# SQANTI3 filter report

Total Genes: 650

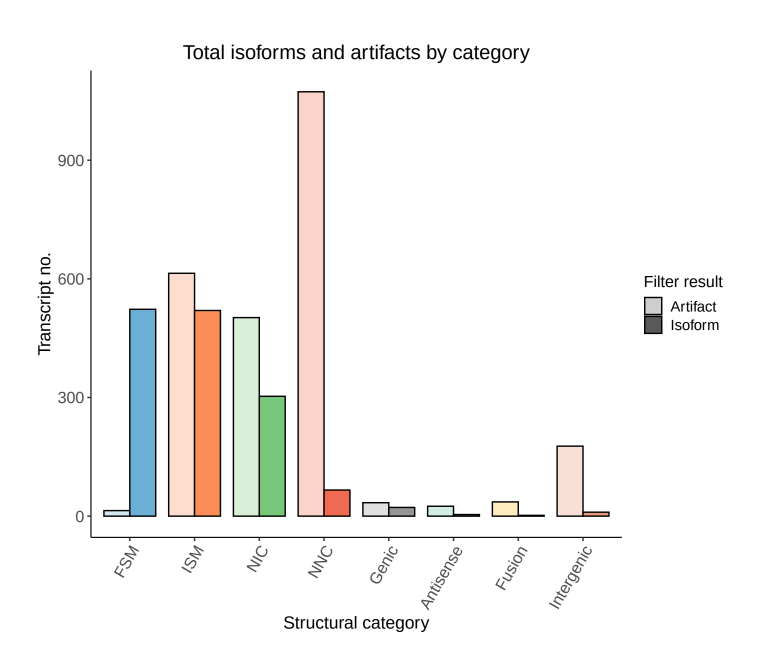
Total Transcripts: 3925

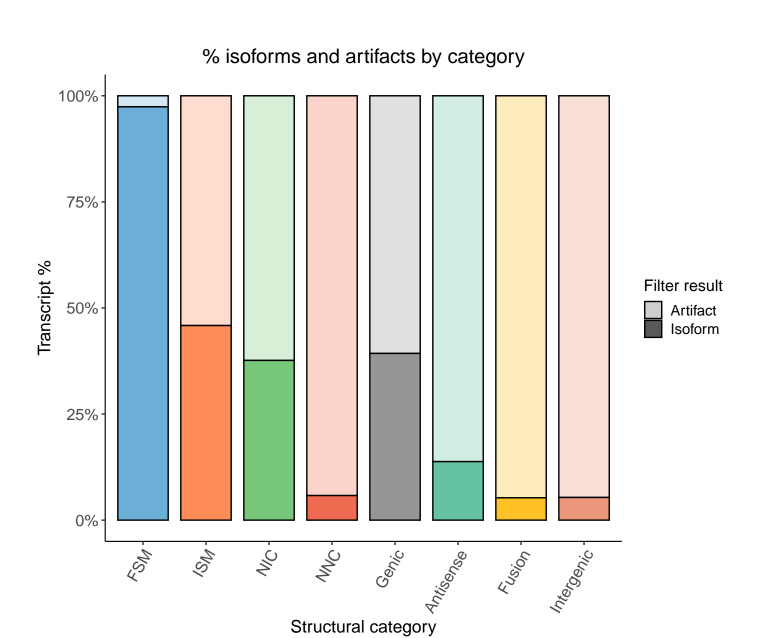
- Isoforms: 1450 (37%)

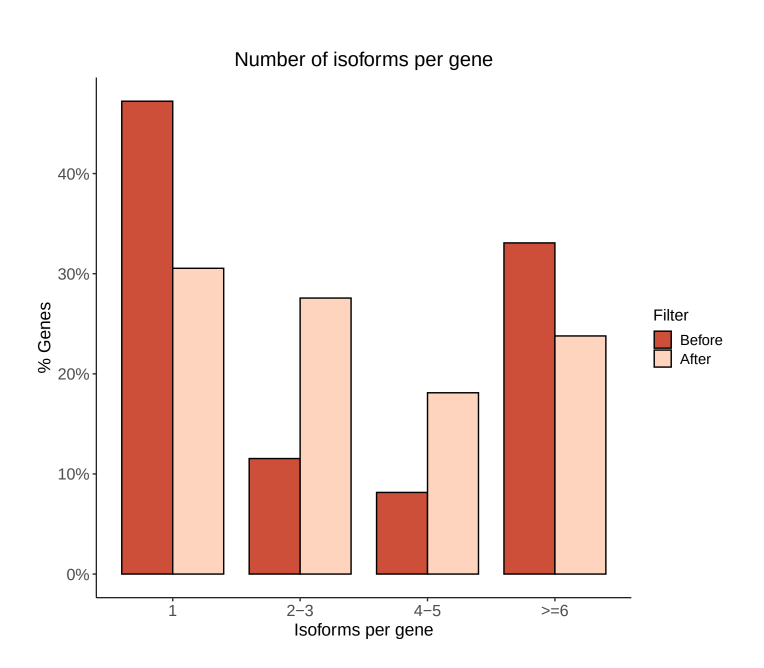
- Artifacts: 2475 (63%)

Gene category	Gene no.	No. of genes with artifacts only
Annotated	437	81
Novel	213	199

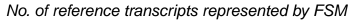
Structural category	Artifact no.	Isoform no.
FSM	14	523
ISM	614	520
NIC	502	303
NNC	1073	66
Genic	34	22
Antisense	25	4
Fusion	36	2
Intergenic	177	10

