HYPE model description

The document describes how HYPE model the flow and transformation of water, nutrients and organic carbon in soil, lakes and rivers. This includes the effect of irrigation, point sources, floodplains and aquifers.

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Processes above ground

Temperature and precipitation

HYPE has many possibilities to adjust the precipitation and temperature given in the Pobs.txt and Tobs.txt files. Adjustment is done because the supplied data can be of different reference height or (known to be) wrong in some areas, or because data vary over the subbasin but the input is given as one value, and or to compensate for non modelled processes.

Temperature adjustments

Subbasin air temperatures \( T_i \) in the input file are normally assumed to be for the subbasin’s average elevation \( \text{elev} \). The same adjustments are made for maximum and minimum temperatures if available.

1. If data with other reference height is used it is possible to adjust for that. The subbasin temperature can be adjusted depending on subbasin elevation with general parameter \( tcelevadd \) \( (\circ / 100m) \). This adjustment assumes observations are located at sea level.

2. Alternatively it can be adjusted based on the difference between subbasin elevation and temperature observation elevation by general parameter \( tcobselev \) \( (\circ / 100m) \). The temperature observation elevation \( \text{elev}_{\text{obs}} \) then need to be supplied to the model.

3. It can be adjusted based on month with the monthly dependent parameter \( \text{monthlapse} \) \( (\circ / 100m) \). These three subbasin elevation adjustments should not be used together. Subbasin temperature can be adjusted equally over all subbasins within a region with the parameter region dependent parameter \( \text{tempcorr} \).

\[
T_i = T_i + \text{tempcorr} \cdot \frac{tcelevadd \times \text{elev}}{100} + \frac{tcobselev \times (\text{elev}_{\text{obs}} - \text{elev})}{100} + \frac{\text{monthlapse} \times \text{elev}}{100}
\]

The temperature can also be adjusted for each class depending on their deviation from the subbasin average elevation \( (\Delta h) \). The class-dependent temperature \( T \) is calculated using the parameter \( tcalt \). The temperature lapse rate often has a value of 0.6 \( (\circ / 100m) \).

\[
T = T_i - \frac{tcalt \times \Delta h}{100}
\]

Precipitation adjustments

Subbasin input precipitation \( P_i \) can be adjusted equally over all subbasins with the general parameter \( pcaddg \) or for some subbasins with the parameter region dependent parameter \( \text{preccorr} \). Additionally it is possible to adjust precipitation for undercatch with different parameters \( (pcurain, pcusnow) \) depending on if the precipitation falls as snow or rain.
Subbasin precipitation ($P_{gc}$) is for subbasin average elevation (elev), but can be adjusted for elevation variations within the subbasin. The precipitation of a class ($P$) is adjusted for classes where the class average elevation is greater than a threshold (general model parameter $pcelevth$). The adjustment is determined by a general parameter ($pcelevadd$) that is the correction per 100m. The class elevation adjustment can alternatively be determined from the basin standard deviation of elevation ($elev_{sd}$) and a parameter $pcelevstd$. The class height adjustment is limited by a general parameter $pcelevmax$. The precipitation of a class can additionally be adjusted with land-use dependent parameter $pcluse$, e.g. for interception evaporation.

\[
P_{gc} = P_i \times \left(1 + pcelevadd\right) \times \left(1 + precorr\right) \times \left(1 + \left(pcurain \times (1 - snfrac) + pcusnow \times snfrac\right)\right)
\]

\[
p_{c\text{\scriptsize{height}}} = \begin{cases} 
MIN \left(\frac{elev + \Delta h - pcelevth}{100} \times pcelevadd + \frac{elev_{sd} - pcelevmax}{100} \times pcelevstd, pcelevmax\right) & \text{elev} + \Delta h < pcelevth \\
0 & \text{else}
\end{cases}
\]

\[
P = P_{gc} \times \left(1 + p_{c\text{\scriptsize{height}}}\right) \times \left(1 - pcluse\right)
\]

Where $\Delta h$ is a class's elevation deviation from the subbasin average elevation and $snfrac$ is the average fraction of precipitation that falls as snow calculated from subbasin temperature ($T$) and temperature thresholds (see equation below), or from input.

**Rainfall and snowfall separation**

The rain/snow fraction of precipitation is calculated based on temperature or given as an input time series. Different temperatures can be used in the equation, i.e. basin average or class temperature. When the air temperature ($T$) is around the threshold temperature for mixed precipitation (land-use dependent parameter $ttmp$ plus general parameter $ttpd$) both rain and snow. The interval for mixed precipitation is given by the parameter $ttpi$. For temperature below threshold minus $ttpi$, the precipitation is assumed to be in solid form only and is added to the snowpack. If the air temperature is greater than the threshold temperature plus $ttpi$, the precipitation is assumed to be solely in liquid form. For intermediate temperatures, the precipitation is assumed to be a mixture of liquid and solid forms i.e. as both rain and snow. The proportion of precipitation ($P_{\text{rain}}$) that falls as rain depends linearly on the temperature.
Alternatively snowfall fraction ($sfrac$) may be read from input file and $a_{\text{rain}}$ calculated as:

$$a_{\text{rain}} = 1 - sfrac$$

The amount of rainfall and snowfall is calculated from the corrected precipitation.

$$rainfall = prec \times a_{\text{rain}}, \quad snowfall = prec \times (1 - a_{\text{rain}})$$

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Evaporation

Potential evaporation \((epot\) in mm) is calculated based on the temperature if it is not read in from file \((Xobs.txt)\). Alternative PET models exist, and is described below. When the air temperature \((T)\) is greater than the threshold temperature \(ttmp\) evaporation is assumed to occur. Snow melting, snow density and evaporation use the same threshold temperature. The basic potential evapotranspiration \((epot\) base) depends on the land use dependent rate parameter \(cevp\).

\[
\text{cseason} = 1 + cevpam \times \sin \left( \frac{2 \times \pi \times \text{dayno} - cevpph}{365} \right)
\]

\[
epot\ base = (cevp \times cseason) \times (T - ttmp)
\]

A seasonal factor \(cseason\) adjusts the potential evaporation rate \((cevp)\) e.g. making it higher in the spring when the air is often dry, and lower in autumn when the air is often more humid than in spring. The factor is sinusoidal with two parameters \(cevpam\) and \(cevpph\). It is not used if \(cevpam\) is zero. A \(cevpph\) around 45 days give a maximum correction in mid May \((\text{dayno}=45+91=136)\). The minimum correction will then be a half year later in September \((\text{dayno}=136+182=318)\). For an earlier maximum, reduce \(cevpph\).

The basic potential evapotranspiration may be adjusted with a regional correction factor \((cevpcorr)\) equally over the year depending on parameter region.

\[
epot = epot\ base \times (1 + cevpcorr)
\]

Evaporation from soil is assumed to occur from the two upper layers. The potential evaporation is assumed to decrease exponentially with depth (depending on the parameter \(epotdist\)). The potential evaporation is divided between the two layers \((epotfrac)\) with the distribution depending on the potential evaporation in the midpoint of each soil layer (figure 1). This is then used by approximating to a rectangle. Since soil layers differ between classes, the evaporation distribution do to.

\[
epot\ i = EXP \left( - epotdist \times soil\ layer\ depth \left( \frac{1}{2} \right) \right)
\]
Figure 1 The distribution of potential evaporation between the top two soil layers.

The actual evaporation from a soil layer (evap) is limited by the availability of water in the soil (soil) above the wilting point (wp, mm). Evaporation is at potential rate only if the water exceeds field capacity (fc, mm) or a (large) proportion (general parameter lp) of field capacity. In between these limits evaporation increase linearly.

\[
evapp = \begin{cases} 
0 & \text{soil-wp} < 0 \\
epot \times \text{epot frac} & \text{soil-wp} > lp \times fc \\
epot \times \text{epot frac} \times \frac{\text{soil-wp}}{lp \times fc} & \text{else}
\end{cases}
\]

\[evap = \text{MAX} \left( \evapp, \text{soil-wp} \right)\]
The actual evaporation may also depend on soil temperature (soiltemp). It is then reduced for temperatures above land use parameter \( t\text{trip} \) and depend on two other land use parameters (\( t\text{redA} \), \( t\text{redB} \)) as well (Figure 3).

The soil temperature evapotranspiration reduction is calculated as:

\[
\text{factor} = 1 - c \\
\text{evapp} = \text{evapp} \times \text{factor}
\]

The actual soil evaporation is set to zero for temperatures below the threshold temperature and for negative potential evaporation estimates (condensation).

A river with an area (i.e. is a class), flooded floodplains and lakes are assumed to evaporate at the potential rate, when the air temperature is above the threshold temperature (\( t\text{tmp} \)). Evaporation is limited by the water body’s volume.
Alternative potential evaporation models

HYPE give the option to exchange the default potential evaporation model for another model. Only the basic potential evaporation ($e_{pot base}$) differs between models. Models 0-2 only use air temperature forcing. Models 3-5 want shortwave radiation and minimum and maximum daily air temperature, although if lacking approximations are made (see section Input to alternative potential evaporation models below). In addition, model 5 wants relative humidity and wind speed if available. Note, regardless of potential evaporation model, the actual evaporation is limited to temperatures above parameter $ttmp$.

Model 0 (default)

As described above; evapotranspiration depends on the rate parameter $cevp$ and air temperature ($T$) above a threshold $ttmp$. If $epot$ is given in Xobs.txt those values are used.

$$e_{pot base} = (cevp \times cseason) \times (T - ttmp)$$

Model 1

Model 1 is the same as model 0, but it will not be using $epot$ from input data, even if it is present.

Model 2 - Modified Jensen-Haise/McGuiness

The modified Jensen-Haise/McGuiness model follow Oudin et al. (2005). The potential evaporation depends on extraterrestrial radiation ($radext$), latent heat of vaporization ($\lambda$) and temperature ($T$). Two general parameters ($jhtadd$ and $jhtscale$) are used and one land use dependent (crop coefficient $kc2$ or $kc$).

$$e_{pot base} = \frac{kc}{jhtscale} \times MAX \left( 0, \frac{radext}{\lambda} \times (T + jhtadd) \right)$$

Model 3 - Modified Hargreaves-Samani

The Hargreaves-Samani evaporation is modified to limit the “turbidity-factor”. The potential evaporation depends on extraterrestrial radiation ($radext$), latent heat of vaporization ($\lambda$), temperature ($T$) and turbidity ($turbidity$). One general parameter ($krs$) is used and one land use dependent (crop coefficient $kc3$ or $kc$).

$$e_{pot base} = MAX \left( 0, kc \times 0.0023 \times \frac{radext}{\lambda} \times \frac{turbidity}{krs} \times (T + 17.8) \right)$$
Model 4 - Priestly-Taylor

The Priestly-Taylor potential evaporation depends on net downward radiation \((\text{netrad})\), slope of saturated vapour pressure curve \((\text{dsatvap})\), latent heat of vaporization \((\lambda)\) and a psychrometric constant \((\gamma)\). One general parameter \((\text{alfapt})\) are used and one land use dependent (crop coefficient \(k_c4\) or \(k_c\)).

\[
e_{\text{pot base}} = \text{MAX} \left(0, k_c \times \text{alfapt} \times \frac{\text{dsatvap} \times \text{netrad}}{\lambda \times (\text{dsatvap} + \gamma)}\right)
\]

Model 5 - FAO Penman-Monteith

The FAO Penman-Monteith potential evaporation depends on net downward radiation \((\text{netrad})\), slope of saturated vapour pressure curve \((\text{dsatvap})\), saturated and actual vapour pressure \((\text{satvap} \text{ and } \text{actvap})\), temperature \((T)\), wind speed \((\text{wind})\) and a psychrometric constant \((\gamma)\). One land use dependent parameter (crop coefficient \(k_c5\) or \(k_c\)) is used.

\[
e_{\text{pot base}} = \text{MAX} \left(0, \frac{k_c \times 0.408 \times \text{dsatvap} \times \text{netrad} + \gamma \times \frac{900}{T + 273} \times \text{wind} \times \left(\frac{\text{satvap} - \text{actvap}}{\text{dsatvap} + \gamma \times \left(1 + 0.34 \times \text{wind}\right)}\right)}{\text{dsatvap} + \gamma \times \left(1 + 0.34 \times \text{wind}\right)}\right)
\]

Input to alternative potential evaporation models

Summary of alternative input to PET models, and link to file reference.

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### Actual vapour pressure

Actual vapour pressure \((actvap)\) is calculated following FAO recommended procedure and function/data priority. Depending on availability of minimum, mean and maximum relative humidity \((rh)\) and minimum, mean and maximum air temperature \((T_{\text{min}}, T, T_{\text{max}})\) equations with different combinations of saturated vapour pressure (calculated from temperature) times relative humidity is used. For example:

\[
actvap = \frac{\text{satvap}(T_{\text{max}}) \times rh_{\text{min}} + \text{satvap}(T_{\text{min}}) \times rh_{\text{max}}}{2}
\]

\[
actvap = \text{satvap}(T_{\text{min}}) \times rh_{\text{max}}
\]

\[
actvap = \text{satvap}(T) \times rh_{\text{mean}}
\]

\[
actvap = \text{satvap}(T_{\text{min}}) \times 1
\]

In case not enough data is available, minimum temperature is calculated from turbidity and a general parameter \((krs)\) and the last equation of those above is used.

\[
T_{\text{min}} = T - 0.5 \times \left( \frac{\text{turbidity}}{krs} \right)^2
\]

Eventually actual vapour pressure is limited by the calculated saturated vapour pressure.
**Air pressure**

Air pressure \((pa)\) is calculated as a function of elevation for the class \((elev)\).

\[
p_a = 101.3 \times \left( \frac{293 - 0.0065 \times elev}{293} \right)^{5.26}
\]

**Shortwave radiation**

Shortwave radiation \((swrad)\) is supplied as input forcing time series or otherwise calculated from extraterrestrial radiation.

\[swrad = radext \times turbidity\]

**Extraterrestrial radiation**

Extraterrestrial solar radiation \((radext)\) is estimated from day of year and latitude. The equations used comes from FAO. The day of the year are used to estimate distance to the sun and declination. The sunset hour angle are calculated from latitude, but with special care for high latitudes (polar night and midnight sun). These variables together with latitude and the solar constant eventually give the current extraterrestrial radiation for each subbasin.

**Latent heat of vaporization**

Latent heat of vaporization \((\lambda)\) is a function of temperature \((T)\).

\[
\lambda = 2.501 - 0.002361 \times T
\]

**Net downward radiation**

The net downward radiation \((netrad)\) is used explicitly for PET model 4 and 5 (Priestly-Taylor and FAO Penman-Monteith). The net radiation is calculated following FAO recommended procedure. It is calculated as net shortwave radiation minus net longwave radiation. Net shortwave radiation \((net_{short})\) is calculated from the shortwave radiation \((swrad)\) and the land use dependent albedo parameter \((alb)\). Net longwave radiation \((net_{long})\) is calculated using temperature \((T_{\text{min}}, T_{\text{max}}, T)\), actual vapour pressure \((actvap)\) and relative shortwave radiation \((relsh)\) if those are available, otherwise it is set to zero. The relative shortwave radiation is shortwave radiation in relation to clear sky shortwave radiation.

\[
net_{short} = swrad \times (1 - alb)
\]
\[
\text{net}_\text{long} = 4.903 \times 10^9 \times \frac{(T_{\text{max}} + 273.15)^4 + (T_{\text{min}} + 273.15)^4}{2} \times \left(0.34 - 0.14 \times \text{actvap}^{0.5}\right) \times \left(1.35 \times \text{relsh} - 0.35\right)
\]

\[
\text{relsh} = \frac{\text{turbidity}}{\text{clearturb}}
\]

**Psychrometric constant**

The psychrometric constant (\(\gamma\)) is a function of air pressure (\(p_a\)) and latent heat of vaporization (\(\lambda\)) following FAO.

\[
\gamma = \frac{0.001013 \times p_a}{0.622 \times \lambda}
\]

**Saturated vapour pressure**

Saturated vapour pressure (\(s\text{atvap}\), kPa) is calculated from temperature (\(\text{temp}\)) following FAO. If daily minimum and maximum air temperature is available \(s\text{atvap}\) is set to the average of the saturated vapour pressure calculated for each of those two temperatures, otherwise it is calculated from daily average air temperature.

\[
s\text{atvap} = 0.6108 \times \text{EXP}\left(17.27 \times \frac{\text{temp}}{\text{temp} + 237.3}\right)
\]

**Slope of saturated vapour pressure curve**

The slope of saturated vapour pressure temperature function (\(d\text{satvap}\)) is used explicitly for PET model 4 and 5 (Priestly-Taylor and FAO Penman-Monteith). The slope is calculated from daily mean air temperature (\(T\)).

\[
d\text{satvap} = 4098 \times 0.6108 \times \text{EXP}\left(\frac{17.27 \times T}{T + 237.3}\right) \times \frac{1}{(T + 237.3)^2}
\]

**Turbidity**

The turbidity factor is used explicitly for the PET model 3 - modified Hargreaves-Samani. If shortwave radiation (\(\text{swrad}\)) has been given as forcing data time serie, the turbidity factor is calculated as

\[
\text{turbidity} = \frac{\text{swrad}}{\text{radext}}
\]

but limited by a minimum turbidity value (0.25) and an estimated clearsky turbidity. The clearsky
turbidity \((\text{clearturb})\) is estimated by the Ångström formula (FAO):

\[
\text{clearturb} = 0.75 + \text{elev} \times 0.00002
\]

where \(\text{elev}\) is subbasin elevation in meter above sea level. If no shortwave radiation time series are given, but time series of daily minimum and maximum temperature are, the turbidity is calculated as in the “ordinary” Hargreaves-Samani turbidity function:

\[
\text{turbidity} = k_{rs} \times S Q R T \left( T_{\text{max}} - T_{\text{min}} \right)
\]

but still limited by the minimum turbidity value and the calculated clearsky turbidity.

**Wind speed**

Wind speed is used in PET model 5 - FAO Penman-Monteith. Wind speed \((\text{wind})\) may be given as a constant general parameter \((mwind)\) or as a forcing data time series. The time serie wind is given for each subbasin \((U_i)\). It is possible to adjust the time serie wind speed to different height than observations. If the general parameters \(z_{\text{wind}}, zwish, \text{roughness}, \text{and } z_{\text{pdh}}\) is set, wind speed is adjusted with the transformation factor \(\text{windtrans}\).

\[
\text{wind} = U_i \times \text{windtrans}
\]

\[
\text{windtrans} = \frac{\ln \left( z_{\text{wind}} \cdot z_{\text{pdh}} \right) - \ln \left( \text{roughness} \right)}{\ln \left( zwish \cdot z_{\text{pdh}} \right) - \ln \left( \text{roughness} \right)}
\]

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### Atmospheric deposition of nitrogen and phosphorus

#### Wet deposition

Atmospheric deposition in the form of wet deposition of IN and SP is added as a concentration of rainfall. You can specify a time series of the concentration in precipitation in Xobs.txt (implemented for IN, SP, and T1). Otherwise nitrogen wet deposition (as a concentration) is specified in GeoData.txt for each subbasin, while the wet deposition of phosphorus is specified by a general model parameters ($wetdepsp$).

If parameter ($aloadconst$) is set, the wet deposition load in considered constant and not dependent on precipitation corrections. Thus the concentration of precipitation is changed to keep the load constant, when precipitation correction is applied.

#### Atmospheric deposition to the soil

Nitrogen dry deposition is specified in GeoData.txt for each subbasin and different vegetation groups, while dry deposition of phosphorus is specified by a model parameter $drydepPP$ (land use dependent).

Dry deposition of nitrogen and phosphorus is added to the snow or, if there is no snow, directly to the ground. Phosphorus deposition is added to PartP-pool in the upper soil layer and nitrogen deposits to the IN dissolved in soil water if the parameter $ponatm$ is not set. A new concentration of IN in the soil water is then calculated. The parameter $ponatm$ indicates that some of the nitrogen deposition will be added to organic nitrogen pool ($fastN$) instead of to the IN pool.

For both nitrogen and phosphorus wet deposition is added through the concentration of precipitation. Precipitation may fall as snow and be added to the snow pack or if it is rain added to the potential infiltration. Depending on the fate of the infiltration, the wet atmospheric deposition will mix with the receiving water (e.g. surface runoff, soil layer water, macropore flow).
**Atmospheric deposition to rivers and lakes**

Nitrogen dry deposition on water surfaces is specified in GeoData.txt for each subbasin, while dry deposition of phosphorus is specified by a model parameter `drydepPP` (land use dependent).

For both nitrogen and phosphorus wet deposition is added through the concentration of precipitation (as described above), while for dry deposition an amount of the nutrient is added to the river (if it has a class-area) or lake water. If the lake is divided atmospheric deposition are added to the fast lake part (FLP).

Alternatively wet deposition of phosphorus on water surfaces can be specified by a general model parameter (`wetdepspl`) as a load. Monthly load of IN atmospheric deposition on water surfaces can be specified in GeoData.txt for each subbasin.

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Land routines

This section explains the computations in the land routines of HYPE. If you want an interactive overview of how the routines simulates runoff please have a look at the HYPE Runoff Explorer.

Basic assumptions

A subbasin in HYPE is divided into classes depending on land use, soil type etc. The classes are called SLC:s which stands for Soil type Land use Combination. This division can be compared to hydrological response units. Lakes and river is also classes, but this section consider only land classes.

HYPE model soil routine contains (up to) three soil layers. The number of soil layers and their respective lower limits (soillayerdepth in meters, figure 1) are listed by class in GeoClass.txt. It is possible to have a different number of layers and depth of these different classes.

The parameters of water retention in the soil are wilting point (wcwp), field capacity (wcfc) and effective porosity (wcep). These are exclusive and are specified in units of depth (figure 2). The model allocates the water retention capacity evenly between soil layers, depending on their thickness, if only one value set. It is also possible to specify one value per parameter and layer. The model uses water holding capacity in mm for the layers. These are represented in the model code and this document (figure 3) as wp(i), fc(i) and ep(i). The model parameters for the water storage capacity wcfc, wcwp, wcep (and layer depending alternative wcwp1, wcwp2, wcwp3, wcfc1, wcfc2, wcfc3, wcep1, wcep2, wcep3) depend on soil type.
The soil layer water content (mm) is represented in the document as $soil(i)$, where $i = \text{layer}$, but the value also depends on the subbasin and class (figure 3). The initial value of soil water is set to $wp + fc$. Optionally the initial value may be set to saturation ($wp + fc + ep$). The upper soil layer can hold more water than the porevolume. Standing water is not treated as a separate pool.

Tile drainage can be placed in any soil layer (figure 4). The depth of the drainage pipe in meters ($tiledepth$) is specified in GeoClass.txt for each class. A depth of 0 m is interpreted as drainage pipes are missing.

Another depth ($streamdepth$) is specified in GeoClass.txt; this is the maximum depth of the drainage to stream (or ditch). Soil water below this level does not contribute to the local runoff. Note that all
land runoff goes through the local river (and possibly local lake) and then the main river before it reaches the outlet lake. There is no direct runoff to lakes.

**Overview of flow paths**

![Flowpaths in the soil in the HYPE model](image)

**Figure 5:** Illustration of flowpaths in the soil in the HYPE model.

**Diagnostic variables**

Some additional output variables are calculated from the soil state variables.

**Groundwater level**

The groundwater level is measured negative from surface (0m) to bottom of the soil layers. A positive groundwater level means that the soil surface is below water. If the ground water table reaches above the surface, the water is calculated with 100% porosity.

The water table is found in the lowest soil layer that is not completely filled with water. Soil layers above this layer may have water in its effective porosity, but that is not included in the groundwater level output variable. The water table for a soil layer is calculated linearly from the proportion of water-filled pores of effective porosity part of the soil pore volume. If the soil moisture of a soil layer is at field capacity (or below), the groundwater level of that soil layer is at the bottom of the layer. If the pore volume is filled, the groundwater level of that soil layer is at the top of the layer.

**Soil moisture deficit**

Soil moisture deficit \((smdf)\) is calculated for the root zone, i.e. the upper two soil layers. It is the water (in \(mm\)) needed to fill the soil (\(soil\)) to field capacity (\(wp+fc\)).
\[ smdf = \sum_{k=1}^{2} \text{MAX} \left\{ fc_k + wp_k - soil_k, 0 \right\} \]

**Frost depth**

Ground frost depth (which is only calculated if the soil temperature is less than zero) depends on soil temperature (\(soill\)), but also on soil water content (\(soil\)), field capacity (\(fc\)), wilting point (\(wp\)) and two parameters \(frost\) and \(sfrost\).

\[ frostdepth = \frac{frost \times sfrost \times soill \times (fc + wp) \times soil}{soil} \]

There are two parameters in order to be able to choose if you want the frost depth to be land use dependent or soil dependent. The not used parameter is set to one.

