

## MOHID Modelling System Description



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# 1 General Overview

## 1.1 Introduction

This document describes the three-dimensional water modeling system *Mohid*. The *Mohid* system is composed by a free surface three dimensional baroclinic hydrodynamic module, a eulerian transport module, a lagrangian transport module, a turbulence module, a zero-dimensional water quality module and an oil dispersion model.

Actually the principal investigators are Ramiro Neves, Paulo Chambel Leitão and Frank Braunschweig, from the Technical University of Lisbon. Contributions from Henrique Coelho, Manuel Villarreal (Turbulence) and Pedro Pina (Water Quality) are also included in the model. A lot of other investigator have given there contribution to this model over the past years.

## 1.2 History

The development of the *Mohid* model started back in 1985, passing since this time through continuously updates and improvements due to its use during different projects of scientific research and engineering projects. Initially the *Mohid* water modeling system was a bi-dimensional hydrodynamic model, called *Mohid 2D* (Neves, 1985). This model was used to study estuaries and coastal areas using a classical finite difference approach. In the following years, a bi-dimensional eulerian and lagrangian transport model were included in this model. The first three-dimensional was introduced by Santos (1995), which used a vertical double Sigma coordinate. This version was called *Mohid 3D*. The limitations of the double Sigma coordinate revealed the necessity to develop a model which could use a generic vertical coordinate, permitting the user to choose the type of vertical coordinate, depending on the study area. Due to this necessity the concept of finite volumes was introduced with the version *Mesh 3D* by Martins (1999). In the *Mesh 3D* model were included a three dimensional eulerian transport model, a three dimensional lagrangian transport model (Leitão, 1996) and the zero-dimensional water quality model (Miranda,

1999). Since the introduction of the finite volumes approach, this discretization remains in the model *Mohid*.

### 1.3 Actual State

With the growing model complexity, it was necessary to introduce a new way in the organization of the information of the *Mohid* model. In 1998 the whole code was submitted to a complete rearrangement, using new the feature of programming languages and also the capacities of the computer to reprogram the whole *Mohid* model. The main goal of this rearrangement was to turn the model more robust, reliable and protect its structure against involuntary programming errors, so it would be more easily “grow able”. To achieve this goal, objected oriented programming in FORTRAN was introduced to the *Mohid* model, like described in Decyk (Decyk, *et al.*, 1997).

The philosophy of the new *Mohid* model (Miranda, *et al.*, 2000), further on simple designated *Mohid*, permits to use the model in any dimension (one-dimensional, two-dimensional or three-dimensional). The whole model is programmed in ANSI FORTRAN 95, using the objected orientated philosophy. The subdivision of the program into modules, like the information flux between these modules was object of a study by the *Mohid* authors.

Actually the model *Mohid* is composed by over 40 modules which complete over 150 mil code lines. Each module is responsible to manage a certain kind of information. The main modules are the modules listed in Table 1.3-1.

Another important feature of *Mohid* is the possibility to run nested models. This feature enables the user to study local areas, obtaining the boundary conditions from the “father” model. The number of nested models is just limited by the available computer power.

---

<i>Module Name</i>	<i>Module Description</i>
Model	Manages the information flux between the hydrodynamic module and the two transport modules and the communication between

---

	nested models.
Hydrodynamic	Full 3D dimensional baroclinic hydrodynamic free surface model. Computes the water level, velocities and water fluxes.
Water Properties (Eulerian Transport)	Eulerian transport model. Manages the evolution of the water properties (temperature, salinity, oxygen, etc.) using a eulerian approach.
Lagrangian	Lagrangian transport model. Manages the evolution of the same properties as the water properties module using a lagrangian approach. Can also be used to simulate oil dispersion.
Water Quality	Zero-dimensional water quality model. Simulates the oxygen, nitrogen and phosphorus cycle. Used by the eulerian and the lagrangian transport modules. Based on a model initially developed by EPA (Bowie, <i>et. al.</i> , 1985).
Oil Dispersion	Oil dispersion module. Simulates the oil spreading due thickness gradients and internal oil processes like evaporation, emulsification, dispersion, dissolution and sedimentation.
Turbulence	One-dimensional turbulence model. Uses the formulation from the GOTM model.
Geometry	Stores and updates the information about the finite volumes.
Surface	Boundary conditions at the top of the water column.
Bottom	Boundary conditions at the bottom of the water column.
Open Boundary	Boundary conditions at the frontier with the open sea.
Discharges	River or Anthropogenic Water Discharges
Hydrodynamic File	Auxiliary module to store the hydrodynamic solution in an external file for posterior usage.

**Table 1.3-1: Principal modules of the model Mohid**

## 1.4 Applications

The MOHID model has been applied to different coastal and estuarine areas and it has showed its ability to simulate complex features of the flows. Several different coastal areas have been modeled with MOHID in the framework of research and consulting projects. Along the Portuguese

coast, different environments have been studied, from river mouths (Douro, Mondego) to coastal lagoons (Ria de Aveiro, Ria Formosa) and broader estuaries (Tagus and Sado estuaries (Martins et al. [2000])). Also the model has been adapted to simulate Galician Rías hydrodynamics (Ría de Vigo (Taboada et al., [1998], Montero et al. [1999], Montero, [1999]), Ría de Pontevedra (Taboada et al.[2000], Villarreal et al. [2000]) and other Rías (Pérez Villar et al. [1999])). Far from the Atlantic coast of the Iberian Peninsula, some European estuaries have been modeled -Western Scheldt (Holland), Gironde (France) (Cancino and Neves, [1999]) and Hellingford (Leitão, [1997])-as well as some coastal estuaries in Brasil (Santos, Fortaleza). Regarding to open sea, MOHID has been applied to the North-East Atlantic region where some processes including the Portuguese coastal current (Coelho et al. [1994]), the slope current along the shelf (Neves et al. [1998]) and the generation of internal tides (Neves et al. [1998]) have been studied and also to the Mediterranean Sea to simulate the seasonal cycle (Taboada, [1999]) or the circulation in the Alboran Sea (Santos, [1995]). More recently *Mohid* has been applied to the Portuguese Monte Novo, Roxo and Alqueva reservoirs (Braunschweig, 2001).

## 2 The Model Module

### 2.1 Introduction

The module Model is the topmost module of the *Mohid* water modeling system and has two main responsibilities, the coordinates of the execution of the hydrodynamic module and the transport modules and the coordination of the father-son communication between nested models. Figure 2-1 shows this coordination.

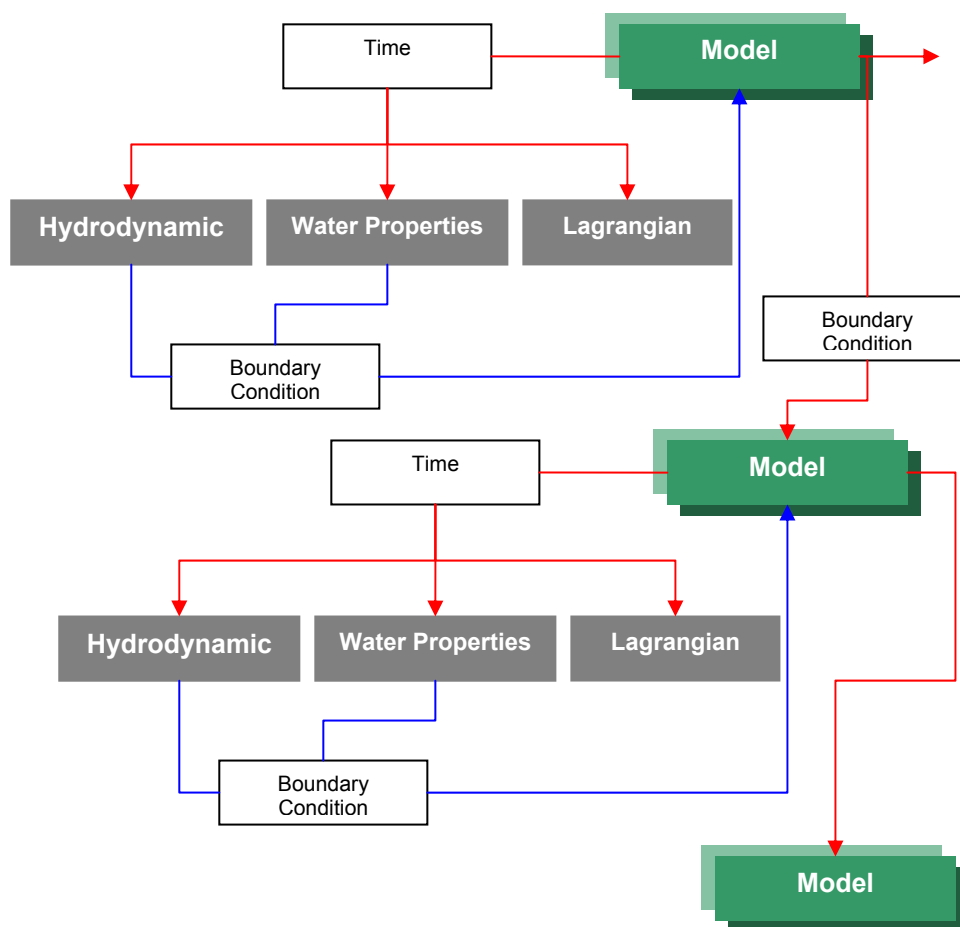


Figure 2-1: Information flux between the nested models

### 2.2 Coordination of the execution of one model

The coordination of the execution of one model consists of the actualization of the global model time and the update of the hydrodynamic

module and the transport models within one model. The transport modules can run with different time steps of the hydrodynamic module (since the time steps of the transport modules are multiplies of the hydrodynamic time step).

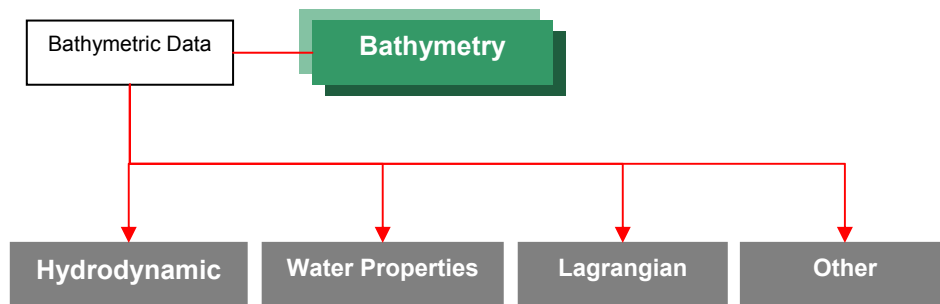
### **2.3 Coordination of the father son communication**

The coordination of the information flux between the nested models includes the synchronization between different nested models, once nested models can also run with different time steps. The coordination of the nested models is done in a hierarchical way. Every model can have one or more nested child models, which recursively can have one or more child models. All the communication is done in one way direction, passing the boundary conditions from the father model to the son(s) model(s).

## 3 The Bathymetry Module

### 3.1 Introduction

The module Bathymetry is one of the lowest modules of the *Mohid* water modeling system. It has basically reads the bathymetric data from an ASCII input file and publishes this data to all client modules.



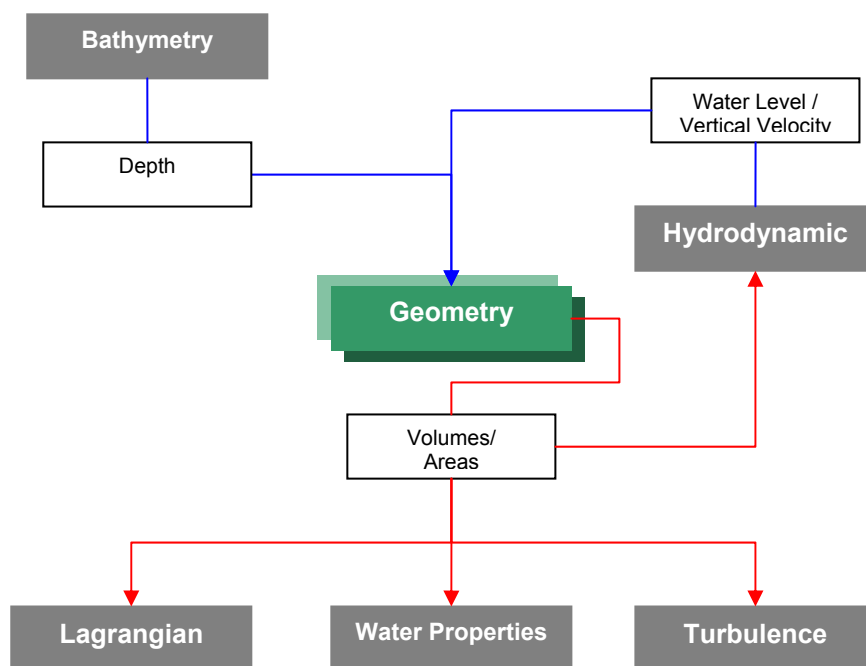
**Figure 3-1: Information flux between the module bathymetry and other modules**

The bathymetric data can be stored in any regular grid, with variable spacing along the X and the Y direction. For every grid point the depth of this point must be given. The horizontal coordinates can be supplied in different types of coordinates. Most common used are metric coordinates and geographic coordinates.

## 4 The Geometry Module

### 4.1 Introduction

The Geometry Module computes the lateral areas and volumes of the finite volume, based upon the surface elevation and the bathymetric data. This information is updated as needed, and published to the other modules of the *Mohid* model. Figure 4-1 represents the information flux between the geometry module and other modules.



**Figure 4-1: Information flux between the Geometry Module and other modules**

### 4.2 Finite Volume

The model *Mohid* uses a finite volume approach (Chippada et al. [1998]; Martins et al. [1999], [2000]) to discretize the equations. In this approach, the discrete form of the governing equations is applied macroscopically to a cell control volume. A general conservation law for a scalar  $U$ , with sources  $Q$  in a control volume  $\Omega$  is then written as:

$$\partial_t \int_{\Omega} U d\Omega + \oint_S \vec{F} d\vec{S} = \int_{\Omega} Q d\Omega \quad \text{Eq. 4-1}$$

where  $F$  are the fluxes of the scalar through the surface  $S$  embedding the volume. After discretizing this expression in a cell control volume  $\Omega_j$  where  $U_j$  is defined, we obtain:

$$\partial_t (U_j \Omega_j) + \sum_{\text{faces}} \vec{F} \cdot \vec{S} = Q_j \Omega_j \quad \text{Eq. 4-2}$$

In this way, the procedure for solving the equations is independent of cell geometry. Actually, the cell can have any shape with only some constraints (see *Montero [1999]* or *Martins [2000]*) since only fluxes among cell faces are required. Therefore, a complete separation between physical variables and geometry is achieved (*Hirsch, [1988]*). As volumes can vary in the course of the calculus, geometry is updated in every time step after computing the physical variables. Moreover, the spatial coordinates are independent, and any geometry can be chosen for every dimension. Cartesian or curvilinear coordinates can be used in the horizontal and a generic vertical coordinate with different sub-domains can be used in the vertical. This general vertical coordinate allows minimizing the errors of some of the classical vertical coordinates (Cartesian, sigma, isopycnal) as pointed in (*Martins et al. [2000]*).

The volume element used in the model MOHID is shown in Figure 4-2. Only a vertical degree of freedom is allowed, and the grid is Cartesian orthogonal in the horizontal. The grid is staggered in the horizontal in an Arakawa C (*Arakawa and Lamb, [1977]*) manner, i.e. horizontal velocities are located in the center of the west (u-velocity) and south (v-velocities) faces, while elevation, turbulent magnitudes and tracers are placed in the center. Also a staggering in the vertical is used, with vertical velocity  $w$ , tracers and turbulent magnitudes vertically placed in the top and bottom faces and horizontal velocities and elevation in the center of the element (in vertical).

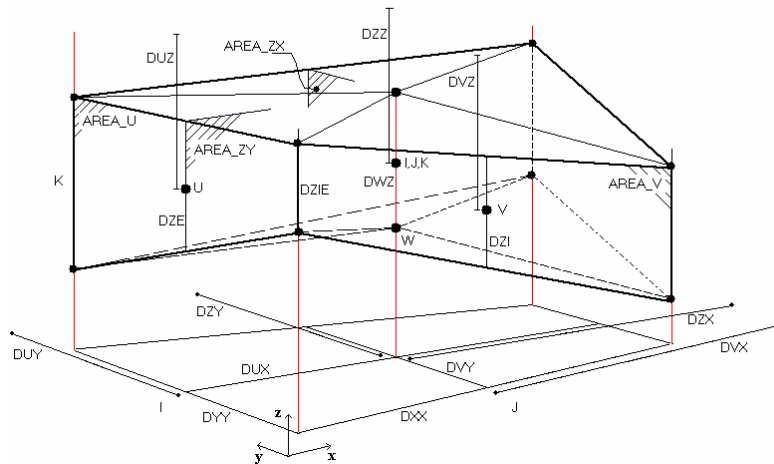
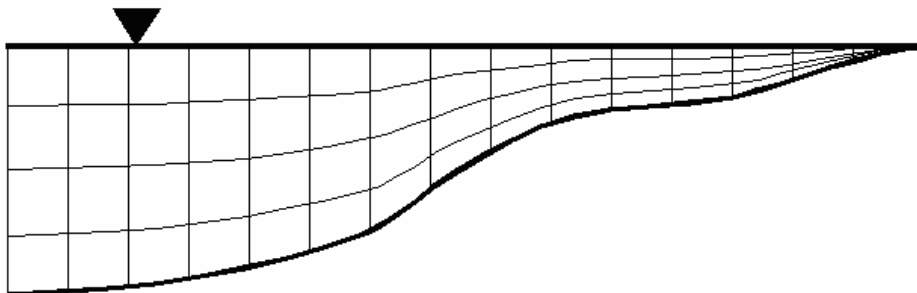


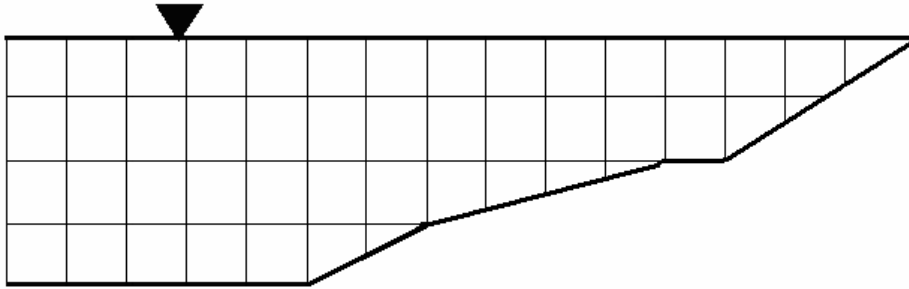
Figure 4-2: Finite volume element of MOHID model

### 4.3 Vertical Coordinates

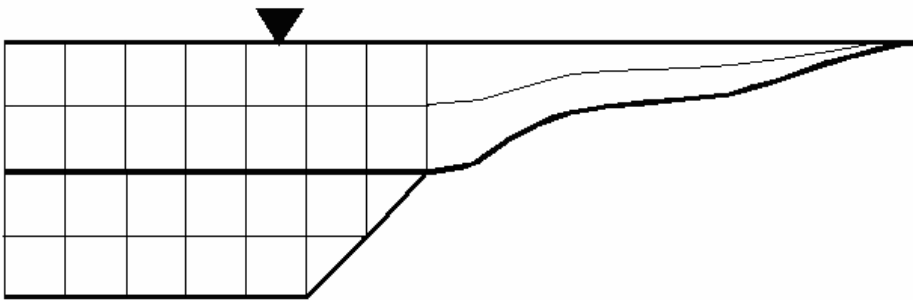
Actually the module Geometry can divide the water column in different vertical coordinates: Sigma, Cartesian, Lagrangian (based on Sigma or based on Cartesian), “Fixed Spacing” and Harmonic. A subdivision of the water column into different domains is also possible. The Sigma and the Cartesian coordinates are the classical ones. The Cartesian coordinate can be used with or without “shaved cells”. The Lagrangian coordinate moves the upper and lower faces with the vertical flow velocity. The “Fixed Spacing” coordinate allows the user to study flows close to the bottom and the Harmonic coordinate works like the Cartesian coordinate, just that the horizontal faces close to the surface expand and collapse depending on the variation of the surface elevation. This coordinate was implemented in the geometry module to simulate reservoirs.



