## Supporting Information Appendix S1

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Journal Title

## 4 Section S1.1 R Code for Consumer-Resource Model

- Below is the R code for our model function, which is represented mathematically in the main text in
- 6 Equations 1-4. The same code, along with all the code to reproduce our results, has been archived on
- Figshare (link) and is available on GitHub (http://github.com/atredennick/Coexistence-Stability/
- 8 releases).

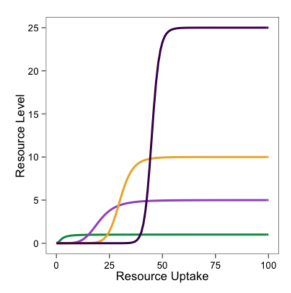
```
simulate_model <- function(seasons, days_to_track, Rmu,</pre>
                           Rsd_annual, sigE, rho,
                           alpha1, alpha2, alpha3, alpha4,
                           beta1, beta2, beta3, beta4,
                           eta1, eta2, eta3, eta4,
                           theta1, theta2, theta3, theta4,
                           nu, r1, r2, r3, r4,
                           a1, a2, a3, a4,
                           b1, b2, b3, b4,
                           eps1, eps2, eps3, eps4,
                           D1, D2, D3, D4,
                           N1, N2, N3, N4, R) {
 require('deSolve') # for solving continuous differential equations
 require('mvtnorm') # for multivariate normal distribution functions
 ## Assign parameter values to appropriate lists
 DNR \leftarrow c(D=c(D1,D2,D3,D4),
                              # initial dormant state abundance
          N=c(N1,N2,N3,N4), # initial live state abundance
                               # initial resource level
           R=R)
 parms <- list (</pre>
   r = c(r1, r2, r3, r4),
                                  # max growth rate for each species
       = c(a1,a2,a3,a4), # rate parameter for Hill function
   b = c(b1, b2, b3, b4),
                                 # shape parameter for Hill function
    eps = c(eps1,eps2,eps3,eps4) # resource-to-biomass efficiency
```

```
####
####
      Sub-Model functions -----
####
## Continuous model (Equations 1-2)
updateNR <- function(t, NR, parms){</pre>
  with(as.list(c(NR, parms)), {
    dN1dt = N1*eps[1]*(uptake_R(r[1], R, a[1], b[1]))
    dN2dt = N2*eps[2]*(uptake_R(r[2], R, a[2], b[2]))
    dN3dt = N3*eps[3]*(uptake_R(r[3], R, a[3], b[3]))
    dN4dt = N4*eps[4]*(uptake_R(r[4], R, a[4], b[4]))
    dRdt = -1 * (dN1dt/eps[1] + dN2dt/eps[2] + dN3dt/eps[3] + dN4dt/eps[4])
    list(c(dN1dt, dN2dt, dN3dt, dN4dt, dRdt)) # output as list
  })
} # end continuous function
## Discrete model (Equations 3-4)
update_DNR <- function(t, DNR, gammas,</pre>
                        alpha1, alpha2, alpha3, alpha4,
                        eta1, eta2, eta3, eta4,
                        beta1, beta2, beta3, beta4,
                        theta1, theta2, theta3, theta4, nu) {
  with (as.list(DNR),{
          <- gammas[1]
    g1
    g2
          <- gammas[2]
    g3
          <- gammas[3]
    g4
          <- gammas [4]
    D1new <- alpha1*N1 + D1*(1-g1)*(1-eta1)
    D2new \leftarrow alpha2*N2 + D2*(1-g2)*(1-eta2)
    D3new \leftarrow alpha3*N3 + D3*(1-g3)*(1-eta3)
    D4new <- alpha4*N4 + D4*(1-g4)*(1-eta4)
    N1new <- beta1*(1-alpha1)*N1 + g1*(D1+(alpha1*N1))*(1-eta1)
    N2new \leftarrow beta2*(1-alpha2)*N2 + g2*(D2+(alpha2*N2))*(1-eta2)
    N3new \leftarrow beta3*(1-alpha3)*N3 + g3*(D3+(alpha3*N3))*(1-eta3)
    N4new \leftarrow beta4*(1-alpha4)*N4 + g4*(D4+(alpha4*N4))*(1-eta4)
    Rnew \leftarrow theta1*(1-alpha1)*N1 + theta2*(1-alpha2)*N2 +
             theta3*(1-alpha3)*N3 + theta<math>4*(1-alpha4)*N4 +
             nu*R + Rvector[t]
```

