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## NEW ALE APPLICATIONS IN NON-LINEAR FAST-TRANSIENT SOLID DYNAMICS

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### ABSTRACT

The arbitrary Lagrangian–Eulerian (ALE) formulation, which is already well established in the hydrodynamics and fluid-structure interaction fields, is extended to materials with memory, namely, non-linear path-dependent materials. Previous attempts to treat non-linear solid mechanics with the ALE description have, in common, the implicit interpolation technique employed. Obviously, this implies a numerical burden which may be uneconomical and may induce to give up this formulation, particularly in fast-transient dynamics where explicit algorithms are usually employed. Here, several applications are presented to show that if adequate stress updating techniques are implemented, the ALE formulation could be much more competitive than classical Lagrangian computations when large deformations are present. Moreover, if the ALE technique is interpreted as a simple interpolation enrichment, adequate – in opposition to distorted or locally coarse – meshes are employed. Notice also that impossible computations (or at least very involved numerically) with a Lagrangian code are easily implementable in an ALE analysis. Finally, it is important to observe that the numerical examples shown range from a purely academic test to real engineering simulations. They show the effective applicability of this formulation to non-linear solid mechanics and, in particular, to impact, coining or forming analysis.

**KEY WORDS** Arbitrary Lagrangian–Eulerian formulation Finite elements Non-linear continuum mechanics Time integration schemes Large boundary motion Applications

### INTRODUCTION

The arbitrary Lagrangian–Eulerian formulation (ALE) is concerned with the kinematic description (i.e. the relationship between the moving media and the computational grid) which is a fundamental consideration in determining a method for the numerical solution of multi-dimensional continuum mechanics problems. This formulation is now fairly well established in the fluid mechanics field, with special emphasis on hydrodynamics and fluid-structure interaction. Obviously, important lines of research are still open; for instance, large boundary motion analysis or optimal meshes and remeshing algorithms. But the most important challenge for the ALE technique lies, perhaps, in its extension to continuum mechanics in general, and, in particular, to non-linear solid mechanics where path dependent material behaviour is fairly common, e.g. plasticity.

Two classical descriptions are normally employed in continuum mechanics. The first one is Lagrangian, in which the mesh points coincide with the material particles. In this description,

no convective effects appear, and this considerably simplifies the numerical calculations; moreover, a precise definition of the moving boundaries and interfaces is obtained – recall that each element contains always the same amount of material. However, the Lagrangian description does not satisfactorily handle the material distortions that lead to element entanglement. On the other hand, the second description is the Eulerian viewpoint, which allows strong distortions without problems because the mesh is fixed with respect to the laboratory frame and the continuum flows through it. However, this latter approach presents two important drawbacks:

- (i) convective effects, which introduce numerical difficulties, arise due to the relative movement between the grid and the particles;
- and
- (ii) sophisticated mathematical mappings are required to follow the interface movement and they often lead to inaccuracies.

Because of the shortcomings of purely Lagrangian and Eulerian descriptions, arbitrary Lagrangian–Eulerian techniques were developed, first in finite differences, Noh<sup>1</sup> and Pracht<sup>2</sup>, among others; and then in finite elements<sup>3,4,5,6,7,8</sup>, among others. This approach is based on the arbitrary movement of the reference frame, which is continuously rezoned in order to allow a precise description of the moving interface and to maintain the element shape.

However, in all references cited previously, the application of the ALE formulation is restricted to inviscid or viscous fluids in both assumptions: compressibility or incompressibility. The advantages and power of this technique in fluid-structure problems are the reason for its introduction and popularization in finite element codes, see Donéa *et al.*<sup>3</sup> and Belytschko *et al.*<sup>4</sup>. But the fluid domain is the only one treated by the ALE formulation while the structure remains associated to the classical Lagrangian description. On the other hand, non-linear viscosity can only be taken into account in the context of generalized Newtonian fluids, Huerta and Liu<sup>7</sup>.

The reason for this bias in the applications of the ALE method, is the fact that in inviscid or viscous fluids, even the generalized Newtonian ones, the stress tensor is determined by the velocity field at every instant: the material has “no memory”. Due to the fact that in the ALE method material points and grid nodes may not coincide, obvious difficulties appear with “memory” materials. That is, materials where the stress is a function of state variables that differ from particle to particle because they are affected by the motion of the material points which possess different stress and strain histories. This is also the case for the Eulerian formulation where the same difficulties have precluded its applications to non-linear solid mechanics.

Only very recently, some attempts to treat non-linear path-dependent materials have appeared in the literature<sup>9,10,11,12</sup>. These approaches have in common that an implicit interpolation technique is needed. Obviously, this implies a numerical burden which may be uneconomical and may induce to give up ALE methods. This is particularly true in fast-transient dynamic analysis of solids where explicit algorithms are usually employed. Here several applications are presented to show that if adequate stress updating techniques are implemented<sup>13</sup> the ALE formulation could be much more competitive than classical Lagrangian computations. Moreover, if the ALE formulation is interpreted as a simple interpolation enrichment, adequate – in opposition to distorted or locally coarse – meshes are employed. Notice also that impossible computations (or at least very involved numerically) with a Lagrangian code are easily implementable in an ALE analysis.

The outline of the present paper is arranged as follows. First, the notation and fundamentals of the ALE description are introduced. A general overview of the governing issues in ALE is presented, it ranges from the kinematics (the fundamental concern of the ALE), conservation

equations, boundary conditions and their implementation in large boundary motion context, to equations of state with their peculiar implementation difficulties and, obviously, remeshing which is inherent to the ALE techniques. Then, the time integration scheme employed for the fast-transient dynamic analysis is discussed, with special emphasis on the particularities introduced by the ALE formulation. Finally, several numerical examples, both academic (designed to prove the accuracy of the computations) and with engineering applications, are described and discussed.

## NOTATION AND FUNDAMENTALS

### *Notation and preliminaries*

A continuous medium under motion is always formed by the same material points; however, its configuration may change with time. In order to describe its motion a one-to-one mapping relating the initial position of a material point  $\mathbf{X}$  with its actual position,  $\mathbf{x}$ , at time  $t$  is needed. This is usually done by means of the displacement vector

$$\mathbf{d} = \mathbf{x}(\mathbf{X}, t) - \mathbf{X} \quad (1)$$

The one-to-one mapping condition is formally insured by requiring that the Jacobian  $J = \det \left[ \frac{\partial x_i}{\partial X_j} \right]$  is non-vanishing. Note that standard indicial notation is adopted; lower-case subscripts denote the components of a tensor and repeated indices, summations over the appropriate range, usually the number of spatial dimensions. The material region is denoted by  $R_{\mathbf{X}}$  and it is related to the particles or material points,  $\mathbf{X}$ , while  $R_{\mathbf{x}}$  and  $\mathbf{x}$  denote the spatial region, also known as the laboratory configuration, and coordinates. They represent the configurations of the continuum at the initial instant and at time  $t$ , respectively.

Two classical viewpoints are considered to describe the motion defined by (1). The first one is Lagrangian in which  $R_{\mathbf{X}}$  is taken as the reference; that is, the reference sticks to the particles. The second one, known as Eulerian, uses the spatial configuration, symbolized by  $R_{\mathbf{x}}$ , as the reference; that is, the reference is fixed in the laboratory. In what follows, the displacements in Lagrangian form are denoted as  $\mathbf{d}^{**}(\mathbf{X}, t)$  while in an Eulerian description they are written as  $\mathbf{d}(\mathbf{x}, t)$ .

In the arbitrary Lagrangian–Eulerian (ALE) description, the computational frame is a reference independent of the particle motion and it may be moving with an arbitrary velocity in the laboratory system; the continuum viewed from this reference is denoted as  $R_{\chi}$ , and the coordinates of any point are denoted as  $\chi$ . Obviously, one-to-one transformations relating the reference to the material and spatial domains are needed. They can be represented, symbolically, as:

$$\Phi \begin{cases} R_{\chi} \times [0, \infty[ \rightarrow R_{\mathbf{x}} \\ (\chi, t) \mapsto \Phi(\chi, t) = \mathbf{x} \end{cases} \quad (2)$$

and

$$\Psi \begin{cases} R_{\chi} \times [0, \infty[ \rightarrow R_{\mathbf{X}} \\ (\chi, t) \mapsto \Psi(\chi, t) = \mathbf{X} \end{cases} \quad (3)$$

where  $t$  denotes time. These transformations present, as previously, non-vanishing jacobians and are schematically represented in *Figure 1*.

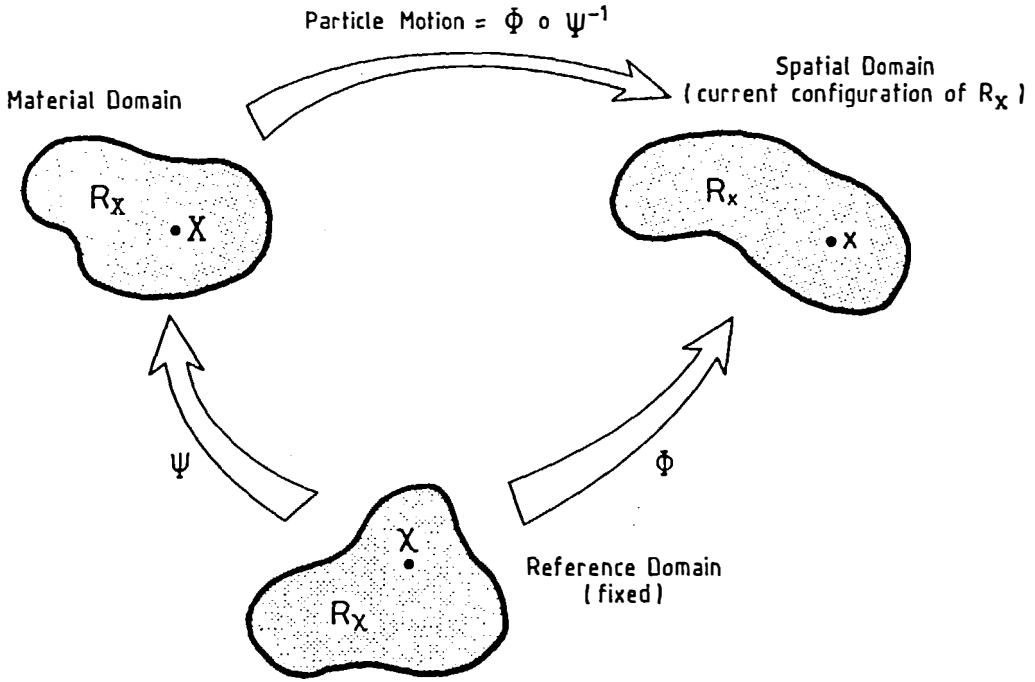


Figure 1 Schematic diagram for the domains and mappings in the ALE description

It is important to notice that the choice of an arbitrary reference does not imply that both  $\Phi$  and  $\Psi$  are chosen arbitrarily. In fact, once  $\Psi$ , or  $\Phi$ , is defined the other is automatically determined by the particle motion. For instance, if  $\Psi$  is the identity function, the reference domain,  $R_x$ , is equivalent to the material region,  $R_x$ ,  $\Phi$  is the particle motion and a Lagrangian description is used. On the other hand, when  $\Phi$  is the identity function at any time, the referential description corresponds to an Eulerian viewpoint and  $\Psi^{-1}$  is the particle motion. In any case, the generality of the ALE formulation allows to prescribe  $\Psi$  in a subset of  $R_x \times [0, \infty[$  while  $\Phi$  is defined on the rest. This will be discussed in detail in the remeshing section.

A graphical interpretation of the three different descriptions is presented in *Figure 2*. In the Lagrangian case, the mesh nodes follow the particle motion, in the Eulerian description the nodal points remain fixed in the laboratory during the complete process and, finally, with the ALE formulation the grid nodes move arbitrarily with respect to the particles and to the laboratory.

Once the reference is defined, the kinematics in the ALE description that link it to the classical Lagrangian and Eulerian descriptions are overviewed. A more detailed treatment is presented in References 5 and 7. For simplicity the fundamental equations governing the motion of the continuum are expressed in the spatial domain, thus involving spatial derivatives. On the other hand, both material and spatial domains are generally in motion with respect to the reference, while  $R_x$  is fixed throughout this formulation. Consequently, it is convenient to express the material time derivatives in a referential form. Finally, it is important to notice that the conservation equations involve material time derivatives. In what follows, a simple procedure to relate material time derivatives with referential time derivatives and spatial derivatives, originally devised in Reference 6, is presented.

