Supporting Information

A.T. Tredennick, P.B. Adler, & F.R. Adler, "The relationship between species richness and..." $Ecology\ Letters$

⁴ Section SI.1 R Code for Consumer-Resource Model

- 5 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 The same code, along with all the code to reproduce our results, has been archived on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).

1

```
simulate_model <- function(seasons, days_to_track, Rmu,</pre>
                           Rsd_annual, sigE, rho,
                           alpha1, alpha2, alpha3, alpha4,
                           eta1, eta2, eta3, eta4,
                           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 <- c(D=c(D1,D2,D3,D4), # initial dormant state abundance
          N=c(N1,N2,N3,N4), # initial live state abundance
          R=R)
                               # initial resource level
 parms <- list (
       = c(r1,r2,r3,r4),
                                 # max growth rate for each species
       = c(a1,a2,a3,a4), # rate parameter for Hill function
      = 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
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
update_DNR <- function(t, DNR, gammas,</pre>
                        alpha1, alpha2, alpha3, alpha4,
                        eta1, eta2, eta3, eta4) {
  with (as.list(DNR),{
          <- gammas[1]
    g1
    g2
          <- gammas[2]
          <- gammas[3]
    g3
          <- gammas[4]
    g4
    D1new \leftarrow (1-g1)*(alpha1*N1 + D1)*(1-eta1)
    D2new \leftarrow (1-g2)*(alpha2*N2 + D2)*(1-eta2)
    D3new \leftarrow (1-g3)*(alpha3*N3 + D3)*(1-eta3)
    D4new <- (1-g4)*(alpha4*N4 + D4)*(1-eta4)
    N1new \leftarrow g1*(alpha1*N1 + D1)*(1-eta1)
    N2new \leftarrow g2*(alpha2*N2 + D2)*(1-eta2)
    N3new \leftarrow g3*(alpha3*N3 + D3)*(1-eta3)
    N4new \leftarrow g4*(alpha4*N4 + D4)*(1-eta4)
    Rnew <- Rvector[t]</pre>
    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)</pre>
  diag(varcov) <- sigE</pre>
  if(sigE > 0) { varcov <- Matrix::nearPD(varcov)$mat } # crank through nearPD to fix roundi
 varcov <- as.matrix(varcov)</pre>
         <- rmvnorm(n = nTime, mean = rep(0,num_spp), sigma = varcov)
         <- exp(e) / (1+exp(e))
  return(g)
}
####
####
              <- c(1:days_to_track)
days
              <- length(parms$r)
num_spp
              <- names(DNR)
nmsDNR
dormants
              <- grep("D", names(DNR))
NR
              <- DNR[-dormants]
nmsNR
              <- names(NR)
              <- getG(sigE = sigE, rho = rho, nTime = seasons, num_spp = num_spp)</pre>
gVec
              <- rlnorm(seasons, Rmu, Rsd_annual)
Rvector
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
          <- ode(y = NR, times=days, func = updateNR, parms = parms)</pre>
           <- output[nrow(output),nmsNR]</pre>
  dormants <- grep("D", names(DNR))</pre>
           <- c(DNR[dormants], NR)
  DNR
  # Save end of season biomasses, before discrete transitions
  saved_outs[season_now+1,] <- DNR</pre>
```

Section SI.2 Exploring Parameter Space

Section SI.3 Eight-Species Storage Effect Model

Section SI.4 Additional Figures

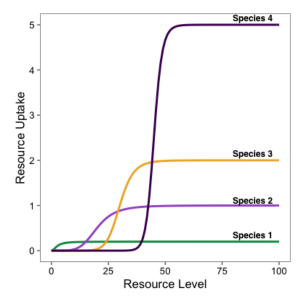


Figure S1-1 Resource uptake curves for each species (represented by different colors) as used in relative nonlinearity simulations. The equation for resource uptake is: $f_i(R) = r_i R^{a_i}/(b_i^{a_i} + R^{a_i})$. Parameter values are as follows. Species 1: r = 0.2, a = 2, b = 2.5; Species 2: r = 1, a = 5, b = 20; Species 3: r = 2, a = 10, b = 30; Species 4: r = 5, a = 25, b = 45.

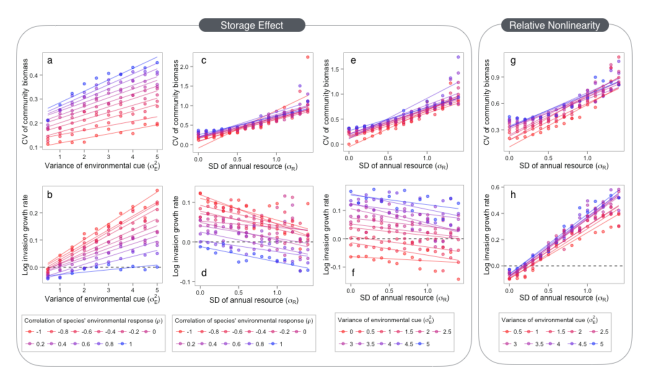


Figure S1-2 Variability of community biomass and invasion growth rates of the inferior competitor in a two-species community under different parameter combinations. Points are mean values from 10,000 growing seasons and lines are linear fits to show trends. In **Storage Effect** plots, resource supply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

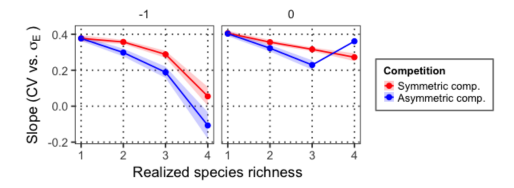


Figure S1-3 Slopes of linear fits for the relationship between $\log(CV)$ and $\log(\sigma_E)$ at different levels of realized species richness from storage effect simulations. The slopes come from linear models fit to log-transformed versions of Figure 3 in the main text. For these simulations, "symmetric 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).

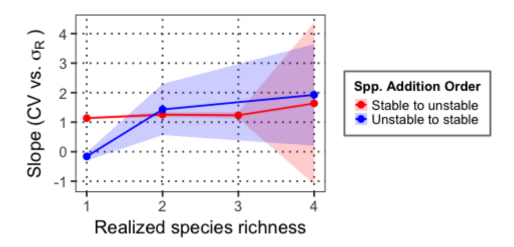


Figure S1-4 Slopes of linear fits for the relationship between $\log(CV)$ and $\log(\sigma_R)$ at different levels of realized species richness from relative nonlinearity simulations. The slopes come from linear models fit to log-transformed versions of Figure 4 in the main text.