Inference on allele frequency (INCLASS-2)

Gustavo & Ana

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```
nAA=3
nAa=16
naa=31
# Sample size (number of diploid genotypes)
n=(nAA+nAa+naa)
# Total number of alleles is 2n
nAlleles=n*2
# Allele frequency
pHat=(2*nAA+nAa)/nAlleles
VAR=pHat*(1-pHat)/nAlleles
SE=sqrt(VAR)
NormalCI=pHat+c(-1,1)*1.96*SE
AlternativeCI=binom.test(x=c(nAA*2+nAa),n=n*2)$conf.int
message(pHat,' +/-',SE)
## 0.22 +/-0.041424630354416
message('Normal CI:', paste(round(NormalCI,4),collapse=' ; '))
## Normal CI:0.1388 ; 0.3012
message('Alternative CI:', paste(round(AlternativeCI,4),collapse=' ; '))
## Alternative CI:0.1433 ; 0.3139
```