

Aberystwyth University

Draft genome assembly of the biofuel grass crop Miscanthus sacchariflorus De Vega, Jose; Donnison, Iain; Dyer, Sarah; Farrar, Kerrie

Published in: F1000Research

DOI: 10.12688/f1000research.44714.1

Publication date: 2021

Citation for published version (APA): De Vega, J., Donnison, I., Dyer, S., & Farrar, K. (2021). Draft genome assembly of the biofuel grass crop Miscanthus sacchariflorus. *F1000Research*, *10*, [29]. https://doi.org/10.12688/f1000research.44714.1

Document License CC BY

General rights

Copyright and moral rights for the publications made accessible in the Aberystwyth Research Portal (the Institutional Repository) are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the Aberystwyth Research Portal for the purpose of private study or research.

- You may not further distribute the material or use it for any profit-making activity or commercial gain
 You may freely distribute the URL identifying the publication in the Aberystwyth Research Portal

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

tel: +44 1970 62 2400 email: is@aber.ac.uk

DATA NOTE



Draft genome assembly of the biofuel grass crop *Miscanthus*

sacchariflorus [version 1; peer review: 2 approved]

Jose De Vega¹, Iain Donnison², Sarah Dyer¹, Kerrie Farrar²

¹Earlham Institute, Norwich, NR4 7UZ, UK

²Institute of Biological, Environmental & Rural Sciences (IBERS) - Aberystwyth University, Aberystwyth, SY23 3EE, UK



Keywords

Miscanthus, biofuel, C4, assembly, annotation



This article is included in the Draft Genomes collection.

Corresponding author: Jose De Vega (jose.de-vega@earlham.ac.uk)

Author roles: De Vega J: Conceptualization, Formal Analysis, Investigation, Writing – Original Draft Preparation; Donnison I: Conceptualization, Funding Acquisition, Project Administration, Writing – Review & Editing; Dyer S: Conceptualization, Funding Acquisition, Project Administration, Writing – Review & Editing; Farrar K: Conceptualization, Funding Acquisition, Project Administration, Writing – Review & Editing

Competing interests: No competing interests were disclosed.

Grant information: This work was funded by core strategic funding from the Biotechnology and Biological Sciences Research Council (BBSRC) in projects BBS/E/T/000PR9818 and BBS/E/W/10963A01A.

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Copyright: © 2021 De Vega J *et al.* This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

How to cite this article: De Vega J, Donnison I, Dyer S and Farrar K. Draft genome assembly of the biofuel grass crop *Miscanthus* sacchariflorus [version 1; peer review: 2 approved] F1000Research 2021, 10:29 https://doi.org/10.12688/f1000research.44714.1

First published: 18 Jan 2021, 10:29 https://doi.org/10.12688/f1000research.44714.1

Introduction

Miscanthus is a genus of C4 perennial rhizomatous grasses native to East Asia and Oceania, and naturally adapted to a wide range of climate zones and land types. *Miscanthus sacchariflorus* is among the most widely distributed species within the genus. It originated in the Yellow Sea region of China and can be predominantly found in cool latitudes of East Asia with varying ploidy¹. *M. sacchariflorus* occurs in both diploid (2n=38) and tetraploid (2n=76) forms, where tetraploid *M. sacchariflorus* genotypes originated by autopolyploidy². *M. sacchariflorus* probably has the greatest winter hardiness among all the Saccharinae³.

Natural interspecific *Miscanthus* hybrids are commonly observed, even between individuals of different ploidy. For example, introgression of *M. sacchariflorus* is often found among cultivated European *M. sinensis* ecotypes^{1,4}. Furthermore, *M. x giganteus*, a sterile triploid hybrid resulting from the hybridization between *M. sinensis* and *M. sacchariflorus*, is the predominant commercially grown species owing to its high biomass productivity and low chemical input requirements. The common occurrence of hybridization events and variable ploidy are challenging to the improvement of these bioenergy grasses and increase the need for genomic resources from different *Miscanthus* species. A chromosomal-scale reference genome using a doubled-haploid *M. sinensis* line was recently published⁴.

We assembled, annotated and validated a draft genome from the diploid *M. sacchariflorus* cv. "Robustus 297" genotype, as well as generating rhizome, stem and leaf RNA-Seq data from the same genotype. This dataset was previously used to verify that both *M. sinensis* and *M. sacchariflorus* share the same A/B ancestral tetraploidy⁴. Here, we present the first draft genome of *M. sacchariflorus*, the second *Miscanthus* genome available after *M. sinensis*⁴.

Methods

Plant materials and sequencing

DNA was extracted from leaves from the diploid *M. sacchariflorus* cv. "Robustus 297" genotype (Biosample SAMN08580354) using the Qiagen DNeasy kit. RNA was also extracted from leaf, stem and root tissues from the same plant. All samples were taken from a plant grown from seed in trays in a glasshouse in 2009. This genotype is established and used in breeding at IBERS (Wales, UK). The RNA-seq libraries were deposited as part of previous work in the BioProject PRJNA639832.

Whole genome sequencing and assembly

We obtained ~5.86e9 pairs of 100 bp paired-end reads from an Illumina paired-end library with a 560 bp insert-size that was sequenced on Illumina HiSeq 2500 machines in rapid run mode by the Earlham Institute. This represents approximately 50X coverage of the heterozygous content and 100X coverage of the homozygous content of the genome. Read quality was assessed, and contaminants and adaptors removed using Kontaminant⁵. These paired-end short-reads were assembled into 17M contigs with a total length of 3.27 Gb using ABySS⁶ version 1.5.1, with default options and a kmer size of 71.

We obtained ~141.1e6 pairs of reads from a Nextera 150 bp mate-pair library with approximately 7 Kb insert-size, which was used for scaffolding the previous contigs together with the paired-end reads, using SSPACE⁷ without "extension" step. Nextera mate-pair reads were required to include a fragment of the adaptor to be used in the scaffolding step⁵, and we filtered out sequences shorter than 500 bp. We obtained 589K scaffolds, a total length of 2.54 Gb with an N50 of 10.2 Kb. This whole-genome assembly was denominated "Msac_v2" and is deposited at NCBI in BioProject PRJNA679435.

Gene model and functional annotations

Our gene structure annotation pipeline⁸ used five sources of evidence that were provided to AUGUSTUS⁹ (version 2.7) for gene annotation: (1) Repetitive and low complexity regions of the scaffolds identified using RepeatMasker¹⁰ (version open-4.0.5) based on homology with the RepBase¹¹ public database (Release 20140131) and a new database of repeat elements identified in the assembly with RepeatModeler¹². The repeats annotation was deposited in Zenodo (See data availability); (2) exon-intron junctions identified by Tophat¹³ (version 2.1.0); (3) de novo and genome-guided ab initio transcripts assembled with Trinity¹⁴ (version 2.6.5)and Cufflinks¹⁵ (version 2.2.1) from RNA-Seq reads obtained from several tissues from the same genotype; (4) ab initio gene models predicted by SNAP¹⁶ (version 29-11-2013) and GeneID¹⁷ (version 1.4.4); and (5) homology-based alignments of transcripts and proteins from Miscanthus sinensis and maize using Exonerate¹⁸ with a minimal identity of 0.7 and coverage of 0.7. Finally, AUGUSTUS9 was run with the options "genemodel=complete" and "alternatives-from-evidence=true" to ensure that the predicted genes were compatible with all the previous provided evidence.

For the functional annotation of these predicted genes, translated gene sequences were compared with the NCBI non-redundant (nr 20170116) proteins and EBI's InterPro (version 5.22.61) databases, and the results were imported into Blast2GO¹⁹ to annotate the GO and GO slim terms, enzymatic protein codes and KEGG pathways. A similar GO annotation from translated gene sequences can be done with eggNOG-mapper²⁰. These functional descriptors were deposited in Zenodo (See *Underlying data*).

Anchoring the whole genome assembly using the Miscanthus sinensis reference

To improve the genome contiguity, we anchored our M. sacchariflorus scaffolds to the Miscanthus sinensis genome⁴. However, no nucleotide content from M. sinensis was incorporated in the M. sacchariflorus assemblies.

Firstly, scaffolds longer than 2 kbps from the whole genome assembly "Msac_v2" were scaffolded again using SSPACE⁷ and the *M. sinensis* mate-pairs reads, the gaps between scaffolds were filled in with Ns. This new whole-genome assembly was

denominated "Msac_v3", and was deposited at NCBI in Bioproject PRJNA435476, under the GenBank accession GCA_002993905. It contains 137,916 scaffolds for a total of 2.074 Gb with an N50 of 25.6 Kbps. The gene annotation was projected to the "Msac_v3" assembly using PASA²¹ (version 2.0.1): genes were aligned to the new assembly using GMAP, requiring a minimum identity of 0.85 and coverage of 0.55, and later validated using the default parameters in PASA.

