TOMAWAC UserManual



Contents

1	TOMAWAC: an overview	6		
1.1	Introduction Generals	6		
1.2	Implementing TOMAWAC			
1.3	TOMAWAC general purposes	6		
2	Representing waves in TOMAWAC	. 8		
2.1	General definition of waves			
2.2	Plane monochromatic waves			
2.3	Random multidirectional waves	9		
2.4	Sea state directional power spectrum	10		
2.5	Directional spectrum of sea state variance	10		
2.6	Sea state directional spectrum of wave action			
2.7	Selecting the directional spectrum discretization variables	11		
3	Hypotheses and application domain of TOMAWAC	13		
3.1	Application domain of the model TOMAWAC	13		
3.2	Wave interactions with other physical factors	14		
3.3	The physical processes modelled in TOMAWAC	14		
4	Mathematical modelling procedures used by TOMAWAC	16		
4.1	Scope of sea state modelling	16		
4.2	Equations solved	18		
4.2.1 4.2.2	Equations solved in a Cartesian spatial coordinate system	18 20		
4.3	TOMAWAC source and sink terms	21		
4.3.1	Generals			
4.3.2 4.3.3	Wind input (term Q_{in})			
4.3.4	Bottom friction-induced dissipations (term O_{hf})			

4.3.5 4.3.6 4.3.7	Non-linear quadruplet interactions (term Qnl)				
4.3.8	Wave blocking effects (upper limit for spectrum or dissipation $Q_{ds,cur}$)				
4.3.9 4.3.10	Dissipation due to vegetation				
4.4	Surface rollers	41			
4.4.1	Theoretical background				
4.4.2					
5	Discretizations used in TOMAWAC	43			
5.1	Spatial discretization				
5.2	Spectro-angular discretization	43			
5.2.1	Frequency discretization				
5.2.2	Directional discretization:				
5.2.3	Spectro-angular grid:	45			
5.3	Temporal discretization	45			
6	Numerical methods used in TOMAWAC	47			
6.1	General solution algorithm	47			
6.2	Processing the advection step	48			
6.2.1	Representation of diffraction in the propagation step	49			
6.3	Processing the source term integration step	49			
6.3.1	Source term integration numerical scheme				
6.3.2	Monitoring the growth of the wave spectrum				
6.4	Processing the boundaries – Boundary conditions	52			
6.4.1 6.4.2	Spatial grid:				
6.5	A set of different models: BAJ modelisation.	53			
7	Inputs-outputs	54			
7.1	Preliminary remark	54			
7.2	The files	55			
7.2.1	The steering (or CAS) file	55			
7.2.2	The geometry file				
7.2.3 7.2.4	The boundary conditions file	57 57			
7.2.4	The tidal water level file	57			
7.2.5	The winds file	58			
7.2.7	The imposed spectra file and the file containing the associated coordinates				
7.2.8	The area definition file	58			
7.2.9	The previous computation file	59			
7.2.10	The global results file	59			
7.2.11	The 2D results file	59			
7.2.12	The time series file	۸n			

7.2.13 7.2.14 7.2.15 7.2.16 7.2.17 7.2.18	The punctual or spectrum results file The printout listing The User FORTRAN file The auxiliary files The dictionary file The libraries	63 63 63 64
7.3	Files standard	64
7.4	Bathymetry data	65
8	Controlling the simulation	66
8.1.1 8.1.2 8.1.3 8.1.4 8.1.5	General parameterisation Computation title Computation time Spectral discretisation Release Environment	66 66 67
8.2	Computation options	67
8.2.1 8.2.2 8.2.3 8.2.4 8.2.5 8.2.6 8.2.7 8.2.8 8.2.9	Co-ordinate system Finite depth Taking a stationary current into account Taking a wind into account Recovering a TELEMAC data item Taking the tide into account Waves-current interactions: direct coupling with Telemac (2D or 3D) flow sir lation Convection step Taking diffraction into account	67 68 68 69 69 mu- 69 71
8.3	Parameterising the source term integration step	72
8.3.1 8.3.2 8.3.3	Introduction	73
8.4	Prescribing the initial conditions	77
8.5 8.5.1 8.5.2 8.5.3 8.5.4	Prescribing the boundary conditions The boundary conditions file	81 81
8.6 8.6.1 8.6.2 8.6.3	Some useful subroutines Modification of bottom topography: USER_TOM_CORFON subroutine Modifying the co-ordinates: USER_CORRXY subroutine Operations on vectors: OVOV subroutine	82
9	API	84
Α	appendix	85
A .1	Output variables of TOMAWAC	85

		5
A.2	List of useful subroutines	87
A.3	Description of the formats being used	88
۹.4	File structure and processing	91
A.5	Parallel computing in TOMAWAC	93
	Bibliography	95

1. Tomawac: an overview

1.1 Introduction Generals

TOMAWAC is a scientific software which models the changes, both in the time and in the spatial domain, of the power spectrum of wind-driven waves and wave agitation for applications in the oceanic domain, in the intracontinental seas as well as in the coastal zone. The model uses the finite elements formalism for discretizing the sea domain; it is part of the TELEMAC system as developed by the EDF R&D's Laboratoire National d'Hydraulique et Environnement (LNHE). The acronym TOMAWAC being adopted for naming the software was derived from the following English denomination:

TELEMAC-based Operational Model Addressing Wave Action Computation (in French: Modèle opérationnel basé sur le système TELEMAC concernant le calcul de l'action d'onde pour les vagues)

TOMAWAC is one of the models making up the TELEMAC system [30], which addresses the various issues that are related to both free surface (either river- or sea-typed) and underground flows, as well as the associated physical processes: bed-load transport, water quality, etc.

1.2 Implementing TOMAWAC

TOMAWAC models the sea states by solving the balance equation of the action density directional spectrum. To serve that purpose, the model should reproduce the evolution of the action density directional spectrum at each node of a spatial computational grid.

In TOMAWAC the wave directional spectrum is split into a finite number of propagation frequencies f_i and directions θ_i . The balance equation of wave action density is solved for each component (f_i, θ_i) . The model is said to be a third generation model (e.g. like the WAM model [61] [36]), since it does not require any parameterization on the spectral or directional distribution of power (or action density). Each component of the action density spectrum changes in time under the effects of the software-modelled processes.

1.3 Tomawac general purposes

TOMAWAC can be used for three types of applications:

1. Wave climate forecasting a few days ahead, from wind field forecasts. This real time type of application is rather directed to weather-forecasting institutes such as Météo-

France, whose one mission consists of predicting continuously the weather developments and, as the case may be, publishing storm warnings.

- 2. **Hindcasting** of exceptional events having severely damaged maritime structures and for which field records are either incomplete or unavailable.
- 3. Study of wave climatology and maritime or coastal site features, through the application of various, medium or extreme, weather conditions in order to obtain the conditions necessary to carry out projects and studies (harbour constructions, morphodynamic coastal evolutions, ...).

During the development of the TOMAWAC model, the LNHE laboratory has been interested mainly on the last two types of applications. It considered also the possibility to carry out research activities focused on the following topics:

- wave-currents and wave-storm surge interactions, especially in those places where tide plays a significant role,
- coastal morphodynamics,
- probability of floods in coastal zone,
- coastal structure stability and coast protection,
- assimilation of wind or wave satellite data during computation...

2. Representing waves in TOMAWAC

2.1 General definition of waves

As stated in paragraph 1.1, the purpose of the TOMAWAC software consists of modelling the generation and the spatiotemporal evolution of waves at the surface of the seas or of the oceans. Then, the main physical process of interest is **the wave** or **the sea states**, these two terms being used interchangeably in this document.

The word waves, generally means all the wind driven free surface waves propagating at the surface of the ocean and the period of which (denoted as T) typically ranges from 2.5 to 25 s, or even, equivalently, whose frequency f=1/T ranges from 0.04 to 0.4 Hz.

The sea state may take various forms, depending on whether the sea is still and quiet or, on the contrary, in a stormy phase, whether the waves are being formed (the so-called wind sea) or, on the contrary, are coming from the ocean after travelling several hundreds or thousand kilometres (the so-called "swell").

2.2 Plane monochromatic waves

The most commonly used way to introduce wave modelling consists of considering simple sinusoidal waves (they are often called regular waves). It is a monochromatic (one period or frequency) and plane (one propagation direction) wave. The free surface elevation, which is denoted as h, depends on the position (x, y) of the point being considered in space, as well as on time t. It is written as:

$$\eta(x, y, t) = a\cos(k(x.\sin\theta + y.\cos\theta) - \omega.t + \phi)$$
 (2.1)

wherein:

a is the wave amplitude (in meters) and corresponds to the distance from the wave crest and the mean level at rest. The wave height, being measured from the crest to the trough of the wave, is used as well: H = 2a.

 ω is the wave frequency (in rad/s). The period (in seconds) $T=2\pi/\omega$ or the frequency f (in hertz) = $1/T=\omega/(2\pi)$ is used as well.

k is the wave number (in rad/m). The wavelength (in meters): $L = 2\pi/k$ is used as well. The wave number k is yielded by the free surface wave linear dispersion relation, according to frequency ω and depth d:

$$\omega^2 = g.k.tanh(k.d) \tag{2.2}$$

 θ is the wave propagation direction (in radians). Conventionally, this direction is measured herein clockwise with respect to Y axis.

 ϕ is the wave phase (in radians).

The energy per unit area of these progressive waves (which consists of kinetic energy and potential energy in halves) amounts to:

$$E = 1/2\rho g a^2 = 1/8\rho g H^2 \tag{2.3}$$

wherein:

g is the gravity acceleration $(g = 9.81m/s^2)$ ρ is the water density (in kg/m^3) ($\rho = 1025kg/m^3$ for seawater).

2.3 Random multidirectional waves

A first representation of waves at the surface of the ocean is possible through the sinusoidal expression being used in the preceding paragraph. When watching an actual sea state, however, not all the waves have the same features, whether it is in terms of height, period or propagation direction. As a matter of fact, the free surface wave energy is distributed over a range of frequencies (waves are then said to be irregular or random) and over a range of propagation directions (waves are then called multidirectional). Mathematically, that irregularity is expressed by writing that a real sea state results from the superposition of an infinite (or large) number of elementary sinusoidal components (i.e. monochromatic and uni-directional components).

Thus, a random multidirectional wave field can be modelled through a superposition method, considering M plane monochromatic components:

$$\eta(x, y, t) = \sum_{m=1}^{M} \eta_m(x, y, t) = \sum_{m=1}^{M} a_m \cos[km(x. \sin \theta_m + y. \cos \theta_m) - \omega_m t + \varphi_m]$$
 (2.4)

A major point in the above expression concerns the phase distribution jm of elementary wave components. The approach used in the TOMAWAC model assumes that these phases are randomly distributed over the $[0;2\pi]$ range with a uniform probability density. The various wave components are then independent, i.e. a linear or phase averaged representation is used.

With the linear representation featuring TOMAWAC and using the random phase hypothesis, the energy per unit area of random multidirectional waves can then be expressed as:

$$E = \sum_{m=1}^{M} \frac{1}{2} \rho g a_m^2 \tag{2.5}$$

It is noteworthy, however, that the distortions of shallow water wave profiles cannot be modelled with such a representation. This is because, as the water depth decreases, the non-linear processes linked to wave propagation and wave interactions with the sea bottom get some importance. The waves become steeper and dissymmetrical: they depart from a sinusoidal profile. A fine modelling of these non-linear effects involves non-linear wave theories (3rd- or 5th-order Stokes waves, cnoidal waves, ...) and/or so-called phase resolving propagation models modelling the evolution of each wave from a train, with a spatial discretization of 20-50 points per wavelength (Boussinesq, Serre equations, ...).

TOMAWAC is a phase averaged model: it is therefore *a priori* hardly suitable for modelling these non-linear effects when the wave profile can no longer be considered as the superposition of a number of independent sinusoidal components. In chapter 4, however, it will be explained how the non-linear effects can be processed and represented through source terms.

2.4 Sea state directional power spectrum

Real waves were introduced in the previous chapter as a discrete sum of elementary components. Actually, the power spectrum over both frequencies and propagation directions is a continuous function. The relevant variable for describing that sea state power spectrum is the **directional spectrum of wave energy** which is also known as wave **directional spectrum of energy** and will henceforth be denoted as E(f,q).

It is a function (in $Joule.Hz^{-1}.rad^{-1}$) that depends on:

- 1. wave frequency f (in Hertz), conventionally only positive (ranging from 0 to +8)
- 2. propagation direction θ , ranging within a 2π length interval.

Correspondence with the discrete case of the previous section is set considering the following equivalence:

$$\sum_{f}^{f+df} \sum_{\theta}^{\theta+d\theta} \frac{1}{2} \rho g a_m^2 = E(f,\theta) df d\theta$$
 (2.6)

In case of a wave propagation in a zero-current medium, a balance equation of the wave energy directional spectrum can be written taking into account some source and sink terms for energy generation or energy dissipation.

2.5 Directional spectrum of sea state variance

The preferred variable for sea state representation and modelling is rather the variance density **directional spectrum**.

This function, noted as F(f,q) and expressed in $m^2.Hz.rad^{-1}$ is simply derived from the directional spectrum of wave energy by the relation:

$$F(f,\theta) = E(f,\theta)/(\rho g) \tag{2.7}$$

Then, in particular, we have:

$$\sum_{f}^{f+df} \sum_{\theta}^{\theta+d\theta} \frac{1}{2} \rho g a_m^2 = E(f,\theta) df d\theta$$
 (2.8)

The relation linking the variance density directional spectrum and the free surface elevation is then written in the following pseudo-integral form:

$$\eta(x, y, t) = \int_{t=0}^{\infty} \int_{\theta=0}^{2\pi} \sqrt{2F(f, \theta)dfd\theta} \cos\left[k(x, \cos\theta + y, \sin\theta) - \omega t + \varphi\right]$$
 (2.9)

It should be reminded that the phases are randomly distributed in that expression over the range $[0;2\pi]$ with a uniform probability density. As regards the amplitude of each elementary component, it is related to the variance density directional spectrum by:

$$a_m = \sqrt{2F(f,\theta)dfd\theta} \tag{2.10}$$

The n-order (n = 0, 1, 2,...) moments mn of the variance density directional spectrum are defined as:

$$m_n = \int_{f=0}^{\infty} \int_{\theta=0}^{2\pi} f^n F(f, \theta) df d\theta$$
 (2.11)

Among these moments, the 0-order moment is equal to the variance of the free surface elevation:

$$<\eta^{2}> = \lim_{t_{o}\to\infty} \frac{1}{t_{o}} \int_{0}^{t_{o}} \eta^{2}(t)dt = m_{0} = \int_{t=0}^{\infty} \int_{\theta=0}^{2\pi} F(f,\theta)dfd\theta$$
 (2.12)

In particular, that moment m_0 affects the determination of the significant spectral wave height H_{mo} (equal to the significant height $H_{1/3}$ assuming that the wave heights are distributed according to a Rayleigh's law) by the relation:

$$H_{mo} = 4\sqrt{m_o} \tag{2.13}$$

The average frequencies f_{01} and f_{02} and $f_{-10}=f_e$ are also used and computed as follows:

$$f_{01} = \frac{m_1}{m_0} f_{02} = \sqrt{\frac{m_2}{m_0}} \text{ and } f_e = \frac{m_0}{m_{-1}}$$
 (2.14)

Further derived parameters can be computed from the variance density directional spectrum (see e.g. in [1].

2.6 Sea state directional spectrum of wave action

In the general case of wave propagation in an unsteady medium (sea currents and/or levels varying in time and space), the directional spectrum of the variance density is no longer kept and a new quantity should be introduced, namely the **directional spectrum of wave action**.

That quantity, noted as N(f,q), will remain constant (without considering the source and sink terms) even though the propagation medium is neither homogeneous nor steady [36] [63] [49] [9].

That action density spectrum is related to the directional spectrum of variance density by the relation:

$$N = F/s \tag{2.15}$$

wherein s denotes the relative or intrinsic angular frequency, i.e. the angular frequency being observed in a coordinate system moving at the velocity of current. Such a frequency is different from the absolute angular frequency w observed in a fixed system of coordinates. The two frequencies are linked by the Doppler effect relation in the presence of a current U:

$$\Omega(\mathbf{k}, \mathbf{x}, t) = \omega = \sigma + \mathbf{k}.\mathbf{U} \tag{2.16}$$

2.7 Selecting the directional spectrum discretization variables

The directional spectra of wave energy, variance or action shall generally be considered as functions depending on five variables:

- time t,
- the pair of coordinates proving the spatial position of the point being considered. In TOMAWAC, these coordinates can be expressed either in a Cartesian coordinate system (x, y) or in a spherical coordinate system (latitude, longitude) according to the dimension of the computational domain.
- the pair of variables applied for directional spectrum discretization, for which several solutions are theoretically possible:
 - (f_a, θ) = (absolute frequency; propagation direction)

- (f_r, θ) = (relative frequency; propagation direction)
- (k, θ) = (wave number; propagation direction)
- $(k_x, k_y) = (k. \sin \theta; k. \cos \theta) =$ (wave number vector)
- For the numerical resolution of equations, the model TOMAWAC uses the pair (f_r, θ) = (relative frequency; propagation direction)

The directional spectra output by TOMAWAC, however, are always expressed in (f_a, θ) . The equations solved by TOMAWAC are thoroughly reviewed in section 4.

3. Hypotheses and application domain of TOMAWAC

3.1 Application domain of the model TOMAWAC

TOMAWAC is designed to be applied from the ocean domain up to the coastal zone. The limits of the application range can be determined by the value of the relative depth d/L, wherein d denotes the water height (in metres) and L denotes the wave length (in metres) corresponding to the peak spectral frequency for irregular waves.

The application domain of TOMAWAC includes:

- the oceanic domain, characterized by large water depths, i.e. by relative water depths of over 0.5 The dominant physical processes are: wind driven waves, whitecapping dissipation and non-linear quadruplet interactions.
- the continental seas and the medium depths, characterized by a relative water depth ranging from 0.05 to 0.5. In addition to the above processes, the bottom friction, the shoaling (wave growth due to a bottom rise) and the effects of refraction due to the bathymetry and/or to the currents are to be taken into account.
- The coastal domain, including shoals or near-shore areas (relative water depth lower than 0.05). For these shallow water areas, such physical processes as bottom friction, bathymetric breaking, non-linear triad interactions between waves should be included. Furthermore, it could be useful to take into account the effects related to unsteady sea level and currents due to the tide and/or to the weather-dependent surges.

Through a so-called finite element spatial discretization, one computational grid may include mesh cells among which the ratio of the largest sizes to the smallest ones may reach or even exceed 100. That is why TOMAWAC can be applied to a sea domain that is featured by highly variable relative water depths; in particular, the coastal areas can be finely represented.

The application domain of TOMAWAC does not include the harbour areas and, more generally, all those cases in which the effects of reflection on structures and/or diffraction may not be ignored.

A first version of a diffraction model is available in TOMAWAC and is able to represent some diffraction effects. The model presents still some limits. It is highly recommended to use phase-resolving models when a detailed simulation of diffraction effects is required (e.g. harbour agitation).

3.2 Wave interactions with other physical factors

Several factors are involved in the wave physics and interact to various extents with the waves changing their characteristics. The following main factors should be mentioned:

- bathymetry and sea bottom geometry (bottom friction, refraction, surf-breaking, non-linear effects of interactions with the bottom, sand rippling...)
- atmospheric circulation (wind and pressure effects)
- tide pattern (variation of currents and water heights),
- three-dimensional oceanic circulation currents.
- over/underelevations caused by exceptional weather events, resulting in sea levels variations up to several meters (storm, surges).

The fine modelling of the interactions between these various physical factors and the waves is generally rather complex and several research projects are currently focused on it. Within the application domain as defined in the previous paragraph, TOMAWAC models the following interactions:

- wave-bathymetry interaction: the submarine relief data input into TOMAWAC are constant in time, but the sea level can change in time. In addition to the effects of the sea level variations in time, TOMAWAC allows to take into account refraction, shoaling, bottom friction and bathymetric breaking. TOMAWAC simulations can take into account some diffraction effects.
- wave-atmosphere interaction: this interaction is the driving phenomenon in the wave generation, takes part in energy dissipation processes (whitecapping, wave propagation against the wind...) and is involved in the energy transfer. To represent the unsteady behaviour of this interaction, TOMAWAC requires 10 m wind fields (specification of the couple of horizontal velocity components) with a time step matched to the weather conditions being modelled. These wind fields can be provided either by a meteorological model or from satellite measurements.
- wave-current interaction: the sea currents (as generated either by the tide or by oceanic circulations) may significantly affect the waves according to their intensity. They modify the refractive wave propagation direction, they reduce or increase the wave height according to their propagation direction in relation to the waves and may influence the wave periods if exhibiting a marked unsteady behaviour. In TOMAWAC, the current field is provided by the couple of horizontal components of its average (or depth-integrated) velocity at the nodes of the computational grid. TOMAWAC allows to model the frequency changes caused either by the Doppler effect or by the unsteady currents, as well as by a non-homogeneous current field.

3.3 The physical processes modelled in TOMAWAC

Those interactions being taken into account by TOMAWAC have been reviewed and a number of physical events or processes have been mentioned in the previous paragraph. These processes modify the total wave energy as well as the directional spectrum distribution of that energy (i.e. the shape of the directional spectrum of energy). So far, the numerical modelling of these various processes, although some of them are now very well known, is not yet mature and keep

on providing many investigation subjects. Considering the brief review of physical interactions given in the previous paragraph, the following physical processes are taken into account and digitally modelled in TOMAWAC:

—> Energy source/dissipation processes:

- <u>wind driven</u> interactions with atmosphere. Those interactions imply the modelling of the wind energy input into the waves. It is the prevailing source term for the wave energy directional spectrum. The way that spectrum evolves primarily depends on wind velocity, direction, time of action and fetch (distance over which the wind is active). It must be pointed out that the energy which is dissipated when the wind attenuates the waves is not taken into account in Tomawac.
- whitecapping dissipation or wave breaking, due to an excessive wave steepness during wave generation and propagation.
- <u>bottom friction</u>-induced dissipation, mainly occurring in shallow water (bottom grain size distribution, ripples, percolation...)
- <u>dissipation through bathymetric breaking</u>. As the waves come near the coast, they swell due to shoaling until they break when they become to steep.
- dissipation through wave blocking due to strong opposing currents.

—> Non-linear energy transfer conservative processes:

- <u>non-linear resonant quadruplet interactions</u>, which is the exchange process prevailing at great depths.
- non-linear triad interactions, which become the prevailing process at small depths.

---> Wave propagation-related processes:

- wave <u>propagation</u> due to the wave group velocity and, in case, to the velocity of the medium in which it propagates (sea currents).
- <u>depth-induced refraction</u> which, at small depths, modifies the directions of the wave-ray and then implies an energy transfer over the propagation directions.
- <u>shoaling</u>: wave height variation process as the water depth decreases, due to the reduced wavelength and variation of energy propagation velocity.
- current-induced refraction which also causes a deviation of the wave-ray and an energy transfer over the propagation directions.
- interactions with unsteady <u>currents</u>, inducing frequency transfers (e.g. as regards tidal seas).
- <u>diffraction</u> by a coastal structure (breakwater, pier, etc...) or a shoal, resulting in an energy transfer towards the shadow areas beyond the obstacles blocking the wave propagation. The current version of the diffraction model implemented in TOMAWAC is able to represent qualitatively some diffraction effects.

These various processes are numerically modelled as presented in Part 4. It should be remembered that, due to the hypothesis adopted in paragraph 3.1 about the TOMAWAC application domain, the <u>reflection</u> (partial or total) from a structure or a pronounced depth irregularity is not addressed by the model.

4. Mathematical modelling procedures used by Tomawac

4.1 Scope of sea state modelling

The directional spectrum of wave action density, as defined in paragraph 2.6, is considered as a function of five variables:

$$N(\boldsymbol{x},\boldsymbol{k},t) = N(x,y,k_x,k_y,t)$$

using, as discretization variables:

- 1. the position vector $\mathbf{x} = (x, y)$ for spatial location in a Cartesian coordinate system
- 2. the wave number vector $\mathbf{k} = (k_x, k_y) = (k.sin\theta, k.cos\theta)$ for directional spectrum discretization, θ denoting the wave propagation direction (direction in which the waves travel).
- 3. the time t.

Under the hypotheses made on the wave representation (see in paragraph 2.6) as well as on the model application domain and the modelled physical processes (see in paragraph 3.3), an equation of evolution of the directional spectrum of wave action can be written in the following form (see in [63] [49] [9] for a detailed demonstration of the way that equation is arranged):

$$\frac{\partial N}{\partial t} + \frac{\partial (\dot{x}N)}{\partial x} + \frac{\partial (\dot{y}N)}{\partial y} + \frac{\partial (\dot{k}_xN)}{\partial k_x} + \frac{\partial (\dot{k}_yN)}{\partial k_y} = Q(k_x, k_y, x, y, t)$$
(4.1)

The equation expresses that, in the general case of waves propagating in a non-homogeneous, unsteady environment (currents and/or sea levels varying in time and space), the wave action is preserved to within the source and sink terms (designated by the term Q).

The following notation is also used in (4.1):

$$\dot{g} = \frac{dg}{dt} = \frac{\partial g}{\partial t} + \frac{\partial x}{\partial t} \frac{\partial g}{\partial x} + \frac{\partial y}{\partial t} \frac{\partial g}{\partial y}$$

In that form (conservative writing in the form of a flux), equation (4.1) can be transposed to other coordinate systems and, for instance, $(k, \theta), (f_a, \theta)$ or else (f_r, θ) can be used for the discretization of directional spectrum [36] [58]

Working in (x, y, k_x, k_y) , however, makes it possible to remain in the canonical coordinate system and to write, for the propagation equations (also named Hamilton's equations):

$$\begin{cases} \dot{x} = \frac{\partial \Omega}{\partial k_x} & \text{and} \quad \dot{y} = \frac{\partial \Omega}{\partial k_y} \\ \dot{k}_x = -\frac{\partial \Omega}{\partial x} & \text{and} \quad \dot{k}_y = -\frac{\partial \Omega}{\partial y} \end{cases}$$
(4.2)

wherein W results from the Doppler relation applied to the wave dispersion relation for the general case with current:

$$\Omega(\mathbf{k}, \mathbf{x}, t) = \omega = \sigma + \mathbf{k}.\mathbf{U} \tag{4.3}$$

wherein: w is the absolute angular frequency observed in a fixed coordinate system.fa = w/(2p) is named absolute frequency.

U denotes the current velocity (depth-integrated).