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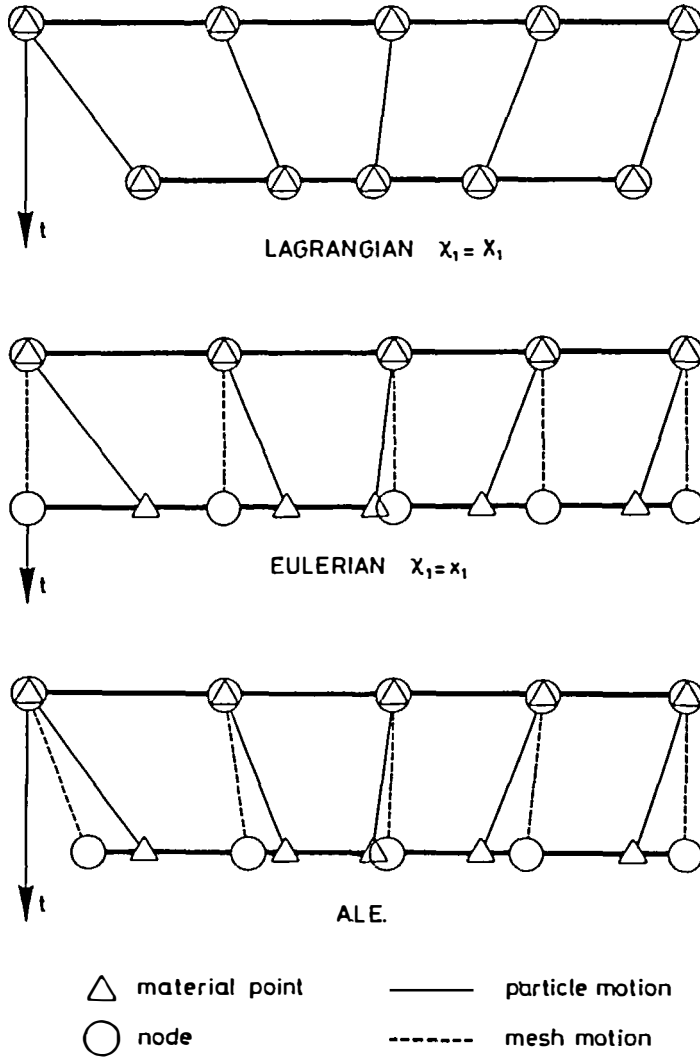


Figure 2 One dimensional example of Lagrangian, Eulerian and ALE mesh and particle motion

Consider a physical property  $F$  expressed as  $f(\mathbf{x}, t)$ ,  $f^*(\boldsymbol{\chi}, t)$ , and  $f^{**}(\mathbf{X}, t)$  in the spatial, referential and material descriptions, respectively. Its material derivative can be written as:

$$\frac{\partial f^{**}(\mathbf{X}, \cdot)}{\partial t} = \frac{\partial f^*(\boldsymbol{\chi}, \cdot)}{\partial t} + w_i \frac{\partial f^*}{\partial \chi_i} \quad (4)$$

where

$$w_i = \frac{\partial \chi_i(\mathbf{X}, \cdot)}{\partial t} \quad (5)$$

is the velocity of the material points in the referential domain.

The formal mathematical notation for time derivatives is usually substituted in the literature, since Donéa<sup>5</sup>, by a more engineering type of notation. That is, given  $f(\xi, t)$ ,

$$\frac{\partial f(\xi, \cdot)}{\partial t} = \frac{\partial f}{\partial t} \Big|_{\xi} \quad (6)$$

meaning, that the partial derivative is taken “holding  $\xi$  fixed”. This remark is important in order not to confuse this notation with the classical mathematical sense of  $|_{\xi}$ , which is “particularized at  $\xi$ ”. Moreover, to simplify the subsequent developments, the *star* notation used to distinguish the three configurations (spatial, referential, or material) will be dropped.

Equation (4) relates the material time derivative with the referential time derivative. However, spatial derivatives are desired instead of derivatives of  $f$  with respect to  $\chi$ . Equation (4) is further simplified by means of the following definitions of material velocity,  $v$ , and mesh velocity,  $\hat{v}$ :

$$v_i = \frac{\partial x_i}{\partial t} \Big|_{\mathbf{x}} \quad (7a)$$

and

$$\hat{v}_i = \frac{\partial x_i}{\partial t} \Big|_{\chi} \quad (7b)$$

respectively. If the physical property is the spatial coordinate  $\mathbf{x}$ , (4) and (7) yield:

$$v_i = \hat{v}_i + w_j \frac{\partial x_i}{\partial \chi_j} \quad (8)$$

or

$$c_i = v_i - \hat{v}_i \quad (9)$$

where

$$c_i = w_j \frac{\partial x_i}{\partial \chi_j} \quad (10)$$

is the convective velocity. While  $\mathbf{w}$  is the material velocity in the reference,  $\mathbf{c}$  is the relative velocity of the particles with respect to the mesh in the laboratory system. Finally, substituting (10) into (4) and using the chain rule, yields the classical relationship between the material time derivative, the referential time derivative and the spatial derivative:

$$\frac{\partial f}{\partial t} \Big|_{\mathbf{x}} = \frac{\partial f}{\partial t} \Big|_{\chi} + c_i \frac{\partial f}{\partial x_i} \quad (11)$$

#### *ALE formulation of the conservation laws*

Under the ALE description, the conservation laws that govern the motion of the continuum are written, in strong form, as:

*continuity*

$$\frac{\partial \rho}{\partial t} \Big|_{\chi} + c_j \frac{\partial \rho}{\partial x_j} = -\rho \frac{\partial v_j}{\partial x_j} \quad (12a)$$



momentum

$$\rho \frac{\partial v_i}{\partial t} \Big|_x + \rho c_j \frac{\partial v_i}{\partial x_j} = \frac{\partial \sigma_{ij}}{\partial x_j} + b_i \quad (12b)$$

energy

$$\rho \frac{\partial e}{\partial t} \Big|_x + \rho c_j \frac{\partial e}{\partial x_j} = \sigma_{ij} \frac{\partial v_i}{\partial x_j} + \rho a - \frac{\partial}{\partial x_i} \left( k_{ij} \frac{\partial \theta}{\partial x_j} \right) \quad (12c)$$

where  $\rho$  is the density,  $\sigma$  is the Cauchy stress tensor,  $b$  is the body force per unit of volume,  $e$  is the specific internal energy,  $a$  is the internal heat generation,  $k$  is the thermal conductivity tensor, and  $\theta$  is the thermodynamic temperature. The derivation of the above equations may be found in References 14, 15 and 7.

It is important to notice that the right hand side of (12) is written in classical Eulerian form while the arbitrary motion of the mesh is only reflected in the left hand side. The purpose of mixing material time derivatives, referential time derivatives and spatial derivatives is now clear from (12): Cauchy stresses are employed (symmetric and defined by the usual engineering constitutive equations), the thermal conductivity is defined in the current configuration (easy implementation for nonlinear anisotropic materials), and the weak form of (12) is obtained after integration over the spatial domain (natural boundary conditions are easily implemented on the actual configuration), see Reference 7.

The origin of (12) and their similarity with the Eulerian equations have induced some authors<sup>4</sup> to name this method the *quasi-Eulerian* description.

At this point the equations that must be satisfied are the conservation laws, (12), and the mesh motion equation, (9), apart from boundary conditions and constitutive equations that will be discussed later. The definition of the mesh velocity,  $\hat{v}$ , is thus mandatory and it is the basis of the remeshing techniques that will be presented in a subsequent section.

### Boundary conditions

One of the main applications of the ALE formulation is on large boundary motion problems. This section is mostly concerned with these analyses and therefore only the boundary conditions directly related with the momentum equation are discussed. In fact, it is assumed that heat flux and/or temperature, as well as density on the inflow boundaries are adequately prescribed. Or, more rigorously, well-posed boundary conditions based on an analysis of the characteristics is done along the boundaries, see for instance Reference 16. In any case, material surfaces which centre the interest of the following discussion, are equivalent to the so-called solid-wall boundaries in the fluid dynamics literature. That is, velocities and heat flux or temperature are given; the *only* particularity is that velocities vary with time and that an extra equation must be solved (the unknown position of the free surface).

In fact, the boundary conditions are obviously related to the problem, not to the description employed. Thus, the same boundary conditions used in Eulerian or Lagrangian descriptions are employed in an ALE formulation. That is, along the boundary of the domain, kinematical and dynamical conditions must be defined. Usually, this is formalized as:

$$v_i = g_i \quad \text{in } \partial R_x^g \quad (13a)$$

$$\sigma_{ij} n_{xj} = h_i \quad \text{in } \partial R_x^h \quad (13b)$$

where  $g$  and  $h$  are the prescribed boundary velocities and tractions, respectively;  $n_x$  is the outward unit normal to  $\partial R_x$ , and  $\partial R_x$  is the piecewise smooth boundary of the *spatial* domain,  $R_x$ .  $\partial R_x$  is in fact composed of two distinct subsets  $\partial R_x^g$  and  $\partial R_x^h$ . As usual, stress conditions on the

boundaries are “natural boundary conditions”, and thus, they are automatically included in the weak, or variational form of (12).

However, if part of the boundary is composed of a material surface, then a mixture of both conditions is sometimes required<sup>17</sup>. Firstly, the conditions required on a material surface are:

- (a) no particles can cross it, and
- (b) stresses must be continuous across the surface (if a net force is applied to a surface of zero mass the acceleration is infinite).

And two types of material surfaces are usually present: free surfaces, and solid wall boundaries which may be frictionless or not.

Along a solid wall boundary the particle velocity is defined (or coupled to a structure system in fluid-structure interaction problems). The requirement that no particles can cross it, can be simply verified if  $w$  is prescribed equal to zero along that boundary, i.e.  $v = \hat{v}$  from (8) which consists in defining the material surface as Lagrangian. However, this condition may be relaxed imposing only the necessary condition:  $w$  equal to zero along the normal to the boundary. The latter allows remeshing tangent to the boundary, the advantages of this type of relaxed boundary condition are evident in the pulling example shown later. The dynamic condition is automatically verified along rigid boundaries, but it presents the classical difficulties in fluid-structure interaction problems when compatibility at nodal level in velocities and stresses is required. An extensive discussion for this case is found in Reference 14.

Along the other type of material surface, i.e. free surface, problems arise because its position is unknown. Thus, the kinematic and dynamic conditions must be imposed and *solved*. The first one, the kinematic condition can be formalized as:

$$\mathbf{c} \cdot \mathbf{n}_x = 0 \quad (14)$$

in an Eulerian description. However, since the boundary is moving, its equivalent in referential form is preferred because the referential domain is fixed, namely:

$$\mathbf{w} \cdot \mathbf{n}_x = 0 \quad (15)$$

where  $\mathbf{n}_x$  is the exterior normal to the referential domain. While the second one, the dynamic condition expresses the stress-free situation:

$$\sigma_{ij} n_{xj} = 0 \quad (16)$$

and, as mentioned earlier, it is directly taken into account by the weak formulation.

In conclusion, free surface problems are the only ones that introduce a new equation, (15), that must be verified along that boundary. It is obvious that this new equation has a strong influence on the remeshing techniques employed.

### *Equations of state*

The initial boundary value problem is not defined until the state equations which reflect the behaviour of the continuum, are specified. Here two of such equations are needed, the first one relates temperature and density to energy:

$$e = e(\rho, \theta) \quad (17)$$

The second relates stress and/or its derivatives to temperature, and velocity and/or its derivatives. The latter constitutive relationship admits several formal representations, here only two of them are discussed. They include, however, most of the engineering materials in fluid or solid mechanics.



First of all, the Cauchy stress tensor is defined as a function of velocity, temperature and density fields:

$$\boldsymbol{\sigma} = \mathbf{s}(\theta, \rho, \mathbf{v}) \quad (18)$$

For instance, any generalized Newtonian fluid, Bird *et al.*<sup>22</sup>, falls in this category because its constitutive relationship is written as:

$$\sigma_{ij} = p\delta_{ij} + \mu \left( \frac{\partial v_i}{\partial x_j} + \frac{\partial v_j}{\partial x_i} \right) \quad (19)$$

where  $\delta_{ij}$  is the Kronecker delta;  $\mu$  is the dynamic viscosity which is shear rate dependent;  $p$  is the pressure which is uniquely determined by density and temperature for compressible fluids or by the velocity field for incompressible fluids. In any case, it is important to notice that (18) represents a class of *no memory* materials. Observe that the stress tensor is uniquely determined at each spatial point given the other instantaneous fields.

