# Mitteilungen aus dem Institut für Pflanzenökologie der Justus–Liebig–Universität Giessen

# Heft 5

# Nitrogen Carbon Interaction Model

# NCIM

Documentation: Model Version 3.00

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# The International Earth System Modelling Community

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# **Chapter 1**

# Introduction

This Version 3.00 of the Nitrogen Carbon Interaction Model (NCIM) is the first NCIM Version which has been released.

NCIM is a global generic model of the terrestrial carbon and nitrogen cycles. It was designed to investigate the coupling in nature of the carbon and nitrogen cycles of terrestrial ecosystems and the influences both cycles have on each other. This is important because carbon and nitrogen compounds are major atmospheric greenhouse gases and the cycles of these two elements, besides others, are strongly affected by human activities. Carbon and nitrogen pools interact considerably with the processes of carbon and nitrogen turnover in the ecosystems und with the climate system. NCIM allows the investigation of these interactions, their importance on the global or regional scale, and the role of natural and anthropogenic influences, on time scales from month to centuries.

NCIM is regionalized on a 0.5 degrees grid of geographical latitude and longitude (62 483 grid elements). The carbon and nitrogen cycles in the model are independent and closed. Mass–balance is maintained. During a regular model run, no amounts of carbon and/or nitrogen or C/N ratios are prescribed. Rather, the C/N ratios of all biospheric and soil compartments are calculated from the respective pools in each model time step.

The coupling of the carbon and nitrogen cycles has been performed in the way we prefer: Information on the system state (pools, state variables) is used for steering or regulating the processes (fluxes). For this purpose the coefficients of the process equations are calculated from the respective state variables by use of empirical functions (for the model structure see Figures 2.1 and 2.2 on pages 12 and 13).

These empirical functions, as well as the flux equations, were calibrated using data sets from our own experiments which were carried out in natural ecosystems of many climate zones of the world, in open top chambers, the phytotron of the Justus–Liebig–University and

Table 1.1: Some terms which are frequently used in this document together with the explanation of their meaning in this documentation. The variables which represent the terms in the model code may be (rarely) scalars or one– to multi–dimensional arrays.

term	synonym	meaning
pool	state variable	the compartments of the system containing matter (carbon,
		nitrogen, etc.) or energy
flux	process	transport of matter or energy from one pool to another.
		Fluxes are used to balance the pools
coefficient		variable in an equation which is updated once at the begin-
		ning or several times during a model time step. The cur-
		rent values of coefficients depend on the state of the sys-
		tem (pools) and/or on the current boundary conditions and
		driving forces (which are state variables in other systems).
parameter		variable in an equation which is set at the very beginning
		of the model run. Parameters depend on either initial con-
		ditions or boundary conditions which are assumed to be
		constant during the model run (i.e., vegetation type, soil
		type and texture, etc.).
driving force	control variable	(independent) variable in an equation which changes dur-
		ing time but is not changed by the model itself. Driving
		forces may either be read from files or provided by other
		coupled model(s). They are pools (state variables) in other
		than the currently modelled systems.
initial condition		Value assigned to a pool before the time evolution of the
		model starts.

the Giessen FACE Experiment (Jäger et al., 2003).

The model makes use of a global digital vegetation data set on the 0.5 degrees grid, derived from the Atlas for Biogeography (Atlas zur Biogeographie, Schmithüsen 1976). This vegetation data set includes 176 vegetation units throughout the world. The soil information on soil types and textures was digitizes from the Soil Map of the World (FAO–Unesco 1974 ff.).

NCIM's structure is modular. Each module is independent of the other modules and consists of a number of "tasks" which again are independent. The tasks work on the entire model grid, so that the state variables, coefficients, parameters, and driving forces are arrays of the respective grid. The dimensions of the commonly used arrays of variables in the model code

igrid	=	$1, \ldots, 62483$	(model grid)
i count	=	$1, \ldots, 149$	(country structure)
ibiome	=	$1, \ldots, 176$	(biomes, vegetation types)
ip	=	$1, \dots, 56$	(pools = state variables)

The documentation follows the Module and Task structure of the model code.

In this documentation, we use several terms which are explained in Table 1.1. The following conventions are used unless differently indicated:

- Variables in mathematical terms are written in *italics*.
- Variables related to the program code are written in typewriter.
- m<sup>2</sup> means square meter of ground surface area.
- g means gram of carbon or nitrogen.

The program code of NCIM has been written in standard FORTRAN 77. Only two exceptions were made: (1) we use the command include to include common block structures in the modules, and (2) we use the non standard character \_ (underline) in names of variables. Either have become standard with FORTRAN 90.

# Chapter 2

# **Model structure**

### 2.1 Structure of the carbon and nitrogen cycles

The carbon cycle in NCIM consists of carbon pools and carbon fluxes. Its nitrogen cycle includes nitrogen pools and nitrogen fluxes. Each pool is balaced by the fluxes which are connected to this pool.

The carbon and nitrogen cycles are calculated for a standard  $m^2$  of each grid element, with a monthly time–step. The dimension of the pools is  $[g \cdot m^{-2}]$ , and the dimension of the fluxes is  $[g \cdot m^{-2} \cdot month^{-1}]$ , therefore.

In addition, NCIM uses a number of pools which we call "dummy"–pools. They are used to collect fluxes for output purposes.

In NCIM the pools are two-dimensional arrays, of the notation p(igrid, ip), with igrid being the respective grid element, and ip the type of the carbon or nitrogen pool (see Tables 2.1 and 2.2 respective 2.3 and 2.4).

For the fluxes (processes) we use one dimensional arrays of the form flux (igrid) with i being a grid element, to balance the pools by differential equations. The flux terms are assigned to these arrays. The carbon and nitrogen fluxes are found in the Tables 2.5 and 2.6.

The differential equations are integrated in time using a forth–order Runge–Kutta method. The time–step of integration is two days.

The Figures 2.1 and 2.2 on pages 12 and 13 show the structure of the carbon and nitrogen cycles of NCIM. Some ecosystem compartments are represented in either the carbon and nitrogen cycles. These include the atmosphere (atmc and atmn), the phytomass (pc and pn), the litter (lc and ln), the soil organisms (micc and micn), and soil humus (humc and humn). Slowly decomposed organic residues (ligc) are assumed to originate from plant lignin compounds and be free of nitrogen. Nitrogen pools without analogous carbon pools include the

Table 2.1: Carbon pools

meaning	ipool			
phytomass carbon herbaceous above-ground	pcha			
phytomass carbon herbaceous below-ground	pchb			
phytomass carbon woody above-ground	pcwa			
phytomass carbon woody below-ground	pcwb			
litter carbon herbaceous above–ground	lcha			
litter carbon herbaceous below-ground	lchb			
litter carbon woody above–ground	lcwa			
litter carbon woody below-ground	lcwb			
carbon in lignin compounds	ligc			
carbon in humic substances				
carbon in the biomass of litter and soil organisms	micc			

Table 2.2: Carbon dummy pools (pools which are used to collect fluxes during a time period for output).

meaning	ipool				
net primary productivity carbon, herbaceous material					
net primary productivity carbon, woody material	nppcw				
litter production carbon from herbaceous material	lpch				
litter production carbon from woody material	lpcw				
litter depletion carbon, herbaceous litter	ldch				
litter depletion carbon, woody litter	ldcw				
lignin production carbon	ligpc				
lignin depletion carbon	ligdc				
humic substances production carbon	humpc				
humic substances depletion carbon	humdc				
carbon ( $CO_2$ ) segregated as excess carbon from soil organisms	misc				
carbon respiration by soil organisms	mirc				
carbon by mortality of soil organisms	mimc				
carbon removed by deforestation	dforc				

meaning	ipool
phytomass nitrogen herbaceous above-ground	pnha
phytomass nitrogen herbaceous below-ground	pnhb
phytomass nitrogen woody above-ground	pnwa
phytomass nitrogen woody below-ground	pnwb
litter nitrogen herbaceous above-ground	lnha
litter nitrogen herbaceous below-ground	lnhb
litter nitrogen woody above-ground	lnwa
litter nitrogen woody below-ground	lnwb
reserve nitrogen stored in plants, which is available for allocation to	resn
phytomass compartments	
nitrogen available (soluble) in litter and soil	avn
nitrogen in the biomass of litter and soil organisms	micn
nitrogen in humic substances	humn
nitrogen in soil buffer pool (immobilized ammonium)	puffn

Table 2.3: Nitrogen pools

mobile nitrogen reserves in plants (resn), the mineral "available" nitrogen in the soil (avn), and the nitrogen adsorbed on clay and humic substances in the soil (puffn).

In the carbon cycle, the main ecosystem turnover includes the atmospheric carbon pool as major object, and the fluxes connected with the atmosphere are the most important processes (net primary productivity nppc and depletion processes miscf, mimcf and mircf). In contrast, in the nitrogen cycle, the main turnover includes soil, plants, and soil organisms. The exchange with the atmospheric nitrogen pool is maintained by relatively minor fluxes like  $N_2$  fixation by symbionts or free living microorganisms (legnf, azonf, actnf) and denitrification processes producing  $N_2$  and  $N_2O$  from organic compounds (n2onf).

Although the carbon and nitrogen cycles are independant of each other and are massbalanced each, they interact strongly. The state of nitrogen pools is used for steering some fluxes in the carbon cycle: net primary productivity nppc, litter depletion ldc, depletion of slowly decomposible organic compounds ligdc, production of excess CO<sub>2</sub> by soil organisms miscf, and the production and depletion of humic substances humpcf and humdcf. On the other side, the state of carbon pools is similarly used for steering nitrogen fluxes: segregation of excess nitrogen by soil organisms misnf, uptake of available nitrogen by plants avunf, Table 2.4: Nitrogen dummy pools (pools which are used to collect fluxes during a time period for output).

meaning	ipool
nitrogen allocated by the net primary productivity	nppn
nitrogen transported with litter production	lpn
nitrogen transported to soil organisms by litter depletion	ldn
nitrogen excreted by soil organisms (humic substances nitrogen)	humpn
nitrogen transported to soil organisms by humus depletion	humdn
nitrogen transported to organisms from available N in litter and soil	mign
nitrogen mineralized by mortality of soil organisms	mimn
nitrogen segregated by soil organisms as excess N	misn
nitrogen uptake by plants from available nitrogen	avun
re-allocation of nitrogen from senescent leaves	raln
nitrogen fixed by symbionts in legume root nodules	legn
nitrogen fixed by Azospirillum symbionts in C4 grasses and cereals	azon
nitrogen fixed by actinomycetes	actn
$N_2O$ and $N_2$ gas eliminated from litter nitrogen pools by denitrification	n2on
processes correlated with gross mineralization	
nitrogen leached from soil to ground water and surface waters	lean
nitrogen deposition as $NO_x$ and $HNO_3$ from anthropogenic sources	depn

and the three fluxes of  $N_2\ fixation$  legnf, azonf, and actnf.

