



Modelling the environmental behaviour of pollutants in Algeciras Bay (south Spain)

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ABSTRACT

An environmental study of Algeciras Bay is carried out through numerical modelling. First, a 2D barotropic model is applied to calculate tides and mean circulation. Results of this model are used by a sediment transport model which provides suspended matter concentrations and sedimentation rates in the Bay. It includes three particle classes. An effective diffusion coefficient has been calibrated simulating temperature distribution inside the Bay. An additional validation is obtained from an independent nitrate dispersion simulation. Then heavy metal dispersion patterns are investigated using a model which includes water–sediment metal interactions and uses the outputs of the hydrodynamic and sediment transport models. The metal transport model has been applied to simulate the dispersion of Zn, Cu and Ni. Results from the hydrodynamic, sediment and metal transport models have been compared with measurements. Model results also indicate that transport inside the Bay is relatively weak. Numerical experiments have been carried out to determine flushing times for conservative and non-conservative pollutants. Flushing time is about 20 days for a conservative tracer, and this value is mainly due to the M_2 residual current. Tides are not effective in removing pollutants.

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1. Introduction

Algeciras Bay (south Spain) is located in Gibraltar Strait, which connects the Atlantic Ocean and the Alboran Sea, in the Mediterranean (Fig. 1 top). Large amounts of pollutants are discharged into the Bay: urban waste from some towns and industrial wastes from one of the largest industrial areas in Andalusia, with a significant number of petrochemical plants and refineries, in addition to the production of steel, paper and power (thermal power plants). The Palmones and Guadarranque rivers empty into the Bay (Fig. 1 bottom), and the Bay is also polluted by heavy maritime traffic (Algeciras port is one of the largest in Spain).

In previous work (Morillo et al., 2008), it has been stated that water in the Bay has a high turnover because of its proximity to the Strait of Gibraltar, where the Mediterranean Sea and the Atlantic Ocean meet, and the strong currents that predominate in the area. However, some preliminary calculations with a low resolution model of the Strait of Gibraltar (Periañez, 2004b) indicated that contaminants released into the Bay tend to remain there. Only north winds are efficient for a rapid cleaning of the Bay.

The objective of this work consists of developing a high resolution numerical model of Algeciras Bay which may be used to understand the dynamics of pollutants in this environment, and to carry out a quantitative study of flushing times. In particular,

we are interested in assessing how water circulation affects such flushing times.

The model consists of a hydrodynamic module, which provides water circulation, a sediment transport model which provides suspended matter concentrations and sedimentation rates, and the pollutant dispersion model. The dispersion model may be applied to non-conservative pollutants (do not remain in solution). Uptake/release reactions between the dissolved and solid phases (suspended matter and bed sediments) are described in a dynamic way, using kinetic transfer coefficients. Results from the hydrodynamic and sediment transport models have been compared with measurements in the Bay. The pollutant dispersion model has been applied to simulate the behaviour of a conservative substance (nitrates) and to heavy metals. In the case of metals, computed concentrations in water and bed sediments are compared with measurements. Model results indicate that, in general, the dynamics of pollutants in the Bay has been correctly described. Then, numerical experiments are carried out to estimate flushing times for both conservative and non-conservative substances. The effects of the different tidal constituents and residuals on flushing times have been investigated. It is worth commenting that this is the first study on environmental conditions, pollutant dynamics and flushing carried out for Algeciras Bay with numerical modelling. Consequently this study completes previous field work carried out in the area (Morillo et al., 2008; Sánchez-Moyano et al., 2002; Guerra-García et al., 2009; Morales-Caselles et al., 2007; Morillo and Usero, 2008; Estacio et al., 1997).

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The three submodels (hydrodynamics, sediments, pollutants) are described separately in the next section, together with some indications on the numerical solution of equations. Later results are presented and discussed, again separately for each submodel. A study on flushing times closes the paper.

2. Model description

2.1. Hydrodynamic model

An important feature of the tidal flow in the Strait is that it can be considered, as a first approach, as barotropic. Indeed, 93% of the current velocity variance in the semidiurnal band has a barotropic character in the Strait (Mañanes et al., 1998). As a consequence, 2D depth-averaged models have already been applied to simulate tides in the Strait (Tejedor et al., 1999; Periañez and Pascual-Granged, 2008). Tsimplis et al. (1995) have even used a 2D barotropic model for simulating tides in the whole Mediterranean Sea. Other authors have also stated that using a 2D model for simulating tides is a reasonable approach (Dyke, 2001; Yanagi, 1999). Inside Algeciras Bay, it has been found that salinity and temperature are essentially homogeneous both horizontally and vertically (University of Cadiz, 2007). Thus, the use of a 2D depth averaged barotropic model is justified. It is based on the following equations:

$$\frac{\partial \zeta}{\partial t} + \frac{\partial}{\partial x}(Hu) + \frac{\partial}{\partial y}(Hv) = 0 \quad (1)$$

$$\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + v \frac{\partial u}{\partial y} + g \frac{\partial \zeta}{\partial x} - \Omega v + \frac{\tau_u}{\rho_w H} = A \left(\frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) \quad (2)$$

$$\frac{\partial v}{\partial t} + u \frac{\partial v}{\partial x} + v \frac{\partial v}{\partial y} + g \frac{\partial \zeta}{\partial y} + \Omega u + \frac{\tau_v}{\rho_w H} = A \left(\frac{\partial^2 v}{\partial x^2} + \frac{\partial^2 v}{\partial y^2} \right) \quad (3)$$

where u and v are the depth averaged water velocities along the x and y axis, h is the depth of water below the mean sea level, ζ is the displacement of the water surface above the mean sea level measured upwards, $H = h + \zeta$ is the total water depth, Ω is the Coriolis parameter ($\Omega = 2w \sin \lambda$, where w is the Earth rotational angular velocity and λ is latitude), g is acceleration due to gravity, ρ_w is water density and A is the horizontal eddy viscosity. τ_u and τ_v are friction stresses which have been written in terms of a quadratic law:

$$\tau_u = k \rho_w u \sqrt{u^2 + v^2} \quad (4)$$

$$\tau_v = k \rho_w v \sqrt{u^2 + v^2}$$

where k is the bed friction coefficient.

The solution of these equations provides the water currents at each point in the model domain and for each time step. Currents are treated through standard tidal analysis (Pugh, 1987, Chapter 4) and tidal constants are stored in files that will be read by the dispersion codes to calculate the advective transport. The model includes the two main tidal constituents, M_2 and S_2 . Thus, the hydrodynamic equations are solved for each constituent and tidal analysis is also carried out for each constituent separately. A residual transport cannot be produced by the pure harmonic currents given by the tidal analysis, thus tidal residuals have been calculated as well, as explained below. Diurnal constituents are not included since most of the variance of current velocities is given by the M_2 signal alone, S_2 being the second important constituent. Therefore they can be used to characterize a very significant fraction of tides in the area (Mañanes et al., 1998).

Once a stable periodic solution of hydrodynamic equations is achieved, tidal analysis is carried out to determine tidal constants that are used by the sediment and dispersion codes. Tidal residual

