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Numerical modeling of the acoustic wave propagation across an homogenized rigid microstructure in the time domain

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Abstract

Homogenization of a thin micro-structure yields effective jump conditions that incorporate the geometrical features of the scatterers. These jump conditions apply across a thin but nonzero thickness interface whose interior is disregarded. This paper aims (i) to propose a numerical method able to handle the jump conditions in order to simulate the homogenized problem in the time domain, (ii) to inspect the validity of the homogenized problem when compared to the real one. For this purpose, we adapt the Explicit Simplified Interface Method originally developed for standard jump conditions across a zero-thickness interface. Doing so allows us to handle arbitrary-shaped interfaces on a Cartesian grid with the same efficiency and accuracy of the numerical scheme than those obtained in an homogeneous medium. Numerical experiments are performed to test the properties of the numerical method and to inspect the validity of the homogenization problem.

\textbf{Keywords:} homogenization, effective jump conditions, time-domain wave propagation, immersed interface method, ADER scheme.

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1. Introduction

The description of the interaction of waves with many scatterers of size much smaller than the wavelength is in principle simple since the scattering is weak, and several approximated methods can be applied, owing to a small parameter being the scattering strength. In the case of many scatterers located periodically, homogenization techniques are well adapted to handle the problem within a rigorous mathematical framework. The classical homogenization of massive media predicts that scatterers occupying a large area can be replaced by an equivalent homogeneous and in general anisotropic medium occupying the same area [5]. The case of scatterers located periodically along a mean line $\Gamma$ (Fig. 1(a)) has been less regarded but it is nowadays accepted that jump conditions across an equivalent interface have to be thought (Fig. 1(b)). Such jump conditions can be established using homogenization techniques which basically rely on the same ingredients than the classical homogenization of massive media, see e.g. [2, 3, 6, 7]; we call them interface homogenizations.

![Figure 1: (a) The real problem involves a structured film composed of a periodic set of sound-hard scatterers in a homogeneous fluid, (b) the equivalent homogenized problem involves a thick interface across which jump conditions apply. The wavefield, on which the jump conditions apply, is not represented.](image)

When the numerical resolution of the problem is sought, such equivalent media are of particular interest since they avoid to mesh the very small scales which are essential in the real problem. Indeed, these small scales define the scales of variation of the evanescent near wavefield excited in the vicinity of the scatterers. This is particularly true for scatterers located along a line since the resulting film is dominated by the effect of the near field, or in other words by boundary layer effects. In the case of sound hard scatterers, such interface homogenization has been proposed recently in [18, 19, 20]. These works follow those developed in
solid mechanics [22] and they are adapted to the wave equation in the harmonic regime [18, 19] and in the time domain [20].

The goal of the present paper is twofold. First, we propose an accurate time-domain numerical scheme for the homogenized problem, thus incorporating the jump conditions across the equivalent thick interface. Next, we inspect the validity of the homogenized result; notably we exemplify the robustness of the jump conditions with respect to the ratio of the array spacing with the typical wavelength and with respect to a possible curvature of the line $\Gamma$. Concerning the proposed numerical scheme, specific aspects will be addressed:

- the capability to handle arbitrary-shaped interfaces on a Cartesian grid, without introducing spurious diffractions due to a naive stair-step discretization of the interfaces;
- the accuracy of the numerical scheme, despite the non-smoothness of the solution across the thick interface;
- the performances of the scheme in terms of computational cost; specifically, the scheme has to guaranty an additional cost in the homogenized problem (due to the treatment of the jump conditions) which is highly negligible compared with the computational cost in the real problem.

To do so, cartesian grid methods such as the Immersed Interface Method (IIM) constitute an attractive strategy. The IIM modifies the stencil at the grid nodes near the interfaces. It was originally developed for elliptic and parabolic equations [10, 11, 12], to handle e.g. flows with irregular domains. Then it has been applied to linear hyperbolic problems in [28]. An overview may be found in the review book [13].

Alternative approaches have been developed in the context of wave propagation by one author of the present paper. The Explicit Simplified Interface Method (ESIM) does not modify explicitly the stencil, which may be a tedious task for a high-order scheme. On the contrary, the numerical values used by the scheme are modified, based on the jump conditions, the conservation laws and the local geometry. Generalization to different constitutive laws on both sides of the interface can be handled in a naturel way, contrary to the IIM. This approach has been applied to a broad class of problems, such as acoustics / elastodynamics [14], seismology [16], poroelasticity [4]. Modeling of imperfect contacts between solids has also been investigated by this approach [14].

The extension of the ESIM to the present homogenized problem involves two ingredients. First, it requires to implement a generalized version of the usual
boundary conditions at an interface, able to involve both the field and its spatial
derivatives. Second, it must handle an interface with a non-zero thickness: the
values of the solution on both sides of the interface are linked together (via the
jump conditions), whereas no field is computed inside the thin interface.

The paper is organized as follows. In section 2, the actual and the homog-
enized problems are presented. Section 3 details the numerical methodology: a
fourth-order ADER scheme combined with the ESIM, this latter constituting the
core of the work. Section 4 presents numerical experiments. Comparisons with
exact solutions confirm the efficiency and the accuracy of the numerical model-
ing. Next, comparisons with direct simulations for a real microstructure confirm
the second-order accuracy of the effective model. Lastly, some perspectives are
drawn in section 5.

2. The real problem and its homogenized version

The real problem concerns the propagation of acoustic waves through sound
hard scatterers located periodically onto a mean line $\Gamma$ (Fig. 1(a)). Periodically
means a constant spacing $h$ between two scatterers defined by the arc length along
$\Gamma$. The resulting curved array is surrounded by a fluid with mass density $\rho$
and isentropic compressibility $\chi$. In the fluid, the linearized Euler equations apply and
Neumann boundary conditions for the pressure apply for sound-hard scatterers.
Denoting $\Omega$ the computational domain containing the array of scatterers, the real
problem consists to solve in $\Omega$

$$
\begin{align*}
\rho \frac{\partial V}{\partial t} &= -\nabla p, \\
\chi \frac{\partial p}{\partial t} &= -\nabla \cdot V, \\
\nabla p \cdot N &= 0 \quad \text{on the boundaries of the scatterers,}
\end{align*}
$$

with $V = (v_x, v_y)^T$ the acoustic velocity and $p$ the acoustic pressure ($\chi$ is often
written in terms of the sound speed $c$: $\chi = (\rho c^2)^{-1}$). The Neumann boundary con-
dition $\nabla p \cdot N = 0$, with $N$ the vector locally normal to the scatterer boundaries,
accounts for a large mass density of the scatterers; it results a vanishing normal
velocity at their boundaries.

If the central wavelength (or the minimum one) $\lambda_0$ imposed by the wave source
is much larger than $h$, the real problem can be replaced by an equivalent homoge-
nized problem, owing to the introduction of the small parameter
\[ \varepsilon \equiv 2\pi \frac{h}{\lambda_0}. \]  

(2)

In [20], such a homogenization has been proposed; the problem ends up with jump conditions for the pressure and for the normal velocity. In this reference, the derivation is performed for a line \( \Gamma \) being straight (Fig. 2(a)). Here, we heuristically extend this result to a curved line \( \Gamma \), just by replacing locally the jump conditions expressed in Cartesian coordinates by jump conditions expressed in the local coordinates defined by the normal and tangent vectors to \( \Gamma \) (Fig. 2(b), and we shall discuss the validity of this extension in this paper).

Next, in [20] (see also the previous works of [6, 7]), the homogenized problem is shown to be associated with a satisfactory equation of energy conservation if the jump conditions are expressed across an “enlarged” interface of interior \( \Omega_e \) delimited by the two lines \( \Gamma^- \) and \( \Gamma^+ \). Doing so allows us to define a positive interface energy, thus guaranties notably the well-posedness in the time domain, which is necessary for our present purpose. The thickness of the interface is a priori arbitrary as soon as it guaranties a positive interface energy. In this paper, following [20], it is taken equal to the thickness \( e \) of the array of Neumann scatterers. We

Figure 2: In the homogenized problem, the array of sound-hard scatterers located periodically along a mean line \( \Gamma \) is replaced by a domain \( \Omega_e \). \( \Omega_e \) is delimited by the lines \( \Gamma^- \) and \( \Gamma^+ \) across which the jump conditions (3) apply. (a) in [20], the jump conditions are expressed for a straight line \( \Gamma \). (b) we extend these conditions to a curved line \( \Gamma \); \( \Gamma^- \) and \( \Gamma^+ \) are deduced from \( \Gamma \) by a translation of \( \pm e/2 \) along the normal vector \( n \) to \( \Gamma \); the normal and tangent vectors to \( \Gamma^\pm \) at the resulting points are the same than at the initial point on \( \Gamma \).

