



Supplement of

Underestimation of boreal soil carbon stocks by mathematical soil carbon models linked to soil nutrient status

Boris Šupek et al.

Correspondence to: Boris Šupek

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1 Source codes of the Yasso07, Q, and CENTURY model

1.1 Yasso07 model

https://code.google.com/p/yasso07ui/source/browse/trunk/y07_subroutine.f90

```
MODULE yasso
IMPLICIT NONE
CONTAINS
SUBROUTINE mod5c(a,t,cl,init,inf,s,leac,z)
!components separately
IMPLICIT NONE
!***** &
!GENERAL DESCRIPTION FOR ALL THE MEASUREMENTS
!***** &
!returns the model prediction for given parameters
! 1-16 matrix A entries: 4*k, 12*p
!17-19 Climate-dependence parameters: b1, b2, g1
!20-22 Leaching parameters: f1, f2, f3 IGNORED IN THE Y07-UI
!23-25 Woody parameters
!26-27 Humus parametens: kH, pH
REAL,DIMENSION(27),INTENT(IN) :: a !parameters
REAL,INTENT(IN) :: t,s,leac !time,size,leaching
REAL,DIMENSION(3),INTENT(IN) :: cl !climatic conditions
REAL,DIMENSION(5),INTENT(IN) :: init
REAL,DIMENSION(5),INTENT(IN) :: inf !infall
REAL,DIMENSION(5),INTENT(OUT) :: z
REAL,DIMENSION(5,5) :: m,mt,m2,mi
INTEGER :: i
REAL,PARAMETER :: pi=3.1415926535
REAL :: tem
REAL,DIMENSION(5) :: te
REAL,DIMENSION(5) :: z1,z2
```

```

!temperature annual cycle approximation
te(1)=cl(1)+4*c1(3)*(1/SQRT(2.0)-1)/pi
te(2)=cl(1)-4*c1(3)/SQRT(2.0)/pi
te(3)=cl(1)+4*c1(3)*(1-1/SQRT(2.0))/pi
te(4)=cl(1)+4*c1(3)/SQRT(2.0)/pi
tem=0.0
DO i=1,4 !Annual cycle, different models
  tem=tem+EXP(a(17)*te(i)+a(18)*te(i)**2.0)/4.0 !Gaussian
END DO
!Precipitation dependence
tem=tem*(1.0-EXP(a(19)*cl(2)/1000))
!Size class dependence -- no effect if sc = 0.0
m(1,1)=a(1)*tem*MIN(1.0,(1.0+a(23)*s+a(24)*s**2.0)**a(25))
m(2,2)=a(2)*tem*MIN(1.0,(1.0+a(23)*s+a(24)*s**2.0)**a(25))
m(3,3)=a(3)*tem*MIN(1.0,(1.0+a(23)*s+a(24)*s**2.0)**a(25))
m(4,4)=a(4)*tem*MIN(1.0,(1.0+a(23)*s+a(24)*s**2.0)**a(25))
!Calculating matrix M, normal
m(2,1)=a(5)*ABS(m(2,2))
m(3,1)=a(6)*ABS(m(3,3))
m(4,1)=a(7)*ABS(m(4,4))
m(5,1)=0.0
m(1,2)=a(8)*ABS(m(1,1))
m(3,2)=a(9)*ABS(m(3,3))
m(4,2)=a(10)*ABS(m(4,4))
m(5,2)=0.0
m(1,3)=a(11)*ABS(m(1,1))
m(2,3)=a(12)*ABS(m(2,2))
m(4,3)=a(13)*ABS(m(4,4))
m(5,3)=0.0
m(1,4)=a(14)*ABS(m(1,1))
m(2,4)=a(15)*ABS(m(2,2))
m(3,4)=a(16)*ABS(m(3,3))
m(5,4)=0.0
m(5,5)=a(26)*tem !no size effect in humus
DO i=1,4
  m(i,5)=a(27)*ABS(m(i,i)) !mass flows EWAN -> H
END DO

```

```

!Leaching
m(1,1)=m(1,1)+leac*c1(2)/1000
m(2,2)=m(2,2)+leac*c1(2)/1000
m(3,3)=m(3,3)+leac*c1(2)/1000
m(4,4)=m(4,4)+leac*c1(2)/1000
!DY solution starts here...
DO i=1,5
  z1(i)=DOT_PRODUCT(m(:,i),init)+inf(i)
END DO
mt=m*t
CALL matrixexp(mt,m2)
DO i=1,5
  z2(i)=DOT_PRODUCT(m2(:,i),z1)-inf(i)
END DO
CALL inverse(m,mi)
DO i=1,5
  z1(i)=DOT_PRODUCT(mi(:,i),z2)
END DO
z=z1
CONTAINS
SUBROUTINE matrixexp(a,b)
IMPLICIT NONE
!returns approximated matrix exponential
!Taylor approximation.. another algorithm perhaps?
REAL,DIMENSION(5,5),INTENT(IN) :: a
REAL,DIMENSION(5,5),INTENT(OUT) :: b
REAL,DIMENSION(5,5) :: c,d
REAL :: p,normiter
INTEGER :: i,q,j
q=10
b=0.0
DO i=1,5
  b(i,i)=1.0
END DO
normiter=2.0
j=1
CALL matrix2norm(a, p)

```

```

DO
  IF (p< normiter) THEN
    EXIT
  END IF
  normiter=normiter*2.0
  j=j+1
END DO
c=a/normiter
b=b+c
d=c
DO i=2,q
  d=MATMUL(c,d)/REAL(i)
  b=b+d
END DO
DO i=1,j
  b=MATMUL(b,b)
END DO
END SUBROUTINE matrixexp
SUBROUTINE matrix2norm(a,b)
IMPLICIT NONE
! returns matrix 2-norm with cartesian vector x,
!| | x| | = 1
!square matrix input (generalize if necessary)
REAL,DIMENSION(5,5),INTENT(IN) :: a
REAL,INTENT(OUT) :: b
INTEGER :: n,i
n=SIZE(a(1,:))
b=0.0
DO i=1,n
  b=b+SUM(a(:,i))**2.0
END DO
b=SQRT(b)
END SUBROUTINE matrix2norm
SUBROUTINE inverse(a,b)
IMPLICIT NONE
! returns an inverse of matrix a
!(column elimination strategy)

```

```

!input has to be a square matrix, otherwise erroneous
REAL,DIMENSION(5,5),INTENT(IN) :: a
REAL,DIMENSION(5,5),INTENT(OUT) :: b
REAL,DIMENSION(5,5) :: c
INTEGER :: n,m,i,j
n=SIZE(a(1,:))
m=SIZE(a(:,1))
IF(m/=n) THEN
    WRITE(*,*) " Does not compute."
    WRITE(*,*) " No square matrix input."
    WRITE(*,*) " Error in function: inverse"
ELSE
    ! ALLOCATE(b(n,n),c(n,n))
    c=a
    b=0.0
    DO i=1,n !setting b a unit matrix
        b(i,i)=1.0
    END DO
    DO i=1,n
        !what if diagonal values are zeros?
        IF(c(i,i)==0.0)THEN!case of singuar matrix, is it?
            b(i,:)=0.0
            c(i,:)=0.0
            b(:,i)=0.0
            c(:,i)=0.0
            !           b(i,i)=1.0
            !           c(i,i)=1.0
        ELSE
            b(i,:)=b(i,:)/c(i,i)
            c(i,:)=c(i,:)/c(i,i)
        END IF
        DO j=1,i-1
            b(j,:)=b(j,:)-b(i,:)*c(j,i)
            c(j,:)=c(j,:)-c(i,:)*c(j,i)
        END DO
        DO j=i+1,n
            b(j,:)=b(j,:)-b(i,:)*c(j,i)
        END DO
    END DO
END

```

```

c(j,:)=c(j,:)-c(i,:)*c(j,i)
END DO
END DO
IF(c(n,n)==0.0) THEN
  b(n,:)=0.0
  b(:,n)=0.0
  !           b(n,n)=1.0
ELSE
  b(n,:)=b(n,:)/c(n,n)
END IF
!now, b is supposed to be the requested inverse
END IF
END SUBROUTINE inverse
END SUBROUTINE mod5c
END MODULE yasso

```

1.2 Q model

```

!***** &
!      Main program to calculate carbon store in forest soils.
!
!      SPRUCE
!***** &
!      Edited by Carina Ortiz, Version 2015-11-16
!
!      Understory vegetation, variable temp, variable litter
      real, dimension(1) :: yr(0:0),carb(0:0),
      +                  totc(0:0),u0y(0:0)
      real, dimension(1,5) :: lit(0:0,5),cssn(0:0,1),
      + vtempsite(0:0,1:244),cssnund(0:0,1),ssi0b(0:0,1),
      + ssi0scr(0:0,1),ssi0s(0:0,1),alfanss(0:0,1),
      z(0:0,1),u0(0:0,1)
      *,ssi0b(0:0,5),ssi0scr(0:0,5),ssi0s(0:0,5)
      * + ssi0scr(0:0,5),cssnund(0:0,5)
      real, dimension (244,0:0) :: nefr !(jj,ii)
      real, dimension (244,0:0) :: br !(jj,ii)
      real, dimension (244,0:0) :: st !(jj,ii)
      real, dimension (244,0:0) :: stpcr !(jj,ii)
      real, dimension (244,0:0) :: und !(jj,ii)

```

```

real, dimension (244,0:0) :: vtemp ! (jj,ii)
! needlesfineroots jj columns(MCSIMS), rows(yr) ii
    real i0, u0sum, ts
    character (len=10) :: dumtext
integer :: r
!
! Monte Carlo Simulations
!
! Read file with parameter setups
!
! Open parameter file
OPEN (unit=111, file='1parvaraccregsprucemean.dat',status='old')
    read(111,*) dumtext
!
! Open output file
open(unit=11, file='q_soil.dat')
!
! Read matrix file with litter production in N simulations
!
! (Monte Carlo), each fraction one separate file
open(unit=1111,file='biom70reg1SpruceNeedlesandfineroots.dat',
+
      status='old')
read(1111,*) dumtext
read(1111,*) nefr
open(unit=11111,file='biom70reg1SpruceBranches.dat',
+
      status='old')
read(11111,*) dumtext
read(11111,*) br
open(unit=111111,file='biom70reg1SpruceStumpCoarseroots.dat',
+
      status='old')
read(111111,*) dumtext
read(111111,*) stpcr
open(unit=111111,file='biom70reg1SpruceStem.dat',
+
      status='old')
read(111111,*) dumtext
read(111111,*) st
open(unit=1111111,file='biom70reg1SpruceUnderveg.dat ',
+
      status='old')
read(1111111,*) dumtext
read(1111111,*) und
!
! Read variable temperature file yearly means
open(unit=1, file='reftempreglsspruce.dat',status='old')

```

```

read(1,*) dumtext
read(1,*) vtemp
! Estimates the mean temperature for equilibrium
*      ts=ts+vtemp
*      ii=ii+1
*      temp=ts/ii !mean temp
! Main loop Number of Monte Carlo simulations
DO ii=1,244
read(111,*) q0n,q0w,e0,etall,beta,maxb,maxs,u00,u01
! Temperature for equilibrium.
! First row temperature, only for one step
do jj=0,0!first row is dumtext !
vtempsite(jj,ii) = vtemp(ii,jj) !
u0(jj,1)=u00+u01*vtempsite(jj,ii)
do i=0,0
u0y(i)=u00+u01*vtempsite(jj,ii) ! each year new temp.
enddo
enddo
! Set some parameters for decomposition functions
ssi0n=0
fC=0.5
do jj=0,0
alfanss(jj,1)=fC*beta*etall*u0(jj,1)*q0n**beta
z(jj,1)=(1.-e0)/(beta*etall*e0)
end do
! Create litter matrix
do jj=0,0!first row is dumtext
lit(jj,1) = nefr(ii,jj)
!litter fraction 1 needles & fine roots
lit(jj,2) = br(ii,jj)
!litter fraction 2 branches & roots
lit(jj,3) = st(ii,jj)
!litter fraction 3 stems
lit(jj,4) = stpcr(ii,jj)
!litter fraction 4 stump & coarse roots
lit(jj,5) = und(ii,jj)
!litter fraction 5 understorey

```

```

    enddo
!Calculates equilibrium litter fraction for needles
do jj=0,0
cssn(jj,1)=(lit(jj,1))*1/(alfanss(jj,1)*(z(jj,1)-1))
enddo
!Steady state understorey vegetation
do jj=0,0
cssnund(jj,1)=(lit(jj,5))*1/(alfanss(jj,1)*(z(jj,1)-1))
enddo
!Steady state input branches
do jj=0,0
ssi0b(jj,1)=lit(jj,2)
enddo
!Steady state input stump and coarse roots
do jj=0,0
ssi0scr(jj,1)=lit(jj,4)
enddo
!Steady state input stem
do jj=0,0
ssi0s(jj,1)=lit(jj,3)
enddo
! Fixed parameters
fC=0.5
! Call for soil decomposition model
call soildecomp(fC,u0,u0y,eta11,e0,beta,ssi0b,ssi0s,ssi0scr,
+ q0n,cssn,q0w,maxb,maxs,itend,lit,carb,nitr,
+ cssnund)

! WRITE TO OUTPUTFILE
* do i=0,0
* write(11,*) carb(i)
* enddo
! End main loop
* enddo !input litter loop
ENDDO !parameter loop
*! Close Temperature file
close(unit=1)
! Close litterfiles

```

```

close(unit=1111)
    close(unit=11111)
close(unit=111111)
close(unit=1111111)
close(unit=11111111)
! Close OUTPUT document
close(unit=11)
! Close parameterrange file
close(unit=111)

!***** &
! Subroutine to calculate carbon stores in forest soils
! Version 2015-11-16
! Undervegetation, variable temp, variable litter
subroutine soildecomp (fC,u0,u0y,etall,e0,beta,ssi0b,ssi0s,
+ ssi0scr,q0n,cssn,q0w,maxb,maxs,itend,lit,carb,nitr,cssnund)
real, dimension(1) :: carb(0:0), totc(0:0),
+ u0y(0:0)
real, dimension(1,5) :: lit(0:0,5), vtempsite(0:0,1:225)
real, dimension (225,0:0) :: nefr !(jj,ii)
real, dimension (225,0:0) :: br !(jj,ii)
real, dimension (225,0:0) :: st !(jj,ii)
real, dimension (225,0:0) :: stpcr !(jj,ii)
real, dimension (225,0:0) :: und !(jj,ii)
real, dimension (225,0:0) :: vtemp !(jj,ii)
* needlesfineroots jj columns(MCSIMS), rows(yr) ii
real i0, u0sum, t, gnv, gnold, gbv, gbold, gsv, gsold
dimension gn(0:0), gb(0:0), gs(0:0)
dimension hn(0:0), hb(0:0), hs(0:0), qn(0:0)
* real, dimension (101,100):: a(0:100,1:100)
integer :: r
! Litter and OM decomposition parameters
z=(1.-e0)/(beta*etall*e0)
zn=1./(beta*etall*e0)
alfa =fC*beta*etall*u0*q0w**beta
!Alfa with constant temp!
alfav=fC*beta*etall*q0w**beta
!Alfa with variable temperature vtemp!