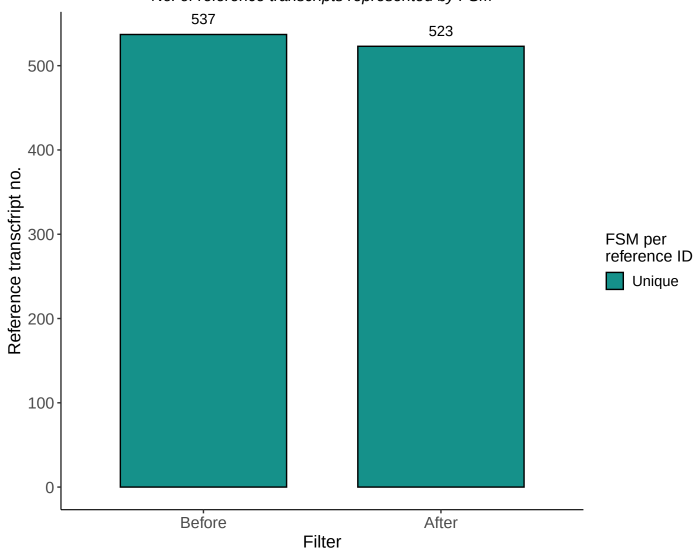




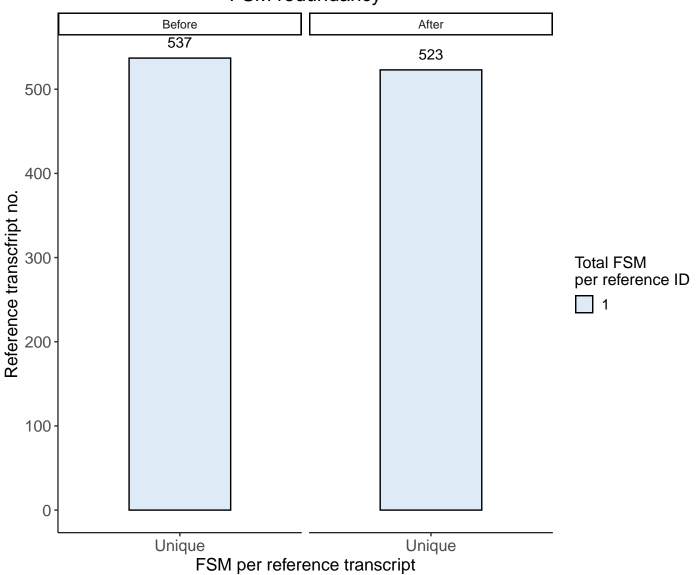


## Reference transcript complexity

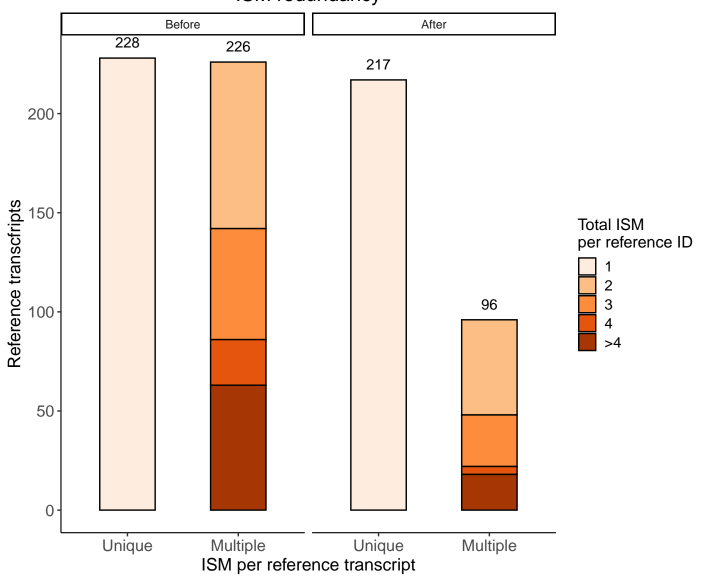




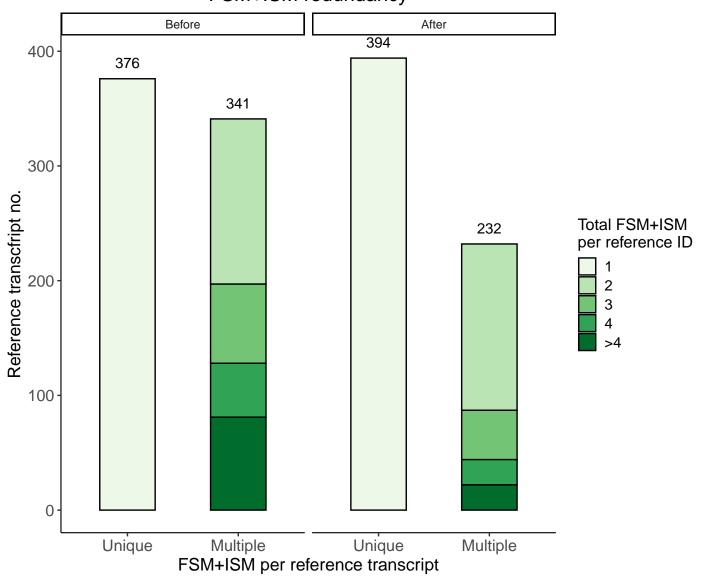
# FSM redundancy

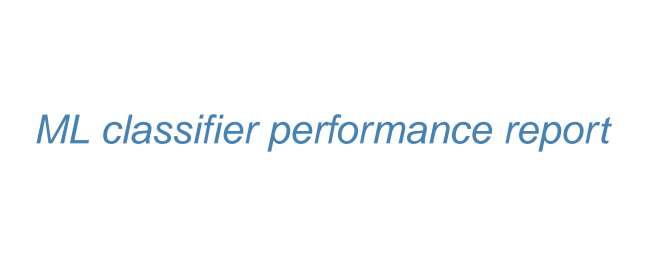


# ISM redundancy



## FSM+ISM redundancy





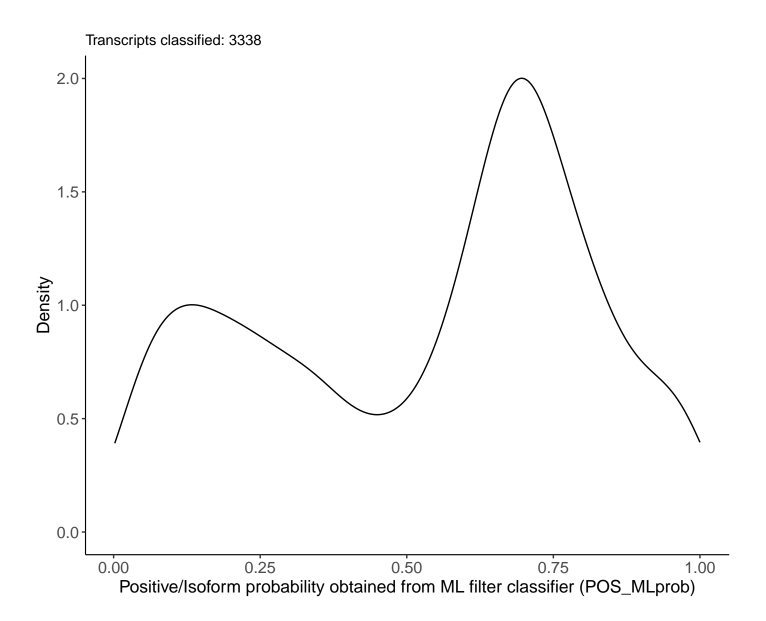
# Classification model performance on test set

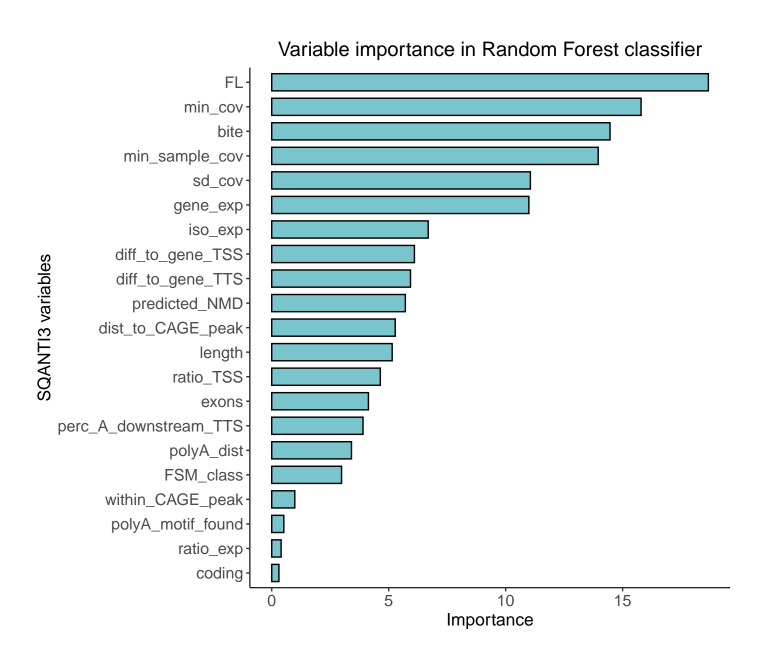
#### Performance metrics

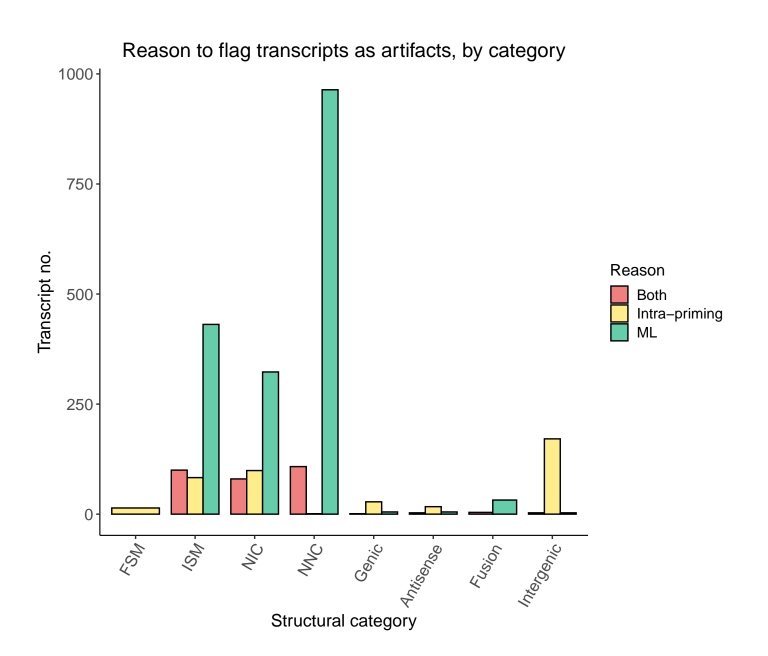
Metric	Value	
Accuracy	9.594595e-01	
Карра	9.189189e-01	
AccuracyLower	8.860627e-01	
AccuracyUpper	9.915602e-01	
AccuracyNull	5.000000e-01	
AccuracyPValue	3.578714e-18	
McnemarPValue	2.482131e-01	
Sensitivity	9.189189e-01	
Specificity	1.000000e+00	
Pos Pred Value	1.000000e+00	
Neg Pred Value	9.250000e-01	
Precision	1.000000e+00	
Recall	9.189189e-01	
F1	9.577465e-01	
Prevalence	5.000000e-01	
Detection Rate	4.594595e-01	
Detection Prevalence	4.594595e-01	
Balanced Accuracy	9.594595e-01	