restrict ourselves to scatterers symmetrical w.r.t. the direction perpendicular to \( \Gamma \). In this case, with \( (n, \tau) \) denoting the local normal and tangent unitary vectors to
Γ, the jump conditions read

\[
\begin{align*}
\begin{cases}
[p] &= B \left\langle \frac{\partial p}{\partial n} \right\rangle, \\
[v_n] &= C_1 \left\langle \frac{\partial v_n}{\partial n} \right\rangle + C_2 \left\langle \frac{\partial v_\tau}{\partial \tau} \right\rangle,
\end{cases}
\end{align*}
\]  

(3)

where \( v_n = \mathbf{V} \cdot \mathbf{n}, v_\tau = \mathbf{V} \cdot \mathbf{\tau}, \) and for any function \( f, \) \( \partial_n f = \nabla f \cdot \mathbf{n}, \) \( \partial_\tau f = \nabla f \cdot \mathbf{\tau}. \) Also, we define

\[
[f] = f_{|\Gamma^+} - f_{|\Gamma^-}, \quad \text{and} \quad \langle f \rangle = \frac{1}{2} \left( f_{|\Gamma^-} + f_{|\Gamma^+} \right)
\]  

(4)

being the jump of \( f \) across the homogenized interface and the mean value of \( f \) respectively (the values \( f_{|\Gamma^-} \) and \( f_{|\Gamma^+} \) are defined locally on \( \Gamma^- \) and \( \Gamma^+ \) along the \( n \) direction). In the above expressions, \( B, C_1 \) and \( C_2 \) are interface parameters which depend on the shape of the scatterers only. As defined in (3), these parameters have the dimension of lengths. Finally, the homogenized problem consists in solving in \( \Omega \setminus \Omega_e \)

\[
\begin{align*}
\rho \frac{\partial \mathbf{V}}{\partial t} &= -\nabla p, \\
\chi \frac{\partial p}{\partial t} &= -\nabla \cdot \mathbf{V}, \\
\text{Jump conditions (3) across } \Omega_e.
\end{align*}
\]  

(5a)  

(5b)  

(5c)

Similar non-trivial jump conditions have been obtained by various authors in related problems. See for instance [25, 8] for a study of acoustics in perforated domains. In the following section, the numerical resolution of the homogenized problem (5) is specifically addressed.

3. Numerical methods

The numerical scheme used to solved (5) is implemented using the first-order hyperbolic system

\[
\frac{\partial}{\partial t} \mathbf{u} + A \frac{\partial}{\partial x} \mathbf{u} + B \frac{\partial}{\partial y} \mathbf{u} = 0,
\]  

(6)

where (6) is deduced from (5) by setting

\[
\mathbf{u} = \begin{pmatrix} v_x \\ v_y \\ p \end{pmatrix}, \quad A = \begin{pmatrix} 0 & 0 & 1/\rho \\ 0 & 0 & 0 \\ 1/\chi & 0 & 0 \end{pmatrix}, \quad B = \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & 1/\rho \\ 0 & 1/\chi & 0 \end{pmatrix}.
\]  

(7)
The ADER-$r$ scheme is used [26] to integrate (6). It is an explicit and two-step finite-difference scheme of order $r$ in both space and time; here we use $r = 4$, which amounts to a fourth-order Lax-Wendroff scheme [17]. It is dispersive of order 4 and dissipative of order 6 [27]. Finally, it is stable under the CFL condition $\beta = c \frac{\Delta t}{\Delta x} \leq 1$ (in two dimensions).

The solution $u$ is discretized in space on a uniform Cartesian grid with mesh sizes $\Delta x = \Delta y$, and in time with a time step $\Delta t$, and we denote $u^n_{i,j}$ the discretized numerical value of $u$ at $M(i,j)$ at $(x = i \Delta x, y = j \Delta y)$ and at time $t = n \Delta t$. The calculation of $u^n_{i,j}$ involves a stencil of $(r+1)^2$ nodes centered at $M$ (25 nodes for ADER 4), which is written formally as the time-marching

$$u^{n+1}_{i,j} = H(u^n_{i_0,j_0}), \quad \{i_0 - i, j_0 - j\} \in \{-2, \cdots, +2\}, \quad (8)$$

with $H$ being deduced from (6) and (7).

The jump conditions (3) of the homogenized problem are discretized by the scheme (8). We introduce now a numerical method that incorporates these conditions in the numerical scheme, for a negligible additional cost.

3.1. Modified ADER scheme in the neighborhoods of $\Omega_e$

To solve the homogenized problem (5), the scheme (8) has to be adapted. Because no constitutive law is defined in the thick interface $\Omega_e$, the solution $u$ in this region is not defined. Thus, we shall first distinguish so-called regular and irregular points $M$ depending on the position of the stencil of $M$ with respect to the interface. In the following, we denote $\Omega^\pm$ the subdomains of $\Omega$ above $\Gamma^+$ and below $\Gamma^-$ (Fig. 3), and $\Omega^+ \cup \Omega^- = \Omega \setminus \Omega_e$.

When all the nodes $P$ of the stencil at $M$ fall in the same physical medium ($\Omega^-$ or $\Omega^+$), the point $M$ is called a regular point. For these points, (8) can be used straightforwardly using $u(M) = u^n_{i,j}$ and $u(P) = u^n_{i_0,j_0}$ being the discretized values of the continuous solution $u$; we call them direct values.

In the neighborhood of $\Omega_e$, it happens that the stencil of $M$ crosses $\Gamma^-$ or $\Gamma^+$ (Fig. 3(a)). It results that nodes $Q$ in the stencil of $M$ belong to $\Omega_e$. Such points $M$ are called irregular since $u(M) = u^n_{i,j}$ in (8) requires the values of $u$ at the nodes $Q$, and $u$ is not defined in $\Omega_e$. In the forthcoming discussion, we focus on an irregular node $Q$ close to $\Gamma^-$: it means that $Q$ is used in the stencils of grid nodes in $\Omega^-$. 

To account for the regular and irregular points, it is possible to adapt (8) in a simple way. This is done by attributing modified values $u^*(Q)$ for the points.
Figure 3: (a) Example of irregular point $M$; in the stencil of $M$ (grey region), $Q$ (●) are in $\Omega_e$ (the usual nodes, in $\Omega^\pm$ are denoted $P$ (×)). (b) Construction of the nodes $P^\pm$ (×) in $\Omega^\pm$ used to calculated the modified value $u^*(Q)$. With $Q^\pm$ the projections of $Q$ on $\Gamma^\pm$ along the normal $n$, the $P^\pm$ are the nodes in the half-disks $D^\pm$ of centers $Q^\pm$ ($P^\pm$ belong to $\Omega^-$ and $\Omega^+$).

$Q$ in $\Omega_e$, and the direct values otherwise. Specifically, the modified ADER 4 is modified as

$$
\begin{align*}
\mathbf{u}_{i,j}^{n+1} &= \mathcal{H} \left( \tilde{\mathbf{u}}_{i_0,j_0}^n \right), \quad \{i_0 - i, j_0 - j\} \in \{-2, \ldots, +2\}, \\
\tilde{\mathbf{u}}_{i_0,j_0}^n &= \mathbf{u}_{i_0,j_0}^n, \quad (x_{i_0}, y_{j_0}) \notin \Omega_e, \\
\tilde{\mathbf{u}}_{i_0,j_0}^n &= \mathbf{u}_{i_0,j_0}^{n*}, \quad (x_{i_0}, y_{j_0}) \in \Omega_e.
\end{align*}
$$

(9)

In (9), the numerical values of the solution are affected to all the nodes in the computational domain (with direct values $\mathbf{u}$ in $\Omega^-$ and $\Omega^+$ and modified values $\mathbf{u}^*$ in $\Omega_e$). We shall see in the following section that the modified values (at the points $Q$) are expressed in terms of the direct values at points $P^\pm$ in $\Omega^\pm$ (Fig. 3(b), the choice of the $P^\pm$ is incidental at this stage). It follows that (9) is solved implicitly on the direct values only: the solution in $\Omega_e$ is not questioned, as expected ($Q$ does not appear as the center of a stencil in (9)).

3.2. Construction of the modified values $\mathbf{u}^*$ at the nodes $Q$ in $\Omega_e$

As previously said, the solution in $\Omega_e$ is undefined ($\mathbf{u}^*(Q)$ does not exist). Only the jump conditions across $\Omega_e$ make sense. Thus, we shall start by defining
and this will be done using $Q^\pm$ being the projections of $Q$ on $\Gamma^\pm$ along $n$, and using the jump conditions (3) which apply between $Q^+$ and $Q^-$. Next, because $Q^\pm$ are not nodes (except by casuality), the solutions at $Q^\pm$ will be expressed in terms of two sets of direct values $u(P^\pm)$. This is the meaning of the nodes $P^\pm$, chosen in the vicinity of $Q^\pm$.

