Figure S14. Comparison between the mapped splice sites and polyadenylation sites in this study and the data set in [17]. - For the splice sites (A) and poly(A) sites (B), we defined a per-gene measure of overlap between the data in this study and the data in Siegel et al. 2010, as follows: define the entire set of sites found by both studies as s_i, for i=1,...,n. Define (in each study) the probability to observe the i'th site, p_i, as the number of reads for that site divided by the total number of reads for the gene. The overlap is the sum over all sites of min(p_i(this study), p_i(Siegel et al. 2010)). This gives 1, if there is perfect overlap and 0, if there is no overlap at all.