

# *SQANTI3 filter report*

*Total Genes: 650*

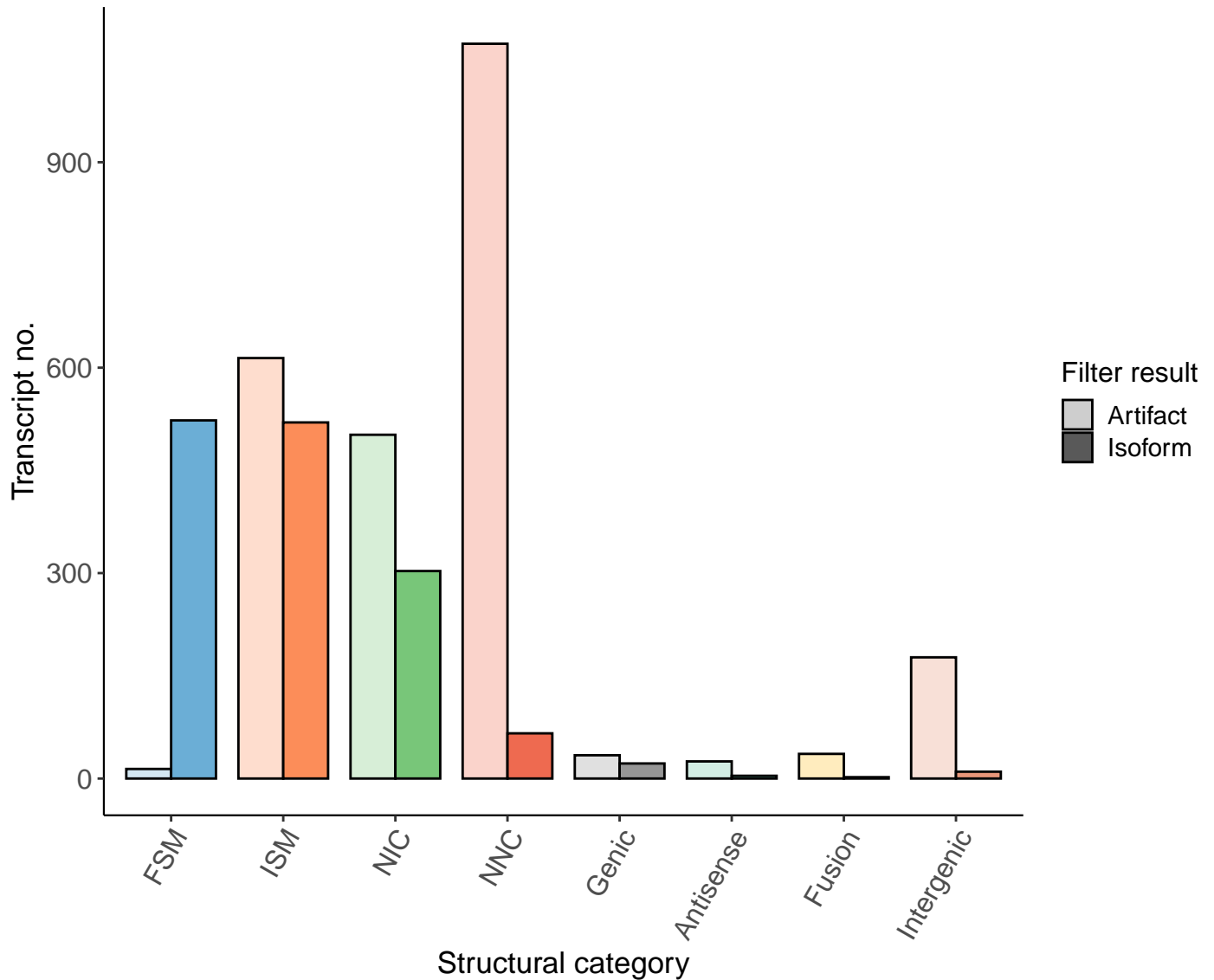
*Total Transcripts: 3925*

*– Isoforms: 1450 (37%)*

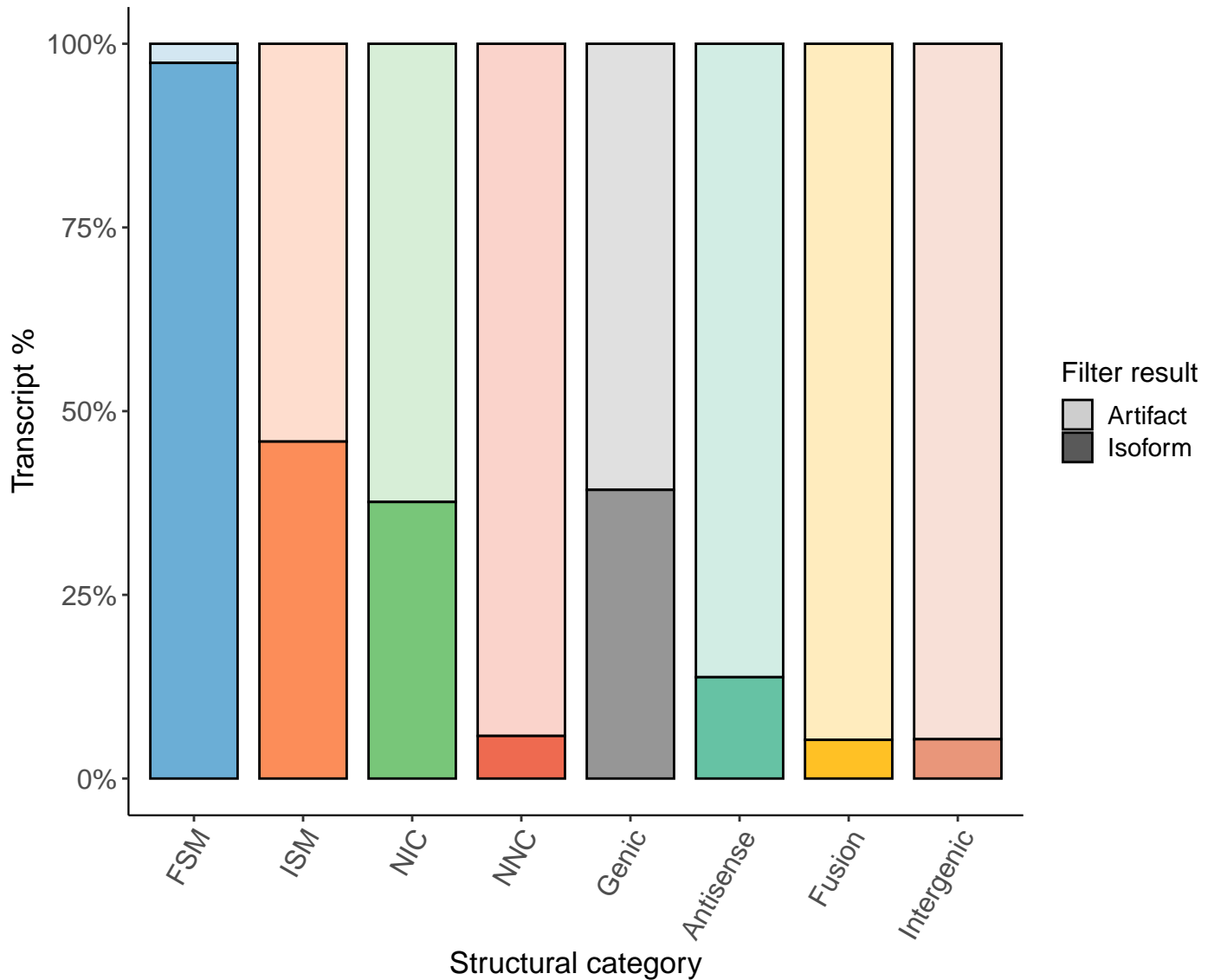
*– Artifacts: 2475 (63%)*

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

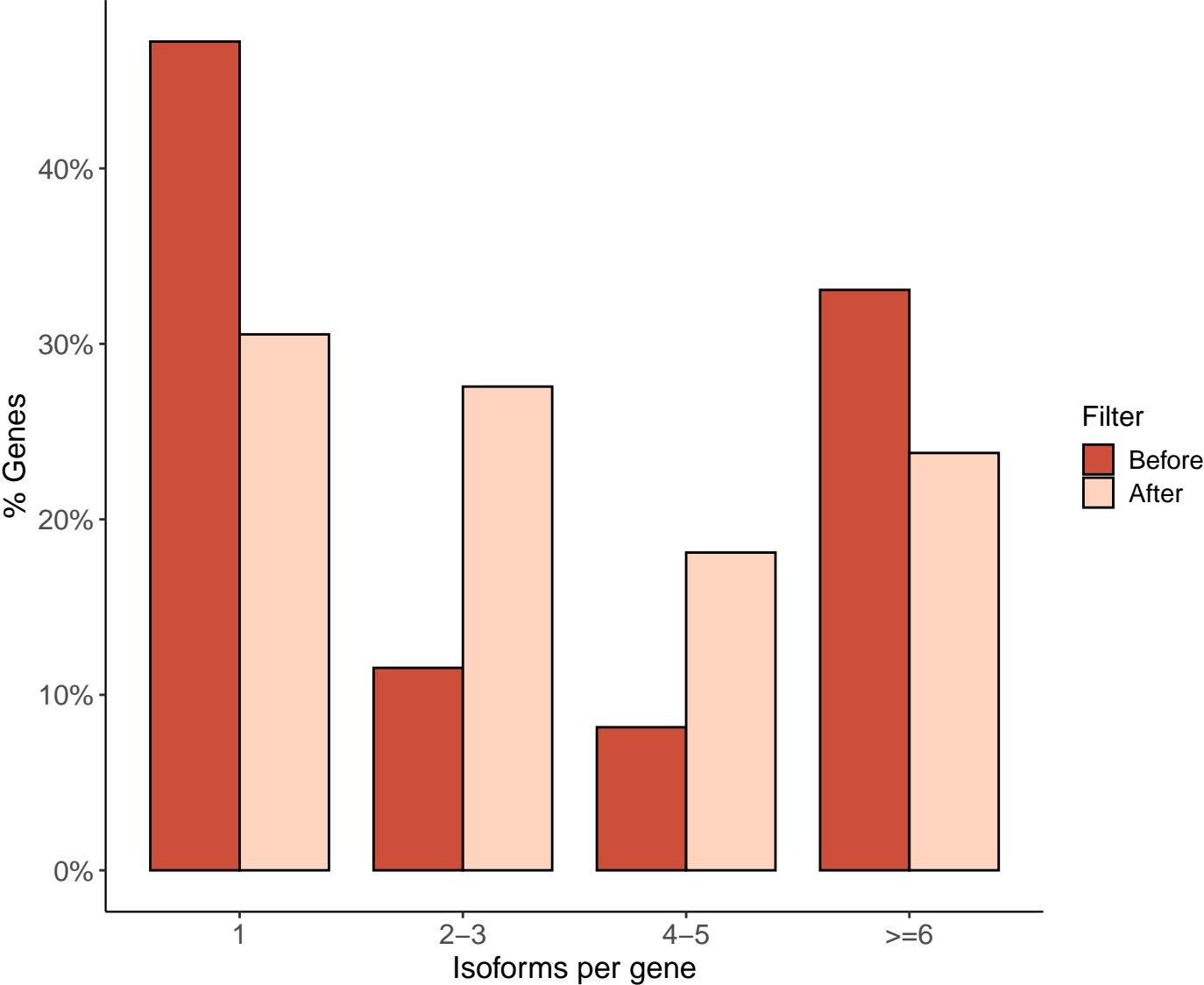
Total isoforms and artifacts by category



