

Micromechanics Analysis Code With Generalized Method of Cells (MAC/GMC) User Guide: Version 3.0

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1.0	Introduction						
2.0	Current Capabilities						
3.0	Background						
	3.1 Micromechanics Models	7					
	3.2 Integration Algorithms1	4					
	3.3 Available Constituent Constitutive Models 1	6					
	3.3.1 Transversely Isotropic Elastic Model1	6					
	3.3.2 Anisotropic Elastic Model	ן או					
	3.3.4 Modified Bodner-Partom Model	9					
	3.3.5 Robinson Viscoplastic Model	20					
	3.3.6 Generalized Viscoplastic Potential Structure (GVIPS) Model 2	2					
	3.3.7 Transversely Isotropic GVIPS Model (TGVIPS)	24					
	3.4 Laminate Theory	26					
	3.5 Fatigue Damage Analysis	30 34					
	3.5.1 Above Initial Fatigue Limit	51 82					
	3.6 User Defined Inelastic Material Model	34					
	3.7 User Defined Functional Form For Material Properties	37					
	3.8 Interface Modeling	12					
	3.9 References	15					
40	Bunning MAC/GMC 4	16					
4.0	4.0.1 MAC/GMC - stand-alone:	16					
	4.0.2 FEAMAC - Accessing MAC through ABAQUS:	6					
	4.1 Input and Output Files	17					
	4.2 Input Requirements (for both MAC/GMC and FEAMAC)5	51					
	4.2.1 Header Line:	;3					
	4.2.2 Output Print Level:	53 54					
	4.2.4 Mechanical Load Control:	57					
	4.2.5 Temperature Control:	58					
	4.2.6 Fatigue Damage Option:6	30					
	4.2.7 Yield and Damage Surface Generation Analysis	32					
		53 54					
	4.2.10 Integrator Identification	56					
	4.2.11 Thermal Conductivity Calculation: (Optional)6	57					
	4.2.12 Constituent Material Model Identification:6	38					
	4.2.13 RVE Data:	7					
	4.2.14 Interface Data: (Optional, for IDP=1, 2, 3, 6 or 11)	94 25					
	4.2.15 Debond Data. (Optional)	,5 39					
	4.2.17 Curve Data:)0					
5.0	Conclusion/Euture Modifications)4					
2.2	5.1 Acknowledgment)4					
60	EXAMPLE PROBLEMS)5					
0.0	6.1 Example A: Pure Mechanical Load	.5					

 6.3 Example C: Thermomechanical Load Problem
6.4 Example D: Transverse Debonding
6.5 Example E: General Loading 120
6.6 Example F: Unidirectional Laminate Problem
6.7 Example G: A Cross-Ply Laminate Problem
6.8 Example H: Triple Periodic GMC; [0/90] 129
6.9 Example I: A User Defined RVE 132
6.10 Example J: A Biaxial Load 135
6.11 Example K: User Defined Inelastic Material Model
6.12 Example L: Fatigue Damage Analysis 156
6.13 Example M: Longitudinal Discrete Fiber Breakage
6.14 Example N: Plain Weave Composite 174
6.15 Example O: FEAMAC Analysis
6.16 Example P: Non-Symmetric Laminate
6.17 Example Q: Yield Surface Plot
6.18 Example R: Ellipsoidal Inclusions
6.19 Example S: PATRAN/MSC MACPOST Output
6.20 Example T: Effective Thermal Conductivity Calculation

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1.0 Introduction

The ability to accurately predict the thermomechanical deformation response of advanced heterogeneous materials (e.g., multi-phase composites, cellular solids, etc.) plays a critical role in the development and practical implementation of these strategic materials. Analytical models that predict the effective behavior of composites are used not only by engineers performing structural analysis of largescale composite components, but also by materials scientists in developing new material systems. For an analytical model to fulfill these two distinct functions it must be based on a micromechanics approach which utilizes physically based deformation and life constitutive models and generates the average (macro) response of a composite material given the properties of the individual constituents and their geometric arrangement. Only then can such a model be used by a materials scientist to investigate the effect(s) of different deformation mechanisms on the overall response of the composite in order to identify the appropriate constituents for a given application. Alternatively, if a micromechanical model is to be used in a large-scale structural analysis it must be 1) computationally efficient, 2) able to generate accurate displacement and stress fields at both the macro and the micro levels and 3) be compatible with the finite element method. Also, as advancements in processing and fabrication techniques make it possible to more accurately engineer (tailor) the architectures of these advanced composite systems, development of a computationally efficient micromechanics analysis tool capable of accurately predicting the effect of microstructural details on the internal and macroscopic behavior of composites becomes even more important. Such computational efficiency is absolutely required because 1) the large number of parameters which must be varied in the course of engineering (or designing) composite (heterogeneous) materials, and 2) the optimization of a material's microstructure requires the integration of the micromechanics model with optimization algorithms. From this perspective, analytical approaches which use closed-form expressions to describe the effect of a material's internal architecture on the overall material behavior are preferable to numerical methods, such as finite element or finite difference schemes.

A number of models presently exist that can fulfill certain aspects of the aforementioned tasks. However, there are very few working models that are both computationally efficient and sufficiently accurate both at the micro- and macro-levels. One such micromechanics model with the potential of fulfilling both tasks is Aboudi's method of cells [1] and its continuous reinforcement [2] and discontinuous reinforcement [3] generalizations. The comprehensive capabilities and efficiency of this method have been documented in references [4] and [5]. Consequently, the recently developed, computationally efficient and comprehensive micromechanics analysis code, MAC/GMC, will now be described: with a predictive capability resting entirely upon the fully analytical micromechanics model, herein referred to as the generalized method of cells, GMC, [2 and 3] . MAC/GMC is a versatile form of research software that "drives" the double or triple periodic micromechanics constitutive models based upon GMC. GMC is capable of predicting the response of both continuous and discontinuous multi-phase composites with arbitrary internal microstructures and reinforcement shapes. **GMC** is a continuum-based micromechanics model that provides closed-form expressions for the macroscopic composite response in terms of the properties, size, shape, distribution, and response of the individual constituents or phases that make up the material. **GMC** also utilizes physically based viscoplastic deformation and life models for each constituent. Furthermore, expressions relating the internal stress and strain fields in the individual constituents in terms of the macroscopically applied stresses and strains are available through strain or stress concentration factors. These expressions make possible the investigation of failure processes at the microscopic level at each step of an applied load history. Similarly, **GMC** provides the capability of studying the influence of bond strength at the fiber/matrix interface which has been shown to be an important damage mechanism in composites [6].

MAC/GMC enhances the basic capabilities of GMC by providing a modular framework wherein 1) various thermal, mechanical (stress or strain control) and thermomechanical load histories can be imposed, 2) different integration algorithms may be selected, 3) a variety of material constitutive models (both deformation and life) may be utilized and/or implemented and 4) a variety of fiber architectures (both unidirectional, laminate and woven) may be easily accessed through their corresponding representative volume elements contained within the supplied library of RVEs or input directly by the user, and 5) graphical post processing of the macro and/or micro field quantities is made available. Consequently, the availability of MAC/GMC now provides industry, academia and government engineers and materials scientists with a comprehensive, computationally efficient, user-friendly micromechanics analysis tool that can easily and accurately design/analyze multi-phase (composite) materials for a given application. MAC/GMC is also ideally suited for conducting sensitivity/parametric studies (i.e., "what-if" scenarios) in the design/analysis of advanced composite materials (e.g., MMC/IMCs, PMCs, and CMCs). Figure 1 illustrates the basic flow diagram for this modular framework code. Furthermore, since GMC is a continuum-based micromechanics model that provides closed form expressions for the macroscopic composite response in terms of the properties, size, shape, distribution, and response of the individual constituents or phases that make up the material, MAC/ GMC can be interfaced directly with standard linear and nonlinear finite element analysis packages (through their respective user definable constitutive routines) for cost-effective large scale component design and analysis. Currently, such an interface exists only for HKS's nonlinear finite element code, ABAQUS. For clarity purposes, when this finite element implementation of MAC/GMC is discussed within this manual it will be referred to as FEAMAC, for example see section 4.1.2.

The capabilities of this version of **MAC/GMC** are listed in section 2, whereas theoretical and background information on the basic capabilities previously itemized are given in section 3. Section 4 describes in detail the required input for using **MAC/GMC** and will be the most referenced section in the entire manual. Section 5 gives some insights into the future modifications planned for **MAC/GMC** and in section 6 examples A-S illustrate and describe the various features of **MAC/GMC**.



Figure1: MAC Flowchart

2.0 Current Capabilities

In this section the current features/capabilities of **MAC/GMC** are itemized, with those new to this version being denoted by **bold type face**.

- Load Types:
 - Thermal, Mechanical, Thermomechanical; Generalized Loading Option
- Aboudi GMC Models (Reformulated for computational speed):
- ..- Double Periodicity Model for continuous reinforcement
 - Triple Periodicity Model for discontinuous/woven reinforcement
- Library of Representative Volume Elements:
 - Double Periodicity Model: circular fiber representations
 - Triple Periodicity Model: ellipsoidal inclusion, open cell
- Input / Graphical Output
 - Free Format Input Data
 - Up to 5 x-y data plot files generated for macro and micro (subcell) quantities
 - PATRAN for subcell geometry and color results evaluation, e.g. stress, strains, inelastic strains, J_2 , etc., via **MACPOST**
- Integration Options:
 - Forward Euler
 - Predictor/Corrector
- Constitutive Models (expanded internal material database)
 - elastic: transversely isotropic and anisotropic
 - inelastic viscoplastic models:
 - Bodner-Partom
 - Robinson
 - GVIPS isotropic form
 - TGVIPS transversely isotropic form
 - user defined constitutive model via subroutines USRMAT (inelastic), USR-FORMDE and USRCPEVAL (elastic); user defined functional dependence of material constants via subroutine USRFUN
- Interface Modeling
 - fiber/matrix interface layer (distinct interfacial material)
 - fiber/matrix debonding conditions (interfacial displacement discontinuity)
- Fatigue Damage Analysis
- Yield Surface Calculations
- Symmetric and Nonsymmetric Laminate Analysis
- FEAMAC allows easy finite element access to MAC/GMC capabilities

3.0 Background

As stated in the introduction, **MAC/GMC**'s predictive capabilities rest upon the fully analytical micromechanics model known as **GMC** which is capable of predicting the inelastic response of both continuous (double periodicity) and discontinuous/woven (triple periodicity) multi-phase composites with arbitrary internal microstructures and reinforcement shapes. Included in this section is a brief overview of the theoretical foundations of **GMC**; a more complete discussion is given in references [1-3]. Also, included in this section are discussions of 1) the available integration algorithms, 2) various constitutive models, 3) theory underlying the laminate, 4) fatigue damage, and 5) interfacial modeling approaches incorporated into **MAC/GMC**. Besides describing the available constitutive models, explanations of the user defined inelastic and elastic material models are included. Finally, the available architectures (Repeating Volume Elements, or RVEs) and required input are then described in section 4.0.

3.1 Micromechanics Models

In the original formulation of the method of cells, a continuously (or discontinuously) reinforced, unidirectional fibrous composite was modeled as a rectangular, double-periodic (or triple-periodic) array of fibers embedded in a matrix phase. The periodic character of the assemblage allowed identification of a repeating unit cell that can be used as a building block to construct the entire composite. The properties of the repeating cell were thus representative of the properties of the entire assemblage. The unit cell consisted of a single fiber subcell surrounded by three matrix subcells for continuous, and seven for discontinuous, composites, hence the name **method of cells**. The rectangular geometry of the repeating unit cell allowed one to obtain an approximate solution for the stresses and strains in the individual subcells given some macroscopically homogeneous state of strain or stress applied to the composite. The approximate solution to the posed boundary value problem was, in turn, used to determine macroscopic (average) properties, traditionally referred to as effective properties, of the composite and the macro (global) stress-strain response in the inelastic region.

In the **generalized method of cells** for continuous (or discontinuous) fibrous composites, the repeating unit cell can consist of an arbitrary number of phases, see Fig. 2. Hence, the generalized method of cells is capable of modeling a multiphase composite. This generalization extends the modeling capability of the original method of cells to include the following: 1) inelastic thermomechanical response of multi-phase metal matrix composite, 2) modeling of various fiber architectures (including both shape and packing arrangements), 3) modeling of porosities and damage, and 4) modeling of interfacial regions around inclusions including interfacial degradation.

The basic homogenization approach taken in the micromechanical analysis consists essentially of four steps. First, the RVE, of the periodic composite is iden-

tified. Second, the macroscopic or average stress and strain states in terms of the individual microscopic (subcell) stress and strain states is defined. Third, the continuity of tractions and displacements are imposed at the boundaries between the constituents. These three steps, in conjunction with micro-equilibrium, establish the relationship between micro (subcell) total, thermal and inelastic strains and macro (composite) strains via the relevant concentration tensors. In the fourth and final step, the overall macro constitutive equations of the composite are determined. These four steps form the basis of the micro-to-macromechanics analysis which describes the behavior of heterogeneous media. The resulting micromechanical analysis establishes the overall (macro) behavior of the multi-phase composite which is expressed as a constitutive relation between the average stress, strain, thermal, and inelastic strains, in conjunction with the effective elastic stiffness matrix.

That is,

$$\bar{\overline{g}} = \underline{B}^* (\bar{\underline{\varepsilon}} - \bar{\underline{\varepsilon}}^I - \bar{\underline{\varepsilon}}^T)$$
(EQ 1)

where, for the most general case of discontinuous reinforcement with N_{α} by N_{β} by N_{γ} number of subcells, the effective elastic stiffness matrix, \underline{B}^{*} , of the composite is given by,

$$\underline{B}^{*} = \frac{1}{dhl} \sum_{\alpha=1}^{N_{\alpha}} \sum_{\beta=1}^{N_{\beta}} \sum_{\gamma=1}^{N_{\gamma}} d_{\alpha} h_{\beta} l_{\gamma} \underline{C}^{(\alpha\beta\gamma)} \underline{A}^{(\alpha\beta\gamma)}$$
(EQ 2)

the composite inelastic strain vector is defined as,

$$\bar{\boldsymbol{\xi}}^{I} = \frac{-\underline{\boldsymbol{B}}^{\ast - 1}}{dhl} \sum_{\alpha = 1}^{N_{\alpha}} \sum_{\beta = 1}^{N_{\beta}} \sum_{\gamma = 1}^{N_{\gamma}} d_{\alpha} h_{\beta} l_{\gamma} \underline{\boldsymbol{\zeta}}^{(\alpha\beta\gamma)} (\underline{\boldsymbol{D}}^{(\alpha\beta\gamma)} \underline{\boldsymbol{\varepsilon}}_{s}^{I} - \bar{\boldsymbol{\varepsilon}}^{I(\alpha\beta\gamma)})$$
(EQ 3)

the average thermal strain vector as,

$$\bar{\boldsymbol{\varepsilon}}^{T} = \frac{-\underline{\boldsymbol{\mathcal{B}}}^{*-1}}{dhl} \sum_{\alpha=1}^{N_{\alpha}} \sum_{\beta=1}^{N_{\beta}} \sum_{\gamma=1}^{N_{\gamma}} d_{\alpha} h_{\beta} l_{\gamma} \underline{\boldsymbol{\mathcal{C}}}^{(\alpha\beta\gamma)} (\underline{\boldsymbol{\mathcal{D}}}^{(\alpha\beta\gamma)} \underline{\boldsymbol{\varepsilon}}_{s}^{T} - \bar{\boldsymbol{\varepsilon}}^{T(\alpha\beta\gamma)})$$
(EQ 4)

and \bar{g} is the uniform applied macro (composite) strain. For the case of continuous reinforcements with N_{β} by N_{γ} number of subcells, EQS. 2 - 4 reduce to the following:

$$\underline{B}^{*} = \frac{1}{hl} \sum_{\beta=1}^{N_{\beta}} \sum_{\gamma=1}^{N_{\gamma}} h_{\beta} l_{\gamma} \underline{C}^{(\beta\gamma)} \underline{A}^{(\beta\gamma)}$$
(EQ 5)



Figure 2: Subcell Dimension Nomenclature

$$\bar{\varepsilon}^{I} = \frac{-\underline{B}^{\ast - 1}}{hl} \sum_{\beta = 1}^{N_{\beta}} \sum_{\gamma = 1}^{N_{\gamma}} h_{\beta} l_{\gamma} \underline{C}^{(\beta\gamma)} (\underline{D}^{(\beta\gamma)} \underline{\varepsilon}_{s}^{I} - \overline{\varepsilon}^{I(\beta\gamma)})$$
(EQ 6)

$$\bar{\boldsymbol{\varepsilon}}^{T} = \frac{-\underline{\boldsymbol{\mathcal{B}}}^{*-1}}{hl} \sum_{\beta=1}^{N_{\beta}} \sum_{\gamma=1}^{N_{\gamma}} h_{\beta} l_{\gamma} \underline{\boldsymbol{\mathcal{C}}}^{(\beta\gamma)} (\underline{\boldsymbol{\mathcal{D}}}^{(\beta\gamma)} \underline{\boldsymbol{\varepsilon}}_{s}^{T} - \bar{\boldsymbol{\varepsilon}}^{T(\beta\gamma)})$$
(EQ 7)

In the above equations matrix notation is employed; where, for example, the average stress, \bar{g} ; average strain, $\bar{\epsilon}$, and inelastic subcell strain, $\bar{\epsilon}'_s$, vectors represent,

$$\overline{\overline{g}} = \{\overline{\sigma}_{11}, \overline{\sigma}_{22}, \overline{\sigma}_{33}, \overline{\sigma}_{23}, \overline{\sigma}_{13}, \overline{\sigma}_{12}\}$$
(EQ 8)

$$\bar{\xi} = \{\bar{\varepsilon}_{11}, \bar{\varepsilon}_{22}, \bar{\varepsilon}_{33}, 2\bar{\varepsilon}_{23}, 2\bar{\varepsilon}_{13}, 2\bar{\varepsilon}_{12}\}$$
(EQ 9)

$$\underline{\varepsilon}_{s}^{I} = \{ \underline{\overline{\varepsilon}}^{I(111)}, ..., \underline{\overline{\varepsilon}}^{I(N_{\alpha}N_{\beta}N_{\gamma})} \}$$
 (EQ 10)

where the six components of the vector $\bar{\varepsilon}^{I(\alpha\beta\gamma)}$ are arranged as in EQ. 9. Similar definitions for $\bar{\varepsilon}_{s}^{T}$, $\bar{\varepsilon}^{T(\alpha\beta\gamma)}$ also exist. Note that the key ingredient in the construction of this macro constitutive law is the derivation of the appropriate concentration matrices, $\underline{A}^{(\alpha\beta\gamma)}$ and $\underline{D}^{(\alpha\beta\gamma)}$ having the dimensions 6 by 6 and 6 by $N_{\alpha}N_{\beta}N_{\gamma}$ respectively, at the micro (subcell) level. The definitions of \underline{A} and \underline{D} , although not given here, may be found in references [2] and [3]. Finally, the matrix $\underline{C}^{(\alpha\beta\gamma)}$ represents the elastic stiffness tensor of each subcell ($\alpha\beta\gamma$) and d_{α} , h_{β} , l_{γ} the respective subcell dimensions (see Fig. 2) wherein,

$$d = \sum_{\alpha=1}^{N_{\alpha}} d_{\alpha} \qquad h = \sum_{\beta=1}^{N_{\beta}} h_{\beta} \qquad l = \sum_{\gamma=1}^{N_{\gamma}} l_{\gamma}$$

Similarly, given the concentration matrices $\underline{A}^{(\alpha\beta\gamma)}$ and $\underline{D}^{(\alpha\beta\gamma)}$, expressions for the average strain in each subcell can be constructed, i. e.,

$$\bar{\boldsymbol{\varepsilon}}^{(\alpha\beta\gamma)} = \boldsymbol{A}^{(\alpha\beta\gamma)} \bar{\boldsymbol{\varepsilon}} + \boldsymbol{\tilde{D}}^{(\alpha\beta\gamma)} (\boldsymbol{\varepsilon}_{s}^{I} + \boldsymbol{\varepsilon}_{s}^{T})$$
(EQ 11)

as well as average stress

$$\overline{g}^{(\alpha\beta\gamma)} = \underline{C}^{(\alpha\beta\gamma)} [\underline{A}^{(\alpha\beta\gamma)} \underline{\overline{\varepsilon}} + \underline{D}^{(\alpha\beta\gamma)} (\underline{\varepsilon}_{s}^{I} + \underline{\varepsilon}_{s}^{T}) - (\underline{\varepsilon}^{I(\alpha\beta\gamma)} + \underline{\varepsilon}^{T(\alpha\beta\gamma)})]$$
(EQ 12)

The analytic constitutive law, see EQ. 1, may be readily applied to investigate the behavior of various types of composites, given knowledge of the behavior of the individual phases. Note that within **MAC/GMC** an incremental tangent solution scheme is utilized, therefore the appropriate rate forms of EQS. 1-12 are employed.

Numerous advantages can be stated regarding the current macro/micro constitutive laws as compared to the other numerical micromechanical approaches in the literature, e.g. the finite element unit cell approach. One advantage is that any type of simple or combined loading (multiaxial state of stress) can be applied irrespective of whether symmetry exists or not, as well as without resorting to different boundary condition application strategies as in the case of the finite element unit cell procedure. Another, advantage concerns the availability of an analytical expression representing the macro elastic-thermo-inelastic constitutive law, thus ensuring a reduction in memory requirements when implementing this formulation into a structural finite element analysis code. Furthermore, this formulation has been shown to predict accurate macro behavior given only a few subcells, within the repeating cell (see references [2], and [4]). Whereas, if one employs the finite element unit cell procedure, a significant number of finite elements are required within a given repeating unit cell to obtain the same level of global accuracy as with the present formulation. Consequently, it is possible to utilize this formulation to efficiently analyze metal matrix composite structures subjected to complex thermomechanical load histories. This is particularly important when analyzing realistic structural components, since different loading conditions exist throughout the structure, thus necessitating the application of the macromechanical equations repeatedly at these locations.

The equations of **GMC-3D** (and consequently through appropriate specialization **GMC-2D**) have recently been reformulated [7] in a way that significantly increases the computational efficiency of the model. This new reformulation has now been implemented into **MAC/GMC** and is the default computational mode. By nature of the traction continuity conditions within the original generalized method, all six stress components are not unique in every subcell. Normal stress components are constant in certain rows of subcells, while shear stress components are constant in certain layers of subcells. The unique subcell stress components are denoted as,

$$T_{11}^{(\beta\gamma)}, T_{22}^{(\alpha\gamma)}, T_{33}^{(\alpha\beta)}, T_{23}^{(\alpha)}, T_{13}^{(\beta)}, T_{12}^{(\gamma)}$$
 (EQ 13)

Consequently, a more efficient formulation of **GMC** can be obtained by applying traction continuity directly (i.e. recognizing that traction continuity conditions require no more and no less than the aforementioned reduction in subcell stress components) and using subcell stresses rather than strains as the basic unknown quantities. Accordingly, the continuity of displacement conditions are formed in terms of subcell stresses (through the use of the subcell constitutive and kinematic equations), and the *mixed concentration equations* for the unit cell are constructed,

$$\tilde{T} = \tilde{Q}^{(\alpha\beta\gamma)}\tilde{\varepsilon} + \tilde{Q}^{IT(\alpha\beta\gamma)}(\tilde{\varepsilon}_{s}^{I} + \tilde{\varepsilon}_{s}^{T})$$
(EQ 14)

Here, \underline{T} is the vector of all subcell stress components (listed in EQ 13), $\underline{G}^{(\alpha\beta\gamma)}$ is

the subcell mixed concentration matrix, and $\mathcal{G}^{IT(\alpha\beta\gamma)}$ is the subcell inelastic-thermal mixed concentration matrix. The term *mixed* is used here because EQ 14 relates local (subcell) stresses to global strains. Clearly this equation contrasts with its original formulation counterpart, EQ 11, which relates local strains to global strains and is thus the unit cell strain concentration equation. As one could obtain EQ 1 from EQ. 12 in the original formulation, similar expressions for the global quantities ($\underline{B}^*, \bar{\epsilon}^I$ and $\bar{\epsilon}^T$) can be determined easily from EQ 14 (see [7] for details).

The increased efficiency of the reformulation of **GMC** emerges mainly due to the increased efficiency of forming EQ 14 versus forming EQ 11. The formation of EQ 14 requires solution (of linear equations) for the unknown independent subcell stress components listed in EQ 13, numbering $N_{\beta}N_{\gamma} + N_{\alpha}N_{\gamma} + N_{\alpha}N_{\beta} +$

 $N_{\alpha} + N_{\beta} + N_{\gamma}$. The formation of EQ 11 requires solution for 6 unknown strain

components for each subcell, or a total of $6N_{\alpha}N_{\beta}N_{\gamma}$ unknowns. Solution of linear equations, in essence, amounts to the inversion of a matrix which has the rank of the number of unknown quantities. Since the computational effort associated with matrix inversion increases approximately as the **cube** of the matrix rank, reducing the number of unknown quantities has a major impact on computational efficiency. The reduction in unknowns due to the reformulation is shown in Figure 3 and can be quite significant. For example, a $10 \times 10 \times 10$ unit cell originally required solution for 6000 unknowns. In the reformulated version of **GMC-3D** the number of unknowns is reduced to 330. Consequently, the corresponding execution times are reduced as well. Table I gives sample CPU times for identical cases executed using the original and reformulated versions of **GMC** with continuous reinforcement. Clearly, as the number of subcells in the repeating unit cell increases, the increase in efficiency attributable to the reformulation becomes astronomical. It is important to note that:

- (1) The reformulation of **GMC** gives **identical results** as the original formulation of GMC **in every case**.
- (2) The significant increase in the computational efficiency of GMC due to the reformulation allows:
 - (a) Analysis of simple unit cells (i.e. few subcells) in a fraction of the time.
 - (b) Analysis of refined unit cells (i.e. many subcells) which was previously impossible (due to excessive execution times and memory requirements) can now be performed in times comparable to those previously required for analysis of simple unit cell, see Table I.



Figure 3. Number of subcells vs. number of unknown variables for the original and reformulated versions of GMC-3D for $N_{\alpha} = N_{\beta} = N_{\gamma}$

Table I: CPU Times (seconds) for Heat-Up of Unidirectional SiC/TiAI

GMC Version	Subcell Discretization								
	2x2	4x4	6x6	8x8	10x10	12x12	20x20	100x100	
original reformulated speed-up ratio	0.87 0.18 4.8	19 0.25 76	182 0.5 364	508 0.9 564	8,679 1.5 5,786	43,781 2.3 19,035	8.3	796	

Note: These execution times are for a specialized version of GMC-2D and not MAC/GMC, therefore one should not expect to see the same execution times when running MAC/GMC due to increased overhead. However, speed-up ratios should be comparable.

3.2 Integration Algorithms

There are two integration algorithms currently available within **MAC/GMC**. The first is the standard, explicit Forward Euler algorithm, which can be expressed as,

$$w_{i+1} = w_i + \Delta t f(t_i, w_i)$$

where

$$w_i = \underline{y}(t_i)$$
$$w_{i+1} = \underline{y}(t_{i+1})$$

and $f(t_i, w_i)$ is the rate of change with respect to time of the vector, $y(t_i)$, i.e.

$$f(t_i, w_i) = \dot{y} = \frac{dy(t_i)}{dt}$$

The second is a predictor/corrector algorithm with a self-adaptive time step which uses:

1) a 4th order Runge-Kutta starter:

$$w_i = w_{i-1} + \frac{(K_1 + 2K_2 + 2K_3 + K_4)}{6}$$

where

$$K_{1} = \Delta t f(t_{i-1}, w_{i-1})$$

$$K_{2} = \Delta t f\left(t_{i-1} + \frac{\Delta t}{2}, w_{i-1} + \frac{K_{1}}{2}\right)$$

$$K_{3} = \Delta t f\left(t_{i-1} + \frac{\Delta t}{2}, w_{i-1} + \frac{K_{2}}{2}\right)$$

$$K_{4} = \Delta t f(t_{i-1} + \Delta t, w_{i-1} + K_{3})$$

2) with an Adams-Bashforth four-step predictor:

$$w_{0} = \alpha_{1} \qquad w_{1} = \alpha_{2} \qquad w_{2} = \alpha_{3} \qquad w_{3} = \alpha_{4}$$

$$w_{i+1}^{P} = w_{i} + \frac{\Delta t}{24} [55f(t_{i}, w_{i}) - 59f(t_{i-1}, w_{i-1}) + 37f(t_{i-2}, w_{i-2}) - 9f(t_{i-3}, w_{i-3})]$$

and 3) an Adams-Moulton four step corrector:

$$w_{i+1}^{c} = w_{i} + \frac{\Delta t}{24} [9f(t_{i+1}, w_{i+1}^{p}) + 19f(t_{i}, w_{i}) - 5f(t_{i-1}, w_{i-1}) + f(t_{i-2}, w_{i-2})]$$

where the α 's come from the 4th order Runge-Kutta starter. Further details may be found in [8].