The construction of $u^*(Q)$ presented in the forthcoming section applies for any sets of $P^\pm$ being nodes of $\Omega^\pm$ in the vicinities of $Q^\pm$. Fig. 3(b) illustrates our choice: $P^\pm$ are the nodes of $\Omega^\pm$ in the disks $D^\pm$ centered at $Q^\pm$ and of radius $d$. In practice, $d \simeq 3.5\Delta x$ and this is discussed further in Sec. 3.3.

### 3.2.1. The ingredients of the construction of $u^*(Q)$

In this section, the construction of $u^*(Q)$ is presented formally. It will be detailed precisely in the Sections 3.2.2 to 3.2.5. Let us recall that $Q$ is used for time-marching at points of $\Omega^-$, and thus is close to $\Gamma^-$. Because the solution is not defined in $\Omega_e$, we start by defining $u^*(Q)$ as the smooth extension of the solution in $\Omega^-$. This is done using a Taylor expansion for $Q$ in the neighborhood of $Q^-$, written formally

$$u^*(Q) = T(Q, Q^-) w(Q^-).$$

In one dimension at order 1, $u^*(Q) = u(Q^-) + (x_Q - x_{Q^-}) \partial_x u(Q^-); w(Q^-)$ encapsulates the weights $(u(Q^-), \partial_x u(Q^-))$ while $T(Q, Q^-)$ are the polynomial forms depending on both $Q$ and $Q^-$, here 1 and $(x_Q - x_{Q^-})$. It is worth noting that (10) introduces a disymmetry between $\Omega^-$ and $\Omega^+$. Indeed, if $u^*(Q)$ is a smooth extension of the solution in $\Omega^-$, it cannot be a smooth extension of the solution in $\Omega^+$ because of the jump conditions. These jump conditions apply between $Q^-$ and $Q^+$ but $Q^\pm$ do not coincide with nodes. Thus, we use the sets of nodes $P^\pm$ in the vicinities of $Q^\pm$, and the Taylor expansions

$$u(P^\pm) = T(P^\pm, Q^\pm) w(Q^\pm),$$

which involves the direct values $u(P^\pm)$.

From (10-11), it is visible that $u^*(Q)$ can be expressed as a function of $u(P^\pm)$ if a relation between $w(Q^\pm)$ and $w(Q^-)$ is established. This relation will obviously involve the jump conditions. Before doing so, the number of terms in $w$ is reduced owing to high-order compatibility conditions, deduced from the initial condition $\nabla \wedge v = 0$ (coming from (5a) and valid in $\Omega^\pm$). The new vector $v$
collects the reduced numbers of unknowns of $w$, with

$$w(Q^\pm) = Gv(Q^\pm), \quad w(P^\pm) = Gv(P^\pm),$$

(12)

and $G$ is a constant matrix. Finally, the jump conditions are used to get a relation between $v(Q^+)$ and $v(Q^-)$, written

$$v(Q^+) = \text{Func} \left[ v(Q^-) \right].$$

(13)

The relations (10) to (13) allow to conclude. First, $u(P^\pm)$ can be expressed as a function of $v(Q^-)$ only. From (11), we have $u(P^-) = T(P^-, Q^-)Gv(Q^-)$. For $u(P^+)$, it starts the same, with $u(P^+) = T(P^+, Q^+)Gv(Q^+)$ and (13) allows to conclude (and (13) is essential since it encapsulates the jump conditions). Using these relations for all the nodes $P^\pm \in D^\pm$ and collecting the direct values $u(P^\pm)$ in a single vector $U = (u(P^-)|_{P^- \in D^-}, u(P^+)|_{P^+ \in D^+})$, the formal relation $U = Mv(Q^-)$ can be inverted to get

$$v(Q^-) = M^{-1} \begin{pmatrix} u(P^-) \\ u(P^+) \end{pmatrix}.$$  

(14)

Finally, (14) is injected in (10), using (12), to get the modified values $u^*(Q)$ as a function of the direct values $u(P^\pm)$

$$u^*(Q) = T(Q, Q^-) G M^{-1} \begin{pmatrix} u(P^-) \\ u(P^+) \end{pmatrix}.$$  

(15)

We shall now detail the steps in the construction of $u^*(Q)$ as implemented numerically for $k$-th order Taylor expansions.

3.2.2. The Taylor expansions, Eqs. (10)-(11)

Here, we simply specify the notations in (10) and (11) to get Taylor expansions at the order $k$. The matrix $T_k$ of $k$-th order expansions for $Q$ near $Q^-$ reads

$$T_k(Q, Q^-) = \begin{pmatrix} I_3, \cdots, \frac{1}{\ell!} (x_Q - x_Q^-)^{\ell-m} (y_Q - y_Q^-)^m I_3, \cdots, \frac{(y_Q - y_Q^-)^k}{k!} I_3 \end{pmatrix},$$

(16)

with $\ell = 0, \cdots, k$ and $m = 0, \cdots, \ell$ and $I_3$ the $3 \times 3$ identity matrix. With $(\ell + 1)$ polynomial forms at the order $\ell$, $T_k$ is a $3 \times n_w$ matrix, with $n_w = 3(k+1)(k+2)/2$. 

10
Next, we collect in a single vector $w_k$ the $n_w$ limit values of $u$ and of its successive spatial derivatives up to the $k$-th order, at $Q^\pm$:

$$w_k(Q^\pm) = \lim_{Q^\pm \in \Omega^\pm} \left( u^T, ..., \frac{\partial^\ell}{\partial x^\ell_m \partial y^m} u^T, ..., \frac{\partial^k}{\partial y^k} u^T \right)^T,$$

(17)

with $\ell = 0, \ldots, k$ and $m = 0, \ldots, \ell$. The modified value $u^*(Q)$ is defined as a smooth extension of the solution in $\Omega^-$

$$u^*(Q) = T_k(Q, Q^-) w_k(Q^-),$$

(18)

and $u^*(Q)$ appears to depend on $k$. As previously said, once this definition is chosen, $u^*(Q)$ cannot be a smooth extension of the solution in $\Omega^+$ because of the jump conditions.

The story is different for the Taylor extensions of the direct values $u(P^\pm)$ written crudely in (11). Because $u(P^\pm)$ are the discretized versions of the exact solution, their Taylor expansions have to be written as approximations, namely

$$u(P^\pm) = T_k(P^\pm, Q^\pm) w_k(Q^\pm) + \mathcal{O}(\Delta x^{k+1}).$$

(19)

3.2.3. High-order compatibility condition, Eq. (12)

The equation (5a) provides a compatibility condition, $\nabla \wedge v = 0$, which tells us that the fluid is irrotational in $\Omega^\pm$. Assuming sufficiently smooth solutions in $\Omega^\pm$, this relation can be differentiated $(\ell - 1)$ times w.r.t. to $x$ and $y$:

$$\frac{\partial^\ell v_y}{\partial x^\ell_m \partial y^{m+1}} = \frac{\partial^\ell v_x}{\partial x^\ell_m \partial y^m}, \quad \ell \geq 1, \ m = 0, \ldots, \ell - 1,$$

(20)

whose version with $\ell = 1, m = 0$ is the originate condition. Doing so for $\ell = 1, \ldots, k$ provides $k(k + 1)/2$ high-order compatibility conditions. The equations are valid everywhere outside $\Omega_e$, and in particular at $Q^\pm$. This allows to use vectors $v_k$ containing only the remaining independent derivatives, thus being reduced in size (to $n_v = (k + 1)(k + 3)$ components). The $v_k$ are linked to $w_k$ by

$$w_k(Q^\pm) = G_k v_k(Q^\pm),$$

(21)

where $G_k$ is a $n_w \times n_v$ matrix (an algorithm to compute $G_k$ can be found in [14], see the appendix A in this reference). Now, (19) can be written as a function of $v_k(Q^\pm)$, using (21),

$$u(P^\pm) = T_k(P^\pm, Q^\pm) G_k v_k(Q^\pm) + \mathcal{O}(\Delta x^{k+1}).$$

(22)
3.2.4. High-order order jump conditions, (13)

To express the jump conditions between $Q^{-}$ and $Q^{+}$, it is useful to come back to the construction of $\Gamma^{\pm}$ and of $Q^{\pm}$ (Fig. 4). Because $\Gamma^{\pm}$ are deduced from $\Gamma$ by the translations of $\pm e/2$ along $n$, the vectors locally normal and tangent to $\Gamma^{\pm}$ at $Q^{\pm}$ are the same, and the same as the vectors $(n, \tau)$, locally normal and tangent to $\Gamma$ at $Q_{\Gamma}$. In practice, we use a parametrization of $\Gamma$ with $s$, with $[X(s), Y(s)]$ the coordinates of $Q_{\Gamma}$. We denote $n = (Y'(s), -X'(s))$ and $\tau = (X'(s), Y'(s))$, with prime denoting the derivative.

