filter	software	purpose
GATK	GATK	Removes putative SNVs with GATK quality scores less than 40 (as part of the GATK
SS	SomaticSniper	processing, with inder realignment and base recampration) Removes putative SNVs with a SomaticScore less than 40
VAQ	SomaticSniper	Removes putative SNVs with Somatic Sniper Varaint Allele Quality scores less than 20
НОП	SomaticSniper, python	Removes putative SNVs that are identified as loss of heterozygosity
10bp-SNV		Removes putative SNVs located within a 10 bp window of any other putative SNV
10bp-INDEL		Removes putative SNVs located within a 10 bp window of indels
IbSNP	python	Removes putative SNVs that overlap with dbSNP coverage
<10%	python	Removes putative SNVs if, in the tumor data, the percentage of reads covering the site with the alternate allele is less than 10%