Building Pangene Sets from Plant Genome Alignments Confirms Presence- Absence Variation

Bruno Contreras-Moreira, Shradha Saraf, Guy Naamati, Sandeep S. Amberkar, Paul Flicek, Andrew R. Jones, Sarah Dyer



European Molecular Biology Laboratory-EBI Estación Experimental de Aula Dei-CSIC





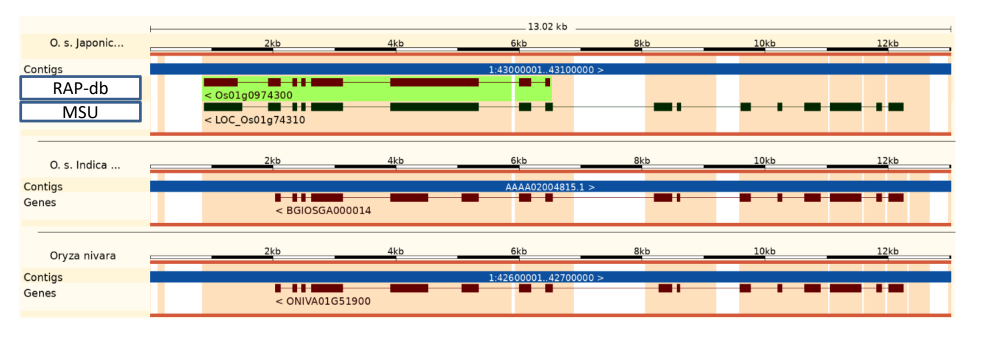




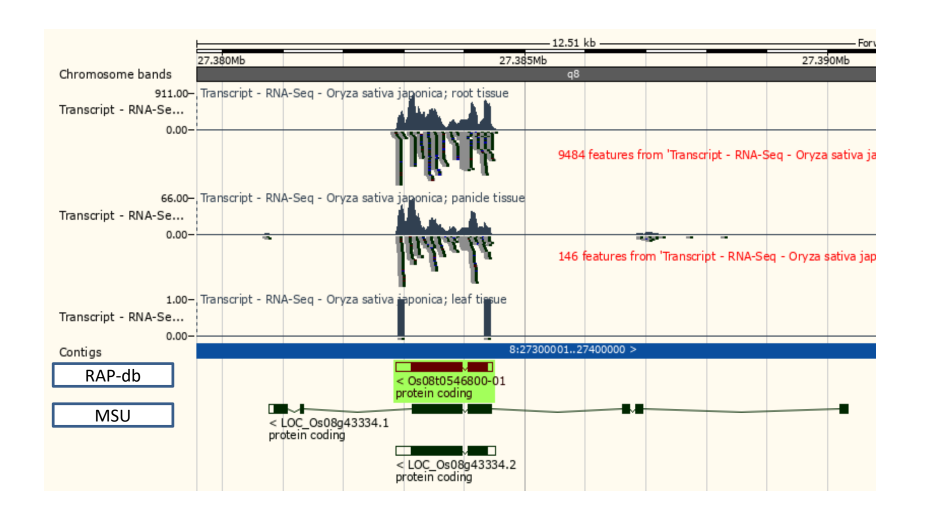


Problem: unconsistent gene annotation

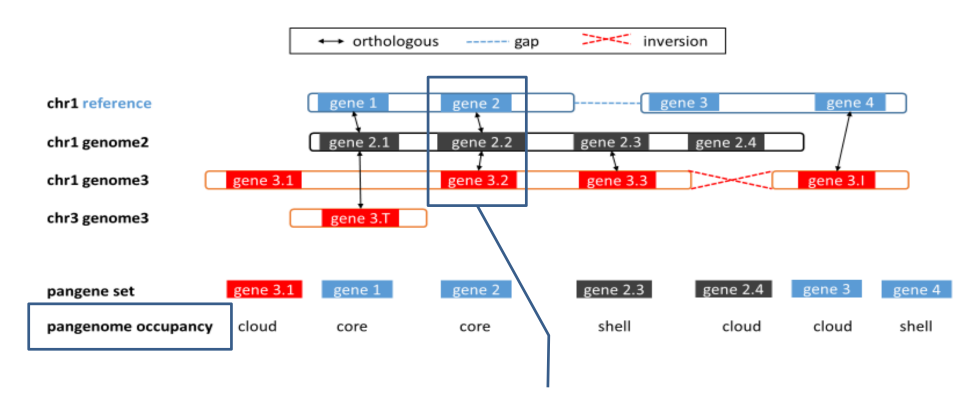
	Gene models MSU (2011)	Gene models RAP-db	
Oryza sativa	55.801	37.859	4.168



Problem: multiple isoforms, are they all valid and relevant?

