Class Notes: Introduction to the Finite Element Methods

Juan David Gómez Cataño jgomezc1@eafit.edu.co Nicolás Guarín-Zapata nicoguarin@gmail.com

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Contents

Sı	Summary				
1	Intr	roduct	ion	2	
2	Rev	Review of Boundary Value Problems			
	2.1	Bound	dary value problems	7	
		2.1.1	Differential formulation-Generalized balance law	7	
2.2 Brief review of the linearized theory of elasticity model		review of the linearized theory of elasticity model	11		
		2.2.1	Equivalence between strong and weak forms	14	
		2.2.2	Variational formulation	21	
		2.2.3	Weighted residual methods	28	
3	The	e Finit	e Element Method	38	
	3.1	Discre	etization of the PVW using FEM	38	
		3.1.1	Basic elements of interpolation theory	44	

CONTENTS	ii

		3.1.2 Formulation of the finite element matrices	55	
4	Cor	mputational aspects		
	4.1	Global assembly		
	4.2	Sparse assembly		
	4.3	Python implementations		
	4.4	Commercial codes and user subroutines	78	
5	Wa	ave propagation problems in the time domain		
	5.1	Problem formulation		
	5.2	Explicit time integration algorithm		
	5.3	Statement of the Problem		
		5.3.1 Damping Assumptions	82	
		5.3.2 Algorithm corresponding to the damping assumption $3 \ldots$.	83	
		5.3.3 Decoupling	85	
	5.4	The DRM approach	89	
6	Wa	ve propagation problems in the frequency domain	90	
	6.1	Problem formulation		
	6.2	Discretization in complex algebra		
	6.3	Discretization in real algebra		

CONT	ENTS	iii
6.4	Scattering by a micropolar solid	90
6.5	Coupling with the BEM	90
7 Per	riodic media problems	91
7.1	Bloch theorem	92
	7.1.1 Bloch theorem in elastodynamics	94
7.2	Bloch theorem as boundary conditions	95
7.3	Variational form for elastodynamics with Bloch-periodicity	96
7.4	Implementation	98
	7.4.1 Bloch-periodicity in FEM	98
	7.4.2 Real algebra implementation	103
8 No	nlinear Equilibrium in FEM	109
8.1	General Equilibrium Equations	109
	8.1.1 Lagrangian description of the equilibrium equations	110
8.2	Non-linear problems	113
Refere	ences	116
Biblio	Bibliography	

Todo list

Check name of the "course"	3
Review notation.	5
Figure: Make a sketch of the DRM	89
Add Periodic media info	91

Summary

Class Notes for the Course Introduction to the Finite Element Methods

Keywords: Scientific Computing, Computational Mechanics, Finite Element Methods, Numerical Methods.

Chapter 1

Introduction

In an introductory graduate course in the Finite Element Method (FEM), the purpose is to develop a basic understanding of the discrete or numerical strategy of Finite Element-(FE) Algorithms when a closed form solution to a Boundary Value Problem (BVP) is not possible due to complexities existing probably: in the boundary conditions, material behavior or in the kinematic description used in the model. In such a course, in order to master and identify the main mathematical and algorithmic aspects of the method at the beginners level, the studied problem is kept lineal. In this sense 4 key aspects make the core of the introductory course (at least for the case of the linearized theory of elasticity boundary value problem):

- Formulation and identification of the strong form of the Boundary Value Problem (BVP) where the continuum mechanics governing equations are revisited and particularized to the case of linear elastic material behavior. The BVP is completed by identifying the correct prescription of boundary conditions for a well posed mathematical problem.
- Formulation and identification of the weak form of the BVP where the governing equations and natural boundary conditions representing equilibrium at the material point level are replaced by an equivalent but weaker form of equilibrium now valid at the global level. In the case of a Continuum Mechanics problem this so-called weak form can be shown to be equivalent to the principle of virtual power and lends itself for the partition of the problem into sub-domains or finite size elements.

- Introduction of the idea of discretization dividing the problem into subdomains and using interpolation theory within each subdomain-In the context of the FE method this is the subject of shape functions. Once the computational specimen has been divided into subdomains (or a mesh of finite elements) the selected primary variable is approximated within each element via interpolation of the known response at selected predefined points or nodes.
- Computational aspects grouped also into 5 points:
 - Formulation of elemental matrices in the physical space.
 - Formulation of elemental matrices in the natural domain-Isoparametric transformation.
 - Numerical integration: Gauss quadrature.
 - Assembly of system matrices and imposition of boundary conditions.
 - Solution of the discrete equilibrium statement and calculation of elemental results.

These 4 key points are perfectly well documented in numerous nicely written textbooks and it could be argued that is not even worth to register for an introductory course since a moderately dedicated student can accomplish the task via self-study. Unfortunately, linearity is scarce—although useful to grasp the basic understanding of the problem—and the real world is full of non-linear behavior and understanding the needed algorithms can be easily justified. In this Advanced Finite Element Methods course the goal is then to understand the basic aspects of non-linear finite element analysis. Like in the linear case there is also a vast amount of literature for the non-linear problem. However, in the non-linear case the kinematic problem itself may take different routes leading to a wide variety of FEM formulations that will difficult a self-study strategy. Considering the above this brief set of class notes is intended as a guide for self-study and more important represents a help towards the implementation of the algorithm for the consideration of Material and Geometric Non-linearities in Solids.

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The basic reference is Professor Bathe's textbook [1] but we will also follow closely ABAQUS Theory Manual [2]. ABAQUS is a multi-physics oriented commercially available finite element analysis tool. Its strength resides in the effective non-linear algorithms and on the capability of taking user subroutines written in Fortran or in C++. The possibility of implementing user subroutines makes it a very powerful

research tool. In the particular case of a stress/displacement analysis problem with non-linearities these are considered through the kinematics contribution or through the material contribution. In the first case the non-linear behavior must be considered at the element level while in the second it corresponds to the response of a material point which in the context of the FEM algorithm corresponds to an integration point. In ABAQUS those two sources of non-linearity can be independently controlled by the user via user subroutines UEL and UMAT. In both cases the non-linearity is primarily solved by the classical Newton-Raphson scheme and that will be the approached followed herein. Although the notes are mainly written for an advanced course the specific problem of a linear solid usually studied in the introductory course can be derived like a particular case of the most general non-linear algorithm.

The current set of Class Notes is organized as follows. First and since we will be dealing with history dependent non-linear problems the most powerful (at least when it works) solution algorithm, namely the Newton-Raphson iteration is studied. The technique is first illustrated for the simple 1D-case and then generalized into the multi-degree of freedom system. In both cases pseudo-codes will be presented preparing the way for the Finite Element Algorithm. The presentation however is not exhaustive in mathematical terms and the reader is referred to excellent treatments like Burden [3] and Press et al. [4] for the mathematical aspects of the Newton method.

In the next section the briefly introduced Newton-Raphson technique is contextualized to the case of a system of equations representing equilibrium between internal and external forces as typically found in a finite element model. Moreover, the non-linearities come into play through a dependence of the internal forces into displacements. At this stage the details of the formulation of the finite element equations via discretization into nodal variables is not presented but emphasis is laid down into the solution algorithm. Interest is then given to the particular form taken by the Newton-Raphson algorithm into the commercial finite element code ABAQUS. That code can be used as a powerful non-linear equation solver where the coefficient matrix and the excitation can be directly controlled by the user. Moreover the solver can be used into a multi-physics context in terms of generalized forces and fluxes. In the particular case of the stress analysis finite element method the user can control the elemental contribution to the coefficient matrix and the contribution of each material point to obtain that element contribution. This is achieved through the so-called user subroutines UEL for element and UMAT for material. Having introduced the Newton-Raphson method the notes concentrate next on general discretization aspects starting from the physical strong form of the equations in the deformed configuration and passing to an arbitrary weak form in the reference configuration. The resulting algorithm is therefore a Total Lagrangian (TL) method. Once the general equations are introduced a particular work conjugate stress-strain pair is chosen and the discrete equations, including kinematic interpolators, are described.

Notation

In a non-linear algorithm the bookkeeping is involved since we have to simultaneously record 4 different fields as follows:

- The physical fields in terms of tensorial descriptions.
- The time field since the problem is solved incrementally and time may appear as an artificial chronological variable or as a real quantity in a dynamic problem.
- The interpolation field. Since all the involved variables will be interpolated we will need to keep track of the way this interpolation is being performed.
- The iterations field needed in the solution of the non-linear problem.

In order to keep this bookkeeping simple we use the following indicial notation with subscripts and superscripts

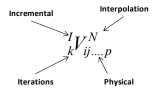


Figure 1.1. General notation to study non-linear finite element problems

Capital superscripts will be reserved for the incremental time description and for the interpolation scheme. For instance an expression like refers to the time instant while refers to . Similarly a variable refers to interpolation over the node. Left

Review notation.

and right subscripts will be used to make reference to the iteration being performed and the order of the tensor variable. For instance refers to a second order tensor corresponding to the iteration.

Chapter 2

Review of Boundary Value Problems

2.1 Boundary value problems

In this section we will define a general initial boundary value problem (I-BVP). In the first part we will introduce the differential formulation given in terms of a set of governing equations and properly specified boundary conditions. The resulting equations are obtained after using a generalized balance law. Following this classical and well known approach we formally re-state these equations in the so-called strong form. Subsequently we re-write and prove an equivalent form of the balance law in the form of an integral representation highly friendly for a numerical solution. Since in the integral description of the problem the order of the derivatives in the field functions decreases by one, the resulting statement is called a weak formulation.

2.1.1 Differential formulation-Generalized balance law

Let ds be a differential surface element; dV a differential volume element; $u(\mathbf{x}, t)$ a scalar (or vector) function of space and time. The flux or rate of flow of the quantity

 $u(\mathbf{x},t)$ through dS at time t is defined like

$$p(\mathbf{x})\nabla u \cdot \hat{n} \, \mathrm{d}S$$
,

where $p(\mathbf{x})$ is a positive function, assumed known and time independent. Similarly, the time rate of change of $u(\mathbf{x}, t)$ in an element dV is given by

$$\rho(\mathbf{x}) \frac{\partial u}{\partial t} \, \mathrm{d}V$$
 ,

where once again $\rho(\mathbf{x})$ is a known, given, time independent positive function. Additional effects occurring in the element dV at the time can be expressed like

$$H(\mathbf{x}, t) dV \equiv -q(\mathbf{x})u(\mathbf{x}, t) + \hat{F}(\mathbf{x}, t)$$

where $\hat{F}(\mathbf{x},t) = \rho(\mathbf{x})F(\mathbf{x},t)$. In the above the term qu represent internal effects due to changes proportional to u while $\hat{F}(\mathbf{x},t)$ are other external influences in the medium.

Balancing the internal and external changes yields

$$\int_{V} \rho(\mathbf{x}) \frac{\partial u}{\partial t} \, dV = \int_{S} p(\mathbf{x}) \nabla u \cdot \hat{n} \, dS + \int_{V} H(\mathbf{x}, t) \, dV ,$$

or equivalently

$$\int\limits_{V} \rho(\mathbf{x}) \frac{\partial u}{\partial t} \, \mathrm{d}V = \int\limits_{S} p(\mathbf{x}) \boldsymbol{\nabla} u \cdot \hat{n} \, \mathrm{d}S - \int\limits_{V} q(\mathbf{x}) u(\mathbf{x},t) \, \mathrm{d}V + \int\limits_{V} \rho(\mathbf{x}) F(\mathbf{x},t) \, \mathrm{d}V \quad .$$

Using the divergence theorem as

$$\int_{S} p(\mathbf{x}) \nabla u \cdot \hat{n} \, dS = \int_{V} \nabla \cdot (p \nabla u) \, dV$$

yields after substitution

$$\int_{V} \left[\rho(\mathbf{x}) \frac{\partial u}{\partial t} - \nabla \cdot (p(\mathbf{x}) \nabla u) + q(\mathbf{x}) u(\mathbf{x}, t) - \rho(\mathbf{x}) F(\mathbf{x}, t) \right] dV = 0$$

Assuming a continuous integrand, the arbitrariness of V implies

$$\rho(\mathbf{x})\frac{\partial u}{\partial t} - \nabla \cdot (p(\mathbf{x})\nabla u) + q(\mathbf{x})u(\mathbf{x}, t) - \rho(\mathbf{x})F(\mathbf{x}, t) = 0$$

Letting

$$\mathcal{L} \equiv -\nabla \cdot p(\mathbf{x})\nabla + q(\mathbf{x})$$

the generalized set of partial differential equations can be written like

$$\rho(\mathbf{x})\frac{\partial u(\mathbf{x},t)}{\partial t} + \mathcal{L}u(\mathbf{x},t) = \rho(\mathbf{x})F(\mathbf{x},t) . \qquad (2.1)$$

And they are categorized as:

• Hyperbolic;

$$\rho(\mathbf{x})\frac{\partial^2 u(\mathbf{x},t)}{\partial t^2} + \mathcal{L}u(\mathbf{x},t) = 0$$

• Parabolic; and

$$\rho(\mathbf{x})\frac{\partial u(\mathbf{x},t)}{\partial t} + \mathcal{L}u(\mathbf{x},t) = 0$$

• Elliptic

$$\mathcal{L}u(\mathbf{x},t) = \rho(\mathbf{x})F(\mathbf{x},t)$$
.

It can be shown that \mathcal{L} satisfies the *symmetry* condition

$$\int_{V} \mathcal{L}(u)v \, dV = \int_{V} \mathcal{L}(v)u \, dV$$

and positive definiteness [5,6]

$$\int_{V} \mathcal{L}(u)u \, dV > 0, \quad \forall u .$$

Strong form

Given $\rho(\mathbf{x})$, $q(\mathbf{x})$, $p(\mathbf{x})$, $F(\mathbf{x},t)$ and \bar{u} find $u(\mathbf{x},t): V \to \mathbb{R}$ such:

$$\rho(\mathbf{x})\frac{\partial u}{\partial t} - \nabla \cdot [p(\mathbf{x})\nabla u] + q(\mathbf{x})u(\mathbf{x},t) - \rho(\mathbf{x})F(\mathbf{x},t) = 0 \quad \forall \mathbf{x} \in V$$

and

$$u = \bar{u} \quad \forall \mathbf{x} \in S_u$$
$$p(\mathbf{x})u_i \hat{n}_i = B(\mathbf{x}, t) \quad \forall \mathbf{x} \in S_t .$$

In the FEM we will look for approximate solutions to u subject to the following conditions:

 $u = \bar{u} \quad \forall \mathbf{x} \in S_u \quad \text{(Essential boundary conditions)}$

and

$$\int\limits_{S} \left(\frac{\partial u}{\partial x_j} \right)^2 \mathrm{d}S < \infty \ ,$$

which corresponds to the functions being square integrable. We will denote this space by $\mathbb H$.