% isoforms and artifacts by category

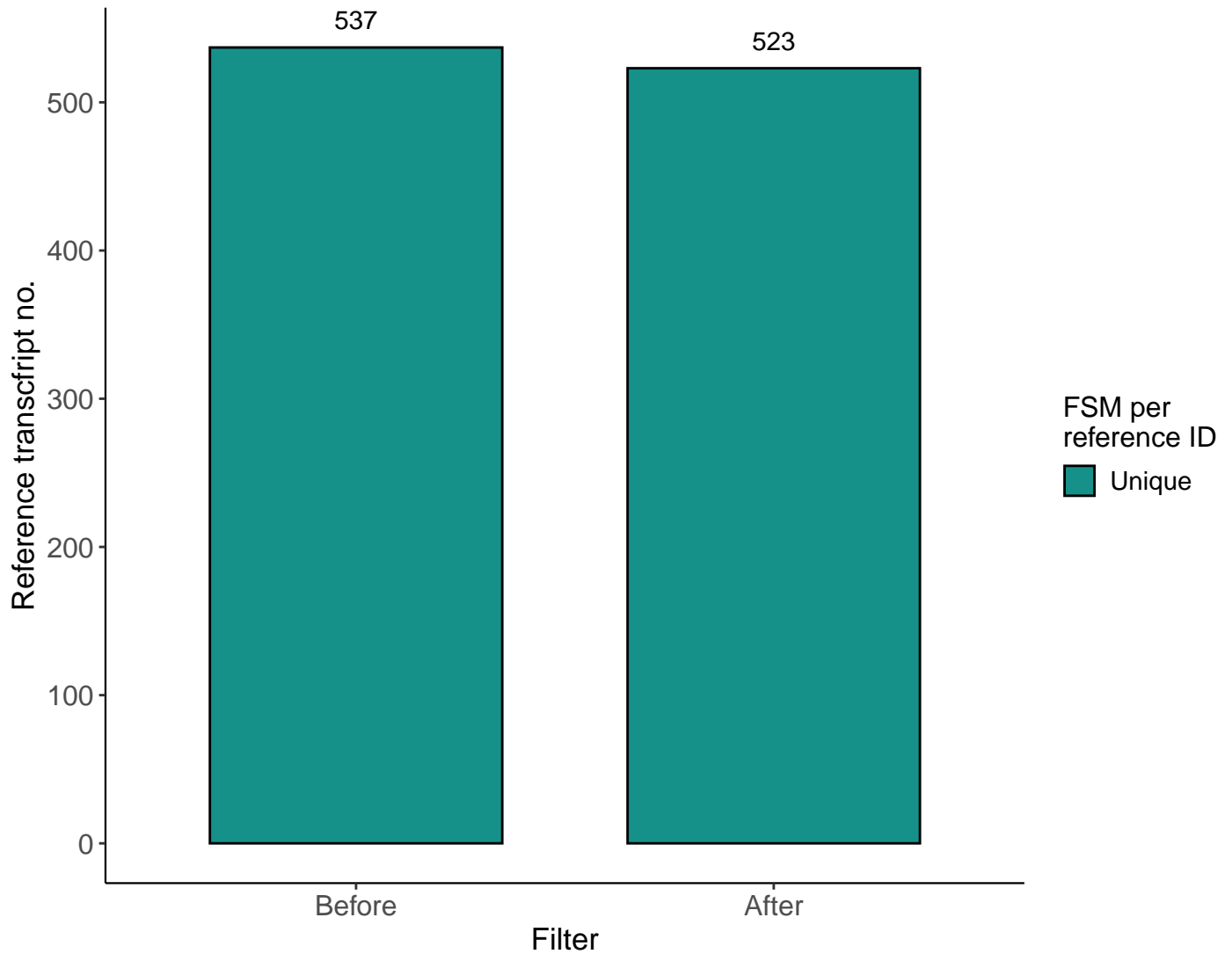


Number of isoforms per gene

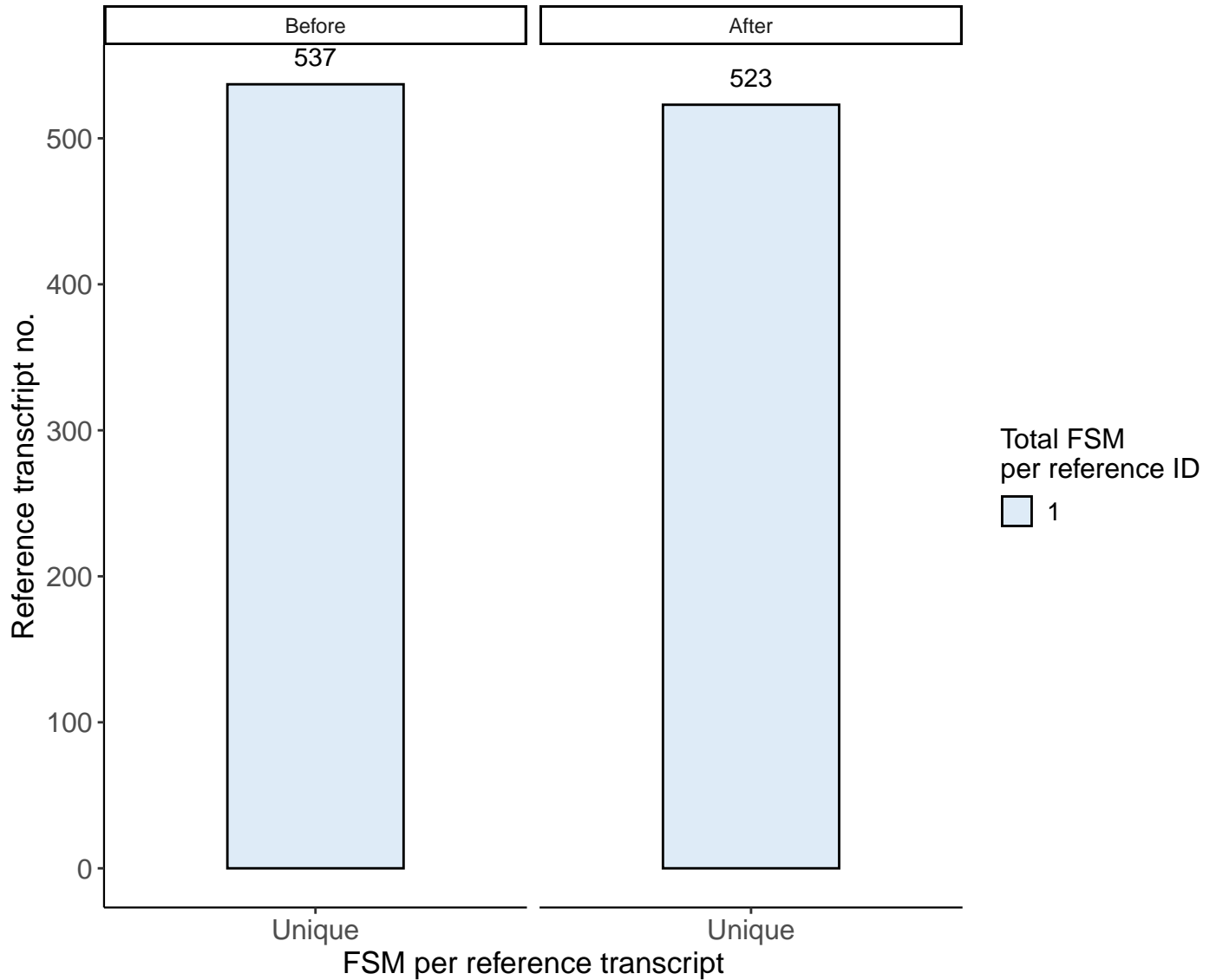


## Reference transcript complexity

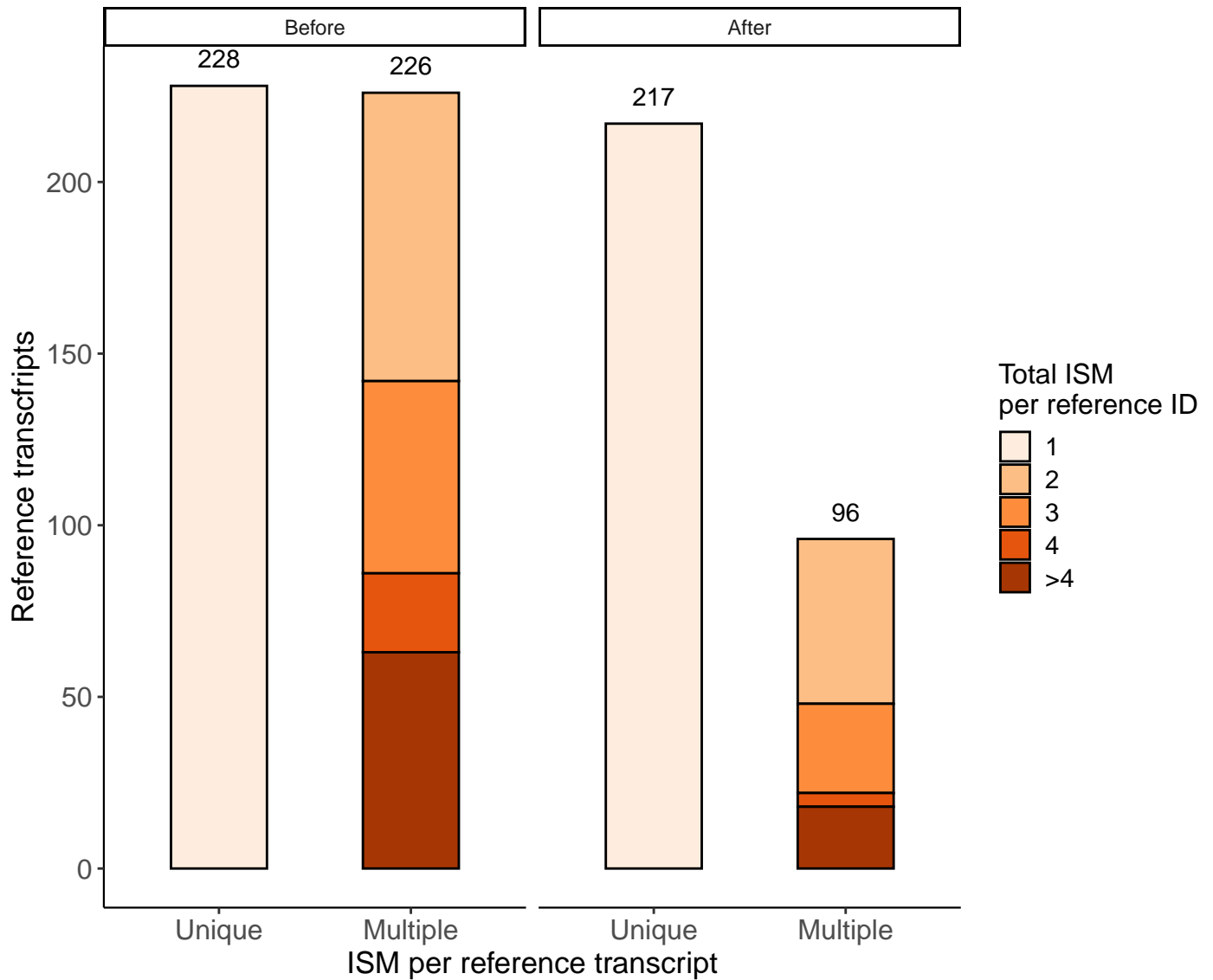
*No. of reference transcripts represented by FSM*



## FSM redundancy

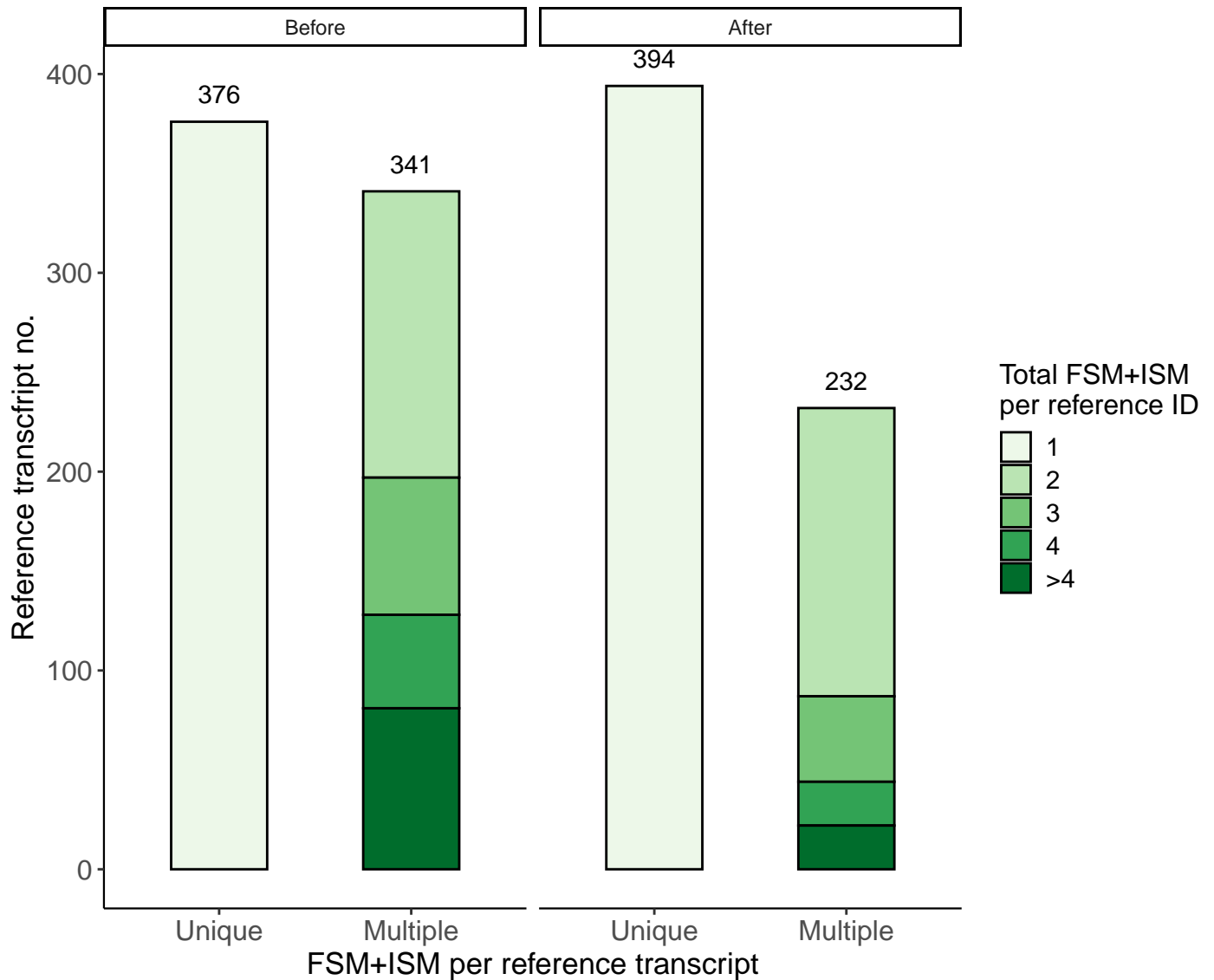


## ISM redundancy





## FSM+ISM redundancy



# *ML classifier performance report*

Classification model performance on test set

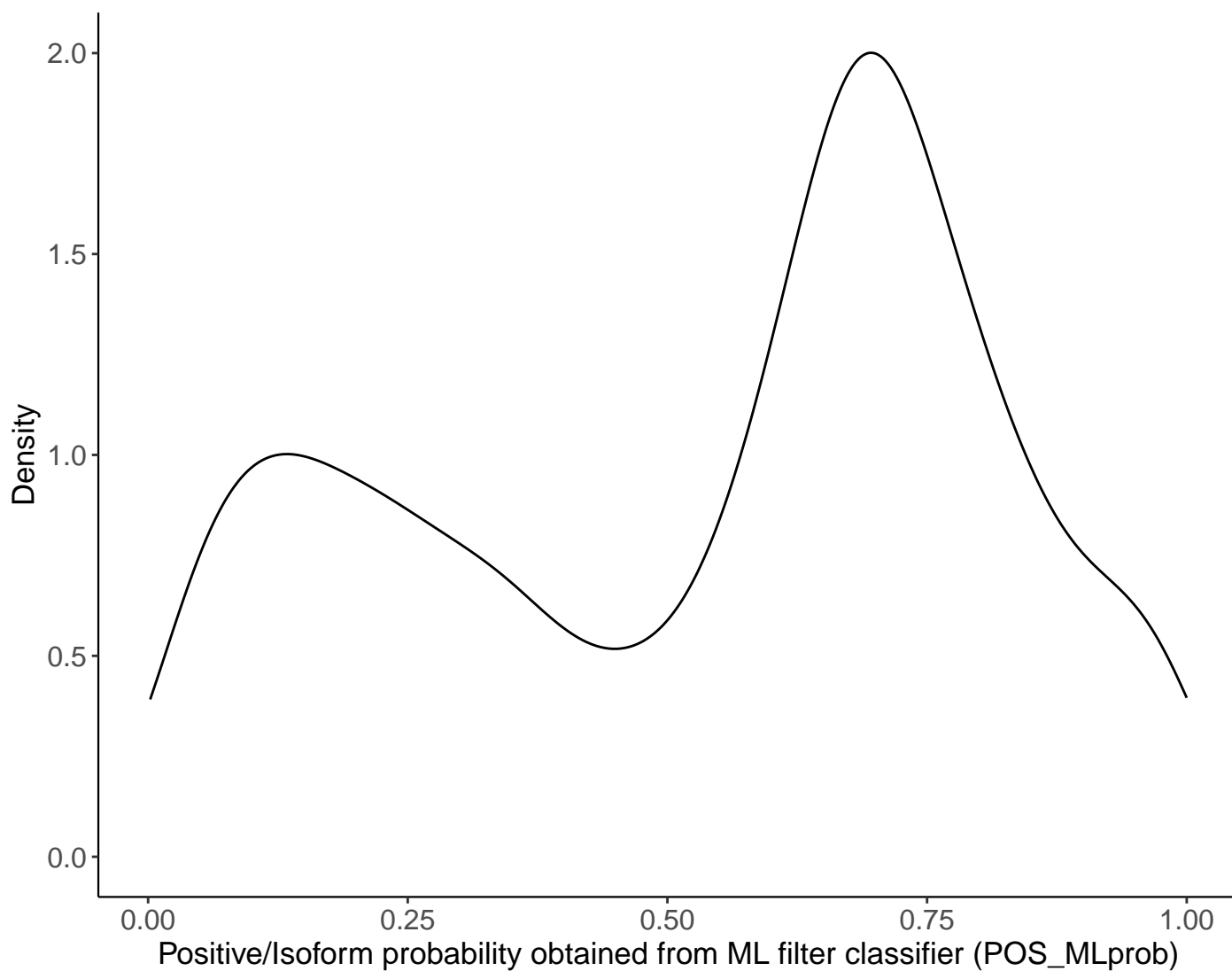
Performance metrics

Metric	Value
Accuracy	9.594595e-01
Kappa	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

