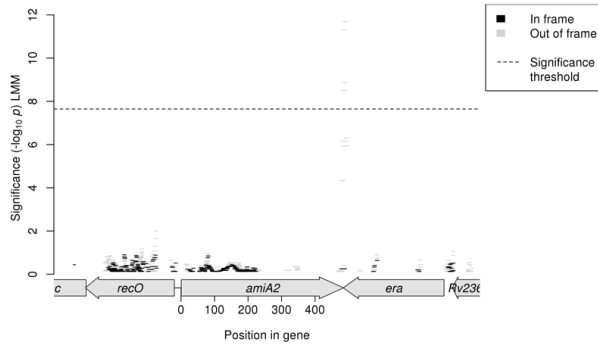
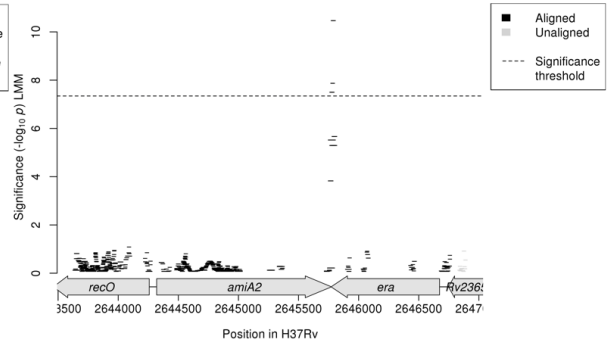


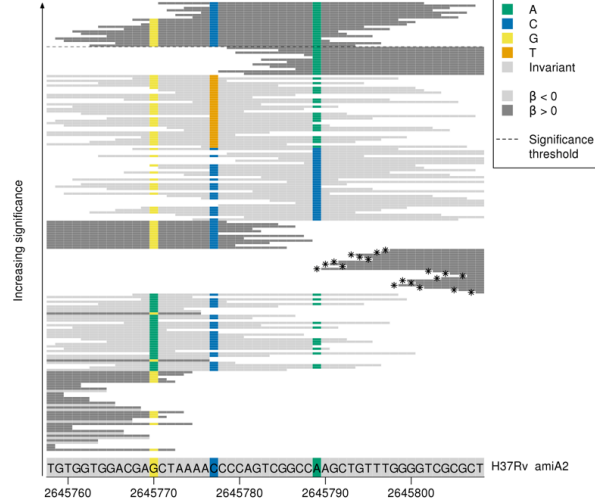
A Oligopeptides



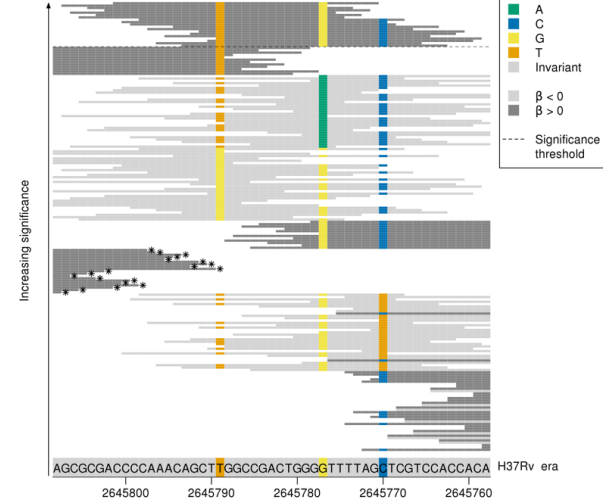
B Oligonucleotides



C *amiA2*



D *era*



S13 Fig. Variants in *amiA2* and *era* associated with bedaquiline MIC. Manhattan plots showing the association results for the *amiA2/era* coding region for the **A** oligopeptides and **B** oligonucleotides, and oligonucleotide alignment plots showing close ups of the significant region in *amiA2/era* in the correct reading frame for **C** *amiA2* **D** *era*. The black dashed lines indicate the Bonferroni-corrected significance thresholds. In the Manhattan plots, oligopeptides are coloured by the reading frame that they align to, black for the correct reading frame for *amiA2*. Oligo-peptides and nucleotides assigned to the region but did not align using BLAST are shown in grey on the right hand side of the plots. In the oligonucleotide alignment plots, the H37Rv reference alleles are shown at the bottom of the figure, grey for an invariant site, coloured at variant site positions. The oligonucleotides that aligned to the region are plotted from least significant at the bottom to most significant at the top. The background colour of the oligonucleotides represents the direction of the b estimate, light grey when $b < 0$ (associated with lower MIC), dark grey when $b > 0$ (associated with higher MIC). Oligonucleotides are coloured by their allele at all variant positions. Oligonucleotides below the MAF threshold and not included in the analysis, but visualised here for signal interpretation, are marked by *s.