The other formal representation of the stress constitutive relationship is associated with path-dependent materials, or materials *with memory*. It relates the material time derivative of the Cauchy stress tensor with the same material fields as previously and with the stress field:

$$\left. \frac{\partial \boldsymbol{\sigma}}{\partial t} \right|_{\mathbf{x}} = \mathbf{r}(\theta, \rho, \mathbf{v}, \boldsymbol{\sigma}) \quad (20)$$

Any of the frequently employed rate type constitutive equations may be written in the previous form. Therefore (20) models a wide range of solid materials and problems, going from small strain linear elasticity to strongly non-linear large strain elasto-plasticity. For instance, any hypo-elastic material can be defined as behaving in the following way:

$$\left. \frac{\partial \sigma_{ij}}{\partial t} \right|_{\mathbf{x}} = \Delta \sigma_{ij}^c + -W_{ik}\sigma_{kj} + \sigma_{ik}W_{kj} \quad (21)$$

where  $\Delta \sigma_{ij}^c$  is the objective increment of stresses and represents the part of  $\boldsymbol{\sigma}$  due to actual straining of the material (“pure deformation”), i.e.:

$$\Delta \sigma_{ij}^c = C_{ijkl}v_{(k,l)} \quad (22)$$

$\mathbf{C}$  is the material response matrix which usually depends on the stress,  $\boldsymbol{\sigma}$ ;  $v_{(k,l)}$  are the components of the velocity stretch tensor,  $v_{(i,j)} = \frac{1}{2} \left( \frac{\partial v_i}{\partial x_j} + \frac{\partial v_j}{\partial x_i} \right)$ . The rest of the terms of (21) are associated to the rotation of the stress tensor. Two formulations are usually employed, the Green–Naghdi formulation if  $\mathbf{W}$  is taken as the rate of rotation matrix, or the more usual Zaremba–Jaumann–Noll or co-rotational formulation when  $\mathbf{W}$  is the spin tensor. In the latter, the components of  $\mathbf{W}$  are simply  $v_{[i,j]} = \frac{1}{2} \left( \frac{\partial v_i}{\partial x_j} - \frac{\partial v_j}{\partial x_i} \right)$ . Usually, these terms associated to the rotation are written as:

$$v_{[i,k]}\sigma_{kj} - \sigma_{ik}v_{[k,j]} = S_{ijkl}v_{[k,l]} \quad (23a)$$

where

$$S_{ijkl} = \frac{1}{2}(\sigma_{ii}\delta_{jk} + \sigma_{jj}\delta_{ik} - \sigma_{ik}\delta_{jl} - \sigma_{jk}\delta_{il}) \quad (23b)$$

The generalized fourth order tensor  $\mathbf{C}$  which represents the material response, can be derived in terms of the Jaumann rate of the Cauchy stress tensor:

$$\mathbf{C}_{ijkl} \equiv \mathbf{C}_{ijkl}^c \quad (24a)$$

or in terms of the Truesdell rate, as:

$$\mathbf{C}_{ijkl} \equiv \mathbf{C}_{ijkl}^t + \mathbf{C}_{ijkl}^* \quad (24b)$$

In the previous equation the tensor  $\mathbf{C}^*$  is needed to insure the objectivity of  $\mathbf{C}$ , and it is defined by:

$$\mathbf{C}_{ijkl}^* = -\sigma_{ij}\delta_{kl} + \frac{1}{2}(\sigma_{il}\delta_{jk} + \sigma_{jl}\delta_{ik} + \sigma_{ik}\delta_{jl} + \sigma_{jk}\delta_{il}) \quad (24c)$$

Finally, in order to be consistent with the ALE formulation the material time derivative in (21) is replaced by the referential one making use of (11), which yields:

$$\left. \frac{\partial \sigma_{ij}}{\partial t} \right|_{\mathbf{z}} = -c_k \frac{\partial \sigma_{ij}}{\partial x_k} + C_{ijkl} v_{(k,t)} + S_{ijkl} v_{[k,\eta]} \quad (25)$$

Notice that now, in opposition to (18), the constitutive equation must be integrated, and the decomposition of the stress rate in three terms, transport, pure deformation and the rotational part, should be exploited during the time integration.

The generalization to elasto-plastic materials is readily obtained by defining a yield surface and supposing that the strain rate is linearly separable into elastic and plastic components. It is very important to notice that the hardening rules which explicitly define the evolution of the yielding surface during plastic deformation, are usually written in incremental or rate form. That is, the yielding limit in isotropic hardening or the back stresses in the kinematic hardening are described by an equation such as (20), and consequently the same time integration algorithm is used for the stress tensor and related variables. Note also that (20) or (25) are not exclusive of path-dependent materials but Hookean linear elasticity, for instance, is also included. Consequently, an efficient integration algorithm for (25) can uniformly treat most of the structural mechanics problems.

Apart from physical considerations, this classification is also numerically sound, because the algorithms associated to (18) or to (20) are completely different; in fact, the classification is intrinsically related to the ALE formulation. Classically, the so-called phenomenological mathematical models represented by (18) where no updating is necessary, are used in fluid mechanics where an Eulerian formulation is usually employed. On the other hand, incremental constitutive relations are common in structural/solid mechanics and are normally associated to a Lagrangian formulation. The updating of any physical property in a Lagrangian description is simple because the reference follows the particles; while in the ALE or the Eulerian formulation such an update is much more involved. This explains why the ALE formulation was naturally developed in fluid mechanics and only recently it is applied in solid mechanics problems<sup>9,10,11,12</sup>.

### Remeshing

The equations that must be solved are (12), (13), (17) and (18) or (20) but also (9) is necessary and therefore, the resolution is only possible if the mesh velocities are given in the domain. The remeshing techniques are concerned with the definition of  $\hat{\mathbf{v}}$ . For instance, if  $\hat{\mathbf{v}} = \mathbf{0}$ , the Eulerian description is imposed, but if  $\hat{\mathbf{v}} = \mathbf{v}$  is prescribed, the Lagrangian description is used. It is obvious that finding the best choice for these velocities and a low cost algorithm for updating the mesh constitutes one of the major problems of the ALE description (cost meaning, here, computer time and computer storage).

As a consequence of the discussion on boundary conditions, two cases are considered. The first one assumes that all the boundaries of the domain have a known position at every instant, this includes Eulerian inflow/outflow boundaries, prescribed boundary motion of material surfaces and solid-wall boundaries. The second is associated to unknown free surfaces on the boundary of the domain (or material surfaces in general), and will be reduced to the former once the position of the free surface is known. This means verifying the kinematic condition of the free surface, namely (14) or (15). These equations are readily verified if the material surface is defined as Lagrangian, i.e.  $\mathbf{c} = \mathbf{w} = \mathbf{0}$ ; this is useful in structural mechanics problems. However, in large boundary motion problems, the element size along these Lagrangian surfaces can induce prohibitive computer cost or numerical inaccuracy. Therefore a relaxed condition expressed by (14) and (15), is recommended, see also Huerta and Liu<sup>17</sup>. In this respect several important remarks should be advanced:

- Equation (14) or (15) are defined along the material surface, therefore, these equations are not solved in the complete domain but only along free surfaces. The extra equation induced by the unknown position of the boundary is consequently less costly than the other field equations.
- The general formulation presented here does not describe the material surface with an elevation (over a datum level) parameter, thus vertical or folded surfaces (such as the ones present in coating flows or shell impact), for example, may be studied (see also the dam break problem in Huerta and Liu<sup>7</sup>).
- Equations (14) and (15) are scalar and only relate mesh and particle motion normal to the material surface. This implies that the user has the choice of deciding the mesh motion along the material surface. This is referred to as mesh sliding along the boundary and has the important advantage of precluding the use of pure Lagrangian nodes on these surfaces. Most of the examples shown later make use of these remeshing features but in the pulling analysis the advantages of such a procedure are clearly outlined. Finally, it should be noticed that the choice between (14) or (15) obeys to numerical efficiency and user's preferences.

At this point, the case of unknown free surfaces is reduced to the one where all the boundaries of the domain are fixed or have a known prescribed motion. Therefore, the continuous remeshing is completely defined once  $\hat{\nu}$  is given in the interior of the domain. This can be done by simple ad hoc formulae, using geometrical considerations<sup>18</sup>, solving potential equations that maintain element regularity<sup>19,11</sup>, or any other mesh generation algorithms that conserve the element connectivity. Most of these remeshings are based on defining new locations for the nodes, and then computing the mesh velocities by finite difference approximation (i.e. increment of displacement over increment of time). If structured meshes are used, the authors recommend to use simple ad hoc formulae where the mesh velocity is linearly interpolated between the velocities at both ends of the inter-element lines. This is an extremely simple and efficient algorithm which maintains element regularity and it is obviously more cost effective than solving potential equations previous to the evaluation of the mesh velocities.

## TIME INTEGRATION

In this section a brief discussion on the time integration schemes is presented. Keeping in mind the fast-transient dynamic context, the second order in time integration scheme presented in Reference 20 is generalized for the integration of the constitutive equation within the ALE formulation. After the finite element integration of the momentum equation, (12b), the system

of algebraic ordinary differential equations that must be integrated, is:

$$\mathbf{M}\mathbf{a} = \mathbf{f}^{\text{ext}} - \boldsymbol{\eta}(\mathbf{v}) - \sum_e \int_{R_e} \mathbf{B}^T \boldsymbol{\sigma} \, dx \quad (26)$$

where  $\mathbf{M}$  is the mass matrix;  $\mathbf{a}$  is the nodal acceleration vector;  $\mathbf{f}^{\text{ext}}$  is the vector of externally applied loads;  $\boldsymbol{\eta}(\mathbf{v})$  is the convective vector (relative motion between mesh and particles); and the last term is the internal force vector which is left as the sum of the element contributions. The element internal force vector is evaluated by means of the matrix of shape function derivatives,  $\mathbf{B}$ , and the Cauchy stress tensor  $\boldsymbol{\sigma}$ .

Since an explicit scheme is sought, the mass matrix,  $\mathbf{M}$ , is substituted by the lumped (diagonalized) mass matrix,  $\mathbf{M}^L$ , uncoupling the system defined by (26). Then the stability condition must be enforced; it is associated to the minimum Courant number of equivalently to the maximum eigenvalue of the system. In fact, in elasto-plastic analysis, this stability condition is related to the minimum element size and, as it will be shown in the numerical examples, the ALE formulation is advantageous because it allows to maintain during the deformation process regular shaped elements.

On the other hand, the constitutive equation, (25) for elasto-plastic materials or (20) in general, must also be integrated. The finite element interpolation and integration for the stresses is different from those of velocities or displacements. In fact, stresses are needed at each quadrature point for further numerical integration of the internal work, see (26). In a similar manner to Reference 11 a multiple stress collocation technique is used in the weak form, where the collocation points do coincide with the quadrature points<sup>13</sup>. The induced system of algebraic equations is explicit and trivial (the mass matrix coincides with the identity matrix), it may be written as:

$$\dot{\boldsymbol{\tau}} = \mathbf{r} - \boldsymbol{\eta}^\sigma \quad (27)$$

where  $\dot{\boldsymbol{\tau}}$  represents the vector of rate of stresses at the quadrature points,  $\mathbf{r}$  is the Lagrangian stress rate which includes only the pure deformation and rotation part, while  $\boldsymbol{\eta}^\sigma$  is associated to the transport of stresses, see (25).

As a matter of fact, a split-step algorithm is used to integrate in time this last system of ordinary differential equations. First of all, a pseudo-Lagrangian stress is obtained by simply integrating the Lagrangian stress rate,  $\mathbf{r}$ ; that is, assuming zero convective velocities. Then, the pseudo-Lagrangian stress is convected adding the contribution due to  $\boldsymbol{\eta}^\sigma$ . The particular expression for  $\boldsymbol{\eta}^\sigma$  depends on the time scheme employed, any numerical formulation for first order hyperbolic or conservation equations may be implemented. Here a Lax–Wendroff and a Godunov-type techniques were adopted to the particular nature of the stress fields (discontinuous element to element) and the desired numerical constraints (explicit code)<sup>13</sup>.