```

```

alfan=fC*beta*eta11*u0*q0n**beta
!Old alfa with constant temp!
alfanv=fC*beta*eta11*q0n**beta
!Alfa with variable temperature vtemp!
alfa0n=fN/fC-(beta*eta11*e0+e0-1)*(fn/fc-r0)/(beta*eta11*e0-1)
! Old SOM from equilibrium org.mat. is calculated and summed
do i=0,0
u0sum=0
do k=1,i !No decomposition the same year as we assess
u0sum=u0sum+u0y(k)
!Integral of u0 when using variable temperature
enddo
gnold= (1+alfanv*u0sum)**(1-(1-e0) /
!Decomposition of old needles
+ (e0*eta11*beta))
t=(i)
! Call for decomposition fuctions
! that calculates the remaining mass of the o.m.
gsold=gold(t,maxs,alfa,alfav,u0sum,z)
gbold=gold(t,maxb,alfa,alfav,u0sum,z)
! Old SOM is summed
carb(i)=
+ gnold*cssn !part of o.m left and the equilibrium input in
fraction
+ +gnold*cssnund
+ +gbold*ssi0b
+ +gsold*ssi0s
+ +gsold*ssi0scr
write(11,*) carb (i)
enddo
! New SOM from is calculated
! and summed with the olds equilibrium carbon
do i=0,0
do mt=0,i
u0sum=0
if (i.ne.mt) then
!No decomposition the same year as we assess

```

```

do k=(mt+1),i
    u0sum=u0sum+u0y(k)
    !Integral of u0 when using variable temperature
enddo
endif
gnv=1./(1.+alfanv*u0sum)**z
!Decomposition of new needles!
t=(i-mt)
gbv=gbran(t,maxb,alfa,alfav,u0sum,z)
!Decomposition of new branches!
if (gbv.gt.1) then
gbv=1
endif
gsv=gbran(t,maxs,alfa,alfav,u0sum,z)
!Decomposition of new stems!
if (gbs.gt.1) then
gbs=1
endif
!      SOM from old equilibrium and new litter is summed
carb(i)=carb(i) +
        +      lit(mt,1)*gnv+
        +      lit(mt,2)*gbv+
        +      (lit(mt,3)+lit(mt,4))*gsv+
        +      lit(mt,5)*gnv
enddo
enddo
!      Litter fractions are 1=needles 2=branches
!      3=stem 4=stump & course roots 5=under vegetation
9991 continue
return
999 end
!      Function for decomposition of old equilibrium carbon
!      for branches (2) and stems(3+4)
function gold(t,itmax,alfa,alfav,u0sum,z)
!Decomposition of old branches & stem!
tmax=real(itmax)
if (t.le.tmax) then

```

```

gold=-2.* (1.+alfav*u0sum)**(2.-z) / (tmax*alfa**2*(1.-z)*(2.-z))
++2.* ((1.+alfav*u0sum)**(3.-z)-1.)
+   / (tmax**2*alfa**3*(1.-z)*(2.-z)*(3.-z))
+   +(1.-t/tmax)**3*tmax/3.-(1.-t/tmax)**2/(alfa*(1.-z))
+   +2.* (tmax-t)/(tmax**2*alfa**2*(1.-z)*(2.-z))

return
else
gold=-2.* (1+alfav*u0sum)**(2.-z) / (tmax*alfa**2*(1.-z)*(2.-z))
+   2.* ((1.+(alfav*u0sum-alfa*(tmax)))** (3.-z)-
+   (1+alfav*u0sum)**(3.-z))
+   / (tmax**2*alfa**3*(1.-z)*(2.-z)*(3.-z))

return
endif
end

function gbran(t,itmax,alfa,alfav,u0sum,z)
!Decomposition of new branches!
tmax=real(itmax)
if (t.le.tmax) then
gbran=2.* ((1+alfav*u0sum)**(1.-z)-(1.-t/tmax)) / (tmax*alfa*(1.-z))
+   +2.* (1.-(1.+alfav*u0sum)**(2.-z)) / ((tmax**2)*(alfa**2)*
+   (1.-z)*(2.-z))+(1.-(t/tmax))**2
return
else
gbran=2.* (1.+alfav*u0sum)**(1.-z) / (tmax*alfa*(1.-z))
+   +2.* ((1.+(alfav*u0sum-alfa*tmax))** (2.-z)-
+   (1.+alfav*u0sum)**(2.-z)) /
+   ((tmax**2)*(alfa**2)*(1.-z)*(2.-z))
return
endif
end

```

1.3 CENTURY model

```

#####
## 
##      SOC sub-model of the CENTURY version 4.0
##
```

```

#####
#
# Coded in R by Shoji Hashimoto (shojih@ffpri.affrc.go.jp)
# edited by Boris Tupek (boris.tupek@luke.fi)
# original model available at
# https://www.nrel.colostate.edu/projects/century/obtain2.htm
#####
#
# Related source files in the original CENTURY model
# Please see /original/source/*.f
#
# adjlig.f, anerob.f, csa_detiv.f, csa_main.f, cycle.f, declig.f
# decomp.f, eachyr.f, h2olos.f, litdec.f, partit.f
# pevap.f, prelim.f, simsom.f, somdec.f, tcalc.f
# wdeath.f, woodec.f, and so on.
#
#####
#
# Simplification:
#
# only for forest ecosystem (not grass, savanna etc)
# no irrigation
# not floating C/N ratio for plant organs.
# cnr_max=cnr_min=cnr_initial in tree.100
# no mineral N cycling: constant N at surface soil
# (xNmineral in f_site.100)
# drain=1, anerb=1
# idef=2 in fix.100 (water function for calculating defac)
# no CO2 effect
#
#####
#
# A bug in the original CENTURY
#
# a bug (please see calfc_wtpt function below)
# The difference in results was small,
# but it depends on the climate and soil.
#
# BFix<-0: with bug as the original CENTURY
# BFix<-1: the bug was fixed

```

```

rm(list=ls())
options(digits=12)
BFix<-1
#define number of years for spinup simulations!
TSTART=1
TEND=500 # 5000 for equilibrium
#####
# Read data
#####
# parameters from fix.100 in the original CENTURY
# environments (site specific temperature,
# precipitation from SMHI), site.100 in the original CENTURY
# parameters describing site conditions(site specific sand,
# silt,clay,bulk density from SFSI data)
# see file site.100 in the original CENTURY
# parameters describing tree,
# see tree.100 in the original CENTURY
# "AND H_J ANDREWS" for conifers
# "Coweeta" for deciduous
# initial conditions from site.100
## READ SITE SPECIFIC data #####
#general parameters (fix.100)
parameters.names<-c("adep1","adep2","adep3","adep4","adep5",
                     "adep6","adep7","adep8","adep9","adep10",
                     "awtl11","awtl12","awtl13","awtl14","awtl15",
                     "awtl16","awtl17","awtl18","awtl19","awtl10",
                     "damr11","damr21","damrmn","dec11,
                     Asrfstr_0","dec21,Asrfmet_0","dec12,
                     Abelstr_0","dec22,Abelmet_0","dec31,
                     Asrfmic_0","dec32,kactv_0","dec5,kslow_0",
                     "dec4,kpass_0","Edepth","Elitst",
                     "Fwloss1","Fwloss2","Fwloss3","Fwloss4",
                     "ntspm,CYCL","OMLECH(1)","OMLECH(2)",
                     "OMLECH(3)","P1CO2A1","P1CO2A2","P1CO2B1",
                     "P1CO2B2","P2CO2","P3CO2","pabres",
                     "Peftxa","Peftxb","pligst1","pligst2",
                     "PMCO21","PMCO22","PmnTmp","PmxBio",

```

```

    "PmxTmp", "PS1CO21", "PS1CO22", "PS1S31",
    "PS1S32", "PS2S31", "PS2S32", "Rsplig",
    "spl1", "spl2", "strmax1", "strmax2",
    "teff1", "teff2", "teff3", "Tmelt1", "Tmelt2")
parameters.values <-c(15,15,15,15,30,30,30,30,0,0,0.8,
                      0.6,0.4,0.3,0.2,0.2,0.2,0.2,0,0,0,
                      0.02,15,3.9,14.8,4.9,18.5,6,7.3,
                      0.2,0.0045,0.2,0.4,0.8,0.8,0.65,
                      0.9,4,0.03,0.12,60,0.6,0.17,0,
                      0.68,0.55,0.55,100,0.25,0.75,3,
                      3,0.55,0.55,0.004,600,-0.0035,
                      0.45,0.55,0.003,0.032,0.003,
                      0.009,0.3,0.85,0.013,5000,
                      5000,0,0.125,0.07,-8,4)
parameters <- data.frame(V1=parameters.values,
                          V2=parameters.names)
#initial parameters (site.100)
init.names<-c("xsrfsrstr", "xsrfmet", "xsrfmic", "xbelstr",
              "xbelmet", "xactv", "xslow", "xpass",
              "xwood1", "xwood2", "xwood3",
              "rwcf_initial1", "rwcf_initial2",
              "rwcf_initial3", "rwcf_initial4",
              "rwcf_initial5", "rwcf_initial6",
              "rwcf_initial7", "rwcf_initial8",
              "rwcf_initial9", "rwcf_initial10",
              "asmos1", "asmos2", "asmos3", "asmos4",
              "asmos5", "asmos6", "asmos7", "asmos8",
              "asmos9", "asmos10", "asmos11", "snql",
              "snow", "srfstrlig", "belstrlig")
init.values <-c(240,60,60,186.5,113.4,130,2570,
                1596,500,500,500,0.5,0.5,0.5,0.5,
                0.5,0.5,0.5,0.5,0.5,0.5,0.2,
                0.2,0.2,0.2,0.2,0.2,0.2,0.2,
                0.2,0.2,0.2,0,0,0.275,0.354)
init <- data.frame(V1=init.values,V2=init.names)
#site (site.100)
site.parameters.names <-c("sitlat", "sitlog",

```

```

"sand", "silt", "clay", "bd",
"nlayer", "nlaypg", "drain",
"basef", "stormf",
"SWFLAGflag_fc_wpt(0useobserved, 1.0Guputa)",
"AWILT1", "AWILT2", "AWILT3",
"AWILT4", "AWILT5", "AWILT6",
"AWILT7", "AWILT8", "AWILT9",
"AWILT10",
"AFIEL1", "AFIEL2", "AFIEL3",
"AFIEL4", "AFIEL5", "AFIEL6",
"AFIEL7", "AFIEL8", "AFIEL9",
"AFIEL10", "elev", "xNmineral")

#site sand, silt, clay, bulk density
site.parameters.values <- c(59.36, 13.47, 0.55, 0.15, 0, 1.226,
                           8, 5, 1, 0.5, 0.9, 1, 0.2, 0.2, 0.2, 0.2,
                           0.2, 0.2, 0.2, 0.2, 0.2, 0.2, 0.3, 0.3,
                           0.3, 0.3, 0.3, 0.3, 0.3, 0.3, 0.3, 0.3,
                           50, 1.65)

site.parameters <- data.frame(V1=site.parameters.values,
                               V2=site.parameters.names)

#climate environment (site.100)
envi.parameters.names <- c("Prec(1)cm", "Prec(2)cm",
                           "Prec(3)cm", "Prec(4)cm", "Prec(5)cm",
                           "Prec(6)cm", "Prec(7)cm",
                           "Prec(8)cm", "Prec(9)cm", "Prec(10)cm",
                           "Prec(11)cm", "Prec(12)cm",
                           "Tmin(1)degree", "Tmin(2)degree",
                           "Tmin(3)degree", "Tmin(4)degree",
                           "Tmin(5)degree", "Tmin(6)degree",
                           "Tmin(7)degree", "Tmin(8)degree",
                           "Tmin(9)degree", "Tmin(10)degree",
                           "Tmin(11)degree", "Tmin(12)degree",
                           "Tmax(1)degree", "Tmax(2)degree",
                           "Tmax(3)degree", "Tmax(4)degree",
                           "Tmax(5)degree", "Tmax(6)degree",
                           "Tmax(7)degree", "Tmax(8)degree",
                           "Tmax(9)degree", "Tmax(10)degree",
                           "Tmax(11)degree", "Tmax(12)degree")

```

```

    "Tmax(11)degree", "Tmax(12)degree")
envi.parameters.values <-c(3.395,2.695,2.884,3.051,3.306,
                           4.471,4.623,6.016,5.494,5.221,
                           5.659,3.858,-6.647,-7.235,-4.201,
                           -0.121,4.97,9.538,11.74,11.038,7.266,
                           3.379,-1.027,-5.32,-0.782,-0.375,3.613,
                           9.369,15.549,19.758,21.351,20.219,
                           15.489,9.915,4.286,0.584)
envi.parameters <- data.frame(V1=envi.parameters.values,
                               V2=envi.parameters.names)

#tree
tree.parameters.names <-c("cerfor(1:2:3,1,1),cnr_fol",
                           "cerfor(1:2:3,3,1),cnr_bra",
                           "cerfor(1:2:3,4,1),cnr_ste",
                           "cerfor(1:2:3,2,1),cnr_fir",
                           "cerfor(1:2:3,5,1),cnr_cor",
                           "DECW1,kwood1_0,bra","DECW2,
                           kwood2_0,ste","DECW3,kwood3_0,cor",
                           "forrtf","leafdr1", "leafdr2",
                           "leafdr3","leafdr4", "leafdr5",
                           "leafdr6","leafdr7", "leafdr8",
                           "leafdr9","leafdr10", "leafdr11",
                           "leafdr12",
                           "wdlig1,cfol_lig","wdlig3,cbra_lig",
                           "wdlig4,cste_lig","wdlig2,cfir_lig",
                           "wdlig5,ccor_lig",
                           "woaddr1fol","woaddr3bra",
                           "woaddr4ste","woaddr2fir", "woaddr5cor")

tree.parameters.values <-c(20,99,140,40,83,1.5,0.5,0.6,
                           0.5,0,0,0,0.002,0.006,0.012,
                           0.013,0.039,0.175,0.664,0.066,
                           0.023,0.223,0.25,0.25,0.25,0.25,
                           1,0.01,0.002,0.04,0.004)

tree.parameters <- data.frame(V1=tree.parameters.values,
                               V2=tree.parameters.names)

# biomass components gC.m-2
biomass.in <- data.frame(id=1,

