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Julian Anthony Camsron Diering

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A dissertation submitted to the Faculty of Engineering University of the Nitwatersrand, Johannesburg for the Degree of Master of Science

Johannesburg 1981.

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I declare that this dissertation is my own, unaided work. It is being submitted for the degree of Master of Science in the University of the Witwatersrand, Johannesburg. It has not been submitted before for any degree or examination in any other University.

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A C K N O W L E D G E M E N T S

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The writer wishes to thank Dr T R Stacey for the advice and many stimulating discussions at all stages of the work. He would also like to thank Steffen, Robertson and Kirsten Inc for use of their computing and drawing office facilities and for allowing time to **complete** these studies.

Special thanks are due to Pam Le Vieux and Anne **Melhuish** for typing most of the dissertation and Denise for her continued interest and moral support.

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FURTHER DEVELOPMENTS OF THE BOUNDARY ELEMENT METHOD WITH APPLICATIONS IN MINING

DIERING, Julian Anthony Cameron, University of Wiwatersrand, 1981.

Three computer programmes designed for the determination of stresses and displacements in and around mine excavations are described. The first is a three-dimensional, boundary element formulation which allows for modelling of large scale **non-homogeneities** in the rock mess surrounding the mine excavations. In addition, shear or tensile failure of the geological interfaces may be modelled in a realistic manner. The second is a "mixed boundary element" formulation comprising three-dimensional boundary and displacement discontinuity **elements** into a single programme. The programme enables the interaction of planar or tabular features with qpen or massive excavations to be modelled efficiently. The third, an extension to an existing programme enables mining in non-hcmogeneous ground to be modelled in two dimensions using the displacement discontinuity method.

Examples are given demonstrating the applicability of these programmes to mining problems. The programmes will run on most mini computers making them practical design aids readily available to the rock mechanics engineer.

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- 1 EQUIVALENCE OF DISPLACEMENT DISCONTINUITY **AND** BOUNDARY **ELEMENT STRESS AND DISPLACEMENT FUNCTIONS**
- **2 PROGRAMME LISTING FOR PROGRAMME BEM**
- **3 PARTIAL LISTING (IF PROGRAMME MBEM**
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CHAPTER 1 ItTTKQDUCTION

: 1,1 Background

The determination of stresses and displacements in and around mine excavations plays an important role in mine planning and mining rock mechanics, Usually, however, the geology surrounding and the geometry of the mine excavation are so complex that (a) analytic solutions are not available and (b) numerous simplifying assumptions have to be made about the geology and geometry before numerical or, sometimes, analytical solutions may be obtained. The geology of the problem is usually simplified by assumptions such as homogeneity, isotropy and linear elastic material while geometric simplifications include two-dimensional or axisymmetric representation of a fully three-dimensional problem, an assumption of an infinite, finite or semi-infinite region of space and smoothing of excavation surfaces.

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The limited applicability of analytic solutions to practical mining problems together with the ready availability of digital computers has resulted in an ever increasing use of computer based stress analysis techniques in mining rock mechanics. Three major classes of numerical Stress analysis have emerged, namely the finite difference, finite element and surface element methods,

The finite difference method finds **some** application in simple time dependent problems but has been largely superceded by the finite element method. This method requires that a sufficiently large volume of material **surrounding** the mine workings being analysed be divided into volume elements. Simplifying assumptions about the stresses and displacements within an element are made and each element only influences its neighbours. As **such,** each element may **be** assigned unique properties so that the finite element **method** is well suited to the analysis of non-homogeneous or **non-linearly** elastic problems.

Surface element methods describe a problem in terms of the excavation surfaces, geological interfaces and very often the surrounding ground surface also. These surfaces are divided into surface or boundary **elements and their mutual interaction calculated so as to satisfy boundary conditions imposed on the surfaces.**

It is immediately apparent that surface area to volume ratio of a problem will determine the relative applicability of a finite or surface element technique. Equally important are the degrees of non-homogeneity and non-linear "behaviour involved. Both methods (and in **fact most stress analysis formulations) usually assume that the host rock is isotropic. The validity of this assumption in most problems is generally accepted even though numerous rook types are grossly anisotropic. The assumption of isotropy is therefore maintained throughout the rest of this dissertation.**

Surface or boundary element methods are based upon the numerical solution of the boundary integral equation (BIB). Different formula**tions** of the BIS include specification of **surface** tractions and displacements, "fictitious forces" and displacements or surface trac**tions** and displacement discontinuities. The displacement discon**tiuuity formulation forms a special class of boundary element method commonly referred to as the displacement discontinuity method. Distinction** is hereafter made between the displacement discontinuity **method** (DEM) and other boundary element methods (BEM) and the finite element **method** (BEM).

Just as the FHM and BEM formulations have their relative merits and disadvantages so do the BEM and DEM formulations. To a first degree of approximation it may be said that the DEM is best suited to the modelling of narrow **or** tabular excavations and their interaction with faults or joints while the BEM is well suited to modelling open excavations with the presence of limited non-homogenities.

Existing formulations

Druse (1969) described a boundary element formulation in three

dimensions for homogeneous bodies. Boundary conditions at the surface elements are specified **Jin terms** of constant tractions and displacements over triangular elements. Examples are **given** to demonstrate the applicability of the formulation to fairly simple problems. Evaluation of influence coefficients (the influence of one component of displacement or traction of one element upon another) **is** done analytical ly. **'the** main draw back of this formulation is that a large number of elements are required for practical problems.

CrUse (1974) described an improved version in which displacements and tractions are allowed to vary linearly over each surface element. This formulation gives **improved** accuracy for the same number of surface elements.

A boundary element formulation in which curved **elements** with linear, quadratic or cubic variation of tractions and displacements is allowed over each element was described by lachat and Ifetson (1976). **A** canputer programme was described, which is capable of handling a wide range of **problems** including thin plate problems. The computer programme is very long (about 10 000 lines of Fortran IV) and might not be well suited to run on small mini-computers. A problem which arises when higher order elements are used is that the integration procedures described to date will only work for finite geometries. It is possible that minor modifications to these **programmes would** enable them to model typical rock mechanics problems, although no literature describing any such modifications was found.

Examples given in the above formulations are related primarily to mechanical engineering and fracture mechanics and solution of the equations is carried out using Gaussian Elimination, a technique not well suited to the solution of large systems of linear equations on a small mini-computer.

Deist and Georgiades (1976) described a slightly different approach in which displacements and "fictitious forces" are taken as constant over flat triangular elements. Evaluation of influence coefficients is done numerically and the equations are solved Using a stationary

second degree iterative solution technique not unlike successive over relaxation (SOR). This iterative technique offers considerable savings of computation time. Machine time is further reduced by the inplenentation of a sophisticated "lumping mechanism" whereby groups of elements are treated as single elements when calculating their influences upon other remote elements. Examples are given shewing the applicability of this programme to mining rock mechanics problems. The programme assumes a 'homogeneous rockmass and is also too large to be easily implemented on a mini-conputer.

Bannerjie and Butterfield (1977) describe, a formulation similar to that of Cruse (1969) and give examples of applications in soil mechanics. None of the above formulations allow for slip or failure to occur at an interface (fault or joint) unless such failure is **implemented** manually step by step. Hocking (1976) has attempted to implement slip on **two-dinensidnal** boundary element interfaces. His approach, **however,** met with little success: "results should be Viewed **with** suspicion until validation is obtained".

The displacement discontinuity method has found wide application for mining **problems** involving tabular excavations. Three-dimensional: formulations have been described by Salaron (1963, 1964 (a), (b), (c)) and Starfield and Crouch (1973) in which, typically, a planar tabular excavation remote from the earth's surface is divided into a large number of square elements. The relative movement between hangingwall and footwall defines the "displacement discontintuity" which is assumed constant over each element. These formulations cannot model the interaction between tabular excavations and the ground surface **or** other non-tabular excavations.

Morris (1976) of **the** Chamber of Mines of South, Africa has extended » the method for tabular excavations close to but not **outcropping** at the earth's surface or for a series of parallel tabular excavations. These formula'.ions cannot model outcropping excavations or any interaction with **non-tabular** excavations or geological discontinuities as is the case with the programmes described in this dissertation.

Crouch **(1976)** extended **the DEM** in **two** -dimens ions to handle excavations of arbitrary shape in a homogeneous rock mass. Failure of faults and joints is realistically modelled by means of a Mohr-Coulomb failure criterion. This very useful extension to the DEM cannot, however, model non-homogeneities.