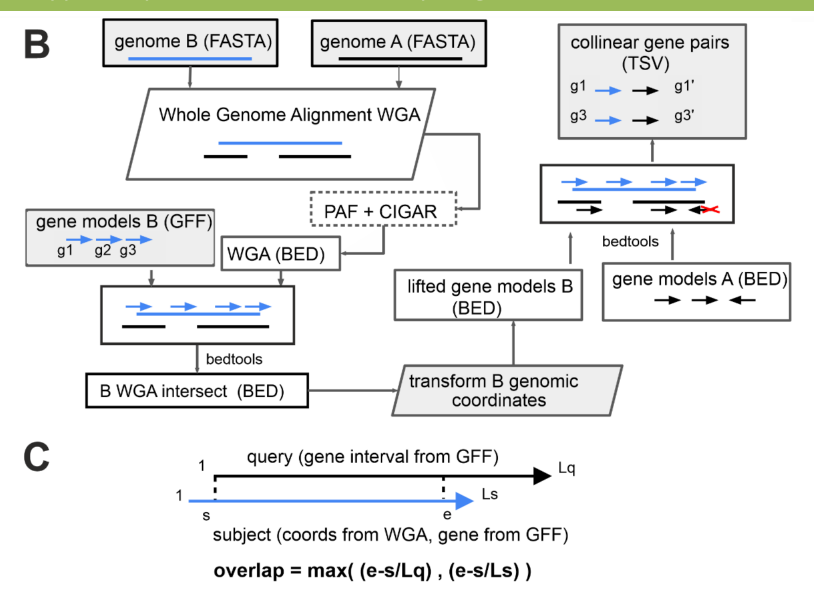


Goal: definition and nomenclature of pangenes

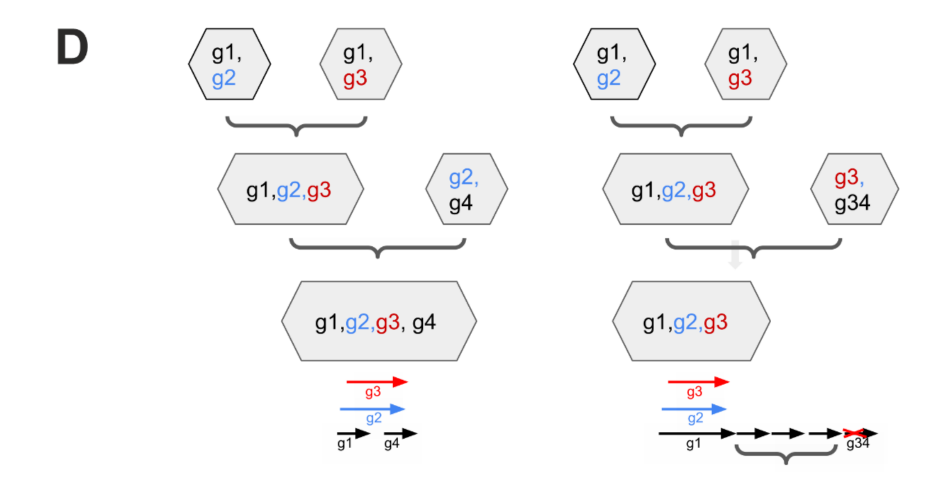


pangene = cluster of overlapping genomic regions containing gene models

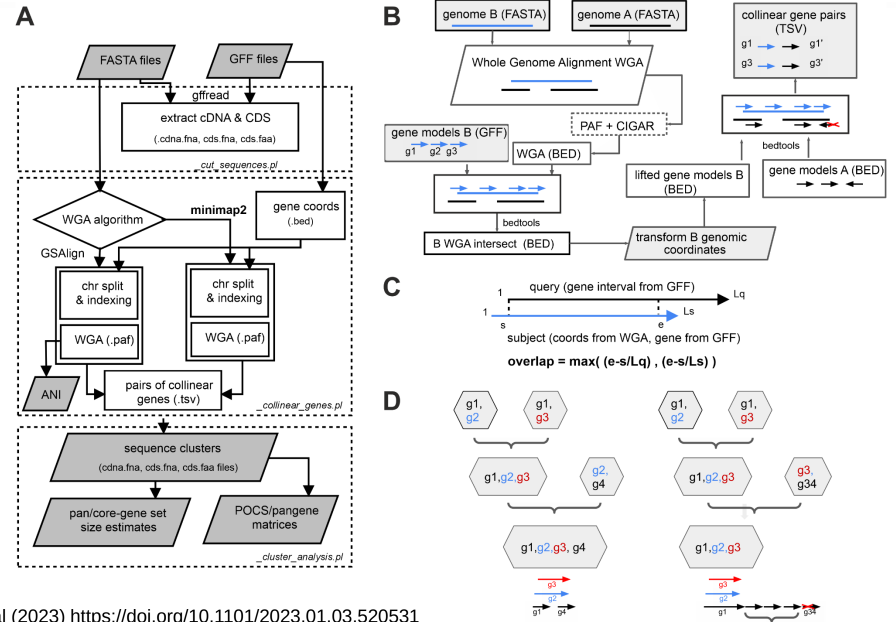
A prototype to produce clusters of pangenes