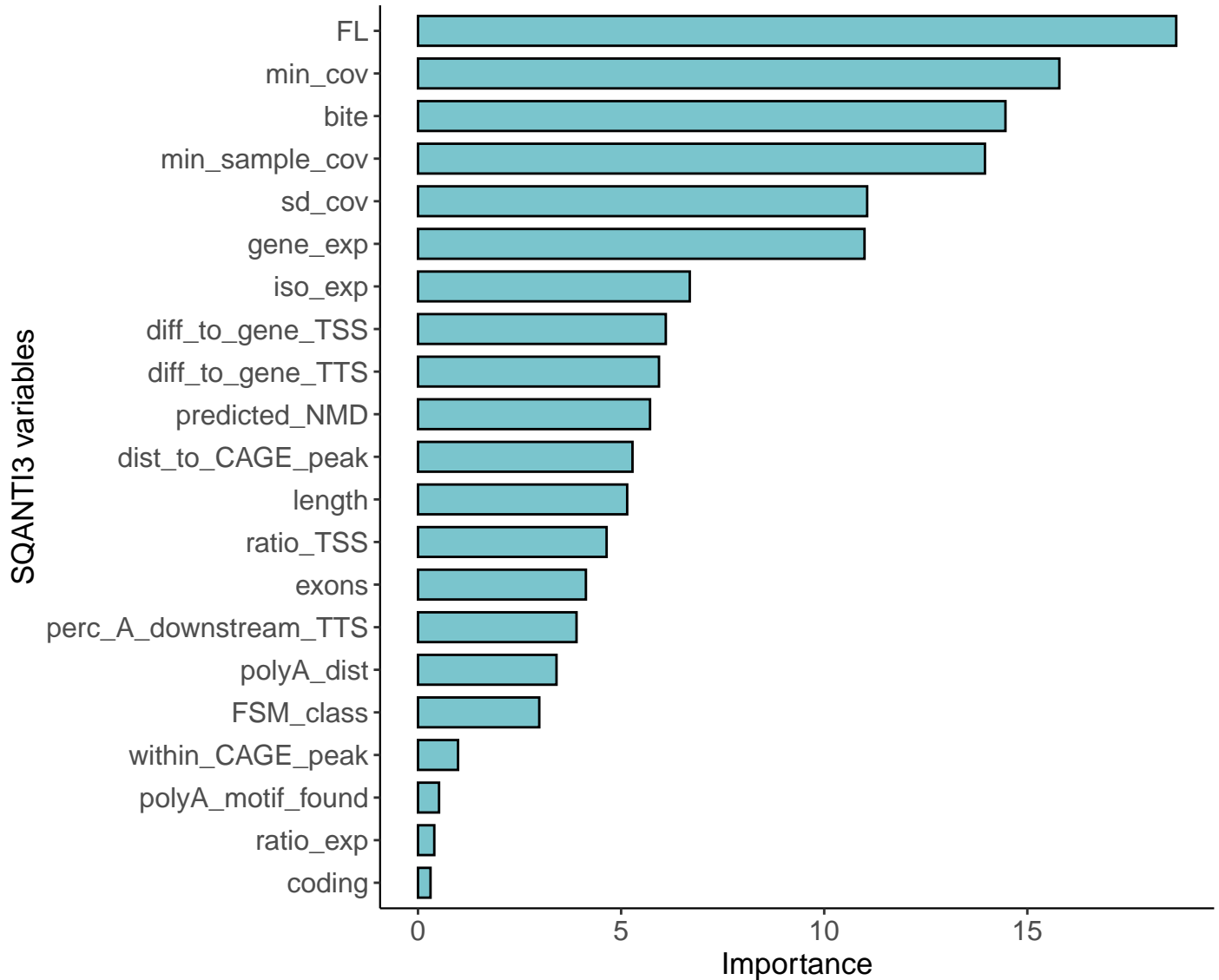
Confusion matrix

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

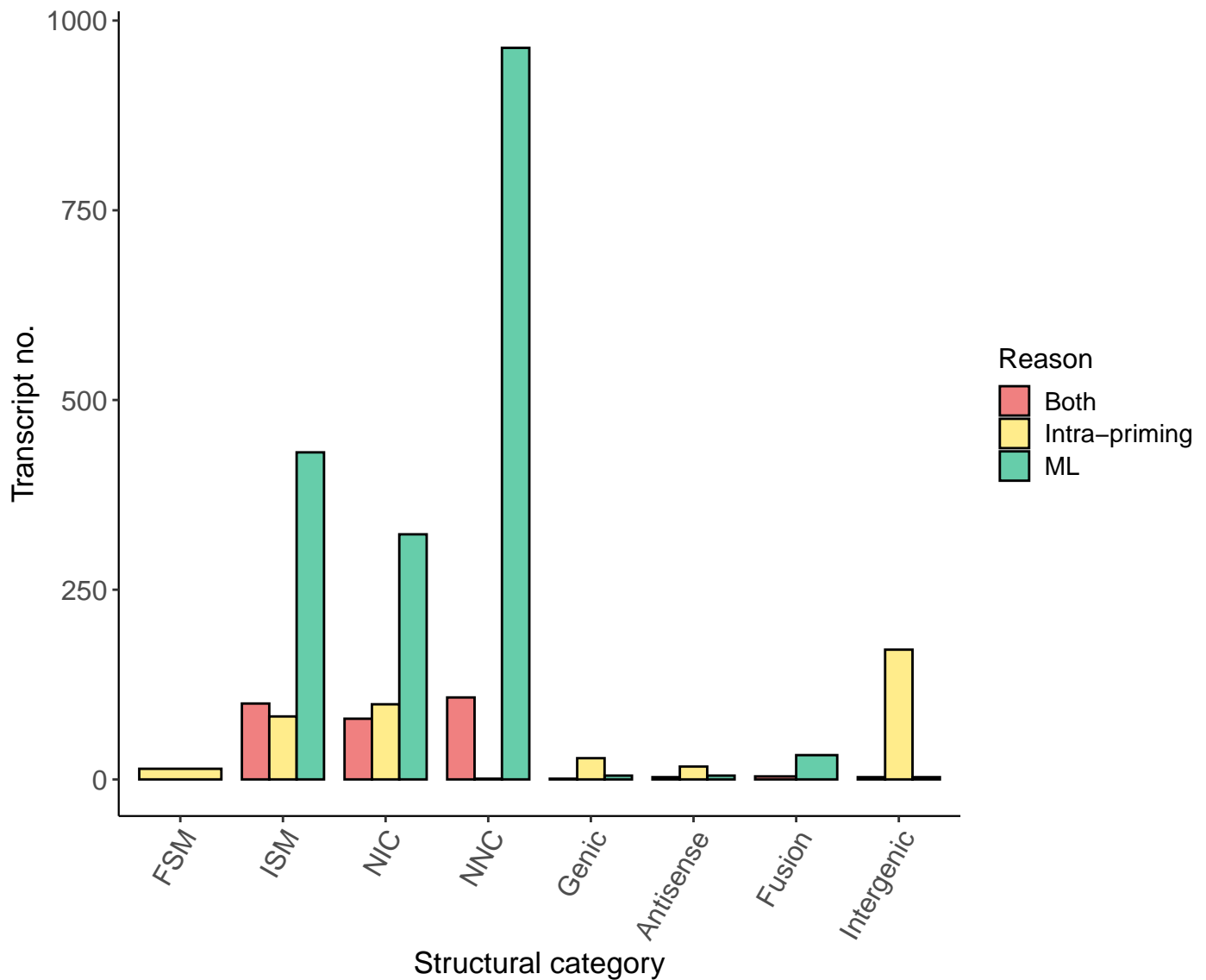
Transcripts classified: 3338



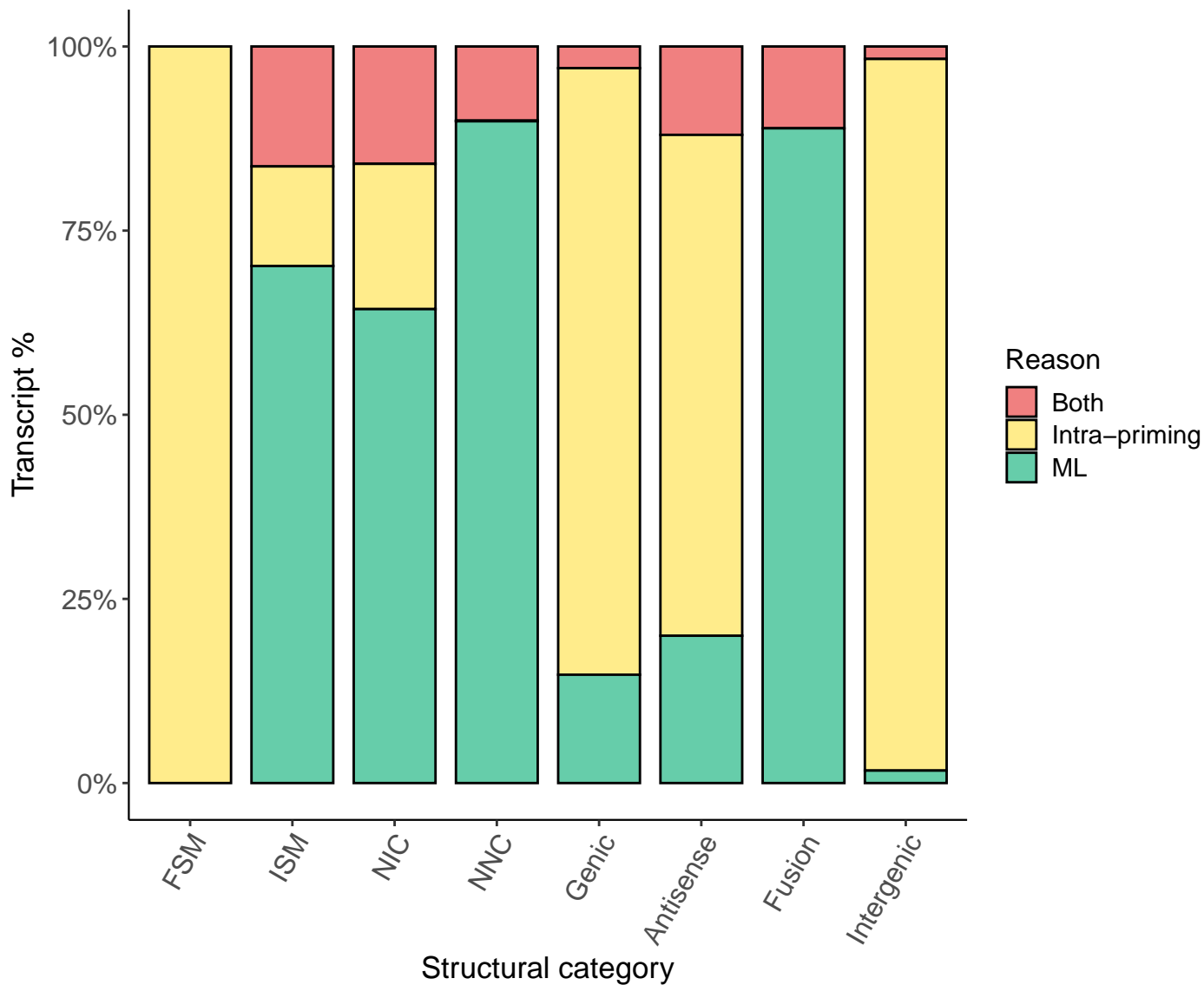
Variable importance in Random Forest classifier



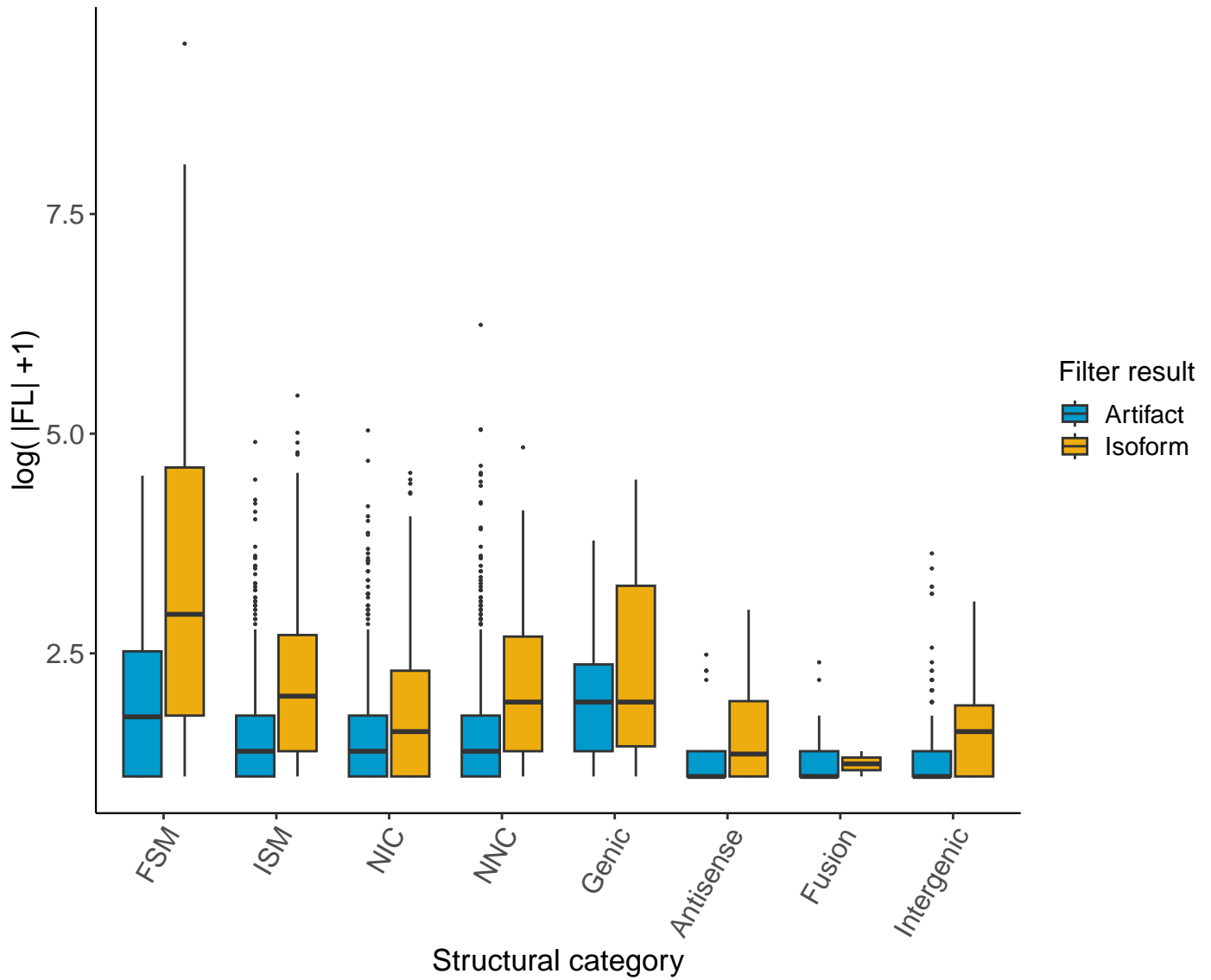
Reason to flag transcripts as artifacts, by category



