Assembly Report

Experiment Details

ExperimentID:	Experiment 69	
Sequencing Coverage:	23	
Number of Partitions:	4	
Allowed Mismatch Count in (Base):	0	
Length of Sliding Window:	51	

Reads Dataset Details

Iteaus Dataset Details	
Total Reads Count:	1212
Total Dataset Size in (Base):	241272
Valid Read Count:	1212
Rejected Read Count:	0
Maximum Read Length:	201
Minimum Read Length:	162

Reference Genome Details

Length of Reference Genome (Base):	10080
Number of K-mers:	9880
Length of K-mer	426

Assembly Details

Number of Contigs:	19
Contig N50:	1026
Contig N90:	916
Number of Scaffold	1
Scaffold N50:	10080
Scaffold N90:	10080
Mis-assembly Count:	0
TotalAssembly Size:	10080

Repeat Details

Total Reads Count(Non-Repeat) :	99
Total Reads Count(Repeat):	289
Retetitve Read Count based on (Partitions Identifier):	288
Retetitve Read Count (Entire Read Frequency Identifier):	4
Retetitve Read Count (Contained Read Frequency Identifier):	3
Total Repetitive Sequences Count	3506
Total Unique Repetitive Sequences Count	5

Total Repeat Size (Base)		8400					
Starting	Ending	Repetitive Sequences	Repeat Count Length Positi			tions	
1	10080	CGGTCGAAAA ACTGCTGGCA GTGGGGCATT ACCTCGAATCT ACCGTCGATAT TGCTGAGTCC ACCCGCCGTA TTGCGGCAAG TCGTATTCCGG CTGATCACATG GTGCTGATGG CAGGTTTCACC GCCGGTAATG AAAAAGGCGA ACTGGTGGTG CTGCTGCTGCC CGGTGCTGGC TGCCTGTTTAC GCGCCGATTG TGCGAGATTT GGACGCATTG TTGCGAGATTT GGACGCATTG TTGCGAGATTT GGACGCATTG TTGCGAGATTT GGACGCATTG TTGCGAGATTT GGACGCGTCAG GTGCCCGATTG TTGCGAGATTT GGACGCGTCAG GTGCCCGATG CCGCGTCAG GTGCCCGATG TGCCCGATG TCCTACCAGG AAGCGATGGA GCTTTCCTACT TCGGCGCTCT AGGTCAGGCC	8	350	Start Positio n 1756 3561 3964 5117 5910 7138 8386 9149	End Positio n 2105 3910 4313 5466 6259 7487 8735 9498	

	1		1			1
		GATTCATTCGG				
		GATGGTCTGT				
		GTGGATTAAAA				
		AAAGAGTGTCT				
		GATAGCAGCTT				
		CTGAACTGGTT				
		ACCTGCCGTG				
		AGTAAATTAAA				
		ATTTTATTGAC				
		TTAGGTCACTA			C4 = "4	□ al
		AATACTTTAAC			Start	End
		CAATATAGGCA				Positio
		TAGCGCACAG			n	n
		ACAGATAAAAA			409	758
		TTACAGAGTAC			2228	2577
		ACAACATCCTC	8 350	350		
1		AAAGCCTACC			3182	3531
		GGTGACAGTG			4730	5079
		CGGGCTTTTTT			5529	5878
		TTCGACCAAAG				
		GTAACGAGGT			6730	7079
		AACAACCATGC			8014	8363
		GAGTGTTGAA			9552	9901
		GTCAGGAGAT			9332	19901
		CCTAAAGGCC				
		TGTACCCGTTA				
		CCTAGCCAGTT				
		GGCATTAAAC				
		GTATACGGTAC				
		CTAGGCATGTA				
		CGTAATCGTAG				
		CCTTAGCAATC				
		TCCAGTCC				

ī						
		GCTTTTCATTC				
		TGACTGCAAC				
		GGGCAATATG				
		TCTCTGTGTGG				
		ATTAAAAAAAAG				
		AGTGTCTGATA				
		GCAGCTTCTG				
		AACTGGTTACC				
		TGCCGTGAGT				
		AAATTAAAATT			04 = =4	
		TTATTGACTTA			Start	End
		GGTCACTAAAT			Positio	Positio
		ACTTTAACCAA			n	n
		TATAGGCATAG			1	350
		CGCACAGACA			854	1203
		GATAAAAATTA				
1	10080	CAGAGTACAC	8	350	1295	1644
		AACATCCATGA		43 63 75	2705	3054
		AACGCATTAGC			4337	4686
		ACCACCATTAC				
		CACCACCATCA			6359	6708
		CCATTACCACA			7574	7923
		GGTAACGGTG			8752	9101
		CGGGCTGACG			0.02	0.0.
		CGTACAGGAA				
		ACACAGAAAAA				
		AGCCCGCACC				
		TGACAGTGCG				
		GGCTTTTTTTT				
		CGACCAAAGG				
		TAACGAGGTAA				
		CAACCATGCG				
		AGTGTTGAAGT				

SRGD Performance Metrics

Initial Reads Count:	1212
Reads Count after Removing Duplication:	388

Reads Count after Overlapping:		331				
Initial Dataset Size in (Base):		241272				
Dataset Sizet after	Removing Duplication	n:	99165			
Dataset Size after 0	Overlapping:		67920			
Overlapping Metri	CS					
Repeat Identificatio	n Time:		00h:00m	n:06s:000ms		
Overlapping Time:			00h:05m	n:48s:000ms		
Reads Alignment T	ime:		00h:02m:48s:000ms			
Total Hybrid Assem	Total Hybrid Assembly Time:		00h:08m:36s:000ms			
Repeat Annotation Time:		00h:40m:56s:000ms				
Overlapping Length	O(N)2 Time Complexity	Hit Index (Overlapping Matched Count RI		RI	
197	9801	11		1	0	
194	9604	35		6	0	
190	8464	22		6	0	
187	7396	18		2	0	
180	7056	14		1	0	
176	6889	60		31	0	
169 2704 5			2	0		
162	2500	3		1	0	
151	2401	7		7	0	