```

```

foliage.tot70=795.954,
branch.tot70=1241.235,
wood.tot70=5110.385,
fineroot.tot70=251.318,
root.tot70=1652.101)

# litter components gC.m-2
litter.in <- data.frame(id=1,
                         foliage.lit.tot70=116.804,
                         branch.lit.tot70=15.515,
                         wood.lit.tot70=12.447,
                         fineroot.lit.tot70=131.778,
                         root.lit.tot70=20.651)

# Define objects from SITE SPECIFIC PARAMETERS: #####
# environment(meteo), site, and tree parameters #####
#site specific parameters
envi <- envi.parameters
tree <- tree.parameters
site <- site.parameters
## define environment #####
# prec: monthly precipitation, cm
# atempmin: monthly minimum air temperature
# atempmax: monthly maximum air temperature
prec<-matrix(0,nrow=12,ncol=1)
atempmin<-matrix(0,nrow=12,ncol=1)
atempmax<-matrix(0,nrow=12,ncol=1)
for(m in 1:12)
{
  prec[m]<-envi[m,1]
  atempmin[m]<-envi[m+12,1]
  atempmax[m]<-envi[m+24,1]
}
## define site parameters #####
# awilt: wilting point
# afiel: field capacity
sitlat<-site[1,1]
sitlog<-site[2,1]
sand<-site[3,1]

```

```

silt<-site[4,1]
clay<-site[5,1]
bd<-site[6,1]
#use mean soil parameters for swedish soils
#(if soil data is not available)
if (is.na(bd)) {
  bd<-1.2
}
if(sum(sand,silt,clay)==0) {
  silt<-0.45
  clay<-0.179
  bd<-0.029
}
nlayer<-as.integer(site[7,1])
nlaypg<-as.integer(site[8,1])
drain<-site[9,1]
basef<-site[10,1]
stormf<-site[11,1]
flag_fc_wpt<-as.integer(site[12,1])
awilt<-matrix(0,nrow=10,ncol=1)
afiel<-matrix(0,nrow=10,ncol=1)
for(i in 1:10)
{
  awilt[i]<-site[12+i,1]
  afiel[i]<-site[22+i,1]
}
elev<-site[33,1]
xNmineral<-site[34,1]
## define init parameters #####
# xsrfstr: surface structural
# xsrfmet: surface metabolic
# xsrfmic: surface microbe
# xbelstr: belowground structural
# xbelmet: belowground metabolic
# xactv: actic pool
# xslow: slow pool
# xpass: passive pool

```

```

# xwood1: branch litter
# xwood2: stem litter
# xwood3: coarse root litter
# rwcf: volumetric soil water content
# asmos: soil water content of the ith soil layer cmh2o
xsrfstr<-init[1,1]
xsrfmet<-init[2,1]
xsrfmic<-init[3,1]
xbelstr<-init[4,1]
xbelmet<-init[5,1]
xactv<-init[6,1]
xslow<-init[7,1]
xpass<-init[8,1]
xwood1<-init[9,1]
xwood2<-init[10,1]
xwood3<-init[11,1]
tawood <- xwood1 + xwood2
tbwood <- xwood3
talit <- xsrfstr + xsrfmet + xsrfmic
tblit <- xbelstr + xbelmet
somsc <- xactv + xslow + xpass
somtc <- xactv + xslow + xpass + xbelstr + xbelmet
rwcf<-matrix(0.1,nrow=10,ncol=1)
for(j in 1:nlayer)
{
  rwcf[j]<-init[11+j,1]
}
asmos<-matrix(0.1,nrow=11,ncol=1)
for(j in 1:(nlayer+1))
{
  asmos[j]<-init[21+j,1]
}
snlq<-init[33,1]
snow<-init[34,1]
srfstrlig<-init[35,1]
belstrlig<-init[36,1]
##
```

```

## define tree parameters #####
# CN ratio of foliage, branch stem, fine roots, coarse roots
# Decomposition constant
# Translocation of N
# Lignin ratios
# Death rate

cnr_fol<-tree[1,1]
cnr_bra<-tree[2,1]
cnr_ste<-tree[3,1]
cnr_fir<-tree[4,1]
cnr_cor<-tree[5,1]
kwood1<-tree[6,1]
kwood2<-tree[7,1]
kwood3<-tree[8,1]
forrtf<-tree[9,1]
leafdr<-matrix(0,nrow=12,ncol=1)
for(j in 1:12)
{
  leafdr[j]<-tree[j+9,1]
}
cfol_lig<-tree[22,1]
cbra_lig<-tree[23,1]
cste_lig<-tree[24,1]
cfir_lig<-tree[25,1]
ccor_lig<-tree[26,1]
wooddr<-matrix(0,nrow=5,ncol=1)
for(j in 1:5)
{
  wooddr[j]<-tree[j+26,1]
}
## define main (FIX) parameters #####
# A: decomposition constant
# k: decomposition constant
adep<-matrix(0.1,nrow=10,ncol=1)
for(j in 1:10)
{
  adep[j]<-parameters[j,1]
}

```

```

}

#
#
awtl<-matrix(0,nrow=10,ncol=1)
for(j in 1:10)
{
  awtl[j]<-parameters[10+j,1]
}
damr11<-parameters[21,1]
damr21<-parameters[22,1]
damrmn<-parameters[23,1]
Asrfstr<-parameters[24,1]
Asrfmet<-parameters[25,1]
Abelstr<-parameters[26,1]
Abelmet<-parameters[27,1]
Asrfmic<-parameters[28,1]
kactv<-parameters[29,1]
kslow<-parameters[30,1]
kpass<-parameters[31,1]
Edepth<-parameters[32,1]
elitst<-parameters[33,1]
fwloss1<-parameters[34,1]
fwloss2<-parameters[35,1]
fwloss3<-parameters[36,1]
fwloss4<-parameters[37,1]
CYCL<-as.integer(parameters[38,1])
omlech<-matrix(0,nrow=3,ncol=1)
omlech[1]<-parameters[39,1]
omlech[2]<-parameters[40,1]
omlech[3]<-parameters[41,1]
P1CO2A1<-parameters[42,1]
P1CO2A2<-parameters[43,1]
P1CO2B1<-parameters[44,1]
P1CO2B2<-parameters[45,1]
Psrfmic<-P1CO2A1
Pactv<-P1CO2A2+P1CO2B2*sand
Pslow<-parameters[46,1]

```

```

Ppass<-parameters[47,1]
pabres<-parameters[48,1]
Peftxa<-parameters[49,1]
Peftxb<-parameters[50,1]
pligst1<-parameters[51,1]
pligst2<-parameters[52,1]
Psrfstr<-parameters[53,1]
Psrfmet<-parameters[54,1]
Pbelstr<-Psrfstr
Pbelmet<-Psrfmet
PmnTmp<-parameters[55,1]
PmxBio<-parameters[56,1]
PmxTmp<-parameters[57,1]
PS1CO21<-parameters[58,1]
PS1CO22<-parameters[59,1]
ps1s31<-parameters[60,1]
ps1s32<-parameters[61,1]
ps2s31<-parameters[62,1]
ps2s32<-parameters[63,1]
RSPLIG<-parameters[64,1]
spl1<-parameters[65,1]
spl2<-parameters[66,1]
strmax1<-parameters[67,1]
strmax2<-parameters[68,1]
teff1<-parameters[69,1]
teff2<-parameters[70,1]
teff3<-parameters[71,1]
Tmelt1<-parameters[72,1]
Tmelt2<-parameters[73,1]
#Biomass data from Swe Forest and Soil Inventory #####
#biomass components gC.m-2
#biomass.in
pools.bfol<- biomass.in[1,2]
pools.bbra<-biomass.in[1,3]
pools.bste<-biomass.in[1,4]
pools.bfir<-biomass.in[1,5]
pools.bcor<-biomass.in[1,6]

```

```

#Litterfall SITE SPECIFIC data
litter.in <- litter.in

# Initialization #####
stempave<-0.0
defac<-0.0
pet<-0.0
anerb<-0.0
CO2out<-0.0
leaching<-0.0
pet<-matrix(0,nrow=12,ncol=1)
avh2o<-matrix(0.0,nrow=3,ncol=1)
amov<-matrix(0.0,nrow=11,ncol=1)
tran<-0.0
evap<-0.0
stream1<-0.0
cleach<-0.0
tcleach<-0.0
#####
#
## Functions of the CENTURY #####
#
##### function (calpet) #####
## potential evapotranspiration
calpet<-function()
{
  # Linacre(1977) from CENTURY source
  #As in the CENTURY
  elev<-0.0
  ave<-matrix(0,nrow=12,ncol=1)
  ave[1]<-(atempmax[1]+atempmin[1])/2.0
  highest<-ave[1]
  lowest<-ave[1]
  for(k in 2:12)
  {
    ave[k]<-(atempmax[k]+atempmin[k])/2.0
    if(ave[k]>highest)

```

```

{
  highest<-ave[k]
}
if(ave[k]<lowest)
{
  lowest<-ave[k]
}
}
if(lowest< (-10.0))
{
  lowest<- (-10.0)
}
ra<-abs(highest-lowest)
for(k in 1:12)
{
  if(atempmin[k]<(-10.0))
  {
    tr<-atempmax[k]-(-10.0)
  }
  else
  {
    tr<-atempmax[k]-atempmin[k]
  }
  t<-tr/2.0+atempmin[k]
  tm<-t+0.006*elev
  td<-0.0023*elev+0.37*t+0.53*tr+0.35*ra-10.9
  e<-((700.0*tm/(100.0-abs(sitlat)))+15.0*td)/(80.0-t)
  monpet<-(e*30.0)/10.0
  if(monpet < 0.5)
  {
    pet[k]<<-0.5*fwloss4
  }
  else
  {
    pet[k]<<-monpet*fwloss4
  }
}
}

```

```

}

## function (calstemp) #####
## soil temperature
calstemp<-function(month)
{
  #For Forest only (e.g. no savana)
  stdead<-0.0
  bio<-(pools.bfol)*2.5+stdead+(xsrfstr+xsrftmet)*2.0*elitst
  if(bio>PmxBio)
  {
    bio<-PmxBio
  }
  else {
    bio<-bio
  }
  stempmax <<-atempmax[month]+
    (25.4/(1+18.0*exp(-0.20*atempmax[month])))*
    (exp(PmxTmp*bio)-0.13)
  stempmin <<-atempmin[month]+PmnTmp*(bio)-1.78
  stempave <<-(stempmax+stempmin)/2.0
}

## function (calfc_wpt) #####
## field capacity and wilting point
calfc_wpt<-function()
{
  #From CENTURY source
  #swflag lets the model user choose between using observed data
  #for awilt and afiel or equations from Gupta and Larson (1979)
  #or Rawls et al (1982).
  #swflag=0
  #Use observed data
  #swflag=1
  #Use G&L for both awilt (-15 bar) and afiel (-0.33 bar)
  #swflag=2
  #Use G&L for both awilt (-15 bar) and afiel (-0.10 bar)
  #swflag=3
  #Use Rawls for both awilt (-15 bar) and afiel (-0.33 bar)
}

```

```

#swflag=4
#Use Rawls for both awilt (-15 bar) and afiel (-0.10 bar)
#swflag=5
#Use Rawls for afiel (-0.33 bar) and observed data for awilt
#swflag=6
#Use Rawls for afiel (-0.10 bar) and observed data for awilt

fcsa<-c( 0.3075,      0.5018,     -0.20,     -0.30,    -0.19,      0.31)
fcsi<-c( 0.5886,      0.8548,      0.0,       0.0,      0.0,       0.0)
fccl<-c( 0.8039,      0.8833,      0.36,      0.23,      0.0,       0.0)
fcom<-c( 2.208E-03,   4.966E-03,   0.0299,   0.0317,   0.0210,   0.0260)
fcbd<-c(-0.1434,     -0.2423,      0.0,       0.0,      0.0,       0.0)
fcwp<-c( 0.0,         0.0,         0.0,       0.0,      0.72,      0.41)
fcin<-c( 0.0,         0.0,         0.2576,   0.4118,   0.2391,   0.4103)
wpsa<-c(-0.0059,     -0.0059,      0.0,       0.0,      0.0,       0.0)
wpsi<-c( 0.1142,      0.1142,      0.0,       0.0,      0.0,       0.0)
wpcl<-c( 0.5766,      0.5766,      0.50,      0.50,      0.0,       0.0)
wpom<-c( 2.228E-03,   2.228E-03,   0.0158,   0.0158,   0.0,       0.0)
wpbd<-c( 0.02671,    0.02671,     0.0,       0.0,      0.0,       0.0)
wpwp<-c( 0.0,         0.0,         0.0,       0.0,      1.0,      1.0)
wpin<-c( 0.0,         0.0,         0.0260,   0.0260,   0.0,       0.0)

#print(somsc)

ompc <- somsc*1.724/(10000*bd*Edepth)
swflag<-flag_fc_wtpt
for(lyr in 1:nlayer)
{
  #Please note:
  #In the original CENTURY model,
  #somsc was not calculated before the call of the prelim.f,
  #so afiel is calculated using somsc=ompc=0.
  #This is a bug of the original CENTURY model
  if(BFix==0)
  {
    omprc<-0.0
  }
  afiel[lyr] <- fcsa[swflag]*sand + fcsi[swflag]*silt +
    fccl[swflag]*clay + fcom[swflag]*ompc +
    fcbd[swflag]*bd + fcwp[swflag]*awilt[lyr] +