Reason to flag transcripts as artifacts, by category

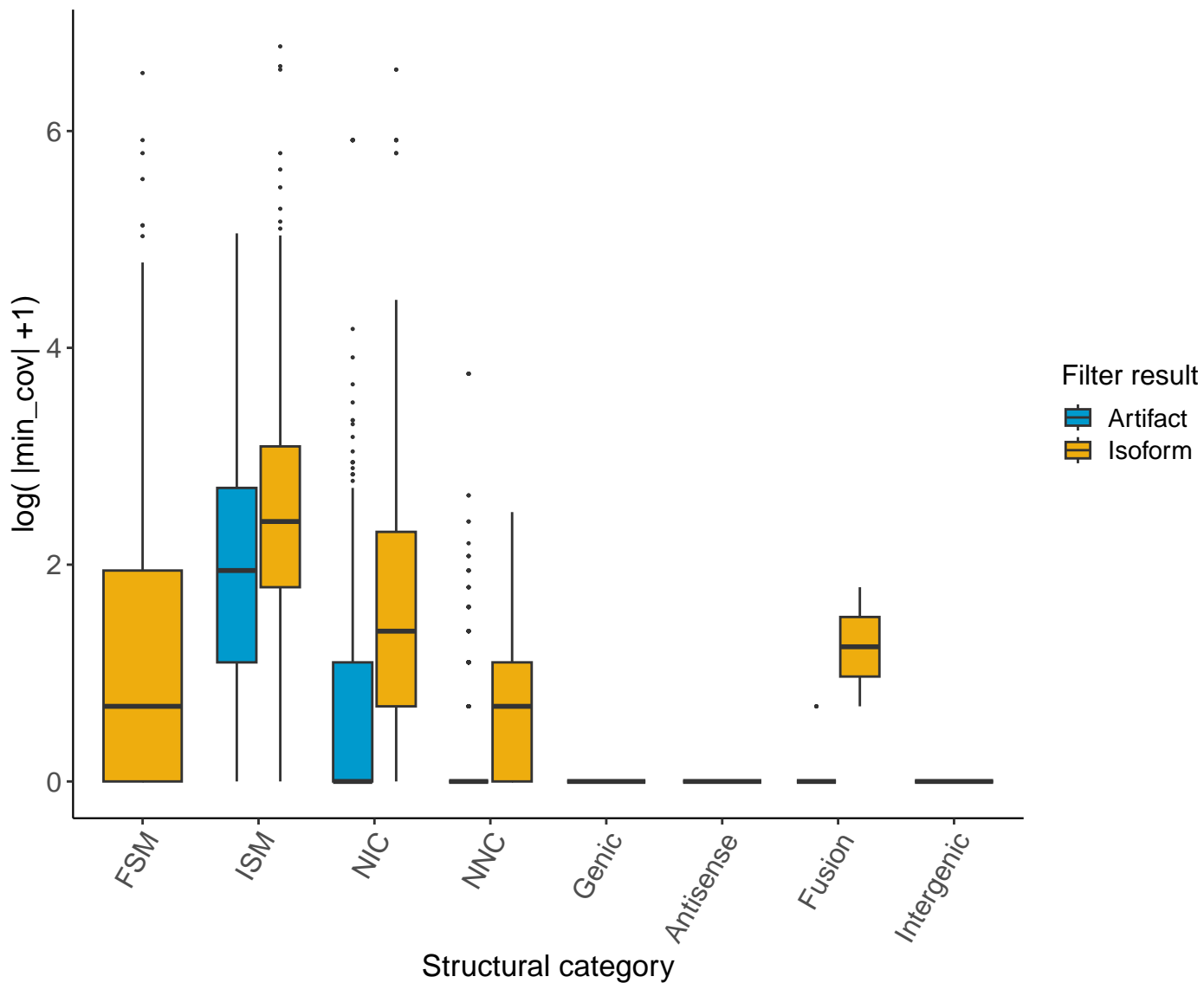


FL – ML importance: 18.67

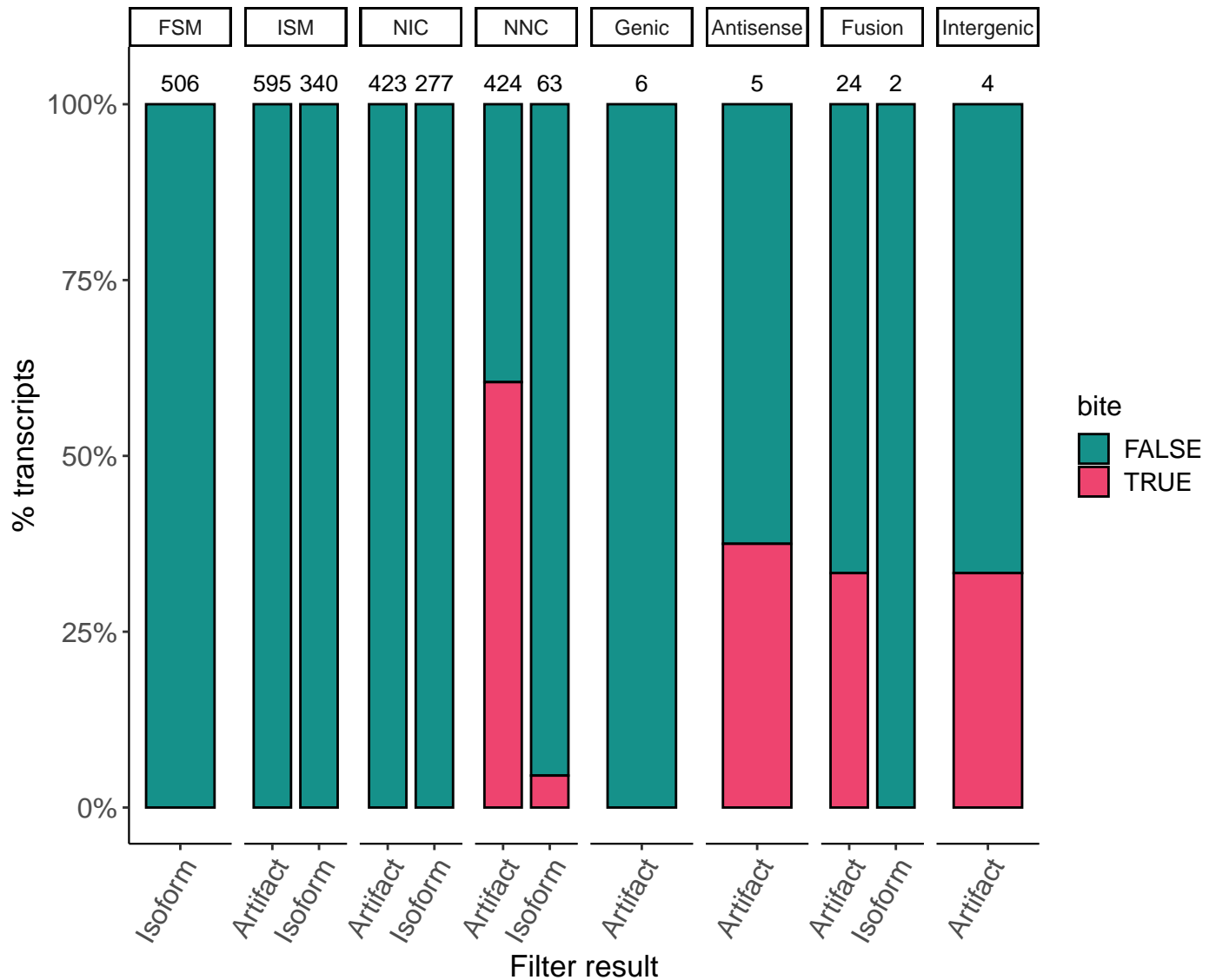




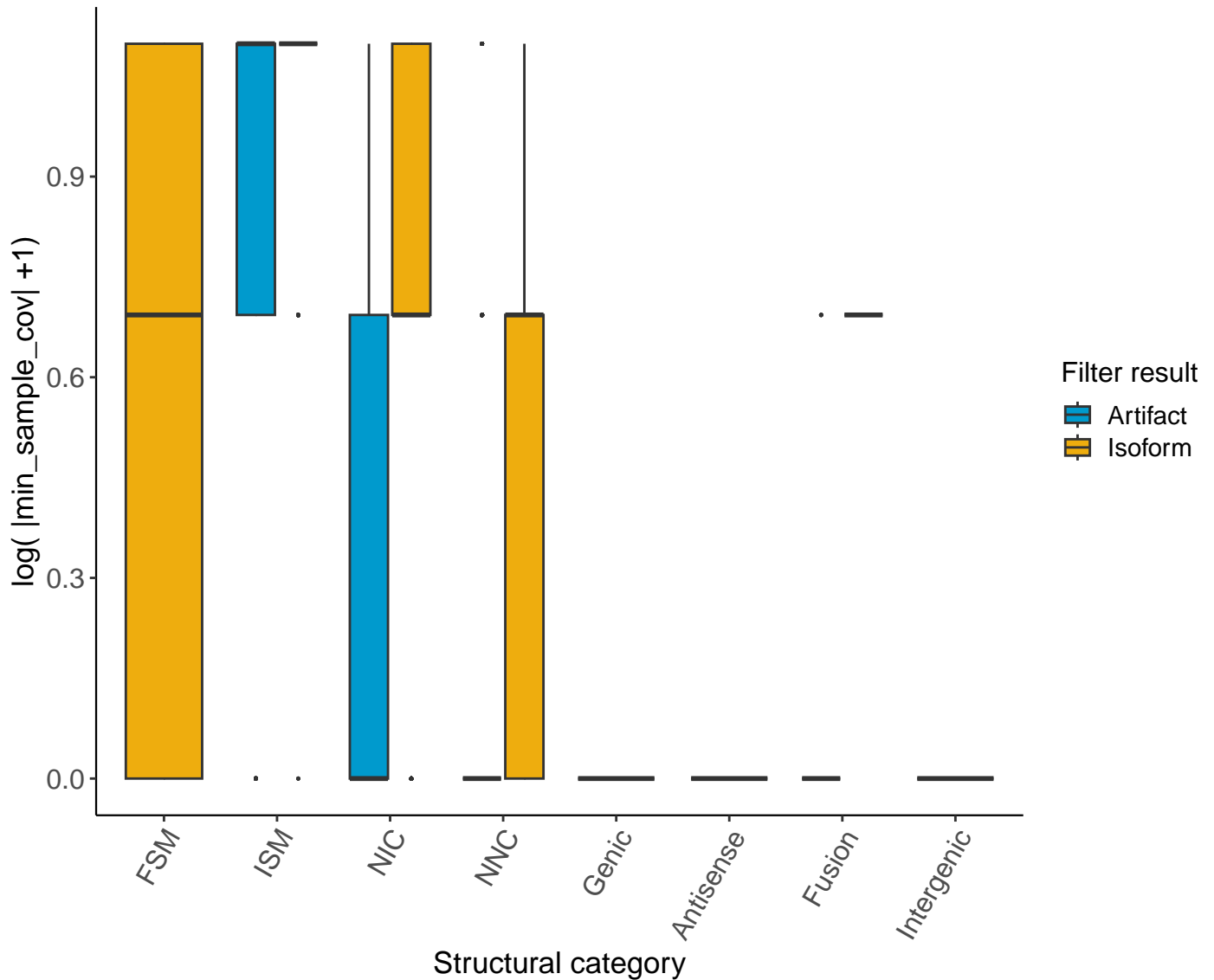
min\_cov – ML importance: 15.79



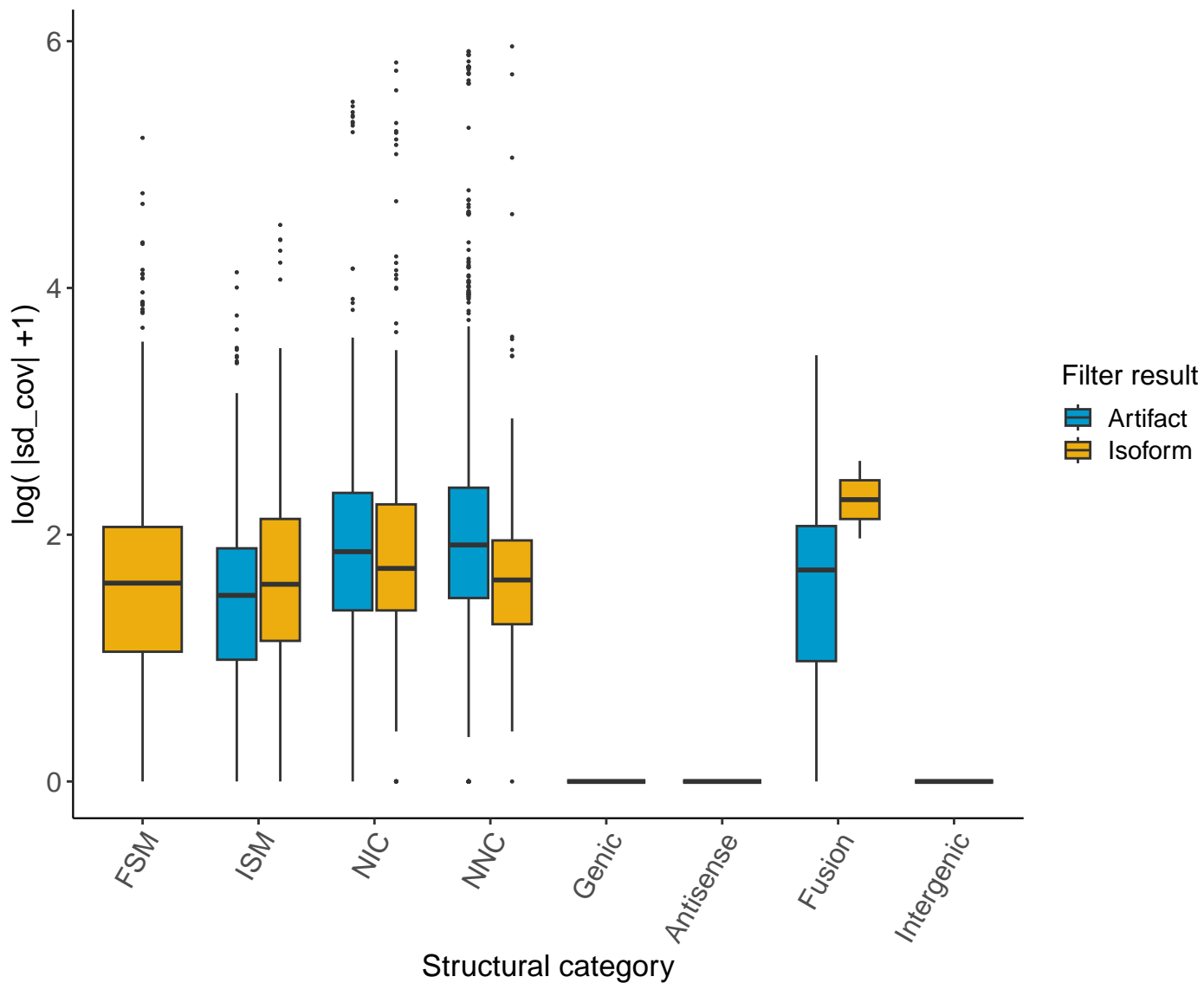
bite – ML importance: 14.46