In the Figures 2.1 on page 12 and 2.2 on page 13 the steering influences on fluxes are indicated by black arrows marked with "N" and "C", respectively.

meaning	flux array name
net primary productivity carbon into herbaceous above-ground parts	nppha
net primary productivity carbon into herbaceous below-ground parts	npphb
net primary productivity carbon into woody above-ground parts	nppwa
net primary productivity carbon into woody below-ground parts	nppwb
litter production carbon of above-ground herbaceous material	lpcha
litter procuction carbon of below-ground herbaceous material	lpchb
litter production carbon of above-ground woody material	lpcwa
litter production carbon of below-ground woody material	lpcwb
carbon losses of above-ground woody material by deforestation	dforca
carbon losses of below-ground woody material by deforestation	dforcb
litter depletion carbon of above-ground herbaceous material	ldcha
litter depletion carbon of below-ground herbaceous material	ldchb
litter depletion carbon of above-ground woody material	ldcwa
litter depletion carbon of below-ground woody material	ldcwb
lignin production carbon	ligpc
lignin depletion carbon	ligdcf
humus production carbon	humpcf
humus depletion carbon	humdcf
carbon loss by respiration of soil organisms	mircf
carbon release by mortality of soil organisms	mimcf
carbon (CO <sub>2</sub> ) segregated as excess carbon from soil organisms	miscf

#### Table 2.5: Carbon fluxes of NCIM.

### Table 2.6: Nitrogen fluxes of NCIM.

meaning	flux array name
allocation of nitrogen due to net primary productivity into herbaceous	nppnha
above–ground parts	
allocation of nitrogen due to net primary productivity into herbaceous	nppnhb
below-ground parts	
allocation of nitrogen due to net primary productivity into woody	nppnwa
above–ground parts	
allocation of nitrogen due to net primary productivity into woody	nppnwb
below-ground parts	
re-allocation of nitrogen from senescent leaves	realnf
litter production nitrogen of herbaceous above-ground material	lpnha
litter production nitrogen of herbaceous below-ground material	lpnhb
litter production nitrogen of woody above-ground material	lpnwa
litter production nitrogen of woody below-ground material	lpnwb
nitrogen losses of above-ground woody material by deforestation	dforna
nitrogen losses of below-ground woody material by deforestation	dfornb
litter depletion nitrogen of above-ground herbaceous material	ldnha
litter depletion nitrogen of below-ground herbaceous material	ldnhb
litter depletion nitrogen of above-ground woody material	ldnwa
litter depletion nitrogen of below-ground woody material	ldnwb
humus depletion nitrogen	humdnf
excretion of nitrogen with humic substances by soil organisms	humpnf
uptake of available nitrogen by soil organisms	mignf
release of nitrogen by mortality of soil organisms	mimnf
segregation of excess nitrogen by soil organisms	misnf
uptake of available nitrogen by plant roots	avunf
fixation of available nitrogen on buffer in soil	avnpuf
return flux of nitrogen from buffer to available nitrogen	puavnf
N <sub>2</sub> fixation by symbionts in legume roots	legnf
$N_2$ fixation by Azospirrillum symbionts of $C_4$ grasses and cereals	azonf
$N_2$ fixation by actinomycetes in tree roots and sandy soils	actnf
nitrogen deposition due to $NO_x$ and $HNO_3$ immissions	depnf
nitrogen flux to the atmosphere by denitrification in soil	n2onf
leaching of soluble N compounds from soil to groundwater	leanf

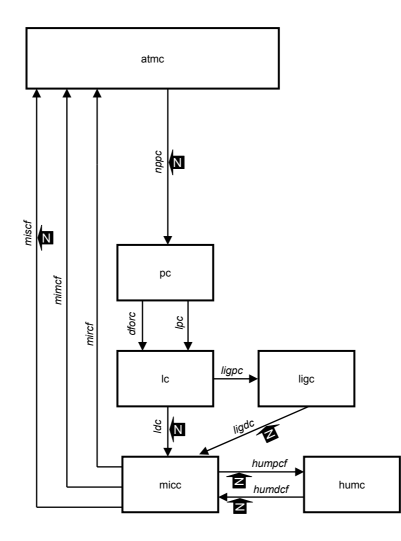


Figure 2.1: Structure of the carbon cycle of NCIM. Pools (state variables) are represented by squares, fluxes by arrows. The notation corresponds with Table 2.1 and Table 2.5. The pools pc, lc and the fluxes nppc, lpc, ldc exist four times each for herbaceous, woody, above ground and below ground materials. The black arrows indicated with "N" mark fluxes (processes) which depend on state variables of the nitrogen cycle.

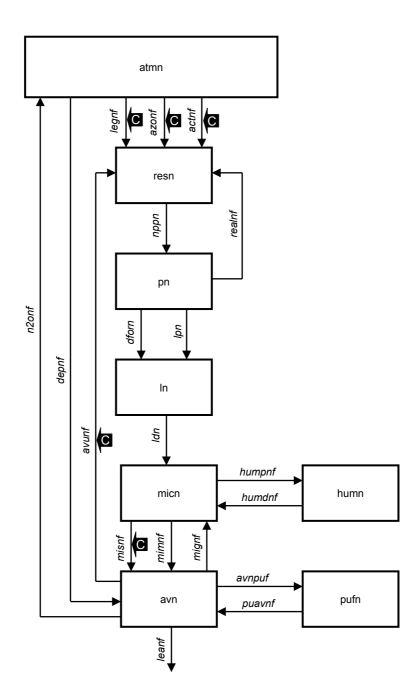


Figure 2.2: Structure of the nitrogen cycle of NCIM. Pools (state variables) are represented by squares, fluxes by arrows. The notation corresponds with Table 2.3 and Table 2.6. The pools pn, ln and the fluxes nppn, lpn, ldn exist four times each for herbaceous, woody, above ground and below ground materials. The black arrows indicated with "C" mark fluxes (processes) which depend on state variables of the carbon cycle.

### 2.2 Structure and function of the model code

The model code consists of modules, which are independent of each other (Figure 2.3 on page 15) and which interact with other modules using well–defined data interfaces. Each module includes one or several to many tasks, pieces of work of some independence, which contribute to the function of the module.

Data exchange between the modules is organized by common blocks. The only exception is the exchange of state variables and their estimates between timint and dgl respective ndgl. These three modules exchange data using a calling list.

The main module ncim 3.00 represents the main program. It organizes the time management of the model, i.e. the calling sequence of the other modules. The module timint includes the time-integration by a 4th-order Runge-Kutta method. The innermost time loop iday of main is subdivided in timint into a number of integration steps istep. Each integration step executes four calls to the two modules which include the mass-balance equations, dgl and ndgl.

The grid element loops (62 483 grid elements of a 0.5 degrees latitude and longitude grid) are placed in the tasks of the modules.

After declaration of the variables, main calls five modules to read driving forces and boundary conditions of the model as input data from files. Next some initial calculations are carried out to get data derived from the input data.

The model pools are filled with initial conditions in the initializing modules. Initial conditions are selected to be able to start a stable model run, i.e. a run without numerical problems. These values are not a steady–state solution of the model for any environmental conditions. The steady state–pool values for a given set of conditions, that means the initial values for a productive model run, are calculated by the model itself during a spin–up process, which we call "prerun". Two different preruns are carried out for about 5 000 model years each. In a first prerun, carbon/nitrogen (C/N) ratios of the relevant model fluxes, mainly the fluxes nppc and nppn, are kept constant. In prerun two, all C/N ratios are released and calculated by the model from the pools, until the model reaches steady–state. It is assumed that steady–state was achieved as soon as long–term trends of pool values are no longer observed (but seasonal changes are always present).

Productive model runs start with a steady-state set of pool values as initial conditions, which were derived by a prerun using the same boundary conditions and driving forces as in the first year of the produktive model run period. Driving forces and boundary conditions change during the model run. Gridded or scalar data sets must be available for the modelling

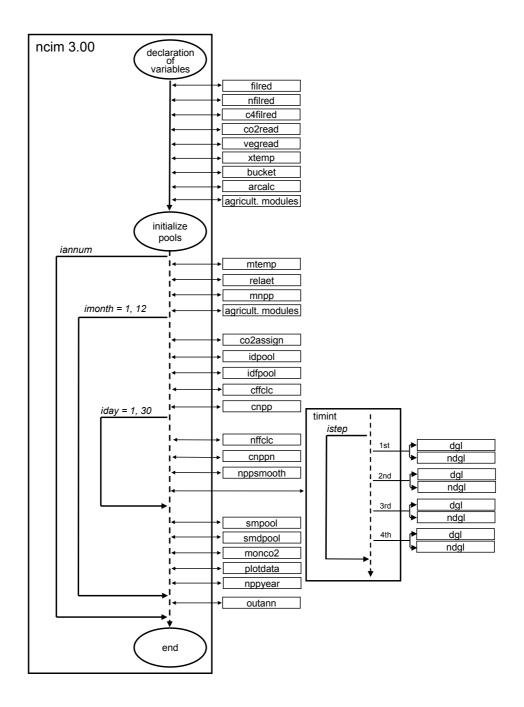


Figure 2.3: Structure of the NCIM program code. The code consists of 42 modules, including main (ncim 3.00), 6 modules in the initialize pools routine, and 7 agricultural modules. A detailed description of the modules in alphabetical order is given in chapter 3 beginning on page 17.

period (we use atmospheric  $CO_2$  concentrations from 1400 AD and climate anomalies and land–use changes from 1860 AD on).

In the annual loop iannum some long-term trends are updated first (land-use on per country level, soil, vegetation). The output of annual model results (summary non-gridded results) occurs at the end of the annual loop.

In the monthly loop imonth monthly changing driving forces are assigned, the dummy pools are set to zero (emptied to be filled again in the following month), and the coefficients of the flux-equations of the carbon cycle are calculated (cffclc, cnpp), thus creating the seasonality of the model. At the end of the monthly loop, upscaling of pools from square-meter to full grid element takes place and the output of gridded results takes place (plotdata).

The inner daily loop iday calls modules for the calculation of the coefficients of the nitrogen cycle. It also calls the integration module timint which evolves the model time by one day (Hoffstadt, Esser, 1997). One day is subdivided into several integration steps istep. The number of istep is selected so that numerical stability of the integration is maintained and integration biasses are minimized. The two modules dgl and ndgl include the mass balance equations for the carbon and nitrogen cycles and the related processes as shown in Figures 2.1 and 2.2 on pages 12 and 13.

# **Chapter 3**

# The NCIM modules

In this chapter, the modules of NCIM 3.00 are explained in alphabetical order. The role of each module in the model, its interface, and, if applicable, important algorithms and equations are explained. In the FORTRAN 77 source code of the model, the modules are subroutines (exept ncim, which is main) and are stored in seperate files each. The filenames are similar to the module names but with the extension \*.f. Detailed comments of the model code may be found in the source code files of the modules.

### 3.1 Agricultural modules

A number of modules are called if land–use is to be considered in a model run. These modules provide information on the areas used for agricultural purposes on country basis in the modelling period, on the grid elements which are used in the different years, and on the probability of the cultivation of C3 or C4 crops.

#### 3.1.1 Module agc4su

This module ranks all grid elements according to their probability to be useful for cultivation of C4 crops. A probability is calulated for each of the following C4 crops: maize, sugar cane, millet, and sorghum.

agc4s	u interface	
In:	monthly temperature	it(igrid,12)
	monthly precipitation	<pre>ipp(igrid,12)</pre>
	annual mean temperature	itann(igrid)
	annual precipitation	ippann(igrid)
	soil fertility	fsoil(igrid)
Out:	probability for cultivation of four C4 crops	prpc4(nc4,igrid)

Call frequency is once at the begin of a model run and once per year from AD 1860 on.