The space of functions satisfying the above two conditions will be denoted by ζ and termed the space of trial functions, formally defined like:

$$\zeta = \{ u \mid u \in \mathbb{H}, u = \bar{u} \quad \forall \mathbf{x} \in S_u \}$$

On the other hand, to validate (or test) the correctness of the approximated or proposed trial functions u it is also necessary to introduce test functions w which are arbitrary except that they satisfy the following conditions:

$$w = 0 \quad \forall \mathbf{x} \in S_n$$

and

$$\int\limits_{S} \left(\frac{\partial w}{\partial x_j} \right)^2 \mathrm{d}S < \infty \ ,$$

which corresponds to the functions being square integrable. The space of functions satisfying the above two conditions will be denoted by \mathcal{L} and termed the space of test functions, formally defined like:

$$\mathcal{L} = \{ w \mid w \in \mathbb{H}, w = 0 \quad \forall \mathbf{x} \in S_u \}$$

Weak form

Given $\rho(\mathbf{x})$, $q(\mathbf{x})$, $p(\mathbf{x})$, $F(\mathbf{x},t)$ and \bar{u} find $u(\mathbf{x},t): V \to \mathbb{R}$ and $\forall w \in \mathcal{L}$ such:

$$\int\limits_{V} p(\mathbf{x})u_{,i}\,w_{,i}\,\mathrm{d}V - \int\limits_{S_t} B(\mathbf{x},t)w\,\mathrm{d}S + \int\limits_{V} q(\mathbf{x})u(\mathbf{x},t)w\,\mathrm{d}V + \int\limits_{V} \rho(\mathbf{x})\frac{\partial u}{\partial t}w\,\mathrm{d}V - \int\limits_{V} \rho(\mathbf{x})F(\mathbf{x},t)w\,\mathrm{d}V = 0$$
 and

$$u = \bar{u} \quad \forall \mathbf{x} \in S_u .$$

Equivalence between the strong and weak forms

$$-\int_{V} [p(\mathbf{x})u_{,i}]_{,i} w \, dV + \int_{S_{t}} [p(\mathbf{x})u_{,i}] \hat{n}_{i} w \, dS - \int_{S_{t}} B(\mathbf{x}, t) w \, dS$$
$$+ \int_{V} q(\mathbf{x})u(\mathbf{x}, t) w \, dV + \int_{V} \rho(\vec{x}) \frac{\partial u}{\partial t} w \, dV - \int_{V} \rho(\mathbf{x}) F(\mathbf{x}, t) w \, dV = 0 \quad (2.2)$$

Grouping together common terms yields

$$\int_{V} \left\{ \rho(\mathbf{x}) \frac{\partial u}{\partial t} - [p(\mathbf{x})u_{,i}]_{,i} + q(\mathbf{x})u(\mathbf{x},t) - \rho(\mathbf{x})F(\mathbf{x},t) \right\} w \, dV + \int_{S_{t}} \left\{ [p(\mathbf{x})u_{,i}] \hat{n}_{i} - B(\mathbf{x},t) \right\} w \, dS = 0$$

from which

$$\rho(\mathbf{x})\frac{\partial u}{\partial t} - [p(\mathbf{x})u_{,i}]_{,i} + q(\mathbf{x})u(\mathbf{x},t) - \rho(\mathbf{x})F(\mathbf{x},t) = 0$$

and

$$p(\mathbf{x})u_{,i}\hat{n}_i = B(\mathbf{x},t) \quad \forall \mathbf{x} \in S_t .$$

2.2 Brief review of the linearized theory of elasticity model

Here we present a brief description of the boundary value problem governing the response of an elastic body. For a full discussion of the model and its mathematical

aspects the reader is referred to [7].

The governing equations (in terms of stresses) stem from the principle of conservation of linear momentum and conservation of moment of linear momentum. The former leads to a set of 3 partial differential equations in the components of the stress tensor while the latter leads to the symmetries in the stress tensor.

$$\sigma_{ij,j} + f_i = \rho \ddot{u}_i \quad \forall \mathbf{x} \in V, t \in \mathbb{R}^+$$

$$\sigma_{ij} = \sigma_{ji}.$$
 (2.3)

In eq. (2.3) σ_{ij} is the stress tensor; f_i is the vector of body forces; and u_i is the displacements vector.

Denoting the tractions vector associated with a surface with normal direction \hat{n}_j by $t_i^{\hat{n}}$ we have the complete BVP as follows:

$$t_i^{\hat{n}} = \sigma_{ii} \hat{n}_i \quad \forall \in \mathbf{x} \in S. \tag{2.4}$$

Equation (2.3) correspond to 6 equations with 12 unknowns (the 9 components of the stress tensor and the 3 components of the displacements vector) and the system is undetermined. In order to have a solvable BVP we must introduce kinematic strain-displacement relations and a stress-strain law. In the case of infinitesimal theory of elasticity the strain-displacement relation is given by:

$$\varepsilon_{ij} = \frac{1}{2}(u_{i,j} + u_{j,i}) \tag{2.5}$$

where the term ϵ_{ij} is the symmetric component of the displacements gradient tensor. The components of the strain tensor describe the distortions and changes in magnitude (volumetric changes) of the material point in the continuum model. Now the simplest stress-strain (constitutive) relationship is given by Hooke's law¹

¹Despite the name of *law* used, this relation is not always valid, but is a good approximation for small strains.

$$\sigma_{ij} = 2\mu\varepsilon_{ij} + \lambda\varepsilon_{kk}\delta_{ij} . {2.6}$$

where μ and λ are material constants. The problem involves now a total of 18 equations and 18 unknowns can be solved if subjected to properly specified boundary conditions.

Displacement formulation

Substituting eq. (2.5) in eq. (2.6) and the result in eq. (2.3) yields after some manipulation:

$$(\lambda + \mu)u_{i,ij} + \mu u_{i,jj} + f_i = \rho \ddot{u}_i \quad \forall \mathbf{x} \in V, \ t \in \mathbb{R}^+. \tag{2.7}$$

Since eq. (2.7) (simultaneously describing equilibrium, kinematic relations and constitutive response) is a second order equation governing the displacement field possible boundary conditions are in terms of the variable itself or its first oder derivatives. For a well-possed problem the following is a set of valid boundary conditions:

$$t_i^{\hat{n}} = \sigma_{ij}\hat{n}_{ij} \quad \forall \mathbf{x} \in S_t u_i = \bar{u}_i \quad \forall \mathbf{x} \in S_u$$
 (2.8)

and where $S_t \cup S_u = S$ and $S_t \cap S_u = \emptyset$.

In the particular case in which u_i is not a function of time, we obtain the static version of the BVP, i.e,

$$(\lambda + \mu) u_{j,ij} + \mu u_{i,jj} + f_i = 0 \quad \forall \mathbf{x} \in V$$

$$t_i^{\hat{n}} = \sigma_{ij} \hat{n}_{ij} \quad \forall \mathbf{x} \in S_t$$

$$u_i = \bar{u}_i \quad \forall \mathbf{x} \in S_u$$

$$(2.9)$$

Notice that the tractions BC

$$t_i^{\hat{n}} = \mu(u_{i,j} + u_{j,i})\hat{n}_j + \lambda u_{k,k}\delta_{ij}\hat{n}_j ,$$

actually involves first order displacements derivative and as such it is a Neumann boundary condition on u_i .

2.2.1 Equivalence between strong and weak forms

Strong form

The strong form corresponds to the differential formulation of the problem, it is denoted by $\{S\}$ and it reads:

Given f_i , $t_i^{\hat{n}}$ and \bar{u}_i find $u_i: V \to \mathbb{R}$ such:

$$(\lambda + \mu)u_{j,ij} + \mu u_{i,jj} + f_i = 0 \quad \forall \mathbf{x} \in V$$

$$t_i^{\hat{n}} = \sigma_{ij}\hat{n}_{ij} \quad \forall \mathbf{x} \in S_t$$

$$u_i = \bar{u}_i \quad \forall \mathbf{x} \in S_u$$

$$(2.10)$$

In eq. (2.10) the boundary conditions specified by the traction vector $t_i^{\hat{n}}$ correspond to the natural boundary conditions, while those specified in terms of the displacements vector \bar{u}_i represent the essential boundary conditions.

- We are interested in developing methods to obtain approximate solutions to $\{S\}$.
- The FEM is formulated starting from a statement equivalent to $\{S\}$ in which we use trial functions until certain prescribed conditions are met.
- We will look for solutions u_i subject to the following conditions:

$$u_i = \bar{u}_i \quad \text{in} \quad S_u$$

$$\int_S \left(\frac{\partial u_i}{\partial x_j}\right)^2 dS < \infty$$

The first condition corresponds to the satisfaction of the essential boundary condition, while the second corresponds to the functions being square integrable. The

space of functions satisfying the above two conditions is denoted by ς and formally defined like

$$\varsigma = \{ u_i \mid u_i \in H, u_i = \bar{u}_i \in S_u \} .$$

On the other hand, in order to validate the introduced trial functions we also need testing functions w_i also called in the FEM literature weighting or distribution functions. These functions are arbitrary apart from having to satisfy the following conditions:

$$w_i = 0 \quad in \quad S_u$$

$$\int_{S} \left(\frac{\partial w_i}{\partial x_j}\right)^2 dS < \infty$$

In what follows we formally denote the space of these functions by V and define it like

$$V = \{ w_i | w_i \in H, u_i = w_i = 0 \in S_u \} .$$

Weak form

Here we will show that the equilibrium statement represented in the differential formulation can be described in alternative forms. In such description the continuity requirement for the trial functions is weaker than in the strong form leading to the term "weak" statement. Here this alternative representation will be denoted like $\{W\}$ and it reads;

Given f_i , $t_i^{\hat{n}}$ and \bar{u}_i find $u_i: V \to \mathbb{R}$ and $\forall w_i \in V$ such:

$$\int\limits_{V} \sigma_{ij} w_{i,j} \, dV - \int\limits_{V} f_i w_i \, dV - \int\limits_{S_t} t_i^{\hat{n}} w_i \, dS = 0$$

Proof 1:

Let $u_i \in \varsigma$ be a solution to $\{S\}$ and let $w_i \in V$. Forming the inner product of the equilibrium statement given in eq. (2.3) with w_i and forcing the integral over the

domain to be zero we have

$$\int\limits_{V} (\sigma_{ij,j} + f_i) w_i \, dV = 0 \ ,$$

expanding the terms in the integrand and integrating by parts the first term on the left we have

$$\int_{V} \sigma_{ij,j} w_i \, dV + \int_{V} f_i w_i \, dV = 0 .$$

$$- \int_{V} w_{i,j} \sigma_{ij} \, dV + \int_{S} \sigma_{ij} \hat{n}_j w_i \, dS + \int_{V} w_i f_i \, dV = 0$$

since $w_i \in V$ it follows that $w_i = 0$ in S_u from which

$$\int_{V} \sigma_{ij} w_{i,j} \, dV - \int_{V} f_i w_i \, dV - \int_{S_t} t_i^{\hat{n}} w_i \, dS = 0$$
 (2.11)

Now, considering that u_i is solution of the strong form $\{S\}$ it must satisfy $u_i = \bar{u}_i \in S_u$ and as a result $u_i \in \varsigma$. On the other hand, since u_i satisfies eq. (2.11) $\forall w_i \in V$ we have that u_i satisfies the definition of weak solution specified in $\{W\}$.

Proof 2:

Let u_i be a solution of $\{W\}$ and thus $u_i \in \varsigma$ which means that

$$u_i = \bar{u}_i \in S_u$$

and that it satisfies

$$\int_{V} \sigma_{ij} w_{i,j} dV - \int_{V} f_i w_i dV - \int_{S_t} t_i^n w_i dS = 0 ,$$

integrating by parts,

$$-\int_{V} \sigma_{ij,j} w_i dV + \int_{S} \sigma_{ij} n_j w_i dS - \int_{V} f_i w_i dV - \int_{S_t} t_i^n w_i dS = 0$$

Since $w_i \in V$ we have that $w_i = 0$ in S_u and therefore

$$\int_{V} w_i(\sigma_{ij,j} + f_i)dV + \int_{S_t} w_i(\sigma_{ij}n_j - t_i^n)dS = 0$$

from which

$$\sigma_{ij,j} + f_i = 0 \quad \mathbf{x} \in V$$

$$t_i^n = \sigma_{ij} n_j \quad \forall \mathbf{x} \in S_t$$

$$u_i = \bar{u}_i \quad \forall \mathbf{x} \in S_u$$

$$(2.12)$$

which is once again the strong form of the problem given in eq. (2.3).

Simple wedge under self-equilibrated loads

Consider the double wedge of side ℓ and internal angle 2ϕ shown in fig. 2.1. It is assumed to be contained in the X-Y plane, with loading conditions satisfying a plane strain (or plane stress) idealization. The material is elastic with Lame constants λ and μ . The wedge is loaded by uniform tractions of intensity S applied over its four faces in such a way that the wedge is self-equilibrated. We wish to find the closed-form elasticity solution for the stress, strain and displacement fields throughout the problem domain.

Under plane strain conditions the general 3D stress equilibrium equations (see eq. (2.3)) reduce to:

$$\frac{\partial \sigma_{xx}}{\partial x} + \frac{\partial \tau_{xy}}{\partial y} = 0$$

$$\frac{\partial \tau_{xy}}{\partial x} + \frac{\partial \sigma_{yy}}{\partial y} = 0$$
(2.13)

while the kinematic relation (eq. (2.5)) reads

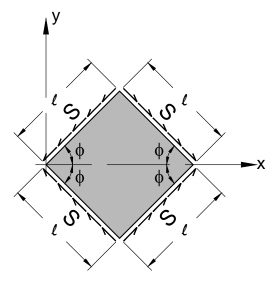


Figure 2.1. 2D Self-equilibrated wedge.

$$\epsilon_{xx} = \frac{\partial u}{\partial x}$$

$$\epsilon_{yy} = \frac{\partial v}{\partial y}$$

$$\gamma_{xy} = \frac{\partial u}{\partial y} + \frac{\partial v}{\partial x}$$
(2.14)

where u and v are the horizontal and vertical displacements respectively.

Stress field

The stress field can be obtained by simple inspection from the traction boundary conditions prescribed over the inclined surfaces yielding;

$$\sum F_x = 0 \longrightarrow -\ell S \cos(\phi) + \sigma_{xx} \ell \sin(\phi) = 0$$
$$\sum F_y = 0 \longrightarrow -\ell S \sin(\phi) - \sigma_{yy} \ell \cos(\phi) = 0$$

and the following stress solution:

$$\sigma_{xx} = S \cot(\phi)$$

$$\sigma_{yy} = -S \tan(\phi)$$

$$\tau_{xy} = 0.$$
(2.15)

In eq. (2.15) the condition $\tau_{xy} = 0$ is due to the symmetries in the problem.

Traction boundary conditions

Let us verify that the above stress solution satisfies the traction BC using the expression:

$$t_i^{\hat{n}} = \sigma_{ij} \hat{n}_{ij}.$$

Denoting the outward normals to the inclined surfaces of the wedge by \hat{n}^1 , \hat{n}^2 , \hat{n}^3 , \hat{n}^4 these are given by;

$$\hat{n}^{1} = -\sin(\phi)\hat{e}_{x} + \cos(\phi)\hat{e}_{y}$$

$$\hat{n}^{2} = -\sin(\phi)\hat{e}_{x} - \cos(\phi)\hat{e}_{y}$$

$$\hat{n}^{3} = +\sin(\phi)\hat{e}_{x} + \cos(\phi)\hat{e}_{y}$$

$$\hat{n}^{4} = +\sin(\phi)\hat{e}_{x} - \cos(\phi)\hat{e}_{y}$$

where \hat{e}_x and \hat{e}_y are the reference unit vectors. Now, the components of the traction vector follow directly like

$$t_i = \sigma_{ij} \hat{n}_j$$

then over the face with normal \hat{n}^1 we have

$$t_x = -S\cos(\phi)$$

$$t_y = -S\sin(\phi)$$

similarly, over the face with normal \hat{n}^2

$$t_x = -S\cos(\phi)$$
$$t_y = +S\sin(\phi)$$

over the face with normal \hat{n}^3

$$t_x = +S\cos(\phi)$$
$$t_y = -S\sin(\phi)$$

and finally, over the face with normal \hat{n}^4 ;

$$t_x = +S\cos(\phi)$$

$$t_y = +S\sin(\phi) .$$

Strain field

The strain field can be obtained after using the stress solution found in eq. (2.15) together with the constitutive law given by eq. (2.6) which for a plane strain idealization takes the form:

$$\epsilon_{xx} = \frac{1}{E} (\sigma_{xx} - \nu \sigma_{yy})$$

$$\epsilon_{yy} = \frac{1}{E} (\sigma_{yy} - \nu \sigma_{xx})$$

$$\gamma_{xy} = \frac{\tau_{xy}}{\mu}$$
(2.16)

which for the particular case yields;

$$\epsilon_{xx} = +\frac{S}{E} \left[\cot(\phi) + \nu \tan(\phi) \right] = +\frac{S}{E} K_1(\nu, \phi)$$

$$\epsilon_{yy} = -\frac{S}{E} \left[\tan(\phi) + \nu \cot(\phi) \right] = -\frac{S}{E} K_2(\nu, \phi)$$

$$\gamma_{xy} = 0.$$
(2.17)

Displacement field

The displacement field is obtained after direct integration of the strains after using the fact that:

$$du_i = \epsilon_{ij} dx_j + \omega_{ij} dx_j$$

and the condition $\omega_{xy} = 0$ also due to symmetries , as follows:

$$u = +\frac{S}{E}K_1(\nu, \phi)x + A$$
$$v = -\frac{S}{E}K_2(\nu, \phi)y + B$$

and where A and B are integration constants.