```

```

    fcin[swflag]

awilt[lyr] <<- wpsa[swflag]*sand + wpsi[swflag]*silt+
            wpcl[swflag]*clay + wpom[swflag]*ompc +
            wpbd[swflag]*bd + wpwp[swflag]*awilt[lyr] +
            wpin[swflag]

ompc<-ompc*0.85
}

}

## function (calwater) #####
## soil water content
calwater<-function(month)
{
  #Initialize
  add<-0.0
  amelt<-0.0
  asimx<-0.0
  avh2o[1]<<-0.0
  avh2o[2]<<-0.0
  avh2o[3]<<-0.0
  avap<-0.0
  evl<-0
  pevp<-0.0
  pttr<-0.0
  rwc1<-0.0
  tran<<-0.0
  trap<-0.01
  aabs<-0.0
  evsnow<-0.0
  evap<<-0.0
  petrem<-pet[month]
  awwt<-matrix(0.0,nrow=11,ncol=1)
  #CO2 effect was not included here
  co2val<-1.0
  irrract<-0.0
  inputs<-prec[month]+irrract
  winputs<-inputs
  atempave<<-(atempmax[month]+atempmin[month])/2.0
}

```

```

aliv<-pools.bfol*2.5
alit<-(xsrfstr+xsrftmet)*2.0
adead<-0.0
*****  

#Snow
#Snowfall
if(atempave <= 0.0)
{
  #snow <- snow + prec[month]
  snow <<- snow + inputs
  winputs<-0.0
}
# melt
if(atempave >= Tmelt1)
{
  melt <- Tmelt2 * (atempave -Tmelt1)
  if(melt>snow)
  {
    melt<-snow
  }
  snow <<-snow-melt
##.....
if((atempave > 0.0) && (snow > 0.0))
{
  snlq<<-snlq+inputs
}
snlq<<-snlq+melt
if(snlq >= (0.05*snow) )
{
  add<-snlq -0.05*snow
  snlq<<-snlq-add
}
}
if(snow > (0.0))
{
  evsnow<-petrem*0.87
  snow1<-snow+snlq
}

```

```

if(evsnow > snow1)
{
    evsnow<-snow1
}
snow<<-snow-evsnow*(snow/snow1)
snlq<<-snlq-evsnow*(snlq/snow1)
evap<<-evap+evsnow
petrem<-petrem-evsnow/0.87
if(petrem < 0.0)
{
    petrem<-0.0
}
}
if(snow <= 0.0)
{
    sd<-aliv+adead
    if(sd > 800.0)
    {
        sd<-800.0
    }
    if(alit > 400.0)
    {
        alit<-400.0
    }
    aint<-(0.0003 * alit +0.0006 *sd) *fwloss1
    aabs<-0.5*exp((-0.002*alit)-(0.004*sd))*fwloss2
    if((aabs+aint)*inputs<0.4*petrem)
    {
        evl<-(aabs+aint)*winputs
    }
    else
    {
        evl<-0.4*petrem
    }
    evap<<-evap+evl
    add<-add+winputs -evl
    trap<-petrem-evl
}

```

```

}

if(atempave < 2.0)
{
  pttr<-0.0
}
else
{
  pttr<-petrem *0.65 *(1.0 -exp(-0.020 *aliv)) *co2val
}
if(pttr <= trap){trap<-pttr}
if(trap <= 0.0){trap<-0.01}
#####
#hptrr is not included
pevp<-petrem -trap -evl
if(evap<0.0){pevp<-0.0}
if((trap-0.01) < add)
{
  #print(add)
  tran<<- trap-0.01
}
else
{
  tran<<- add
}
trap<-trap-tran
add<-add-tran
strm<-0.0
base<-0.0
stream1<<-0.0
for(j in 1:nlayer)
{
  asmos[j]<<-asmos[j]+add
  afl<-adep[j]*afiel[j]
  if(asmos[j]>afl)
  {
    amov[j]<<-asmos[j]-afl
    asmos[j]<<-afl
}
}

```

```

if(j == nlayer)
{
    strm<-amov[j]*stormf
}
else
{
    amov[j]<<-0.0
}
add<-amov[j]
}

asmos[nlayer+1]<<-asmos[nlayer+1]+add-strm
base<-asmos[nlayer+1]*basef
asmos[nlayer+1]<<-asmos[nlayer+1]-base
stream1<<-strm+base
asimx<-asmos[1]
rwcl<-0.0
tot<-0.0
tot2<-0.0
for(j in 1:nlayer)
{
    avw<-asmos[j]-awilt[j]*adep[j]
    if(avw < 0.0)
    {
        avw<-0.0
    }
    awwt[j]<-avw*awtl[j]
    tot<-tot+avw
    tot2<-tot2+awwt[j]
}
if(tot<trap)
{
    trap<-tot
}
else
{
    trap<-trap
}

```

```

}

if(tot2 > 0.0)
{
  for(j in 1:nlayer)
  {
    avinj<-asmos[j]-awilt[j]*adep[j]
    if(avinj < 0.0)
    {
      avinj<-0.0
    }
    trl<-(trap*awwt[j])/tot2
    if(trl > avinj)
    {
      trl<-avinj
    }
    asmos[j]<<-asmos[j]-trl
    #if(year==5 && month==1){cat(asmos[j], trl, "\n") }
    avinj<-avinj-trl
    tran<<-tran+trl
    rwcf[j]<-(asmos[j]/adep[j]-awilt[j])/(afiel[j]-awilt[j])
    if(j<=nlaypg)
    {
      avh2o[1]<<-avh2o[1]+avinj
    }
    avh2o[2]<<-avh2o[2]+avinj
    if(j <= (2))
    {
      avh2o[3]<<-avh2o[3]+avinj
    }
  }
}
fwlos<-0.25
evmt<-(rwcf[1]-fwlos)/(1.0-fwlos)
if(evmt <= (0.01))
{
  evmt<-0.01
}

```

```

evlos<-evmt*pевп*aabs*0.10
avinj<-asmos[1]-awilt[1]*adep[1]
if(avinj < 0.0)
{
  avinj<-0.0
}
if(evlos > avinj)
{
  evlos<-avinj
}
asmos[1]<<-asmos[1]-evlos
evap<<-evap+evlos
avhsm<-(asmos[1]+rwcl*asimx)/(1.0+rwcl)
rwcf[1]<<-(avhsm/adep[1]-awilt[1])/(afiel[1]-awilt[1])
avh2o[1]<<-avh2o[1]-evlos
avh2o[2]<<-avh2o[2]-evlos
avh2o[3]<<-avh2o[3]-evlos
}
## function (caldefac) #####
## decomposition factor
caldefac<-function()
{
  if(snow > 0.0)
  {
    stempave<-0.0
  }
  # Cal defac
  tfunc<-teff1+teff2*exp(teff3 * stempave)
  rprpet <<- (avh2o[3] + prec[month] ) / pet[month]
  #* idef in fix.100 in Century control linear 1 or ratio 2 option
  #* this is idef==2
  if(rprpet > 9.0 )
  {
    wfunc<<-1.0
  }
  else
  {

```

```

wfunc<<-1.0/(1.0+30.0*exp(-8.5*rprpet))
}

#if(wfunc>1.0)
#{
#  wfunc<<-1.0
#}
defac<<-tfunc*wfunc
##
# If you want to use the defac from the original CENTURY, then.
# defac<<-centdefac[1+(year-1)*12+month,2]
#*** Cal anerb
anerb <<- 1.0
}

## functions to Divide litter inputs#####
## 
## function (calcenturyinput) #####
## litter inputs into each soil carbon pool:1
calcenturyinput<-function()
{
  if(flows.lfinfol>0.0)
  {
    #centurypartit(1, cnr_srflit)
    centurypartit(1, cnr_fol)
  }
  else
  {
    usrfstr <<-0.0
    usrfmet <<-0.0
    usrfstr_lig <<- 0.0
  }
  if(flows.lfinfir>0.0)
  {
    #centurypartit(2, cnr_bellit)
    centurypartit(2, cnr_fir)
  }
  else
  {

```

```

ubelstr <<- 0.0
ubelmet <<- 0.0
ubelstr_lig <<- 0.0
}
uwood1 <<- flows.lfinbra
uwood2 <<- flows.lfinste
uwood3 <<- flows.lfincor
uwood1_lig <<- flows.lfinbra * cbra_lig
uwood2_lig <<- flows.lfinste * cste_lig
uwood3_lig <<- flows.lfincor * ccor_lig
}

## function (centurypartit) #####
## litter inputs into each soil carbon pool:2
centurypartit<-function(p, cnr)
{
  #translocation
  #forrtf
  if(p==1)
  {
    cpart<- flows.lfinfo1
    epart<- flows.lfinfo1*(1.0/cnr)*(1.0-forrtf)
    #amax1
    if(cpart/pabres > 1.0)
    {
      s<-cpart/pabres
    }
    else
    {
      s<-1.0
    }
    #damr11<-0.0
    dirabs<- damr11 * xNmineral * s
    if((epart+dirabs) <= 0.0)
    {
      rcetot<-0.0
    }
  }
  else

```

```

{
  rcetot<-cpart/(epart+dirabs)
}
if(rcetot < damrmn)
{
  dirabs<-cpart/damrmn-epart
}
if(dirabs<0.0)
{
  dirabs<-0.0
}
frlign<- cfol_lig
}
else if (p==2)
{
  cpart<- flows.lfinfir
#epart<- flows.lfinfir*(1.0/cnr)*(1.0-forrtf)
  epart<- flows.lfinfir*(1.0/cnr)
#amax1
  if(cpart/pabres > 1.0)
  {
    s<-cpart/pabres
  }
  else
  {
    s<-1.0
  }
  dirabs<- damr21 * xNmineral * s
  if((epart+dirabs) <= 0.0)
  {
    rcetot<-0.0
  }
  else
  {
    rcetot<-cpart/(epart+dirabs)
  }
  if(rcetot < damrmn)

```

```

{
  dirabs<-cpart/damrmn-eprt
}
if(dirabs<0.0)
{
  dirabs<-0.0
}
fralign<- cfir_lig
}
else
{
  printf("error")
}
###  

frn<- (eprt + dirabs) / (cpart*2.5)
rlnres<-fralign/frn
frmst<- spl1 -spl2 *rlnres
if(fralign > (1.0-frmst))
{
  frmst<- 1.0-fralign
}
if(frmst<0.20)
{
  frmst<-0.20
}
caddm <- cpart*frmst
if(caddm < 0.0)
{
  caddm<-0.0
}
cadds <- cpart-caddm
fligst <- fralign/(cadds/cpart)
if(fligst > 1.0)
{
  fligst <- 1.0
}
if(p==1)

```

```

{
  usrfstr <<- flows.lfinfo1 *(1.0-frmet)
  usrfmet <<- flows.lfinfo1 *frmet
  usrfstr_lig <<- fligst
}
else if (p==2)
{
  ubelstr <<- flows.lfinfir *(1.0-frmet)
  ubelmet <<- flows.lfinfir *frmet
  ubelstr_lig <<- fligst
}
}

#####
## functions (calcentury)
## to calculate soil carbon dynamics
## ****
calcentury<-function()
{
  uwood1<-flows.lfinbra
  uwood2<-flows.lfinste
  uwood3<-flows.lfincor
  #*****
  # ** Dead branch = Wood 1
  #strlig=(xwood1*wood1strlig+uwood1_lig)/(xwood1+uwood1)
  #wood1strlig= strlig
  strlig <-cbra_lig
  if(xwood1>0.000001)
  {
    tcflow <- xwood1*defac*kwood1*exp(-pligst1*strlig)*DT
    if(tcflow>xwood1)
    {
      tcflow<-xwood1
    }
  }
  else
  {
    tcflow<-0.0
  }
}

```

```

}

tsom2_fwood1 <- tcflow * strlig

#*Respiration associated with decomposition to som2
co2los <- tsom2_fwood1 * RSPLIG
CO2out <<- CO2out+co2los

#*Net C flow to SOM2
tsom2_fwood1 <- tsom2_fwood1 - co2los
tsom1_fwood1 <- tcflow - tsom2_fwood1 - co2los
#*Respiration associated with decomposition to som1
co2los <- tsom1_fwood1 * PS1C021
tsom1_fwood1 <- tsom1_fwood1 -co2los
CO2out<<-CO2out+co2los

*****
xwood1_new <- xwood1 + uwood1 - tcflow
*****  

# ** Dead Stem = Wood 2
#strlig=(xwood2*wood2strlig+uwood2_lig)/(xwood2+uwood2)
#wood2strlig= strlig
strlig<-cste_lig
if(xwood2>0.000001)

{
  tcflow<- xwood2*defac*kwood2*exp(-pligst1*strlig)*DT
  if(tcflow>xwood2)
  {
    tcflow<-xwood2
  }
}
else
{
  tcflow<-0.0
}
tsom2_fwood2 <- tcflow * strlig

#*Respiration associated with decomposition to som2
co2los <- tsom2_fwood2 * RSPLIG
CO2out<<-CO2out+co2los

#*Net C flow to SOM2
tsom2_fwood2 <- tsom2_fwood2 - co2los

```

```

tsom1_fwood2 <- tcflow - tsom2_fwood2 - co2los
#*Respiration associated with decomposition to som1
co2los <- tsom1_fwood2 * PS1C021
tsom1_fwood2 <- tsom1_fwood2 -co2los
CO2out<<-CO2out+co2los
*****
xwood2_new <- xwood2 + uwood2 - tcflow
*****
# ** Dead Coarse root = Wood 3
#strlig=(xwood3*wood3strlig+uwood3_lig)/(xwood3+uwood3)
#wood3strlig= strlig
strlig<-ccor_lig
if(xwood3>0.000001)
{
  tcflow<- xwood3*defac*kwood3*exp(-pligst2*strlig)*anerb*DT
  if(tcflow>xwood3)
  {
    tcflow<-xwood3
  }
}
else
{
  tcflow<-0.0
}
tsom2_fwood3 <- tcflow * strlig
#*Respiration associated with decomposition to som2
co2los <- tsom2_fwood3 * RSPLIG
CO2out<<-CO2out+co2los
#*Net C flow to SOM2
tsom2_fwood3 <- tsom2_fwood3 - co2los
tsom1_fwood3 <- tcflow - tsom2_fwood3 - co2los
#*Respiration associated with decomposition to som1
co2los <- tsom1_fwood3 * PS1C021
tsom1_fwood3 <- tsom1_fwood3 -co2los
CO2out<<-CO2out+co2los
*****
xwood3_new <- xwood3 + uwood3 - tcflow