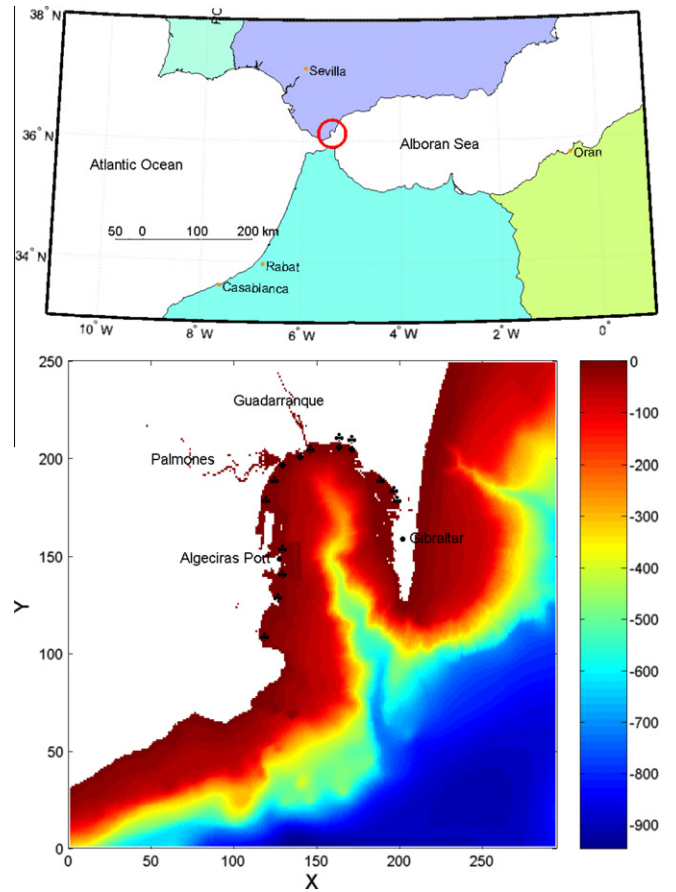


Fig. 1. Up: general localization of Algeciras Bay (inside the red circumference). Down: topography (m) of the Bay. Each unit in the axis corresponds to 100 m (grid cell spacing). \blacktriangle indicate urban waste release points and \diamond industrial waste release points. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

transports are also calculated. This is done for the M_2 and S_2 tides separately. Tidal residuals for each constituent are calculated from the equation:

$$\bar{q}_r = \frac{\langle H \bar{q}_t \rangle}{\langle H \rangle} \quad (5)$$

which corresponds to the Eulerian residual transport velocity (Delhez, 1996). In this equation $\langle \rangle$ is the time averaging operator, \bar{q}_r is the tidal residual and \bar{q}_t is the instantaneous tidal current. In addition, the mean current due to geostrophic flow is calculated from an additional run of the hydrodynamic model. In this run constant values of the sea surface elevation along the open boundaries of the domain are specified (Sannino et al., 2004; Periañez and Pascual-Granged, 2008). The total mean current is obtained adding the tidal residuals of the M_2 and S_2 constituents plus the mean geostrophic flow. Current \bar{q} at any time and position in the Strait is obtained adding this total mean current and the instantaneous tidal currents at the corresponding point. This is the water velocity value used in the dispersion calculations. This procedure is the same used in Periañez and Pascual-Granged (2008).

2.2. Sediment transport

The transport of sediments is described by a 2D advection-diffusion equation to which some terms are added. These are external sources of particles, terms describing particle deposition on the seabed and erosion from the bed to the water column.

The formulation of these processes is based upon standard formulae. Thus, the erodability constant is used for the erosion term. Particle deposition is described using the settling velocity, which is obtained from Stoke's law. Critical erosion and deposition stresses are applied as usually (Periañez, 2005a,b; Liu et al., 2002a; Lumborg and Windelin, 2003; Cancino and Neves, 1999).

Three particle classes are considered in the model, each one characterized by a mean size, to obtain a better representation of suspended particles dynamics and sedimentation in the Bay.

The equation for suspended sediment transport is:

$$\frac{\partial(Hm_k)}{\partial t} + \frac{\partial(uHm_k)}{\partial x} + \frac{\partial(vHm_k)}{\partial y} = \frac{\partial}{\partial x} \left(HK_h \frac{\partial m_k}{\partial x} \right) + \frac{\partial}{\partial y} \left(HK_h \frac{\partial m_k}{\partial y} \right) + (ER_k - DEP_k) + S_k \quad (6)$$

where $k = 1, 2, 3$ denote each sediment class, m_k is the suspended matter concentration, K_h is an effective horizontal diffusion coefficient (as explained in Section 3.2), S_k is the external particle source and ER_k and DEP_k are the erosion and deposition terms, respectively.

The deposition term is written in the following form:

$$DEP_k = w_k m_k \left(1 - \frac{|\bar{\tau}|}{\tau_{cd}} \right) \quad (7)$$

where $\bar{\tau}$ is the total bottom friction stress (tides plus mean flow) and τ_{cd} is a critical deposition stress above which no deposition occurs since particles are kept in suspension by turbulence. The settling velocity of particles is obtained from Stokes's law as commented above:

$$w_k = \frac{\rho - \rho_w}{\rho_w} \frac{g D_k^2}{18 \nu} \quad (8)$$

where ρ and D_k are suspended particle density and diameter, respectively, and ν is the kinematic viscosity of water.

The erosion rate is written in term of the erodability constant:

$$ER_k = E f_k \left(\frac{|\bar{\tau}|}{\tau_{ce}} - 1 \right) \quad (9)$$

where E is the erodability constant, f_k gives the fraction of k class particles in the bed sediment and τ_{ce} is a critical erosion stress below which no erosion occurs. The model can also calculate sedimentation rates (SR) as the balance between the deposition and erosion terms.

2.3. Pollutant transport

Non-conservative pollutants are those which do not remain dissolved in the water column, but have a certain affinity to be fixed to particles. If the pollutant is introduced in the water column, it will be fixed to settling suspended particles and their deposition on the sea bed will contaminate the bottom sediment. Of course there are also advection/diffusion processes in water and direct adsorption of pollutants on the seabed. The exchanges between the dissolved and solid phases may be described in terms of kinetic transfer coefficients. Thus, assuming that adsorption/release reactions are governed by a single reversible reaction, a coefficient k_1 governs the transfer from the liquid to the solid phase and a coefficient k_2 governs the inverse process. Also, the migration of contaminants to the deep sediment may be included. Thus, contaminants deposited on the sediment surface will be buried by particle deposition and will migrate below the mixed sediment layer which directly interacts with the dissolved phase (see for instance the discussion in Monte et al., 2006). This effect may be easily treated as a decay process with constant λ_{burial} given by Periañez (2009):

$$\lambda_{burial} = \frac{SR}{\rho_s L} \quad (10)$$

where L is the sediment mixing depth (the distance to which the dissolved phase penetrates the sediment) and ρ_s is the sediment bulk density (dry mass divided by wet volume). Nevertheless, this process is not significant in Algeciras Bay due to the low SR values and the typical simulation times (several months). Sediment bulk density is related to particle density through porosity p : $\rho_s = \rho(1 - p)$.

The adsorption process is a surface phenomenon that depends on the surface of particles per water volume unit into the grid cell. This quantity has been denoted as the exchange surface (Periañez, 2003a, 2004a, 2008, 2009). Thus in general:

$$k_{1,k} = \chi(S_{m,k} + S_{s,k}) = k_{1,k}^{spm} + k_{1,k}^{sed} \quad (11)$$

where $S_{m,k}$ and $S_{s,k}$ are the exchange surfaces for suspended matter and bottom sediments, respectively (dimensions $[L]^{-1}$) for the corresponding particle class k . χ is a parameter with the dimensions of a velocity. It is denoted as the exchange velocity (Periañez, 2004a, 2008, 2009).

Assuming spherical particles, the exchange surfaces are written as (see references cited above):

$$S_{m,k} = \frac{3m_k}{\rho R_k} \quad (12)$$

and

$$S_{s,k} = \frac{3L f_k (1 - p) \phi}{R_k H} \quad (13)$$

where R_k is particle radius, p is sediment porosity and ϕ is a correction factor that takes into account that part of the sediment particle surface may be hidden by other sediment particles. This formulation has been successfully used in all modelling works cited above. Real particles are not spheres, but with this approach it is possible to obtain an analytical expression for the exchange surface (Duursma and Carroll, 1996).

The equation that gives the temporal evolution of pollutant concentration in the dissolved phase, C_d , is:

$$\begin{aligned} \frac{\partial(HC_d)}{\partial t} + \frac{\partial(uHC_d)}{\partial x} + \frac{\partial(vHC_d)}{\partial y} = & \frac{\partial}{\partial x} \left(HK_h \frac{\partial C_d}{\partial x} \right) \\ & + \frac{\partial}{\partial y} \left(HK_h \frac{\partial C_d}{\partial y} \right) - H \\ & \times \sum_k \left(k_{1,k}^{spm} + k_{1,k}^{sed} \right) C_d + k_2 H \\ & \times \sum_k m_k C_{s,k} \\ & + k_2 L \rho_s \phi \sum_k f_k A_{s,k} \end{aligned} \quad (14)$$

where $A_{s,k}$ and $C_{s,k}$ are concentrations in bed sediment and suspended matter class k , respectively.