1.3 Scope of the dissertation

The major portion of this dissertation is concerned with stress analysis formulations in three- dimensions. Numerical and computational problems associated with three-dimensional analyses are in general mich greater than for equivalent two-dimensional analyses. Execution times, storage requirements, data preparation tines and degrees of freedom are usually an order of magnitude greater than for two-dimensional formulations. As a result, the cost of a threedimensional stress analysis is usually high and often prohibitive.

One approach used to alleviate the problem centres around the introduction of sophisticated elements. **%ienkiewicz** (1971) has used sophisticated finite elements to great advantage while Lachat and Watson (1976) and Cruse (1973) have introduced improved boundary elements with equal success. With this approach it is still necessary to solve most problems on large main **frame** systems.

The approach adopted in this dissertation relies on efficient handling of a large number of simple elements. Reduction of main and disk storage requirements, programme size and execution times were main goals of this dissertation. In particular it was necessary that any formulation be able to run on a small 16- bit mini-computer. Satisfaction of this requi rement results in a great cost reduction to those users who have mini-computers but have to rely cn commercial computing* beureux for three-dimensional problems. A similar approach has been used by Deist and Geordiodes (1976). Much of the experience gained in efficient handling of a large number of simple elements is directly applicable to the more sophisticated boundary elements.

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The first formulation described here is based on that of Cruse (1969). He introduced a simple triangular boundary element. Calculation of influence coefficients - the effect of one element on another - is done analytically and the resulting equations are solved using Gaussian elimination with iteration on the residues. His examples are concerned with problems in fracture mechanics. The following changes are made to his formlation;

- (i) Equations are solved iteratively using the method of successive over relaxation.
- (ii) Elements are grouped into "lump" elements.
- (iii) Non-homogeneous problems may be analysed.

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- (iv) Slip or failure of interfaces may be modelled by a Mohr-Coulamb failure criterion.
- (v) A variety of symmetry conditions **may be** imposed.
- (vi) The prograntre will run **on** a small itini-conputer.

The second formulation combines the above boundary element programme with a displacement discontinuity formlation based upon that of Starfield and Crouch (1976). Displacement discontinuity **elements** are used to **model** a fault or a tabular excavation while the boundary elements may be used to model the earth's surface, an open pit or a massive excavation. This formulation has significant advantages over the first for many problems.

Finally an improvement to the two-dimensional displacement discontinuity formulation of Crouch (1976) is described here. He described modelling of mining in faulted ground which is homogeneous. The prograrnre M1NAP of Crouch is modified here enabling modelling of a large number of non-honogeneous problems.

Examples are given to **test** the accuracy of the **three-dimensional** formulations against analytic or other formulations and which demonstrate the applicability of these programmes to practical rock mechanics problems. These include:

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- **(i) & long rectangular tunnel**
- (ii) Two massive excavations close to the earth^{'s} surface
- (iii) Interaction between an underground tabular and open pit exca**vations**
- **Tabular excavation mining up to a fault**
- **Interaction beWeeh massive underground and open pit excavations**
- **(vi) A pile socketed in rock with slip on the pile/rock inter**face.

The three farnulaticns are henceforth referred to by the programme names:

Briefly the contents of the dissertation are as follows:

Chapter 2 gives the basic equations for the BFM and MBEM programmes.

Chapter 3 describes the numerical integration procedures used for evaluating influence coefficients.

- Chapter 4 describes **the implementation** of lumping into the. BEM and MBEM **programmes,** The **primary** objectives of the lumping $mechanism area:$
	- (i) to convert a full system of equations into one **which is about 20% to 50% populated thus reducing disk storage and execution time,**

(ii) to produce additional checks on input data and

(iii) to reduce main memory required

- Chapter 5 describes the **implementation of** interface elements and symmetry conditions. When **two** or more subregions with **different elastic properties are being modelled, it is possible to allow the material interface to fail in shear or in tension.**
- Chapter 6 describes a similar implementation of interface elements into the two-dimensional programme MINAP of Crouch (1976).

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This allows for modelling of a wide range of nog-hanogenedus problems while **allowing** for possible tensile or shear failure of the material interfaces. The contents of **this** chapter form the basis of a recent publication Diering (1980a). **Chapter 7 describes programme validation, with a few examples to** assess programme accuracy and numerical integration sensitivity. Chapter 8 gives a brief discussion of some of the programming considerations. Particular attention is given to disk **storage and disk access considerations^** Chapter 9 contains various examples **demonstrating** a wide variety of

Chapter 10 gives conclusions and a general discussion of the dissertation.

APPENDICES

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1 Equivalence of displacement discontinuity and boundary element stress and displacement functions.

2-4 Complete or partial listings of the various programmes.

applications in rock mechanics,

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CHAPTRR 2 DEFINITION OF TERMS AND GOVERNING EQUATIONS

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The notation used here is the normal Cartesian tensor notation with summation over repeated indices and the comma representation of partial differentiation. The basic equations presented in 2.1 and 2.2 are derived by **Cruse (1969), Starfield and Crouch (1973) or Lachat and Matson (1976).**

2 .1 Boundary element formulation

The region of interest or rock mass may be divided into a number of subregions &(%), each of vhich may have different elastic constants $E^{(k)}$ and $v^{(k)}$. Let $x = (x_1, x_2, x_3)$ be **the global co-ordinates of a point in ortdiogonal Cartesian co-ordinates.**

A surface denotes the interface area between two subregions or any surface upon which tractions and/or displacements are specified. The Surfaces of each subregion are divided into a number N of triangular or quadrilateral planar elements Δs _m over which tractions t_1 (m) and displacements **U**_i (m) are constant (1=1,2,3) (m=1,2...N), The **system of integral equations \diich governs tlie interaction between tractions and displacements is given (Ciruse, 1969) by**

$$
\frac{f(x)}{f(x)} = \sum_{n=1}^{M} u_i(n) \int_{\mathbb{R}^n} T_{x_i}^{(k)}(m,n) dS(n)
$$

=
$$
\sum_{n=1}^{M} t_i(n) \int u_{i_i}^{(m,n)} dS(n)
$$
 (1)

for the kth subregion.

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The tractions and **displacements** appear outside **the** integral signs in (¹) because of the **assumption** of constant tractions and **displacements.** over each element. In (1) element m is termed a receiving element while elements n are termal emitting elements, ie an emitting **element** n influences the **displacements and** tractions of a receiving **element m via the influence coefficients**

$$
\int\limits_{\Delta s_{\infty}}\mathcal{T}_{ij}\stackrel{\text{(A)}}{=} (m,n)\ d\mathcal{S}(n)
$$

$$
\int u_{ij}^2 (m,m) \, dS(m) \tag{2}
$$

The subscripts i and j relate the relevant components of traction or evaluates the integrals in (2) analytically. **It** is possible to evaluate these integrals numerically (except with $m = n$) with considerable time saving in most cases. The integrals in which $m = n$ are termed "element self effects" and are evaluated as described by Cruse (1969). Numerical evaluation of the integrals in (2) is discussed in more detail in Chapter 3, displacement in the global co-ordinate system. Cruse (1969)

Once the surface tractions and displacements are known the stresses $\sigma_{ij}^{\ldots}(\gamma)$ and displacements $u_i(\gamma)$ at other points y in the kth subregion are given (1) by

$$
u_i(y) = -\sum_{n=1}^{N} u_j(n) \Delta T_{ij}(y_m)
$$

+
$$
\sum_{n=1}^{N} T_{ij}(x) \Delta U_{ij}(y_m)
$$
 (3)

$$
\sigma_{ij}^{(k)}(y) = -\sum_{n=1}^{\infty} u_{\mathcal{A}}^{(k)}(x) \Delta S_{\mathcal{A}}(y, \eta)
$$

+ $\sum_{n=1}^{\infty} t_{\mathcal{A}}^{(k)}(x) \Delta D_{\mathcal{A}}(y, \eta)$ (4)

where $\Delta T_{ij}(y,n)$ and $\Delta U_{ij}(y,n)$ are the integrals in (2)

and
$$
\triangle S_{\ell,j}^{(k)}(\gamma,\eta) = \int_{\Delta S_{\ell}} S_{\ell,j}^{(k)}(\gamma,\eta) dS(\eta)
$$

$$
\Delta Da_{ij}^{(k)}(y,n) = \int_{\Delta S_n} D_{i,j}^{(k)}(y,n) dS(n)
$$

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The (k) superscript for the kth subregion is dropped henceforth for convenience where not necessary. The functions T,U,S and D are given (Cruse, 1969) by

 $\mathbf{S} = \begin{bmatrix} \mathbf{A}^T \\ \mathbf{A}^T \end{bmatrix}$

$$
T_{i,j}(m,n) = \frac{c}{4\pi r^{2}} \left[\frac{\partial r}{\partial n} \left(\delta_{ij} + \frac{3}{1-2D} r_{i} r_{ij} \right) - n_{j} r_{i} + n_{i} r_{j} \right]
$$