**Figure 4-3: Sigma domain with 4 Layers**



**Figure 4-4: Cartesian domain with 4 Layers (shaved cells)**



**Figure 4-5: Sub-division of the water column in a Cartesian domain (inferior) and a Sigma domain (superior)**

## 5 The Hydrodynamic Module

### 5.1 Introduction

In this section the hydrodynamic module of the model MOHID is described. The information flux of the hydrodynamic module, relative to the other modules of *Mohid*, is shown in Figure 5-1.

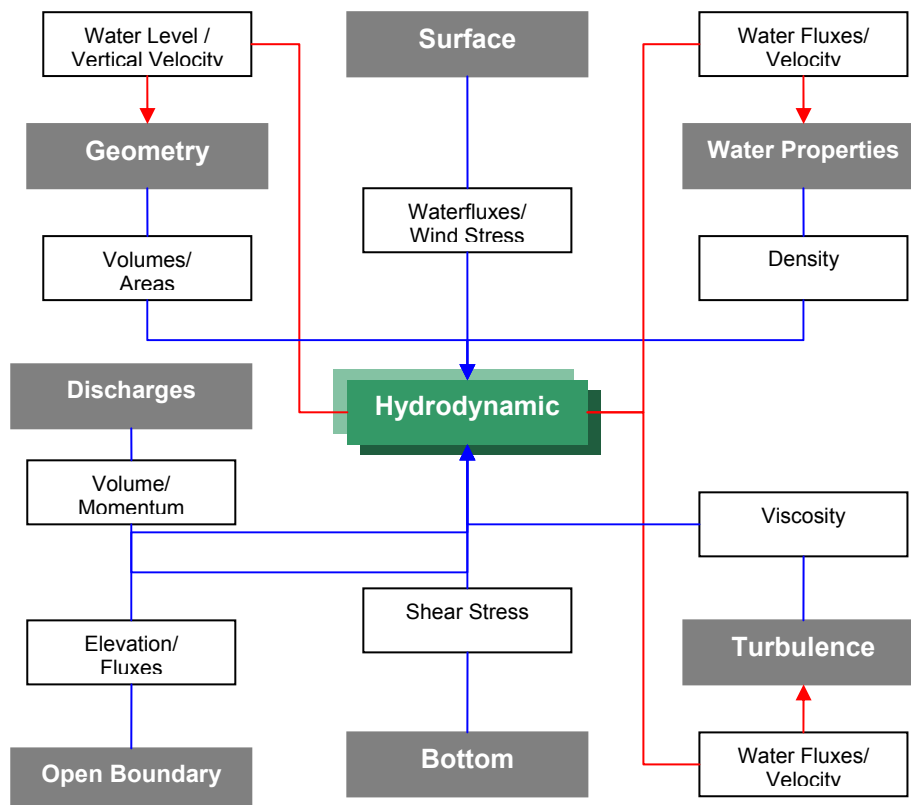


Figure 5-1: Information flux between the Hydrodynamic Module and other modules

### 5.2 Equations

The model solves the three-dimensional incompressible primitive equations. Hydrostatic equilibrium is assumed as well as Boussinesq and Reynolds approximations. All the equations below have been derived taken into account these approximations. The momentum balance equations for mean flow horizontal velocities are, in Cartesian form:

$$\begin{aligned} \partial_t u = & -\partial_x(uu) - \partial_y(uv) - \partial_z(uw) + fv - \frac{1}{\rho_0} \partial_x p \\ & + \partial_x((v_H + \nu) \partial_x u) + \partial_y((v_H + \nu) \partial_y u) + \partial_z((v_t + \nu) \partial_z u) \end{aligned} \quad \text{Eq. 5-1}$$

$$\begin{aligned} \partial_t v = & -\partial_x(vu) - \partial_y(vv) - \partial_z(vw) - fu - \frac{1}{\rho_0} \partial_y p \\ & + \partial_x((v_H + \nu) \partial_x v) + \partial_y((v_H + \nu) \partial_y v) + \partial_z((v_t + \nu) \partial_z v) \end{aligned} \quad \text{Eq. 5-2}$$

Where  $u$ ,  $v$  and  $w$  are the components of the velocity vector in the  $x$ ,  $y$  and  $z$  directions respectively,  $f$  the Coriolis parameter,  $v_H$  and  $v_t$  the turbulent viscosities in the horizontal and vertical directions,  $\nu$  is the molecular kinematic viscosity (equal to  $1.3 \cdot 10^{-6} \text{ m}^2 \text{ s}^{-1}$ ),  $p$  is the pressure. The temporal evolution of velocities (term on the left hand side) is the balance of advective transports (first three terms on the right hand side), Coriolis force (forth term), pressure gradient (next three terms) and turbulent diffusion (last three terms).

The vertical velocity is calculated from the incompressible continuity equation (mass balance equation):

$$\partial_x u + \partial_y v + \partial_z w = 0 \quad \text{Eq. 5-3}$$

by integrating between bottom and the depth  $z$  where  $w$  is to be calculated:

$$w(z) = \partial_x \int_{-h}^z u dx + \partial_y \int_{-h}^z v dy \quad \text{Eq. 5-4}$$

The free surface equation is obtained by integrating the equation of continuity over the whole water column (between the free surface elevation  $\eta(x,y)$  and the bottom  $-h$ ):

$$\partial_t \eta = -\partial_x \int_{-h}^{\eta} u dz - \partial_y \int_{-h}^{\eta} v dz \quad \text{Eq. 5-5}$$

The hydrostatic approximation is assumed with:

$$\partial_z p + g\rho = 0 \quad \text{Eq. 5-6}$$

where  $g$  is gravity and  $\rho$  is density. If the atmospheric pressure  $p_{atm}$  is subtracted from  $p$ , and density  $\rho$  is divided into a constant reference density  $\rho_0$  and a deviation  $\rho'$  from that constant reference density, after integrating from the free surface to the depth  $z$  where pressure is calculated, we arrive to:

$$p(z) = p_{atm} + g\rho_0(\eta - z) + g \int_z \eta \rho' dz \quad \text{Eq. 5-7}$$

Eq. 5-7 relates pressure at any depth with the atmospheric pressure at the sea surface, the sea level and the anomalous pressure integrated between that level and the surface. By using this expression and the Boussinesq approximation, the horizontal pressure gradient in the direction  $x_i$  can be divided in three contributions:

$$\partial_{x_i} p = \partial_{x_i} p_{atm} - g\rho_0 \partial_{x_i} \eta - g \int_z \eta \partial_{x_i} \rho' dz \quad \text{Eq. 5-8}$$

The total pressure gradient is the sum of the gradients of atmospheric pressure, of sea surface elevation (barotropic pressure gradient) and of the density distribution (baroclinic pressure gradient). This decomposition of the pressure gradient is substituted in Eq. 5-1 and Eq. 5-2.

The density is obtained from the salinity and from the temperature, which are transported by the water properties module.

### 5.3 Discretization

#### 5.3.1 Spatial discretization: Finite volume approach

The spatial discretization is described in the geometry module.

#### 5.3.2 Temporal discretization: semi-implicit ADI algorithm

The temporal discretization is carried out by means of a semi implicit ADI (Alternate Direction Implicit) algorithm, introduced by Peaceman and

Racford in 1955 (Fletcher, [1991]). This algorithm computes alternatively one component of horizontal velocity implicitly while the other is calculated explicitly. The resulting equation system is a tridiagonal one that can be solved by Thomas algorithm in an efficient and quick way. This allows preserving the stability advantages of implicit methods without the drawbacks of computational expensiveness and associated phase errors. A longer time-step can therefore be used. Two different discretizations are coded in the model: a 4 equations one with two time levels per iteration- the S21 scheme (Eq. 5-9) by Abbott et al. [1973]- and the 6 equation algorithm by Leendertsee, [1967], more convenient when intertidal zones are to be modeled, since velocities are updated every half time step. The S21 scheme is shown by Eq. 5-9:

$$\begin{aligned}
 & \eta^{t+1/2}(u^{t+1}, u^t, v^{t+1/2}, v^{t-1/2}) \rightarrow u^{t+1} \rightarrow \\
 & w^{*t+1/2} \xrightarrow{\text{GeometryUpdate}} w^{t+1/2} \rightarrow S^{t+1/2}, T^{t+1/2} \rightarrow \\
 & \eta^{t+1}(u^t, u^{t+1}, v^{t+3/2}, v^{t+1/2}) \rightarrow u^{t+3/2} \rightarrow \\
 & w^{*t+1} \xrightarrow{\text{GeometryUpdate}} w^{t+1} \rightarrow S^{T+1}, T^{T+1}
 \end{aligned}
 \tag{Eq. 5-9}$$

Each iteration is divided in two half steps. In the first half step, the free surface elevation  $\eta$  and then one of the horizontal velocities ( $u$ ) are computed in an implicit way. The required value of the other velocity is taken from the previous time step. A vertical velocity  $w^*$  is computed from the continuity equation. Then, geometry is updated and the vertical velocity is corrected. The same process is followed in the next half step, but for the other component of horizontal velocity. In this diagram, salinity and temperature are computed each half step. As internal modes are much slower than external modes,  $S$  and  $T$  can be updated with a longer time step without losing accuracy and stability.

### 5.3.3 Discretization of the different processes

A sketch of the discretization will be given below. A full description of the discretization may be found in Martins, [2000] and Montero, [1999].

### 5.3.3.1 Free surface equation

Free surface elevation is calculated by integrating the continuity equation (Eq. 5-3) over the whole water column. In the finite volume approach, this integration is done via a summation the volume fluxes over all water column cells. For the S21 discretization and the first half time step, it reads:

$$\frac{\eta_{ij}^{t+1/2} - \eta_{ij}^t}{\Delta t / 2} = \frac{1}{A_{hij}} \left[ \frac{1}{2} \left( \sum_{kbot}^{kmax} U_{ijk}^{t+1} A_{u_{ijk}}^t + \sum_{kbot}^{kmax} U_{ijk}^t A_{u_{ijk}}^t \right) - \frac{1}{2} \left( \sum_{kbot}^{kmax} U_{ij+1k}^{t+1} A_{u_{ij+1k}}^t + \sum_{kbot}^{kmax} U_{ij+1k}^t A_{u_{ij+1k}}^t \right) \right] + \frac{1}{A_{hij}} \left[ \frac{1}{2} \left( \sum_{kbot}^{kmax} V_{ijk}^{t+1/2} A_{v_{ijk}}^t + \sum_{kbot}^{kmax} V_{ijk}^{t-1/2} A_{v_{ijk}}^t \right) - \frac{1}{2} \left( \sum_{kbot}^{kmax} V_{i+1jk}^{t+1/2} A_{v_{i+1jk}}^t + \sum_{kbot}^{kmax} V_{i+1jk}^{t-1/2} A_{v_{i+1jk}}^t \right) \right] \quad \text{Eq. 5-10}$$

where  $A_{hij} = DUX_{ij} * DVY_{ij}$  is the area projected on the horizontal plane. Fluxes are temporally averaged, so the calculus is centered in  $t+1/2$ . An analogous discretization is carried out for the next half step. The fluxes  $U \cdot AU$  and  $V \cdot AV$  are obtained from the momentum equation. The discretization of the different terms will be discussed below.

### 5.3.3.2 Velocity equation

If we discretize equation (Eq. 5-1) making use of (**Error! Reference source not found.**) and S21 discretization, we arrive to (an equivalent equation is derived for  $v$ , Eq. 5-2) for every cell  $u_{ijk}$  of the grid:

$$\frac{\Omega_{u_{ijk}}^t (U_{ijk}^{t+1} - U_{ijk}^t)}{\Delta t} + \sum_{m=1}^{Nfaces} \vec{F}_m \cdot \vec{n}_m A_m = f_{u_{ijk}} \Omega_{u_{ijk}}^t \bar{V}_{ijk}^t \quad \text{Eq. 5-11}$$

where  $\Omega_{u_{ijk}}^t$  is the volume of the computation cell for  $U_{ijk}$  and  $f_{u_{ijk}}$  is the value of the Coriolis parameter for that cell. The value  $\bar{V}_{ijk}^t$  represents the average value of the  $v$ -component of the flow on this cell. The second term on the left hand side represents the fluxes of the forces  $F_m$  through the surface  $A_m$  of the cell  $m$ . The Coriolis force is the term on the right hand

side and the other terms in the equation are included in the summation on the left hand side.

### 5.3.3.2.1 Coriolis term

As we can see on the right hand side of Eq. 5-11, the Coriolis term is discretized explicitly, although it is well-known that this implies a restriction on  $\Delta t$  ( $\Delta t \leq 2/f$ , with  $f$  the Coriolis parameter). This limitation is not critical for coastal applications -for latitude of  $43^\circ$   $\Delta t \leq 2000$  s  $\approx$  5h 30min, that is much bigger than the time steps chosen in these applications.

The other terms in this formulation are seen as fluxes through the surfaces of the control volume, and therefore enter in the second term on the left hand side.

### 5.3.3.2.2 Advective terms

In order to guarantee momentum conservation, fluxes into the element must have null divergence. This is accomplished by using in the convective terms the same fluxes obtained in the last computation of elevation and vertical velocity. Convective fluxes are computed in every face of the cell:

$$\begin{aligned}
 - \sum_{m=1}^{N_{faces}} \vec{F}_m \cdot \vec{n} A_m = & \left[ (U \cdot ufluxU)_{ij+1k}^t - (U \cdot ufluxU)_{ijk}^t + \right. \\
 & (U \cdot ufluxV)_{i+1jk}^t - (U \cdot ufluxV)_{ijk}^t + \\
 & \left. (U \cdot ufluxW)_{ijk+1}^t - (U \cdot ufluxW)_{ijk}^t \right] \quad \text{Eq. 5-12}
 \end{aligned}$$

with  $ufluxU_i$  denotes the flux of  $U_i$  through the cell of calculus of  $u$ . A mixed scheme upwind-central differences is used for computing  $ufluxU_i$  (James, [1987], Santos [1995]). Horizontal advective fluxes are discretized explicitly as the restriction that surface waves impose on stability is small for the characteristic range of velocities. The vertical advective term can give problems if the layer thickness is small, as can happen in shallow zones with sigma grids. Two solutions to this problem have been introduced in the model: an implicit discretization or neglecting this term in those regions.

### 5.3.3.2.3 Barotropic pressure gradient

The restriction of surface waves on stability lead to the implementation of the semi-implicit algorithm so this term limits stability and consequently is discretized implicitly. For the cell  $u_{ijk}$  and the first semi-step:

$$-\sum_{m=1}^{N_{faces}} \vec{F}_m \cdot \vec{n} A_m = \frac{1}{\rho_0} \left[ (P_{atm_{j-1}}^{t+1/2} - P_{atm_j}^{t+1/2}) + \rho_0 g (\eta_{ij-1}^{t+1/2} - \eta_{ij}^{t+1/2}) \right] A_{u_{ijk}}^t \quad \text{Eq. 5-13}$$

This expression, when substituted in the equation for the free surface, results in a tridiagonal system, which is solved by Gaussian elimination. In the equation for velocities, the values of  $\eta$  are already known, which allows the explicit discretization of this term for introduction in momentum equations.

### 5.3.3.2.4 Baroclinic pressure gradient

Internal modes do not introduce a stringent restriction on stability, so they can be discretized explicitly. The fluxes induced by this term through the faces of a  $u_{ijk}$  cell are:

$$-\sum_{m=1}^{N_{faces}} \vec{F}_m \cdot \vec{n} S_m = \frac{g}{\rho_0} \left( \sum_{l=k+1}^{k_{max}} (\rho'_{ij-1l} DWZ_{ij-1l}) + \rho'_{ij-1k} \Delta z'_{ij-1k} \right) A_{u_{ijk}}^t - \left( \sum_{l=k+1}^{k_{max}} (\rho'_{ijl} DWZ_{ijl}) + \rho'_{ijk} \Delta z'_{ijk} \right) A_{u_{ijk}}^t \quad \text{Eq. 5-14}$$

where  $\Delta z'_{ijk}$  represents the vertical distance from the cell top to the velocity point and arises as a consequence of the vertical staggering of the grid ( $\rho'$  is not defined in the same point as the u-velocity).

### 5.3.3.2.5 Horizontal diffusive fluxes

Horizontal diffusive fluxes are computed in every vertical face of the cell, applying that fluxes are normal to these faces:

$$-\sum_{m=1}^{N_{faces}} \vec{F}_m \cdot \vec{n} A_m = (F_{ij-1/2k}^t Azx'_{ij-1k} - F_{ij+1/2k}^t Azx'_{ijk}) +$$

$$\left( F_{i-1/2,jk}^t \frac{Av_{ijk}^t + Av_{ij-1k}^t}{2} - F_{i+1/2,jk}^t \frac{Av_{i+1,jk}^t + Av_{i+1,j-1k}^t}{2} \right) \quad \text{Eq. 5-15}$$

Fluxes for x direction are:

$$F_{ij-1/2k}^t = v_{H_{ij-1k}}^t \frac{U_{ijk}^t - U_{ij-1k}^t}{DUX_{ij-1}} \quad \text{Eq. 5-16}$$

and for the y-direction:

$$F_{i-1/2,jk}^t = v_{H_{i-1/2,j-1/2k}}^t \frac{U_{ijk}^t - U_{ij-1k}^t}{(DYY_{ij} + DYY_{i-1,j})/2} \quad \text{Eq. 5-17}$$

where the horizontal viscosity coefficient  $v_H^t$  is interpolated to the appropriate point.

#### 5.3.3.2.6 Vertical diffusion

These terms must be discretized implicitly as the restriction imposed by an explicit discretization on the time step is strict for the resolution we will use.

$$- \sum_{m=1}^{N_{faces}} \vec{F}_m \cdot \vec{n} A_m = (F_{ijk-1/2}^{t+1} - F_{ijk+1/2}^{t+1}) A_{h_{ij-1/2}}^t + \quad \text{Eq. 5-18}$$

with fluxes given by the equation:

$$F_{ijk-1/2}^{t+1} = v_{ij-1/2k-1}^t \frac{U_{ijk}^{t+1} - U_{ijk-1}^{t+1}}{DUZ_{ijk-1}^t} \quad \text{Eq. 5-19}$$

## 5.4 Boundary conditions

### 5.4.1 Free surface

All advective fluxes across the surface are assumed to be null. This condition is imposed by assuming that the vertical flux of  $W$  at the surface is null:

$$Wflux|_{surface} = 0 \quad \text{Eq. 5-20}$$

Diffusive flux of momentum is imposed explicitly by means of a wind surface stress,  $\tau_w$ :

$$v \frac{\partial \overline{v_H}}{\partial z} \Big|_{surface} = \vec{\tau}_w \quad \text{Eq. 5-21}$$

Wind stress is calculated according to a quadratic friction law:

$$\vec{\tau}_w = C_D \rho_a \vec{W} \left| \vec{W} \right| \quad \text{Eq. 5-22}$$

where  $C_D$  is a drag coefficient that is function of the wind speed,  $\rho_a$  is air density and  $W$  is the wind speed at a height of 10 m over the sea surface.