 σ denotes the intrinsic or relative angular frequency, which is observed in a coordinate system moving at the velocity U. It is given by the dispersion relation in the zero-current case:

$$\sigma^2 = g.k.tanh(k.d) \tag{4.4}$$

 $f_r = \sigma/(2\pi)$ is named intrinsic or relative wave frequency. d denotes the water height. Through the Hamilton's equations (4.2), it can be demonstrated that we have:

$$\frac{\partial \dot{x}}{\partial x} + \frac{\partial \dot{y}}{\partial y} + \frac{\partial \dot{k}_x}{\partial k_x} + \frac{\partial \dot{k}_y}{\partial k_y} = 0 \tag{4.5}$$

or $div(\mathbf{V}) = 0$ when defining: $\mathbf{V} = (\dot{x}, \dot{y}, \dot{k}_x, \dot{k}_y)$

The evolution equation (4.1) can then alternatively be written in the following form (the so-called transport form):

$$\frac{\partial N}{\partial t} + \dot{x}\frac{\partial N}{\partial x} + \dot{y}\frac{\partial N}{\partial y} + \dot{k}_x\frac{\partial N}{\partial k_x} + \dot{k}_y\frac{\partial N}{\partial k_y} = Q(k_x, k_y, x, y, t)$$

$$\frac{\partial N}{\partial t} + \mathbf{V}.grad_{\mathbf{x},\mathbf{k}}(N) = Q$$
(4.6)

The transfer rates are given by the linear wave theory [11] [36] [43] [58]

$$\dot{x} = C_g \frac{k_x}{k} + U_x
\dot{y} = C_g \frac{k_y}{k} + U_y
\dot{k}_x = -\frac{\partial \sigma}{\partial d} \frac{\partial d}{\partial x} - \mathbf{k} \cdot \frac{\partial \mathbf{U}}{\partial x}
\dot{k}_y = -\frac{\partial \sigma}{\partial d} \frac{\partial d}{\partial y} - \mathbf{k} \cdot \frac{\partial \mathbf{U}}{\partial y}$$
(4.7)

 C_g is the relative (or intrinsic) group velocity of waves, i.e. as is observed in a coordinate system moving at the velocity of the current:

$$C_g = \frac{\partial \sigma}{\partial k} = n \frac{\sigma}{k} \text{ with } n = \frac{1}{2} \left(1 + \frac{2kd}{\sinh(2kd)} \right)$$
 (4.8)

The relative (or intrinsic) phase velocity C of waves is also introduced: $C = \frac{\sigma}{L}$

The sea state spectral modelling will then consist of solving the evolution equations (4.1) or (4.6), using the kinematic equations (4.7).

The transport equation formulation (4.6) has been adopted in TOMAWAC, since it is closely related to other equations applied in hydraulics, which have already been treated at the LNHE and for which methods and a know-how have been developed long ago.

As regards the discretization variables being used in TOMAWAC, we have already mentioned in paragraph 2.7 that:

- spatial discretization can be based either on a Cartesian coordinate system in (x, y) or on a spherical coordinate system at the Earth's surface in (λ, ϕ) = (longitude, latitude).
- Discretization of angular spectrum uses the pair (f_r, θ) = (relative frequency; propagation direction).

The following conventions are adopted for writing the equations:

- the x-axis (in the Cartesian coordinate system) or the l-axis of longitudes (in the spherical coordinate system) is assumed to be horizontal, directed to the right, whereas the y-axis (in the Cartesian coordinate system) or the j-axis of latitudes (in the spherical coordinate system) is assumed to be vertical, upwardly directed. Then, in spherical coordinates, the vertical axis points at the north, whereas the horizontal axis points to the East.
- In either case, the wave propagation directions q are defined with respect to the vertical axis in the clockwise direction.

These conventions are illustrated below in Figure 4.1 Those equations that correspond to the two spatial discretizations options are developed in the next paragraphs.

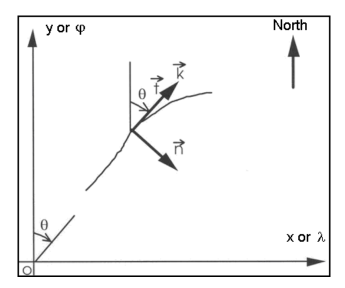


Figure 4.1: definition of location conventions as used in TOMAWAC

4.2 Equations solved

4.2.1 Equations solved in a Cartesian spatial coordinate system

By switching the variable from (x, y, k_x, k_y) to (x, y, f_r, q) , it can be shown that the following relation exists for the directional spectrum of wave action as expressed in both coordinate systems:

$$N(x, y, k_x, k_y, t) = \frac{CC_g}{2\pi\sigma} \tilde{N}(x, y, f_r, \theta, t) = \tilde{B}.\tilde{F}(x, y, f_r, \theta, t)$$
(4.9)

putting:
$$\tilde{B} = \frac{C C_g}{2\pi\sigma^2} = \frac{C_g}{(2\pi)^2 k f_r}$$
 (4.10)

The evolution equation (4.6) is then written as:

$$\frac{\partial(\tilde{B}\tilde{F})}{\partial t} + \dot{x}\frac{\partial(\tilde{B}\tilde{F})}{\partial x} + \dot{y}\frac{\partial(\tilde{B}\tilde{F})}{\partial y} + \dot{\theta}\frac{\partial(\tilde{B}\tilde{F})}{\partial \theta} + \dot{f}_r\frac{\partial(\tilde{B}\tilde{F})}{\partial f_r} = \tilde{B}.\tilde{Q}(x,y,\theta,f_r,t)$$
(4.11)

with the following transfer rates, as computed from the linear wave theory:

$$\dot{x} = C_g \cdot \sin \theta + U_x
\dot{y} = C_g \cdot \cos \theta + U_y
\dot{\theta} = -\frac{1}{k} \frac{\partial \sigma}{\partial d} \tilde{G}_n(d) - \frac{\mathbf{k}}{k} \cdot \tilde{G}_n(\mathbf{U})
\dot{f}_r = \frac{1}{2\pi} \left[\frac{\partial \sigma}{\partial d} \left(\frac{\partial d}{\partial t} + \mathbf{U} \cdot \nabla d \right) - C_g \mathbf{k} \cdot \tilde{G}_t(\mathbf{U}) \right]$$
(4.12)

The operators \tilde{G}_n and \tilde{G}_t refer to the computation of a function gradient in directions that are respectively normal and tangential to the characteristic curve with the direction q:

$$\tilde{G}_{n}(g) = \mathbf{n} \cdot \nabla g = \cos \theta \frac{\partial g}{\partial x} - \sin \theta \frac{\partial g}{\partial y}
\tilde{G}_{t}(g) = \mathbf{t} \cdot \nabla g = \sin \theta \frac{\partial g}{\partial x} + \cos \theta \frac{\partial g}{\partial y}$$
(4.13)

Besides, using the dispersion relation 4.4, it can be demonstrated that:

$$\frac{\partial \sigma}{\partial d} = \frac{\sigma k}{\sinh(2kd)} \tag{4.14}$$

The spatial transfer rates \dot{x} and \dot{y} (equations 4.12) model the spatial wave propagation and the shoaling. The directional transfer rate $\dot{\theta}$ models the refraction-induced change of wave propagation direction. Refraction is generated by the spatial variations of those properties of the environment in which the waves propagate and can result either from a bathymetric variation (first term) or from current gradients (second term). The relative frequency transfer rate \dot{f}_r models the relative frequency changes resulting from sea level variations both in space and time and/or from current variations in space.

It is noteworthy that this last term is zero in the case of zero-current and of no variation of sea level in time: the advection equation is then reduced to a three-dimensional equation.

Lastly, as regards the source terms, it should be mentioned that changing the coordinate system and using the factor \tilde{B} allows to switch from the term Q to a term \tilde{Q} that is directly expressed in terms of the directional variance spectrum with a variance $\tilde{F}(f_r,\theta)$. The content of that term is explained in paragraph 4.3

Consideration of diffraction

If diffraction is taken into account, a new wave number K is computed, whose modulus is the sum of the wave number k calculated by the linear wave theory plus a term that takes into account the effects of the wave diffraction. This modified wave number can be used to represent the effect of diffraction in a phase-averaged spectral model, as proposed by Holthujisen (2003). Two different mathematical formulations of diffraction are considered:

• the equation of Berkhoff (1972), also called Mild-Slope Equation (MSE), which gives the velocity potential of a monochromatic and unidirectional wave propagating over a slowly-varying bathymetry and describes the combined effects of diffraction and refraction;

• a modified form of the MSE, the Revised Mild Slope Equation (RMSE) proposed by Porter (2003).

In the presence of diffraction the effective wave number *K* is defined as:

$$|\mathbf{K}|^2 = k^2 (1 + \delta) \tag{4.15}$$

where δ is a diffraction parameter defined as when referring to the **Mild-Slope Equation** [6]

$$\delta_{MSE} = \frac{\nabla \cdot (CC_g \nabla a)}{k^2 CC_o a} \tag{4.16}$$

when referring to the **Revised Mild-Slope Equation** [51] (with $A = ak\sqrt{CC_g}$)

$$\delta_{RMSE} = \frac{\nabla \cdot \left(k^{-2}\nabla A\right)}{A} \tag{4.17}$$

In the presence of diffraction, the energy propagation speed in geographic space C_{gd} is [31]:

$$C_{gd} = \frac{K}{k}C_g \tag{4.18}$$

Under the hypothesis of zero current and of no varying water levels (i.e. \dot{f}_r =0) and inserting (4.15) in (4.12), the transfer rates in presence of diffraction are:

$$\dot{x} = \sqrt{(1+\delta)}C_g.\sin\theta\tag{4.19}$$

$$\dot{y} = \sqrt{(1+\delta)}C_g \cdot \cos\theta \tag{4.20}$$

$$\dot{\theta} = -\frac{(1+\delta)^{1/2}}{k} \frac{\partial \sigma}{\partial d} \tilde{G}_n(d) - \frac{C_g}{2(1+\delta)^{1/2}} \tilde{G}_n(\delta)$$
(4.21)

The above terms (4.19) to (4.21) have been implemented in TOMAWAC, in both the MSE and RMSE formulations, to represent the effect due to diffraction.

4.2.2 Equations solved in a spherical spatial coordinate system

By switching the variables from (x, y, k_x, k_y) to $(\lambda, \phi, f_r, \theta)$, it can be shown that the following relation exists for the directional spectrum of wave action as expressed in both coordinate systems:

$$N(x, y, k_x, k_y, t) = \frac{CC_g}{2\pi\sigma R^2 \cos\phi} \hat{N}(\lambda, \phi, f_r, \theta, t) = \hat{B}.\hat{F}(\lambda, \phi, f_r, \theta, t)$$
(4.22)

putting:
$$\hat{B} = \frac{C C_g}{2\pi\sigma^2 R^2 \cos\phi} = \frac{C_g}{(2\pi)^2 k f_r R^2 \cos\phi}$$
 (4.23)

R denotes the Earth's radius (R = 6400 km) and, once more, λ and ϕ are respectively the longitude and the latitude of the point being considered.

The evolution equation (4.6) is then written as:

$$\frac{\partial(\hat{B}\hat{F})}{\partial t} + \dot{\lambda}\frac{\partial(\hat{B}\hat{F})}{\partial\lambda} + \dot{\phi}\frac{\partial(\hat{B}\hat{F})}{\partial\phi} + \dot{\theta}\frac{\partial(\hat{B}\hat{F})}{\partial\theta} + \dot{f}_r\frac{\partial(\hat{B}\hat{F})}{\partial f_r} = \hat{B}.\hat{Q}(\lambda,\phi,\theta,f_r,t)$$
(4.24)

with the following transfer rates:

$$\dot{\lambda} = \frac{1}{R\cos\phi} \left(C_g \cdot \sin\theta + U_{\lambda} \right)
\dot{\phi} = \frac{1}{R} \left(C_g \cdot \cos\theta + U_{\phi} \right)
\dot{\theta} = \frac{1}{R} \left[C_g \sin\theta \tan\phi - \frac{1}{k} \frac{\partial\sigma}{\partial d} \hat{G}_n(d) - \frac{\mathbf{k}}{k} \cdot \hat{G}_n(\mathbf{U}) \right]
\dot{f}_r = \frac{1}{2\pi R} \left[\frac{\partial\sigma}{\partial d} \left(\frac{\partial d}{\partial t} + \frac{U_{\lambda}}{\cos\phi} \frac{\partial d}{\partial \lambda} + U_{\phi} \frac{\partial d}{\partial \phi} \right) - C_g \quad \mathbf{k} \cdot \hat{G}_t(\mathbf{U}) \right]$$
(4.25)

As in the previous case, the operators \hat{G}_n and \hat{G}_t refer to the computation of a function gradient in directions that are respectively normal and tangential to the characteristic curve with the direction q:

$$\hat{G}_n(g) = \frac{\cos \theta}{\cos \phi} \frac{\partial g}{\partial \lambda} - \sin \theta \frac{\partial g}{\partial \phi}$$

$$\hat{G}_t(g) = \frac{\sin \theta}{\cos \phi} \frac{\partial g}{\partial \lambda} + \cos \theta \frac{\partial g}{\partial \phi}$$
(4.26)

As previously, the spatial transfer rates λ and $\dot{\phi}$ (equations 4.25) model the wave propagation in space and the shoaling. In that coordinate system, the directional transfer rate $\dot{\theta}$ has an additional term (the first term) compared to the case in Cartesian coordinates. That term results from the propagation in spherical coordinates, in such a way that waves are located with respect to the North change during the propagation over a large circle arc at the Earth's surface [61] [36]. Both second and third terms $\dot{\theta}$ model the refraction caused respectively by bathymetry and currents. The relative frequency transfer rate \dot{f}_r models the changes of relative frequency resulting from variations of the sea level or of the current in both space and time. It is noteworthy that this last term is zero in the case of zero current and of no variation of the sea level in time: the advection equation is then reduced to a three-dimensional equation.

4.3 TOMAWAC source and sink terms

4.3.1 Generals

The source and sink terms that compose \tilde{Q} and \hat{Q} in the right-hand members of evolution equations (4.11) and (4.24) of directional spectrum of wave action gather the contributions from the physical processes listed in paragraph 3.3. for the application domain of TOMAWAC:

$$Q = Q_{in} + Q_{ds} + Q_{nl} + Q_{bf} + Q_{br} + Q_{tr} + Q_{ds,cur} + Q_{veg} + Q_{porous}$$
(4.27)

wherein

- Q_{in} : wind-driven wave generation
- Q_{ds} : whitecapping-induced energy dissipation
- Q_{nl} : non-linear quadruplet interactions
- Q_{bf} : bottom friction-induced energy dissipation
- Q_{br} : bathymetric breaking-induced energy dissipation
- Q_{tr} : non-linear triad interactions
- $Q_{ds,cur}$: enhanced breaking dissipation of waves on a current

- Q_{veg} : dissipation due to vegetation.
- Q_{porous} : dissipation due to porous media.

These source and sink terms are numerically modelled and parameterized as detailed in the next paragraphs. For most of these processes, several models or formulations are proposed and available in TOMAWAC.

4.3.2 Wind input (term Q_{in})

Three wind generation models are available in TOMAWAC. The model to be activated is selected through the keyword *WIND GENERATION* in the steering file, which can take four values, namely:

- 0 no wind input (default value)
- 1 Janssen's model [33] [34] (WAM cycle 4)
- 2 Snyder et al. model [55]
- 3 Yan's model [65]

Beside those exponential growth-type wind generation models, a linear growth model is also available in TOMAWAC, which has been proposed by Cavaleri & Malanotte-Rizzoli [10] (see last paragraph in this section The model can be activated through the keyword *LINEAR WAVE GROWTH*, and can be used together with one of the three above mentioned models. Its main feature is that it permits to start a wave simulation from a nil wave spectrum (whereas the three above mentioned models need some initial energy level for the wave spectrum to grow under wind action).

Option 1 for wind input: Janssen's model

With that option, the model implemented for the wind input term is based upon the Janssen's works [33] [34]; Janssen proposed a quasi-linear theory for modelling the ocean/atmosphere interactions. The linear growth term is ignored and only an exponential energy growth is taken into account, following Miles' results [45].

A quasi-linear source term is obtained as a function of the directional variance spectrum:

$$Q_{in} = \sigma.\varepsilon.\beta. \left(\left[\frac{u_*}{C} + z_{\alpha} \right] \max \left[\cos(\theta - \theta_w); 0 \right] \right)^2 F$$
 (4.28)

with the following notations:

 $\varepsilon = \rho_{air}/\rho_{water}$ is the ratio of air and water specific gravities ($\varepsilon = 1.25 \ 10^{-3}$).

 $C = \sigma/k$ is the wave phase velocity

 θ_w is the local wind direction (direction in which it blows)

 u_* is the friction velocity, being linked to the surface stress τ_s by the following relation:

$$u_* = \sqrt{\frac{\tau_s}{\rho_{air}}} \tag{4.29}$$

 z_{α} is a constant allowing to offset the growth curve.

The operator 'max' in the source term expression limits the wave generation for the propagation directions included within a $\pm 90^{\circ}$ angular sector with respect to the local wind direction qw. For the wave directions making an angle in excess of 90° with respect to the wind direction θ_w , the wind input term is void.

In the Janssen's model [34], the Miles' parameter β is a function of two non-dimensional variables:

- the wave age: $A = \frac{u_*}{C}$
- the wind profile parameter: $\Omega = \frac{g.z_0}{u_*^2}$

It is written as
$$\beta = \frac{\beta_m}{\kappa^2} \mu \ln^4 \mu$$
 (4.30)

where κ is the Von Karman's constant

 β_m denotes a coefficient set to 1.2 by Janssen [34]

z₀ denotes the roughness length

 μ denotes the non-dimensional critical height:

$$\mu = \min \left[\frac{g \cdot z_0}{C^2} \exp \left(\frac{\kappa}{\left[\frac{u_*}{C} + z_{\alpha} \right] \cos(\theta - \theta_w)} \right); 1 \right]$$

$$= \min \left[\Omega \cdot A^2 \exp \left(\frac{\kappa}{\left[A + z_{\alpha} \right] \cos(\theta - \theta_w)} \right); 1 \right]$$
(4.31)

The Janssen's model [33] [34] is characterized by the method it uses for computing u* and z0. The surface stress τ_s is a function depending, on the one hand, on the wind velocity U_{10} at 10 m and, on the other hand, on the sea state roughness through the wave stress τ_w . It is obtained by solving the following system of equations:

$$U_{10} = \frac{u_*}{\kappa} \ln \left(\frac{10 + z_0 + \tilde{z}_0}{z_0} \right) \approx \frac{u_*}{\kappa} \ln \left(\frac{10}{z_0} \right)$$

$$\tag{4.32}$$

$$z_0 = \frac{\tilde{z}_0}{\sqrt{1 - \tau_w/\tau_s}} \text{ avec } \tilde{z}_0 = \alpha \frac{u_*^2}{g} \text{ et } u_* = \sqrt{\frac{\tau_s}{\rho_{air}}}$$

$$(4.33)$$

The solution of the system of equations through a Newton-Raphson's iterative method yields the surface stress τ_s , the friction velocity u_* and the roughness length z_0 .

The initial value of friction velocity u_* being used in the iterative algorithm is obtained considering a constant drag coefficient:

 $u_* \sqrt{C_D} U_{10}$ where: $C_D = 1.2875 \ 10^{-3}$ by default.

The wave stress τ_w itself is computed from the variance spectrum F (via the source term Qin) using the following relation:

$$\tau_{w} = \left| \iint \rho_{water} \, \sigma \, Q_{in}(f_{r}, \theta) \, (\cos \theta, \sin \theta) \, df_{r} \, d\theta \right| \tag{4.34}$$

That integral is numerically computed over the discretized portion of the spectrum and a parametrization, based upon a decrement of variance in f^{-n} , is used for the high frequencies portion of the spectrum.

In fact, that source term has eight parameters, namely:

- coefficient β_m (corresponding to the keyword WIND GENERATION COEFFICIENT in the steering file). Its default value is taken as 1.2, in accordance with the Janssen's proposal [34] and the value adopted in the model WAM-Cycle 4.
- air specific gravity ρ_{air} (corresponding to the keyword *AIR DENSITY* in the steering file. Its default value is taken as 1.225 kg/m³.
- water specific gravity ρ_{water} (corresponding to the keyword WATER DENSITY in the steering file). Its default value is taken as $1{,}000kg/m^3$.

- constant α (corresponding to the keyword *CHARNOCK CONSTANT* in the steering file). Its default value is taken as 0.01, in accordance with the Janssen's proposal and the standard value adopted in the model WAM-Cycle 4.
- constant κ (corresponding to the keyword *VON KARMAN CONSTANT* in the steering file). Its default value is taken as 0.41, i.e. the typical value.
- initial drag coefficient C_D (corresponding to the keyword WIND DRAG COEFFICIENT in the steering file). This drag coefficient is provided for initializing the iterative computation of friction velocity u_* . Its default value is taken as 1.2875 10^{-3} .
- offset constant z_{α} (corresponding to the keyword *SHIFT GROWING CURVE DUE TO WIND* in the steering file). Its default value is taken as 0.011, in accordance with the value adopted in the model WAM-Cycle 4.
- elevation at which the wind is recorded (corresponding to the keyword WIND MEASURE-MENTS LEVEL in the steering file). Its default value is taken as 10 m: its corresponds to the typical value and to the value being adopted in the above explanations.

Option 2 for wind input: Snyder et al. model

In that option, the model implemented for the wind input term is based upon the works conducted by Snyder *et al.* [55], as amended by Komen *et al.* [35] to take into account the friction velocity u* instead of the wind velocity at 5 m. It corresponds to the formulation being used in the cycle 3 release of WAM model. The formulation is simpler than the Janssen's theory which Option 1 is based upon (see in preceding paragraph):

As in Option 1, the linear growth term is ignored and only an exponential energy growth is taken into account, following the Miles' results [45]:

$$Q_{in} = \beta F \text{ where: } \beta = \max \left[0; 0.25 \frac{\rho_{air}}{\rho_{water}} \left(28 \frac{u_*}{C} \cos(\theta - \theta_w) - 1 \right) \right] \sigma \tag{4.35}$$

The shear velocity value u_* used is obtained considering a drag coefficient linearly depending on the wind velocity:

$$u_* = \sqrt{C_D}U_{10}$$
 where : $C_D = 6.5 \ 10^{-5} + 8 \ 10^{-4}U_{10}$ if $U_{10} > 7.5m/s$.
 $C_D = 1.2875 \ 10^{-3}$ if $U_{10} < 7.5m/s$. (4.36)

That source term only uses two parameters, namely:

- air density ρ_{air} (corresponding to the keyword AIR DENSITY in the steering file. Its default value is taken as 1.225 kg/m^3 .
- water density ρ_{water} (corresponding to the keyword WATER DENSITY in the steering file). Its default value is taken as 1,000 kg/m^3 .

Option 3 for wind input: Yan's model

The Yan's model [65] consists of a combination of u_*/C and $(u_*/C)^2$ terms. It is valid over a wide range of frequencies and wind speeds:

$$Q_{in} = \beta F$$
with $\beta = \left[D \left(\frac{u_*}{C} \right)^2 \cos(\theta - \theta_w) + E \frac{u_*}{C} \cos(\theta - \theta_w) + F \cos(\theta - \theta_w) + H \right] \sigma$
(4.37)

To select this model, the keyword *WIND GENERATION* must be set to 3 in the steering file. This source term makes use of four parameters. The default values of those parameters correspond to the coefficients proposed by Westhuysen [15].

- The coefficient D, corresponding to the keyword YAN GENERATION COEFFICIENT D, has a default value of $4.0 \ 10^{-2}$;
- The coefficient E, corresponding to the keyword YAN GENERATION COEFFICIENT E, has a default value of $5.52 ext{ } 10^{-3}$;
- The coefficient F, corresponding to the keyword YAN GENERATION COEFFICIENT F, has a default value of $5.2 ext{ } 10^{-5}$;
- The coefficient H, corresponding to the keyword YAN GENERATION COEFFICIENT H, has a default value of $-3.02 \ 10^{-4}$.

Linear wave growth: Cavaleri and Malanotte-Rizzoli model

The linear growth mechanism described by Phillips [47], [48] is useful to initialise the wave growth. If this term is neglected, it is necessary to set a non-zero sea-state as initial condition in order to enable the wave energy spectrum to grow.

The term that has been implemented in TOMAWAC is the linear wave growth term of Cavaleri & Malanotte-Rizzoli [10], as formulated by Tolman [59]:

$$Q_{in}(f,\theta) = \alpha(f,\theta) = 1,5.10^{-3} g^{-2} \left[u_* \max(0,\cos(\theta - \theta_w)) \right]^4 \exp\left[-\left(\frac{f}{f_{PM}}\right)^{-4} \right]$$
(4.38)

where u_* is the friction wind velocity, θ_w the wind direction and f_{PM} is a peak frequency called Pierson-Moskowitz frequency [50], defined as:

$$f_{PM} = \frac{1}{2\pi} \frac{g}{28.u_*} \tag{4.39}$$

To select this model, the keyword *LINEAR WAVE GROWTH* must be set to 1 in the steering file. This model does not require any input parameter.

4.3.3 Whitecapping-induced dissipations (term Q_{ds})

Two models are available in TOMAWAC. The whitecapping or the free surface slope-induced breaking is activated through the keyword *WHITE CAPPING DISSIPATION* in the steering file; the keyword can take three values, namely:

- 0 no whitecapping-induced dissipation (default value)
- 1 Komen et al. [35] and Janssen's [34] dissipation model
- 2 Westhuysen et al. dissipation model [15].

For a more detailed description of the issues related to the whitecapping dissipation modelling and of the recent advances in this field, reference can be made to [64].

Option 1 for whitecapping: Komen and Janssen dissipation model

In deep water, that term is written as follows in TOMAWAC:

$$Q_{ds} = -\frac{1}{g^4} C_{dis} \,\bar{\sigma}^9 \, m_0^2 \, \left(\delta \left(\frac{\sigma}{\bar{\sigma}} \right)^2 + (1 - \delta) \left(\frac{\sigma}{\bar{\sigma}} \right)^4 \right) \, F \tag{4.40}$$

With a finite water height, TOMAWAC uses the following formulation:

$$Q_{ds} = -C_{dis} \,\bar{\sigma} \,\bar{k}^4 \,m_0^2 \,\left(\delta \frac{k}{\bar{k}} + (1 - \delta) \left(\frac{k}{\bar{k}}\right)^2\right) F \tag{4.41}$$

 m_0 denotes the total variance, $\bar{\sigma}$ denotes the average intrinsic frequency and \bar{k} denotes the average wave number; they are respectively computed as followings:

$$m_0 = \int_{f_r=0}^{\infty} \int_{\theta=0}^{2\pi} F(f_r, \theta) df_r d\theta \tag{4.42}$$

$$\bar{\sigma} = \left(\frac{1}{m_0} \int_{f_r=0}^{\infty} \int_{\theta=0}^{2\pi} \frac{1}{\sigma} F(f_r, \theta) df_r d\theta\right)^{-1}$$
(4.43)

$$\bar{k} = \left(\frac{1}{m_0} \int_{f_r=0}^{\infty} \int_{\theta=0}^{2\pi} \frac{1}{\sqrt{k}} F(f_r, \theta) df_r d\theta\right)^{-2}$$
 (4.44)

The formulas for computing the average frequency and the average wave number are derived from those in use in WAM-cycle 4 [36]. These averages are not directly weighted by the variance spectrum, since it was found, when WAM-cycle 3 [61] was being developed, that the above expressions yielded more stable results than the conventional weighted averages. Lastly, it should be pointed out that in TOMAWAC, the above average quantities are computed not only on the discretized portion of the variance spectrum, but also analytically on the high frequency portion (up to + 8) considering a decreasing variance in f^{-n} .