```

```

*****  

# ** surface structural  

#strlig=(pools.xlig1_fol + pools.xlig1_bra +  

#          pools.xlig1_ste)/(pools.talit)  

#srfstrlig = xsrfstr*srfstrlig/xsrfstr  

#strlig=(xsrfstr*srfstrlig + usrfstr_lig)/(xsrfstr+usrfstr)  

strlig<- (xsrfstr*srfstrlig + usrfstr_lig*usrfstr)/  

          (xsrfstr + usrfstr)  

srfstrlig <<- strlig  

if(xsrfstr>0.000001)  

{  

  if(xsrfstr>strmax1)  

  {  

    mass<-strmax1  

  }  

  else  

  {  

    mass<-xsrfstr  

  }  

  tcflow <-mass*defac*Asrfstr*exp(-pligst1*strlig)*DT  

  if(tcflow>xsrfstr)  

  {  

    tcflow<-xsrfstr  

  }  

}  

else  

{  

  tcflow<-0.0  

}  

tsom2_fsrfstr <- tcflow * strlig  

#*Respiration associated with decomposition to som2  

co2los <- tsom2_fsrfstr * RSPLIG  

CO2out <<- CO2out+co2los  

#*Net C flow to SOM2  

tsom2_fsrfstr <- tsom2_fsrfstr - co2los  

tsom1_fsrfstr <- tcflow - tsom2_fsrfstr - co2los  

#*Respiration associated with decomposition to som1

```

```

co2los <- tsom1_fsrstr * PS1CO21
tsom1_fsrstr <- tsom1_fsrstr -co2los
CO2out <<- CO2out+co2los
*****
xsrfstr_new <- xsrfstr + usrfstr - tcflow
*****
# ** soil structural
#strlig=(pools.xlig1_fir+pools.xlig1_cor)/(pools.tblit)
#belstrlig = xbelstr*belstrlig/xbelstr
strlig<- (xbelstr*belstrlig + ubelstr_lig*ubelstr) /
(xbelstr + ubelstr)
belstrlig<<-strlig
if(xbelstr>0.000001)
{
  if(xbelstr>strmax2)
  {
    mass<-strmax2
  }
  else
  {
    mass<-xbelstr
  }
  tcflow<-mass*defac*Abelstr*exp(-pligst2*strlig)*anerb*DT
  if(tcflow>xbelstr)
  {
    tcflow<-xbelstr
  }
}
else
{
  tcflow<-0.0
}
tsom2_fbelstr <- tcflow * strlig
#*Respiration associated with decomposition to som2
co2los <- tsom2_fbelstr * RSPLIG
CO2out <<- CO2out+co2los
#*Net C flow to SOM2

```

```

tsom2_fbelstr <- tsom2_fbelstr - co2los
tsom1_fbelstr <- tcflow - tsom2_fbelstr - co2los
#*Respiration associated with decomposition to som1
co2los <- tsom1_fbelstr * PS1CO22
tsom1_fbelstr <- tsom1_fbelstr -co2los
CO2out<<-CO2out+co2los
*****
xbelstr_new <- xbelstr + ubelstr - tcflow
*****
# ** surface metab
if(xsrfmet>0.000001)
{
  tcflow<-xsrfmet * defac * Asrfmet * DT
  if(tcflow>xsrfmet)
  {
    tcflow<-xsrfmet
  }
}
else
{
  tcflow<-0.0
}
co2los<-tcflow*Psrfmet
tsom1_fsrfmet <- tcflow-co2los
CO2out <<- CO2out+co2los
xsrfmet_new <- xsrfmet +usrfmet -tcflow
*****
# ** belowground metab
if(xbelmet>0.000001)
{
  tcflow<-xbelmet * defac * Abelmet* anerb * DT
  if(tcflow>xbelmet)
  {
    tcflow<-xbelmet
  }
}
else

```

```

{
  tcflow<-0.0
}
co2los<-tcflow*Pbelmet
tsom1_fbelmet<-tcflow-co2los
CO2out<<-CO2out+co2los
xbelmet_new<-xbelmet + ubelmet -tcflow
*****#***** surface microbe
if(xsrfmic>0.000001)
{
  tcflow <- xsrfmic * defac * Asrfmic *DT
  if(tcflow>xsrfmic)
  {
    tcflow <- xsrfmic
  }
}
else
{
  tcflow<-0.0
}
co2los<-tcflow*Psrfmic
tsom2_fsrfmic<-tcflow-co2los
CO2out<<-CO2out+co2los
xsrfmic_new <- xsrfmic + tsom1_fsrfstr + tsom1_fsrfmet +
           tsom1_fwood1 + tsom1_fwood2 -tcflow
#xsrfmic_new= -tcflow + xsrfmic + tsom1_fsrfstr+tsom1_fsrfmet
#xsrfmic_new= -tcflow + xsrfmic + tsom1_fsrfmet
*****#***** active
eftext <- Peftxa + Peftxb * sand
if(xactv>0.000001)
{
  tcflow<-xactv* defac * eftext * kactv * anerb *DT
  if(tcflow>xactv)
  {
    tcflow<-xactv

```

```

        }
    }
else
{
    tcflow<-0.0
}
co2los<-tcflow*Pactv
#*cfsfs2=tcflow-co2los
#*tcflow=tcflow-co2los
CO2out<<-CO2out+co2los
fps1s3 <- ps1s31 + ps1s32 * clay
tsom3_factv<-tcflow * fps1s3
#leaching
if(amov[2]>0.0)
{
    orglch<-omlech[1]+omlech[2]*sand
    t<-1.0-(omlech[3]-amov[2])/omlech[3]
    if(t>1.0)
    {
        linten<-1.0
    }
    else
    {
        linten<-t
    }
    cleach<<-tcflow * orglch * linten
}
else
{
    cleach<<-0.0
}
tcleach<<-tcleach+cleach
tsom2_factv<-tcflow -co2los -tsom3_factv -cleach
#* Updated at the end.
#xactv_new = xactv + tsom1_fbelstr +tsom1_fbhelmet +
#           tsom1_fwood3 +tsom1_fslow +tsom1_fpass -tcflow
xactv_new <- xactv + tsom1_fbelstr +tsom1_fbhelmet +

```

```

tsom1_fwood3 -tcflow
*****
#**** Slow
if(xslow>0.000001)
{
  tcflow<-xslow *defac * kslow * anerb *DT
  if(tcflow>xslow)
  {
    tcflow<-xslow
  }
}
else
{
  tcflow<-0.0
}
co2los<-tcflow*Pslow
#*cfsfs2=tcflow-co2los
#*tsom3_fslow=tcflow-co2los
CO2out<<-CO2out+co2los
xslow_new <- xslow + tsom2_fsrfstr +tsom2_fsrfmic +
           tsom2_fbclstr +tsom2_factv + tsom2_fwood1 +
           tsom2_fwood2 + tsom2_fwood3 -tcflow
fps2s3 <- ps2s31 + ps2s32 * clay
tsom3_fslow<-tcflow * fps2s3
tsom1_fslow<-tcflow -co2los -tsom3_fslow
*****
#**** Passive
if(xpass>0.000001)
{
  tcflow<-xpass *defac * kpass* anerb *DT
  if(tcflow>xpass)
  {
    tcflow<-xpass
  }
}
else
{

```

```

    tcflow<-0.0
}
co2los<-tcflow*Ppass
#*cfsfs2=tcflow-co2los
tsom1_fpass<-tcflow-co2los
CO2out<<-CO2out+co2los
xpass_new <- xpass + tsom3_factv +tsom3_fslow -tcflow
*****
#***** Active new
#xactv_new = xactv + tsom1_fpass
#xactv_new = xactv + tsom1_fslow
xactv_new <- xactv_new + tsom1_fslow + tsom1_fpass
*****
#***** UPDATE
xsrfstr <<- xsrfstr_new
xsrfmet <<- xsrfmet_new
xsrfmic <<- xsrfmic_new
xbelstr <<- xbelstr_new
xbelmet <<- xbelmet_new
xactv <<- xactv_new
xslow <<- xslow_new
xpass <<- xpass_new
xwood1 <<- xwood1_new
xwood2 <<- xwood2_new
xwood3 <<- xwood3_new
*****
somsc <<- xactv + xslow + xpass
talit <<- xsrfstr + xsrfmet + xsrfmic
tblit <<- xbelstr + xbelmet
somtc <<-xactv + xslow + xpass + xbelstr + xbelmet
tawood <<- xwood1 + xwood2
tbwood <<- xwood3
}
#endif of functions
## calculate field capacity, wilting point #####
awilt
afiel

```

```

if(flag_fc_wpt>0.0)
{
  somsc <- xactv + xslow + xpass
  calfc_wpt()
}
awilt
afiel
## Initialize soil water condition #####
# essential to calculate deep asmos correctly
pet
calpet()
pet
for(month in 1:12)
{
  calwater(month)
}
obj.s <- ls()
#obj.s
#####
## MAIN CENTURY SIMULATION #####
carbon.out <- NULL
for(s in 1:1){
  id<-s
  soil.carbon.year.out <-NULL
  for(year in TSTART:TEND){
    CO2out<-0.0
    calpet()
    #month loop
    for(month in 1:12){
      #month=1
      tclease<-0.0
      DT<-1.0/(12.0*CYCL)
      #####.
      #Litterfall SITE SPECIFIC data
      flows.lfinfol<-litter.in [1,2]*leafdr[month]*(1.0/CYCL)
      flows.lfinbra<-litter.in [1,3]*(1.0/(12*CYCL))
      flows.lfinste<-litter.in [1,4]*(1.0/(12*CYCL))
    }
  }
}

```

```

flows.lfinfir<-litter.in [1,5]*(1.0/(12*CYCL))
flows.lfincor<-litter.in [1,6]*(1.0/(12*CYCL))
talit <-xsrfstr + xsrfmet +xsrfmic
tblit <- xbelstr + xbelmet
tawood <- xwood1 + xwood2
tbwood <- xwood3
somsc <- xactv + xslow + xpass
somtc <- xactv + xslow + xpass + xbelstr + xbelmet
##.....
calstemp(month)
calwater(month)
if(snow>0.0)
{
  stempmax <-0.0
  stempmin <-0.0
  stempave <-0.0
}
##.....
caldefac()
calcenturyinput()
# CENTURY CARBON FUNCTION SIMULATIONS
# updated 4 times per month (CYCL=4)
for(i in 1:CYCL)
{
  calcentury()
}
#end of centurycal CYCL loop
}
#end of month loop ()
##.....
## site specific output of CENTURY carbon
if(year==year) #TEND)
{
  soil.carbon0 <- data.frame(id,year, month,
                                xsrfstr, xsrfmet,
                                xsrfmic, xbelstr, xbelmet,
                                xactv, xslow, xpass, somsc,

```

```

        xwood1, xwood2, xwood3,
        CO2out, somtc)
    }

soil.carbon.year.out <- rbind(soil.carbon.year.out,
                                soil.carbon0)
}

#end of year loop
carbon.out <-rbind(carbon.out,soil.carbon.year.out )
}

#end of site.group for loop
options(digit=12)
century.out <-carbon.out[,c("id","year","month",
                            "CO2out","somsc",
                            "xbelstr","xbelmet",
                            "xactv","xslow","xpass",
                            "somtc")]

#convert gC.m-2 to tC.ha-1 by 1/1e6*1e4
century.out[,4:11]<-century.out[,4:11]/100
##.....
#plot components of soil carbon stock
#somtc <- xactv + xslow + xpass + xbelstr + xbelmet
#figure
par(mfrow=c(1,1), mar=c(5,5,1,1))
plot(century.out$year,century.out$somtc,
      log="y", ylim=c(0.3,round(max(century.out$somtc),1)+50),
      ylab="CENTURY soil carbon pools (tC/ha)",
      xlab="year")
lines(century.out$year,century.out$somtc,
      lwd=2)
lines(century.out$year,century.out$xactv,
      col="blue", lwd=2)
lines(century.out$year,century.out$xslow,
      col="red", lwd=2)
lines(century.out$year,century.out$xpass,
      col="orange", lwd=2)
lines(century.out$year,century.out$xbelstr,

```

```
col="grey", lwd=2)
lines(century.out$year,century.out$xbelmet,
      col="magenta", lwd=2)
legend("bottomright",
       c("total","active","slow","passive",
         "bg.structural","bg.metabolic"),
       col=c("black","blue","red","orange","grey","magenta"),
       pch=c(1,NA,NA,NA,NA,NA),
       lwd=2, lty=1, border="white",bg="white")
```

Table S1. Statistical characteristics (mean, standard error) of basic variables for groups of soils derived by recursive partitioning including soil variables (see Fig. 2a), compared with interpretation of carbon, moisture, and fertility of groups.

N of soil samples in groups		959	909	136	335	182	296	180	8	142	83
Total SOC ($tC\ ha^{-1}$)	Mean	65.1	81.8	130.2	86.2	126.4	103.9	136.8	268.6	143.7	203.1
	SE	1	1.3	5.1	2.6	4.8	2.8	4.2	23.7	6.7	9.8
SOC mineral ($tC\ ha^{-1}$)	Mean	45.4	56.4	86.9	68.5	98.4	73.2	92.4	230.6	108.8	153.3
	SE	1.6	2.1	8.2	4.9	9.1	4.5	7	33	12.8	21.6
C/N	Mean	13.4	18.6	15.4	10.7	8.1	23.8	21.7	23.1	23.3	32.6
	SE	0.3	0.4	0.8	0.2	0.2	0.5	0.5	1.9	0.6	2
H100 (m)	Mean	20.8	24.3	24.7	25.4	26.6	22.2	29.9	31	23.6	24
	SE	0.2	0.2	0.5	0.4	0.4	0.3	0.3	0.5	0.5	0.6
Total Litter ($tC\ ha^{-1}$)	Mean	2.7	3.3	3.4	3.5	3.6	3.1	4	3.8	3.2	3.5
	SE	0	0	0.1	0.1	0.1	0	0.1	0.2	0.1	0.1
Temperature air (C)	Mean	3.3	4.8	5.1	5.1	5.8	4.5	7.3	7.3	5.3	6.4
	SE	0.1	0.1	0.2	0.1	0.1	0.1	0.1	0.2	0.2	0.2
Long-term moisture (%)	Mean	20.2	22.4	26	23.6	26.2	22.9	22.9	21.3	21.8	25.7
	SE	0.2	0.2	0.6	0.3	0.5	0.4	0.4	1.3	0.5	0.7
Precipitation (mm y-1)	Mean	698.8	712.9	697.1	644.1	630.3	693	817.2	1173.2	687.9	619.4
	SE	5.7	7	18.8	10.4	12.5	10.8	21.2	162	18.1	27
CEC ($mmol_c\ kg^{-1}$)	Mean	4.7	12.1	11.6	49.2	91.5	24	23.5	24.5	59.7	98.7
	SE	0.1	0.1	0.2	2.6	5.1	0.3	0.3	1.7	3.2	7.6
pH	Mean	5.2	5.1	5.1	5.5	5.6	5	4.8	4.6	4.9	6
	SE	0	0	0	0	0.1	0	0	0.1	0.1	0.2
Clay content (%)	Mean	0.8	1.1	4.2	5.9	21.5	1.7	2.4	2.5	2.4	8
	SE	0.1	0.1	0.5	0.5	1	0.3	0.3	1.3	0.4	1.2
Silt content (%)	Mean	15.1	14.5	29	27.2	57.8	16.4	17.9	18.8	17.1	32.5
	SE	0.3	0.3	1.5	1.2	1.4	0.7	0.7	3.2	1	3
Carbon	low	medium	high	medium	high	medium	high	extra	high	extra	
Moisture	dry-fresh	fresh	moist-fresh	fresh	moist-fresh	fresh	fresh	fresh	fresh	moist-fresh	
Fertility	low	medium	medium	medium	high	low	high	high	medium	medium	
Soil group	1	2	3	4	5	6	7	8	9	10	

Table S2. Statistical characteristics (mean, standard error) of basic environmental and soil variables for regression tree of data groups classified by recursive partitioning with data excluding soil variables (see Fig. 3).