#### 5.4.2 Bottom boundary

Also at the bottom, advective fluxes are imposed as null and diffusive flux of momentum is estimated by means of a bottom stress that is calculated by a non-slip method with a quadratic law that depends on the near-bottom velocity. So, the diffusive term at the bottom is written as:

$$v \frac{\partial \overline{v_H}}{\partial z} \Big|_{bottom} = C_D \vec{v}_H \left| \vec{v}_H \right| \quad \text{Eq. 5-23}$$

$C_D$  is the bottom drag coefficient that is calculated with the expression:

$$C_D = \left( \frac{\kappa}{\log \left( \frac{z + z_0^b}{z_0^b} \right)} \right)^2 \quad \text{Eq. 5-24}$$

where  $\kappa$  is von Karman constant and  $z_0^b$  is the bottom roughness length. This quadratic law is derived from the logarithmic law of the wall near boundaries characteristic of boundary layers, as the bottom velocities are located half a grid box above the bottom. This term is calculated semi-implicitly following Backhaus [1985] for the sake of numerical stability.

No fluxes of salinity and temperature are considered at the bottom.

#### 5.4.3 Lateral closed boundaries

At these boundaries, the domain is limited by land. For the resolution we are using, this lateral boundary layer is resolved, so a impermeable, free slip condition can be used:

$$\frac{\partial \vec{v}_H}{\partial \eta} = 0 \quad \text{Eq. 5-25}$$

$$\vec{v} \cdot \vec{n} = 0 \quad \text{Eq. 5-26}$$

In the finite volume formalism, these conditions are implemented straightforwardly by specifying zero normal water fluxes and zero momentum diffusive fluxes at the cell faces in contact with land.

#### 5.4.4 Open boundaries

Open boundaries arise from the necessity of confining the domain to the region of study. The values of the variables must be introduced there such that it is guaranteed that information about what is happening outside the domain will enter the domain in a way that the solution inside the domain is not corrupted. Also, waves generated inside the domain should be allowed to go out. There exists no perfect open boundary condition and the most suitable would depend on the domain and the phenomena to be modeled. A recent review paper comparing open boundary conditions in test cases can be found in Palma and Matano [1999]. Some different open boundaries are already introduced in MOHID 3D (Santos, [1995], Montero, [1999]) and some others like FRS (Flow Relaxation Scheme) are in progress.

#### 5.4.5 Moving boundaries

Moving boundaries are closed boundaries that change position in time. If there are inter tidal zones in the domain, some points can be alternatively covered or uncovered depending on tidal elevation. A stable algorithm is required for modeling these zones and their effect on hydrodynamics of

estuaries. A detailed exposition of the algorithms used in MOHID can be found in Martins et al. [1999] and Martins [1999].

## 6 The Lagrangian Module

### 6.1 Introduction

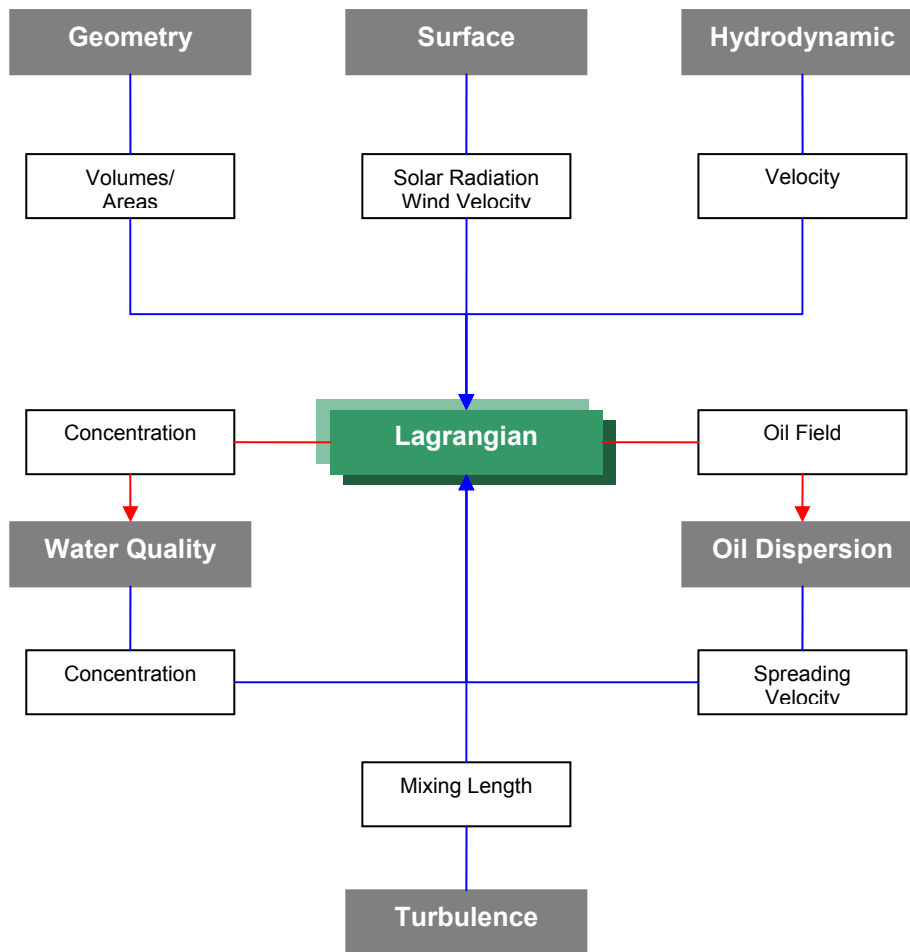
Lagrangian transport models are very useful to simulate localized processes with sharp gradients (submarine outfalls, sediment erosion due to dredging works, hydrodynamic calibration, oil dispersion, etc.).

*Mohid's* Lagrangian module uses the concept of tracer. The most important property of a tracer is its position (x,y,z). For a physicist a tracer can be a water mass, for a geologist it can be a sediment particle or a group of sediment particles and for a chemist it can be a molecule or a group of molecules. A biologist can spot phytoplankton cells in a tracer (at the bottom of the food chain) as well as a shark (at the top of the food chain), which means that a model of this kind can simulate a wide spectrum of processes.

The movement of the tracers can be influenced by the velocity field from the hydrodynamic module, by the wind from the surface module, by the spreading velocity from oil dispersion module and by random velocity.

At the present stage the model is able to simulate oil dispersion, water quality evolution and sediment transport. To simulate oil dispersion the lagrangian module interacts with the oil dispersion module, to simulate the water quality evolution the lagrangian module uses the feature of the water quality module. Sediment transport can be associated directly to the tracers using the concept of settling velocity.

Figure 6-1 represents the information flux between the lagrangian module and other modules of *Mohid*.



**Figure 6-1: Information flux between the Lagrangian module and other modules**

Another feature of the lagrangian transport model is the ability to calculate residence times. This can be very useful when studying the exchange of water masses in bays or estuaries.

## 6.2 Tracer concept

Like referred above, the *Mohid's* Lagrangian module uses the concept of tracer. The tracers are characterized by their spatial coordinates, volume and a list of properties (each with a given concentration). The properties can be the same one like the ones described in the water properties module or coliform bacteria. Each tracer has associated a time to perform the random movement.

The tracers are “born” at origins. Tracers which belong to the same origin have the same list of properties and use the same parameters for random walk, coliform decay, etc. Origins can differ in the way they emit tracers. There are three different ways to define origins in space:

- a “Point Origins” emits tracers at a given point;
- a “Box Origins” emits tracers over a given area;
- a “Accident Origins” emit tracers in a circular form around a point;

There are two different ways in which origins can emit tracers in time:

- a “Continuous Origins” emits tracers during a period of time;
- a “Instantaneous Origins” emits tracers at one instant;

Origins can be grouped together in Groups. Origins which belong to the same group are grouped together in the output file, so it is more easy to analyze the results.

## 6.3 Equations

### 6.3.1 Tracer Movement

The major factor responsible for particle movement is generally the mean velocity. The spatial co-ordinates are given by the definition of velocity:

$$\frac{dx_i}{dt} = u_i(x_i, t) \quad \text{Eq. 6-1}$$

where  $u$  stands for the mean velocity and  $x$  for the particle position.

Equation 3.1 is solved using a simple explicit method:

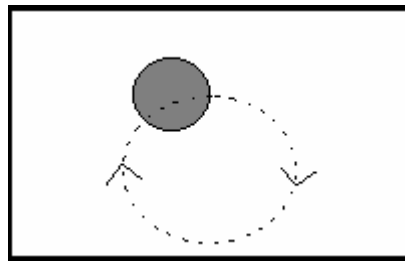
$$x_i^{t+\Delta t} = x_i^t + \Delta t \cdot u_i^t \quad \text{Eq. 6-2}$$

Higher order accuracy requires the use of an iterative procedure. The scheme adopted by Monteiro (1995) uses second order accuracy. Costa (1991) concluded that higher order schemes are important whenever curvature of the flow exists and a large time step is used. For most of the

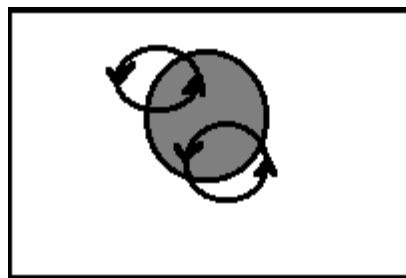
natural flows the explicit method is accurate enough. Velocity at any point of space is calculated using a linear interpolation between the points of the hydrodynamic model grid. The lagrangian module permits to divide the calculation of the trajectory of the tracers into sub-steps of the hydrodynamic time step.

### 6.3.2 Turbulent Diffusion

Turbulent transport is responsible for dispersion. The effect of eddies over particles depends on the ratio between eddies and particle size. Eddies bigger than the particles make them move at random as explained in Figure 6-2. Eddies smaller than the particles cause entrainment of matter into the particle, increasing its volume and its mass according to the environment concentration, like shown in Figure 6-3.



**Figure 6-2: Random movement forced by an eddy larger than the particle**



**Figure 6-3: Random movement forced by an eddy larger than the particle**

The random movement is calculated following the procedure of Allen (1982). The random displacement is calculated using the mixing length and the standard deviation of the turbulent velocity component, as given by the turbulence closure of the hydrodynamic model. Particles retain that velocity during the necessary time to perform the random movement, which is dependent on the local turbulent mixing length.

The increase in volume is associated with small-scale turbulence and is reasonable to assume it as isotropic. In these conditions, small particles keep their initial form and their increase in volume is a function of the volume itself.

### 6.3.3 Mass Decay rate

The decay rate of coliform bacteria, which are can associated to the tracers, is computed by the following equation:

$$\frac{dC}{dt} = -\frac{\ln 10}{T_{90}} C \quad \text{Eq. 6-3}$$

where C represents the concentration, and  $T_{90}$  the time interval for 90% of the coliform bacteria to die.

An implicit method is used to solve Eq. 6-3 numerically, preventing a negative number of coliform bacteria.

### 6.3.4 Monitoring Boxes

The lagrangian module permits to monitor the distribution of particles inside “monitoring boxes”. This feature is very useful to compute the residence time of water inside these monitoring boxes and the origins of the water present inside each box at each moment. The lagrangian module “monitors” the boxes the following way:

- In every instant the volume of each box  $b$ ,  $InstBoxVol(b)$  is calculated;

$$InstBoxVol(b) = \int (h + Z) dx dy$$

- In every instant the origin “o” of the water inside each monitoring box “b” is identified and the volume of the water from each origin is stored in the variable  $InstVolumeByOrigin(b, o)$ :

$$InstVolumeByOrigin(b, o) = \sum_o Vol_j^b$$

- **In case of instantaneous emissions in boxes**, these contributions are integrated over the time, given the integrated contribution over the time,  $IntgVolumeByOrigin(b, o)$

$$IntgVolumeByOrigin(b, o) = \int InstVolumeByOrigin(b, o) dt$$

A measure of the residence time of the water emitted into box “o” in monitoring box “b” is given by:

$$ResidenceTimePerBox(b, o) = IntgVolumeByOrigin(b, o) / InitialVol(o)$$

Adding the values for all monitoring boxes inside the estuary one gets the residence time inside the whole system of the water emitted into box “o”:

$$ResidenceTime(o) = \sum_b ResidenceTimePerBox(b, o)$$

These values also permit to compute how each monitoring box is influenced by each emitting box:

$$InfluenceOverBox(b, o) = IntgVolumeByOrigin(b, o) / InitialVol(b)$$

In case of a continuous emission, the residence time can be computed as:

$$ResidenceTimePerBox(b, o) = InstVolumeByOrigin(b, o) / DischargeRate(o)$$

Again, the addition of the values of the residence time in each box gives the Residence time inside the System

- The Output is done in four ways:
  - Time Series in ASCII columns for every monitoring box and every time step. For every monitoring box a file is written where the first column represents the box volume and others represent the contributions to this box from every origin. Both, the instantaneous and the integrated values are written
  - Time Series in ASCII of the variable

*ResidenceTimeperBox(b,o)*

- HDF Matrix for every Origin, every output instant. The relative contribution of each emitting box “o” for the instantaneous volume in each monitoring box “b” is written as the percentage of the instantaneous volume of the monitoring box:

$$\text{Matrix}(b,o) = 100 * \text{InstVolumeByOrigin}(b,o) / \text{InstBoxVol}(b)$$

HDF Matrix, one for all origins, every output time. The output Matrix is filled depending on the instantaneous contributions (in volume) of particle to a given monitor box. The missing volume is filled with 0 (freshwater).

## 7 The Oil Module

### 7.1 Introduction

The prediction and simulation of the trajectory and weathering of oil spills are essential to the development of pollution response and contingency plans, as well as to the evaluation of environmental impact assessments.

In order to predict the behaviour of the oil products spilled in coastal zones, an oil weathering model was developed, which predicts the evolution and behaviour of the processes (transport, spreading and behaviour) and properties of the oil product spilled in water. Some pollution response methods are also integrated in the model.

### 7.2 Implementation

Oil density and viscosity, and many different processes are included in oil module, such as oil spreading, evaporation, dispersion, sedimentation, dissolution, emulsification, oil beaching and removal techniques.

Different alternative methods were coded for the prediction of some processes like oil spreading, evaporation, dispersion, sedimentation and emulsification. Therefore, when using the model, there is more than one way of simulating the same process, depending, for example, on the characteristics of the computational mesh or on the magnitude of the spill.

The oil weathering module (OWM) uses mainly the 3D hydrodynamics and 3D lagrangian transport modules. The hydrodynamic module simulates the velocity field necessary for the lagrangian module to calculate oil trajectories. These oil trajectories are computed assuming that oil can be idealized as a large number of particles that independently move in water. Water properties and atmospheric conditions are introduced in lagrangian module and used by oil module for determination of oil processes and properties. Excepting spreading and oil-beaching, all weathering processes and properties are assumed uniform for all tracers, like water properties and atmospheric conditions, which are considered equal to these environmental conditions determined in accident origin.

As it was already mentioned, the movement of the oil tracers can be influenced by the velocity field from the hydrodynamic module, by the wind from the surface module, by the spreading velocity from oil module and by random velocity.

Oil temperature is assumed equal to water temperature, neglecting solar radiation or any other energy transfer process that may influence oil temperature.

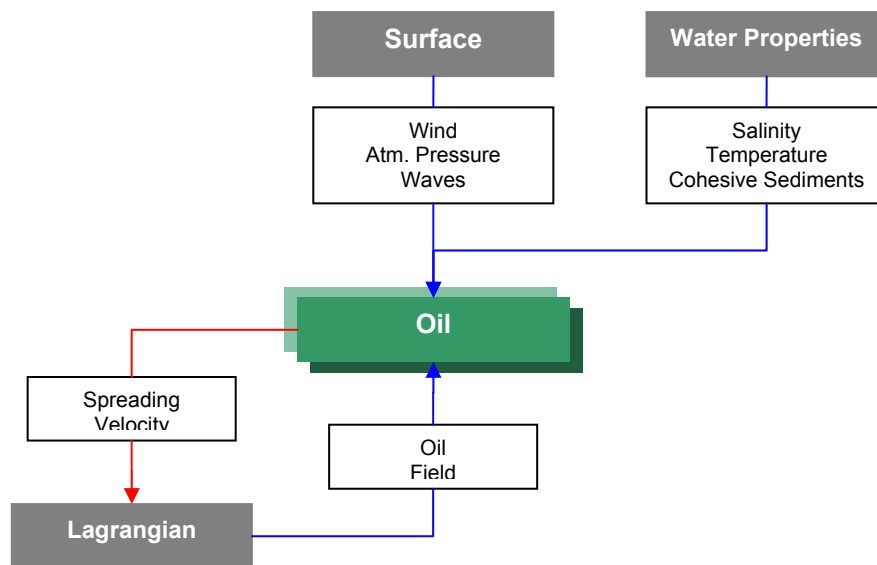


Figure 7-1 Information flux between the oil module and other modules

### 7.3 Equations

#### 7.3.1 Spreading

In case of an instant accident, the initial area of spilled oil is determined by an equation deduced from Fay’s solutions (Fay, 1969). Once initial phase of spreading (gravity-inertial phase) is too short, initial area is calculated when that phase ends, and gravity-viscous phase starts:

$$A_0 = \pi \frac{k_2^4}{k_1^2} \left( \frac{\Delta g V_0^5}{v_w} \right)^{1/6} \tag{Eq. 7-1}$$

Where:

$A_0$  – initial area

$$\Delta = (\rho_w - \rho_o) / \rho_w$$

$\rho_w$  – water density

$\rho_o$  – oil density

$g$  – gravity acceleration

$V_0$  – volume of spilled oil

$\nu_w$  – water kinematic viscosity

$k_1 = 0.57$  and  $k_2 = 0.725$  (as recommended by Flores et al 1999)

Two different algorithms were implemented to estimate oil spreading.

One of the algorithms determines random velocities  $u_d$  e  $v_d$  (with uniform distribution) inside range  $[-U_r, U_r]$ ,  $[-V_r, V_r]$  (in directions  $x$  and  $y$ , respectively) proportional to diffusion coefficients, which are calculated assuming that lagrangian tracers spreading is equivalent to Fay's formulas solution (Fay, 1969). The following relationship between diffusion coefficients  $D_x$  and  $D_y$  and the velocity fluctuation range  $[-U_r, U_r]$ ,  $[-V_r, V_r]$  is adopted according to Leitão (1996):

$$U_r = \sqrt{\frac{2D_x}{\Delta t}} \quad \text{Eq. 7-2}$$

$$V_r = \sqrt{\frac{2D_y}{\Delta t}} \quad \text{Eq. 7-3}$$

Random velocities are therefore determined in the following way, like suggested by Proctor *et al.* (1994):

$$u_d = R_1 \cos(2\pi R_2) \cdot U_r \quad \text{Eq. 7-4}$$

$$v_d = R_1 \sin(2\pi R_2) \cdot V_r \quad \text{Eq. 7-5}$$

where  $R_1$  e  $R_2$  are randomly generated numbers between 0 and 1.

The only phase simulated in spreading is the gravity-viscous phase, from solutions proposed by Fay, where diffusion coefficients  $D_x$  and  $D_y$  have the following formulation (this model uses a numerical solution of this equation):

$$D_x = D_y = \frac{\pi k_2^2}{16} \left( \frac{\Delta g V^2}{\nu_w^{1/2}} \right)^{1/3} \cdot \frac{1}{\sqrt{t}} \quad \text{Eq. 7-6}$$

Where:

$V$ - volume of spilled oil

$t$  – time after spill

The other algorithm proposed for oil spreading is based in thickness differences inside oil slick, presuming that the existence of a thickness gradient generates a “spreading force” in the direction of minor thickness. Therefore, a tracer will move from the computational cell with larger oil thickness to the one thinner.

