



| location | <i>M</i> (bits) | <i>z-score</i> | <i>robustness</i> | <i>position bias</i> | <i>orientation bias</i> | <i>conservation index</i> | <i>seed</i> | <i>motif name</i> | protein array |
|----------|-----------------|----------------|-------------------|----------------------|-------------------------|---------------------------|-------------|--------------------------------------|---------------|
| 5' | 0.008 | 44.2 | 10/10 | - | ↔ | 0.99 | ACGACGA | PIE_1 | - |
| 5' | 0.008 | 39.5 | 10/10 | - | ↔ | 1.00 | AATCGAT | CEH-38, CEH_28, CEH-28 ... | - |
| 5' | 0.004 | 17.9 | 10/10 | Y | → | 0.97 | CTCTCAC | PIE_1 | - |
| 5' | 0.005 | 26.8 | 10/10 | Y | ↔ | 1.00 | ACAGGTG | SNAI_1, HLH_1.4 | - |
| 5' | 0.005 | 25.4 | 10/10 | - | ↔ | 0.74 | CATATCA | HPO_39 | - |
| 5' | 0.004 | 22.6 | 10/10 | - | ↔ | 0.90 | TACCAA | TBX_40 | - |
| 5' | 0.004 | 22.7 | 10/10 | - | ↔ | 0.37 | TGCCTAC | NHR_256_NHR_79, NHR_28_NHR_53_NHR_70 | - |
| 5' | 0.004 | 21.5 | 10/10 | - | ↔ | 0.37 | CTATGTA | - | - |