\n
$$
U_{i,j}(m,n) = \frac{1}{4\pi r^{2}} \left[\frac{3-4\nu}{4(1-\nu)} \delta_{ij} + \frac{1}{4(1-\nu)} r_{i} r_{j} \right]
$$

\n
$$
S_{4ij}(m,n) = \frac{c_{m}}{2\pi r^{2}} \left\{ 3\frac{\partial r}{\partial n} \left[\delta_{ij} r_{j} \left(1 + \frac{\nu}{1-2D} \left(\delta_{4i} r_{ij} + \delta_{4j} r_{j} \right) \right) \right.\right.
$$

\n
$$
= \frac{c_{m}}{1-2\nu} r_{i} r_{j} r_{i} \left[1 + \frac{3\nu}{1-2\nu} \left(n_{i} r_{j} r_{i} + n_{j} r_{i} r_{i} \right) \right]^{(6)}
$$

\n
$$
+ 3n_{4} r_{i} r_{i} r_{j} + n_{j} \delta_{4i} + n_{i} \delta_{4j} - \frac{1-4\nu}{1-2\nu} r_{i} \delta_{ij} \right\}
$$

\n
$$
D_{4i,j}(m,n) = \frac{c}{4\pi r^{2}} \left(\delta_{4i} r_{j} + \delta_{4j} r_{i} - \delta_{ij} r_{j} \right)
$$

where j_j is the Kronecker Delta Function

$$
\mathbf{r} = \left[\left(\mathbf{x}_t(\mathbf{w}) - \mathbf{x}_t(\mathbf{w}) \right) \left(\mathbf{x}_t(\mathbf{w}) - \mathbf{x}_t(\mathbf{w}) \right) \right]^T
$$
\n
$$
\mathbf{T} \mathbf{i} = \frac{\partial \mathbf{r}}{\partial \mathbf{x}} \left(\mathbf{x}_t(\mathbf{w}) - \mathbf{x}_t(\mathbf{w}) \right)
$$
\n
$$
= \frac{1}{T} \left(\mathbf{x}_t(\mathbf{w}) - \mathbf{x}_t(\mathbf{w}) \right)
$$
\n(8)

$$
C = (1 - 2 \nu) / 2 (1 - \nu)
$$

 $m_i = m_i(\omega)$ = outward unit normal vector to the emitting element

주장으로 이용할 수 있다. 그런 사람들이 아르치는 아이의 모임을 하면 없이 이번 이 사람 위에
2017년 - 대한민국의 대한민국의 학생들은 대한민국의 대통령을 하는 아이의 사람들이 있다.

μ = shear modulus

- Hie equations (1), (3) and (4) form the **basis** of the first programme BBM of this dissertation. U₁ (n) or t_1 (n) is specified for each **element for 1 = 1,2,3 except where the element represents an** interface to another subregion. The boundary conditions at interface **elements (between subregions (k) and (ktl) say) are**

$$
u_{\pm} (k) (n) = u_{\pm} (k+1) (m)
$$
\n
$$
t_{\pm} (k) (n) = -t_{\pm} (k+1) (m)
$$
\n(10)

unless failure of the interface (occurs. A mechanism for allowing failure to occur is discussed in Chapter 5. The equations (1), (3), and (4) are further modified to include the "lumping **mechanism",** various symmetry conditions and the above interface elements.

2 Displacement discontinuity formulation

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Consider now a plane, relatively thin excavation **which** is treated here as a single plane surface of negligible thickness. This plane may be divided up into a mesh or grid of square displacement discontinuity **elements. A** particular element is denoted by its row and column number in the grid (Fig 2.1), ie element ij lies in the ith row and jth column of the grid. Define a local coordinate system for this grid (Fig 2.1) so that the x- and y-axes point in the directions of increasing row **and** column numbers **respectively. Let the x-y plane be separated into two surfaces within the** mesh the top surface (outward normal points down) being denoted by the + superscript and the **bottom** surface by a-. A displacement discontinuity arises **when** these **two** surfaces move relative to one another. If, as before, it is assumed that displacements and surface tractions are constant over each element then constant displacement **discontinuity components may be defined for each element ij by**

$$
d_{\alpha}(t, j) = u_{\alpha}(t, j) - u_{\alpha}(t, j) \qquad (\alpha = i, 2, 3) \qquad (11)
$$

The surface tractions acting on the + and - surfaces of any displacement discontinuity element have equal magnitude but opposite sign. an element. The sign conventions adopted are positive normal stresses denoting compression and the normal displacement discontin**uity is 'positive if: the + and - surfaces move towards one another (as** It is more coveniant therefore to consider stresses acting "within" **is normally the ease under theaction of a compressive stress field).**

The normal and shear stresses acting on a displacement discontinuity element if are given by
 σ_{13} (1, j)

-
- σ ₂₃ (i,j)
- $0.33 (i,j)$

in the local co-ordinate system chosen. Stanfield and Grouch (1973) give equations relating these normal and shear stresses to the normal and shear displacement discontinuity components

$$
\sigma_{\varsigma_{\beta}}(t, j) = \sum_{k=1}^{M} \sum_{\ell=1}^{N} K_{\varsigma_{\beta}}(t, j, \Delta, L) d_{\beta}(L, L)
$$
\n
$$
(\alpha, \beta = 1, 2, 3)
$$
\n(12)

earlier. These coefficients K may be evaluated in closed form (Starfield and Crouch, 1973) for square displacement discontinuity **elements lying in the same plane and may be expressed in terms of row** and column differences i-k and j-1. where **there are** M rows and **columns** in **the** grid - and K denotes an influence coefficient similar to the T coefficients described

The stresses and displacements at points y outside the grid are given (see Appendix 1) by

$$
u_{\kappa}(y) = -\sum_{k=1}^{M} \sum_{\ell=1}^{M} d_{\beta}(\ell, \ell) \triangle T_{\kappa \beta}(\gamma, \ell, \ell)
$$
 (13a)

$$
\sigma_{\alpha,\beta}(\gamma) = \sum_{\Delta=1}^{M} \sum_{\ell=1}^{M} d_{\delta}(\mathbf{A}, \ell) \Delta S_{\delta \kappa \beta}(\gamma, \mathbf{A}, \ell) \qquad (13b)
$$

The form of $(13a)$ and $(13b)$ is very similar to that of (3) and (4) . The coefficients T (y,k,l) and $U(y,k,l)$ are evaluated numerically using the functions given in (6). The additional index in these terms is. used purely to indicate the row and column of a. displacement discontinuity element as opposed to an element number for boundary elements. The surface tractions acting upon a displacement discontinuity element do not affect the stresses and displacements elsewhere in the body so that the U and D terms in (3) and (4) are not present in (13a) and (13b). **When** evaluating the T and S functions for a displacement discontinuity element, the outward normal of the bottom (-) surface is chosen in keeping with the definition of $d(i,j)$ in (11) .

Equations for mixed boundary element method

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In order, to derive the equations for the mixed boundary element method, it is necessary first. to convert, the co-ordinates, tractions, displacements and normal vectors of the boundary elements to the local co-ordinate system of the displacement discontinuity elements. The necessary transformations are '

$$
u_{\alpha} = \ell_{\alpha \pm} u_{\pm}
$$

\n
$$
x_{\alpha} = \ell_{\alpha \pm} (x_{\pm} - x_{\pm}')
$$

\n
$$
n_{\alpha} = \ell_{\alpha \pm} n_{\pm}
$$
 etc (14)

where $\ell_{\alpha,i}$ are the direction cosines of the local with respect to the global co-ordinate system,

and x'_1 is the origin of the local with respect to the global co-ordinate system.

Once these quantities have been evaluated, there is no further need to consider the global co-ordinate system and the i,j and k subscripts of equations (1) to (9) are merely replaced by Greek subscripts \forall , β , γ etc. (This avoids confusion with the i, j, k and 1 values for rows and columns).

The equations (1) , (3) and (4) are written for a tension positive stress convention While the corresponding equations for the displacement discontinuity elements are written for a compression positive convention* The equations which **follow** take this into account and -- re written for the latter convention.