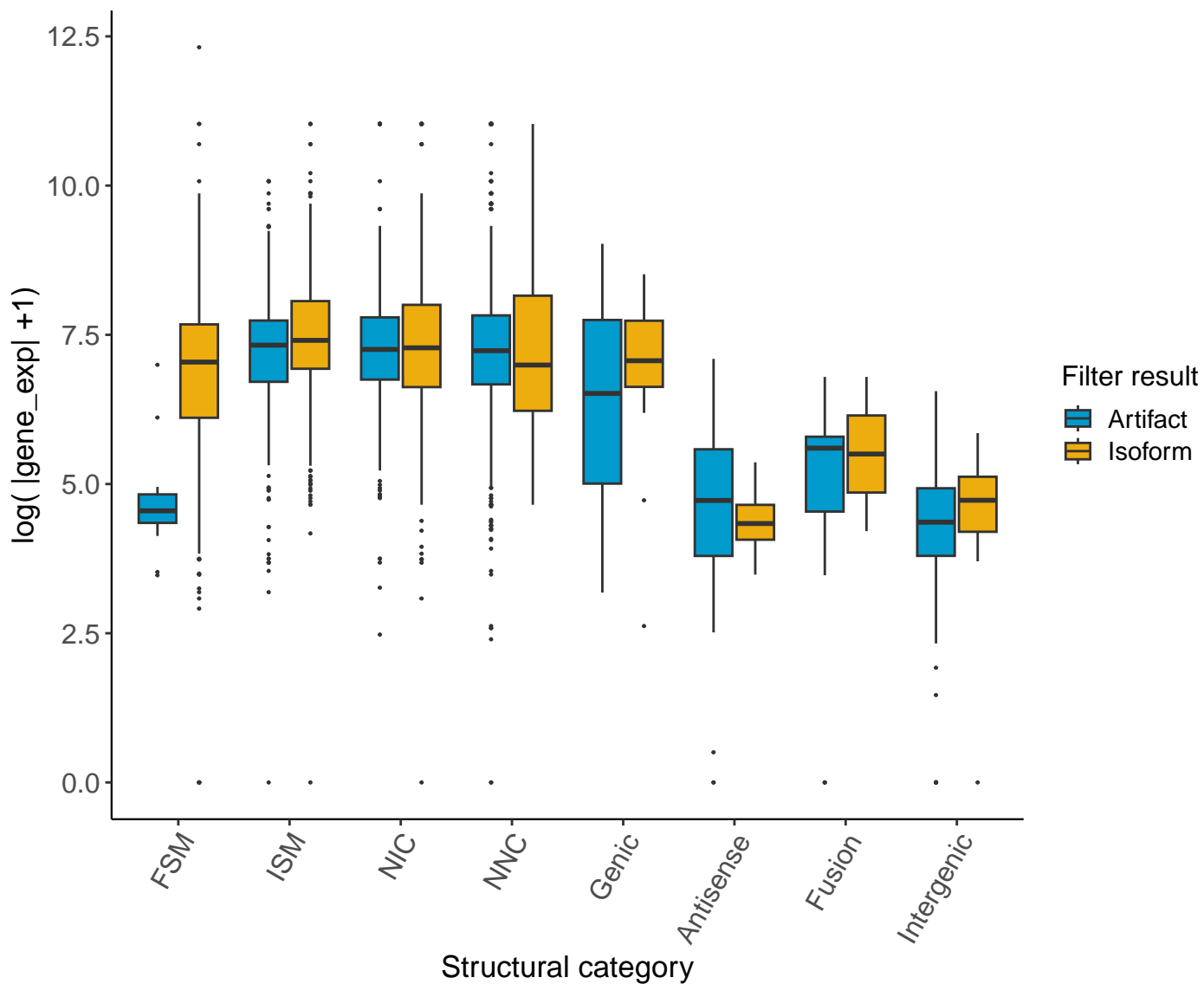
min\_sample\_cov – ML importance: 13.96



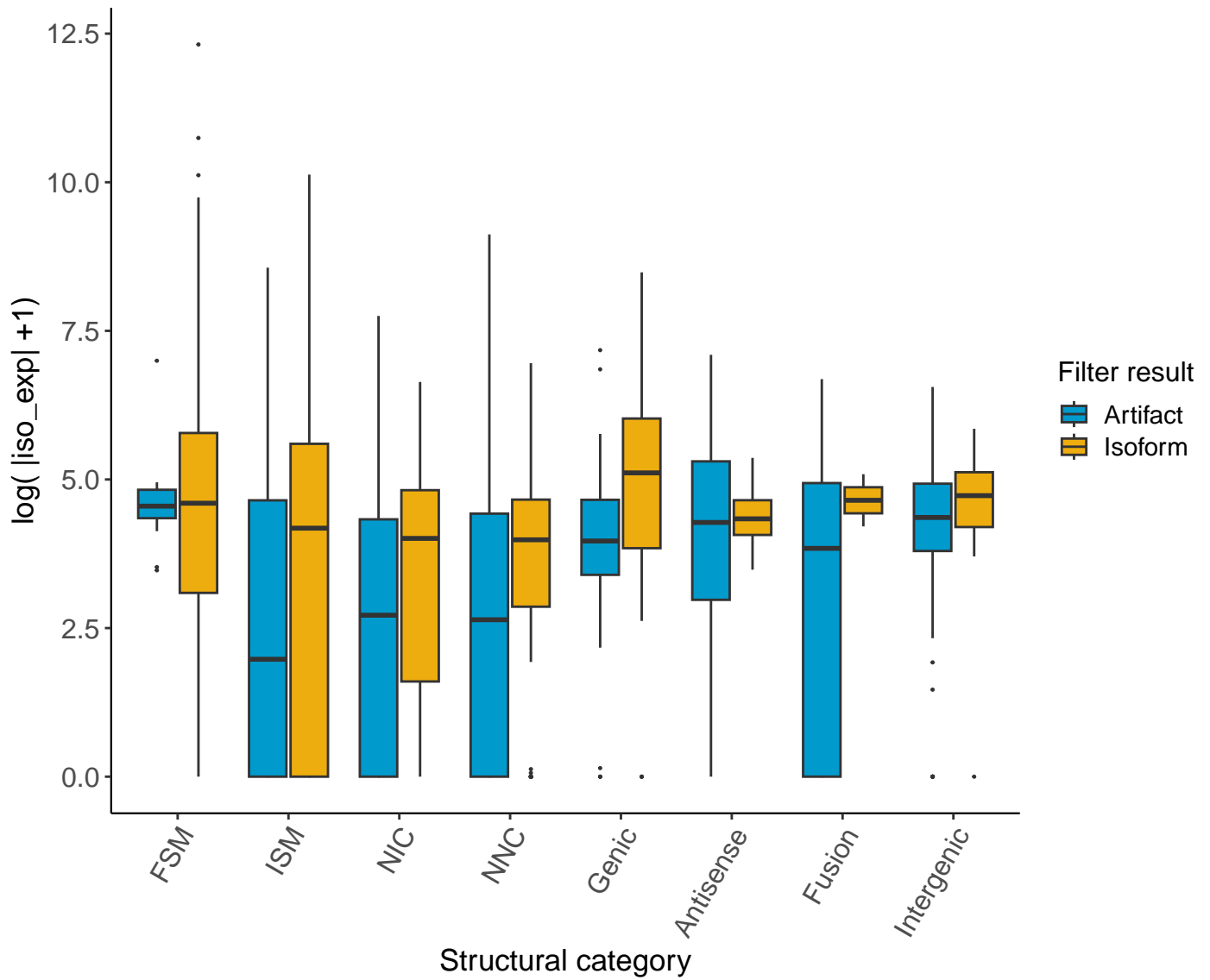
sd\_cov – ML importance: 11.06



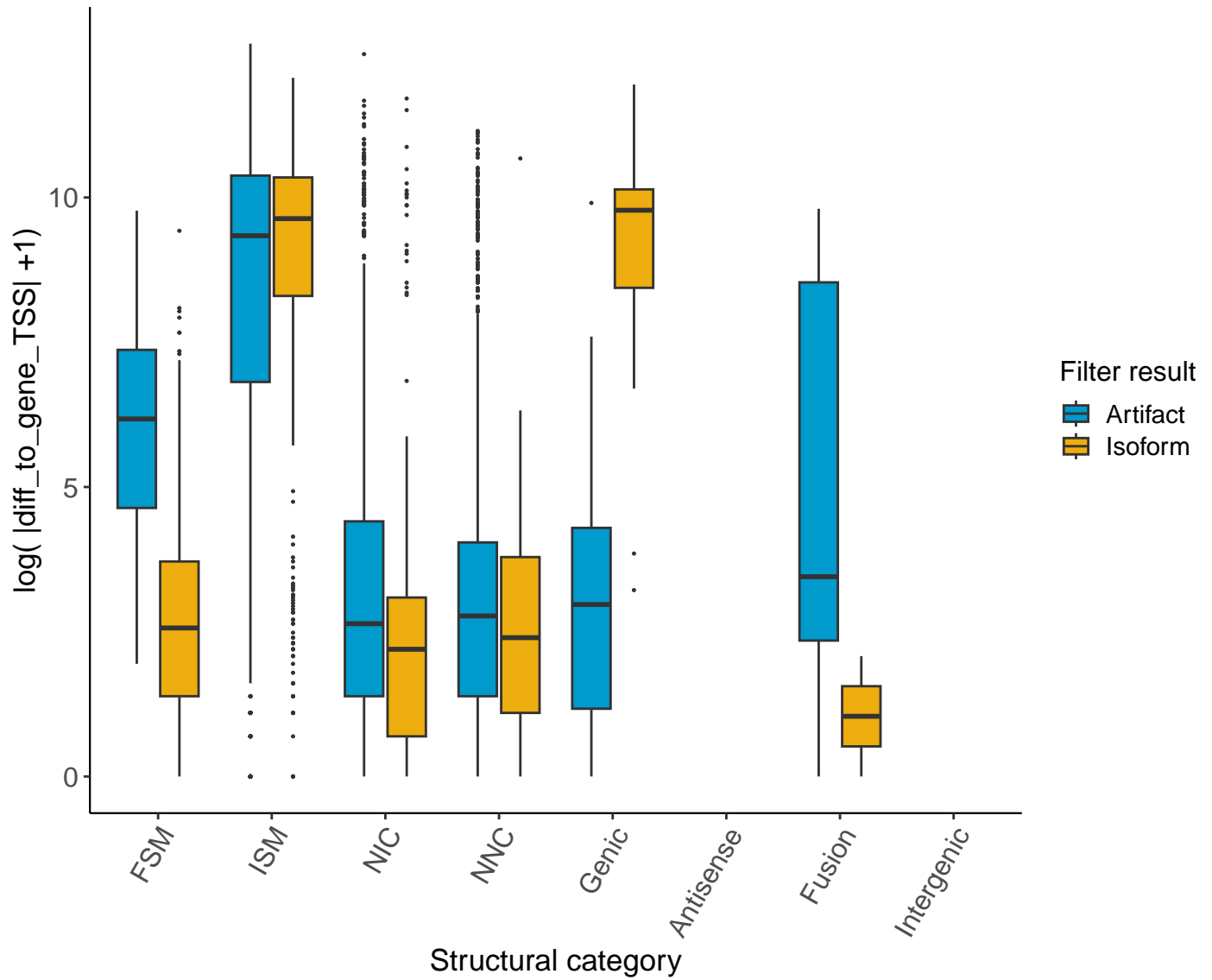
gene\_exp – ML importance: 10.99



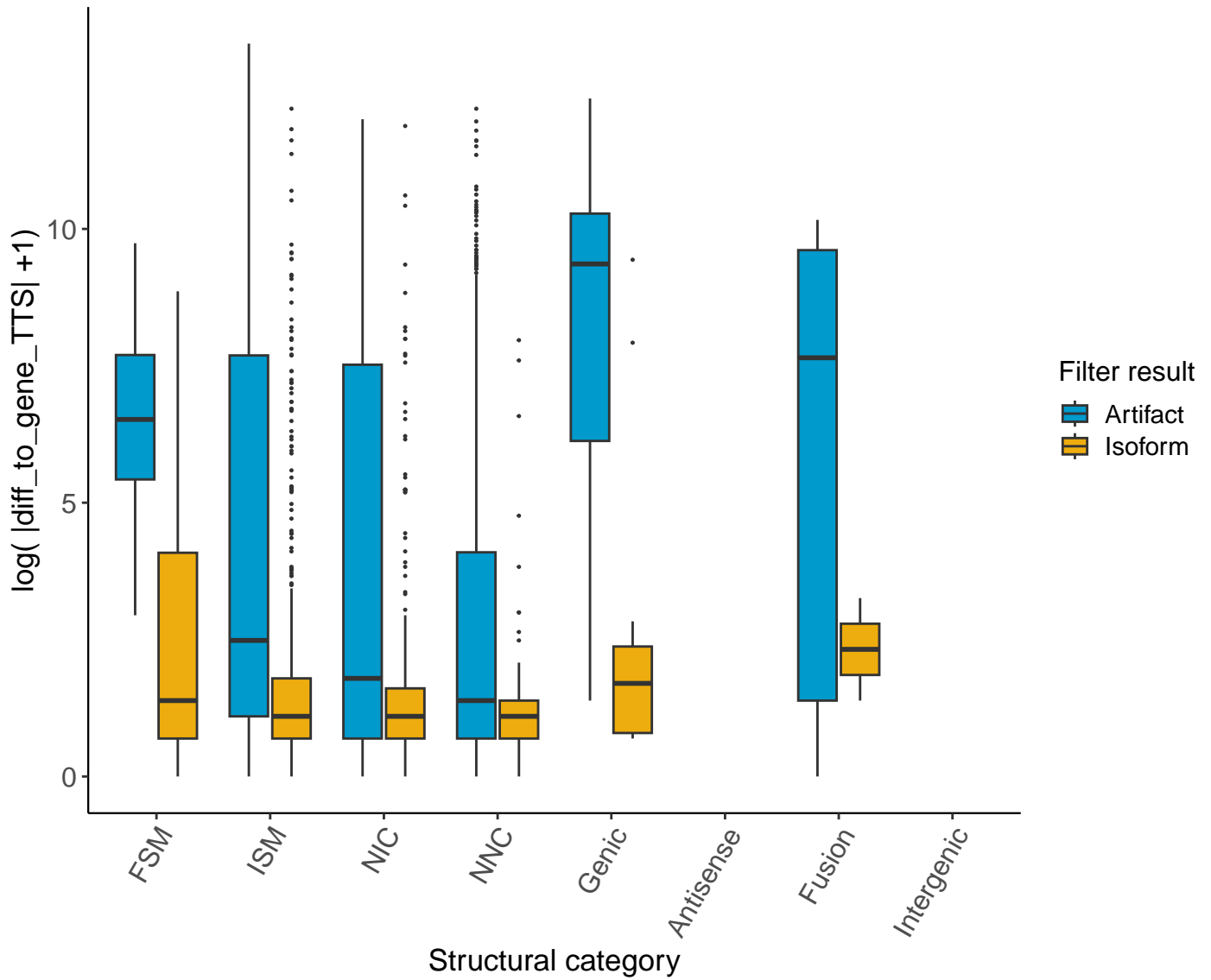
iso\_exp – ML importance: 6.69