This formulation uses a coefficient to approach the solution to the Fay solution, in order to make results sensible to some factors, like different oil densities, originating different behaviours.

Spreading coefficient is given by:

$$k = k_1 \cdot \frac{\Delta^{1.6} g V^2}{\nu_w^{1/6}} \quad \text{Eq. 7-7}$$

where  $k_1$  is a parameter introduced by the user, with a default value of 10.0.

Therefore, in oil module velocities are calculated in the faces of cells where oil is present, in directions  $x$  and  $y$ , in the following way:

$$u_{cell} = k \cdot \frac{\Delta h}{\Delta x} \quad \text{Eq. 7-8}$$

$$v_{cell} = k \cdot \frac{\Delta h}{\Delta y} \quad \text{Eq. 7-9}$$

where  $\frac{\Delta h}{\Delta x}$  and  $\frac{\Delta h}{\Delta y}$  are the thickness gradients of a cell, in directions  $x$  and  $y$ . Subsequently, in lagrangian module tracers velocities are interpolated based on cell faces velocities and tracers position.

If average oil thickness becomes too thin – less than a value between 0.1 and 0.01 mm, depending of product viscosity –, oil spreading is stopped, according to Reed (1989).

### 7.3.2 Density

Density can be estimated by:

$$\rho_e = F_{wv} \cdot \rho_w + \rho_{oil}(1 - F_{wv})(1 + c_{DE}F_e)[1 - c_{DT}(T - T_0)] \quad \text{Eq. 7-10}$$

where  $\rho_e$  is the density of the emulsion at temperature  $T$ ,  $\rho_{oil}$  is the density of fresh oil at reference temperature  $T_0$ ,  $\rho_w$  is the water density,  $c_{DE}$  e  $c_{DT}$  are empirical constants (NOAA (1994) recommends the following values:  $c_{DE} = 0.18$  and  $c_{DT} = 8 \times 10^{-4}$ ).

The oil initial density is obtained from API density. Only oil products with lower density than water are modelled, because the remainder will sink.

### 7.3.3 Viscosity

Viscosity is changed by three major processes: temperature, evaporation and emulsification.

The influence of temperature can be calculated by Andrade's correlation:

$$\mu = \mu_0 e^{c_T \left( \frac{1}{T} - \frac{1}{T_0} \right)} \quad \text{Eq. 7-11}$$

where  $\mu$  is the oil viscosity at temperature  $T$ ,  $\mu_0$  is the initial oil viscosity at reference temperature  $T_0$  and  $c_T$  is an empirical constant whose recommended value by NOAA (1994) is 5000 K.

Viscosity modification due to emulsification is defined by Mooney's equation (1951):

$$\mu = \mu_0 e^{\left[ \frac{c_V F_{wv}}{(1-c_M F_{wv})} \right]} \quad \text{Eq. 7-12}$$

where  $F_{wv}$  is water volume fraction the emulsion,  $c_V$  is an adimensional empirical constant (Mackay et al., 1980 recommends the value of 2.5) and  $c_M$  is an additional Mooney's constant with the value of 0.65.

The effect of evaporation on viscosity is calculated by the following equation (Mackay et al., 1980):

$$\mu = \mu_0 \cdot e^{(c_E F_{em})} \quad \text{Eq. 7-13}$$

$F_{em}$  is the mass fraction of evaporated oil, and the adimensional empirical constant  $c_E$  varies with oil type, between 1 and 10, with higher values for more viscous products. In this model, when fresh oils at 15°C have a cinematic viscosity greater than 38 cSt,  $c_E$  is always considered 10. In case of less viscous oils,  $c_E$  is estimated by a second degree polynomial regression:

$$c_E = -0.0059 \cdot V_{cin15}^2 + 0.4461 \cdot V_{cin15} + 1.413 \quad \text{Eq. 7-14}$$

where  $V_{cin15}$  is the oil cinematic viscosity at 15°C.

The three previous equations (Eq. 7-11, Eq. 7-12 and Eq. 7-13) can be joined in a single equation:

$$\mu = \mu_0 \cdot e^{\left[ (c_E F_{em}) + \frac{c_V F_{wv}}{(1-c_M F_{wv})} + c_T \left( \frac{1}{T} - \frac{1}{T_0} \right) \right]} \quad \text{Eq. 7-15}$$

### 7.3.4 Evaporation

In MOHID, the oil evaporation process can be estimated by two different methods: an analytical method, also known as the evaporative exposure method (developed by Stiver & Mackay, 1984), and by a more recent methodology proposed by Fingas (1998).

Evaporative exposure method is given by the formula:

$$\frac{dF_e}{dt} = \frac{K_e A_s}{V_0} \cdot \exp\left(A - \frac{B}{T} (T_0 + T_G F_e)\right) \quad \text{Eq. 7-16}$$

$F_e$  is the volume fraction of evaporated oil,  $T$  is oil temperature,  $A_s$  is the oil slick area,  $V_0$  is the initial oil volume,  $K_e$  is the mass transfer coefficient, determined by a simple formulation proposed by Buchanan & Hurford (1988):

$$K_e = 2.5 \times 10^{-3} W^{0.78} \quad \text{Eq. 7-17}$$

$A$  and  $B$  are empirical constants,  $T_0$  is the initial boiling point and  $T_G$  is the distillation curve gradient. All these parameters depend of oil type. In this model, they are estimated, and  $T_0$  e  $T_G$  are obtained from  $API$  density, according to version 1.1 of ADIOS model (NOAA, 1994):

$$A = 6.3; B = 10.3$$

For crude oils:

$$T_0 = 532.98 - 3.1295 \cdot API \quad \text{Eq. 7-18}$$

$$T_G = 985.62 - 13.597 \cdot API \quad \text{Eq. 7-19}$$

For refined products:

$$T_0 = 654.45 - 4.6588 \cdot API \quad \text{Eq. 7-20}$$

$$T_G = 388.19 - 3.8725 \cdot API \quad \text{Eq. 7-21}$$

Mervin Fingas proposed other method for evaporation calculus. He proposed a simplified formulation, where the relevant factors are time and temperature.

For many oil types, Fingas determined specific empirical equations in the following forms (this model uses the numerical solutions of the following equations):

$$\%Ev = (\alpha + \beta \cdot T) \ln(t) \quad \text{Eq. 7-22}$$

or

$$\%Ev = (\alpha + \beta \cdot T) \sqrt{t} \quad \text{Eq. 7-23}$$

where  $\%Ev$  is the percentage (by weight) of evaporated oil,  $\alpha$  and  $\beta$  are empirical constants specific for each oil type,  $T$  is oil temperature,  $t$  is time after spill (minutes).

If empirical data is unknown, generically equations can be used:

$$\%Ev = [(0.165(\%D) + 0.045 \cdot (T - 15))] \ln(t) \quad \text{Eq. 7-24}$$

or

$$\%Ev = [(0.0254(\%D) + 0.01 \cdot (T - 15))] \sqrt{t} \quad \text{Eq. 7-25}$$

$\%D$  is the percentage (by weight) distilled at 180°C.

Square root equations are used in some refined oils and in short term simulations (1-2 days).

### 7.3.5 Emulsification

This process consists in incorporation of water in oil. This process usually starts after an amount of oil has evaporated. An emulsification constant is used, which means the percentage of oil evaporated before emulsification starts. By default, this constant is 0%.

When emulsification starts, incorporation of water in oil can be simulated by two different processes.

An equation widely used, proposed by Mackay *et al.* (1980), is implemented in this model:

$$\frac{dF_{wv}}{dt} = K_w (1+W)^2 \left( 1 - \frac{F_{wv}}{F_{wv}^{final}} \right) \quad \text{Eq. 7-26}$$

where  $F_{wv}$  is the water volume fraction incorporated in emulsion;  $F_{wv}^{final}$  is the final volume fraction of water incorporated in emulsion;  $K_w$  is an empirical constant, introduced by the model user. Usually this constant assume values between  $1.0 \times 10^{-6}$  and  $2.0 \times 10^{-6}$ . MOHID default value is  $1.6 \times 10^{-6}$ , which is also used in ADIOS model (NOAA, 1994).

The other algorithm used is Rasmussen equation (Rasmussen, 1985).

$$\frac{dF_{wv}}{dt} = R_1 - R_2 \quad \text{Eq. 7-27}$$

where:

$R_1$  - water incoming rate ( $s^{-1}$ ), given by:

$$R_1 = \frac{K_1}{\mu_0} (1+W)^2 (F_{wv}^{final} - F_{wv}) \quad \text{Eq. 7-28}$$

$R_2$  - water outgoing rate ( $s^{-1}$ ), given by:

$$R_2 = \frac{K_2}{Asph \cdot Wax \cdot \mu_0} F_{wv} \quad \text{Eq. 7-29}$$

$Asph$  is the asphaltene content in oil (%),  $Wax$  is the wax content (%), and  $K_1$  e  $K_2$  are experimentally determined constants by Rasmussen (1985):

$$K_1 = 5 \times 10^{-7} \text{ kg.m}^{-3}; K_2 = 1.2 \times 10^{-7} \text{ kg.m}^{-1} \cdot \text{s}^{-2}.$$

### 7.3.6 Dispersion

This is the process where oil droplets entrain in water column.

Two different methods are available to predict this weathering process. One of them is Delvigne & Sweeney (1988) method:

$$\frac{dm_d}{dt} = c_{oil} \cdot D_{ba}^{0.57} \cdot f_s \cdot F_{wc} \cdot d_0^{0.7} \cdot \Delta d \quad \text{Eq. 7-30}$$

This equation estimates mass transfer rate per time unit, where  $f_s$  is the surface fraction covered by oil (considered equal to oil content in emulsion water + oil);  $d_0$  is the droplet diameter;  $\Delta d$  is the oil droplets diameters range around  $d_0$  (model assumes a droplet size range between 5 – 70 microns. Bigger droplets will tend to resurface - NOAA, 1994);  $c_{oil}$  is a parameter experimentally determined which depends on oil type. This model uses a logarithmical regression based on oil cinematic viscosity:

$$c_{oil} = -312.25 \cdot \ln(V_{cin}) + 2509.8 \quad \text{Eq. 7-31}$$

where  $V_{cin}$  is the oil cinematic viscosity

(if this regression gives negative values,  $c_{oil}$  is considered 0)

$D_{ba}$  is the wave dissipation energy per unit of surface area, which can be calculated by:

$$D_{ba} = 0.0034 \rho_w g H_{rms}^2 \quad \text{Eq. 7-32}$$

$H_{rms}$  is:

$$H_{rms} = \frac{1}{\sqrt{2}} H_0 \quad \text{Eq. 7-33}$$

where  $H_0$  is wave height.

$F_{wc}$  is the fraction of the sea surface that is covered with whitecaps per time unit, given by:

$$F_{wc} = \frac{C_b(W - W_i)}{T_w} \quad \text{Eq. 7-34}$$

where  $C_b = 0.032 \text{ s.m}^{-1}$ ,  $W_i$  is the wind velocity to start whitecaps ( $4 \text{ m.s}^{-1}$ );  $T_w$  is the wave period.

If wave period and wave height are unknown, these properties can be

empirically determined as function of wind speed, according to ADIOS model formulations (NOAA, 1994):

$$H_0 = 0.243 \frac{W^2}{g} \quad \text{Eq. 7-35}$$

and

$$T_w = 8.13 \frac{W}{g} \quad \text{Eq. 7-36}$$

Once turbulent energy is difficult to determine, other simplified algorithms have been developed for vertical dispersion in function of square wind velocity. One of them is used in this model – the formulation proposed by Mackay *et al.* (1980):

$$\frac{dm_d}{dt} = 0.11 m_{oil} \frac{(1+W)^2}{1+50\mu^{1/2}h\sigma} \quad (\text{kg}\cdot\text{h}^{-1}) \quad \text{Eq. 7-37}$$

, where  $m_{oil}$  is the oil mass that remains in surface,  $\mu$  is the oil dynamic viscosity (cP),  $h$  is the slick thickness (cm),  $W$  is the wind velocity ( $\text{m}\cdot\text{s}^{-1}$ ) and  $\sigma$  is oil-water interfacial tension ( $\text{dyne}\cdot\text{cm}^{-1}$ ).

### 7.3.7 Sedimentation

Although oil sedimentation process is relatively complicated and difficult to estimate, a formulation developed by Science Applications International (Payne *et al.*, 1987) is used in MOHID:

$$\frac{dm_{sed}}{dt} = 1.3 \sqrt{\frac{E}{V_w}} K_a \cdot C_{oil} \cdot C_{sed} \cdot z_i \cdot A_s \quad \text{Eq. 7-38}$$

This equation gives the mass of sedimented oil per time unit ( $\text{kg}\cdot\text{s}^{-1}$ ), where:

$V_w$  is the water dynamic viscosity ( $\text{kg}\cdot\text{m}^{-1}\cdot\text{s}^{-1}$ );  $K_a$  is the stick parameter with value  $1 \times 10^{-4} \text{ m}^3\cdot\text{kg}^{-1}$ ;  $z_i$  is the intrusion depth of oil droplets in the water column due to breaking waves, given by Delvigne & Sweeney (1988):

$$z_i = 1.5 \cdot H_0 \quad \text{Eq. 7-39}$$

$E$  is the rate of dissipated energy from water surface ( $\text{J.m}^{-3}.\text{s}^{-1}$ ). This is estimated from the wave dissipation energy ( $D_{ba}$ ), previously explained in dispersion section:

$$E = \frac{D_{ba}}{z_i \cdot T_w} \quad \text{Eq. 7-40}$$

$C_{sed}$  is the sediment concentration in water column ( $\text{kg.m}^{-3}$ ),  $C_{oil}$  is the oil droplet concentration in water column ( $\text{kg.m}^{-3}$ ). This concentration can be determined from dispersion rate proposed by Delvigne & Sweeney (1988) (explained in dispersion section), integrating this rate for wave period and intrusion depth of oil droplets:

$$\frac{dC_{oil}}{dt} = \frac{dm_d}{dt} \cdot \frac{1}{z_i} \quad \text{Eq. 7-41}$$

Only droplets greater than 70 microns and smaller than 200 microns are considered for sedimentation. Bigger droplets are less probable to stick to sedimented particulate matter, and smaller than 70 microns are already estimated in dispersion process.

### 7.3.8 Dissolution

This process may be quantified through Cohen method, where dissolution rate is estimated by:

$$\frac{dDiss}{dt} = K \cdot f_s \cdot A_s \cdot S \quad (\text{g.h}^{-1}) \quad \text{Eq. 7-42}$$

$f_s$  is the surface fraction covered by oil (considered equal to oil content in emulsion water + oil);  $A_s$  is the oil slick area ( $\text{m}^2$ ) and  $S$  is the oil solubility in water. Huang & Monastero (1982) proposed an analytical solution for the solubility of a typical oil (this model uses the numerical solutions of the following equation):

$$S = S_0 \cdot e^{-\alpha t} \quad \text{Eq. 7-43}$$

where  $S_0$  is the solubility of the “fresh” oil (30 g.m<sup>3</sup>);  $\alpha$  is a decayment constant (0.1);  $t$  is the time after spill (h) and  $K$  is the dissolution mass transfer coefficient (0.01m.h<sup>-1</sup>)

### 7.3.9 Oil-Beaching

When oil reaches a coastal zone, it might become beached. This model estimates the amount of beached oil when the model user predefines a beaching probability (or different beaching probabilities for different coastal zones).

### 7.3.10 Removal techniques

Some removal techniques like chemical dispersion or mechanical cleanup are also included in model.

#### 7.3.10.1 Chemical Dispersion

The application of chemical dispersants is simulated since dispersant efficiency, percentage of oil slick sprayed, and application period are known. The chemical dispersed rate is predicted by the following equation:

$$\frac{dm_{Qchem}}{dt} = \frac{m_{oil} \cdot \left( \frac{\%A_{spr}}{100} \right) \cdot \left( \frac{\%Ef}{100} \right)}{\Delta t} \quad \text{Eq. 7-44}$$

$m_{oil}$  is the instant mass of oil,  $\%A_{spr}$  is the percentage of total slick area sprayed by the chemical dispersant, and  $\%Ef$  is the efficiency of the chemical product.

#### 7.3.10.2 Mechanical Cleanup

Mechanical Cleanup is also simulated for a certain time period, where the volume rate or total emulsion removed during that period must be known.

If emulsion volume rate removed by the skimmer is unknown, it is obtained from the total volume of emulsion mechanically removed in the operation

time period:

$$\frac{dV_{mec}}{dt} = \frac{V_{TotMec}}{\Delta t} \quad \text{Eq. 7-45}$$

Where  $V_{TotMec}$  is the total volume of emulsion mechanically removed ( $\text{m}^3/\text{h}$ )

and  $\frac{dV_{mec}}{dt}$  is the volume rate of emulsion mechanically removed ( $\text{l/h}$ ).

After a conversion of this emulsion volume rate to  $\text{m}^3/\text{s}$ , the rate of oil volume removed is estimated by:

$$\frac{dV_{OilMec}}{dt} = \frac{dV_{mec}}{dt} \cdot (1 - Y) \quad \text{Eq. 7-46}$$

Where  $Y$  is the water content in emulsion water + oil.

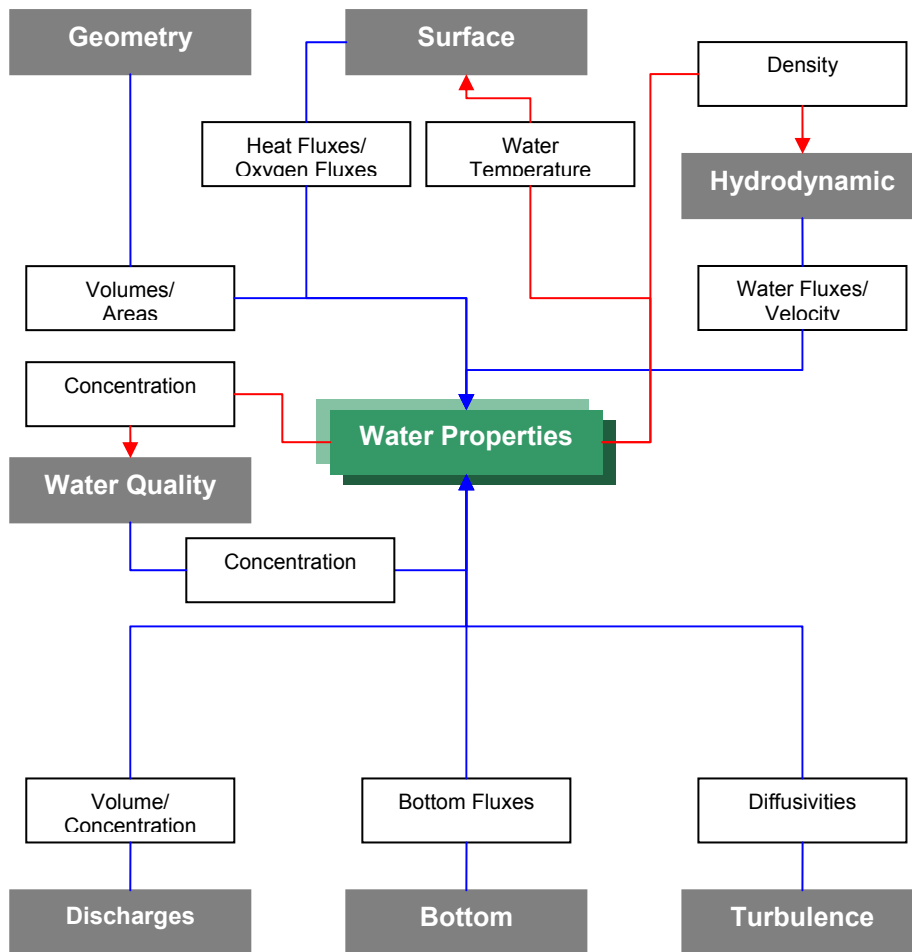
## 8 The Water Properties Module

### 8.1 Introduction

The water properties module coordinates the evolution of the water properties in the water column, using a eulerian approach. This coordination includes the transport due advective and diffuse fluxes, water discharges from rivers or anthropogenic sources, exchange with the bottom (sediment fluxes) and the surface (heat fluxes and oxygen fluxes), sedimentation of particulated matter and the internal sinks and sources (water quality).