Finally, we obtained the chromosomal position in the *M. sinensis* chromosomes of the scaffolds from the "Msac_v3" assembly. Using Satsuma2²² (version untagged-330e3341a1151a978b37), we identified every perfect-identify match between both assemblies (3,635,504 matches in total). The coordinates of these matches in BED 8 format were used as input to the "OrderOrientBySynteny" script from Satsuma2, which identifies the best chromosomal position for each scaffold. These position coordinates are available as an AGP file as part of

GCA_002993905, which anchors our final whole-genome assembly to 19 chromosomes (accessions CM00959 to CM009609 in NCBI).

Completeness assessment

RNA-seq cleaned reads from each tissue were independently aligned to both assembly versions using STAR²³ (version 2.6.0c). BUSCO²⁴ (version 4.1.4) was used to assess completeness with the single-copy orthologs database for green plants (Viridiplantae, version 2020-09-10). Orthologs were identified using Orthofinder2²⁵ (version 2.3.12) with default parameters and the option "-msa", which directly provided comprehensive statistics comparing the provided proteomes. All the proteomes from the other species used (Table 1) were downloaded from Phytozome (v7.1 DOE-JGI). Genomes were aligned using Minimap2²⁶ (version 2.17) with the "asm10" parameter for related genomes, secondary alignments (tp:A:S) filtered out, and results visualised using dotPlotly²⁷ (Github version, latest updated on 4 May 2018).

Table 1. Completeness statistics of the unanchored and anchored *M. sacchariflorus* whole-genome assemblies in comparison to the *M. sinensis* reference.

	<i>Msac_v2</i> (unanchored)	Msαc_v3 (anchored by M. sinensis)	Reference: <i>M. sinensis</i> ⁴ .	
NCBI bioproject	PRJNA679435	PRJNA435476 (GCA_002993905)	v.7.1 from Phytozome	
Length	2.539 Gb	2.074 Gb	1.68 Gbps	
Scaffolds	588,758 scaffolds	137,931 scaffolds*	19 Chrs and 14,414 scaffolds	
N20	25.39 Kbps	62.61 Kbps	146.1 Mbps	
N50	10.25 Kbps	25.63 Kbps	88.51 Mbps	
N80	2.79 Kbps	9.42 Kbps	75.06 Mbps	
Max	378.48 Kbps	458.83 Kbps	160.9 Mbps	
ANNOTATION	<u>Msac_v2</u>	<u>Msac_v3</u>	<u>M. sinensis</u>	
Gene models	81,431	68,578	67,967	
Proteins	86,767	68,578**	67,789	
BUSCO	<u>Msac_v2</u>	<u>Msac_v3</u>	<u>M. sinensis</u>	
Complete	55.5% (48% in single copy)	59.8% (50.4% in single copy)	97.6% (36.2% in single copy)	
Fragmented	32.2%	26.4%	1.6%	
Missing	12.3%	13.6%	0.8%	
<u>RNA MAPPING</u>	<u>Msac_v2</u>	<u>Msac_v3</u>	<u>M. sinensis</u>	
Reads mapping in the genome once (root, stem and leaf)	76.2% 76.4% 78.8%	75% 76.7% 78.1%	78.8% *** 83.5% 82.5%	
Reads mapping in the genome multiple times (root, stem and leaf)	22.5% 23% 20.7%	19.5% 18.8% 17.3%	19.7% *** 15.5% 16.6%	

*15 scaffolds from plastids were discarded during the deposit in NCBI resulting in 137,916 scaffolds. ** Only the longest transcript was considered in each projected locus. *** Cross-species alignments.

Results

We produced two whole-genome assemblies for *M. sacchariflorus* that we named "Msac_v2" and "Msac_v3", with total lengths of 2.54 Gbps and 2.074 Gbps, respectively (Table 1). The difference in size is mainly a result of filtering 402 Mb from sequences under 2 kb in the latter before anchoring to the *M. sinensis* genome. Our "Msac_v2" assembly covered ~59 % of *M. sacchariflorus* genome size, which is estimated to be 4.3 Gb²⁸. Approximately 40% of the assembly was composed by transposable elements (987.3 Mb; Table 2), including 491 Mb (19.4%) and 154 Mb (6.1%) by copies of the Gypsy and Copia LTRs, respectively; and 180 Mb (7.1%) by several class 2 DNA transposons (MULE, CMC, Harbinger, etc.)