The temporal evolution of pollutant concentration in suspended particles is given, for each class k , by:

$$\begin{aligned} \frac{\partial(HC_s m)_k}{\partial t} + \frac{\partial(uHC_s m)_k}{\partial x} + \frac{\partial(vHC_s m)_k}{\partial y} = & \frac{\partial}{\partial x} \left(HK_h \frac{\partial(C_s m)_k}{\partial x} \right) + \frac{\partial}{\partial y} \left(HK_h \frac{\partial(C_s m)_k}{\partial y} \right) + k_{1,k}^{spm} C_d H \\ & - k_2 m_k C_{s,k} H + SED_k \end{aligned} \quad (15)$$

where SED_k expresses the pollutant exchange between suspended particles and the bed sediment resulting from erosion/deposition:

$$SED_k = \begin{cases} -SR_k C_{s,k} & SR_k > 0 \\ -SR_k A_{s,k} & SR_k < 0 \end{cases} \quad (16)$$

The equation for the temporal evolution of concentration in the bed sediment is:

$$\frac{\partial A_{s,k}}{\partial t} = k_{1,k}^{sed} \frac{C_d H}{L \rho_s f_k} - k_2 A_{s,k} \phi + SED_k \quad (17)$$

where now the exchange due to erosion/deposition of suspended particles is written as

$$SED_k = \begin{cases} \frac{SR_k C_{s,k}}{L \rho_s f_k} & SR_k > 0 \\ \frac{SR_k A_{s,k}}{L \rho_s f_k} & SR_k < 0 \end{cases} \quad (18)$$

The total concentration of pollutants in the sediment, A_{tot} , is computed from:

$$A_{tot} = \sum_k f_k A_{s,k} \quad (19)$$

and in the case of suspended matter it is:

$$C_s^{total} = \frac{\sum_k m_k C_{s,k}}{\sum_k m_k} \quad (20)$$

2.4. Numerical solution

All equations are solved using explicit finite difference schemes (Kowalick and Murty, 1993) on a grid with resolution $\Delta x = \Delta y = 100$ m. Second order accuracy schemes are used for advective and diffusive terms. Time step in the hydrodynamic equations, limited by the CFL condition, is $\Delta t = 0.5$ s. Since the sediment transport and pollutant dispersion models are run off-line, time step could be increased to 30 s.

Open boundary conditions consist of, in hydrodynamic calculations, specifying sea surface elevations compiled from observations and from a larger model covering the complete Strait of Gibraltar (Tejedor et al., 1999; Periañez and Caravaca, 2010).

The suspended matter model (without the erosion term) is started from a sea bottom containing no sediments. Then the accumulation of particles of each class is calculated to have a first estimation of the distribution of sediment particle sizes over the

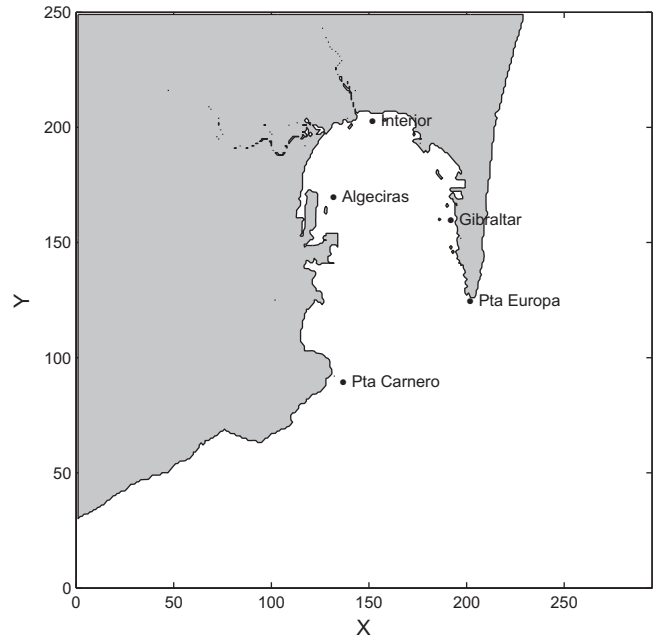


Fig. 3. Map indicating points where results from the hydrodynamic calculations have been compared with observations.

Table 1

Observed, index *obs*, and computed, index *comp*, amplitudes (A, m) and phases (ϕ , deg) of tidal elevations at several locations indicated in Fig. 3.

Station	M_2		S_2		M_2		S_2	
	A_{obs}	ϕ_{obs}	A_{comp}	ϕ_{comp}	A_{obs}	ϕ_{obs}	A_{comp}	ϕ_{comp}
Pta Carnero	0.311	47.5	0.318	46.3	0.115	71.0	0.119	71.2
Algeciras	0.310	48.0	0.291	45.5	0.111	73.9	0.112	72.4
Gibraltar	0.298	46.0	0.291	45.5	0.107	72.0	0.112	72.4

model domain. Next, the suspended matter model is started (with erosion) from the estimated particle class distribution until a

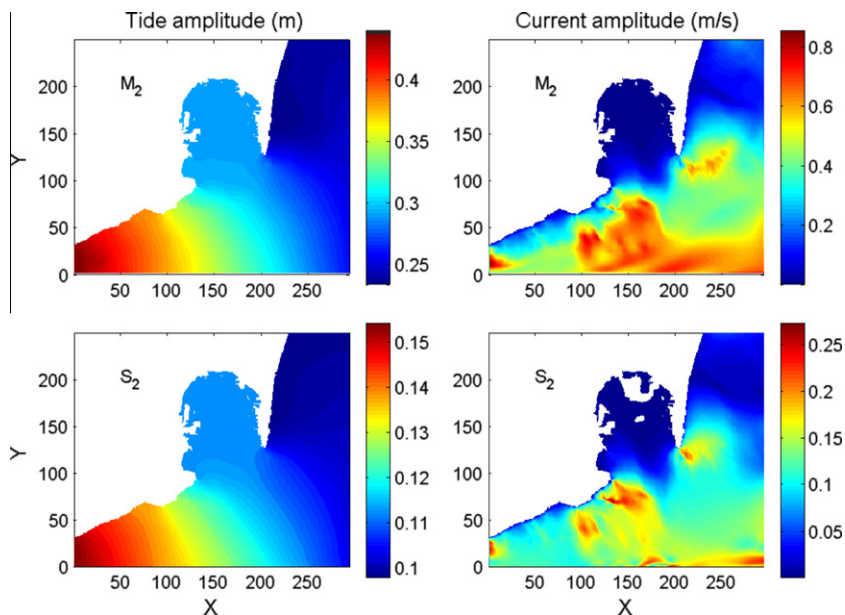


Fig. 2. Maps of computed tide amplitude and tidal current amplitude for both constituents included in the model.

Table 2
Observed, index *obs*, and computed, index *comp*, magnitudes (cm/s) of *u* and *v* tidal currents at some locations indicated in Fig. 3.

Station	M_2				S_2			
	u_{obs}	v_{obs}	u_{comp}	v_{comp}	u_{obs}	v_{obs}	u_{comp}	v_{comp}
Pta Europa	24.3	19.3	28.3	19.6	7.26	8.06	8.50	4.20
Pta Carnero	3.33	13.5	1.40	12.9	1.73	1.65	1.80	6.70
Interior	2.88	0.55	0.10	0.30	0.53	0.40	0.10	0.10

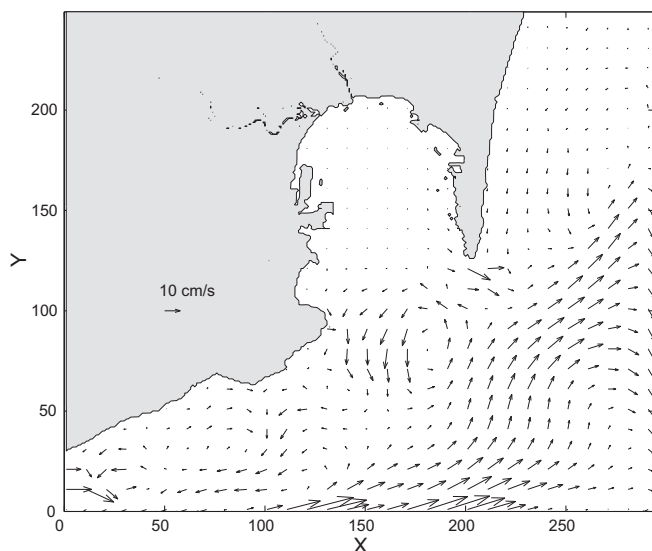


Fig. 4. Computed M_2 residual. Only one of each 100 computed vector is drawn for clarity.

steady state is reached. This way a self-consistent distribution of different particle classes, f_k on the sea bed over the model domain can be obtained. This procedure was successfully applied in Periañez (2005a). Suspended matter concentrations are defined along open boundaries for solving the sediment transport model (Dirichlet boundary condition) according to measured values in the Strait of Gibraltar (León-Vintró et al., 1999).