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**Soil water**

**Groundwater runoff**

Runoff depends on the water table in relation to the drainage level. Runoff occurs when soil water reaches above field capacity in the soil layers. Runoff depends on soil water in the effective porosity (also used to calculate the groundwater table) and a recession coefficient (\(rc\)). If the soil is not saturated, runoff from the soil layer depends only on the water of that soil layer. Runoff occurs from all three soil layers (\(runoff(k), k=1\)-3) down to the drainage level, which is the depth of the stream (\(stream\)).
For the soil layer at drainage level, e.g. in the third layer of figure 6, and if the soil layer is not saturated, the runoff depends on the water level above the stream depth and the following equation replaces the one above.

\[ \text{runoff}(k) = \begin{cases} \text{rc}(k) \times \left( \text{soil}(k) - \text{wp}(k) - \text{fc}(k) \right) \times \left( \text{wp}(k) + \text{fc}(k) < \text{soil}(k) < \text{wp}(k) + \text{fc}(k) + \text{ep}(k) \right) \\ 0 \end{cases} \]

Soil layers that lie entirely below the stream depth have no groundwater runoff.

If a soil layer is saturated, i.e. \( \text{soil} > \text{fc} + \text{wp} + \text{ep} \), the runoff of the soil layer depends also on the water in the soil layer(s) above. For example if the drainage is in soil layer 3 and both soil layer 2 and 3 are saturated, the groundwater table in soil layer 1 determines the runoff of soil layer 3. The runoff is limited to the water above field capacity in the third layer though.

\[ \text{runoff}(3) = \text{MIN} \left( \text{soil}(3) - \text{wp}(3) - \text{fc}(3), \text{rc}(3) \times \frac{\text{deltah}}{\text{soillayerthick}(3)} \times \text{ep}(3) \right) \]
If the stream depth is below the bottom of the lowest soil layer. The extra distance will act as a level to increase $\Delta h$ and the runoff from the lowest soil layer.

**Recession coefficient of groundwater runoff**

The recession coefficient is calculated from two parameters, $rrcs1$ and $rrcs2$ which depend on soil type and a parameter that is general $rrcs3$. The recession coefficient is assumed to decrease with depth and the parameters indicate the coefficient value in the topmost layer ($rrcs1$) and in the bottom layer ($rrcs2$). If $rrcs2$ is not specified it is assumed to be similar to the recession in the topmost layer. The third parameter adjusts the upper layer recession due to the subbasin gradient ($slope$).

$$rrcs1 = rrcs1 \times \left(1 + rrcscorr \right) + rrcs3 \times slope$$

$$rrcs2 = rrcs2 \times \left(1 + rrcscorr \right)$$

The correction factor corrects $rrscorr$ parameters $rrcs1$ and $rrcs2$ for different parameter regions ($parreg$). It is defined as an increase. Note that the recession is limited to one. The recession is assumed to diminish exponentially and values of $rrcs1$ and $rrcs2$ applies to the midpoint of each layer ($d_k$).

$$rc\left(k\right) = rrcs \times e^{-b \times d_k}$$

The variable $b$ is an auxiliary variable.

$$b = \log\left(\frac{rrcs1}{rrcs2}\right)$$

$$b = \left(\frac{soillayerdepth(3) \times soillayerthick(3)}{2} \times \frac{soillayerthick(1)}{2}\right)$$

The result is:

$$rc\left(1\right) = rrcs1$$

$$rc\left(2\right) = rrcs1 \times \exp\left(-b \times \left(\frac{soillayerthick(1)}{2} + \frac{soillayerthick(2)}{2}\right)\right)$$
Runoff through drainage pipes

Runoff in the drainage pipes occurs when the water table (the percentage of filled pores of the effective porosity) rises above the pipe's depth (figure 7). Runoff depends on the groundwater surface elevation over the pipe (\(\text{deltah}, \text{m}\)), and a recession coefficient \(\text{trrcs}\). Recession parameter \(\text{trrcs}\) depends on soil type, while drainage pipe level depends on the class. The recession parameter is adjusted with the correction parameter \(\text{rrcscorr}\) for different parameter regions (\(\text{parreg}\)). It is defined as an increase.

\[
\text{trrcs} = \text{trrcs} \times (1 + \text{rrcscorr})
\]

Depending on which soil layer drainage pipe is in, the runoff will be calculated for water in that soil layer. For the soil layer \(k\) (\(\text{soil}(k)\) is the water content in soil layer \(k\)) runoff is calculated as the parameter \(\text{trrcs}\) times the water found in the effective porosity of the layer and of the overlying soil layers if it is full.

\[
\text{deltah} = \frac{(\text{soil}(k) - \text{wp}(k) - \text{fc}(k))}{\text{ep}(k)} \times \text{soillayerthick}(k) - \left(\frac{\text{soillayerdepth}(k)}{\text{tiledepth}}\right)
\]

\[
\text{IF(soil}(k) - \text{wp}(k) - \text{fc}(k) - \text{ep}(k) \geq 0.) \quad \text{deltah} = \text{deltah} + \frac{(\text{soil}(k-1) - \text{wp}(k-1) - \text{fc}(k-1))}{\text{ep}(k-1)} \times \text{soillayerthick}(k-1)
\]

\[
\text{IF(deltah} > 0.) \quad \text{runoffd} = \text{trrcs} \times \text{deltah} / \text{soillayerthick}(k) \times \text{ep}(k)
\]

Figure 7: Illustration for calculation of runoff through the drainage pipes.

Infiltration

Infiltration is calculated from the sum of rain and snowmelt (\(\text{infilt0}, \text{mm/time step}\)).

\[
\text{infilt0} = \text{rainfall} + \text{melt}
\]

Part of the available water for infiltration (\(\text{infilt0}\)) may not infiltrate into the soil, due to limitations by the soil's infiltration capacity and other properties of the soil. The calculation of actual infiltration will consider effects of surface runoff, macropore flow and frozen soil. If the finally calculated infiltration is greater than zero, it is added to the upper layer soil water. This is done regardless of whether there is space in the soil pores there or not. If the water exceeds the water pore volume it is assumed to lie on the ground, but it still belongs to the upper soil layer, is totally mixed and thus has the same concentrations.
HYPE has an option for alternative calculation order of soil processes during a timestep. As default it calculates and add infiltration (and let the soil water percolate) before runoff and evaporation is calculated and removed from the soil water. Alternatively runoff and evapotranspiration is calculated before infiltration and percolation to slow the response of soil runoff. These options is tested during development of the soil routine.

**Diversion of surface runoff and macropore flow**

Surface runoff due to excess infiltration and macropore flow are calculated from the sum of snow melt and rainfall; the water available for infiltration ($infilt0$).

If the current infiltration rate is greater than a threshold ($mactrinf$, mm/timestep) then macropore flow ($macroflow$) and surface runoff ($infoverflow$) may occur. In addition, the water in the upper soil layer needs to be larger than another threshold ($mactrsm$) for surface runoff and macropore flow to occur. The two flows are calculated as a percentage ($macrate$ respective $srrate$) of the infiltration above the first threshold;

\[
macroflow = macrate \times (infilt0 - mactrinf) \\
infoverflow = srrate \times (infilt0 - mactrinf)
\]

All the four aforementioned parameters are soil type dependent. If $macrate$ and $srrate$ together are greater than one, they are weighted so that their sum is one prior to calculation of the surface runoff and macropore flow;

\[
macroflow = \left( \frac{macrate}{macrate + srrate} \right) \times (infilt0 - mactrinf) \\
infoverflow = \left( \frac{srrate}{macrate + srrate} \right) \times (infilt0 - mactrinf)
\]

The actual infiltration is calculated by subtracting the macropore flow and surface runoff from the sum of snow melt and rain.

\[
infilt = infilt0 - macroflow - infoverflow
\]

**Additional infiltration limitation by frozen soil**

An optional model for infiltration limitation and diversion of flow considers the effect of frozen soil. It is developed based on Zhao and Gray (1999). This model redirects all or part of the remaining infiltration, after calculating the diversion of surface runoff and macropore flow as described above.

If the minimum daily temperature is less than 10 degrees and the infiltration is larger than 5mm/d an ice lens is created in the soil. In this case, and as long as the maximum daily temperature is below zero, the ice lens redirect all infiltration to surface runoff and macropore flow.
If there is no ice lens, but the soil temperature of the upper soil layer (soiltemp) is below zero the infiltration is restricted but not blocked. The infiltration is restricted by a potential infiltration adapted from Zhao and Gray (1999). The potential infiltration (potinfilt) depends on a model parameter (bfroznsoil) that is soil type dependent. It also depend on the “opportunity time” (t0), which is an estimate of the time with possible infiltration in hours;

\[
potinfilt = \begin{cases} 
0 & \text{for } soil \geq pw \\
\frac{bfroznsoil \times 0.99^{2.92} \times \left(1 - \frac{soil}{pw}\right)^{1.1} \times \left(\frac{0 - soiltemp}{273.15}\right)^{-0.45} \times t0^{0.44}}{24} & \text{for } soil < pw
\end{cases}
\]

\[
t0 = \begin{cases} 
1 & \text{for } 0.05 \times snow < 6 \\
0.05 \times snow - 5 & \text{for } 0.05 \times snow > 6
\end{cases}
\]

Here soil is soil water of the upper soil layer (mm), pw is pore volume of the upper soil layer (mm) and calculated from the model parameters for water holding capacity (wcwp, wcfc and wcep), and snow is snow water equivalent (mm).

If the actual infiltration is greater than the potential infiltration then the overshoot is redirected to surface runoff and macropore flow.

\[
redirect = infilt - potinfilt
\]

The here redirected infiltration is added to the macropore flow and overland flow of the basic model proportionally to their respective model parameters (macrate and srrate).

\[
macroflow = macroflow + \left(\frac{macrate}{macrate + srrate}\right) \times redirect
\]

\[
inflowflow = inflowflow + \left(\frac{srrate}{macrate + srrate}\right) \times redirect
\]

**Percolation**

The flow of water downward through the soil layers is only done by water over field capacity (water in the effective porosity). A maximum percolation (mm/d) limits the flow between soil layers. For the upper soil layer it is mperc1, and for the second soil layer it is mperc2. These parameters are soil type dependent. Flow is also limited by how much water the lower layer can receive.

Drainage from soil layer 1 to soil layer 2 is
\[ \text{perc1x} = \text{MIN}\left(\left(\text{soil}\left(1\right) - \text{wp}\left(1\right) - \text{fc}\left(1\right)\right), \text{mperc1}\right) \]

but if there is not enough capacity in soil layer 2 the drainage is instead
\[ \text{perc1} = \text{wp}\left(2\right) + \text{fc}\left(2\right) + \text{ep}\left(2\right) - \text{soil}\left(2\right) + \text{perc2} \]

and fills the second soil layer.

Drainage from soil layer 2 to soil layer 3 can be at most
\[ \text{perc2x} = \text{MIN}\left(\text{wp}\left(3\right) + \text{fc}\left(3\right) + \text{ep}\left(3\right) - \text{soil}\left(3\right), \text{mperc2}\right) \]

for that is what the soil layer 3 can receive. If soil layer 2 does not reach field capacity with \text{perc1x} added
\[ \text{perc2} = 0 \]

but if soil layer 2 with \text{perc1x} added exceeds field capacity
\[ \text{perc2} = \text{MIN}\left(\text{soil}\left(2\right) + \text{perc1} - \text{wp}\left(2\right) - \text{fc}\left(2\right), \text{perc2x}\right) \]

Regional groundwater flow is created by additional percolation from soil (see Section on Regional groundwater flow).

**Upwelling**

Flow may enter the lowest soil layer, i.e. regional groundwater flow. Upwelling to soil layers above may occur if the soil layer is filled.

**Saturated surface runoff**

Surface runoff due to a high ground water table \(q\), mm/time step) occurs when the water table in the upper soil layer reaches above the surface. It depends on a parameter \text{srrcs} which is dependent on land use. The recession parameter is corrected with the correction factor \text{rrcscorr} for different parameter regions (parreg). It is defined as an increase.

\[ \text{srrcs} = \text{srrcs} \times \left(1 + \text{rrcscorr}\right) \]
\[ q = \text{MAX} \left( srrcs \times \left( \text{soil}(1) - \text{wp}(1) - \text{fc}(1) - \text{ep}(1) \right), 0 \right) \]

Runoff is removed from the uppermost soil layer. The total surface runoff (due to high ground water table and low infiltration capacity) is calculated and printed.

**Macropore flow**

Macropore flow occurs when the potential infiltration and water in the upper soil layer is large (see Diversion of surface runoff and macropore flow above). It is caused by a limited infiltration capacity of the soil. Macropore flow (macroflow) is added to the layer in which the water table is located (see Diagnostic variables above). The water is added to this layer only until it is full and the excess is trapped in the layer above.

Thus water does not flow up into the layer above when macropore flow is larger than the empty space in the soil layer with the water table, as in the case of groundwater inflow. Instead the excess flow stays in the soil layer above before reaching the soil layer of the water table. This distinction is important for the substances following the macropore flow.

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Snow routines

Snow melt

As default, snow melting occurs when the temperature is greater than the threshold temperature. The amount of snow (in mm) that melts \((melt)\) depends on the snowmelt parameter \(cmlt\), threshold temperature parameter \(ttmp\) and air temperature \((temp)\).

\[
melt = \text{MIN} \left( cmlt \times \left( temp - ttmp \right), \text{snow} \right)
\]

The parameters \(cmlt\) and \(ttmp\) are related to land use.

Snow melt models

Alternative snowmelt models exist, but are not fully described here yet.

Model 0 (default)

Temperature index model, without snow cover scaling

\[
melt = \text{MIN} \left( cmlt \times \left( temp - ttmp \right), \text{snow} \right)
\]

Model 1

Temperature index model, with snow cover scaling

\[
melt = \text{MIN} \left( cmlt \times \left( temp - ttmp \right), \text{snow} \right) \times \text{snowcover}
\]
Model 2

Temperature and radiation index model, with snow cover scaling

Snow cover

Normally snow is assumed to cover the whole class if present. Alternatively if parameters are given, snow cover fraction (fsc) within a class is calculated based on snow water equivalent (snow). The formulation is based on Samuelsson et al. (2006). During snow build up the snow cover increase as a function of snow water equivalent until a maximum value (general parameter fscmax) is reached.

\[ f_{sc} = f_{scmax} \times \tanh \left( 0.1 \times \text{snow} \right) \]

It is also possible to specify a minimum snow cover (general parameter fscmin). As soon as the fractional snow cover area reaches above a certain threshold (fscmax-fsclim), the snow cover area is determined by another relation that represents the redistribution of snow during winter. In this case snow cover is dependent on maximum snow pack during the winter (snowmax) and a snow cover redistribution factor that is dependent on variation in elevation (stdelev, the standard deviation of elevation within the subbasin) and land use.

\[ f_{sc} = \frac{\text{snow}}{\text{snowmax} \times f_{scdist}} \]

\[ f_{scdist} = f_{scdist0} + f_{scdist1} \times \text{stdelev} \]

The snow distribution factor (fscdist) is determined by three land use dependent parameters; fscdist0 and fscdist1 in the linear equation and a maximum value (fscdistmax). Also in this case the snowcover is limited by the maximum and minimum value parameters. When the end of the snow season approaches (defined by general parameter fsck1) the snowmax variable is gradually decreased in order to be reset before next winter season:

\[ \text{snowmax} = \text{snowmax} - \frac{\left( f_{sc1} \times \text{snowmax} - \text{snow} \right)}{f_{sc1}} \times e^{-f_{scexp} \times ts} \]

\[ \frac{\text{snow}}{\text{snowmax}} < f_{sc1} \]

The equation depends on two general parameters, fsck1 and fsckexp, where fsckexp depend on time (ts is seconds per timestep of simulation).

For winters when the snow pack not reach the definition of large snow pack, the first equation is used during the whole season.

Soil temperature and snow depth

Soil layer temperature (soiltemp) is calculated as a balance of three temperatures; previous time step soil layer temperature, soil temperature at deep depth (deeptemp) and air temperature (T). The weight of the deep soil is constant (0.001), while the weight of the air temperature (weightair)
depends on snow depth (snowdepth) and parameters. The soil memory (soilmem) depends on depth
and land use, with parameters surfmem and depthrel. The memory of deep soil temperature is a
general parameter (deepmem).

\[
soilmem = \begin{cases} 
    deepmem & \text{for deepmem} \\
    surfmem \times e^{-depthrel \times depth(k)} & \text{for soil layer } k 
\end{cases}
\]

\[
weight_{air} = \frac{1}{soilmem + 10 \times snowdepth}
\]

\[
deeptemp = weight_{air} \times T + \left(1 - weight_{air}\right) \times deeptemp
\]

\[
soiltemp = weight_{air} \times T + \left(1 - weight_{air} - weight_{deep}\right) \times soiltemp + weight_{deep} \times deeptemp
\]

In the default snow depth model, snow density (snowdens) depends on the snow's age in days
(snowage). Snow density for fresh snow (sdnsnew) and the increase of density with snow age
(snowdensdt) are general parameters (~ 0.1 and ~0.002). The snow's age increases by one every
time step, but are weighted with age (0) for any new snow.

\[
snowage = \frac{\left(snowage + 1\right) \times oldsnow}{oldsnow + snowfall}
\]

\[
snowdens = sdnsnew + snowdensdt \times snowage
\]

\[
snowdepth = \frac{0.1 \times snow}{snowdens}
\]

In the alternative snow depth model, snow density is calculated by a compacting factor. Snow density
for fresh snow (sdnsnew), maximum snow density (sdnsmax), compaction rate for low temperatures
(sdnsrate) and additional compaction for high temperature (sdnsradd) are all general parameters.
The change in snowdensity (densdt) due to compaction each time step is calculated as:

\[
densdt = sdnsrate \times \left(sdnsmax - snowdens\right)
\]

for cold days (temperature is below threshold temperature parameter tttmp), and

\[
densdt = \left(sdnsrate + sdnsradd\right) \times \left(sdnsmax - snowdens\right)
\]

for warm days.

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**Glaciers**

The glacial class' area \( \text{class area} \) is divided into snowfield and glacier \( \text{glac area} \). The division may vary with simulation time, but the glacier cannot be larger than the slc class' area. At simulation start the area is considered to be all glacier, unless specific information on which year the class area is valid is given. The initial glacier volume is calculated based on glacier area or set for each glacier. If the initial year is given the initial volume is valid for that date. In this case, year of the area and/or volume and the average annual change in massbalans \( \text{annmb} \) in m/yr is given as input data or model parameters. The initial glacier volume is then adjusted according to the difference in years \( \text{yeardiff} \) between simulations start and glacier area data.

The glacier volume - area relationship:

\[
\text{glac vol} = \text{coef} \times \text{glac area}^\text{exp}
\]

\[
\text{coef} = \text{coef} \times e^c
\]

The initial glacier volume if calculated from class area:

\[
\text{glac vol} = \text{coef} \times \text{class area}^\text{exp} + \text{yeardiff} \times \frac{\text{annmb} \times \text{class area}}{\text{glac dens} \times 1000}
\]

The glacier area is correspondingly calculated from the glacier ice volume, but is limited by the class area. The relation is:
The equation coefficients \( \text{coef} \) and \( \text{exp} \) can have different values for specific glaciers. The first coefficient \( \text{coef} \) is calculated as the product of \( \text{EXP}(\text{c}) \), where \( \text{c} \) is a glacier volume correction, and a general parameter \( \text{(glacvcoef/glacvcoef1)} \) depending on glacier type. The second coefficient, \( \text{exp} \) is a general parameter \( \text{(glacvexp/glacvexp1)} \) depending on glacier type. Glacier density \( \text{(glacdens)} \) is a general model parameter \( (\text{m}^3 \text{ water}/ \text{m}^3 \text{ ice}) \).

Glaciers are divided into four types. The default type is mountain glacier, the alternatives are ice cap, ice sheet and infinite glacier. Glacier type is given as input, or determined by the glacier area (a threshold \( \text{(glac2arlim)} \), a general parameter). The glacier area is used to determine the glacier type if it is not given as input and the threshold parameter is set. The glaciers will then be divided into mountain glaciers and ice caps.

The default glacier and the ice cap glacier type differ by having different parameter values for the volume-area relationship. The default values for the volume-area relationship are based on Radic and Hock (2010).

The ice sheet glacier do not simulate an snow field, and the glacier will have a constant area. The ice sheet glacier type is thus not depending on the volume-area relationship, except for initial volume if that is not given. In that case the initial volume of the ice sheet glacier will used the ice cap parameter values.

The infinite glacier type is independent on glacier volume, because the glacier melting is not limited by the glacier volume. It has a constant area (the class area) and do not simulate an snow field.

For snowfields, all soil processes are calculated as for the common land classes. Snow pack is only calculated for the snowfield, i.e. snow depth is assumed zero on glaciers. For glaciers, all precipitation is added to the glacier ice. Glacier melting is calculated as:

\[
melt = \text{cmlt} \times \left( T - \text{ttmp} \right)
\]

where \( \text{cmlt} \) is the general parameter \( \text{glaccmlt} \) and \( \text{ttmp} \) the general parameter \( \text{glacttmp} \).

An alternative glacier melting model exist, which depend on class' temperature \( (T) \) and radiation \( (\text{swrad}) \) and has a refreezing component (governed by general parameter \( \text{crefr} \)). The radiation component depend on the albedo of the glacier, which in turn depends on if the glacier is snow covered \( (\text{snowcov}) \) or not, and general parameters for rate \( (\text{cmrad}) \) and glacier ice albedo \( (\text{glacalb}) \). The refreezing component is limited to the glacier melt.

\[
melt = \begin{cases} 
\text{cmrad} \times \text{swrad} \times \left( 1 - \text{albedo}_{\text{glacier}} \right) + \text{cmlt} \times \left( T - \text{ttmp} \right) & \text{for } T \geq \text{ttmp} \\
\text{cmrad} \times \text{swrad} \times \left( 1 - \text{albedo}_{\text{glacier}} \right) - \text{crefr} \times \text{cmlt} \times (\text{ttmp} - T) & \text{for } T < \text{ttmp}
\end{cases}
\]

\[
\text{albedo}_{\text{glacier}} = \text{albedo}_{\text{snow}} \times \text{snowcov} + \text{glacalb} \times \left( 1 - \text{snowcov} \right)
\]

Glacier evaporation is optional and set by an model option \( \text{(snowevaporation)} \). The evaporation is determined as a fraction of the potential evaporation by the general model parameter \( \text{fepotglac} \).

\[
\text{glac}_{\text{area}} = \left( \text{glac}_{\text{vol}} \times \frac{1}{\text{coef}} \right)^{\text{exp}}
\]
Glacier melting, snow melting on snowfield and/or rain are added to the soil. Thus the soil represents the whole glacier class area. The concentrations in the glacial melt water are zero. This means that any atmospheric deposition of nutrients is lost on the glacier.

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References


Rivers and lakes

Basic assumptions

The HYPE model can contain two types of rivers, local stream and main river, and two types of lakes, local lakes and outlet lakes (Figure 1). Local and main rivers are present in all subbasins and the length of each is calculated as the square root of the subbasin area. The length of the watercourses can be given as input. The river can be a SLC class and is then given an area, but rivers can also be one-dimensional (i.e. no fraction of the subbasin area are occupied by the river and no precipitation added to the river). All local runoff is entering the local river. Local lakes (ilake) receive a portion of the local runoff. The flow leaving the local river (including flow from local lake) goes to the main river of the same subbasin. If there are upstream subbasins their flow is added to the local flow when both flows flow into the main river. Outlet lakes (olake) receive the outflow from the main river, i.e. all upstream and local flows.

![Figure 1](image)

Figure 1: Schematic representation of streams and lakes in HYPE, and the link between them.

The two lake types are separate classes. The lake classes have characteristics such as land use and soil type, which are defined together with the other classes' characteristics (in GeoClass.txt). Precipitation, atmospheric deposition and evaporation of rivers and lakes are calculated first, while river flow and inflow, transformation processes and the outflow of the lakes is calculated thereafter. Lakes and rivers are calculated in the model's routing part after all classes are calculated for the subbasin.

An outlet lake can be part of a larger lake. It is then called a lake basin. Lake basins are olakes in nearby subbasins.

A simple outlet lake has a threshold. The outflow ends if the water level drops below the threshold. Lake mean depth below the threshold is specified in GeoData.txt or LakeData.txt as `lake_depth` in meters. Lake depth can also be set by parameters, i.e. general parameter `gldepo` or olake region parameter `olldepth`. The threshold is also the the water level of the lake at the start of a simulation. The current water level is denoted as `wlm` in Fig. 2. For printing, the outlet lake water level (output
variable $w_{com}$) is calculated in meters and you can set a reference level ($w_{0ref}$) in LakeData.txt to get the same height system as any observations of the lake’s water level. The lake’s $w_{0ref}$ is added to the water level above the threshold. A regulated lake (dam) has two thresholds. One, same as for a simple lake, is used for spill, and one lower threshold were outflow ends completely. The distance between the thresholds are determined by the regulation volume. HYPE assumes the lake/dam has vertical sides in the calculations, thus the observed variation in water level may be larger than the simulated variation. It is therefore possible to adjust the output $w_{com}$ (and $w_{cav}$) for the actual amplitude of the regulation volume ($w_{amp}$). This will make the simulated and recorded water stage comparable below the threshold for a regulated lake.

![Lake diagram](image.png)

Figure 2: An outlet lake with some variables.

A local lake also has a threshold depth that is used as start value. The depth is given by general parameter $gldepi$ and is then the same for all the local lakes, or by ilake region parameter $illdepth$. It is measured in meters. A percentage of flow from the local stream flows into the local lake. The rest of the local flow runs directly to main river watercourse.