We identified 219,394 primary alignments longer than 2 kb between the unanchored *M. sacchariflorus* ("Msac_v2") and *M. sinensis*. The resulting dotplot (Figure 1) shows the conserved synteny between both species, which diverged 1.6 Mya⁴. Figure 1 also shows the highly conserved synteny between the pairs of homoeologous chromosomes (e.g. green boxes in chromosomes one and two), and the fusion in chromosome 7 of the chromosome homeolog to chromosome 13; which was also reported in *M. sinensis*⁴. There are several large inversions between chromosomes 9 and 10, and 3 and 4 (cyan boxes in Figure 1). Our assembly of a heterozygous

genotype resulted in multiple heterotigs (heterozygous contigs) containing the alternative or secondary haplotypes (e.g. pink boxes in Figure. 1).

The utility of our assemblies for genomic studies is evidenced by the proportion of RNA-seq from three different tissues from the same *M. sacchariflorus* genotype that aligned to the assemblies. On average 99% and 95% of the RNA-seq reads aligned in "Msac_v2" and "Msac_v3", respectively (Table 1).

We estimated that we assembled more than 85% of the *M. sacchariflorus* genes. Furthermore, our assemblies include several alleles of genes in the heterozygous regions of the genome, while the *M. sinensis* reference was generated from a double-haplotyped genotype. The estimation of the proportion of assembled genes (~85%) was supported by (1) the results from BUSCO, which reported 86.4-87.7% of presented core genes, of which ~2/3rds were complete (Table 1); and (2) the difference in the number of proteins from related species for which we can identify an ortholog in *M. sacchariflorus* compared to *M. sinensis*, as control, using Orthofinder2 (Table 3).

Based on the results from Orthofinder2 (Table 3), we found orthologs in M. sacchariftorus for 64.5% of the M.sinensis

Category	Superfamily	Coverage(bp)	Fraction (2.539 Gb)
Class 1 TEs: retrotransposons (copy and paste)	Gypsy LTR	491,915,558	19.37%
	Copia LTR	154,244,411	6.08%
	Other LTRs	87,661,401	3.45%
	SINEs	5,029,476	0.20%
	LINEs	25,076,275	0.99%
	Other non-LTR retrotransposons	29,192,841	1.15%
Class 2 TEs: DNA transposons (cut and paste)	hAT	10,722,644	0.42%
	Harbinger/PIF	24,553,614	0.97%
	MULE/MuDR	29,733,691	1.17%
	Stowaway/TcMar	14,112,359	0.56%
	CMC_EnSpm	56,449,907	2.22%
	Helitron	10,601,152	0.42%
	Other	34,676,501	1.37%
Unclassified TEs	Unclassified TEs	5,934,794	0.23%
	Satellites	5,339,464	0.21%
Non IEs	snRNAs	23,147	0.00%
TOTAL		985,267,235	38.81%

Table 2. Transposable elements identified in the Miscanthus sacchariflorus genome.



Figure 1. Conserved synteny between *M. sacchariflorus* and *M. sinensis* genomes. The plot shows the primary alignments longer than 2 kbps between both species. The *M. sacchariflorus* scaffolds (Y-axis) have been sorted by their coordinates in *M. sinensis* chromosomes (X-axis). Large homoeologous blocks and chromosomal rearrangements are highlighted in boxes.

Orthologs	Msac_v2	Msac_v3	Msin	Sita	Sbic	Zma	Pvi
From Msac_v2 (86,767)	-	NA	44,151 (50.9%)	36,904 (42.5%)	37,219 (42.9%)	38,478 (44.3%)	45,792 (52.8%)
From Msac_v3 (68,578)	NA	-	38,122 (55.6%)	32,273 (47.1%)	32,296 (47.1%)	33,395 (48.7%)	38,755 (56.5%)
From Msin (67,789)	43,739 (64.5%)	37,501 (55%)	-	41,532 (64.1%)	43,475 (64.1%)	39,986 (58.9%)	45,913 (67.7%)
From Sita (40,599)	26,846 (66.1%)	23,559 (58%)	28,473 (70.1%)				
From Sbic (39,441)	27,877 (70.7%)	24,125 (61.2%)	30,907 (78.4%)				
From Zma (88,760)	41,530 (46.8%)	35,955 (40.5%)	41,784 (47.1%)				
From Pvi (125,439)	63,692 (50.8%)	56,120 (44.7%)	64,271 (51.2%)				

