Biology. California, Menlo Park: AAAI Press. pp. 28–36.
- Cermakian N, Ikeda TM, Miramontes P, Lang BF, Gray MW, Cedergren R. 1997. On the evolution of the single-subunit RNA polymerases. J Mol Evol 45:671–81.
- Černý J, Bolfiková BČ, Valdés JJ, Grubhoffer L, Růžek D. 2014. Evolution of tertiary structure of viral RNA dependent polymerases. *PLoS ONE* 9(5):e96070. http:// dx.doi.org/10.1371/journal.pone.0096070.
- Chan BS-S, Court DA, Vierula PJ, Bertrand H. 1991. The kalilo linear senescenceinducing plasmid of *Neurospora* is an invertron and encodes DNA and RNA polymerases. *Curr Genet* 20:225–37.
- Draculic S, Wang L, Cuéllar J, Guo Q, Velázquez G, Martín-Benito J, Sousa R, Valpuesta JM. 2014. Yeast mitochondrial RNAP conformational changes are regulated by interactions with the mitochondrial transcription factor. Nucl Acids Res 42(17):11246–60. http://dx.doi.org/10.1093/nar/gku795.
- Enan MR, Ahamed A. 2014. DNA barcoding based on plastid matK and RNA polymerase for assessing the genetic identity of date (*Phoenix dactylifera* L.) cultivars. *Genet Mol Res* 13(2):3527–36. http://dx.doi.org/10.4238/2014.
- Goddard JM, Cummings DJ. 1975. Structure and replication of mitochondrial DNA from Paramecium aurelia. J Mol Biol 97(4):593–609.
- Griffiths AJ. 1995. Natural plasmids of filamentous fungi. Microbiol Mol Biol Rev 59(4):673-85.
- Hausner G. 2003. Fungal mitochondrial genomes, plasmids and introns. Applied Mycology and Biotechnology 3:101–31.
- Kelley LA, Sternberg MJE. 2015. The Phyre2 web portal for protein modeling, prediction and analysis. *Nature Protocols* 10:845–58. http://www.sbg.bio.ic.ac.uk/ phyre/.
- Kempken F, Hermanns J, Osiewacz HD. 1992. Evolution of linear plasmid. J Mol Evol 35:502-13.

- Kim K, Jeong J, Youn HS, Koo YB, Roe J. 2000. The terminal protein of a linear mitochondrial plasmid is encoded in the N-terminus of the DNA polymerase gene in white-rot fungus *Pleurotus ostreatus*. *Curr Genet* 38:283–90.
- Krawczyk K, Sawicki J. 2013. The uneven rate of the molecular evolution of gene sequences of DNA-dependent RNA polymerase I of the genus *Lamium L. Int J Mol Sci* 14:11376–91. http://dx.doi.org/10.3390/ijms140611376.
- Marchler-Bauer A, Lu S, Anderson JB, Chitsaz F, Derbyshire MK, DeWeese-Scott C, Fong JH, Geer LY, Geer RC, Gonzales NR, Gwadz M, Hurwitz DI, Jackson JD, Ke Z, Lanczycki CJ, Lu F, Marchler GH, Mullokandov M, Omelchenko MV, Robertson CL, Song JS, Thanki N, Yamashita RA, Zhang D, Zhang N, Zheng C, Bryant SH. 2011. CDD: a Conserved Domain Database for the functional annotation of proteins. *Nucleic Acids Res* 39(D):225–9.
- Masters BS, Stohl LL, Clayton DA. 1987. Yeast mitochondrial RNA polymerase is homologous to those encoded by bacteriophages T3 and T7. *Cell* 51:89–99.
- Meinhard F, Kempken F, Kämper J, Esser K. 1990. Linear plasmids among eukaryotes: fundamentals dan application. Curr Genet 17:89–95.
- Oeser B. 1988. S2 plasmid from Zea mays encodes a specific RNA polymerase: an alternative alignment. *Nucleic Acids Research* 16(17):8729.
- Oeser B, Tudzynski P. 1989. The linear mitochondrial plasmid pClK1 of the phytopathogenic fungus *Claviceps purpurea* may code for a DNA polymerase and an RNA polymerase. *Mol Gen Genet* 217(1):132–40.
- Rost B. 1999. Twilight zone of protein sequence alignments. *Protein Eng* 12(2): 85-94.
- Shi Y. 2013. Linking Transcription to DNA Replication in Mammalian Mitochondria [thesis]. Stockhol: Karolinska Institute.
- Singh HK, Parveen I, Raghuvanshi S, Babbar SB. 2012. The loci recommended as universal barcodes for plants on the basis of floristic studies may not work with congeneric species as exemplified by DNA barcoding of Dendrobium species. *BMC Research Notes* 5:42. http://dx.doi.org/10.1186/1756-0500-5-42.
- Swart EC, Nowacki M, Shum J, Stiles H, Higgins BP, Doak TG, Schotanus K, Margrini VJ, Minx P, Mardis ER, Landweber LF. 2012. The Oxytricha trifallax mitochondrial genome. Genome Biol Evol 4(2):146–54. http://dx.doi.org/ 10.1093/gbe/evr136.
- Tallei TE, Moeis MR, Endoh H. 2002. Determinasi urutan nukleotida pada plasmid mitokondria Paramecium caudatum stok GT704. Jurnal Bionatura 4(3):174–88.
- Tsukii Y, Endoh H, Yazaki K. 1994. Distribution and genetic variability of mitochondrial plasmid-like DNAs in *Paramecium*. Jpn J Genet 69:685–96.
- Tunitskaya VL, Kochetkov SN. 2002. Structural-functional analysis of bacteriophage T7 RNA polymerase. *Biochemistry (Mosc)* 10:1124–35.
- van Diepeningen AD, Debets AJ, Slakhorst SM, Hoekstra RF. 2008. Mitochondrial pAL2-1 plasmid homologs are senescence factors in Podospora anserina independent of intrinsic senescence. *Biotechnol J.* 3(6):791–802 http://dx.doi.org/ 10.1002/biot.200800005.
- Vickery DB. 2011. A functional analysis of the kalDNA plasmid from senescent strains of Neurospora intermedia [Dissertation]. British Columbia, Canada: University of British Columbia. http://dx.doi.org/10.14288/1.0101043.
- Youn HS, Koo YB, Roe JH. 2001. Nucleotide sequence of 7.2 kb mitochondrial linear plasmid DNA in *Pleurotus ostreatus*. The Korean Journal of Microbiology 37(1): 37–41.