Let an entire displacement discontinuity grid 'be placed within the first subregion of boundary elements. Stresses and displacements at points y inside this subregion are given by a summation of equations (3) and (4) with $(13a)$ and $(13b)$ -

$$
u_{\alpha}(y) = -\sum_{n=1}^{\infty} \left[u_{\beta}(n) \Delta \prod_{\alpha_{\beta}} (y_{i}n) + \mathcal{F}_{\beta} \Delta U_{\alpha_{\beta}} (y_{i}n) \right]
$$
\n
$$
-\sum_{k=1}^{\infty} \sum_{\ell=1}^{n} d_{\beta}(\beta, L) \Delta_{\alpha_{\beta}} (y_{j} \Delta, L)
$$
\n(15)

$$
\sigma_{\kappa_{\beta}}(y) = \sum_{n=1}^{\infty} \left[u_{\gamma}(n) \Delta_{\sigma_{\kappa_{\beta}}}(y, y) - t_{\kappa}(n) \Delta D_{\gamma_{\kappa_{\beta}}}(y, n) \right]
$$

+
$$
\sum_{\delta=1}^{M} \sum_{\ell=1}^{N} d_{\gamma}(A, L) \Delta S_{\gamma_{\kappa_{\beta}}}(y, A, L)
$$
 (16)

Similarly, the displacements induced at centroids of boundary elements by displacement discontinuity elenents must be included in equation (1) and stresses $({\sigma}_{13},$ ${\sigma}_{23}$ and ${\sigma}_{33}$ only) induced at displacement discontinuity elements by boundary **elements** must be included in equation (11) giving

$$
\frac{1}{2} u_{\alpha}(x) + \sum_{n=1}^{N} u_{\beta}(x) \Delta T_{\alpha\beta}(m, x) + \sum_{n=1}^{N} \sum_{\ell=1}^{M} d_{\beta}(\ell, l) \Delta T_{\alpha\beta}(m, \ell, \ell)
$$
\n(17)

$$
=\sum_{n=1}^{\infty}t_{\alpha}\Delta U_{\alpha\beta}(m,n)
$$

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for boundary elements and

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$$
\sigma_{\alpha_{2}}(t_{1}) = \sum_{\beta=1}^{M} \sum_{\ell=1}^{M} K_{\alpha_{\beta}}(t_{1}, \beta, \ell) d_{\beta}(\beta, \ell)
$$
\n
$$
+ \sum_{\alpha=1}^{M} u_{\alpha}(\alpha) \Delta_{\gamma \alpha}(\gamma, \alpha) - \sum_{\alpha=1}^{M} \sum_{\alpha} (u_{\alpha}) \Delta_{\gamma \alpha}(\gamma, \alpha)
$$
\n(18)

17

for displacement discontinuity elements

The equations (15) to (18) form the basis of the mixed boundary element programme MBEM. For each element, the stresses or displacements or tractions are specified in (17) and (18) in the x, y and *z* directions and a linear system of equations results. These equations are then solved iteratively using the method of successive over relaxation (SOR) for the unknown displacements, tractions and displacement discontinuities.

The following points are worth noting about these equations:

- (1) They do not include the effects of lumping
- (2) Tractions and displacements are both not known a priori at interfaces between subregions but may be found iteratively as described in the fifth chapter.
- (3) The numbers of coefficients T, U etc calculated in or used by equations (17) and (18) are

Since the coefficients $K(i,j,k,l)$ in (18) depend only upon the differences $1 - k$ and $1 - 1$, and since some of the coefficients may be collected into the vector of known boundary conditions, the number of coefficients **which** must be stored. by the computer in the absence of **lumping is at best**

$$
(9N^2 + 9 M^2N) + 9 M^2 + 3 M^2 \tag{19}
$$

while the number of degrees of freedom in the system is

 $3(N + M^2)$

For practical problems, $N \geq 150$ while $M \geq 20$ so that excessive amounts of storage are required. The need, for a means of reducing storage requirements is evident.

.4 Discussion of mixed boundary **element** method equations

Much attention is currently being given to "hybrid" or mixed stress analysis techniques. Zienkiewicz (1979) gives a comprehensive summary of techniques currently in use for combining finite element and boundary element formulations. Each element type, is used to model that part of the problem to Which it is best suited. Grouch (1976) has demonstrated hew, in two-dimensional problems, the **displacement** discontinuity **method** nay model **both** crack or fault type problems as well as open cavity problems, This formulation uses equations similar to (12). It is seen from equation (12) that calculation of "stress" influence coefficients is required as compared with "displacement" influence coefficients (3) required for a boundary **element** formulation. Use of the latter type of influence coefficient for open cavity type problems is to be preferred for the **following** reasons:

(i) If influence coefficients are being evaluated using numerical integration (no closed form solution to the integral $K_{\omega, \rho}$ (i, j,i, j) in (12) for an arbitrary quadrilateral or triangle was found in the literature) then the time required to evaluate "displacement" coefficients is significantly less than that required for **the** equivalent stress coefficients. In addition, all integrations may be done with respect to a single co-ordinate system whereas the "stress" coefficients have to be transformed to the local co-ordinate system of each element if this does not coincide with the global co-ordinate system.

(ii) Grouch (1976) shows how, when dealing with an open cavity, the displacement discontinuity formulation produces an interior and an exterior region. Some problems, arise with the interior region if no restraints are rnade to prevent rigid body motion. No such problems arise with the boundary elements since there is not more than one region under consideration.

Conversely, numerous **problems** arise when attempts are rnade to use boundary elements to model a tabular excavation or a crack type problem.

It is logical therefore, to match element types to the problem. Moreover, since most tabular excavations are nearly planar it is economic to model such an excavation with a regular grid of Square displacement discontinuity elements. **An** open pit or open excavation is likely to have an irregular shape necessitating the use of triangular or quadrilateral elements.

Equations (15) to (18) are derived for the class of problem in which tabular and open excavations are present. This is a class of problems **which** arises fairly frequently in mining rock mechanics.

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CHAPTER 3 NUMERICAL INTEGRATION OF INFLUENCE COEFFICIENTS

The follcwing Integrals or influence coefficients have to be evaluated numerically in the boundary element or mixed boundary element formulations:

 Δ $T_{\leq B}$ (y,n) Δ U_{sep} (y,n) Δ S_{rd} (y,n) $\Delta D_{\text{y} \alpha \beta}(\text{y},\text{n})$

The method of integrating **these** functions over a flat triangle is **the** same for each function even if the resulting accuracies differ slightly. It is therefore only necessary to describe the evaluation of $T(y,n)$. Three separate cases exist:

- (i) Triangular boundary elements
- (ii) Guadrilaterial boundary elements
- (iii) Square displacement discontinuity elements

3.1 Boundary elements

The quadrilateral elements are simply divided into two triangular elements which are then treated separately. The function $T(y,n)$ varies over the surface of the nth triangle, (an emitter triangle). The rate of variation depends primarily **on** the distance separating the emitting triangle n from the receiving point y and the size of the element and to a lesser extent upon the orientation, and shape of the emitting element.

The method of evaluating the integral is equivalent to first estimating an average value of the function over the element and then multiplying this value by the area of **the** triangle.

If a is the area of the emitting triangle and r the distance to the point y then a measure of the variation of the function T over the triangle is given by the ratio R.

$$
\mathbf{R} = \mathbf{r}^2/\mathbf{a}
$$

(See Fig 3.1)

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In each case R is evaluated and the number of points at which T is to be evaluated over the triangle in order to give sufficient accuracy is determined. The options are 1, 3, 7, 21 and 42 points. The points at which the function is to be evaluated are given in Table 3.1 for the 1, 3 and 7 point cases Zienkiewicz (1971)

TABLE 3.1 : CO-ORDINATES AND WEIGHTS FOR 1, 3 AND 7 POINT

A point x say, within triangle n, at which the function T is to be evaluated is given by

$$
\times_{\kappa} (k_{\kappa}) = \frac{c}{4} (k) \times_{\kappa} (\kappa) + \frac{c}{4} (k) \times_{\kappa} (\kappa) + \frac{c}{4} (k) \times_{\kappa} (\kappa)
$$

$$
= \frac{c}{4} (k) \times_{\kappa} (\kappa) \qquad \qquad \kappa = 1, 2, 3 \qquad (21)
$$

are the co-ordinates of the node β of triangle n and $\oint_A (k)$ are the $\cdot 1$). point, (Table 3.1). θ
where x_{α}

(22)

Thus

$$
\Delta T_{\alpha\beta}(y,\eta) = \alpha \sum_{\beta=1}^{T} w_{\beta} T_{\alpha\beta}(y, x(\beta_{\alpha}))
$$

where $I = 1,3$ or 7

Mien 7 point integration is inadequate (ie value of R too small), the triangle is subdivided further into three or six smaller triangles, each of equal area and the 7 point formula is applied in turn to each of these. The nodes of the smaller triangles are merely the nodes or centroids of the original triangle or the midpoints of its sides, $(Fig 3.1)$