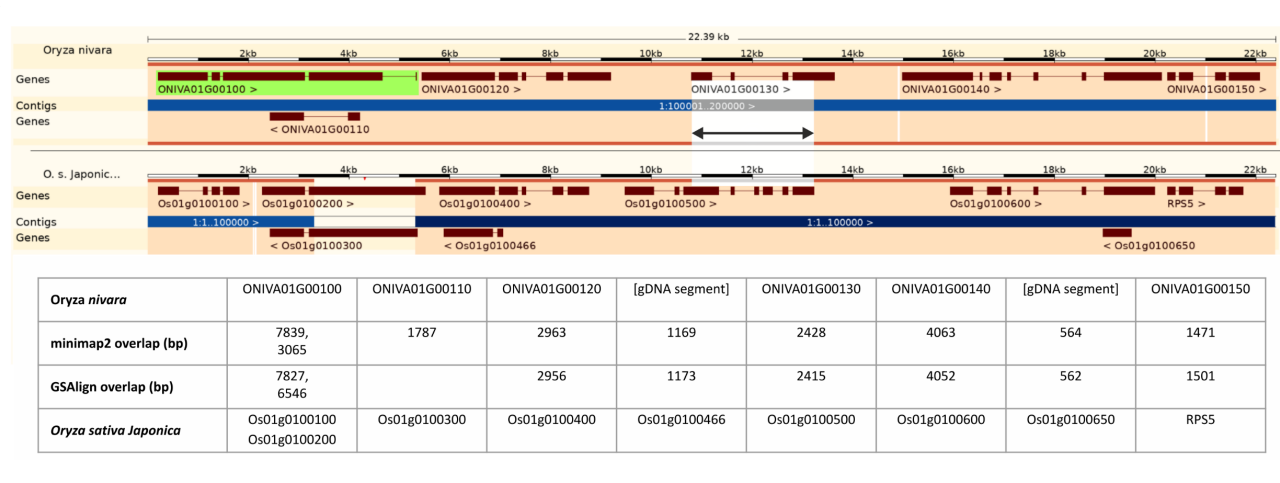
A prototype to produce clusters of pangenes



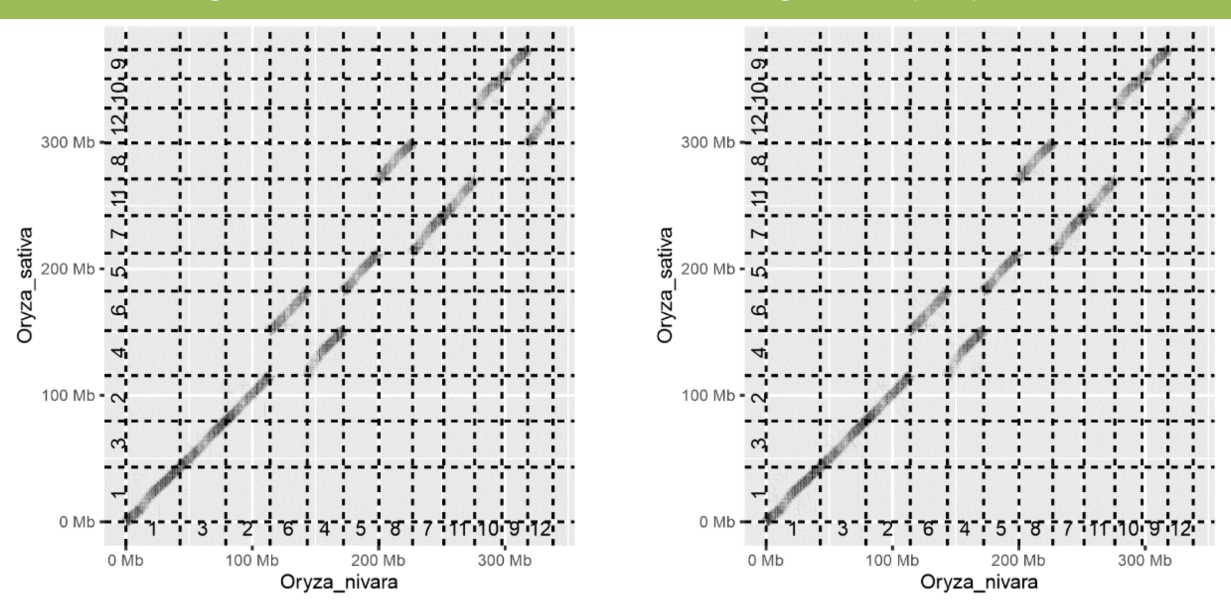
A prototype to produce clusters of pangenes



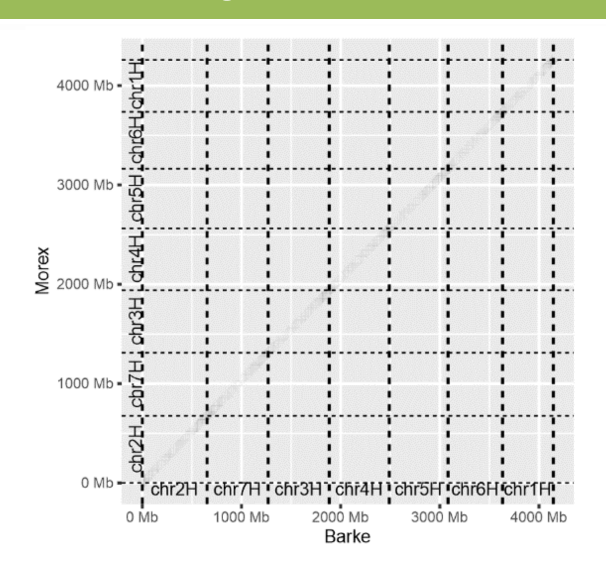
Region in chr1 of Oryza nivara (top) and Oryza sativa (bottom) in Ensembl

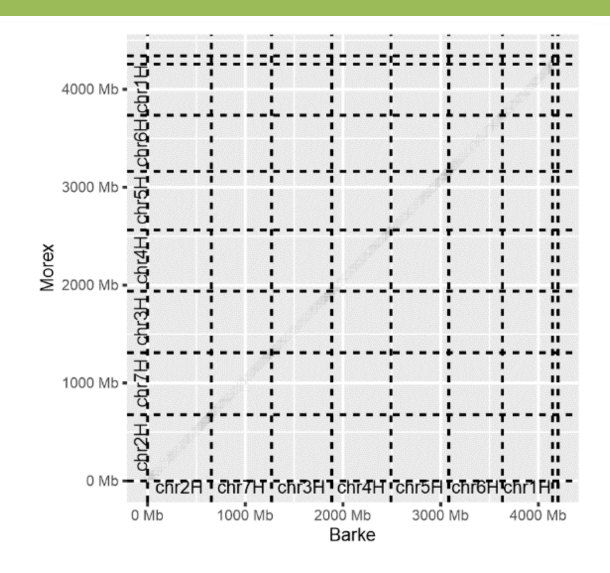


Pangenes are collinear in Whole Genome Alignments (rice)

