## **Assembly Report**

Experiment Details

ExperimentID:	Experiment 25
Sequencing Coverage:	31
Number of Partitions:	5
Allowed Mismatch Count in (Base):	0
Length of Sliding Window:	61

**Reads Dataset Details** 

reads Dataset Details	
Total Reads Count:	1045
Total Dataset Size in (Base):	313765
Valid Read Count:	1045
Rejected Read Count:	0
Maximum Read Length:	301
Minimum Read Length:	262

Reference Genome Details

Length of Reference Genome (Base):	10080	
Number of K-mers:	9780	
Length of K-mer	501	

Assembly Details

Number of Contigs:	13
Contig N50:	1376
Contig N90:	1251
Number of Scaffold	1
Scaffold N50:	10080
Scaffold N90:	10080
Mis-assembly Count:	0
TotalAssembly Size:	10080

Repeat Details

Total Reads Count(Non-Repeat) :	337
Total Reads Count(Repeat):	57
Retetitve Read Count based on (Partitions Identifier):	57
Retetitve Read Count (Entire Read Frequency Identifier):	1
Retetitve Read Count (Contained Read Frequency Identifier):	1
Total Repetitive Sequences Count	450
Total Unique Repetitive Sequences Count	1

Total Repeat Size (Base)		1400				
Starting	Ending	Repetitive Sequences	Repeat Count	Length	Position	s
1	10080	GCTTTTCATTC TGACTGCAAC GGGCAATATG TCTCTGTGTGG ATTAAAAAAAA GAGTGTCTGATA GCAGCTTCTG AACTGGTTACC TGCCGTGAGT AAATTAAAATT TTATTGACTTA GGTCACTAAAT ACTTTAACCAA TATAGGCATAG CGCACAGACA GATAAAAATTA CAGAGTACAC AACATCCATGA AACGCATTAC CACCACTACA CCACCATCA CCATTACCACA GGTAACGGTG CGGCTGACG CGGCTGACG CGGCTGACG CGTACAGGAA ACACCATCA CCATTTACCACA GGTAACGGTG CGGCTGACG CGTACAGGAA ACACAGGAAAAA AGCCCGCACC TGACAGTGCG GGCTTTTTTT CGACCAAAGG TAACGAGGTAA CAACCATGCG AGTGTTGAAGT	4	350	Start Positio n 1 799 1434 8510	End Positio n 350 1148 1783 8859

Initial Reads Count:		1045			
Reads Count after Removing Duplication:		394			
Reads Count after Overlapping:		167			
Initial Dataset Size in (Base):		313765			
Dataset Sizet after Removing Duplication:		129553			
Dataset Size after Overlapping:		58971			
Overlapping Metrics			Γ		
Repeat Identification Time:		00h:00m:12s:000ms			
Overlapping Time:		00h:03m:53s:000ms			
Reads Alignment Time:		00h:00m:18s:000ms			
Total Hybrid Assembly Time:		00h:04m:11s:000ms			
Repeat Annotation Time:		00h:00m:21s:000ms			
Overlapping Length	O(N)2 Time Complexity	Hit Index Count		Overlapping Matched Count	RI
297	113569	45		1	0
276	112896	350		164	0
251	29584	8		2	0
226	28900	165		60	0