The quadrature algorithm may be **summarized** as **follows:-**

- (1) If element n is quadrilaterial, divide into two triangular elements, n and n' say
- (2) Calculate centroid of triangle n $x^{\circ}(n)$ (or $x^{\circ}(n')$)
- (3) Calculate $r^2 = (y_i x_i^o(n))(y_i x_i^o(n))$
- (4) Calculate $R = r^2/a$ and decide on the required accuracy 1,3 or *7* point etc (a = triangle area)
- (5) Evaluate the outward normal to triangle $n = n(n)$
- (⁶) Subdivide triangle n into **1,3** or 6 subtriangles and calculate additional nodal co-ordinates for the subtriangles (m) if **necessary**
- (7) There are 1,3 or 7 sample points **within** each subtriangle (m)

For each sample point k inside subtriangle m:

(a) evaluate its co-ordinate x_{α} (k_m) from (21)

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- (b) evaluate r, r_{jet} and $\frac{dr}{r}$ from (7), (8) and (9)
- **dn** (c) evaluate the functions $\mathbb{T}_{\alpha,\beta}(y,x(k_m))$, $\mathbb{U}_{\alpha,\beta}(y,x(k_m))$ etc as **required**
- (d) continue evaluation of Δ $T_{\alpha\beta}(y,n)$ etc from (22)

Although the different functions T, t), S and D are inversely proportional to r, r^2 or r^3 , it is convenient from a programming point of view to evaluate them together as described above.

There are several secondary benefits arising from this point Integra-' tion scheme. A continuous check (31 the ratio R at each integrand point enables errors in the data input to be easily detected.

Fig 3.2 shows a common example in which α 's node of an element is **incorrectly specified. Such errors are easily overlooked when checking the data manually since the area, outward normal and position of the element may ail be correct. If the ratio R drops below some threshold value R^ln* say, during the integration procedure, the error is easily detected. If elements are so close that a 42 point integration formula is unreliable, then it is highly probable that a bad choice of element sizes has been made and that the iterative solution.would converge very slowly or not at all.**

When lumping is implemented, it is possible, by using only 1 point integration, to quickly assess the amount of storage which Will be required. If the maximum available storage is exceeded, then a coarser lumping mechanism may be adopted without wasting too much time,

3.2 Displacement discontinuity elements

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The displacement discontinuity elements are all square so that inple**mentation of a Gauss Quadrature focmila is simple, _ accurate and efficient. For each element the ratio R is evaluated as before and 1, 4, 9 or 64 point formulae are selected accordingly. (22) is rewritten for displacement dis continuity element ij as -**

" ; ;

$$
\Delta T_{\alpha\beta}(y, i_{1j}) = \alpha \sum_{\alpha=1}^{m} w_{\beta} T_{\alpha\beta}(y, x(\beta, i, j))
$$
 (23)

The use of square elements enables the integrand points x(k,i,i) to be evaluated efficiently in terms of the element oentoid and element half width. The Gauss Quadrature coefficients were taken from Zienkiewicz (1971). Also since **it** is known that the outward normal **to** all **displacement** discontinuity elements- is (⁰ ,⁰ ,¹), considerable simplifications may be made to the functions T and S.

3.3 <u>Discussion</u>

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Briefly, the advantanges of the numerical integration may be **summarized as follows:**

- A comprehensive check cn the input data is **made** avaiable -
- **For** most problems, the numerical ; tegration is quicker than analytic integration. For some geometries, this might not be **true, however**

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- **(iii)** It is not necessary to evaluate or invert the Jacobian matrix **at every Integration point**
- **& trade off between accuracy and. execution time is available.** This was found to **be** very useful in the development stage of **the programmes.**

It should also be noted that the element self effects (for Which r=o) are evaluated analytically due to the presence of the l/r singularities. The displacement discontinuity coefficients K (18) are also evaluated analytically facilitating the use of a recurrence formula, described by Starfield and Croudh (1973).
CHAPTER 4: DESCRIPTION OF LUMPING MECHANISM

The- lumping mechanism described here is essentially an extension of the numerical quadrature procedure described in. Chapter 3. tvhen emitting, elements are remote from receiving elements, then the functions

> $T_{\alpha, \beta}$ (y,n) $U_{\alpha\beta}$ (y,n) $S_{\gamma\alpha\beta}(y,n)$ $D_{\gamma\alpha\beta}(y,n)$

vary slowy over the emitting elements. When this occurs, the coefficients for a number of elements may be σ 'wire? into a single lump coefficient. The *jumping* mechanism has alward been put to great use in existing boundary and displacement discontinuity formulations (Starfield and Crouch (1973) and Deist e ^ Georgiadis (1976)) but **tliese** schemes differ somewhat from the scheme * tlined below.

4.1 Boundary: element lumping

When a mesh of boundary elements is being drawn up, the user is required to group elements with similar orientation, size and location into "lump elements" containing from one to twelve boundary elements. (The extra effort required to do this is more than off-set by the additional error checks which become available). Consider two Imp elements with **4** boundary **elements** in each (Fig 4.1) . let the "receiver" lump contain 4 potential receiving elements and the "emitter" lump 4 potential emitter elements. In the absence of any Imping mechanism, 4x4 = 16 sets of coefficients have to be calculated (there are 18 coefficients in each set). If the 4 emitting elements are grouped together then each receiving **element** requires a different set of coefficients, ie. 4 **tz^ts** are required.. If **the** receiving elements are grouped together then only 1 coefficient set is required.

The three types of coefficient set are referred to as element-element, lump-element and Imp-lump coefficients respectively

(See Fig 4.2). If lumps are chosen to be nearly planar or planar then the centroids, areas and outward normals of the lump elements may be calculated just as for normal elements.

In deciding Which coefficient type to use, the ratio

$$
R = \frac{r^2}{a}
$$
 is used

as before where $r =$ distance separating the lump centroids a = area of emitting lump

The potential time and storage savings of this lumping scheme improve **as the number of elements increase. If boundary elements alone are** considered, then the number of coefficients required for an N element problem in the absence of lumping is

18

If the average number of elements per lump is 6, say, then the number **of lumps is**

$$
M \approx N/6
$$

and if approximately 40 element-element coefficients are required per element, then the approximate number of coefficients required with **lumping** is

$$
18 \text{ N} \left[\frac{\text{N}}{\text{6}} + 40 + \frac{\text{N}}{36} \right]
$$

The storage **and** time-saving factor for **N** = 300 is therefore about 3.

The lump elements are treated just as ordinary elements, and lump displacements, tractions, normals and areas are calculated as averages weighted with element areas. The lumping mechanism is directly applicable to the evaluation of interior stresses and displacements. At present all lump coefficients are evaluated by one point integral formulae, but it is expected that greater savings would be obtained by using higher order formulae for these coefficients, since emitting lump elements could effectively be brought closer to receiving elements, thus further reducing the total number of coefficients.

.2 Displacement discontinuity lumping

Two separate schemes are adopted for evaluation of element-element interactions and for evaluation of stresses and displacements at, interior points. The former scheme is based on that of Starfield and Grouch (1973) while the latter is essentially **that** described above.

For evaluation of element-element coefficients, groups of 1,4, 9 or 25 displacement discontinuity elements are grouped into square **lump elements** and **lump-lump** or element-elernent coefficients only are calculated. The need for the lump-element coefficients described above is obviated apparently because the relevant integrals are evaluated in closed*form, **not** numerically-

3 Reduction of storage requirements

While the lumping scheme described above was initially implemented to reduce disk storage requirements and execution time, a number of other benefits also result, the most important being a reduction in core storage requirements.

In the absence of lumping / **it** is expedient to hold in main memory the following arrays.

nodal co-ordinates element areas

element displacements element tractions element-direction cosines **element centroids element-node numbering element codes**

Once Imping is introduced it becomes necessary to keep track of the integration scheme (lunp^lump, lunp-elsment or element-elament) used for each lump element. If there are M lumps, then this array is of dimension *lxM.

Once interface elements are introduced, it is necesary also to store for each lump element information such as cohesion and angle, of friction (Chapter 5) as well as the direction cosines of a local **Co-ordinate system for each element.**

Before **implementation** of the storage reduction scheme, it was found that core storage limited the maximum number of elements to about 500. With 500 **elements** however, execution time was increased since it was easier to calculate element centroids and direction cosines as required rather than store them permanently.

Since elements are always accessed through their parent lump element, it is possible to retain in main memory element properties only for those lumps under **consideration**« For example, a receiver lump and an emitting lump element are retained in main memory during calculation of influence coefficients, to reduce the number of **disk** transfers required to implement this scheme, all of **the** lump element arrays (lump areas, displacements etc) are stored in main memory. Main memory requirements are then restricted by the number of lump elements, rather than ordinary elements.