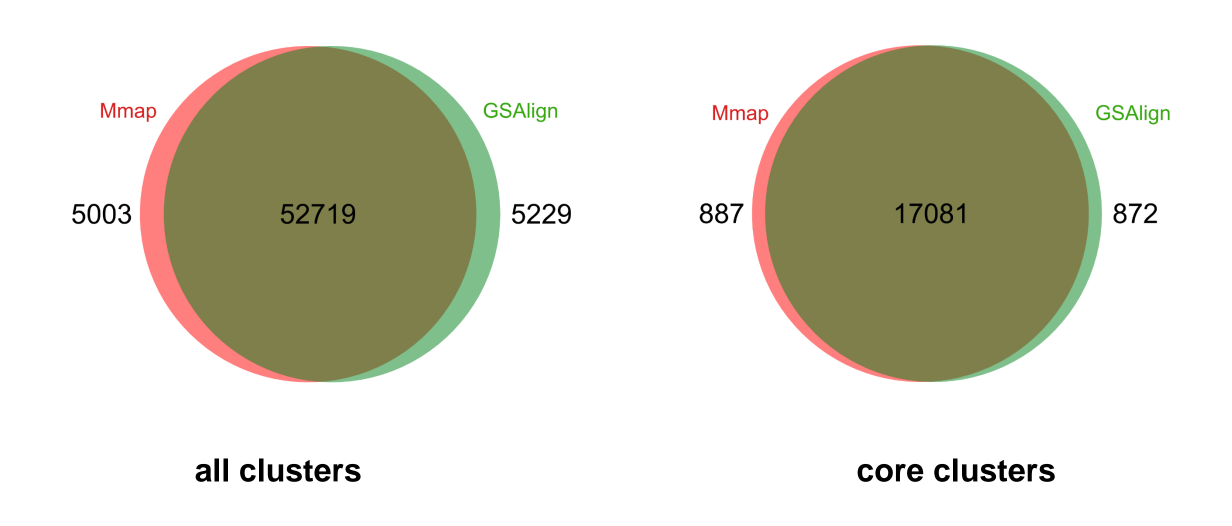


Pangenes are collinear in Whole Genome Alignments (barley)





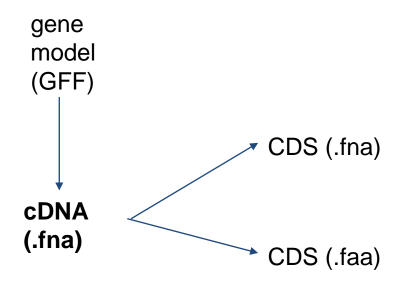
Minimap2 and GSAlign produce similar pangenes (rice3)



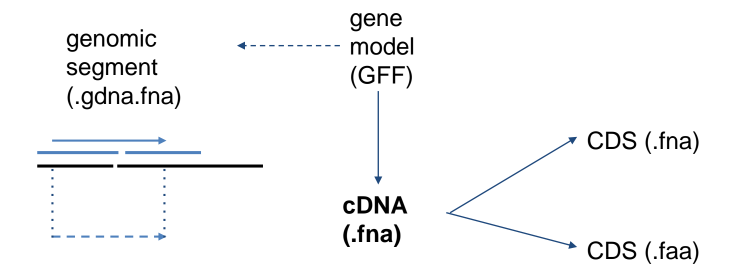
How do pangene clusters look like? GIGANTEA Os01g0182600, 3'



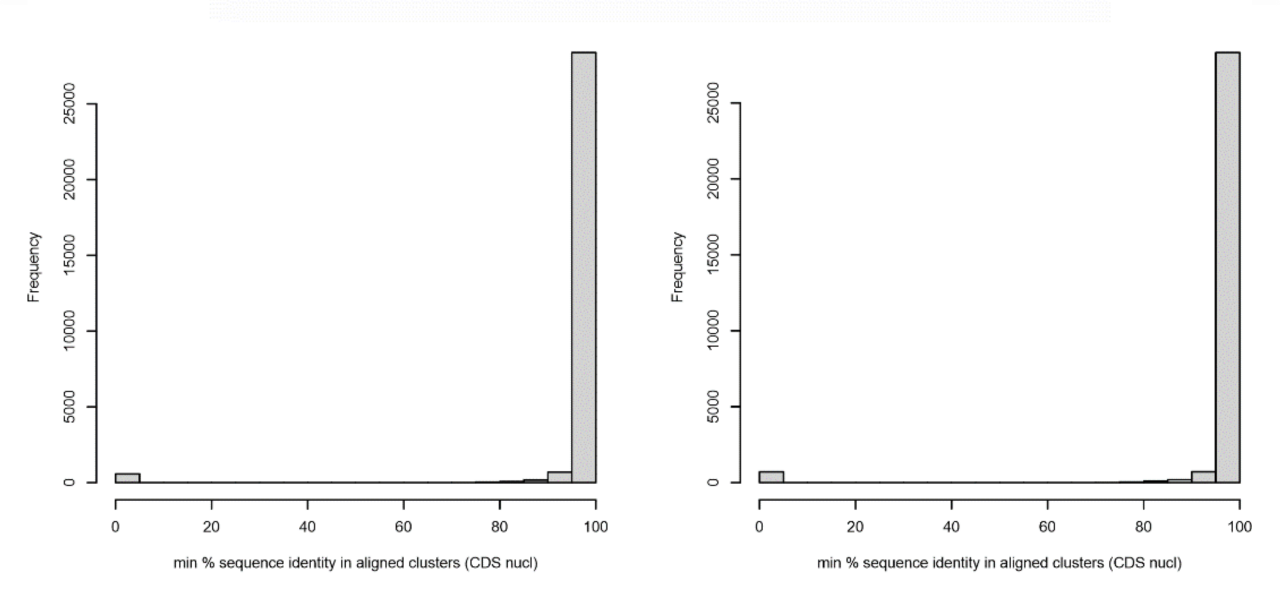
Different versions of pangene clusters: cDNA, CDS nucl, CDS pep