From the condition u=0 at $x=\ell\cos(\phi)$ we have that $A=-\frac{S}{E}K_1(\nu,\phi)\ell\cos(\phi)$ then it follows that

$$u = \frac{S}{E}K_1(\nu, \phi)(x - \ell \cos(\phi)).$$

Similarly, from the condition v = 0 at y = 0 we have that B = 0 from which

$$v = -\frac{S}{E}K_2(\nu, \phi)y$$

2.2.2 Variational formulation

In this section we formulate the boundary value problem using the approach of the calculus of variations in which the governing PDEs and boundary conditions are obtained after finding the minimum (or maximum) of a functional according to a variational principle².

²According to Wikipedia [8]

We will see that the weak form (and therefore also the strong form) can be obtained alternatively through the process of finding extreme values for a functional. We will illustrate this idea for the general case of theory of elasticity and then we will present particular examples.

Some vague definitions in the calculus of variations

In variational calculus a **functional** can be understood as a "function" having as independent variables or arguments a space of vector functions and producing as a result (or dependent variable) a scalar. For instance, in the particular case of the theory of elasticity such a "function" corresponds to the total potential energy functional Π given by;

$$\Pi(u_i) = \frac{1}{2} \int_V \sigma_{ij} \varepsilon_{ij} \, dV - \int_V f_i \, u_i \, dV - \int_S t_i^{(n)} u_i \, dS$$
 (2.18)

and where the first term in the left hand side corresponds to the internal strain energy, while the last two terms are the work done by the external body and traction forces. The above functional has as independent variables the displacement vector and its spatial derivatives. This is indicated by the presence of the displacement vector u_i in the expression $\Pi(u_i)$.

In variational calculus we are interested in finding a function u_i that renders the functional Π a maximum or a minimum. In loose terms, the analogous to the differential operator in calculus of functions is now termed the variational operator δ (i.e., δ is analogous to $\frac{\partial}{\partial x_i}$). As such $\delta\Pi$ acts over the function u_i and its derivatives

A variational principle is a scientific principle used within the calculus of variations, which develops general methods for finding functions which minimize or maximize the value of quantities that depend upon those functions. For example, to answer this question: "What is the shape of a chain suspended at both ends?" we can use the variational principle that the shape must minimize the gravitational potential energy.

According to Cornelius Lanczos, any physical law which can be expressed as a variational principle describes an expression which is self-adjoint. These expressions are also called Hermitian. Such an expression describes an invariant under a Hermitian transformation.

as follows

$$\delta\Pi = \frac{\partial\Pi}{\partial u_i} \delta u_i + \frac{\partial\Pi}{\partial \left(\frac{\partial u_i}{\partial x_j}\right)} \delta \left(\frac{\partial u_i}{\partial x_j}\right) + \dots + \frac{\partial\Pi}{\partial \left(\frac{\partial^n u_i}{\partial x_j \cdots \partial x_k}\right)} \delta \left(\frac{\partial^n u_i}{\partial x_j \cdots \partial x_k}\right) .$$

The following rules apply to the variational operator δ :

• For functionals Π and Φ it follows that

$$\delta(\Pi + \Phi) = \delta\Pi + \delta\Phi$$

• For functionals Π and Φ it follows that

$$\delta(\Pi\Phi) = \delta\Pi\,\Phi + \Pi\,\delta\Phi$$

• For a functional Π and an integer n it follows that

$$\delta(\Pi^n) = n(\Pi^{n-1}) \, \delta\Pi$$

• For a functional Π it follows that

$$\delta \int \Pi \, \mathrm{d}x = \int \delta \Pi \, \mathrm{d}x$$

If the variational operator is applied to the functional $\Pi(u_i)$ it produces functions or variations in u_i which are arbitrary and such $\delta u_i \in V$ and $\delta u_i = 0$ in S_u .

To find an extreme function in the calculus of variations we proceed like in differential calculus. Here we compute the first variation of the functional $\delta\Pi$ and solve the variational equation

$$\delta\Pi = 0 \tag{2.19}$$

in the unknown function u_i .

In the particular case of the total potential energy functional Π this yields

$$\int_{V} \sigma_{ij} \, \delta \varepsilon_{ij} \, dV - \int_{V} f_i var u_i \, dV - \int_{S_t} t_i^{(n)} \, \delta u_i \, dS = 0$$
 (2.20)

where we recognize the weak form of the BVP stated previously. It becomes evident that the functions δu_i in eq. (2.20) play the role of the test functions w_i introduced in the weak form. On the other hand, since we have already shown that the weak and strong forms are equivalent we conclude that having the functional and the essential boundary conditions is equivalent to having the strong form of the problem.

Principle of minimum potential energy

In the theory of elasticity the total potential energy Π is the result of adding the elastic strain energy which is stored in the body upon deformation and the potential energy (work) imparted to the body by the applied forces. The principle states that the body is in equilibrium when this total potential energy reaches a minimum. This is equivalent to stating that an equilibrium configuration is attained when an infinitesimal variation from the position of minimum potential energy involves null changes in energy. This implies the variational condition:

$$\delta\Pi = 0. (2.21)$$

The above principle leads to the so-called principle of virtual displacements stated as follows³: "The equilibrium of the body requires that for any compatible small virtual displacements satisfying the condition of being zero at S_u , imposed on the body in its state of equilibrium, the total internal virtual work is equal to the total external virtual work"

$$\int_{V} \sigma_{ij} \, \delta \varepsilon_{ij} \, dV - \int_{V} f_i \delta u_i \, dV - \int_{S_i} t_i^{(n)} \, \delta u_i \, dS = 0$$

where δu_i are the virtual displacements and $\delta \varepsilon_{ij}$ are the corresponding virtual strains.

Comparing the virtual work principle with the weak formulation given in eq. (2.11) we identify δu_i with the test functions w_i . As such the PVW takes the form of a powerful tool to test if a body is in equilibrium for a given solution (represented by the trial functions). In what follows we illustrate the use of the principle through some examples corresponding to problems in Bathe's textbook.

 $^{^3 \}mathrm{see}$ Bathe pp156

Problem⁴

Establish the differential equation of equilibrium of the problem shown and the boundary conditions. Determine whether the differential operator of the problem is symmetric and positive definite and prove your answer. [1]

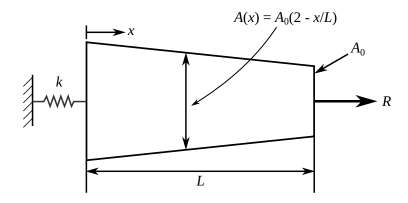


Figure 2.2. Rod with varying cross-sectional area. The Young's modulus is E.

$$\Pi = \frac{1}{2} \int_0^L \sigma_{xx} \varepsilon_{xx} A(x) dx + \frac{1}{2} k u_0^2 - R u_L$$
$$= \frac{1}{2} \int_0^L EA(x) \left(\frac{du}{dx}\right)^2 dx + \frac{1}{2} k u_0^2 - R u_L .$$

The first variation for this functional is

$$\delta \Pi = \int_{0}^{L} EA(x) \frac{\mathrm{d}u}{\mathrm{d}x} \frac{\mathrm{d}\delta u}{\mathrm{d}x} \,\mathrm{d}x + ku_0 \,\delta u_0 - R \,\delta u_L .$$

⁴3.15 from Finite Element Procedures

If we integrate by parts, we obtain

$$\delta\Pi = -\int_{0}^{L} \frac{\mathrm{d}}{\mathrm{d}x} \left[EA(x) \frac{\mathrm{d}u}{\mathrm{d}x} \right] \delta u \, \mathrm{d}x + EA(x) \frac{\mathrm{d}u}{\mathrm{d}x} \, \delta u \bigg|_{0}^{L} + ku_{0} \, \delta u_{0} - R \, \delta u_{L}$$

from which

$$\frac{\mathrm{d}}{\mathrm{d}x} \left[EA(x) \frac{\mathrm{d}u}{\mathrm{d}x} \right] = 0$$

$$EA(x) \frac{\mathrm{d}u}{\mathrm{d}x} \Big|_{x=0} = ku_0$$

$$EA(x) \frac{\mathrm{d}u}{\mathrm{d}x} \Big|_{x=L} = R$$

Problem: The Hu-Washizu Variational Principle⁵

Consider the Hu-Washizu functional:

$$\Pi^* = \Pi - \int_{V} \lambda_{ij}^{\varepsilon} (\varepsilon_{ij} - L_{ijk} u_k) \, dV - \int_{S_u} \lambda_i^u (u_i^{S_u} - \bar{u}_i) \, dS$$
 (2.22)

where

- Π: is the potential energy functional.
- L_{ijk} is a differential operator such $\varepsilon_{ij} = L_{ijk}u_k$.
- \bullet S_u surface where essential boundary conditions are prescribed.
- $\lambda_{ij}^{\varepsilon}$ and λ_i^u are Lagrange multipliers.

Using the condition $\delta\Pi=0$ derive for the interior of the body the equilibrium equations

$$\sigma_{ij,j} + f_i = 0$$

⁵4.35 from Finite Element Procedures

the strain-displacement relationship

$$\varepsilon_{ij} = L_{ijk} u_k$$

and the constitutive equation

$$\sigma_{ij} = C_{ijkl} \varepsilon_{kl}$$

and at the surface of the body the relation between the stress tensor and the applied tractions vector at S_t

$$t_i = \sigma_{ij} n_j$$

the relation between the stress tensor and the unknown tractions vector (or reactions) at S_u

$$t_i = \tilde{\sigma_{ij}} n_j$$

and the essential boundary condition at S_u

$$u_i = \tilde{u}_i$$

We want to determine the Euler equations resulting from the condition $\delta \Pi^* = 0$. Applying the variational operator we have

$$\delta\Pi^* = \delta\Pi - \int_{V} \delta\lambda_{ij}^{\varepsilon} \left(\varepsilon_{ij} - L_{ijk}u_k\right) dV - \int_{V} \lambda_{ij}^{\varepsilon} \left(\delta\varepsilon_{ij} - L_{ijk}\delta u_k\right) dV$$

$$- \int_{V} \delta\lambda_{i}^{u} \left(u_i^{S_u} - \bar{u}_i\right) dS - \int_{V} \lambda_{i}^{u} \delta u_i^{S_u} dS$$
(2.23)

and

$$\delta\Pi^* = \int_{V} C_{ijkl} \varepsilon_{kl} \, \delta\varepsilon_{ij} \, dV - \int_{S_t} t_i \, \delta u_i \, dS - \int_{V} f_i \, \delta u_i \, dV - \int_{V} \lambda_{ij}^{\varepsilon} \, \delta\varepsilon_{ij} \, dV + \int_{V} \lambda_{ij}^{\varepsilon} L_{ijk} \, \delta u_k \, dV$$
$$- \int_{V} \delta\lambda_{ij}^{\varepsilon} (\varepsilon_{ij} - L_{ijk} u_k) \, dV - \int_{S} \delta\lambda_{i}^{u} (u_i^{S_u} - \bar{u}_i) \, dS - \int_{S_u} \lambda_{i}^{u} \, \delta u_i \, dS = 0$$

$$(2.24)$$

using

$$\int_{V} (\lambda_{ij}^{\varepsilon} \, \delta u_i)_{,j} \, dV = \int_{V} \lambda_{ij}^{\varepsilon} \, \delta u_{i,j} \, dV + \int_{V} \lambda_{ij,j}^{\varepsilon} \, \delta u_i \, dV$$

In the above we can write

$$\int_{V} \lambda_{ij}^{\varepsilon} L_{ijk} \, \delta u_k \, dV = \int_{V} (\lambda_{ij}^{\varepsilon} \, \delta u_i)_{,j} \, dV - \int_{V} \lambda_{ij,j}^{\varepsilon} \delta u_i \, dV$$

$$= \int_{S_t} \lambda_{ij}^{\varepsilon} \, \delta u_i \, \hat{n}_j \, dS - \int_{V} \lambda_{ij,j}^{\varepsilon} \, \delta u_i \, dV$$

therefore

$$\begin{split} \delta\Pi^* &= \int\limits_{V} \left(C_{ijkl} \varepsilon_{kl} - \lambda_{ij}^{\varepsilon} \right) \delta\varepsilon_{ij} \, \mathrm{d}V - \int\limits_{S_t} t_i \, \delta u_i \, \mathrm{d}S - \int\limits_{V} f_i \, \delta u_i \, \mathrm{d}V + \int\limits_{S_t} \lambda_{ij}^{\varepsilon} \, \delta u_i \, \hat{n}_j \, \mathrm{d}S - \int\limits_{V} \lambda_{ij,j}^{\varepsilon} \delta u_i \, \mathrm{d}V \\ &- \int\limits_{V} \delta\lambda_{ij}^{\varepsilon} (\varepsilon_{ij} - L_{ijk} u_k) \, \mathrm{d}V - \int\limits_{S_u} \delta\lambda_i^u \, (u_i^{S_u} - \bar{u}_i) \, \mathrm{d}S - \int\limits_{S_u} \lambda_i^u \, \delta u_i \, \mathrm{d}S = 0 \\ &= \int\limits_{V} \left(C_{ijkl} \varepsilon_{kl} - \lambda_{ij}^{\varepsilon} \right) \delta\varepsilon_{ij} \, \mathrm{d}V + \int\limits_{S_t} \left(\lambda_{ij}^{\varepsilon} \hat{n}_j - t_i \right) \delta u_i \, \mathrm{d}S \\ &- \int\limits_{V} \left(\lambda_{ij,j}^{\varepsilon} + f_i \right) \delta u_i \, \mathrm{d}V - \int\limits_{V} \left(\varepsilon_{ij} - L_{ijk} u_k \right) \delta\lambda_{ij}^{\varepsilon} \, \mathrm{d}V - \int\limits_{S_u} \left(u_i^{S_u} - \bar{u}_i \right) \delta\lambda_i^u \, \mathrm{d}S - \int\limits_{S_u} \lambda_i^u \, \delta u_i^{S_u} \, \mathrm{d}S = 0 \end{split}$$

Now, imposing the conditions $\delta \varepsilon_{ij} \neq 0$, $\delta \lambda_{ij}^{\varepsilon} \neq 0$, $varu_i \neq 0$ in S_t , $\delta u_i \neq 0$ in V and $\delta \lambda_i^u \neq 0$ in S_u we have

$$\lambda_{ii}^{\varepsilon} = C_{iikl} \varepsilon_{kl} \tag{2.25}$$

$$\varepsilon_{ii} = L_{iik} u_k \tag{2.26}$$

$$t_i = \lambda_{ij}^{\varepsilon} \hat{n}_j \tag{2.27}$$

$$\lambda_{ij,j}^{\varepsilon} + f_i = 0 \tag{2.28}$$

$$u_i^{S_u} = \bar{u}_i \quad . \tag{2.29}$$

2.2.3 Weighted residual methods

This section introduces the concept of residual or difference from zero in a differential equation once its solution is approximated. For that purpose we will take as

prototype equation the one obtained as our general model of BVP (see eq. (2.1)) and recalled here for completeness

$$\rho(\mathbf{x})\frac{\partial u(\mathbf{x},t)}{\partial t} + \mathcal{L}u(\mathbf{x},t) = \rho(\mathbf{x})F(\mathbf{x},t) . \qquad (2.30)$$

We will assume that the actual solution to the generalized BVP given by eq. (2.30) is approximated by $\tilde{u}(\mathbf{x})$ through a superposition like

$$\tilde{u}(\mathbf{x}) = N^I(\mathbf{x})u^I \tag{2.31}$$

where $N^{I}(\mathbf{x})$ are interpolating functions and I denotes a superposition index varying like I = 1, 2, ..., K with K being the number of points where the solution is known. In what follows we will use $u(\mathbf{x})$ instead of $\tilde{u}(\mathbf{x})$ but will keep in mind that we are actually using the approximation given by eq. (2.31). Similarly, in order to keep the discussion simple for the time being we will drop the time effects reducing the generalized PDE to the simple form:

$$\mathcal{L}u(\mathbf{x}) = \rho(\mathbf{x})F(\mathbf{x}) . \tag{2.32}$$

Now, since we are using the approximation given by eq. (2.31) this equation is not strictly satisfied but instead we will have the following "unbalanced" condition

$$\mathcal{L}u(\mathbf{x}) - \rho(\mathbf{x})F(\mathbf{x}) \equiv R \neq 0$$

where the term R corresponds to a residual error which is to be distributed throughout the solution domain. The so-called weighted residual methods differ in the form in which they distribute the residual between the different K points conforming the computational domain.