Table 3. Number of orthologs between *Miscanthus sinensis* (Msin), *Setaria italica* (Sita; foxtail millet), *Sorghum bicolor* (Sbic; sorghum), *Zea mays* (Zma; maize), and *Panicum virgatum* (Pvi; switchgrass) obtained using Orthofinder 2.

annotated proteins, so we estimated ~1/3rd of the Miscanthus proteins to be specific to each species. On the other hand, we estimated that ~3,000 genes may be missing in the "Msac_v2" annotation based on the number of Sorghum bicolor proteins with orthologues in *M. sinensis* but absent in *M. sacchariflorus*. Better estimations were obtained with the other four species, where the genes absent in Msac_v2 compared with M. sinensis were estimated to be 254, 579 and 1627 (Table 3). Additionally, ~6,000 genes could be missed in "Msac_v3" compared to "Msac_v2" based on the difference in the number of M. sinensis orthologues in each assembly (Table 3). This is likely from genes in the sequences shorter than 2 Kbps (totalling 402 Mbps) that were filtered out before anchoring. There was a large difference in the proportion of "fragmented" BUSCO genes found in the M. sacchariflorus (32.2%) and M. sinensis (1.6%) assemblies (Table 1). To assess if that difference had an effect on the quality of the annotation, we compared the number of proteins from M. sacchariflorus and M. sinensis for which we can identify an ortholog in another species (Table 3); we found the difference between both Miscanthus species ranged between 6,571 proteins when compared to sorghum (43,475 to 37,219; Table 2) to only 121 when compared to maize (39,986 to 38,478, Table 3).

In conclusion, our *M. sacchariflorus* genome can served as the basis for functional genetic analyses on *Miscanthus*, one of the main biofuel grass crops used in temperate latitudes. However, there are opportunities to improve it using new approaches, such as long-reads.

Data availability

Underlying data

NCBI BioProject: Miscanthus sacchariflorus cultivar:Robustus 297. Accession number PRJNA435476; https://identifiers.org/bioproject:PRJNA435476.

This BioProject contains the raw paired-end and mate-pair reads.

NCBI BioProject: RNA-seq Miscanthus hybrids with contrasting phenotypes. Accession number PRJNA639832; https://identifiers. org/bioproject:PRJNA639832.

This BioProject contains RNA-seq reads, deposited as part of a previous project²⁹.

NCBI BioProject: Miscanthus sacchariflorus cultivar:Robustus 297. Accession number PRJNA679435; https://identifiers.org/ bioproject:PRJNA679435.

This Bioproject contains the unanchored "Msac_v2" assemblies and gene annotations under accession JADQCR000000000.

The anchored "Msac_v3" assemblies and gene annotations are deposited under accession accession GCA_002993905 under Bioproject PRJNA435476.

The chromosomal positions in the *M. sinensis* chromosomes of the scaffolds from the "Msac_v3" assembly are available in an AGP file as part of GCA_002993905, which places the scaffolds in 19 chromosomes (accessions CM009591 to CM009609 in NCBI).

Zenodo: Supplementary dataset to "Draft genome assembly of the biofuel grass crop Miscanthus sacchariflorus". http://doi. org/10.5281/zenodo.4270235.

This project contains the assemblies in FASTA format, gene annotations in GFF3 format, functional annotations in tabulated text format, and AGP file with anchoring information.

Data deposited with Zenodo are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