diff\_to\_gene\_TSS – ML importance: 6.1

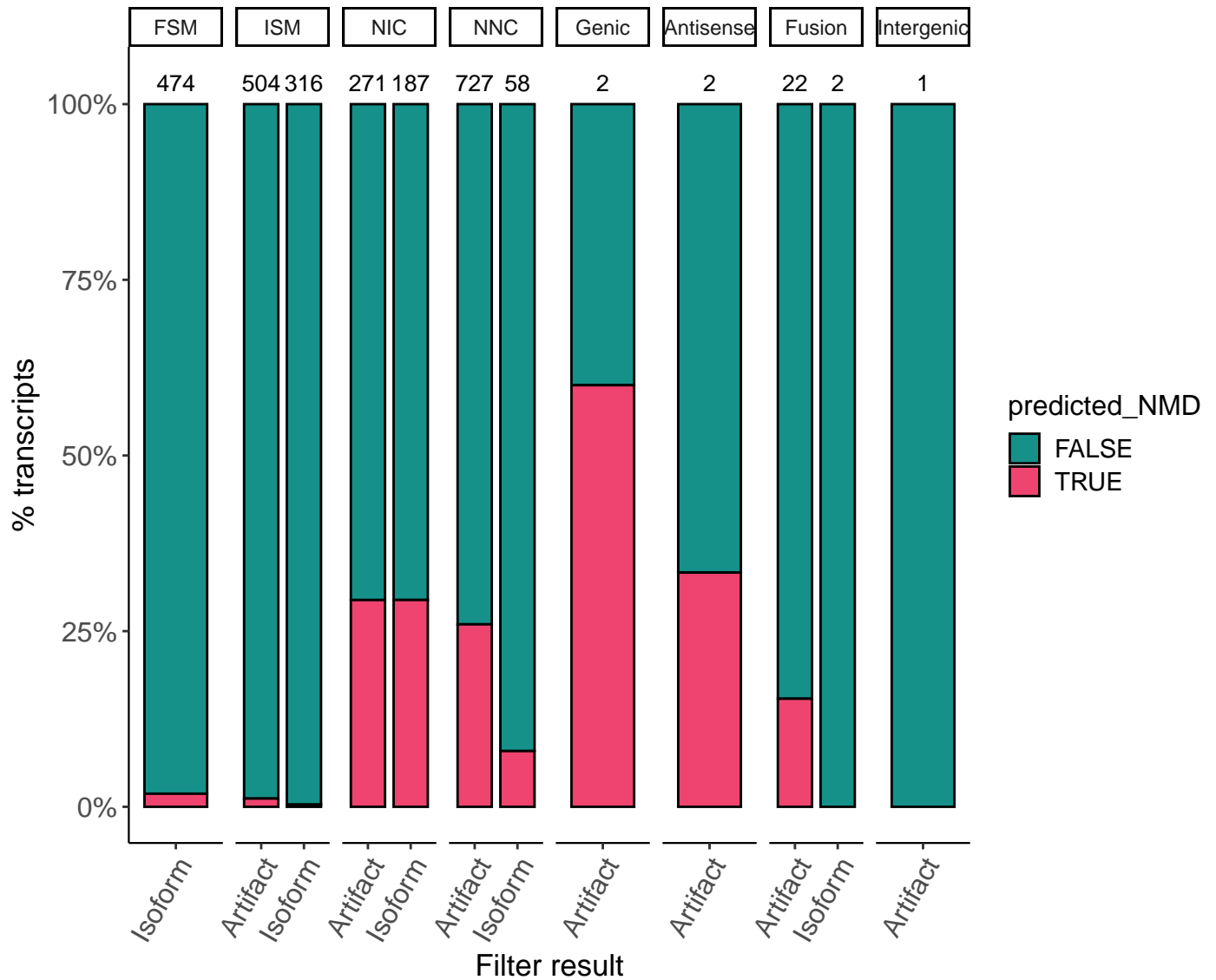


diff\_to\_gene\_TTS – ML importance: 5.93

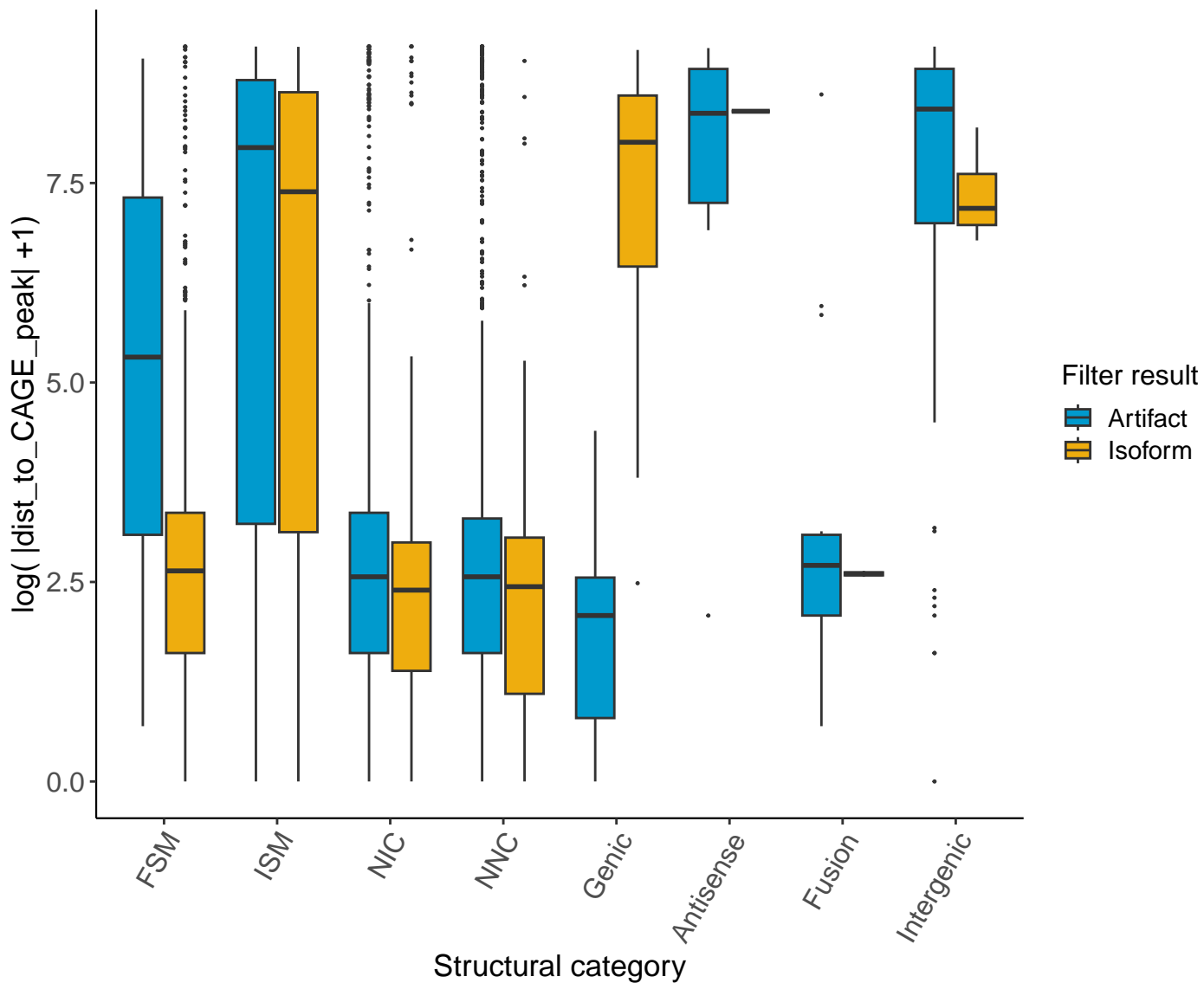




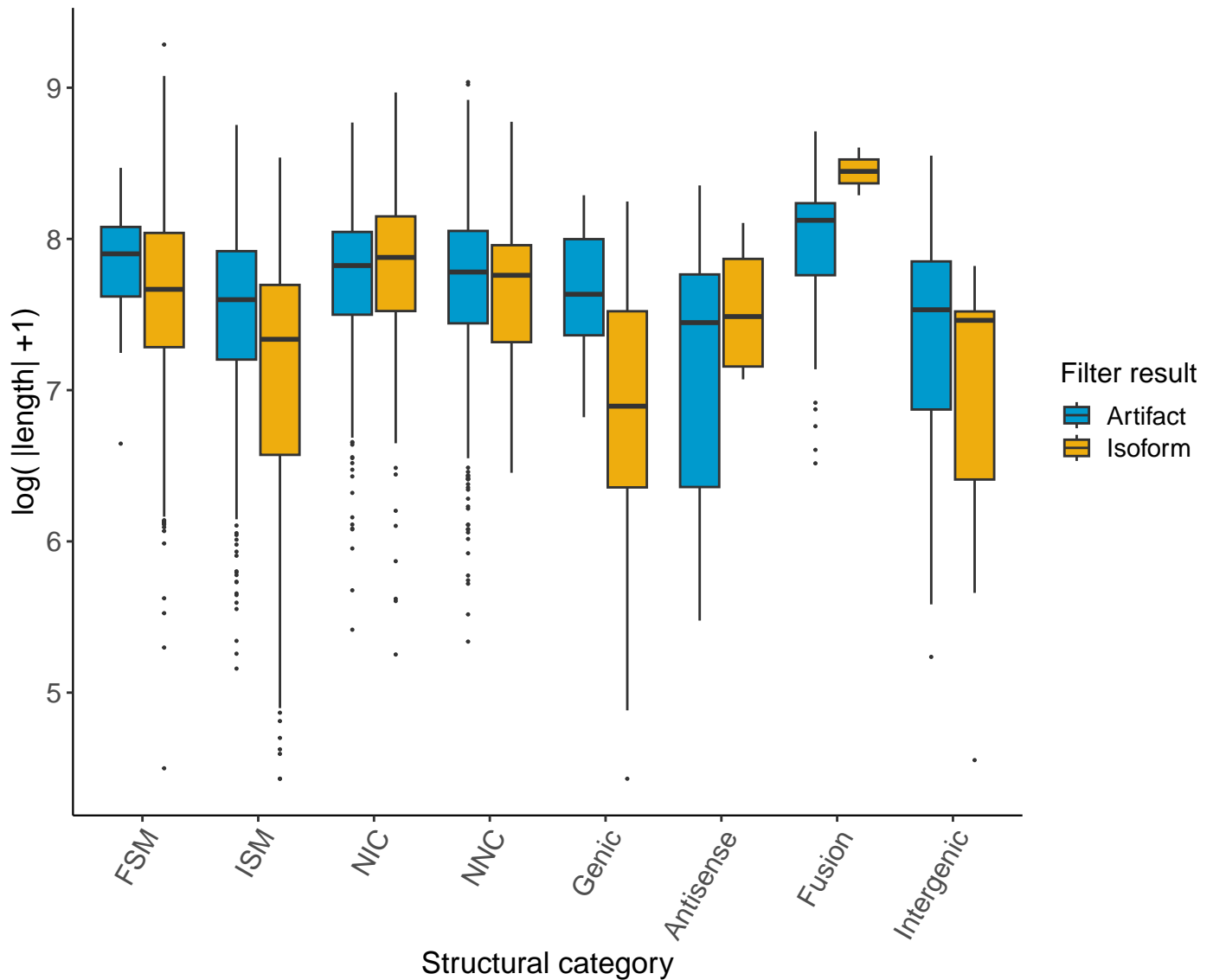
predicted\_NMD – ML importance: 5.71



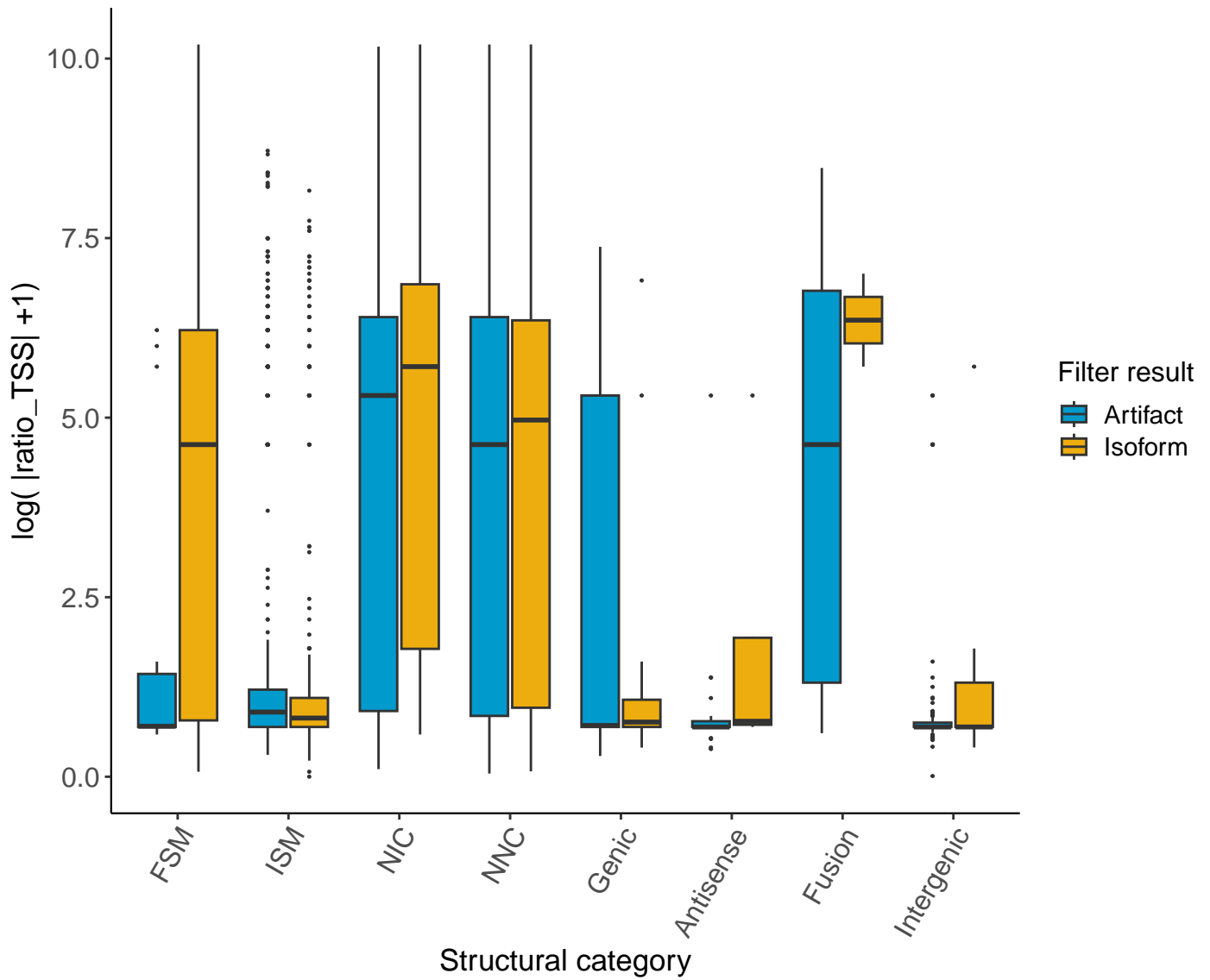
dist\_to\_CAGE\_peak – ML importance: 5.28