Using parameters, you can divide the lake into two parts, one with faster flows (FLP) and one with slower flows (SLP) (Figure 3). This function is used for the simulation of nutrients to simulate stratification, strangulation or other phenomena that may limit the mixing of a lake. With this feature, the flows through the lake follow the schedule below. The split is determined by the parameter $deeplake$, which is the fraction of the lake’s initial volume SLP, the remaining (varying) volume in the lake is the FLP. The parameter $fastlake$ determines where the outflow will be coming from. Default is that outflow is taken from the slow lake part. Increasing $fastlake$ will let the FLP contribute to outflow. $Fastlake$ equal to one gives the maximum contribution of FLP, and the outflow will be taken proportionally from the two lake parts according to their volume.
The streams may have a delay and an attenuation of the flow. The former shifts the peaks, but leaves them otherwise untouched, while the attenuation both delays and smooths out peaks. The streams have a dead volume in which the residence time of solutes is increased relative to the water and the concentration smoothed out further over time.

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Rivers
Common rivers processes

Precipitation

Daily precipitation is added to the river if it has an area (is a class) and a new concentration is calculated. Precipitation is divided between river water in damping box and queue according to the respective volumes.

Evaporation

If the river has an area (is a class), it evaporates and a new concentration is calculated. Normally the river area is constant over time, but with parameters a reduction of riverarea can be simulated for low volume/flow. The reduced river area is also used for heat exchange calculations.

\[
frac{area} = \begin{cases} 
0 & x \leq 0 \\
\left(1 + \frac{x - \text{fraxe}}{\text{fraxe} - \text{fraxm}}\right) \times \left(\frac{x}{\text{fraxe}}\right)^{\frac{x}{\text{fraxm}} - 1} & 0 < x < \text{fraxe} \\
1 & x \geq \text{fraxe}
\end{cases}
\]

The parameters \(\text{fraxe}\) and \(\text{fraxm}\) are general, and \(\text{fraxe}\) is (minimum) mean river depth (m) where fractional river area = 1 and \(\text{fraxm}\) is mean river depth (m) where the slope of the fractional river area has its maximum (must be in the range between 0 and \(\text{fraxe}\)). \(x\) is the current mean river depth (based on full area extention).

Pure delay

The delay in the watercourse (\(\text{transtime}\)) in days is determined by the length of the watercourse (\(\text{rivlen}\)) and the water’s maximum velocity (\(\text{rivvel}\)). The maximum velocity is a general parameter with unit m/s. The delay in the river is dependent on subbasin land area if the default river length is used. The delay is a pure translation. The delay is divided into whole days (\(\text{ttday}\)) and parts of the day (\(\text{ttpart}\)).

\[
\text{transtime} = \frac{\text{rivlen}}{\text{rivvel} \times 8.64 \times 10^4}
\]

\(\text{ttday} = \text{INT}(\text{transtime})\)

\(\text{ttpart} = \text{TRANSTIME} - \text{REAL}(\text{ttday})\)

The inflow of the river is stored in two arrays (\(\text{riverq}\) and \(\text{rierc}\)) until it is time for it to flow out of the
river stretch. The outflow is weighted by using the parts of the time step (\( ttpart \)) that are to flow out during the time step.

\[
\begin{align*}
\text{transq} &= (1-ttpart) \times \text{riverq}(\text{ttday}) + ttpart \times \text{riverq}(\text{ttday}+1) \\
\text{IF}(\text{transq}>0) \text{THEN} & \quad \text{transc} = ((1-ttpart) \times \text{riverq}(\text{ttday}) \times \text{riverc}(\text{ttday}) + ttpart \times \text{riverq}(\text{ttday}+1) \times \text{riverc}(\text{ttday}+1)) / \text{transq} \\
\text{ELSE} & \quad \text{transc} = 0.
\end{align*}
\]

After the calculation of outflow the values in the arrays are shifted forward one time step.

**Delay and attenuation**

If the delay in the watercourse also includes a damping of the peaks then only part of the delay is considered as translation, while the rest of the delay lies in damping. The translation is calculated first and then the flow goes through a linear box that creates attenuation. The parameter damp indicates how much of the delay that will occur in the attenuation box (or damping box), and translation time is reduced accordingly. Otherwise the translation is calculated in the same manner as above.

\[
totalltime = \frac{\text{rivlen}}{\text{rivvel} \times 8.64 \times 10^1}
\]

\[
\text{transtime} = (1-\text{damp}) \times \text{totalltime}
\]

The result from the translation of water (\( \text{transq} \) and \( \text{transc} \)) flows into the attenuation box, which is assumed to be completely mixed. The delay time in the box (\( kt \)) is recalculated to a corresponding recession coefficient (\( \text{riverrc} \)). The recession coefficient (\( \text{riverrc} \)) used to calculate the outflow from the box (\( \text{dampq} \)) is a function of volume in the box (\( \text{riverbox} \)).

\[
kt = \text{damp} \times \text{totalltime}
\]

\[
\text{riverrc} = 1 - \text{kt} + \text{kt} \times e^{-1/\text{kt}}
\]

\[
\text{dampq} = \text{riverrc} \times \text{riverbox}
\]

The delay in the attenuation box is at the centre of gravity of the flow (not the tip) and is not calculated exactly (see Derivation of the equation of recession in the attenuation box below). Therefore there may be a difference in a peak delay between the use of pure delay and delay with damping, although the maximum velocity is set to the same value. In the case of damping the max velocity parameter \( \text{rivvel} \) may be approximated.

**Additional smoothing of dead volume**

A dead volume (\( \text{deadriver} \)) can be used to get extra equalization of concentrations in the watercourse. Then the outflow equation for the attenuation box (\( \text{riverbox} \)) is:
In this case, the initial value of the attenuation box is equal to the dead volume. Inflows into the box are mixed with the entire volume. At present this represents a dilution of the concentration since the initial value of concentration in the box is zero. At present the estimated dead volume as dependent on the upstream area (including the area of the subbasin itself). Depth is calculated from the parameter dead (deadl or deadm, m²/ km²) and the upstream area (km²), the width is assumed to be 10 times the depth and the river length (m) is estimated to be the root of the subbasin’s area unless it is set among the input data. This gives the dead volume:

\[ \text{deadriver} = \text{dead} \times \text{upparea} \times \text{rivlen} \]

Dead volume can be used even without suppression of water (damp = 0).

**Derivation of the equation of recession in the attenuation box**

The continuous-time equation for the change in volume \( S \) in a reservoir is:

\[
\frac{dS}{dt} = i(t) - q(t)
\]

where \( i(t) \) is influx and \( q(t) \) is outflow. If we assume that the outflow is proportional to the volume we get another expression for the change in \( S \). \( k \) is in the unit of time.

\[
q(t) = \frac{1}{k} S,
\]

\[
\frac{dS}{dt} = k \frac{dq}{dt}
\]

Combining these two equations gives a first degree equation that can be solved using the method of “integrating factor”:

\[
\frac{dq}{dt} + \frac{1}{k} q = \frac{i}{k},
\]

\[
\frac{d}{dt} e^{t/k} q = \frac{i}{k} e^{t/k},
\]

\[
q(t) = \frac{1}{k} \int i(t) e^{\tau - t/k} d\tau
\]

In our case, we have a constant influx during a time steps of length 1 day. Assume that the influx during a time steps are \( I \) and the outflow \( Q \). These flows are thus the equivalent discrete flows.

Because \( I \) is constant during the time step \( (i(t) = I) \) \( Q \) can be calculated. Define

\[
y(t) = e^{t/k} q(t)
\]

and integrate the second equation above, from 0 to \( t \) (\( t = 1 \), one time step).
\[
\int \frac{dy}{dt} \, dt = \int_0^t \frac{i(t)}{k} e^{\tau/k} d\tau,
\]
\[
y(t) - y(0) = \frac{I}{k} \int_0^t e^{\tau/k} d\tau = I \left( e^{t/k} - 1 \right)
\]
\[
e^{t/k} q(t) - q(0) = I \left( e^{t/k} - 1 \right)
\]
\[
q(t) = q(0) e^{-t/k} + \left( 1 - e^{-t/k} \right)
\]

Calculate \( Q \) for time step (the mean of \( q(t) \)) and replace the outflow with the volume.

\[
Q = \frac{1}{c} \int_0^1 q(t) dt = \frac{1}{c} \int_0^1 q(0) e^{-t/k} + I \left( 1 - e^{-t/k} \right) dt
\]
\[
= q(0) \left[ 1 - e^{-t/k} \right] dt + I \left[ 1 - e^{-t/k} \right] dt
\]
\[
= q(0) \left[ e^{-t/k} - 1 \right] + I \left[ e^{-t/k} - 1 \right] - I \left[ e^{-t/k} - 1 \right]
\]
\[
= I \left( e^{-1/k} - 1 \right) \left( kI - kq(0) \right)
\]
\[
Q = \left( I + e^{-1/k} - 1 \right) \left( kI - S(0) \right)
\]

Recession coefficient is defined as the proportion of outflow after an inflow pulse. It is therefore calculated here as \( r = Q/I \) under the assumption that the starting volume can be neglected, \( s(0) = 0 \).
For a real flow the volume can of course not be neglected in this way and the relationship is only approximate.

**Bank-full flow**

Bank-full flow is used for erosion of particulate phosphorus in rivers (see Sedimentation/Resuspension). It is approximated by the second highest daily flow during the last year.

**Local river**

The local river has a length equal to the square root of the subbasin area, if not specified as input. Runoff from the land area of the subbasin forms the inflow to the local river. The flow in the local river is delayed and attenuated as described above. Of the resulting flow from the stream a constant percentage goes to the local lake (icatch), and the rest directly to the main river.

**Main river**

A main river is present in all subbasins. The length is equal to the square root of the subbasin area, if it is not specified as input. In areas without incoming water from upstream there is still a main river, but it receives only local river flow after the local lake. In subbasins with upstream incoming water, the flow to the main river will be the sum of outflow from the local lake, the proportion of flow in the local river not flowing into the local lake and the water from upstream.

The return flow from an aquifer is added to the inflow of the main river.

The flow in the main river is delayed and attenuated as described above. The resulting effluent from the river flows to the outlet lake or to the next subbasin if there is no outlet lake.

**Inflow from upstream subbasins**

In input files it is given to which subbasin(s) the outflow from each subbasin flows. The upstream flow enters the main river of the downstream subbasin. Inflow into the main river of a subbasin is calculated by adding outflows from upstream areas. Concentrations are flow-weighted by their relative share. The upstream flow enters the main river, except if the upstream subbasin has a lakebasin that is part of the downstream subbasin's lake. In this case the upstream lakebasin outflow goes directly into the downstream lakebasin.

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### Lakes

#### Common lake processes

**Precipitation**

Daily precipitation is added to the lake and a new concentration is calculated. Lakes are assumed to be completely mixed if the parameter `deeplake` is not set to a value greater than zero.

**Evaporation**

The lake is assumed to evaporate and a new concentration calculated.

**Outflow with rating curve**

The rating curve is used for calculation of outflow above a threshold at several occasions.

\[ q(t) = k \times \left( w(t) - w_0 \right)^p \]

In the program the equation is solved with linearization (Lindström, 2016). For rating curve exponent
equal to one, the solution is exact, while for other exponent an approximation is used.

**Local lake (ilake)**

The local lake has an initial volume which is determined by its area and threshold depth. The lake depth is determined by model parameters \((gldepi\) or \(illdepth\)).

The inflow to the lake is a percentage of the flow in the local stream \((icatch\). The percentage is determined by the percentage of the subbasin area that drain to the lake. This percentage can be given as a fraction in GeoData.txt for each subbasin with an internal lake, or be given by a regional parameter \(ilicatch\), or be given by a general parameter \(gicatch\). If not set at all the default value is 1, i.e. the local river runs through the local lake. The flow from the local river is added to the lake. The lake water is assumed completely mixed if \(deeplake = 0\).

Water outflow is calculated with the universal rating curve using general or region specific model parameters.

**Simple outlet lake or dam (olake)**

**Inflow**

The outlet lake receives both local runoff and inflow from upstream areas via the main river. An outlet lake may receive inflow from regional groundwater flow.

**Outflow with universal rating curve**

If the water level \((wlm\)) is higher than the threshold \((lake_depth\)) the outflow is calculated with the following equation:

\[
outflow = \begin{cases} 
0, & \text{for } wlm < lake_depth \\
gratk \times uparea^{grata} \times ratcorr \times \left( \frac{wlm - lake_{depth}}{lake_{depth}} \right)^{gratp}, & \text{for } grata > 0 \\
gratk \times ratcorr \times \left( \frac{wlm - lake_{depth}}{lake_{depth}} \right)^{gratp}, & \text{for } grata = 0
\end{cases}
\]

\(gratk\), \(gratp\) and \(grata\) are general parameters that apply to all lakes in the model application. \(ratcorr\) is parameter region dependent model parameter for adjusting \(gratk\) to different regions. The upstream area \((uparea, \text{km}^2)\) is included in the equation if parameter \(grata\) is \(> 0\). If the water level is below the threshold then the outflow is zero.

**Outflow with specific rating curve**
In LakeData.txt a specific rating curve may be set instead of the general parameters. This is done by setting the rate and exp larger than zero in LakeData.txt.

\[
out\ flow = rate \times \left( wlm - lake\ depth \right)^{exp}
\]

**Regulated lake**

A simple regulation routine can be given in LakeData.txt for selected outlet lakes. There are two main versions of regulations. The first determine a (constant) production flow between the lake threshold and a lower water stage. The second determine different rating curves for two production periods.

**Production flow**

The regulated outflow is independent of water level \((wlm (m))\) between the threshold \((w0=\text{lake\ depth})\) and a minimum water level \((wmin)\). For this interval the production flow \((qprod)\) is used. Production flow may depend on time of year and the water level. The minimum water stage is not really a water stage, but is calculated from the regulation volume \((\text{regvol})\) and the lake (surface) area. Thus it does not consider the area changing with depth.

For water levels above the threshold, the flow is calculated with a rating curve (typically flow through spillways) or all the water is discharged, but it is at least equal to the production flow \((qprod)\).

Compiled together as one equation, the outflow from a regulated lake is:
Input variables $rate$, $exp$, $qprod$, $regvol$, $w0$, $qamp$ and $qpha$ can be found in LakeData.txt. The variable $wmin$ is calculated by the program from $regvol$ and lake area:

$$wmin = w0 - \frac{regvol \times 1000000}{area}$$

Production flow can have two different values during the year, which depends on the day of the year. This is determined by the input variables $qprod1$, $qprod2$, $datum1$ and $datum2$. Regulation period 1 between $datum1$ and $datum2$ has production flow $qprod1$, while the rest of the year has production flow $qprod2$. Not setting the dates gives the same production flow the whole year ($qprod1$).

Production flow can alternatively be made to vary sinusoidal over the years, with a peak in December, when power output is normally high, and a minimum in June. This is done with the input variable $qamp$. If you want a different seasonal variation set $qpha$ (default = 102).

$$qprod = qprod \times \left(1 + qamp \times \sin \left(\frac{2 \times \pi \times (\text{dayno} + qpha)}{365}\right)\right)$$

In addition, the production flow is reduced from this level when there is a low water level in the dam. The flow will then be reduced linearly from full production flow for that time of year when water level is over the limit ($limqprod$ (percentage of volume)) to zero when no water is left in the dam over $wmin$ level.

Two rating curves

Regulated outflow can also be calculated from different rating curves for the regulation periods. The rating curve parameters are the specific rate and exponent of is lacking the general rate and exponent.
Dams of specific purpose

In addition to regulated lakes (described above), dams in HYPE can instead be regulated by a specific purpose. Dams are then governed by the main purpose of the dam, e.g. hydropower, flood control. The rules of calculating the outflow of the dam then depends on the specified purpose of each dam. The outlet lakes that are regulated by a specified purpose is given in DamData.txt. A dam without defined purpose may use the methods given in previous sections (and be defined in LakeData.txt).

Dams are characterised by their regulation volume ($\text{regvol}$) between a minimum water stage ($\text{wmin}$) and the spill threshold ($\text{w0}$). The minimum water stage is not a real a water stage, but is calculated from the regulation volume ($\text{regvol}$) and the dam (surface) area. Thus it does not consider the area changing with depth. Below the minimum water level ($\text{wmin}$) no outflow occur.

For dams a preferred production flow is calculated (see below). This is released if the dam in within the regulation volume, but no more water than to the minimum threshold is released. If instead the water level of the dam is above the threshold $\text{w0}$, the spillway equation (rating curve) is used. The outflow is the maximum of the spillway equation and the preferred production flow.

Input variables $\text{rate}$, $\text{exp}$, $\text{regvol}$, $\text{qamp}$ and $\text{qpha}$ is given in DamData.txt.

General preferred production flow

For all dams the general preferred production flow is calculated from input data on the dam. The flow that is preferred to be released from the dam today is then adjusted depending on purpose. The general production flow ($\text{qprod}$) may be constant or vary between two values per year. This is determined by the input variables $\text{qprod1}$, $\text{qprod2}$, $\text{datum1}$ and $\text{datum2}$. Regulation period 1 between $\text{datum1}$ and $\text{datum2}$ has production flow $\text{qprod1}$, while the rest of the year has production flow $\text{qprod2}$. Not setting the dates gives the same production flow the whole year ($\text{qprod1}$). If $\text{qprod1}$ is not given $\text{qprod}$ is estimated from the yearly average inflow of the dam, which is given as input data.

The production flow may be reduced from this level when there is a low water level in the dam. The flow will be reduced linearly from full production flow for that time of year when water level is over the limit ($\text{limqprod}$ (percentage of volume)) to zero when no water is left in the dam over $\text{wmin}$ level.

1 - Irrigation dam and 2 - Water supply dam

Dams of these purposes uses the general preferred production flow as it is.
3 - Flood control dam

The flood control dam will try to stay empty and ready to delay large inflows. The preferred production flow is calculated from the inflow of the day ($Q_{\text{inftoday}}$) for low inflow and water stages, but set higher than the inflow for high flows and water levels. The maximum flow ($q_{\text{thresh}}$, the threshold flow) is determined from input data ($Q_{\text{inft}}$ is average inflow to dam for each month) and a general parameter ($k_{\text{thr}}$), while the water level threshold ($w_{\text{thres}}$) is determined by a general parameter ($k_{\text{low}}$) for the fraction of the regulation depth.

$$q_{\text{thresh}} = k_{\text{thr}} \times \text{MAX} \left( Q_{\text{inf}} \right)$$

$$w_{\text{thres}} = w_0 \cdot k_{\text{low}} \times \left( w_0 - w_{\text{min}} \right)$$

If the current inflow is larger than the threshold inflow ($q_{\text{thresh}}$), the maximum allowable discharge is released:

$$q_{\text{prod}} = q_{\text{thresh}}$$

If the current inflow is less than the threshold inflow ($q_{\text{thresh}}$) and the current water level is lower than the threshold level ($w_{\text{thresh}}$), the inflow is released from the dam:

$$q_{\text{prod}} = Q_{\text{inftoday}}$$

If the water level is above the threshold ($w_{\text{thresh}}$), more water is released than inflow in an attempt to empty the dam. The factor multiplied by inflow ($k_{\text{rel}}$) is a general parameter:

$$q_{\text{prod}} = \text{MIN} \left( k_{\text{rel}} \times Q_{\text{inftoday}}, q_{\text{thresh}} \right)$$

4 - Hydropower dam

Dams for hydropower uses the general preferred production flow, but it may be adjusted with a seasonal factor. The factor is sinusoidal over the year, with a peak in December, when power output is normally high, and a minimum in June. If you want a different seasonal phase set $q_{\text{pha}}$ to another value (default = 102). The seasonal adjustment is set with the input variable $q_{\text{amp}}$ (>0, <1) for each dam.

$$q_{\text{prod}} = q_{\text{prod}} \times \left( 1 + q_{\text{amp}} \times \sin \left( \frac{2 \times \pi \times \left( \text{dayno} + q_{\text{pha}} \right)}{365} \right) \right)$$

Alternatively the seasonal variation adjustment can be determined by the climate. If this option is used $q_{\text{amp}}$ is set to zero (or left out) in the file, and the value used in the calculations is determined by the fraction of snow fall ($\text{snowfrac}$) for each lake. If the snow fall fraction is above 0.35, a $q_{\text{amp}}$ of 0.71 is used.
Outlet lake with two outlets

An outlet lake that is not part of a lake composed of lakebasins, may have two defined outlets in LakeData.txt. The outlets can be divided into different outlet types depending on method for determining the outflow. HYPE will define an outlet's type based on variables present in LakeData.txt. Below the defining (necessary) variables are given for each outlet type. Note: The threshold (w0ref) is given for first outlet. For the second outlet this variable instead is given relative to the threshold of outlet 1 (i.e usually zero).

Outlet types

1. An outlet with outflow as production flow (only). The production flow can be determined by constants for different seasons or with a sinus adjustment; see Production flow above. Defining variables in LakeData.txt: regvol, (rate=0, maxQprod=0, minflow=0)

2. An outlet with outflow as production flow (same as type 1), but with an allowed (higher) maximum production flow. If the outflow from the other outlet is above zero, part of that flow will be sent through this outlet as part of production flow instead. Defining variables in LakeData.txt: regvol, maxQprod.

3. Rating curve determined outflow with constant threshold. Defining variables in LakeData.txt: rate (deltaw0=0, regvol=0, maxQprod=0)

4. Rating curve determined outflow with constant threshold relative to the threshold of the first outlet. This type is only used for the second outlet. Defining variables in LakeData.txt: rate, w0ref (regvol=0, deltaw0=0)

5. Flow determined by production flow below the threshold, and by a rating curve above the threshold. This could be production and overflow from a hydropower plant that goes too the same downstream subbasin. Defining variables in LakeData.txt: regvol, rate (maxQprod=0, minflow=0)

6. An outlet with outflow as production flow determined by different rating curves for different seasons; see Two rating curves above. Defining variables in LakeData.txt: rate, deltaw0 (regvol=0, maxQprod=0)

7. An outlet with outflow as production flow determined by two rating curves (same as type 6), but with an allowed (higher) maximum production flow. If the outflow from the other outlet is above zero, part of that flow will be sent through this outlet as part of production flow instead. Defining variables in LakeData.txt: rate, deltaw0, maxQprod (regvol=0)

8. An outlet with outflow as minimum flow. The minimum flow is determined same as production flow by constants for different seasons or with a sinus adjustment. Defining variables in LakeData.txt: regvol, minflow (rate=0, maxQprod=0)

9. Minimum flow determined by a production flow below the threshold. In addition overflow is determined by a rating curve above the threshold. This could be a minimum environmental flow together with overflow. Defining variables in LakeData.txt: regvol, rate, minflow (maxQprod=0).

10. An outlet with outflow determined from a time series in Xobs (dwtr). This type is only used for the second outlet. Defining variables in LakeData.txt: obsflow=1
**Combinations of outlet types**

Examples of common combinations of outlets:

- A hydropower plant with an estimated average production flow and a spill overflow at high water levels are simulated by combining production flow (outlet type 1) with rating curve determined overflow (outlet type 3).

- Same as above a hydropower plant with production flow in one branch and overflow in another (outlet type 3), but in this case the maximum allowed production is assumed higher than the average production (outlet type 2). This means that if overflow is calculated due to high water level, part of that flow is lead to the powerplant and through production to the first branch.

- A hydropower plant where the production flow and spill goes to the same downstream subbasin (outlet type 5), but where a second spill outlet goes to another subbasin (outlet type 4).

- A hydropower plant with production flow in one branch (outlet type 1), and a minimum (environmental) flow plus overflow outlet in the old river course that goes to another subbasin (outlet type 9).

- A lake with two outlets with flow determined by a rating curve for each of them (may have different threshold also) are simulated by a combination of outlet type 3 and 4.

Many other combinations can be used, but not all outlet types are natural to combine. The following are **not allowed**:

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<td>3-5, 8, 9</td>
</tr>
<tr>
<td>10</td>
<td>all</td>
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**Change of branched flow after updating total flow**

After calculating the outflow from the two outlets, which is done separately, the total outflow could be higher than the volume above the threshold. In this case the outflow will be reduced. When updating the total outflow from a lake with two outlets against observed total flow, the flow in respective branch has to be adjusted too. Depending on the outlet types this is done by different methods.

Lakes with a clear defined division between production flow and spill branch is handled so that production flow takes priority. For outlets with a maximum production flow, flow are diverted into this branch up to the maximum value and the rest into the other branch. For other lakes with a clear defined division between production flow and spill branch, the same method is applied but with the current production flow as the maximum value. For lakes with a minimum flow in one branch the
minimum flow is given highest priority. Second priority is given to the production flow of the other branch (if any). At last the remaining flow is given to the branch for spill (outlet type 9) or the non minimum flow branch (outlet type 8). For the rest of the outlet type combinations the new flows are changed to be proportional to the old flows.

**Outlet lake (olake) as a lake basin**

An olake can be part of a larger lake. It is then called a lake basin of the larger lake. Lake basins are treated differently if they are located within the lake or if they are the last one, i.e. the one with the lake outflow.

**Inflow**

A lake basin can get local inflow from upstream areas and from the regional groundwater. Inflow works in the same way as for a simple olake (see above).