#### 3.1.2 Module agrc4pat

This subroutine determines the pattern of grid elements used for cultivation of C4 crops in a given model year.

agrc4	pat interface	
In:	required area for four C4 crops (AD 1860-1991)	agc4re(nc4,icount,1860:1991)
	standard map of the distribution of four C4 crops	usec4std(igrid)
	probability for cultivation of four C4 crops	prpc4(nc4,igrid)
	duration of land-use for each grid element (years)	useyr(igrid)
Out:	use-matrix of grid elements for the four C4 crops	useann(igrid)

The integer numbers assigned to the matrix useann mean:

0 natural vegetation, no use

1 C3 field crops

- 2 sugar cane
- 3 maize
- 4 millet
- 5 sorghum

Call frequency is once at the begin of a model run and once per year from AD 1860 on.

#### 3.1.3 Module agrfut

Module to provide data on the possible change of the use-matrix useann in the period 1991 to 2050 (land-use scenario). Based on a given ratio of agricultural net primary production to the population number of each country, it changes the use-matrix once a year so that the ratio is as close as possible to the given value.

Entire grid elements are transformed from useann = 0 to useann = 1, or vice versa. This is done if the required change of the country's agricultural production exceeds 60% of the productivity of the next grid element to be taken into or out of use.

The population data per country are calculated by use of polynomials. Those were calculated from the FAO population estimates for the period 1980 to 2050.

agrfut interface			
In:	model year (period AD 1991-2050)	iannum	
	area of each grid element	area(igrid)	
	10 years running mean of potential agricultural npp of a grid	nppyrm(igid)	
	element		
	four parameters for population polynomial per country	apop(icount,iparam)	
	relative agricultural productivity per country	rap(icount)	
	annual modification factor for $rap$ in the period 1991 to	fymult(icount)	
	2050		
	relative agricultural productivity per grid element	rapg(igrid)	
	probability of land-use of a given grid element in the given	agrprp(igrid)	
	year		
Out:	land-use-matrix of grid elements	useann(igrid)	
	duration of land-use for each grid element (years)	useyr(igrid)	

The call frequency is once per year for the period 1991 to 2050.

#### 3.1.4 Module agrpat

This module provides the actual pattern of grid elements used for agricultural purposes in a given model year in the matrix useann (igrid). The matrix useann includes values of zero for grid elements presently not in use, and 1 for those in use. If the module agrc4pat is called in advance of agrpat, the matrix useann may also include the values 2 through 5 which correspond with a variety of C4 field crops.

This module applies to model years up to 1990. Compare also module agrfut which provides land-use scenarios from 1991 on.

agrpat	interface	
In:	correction matrix to calculate the agriculturally used areas	korri(icount,1860:1990)
	in each country in the current model year	
	standard map of the distribution of agricultural crops in AD	usestd(igrid)
	1980	
	probability of each grid element to be agriculturally used in	agrprp(igrid)
	the current year	
	actual model year	iannum
	year of start of model run AD	ianns
	area of each grid element	area(igrid)
Out:	use-matrix of grid elements	useann(igrid)
	duration of land-use for each grid element (years)	useyr(igrid)

The module agrpat is called in the first model year and once every year in the period 1860 through 1990.

#### 3.1.5 Module agryie

This module provides the relative agricultural net primary productivity for each country and assignes the value rap(icount) to each grid element of the country rapg(igrid). The computation of rap depends on the current model year iannum.

- If *iannum* is equal to or lower than 1980, rap is set to the value of AD 1980.
- If iannum is greater than 1980 but equal to or lower than 1990, rap is determined by linear interpolation between the values of 1980 and 1990.
- For further scenarios, rap in 2050 of each country can be modified by a special factor considering individual developments fymult. Alternatively, rap in 2050 of all countries can be set to a fixed value scfut. The value rap for the current year is computed by linear interpolation between the value of AD 1990 and of AD 2050.

agryie interface			
In:	global value for relative agricultural productivity in AD	scfut	
	2050		
	current model year	iannum	
	relative agricultural productivity in AD 1980	rap80(icount)	
	relative agricultural productivity in AD 1990	rap90(icount)	
	factor to modify relative agricultural productivity in each	fymult(icount)	
	country		
Out:	relative agricultural productivity in the current model year	rap(icount)	
	for each country		
	relative agricultural productivity assigned to grid elements	rapg(igrid)	
	in current model year		

Module agryie ist called once per year in the period 1980 through 2050.

#### **3.1.6** Module agrsuit

In this module, the grid elements are ranked according to a "clearing probability" agrprp, which is defined as a product of several individual probabilities. At the present state, these individual probabilities include:

- probability based on the production potential: the higher the natural productivity, the higher the probability to be taken into use, the productivity is estimated by the Miami model (Lieth 1975).
- probability from soil quality: the higher the soil fertility, given by the array soil factor fsoil, the higher the probability to be used for agriculture,
- agricultural status of surrounding grid elements.

agrsui	agrsuit interface			
In:	standard map of the distribution of agricultural crops in AD 1980	usestd(igrid)		
	use-matrix of grid elements	useann(igrid)		
	latitude of grid element (lower left corner)	lat(igrid)		
	longitude of grid element (lower left corner)	lon(igrid)		
	annual mean temperature of grid element	itann(igrid)		
	annual precipitation of grid element	ippann(igrid)		
	soil fertility factor	fsoil(igrid)		
	actual model year	iannum		
Out:	probability of each grid element to be agriculturally used in	agrprp(igrid)		
	the current year			

The call frequency is once in the first model year and once a year from AD 1860 on.

#### 3.1.7 Module c4filrd

Module to read from files the information on the areas per country which are cultivated with C4 crops in the period 1961–1991, and a standard map of the distribution of grid elements dominated by C4 crops.

c4filrd interface			
In:	file dc4map	usec4std(igrid)	
	file dsugar	agc4re(sug,icount,iannum)	
	file dmaize	agc4re(mai,icount,iannum)	
	file dmillet	agc4re(mil,icount,iannum)	
	file dsorghum	agc4re(sor,icount,iannum)	
Out:	standard map of grid elements cultivated with C4 crops	usec4std(igrid)	
	required agricultural area for sugar cane	agc4re(sug,icount,iannum)	
	required agricultural area for maize	agc4re(mai,icount,iannum)	
	required agricultural area for millet	agc4re(mil,icount,iannum)	
	required agricultural area for sorghum	agc4re(sor,icount,iannum)	

Call frequency is once in the first model year and once per year in the period 1961 through 1991.

## 3.2 Module arcalc

This module computes the area of each grid element from the latitude of its lower left (southwestern) corner and the grid width. In addition it provides some information on the extremes of grid element sizes per country and global.

arcalc interface			
In:	latitude of lower left corner of grid element	lat(igrid)	
	number of the first grid element of each country	lcgrid(icount)	
	number of the last grid element of each country	mcgrid(icount)	
Out:	area [m <sup>2</sup> ] of each grid element	area(igrid)	
	area of the smallest grid element	aremin	
	area of the smallest grid element of each country	aremi(icount)	
	area of the largest grid element of each country	arema(icount)	
	mean area of the grid elements of each country	aremea(icount)	

arcalc is called once at the beginning of the model run.

## 3.3 Module bucket

bucket estimates monthly potential and actual evapotranspiration from a simple "bucket" soil moisture accounting scheme, using the Priestley/Taylor evapotranspiration formula. This module was adopted from the IIASA biome model (Prentice et al. 1992). In NCIM 3.00 actual evapotranspiration is used to influence seasonality, especially in the tropics and subtropics.

bucket interface			
In:	latitude of grid element (lower left corner)	lat(igrid)	
	longitude of grid element (lower left corner)	lon(igrid)	
	annual mean temperature of grid element	itann(igrid)	
	annual precipitation of grid element	ippann(igrid)	
	monthly temperatures of grid element	it(igrid,imonth)	
	monthly precipitation of grid element	<pre>ipp(igrid, imonth)</pre>	
	monthly mean cloudfreeness of grid element	<pre>icloud(igrid,imonth)</pre>	
Out:	monthly actual evapotranspiration of each grid element	aetm(igrid, imonth)	

bucket is called once per model year.

### **3.4 Module cffclc**

Module to provide the coefficients for the fluxes litter production and litter depletion, soil organic carbon production and soil organic carbon depletion and for the additional deforestation flux. cffclc determines the coefficients for each grid element in a given month.

These coefficients are required to calculate carbon and nitrogen fluxes in the modules dgl and ndgl. In cffclc the coefficients of the slower processes are updated monthly. The faster processes are updated daily in the module nffclc.

cffclc translates the driving forces on a grid element level into coefficients which occur in flux equations. This module together with nffclc is an important interface between environmental conditions an the biosphere, therefore.

Due to the relative importance of this module a more detailed description is given here.

While this module ist called the first time, the C/N ratio of litter production material is set to the standard value 32. Thus, it remained fixed during the first prerun (spin–up) procedure.

Next, cdfor is set to 0.06 for all grid elements under land-use. This results in a half-life time of the clearing progress of one year.

Next, the litter production coefficients for harbaceous material are set. It is assumed, that litter production only occurs in month with decreasing evapotranspiration. The coefficient's share in a given month is proportional to the amount of decrease of aetm in comparison to the total decrease in sequence aetdsu:

$$clpha(igrid) = \frac{aetdm(igrid, imonth)}{aetdsu(igrid)} \cdot \frac{herb(ibiome(igrid))}{0.59181 \cdot ageh(ibiome(igrid))^{0.79216}}$$
(3.1)

If no month with decreasing evapotranspiration occurs, litter production coefficients are evenly distributed over the months of a year:

$$clpha(igrid) = \frac{1}{12} \cdot \frac{herb(ibiome(igrid))}{0.59181 \cdot ageh(ibiome(igrid))^{0.79216}}$$
(3.2)

Litter production coefficients for woody material are evenly distributed throughout a year and are calculated analogously:

$$clpwa(igrid) = \frac{1}{12} \cdot \frac{1 - herb(ibiome(igrid))}{0.59181 \cdot agew(ibiome(igrid))^{0.79216}}$$
(3.3)

The coefficients for below-ground material are set equal to the analogous above-ground materials.

cffclc interface

In: monthly decrease of aetm aetdm (igrid, im	onth)
aetm in december of the last year aetmdc (igrid)	onen)
annual sum of negative monthly values of aetdm aetdsu (igrid)	
use-matrix of grid elements for agriculture useann (igrid)	
soil fertility factor fsoil (igrid)	
share factor of herbaceous productivity of each biome herb (ibiome (ig	
share factor of above-ground productivity of each biome abvgrd (ibiome (	2
mean age of herbaceous parts of each biome ageh (ibiome (ig	
mean age of woody parts of each biome agew (ibiome (ig	rid))
long-term mean temperature of warmest months mtwa (igrid)	
long-termmean temperature of coldest month mtco(igrid)	
vegetation type of each grid element ibiome (igrid)	
monthly temperatures of grid element it (igrid, imont	h)
monthly precipitation of grid element ipp (igrid, imon	th)
most important soil unit of the grid element soilat (igrid)	
current month imonth	
Out:coefficient litter production herbaceous above groundclpha(igrid)	
coefficient litter production herbaceous below ground clphb (igrid)	
coefficient litter production woody above ground clpwa (igrid)	
coefficient litter production woody below ground clpwb (igrid)	
coefficient litter depletion herbaceous above ground cldha (igrid)	
coefficient litter depletion herbaceous below ground cldhb (igrid)	
coefficient litter depletion woody above ground cldwa (igrid)	
coefficient litter depletion woody below ground cldwb (igrid)	
coefficient soil organic carbon production from herbaceous csocph (igrid)	
material	
coefficient soil organic carbon production from woody ma- csocpw(igrid)	
terial	
coefficient soil organic carbon depletion socd (igrid)	
coefficient of deforestation cdfor (igrid)	
counter for the number of month with litter production in ifall(igrid)	
sequence	
flag indicating month with litter production lfall (igrid)	

For the biomes of the temperate and boreal zones, i.e. cold-deciduous biomes, the monthly coefficient of litter production will be set to  $(2 \cdot \ln 2)$  in the first three month in which the monthly temperature drops (NOT rises) to or below the half of the mean temperature of the warmest month.