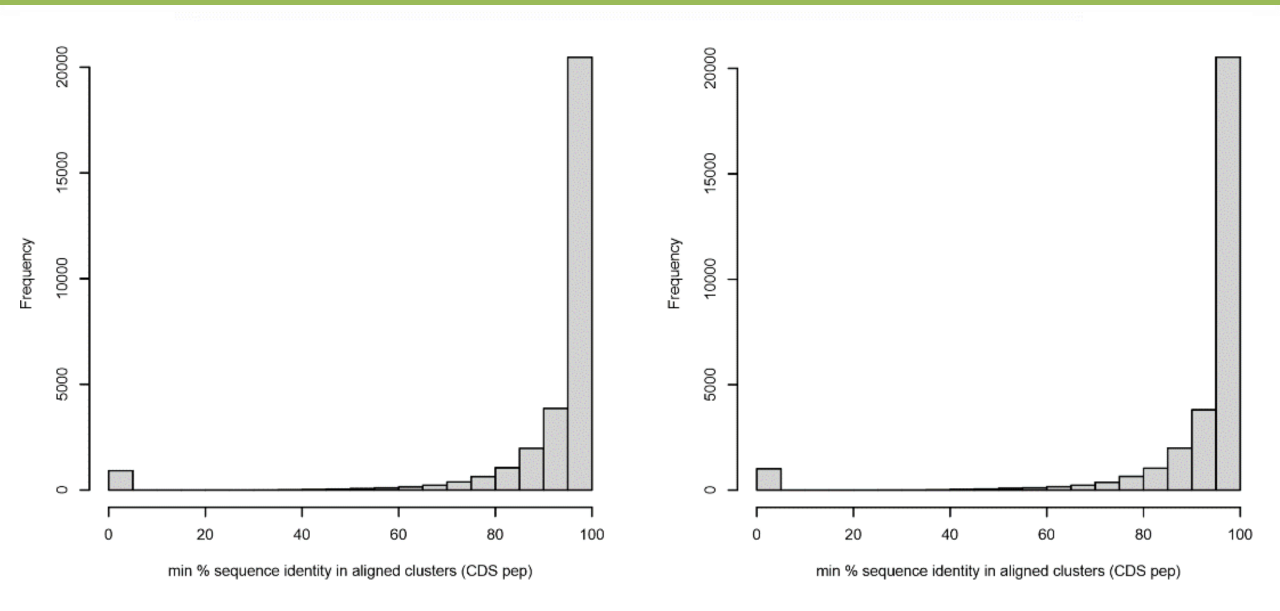
Different versions of pangene clusters: cDNA, CDS nucl, CDS pep + gDNA



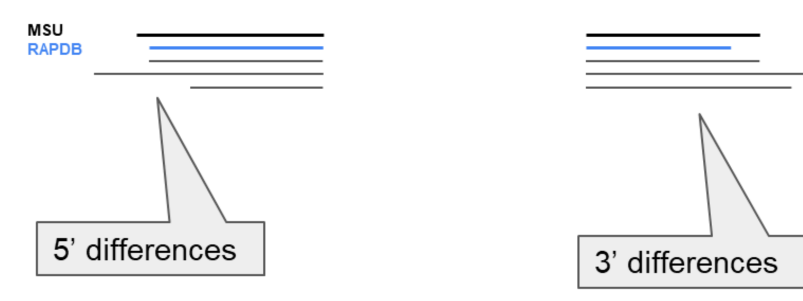
Minimap2 & GSAlign make pangene clusters with high sequence identity (nucl)

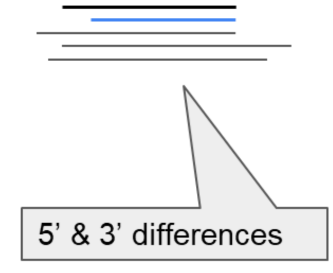


Minimap2 & GSAlign make pangene clusters with high sequence identity (pep)