![Figure 4: Parametrization of $\Gamma$ with the parameter $s$; the vectors $(n, \tau)$ locally normal and tangent to $\Gamma$ at $Q_{\Gamma}[X(s), Y(s)]$ are also normal and tangent to $\Gamma^{\pm}$ at $Q^{\pm}$ by construction.](image)

From (3) and (4), the fields at $Q^{\pm}$ are related by

\[
\begin{align*}
\begin{cases}
p(Q^{+}) - p(Q^{-}) = \frac{B}{2} \left( \frac{\partial p}{\partial n}(Q^{+}) + \frac{\partial p}{\partial n}(Q^{-}) \right), \\
v_{n}(Q^{+}) - v_{n}(Q^{-}) = \frac{C_{1}}{2} \left( \frac{\partial v_{n}}{\partial n}(Q^{+}) + \frac{\partial v_{n}}{\partial n}(Q^{-}) \right) + \frac{C_{2}}{2} \left( \frac{\partial v_{\tau}}{\partial \tau}(Q^{+}) + \frac{\partial v_{\tau}}{\partial \tau}(Q^{-}) \right).
\end{cases}
\end{align*}
\]

(23)

The normal and tangential derivatives involved in (23) are expressed in terms of
the parametric representation of $\Gamma$: from the components of $(n, \tau)$:

$$v_n = Y' v_1 - X' v_2,$$
$$v_\tau = X' v_1 + Y' v_2,$$

$$\frac{\partial v_n}{\partial n} = (Y')^2 \frac{\partial v_1}{\partial x} - X' Y' \left( \frac{\partial v_2}{\partial x} + \frac{\partial v_1}{\partial y} \right) + (X')^2 \frac{\partial v_2}{\partial y}, \quad (24)$$
$$\frac{\partial v_\tau}{\partial \tau} = (X')^2 \frac{\partial v_1}{\partial x} + X' Y' \left( \frac{\partial v_2}{\partial x} + \frac{\partial v_1}{\partial y} \right) + (Y')^2 \frac{\partial v_2}{\partial y},$$
$$\frac{\partial p}{\partial n} = Y' \frac{\partial p}{\partial x} - X' \frac{\partial p}{\partial y}. \quad (24)$$

Injecting (24) into (23) leads to lengthy expressions, which are summed up by the matrix form

$$E_0 \left[ w_0(Q^+) - w_0(Q^-) \right] = \frac{1}{2} F_1 \left[ w_1(Q^+) + w_1(Q^-) \right], \quad (25)$$

where the matrices $E_0$ and $F_1$ depend on the geometry of the interface because of their dependance on $(n, \tau)$. $E_0$ is a $2 \times 3$ matrix defined by

$$E_0 = \begin{pmatrix} Y' & -X' & 0 \\ 0 & 0 & 1 \end{pmatrix}, \quad (26)$$

and $F_1$ is a $2 \times 9$ matrix whose non-zero components are

$$F_1(1, 4) = C_1 \left( Y' \right)^2 + C_2 \left( X' \right)^2,$$
$$F_1(1, 5) = F_1(1, 7) = -C_1 X' Y' + C_2 X' Y',$$
$$F_1(1, 8) = C_1 \left( X' \right)^2 + C_2 \left( Y' \right)^2,$$
$$F_1(2, 6) = BY',$$
$$F_1(2, 9) = -BX'. \quad (27)$$

Now, the goal is to obtain an extended version of the jump conditions involving derivatives of $u$ up the $k$-th order, namely involving $w_k$. For this purpose, (25) is differentiated w.r.t the time $t$ and the parameter $s$, and using

$$\begin{cases} \frac{\partial}{\partial t} w_\ell = -A \frac{\partial}{\partial x} w_\ell - B \frac{\partial}{\partial y} w_\ell, \\ \frac{\partial}{\partial s} = X'(s) \frac{\partial}{\partial x} + Y'(s) \frac{\partial}{\partial y}, \end{cases} \quad (28)$$
where the first relation holds for any $\ell$. Applying to (25) the chain-rule
\[
\frac{\partial^{(\ell-i)}}{\partial t^{(\ell-i)}} \frac{\partial}{\partial s^i} \left[ E_0 \left( w_0(Q^+) - w_0(Q^-) \right) - \frac{1}{2} F_1 \left( w_1(Q^+) + w_1(Q^-) \right) \right] = 0,
\]
where $i = 0, \ldots, \ell$, provides ($\ell+1$) matrix relations, or $2(\ell+1)$ scalar relations, at each order $\ell$. These relations involve spatial derivatives up to the order $(\ell+1)$ and the chain-rule is stopped at the order $k$. The $n_j = (k+1)(k+2)$ obtained relations for $\ell = 0$ to $k$ are collected in the matrix relation
\[
E_k \left[ w_k(Q^+) - w_k(Q^-) \right] = \frac{1}{2} F_{k+1} \left[ w_{k+1}(Q^+) + w_{k+1}(Q^-) \right].
\] (30)
For instance, at $k = 1$ one gets
\[
E_1 = \begin{pmatrix}
E_0 & 0 & E_0 \\
0 & -E_0A & -E_0B \\
E_0' & X'E_0 & Y'E_0
\end{pmatrix}.
\] (31)
Computing the matrices $E_k$ and $F_{k+1}$ for $k > 1$ is a tedious task, that has been automatized thanks to computer algebra tools. It is worth noting that they depend on the first $k$-th derivatives of $X(s)$ and $Y(s)$, thus they encapsulate informations on the interface shape well beyond its position only.

In (30), $w_{k+1}$ involves spatial derivatives up to the order $(k+1)$, and this order is not considered in the resolution at the order $k$. Considering $F_{k+1}$ the restriction of $F_{k+1}$ obtained by removing the last $(n_w + 1)$-th column in the matrix $F_{k+1}$, (30) simplifies to
\[
E_k^+ w_k(Q^+) = E_k^- w_k(Q^-),
\] where $E_k^\pm \equiv E_k \mp \frac{1}{2} F_{k+1}$ are two $n_j \times n_w$ matrices. Combining (21) and (32) yields
\[
S_k^+ v_k(Q^+) = S_k^- v_k(Q^-),
\] (33)
with the $n_j \times n_v$ matrices $S_k^+ = E_k^+ G_k$ being smaller than the $n_j \times n_w E_k^\pm$. With $n_j = (k+1)(k+2)$ and $n_v = (k+1)(k+3)$, the system (33) is underdetermined. It is inverted in the sense of the least-squares sense using Singular Value Decomposition, leading to
\[
v_k(Q^+) = \left( \left( S_k^+ \right)^{-1} S_k^- \ | \ K S_k^+ \right) \left( \begin{array}{c} v_k(Q^-) \\ \Lambda_k \end{array} \right),
\] (34)
where \((S_k^+)^{-1}\) is the least-square pseudo-inverse of \(S_k^+\), \(K_{S_k^+}\) is the matrix filled with the kernel of \(S_k^+\), and \(\Lambda_k\) is a set of the \((n_v-n_i)\) Lagrange multipliers which are the coordinates of \(v_k(Q^+)^{-1}\) onto the kernel. A singular value decomposition of \(S_k^+\) is used to build \((S_k^+)^{-1}\) and the kernel \(K_{S_k^+}\) [24].

3.2.5. Final step in the construction of modified values, (14)-(15)

We want to express \(v_k(Q^-)\) as a function of the set of direct values \(u(P^\pm)\) (we denote \(n^\pm\) the numbers of nodes \(P^\pm\)). For the set of nodes \(P^-\) in \(\Omega^-\), (19) is simply re-written

\[
\begin{align*}
u(P^-) &= T_k(P^-, Q^-)G_k \left( 1 | 0 \right) \begin{pmatrix} v_k(Q^-) \\ \Lambda_k \end{pmatrix} + \mathcal{O}(\Delta x^{k+1}). \tag{35}
\end{align*}
\]

In the above relation, 1 stands for the \(n_v \times n_v\) identity matrix and 0 for the \(n_v \times (n_v-n_i)\) zero matrix. For the set of nodes \(P^+\) in \(\Omega^+\), we use (34) in (22), whence

\[
\begin{align*}
u(P^+) &= T_k(P^+, Q^+)G_k \left[ (S_k^+)^{-1} S_k^- | K_{S_k^+} \right] \begin{pmatrix} v_k(Q^-) \\ \Lambda_k \end{pmatrix} + \mathcal{O}(\Delta x^{k+1}). \tag{36}
\end{align*}
\]