Note: It has been found, based on experience, that for relatively rapid monotonic or cyclic loadings it may be more efficient to use the Forward Euler integrator since the predictor/corrector requires 5 evaluations per step, as shown above. However, in the case of creep, relaxation or slow monotonic or cyclic loading histories, significant increases in solution speeds can be obtained using the predictor/corrector algorithm.

Finally, within **MAC/GMC** the vector w as used above contains the following macro quantities:

<u>position</u>	<u>contents</u>
1 - 6	Macro Total Strain
7 - 12	Macro Stress
13 - 18	Macro Inelastic Strain
19 - 30	(currently empty
	space for 2 6x1 vectors)
31 - 36	Macro Thermal Strain
37	Current Temperature
38- 46	Future Use

and the quantities associated with each subcell are stored sequentially in w, such that

position	<u>contents</u>
47 - 52	Micro Total Strain
53 - 58	Micro Stress
59 - 64	Micro Inelastic Strain
65 - 76	Micro Internal State Variables
	(space for 2 6x1 vectors)
77 - 82	Micro Thermal Strain
83 - 88	Debonding Parameters

The above 42 positions (i.e., positions 47-88) are repeated for the total number of subcells (N) thus bringing the total length of the w vector to 46+42N. It follows that a second vector of similar length that contains the corresponding macro and micro rates, e.g., $f(t_i, w_i)$, is also utilized.

3.3 Available Constituent Constitutive Models

Currently **MAC/GMC** provides two elastic and five inelastic constitutive models. These models have been selected purely based upon the availability of material parameters for the materials of interest. However, **MAC/GMC** is designed in a modular fashion thus allowing the implementation of additional constitutive models through a user defined subroutine. Two of the five available inelastic models are capable of representing transversely isotropic material behavior, thus allowing one to investigate the reinforcement of an anisotropic matrix allowing idealization of a heterogeneous material via a pseudo-homogenous anisotropic material (e.g. fiber tow). In all five inelastic models a purely elastic response is possible by modifying a single material parameter for each model as noted below.

3.3.1 Transversely Isotropic Elastic Model

Reference: Jacob Aboudi, Mechanics of Composite Materials, Elsevier, 1991

The following transversely isotropic model is provided for those materials that have an elastic only response and whose strong direction is aligned with the 1 axis shown in Fig. 6, such as the fiber constituent in a composite.

$$\begin{bmatrix} \sigma_{11} \\ \sigma_{22} \\ \sigma_{33} \\ \sigma_{23} \\ \sigma_{13} \\ \sigma_{12} \end{bmatrix} = \begin{bmatrix} C_{11} \ C_{12} \ C_{12} \ 0 \ 0 \ 0 \\ C_{12} \ C_{22} \ C_{23} \ 0 \ 0 \ 0 \\ C_{12} \ C_{23} \ C_{22} \ 0 \ 0 \ 0 \\ 0 \ 0 \ 0 \ C_{44} \ 0 \ 0 \\ 0 \ 0 \ 0 \ C_{44} \ 0 \\ \gamma_{13} \\ \gamma_{12} \end{bmatrix} - \begin{bmatrix} \alpha_L \Delta T \\ \alpha_T \Delta T \\ \alpha_T \Delta T \\ 0 \\ 0 \\ 0 \end{bmatrix}$$

where components C_{ii} can be expressed in terms of five independent constants,

$$E_A, E_T, v_A, v_T, G_A$$

thus,

$$C_{11} = E_A + 4\kappa v_A^2$$

$$C_{12} = 2\kappa v_A$$

$$C_{22} = \kappa + \frac{0.5E_T}{(1 + v_T)}$$

$$C_{23} = \kappa - \frac{0.5E_A}{(1 + v_T)}$$

$$C_{44} = G_A$$

$$C_{66} = \frac{(C_{22} - C_{23})}{2}$$

with

$$\kappa = 0.25 E_A / [0.5(1 - v_T)(E_A / E_T) - v_A^2]$$

3.3.2 Anisotropic Elastic Model

Reference: S. M. Arnold, A Transversely Isotropic Thermoelastic Theory, NASA TM 101302, 1988

An alternative transversely isotropic model has also been provided for those materials that have an elastic only response, but whose strong direction may not be aligned with the 1 axis shown in Fig. 6 (i.e., the plane of isotropy is allowed to rotate), for example in the case of woven composites, see **Example N**. This elastic model is consistent with the anisotropic inelastic models described subsequently and is defined within the global coordinate system via a direction cosine vector, d_i .

$$\sigma = [C] \{ \varepsilon - \varepsilon^{in} \}$$

.1.

where the stiffness matrix, C, is a function of the five independent material parameters E_A , E_T , v_A , v_T , G_A and the direction cosine vector, d_i , and, in general is fully populated. Note the multiaxial thermal strain tensor is assumed to have the following form,

$$\varepsilon_{ij}^{th} = [(\alpha_L - \alpha_T)d_id_j + \delta_{ij}\alpha_T]\Delta T$$

Further details can be found in the above reference.

3.3.3 Bodner-Partom Model

Reference: Jacob Aboudi, Mechanics of Composite Materials, Elsevier, 1991

This model represents a Bodner-Partom viscoplastic material with isotropic hardening, Z, and can be used for an initially isotropic metallic material.

The flow law is given as:

$$\dot{\varepsilon}_{ij}^{I} = \Lambda s_{ij}$$

where

$$\Lambda = \sqrt{\frac{D_2^{PL}}{J_2}}$$
$$D_2^{PL} = D_0^2 exp \left[-\left(\frac{A^2}{J_2}\right)^n \right]$$
$$A^2 = \frac{1}{3} Z_{eff}^2 \left(\frac{n+1}{n}\right)^{\frac{1}{n}}$$
$$J_2 = \frac{1}{2} S_{ij} S_{ij}$$
$$S_{ij} = \sigma_{ij} - \frac{1}{3} \sigma_{kk} \delta_{ij}$$

The evolution law for isotropic hardening is given as:

$$\dot{Z} = m(Z_1 - Z_{eff}) \frac{\dot{W}^{PL}}{Z_0}$$

where Z_0 , Z_1 and *m* are inelastic constants and the plastic work rate, \dot{W}^{PL} , is given by;

$$\dot{W}^{PL} = \sigma_{ij} \dot{\varepsilon}_{ij}^{I}$$

$$Z_{eff} = Z_0 + q \int_0^t \dot{Z}(\tau) d\tau + (1 - q) \sum_{i, j = 1}^3 r_{ij} \int_0^t \dot{Z}(\tau) r_{ij}(\tau) d\tau$$

$$r_{ij}(t) = \sigma_{ij}(t) / [\sigma_{kl}(t)\sigma_{kl}(t)]^{1/2}$$

An elastic only response may be obtained by setting the material parameter D_0 to zero.

3.3.4 Modified Bodner-Partom Model

Reference: R. W. Neu, "Nonisothermal Material Parameters For the Bodner-Partom Model", MD-Vol. 43, Material Parameter Estimation for Modern Constitutive Equations, L. A. Betram, S.B. Brown, and A.D. Freed, Eds., ASME, Book No. H00848, 1993.

This model represents a nonisothermal Bodner-Partom viscoplastic material with isotropic, Z^{I} , and directional hardening, Z^{D} , and can be used for an initially isotropic metallic material.

The flow law is given as:

$$\dot{\boldsymbol{\varepsilon}}_{ij}^{I} = \Lambda \boldsymbol{s}_{ij}$$

where

$$\Lambda = D_0 exp \left[-0.5 \left(\frac{Z^2}{3J_2} \right)^n \right] \frac{1}{\sqrt{J_2}}$$
$$Z = Z^I + Z^D$$
$$J_2 = \frac{1}{2} S_{ij} S_{ij}$$
$$S_{ij} = \sigma_{ij} - \frac{1}{3} \sigma_{kk} \delta_{ij}$$

The evolution law for the isotropic hardening, Z', is given as:

$$\dot{Z}^{I} = m_{1}(Z_{1} - Z^{I})\dot{W}^{PL} - A_{1}Z_{1}\left(\frac{Z^{I} - Z_{2}}{Z_{1}}\right)^{r_{2}} + \left(\left(\frac{Z^{I} - Z_{2}}{Z_{1} - Z_{2}}\right)\frac{\partial Z_{1}}{\partial T} + \left(\frac{Z_{2} - Z^{I}}{Z_{1} - Z_{2}}\right)\frac{\partial Z_{2}}{\partial T}\right)\dot{T}$$

where Z_0 , Z_1 , Z_2 , A_1 , r_1 and m_1 are the material parameters associated with isotropic hardening. The magnitude of the directional hardening is defined as the scalar product of a state variable, β_{ij} , and unit stress vector, u_{ij} , as given below.

$$Z^{D} = \beta_{ij} u_{ij}$$
$$\dot{\beta_{ij}} = m_2 (Z_3 u_{ij} - \beta_{ij}) \dot{W}^{PL} - A_1 Z_1 \left(\frac{\sqrt{\beta_{ij}\beta_{ij}}}{Z_1}\right)^{r_2} v_{ij} + \left(\left(\frac{\beta_{ij}}{Z_3}\right) \frac{\partial Z_3}{\partial T}\right) \dot{T}$$

$$u_{ij} = \frac{\sigma_{ij}}{\sqrt{\sigma_{kl}\sigma_{kl}}}$$
$$v_{ij} = \frac{\beta_{ij}}{\sqrt{\beta_{kl}\beta_{kl}}}$$

where, the plastic work rate is defined as; $W^{PL} = \sigma_{ij} \varepsilon_{ij}^{I}$ and with, Z_3 , A_2 , r_2 and m_2 being the material parameters defining the directional evolution. An elastic only response may be obtained by setting the material parameter D_0 to zero.

3.3.5 Robinson Viscoplastic Model

Reference: S.M. Arnold, D.N. Robinson, and P.A. Bartlotta, "Unified Viscoplastic Behavior of Metal Matrix Composites", NASA TM 105819, 1992

This model represents a **transversely isotropic material** wherein the vector of direction cosines, d_i , defines the preferred material direction. In this model the strength of anisotropy is specified by the parameters ω and η ; where ω is the ratio of the normal longitudinal and transverse yield stress and η is the ratio of longitudinal and transverse threshold shear stress.

Flow Law:

$$\dot{\varepsilon}_{ij}^{I} = \frac{\langle F^{n} \rangle}{2\mu} \Gamma_{ij}$$

Evolution Law:

$$\dot{a}_{ij} = \frac{H}{G^{\beta}} \dot{\varepsilon}_{ij}^{l} - RG^{m-\beta} \Pi_{ij}$$

where

$$\begin{split} \Gamma_{ij} &= \Sigma_{ij} - \xi (D_{ki} \Sigma_{jk} + D_{jk} \Sigma_{ki} - 2I_0 D_{ij}) - \frac{1}{2} \zeta I_0 (3D_{ij} - \delta_{ij}) \\ \Pi_{ij} &= a_{ij} - \xi (D_{ki} a_{jk} + D_{jk} a_{ki} - 2\hat{I}_0 D_{ij}) - \frac{1}{2} \zeta \hat{I}_0 (3D_{ij} - \delta_{ij}) \end{split}$$

and

$$F = \frac{1}{\kappa_T^2} \left[I_1 + \frac{1}{\eta^2} I_2 + \frac{9}{4(4\omega^2 - 1)} I_3 \right] - 1$$
$$\hat{G} = \frac{1}{\kappa_T^2} \left[\hat{I}_1 + \frac{1}{\eta^2} \hat{I}_2 + \frac{9}{4(4\omega^2 - 1)} \hat{I}_3 \right]$$
$$G = \langle \hat{G} - \hat{G}_0 \rangle H \nu [S_{ij} \pi_{ij}] + \hat{G}_0$$
$$I_1 = J_2 - I - \frac{1}{4} I_3 \qquad I_2 = I - I_3 \qquad I_3 = I_0^2$$

$$J_2 = \frac{1}{2} \sum_{ij} \sum_{ji} \qquad I = D_{ij} \sum_{ji} \qquad D_{ij} = d_i d_j \qquad \sum_{ij} = S_{ij} - a_{ij}$$
$$\xi = \frac{\eta^2 - 1}{\eta^2} \qquad \zeta = \frac{4(\omega^2 - 1)}{4\omega^2 - 1}$$

with κ_T , μ , n, H, β , R, m, G_o , η and ω representing the associated required inelastic material parameters and < > denoting a Macauley bracket, as defined below.

$$\langle \rangle = \left\{ \begin{array}{c} 0 \ x < 0 \\ x \ x \ge 0 \end{array} \right\}$$

The invariants \hat{I}_1 , \hat{I}_2 , \hat{I}_3 are the same I_1 , I_2 , I_3 as those given above but with Σ_{ij} replaced by a_{ij} . Special cases involving an isotropic material and/or elastic only response can be obtained by defining $\omega = \eta = 1$ and/or by setting κ_T to an extremely large number.

3.3.6 Generalized Viscoplastic Potential Structure (GVIPS) Model

References: S.M. Arnold, A.F. Saleeb, and M.G. Castelli, "A Fully Associative, Non-Linear Kinematic, Unified Viscoplastic Model for Titanium Based Matrices", NASA TM 106609, 1994.

S.M. Arnold, A.F. Saleeb, and M.G. Castelli, "A Fully Associative, Nonisothermal, NonLinear Kinematic, Unified Viscoplastic Model for Titanium Alloys", NASA TM 106926, 1994.

This model is a fully associative, multiaxial, nonlinear kinematic hardening viscoplastic model for use with initially isotropic metallic materials. A unique aspect of this model is the inclusion of non-linear hardening through the use of a compliance operator Q_{ijkl} in the evolution law for the back stress. This non-linear tensorial operator is significant in that it allows both the flow and evolutionary laws to be fully associative (and therefore easily integrated) and greatly influences the multiaxial response under non-proportional loading paths.

Flow Law:

$$\dot{\varepsilon}_{ij}^{I} = \frac{3}{2} \left\| \dot{\varepsilon}_{ij}^{I} \right\| \frac{\Sigma_{ij}}{\sqrt{J_2}} \qquad if \qquad F \ge 0$$

where

$$\left\|\dot{\varepsilon}_{ij}^{I}\right\| = \sqrt{\frac{2}{3}}\dot{\varepsilon}_{ij}^{I}\dot{\varepsilon}_{ij}^{I} = \frac{\mu F^{n}}{\kappa}$$

Internal constitutive rate equation:

$$\dot{a}_{ij} = L_{ijrs}\dot{A}_{rs}$$

Evolution Law:

$$\dot{A}_{rs} = \dot{\varepsilon}_{rs}^{I} - \frac{3}{2} \frac{\beta \kappa}{\kappa_{o}^{2}} \left\| \dot{\varepsilon}_{ij}^{I} \right\| \frac{a_{rs}}{\sqrt{G}} Hv[Y] - \frac{3R_{\alpha}B_{0}G^{4}}{\kappa_{o}^{2}} a_{rs} \quad if \quad a_{ij}\Sigma_{ij} \ge 0$$

$$\dot{A}_{rs} = Q_{rslm} E_{lmnp} \left(\dot{\varepsilon}_{np}^{I} - \frac{3}{2} \frac{\beta \kappa}{\kappa_{o}^{2}} \right\| \dot{\varepsilon}_{ij}^{I} \left\| \frac{a_{np}}{\sqrt{G}} Hv[Y] - \frac{3R_{\alpha}B_{0}G^{4}}{\kappa_{o}^{2}} a_{np} \right) \quad if \quad a_{ij}\Sigma_{ij} < 0$$

where

$$F = \left\langle \frac{\sqrt{J_2}}{\kappa} - Y \right\rangle$$
$$Y = \left\langle 1 - \beta \sqrt{G} \right\rangle$$
$$G = \frac{I_2}{\kappa_o^2}$$

$$L_{ijrs} = [Q_{ijrs}]^{-1} = \frac{\kappa_o^2}{3B_0(1+B_1pG^{p-1})} \left(I_{ijrs} - \frac{3B_1p(p-1)G^{p-2}}{\kappa_o^2(1+B_1pG^{p-1}(6p-5))} a_{rs}a_{ij} \right)$$

and

$$I_{2} = \frac{3}{2}a_{ij}a_{ij} \qquad \Sigma_{ij} = S_{ij} - a_{ij}$$
$$J_{2} = \frac{3}{2}\Sigma_{ij}\Sigma_{ij} \qquad S_{ij} = \sigma_{ij} - \frac{1}{3}\sigma_{kk}\delta_{ij}$$

with $\kappa, \mu, n, \kappa_o, B_0, B_1, p, R_\alpha, q$ and β being the associated required inelastic material parameters. Typically, $\kappa, \mu, B_0, R_\alpha$ and β are taken to be functions of temperature and $\kappa_o = \kappa(T_{ref})$ is the initial drag stress at the reference temperature. The special case of an elastic only response maybe obtained by setting κ to an extremely large value.

3.3.7 Transversely Isotropic GVIPS Model (TGVIPS)

Reference: A.F. Saleeb and T.E. Wilt, Analysis of the Anisotropic Viscoplastic-Damage Response of Composite Laminates-Continuum Basis and Computational Algorithms, Int. J. Numer. Meth. Enging., Vol. 36, pp. 1629-1660, 1993.

This model is a fully associative, multiaxial, isothermal, nonlinear kinematic hardening viscoplastic model for use with **initially transversely isotropic** metallic materials. A unique aspect of this model is the inclusion of non-linear hardening through the use of a compliance operator Q_{ijkl} in the evolution law for the back stress. This non-linear tensorial operator is significant in that it allows both the flow and evolutionary laws to be fully associative (and therefore easily integrated) and greatly influences the multiaxial response under non-proportional loading paths.

The flow law for the inelastic strain, $\dot{\epsilon}^{\prime}$, is given by,

$$\dot{\boldsymbol{\xi}}^{I} = \frac{\langle \boldsymbol{F}^{n} \rangle}{2\mu} \boldsymbol{\Gamma}$$

and the evolution law for internal stress, α is given by,

$$\dot{\alpha} = \left[\tilde{Z}_{m} + \frac{h'}{h(1+2\beta)} (\tilde{\alpha} \otimes \tilde{\alpha}) \right] \left(\frac{H}{G^{\beta}} \dot{\varepsilon}' - RG^{m} \tilde{\Pi} \right)$$

where h = $H/G^{\beta},\,h'$ = $-\beta/G$ and Z_m = \underline{M}^{-1}

$$F = \frac{1}{2\kappa_t^2} (\tilde{g} - \tilde{g}) : \tilde{M} : (\tilde{g} - \tilde{g}) - 1$$

$$G = \frac{1}{2\kappa_t^2} \tilde{g} : \tilde{M} : \tilde{g}$$

$$\tilde{\Gamma} = \tilde{M} : (\tilde{g} - \tilde{g})$$

$$\tilde{\Pi} = \tilde{M} : \tilde{g}$$

$$\tilde{M} = \tilde{P} - \xi \tilde{Q} - \frac{1}{2} \zeta \tilde{R}$$

The anisotropy of the material is introduced through the M matrix, specifically the parameters, ζ , ξ which are defined as,

$$\xi = \frac{\eta^2 - 1}{\eta^2} \qquad \zeta = \frac{4(\omega^2 - 1)}{4\omega^2 - 1}$$
$$\eta = \frac{K_l}{K_l} \qquad \omega = \frac{Y_l}{Y_l}$$

In the above, $0 \le \xi \le 1$ and $0 \le \zeta \le 1$ are the material strength ratios, in which the constants K_l (K_t) are the threshold strengths in longitudinal (transverse) shear, and Y_l (Y_t) are the threshold strengths in longitudinal (transverse) tension. Note that for an isotropic material, $\omega = \eta = 1$.

In addition, the fourth-order tensors \underline{P} , \underline{Q} , and \underline{R} are defined as,

$$\begin{split} \mathbf{P} &= \mathbf{I} - \frac{1}{3} (\mathbf{\delta} \otimes \mathbf{\delta}) \qquad \mathbf{I}_{ijkl} = \frac{1}{2} (\delta_{ik} \delta_{ji} + \delta_{il} \delta_{jk}) \\ \mathbf{Q}_{ijkl} &= \frac{1}{2} (D_{ik} \delta_{jl} + D_{il} \delta_{jk} + D_{jk} \delta_{il} + D_{jl} \delta_{ik}) - 2D_{ij} D_{kl} \\ \mathbf{R}_{ijkl} &= 3D_{ij} D_{kl} - (D_{ij} \delta_{kl} + \delta_{ij} D_{kl}) + \frac{1}{3} (\delta_{ij} \delta_{kl}) \end{split}$$

In the above, the vector of direction cosines d_i defines the orientation of the material fiber, which leads to the material directionality tensor D, $(D = d_i d_j)$. Also, δ is the Kronecker delta (second-order identity tensor) and I is the fourth-order identity tensor. Finally, the symbol : indicates double-contraction and \otimes cross product.

Note: When calculating Z_m for the three dimension case one needs to replace \underline{P} with $\underline{\hat{P}} = \text{diag}[1,1,1,2,2,2]$ as \underline{P} is singular for the three dimensional case.

3.4 Laminate Theory

MAC/GMC includes the capability to analyze general (symmetric and nonsymmetric) composite laminates [9], see Fig. 4. Mid-plane strains and resultant forces in the plane of the laminate may be applied. That is, the global laminate stress-strain relation that is solved within **MAC/GMC** is expressed as,

$$\begin{bmatrix} \overline{N}_{XX} \\ \overline{N}_{YY} \\ \overline{N}_{XY} \end{bmatrix} = \begin{bmatrix} A_{11} & A_{12} & A_{13} \\ A_{21} & A_{22} & A_{23} \\ A_{31} & A_{32} & A_{33} \end{bmatrix} \begin{bmatrix} \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{yy} \\ \overline{\gamma}_{xy} \end{bmatrix} + \begin{bmatrix} B_{11} & B_{12} & B_{13} \\ B_{21} & B_{22} & B_{23} \\ B_{31} & B_{32} & B_{33} \end{bmatrix} \begin{bmatrix} \overline{\kappa}_{xx} \\ \overline{\kappa}_{yy} \\ \overline{\kappa}_{xy} \end{bmatrix} - \begin{bmatrix} \overline{N}_{XX}^{T} \\ \overline{N}_{YY}^{T} \\ \overline{N}_{XY}^{T} \end{bmatrix}$$
(EQ 15)
$$\begin{bmatrix} \overline{M}_{XX} \\ \overline{N}_{YY} \\ \overline{N}_{XY} \end{bmatrix} = \begin{bmatrix} B_{11} & B_{12} & B_{13} \\ B_{12} & B_{13} \\ \overline{\epsilon}_{xx} \end{bmatrix} \begin{bmatrix} \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{xy} \end{bmatrix} + \begin{bmatrix} D_{11} & D_{12} & D_{13} \\ D_{11} & D_{12} & D_{13} \\ \overline{\epsilon}_{xx} \end{bmatrix} \begin{bmatrix} \overline{M}_{XX} \\ \overline{k}_{xy} \\ \overline{k}_{xy} \end{bmatrix} = \begin{bmatrix} \overline{M}_{XX} \\ \overline{k}_{xy} \\ \overline{k}_{xy} \end{bmatrix} = \begin{bmatrix} B_{11} & B_{12} & B_{13} \\ \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{xx} \end{bmatrix} = \begin{bmatrix} B_{11} & B_{12} & B_{13} \\ \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{xx} \\ \overline{\epsilon}_{xx} \end{bmatrix} = \begin{bmatrix} B_{11} & B_{12} & B_{13} \\ \overline{\epsilon}_{xx} \\ \overline$$

$$\begin{bmatrix} \overline{M}_{YY} \\ \overline{M}_{YY} \\ \overline{M}_{XY} \end{bmatrix} = \begin{bmatrix} B_{21} & B_{22} & B_{23} \\ B_{31} & B_{32} & B_{33} \end{bmatrix} \begin{bmatrix} \overline{k}_{yy} \\ \overline{\gamma}_{xy} \end{bmatrix} + \begin{bmatrix} D_{21} & D_{22} & D_{23} \\ D_{31} & D_{32} & D_{33} \end{bmatrix} \begin{bmatrix} \overline{k}_{yy} \\ \overline{k}_{xy} \end{bmatrix} - \begin{bmatrix} \overline{M}_{YY}^{I} \\ \overline{M}_{XY}^{I} \end{bmatrix} - \begin{bmatrix} \overline{M}_{YY}^{T} \\ \overline{M}_{XY}^{T} \end{bmatrix}$$
(EQ 16)

or

$$\begin{bmatrix} \overline{N} \\ \overline{M} \\ \overline{M} \end{bmatrix} = \begin{bmatrix} A & B \\ B & D \end{bmatrix} \begin{bmatrix} \overline{e} \\ \overline{k} \\ \overline{k} \end{bmatrix} - \begin{bmatrix} \overline{N}^{I} \\ \overline{M}^{I} \end{bmatrix} - \begin{bmatrix} \overline{N}^{T} \\ \overline{M}^{T} \end{bmatrix}$$
(EQ 17)

where \overline{N} , \overline{N}^{I} , \overline{N}^{T} , and \overline{M} , \overline{M}^{I} , \overline{M}^{T} are the global laminate total, inelastic and thermal force and moment resultants, respectively. The matrices \underline{A} , \underline{B} , and \underline{D} are the global laminate extensional, coupling and bending stiffnesses, respectively, and, $\overline{\underline{e}}$ and $\overline{\underline{K}}$ the global laminate mid-plane strain and mid-plane curvature, respectively.

In forming the laminate extensional stiffness \underline{A} the generalized method of cells model, **GMC**, is utilized to calculate the individual lamina properties. In this regard, the individual laminate stiffness, in lamina coordinates, \underline{Q} is given by,

in which the components of Q are given as,

$$Q_{11} = C_{11} - \frac{C_{13}C_{31}}{C_{33}} \qquad Q_{12} = C_{12} - \frac{C_{13}C_{23}}{C_{33}}$$

$$Q_{22} = C_{22} - \frac{C_{23}C_{32}}{C_{33}} \qquad Q_{33} = C_{66}$$
(EQ 19)

The C_{ij} in the above are the <u>effective</u> macro properties for the unidirectional composite lamina obtained from **GMC**.

It follows, employing a Kirchhoff-Love hypothesis, that the lamina stress-strain relation in global (laminate) coordinates denoted by x-y is given by the relation,

$$\begin{bmatrix} \sigma_{x} \\ \sigma_{y} \\ \sigma_{xy} \end{bmatrix}_{k} = \begin{bmatrix} \overline{Q}_{11} \ \overline{Q}_{12} \ \overline{Q}_{13} \\ \overline{Q}_{12} \ \overline{Q}_{22} \ \overline{Q}_{23} \\ \overline{Q}_{13} \ \overline{Q}_{23} \ \overline{Q}_{33} \end{bmatrix}_{k} \begin{bmatrix} \overline{\varepsilon}_{xx} \\ \overline{\varepsilon}_{yy} \\ \overline{\gamma}_{xy} \end{bmatrix} + z \begin{bmatrix} \overline{\kappa}_{xx} \\ \overline{\kappa}_{yy} \\ \overline{\kappa}_{xy} \end{bmatrix} - \begin{bmatrix} \varepsilon_{xx}^{I} \\ \varepsilon_{yy}^{I} \\ \varepsilon_{yy}^{I} \\ \gamma_{xy}^{I} \end{bmatrix}_{k} \begin{bmatrix} \overline{\varepsilon}_{xx} \\ \varepsilon_{xy}^{T} \\ \gamma_{xy}^{I} \end{bmatrix}_{k} \begin{bmatrix} \varepsilon_{xx} \\ \varepsilon_{xy} \\ \varepsilon_{yy}^{I} \\ \gamma_{xy}^{I} \end{bmatrix}_{k} \begin{bmatrix} \varepsilon_{xx} \\ \varepsilon_{xy} \\ \varepsilon_{yy}^{I} \\ \gamma_{xy}^{I} \end{bmatrix}_{k} \begin{bmatrix} \varepsilon_{xx} \\ \varepsilon_{xy} \\ \varepsilon_{yy} \\ \gamma_{xy}^{I} \end{bmatrix}_{k} \end{bmatrix}$$
(EQ 20)

or

$$\mathfrak{g}_{k} = \overline{Q}_{k} (\overline{\mathfrak{g}} + z \overline{\mathfrak{K}} - \mathfrak{g}_{k}^{T} - \mathfrak{g}_{k}^{T})$$
(EQ 21)

where

$$\overline{\underline{Q}_{k}} = R_{k}^{-1} \underline{Q}_{k} R_{k}$$
(EQ 22)

$$R_{k} = \begin{bmatrix} \cos^{2}\theta & \sin^{2}\theta & 2\sin\theta\cos\theta \\ \sin^{2}\theta & \cos^{2}\theta & (-2)\sin\theta\cos\theta \\ -\sin\theta\cos\theta & \sin\theta\cos\theta & \cos^{2}\theta - \sin^{2}\theta \end{bmatrix}$$
(EQ 23)

and θ is the orientation of the longitudinal lamina axis with respect to the global xdirection, see Fig. 4, thus \overline{Q}_k is the transformed lamina stiffness, i.e. from local lamina to global laminate coordinates. In addition, σ_k is the lamina stress vector in laminate coordinates. It then follows that the global laminate extensional stiffness \underline{A} is given by,

$$\underline{A} = \sum_{k=1}^{nly} \overline{\underline{Q}}_k t_k \tag{EQ 24}$$

in which nly is the total number of layers in the laminate and t_k is the thickness of the k^{th} lamina. The coupling and bending stiffnesses can be similarly developed and given by the following expressions:

$$\underline{B} = \frac{1}{2} \sum_{k=1}^{nly} \overline{\underline{Q}}_k (z_k^2 - z_{k-1}^2)$$
(EQ 25)

$$\bar{D} = \frac{1}{3} \sum_{k=1}^{nly} \bar{Q}_k (z_k^3 - z_{k-1}^3)$$
(EQ 26)

where z_k is the distance (considering the sign) to the top of layer k from the midplane.