That source term has two parameters:

- constant C_{dis} (corresponding to the keyword WHITE CAPPING DISSIPATION COEF-FICIENT in the steering file). Its default value is taken as 4.5, in accordance with the proposal made by Komen *et al.* [35] and the standard value adopted in the model WAM-Cycle 4.
- weighting parameter δ (corresponding to the keyword WHITE CAPPING WEIGHTING COEFFICIENT in the steering file). Its default value is taken as the 0.5 average value.

Option 2 for whitecapping: Westhuysen dissipation model

The Westhuysen dissipation model [15] is based on a saturation-based model formulation, which defines the Q_{ds} term as depending on the saturation threshold B_r .

The expression proposed by Westhuysen is:

$$Q_{ds} = -C_{dis,break} \left(\frac{B(k)}{B_r}\right)^{p_0/2} g^{1/2} k^{1/2} F(f,\theta)$$
(4.45)

where
$$B(k) = \frac{1}{2\pi} \int_0^{2\pi} C_g k^3 F(f, \theta) d\theta = C_g k^3 \frac{E(f)}{2\pi}$$

and
$$p_0\left(\frac{u_*}{C}\right) = 3 + \tanh\left[w\left(\frac{u_*}{C} - 0, 1\right)\right]$$

The variable *w* is set equal to 25.

This model is implemented in TOMAWAC in its most recent version, as formulated by Westhuysen [13], which combines the terms of Komen [35] (Q_{ds}^K) with that of Westhuysen [15] as follows:

$$Q_{ds} = f_{br}(f) \cdot Q_{ds}^{W} + (1 - f_{br}(f)) \cdot Q_{ds}^{K}$$
(4.46)

with
$$f_{br}=rac{1}{2}+rac{1}{2} anh\left\{10\left[\left(rac{B(k)}{B_r}
ight)^{1/2}-1
ight]
ight\}$$

This model is selected by setting the keyword WHITE CAPPING DISSIPATION to 2 in the steering file.

This source term makes use of 4 parameters. Their default values correspond to the coefficients proposed by Westhuysen [13].

- The coefficient $C_{dis,break}$, corresponding to the keyword WESTHUYSEN DISSIPATION COEFFICIENT, has a default value of 5.0 10^{-5} ,
- The coefficient B_r , corresponding to the keyword SATURATION THRESHOLD FOR THE DISSIPATION, has a default value of 1.75 10^{-3} ,
- The coefficient $C_{dis,non-break}$, corresponding to the keyword WESTHUYSEN WHITE CAP-PING DISSIPATION, has a default value of 3.29,
- The coefficient δ , corresponding to the keyword WESTHUYSEN WEIGHTING COEF-FICIENT, has a default value of 0.0.

4.3.4 Bottom friction-induced dissipations (term Q_{bf})

A single model is available in TOMAWAC. The bottom friction-induced dissipation is activated through the keyword *BOTTOM FRICTION DISSIPATION* in the steering file; the keyword can take two values, namely:

- 0 no bottom friction-induced dissipation (default value)
- 1 expression obtained during the JONSWAP campaign (Hasselmann *et al.* [24]) and taken up by Bouws and Komen [8].

The model used for the bottom friction-induced energy losses is an empirical expression globally representing the various contributions from the wave-bottom interaction (percolation, friction...):

$$Q_{bf} = -\Gamma \left(\frac{\sigma}{g.\sinh(k.d)}\right)^2 F \tag{4.47}$$

That (linear) expression is programmed in TOMAWAC using the following alternative form, which involves the dispersion relation:

$$Q_{bf} = -\Gamma \frac{2k}{g \cdot \sinh(2.k.d)} F \tag{4.48}$$

That source term has a single parameter:

• constant G (corresponding to the keyword *BOTTOM FRICTION COEFFICIENT* in the steering file). Its default value is taken as 0.038 m2.s-3, in accordance with what had been obtained during the JONSWAP campaign [24] and with the standard value being used in the model WAM-Cycle 4.

4.3.5 Bathymetric breaking-induced dissipations (term Q_{br})

In TOMAWAC, four parametric formulas are proposed for reproducing the effects of the bathymetric breaking-induced energy dissipation on the spectrum. The bathymetric breaking-induced dissipation is activated through the keyword *DEPTH-INDUCED BREAKING DISSIPATION* in the steering file; the keyword can take five values:

- No breaking-induced dissipation (default value)
- Battjes and Janssen's model [2]
- Thornton and Guza's model [57]
- Roelvink's model [54]

• Izumiya and Horikawa's model [32]

The first three models are parametric spectral models developed for studying the random waves, whereas the fourth one is a turbulence model initially developed for studying the regular waves. The general principle of the parametric spectral models consists in developing an expression for the total dissipation of wave energy by combining a rate of breaker-induced dissipation with a breaking probability.

Whatever model is adopted, the directional spectrum version of the bathymetric breakinginduced dissipation term is based on the assumption that breaking does not affect the energy frequency and direction distributions.

Battjes and Janssen's model (1978)

The Battjes and Janssen's breaking model [2] is based on the analogy with a hydraulic jump. Besides, it assumes that all the breaking waves have a height H_m , which is of the same order of magnitude as the water depth. The total energy dissipation term D_{br} is expressed as follows

$$D_{br} = -\frac{\alpha Q_b f_c H_m^2}{4} \tag{4.49}$$

where H_m denotes the maximum wave height being compatible with the water depth, Q_b is the fraction of breaking wave, f_c is a characteristic wave frequency and a is a numerical constant of order 1.

 H_m can be computed either through the relation:

$$H_m = \gamma_2 d \tag{4.50}$$

or through a relation derived from the Miche's criterion

$$H_m = \frac{\gamma_1}{k_c} \tanh\left(\frac{\gamma_2 k_c d}{\gamma_1}\right) \tag{4.51}$$

where k_c is linked to f_c by the linear wave dispersion relation.

 Q_b is estimated, according to the Battjes and Janssen's theory, as a solution of the implicit equation:

$$\frac{1 - Q_b}{\ln Q_b} = -\frac{H_{m0}^2}{2H_m^2} \tag{4.52}$$

In TOMAWAC, that equation can be solved either in a dichotomous way or through explicit approximations as proposed by Dingemans [16]. The latter are expressed as follows when putting:

$$b = \frac{H_{m0}}{\sqrt{2}H_{m}}$$

- version 1:

$$Q_b = 0 \qquad \text{if } b < C_b, (C_b = 0.5)$$

$$Q_b = \left(\frac{b - C_b}{1 - C_b}\right)^2 \quad \text{if } b \ge C_b$$

$$(4.53)$$

- version 2:

$$q_{0} = (2b-1)^{2} \qquad \text{if } 0.5 < b < 1$$

$$q_{0} = 0 \qquad \text{if } b \le 0.5$$

$$q_{1} = q_{0} - b^{2} \frac{q_{0} - e^{\left[(q_{0}-1)/b^{2}\right]}}{b^{2} - e^{\left[(q_{0}-1)/b^{2}\right]}}$$

$$Q_{b} = 0 \qquad \text{if } b \le C_{b}, (C_{b} = 0.3)$$

$$Q_{b} = q_{1} \qquad \text{if } C_{b} < b < 0.9$$

$$Q_{b} = q_{0} \qquad \text{if } 0.9 \le b \le 1.0$$

$$(4.54)$$

- version 3:

$$Q_b = 2.4 * b^7 (4.55)$$

The directional spectrum version of the sink term is based on the assumption that breaking does not modify the frequency and directional distribution of energy. The source term Q_{br} is then written as:

$$Q_{br}(f,\theta) = -\frac{\alpha Q_b f_c H_m^2}{4} \frac{F(f,\theta)}{m_0}$$
(4.56)

Three constants can be modified using keywords:

- constant α corresponds to the keyword *DEPTH-INDUCED BREAKING 1 (BJ) COEF-FICIENT ALPHA* in the steering file. Its default value is taken as 1, in accordance with the value as recommended by Battjes and Janssen [2]
- constant g₁ corresponds to the keyword *DEPTH-INDUCED BREAKING 1 (BJ) COEF-FICIENT GAMMA1* in the steering file. Its default value is taken as 0.88, in accordance with the value as recommended by Battjes and Janssen [2]
- constant g₂ corresponds to the keyword *DEPTH-INDUCED BREAKING 1 (BJ) COEF-FICIENT GAMMA2* in the steering file. Its default value is taken as 0.8, in accordance to the value as recommended by Battjes and Janssen [2]

The following keywords are for selecting the model options:

- The characteristic wave frequency is selected through the keyword *DEPTH-INDUCED BREAKING 1 (BJ) CHARACTERISTIC FREQUENCY*. Six values are possible:
 - 1. average frequency: $\bar{f} = \frac{\bar{\sigma}}{2\pi}$ (refer to equation (4.36.b))
 - 2. average frequency: f_{01} , computed from the spectrum moments m_0 and m_1 (default value)
 - 3. average frequency: f_{02} , computed from the spectrum moments m_0 and m_2
 - 4. discrete peak frequency: f_p
 - 5. peak frequency computed through the Read's method to order 5: f_{R5}
 - 6. peak frequency computed through the Read's method to order 8: f_{R8}
- The computation mode for breaking probability Q_b (exact computation or utilization of an explicit approximation as proposed by Dingemans [16]) is selected through the keyword $DEPTH-INDUCED\ BREAKING\ 1\ (BJ)\ QB\ COMPUTATION\ METHOD$. By default, version 2 of the explicit formulations as proposed by Dingemans is used (see above). For applications, it is recommended **not** to modify the value of that keyword.

- The computation mode for the maximum height compatible with the local water depth, H_m , is selected through the keyword *DEPTH-INDUCED BREAKING 1 (BJ) HM COM-PUTATION METHOD*. Two values are possible:
 - 1. Relation: $H_m = \gamma_2 d$ (default value)
 - 2. Miches' relation (see in (4.51) above)

Thornton and Guza's model (1983)

The Thornton and Guza's breaking model [57] is based on the analogy with a hydraulic jump and on two types of breaking wave height distribution. The energy sink term is written according to the breaking wave height distribution being retained:

function 1:
$$Q_{br1}(f,\theta) = -48\sqrt{\pi}B^3 f_c \frac{(2m_0)^{5/2}}{H_m^4 d} F(f,\theta)$$
 (4.57)

function 2:
$$Q_{br2}(f,\theta) = -12\sqrt{\pi}B^3 f_c \frac{(2m_0)^{3/2}}{H_m^2 d} \left[1 - \left(1 + \left(\frac{8m_0}{H_m^2}\right)\right)^{-5/2}\right] F(f,\theta)$$
 (4.58)

 f_c is the characteristic wave frequency (average frequency, f_{01} , f_{02} or peak frequency) and B is a parameter ranging from 0.8 to 1.5 (its default value in TOMAWAC is B=1.0). The maximum wave height compatible with the water depth, H_m , is governed by the parameter γ ($H_m = \gamma d$). The breaking model as proposed by Thornton and Guza can then be parameterized by the user via the following 4 keywords:

- DEPTH-INDUCED BREAKING 2 (TG) WEIGHTING FUNCTION Two values are possible:
 - 1. weighting function 1 (see in 4.57)
 - 2. weighting function 2 (see in 4.58) (default value)
- DEPTH-INDUCED BREAKING 2 (TG) CHARACTERISTIC FREQUENCY Six values are possible:
 - 1. average frequency: $\bar{f} = \frac{\bar{\sigma}}{2\pi}$ (refer to equation (4.36.b))
 - 2. average frequency: f_{01} , computed from the spectrum moments m_0 et m_1
 - 3. average frequency: f_{02} , computed from the spectrum moments m_0 et m_2
 - 4. discrete peak frequency: f_p
 - 5. peak frequency computed through the Read's method to order 5: f_{R5} (default value)
 - 6. peak frequency computed through the Read's method to order 8: f_{R8}
- DEPTH-INDUCED BREAKING 2 (TG) COEFFICIENT B corresponding to the B variable. Its default value in the model is taken as 1.
- DEPTH-INDUCED BREAKING 2 (TG) COEFFICIENT GAMMA corresponding to the γ variable. Its default value in the model is taken as 0.42.

Roelvink's model (1993)

The Roelvink's breaking model [54] is based on the analogy with a hydraulic jump and on two types of wave height distribution in the breaking zone (Weibull or Rayleigh). The energy sink term is written according to the wave height distribution in the breaking zone:

• Weibull's distribution:

$$Q_{br1}(f,\theta) = -\alpha f_c m A \sqrt{\frac{2}{m_0}} F(f,\theta) \int_0^\infty \left(\frac{H}{\sqrt{8m_0}}\right)^{2m+1} \exp\left[-A\left(\frac{H}{\sqrt{8m_0}}\right)^{2m}\right] \cdot \left[1 - \exp\left(-\left(\frac{H}{\gamma d}\right)\right)^N\right] dH$$
(4.59)

$$A = \left[\Gamma\left(1 + \frac{1}{m}\right)\right]^{m} \text{ with } m = 1 + 0.7\tan^{2}\left(\frac{\pi}{2}\frac{1}{\gamma_{2}}\frac{\sqrt{8m_{0}}}{d}\right)$$
(4.60)

The coefficient γ_2 is usually set to 0.65.

• Rayleigh's distribution:

$$Q_{br2}(f,\theta) = -\alpha f_c \sqrt{\frac{2}{m_0}} F(f,\theta) \int_0^\infty \left(\frac{H}{\sqrt{8m_0}}\right)^3 \exp\left[-\left(\frac{H}{\sqrt{8m_0}}\right)^2\right] \left[1 - \exp\left(-\left(\frac{H}{\gamma d}\right)\right)^N\right] dH$$
(4.61)

 f_c denotes the characteristic wave frequency (average frequency, f_{01} , f_{02} or peak frequency), a is a numerical constant of order 1, γ is the proportional control factor between the allowable wave height and the water depth (by default, $\gamma = 0.54$) and N is an exponent in the wake breaking weighting function (typically N=10).

Thus, the Roelvink's breaking model can be parameterized by the user via the following 5 keywords:

- DEPTH-INDUCED BREAKING 3 (RO) COEFFICIENT ALPHA, corresponding to the a variable. Its default value in the model is taken as 1.0.
- DEPTH-INDUCED BREAKING 3 (RO) COEFFICIENT GAMMA, corresponding to the γ variable. Its default value in the model is taken as 0.54.
- DEPTH-INDUCED BREAKING 3 (RO) COEFFICIENT GAMMA2, corresponding to the γ_2 variable. Its default value in the model is taken as 0.65.
- DEPTH-INDUCED BREAKING 3 (RO) WAVE HEIGHT DISTRIBUTION provided for retaining either a Weibull distribution (4.48) if the (default) value of the parameter is 1 or a Rayleigh distribution (4.50) if the parameter value is 2.
- DEPTH-INDUCED BREAKING 3 (RO) EXPONENT WEIGHTING FUNCTION, corresponding to the N variable. Its default value in the model is 10.
- DEPTH-INDUCED BREAKING 3 (RO) CHARACTERISTIC FREQUENCY Six values are possible:
 - 1. average frequency: $\bar{f} = \frac{\bar{\sigma}}{2\pi}$ (refer to equation (4.36.b))
 - 2. average frequency: f_{01} , as computed from the spectrum moments m_0 and m_1
 - 3. average frequency: f_{02} , as computed from the spectrum moments m_0 and m_2
 - 4. discrete peak frequency: f_p
 - 5. peak frequency as computed through the Read's method to order 5: f_{R5} (default value)
 - 6. peak frequency as computed through the Read's method to order 8: f_{R8}

Izumiya and Horikawa's turbulence model (1984)

Izumiya and Horikawa [32] sought an estimate of the dissipation by breaking-induced turbulence in the case of regular waves. From the Reynolds' equations and only considering a one-dimensional condition, they obtained an expression of the breaking-induced dissipation of wave energy in the following form:

$$\frac{d}{dx}(EC_g) = -\alpha \frac{E^{3/2}}{\rho^{1/2} d^{3/2}} \left(\frac{2C_g}{c} - 1\right)^{1/2}$$
(4.62)

E denotes the total wave energy, C_g and c are respectively group and phase velocities associated to the characteristic wave frequency f_c (average frequency f_{01} , f_{02} or peak frequency), α is a parameter governing the magnitude of the energy dissipation to be determined. For any profile, a shoal may induce the wave reforming. In order to take that process into account, Izumiya and Horikawa express the factor α in terms of a deviation from a steady state:

$$\alpha = \beta_0 \left(M_*^2 - M_{*S}^2 \right)^{1/2}$$

where M_* is a dimensionless quantity in the form: $M_*^2 = \frac{C_g}{c} \cdot \frac{E}{\rho g d^2}$

From laboratory data, Izumiya and Horikawa set M_{*S}^2 to 9 10^{-3} and β_0 to 1.8.

Assuming that the breaking does not affect the frequency and direction distribution of energy, the dissipation term is lastly written as:

$$Q_{br}(f,\theta) = -\beta_0 \left(\frac{C_g}{c} \cdot \frac{m_0}{d^2} - M_{*S}^2\right)^{1/2} \frac{g^{1/2} m_0^{1/2}}{d^{3/2}} \left(\frac{2C_g}{c} - 1\right)^{1/2} F(f,\theta)$$
(4.63)

Thus, the breaking model as proposed by Izumiya and Horikawa can be parameterized by the user through the three following keywords:

- DEPTH-INDUCED BREAKING 4 (IH) COEFFICIENT BETA0, corresponding to the β_0 variable. The default value in the model is 1.8.
- DEPTH-INDUCED BREAKING 4 (IH) COEFFICIENT M2STAR, corresponding to the M_{*S}^2 variable. The default value in the model is 0.009.
- DEPTH-INDUCED BREAKING 4 (IH) CHARACTERISTIC FREQUENCY Six values are possible:
 - 1. average frequency: $\bar{f} = \frac{\bar{\sigma}}{2\pi}$ (refer to equation (4.29.b))
 - 2. average frequency: f_{01} , as computed from the spectrum moments m_0 and m_1
 - 3. average frequency: f_{02} , as computed from the spectrum moments m_0 and m_2
 - 4. discrete peak frequency: f_p
 - 5. peak frequency as computed through the Read's method to order 5: f_{R5} (default value)
 - 6. peak frequency as computed through the Read's method to order 8: f_{R8}

4.3.6 Non-linear quadruplet interactions (term Qnl)

Three non-linear quadruplet interactions models are available in TOMAWAC. The non-linear quadruplet interactions are activated through the keyword *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* in the steering file; the keyword can take four values, namely:

- 0 no non-linear quadruplet interaction (default value)
- 1 DIA method (Discrete Interaction Approximation) of Hasselmann *et al.* [25] [26] which is a discrete parameterization of the exact computation operator as proposed by Hasselmann [22] [23]
- 2 MDIA method (Multiple DIA) as proposed by Tolman [60]
- 3 Quasi exact GQM method (Gaussian Quadrature Method) as introduced by Lavrenov [38] and implemented by Gagnaire-Renou et al. [20]

Option 1 for non-linear quadruplet interactions: DIA method

The method and its implementation in TOMAWAC have been the subject of a specific report [4] which the reader is invited to refer to for further information. The major teachings of the DIA method are summarized below.

The exact expression of the deep water interactions term as set by Hasselmann [22] [23], expressed herein for convenience as a function of the wave number vector, is analogous to a Boltzmann integral:

$$Q_{nl}^{exact} = \iiint \sigma_4.G.\delta\left(\mathbf{k}_1 + \mathbf{k}_2 - \mathbf{k}_3 - \mathbf{k}_4\right)\delta\left(\sigma_1 + \sigma_2 - \sigma_3 - \sigma_4\right)$$

$$\left[\frac{F(\mathbf{k}_1)}{\sigma_1}\frac{F(\mathbf{k}_2)}{\sigma_2}\left(\frac{F(\mathbf{k}_3)}{\sigma_3} + \frac{F(\mathbf{k}_4)}{\sigma_4}\right) - \frac{F(\mathbf{k}_3)}{\sigma_3}\frac{F(\mathbf{k}_4)}{\sigma_4}\left(\frac{F(\mathbf{k}_1)}{\sigma_1} + \frac{F(\mathbf{k}_2)}{\sigma_2}\right)\right]d\mathbf{k}_1d\mathbf{k}_2d\mathbf{k}_3 \quad (4.64)$$

The energy exchanges, in that integral (*a* priori rather uneasily computable), take place between quadruplets meeting the resonance conditions:

$$\sigma_1 + \sigma_2 = \sigma_3 + \sigma_4 \text{ and } \mathbf{k}_1 + \mathbf{k}_2 = \mathbf{k}_3 + \mathbf{k}_4$$
 (4.65)

as evidenced by the two Dirac functions δ in the integral.

G denotes the value of the coupling term for the resonant quadruplet interactions $(\mathbf{k}_1, \mathbf{k}_2, \mathbf{k}_3, \mathbf{k}_4)$. Establishing and computing its expression is also an awkward task. Hasselmann [22] proposed a computation mode that was also taken up and given a more concise form by such other authors as Webb [62].

The exact computation of the above Boltzmann integral is too complex and time-consuming for such a sea state operational model as TOMAWAC (see e.g. [25]). That is why, starting from the experiment as developed in WAM [61] [36], TOMAWAC uses the DIA (Discrete Interaction Approximation) approximate computation method as proposed by Hasselmann *et al.* [25]. Whereas the exact computation requires the summation of the contributions from a great number of quadruplets, the approximate computation implements only a small number of quadruplet configurations which are all similar.

That standard interaction configuration is defined as follows:

- two of the wave numbers are alike: $\mathbf{k}_1 = \mathbf{k}_2 = \mathbf{k}$, which also involves that the two related frequencies are identical: $\sigma_1 = \sigma_2 = \sigma$
- the other two frequencies s3 and s4 are defined by:

$$\sigma_3 = (1 + \lambda)\sigma = \sigma^+$$

 $\sigma_4 = (1 - \lambda)\sigma = \sigma^-$

Through the value $\lambda=0.25$, a good correlation with the exact computation of the integral (see[25]) could be achieved. That value is used in the model WAM [61] [36] and is taken up in TOMAWAC.

- since the wave vectors $\mathbf{k}_3 = \mathbf{k}^+$ and $\mathbf{k}_4 = \mathbf{k}^-$ should observe the resonance condition, it can be shown they are featured by angles $\theta_3 = 11.5^\circ$ and $\theta_4 = -33.6^\circ$ with respect to the common direction of $\mathbf{k}_1 = \mathbf{k}_2 = \mathbf{k}$ (refer to [25]).
- Furthermore, the mirror image is taken into account by considering the vectors as symmetrical with respect to the direction of $\mathbf{k}_1 = \mathbf{k}_2 = \mathbf{k}$.

The standard interaction configuration (in full line) and its mirror image (in dotted line) are shown schematically in Figure 4.2.

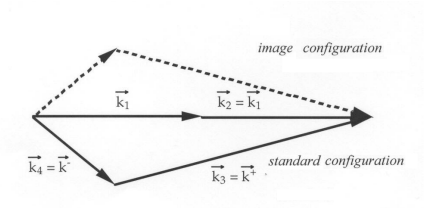


Figure 4.2: Schematic standard interaction configuration for the DIA method

With this standard configuration, the non-linear interaction term for all four resonant wave numbers is written as [25]:

$$\begin{bmatrix} Q_{nl} \\ Q_{nl}^- \\ Q_{nl}^+ \end{bmatrix} = \begin{bmatrix} -2 \\ 1 \\ 1 \end{bmatrix} \Pi g^{-4} f_r^{11} \left(F^2 \left(\frac{F^+}{(1+\lambda)^4} + \frac{F^-}{(1-\lambda)^4} \right) - \frac{2FF^+F^-}{(1-\lambda^2)^4} \right)$$
(4.66)

With such a computation method, the vector \mathbf{k} scans all the discretization nodes of the directional spectrum mesh. The number of configurations being considered is then twice as large as the number of points in that mesh. In relation to the full computation, the 5-dimensional space (three integration dimensions and two dimensions for \mathbf{k}_4) of all the possible resonant quadruples is reduced to a 2-dimensional space.

In a finite water depth, from exact computations of the Boltzmann integral, Herterich and Hasselmann [29] suggested to make a deep water computation based on the previous method, then to multiply it by a coefficient R representing the effect of the finite water height:

$$Q_{nl}(d) = R.Q_{nl}(d = \infty) \tag{4.67}$$

Coefficient R is a function of the normalized water height $\bar{k}.d$ and is expressed as follows:

$$R(\chi) = 1 + \frac{5.5}{\chi} (1 - \frac{5}{6}\chi) \exp\left(-\frac{5}{4}\chi\right) \text{ where } \chi = \frac{3}{4}\bar{k}.d$$
 (4.68)

The average wave number \bar{k} was defined in the previous paragraph (see 4.44).

In its authors' opinion, that relation remains valid as long as $\bar{k}.d > 1$. It is used as such in TOMAWAC for the finite water depth computations.

That source-term has a single parameter:

• constant I (corresponding to the keyword STANDARD CONFIGURATION PARAMETER of the steering file). Its default value is taken as 0.25, in accordance with the proposal made by Hasselmann et al. [25] and with the standard value in the model WAM-Cycle 4.

Option 2 for non-linear quadruplet interactions: MDIA method

The MDIA method (multiple DIA) is an extension of the DIA algorithm. We use here the version proposed by Tolman [60]

This method can give very reasonable results in simple situations, but in case of unsteady or rapidly changing sea-state conditions (e.g. in the situation of abrupt changes of wind direction) it results in significant qualitative and quantitative differences when compared with exact methods [5].

The MDIA method consists of using a quadruplet depending on 2 parameters, λ and μ , defined as follows:

$$\mathbf{k}_0 + \mathbf{k}_1 = \mathbf{k}_2 + \mathbf{k}_3 = 2\mathbf{k} \tag{4.69}$$

$$\sigma_0 = (1 + \mu)\sigma, \quad \sigma_1 = (1 - \mu)\sigma$$

$$\sigma_2 = (1 + \lambda)\sigma, \quad \sigma_3 = (1 - \lambda)\sigma$$
(4.70)

The λ and μ values proposed by Tolman that allow to best estimate the Q_{nl4} source term in the case of 4 interacting quadruplets are shown in Figure 4.3

Component	λ	μ
1	0.075	0.023
2	0.219	0.127
3	0.299	0.184
4	0.394	0.135

Figure 4.3: Values of the parameters λ and μ proposed by Tolman to best estimate quadruplet interactions with the MDIA method, in the case of 4 interacting quadruplets [60].

To select this model in TOMAWAC the keyword *NON-LINEAR TRANSFERS BETWEEN FRE-QUENCIES* must be set to 2 in the steering file.

This model does not require any other parameter: the values of the λ and μ parameters are set as constants in the code. However they can be modified when considering a larger number of interacting quadruplets.