In order to maintain a central difference scheme for the time integration of (26) and (27), an uncoupled algorithm is devised. If accelerations are computed from (26) at time  $t^n$  then velocities are evaluated at  $t^{n+1/2}$  by:

$$\mathbf{v}^{n+1/2} = \mathbf{v}^{n-1/2} + \Delta t \mathbf{a}^n + \mathcal{O}(\Delta t^2) \quad (28a)$$

Once the velocities are known at the half step, (27) is enforced. The stress rate is then computed at  $t^{n+1/2}$  and consequently stresses and displacements are simply updated by equations similar to the previous one, namely:

$$\boldsymbol{\tau}^{n+1} = \boldsymbol{\tau}^{n-1} + \Delta t \dot{\boldsymbol{\tau}}^{n+1/2} + \mathcal{O}(\Delta t^2) \quad (28b)$$

$$\mathbf{d}^{n+1} = \mathbf{d}^{n-1} + \Delta t \mathbf{v}^{n+1/2} + \mathcal{O}(\Delta t^2) \quad (28c)$$

With these new values, the right hand side term in (26) as well as the mass matrix can be evaluated at  $t^{n+1}$ ; thus, the acceleration is computed at the new time step ( $n + 1$ ) and the process repeated as many times as needed.

The time scheme presented has the advantage that both conservation of momentum and the constitutive equation are solved separately with the best suited technique in each case. For instance while simple updating techniques can be used in (26) if the relative motion of the mesh and the particles is small, on the contrary the stress (and stress related variables such as the yield stress) sensitivity to convection imposes sophisticated updating schemes. Moreover, as mentioned previously, the interest in interpolating the stresses directly at the quadrature points, precludes a simple coupling between the second order ordinary differential equations in (26) and the first order ones (27). Finally, such an algorithm allows second order accuracy and it is easily generalized to time partitioning computations. It must be noticed, however, that the second order accuracy is lost in variable time stepping computations.

## NUMERICAL EXAMPLES

In this section the ALE formulation and the explicit time integration procedure are applied to several problems that range from purely academic test to engineering simulations. The first one is an elastic–plastic one-dimensional wave propagation problem first reported in Reference 9. This problem which has an obvious analytical solution in the elastic case allows to compare the numerical results with the exact solution. Moreover, the effectiveness of the formulation and in particular of the convection procedures, can be easily established.

The second example is the well known bar impact benchmark test. It is a classical test for impact codes and a full large strain demonstrative computation. It allows the comparison of Lagrangian and ALE results and presents a full range of boundary surfaces going from an axis of symmetry to free material surfaces with large boundary motion. A simple transformation of this example induces the following numerical simulation where necking appears and the effectiveness of the ALE formulation is demonstrated compared with the Lagrangian description.

Finally, an engineering example is shown. It simulates a coining process which is difficult and tedious to model a classical update Lagrangian description. However, it does not present any difficulty in the context of an ALE formulation and comparisons between a fast (dynamic) die velocity and a slow (quasi-static) die velocity are shown.

### *One-dimensional stress wave problem*

This elastic(-plastic) one-dimensional stress wave problem is presented here to assess the ALE formulation in the structural mechanics context. The material properties and element size are chosen equal to those used in Reference 9, for easier comparisons. Note, however, that the stress update schemes implemented here do not need special numerical parameters “à la” Streamline Upwind Petrov–Galerkin.

A schematic problem statement is shown in *Figure 3*, where the constants for the isotropic hardening material are given. The infinitely long elastic(-plastic) rod is discretized in 400 elements of 0.1 size. The problem is assumed isothermal and constant density is supposed throughout. The propagation of a square compression stress wave (–100 in amplitude, 4.5 in width) is simulated under several descriptions: Lagrangian, Eulerian and ALE. For the latest two arbitrarily chosen mesh velocities are imposed, one negative (CASE A) and one positive (CASE B). Since the purpose of this simulation is to provide a severe test to the formulation and the algorithms the mesh velocity is taken equal to one fourth of the speed of sound, notice that the

# 1-D ELASTIC (-PLASTIC) WAVE PROPAGATION

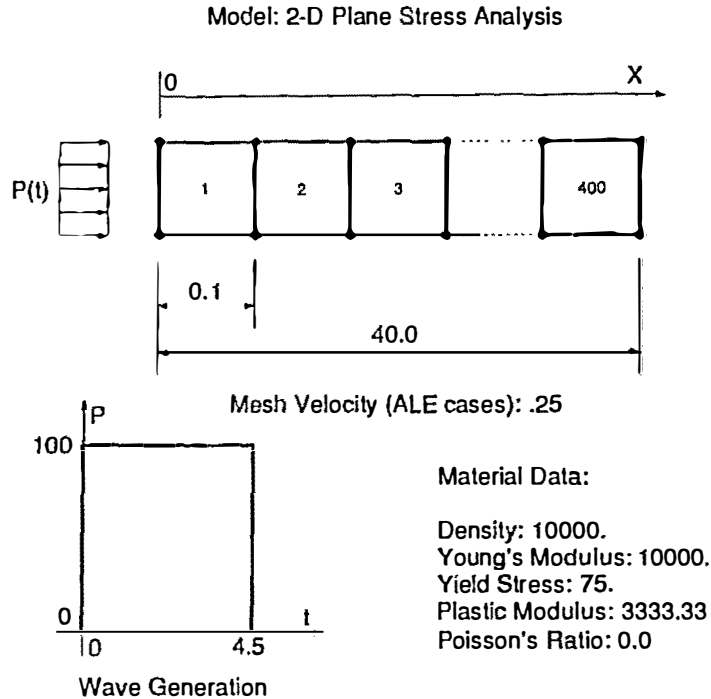


Figure 3 Schematic statement of the one-dimensional stress wave problem

material velocity of the particles is almost negligible in front of the mesh velocity. In fact, in the ALE cases the mesh velocity is initially set to zero (Eulerian description) and suddenly modified to a constant value (0.25 in modulus) after  $t = 24$ . The domain and duration of the simulation are such that no reflected waves appear, and the stress waves are superimposed in the figures to compare them.

Figure 4 shows the comparison between all cases in elastic and elasto-plastic regimes for the two stress update procedures, the Lax-Wendroff (L-W) type algorithm and the Godunov-type (G) update. Several remarks can be advanced, first of all, the implemented stress update algorithms introduce high frequency damping, as expected, and thus, the *wiggles* are almost suppressed. On the other hand, the monotonic algorithm of the Lagrangian description (no artificial damping was implemented) produces the unrealistic oscillations. Note that the Eulerian description, where the convection velocity is the particle velocity, presents a similar behavior because the particle velocity is almost negligible; the amplitude of the oscillations is nonetheless reduced.

To see the influence of the mesh motion direction, i.e. CASE A versus CASE B, Figure 5 presents the superposition of both cases for every stress update algorithm. Notice that the Godunov-type algorithm is insensitive to the mesh direction. On the other hand, the Lax-Wendroff technique which captures sharper fronts, shows small differences between both cases.

Finally, *Figure 6* shows the influence of the time increment in the solution. As expected, the lower the Courant number the larger the relative phase error, in the L-W scheme. However, the solution is still very adequate since the dissipative character of the scheme damps out the shorter wavelengths. Note also that this time dependency is negligible in the Godunov-type of update.

#### *Bar impact benchmark test*

The bar impact problem is chosen here for several reasons: it is a standard benchmark problem for fast-transient dynamic computer codes, although no analytical solution exist the ALE results can be compared to the Lagrangian ones, and the advantages of the proposed formulation are clearly outlined.

A cylindrical bar of radius 3.2 mm and length 32.4 mm impacts a rigid frictionless wall at an initial velocity of 227 m/s. The material is assumed elasto-plastic with  $E = 117$  Gpa,  $\nu = 0.350$ ,  $\sigma_y = 400$  MPa,  $\rho = 8930$  kg/m<sup>3</sup>, and a plastic modulus  $E_p = 100$  MPa. The time-step is variable and automatically chosen by the code to maintain the numerical stability, a maximum Courant number of 0.5 is imposed in the *smaller* element. An axisymmetric mesh of 250 bilinear elements is employed.

In *Figure 7*, the calculated deformed shapes for the Lagrangian (left) and ALE (right) descriptions are compared at every 10  $\mu$ s (only the mesh is shown) up to the final 80  $\mu$ s. Only the Godunov-type stress update algorithm is shown because negligible differences are found comparing with the Lax–Wendroff technique. The difference between the Lagrangian and ALE calculations for the radius and axial length are 0.5% and 0.4%, respectively.

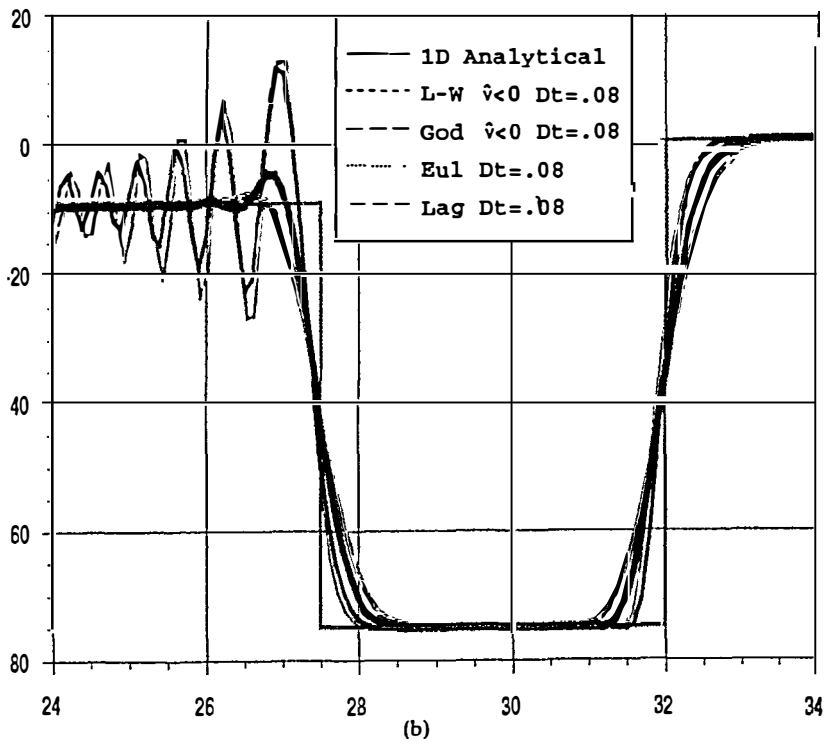
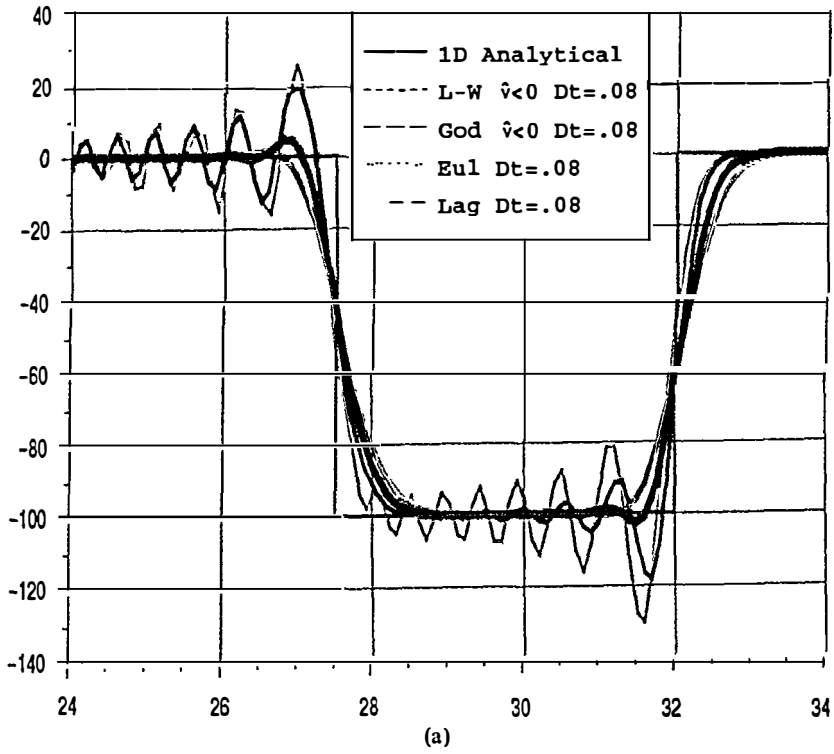
Important differences, however, are observed in the time-step,  $\Delta t$ , employed, see *Figure 8*. The uniformity of the mesh, maintained by the Giuliani<sup>18</sup> algorithm, allows larger and more uniform time increments. This obviously induces an important difference in the computer time required for the simulation. While the Lagrangian description needed 3 hours 44 min (31 700 time-steps), the ALE formulation used 27 min (1 900 time-steps).