Once currents, suspended matter concentrations and sedimentation rates have been calculated and stored in files, the pollutant transport model may be run. In this case a no-gradient open boundary condition has been used.

3. Results

Results from the hydrodynamic, sediment transport and pollutant dispersion models are presented separately in the following subsections. Next, results from the flushing numerical experiments are discussed.

3.1. Hydrodynamics

As an example, computed tide amplitudes and tidal current amplitudes for M_2 and S_2 constituents are presented in Fig. 2. It may be seen that there is a significant reduction in the tide amplitude from the Strait into Algeciras Bay and towards the Mediterranean. Tidal currents are also weak inside the Bay, specially in the case of S_2 . Indeed, negligible currents are obtained for this constituent in the inner part of the Bay. A comparison between computed and measured (Tsimplis et al., 1995; Tejedor et al., 1999) tide amplitudes and phases at some points indicated in Fig. 3 may be

seen in Table 1. Current amplitudes are also compared with available observations (Universidad de Cadiz, 2007) in Table 2. It may be seen that, generally speaking, the model is giving a realistic representation of tides in the Bay. A map showing the computed M_2 residual transport velocity may be seen in Fig. 4. Currents are of the order of 10 cm/s in the Strait of Gibraltar, but extremely weak inside the Bay. It is interesting to notice the cyclonic gyre which is apparent at the mouth of Algeciras Bay.

3.2. Sediment transport

Winds and waves are not explicitly included in the hydrodynamic model. It is evident that they will produce an enhanced mixing inside the Bay (although waves inside the Bay are small, predominantly <0.5 m height; University of Cadiz, 2004). Thus an effective diffusion coefficient K_h has been defined, which would include mixing induced by wind and waves. To select its value, the water temperature distribution has been simulated. Water temperature T is described by an advection/diffusion equation in the form:

$$\frac{\partial T}{\partial t} + u \frac{\partial T}{\partial x} + v \frac{\partial T}{\partial y} = \frac{\partial}{\partial x} \left(K_h \frac{\partial T}{\partial x} \right) + \frac{\partial}{\partial y} \left(K_h \frac{\partial T}{\partial y} \right) \quad (21)$$

The model is started from an uniform water temperature over all the domain, but temperature is defined at the release points of the thermal power plants which are cooled with water from the Bay (these are points 4 and 6 in Fig. 5). The effects of these heat emissions are restricted to the shallow waters along the coastline. Thus, we assume that they do not affect the general circulation of the Bay and the barotropic approach is still valid. Indeed, it was commented in Section 2.1 that measurements of salinity and temperature showed generally homogeneous distributions (horizontally and vertically). These measurements were made in the main water body of the Bay and do not reflect the heat emissions from power plants.

The model is run until a steady temperature distribution is obtained. The diffusion coefficient K_h is changed by trial and error until the computed distribution agrees with measurements. When this occurs we would have the value of the effective diffusion coefficient which implicitly includes the average mixing induced by wind and waves. Such value resulted $10 \text{ m}^2/\text{s}$, and a comparison between measured (Guerra-García et al., 2009) and computed water temperatures may be seen in Fig. 5. This approach consisting of not including winds explicitly in the simulation has been successfully used in previous modelling works (Periañez, 2002, 2003a,b, 2005a, 2008, 2009). Although it may be a too simple approach, it can give a realistic view of the mean transport processes in the Bay, given the generally good agreement between measured and computed suspended sediment concentrations and metal distributions that will be shown below. However, the sensitivity of suspended sediment and pollutant transport in Algeciras Bay to changing winds is an interesting problem that has to be addressed in the next future, since dispersion processes related to meteorological conditions may become apparent. Moreover, erosion events may be induced by wind waves in the shallower areas. Nevertheless, wave-induced erosion is not included in the model since calculations are carried out under calm conditions.

Once the effective diffusion coefficient has been defined, sediment transport simulations may be carried out. Three particle classes have been used. Mean particle radius defining each class are 0.1, 1.0 and $15 \mu\text{m}$. The smallest class represents the background suspended sediment in the water column (as in Liu et al., 2002b). The following classes correspond to clay and medium silt, similarly to, for instance, Liu et al. (2002b), Jiang et al. (2000) and Holt and

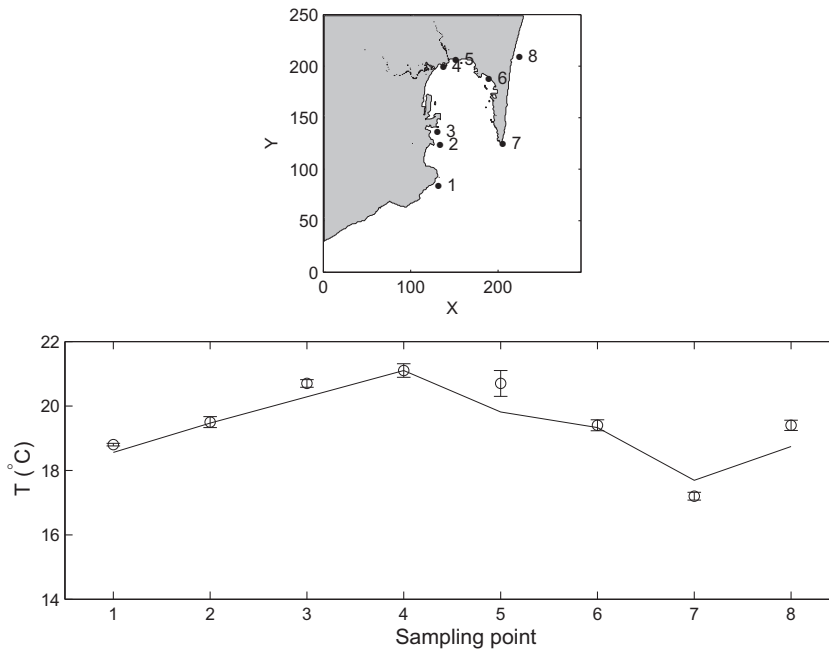


Fig. 5. Computed and measured water temperature along the Bay's shoreline. Measurement points are indicated on the top panel.

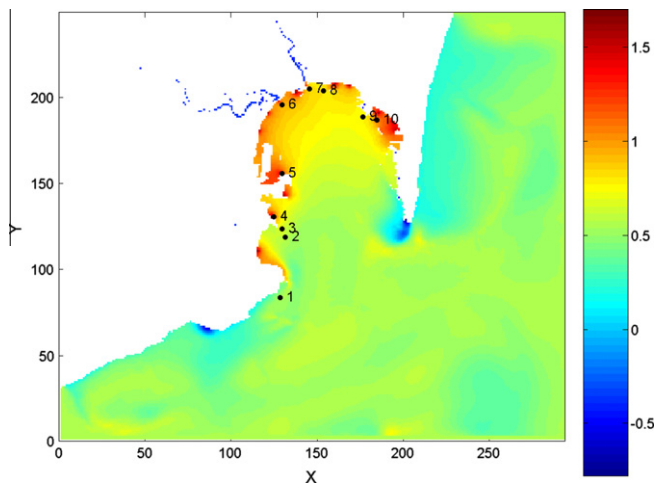


Fig. 6. Computed distribution of total suspended matter concentrations (g/m^3) over the domain. A logarithmic scale is used to appreciate details more clearly. Numbers indicate points where suspended matter concentrations and sedimentation rates have been compared with measurements.