Using eq. (2.31) in eq. (2.30) and the linearity in the differential operator yields

$$R = \mathcal{L}(N^P)u^P - \rho F.$$

We can see that the residual R is a function defined over the domain of interest. The residual would be exactly zero for the solution of the differential equation, but it will not be zero in general. Thus, we want to make the function R as close to zero as possible. To make R as small as possible we need a function (a functional) where

we can compare different approximation functions. After getting this functional we can minimize its value. For this minimization we could use the norm of the function, another option is to compute a weighted *average* of the function over the domain. This is what we call a weighted residual

$$\Pi[u, w] = \int_{V} wR(u) \, dV ,$$

and we want to minimize it by making

$$\delta\Pi\left[u,w\right] = 0\,,$$

In what follows we will consider different strategies to distribute or weight the residual R over the computational domain.

Galerkin method

In the Galerkin scheme the interpolation functions are used also as weighting functions leading to:

$$\int\limits_V N^Q R \, \mathrm{d}V = 0$$

or explicitly

$$\int_{V} N^{Q} \mathcal{L}(N^{P}) \, dV \, u^{P} = \int_{V} N^{Q} \rho F \, dV . \qquad (2.33)$$

Imposing eq. (2.33) in the K points conforming the computational domain or equivalently ranging Q from 1 to K leads to the following system of algebraic equation

$$K^{QP}U^P = f^Q (2.34)$$

where U^P is a vector that stores the point values of the function u along te K points of the computational domain, while f^Q stores the corresponding point excitations.

Least squares method

In this method the integral of the square of the residual is minimized with respect to the K point parameters or nodal values of the function. Accordingly,

$$\frac{\partial}{\partial u^I} \int_V R^2 \, dV = 0$$
$$\int_V R \frac{\partial R}{\partial u^I} \, dV = 0,$$

The least squares method is a special case of the weighted residual method for the weight functions

$$w^I = \frac{\partial R}{\partial u^I} \,.$$

Expanding the residual, and considering the operator \mathcal{L} as linear, we obtain

$$\frac{\partial}{\partial u^I} \int_{V} [\mathcal{L}(N^P u^P) - \rho F]^2 \, dV = 0$$

$$\int_{V} [\mathcal{L}N^P u^P - \rho F] \mathcal{L}(N^I) \, dV = 0$$

$$\int_{V} \mathcal{L}(N^I) \mathcal{L}(N^P) \, dV \, u^P - \rho \int_{V} \mathcal{L}(N^I) F \, dV = 0$$

which can be written like

$$K^{IP}U^P = f^I (2.35)$$

Collocation method

In the collocation method the coefficients of the approximation are determined by forcing the residual to be exactly zero at K points over the computational domain, i.e.,

$$\mathcal{L}(N^I)u^I - \rho F = 0 \,,$$

or

$$\mathcal{L}[N^I(x^J)]u^I - \rho F(x^J) = 0,$$

where J ranges between 1 and K. This equation can be rewritten as a weightedresidual if we consider the residual to be $\delta(x-x^I)$, the Dirac delta function over the selected points

The resulting system of algebraic equation can be written as

$$K^{IP}U^P = f^I. (2.36)$$

Subdomain method

The zero value of the residual is imposed upon K subdomains

$$\int_{V^I} \mathcal{L}(N^P) \, dV^I \, u^P - \rho \int_{V^I} F \, dV^I = 0 \qquad I = 1, \cdots, K.$$

For instance, for the N-th element it follows that

$$\int_{V^N} \mathcal{L}(N^P) dV^N u^P - \rho \int_{V^N} F dV^N = 0 \qquad P = 1, \dots, K.$$

Applying the equation over the K subdomains leads to the discrete system;

$$K^{IP}U^P = f^I I = 1, \cdots, K.$$
 (2.37)

Ritz method

It operates directly upon the variational statement of the problem. For a given functional

$$\Pi = \Pi(N^Q u^Q)$$

the variational equation reads

$$\delta\Pi \equiv \frac{\partial\Pi}{\partial u^Q} \,\delta u^Q = 0$$

from which

$$\frac{\partial \Pi}{\partial u^Q} = 0.$$

Problem: Discretization of the generalized parabolic equation.

Let us consider the case of the generalized parabolic equation and its discretization following the Galerkin method

$$\rho(\mathbf{x})\frac{\partial u(\mathbf{x},t)}{\partial t} + \mathcal{L}u(\mathbf{x},t) = 0$$

which can also be written using indicial notation

$$\frac{\partial}{\partial x_i} \left[p(x) \frac{\partial u}{\partial x_i} \right] + q(x)u + \rho \frac{\partial u}{\partial t} = \rho F.$$

Assuming that p(x) = 1 yields

$$-\int\limits_{V}N^{P}N_{,ii}^{Q}\,\mathrm{d}V\,u^{Q}+\int\limits_{V}qN^{P}N^{Q}dVu^{Q}+\rho\int\limits_{V}N^{P}N^{Q}dVv^{Q}-\rho\int\limits_{V}N^{P}FdV=0$$

$$\int\limits_{V}N_{,i}^{P}N_{,i}^{Q}dVu^{Q}-\int\limits_{S}N^{P}N_{,i}^{Q}\hat{n}_{i}dSu^{Q}+\int\limits_{V}qN^{P}N^{Q}dVu^{Q}+\rho\int\limits_{V}N^{P}N^{Q}dVv^{Q}-\rho\int\limits_{V}N^{P}FdV=0$$

$$\int\limits_{V} \left(N_{,i}^{P} N_{,i}^{Q} + q N^{P} N^{Q}\right) dV u^{Q} + \rho \int\limits_{V} N^{P} N^{Q} dV V^{Q} = \int\limits_{S} N^{P} N_{,i}^{Q} \hat{n}_{i} dS u^{Q} + \rho \int\limits_{V} N^{P} F dV$$

which can be written in discrete form;

$$K^{PQ}U^Q + C^{PQ}V^Q = f^p$$

Problem: Discretization of Navier equations.

In this case the differential equations are written as

$$(\lambda + \mu)u_{j,ij} + \mu u_{i,jj} + f_i = 0.$$

We can write the differential operator as

$$L_{ij} \equiv (\lambda + \mu) \frac{\partial^2}{\partial x_i \partial x_j} + \mu \frac{\partial^2}{\partial x_k \partial x_k} \delta_{ij}$$

$$\begin{split} r_i &= -f_i \\ u_i &= N_i^Q u^Q \\ \mathcal{L}_{ij}(u_j) &\equiv (\lambda + \mu)(N_j^Q u^Q)_{,ij} + \mu(N_j^Q u^Q)_{,kk} \delta_{ij} \\ \mathcal{L}_{ij}(u_j) &\equiv (\lambda + \mu)N_{j,ij}^Q u^Q + \mu N_{i,kk}^Q u^Q \\ \mathcal{L}_{ij}(u_j) &\equiv \mathcal{L}_{ij}(N_j^Q) u^Q \end{split} .$$

In the Galerkin scheme we use the trial function as weighting function.

$$R_i \equiv L_{ij}(N_i^Q)u^Q + f_i$$

and we state ...

$$\int_{V} N_i^P R_i dV = 0 \qquad P = 1, 2, ..., N.$$

$$\int_{V} N_i^P \mathcal{L}_{ij}(N_j^K) dV u^K + \int_{V} N_i^P f_i dV = 0$$

$$(\lambda + \mu) \int_{V} N_i^P N_{j,ij}^K dV u^K + \mu \int_{V} N_i^P N_{i,kk}^K dV u^K + \int_{V} N_i^P f_i dV = 0$$

integrating by parts;

$$\begin{split} -(\lambda + \mu) \int\limits_{V} N_{i,j}^{P} N_{j,i}^{K} dV u^{K} + (\lambda + \mu) \int\limits_{S} N_{i}^{P} N_{j,i}^{K} \hat{n}_{j} dS u^{K} - \mu \int\limits_{V} N_{i,k}^{P} N_{i,k}^{K} dV u^{K} \\ + \mu \int\limits_{S} N_{i}^{P} N_{i,k}^{K} \hat{n}_{k} dS u^{K} + \int\limits_{V} N_{i}^{P} f_{i} dV &= 0 \end{split}$$

which can be written like;

$$K^{PQ}U^Q = F^P$$

where

$$K^{PQ} = (\lambda + \mu) \int_{V} N_{i,j}^{P} N_{j,i}^{Q} dV + \mu \int_{V} N_{i,k}^{P} N_{i,k}^{Q} dV$$

$$F^{P} = \int_{S} N_{i}^{P} t_{i}^{(\hat{n})} dS + \int_{V} N_{i}^{P} f_{i} dV = 0$$

Problem: Discretization of the acoustic wave equation.

$$\vec{\nabla} \cdot \left[\frac{1}{\rho} \vec{\nabla} p(\mathbf{x}) \right] - \frac{\partial}{\partial t} \left(\frac{1}{\lambda} \frac{\partial \rho}{\partial t} \right) - q(\mathbf{x}) = 0$$

where we recognize;

$$\mathcal{L}() \equiv \vec{\nabla} \cdot \left(\frac{1}{\rho} \vec{\nabla}\right) - \frac{\partial}{\partial t} \left(\frac{1}{\lambda} \frac{\partial}{\partial t}\right)$$

Let

$$p(x) = N^K p^K$$

then

$$\mathcal{L}(p) \equiv \vec{\nabla} \cdot \left(\frac{1}{\rho} \vec{\nabla} N^K p^K \right) - \frac{\partial}{\partial t} \left(\frac{1}{\lambda} \frac{\partial N^K p^K}{\partial t} \right)$$

or in indicial notation

$$\mathcal{L}(p) \equiv \left(\frac{1}{\rho} N_{,i}^{K}\right)_{,i} p^{K} - \frac{\partial}{\partial t} \left(\frac{1}{\lambda} \frac{\partial N^{K}}{\partial t}\right) p^{K}$$

which is equivalent to having;

$$\mathcal{L}(p) \equiv \mathcal{L}(N^K)p^K$$

using the trial functions as weighting function and recalling the definition of the residual which in this case reads;

$$R = \mathcal{L}(N^K)p^K - q$$

yields;

$$\int\limits_{V} N^{J}RdV = 0 \qquad J = 1, 2, ..., K$$

$$\int\limits_V N^J L(N^K) dV p^K - \int\limits_V N^J q dV = 0$$

$$\int\limits_{V} N^{J} \bigg(\frac{1}{\rho} N_{,i}^{K}\bigg)_{,i} dV p^{K} - \int\limits_{V} N^{J} \frac{\partial}{\partial t} \left(\frac{1}{\lambda} \frac{\partial N^{K}}{\partial t}\right) dV p^{K} - \int\limits_{V} N^{J} q dV = 0$$

Integrating by parts the first term on the R.H.S gives us;

$$-\int\limits_{V}N_{,i}^{J}\frac{1}{\rho}N_{,i}^{K}dVp^{K}+\int\limits_{S}N^{J}\frac{1}{\rho}N_{,i}^{K}\hat{n}_{i}dSp^{K}=\int\limits_{V}N^{J}\frac{\partial}{\partial t}\left(\frac{1}{\lambda}\frac{\partial N^{K}}{\partial t}\right)dVp^{K}+\int\limits_{V}N^{J}qdV$$

$$\int\limits_{V} N_{,i}^{J} \frac{1}{\rho} N_{,i}^{K} dV p^{K} + \int\limits_{V} N^{J} \frac{1}{\lambda} N^{K} dV \ddot{p}^{K} = \int\limits_{S} N^{J} \frac{1}{\rho} N_{,i}^{K} \hat{n}_{i} dS p^{K} + \int\limits_{V} N^{J} q dV$$

$$K^{JK}P^K + M^{JK}\ddot{P}^K + f^J = 0$$

Chapter 3

The Finite Element Method

3.1 Discretization of the PVW via the FEM

A simple spring-mass system

The simple problem of a spring-mass system considered next resembles most of the algorithmic aspects of a finite element code with the advantage that the problem is already a discrete mechanical system. The problem consists of an assemblage of masses joined by different springs submitted to time varying loads. Each spring plays the role of a finite element and each mass is analogous to a nodal point in a finite element algorithm. For instance the full system may be like the one shown in fig. 3.1

Consider a typical spring (finite element) like the one shown in fig. 3.2

The relation between the force and the relative displacement can be written like

$$f_1 = K(u_1 - u_2)$$

and from equilibrium we have

$$f_1 + f_2 = 0$$

which yields the following force-displacement relationship for a typical spring ele-

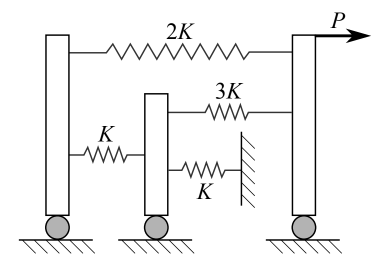


Figure 3.1. Typical assemblage of springs and masses.

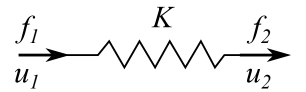


Figure 3.2. Typical spring element.

ment:

$$\begin{cases} f_1 \\ f_2 \end{cases} = K \begin{bmatrix} 1.0 & -1.0 \\ -1.0 & 1.0 \end{bmatrix} \begin{Bmatrix} u_1 \\ u_2 \end{Bmatrix}$$
(3.1)

On the other hand, the equilibrium equation for a typical mass with displacement u_j (see fig. 3.3) and attached to springs i and i+1 reads

$$f_2^i + f_1^{i+1} + m_j \frac{\mathrm{d}V_j}{\mathrm{d}t} = P_j.$$
 (3.2)

which can be written in terms of displacements using eq. (3.1) like

$$(K^{i} + K^{i+1})u_{j} - K^{i}u_{j-1} - K^{i+1}u_{j+1} + m_{j}\frac{\mathrm{d}V_{j}}{\mathrm{d}t} = P_{j}.$$

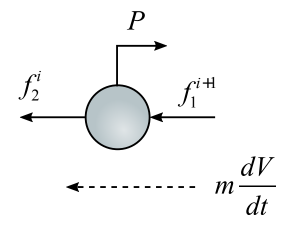


Figure 3.3. Free body diagram for a typical mass connected to springs i and i+1.