Option 3 for non-linear quadruplets interactions: GQM method

The Gaussian Quadrature Method (GQM) is based on the use of Gaussian quadratures for the different numerical integrations arising in evaluating Equation 4.64. This technique, proposed by Lavrenov [38], has been developed and optimised to adequate results regarding both precision and CPU time [5], [19], [20]

Several steps are needed to transform Equation 4.64 into an expression that can be integrated via Gaussian quadrature method. They can be summarized as follows (for a detailed description reference can be made to [19]):

1. Elimination of the Dirac function on the wave numbers $\delta(\mathbf{k}_0 + \mathbf{k}_1 - \mathbf{k}_2 - \mathbf{k}_3)$, by imposing $\mathbf{k}_3 = \mathbf{k}_0 + \mathbf{k}_1 - \mathbf{k}_2$ (see the resonance condition, Equation 4.65). Equation 4.64 is therefore reduced to an integral with 4 dimensions, including a single Dirac function on the frequency σ .

- 2. Variable change, to work with (σ, θ) instead of k, and reformulation of the equation in terms of variance density (F) instead of wave action density (N).
- 3. Integration over θ_2 and elimination of the Dirac function on the frequency. A 3-dimension integral is obtained, without any Dirac function.
- 4. Final expression of the non-linear transfer term: the variables σ_3 , θ_2 and θ_3 , are expressed as functions of σ_1 , θ_1 and σ_2 , and of σ_0 and θ_0 , which Q_{nl4} term depends on. The variables σ_a , k_a et ε_a defined by $\sigma_a = \sigma_0 + \sigma_1$, $k_a = k_0 + k_1$ and $\varepsilon_a = 2gk_a/\sigma_a^2$, and depending only on $(\sigma_0$ and $\theta_0)$ and $(\sigma_1$ and $\theta_1)$, are used as well.

$$Q_{nl4} = \int_{\sigma_{1}=0}^{+\infty} \int_{\theta_{1}=0}^{2\pi} \int_{\sigma_{2}=0}^{\sigma_{a}/2} 2 \frac{\sigma_{a}^{4} T}{\sigma_{1} \sigma_{2} \sigma_{3}} \frac{F_{2} F_{3} (F_{0} \sigma_{1}^{4} + F_{1} \sigma_{0}^{4}) - F_{0} F_{1} (F_{2} \sigma_{3}^{4} + F_{3} \sigma_{2}^{4})}{\sqrt{\tilde{B}_{0} (\varepsilon_{a}, s_{2}) \tilde{B}_{1} (\varepsilon_{a}, s_{2}) \tilde{B}_{2} (\varepsilon_{a}, s_{2})}} d\sigma_{1} d\theta_{1} d\sigma_{2}$$

$$(4.71)$$

where s_2 is defined as $s_2 = \sigma_2/\sigma_a$.

Equation 4.71 is then integrated using different quadrature methods:

- Gauss-Legendre or Gauss-Chebyshev quadratures are used for the integration over σ_2 , depending on the ε_a values determining number and type of singularities.
- Gauss-Chebyshev quadratures are used for the integration over θ_1 .
- The integration over σ_1 is realized using the trapezoidal rule.

Three different GQM method resolutions have been tested:

- A "fine" resolution, considered as the exact calculation of the non-linear transfer term, as no improvement in the results is noticed when further increasing the method resolution.
- An "intermediate" resolution.
- A "coarse" resolution, whose parameters are given as default values in TOMAWAC, which represents the best compromise between accuracy of the solution and CPU time.

The configurations that do not effect significantly the global computation of Q_{nl4} are neglected. This configuration selection allows to reduce the CPU time. The threshold values set as default in TOMAWAC reduce the number of configuration:

- by 21% in the "fine" resolution case
- by 34% in the "intermediate" resolution case
- by 64% in the "coarse" resolution case

The GQM method is selected by setting to 3 the keyword *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* in the steering file.

This method makes use of 6 parameters. The default values of those parameters correspond to the "coarse" resolution case:

• The three keywords SETTING FOR INTEGRATION ON OMEGA1, SETTING FOR INTEGRATION ON OMEGA2 and SETTING FOR INTEGRATION ON THETA1 determine the number of integration points over the three variables σ_1 , θ_1 and σ_2 and their default values are respectively 3, 3 and 6 ("coarse" resolution). The values 1, 4, 8 and 2, 8, 12 correspond respectively to the "intermediate" and "fine" resolution cases.

• The three keywords *THRESHOLDO FOR CONFIGURATIONS ELIMINATION*, *THRESH-OLD1 FOR CONFIGURATIONS ELIMINATION* and *THRESHOLD2 FOR CONFIGU-RATIONS ELIMINATION* affect the percentage of discarded configurations. Their default values are respectively 0, 10¹⁰ and 0.15. For the "intermediate" and "fine" resolution cases, the first two values are the same, and the threshold2 values is equal respectively to 0.01 and 0.001.

4.3.7 Non-linear transfers between triads (Qtr term)

Two non-linear triads interactions model are available in TOMAWAC. The triad model is activated through the keyword *TRIAD INTERACTIONS* in the steering file. The keyword can take three values

- 0 No triad interaction (default value)
- 1 LTA model interaction, a model proposed by Eldeberky [17] see 4.3.7.
- 2 SPB model proposed by Becq [3] see 4.3.7.

LTA (Lumped Triad Approximation) model

A parametric model allowing to take into account the non-linear triad interactions in the averaged-phase models has been proposed by Eldeberky and Battjes [17]. The LTA model is a parametric approach that is based on the Madsen and Sorensen's deterministic spectral model [42]. Simplifying hypotheses are introduced for reducing the computation cost. Thus, a parametric formulation is given for the biphase as a function of the Ursell's number and the model is restricted to the self-interactions.

The source term is written as:

$$Q_{LTA}(f,\theta) = Q_{LTA}^{+}(f,\theta) + Q_{LTA}^{-}(f,\theta)$$

$$Q_{LTA}^{+}(f,\theta) = \alpha_{LTA} c C_g g^2 \frac{R_{f/2,f/2}^2(f/2,f/2)}{S_f^2} \sin \left| \beta_{f/2,f/2} \right| \left[F^2(f/2,\theta) - 2F(f,\theta)F(f/2,\theta) \right]$$

$$Q_{LTA}^{-}(f,\theta) = -2 Q_{LTA}^{+}(2f,\theta)$$
(4.72)

 α_{LTA} is the model adjustment coefficient; c and C_g denote the phase and group velocities, respectively.

R is the self-interaction coefficient:

$$R_{f,f} = (2k)^2 \left[\frac{1}{2} + \frac{(2\pi f)^2}{gd \, k^2} \right] \tag{4.73}$$

S is given by the relation:

$$S_f = -2k \left(gd + 2Bgd^3k^2 - (B+1/3)d^2(2\pi f)^2 \right)$$
 (4.74)

The biphase β is given by the relation:

$$\beta(f,f) = -\frac{\pi}{2} + \frac{\pi}{2} \tanh\left(\frac{0.2}{Ur}\right) \tag{4.75}$$

where Ur denotes the Ursell's number:

$$Ur = \frac{g}{8\pi^2\sqrt{2}} \frac{H_{m0}T_m^2}{d^2} \tag{4.76}$$

with H_{m0} being the significant spectral height and T_m being the average wave time.

 Q_{LTA}^{\pm} denotes the negative and positive contributions of the self-interactions. Since Q_{LTA}^{+} denotes the positive contributions to the first upper harmonic, it should be positive. The negative values of Q_{LTA}^{+} are replaced by the zero value. In the numerical integration of the energy equation, the source term for the triad interactions is generally only computed for frequencies that are lower than $R_{fm}f_m$ (Ris [52]) where $R_{fm}=2.5$.

Two constants can me modified through keywords:

- constant α_{LTA} corresponding to the keyword TRIADS 1 (LTA) COEFFICIENT ALPHA. Its default value is $\alpha_{LTA} = 0.5$
- constant R_{fm} corresponding to the keyword TRIADS 1 (LTA) COEFFICIENT RFMLTA. Its default value is $R_{fm} = 2.5$

SPB model

The SPB model was developed by Becq [Becq, 1998] from the extended Boussinesq equations as proposed by Madsen and Sorensen [41]. The model is for simulating the effects induced by the collinear and non-collinear interactions of spectral components. The source term is written as:

$$Q(f,\theta) = \frac{B'g}{2S_{1,f}} \int_{0}^{f} \int_{0}^{2\pi} \int_{0}^{f} \int_{0}^{2\pi} df_{1} df_{2} d\theta_{1} d\theta_{2} T_{f_{1},f_{2}} \delta\left(\theta_{k} - \theta_{k_{1}+k_{2}}\right) \delta\left(f - f_{1} - f_{2}\right) + \frac{B'g}{S_{1,f}} \int_{0}^{\infty} \int_{0}^{2\pi} \int_{0}^{\infty} \int_{0}^{2\pi} df_{1} df_{2} d\theta_{1} d\theta_{2} T_{-f_{2},f_{1}} \delta\left(\theta_{k_{1}} - \theta_{k+k_{2}}\right) \delta\left(f_{1} - f - f_{2}\right)$$

$$(4.77)$$

with:

$$T_{f_1,f_2} = \frac{gK}{K^2 + \Delta k^2} R_{f_1,f_2} \left[-\frac{R_{-f_2,f}}{S_{2,f_1}k_1} F F_2 - \frac{R_{-f_1,f}}{S_{2,f_2}k_2} F F_1 + \frac{R_{f_1,f_2}}{S_{2,f}k} F_1 F_2 \right]$$
(4.78)

$$B' = \frac{Cg}{2\pi k} \tag{4.79}$$

$$R_{f1,f2} = (k_1 + k_2)^2 \left[\frac{1}{2} + \frac{(2\pi)^2 f_1 f_2}{g d k_1 k_2} \right]$$
 (4.80)

$$S_f = -2k \left(gd + 2Bgd^3k^2 - (B+1/3)d^2(2\pi f)^2 \right)$$
 (4.81)

F denotes the variance spectrum in terms of frequencies and directions, $F(f, \theta)$. T_{f_1, f_2} and T_{-f_2, f_1} respectively correspond to the sum and difference interactions. K is the model adjustment parameter.

Since the model was designed for taking into account the energy transfers for all the possible triad configurations within the spectrum, the computation times are very long. In order to shorten these computation times, the interactions can be restricted over a range of spectral components that are included within a given angular sector. Thus, directional limits can be user-prescribed.

Three constants can be modified through keywords:

- constant K corresponding to the keyword 'TRIADS 2 (SPB) COEFFICIENT K. Its default value is K=0.34
- the lower and upper directional markers corresponding to the keywords *TRIADS 2 (SPB) LOWER DIRECTIONAL BOUNDARY* and *TRIADS 2 (SPB) UPPER DIRECTIONAL BOUNDARY*. Their respective default values are 0 and 360.

4.3.8 Wave blocking effects (upper limit for spectrum or dissipation $Q_{ds,cur}$)

When water waves meet a strong adverse current, with a velocity that approaches the wave group velocity, waves are blocked. Two options can be considered in TOMAWAC to take into account wave blocking effects.

Option 1: consider an equilibrium range spectrum (in the presence of ambient flow) applied as an upper limit for the spectrum (Hedges et al. [27])

Option 2: add a dissipative term on the right-hand side of the action balance equation [14] The inclusion of wave-blocking effects is activated through the keyword *DISSIPATION BY STRONG CURRENT* in the steering file; the keyword can take three values, namely:

- 0 No wave-blocking effects (default value)
- 1 Upper limit for spectrum Hedges et al [27] see 4.3.8.
- 2 Westhuysen *et al.* [14] enhanced dissipation of waves on a strong current see4.3.8.

option 1 for wave blocking: upper limit for the spectrum

An upper limit is imposed to the spectrum, using a Phillips [49] shape, i.e.

If
$$E(f) > E \max = \frac{\alpha g^2}{(2\pi)^4 f^5}$$
 then $F(f, \theta) = \frac{E \max}{E(f)} F(f, \theta)$

with α the Phillips's constant in the Pierson-Moskowitz spectrum, equal to = 0.0081.

This model is selected by setting the keyword *DISSIPATION BY STRONG C URRENT* to 1 in the steering file.

option 2 for wave blocking: Westhuysen formulation

The expression proposed by Westhuysen is:

$$Q_{ds,cur} = -C_{ds,cur} \max \left(\frac{f_r}{f}, 0\right) \left(\frac{B(k)}{B_r}\right)^{p_0/2} F(f, \theta)$$

where

$$B(k) = \frac{1}{2\pi} \int_0^{2\pi} C_g k^3 F(f, \theta) d\theta = C_g k^3 \frac{E(f)}{2\pi}$$

and

$$p_0\left(\frac{u_*}{C}\right) = 3 + \tanh\left[w\left(\frac{u_*}{C} - 0, 1\right)\right]$$

The variable w is set equal to 25.

This model is selected by setting the keyword *DISSIPATION BY STRONG CURRENT* to 2 in the steering file.

This source term makes use of 2 parameters. Their default values correspond to the coefficients proposed by Westhuysen [14]:

- The coefficient $C_{ds,cur}$, corresponding to the keyword DISSIPATION COEFFICIENT FOR STRONG CURRENT has a default value of 0.65,
- The coefficient B_r , corresponding to the keyword SATURATION THRESHOLD FOR THE DISSIPATION, has a default value of 1.7510^{-3} .

4.3.9 Dissipation due to vegetation.

When the ratio between vegetation height and water depth is important, vegetation can imply some dissipation. Some methods exist to modelize this dissipation. The method we chose is based on the formulation proposed by Suzuki et al. [56]

$$Q_{veg} = -\sqrt{rac{2}{\pi}} ilde{C}_D b_{v} N_{v} \left(rac{ar{k}}{ar{\sigma}}
ight)^3 rac{\sinh^3(ar{k}lpha h) + 3\sinh(ar{k}lpha h)}{3ar{k}\cosh^3(ar{k}h)} \sqrt{m_0} F(oldsymbol{\sigma}, oldsymbol{ heta})$$

 m_0 denotes the total variance, $\bar{\sigma}$ denotes the average intrinsic frequency and \bar{k} denotes the average wave number; they are respectively computed as followings:

$$m_0 = \int_{f_r=0}^{\infty} \int_{\theta=0}^{2\pi} F(f_r, \theta) \, df_r \, d\theta \tag{4.82}$$

$$\bar{\sigma} = \left(\frac{1}{m_0} \int_{f_r=0}^{\infty} \int_{\theta=0}^{2\pi} \frac{1}{\sigma} F(f_r, \theta) \, df_r \, d\theta\right)^{-1}$$
 (4.83)

$$\bar{k} = \left(\frac{1}{m_0} \int_{f_r=0}^{\infty} \int_{\theta=0}^{2\pi} \frac{1}{\sqrt{k}} F(f_r, \theta) \, df_r \, d\theta\right)^{-2} \tag{4.84}$$

Where \tilde{C}_D is the vegetation drag coefficient (BULK DRAG COEFFICIENT), $b_v[m]$ is the stem diameter of cylinder (plant) (STEM DIAMETER), N_v is the number of plants per square meter (NUMBER OF PLANTS M2) and α is the ratio between vegetation height (VEGETATION HEIGHT) and water depth.

The vegetation is activated through the keyword *VEGETATION TAKEN INTO ACCOUNT*The area can be defines through a list of polygons using a files. *FILE WITH DEFINITION OF POLYGONS*

4.3.10 Dissipation due to porous media.

The propagation of wave over a porous medium implies some dissipation. The dispersion equation for waves that propagate over a dissipative medium derived by [40] was expressed in general form by [44]

$$\Gamma - x \tanh(\beta x) - \Phi \tanh(\alpha x) [x - \Gamma \tanh(\beta x)] = 0 \tag{4.85}$$

where x = Kh a dimensionless complex wave number, $\Gamma = \sigma^2 h/g$ the relative wave frequency, αh the thickness of the porous layer, h the water depth, g the acceleration due to gravity, $\Phi = \tau/(s-if)$ the dissipation factor, s the inertial coefficient $s = 1 + C_a(1-\tau)/\tau$, with τ the porosity rate and C_a the damping mass coefficient, f the linear friction coefficient and $g = 1 - \alpha$. If Γ , Φ and σ are known, it can be shown that the solution of equation 4.85 gives the sink term of the equations used by TOMAWAC under the form.

$$Q_{porous} = -2K_i \tilde{C}_g F(f, \theta) \tag{4.86}$$

where K_i is the imaginary part of the complex solution of the equation 4.85 and \tilde{C}_g is the group velocity.

Solving 4.85 is not so easy but in [12], the authors proposed a method based on taylor developments that solves it. That is what is implemented in TOMAWAC

An approximation is done with the calculation of the group velocity in that case, we calculate it considering that the depth is βh . which is in between the velocity for a depth of h and the real velocity of a porous media which would be difficult to calculate.

The area can be defines through a list of polygons using a files. FILE WITH DEFINITION OF POLYGONS

4.4 Surface rollers 41

4.4 Surface rollers

4.4.1 Theoretical background

When waves break, a part of the wave energy is transferred to surface rollers. These propagate shoreward, thus causing a delay between the point, where the waves begin to break and the point, where wave setup and longshore currents develop. In additional the surface rollers transfer mass towards the coast, thus influencing the return current that occurs. Finally, they can influence the stirring up of sediment. In the current implementation, only the effect of surface rollers on the longshore current is taken into account.

The surface rollers can be switched on by setting $SURFACE\ ROLLERS$ to TRUE. In the implementation, we follow the work by [53]. According to this reference, the surface roller energy E_r per unit of volume, integrated over the water column is given by:

$$E_r = \frac{1}{2}A \frac{\overline{U_{roller}^2 + W_{roller}^2}}{L} \tag{4.87}$$

Here, L is the wave length, A is cross-sectional area of the surface roller, and U_{roller} and W_{roller} the velocity components in the horizontal and vertical direction respectively. The evolution of the surface roller energy is given by the following differential equation:

$$\frac{\partial E_r}{\partial t} + \frac{\partial c_x E_r}{\partial x} + \frac{\partial c_y E_r}{\partial y} = D_w - D_r \tag{4.88}$$

Here c_x and c_y are the x and y components of the phase velocity of the waves. Using this formulation, effects of wave current interaction are not taken into account. D_w is the energy dissipation due to breaking of the waves, and D_r is the energy dissipation of the surface rollers, which is parametrized as:

$$D_r = 2\frac{\beta_s}{\beta_2} \frac{g}{c} E_r \tag{4.89}$$

Here, β_s and β_2 are calibration parameters, and g is the gravitational acceleration. The parameters are set with the keywords *BETA S SURFACE ROLLERS* and *BETA 2 SURFACE ROLLERS*. The effect of the surface rollers of the current comes from its effect on the radiation stresses, which are given by:

$$S_{xx} = S_{xx,waves} + E_r \cos^2 \theta \tag{4.90}$$

$$S_{xy} = S_{xy,wayes} + E_r \cos \theta \sin \theta \tag{4.91}$$

$$S_{yy} = S_{yy,waves} + E_r \sin^2 \theta \tag{4.92}$$

Here, S_{xx} , S_{xy} and S_{yy} are the different components of the radiation stress tensor, $S_{xx,waves}$, $S_{xy,waves}$ and $S_{yy,waves}$ the radiation stress components from the surface waves (as calculated in TOMAWAC), and θ the mean wave direction.

In TOMAWAC, the surface roller energy E_r , breaking wave dissipation D_w , and roller wave dissipation D_r can all be exported to an output file by adding respectively SRE, DBR, and DSR to the keyword $VARIABLES\ FOR\ 2D\ GRAPHIC\ PRINTOUTS$.

4.4.2 Implementation

The surface roller energy balance is calculated after each time step in TOMAWAC. It is possible to perform multiple substeps for the calculation of the surface roller energy balance by setting the keyword *NUMBER OF SURFACE ROLLER TIME STEPS*. The surface roller energy balance is calculated using a fractional step method. The advection step is solved first using the method of characteristics. The velocity field is determined every time step from the peak period, water depth and mean wave direction calculated by TOMAWAC (for the moment ignoring

the effect of wave-current interaction). After the advection step, the source and sink terms are applied. Here, D_r is calculated using an implicit numerical discretisation while D_w uses an explicit discretisation. D_w is exported directly from the calculation of the depth induced breaking source term in TOMAWAC.

5. Discretizations used in TOMAWAC

The main aspects concerning the numerical discretization in TOMAWAC are presented and discussed herein for the two spatial variables (paragraph 5.1), for the two spectro-angular variables (paragraph 5.2) and for the time domain (paragraph 5.3).

5.1 Spatial discretization

The spatial coordinate system, whether it is Cartesian or spherical, is a planar two-dimensional domain that is meshed by means of triangular finite elements. Only the maritime portion of the computational domain is meshed, so that all the computational points of the spatial grid are provided with a water depth that is strictly above zero. Through this discretization technique, the mesh size may naturally be variable over the spatial domain, particularly enabling to get a fine grid in the areas of specific interest, featured either by complex geometries (straits, intracontinental seas, bays...) or by high bathymetric gradients. Furthermore, that spatial grid may include one or more islands.

The number of discretization points is only limited by the RAM capacities of the computing machine. The equation solved by Tomawac does not prescribe *a priori* any conditions about the number of grid points per wave length. The density of spatial discretization points is left at the user's will. It should match, however, both spatial and temporal scales of variation of the physical characteristics of the domain being studied, in particular bathymetry and wind field. In the general case, this spatial grid is realised on a workstation using one of the mesh generators associated to the TELEMAC system (refer to the 7.2.2 for further details about the preparation of the grid). Two examples of spatial grids developed for Tomawac for simulated storms in the North Atlantic Ocean, the Channel and the North Sea are illustrated in Figure 5.1.

5.2 Spectro-angular discretization

5.2.1 Frequency discretization

In TOMAWAC, the frequency domain is discretized considering a series of NF frequencies in a geometric progression:

 $f_n = f_1.q^{n-1}$ with n ranging from 1 to NF

The minimum frequency is then f_1 and the maximum frequency is $f_1 ext{.} q^{NF-1}$.

In order to define the frequency discretization, the user should specify as an input into the steering file:

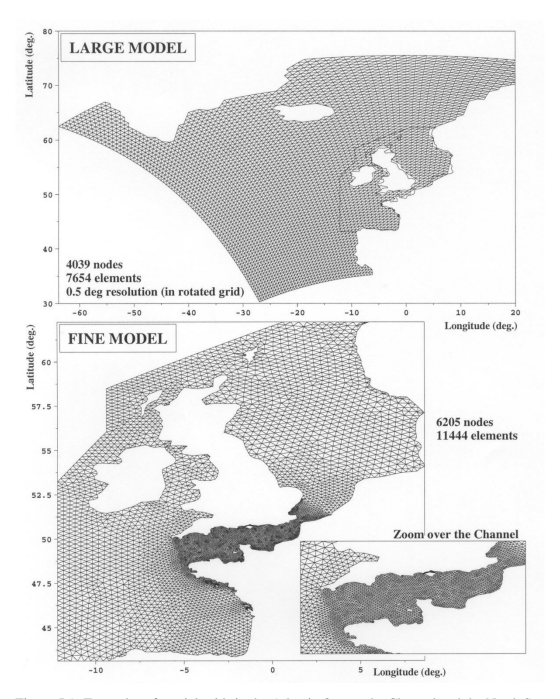


Figure 5.1: Examples of spatial grids in the Atlantic Ocean, the Channel and the North Sea

- the frequency number: *NF* (corresponding to the keyword *NUMBER OF FREQUEN-CIES* in the steering file)
- the minimum frequency: f_1 (in Hertz) (corresponding to the keyword MINIMAL FRE-QUENCY in the steering file)
- the frequential ratio: q (corresponding to the keyword *FREQUENTIAL RATIO* in the steering file)

5.2.2 Directional discretization:

The interval of propagation direction [0,360°] is discretized into ND evenly distributed directions, so that these directions are:

 $\theta_m = (m-1).360/ND$ with m ranging from 1 to ND

In order to define the directional discretization, the user should specify as an input into the steering file:

- the direction number: *ND* (corresponding to the keyword *NUMBER OF DIRECTIONS* in the steering file).
- The direction convention selected for the input/output directional variables: either nautical or counterclockwise (corresponding to the keyword *TRIGONOMETRICAL CONVENTION* in the steering file, the default value of which is NO). The nautical convention sets the wave propagation directions (towards which the waves are propagating) in relation to the true North or the vertical axis and opposite to the counterclockwise direction. The counterclockwise convention sets the wave propagation directions in relation to the horizontal axis.
- Note that the convention selected for computing the directions within the FORTRAN model always defines the propagation directions in the clockwise direction from the true North, even though the keyword *TRIGONOMETRICAL CONVENTION*=YES!

5.2.3 Spectro-angular grid:

A two-dimensional grid for spectro-angular discretization is achieved by combining the above defined frequency and directional discretizations. That grid has NF.ND points.

A polar representation is used in TOMAWAC, where the wave frequencies are measured radially and where the propagation direction corresponds to the value of the angle in relation to the axis selected by the user as (vertical or horizontal) origin. An example of a spectro-angular grid having 25 frequencies and 12 directions is illustrated in Figure 5.2

5.3 Temporal discretization

In TOMAWAC, each computation begins at the internal date 0, to which an actual date being defined by the keyword *DATE OF COMPUTATION BEGINNING* in the steering file can be associated. That date is specified as per the yymmddhhmm format which corresponds to the moment dd/mm/yy at hh:mm (for example, 9505120345 corresponds to May 12, 1995 at 3.45). The evolution equation of the directional spectrum of wave action density is integrated with a constant time step which is expressed in seconds through the keyword *TIME STEP* in the steering file. Sub-iterations of that time step can also be made for computing the source terms (refer to paragraph 6.3). That number of sub-time steps per time step is defined in the steering file through the keyword *NUMBER OF ITERATIONS FOR THE SOURCE TERMS*.

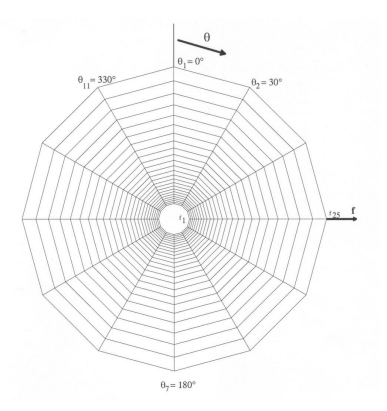


Figure 5.2: Example of a spectro-angular grid as used by TOMAWAC (25 frequencies and 12 directions in this case)

6. Numerical methods used in TOMAWAC

6.1 General solution algorithm

As stated in Section 4, the equation to be solved by the TOMAWAC software is a transport (convection) equation with source terms that can be written in the following general form:

$$\frac{\partial (BF)}{\partial t} + \mathbf{V} \cdot \nabla (BF) = BQ \tag{6.1}$$

Both functions F and Q are functions of five variables and depend, e.g. in Cartesian coordinates, on x, y, θ, f_r and t. The above equation is then to be solved on a four-dimensional grid in (x, y, θ, f_r) and \mathbf{V} is a transport vector, which is a dimension-4 vector in the general case. It is reduced, however, to a three-dimensional vector (\dot{f}_r) is zero) when there is neither a current nor a variation of water depth in time.

$$\mathbf{V} = \begin{pmatrix} \dot{x} \\ \dot{y} \\ \dot{\theta} \\ \dot{f}_r \end{pmatrix} \tag{6.2}$$

Equation (6.1) is solved in TOMAWAC through a fractional step method, i.e. the convection and the source term integration steps are solved successively and separately. Thus, the following steps are successively solved from a current state at the date $t = n.\Delta t$, in which the variance spectrum F^n is known in all points:

an **advection step** without source terms (refer to paragraph 6.2):

$$\frac{\partial (BF)}{\partial t} + \mathbf{V} \cdot \nabla (BF) = 0 \tag{6.3}$$

discretized as follows:

$$\frac{(B.F) * - (B.F)^n}{\Delta t} = \left[\mathbf{V}.grad(B.F) \right]^n \tag{6.4}$$

from which a value of $(B.F)^*$, then of F^* , intermediate after the convection step, is derived a **source term integration step** (refer to paragraph 6.3):

$$\frac{\partial F}{\partial t} = Q \tag{6.5}$$

discretized as follows:

$$\frac{F^{n+1} - F^*}{\Delta t} = \frac{Q^{n+1} + Q^*}{2} \tag{6.6}$$

since coefficient B is time independent.