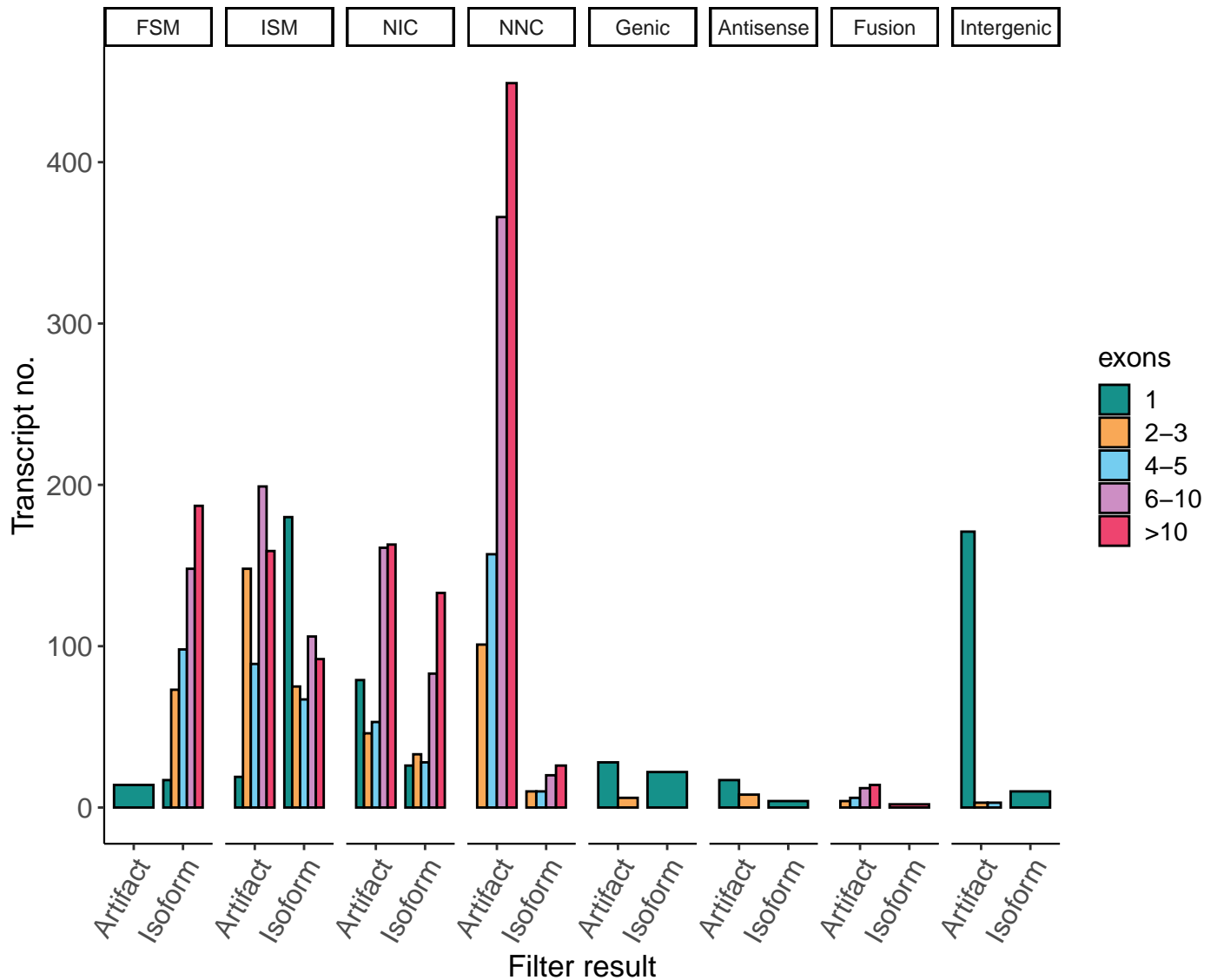
length – ML importance: 5.15



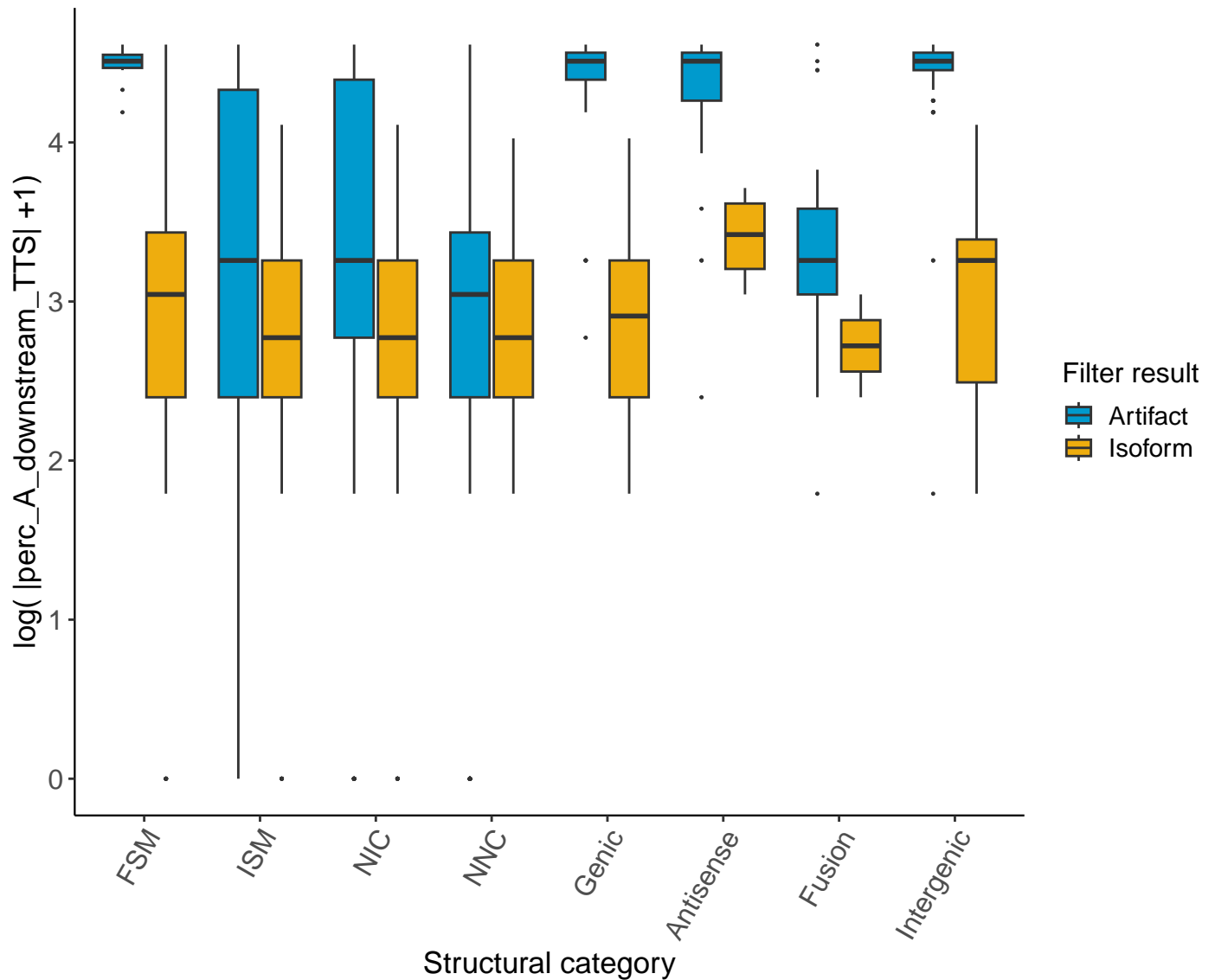
ratio\_TSS – ML importance: 4.64



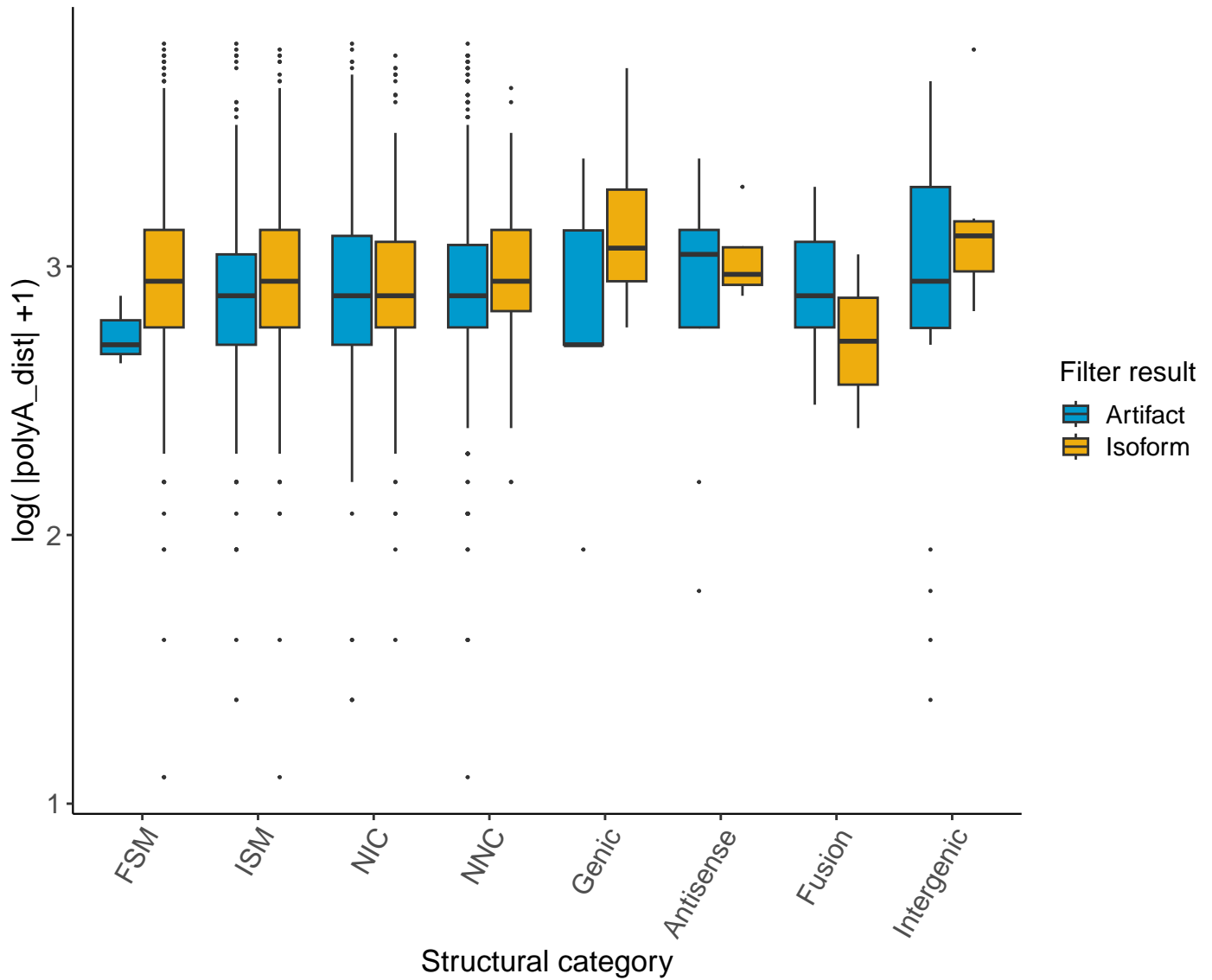
# exons – ML importance: 4.13



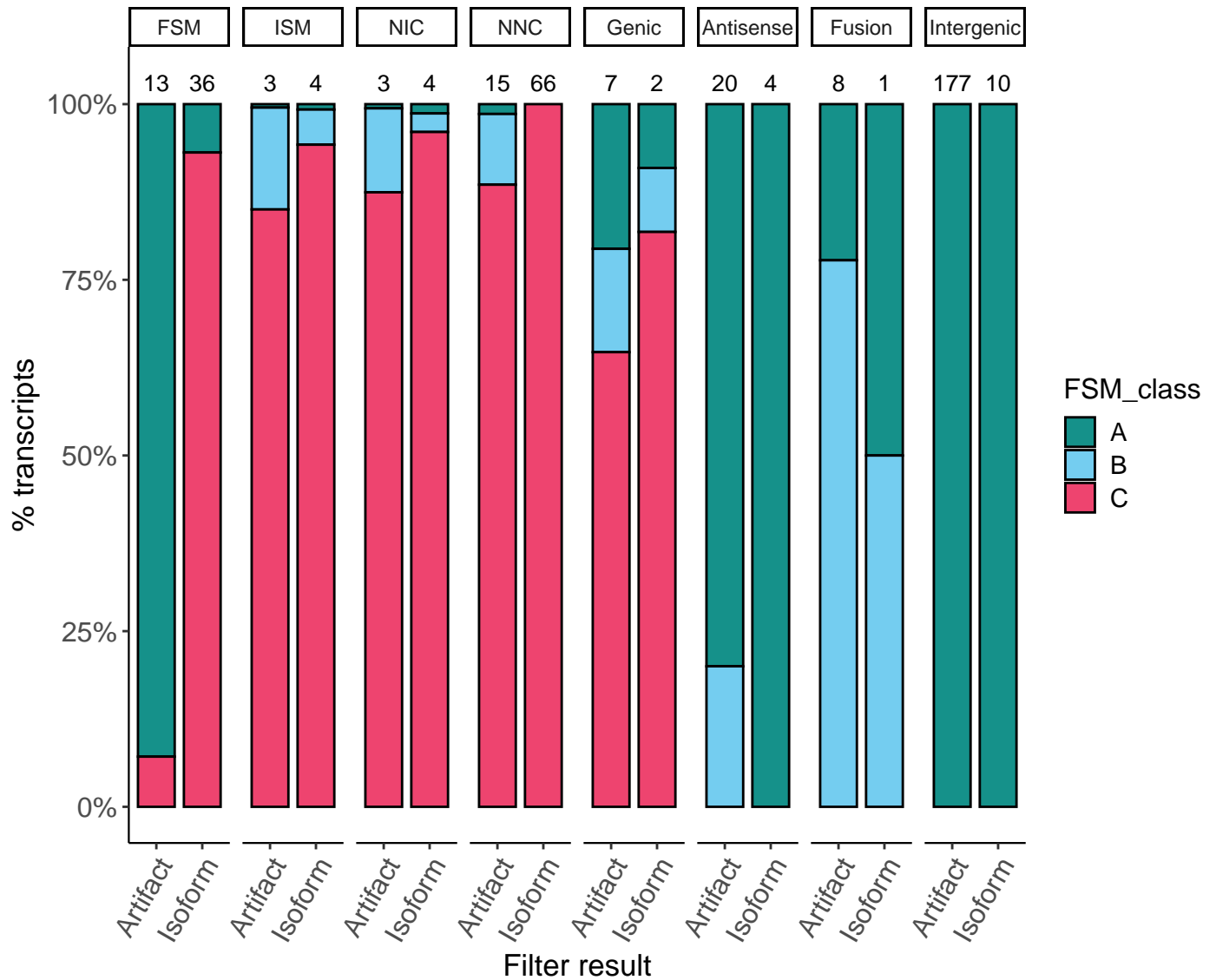
perc\_A\_downstream\_TTS – ML importance: 3.9



polyA\_dist – ML importance: 3.41

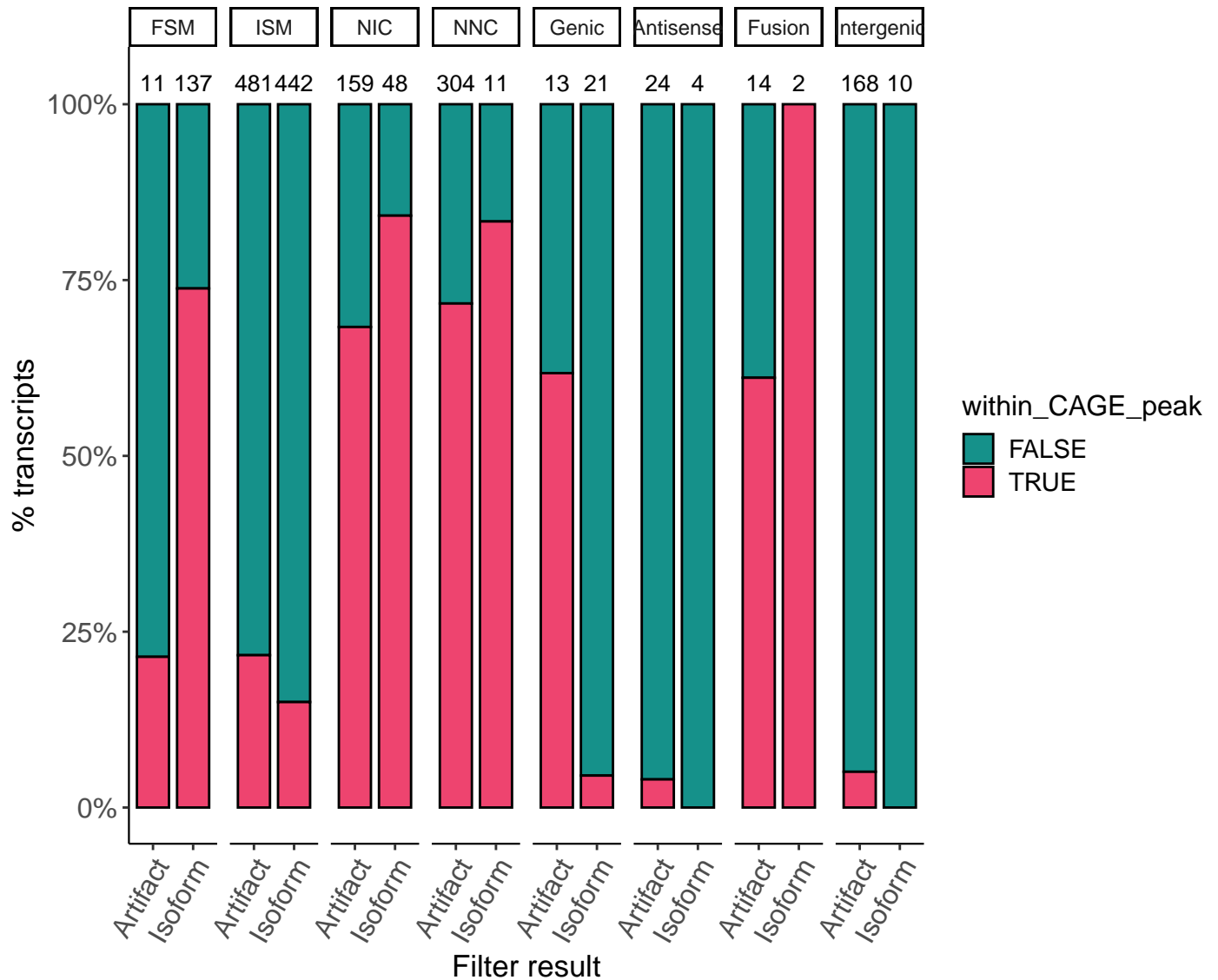


# FSM\_class – ML importance: 2.99

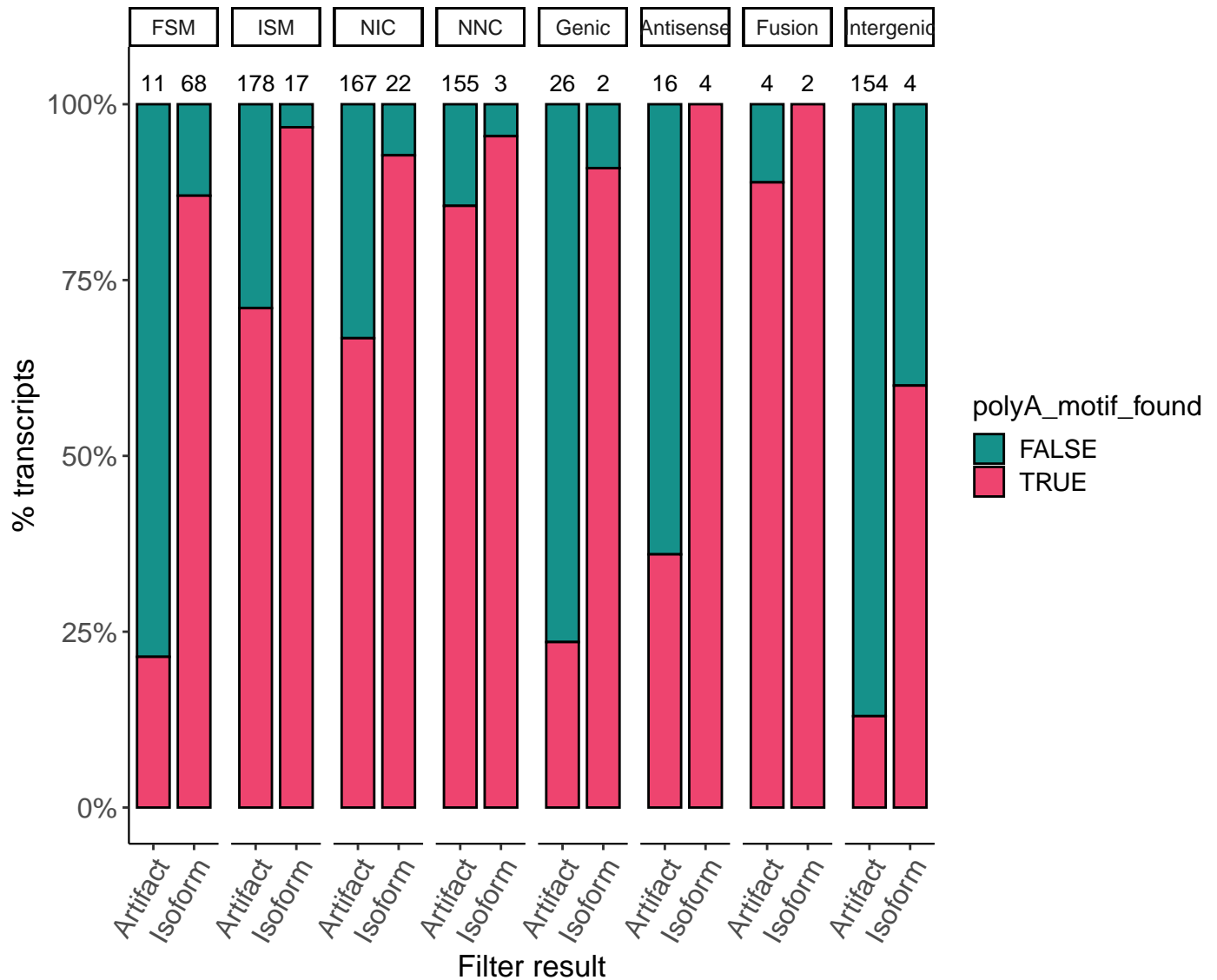




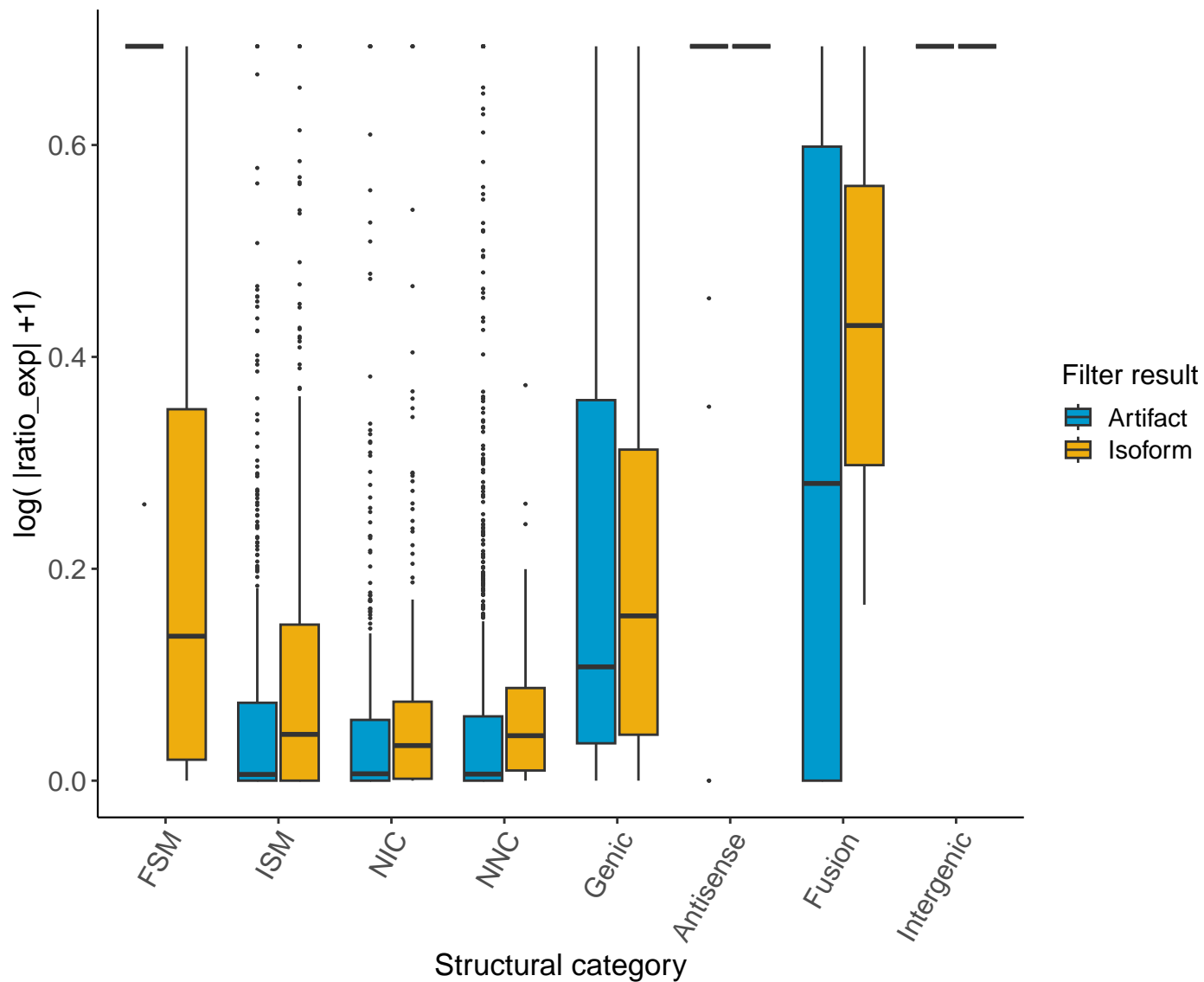
within\_CAGE\_peak – ML importance: 0.99