Returning to EQ. 17, the quantities \overline{N}^{I} and \overline{N}^{T} (the laminate inelastic and thermal force resultants, respectively) are calculated from the individual lamina contributions through the following relations,

$$\overline{\underline{N}}^{I} = \sum_{k=1}^{nly} \overline{\underline{Q}}_{k} \int_{z_{k-1}}^{z_{k}} \underline{\varepsilon}_{k}^{I} dz \qquad \overline{\underline{M}}^{I} = \sum_{k=1}^{nly} \overline{\underline{Q}}_{k} \int_{z_{k-1}}^{z_{k}} \underline{\varepsilon}_{k}^{I} z dz \qquad (EQ 27)$$

and

$$\overline{\underline{N}}^{T} = \sum_{k=1}^{nly} \overline{\underline{Q}}_{k} \underline{\underline{\varepsilon}}_{k}^{T} t_{k} \qquad \overline{\underline{M}}^{T} = \frac{1}{2} \sum_{k=1}^{nly} \overline{\underline{Q}}_{k} \underline{\underline{\varepsilon}}_{k}^{T} (z_{k}^{2} - z_{k-1}^{2})$$
(EQ 28)

where the integrals in EQ. 27 are performed using second order gauss quadrature which requires two integration points per layer. Thus all field quantities are tracked at the two gauss quadrature points in each layer of the laminate in **MAC/GMC**.



Figure 4: Laminate Coordinate System

3.5 Fatigue Damage Analysis

The fatigue damage calculations utilize a recently developed multiaxial, isothermal, continuum damage mechanics model for the fatigue of unidirectional metal matrix composites [10]. The model is phenomenological, stress based, and assumes a single scalar internal damage variable, *D*. Note that for an initially anisotropic material, the evolution of the damage, although a scalar, is directionally dependent. As will be shown, this directional dependence is accounted for in the terms, \hat{F}_m , Φ_{fl} , and Φ_u . The present multiaxial, isothermal, continuum damage model for **initially transversely isotropic materials** (e.g., unidirectional metal matrix composites) may be expressed as, [10]

$$\int_{D_{k-1}}^{D_k} dD = \int_{0}^{N} [1 - (1 - D)^{\beta + 1}]^{\alpha} \left[\frac{\hat{F}_m}{1 - D}\right]^{\beta} dN$$
 (EQ 29)

where N is the number of cycles at the current stress state, (σ_k) , and D_k and D_{k-1} are the amount of damage at the current and previous increments, respectively. The quantity α which is a function of the current stress state is defined as,

$$\alpha = 1 - a \frac{\langle \Phi_{fl} \rangle}{\langle \Phi_{\mu} \rangle} \tag{EQ 30}$$

where $\langle \rangle$ are the Macauley brackets. In the above, the fatigue limit surface, Φ_{fl} , and the static fracture surface, Φ_{μ} , are defined as

$$\Phi_{fl} = \frac{1}{2} \frac{max}{t_0} \frac{max}{t} F_{(\sigma_{fl})}(\sigma_{ij}(t) - \sigma_{ij}(t_0)) - 1$$
(EQ 31)

$$\Phi_u = 1 - \frac{max}{t} F_{(\sigma_u)}(\sigma_{ij}(t))$$
(EQ 32)

and the quantity \hat{F}_m , used in EQ. 29 is the normalized stress amplitude, and is defined as,

$$\hat{F}_{m} = \frac{1}{2} \frac{max}{t} \frac{max}{t_{0}} F_{(M)}(\sigma_{ij}(t) - \sigma_{ij}(t_{0}))$$
(EQ 33)

Note, the case $\langle \Phi_u \rangle = 0$ indicates static fracture, which is failure, making it unnecessary to perform the fatigue calculations as in this case the subcell is considered to have failed completely. Thus, having to consider the possibility of α being undefined is unnecessary. The case $\langle \Phi_{fl} \rangle = 0$ indicates that the current stress state is below the fatigue limit and thus α is set equal to 1. This presents a special case when integrating the fatigue damage expression, EQ. 29, and will be considered later in this section.

In the above equations, t_0 is the time at the beginning of the current load cycle, and t, is some time during the load cycle. The general form for $F_{(\sigma_{fl}), (\sigma_u), or (M)}$ may be expressed as,

$$F_{()} = \sqrt{\frac{1}{\left(\right)_{L}^{2}} \left\{ (4\omega_{()}^{2} - 1)I_{1} + \frac{4\omega_{()}^{2} - 1}{\eta_{()}^{2}}I_{2} + \frac{9}{4}I_{3} \right\}}$$
(EQ 34)

It is in the above expression, in which the evolution of the damage becomes directionally dependent. This simply amounts to the assumption of partial anisotropy, where the "extent" (magnitude) of damage is affected by the directionality of the stress state. Specifically, the directional dependence enters through the quantities, I_1 , I_2 , I_3 , $\omega_{()}$, and $\eta_{()}$. The quantities, I_1 , I_2 , I_3 are invariants having the form,

$$I_{1} = \frac{1}{2} S_{ij} S_{ij} - d_{i} d_{j} S_{jk} S_{ki} + \frac{1}{4} (d_{i} d_{j} S_{ij})^{2}$$

$$I_{2} = d_{i} d_{j} S_{jk} S_{ki} - (d_{i} d_{j} S_{ij})^{2}$$

$$I_{3} = (d_{i} d_{j} S_{ij})^{2}$$
(EQ 35)

which are a function of the current deviatoric stress state, $s_{ij}^k = \sigma_{ij}^k - \frac{1}{3}\sigma_{mm}^k \delta_{ij}$, as well as the vector d_i which defines the materials' preferred direction (e.g., fiber orientation in a composite). In addition, the terms $\omega_{(\)}$ and $\eta_{(\)}$ represent the ratios of longitudinal to transverse normal and shear stresses, respectively. Note, the longitudinal direction is parallel to the preferred direction and transverse is perpendicular to the preferred direction. For initially transversely isotropic materials, $\omega_{(\)}$ and $\eta_{(\)}$ are > 1 and for isotropic materials $\omega_{(\)}$ and $\eta_{(\)}$ are = 1.

In the context of micromechanics analysis within **MAC/GMC**, the isotropic simplification of the above representation will be predominately used for the various constituent phases [11]. This isotropic representation is the previously validated NonLinear Cumulative Damage Rule (NLCDR) developed at **ONERA** (Office Nationale d'Etudes et de Recherches Aerospatiales) for isotropic monolithic metals. However, it maybe desirable to use the transverse isotropic form when dealing with fiber tows in woven composites systems.

3.5.1 Above Initial Fatigue Limit

Given a current state of stress, σ_k , above the fatigue limit, i.e. $\alpha \neq 1$ and integrating EQ. 29 results in an expression for the number of cycles, *N*, i.e.,

$$N = \frac{\left(\left[1 - \left(1 - D_{k}\right)^{\beta+1}\right]^{1-\alpha} - \left[1 - \left(1 - D_{k-1}\right)^{\beta+1}\right]^{1-\alpha}\right)}{\hat{F}_{m}^{\beta}(1-\alpha)(\beta+1)}$$
(EQ 36)

Note that D_{k-1} is the total amount of damage at the beginning of the load block and D_k is the total amount of damage at the end of this load block. Alternatively, rewriting EQ. 36 an expression for the damage, D_k , in terms of the number of cycles and previous damage can be obtained, i.e.,

$$D_{k} = 1 - \left(1 - \left\{\left[1 - (1 - D_{k-1})^{\beta+1}\right]^{1-\alpha} + (1 - \alpha)(\beta + 1)\hat{F}_{m}^{\beta}N\right\}^{\frac{1}{1-\alpha}}\right)^{\frac{1}{\beta+1}}$$
(EQ 37)

In the present computational scheme, since the damage increment is controlled, both D_k and D_{k-1} are known. That is, $D_k = D_{k-1} + \Delta D$ where ΔD is the user specified increment in damage. Thus EQ. 36 is used to predict the increment in the number of cycles for each subcell, N^e , due to the imposed increment in damage.

To calculate the number of cycles to failure, for an initial damage amount, D_{k-1} , let $D_k = 1$, which results in the following,

$$N_F = \frac{\left(1 - \left[1 - \left(1 - D_{k-1}\right)^{\beta - 1}\right]^{1 - \alpha}\right)}{\hat{F}_m^{\ \beta} (1 - \alpha)(\beta + 1)}$$
(EQ 38)

3.5.2 Below Initial Fatigue Limit

Now consider the case in which the current stress state is below the initial fatigue limit, i.e. $\langle \Phi_{fl} \rangle = 0$, which leads to $\alpha_k = 1$. Thus, EQ. 29 takes the form,

$$\int_{D_{k-1}}^{D_k} \frac{(1-D)^{\beta}}{1-(1-D)^{\beta+1}} dD = \int_{0}^{N} \hat{F}_m^{\beta} dN$$
 (EQ 39)

Upon integrating the above equation, the increment in cycles, N, with initial damage, D_{k-1} , may be expressed as,

$$N = \left(\frac{\log[1 - (1 - D_k)^{\beta + 1}] - \log[1 - (1 - D_{k-1})^{\beta + 1}]}{\hat{F}_m^{\beta}(\beta + 1)}\right)$$
(EQ 40)

Alternatively, the following expression for the damage, D_k , may be expressed as:

$$D_{k} = 1 - \{1 - [1 - (1 - D_{k-1})^{\beta+1}]exp((\beta+1)\hat{F}_{m}^{\beta}N)\}^{\frac{1}{\beta+1}}$$
(EQ 41)

For the number of cycles to failure, let $D_k = 1$,

$$N_F = \frac{-\log[1 - (1 - D_{k-1})^{\beta+1}]}{\hat{F}_m^{\beta}(\beta+1)}$$
(EQ 42)

The effect of damage is included in the present micromechanics analysis utilizing the concept of <u>effective stress</u> and the hypothesis of strain-equivalence [12].

$$\hat{g} = \frac{g}{(1 - D_k)} \tag{EQ 43}$$

3.6 User Defined Inelastic Material Model

MAC/GMC, has the option for a user to implement their own inelastic constitutive model. This is accomplished by using the subroutine **USRMAT** into which the user writes the necessary FORTRAN code for the particular constitutive model being implemented. The **USRMAT** subroutine is as shown below and is always called whenever ncmd=99 (see section **4.2.12**).

```
SUBROUTINE USRMAT (DSA, SA, PE, PV, D, LOCTISO, TIME, TSTEP,
   ÷
        CTEMP, DTEMPR, NIO, NE, NV, NS, MN, CDUM, DMGF,
   &
        NEP, NVP, NSASIZE)
           user material constitutive model for determination of
С
    purpose:
            the inelastic strain and state variable rates
С
С
            (used when ncmd = 99)
    IMPLICIT DOUBLE PRECISION (A - H, O - Z)
    CHARACTER*2 CDUM
    DIMENSION SS(6), S(6), R(6)
    DIMENSION DSA(NSASIZE), SA(NSASIZE)
    DIMENSION PV(NVP), PE(NEP), D(3)
************************
 note: 1) in this subroutine, [SA] and [DSA] contain the
С
С
        micro (subcell) quantities for aboudi's micromechanics model
С
С
      2) arrangement of [dsa] & [sa] arrays:
        variable
С
                       location
С
       Ç
       | strain rate
                       (1-6) (contains ENGINEERING shears)
С
        С
       stress rate
                        (7 - 12)
С
       | inelastic
С
С
       strain rate
                        (13-18) (contains ENGINEERING shears)
С
       С
       12 "slots"
                        (19 - 30)
С
       | for state variables
С
       |-----
С
       thermal strain rate (31-36)
       +-----
С
         ***
c NOTE: quantities in [SA] and [DSA] are SUBCELL quantities - the
      values on entry are for the first subcell containing material
С
С
      # MN - the values on exit of this subroutine will be applied to
      ALL SUBCELLS containing material # MN. It is thus recommended
С
      that, if using the field variables, you assign the appropriate
С
      material # to ONE SUBCELL ONLY. Use of [SA] and [DSA] in this
С
С
      context in conjunction with bending in laminate theory will
С
      result in erroneous results as field variables become dependent
```
C	on throug	h-thickness position.
~ * *	on ontru:	
c c	SA	- vector of total (integrated) quantities (see above)
c	PE(NEP)	- vector of elastic constants for material MN
c c		(where NE = # of elastic constants > 9 MAX)
c c	PV (NVP)	- vector of viscoplastic constants for material MN
c	- • ((where NV = # of viscoplastic constants> 19 MAX)
c	D(3)	- vector of direction cosines (for models 3, 7, & 9)
c	LOCTISO	- flag indicating if ANY material exhibits
c		local transverse isotropy (and global anisotropy)
с		= 0 - all materials are at most globally transversely
с		isotropic (D not used)
с		= 1 - at least one material is locally transversely
с		isotropic (D used)
с	TIME	- current time
с	TSTEP	- current time step
с	CTEMP	- current temperature
с	DTEMPR	 time rate of change of temperature
с	NIO	- unit number of .out file
с	NE	- # of elastic constants> 9 MAX
с	NV	- # of viscoplastic constants> 19 MAX
С	NS	- subcell number
С	MN	- material number
с	CDUM	- material character/number designation
с		(i.e. Fl = fiber #1)
с	DMGF	- damage factor - if damage is included the user
С		should multiply material stillness terms by Digr
С		when using such terms in their inerastic model.
С		
С	expected on e	XIC:
C ***	USA ************	- Vector of face qualities (see above)
с		
***	*****	* * * * * * * * * * * * * * * * * * * *
*		BEGIN USER EDITS *
***	* * * * * * * * * * * * * * * *	* * * * * * * * * * * * * * * * * * * *
С	place co	de defining model here
***	* * * * * * * * * * * * * * * *	***************************************
*		END USER EDITS *
1	* * * * * * * * * * * * * * *	**********
	RETURN	
	END	

A description of the input and output required for the usrmat subroutine is as follows

Data supplied to USRMAT:

SA	array containing current total quantities for all
	of the state variables
PE	array containing elastic constants

PV D(3) LOCTISO	array containing inelastic constants vector of direction cosines (for models 3, 7, & 9) flag indicating if ANY material exhibits local transverse isotropy (and global anisotropy) = 0 - all materials are at most globally transversely isotro- pic (D not used) = 1 - at least one material is locally transversely isotropic (D used)
TIME TSTEP CTEMP DTEMPR NE NV NIO NCE NS MN CDUM DMGF	current time current time increment (step) current temperature current temperature rate total number of elastic constants total number of inelastic constants output file unit number current subcell number subcell number material number of current constituent Character string identifying current constituent damage factor - if damage is included the user should multiply material stiffness terms by DMGF when using such terms in their inelastic model.

Output expected from USRMAT:

DSA current increments in all state variables

The state variables are arranged in SA and DSA in the following order:

Position	Quantity
1-6	strain
7-12	stress
13-18	inelastic strain
19-30	available space for 2 6x1 vectors for model dependent internal state variables
31-36	thermal strain
37-42	debonding parameters

Again, SA contains the total quantities and DSA contains the rates.

- Note: Example K contains a sample input file and the USRMAT subroutine containing an implementation of the Bodner-Partom viscoplastic material model as well as two elastic models.
- Note: GMC utilizes engineering strains, in SA and DSA consequently it is the users responsibility to convert tensorial strain quantities to engineering before exiting the USRMAT routine.

3.7 User Defined Functional Form For Material Properties

An addition to this version of **MAC/GMC** is the option for a user to implement functionally dependent material properties. With this the elastic and/or inelastic material properties may be defined as a function of any variable contained in SA and DSA (e.g., stress, strain, time, temperature, etc.) In certain cases, the user must also provide code for determining the stiffness matrix (USRFORMDE) or the time derivative of the stiffness matrix (USRCPEVAL). This is accomplished by using the subroutines **USRFUN, USRFORMDE, and USRCPEVAL** into which the user writes the necessary FORTRAN code for the particular constitutive model being implemented. All three routines are shown below.

```
SUBROUTINE USRFUN (MN, TIME, TSTEP, CTEMP, DTEMPR, SA, DSA,
       DOLD, PEM, PVM, D, LOCTISO, ALPA, ALPT,
   æ
       NE, NV, NMTS, NEP, NVP, NSASIZE)
   &
    purpose: user subroutine to allow elastic and viscoplastic
С
           material properties to be functions of TEMP or
С
           field variables. Used for user defined functional
С
           form material properties, that is, when:
С
           (mat .eq. 'U') .and. (ifm .eq. 2)
С
С
           can be used in conjunction with a provided material
С
   note:
           constitutive model, or a constitutive model input
С
С
           by the user in USRMAT
    INCLUDE 'parm.inc'
    IMPLICIT DOUBLE PRECISION (A - H, O - Z)
    DIMENSION DOLD(6, 6), D(3)
    DIMENSION PEM(NEP, NMTS), PVM(NVP, NMTS)
    DIMENSION ALPA(NMTS), ALPT(NMTS)
    DIMENSION DSA (NSASIZE), SA (NSASIZE)
******
                            c note: 1) in this subroutine, [SA] and [DSA] contain the
С
        micro (subcell) quantities for aboudi's micromechanics model
С
С
      2) arrangement of [dsa] & [sa] arrays:
        variable
С
                       location
С
       +------
                       (1-6) (contains ENGINEERING shears)
       | strain rate
С
       С
С
       stress rate
                       (7 - 12)
       С
       | inelastic
С
       strain rate
С
                     (13-18) (contains ENGINEERING shears)
С
       _____
       12 "slots"
С
                       (19-30)
С
       for state variables
       |------
С
С
       thermal strain rate (31-36)
С
```

c NOTE: quantities in [SA] and [DSA] are SUBCELL quantities - the values on entry are for the first subcell containing material С с # MN - the values on exit of this subroutine will be applied to С ALL SUBCELLS containing material # MN. It is thus recommended that, if using the field variables, you assign the appropriate С С material # to ONE SUBCELL ONLY. Use of [SA] and [DSA] in this context in conjunction with bending in laminate theory will С С result in erroneous results as field variables become dependent С on through-thickness position. ***** ****** С on entry: С MN - material number С TIME - current time С TSTEP - current time step С CTEMP - current temperature DTEMPR С - time rate of change of temperature С SA - vector of total (integrated) quantities (see above) С DSA - vector of rate quantities (see above) С DOLD(6, 6) - previous elastic material stiffness matrix PEM(NE, MN) - vector of previous elastic constants for material С С # MN (where NE = # of elastic constants --> 9 MAX) PVM(NV, MN) - vector of previous viscoplastic constants for C С material # MN С (where NV = # of viscoplastic constants --> 19 MAX) С expected on exit: С PEM(NE, MN) - vector of current elastic constants for material MN С PVM(NV, MN) - vector of current viscoplastic constants for С С material MN D(3) С - vector of direction cosines С (required for models 3, 7, & 9) - flag indicating if ANY material exhibits С LOCTISO С local transverse isotropy (and global anisotropy) = 0 - all materials are at most globally transversely С С isotropic (D not used) = 1 - at least one material is locally transversely С С isotropic (D used) С ALPA(MN) - longitudinal cte for material MN C ALPT (MN) - transverse cte for material MN ******************* BEGIN USER EDITS С place code here END USER EDITS RETURN END

```
SUBROUTINE USRFORMDE (MN, PEM, PVM, D, LOCTISO, DNEW,
        NE, NV, NEP, NVP, NMTS)
   æ
    purpose: user subroutine to allow formation of material stiffness
С
            matrices based on a user constitutive model (used when
С
С
            ncmd = 99
С
    INCLUDE 'parm.inc'
С
    IMPLICIT DOUBLE PRECISION (A - H, O - Z)
    DIMENSION DNEW(6, 6)
    DIMENSION PEM(NEP, NMTS), PVM(NVP, NMTS)
    DIMENSION D(3)
on entry:
С
              - material number
С
     MN
     PEM(NE, MN) - vector of elastic constants for material
С
                # MN (where NE = # of elastic constants --> 9 MAX)
С
     PVM(NV, MN) - vector of viscoplastic constants for material # MN
С
                (where NV = # of viscoplastic constants --> 19 MAX)
с
              - vector of direction cosines
    D(3)
¢
               (required for models 3, 7, & 9)
С
              - flag indicating if ANY material exhibits
     LOCTISO
С
                local transverse isotropy (and global anisotropy)
С
              = 0 - all materials are at most globally transversely
С
                   isotropic (D not used)
С
              = 1 - at least one material is locally transversely
С
                   isotropic (D used)
С
С
С
   expected on exit:
     DNEW(6, 6) - current elastic material stiffness matrix
С
******
                 BEGIN USER EDITS
                               *****
******
                       place code here
С
END USER EDITS
```

RETURN END

***** SUBROUTINE USRCPEVAL (DSA, SA, MN, TIME, TSTEP, CTEMP, DTEMPR, DNEW, DOLD, PEM, PVM, D, LOCTISO, ALPA, ALPT, DDOT, & æ NE, NV, NMTS, NEP, NVP, NSASIZE) С purpose: user subroutine to allow formation of the TIME derivative of the material stiffness matrix. С this subroutine is used when: С a) material properties are user defined and functional С form. That is: (mat .eq. 'U') .and. (ifm .eq. 2) С b) the constitutive model is user-defined, and the С material properties are not functional form, and С С the material properties are temperature-dependent С that is: (ncmd .eq. 99) .and. (ifm .ne. 2) .and. С (ndpt .eq. 2) IMPLICIT DOUBLE PRECISION (A - H, O - Z) DIMENSION DNEW(6, 6), DOLD(6, 6) DIMENSION DDOT(6, 6) DIMENSION PEM(NEP, NMTS), PVM(NVP, NMTS) DIMENSION ALPA(NMTS), ALPT(NMTS) DIMENSION DSA(NSASIZE), SA(NSASIZE) c note: 1) in this subroutine, [SA] and [DSA] contain the С micro (subcell) quantities for aboudi's micromechanics model С 2) arrangement of [dsa] & [sa] arrays: С location С variable +-----С (1-6) (contains ENGINEERING shears) С strain rate С |------С stress rate (7 - 12)С С | inelastic | strain rate С (13-18) (contains ENGINEERING shears) С | 12 "slots" С (19 - 30)| for state variables С |-------С С | thermal strain rate (31-36) +-----С c NOTE: quantities in [SA] and [DSA] are SUBCELL quantities - the values on entry are for the first subcell containing material С # MN - the values on exit of this subroutine will be applied to C С ALL SUBCELLS containing material # MN. It is thus recommended С that, if using the field variables, you assign the appropriate material # to ONE SUBCELL ONLY. Use of [SA] and [DSA] in this С context in conjunction with bending in laminate theory will С result in erroneous results as field variables become dependent С С on through-thickness position.

С on entry: - vector of total (integrated) quantities (see above) SA С - vector of rate quantities (see above) DSA С - material number MN С С TIME - current time TSTEP - current time step С - current temperature CTEMP С DTEMPR - time rate of change of temperature С DNEW(6, 6) - current elastic material stiffness matrix С DOLD(6, 6) - previous elastic material stiffness matrix С PEM(NE, MN) - vector of current elastic constants for material MN С (where NE = # of elastic constants --> 9 MAX) С PVM(NV, MN) - vector of current viscoplastic constants for С material MN С (where NV = # of elastic constants --> 19 MAX) С D(3) - vector of direction cosines С (required for models 3, 7, & 9) С - flag indicating if ANY material exhibits LOCTISO С local transverse isotropy (and global anisotropy) С = 0 - all materials are at most globally transversely С isotropic (D not used) С = 1 - at least one material is locally transversely С isotropic (D used) С - longitudinal cte for material MN ALPA(MN) С ALPT(MN) - transverse cte for material MN С С С expected on exit: DDOT(6, 6) - derivative with respect to TIME of stiffness matrix С BEGIN USER EDITS ***** place code here С END USER EDITS *******

RETURN END

A description of the input and output required for each subroutine is given in the comment statements at the top of each subroutine.

- Note: MAC/GMC assumes the longitudinal direction for the material is oriented in the x₁- coordinate direction, see Fig. 6.
- Note: Example K illustrates how one could use each of these routines to define material properties as well as the associated elastic stiffness matrix.

3.8 Interface Modeling

Interfaces in composite materials play a major role in the determination of their mechanical and thermal properties. Consequently, it is important to have the ability to model interface behavior accurately. This is accomplished in **MAC/GMC** in one of two ways. The first is to define an actual interface region with its own constitutive behavior. In this way the influence of initial imperfections (flaws, voids, improper wetting, etc.) and induced interfacial damage (due to stress, environment, chemical reactions, etc.) may be incorporated into the micromechanical analysis of the overall behavior of the composite. The development of proper interfacial constitutive models is an active area of research, and **MAC/GMC**, through the use of its **USRMAT** routine, provides the researcher with a convenient tool for testing new and existing interfacial constitutive models.

The second approach to modeling the effect of imperfect (weak) bonding between two phases (e.g. a fiber and a matrix) is to assume that a jump in the displacement field at an interface may occur given certain conditions, while still maintaining continuity of the traction vector. In the spirit of Jones and Whittier [13] and Achenbach and Zhu [14] we have assumed the following flexible interface model.

$$\begin{bmatrix} u_n^{\ I} = R_n \cdot \sigma_n^{\ I} \\ (u_t^{\ I} = R_t \cdot \sigma_t^{\ I}) \end{bmatrix} \qquad if \qquad \begin{bmatrix} \sigma_n^{\ I} \ge \sigma_{DBn} \\ \sigma_t^{\ I} \ge \sigma_{DBt} \end{bmatrix}$$
(EQ 44)

$$\begin{bmatrix} (u_n^{\ l} = 0) \\ (u_t^{\ l} = 0) \end{bmatrix} \qquad if \qquad \begin{bmatrix} \sigma_n^{\ l} < \sigma_{DBn} \\ \sigma_t^{\ l} < \sigma_{DBt} \end{bmatrix}$$
(EQ 45)

where R_n , R_t , σ_{DBn} and σ_{DBt} are the interfacial normal and shear, compliance and debond stresses, respectively. Note, the implementation of the various forms for R_n and R_t described below will impact the definition of the concentration matrices, $A^{(\alpha\beta\gamma)}$ in the original **GMC** formulation or its counterpart in the reformulated version, see section 3.1.

This approach to debonding has been implemented in **MAC/GMC** in two forms. In the first, R_n and R_t are assumed to be constants that are independent of time. Therefore, when the time derivative of EQ. 44 is taken (as it is for implementation in the incremental formulation of **MAC/GMC**) the expression becomes,

$$\begin{bmatrix} (\dot{u}_n^{\ I} = R_n \cdot \dot{\sigma}_n^{\ I}) \\ (\dot{u}_t^{\ I} = R_t \cdot \dot{\sigma}_t^{\ I}) \end{bmatrix} \qquad if \qquad \begin{bmatrix} \sigma_n^{\ I} \ge \sigma_{DBn} \\ \sigma_t^{\ I} \ge \sigma_{DBt} \end{bmatrix}$$
(EQ 46)

Hence, the debonding is instantaneous, and immediately reaches its full extent. If R_n and R_t are chosen to be sufficiently large (as is customary), the stress at the interface will remain constant, with a value of σ_{DB} after debonding occurs.