**Flow between lake basins**

Flow through a lake divided into lake basins occurs only in one direction according to the maindown (and possibly branchdown), no two-way exchange between lake basins is implemented. For upstream lake basins that run into another lake basin, all the water above the threshold is drained. The threshold of the lake (same as last lake basin) is used, not any internal threshold between lake basins. If the lake water level ($w_{lm}$) is higher than the threshold ($w_0$ or $w_{min}$) outflow is obtained.

**Outflow of lake composed of lake basins**

For the last lake basin outflow is estimated by a rating curve, either a specific equation or with the general equation, or by regulation routine. Since all the water (above threshold) in the lake (from all lake basins) accumulates in the last lake basin the water level cannot be used directly, but it is recalculated to calculate the outflow from the lake. The last lake basin water level ($w_{lm}$) is scaled by the subbasin lake area ($area$) to the whole lake area ($lakearea$). If the calculated lake water level ($w_{lake}$) is higher than the threshold ($w_{thresh}$) the outflow is estimated by the rating curve.

\[
\begin{align*}
  w_{lake} &= (w_{lm} - w_{thresh}) \times \frac{area}{lakearea} + w_{thresh} \\
  \text{outflow} &= \text{rate} \times (w_{lake} - w_{thresh})^{\text{exp}}
\end{align*}
\]

The threshold may be changing over the year as described in Section Two rating curves above. If the water level is below the threshold then the outflow is zero or if the production flow regulation is used there is production flow. See Section Production flow above. Production flow is also applied on the recalculated water stage.

**Initialisation of lake volume**

Unless a starting state is given from a file, the lakes start the simulation filled with water to their
outflow threshold. That means for most lakes a water level equal to \textit{lake\_depth}. Dams are filled to the
dam's maximum elevation (equal to \textit{lake\_depth}), except for flood control dams which are initialized
with an empty regulation volume.

For lakes with outflow determined by a rating curve, the water level of the lake will be higher than the
outflow threshold level. The equilibrium level will depend on the size of the inflow and the outflow
rating curve parameters. Depending on the residence time of water in the lake it may take time for
this level to be established, and until then the outflow of the lake will be simulated lower than it
should be. Thus a spin-up time is needed for a model simulation.

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References


Floodplains

Floodplains may be simulated adjacent to a main river or outlet lake. Floodplain is a land area close to the surface water that may be temporary flooded. Water may flow from the surface water to the floodplain and back. The floodplain is a part (fpf) of the classarea of the main river or the lake. The lake/river surface area is the rest of the classarea. The floodplain is flooded when the water in the river or lake reaches above a certain threshold. The water on the floodplain may return to the river lake or may be trapped when the water level of the floodplain sink.

Figure 7: Five possible scenarios of flooding.
Common floodplain processes

The water level of the floodplain and the extent of the flooded area (a\textsubscript{\text{plain}}, m\textsuperscript{2}) are calculated from the volume (v\textsubscript{\text{vol\textsubscript{\text{plain}}}}, m\textsuperscript{3}), maximum area (a\textsubscript{\text{max}}) and water level at maximum areal extent (f\textsubscript{ym}, m). The actual water level will be compared to the equilibrium water level to determine the water exchange. The equilibrium water level is the water level we would have if water was evenly distributed in the lake and the floodplain. It is solved from a second degree equation.

\[
w_l^{\text{plain}} = \sqrt{\frac{v_{\text{vol\textsubscript{\text{plain}}}} \times f_{ym} \times 2}{a_{\text{max}}}}
\]

\[
a_{\text{plain}} = \frac{w_l^{\text{plain}} \times a_{\text{max}}}{f_{ym}}
\]

\[
a_{\text{max}} = f_x \times \text{classarea}
\]

Figure 8: Illustration of floodplain variables of a lake.

The exchange of water between lake (or river) and floodplain is determined by the respective water levels (w\textsubscript{body} and w\textsubscript{plain}), the equilibrium water level (w\textsubscript{equil}) and the thresholds between the water bodies (\textit{h}_{body/\text{plain}}). If the water level is over the threshold and higher than the water level of the floodplain the flow from water body to floodplain is:

\[
q = r_{\text{bodyToPlain}} \times MIN \left( w_{\text{lake}} - f_{body} - w_{\text{equil}}, w_{\text{lake}} - f_{\text{body}} \right)
\]

If the floodplain water level is over the threshold and higher than the water level of the water body the flow from floodplain to body is similarly:

\[
q = r_{\text{plainToBody}} \times MIN \left( w_{\text{plain}} - f_{\text{plain}} - w_{\text{equil}}, w_{\text{plain}} - f_{\text{plain}} \right)
\]

\[
\frac{a_{\text{plain}}}{a_{\text{max}}}
\]

Floodplain water level (w\textsubscript{\text{plain}}) and degree of flooded area (\frac{a_{\text{plain}}}{a_{\text{max}}}) are possible to print out from a simulation with an outlet lake or main river with floodplain.
Simple floodplain model

The simple floodplain model simulates precipitation on the floodplain and evaporation from the flooded water, in addition to the exchange of water between the river or lake and the flood plain. All precipitation on the floodplain is contributing to flooded water.

Floodplain model with soil routines

The floodplain model with soil routines is calculating all soil flows for the non-flooded part of the floodplain. The runoff from the floodplain soil goes to the flooded water, and not to the local stream as it does for ordinary land classes. Precipitation and evapotranspiration is calculated for the flooded water and the non-flooded soil. For the soil part, snow, infiltration and surface runoff, percolation, tile drainage and groundwater runoff is calculated as for an ordinary soil. Infiltration from the flooded water to the soil may occur. Regional groundwater percolation is only calculated for the flooded part together with percolation from the river.

Connected floodplains

It is possible for flooded floodplains to limit the water flow from upstream rivers and lakes. This is determined based on the water levels in the floodplains, a higher water level of a downstream floodplain compared to a upstream floodplain may reduced the flow from the upstream compartment (main river or outlet lake) to the downstream compartment (main river or outlet lake). The damming flow is calculated as the flow to reach equilibrium water level between the two floodplains. The relative level of the two floodplains location is determined based on the classes average elevation and thresholds or given as indata separately. The function of connecting floodplains works within a subbasin or between connected subbasins. This floodplain features may be used together with the Floodplain model with soil routines.

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**Bifurcations**

Outflow from a subbasin may flow in one or two directions. The main flow follows the main channel, which is the downstream subbasin given by the path in GeoData.txt. If there is a bifurcation, the branch flow goes to another downstream subbasin (with subid branchid). Any of the flows may go outside the model set-ups area, they are then no longer a concern of the model.

There are three ways to determine the flow in the different channels. 1) The division of the total outflow is determined in BranchData.txt. 2) The outflow is calculated for two outlets separately and then divided into the channels. For the second alternative see Outlet lake with two outlets section above. 3) The demanded flow in the branch is prescribed (Qbranch). If the total flow is not enough to fulfill the need, less water goes into the branch. The main channel gets the rest of the total flow.

For the first method the total outflow has been calculated the ordinary way. The flows division into the two channels is determined by four parameters; mainpart, maxQmain, minQmain and maxQbranch, which are set in BranchData.txt. Zero values of the input data mean they are not used. The main flow (mainflow) is calculated from the totalflow (q) as:
main flow = \begin{cases} 
q_{\text{main part}} x \left(q - minQ_{\text{main}}\right) + minQ_{\text{main}} & \text{if } q < minQ_{\text{main}} \\
maxQ_{\text{main}} & \text{if } minQ_{\text{main}} < q < q_{\text{thresh}} \\
q - maxQ_{\text{branch}} & \text{if } q > q_{\text{thresh}}, q_{\text{thresh}} = \frac{maxQ_{\text{main}} - minQ_{\text{main}}}{\text{main part}} + minQ_{\text{main}} \\
0 & \text{if } q > q_{\text{thresh}}, q_{\text{thresh}} = \frac{maxQ_{\text{branch}}}{1 - \text{main part}} + minQ_{\text{main}} 
\end{cases}

q_{\text{thresh}} = \text{MIN} \left( \frac{maxQ_{\text{main}} - minQ_{\text{main}}}{\text{main part}} + minQ_{\text{main}}, \frac{maxQ_{\text{branch}}}{1 - \text{main part}} + minQ_{\text{main}}, 0 \right)

The rest of the total flow goes in the branch. The following figures give some examples of parameter combinations.

- Serie 1: main part = 0.5, minQmain = 2, maxQmain = 10, maxQbranch = 0
- Serie 2: main part = 0.5, minQmain = 2, maxQmain = 0, maxQbranch = 5
- Serie 3: main part = 0.5, minQmain = 0, maxQmain = 10, maxQbranch = 0
- Serie 4: main part = 0, minQmain = 2, maxQmain = 0, maxQbranch = 5

Figure 5: Main flow in relation to total flow for four different parameter combinations.

Figure 6: Branch flow in relation to total flow for four different parameter combinations.
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Nitrogen and phosphorus in land routines

HYPE simulate nitrogen and phosphorus in soil both as immobile soil pools and as fractions dissolved in soil water. The pools are changed by outside sources and internal soil processes. The following fractions are simulated:

- **fastN** - immobile pool of organic nitrogen in the soil with rapid turnover ($kg/km^2 = mg/m^2$)
- **fastP** - immobile pool of organic phosphorus in the soil with rapid turnover ($kg/km^2 = mg/m^2$)
- **humusN** - immobile pool of organic nitrogen in the soil with slow turnover ($kg/km^2 = mg/m^2$)
- **humusP** - immobile pool of organic phosphorus in the soil with slow turnover ($kg/km^2 = mg/m^2$)
- **IN** - concentration of inorganic nitrogen in soil water ($mg/L$)
- **ON** - the concentration of (dissolved) organic nitrogen in soil water ($mg/L$)
- **partP** - immobile pool of inorganic phosphorus adsorbed to soil particles ($kg/km^2 = mg/m^2$)
- **PP** - concentration of particulate phosphorus (of organic origin) in soil water ($mg/L$)
- **SP** - concentration of soluble (reactive) phosphorus in the soil water ($mg/L$)

The figure below give an overview of the nitrogen and phosphorus in soil in HYPE. The internal processes are shown with broad arrows and described in the section **Nutrient soil processes** below. Sources and sinks are shown with arrows and text, and are further described in sections **Nutrient sources** and **Vegetation and soil surface processes**.

Figure 1: Sources and sinks of nitrogen and phosphorus in the soil. The humusN and humusP pools are called slowN and slowP in this figure.
Nutrient Sources

Fertilizer

Fertilization can be specified to be applied up to two times per year for each of the two fertilization forms, (commercial) fertilizer (fert) and manure (man). For each fertilization event the application date is specified, the amount of N and P (fert(inut,ife), man(inut,ife), where inut=N,P and ife=1,2 the fertilization event) to be added, and a management indicator (fdown(ife), mdown(ife)). The last information indicates how much of the fertiliser is to be added to the second soil layer (the rest is added to the first soil layer). These parameters are given in CropData.txt and are crop and region dependent. For manure 50% of the nutrients are assumed to be in organic form (inorgpart = 0.5).

Fertilizer and the inorganic part of the manure are added to the IN and SP pools of the soil water. The total amount is spread out over a specified period (given by the general parameter fertdays) after the date of fertilization. An equal amount is applied each day for the specified period. The organic part of the manure is added to the fastN and fastP pools.

Each class can have two crops (icrop=1,2), the main crop and the secondary crop. The secondary crop do not have to cover (a) the whole class, e.g. if only 50% of the spring wheat (main crop) area have catch crop (second crop) (a(1)=1, a(2)=0.5). The total amount of added nutrients (common add) is calculated as the sum of fertiliser additions from the main and secondary crops.

Inorganic nutrients to be added to IN and SP pools:

\[
\text{common add}_{\text{inut}, \text{layer}=1} = \sum_{\text{icrop}=1}^{2} \sum_{\text{ife}=1}^{2} a_{\text{icrop}} x \left( \text{fert}_{\text{inut}, \text{ife}} \times \left( 1 - \text{fdown}_{\text{ife}} \right) \right) \times \left( \text{fertdays} \right)
\]

Organic nutrients to be added to fastN and fastP pools:

\[
\text{common add}_{\text{inut}, \text{layer}=1} = \sum_{\text{icrop}=1}^{2} \sum_{\text{ife}=1}^{2} a_{\text{icrop}} x \left( \text{man}_{\text{inut}, \text{ife}} \times \left( 1 - \text{mdown}_{\text{ife}} \right) \right) \times \left( 1 - \text{inorgpart} \right) \times \left( \text{fertdays} \right)
\]
Note that the terms for application of fertilizer and manure only are added during the active time, which is from the application date and for fertdays number of days thereafter.

**Plant residues**

Plant residues are added at a specified date each year. They are added to the immobile soil pools (fastN, fastP, humusN, and humusP) to get a delay of the leakage and an accumulation of humus in the soil. The amount of crop residues (resamount(inut), inut=N,P) and the date they are added on depends on crop and region and is given in the CropData.txt file. Part of the plant residues are added to the fast pool (resfast), and part to the humus pool. The plant residues are added to the upper two soil layers with a distribution (resdown) also specified in CropData. For each class the plant residues are weighted (common_res) for main and secondary crop together every day with the help of share for each crop (a(icrop), where icrop=1,2).

\[
\text{common}_{\text{res}}(\text{fast-inut}, \text{layer=1}) = \sum_{\text{icrop}=1}^{2} a(\text{icrop}) \times \text{resfast}(\text{icrop}) \times \text{resamount}(\text{inut, icrop}) \times \left(1 - \text{resdown}(\text{icrop})\right)
\]

\[
\text{common}_{\text{res}}(\text{fast-inut}, \text{layer=2}) = \sum_{\text{icrop}=1}^{2} a(\text{icrop}) \times \text{resfast}(\text{icrop}) \times \text{resamount}(\text{inut, icrop}) \times \text{resdown}(\text{icrop})
\]

\[
\text{common}_{\text{res}}(\text{humus-inut}, \text{layer=1}) = \sum_{\text{icrop}=1}^{2} a(\text{icrop}) \times \left(1 - \text{resfast}(\text{icrop})\right) \times \text{resamount}(\text{inut, icrop}) \times \left(1 - \text{resdown}(\text{icrop})\right)
\]

\[
\text{common}_{\text{res}}(\text{humus-inut}, \text{layer=2}) = \sum_{\text{icrop}=1}^{2} a(\text{icrop}) \times \left(1 - \text{resfast}(\text{icrop})\right) \times \text{resamount}(\text{inut, icrop}) \times \text{resdown}(\text{icrop})
\]

All crop residues are transferred into the soil at a given date as determined by the user in CropData.txt. Note that the terms for application of residues only are added that day.

**Rural household diffuse source**

Information on rural household diffuse source (private sewers) is located in the file GeoData.txt. The source has a flow and concentrations of total nitrogen and phosphorus and fraction of IN and SP. The diffuse source is divided into two parts, where the distribution is determined by a general parameter (locsoil). One part is added directly to the local river. The other part is added to soil water in the bottom soil layers in the catchment area of all land classes with a distribution proportional to the classes’ area.
Vegetation and soil surface processes

The CropData.txt file contains information on the various crops that are simulated. Each soil type-land use class (SLC-class) can have a main crop and a secondary crop. To reduce the number of SLC-classes in a simulation the individual crops with similar properties are brought together in a grouped crop (e.g. row crops). In CropData.txt there is information such as the amount of fertilizer, manure and crop residues introduced into the soil for different crops in different regions. The file also provides coefficients for nutrient uptake capabilities.

Potential vegetation uptake of nitrogen

The potential uptake is used to calculate the absorption of IN and SP from the soil water (see section Vegetation nutrient uptake). The actual uptake will be limited by the nutrients available. Potential uptake of nitrogen by plants is based on a three-parameter equation (logistic growth) from SOILN (Eckersten et al., 1994). Nitrogen uptake \( \text{uptake} \) is calculated for the main and secondary crop and added to the common potential absorption function \( \text{common_uptake}, \text{mg/m2/d} \) using the percentage of the area of the main crop \( \text{part} = 1.0 \) and secondary crop \( \text{part} < 1 \). The potential uptake is divided between the upper soil layer and second soil layer by a crop dependent input variable for the fraction of the upper soil layer. If the crop is growing in a class of only one soil layer, only the upper soil layer fraction of the potential uptake will be used in the calculations. The growing period with simulated nutrient uptake is determined by the sowing date \( \text{bd2} \) and continues until harvest \( \text{bd3} \). Potential
uptake depends on three parameters (up1, up2 and up3) and the number of days after sowing date. The uptake for a specific crop is calculated with the parameter values for that crop as

\[ h_{elp} = \left( up1 - up2 \right) \times \frac{1}{c} \times \left( dayno - bd2 \right) \]

\[ uptake = \frac{up1 \times up2 \times up3 \times h_{elp}}{\left( up2 + h_{elp} \right) \times \left( up2 + h_{elp} \right)} \]

for the growing period. Outside the growing period the uptake is assumed to be zero.

Autumn-sown crops may take up IN and SP for a while after sowing in autumn. The same potential uptake of nitrogen as the main growing season are used, but uptake is limited by a temperature function. This uptake will run from the autumn sowing date (bd5) to the mid winter (31 December or 30 June depending on the autumn sawing date).

\[ h_{elp} = \left( up1 - up2 \right) \times \frac{1}{c} \times \left( dayno - bd5 + 25 \right) \]

\[ uptake = \frac{tmpfcn \times up1 \times up2 \times up3 \times h_{elp}}{up2 + h_{elp}} \]

where the temperature function (tmpfcn) is calculated as

\[ tmpfcn = \text{delim} \begin{cases} MIN\left[1,\left(\frac{T-5}{20}\right)\right] & T > 5 \\ 0 & T < 5 \end{cases} \]
The growing period will then continue next season from \( bd2 \) as described above.

The sowing date (\( bd2 \)) may be given as a constant, or calculated dynamically depending on temperature. If it is dynamically determined it is set to the first day of the year which has a degreeday sum (\( GDD \)) above a threshold (\( gddsow \)). The degreeday sum (\( GDD \)) is calculated as

\[
GDD(d+1) = GDD(d) + \max\left(0, T - \text{basetemp}\right)
\]

where \( d \) is day of year, \( T \) is air temperature (degree Celcius), \( \text{basetemp} \) is a temperature threshold. The \( GDD \) is accumulated for each day after \( \text{firstday} \) with day length larger than \( \text{daylength} \). The \( GDD \) is zeroed at \( \text{firstday} \).

### Soil erosion

The erosion of soil particles is calculated by HYPE for transport of particulate phosphorus (PP) or for the simulation of sediment material (SS). For transport of PP an erosion model depending on mobilization of particles by rain and by surface runoff is often used. This formulation is the default model for sediment transport, but another erosion model based on catchment erosion index can also be used.

In the first model PP can leave the land by two means, by surface runoff transport and by macropore flow through drainage pipes. The calculation of PP transport is done in three steps: first the erosion (mobilization) of soil particles from the land surface is calculated, secondly how much of the mobilized particles that are leaving the field is calculated, finally the amount of soil particles is converted to phosphorus.

Mobilization (\( \text{MobilisedRain} \) and \( \text{MobilisedSR} \)) occurs due to energy from falling raindrops (\( \text{Rain\_fall\_energy} \)) and/or from surface runoff. The kinetic energy in rainfall is calculated as a function of rainfall and day of the year (\( \text{dayno} \)). If the precipitation falls as snow, or if it falls on snow-covered ground or if it is smaller than 5 mm/day no mobilization occurs in the model. Some of the rain drop’s energy can be absorbed by vegetation. Cropcover is defined as the portion of land that is sheltered from raindrops; for a description of how this is calculated, see next chapter. The factor \( \text{common\_cropcover} \) is the total fraction of cover that the main and secondary crop give together. It varies over the year due to crop growth and management. The mobilization is also influenced by soil erodibility (soil dependent parameter \( \text{soilerod} \)).

\[
\text{Rain\_fall\_energy} = \text{prec} \times \left(\frac{8.95 + 8.44 \times \log_{10}(\text{prec} \times 2 \times \left(0.257 + 0.09 \times \sin\left(2\pi \times \left(\frac{\text{dayno} - 70}{365}\right)\right)\right))}{\text{soilerod}}\right)
\]

\[
\text{MobilisedRain} = \text{Rain\_fall\_energy} \times \left(1 - \text{common\_cropcover}\right) \times \text{soilerod}
\]

When surface runoff occurs, soil particles are eroded and carried away as the soil surface is exposed to shear forces. The mobilization is calculated from the surface runoff (\( \text{sflow} \)), land slope, a parameter for soil cohesion (\( \text{soilcoh} \) (kPa) soil type dependent), and a general parameter (\( \text{sreroexp} \)). This type of
erosion can be mitigated by protective vegetation or vegetation residues that are in contact with the ground. The calculation of this factor (groundcover) is described in next chapter. The variable common groundcover is the total fraction of cover that the main and secondary crop give.

\[
\text{Mobilised}_{SR} = \frac{(s_{\text{flow}} \times 365)^{\text{erosexp}} \times \left(1 - \text{common groundcover}\right) \times \frac{1}{0.5 \times \text{soilcob}} \times \sin \left(\frac{\text{slope}}{100}\right)}{365}
\]

All mobilised particles is not removed from the field, because sometimes the transport capacity of the particle-bearing water (eflow) will not suffice for the task. A transportfactor will reduce the particle amount:

\[
\text{transportfactor} = \text{MIN} \left(1.0, \left(\frac{\text{eflow}}{4}\right)^{ \text{1.3}} \right)
\]

Mobilised sediment (kg/km2) is calculated as:

\[
\text{mobilSed} = 1000 \times \left(\text{Mobilised}_{\text{Rain}} + \text{Mobilised}_{SR} \times \text{transportfactor} \right)
\]

The alternative erosion model calculates mobilised sediment (mobilSed (kg/km2)) based on rainfall (rain) and a number of model parameters and subbasin input data.

\[
\text{mobilSed} = 1000 \times \left(\frac{\text{slope}}{5}\right)^{\text{erodslope}} \times \text{erodluse} \times \text{erodsoil} \times \frac{\text{EI}}{\text{erodindex}} \times \text{rain}^{\text{erodexp}}
\]

The parameters erodslope, erodexp and erodindex are general. The parameters erodluse and erodsoil are land-use and soil type dependent. Subbasin input is needed on slope, the subbasins average slope, and an erosion index, EI.

A selective process is affecting the soil erosion of phosphorus. Smaller and lighter particles are eroded easier than larger ones. The tiny particles contain more P per unit weight than the average particle of the soil. Therefore an enrichment factor (enrichment) is used. The enrichment factor is calculated from three parameters (ppenrmax, ppenrstab, ppenrflow), one of which is soil type dependent (ppenrmax), and the particle bearing flow. Typical values of the parameters, here called max, stab and flowstab, are given in the example in Figure 3.
Eventually the mobilised PP (kg/km2) is calculated as:

$$mobilP = 0.000001 \times mobilScd \times PartP + HumusP \times \frac{thickness \times bulkdensity}{enrichment}$$

The calculated mobilised phosphorus ($mobilP$) is now available to add to the particulate phosphorus (PP) of surface runoff.

A portion of PP in surface runoff ($sflow$) is filtered out (for example by buffer zones). Filtering ($srfilt$) is determined by land use dependent parameters ($bufferfilt$, $innerfilt$, $otherfilt$), percentage of agricultural land close to a watercourse ($alfa$), and proportion of agricultural land near the rivers which have a protective buffer zone ($bufferpart$).

$$srfilt = otherfilt + alfa \times \left(1 + bufferpart \times (bufferfilt - 1)\right) + innerfilt \times (1 - alfa)$$

Similarly, part of the PP which is transported by macropore flow ($mflow$) is filtered away between the soil surface and drainage pipes. The parameter that determines this effect ($macrofilt$) is soil dependent. The PP transported ($erodedP$) by surface runoff and macropore flow is collected in a temporary storage pool ($relpool$ (kg/km2)) together with PP in tile runoff ($tilep$).

$$erodedP = \frac{srfilt \times sflow + macrofilt \times mflow}{eflow} \times mobilP$$

$$eflow = sflow + mflow$$

$$relpool = relpool + erodedP + tilep$$

From the temporary pool phosphorus is released ($release$ (kg/km2)) and then transported to the local river depending on the size of the total runoff ($runoff$ (mm)). The parameters $pprelmax$ and $pprelexp$ are general.

$$release = MIN \left( relpool, relpool \times (runoff / pprelmax)^{pprelexp} \right)$$
Crop cover and ground cover

Erosion can be mitigated by protective vegetation or vegetation residues that are in contact with the ground. Crop cover and ground cover reduce erosion by rain and surface runoff for particulate phosphorus (PP) and the default sediment transport model (SS). Each crop covers a fraction of the ground, thus for simultaneous crops (1st and 2nd crop) their respective crop/ground cover is combined to a common crop/ground cover.

