# Analisi di dati trascrittomici di RNA-seq da frazioni polisomiale e totale da una linea cellulare epiteliale tumorale

Supervisore

Alberto Inga

Co-supervisori Alessandro Romanel Samuel Valentini Laureando

Giacomo Fantoni

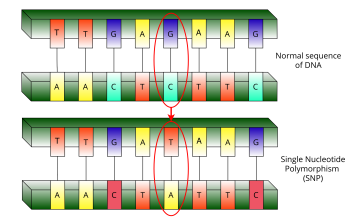


Dipartimento di Ingegneria e Scienza dell'informazione Corso di Laurea in Informatica

- 4 ロ b 4 個 b 4 き b 4 き b 9 Q C

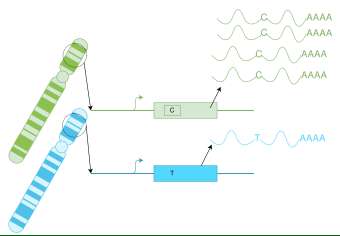
Università di Trento

#### **TransSNP**





## Individuare transSNP



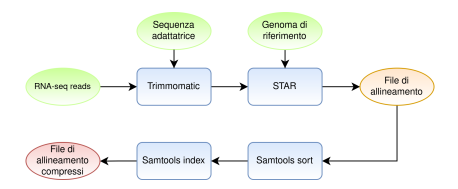
Giacomo Fantoni

Università di Trento

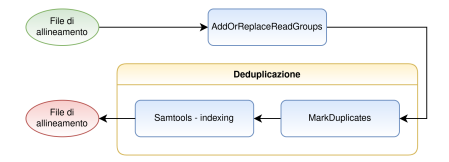
# Campioni

Nome	Trattamento	Linea cellulare
scr_DMS0	DMSO	HCT116
scr_NUTLIN	NUTLIN	HCT116
shDHX30_DMSO	DMSO	HCT116 con knockdown di DHX30
shDHX30_NUTLIN	NUTLIN	HCT116 con knockdown di DHX30
shPCBP2_DMSO	DMSO	HCT116 con knockdown di PCBP2
shPCBP2_NUTLIN	NUTLIN	HCT116 con knockdown di PCBP2

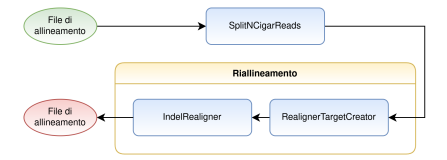
#### Allineamento



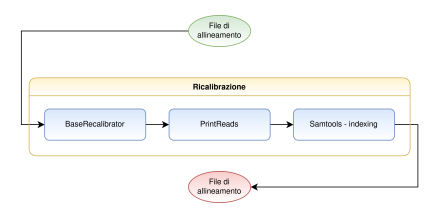
#### Deduplicazione



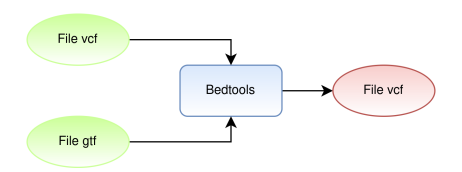
#### Riallineamento



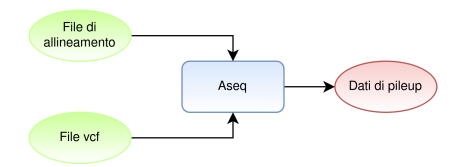
#### Recalibrazione



#### Analisi dati WES

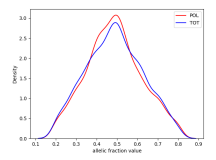


## **ASEQ**



# Analisi dati di pileup

dbsnp	ref	alt	Α	С	G	T	af	cov
rs1423991279	G	Α	6	0	5	0	0.545455	11
rs2272757	G	Α	118	0	128	0	0.479675	246
rs975483396	G	Α	6	0	6	0	0.500000	12



Filtri sugli SNP	SNP individuati		
Input di ASEQ	40418219		
$cov \geq 10$ e $0.2 \leq af \leq 0.8$	13681		
Dati per 3 replicati	851		

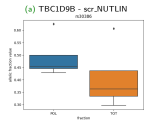
#### Analisi dei risultati

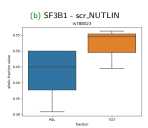
Condizione	Totali	3'-UTR	5′-UTR
scr_DMSO	27	8	2
scr_NUTLIN	33	11	2
shDHX30_DMSO	22	9	1
shDHX30_NUTLIN	25	11	0
shPCBP2_DMSO	24	10	1
shPCBP2_NUTLIN	30	11	2

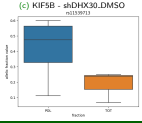
Table: SNP individuati con p-value nominale inferiore a 0.05



#### Geni individuati







# Grazie per l'attenzione.