In the second form, R_n and R_t are assumed to be functions of time. Thus we obtain

$$\begin{bmatrix} \dot{u}_n^{\ l} = R_n \cdot \dot{\sigma}_n^{\ l} + \dot{R}_n \cdot \sigma_n^{\ l} \\ \dot{u}_t^{\ l} = R_t \cdot \dot{\sigma}_t^{\ l} + \dot{R}_t \cdot \sigma_t^{\ l} \end{bmatrix} \qquad if \qquad \begin{bmatrix} \sigma_n^{\ l} \ge \sigma_{DBn} \\ \sigma_t^{\ l} \ge \sigma_{DBt} \end{bmatrix}$$
(EQ 47)

in lieu of EQ 46. The additional term present in EQ. 47 for both normal and tangential debonding is significant. If the time-dependence of R_n and R_t is chosen wisely, these additional terms will enable local unloading in the composite. For implementation in **MAC/GMC**, the following functional form of the time-dependence has been employed,

$$R(t) = \Lambda \left[\exp\left(\frac{\hat{t}}{B}\right) - 1 \right]$$
 (EQ 48)

where \hat{t} is the time since debonding, and Λ and B are functional parameters which characterize how the interface unloads.

Figure 5 shows a simple example to illustrate the differences between the two implementations of this debond model. The repeating unit cell used to generate the results shown in this figure, is IDP= 1 as illustrated in Fig. 9. As Fig. 5 shows, the results generated using both implementations of the debond model are the same until debonding occurs at an interfacial stress of approximately 15 ksi. At this point, the stress at the interface in the first implementation becomes constant, while, in the case of the second implementation, the slope of the interfacial stress verses applied strain curve decreases and then the interface begins to unload. The effect of the difference between the two implementations on the predicted composite stress-strain response is shown clearly in Fig. 5, where the softer composite response is a result of additional inelasticity in the remaining matrix subcells due to the local stress redistribution from the debonded subcells.

,



Figure 5. Simulated transverse behavior using the two implementations of the debond model in **MAC/GMC**.

3.9 References

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4.0 Running MAC/GMC

MAC/GMC can be executed in one of two ways; 1) as a stand-alone code, and 2) through the UMAT facilities of the nonlinear finite element code, **ABAQUS**. Upon linking the associated object modules that comprise **MAC/GMC**, an executable version of **MAC/GMC** is generated as well as the file "feamac.o". The file "feamac.o" is one of the required files to be provided to **ABAQUS** for finite element access to **MAC**.

4.0.1 MAC/GMC - stand-alone:

The execution of **MAC/GMC** as a stand-alone code utilizes I/O redirection. Execution of **MAC/GMC** is initiated by typing a line at the command prompt such as;

MAC < infilename > outfilename

4.0.2 FEAMAC - Accessing MAC through ABAQUS:

To execute **MAC/GMC** through **ABAQUS** several files are required, these are summarized in the next section. The execution of **ABAQUS** for **MAC/GMC** access is as follows;

abaqus job=abaqus_input_deck user=feamac.o

Solution In the second sec

OR you can check our web page at

www.lerc.nasa.gov/WWW/LPB/mac/index.html

4.1 Input and Output Files

The input and output files required for the stand-alone version of **MAC/GMC** and **FEAMAC** are specified in different ways. The differences between these two methods of running **MAC/GMC** are detailed below.

4.1.1 MAC stand-alone

Since **MAC/GMC** reads from standard in and writes to standard out, the name of the files used as input and output are left to the user's discretion, i.e., will have no impact on the performance of the **MAC/GMC** code. As will be explained in the following sections, the user will also need to specify the names of the **PATRAN** output files and the files containing the X-Y data for plotting.

Note that a "debug" file may be generated depending on the "PRINT" level the user specifies in the input file (see **4.2.2**). This file will have the name "mac_debug".

<u>Input</u>	<u>Output</u>
infilename	outfilename
	debug file (mac_debug)
	PATRAN files (optional, see section 4.2.9)
	plot data files (see section 4.2.16 and 4.2.17)
	damage files (optional, see section 4.2.6)
	yield surface files (optional, see section 4.2.7)

4.1.2 FEAMAC

Almost all of the capabilities of MAC/GMC are available through the user defined material facilities of the ABAQUS finite element code. Access to MAC/ GMC from within ABAQUS is outlined here. Currently, MAC/GMC, can only be access using 3D elements; however, FEAMAC has only been specifically tested using the C3D8 element within ABAQUS. Note any options not available or modified for finite element implementation are denoted subsequently by the following notation, for example:

FEAMAC Note: Not required by FEAMAC

An example containing an **ABAQUS** input deck, required FORTRAN subroutines, and **FEAMAC** input files are given in **Example O**.

ABAQUS Input Deck:

To utilize FEAMAC, the ABAQUS input deck must contain the "*USER

SUBROUTINE", "*USER DEFINED FIELD", "*INITIAL CONDITIONS, TYPE=FIELD", "*EXPANSION" (if conducting nonisothermal analysis) and "*USER MATERIAL" card. Where the USER SUBROUTINE card identifies the name of the file containing the FORTRAN subroutines required to access MAC/ GMC and the "USER MATERIAL" card instructs ABAQUS to obtain the materials stress/strain behavior from the supplied subroutines. The name of the user material (defined in the ABAQUS input deck), converted to upper case, will be appended with the file extension that has been defined in the user supplied FORTRAN routines, consequently this name must be used as the MAC/GMC input filename for the given material. Note: the material name is limited by ABAQUS to be 8 characters or less. See the ABAQUS Users' Manual for a complete descriptions of these cards

ABAQUS FORTRAN Subroutine Files:

The name of this file is to be provided on the "*USER SUBROUTINE" card inside the **ABAQUS** input deck. These subroutines will be executed for every integration point of every element in the finite element model associated with a given material group. The required subroutine are listed in **Example O**. Only a portion of the FEAMAC_INIT routine should be edited by the user. This portion is bounded by the comments;

C *** BEGIN USER EDITS *** and C *** END OF USER EDITS ***

The seven variables within this section that can be edited are as follows:

PATH:	CHARACTER*80
	Path to working directory
EXTENSION:	CHARACTER*80
	File name extension to be appended to material name to form the name of the MAC/GMC input deck.
DFNAME:	CHARATER*80
	Name of diagnostic file.
NPEL:	INTEGER
	Number of integration points whose data is to be plotted.
N_PEN(N):	INTEGER ARRAY (N:1->NPEL)
	Element numbers whose data is to be plotted.
N_PIN(N):	INTEGER ARRAY (N:1->NPEL)
	Integration numbers whose data is to be plotted.

Note: N_PEN() and N_PIN() pairs define locations of data extraction to be utilized with the *MACRO or *MICRO MAC/GMC options

AB_PRINT:

INTEGER Diagnostic print level for FEAMAC.

- 0 NONE
- 1 UMAT: STRESS, STRAIN, INCREMENTAL AND EXECUTION TRACING INFORMATION
- 3 FEAMAC_PRE: STRESS, STRAIN, INCREMENTAL AND EXECUTION TRACING INFORMATION
- 4 TEMPERATURE INFORMATION
- 5 STATE INFORMATION BEFORE AND AFTER CALL TO FEAMAC
- 25 CONSTITUTIVE MATERIAL PROPERITES

Note: Higher values of AB_PRINT provide all information provided by lower values plus additional information as described above.

FEAMAC Input Deck:

The FEAMAC input deck can be identical to the stand-alone MAC/GMC input deck as discussed in section 4.2. The associated file name is obtained by converting the material property name, defined within the ABAQUS input deck, to uppercase and appending the file extension specified in the FEAMAC_INIT subroutine described previously. Since the load history definition and time integration are dictated by the associated ABAQUS, input deck, information in the MAC/GMC deck that relates to these issues will be ignored. All entries in section 4.2 that describe MAC/GMC options should be considered to be required by FEAMAC unless denoted otherwise.

Note: Multiple FEAMAC input decks can be specific per ABAQUS run, so that multiple material systems can be analyzed within a given structure. Utilization of this option will allow the most efficient execution of large problems. For example, if one knows that only a small portion of the problem will be inelastic then define the zone of elements that may go inelastic by a given material group and the remainder by another. In this way maximum speed can be achieved, as a purely isothermal, elastic, problem with no micro fields being printed executes the fastest.

FEAMAC Output Files:

Most diagnostic messages from **FEAMAC** appear in the **ABAQUS** msg file. Files containing X-Y plotting information are generated for each N_PEN/N_PIN pair. The names for these files correspond to those defined in the **MAC/GMC** deck except that the N_PEN and N_PIN number are append to the name.

Example:

If in the FEAMAC_INIT subroutine the user specified;

NPEL = 1 N_PEN(1) = 18 N_PIN(1) = 7

and in the FEAMAC deck one specified;

FEAMAC would provide the stress and strain data in the 11direction for integration point 7 of element number 18 in a file named

"PLOT_FILE_SVE.18.7.feamacro.data".

Note: see section 4.2.17 regarding curve data input

4.2 Input Requirements (for both MAC/GMC and FEAMAC)

This section describes how the data should appear in the input file, *infilename*, where each line in the input file is limited to 80 characters. In the following, each block of input data will have its own subsection and will typically contain the following information:

- 1) statement of purpose
- 2) declaration of input data block
- 3) example(s)
- 4) notes

A data block has the following general format:

1) ***KEYWORD** beginning of data block
 2) input line(s)
 3) % end of data block

where

1) ***KEYWORD**: (denotes beginning of data block)

Each input data block begins with the appropriate keyword, which starts with the * symbol. For example, the keyword for selecting the method of integration is *SOLVER. The input routine scans the input file and locates the appropriate keyword and then reads the corresponding input data

2) Input line(s):

The input lines contain the necessary input data. The specific format of these input lines will be given in the following subsections. The capital letters denote actual variable names and lower case letters represent the possible input choices. Multiple inputs can be on one line, just as shown in the following. However, at **least one space must be used** to separate **data sets** on a single line.

Two special characters (& and #) are provided for entering the input.

The "&" (continuation) symbol:

For input data that is too long to fit on a **single 80 character line**, the "&" symbol is used as the continuation character. Thus, a line of data may be divided into a series of lines. For example, when entering the material properties all of the data cannot fit on a single line, thus the continuation character is required:

EL=11700.,11700.,0.365,0.365,4287.5,1.,1. & VI=0.8E-8,0.1,0.1E-5,0.,0.85E-3,0.05,1.,1.,1.,3.3,1.8,1.35,1.,0.01

The "#" (comment) symbol:

The input file may also contain "comments" for the users convenience. The "#" symbol is used to mark a comment line, with the requirement that the "#" symbol appears in the <u>first</u> column.

3) % (denotes end of data block)

Each input block ends with the "%" symbol. The % symbol <u>should be</u> included as it signals the input routine that a particular data block has been completed.

A sample data block is as follows,

*****SOLVER

NTF=2 ISTM=0.0001 ERR=0.1E-3
NTF=2 ISTM=0.0000024 ERR=0.1E-2 %

Note how the # symbol is used to comment out a line of input data. Thus, the user can change various parameters by simply commenting out input lines containing different input data. Also, note how the data block is terminated with the % symbol and how <u>it need not</u> appear on a separate line.

Note: It is suggested that the order of keywords in a given input file follow that given in this manual as some keyword ordering may give rise to problems, since in certain cases, MAC/GMC uses input from previous keywords to determine which input to read from later keywords.

4.2.1 Header Line:

Purpose: Define the title of this particular job (80A format).

Note: the Header Line is limited to <u>one</u> 80 character line, and is always taken as the first line in the input file.

FEAMAC Note: Required by FEAMAC

problem title

Example: Transverse tensile response of 35% SCS6/Ti-6-4

4.2.2 Output Print Level:

Purpose: To control the output generated.

FEAMAC Note: Required by FEAMAC

***PRINT**

NPL=nplvl

%

Where:

nplvl:

-1	= print out macro stiffness matrix, engineering constants and macro-
	thermal expansion coefficients and stop
0	= Input echo, minimal output (most commonly used)
1	= time and iteration information
3	= Material properties/Stiffness matrix output
5	= Lamination information
7	= Damage information
9	= Ancillary data
10	=program execution trace and all array data

CAUTION: the choice 10 generates a very large output file

Note: Print levels are inclusive, such that the current level information as well as all lower level information is printed.

Example: (minimal print out)

*PRINT NPL=0 %

4.2.3 Load Type:

Purpose: To select load type

FEAMAC Note: Only LCON is required for FEAMAC, LOP, and LSS are Ignored by FEAMAC.

***LOAD**

LCON=nsel LOP=lop LSS=iopt

%

Where

nsel:

1 = Thermal Load

2 = Mechanical Load

3 = Thermomechanical Load

lop: (for 2-D and 3-D GMC)

- 1 = axial load in 1-direction
- 2 = axial load in 2-direction

3 = axial load in 3-direction

- 4 = shear load 23-direction 5 = shear load 13-direction
- 6 = shear load 12-direction
- 99 = generalized loading in potentially all 6 directions

For 2-D GMC:

7 = axial loads in 1 and 2 directions

8 = axial loads in 2 and 3 directions

9 = axial load in 1-direction and shear load in 23-direction

10 = axial load in 2-direction and shear load in 13-direction

For 3-D GMC:

- 7 = axial loads in 1 and 2 directions
- 8 = axial loads in 2 and 3 directions
- 9 = axial loads in 1 and 3 directions
- 10 = axial load in 1-direction and shear load in 23-direction
- 11 = axial load in 2-direction and shear load in 13-direction
- 12 = axial load in 3-direction and shear load in 12-direction

iopt.

- 1 = Strain control
- 2 = Stress control
- Note: If lop = 99 one must now enter six iopt values, for example: LCON= nsel LOP=lop LSS = *iopt1*, *iopt2*, *iopt3*, *iopt4*, *iopt5*, *iopt6*

Note: see Fig. 6 for the definitions of the directions mentioned above

Note: If using laminate option; lop = 1, 2 or 6 are the only valid selections

Example: General loading option

***LOAD** LCON=1 LOP=99 LSS=2,1,2,2,2,2 %

Note: This would correspond to specifying mixed stress and strain control, i.e.,:

 $\sigma_{11}, \epsilon_{22}, \sigma_{33}, \tau_{23}, \tau_{31}, \tau_{12}$

If however, the loads in the ***MECH** section were set to 0, 0.01, 0,0,0,0 this would correspond to a pure strain control problem in the 22 direction.



Figure 6: Coordinate Systems

4.2.4 Mechanical Load Control:

Purpose: Select type of load control for mechanical load

Note: This block is only required if LCON = 2 or 3

FEAMAC Note: All the data on the MECH card is ignored by FEAMAC

***MECH** NPTW=*nptw* $TI=t_1, t_2....t_{nptw}$ $LO=I_1, I_2, ..., I_{nptw}$

%

Where:

nptw: - number of points on load curve $t_1, t_2, ...t_{nptw}$: time values $l_1, l_2, ...l_{nptw}$: load curve values

Example:

*MECH

NPTW=3 TI=0., 1., 2. LO=0., 20., 30. %



Note: For LOP=7,8,...etc (i.e., two load/displacement components) the following format is required (See Example J for an example):

*MECH

NPTW=2	TI=0., 1.5	LO=0., 0.015	component 1 curve
NPTW=2	TI=0., 1.5	LO=0., 0.01 %	component 2 curve

For LOP=99, six load/displacement components are required.

4.2.5 Temperature Control:

Purpose: Select control for temperature.

- I Note: This block is only required if LCON = 1 or 3
- IF FEAMAC Note: All the data on the THERM card is ignored by FEAMAC
 - ***THERM**

NPTT=nptt TI= $t_1, t_2, \dots, t_{nptw}$ TE= $te_1, te_2, \dots, te_{nptt}$

%

÷

Where:

nptt: - number of points on temperature curve $t_1, t_2, ...t_{nptw}$: time values $te_1, te_2, ...te_{nptt}$: temperature curve values

Example:

***THERM**

NPTT=3 TI=0., 1., 2. TE=100., 200., 150. %



Note: For the thermomechanical load $t_{nptw} \equiv t_{nptt}$, and both curves must have $t_1 \equiv 0$. But the number and time value of the data points in-between maybe different, see figure 7.



Figure 7: Load History Specification

4.2.6 Fatigue Damage Option:

Purpose: Activate the Fatigue Damage analysis option

- Note: Currently this option is only available for continuous reinforcements, modid=1 see section 4.2.8
- FEAMAC Note: This option is currently not available from within FEAMAC

*DAMAGE

NCY=ncycle D=deld DMAX=dmax FG=fg FL=fl

where:

ncycle - number of "blocks" of cycles to be run (see Fig. 5)

deld - damage increment

dmax - maximum amount of damage allowed $(0 \le dmax \le 1)$

fg:

- 0 no macro (global) failure check
- 1 macro failure check
- fl:
- 0 no micro failure check
- 1 micro failure check

Note: For FG=1, the following data is required on a new line;

T=ss IC=comp V=val

Where

SS:

- 1 stress
- 2 strain

comp:

- 1 component 11
- 2 component 22
- 3 component 33
- 4 component 23
- 5 component 13
- 6 component 12

val - value of failure stress or strain (depending on value of *ss*)

Note: For FL=1, see description of additional required data in section 4.2.12

Note: the actual time vs. load data for the "block" of cycles is specified using the previously defined ***MECH** input data (section **4.2.4**).



Figure 8

Note: When the fatigue damage analysis option is activated the following additional files will be generated:

dam1.data	intermediate damage calculations
dam2.data	print-out of cycles to failure at the end of each load block
dam3.data	print-out of current damage in each subcell at the end of each load block

Example:

***DAMAGE**

NCY=2 D=0.25 DMAX=0.95 FG=1 FL=0 T=1 IC=2 V=100.

Note: In the above example:

- 2 "blocks" of cycles have been specified via NCY
- the damage increment is 0.25 (25%) with a maximum allowable amount of damage set to 0.95 (95%)
- a <u>Global</u> failure criteria is used: stress component -22 with a failure value of 100.

4.2.7 Yield and Damage Surface Generation Analysis

Purpose: Initiate probing history so as to generate flow/damage surface plots

Note: Currently this option is only available for continuous reinforcements, modid=1 see section 4.2.8

FEAMAC Note: All the data on the SURF card is <u>ignored</u> by FEAMAC; option is meaningless in the context of finite element.

*SURF

NPRE=npre ISP=isp IAN=ian C1=c1 C2=c2 C3=c3 C4=c4

where:

- *npre* the number of preloading steps before probing for the yield surface begins (e.g., this is allows one to represent stress-free cooling)
- *isp* stress space
 - = 1 transverse-axial $(\sigma_{22} \sigma_{11})$
 - = 2 transverse-transverse $(\sigma_{22} \sigma_{33})$
 - = 3 shear-axial $(\sigma_{12} \sigma_{11})$
- *ian* probe angle increment in degrees

"yield" criteria:

- c1 equivalent plastic strain, $\sqrt{\frac{2}{3}}\Delta\epsilon^{i}_{ij}\Delta\epsilon^{i}_{ij}$
- c2 Surface of Constant Dissipation Rate, SCDR, $\Sigma_{ii} \cdot \dot{\epsilon}_{ii}^{1}$
- c3 Surface of Constant Inelastic Strain Rate, SCISR, $\dot{\epsilon}_{ii}^i \cdot \dot{\epsilon}_{ii}^i$
- c4 Surface of Constant Inelastic Power, SCIP, $\overline{\sigma} \cdot \Delta \overline{\epsilon}^{i}$
- Note: Results from application of this option are described in: Lissenden C. J. and Arnold, S. M.; "Theoretical and Experimental Considerations in Representing Macroscale Flow/Damage Surfaces For Metal Matrix Composites", Int. Jnl. of Plasticity, Vol. 13, No. 4, pp. 327-358, 1997.
- Note: Upon fulfillment of each criteria (i.e., c1, c2, c3, and c4) the probe angle and stress vector are written to output files surf1.dat, surf2.dat, surf3.dat, and surf4.dat, respectively, for post processing by the user.
- Note: Probing continues until all four criteria are satisfied or <u>the specified load</u> <u>history is completed for a given probe angle</u>. Thus, to render a particular criterion inactive, use a value of 0 for the criterion so it is fulfilled immediately, thereby rendering the associated output meaningless. Also, it is suggested that a large load history be imposed to ensure yielding along a given probe angle occurs.

4.2.8 Micromechanics Model Identification:

Purpose: Select desired GMC micromechanics model

***MODEL**

MOD=modid

- Note: If modid=3 (laminate option) you must enter the following line of data, MATSYS=matsys NLY=nly THK=thk₁, thk₂,... thk_{nly} CON=c₁, c₂,...c_{nly} SYS=s₁,s₂,...s_{nly} ANG=a₁, a₂,... a_{nly}
- **FEAMAC Note:** Currently laminate option (modid=3) is not available within the finite element implementation.

%

Where

modid:	 defines micromechanics analysis 1 = double periodicity 2 = triple periodicity 3 = laminate option
matsys:	- number of different material systems in the analysis
nly:	- number of layers in the laminate
thk ₁ , thk ₂ , thk _{nly} :	- thickness of each layer
C ₁ , C ₂ , C _{nly} :	- layer material type id: 1 = isotropic 2 = anisotropic
s ₁ , s ₂ , s _{nly} :	- material system id number
a ₁ , a ₂ , a _{nly} :	- angles of each layer (in degrees)

Example 1: double periodicity model

*MODEL

MOD=1 %

Example 2: laminate option with 1 material system

***MODEL**

MOD=3 MATSYS=1 NLY=4 THK=0.25,0.25,0.25,0.25 CON=2,2,2,2 & SYS=1,1,1,1 ANG=45.,-45.,45. %

4.2.9 PATRAN Output: (Optional)

Purpose: To enable generation of additional output files for use within the MSC/ PATRAN graphical pre- and post-processing environment - when running the MACPOST program. For details on using MACPOST, see NASA TM 209062

Note: <u>This is an optional input line</u>. If output for use in the MACPOST program is not desired, do NOT include this data block. Inclusion of this data block will <u>increase execution time</u> significantly, particularly for complex RVE's, as microfield data for each subcell are written to additional output files.

***PATRAN**

FN=*prefix* TPRE=*tpre* STP= npstp

%

Where:

- *prefix*: filename prefix that will be assigned to the geometry file for use in **MACPOST**.
- tpre: preloading time after which output to data files begins
- *npstp*: time step interval for output to the MACPOST data files (output occurs every npstp time steps).
 CAUTION: a small value for npstp will cause large output files to be generated, which can result in significantly longer execution times.

Example 1: enable the generation of **MACPOST** output files using prefix "run1"

***PATRAN**

fn=run1 TPRE=57600 STP=300 %

Note: If this option is chosen, MAC/GMC generates 14 additional output files for use in the MACPOST program within MSC/PATRAN. After the time reaches 57600 units, data is written every 300 time steps. The following 14 output files are generated:

prefix.macgeo:	Contains unit cell (RVE) geometry data.
prefix.total_pat.data:	Contains number of time steps and number of subcells data

209062.

prefix.macro1_pat.data:	Macroscopic strain data at different times
prefix.macro2_pat.data:	Macroscopic stress data at different times
prefix.macro3_pat.data:	Macroscopic inelastic strain data at different times
prefix.macro4_pat.data:	Macroscopic thermal strain, creep time, temperature, and stress invariant data at dif- ferent times
prefix.micro1_pat.data:	Microscopic (subcell) strain data at different times
prefix.micro2_pat.data:	Microscopic (subcell) stress data at different times
prefix.micro3_pat.data:	Microscopic (subcell) inelastic strain data at different times
prefix.micro4_pat.data:	Microscopic (subcell) thermal strain, creep time, temperature, and stress invariant data at different times
prefix.micro1_pat.contour	Microscopic (subcell) strain data for contour plots
prefix.micro2_pat.contour	: Microscopic (subcell) stress data for contour plots
prefix.micro3_pat.contour	: Microscopic (subcell) inelastic strain data for contour plots
prefix.micro4_pat.contour	: Microscopic (subcell) thermal strain, creep time, temperature, and stress invariant data for contour plots
See Example S for an example i deck that utilizes the PATRAN	llustrating how to construct the required input I option and typical results.

Note: All 14 files listed above must be present in the active directory used in MSC/PATRAN in order to use the MACPOST program. For more details on the content of the various MACPOST files, see NASA TM

FEAMAC Note: Invoking this option within FEAMAC, will cause all "_pat" extensions in the above output files to be changed to "_feapat"

4.2.10 Integrator Identification

Purpose: Select type of integration scheme

re FEAMAC Note: All the data on the *SOLVER card is ignored by FEAMAC

***SOLVER**

NTF=ntf ... (additional data as specified below) %

- If NTF=1, enter the following: NTF=1 NPTS=*npts* TIM= $t_1, t_2, ..., t_{npts}$ STP= $st_1, st_2, ..., st_{npts-1}$ %
- If NTF=2, enter the following:

NTF=2 ISTM=istpm ISTT=istpt ERR=errtol %

Where:

ntf:

1 = Forward Euler method (specified time step)

- 2 = Predictor/Corrector method (self-adaptive time step)
- *npts:* = number of time points
- t_i: = start and end times for load increments
- st_{i:} = time step for that load increment
- *istpm*: = initial mechanical load time step
- *istpt*: = initial thermal load time step
- *errtol*: = error tolerance for predictor/corrector
- Note: <u>only</u> istpm is required for a mechanical load (*nsel=2*, see section 4.2.3), <u>only</u> istpt is required for a thermal load (*nsel=1*) and <u>both</u> istpm <u>and</u> istpt are required for a thermomechanical load (*nsel=3*)
- Note: errtol is <u>only</u> required when using predictor/corrector (ntf = 2) Suggested errtol = 0.1, if one sees oscillations in predicted response curve the errtol should be lowered.
- Image: Second and cases using the second debonding implementation <u>must</u> use forward euler.

Example 1: for a mechanical load, select predictor/corrector, with initial time step = 0.001 and error tolerance = 0.01

***SOLVER**

NTF=2 ISTM=0.001 ERR=0.01%

Example 2: select Forward Euler, time step = 0.01 and 0.02

***SOLVER**

NTF=1 NPTS=3 TIM=0.,1.,2. STP=0.01,0.02%



4.2.11 Thermal Conductivity Calculation: (Optional)

Purpose: Calculate the coefficients of thermal conductivity for the composite

*COND

- Note: If the *COND keyword is not used, the thermal conductivity calculations are not performed.
- Note: To use this option, all materials must be user input material properties (mat=U, see section 4.2.12) and must be at the same temperatures.
 MAC/GMC will then calculate the global thermal conductivities at the material property input temperatures, and write them to the output file, see Example T.

4.2.12 Constituent Material Model Identification:

Purpose: To select the model for the fiber and matrix constituents.

*FIBER NFIBS=nfibs

The following new line is to be repeated for each fiber (nfibs):

NF=nf_f **MS=ms_f** MF=ncmd_f NDPT=dpt **TEMP=mtemp** MAT=mat_f & IFM=ifm **D=d₁, d₂, d₃**

***MATRIX**

NMATX=*nmatx*

The following new line is to be repeated for each matrix (nmatx):

NM=nm_m **MS=ms**_m MM=ncmd_m NDPT=dpt **TEMP=mtemp** MAT=mat_m & IFM=ifm D=d₁,d₂,d₃

***MONOL (available for laminate option only)** NMON=*nmon*

The following new line is to be repeated for each monolithic material (nmon): NMO=nmo_{iso} MS=ms_{iso} MMO=ncmd_{iso} NDPT=dpt **TEMP=mtemp** & MAT=mat_{iso} **IFM=ifm**

%

Where:

- nfibs: number of different fibers
- nmatx: number of different matrices
- *nmon*: number of different monolithic layers
- *nf*; fiber material designation number running from 1 to *nfibs*, sequentially.
- *nm_m*: matrix material designation number running from 1 to *nmatx*, sequentially.
- *nmo_{iso}*: monolithic material designation number running from 1 to *nmon*, sequentially.
- **msf:** fiber material system ID (required only when using laminate option)
- *ms_m*: matrix material system ID. (**required only when using laminate option**)
- msiso: monolithic material system ID

- *ncmd:* material model identifier for either fiber, matrix or monolithic layer: 1 = Bodner-Partom Model
 - T = Douller-Partonn Moder
 - 2 = Modified Bodner-Partom Model
 - 3 = Robinson Viscoplastic Model
 - 4 = Generalized Viscoplastic Potential Structure (GVIPS) Model
 - 6 = Transversely Isotropic Elastic Model (2-3 isotropic plane)
 - 7 = Transversely Isotropic GVIPS Model (TGVIPS)
 - 9 = Local Transversely Isotropic Elastic Model
 - 99 = User defined model (see note at the end of section for special format instructions)
- *mat:* material identification letter for either fiber or matrix, selected from material database, see **Table II**.

■ Note: By specifying MAT=U allows the user to specify the material constants according to the formats specified below.

- *dpt*: flag indicating whether material constants should be temperature independent or temperature dependent
 - 1 = Temperature Independent
 - 2 = Temperature Dependent
- mtemp: the constant temperature at which material properties are to be taken (only required for dpt= 1 and when using database properties)
- *ifm* flag indicating whether material properties will be read from input file or taken from a user defined function (provided in the USRFUN subroutine) **ONLY NEEDED when MAT=U**.
 - 1 = read from input file
 - 2 = functional form taken from USRFUN routine
- d_i: direction vector defining the normal to the plane of local isotropy (only required for ncmd = 3, 7 or 9)
- Note: If modid=1 (Double periodicity) and one desires transverse isotropy using ncmd= 3,7 or 9 then the strong material direction must be specified in the 1direction.