The adequacy of the stress updating algorithm is stressed in the last figure of this example. The distribution of final yield stress (directly related to the plastic strain) is plotted for the Lagrangian case and the ALE analysis. While *Figure 9a* shows a remarkable similarity between both simulation (observe that the contour lines are almost coincident), *Figure 9b* shows important differences between the plastic zones at the final instant. In the latest case, no stress update procedure is implemented but the remeshing is performed. This is obviously a *wrong* calculation because no history dependent state variables are transported and the path-dependency is not reflected in the computations. However the height and width of the final bar present differences of only 1.3% and 6.5%, respectively, with the Lagrangian results. This suggests that the measures of final height and width are weak indicators of the goodness of the algorithm and that other results, such as the distribution of final yield stress, must be compared when dealing with this benchmark test.

#### *Bar pulling and necking analysis*

This example is a simple extension of the previous one. The same geometry and material are taken, even the same initial velocity is prescribed but now the sign is opposite. Thus the present example simulates a sort of pulling test. It has been chosen because it presents another possible advantage of the ALE formulation. While the Lagrangian analysis suffers from excessive element distortion, precisely where the necking occurs, the ALE description allows a regular element size distribution everywhere, see *Figure 10*. *Figure 11* compares the yield stress distribution at





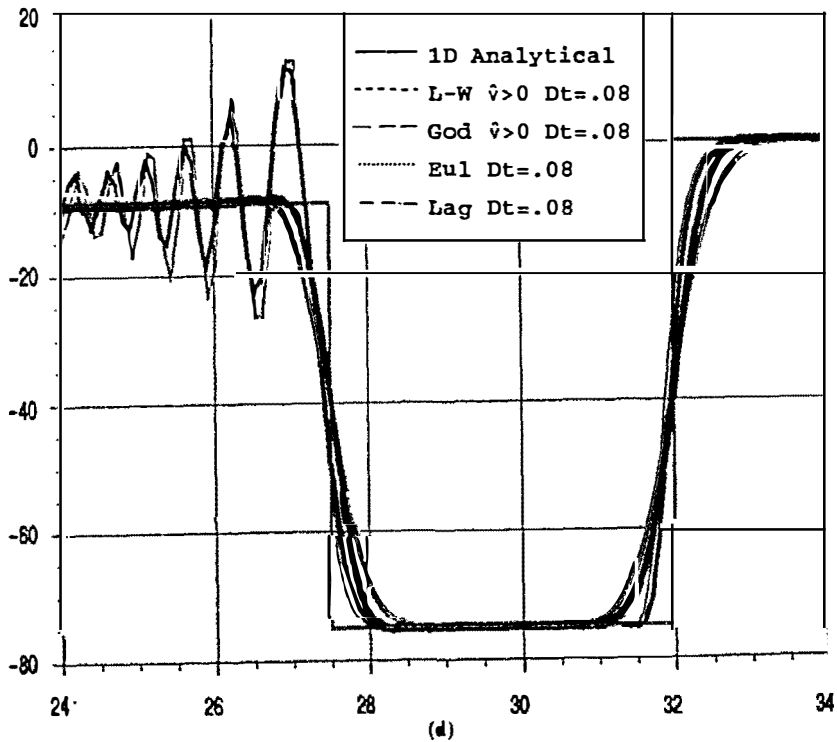
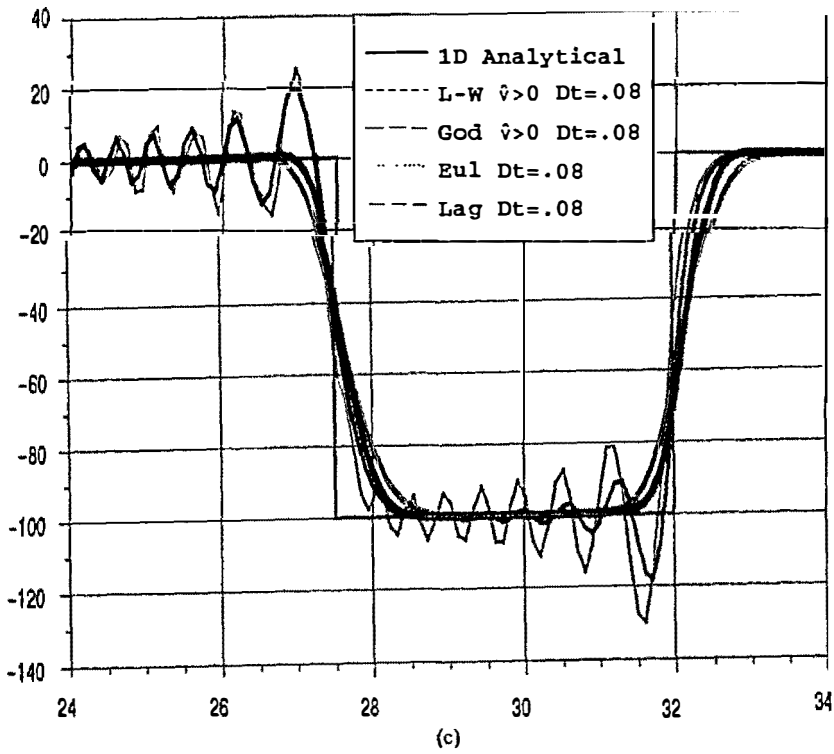
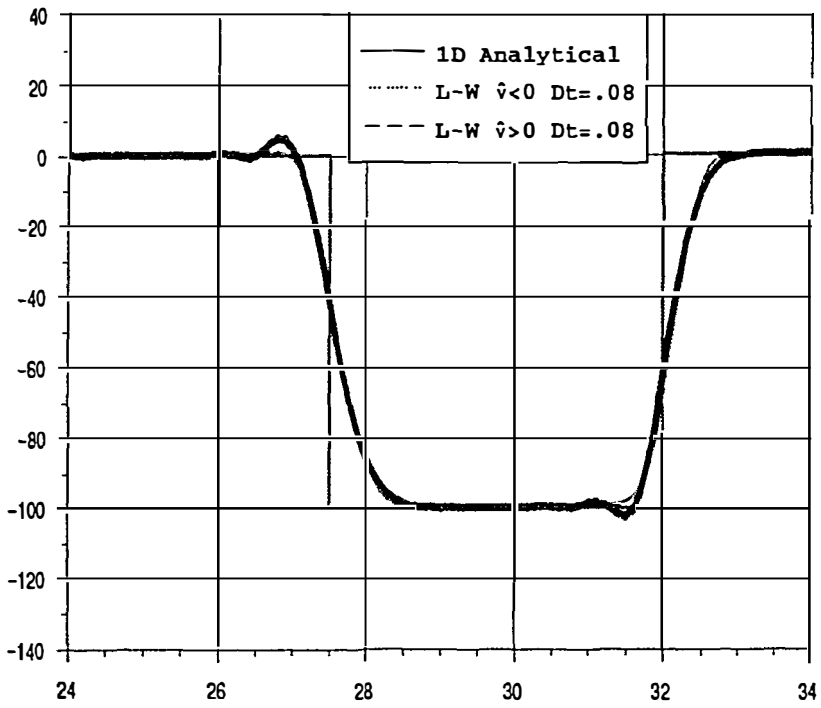
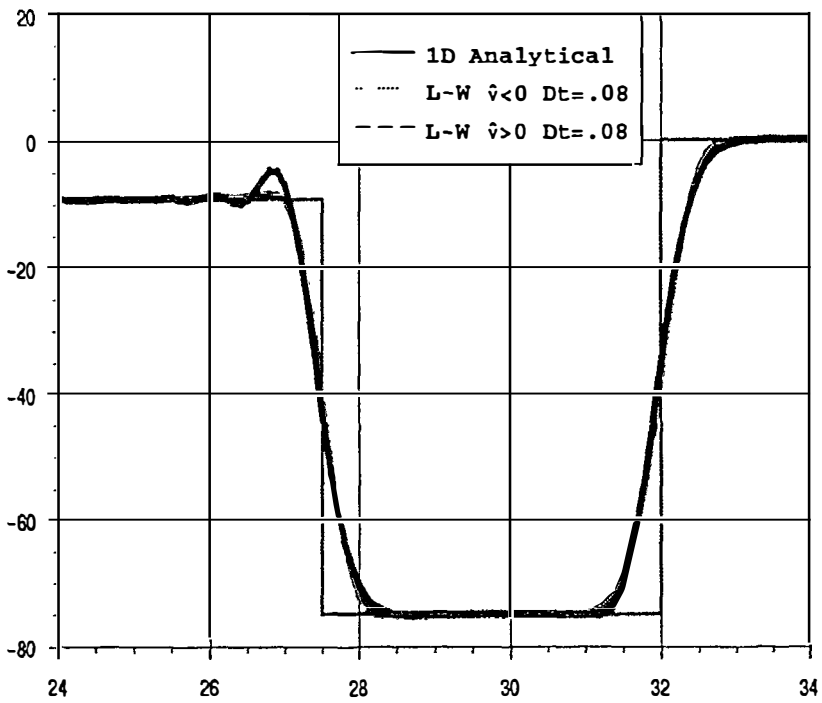


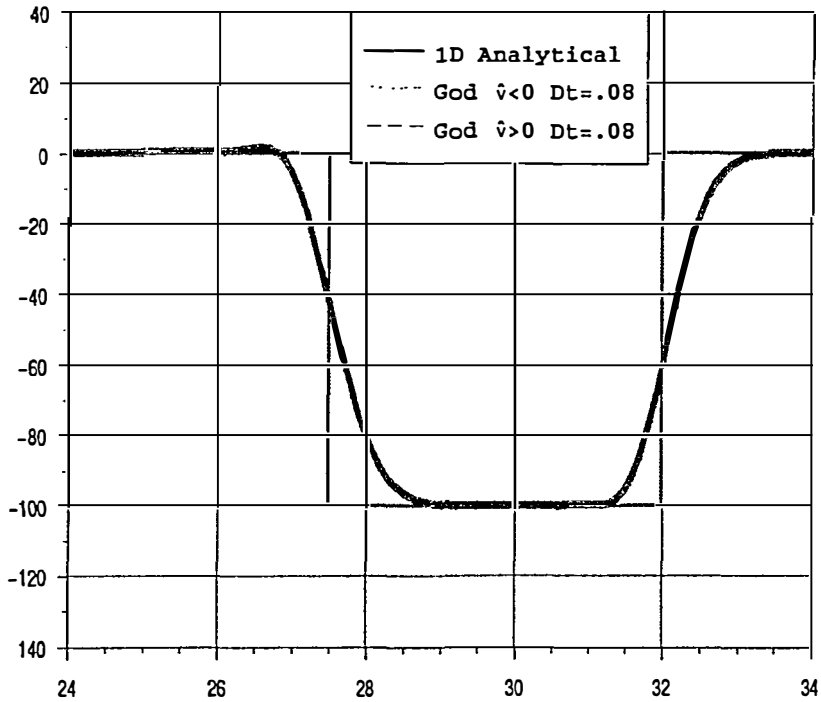
Figure 4 Comparison between the obtained results, (a) CASE A (negative mesh velocity) with elastic material, (b) CASE A with elasto-plastic material, (c) CASE B (positive mesh velocity) with elastic material, (d) CASE B with elasto-plastic material