Now, we collect in a single vector \(U\) the \((n^- + n^+)\) vectors \(u(P^\pm)\)

\[
U = \begin{pmatrix} u(P_1^-) \\ \vdots \\ u(P_n^-) \\ u(P_1^+) \\ \vdots \\ u(P_n^+) \end{pmatrix}, \tag{37}
\]

and use (35) and (36) written in the matrix form

\[
U = M \begin{pmatrix} v_k(Q^-) \\ \Lambda_k \end{pmatrix} + \mathcal{O}(\Delta x^{k+1}), \tag{38}
\]

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where

\[
M \equiv \begin{pmatrix}
T_k(P_1^-, Q^-)G_k (1 | 0) \\
\vdots \\
T_k(P_n^-, Q^-)G_k (1 | 0) \\
T_k(P_1^+, Q^+)G_k \left( \left( S_k^+ \right)^{-1} S_k^- | K_{S_k^+} \right) \\
\vdots \\
T_k(P_n^+, Q^+)G_k \left( \left( S_k^+ \right)^{-1} S_k^- | K_{S_k^+} \right)
\end{pmatrix}
\]  

(39)

is a \((3(n^- + n^+) \times (2n_v - n_s))\) matrix. To ensure that the system \((38)\) is overdetemined, the radius \(R_D\) of the discs \(D^\pm\) has to be chosen in order to ensure that

\[
\gamma(R_D, k) \equiv \frac{3(n^- + n^+)}{(k + 1)(k + 4)} \geq 1,
\]  

(40)

and this condition will be further discussed in Section 3.3. The condition \((40)\) being ensured, \((38)\) is solved using the least-squares inverse \(M^{-1}\) of \(M\). The Lagrange multipliers \(\Lambda_k\) have been incorporated in the construction of \(M\), but they are not needed to build the modified value. Thus, they are removed using the \((n_w - n_v) \times 3(n^- + n^+)\) restriction \(\overline{M}^{-1}\) of \(M^{-1}\), to get only

\[
v_r(Q^-) = \overline{M}^{-1}U,
\]

and finally using \((18)\)

\[
u^*(Q) = T_k(Q, Q^-) G_k \overline{M}^{-1} U.
\]  

(41)

The matrices \(T_k(P^\pm, Q^\pm)\), \(G_k\) and \(\left( \left( S_k^+ \right)^{-1} S_k^- | K_{S_k^+} \right)\) are involved in \((41)\) through \(\overline{M}^{-1}\).

3.3. Comments and practical details

1. The described algorithm is applied to the irregular points in \(\Omega^\pm\). The sizes of the matrices involved are summarized in table 1. Since the jump conditions do not vary with time, the evaluation of the matrices in \((41)\) is done during a preprocessing step. Only small matrix-vector products are therefore required at each time step. After optimization of the computer codes, this additional cost is made negligible, lower than 1% of the time-marching of the ADER 4 scheme.
\[ W = \frac{3 (k + 1)(k + 2)}{2} \]
\[ V = (k + 1)(k + 3) < W \]
\[ J = (k + 1)(k + 2) < V \]

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Size</th>
</tr>
</thead>
<tbody>
<tr>
<td>( T_k )</td>
<td>( 3 \times n_w )</td>
</tr>
<tr>
<td>( G_k )</td>
<td>( n_w \times n_v )</td>
</tr>
<tr>
<td>( S^\pm_k )</td>
<td>( n_j \times n_v )</td>
</tr>
<tr>
<td>( M )</td>
<td>( 3 (n^+ + n^-) \times (2n_w - 2n_v - n_j) )</td>
</tr>
<tr>
<td>( M^{-1} )</td>
<td>( 3 (n^+ + n^-) \times (n_w - n_v) )</td>
</tr>
</tbody>
</table>

Table 1: Quantities involved in the computation of the modified values (section 3.2).

2. The matrix \( M \) in (38) depends on the subcell positions of \( Q^\pm \) inside the mesh and on the jump conditions expressed between \( Q^+ \) and \( Q^- \), involving the local geometry and the curvature of \( \Gamma^\pm \) at \( Q^\pm \). Consequently, all these insights are incorporated in the modified value (41), and thus also in the scheme.

3. The optimal order \( k \) depends on the order \( r \) of the scheme and on the jump conditions. Let us begin with the classical case of acoustics where the jump conditions do not involve spatial derivatives: for instance, \([v_n] = 0\) and \([p] = 0\). In this case, taking \( k = r \) maintains a \( r \)-th order global accuracy [4] (the criterion \( k = r - 1 \) is even sufficient [9]). In the non-classical case studied here, the jump conditions involve first-order spatial derivatives. After successive derivations, the higher-order terms are canceled in (32), which introduces a loss of accuracy. To maintain the \( r \)-th order convergence, the order of the ESIM must be increased of one unity: \( k = r + 1 \equiv 5 \) for the ADER 4 scheme.

4. The simulations indicate that overestimation of \( \gamma \) in (40) ensures the stability of the ESIM. Numerical experiments have shown that \( d = 3.5 \Delta x \) is a good choice. Typically, it gives \( n^\pm \approx 10 \) and \( \gamma \approx 4 \).

3.4. Summary of the algorithm

The numerical strategy presented all along this section couples two numerical methods: a finite difference ADER scheme and the Explicit Simplified Interface Method. To clarify the interactions between these methods, the algorithm can be summarized as follows:

- **Preprocessing**
• Detection of irregular grid points

• At each irregular point $M$:
  - detection of the orthogonal projections $Q^\pm$
  - computation of the matrices $S_k^\pm$ (33)
  - filling of $M$ (39)
  - computation of extrapolation matrix in (41)

• Initialization of the computation at $t = 0$

▷ Time iterations

• Computation of the modified values (41)

• Time-marching at the regular points (8)

• Time-marching at the irregular points (9)

4. Numerical experiments

In this section, we first validate the ESIM implemented on the thick interface as presented in the preceding section. This is done for a plane wave at normal and oblique incidences on the interface. These cases allow for analytical solutions of the homogenized problem, already validated in [18, 21] in the harmonic regime. The extension to the time domain being done by means of discrete inverse Fourier transforms, we do not repeat the validation by comparison of the solutions of the homogenized and of the direct problems. We inspect the accuracy of the numerical scheme varying the order $k$ in the scheme, and discuss the convergence obtained with the estimated optimal value $k = 5$.

Next, we exemplify the capability of the homogenized problem (5) to mimic the real problem (1) in the time domain. The solutions of the direct and the homogenized problems are compared in the case of a source emitting a short pulse with a central frequency that we vary to inspect the robustness of the homogenized solution w.r.t. the small parameter $\epsilon$, defined in (2) (and $\lambda_0$ will be defined latter). This is done for a straight and curved line array.

The following characteristics hold for all our simulations:
• The real problem is an array of 50 rectangular sound-hard scatterers in water. The periodicity of the array is \( h = 20 \text{ m} \) with a filling ratio of the scatterers \( \varphi = 0.5 \); the thickness of the array is \( e = 20 \text{ m} \). For these dimensions of the scatterers, the interface coefficients entering in the jump conditions of the homogenized problem are

\[
B = 44.412, \quad C_1 = 10, \quad C_2 = 8.338. \tag{42}
\]

(see Appendix A). For water, we use \( \rho = 1000 \text{ kg/m}^3 \) and \( c = 1500 \text{ m/s} \).

• The numerical results have been performed in a domain of extension 1200 m \( \times \) 1200 m, discretized on \( N_x \times N_y \) points, with \( N_x = N_y \). The time step follows from the CFL condition: \( \Delta t = \beta \Delta x / c \), and we used \( \beta = 0.95 \).

• Discrepancies between two solutions yielding the pressure fields \( p_1 \) and \( p_2 \) are given by the relative difference \( \| p_1 - p_2 \| / \| p_1 \| \) (and \( \| . \| \) refers to the discrete L^2- norm).

• The temporal signals are built using \( h(t) \) being a combination of truncated sinusoids

\[
h(t) = \begin{cases} 
\sum_{m=1}^{4} a_m \sin(\beta_m \omega_0 t) & \text{if } 0 < t < \frac{1}{f_0}, \\
0 & \text{otherwise},
\end{cases}
\tag{43}
\]

where \( \beta_m = 2^{m-1} \), \( \omega_0 = 2\pi f_0 \); the coefficients \( a_m \) are: \( a_1 = 1, a_2 = -21/32, a_3 = 63/768, a_4 = -1/512 \), ensuring \( C^6 \) smoothness of \( h(t) \). The Fourier transform of (43) reads

\[
\hat{h}(\omega) = \frac{\omega_0}{2\pi} \sum_{m=1}^{4} a_m \beta_m e^{2\pi i \omega_0 t} - \frac{1}{\omega^2 - \beta_m^2 \omega_0^2},
\tag{44}
\]

with a maximum slightly greater than \( f_0 \) and a cut-off frequency \( f_m \) at around \( 5f_0 \). \( h(t) \) and \( \hat{h}(\omega) \) are shown in Figs. 5 for \( f_0 = 10 \text{ Hz} \).