Note: Additional Input is required if the ***DAMAGE** (section **4.2.6**) option is invoked

The following additional data is entered on a new line:

ANG= θ BN=b BP=b' OMU= ω_u OMFL= ω_{fl} OMM= ω_m ETU= η_u & ETFL= η_{fl} ETM= η_m BE= β A=a SFL= σ_{fl} XML=M SU= σ_u SK=sk %

where:

sk:

= 1 skip fatigue damage calculations for this material= 0 perform fatigue damage calculations

The remaining constants are described in section 3.5

- Note: In addition, for FL=1 (micro failure criteria) in section 4.2.6, the additional data is also required:
- T=ss IC=icomp V=val

SS:

1 - stress

2 - strain

comp:

- 1 component 11
- 2 component 22
- 3 component 33
- 4 component 23
- 5 component 13
- 6 component 12

val - value of failure stress or strain (depending on value of *ss*)

Example 1: select 1 fiber, SCS-6, and 1 matrix material, TIMETAL 21S, both read from database; i.e., SCS-6/TIMETAL 21S composite system

***FIBER**

NFIB=1 NF=1 MF=6 NDPT=1 TEMP = 23 MAT=D %

***MATRIX**

NMATX=1 NM=1 MM=4 NDPT=1 TEMP = 23 MAT=A %

r Note: See Section 3.3 for a mathematical description of each material model.

Example 2: (select 2 matrix materials; material 1: Boron, read from database, material 2: user supplied properties)

***MATRIX**

NMATX=2 NM=1 MM=6 NDPT=1 MAT=U IFM=1 EL=E_L, E_T, v_A , v_T , & G_A, α_A , α_T VI= D₀, Z₀, Z₁, m, n,q K= κ_A , κ_T NM=2 MM=1 NDPT=1 MAT=U IFM=1 EL=E_L, E_T, v_A , v_T , & G_A, α_A , α_T VI= D₀, Z₀, Z₁, m, n,q K= κ_A , κ_T %
Note: The κ_A , κ_T data is only required if the ***COND** keyword has been used; where κ_A , κ_T are the axial and transverse thermal conductivities, respectively.

Required Material constants for each model is as follows:

Bodner-Partom: ncmd = 1 Elastic: Inelastic: $EL=E_I$, E_T v_A , v_T , G_A , α_A , α_T $VI=D_0$, Z_0 , Z_1 , m, n,q Modified Bodner-Partom: ncmd = 2 Elastic: Inelastic: Dm_1 , Dm_2 Robinson Viscoplastic: *ncmd* = 3 Inelastic: Elastic: EL=E_L, E_T, v_A , v_T , G_A, α_A , α_T VI= n, m, μ , κ_T , β , R, H, \hat{G}_0 , η , ω **Directions**: $D=d_1, d_2, d_3$ GVIPS: ncmd = 4 Elastic: Inelastic: $VI = \mu$, κ , R_{α} , R_{κ} , B_0 , B_1 , L_0 , L_1 , m, n, EL=E, v, α p, q, w, z₀ Elastic Model: *ncmd* = 6 $EL=E_L, E_T, v_A, v_T, G_A, \alpha_A, \alpha_T$ TGVIPS: ncmd = 7 Inelastic: Elastic: $\mathsf{EL}=\mathsf{E}_{\mathsf{L}}, \mathsf{E}_{\mathsf{T}}, \mathsf{v}_{A}, \mathsf{v}_{T}, \mathsf{G}_{\mathsf{A}}, \alpha_{A}, \alpha_{T} \quad \mathsf{VI}=\kappa, \mathsf{n}, \mu, \mathsf{m}, \beta, \mathsf{R}, \mathsf{H}, \mathsf{G}_{0}, \omega, \eta$ Directions: $D=d_{1}, d_{2}, d_{3}$ Transversely Isotropic Elastic Model: ncmd = 9 Elastic Directions: $EL=E_L, E_T, v_A, v_T, G_A, \alpha_A, \alpha_T$ $D=d_1, d_2, d_3$

Model	Material	Temperature Dependent ?	Units	mat
Bodner-Partom	Aluminum (2024-T4)	Yes	Pa, sec, ^o C	A
<i>ncmd</i> = 1	Aluminum (2024-0)	Yes	Pa, sec, ^o C	В
all properties	Aluminum (6061-0a)	Yes	Pa, sec, ^o C	С
taken from ref [1]	Aluminum (6061-0b)	Yes	Pa, sec, ^o C	D
	Aluminum (pure)	Yes	Pa, sec, ^o C	Е
	Titanium (pure)	No	Pa, sec, ^o C	F
	Copper (pure)	No	Pa, sec, ^o C	G
Modified Bodner- Partom	TIMETAL 21S	Yes	Pa, sec, ^o C	A
<i>ncmd</i> = 2				
Robinson Visco-	Kanthal	No, 600°C	ksi, hr,ºC	A
plastic	FeCrAlY	Yes	ksi, hr, °C	в
<i>ncmd</i> = 3	W/Kanthal (vf=35%)	No, 600°C	ksi, hr, °C	с
GVIPS	TIMETAL 21S	Yes	ksi, sec, ^o C	Α
<i>ncmd</i> = 4				
Linear Elastic	Boron	No	Pa, ^o C	A
<i>ncmd</i> = 6	SCS-6	Yes	Pa, ⁰C	В
All properties are	Tungsten (W)	No	Pa, ^o C	С
assumed isotropic	Boron	No	ksi, ^o C	D
	SCS-6	Yes	ksi, ^o C	Е
	Tungsten (W)	No	ksi, ^o C	F
TGVIPS	Ti-6-4	Yes	ksi, sec, ^o C	Α
<i>ncmd</i> = 7				
Linear Elastic	T50 Graphite	No	Pa, ^o C	A
<i>ncmd</i> = 9	T300 Graphite	No	Pa, ^o C	В
Transversely Isotropic	P100 Graphite	Yes	Pa, ⁰C	С
	T50 Graphite	No	ksi, ^o C	D
	T300 Graphite	No	ksi, ^o C	E
	P100 Graphite	Yes	ksi, ^o C	F

Table II. MAC/GMC Material Constant Database

Note: Warning: It is the user's responsibility to ensure that consistent material property units are being employed within a given problem. Particularly, when mixing database and user supplied material properties.

- Note: Even if a material model is temperature independent, it can still be used in a nonisothermal analysis (ndpt=2). Its properties will just not vary with temperature.
- Note: Required Format for User Supplied <u>Non-Isothermal</u> Material Constants: each of the following data statements are on <u>separate</u> lines.

Note: The total number of viscoplastic constants (V1, V2, V3, ... VN) required for each model are described on the bottom of page 71. For ncmd=4, three extra lines (V15= κ_o V16= B_o' V17= β_κ) must be added. Also, D= d₁, d₂, d₃ is only required when ncmd= 3, 7 or 9.

■ Note: Format for <u>User defined material model</u> (ncmd=99):

Given User Supplied Isothermal Material Constants, the following special format is required:

*FIBER

NFIBS=*nfibs*

The following line is to be repeated for each fiber (nfibs): $NF=nf_f$ $MS=ms_f$ MF=99 NDPT=dpt NPE=npe $EL=e_1,e_2,...e_{npe}$ & $ALP=\alpha_A, \alpha_T$ NPV=npv $VI=v_1,v_2,...v_{npv}$ $K=\kappa_A, \kappa_T$ %

***MATRIX**

NMATX=nmatx

The following line is to be repeated for each matrix (nmatx): $NM=nm_m MS=ms_m MM=99 NDPT=dpt NPE=npe EL=e_1, e_2, \dots e_{npe} \& ALP= \alpha_A, \alpha_T NPV=npv VI=v_1, v_2, \dots v_{npv} K=\kappa_A, \kappa_T \%$

Where:

- *npe*: total number of elastic constants (maximum of 9)
- *npv*: total number of inelastic constants (model specific, max of 19)
- e₁, e₂,.. elastic constants
- v_1, v_2, \dots inelastic constants (model specific)
- α_A, α_T ... longitudinal and transverse thermal expansion coefficients
- κ_A, κ_T ... thermal conductivities (if ***COND** only)

OR given User Supplied Non-Isothermal Material Constants:

*FIBER NFIBS=nfibs

The following line is to be repeated for each fiber (nfibs): $NF=nf_f MS=ms_f MF=99 NDPT=2 MAT= U IFM= 1$ NPE=npe NPV=npv

with the following data statements immediately following each material declaration on a separate line.

$$V2= V2_{T_1}, V2_{T_2}, \dots V2_{T_{n-1}}$$

***MATRIX**

NMATX=*nmatx*

The following line is to be repeated for each matrix (nmatx): $NM=nm_m$ MS= ms_m MM=99 NDPT=2 MAT=U IFM = 1 NPE=npe NPV=npv

with the following data statements immediately following each material declaration on a <u>separate</u> line.

:

NTP=	ntpts
TEM=	T ₁ , T ₂ ,, T _{ntpts}
E1=	$E_{1T_1}, E_{1T_2}, \dots E_{1T_{nipis}}$
E2=	$E_{2T_1}, E_{2T_2}, \dots E_{2T_{nipls}}$

Enpe=
$$E_{npeT_1}, E_{npeT_2}, \dots E_{npeT_{nipis}}$$

ALPA= $\alpha_{AT_1}, \alpha_{AT_2}, \dots \alpha_{AT_{nipis}}$
ALPT= $\alpha_{TT_1}, \alpha_{TT_2}, \dots \alpha_{TT_{nipis}}$
V1= $V1_{T_1}, V1_{T_2}, \dots V1_{T_{nipis}}$
V2= $V2_{T_1}, V2_{T_2}, \dots V2_{T_{nipis}}$

: Vnpv= $Vnpv_{T_1}, Vnpv_{T_2}, \dots Vnpv_{T_{nipis}}$ KA= $\kappa_{AT_1}, \kappa_{AT_2}, \dots \kappa_{AT_{nipis}}$ KT= $\kappa_{TT_1}, \kappa_{TT_2}, \dots \kappa_{TT_{nipis}}$

- Note: Required material user subroutines; see Example K for details on how they integrate together
 - USRFUN: Routine that defines functional form for material properties Required when: ifm=2 and mat=U
 - USRFORMDE: Routine that forms the stiffness matrix. Required when ncmd = 99
 - USRCPEVAL: Routine that defines the time derivative of the stiffness matrix. Required when: a) **ifm=2 and mat=U** b) **ncmd=99 AND ifm=1 AND ndpt=2**

4.2.13 RVE Data:

Purpose: Select RVE representing desired fiber packing arrangement/architecture.

*MRVE

IDP=*idp*

For Double Periodic RVE (i.e., MOD=1):

Note: The following data is entered on the <u>same</u> line as IDP (except where noted)

Without Interface:

```
• For IDP = 0
```

no further data required (only used for a monolithic layer in a laminate)

```
• For IDP = 1, 2, or 3
  VF=vf
• For IDP = 4
  VF=vf
          XA=xa
• For IDP = 6
  VF=vf R=R
• For IDP = 7
  VF=vf R=R
• For IDP = 9
  VF1=vf1 RAD1=rad1 VF2=vf2
                                 RAD2=rad2
                                                R=R
• For IDP = 11
  VF=vf RAD=rad1 R=R
• For IDP = 13
  VF=vf R=R
With Interface:
•For IDP = 1, 2, or 3
  VF=vf RAD=rad1 CPER=cper
• For IDP = 4
  Currently NOT Available
• For IDP = 6
  VF=vf R= R CPER=cper
• For IDP = 7
  Currently NOT Available
```

For IDP = 9 VF1=vf1 RAD1=rad1 CPER1=cper1 VF2=vf2 & RAD2=rad2 CPER2=cper2 R=R
IDP = 11 VF=vf RAD=rad1 R=R CPER=cper
For IDP = 13 For IDP = 13 For IDP = 13

%

where:

vf, vf1, vf2 =	the fiber volume ratios
rad, rad1, rad2 =	fiber radii
cper, cper1, cper2 =	ratios of interface thickness to fiber radius
xa =	length of the cross, see Fig. 10.
R - VN which defined	, the ratio of distances between fiber centers w

R = X/Y which defines the ratio of distances between fiber centers within a "ply" and those between a "ply" (see Fig. 11)

Note: For the laminate option the RVE data has the following input format:

*MRVE
IDP=idp₁₁, idp₁₂, ... idp_{nly}
L=1 VF= ... (rve data for layer 1)
L=2 VF= ... (rve data for layer 2)
:
L=nly VF= ... (rve data for layer nly)

%

Where:

idp:- unique identifying number of each **Double Periodic** RVE as given below

idpDescription• 1 =Square Fiber, Square Pack (original 4-cell model)

$$V_f \le 1/(1+\Delta)^2$$

RVE shown in Fig. 9

idpDescription• 2 =Square Fiber, Triangular (hexagonal) Pack $V_f \le 0.86602/(1 + \Delta)^2$
RVE shown in Fig. 9• 3 =Square Fiber, Square Diagonal Pack;

 $V_f \leq 0.5/(1+\Delta)^2$

RVE shown in Fig. 9











Fiber



Figure 9: RVE's Available in MAC/GMC







IDP = 7



IDP = 9



IDP = 11



IDP = 13



Fiber

Figure 9 con't: RVE's Available in MAC/GMC

idp Description

$$V_f \le 1 - 4(xa)^2$$

RVE shown in Fig. 9

• 6 = 7x7 Circular Fiber Approximation Rectangular or Square Pack

$$V_f \le \frac{0.8125}{R(1+\Delta)^2}$$
 if $R > 1.0$
 $V_f \le \frac{R(0.8125)}{(1+\Delta)^2}$ if $R < 1.0$
RVE shown in Fig. 9

• 7= 14x14 Circular Fiber Approximation Rectangular or Square Pack

$$V_f \le \frac{0.8148}{R} \qquad if \qquad R > 1.0$$

 $V_f \le R(0.8148)$ if R < 1.0

RVE shown in Fig. 9

• 9 = Two Different Size Square Fibers, Rectangular or Square Pack

$$V_{f_{2}} \leq \frac{1}{R \left[(1 + \Delta_{2}) + (1 + \Delta_{1}) \frac{R_{f_{1}}}{R_{f_{2}}} \right]^{2}}$$

$$V_{f_{1}} \leq \frac{2\sqrt{RV_{f_{2}}}}{\left[\left(\frac{R_{f_{2}}}{R_{f_{1}}} \right)^{2} (1 + \Delta_{2}) + \left(\frac{R_{f_{2}}}{R_{f_{1}}} \right) (1 + \Delta_{1}) \right]}$$

$$\frac{V_{f_{1}}}{V_{f_{2}}} \left[\frac{R_{f_{1}}}{R_{f_{2}}} \right]^{2} \geq 1$$

RVE shown in Fig. 9

- Solution Note: Two fibers and two interfaces must be defined in *Fiber and *Interface.
- Solution Note: V_{f_1} will be altered, if necessary, such that an integer number of small fibers fit within the RVE.

• 11 = Square Fiber, Rectangular Pack

$$V_{f} \leq \frac{1}{R(1+\Delta)^{2}} \qquad if \qquad R > 1.0$$
$$V_{f} \leq \frac{R}{(1+\Delta)^{2}} \qquad if \qquad R < 1.0$$

RVE shown in Fig. 9

Description

26 x 26 Circular Fiber, Rectangular or Square Pack

 $V_f \le \frac{0.80613}{R}$ if R > 1.0

 $V_f \le 0.80613R$ if R < 1.0

RVE shown in Fig. 9

Note: $R_{f_1} = \text{rad1}, R_{f_2} = \text{rad2}$ and if no interface is present, $\Delta = cper$, $\Delta_1 = cper1$ and $\Delta_2 = cper2$ should be taken to be zero, in the above formulas.

99 = User Defined RVE
 Example of RVE representing random packing shown in Fig.
 12. Required input shown in Example I.

Example: Triangular packing with interface thickness 1% of fiber radius.

*MRVE

IDP=2 VF=35 RAD=0.07 CPER=0.01 %

• 13 =

<u>idp</u>







Figure 11: Hybrid Composite RVE; IDP = 9. Large Fiber Spacing Ratio, R=X/Y



IDP,= 99

Figure 12: User defined RVE (see Example I)

For Triple Periodic RVE (i.e., MOD=2):

Note: The following data is entered on the <u>same</u> line as the IDP data, except where noted.

Thus for

```
For IDP = 1
VF=vf ASP=asp
For IDP = 2
VF=vf ASP1=asp1 ASP2=asp2
For IDP = 3
VF=vf ASP1=asp1 ASP2=asp2 DR=dr
```

where,

- asp,asp1 are the aspect ratio of the short fiber (i.e., fiber length/fiber diameter; e.g. in IDP=3 this is $(d_1 + d_2 + d_3)/h_1$)
- asp2 is the aspect ratio of the unit cell, d/h (see Fig. 2)
- dr is the ratio d_1/d_3 which quantifies the fiber off-set (see Fig. 13)
- For IDP = 4

One of the following lines should be entered <u>after</u> the IDP data depending upon the option desired OPT=1 A=a B=b C=c D=d H=h L=l % OPT=2 VF=vf A=a B=b C=c RD=rd RL=rl % OPT=3 VF=vf RA=ra RC=rc D=d H=h L=l % OPT=4 VF=vf A=a B=b C=c RD=rd RLC= rlc %

where,

a,b, and c are the ellipsoid's semi-major axes (see Fig. 13)
d, h and I are the unit cell dimensions (see Fig. 13)
rd = aspect ratio of d/h
rl = aspect ratio of I/h
ra = aspect ratio of a/b
rc = aspect ratio of c/b
rlc = aspect ratio of I/(2c)
•For IDP = 10
VF=vf

Where

idp:- unique identifying number of each Triple Periodic RVE given Fig. 13

Description

Short fibers in Square array, with equal spacing in all directions (i.e., $d_2 = h_2 = l_2$ see Fig. 13)

 $0 \le V_f \le 1$

RVE shown in Fig. 13

Short fibers in Diagonal array, with variable inclusion spacing in the x_1 - direction

 $V_{f} < \frac{1}{2} \left(\frac{A s p 1}{A s p 2} \right) \qquad if \qquad A s p 1 < A s p 2$ $V_{f} < \frac{1}{2} \left(\frac{A s p 2}{A s p 1} \right)^{2} \qquad if \qquad A s p 1 > A s p 2$

RVE shown in Fig. 13

Off-set short fibers in square array, variable fiber spacing in the x_1 - direction --- NOTE Asp1 > Asp2

$$V_{f} < 4 \left(\frac{Asp1}{Asp2}\right)^{2} \qquad if \qquad \frac{Asp1}{Asp2} > 2$$
$$V_{f} < \frac{1}{2} \left(\frac{Asp2}{Asp1}\right) \qquad if \qquad \frac{Asp1}{Asp2} < 2$$
$$V_{f} > \frac{1}{2} \left(\frac{Asp2}{Asp1}\right)^{2}$$

RVE shown in Fig. 13

- 4= Ellipsoidal Inclusion
- Note: Analytical expressions for Vf limits are unavailable as forming this unit cell geometry requires solution of non-linear equations.
- Note: Use of non-physical ellipsoid unit cell dimensions (e.g., 2b > h) will cause execution of MAC/GMC to stop.
- 10= Open cell

 $0 \le V_f \le 1$

- Note: Solid material (see Fig. 13) is the matrix material specified in *MATRIX. whereas, the "open" material is the fiber material specified in *FIBER. To simulate a truly "open cell" material, specify a fiber material with the properties of air, thus fiber volume fraction becomes the void fraction
- 99= User defined RVE

• 3=

idp

1 =

• 2=

Note: The following is the format for the user defined RVE's, IDP = 99 (see Fig. 12 for an example).

1) For 2-D RVE: (Each line of data must be on a separate line)

NB=nb NG=ng H= h_1 , h_2 , h_{ng} L= l_1 , l_2 , l_{nb} CM= $ss_{nb,1}$, $ss_{nb,2}$,..., $ss_{nb,ng}$ repeat data for the (nb x ng) 2-D RVE CM= $ss_{1,1}$, $ss_{1,2}$,..., $ss_{1,ng}$

2) For <u>3-D</u> RVE: (Each line of data must be on a separate line)

NA=
$$na$$
 NB= nb NG= ng
D= $d_1, d_2, ..., d_{na}$
H= $h_1, h_2, ..., h_{ng}$
L= $l_1, l_2, ..., l_{nb}$
CM= $ss_{na,1,1}, ss_{na,2,1}, ..., ss_{na,nb,,1}$
CM= $ss_{na-1,1,1}, ss_{na-1,2,1}, ..., ss_{na-1,nb,,1}$
repeat data for the ($na \ge nb \ge ng$) 3-D RVE

Where:

 $ss_{i,j}$ and $ss_{i,j,k}$ - are the identifying material labels which are given in the following format: first character: F - for a fiber or M - for matrix second character: 1, 2, 3, ... for fiber/matrix number 1, 2, 3 ... - number of subcells in the alpha direction na - number of subcells in the beta direction nb - number of subcells in the gamma direction ng - the height of each subcell (x₂-dimension) h - the length of each subcell (x₃- dimension) 1 - the depth of each subcell (x1- dimension) d







IDP = 2



Figure 13 Available 3-D RVE's



Ellipsoidal Inclusion



Ellipsoidal Inclusion RVE (IDP = 4)

Figure 13 con't: Available 3-D RVE's





Figure 13 con't: Available 3-D RVE's

■ Note: A 2-D (continuous reinforcement) and 3-D (discontinuous reinforcement) example are described next with a detailed example of the user defined inputs being found in Example I, 2-D case, and in Examples H and N, for the 3-D case.





Example: 2-D user defined RVE (as shown above)

*MRVE

IDP=99 NB=3 NG=3 H=1.,1.,1. L=1.,1.,1. CM=M1,M1,M1 CM=M1,M1,F1 CM=F1,M1,M1 %

Note: the fiber ID is F1 and the matrix ID is M1



Figure 15

- **Example:** A 3-D User defined RVE (as shown above), representing a [0/90] continuous reinforced laminate composite.
- Note: The fiber is material number F1 and the matrix is material number M1.

.





*MRVE	
IDP=99 ASP=1.	
NA=4 NB=2 NG=2	
D=1.,1.,1.,1.	
H=1.,1.	
L=1.,1.	
CM=M1,M1	rows 1-4 are for cross-section 1 (γ =1)
CM=M1,M1	
CM=M1,M1	
CM=F1,M1	
CM=M1,M1	rows 5-8 are for cross-section 2 (γ =2)
CM=F1,F1	
CM=M1,M1	
CM=F1,M1 %	

4.2.14 Interface Data: (Optional, for IDP=1, 2, 3, 6 or 11)

Purpose: Specify interface layer properties

*INTERFACE

NINT=*nint*

The following line is to be repeated for each interface(nint): NI= nm_i MS= ms_i MI= $ncmd_i$ NDPT=dpt TEMP=mtemp MAT= mat_i IFM= ifm_i D= d_1 , d_2 , d_3

%

Solution If MAT= U, see section 4.2.12 for format of additional input required.

Where

nint:	- number of different interfaces
nm _i :	- i th interface number
ms _i : <i>ncmd_i: mat._i</i>	- matrix material system ID. (required only when using laminate option) - material model identifier for the <i>ith</i> interface - material id letter for the <i>ith</i> interface
dpt.	 flag indicating whether material constants should be temperature independent or temperature dependent 1 = Temperature Independent 2 = Temperature Dependent
mtemp:	 the constant temperature at which material properties are to be taken (only required for dpt= 1 and when using database properties)
ifm	 flag indicating whether material properties will be read from input file or taken from a user defined function (provided in the USRFUN subroutine) ONLY NEEDED when MAT=U. 1 = read from input file 2 = functional form taken from USRFUN routine
d _i :	 direction vector defining the normal to the plane of local isotropy (only required for ncmd = 3, 7 or 9)

Note: Please refer to section 4.2.12 for ncmd and mat values

Example: (Assuming one interface with user supplied material constants)

*INTERFACE NINT=1 NI=1 MI=4 NDPT=1 MAT=U IFM=1 & EL=11700.,0.365,1.0E-06 & VI=0.8E-8,0.1,0.1E-5,0.,0.85E-3,0.05,1.,1.,1.,3.3,1.8,1.35,1.,0.01 %

4.2.15 Debond Data: (Optional)

Purpose: Specify the subcell faces where debonding can take place (see section **3.8**)

2-D GMC Model Format

***DEBOND**

NII=*nii*

Option 1

```
DBCH= 1 NBI=nbi NGI=ngi FACE=nfc RN=rn BDN=bdstrn RS=rs & BDS=bdstrs TI=tmd
```

OR

```
# Option 2
```

```
DBCH= 2 NBI=nbi NGI=ngi FACE=nfc BDN=bdstrn GCN=gamman & BCN= betan TOLN=toln BDS= bdstrs GCS=gammas BCS=betas TI=tmd
```

:

repeat nii times

%

3-D GMC Model Format

*DEBOND

NII=nii

Option 1

```
DBCH= 1 NAI=nai NBI=nbi NGI=ngi FACE=nfc RN=rn BDN=bdstrn & RS=rs BDS=bdstrs TI=tmd
```

OR

Option 2

```
DBCH= 2 NAI=nai NBI=nbi NGI=ngi FACE=nfc BDN=bdstrn & GCN=gamman BCN= betan TOLN=toln BDS= bdstrs GCS=gammas & BCS=betas TI=tmd
```

%

Laminate Model Format

Note: Currently this is only available for continuous reinforcements, modid=1 see section 4.2.8

***DEBOND**

```
L=1 NII=nii (data for layer 1)
see 2-D GMC format above --- for required input here
L=2 Nii=nii (data for layer 2)
see 2-D GMC format above --- for required input here
:
L=nly Nii=nii (data for layer nly)
see 2-D GMC format above --- for required input here
%
```

Where:			
nii.	- number o	f subcell interfaces with de	bonding
<i>dbch</i> : - debond model implementation choice, =1 (for first) or (for second), see section 3.8			$e_{r} = 1$ (for first) or $= 2$
<i>nai, nbi</i> and <i>ngi</i> :	- subcell in	dices (α, β, γ)	
nfc:	 used specify which "face" of the subcell is to be deb- onded (See Fig. 17 and 18) 		
<u>Fo</u> = 1 = 2 = 3	<u>r 2-D GMC</u> Top Face Right Face Both Faces		
<u>For</u> = 1 = 2 = 3	<u>3-D GMC</u> Top Face Right Face Back Face	= 4 Top & Right Face = 5 Top & Back Face = 6 Right & Back Face	= 7 ALL Faces
rn:	- R _n value (see section 3.8)	

bdstrn:	 debond stress, normal component
rs:	- R _s value (see section 3.8)
bdstrs:	- debond stress, shear component
tmd:	 time at which debond criteria becomes active (defaults to <i>tmd</i>=0. if not specified)
gamman:	- Λ_n value (see section 3.8)
gammas:	- Λ_s value (see section 3.8)
betan:	- B _n value (see section 3.8)
betas:	- B _s value (see section 3.8)
toln:	 Normal stress reversal tolerance for unloading (i.e., stress value that is low enough to treat as zero stress interface has closed)
ræ Note:	For the Laminate Model Only: NII=0 specifies NO debonding for that layer and no additional debond data is required.

Note: All debond parameters must be calibrated for a given composite system. **Example M** provides some initial estimates for an SCS-6/Ti system.

Example 1:

allow debonding of subcell 1,1 top face normal component starting at time = 11

NII=1

DBCH= 1 NBI=1 NGI=1 FACE=1 RN=1. BDN=100. RS=0. BDS=0. TI= 11.

Example 2:

allow debonding of subcell 1,1 right face shear component starting at time = 15

NII=1 DBCH= 1 NBI=1 NGI=1 FACE=2 RN=0. BDN=0. RS=1. BDS=100. TI= 15.