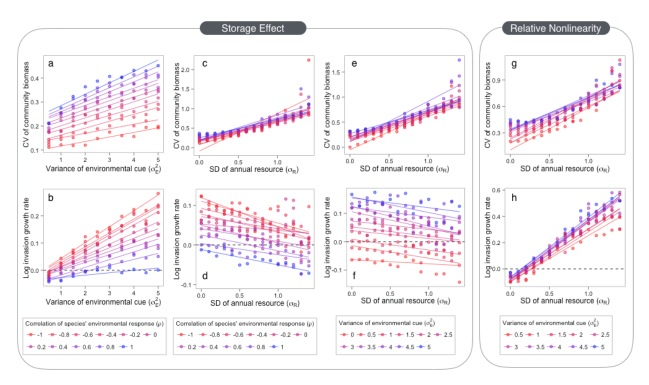
```
return(c(D1new, D2new, D3new, D4new, N1new, N2new, N3new, N4new, Rnew))
  })
}
## Resource uptake function (Hill function)
uptake_R <- function(r, R, a, b) {
  return((r*R^a) / (b^a + R^a))
}
## Generate germination fractions
getG <- function(sigE, rho, nTime, num_spp) {</pre>
               <- matrix(rep(rho*sigE,num_spp*2), num_spp, num_spp)
  diag(varcov) <- sigE</pre>
  if(sigE > 0) { varcov <- Matrix::nearPD(varcov)$mat } # coerce matrix to PD</pre>
  varcov <- as.matrix(varcov)</pre>
        <- rmvnorm(n = nTime, mean = rep(0,num_spp), sigma = varcov)</pre>
         <- exp(e) / (1+exp(e))
  return(g)
}
####
#### Simulate model -----
####
              <- c(1:days_to_track)</pre>
days
num_spp
              <- length(parms$r)</pre>
              <- names(DNR)
nmsDNR
               <- grep("D", names(DNR))
dormants
NR.
               <- DNR[-dormants]
               <- names(NR)
nmsNR
gVec
               <- getG(sigE = sigE, rho = rho, nTime = seasons, num_spp = num_spp)</pre>
Rvector
              <- rlnorm(seasons, Rmu, Rsd_annual)</pre>
saved_outs
               <- matrix(ncol=length(DNR), nrow=seasons+1)
saved_outs[1,] <- DNR</pre>
## Loop over seasons
for(season_now in 1:seasons) {
  # Simulate continuous growing season
  output <- ode(y = NR, times=days, func = updateNR, parms = parms)</pre>
```

```
<- output[nrow(output),nmsNR]</pre>
    dormants <- grep("D", names(DNR))</pre>
    DNR
              <- c(DNR[dormants], NR)
    # Save end of season biomasses, before discrete transitions
    saved_outs[season_now+1,] <- DNR</pre>
    names(DNR) <- nmsDNR</pre>
    DNR <- update_DNR(season_now, DNR, gVec[season_now,],</pre>
                        alpha1 = alpha1, alpha2 = alpha2,
                        alpha3 = alpha3, alpha4 = alpha4,
                        eta1 = eta1, eta2 = eta2, eta3 = eta3, eta4 = eta4,
                       beta1 = beta1, beta2 = beta2,
                       beta3 = beta3, beta4 = beta4,
                       theta1 = theta1, theta2 = theta2,
                       theta3 = theta3, theta4 = theta4,
                       nu=nu)
    names(DNR) <- nmsDNR</pre>
               <- DNR[-dormants]</pre>
    names(NR) <- nmsNR</pre>
  } # next season
  return(saved_outs)
} ## End simulation function
```

## Section S1.2 Additional Figures



**Figure S1-1** Resource uptake curves for each species (represented by different colors) as used in relative nonlinearity simulations.



**Figure S1-2** Variability of community biomass and invasion growth rates of the inferior competitor under different parameter combinations. Points are mean values from 10,000 growing seasons and lines are linear fits to show trends. In **Storage Effect** plots, resource supply is held constant between growing seasons, whereas resource supply varies each year in **Relative Nonlinearity** simulations.

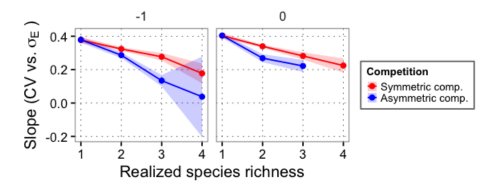


Figure S1-3 Slopes of linear fits for the relationship between  $\log(CV)$  and  $\log(\sigma_E)$  at different levels of realized species richness from storage effect simulations. The slopes come from linear models fit to log-transformed versions of Figure 3 in the main text. For these simulations, "symmetric competion" ( $\bullet$ ) refers to similar live-to-dormant biomass allocation fractions ( $\alpha = [0.5, 0.495, 0.49, 0.485]$  for the four species), and "asymmetric competition" ( $\bullet$ ) refers to more dissimilar live-to-dormant biomass allocation fractions ( $\alpha = [0.5, 0.49, 0.48, 0.47]$  for the four species).

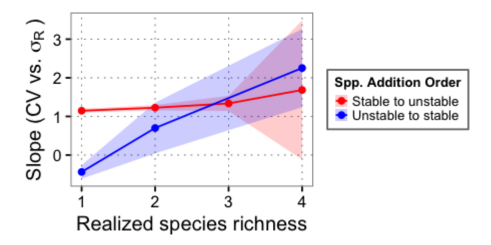


Figure S1-4 Slopes of linear fits for the relationship between  $\log(CV)$  and  $\log(\sigma_R)$  at different levels of realized species richness from relative nonlinearity simulations. The slopes come from linear models fit to log-transformed versions of Figure 4 in the main text.