# Supplementary Materials for 

## A Genome-wide Survey of Mutations in the Jurkat Cell Line

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Figs. S1 to S8

Captions for Tables S1 and S2

Figure S1: Raw CNVnator read depth density
Distribution of raw CNVnator read depth calls.


Figure S2: Deletion size distributions by variant caller (1 - 200bp)
Distributions of deletion call sizes between 1bp and 200bp for each variant caller. The bin size is 1 bp .


Figure S3: Deletion size distributions by variant caller (200 - 1,000bp)
Distributions of deletion call sizes between 200bp and 1 kb for each variant caller. The bin size is 10bp.


Figure S4: Deletion size distributions by variant caller (1,000 - 100,000,000bp)
Distributions of deletion call sizes between 1 kb and 100 Mb for each variant caller. The bin size is logscaled with 200 bins.


Figure S5: Insertion size distributions by variant caller
Distributions of insertion call sizes for each variant caller. The bin size is 1 bp .


Figure S6: Duplication size distributions by variant caller
Distributions of duplication call sizes for each variant caller. The bin size is log-scaled with 200 bins.


Figure S7: Inversion size distributions by variant caller
Distributions of inversion call sizes for each variant caller. The bin size is log-scaled with 100 bins.


Figure S8: Intra-chromosomal translocation size distributions by variant caller
Distributions of intra-chromosomal translocation call sizes for each variant caller. The bin size is logscaled with 100 bins.


Table S1: Long deletions found in Jurkat with matches to pathogenic variants in dbVar Matches were determined by $90 \%$ reciprocal overlap. All database matches are included with one dbVar entry per row.

Table S2: Duplications found in Jurkat with matches to pathogenic variants in dbVar Matches were determined by $90 \%$ reciprocal overlap. All database matches are included with one dbVar entry per row.