Actually the model *Mohid* can simulate 24 different water properties: temperature, salinity, phytoplankton, zooplankton, particulate organic phosphorus, refractory dissolved organic phosphorus, non-refractory dissolved organic phosphorus, inorganic phosphorus, particulate organic nitrogen, refractory organic nitrogen, non-refractory organic nitrogen, ammonia, nitrate, nitrite, biological oxygen demand, oxygen, cohesive sediments, ciliate bacteria, particulate arsenic, dissolved arsenic, larvae and fecal coli-forms. Any new property can be added very easily, due to the object orientated programming used within the *Mohid* model.

In the water quality module, the nitrogen, oxygen and phosphorus cycle can simulate the terms of sink and sources. Figure 8-1 represents the information flux of the water properties module.



**Figure 8-1: Information flux between the Water Properties Module and other modules**

## 8.2 Equations

### 8.2.1 Transport

The transport due advective and diffusive fluxes, of a given property  $A$ , is resolved by the following equation:

$$\begin{aligned}
 \partial_t A = & -\partial_x(uA) - \partial_y(vA) - \partial_z(wA) \\
 & + \partial_x(v'_H \partial_x A) + \partial_y(v'_H \partial_y A) + \partial_z((v'_t + v'_A) \partial_z A)
 \end{aligned}
 \tag{Eq. 8-1}$$

where  $u$ ,  $v$  and  $w$  are the velocity in  $x$ ,  $y$  and  $z$  direction,  $v'_H$  and  $v'_t$  the horizontal and vertical eddy diffusivities, and  $v'_A$  the molecular diffusivity.

The temporal evolution of A is the balance of advective transport by the mean flow and turbulent mixing and the possible sink and sources the property may have.

### 8.2.2 Density

The density  $\rho$  is calculated as a function of temperature and salinity by a simplified equation of state (*Leendertsee and Liu, [1978]*):

$$\rho = (5890 + 38T - 0.375T^2 + 3S) /$$
$$((1779.5 + 11.25T - 0.0745T^2) - (3.8 + 0.01T)S +$$
$$0.698(5890 + 38T - 0.375T^2 + 3S)) \quad \text{Eq. 8-2}$$

That is an approximation for shallow water of the most widely used UNESCO equation (UNESCO, [1981]).

## 9 The Water Quality Module

### 9.1 Introduction

Today, efforts towards ecological modeling are being made in most countries where water quality management is a major concern. Fransz *et al.*, (1991) notice that most new generation models tend to become much more biologically and chemically diversified than earlier models, as it is now largely recognized that there is no way to simulate in sufficient detail the ecosystem behavior without an in-depth treatment of the full cycle of organic matter.

These processes are not foreign to the preoccupations caused by the eutrophication and its various manifestations. Although there is general consensus that the inputs of nutrients to the sea must be reduced there is so far no firm scientific basis to decide upon the extent of such reductions.

An appropriate way of addressing the problem of eutrophication and of testing nutrient reduction scenarios is to simulate the phenomenon with mathematical models. It is probably correct to assume that any ecological model with a sufficiently complex internal structure and the multiple relationships that are found at the lower trophic levels will come close to an answer, provided the right time scale is applied.

The ecological model included in *Mohid* is adapted from EPA, (1985) and pertains to the category of ecosystem simulation models i.e. sets of conservation equations describing as adequately as possible the working and the interrelationships of real ecosystem components. It's not correct to say that the model describes the lower trophic levels with great accuracy. In fact the microbial loop that plays a determinant role in water systems in the recycling processes of organic waste is very simplified in our model.

Lower trophic levels appear in nearly all marine ecosystem simulation models since there is at least a compartment "phytoplankton" required to drive the organic matter cycle. Some early models applied in the North Sea were one-compartment models, especially endeavouring to simulate phytoplankton growth, in relation with the physical environment and with

grazing pressure (treated as a forcing variable). Both the influence of the Lotka-Volterra equations – developed in the 1920s – and that of findings in the field of plant physiology (photosynthesis-light relationship) were discernible. It was not long before limiting nutrient and herbivorous zooplankton were incorporated as well, as state variables in simulation models. (Fransz *et al.*, 1991)

## 9.2 The general model

Franz *et al.* (1991) defined the general conservation equations for an idealized marine ecosystem model. Here we have adapted their definitions and establish a system that consists in five general state variables including phytoplankton, zooplankton, dissolved nutrient, organic matter in pelagic phase, organic matter in benthic phase, pelagic bacteria, benthic bacteria.

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$$dN/dt = -f_{12} \text{ (uptake by phytoplankton)} - f_{15} \text{ (uptake by pelagic bacteria)} + f_{51} \text{ (pelagic mineralization)} + f_{61} \text{ (benthic mineralization)} + f_{01} \text{ (excretion by zooplankton)} + \text{advection and diffusion}$$

$$dP/dt = +f_{12} \text{ (phytoplankton growth)} - f_{23} \text{ (excretion of pOM)} - (f'_{23} + f'_{24}) \text{ (natural mortality)} - f_{20} \text{ (grazing)} - f_{24} \text{ (phytoplankton sinking)} + \text{advection and diffusion.}$$

$$dZ/dt = +f_{20} \text{ (zooplankton growth)} - f_{01} \text{ (excretion of nutrients)} - f_{04} \text{ (excretion of bOM)} - f_{03} \text{ (excretion of pOM)}$$

$$dpOM/dt = +f_{23} \text{ (excretion of pOM)} + f'_{23} \text{ ((1-a).natural mortality of phytoplankton)} + f_{53} \text{ ((1-b).natural mortality of pelagic bacteria)} + f_{03} \text{ ((1-c).fecal pellets and detritus from upper trophic levels)} - f_{35} \text{ (pOM degradation by pelagic bacteria)} + \text{advection and diffusion.}$$

$$dbOM/dt = +f'_{24} \text{ ((a).natural mortality of phytoplankton)} + f_{24} \text{ (phytoplankton sinking)} + f_{54} \text{ ((b). natural mortality of pelagic bacteria)} + f_{64} \text{ (natural mortality of benthic bacteria)} + f_{04} \text{ ((c).fecal pellets and detritus from upper trophic levels)} - f_{46} \text{ (bOM degradation by benthic bacteria)} + \text{advection and diffusion.}$$

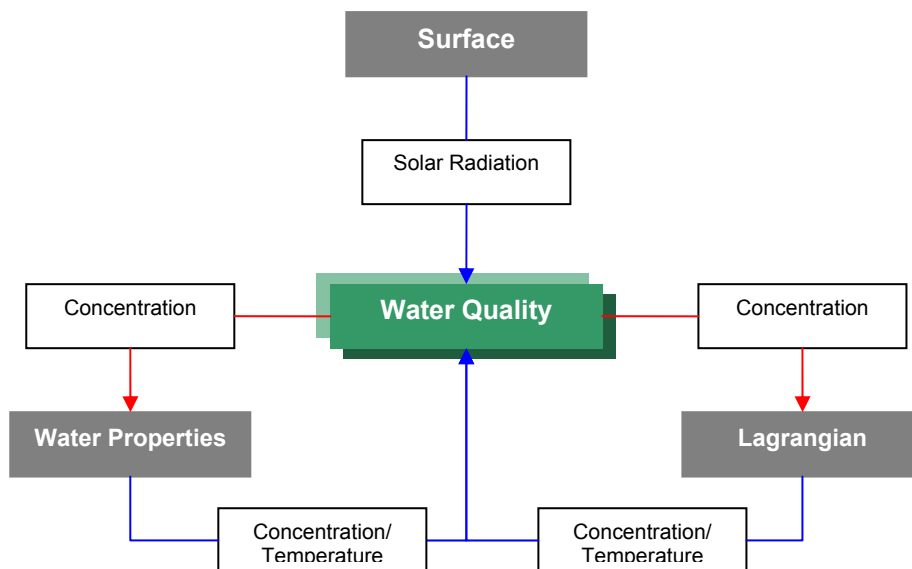
$$dpB/dt = +f_{35} \text{ (pOM degradation)} + f_{15} \text{ (N uptake)} - f_{51} \text{ (pelagic mineralization)} - (f_{53} + f_{54}) \text{ (natural mortality)} + \text{advection and diffusion.}$$

$$dpB/dt = +f_{46} \text{ (bOM degradation)} - f_{61} \text{ (benthic mineralization)} - f_{64} \text{ (natural mortality)}.$$

where  $N$  represents the concentration of dissolved inorganic nutrient,  $P$  the concentration of concentration of phytoplankton,  $Z$  the concentration of zooplankton,  $pOM$  the concentration of pelagic organic matter,  $bOM$  the concentration of benthic organic matter,  $pB$  the concentration of pelagic bacteria,  $bB$  the concentration of benthic bacteria and  $a, b, c$  factors comprised between 0 and 1.

The primary production process, powered by light energy, is the necessary engine for all transfers of mass between biological compartments. Zooplankton that on early days was not explicitly modeled it's now considered an important state variable.

The *Mohid* Water Quality module is a zero-dimensional ecological model, which can be used by the eulerian or the lagrangian transport modules. The nitrogen cycle, oxygen cycle and the phosphorus cycle are included. A brief description of these cycles is presented in the next sections. Figure 9-1 represents the information flux between the water quality module and other modules.



**Figure 9-1: Information flux between the water quality module and other modules**

The water quality module has been developed in terms of sinks and sources. Such an approach is convenient to give these models the desired flexibility. Because of the properties interdependency a linear equation system is computed for each control volume and this system can be compute forward or backward in time.

Many of the equations described in the next sections are written as dependent on a regulating factor, which contains the functional response of the organism to some environmental parameters such as light, nutrients or temperature. When growth is a function of many resources, there is a large range of functional forms that might express the joint dependence. To control the various possibilities, it is common to think of separate resources as limiting factors reducing some theoretical maximum growth rate - factors that can be determined separately and the combined by a small number of ways.

Each growth limitation factor can range from a value of 0 to 1. A value of 1 means the factor does not limit growth (i.e. is at optimum intensity, nutrients are available in excess, etc) and a value of 0 means the factor is so severely limiting that growth is inhibit entirely.

Four major approaches have been used to combine the limiting factors:

- A multiplicative formulation in which all factors are multiplied together. This approach assumes that several nutrients in short supply will more severely limit growth than a single nutrient in short supply. The major criticism of this approach is that the computed growth rates may be excessively low when several nutrients are limiting. Also, the severity of the reduction increases with the number of limiting nutrients considered in the model, making comparison between models difficult.
- A minimum formulation in which the most severely limiting factor alone is assumed to limit growth. This formulation is based on "Liebig's law of the minimum" which states that the factor in shortest supply will control the growth of algae. The minimum formulation is often used only for nutrient limitation, with a multiplicative formulation for the light limitation factor.

- A harmonic mean formulation that combines the reciprocal of each limiting factor in the following manner:

$$f(\text{Light}, \text{Nutrient}_1, \text{Nutrient}_2, \dots, \text{Nutrient}_n) = \frac{n}{\frac{1}{f(\text{Light})} + \frac{1}{f(\text{Nutrient}_1)} + \dots + \frac{1}{f(\text{Nutrient}_n)}} \quad \text{Eq. 9-1}$$

where  $n$  = number of limiting factors.

This formulation is based on an electronic analogy of several resistors in series. The rationale for this formulation is that it includes some interactions between multiple limiting nutrients, but is not as severely limiting as the multiplicative formulation. Under a wide range of conditions, the harmonic mean formulation and minimum formulation produce similar growth response curves (Swartzman and Bentley, 1979 in EPA, 1985).

- An arithmetic mean formulation that uses the average of each limiting factor. The rationale for this formulation is the same as for the harmonic mean formulation. However, this formulation is rarely used since it does not restrict growth enough. For example, the arithmetic mean formulation allows growth even if a critical nutrient such nitrogen is totally absent, as long other nutrients are available.

### 9.3 Phytoplankton

The growth of phytoplankton is limited to several factors, like described in the following sections.

#### 9.3.1 Nutrient limitation

The model considers nitrogen (ammonia and nitrate) and phosphorus to be the nutrients that limits phytoplankton growth. Nitrate and ammonia are considered in the same pool. But difficulties could be encountered to subtract phytoplankton uptake from the ammonia and nitrate pool and this problem is solved by introduction of the ammonia preference factor ( $\beta_{NH4}$ ). The nutrient limitation is expressed in a Michaelis-Menten form, with half saturation constant ( $K_N$ ). In the case of ammonia and nitrate, the model considers:

$$\Psi(N)_{Phy} = \frac{NH_4 + NO_3}{K_N + NH_4 + NO_3} \quad \text{Eq. 9-2}$$

where  $\Psi(N)_{Phy}$  represents the nutrient limitation due nitrogen presence,  $NH_4$  and  $NO_3$  the ammonia and nitrate concentrations (mg N.L<sup>-1</sup>) and  $K_n$  the half-saturation constant for nitrogen limitation (mg N.L<sup>-1</sup>).

In the case of phosphorus the above equation takes the form:

$$\Psi(P)_{Phy} = \frac{PO_4}{K_p + PO_4} \quad \text{Eq. 9-3}$$

where  $\Psi(P)_{Phy}$  represents the nutrient limitation due phosphorus presence,  $PO_4$  the phosphorus concentration (assumed to be completely available as orthophosphate) (mg P.L<sup>-1</sup>) and  $K_p$  the half-saturation constant for phosphorus limitation (mg P.L<sup>-1</sup>).

The nutrient limitation factor is given by the minimum of  $\Psi(N)_{Phy}$  and  $\Psi(P)_{Phy}$ .

### 9.3.2 Temperature limitation

The concept of Thornton and Lessen, (1978) is adopted to represent temperature limitation factor ( $\Psi(T)$ ) on autotrophy and heterotrophy organisms:

$$\Psi(T) = K_A(T) \cdot K_B(T) \quad \text{Eq. 9-4}$$

where  $K_A(T)$  is defined by:

$$K_A(T) = \frac{K_1 \cdot e^{\gamma_1 \cdot (T - T_{\min})}}{1 + K_1 \cdot (e^{\gamma_1 \cdot (T - T_{\min})} - 1)} \quad \text{Eq. 9-5}$$

with

$$\gamma_1 = \frac{Ln \frac{K_2(1 - K_1)}{K_1(1 - K_2)}}{T_{opt_{\min}} - T_{\min}} \quad \text{Eq. 9-6}$$

where  $K_B(T)$  is defined by:

$$K_B(T) = \frac{K_4 \cdot e^{\gamma_2 \cdot (T_{\max} - T)}}{1 + K_4 \cdot (e^{\gamma_2 \cdot (T_{\max} - T)} - 1)} \quad \text{Eq. 9-7}$$

with

$$\gamma_2 = \frac{\text{Ln} \frac{K_3(1 - K_4)}{K_4(1 - K_3)}}{T_{\max} - \text{Topt}_{\max}} \quad \text{Eq. 9-8}$$

$\text{Topt}_{\min}$  (°C) and  $\text{Topt}_{\max}$  (°C) represent the temperature interval for an optimal process, and  $T_{\max}$  (°C) and  $T_{\min}$  (°C) the maximum and minimum tolerable temperature where processes are completely inhibited. Remaining constants ( $K_1$ ,  $K_2$ ,  $K_3$  and  $K_4$ ) control the shape of the response curve of temperature effect; these values are assumed equal for all organisms in this model.

### 9.3.3 Light limitation

Photosynthesis is possible only when light reaching the algae cell is above certain intensity. This means that phytoplankton is limited to the uppermost layers of the water column where light intensity is sufficient for photosynthesis to occur. The depth to which light will penetrate in water, and hence the depth at which production can occur, is dependent on a number of factors; these include absorption of light by water, the wavelength of light, transparency of water, reflection from the surface of the water, reflection from suspended particles, latitude, and season of the year.

The solar radiation depends on factors such as clouds and dust in the atmosphere and the solar elevation. The calculation of the solar radiation is described in the surface module.

When light strikes the surface of water, a certain amount of light is reflected back; the amount depends on the angle at which the light strikes the surface of water. If the angle from the horizontal is low, a large amount will be reflected. Conversely, the nearer the angle is to 90° (that is

perpendicular to the horizontal surface of the water), the greater will be the penetration and the lesser will be the reflection (Nybakken 1993). The angle at which the light strikes the surface of the water is directly related to the maximum height of the sun above the horizon.

The extinction of light in the marine environment is one of the important water quality variables often addressed by aquatic scientists and oceanographers. The characteristics of the underwater light field itself are a classical subject of oceanographic optics (Rivera, 1997). The available light is one of the primary limiting variables in the growth of submerged flora, besides nutrients and temperature. Light availability is of major importance not only in determining how much plant growth will be but also which kind of species will predominate and which kind will evolve (Rivera, 1997). Vertical light attenuation and its spectral distribution are related to the absorption by the water itself and the following additional components of the water column: photosynthetic organisms, suspended particles and soluble compounds. Modeling light attenuation is the basis to predict the intensity and spectral composition of available light for phototropic populations (Vila *et al*, 1996).

The rate of the light reaction of photosynthesis is strictly dependent on light intensity. Increases in light intensity lead to greater photosynthetic rates until some maximum is reached. At this point the producers cannot use any more light. The enzymes involved in photosynthesis cannot act fast enough to process light quanta any faster, so rate of photosynthesis reaches an asymptote. Increasingly higher light intensities usually inhibit photosynthesis (Valiela, 1995).

#### **9.3.3.1 Light extinction in water**

Kirk (1980) defines the inherent optical properties as the absorption, scattering and beam attenuation coefficients of a medium. The absorption coefficient is defined as the fraction absorbed per unit of path length from a parallel beam of monochromatic light directed normal to an infinitesimally thin layer of medium. Similarly, the scattering coefficient is defined as the fraction scattered of the incident parallel beam divided by the path length. The beam attenuation coefficient is defined as the sum of the absorption

and scattering coefficients.

By definition, the incident light field or downward irradiance in a water column refers to the instantaneous value of the down-welling radiant flux in a horizontal unit area.

Kirk (1980) differentiates between downward and upward irradiance, the first being that due to down-welling stream of light and the second due to the upwelling stream of light. In light extinction studies, the desirable quantity is the down-welling PAR which is referred to as the downward irradiance covering the 400 – 700 nm range of the wave spectrum. The down-welling PAR is attenuated due to both scattering and absorption processes by the optically active components in the water column.

The major light absorbing and scattering components in the water column include dissolved organic substances, dead and living plankton material, suspended inanimate particles, and water itself. These components differ in the way they absorb and scatter downward irradiance across the photosynthetic wave band.

Generally, the strong absorption in inland and estuarine waters is attributed to organic substances, gelatin and/or phytoplankton. On the other hand, scattering, as pointed by Kirk (1980), does not itself “remove” light since a scattered photon is still available for photosynthesis. However, by making the photons follow a zig-zag path, the probability of being absorbed by the absorbing components in the aquatic medium is increased. Hence, with the scattering contribution of suspended particles for example, the vertical attenuation is intensified through this mechanism.