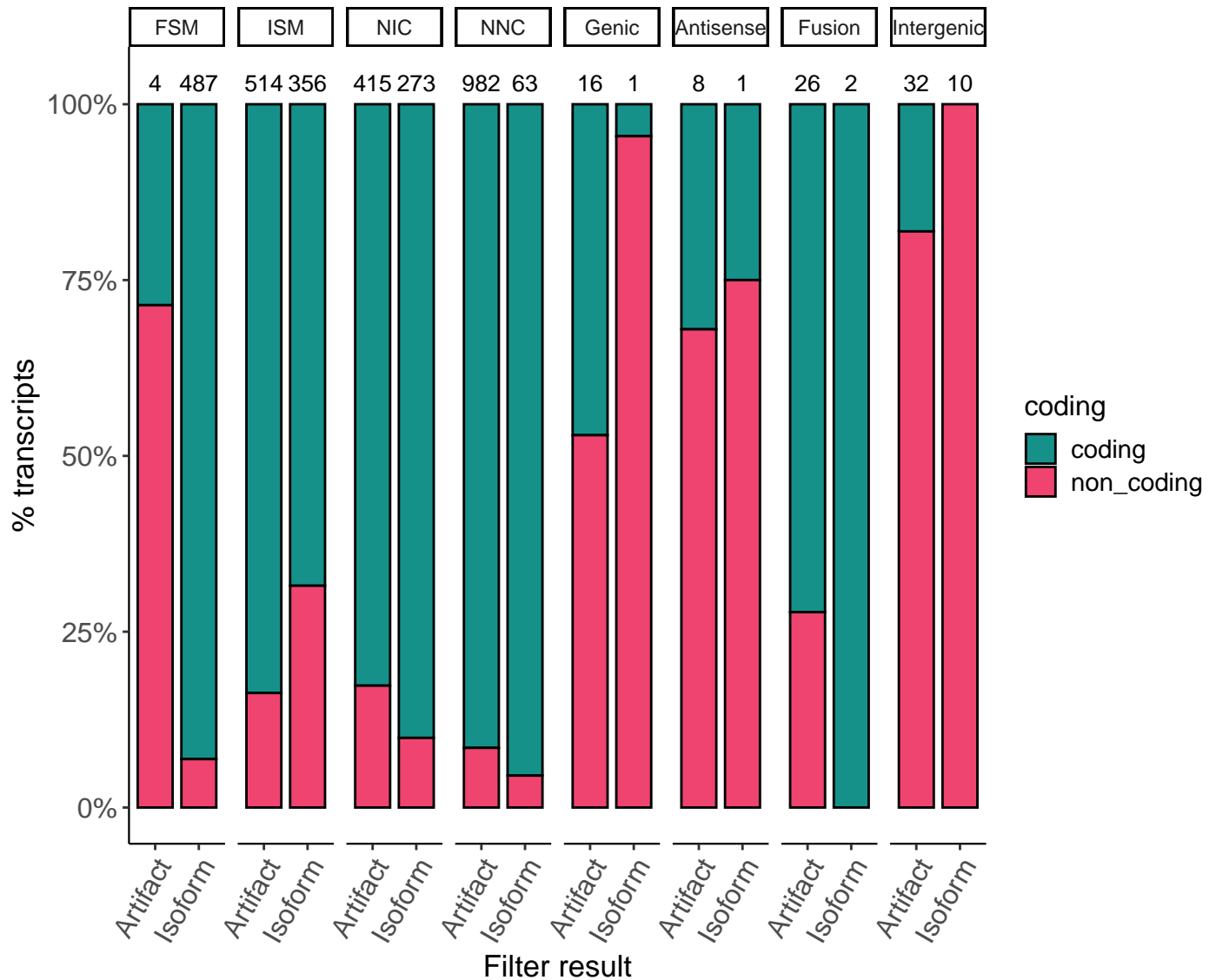
polyA\_motif\_found – ML importance: 0.52



ratio\_exp – ML importance: 0.4



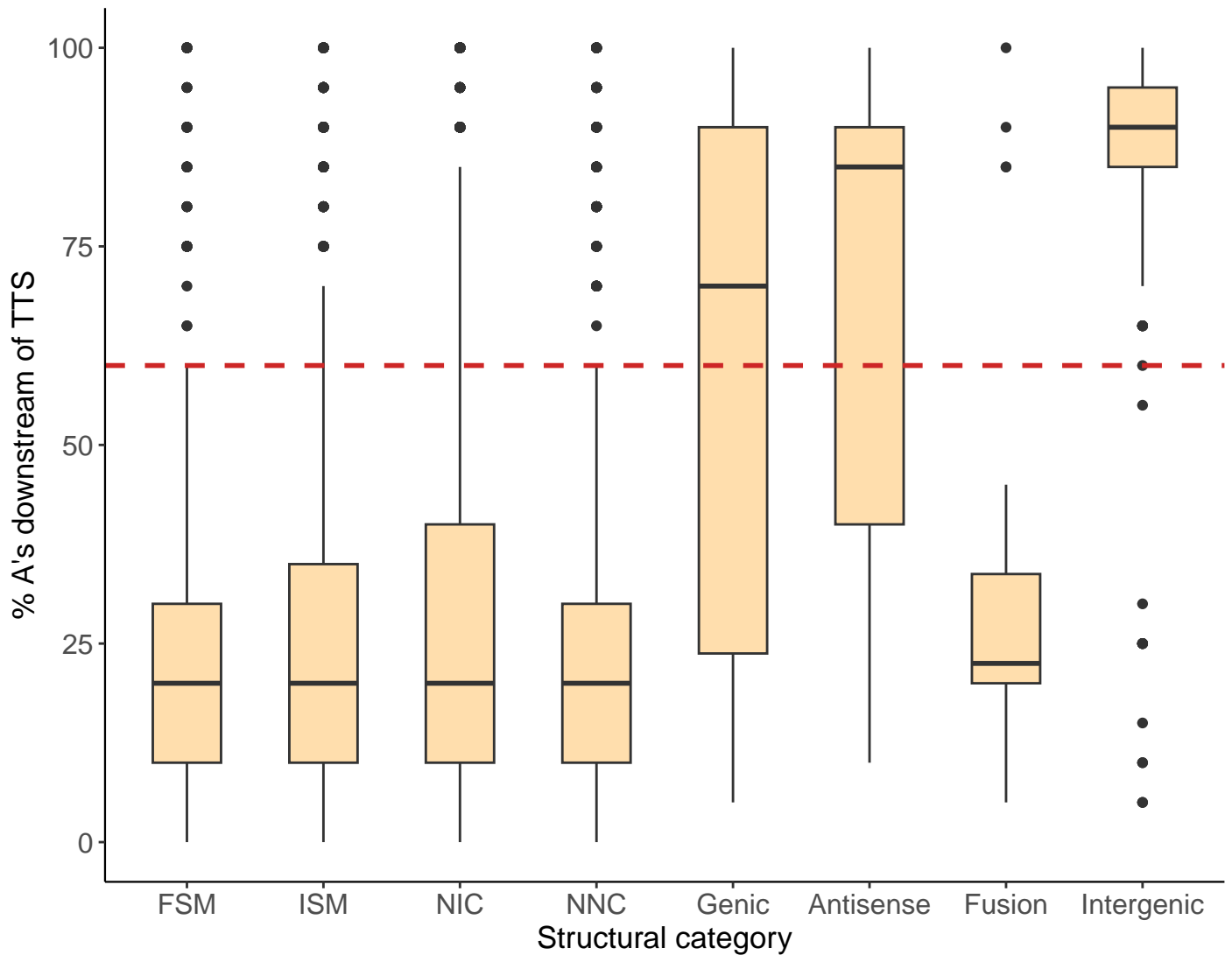
coding – ML importance: 0.31



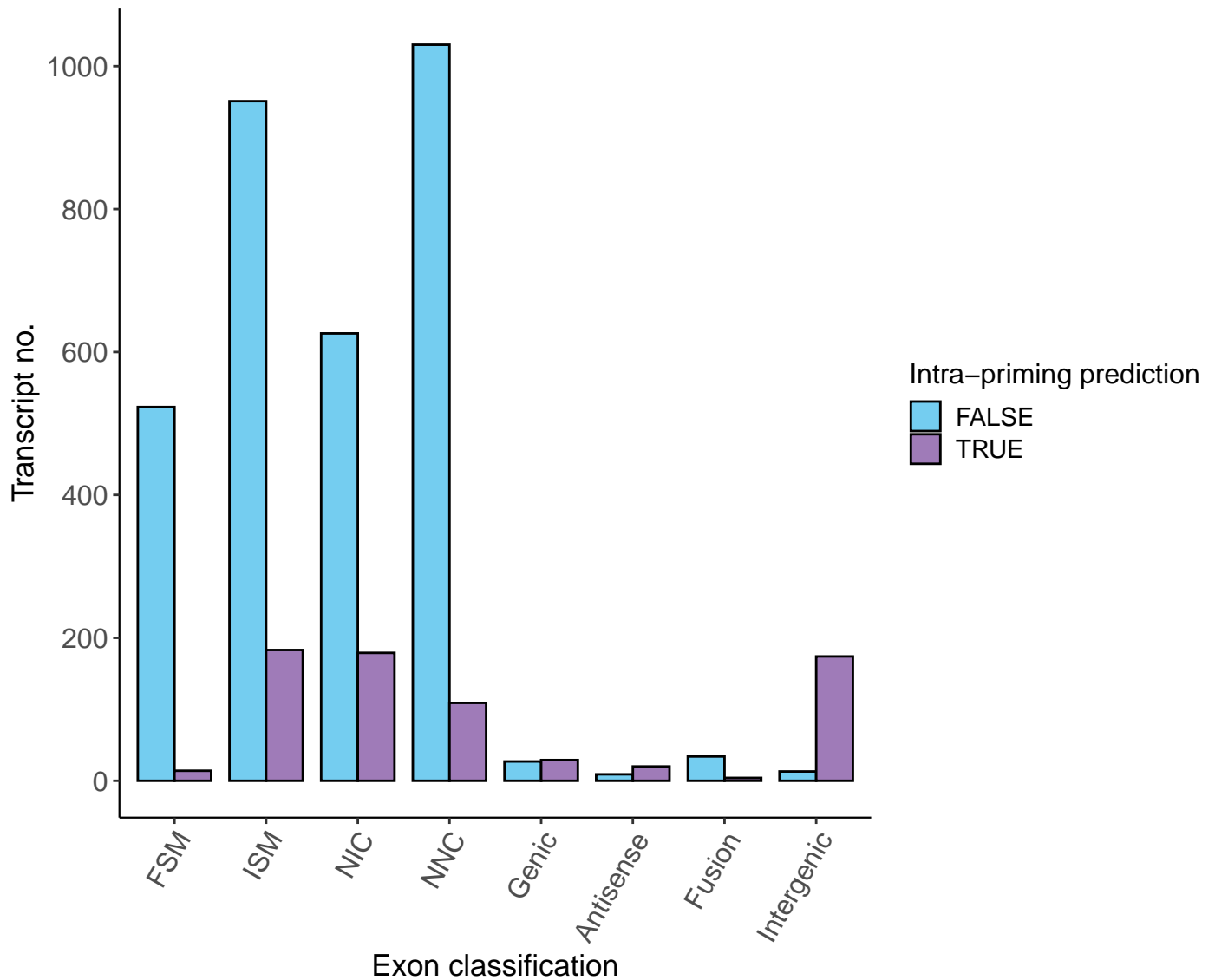
## *Intra-priming filter report*

## A % by category

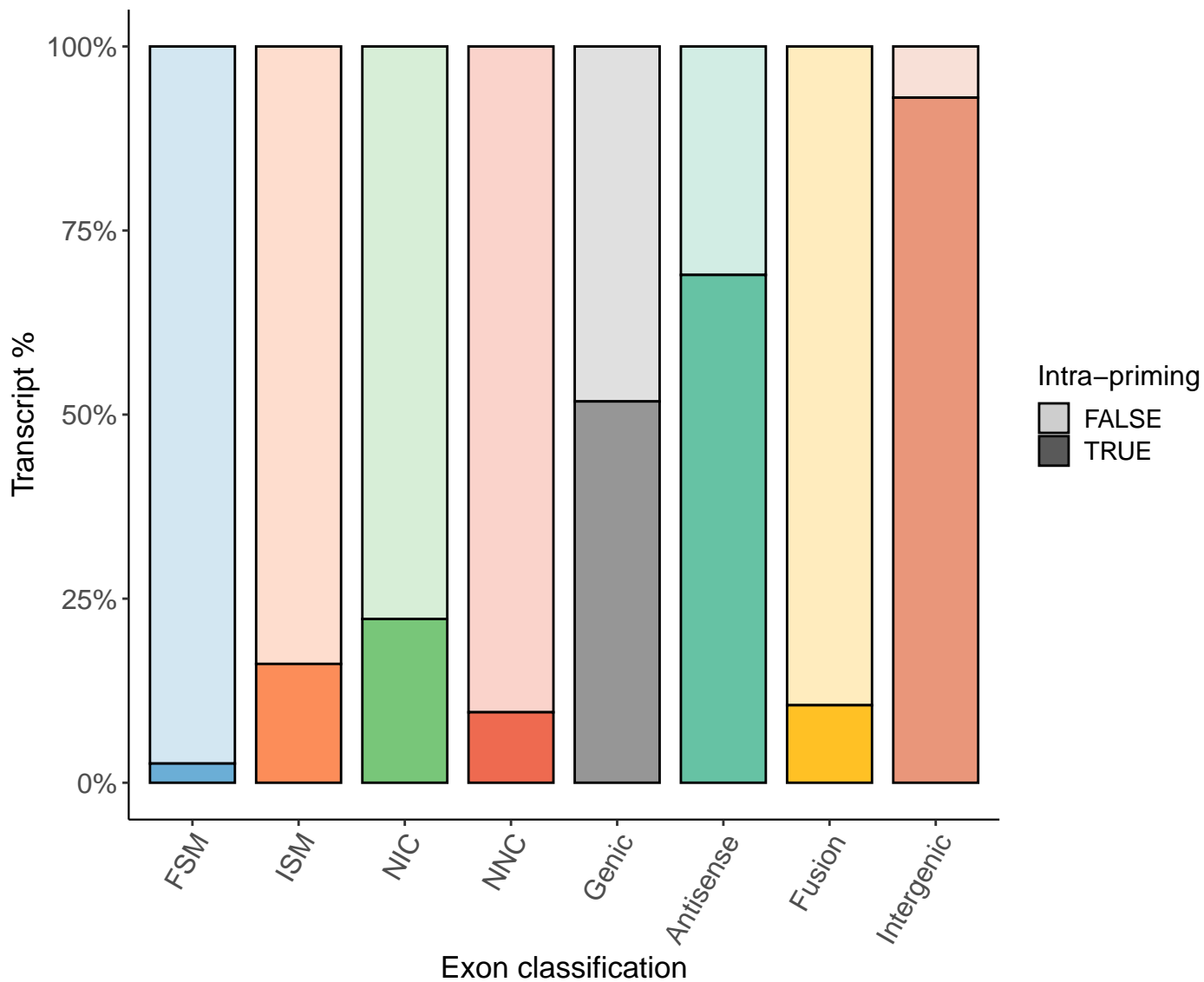
Red line indicates threshold employed in ML filter



Isoforms flagged as intra-priming, by category

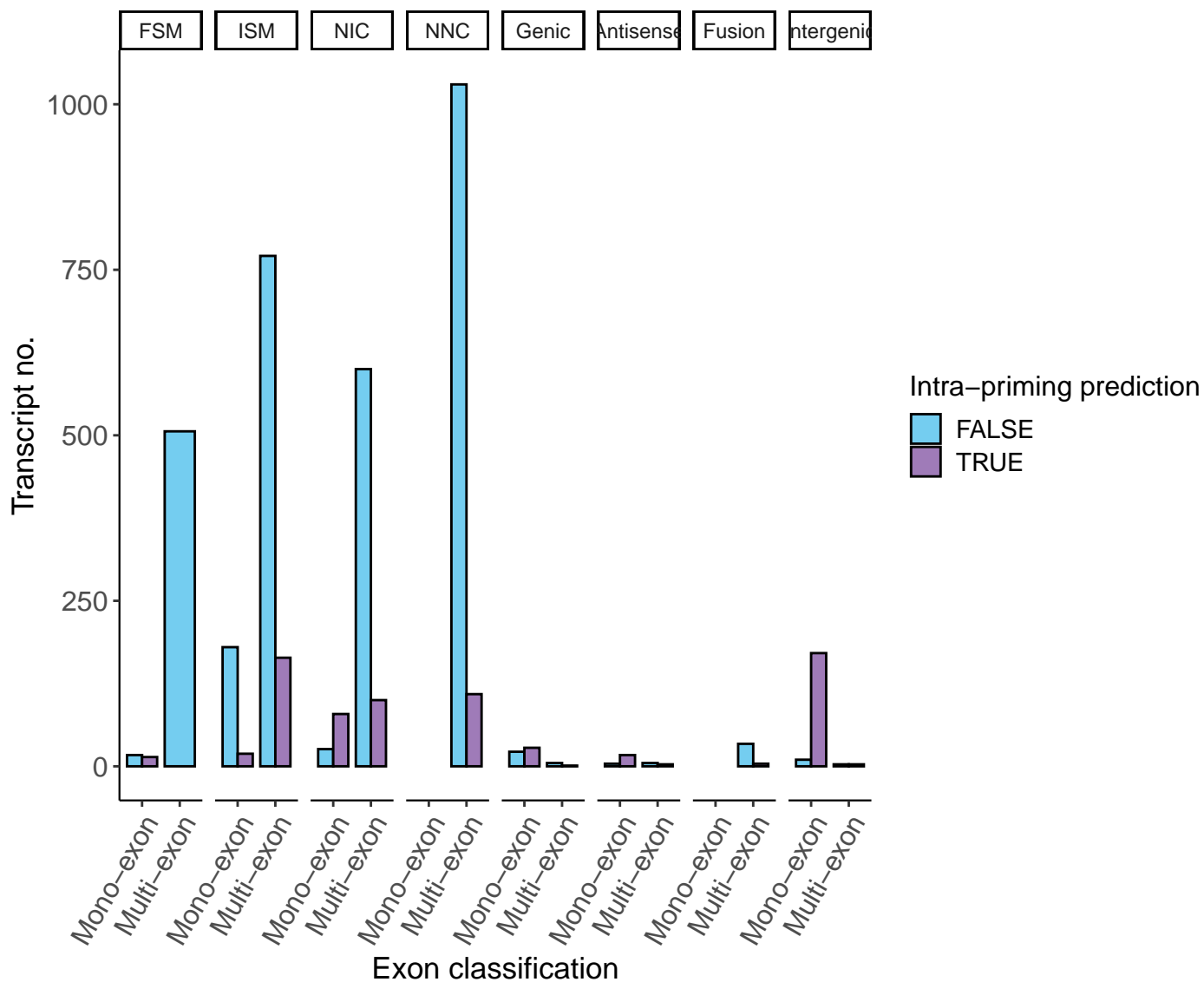


Isoforms flagged as intra-priming, by category (%)





Isoforms flagged as intra-priming, by exon number



Isoforms flagged as intra-priming, by exon number (%)