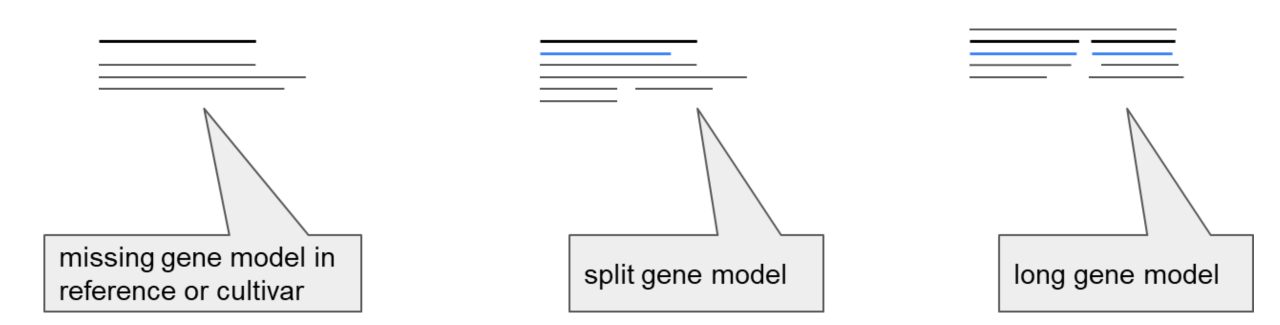
Apps: pangene clusters provide evidence for gene model annotation (rice)





benchmark set	isoforms	in pangenes	match cluster mode
Oryza sativa genes curated in RAP-db	3895	2937	1700 (57.8%)
Oryza proteins curated in SwisProt	5685	3876	1650 (42.5%)

Apps: pangene clusters provide evidence to fix annotation errors (rice)



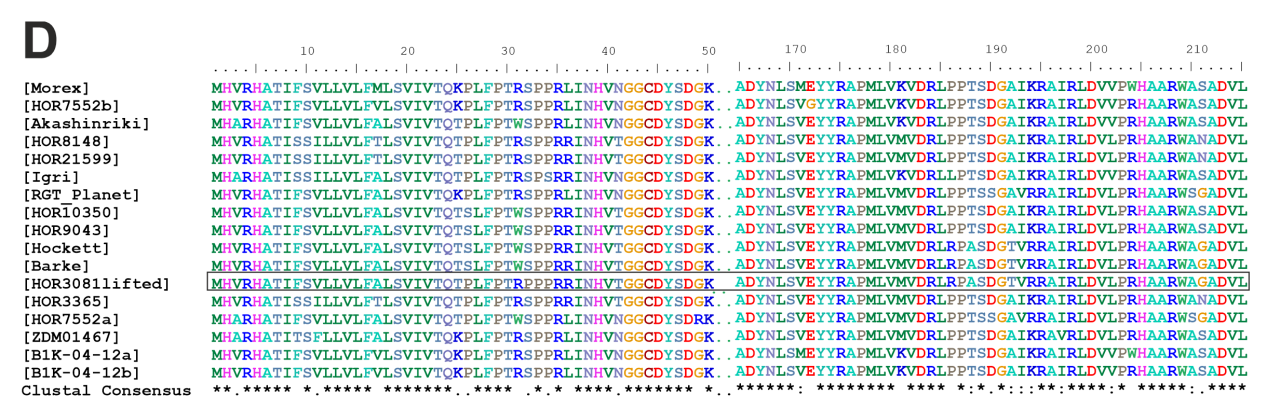
Example: validating PAV in barley

,					
	length	pairs	overlap	genename	taxon
	892	14	12005	Horvu_AKASHIN_1H01G016200	Akashinriki
	898	13	11167	Horvu_21599_1H01G015900	HOR21599
	904	13	11258	Horvu_10350_1H01G020000	HOR10350
	892	11	9519	Horvu_PLANET_1H01G014600	RGT_Planet
	733	11	8040	Horvu_BARKE_1H01G018700	Barke
	904	11	9434	Horvu_9043_1H01G017700	HOR9043
	1380	10	8989	Horvu_HHOR_1H01G019100	HOR3365
	898	10	8152	Horvu_8148_1H01G016700	HOR8148
	733	9	6558	Horvu_HOCKETT_1H01G013500	Hockett
	771	8	6166	Horvu_MOREX_1H01G020700	Morex
	904	7	5961	Horvu_IGRI_1H01G016300	Igri
	897	7	5962	Horvu_7552_1H01G018700	HOR7552
	1444	6	5769	Horvu_HUANG_1H01G010400	ZDM01467
	1374	6	5344	Horvu_7552_1H01G018300	LIODZEES
	1380	2	2247	Horvu_FT11_1H01G019600	# missing
	1380	2	1661	Horvu_FT11_1H01G019300	# source=

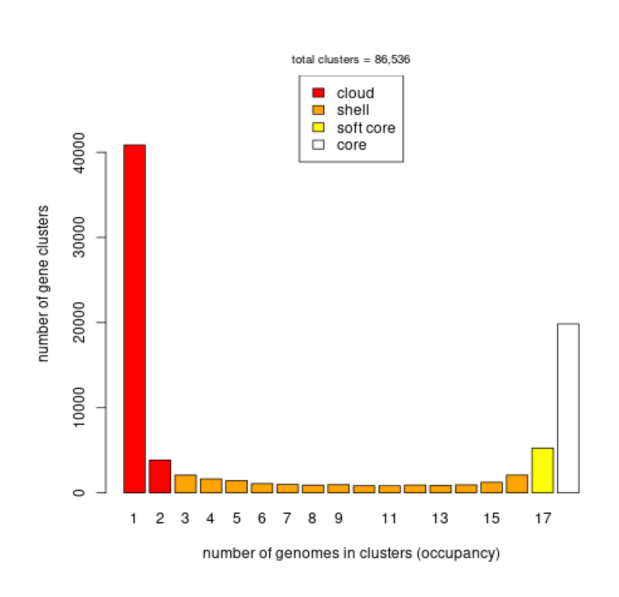
GMAP lift-over
Genomic segment

ng gene model: corrected chr1H:3798001-3798892(-) [HOR3081] ce=Hockett matches=639 mismatches=9 indels=0 chr1H 3798173 3798892 gmap gene chr1H mRNA gmap 3798173 3798892 chr1H gmap 3798591 3798892 97 exon chr1H gmap 3798173 3798518 100 exon chr1H gmap CDS 3798591 3798892 97 chr1H gmap CDS 3798173 3798518 100

