

C706407:

Tel: 1-888-762-2568 Fax: 1-510-783-5386 Email: info@biochain.com

Data Sheet

Product Name: Paraffin Tissue Section - Human Tumor: Breast, Visium-Characterize
Catalog No.: <u>T2235086-SV</u> Lot No.: C706407
Species: ■ Human □ Mouse □ Rat □ Monkey (Rh) □ Guinea Pig □ Porcine □ Bovine □ Hamster □ Dog □ Monkey (Cy) □ Rabbit □ Plant
Tissue Type: □ Normal □ Adult □ Fetal ■Tumor □ Disease □ Cell line Tissue Name: Breast Tumor Size: N/A
Donor Information: H50523T, Race: Caucasian Male: year(s) old Female:68 year(s) old
Pathological Diagnosis: Invasive carcinoma, Stage IIIB T1cN3aM0
Location: Breast
Visium Characterization:





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Summary		
Number of Spots Under Tissue	2,353	
Mean Reads per Spot	64,778	
Median Genes per Spot	3,061	
Sequencing		
Number of Reads	152,423,699	
Valid Barcodes	98.4%	
Valid UMIs	100.0%	
Sequencing Saturation	87.2%	
Q30 Bases in Barcode	96.4%	
Q30 Bases in Probe Read	68.7%	
Q30 Bases in UMI	96.3%	
Mapping		
Reads Mapped to Probe Set	98.8%	
Reads Mapped Confidently to Probe Set	98.1%	
Reads Mapped Confidently to the Filtered Probe Set	83.2%	
Spots		
Fraction Reads in Spots Under Tissue	84.2%	
Mean Reads per Spot	64,778	
Mean Reads Under Tissue per Spot	53,620	
Median UMI Counts per Spot	5,451	
Median Genes per Spot	3,061	
Genes Detected	16,106	



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Summary Legend

Column Header	Definition	
	Total number of read pairs that were assigned to this library in	
Number of Reads	demultiplexing.	
	Fraction of reads with barcodes that match the whitelist after barcode	
Valid Barcodes	correction.	
	Fraction of reads with valid UMIs; i.e. UMI sequences that do not contain	
Valid UMIs	Ns and that are not homopolymers.	
	The fraction of reads originating from an already-observed UMI. This is a	
	function of library complexity and sequencing depth. More specifically, this	
	is the fraction of confidently mapped, valid spot-barcode, valid UMI reads	
Sequencing Saturation	that had a non-unique spot-barcode/UMI/gene combination.	
OOO Bassa in Baysada	Fraction of spot barcode bases with Q-score >= 30, excluding very low	
Q30 Bases in Barcode	quality/no-call (Q <= 2) bases from the denominator.	
	Fraction of RNA read bases with Q-score >= 30, excluding very low	
Q30 Bases in Probe Read	quality/no-call (Q <= 2) bases from the denominator. This is Read 2 for the Visium v1 chemistry.	
Q30 Bases III Flobe nead	Fraction of UMI bases with Q-score >= 30, excluding very low quality/no-	
Q30 Bases in UMI	call ($Q \le 2$) bases from the denominator.	
Reads Mapped to Probe Set	Fraction of reads that mapped to the probe set.	
Reads Mapped Confidently to Probe		
Set	Fraction of reads that mapped uniquely to a probe in the probe set.	
	Fraction of reads from probes that map to a unique gene. These reads are	
Reads Mapped Confidently to the	considered for UMI counting by default. This metric will be None when	
Filtered Probe Set	probe filtering is disabled.	
Fraction Reads in Spots Under	The fraction of valid-barcode, confidently-mapped-to-transcriptome reads	
Tissue	with tissue-associated barcodes.	
	The number of reads, both under and outside of tissue, divided by the	
Mean Reads per Spot	number of barcodes associated with a spot under tissue.	
	The number of reads under tissue divided by the number of barcodes	
Mean Reads Under Tissue per Spot	associated with a spot under tissue.	
Median UMI Counts per Spot	The median number of UMI counts per tissue covered spot.	
	The median number of genes detected per spot under tissue-associated	
Median Genes per Spot	barcode. Detection is defined as the presence of at least 1 UMI count.	
	The number of unique genes from the filtered probe set with at least one	
Genes Detected	UMI count in any tissue covered spot.	

Components:

- 5 slides per package
 Certificate of Analysis

APPROVED BY: