## **Assembly Report**

**Experiment Details** 

ExperimentID:	Experiment 53
Sequencing Coverage:	23
Number of Partitions:	3
Allowed Mismatch Count in (Base):	0
Length of Sliding Window:	67

**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	326

Assembly Details			
Number of Contigs:	24		
Contig N50:	1026		
Contig N90:	826		
Number of Scaffold	1		
Scaffold N50:	10080		
Scaffold N90:	10080		
Mis-assembly Count:	0		
TotalAssembly Size:	10080		

Repeat Details

Total Reads Count(Non-Repeat) :	317
Total Reads Count(Repeat):	82
Retetitve Read Count based on (Partitions Identifier):	80
Retetitve Read Count (Entire Read Frequency Identifier):	3
Retetitve Read Count (Contained Read Frequency Identifier):	3
Total Repetitive Sequences Count	537
Total Unique Repetitive Sequences Count	3

Total Repeat Size (Base)		4200				
Starting	Ending	Repetitive Sequences	Repeat Count	Length	Position	S
1	10080	CGGTCGAAAA ACTGCTGGCA GTGGGGCATT ACCTCGAATCT ACCGTCGATAT TGCTGAGTCC ACCCGCCGTA TTGCGGCAAG TCGTATTCCGG CTGATCACATG GTGCTGATGG CAGGTTTCACC GCCGGTAATG AAAAAGGCGA ACTGGTGGTG CTTGGACGCA ACTGCTGGTG CTGCTGTTAC GCGCGATTTAC GCGCCGATTT GGACGCA CTACTCTGCTG CGGTGCTGGC TGCCTGTTTAC GCGCCGATTT GGACGGACGT TGCGAGATTT GGACGGACGT TGCGAGATTT GGACGGACGT TGCGAGATTT GGACGGACGT TGCGAGATTT TGCGAGATTT GGACGGATG TTGCGAGTC TTGCGAGTC TTGCGAGTC TTGCGAGTC TTGCGAGTC TTGCGAGTC TTGCGAGTT TTGCGCGTCAG GTGCCCGATG TCCTACCAGG AAGCGATGGA GCTTTCCTACT TCGGCGCTCT AGGTCAGGCC	4	350	Start Positio n 1376 4127 6570 8665	End Positio n 1725 4476 6919 9014

	1	
GATTCATTCGG		
GATGGTCTGT		
GTGGATTAAAA		
AAAGAGTGTCT		
GATAGCAGCTT		
CTGAACTGGTT		
ACCTGCCGTG		
AGTAAATTAAA		
ATTTATTGAC		
TTAGGTCACTA		
AATACTTTAAC		
CAATATAGGCA		
TAGCGCACAG		Start End
ACAGATAAAAA		Positio Positio
TTACAGAGTAC		n n
ACAACATCCTC		
1 10080 AAAGCCTACC 4	350	3285 3634
GGTGACAGTG		4858 5207
CGGGCTTTTT		7202 7551
TTCGACCAAAG		
GTAACGAGGT		9332 9681
AACAACCATGC		
GAGTGTTGAA		
GTCAGGAGAT		
CCTAAAGGCC		
TGTACCCGTTA		
CCTAGCCAGTT		
GGCATTAAAC		
GTATACGGTAC		
CTAGGCATGTA		
CGTAATCGTAG		
CCTTAGCAATC		
TCCAGTCC		

	<u> </u>	i	1	i	1
		GCTTTTCATTC			
		TGACTGCAAC			
		GGGCAATATG			
		тстствтвт			
		ATTAAAAAAAG			
		AGTGTCTGATA			
		GCAGCTTCTG			
		AACTGGTTACC			
		TGCCGTGAGT			
		AAATTAAAATT			
		TTATTGACTTA			
		GGTCACTAAAT			
		ACTTTAACCAA		Start	End
		TATAGGCATAG		Positio	Positio
		CGCACAGACA		n	n
		GATAAAAATTA		1	
1	10080	CAGAGTACAC 4	350	1	350
		AACATCCATGA		799	1148
		AACGCATTAGC		2039	2388
		ACCACCATTAC			
		CACCACCATCA		8043	8392
		CCATTACCACA			
		GGTAACGGTG			
		CGGGCTGACG			
		CGTACAGGAA			
		ACACAGAAAAA			
		AGCCCGCACC			
		TGACAGTGCG			
		GGCTTTTTTTT			
		CGACCAAAGG			
		TAACGAGGTAA			
		CAACCATGCG			
		AGTGTTGAAGT			

SRGD Performance Metrics

Initial Reads Count:	1212
Reads Count after Removing Duplication:	399

Reads Count after Overlapping:		227			
Initial Dataset Size in (Base):		241272			
Dataset Sizet after	Removing Duplication	n:	101376		
Dataset Size after 0	Overlapping:		49625		
Overlapping Metri	CS				
Repeat Identificatio	n Time:		00h:00m	n:08s:000ms	
Overlapping Time:			00h:04m	n:00s:000ms	
Reads Alignment T	ime:		00h:00m	n:24s:000ms	
Total Hybrid Assem	Total Hybrid Assembly Time:		00h:04m:24s:000ms		
Repeat Annotation	Time:		00h:00m:42s:000ms		
Overlapping Length	O(N)2 Time Complexity	Hit Index Count		Overlapping Matched Count	RI
200	100489	61		7	0
197	96100	38		1	0
196	95481	62		5	0
195	92416	62		4	0
194	90000	58		1	0
191	89401	44		1	0
190	88804	38		1	0
188	88209	49		3	0
187	86436	43		1	0
186	85849	42		1	0
183	85264	44		1	0
177	84681	56		1	0

176	84100	296	138	0
152	23104	13	1	0
151	22801	11	5	0
150	21316	17	1	0