Example: validating PAV in barley



Apps: pangene clusters capture pangenome dynamics (rice)



softcore (all isoforms)	96.9
oryza_sativa_RAPDB.cds	85.3
oryza_sativa_MSU.cds	94.7
oryza_sativa_arc.cds	94.7
oryza_sativa_azucena.cds	95.3
oryza_sativa_chaomeo.cds	95.4
oryza_sativa_gobolsailbalam.cds	94.4
oryza_sativa_ir64.cds	94.8
oryza_sativa_ketannangka.cds	95.0
oryza_sativa_khaoyaiguang.cds	95.2
oryza_sativa_larhamugad.cds	94.1
oryza_sativa_lima.cds	94.5
oryza_sativa_liuxu.cds	94.7
oryza_sativa_natelboro.cds	93.4
oryza_indica.cds	95.1
oryza_sativa_ZS97.cds	94.5
oryza_sativa_n22.cds	95.3
oryza_sativa_mh63.cds	94.5

% BUSCO complete

TO BE DONE

 PanOryza project: define rice pangenes that will be curated in UniProt in collaboration with the rice community

- Create rules to name pangenes:
 - stable and consistent
 - support future addition of new annotation sets

For you: try it out at https://github.com/Ensembl/plant-scripts

Building Pangene Sets from Plant Genome Alignments Confirms Presence-Absence Variation

Bruno Contreras-Moreira, Shradha Saraf, Guy Naamati, Sandeep S. Amberkar, Paul Flicek, Andrew R. Jones, Sarah Dyer

@BrunoContrerasM











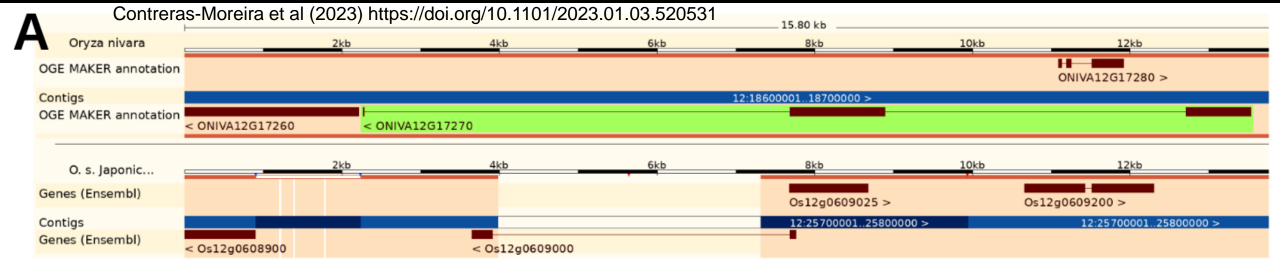


EXTRA SLIDES

	max RAM (GB)	WGA N50 (Kbp)	% genes blocks3+	total clusters	(soft) core clusters	% BUSCO complete
ACK2	4.5	6.1	34.3	43,951	15,768	74.3
rice3	1.4	[27.4, 29]	[74.4, 78.4]	62,915	18,681	83.9
chr1wheat10 (- H)	64.5	[80.8, 142.4]	[38.8, 54.2]	30,173	7,872	
barley20 (-H)	46.3	[43.6, 75.7]	[24.9, 35.4]	180,667	15,674 (23,888) {34,074}	

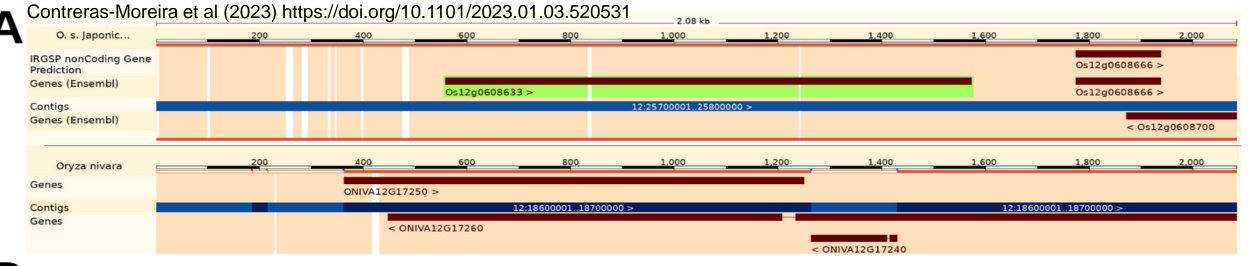
	max RAM (GB)	WGA N50 (Kbp)	% genes blocks3+	total clusters	(soft) core clusters	% BUSCO complete	% ANI
ACK2	4.5	4.3	23.4	43,340	16,432	74.7	84.7
rice3	3.3	[15.2, 16.9]	[51.6, 56.6]	62,844	18,626	84.2	[96.4, 9 7.6]
chr1wheat10	83.4	[40.9, 72.1]	[20.2, 34.2]	30,135	7,723		[98.9, 9 9.4]
barley20	113.1	[17.1, 34.3]	[10.5, 15.9]	173,984	13,934 (21,171)		, , , , , , , , , , , , , , , , , , ,
barley20 -H	110.1	[16.6, 32.9]	[10.5, 15.9]	188,289	13,957		[96.8, 9 9.3]