Example 3: Three Layer Laminate Analysis

debond subcell 1,1 right face shear component for layer 2 only

L=1 NII=0 (no debonding specified) L=2 NII=1 DBCH=1 NBI=1 NGI=1 FACE=2 RS=1. BDN=100. L=3 NII=0 (no debonding specified)







Figure 17: Subcell Faces For 2-D GMC



Figure 18: Subcell Faces For 3-D GMC

4.2.16 Plot Point Information:

Purpose: Specify the frequency at which data will be written to output files for both X-Y data

*CURVE

NP=npmax

%

Where:

npmax: - plot point increment

Example: (print out every fifth data point)

*CURVE

NP=5 %

4.2.17 Curve Data:

Note: Currently a maximum of <u>5</u> macro and <u>5</u> micro curves may be specified per problem

Specifying Curve Data For Macro (composite) Quantities:

*MACRO NT=nucuv

Repeat the following line *nucuv* times: NC=*nocu* X=*maidx* Y=*maidy* NAM=*tname*

%

Where:

nucuv - total number of curves

nocu: - curve number

maidx and maidy variable options are:

$1 - e_{11}$	7 - σ ₁₁	13 - ε ₁₁	19 - 30 n.a.	36 - γ_{13}^{in}
2 - e ₂₂	8 - σ ₂₂	14 - ε ⁱ ₂₂	31 - ϵ_{11}^{th}	37 - Total Time
3 - <i>e</i> ₃₃	9 - σ ₃₃	15 - ϵ_{33}^{i}	32 - $\epsilon_{22}^{'h}$	38 - Creep Time
4 - γ ₂₃	10 - τ ₂₃	16 - γ ⁱ ₂₃	33 - $\epsilon_{33}^{\prime h}$	39 - Temperature
5 - γ ₁₃	11 - τ ₁₃	17 - γ ⁱ ₁₃	34 - γ_{23}^{th}	
6 - γ ₁₂	12 - τ ₁₂	18 - γ_{12}^{i}	35 - γ_{12}^{th}	

tname: - name of plot file

Note: The file(s) generated will be of the form *tname_macro.data*. If the user desires to use the same *tname* for files 2-5, a double quote, ", is entered for *tname*. Those files then will have the form:

*tname*2_macro.data *tname*3_macro.data etc.

. 1.

(see the example on the next page for more details)

FEAMAC Note: The file(s) generated when running **FEAMAC** will be of the form *tname*.element#.integrationpt#.feamacro.data, see section **4.1.2**.

EXAMPLE:

*MACRO

NT=3				
NC=1	X=1	Y=7	NAM=stress	file created: stress_macro.data
NC=2	X=37	Y=13	NAM ="	file created:stress2_macro.data
NC=3	X=38	Y=13	NAM= plot %	file created: plot_macro.data

Note: For the **Laminate option**, the *maidx* and *maidy* variable options are modified as follows:

1 - ē _{xx}	7 - \overline{N}_{XX}	15 - \overline{N}'_{XY}	21 - 23 n.a.	31 - \overline{N}_{XX}^T	37 - Total Time
2 - ē _{yy}	8 - \overline{N}_{YY}	16 - \overline{M}_{XX}^{I}	24 - $\bar{\kappa}_{xy}$	32 - \overline{N}_{YY}^T	38 - Creep Time
3 - ē ₃₃	9 - 11 n.a.	17 - \overline{M}_{YY}^{I}	25 - \overline{M}_{XX}	33 - \overline{N}_{XY}^T	39 - Temperature
4 - n.a.	12 - \overline{N}_{XY}	18 - \overline{M}_{XY}^{I}	26 - \overline{M}_{YY}	34 - \overline{M}_{XX}^T	
5 - n.a.	13 - \overline{N}_{XX}^{I}	19- κ _{xx}	27 - 29 n.a.	$35 - \overline{M}_{YY}^T$	
6 - γ _{xy}	14 - \overline{N}_{YY}^{I}	20 - κ _{yy}	30 - \overline{M}_{XY}	$36 - \overline{M}_{XY}^T$	

Note: When using laminate option, <u>resultant forces</u> (\overline{N}_{XX} , \overline{N}_{YY} , or \overline{N}_{XY}) are output in place of stress components. To obtain associated stresses one must merely divide resultant force by overall laminate thickness. Also, the strains are <u>mid-plane strains</u> while $\overline{\epsilon}_{33}$ represents the average strain through the thickness.

Specifying Curve Data for Micro (subcell) Quantities:

***MICRO**

NT=nucuv

Repeat the following line *nucuv* times: NC=nocu LYR=lyr CELL=nssel X=maidx Y=maidy NAM=tname2

%

Where:

nucuv - total number of curves

nocu:	- curve number
lyr.	 layer number (only required when using laminate option)
nssel:	 subcell number (see next page for details on numbering)
tname2:	- name of plot file

maidx and maidy variable options are:

- **Note:** The quantities Φ and Ψ are the possible internal state variables (constitutive model dependent).
- Note: For the Laminate option, the data will be output for both integration points within a layer. Thus 4 rather than 2 columns of data will appear in the associated output file(s), with the first 2 columns representing the x and y quantities at the lower quadrature point and the last 2 columns those associated with the upper quadrature point (see Example P).
- Note: The file(s) generated will be of the form *tname*2_micro.data. If the user desires to use the same *tname*2 for files 2-5, a double quote, ", is entered for *tname*2. Those files then will have the form:

*tname*22_micro.data *tname*23_micro.data etc.

- Note: The file(s) generated when running the laminate option will be of the form tname.l#.micro.data, where # indicates the layer number, see example below
- FEAMAC Note: The file(s) generated when running FEAMAC will be of the form tname.element#.integrationpt#.feamicro.data, again see section 4.1.2.

Note: Subcell numbering is assigned according to the following algorithm.

2-D case

Do * IB=1,NB Do * IG=1,NG *subcell number* = NG*(IB-1)+IG * continue

IN Note: Refer to numbers in upper left corners of subcells in Fig. 14 for example of 2-D subcell numbering scheme.

<u>3-D case</u>

Do * IA=1,NA Do * IB=1,NB Do * IG=1,NG *subcell number* = IA+(IB-1)*NA+(IG-1)*NA*NB * continue

■ Note: Refer to numbers in upper left corners of subcells in Fig. 16 for example of 3-D subcell numbering scheme.

EXAMPLE:

*MICRO

NT = 1 NC=1 CELL=1 X=1 Y=7 NAM=cell % f

file created:cell_micro.data

EXAMPLE: Assuming laminate comprised of 2 layers

***MICRO**

NT = 2 NC=1 LYR=1 CELL=2 X=3 Y=2 NAM=lam-x2 % file created:**lam-x2.l1.micro.data** NC=2 LYR=2 CELL=2 X=3 Y=1 NAM=lam-x3 %

file created:lam-x3.l2.micro.data

5.0 Conclusion/Future Modifications

A computationally efficient, user-friendly, comprehensive, micromechanics analysis tool, **MAC/GMC**, has been presented that admits physically based viscoplastic deformation and life models, can analyze continuous or discontinuous multiphased materials of interest in advanced propulsion systems, and can assist both the materials scientist and structural analyst in developing, designing and analyzing strategic materials. **MAC/GMC's** most outstanding feature is its ability to accurately model various laminated fiber architectures (including both shape and packing arrangements) at minimal expense both from a computational and required user input standpoint. However, the development of this tool is far from complete. A number of possible future enhancements that are planned or currently underway include:

- Provision of an implicit integration algorithm to improve computational efficiency.
- The incorporation of woven composite IDP architectures.
- The ability to analyze smart composites, i.e. piezoelectric, etc.
- The addition of debond unit cells, so as to minimize user input.
- The incorporation of additional damage evolution laws and failure criteria (e.g., ultimate longitudinal and transverse tensile strength) so as to automate life estimates.
- The incorporation of viscoelastic behavior models for the analysis of elevated temperature polymeric composites.
- The ability to use the triple-periodicity model in conjunction with laminate theory allowing individual lamina to be reinforced with particulates or weaves.

5.1 Acknowledgment

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6.0 EXAMPLE PROBLEMS

It is recommended that a new user construct a **MAC/GMC** input file using the data given in these Examples and then check to see if the same result plots and/or files are obtained.

6.1 Example A: Pure Mechanical Load

Sample Input File For A Mechanical Load Problem

The following example is used to explain the control blocks in more detail.

Problem Summary:

Load Type:	Mechanical
Load Component:	22-direction (transverse to fiber)
Load History:	Monotonic
Load Control:	Strain
Load History Data:	$\dot{\epsilon} = 8.333 \times 10^{-4}$ /sec,
	$\varepsilon_{max} = 0.015$,
	$\varepsilon_{min} = 0.$
	$\Delta t_{initial} = 0.0000024 \text{ sec}$
Micromechanics model:	Double Periodicity
Fiber Packing Arrangement:	Hexagonal Pack at 35% fiber volume ratio
Integration Algorithm:	Predictor/Corrector
Constituent Material Model:	GVIPS - isotropic form
Constituents:	Fiber: SCS-6 Matrix: TIMETAL21S Interface: fictitious weak interface for TIMETAL21S

Note: This example will take an extremely long time to run due to the elastic-perfectly viscoplastic definition of the fictitious weak interface, as the overall time step is limited to that of the allowable interface time step. This slow computational response is NOT indicative of GMC. Mechanical load, no residual, perfectly viscoplastic interface, takes long time *PRINT NPL=0 % *LOAD LCON=2 LOP=2 LSS=1 % *MECH NPTW=2 TI=0.,18. LO=0.,0.015 % *MODEL MOD=1 % *SOLVER NTF=2 ISTM=0.0000024 ERR=0.1E-2 % *FIBER NFIBS=1 NF=1 MF=6 NDPT=1 MAT=D TEMP=650 % *MATRIX NMATX=1 NM=1 MM=4 NDPT=1 MAT=A TEMP=650 % *MRVE IDP=2 VF=0.35 RAD=0.07 CPER=0.1 % *INTERFACE NINT=1 NI=1 MI=4 NDPT=1 MAT=U IFM=1 & EL=11700.,0.365,1. & VI=0.8E-8,0.1,0.1E-5,0.,0.85E-3,0.05,1.,1.,1.,3.3,1.8,1.35,1.,0.01 % *CURVE NP=1 % *MACRO NT=1 NC=1 X=2 Y=8 NAM=apdxa % *END


Note: Using this fictitious interface model the qualitative feature of a weak bond (reduction below that of the matrix only) is obtained, however an accurate fit of the experimental data is still lacking. For a more accurate prediction of transverse behavior see Examples C and D.

6.2 Example B: Pure Thermal Problem

Sample Input File For A Thermal Load Problem

The following example is used to explain the control blocks in more detail.

Problem Summary:

4

Load Type:	Thermal
Load History Data:	$\dot{T} = 0.01 ^{\circ}\text{C/sec},$
	$T_{max} = 371.1 {}^{\circ}\text{C}$
	$T_{min} = 21.1 {}^{\circ}\text{C}$
	$\Delta t_{initial} = 17.505 \text{ sec}$
Micromechanics model:	Double Periodicity
Fiber Packing Arrangement:	Square Pack at 35% fiber volume ratio
Integration Algorithm:	Predictor/Corrector
Constituent Material Model:	Bodner-Partom
Constituents:	Fiber: T50 Graphite Matrix: Aluminum (2024-T4)

Note: This problem is taken from reference 1., pg. .238

```
cooldown heatup bodner model aboudi prob. pre/cor gmc2d
*PRINT
NPL=0 %
*LOAD
LCON=1 %
*THERM
NPTT=3 TI=0.,35010.,70020. &
TE=371.1,21.1,371.1 %
*MODEL
 MOD=1 %
*SOLVER
NTF=2 ISTT=17.505 ERR=0.001 %
*FIBER
NFIBS=1
NF=1 MF=9 NDPT=2 MAT=A D=1.,0.,0.%
*MATRIX
NMATX=1
NM=1 MM=1 NDPT=2 MAT=A %
*MRVE
IDP=1 VF=0.30 %
*CURVE
NP=5 %
*MACRO
NT=1
NC=1 X=39 Y=1 NAM=apdxb %
*END
```



Note: It is recommended that a new user construct a mac input file using the data given in this Example and then check to see if the same result plot is obtained.

6.3 Example C: Thermomechanical Load Problem

Sample Input File For A Thermomechanical Load Problem

The following example is used to explain how to impose thermal residual stresses due to manufacturing into mechanical analysis.

Problem Summary:

Load Type:	Thermomechanical		
Load Component:	33-direction (transverse to fiber)		
Load History:	Cyclic		
Load Control:	Strain		
Load History Data:	Cool-down from 900 °C to 23 °C then hold tem- perature constant during mechanical loading		
	$\dot{T} = 1.52 \times 10^{-2}$ °C/sec		
	$\dot{\epsilon} = 1.666 x 10^{-4}$ /sec, $\epsilon_{max} = 0.015$,		
	$\varepsilon_{min} = 0.$		
	$\Delta t_{thermal} = 100. \text{ sec}, \Delta t_{mech} = 0.1 \text{ sec}$		
Micromechanics model:	Double Periodicity		
Fiber Packing Arrangement:	Rectangular Pack at 33% fiber volume ratio		
Integration Algorithm:	Forward Euler		
Constituent Material Model:	GVIPS - isotropic form		
Constituents:	Fiber: SCS-6 (properties input manually) Matrix: TIMETAL21S		
Debonding:	First implementation of debond model		

Note: This example demonstrates how one would included the effects of residual stresses on the composite behavior. Also, although strain control is prescribe, during a pure cool down MAC/GMC allows Poisson's strains to be incurred.

```
test of residual stress idp=11 23C apply transverse loading
*PRINT
NPL=0 %
*LOAD
LCON=3 LOP=3 LSS=1 %
# LCON=3 LOP=1 LSS=1 %
*MECH
NPTW=3 TI=0.,57600.,57690. LO=0.,0.,0.015 %
# NPTW=3 TI=0.,57600.,57750. LO=0.,0.,0.015 %
*THERM
 NPTT=3_TI=0.,57600.,57690. TE=900.,23.,23. %
# NPTT=3 TI=0.,57600.,57750. TE=900.,23.,23. %
*MODEL
 MOD=1 %
*SOLVER
 NTF=1 NPTS=3 TIM=0.,57600.,57690. STP=100.,0.1 %
# NTF=1 NPTS=3 TIM=0.,57600.,57750. STP=100.,0.1 %
*FIBER
 NFIBS=1
# Use GVIPS but make it elastic k=1E20, so it represents SCS-6 fiber
 NF=1 MF=4 NDPT=1 MAT=U IFM=1 &
 EL=58.E3, 0.32, 3.5E-06 &
 VI=0.8E-9,1.E20,0.1E-5,0.,0.85E-4,0.05,1.,1.,1.,3.3,1.8,1.35,1.,0.01 %
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=2 MAT=A %
*MRVE
 IDP=11 VF=0.33 RAD=0.07 R=1.1111 %
*DEBOND
 NII=2
 DBCH=1 NBI=1 NGI=1 FACE=2 RN=0.1 BDN=14. TI=57600.
 DBCH=1 NBI=1 NGI=2 FACE=2 RN=0.1 BDN=14. TI=57600. %
*CURVE
 NP=2 %
*MACRO
 NT=2
 NC=1 X=3 Y=9 NAM=apdxc-t
 NC=2 X=1 Y=7 NAM=apdxc-I
*END
```



- Note: In order to produce the longitudinal response curve one must change the direction of loading (LOP=1 instead of 3) and the mechanical strain rate imposed, so as to agree with the imposed experimental history by commenting and uncommenting the appropriate lines under *LOAD, *MECH, *THERM, and *SOLVER.
- Note: It is recommended that a new user construct a MAC/GMC input file using the data given in this Example and then check to see if the same resulting plot can be obtained.

6.4 Example D: Transverse Debonding

Broblem Summany	Input File for	Transverse Debonding	
Problem Summary			
Load Type:	Thermomechanical		
Load History Data:	33-direction (transverse to fiber)		
Load History:	Cyclic		
Load Control:	Strain		
Load History Data:	Cool-down from 900°C to 534°C to 23°C, heat-up to 650°C, then hold temperature during mechanical load- ing $\dot{T} = 0.0152$ °C/sec, 0.0871 °C/sec $\dot{\epsilon} = 1.666 \times 10^{-4}$ /sec, $\epsilon_{max} = 0.018$, $\epsilon_{min} = 0$. $\Delta t_{thermal} = 250.$, 40. sec.; $\Delta t_{mech} = 0.4$ sec.		
Micromechanics Model:	Double Periodicity		
Fiber Packing Arrangement:	Rectangular Pack, R = 1.1111, 35% fiber volume ratio		
Repeating Unit Cell:	26x26 circular fiber cross-section approximation		
Integration Algorithm:	Forward Euler		
Constituent Material Model:	Fiber: Matrix:	Elastic, transversely isotropic GVIPS - isotropic form	
Constituents:	Fiber: Matrix:	SCS-6 (temp. dep. properties input man- ually) TIMETAL21S	
Debonding:	Second implementation of debond model		
Debond Parameters:	$\sigma_{DB} = \sigma_{DB}^{o} + C_1 / slope $, $\sigma_{DB}^{o} = 7.ksi, C_1 = 15.ksi$		
	$\Lambda = \Lambda_o - C_2$ $C_2 = 0.0399$	$/ slope $, $\Lambda_o = 0.06 \frac{in^3}{kip}$, $p_7 \frac{in^3}{2}$ B= 16.5 1/sec	
	- 2	kip	

Note: The debond parameters σ_{DB}^{o} and Λ have been weighted with respect to the local slope (i.e., x_2/x_3) of the representative circular fiber at each interface (see IDP=13 Figure 9). This helps correct for the fact that the representative circle has interfaces normal to the loading direction (due to the necessarily rectangular-shaped subcells), while an actual circular fiber does not. For an actual circular fiber, the fiber matrix interface is only normal to the loading direction at two points, and as we trace the fiber's perimeter away from these points, the interface becomes increasingly parallel to the loading direction. In our representative fiber, we thus provide interfaces that are more parallel to the loading direction with larger debond strengths since the component of stress normal to the actual fiber interface (at this point) is lower. Similarly, we provide these more parallel interfaces with lower values of Λ so that, once debonding occurs, the stress at the simulated interface (which is in the global loading direction) unloads more slowly. The weighting has been chosen as linear (with respect to the local interfacial slope) for simplicity and

because this introduces only one additional parameter for each of σ_{DB}^{o} and Λ . Note that modeling of interfacial debonding is an area of active research using **MAC/GMC**, and thus the techniques employed in this example are still developing.

The results of this example shown in the following figure have been plotted only to an applied strain level of 0.016 because at this point the slope of the predicted global stress-strain curve becomes negative. We are treating this as a criterion for simulated global failure of the composite.

The interfacial curves shown in the figure are **NOT** simple plots of the ***MICRO** output files. Rather, the microlevel stress from these output files is plotted versus the macro level strain applied to the composite from the first ***MACRO** output file.

```
test of refined subcell transverse debonding
*PRINT
 NPL=1 %
*LOAD
 LCON=3 LOP=3 LSS=1 %
*MECH
 NPTW=5 TI=0.,24000.,57600.,64800.,64908. LO=0.,0.,0.,0.,0.018 %
*THERM
 NPTT=5 TI=0.,24000.,57600.,64800.,64908. TE=900.,534.583,23.,650.,650. %
*MODEL:
 MOD=1 %
*SOLVER
 NTF=1 NPTS=5 TIM=0..24000..57600..64800..64908. STP=250..40..40..0.4 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=2 MAT=U IFM=1
 NTP=6
 TEM=21.1,204.44,315.56,426.67,537.78,871.11
 EA=57.0E3.55.98E3.55.4E3.54.82E3.54.24E3.53.36E3
 ET=57.0E3,55.98E3,55.4E3,54.82E3,54.24E3.53.36E3
 NUA=0.25,0.25,0.25,0.25,0.25,0.25
 NUT=0.25,0.25,0.25,0.25,0.25,0.25
 GA=22.8E3,22.392E3,22.16E3,21.928E3,21.696E3,21.344E3
 ALPA=3.564E-6,3.618E-6,3.726E-6,3.906E-6,4.068E-6,4.572E-6
 ALPT=3.564E-6.3.618E-6.3.726E-6.3.906E-6.4.068E-6.4.572E-6
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=2 MAT=A %
*MRVE
 IDP=13 VF=0.35 R=1.111111111
*DEBOND
 NII=48
 DBCH=2 NBI=13 NGI= 1 FACE= 2 BDN=7. GCN=0.060 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
 DBCH=2 NBI=12 NGI= 2 FACE= 2 BDN=11.3 GCN=0.0486 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
 DBCH=2 NBI=11 NGI= 3 FACE= 2 BDN=12. GCN=0.0467 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
 DBCH=2 NBI=10 NGI= 4 FACE= 2 BDN=17. GCN=0.0334 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
 DBCH=2 NBI= 9 NGI= 5 FACE= 2 BDN=18.3 GCN=0.0300 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
 DBCH=2 NBI= 8 NGI= 6 FACE= 2 BDN=19. GCN=0.0280 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
 DBCH=2 NBI= 7 NGI= 7 FACE= 2 BDN=22, GCN=0.0200 BCN=16.5 TOLN=1 &
 BDS= 40 GCS=0.1 BCS=100 TI=64800
```

DBCH=2 NBI= 6 NGI= 8 FACE= 2 BDN=25.8 GCN=0.0100 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 5 NGI= 9 FACE= 2 BDN=27. GCN=0.00671 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 4 NGI=10 FACE= 2 BDN=29.5 GCN=0.000045 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 3 NGI=11 FACE= 2 BDN=52. GCN=0.0 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 2 NGI=12 FACE= 2 BDN=59.5 GCN=0.0 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 2 NGI=14 FACE= 2 BDN=59.5 GCN=0.0 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 3 NGI=15 FACE= 2 BDN=52. GCN=0.0 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 4 NGI=16 FACE= 2 BDN=29.5 GCN=0.000045 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 5 NGI=17 FACE= 2 BDN=27. GCN=0.00671 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 6 NGI=18 FACE= 2 BDN=25.8 GCN=0.0100 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 7 NGI=19 FACE= 2 BDN=22. GCN=0.0200 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 8 NGI=20 FACE= 2 BDN=19. GCN=0.0280 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI= 9 NGI=21 FACE= 2 BDN=18.3 GCN=0.0300 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=10 NGI=22 FACE= 2 BDN=17. GCN=0.0334 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=11 NGI=23 FACE= 2 BDN=12. GCN=0.0467 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=12 NGI=24 FACE= 2 BDN=11.3 GCN=0.0486 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=13 NGI=25 FACE= 2 BDN=7. GCN=0.060 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=14 NGI=25 FACE= 2 BDN=7. GCN=0.060 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=15 NGI=24 FACE= 2 BDN=11.3 GCN=0.0486 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=16 NGI=23 FACE= 2 BDN=12. GCN=0.0467 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=17 NGI=22 FACE= 2 BDN=17. GCN=0.0334 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=18 NGI=21 FACE= 2 BDN=18.3 GCN=0.0300 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800 DBCH=2 NBI=19 NGI=20 FACE= 2 BDN=19. GCN=0.0280 BCN=16.5 TOLN=1 & BDS= 40 GCS=0.1 BCS=100 TI=64800

```
DBCH=2 NBI=20 NGI=19 FACE= 2 BDN=22. GCN=0.0200 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=21 NGI=18 FACE= 2 BDN=25.8 GCN=0.0100 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=22 NGI=17 FACE= 2 BDN=27. GCN=0.00671 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=23 NGI=16 FACE= 2 BDN=29.5 GCN=0.000045 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=24 NGI=15 FACE= 2 BDN=52. GCN=0.0 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=25 NGI=14 FACE= 2 BDN=59.5 GCN=0.0 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=25 NGI=12 FACE= 2 BDN=59.5 GCN=0.0 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=24 NGI=11 FACE= 2 BDN=52. GCN=0.0 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=23 NGI=10 FACE= 2 BDN=29.5 GCN=0.000045 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=22 NGI= 9 FACE= 2 BDN=27. GCN=0.00671 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=21 NGI= 8 FACE= 2 BDN=25.8 GCN=0.0100 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=20 NGI= 7 FACE= 2 BDN=22. GCN=0.0200 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=19 NGI= 6 FACE= 2 BDN=19. GCN=0.0280 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=18 NGI= 5 FACE= 2 BDN=18.3 GCN=0.0300 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=17 NGI= 4 FACE= 2 BDN=17. GCN=0.0334 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=16 NGI= 3 FACE= 2 BDN=12. GCN=0.0467 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=15 NGI= 2 FACE= 2 BDN=11.3 GCN=0.0486 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
DBCH=2 NBI=14 NGI= 1 FACE= 2 BDN=7. GCN=0.060 BCN=16.5 TOLN=1 &
BDS= 40 GCS=0.1 BCS=100 TI=64800
*CURVE
NP=3 %
*MACRO
NT=2
NC=1 X=3 Y=9 NAM=TRAN-M
NC=2 X=24 Y=3 NAM=TRAN-T
*MICRO
NT=5
# 13.1
NC=1 CELL=313 X=3 Y=9 NAM=TRAN-1
```

2,14 NC=2 CELL=40 X=3 Y=9 NAM=TRAN-2 # 4,16 NC=3 CELL=94 X=3 Y=9 NAM=TRAN-3 # 7,19 NC=4 CELL=175 X=3 Y=9 NAM=TRAN-4 # 5,17 NC=5 CELL=121 X=3 Y=9 NAM=TRAN-5 % *END

The following figure was obtained from the x-y plot data file produced by the present example.



6.5 Example E: General Loading

Sample Input File for General Loading

Problem Summary

Load Type:	Thermo-M	echanical
Load History Data:	Combined 33 13 12 12 11, Hea	: • normal strain (transverse to fiber) • shear strain • shear strain 22, and 23 component stress-free at-up from 23 °C to 223 ° C
Load History:	Monotonic	
Load Control:	Combined	stress and strain
Load History Data:	$\dot{T} = 1.85$ $\dot{\epsilon}_{22} = 9.26x$	^o C/sec 10^{-5} /sec, $\varepsilon_{22max} = 0.01$, $\varepsilon_{22max} = 0$
	$\epsilon_{13} = 5.56x$	10^{-5} /sec, $\varepsilon_{13max} = 0.006$, $\varepsilon_{13min} = 0$
	$\dot{\varepsilon_{12}} = 1.11x$	10^{-4} /sec, $\varepsilon_{12max} = 0.012$, $\varepsilon_{12min} = 0$
	$\Delta t = 0.4$	Sec
Micromechanics Model:	Double Pe	riodicity
Fiber Packing Arrangement:	Square Pa	ack, R = 1., 45% fiber volume ratio
Repeating Unit Cell:	2x2 subce	lls, square pack using IDP=11
Integration Algorithm:	Forward E	uler
Constituent Material Model:	Fiber: Matrix:	Elastic, isotropic GVIPS - isotropic form
Constituents:	Fiber: Matrix:	SCS-6 (temp. dep. properties input manually) TIMETAL21S

```
example of general loading
*PRINT
 NPL=1 %
*LOAD
LCON=3 LOP=99 LSS=2,2,1,2,1,1 %
*MECH
 NPTW=2 TI=0.,108. LO=0.,0.
 NPTW=2 TI=0.,108. LO=0.,0.
 NPTW=2 TI=0.,108. LO=0.,0.01
 NPTW=2 TI=0.,108. LO=0.,0.
 NPTW=2 TI=0.,108. LO=0.,0.006
 NPTW=2 TI=0.,108. LO=0.,0.012
*THERM
 NPTT=2 TI=0.,108. TE=23.,223.
*MODEL
 MOD=1 %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,108. STP=0.4 %
*FIBER
 NFIBS=1
 NF=1 MF=9 NDPT=2 IFM= 1 MAT=U
 NTP=6
 TEM=21.1,204.44,315.56,426.67,537.78,871.11
 EA=57.0E3,55.98E3,55.4E3,54.82E3,54.24E3,53.36E3
 ET=57.0E3,55.98E3,55.4E3,54.82E3,54.24E3,53.36E3
 NUA=0.25,0.25,0.25,0.25,0.25,0.25
 NUT=0.25,0.25,0.25,0.25,0.25,0.25
 GA=22.8E3,22.392E3,22.16E3,21.928E3,21.696E3,21.344E3
 ALPA=3.564E-6.3.618E-6.3.726E-6.3.906E-6.4.068E-6.4.572E-6
 ALPT=3.564E-6,3.618E-6,3.726E-6,3.906E-6,4.068E-6,4.572E-6
 D=1..0..0.
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=2 MAT=A %
*MRVE
 IDP=11 VF=0.45 RAD=0.07 R=1.0
*CURVE
 NP=5 %
*MACRO
 NT=4
 NC=1 X=3 Y=9 NAM=GENLD-33
 NC=2 X=5 Y=11 NAM=GENLD-13
 NC=3 X=6 Y=12 NAM=GENLD-12
 NC=4 X=39 Y=1 NAM=GENLD-T %
*END
```





122

6.6 Example F: Unidirectional Laminate Problem

Sample Input File For A Laminate Problem

The following example is used to explain the control blocks in more detail.