Harvested crops have seasonally varying ground and crop cover, while permanent vegetation (e.g. forest) has constant values for these parameters. The input data needed for calculations (ccmax1, ccmax2, gcmax1, gcmax2) are given in CropData.txt. Parameters ccmax1 and gcmax1 describe the maximum crop and ground cover during spring-summer growing season, parameters ccmax2 and gcmax2 are corresponding maxima for winter crop’s growth. These maximum ratios are reached at maxday1 and maxday2, which are defined as halfway between planting and harvest, and halfway between autumn planting and midwinter (1 January or 30 June depending on autumn planting date), respectively. From sowing the coverage fractions increase linearly up to their maximum values. After these dates maximum coverage is maintained to the next ploughing, harvest, or until the growing season starts again in the spring (for winter crops) (Figure 4). During the period between harvesting and ploughing, crop cover is equal to ground cover (gcmax1). At ploughing, ground and crop cover are reduced to zero. Parameters bd1 and bd4 describe the dates of spring and autumn ploughing. In the case of spring sowing, when no winter crop is crowing, either one of the ploughing parameters can be set for ploughing date.
Transformation of nitrogen from atmospheric deposition

The parameter $ponatm$ indicates that some of the nitrogen deposition will be organic instead of only inorganic. For wet deposition and deposition on snow, this is adjusted when there is infiltration of rain and melted snow into the soil. A portion ($ponatm$) of IN in the infiltration are added to fastN in the top layer instead of following the water to the soil water.

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**Nutrient soil processes**

**Soil pools - initial values**

The initial pools of immobile soil nutrients are dependent on land use and is determined by the user and very important because they are changing slowly. The initial pools of the soil layers are calculated from a parameters. The parameters (humusn0, humusp0, partp0, fastn0, fastp0) are representative for the uppermost soil layer; the unit for these parameters is mg/m³. The initial humusN, humusP and partP-pools decreases with depth, with half depth determined by land use dependent parameters (hnhalf, hphalf and pphalf).

The equation for a depth dependent soil nutrient is:

\[ \text{nut}(d) = \text{nut}0 \times e^{-\text{par} \times d} \]

where \( d \) is depth below zero-level, \( \text{nut}0 \) is nutrient at zero-level (i.e. model parameter for uppermost soil layer) and \( \text{par} \) is calculated from respective nutrient half depth model parameter (NNhalf) as:

\[ \text{par} = \frac{\log 2}{\text{NNhalf}} \]
As the model parameters that represent zero-level nutrient \((nut0)\) is defined for the uppermost soil layer, the zero-level is the middle of uppermost soil layer. The following depths \((d)\) represent the soil layers (1-3):

\[
d(1) = 0
\]

\[
d(2) = \frac{soilthick(1)}{2} + \frac{soilthick(2)}{2}
\]

\[
d(3) = \frac{soilthick(1)}{2} + \frac{soilthick(2)}{2} + \frac{soilthick(3)}{2}
\]

The nutrient is transformed into model soil nutrient pools (unit \(kg/km^2\)) by taking into account the thickness of the soil layers.

\[
pool(k) = nut(d(k)) \times soilthick(k), \quad k = 1..3
\]

Pools with rapid turnover \((fastN, fastP)\) have initial values independent of depth.

\[
pool(k) = nut_{0} \times soilthick(k), \quad k = 1..3
\]

The nutrients dissolved in soil water is initialized independent of depth. The initial IN and SP concentrations are zero. The initial ON and PP concentration are determined by land use dependent parameters \((onconc0\) and \(ppconc0, mg/L\) - valid for all three soil layers.

**Common functions**

Many soil processes depend on temperature and soil moisture. The following equations are used in these cases. The temperature function (figure 5) depends on the estimated soil layer temperature \((soiltemp)\). The soil temperature requires some parameters to be simulated, see Section Soil temperature and snow depth.

\[
tmpfcn = 2**((soiltemp - 20.0) / 10.0)
\]

\[
IF(temp < 5.0) \quad tmpfcn = tmpfcn \times (soiltemp / 5.0)
\]

\[
IF(temp < 0.0) \quad tmpfcn = 0.0
\]

The humidity function (figure 6) depends on soil moisture \((soil)\) in the soil layer and the parameters of wilting limit \((wp)\), field capacity \((fc)\) and effective porosity \((ep)\) transformed to unit \(mm\). All these humidities are specified as percentages. The function includes coefficients \(thetaupp = 0.12\), \(thetalow = 0.08\), \(thetapow = 1.0\) and \(satact = 0.6\). Note that another function is used in the calculation of denitrification. For soil layers \(k = 1..3\) the equation is:

\[
IF(soil >= wp + fc + ep) \quad THEN
\]

\[
smfcn = satact
\]

ELSE
smfcn = min(1.,(1-satact)*((wp+fc+ep-soil) / (thetaupp*soilthick)) **thetapow + satact,((soil-wp)/(thetalow*soilthick)) **thetapow)
ENDIF
IF(soil < wp) THEN
  smfcn=0.0
ENDIF

The humidity function (figure 6) is always less than or equal to one while the temperature function may be greater than one when the temperature exceeds 20 degrees.

Vegetation nutrient uptake

Vegetation nutrient uptake is a sink of the IN and SP concentrations in the two upper soil layers. The uptake (uptake / uptakeP, mg/m2/d) depends on the potential uptake of nitrogen \( \text{uptake}^{\text{common}} \), calculated for the combination of main crop and any secondary crop of the specified class. Vegetation nutrient uptake also depends on uptsoil1 which gives the distribution of uptake between the two layers for a crop, the pool of IN (INpool) and SP (SPpool) and for phosphorus by P / N ratio (pnratio). The pool of dissolved nutrients are calculated as the concentration multiplied by the water in the soil layer (in mm). The main and secondary crop of a class is specified in GeoClass.txt.

\[
\text{uptake}N(1) = \text{MIN} \left( \frac{\text{uptsoil1} \times \text{uptake}^{\text{common}}}{a \times \text{INpool}(1)} \right)
\]
The plants can take up a maximum factor $a$ of the pool. This factor is calculated as a percentage of soil water over the wilting point. The equations for the potential uptake of nitrogen ($\text{uptake}_{\text{N}}(2)$) are described under that section above. The constants $\text{uptsoil1}$ (percentage of plant nutrients absorbed top layer) and $\text{pnratio}$ (ratio of N and P uptake) are given for each crop.

### Denitrification

Denitrification is a sink for IN in all three soil layers. Denitrification ($\text{denitr}, \text{mg/m}^2/\text{d}$) depends on the denitrification rate and the pool of IN ($\text{INpool}$) in each soil layer ($k = 1.3$). The denitrification rate depends on a rate coefficient ($\text{drate}$), soil temperature, IN concentration and soil moisture.

$$\text{denitr}(k) = \text{drate} \times \text{INpool}(k) \times \text{tmpfcn}(k) \times \text{smfcn}(k) \times \text{concfcn}(k)$$

The coefficient $\text{drate}$ is determined by land use dependent model parameters $\text{denitrlu}$ and $\text{denitrlu3}$. The temperature dependence ($\text{tmpfcn}$) is described above. The soil moisture function (figure 7) is exponential and thus different from the general soil moisture function.

$$\text{smfcn} = \begin{cases} 0, & \text{soil} < \text{pw} \times \text{lim} \\ \exp\left(\frac{\text{soil}}{\text{pw} \times \text{dlim}}\right), & \text{else} \end{cases}$$

where $\text{pw} = \text{wp} + \text{fc} + \text{ep}$

The function depends on soil moisture ($\text{soil}$) and pore volume ($\text{pw}$). It also depends on two constants; the limit where moisture is high enough to allow denitrification to occur ($\text{dlim} = 0.7$) and the exponent ($\text{exp} = 2.5$). These cannot currently be changed. The dependence of the denitrification rate on the IN concentration is described by a function with a half-saturation concentration (general parameter $\text{hsatINs}$ was in earlier HYPE versions a constant equal to 1 mg/L) (Figure 8).

$$\text{concfcn} = \frac{\text{conc}}{\text{conc} + \text{hsatINs}}$$
Immobile soil nutrient pool transformations

Turnover of fastN

Turnover of fastN is a sink for fastN and a source of IN in all three soil layers (k = 1..3). Turnover \((transfN, \text{mg/m}^2/\text{d})\) depends on a general parameter \((\text{minerfn})\), temperature function \((\text{tmpfcn})\), humidity function \((\text{smfcn})\) and the pool of fastN \((\text{fastN})\).

\[
transfN(k) = \text{minerfn} \times \text{tmpfcn}(k) \times \text{smfcn}(k) \times \text{fastN}(k) 
\]

Turnover of fastP

Turnover of fastP is a sink for fastP and a source of SP in all three soil layers \((k = 1..3)\). Turnover \((transfP, \text{mg/m}^2/\text{d})\) depends on a general parameter \((\text{minerfp})\), temperature function \((\text{tmpfcn})\), humidity function \((\text{smfcn})\) and the pool of fastP \((\text{fastP})\).

\[
transfP(k) = \text{minerfp} \times \text{tmpfcn}(k) \times \text{smfcn}(k) \times \text{fastP}(k) 
\]
Turnover of humusN

Turnover of humusN is a sink for humusN and a source of fastN in all three soil layers \((k = 1..3)\). It is supposed to be a very slow process. Turnover \((\text{degradhN}, \text{mg/m2/d})\) depends on a general parameter \((\text{degradhn})\), temperature function \((\text{tmpfcn})\), humidity function \((\text{smfcn})\) and the pool of humusN \((\text{humusN})\).

\[
d_{\text{degradhN}}(k) = \text{degradhn} \times \text{tmpfcn}(k) \times \text{smfcn}(k) \times \text{humusN}(k)
\]

Turnover of humusP

Turnover of humusP is a sink for humusP and a source of fastP all three soil layers \((k = 1..3)\). It is supposed to be a very slow process. Turnover \((\text{degradhP}, \text{mg/m2/d})\) depends on a general parameter \((\text{degradhp})\), temperature function \((\text{tmpfcn})\), humidity function \((\text{smfcn})\) and the pool of humusP \((\text{humusP})\).

\[
d_{\text{degradhP}}(k) = \text{degradhp} \times \text{tmpfcn}(k) \times \text{smfcn}(k) \times \text{humusP}(k)
\]

Dissolution of soil organic pools

All organic soil pools of nutrient (humusN, fastN, humusP and fastP) contribute to dissolved organic nutrients in soil water (i.e. ON and PP). This process is a sink of the soil pools and a source of ON and PP in all three soil layers \((k = 1..3)\). Dissolution \((\text{mg/m2/d})\) depends on a model parameter (one for each soil pool, \(\text{dispar}\)), temperature function \((\text{tmpfcn})\), humidity function \((\text{smfcn})\) and the size of the soil pool \((\text{soilpool})\).

\[
d_{\text{dissolution}}(k) = \text{dispar} \times \text{tmpfcn}(k) \times \text{smfcn}(k) \times \text{soilpool}(k)
\]

The dissolution model parameters \((\text{dispar})\) is land use dependent and named \(\text{dissolfp}, \text{dissolfn}, \text{dissolhp}, \text{and dissolhn}\) for dissolution of nutrient pools fastN, fastP, humusN and humusP respectively.

Percolation

ON and PP concentration are reduced as the water percolates down through the soil layers. The decrease in concentration depends on land use dependent model parameters \((\text{par})\).

\[
\text{conc} = \text{conc} \times (1 - \text{par})
\]

Balance SP - PartP

SP is in a dynamic equilibrium with P adsorbed to soil particles \((\text{partP})\). The equilibrium is described by the Freundlich equation. From the total amount of P \((\text{SP} + \text{partP})\) in the soil an equilibrium concentration of SP is calculated in the soil solution using the iterative Newton-Raphson method.
coeff = Kfr * bulkdensity * soilthick

DO WHILE
  fxn = xn * vol + coeff * (xn ** Nfr) - totalP
  fprimxn = vol + Nfr * coeff * ( xn **(Nfr-1))
  xn_1 = xn - fxn / fprimxn
  xn = xn_1
ENDDO

The coefficients $K_{fr}$, and $N_{fr}$ are soil dependent, bulkdensity is the dry density of soil (= 1300 kg/m3) and soilthick is the soil layer thickness. $fxn$ is function value for the concentration $xn$, $fprimxn$ is the function value of the derivative of $fxn$, and $x_{n+1}$ is the concentration of the next iteration. The resulting SP equilibrium concentration ($x_{n+1}$) is used to calculate the PP equilibrium concentration ($SP_{n+1}$). A new SP concentration ($SP_{n}$) is calculated from the previous concentration ($SP_{n-1}$) and the equilibrium concentration, taking into account the balance is not reached during the time step.

$$PP_{equi} = K_{fr} \times \left( x_{n+1}^{N_{fr}} \right)$$

$$AdsDes = \left( PP_{equi} - concSolid \right) \times \left( 1 - e^{-K_{ads}} \right)$$

$$SP_{conc} = SP_{conc_{old}} \left( AdsDes \times bulkdensity \times soilthick \right)$$

$AdsDes$ is the amount of adsorbed/desorbed P, $KadsDes$ is a soil dependent rate constant, $equi_{concSolid}$ is the solid phase equilibrium concentration, and $concSolid$ is the initial concentration the solid phase.

**Crop rotation effect on soil pools**

The soil pools of nutrients are shared by crops on rotation on a field. This is modelled by averaging the soil pools once per year for the classes which are in the same crop rotation group.

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Transport with soil water

IN, ON, SP and PP comes with water when it flows through the soil, but remain when water evaporates. Different flow paths contribute to the runoff of water and concentration of IN, ON, SP and PP from the corresponding soil layer. The flow paths possible today are: surface runoff (soil layer 1), runoff (soil layer 1, 2 and 3) and runoff through drainage pipe (soil layer 1, 2 or 3).

Diagnostic output variables of soil nutrients

Some additional output variables are calculated from the soil variables. Gross and net soil load is calculated for the “root zone”, e.g. soil layer one and two together, and for soil layer three separate. A third option for soil load is calculated not strictly based on the soil layers but as an approximation for “groundwater” based on third soil layer but manipulated with tile drainage.
Gross soil loads

Gross load to soil layer one and two (root zone). Components included: atmospheric deposition (wet and dry), fertilizers, crop residues, rural load if number of soil layers is less than three. HYPE variable IDs: sl01=IN, sl03=ON, sl05=sl01+sl03=TN, sl07=SP, sl09=PP, sl11=sl07+sl09=TP

Gross load to soil layer three (lower soil). Components included: percolation from soil layer two to three, rural load if number of soil layers is three. HYPE variable IDs: sl13=IN, sl15=ON, sl17=sl13+sl15=TN, sl19=SP, sl21=PP, sl23=sl19+sl21=TP

Gross load to “groundwater” (modified soil layer three). Components included: percolation from soil layer two to three, rural load if number of soil layers is three, tile drain runoff if tile is in soil layer one or two. HYPE variable IDs: sl25=IN, sl27=ON, sl29=sl25+sl27=TN, sl31=SP, sl33=PP, sl35=sl31+sl33=TP

Net soil loads

Net load from soil layer one and two (root zone). Components included: percolation from soil layer two to three, runoff from soil layer one and two, tile drain runoff if tile is in soil layer one or two, surface runoff. HYPE variable IDs: sl02=IN, sl04=ON, sl06=sl02+sl04=TN, sl08=SP, sl10=PP, sl12=sl08+sl10=TP

Net load from soil layer three (lower soil). Components included: runoff from soil layer three, tile drain runoff if tile is in soil layer three. HYPE variable IDs: sl14=IN, sl16=ON, sl18=sl14+sl16=TN, sl20=SP, sl22=PP, sl24=sl20+sl22=TP

Net load from “groundwater” (modified soil layer three). Components included: runoff from soil layer three, tile drain runoff if tile is in any soil layer. HYPE variable IDs: sl26=IN, sl28=ON, sl30=sl26+sl28=TN, sl32=SP, sl34=PP, sl36=sl32+sl34=TP

Figure 9: Components of calculated gross (brown) and net (green) loads of soil.
Nutrient soil leakage from outer source

There is a model option to skip calculating nutrients with HYPE and instead use nutrient soil leakage from an outer source. In this case the calculations above is skipped, while soil water and runoff processes is calculated as usual. HYPE output variables related to soil nutrients are set to missing values.

The first implemented soil leakage model was

1. monthly seasonal variation of concentrations for each subbasin

The soil leakage is introduced into the model as inorganic and organic nitrogen, and soluble reactive phosphorus and particulate phosphorus concentrations of the total runoff. Different possibilities exist on how to introduce these regarding time variations (monthly seasonal variation, daily), spatial variations (subbasin, regions) and land use dependence (class groups). So far the only implemented alternative is the monthly seasonal variation.

The nutrient concentration of total runoff (HYPE variable ID crun) is set, and then used for all further calculations in the streams and lakes.

The default is to use HYPE calculated runoff concentrations with the processes described above (modeloption soilleakage 0).

References

Nitrogen and phosphorus processes in rivers and lakes

HYPE simulates concentration of inorganic nitrogen (IN), organic nitrogen (ON), soluble reactive phosphorus (SP) and particulate phosphorus (PP) in discharge and other surface waters. As output also the total nitrogen (TN) and phosphorus (TP) concentration can be had. In addition total suspended sediments (TS) can be simulated. It consists of the sum of suspended sediments (SS) and algae (AE) simulated concentrations.

Basic assumptions

Transformations of nutrients take place in lakes and rivers. For lakes, which are divided into fast (FLP) and slow (SLP) lake parts, the process is performed only in SLP (Fig. 1). For rivers, which hold delayed water in a queue and in the damping box, the processes is performed only in the damping box.

![Figure 1: Nutrient flows in a lake that is affected by nutrient processes.](image)

The processes of denitrification, primary production and mineralization have been implemented for both rivers and lakes. For particulate phosphorus (PP) there is an exchange with the river sediments. The rivers dimensions are used in the calculation of these processes. The width and depth of the watercourse are calculated from a number of empirical equations (for more information on these equations see “Modelling phosphorus transport and retention in river networks” by Jörgen Rosberg).

\[
velocity = 10^{vel_1} \times meanflow^{vel_2} \times \left( \frac{flow}{meanflow} \right)^{vel_3}
\]

\[
width = 10^{width_1} \times \left( \frac{flow}{velocity} \right)^{width_2 + width_3 \times \log_{10} \left( \frac{flow}{velocity} \right)}
\]
\[
\text{depth} = \left( \frac{\text{flow}}{\text{velocity}} \right) / \text{width}
\]

where vel1-vel3 and width1-width3 are lake region dependent parameters, flow is the flow of water in the watercourse \((m^3/s)\) and meanflow is a 365-day rolling average flow. The river’s width is limited by the dead volume width and a parameter maxwidth. The rivers length is estimated at square root of catchment area or is given as input. The watercourses bottom area is calculated as the length times the width, where the maximum of the above-calculated width and dead volume width is used. The bottom area is used for the nutrient processes. If the river is a class and thus has a defined class area, the class area is used to estimate the bottom area instead.

The water temperature \((T_w)\) is used in some of the process calculations. It is calculated through weighting the air temperature \((T)\) and yesterday’s water temperature. The weighting constant (similar to moving average period) \(T_{\text{wdays}}\) is set to 20 days for rivers and is by default 5 days for lakes. For lakes a depth dependent \(T_{\text{wdays}}\) can also be used. Then the weighting constant is set to the equivalent days of the lake's depths up to a maximum (maxpar, a parameter) plus 5 days. Note that the water temperature can fall below \(0^\circ C\).

\[
T_w = \left( 1 - \frac{1}{T_{\text{wdays}}} \right) \times T_w + \frac{1}{T_{\text{wdays}}} \times T
\]

\[
T_{\text{wdays}} = \begin{cases} 
\text{MIN} \left( \text{MAX} \left( \text{lake}_{\text{depth}} \times 5, 5 + \text{maxpar} \right) \right) & \text{lake} \\
20 & \text{river}
\end{cases}
\]

Alternatively the water temperature calculated as a tracer (called T2) may be used. This model option is set in info.txt.

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Denitrification

Denitrification, a sink for inorganic nitrogen in lakes and rivers, is a function of the bottom area, the IN concentration \( (conc) \) in water volume, water temperature \( (T_w) \) and a rate parameter. The dependence on concentration is formulated as a half saturation equation. In the concentration function, the half saturation parameter \( (par_{half}) \) is a general parameter, but it was in earlier HYPE versions a constant equal to 1.5 mg/L. Denitrification \( (denitr, kg/day) \) is limited to a maximum of 50% of the available IN pool (i.e. in SLP).

\[
tmpfcn = \begin{cases} 
0 & T_w < 0 \\
\frac{T_w}{5} \times 2^{10} & 0 < T_w \leq 5 \\
\frac{T_w-20}{2} \times 10 & else 
\end{cases}
\]

\[
concfcn = \frac{conc}{conc + par_{half}}
\]

\[
denitr = MIN \left( 0.5 \times INpool \times par_{den} \times area \times concfcn \times tmpfcn \right)
\]

The bottom area \( (area) \) is equal to the lake area for lakes, and watercourse width multiplied by the length for rivers (see above). The rate parameter \( (par_{den}) \) depend on the water body. Local and main rivers have general parameters \( (denitrwl, denitwrm) \), while lakes in addition to use a general parameter \( (denitwl) \) can have lake specific values.

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Primary production and mineralization

Primary production in lakes and rivers is a source of organic nitrogen and particulate phosphorus and a sink for inorganic nitrogen and soluble reactive phosphorus in the model. The reverse is true for mineralization. The processes are modelled together and only one is active at the time. Primary production and mineralization are controlled by two temperature functions. The first \((\text{tmpfcn1})\) is solely dependent on the water temperature \(T_w\). It simulates the increased activity at warmer temperatures. The second \((\text{tmpfcn2})\) governs the relationship between primary production and mineralization and determines which one dominates. Net primary production is highest in spring (northern hemisphere) and changes into net mineralization when the temperature \(T_{10}\) is less than the temperature \(T_{20}\) in autumn. These two temperatures are calculated as the average water temperature of 10 and 20 days.

\[
\text{tmpfcn1} = \frac{T_w}{20}.
\]

\[
\text{tmpfcn2} = \frac{(T_{10} - T_{20})}{5}.
\]

\[
\text{tmpfcn} = \text{tmpfcn1} \times \text{tmpfcn2}
\]

The primary production and mineralization processes are also governed by long-term average concentrations of modelled total phosphorus \((TP)\) in the lake or watercourse through a concentration function \((\text{TPfcn})\) with the general half saturation parameter \(h_{satTP}\). The half saturation parameter was in earlier HYPE versions a constant equal to 0.05 mg/L. If phosphorus is not modelled a long-term average total phosphorus concentration as a lake region dependent parameter \((tpmean)\) is used. A limiting concentration \((\text{lim})\) can be set, and then the long-term average concentration is reduced before using it in the concentration function.

\[
\text{TPfcn} = \frac{TP \cdot \text{lim}}{(TP \cdot \text{lim}) + h_{satTP}}
\]

For lakes, the process is acting only in lake part SLP, while the processes are active throughout the watercourse volume. The estimated production/mineralization \((\text{minprodNpot}, \text{kg / day})\) is the potential transformation, and may be limited by the availability of nutrients. Only 50% of the available IN the pool (at the primary production) or 50% of the ON-pool (for mineralization) can be transformed. The potential phosphorus conversion \((\text{minprodPpot})\) is calculated in the same way, but with its own parameter \((wprodp)\) and a factor for phosphorus/nitrogen ratio \((\text{NPratio} = 1/7.2)\). Similarly, there is a restriction against transforming maximum 50% of the SP and PP pools. The parameters \(wprodn\) and \(wprodp\) is generic or can be specified for each lake. The area is equal to lake area for lakes and bottom area for rivers (width multiplied by the length of the watercourse, see Basic assumptions, or if the river is a class, the class' area). The water depth \((\text{depth})\) is the SLP lake part, and for the river the depth calculated above.
*minprodNpot* = \( wprodn \times TP_{fen} \times tmppf_{en} \times area \times depth \)

*minprodPpot* = \( wprod\times TP_{fen} \times tmppf_{en} \times area \times depth \times NPratio \)

For simulations of total suspended sediments, algae is simulated as a component of the suspendend material. Nitrogen in algae is assumed to grow and decline with the same function as production and mineralisation of organic nitrogen. If nitrogen is simulated it uses the actual estimated production/mineralisation, but otherwise the potential production/mineralisation (\(minprodNpot\)) is used. The mineralisation of algae is limited to available amount, but the production is then unlimited.