Number of samples		735.0	932.0	796.0	711.0	56.0
Total soil carbon (tC ha^{-1})	Mean	67.1	85.4	96.5	120.1	179.0
	SE	1.4	1.5	2.0	2.4	12.5
C/N	Mean	15.0	15.8	16.8	18.1	35.2
	SE	0.3	0.3	0.4	0.4	3.2
H100 (m)	Mean	19.0	19.7	27.0	30.1	18.7
	SE	0.1	0.1	0.1	0.2	0.3
Total Litter (tC ha^{-1})	Mean	2.3	2.8	3.6	4.0	3.0
	SE	0.0	0.0	0.0	0.0	0.1
Soil water content (%)	Mean	20.2	24.2	22.0	22.6	22.3
	SE	0.2	0.2	0.2	0.2	1.0
Temperature air (C)	Mean	2.4	2.7	6.4	7.2	7.2
	SE	0.1	0.1	0.0	0.0	0.0
pH	Mean	5.2	5.2	5.3	4.9	7.2
	SE	0.0	0.0	0.0	0.0	0.1
Sand content (%)	Mean	52.7	51.5	47.2	55.1	36.2
	SE	0.5	0.5	0.8	0.6	4.0
Clay content (%)	Mean	0.8	2.6	6.1	3.0	5.8
	SE	0.1	0.2	0.4	0.2	1.1
Silt content (%)	Mean	14.6	19.2	25.3	20.0	29.5
	SE	0.3	0.5	0.8	0.5	4.1
Group acronym		low-C cold.pine	median-C cold.other	median-C warm.rainy low-N	high-C warm.rainy high-N	extra-C warm.dry

Table S3. Species and classes of ground vegetation grouped into functional types (1-dwarfshrubs, 2-herbs, 3-grasses, 4-mosses and 5-lichens).

Functional type	Ground vegetation
1	<i>Vaccinium myrtillus</i> , <i>Vaccinium vitis-idaea</i> , <i>Arctostaphylos uva-ursi</i> , <i>Empetrum nigrum</i> ssp, <i>Calluna vulgaris</i> , <i>Erica tetralix</i> , <i>Vaccinium uliginosum</i> , <i>Rhododendron tomentosum</i> , <i>Andromeda polifolia</i> , <i>Vaccinium oxycoccus/microcarpum</i> , Other field layer plants
2	<i>Gymnocarpium dryopteris</i> , <i>Oxalis acetosella</i> , <i>Anemone nemorosa</i> , <i>Maianthemum bifolium</i> , <i>Chamaenerion angustifolium</i> , <i>Anthriscus sylvestris</i> , <i>Melampyrum pratense/sylvaticum</i> , <i>Equisetum sylvaticum</i> , <i>Menyanthes trifoliata</i> , <i>Rubus chamaemorus</i> , <i>Phegopteris connectilis</i> , <i>Hepatica nobilis</i> , <i>Geum rivale</i> , <i>Urtica dioica</i> , <i>Rumex acetosa</i> , <i>Stellaria nemorum</i> , <i>Stellaria holostea</i> , <i>Silene dioica</i> , <i>Aconitum lycoctonum subsp. septentrionale</i> , <i>Actaea erythrocarpa</i> , <i>Trollius europaeus</i> , <i>Cardamine bulbifera</i> , <i>Filipendula ulmaria</i> , <i>Mercurialis perennis</i> , <i>Sanicula europaea</i> , <i>Aegopodium podagraria</i> , <i>Librar</i> , <i>Galium odoratum</i> , <i>Lamiastrum galeobdolon</i> , <i>Stachys sylvatica</i> , <i>Cirsium palustre</i> , <i>Cirsium heterophyllum</i> , <i>Lactuca alpina</i> , <i>Lactuca muralis</i> , <i>Crepis paludosa</i> , <i>Paris quadrifolia</i> , <i>Neottia ovata</i> , <i>Geranium sylvaticum</i> , <i>Rubus idaeus</i> , Other large grown ferns
3	Broad-leaved grass, Narrow-leaved grass, <i>Carex globularis</i> , Other sedges
4	<i>Pteridium aquilinum</i> , <i>Lycopodiaceae</i> , <i>Spagnum spp</i> , <i>Polytrichum commune</i> , <i>Pleurozium schreberi</i> , <i>Hylocomium splendens</i> , Other bryophytes
5	<i>Cladonia</i> , <i>Stereocaulon spp</i> , <i>Cladina spp</i> , <i>Cladonia and Cladina</i> , Other lichens

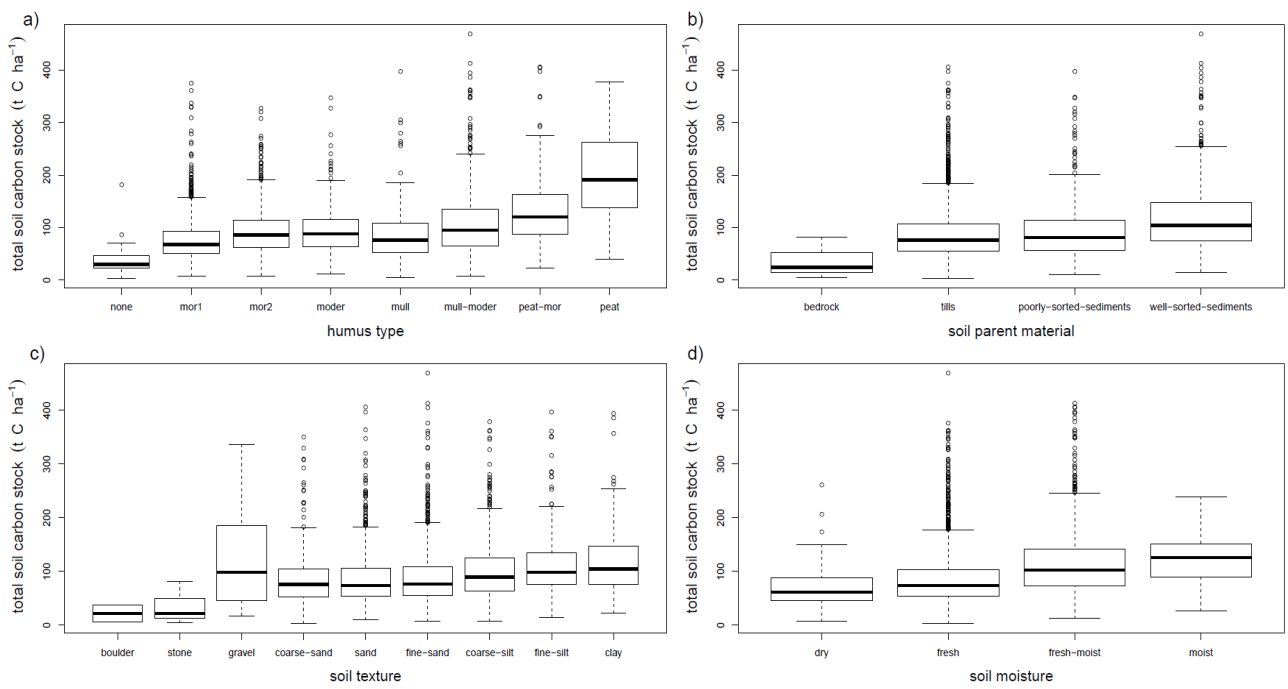


Figure S1. Boxplot main levels (minimum, 1st quantile, median, 3rd quantile, maximum, and dots for outliers) of the total soil carbon stock ($t\text{C ha}^{-1}$) for SFSI categorical data on a) humus type, b) soil parent material, c) soil texture, and d) soil moisture.

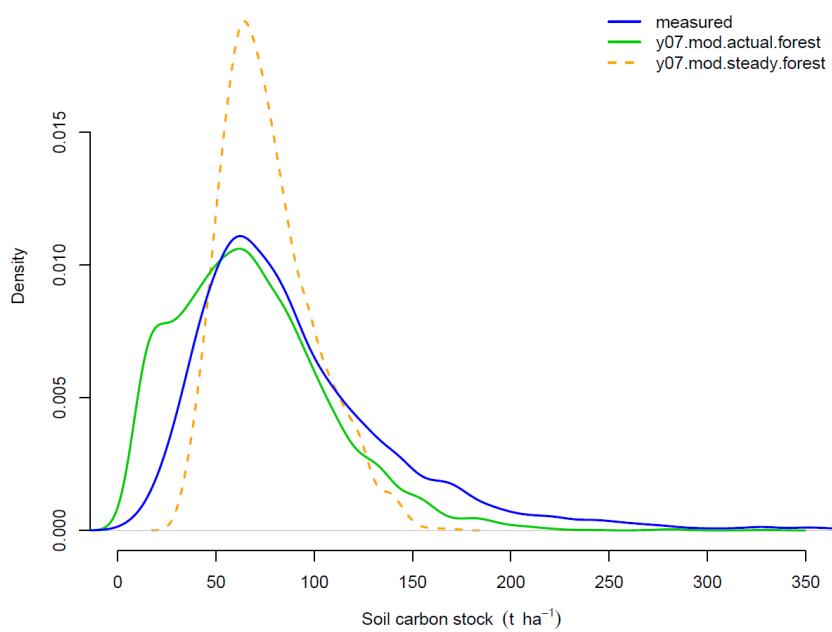


Figure S2. Density function of soil carbon stock measurements (measured) and the simulated soil carbon by the soil carbon model Yasso07 run with observed state forest and litter inputs (y07.mod.observed.forest) and with the equilibrium forest and litter inputs (y07.mod.steady.forest).

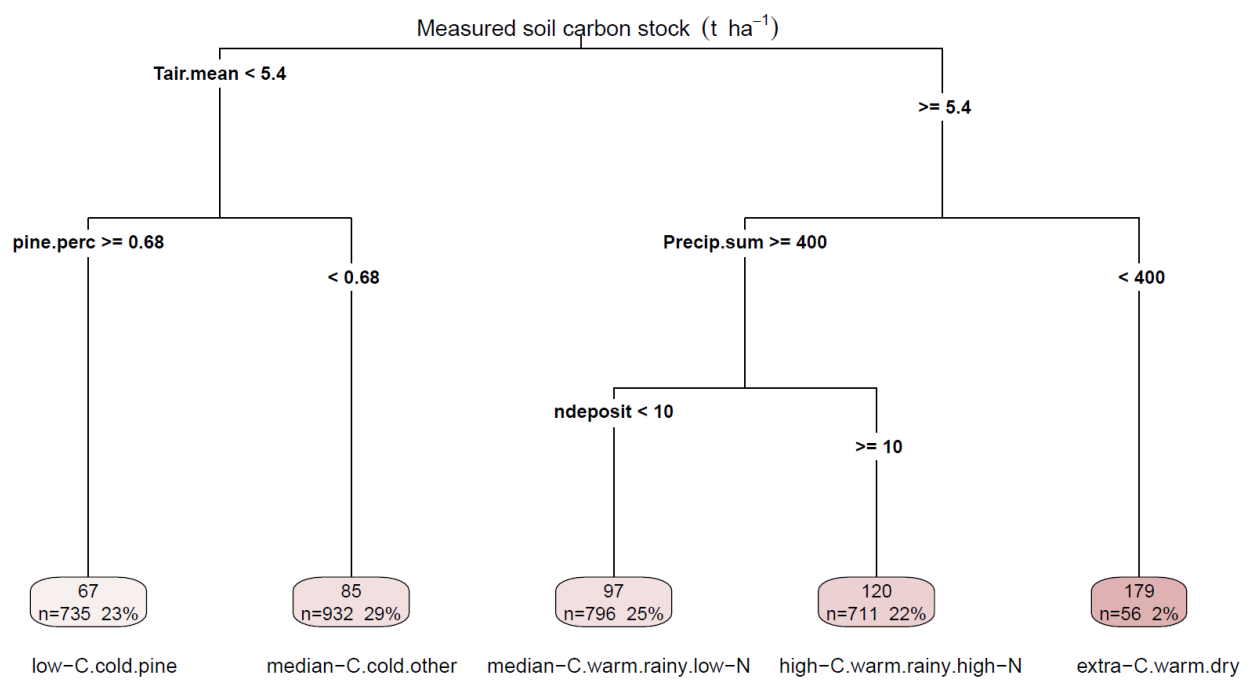


Figure S3. Classification/regression tree for the measured soil carbon stock ($t\text{C ha}^{-1}$) and site environmental characteristics excluding soil physicochemical properties; the annual air temperature (Tair.mean, $^{\circ}\text{C}$), the fraction of pine trees of the total canopy (pine.perc), the annual precipitation sum (Precip.sum, mm), and the nitrogen deposition (ndeposit, $\text{kgN ha}^{-1} \text{y}^{-1}$). The values in the leaves of the tree show for the distinct environmental conditions mean soil carbon stock ($t\text{C ha}^{-1}$), number and percentage of samples. The group acronyms are shown below the leaves of the regression tree.