The element properties for any lump element are stored on disk using labelled **common** arrays (standard for Ascii Fortran **IV),** This enables the use of a direct disk access routine reducing further the disk access times.

4.4 Lumping - a brief discussion

The lumping system described here is designed specifically for the boundary integral type of equation. It is ideal for systems **requiring disk or tape storage because of the sequential access of data, Tt is also ideal for systems of equations which, are diagonally dominant and hence well suited to iterative solution techniques, The lump variables (displacements and tractions) are calculated as weighted averages of their constituent elements. As such it is possible to implement this lumping mechanism into systems of equations which are being aabwxl using an elimination rather than an iterative technique. . The total number of degrees of freedom of the system would be increased by about 10 percent. This new system of** equations would also be about 15 to 50 percent populated but unfortunately would not be a banded system. Special elimination techniques which minimize the amount of "fill in" (zero coefficients which **become non-zero in the elimination process) would be required. In** addition, the simple sequential access of coefficients used in the iterative solution is not applicable to elimination schemes. Disk or coefficient access tends **to be** more random. Finally, elimination schemes are not well suited to the modelling of non-linear behaviour **which occurs When failure of material interfaces is initiated.**

Gaussian elimination may be compared with successive over relaxation (SOE) for a problem of 1 000 **elements** or 3 000 degrees of freedom as follows:

Gaussian Elimination (SOR)

Mumber of coefficients 9x10⁶ 2x10⁶ adds and multiplies ie $9x10^9$ ie \pm $6x10^7$

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Angelska († 1705)
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Number of arithmetic $1/3$ \mathbb{N}^3 2x10⁶x2 per iteration

For such **a problem,** the iterative solution is up to ISO times more efficient than elimination without the lumping mechanisms.

The. lumping scheme, as **implemented** in this dissertation is applied to the simplest boundary element type, namely the constant displacement/traction element. **Although** not an express aim of this dissertation, it is felt that lump elements provides a reasonable alterna**tive to the more sophisticated element types (quadratic and cubic** variation of unknowns over each element) of Xachat and Watson **(1976).**

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Alternatively, a marked improvement in the performance of these higher order elements could be expected if they could be lumped.

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CHAPTER 5. DESCRIPTION OF BOUNDARY INTERFACE ELEMEOtS AND SYMMETRY CONDITIONS

Consider the simple problem shown in Figure 5.1 of two subregions each containing 2 elements (after Lachat and Watson (1976)). Each element is a **schematic representation of a number of simple planar elements which would constitute each subregion. The boundary conditions are such that trac-» tions are specified at elements 1 and 4 While stresses and displacements are continuous across the interface between the two subregions. Assume** further, for the moment, that the problem only has displacements and tractions in one dimension. Equation (!) may be written in matrix form for the problem as -

or by using subscripts for the matrix coefficients:

The zero coefficients arise because there is no direct interaction between the two subregions other than the displacement and traction boundary conditions at the interface. For this problem these may be written as:-

$$
u_2 = u_3
$$

$$
t_2 = -t_3
$$

(25)

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By sorting known and unknown quantities to the left and righthand sides respectively and applying (25) to (24), (24) may be rewritten as -

where superscripts denote the subregion

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It has been stated above that the equations (1) or (26) are solved using an iterative scheme (Successive over relaxation). A sufficient, but not necessary, condition for convergence of this scheme is that the system matrix is diagonally dominant (Froberg 1970). Experience has shown that it is only necessary to maintain an approximate degree of diagonal dominance in the system matrix. (The rate of convergence gradually decreases as diagonal dominance decreases), New the magnitudes of the element self effects T_{11} and U_{11} are approximately

 (27)

 $\mathbb{T}\begin{matrix} \text{(k)}\ \mathbb{T}\ \text{11}\ \text{22}\ \end{matrix}\approx\frac{1}{2}$ $v\frac{(k)}{11}$ $\approx \frac{1}{G(k)}$

where $G(k)$ is the shear modulus for the kth subregion. Also, within any subregion for a well posed problem

Let the coefficients $T_{i,j}(k)$ and $U_{i,j}(k)$ it) be denoted by δ and substitute (27) into (26). Then

In (28) approximate diagonal dominance may be obtained by scaling the shear moduli $_G^{(1)}$ or $_G^{(2)}$ if they are approximately equal.

Assume firstly that

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$$
G(2) = 100 G(1)
$$

$$
G(1) = 2
$$

The system matrix in (28) becomes

It is seen that the third row is definitely not diagonally dominant while the second row is almost diagnoally dominant. Improved diagonal dominance **may be obtained in (29) however by swapping the second and third rows or** by a choice of shear moduli so that

(29)

 $G(2) < < G(1)$

Since the programming of row swapping is inconvenient and since it is not generally possible to choose suitable shear moduli the following iterative procedure has been adopted for subregions with very different elastic properties.

- (1) For the softer subregion, estimate tractions at the interface.
- (2) Use these tractions as specified boundary conditions for the stiffer subregion.
- (3) For the stiffer subregion, estimate displacements at the interface.
- (4) Use these displacements as specified boundary conditions for the softer subregion.

This iterative cycle is easily included **in** the overall iterative solution.

Intuitively, large displacements **in** a soft material produce small stresses while large stresses in a stiff material produce small displacements. The diagonal dominance is therefore interpreted as a large cause producing a **small** effect rather than vice versa.

A somewhat, unfortunate consequence of this limitation of allowable boundary conditions is that it is not possible to model a stiff subregion completely enclosed by softer subregions because there is then no restriction of rigid body displacement in the stiff subregion

5.1 Failure at an interface

It is possible at any stage during the iterative solution (for the tractions and displacements) to calculate- the normal and shear displacements and tractions at any element. Lachat and Watson (1976) show how the equations (1) can be rewritten to give tractions and displacements in a local co-ordinate system for each element. This **represents a large amount of additional calculation and an alterna**tive approach is to transform tractions and displacements to some local co-ordinate system only when required. Consider the **problem** of **Fig 5.1 again. Let a local co-ordinate system for any element be** defined so that the **2-axis** is the outward normal and y-axis is horizontal. Let the direction cosines of this "elemental" local **co-ordinate system with respect to the oo-ordinate system of the dis**placement discontinuity grid be $\ell_{\mathcal{A}}$ and the local displacements and i i */ t* tractions be t χ and $u \chi$ respectively. (tg and u3 are then **normal tractions and displacements).**

Then

and

 $t'_{\alpha\beta} = \ell_{\alpha\beta} t_{\beta}$ $u'_{\alpha} = \ell_{\alpha\beta} u_{\beta}$ $t_{\beta} = -\ell \kappa_{\beta} - t_{\alpha}$ $u_{\beta} = -\ell \alpha \beta - u_{\alpha}$

(30)

The boundary conditions for interface elements are: الواردة كالكالم الكامية tractions specified for stiffer elements displacements specified for softer elements

A Mohr Coulomb failure criterion is implemented in the iterative solution as follows, Let the cohesion and angle of friction of the interface be c and ϕ respectively. Then the shear strength $\sigma_{\rm s}$ of the interface is given by

$$
\mathbf{r}_s = \mathbf{c} + \mathbf{c} \quad \text{tan } \phi \tag{31}
$$

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where $\sigma_n = -t_3' + p_3$ = Total normal stress P3 = primitive normal stress

The **maximum** total shear stress component is given by

$$
\mathcal{L}_{\text{max}} = \sqrt{(-t_1 + p_1)^2 + (-t_2 + p_2)^2}
$$
 (32)

If $\chi_{\text{max}} \geq \sigma$ s then failure occurs. If σ_n is tensile when failure does occur tlien the node of failure is also tensile. If not, then the failure node is in shear.

5.1.1 Shear failure

Shear failure is inplenented singly as a change of boundary conditions for the interface elements. Continuity of normal displacements and stresses must be maintained, but the shear displacements are unknown for both interface elements..

Ihe shear tractions are given by

$$
t_1'_{new} = t_1'_{old} \times \frac{\sigma_s}{\tau_{max}} + p_1
$$
\n
$$
t_2'_{new} = t_2'_{old} \times \frac{\sigma_s}{\tau_{max}} + p_2
$$
\n(34)

5.1.2 Tensile failure

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If tensile failure occurs at an interface, then the newly **created void becomes indistinguishable from an open excava-^ tion. Tractions (equal but opposite in sign) must then be specified at each interface element so that the total resultant tractions (primitive and induced) at these elements are** $zero, i.e set t_1' = -p_1'$.