Next, C/N ratios for litter material are calculated from the respective carbon and nitrogen pools. This is done after the two preruns (spin–up) have been completed (the ratios for the preruns are set in module initn to standard values). The C/N ratios are only calculated if the amount of carbon in a litter pool is not less than  $10^{-5}$  g. If it is less, C/N ratios are set to standard values.

The depletion coefficients for litter and soil organic materials are calculated from climate variables and depend on the nitrogen content of the material. They depend on climate in a complex way:

$$cldh = \exp\left(p_1 + p_2 \cdot \ln(ipp) - p_3 \cdot ipp^{p_4}\right) + \exp(p_5) \cdot \tanh(p_6 \cdot ipp)$$
(3.4)

with the parameters

$$p_{1} = -1.96628 \cdot (it - 5) - 12.39641$$

$$p_{2} = 0.002236189 \cdot (it + 55)^{2}$$

$$p_{3} = 4.568434 \cdot \exp(-0.1041649 \cdot (it - 5))$$

$$p_{4} = 0.0001132567 \cdot (it + 55)^{2}$$

$$p_{5} = 0.07315304 \cdot (it - 5) - 3.51145$$

$$p_{6} = \exp\left(\frac{15000}{(it + 55)^{2}} - 6.5\right)$$

and the restrictions

If 
$$ipp = 0.0$$
 then  $cld = 0.0$   
If  $it < -30.0$  then  $cld = \exp(p_5) \cdot \tanh(p_6 \cdot ipp)$ 

Function (3.4) was developed using data for above ground herbaceous material (see Esser, 1986, for a list of the data used). It is modified for each compartment of litter according to

$$cldha = cldhb = cldh$$
  
 $cldwa = cldwb = 0.3 \cdot cldh$ 

The plot of function (3.4) is given in Figure 3.1.

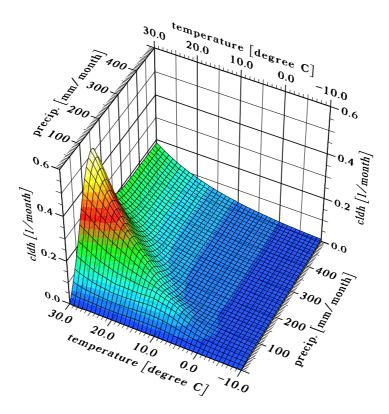


Figure 3.1: Plot of the litter depletion coefficient as a function of monthly precipitation and temperature (eq. 3.4).

Since function (3.4) requires a considerable amount of computing time, we have precalculated the depletion coefficients for a  $100 \times 100$  matrix of monthly temperature and precipitation intervals, in the range of temp -10 to 30 °C, precip 0 to 500 mm·month<sup>-1</sup>. Access to this matrix is then possible randomly by temp and precip classes.

Litter depletion coefficients as calculated from climate variables are then corrected according to the deviation of the actual nitrogen content of the material from a mean content:

$$cldha(igrid) = cldh \cdot \left(\frac{1.7}{1 + 1.4 \cdot \exp(0.1 \cdot (cnlha(igrid) - cnmean))} + 0.3\right)$$
(3.5)

The depletion coefficients for the materials woody above–ground, herbaceous below–ground, and woody below–ground are corrected analogously.

The depletion coefficients for humic substances and for lignin compounds are derived from the depletion coefficient for herbaceous material as calculated by equation (3.4). Since lignin compounds are nitrogen free, the nitrogen status of the soil organisms is considered for the depletion coefficient of lignin:

$$chumd(igrid) = cldh \cdot 0.0008$$
 (3.6)

$$cligd(igrid) = \frac{6}{cnmic(igrid)} \cdot cldh \cdot 0.05$$
 (3.7)

For certain soil types, *chumd* and *cligd* are modified:

$$chumd \begin{cases} \cdot 0.2 & \text{for Histosols} \\ \cdot 0.5 & \text{for gelic Gleysols} \end{cases}$$

 $cligd \begin{cases} \cdot 0.038 & \text{for Histosols} \\ \cdot 0.5 & \text{for gelic Gleysols} \end{cases}$ 

Finally, the lignin production coefficients are set. It was assumed therefore, that the lignin content of plant material will contribute to the formation of slowly decomposed humic acids in the soil. Thus, the lignin production coefficients are set to the mean share of lignin in herbaceous and woody plant material:

$$cligph(igrid) = 0.176 \tag{3.8}$$

$$cligpw(igrid) = 0.48 \tag{3.9}$$

The module cffclc is called once every month.

### 3.5 Module cnpp

This subroutine provides monthly net primary productivity for the array of grid elements. It considers climate on an annual basis, according to the Miami model (Lieth 1975). From the annual NPP value a part is allocated to the actual month according to the share of actual evapotranspiration of the current month, weighted by an exponent.

NPP from climate is then modified by correction factors for soil influence (fsoil), and the  $CO_2$  fertilization effect (co2fak). If the grid element is considered to be agriculturally used, the npp is further corrected by a factor (rapg) which represents the "Relative Agricultural Productivity" of the respective country. For further explanation see Esser (1995) or Esser (1994).

cnpp interface

In:	annual mean temperature of grid element	itann(igrid)
	annual precipitation of grid element	ippann(igrid)
	monthly temperatures of grid element	it(igrid,imonth)
	monthly actual evapotranspiration of each grid element	<pre>aetm(igrid,imonth)</pre>
	soil fertility factor of grid element	fsoil(igrid)
	relative agricultural productivity of grid element	rapg(igrid)
	share factor of herbaceous productivity of each biome	herb(ibiome(igrid))
	share factor for above-ground productivity of each biome	<pre>abvgrd(ibiome(igrid))</pre>
	potential natural vegetation of grid element	ibiome(igrid)
	CO <sub>2</sub> concentration in the atmosphere of a given month	co2
	$[\mu \mathbf{l} \cdot \mathbf{l}^{-1}]$	
Out:	monthly net primary productivity of herbaceous above-	nppha(igrid)
	ground material	
	monthly net primary productivity of herbaceous below-	npphb(igrid)
	ground material	
	monthly net primary productivity of woody above-ground	nppwa(igrid)
	material	
	monthly net primary productivity of woody below-ground	nppwb(igrid)
	material	

Net primary productivity is calculated as the minimum of two potential productivities which were derived from temperature and precipitation. The fraction of NPP in a given month is calculated using information on actual evapotranspiration. The factor 0.45 transforms dry matter into carbon.

$$nppt = \frac{3000}{1 + \exp(1.315 - 0.119 \cdot itann(igrid))}$$
(3.10)

$$npppp = 3000 \cdot (1 - \exp(-0.000664 \cdot ippann(igrid)))$$
 (3.11)

$$nppann = \min(nppt, npppp)$$
 (3.12)

$$npp(igrid) = nppann \cdot \frac{aetml(igrid, imonth)^{3}}{aetsu2(igrid)} \cdot fsoil(igrid) \cdot rapg(igrid) \cdot fco2 \cdot 0.45$$
(3.13)

In equation (3.13) aetml is equal to aetm, except in the case that aetm is less than 45 mm month<sup>-1</sup> in the given month and at least three month exist having aetm greater than 45 mm month<sup>-1</sup>. aetsu2 is the annual sum of  $aetml^3$ .

The factor fco2 in equation (3.13) represents the CO<sub>2</sub> fertilization effect. In NCIM 3.00 it is calculated from the actual atmospheric CO<sub>2</sub> concentration  $[\mu l \cdot l^{-1}]$  and the soil fertility according to equation (3.14).

$$co2fak = a \cdot [1 - \exp\{-r \cdot (co2 - 80)\}]$$
 (3.14)

where a = 1 + fsoil(igrid)/4 and  $r = -\ln(1 - 1/a)/240$ .

The  $CO_2$  fertilization effect is only applied to natural vegetation. If the grid element is agriculturally used, the factor will be set to 1 assuming supply limited by minerals rather than by the  $CO_2$  concentration.

Finally, monthly net primary productivity is distributed to the four compartments herbaceous and woody and above–ground and below–ground:

$$nppha(igrid) = npp(igrid) \cdot herb(ibiome(igrid)) \cdot abvgrd(ibiome(igrid)) (3.15)$$
  

$$npphb(igrid) = npp(igrid) \cdot herb(ibiome(igrid))$$
  

$$\cdot (1 - abvgrd(ibiome(igrid))) (3.16)$$

$$nppwa(igrid) = npp(igrid) \cdot (1 - herb(ibiome(igrid)))$$
$$\cdot abvgrd(ibiome(igrid))$$
(3.17)

$$nppwb(igrid) = npp(igrid) \cdot (1 - herb(ibiome(igrid)))$$
$$\cdot (1 - abvgrd(ibiome(igrid)))$$
(3.18)

Module cnpp is called once a month.

### 3.6 Module cnppn

This module saves the monthly NPP as provided by cnpp after the soil correction has been eliminated. As often as called, cnppn corrects the monthly saved NPP according to the actual C/N ratio in the plant material. If cnpp was called, cnppn updates the saved NPP values. In this module, the actual C/N ratios in the plant material are updated first.

$$cnacha(igrid) = \frac{pool(igrid, pcha)}{pool(igrid, pnha)}$$
(3.19)

$$cnachb(igrid) = \frac{pool(igrid, pchb)}{pool(igrid, pnhb)}$$
 (3.20)

$$cnacwa(igrid) = \frac{pool(igrid, pcwa)}{pool(igrid, pnwa)}$$
(3.21)

$$cnacwb(igrid) = \frac{pool(igrid, pcwb)}{pool(igrid, pnwb)}$$
(3.22)

In the next step, NPP is corrected according to the deviation of the actual C/N ratio in leaves from a biome-typical standard C/N ratio cnh(ibiome). The variable names ending with "o" refer to the old values of these variables.

$$x = \frac{cnh(ibiome(igrid))}{cnacha(igrid)}$$

$$fn = 1.7465 \cdot (1 - e^{-1.7 \cdot (x - 0.5)})$$

$$fnmean(igrid) = \frac{fnmean(igrid) \cdot (2 \cdot 15 - 1) + fn}{2 \cdot 15}$$

$$nppha(igrid) = fnmean(igrid) \cdot npphao(igrid)$$
(3.24)

$$npphb(igrid) = fnmean(igrid) \cdot npphbo(igrid)$$
 (3.25)

$$nppwa(igrid) = fnmean(igrid) \cdot nppwao(igrid)$$
 (3.26)

$$nppwb(igrid) = fnmean(igrid) \cdot nppwbo(igrid)$$
 (3.27)

### 3.7 Module co2assign

The model uses values for the  $CO_2$  concentration of the atmosphere from AD 1400 on. These values are global means of each year. The module colassign uses a simple interpolation procedure to calculate monthly means of the atmospheric  $CO_2$  concentration. The seasonal cycle is not considered. The monthly values help prevent high annual steps which would cause instabilities in the model. Prior to AD 1400 the monthly  $CO_2$  concentration is set to the annual value for AD 1400.

co2ass	co2assign interface		
In:	annual mean CO <sub>2</sub> concentration $[\mu l \cdot l^{-1}]$	co2ann(iannum)	
Out:	mean CO <sub>2</sub> concentration of current month $[\mu l \cdot l^{-1}]$	co2	

This module is called monthly.