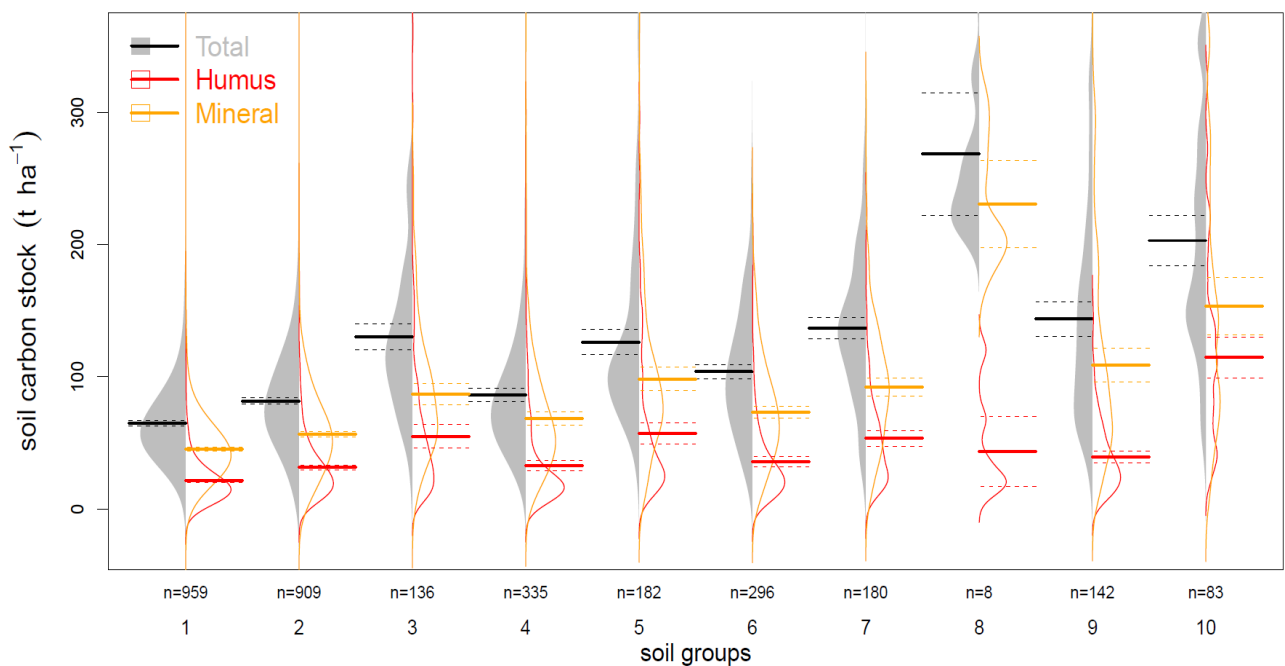


Figure S4. Density functions for 10 physicochemical groups of the soil carbon (SOC) stock ($tC\text{ ha}^{-1}$) Swedish forest soil inventory measurements for soil depth up to 1 m (total, grey fill) and for the soil humus horizon and mineral soil horizon. The thin lines are the density distributions. The thick lines are the group means and dashed lines are their confidence intervals. The n is number of samples. For description of group levels of SOC stocks, moisture, and fertility see Fig.2 and Table S1.

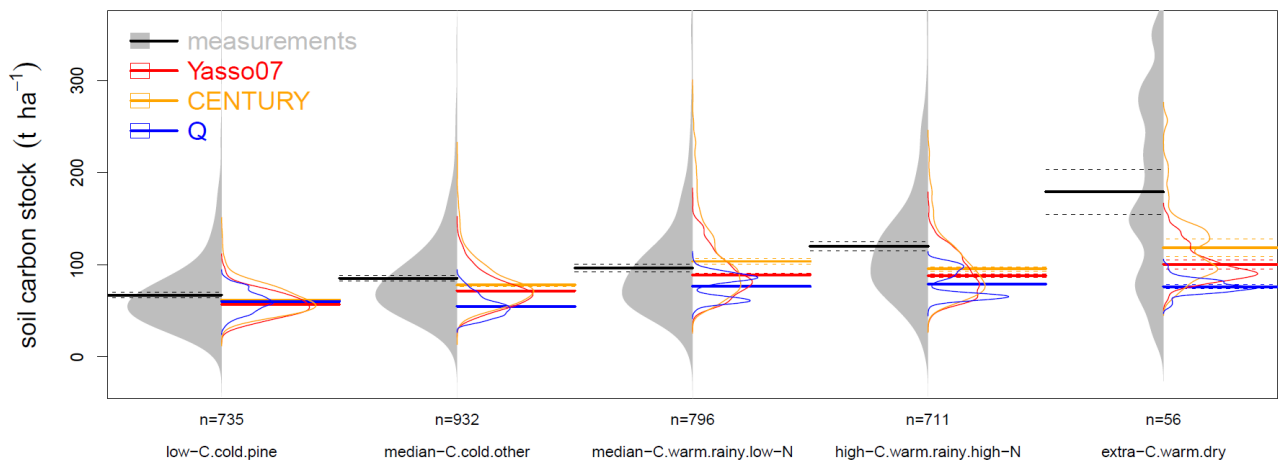


Figure S5. Bean plot of density functions for 5 groups of the soil carbon ($tC\ ha^{-1}$) measurements (C.measured, grey fill) and soil carbon estimates simulated by the soil carbon models Yasso07, CENTURY, and Q with the litter input derived from the equilibrium forest. The thin lines are the density distributions. The thick lines are the group means and dashed lines are their confidence intervals. The n is number of samples. For description of group acronyms based on levels of SOC stocks, temperature, percentage of pine in canopy, precipitation, and nitrogen deposition see Fig.S3 and Table S2.

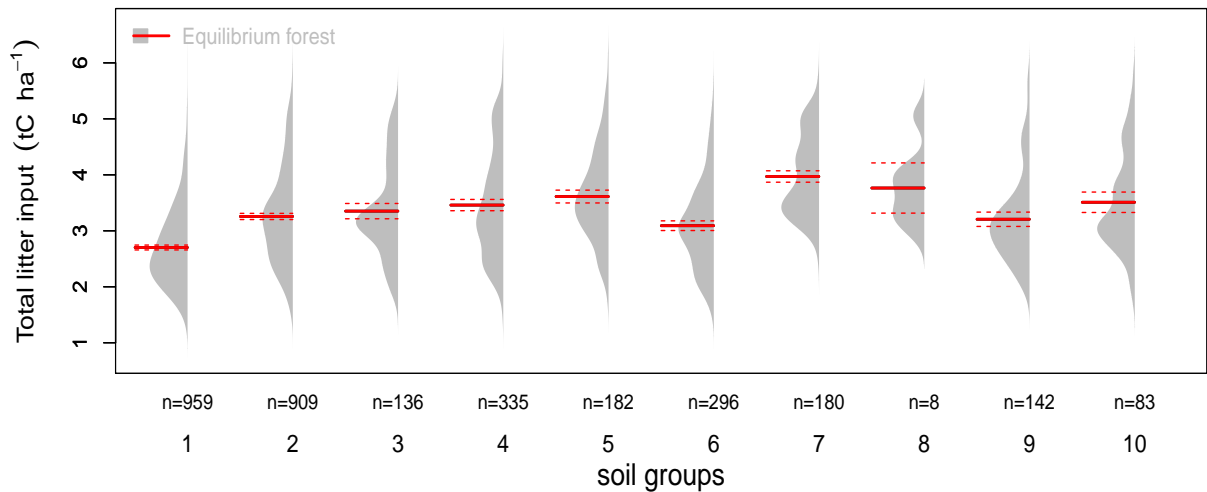


Figure S6. Density functions for 10 physicochemical groups of the total annual plant litter input ($tC\ ha^{-1}$) of equilibrium forest. The thick lines are the group means and dashed lines are their confidence intervals. The n is number of samples. For description of group levels of SOC stocks, moisture, and fertility see Fig.2 and Table S1.

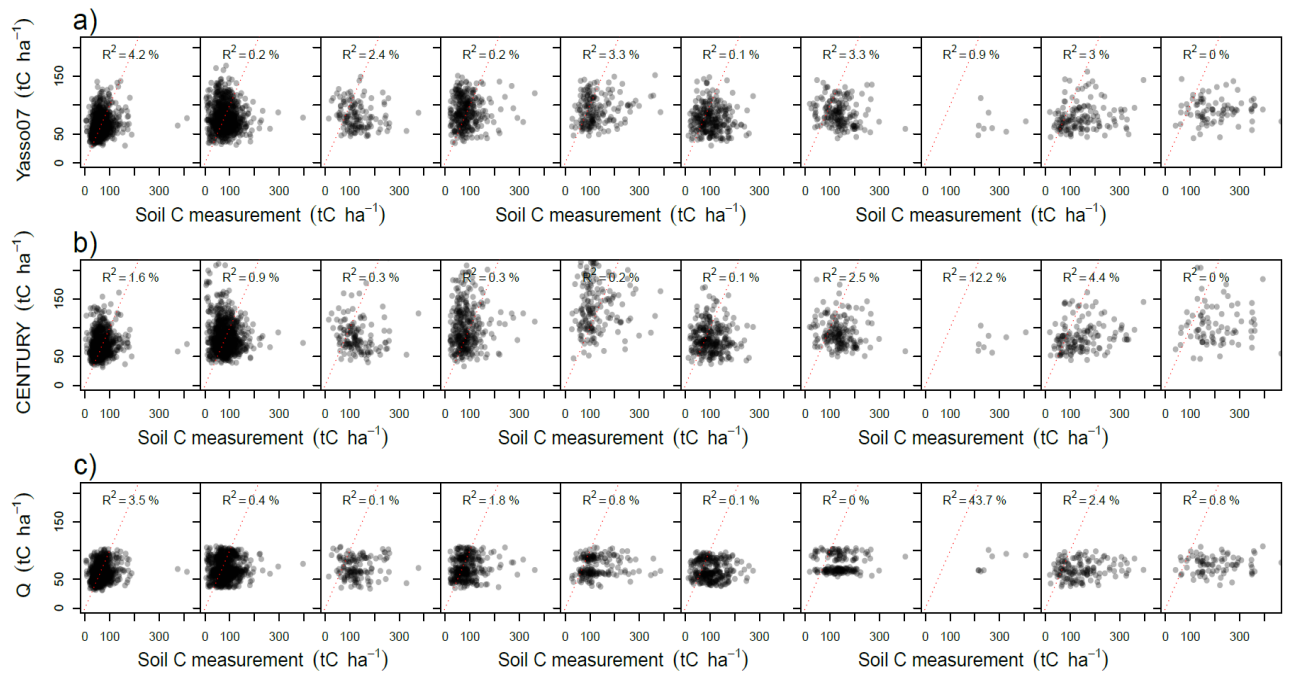


Figure S7. Scatter plots between model soil organic carbon stock ($tC\ ha^{-1}$) measurements and a) Yasso07 and CENTURY, b) Yasso07 and Q, and c) CENTURY and Q for 10 physicochemical groups of Fig.2.

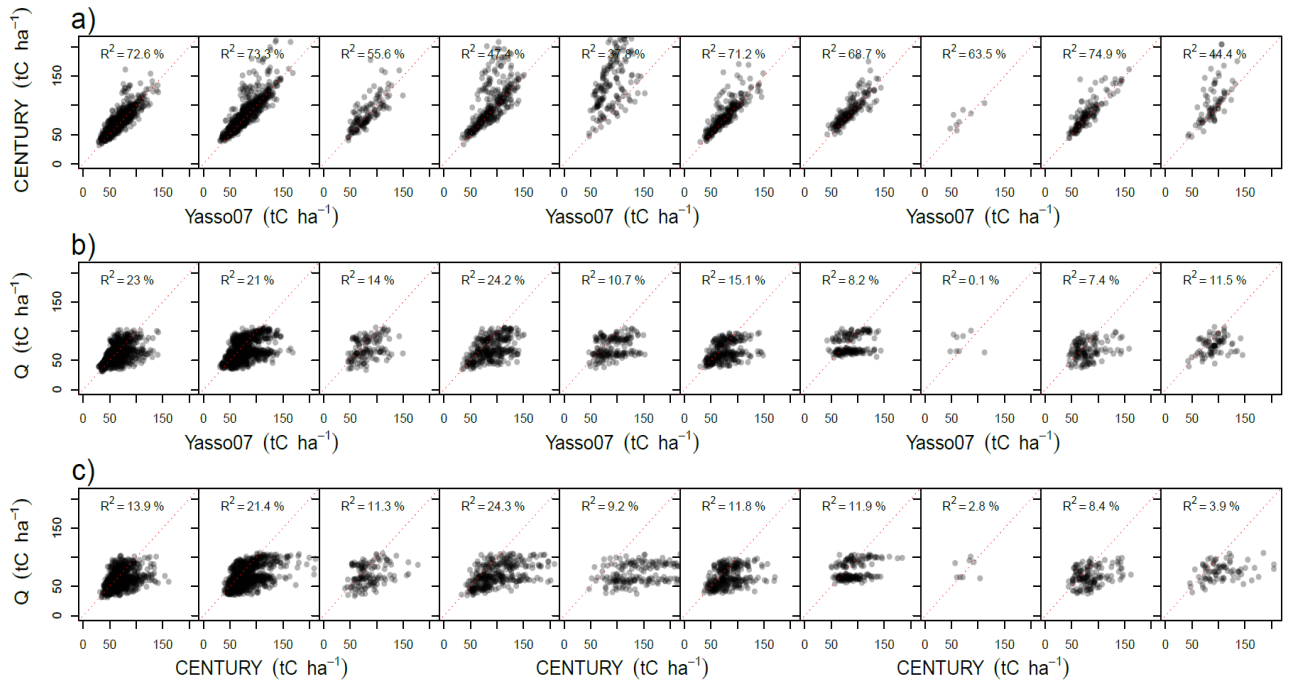


Figure S8. Scatter plots between model soil organic carbon estimates ($tC\ ha^{-1}$) of a) Yasso07 and CENTURY, b) Yasso07 and Q, and c) CENTURY and Q for 10 physicochemical groups of Fig.2.