#### Confusion matrix

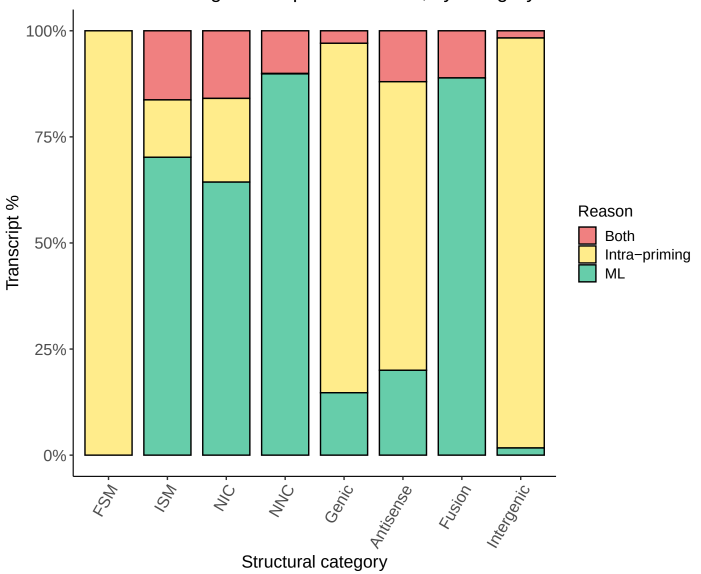
Prediction	Reference	Freq
POS	POS	34
NEG	POS	3
POS	NEG	0
NEG	NEG	37