### 3.8 Module co2read

Read annual data of the atmospheric CO<sub>2</sub> concentration co2ann (iannum)  $[\mu l \cdot l^{-1}]$  for the period AD 1400 including AD 2002 from file.

co2rea	d interface	
In:	file CO2.data	co2ann(iannum)
Out:	annual mean $CO_2$ concentration $[\mu l \cdot l^{-1}]$	co2ann(iannum)

Called once at the beginning of a model run.

# 3.9 Module dgl

Module dgl contains, together with its "sister" module ndgl (see page 38), the calculation of the fluxes (processes) and the mass-balance equations for carbon and nitrogen. Module dgl includes the "slow" fluxes of carbon, while ndgl includes the "fast" carbon fluxes and the nitrogen fluxes.

dgl interface		
In:	monthly net primary productivity of herbaceous above-	nppha(igrid)
	ground material	
	monthly net primary productivity of herbaceous below-	npphb(igrid)
	ground material	
	monthly net primary productivity of woody above-ground	nppwa(igrid)
	material	
	monthly net primary productivity of woody below-ground	nppwb(igrid)
	material	
	coefficient litter production herbaceous above ground	clpha(igrid)
	coefficient litter production herbaceous below ground	clphb(igrid)
	coefficient litter production woody above ground	clpwa(igrid)
	coefficient litter production woody below ground	clpwb(igrid)
	coefficient litter depletion herbaceous above ground	cldha(igrid)
	coefficient litter depletion herbaceous below ground	cldhb(igrid)
	coefficient litter depletion woody above ground	cldwa(igrid)
	coefficient litter depletion woody below ground	cldwb(igrid)
	coefficient lignin production from herbaceous material	cligph(igrid)
	coefficient lignin production from woody material	cligpw(igrid)
	coefficient of deforestation	cdfor(igrid)
In by list:	array of C and N pools	p(igrid,ip)
Out by list:	array of time-derivatives of pools	dp(igrid,ip)

### **3.9.1** Calculation of fluxes (processes)

To prevent redundancy, fluxes are calculated first and assigned to arrays of local variables.

Litter production carbon fluxes:

$$lpcha(igrid) = clpha(igrid) \cdot p(igrid, pcha)$$
 (3.28)

$$lpchb(igrid) = clphb(igrid) \cdot p(igrid, pchb)$$
 (3.29)

$$lpcwa(igrid) = clpwa(igrid) \cdot p(igrid, pcwa)$$
 (3.30)

$$lpcwb(igrid) = clpwb(igrid) \cdot p(igrid, pcwb)$$
 (3.31)

Deforestation carbon fluxes:

$$dforca(igrid) = cdfor(igrid) \cdot p(igrid, pcwa)$$
(3.32)

$$dforcb(igrid) = cdfor(igrid) \cdot p(igrid, pcwb)$$
 (3.33)

Litter depletion carbon fluxes:

$$ldcha(igrid) = cldha(igrid) \cdot p(igrid, lcha)$$
 (3.34)

$$ldchb(igrid) = cldhb(igrid) \cdot p(igrid, lchb)$$
 (3.35)

$$ldcwa(igrid) = cldwa(igrid) \cdot p(igrid, lcwa)$$
(3.36)

$$ldcwb(igrid) = cldwb(igrid) \cdot p(igrid, lcwb)$$
 (3.37)

Lignin depletion carbon flux:

$$ligdcf(igrid) = cligd(igrid) \cdot p(igrid, ligc)$$
(3.38)

#### **Time-derivatives of the pools (mass-balance equations)** 3.9.2

After calculating the carbon fluxes, the mass-balance equations follow. Mass-balance of carbon pools is maintained by balancing the input aud output fluxes:

Phytomass carbon:

$$dp(igrid, pcha) = nppha(igrid) - lpcha(igrid)$$
(3.39)

$$dp(igrid, pchb) = npphb(igrid) - lpchb(igrid)$$
(3.40)

$$dp(igrid, pcwa) = nppwa(igrid) - lpcwa(igrid) - dforca(igrid)$$
(3.41)

$$dp(igrid, pcwb) = nppwb(igrid) - lpcwb(igrid) - dforcb(igrid)$$
(3.42)

Litter carbon:

$$dp(igrid, lcha) = lpcha(igrid) - ldcha(igrid)$$
(loss to soil lignin pool:)
$$- cligph(igrid) \cdot lpcha(igrid)$$
(3.43)

(2, 25)

$$dp(igrid, lchb) = lpchb(igrid) - ldchb(igrid) - cligph(igrid) \cdot lpchb(igrid)$$
(3.44)  

$$dp(igrid, lcwa) = lpcwa(igrid) + df orca(igrid) - ldcwa(igrid) - cligpw(igrid) \cdot lpcwa(igrid)$$
(due to clearing:)  

$$- cligpw(igrid) \cdot df orca(igrid)$$
(3.45)  

$$dp(igrid, lcwb) = lpcwb(igrid) + df orcb(igrid) - ldcwb(igrid) - cligpw(igrid) \cdot lpcwb(igrid)$$
(due to clearing:)

$$- cligpw(igrid) \cdot dforcb(igrid) \tag{3.46}$$

Soil lignin pool (residuals of litter pools):

$$\begin{aligned} dp(igrid, ligc) &= cligph(igrid) \cdot lpcha(igrid) \\ &+ cligph(igrid) \cdot lpchb(igrid) \\ &+ cligpw(igrid) \cdot lpcwa(igrid) \\ &+ cligpw(igrid) \cdot lpcwb(igrid) \\ &+ cligpw(igrid) \cdot df orca(igrid) \\ &+ cligpw(igrid) \cdot df orcb(igrid) \end{aligned}$$

(lignin depletion, the carbon uptake by organisms while feeding on lignin:) -ligdcf(igrid) (3.47)

In the following, the equations for the collection of fluxes in "dummy" pools are found (for the list of dummy pools and their notation see Tables 2.2 and 2.4 on pages 7 and 9).

$$dp(igrid, nppch) = nppha(igrid) + npphb(igrid)$$
 (3.48)

$$dp(igrid, nppcw) = nppwa(igrid) + nppwb(igrid)$$
(3.49)

$$dp(igrid, lpch) = lpcha(igrid) + lpchb(igrid)$$
(3.50)

$$dp(igrid, lpcw) = lpcwa(igrid) + lpcwb(igrid)$$
(3.51)

$$dp(igrid, ldch) = ldcha(igrid) + ldchb(igrid)$$
 (3.52)

$$dp(igrid, ldcw) = ldcwa(igrid) + ldcwb(igrid)$$
(3.53)

$$dp(igrid, ligpc) = cligph(igrid) \cdot lpcha(igrid)$$
  
+  $cligph(igrid) \cdot lpchb(igrid)$ 

$$+ cligpw(igrid) \cdot lpcwa(igrid)$$
$$+ cligpw(igrid) \cdot lpcwb(igrid)$$
(3.54)

$$dp(igrid, ligdc) = ligdcf(igrid)$$
 (3.55)

$$dp(igrid, humdc) = humdcf(igrid)$$
 (3.56)

$$dp(igrid, dforc) = dforca(igrid) + dforcb(igrid)$$
 (3.57)

The module dgl is called istep times every model day (iday), see module timint and Figure 2.3 on page 15.

### **3.10** Module filred

This module is one of several which organize the input of data into the model. Those data include driving forces and boundary conditions. Files having names that end with the extension \*.05 contain gridded (0.5 degrees) data which are organized countrywise. For each country, a header record with the continent code, the country code, and the number of grid elements that belong to the country, starts the country data, followed by the data records for this country (each record includes data for one grid element).

Module filred is called once at the beginning of a model run.

### **3.11** Modules idpool and idfpool

In the model, we use so called "dummy" pools to add up fluxes of the model during a month for output (see Table 2.2 on page 7 and Table 2.4 on page 9). At the end of the month, those "dummy" pools contain the monthly accumulated sum of the material transported by the flux. The "dummy" pools are handeled like ordinary pools, i.e. they are balanced by the flux(es) they should collect, and integrated in the modules dgl or ndgl.

At the beginning of each month, "dummy" pools are set to zero, which means they are made empty. This is done in the two modules idpool and idfpool. These modules are called every month at the beginning of the monthly loop.

filred	interface	
In:	file korrmatrix_1990	korri(icount,year)
	file fyield8090.country	rap80(icount)
		rap90(icount)
	file popparam	apop(icount,iparam)
	file fymult.2050	fymult(icount)
	file coord.05	lat(igrid),lon(igrid)
	file temp_cramer.05	it(igrid,imonth)
		itann(igrid)
	file prec_cramer.05	<pre>ipp(igrid, imonth)</pre>
		ippann(igrid)
	file clou_cramer.05	<pre>icloud(igrid,imonth)</pre>
	file soilfactors.05.mod1805	soilasc,fsoil(igrid)
	file usestd_1980_circl0.05	usestd(igrid)
	file fossil.year	fossc(iannum)
	file cldmatrix	<pre>cld(nx,ny)</pre>
Out:	correction matrix to calculate the agriculturally used areas	korri(icount,iannum)
	in each country in the current model year	
	relative agricultural productivity in AD 1980	rap80(icount)
	relative agricultural productivity in AD 1990	rap90(icount)
	four parameters for population polynomial per country	apop(icount,iparam)
	annual modification factor for rap, 1991 to 2050	fymult(icount)
	latitude of the lower left corner of a grid element	lat(igrid)
	longitude of the lower left corner of a grid element	lon(igrid)
	monthly temperature	<pre>it(igrid, imonth)</pre>
	mean annual temperature	itann(igrid)
	monthly precipitation	<pre>ipp(igrid,imonth)</pre>
	annual precipitation	ippann(igrid)
	"cloudfreeness" of the month	<pre>icloud(igrid,imonth)</pre>
	soil type (coded)	soilat(igrid)
	soil fertility factor	fsoil(igrid)
	standard map of agricultural crops in 1980	usestd(igrid)
	emissions of carbon (CO <sub>2</sub> ) from fossil sources	fossc(iannum)
	depletion coefficients for a $100 \times 100$ matrix of monthly	cld(nx,ny)

temperature and precipitation values

## 3.12 Module mnpp

Module calculates 10–years running mean of the annual sum of net primary productivity of agriculturally used grid elements (NPP<sub>agri</sub>). If a grid element is coverd with natural vegetation, the NPP<sub>agri</sub> is calculated as if it were used.

mnpp	mnpp interface			
In:	annual sum of net primary productivity	nppyr(igrid)		
	relative agricultural productivity	rapg(igrid)		
	vegetation type	ibiome(igrid)		
Out:	10-year running mean of annual NPP	nppyrm(igrid)		

Module called once a year.