References

- Clark LV, Jin X, Petersen KK, et al.: Population structure of Miscanthus sacchariflorus reveals two major polyploidization events, tetraploidmediated unidirectional introgression from diploid M. sinensis, and diversity centred around the Yellow Sea. Ann Bot. 2019; 124(4): 731-48. PubMed Abstract | Publisher Full Text | Free Full Text
- Dwiyanti MS, Rudolph A, Swaminathan K, et al.: Genetic analysis of putative triploid Miscanthus hybrids and tetraploid M. sacchariflorus collected from sympatric populations of Kushima, Japan. Bioenergy Res. 2013; 6(2): 486–93. Publisher Full Text
- Clark LV, Dzyubenko E, Dzyubenko N, et al.: Ecological characteristics and in situ genetic associations for yield-component traits of wild Miscanthus from eastern Russia. Ann Bot. 2016; 118(5): 941–55. PubMed Abstract | Publisher Full Text | Free Full Text
- Mitros T, Session AM, James BT, et al.: Genome biology of the paleotetraploid perennial biomass crop Miscanthus. Nat Commun. 2020; 11(1): 5442.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Leggett RM, Ramirez-Gonzalez RH, Clavijo BJ, et al.: Sequencing quality assessment tools to enable data-driven informatics for high throughput genomics. Front Genet. 2013; 4: 288.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Simpson JT, Wong K, Jackman SD, et al.: ABySS: a parallel assembler for short read sequence data. Genome Res. 2009; 19(6): 1117–23.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Boetzer M, Henkel CV, Jansen HJ, et al.: Scaffolding pre-assembled contigs using SSPACE. Bioinformatics. 2011; 27(4): 578–9.
 PubMed Abstract | Publisher Full Text
- De Vega JJ, Ayling S, Hegarty M, et al.: Red clover (*Trifolium pratense* L.) draft genome provides a platform for trait improvement. Sci Rep. 2015; 5: 17394. PubMed Abstract | Publisher Full Text | Free Full Text
- Stanke M, Keller O, Gunduz I, et al.: AUGUSTUS: ab initio prediction of alternative transcripts. Nucleic Acids Res. 2006; 34(Web Server issue): W435–W9. PubMed Abstract | Publisher Full Text | Free Full Text
- Tarailo-Graovac M, Chen N: Using RepeatMasker to identify repetitive elements in genomic sequences. *Curr Protoc Bioinformatics*. 2009; 25(1): Chapter 4: Unit 4.10.
 PubMed Abstract | Publisher Full Text
- Jurka J, Kapitonov VV, Pavlicek A, et al.: Repbase Update, a database of eukaryotic repetitive elements. Cytogenet Genome Res. 2005; 110(1–4): 462–7. PubMed Abstract | Publisher Full Text
- 12. Smit AF, Hubley R: RepeatModeler Open-1.0. 2008. Reference Source
- Trapnell C, Pachter L, Salzberg SL: TopHat: discovering splice junctions with RNA-Seq. Bioinformatics. 2009; 25(9): 1105–11.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Grabherr MG, Haas BJ, Yassour M, et al.: Trinity: reconstructing a full-length transcriptome without a genome from RNA-Seq data. Nat Biotechnol. 2011; 29(7): 644–52.
 - PubMed Abstract | Publisher Full Text | Free Full Text
- 15. Trapnell C, Roberts A, Goff L, et al.: Differential gene and transcript expression

analysis of RNA-seq experiments with TopHat and Cufflinks. *Nat Protoc.* 2012; 7(3): 562–78. PubMed Abstract | Publisher Full Text | Free Full Text

- Bromberg Y, Rost B: SNAP: predict effect of non-synonymous polymorphisms on function. Nucleic Acids Res. 2007; 35(11): 3823–35. PubMed Abstract | Publisher Full Text | Free Full Text
- Blanco E, Parra G, Guigó R: Using geneid to identify genes. Curr Protoc Bioinformatics. 2007; Chapter 4: Unit 4.3.
 PubMed Abstract | Publisher Full Text
- Slater GSC, Birney E: Automated generation of heuristics for biological sequence comparison. BMC Bioinformatics. 2005; 6(1): 31.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Conesa A, Götz S, García-Gómez JM, et al.: Blast2GO: a universal tool for annotation, visualization and analysis in functional genomics research. *Bioinformatics.* 2005; 21(18): 3674–6.
 PubMed Abstract | Publisher Full Text
- Huerta-Cepas J, Forslund K, Coelho LP, *et al.*: Fast Genome-Wide Functional Annotation through Orthology Assignment by eggNOG-Mapper. *Mol Biol Evol*, 2017; 34(8): 2115–22.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Haas BJ, Delcher AL, Mount SM, et al.: Improving the Arabidopsis genome annotation using maximal transcript alignment assemblies. Nucleic Acids Res. 2003; 31(19): 5654–66.
- PubMed Abstract | Publisher Full Text | Free Full Text 22. Clavijo B, Wright J, Yanes L.
- Reference Source
- Dobin A, Davis CA, Schlesinger F, et al.: STAR: ultrafast universal RNA-seq aligner. Bioinformatics. 2013; 29(1): 15–21.
 PubMed Abstract | Publisher Full Text | Free Full Text
- Simão FA, Waterhouse RM, Ioannidis P, et al.: BUSCO: assessing genome assembly and annotation completeness with single-copy orthologs. Bioinformatics. 2015; 31(19): 3210–2.
 PubMed Abstract | Publisher Full Text
- 25. Emms DM, Kelly S: OrthoFinder: phylogenetic orthology inference for comparative genomics. *Genome Biol.* 2019; 20(1): 238. PubMed Abstract | Publisher Full Text | Free Full Text
- Li H: Minimap2: pairwise alignment for nucleotide sequences. Bioinformatics. 2018; 34(18): 3094–100.
 PubMed Abstract | Publisher Full Text | Free Full Text
- PubMed Abstract | Publisher Full Text | Free Full Text