James (1999). Settling velocities are then obtained from Stoke's law, as mentioned above.

Other parameters are required to run the sediment transport module. The critical deposition stress typically ranges between 0.04 and 0.1 N/m^2 (Tattersall et al., 2003), while the critical erosion stress ranges 0.1–1.5 N/m^2 (Tattersall et al., 2003). In the present application intermediate values of 0.06 and 1.0 N/m^2 have been taken for the critical deposition and erosion stresses, respectively, as in the models for southern Spain described in Perriñez (2008, 2009). The erodability constant is fixed as $E = 1.6 \times 10^{-3} \text{ kg}/\text{m}^2\text{s}$. This parameter typically varies between 2×10^{-4} and $3 \times 10^{-3} \text{ kg}/\text{m}^2\text{s}$ (Tattersall et al., 2003). A standard value $\rho = 2600 \text{ kg}/\text{m}^3$ has been used for particle density.

Sediment discharges by Palmones and Guadarranque rivers, as well as from waste release have been defined by trial and error,

until the model showed a reasonable agreement with observations. Of course, these magnitudes present seasonal (and at a shorter scale as well) variations. However, these are not quantified and our objective has consisted, simply, of having a correct estimate of mean suspended matter concentrations over the domain to be able to simulate the dispersion of reactive pollutants. It must be considered that available sediment concentrations and SR to be compared with model output are mean (annual) averages (Sánchez-Moyano et al., 2002), which justifies our approach.

The sediment model is started from an uniform suspended matter concentration of $3 \text{ g}/\text{m}^3$ (total) and integrated until a steady state is achieved (see also Section 2.4). The computed distribution of suspended sediments over the domain may be seen in Fig. 6 and a comparison between computed and measured (Sánchez-Moyano et al., 2002) suspended matter concentrations and sedimentation rates is presented in Fig. 7. The distribution in Fig. 6 shows an increase in suspended matter towards the interior of the Bay due to inputs from rivers and urban wastes, as stated in Sánchez-Moyano et al. (2002). The magnitude of suspended sediment concentration is in general correctly predicted by the model, except along the southwest shore of the Bay (points 1 and 2), where suspended matter concentration is underestimated. Sedimentation rates are, in general, underestimated. This is not surprising given the approximations made. Although constant suspended matter concentrations have been defined in the rivers, there will be seasonal variations depending on pluviometry, for instance. Heavy rain episodes will enhance mean sedimentation rates, and this is not described in the model. Sediment bed load transport is neglected. Finally, atmospheric deposition events of particles coming from the Sahara Desert have not been considered in the model since they cannot be easily quantified (as in Perriñez, 2008).

3.3. Pollutant transport

The dispersion of metals Zn, Cu and Ni has been simulated since there are field measurements which can be used to test the model results. These three metals were studied in a previous modelling work of the Gulf of Cadiz (Perriñez, 2009), at the west side of Gibraltar Strait. Thus, the same parameters as in such model have

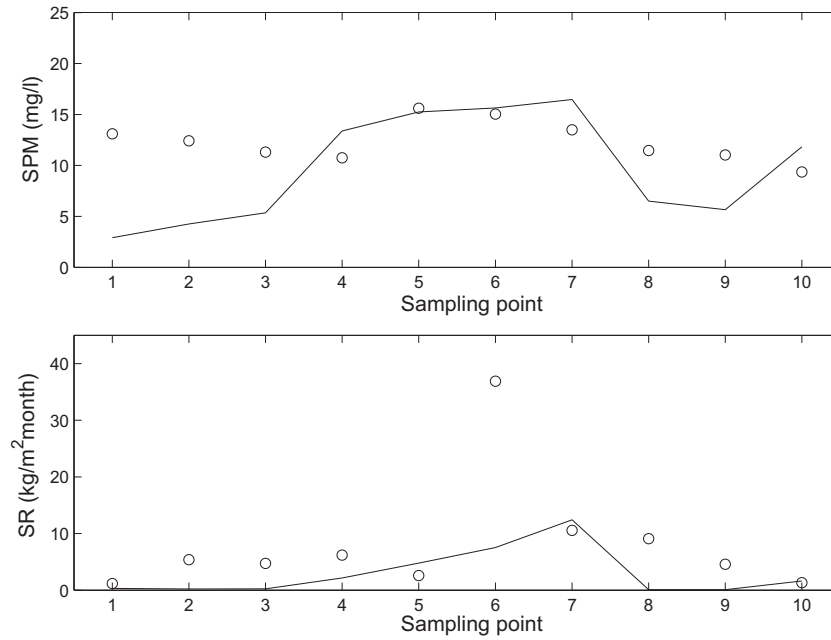


Fig. 7. Comparison between measured (points) and computed (lines) suspended matter concentrations and sedimentation rates. Sampling points are indicated in Fig. 6. Measured values correspond to annual mean values (one sample per month).

Table 3
Summary of model parameters. Details and justification may be seen in Periañez (2009).

Parameter description	Value
sediment mixing depth	$L = 0.1 \text{ m}$
particle density	$\rho = 2600 \text{ kg/m}^3$
correction factor	$\phi = 0.1$
sediment porosity	$p = 0.5$
desorption kinetic coefficient	$k_2 = 1.16 \times 10^{-5} \text{ s}^{-1}$
Zn distribution coefficient	$k_d = 7.0 \times 10^4$
Ni distribution coefficient	$k_d = 2.0 \times 10^4$
Cu distribution coefficient	$k_d = 2.6 \times 10^4$

been used in the present work. Essentially, we require the exchange velocity χ and the kinetic coefficient k_2 for each metal. Also, a mean porosity, sediment mixing depth L and correction factor ϕ have to be specified. The exchange velocity is deduced from the equilibrium partition coefficient k_d as described in detail in Periañez (2009). Values for the different parameters are summarized in Table 3, and detailed justification may be seen in Periañez (2009), thus it is not repeated here. Sensitivity analysis of model formulation to these parameters may also be seen in such reference.

Nevertheless, a first numerical experiment has been carried out applying the model to simulate the dispersion of a conservative contaminant (thus $\chi = k_2 = 0$). Releases from Acerinox plant have been simulated. It is a stainless steel production plant located between Palmones and Guadarranque rivers. Wastes released to the Bay contain significant amounts of nitrates (University of Cadiz, 2007). The model is started from zero concentrations over the whole domain and nitrate concentration in the release point from Acerinox has been fixed as 11.5 mg/l, according to University of Cadiz (2007). A simulation over 40 days is carried out. In this time steady state is reached. Model results are compared with the observed nitrate distribution in September 2006, when winds in the area were weak. It must be commented that concentration in the release point, 11.5 mg/l, is the measured one in this date. A

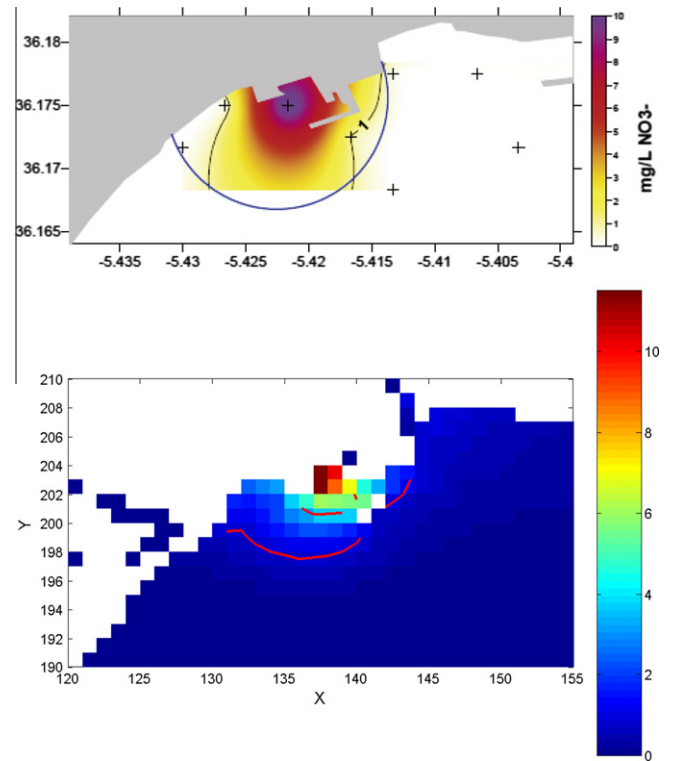


Fig. 8. Top: measured nitrate concentrations. The 1 mg/l concentration isoline is shown, as well as the 0.5 miles circumference centered in Acerinox. Bottom: computed concentrations. Lines are the 5 (closer to the source) and 1 mg/l isolines.