Writing the elemental equilibrium equations in terms of u_{j-1} , u_j and u_{j+1} we have:

$$\left\{ \begin{array}{c} f_1^i \\ f_2^i \end{array} \right\} = \left[\begin{array}{cc} k_{11}^i & k_{12}^i \\ k_{21}^i & k_{22}^i \end{array} \right] \left\{ \begin{array}{c} u_{j-1} \\ u_j \end{array} \right\}$$

and

$$\left\{ \begin{array}{c} f_1^{i+1} \\ f_2^{i+1} \end{array} \right\} = \left[\begin{array}{cc} k_{11}^{i+1} & k_{12}^{i+1} \\ k_{21}^{i+1} & k_{22}^{i+1} \end{array} \right] \left\{ \begin{array}{c} u_j \\ u_{j+1} \end{array} \right\}$$

which gives for the equilibrium equation of the m_j mass:

$$k_{21}^{i}u_{j-1} + (k_{22}^{i} + k_{11}^{i+1})u_{j} + k_{12}^{i+1}u_{j+1} + m_{j}\frac{dV_{j}}{dt} = P_{j}.$$

Considering also the contributions from the springs K^i and K^{i+1} to the equilibrium of masses m_{j-1} and m_{j+1} respectively we have the following matrix block:

$$\begin{bmatrix} k_{11}^{i} & k_{12}^{i} \\ k_{21}^{i} & k_{22}^{i} + k_{11}^{i+1} & k_{12}^{i+1} \\ k_{21}^{i+1} & k_{22}^{i+1} & k_{22}^{i+1} \end{bmatrix}$$

Considering now the complete system of masses and springs leads to a system of linear equations of the form

$$[K_G] \{U_G\} + [M] \{A_G\} = \{F_G\}.$$
 (3.3)

where each equation represents the equilibrium of a given mass. The coefficient matrix in eq. (3.3) can be assembled in a systematic way by establishing the connection between the global and local degrees of freedom. This can be accomplished through an operator storing in each row the global degrees of freedom corresponding to each element. For instance, fig. 3.4 shows elements K^i and K^{i+1} and the global degrees of freedom corresponding to masses m_{j-1} , m_j and m_{j+1} . The corresponding entries of the DME operator for these elements are given by:

$$DME = \left[egin{array}{ccc} j-1 & j \ j & j+1 \end{array}
ight]$$

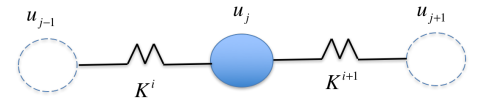


Figure 3.4. Global degrees of freedom connected to the spring elements K^i and K^{i+1} respectively.

and the contribution from these elements to the global coefficient matrix for elements i and i + 1 reads respectively:

$$K_{j-1,j-1} \leftarrow K_{j-1,j-1} + k_{11}^{i}$$

$$K_{j-1,j} \leftarrow K_{j-1,j} + k_{12}^{i}$$

$$K_{j,j-1} \leftarrow K_{j,j-1} + k_{21}^{i}$$

$$K_{j,j} \leftarrow K_{j,j} + k_{22}^{i}$$

and

$$K_{j,j} \leftarrow K_{j,j} + k_{11}^{i+1}$$

$$K_{j,j+1} \leftarrow K_{j,j+1} + k_{12}^{i+1}$$

$$K_{j+1,j} \leftarrow K_{j+1,j} + k_{21}^{i+1}$$

$$K_{j+1,j+1} \leftarrow K_{j+1,j+1} + k_{22}^{i+1}$$

The system given by eq. (3.3), assembled with the aid of the DME operator can be solved for the global displacements U_G . The pseudo-code shown in line 1 presents all the steps required to solve the problem in the context of the finite element method. In that code the so-called DME operator is an equation assembly array indicating how each element contributes to the global stiffness and mass matrix.

```
Algorithm 1: Springs Algorithm.
```

Find internal forces

```
Data: Problem parameters; NUMNP, NUMEL, NMATP Result: Displacements and spring forces Create DME operator; Assemble K^G, F^G; while j \leq 1, NUMEL do K^G \leftarrow K^G + K^i + K^G \leftarrow F^G + F^i end Solve K^G = K^G + K^G
```

```
11 11 11
Computes the displacement solution for an assembly
of 1D spring elements under point loads.
import numpy as np
import starter as sta
import assembler as ass
from datetime import datetime
start_time = datetime.now()
   PRE-PROCESSING
0.00
nodes , mats , elements , loads = sta.readin()
ne , nn , nm , nl = sta.proini(nodes , mats , elements , loads)
DME , IBC , neq = ass.DME(nn , ne , nodes , elements)
   ASSEMBLY
KG = np.zeros([neq, neq])
for i in range(ne):
    kloc , ndof = ass.retriever(elements , mats , nodes , i)
    KG = ass.assembler(KG , neq , kloc , ndof , DME , i)
RHSG = ass.loadasem(loads, IBC, neq, nl)
   SYSTEM SOLUTION
UG = np.linalg.solve(KG, RHSG)
if not(np.allclose(np.dot(KG, UG), RHSG)):
    print("The system is not in equilibrium!")
end_time = datetime.now()
print('Duration for system solution: {}'.format(end_time - start_time))
```

3.1.1 Basic elements of interpolation theory

Let f(x) be a function whose values are known at n discrete points $x_1, x_2, ..., x_n$. We want to know (interpolate) the value of f(x) at an arbitrary point $x \in [x_1, x_n]$.

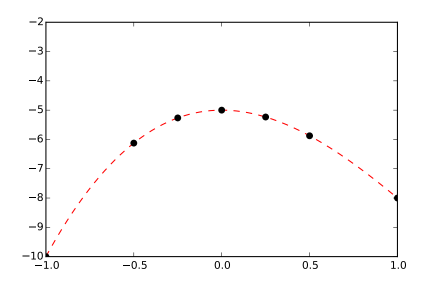


Figure 3.5. Global interpolation of a function

The process of interpolation or computation of the unknown value of f(x) using the known values $\{f^1, f^2, ..., f^n\}$ involves two steps:

- i Fitting an interpolating function to the known data points.
- ii Evaluating the function at the arbitrary point.

We can (i) use all the n-data points and fit an (n-1)-th order polynomial (which is cumbersome and difficult to code) or (ii) split the domain in sub-intervals and use local polynomials within each sub-interval. This last approach involves only a couple of polynomials and it is easy to code, however it may have some continuity issues.

In finite element analysis local interpolation is used in order to proceed systematically. Local interpolation uses a finite number of nearest-neighbors and generates interpolated values f(x) that do not in general have continuous first or higher derivatives.

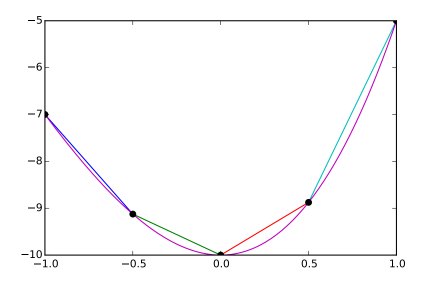


Figure 3.6. Local or piecewise interpolation of a function

Lagrange interpolation theorem

Given a set of n-points $\{(x^1, y^1), \dots, (x^n, y^n)\}$ where $y^n \equiv f(x^n)$ then: "there exists a unique polynomial p(x) of order at most (n-1) such $p(x^I) = f(x^I)$ for $I = 1, 2, \dots, n$ ". The polynomial is given by;

$$p(x^I) = L^I(x)f(x^I) \tag{3.4}$$

for I = 1, 2, ..., n where

$$L^{I}(x) = \prod_{\substack{J=1\\I \neq J}}^{n} \frac{(x - x^{J})}{(x^{I} - x^{J})}$$
(3.5)

and where it should be noticed that

$$L^{I}(x^{J}) = \delta^{IJ}.$$

Example for n=3

Consider the domain [-1,1] and the data points at $x^1 = -1.0$, $x^2 = +1.0$ and $x^3 = 0.0$. We have

$$L^{1}(x) = \frac{(x - x^{2})(x - x^{3})}{(x^{1} - x^{2})(x^{1} - x^{3})} \equiv -\frac{1}{2}(1 -)x$$
$$L^{2}(x) = \frac{(x - x^{1})(x - x^{3})}{(x^{2} - x^{1})(x^{2} - x^{3})} \equiv +\frac{1}{2}(1 + x)x$$

and

$$L^{3}(x) = \frac{(x - x^{1})(x - x^{2})}{(x^{3} - x^{1})(x^{3} - x^{2})} \equiv 1 - x^{2}.$$

The resulting interpolating polynomials $L^{I}(x)$ and the interpolating function are shown in fig. 3.7 below

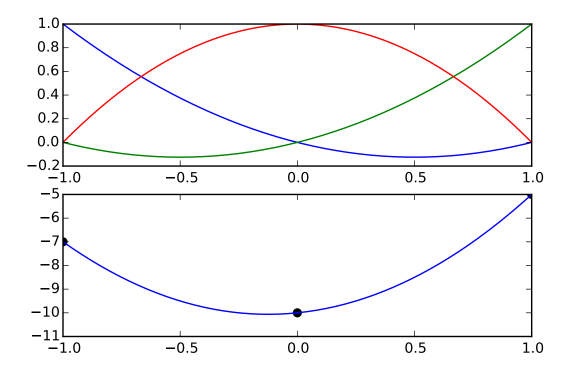


Figure 3.7. Interpolating polynomials and the resulting interpolating function

Example: Interpolation of a function using a global and a local scheme

Assume we have known values of the function:

$$f(x) = x^3 + 4x^2 - 10$$

in the interval [-1.0, 1.0] and we wish to obtain an interpolated version of the function using different schemes.

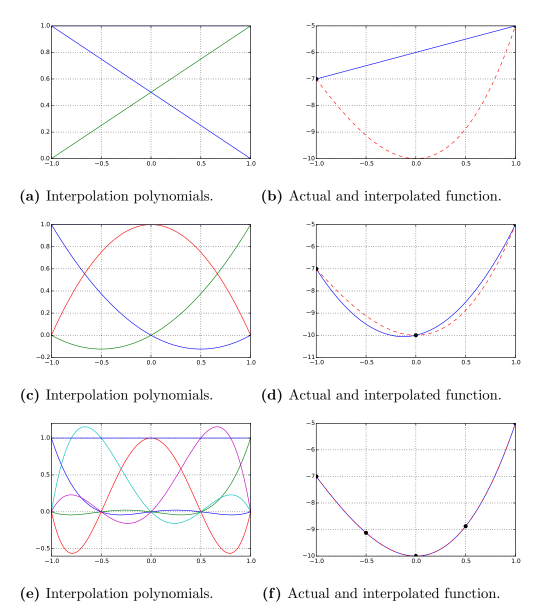
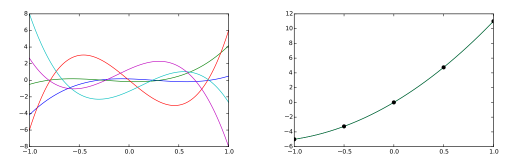


Figure 3.8. Lineal interpolation of the function $f(x) = x^3 + 4x^2 - 10$.

Figure 3.8 shows the interpolating polynomials and the resulting function for the case of first, second and fourth order polynomials respectively. Similarly, the first order derivative obtained out of the interpolated function is displayed in fig. 3.9.



(a) First order derivatives of the inter- (b) Interpolated first order derivative of polation polynomials. the function.

Figure 3.9. Lineal interpolation of the function $f(x) = x^3 + 4x^2 - 10$.

We now proceed like in the finite element method and use the local first order polynomials shown in fig. 3.10 to interpolate the function under study.

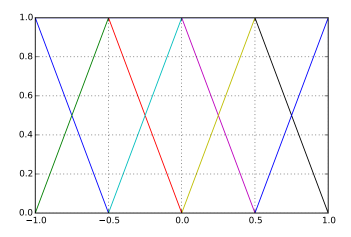
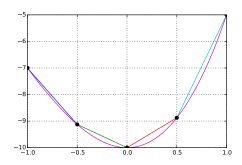
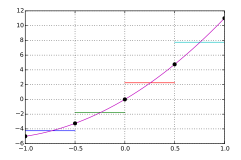


Figure 3.10. Local interpolating polynomials

The resulting function and its numerically obtained first order derivateive are shown in fig. 3.11. It is clear how the local interpolation destroys the global continuity in the first order derivative while the function itself remains continuous.





(a) Function interpolated with local first (b) Interpolated first order derivative of order polynomials. the function.

Figure 3.11. Lineal interpolation of the function $f(x) = x^3 + 4x^2 - 10$.

Extension to 2D domains

Assume we are now interested in conducting interpolation of a function over a spatial 2-dimensional domain where every point is specified by a position vector of the form $\mathbf{x} = x\hat{\imath} + y\hat{\jmath}$. We want to know, via interpolation, the value of a function $f(\mathbf{x})$ at an arbitrary point \mathbf{x} provided we know the set of n-points $\{(\mathbf{x}^1, f^1), \dots, (\mathbf{x}^n, f^n)\}$.

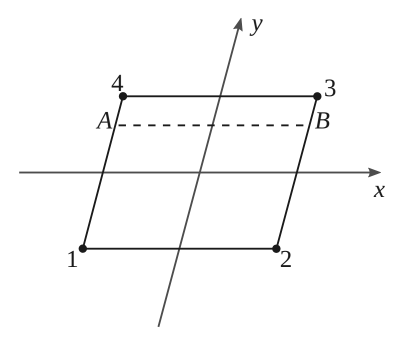


Figure 3.12. Basic square domain

We first fix $x = x^A$ and conduct 1-dimensional interpolation along the y direction as discussed in the previous section as follows (see fig. 3.13)

$$f(x^A, y) = L^1(y)f^1 + L^4(y)f^4$$

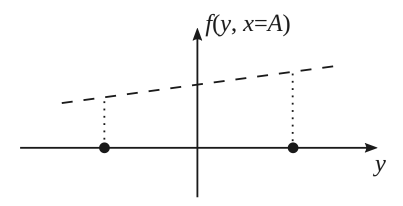


Figure 3.13. Interpolation along the y-direction

Similarly, we can fix $x = x^B$ and interpolate once again along the y direction

$$f(x^B, y) = L^2(y)f^2 + L^3(y)f^3.$$

We now conduct the interpolation along the x-direction using the functions $f(x^A, y)$ and $f(x^B, y)$ respectively as follows

$$\begin{split} f(x,y) &= L^A(x) f(x^A,y) + L^B(x) f(x^B,y) \\ f(x,y) &= L^A(x) \{ L^1(y) f^1 + L^4(y) f^4 \} + L^B(x) \{ L^2(y) f^2 + L^3(y) f^3 \} \\ f(x,y) &= L^A(x) L^1(y) f^1 + L^A(x) L^4(y) f^4 + L^B(x) L^2(y) f^2 + L^B(x) L^3(y) f^3 \ , \end{split}$$

where

$$L^{A}(x) \equiv L^{1}(x)$$

$$L^{B}(x) \equiv L^{2}(x)$$

$$L^{1}(y) \equiv L^{1}(y)$$

$$L^{2}(y) \equiv L^{1}(y)$$

$$L^{3}(y) \equiv L^{2}(y)$$

$$L^{4}(y) \equiv L^{2}(x)$$
.

The function of two variables is then written as the product of one-dimensional interpolations

$$f(x,y) = N^{1}(x,y)f^{1} + N^{2}(x,y)f^{2} + N^{3}(x,y)f^{3} + N^{4}(x,y)f^{4}$$

with

$$\begin{split} N^1(x,y) &= L^1(x)L^1(y) \\ N^2(x,y) &= L^2(x)L^1(y) \\ N^3(x,y) &= L^2(x)L^2(y) \\ N^4(x,y) &= L^1(x)L^2(y) \ . \end{split}$$

See Figure 3.14 for the shape functions of a 4-nodes element.

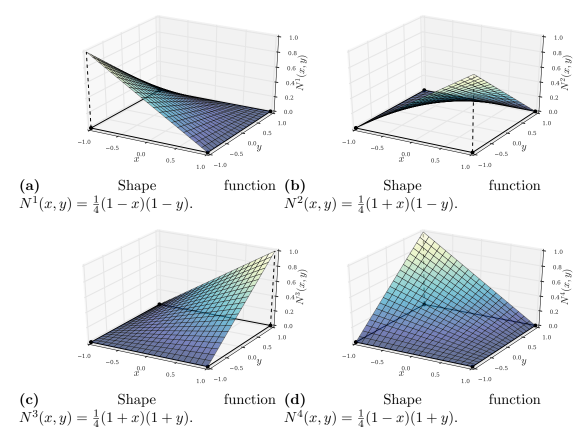


Figure 3.14. Shape functions for a 4-nodes element.

See Figure 3.15 for the shape functions of a 9-nodes element.

