PRELIMENARY STEP

- The computer used for executing the steps should conform with the system data provided in the file os-release.txt
- Extract the archive **HDVDB.tar**
- Place ACTAT_PERMS, newpos.sh, newpos.awk, dist.txt, sl1.sh, sl2.awk and the directory gx at the same location as the untarred HDVDB.tar
- The script sl1.sh uses RNAfold verify ViennaRNA is installed on computer and RNAfold is located at a directory included in the machine's \$PATH
- Place sorted_candidates_genotypes_3_7_und.xlsx at a location of your convenience on a Windows computer. This spreadsheet is provided for easy reference to the Results section.

At the same Linux location of the files from this step, run:

STEP 1

./newpos.sh

STEP 2

cat dist.txt | while read L; do ./sl1.sh \$L; done > candidates.txt &

STEP 3

• For genotype 3:

while IFS=\$'\t' read -r amber id _ sl1p _; do awk -v ID="\$id" -v AW="\$amber" -v SL1P="\$sl1p" -v SL2="CCGCAG" -f sl2.awk 1_xx\$id; done < ./gx/g3_44 | sort -nk10

• For genotype 7:

while IFS=\$'\t' read -r amber id_ sl1p _; do awk -v ID="\$id" -v AW="\$amber" -v SL1P="\$sl1p" -v SL2=" CCGAGG" -f sl2.awk 1_xx\$id; done < ./gx/g7_44 | sort -nk10

For undefined genotype:

```
while IFS=$'\t' read -r amber id_ sl1p_; do awk -v ID="$id" -v AW="$amber" -v SL1P="$sl1p" -v SL2="CCGCAG" -f sl2.awk 1_xx$id; done < ./gx/gund_44 | sort -nk10 while IFS=$'\t' read -r amber id_ sl1p_; do awk -v ID="$id" -v AW="$amber" -v SL1P="$sl1p" -v SL2=" CCGAGG" -f sl2.awk 1_xx$id; done < ./gx/gund_44 | sort -nk10
```