## 3.13 Module monco2

Assign the monthly  $CO_2$  concentration to an annual array at the end of the month.

monco2 interface		
In:	$CO_2$ concentration of the month $[\mu l \cdot l^{-1}]$	co2
Out:	array of monthly CO <sub>2</sub> concentrations	co2mon(imonth)

To be called monthly at the end of the month.

## 3.14 Module mtemp

This module assigns the the mean temperature of the warmest month of a given model year and grid element to the array mtwa (igrid), and the temperature of the coldest month to the array mtco(igrid).

Besides that, it calculates a 50-years running mean of mtwa and assigns it to the array mtwam(igrid).

mtemp interface			
In:	monthly temperature	it(igrid,imonth)	
Out:	mean temperature of warmest month	mtwa(igrid)	
	mean temperature of coldest month	mtco(igrid)	
	50-years running mean of mtwa	mtwam(igrid)	

Module is called once a year.

## **3.15 Module ncim 3.00**

This module represents the main program in FORTRAN code. Its main responsibility is to organize the call sequence and call frequencies of the modules of the model. For details and the model structure see Figure 2.3 on page 15.

Besides that ncim reads some data which are required for the configuration of an individual model run, from a configuration file ncimdata. This file is expected to be accessible at the beginning of a model run in the directory in which the model is started. The variables read and their possible values are as follows:

variable	meaning	values
ipresw	prerun switch	0 (no prerun performed)
		1 (prerun, pools will be initialized)
		2 (prerun, initial pools read from file)
iprern	prerun duration (years)	$0, \ldots, 10000$
iannb	starting year AD	$1860 \rightarrow$
iannend	ending year AD	$\rightarrow 2050$
fire	fire switch	0 (no vegetation fires considered)
		1 (vegetation fires considered)
co2	initial CO <sub>2</sub> concentration $[\mu l \cdot l^{-1}]$	$200, \dots, 600$
		preindustrial value 280.86
nannpr	nitrogen prerun 1 (years)	$0, \ldots, 10000$
nannzu	nitrogen prerun 2 (years)	$0, \ldots, 10000$
agri	agriculture switch	.true.or.false.

There are no other input data of this module. Output data include the model year iannum, the current month imonth, and the current day iday.

## 3.16 Module ndgl

Module ndgl contains, together with its "sister" module dgl (see page 32), the calculation of the fluxes (processes) and the mass-balance equations for nitrogen and carbon. Module ndgl includes the "fast" fluxes of carbon and the nitrogen fluxes, while dgl includes the "slow" carbon fluxes.

ndgl interface

Out:	array of time-derivatives of pools	dp(igrid, ip)
In:	array of C and N pools	p(igrid,ip)
by list:	current model year	iannum
	portion of recycled N before leaves are shed	<pre>rcych(ibiome(igrid)) ionnum</pre>
	standard C/N ratio of woody material	cnw(ibiome(igrid)
	standard C/N ratio of herbaceous material	<pre>cnh(ibiome(igrid))</pre>
	change of N deposition period 1860–1990	kordep(land,iannum)
	N deposition standard year	<pre>atmdep(igrid,imonth)</pre>
	share of NPP by C4 plants	c4npp(ibiome(igrid))
	share of NPP by legumes	<pre>legnpp(ibiome(igrid)</pre>
	respiration coefficient of soil organisms	crmic(igrid)
	leaching coefficient of available N to groundwater	cleach(igrid)
	coefficient of desorption of N from soil	cpuavn(igrid)
	coefficient of fixation of $NH_4^+$ in soil	cavnpu(igrid)
	coefficient of N (avn) uptake by plants	cassn(igrid)
	coefficient of mortality of soil organisms	cmmic(igrid)
	coefficient of N (avn)uptake by soil organisms	cgmic(igrid)
	material allocation coefficient for N to woody below–ground material	calwb(igrid)
	ground material allocation coefficient for N to woody above–ground	calwa(igrid)
	ground material allocation coefficient for N to herbaceous below-	calhb(igrid)
	allocation coefficient for N to herbaceous above-	calha(igrid)
	depletion coefficient humic substances	chumd(igrid)
	coefficient of deforestation	cdfor(igrid)
	rial coefficient lignin production from woody material	cligpw(igrid)
	coefficient lignin production from herbaceous mate-	cligph(igrid)
	coefficient litter depletion woody below ground	cldwb(igrid)
	coefficient litter depletion woody above ground	cldwa(igrid)
	coefficient litter depletion herbaceous below ground	cldhb(igrid)
	coefficient litter depletion herbaceous above ground	cldha(igrid)
	coefficient litter production woody below ground	clpwb(igrid)
	coefficient litter production woody above ground	clpwa(igrid)
	coefficient litter production herbaceous below ground	clphb(igrid)
	coefficient litter production herbaceous above ground	clpha(igrid)
	monthly net primary productivity of woody below- ground material	nppwb(igrid)
	monthly net primary productivity of woody above– ground material	nppwa(igrid)
	monthly net primary productivity of herbaceous below–ground material	npphb(igrid)
In:	monthly net primary productivity of herbaceous above–ground material	nppha(igrid)
nagi inte	Hatt	

)

In a first step, the C/N ratios for the different materials are calculated. Before the model run starts, i.e. during the two prerun procedures (spin–up), C/N ratios are set to standard values. After the start of the model run, C/N ratios for the 4 phytomass compartments, for the 4 litter compartment, for humic substances, and for the soil organisms are calculated by division of the respective carbon and nitrogen pools.

#### **3.16.1** Calculation of fluxes (processes)

The next task in this module ndgl is to calculate the fluxes (processes):

Depletion of humic substances:

$$humdcf(igrid) = chumd(igrid) \cdot p(igrid, humc)$$
(3.58)

Allocation of nitrogen from the N-reserve pool in plants to structural N compounds in the herbaceous above–ground phytomass:

$$nppnha(igrid) = calha(igrid) \\ \cdot \left(\frac{p(igrid, resn)}{resn0(igrid)}\right)^{zalloc} \\ \cdot 10^{min\left(1, \frac{\ln \frac{cnacha(igrid)}{cnh(ibiome(igrid))}}{(\ln 2.)^3}\right)}$$
(3.59)

The coefficient *calha* is provided by the module nffclc. The second term in equation (3.59) is the correction of allocation due to the availability of mobile nitrogen in the N–reserve pool of the plants. In the third term, the actual C/N ratio of the plant material (*cnacha*) is compared with the "normal" C/N ratio (*cnh*) of the material under consideration, thus detecting N deficiency or N abundance. This term provides an non–linear correction of allocation depending on the actual N–state of the tissues.

Similar equations are used for the allocation to the other three phytomass compartments *nppnhb*, *nppnwa*, and *nppnwb*.

During the preruns (spin–up procedure) the allocation of nitrogen is calculated differently, providing an input of carbon and nitrogen into the plant material with fixed C/N ratios:

$$nppnha(igrid) = \frac{nppha(igrid)}{cnh(ibiome(igrid))}$$
(3.60)

$$nppnhb(igrid) = \frac{npphb(igrid)}{cnh(ibiome(igrid))}$$
(3.61)

$$nppnwa(igrid) = \frac{nppwa(igrid)}{cnw(ibiome(igrid))}$$
(3.62)

$$nppnwb(igrid) = \frac{nppwb(igrid)}{cnw(ibiome(igrid))}$$
 (3.63)

$$lpnha(igrid) = clpha(igrid) \cdot p(igrid, pnha)$$
 (3.64)

$$lpnhb(igrid) = clphb(igrid) \cdot p(igrid, pnhb)$$
 (3.65)

$$lpnwa(igrid) = clpwa(igrid) \cdot p(igrid, pnwa)$$
 (3.66)

$$lpnwb(igrid) = clpwb(igrid) \cdot p(igrid, pnwb)$$
 (3.67)

Deforestation Nitrogen:

$$dforna(igrid) = cdfor(igrid) \cdot p(igrid, pnwa)$$
(3.68)

$$dfornb(igrid) = cdfor(igrid) \cdot p(igrid, pnwb)$$
(3.69)

Litter Depletion Nitrogen:

$$ldnha(igrid) = cldha(igrid) \cdot p(igrid, lnha)$$
 (3.70)

$$ldnhb(igrid) = cldhb(igrid) \cdot p(igrid, lnhb)$$
 (3.71)

$$ldnwa(igrid) = cldwa(igrid) \cdot p(igrid, lnwa)$$
 (3.72)

$$ldnwb(igrid) = cldwb(igrid) \cdot p(igrid, lnwb)$$
 (3.73)

Humus Depletion Nitrogen:

$$humdnf(igrid) = chumd(igrid) \cdot p(igrid, humn)$$
(3.74)

Excretion of humic substances by microorganisms, assumed to be 5% of total mineralized litter N:

$$humpnf(igrid) = (ldnha(igrid) + ldnhb(igrid) + ldnwa(igrid) + ldnwb(igrid)) \cdot 0.05$$
(3.75)

Uptake of available nitrogen (avn) by organisms:

$$mignf(igrid) = cgmic(igrid) \cdot max(0, p(igrid, avn))^{zmign}$$
(3.76)

Mortality of microorganisms:

$$mimnf(igrid) = cmmic(igrid) \cdot p(igrid, micn)$$
(3.77)

Segregation of excess nitrogen, if the C/N ratio is < 6. At the moment, the maximum coefficient (at cnmic = 0) is 3/4 per month, or 0.025 per day:

$$misnf(igrid) = \frac{3}{4} \cdot \frac{6 - min(6, cnmic(igrid))}{6} \cdot p(igrid, micn)$$
(3.78)

Uptake of available nitrogen avn by plants:

$$avunf(igrid) = cassn(igrid) \cdot p(igrid, pcha) \\ \cdot \frac{2}{1. + exp(1 \cdot (p(igrid, resn) - resn0(igrid)))} \\ \cdot p(igrid, avn)^{zavn}$$
(3.79)

Ammonium fixation on buffer in soil:

$$avnpuf(igrid) = cavnpu(igrid) \cdot p(igrid, avn)$$
 (3.80)

And return flux from buffer to avn:

Calculate actual adsorption-capacity in grid element:

$$cap(igrid) = (1 - sand(igrid)) \cdot fsoil(igrid) \cdot 105$$
 (from clay minerals)  
+  $p(igrid, humc) \cdot 3.5 \cdot 10^{-2}$  (from humic compounds) (3.81)

Correct the desorption coefficient by the calculated free capacity:

$$puavnf(igrid) = cpuavn(igrid) 
\cdot \frac{cap(igrid)}{cap(igrid) - p(igrid, puffn)} 
\cdot p(igrid, puffn)$$
(3.82)

Respiration of microorganisms:

$$mircf(igrid) = crmic(igrid) \cdot p(igrid, micc)$$
 (3.83)

Mortality of microorganisms:

$$mimcf(igrid) = cmmic(igrid) \cdot p(igrid, micc)$$
 (3.84)

Output of excess  $CO_2$  by microorganisms if they feed on carbon-rich compounds: Use a saturation function. The saturation function has half-saturation at cnmic = 9. The coefficient is scaled to 3/4 per month or 0.025 per day.

$$miscf(igrid) = \frac{max(6, cnmic(igrid)) - 6}{max(6, cnmic(igrid)) - 3}$$
$$\cdot \frac{3}{4} \cdot p(igrid, micc)$$
(3.85)

There are "open" fluxes, which connect the atmospheric  $N_2$  pool to the biospheric nitrogen pools. These "open" fluxes are the only way to add to the biosphere additional nitrogen in order to compensate for the additional carbon (CO<sub>2</sub> from fossil sources) which is incorporated into the biosphere. This makes these fluxes most important. Equations (3.86) through (3.88) provide *preliminary* descriptions of these fluxes.