A common method often employed in modeling the extinction of downward irradiance is to consider the influence of the major optically active components separately giving partial extinction coefficients for each component. The sum of all the partial extinction coefficients gives the average extinction coefficient of the water column (Rivera, 1997).

Light extinction in natural waters is affected by four primary groups of substances whose composition and concentration differ in each water body giving different values of the extinction coefficient. Further more, the

extinction coefficient may change with time due to the varying composition and concentrations of the primary factors. These factors, which are referred to as optically active components of the water column, include inanimate suspended solids, dead or living phytoplankton (algae), gelvins and water itself (Rivera, 1997). Parson *et al.* (1984) uses this concept when defining the extinction coefficient in the water column ( $k$ ) as follows:

$$k = k_w + k_p + k_d + k_s \quad \text{Eq. 9-9}$$

where  $k_w$ ,  $k_p$ ,  $k_d$ , and  $k_s$  represent diffusion and scattering of light energy due to water (w), phytoplankton (p), suspended particles other than phytoplankton (d), and dissolved matter (s), respectively. The suspended particles include many different forms such as clay particles, organic detritus, and organisms varying in size. Each of these extinction coefficients are highly dependent on wavelength, however according to Parson *et al.* (1984), for the purpose of most biological events, the average extinction coefficient in the wavelength of PAR rather than the value at particular wavelengths is probably the most practical.

The partial extinction coefficients can be determined from the specific extinction coefficient and the concentration of the optically active components of the water column by the relation:

$$k_n = \kappa_n c_n \quad \text{Eq. 9-10}$$

where  $k_n$  is the extinction coefficient of a particular component  $n$ ,  $\kappa_n$  the specific extinction of that component and  $c_n$  the observed concentration.

The majority of the water quality models revised (e.g. Vila e Garcia-Gil, 1996 Arhonditsis *et al.* 2000, Napolitano *et al.*, 2000 Nakata *et al.*, 2000, Kawamiya *et al.*, 2000, Humborg *et al.*, 2000, Neuman, 2000, Tett and Wilson, 2000) compute the extinction coefficient considering only phytoplankton self-shading effect. The general form of the established relation is usually like the next equation, with different set of parameters determined according to local measurements.

$$k = k_w + \kappa_{phy} c_{phy} \quad \text{Eq. 9-11}$$

Cole and Buchak (1995) and Somlyódy and Koncsos (1991) are some examples where the extinction coefficient is computed considering not only the phytoplankton concentration but also sediment concentration.

Each of these specific extinction values can represent a problem of there one in terms of modeling. A usual solution is to develop a relationship based in local measurements that allow us to determine the overall extinction coefficient. This kind of relationship can be dependent on one of the factors already described (usually phytoplankton) but does not specifically distinguish between the chosen factor and other materials. Parson *et al.*, (1984) presents a equation of this kind derived from field observations carried out in the western North Atlantic, which is used by several authors (Yanagi *et al.*, 1997; Miranda 1997). This equation relates the average extinction coefficient (k) to the chlorophyll a concentration (C) for natural phytoplankton community as follows:

$$k = 0.04 + 0.0088C + 0.054C^{2/3} \quad \text{Eq. 9-12}$$

The coefficients to compute the extinction parameter are determined by the local light conditions of the study area. Portela (1996) following the observations made by Martins e Duffner, (1982) on the Tagus estuary obtained an average value for the extinction coefficient of 4.5 m<sup>-1</sup> and a median value of 3.4 m<sup>-1</sup>. Portela (1996) applied a linear regression model to the observed values of extinction coefficient and the concentration of suspended sediments measured in the Tagus estuary in 1980 (Martins e Duffner, 1982). As expected, a close relation between the two variables is observed. The final regression equation is:

$$k=1.24 + 0.036C_{ss} \quad \text{Eq. 9-13}$$

Another formulation included in the Mohid water quality module, which calculates the effect on light attenuation, depending on phytoplankton and sediment concentration, was presented by Pina (2001):

$$k = 0.04 + 0.0088C + 0.054C^{2/3} + 0.036C_{ss} \quad \text{Eq. 9-14}$$

### 9.3.3.2 Phytoplankton reaction to light

The rate of the light reaction of photosynthesis is strictly dependent on light

intensity. Increases in light intensity lead to greater photosynthetic rates until some maximum is reached. At this point the producers cannot use any more light, the enzymes involved in photosynthesis cannot act fast enough to process light quanta any faster, so rate of photosynthesis reaches an asymptote. Increasingly higher light intensities usually inhibit photosynthesis (Valiela, 1995).

During the last decades a considerable amount of research has been carried out on primary productivity modeling (e.g. Steele, 1962; Jassby and Platt, 1976; Platt *et al.*, 1980; Falkowski & Wirick, 1981; Eilers and Peeters, 1988). In most of these works formulations of the relationship between primary productivity and light intensity were proposed and tested against field and/or laboratory data. Most of these equations are empirical, only a few of them being deduced from the physiology of photosynthesis (e.g. Fasham and Platt, 1983; Eilers and Peeters, 1988). These formulations have been used for several years in ecological models.

The light intensity affects only the photosynthesis, its representation use the formulation of Steele (1962) integrated on the depth, Parsons *et al.*, (1995) for this zero-dimensional model and a classic Beer-Lambert function for the light intensity:

$$E(z) = E_0 \cdot e^{(-k(p)z)} \quad \text{Eq. 9-15}$$

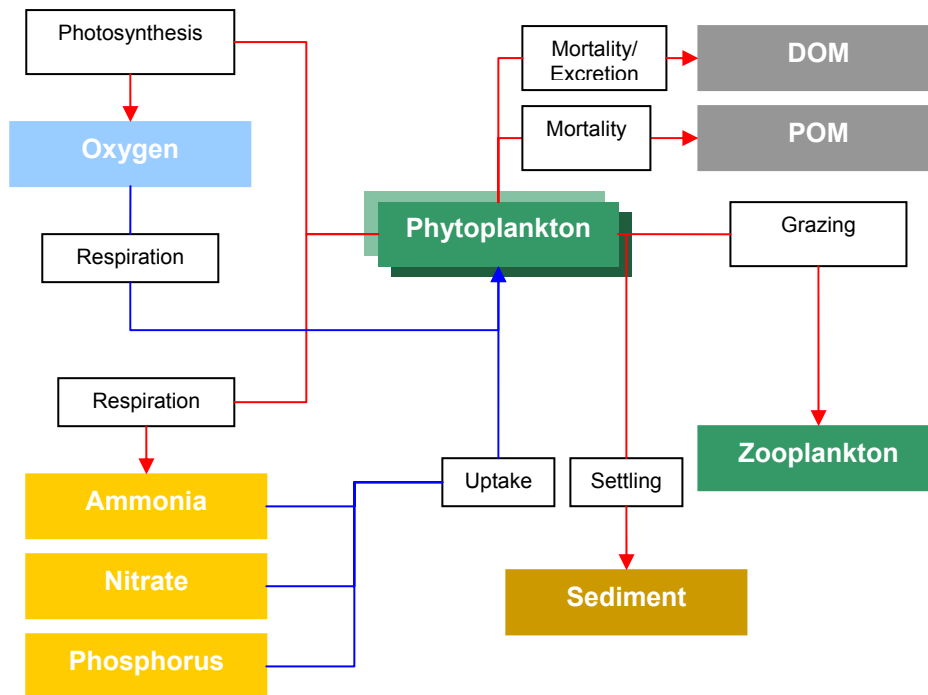
with

$$\Psi(E) = \frac{e^1}{k(p).z} \cdot \left( e^{\frac{E_0}{E_{opt}} \cdot e^{(-k(p)z)}} - e^{\frac{E_0}{E_{opt}}} \right) \quad \text{Eq. 9-16}$$

$E_0$  represents the effective solar radiation at the water surface ( $W.m^{-2}$ ),  $k(p)$  the light extinction factor ( $m^{-1}$ ),  $E_{opt}$  the optimal light intensity for photosynthesis and  $z$  the depth (m).

### 9.3.4 Equations

Figure 5-1 represents the internal fluxes of phytoplankton modeled by *Mohid's* water quality module.



**Figure 9-2: Internal Flux of Phytoplankton**

Phytoplankton is described in terms of carbon concentration ( $\text{mgC} / \text{l}$ ). The model assumes three limitations affecting the maximum phytoplankton growth rate,  $\mu_{max}$ . Temperature  $\Psi(T)$ , light effect  $\Psi(E)$  and nutrient limitation (minimum of  $\Psi(N)_{Phy}$  and  $\Psi(P)_{Phy}$ ), like described in the previous chapter.

The simulation of the phytoplankton is developed with the following considerations: it consumes inorganic nutrients (ammonia and nitrate from the nitrogen cycle and inorganic phosphorus from the phosphorus cycle) depending on their availability. Another factor which influences the growth of phytoplankton is the availability of light as a source of energy for photosynthesis. During the process photosynthesis dissolved oxygen is produced. The respiration process consumes oxygen and produces ammonia. The Excretion of Phytoplankton produces dissolved organic material (Refractory Dissolved Organic Nitrogen, Non-Refractory Dissolved Organic Nitrogen, Refractory Dissolved Organic Phosphorus and Non-Refractory Dissolved Organic Phosphorus). By mortality phytoplankton increases the dissolved organic material and the particulate organic material (Particulate Organic Nitrogen and Particulate Organic

Phosphorus). By the grazing of phytoplankton by zooplankton, the concentration of phytoplankton decreases. The settling process is modeled in the water properties module.

The rate equation of phytoplankton, used by *Mohid*, can be written as:

$$\frac{\partial \Phi_{Phy}}{\partial t} = (\mu_{Phy} - r_{Phy} - ex_{Phy} - m_{Phy}) \Phi_{Phy} - G \quad \text{Eq. 9-17}$$

The growth rate,  $\mu_{Phy}$  ( $\text{day}^{-1}$ ), is given by:

$$\mu_{Phy} = \mu_{max} \cdot \min(\Psi(N)_{Phy}, \Psi(P)_{Phy}) \cdot \Psi(E)_{Phy} \cdot \Psi(T)_{Phy} \quad \text{Eq. 9-18}$$

where  $\mu_{max}$  represents the maximum growth rate.

The respiration,  $r_{Phy}$  ( $\text{day}^{-1}$ ), is given by:

$$r_{Phy} = k_{er} \cdot \exp(0.069 \cdot T^{\circ}) + k_p \mu_{Phy} \quad \text{Eq. 9-19}$$

where  $k_{er}$  represents the endogenous respiration constant and  $k_p$  the photorespiration factor.

The excretion,  $ex_{Phy}$  ( $\text{day}^{-1}$ ), is given by:

$$ex_{Phy} = \varepsilon_{Phy} \cdot \mu_{Phy} (1 - \Psi(E)_{Phy}) \quad \text{Eq. 9-20}$$

The natural mortality,  $m_{Phy}$  ( $\text{day}^{-1}$ ), is given by:

$$m_{Phy} = m_{max} \cdot \frac{\frac{\Phi_{Phy}}{\mu_{Phy}}}{Km + \frac{\Phi_{Phy}}{\mu_{Phy}}} \quad \text{Eq. 9-21}$$

where  $m_{max}$  represents the maximum mortality and  $K_m$  the mortality half saturation rate.

The grazing,  $G$ , is given by:

$$G = \frac{g_z}{E} \Phi_z \quad \text{Eq. 9-22}$$

where  $g_z$  represents the net growth rate of zooplankton,  $E$  the assimilation efficiency and  $\Phi_z$  the concentration of zooplankton.

Symbol	Coefficient	Unit	Value
$\mu_{max}$	Maximal growth Rate	$d^{-1}$	2.2
$\varepsilon_{Phy}$	Excretion constant	-	0.07
$k_{er}$	Endogenous respiration constant	-	0.0175
$k_p$	Photorespiration factor	-	0.125
$K_m$	Mortality half saturation rate	$mgC.d.L^{-1}$	0.3
$m_{max}$	Maximal mortality rate	$d^{-1}$	0.02
$E$	Assimilation efficiency		0.6
<u>Nutrient limitation</u> $K_N$	Half saturation constant for nutrient limitation	$mgN.L^{-1}$	0.014
<u>Temperature limitation</u>			
$Topt_{minPhy}$	Minimal temperature for an optimal photosynthesis	$^{\circ}C$	25.0
$Topt_{maxPhy}$	Maximal temperature for an optimal photosynthesis	$^{\circ}C$	26.5
$Temp_{minPhy}$	Minimal temperature for photosynthesis	$^{\circ}C$	4.0
$Temp_{maxPhy}$	Maximal temperature for photosynthesis	$^{\circ}C$	37.0
<b><math>Eopt</math></b>	Optimal light intensity for photosynthesis	$Wm^{-2}$	100
$\alpha_{Phy}$	Redfield ratio (N:C)	-	0.18
$K_{PhyNutReg}$	phytoplankton nutrient regeneration half saturation rate	$mgN.L^{-1}$	1.0
$F_{org P/Ph}$	Fraction of organic particulate excretion		0.7
$f_{in/Phy}$	Fraction of inorganic excretion	-	0.5
$f_{orgD/Phy}$	Fraction of organic soluble excretion	-	0.4

**Table 9.3-1: Values used in the standard model for the phytoplankton variable**

## 9.4 Zooplankton

Zooplankton is described in terms of carbon concentration ( $mg C l^{-1}$ ) and the net growth rate,  $g_z$  ( $day^{-1}$ ), is obtained from Ivlev, (1945) adapted by Parsons *et al.*, (1967). Respiration and non-predatory mortality of the

zooplankton ( $\text{day}^{-1}$ ),  $r_z$  and  $m_{Z_0}$  are considered functions of temperature, being treated as one variable. The predatory mortality,  $G_z$ , depends on the zooplankton concentration.

Figure 9-3 represents the internal flux of zooplankton.

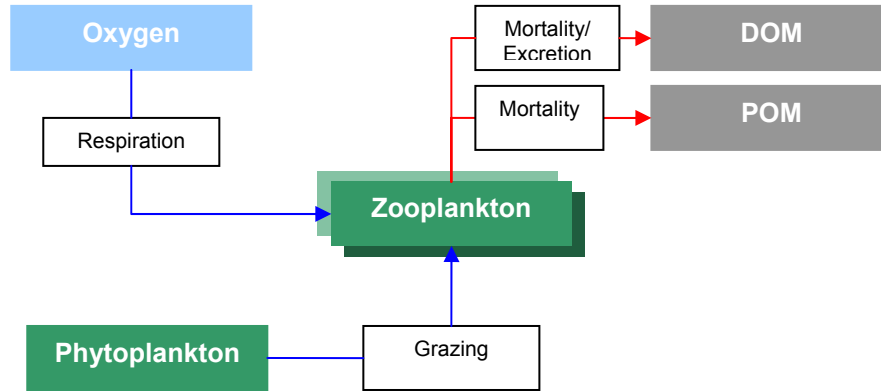


Figure 9-3: Internal Flux of Zooplankton

#### 9.4.1 Equations

The growth of zooplankton is given by:

$$\frac{\partial \Phi_Z}{\partial t} = (g_z - r_z - m_z) \cdot \Phi_Z \cdot G_Z \quad \text{Eq. 9-23}$$

The growth rate,  $g_z$  ( $\text{day}^{-1}$ ), is given by:

$$g_z = g_{\max} \cdot (T_{\text{ref}}) \cdot \Psi(T)_Z \cdot \left(1 - e^{-\Delta(\Phi_{\text{phy}} - \Phi_{\text{phy0}})}\right) \quad \text{Eq. 9-24}$$

where  $g_{\max}$  represents the maximum growth rate,  $\Delta$  stands for the Ivlev constant,  $\Omega_{\text{phy0}}$  represents the minimum concentration of phytoplankton for grazing. The temperature limitation is calculated in the same way as for phytoplankton, but with other constants.

The natural mortality and respiration,  $r_z + m_z$  ( $\text{day}^{-1}$ ), is given by:

$$r_z + m_z = d_z \cdot (T_{\text{ref}}) \cdot \Psi(T) \quad \text{Eq. 9-25}$$

where  $d_z$  represents the natural mortality and respiration rate.

Grazing,  $G_z$  ( $\text{day}^{-1}$ ), is given by:

$$G_z = e_z Z \quad \text{Eq. 9-26}$$

where  $e_z$  represents the predatory mortality rate.

Symbol	Coefficient	Unit	Value
$g_{max}$	Maximal growth Rate	$\text{d}^{-1}$	0.1
$\Delta$	Ilev constant	-	13.0
$Phy_0$	Minimum phytoplankton concentration for grazing	mgC/l	0.045
$d_z$	Non predatory mortality and respiration rate	$\text{d}^{-1}$	0.036
$e_z$	Predatory mortality rate	$\text{d}^{-1}$	0.01
<u>Temperature limitation</u>			
$Topt_{minZo}$	Minimal temperature for an optimal zoo growth	$^{\circ}\text{C}$	24.8
$Topt_{maxZo}$	Maximal temperature for an optimal zoo growth	$^{\circ}\text{C}$	25.1
$Temp_{minZo}$	Minimal temperature for zoo growth	$^{\circ}\text{C}$	5.0
$Temp_{maxZo}$	Maximal temperature for zoo growth	$^{\circ}\text{C}$	35.0
$\alpha_{Phy}$	Redfield ratio (N:C)	-	0.18

**Table 9.4-1: Values used in the standard model for the zooplankton variable**

## 9.5 Nitrogen

In the Mohid water quality module, the nitrogen appears as organic and inorganic nitrogen.

The inorganic nitrogen is divided into ammonia ( $\text{NH}_4$ ), nitrite ( $\text{NO}_2$ ) and nitrate ( $\text{NO}_3$ ).

The organic nitrogen is divided into particulate organic nitrogen (PON), dissolved organic nitrogen non refractory (DONnr) and dissolved organic nitrogen refractory (DONre). DONnr includes small molecular substrates, assumed to be degraded in the day of production and DONre with a longer

turn over.

### 9.5.1 Ammonia

The sources of ammonia are the organic forms of nitrogen (PON, DONnr and DONre) due to decay and phytoplankton due to the dark respiration process. The sinks of ammonia are nitrite (nitrification) and phytoplankton (uptake).

Figure 9-4 represents the internal fluxes of ammonia modeled by *Mohid's* water quality module.