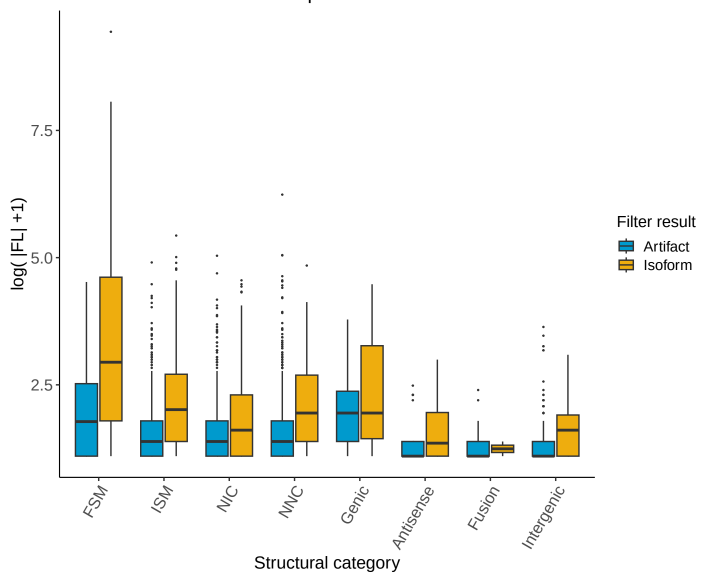


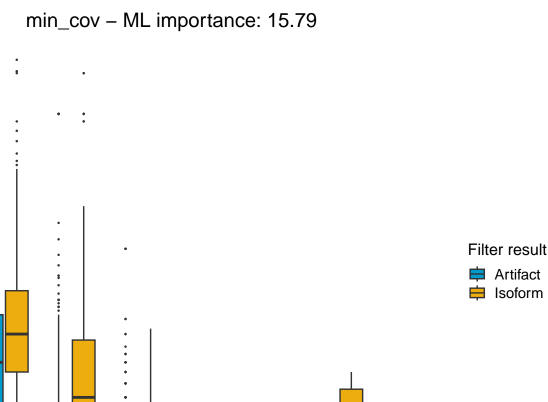


## Reason to flag transcripts as artifacts, by category



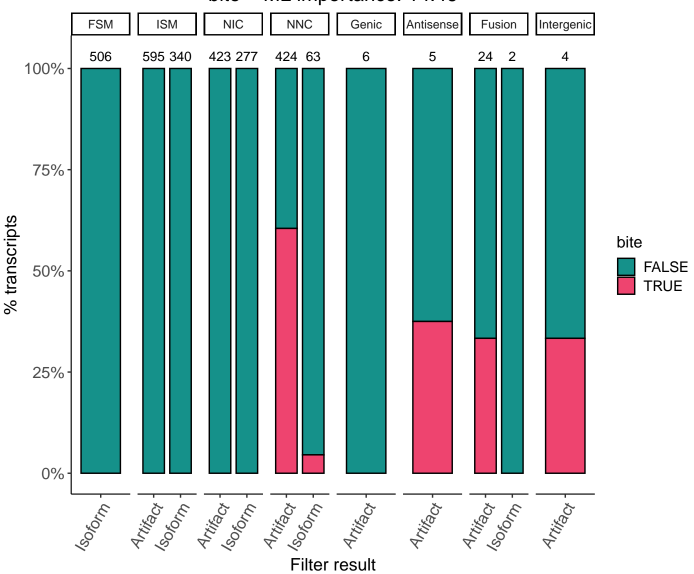
FL – ML importance: 18.67

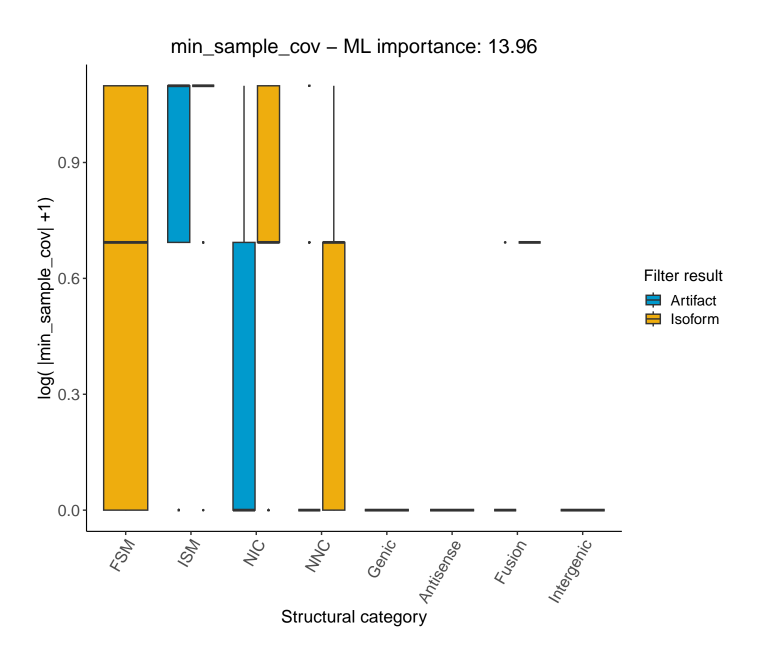


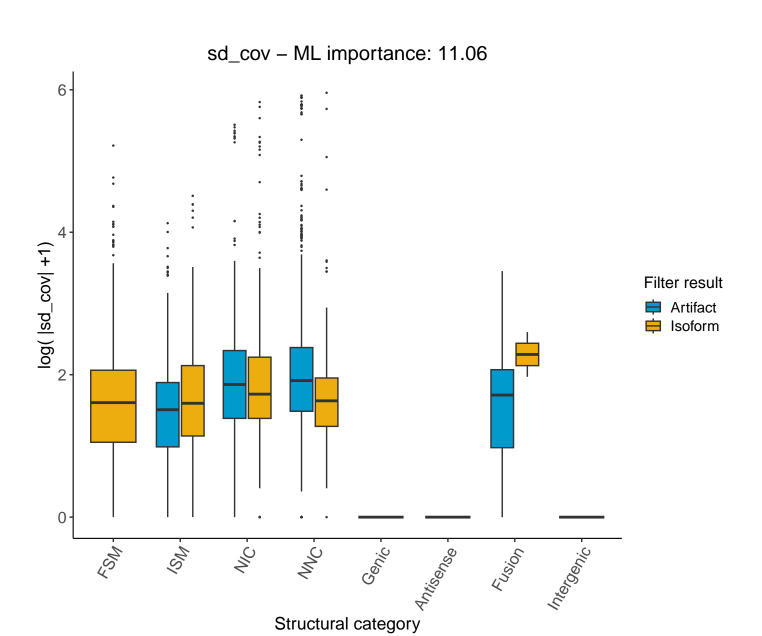


log( |min\_cov| +1) Filter result 0 -Structural category

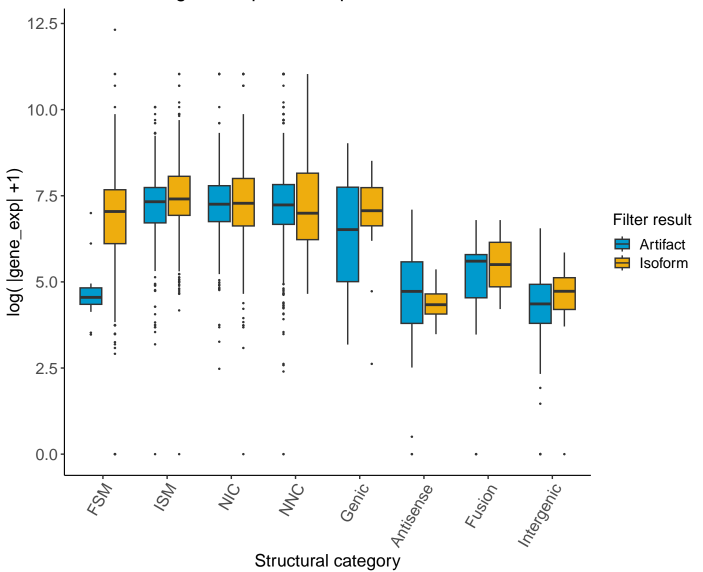