The variance density spectrum F^{n+1} for a time step (time $t = (n+1).\Delta t$) is then obtained. That operation is then repeated for the next time step and as many times as necessary for covering the simulation period being considered.

6.2 Processing the advection step

The propagation step is solved in TOMAWAC by means of the method of characteristics which is largely used at the LNHE for processing various convection equations (refer for example to [18]). The application of that method to TOMAWAC has a specific feature: the method should be applied to a dimension-4 space in the general case and to a dimension-3 space when there is no current and the depth is constant over time; furthermore the domain in propagation directions is periodic.

It should be reminded that equation (6.3) without source terms is processed in that step, being discretized as follows for a time step $\Delta t p$:

$$\frac{(B.F)^* - (B.F)^n}{\Delta t_p} = [\mathbf{V}.grad(B.F)]^n$$
(6.7)

The convector field V, whose expression was given in Section 4, is not time dependent when there is no tide, just like factor B (refer to paragraph 4). The equation to be processed can be simplified as follows in that case:

$$\frac{B.F * - B.F^n}{\Delta t_p} = \mathbf{V}. \left[grad(B.F) \right]^n \tag{6.8}$$

This is a major advantage, since the characteristics can be traced back only once, at the beginning of the simulation. It is sufficient to store the origin of the characteristic pathlines and to retrieve them whenever the convection step is called. For each quadruplet $(x_Q, y_Q, \theta_Q, fr_Q)$ of the discretized spatial and spectro-angular variables, the characteristic curve is traced back to the time step Dtp and the "arrival" point $(x_P, y_P, q_P fr_P)$, which is called foot of the characteristic pathline, is stored. Actually, the numbers of the discretization elements (triangular elements for the spatial grid and quadrangular elements for the spectro-angular grid) including that foot of the characteristic pathline, as well as the linear interpolation coefficients allowing to obtain the values in that point from the values at the apices of the elements (barycentric coordinates), are kept. Thus, the convection step can be reduced in the form:

$$(B.F) * (x_O, y_O, q_O, fr_O) = (B.F)^n (x_P, y_P, \theta_P, fr_P)$$
(6.9)

That step requires a short computation time since it only consists of an interpolation operation over each time step, once the characteristics have been traced back at the beginning of a computation.

When there is a tide, the principle remains unchanged, but the characteristics should be traced back after every depth and current update.

Such a method has the advantage of being unconditionally stable, enabling to revoke the condition that requires a Courant number below 1 and which is implemented, for example, in the upstream off-centred first-order propagation scheme being used in the WAM-cycle 4 model [61] [36]. The finite element grid generation technique is provided for achieving a locally finer computational grid in order to represent irregular bathymetric features or an irregular coastline. Thanks to the applied propagation scheme, the time step does not necessarily have to be much reduced, so that reasonable computation times can be kept. It should actually be clear that, rather than the propagation step, the source term integration step (particularly the computation

of non-linear interactions) does consume most of the computation time. As regards the numerical schemes in which the propagation step implies a shorter time step when making the grid finer (e.g. as in the case of the WAM model), the overall computation time happens to become much longer because of the source terms and the model becomes less attractive for the practical applications. Owing to the method of characteristic, on the contrary, the TOMAWAC model allows to overcome that restriction and is therefore attractive even for grids with a rather fine spatial resolution.

The method of characteristic, however, has some drawbacks due to the fact that, in the general case, it has a significant level of numerical diffusion and is not conservative.

6.2.1 Representation of diffraction in the propagation step

If diffraction is taken into account, the transfer rates are time dependent and therefore the characteristics must be traced back at each time step, which is computationally expensive. Furthermore, when diffraction is taken into account, the directional propagation rates change significantly quicker in space and a very large number of iterations is required in order to solve the ODEs system when tracing back the characteristics. The system of ODEs becomes even stiffer than in the stationary case and increases the computational time drastically. Tomawac uses the Runge-Kutta method to solve iteratively the ODEs system.

The implementation of the diffraction terms requires the computation of second derivatives (respectively of the variables a and A for the MSE the RMSE cases). For the calculation of derivatives TOMAWAC uses a standard library (BIEF) of linear finite element routines, provided as part of the TELEMAC suite. The second derivative method provided within BIEF is not very accurate when it is used with a linear finite element grid and produces significant noise around the boundaries,

An alternative method for determining the second order derivative, proposed and implemented in TOMAWAC by Kriezi, is a meshfree technique. This method, based on a Radial Point Interpolation technique [39], allows to improve the computation of the second derivative while avoiding higher order finite elements. This method is described in [37].

6.3 Processing the source term integration step

6.3.1 Source term integration numerical scheme

The source and sink terms in the equation of variance density spectrum evolution are integrated using the following scheme, which can be changed by the user from fully explicit to fully implicit:

$$\frac{F^{n+1} - F*}{\Delta t} = (1 - \chi) Q * + \chi Q^{n+1}$$
(6.10)

where the exponent * denotes the values of the variables after the propagation step (but before the source term integration step) and the exponent $^{n+1}$ denotes the values of the variables after the source term integration step. The parameter χ , set by the user through the keyword *IMPLICITATION COEFFICIENT FOR SOURCE TERMS* in the steering file, should lie in the range [0; 1]:

 $\chi = 0$ corresponds to the fully explicit scheme (only the value of source terms Q^* are used). This option is not advised for practical applications.

 $\chi = 0.5$ corresponds to the semi-explicit scheme, used in the WAM-Cycle 4 model [61] [36]. It enables to use fairly long time steps (about 20-30 min in an oceanic environment). This value was the only one considered in earlier versions of TOMAWAC. It is the default value of the keyword.

 $\chi = 1$ corresponds to the fully implicit scheme, as used for instance in a modified version of WAM [28].

Emphasis should be laid on the fact that the source term integration step is local, i.e. it is carried out independently for each point in the 2D spatial grid.

The numerical implementation of the scheme is briefly presented below.

We first define $\Delta F = F^{n+1} - F^*$, the change in the spectrum value. Then the source/sink terms are classified as linear (Q_l) or non-linear terms (Q_{nl}) in the wave spectrum F:

$$Q = Q_l + Q_{nl} \tag{6.11}$$

As regards the source terms that are linear in F, note that: $Q = \beta F$, hence we get:

$$Q_I^{n+1} = \beta^{n+1} F^{n+1} = \beta^{n+1} F^* + \beta^{n+1} \Delta F$$
 (6.12)

As regards the source terms that are non-linear in F, a Taylor's expansion is done keeping only the first-order term:

$$Q_{nl}^{n+1} \approx Q_{nl}^* + \frac{\partial Q_{nl}^*}{\partial F} \Delta F \tag{6.13}$$

 $\frac{\partial Q_{nl}^*}{\partial F}$ is a matrix of differential increments that is broken down into a diagonal component $[\Lambda^*]$ and an extra-diagonal component $[N^*]$:

$$\frac{\partial Q_{nl}^*}{\partial F} = [M^*] = [\Lambda^*] + [N^*] \tag{6.14}$$

Substituting into the expression of Q_{nl}^{n+1} , this yields:

$$Q_{nl}^{n+1} \approx Q_{nl}^* + ([\Lambda *] + [N *]) \Delta F \tag{6.15}$$

Adding the contributions from the linear and nonlinear terms, we obtain:

$$Q^* = \beta * F * + Q_{nl}^* \tag{6.16}$$

$$Q^{n+1} = Q_l^{n+1} + Q_{nl}^{n+1} = \beta^{n+1}F * + \beta^{n+1}\Delta F + Q_{nl}^* + ([\Lambda *] + [N*])\Delta F$$
 (6.17)

The variation of the variance density spectrum due to the source terms is written as:

$$\Delta F = F^{n+1} - F * = \Delta t \left((1 - \chi) Q^{n+1} + \chi Q * \right)$$
 (6.18)

i.e., after substitution of the source term expressions:

$$\Delta F = \Delta t \left\{ (1 - \chi) \left[\beta^* F^* + Q_{nl}^* \right] + \chi \left[\beta^{n+1} F * + \beta^{n+1} \Delta F + Q_{nl}^* + ([\Lambda *] + [N *]) \Delta F \right] \right\} \quad (6.19)$$

$$\Delta F\left[1-\chi\Delta t\left(\beta^{n+1}+\left(\left[\Lambda*\right]+\left[N*\right]\right)\right)\right]=\Delta t\left(\left[(1-\chi)\beta*+\chi\beta^{n+1}\right]F^*+Q_{nl}^*\right) \tag{6.20}$$

The matrix between brackets in the left-hand member of the latter equation cannot be easily inverted in the general case. The designers of the WAM model [61], however, demonstrated that the diagonal portion $[L^*]$ usually prevails over the extra-diagonal portion $[N^*]$. Relying on comparative tests, they conclude that the extra-diagonal portion can be ignored in favour of the diagonal portion, even with time steps of 20 min or so. Due to that simplification, the inversion is much easier and we finally obtain:

$$\Delta F \approx \Delta t \frac{\left((1 - \chi)\beta * + \chi \beta^{n+1} \right) F * + Q_{nl}^*}{1 - \chi \Delta t \left(\beta^{n+1} + \Lambda * \right)}$$

$$(6.21)$$

Source/sink terms	Linear or	Remarks	Type of contribution	Type of contribution
Source/sink terms	non-Linear	Kemarks	to Q_DER	to Q_{TOT}
	Linear	β		
Wind input	or	depends	$oldsymbol{eta}^{n+1}$	$((1-\boldsymbol{\chi})\boldsymbol{\beta}^*$
	quasi-linear	on time		$+\chi\beta^{n+1})F^*$ Λ^*F^*
Whitecapping	Quasi-linear	Slightly	Λ^*	Λ^*F^*
		non-linear		
Bottom friction	linear	β does not	β	$\beta F*$
		depend on time		
Non-linear transfers				
between frequency	non-lin.		Λ^*	Q_{nl4}^*
quadruplets				
Bathymetric		b does not		
breaking	linear	depend on time	β	$eta F^*$
		$\beta^{n+1} = \beta^n = \beta$		
Non-linear transfers	non-lin		Λ^*	Q_{tr}^*
between triads				

Table 6.1: Contributions of the different source/sink terms implemented in TOMAWAC.

For the sake of convenience, that expression is rewritten as:

$$\Delta F = \frac{\Delta t. Q_{TOT}}{1 - \chi \Delta t Q_{DER}} \tag{6.22}$$

where: $Q_{TOT} = ((1 - \chi)\beta^* + \chi\beta^{n+1})F^* + Q_{nl}^*$ denotes a total source term and $Q_{DER} = \beta^{n+1} + \Lambda^*$ denotes a source term derived with respect to F.

The contributions of the various source terms implemented in TOMAWAC and described in paragraph 4.3 are schematically illustrated in the Table 6.1

The source term integration time step may be different from the propagation time step in TOMAWAC, but it should be a sub-multiple of it. Thus, several source term integration time sub-steps per propagation time step can be defined. That option is governed by the keyword *NUMBER OF ITERATIONS FOR THE SOURCE TERMS* in the steering file. The default value of that parameter is set to 1.

It was found experimentally that the depth-induced breaking source term, which is sometimes very strong, can still be overestimated if the time step that was selected for the source term integration is too long. In order to avoid that, TOMAWAC gives an opportunity to make a number of time sub-steps that are specific to that source term. These time sub-steps are in a geometric progression. In order to limit that number of time sub-step, TOMAWAC first clips the wave height by setting a maximum H_{m0}/d ratio d being the depth) to 1.

Subsequently, a Euler's explicit scheme is used at each time step:

$$\frac{F^{n+1} - F^*}{\Delta t_2} = Q^* \text{ i.e. } \Delta F = \Delta t_2 Q^*$$
 (6.23)

The H_{m0}/d ratio can be modified through the keyword MAXIMUM VALUE OF THE RATIO HM0 ON D (however, this it not advisable). The number of time sub-steps is specified through the keyword NUMBER OF BREAKING TIME STEPS. The geometric ratio is given by the keyword COEFFICIENT OF THE TIME SUB-INCREMENTS FOR BREAKING.

6.3.2 Monitoring the growth of the wave spectrum

In order to limit the possible risks of numerical instabilities related to the source term integration, TOMAWAC is provided with various options for limiting the growth of the directional spectrum per source term integration time step. The choice of option is done by the user in the steering file through the keyword *WAVE GROWTH LIMITER*, which can bet set to 0, 1 (default value) or 2:

0 : no limiter is applied to the computed ΔF from the various source/sink terms.

1: the limiter is directly inspired by the criterion proposed by the WAM group [61]. The absolute variation of the variance density spectrum as it was computed by the scheme in paragraph 6.3.1. should remain lower than a fraction of an equilibrium spectrum ΔF_{lim} :

$$\Delta F_{\text{lim}} = 6.4 \ 10^{-7} \, g^2 \frac{\Delta t}{1200} f^{-5} \tag{6.24}$$

This is presently the default value in TOMAWAC, as it was the only possibility in earlier versions of the code.

2: the limiter is computed with an updated expression, proposed by Hersbach et al. [28]:

$$\Delta F_{lim} = 3.0 \ 10^{-7} g \max(u*, g \tilde{f}_{PM}/f) f^{-4} f_{NF} \Delta t$$
 (6.25)

where u^* is the friction velocity at the water surface and $\tilde{f}_{PM} = 5.6 \ 10^{-3}$ is the dimensionless Pierson–Moskowitz peak frequency.

This option should now be preferably used for practical applications.

3: In this option we use the same formula as for 2, but we take the mean of windsea frequencies f_{meanWS} (see 6.5) instead of f_{NF}

6.4 Processing the boundaries – Boundary conditions

6.4.1 Spatial grid:

Two types of boundary conditions are considered in TOMAWAC for the finite element spatial grid:

- The former corresponds to <u>a free boundary condition</u>, i.e. that absorbs the whole wave energy. It may be a sea boundary, hence it is assumed that the waves propagate beyond the domain and nothing enters it. It may be a solid boundary, hence it is assumed that the coast absorbs completely the wave energy (no reflection).
- The latter corresponds to a <u>prescribed value boundary condition</u>. The whole wave spectrum is then prescribed at each point along that boundary and for each step. Energy enters into the computational domain.

6.4.2 Spectro-angular grid:

As regards the propagation directions, the grid generation is periodical over the range $[0;360^{\circ}]$: hence there are no directional boundary conditions.

As regards the wave frequencies that are discretized, the minimum and maximum frequency markers are considered as "open boundary limits", where the energy can be transferred to lower or higher frequencies, exiting the discretized frequency range.

6.5 A set of different models: BAJ modelisation.

In [7], the authors proposed a new formulation in terms of mean wave parameters to emphasis the high frequency part in interaction between windsea and swell. That formula has been implemented in Tomawac. It consist firstly in choosing a set of formulation in term of simulation. This modelisation includes WAM 4 for Janssen wind modelisation, (i.e. option 1 in 4.3.2), Komen whitecapping dissipation (i.e. option 1 in 4.3.3,with $C_{dis} = 2.1$ and $\delta = 0.4$). In the initial paper, BAJ formulation also includes Discrete Integration Approximation for non linear resonant quadruplet interactions, (i.e. option 1 in 4.3.6), but since TELEMAC-V 8.2, it doesn't include automatically this modele, which allows user to use BAJ with another formulation for Non Linear Interaction.

In those formulations the mean wave number \sqrt{k} and the mean angular frequency σ are defined using weighted spectral integrals that put more emphasis on the high frequencies:

$$\sqrt{\langle k \rangle} = \frac{\int d\mathbf{k} \sqrt{\mathbf{k}} F(\mathbf{k})}{\int d\mathbf{k} F(\mathbf{k})}$$
$$\langle \sigma \rangle = \frac{\int d\mathbf{k} \sigma F(\mathbf{k})}{\int d\mathbf{k} F(\mathbf{k})}$$

Another point is the frequency cut off to calculate the discrete integration. This frequency is no longer a function of the mean frequency but the mean frequency of the windsea only, denoted by f_{meanWS} , where only frequency with $S_{input} > 0$ or $\frac{28}{c}u_*cos(\Delta\theta) \ge 1$ are considered. Where S_{input} is the wind input source term, c is the wave phase speed, u_* is the friction velocity and $\Delta\theta$ is the difference between the wind direction and the wave propagation direction. Then only frequencies such that:

$$f_{min} \le f \le min(2.5 f_{meanWS}, f_{max})$$

will be considered for the integration. Frequencies above will follow a f^{-5} shape. The consequence of that choice is the modification of the charnock constant in the wind generation (0.0095).

The last point concern the wave growth limiter that will use the mean frequency of the windsea (option 3 in 6.3.2).

7. Inputs-outputs

7.1 Preliminary remark

During a computation, the TOMAWAC software uses a number of files, some of which are optional, as inputs and outputs.

The input files are:

- The steering or CAS file (mandatory),
- The mesh or geometry file (mandatory),
- The boundary conditions or CONLIM file (mandatory),
- The seabed, bottom or bathymetry file (optional),
- The FORTRAN or PRINCI file (optional),
- The currents file (optional),
- The winds file (optional),
- The imposed spectra on the boundary (optional),
- The coordinates of the nodes for which a spectra will be imposed (optional),
- The previous computation file (optional),
- The binary user file (optional),
- The formatted user file (optional),
- The coordinates of the nodes for which spectra will be written (mandatory if a punctual or a spectra file is defined),
- The time series coordinates file which contains coordinates and periods of time where time series are extracted and written in the time series file.

The output files are:

• The 2D results or grid file (mandatory),

7.2 The files 55

- The punctual results or spectra file (optional),
- The next computation file (optional),
- The listing printout (either on the display screen or in the file, see in A.2),
- The binary user file (optional),
- The formatted user file (optional),
- The time series file which contains time series at locations defined in the time series coordinates file.

7.2 The files

7.2.1 The steering (or CAS) file

The steering file name is specified in the steering file through the keyword: *STEERING FILE*. It is a text file created by means of a text editor. In a way, it serves as the computation control panel. It includes a set of keywords to which values are assigned. If a keyword does not appear in this file, then TOMAWAC will assign to it the default value as defined in the dictionary file (refer to the reference manual). If such a default value is not defined in the dictionary, then the computation will come to a halt and display an error message. For instance, the command *NUMBER OF DIRECTIONS* = 12 is for specifying that the direction spectrum of wave action or its moments will be discretised over 12 propagation directions.

TOMAWAC reads the steering file at the beginning of the computation.

Both dictionary file and steering file are read by the so-called DAMOCLES utility which is included in TOMAWAC. The syntactic rules of DAMOCLES should then be observed upon the creation of the steering file These rules are described here below.

The write rules are as follows:

- The keywords can be of the Integer, Real, Logical or Character format type.
- The keyword sequence order in the steering file is of no importance.
- Each line has a maximum of 72 characters. However, as many linefeeds as one wants are allowed provided that the keyword name does not run from one line to the next.
- For the table-like keywords, the successive values are separated by a semi-colon. A number of values equal to the table dimension should not necessarily be given; in such a case, DAMOCLES returns the number of values being read. For example:

ABSCISSAE OF SPECTRUM PRINTOUT POINTS = 1.2;3.4 (that keyword is declared as a 19-valued table)

• The symbols ":" or "=" are indiscriminately used to separate a keyword from its value. They can be either preceded or followed with any number of blanks. The value itself may appear on the next line. For example:

```
NUMBER OF DIRECTIONS = 12
or
NUMBER OF DIRECTIONS: 12
or else
NUMBER OF DIRECTIONS =
12
```

- The characters occurring between a pair of "/" on one line are regarded as comments. Likewise, the characters occurring between a "/" and a the end of a line are regarded as comments. For example: TYPE OF BOUNDARY DIRECTIONAL SPECTRUM = 1 / Jonswap spectrum
- A whole line beginning with a "/" in the first column is regarded as a comment, even though another / occurs on the line. For example: / The geometry file is . /maillage/geo
- Integer writing: Do not exceed the maximum size being allowed by the machine (in a machine with 32 bit architecture, the values range from -2 147 483 647 to +2 147 483 648. Do not enter a blank between the sign (optional for the + sign) and the number. A dot at the end of the number is tolerated.
- Real writing: A dot or a comma is allowed as a decimal point, as well as the FORTRAN E and D formats (1.E-3 0.001 0,001 1.D-3 denote the same value).
- Logical value writing: The values 1, YES, OUI, .TRUE., TRUE, VRAI on the one hand, and 0, NON, NO, .FALSE., FALSE, FAUX on the other hand are allowed.
- Character string writing: Those strings including blanks or reserved symbols ("/",":", "=", "&") should be put in single quotes ('). The value of a character keyword may include up to 144 characters. As in FORTRAN, the quotes occurring within a string should be doubled. A string may neither begin nor end with a blank. For example: TITLE = 'HOULE D'OUEST'

In addition to the keywords, a number of directives or metacommands that are interpreted during the sequential readout of the steering file may be used as well:

- The &FIN command indicates the end of file (even though the file is not completed). Thus, some keywords can be disabled simply by placing them behind that command for easily making it possible to enable them again subsequently.
- The &ETA command prints the list of keywords and the relevant values at the time when DAMOCLES meets that command. This display will occur at the beginning of listing printout.
- The &LIS command prints the list of keywords. This display will occur at the beginning of listing printout.
- The &IND command prints the detailed list of keywords. This display will occur at the beginning of listing printout.
- The &STO command causes the interruption of the program, the computation does not go on.

7.2.2 The geometry file

The geometry file name is specified in the steering file through the keyword: *GEOMETRY FILE*. It is either a SERAFIN-formatted or a MED-formatted binary file: it can be read by FUDAA PRE-PRO or RUBENS (for SERAFIN) or Salome (for med) and it can be created by the STB-TEL module from the file(s) as produced by the mesh generator. The format structure is described in A.4

This file includes the complete information about the horizontal mesh, i.e. the number of mesh points (variable NPOIN2), the number of elements (variable NELEM2), the X and Y vectors

7.2 The files 57

containing the co-ordinates of all the points and, lastly, the IKLE2 vector containing the connectivity table.

Furthermore, this file may also include bathymetry information in each point of the mesh, provided that the interpolation of the bathymetry was carried out during the execution of the STB-TEL module or during the generation of the mesh.

TOMAWAC reproduces the information regarding the geometry at the beginning of the 2D results. Any computation results file can then be used as a geometry file when one wants to perform a further simulation on the same mesh.

7.2.3 The boundary conditions file.

The boundary conditions file name is specified in the steering file through the keyword: *BOUNDARY CONDITIONS FILE*.

It is a formatted file that can be created automatically by STBTEL and can be modified by means of a text editor. If the geometry file is a serafin, Each line in this file is assigned to one point of the boundary and listed in sequential order in terms of the boundary node numbers. The numbering of the boundary points first delineates the domain contour in the counterclockwise direction, then the islands in the clockwise direction. This file is described in detail in 8.5.1 If the geometry file format is med, then each line correspond to a group of segments defined in the geometry file.

7.2.4 The currents file

According to its type – binary or formatted- the currents file name is specified in the steering file through the keywords: *BINARY CURRENTS FILE* and *FORMATTED CURRENTS FILE*. It is the file from which TOMAWAC reads the current field components. The current field may be either stationary or non-stationary. The current field will be non-stationary when the keyword *CONSIDERATION OF TIDE* is set to TRUE. When the current field is stationary, the keyword *CONSIDERATION OF A STATIONARY CURRENT* should be set to TRUE. By default, both keywords will be set to FALSE. When both are set to TRUE, the keywords will be inconsistent, and the program will halt.

Several commonly used formats can be read. This selection is made through the integer keyword *CURRENTS FILE FORMAT*. It can be set to a value from 3 to 4.

- The format is 3: it is a TELEMAC result file of the SERAFIN or MED standard. It is a binary file the name and its name be assigned to the keyword: *BINARY CURRENTS FILE*. If the current is assumed to be stationary, then the additional keyword *TIME IN-CREMENT NUMBER IN TELEMAC FILE* should be used in order to find the time step number related to the desired record. TELEMAC data other than the current components e.g. water levels, can also be read by means of this format (refer to 8.2.5).
- The format is 4: data written in a different format can be read provided that the user supplies the relevant subroutine in the relevant FORTRAN file (see 8.2.3 and 8.2.6).

7.2.5 The tidal water level file

According to its type – binary or formatted- the tidal water level file name is specified in the steering file through the keywords: *BINARY TIDAL WATER LEVEL FILE* or *FORMATTED TIDAL WATER LEVEL FILE*.

This is the file from which TOMAWAC reads the tidal water level being referred to the *INITIAL STILL WATER LEVEL*. Several commonly used formats can be read. This selection is made by means of the integer keyword *TIDAL WATER LEVEL FILE FORMAT*. It can be set to a value from 3 to 4.

- The format is 3: it is a TELEMACTELEMAC result file of the SERAFINSERAFIN standard. It is a binary file and its name should be assigned to the keyword: *BINARY TIDAL WATER LEVEL FILE*.
- The format is 4: data written in a different format can be read provided that the user supplies the relevant subroutine in the relevant FORTRAN file (see in 8.2.6).

7.2.6 The winds file

According to its type – binary or formatted- the wind file name is specified in the steering file through the keywords: *BINARY WINDS FILE* or *FORMATTED WINDS FILE*.

This is the file from which TOMAWAC reads the information about the wind fields. As in the case of the current, several read formats are allowed. The integer keyword *WINDS FILE FORMAT* can be set to values from 1 to 4.

- The format is 3: it is a TELEMAC result file of the SERAFIN standard. It is a binary file and its name should be assigned to the keyword: *BINARY WINDS FILE*. If the wind is assumed to be stationary, then the additional keyword *TIME STEPTIME STEP NUMBER IN TELEMAC FILE* should be used in order to find the time step number related to the desired record.
- The format is 4: data written in a different format can be read provided that the user supplies the relevant subroutine in the relevant FORTRAN file (see in 8.2.4).