comparison between the observed and calculated distributions may be seen in Fig. 8. Measurements indicate that the 1 mg/l isoline is within a 0.5 miles radius circumference centered in the industrial plant. Model results are in agreement with these observations. The nitrate patch has an almost circular shape, due to the

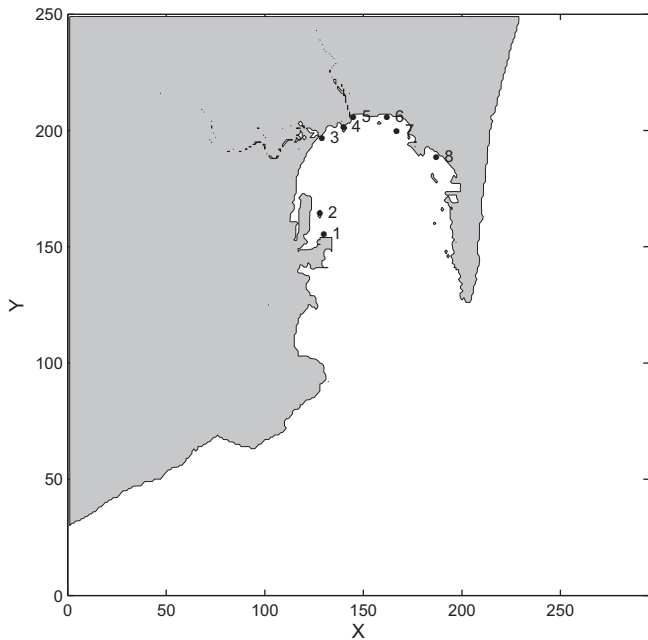


Fig. 9. Map indicating points where metal concentrations in water and sediments have been compared with measurements.

very weak currents in this part of the Bay. Also, the dilution degree, in close agreement with the observed one, confirms the correct calibration of the effective diffusion coefficient previously carried out through the temperature simulations.

To simulate reactive metal dispersion the model is again started from zero concentrations in water, suspended matter and bed sediments and integrated until steady concentrations are obtained over the domain in the three phases. In practice, simulations over some 100 days are required, longer than in the case of nitrates because of water–sediment interactions and the presence of multiple sources. Metal concentrations in water and bed sediments are

compared with measurements in the Bay (Morillo et al., 2008; Morales-Caselles et al., 2007). A mean metal discharge, in dissolved form, is introduced from industrial and urban waste sources. For each metal, the same magnitude is used for all urban sources and for all industrial sources. These average inputs were selected by calibration. Of course, this is a very rough approach: inputs are not the same for all sources and will present temporal variability. However, it allows obtaining an estimation of average metal concentrations in the Bay and it may also be observed if the metal partition between water and sediment is correctly reproduced by the model. Using a different source magnitude for each release point, obtained from calibration, may look tricky since such magnitudes could be fitted to improve model agreement with observations. Thus it was decided to use the same magnitude for sources of the same nature (industrial or urban), although this will clearly make model/measurement comparisons worse. It must finally be taken into account that sediments integrate contamination over a long temporal scale, thus short period fluctuations are not relevant. This approach to the source term has been used in previous modelling works (Periañez, 2002; Periañez et al., 2005).

Points where metals have been measured in the dissolved phase and bed sediments are shown in Fig. 9. A comparison between calculated and measured concentrations may be seen in Figs. 10–12 for Zn, Cu and Ni, respectively. The same pattern is obtained in the two phases for the three metals. This is a logical result since the bed sediment is more contaminated as concentration in water over it increases. However, it is worth mentioning that, in spite of the procedure used to define the source term, realistic concentration levels are in general simultaneously obtained for the dissolved phase and the sediment, and for the three metals. This indicates that metal partition between the dissolved and solid phases has been correctly described.

Metal distribution over the Bay may be seen, in the case of Zn as an example, in Figs. 13–15, where the computed distributions in water, suspended matter and bed sediments are presented. Concentrations decrease quickly, even inside the Bay, specially in the case of the dissolved phase. This confirms the weak transport inside the Bay which was observed when simulating nitrates. Metals

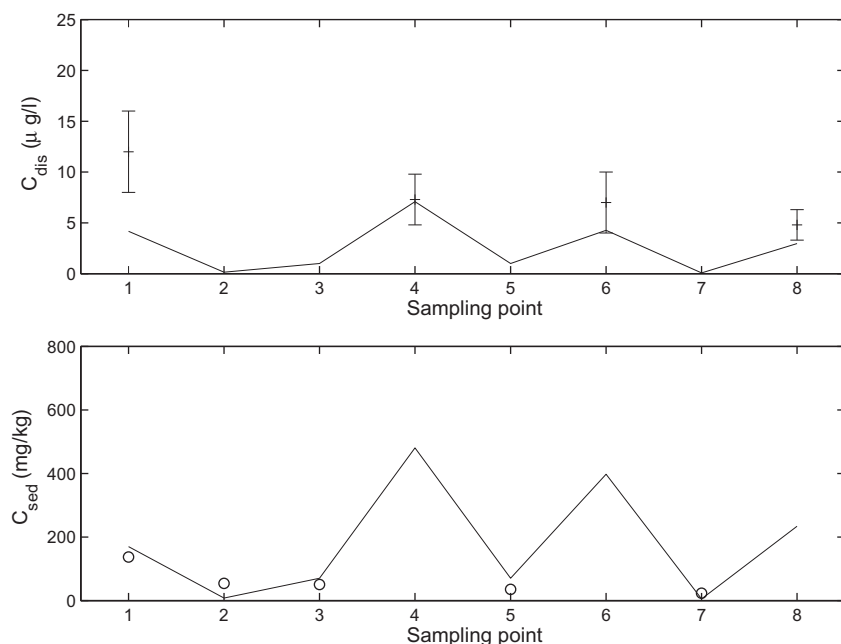


Fig. 10. Computed (lines) and measured (points) Zn concentrations along the Bay's shoreline in water (top) and bed sediments (bottom). Measurement points are indicated on Fig. 9.

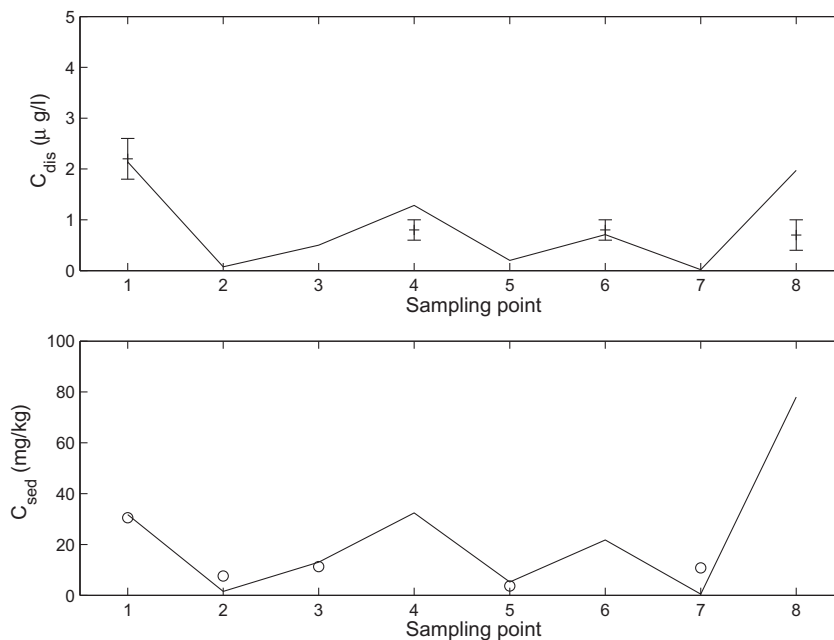


Fig. 11. Computed (lines) and measured (points) Cu concentrations along the Bay's shoreline in water (top) and bed sediments (bottom). Measurement points are indicated on Fig. 9.