In the simulations, various central frequencies are considered: \( f_0 = 2.5, 5 \) and 10 Hz. The resulting values of \( \varepsilon \) calculated using \( k_0 = 2\pi f_0 / c \) are \( \varepsilon = 0.21, 0.41 \) and 0.83. However the spectral content of the source being large, the wave packet contains wavenumbers 5 times smaller than \( k_0 \); these small scales are associated to \( \varepsilon \) larger than 1 (up to 4).
4.1. Validation of the ESIM for large interface

To begin with, we examine how accurately the ESIM discretizes the homogenized jump conditions. To that aim, we consider a plane wave at oblique incidence \( \theta_I \) on the interface. This case allows for an exact, or say reference, solution of the homogenized problem in the frequency domain (see Appendix B)

\[
u(x, y, \omega) = u_I(x, y, \omega) + R(\omega)u_R(x, y, \omega) + T(\omega)u_T(x, y, \omega),
\]

with \( u_{I,R,T} \) given by (B.2) and \( (R, T) \) by (B.5). Afterwards the solution \( u(x, y, t) \) in the time domain (called reference homogenized solution in the following) is deduced by discrete inverse Fourier transform of \( u(x, y, \omega) \).

Numerically, this solution has to be recovered by imposing, at each time step, the reference solution \( u(i\Delta x, j\Delta x, n\Delta t) \) on the 2 lines \( (i = 0, 1 \text{ and } (N_x - 1), N_x) \) and the 2 columns \( (j = 0, 1 \text{ and } (N_y - 1), N_y) \) at the edges of the computational domain. Once these boundary conditions have been imposed, the numerical scheme has to be able to produce the solution in the whole domain. This numerical solution is referred as the numerical homogenized solution in the following.

In the simulations, we used a temporal signal with Fourier dependence given by (44) at the central frequency \( f_0 = 10 \) Hz, resulting in a wave packet with the central and smallest wavelengths of 150 m and 30 m respectively. The homogenized interface mimicking the array of Neumann rectangles (with spacing 20 m, filling fraction 0.5 and thickness 20 m) is thus 20 m large, and associated to the interface parameters (42). The Fourier synthesis is done using \( N_f = 256 \) modes with a uniform frequency step \( \Delta f = 0.4 \) Hz around \( f_0 \) and yields the reference signal.

Figure 5: (a) Temporal signal at a source point (43) for \( f_0 = 10 \) Hz, and (b) corresponding spectrum.
solution $u$ imposed at the 2 points boundaries of the computational domain, as previously described. The computational domain is 1200 m × 1200 m large and it is discretized using $\Delta x = \Delta y = 2$ m ($N_x = N_y = 600$), thus from the CFL condition $\Delta t = 0.0013$ s.

4.1.1. Plane wave at normal incidence

We first consider a normal incidence, for which $u$ is independent of $x$. Fig. 6 shows the pressure field of the reference solution which is imposed in the whole domain at the initial time $t = 0$; this initial time has been chosen before the wave hits the interface (the profile of this one-dimensional field is also reported).

![Figure 6](image1.png)

Figure 6: (a) Pressure field of the incident wave (for the reference solution) imposed at the initial time of the simulation and (b) corresponding $y$-profile.

The reference and the numerical homogenized solutions are then computed in time; the pressure fields of the numerical solution after 158 iterations ($t = 0.2$ s) is reported in Fig. 7(a). Fig. 7(b) reports the profiles along $y$ of the two solutions. The discrepancy between both solutions is 0.2%, a very low error which can be attributable to the discretization. We conclude that the ESIM discretized correctly the jump conditions.

To further inspect the accuracy of the method, we report results varying the $k$-values and the spatial discretization. With the same $\Delta x = 2$ m, we used the orders $k = 1$ to 5 to compute the numerical homogenized solution. Fig. 8(a) reports the resulting pressure profiles along $y$ at $t = 0.2$ s (the profile for $k = 5$ is indiscernible of the one for $k = 4$ and it is not reported). Fig. 8(b) reports the errors between these profiles and their reference counterpart as a function of $k$. 

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While the solution computed with \( k = 1 \) misses the correct order of magnitude of the reference homogenized solution, good results are obtained for \( k > 1 \), and the error becomes incidental for \( k = 5 \) as used later on.

Next, we used \( k = 5 \) and various discretizations for \( N_x = 300 \) up to \( N_x = 4800 \). The error is reported in Fig. 9 as a function of \( N_x \). The order measured (being the slope of the curve) is close to 4 at this order as soon as \( N_x = 300 \) (\( \Delta x = 4 \) m) and this corresponds to the best convergence that we can expect since the order 4 is the order of ADER 4 in homogeneous medium. It is worth noting that this order 4 is not found for \( k < 5 \), and such high order has a numerical cost. However, the difference in accuracy between \( k = 4 \) and \( k = 5 \) being very small, a good compromise in practice is to choose \( k = 4 \). In terms of the spatial resolution, we have said that the smallest wavelength is about 30 m; with an error less than 1% for \( N_x = 600 \) (\( \Delta x = 2 \) m), we can estimate that 15 grid nodes per wavelength calculated for the smallest wavelength is a good criterion to fix the spatial resolution (and this conclusion holds for \( k = 4 \) as well).

4.1.2. Plane wave at oblique incidence and tilted interface

The same comparison between the reference and computed homogenized solutions is performed in the case of a wave at oblique incidence and, more importantly, using a tilted interface. As in the previous section, the reference homogenized solution is obtained by discrete inverse Fourier transform of (45) with \( \theta_I \) the
Figure 8: Influence of the order \( k \) in the computation of the numerical homogenized solution. (a) Zoom on the pressure profile of Fig. 7(b); the profiles for \( k = 1, 2, 3 \) and 4 are shown (the profiles for \( k = 4 \) and 5 are indiscernable). (b) Errors of the numerical solution compared to the reference solution (calculated for \( y \in [0, 1200] \)) as a function of the order \( k \).

Figure 9: Error as a function of the mesh size \( N_x \) (for \( k = 5 \)). The numerical values indicate the slopes between two successive points, the slope -4 is shown in dotted line.

angle between the incident wave and the tilted interface (see Appendix B). The case of a tilted interface allows us to inspect another aspect of the efficiency of the ESIM, namely its capacity to account for the interface shape with a subcell resolution on a Cartesian grid. In the present case, this means that the real slope of the interface is accounted for, instead of a crude stair-step discretization.
We considered the interface with a tilt angle of about $10^\circ$ with $Ox$ and the incident wave packet makes an angle $-30^\circ$ with $Ox$. Figs. 10 show the pressure fields of the numerical homogenized solution at the initial time (identical to the reference one) and after the wave packet has propagated ($t = 0.2$ s corresponding to 158 iterations). As expected, no spurious diffractions have appeared.

Figure 10: (a) Pressure field of the numerical homogenized solution (identical to the reference one) imposed at the initial time of the simulation, (b) Pressure field of the numerical homogenized solution at $t = 0.2$ s; the vertical slice (dotted lines) at $x = 740$ m, $y \in [100, 800]$ m is used in Fig. 11.

More quantitatively, the pressure profile along the vertical slice ($x = 740$ m and $y \in [100, 800]$ m) is reported in Fig. 11 together with the corresponding reference solution (the scattered wave packets (R and T) are visible on these profiles). The discrepancy between the two profiles is of 0.5%, as small as in the case of a non tilted interface. This accuracy could not be obtained with a piecewise constant approximation of the interface shape, and it confirms that the slope of interface is accurately accounted for in the numerical scheme.

4.2. Validation of the homogenized problem in the time domain

In the previous section, we have inspected the ability of the ESIM to properly account for the homogenized jump conditions (3). Another question is whether or not the homogenized problem (5) is a good approximation of the real one (1); this question is addressed now through two examples. First, we consider an array of scatterers along a straight line $\Gamma$, which corresponds to the configuration for which the jump conditions have been derived [20]. Next, we consider an array located
onto a curved line $\Gamma$, for which we extended heuristically the jump conditions. This allows us to inspect the intuitive idea that such extension is possible for small curvature of $\Gamma$.

The real problem is solved numerically following the method presented in [16]. The numerical method uses a ADER 4 scheme, and is able to accurately account for free boundaries at an interface by affecting so-called fictitious values of the solution inside the sound-hard scatterers. In the spirit, these fictitious values are the equivalent of the modified values presented in this paper, and used in the homogenized thick interface.