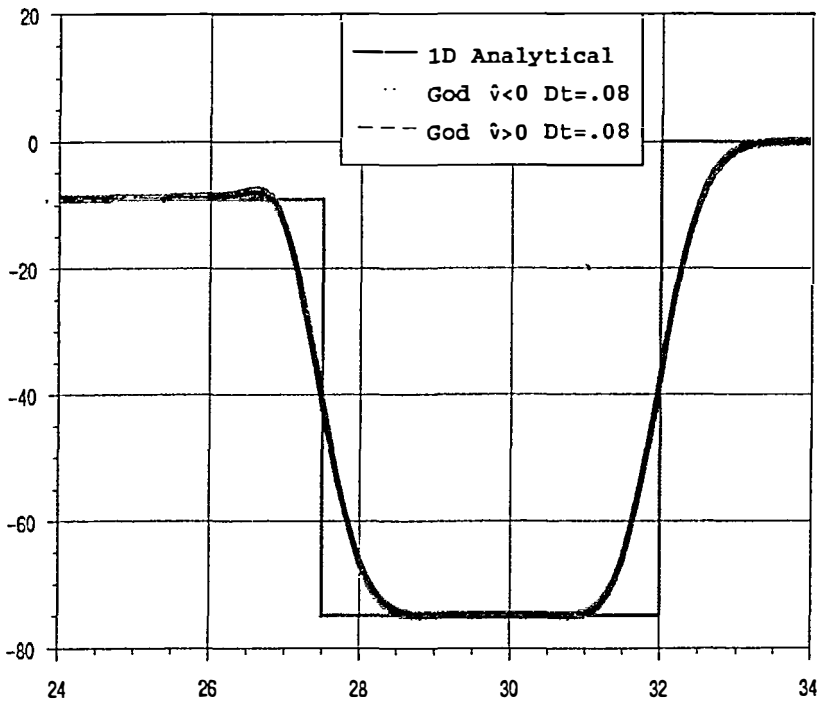
(a)



(b)

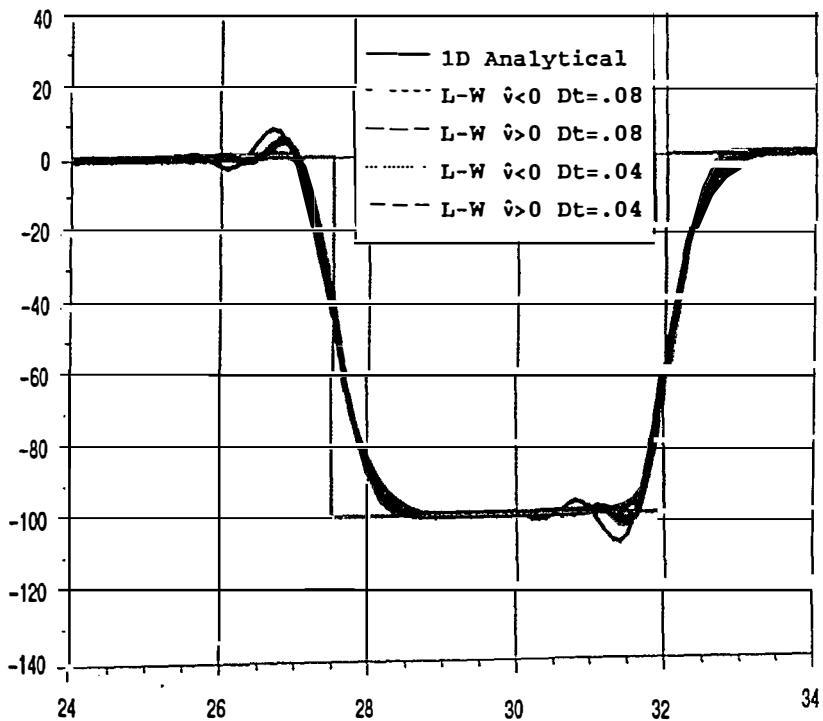


(c)

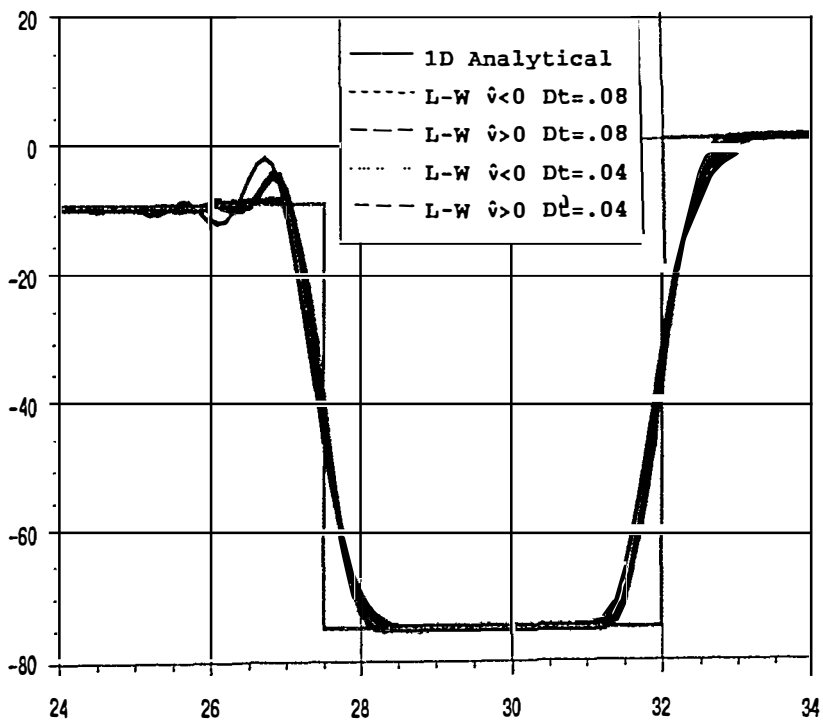


(d)

Figure 5 Influence of the mesh velocity direction on the stress update algorithms. (a) L-W and Elastic material; (b) L-W and Elasto-plastic material; (c) G and Elastic material; (d) G and Elasto-plastic material



(a)



(b)

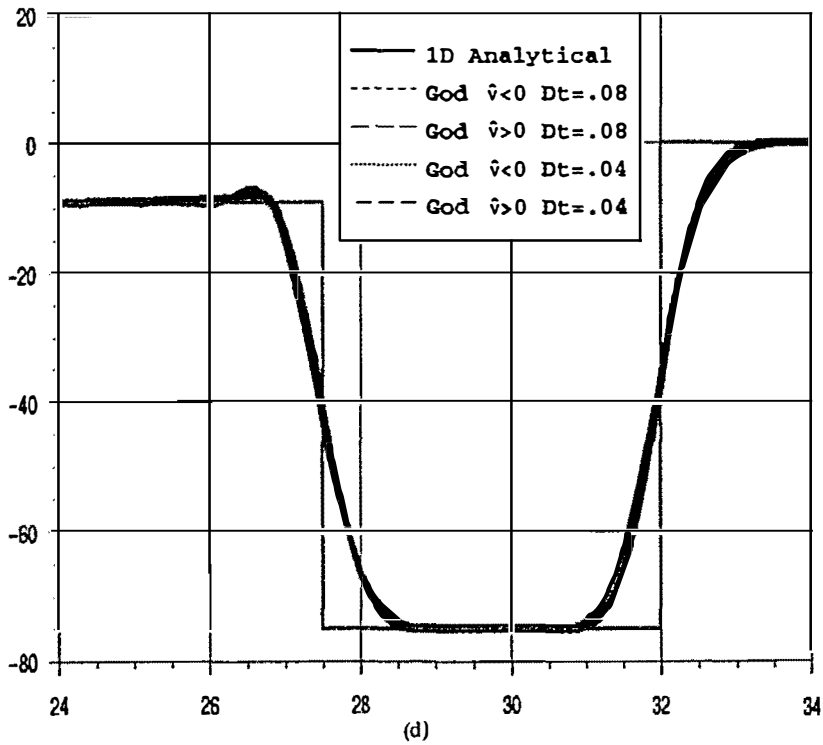
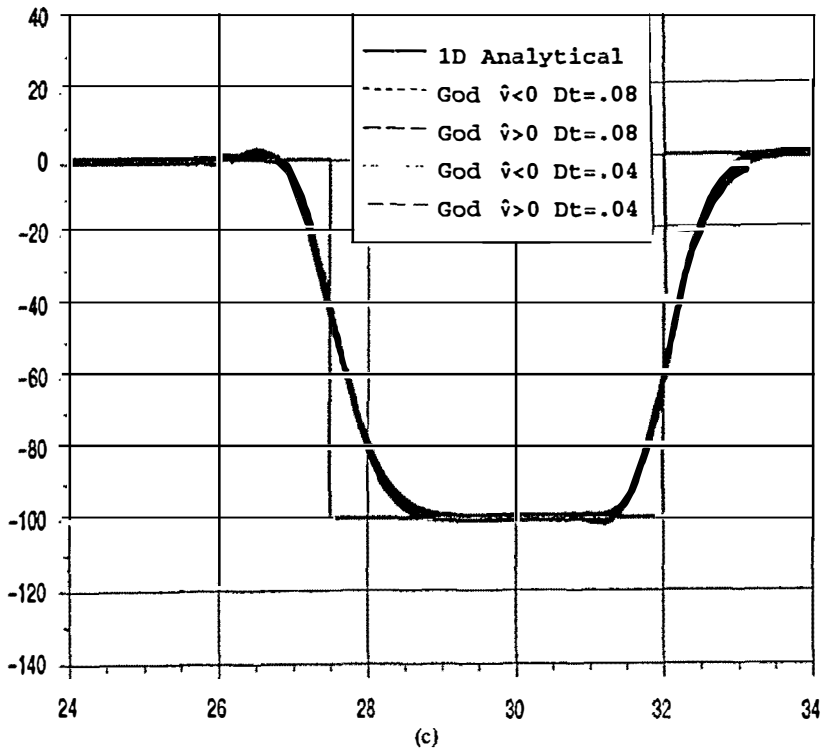


Figure 6 Influence of the time increment. (a) L-W and Elastic material; (b) L-W and Elasto-plastic material; (c) G and Elastic material; (d) G and Elasto-plastic material

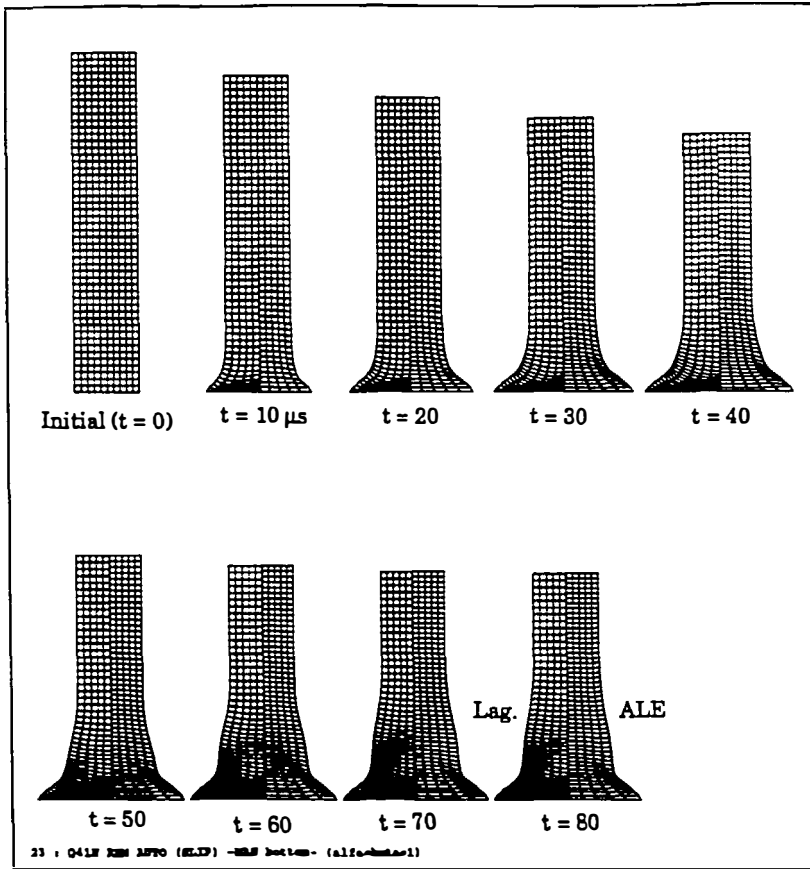


Figure 7 Comparison between the deformed meshes obtained with the Lagrangian description (left) and ALE formulation (right)