Problem Summary:

Load Type:	Mechanic	al
Load History:	Monotonio	C
Load Control:	Strain	
Load History Data:	$\dot{\epsilon} = 0.01/$	sec,
	$\varepsilon_{max} = 0.$	01,
	∆t _{constant} =	= 0.00025 sec
Micromechanics model:	Laminate	Theory
Fiber Packing Arrangement:	Square P	ack at 46% fiber volume ratio
Integration Algorithm:	Forward E	Euler
Constituent Material Model:	Bodner-P	artom
Constituents:	Fiber: Matrix:	Boron Aluminum (6061-0a)

```
test of laminate strain control
*PRINT
 NPL=0 %
*LOAD
 LCON=2 LOP=1 LSS=1 %
*MECH
 NPTW=2 TI=0.,1. LO=0.,0.01 %
*MODEL
 MOD=3 MATSYS=1 NLY=1 THK=1. CON=2 SYS=1 ANG=45. %
# MOD=3 MATSYS=1 NLY=1 THK=1. CON=2 SYS=1 ANG=0. %
# MOD=3 MATSYS=1 NLY=1 THK=1. CON=2 SYS=1 ANG=90. %
# MOD=3 MATSYS=1 NLY=1 THK=1. CON=2 SYS=1 ANG=10. %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,1. STP=0.00025 %
*FIBER
 NFIBS=1
 NF=1 MS=1 MF=6 NDPT=1 TEMP= 21, MAT=A %
*MATRIX
 NMATX=1
 NM=1 MS=1 MM=1 NDPT=1 TEMP= 21. MAT=C %
*MRVE
IDP=1
 L=1 VF=0.46 %
*CURVE
 NP=10 %
*MACRO
NT=1
 NC=1 X=1 Y=7 NAM=apdxf %
*END
```

The following figures were obtained from the x-y plot data file produced by the present example. The validity of these predictions were assessed by comparing these results to those previously obtain from:



reference 1 Fig. 8-5, pg. 235



reference 1 Fig. 8-7, pg. 237

6.7 Example G: A Cross-Ply Laminate Problem

Sample Input File For A Laminate Problem

The following example is used to explain the control blocks in more detail.

Problem Summary:

Load Type:	Mechanical		
Load History:	Monotonic		
Load Control:	Stress		
Load History Data:	$\dot{\sigma} = 0.0$	1 ksi/sec,	
	$\sigma_{max} =$	45. ks i,	
	∆t _{constat}	_{nt} = 1.125 sec	
Micromechanics model:	[±45] _s L	aminate with Double Periodicity	
Fiber Packing Arrangement:	Square Pack at 45% fiber volume ratio		
Integration Algorithm:	Forward Euler		
Constituent Material Model:	Bodner-Partom		
Constituents:	Fiber: Matrix:	Boron (properties input manually) Aluminum (properties input manually)	

Note: This problem is taken from the reference: Analysis of Metal-Matrix Composite Structures-II. Laminate Analysis, Arenburg, R. T. and Reddy, J. N., Computers and Structures, Vol. 40, N.6, pp. 1369-1385, 1991.

```
test of [+-45]2s laminate stress control
*PRINT
 NPL=0 %
*LOAD
 LCON=2 LOP=1 LSS=2 %
*MECH
 NPTW=2 TI=0.,4500. LO=0.,45. %
*MODEL
 MOD=3 MATSYS=1 NLY=4 &
 THK=0.25.0.25.0.25.0.25 &
 CON= 2, 2, 2, 2 &
 SYS= 1, 1, 1, 1 &
 ANG=45.,-45.,-45.,45. %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,4500. STP=1.125 %
*FIBER
 NFIBS=1
 NF=1 MS=1 MF=6 NDPT=1 MAT=U IFM=1 &
 EL=58.E3,58.E3,0.20,0.20,24.17E3,6.3E-6,8.28E-6 %
*MATRIX
 NMATX=1
 NM=1 MS=1 MM=1 NDPT=1 MAT=U IFM=1 &
 EL=9.53E3,9.53E3,0.33,0.33,3.58E3,21.06E-6,21.06E-6 &
 VI=1.E4,49.,63.,300.,4.,1. %
*MRVE
IDP=1,1,1,1
L=1 VF=0.45 %
L=2 VF=0.45 %
L=3 VF=0.45 %
L=4 VF=0.45 %
*CURVE
 NP=10 %
*MACRO
 NT=1
 NC=1 X=1 Y=7 NAM=apdxg %
*END
```

The following figure was obtained from the x-y plot data file produced by the present example and for validation was compared to Arenburg and Reddy, Fig. 16, pg. 1382.



6.8 Example H: Triple Periodic GMC; [0/90]

		•
Problem Summary:		
Loading:	Mechanic	al, Strain control
Load History Data:	$\dot{\epsilon} = 0.01/$	/sec,
	$\varepsilon_{max} = 0.$	01,
	∆t _{constant}	= 0.00025 sec
Micromechanics model:	Triple Per	iodic GMC
Fiber Packing Arrangement:	Rectangu (input ma	llar Pack at 46% fiber volume ratio nually, see figure below)
Integration Algorithm:	Forward I	Euler
Constituent Material Model:	Bodner-P	artom
Constituents:	Fiber: Matrix:	Boron Aluminum (6061-0a)

Sample Input File For Triple Periodic GMC

This problem uses the 3-D **GMC** model to simulate a [0/90] laminate as shown below.



```
test of gmc3d model 0/90 laminate simulation
*PRINT
 NPL=10 %
*LOAD
 LCON=2 LOP=3 LSS=1 %
*MECH
 NPTW=2 TI=0.,1. LO=0.,0.01 %
*MODEL
 MOD=2 %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,1. STP=0.00025 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=1 TEMP= 21. MAT=A %
*MATRIX
 NMATX=1
 NM=1 MM=1 NDPT=1 TEMP= 21. MAT=C %
*MRVE
 IDP=99
 NA=4 NB=2 NG=2
 D=0.67823,0.32177,0.67823,0.32177
 H=0.67823,0.32177
 L=0.32177,0.67823
 CM=M1,M1
 CM=M1,M1
 CM=M1,M1
 CM=F1,M1
 CM=M1,M1
 CM=F1.F1
 CM=M1,M1
 CM=F1,M1 %
*CURVE
 NP=10 %
*MACRO
 NT=1
 NC=1 X=3 Y=9 NAM=apdxh %
*END
```



Note: It is recommended that a new user construct a MAC/GMC input file using the data given in this Example and then check to see if the same result plot is obtained.

6.9 Example I: A User Defined RVE

Sample Input File For A User Defined RVE

The following example is used to explain how to enter a user defined architecture.

Problem Summary:

. .

Load Type:	Mechanical		
Load History: Load Control:	Monotonic Strain		
Load History Data:	$\dot{\epsilon} = 0.1$ /sec,		
	$\varepsilon_{max} = 0.01$,		
	∆t _{constant} = 0.1 sec		
Micromechanics model:	Double periodic		
Fiber Packing Arrangement:	Random composite (see figure 13)		
Integration Algorithm:	Forward Euler		
Constituent Material Model:	Bodner-Partom		
Constituents:	Fiber: Boron Matrix: Aluminum (6061-0a)		

```
random composite rve using 2-d gmc
*PRINT
NPL=-1 %
*LOAD
LCON=2 LOP=2 LSS=1 %
*MECH
NPTW=2 TI=0.,0.1 LO=0.,0.01 %
*MODEL
MOD=1 %
*SOLVER
NTF=1 NPTS=2 TIM=0.,0.1 STP=0.00005 %
*FIBER
NFIBS=1
NF=1 MF=6 NDPT=1 TEMP=21. MAT=A %
*MATRIX
NMATX=1
NM=1 MM=1 NDPT=1 TEMP=21. MAT=C %
*MRVE
IDP=99
NB=14 NG=14
CM=F1,M1,M1,M1,M1,F1,F1,M1,M1,F1,F1,M1,F1
CM=F1,M1,F1,F1,M1,M1,F1,F1,M1,M1,M1,M1,M1,F1
CM=M1,M1,F1,F1,M1,M1,M1,M1,M1,M1,F1,F1,M1,M1
CM=M1,M1,M1,M1,M1,M1,F1,F1,M1,F1,F1,M1,M1
CM=M1,M1,F1,F1,M1,M1,M1,F1,F1,M1,M1,M1,M1,M1,M1
CM=F1.M1.F1.F1.M1.M1.M1.M1.M1.M1.F1.F1.M1.F1
CM=F1,M1,M1,M1,M1,F1,F1,M1,M1,M1,F1,F1,M1,F1
CM=M1.M1.M1.M1.M1.F1.F1.M1.M1.M1.M1.M1.M1.M1.M1
CM=M1,F1,F1,M1,M1,M1,M1,M1,F1,F1,M1,F1,F1,M1
CM=M1,F1,F1,M1,F1,F1,M1,M1,F1,F1,M1,F1,F1,M1
CM=M1,M1,M1,M1,F1,F1,F1,F1,M1,M1,M1,M1,M1,M1,M1
CM=F1,F1,M1,M1,M1,M1,F1,F1,M1,M1,M1,F1,F1,M1
CM=F1,F1,M1,M1,M1,M1,M1,M1,M1,M1,M1,F1,F1,M1
CM=M1,M1,M1,M1,M1,M1,M1,M1,M1,M1,F1,F1,M1,M1
                                             %
*CURVE
NP=1 %
*MACRO
NT=1
NC=1 X=2 Y=8 NAM=apdxi %
*END
```

The corresponding output produced: VOLUME RATIO= 0.347E+00 MATERIAL NO.= 1 MATERIAL NO. = 2 VOLUME RATIO= 0.653E+00 CG - Effective/Macro Stiffness Matrix 0.221E+12 0.610E+11 0.608E+11 0.610E+11 0.148E+12 0.675E+11 0.608E+11 0.675E+11 0.147E+12 0.384E+11 **.** . 0.392E+11 0.398E+11 CI - Inverse of Effective/Macro Stiffness 0.537E-11 -0.152E-11 -0.153E-11 -0.152E-11 0.897E-11 -0.350E-11 -0.153E-11 -0.350E-11 0.906E-11 0.260E-10 0.255E-10 0.251E-10 Effective Engineering Moduli E11S= 0.186E+12 N12S= 0.282E+00 E22S= 0.111E+12 N23S= 0.390E+00 E33S= 0.110E+12 G23S= 0.384E+11 G13S= 0.392E+11 G12S= 0.398E+11 Effective Thermal Expansion Coefficients 0.120E-04 0.192E-04 0.196E-04 STOPPING: NPLVL == -1//__//

/____/_/_/__/

Fiber Packing Arrangement:

Constituent Material Model:

Integration Algorithm:

Constituents:

6.10 Example J: A Biaxial Load

Sample Input File For A Biaxial Load

Forward Euler

GVIPS

Fiber:

Matrix:

Square Pack at 35% fiber volume ratio

TIMETAL 21S

SCS-6 (GVIPS forced to be elastic)

The following example is used to explain how to impose a biaxial load state.

Problem Summary:	
Load Type:	Mechanical
Load History:	Monotonic
Load Control:	Strain
Load History Data:	$\dot{\epsilon_{11}} = 0.01$ /sec , $\dot{\epsilon}_{22} = 0.0067$ /sec
	$\epsilon_{11max} = 0.015$ $\epsilon_{22max} = 0.01$,
	$\Delta t_{constant} = 0.015 \text{ sec}$
Micromechanics model:	Double Periodic

135

```
test of biaxial load
*PRINT
NPL=7 %
*LOAD
 LCON=2 LOP=7 LSS=1 %
*MECH
 NPTW=2 TI=0.,1.5 LO=0.,0.015
 NPTW=2 TI=0.,1.5 LO=0.,0.010 %
*MODEL
 MOD=1 %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,1.5 STP=0.015 %
*FIBER
 NFIBS=1
 NF=1 MF=4 NDPT=1 MAT=U IFM=1&
 EL=58.E3,0.32,3.5E-06 &
 VI=0.8E-9,1.E20,0.1E-5,0.,0.85E-4,0.05,1.,1.,1.,3.3,1.8,1.35,1.,0.01 %
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=1 TEMP=650. MAT=A %
*MRVE
 IDP=1 VF=0.35 %
*CURVE
 NP=1 %
*MACRO
 NT=2
 NC=1 X=1 Y=7 NAM=biaxl
 NC=2 X=2 Y=8 NAM=biaxt %
*END
```



6.11 Example K: User Defined Inelastic Material Model

Problem Summary:			
Load Type:	Thermomechanical		
Load History:	Cyclic		
Load Control:	Strain		
Load History Data:	Cool-down from 400°C to 23°C, hold temperature during mechanical loading \dot{T} =0.419 °C/sec $\dot{\epsilon}$ =1.667x10 ⁻⁴ /sec, ϵ_{max} =0.015, ϵ_{min} =0. $\Delta t_{thermal}$ = 0.5 sec.; Δt_{mech} =0.05 sec.		
Micromechanics model:	Double Periodicity		
Fiber Packing Arrangement:	User Input 4x2 RVE with 2 fibers		
Integration Algorithm: Constituent Material Model:	Forward Euler Fiber 1: User Model (elastic) Fiber 2: User Model (elastic) Matrix: User Model (Bodner-Partom via USRMAT)		
Constituents:	Fiber 1:	Imaginary material, user function material properties. Elastic modulus is a function of longitudinal strain:	
		$E_{new} = 700-80.(\epsilon_{11})(E_{previous})$ with units of MPa	
	Fiber 2:	Imaginary material, temperature depen- dent properties input manually.	
	Matrix:	Imaginary material, user function tempera- ture dependent material properties. (see USRFUN subroutine)	

Sample Input File For A User Defined Material Model

Note: The purpose here is to demonstrate how to use the various user definable subroutines, i.e., **USRMAT, USRFUN**, **USRFORMDE**, **USRCPEVAL** see section **4.2.12** for more information.

```
test of user subroutines/input
*PRINT
NPL=10 %
*LOAD
LCON=3 LOP=1 LSS=1 %
*MECH
 NPTW=3 TI=0.,900.,990. LO=0.,0.,0.015 %
*THERM
 NPTT=3 TI=0.,900.,990. TE=400.,23.,23. %
*MODEL
 MOD=1 %
*SOLVER
 NTF=1 NPTS=3 TIM=0.,900.,990 STP=0.5,0.05 %
*FIBER
 NFIBS=2
 NF=1 MF=99 NDPT=2 MAT=U IFM=2
 NF=2 MF=99 NDPT=2 MAT=U IFM=1 NPE=2 NPV=0
 NTP=3
 TEM=21.,200.,400.
 E1=314.1E9,293.2E9,253.0E9
 E2=0.33,0.33,0.33
 ALPA=4.5E-6,5.3E-6,6.1E-6
 ALPT=4.5E-6,5.3E-6,6.1E-6
*MATRIX
 NMATX=1
 NM=1 MM=99 NDPT=2 MAT=U IFM=2 %
*MRVE
 IDP=99
 NB=4 NG=2
 H=1.,1.,1.,1.
 L=1.,1.
 CM=M1,M1
 CM=F2,M1
 CM=M1,M1
 CM=F1,M1
*CURVE
 NP=2 %
*MACRO
 NT=1
 NC=1 X=1 Y=7 NAM=apdx-k
```

*END



Figure was obtained from the x-y plot data file produced by the present example.

The USRMAT subroutine

The USRMAT subroutine is used here to implement the Bodner-Partom Viscoplastic model currently available in **MAC/GMC**

SUBROUTINE USRMAT (DSA, SA, PE, PV, D, LOCTISO, TIME, TSTEP, CTEMP, DTEMPR, NIO, NE, NV, NS, MN, CDUM, DMGF, æ NEP, NVP, NSASIZE) δc purpose: user material constitutive model for determination of С the inelastic strain and state variable rates С (used when ncmd = 99) С IMPLICIT DOUBLE PRECISION (A - H, O - Z) CHARACTER*2 CDUM DIMENSION SS(6), S(6), R(6)DIMENSION DSA(NSASIZE), SA(NSASIZE) DIMENSION PV(NVP), PE(NEP), D(3) c note: 1) in this subroutine, [SA] and [DSA] contain the micro (subcell) quantities for aboudi's micromechanics model С С 2) arrangement of [dsa] & [sa] arrays: С variable location С +-----С (1-6) (contains ENGINEERING shears) С strain rate |-----С (7-12) С | stress rate С | inelastic С strain rate (13-18) (contains ENGINEERING shears) С -----С | 12 "slots" (19 - 30)С С | for state variables |-----С thermal strain rate (31-36) С С ********* * * * * * * * * * * * С on entry: - vector of total (integrated) quantities (see above) SA С - vector of elastic constants for material MN PE(NE) С (where NE = # of elastic constants --> 9 MAX) С - vector of viscoplastic constants for material MN С PV(NV) (where NV = # of viscoplastic constants --> 19 MAX) С - vector of direction cosines (for models 3, 7, & 9) D(3) С - flag indicating if ANY material exhibits local LOCTISO С transverse isotropy (and global anisotropy) С = 0 - all materials are at most globally transversely С С isotropic (D not used)

```
= 1 - at least one material is locally transversely
С
                   isotropic (D used)
С
      TIME
              - current time
С
              - current time step
С
      TSTEP
      CTEMP
              - current temperature
С
      DTEMPR
              - time rate of change of temperature
С
              - unit number of .out file
      NIO
С
              - # of elastic constants --> 9 MAX
С
      NE
              - # of viscoplastic constants --> 19 MAX
      NV
С
              - subcell number
С
      NS
              - material number
    MN
С
              - material character/number designation
С
     CDUM
                (i.e. F1 = fiber #1)
С
             - damage factor - if damage is included the user
     DMGF
С
                should multiply material stiffness terms by DMGF
С
                when using such terms in his inelastic model.
С
С
c expected on exit:
     DSA - vector of rate quantities (see above)
С
             *****
****
    IF ( NE .GT. NEP ) THEN
       CALL FATALERROR (NIO)
       WRITE(NIO, *) 'TO MANY ELASTIC PROPERTIES FOR MATERIAL # ', MN
       WRITE(NIO, *) ' # USED
                           = `, NE
       WRITE(NIO, *) ' # ALLOCATED = ', NEP
       STOP
    ENDIF
    IF ( NV .GT. NVP ) THEN
       CALL FATALERROR(NIO)
       WRITE(NIO, *) 'TO MANY NON-LINEAR PROPERTIES FOR MATERIAL # ',
          MN
    &
                            = `, NV
       WRITE(NIO, *) ' # USED
       WRITE(NIO, *) ' # ALLOCATED = ', NVP
       STOP
    ENDIF
*****
                 BEGIN USER EDITS
*****
     WRITE(6, *) ' PV=', PV
С
     WRITE(6, *) ' PE=', PE
С
C-----
   MATERIALS #1 & #2 ARE ELASTIC
С
    (set inelastic strain rates to zero)
С
IF ((MN .EQ. 1) .OR. (MN .EQ. 2)) THEN
       DSA(13) = 0
       DSA(14) = 0
       DSA(15) = 0
```
```
DSA(16) = 0
     DSA(17) = 0
     DSA(18) = 0
     RETURN
   ENDIF
C-----
  MATERIAL #3 --> USE BODNER-PARTOM
С
C-----
C------
c copy appropriate viscoplastic material constants
C-----
   IF( NV .LT. 6 ) THEN
     CALL FATALERROR(NIO)
     WRITE(NIO, *) ' NOT ENOUGH PV SPACE: NV =', NV
     STOP
   ENDIF
   D0 = PV(1)
   ZO = PV(2)
   Z1 = PV(3)
   BM = PV(4)
   AN = PV(5)
   Q = PV(6)
C-----
c copy stress from [sa] to [ss]
C-----
   SS(1) = SA(7)
   SS(2) = SA(8)
   SS(3) = SA(9)
   SS(4) = SA(10)
   SS(5) = SA(11)
   SS(6) = SA(12)
C-----
c compute the deviatoric stress [s] in the subcell
TEMP = (SS(1) + SS(2) + SS(3)) / 3.0
   S(1) = SS(1) - TEMP
   S(2) = SS(2) - TEMP
   S(3) = SS(3) - TEMP
   S(4) = SS(4)
   S(5) = SS(5)
   S(6) = SS(6)
C------
C-----
   AJ2 = 0.5 * (S(1)**2 + S(2)**2 + S(3)**2) + S(4)**2 + S(5)**2 +
      S(6)**2
   &
   SQ3AJ = DSQRT(SS(1)**2 + SS(2)**2 + SS(3)**2 + 2 * (SS(4)**2 + 2)
      SS(5)**2 + SS(6)**2))
   &
   SQ2 = 1.414215
   IF (SQ3AJ .EQ. 0.0) THEN
     CALL ZEROR (R, 6)
   ELSE
```

```
R(1) = SS(1) / SQ3AJ
     R(2) = SS(2) / SQ3AJ
     R(3) = SS(3) / SQ3AJ
     R(4) = SQ2 * SS(4) / SQ3AJ
     R(5) = SQ2 * SS(5) / SQ3AJ
     R(6) = SQ2 * SS(6) / SQ3AJ
   ENDIF
C-----
 if d0=0 then assume elastic and zero-out
С
c [dsa(13-30)] (inelastic strain rate and
c internal variable rates), then return
C-----
   IF (DO .EQ. 0) THEN
     DO 100 JJ = 13, 30
       DSA(JJ) = 0.0
     CONTINUE
 100
     RETURN
C-----
c inelastic
C-----
   ELSE
      ZEF = ZO + Q * SA(2O) + (1 - Q) * (R(1) * SA(21) + R(2) *
         SA(22) + R(3) * SA(23) + R(4) * SA(24) + R(5)
   Se .
         * SA(25) + R(6) * SA(26))
   &
      IF (AJ2 .EQ. 0.0) THEN
       AL = 0.0
      ELSE
        ARG1 = ZEF^{*2.0} / (3.0 * AJ2)
        IF (ARG1 . GT. 1E6) ARG1 = 1E6
       CON = .5 * (AN + 1.0) / AN
        ARG = CON * (ARG1) * AN
        IF (ARG .GT. 50.0) ARG = 50.0
        AL = D0 / (DEXP(ARG) * DSQRT(AJ2))
     ENDIF
C-----
c inelastic strain rates
C-----
      DSA(13) = AL * S(1)
      DSA(14) = AL * S(2)
      DSA(15) = AL * S(3)
      DSA(16) = 2 * AL * S(4)
      DSA(17) = 2 * AL * S(5)
      DSA(18) = 2 * AL * S(6)
C-----
c plastic work rate
WPD = S(1) * DSA(13) + S(2) * DSA(14) + S(3) * DSA(15) + S(4) *
      DSA(16) + S(5) * DSA(17) + S(6) * DSA(18)
   &
C-----
c state variable rates
```

```
C-----
    DSA(19) = WPD
    ZOM = BM / ZO
     ZD = ZOM * (Z1 - ZEF) * WPD
     DSA(20) = ZD
    DSA(21) = ZD * R(1)
     DSA(22) = ZD * R(2)
     DSA(23) = ZD * R(3)
     DSA(24) = ZD * R(4)
     DSA(25) = ZD * R(5)
     DSA(26) = ZD * R(6)
   ENDIF
***********
              END USER EDITS
  RETURN
   END
```

The USRFUN subroutine

The USRFUN subroutine is used here to allow material properties to be entered in functional form instead of having to be linearly interpolated within **MAC/GMC**

c###	############ SUBROUTIN & DOLD & NE, I	*#####################################
с с с с	purpose:	user subroutine to allow elastic and viscoplastic material properties to be functions of TEMP or field variables. Used for user defined functional form material properties, that is, when: (mat .eq. 'U') .and. (ifm .eq. 2)
с с с	note:	can be used in conjunction with a provided material constitutive model, or a constitutive model input by the user in USRMAT
	IMPLICIT	DOUBLE PRECISION (A - H, O - Z)
	DIMENSION	DOLD(6, 6), D(3)
	DIMENSION	PEM(NEP, NMTS), PVM(NVP, NMTS)
	DIMENSION	ALPA (NMTS), ALPT (NMTS)
	DIMENSION	DSA(NSASIZE), SA(NSASIZE)

c note: 1) in this subroutine, [SA] and [DSA] contain the С micro (subcell) quantities for aboudi's micromechanics model С 2) arrangement of [dsa] & [sa] arrays: С variable location С +-----С (1-6) (contains ENGINEERING shears) С | strain rate С (7-12) stress rate С С | inelastic С (13-18) (contains ENGINEERING shears) С | strain rate С 12 "slots" (19-30) С for state variables С |-----С | thermal strain rate (31-36) с С ***** c NOTE: quantities in [SA] and [DSA] are SUBCELL quantities - the values on entry are for the first subcell containing material С # MN - the values on exit of this subroutine will be applied to С ALL SUBCELLS containing material # MN. It is thus recommended С that, if using the field variables, you assign the appropriate С material # to ONE SUBCELL ONLY. Use of [SA] and [DSA] in this С context in conjunction with bending in laminate theory will С result in erroneous results as field variables become dependent С on through-thickness position (while the material # does not). С С on entry: - material number С MN TIME - current time С TSTEP - current time step С С CTEMP - current temperature С DTEMPR - time rate of change of temperature - vector of total (integrated) quantities (see above) С SA DSA - vector of rate quantities (see above) С DOLD(6, 6) - previous elastic material stiffness matrix С PEM(NE, MN) - vector of previous elastic constants for material С С # MN (where NE = # of elastic constants --> 9 MAX) С PVM(NV, MN) - vector of previous viscoplastic constants for material # MN С (where NV = # of viscoplastic constants --> 19 MAX) С С С expected on exit: С PEM(NE, MN) - vector of current elastic constants for material MN PVM(NV, MN) - vector of current viscoplastic constants for С material MN С - vector of direction cosines D(3) С С (required for models 3, 7, & 9) С LOCTISO - flag indicating if ANY material exhibits local transverse isotropy (and global anisotropy) С = 0 - all materials are at most globally transversely С

```
isotropic (D not used)
С
            = 1 - at least one material is locally transversely
С
               isotropic (D used)
С
          - longitudinal cte for material MN
С
   ALPA(MN)
С
    ALPT(MN) - transverse cte for material MN
            - NUMBER OF ELASTIC PROPERTIES USED
       NE
С
            - NUMBER OF VISCOPLASTIC PROPERTIES USED
       NV
С
С
*****
              BEGIN USER EDITS
WRITE(6, *) 'TOP OF USRFUN'
С
C-----
C MATERIAL # 1: E = FUNCTION OF STRAIN & PREVIOUS E
          FOR USE WITH USER CONSTITUTIVE MODEL
С
C-----
   IF (MN .EQ. 1) THEN
     IF (SA(1) .GT. 0) THEN
       EAOLD = PEM(1, MN)
       EA = 700.E9 - EAOLD * SA(1) * 80.0
     ELSE
       EA = 700.E9
     ENDIF
     ET = EA
     FNA = 0.41
     FNT = 0.41
     GA = EA / (2.0 * (1.0 + FNA))
     ALPA(MN) = 4.5E-6
     ALPT(MN) = 4.5E-6
     NE = 5
     PEM(1, MN) = EA
     PEM(2, MN) = ET
     PEM(3, MN) = FNA
     PEM(4, MN) = FNT
     PEM(5, MN) = GA
c MATERIAL # 2: PROPERTIES = INTERPOLATED AT INPUT TEMPERATURES
            THIS SUBROUTINE IS NOT CALLED FOR MN #2 SINCE THE
С
            MATERIAL PROPERTIES ARE NOT FUNCTIONAL FORM
С
c MATERIAL # 3: PROPERTIES = LINEAR FUNCTION OF TEMPERATURE
           FOR USE WITH USER CONSTITUTIVE MODEL
С
C-----
```

```
ELSEIF (MN .EQ. 3) THEN
         IF (CTEMP .LT. 21.0) CTEMP = 21.0
         IF (CTEMP .GT. 400.) CTEMP = 400.
C -- ELASTIC
         E = 72.4E9 - 81.53E6 * (CTEMP - 21.)
         FN = 0.33 + 7.916E-5 * (CTEMP - 21.)
         ALP = 22.5E-6 + 3.958E-9 * (CTEMP - 21.)
         NE = 5
         PEM(1, MN) = E
         PEM(2, MN) = E
         PEM(3, MN) = FN
         PEM(4, MN) = FN
         PEM(5, MN) = E / (2. * (1. + FN))
         ALPA(MN) = ALP
         ALPT(MN) = ALP
C -- VISCOPLASTIC
         D0 = 1.E4
         Z0 = 340.E6
         Z1 = 435.E6
         BM = 300.0
         AN = 10.0 - 0.02493 * (CTEMP - 21.)
         Q = 1.0
         NV = 6
         PVM(1, MN) = D0
         PVM(2, MN) = Z0
         PVM(3, MN) = Z1
         PVM(4, MN) = BM
         PVM(5, MN) = AN
         PVM(6, MN) = Q
      ENDIF
```