7.2.7 The imposed spectra file and the file containing the associated coordinates

It is possible to impose spectra on the open boundary of the mesh. To do so, the boundary condition for those nodes need to follow the format 5 4 4 4, see section 8.5.1. There are then two files that need to be given:

- *IMPOSED SPECTRA FILE*: which is a binary file, such as outputted in the punctual or spectrum results file, see section 7.2.13.
- FILE WITH COORDINATES OF SPECTRA TO IMPOSE: which is an ascii file with the coordinates associated with each spectrum in the IMPOSED SPECTRA FILE.

Since the *IMPOSED SPECTRA FILE* follows the format of the TELEMAC SYSTEM results, the format can be given using the keyword *IMPOSED SPECTRA FILE FORMAT*. In addition, since imposed spectra can change in time, the *TIME UNIT OF IMPOSED SPECTRA FILE* and *TIME SHIFT OF IMPOSED SPECTRA FILE* can be modified.

Finally, the *FILE WITH COORDINATES OF SPECTRA TO IMPOSE* follows the format of DAT outputs of the module SMESH in SALOME. For a list of nodes, it follows this structure:

```
<Number of Points> 0
<Point Id> <X> <Y> <Z>
...
```

7.2.8 The area definition file.

When a user is using vegetation or porous medi he has the opportunity to define areas that will contains vegetation or porous media through a list of polygons. Those polygones are defined through the *FILE WITH DEFINITION OF POLYGONS*. This file is an i2s format file. The first line contains the number of polygons. The second one, the numbers of points of each polygon and then each line contains the coordinates of each point.

7.2 The files 59

7.2.9 The previous computation file

This previous computation file name is specified in the steering file through the character keyword: *PREVIOUS COMPUTATION FILE*.

If a *NEXT COMPUTATION* is done, TOMAWAC fetches this file in order to initialize the directional spectrum of wave action at every point. This file's format, which is specific to TOMAWAC, is described in A.3. It is a binary file.

7.2.10 The global results file

The global results file name is specified in the steering file through the keyword: *GLOBAL RESULTS FILE*.

This file is created when a *GLOBAL OUTPUT AT THE END* is requested. It saves the wave action density directional spectrum at every point in the last time step. This file format is described in A.3.

7.2.11 The 2D results file

The 2D results file name is specified in the steering file through the character keyword: 2D RESULTS FILE.

This is the file into which TOMAWAC writes the results of the 2-dimensional variables during the computation. It is a binary file of the TELEMAC-S tandard (SERAFIN or MED). The data contained in it are in the following order:

- all the data about the mesh geometry;
- the names of the variables being stored;
- for each time step, the time and the values of the variables are given for each point of the 2D mesh.

Its content varies according to the values of the following keywords:

- NUMBER OF FIRST ITERATION FOR GRAPHICS PRINTOUTS: provided for determining from which time step will the data storage desirably begin, so that the file size will not be too large.
- *PERIOD FOR GRAPHICS PRINTOUTS*: sets the period, as a number of propagation time increments, of printouts so that the file size will not be too large.
- VARIABLES FOR 2D GRAPHICS PRINTOUTS: provided for specifying the list of variables to be stored into the 2D results file. Each variable is identified by 2, 3 or 4 letters (refer to 7.1 that lists the available variables).

Table 7.1: List of 2D results variables

Identifier	Definition
M0	Total variance
HM0	Spectral significant wave height
DMOY	Mean wave direction
SPD	Mean directional spreading
ZF	Sea bottom level

Table 7.1: (continued)

Identifier	Definition	
WD	Water depth	
UX	Current along X	
UY	Current along Y	
VX	Wind along X	
VY	Wind along Y	
FX	Driving force along X	
FY	Driving force along Y	
SXX	Radiation stress along xx	
SYY	Radiation stress along yy	
SXY	Radiation stress along xy	
UWB	Bottom celerity	
PRI	Private table	
FMOY	Mean frequency FMOY	
FM01	Mean frequency FM01	
FM02	Mean frequency FM02	
FPD	Discrete peak frequency	
FPR5	Peak frequency by Read method of order 5	
FPR8	Peak frequency by Read method of order 8	
US	Surface friction velocity u*	
CD	Surface drag coefficient CD	
Z 0	Surface roughness length Z0	
WS	Surface wave stress	
TMOY	Mean period Tmoy	
TM01	Mean period Tm01	
TM02	Mean period Tm02	
TPD	Discrete peak period	
TPR5	Peak period by Read method of order 5	
TPR8	Peak period by Read method of order 8	
POW	Wave power (per meter along wave crest)	
BETA	Breaking waves rate	
BETAWC	White Capping rate	
SRE	Surface Roller Energy	
DBR	Breaker Dissipation	
DSR	Roller Dissipation	

For instance, if the significant wave heights, the water depths and the average wave propagation directions are desired,

VARIABLES FOR 2D GRAPHICS PRINTOUTS = HM0,WD,DMOY must be entered in the steering file.

7.2.12 The time series file

The time series file is a file (in SERAFIN or MED format), in which time series are written for certain points in the domain of the simulation. This feature has been introduced since release 8.5. In this way, one can obtain results with a high time frequency, without using a large amount of disk space. The time series file module interpolates data (using linear interpolation even if

7.2 The files 61

using quasi-bubble or quadratic elements) to the required output coordinates.

• TIME SERIES FILE The name of the output file to write. The time and coordinates correspond to those given in TIME SERIES COORDINATES FILE,

- TIME SERIES FILE FORMAT The file format of TIME SERIES FILE (single or double precision SERAFIN, or MED, default = SERAFIN),
- TIME SERIES COORDINATES FILE The file which contains the output times and output coordinates for TIME SERIES FILE. The format is described next.

The TIME SERIES COORDINATES FILE has following format:

- On the first line, there are two numbers indicating:
 - The number of time periods for which data is written to the TIME SERIES FILE,
 - The number of coordinates for which output is generated.
- On the next lines, for each output period, there are three numbers indicating:
 - Start of the output period (in seconds since the start of the computation),
 - End of the output period (in seconds since the start of the computation),
 - Output time interval (in seconds).
- For each coordinate, three numbers and a string, containing respectively:
 - x-coordinate of the output point,
 - y-coordinate of the output point,

2 3

- Unique ID of each station (in order to easily encounter the output data in the output files). This ID will be written to the IKLE array in the output file, which has a size [NPOINTS × 1]. Note that the time series file consists of a list of points. There is no mesh information in a time series file. Therefore, the length of the IKLE array is the same as the number of points in the time series file,
- The name of the station (for better readability of the coordinate file).
- Anything after the list of coordinates is considered comment and ignored by TELEMAC.

An example of a TIME SERIES COORDINATES FILE is given below.

```
0 100 10
500 1000 20
250 175     01 output_point_01
500 175     02 output_point_01
750 175     03 output_point_02

Description of the file
1st line: number of periods and number of points
2nd line: period 1: start time, end time and interval (in seconds)
3rd line: period 2: start time, end time and interval (in seconds)
4th-6th line: output points; x and y coordinates, unique ID, and station name
```

7.2.13 The punctual or spectrum results file

This file's name is specified in the steering file through the character keyword: *PUNCTUAL RESULTS FILE*.

This is the file into which the directional spectra of wave action at some previously specified points are stored by TOMAWAC during the computation. To define those points, a list of coordinates needs to be given, and the spectra of the closest node will be outputted. This means that there is no spatial interpolation done. There are two methods that can be used to define those points:

1. Through keywords in the steering file:

The *X* and *Y* coordinates are given through the following keywords: *ABSCISSAE OF SPECTRUM PRINTOUT POINTS* and *ORDINATES OF SPECTRUM PRINTOUT POINTS*. These are chart keywords, which means that each value is seperated with semicolon, and the number of printout points is equal to the size of the array given. For example:

```
Keywords to write a spectra using method 1:

/----/
/ WRITING SPECTRA
/----/
PUNCTUAL RESULTS FILE = <FileName>
ABSCISSAE OF SPECTRUM PRINTOUT POINTS = <X1;X2;X3;X4;X5>
ORDINATES OF SPECTRUM PRINTOUT POINTS = <Y1;Y2;Y3;Y4;Y5>
```

2. Through an external ASCII file:

The *X* and *Y* can also be defined in an external ASCII file which follows the format of DAT outputs of the module SMESH in SALOME. For a list of nodes, it follows this structure:

```
<Number of Points> 0
<Point Id> <X< <Y> <Z>
...
```

The name of the file is referenced using the *FILE WITH COORDINATES OF SPECTRA TO WRITE* keyword. For example:

Warning:

If both methods are used in the same steering file, the simulation will be terminated and an error message will be printed.

7.2 The files 63

This result file is a SERAFIN formatted file. It first includes all the data about the spectral mesh geometry, then the names-codes of displayed points. This name-code is of the type: Fa_PT2Db , where a denotes the point's sequence order number within the list of coordinates and b denotes the number of the closest 2D point to the specified position. Subsequently, for each graphic printout, it contains the time and the value of the directional spectrum of wave action for each pair (direction, frequency) in the spectral mesh.

The keywords *PERIOD FOR GRAPHICS PRINTOUTS* and *NUMBER OF FIRST ITERATION FOR GRAPHICS PRINTOUTS* are shared by the two results files; thus, the printouts are synchronous for either file.

Finally, there is another KEYWORD to store punctual results, it is called *1D SPECTRA RE-SULTS FILE*. The difference with the previous one is that the spectral energy is summed over directions and that the file is stored in *ASCII* format. The points where information are stored are defined by the same variables.

7.2.14 The printout listing

This file contains all the messages as generated by TOMAWAC during the computation. It is the main report of a TOMAWAC run. Its content depends on the value of the following keyword:

• *PERIOD FOR LISTING PRINTOUTS*: this sets the time between two time steps of message transmission. This value is given in terms of the number of iterations. For example, the following sequence:

TIME INCREMENT = 30.
PERIOD FOR LISTING PRINTOUTS= 2

will result in a print in the output listing every 60 seconds of simulation.

7.2.15 The User FORTRAN file

This User FORTRAN file name is specified in the steering file through the character keyword: *FORTRAN FILE*. Notice that if the user gives a directory for that keyword, all the fortran file from this directory will be used.

The FORTRAN contains all the user-modified TOMAWAC subroutines as well as the specifically developed routines for that computation.

This file is compiled and linked during run time in order to generate the executable being used for the simulation.

7.2.16 The auxiliary files

Other input/output files may be used by TOMAWAC.

- A binary data or results file: its name is specified through the character keyword *BINARY FILE 1* (Channel unit No. 24).
- A formatted data or results file: its name is specified through the character keyword *FORMATTED FILE 1* (Channel unit No. 26).

These files can be used either for supplying data to the program or for allowing data to be processed that are not available in the standard results files; obviously, the user must manage the read and write operations of these files within the FORTRAN program.

7.2.17 The dictionary file

This dictionary file contains all the information about the keywords (French/English name, default values, type). This file can be viewed in a text editor by the user, but it must not be modified in any way.

7.2.18 The libraries

At the beginning of a computation, the main user-written FORTRAN routine is compiled, then linked in order to generate the executable program that is subsequently run.

The following libraries are used during the link editing operation:

- TOMAWAC library: contains the specific subroutines of the TOMAWAC computation model.
- telemac libraries: contains the specific subroutines of the TELEMAC-2D or TELEMAC-3D computation model.
- utils library: contains a number of utility subroutines, such as the partitioning and parallel library.
- damocles library: contains the subroutines handling the steering file reading.
- hermes library: contains the subroutines that permits to read and write data according to the file format chosen.
- gaia library: contains the subroutines that permits to simulate sediment transport
- api library: contains the subroutines that permits to the modulus to communicate
- BIEF library: contains the computation modules related to the finite element-typed operations (operations on both matrixes and vectors). This library is shared by all the simulation models as developed by the LNHE within the TELEMAC structure (BIEF means "BIbliothèque d'Eléments Finis", i.e. Finite Element Library).

7.3 Files standard

Almost all files that were in Serafin format in previous versions of TOMAWAC, have been given a key-word for the file format.

If the name of the file is: "GEOMETRY FILE" ("FICHIER DE GEOMETRIE"), the new keyword will be: "GEOMETRY FILE FORMAT" ("FORMAT DU FICHIER DE GEOMETRIE"). This format is given in 8 characters. Three choices are possible so far:

- 'SERAFIN '(do not forget the space at the end): it is the default standard within the TELEMAC processing chain. The format is recognized by the FUDAA PRE-PRO graphics post-processor. The RUBENS graphics post-processor reads the SERAFIN format as well, but it won't be developed anymore and it is bound to disappear. The SERAFIN file format is described in detail in A.3.
- 'SERAFIND': Serafin format, but with double precision. Can be used for a more accurate "computation continued" or for more accurate validations. Neither FUDAA PRE-PRO nor Rubens can read this format.

• 'MED ': this is an EDF-CEA format used in the Salomé platform, that enables to use the post-processors of this platform. It is based on hdf5. This new format is not activated if you use the default subroutine med.f provided, which is mostly void. If you take instead the file med.edf and rename it med.f, med formats will be available, but two additional libraries are necessary to use this format and have to be specified in the systel.ini file. Full instructions will be given in further releases, this is so far for internal use at EDF.

A new file structure has been added to library BIEF for simplifying the opening/closing and reading/writing operations with these file formats, as well as for simplifying the coupling between programmes,. The description of this file structure and of the operations on those files are given in A.4

As specified in section 7.2.7, a fourth binary format exists, which is specific to TOMAWAC and is used only for saving the results when they are used to initialize a next computation. This binary file format cannot be read by the RUBENS post-processor, or by FUDAA PRE-PRO graphics post-processor.

7.4 Bathymetry data

The bathymetry information can be supplied to TOMAWAC at two levels:

- Directly in the geometry/mesh file by a bathymetry value being assigned to each node in the mesh. In this case the bathymetry data have been processed previously, running the STBTEL module or mesh generator. For example, STBTEL reads the information from one or more bottom topography files (up to 5 files) and performs an interpolation at every point within the domain;
- In the form of an irregular pattern of spot heights without any necessary relation to the mesh nodes, during the TOMAWAC computation. The interpolation is then performed directly by TOMAWAC with the same algorithm as used by STBTEL. The bathymetry file name is given by the character keyword *BOTTOM TOPOGRAPHY FILE*. Unlike STBTEL, TOMAWAC only handles one bottom topography file. The file can be in SINUSX format or can consist of three columns X,Y,Z.

TOMAWAC also provides an opportunity to carry out a smoothing of the bathymetry in order to get a more consistent geometry. The smoothing algorithm can be iterated several times in order to achieve more or less extensive smoothing. The number of iterations is set using the keyword *BOTTOM SMOOTHINGS* and is carried out within the USER_TOM_CORFON subroutine. This keyword's default value is 0. (also refer to the programming of the CORFON user subroutine in 8.6.1).

NOTE: the bathymetry data should preferably be supplied to TOMAWAC in the form of water depth and not of water height. If necessary, a conversion can be performed in the USER_TOM_CORFON subroutine.

8. Controlling the simulation

8.1 General parameterisation

The general parameterisation of the computation is controlled using the steering file.

8.1.1 Computation title

The computation case title is specified by the keyword TITLE

8.1.2 Computation time

The time data are prescribed using the two keywords *TIME step* (real) and *NUMBER OF TIME INCREMENTS* (integer). The former sets the elapsed time, in seconds, between two consecutive computation instants (but not necessarily two outputs in the results file). The number of time steps is for setting the overall computation time (which is obviously equal to the time increment value multiplied by the number of time steps).

An additional keyword also refers to time. This is the *DATE OF COMPUTATION BEGIN-NING*, which is used to identify the time in relation to the date/times written in the *WINDS FILE* (refer to A.3, WAM-typed format). The convention adopted for writing the *DATE OF COMPUTATION BEGINNING* is yyyymmddhhmm. For instance, 190311051110 corresponds to November 5th, 1903, at 11.10 AM.

Note that when a computation is resumed, the initial time of the new computation corresponds to the last time increment in the previous computation (i.e. the computation is not resumed at time zero).

8.1.3 Spectral discretisation

The spectral discretisation is defined by the following 5 keywords:

- NUMBER OF DIRECTIONS,
- NUMBER OF FREQUENCIES,
- MINIMUM FREQUENCY,
- FREQUENTIAL RATIO,
- SPECTRUM TAIL FACTOR.

It should be reminded that the directions are evenly distributed from 0 to 360 degrees. Two conventions can be chosen by the user for expressing the wave propagation by means of the keyword *TRIGONOMETRICAL CONVENTION* (logical). The trigonometrical convention locates the wave propagation from the positive X axis and the direction of rotation is in a counterclockwise direction. The default convention is the nautical convention (*TRIGONOMETRICAL CONVENTION* = NO) that locates the propagation direction in relation to the true "North", i.e. the Y axis. The selected direction of rotation is a clockwise direction. The direction will then correspond to the "heading" in the sense of the navigational maps, i.e. the direction the waves are propagating towards.

The frequencies are distributed geometrically in accordance with the following relation:

$$f_k = f_0 r^{k-1}$$
 $(k = 1, NF)$

where f_0 is the MINIMUM FREQUENCY r is the FREQUENTIAL RATIO NF is the NUMBER OF FREQUENCIES

We can choose how to treat the high frequencies. If we want to impose the frequencies higher than the frequency cut off to decay with a law of the type f^{-n} , then we set the *OPTION FOR DIAGNOSTIC TAIL* to 1, and the *SPECTRUM TAIL FACTOR* corresponds to the value of n. The frequency cut off is the max of 4 Pierson-Moskowitz frequency and 2.5 mean frequency. The Pierson-Moskowitz frequency is defined as $\frac{g}{2\pi 28u^*}$ where u^* is the friction wind velocity.

8.1.4 Release

When generating the executable, the release number of libraries being used for editing the links is indirectly provided by the keyword TOMAWAC *RELEASE NUMBER*. By default, TOMAWAC release 7.1 utilises the 7.1 releases of the TELEMAC system libraries.

8.1.5 Environment

When a vector computer is used, the CPU vector length used in the forced vectorisation technique can be specified by means of the keyword *VECTOR LENGTH*. The default value is 1 and is appropriate for scalar machines such as the present workstations. If a value of 1 is used on a vector machine, then the advantage of the vectorisation loops (although they are few in TOMAWAC) is lost.

8.2 Computation options

8.2.1 Co-ordinate system

Cartesian co-ordinates (expressed in meters) are used by default. For domains of a large extent, working in spherical co-ordinates may become necessary. The value of the logical keyword *SPHERICAL COORDINATES* should then be set to "TRUE". The co-ordinates are then expressed in degrees.

8.2.2 Finite depth

In nearshore areas, wave conditions will be influenced by the water depth and therefore the bottom effect can no longer be ignored. This is the default case in TOMAWAC: the keyword *INFINITE DEPTH* is set to "FALSE". In situations where depths effects are to be explicitly ignored the keyword *INFINITE DEPTH* should be set to "TRUE".

8.2.3 Taking a stationary current into account

A stationary current can be taken into account in the TOMAWAC. The relevant logical keyword is *CONSIDERATION OF A STATIONARY CURRENT*. The current affects mainly the convection step.

The current can be specified in various ways:

When the current is either constant over the domain or can be described analytically, the ANA-COS.f subroutine can be included in the FORTRAN file and modified accordingly. In this subroutine the UC and VC are NPOIN2-sized (number of points in the horizontal mesh) vectors and correspond to the components along the X and Y axes of the current, respectively. This is how the current is specified when the keyword *FORMATTED CURRENTS FILE* or *BINARY CURRENTS FILE* is not specified, whereas the keyword *CONSIDERATION OF A STATION-ARY CURRENT* is "TRUE".

TOMAWAC can also take into account a current provided in a binary or formatted file. The keyword *BINARY CURRENTS FILE* or *FORMATTED CURRENTS FILE* should then be given a value (the name of the file). Three different formats are available to read this data. The value corresponds to the keyword *CURRENTS FILE FORMAT* (see 7.2.4). When the currents file is taken from TELEMAC-2D, then an additional keyword should be specified, namely the *TIME INCREMENT NUMBER IN TELEMAC FILE*. This locates the desired record.

If the predefined formats cannot be used, the USER_CURRENT.f subroutine can be included in the FORTRAN and modified accordingly, specifying the format 4 for the INDIC FORTRAN variable in the CAS file. The current data are read from the file and are interpolated onto the nodes of the computation mesh.

8.2.4 Taking a wind into account

Consideration of a wind is specified by the logical keyword *CONSIDERATION OF A WIND*. The wind may be either stationary or variable in time and is specified by means of the logical keyword *STATIONARY WIND*.

To Specify a constant wind in space, one can use keyword WIND VELOCITY ALONG X and WIND VELOCITY ALONG Y

When the wind can be described analytically, the user subroutine USER_ANAVEN.f can be used. The wind is fully specified when the keywords *FORMATTED WINDS FILE* and *BINARY WINDS FILE* do not have any value, whereas the keyword *CONSIDERATION OF A WIND* is "TRUE".

TOMAWAC can also take into account a wind given in a binary or formatted file. In this case a value (the name of the file) should be assigned to the keyword *FORMATTED WINDS FILE* or *BINARY WINDS FILE*. The available formats for reading out these data correspond to the keyword *WINDS FILE FORMAT* (see in section 7.2.6 and A.3).

When these predefined formats cannot be used, the subroutine USER_WIND.f can be included in the FORTRAN file and modified accordingly, specifying the format 4 for the INDIV FORTRAN variable. On completion of reading the winds file, the wind components are used as such if provided on the computational mesh, or interpolated over that mesh in provided on a different grid.

NOTE: an interpolation between two different meshes of equivalent sizes is usually computationally very expensive. Although possible, it is highly inadvisable, particularly as regards to the wind, since this is a time-varying data item. In such cases a pre-interpolation over the computation mesh, e.g using STBTEL is recommended, followed by the reading of the wind data in format 3. Alternatively this pre-interpolation can be performed by means of the FASP subroutine from the utils library.

8.2.5 Recovering a TELEMAC data item

Recovering a 2D result data item from a TELEMAC-2D hydrodynamic computation might be of interest, e.g. the value of wind-driven surge at every point. To avoid an increase in the number of files the BINARY CURRENTS FILE is used to specify this input file. The keyword CURRENTS FILE FORMAT should then be set to 3. This option is further specified using the logical keyword RECOVERY OF TELEMAC DATA ITEM. The data item is located within the file through the TIME INCREMENT NUMBER IN TELEMAC FILE.

NOTE: a TELEMAC data item and the components of a current can both be read simultaneously provided that they occur in the same file at the same record.

8.2.6 Taking the tide into account

Tide-induced effects, i.e. unsteady/non-stationary water levels and currents can be taken in to account. The relevant logic keyword is *CONSIDERATION OF TIDE*.

In order to take tide into account, a current and a tide water depth that is referenced in relation to the "INITIAL STILL WATER LEVEL" must be specified. These data can be initialized in various ways:

Should the tide be easy to describe analytically, the USER_ANAMAR.f subroutine, can be included in the FORTRAN file and modified accordingly. In the subroutine the terms UC and VC, ZM and DZHDT are NPOIN2-sized (number of points in the horizontal mesh) vectors and correspond to the current components along the X and Y axes, the tidal water level in relation to the "INITIAL STILL WATER LEVEL" and the water depth variation in time, respectively. An analytical expression must be assigned to all of these vectors.

TOMAWAC can also take into account a current that is given in a binary or formatted file. A value (the name of the currents file) should be assigned to the keyword *BINARY CURRENTS FILE* or *FORMATTED CURRENTS FILE*. Two different formats are available for reading these data. This format is specified using the keyword *CURRENTS FILE FORMAT* (see 7.2.4). When these predefined formats cannot be used, the user subroutine USER_CURRENT.f can be included in the FORTRAN file and modified accordingly. In such cases the *CURRENTS FILE FORMAT* (FORTRAN variable INDIC) must be set to 4 in the CAS file. Once the data of the currents file are read, the current components are interpolated over the computation mesh.

The tidal water level can be also provided in a binary or formatted file. A value (the name of the water level file) should be assigned to the keyword *BINARY TIDAL WATER LEVEL FILE* or *FORMATTED TIDAL WATER LEVEL FILE*. Two different predefined formats are available for reading this data. The format type is specified using the keyword *TIDAL WATER LEVEL FILE FORMAT* (refer to 7.2.5). If the user chooses the Serafin format (i.e. TELEMAC-2D format), then the *RANK OF THE WATER LEVEL DATA IN THE TELEMAC FILE* must also be specified. When the predefined formats cannot be used, the user subroutine USER_TIDE.f file can be included in the FORTRAN file and modified accordingly. In such cases the *TIDAL WATER LEVEL FILE FORMAT* (the INDIM FORTRAN variable) must be set to 4 in the CAS file

Both currents and tidal water levels will be updated upon each *TIDE REFRESHING PERIOD*. This keyword corresponds to an integer multiple of the propagation *TIME STEP*, i.e. the currents and tidal levels cannot be specified at time steps less than the model TIME STEPS.

8.2.7 Waves-current interactions: direct coupling with Telemac (2D or 3D) flow simulation

It is possible to directly couple a TOMAWAC and a TELEMAC- simulation (either TELEMAC-2D or TELEMAC-3D) to represent the wave-current interactions more precisely.

In TOMAWAC, when the keywords *CONSIDERATION OF A TIDE* or *CONSIDERATION OF A STATIONARY CURRENT* are used, the current is imposed as an input data: in this case only the effect of the current on the waves is taken into account, but not the effect of the waves on the current. The current imposed, therefore, is not affected by the waves and can differ from the real current interacting with the waves.

By using a direct coupling TELEMAC-TOMAWAC it is possible to represent wave current interactions in both directions: TELEMAC transfers to TOMAWAC the updated values of current velocities and water depths, while TOMAWAC solves the wave action density conservation equation with reference to those current and water depth values and returns to TELEMAC-T he updated values of the wave driving forces FX and FY acting on the current.

To directly couple a TELEMAC model with a TOMAWAC model, the following conditions must be satisfied:

- the TELEMAC and TOMAWAC simulations must have the same simulation time length (given by the time step multiplied by the number of time steps)
- the time steps set for the two simulations must be equal or multiple of each other
- current and/or water level file cannot be used as input data for the TOMAWAC simulation
- the driving force along X and Y (FX, FY) must be set among the 2D output variables in the steering file

There are two kinds of coupling, in the first one the TELEMAC- and TOMAWAC models must have the same horizontal mesh. Using tel2tom you are allowed to have different meshes with interpolation of variables between the 2 meshes. In that case the weight of interpolations must have been calculated before the simulation and must be recorded in the meshes. This can be done using the command $run_telfile.py\ tel2tom$. An example of use of this command is done in the notebook: pretel/tel2tom.ipynb

In case of direct coupling TELEMAC-TOMAWAC, TELEMAC is the main programme and calls the TOMAWAC subroutine WAC, which is the main subroutine of TOMAWAC and solves the equation of generation and propagation of the directional wave spectrum.

