Priors_production_AM.Rmd

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production des priors AM

mon_script library(stats) library("KScorrect", lib.loc="~/R/x86_64-pc-linux-gnu-library/3.3") #####partie locus #variables locus #-L=taille du gene #-t=theta#-r=rho#-delta=taille du track recombinant #boucle de 1000000 iterations(1000000 tirage demographique) demo<-NULL locus<-NULL tbs<-NULL #####TIRER un prior locus dans une distribution uniforme de bornes L<-scan("/home/kadurand/partage_windows/Xylella/analyses_genomiques/ABC/1368oRTHOLOGUES_summarystats/le t<--runif(1368,0, 0.001)#bound_theta=[0-0.0003]bornes vrai pour 13pauca_multiplex augmenter la borne su r<-runif(1368,0,0.001)#bound_theta=[0-0.0003]bornes vrai pour 13pauca_multiplex augmenter la borne sup delta<-round(runif(1368,10, 1000))#bound=[10-1000] #print(L, t, r, delta)m_locus=matrix(c(L,t,r,delta),ncol=4) m_locus=as.data.frame(m_locus) for (i in 1:10000) {#tirage des priors demographiques #variables demographique modéle SI ##Param_demo (5) = Ts N1, N2, M12, M21 Ts<-rlunif(1,100,1E+8)#bound=[1,100,1E+8] N1 < -rlunif(1,100,1E+6) #bound = [100,1E+6]N2 < -rlunif(1,100,1E+6) #bound = [100,1E+6]Na < -rlunif(1, 100, 1E+6) #Bound = [100, 1E+6]M12 < -runif(1,0.01,30) #bound = [0.01-30]M21 < -runif(1, 0.01, 30) #bound = [0.01-30]Tam < -rlunif(1,100,Ts) #) #bound = [0-100]#print(Ts , N1, N2, M12, M21, Tam) m_demo=matrix(c(Ts,N1,N2,Na,M12,M21,Tam),ncol=7) m_demo=as.data.frame(m_demo) locus<-cbind(m_locus,m_demo)</pre> path <- "/home//kadurand/partage_windows/Xylella/analyses_genomiques/ABC/fastSimBac_linux/Priors_AM_100</pre> write.table(locus,file= paste(path,i, sep="-"),col.names=FALSE,row.names =FALSE)