These Updated tractions and displacements are then transformed back to the global co-ordinate system and tlie process continued. Other Ix^undary conditions may also be implemented at interfaces. These liave been described in detail by Crouch (1976) and Starfield and Crouch (1973). Essentially an interface may or may not have a filling or the interface may be treated as part of a tabular excavation. If the interface **lias no filling it may still fail in shear or tension. If the** interface has a filling then the relative displacement of the **interface surfaces is controlled by the stiffness of the fill** unless failure occurs. If **the** interface is mined or open then **convergence or separation of tire surfaces occurs but a .limit: to the maximum amount of convergence may be specified. Inter face elements are therefore assigned different codes to distinguish their different properties.**

- Code **7** Interface element with no **infill**
- Code 9 Mined or open with a limit on maximum convergence
- **Code 10 Tensile failed element**
- **Code ll Shear failed element**
- **Code 12 Interface element with infill**
- Code 1 Open element with no limit on maximum convergence

These elements are distinguished from other elements which do not belong to interfaces by their codes. Codes for the other elements are;

- Code ¹ Open or mined element (tractions specified)
- Code 2 Zero displacement or fixed element
- Code 3 Element fixed **in** x direction
- **Code** 4 Element fixed in y direction
- Code 5 Element fixed in a z direction
- **Code** ⁶ **Any other specified mixture of boundary conditions**
- Code ⁸ Represents the earth's surface (**z-co-ordinate** = 0)

5.2 Symmetry conditions

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Symmetry conditions are easily incorporated into **the** boundary element programme but have not as yet been incorporated into the displacement **discontinuity elements of the mixed boundary element programme.** Fig 5.2 shows a two-dimensional **example** containing 4 subregions in which there are two planes of symmetry, XSYM and YSYM. Subregion 3 is entirely contained within subregion 1. It is therefore not directly affected by its symmetry **images** in the other three quadrants and therefore does not have any symmetry in itself. It is necessary, therefore, to assign separate symmetry conditions to each subregion. The following codes and symmetry types are catered for:

in the example in **Fig 5.2** the following symmetry codes would apply:

Fig 5.3 shows an example with x and y symmetry. Each of the three images has to be treated separately since for the x image, only x-components of traction, displacement, position etc change Wile **only y-components are affected in the y-image and so on. This is done in the programme by means of two arrays. The first 8x& array relates different symmetry images to the symmetry code While the** second array relates a particular image (x,y or xy etc) to the compo**nents of traction, displacement etc that depend upon that image** (Table 5.2). In Tables 5.1 and 5.2 , 1 denotes "yes" and 0 denotes $"$ no".

	Symmetry Code	
Synnetry Image		
Object		
12		
IΧ		
xz		

TABLE 5.1: SYMMETRY IMAGE - SYMMETRY CODE TABLE (1=Yes, 0=No)

TABLE $5.2:$ SYMMETRY IMAGE - x, y, z COMPONENT TABLE (1=Yes, 0=No)

y% 0 0 0 0 0 0 1 1 xyz 0 0 0 0 0 0 0 1

Nb detailed description of symmetry *as found Jin the literature but it **is expected** that this algorithm for **the implementation** of symmetry is possibly novel (ie different codes for different subregions).

CHAPTER 6 MODIFICATION OF PROGRAMME MlNAP FOR NON-HOMOGENEOUS PROBLEMS

The application of the displacement discontinuity method to mining problems in two dimensions has been well demonstrated by Crouch (1976) with his programme MlNAP. A restriction of this programme is that it cannot model mining problems in non-homogeneous ground. The programme **allows** for specification of mixed boundary conditions which makes the incorporation of **non-homogeneous** subregions into the programme relatively easy.

The basic equations for the two-dimensional displacement discontinuity method have been given, Crouch (1976), for a problem containing N elements .(no tensor rotation for summation here).

 (35)

 (36)

Where stresses are specified as boundary conditions

and j represents an emitting element

- **i represents a receiving element**
- s represents a shear effect
- **n represents a normal effect**
	- are induced stresses
- d **are displacement** discontinuities
- A are stress **influence** coefficients derived by Crouch (1976)

where displacements are specified as boundary conditions,

where B arc displacement influence coefficients

u are displacements on **one** or **other** of the two displacement discontinuity surfaces.

زنه For example, B gives the shear displacement induced at element i by the sn normal displacement discontinuity component of element j.

Consider now the hypothetical problem of Fig 5.1 discussed also in Chapter 5. Stresses are specified at elements 1 and 4, but neither stresses nor displacements are known at elements 2 and 3. Equations (35) and (36) may be written for this problem as

 $\frac{1}{\sigma}$

 $\overline{2}$ \mathbf{C}^{\star}

 \mathbf{z} \mathbf{u}

 $\mathbf{3}$ σ

 $\overline{3}$ $\mathbf u$

 $\frac{4}{5}$

where A, B are submatrics given by

 (37)

Applying the boundary conditions for an interface (10) to (37) gives

(38) has **the same** form as (26) with T replaced by B and U replaced by \mathbf{A}

Ihe equations (38) are not diagonally dominant in general and so the solution of (38) using an interative scheme is also not always possible* These equations may also not be solved by any elimination schemes because of the non-linearities introduced by the fault **elements** or total closure restrictions essential for most mining applications. As the magnitudes

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of the B and B terms are always equal (they only depend upon the element **sise and orientation, approximate diagonal dominance may be achieved if**

22 *33 33 33 33 33 33 33* **the A terms are greater the A terns. .As with the boundary elements,** this is achieved in practice by incorporation of the following **algorithm** into the interative solution for the displacement discontinuity components.

- **(1) Fbr the softer subregion, estimate normal and shear stresses at the** interface.
- (2) Use these stresses as specified boundary conditions for the stiffer subregion,
- **(3) fbr the stiffsr subregion, estimate normal and shear displacements at** the interface.
- (4) the these displacements as specified boundary conditions for the **softer subregion..**

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Crouch (1976) describes how the displacement discontinuity element may be used to wodel solid or mined seam elements, mined seam elements which have whsequently been back-filled or fault elements which have failed in shear or tension. These features may be incorporated into the interfaces Jescribed above in the same manner as described by Crouch. This is not discussed further hare.

I"- Men beau &aand that the gate of convergetice of the equations in (38) is nojut half that for Ixamcgeneoua problems aryl that an over-relaxation factor preater than about 1,15 tends to diverge. It is also not possible to use **this alogwlthm fbr nonrhcmogaaeous problems in Which, a stiffer subregion is completely enclosed within & softer aUbregion,**

The elterations required to the programme MINAP are minimal and a wide range of non-honogenous problems may be solved without violating the **rnuhricfion imposed above.**

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CHAPTER 7 PROGRAMME VALIDATION AND NUMERICAL ACCURACY

7.1 Programme BEM

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The BEM programme was tested using a homogeneous unit cube under uniaxial tension. The following tests were run:

- 12 Triangular Elements
- 24 Triangular Elements
- 24 Square Elements
- 96 Triangular Elements

The case with 12 triangular elements was the same as that used by Cruse (1969) (See Fig 7.1). Results from these tests are summarized in Table 7.1. For the 12 triangular element test, a combination of 21 and 42 point integration formulae were used and the results of Cruse (1969) are given for comparison of the numerical and analytical integration of the influence coefficients.

TABLE 7.1: UNIT CUBE UNDER UNIAXIAL TENSION

From these results and similar results from the 24 and 96 element cubes (fable 7.2) it was concluded that the boundary element method programme BEM was working for homogeneous bodies at least. Imping was used in the 96 element run without seriously affecting the accuracy. As no analytic solution with which to test the programme for non-homogeneous problems was available, the test case of two unit cubes Under uniaxial tension, in which one of the loaded ends **was** rigid, was used (Fig 7.2), Each cube consisted of 12 elements. Answers appeared reasonable when compared with expected answers **(Table 7.3).**

TABLE 7.2: UNIT CUBE UNDER UNIAXIAL TENSION MODELLED WITH 24 OR 96 ELEMENTS

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TABLE 7.3: RESULTS FOR TWO JOINED CUBES UNDER UNIAXIAL COMPRESSION

The effect of lumping and integration accuracy on accuracy and running time was.also checked against two-dimensional solutions obtained with the displacement discontinuity programme MIMAP. 116 elements were used to represent one eighth of a rectangular tunnel measuring 34 m x 3,2 m x 6 m (the height being 3,6m). The vertical stress was 60 MPa and the horizontal streses 30 MPa each (Fig 7.3). Element sizes were graded further from the tunnel centre where 8 elements were used to span half the hangingwall and 8 for half **the** sidewall. Table 7.4 shows a comparison of **displacements** for two MIMAP and 4 boundary element runs (the displacements represent vertical displacements along the hangingwall section of the tunnel). Details of the integration constants and running times are given in Table 7.5.