The TELEMAC-TOMAWAC coupling is set via four keywords in the TELEMAC steering file:

- *COUPLING WITH*, to which the value 'TOMAWAC' must be assigned for the first kind of coupling. If you want to use tel2tom and have different meshes for TOMAWAC and TELEMAC- the value TOMAWAC2 must be assigned.
- WAVE DRIVEN CURRENT must be set to 1
- TOMAWAC STEERING FILE, which specifies the name of the TOMAWAC steering file: its path must be specified with reference to the working directory of the TELEMAC steering file.
- COUPLING PERIOD FOR TOMAWAC (variable PERCOU_WAC), which specifies how many TELEMAC time steps TOMAWAC is called:
 - The TOMAWAC time step must be less or equal TELEMAC simulation time step multiplied by PERCOU_WAC.
 - The ratio of the two time steps must be an integer.

Different examples are given in gaia/littoral-t2d-tom and Coupling_Wind.

In the case of coupling with TELEMAC-3D, you can also make the difference between a 2D coupling and a real 3D coupling. To specify that you want a 3D coupling you have to add *T3D* after coupling with tomawac (see the documentation of TELEMAC-3D). Some examples of this 3D coupling are given in the test cases (*Rip* and *3Dcoupling*).

All the file names specified in the TOMAWAC steering file (geometry file, boundary conditions file, Fortran file, ...) must be given as paths relative to the working directory of the TELEMAC steering file. Note that, when you use direct coupling, you are not allowed to read a wind in TOMAWAC: wind must be transmitted by TELEMAC-2D or TELEMAC-3D

For more information concerning the modelling options in TELEMAC-, please refer to the TELEMAC- documentation.

8.2.8 Convection step

For specific validation tests, for example, it may be interesting to drop the convection step and only consider the effect of the source terms. To do this requires assigning the "FALSE" value to the keyword *CONSIDERATION OF PROPAGATION*.

8.2.9 Taking diffraction into account

Diffraction can be included in TOMAWAC simulations only if the following conditions are fulfilled:

- The spatial mesh is defined in terms of Cartesian coordinates.
- Neither currents nor time-varying levels are taken into account.
- The simulation is run sequentially and not in parallel mode.

Four keywords can be used to parameterize the representation of diffraction in TOMAWAC:

- *DIFFRACTION*: its default value is zero, and corresponds to the case in which diffraction is not taken into account. If *DIFFRACTION* is set to one, diffraction is considered and the Mild Slope Equation (MSE) approach is applied. If *DIFFRACTION* is set to two, diffraction is considered and the Revised Mild Slope Equation (RMSE) approach is applied.
- STARTING TIME STEP FOR DIFFRACTION: it indicates at which time step of the simulation diffraction is taken into account. Its default value is 1 (i.e. diffraction is solved from the first time step). When diffraction is considered in TOMAWAC simulations, the computing time is longer. To reduce it, it is recommended to set STARTING TIME STEP FOR DIFFRACTION at a value corresponding approximately to the time step at which the same TOMAWAC model, without the consideration of diffraction, reaches a steady-state.
- VARIANCE THRESHOLD FOR DIFFRACTION: it indicates the minimum spectral variance threshold considered when diffraction is taken into account. The default value is 1E-12.
- *DIFFRACTION FILTER*: if TRUE, it indicates that the filtering of the local amplitudes of directional spectra is carried out when diffraction is considered.

The diffraction module implemented in TOMAWAC is able to capture the diffusion of the wave height behind diffracting structures, but it is not adapted to represent in detail the diffracted wave field.

The first version of the diffraction module implemented in TOMAWAC presents still several limits that must be overcome:

- Unrealistic energy build-up and numerical noise are generated when diffraction is taken into account: at a constant Courant number, the smaller the mesh element size to wavelength ratio, the larger the numerical instability.
- The simulation results are very sensitive to the Courant number and to the ratio between the mesh element size and the wavelength associated to the peak period.
- In the case of a complex and rapidly varying bathymetry (e.g. a submerged shoal case), the model underestimates the effect of diffraction.

For problems in which diffraction must be simulated precisely (e.g. harbor agitation), the use of phase-resolving model is recommended.

The following recommendations can be made, in order to build and run a TOMAWAC model in which diffraction is taken into account:

- A too fine mesh can lead to numerical noise and unsteadiness in the results, while with a mesh too coarse the diffraction effect is underestimated. The best compromise found so far for the parameterization of TOMAWAC is to set the ratio between the smallest mesh element size and the wavelength associated to the peak period $\Delta X/\Lambda$ within the interval 0.2-0.25 (i.e. approximately 4 to 5 mesh elements per wavelength)
- When setting the Courant number Cr, a compromise must be found between the computing time required (large values of Cr imply shorter computing times) and a low numerical noise in the solution (the lower Cr, the lower the numerical noise). For values of $\Delta X/\Lambda$ within the interval 0.2-0.25 a Courant number Cr not larger than 0.1 is recommended.
- A numerical smoothing filter can improve the simulation results, especially when using a mesh with a very high spatial resolution (small $\Delta X/\Lambda$, i.e. when numerical noise and energy build-up are generated during the simulation. The filter smoothens in the spatial domain the local amplitude of the directional spectrum.

8.3 Parameterising the source term integration step

8.3.1 Introduction

When it is required to take into account the source/sink terms, the logic keyword *CONSIDER-ATION OF SOURCE TERMS* should be set to "TRUE".

It has been shown that the source term integration may require a shorter time step than the time step that is used for convection. The *TIME STEP* in the steering file corresponds to the convection time step. The source term integration step is controlled using the integer keyword *NUMBER OF ITERATIONS FOR THE SOURCE TERMS*. This keyword is set to the number of source terms integration time steps that will be conducted after each convection step (default value = 1). The effective time-step used for source term integration is thus:

(TIME STEP)/(NUMBER OF ITERATIONS FOR THE SOURCE TERMS).

Depending on the source/sink terms, two different schemes are used for time integration:

• For the source/sink terms which are dominant in large and medium water depths (namely wind input, white-capping dissipation, nonlinear quadruplet interactions and bottom friction) a scheme with variable implicitation level is used (see section 8.3.2).

• For the source/sink terms which are dominant in shallow water depths (namely depth-induced breaking and, nonlinear triad interactions) an explicit scheme is used, possibly with sub-steps to cover one source term time-step (see section 8.3.3).

8.3.2 Source/sink terms in large and medium water depth Wind input

In TOMAWAC three wind generation models and a linear wave growth model are available (see section 4.2.3.2 for details).

If the keyword WIND GENERATION is set to 0, no wind input will be taken into account. If, on the other hand, a strictly positive value is chosen, the corresponding wind input formulation will be taken into account.

<u>The Janssen formulation</u> is activated using the value 1 for the WIND GENERATION keyword. Janssen's formulation requires several additional data, specified by the following keywords:

- AIR DENSITY,
- WATER DENSITY,
- WIND GENERATION COEFFICIENT,
- VON KARMAN CONSTANT,
- CHARNOCK CONSTANT,
- SHIFT GROWING CURVE DUE TO WIN,
- WIND DRAG COEFFICIENT,
- WIND MEASUREMENT LEVEL.

As a general rule, the default values for these keywords shall not be modified.

The Snyder formulation is activated setting to 2 the value for the WIND GENERATION keyword and uses two parameters, specified by the following keywords:

- AIR DENSITY,
- WATER DENSITY,

<u>The Yan formulation</u> is activated setting to 3 the value for the WIND GENERATION keyword and uses four parameters, specified by the following keywords:

- YAN GENERATION COEFFICIENT D
- YAN GENERATION COEFFICIENT E
- YAN GENERATION COEFFICIENT F
- YAN GENERATION COEFFICIENT H

As a general rule, the default values for these keywords shall not be modified.

The linear wave growth model, based on the Cavaleri & Malanotte-Rizzoli formulation, is activated setting to 1 the keyword *LINEAR WAVE GROWTH*. This formulation does not require any additional data to be specified by the user.

White capping dissipation

In TOMAWAC two whitecapping dissipation models are available (see section 4.2.3.2 for details).

If the integer keyword *WHITE CAPPING DISSIPATIO* is set to 0, this source term will be ignored. If a strictly positive value is selected, the corresponding formulation will be taken into account.

<u>The Komen's formulation</u> corresponds to the value 1 of the *WHITE CAPPING DISSIPATION* keyword. This formulation requires two complementary data, specified by the following keywords:

- WHITE CAPPING DISSIPATION COEFFICIENT
- WHITE CAPPING WEIGHTING COEFFICIENT.

As a general rule, the default values for these keywords shall not be modified.

The Westhuysen formulation corresponds to the value 2 of the WHITE CAPPING DISSIPATION keyword. This formulation requires four complementary data, specified by the following keywords:

- WESTHUYSEN DISSIPATION COEFFICIENT
- SATURATION THRESHOLD FOR THE DISSIPATION
- WESTHUYSEN DISSIPATION MODEL WHITE CAPPING DISSIPATION
- WESTHUYSEN DISSIPATION MODEL WEIGHTING COEFFICIENT

As a general rule, the default values for these keywords shall not be modified.

Bottom friction dissipation

If the integer keyword *BOTTOM FRICTION DISSIPATION* is set to 0, this source term will be ignored. If a strictly positive value is chosen, the corresponding formulation will be taken into account. This source term is only taken account if the keyword *INFINITE DEPTH* is set to "FALSE".

The only formulation implemented is from Hasselmann (see section 4.2.3.4 for details) This formulation is specified by setting the *BOTTOM FRICTION DISSIPATION* keyword to 1. This formulation requires the specification of the keyword:

- BOTTOM FRICTION COEFFICIENT.

As a general rule, the default value for this keyword shall not be modified.

Non-linear transfers between quadruplets

Three non-linear quadruplet interactions models are available in TOMAWAC (see in section 4.2.3.6). The non-linear quadruplet interactions are activated through the keyword *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* in the steering file; the keyword can take four values. If the integer keyword *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* is set to 0, this source term will not be taken into account. If a strictly positive value is chosen, the corresponding formulation will be taken into account.

<u>The DIA method</u> (formulation from [25]) corresponds to the value 1 for the *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* keyword. This formulation does not require any additional data to be specified by the user.

<u>The MDIA method</u> (formulation from [60]) corresponds to the value 2 for the *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* keyword. This formulation does not require any additional data to be specified by the user.

The quusi-excat GQM method (formulation based on [38]) corresponds to the value 3 for the *NON-LINEAR TRANSFERS BETWEEN FREQUENCIES* keyword. This formulation requires the specification of six keywords:

- SETTING FOR INTEGRATION ON OMEGA1
- SETTING FOR INTEGRATION ON THETA1
- SETTING FOR INTEGRATION ON OMEGA2
- THRESHOLDO FOR CONFIGURATIONS ELIMINATION
- THRESHOLD1 FOR CONFIGURATIONS ELIMINATION
- THRESHOLD2 FOR CONFIGURATIONS ELIMINATION

8.3.3 Source/sink terms in shallow water depth

Time integration scheme and time step

As mentioned above, the depth-induced breaking and nonlinear triad interaction terms are time-integrated with an explicit scheme.

As found practically, contributions from these source terms can be overestimated if the selected time step for source term integration is too long. In order to avoid this, TOMAWAC can perform a number of time sub-steps which are specific to these source terms through the keyword *NUMBER OF BREAKING TIME STEPS*.

These time sub-steps are arranged in a geometrical progression, i.e. they are defined in the following way:

 $dt_{i+1} = q dt_i$

where the geometrical ratio q is specified through the keyword: *COEFFICIENT OF THE TIME SUB-INCREMENTS FOR BREAKING*

In order to limit this number of time-steps, TOMAWAC first clips the wave height by setting a maximum H_{m0}/D ratio (D is the water depth) to 1. This ratio can be modified by means of the keyword *MAXIMUM VALUE OF THE RATIO HM0 TO D*. However, this is generally not advisable.

Wave breaking dissipation

If the integer keyword *DEPTH-INDUCED BREAKING DISSIPATION* is taken as 0, this source term will be ignored. If a strictly positive value is chosen, the corresponding formulation will be taken into account.

Four formulations have been implemented (see section 4.2.3.5 for details):

1: Battjes and Janssen's model (1978)

This formulation requires additional data to be provided, specified by the following keywords:

- DEPTH-INDUCED BREAKING 1 (BJ) COEFFICIENT ALPHA
- DEPTH-INDUCED BREAKING 1 (BJ) COEFFICIENT GAMMA1
- DEPTH-INDUCED BREAKING 1 (BJ) COEFFICIENT GAMMA2
- DEPTH-INDUCED BREAKING 1 (BJ)CHARACTERISTIC FREQUENCY
- DEPTH-INDUCED BREAKING 1 (BJ)QB COMPUTATION METHOD

2: Thornton and Guza's model (1983)

This formulation requires additional data to be provided, specified by the following keywords:

- DEPTH-INDUCED BREAKING 2 (TG) COEFFICIENT B
- DEPTH-INDUCED BREAKING 2 (TG) COEFFICIENT GAMMA
- DEPTH-INDUCED BREAKING 2 (TG) WEIGHTING FUNCTION
- DEPTH-INDUCED BREAKING 2 (TG) CHARACTERISTIC FREQUENCY

3: Roelvink's model (1993)

This formulation requires additional data to be provided, specified by the following keywords:

- DEPTH-INDUCED BREAKING 3 (RO) COEFFICIENT ALPHA
- DEPTH-INDUCED BREAKING 3 (RO) COEFFICIENT GAMMA
- DEPTH-INDUCED BREAKING 3 (RO) COEFFICIENT GAMMA2
- DEPTH-INDUCED BREAKING 3 (RO) WAVE HEIGHT DISTRIBUTION
- DEPTH-INDUCED BREAKING 3 (RO)EXPONENT WEIGHTING FUNCTION
- DEPTH-INDUCED BREAKING 3 (RO)CHARACTERISTIC FREQUENCY

4: Izumiya and Horikawa's model (1984)

This formulation requires additional data to be provided, specified by the following keywords:

- DEPTH-INDUCED BREAKING 4 (IH) COEFFICIENT BETAO

- DEPTH-INDUCED BREAKING 4 (IH) COEFFICIENT M2STAR
- DEPTH-INDUCED BREAKING 4 (IH) CHARACTERISTIC FREQUENCY

Triad interactions

If the integer keyword *TRIAD INTERACTIONS* is set to 0, this source term will be ignored. If a strictly positive value is specified, the corresponding formulation will be taken into account. Two formulations (see section 4.2.3.7) have been implemented.

1: LTA model

This formulation requires additional associated data to be specified using the following keywords:

- TRIADS 1 (LTA) COEFFICIENT ALPHA
- TRIADS 1 (LTA) COEFFICIENT RFMLTA

2: SPB model

This formulation requires additional associated data to be specified using the following keywords:

- TRIADS 2 (SPB) COEFFICIENT KTRIADS 2 (SPB) COEFFICIENT K
- TRIADS 2 (SPB) LOWER DIRECTIONAL BOUNDARY
- TRIADS 2 (SPB) UPPER DIRECTIONAL BOUNDARY

ATTENTION: the SPB model is very time-consuming; compared to the LTA model formulation, it requires a computational time approximately 700 times higher.

Wave blocking

If the integer keyword *DISSIPATION BY STRONG CURRENT* is set to 0, this source term will be ignored. If a strictly positive (1 or 2) is specified, the corresponding formulation will be taken into account.

1: Upper limit for spectrum: We locally use a Phillips's constant of 0.0081 to build a limit spectrum. This does not change the value of Phillip's constant in the entire code.

2: Westhuysen formulation:

This Westhuysen formulation requires 2 parameters.

- DISSIPATION COEFFICIENT FOR STRONG CURRENT
- SATURATION THRESHOLD FOR THE DISSIPATION

Caution: the second coefficient is the one used in the white capping formulation of Westhuysen [13].

Vegetation dissipation

If logical keyword *VEGETATION TAKEN INTO ACCOUNT* is set to true, then vegetation is taken into account. As the parameters are case-specific, the parameters should be modified through the fortran user subroutine *QVEG*. In this subroutine the user will give the value \tilde{C}_D , the vegetation drag coefficient, b_v [m] the stem diameter of cylinder (plant), N_v [-] the number of plants per square meter and α the ratio between vegetation height and water depth.

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Porous Media

If logical keyword *POROUS MEDIA* is set to true, then vegetation is taken into account. As the parameters are case-specific, the parameters should be modified through the fortran user subroutine QPOROS. In this subroutine the user will give the value of porosity rate, (TAU), the damping mass coefficient (CA), the linearised friction coefficient (coeff) and the thickness rate of the porous medium (alpha).

8.4 Prescribing the initial conditions

Initial conditions can be prescribed using the integer keyword TYPE OF INITIAL DIREC-TIONAL SPECTRUM.

Table 8.1 shows all the available options in TOMAWAC for computing the frequential and directional distribution of wave action. It should be remembered that the variance density directional spectrum is computed as the product:

$$F(f, \theta) = E(f).D(\theta)$$

where E(f) here denotes the variance density spectrum and $D(\theta)$ denotes the angular distribution function.

It is reminded that the parameterised JONSWAP spectrum is defined as:

$$E(f) = \alpha_{phil} g^{2} (2\pi)^{-4} f^{-5} \exp \left[-\frac{5}{4} \left(\frac{f_{p}}{f} \right)^{4} \right] \gamma^{\exp \left[-\frac{(f - f_{p})^{2}}{2\sigma^{2} f_{p}^{2}} \right]}$$

where: $\sigma = \sigma_a = 0.07$ for $f < f_p$ and $\sigma = \sigma_b = 0.09$ for $f > f_p$. Here, α_{phil} is the Phillips constant that can be specified in the parameter file (keyword INITIAL PHILLIPS CONSTANT). If one wants to parametrize the formula with H_{m0} (case 5 with wind and case 6), one writes

$$E(f) = \alpha_{phil} H_{mo}^2 \frac{f_p^4}{f^5} \exp\left[-\frac{5}{4} \left(\frac{f_p}{f}\right)^4\right] \gamma^{\exp\left[-\frac{\left(f - f_p\right)^2}{2\sigma^2 f_p^2}\right]}$$

With
$$\alpha_{phil}=\frac{0.0624}{0.23+0.0336\gamma-\frac{0.185}{1.9+\gamma}}$$
 as done in Tomawac.

In case 7, the TMA spectrum is a depth-corrected JONSWAP-type spectrum.

A parameterised spectrum with two directional peaks can also be generated. In such case, both main propagation directions (θ_1 and θ_2) as well as the weighting factor (λ) between the two power peaks:

$$D(heta) = rac{\lambda}{\Delta_1} ilde{D}(heta - heta_1) + rac{1-\lambda}{\Delta_2} ilde{D}(heta - heta_2)$$

must be specified.

 Δ_1 and Δ_2 in the equation above, are automatically computed by TOMAWAC in order to normalize the angular distribution function.

Three angular distribution functions may be chosen using the keyword INITIAL ANGULAR DISTRIBUTION FUNCTION, which correspond to the following options:

1: model in
$$\cos^{2s}(\theta-\theta_0)$$
 ; $\theta\in[\theta_0-\pi/2$; $\theta_0+\pi/2]$

2: model in
$$\exp(-0.5((\theta - \theta_0)/s)^2)$$
; $\theta \in [\theta_0 - \pi/2; \theta_0 + \pi/2]$

2: model in
$$\exp\left(-0.5\left((\theta-\theta_0)/s\right)^2\right)$$
; $\theta \in [\theta_0-\pi/2; \theta_0+\pi/2]$
3: model in $\cos^{2s}\left((\theta-\theta_0)/2\right)$; $\theta \in [\theta_0-\pi; \theta_0+\pi]$ Mitsuyasu [46]

4: same than 3 but with
$$s = \begin{cases} s_{max} \left(\frac{f}{f_p}\right)^5 : f \le f_p \\ s_{max} \left(\frac{f}{f_p}\right)^{-2.5} : f > f_p \end{cases}$$
 as proposed by Goda et al. [21].

with θ_0 being the main sea-state propagation direction and s being the directional spread. According to [21], s_{max} should be 10, 25 and 75 for wind waves, swell with short decay distance, and swell with long decay distance.

The forementioned constants can be specified using the following of keywords:

 H_{m0} : INITIAL SIGNIFICANT WAVE HEIGHT,

f_p: INITIAL PEAK FREQUENCY,

γ: INITIAL PEAK FACTOR,

 σ_a : INITIAL VALUE OF SIGMA-A FOR SPECTRUM,

 σ_b : INITIAL VALUE OF SIGMA-B FOR SPECTRUM,

 α_{phil} : INITIAL PHILLIPS CONSTANT,

Fetch: INITIAL MEAN FETCH VALUE,

fpmax: INITIAL MAXIMUM PEAK FREQUENCY,

 θ_1 : INITIAL MAIN DIRECTION 1,

s1: INITIAL DIRECTIONAL SPREAD 1,

 θ_2 : INITIAL MAIN DIRECTION 2,

s2: INITIAL DIRECTIONAL SPREAD 2,

 λ : INITIAL WEIGHTING FACTOR FOR ADF,

The keyword *SPECTRUM ENERGY THRESHOLD* is used whatever option is chosen. It is only useful for comparisons with the WAM model.

Specific initial conditions can be prescribed directly for the directional spectrum of wave action using the condiw.f subroutine which can be called from the speini.f subroutine.

8.5 Prescribing the boundary conditions

The boundary conditions are prescribed over the <u>relative</u> spectrum of wave action, i.e. expressed in co-ordinate system that moves with the current.

There are three kinds of boundary conditions are available in TOMAWAC.

The first one corresponds to a free boundary i.e. a boundary that fully absorbs the wave energy. It may be a liquid boundary: it is then assumed that the waves propagate beyond the domain and nothing else enters it. It may be a solid boundary: it is then assumed that the shore fully absorbs the wave energy. The colour of this condition will begin by 2, and will be 2 2 2 2 to be consistent with telemac.

The second one corresponds to a boundary with a prescribed value. In this case the wave action spectrum is then strictly imposed at each point along that boundary. This boundary condition allows wave energy to enter the computational domain. The imposed spectro-angular spectrum can be given by analytical function such as a jonswap (see 8.1) or imposed by an imposed spectra file (see). The colour of this condition will start with 5 and will be 5 4 4 4 to be consistent with telemac.

The third one corresponds to a reflective boundary. As there are no natural expression to have a reflective condition with the equations governing waves propagation in Tomawac, this condition is more a source dependending on the incoming wave. At each point of this boundary for each frequency and each direction of the incoming wave, we construct a source with the direction transmitted following the law of Snell Descartes(see 8.1) and projected on the discretised directions.

	ı		I
Key-			
word		Spectrum	Constants being used
value			
0		Zero spectrum	none
1	Wind $\neq 0$	- Frequencies: Jonswap according to	f_{pmax} , γ , σ_a , σ_b , $Fetch$, s_1
		wind	
		- Directions: unimodal about the	
		wind $(\theta_1 = \theta_w)$	
	Wind=0	Zero spectrum	none
2	Wind $\neq 0$	- Frequencies: Jonswap according to	f_{pmax} , γ , σ_a , σ_b , $Fetch$, s_1
		wind	
		- Directions: unimodal about the	
		wind $(\theta_1 = \theta_w)$	
	Wind=0	- Frequencies: parameterised Jon-	$\alpha_{phil}, f_p, \gamma, \sigma_a, \sigma_b, s1, \theta_1$
		swap (α, f_p)	
		- Directions: parameterised uni-	
		modal (same spectrum at every	
		point)	
3	Wind $\neq 0$	- Frequencies: parameterised Jon-	$\alpha_{phil}, f_p, \gamma, \sigma_a, \sigma_b, s1$
		swap (α)	
		- Directions: unimodal about the	
		wind $(\theta_1 = \theta_w)$	
	Wind=0	Zero spectrum	none
4		- Frequencies: parameterised Jon-	$\alpha_{phil}, f_p, \gamma, \sigma_a, \sigma_b, s1, \theta_1, s2,$
		swap (α, f_p)	θ_2, λ
		- Directions: parameterised angular	
		distribution function. Same spec-	
		trum at every point	
γ , σ_a , σ_b , $s1$, θ_1 , $s2$, θ_2 , λ			
5	Wind $\neq 0$	- Frequencies: parameterised Jon-	H_{m0} , f_p , γ , σ_a , σ_b , s1
		swap (H_{m0})	•
		- Directions: unimodal about the	
		wind $(\theta_1 = \theta_w)$	
	Wind=0	Zero spectrum	none
6		- Frequencies: parameterised Jon-	H_{m0} , f_p , γ , σ_a , σ_b , $s1$, θ_1 , $s2$,
		swap (H_{m0}, f_p)	θ_2, λ
		- Directions: parameterised angular	
		distribution function. Same spec-	
		trum at every point	
7		- Frequencies: parameterised TMA	H_{m0} , f_p , γ , σ_a , σ_b , $s1$, θ_1 , $s2$,
		(H_{m0},f_p)	θ_2,λ
		- Directions: parameterised angular	
		distribution function. Same spec-	
		trum at every point	

Table 8.1: Summary table of the spectrum types as proposed in TOMAWAC

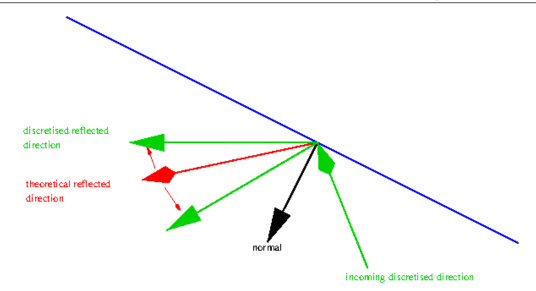


Figure 8.1: scheme of reflection in TOMAWAC

To have reflection the user must set the keyword *REFLECTION* to *YES* (default is *NO*) The user can also specify a coefficient of reflection, by the keyword *REFLECTION COEFFICIENT* (the default is 1). The colour of reflection condition must start by 8 and will be 8 4 4 4 to be consistent with telemac.

The boundary conditions are specified using the boundary conditions (CONLIM) file, the steering (CAS) file and the LIMWAC.f.

8.5.1 The boundary conditions file

The Serafin boundary condition file

The boundary condition (CONLIM) file is normally supplied by STBTEL (or other TELEMAC mesh generators), but it can also be generated by means of a text editor. Each line in this file is assigned to one point of the boundary and listed in sequential order in terms of the boundary node numbers. The numbering of the boundary points first delineates the domain contour in a counterclockwise direction, then the islands in the clockwise direction. The total number of edge points is noted as NPTFR.

13 values are given for each point. Only data in colums 1, 12 and 13 are used by TOMAWAC:

- The 13th data column (integer variable IPTFR) corresponds to the boundary point number ranked in terms of the boundary point numbering.
- The 12th data column (integer variable IPOIN) corresponds to the global number of the point in the 2D mesh.
- Lastly, the 1st data column (integer variable LIFBOR) corresponds to the kind of boundary condition. Consistent with TELEMAC-2D, its value is 2 in the case of a free boundary and 5 in the case of a boundary with a prescribed value.

The Med boundary condition file

In that case each line of the file gives the colour of a group of segment. To be consistent with TELEMAC-2 d 4 colours has to be given but only the first one is taken into account by tomawac. Each points belonging to the segments of the group will then have the colour designed.

8.5.2 Prescribing the boundary conditions in the CAS file

Boundary conditions prescribed using the CAS file will necessarily be homogeneous all along the domain entry boundaries.