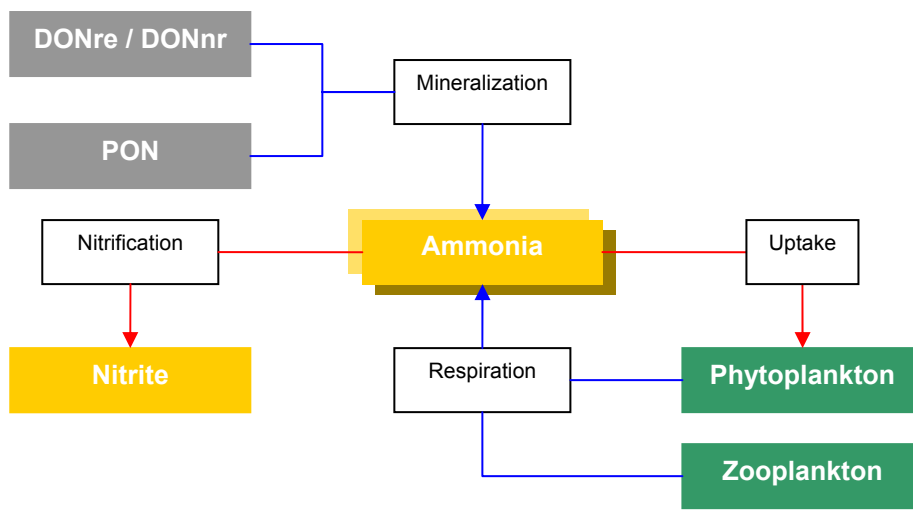


Figure 9-4: Internal Flux of Ammonia

The rate equation of ammonia is given by:

$$\frac{\partial \Phi_{NH_4}}{\partial t} = (f_{in/Phy} ex_{Phy} - \mu_{NH_4}) \Phi_{Phy} + f_{in/Z} ex_z \Phi_z + \varphi_{Nre} \Phi_{DONre} + \varphi_{Nnr} \Phi_{DONnr} + f_{orgP/Phy} \varphi_{det} \Phi_{PON} \quad \text{Eq. 9-27}$$

The assimilation rate of  $NH_4$ ,  $\mu_{NH_4}$ , is given by:

$$\mu_{NH_4} = \beta_{NH_4} \alpha_{N:C} \mu_{Phy} \quad \text{Eq. 9-28}$$

where  $\beta_{NH_4}$  is the ammonia preference factor given by:

$$\beta_{NH_4} = \frac{\Phi_{NH_4} \cdot \Phi_{NO_3}}{(K_N + \Phi_{NH_4})(K_N + \Phi_{NO_3})} \cdot \frac{\Phi_{NH_4} \cdot K_N}{(K_N + \Phi_{NH_4})(K_N + \Phi_{NO_3})} \quad \text{Eq. 9-29}$$

and  $\alpha_{N:C}$  represents the Redfield ratio between N:C.

The mineralization rate of DONre,  $\varphi_{Nre}$  is given by:

$$\varphi_{Nre} = M_{DONre} \theta_{DONre} (T - T_{ref}) \frac{\Phi_{Phy}}{K_{PhyNutRege} + \Phi_{Phy}} \quad \text{Eq. 9-30}$$

where

$M_{DONre}$  – reference rate for the mineralization of DONre

$\theta_{DONre}$  – temperature coefficient for the mineralization of DONre

$T_{ref}$  – reference temperature

$K_{PhyNutRege}$  – half saturation constant for the regeneration of phytoplankton

The mineralization rate of DONnr,  $\varphi_{Nnr}$  is given by:

$$\varphi_{Nnr} = M_{DONnr} \theta_{DONnr} (T - T_{ref}) \quad \text{Eq. 9-31}$$

where

$M_{DONnr}$  – reference rate for the mineralization of DONnr

$\theta_{DONnr}$  – temperature coefficient for the mineralization of DONnr

The dissolution rate of PON,  $\varphi_{det}$ , is given by:

$$\varphi_{det} = M_{det} \theta_{det} (T - T_{ref}) \quad \text{Eq. 9-32}$$

where

$M_{det}$  – reference rate for the dissolution of PON

$\theta_{det}$  – temperature coefficient for the dissolution

### 9.5.2 Nitrite

The source of nitrite, modeled by *Mohid*, is ammonia and the sink is nitrate. Figure 9-5 represents the internal fluxes of nitrite modeled by *Mohid's* water quality module.

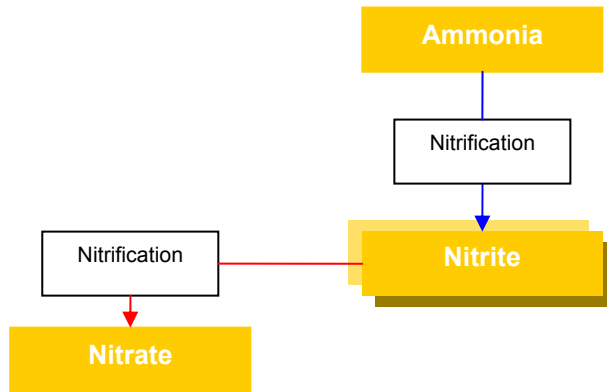


Figure 9-5: Internal Flux of Nitrite

The rate equation of nitrite is given by:

$$\frac{\partial \Phi_{NO_2}}{\partial t} = \varphi_{2N} \Phi_{NH_4} - \varphi_{2N} \phi_{NO_2} \quad \text{Eq. 9-33}$$

with the rate of nitrification,  $\varphi_{2N}$  is given by:

$$\varphi_{2N} = M_{nit} \theta_{nit} (T - T_{ref}) \frac{\Phi_{O_2}}{K_{nit} + \Phi_{O_2}} \quad \text{Eq. 9-34}$$

where

$M_{nit}$  – reference rate of nitrification

$\theta_{nit}$  – temperature coefficient for nitrification

$K_{nit}$  – half saturation constant for nitrification

### 9.5.3 Nitrate

The source of nitrate, modeled by *Mohid*, is nitrite and the sink the uptake by phytoplankton. Figure 9-6 represents the internal fluxes of nitrate

modeled by *Mohid's* water quality module.

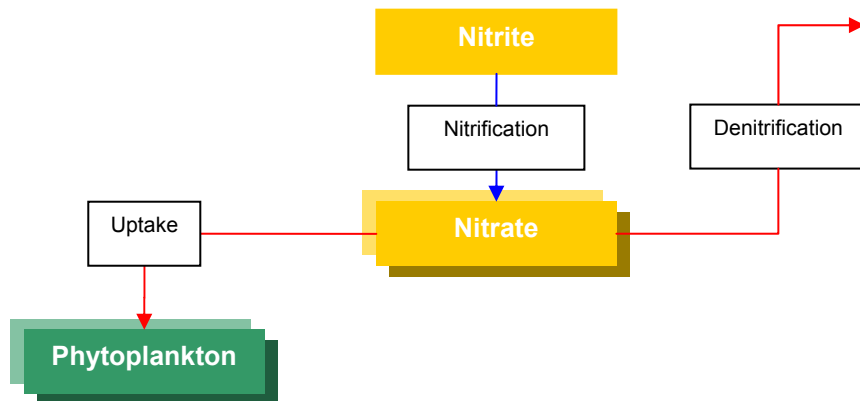


Figure 9-6: Internal Flux of Nitrate

The rate equation of nitrate is given by:

$$\frac{\partial \Phi_{NO_3}}{\partial t} = \varphi_{2N} \Phi_{No_2} - \varphi_{2N} \phi_{NO_3} - \mu_{NO_3} \quad \text{Eq. 9-35}$$

The assimilation rate of  $NO_3$ ,  $\mu_{NO_3}$ , is given by:

$$\mu_{NO_3} = (1 - \beta_{NH_4}) \alpha_{N:C} \mu_{Phy} \quad \text{Eq. 9-36}$$

#### 9.5.4 Particulate organic nitrogen – PON

The sources of PON are the mortality of phytoplankton and zooplankton and the sinks are the mineralization to ammonia and the decomposition to DONre.

Figure 9-7 represents the internal fluxes of PON modeled by *Mohid's* water quality module.

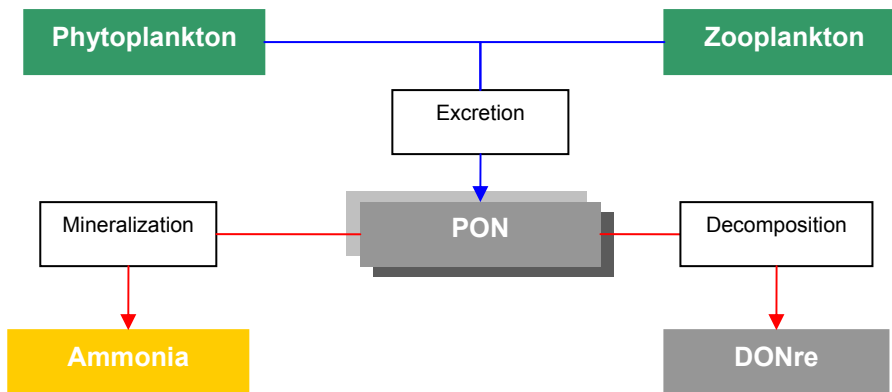


Figure 9-7: Internal Flux of PON

The rate equation of the PON is given by:

$$\frac{\partial \Phi_{PON}}{\partial t} = \left[ (1 - f_{orgD/Phy}) (1 - f_{in/Phy}) ex_{Phy} + m_{Phy} \right] \Phi_{Phy} + \left[ (1 - f_{orgD/Z}) (1 - f_{in/Z}) ex_Z + m_Z \right] \Phi_Z - \varphi_{det} \Phi_{PON} \quad \text{Eq. 9-37}$$

All variables have the same meaning as in the previous paragraphs.

### 9.5.5 Dissolved organic nitrogen non refractory – DONnr

The sources of DONnr are the mortality and the excretions of phytoplankton and zooplankton and the sinks are the mineralization to ammonia.

Figure 9-8 represents the internal fluxes of DONnr modeled by *Mohid's* water quality module.

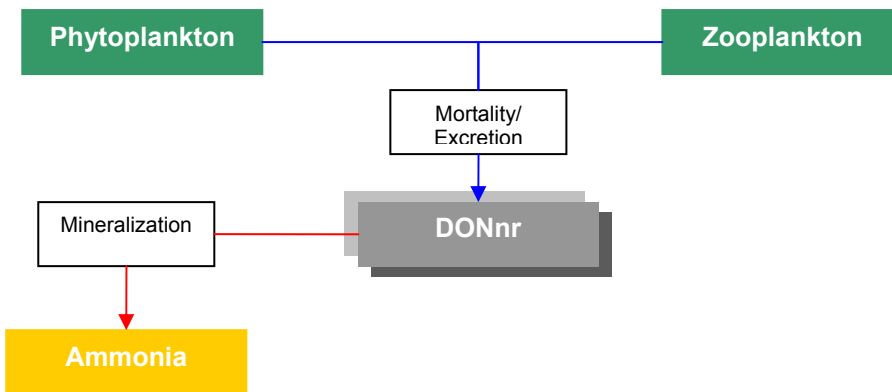


Figure 9-8: Internal Flux of DONnr

The rate equation of the DONnr is given by:

$$\frac{\partial \Phi_{DONnr}}{\partial t} = f_{orgD/Phy} (1 - f_{in/Phy}) ex_{Phy} \Phi_{Phy} + f_{orgD/Z} (1 - f_{in/Z}) ex_Z \Phi_Z - \varphi_{Nnr} \Phi_{DONnr} \quad \text{Eq. 9-38}$$

All variables have the same meaning as in the previous paragraphs.

### 9.5.6 Dissolved organic nitrogen refractory – DONre

The source of DONre is the decomposition of the PON and the sink is the mineralization to ammonia.

Figure 9-9 represents the internal fluxes of DONre modeled by *Mohid's* water quality module.

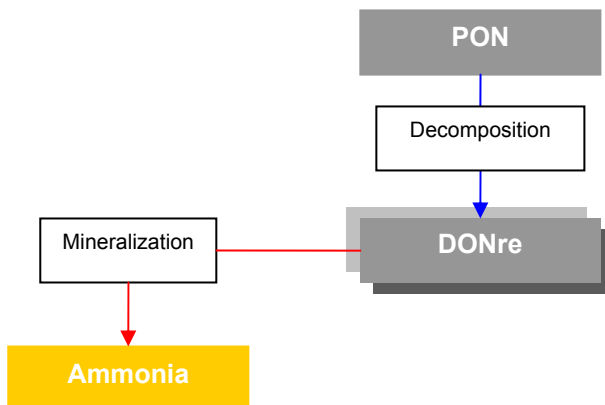


Figure 9-9: Internal Flux of DONre

The rate equation of the DONre is given by:

$$\frac{\partial \Phi_{DONre}}{\partial t} = \varphi_{det} \Phi_{PON} (1 - f_{orgP/Phy}) - \varphi_{Nr} \Phi_{DONre} \quad \text{Eq. 9-39}$$

All variables have the same meaning as in the previous paragraphs.

Symbol	Coefficient	Unit	Value
$T_{ref}$	Reference temperature	°C	25
$M_{DONre}$	Reference rate for mineralization	$d^{-1}$	0
$M_{DONnr}$	Reference rate for mineralization	$d^{-1}$	0.1
$\theta_{DONre}$	Temperature coefficient for mineralization of DONre	-	1.02
$\theta_{DONnr}$	Temperature coefficient for mineralization of DONnr	-	1.02
$K_{PhNutRege}$	Half saturation constant for phytoplankton regeneration	-	1.
$M_{nitri}$	Reference rate for nitrification		0.1
$\theta_{nitri}$	Temperature coefficient for nitrification	-	1.08
$K_{nitri}$	Half saturation constant for nitrification		2.0
$M_{det}$	reference rate for PON dissolution	$d^{-1}$	0.1
$\theta_{det}$	Temperature coefficient for decomposition	-	1.02
$M_{denit}$	reference rate for denitrification	$d^{-1}$	0.1
$\theta_{denit}$	Temperature coefficient for denitrification	-	1.046
$K_{denit}$	Half saturation constant for denitrification	$mgO_2.L^{-1}$	0.1
$\alpha_{Phy}$	N:C ratio for phyto	-	0.18
$\alpha_{Zo}$	N:C ratio for zoo		0.24

**Table 9.5-1: Values used in the standard model for the nitrogen variables**

## 9.6 Phosphorus

In the Mohid water quality module, phosphorus appears, like nitrogen, in an organic and an inorganic form.

The inorganic phosphorus is assumed to be available as orthophosphate ( $PO_4$ ) for uptake by phytoplankton.

The organic phosphorus is divided into particulate organic phosphorus (POP), dissolved organic phosphorus non refractory (DOPnr) and dissolved organic phosphorus refractory (DOPre). The rate equations of phosphorus are implemented in the same way as the nitrogen cycle, except that there is just one compartment of inorganic phosphorus.

### 9.6.1 Inorganic Phosphorus

Figure 9-10 represents the internal fluxes of inorganic phosphorus modeled by *Mohid's* water quality module.

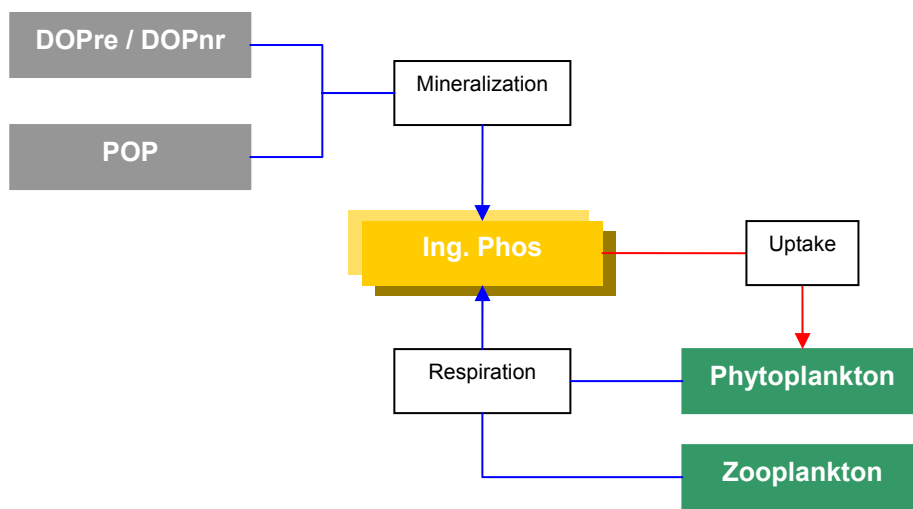


Figure 9-10: Internal Flux of Inorganic Phosphorus

The rate equation of inorganic phosphorus is given by:

$$\frac{\partial \Phi_{PO_4}}{\partial t} = (f_{in/Phy} ex_{Phy} - \mu_{PO_4}) \Phi_{Phy} + f_{in/Z} ex_z \Phi_z + \varphi_{Pre} \Phi_{DOPre} + \varphi_{Pnr} \Phi_{DOPnr} + f_{orgP/Phy} \varphi_{det} \Phi_{POP} \quad \text{Eq. 9-40}$$

The assimilation rate of  $PO_4$ ,  $\mu_{PO_4}$ , is given by:

$$\mu_{PO_4} = \alpha_{N:C} \mu_{Phy} \quad \text{Eq. 9-41}$$

$\alpha_{Phy}$  represents the Redfield ratio between N:P.

### 9.7 Oxygen

Figure 9-11 represents the internal fluxes of oxygen by *Mohid's* water quality module.

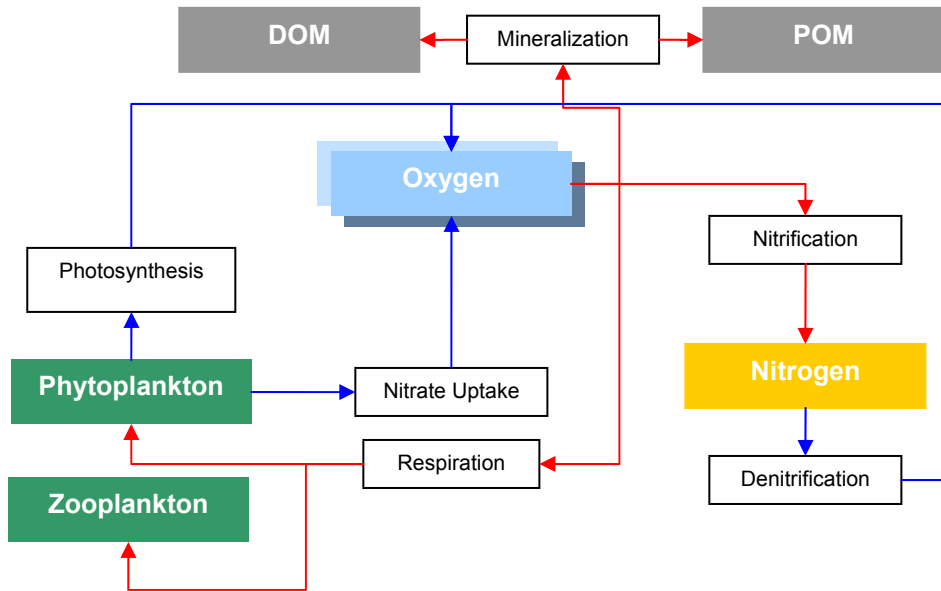


Figure 9-11: Internal Flux of Inorganic Oxygen

$$\frac{\partial \Phi_O}{\partial t} = (\mu_{phy} \alpha_{OC} + \mu_{phy} \alpha_{O:N} \alpha_{N:C} (1 - \beta_{NH4}) - \alpha_{OC} r_{Phy}) \Phi_{Phy} + r_z \alpha_{OC} \Phi_{Zoo} + \phi_{2N} \alpha_{N:O} \Phi_{NH4} - \left( \alpha_{MinO:N} \phi_{Nre} \Phi_{DONre} + \alpha_{MinO:N} \phi_{Nnr} \Phi_{DONnr} + \alpha_{MinO:N} \phi_{det} \Phi_{PON} + \alpha_{MinO:P} \phi_{Pre} \Phi_{DOPre} + \alpha_{MinO:P} \phi_{Pnr} \Phi_{DOPnr} + \alpha_{MinO:P} \phi_{det} \Phi_{POP} \right) \Phi_O \quad \text{Eq. 9-42}$$

$\alpha_{MinO:N}$  Mineralization oxygen/ nitrogen ratio

$\alpha_{MinO:P}$  Mineralization oxygen/ phosphorus ratio

## 10 The Surface Module

### 10.1 Introduction

The surface module stores the boundary conditions at the surface of the water column. These boundary conditions can be divided in two types of boundary condition. One type of boundary condition which are given directly by the user, usually meteorological data (wind velocity, air temperature, dew point, evaporation, cloud cover) or boundary conditions calculated automatically by the model from the meteorological data/conditions of the water column (wind stress, solar radiation, latent heat, infrared radiation, sensible heat, oxygen flux). The information flux between the surface module and other modules is shown in Figure 10-1.