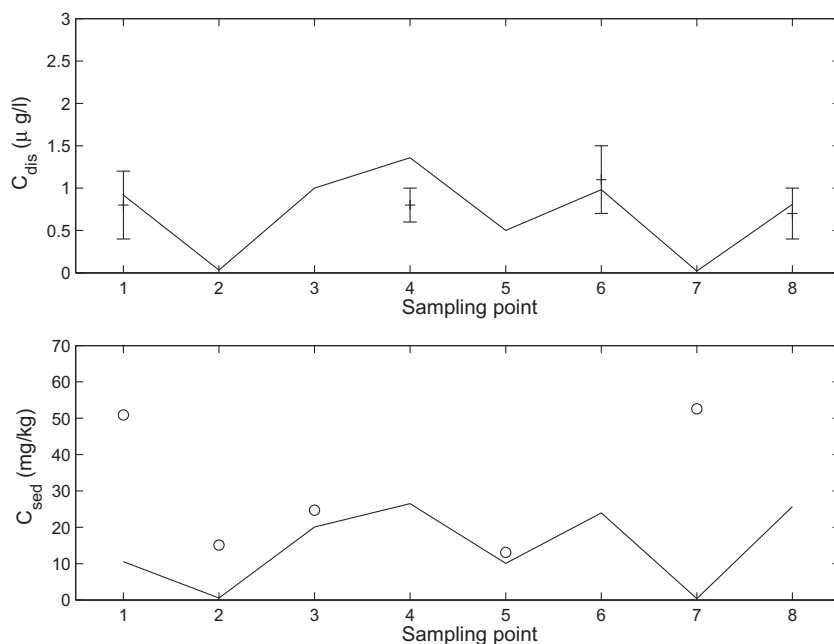


Fig. 12. Computed (lines) and measured (points) Ni concentrations along the Bay's shoreline in water (top) and bed sediments (bottom). Measurement points are indicated on Fig. 9.

which leave the Bay are mainly directed to the east (Mediterranean Sea) due to the prevailing residual currents in the Strait (Periañez and Caravaca, 2010). However, tidal mixing in the Strait makes a metal fraction to move to the west as well. High metal concentrations in suspended matter ($\sim 10^2$ mg/kg) are due to the low particle concentrations (Fig. 6), but a very significant portion of the metal content, per water volume, is in dissolved form. Bed sediments are contaminated as water and suspended matter containing metals travel above them. A smooth distribution is obtained outside the Bay (Fig. 15) since sediments integrate long-scale processes as commented above.

Model results, however, have to be interpreted with care (as commented before for suspended matter transport) since plumes of suspended particles and hence contaminants are sensitive to changes in wind speed and direction as well as they depend on pluviometry. Moreover, other meteorological conditions, such as atmospheric pressure differences between the Atlantic and Mediterranean, can induce flow variations through the Strait of Gibraltar. These facts make the comparison of computed metal concentrations in the water column with the corresponding measurements specially difficult (as already discussed by Dyke, 2001). However, bed sediments integrate all this variability and

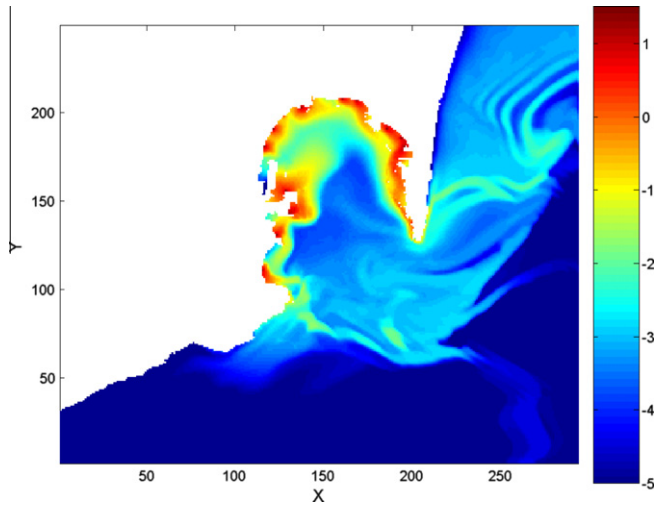


Fig. 13. Computed Zn concentrations in the dissolved phase (µg/l). A logarithmic scale is used to appreciate details more clearly.

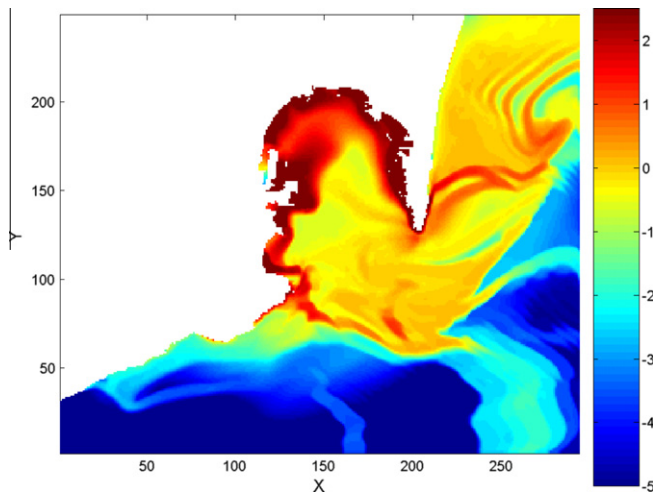


Fig. 14. Computed Zn concentrations in suspended matter (mg/kg). A logarithmic scale is used to appreciate details more clearly.

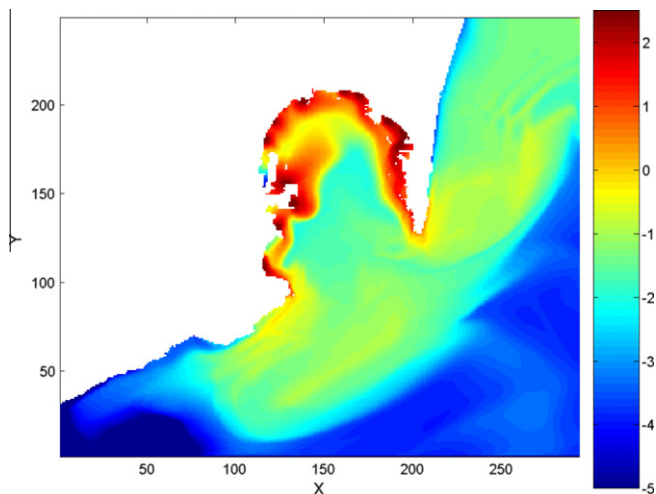


Fig. 15. Computed Zn concentrations in bed sediments (mg/kg). A logarithmic scale is used to appreciate details more clearly.

the generally good behaviour of the model in reproducing sediment contamination gives some confidence on the model.

3.4. Bay flushing times

Some numerical experiments have been carried out to estimate Algeiras Bay flushing times. Flushing time is defined as the time in which the tracer inventory in the water column decreases in a factor e (Prandle, 1984) and the sediment half-life is defined as the time in which the tracer inventory in the sediment decreases in a factor 2 (Periañez, 2003b). These parameters are relevant for the water quality of a system, and it is important to know the time scale for a pollutant discharged into a water body to be transported out of the system (Shen and Haas, 2004). Thus, flushing times have been recently determined for a number of water bodies using numerical experiments (Shen and Haas, 2004; Choi and Lee, 2004). On the other hand, it is also known that a contaminated sediment may act as a long-term delayed source of previously released contaminants (Cook et al., 1997). Consequently, it is also relevant to have estimations of the sediment half-life (Periañez, 2003b).

Initially, some experiments have been carried out with a conservative pollutant, remaining dissolved, to assess the effects of tidal constituents and residuals in flushing times. Later, flushing times in the case of a reactive contaminant (Zn has been used as an example) are investigated.

In the case of a conservative tracer, model initial conditions consist of setting a 100 units/m³ concentration at every grid cell inside the Bay and zero concentration elsewhere. The model is integrated and the time evolution of the pollutant inventory inside the Bay written to an output file. Fitting to exponential decay curves provides the flushing time. Thus, the system-wide (as defined by Choi and Lee, 2004) flushing time has actually been determined. These authors found that this parameter is useful for determining the long-term water quality of a system.