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**Sedimentation/Resuspension**

Sedimentation in lakes is a sink for particulate phosphorus (PP) and organic nitrogen (ON), as well as for suspended sediments (SS) and algae (AE). Sedimentation (\(sed, \text{kg/day}\)) is calculated as a function of concentration (\(conc\)) in the lake and lake area (\(area\)). The sedimentation rate (\(par_{sed}\)) is given by parameters (\(sedon, sedpp, sedss, sedae\)) which are generic, but ON and PP sedimentation can be specified for each lake. The concentration used in the equation may be limited (\(lim\)) by general parameters (\(\text{limsedON, limsedPP, limsedSS}\)), but not for AE (\(lim=0\)).

\[
\text{sed} = par_{sed} \times \left(\frac{\text{conc}-\text{lim}}{\text{lim}}\right) \times \text{area}
\]

In the river no particles are removed, but they are redistributed over time. This process is used for particulate phosphorus (PP) and suspended sediments (SS). The process is a combination of sedimentation (\(sed, \text{m/day}\)) and resuspension (\(resusp, \text{m/day}\)). Particles in the sediments is collected in a pool (\(\text{sedimentpool}\)) which will increase as particles from the water volume (\(\text{waterpool}\)) settle at low water flows. The higher the water flow the lower the sedimentation and more particles returns to the water. This combined process is governed by the general parameter \(sedexp\). The net effect is determined by the sign of the variable (-1 <\(\text{sedresp}\)<1).
\[
\text{sedresp} = \max \left( -1, \min \left( 1, \frac{q_{\text{bank}} \cdot \text{flow}}{q_{\text{bank}}} \cdot \frac{\text{sedexp}}{\text{flow}} \right) \right)
\]

\[
\text{sed} = \begin{cases} 
\text{sedresp} \times \text{waterpool} & \text{sedresp} > 0 \\
0 & \text{sedresp} < 0 
\end{cases}
\]

\[
\text{resusp} = \begin{cases} 
-\text{sedresp} \times \text{sedimentpool} & \text{sedresp} < 0 \\
0 & \text{sedresp} > 0 
\end{cases}
\]

where flow is the current river flow (m³/s) and q_{\text{bank}} is the flow when river is filled to the brim. This flow is calculated as the second largest simulated flow in the last year. It is adjusted with a correction factor of 0.7 before use in the sedresp equation.

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**Internal load**

Lakes can leak phosphorus from the bottom e.g. release of phosphorus during oxygen deficiency or the mixing of the previously sedimented emissions. Internal load can be simulated for SP or PP or both fractions of phosphorus. With two parameters the release of PP or SP is obtained for a specific lake (par_{prod} m/d). The load to the lake (load, kg/day) is calculated as a function of lake water temperature \( T_w \), average concentration of phosphorus in the lake (TPfcn) and lake area (area). TPfcn is currently not depending on simulated concentration as it is set to constant 0.1 mg/L.

\[
\text{load} = \frac{\text{par}_{\text{prod}} \times \text{TPfcn} \times \text{tmpfcn} \times \text{area}}{1000}
\]

\[
\text{tmpfcn} = 0.86 \left[ T_w - 15 \right]
\]
Macrophyte uptake

For shallow waters in lakes macrophytes can grow and take up inorganic nutrients (IN and SP). The nutrients are lost to the model. Macrophyte uptake are controlled by a temperature function ($tmpfcn$) and a concentration function ($TPfcn$) for total phosphorus. The temperature function is composed of two parts; one exponential and one dependent on the water temperature ($Tw$) above the average temperature of the last twenty days ($T20$). The concentration function is the same half-saturation function as for production and mineralisation above.

$$tmpfcn = \frac{\left(\frac{Tw}{20}\right)^{0.3} \times (Tw - T20)}{5}, Tw > 0 \text{ and } tmpfcn > 0$$

The lake area ($fracarea$) that is shallower than a production depth ($proddep$, general parameter $muptdep$) is assumed to be active with macrophyte uptake. The lake is for this purpose assumed to be decreasing linearly with depth until twice the average depth of the lake.

$$fracarea = \frac{proddep}{\left(\frac{2 \times vol}{area}\right)} \times area$$

The uptake ($upt$) is limited to maximum 50% of the available nutrients and the whole equation becomes:

$$upt = uptpar \times tmpfcn \times TPfcn \times fracarea$$

The uptake rate parameters ($uptpar$), different for IN and SP, are general ($muptn, muptp$). The process is similar to the modelled macrophyte uptake in wetlands.
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Water management

Constructed wetlands for nutrient retention

For an overview of basic assumptions and explanation of variables see the Basic assumptions section in the Rivers and lakes chapter.

The wetlands that are simulated are small artificial ponds. They have an area and depth (dep), but their area is not taken into account in terms of precipitation and evaporation. The water flow passes through the wetlands without being affected, so it's just as nutrient traps that the wetland model is significant. There are two types of wetlands, just as for the rivers and lakes. They are situated before the river in the calculation scheme. The local wetland (lrwet) receives a share of the local runoff (part) the rest passes by unaffected. Wetlands in main rivers (mrwet) receive a portion of the flow in the main river and the rest passes unaffected.

Wetland nutrient processes

In wetlands, retention of inorganic nitrogen is modelled (denitrification). For total phosphorus retention (sedimentation) and production (or release from sediments) of TP are modelled. The rates of these processes are constant coefficients (teta=1.2, tkoeff=20, inpar=2.3, sedpar= 0.09, and uptpar=0.1). The retention is limited to 99.9% of the substance in the wetland. The retention (retIN, retTP, g/d) depends on the rate parameter, the concentration in the wetland, wetland area, and for inorganic nitrogen also on 5-day-mean air temperature (T5). The production (prodTP, g/d) depends on a rate parameter, the concentration of the inflow to the wetland, wetland area, and a temperature function (30-day-mean air temperature, T30). The change in TP are divided equally between SP and PP.

\[
retIN = \frac{\text{inpar} \times \text{INconc} \times T_5 \times \text{area}}{1000} \\
retTP = \text{sedpar} \times \text{TPconc} \times \text{area} \\
prodTP = \text{uptpar} \times \text{TPin} \times \text{area} \times teta \left( T_{30} - \text{tkoeff} \right)
\]

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References


**Constructed wetlands with water regulation capability**

The wetlands are water classes, but simulated as a land class with special functions. If the soil is over saturated the standing water is the water volume of the wetland ($vol$, m$^3$). If the soil is not over saturated, the wetland is dried out. The wetlands area ($area$, m$^2$) is defined by the class area, and the depth ($w$) varies with flow. A threshold ($w_0$) for the wetland outflow above the soil surface keep water in the wetland. The wetland outflow is determined by a rating curve above this threshold (see also lake outflow). The thresholds can be set by parameters or if parameters are not set it is equal to minus the streamdepth (from GeoClass).

$$out\ flow = k \times \left( w - w_0 \right)^p$$

There are two types of wetlands; internal wetlands ($iwet$) and wetlands at the outlet of the subbasin ($owet$). Internal wetlands receive a fraction ($ifraction$) of the runoff from other land classes. Outlet wetlands cannot be present in the same subbasin as an outlet lake. They receive the flow from the main river of the subbasin.

**Wetland nutrient processes**

The concentration of the wetland ($conc$) is the concentration of soil water in soil layer 1. While calculating wetland nutrient processes only the nutrients in the water volume of the wetland is considered though. After that the nutrient concentration of the upper soil layer is updated.

Denitrification of inorganic nitrogen in the wetland is modelled as denitrification in the soil water. Sedimentation of organic nitrogen, particulate phosphorus and suspended sediments are simulated ($sed$, g/d). Uptake of inorganic nutrients (IN and SP) are modelled as macrophyte uptake. The macrophytes are assumed to cover a part of the wetland area ($fracarea$). The covered fraction is calculated as the part that is shallower than a production depth ($proddep$) assuming the wetland area is decreasing linear with depth until twice the average depth of the wetland. The macrophytes are assumed to give residuals of equal amount of nutrient back to the sediment (i.e. immobile organic nutrient pools of soil layer one). The macrophyte uptake process ($upt$, g/d) depends on a rate parameter ($uptpar$), macrophyte fraction of wetland area, temperature ($tmpfcn$) and total phosphorus concentration ($TPfunc$). The temperature and TP functions are similar to the ones used by primary production in lakes. The temperature function use 5- and 30-day mean air temperature ($T5$, $T30$). The half saturation concentration of TP is 0.05 mg/L ($hsatTP$). The sedimentation is limited to 99.9% of the substance in the wetland water, while macrophytes are limited to 50% of the dissolved inorganic nutrients.

$$sed = velpar \times area \times conc$$
\[ u_{pt} = u_{tp} \times \text{tmpfcn} \times \text{TPfcn} \times \text{fracarea} \]

\[ \text{tmpfcn} = \left( \frac{T_5}{20} \right)^{\text{tmppar}} \times \left( \frac{T_5 - T_30}{5} \right) \]

\[ \text{TPfcn} = \frac{\text{TPconc}}{\text{TPconc} + \text{hsatTP}} \]

\[ \text{fracarea} = \left( \frac{\text{proddep}}{\text{area}} \right) \times \text{area} \]

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**Irrigation**

**General principles**

Irrigation constitutes a key water management activity in many parts of the world. Therefore, the HYPE model has a routine to simulate irrigation. The representation of irrigation in the model is based on a set of principles. Firstly, the irrigation water demand is assessed. Subsequently, the demanded water is withdrawn from the defined irrigation water sources. HYPE can either withdraw water from defined sub-basins in the model domain (subject to availability), or from unlimited sources outside the domain. Finally, the withdrawn water is applied onto the classes from which the demand originated. In addition, water losses between demand, withdrawal, and application are taken into account (for withdrawals within the model domain).

A class is irrigated if the crop type associated with it is irrigated (defined in GeoClass.txt). A crop is
irrigated if the irrigation input variables in the CropData.txt file are defined and non-zero (plantday, lengthini, kcbini, lengthdev, lengthmid, kcbmid, lengthlate, kcbend, dlref). Irrigation also requires appropriate values in the MgmtData.txt file (gw_part, regsrcid, irrdam, region_eff, local_eff, demandtype) and the par.txt file (pirrs, pirrg, sswcorr etc.). See the File Reference for more details on each file and each parameter.

**Irrigation water demand**

The irrigation water demand \( W_{i,D,j} \) is calculated each day for each irrigated class \((j)\) at the end of the soil water balance calculations. Two approaches to calculate \( W_{i,D,j} \) are implemented in HYPE, one for submerged crops (e.g. paddy rice) and one for non-submerged crops. The input variables \textit{imm_start} and \textit{imm_end} in CropData.txt define (1) the beginning and end of the submerged season, and (2) if crops are submerged or not (zero is interpreted as a non-submerged crop).

**Non-submerged crops**

For non-submerged crops, the calculations are based on the FAO-56 crop coefficient methods (Allen et al., 1998). The dual crop coefficient method is used because it is more specific than the single crop coefficient method, and more suitable for daily water balance models. Since transpiration is of primary interest in estimating crop water demand, the irrigation routine focuses on estimating potential transpiration \( T_P \) with the basal crop coefficient \( K_{CB} \) and the reference potential crop evapotranspiration \( ET_0 \):

\[
T_P = K_{CB} \times ET_0
\]

\( ET_0 \) follows the dynamics described above (here following Wisser et al. (2008)). \( K_{CB} \) depends on crop type and phenological stage, which is defined in CropData.txt. \( K_{CB} \) is constant during the initial development stage, then increases linearly during the development stage until it reaches the mid-season stage during which it is again constant. Finally, \( K_{CB} \) decreases linearly from the end of the mid-season stage until the end of the season. The dynamics of \( ET_0 \) and \( K_{CB} \) produces a dynamic \( T_P \) profile (Figure 1). Allen et al. (1998) provide indicative values for \( K_{CB} \) (cf. their Table 17).
On any given day, the model first calculates whether irrigation is needed, and then the amount required. The irrigation need is assessed by comparing the current soil water content \((H)\) with a dynamic irrigation threshold \((S_{SW})\), the soil water stress threshold):

\[
i \text{If } H < \left( S_{SW} \times P_{SSWCor} \times AWC \right) \rightarrow \text{irrigate}
\]

\(H\) is the plant-available soil moisture (i.e. soil water above \(wcwp1\) and \(wcwp2\) in soil layers 1 and 2 respectively). \(AWC\) is the maximum plant-available water content in soil layers 1 and 2 (i.e. the sum of \(fc1\) and \(fc2\)). \(S_{SW}\) is a fraction of \(AWC\) (defined upwards from \(wcwp\)). Below \(S_{SW}\) the crop experiences water stress, creating a need for irrigation. \(S_{SW}\) varies from day to day and depends on the crop type and \(T_P\):

\[
S_{SW} = 1 - \left( DL_{ref} + 0.04 \times \left( 5 \times \frac{T_P}{0.95} \right) \right)
\]

\(DL_{ref}\) is a crop-type specific reference depletion level (essentially the fraction of \(AWC\) that can be depleted before stress occurs, defined downwards from \(wcfc\)). Allen et al. (1998) provide indicative...
values for $DL_{\text{ref}}$ (cf. their Table 22). The $S_{SW}$ equation is a slightly modified form of the original FAO-56 equation to account for the fact that only $TP$ is used here. A typical $S_{SW}$ profile is shown in Figure 3.1. By default, $S_{SW}$ is limited to the range 0.2 - 0.9, but it can be further refined with the parameter $P_{SSWCORR}$ ($sswcorr$ in par.txt) to maximum 1.

If irrigation is needed, the required irrigation amount ($W_{I,D,j}$) can be calculated with three alternative methods in HYPE (chosen by the demandtype variable in MgmtData.txt):

1. A constant $W_{I,D,j}$ (defined by the $irrdemand$ parameter in par.txt)

2. Up to the field capacity: $W_{I,D,j} = AWC - H$

3. Up to a defined fraction of $S_{SW}$ ($P_{iwdfrac}$, $iwdfrac$ in par.txt):

$$W_{I,D,j} = \min \left( S_{SW} \times P_{SSWCORR} \times AWC - H \right) \times P_{iwdfrac} \times \left( AWC - H \right)$$

The fraction can be larger than 1. For example, to irrigate to a level 10% above $S_{SW}$, $P_{iwdfrac} = 1.1$. $W_{I,D,j}$ is, however, limited to $AWC$.

**Submerged crops**

The irrigation of submerged crops aims to satisfy a target flooding level above the soil surface (Wisser et al., 2008). The target flooding level is a constant input parameter ($P_{immdepth}$, $immdepth$ in par.txt). Irrigation is required if the water level of the top soil layer (H1) falls below the target flooding level:

$$i_f H_1 < WP_1 + FC_1 + EP_1 + P_{immdepth} \rightarrow \text{irrigate}$$

The irrigation water demand is equal to the amount needed to reach the target flooding level:

$$W_{I,D,j} = WP_1 + FC_1 + EP_1 + P_{immdepth} - H_1$$

If the submerged season is shorter than the crop season, $W_{I,D}$ during the non-submerged period is calculated in the same way as for non-submerged crops. To maintain a desired flooding level and simulate terracing, for example, it may be necessary to adjust the soil and runoff parameters of the class with the submerged crops. When $W_{I,D,j}$ has been calculated for each irrigated class in the sub-basin, the total field-scale irrigation water demand for the sub-basin ($W_{I,D}$) is calculated:

$$W_{I,D} = \sum_{j=1}^{N} W_{I,D,j}$$
**Irrigation water withdrawal**

Irrigation water can be abstracted from a set of water sources (Figure 2). Within a given sub-basin, water can be abstracted from the olake, the ilake, the main river, and from groundwater in a deep aquifer. In addition, water can be withdrawn from the olake and the main river of another sub-basin. These sources can be used on their own or in combination. Alternatively, HYPE can withdraw water from an unlimited source outside the model domain. This is specified with the `irrunlimited` code word in info.txt, and applies to all irrigated sub-basins.

Withdrawals are calculated directly after the local discharge and the upstream discharge has been combined to flow into the main river of a given sub-basin.

![Figure 2: Schematic illustration of the available irrigation sources in HYPE. Irrigation water can be withdrawn from: (i) dams in the sub-basin (olake and ilake), (ii) the main river in the sub-basin, (iii) groundwater in a deep aquifer, and (iv) dams (olake) and main rivers in other sub-basins.](image)

**Irrigation inefficiencies within the sub-basin**

Before any withdrawal occurs, the field-scale \( W_{I,D} \) is scaled to sub-basin scale \( W_{L,D} \), the local sub-basin irrigation water demand. This is done in order to account for the often significant water losses between withdrawals and soil moisture replenishment. A simple user-provided scaling factor is...
The local efficiency \( E^L \), \( local\_eff \) in MgmtData.txt represents the fraction of the withdrawn water within the sub-basin that infiltrates the irrigated soil. \( E^L \) accounts for losses in irrigation canals, ponds etc. within the sub-basin, and for on-field losses from irrigation equipment (e.g. sprinkler systems). \( W_{L,D} \) is not scaled if all withdrawals are from an unlimited source outside the model domain (i.e. \( E^L = 1 \) if \( irr\_unlimited \) is \( y \)).

**Withdrawal from sources within the sub-basin**

The model first attempts to withdraw water from sources within the sub-basin where the demand originated. The user specifies the proportion of water to be withdrawn from surface water and deep aquifer groundwater sources, respectively, in the MgmtData.txt file (\( gw\_part \)). A \( gw\_part \) value between 0 and 1 represents the long-term average proportion of surface and deep aquifer groundwater withdrawals for irrigation within the sub-basin. A \( gw\_part \) value of 0 indicates that all irrigation water exclusively comes from surface water sources, while a \( gw\_part \) value of 1 indicates that all irrigation water exclusively comes from deep aquifers. Based on \( gw\_part \), \( W_{L,D} \) is split into the groundwater demand \( W_{L,D,g} \), and the surface water demand \( W_{L,D,s} \).

If any surface water demand exists, the model sequentially attempts to withdraw water from the olake, the ilake, and the main river. However, water withdrawal from olakes and ilakes is only calculated if the variable \( irr\_dam \) in MgmtData.txt is set to 1 for the sub-basin. Water withdrawals from the main river occur both from the inflow to the river reach and from the volume stored in the reach.

If the \( W_{L,D,s} \) volume is available at the source, the demanded water is withdrawn. Otherwise, only the available volume \( V^s \) is withdrawn. The withdrawal can also be scaled with the user-defined parameter \( P_{I,S} \) (\( pirrs \) in par.txt):

\[
W_{L,A,s} = \min\left(W_{L,D,s}, V^s\right) \times P_{I,S}
\]

\[
W_{L,D,s2} = W_{L,D,s} - W_{L,A,s}
\]

where \( W_{L,A,s} \) is the abstracted water from the first surface water source in the sub-basin, and \( W_{L,D,s2} \) is the residual surface water demand. The residual demands \( W_{L,D,s1}, W_{L,D,1} \) and \( W_{R,D2} \) are calculated without the \( P_{I,S} \) scaling in order to prevent erroneous source compensation due to scaling. If any demand remains, the next surface water source
is probed in the same manner:

$$W_{L,A,s} = \min \left( W_{L,D,s_2}, V_{s_2} \right) \Box P_{I,S}$$

The total surface water withdrawal within the sub-basin ($W_{L,A,s}$), and the remaining surface water demand ($W_{L,D,s,1}$) is calculated accordingly, based on the abstractions from each source ($k$):

$$W_{L,A,s} = \sum_{k=1}^{N} W_{L,A,s}^{(k)}$$

If any groundwater demand exists, the model withdraws a user-specified fraction of $W_{L,D,g}$ from an unlimited source outside the model domain or if an aquifer is simulated connected to the subbasin from this aquifer. The first case conceptually represents a large deep aquifer source, which is currently outside of the model domain. $W_{L,A,g}$ is the abstracted groundwater and $P_{I,G}$ (pirrg in par.txt) is the groundwater withdrawal fraction:

$$W_{L,A,g} = W_{L,D,g} \times P_{I,G}$$

To simulate a more dynamic conjunctive use of groundwater and surface water sources, the model allows for compensation of remaining surface water demands from the groundwater source. This compensation is only allowed if both groundwater and surface water sources are used ($0 < gw_part < 1$), and if the $irrcomp$ parameter is >0. The $irrcomp$ parameter defines the degree of compensation allowed, i.e. the fraction of the residual surface water demand which can be met through source compensation. The compensation algorithm is as follows: if any surface water demand remains ($W_{L,D,s,1} > 0$) and the groundwater is not depleted, the groundwater withdrawal cycle is calculated once more using the scaled residual surface water demand. Finally, after possible source compensation, the remaining (surface) water demand at the sub-basin scale ($W_{L,D,1}$) is calculated.

**Withdrawal from another sub-basin**

The model can also simulate withdrawal from another sub-basin in the model domain ($D_R$, defined with the $regrsid$ input variable in MgmtData.txt). This withdrawal is calculated when the model reaches $D_R$ in the calculation order, after possible local irrigation water abstractions in $D_R$ and only if any irrigation demand remains in the sub-basin(s) connected to the regional source ($W_{L,D,1} > 0$). For each connected sub-basin ($i$), $W_{L,D,i}$ is scaled to represent the regional-scale demand from that sub-basin ($W_{R,D,i}$). Again, a simple scaling factor is applied:
The regional efficiency ($E_{R,i}$, `region_eff` in MgmtData.txt) represents the fraction of the withdrawn water at the regional source that reaches the connected sub-basin. $E_{R,i}$ refers to the connected sub-basin. The regional scaling accounts for often significant water conveyance losses in large irrigation networks (in canals and dams etc.).

The total water demand from the regional source ($W_{R,D}$) is then calculated as the sum of the demand from each connected sub-basin, scaled by a parameter controlling the strength of the regional connection ($P_{regirr}$, `regirr` in par.txt):

$$W_{R,D} = \left( \sum_{i=1}^{N} W_{R,D,i} \right) \times P_{regirr}$$

The regional demand can be met from two sources in sub-basin $D_{R}$: the lake and the main river. If the regional source sub-basin has an lake, and if the `irrdam` input variable is set to 1 for that sub-basin, the model attempts to withdraw $W_{R,D}$ first from the lake and then the residual from the main river. If not, the model only attempts to withdraw $W_{R,D}$ from the main river. The regional abstraction ($W_{R,A}$) is limited by the volume available at the source ($V_{r}$) and the scaling parameter $P_{I,S}$:

$$W_{R,A1} = \min\left( W_{R,D}, V_{r1} \right) \times P_{I,S}$$

$$W_{R,D2} = W_{R,D} - \frac{W_{R,A1}}{P_{I,S}}$$

$$W_{R,A2} = \min\left( W_{R,D2}, V_{r2} \right) \times P_{I,S}$$

$$W_{R,A} = W_{R,A1} + W_{R,A2}$$

where $W_{R,A1}$ is the abstracted water from the first water source in $D_{R}$, $V_{r1}$ the volume available at the first source, $W_{R,D2}$ the residual regional water demand after withdrawal from the first source (but prior to the $P_{I,S}$ scaling), $W_{R,A2}$ the abstracted water from the second water source in $D_{R}$, and $V_{r2}$ the volume available at the second source.

**Substance concentrations of irrigation water withdrawals**

The concentrations of the withdrawn water are the same as that of the irrigation water source. If the water originates from several sources, the volume-weighted concentration is calculated. If desired,
the model can simulate sedimentation tanks, in which a defined fraction of the particulate phosphorous ($pp$) and organic nitrogen ($on$) settles:

\[ C_{a,pp} = C_{src,pp} \times \left(1 - P_{cirrsink}\right) \]

\[ C_{a,on} = C_{src,on} \times \left(1 - P_{cirrsink}\right) \]

where $C_{a}$ is the concentration of the abstracted water after settling, $C_{src}$ is the concentration of the source, and $P_{cirrsink}$ is the concentration reduction fraction (cirrsink parameter in par.txt). To use sedimentation tanks in a region, the concentration reduction fraction needs to be set in par.txt ($0 < cirrsink \leq 1$).

**Irrigation water application**

In the calculation order, the irrigation water application occurs the next time step the model reaches the sub-basin from which the demand originated (typically the following day). The water is applied to the soils at the beginning of the soil balance calculations, before the calculation of the natural processes.

The regionally abstracted water ($\mathcal{W}_{R,A}$) is first distributed to each connected sub-basin ($i$) according to their proportional demand, and then scaled to the local scale using the respective regional efficiency:

\[ \mathcal{W}_{R,A,i} = \frac{\mathcal{W}_{R,A} \times \mathcal{W}_{R,D,i} \times P_{regiwe}}{\mathcal{W}_{R,D} \times E_{R,i}} \]

For a given sub-basin, the total amount of abstracted water available at the local scale ($\mathcal{W}_{L,A,i,\text{tot}}$) is calculated and then scaled, using the local efficiency, to represent the water applied to the soil ($\mathcal{W}_{I,A,i}$):

\[ \mathcal{W}_{L,A,i,\text{tot}} = \mathcal{W}_{L,A,i} + \mathcal{W}_{R,A,i} \]

\[ \mathcal{W}_{I,A,i} = \mathcal{W}_{L,A,i,\text{tot}} \times E_{L,i} \]

$\mathcal{W}_{I,A,i}$ is then distributed onto each irrigated class in proportion to its water demand:

\[ \mathcal{W}_{I,A,j} = \frac{\mathcal{W}_{I,A,i} \times \mathcal{W}_{I,D,i}}{\mathcal{W}_{I,D}} \]

$\mathcal{W}_{I,A,j}$ is added to the soil water of class $j$ as additional infiltration. $\mathcal{W}_{I,A,j}$ is divided between the top two soil layers according to the epotdist function, beginning with the second layer. For unlimited irrigation: $\mathcal{W}_{I,A,j} = \mathcal{W}_{I,D,j}$, $P_{I,S} = 1$ and $P_{I,G} = 1$.

