Chloroplast genomics: Expanding resources for an evolutionary conserved miniature molecule with enigmatic applications

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Abstract

Chloroplast, methylation deprived uniparental organelle genome is the most studied organelle genome from the perspective of evolution and functional omics. Recent advances in organelle genome sequencing both in terms of genome or transcriptome sequencing has opened a wide range of opportunities to understand the transcriptional and translational role of the genes mainly involved in the light harvesting apparatus and the evolution of the inverted repeats across the lineage. However, as compared to the nuclear genome, limited resources are available in case of organelle genome. In this review, we discuss the recent advances in the chloroplast genomics and the resources that have been developed for understanding the evolution, repeat patterns, functional genomics of this miniature molecule with enigmatic applications.

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1. Chloroplast genomes: dynamic organization and evolutionary fluctuations

Photosynthesis is critical to all aspects of plant life and to combat the environmental fluctuations. Chloroplast, evolutionarily conserved and endomictobiotically originated molecule play a major role in photosynthesis by acting as host to three major complex such as photosystem II (PSII), the cytochrome b6f complex (Cytb6f), and photosystem I (PSI) [1]. Evolutionary conservation of these complexes in chloroplast genome thylakoid membrane represents the main sites of the light capture and the oxygen production as well as playing a major role in the light state transitions with plastid division apparatus responsible for the binary fission spatially distributed between the stromal and cytosolic space [2]. Among the spatially distributed genes in circular fashion, chloroplast represents a set of genes, which are vital for controlling the photosynthetic efficiency and to determine the dynamic organization of the thylakoid membrane and cyclic electron flow [3]. Evolutionary conserved organization of chloroplast genomes, which is circular in nature and follows a D-loop replication model is structured in a quadrupartite structure, which is partitioned into two repeat regions, which are defined by the differences in the length as large single copy (LSC) regions spanning across a length of 80–90 kb and a short single copy region (SSC), representing a 16–27 kb region. Organization of these LSC and SSC regions is a dynamic process and has been widely reported to undergo expansion and contraction [4]. Although the evolutionary conservation of the chloroplast genic regions has been widely reported as exemplified by their use as molecular barcodes, few instances of rapidly evolving genes such as rbcL, which encodes the large subunit of ribulose-1,5-bisphosphate carboxylase/oxygenase (RUBISCO), and plays a major role in carbon assimilation and other genes such as matK (maturase K), ndhB and psbA-J, which are involved in modulating the state transitions has also been seen [5]. In contrast, the repeat organization is very dynamic and although conserved across the angiosperms, dynamic loss of the inverted repeat copies has been widely documented amongst the gymnosperms [6]. Taking all these structural variations with in the size of (150–160 kb), it worth to highlight the role of evolutionary conserved, distinct and model genome to understand the genome fluctuations (Fig. 1). From the viewpoint of regulatory genomes, transcriptional and, transcriptional flux, chloroplast genomes have been widely explored in addition to point mutants and also the identification of the RNA Editing events.
2. Resources and tools for chloroplast genomics and functional genomics

In the past few years, considerable focus has been leveraged on the development of tools and genomic resources for advancing the chloroplast genomics. Here we outline the recent tools that have been developed for the chloroplast genomics from the viewpoint of genome assembly, annotation, evolutionary aspects, repeats, markers, and functional genomics (Table 1).

2.1. Organelle genome assembly

ORGanelle ASseMbler [7] is a useful tool for assembling of organelle genome sequences such as mitochondria and plant plastid genomes. The tool can be used to assemble small sequences that are over-represented in a whole genome shotgun sequence dataset. ORGanelle ASseMbler is a command line open source software tool developed using Python libraries and works in Linux and MacOSX systems. The implemented algorithm is linearized in a three step model: 1. The first step indexes the sequencing reads and then assembles the organelle genome with an option of the assembling graph in GML format. The assembling graph can be visualized using any graph visualization tools. The last step involves the extraction of the organelle specific sequences from the graph in a single FASTA file. The limitation of the algorithm is that the implemented algorithm is not capable of reorienting the circular structure of the chloroplast genome and thus manual curation is required to circularize the genome.

NOVOPlasty [8] is a recently published algorithm, which uses seed-extend based assembler and perform the organelle genome assembly by hashing the sequences from the whole genome sequencing (WGS) runs into a table, which allows for the rapid sorting of the sequences. Assembling of the organelle genomes starts with the seed sequence, which acts as an anchor to the extend the seed bi-directionally. The unique feature of the NOVOPlasty is that the user can specify the sequences, which can be a single organelle genome reads, a closely related organelle gene or an evolutionary related or distinct chloroplast genome. The unique feature of NOVOPlasty is that using the seed-extend approach, the assembler assembles the genome in circularized format if both the ends of seed-extend overlap by 200 bp [8]. The tool works on both Linux and MacOSX operating system.