	dataset	core clusters	multiple copies	shell clusters	gDNA segments	match Compara	share InterPro domains
Compara orthogroups	ACK2	20,192	161				[18,259]
minimap2 clusters	ACK2	15,768	490			14,044	[14,371]
GSAlign clusters	ACK2	16,432	446			14,145	[14,790]
Compara orthogroups	rice3	13,020	219	6,386			16,766 [11,571]
minimap2 clusters	rice3	20,419	3,022	9,503	5,593	17,317	22,796 [17,232]
GSAlign clusters	rice3	20,224	2,831	9.863	6,173	17,103	22,818 [16,957]



>transcript:ONIVA12G17270.1 gene:ONIVA12G17270 12:18626386-18637647(-) [Oryza_nivara]
MQLLFFSFLFLLLARETSAVAADGCSRRCGGLVVPYPFGFSGSCPIMLSCNVDGGSNSTAALILQGNDAT
TTDRSYTVVDGSFNSTASTFTVSVPPSCNRTVSDARRWLSGANYGVSSRTGLFLRGCRNATSSDCSVPVE
TMLRTTRCSGGGGNETASSSLTCIASLSPATPAERGLGGLFAQWEKVEEPRCENLLTSVYGDTREGVFSL
EFAAAEMRWWVNGSCGGGVDDLGRCAANATCIPMQTPSGNWGHRCECLPWMAGDGFAAGEGCYAGKRRRM
RVVEFATAGSVAFLLCLALSVWCLLRRRQWRRNNAKLTVKMARKHLPKDARFFRGKPIEDELELEAAGPR
RFHYGELAAATANFSDDRRLGSGGFGSVYRGFLNGGDVAVKRVAETSRQGWKEFVAEVRIISRLRHRNLV
PLVGWCHDGGDELLLVYELMPNGSLDAHIHSSGNVLPWPARYEVVLGVGAALMYLHHEAEQRVVHRDIKP
SNVMLDASFSARLGDFGLARLIDDGRRSRTTGIAGTMGYIDAECFLLAGRASVESDVYSFGVVLLEVACG
RRPAVVINGGEDAIHLTQWVWDTHGGAAGGGILDAADTRLNGEFDVAEMERVLAVGLWCAHPDRGLRPSI
ROAVSVLRFEAPLPSLPVRMPVATYGPPVSTASAPTSNDTSAGRLHP

>transcript:0s12t0609000-00 gene:0s12g0609000 12:25722111-25722886(-) [Oryza_sativa] MPVATYGPPVSTASAPTSNDTSAGRDSATRTVKSEDPLPPRLYARQGQLDSHLFPLAFIEPPFVEHLACM PIELAIAICLALHLVRRRAPPACACHPLAVLAFLSPWRPSA



>transcript:ONIVA12G17250.1 gene:ONIVA12G17250 12:18611207-18616986(+) [Oryza_nivara] CTTCGCTTGGAATATACAACAAGATGGGATAGTTTATGATGGAAAGGGACTATTTAGCGCTGCTCGTGTTTTGGTGGATGTTTTCGGACATGATAAAGTTTGGGTGTTTGGATGGTTTTGCCCCAATG CATCTTATTTGGAATCTTGACCAGGTAAAAGAGAACATTGATGCCAAGACACATCAACTTATAGATGCTCGAGGAAAGCCTAGATTTGATGGTGCAGTTCCAGAGCCACGGAAAGGAATAAGAAGTGGGCATGTGCCTG AGCATGTATATTAGCTCTGGGCCTCCACCGCCTCGGCAAAACCGATGTTCCTGTATATGATGGATCATGGACTGAATGGGGGAGCCCATCCTGACACTCCAGTTGCCACTGCTGATTAGTACCAGTTACAATCTTT CCACCGGCATCCTCGCCGGGAGGCTCGGCGCGCGCCCCTCGCCCCTCAGCACGCCGATGGCCTGATCGTCGGCCGCAGGCTCCGGTCAGGGTGCGCGCACCAGAGCCCGACGACCATGACGGCCTCCATCTCCCC GCCGTCGAACTCGCCGGTGAGCCGCCGGTCAGCGCGTCAAGAATCCTCCCACTGCCGTACAAATTCCAGACCCACTGAGCAATGTGGATCCGGTCTTCGTCTACCTCGGATTGGTGGTGGTGGTCCCCCATTATTGGT CGCCGGCCGCAGGCGATCTCAAGGAGGACGCCGAAGCTGTACACGTCGGACTCGGCGTTGGCCCTACCGGTGATCATGCACTCCGGGTCCATGTACCCCCATCGTGCCGCGAGCACCCGTGGTGTGAGCCCCGGC CATGGTCGACGAGCCTGGCGAGGCCGAAGTCGCCGAGCTTGGCGTTGAAGGCGGCGTCGAGCATGATGTTGCTCGGCTTGATGTCGCGGTGCACCACCGCACTGCTCCCACTCCTCGTGCAGGTACAGCAGCGCCGAGCC