The water withdrawn from a regional source that does not reach connected sub-basins ($\mathcal{W}_{R,L}$)
Evaporates at the regional source \( D_R \):

\[
W_{R,L} = W_{R,A} \sum_{i=1}^{N} W_{R,A,i}
\]

Similarly, water losses due to local inefficiencies \( W_{L,L} \) evaporate within the sub-basin itself. This applies to local losses from abstractions both within the sub-basin and from the regional source (where applicable):

\[
W_{L,L} = W_{L,A,i,tot} - W_{I,A,i}
\]

Evaporation due to regional and local inefficiencies proportionally concentrates substances in the withdrawn water. The substance concentrations in the irrigation water applications are hence higher than at the points of withdrawal (the mass remains the same while the volumes are reduced). However, if unlimited irrigation is simulated, the concentrations of the applied water are the same as in the layers to which water is added (i.e., causing no change in concentration).

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References


Point sources

Information on point sources is located in the file PointSourceData.txt.

Nutrient point sources

The model can handle up to three different types of point sources. They can used to simulate e.g. treatment plants, stormwater outlets, and industrial sources as separate types. All point sources have a constant flow, concentrations of total nitrogen and phosphorus, and fractions of IN and SP for a period of time. The time may be the whole simulation period, or different sources may be active during different parts of the simulation period. Point sources are added to the water in the main river.

Sediment point source

Sediment point sources are handled similar to nutrient point sources. A point source has a constant flow, concentration of total suspended sediment (TS), and fraction of suspended sediments (SS). The point source may be active the whole simulation period, or different sources may be active during different parts of the simulation period. Point sources are added to the water in the main river.
Tracer T2 (water temperature) point sources

Water temperature may be added to the flow of nutrient point sources if T2 is simulated together with N and P. Water temperature point source may also be added on its own in the same way as nutrient point sources.

Tracer T1 point sources

Tracer T1 point source may be added in the same way as nutrient point sources. In addition, point sources of tracer T1 can be added to the local stream, the local lake, the main river or the outlet lake.

Negative point source

A point source with negative flow denotes an abstraction of water. The abstraction of water can be made from three different points in the river network. The default is to remove it from the main river volume (including the queue) after all river processes have been calculated except outflow from the river. Alternatives are to abstract the water from the outlet lake volume or from the main river inflow from upstream and from the river volume (and queue) proportionally. In the latter case the removal is done before any inflow to the main river is added (e.g. from upstream, point sources, or precipitation). The water is removed from the source, while the concentration is kept.

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Water abstraction and transfer

Water transfer can be simulated by HYPE in different ways. One way is to represent abstraction by defining negative point source discharge. Another method is to use the bifurcation functionality that defines a branch through which the water is transferred to a downstream receiving subbasin. A third way is to define water transfer through water management (abstracting from subbasins with outlet lakes and transferring to any other subbasin). The negative point source method has the disadvantage that the water transfer is constant and with fixed concentrations specified by the user (although you can let it be constant at different values for different periods), but the abstraction can be located at different points throughout the catchment. The bifurcation method has the limitation that the receiving subbasin must be downstream. The management method can only take water from an outlet lake, and the water transfer will be delayed one time step. In all methods, the amount of water available on any given day will limit the amount that can be withdrawn or transferred.

Water transfer through bifurcation

Water transfer through bifurcation lets you divert a part of the outflow from a subbasin through a branch to another downstream subbasin specified in BranchData. The flow diverted may be defined by BranchData parameters (fraction, max-, or minflow). The water transfer can alternatively be given as a recorded time series of demanded water transfer (given as dwtr in Xobs). This water will then be taken from the ordinary simulated outflow of the subbasin if there is enough flow, the remaining flow will go through the main channel. A third method, if you have an outlet lake, is to use two outlets. The flow of outlet 2 can be defined as the flow given by dwtr.

See also the general description of bifurcations and Outlet lake with two outlets.

Water transfer through negative point source

Abstraction of water can be defined as a negative point source. The abstracted water is constant (although it can be constant at different values for different periods), and thus the same amount of water is removed from the model every time step. The abstraction can be made from different points in the river network within the subbasin. The water is removed while the concentration is kept in the water remaining in the river. To simulate water transfer within the model domain, the same amount of water need to be added as a (positive) point source in another subbasin. This subbasin do not have to be downstream, since its (constant) flow and concentrations is defined in the file beforehand. However, there is no direct connection between the subbasins in which water is abstracted and released, respectively. In order to preserve the water balance, the user must define equally large flows abstracted and added. Since the concentration of the negative point source is defined in the file beforehand, the connection to the quality of the water abstracted water is not automatically preserved.

Water transfer through water management

Water transfer between two subbasins can be defined through water management (MgmtData.txt) if the source is an outlet lake. The source subid and subid of the receiving subbasin is defined together with a constant flow. The water is taken from the outlet lake of the source subbasin and is given the
concentration of the lake. The water is then transported to the other subbasin's main river and arrive there the next time step. The delay in flow is necessary to handle transfer upstream within a catchment. Several water transfer can be specified for the same source or receiving subbasin.

Alternatively a demanded flow time series from Xobs can be used instead of a constant flow. In this case only one water transfer can be specified for each source lake.

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Dams

Regulation of flow through dams is described in the Chapter about Rivers and lakes. Dams of different purposes and regulation management can be simulated. See details in the sections on Simple outlet lake or dam and the special case of an Outlet lake with two outlets.
Deep processes

There are two model options for processes deep below ground; regional groundwater flow and aquifers. The regional groundwater flow model simulates groundwater flow between subbasins, but no explicit storage of deep groundwater. The aquifer model simulates aquifer storage and delay before regional groundwater reaches the destination.

Regional groundwater flow

Soil regional groundwater flow creation

Regional groundwater flow from a subbasin is calculated before the subbasin and class loops. This water is added to the downstream subbasin (in the class-loop part of the code) and/or to the outlet lake of the present subbasin (in the routing calculation part of the code). A subbasin dependent variable (grwtolake) governs the allocation between them. The outflow from a subbasin is calculated as a percentage of the water-retention capacity with a recession coefficient (rcgrw) e.g. for the lowest soil layer (k) the groundwater outflow for a class becomes:

\[ q_{grw}(t) = rcgrw \times \left( \text{soil}(k) - \text{wp}(k) + fc(k) \right) \]

Groundwater flows are summarized for all classes in proportion to their areas and converted to volume flow out of the subbasin.

Soil addition of regional groundwater flow

The groundwater flow to be added to the bottom layer (and mixed) is added in the class loop portion of the code. If the layer then becomes full of water, it will be pressed into the layer above and mixed, etc. Please note that if downstream subbasin is much smaller than the upstream subbasin, groundwater inflows can become large. If part of the groundwater flow is to go to the outlet lake in the subbasin this is added to the volume of the lake in the routing part of the code.

Regional groundwater to outlet lake

With an input variable in GeoData.txt, an outlet lake may be flagged to receive inflow from the regional groundwater. The coefficient (grwtolake) specifies the portion of the produced regional groundwater flow which is led to the subbasins outlet lake.

Links to file reference
Aquifers

Deep percolation

Deep percolation is determined by a recession coefficient \( r_{c_{\text{grw}}} \). The recession coefficient is set based on the recession parameter \( r_{c_{\text{grw}st}} \) which is soil type dependent, but if it is not set the general parameter \( r_{c_{\text{grw}}} \) is used instead. The recession coefficient can also be regionally adjusted by the correction parameter \( a_{\text{qperc}} \).

\[
r_{c_{\text{grw}}} = r_{c_{\text{grw}}} \left( 1 + a_{\text{qperc}} \right)
\]

Deep percolation from lowest soil layer in model is determined by the recession coefficient and the available water for runoff in the soil layer \( a_{\text{qperc} \text{ (mm/timestep)}} \).

\[
q_{\text{perc}}(t) = r_{c_{\text{grw}}} \times \left( sm - fc - wp \right) \quad sm > fc + wp
\]

Deep percolation from main river in model is determined by the recession coefficient and the water in the river \( a_{\text{qperc} \text{ (m3/timestep)}} \).

\[
q_{\text{perc}}(t) = r_{c_{\text{grw}}} \times \left( water \cdot depth \right)
\]

The water is removed from river water and the queue in proportion to their relative volumes.
Deep percolation delay and aquifer recharge

Deep percolation from soil and rivers in all recharge subbasins of an aquifer is added together. This total deep percolation \( q_{\text{perc}} \) is delayed before it recharges the aquifer \( q_{\text{rech}} \).

The delay property \( \text{delay} \), in days is specific for each aquifer. It can be adjusted by the parameter region dependent model parameter \( \text{aqdelcor} \).

Return flow from aquifer

Outflow from the aquifer depends on the water in the aquifer \( \text{vol} \) and a rate. The rate is determined by a aquifer property \( \text{retrate} \) and a regional adjustment parameter \( \text{aqretcor} \), parameter region dependent.

\[
q_{\text{out}} = \text{retrate} \times \left( 1 + \text{aqretcor} \right) \times \text{vol}
\]

The outflow may be divided between several receiving subbasins \( \text{retfrac} \). The water is added to the inflow of the main river of these subbasins.

The return flow is added to the main river of the receiving subbasin.

Nutrient processes related to aquifers

The organic nitrogen, particulate phosphorus, and organic carbon in water percolating from soil and river stay in the soil/river, while temperature, inorganic nitrogen and SRP follow the water to the aquifer. Inorganic nitrogen is subjected to denitrification in the aquifer (see Chapter Nitrogen and phosphorus in land routines - Denitrification).

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Organic carbon

Figure 1: Schematic figure of the OC-model.

Source of organic material

Litter fall in the form of plant residues add organic material to HYPE. It increases the levels of fastC in top two layers in soil. The organic carbon addition by litter fall is defined based on crop. Input, resc (kg/ha/yr), gives a daily supplement to the pool during the number of days determined by parameter litterdays.

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Soil processes

Soil pool initial values

The initial size of organic carbon pools in the soil is dependent on land use and determined by the user. The parameters (humusc1, humusc2, humusc3, fastc1, fastc2, fastc3) give OC content of the three soil layers. The unit for these parameter values is mg/m3. With this information, the pools the size in the different layers are calculated. The model transforms pools into the unit kg/km2 by taking into account the thickness in the layers.

Common functions

Many soil processes depend on temperature and soil moisture. They use the same common functions as nitrogen and phosphorus. Organic carbon soil transformations (production of humusC from fastC, turnover of fastC and turnover of humusC) use the soil moisture function with parameters given by the user instead of the coefficients described for nutrients. The coefficient $\theta_{low}$ is replaced by the
land-use dependent parameter `ocsoilslp`, and the coefficient `satact` is replaced by land-use dependent parameter `ocsoilsat`.

![Diagram of organic carbon processes in soil](image)

**Figure 2: Organic carbon processes in soil.**

### Production of humusC from fastC

Some of the litter fall is converted into humus. For HYPE this means that fastC (the pool where litter fall was added) is transformed to humusC in the uppermost soil layer. In Figure 2 the pool of humusC is denoted slowC because of its slower transformation rates.

The other soil layers \( (k) \) also have a transition from fastC to humusC. The loss of fastC does not all end up in the humusC pool, but a proportion (parameter `minc`) is mineralized in the process. The transformation \( (fasttohumus, \text{mg/m}^2/\text{d}) \) depends on soil moisture \( (smfcn) \) and temperature \( (tmpfcn) \), amount of Oc in the pool \( (fastC) \) and a vegetation dependent parameter \( kih \).

\[
fasttohumus(k) = kih\times tmpfcn(k)\times smfcn(k)\times fastC(k)
\]

### Turnover of fastC

Turnover of fastC is a sink for fastC and a source of dissolved OC in soil water in all soil layers \( (k = 1-3) \). The loss of fastC does not all go to the OC, but a proportion (parameter `minc`) is mineralized. Turnover \( (transFC, \text{mg/m}^2/\text{d}) \) depends on a general parameter \( (klo) \), the temperature function \( (tempfcn) \), humidity function \( (smfcn) \) and the pool of fastC \( (fastC) \).

\[
transFC(k) = klo\times tempfcn(k)\times smfcn(k)\times fastC(k)
\]

In dry conditions a transfer in the opposite direction can also occur. The transformation of OC to fastC is a decrease of OC and a source of fastC in all soil layers \( (k = 1-3) \). The loss of OC is not all to fastC but a proportion (parameter `minc`) is mineralized. Turnover \( (doctofast, \text{mg/m}^2/\text{d}) \) depends on a general parameter \( (kof) \) and the pool of OC \( (OCpool) \). The transfer is limited that the soil layer temperature must be less than 5 °C, the soil moisture \( (sm) \) must be less than field capacity and moisture function \( (smfcn) \) must be less than the parameter \( koflim \).
Turnover of humusC

Turnover of humusC is a sink for humusC and a source of OC in all soil layers (k = 1.3). The turnover rate of humusC is lower than that of fastC, why it is also called slowC (e.g. in Figure 2). The loss of humusC does not all go to the DOC, but a proportion (parameter minc) is mineralized. Turnover (transhC, mg/m2/d) depends on a general parameter (kho), temperature function (tempfcn), humidity function (smfcn) and the pool of humusC (humusC).

\[ \text{transhC}(k) = \text{kho} \times \text{tempfcn}(k) \times \text{smfcn}(k) \times \text{humusC}(k) \]

Percolation

Organic carbon is lost from the soil water as it flows down through the soil layers and where it is dissipated to become a regional groundwater flow. The decrease in concentration depends on soil moisture and temperature and a calibration parameter.

\[ \text{conc} = \text{conc} \times \left( 1 - \text{par} \times \text{tempfcn} \times \text{smfcn} \right) \]

The soil moisture function and temperature function are the general functions described for soil processes. Percolation uses the nutrient coefficients for the soil moisture function, not the parameters that the OC transformations uses. The parameter, par in the equation, is kcgwreg for regional groundwater flow formation and koc for percolation between soil layers. Both are general parameters.

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Riparian zone

Runoff from soil may flows through a riparian zone before it reaches the local river. Surface runoff and drainage water from drainage pipes reaches the local river without passing through the riparian zone. In the riparian zone the levels of OC are affected, while flows remain unchanged. The change depends on soil temperature, class altitude (elev (in masl)), the water table (gwat) and its recent change, season and soil moisture (sm). The runoff concentration (conc(i)) of each soil layer (k) increases with the factor:

\[ f(k) = 1 + \text{r iPz} \times \text{tempfcn}(k) \times \left( \frac{\text{elev}}{100} \right) \times \text{f grw} \times \text{f season} \times \text{f sm} \]
The temperature function \((\text{tmpfcn})\) is the usual of soil processes (see above). The following equations describe the other process functions:

\[
f_{\text{grw}} = c^{\text{ripe} \times \text{gwnt}}
\]

\[
f_{\text{season}} = \begin{cases} 
\text{rips autumn} & \text{sm} \leq \text{wp} \\
1 & \text{otherwise}
\end{cases}
\]

\[
f_{\text{sm}} = \begin{cases} 
0 & \text{sm} \leq \text{wp} \\
f_2(\text{sm}) & \text{wp} < \text{sm} < \text{pw} \\
\text{satact} & \text{sm} \geq \text{pw}
\end{cases}
\]

\[
f_2(sm) = \begin{cases} 
\min \left( \frac{1}{d \times \Theta_{\text{up}}}, \frac{1}{d \times \Theta_{\text{low}}}, \frac{\text{satact} \times (\text{satact} - \text{sm} - \text{wp})}{\text{satact} \times \text{wp} - \text{sm}} \right) & \text{rising grw} \\
\min \left( \frac{1}{d \times \Theta_{\text{up}}}, \frac{1}{d \times \Theta_{\text{low}}}, \frac{\text{satact} \times (\text{satact} - \text{sm} - \text{wp})}{\text{satact} \times \text{sm} - \text{wp}} \right) & \text{sinking grw}
\end{cases}
\]

The activation of riparian zone processes is based on land use. It is primarily though to act on forest runoff. The land use dependent parameter \(\text{ripz}\) determines the overall level of increase in concentration in the riparian zone, and if set to zero no riparian zone processes are used. In addition two general parameters can influence the effect of the riparian processes; \(\text{ripe}\) which determines the groundwater level dependence, and \(\text{rips}\) which determines the seasonal influence. Season division is determined by ten-day and twenty-day averages of air temperature (T10, T20). Autumn is defined as the period when T10 is less than T20. The soil moisture function is different for an increasing (rising) and sinking ground water table (figure 2). It contains coefficients \(\Theta_{\text{up}} = 0.12\), \(\Theta_{\text{low}} = 0.08\) and saturation \(\text{satact} = 0.6\). It depends on the soil moisture of all layers together \((\text{sm})\) and the water-holding capacity parameters; \(\text{wp}\) - wilting border and \(\text{pw}\) - total pore volume, in fractions of total soil layer thickness \((d)\).
Rivers and lakes

The initial organic carbon concentration in rivers are assumed to be zero, while the lakes' concentration are set by the user. The parameter, $tocmean$, is lakeregion dependent, but can also be set for each lake separately.

Primary production and mineralization

Primary production is a source of organic carbon in rivers and lakes, while mineralization is a sink. Primary production and mineralization is calculated the same way as for nitrogen, but with its own calibration parameter ($wprodc$). The equations are repeated below. The production/mineralization depend on temperature and total phosphorus and lake area ($area$). The potential carbon transformation ($minprodCpot$, kg / day) is proportional to the potential nitrogen transformation ($minprodNpot$, see also NP section) with a transformation rate that depends on the carbon-nitrogen ratio ($NCratio = 5.7$). The calculated mineralization of organic carbon is limited to a maximum of 50% of the available OC pool. If phosphorus is not modelled a long-term average total phosphorus concentration as a lake region dependent parameter ($tpmean$) is used. If set, the long-term average concentration is reduced by the general parameter $limsedPP$ before using it in the concentration function thus reducing the production/mineralisation of OC.

$$tmpfcn1 = \frac{water temp}{20}$$

$$tmpfcn2 = \left(\frac{T1o - T2o}{5}\right)$$

$$tmpfcn = tmpfcn1 \times tmpfcn2$$
$TPcn = \frac{\left( TPC\text{nc}\cdot \text{limsedPP} \right)}{\left( TPC\text{nc}\cdot \text{limsedPP} + \text{halfsatTPwater} \right)}$

$\text{minprodNpot} = \text{wprodC} \times TPcn \times TGcn \times \text{area}$

$\text{minprodCpot} = \text{minprodNpot} \times NCratio$

**Sedimentation**

Sedimentation in lakes is a sink for OC and works the same way as for organic nitrogen and particulate phosphorus. Sedimentation ($\text{sedOC, kg/day}$) is calculated as a function of OC concentration in lake water ($\text{conc}$) and lake area ($\text{area}$). The settling velocity parameter $\text{sedoc}$ is general or can be specified for each lake.

$\text{sedOC} = \text{sedoc} \times \text{conc} \times \text{area}$

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Tracers

HYPE can simulate tracers in addition to water flows, nutrients and organic carbon. Tracers are simulated as a substance given in info.txt and follow the water as a concentration. Two tracers are implemented:

- T1 is a general tracer that can have different characteristics that is governed by model parameters
- T2 is water temperature

General tracer (T1)

T1 is the substance name for a general tracer. It can be used to simulate different substances, e.g. patogens or O18 isotope. The tracer is simulated as a concentration in HYPE's water bodies, e.g. snow, soil water, surface water. In addition T1 can be simulated as adsorbed to soil, as a store above ground, and in river sediments.

The following table shows how the unit of different tracers relates to each other and how they are related to input and output parameters and variables. The general unit (U) is used in tables of parameters and input data where the substance is not defined.

<table>
<thead>
<tr>
<th>Variabel/parameter</th>
<th>General unit (U)</th>
<th>Weight (kg)</th>
<th>Number of patogens (#)</th>
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<tbody>
<tr>
<td>Amount in manure/residuals (t1amount)</td>
<td>U/ha</td>
<td>kg/ha</td>
<td>#/ha</td>
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<tr>
<td>Point source concentration (ps_t1)</td>
<td>mU/m3 or µU/L</td>
<td>mg/L</td>
<td>thousandth part/m3 or millionth part/L</td>
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<tr>
<td>Concentration of water as incoming source (cpt1, t1leaksoil/t1leakluse)</td>
<td>mU/m3 or µU/L</td>
<td>mg/L</td>
<td>thousandth part/m3 or millionth part/L</td>
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<tr>
<td>Initial concentration parameters (init1, init1sw)</td>
<td>mU/m3 or µU/L</td>
<td>mg/L</td>
<td>thousandth part/m3 or millionth part/L</td>
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<tr>
<td>Simulated amount T1 in or above soil (aT11, aT12, aT13, sT11, sT12, sT13, T1sf)</td>
<td>U/km2</td>
<td>kg/km2</td>
<td>#/km2</td>
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<tr>
<td>Variabel/parameter</td>
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<td>------------------------</td>
</tr>
<tr>
<td>Simulated amount T1 in river sediment ($T_{smr}$, $T_{slr}$)</td>
<td>U</td>
<td>kg</td>
<td>#</td>
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<td>Simulated concentration of water ($co_{T1}$, $ce_{T1}$, $cs_{T1}$, $cl_{T1}$, $Tc_{r1}$, $Tc_{r2}$, $Tc_{r3}$, $Tc_{rd}$, $Tc_{rs}$)</td>
<td>mU/m3 or µU/L</td>
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<td>thousandth part/m3 or millionth part/L</td>
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<tr>
<td>Observed concentration ($re_{T1}$)</td>
<td>mU/m3 or µU/L</td>
<td>mg/L</td>
<td>thousandth part/m3 or millionth part/L</td>
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</table>

### Sources

There are four different ways to introduce the tracer T1 to the model:

1) T1 may be added as a concentration in precipitation ($cpt_{1}$). This concentration is added as a time series in $X_{obs}.txt$.

2) Point sources may be a source of T1 to surface waters. Point sources of tracer T1 can be added to the local stream, the local lake, the main river or the outlet lake.

3) A pool laid down on land can be a source of the tracer. The pool is defined similar to the manure for nutrients ($t_{1amount}$). It can be added at a specific point in time or equally distributed over a period of time. The pool may be tilled down into the top soil layers also in similar fashion as manure.

4) The tracer can be introduced to HYPE in the form of typical concentrations for leakage from different land-uses and/or soil types. The concentration of the runoff from a class ($conc$) is calculated as the product of two model parameters:

$$conc = t_{1leakluse} \times t_{1leaksoil}$$

One parameter is land use dependent ($t_{1leakluse}$) and one is soil type dependent ($t_{1leaksoil}$).

The source of tracer by a typical concentration (introduction way 4) is not compatible with the sources concentration of precipitation (introduction way 1) and pool on the ground (introduction way 3).

### Processes

In its basic state, T1 act as a tracer passively following the water through the model (except for evaporation). By supplying input data and parameter values this behaviour may be changed to simulate a non-conservative substance, e.g., a patogen.

### Evaporation

As default T1 do not follow evaporating water. This is an exception from the basic assumption that T1 is a conservative tracer if no parameters are set. The tracer will follow evaporating water (with same concentration as the water it is evaporating from) if the model parameter $t_{1evap}$ is set to one. Default is zero. It is possible to set a value between zero and one so that the tracer partly follows evaporating water.
Release from above ground storage

T1 can be released from a storage above ground (e.g. manure) by rain and snowmelt (flow \( q \) (mm/day)). The released tracer will follow the flow, as a concentration, and depending on the fate of the flow, the tracer may follow surface runoff, infiltration or other pathways. The release is governed by the model parameter \( t1rel \) (1/mm). A fraction (\( a_{rel} \)) of the total amount of tracer in the store above ground is released and follows the flow according to equation:

\[
a_{rel} = 1 - e^{-t1rel \times q}
\]

Exponential decay

Pathogens grow and die over time. To simulate this, HYPE supplies a process for the combined effect as an exponential decay. The process is governed by the model parameter \( t1expdec \), which denote the halv life time of the tracer in days. Exponential decay is applied to tracer T1 in most forms; in soil water, river, lakes, the above ground storage, tracers adsorbed to soil and tracers in river sediment.

Adsorption/desorption

Tracer adsorption/desorption to/from soil particles is calculated based on an equilibrium concentration (\( c_{eq} \) (e.g. number of patogens/L or U/L)) of soil water. The equilibrium concentration is assumed to be reached within one time step of the model, typically one day, and this gives the amount of tracer adsorbed or desorbed during the time step is calculated. The equilibrium concentration (\( c_{eq} \)) is calculated from the amount of tracers and soil characteristics. The equation is:

\[
c_{eq} = \frac{t1freuc \times Atot}{v + t1freuc \times b_{density} \times d}
\]

where \( Atot \) is the sum of tracer adsorbed and solved in soil water of the soil layer (U/km2), \( v \) the volume of water in the soil layer (mm), \( b_{density} \) is soil density (1300 kg soil/m3), and \( d \) is the thickness of the soil layer (m). The general model parameter \( t1freuc \) is an equilibrium concentration coefficient. In the case of pathogens, the model parameter describes the number of pathogens adsorbed to one kg of soil per pathogens in one litre of soil water (or in general unit (U/kg soil)/(U/L)).