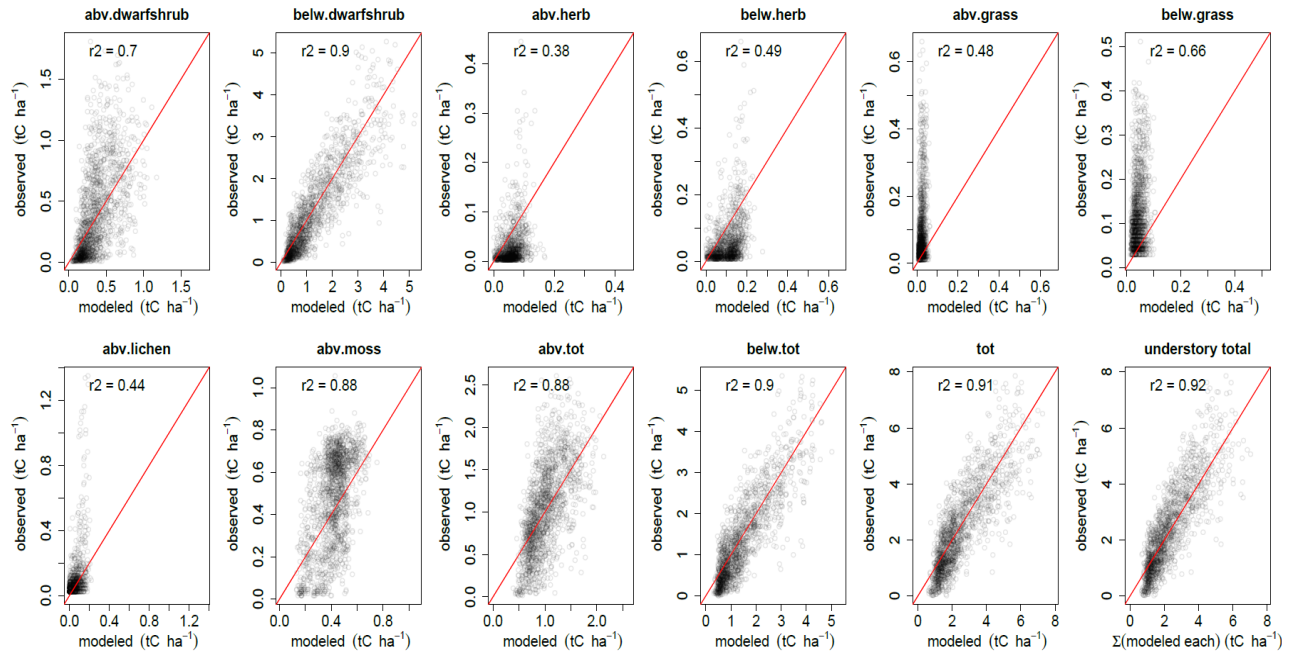


Figure S9. Scatter plots for the dry weight biomass ($tC\text{ ha}^{-1}$) of the functional types of understory vegetation for Swedish Forest Inventory plots in observed state being close to the estimated long-term mean conditions “equilibrium”. On the x-axis is the biomass modelled by the understory vegetation dry weight biomass ($tC\text{ ha}^{-1}$) models and on the y-axes is the observed coverage multiplied by the coverage/biomass conversion functions. The abbreviations “abv”, “belw”, and “tot” mean aboveground, belowground and total. The last panel for “understory total” shows high agreement between the sums of each modeled functional types and the sums of all functional types. The r^2 values represent the coefficient of determination indicating how close the modeled values fit the observed values.

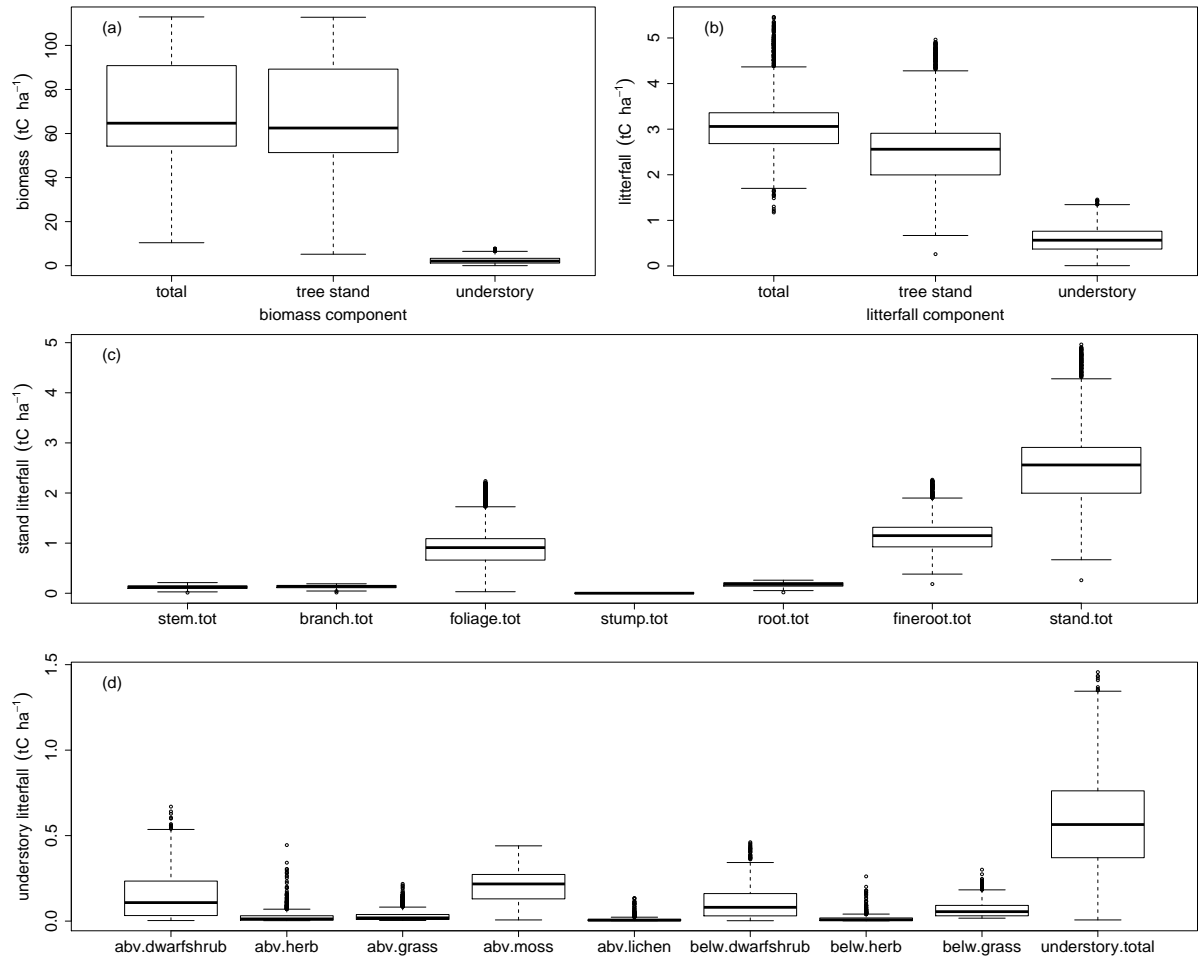


Figure S10. The tree stand and understory forest (a) biomass, (b) litterfall, (c) stand litterfall and (d) understory litterfall (all in tC ha^{-1}) for Swedish Forest Inventory plots with available understory coverage observations and in their observed state close to the estimated long-term mean conditions “equilibrium”.

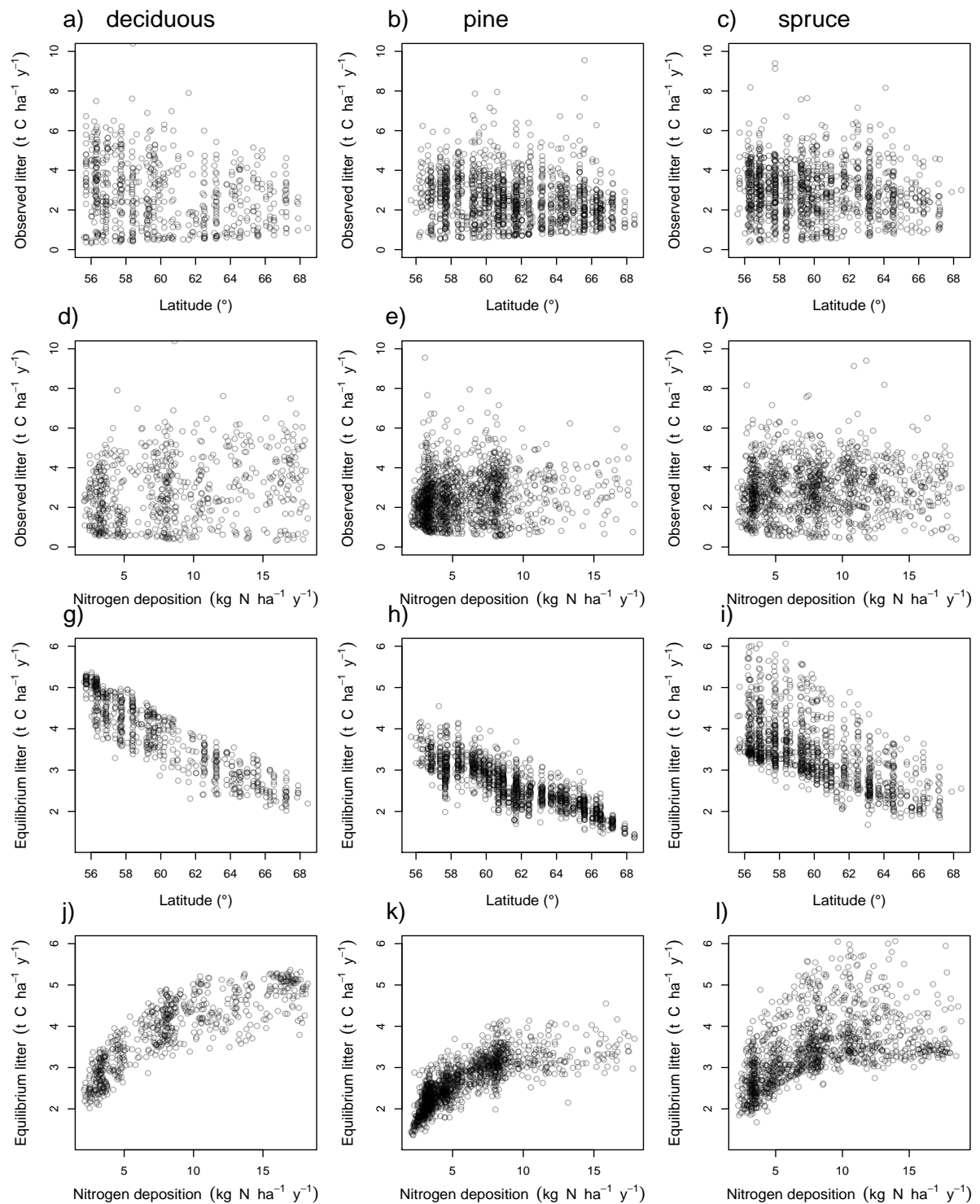


Figure S11. Scatterplots between Latitude ($^{\circ}$) and the observed state forest litterfall ($\text{tC ha}^{-1} \text{y}^{-1}$) a), b), c) or long-term mean “equilibrium” forest litterfall ($\text{tC ha}^{-1} \text{y}^{-1}$), g) h), i); and scatterplots between Nitrogen deposition ($\text{kgN ha}^{-1} \text{y}^{-1}$) and the observed forest litterfall ($\text{tC ha}^{-1} \text{y}^{-1}$) d), e), f) or long-term mean “equilibrium” forest litterfall ($\text{tC ha}^{-1} \text{y}^{-1}$) j), k), l) for deciduous species, Scots pine, and Norway spruce dominated stands.

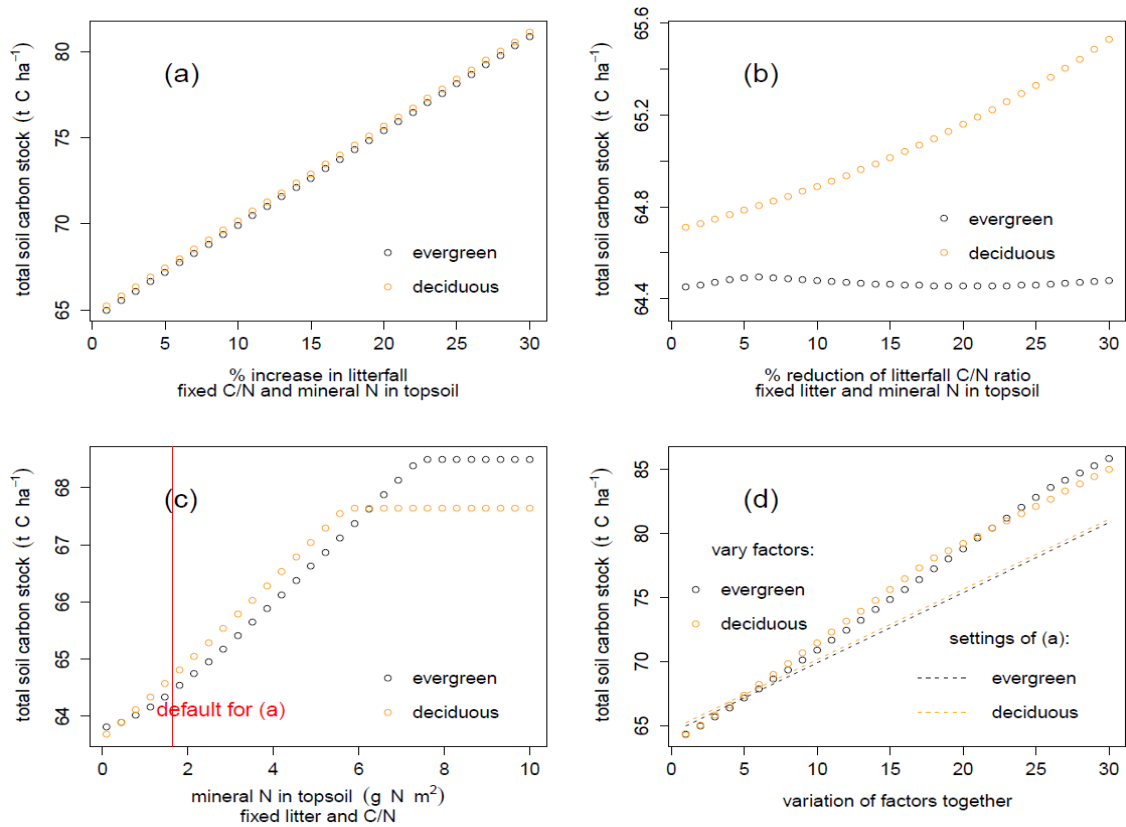


Figure S12. Sensitivity of simulated SOC stocks (tC ha^{-1}) of CENTURY model to variation in litterfall (a), C/N ratio of litterfall (b), topsoil mineral N (gN m^{-2}) (c), and to variation of factors together (d). SOC stocks of CENTURY are output of spin up simulation up to 1000 years.