The boundary conditions can be prescribed by means of the integer keyword TYPE OF BOUND-ARY DIRECTIONAL SPECTRUM.

Table 8.1 (see section 8.4) presents all the spectrum types available in TOMAWAC. The constants given in this table can be prescribed using the following keywords:

H_{m0}: BOUNDARY SIGNIFICANT WAVE HEIGHT,

 f_p : BOUNDARY PEAK FREQUENCY,

γ: BOUNDARY PEAK FACTOR,

 σ_a : BOUNDARY SPECTRUM VALUE OF SIGMA-A,

 σ_b : BOUNDARY SPECTRUM VALUE OF SIGMA-B,

 α_{nhil} : BOUNDARY PHILLIPS CONSTANT,

Fetch: BOUNDARY MEAN FETCH VALUE,

fpmax: BOUNDARY MAXIMUM PEAK FREQUENCY,

 θ_1 : BOUNDARY MAIN DIRECTION 1,

s1: BOUNDARY DIRECTIONAL SPREAD 1,

 θ_2 : BOUNDARY MAIN DIRECTION 2,

s2: BOUNDARY DIRECTIONAL SPREAD 2,

 λ : BOUNDARY WEIGHTING FACTOR FOR ADF,

Three angular distribution functions have been implemented and can be selected using of the keyword: BOUNDARY ANGULAR DISTRIBUTION FUNCTION, which corresponds to the following options:

1: model in $\cos^{2s}(\theta - \theta_0)$; $\theta \in [\theta_0 - \pi/2; \theta_0 + \pi/2]$

2: model in $\exp(-0.5((\theta - \theta_0)/s)^2)$; $\theta \in [\theta_0 - \pi/2; \theta_0 + \pi/2]$

3: model in
$$\cos^{2s}((\theta - \theta_0)/2)$$
; $\theta \in [\theta_0 - \pi; \theta_0 + \pi]$ [46]
4: same than 3 but with $s = \begin{cases} s_{max} \left(\frac{f}{f_p}\right)^5 : f \leq f_p \\ s_{max} \left(\frac{f}{f_p}\right)^{-2.5} : f > f_p \end{cases}$ as proposed by Goda et al. [21]

Since the boundary spectrum computation procedures are similar to those for the initial spectrum, refer to section 8.4 for further details.

8.5.3 Imposing spectra along an open boundary

It is also possible to impose spectra along the nodes of an open boundary; i.e. one that follows the format 5 4 4 4, so that LIFBOR is equal to 5. The keywords are described in section 8.5, and an example is given bellow:

```
Keywords to impose spectra along an open boundary:
 IMPOSE SPECTRA ON THE OPEN BOUNDARY
IMPOSED SPECTRA FILE = <FileName>
/IMPOSED SPECTRA FILE FORMAT = 'SERAFIN'
FILE WITH COORDINATES OF SPECTRA TO IMPOSE =
/TIME UNIT OF IMPOSED SPECTRA FILE = 1.
TIME SHIFT OF IMPOSED SPECTRA FILE = 0.
```

The imposed spectra are interpolated linearly in time, which implies that the first recorded value needs to be at least equal to the begining of the simulation, and the last recorded value needs to be at least equal to the last time step. Furthermore, the open boundary nodes of the mesh will take the value of the closest imposed spectra point, as defined by *FILE WITH COORDINATES OF SPECTRA TO IMPOSE*. It is therefore recommended to have in the *IMPOSED SPECTRA FILE* one spectra per open boundary node.

Finally, this method of imposing spectra on the open boundary is particularly useful when the user wants to use nested meshes, such as described in the impose_spectra example. In this case, the bigger mesh needs to output one spectra per node in the smaller mesh. This can be easily done using the keywords associated with the *PUNCTUAL RESULTS FILE*, see section 7.2.13. Care should also be given on the frequency with which the spectra will be outputted, as a linear interpolation in time might not follow the evolution of the boundary if the imposed values are too far appart.

8.5.4 The LIMWAC user subroutine

It should be reminded that the spectrum is discretised over both frequencies and directions and that it is a <u>relative</u> spectrum, i.e. expressed in a coordinate system that moves with the current. The subroutine LIMWAC, in its original version, allows to impose the spectrum components at each point of a boundary with a prescribed value. The spectrum components are calculated from the parameters specified in the CAS file (see section 7.2.1). This subroutine, however, can easily be modified to specify e.g. non-homogeneous (in space) boundary conditions. When such specific boundary conditions are required, these will ideally be incorporated in the user part provided in the code of the LIMWAC procedure. The keyword *BOUNDARY SPECTRUM MODIFIED BY USER* must also be set to YES.

8.6 Some useful subroutines

8.6.1 Modification of bottom topography: USER_TOM_CORFON subroutine

The seabed levels can be modified in two different ways, as already stated in section 7.4 The seabed levels can be modified at the beginning of the computation using the USER_TOM_CORFON subroutine, which is called once at the beginning of the computation. This subroutine allows the value of the ZF variable to be modified at each mesh point. For this purpose, a number of variables such as, for instance, the point coordinates, the element area values, the connectivity table, etc., are provided.

By default, the USER_TOM_CORFON subroutine performs the same number of bottom smoothing iterations as LISFON, i.e. the same value as specified by the integer keyword *BOTTOM SMOOTHINGS*.

Note that the USER_TOM_CORFON subroutine is not called in case the computation is initialized with the result of a former TOMAWAC run ("hot start").

This subroutine is listed in A.2

8.6.2 Modifying the co-ordinates: USER_CORRXY subroutine

TOMAWAC allows the mesh point co-ordinates to be modified at the beginning of the computation, so that an up-scaling (switching from a small scale model to a full-size model), a rotation or a translation can be performed.

Such changes are made using the CORRXY subroutine from the BIEF library, which is called in at the beginning of the computation. This subroutine is void by default and provides, in the form of a comment, an example of programming relevant to a change of scale and origin. It is part of the TELEMAC-2D library and is listed in A.2

8.6.3 Operations on vectors: OVOV subroutine

The BIEF library has a range of very useful subroutines including, in particular, subroutines for operations on vectors. A number of relations have been programmed so that loops can be replaced by a mere procedure call.

The syntax is as follows:

CALL OVOV(OP, X, Y, Z, C, NPOIN)

Where OP is a string of exactly 8 digits that is indicative of the operation about to be performed on the X, Y, Z vectors and the constant C. The result is the vector X.

Example: CALL OVOV('X=X+Y ', X, Y, Z, C, NPOIN) Y is added to X, the result will be stored in X.

A full list of available operations are given in the Guide to programming in the Telemac system version 6.0.

9. API

Information on the TOMAWAC API can be found in the telapy user documentation.

A. appendix

A.1 Output variables of TOMAWAC

Table A.1: Output variables of TOMAWAC

N°	Symbol	Description of the variable	Unit	Output name
1	M0	Variance of the sea state m_o . moment of order 0 of the wave spectrum. $m_0 = \int_0^\infty \int_0^{2\pi} F(f,\theta) df d\theta = \int_0^\infty S(f) df$	m^2	VARIANCE M0
2	HM0	Spectral significant wave height H_{mo} $H_{mo} = 4\sqrt{m_0}$	m	WAVE HEIGHT HM0
3	DMOY	Mean wave direction θ_m with respect to the North or to the X axis depending on the adopted choice. $\theta_m = Arc \tan(\frac{b}{a}) * (180/\pi)$ $a = \frac{\int_0^\infty \int_0^{2\pi} \cos(\theta) F(f,\theta) df d\theta}{m_o}$ $b = \frac{\int_0^\infty \int_0^{2\pi} \sin(\theta) F(f,\theta) df d\theta}{m_o}$	deg.	MEAN DIRECTION
4	SPD	Mean angular spreading σ of the directional wave spectrum. $\sigma = \sqrt{2\left(1 - \sqrt{a^2 + b^2}\right)} * (180/\pi)$	deg.	WAVE SPREAD
5	ZF	Sea bottom elevation.	m	BOTTOM
6	WD	Water depth.	m	WATER DEPTH
7	UX	Component along X axis of the marine flow velocity vector (input data)	m/s	VELOCITY U
8	UY	Component along Y axis of the marine flow velocity vector (input data)	m/s	VELOCITY V
9	VX	Component along X axis of the wind velocity vector (input data)	m/s	WIND ALONG X
10	VY	Component along Y axis of the wind velocity vector (input data)	m/s	WIND ALONG Y

Table A.1: (continued)

N°	Symbol	Description of the variable	Unit	Output name
11	FX	Component along X axis of the radiation force due to waves.	m/s ²	FORCE FX
12	FY	Component along Y axis of the radiation force due to waves.	m/s ²	FORCE FY
13	SXX	Component Sxx of the radiation stress tensor.	m^3/s^2	STRESS SXX
14	SXY	Component Sxy of the radiation stress tensor.	m^3/s^2	STRESS SXY
15	SYY	Component Syy of the radiation stress tensor.	m^3/s^2	STRESS SYY
16	UWB	Mean orbital wave velocity at the bottom.	m/s	BOTTOM VELOCITY
17	PRI	Private variable as chosen by the user.	(user)	PRIVATE 1
18	FMOY	Mean wave frequency, computed from the moments of order -1 and 0 of the wave spectrum. $f_{moy} = f_{-10} = \left(\frac{m_{-1}}{m_0}\right)^{-1} = \left(\frac{\int_0^\infty \frac{S(f)}{f} df}{\int_0^\infty S(f) df}\right)^{-1}$	Hz	MEAN FREQ
19	FM01	Mean wave frequency, computed from the moments of order 0 and 1 of the wave spectrum. $f_{01} = \frac{m_1}{m_0} = \frac{\int_0^\infty f . S(f) df}{\int_0^\infty S(f) df}$	Hz	MEAN FREQ FM01
20	FM02	Mean wave frequency, computed from the moments of order 0 and 2 of the wave spectrum. $f_{02} = \sqrt{\frac{m_2}{m_0}} = \sqrt{\frac{\int_0^\infty f^2 . S(f) df}{\int_0^\infty S(f) df}}$	Hz	MEAN FREQ FM02
21	FPD	Discrete peak frequency (among the frequencies used to discretise the spectrum).	Hz	PEAK FREQ FPD
22	FPR5	Peak frequency computed by the Read method of order 5. $f_{pR5} = \frac{\int_0^\infty f . S^5(f) df}{\int_0^\infty S^5(f) df}$	Hz	PEAK FREQ FPR5
23	FPR8	Peak frequency computed by the Read method of order 8. $f_{pR8} = \frac{\int_{0}^{\infty} f . S^{8}(f) df}{\int_{0}^{\infty} S^{5}(f) df}$	Hz	PEAK FREQ FPR8
24	US	Surface friction velocity u*.	m/s	USTAR
25	CD	Surface drag coefficient C_D .	-	CD
26	Z 0	Surface roughness length z_o .	m	Z0
27	WS	Surface stress due to waves.	kg/(m.s ²)	WAVE STRESS

Table A.1: (continued)

N°	Symbol	Description of the variable	Unit	Output name
28	TMOY	Mean wave period, computed from the moments of order -1 and 0 of the wave spectrum. $T_{moy} = T_{-10} = \frac{m_{-1}}{m_0} = \frac{\int_0^\infty T.S(f) df}{\int_0^\infty S(f) df}$ Mean wave period, computed from the mo-	S	MEAN PERIOD TMOY
29	TM01	ments of order 0 and 1 of the wave spectrum. $T_{01} = \frac{1}{f_{01}}$	S	MEAN PERIOD TM01
30	TM02	Mean wave period, computed from the moments of order 0 and 2 of the wave spectrum. $T_{02} = \frac{1}{f_{02}}$ Discrete peak period.	S	MEAN PERIOD TM02
31	TPD	$T_{pD} = rac{1}{f_{pD}}$	S	PEAK PERIOD TPD
32	TPR5	Peak period computed by the Read method of order 5. $T_{pR5} = \frac{1}{f_{pR5}}$	S	PEAK PERIOD TPR5
33	TPR8	Peak period computed by the Read method of order 8. $T_{pR8} = \frac{1}{f_{pR8}}$	S	PEAK PERIOD TPR8
34	POW	Unit wave power rate (per meter of crest length).	kW/m	WAVE POWER
35	BETA	Breaking wave rate	-	BREAKING RAT
36	BETAWC	White Capping rate	-	WHITE CAPING
37	SRE	Surface Roller Energy	M3/S2	SURFACE ROLLER E
38	DBR	Breaker Dissipation	M2/S	BREAKER DIS- SIP
39	DSR	Roller Dissipation	M3/S3	ROLLER DISSIP

A.2 List of useful subroutines

The following subroutines can be modified by the user:

- USER_ANACOS.f: Specification of an analytical stationary current.
- USER_ANAMAR.f: Specification of an analytical tide (current and water level).
- USER_ANAVEN.f: Specification of an analytical wind.
- USER_TOM_CORFON.f: Modification of bottom topography.
- USER_CORRXY.f: Modification of coordinates.

- USER_CURRENT.f: Currents file reading from a user-defined format.
- USER_DUMP2D.f: To define a new output defined by the user.
- USER_LIMWAC.f: Specification of the boundary conditions.
- USER_TIDE.f: Tide level file reading from a user-defined format.
- USER_WIND.f: Winds file reading from a user-defined format.

When these subroutines are used, they should be integrated into the user's FORTRAN file, or directory, then modified.

A.3 Description of the formats being used

SERAFIN-formatted file It is a binary file. This format is used for the 2D RESULTS FILE2D RESULTS FILE, the CURRENTS FILE FORMAT (format 3) and the WINDS FILE FORMAT (format 3).

The list of records is as follows:

- a record including the study title (80 digits),
- a record including the pair of integers NBV(1) and NBV(2) (number of variables of linear and quadratic discretisations, NBV(2) being 0),
- NBV(1) + NBV(2) records including (in 32 digits) each variable's name and unit,
- a record including the integers 1,0,0,0,0,0,0,0 (10 integers, only the first of which is presently used),
- a record including the integers NELEM,NPOIN,NDP,1 (number of elements, number of points, number of points per element and the value 1),
- a record including the integer array IKLE ((NDP,NELEM)-dimensioned array), the connectivity table. WARNING: the dimensions of this array are (NELEM,NDP) in TOMAWAC),
- a record including the integer array IPOBO (NPOIN-dimensioned array). An item value is 0 for an inner point, and provides the edge point numbers for the others),
- a record including the X real array (NPOIN-dimensioned array of point abscissae),
- a record including the Y real array (NPOIN -dimensioned array of point ordinates),

The following can then be found for each time step:

- a record including time T (real),
- NBV(1)+NBV(2) records including the results arrays for each variable at time T.

TOMAWAC-formatted file

It is a binary file. That format is used for the *PREVIOUS COMPUTATION FILE* and the *GLOBAL RESULT FILE*

The list of records is as follows:

• a record including the study title (80 digits).

- a record including the pair of integers NPLAN and NF corresponding respectively to the number of propagation directions and the number of frequencies.
- a record including the pair of integers NELEM2 and NPOIN2 corresponding respectively to the numbers of elements and 2D points.
- a record including end-of-computation time t (real).
- a record including the NPLAN-dimensioned real array TETA (directions of propagation, as expressed in radians).
- a record including the NF-dimensioned real array FREQ (propagation frequencies, as expressed in Hz).
- a record including the NPOIN*NPLAN*NF –dimensioned real area F (directional spectrum of wave action) at time t.

When a current is taken into account, then one can find:

- a record including the NPOIN–dimensioned real array UC (component of current along X) upon time t.
- a record including the NPOIN-dimensioned real array VC (component of current along Y) at time t.

When a wind is taken into account, then one can find:

- a record including the NPOIN–dimensioned real array UV (component of wind along X) at time t.
- a record including the NPOIN–dimensioned real array VV (component of wind along Y) at time t.

When the tide is taken into account, then one can find:

• a record including the NPOIN-dimensioned real array DEPTH at time t.

VENTS-WAM-Cycle 4-formatted file

It is a formatted file. That user would have to implement the reading if wind file format equals 4.

The list of records is as follows:

- 1- Winds grid input dimensions:

KCOL, KROW, RLATS, RLATN, RLONL, RLONR, ICOORD, IWPER

Read with the format (2I4,4F9.3,2I2)

KCOL: Number of longitudes in the winds grid

KROW: Number of latitudes in the winds grid

RLATS: Latitude of southern grid boundary (degrees)

RLATN: Latitude of northern grid boundary (degrees)

RLONL: Longitude of western grid boundary (degrees)

RLONR: Longitude of eastern grid boundary (degrees)

ICOORD: Code of coordinates (ICOORD=1 mandatory)

IWPER: Code of periodicity (IWPER=0 mandatory)

The latitude and longitude increments are respectively computed by:

DPHI = (RLATN-RLATS)/(KROW-1)

DLAM = (RLONR-RLONL)/(KCOL-1)

Subsequently, for each wind field date:

- 2- Wind field date:

IDTWIR read with format (I10)

WAM-formatted wind field date WAM (yymmddhhmm)

3- Horizontal (W-E) wind components at the grid points:

ICODE read with format (I2)

Input field type flag

1: friction velocities

2: surface stresses

3: wind velocities at 10 m

(UWND(ILON,ILAT), (ILON=1,KCOL), ILAT=1,KROW)

read with format (10F6.2)

Horizontal (W-E) component value

- 4- Vertical (S-N) wind components at the grid points:

KCODE read with format (I2)

Input field type flag

KCODE should be equal to ICODE.

(VWND(ILON,ILAT), (ILON=1,KCOL), ILAT=1,KROW)

read with format (10F6.2)

Vertical (S-N) component value

The longitudes are scanned from the West eastwards and the latitudes from the South northwards

"finite differences" type- formatted file

It is a formatted file. That file is possibly used for the *FORMATTED CURRENTS FILE*, the *FORMATTED WINDS FILE* or the *FORMATTED TIDAL WATER LEVEL FILE* when the option 1 is chosen for the format.

The list of records is as follows:

- a record including eight integers NCOL, NLIG, YMIN, YMAX, XMIN, XMAX, BID1 and BID. These variables respectively correspond to the number of columns, the lines of the mesh, the minimum and maximum ordinates, the minimum and maximum abscissae of the mesh, followed by 2 variables that are left unused by TOMAWAC.
- an empty record
- a record including the first component of the variable to be retrieved (for example Ux)
- a record including, if any, the second component of the variable to be retrieved (for example Uy)

SINUSX -formatted file

This format is used for the BOTTOM TOPOGRAPHY FILE and possibly for the CURRENTS FILE.

It is a quite simple format, consisting in successive records of the X, Y, ZF type for the bottom topography file and of the X, Y, UC, VC type for the currents file.

A.4 File structure and processing

All files in the Serafin format may now be built in 3 different formats (given in 8 characters):

'SERAFIN': old format, understood by Rubens

'SERAFIND': the same in double precision, not understood by Rubens

'MED ': understood by the Salome Platform.

Consequently, for every previous Serafin file called "NAME-OF-FILE", a key-word "NAME-OF-FILE FORMAT" has been added. In French "FORMAT DU FICHIER DE ...".

For simplifying the implementation of this new possibility, as well as simplifying the coupling between programmes, a new file structure has been added to library BIEF. The goal is to store all the information related to a file in a single structure. The previous names of logical units and file names, such as NGEO, NRES,... and NOMGEO, NOMRES,... have been replaced by this new structure. It concerns ALL the files, not only the Serafin format. As a consequence, all the subroutines reading or writing to files have been modified.

More details are given here below.

Inputs and outputs: opening and closing files

Structure of files The new Fortran 90 structure for files is as follows: C C C STRUCTURE OF FILE C C== C TYPE BIEF_FILE C C LU: LOGICAL UNIT TO OPEN THE FILE INTEGER LU \mathbf{C} C NAME: NAME OF FILE CHARACTER(LEN=144) NAME C C TELNAME: NAME OF FILE IN TEMPORARY DIRECTORY CHARACTER(LEN=6) TELNAME C FMT: FORMAT (SERAFIN, MED, ETC.) CHARACTER(LEN=8) FMT C ACTION: READ, WRITE OR READWRITE CHARACTER(LEN=9) ACTION C BINASC: ASC FOR ASCII OR BIN FOR BINARY CHARACTER(LEN=3) BINASC C C TYPE: KIND OF FILE CHARACTER(LEN=12) TYPE END TYPE BIEF FILE

The various data and results files of TOMAWAC are described in its dictionary. The information relevant to files will be read with the subroutine READ_SUBMIT, which is called in subroutine LECDON_TOMAWAC, and stored in an array of file structures (called, WAC_FILES). Hereafter is given an excerpt of TOMAWAC dictionary regarding the results file:

NOM = 'FICHIER DES RESULTATS 2D'

NOM1 = '2D RESULTS FILE2D RESULTS FILE'

TYPE = CARACTERE

INDEX = 08

MNEMO = 'WAC FILES(WACRES)%NAME'

SUBMIT = 'WACRES-READWRITE-08; WACRES; OBLIG; BIN; ECR; SELAFIN'

DEFAUT = ' '

DEFAUT1 = ' '

The character string called SUBMIT is used both by the perl scripts and, through Damocles, by the Fortran programme. It is composed of 6 character strings.

The first string, here WACRES-READWRITE-08, is made of:

- 1) the fortran integer for storing the file number: WACRES (which is declared in declarations_telemac2d.f)
- 2) the argument ACTION in the Fortran Open statement that will be used to open the file. ACTION may be READ, WRITE, or READWRITE. Here it is READWRITE because the results file is written, and in case of validation it is read at the end of the computation. It will be stored into WAC FILES(WACRES) %ACTION
- 3) the logical unit to open the file. This a priori value may be changed in case of code coupling. It is stored into WAC_FILES(WACRES) %LU

The **second** string, here WACRES, is the name of the file as it will appear in the temporary file where the computation is done.

The **third** string may be OBLIG (the name of the file must always be given), or FACUL (this file is not mandatory).

The **fourth** string (here BIN) says if it is a binary (BIN) or ASCII (ASC) file.

The **fifth** string is just like the READWRITE statement and is used by the perl scripts.

The **sixth** string is also used by the perl scripts and gives information on how the file must be treated. 'SELAFIN' means that the file is a Selafin format, it will have to be decomposed if parallelism is used. Other possibilities are:

SELAFIN-GEOM: this is the geometry file

FORTRAN: this is the Fortran file for user subroutines

CAS: this is the parameter file

CONLIM: this is the boundary conditions file

PARAL: this file will have an extension added to its name, for distinguishing between processors

DICO: this is the dictionary

SCAL: this file will be the same for all processors

The following sequence of subroutines is used for opening, using and closing files:

<u>Note:</u> subroutine INIT_FILES2 in BIEF version 5.9 has been renamed BIEF_INIT in version 6.0 and has from now on nothing to see with files.

1) opening files

IFLOT=0

CALL BIEF_OPEN_FILES(CODE,WAC_FILES,44,PATH,NCAR,COUPLAGE,IFLOT,ICODE)

CODE: name of calling program in 24 characters

WAC_FILES: the array of BIEF_FILE structures

44: the size of the previous array

PATH: full name of the path leading to the directory the case is

NCAR: number of characters of the string PATH

COUPLAGE: logical stating if there is a coupling between several programs.

IFLOT: in case of coupling, will be the last logical unit taken by a file

ICODE: code number in a coupling. For example in a coupling between TELEMAC-2D and GAIA, TELEMAC-2D will be code 1 and GAIA will be code 5.

2) using files:

Most operations on files consist on reading and writing, which always uses the logical unit. Every file has a name in the temporary folder where the program is executed, e.g. WACRES. The associated file number is an integer with the same name. The logical unit of this file will be equal to WACRES if there is no coupling, but more generally it is stored into WAC_FILES(WACRES)%LU. The logical unit of the geometry file in GAIA will be GAI_FILES(GAIGEO)%LU.

Sometimes the real name of files in the original is also used, for example to know if it exists (= has been given in the parameter file). This name is retrieved in the component NAME. For example the name of the geometry file in GAIA will be GAI_FILES(GAIGEO)%NAME.

3) closing files:

CALL BIEF_CLOSE_FILES(CODE,WAC_FILES,44,PEXIT)

CODE: name of calling program in 24 characters

WAC_FILES: the array of BIEF_FILE structures

44: the size of the previous array

PEXIT: logical, if yes will stop parallelism (in a coupling the last program calling BIEF_CLOSE_FILES will also stop parallelism).

A.5 Parallel computing in TOMAWAC

This document presents some basic concepts about the parallelism in TOMAWAC. It does not describe the parallel algorithm.

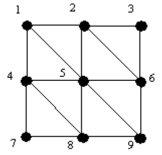
The parallelism in TOMAWAC is based on the message passing paradigm in order to run on a distributed memory architecture. The MPI library is used to manage the communication between parallel processors. Each MPI processor performs operations on its local memory independently from the other ones. The synchronisation and the data transfer between MPI processors are realized by sending or receiving messages.

Suppose that the parallel computing is performed on n MPI processus. The user has to add the following statement in the steering file:

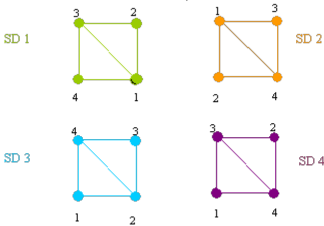
PARALLEL PROCESSORS = n

The domain (the finite element mesh) is split into n nonoverlapping subdomains. Each subdomain is assigned to a single MPI processor; in other words there is only one subdomain per MPI processor. The mesh partitioning is ensured by the PARTEL program included in the TELEMAC system. It generates the n local geometries and the boundary condition files required by the n MPI processus.

Let's consider a finite element mesh having 8 finite elements and 9 nodes



and suppose that the PARTEL program produces this following mesh partitionning (partition into 4 subdomains SD1 to SD4):



One could remark that a node does not have necessarily the same number in the global finite element mesh and in each subdomain. The node number in the global mesh is called global number whereas the node number in a subdomain is called local number.

There used to be two arrays (knogl and knolg) to link the global numbering and the local numbering, but since knogl could be huge if the number of processors was increasing, in version V7P1 we replaced the array knogl by a function global_to_local_point.

j=global_to_local_point(k,mesh) indicates that the local number of the node having global number k is j ("gl" means global numbering to local numbering, j=0 indicates that the node having global number k does not belong to the subdomain);

knolg(j)=k indicates that the global number of the node having local number j is k in the global mesh ("lg" means global numbering to local numbering).

It is important to understand that the global numbering does not exist in parallel: the user files **need to be modified** if global node numbering is used. Suppose that in the user file there is the following instruction:

c The D-value is set to 0.0 for the node number 200 D(200)=0.0

This instruction needs to be rewritten as follows to be used both in sequential or in parallel: IF (NCSIZE.GE.1) THEN

c NCSIZE is the number of MPI processus, so if NCSIZE > 1, the c code runs in parallel IF (global to local point(100,mesh) .NE. 0) THEN

c the node having global number 100 belongs to the subdomain

D(global_to_local_point(100,mesh))=0.0

END IF

ELSE

c here the code runs in sequential

D(100)=0.0

END IF

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