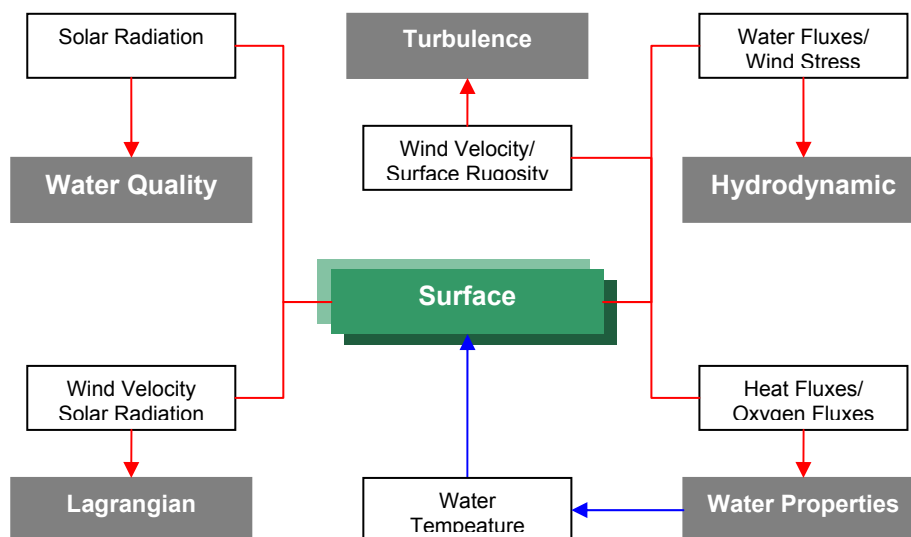


Figure 10-1: Information flux between the Surface Module and other modules

### 10.2 Wind

Wind stress is calculated according to a quadratic friction law:

$$\vec{\tau}_w = C_D \rho_a \vec{W} \left| \vec{W} \right| \quad \text{Eq. 10-1}$$

where  $C_D$  is a drag coefficient that is function of the wind speed,  $\rho_a$  is air

density and  $W$  is the wind speed at a height of 10 m over the sea surface.

The drag coefficient is computed according to Large and Pond [1981]:

$$C_D = 1.14e^{-3} \quad (W < 10 \text{ m/s}) \quad \text{Eq. 10-2}$$

$$C_D = 4.4e^{-4} + 6.5e^{-5} \frac{\vec{W}|\vec{W}|}{|\vec{W}|} \quad (10 \text{ m/s} < W < 26 \text{ m/s}) \quad \text{Eq. 10-3}$$

### 10.3 Heat fluxes

The heat fluxes at the surface can be separated into five distinctive fluxes: solar shortwave radiation, atmospheric long-wave radiation, water long-wave radiation, sensible heat flux and latent heat flux. These fluxes can be grouped into two ways: in (i) radiative fluxes (first three fluxes) and (ii) non-radiative fluxes (last two fluxes) or in (iii) fluxes independent of the water temperature (first two fluxes) and in (iv) fluxes dependent of the water temperature (last three fluxes).

#### 10.3.1 Solar radiation

Solar radiation is an important ecological parameter, and is often the key driving force in ecological processes (Brock, 1981). The solar radiation flux of short wavelength is compute by:

$$Q = Q_0 A_t (1 - 0.65 C_n^2) (1 - R_s) \quad \text{Eq. 10-4}$$

where  $Q_0$  is the solar radiation flux on top atmosphere ( $\text{Wm}^2$ ),  $A_t$  the coefficient for atmospheric transmission,  $C_n$  the cloud cover percentage and  $R_s$  stands for albedo (0.055). The solar radiation flux on top atmosphere can be expressed as:

$$Q_0 = \frac{I_0}{r^2} \text{senz} \quad \text{Eq. 10-5}$$

where  $I_0$  stands for the solar constant which is the energy received per unit time, at Earth's mean distance from the Sun, outside the atmosphere, a standard value, used is  $1353 \text{ Wm}^{-2}$  (Brock, 1981),  $r$  stands for the radius vector and  $z$  stands for the solar high.

### 10.3.1.1 Radius vector, $r$

During its revolution around the Sun, the Earth's distance varies with time of year by 3.0%, due to the Earth's eccentric orbit. This eccentricity influences in a minor way the amount of solar radiation impinging on the Earth's surface. The radius vector of Earth,  $r$ , expresses this ellipticity and can be calculated approximately from the following equation (Nicholls and Child, 1979 in Brock, 1981):

$$r = 1.0 + 0.017 \cos \left[ 2\pi \frac{(186 - d)}{365} \right] \quad \text{Eq. 10-6}$$

where  $d$  stands for Julian Day.

### 10.3.1.2 Solar High

Solar radiation at any location on Earth is influenced by the motions which the Earth makes in relation to the Sun. The Earth is tilted  $23.45^\circ$  from the plane of the Earth's orbit. The declination of Earth is the angular distance at solar noon between the Sun and the Equator, north-positive. Declination depends only on the day of the year, and will be opposite in the Southern Hemisphere. The declination is obtained precisely from ephemeris tables, but can be calculated close enough for all practical purposes from the equation given by Cooper (1969 in Brock, 1981):

$$D1(\text{declination}) = 23.45 \sin [2\pi(284 + d)/365] \quad \text{Eq. 10-7}$$

The other major motion is the daily rotation of the Earth around itself. The Earth moves  $15^\circ$  per hour and the sunset (or sunrise) hour-angle,  $W1$ , is the angle between the setting Sun and the south point. The value  $W1$  can be calculated if the latitude ( $L$ ) and declination are known:

$$W1 = \arccos (-\tan(L)\tan(D1)) \quad \text{Eq. 10-8}$$

In this equation, if  $L$  and  $D1$  are in degrees then  $W1$  will be given in degrees. From  $W1$ , the daylength in hours,  $L1$ , can be calculated from the equations:

$$\text{Sunrise} = 12 - \frac{1}{2} L1 \quad \text{Eq. 10-9}$$

$$\text{Sunset} = 12 + \frac{1}{2} L1 \quad \text{Eq. 10-10}$$

The hour-angle at any given time can be calculated from one of the following equations:

$$W2 = (T+12)*\pi/12, T < 12 \quad \text{Eq. 10-11}$$

$$W2 = (T-12)*\pi/12, T > 12$$

Where T is the time (h) from midnight.

The Zenith angle or angular elevation of the Sun above the horizon, Z, can be calculated if the declination, D1, the latitude, L1, and the hour-angle, W2, are known:

$$\text{Cos}(Z) = \sin(D1)\sin(L) + \cos(D1)\cos(L)\cos(W2) \quad \text{Eq. 10-12}$$

As a consequence of attenuation, radiation has two distinct directional properties when it reaches the ground.

### 10.3.1.3 Direct Radiation

Direct radiation arrives from the direction of the solar disc and includes a small component scattered directly forward. The term diffuse describes all other scattered radiation received from the blue sky and from clouds, either by reflection or by transmission. Direct radiation at the ground, measured at right angles to the beam, rarely exceeds 75% of the Solar Constant, i.e. about  $1030 \text{ Wm}^{-2}$ . The minimum loss of 25% is attributable to molecular scattering and to absorption in almost equal proportions. (Monteith and Unsworth, 1990)

### 10.3.1.4 Diffuse radiation

Beneath a clean, cloudless atmosphere, the absolute amount of diffuse radiation increases to a maximum somewhat less than  $200 \text{ Wm}^2$  when the zenith angle of the sun (Z) is less than  $50^\circ$  and the ratio of diffuse ( $Q_{\text{dif}}$ ) to total radiation ( $Q_0$ ) falls between 0.1 and 0.15. With increasing cloud amount also,  $Q_{\text{dif}}/Q_0$  increases and reaches unity when the sun is obscured by dense cloud: but the absolute level of  $Q_d$  is maximal when

cloud cover is about 50%.

The coefficient for atmospheric transmission is computed by the method followed by Rosati and Miyakoda (1988 in Portela, 1996):

$$A_t = A_{dir} + A_{dif} \quad \text{Eq. 10-13}$$

where  $A_{dir}$  is the direct fraction and  $A_{dif}$  is the diffuse fraction of solar radiation on top atmosphere that reaches the surface under a clear sky.

The direct fraction  $A_{dir}$  is given by:

$$A_{dir} = \tau^m \quad \text{Eq. 10-14}$$

where  $\tau = 0.74$  is atmospheric transmission coefficient for direct radiation and  $m$  the sectional mass, compute by the following expression:

$$m = 1 / \text{sen}(Z) \quad \text{Eq. 10-15}$$

where  $Z$  is the zenith angle in radians.

The diffuse fraction  $A_{dif}$  is given by:

$$A_{dif} = \frac{1 - A_a - A_{dir}}{2} \quad \text{Eq. 10-16}$$

where  $A_a = 0.09$  is the absorption coefficient due to water vapour and ozone.

### 10.3.2 Infrared radiation flux

The infrared radiation flux is computed in concordance with the *Stefan-Boltzman* law:

$$R_{br} = \varepsilon * \sigma * (273.15 + T_w)^4 \quad \text{Eq. 10-17}$$

where  $R_{br}$  represents the infrared radiation ( $W/m^2$ ),  $\varepsilon$  the emissivity of water (0.97),  $\sigma$  the Stefan-Boltzman constant ( $5.669 \cdot 10^{-08} W/m^2/K^4$ ) and  $T_w$  the water temperature.

### 10.3.3 Latent heat flux

The latent heat flux decreases the heat inside the water body. It represents the quantity of heat for the evaporation. The equation implemented in the model *Mohid* is known by the law of *Dalton*:

$$H_L = (19.0 + 0.95U_w^2) * (e_{s,w} - r_h * e_{s,a}) \quad \text{Eq. 10-18}$$

where  $H_L$  represents the latent heat flux (m/s),  $e_{s,w}$  water pressure of saturation (mmHg),  $r_h$  for the relative humidity and  $e_{s,a}$  the air pressure of saturation.

The model just considers the latent heat in the case of evaporation. In the inverse process, the model considers that the heat gain remains in the atmosphere.

### 10.3.4 Sensible heat flux

The difference between the air temperature and the water temperature is responsible for the sensible heat flux. The equation implemented in the model *Mohid* is known by the law of *Bowen*:

$$H_S = C_b * (19.0 + 0.95 * U_w^2) * (T_w - T_a) \quad \text{Eq. 10-19}$$

Where  $H_s$  represents the sensible heat flux ( $W/m^2$ ),  $C_b$  is the *Bowen's* coefficient (0.47mmHg/K),  $U_w$  the wind speed 10m above the surface of the water,  $T_w$  the water temperature and  $T_a$  the air temperature (K).

## 10.4 Gas flux

Actually just the oxygen flux is implemented. The formula used is indicated below:

$$K_L = \alpha U_w^\beta \quad \text{Eq. 10-20}$$

where  $K_L$  represents the velocity of the gas transfer (m/s),  $\alpha$  and  $\beta$  are coefficients depending in on the wind velocity,  $U_w$ :

$$\alpha = 0.2 \text{ if } W < 3.5 \text{ and } \alpha = 0.057 \text{ if } W > 3.5$$

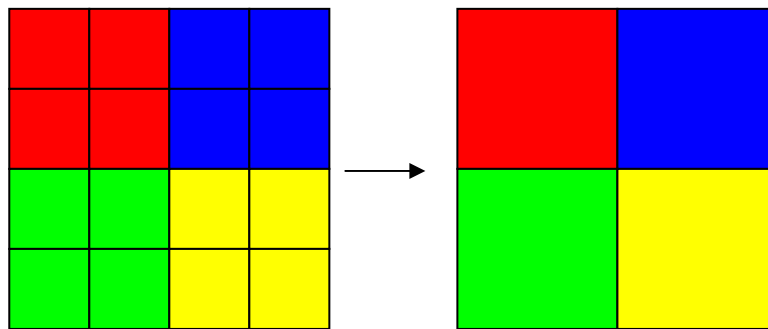
$\beta = 1.0$  if  $W < 3.5$  and  $\beta = 2.0$  if  $W > 3.5$

## 11 The Hydrodynamic File Module

### 11.1 Introduction

In this section the hydrodynamic file module of the model *Mohid* is described. This module can be seen as an auxiliary module, which permits the user of the model *Mohid* to integrate the hydrodynamic solution in space and in time and store this solution in a file. This file can later be used to simulate longer periods, like water quality simulation which needs simulation times for at least one year.

The special integration consists in the integration of several grid cells into one single cell. This grouping can be done for any quadratic group of grid cells, like 2x2 or 3x3. This grouping results in a drastic reduction of computing points. In the case of space integration 2x2, the resulting domain will just contain one fourth of grid points, each with the double size in the horizontal and vertical extension. Figure 11-1 shows a schematic representation of the space integration 2x2. The number of the overall grid points reduces from 16 to 4.



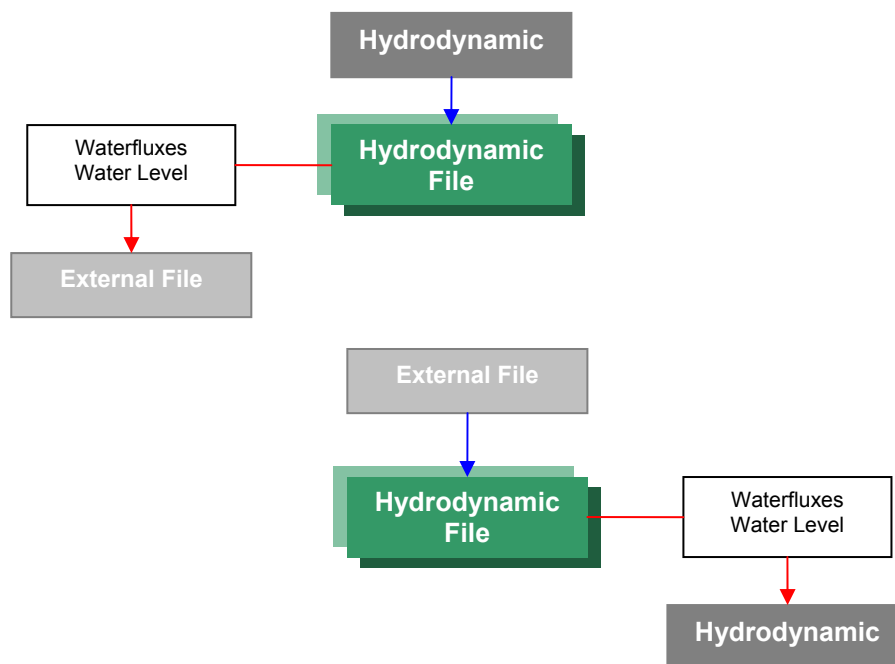
**Figure 11-1: Schematic representation of the space integration**

The time integration consists in the integration along several time steps of the hydrodynamic solution. The time integration can be directly connected to the space integration, once the larger grid spacing obtained by the space integration, allows the model to run with a larger time step. In the case of mass transport, the celerity which controls the stability of the model isn't the propagation of the pressure wave, but the maximum flow velocity.

In the resulting file, the hydrodynamic solution of the in time and in space integrated hydrodynamic solution is stored, which in posterior simulation can be used to obtain results more quickly. Regarding time, there are two different ways of storing this information: the information can be stored as a integrated solution with a given start date and a given end date, or as a solution which repeats itself (one tidal cycle).

The usage of the hydrodynamic file module has shown that the errors introduced in the integrated solution are usually small, and not significant for long-term water quality simulations.

The information flux of the hydrodynamic file module, relative to the other modules of *Mohid*, is shown in Figure 11-2. At the top the process of the file recording is shown and at the bottom the process of getting the solution from the file is shown.



**Figure 11-2: Information flux between the Hydrodynamic File Module and other modules**

## 11.2 Methodology

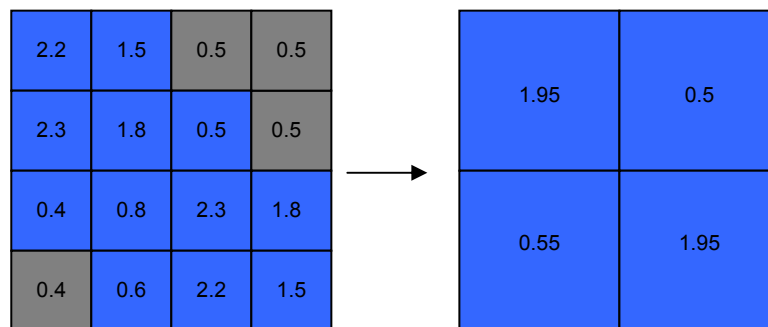
The space integration is divided into to steps:

1. Compute an integrated bathymetry, base on the bathymetry with a higher resolution
2. In each integration step, the sum of all water fluxes along a given face is kept in the output file. The average surface elevation is also kept in the output file.

### 11.2.1 Integration of the bathymetry

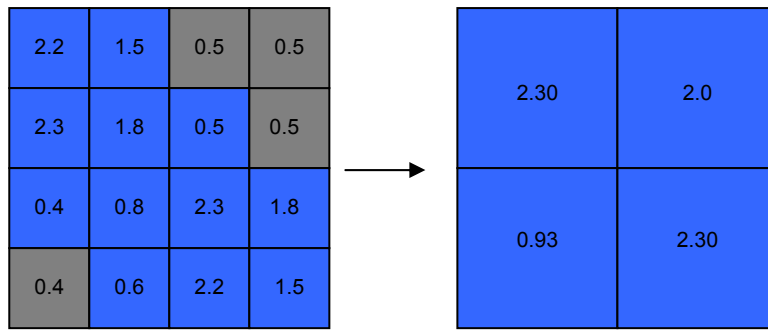
The integration of the bathymetry can be done in two different ways. In both ways, before the bathymetry is integrated, the land points with are filled with the minimum depth of the cells which are to be integrated.

The first way of the integration is designated as “Mean Integration”. This methodology calculates the average depth of the cells to be integrated, using this average as the depth for the new bathymetry. Figure 11-3 shows an integration of the bathymetry by the “Mean Integration” methodology. Water points are colored blue and land points are colored grey.



**Figure 11-3: Integration of the bathymetry using the “Mean Integration”**

The second way of integration is designated as “Maximum Integration”. This methodology uses the maximum depth of all cells, multiplied by the area of each cell and then divided by the total area of the water points.



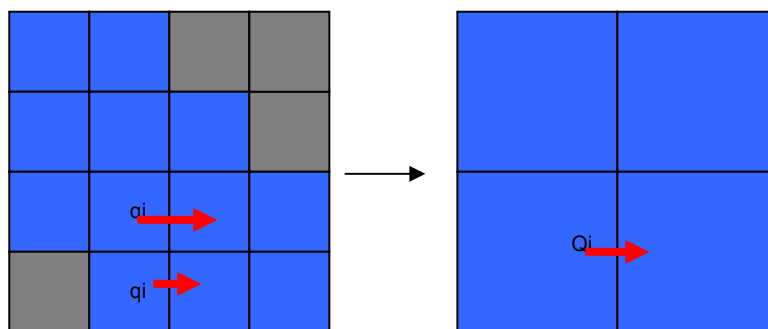
**Figure 11-4: Integration of the bathymetry using the “Maximum Integration”**

### 11.2.2 Integration of the water fluxes

The water flux between two cells is calculated from the average water flux between these cells. These fluxes can be integrated over several time steps.

$$Q_i = \sum_1^{nCells} \sum_1^{nSteps} q_i / nSteps \quad \text{Eq. 11-1}$$

Figure 11-5 shows a schematic representation of the integrated water fluxes over the integrated bathymetry.



**Figure 11-5: Schematic representation of the water flux integration**

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