bite – ML importance: 14.46



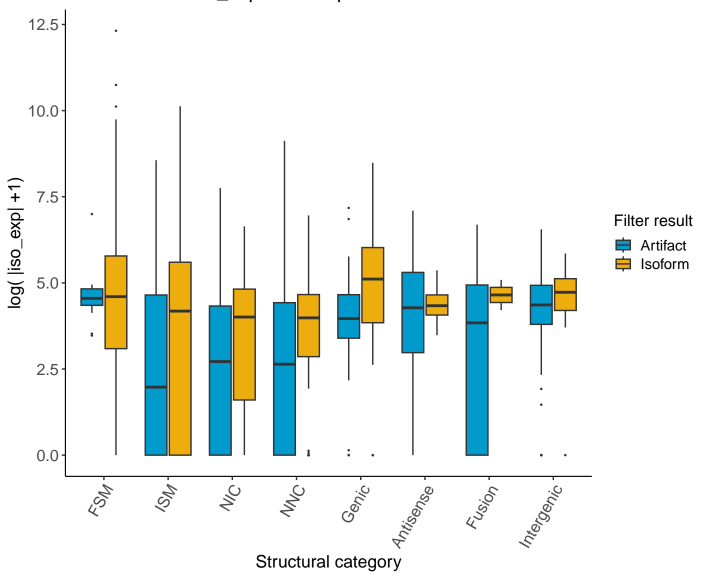




gene\_exp - ML importance: 10.99

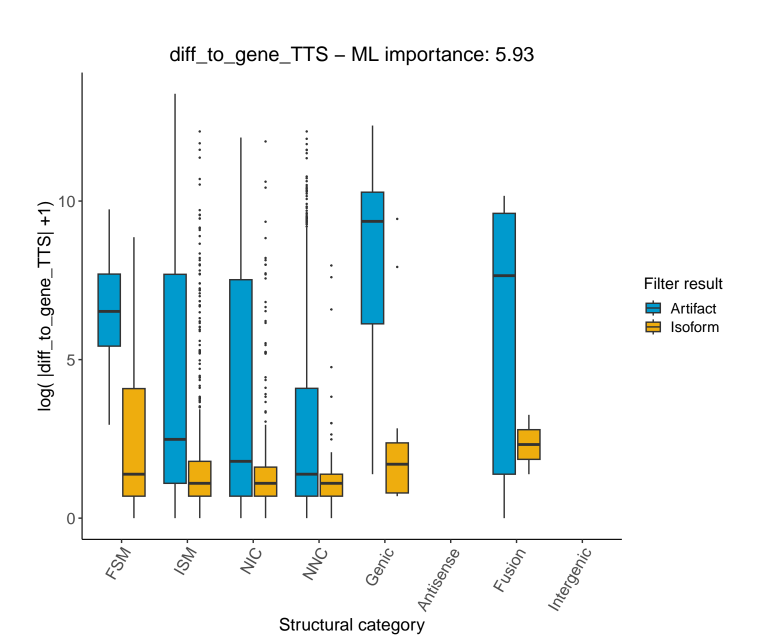


iso\_exp - ML importance: 6.69

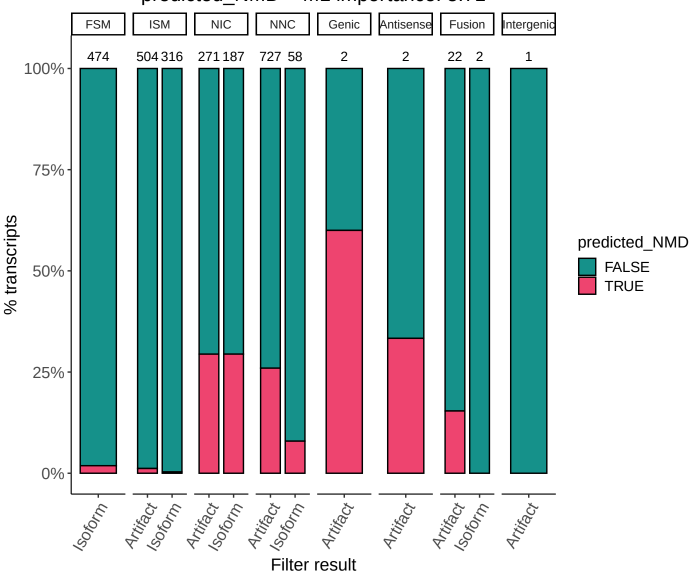


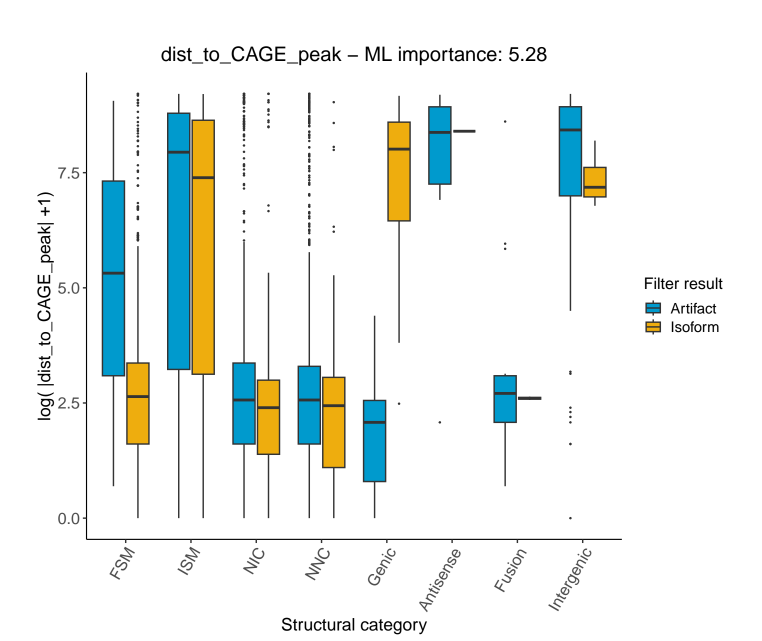
diff\_to\_gene\_TSS - ML importance: 6.1 10log( |diff\_to\_gene\_TSS| +1) Filter result Artifact Isoform 0. Genic -C