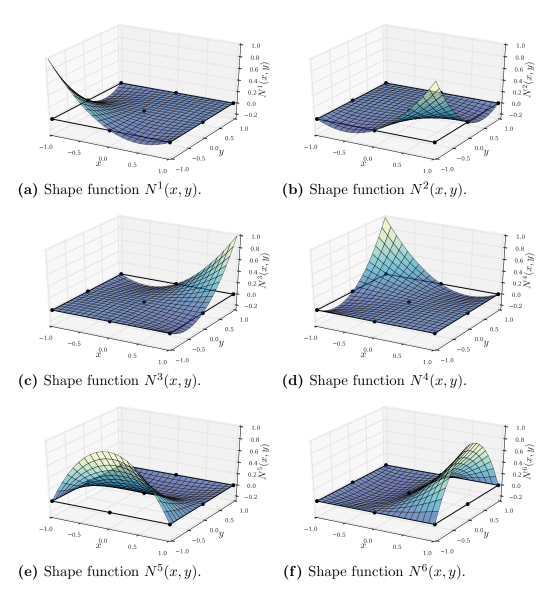


Figure 3.15. Shape functions for a 9-nodes element.

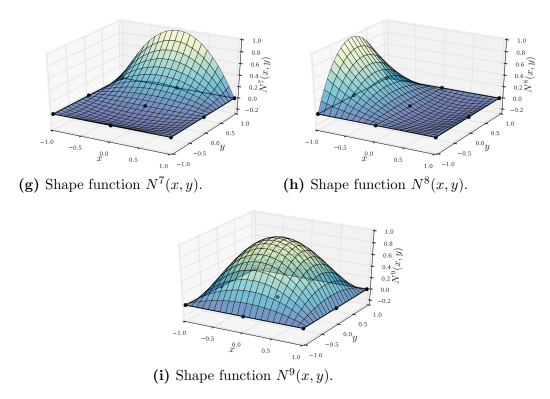


Figure 3.15. Shape functions for a 9-nodes element. (Continued)

3.1.2 Formulation of the finite element matrices

We now discretize the principle of virtual work repeated below for completeness:

$$\int_{V} \sigma_{ij} \delta u_{i,j} \, dV - \int_{V} f_i \delta u_i \, dV - \int_{S_t} t_i^n \delta u_i \, dS = 0.$$
(3.6)

For that purpose we will divide the complete domain V into N-finite non-overlapping sub-domains over each one of which we will approximate the solution in terms of local interpolating functions (see fig. 3.11). Since the PVW (or weak form of the BVP) has been cast into an integral representation, it is possible to build the total integral

considering the contribution of the N-sub-domains like:

$$\sum_{e=1}^{NEL} \int_{V_e} \sigma_{ij} \,\delta u_{i,j} \,\mathrm{d}V^e - \int_{V} f_i \,\delta u_i \,\mathrm{d}V^e - \int_{S_t} t_i^n \,\delta u_i \,\mathrm{d}S^e = 0 \tag{3.7}$$

For simplicity we consider only a single element or sub-domain, thus;

$$\int_{V} \sigma_{ij} \delta \epsilon_{i,j} dV - \int_{V} f_i \delta u_i dV - \int_{S_t} t_i^n \delta u_i dS = 0$$
(3.8)

where it is assumed that the discretized version of each element is later added up (or assembled) into the global equations for the complete model and where we have already used the fact that σ_{ij} is symmetric impliying that $\sigma_{ij}\delta u_{i,j} = \sigma_{ij}\delta\epsilon_{i,j}$.

The involved functions (e.g., displacements, strain, stresses) will be approximated via interpolation of the solution over a determined number of points termed in what follows nodes. Assume for instance that over element e containing n such nodes we know the displacements vector u_i . For instance, fig. 3.16 below shows a typical square 4-noded element of side 2h. The rectangular components of the displacement vector at each node are labeled like $[u_1, u_2, ..., u_7, u_8]$.

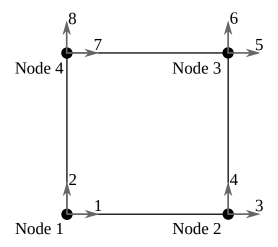


Figure 3.16. Square element of side 2h.

Furthermore, in a more general treatment, we let the displacements for an arbitrary p-node of a 3D problem $u^P = [u^P, v^P, w^P]$. Using ideas from interpolation theory it is now possible to approximate the displacements vector over an arbitrary point \mathbf{x} inside the element by;

$$u_i(\mathbf{x}) = N_i^1(\mathbf{x})u^1 + N_i^2(\mathbf{x})u^2 + \dots + N_i^P(\mathbf{x})u^P + \dots + N_i^n(\mathbf{x})u^n$$

or in more general form

$$u_i(\mathbf{x}) = N_i^Q(\mathbf{x})u^Q \tag{3.9}$$

and where the caption superscripts indicate summation over the number of nodes of the element while the subscript refers to the physical character of the variable being interpolated.

Now let us use the kinematic relationship between strains and displacements:

$$\varepsilon_{ij}(\mathbf{x}) = \frac{1}{2}(u_{i,j} + u_{j,i})$$

together with eq. (3.9). Carrying the derivatives into the shape functions yields;

$$\varepsilon_{ij}(\mathbf{x}) = \frac{1}{2} \left(\frac{\partial N_i^Q}{\partial x_j} + \frac{\partial N_j^Q}{\partial x_i} \right) u^Q$$

which can be written like

$$\varepsilon_{ij}(\mathbf{x}) = B_{ij}^Q(\mathbf{x})u^Q \tag{3.10}$$

after letting

$$B_{ij}^{Q} = \frac{1}{2} \left(\frac{\partial N_{i}^{Q}}{\partial x_{j}} + \frac{\partial N_{j}^{Q}}{\partial x_{i}} \right).$$

Proceeding analogously for the virtual fields after using

$$\delta u_i = N_i^Q(\mathbf{x}) \delta u^Q$$

gives us:

$$\int\limits_{V} C_{ijkl} B_{kl}^{P} u^{P} B_{ij}^{Q} \delta u^{Q} dV - \int\limits_{V} f_{i} N_{i}^{Q} \delta u^{Q} dV - \int\limits_{S_{t}} t_{i}^{n} N_{i}^{Q} \delta u^{Q} dS = 0$$

which can be re-organized into

$$\delta u^Q \int_V B_{ij}^Q C_{ijkl} B_{kl}^P \, dV \, u^P - \delta u^Q \int_V N_i^Q f_i \, dV - \delta u^Q \int_{S_t} N_i^Q t_i^n \, dS = 0$$

which is the generalized discrete version of the PVW for a single element consistent with eq. (3.8). Defining the terms corresponding to the integrals like:

$$K^{QP} = \int_{V} C_{ijkl} B_{kl}^{P} B_{ij}^{Q} dV$$

$$f_{c}^{Q} = \int_{S} N_{i}^{Q} t_{i}^{n} dS$$

$$f_{V}^{Q} = \int_{V} N_{i}^{Q} f_{i} dV$$

$$(3.11)$$

allows us to write:

$$\delta u^Q f_\sigma^Q - \delta u^Q f_V^Q - \delta u^Q f_c^Q = 0.$$

This equation is once again the generalized discrete version of the PVW corresponding to the Q-displacement degree of freedom. The equation quantifies the work

of the forces along the Q-th degree of freedom over the virtual displacements δu^Q . If we now use the arbitrary character of the virtual field δu^Q we can write;

$$f_{\sigma}^{Q} - f_{V}^{Q} - f_{c}^{Q} = 0. {(3.12)}$$

This is now a mechanical equilibrium equation relating the forces along the Q-th degree of freedom associated to the internal element stresses f_{σ}^{Q} ; the external body forces f_{V}^{Q} and the applied external tractions f_{c}^{Q} . Introducing the stiffness of the element gives:

$$K^{QP}u^P = f_V^Q + f_c^Q. (3.13)$$

Formulation for square elements

In the previous section we recognized:

$$K^{QP} = \int_{V} C_{ijkl} B_{kl}^{P} B_{ij}^{Q} dV$$

$$f_{c}^{Q} = \int_{S} N_{i}^{Q} t_{i}^{n} dS$$

$$f_{V}^{Q} = \int_{V} N_{i}^{Q} f_{i} dV$$

$$(3.14)$$

as the elemental stiffness matrix and the vectors of consistent contact and body forces for a single element. Let us assume that the domain of the single element is a perfect square of side 2h (see fig. 3.17), thus;

$$K = \int_{-h}^{+h} \int_{-h}^{+h} B^{T}CBdxdy$$

$$f_{c} = \int_{-h}^{+h} N^{T}t^{n}ds$$

$$f_{v} = \int_{-h}^{+h} \int_{-h}^{+h} N^{T}fdxdy$$

$$(3.15)$$

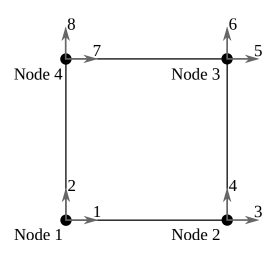


Figure 3.17. Square element of side 2h.

and where N and B are the displacements interpolation matrix and the strain-displacements interpolation matrix respectively. For the linear 4-noded element shown in fig. 3.17 these matrices have the form defined in the following interpolation equations:

$$\left\{ \begin{array}{l} u(\vec{x}) \\ v(\vec{x}) \end{array} \right\} = \left[\begin{array}{ccccc} N^{1}(\vec{x}) & 0 & N^{2}(\vec{x}) & 0 & N^{3}(\vec{x}) & 0 & N^{4}(\vec{x}) & 0 \\ 0 & N^{1}(\vec{x}) & 0 & N^{2}(\vec{x}) & 0 & N^{3}(\vec{x}) & 0 & N^{4}(\vec{x}) \end{array} \right] \left\{ \begin{array}{l} u^{1} \\ u^{2} \\ u^{3} \\ u^{4} \\ u^{5} \\ u^{6} \\ u^{7} \\ u^{8} \end{array} \right\}$$

with the independent shape functions being:

$$N^{1}(x) = \frac{1}{4}(1-x)(1-y)$$

$$N^{2}(x) = \frac{1}{4}(1+x)(1-y)$$

$$N^{3}(x) = \frac{1}{4}(1+x)(1+y)$$

$$N^{4}(x) = \frac{1}{4}(1-x)(1+y)$$

Notice that in the computation of the stiffness matrix we actually require the spatial derivatives of the shape functions given by;

$$\frac{\partial N^1(x)}{\partial x} = -\frac{1}{4}(1-y)$$

$$\frac{\partial N^2(x)}{\partial x} = +\frac{1}{4}(1-y)$$

$$\frac{\partial N^2(x)}{\partial x} = +\frac{1}{4}(1-y)$$

$$\frac{\partial N^3(x)}{\partial x} = +\frac{1}{4}(1+y)$$

$$\frac{\partial N^3(x)}{\partial y} = -\frac{1}{4}(1+x)$$

$$\frac{\partial N^3(x)}{\partial y} = +\frac{1}{4}(1+x)$$

$$\frac{\partial N^4(x)}{\partial y} = -\frac{1}{4}(1-y)$$

$$\frac{\partial N^4(x)}{\partial y} = +\frac{1}{4}(1-y)$$

The contribution to the element matrix from a typical nodal point is thus given by;

$$K = \int_{-h-h}^{+h+a} \begin{bmatrix} \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ \cdots & \frac{\partial N^K(x)}{\partial x} & 0 & \frac{\partial N^K(x)}{\partial y} & \cdots & \cdots \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \end{bmatrix} \begin{bmatrix} A & B & 0 \\ B & A & 0 \\ 0 & 0 & C \end{bmatrix} \begin{bmatrix} \cdots & \frac{\partial N^Q(x)}{\partial x} & 0 & \cdots \\ \cdots & 0 & \frac{\partial N^Q(x)}{\partial y} & \cdots \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \end{bmatrix} dxdy$$

while a typical element of the resultant stiffness matrix takes the general form:

$$K \equiv A \int_{-h}^{+h} \int_{-h}^{+h} \frac{\partial N^K}{\partial x} \frac{\partial N^Q}{\partial x} dx dy + C \int_{-h}^{+h} \int_{-h}^{+h} \frac{\partial N^K}{\partial y} \frac{\partial N^Q}{\partial x} dx dy.$$

It should be noticed that for the considered perfectly square element the resulting stiffness matrix can be partitioned into factors which depend upon the material properties only and into a single factor which depends on the size h. This last contribution can be obtained analytically.

Once the elemental matrices are obtained these are assembled into the final global system of equations. Details of the assembly process are discussed in chapter 4. The following Python script computes the stiffness matrix for the 4-noded element.

11 11 11

```
0.00
from __future__ import division
import numpy as np
from sympy import *
def umat(nu,E):
   """2D Elasticity constitutive matrix"""
    C = zeros(3, 3)
    G = E/(1 - nu**2)
    mnu = (1 - nu)/2.0
    C[0, 0] = G
    C[0, 1] = nu*G
    C[1, 0] = C[0, 1]
    C[1, 1] = G
    C[2, 2] = G*mnu
    return C
def stdm4(x,y):
   """Four noded element strain-displacement matrix"""
   N = zeros(4)
    B = zeros(3,8)
    N = S(1)/4*Matrix([
         (1 - x)*(1 - y),
         (1 + x)*(1 - y),
         (1 + x)*(1 + y),
         (1 - x)*(1 + y)])
    dhdx=zeros(2,4)
    for i in range(4):
        dhdx[0,i]=diff(N[i],x)
        dhdx[1,i]=diff(N[i],y)
    for i in range(4):
        B[0, 2*i] = dhdx[0, i]
        B[1, 2*i+1] = dhdx[1, i]
        B[2, 2*i] = dhdx[1, i]
```

```
B[2, 2*i+1] = dhdx[0, i]
    return B
# Assign symbols
x , y = symbols('x y')
nu, E = symbols('nu, E')
h=symbols('h')
K = zeros(8,8)
# Symbolically compute matrices
C = umat(nu, E)
B = stdm4(x, y)
K_{int} = B.T*C*B
# Integrate final stiffness
for i in range(8):
    for j in range(8):
        K[i,j] = integrate(K_int[i,j], (x,-h,h), (y,-h,h))
knum = K.subs([(E, S(1)), (nu, S(1)/3.0), (h, S(2))])
print knum
```

Formulation in the natural space: the continuum mechanics analogy

In typical finite element equilibrium equations we need to perform integration over the reference element domain $V_0(\mathbf{x})$ corresponding to originally arbitrarily shaped sub-domains as created during the meshing process. In order to proceed with this integration it is useful to consider the following continuum mechanics analogy.

First assume that the actual physical domain $V_0(\mathbf{x})$ is the result of a deformation process imparted upon the natural domain as shown in fig. 3.18. In this analogy, the physical domain $V_0(\mathbf{x})$ is regarded like a "deformed" configuration at an imaginary

time t = t, while the natural "undeformed" domain $V(\mathbf{r})$ is treated like a reference undeformed configurations at time t = 0. Both configurations are assumed to be connected through a deformation process

$$\mathbf{X} = \mathbf{X}(\mathbf{r})$$

$$\mathbf{r} = \mathbf{r}(\mathbf{X})$$
(3.16)

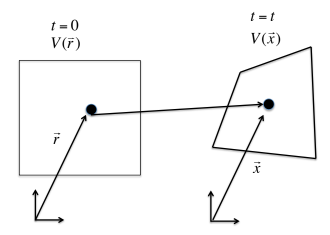


Figure 3.18. Definition of the natural domain

In eq. (3.16) we can understand ${\bf r}$ like a material (Lagrangian) variable and ${\bf X}$ like a spatial (or Eulerian) variable. Using the continuum mechanics analogy it is clear that the "deformation" process at the continuum level is fully characterized by the "deformation" gradient or Jacobian of the transformation eq. (3.16) and defined according to

$$dX_i = \frac{\partial X_i}{\partial r_J} dr_J \equiv J_{iJ} dr_J$$
 (3.17)

where dr_J and dX_i represent material vectors in the original and deformed configuration. From eq. (3.17) it is evident that the Jacobian contains all the information describing the change of the physical sub-domain with respect to the natural element. For the element integration process we will assume that every element $V(\mathbf{r})$ in the natural domain deforms into the physical element $V_0(\mathbf{X})$, thus allowing us to write typical terms like the ones in the material stiffness matrix

$$\int_{V(\mathbf{X})} \hat{B}_{ij}^{K}(\mathbf{X}) C_{ijkl} \hat{B}_{kl}^{P}(\mathbf{X}) dV(\mathbf{X}) \equiv \int_{V_0(\mathbf{r})} \hat{B}_{ij}^{K}(\mathbf{r}) C_{ijkl} \hat{B}_{kl}^{P}(\mathbf{r}) J dV_0(\mathbf{r})$$
(3.18)

where we have used $dV(\mathbf{X}) = JdV(\mathbf{r})$, with J being the determinant of the deformation gradient and in general we transform functions between the natural and physical space making use of eq. (3.16) according to

$$f(\mathbf{r}) = F[\mathbf{X}(\mathbf{r})] \tag{3.19}$$

Interpolation scheme

Having identified the fact that the integration process will take place in the natural domain, we will approach the interpolation process directly in this natural space. In the case of the displacement based finite element method all the involved variables will then be obtained via interpolation of nodal displacements. For instance, assume that a given problem variable is defined in the physical space by the tensor $\Phi_{ik...p}(\mathbf{X})$. The interpolated variable is then written like;

$$\Phi_{ij\dots p}(\mathbf{X}) = H_{ij\dots p}^K(\mathbf{r})\hat{u}^K \tag{3.20}$$

where \hat{u}^K represents a vector of nodal points displacements, see fig. 3.19, and $H_{ij...p}^K(\mathbf{r})$ is an interpolator which keeps the tensorial character of the original physical variable $\Phi_{ik...p}(\mathbf{X})$ and where the super-index makes reference to a nodal identifier (with the summation convention in place).