 27.
 Poorten T.
- Reference Source
- Rayburn AL, Crawford J, Rayburn CM, et al.: Genome Size of Three Miscanthus Species. Plant Mol Biol Report. 2009; 27(2): 184.
 Publisher Full Text
- De Vega JJ, Peel N, Purdy SJ, et al.: Differential expression of starch and sucrose metabolic genes linked to varying biomass yield in *Miscanthus* hybrids. *BioRxiv*. 2020; 2020–08.
 Publisher Full Text

Open Peer Review

Current Peer Review Status:

Version 1

Reviewer Report 01 March 2021

https://doi.org/10.5256/f1000research.47786.r78040

© **2021 Dwiyanti M.** This is an open access peer review report distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.



Maria Stefanie Dwiyanti 匝

Research Faculty of Agriculture, Hokkaido University, Sapporo, Japan

The availability of *Miscanthus sacchariflorus* genome sequence will be useful for *Miscanthus* related research, particularly in bioenergy related topics.

"Better estimations were obtained with the other four species, where the genes absent in Msac_v2 compared with M. sinensis were estimated to be 254, 579 and 1627 (Table 3)."

- I found that the difference between number of genes in "*Msac_v2*" compared to other four species is larger than 254, 579, and 1627; or the way I look into the table is wrong?
- Perhaps the sentence above can be reworded so we can easily compare with the Table 3 content.
- Also, what are the predicted functions of genes absent in "Msac_v2" compared to M.sinensis?
 - This information may provide some clues to trait difference between *M. sacchariflorus* and *M.sinensis*.

Is the rationale for creating the dataset(s) clearly described?

Yes

Are the protocols appropriate and is the work technically sound?

Yes

Are sufficient details of methods and materials provided to allow replication by others? Yes

165

Are the datasets clearly presented in a useable and accessible format?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Plant genetics and genomics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 25 February 2021

https://doi.org/10.5256/f1000research.47786.r77622

© **2021 Riaño-Pachón D.** This is an open access peer review report distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.



Diego Mauricio Riaño-Pachón 匝

Computational, Evolutionary and Systems Biology Laboratory, Center for Nuclear Energy in Agriculture (CENA), University of São Paulo, Piracicaba, Brazil

The Data Note, "Draft genome assembly of the biofuel grass crop *Miscanthus sacchariflorus*", introduces two *Miscanthus sacchariflorus* genome assemblies, which have been deposited in NCBI under the bioprojects: PRJNA679435 and PRJNA435476. Genome sequencing was carried out with Illumina paired end reads and mate-pairs, the assemblies are greatly fragmented, which is expected due to the sequencing technologies used. This is the first *Miscanthus sacchariflorus* genome assembly, which is of interested for the bioenergy community, and can be used to generate insigths with the genomes of other bioenergy crops, like sorghum and sugarcane.

Suggestions:

- Look for contaminant organisms in the final assemblies using BlobPlots.
- Provide GenomeScope and Smudgeplots for the clean reads, to generate further statistics prior to assembly.

Is the rationale for creating the dataset(s) clearly described?

Yes

Are the protocols appropriate and is the work technically sound?

Yes

Are sufficient details of methods and materials provided to allow replication by others? $\ensuremath{\mathbb{No}}$

Are the datasets clearly presented in a useable and accessible format?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Bioinformatics, genome assembly and annotation.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

The benefits of publishing with F1000Research:

- Your article is published within days, with no editorial bias
- You can publish traditional articles, null/negative results, case reports, data notes and more
- The peer review process is transparent and collaborative
- Your article is indexed in PubMed after passing peer review
- Dedicated customer support at every stage

For pre-submission enquiries, contact research@f1000.com

F1000 Research