Structural category

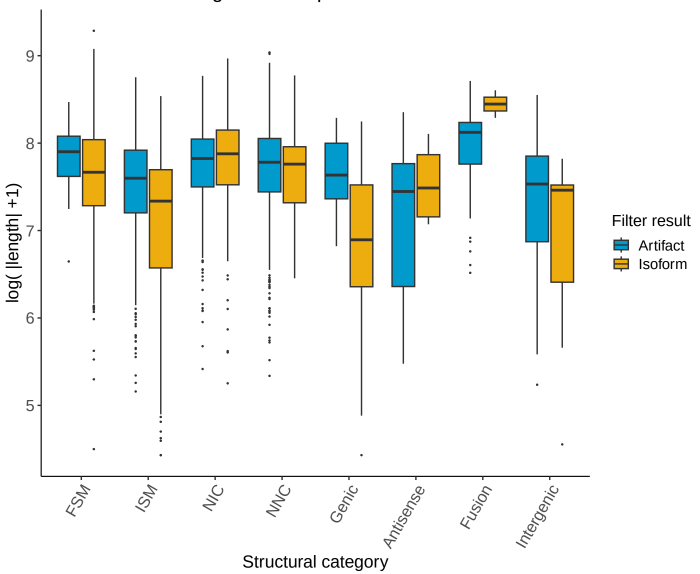


## predicted\_NMD - ML importance: 5.71

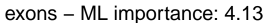


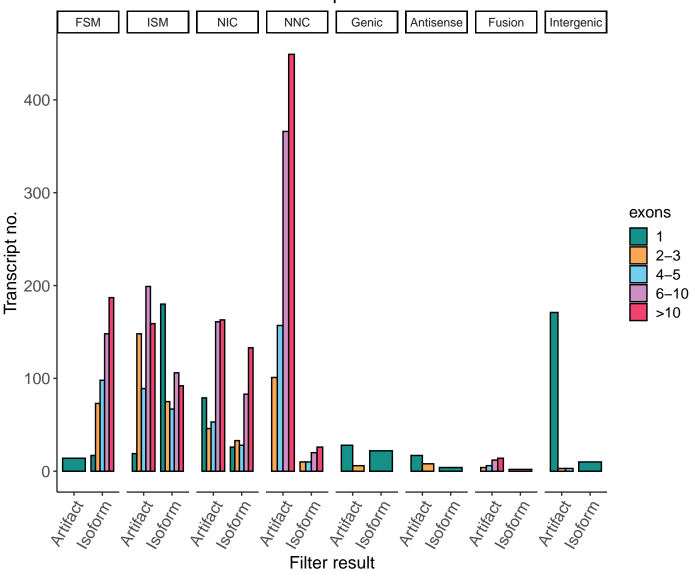


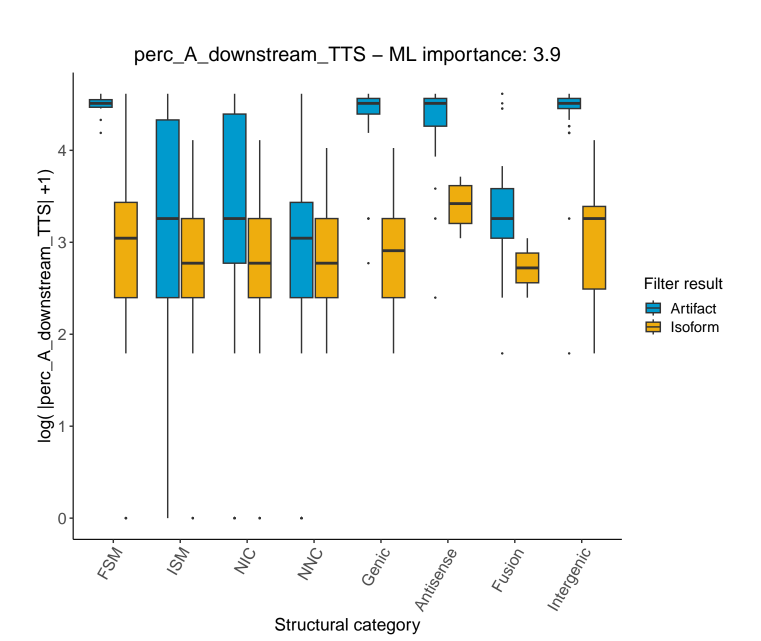
length – ML importance: 5.15



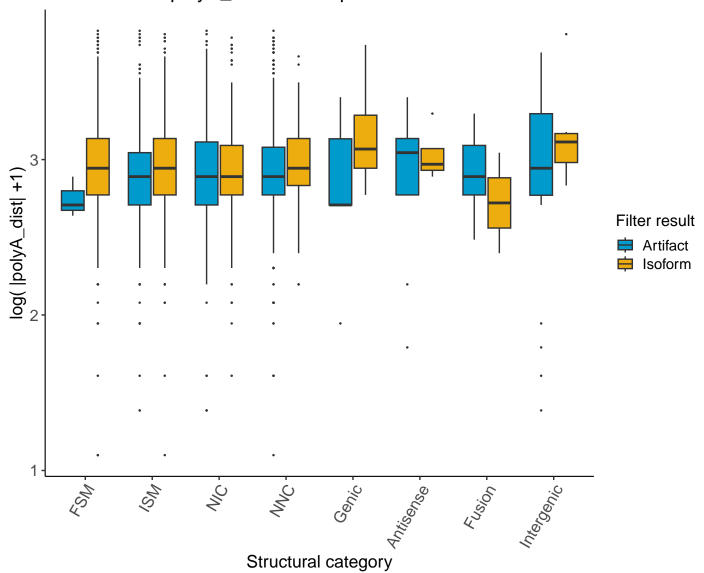
ratio\_TSS - ML importance: 4.64 10.0 7.5 log( |ratio\_TSS| +1) Filter result 5.0 Artifact Isoform 2.5 0.0 -WC-181 Structural category