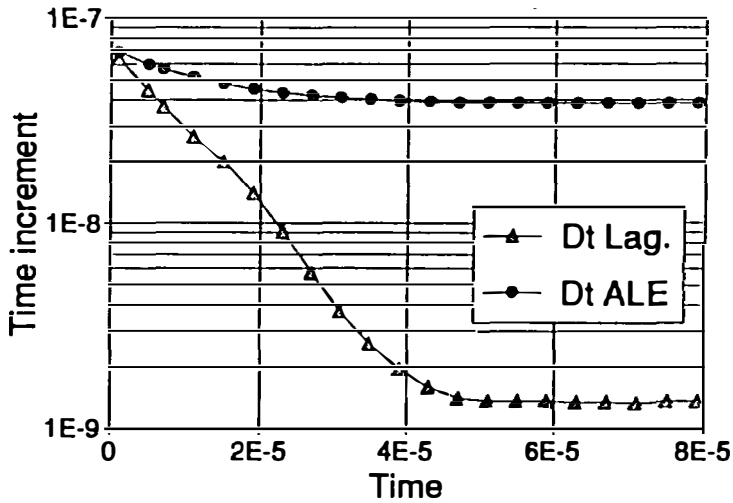
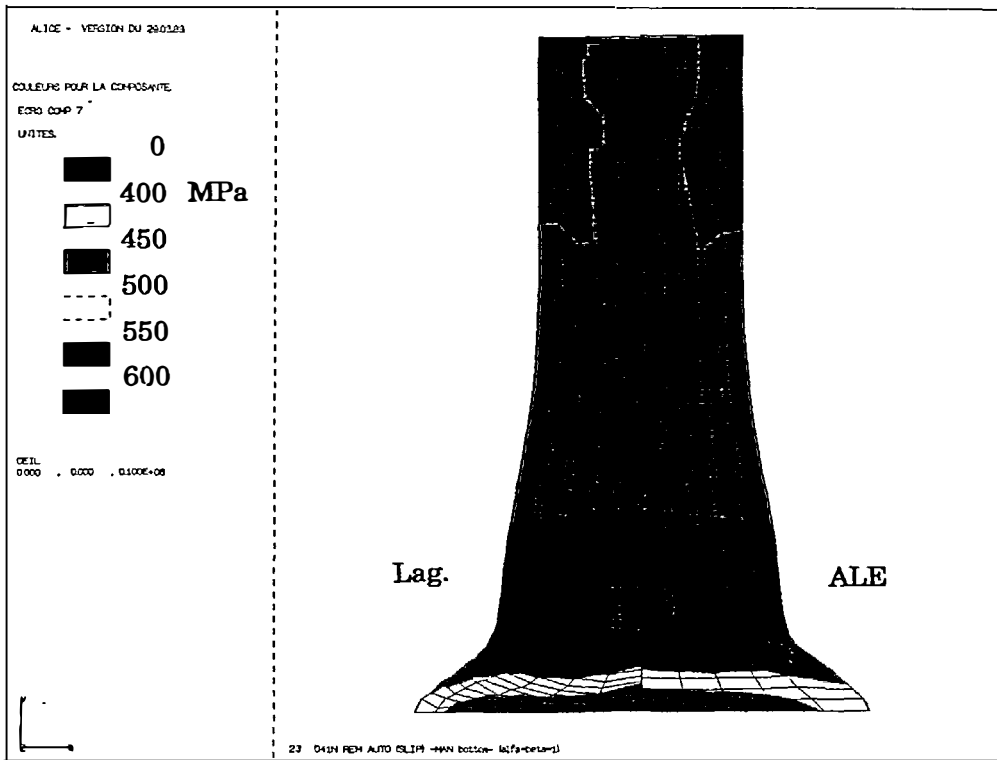
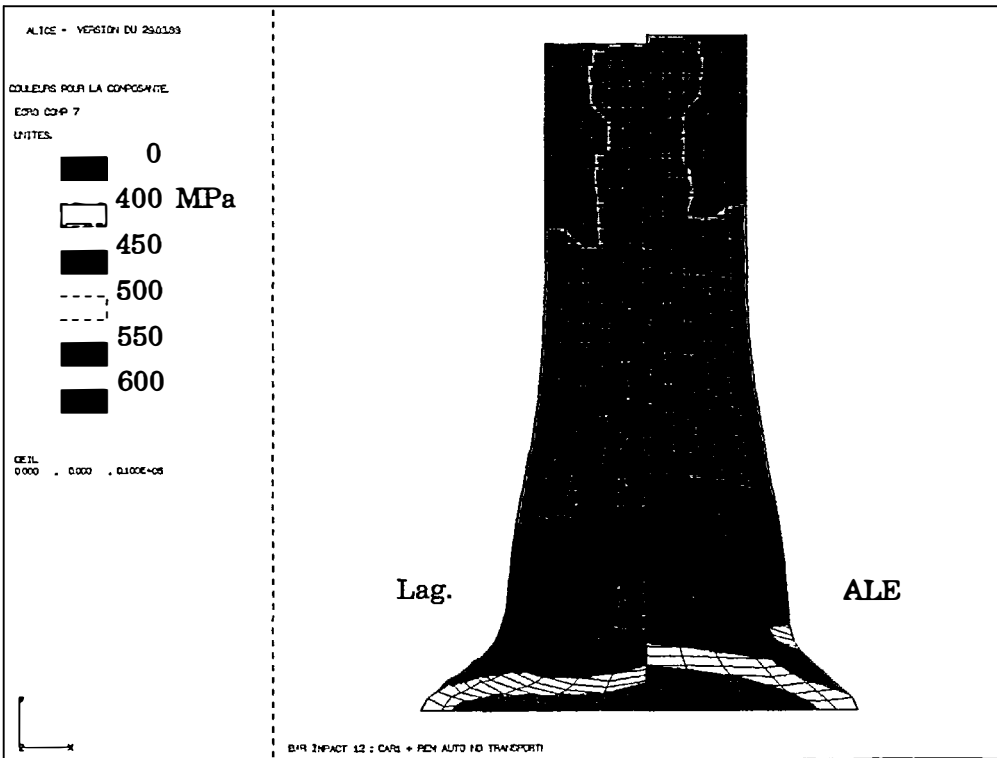


Figure 8 Variation of the time-increment,  $\Delta t$ , with time (during the deformation process)





(a)



(b)

Figure 9 Comparison between final yield stress distributions: (a) Lagrangian (left) and ALE (right); (b) Lagrangian (left) and incorrect computation without stress update during remeshing (right)

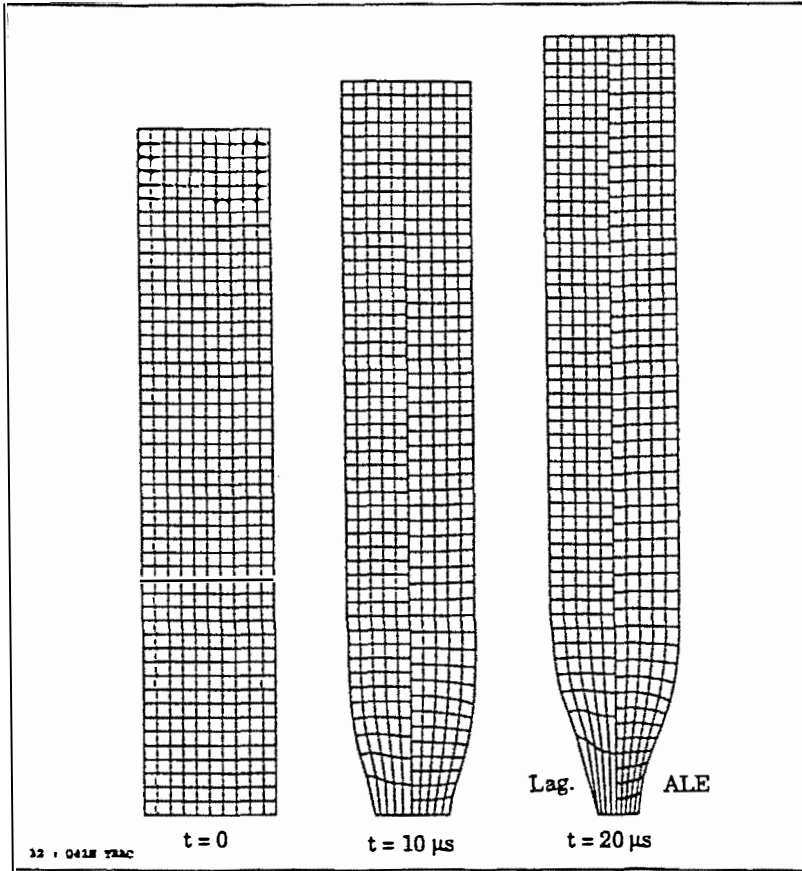


Figure 10 Pulling analysis: comparison of the deformed meshes at various instants, Lagrangian (left) and ALE (right)

different instants and, obviously, the Lagrangian computation shows large inaccuracies due to the mesh distortion. It should be noticed, in this figure, that the free surface nodes are not defined as Lagrangian and that the remeshing allows tangential *sliding* of the mesh nodes along material surfaces even under large boundary motion.

#### Simulation of a forming process

Finally, an engineering problem is presented, a similar example is presented in Reference 21 but in a static analysis. It consists in an elasto-plastic material with  $E = 200$  GPa,  $\nu = 0.30$ ,  $\sigma_y = 250$  MPa,  $\rho = 8930$  kg/m<sup>3</sup>, and a plastic modulus  $E_p = 1$  GPa. The body is deformed by a rigid frictionless tool with a prescribed velocity, only a quarter of the domain (a rectangular region of 3 cm by 1 cm) is studied because two axes of symmetry are supposed, and a plane strain analysis is conducted using  $2 \times 2$  Gauss integration. The analysis is performed up to a 60% reduction in height of the original piece. A schematic statement of this problem is presented in Figure 12.

Note that the frictionless boundary condition is rather difficult to implement in the Lagrangian case. Moreover, different meshes are needed (see, for instance, Reference 21) depending on the

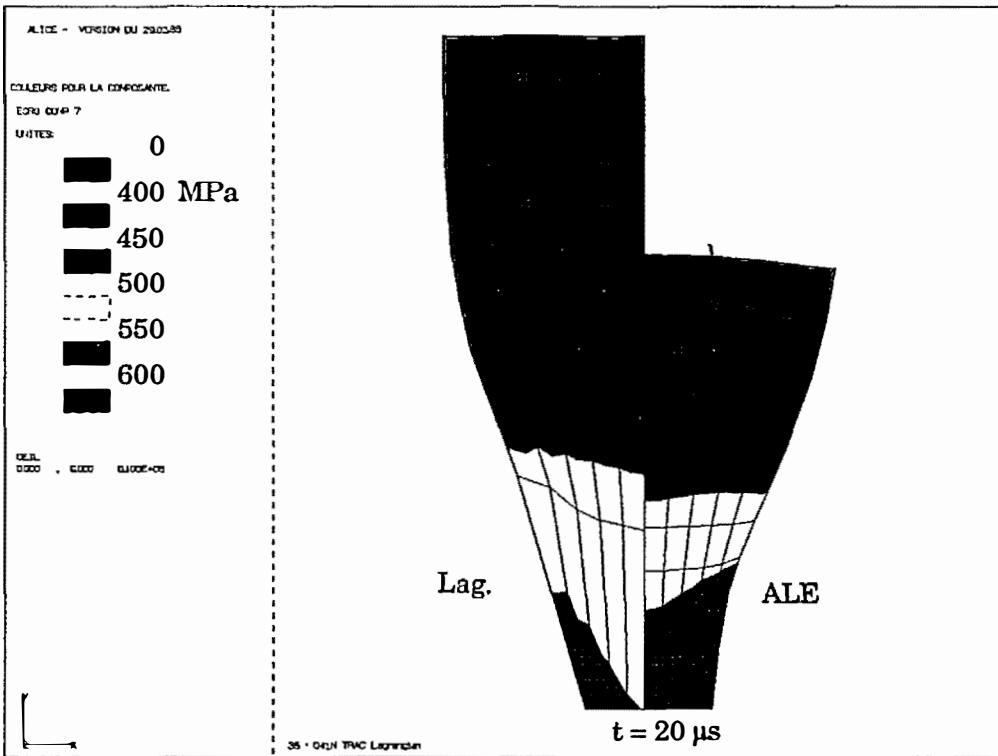
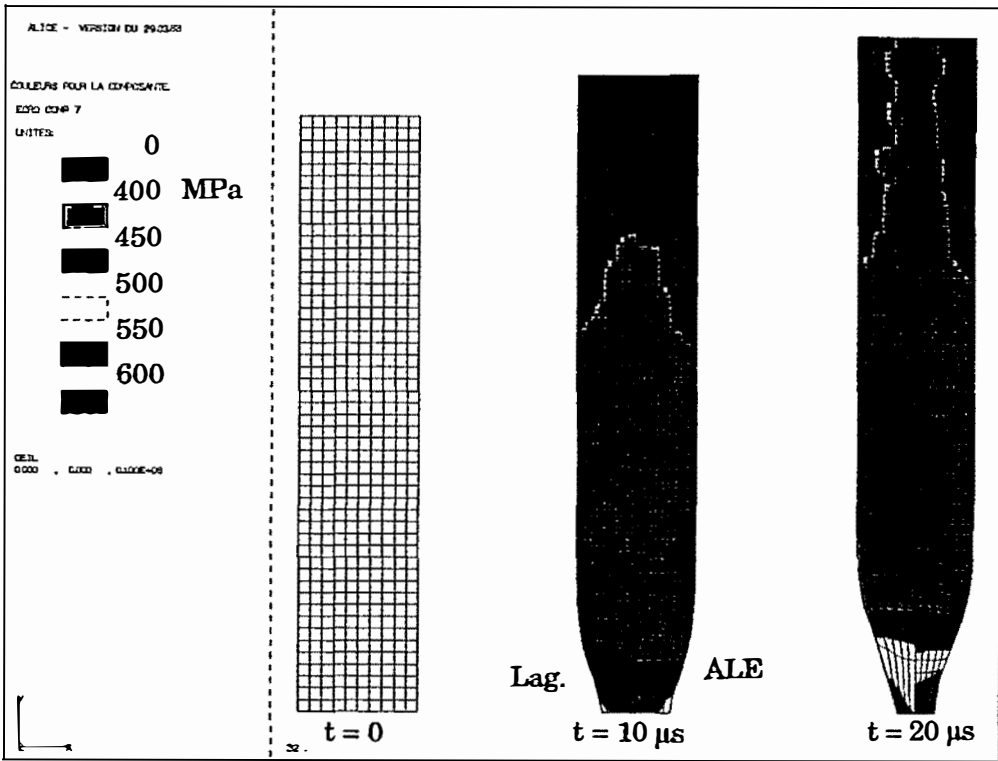