2.2. Functional annotation

Functional annotation of chloroplast genome is an important process, as the rate of molecular evolution depends on the well annotated genome. Previously DOGMA [9] has been developed and has been the gold standard for the annotation of chloroplast genomes. Here, we describe some of the recent tools that have been developed for the functional annotation:

PLANN (Plastome Annotator) [10] is a command-line tool developed for the automated annotation of the assembled chloroplast genomes by comparing the user given chloroplast genome to a well annotated chloroplast genome. The tool has been designed to work on unix-based operating systems including Linux and MacOSX. PLANN uses NCBI tools BLASTN, tbl2asn and Sequin to perform the annotation. The graphical user interface application Sequin is first used to generate a template file for the output. The input files include a new plastome fasta sequence, a reference plastome file in GenBank format and a Sequin template file. PLANN annotates the new plastome by matching sequences with the gene sequences of reference genome using BLASTN and tbl2asn, and then transforms them into corresponding genomic locations of the new plastome in Sequin format.
Table 1
List of the recently developed resources for chloroplast genomics.

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<tr>
<td>NOVOPlasty</td>
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<td>ChloroSSRdb</td>
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CpgAVAS (Chloroplast Genome Annotation, Visualization, Analysis and GenBank Submission) [11] is an online webserver meant to provide standard functions to annotate and analyze chloroplast genome sequences. Additionally, it can generate circular genome maps, summary statistics of annotated genome and creation of files for GenBank submission. CpgAVAS has been developed to overcome the limitations of DOGMA, the popularly used chloroplast genome annotation web server. CpgAVAS has been implemented using Perl Catalyst Web Application Framework and a combination of Perl programs. The CpgAVAS server accepts a completely sequenced chloroplast genome as input and predicts its protein coding regions, rRNA genes, tRNA genes and inverted repeats through the comparative annotation with well annotated chloroplast genomes. It also includes tRNAscan [12] for the prediction of the tRNA in the chloroplast genomes. Protein coding regions are predicted using ab initio gene prediction tools and similarity based approaches. GenBank annotations of chloroplast genomes are first used to cluster the protein, CDS and rRNA gene sequences into homologous groups and formed into one blast-able database. The database is further used to create reference protein and cDNA/rRNA gene dataset for each input genome sequence. The reference protein, cDNA and rRNA genes are searched using Blastx, Blastn, protein2genome, est2genome programs and corresponding best hits are used to annotate the input genome sequence. Further, inverted repeats of the input genome are identified using the Vmatch [13] software tool and tRNAs are identified using tRNAscan [12].

Verdant [14] is a new developed database driven suite of tools specifically designed for annotation, alignment and tree generation of chloroplast genomes. It is a web-based software connected to a database that provides accurate annotation of chloroplast genomes without manual intervention. Verdant uses different programs namely annoBTD (unpublished), MAFFT [15], RaxML [16], Circos [17] and JBrowse [18] to perform defined functions. AnnoBTD has been implemented to automate annotation without any manual editing. Protein coding regions of the input genome are identified by using de novo ORF identification, which is a novel feature of AnnoBTD. rRNAs and tRNAs are detected and annotated by blastn. Very few exons which are missed by other annotation programs are also detected by AnnoBTD. The extensive features of Verdant not only includes the annotation of the chloroplast genome but also allows for the automated alignment of the annotated genes, rRNAs and tRNAs, introns and intergenic regions using progressive and iterative refinement algorithms as implemented in MAFFT. Alignments done using the MAFFT can be passed to RaxML for phylogeny estimation. For the visualization of annotations, Circos and JBrowse has been implemented, which allows the visualization of the circular features. Verdant is developed using PHP, MySQL, Perl, JS, HTML and CSS. Users can create their own projects and can perform taxon selection, feature selection, alignment and phylogenetic tree reconstruction.

CGAP (Chloroplast Genome Analysis Platform) [19] is a comprehensive resource developed for comparative analysis of chloroplast genomes. CGAP is an interactive web-based tool with features like genome collection, visualization, phylogenetic analysis, content comparison and annotation of complete chloroplast genomes. It contains a back-end database of hundreds of complete chloroplast genomes including their annotation features such as genes, CDS, tRNA, rRNA, promoter, exon/intron regions, and repeats. The visualization module of CGAP can be used to create high quality genome maps to visualize circular complete genomes, linear regional genomes, modified published genomes and user unpublished genomes. CGAP can also be used to compare the similarities and differences of the feature content between different chloroplast genomes. CGAP is also integrated with phylogeny tools that uses an alignment free method for tree generation and comparison. The Genome Annotation module of CGAP can be used to annotate new chloroplast genomes based on the reference chloroplast genomes in the CGAP database by using BLAST programs. CGAP has been developed using Python language and Web2py web framework.

