Species (kingdom, phylum)	Genomic location (strand)	Frameshifting signal sequence	Length	Efficiency
Methanocaldococcus fervens AG86 (Archaea, Euryarchaeota)	NC_013156.1:1121859-1123902(+)	AAA AAG GAA AAT AAT GAC GTA AAA AAA AAA ACA AGA ATA AAT TAA ATA ATG AAT CAA ATA ATA ATG AGA ATA ATA AT	77 bp	63%
Methanocaldococcus sp. FS406-22 (Archaea, Euryarchaeota)	NC_013887.1:1224237-1224313(+)	GAG_CAA_AAA_AAT_GAT_GAC_GTA_AAA_AAA_AAC_ATG_ATG_AAA_TAA_GAA_ATG_AGG_ATT_CAA_AT	77 bp	40%
Delftia acidovorans SPH-1 (Bacteria; Betaproteobacteria)	NC_010002.1:6395711-6395640(-)	AAG_GTG_GGA_GCG_CCC_GCC_ACA_AAA_AAA_GCC_TGA_GCC_CCC_GCG_GCC_CCG_CGC_CCG_CGC_CCG_CGC_CCG_CGC_CCG_CGC_CCG_CGC_CCG_CGC_CCG_CGC_CAC_AGG_CAA_GGG_CTG_CGG_GGG	72 bp	10%