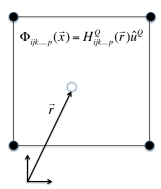


Figure 3.19. General interpolation strategy in the natural domain

Since the primary variable corresponds to displacements it must be kept in mind that $H_{ij...p}^{K}(\mathbf{r})$ corresponds to combinations of derivatives (or other arbitrary combinations) of the basic element shape functions defined in;

$$u_i(\mathbf{X}) = N_i^K(\mathbf{r})\hat{u}^K \tag{3.21}$$

For the general interpolation process we need two kinds of transformations. First we need to transform integrals over the physical space into integrals into the natural space which corresponds to

$$\int_{V(\mathbf{X})} F(\mathbf{X}) dV(\mathbf{X}) \equiv \int_{V_0(\mathbf{r})} f(\mathbf{r}) J dV_0(\mathbf{r})$$
(3.22)

Second we need to relate spatial differentiation in both, the physical and spatial domains. Let us define these operators like ∇_i^X and ∇_I^r respectively. It then follows from eq. (3.19) that

$$\frac{\partial F}{\partial X_i} = \frac{\partial f}{\partial r_J} \frac{\partial r_J}{\partial X_i} \tag{3.23}$$

from where we can establish the connection between the two operators like

$$\nabla_i^X = J_{iJ}^{-1} \nabla_J^r \tag{3.24}$$

The fundamental interpolator

We further define the fundamental interpolator giving rise to gradients of the primary displacement variable in the physical space according to

$$u_{i,j}(\mathbf{X}) = L_{ij}^K(\mathbf{r})\hat{u}^K \tag{3.25}$$

This fundamental interpolator $L_{ik}^K(\mathbf{r})$ is derived after using eq. (3.21) and eq. (3.24) in the physical displacement gradient definition as shown next

$$u_{i,j}(\mathbf{X}) = \nabla_j^X u_i(\mathbf{X})$$

$$u_{i,j}(\mathbf{X}) = \nabla_j^X N_i^K(\mathbf{r}) \hat{u}^K$$

$$u_{i,j}(\mathbf{X}) = J_{jQ}^{-1} \nabla_Q^r N_i^K(\mathbf{r}) \hat{u}^K$$

$$u_{i,j}(\mathbf{X}) = J_{jQ}^{-1} N_{i,Q}^K(\mathbf{r}) \hat{u}^K$$

then

$$L_{ij}^{K}(\mathbf{r}) = J_{jQ}^{-1} N_{i,Q}^{K}(\mathbf{r})$$
(3.26)

Elemental stiffness matrix

The elemental material stiffness matrix computed in the natural domain of fig. 3.20 reads

$$K^{KP} = \int_{V_0(\mathbf{r})} \hat{B}_{ij}^K(\mathbf{r}) C_{ijkl} \hat{B}_{kl}^P(\mathbf{r}) J dV_0(\mathbf{r}) \equiv \int_{r=-1}^{r=+1} \int_{s=-1}^{s=+1} \hat{B}_{ij}^K(r,s) C_{ijkl} \hat{B}_{kl}^P(r,s) J(r,s) dr ds$$

$$(3.27)$$

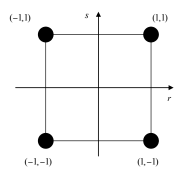


Figure 3.20. Natural domain of integration

Once the interpolator $\hat{B}_{ij}^{K}(\mathbf{r})$ has been identified the elemental stiffness matrix is obtained via numerical integration (quadrature) as described in (3.28)

$$\int_{r=-1}^{r=+1} \int_{s=-1}^{s=+1} \hat{B}_{ij}^{K}(r,s) C_{ijkl} \hat{B}_{kl}^{P}(r,s) J(r,s) dr ds \approx \sum_{i,j=1}^{\text{NGPTS}} \alpha_{i} \alpha_{j} \hat{B}_{kl}^{K}(r_{i},s_{j}) C_{ijkl} \hat{B}_{kl}^{P}(r_{i},s_{j}) J(r_{i},s_{j})$$
(3.28)

and where NGPTS corresponds to the number of integration points, α_j is a weighting factor and r_i, s_j are the coordinates of a typical point \mathbf{r} in the natural space of fig. 3.20.

One important aspect of the numerical integration that has to be kept in mind is

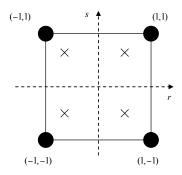


Figure 3.21. Natural integration domain showing quadrature evaluation nodes

accuracy. Depending on the particularly selected integration scheme, the number of introduced integration points fixes the maximum polynomial order of the considered functions that can be integrated accurately. In the case of the integrand in eq. (3.28), it is clear that this order increases as the distortion of the physical element with respect to the natural element increases. One way of dealing with this dependency of accuracy with element distortion is to make use of adaptative integration techniques which are numerically expensive. What is actually done in standard FEM analysis is to choose the number of quadrature points beforehand and introduce distortion related error criteria inside the code in such a way that some sort of validation is performed before the numerical integration process is started.

Strain displacement interpolator for the infinitesimal strain tensor

The Q-th nodal contribution to the infinitesimal strain-displacement interpolator can be obtained in explicit form as follows. Let L_x^Q and L_y^Q be the spatial differential operators in x and y respectively. We have after expanding eq. (3.26)

$$L_x^Q = J_{xP}^{-1} \frac{\partial N^Q}{\partial r_P} \equiv J_{xr}^{-1} \frac{\partial N^Q}{\partial r} + J_{xs}^{-1} \frac{\partial N^Q}{\partial s}$$
$$L_y^Q = J_{yP}^{-1} \frac{\partial N^Q}{\partial r_P} \equiv J_{yr}^{-1} \frac{\partial N^Q}{\partial r} + J_{ys}^{-1} \frac{\partial N^Q}{\partial s}$$

or in matrix form

The Q-th nodal contribution is then assembled as follows;

Algorithm 2: Strain-displacement interpolator

Data: Nodal coordinates x^Q

Result: Strain-displacement interpolator B_{ij}^Q

Compute Jacobian $J_{iJ} = \frac{\partial^2 N_i^Q}{\partial r_J \partial \hat{x}}^Q$

Invert Jacobian $J_{iJ} \to J_{iJ}^{-1}$

Compute fundamental interpolator $L_{ij}^Q = J_{jP}^{-1} \frac{\partial N_i^Q}{\partial r_P}$

Assemble $B_{ij}^Q = \frac{1}{2} \left(L_{ij}^Q + L_{ji}^Q \right)$

Chapter 4

Computational aspects

4.1 Global assembly

In this section we will discuss the fundamental process of building the system of algebraic equations through the addition or assembly of elemental coefficient matrices as described by eq. (4.1) where K^G is the global coefficient matrix; k^i is the local elemental matrix for the *i*-th element; $\bigwedge_{i=1}^{Numel}$ is the assembly operator and *i* is an element index ranging between 1 and the total number of elements Numel that conform the finite element model. Notice that the assembly operator is analogous to the sum operator commonly used in the representation of a series of N terms but it contains information indicating the position of each single term from the element coefficient matrix within the global system.

$$K^G = \bigwedge_{i=1}^{Numel} k^i. \tag{4.1}$$

In this section we will describe the fundamental algorithmic steps to perform that process. We first summarize the basic general steps and then illustrate the process through a simple example involving a 2D mesh of 4-noded elements. The section is complemented with the Python scripts **GLOBAL.py** and the secondary module **assemutil.py**.

The assembly algorithm

The process of assembly of the elemental coefficient matrices into the global system involves (i) the identification of the active degrees of freedom (or equation numbers) assigned to each node in the mesh (ii) the identification of the relationship between the elemental degrees of freedom and the global degrees of freedom and (iii) the computation of the coefficient matrix for each element in the model.

The first step is easily accomplished by assigning a boundary condition flag to the degrees of freedom existing at each node. Here we use a 0 value to indicate an active degree of freedom and a -1 to indicate a prescribed degree of freedom. This information is stored in a boundary condition array IBC, of dimension $nn \times MDIM$, where nn corresponds to the total number of nodal points and MDIM represents the problem dimensionality. The IBC array is first input by the user and later modified by the program in a process where the boundary condition flag is read and equations are counted and assigned according to the result. If the boundary condition flag is equal to 0 the program assigns an equation number while it produces a 0 value if the flag is equal to -1.

In the second step the nodes conforming each element are stored in a connectivity array IELCON of dimension $Numel \times MxNNel$ where Numel is the number of elements in the model and MxNNel is the maximum number of nodes in a given element. Thus each row in the IELCON array stores the nodal point data for the element. Each entry in the IELCON array can now be directly translated into equation numbers using the processed IBC array. This results in the discrete version of the assembly operator $\bigwedge_{i=1}^{Numel}$ also called the assembly list or the DME operator in our codes.

In the final step the mesh is covered one element at a time and each elemental coefficient matrix is assembled into the global matrix as indicated by the DME operator. The actual computation of the elemental matrix is conducted by an element based subroutine, called here UEL, which may be different for each element in the mesh according to different kinematic o material assumptions. The complete process is summarized in line 3 where we describe for completeness additional steps involved in the finite element algorithm. We have introduced additional global and elemental arrays RHS^G and rhs^i respectively, storing the element nodal excitation and the resulting vector of global excitation. This vector is assembled simultaneously, and using the same data, with the global coefficient matrix. In the final step in the

finite element algorithm the system of equations is solved. Notice that the essential boundary conditions were already considered during the assembly process and as a result the global algebraic system of equations is ready to be solved. Finally, after the system has been properly solved the nodal results are scattered through the elements in a process which is inverse to the assembly operation but that uses the same information contained in the DME operator.

Algorithm 3: Summarized algorithm for the finite element method

```
Data: Finite element model

Result: Field function

READ IBC and IELCON arrays;

Compute modified IBC array;

Compute DME operator (using IBC and IELCON arrays);

K^G \leftarrow 0.0;

for i \leftarrow 1 to Numel do

Call UEL(element parameters; K^i);

K^G \leftarrow K^G + K^i (Assemble each k^i into K^G according to the DME operator);

RHS^G \leftarrow RHS^G + rhs^i (Assemble each rhs^i into RHS^G);

end

Solve K^GU^G = RHS^G;

Scatter solution to the elements
```

Sample problem

Consider the mesh shown in section 4.1. The problem parameters in this case are nn = 9, Numel = 4 and MxNNel = 4.

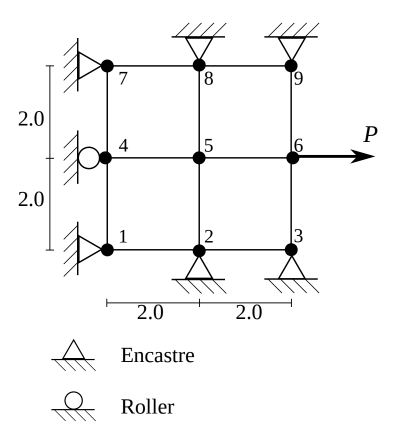


Figure 4.1. Finite element mesh of 4-noded elements. The global matrix can be assembled using the python scripts **GLOBAL.py** and the secondary module **assemutil.py**. The input files required to run the script are called **nodes.txt** and **eles.txt**

The array of boundary conditions as input from the user corresponds to

$$IBC = \begin{bmatrix} -1 & -1 \\ -1 & -1 \\ -1 & -1 \\ -1 & 0 \\ 0 & 0 \\ 0 & 0 \\ -1 & -1 \\ -1 & -1 \\ -1 & -1 \end{bmatrix}$$

while its code-modified version reads

$$IBC = \begin{bmatrix} 0 & 0 \\ 0 & 0 \\ 0 & 0 \\ 0 & 1 \\ 2 & 3 \\ 4 & 5 \\ 0 & 0 \\ 0 & 0 \\ 0 & 0 \end{bmatrix}$$

Notice that the program changes each -1 entry to a 0 valued entry and assigns an equation number to the position containing an original 0 entry during an equation-assigning and equation-counting operation. Similarly, the element connectivity for the current example read

$$IELCON = \begin{bmatrix} 1 & 2 & 5 & 4 \\ 2 & 3 & 6 & 5 \\ 4 & 5 & 8 & 7 \\ 5 & 6 & 9 & 8 \end{bmatrix}$$

The so-called DME operator is actually the same IELCON but translated into

equation numbers for each element. Accordingly in the current example we have

$$DME = \begin{bmatrix} 0 & 0 & 0 & 0 & 2 & 3 & 0 & 1 \\ 0 & 0 & 0 & 0 & 4 & 5 & 2 & 3 \\ 0 & 1 & 2 & 3 & 0 & 0 & 0 & 0 \\ 2 & 3 & 4 & 5 & 0 & 0 & 0 & 0 \end{bmatrix}$$

The definition of the element connectivity and its subsequent translation into degrees of freedom is carried out according to a local element definition like the one shown in section 4.1

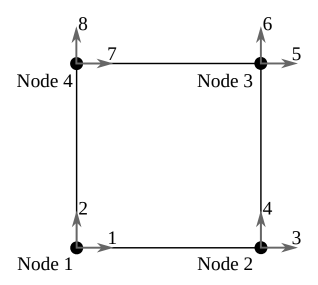


Figure 4.2. Local definition for a 4-noded element.

As a result the (i, j) entry in the DME corresponds to the global equation number associated with the local degree of freedom j of the i element. For instance the value of 4 stored at position (2, 5) in the current DME indicates that the global equation 4 corresponds to the local equation 5 (column index) in element 2 (row index).

The assembly process is then conducted by identifying the relation between the entries in each row of the DME operator and the list of local degrees of freedom for the reference element shown in section 4.1. Accordingly, for element 2 it follows that

the assembly of row 5 of the local stiffness matrix proceeds as follows

$$K_{4,4}^G \leftarrow K_{4,4}^G + k_{5,5}^2$$

$$K_{4,5}^G \leftarrow K_{4,5}^G + k_{5,6}^2$$

$$K_{4,3}^G \leftarrow K_{4,3}^G + k_{5,8}^2$$

In line 3 it must be noticed that the local stiffness matrix for each i element is obtained by the call to the local element subroutine UEL. In the script GLOBAL.py these elemental routines produce a fictitious matrix filled out with ones and the actual computation of the elemental coefficient matrix is discussed later.

4.2 Sparse assembly

In Finite Elements is common to have stiffness and mass matrices that are sparse, i.e., matrices in which most of the elements are zero.