Sedimentation/resuspension in river

Sedimentation or resuspension of tracers in rivers is calculated the same way as particulate phosphorus (described here). The process depends on the current flow (\( q \)) in the river in relation to bankful flow (\( q_{bank} \)) and the general model parameter \( t1sedexp \). Bankful flow is the flow when the river is filled to the brim. This flow is calculated as the second largest simulated flow in the last year. The fraction of the tracer that is resuspended or sedimented is determined by the equation:
For low flow, the fraction \( a_{sres} \) is positive and sedimentation \( F_{sed} \) (U/day) occur, for high flow the fraction is negative and there will be resuspension from the sediments instead \( F_{resusp} \) (U/day):

\[
F_{sed} = a_{sres} \times c_{river} \times v_{river}, a_{sres} > 0
\]

\[
F_{resusp} = -a_{sres} \times Ased, a_{sres} < 0
\]

where \( c_{river} \) is the concentration of T1 in river (U/m³), \( v_{river} \) is the volume of river (m³), and \( Ased \) is amount of tracer T1 in the sediment (U).

### Sedimentation in lakes

Sedimentation of tracer T1 in lakes is calculated the same way as e.g. particular phosphorus. The sedimentation is governed by sink velocity (m/d), which is given by the general model parameter \( t1sedvel \).

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### Links to relevant procedures in the code
Water temperature (T2)

Water temperature can be simulated by HYPE with the substance/tracer called T2. Water temperature is suitable to be simulated in a catchment model, because it relates to the hydrology and the landscape. The substance follows the water paths in HYPE and is calculated in degree Celsius.

Sources

There are different ways to introduce heat into the water of HYPE.

Initial value of water temperature of HYPE's water bodies can be set by model parameters. The general parameter iniT2 is used for some water bodies, i.e. soil, rivers and lakes. Aquifer temperature is given in AquiferData.txt (temp).

Snow and glacier melt water is assumed to hold zero degrees.

Precipitation

Temperature of precipitation on land is set to air temperature (but limited to zero).

For water surfaces the temperature setting is more complex. Temperature of rain on water is set to air temperature (but limited to zero). Snowfall are assumed to have the average temperature of the water (uppertemp for lakes, mean temperature for rivers) for ice conditions, i.e. not to influence the temperature of the water. For snowfall on open water the temperature is calculated as dependent on sensible heat content and latent heat of freezing, taken into account different specific heat capacity of ice and water. Temperature of precipitation is a weighed average of these three components.

Point source

Point sources influence the water temperature T2. Point sources may have a constant temperature set from indata (ps_t2) or otherwise the water is assumed to be zero degrees. Point sources of tracer
T2 can be added to the local or main river, or internal or outlet lake.

**Local diffuse source**

The water of local diffuse source is not allowed to influence the water temperature of the receiving water body (soil or local river).

**Soil processes**

The primary process for changing water temperature is exchange of heat over a water surface. Apart from that, surface water temperature is determined by mixing water of different temperature. Exceptions have been made for some processes not to influence the water temperature. This because their effect on water temperature has not yet been included in full.

**Soil temperature**

In addition to the “concentration” of T2 in soil water, the (solid) soil layer has their own temperature state variables. Temperature of soil, i.e. solid material, is calculated separately from the T2 temperature of soil water. The temperature of a soil layer depends on the air temperature and the deep soil layer temperature to different degree. Deep soil layer temperature is calculated separately. See Chapter Soil temperature and snow depth.

Temperature T2 of soil water is not yet developed and is simply set to the temperature of soil. Still, snow melt are assigned the temperature T2 zero. Evaporation of soil water is not affected by T2 or affecting soil water T2.

**Runoff**

Soil runoff temperature is set from soil layer temperatures. Tile drainage temperature is set to the temperature of the soil layer it originates from. Surface runoff gets the temperature from soil layer one (saturated overland flow) or rainfall, snowmelt and glaciermelt (excess infiltration).

**Lake and river ice**

The river ice routines are performing best if the special classes for local and main river is set. The routines then use the forcing data and snow parameters defined for these classes. If the river classes is not set, forcing and snow parameters for lakes are used as a replacement. If no lake classes are provided, forcing and snow parameters are taken from slc class one, whatever that is.

**Floating snow covered ice sheet model**

Lake and river ice is simulated with a simple thermodynamic ice model driven by air temperature and precipitation. It is to a large extent adopted from the models presented by Leppäranta (1983, 1993),
in which lake and river ice is represented by a floating snow covered ice sheet characterized by three layers of frozen water: black ice, snow ice, and snow (Figure 2, left). If the top of the ice layer is submerged under water, the depth of a slush layer (wet snow) is also defined (Figure 2, right).

Figure 2: A floating snow covered ice sheet with (left) three layers of frozen water black ice, snow ice and snow and (right) with a slush (wet snow) layer formed when the top of the ice is submerged under water due to the load of the snow pack.

**Influence of the ice model on lake and river evaporation and water temperatures**

It should be noted that in the current version of the HYPE model, the ice mass balance is independent from the mass balance of the lakes and rivers. No water is removed from lake or river water to form the ice cover and no water is added to the lake and river when the ice is melting. This simplification is reasonable only for rivers and lakes large enough to never freeze to the bottom. The ice model influences lake and river water only through its' impact on water temperatures and evaporation:

- evaporation from lakes and rivers is set to 0 in the presence of ice
- heating and cooling of lakes and rivers through surface heat exchange with the air is set to 0 in the presence of ice
- lake and river water temperature is adjusted to take into account the inflow of 0°C melt water during ice melt

Water temperature is also adjusted to take into account the heat needed to melt snowfall on ice-free lakes and rivers.

**Black ice growth**

\[
\frac{dH_i}{dt} (\text{cm/s}) \]

Black ice growth \((\text{cm/s})\) is derived from the modified Stefan’s equations as described by Leppäranta (1983):

\[
\frac{dH_i}{dt} = \frac{k_i}{\rho_i \times L_f} \times \frac{\left(T_f - T_a\right)}{\left(H_i + H_s \times \frac{k_i}{k_e} + \frac{k_i}{k_a}\right)}
\]

The first part is constant in HYPE. \(k_i\) is thermal conductivity of ice (0.022 J/°C/cm/s), \(\rho_i\) is density of ice (0.917 g/cm³), and \(L_f\) is latent heat of freezing (334 J/g).

\(T_f\) is freezing temperature of lake or river water (°C), a parameter. \(T_a\) is air temperature (°C). \(H_i\) is ice
depth (cm) and $H_s$ is snow depth on the ice (cm). $k_a$ is heat exchange coefficient with atmosphere (J/°C/cm²/s). $k_s$ is thermal conductivity of snow (J/°C/cm/s). It is calculated from thermal conductivity of ice and snow and ice density, and a parameter $k_{exp}$:

$$k_s = k_i \times \left( \frac{\rho_s}{\rho_i} \right)^{k_{exp}}$$

where $\rho_s$ is density of snow (g/cm³). Snow density changes over time depending on a change parameter ($sndens$).

The heat released when the black ice is formed at the bottom of the ice layer $(\rho_i \times L_f \times \frac{dH_i}{dt})$ is conducted through the ice and the overlaying snow layer to the atmosphere. This heat flow is driven by the temperature gradient between the air above the ice or snow surface and the freezing temperature of the water assumed at the bottom of the ice layer, and is governed by the depths and thermal conductivities in the ice and snow layers, and the heat exchange coefficient in the air. It should also be noted that a term representing the heat flow from the water has been neglected compared to Leppäranta (1983).

The black ice growth equation is only used if the ice surface is above the water surface (in other words, if there is no slush layer on top of the ice to be frozen first) and if air temperature is below the freezing temperature $T_f$.

### Snow ice growth

If the mass of the snow layer exceeds the floating capacity of the ice, the bottom of the snow layer will be submerged in lake water, and a slush layer ($H_{sl}$) will form with a depth according to:

$$H_{sl} = \frac{H_s \times \rho_s - H_i \left(1 - \rho_s \right)}{\rho_s / \rho_i}$$

As pointed out by Leppäranta (1993), the freezing of slush into snow ice releases less heat compared to black ice growth since the slush already consists of frozen snow particles, and this heat only need to be conducted through the snow pack. The growth of snow ice can thus be calculated by the following equation from Leppäranta (1993):

$$\frac{dH_{bi}}{dt} = \frac{k_s}{\left( \rho_i \times L_f \left(1 - \frac{\rho_s}{\rho_i} \right) \right)} \times \left( T_f - T_\alpha \right)$$

$$\left( H_s + \frac{k_i}{k_a} \right)$$

where all variables are defined above for black ice growth.
Ice melt and final ice break-up

Snow and ice melt is calculated with a simple temperature index model when air temperature is above 0°C. The snow on the ice has to be completely melted before any ice melt is calculated:

\[
\frac{dH_i}{dt} = -p_m \times T_a
\]

Ice can be melted from below if the water contain enough heat. This process is limited by a melt efficiency parameter \((m_{eff})\).

Final ice break-up is defined as the time step when \(H_i = 0\). Although it is well known that ice tend to fall apart before it is completely melted, this is not considered by the present model.

Initiation of ice growth

Ice growth is initiated when the solution of the surface heat balance for the open water results in a water temperature below the freezing point. The fraction of the water surface with new ice formation is then estimated by reducing the open water surface area until the surface heat balance solution results in a water temperature equal to the freezing point. The transition from fully ice-free to fully ice-covered conditions is thus smoothed by the gradual increase in fractional ice cover. For deep lakes, the temperature of an upper layer (EPI, Figure 3 below) is used for the onset of ice growth, whereas for shallow lakes and rivers, the mean water temperature is used.

Surface water processes

Temperature \(T_2\) of a river is not affected by irrigation withdrawal, abstractions, rural households or point source additions, constructed wetlands, or water returning from aquifers.

Lake basic assumptions

For substances the lake volume is divided into two parts FLP and SLP (see Chapter Rivers and lakes - Basic assumptions)). In addition to simulating temperature as the substance \(T_2\) in the two lake parts, states of lake temperature in a hypothetical epilimnion and hypolimnion \((\text{uppertemp, lowertemp})\) is handled in the model. The average temperature of FLP+SLP is equal to the average temp of the EPI+HYPO. The size of volumes related to the upper and lower temperatures are determined by the thermocline which is estimated for each olake.
Thermocline depth is estimated from lake area (Hanna, 1990). This average depth is adjusted for current changes by adding precipitation ($\text{prec}$) and inflow ($\text{qin}$) to the lake, and remove evaporation ($\text{evap}$).

$$\text{epidepth} = 6.95 \times \text{lakearea}^{0.185} + \text{prec} - \text{evap} + \text{qin}$$

where $\text{lakearea}$ is (km$^2$) and $\text{epidepth}$ is in m below lake surface. Precipitation, evaporation and inflow is also in meter.

The adjusted depth ($\text{epidepth}$) is then used as the division of the lake volume into upper and lower if the lake water depth is larger than the $\text{epidepth}$. Otherwise the lake is treated as one single mixed volume in regard to T2 calculations.

**Evaporation**

Lakes and rivers with given class area may have evaporation. The actual evaporating area is reduced if it is ice covered. Evaporation is not changing the T2 temperature of the river or lake. The latent heat is assumed to be taken care of by the water - atmosphere T2 exchange routine. River evaporation is removed proportionally from the water stores making up the river volume. Lake evaporation may be removed from one or both lake parts (FLP and SLP).

Evaporation from flooded floodplain of rivers or lakes also keep the temperature of the flooded water unchanged.

**Water - atmosphere T2 exchange**

River and lake water atmosphere exchange is calculated during ice free conditions. Water surface heat balance is calculated based on a model by Piccolroaz et al (2013), with modifications for fractional ice cover, and calculation of fractional freezup area. This equation depends on four general parameters; $\text{tcf}$-temperature difference coefficient, $\text{scf}$ - solar radiation coefficient, $\text{ccf}$ - constant coefficient and $\text{lcf}$ - linear coefficient. The parameters comes in one set for rivers ($\text{tcfriver}$, $\text{scfriver}$, $\text{ccfriver}$, $\text{lcfriver}$) and one set for lakes ($\text{tcflake}$, $\text{scflake}$, $\text{ccflake}$, $\text{lcflake}$).

A simpler T2 exchange model based only on the temperature difference between air and water is also available. It has two general heat transfer parameters ($\text{t2trlake}$, $\text{t2trriver}$, unit J/m$^2$/s/degree).
For rivers with no specified river area as a class, a river area is calculated for use in the water-atmosphere T2 exchange and ice processes. The river area is calculated as described in Section Nitrogen and phosphorus processes in rivers and lakes. In addition the river area may be reduced at low volume/flow. This is controlled with parameters (see River and Lakes - Evaporation).

**T2 stratification**

At spring and autumn circulation, T2 temperature is averaged out between upper and lower temp. During periods with stratification a heat transfer between the upper temp and lower temp is assumed. This heat transfer depend on the general parameter upper2deep (J/m²/s/degree).

**Lake outflow**

Lake outflow T2 temperature may be determined by the T2 of the lake parts (FLP and SLP) contributing to outflow (see Figure 2). To use this method set parameter t2mix. For ilakes the t2mix method is always used.

For outlet lakes another method for determining lake outflow temperature is the default method. This method assigns upper temp to outflow water if the lake is stratified and the thermocline is deeper than the lake threshold (thres). For other conditions a mixture of upper and lower temp is used to set the outflow temperature (t2out).

\[
\text{t2out} = \text{fraction} \times \text{upper temp} + \left(1 - \text{fraction}\right) \times \text{lower temp}
\]

\[
\text{fraction} = \frac{\text{epidepth}}{\left(\text{lakewst} - \text{thres}\right)}
\]

where lakewst is the current depth of the lake.

**References**


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Sediment

Introduction

Sediments are simulated as suspended sediments (SS) and nitrogen in algae (AE). The sum of the two substances is an additional output (total suspended sediments, TS). Suspended sediments are not simulated in the soil of HYPE. Suspended sediments are first introduced into the runoff of the soil by soil erosion.

The main states are concentration of SS and AE in all water stores of HYPE, i.e. soil, river, lakes, but also snow, aquifers (in the store and on the move there or away), irrigation water, floodplain water, and water transfer (that is delayed one time step) could hold suspended sediments and algae. In the current model though, concentration of SS and AE is only positive in river and lakes, and flows originating from them. In addition two pools of settled sediment are simulated; a pool of delayed sediment in runoff, and a pool of (temporarily) settled sediment in river. Note: No store of “settled sediment” of the soil is simulated, i.e. HYPE has an unlimited source of soil for erosion.

Sources of sediment load

Soil erosion

The main source of sediment in HYPE is from soil erosion. Soil erosion is modelled in several steps. First particles are mobilized from the soil by rain or surface runoff. The result is here called “mobilized sediments”. The mobilized sediments are assumed suspended in an “eroding flow”, which is surface runoff plus macropore flow. Secondly, if there is surface runoff, the mobilized sediments is transported away from the field with the surface runoff and macropore flow (“eroded sediment”). A filtering can be applied to reduce the amount of eroded sediment. If there is no surface runoff, there will be no transport of eroded sediments. For SS, the eroded sediments are taken from an infinite pool. For PP, the eroded particulate phosphorus is taken from the soil pools. Thirdly, the eroded sediments in macropore flow and surface runoff (and tile runoff though that is generally zero) are delayed in a temporary pool. The release of suspended sediments from this pool is determined by the
total runoff from the class, and is following the total runoff off the land.

**Step 1: Mobilization of particles from soil**

HYPE has two alternative models for mobilization; soil erosion model 1 and soil erosion model 2.

**Soil erosion model 1 (MMF-based model):** The first model is based on Morgan-Morgan-Finney erosion model (Morgan et al., 1984) and calculates particles mobilized by rainfall energy and surface runoff. The kinetic energy in rainfall is calculated as a function of rainfall \( \text{rain, mm/ts} \) and day of the year \( \text{dayno} \). If the precipitation falls as snow, or if it falls on snow-covered ground or if it is smaller than 5 mm/day no mobilization occurs in the model. Some of the raindrop's energy can be absorbed by vegetation. Crop cover is defined as the portion of land that is sheltered from raindrops; for a description of how this is calculated, see Chapter Crop cover and ground cover. The factor \( \text{common} \) is the sheltering effect that the main and secondary crops give together. It varies over the year due to crop growth and management. The mobilization by rain \( \text{MobilisedRain, g/m}^2\text{ts} \) is also influenced by soil erodibility (soil dependent parameter \( \text{soilerod (g/J)} \)).

\[
\text{Rain fall energy} = \text{rain} \times \left( 8.95 + 8.44 \times \log_{10} \left( \text{rain} \times 2 \times \left( 0.257 + 0.90 \times \sin \left( 2 \times \left( \text{dayno} - 70 \right) / 365 \right) \right) \right) \right)
\]

\[
\text{MobilisedRain} = \text{Rain fall energy} \times \left( 1 - \text{common cropcover} \right) \times \text{soilerod}
\]

When surface runoff occurs, soil particles are eroded and carried away as the soil surface is exposed to shear forces. The mobilization \( \text{MobilisedSR, g/m}^2\text{day} \) is calculated from the surface runoff \( \text{sflow, mm/day} \), subbasin average slope, a parameter for soil cohesion \( \text{soilcoh (kPa) soil type dependent} \), and a general parameter \( \text{sreroexp} \). This type of erosion can be mitigated by protective vegetation or vegetation residues that are in contact with the ground. The calculation of this reducing factor \( \text{groundcover} \) is described in Chapter Crop cover and ground cover. The factor \( \text{common groundcover} \) is the combined effect of the main and secondary crops.

\[
\text{MobilisedSR} = \frac{\left( \text{sflow} \times 365 \right)^{\text{sreroexp}} \times \left( 1 - \text{common groundcover} \right) \times \frac{1}{0.5 \times \text{soilcoh}} \times \sin \left( \frac{\text{slope}}{100} \right)}{365}
\]

All mobilized particles are not staying mobilized, because sometimes the transport capacity of the particle-bearing water \( \text{eflow} \) will not suffice for the task. If this is the case, a transport factor reduces the particle amount mobilized:

\[
\text{transport factor} = \text{MIN} \left( 1.0, \left( \frac{\text{eflow}}{4} \right)^{1.3} \right)
\]

Finally mobilized sediment \( \text{mobilSed, kg/km}^2\text{day} \) is calculated as the sum of rain and surface runoff caused mobilization as:
Soil erosion model 2 (HBV-SED based model): The second model is based on HBV-SED model (Lidén, 1999; Lidén et al., 2001) and calculates particles mobilized by rain (\(\text{rain}\)). When the ground is frozen or there is snow on the ground, the no particles are mobilized. The mobilization also depends on soil and land characteristics in the form of model parameters and data on subbasin characteristics.

\[
mobil\text{Sed} = 1000 \times \left( \frac{\text{erosslope}}{5} \right) \times \text{erodluse} \times \text{erodsoil} \times \frac{\text{EI}}{\text{erodindex}} \times \text{rain}^{\text{erodexp}}
\]

The parameters \(\text{erosslope}\), \(\text{erodexp}\) and \(\text{erodindex}\) are general and thus same for the whole model domain. The parameters \(\text{erodluse}\) and \(\text{erodsoil}\) are land-use and soil type dependent. Subbasin input are: \(\text{slope}\), the subbasins' average slope, and an erosion index, \(\text{EI}\). The many parameters give the possibility to simulate erosion as dependent on slope, soil type, land use or subbasin.

**Step 2: Transport of eroded sediments off the field**

If at one time step there is no surface runoff, there will be no transport of suspended sediment. Still, calculation continues with step 3, and earlier delayed eroded sediment may reach the local stream. If there is surface runoff, a fraction of the mobilised sediments is assumed to go with surface runoff (\(\text{sflow}\)). If there is surface runoff and tile runoff and macropore flow, a fraction of the mobilized sediments is assumed to travel with the macropore flow (\(\text{mflow}\)). The respective fractions depend on the respective size of the two flows, but are reduced by filtering of the particles. The filtering depends on landuse, the proximity of agricultural land to water, presence of buffer strips etc. The filtering effect on suspended sediments in surface runoff (\(\text{srfilt}\)) is parameterized with land use parameters and subbasin input, while the filtering effect on suspended sediments of macropore flow is parameterized with a soil type parameter (\(\text{macrofilt}\)).

\[
\text{sr filt} = \text{other filt} + \text{alpha} \times \left( 1 + \text{bufferpart} \times \left( \text{buffer filt} - 1 \right) \right) + \text{inner filt} \times \left( 1 - \text{alpha} \right)
\]

\[
\text{erodedSed} = \left( \text{sr filt} \times \text{s flow} + \text{macro filt} \times \text{m flow} \right) \times \text{mobilSed}
\]

\[
\text{e flow} = \text{s flow} \times \text{m flow}
\]

The total amount of eroded sediment (\(\text{erodedSed}\), kg/km2/day) is the sum of contributions from surface flow and macropore flow. For SS, the eroded sediments are taken from an infinite pool. For PP, the eroded particulate phosphorus is calculated from the eroded sediment (see Soil erosion) and then taken from the solid soil pools (\(\text{humusP}\) and \(\text{partP}\)). This difference exists because HYPE does not simulate any pool of “soil sediments”.

**Step 3: Suspended sediment reaching the local stream**

The concentration of suspended sediments of the runoff leaving the soil is calculated based on total runoff (\(\text{runoff}\), mm/ts) and a pool of sediment particles (\(\text{relpool}\)). The pool collects the suspended sediments in the respective runoff pathways, i.e. primarily the eroded sediment from surface runoff and macropore flow calculated in the previous step. The pool constitutes a temporary delay. The
release of sediment \((\text{release} \, \text{kg/km2})\) from the pool temporary is calculated with two general parameters \((\text{pprelmax} \, \text{and} \, \text{pprelexp})\):

\[
\text{release} = \text{MIN}\left(\text{relpool}, \text{relpool} \times \left(\frac{\text{runoff}}{\text{pprelmax}}\right)^{\text{pprelexp}}\right)
\]

The suspended sediments released are following the total runoff off the land. The release divided by the total runoff gives a suspended sediment concentration, which is used for all runoff pathways. This concentration is set to all runoff from the class, i.e. the sediments are transported by all flow paths off the land.

**Rural household outlets and point sources**

Suspended sediments may be added to the model as diffuse rural households outlets (private sewers) or point sources. Similar to other substances in HYPE, they are added as flow with concentration of the total suspended sediments and with a fraction determining the suspended part, while nitrogen in algae make up the rest \((\text{GeoData.txt PointSourceData.txt})\). To add only suspended sediments set the fraction to 1. The diffuse source is divided into two parts, where the division is determined by a general parameter \((\text{locsoil})\). One part is added directly to the local river. The other part is added to the water of the bottom soil layers of each class in the subbasin. The distribution between classes is done proportional to the area of each class. Point sources are added in the main river. Point sources are described in the Chapter on Water management \((\text{point sources})\) and rural household diffuse source in the Chapter on Nitrogen and phosphorus in land routines \((\text{rural sources})\).

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**Soils**

Suspended sediments are not simulated in the soil calculations of HYPE. Suspended sediments are first introduced into the runoff of the soil.

**Sediment in rivers and lakes**

**Primary production and mineralization**

Primary production in lakes and rivers is affecting sediment simulation by being a source of algae organic nitrogen (AE). Mineralisation, as a sink of AE, is the reverse process. Nitrogen in algae is assumed to grow and decline with the same function as production and mineralisation of organic nitrogen (ON). The production and mineralisation processes are modelled together and only one is active at the time. If nitrogen is simulated by HYPE it uses the actual estimated production/mineralisation for AE, otherwise the potential production/mineralisation (minprodNpot) is used for AE. The actual production is limited by the availability of inorganic nutrients. If sediments are simulated without simultaneously simulating nitrogen, the mineralisation of algae is limited to available amount, but the production is unlimited.

The potential production/mineralisation is depending on temperature ($T_w$, $T10$, $T20$), long-term total phosphorus concentration ($TP$), surface area of water body ($area$) and a parameter ($wprodn$).

$$minprodNpot = wprodn \times TPfcn \times tmpfcn \times area$$

$$TPfcn = \frac{TP_{limsedPP}}{(TP_{limsedPP} + hsatTP)}$$

$$tmpfcn = tmpfcn1 \times tmpfcn2$$

$$tmpfcn1 = \frac{T_w}{20}$$

$$tmpfcn2 = \frac{(T10 - T20)}{5}$$

Temperatures $T10$ and $T20$ are calculated as the average water temperature ($T_w$) of 10 and 20 days. They determine if production or mineralization dominates at the current time. The water temperature is calculated through weighting the air temperature ($T$) and yesterday's water temperature (see basic
assumptions of rivers and lakes).

The phosphorus function \((TPfcn)\) is dependent on a general half saturation parameter \(hsatTP\), and a limitation parameter \(limsedPP\). If phosphorus is simulated by HYPE the long-term average concentration \((TP)\) is calculated from that, otherwise a lakeregion parameter is used \((tpmean)\).

**Sedimentation**

Sedimentation in lakes is a sink for SS and AE and works the same way as for nutrients. Sedimentation \((sed, \text{kg/day})\) is calculated as a function of concentration in lake water \((conc)\) and lake area \((area)\). The settling velocity parameter \((sedss, sedae)\) is general. The concentration used in the equation may be limited \((lim)\) by general parameter \((limsedSS)\) for SS, but not for AE \((lim=0)\).

\[
sed = \text{par}_\text{sed} \times \left(\frac{conc}{lim}\right) \times area
\]

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