Choi and Lee (2004) have found that the system-wide flushing time is better obtained from a double exponential decay curve rather than a single exponential decay. The temporal evolution of the mass within the system, $M(t)$, is thus given by the following equation:

$$\frac{M(t)}{M_0} = (1 + \beta)e^{-\alpha_1 t} - \beta e^{-\alpha_2 t} \quad (22)$$

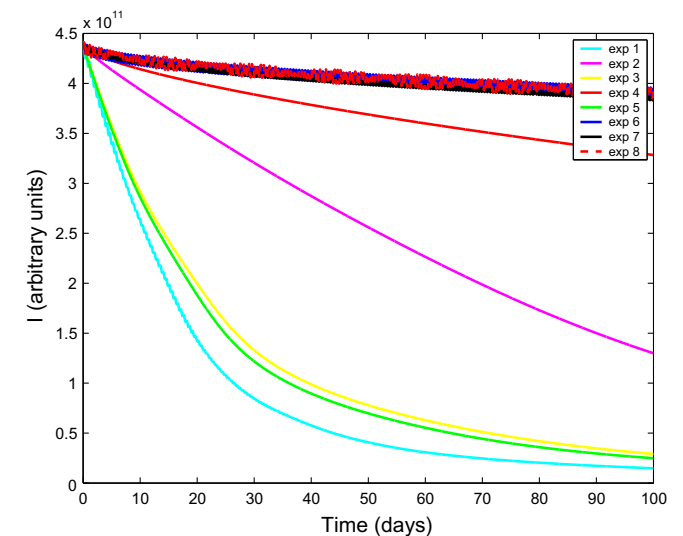


Fig. 16. Temporal evolution of the conservative pollutant inventory I inside the Bay for the flushing experiments.

Table 4

Flushing times (days) for the different experiments involving a conservative pollutant. The regression coefficient of the numerical fitting to the double exponential decay function is also given.

	Experiment							
	1	2	3	4	5	6	7	8
M_2 tide	✓					✓		✓
M_2 residual	✓		✓		✓			
S_2 tide	✓						✓	✓
S_2 residual	✓			✓	✓			
Sea level current	✓	✓						
T_f (days)	19.2	81.5	27.6	310	25.6	677	617	680
r^2	0.9904	0.9993	0.9825	0.9323	0.9846	0.8700	0.9490	0.8717

where M_0 is the initial mass within the system. If the three parameters β , α_1 and α_2 are determined from numerical fitting, the system-wide flushing time is given by Choi and Lee (2004):

$$T_f = \frac{1 + \beta}{\alpha_1} - \frac{\beta}{\alpha_2} \quad (23)$$

The temporal evolution of pollutant inventories inside the Bay for all the experiments may be seen in Fig. 16. Experiments are summarized in Table 4, where computed flushing times are given together with the r^2 coefficient for each fitting, carried out at 95% confidence level.

Depending on the assumed water circulation, flushing times range from some 20 to almost 700 days. If tides and residuals are included in the simulation (exp 1), the flushing time results 19.2 days. Similar values are obtained in exp 3 and 5, which include the M_2 residual and the M_2 plus S_2 residuals, respectively. This indicates that flushing, and thus cleaning of the Bay, is primarily controlled by the M_2 residual. Flushing induced by the S_2 residual alone (exp 4) is weak (310 days) and the produced by sea-level difference (geostrophic) induced currents (exp 2) is intermediate. Flushing times due to the tidal constituents by themselves are long (exp 6, 7 and 8), over 600 days, due to the forward and backward periodic currents produced by the tides, which make them little efficient to flush pollutants.

In summary, flushing time of Algeciras Bay is of the order of 20 days, and this value is mainly due to the M_2 residual current. The second most important component of water circulation in inducing flushing off the Bay is current produced by sea level differences between the Atlantic and Mediterranean.

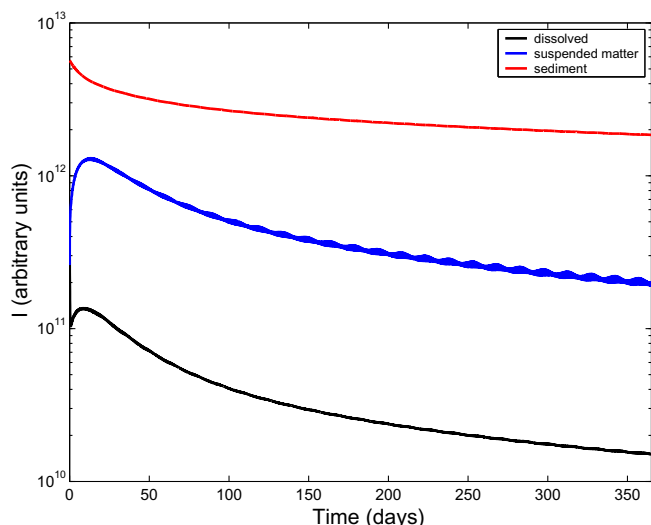


Fig. 17. Temporal evolution of Zn inventory I inside the Bay for the three phases.

In the experiment carried out for a reactive pollutant (Zn) initial conditions are the same as above for the dissolved phase, but concentrations in suspended matter and bed sediments are assumed to be initially at equilibrium (through the corresponding k_d , see Table 3) with the dissolved phase. All components in water circulation are considered (as exp 1 in the conservative pollutant simulations). A simulation over one year has been carried out. Evolution of total inventories in each phase may be seen in Fig. 17. Flushing times for the dissolved phase and for contaminants attached to suspended particles are, respectively, 110 and 144 days. Thus, they are considerably longer than in the case of a conservative pollutant, as should be expected. Sediment half-time is 316 days, indicating that sediments retain contaminants which then are slowly released. This sediment half-time is similar to that of Huelva estuary (southwest Spain) for ^{226}Ra (510 days; Perriñez et al., 2005), although this element is more mobile in the environment than Zn. In a more energetic environment as the English Channel, sediment half-time, in the case of plutonium (whose mobility is closer to that of Zn than in the case of Ra), is 70 days (Perriñez, 2003b), significantly shorter than in Algeciras Bay.

We can conclude that flushing of the Bay is not as fast as could be expected due to the strong currents in the Strait of Gibraltar, as was mentioned in Morillo et al. (2008).

4. Conclusions

A numerical model has been developed to study the behaviour of pollutants in Algeciras Bay. The model includes a hydrodynamic module, a sediment transport module and the dispersion module itself.

Results from the hydrodynamic model have been compared with observed tides and currents in the domain. The sediment transport model includes three particle sizes for a better representation of suspended particle dynamics. An effective diffusion coefficient, implicitly including mixing by winds and waves, has been used. It has been obtained through a calibration process simulating the temperature distribution in the Bay. Calculated suspended matter concentrations are in relative agreement with observations. A further validation of the calibrated effective diffusion coefficient has been obtained through the simulation of the dispersion of a conservative pollutant (nitrates released from a stainless steel production plant). Its value is $10 \text{ m}^2/\text{s}$.

Water–sediment interactions in the dispersion model have been described in a dynamic way, using kinetic transfer coefficients. Calculated Zn, Cu and Ni distribution in water and sediments are, generally speaking, in agreement with observations.

Model results indicate that transport inside the Bay is relatively weak. Indeed, the nitrate simulation shows that most of the contaminant plume stays close to the source. In the case of metals there is a strong concentration gradient inside the Bay, which again is an indication of relatively weak mixing. Pollutants which are

able to leave the Bay are mainly transported towards the Mediterranean Sea by the dominant mean currents in the area.

Some numerical experiments have been carried out to estimate flushing times of the Bay for conservative and non-conservative tracers, as well as sediment half-times. In the case of a conservative tracer, it has been found that flushing time of Algeciras Bay is of the order of 20 days, and this value is mainly due to the M_2 residual current. The second most important component of water circulation in inducing flushing off the Bay is current produced by sea level differences between the Atlantic and Mediterranean. Tides are not effective in removing pollutants due to the periodic forward/backward motion which they produce. Flushing times in the case of a reactive pollutant are significantly longer.

Acknowledgement

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