For instance, in a regular mesh formed with bilinear quadrilaterals the number of nonzero entries is given by the expression

$$storage = 9n_x n_y - 9n_x - 3n_y + 4,$$

where n_x is the number of nodes in the x coordinate and n_y is the number of nodes in the y coordinate. Figure 4.3 presents the needed storage for a mesh with $n_x = n_y$ for different sizes.

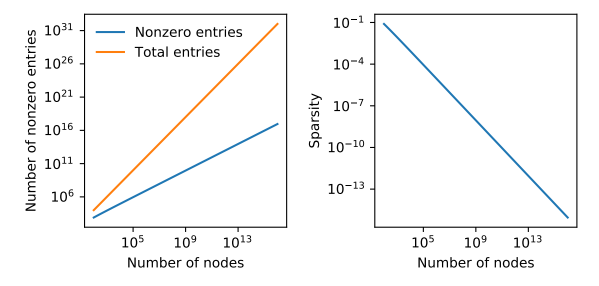


Figure 4.3. Nonzero entries in a sparse matrix for a structured mesh of bilinear elements.

4.3 Python implementations

4.4 Commercial codes and user subroutines

Chapter 5

Wave propagation problems in the time domain

- 5.1 Problem formulation
- 5.2 Explicit time integration algorithm
- 5.3 Statement of the Problem

Consider the discrete dynamic equilibrium equations at time t

$$M^t A + C^t V + K^t U = {}^t F (5.1)$$

where M, C, K are the assembled mass, damping and stiffness matrix respectively and similarly tA , tV , tU , tF are the nodal accelerations, velocities, displacements and external loads vectors at time t. In terms of forces, eq. (5.1) can be written like;

$${}^{t}F^{I} + {}^{t}F^{D} + {}^{t}F^{s} = {}^{t}F$$
 (5.2)

where ${}^tF^I$, ${}^tF^D$ and ${}^tF^s$ are inertial, damping and elastic nodal forces respectively.

Expanding the acceleration and velocity terms at time t in a consistent finite central differences scheme we have;

$${}^{t}A = \frac{1}{\Delta t^{2}} \left({}^{t-\Delta t}U - 2^{t}U + {}^{t+\Delta t}U \right)$$

$${}^{t}V = \frac{1}{2\Delta t} \left({}^{-t-\Delta t}U + {}^{t+\Delta t}U \right)$$

$$(5.3)$$

It is convenient to write eq. (5.3) like;

$${}^{t}A = {}^{t}\hat{A} + \frac{1}{\Delta t^{2}} {}^{t+\Delta t}U$$

$${}^{t}V = {}^{t}\hat{V} + \frac{1}{2\Delta t} {}^{t+\Delta t}U$$

$$(5.4)$$

where:

$${}^{t}\hat{A} = \frac{1}{\Delta t^{2}} \left({}^{t-\Delta t}U - 2^{t}U \right)$$

$${}^{t}\hat{V} = -\frac{1}{2\Delta t} {}^{t-\Delta t}U$$

$$(5.5)$$

are termed predictors, while $\frac{1}{\Delta t^2}{}^{t+\Delta t}U$ and $\frac{1}{2\Delta t}{}^{t+\Delta t}U$ are termed correctors.

Using eq. (5.3) in eq. (5.1) yields;

$$\left(\frac{1}{\Delta t^2}M + \frac{1}{2\Delta t}C\right)^{t+\Delta t}U = {}^tF - M^t\hat{A} - C^t\hat{V} - K^tU$$
(5.6)

or after letting;

CHAPTER 5. WAVE PROPAGATION PROBLEMS IN THE TIME DOMAIN 81

$$\hat{K} = \frac{1}{\Delta t^2} M + \frac{1}{2\Delta t} C$$

and

$${}^t\hat{F} = {}^tF - M^t\hat{A} - C^t\hat{V} - K^tU$$

we have;

$$\hat{K}^{t+\Delta t}U = {}^t\hat{P}$$

which can be solved for $^{t+\Delta t}U$.

Algorithm 4: Explicit algorithm

Data: Values at t = 0, Geometry, Material Paramters

Result: Displacements, Velocity and Acceleration at any time t = t

Compute predictors ${}^{t}\hat{A}$, ${}^{t}\hat{V}$, ${}^{t}\hat{U}$

Assemble \hat{K} , $^t\hat{P}$

Solve $\hat{K}^{t+\Delta t}U = {}^t\hat{P}$

Perform Corrections ${}^tA \leftarrow {}^t\hat{A} + \frac{1}{2\Delta t^2}{}^{t+\Delta t}U$ and ${}^tV \leftarrow {}^t\hat{V} + \frac{1}{2\Delta t}{}^{t+\Delta t}U$

Redefine forces as follows

$${}^{j}F^{I} = \frac{1}{\Delta t^{2}}M^{j}U$$

$${}^{j}F^{D} = \frac{1}{2\Delta t}C^{j}U$$

$${}^{j}F^{S} = K^{j}U$$

$$(5.7)$$

and write (5.6) as

$$^{t+\Delta t}F^I + ^{t+\Delta t}F^D = ^tF - ^tF^s + 2^tF^I - ^{t-\Delta t}F^I + ^{t-\Delta t}F^D$$
 (5.8)

• (5.8) is an equilibrium equation at time t = t allowing to predict the displacements at time $t = t + \Delta t$ in terms of previously known values at times t and

$$t = t - \Delta t$$
.

• The equation is exact within the error introduced by the expansion used in (5.3).

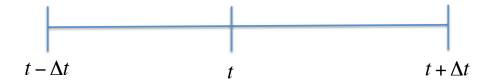


Figure 5.1. Definition of the general iteration

• The first predicted solution is at $t = \Delta t$ and we require data at $t = -\Delta t$ and at t = 0.

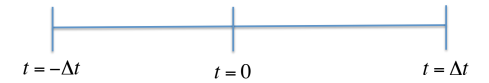


Figure 5.2. Definition of the general iteration

5.3.1 Damping Assumptions

1 Use Rayleigh Damping and retain the velocity expansion used in (5.3). That is;

$$C = \alpha M + \beta K \tag{5.9}$$

then we have (in terms of forces);

$$(1 + \beta \Delta t^2)^{t+\Delta t} F^I + \frac{\alpha}{2\Delta t}^{t+\Delta t} F^S = {}^t \hat{F}$$
 (5.10)

where;

$${}^{t}\hat{F} = {}^{t}R - {}^{t}F^{S} + 2{}^{t}F^{I} - {}^{t-\Delta t}F^{I} + {}^{t-\Delta t}F^{D}$$

CHAPTER 5. WAVE PROPAGATION PROBLEMS IN THE TIME DOMAIN 83

Solution in equation (5.10) requires the full assembly and factorization of an effective stiffness matrix.

2 Neglect damping (This is however inconvenient for finite domains)

$$^{t+\Delta t}F^I = ^t F - ^t F^S + 2^t F^I - ^{t-\Delta t} F^I$$
 (5.11)

3 Use Rayleigh damping but modify the velocity expansion introduced in (5.3). Using

$${}^{t}V = \frac{1}{\Delta t} ({}^{t}U - {}^{t-\Delta t}U) \tag{5.12}$$

yielding;

$$^{t+\Delta t}F^I = ^t F - ^t F^S + 2^t F^I - ^t F^D + ^{t-\Delta t} F^D - ^{t-\Delta t} F^I$$
 (5.13)

Defining a set of forces associated to the initial conditions like;

$$^{t-\Delta t}F^{IC} = ^{t-\Delta t}F^{I} - ^{t-\Delta t}F^{D}$$

we have

$$^{t+\Delta t}F^I = ^t F - ^t F^S + 2^t F^I - ^t F^D - ^{t-\Delta t} F^{IC}$$
 (5.14)

5.3.2 Algorithm corresponding to the damping assumption 3

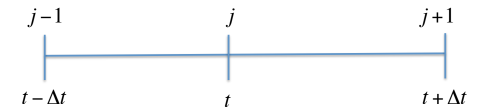


Figure 5.3. Definition of the general iteration

Let us write (5.14) like

$$^{j+1}F^I = ^j F - ^j F^S + 2^j F^I - ^j F^D - ^{j-1} F^{IC}$$
 (5.15)

where the initialization process corresponds to;

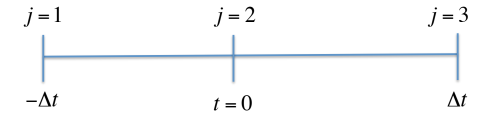


Figure 5.4. Definition of the initial iteration

Applying (5.15) for t = 0 we have;

$${}^{0}F^{I} = {}^{0}F - {}^{0}F^{S} + 2{}^{0}F^{I} - {}^{0}F^{D} - {}^{-\Delta t}F^{IC}$$

from which it is clear that we require $-^{\Delta t}U$. Applying the central difference expansion at t=0 and solving for $-^{\Delta t}U$ yields;

$$^{-\Delta t}U = {}^{0}F - \Delta t^{0}V + \frac{\Delta t^{2}}{2}^{0}A$$
 (5.16)

Using (5.16) in (5.15) allows us to start up the algorithm.

Particulars

In what follows we concentrate on this last algorithm and in order to study some details we return to its standard displacements form. Writing (5.15) in terms of displacements and re-arranging yields;

$$\frac{1}{\Delta t^2}M^{t+\Delta t}U = {}^tF - (1 + \frac{\beta}{\Delta t})K^tU + (\frac{2}{\Delta t^2} - \frac{\alpha}{\Delta t})M^tU - (\frac{1}{\Delta t^2} - \frac{\alpha}{\Delta t})M^{t-\Delta t}U + (\frac{\beta}{\Delta t})K^{t-\Delta t}U$$
 (5.17)

Let;

$$a_1 = 1 + \frac{\beta}{\Delta t}$$

$$a_2 = \frac{2}{\Delta t^2} - \frac{\alpha}{\Delta t}$$

$$a_3 = \frac{1}{\Delta t^2} - \frac{\alpha}{\Delta t}$$

$$a_4 = \frac{\beta}{\Delta t}$$

$$^{t+\Delta t}F^I = ^t F - a_1K^tU + a_2M^tU - a_3M^{t-\Delta t}U + a_4K^{t-\Delta t}U$$
 (5.18)

5.3.3 Decoupling

Consider the equation for the i-th d.o.f;

$$\frac{1}{\Delta t^2} M_{ij}^{t+\Delta t} U_j = {}^t F_i - a_1 K_{ij}^t U_j + a_2 M_{ij}^t U_j - a_3 M_{ij}^{t-\Delta t} U_j + a_4 K_{ij}^{t-\Delta t} U_j$$
 (5.19)

where we keep i fixed in (5.19). For a lumped mass matrix we can write;

$$M_{ij} = m_I \delta_{ij}$$

then (5.19) becomes;

$$\frac{1}{\Delta t^2} m_I^{t+\Delta t} U_i = {}^t F_I - a_1 K_{ij}^t U_j + a_2 m_I^t U_i - a_3 m_I^{t-\Delta t} U_i + a_4 K_{ij}^{t-\Delta t} U_j$$
 (5.20)

Let;

$$t^{t+\Delta t}F_{i}^{I} = \frac{1}{\Delta t^{2}}m_{I}^{t+\Delta t}U_{i}$$

$$t^{t}\hat{F}_{i}^{S} = a_{1}K_{ij}^{t}U_{j}$$

$$t^{t}\hat{F}_{i}^{I} = a_{2}m_{I}^{t}U_{i}$$

$$t^{t-\Delta t}\hat{F}_{i}^{I} = a_{3}m_{I}^{t-\Delta t}U_{i}$$

$$t^{t-\Delta t}\hat{F}_{i}^{S} = a_{4}K_{ij}^{t-\Delta t}U_{j}$$

$$(5.21)$$

so the recursive equation takes the form;

$$^{t+\Delta t}F_{i}^{I} = ^{t}F_{i} - ^{t}\hat{F}_{i}^{S} + ^{t}\hat{F}_{i}^{I} - ^{t-\Delta t}\hat{F}_{i}^{I} + ^{t-\Delta t}\hat{F}_{i}^{S}$$

$$(5.22)$$

and the algorithm then reduces to;

Algorithm 5: Summarized Algorithm

Data: Time span, Geometry, Material Parameters

Result: Displacements, Velocity and Acceleration time histories $t + \Delta t R^{I}$

Compute ${}^{t+\Delta t}F_i^I$

Solve for
$${}^{t+\Delta t}U_i = \left(\frac{\Delta t^2}{m_I}\right)^{t+\Delta t} F_i^I$$

Update tV_i , tA_i

To initialize the algorithm we apply the FD's equations at t=0

$$\frac{1}{\Delta t^2} m_I^{\Delta t} U_i = {}^{0} F_i - a_1 K_{ij}^{0} U_j + a_2 m_I^{0} U_i - a_3 m_I^{-\Delta t} U_i + a_4 K_{ij}^{-\Delta t} U_j$$

where $^{-\Delta t}U_i$ is obtained from (5.16)

CHAPTER 5. WAVE PROPAGATION PROBLEMS IN THE TIME DOMAIN 87

$$^{-\Delta t}U_i = ^{0}U_i - \Delta t^0 V_i + \frac{\Delta t^2}{2} A_i$$
 (5.23)

The initial acceleration is obtained after assuming homogeneous IC's;

$$m_I^0 A_i + C_{ij}^0 V_j + K_{ij}^0 U_j = {}^0 F_i$$

therefore

$$m_I^0 A_i = \frac{{}^0F_i}{m_I}$$

$${}^{-\Delta t}U_i = \frac{\Delta t^2}{2m_I}^0 F_i$$

Moreover, neglecting the damping effects on the prediction of $^{\Delta t}U_i$ yields;

$$^{\Delta t}U_i = \frac{\Delta t^2}{2m_I}^0 F_i$$

Algorithm 6: Full Algorithm

Data: Time span, Geometry, Material Parameters

Result: Displacements, Velocity and Acceleration time histories

Initialize solution vectors (j = 1);

$${}^{0}U_{i} \longrightarrow {}^{1}U_{i} = 0, {}^{0}V_{i} = 0, {}^{1}A_{i} = \frac{{}^{1}R_{i}}{m_{I}};$$

Select Δt and integration constants a_1, a_2, a_3, a_4 ;

Fix 1-st predicted value (let j = 2);

$$^{\Delta t}U_i \longleftarrow \frac{\Delta t^2}{2m_I}^0 F_i \longleftrightarrow \left[^2U_i \longleftarrow \frac{\Delta t^2}{2m_I}^1 F_i \right]$$

Time Integration Phase;

while
$$j \leq N$$
 do

end

Nodal Assembler

In the uncoupled explicit finite element formulation the equation solving process proceeds one degree of freedom at a time. This implies a different assembly process to the one used in an implicit algorithm where a formal coefficient matrix is assembled and inverted. Now the mass, damping and stiffness elemental matrices are used to obtain effective nodal forces at each degree of freedom. In summary the mesh is not covered in an element by element basis, but in a node by node basis. In the following algorithm we discuss this nodal assembly process where in order to solve the displacement at a given degree of freedom prior knowledge of the element contributing to the given node is necessary. In the nodal assembler algorithm the

CHAPTER 5. WAVE PROPAGATION PROBLEMS IN THE TIME DOMAIN 89

following arrays are needed.

ILIST(): Stores the elements connected to the current node.

LPLIST():Stores the local position of the current node in each one of the elements of ILIST().

NIEL():Number of elements at the current node. This array is used to access ILIST().

Algorithm 7: Nodal Assembler

Data: Number of nodal points, number of elements, Model

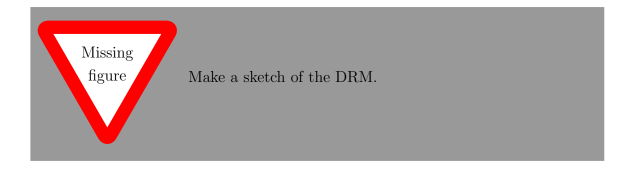
Result: Displacements, Velocity and Acceleration time histories

while $i \leq NUMNP$ do

 $K \leftarrow NIEL(i)$

end

5.4 The DRM approach



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