Nitrogen fixation by symbionts in legume roots:

$$legnf(igrid) = legnpp(ibiome(igrid)) \cdot 0.3$$

$$\cdot \left(\frac{nppha(igrid) + npphb(igrid)}{cnh(ibiome(igrid))} + \frac{nppwa(igrid) + nppwb(igrid)}{cnw(ibiome(igrid))}\right)$$

$$\cdot \frac{resn0(igrid)}{p(igrid, resn)}$$
(correct for N-requirements) (3.86)

N fixation by Azospirillum symbionts in C4 grasses and cereals:

$$azonf(igrid) = c4npp(ibiome(igrid)) \cdot 0.03$$
  

$$\cdot \frac{nppha(igrid) + npphb(igrid)}{cnh(ibiome(igrid))}$$
  

$$\cdot \frac{resn0(igrid)}{p(igrid, resn)}$$
(3.87)

N fixation by Actinomycetes in tree roots on sandy soils. (Assumed to be proportional to the respiration flux of soil organisms and to the sand fraction in the soil):

$$actnf(igrid) = 0.01 \cdot mircf(igrid) \cdot sand(igrid)$$
 (3.88)

Anthropogenic nitrogen immissions (deposition of gaseous and particulate compounds) from emissions by traffic, heating devices, power plants, and others. For years prior to 1860 the deposition values of AD 1860 are assumed. Monthly values on per country level are calculated:

$$depnf(igrid) = atmdep(igrid, imonth) \cdot kordep(land(igrid), max(1860, iannum))$$
(3.89)

Leaching of available nitrogen (avn) into groundwater. The leaching coefficient *cleach* is provided by the module nffclc.

$$leanf(igrid) = cleach(igrid) \cdot p(igrid, avn)$$
(3.90)

Calculate the denitrification flux (N2ONF) of a grid element from the net mineralization. This flux includes the  $N_2O$  and  $N_2$  production from soil. We use an algorithm proposed by Nevison et al. (1996) which assumes that denitrification depends on net mineralization. It may be replaced in the near future.

$$n2onf(igrid) = \begin{cases} 9.5 \cdot 10^{-4} \cdot mimnf(igrid) & \text{for } mimnf \le 2.3 \\ 7.2 \cdot 10^{-5} \cdot mimnf(igrid)^2 \\ + 3.3 \cdot 10^{-4} \cdot mimnf(igrid) \\ + 1.0 \cdot 10^{-3} \end{cases} \quad \text{for } mimnf > 2.3 \qquad (3.91)$$

## **3.16.2** Time-derivatives of the pools (Mass-balance equations)

In the module ndgl the time-derivatives of the nitrogen pools and the "fast" carbon pools are calculated by balancing of the fluxes.

Humus Carbon:

$$dp(igrid, humc) = humpnf(igrid) \cdot cnhum(igrid)$$
(C flux from N flux)  
$$\cdot cnmic(igrid)/6$$
(regulating production)  
$$- humdcf(igrid)$$
(3.92)

Carbon content of organisms in litter and soil:

$$dp(igrid, micc) = \\ ldcha(igrid) + ldchb(igrid) + ldcwa(igrid) + ldcwb(igrid) \\ + ligdcf(igrid) \\ + humdcf(igrid) \\ - humpnf(igrid) \cdot cnhum(igrid) \qquad (excretion of humic substances) \\ - mircf(igrid) \\ - mincf(igrid) \qquad (output as CO2 due to mortality) \\ - miscf(igrid) \qquad (output of excess CO2) \\ (3.93)$$

#### Phytomass Nitrogen:

$$dp(igrid, pnha) = nppnha(igrid) - lpnha(igrid)$$
 (3.94)

$$dp(igrid, pnhb) = nppnhb(igrid) - lpnhb(igrid)$$
(3.95)

$$dp(igrid, pnwa) = nppnwa(igrid) - lpnwa(igrid) - dforna(igrid)$$
(3.96)

$$dp(igrid, pnwb) = nppnwb(igrid) - lpnwb(igrid) - dfornb(igrid)$$
(3.97)

#### Litter Nitrogen:

$$dp(igrid, lnha) = lpnha(igrid)$$

$$\cdot (1 - rcych(ibiome(igrid))) \qquad (recycle N before leaves shed)$$

$$- ldnha(igrid) \qquad (3.98)$$

$$dp(igrid, lnhb) = lpnhb(igrid) - ldnhb(igrid)$$
(3.99)

$$dp(igrid, lnwa) = lpnwa(igrid) + dforna(igrid) - ldnwa(igrid)$$
(3.100)

$$dp(igrid, lnwb) = lpnwb(igrid) + dfornb(igrid) - ldnwb(igrid)$$
(3.101)

Humus Nitrogen (excretion of humic substances minus uptake by organisms):

$$dp(igrid, humn) = humpnf(igrid) - humdnf(igrid)$$
(3.102)

Nitrogen in organisms in litter and soil. This pool is one of the most important "fast" nitrogen pools in litter and soil. It absorbs very fast (hours) available nitrogen (soluble nitrogen compounds) thus preventing or minimizing losses by leaching. Nitrogen in organisms is freely available for plants due to the short life–span of these organisms:

$$dp(igrid, micn) =$$

$$ldnha(igrid) + ldnhb(igrid) + ldnwa(igrid) + ldnwb(igrid)$$
(input from litter)

+ mignf(igrid)	(input from available N)	
+ humdnf(igrid)	(input from depletion of humic substances)	
-humpnf(igrid)	(loss from production of humic substances)	
-mimnf(igrid)	(loss (mineralization) by mortality)	
-misnf(igrid)	(segregation of excess N if C/N< 6)	
		(3.103)

Available nitrogen in soil. This pool includes the soluble nitrogen compounds in soil (and litter) which are produced by mineralization or added by immissions or fertilizers, i.e. mainly the  $NH_4^+ \leftrightarrow NO_2^- \leftrightarrow NO_3^-$  system.

$$\begin{aligned} dp(igrid, avn) &= \\ mimnf(igrid) & (\text{mineralization by mortality of organisms}) \\ &+ misnf(igrid) & (\text{segregation of excess N from organisms if C/N < 6)} \\ &+ depnf(igrid) & (\text{immission of NO}_{x}) \\ &- mignf(igrid) & (\text{uptake by microorganisms}) \\ &- avunf(igrid) & (\text{uptake by plants}) \\ &- leanf(igrid) & (\text{leaching into groundwater}) \\ &- n2onf(igrid) & (\text{denitrification from net mineralization}) \\ &- avnpuf(igrid) & (\text{fixation by adsorption on buffer}) \\ &+ puavnf(igrid) & (\text{desorption}) \end{aligned}$$

Balance of ammonium in the soil buffer system. This buffer system cuts peaks of available nitrogen, which would increase the leaching losses.

$$dp(igrid, puffn) = avnpuf(igrid) - puavnf(igrid)$$
(3.105)

Balance of reserve Nitrogen in plants. This pool serves as mobile nitrogen source for the allocation to the nitrogen in the phytomass compartments. Nitrogen uptake from soil, fixation of atmospheric  $N_2$  by symbionts and re–allocation from senescent plant parts provide nitrogen for this pool.

dp(igrid, resn) =	
avunf(igrid)	(available N uptake from soil)
$+ lpnha(igrid) \cdot rcych(ibiome(igrid))$	(re-allocation from senescent leaves)
+ legnf(igrid)	(fixation by symbionts in legume roots)
+ a zon f(i grid)	(fixation by Azospirillum symbionts)
+ actnf(igrid)	(fixation by Actinomecetes)
- nppnha(igrid) - nppnhb(igrid) - nppnwa(igrid) - nppnwb(igrid)	(allocation to phytomass compartments)
	(3.106)

The module ndgl is called once per integration step istep.

## 3.17 Module nffclc

In this module the coefficients of the "fast" fluxes are calculated from their driving forces.

The allocation coefficients for nitrogen to the compartments of phytomass are assumed to be a standard fraction of the respective share of net primary productivity to this compartment:

$$calha(igrid) = \frac{nppha(igrid)}{cnh(ibiome(igrid))}$$
(3.107)

$$calhb(igrid) = \frac{npphb(igrid)}{cnh(ibiome(igrid))}$$
(3.108)

$$calwa(igrid) = \frac{nppwa(igrid)}{cnw(ibiome(igrid))}$$
(3.109)

$$calwb(igrid) = \frac{nppwb(igrid)}{cnw(ibiome(igrid))}$$
(3.110)

The coefficient of growth of soil organisms is assumed to be  $15 \times$  higher than the depletion coefficient of herbaceous litter:

$$cgmic(igrid) = 15 \cdot cldha(igrid)$$
 (3.111)

In:	standard C/N ratio of herbaceous plant material	<pre>cnh(ibiome(igrid))</pre>
	standard C/N ratio of woody plant material	<pre>cnw(ibiome(igrid))</pre>
	monthly net primary productivity of herbaceous above–	nppha(igrid)
	ground material	
	monthly net primary productivity of herbaceous below-	npphb(igrid)
	ground material	
	monthly net primary productivity of woody above-ground	nppwa(igrid)
	material	
	monthly net primary productivity of woody below-ground	nppwb(igrid)
	material	
	coefficient litter depletion herbaceous above-ground	cldha(igrid)
	monthly actual evapotranspiration of each grid element	<pre>aetm(igrid,imonth)</pre>
	monthly temperatures of grid element	it(igrid,imonth)
	monthly precipitation of grid element	<pre>ipp(igrid, imonth)</pre>
	mean temperature of the warmest month	mtwam(igrid)
	sand fraction of the soil $(0, \ldots, 1)$	<pre>sand(igrid)</pre>
Out:	allocation coefficient for N to herbaceous above-ground	calha(igrid)
	material	
	allocation coefficient for N to herbaceous below-ground	calhb(igrid)
	material	
	allocation coefficient for N to woody above-ground mate-	calwa(igrid)
	rial	
	allocation coefficient for N to woody below-ground mate-	calwb(igrid)
	rial	
	coefficient of growth of soil organisms	cgmic(igrid)
	coefficient of mortality of soil organisms	cmmic(igrid)
	coefficient of respiration of soil organisms	crmic(igrid)
	coefficient of N uptake by plant roots	cassn(igrid)
	coefficient of leaching of available N to ground water	cleach(igrid)
	coefficient of transport (adsorption) between available ni-	cavnpu(igrid)
	trogen and the soil buffer	
	coefficient of desorption from soil buffer	cpuavn(igrid)

It is assumed that the life span of soil-organisms depends on cldha.

$$cmmic(igrid) = 0.1 \cdot cldha(igrid)$$
 (3.112)

The monthly respiration coefficient for soil organizms. This is the basic metabolic turnover of the soil organisms. It depends on temperature according to  $Q_{10} = 2$  with a base-temperature of 20°C. It is assumed that it is not influenced by humidity!