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TABLE 7.4: COMPARISON OF BOUNDARY DISPLACEMENTS FOR RECTANGULAR .TONNEL

Table 7,6 shews comparisons of stresses at interior points. The x-y co-ordinates are such that the hangingwall lies at y = 1,8 and the sidewall at x = 1,6, The tables clearly show the high order of integration required to obtain reasonable results for stresses at interior points. Such accurate integration is unnecessary when **solving for surface displacements and tractions and the time savings obtained by lumping and variable integration accuracy are clearly demonstrated by Table 7.5.**

TABLE 7.5: RUNNING TIMES AND INTEGRATION SCHEME COMPARISON

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TABLE 7.6: COMPARISON OF STRESSES AT INTERIOR POINTS

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* Using same integration constants for interior points as in BEN 1 ran.

7.2 Mixed boundary element programme MBEM

Before **combining** the boundary elements **with** displacement discontinuity elements, a test was done to compare the two methods. The displacement discontinuity programme RIDE used is described elsewhere (Starfield and Crouch 1973, Diering 1977). A square flat planar tabular excavation of dimensions 80 m x 80 m subjected to a normal load of 60 MPa was used as the test case and 98 boundary elements with dimensions ranging from 40 m square to 10 m square were used. **Two displacement discontinuity runs were done with 64 10 m square** elements and 16 20 m square elements. The region discretized for the boundary element run was 160 m x 160 m square. Tables 9 and 10 show a comparison of (hangingwall/footwall convergence) and interior stresses and displacements (in **the** hangingwall).

TABLE 10: COMPARISON OF CLOSURES - PROGRAMMES BEM AND RIDE

TABLE 7.8: COMPARISON OF INTERIOR STRESSES AND **DISPLACEMENTS** PROGRAMMES BEM AND RIDE

The discrepancy- in displacements 50 m above the excavation probably arises from the limited **size** of the boundary element mesh While the large tensile stress given by the displacement discontinuity **method** 1 0 m in the hangingwall is as a result of the 1 point integration formula used in evaluating the stress, The otherwise good agreement between the displacements prompted the combination of the two element types. Running times were:

In modelling the excavation **with** boundary elements, it was possible to discretize only the hangingwall and surrounding solid areas of the excavation. If hangingwall and footwall movements were unequal, then twice as many boundary elements would have been required, **the** great **advantages of the displacement discontinuity elements over the boun**dary elements for this type of problem are evident.

A direct verification of the MBEM programme appeared very difficult but was not necessary since all of the programming logic appeared in one or other of the programmes RIDE or **BEM.** A number of **direct** test **examples have been run and answers from these tests have appeared to** be reasonable. One example is given here - **a 4 x 4** array of displacement discontinuity elements. As the depth B increases **the** displacements of **the** boundary element tend to zero while the closures in the displacement discontinuity elements tend cowards those of an independent RIDE run. When **the** depth 3 is small, then the hangingwall movements become significantly greater **than the footwall** move-

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ments (as has 'been demonstrated "by Grouch (1976) in two dimensions). When the depth 3 becomes less than about 0.75 of the maximum element width, then numerical **convergence** is lost. As the constant dis**placement and constant traction assumptions for each element are MG longer valid in this case it is fortunate that the programme automatically rejects such ill-conditioned or badly specified problems. Indeed, the accuracy achieved in any problem, appears to be strongly related to the rate of convergence of the numerical solution* There are of course many excepkiona to this general rule,**

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A second method of testing the programma was to vary all of the integration parameters to check the dependence of the solutions upon the accuracy of numerical integration.

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CHAPTER 8 PROGRAMMING CONSIDERATIONS

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The size of a prograrnme is governed: by the number of programming stega malcing up the programme and by the amount of data required or used by the programme.

8.1 Programme structure

The core requirements of a programme are easily reduced by using programme overlays or "swaps". Only that portion of the programme which **is in use is held in main memory. Provided the programme is well structured, the amount of swapping of programmes in and cut of mala memory is minimal and results in negligible increase in total running time. Fortunately, boundary and finite element formulations are well structure! and the BEM and MBEM programmes were easily divided into the following swaps.**

- **(i) Control pf^ramme**
- **(ii) Input and data checking**
- **(iii) Calculation of influence coefficients**
- **(iv) Iterative solution for unknown displacements and tractions**
- **(v) Output of displacements and tractions**
- **(vi) calculation of stressns and displacements at specified points**

It is possible to stop or start the programme at any stage. Thus, for example, **it is** possible to store the displacement and traction **solutions for a number of runs (iv) while overwriting the very large** influence coefficient file (iii), if necessary for successive runs. **Stresses and displacements may subsequently be calculated at any point for any of the displacement/traction solutions, A typical application of the above procedure arises when a number of different geometries for a problem are studied. After the initial analysis, it becomes necessary to determine stresses and displacements at a few additional points without repeating the entire analysis. This is easily achieved with the above programme structuring.**

8.2 Data storage

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8.2.1 The influence coefficient file (IGF)

The file containing the influence coefficients (the ICF) can easily become very large. For example, a problem containing **1 000 elements (in which the lumping mechanism was not used to reduce the number of influence coefficients, would require** about 36 M bytes.

In designing the programme, it was decided to use sequential access for the ICF. That is to say, the influence coeffi-' cients are calculated in precisely the same order as needed during the iterative solution stage. The use of sequential access for the ICF makes it possible to store the ICP on tape. This will be done shortly, and it Is expected that the increase in running time will be more Chan offset by the reduced cost of cheaper peripheral storage.

Access time to the ICP is reduced by retaining in core a fairly large buffer array (8 k byte in the present versions). This buffer array is also accessed sequentially and a disk/tape transfer is only required every 2 000 coefficients (l coefficient = 4 bytes).

All access to the ICP is controlled by one small subroutine so that the programmes are not too machine specific.

8.2.2 Storage of **lump** elements

All the information for the elements of any one lump is stored on two disk blocks of 512 bytes each. This information includes displacements, tractions, direction cosines, areas and codes. The nodal co-ordinates Which are used by more than one lump in many cases are retained in main memory.
Since it is never necessary to retain in main memory the detailed information of more than two lump elements at any one time, the two lump elements are referred to as a receiving and an emitting lump element. Each has assigned to it a labelled common block of 512 words. It is thus a simple matter to read **or write the information for all the elements within a lump to** or from disk.

Programme listings

A complete listing of programme BEM is given in Appendix 2 while partial listings of programmes MBEM and MINAPH are given in Appendix 3 and 4. A partial listing of MBEM is given to **avoid** repetition since numerous subroutines are either common to BEM and MBEM or **at** least similar to one another.

A partial listing of MINAPH is given because the original programme was developed by Crouch (1976). Only those sections which were changed. to facilitate modelling of **non-homogenous** problems are given.

CHAPTER 9 EXAMPLES OF PRACTICAL APPLICATIONS

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Before describing some of the practical applications of BEM and MBEM, it is useful to briefly summarise some of the uncertainties Which accompany a typical stress analysis problem in.rock mechanics. In addition, a number **of useful rules of thumb have evolved from the use of these programmes.**

The loading for a problem arises primarily **from** the primitive or in **situ stresses** Which are present in **the** rock mass before mining commences. It is **possible to** measure these stresses with reasonable accuracy, (Gay, 1975) but the presence of dykes, faults etc can result in a fairly irregular stress distribution in the rock mass even before any mining commences. A more Usual approach is to assume **that** the overburden stresses increase linearly with depth below surface and that horizontal stresses are a constant fraction of the vertical stress. Gay (1975) shews how the horizontal **to** vertical stress ratio varies **with** depth on average in Southern Africa.

The excavation geometry for a typical problem is usually very **complex** so that a number of simplifying assumptions have to be made. Usually, service excavations are much smaller than the production excavations, so that only the latter are modelled. Sometimes the extent of mining is such that it is not possible to model all the production excavations. Typical examples of this occur in the Witwatersrand gold fields where mining is more or less continuous for distances exceeding twenty five kilometers.

The geology of most **problems** is usually complex and is also usually based on a number of boreholes. It is practical therefore only to consider major geological horizons with significantly different material properties.

The material properties are seldom **known** with any great degree of certainty. Apart from material anisotropies Which are difficult to measure, a number of other problems arise in assessing suitable material properties. Elastic moduli and material strength are usually determined from small specimen tests. It is known that actual large scale properties

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