4.2.1. The case of a straight array of sound-hard scatterers

The array of rectangular scatterers is placed at $y = 500$ m along the $x$-axis. Each rectangle is 10 m large along $x$, with spacing $h = 20$ m, and $e = 20$ m thick along $y$. In the homogenized problem, the interface is $e = 20$ m thick (the region $y = [490, 510]$ m is not resolved), and associated to the interface parameters (42). We considered a source point at $(x_s = 600, y_s = 620)$ m emitting the short pulse $h(t)$ in (43), with central frequency $f_0$. When not specified, we used a fine grid $\Delta x = 0.125$ m for the real problem and a coarse grid $\Delta x = 2$ m for the homogenized problem (in both cases, $k = 4$ has been considered). These are the mesh sizes required to get converged solutions (from $\Delta x = 2$ m, reducing the mesh size to $\Delta x = 0.5$ m produces about 20% variations of the solution of the real problem,
Figure 12: Pressure fields $p^f$ (real problem with the array) and $p$ (homogenized problem) computed with ADER 4 at $t = 0.38$ s for $f_0 = 2.5$, 5 and 10 Hz. $f_0$ is the central frequency of the signal $h(t)$ in (43) imposed by the source point (indicated by the black point).

while the solutions of the homogenized have already converged, with variations less than 0.1%).
The pressure fields $p^r$ computed in the real problem and $p$ in the homogenized one are reported in Figs. 12 for central frequencies $f_0 = 2.5$, 5 and 10 Hz, at $t = 0.38$ s (corresponding profiles along the centerline $x = 600$ m are reported in Figs. 13). The discrepancy between the two fields is of about 5% for $f_0 = 2.5$ and 5 Hz, and it is of 10% for $f_0 = 10$ Hz, and these orders of magnitude are in agreement with those reported in the frequency regime, see [18] (we measured the discrepancy between both fields outside the thick interface $y \in [490, 510]$ m where $p$ is not defined). Note that, at $f_0 = 10$ Hz, the central and smallest wavelengths are 150 and 30 m, leading to $\varepsilon \sim 0.8 - 4$, so overcoming the intuitive limit $\varepsilon = 1$ for the validity of the homogenization. We report in Fig. 13(d) the relative errors for increasing $\varepsilon$-value. Here, the error has been calculated on the part of the profiles corresponding to the transmitted wave packet $y \in [0, 490]$ m (thus avoiding to cross the interface) and with $\varepsilon = 2\pi f_0 h/c$ (an extra point at $f_0 = 20$ Hz has been added). The observed $\varepsilon^2$ scaling is consistent with the expected second-order accuracy of the interface homogenization model.

The computations have been performed using a fine grid $\Delta x = 0.125$ m for the real problem and a coarse grid $\Delta x = 2$ m for the homogenized problem; as previously said, we checked that these meshes are required to get converged solutions. This is not obvious at $f_0 = 10$ Hz where the source generates wavelengths of the same order of magnitude than the size of the array; thus, we could expect that the two problems, real and homogenized, require the same mesh size, but we observe that it is not the case. Heuristically, this can be explained as follows. The fine grid needed for the real problem resolves the smallest scale, and it turns out that this smallest scale is associated to the evanescent field, excited in the vicinity of the array (this small scale is visible in Fig. 12(a) for $f_0 = 10$ Hz). This means that the usual rule of say 15 nodes per wavelength has to apply to this near field scale and not only to the incident wavelength; however, the near field scale is not known a priori and it depends on how deeply the evanescent field is excited, so it is difficult to anticipate how fine has to be the grid (a discussion of this point is presented in [21]).

In the homogenized problem, the near field effect is encapsulated in the interface parameters and this is possible since the near field is essentially a static field. Thus, the mesh size is limited by the usual rule on the incident field only. This is confirmed here; with a minimum incident wavelength of 30 m, a mesh size $\Delta x = 2$ m is sufficient to get a converged homogenized solution; next the validity of the field depends on how close the real near field is close to the static one, and this is lost progressively by increasing the frequency.
Now, let us inspect how different is the story for the mesh size needed in the real problem. Fig. 14 shows a zoom of the profiles along the centerline \( x = 0 \) \((y \in [50, 250] \text{ m})\) for decreasing mesh size \( \Delta x \). It is visible that the solution of the real problem continues to converge toward the solution of the homogenized problem (up to the error due to model) for \( \Delta x \) well beyond \( \Delta x = 2 \text{ m} \), and it has not converged before \( \Delta x = 0.25 \text{ m} \). This means that the evanescent field experiences rapid variations with a typical exponential decrease over lengths of few meters (\( \Delta x = 0.25 \text{ m} \) solves, as a rule of the thumb, variations over 4 meters of the evanescent field, to be compared to the 30 meters wavelength).

Figure 13: (a-c) Pressure profiles along the centerline \( x = 600 \text{ m} \) from Figs. 12, \( p^r \) (blue symbols) and \( p \) (black lines). (d) Error \(|p^r - p|/|p|\) calculated on the transmitted wave \( y \in [0490] \text{ m} \) as a function of \( \varepsilon = 2\pi f_0/c \) (an extra point at \( f_0 = 20 \text{ Hz} \) has been considered); dotted grey line shows the \( \varepsilon^2 \) law.

\( f_0 = 2.5 \text{ Hz} \)

\( f_0 = 5 \text{ Hz} \)

\( f_0 = 10 \text{ Hz} \)
In conclusion, the gain in replacing the real problem by the homogenized one at low frequency is simply given by the ratio between the geometrical size of the array \((h, e)\) on the wavelength. Besides, the smaller is the frequency, the better is the agreement between the homogenized and the real problem. It is obviously for these small \(\varepsilon\) values that the homogenized problem can mimic the real problem in the most efficient way.

The situation is more involved for intermediate frequencies (say \(\varepsilon\) of order unity). In this regime, the homogenized solution becomes less efficient to describe the real problem. Nevertheless, the gain in the numerical cost remains important; this is because the smallest scale to be resolved in the real problem is associated to the near field variation, and not anymore to the array size. Because higher frequency produces stronger scattering, the evanescent field may contain scales significantly smaller than the array size. For these higher frequencies, a compromise between the accuracy of the homogenized solution and the numerical gain has to be found, and this depends on the wanted precision. As an indication in the numerical gain at \(f_0 = 10\) Hz, a computational time of 1 minute for the homogenized problem corresponds to a computational time of 10 hours in the real problem. The extra time needed in the real problem is not only due to the smaller
mesh grid, but also to the smaller time step imposed by the CFL condition.

### 4.3. Variable homogenized interface

As previously said, the numerical implementation of the jump conditions (3) have been extended to the case of a curved line $\Gamma$. Below, we report results varying the curvatures. For small curvatures, this allows us to validate the numerical implementation of the jump conditions along a curved line; also, we inspect the error due to the model when increasing the curvature. For high curvatures, we expect the jump conditions to be modified. It is outside the scope of the present paper to derive such conditions, but let us estimate the maximum curvature below which we expect the jump conditions (3) to be unaffected. The parameters $(B, C_1, C_2)$ have been calculated in static problems to account for the boundary layers effects near the scatterers; if these boundary layers are significantly modified because of the local curvature, the parameters will be affected as well; it is worth noting that, if the case, the homogenized problem becomes more tricky since the parameters will vary along $\Gamma$ (if the curvature varies, that is for any curve $\Gamma$ except a circle). Fig. 15 shows the relative position of two -rectangular- scatterers for a local curvature $1/R$, producing a minimum distance of $\delta h$ between them (and $\delta h = h$ for infinite $R$, or zero curvature).

![Figure 15: Effect of the local curvature on the relative positions of two scatterers. $R$ is the local radius of curvature, producing a smallest distance $\delta h \leq h$ between the two scatterers (and $\delta h = h$ for $1/R = 0$).](image)

The modification of the boundary layers in the static problems involved to calculate $(B, C_1, C_2)$ are expected to depend of how $\delta h/h$ is far from 1. Note also the condition $\delta h > \varphi h$ for no overlapping of the scatterers. It is easy to see that this leads to

$$1 \geq \frac{\delta h}{h} = \frac{R - e/2}{R} > \varphi.$$  (46)
Now, in addition to the condition of no-overlapping which imposes \( e/R < 2(1 - \varphi) \), small values of \( e/R \)-values (or small \( 1 - \delta h/h \)) are expected to ensure the validity of (3), and this is what we shall inspect further.

To easily increase the curvature, we consider a curved line \( \Gamma \) in the form of a sinusoid with mean value at \( y_m = 500 \text{ m} \) and \( y_{\Gamma} = y_m + A \cos 2\pi(x - x_s)/D \) \((x_s = 600 \text{ m})\). We kept a fixed \( A = 10 \text{ m} \) value and varied \( D \) from \( D = 1000 \text{ m} \) and 125 m. The corresponding maximum local curvatures are given by \( \kappa = A(2\pi/D)^2 \), leading to a minimum local radius of curvature \( R \) from \( R = 2500 \text{ m} \) to 40 m. This smallest value of \( R \) corresponds to twice the minimum value imposed by non overlapping (46).