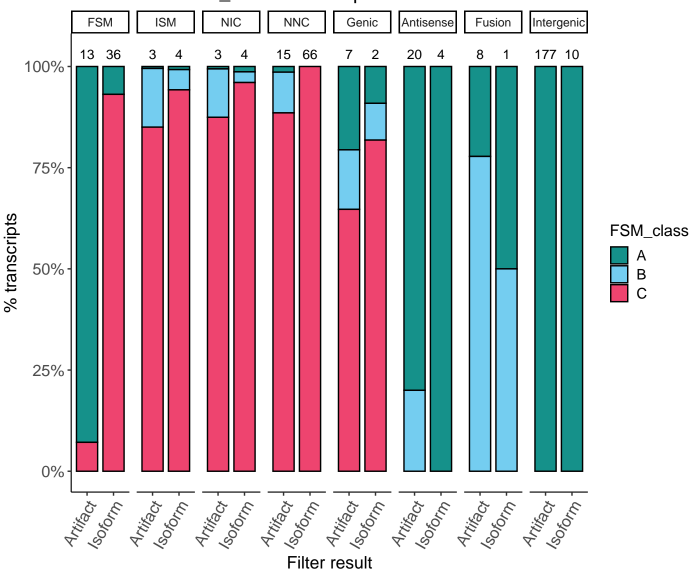




polyA\_dist - ML importance: 3.41



FSM\_class - ML importance: 2.99

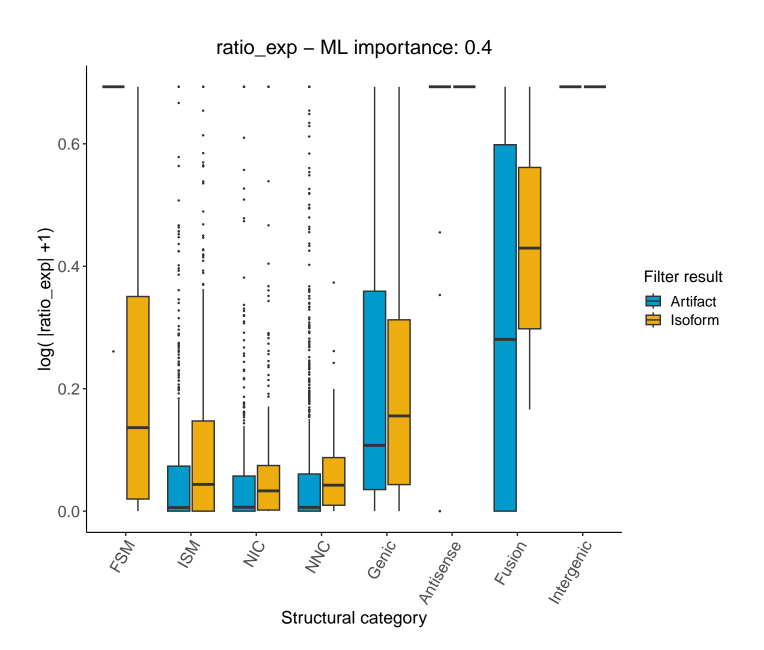


## within\_CAGE\_peak - ML importance: 0.99 **FSM** Fusion Genic Antisense ntergenio 11 137 481442 159 48 304 11 13 21 24 14 2 168 10 4 100% 75% % transcripts within\_CAGE\_peak **FALSE** 50% TRUE 25% 0%

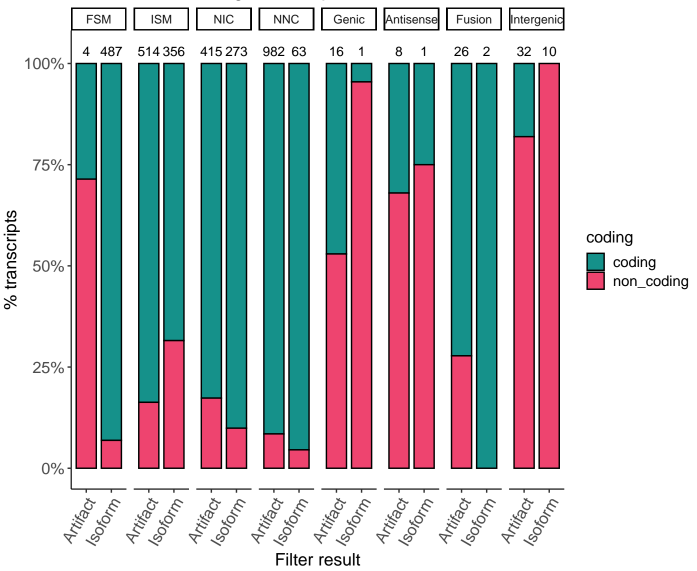
Filter result

polyA\_motif\_found - ML importance: 0.52 **FSM** Genic **Fusion** Antisense ntergenic 154 4 11 68 178 17 167 22 155 3 26 2 16 4 2 4 100% 75% % transcripts polyA\_motif\_found **FALSE** 50% **TRUE** 25% 0% 

Filter result



### coding - ML importance: 0.31





A % by category Red line indicates threshold employed in ML filter 100 75 % A's downstream of TTS 50 25 0 FSM ISM NiC NNC Genic Antisense Fusion Intergenic

Structural category