2.3. Visualization

Visualization of the chloroplast genome characteristics is an important feature that allows the display of chloroplast genes in circular fashion with additional features such as rRNAs, tRNAs, genic regions and IR boundaries. For the visualization of the chloroplast genome, OrganellarGenomeDraw (OGDRAW) [20,21] has been developed, which allows for the display of the genomic features and allows the user to create high quality circular and linear graphs. An important feature present in OGDRAW allows for the visualization of the expression data from the transcript profiling, polysome profiling or from the proteomics experiments. The software allows to display the transcriptional and translational status of the chloroplast encoded genes. Besides, web-based, OGDRAW is also available as a Perl module, which can be integrated into the annotation pipelines. CpgAVAS [22] and Verdant [14] provides inbuilt visualization of the annotated chloroplast genome using the OGDRAW. Colour settings in the OGDRAW for the clockwise and counter clockwise genes can be easily edited using the configuration file and java enabled OGDRAWConfig [20].

2.4. Markers and codon usage

The main application of the chloroplast genome has been attributed to the development of the molecular markers primarily due to the conserved gene regions and the ease of the development of the polymorphic markers. ChloroMitoSSRDB [23] and ChloroMitoSSRDB2.00 [24] is the first repository that has been developed for the large-scale visualization of the simple sequence repeats (SSRs) across the chloroplast genomes. The developed platform offers several features such
as the visualization of the distribution of repeat patterns using dynamic graphs, and the cross-linking of the identified repeats to the genic or non-genic regions. The developed platform also offers a comparative assessment of the two repeat mining algorithms IMEx [25,26] and MISA [27] and allows the repeat mining using the commonly used tools under one comparative framework. ChloroSSRdb [28] is a repeat mining framework, which is focused primarily on green plants.

Another aspect that made the chloroplast genomes distinct is the use of the chloroplast genomes for functional genomics by over-expressing the gene of interest. Chloroplast based plant functional factories has been widely exploited to develop over-expression of immunogenic vaccines. Recently developed ChloroMitoCU [29] offers a comparative assessment of the codon usage profiles across the chloroplast genomes. Currently, ChloroMitoCU contains 29,960 complete (full-length) protein-coding genes (CDGs) from all reference clades of chloroplasts genomes. The unique features of ChloroMitoCU involves the comparative assessment of the codon usage profiles across phylogenetic distant and related chloroplast genomes. Additionally, ChloroMitoCU allows for the comparative assessment of the codon usage patterns across the previously analysed chloroplast genome and the user submitted chloroplast genes.

2.5. Transcriptional profiling and RNA-editing

RNA polymerases and association of six sigma factors play a major role in maintaining the transcriptional based expression profiling [30]. Associative role of these polymerases and sigma factors provide important understanding of the role of splicing, gene editing and expression profiling of mutants in response to environmental stresses. Recently developed ChloroSeq [30] presents an optimized pipeline, which combines the spliced alignment tool tophat, bowtie, and bedtools, which have been developed previously for the genomic architecture visualization to process chloroplast RNA-seq expression profiles and allows for the estimation of the expression quantification across the exon and introns, splicing efficiency and the putative RNA Editing sites. RNA editing is a post-transcriptional process, which mainly involves the conversion of cytidine-to uridine and forms an important part of the RNA maturation process [31,32]. Across the plant lineage, evolutionary conserved RNA editing factors such as CRR28 and RARE1 have been widely shown to affect the cytidine-to-uridine conversion in sites mainly associated with ndh genes such as ndhBeU467PL, ndhDeU878SL and accDeU794SL [32]. PREPACT [33] has been widely used for the estimation of the RNA-Editing events from angiosperms like Arabidopsis thaliana, Orzya sativa to the early branching angiosperms such as Amborella. PREPACT allows the detection of the RNA Editing by comparative analysis across the 17 pre-implemented chloroplast transcriptomes and provides a user adjustable stringency threshold of 90 – 70% for the detected RNA editing events. PREPACT operates in three distinct modes which allows for the prediction of the RNA-Editing events based on the alignment prediction, CDA or BLASTx predictions. Recent implication of the RNA-Editing events allowed for the construction of the first synthetic operon in chloroplast genomes [34].

3. Conclusion

Chloroplast genomics is at the forefront of biology with the advent of the next generation sequencing, coupled with the advances in the assembling strategies applying novel seed-extend approaches to allow the assembly of the completely circularized genome. Recent advances have mainly focussed on the annotation and the comparative genomics of chloroplast genomes, thus allowing a better development of chloroplast genomes as plant factories.

Authors Contribution

GS conceived, designed the research and wrote the MS; GB contributed to the MS writing; DE and PJR provided the revisions and edits to the MS.

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