To begin with, we report in Figs. 16 and 17(a-c) the wavefields and the profiles on the centerline \( x = 600 \text{ m} \) in the real problem and in the homogenized problem for \( f_0 = 10 \text{ Hz} \). The reported time is \( t = 0.38 \) s, and the calculations have been performed in the same conditions as in Figs. 12-13). Here, we have considered \( D = 250 \text{ m} \) \((R = 160 \text{ m})\), \( D = 160 \text{ m} \) \((R = 65 \text{ m})\) and \( D = 125 \text{ m} \) \((R = 40 \text{ m})\). Corresponding values of \( 1 - \delta h/h \) are 0.06, 0.15 and 0.25 respectively, for a maximum allowed value of 0.5. By comparison of the results in Figs. 16-17 with those obtained with a straight line \( \Gamma \) in Figs. 12-13, it is visible that even a small curvature produces significant modification in the field pattern. This is particularly noticeable regarding the signal often referred as the "coda," which corresponds to the signal between the two main wavefronts directly transmitted and reflected by the interface. In the real problem, increasing the curvature of \( \Gamma \) enhances multiple scattering effects in the region of the scatterers, which feeds the coda region. Also noticeable is the fact that the homogenized solutions reasonably reproduce the main features of this coda region, although they are unable to reproduce its finest scales.

Finally, Fig. 17(d) shows the error as a function of \( 1 - \delta h/h \) where additional \( D \) values have been considered. The result is as expected: for \( \delta h \simeq h \), the effect of the curvature is incidental and the error remains the same as for a straight line \( \Gamma \); increasing further the curvature produces an increase in the error. In the presented case, this happens for \( 1 - \delta h/h \) above 10%, thus for a relative minimum distance between the scatterers 5 times smaller than the minimum distance of overlapping.

5. Conclusion

We have proposed a numerical method to implement jump conditions established within a homogenization approach. These jump conditions allow us to
Figure 16: Wavefields for a curved mean line $\Gamma$. Expect the shape of $\Gamma$, the calculations are identical to those of Fig. 12 for $f_0 = 10$ Hz. On (a-c), the left panels show the wavefields $p'$ in the real problem (with dotted white lines indicating $\Gamma$); the right panels the wavefields $p$ in the homogenized problem (with the white region indicating the homogenized interface).

replace the real problem of the wave propagation though an array of sound hard scatterers by an equivalent homogenized problem, much simpler to handle numerically. The numerical method has been validated considering a scattering problem for which an explicit solution is available, and comparisons of the solutions in the homogenized problem and in the real problem have confirmed the ability of the homogenized problem to advantageously replace the real one. Advantageously is meant with a computational time highly reduced; in the cases reported here, the computational time is typically reduced by a factor 500. In fact, and as previously said, accounting for the jump conditions requires additional calculations being time independent. Thus, they are done once and for all independently of the wave source and before the time resolution is considered. Then, the time resolution is not more demanding than for a problem of wave propagation in a homogeneous medium.

In terms of the possible extensions of the numerical method, several directions seem of interest to us. The first is rather incremental. We have considered thicknesses of the scatterers being small but sufficiently large with respect to the expected mesh size; this is because we assumed that points of certain stencils may fall within the homogenized interface (which is not resolved). If the scatterer thickness passes below the mesh size, the region of the interface will always be contained between two nodes; this case has been considered already, notably for
vanishing scatterer thickness [1] (the harmonic regime is considered in this reference) and it is strictly more simple to handle numerically. Nevertheless, because of the practical interest in such thin arrays (the gain in the numerical implementation is even more significant), this has to be done. The second extension is numerically not so demanding but it requires to adapt the homogenization approach. The jump conditions have been established for the scatterers being located along a straight line and we have inspected their possible extension to the case of a curved line. This has confirmed the intuitive idea that large local curvatures require a modified version of the jump conditions. It is worth noting that this would lead to a more involved homogenized problem since the curvature being defined locally along the mean line, the parameters entering in the jump conditions would be local as well. Nevertheless, and again in regard with the practical interest of such configurations, a generalization to scatterers located onto curved lines deserves interest.

Finally, several extensions concern the nature of the scatterers, and let us men-
tion two situations which are not trivial extensions of the present work. The first concerns scatterers associated to Dirichlet boundary conditions: it is typically metallic arrays illuminated by a polarized electromagnetic wave in the far infrared regime. In this case, effective boundary conditions at each side of the interface have to be considered, rather than jump conditions [18]. The second is a bolder extension of the present work. It concerns scatterers with material properties having high contrasts with respect to the surrounding medium, such that resonances inside the scatterers are possible. In this case, the jump conditions cannot be derived in the time domain. The calculations are done in the harmonic regime, revealing interface parameters being frequency dependent. Thus, the numerical implementation of these jump conditions in the time domain requires to handle these frequency dependent parameters.

Appendix A. Interface parameters \((B, C_1, C_2)\) for rectangular sound-hard scatterers

The interface parameters involved in the jump conditions (3) have been calculated for rectangular sound-hard scatterers in [20, 21]. As they are written in (3), they have the dimension of lengths and they read

\[
B = \frac{e}{1 - \varphi} - \frac{2}{\pi} h \log \left( \sin \frac{\pi(1 - \varphi)}{2} \right), \\
C_1 = e(1 - \varphi), \\
C_2 \simeq e(1 - \varphi) - \frac{\pi}{8} h (1 - \varphi)^2, \quad \text{if this leads to a positive value},
\]

and, as a rule of the thumb for the last parameter

\[
\begin{cases}
C_2 \simeq e(1 - \varphi) - \frac{\pi}{8} h (1 - \varphi)^2, & \text{if this leads to a positive value}, \\
C_2 \simeq e(1 - \varphi) - \frac{\pi}{8} h (1 - \varphi)^2, & \text{if this leads to a positive value},
\end{cases}
\]

for rectangular scatterers of length \(\varphi h\), with spacing \(h\) and thickness \(e\). Here \(\varphi\) is the filling fraction of the inclusion. A different definition is used in [20], where it refers to the filling fraction of the matrix, so \(\varphi\) is replaced here by \(1 - \varphi\). A more accurate estimation of \(C_2\) can be obtained by solving a so-called elementary problem, and a simple script to do so is provided in [20].

Appendix B. Scattering of a plane wave at oblique incidence on a plane homogenized interface

We consider a plane wave at oblique incidence \(\theta\) on the thick interface and the problem to solve is (5). We want to determine \(u(x, y, \omega)\) in (45), and \(u\) is defined
in (7). Below, we shall calculate the pressure $p$ afterwards $u = (v_x, v_y, p)^T$ is deduced using (5) written in the harmonic regime, with time dependence $e^{-i\omega t}$, whence

\[
\begin{align*}
  v_x(x, y, \omega) &= -\frac{i}{\rho \omega} \frac{\partial p}{\partial x}(x, y, \omega), \\
  v_y(x, y, \omega) &= -\frac{i}{\rho \omega} \frac{\partial p}{\partial y}(x, y, \omega).
\end{align*}
\]

(B.1)

For this one dimensional problem, the pressure field (7) reads

\[
p(x, y, \omega) = e^{i\omega x} \sin \theta/c \times \begin{cases}
  e^{-i\omega(y-e/2) \cos \theta/c} + &Re^{i\omega(y-e/2) \cos \theta/c}, & x > e/2, \\
  Te^{-i\omega(y+e/2) \cos \theta/c}, & x < e/2,
\end{cases}
\]

(B.2)

and we want to determine the scattering coefficients $(R, T)$. It is sufficient to inject (B.2) in the jump conditions (3) using (5), and setting the impedance $Z = \rho c \cos \theta$ and the parameters $(\alpha, \beta)$

\[
\alpha = \frac{\rho}{2} \left( C_1 \cos^2 \theta + C_2 \sin^2 \theta \right), \quad \beta = \frac{B}{2c} \cos \theta,
\]

(B.3)

to get the system satisfied by $R$ and $T$

\[
\begin{align*}
  (T - R) (1 + i\omega \beta) &= (1 - i\omega \beta), \\
  (T + R) (Z - i\omega \alpha) &= (Z + i\omega \alpha).
\end{align*}
\]

(B.4)

We get

\[
\begin{align*}
  R &= \frac{i\omega (\alpha + Z\beta)}{(Z + i\omega \alpha)(1 + i\omega \beta)}, \\
  T &= \frac{Z - \omega^2 \alpha \beta}{(Z + i\omega \alpha)(1 + i\omega \beta)},
\end{align*}
\]

(B.5)
from which \( |R|^2 + |T|^2 = 1 \). These expressions of \((R, T)\), together with (B.2) and (B.1) give the reference solution \( u(x, y, \omega) \) used in (45).