Figure 11 Pulling analysis: comparison between yield stress distributions

## Symmetric Coining

Model: 2-D Plane Strain Analysis

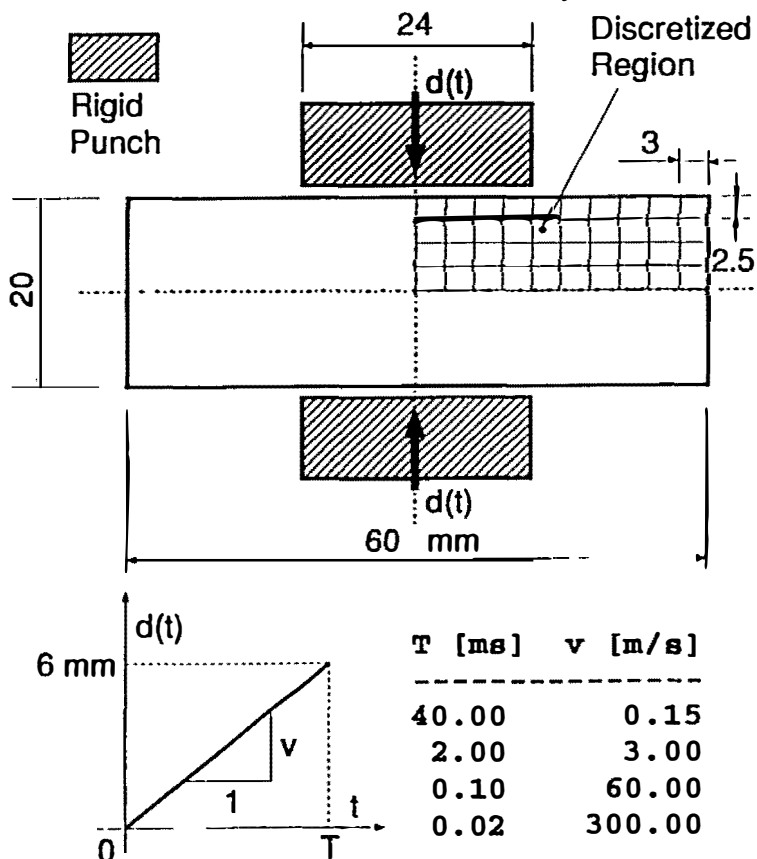


Figure 12 Schematic statement of the symmetric coining problem

reduction required because every reduction induces different deformations and consequently, diverse element distortions. And finally, the Lagrangian description needs an ad hoc mesh, particularly refined under the edge of the tool where most of the deformation occurs.

Figure 13 shows the evolution of the yield stress (equivalent plastic strain) during the deformation process for a punching velocity of 60 m/s. The ALE formulation allows to maintain element regularity and an accurate description of the boundary motion. The evolution of the plastic areas are always led by the right corner of the tool. Notice however that as expected, a plastic band is clearly detected between the previously mentioned corner and the center of the specimen.

To compare the influence of the dynamic effects different punch velocities are studied. Figure 14 shows the comparison between the final deformed shapes. The influence of the dynamic effects is negligible for die velocities under 3 m/s; but, as expected, the fast velocity cases induce a larger back extrusion while the quasi-static simulations allow larger amount of material to flow outward. The external free surface shape is clearly dependent on the dynamic effects.

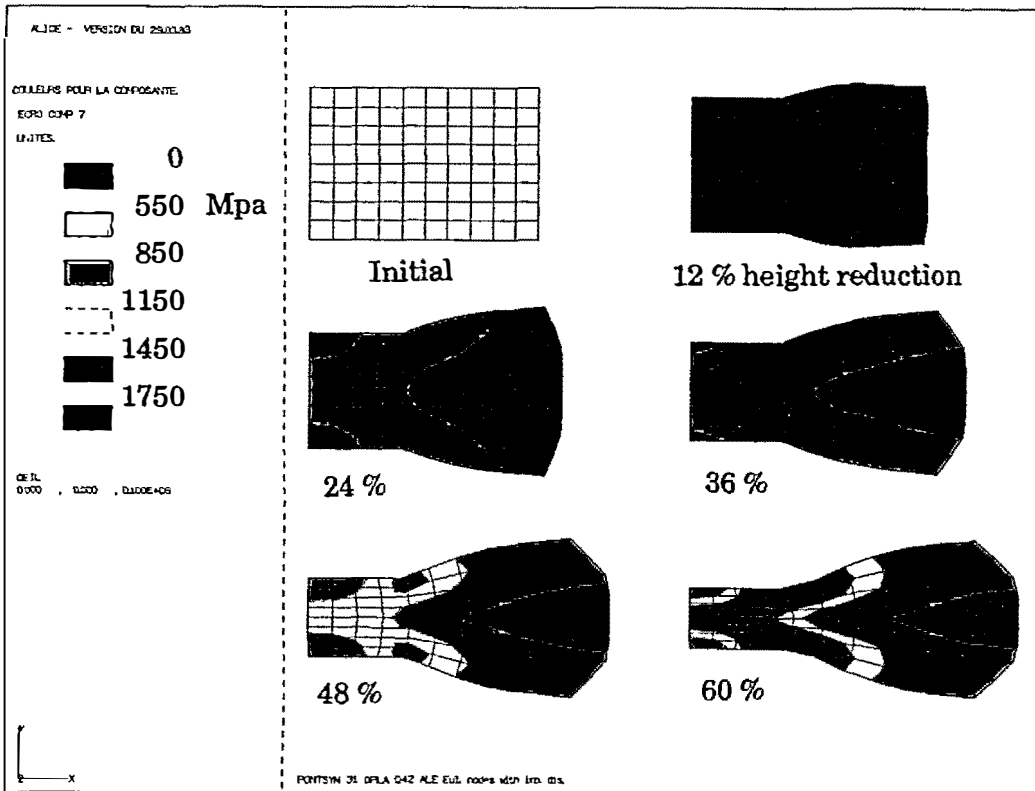


Figure 13 Evolution of the yield stress field during the deformation process, die velocity of 60 m/s

### SUMMARY AND CONCLUSIONS

The general overview and summarized discussion of the ALE formulation presented here indicates the applicability and effectiveness of this technique in generalized continuum mechanics problems. The applications shown are focussed in non-linear solid mechanics because the ALE formulation is already well expounded in the hydrodynamic and fluid-structure interaction fields.

This technique enhances the basic advantages of the finite element method for modelling complex geometries and boundary conditions. It allows a uniform and simple treatment of both confined boundaries and large boundary motion of free surfaces, as well as an excellent flexibility in moving the computational mesh. The result is a very versatile modelling technique which permits accommodation of large continuum distortion and boundary motion, numerical efficiency (from a computational cost point of view), and accurate numerical modelling in particular on material surfaces mapping and interpolation enrichment over determined areas.

After the introduction to the ALE notation and fundamentals, a discussion of the primary governing issues has been presented. Kinematics which are the fundamental concern in ALE, are necessary to develop the governing equations of the continuum problem. Then several specific and inherent concerns of the ALE technique, boundary conditions implementation for large boundary motion, equations of state, and remeshing are discussed. Their adequate

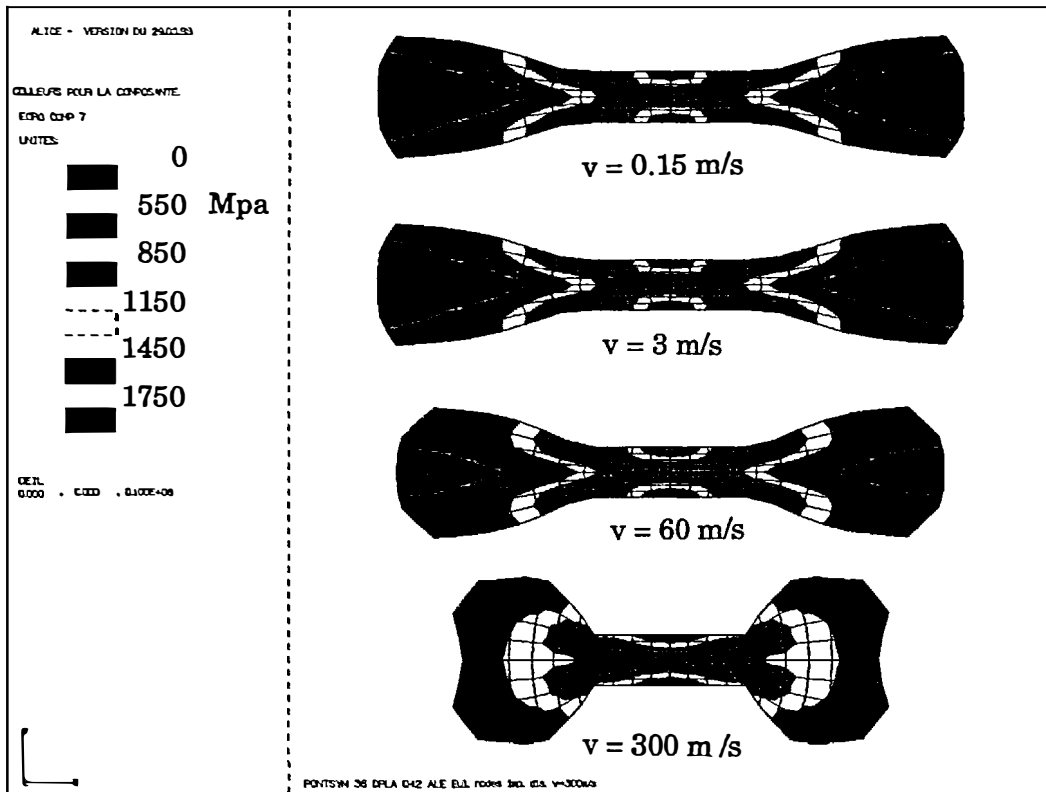


Figure 14 Comparison between the final deformed meshes and yield stress distributions at different punching velocities

implementation is crucial for the applicability and effectiveness of the proposed technique; thus the discussion focusses on their advantages and drawbacks. All of the previous remarks are general in the ALE formulation, while the time integration algorithm employed is especially designed for the implementation of the ALE formulation in a fast-transient dynamics code.

Finally, it is important to notice that the numerical examples shown range from purely academic tests to real engineering simulations. They show the applicability of this formulation to non-linear solid mechanics and in particular to impact, coining or forming analysis.

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