$$q_{10} = 2.0$$

$$x = \frac{\ln q_{10}}{10}$$

$$crmic(igrid) = 0.05 \cdot e^{x \cdot (it(igrid, imonth) - 20)}$$
(3.113)

Nitrogen uptake coefficient by plant roots (N assimilation rate) cassn. We assume a maximum rate of 0.013 [gN/gC/month]. It is assumed that soil moisture must not be considered since green leaves phytomass, which drives the flux, already depends on moisture. cassn is corrected for the temperature influence: The maximum rate is assumed to be valid for the "temperature of adaptation", i.e. the mean temperature of the warmest month (mtwam) and modified with  $Q_{10} = 2$ :

$$q_{10} = 2.0$$

$$x = \frac{\ln q_{10}}{10}$$

$$cassn(igrid) = 0.013 \cdot e^{x \cdot (it(igrid,imonth) - mtwam(igrid))}$$
(3.114)

Leaching rate of nitrogen into the ground water. The coefficient cleach depends on the water saturation of the soil and the composition (sand fraction) of the soil material. It is zero if the precipitation of the month is zero or less than the actual evapotranspiration or if the temperature is less than zero (frozen water):

$$cleach(igrid) = 0 \begin{cases} \text{for } ipp(igrid, imonth) = 0 \\ \text{or } ipp(igrid, imonth) < aetm(igrid, imonth) \\ \text{or } it(igrid, imonth) < 0 \\ \text{else:} \end{cases}$$

$$cleach(igrid) = (0.1 + sand(igrid)) \cdot \left(1 - \frac{aetm(igrid, imonth)}{ipp(igrid, imonth)}\right)$$
(3.115)

Transport coefficients between available nitrogen in soil (avn) and soil buffer (mainly for ammonium). These are preliminary tentative values (Müller, Giessen, 2004, pers. comm.) and at present subject to further investigation.

$$cavnpu(igrid) = 10 \tag{3.116}$$

$$cpuavn(igrid) = \frac{cavnpu(igrid)}{60}$$
 (3.117)

The module nffclc is called every day.

## 3.18 Module nfilrd

This module is one of several which organize the input of data into the model. This module reads data from files which are required to run the nitrogen cycle. Those data include driving forces and boundary conditions. Files having names that end with the extension \*.05 contain gridded (0.5 degrees) data which are organized countrywise. For each country, a header record with the continent code, the country code, and the number of grid elements that belong to the country, starts the country data, followed by the data records for this country (each record includes data for one grid element).

nfilrd interface			
In:	file texture.05	itext(igrid)	
	file atmdep.05	depmax(igrid)	
		<pre>scal(igrid,imonth)</pre>	
	file korratm.1860-1986.icount	kordep(icount,iannum)	
Out:	sand–fraction in the soil of a grid element $(0, \ldots, 1)$	sand(igrid)	
	nitrogen deposition for AD 1980 $[g \cdot m^{-2} \cdot month^{-1}]$	atmdep(igrid,imonth)	
	correction matrix to calculate the deposited amounts of N in	kordep(icount,iannum)	
	each country in the current model year		

## 3.19 Module nppsmooth

Module nppsmooth takes the net primary productivity (NPP) arrays (nppha, npphb, nppwa, nppwb) as soon as they are provided by modules cnpp and cnppn. It stores these original values and then overwrites the arrays with the smoothed NPP values. Smoothing means calculating a 12–day running mean of the NPP arrays in order to prevent steps of NPP values as a new month begins (NPP is updated from driving forces every month.

The Module nppsmooth is called daily.

nppsn	nooth interface	
In:	monthly net primary productivity of herbaceous above– ground material	nppha(igrid)
	monthly net primary productivity of herbaceous below-	npphb(igrid)
	ground material	
	monthly net primary productivity of woody above-ground	nppwa(igrid)
	material	
	monthly net primary productivity of woody below–ground material	nppwb(lgrid)
Out:	smoothed values of:	
	monthly net primary productivity of herbaceous above-	nppha(igrid)
	ground material	
	monthly net primary productivity of herbaceous below-	npphb(igrid)
	ground material	<i>.</i>
	monthly net primary productivity of woody above–ground	nppwa(igrid)
	material monthly net primary productivity of woody below–ground	nonub (igrid)
	material	πΡρωρ(τάττα)
	******	

## 3.20 Module nppyear

This module adds the dummy pools of net primary productivity for annual sums on a grid element basis (for output). This module is called every day.

nppye	nppyear interface		
In:	dummy pools of monthly net primary productivity	p(igrid,nppch)	
		p(igrid,nppcw)	
Out:	annual npp $[g \cdot m^{-2} \cdot yr^{-1}]$	nppyr(igrid)	

## 3.21 Module outann

This subroutine produces model output of global sums of nitrogen and carbon pools. Dummy pools are also added to produce global annual sums of fluxes. Since the output requirements depend on the investigated problem, we assume that this subroutine is re-written for any new research activity. This module imports fields (arrays) of the form poolsm(imonth) from

the modules smpool and smdpool. These fields contain the global sums of nitrogen or carbon for the 12 month of a year.

This module is called at the end of a model year. It may be called at the end of any month of any model year for which model results are required.

## **3.22 Module plotdata**

This module is used to write a complete model system state to an output file. The pools and dummy pools together with running means of certain pools are saved in this file. The values are on grid element level and have the dimension  $[g \cdot m^{-2} \cdot yr^{-1}]$ . The file may be used for several purposes, mainly (1) to restart the model if the computer run had to be stopped for any reason, (2) to produce thematic maps which show results on a 0.5 degrees resolution. The module plotdata may be called in regular intervals to save the model state for security. It may be called on request to produce output results at a certain time.

## 3.23 Module relaet

The module relact provides values that are necessary to distribute net primary productivity and the litter production over the months of a year.

relaet interface						
In:	monthly actual evapotranspiration of each grid element	aetm(igrid,imonth)				
Out:	monthly decrease of aetm	<pre>aetdm(igrid,imonth)</pre>				
	aetm in december last year	aetmdc(igrid)				
	annual sum of negative monthly values of aetdm	aetdsu(igrid)				
	annual sum of actual evapotranspiration	aetsum(igrid)				

Module relact is called once a year at the beginning of the year.

## **3.24** Modules smpool and smdpool

In these modules, the fields (arrays) of the form poolsm(imonth) are produced, which are used by the output module outann. The pools respective dummy-pools on grid element level of the form p(igrid, ip) (see Tables 2.1 and 2.2 respective 2.3 and 2.4) are of the dimension  $[g \cdot m^{-2}]$ . In these modules, the values for entire grid elements are calculated by

multiplication with the grid element area from the module arcalc. Then these values are globally added for global sums.

smpool and smdpool are called monthly.

## **3.25** module timint

This subroutine is a special module for time integration. It propagates the system state by SEC seconds. It uses the Runge–Kutta method (4th order). The time derivatives of the state variables (pools and dummy pools) are provided by the subroutines dgl and ndgl.

If we assume, that the basic time interval, one day, is subdivided into nstep integration steps, we can define several time steps for integration:

$$deltat = \frac{1}{nstep}$$

$$dt2 = \frac{deltat}{2}$$

$$dt3 = \frac{deltat}{3}$$

$$dt6 = \frac{deltat}{6}$$
(3.118)

The integration of the mass balance system over one time step by nstep integration steps istep is then done as follows (dp being the time derivative of pool, npool the total number of state variables):

```
do 42 istep = 1, nstep
            call dgl (pool,dp)
            call ndgl (pool,dp)
            do 421 ip = 1, npool
            do 421 igrid = lgrid, mgrid, jgrid
            ptmp(igrid,ip) = pool(igrid,ip) + dp(igrid,ip) * dt2
            psum(igrid,ip) = pool(igrid,ip) + dp(igrid,ip) * dt6
421 continue
            call dgl (ptmp,dp)
            call ndgl (ptmp,dp)
            do 422 ip = 1, npool
            do 422 igrid = lgrid, mgrid, jgrid
```

```
ptmp(igrid, ip) = pool(igrid, ip) + dp(igrid, ip) * dt2
          psum(igrid, ip) = psum(igrid, ip) + dp(igrid, ip) * dt3
422
        continue
        call dgl (ptmp, dp)
        call ndql (ptmp,dp)
        do 423 ip = 1, npool
        do 423 igrid = lgrid, mgrid, jgrid
          ptmp(iqrid,ip) = pool(iqrid,ip) + dp(iqrid,ip) * deltat
          psum(igrid, ip) = psum(igrid, ip) + dp(igrid, ip) * dt3
423
        continue
        call dgl (ptmp, dp)
        call ndgl (ptmp,dp)
        do 424 ip = 1, npool
        do 424 igrid = lgrid, mgrid, jgrid
          pool(igrid, ip) = psum(igrid, ip) + dp(igrid, ip) * dt6
424
        continue
```

53

```
42 continue
```

In this algorithm, the two-dimensional array pool (igrid, ip) always contains the total system state.

Module timint is called every model day.

## 3.26 Module vegread

This module organizes the input into the model of the biomes and biome related parameters. Files having names that end with the extension \*.05 contain gridded (0.5 degrees) data which are organized countrywise. For each country, a header record with the continent code, the country code, and the number of grid elements that belong to the country, starts the country data, followed by the data records for this country (each record includes data for one grid element). The information is based on tha "Atlas of Biogeography" by Schmithüsen (1976) which distinguishes 176 vegetation units of potential natural vegetation (biomes). We have aggregated these biomes into 31 biome–groups, for which vegetation parameters were established. These parameters are assigned to the 176 biomes.

vegrea	vegread interface				
In:	file schmithuesen.05	ibiome(igrid)			
	file vegets	ibiome, igroup(ibiome)			
	file groupparameters_c	grpno, gagew(grpno)			
		<pre>gageh(grpno), gherb(grpno)</pre>			
		gabvgr(grpno)			
	file groupparameters_n	grpno, gcnh(grpno)			
		grcych(grpno), gcnw(grpno)			
		glegnp(grpno), gc4npp(grpno)			
Out:	potential natural vegetation of grid element	ibiome(igrid)			
	standard C/N ratio of herbaceous plant material	<pre>cnh(ibiome(igrid))</pre>			
	standard C/N ratio of woody plant material	<pre>cnw(ibiome(igrid))</pre>			
	share factor of herbaceous productivity of each biome	herb(ibiome(igrid))			
	share factor of above-ground productivity of each biome	<pre>abvgrd(ibiome(igrid))</pre>			
	mean age of herbaceous parts of each biome	ageh(ibiome(igrid))			
	mean age of woody parts of each biome	agew(ibiome(igrid))			
	portion of recycled N before leaves are shed	<pre>rcych(ibiome(igrid))</pre>			
	share of NPP by legumes	<pre>legnpp(ibiome(igrid))</pre>			
	share of NPP by C4 plants	c4npp(ibiome(igrid))			

The module  ${\tt vegread}$  is called once at the beginning of a model run.

## 3.27 Module xtemp

Module to calculate growing degree days on a zero degree basis and growing degree days on

а	5	degree	basis.
---	---	--------	--------

xtemp	xtemp interface				
In:	monthly temperature	it(igrid,imonth)			
Out:	growing degree days on zero basis	gdd0(igrid)			
	growing degree days on 5 degree basis	gdd5(igrid)			

Called once at the beginning of a model run.

# **Chapter 4**

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