RETURN END

The USRFORMDE subroutine

The USRFORMDE subroutine is used here to form the elastic material stiffness matrix in **MAC/GMC** when a user defined constitutive model is used.

```
SUBROUTINE USRFORMDE (MN, PEM, PVM, D, LOCTISO, DNEW,
        NE, NV, NEP, NVP, NMTS)
    &
    purpose: user subroutine to allow formation of material stiffness
С
            matrices based on a user constitutive model, which may use
С
             arbitrary material properties (used when ncmd = 99)
С
С
    IMPLICIT DOUBLE PRECISION (A - H, O - Z)
    DIMENSION DNEW(6, 6)
    DIMENSION PEM(NEP, NMTS), PVM(NVP, NMTS)
    DIMENSION D(3)
on entry:
С
               - material number
С
     MN
     PEM(NE, MN) - vector of elastic constants for material
С
                 # MN (where NE = # of elastic constants -> 9 MAX)
С
     PVM(NV, MN) - vector of viscoplastic constants for material # MN
С
С
                 (where NV = # of viscoplastic constants --> 19 MAX)
               - vector of direction cosines
     D(3)
С
               (required for models 3, 7, & 9)
С
С
     LOCTISO
               - flag indicating if ANY material exhibits local
С
                transverse isotropy (and global anisotropy)
               = 0 - all materials are at most globally transversely
С
С
                    isotropic (D not used)
               = 1 - at least one material is locally transversely
С
                    isotropic (D used)
С
С
   expected on exit:
С
     DNEW(6, 6) - current elastic material stiffness matrix
С
BEGIN USER EDITS
С
     WRITE(6, *) 'TOP OF USRFORMDE'
C NOTE: In the examples shown here, standard engineering material
С
       elastic constants (E, nu) are used. However, the user is
С
       free to employ any material elastic constants with his
С
       constitutive model (i.e., bulk modulus, etc.). Thus, the
С
       user must provide the equations to determine the stiffness
C
       components required by GMC from his elastic constants.
c MATERIAL # 1: E = FUNCTION OF STRAIN & PREVIOUS E
```

FOR USE WITH USER CONSTITUTIVE MODEL С c-----IF (MN .EQ. 1) THEN EA = PEM(1, MN)ET = PEM(2, MN)FNA = PEM(3, MN)FNT = PEM(4, MN)GA = PEM(5, MN)DO 100 I = 1, 6 DO 100 J = 1, 6 DNEW(I, J) = 0100 CONTINUE GT = 0.5 * ET / (1 + FNT)FK = 0.25 * EA / (0.5 * (1 - FNT) * (EA / ET) - FNA**2)DNEW(1, 1) = EA + 4.0 * FK * FNA**2DNEW(2, 1) = 2.0 * FK * FNADNEW(3, 1) = 2.0 * FK * FNADNEW(1, 2) = 2.0 * FK * FNADNEW(2, 2) = FK + GTDNEW(3, 2) = FK - GTDNEW(1, 3) = 2.0 * FK * FNADNEW(2, 3) = FK - GTDNEW(3, 3) = FK + GTDNEW(4, 4) = GTDNEW(5, 5) = GADNEW(6, 6) = GAC----c MATERIAL # 2: PROPERTIES = INTERPOLATED AT TEMPERATURES FOR USE WITH USER CONSTITUTIVE MODEL С c-----ELSEIF (MN : EQ. 2) THEN E = PEM(1, MN)FN = PEM(2, MN)GA = E / (2. * (1. + FN))DO 320 I = 1, 6DO 320 J = 1, 6DNEW(I, J) = 0320 CONTINUE GT = 0.5 * E / (1 + FN)FK = 0.25 * E / (0.5 * (1 - FN) - FN**2)DNEW(1, 1) = E + 4.0 * FK * FN**2 DNEW(2, 1) = 2.0 * FK * FN

.

```
DNEW(3, 1) = 2.0 \times FK \times FN
      DNEW(1, 2) = 2.0 * FK * FN
      DNEW(2, 2) = FK + GT
      DNEW(3, 2) = FK - GT
      DNEW(1, 3) = 2.0 * FK * FN
      DNEW(2, 3) = FK - GT
      DNEW(3, 3) = FK + GT
      DNEW(4, 4) = GT
      DNEW(5, 5) = GA
      DNEW(6, 6) = GA
c MATERIAL # 3: PROPERTIES = LINEAR FUNCTION OF TEMPERATURE
            FOR USE WITH USER CONSTITUTIVE MODEL
С
c-----
    ELSEIF (MN .EQ. 3) THEN
      E = PEM(1, MN)
      FN = PEM(3, MN)
      GA = PEM(5, MN)
      DO 300 I = 1, 6
         DO 300 J = 1, 6
           DNEW(I, J) = 0
300
      CONTINUE
      GT = 0.5 * E / (1 + FN)
      FK = 0.25 * E / (0.5 * (1 - FN) - FN**2)
      DNEW(1, 1) = E + 4.0 * FK * FN**2
      DNEW(2, 1) = 2.0 * FK * FN
      DNEW(3, 1) = 2.0 * FK * FN
      DNEW(1, 2) = 2.0 * FK * FN
      DNEW(2, 2) = FK + GT
       DNEW(3, 2) = FK - GT
       DNEW(1, 3) = 2.0 * FK * FN
      DNEW(2, 3) = FK - GT
       DNEW(3, 3) = FK + GT
       DNEW(4, 4) = GT
       DNEW(5, 5) = GA
       DNEW(6, 6) = GA
    ENDIF
    END USER EDITS
************
     RETURN
    END
```

The USRCPEVAL subroutine

The USRCPEVAL subroutine is used here to form the time derivative of the material stiffness matrix in **MAC/GMC**

C#####################################						
<pre>c purpose: user subroutine to allow formation of the TIME c derivative of the material stiffness matrix. c this subroutine is used when: c a) material properties are user defined and functional form. That is: (mat .eq. 'U') .and. (ifm .eq. 2) c b) the constitutive model is user-defined, and the material properties are not functional form, and the material properties are temperature-dependent c that is: (ncmd .eq. 99) .and. (ifm .ne. 2) .and. c (ndpt .eq. 2)</pre>						
IMPLICIT DOUBLE PRECISION (A - H, O - Z)						
DIMENSION DNEW(6, 6), DOLD(6, 6) DIMENSION DDOT(6, 6) DIMENSION PEM(NEP, NMTS), PVM(NVP, NMTS) DIMENSION ALPA(NMTS), ALPT(NMTS) DIMENSION DSA(NSASIZE), SA(NSASIZE)						
<pre>************************************</pre>						
c 2) arrangement of [dsa] & [sa] arrays: c variable location						
c + c strain rate (1-6) (contains ENGINEERING shears)						
c stress rate (7-12)						
c inelastic c strain rate (13-18) (contains ENGINEERING shears)						
c 12 "slots" (19-30) c for state variables						
c thermal strain rate (31-36)						
· · · · · · · · · · · · · · · · · · ·						
c on entry: c SA - vector of total (integrated) quantities (see above)						

```
- vector of rate quantities (see above)
     DSA
С
              - material number
С
     MN
              - current time
     TIME
С
     TSTEP
              - current time step
С
               - current temperature
     CTEMP
С
     DTEMPR
              - time rate of change of temperature
С
     DNEW(6, 6) - current elastic material stiffness matrix
С
     DOLD(6, 6) - previous elastic material stiffness matrix
С
     PEM(NE, MN) - vector of current elastic constants for material MN
С
                (where NE = # of elastic constants --> 9 MAX)
с
     PVM(NV, MN) - vector of current viscoplastic constants for
С
                material MN
С
                 (where NV = # of elastic constants --> 19 MAX)
С
               - vector of direction cosines
     D(3)
С
                (required for models 3, 7, & 9) local
С
               - flag indicating if ANY material exhibits
     LOCTISO
С
                transverse isotropy (and global anisotropy)
С
               = 0 - all materials are at most globally transversely
С
                    isotropic (D not used)
С
               = 1 - at least one material is locally transversely
С
                    isotropic (D used)
С
               - longitudinal cte for material MN
     ALPA(MN)
С
               - transverse cte for material MN
     ALPT(MN)
С
С
  expected on exit:
С
    DDOT(6, 6) - derivative with respect to TIME of stiffness matrix
С
BEGIN USER EDITS
***********
     DIMENSION DHOLD(6, 6, 3), DNEW1(6, 6), PEM1(9, 10), PVM1(19, 10)
     DIMENSION ALPA1(10), ALPT1(10)
     WRITE(6, *) 'TOP OF USRCPEVAL'
С
C-----
C MATERIAL # 1: E = FUNCTION OF STRAIN & PREVIOUS E
              FOR USE WITH USER CONSTITUTIVE MODEL
С
C-----
                                           _____
     IF (MN .EQ. 1) THEN
c --- method one - simply divide change in stiffness components by TSTEP
       DO 200 I = 1, 6
          DO 200 J = 1, 6
            DDOT(I, J) = (DNEW(I, J) - DOLD(I, J)) / TSTEP
  200
       CONTINUE
C-----
C MATERIAL # 2: PROPERTIES = INTERPOLATED AT TEMPERATURES
               FOR USE WITH USER CONSTITUTIVE MODEL
С
```

```
c-----
     ELSEIF (MN .EQ. 2) THEN
c --- method 2 - if material props are only function of temperature,
c --- calculate stiffness at CTEMP - 0.5 & CTEMP + 0.5, difference
c --- equals change per temp, multiply by DTEMPR equals change per time
        IF (CTEMP .LT. 21.0) CTEMP = 21.0
        IF (CTEMP .GT. 400.) CTEMP = 400. .
        DO 300 I = 1, 6
          DO 300 J = 1, 6
             DO 300 K = 1, 2
               DHOLD(I, J, K) = 0
 300
       CONTINUE
       DO 420 K = 1, 2
          IF (K .EQ. 1) CTEMP1 = CTEMP - 0.5
          IF (K .EQ. 2) CTEMP1 = CTEMP + 0.5
c --- this subroutine interpolates properties for material MN at
c --- the temperature CTEMP1
          CALL INTTMP(MN, CTEMP1, PEM1, ALPA1, ALPT1, PVM1)
c --- calculate the stiffness components
          CALL USRFORMDE (MN, PEM1, PVM, D, LOCTISO, DNEW1,
               NE, NV, NEP, NVP, NMTS)
    &
c --- note: for internal constitutive models (ncmd .ne. 99), use
c --- subroutine FORMDE to calculate stiffness components.
c --- syntax: CALL FORMDE(MN, PEM1, D)
          DO 480 I = 1, 6
             DO 480 J = 1, 6
                DHOLD (I, J, K) = DNEW1(I, J)
  480
          CONTINUE
  420
        CONTINUE
        DO 520 I = 1, 6
           DO 520 J = 1, 6
             DDOT(I, J) = (DHOLD(I, J, 2) - DHOLD(I, J, 1)) * DTEMPR
  520
        CONTINUE
C-----
c MATERIAL # 3: PROPERTIES = LINEAR FUNCTION OF TEMPERATURE
               FOR USE WITH USER CONSTITUTIVE MODEL
С
C-----
     ELSEIF (MN .EQ. 3) THEN
c --- method 2 - if material props are only function of temperature,
c --- calculate stiffness at CTEMP - 0.5 & CTEMP + 0.5, difference
c --- equals change per temp, multiply by DTEMPR equals change per time
        IF (CTEMP .LT. 21.0) CTEMP = 21.0
        IF (CTEMP .GT. 400.) CTEMP = 400.
```

```
DO 330 I = 1, 6
          DO 330 J = 1, 6
             DO 330 K = 1, 2
                DHOLD(I, J, K) = 0
330
        CONTINUE
        DO 400 \text{ K} = 1, 2
           IF (K .EQ. 1) CTEMP1 = CTEMP - 0.5
           IF (K .EQ. 2) CTEMP1 = CTEMP + 0.5
c --- determine material props at the appropriate temperature
           CALL USRFUN(MN, TIME, TSTEP, CTEMP1, DTEMPR, SA, DSA,
               DOLD, PEM1, PVM1, D, LOCTISO, ALPA1, ALPT1,
    &
               NE, NV, NMTS, NEP, NVP, NSASIZE)
    &
c --- calculate the stiffness components
           CALL USRFORMDE (MN, PEM1, PVM, D, LOCTISO, DNEW1,
               NE, NV, NEP, NVP, NMTS)
    &
c --- note: for internal constitutive models (ncmd .ne. 99), use
c --- subroutine FORMDE to calculate stiffness components.
c --- syntax: CALL FORMDE(MN, PEM1, D)
           DO 450 I = 1, 6
              DO 450 J = 1, 6
                DHOLD (I, J, K) = DNEW1(I, J)
  450
           CONTINUE
  400
        CONTINUE
        DO 500 I = 1, 6
           DO 500 J = 1, 6
              DDOT(I, J) = (DHOLD(I, J, 2) - DHOLD(I, J, 1)) * DTEMPR
        CONTINUE
  500
      ENDIF
```

RETURN END

6.12 Example L: Fatigue Damage Analysis

Sample Input File For A Fatigue Damage Analysis

The following example is used to explain how to conduct fatigue damage analysis within **MAC/GMC**.

Problem Summary:

Load Type:	Mechanical		
Load History:	Cyclic		
Load Control:	Stress		
Load Component:	11 - direction (in the fiber direction)		
Load History Data: $\Delta D = 0.15$	$\sigma_{11_{max}} =$	115. ksi, ἀ ₁₁ = 128. ksi/sec,	
	$\Delta t_{constant}$	= 0.01797 sec	
Micromechanics model:	Double Periodic		
Fiber Packing Arrangement:	Square Pack at 35% fiber volume ratio		
Integration Algorithm:	Forward Euler		
Constituent Material Model:	Elastic (fiber) and Bodner-Partom (matrix)		
Constituents:	Fiber: Matrix:	SCS-6 (input manually) Ti - 15 - 3 (at 800°F, input manually)	

Note: A similar example problem is described in more detail in: Wilt, T.E., Arnold, S. M., and Saleeb, A.F.: A Coupled/Uncoupled Computational Scheme for Deformation and Fatigue Damage Analysis of Unidirectional Metal Matrix Composites, <u>Applications of Continuum</u> <u>Damage Mechanics to Fatigue and Fracture</u>, ASTM STP 1315, D.L. McDowell, Ed., American Society for Testing and Materials, pp. 65-82, 1997.

The purpose here is to demonstrate how to use the ***DAMAGE** option.

```
Longitudinal fatigue analysis of square packed SCS6/Ti-15-3 system
*PRINT
 NPL=0%
*LOAD
 LCON=2 LOP=1 LSS=2 %
*MECH
NPTW=3 TI=0.,0.89844,1.79688 LO=0.,115.,0. %
*MODEL
 MOD=1 %
*DAMAGE
 NCY=12 D=0.15 DMAX=1.0 FG=0 FL=1%
*SOLVER
 NTF=1 NPTS=3 TIM=0.,0.89844,1.79688 STP=0.01797,0.01797 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=1 MAT=U IFM=1 &
 EL=58.E3,58.E3,0.25,0.25,23.2E3,3.5E-6,3.5E-6
 ANG=1. BN=2. BP=4. OMU=1. OMFL=1. OMM=1. ETU=1. &
 ETFL=1. ETM=1. BE=1. A=1. SFL=1. XML=1. &
 SU=1. SK=1
# Note sk=1 means skip fatigue damage for fiber, so damage properties are
# meaningless - but failure criteria is still applicable.
 T=1 IC=1 V=320. %
*MATRIX
 NMATX=1
 NM=1 MM=1 NDPT=1 MAT=U IFM=1 &
 EL=12770.,12770.,0.32,0.32,4837.12,21.06E-6,21.06E-6 &
 VI=1000.,120.,120.,10.,3.,1.
 ANG=0. BN=0. BP=0. OMU=1. OMFL=1. OMM=1. ETU=1. &
 ETFL=1. ETM=1. BE=2.27 A=0.2302 SFL=20.3 XML=900. &
 SU=128. SK=0
 T=2 IC=1 V=0.01 %
*MRVE
  IDP=1 VF=0.35 %
*CURVE
  NP=1 %
*MACRO
  NT=1
  NC=1 X=1 Y=7 NAM=apdxl %
```

```
*END
```

The following results is contained in the output file: dam2.data

Calculated Cycles To Failure At Increment 100 Applied Load Cycle 1 Subcell # Nf 0.100000E+10 1 0.613603E+05 2 0.613603E+05 3 0.664385E+05 4 ***** ****** 2 ***** ***** CONTROLLING SUBCELL 60494.2250830768 NF => 60494.0000000000 NF => **** CURRENT TOTAL # CYCLES = 60494.000000000 *** AFTER APPLIED LOAD CYCLE 1 ******* Calculated Cycles To Failure At Increment 200 Applied Load Cycle 2 Subcell # Nf 1 0.100000E+10 2 0.116736E+04 3 0.116736E+04 4 0.545641E+04 ****** ****** 2 ***** ***** CONTROLLING SUBCELL 674.890587838351 NF => 674.000000000000 NF => ****** ***** **** CURRENT TOTAL # CYCLES = 61168.000000000 *** AFTER APPLIED LOAD CYCLE 2 **** Calculated Cycles To Failure At Increment 300 Applied Load Cycle 3 Subcell # Nf 0.100000E+10 1 2 0.709630E+03 3 0.709630E+03 4 0.438086E+04 2 ***** ***** CONTROLLING SUBCELL NF => 419.320819753184 NF => 419.00000000000 **** **** CURRENT TOTAL # CYCLES = 61587.000000000

*** AFTER APPLIED LOAD CYCLE Calculated Cycles To Failure At Increment 400 Applied Load Cycle 4 Nf Subcell # 0.100000E+10 1 0.463247E+03 2 3 0.463247E+03 0.361975E+04 4 **** ****** 3 ***** ***** CONTROLLING SUBCELL 307.516313605484 NF => 307.000000000000 NF => **** CURRENT TOTAL # CYCLES = 61894.000000000 *** AFTER APPLIED LOAD CYCLE 4 **** Calculated Cycles To Failure At Increment 500 Applied Load Cycle 5 Subcell # Nf 0.100000E+10 1 2 0.295356E+03 0.295356E+03 3 0.301874E+04 4 ***** 2 ***** ***** CONTROLLING SUBCELL 232.831665300220 NF => 232.000000000000 NF => **** CURRENT TOTAL # CYCLES = 62126.000000000 5 *** AFTER APPLIED LOAD CYCLE ***** **** Calculated Cycles To Failure At Increment 600 Applied Load Cycle 6 Subcell # Nf 0.100000E+10 1 0.166907E+03 2 0.166907E+03 3 4 0.253239E+04 ***** 2 ***** ***** CONTROLLING SUBCELL NF => 158.292840626614 NF => 158.0000000000

********* **** CURRENT TOTAL # CYCLES = 62284.000000000 *** AFTER APPLIED LOAD CYCLE 6 ******* Calculated Cycles To Failure At Increment 700 Applied Load Cycle 7 Subcell # Nf 0.100000E+10 1 2 0.620698E+02 3 0.620698E+02 0.214841E+04 4 ********* **** 3 ***** ***** CONTROLLING SUBCELL 62.0697834887934 NF => NF => 62.00000000000000 **** CURRENT TOTAL # CYCLES = 62346.000000000 *** AFTER APPLIED LOAD CYCLE 7 Calculated Cycles To Failure At Increment 800 Applied Load Cycle 8 Subcell # Nf 1 0.100000E+10 2 0.735691E+01 0.735691E+01 3 4 0.195843E+04 2 ***** ***** CONTROLLING SUBCELL NF => 7.35691370876420 NF => 7.00000000000000 **** CURRENT TOTAL # CYCLES = 62353.000000000 *** AFTER APPLIED LOAD CYCLE 8 Calculated Cycles To Failure At Increment 900 Applied Load Cycle 9 Subcell # Nf 0.100000E+10 1 2 0.290197E+01 3 0.290197E+01 4 0.194075E+04 ********

```
2 *****
***** CONTROLLING SUBCELL
NF => 2.90196867892969
    2.000000000000000
NF =>
**** CURRENT TOTAL # CYCLES = 62355.000000000
*** AFTER APPLIED LOAD CYCLE
                         9
*****
Calculated Cycles To Failure At Increment *** Applied Load Cycle 10
        Nf
Subcell #
  1
     0.100000E+10
     0.202802E+01
  2
     0.202802E+01
  3
     0.193669E+04
  4
*****
*****
                       2 *****
***** CONTROLLING SUBCELL
NF => 2.02801986635817
     2.000000000000000
NF =>
**** CURRENT TOTAL # CYCLES = 62357.000000000
                        10
*** AFTER APPLIED LOAD CYCLE
Calculated Cycles To Failure At Increment *** Applied Load Cycle 11
Subcell #
         Nf
  1
      0.100000E+10
     -0.999000E+03
  2
     -0.999000E+03
  3
      0.193110E+04
  4
        2 FAILED ... SKIPPED
SUBCELL
                              2
 **** WARNING **** DELN < 1 FOR CELL
            3 FAILED ... SKIPPED
SUBCELL
 **** WARNING **** DELN < 1 FOR CELL
                              3
 **** WARNING **** DELN < 1 FOR CELL
                              4
 SORTN: **** STOP **** DELN < 1
```

At this point the matrix subcells have all failed and the analysis stops.

Thus the life of the composite is 62357 cycles.

```
The following results is contained in the output file: dam3.data
```

```
*****
SUBCELL #
             1
IDAM F
               100000000.00000
CYCLES TO FAILURE XNF
*****
*****
SUBCELL #
             2
IDAM F
CYCLES TO FAILURE XNF 61360.2956047440
****
SUBCELL #
             3
IDAM F
CYCLES TO FAILURE XNF 61360.2956047440
******
             4
SUBCELL #
IDAM F
CYCLES TO FAILURE XNF 66438.5151084627
******
*****
** Current Damage At Increment 100 Applied Load Cycle
                                   1
SUBCELL # D
             1-D
 1 0.00000E+00 0.10000E+01
  2 0.149958E+00 0.850042E+00
  3 0.149958E+00 0.850042E+00
  4 0.632021E-03 0.999368E+00
SUBCELL #
             1
IDAM F
CYCLES TO FAILURE XNF 100000000.00000
****
SUBCELL #
             2
IDAM T
CYCLES TO FAILURE XNF 1167.36343322662
******
******
SUBCELL #
             3
IDAM T
CYCLES TO FAILURE XNF 1167.36343322663
******
SUBCELL #
             4
IDAM T
CYCLES TO FAILURE XNF 5456.41016344678
*****
*****
** Current Damage At Increment 200 Applied Load Cycle
                                   2
SUBCELL # D
             1-D
  1 0.00000E+00 0.100000E+01
  2 0.299638E+00 0.700362E+00
```

3 0.299638E+00 0.700362E+00 4 0.141115E-02 0.998589E+00 1 SUBCELL # IDAM T CYCLES TO FAILURE XNF 1000000000.00000 SUBCELL # 2 IDAM T CYCLES TO FAILURE XNF 709.630361973297 **** 3 SUBCELL # IDAM T CYCLES TO FAILURE XNF 709.630361973300 4 SUBCELL # IDAM T CYCLES TO FAILURE XNF 4380.85967672225 ** Current Damage At Increment 300 Applied Load Cycle 3 SUBCELL # D 1-D 1 0.00000E+00 0.100000E+01 0.449466E+00 0.550534E+00 2 3 0.449466E+00 0.550534E+00 4 0.243012E-02 0.997570E+00 ***** 1 SUBCELL # IDAM T CYCLES TO FAILURE XNF 100000000.00000 2 SUBCELL # IDAM T CYCLES TO FAILURE XNF 463.247417312908 **** SUBCELL # 3 IDAM T CYCLES TO FAILURE XNF 463.247417312908 **** 4 SUBCELL # IDAM T CYCLES TO FAILURE XNF 3619.74747832020 ****** ** Current Damage At Increment 400 Applied Load Cycle 4 SUBCELL # D 1-D 1 0.00000E+00 0.100000E+01 2 0.599071E+00 0.400929E+00

. .

```
3 0.599071E+00 0.400929E+00
  4 0.375666E-02 0.996243E+00
******
SUBCELL #
              1
IDAM T
CYCLES TO FAILURE XNF 100000000.00000
*******
******
SUBCELL #
              2
IDAM T
CYCLES TO FAILURE XNF 295.355917088531
********
******
SUBCELL #
              3
IDAM T
CYCLES TO FAILURE XNF 295.355917088531
*****
****
SUBCELL #
              4
IDAM T
CYCLES TO FAILURE XNF
                3018.74014755274
*****
*****
** Current Damage At Increment 500 Applied Load Cycle
                                      5
SUBCELL # D 1-D
  1 0.000000E+00 0.100000E+01
  2 0.748061E+00 0.251939E+00
  3 0.748061E+00 0.251939E+00
  4 0.539214E-02 0.994608E+00
SUBCELL #
              1
IDAM T
CYCLES TO FAILURE XNF 100000000.00000
*****
******
SUBCELL #
              2
IDAM T
CYCLES TO FAILURE XNF 166.907472034450
**********************
******
SUBCELL #
              3
IDAM T
CYCLES TO FAILURE XNF 166.907472034450
******
*****
SUBCELL #
              4
IDAM T
CYCLES TO FAILURE XNF 2532.39380892976
******
******
** Current Damage At Increment 600 Applied Load Cycle 6
SUBCELL # D 1-D
  1 0.00000E+00 0.100000E+01
  2 0.897013E+00 0.102987E+00
  3 0.897013E+00 0.102987E+00
  4 0.707338E-02 0.992927E+00
```

```
*****
SUBCELL #
             1
IDAM T
CYCLES TO FAILURE XNF 100000000.00000
****
****
             2
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF 62.0697834887973
*****
*********
             3
SUBCELL #
IDAM T
              62.0697834887934
CYCLES TO FAILURE XNF
4
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF 2148.40603399676
** Current Damage At Increment 700 Applied Load Cycle 7
      D
             1-D
SUBCELL #
  1 0.00000E+00 0.100000E+01
  2 0.987089E+00 0.129106E-01
  3 0.987089E+00 0.129106E-01
  4 0.796197E-02 0.992038E+00
1
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF 100000000.00000
******
2
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF 7.35691370876420
*****
3
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF 7.35691370914608
*****
*****
SUBCELL #
             4
IDAM T
CYCLES TO FAILURE XNF 1958.42907053144
****
** Current Damage At Increment 800 Applied Load Cycle
                                    8
SUBCELL # D 1-D
  1 0.000000E+00 0.100000E+01
  2 0.994882E+00 0.511765E-02
  3 0.994882E+00 0.511765E-02
  4 0.807672E-02 0.991923E+00
```

```
*****
SUBCELL #
             1
IDAM T
CYCLES TO FAILURE XNF
               100000000.00000
*****
             2
SUBCELL #
IDAM T
               2.90196867892969
CYCLES TO FAILURE XNF
******
******
SUBCELL #
             3
IDAM T
CYCLES TO FAILURE XNF 2.90196868203595
******
SUBCELL #
             4
IDAM T
CYCLES TO FAILURE XNF 1940.74760195980
*****
** Current Damage At Increment 900 Applied Load Cycle
                                    9
SUBCELL # D
              1-D
 1 0.00000E+00 0.100000E+01
 2 0.996420E+00 0.357991E-02
 3 0.996420E+00 0.357991E-02
 4 0.811000E-02 0.991890E+00
*****
SUBCELL #
            1
IDAM T
CYCLES TO FAILURE XNF 100000000.00000
******
*******
SUBCELL #
             2
IDAM T
CYCLES TO FAILURE XNF 2.02801986635817
*******
******
SUBCELL #
             3
IDAM T
CYCLES TO FAILURE XNF 2.02801987334260
*****
******
SUBCELL #
             4
IDAM T
CYCLES TO FAILURE XNF 1936.69355062971
******
** Current Damage At Increment *** Applied Load Cycle 10
SUBCELL # D 1-D
 1 0.00000E+00 0.100000E+01
 2 0.999034E+00 0.966462E-03
 3 0.999034E+00 0.966462E-03
  4 0.814345E-02 0.991857E+00
```

```
SUBCELL #
        1
IDAM T
CYCLES TO FAILURE XNF 100000000.00000
****
2
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF -999.0000000000
SUBCELL #
           3
IDAM T
CYCLES TO FAILURE XNF -999.0000000000
****
****
          4
SUBCELL #
IDAM T
CYCLES TO FAILURE XNF 1931.10439061903
```

6.13 Example M: Longitudinal Discrete Fiber Breakage

Problem Summary	ion Longitadi	<u>Har Brounder (Ber Broundge</u>	
Load Type:	Thermomechanical		
Load History:	Cyclic		
Load Control:	Strain		
Load Component:	11-direction (in fiber direction)		
Load History Data:	Cool-downfrom 900°C to 23°C, heat-up to 650°C, then hold temperature during mechanical load- ing $\dot{T} = 0.0152$ °C/sec, 0.0871 °C/sec $\dot{\varepsilon} = 1.0 \times 10^{-4}$ /sec, $\varepsilon_{max} = 0.012$, $\varepsilon_{min} = 0$. $\Delta t_{thermal} = 500.$, 40. sec.; $\Delta t_{mech} = 0.4$ sec.		
Micromechanics Model:	Triple Periodicity		
Fiber Packing Arrangement:	Square Pack, 35% fiber volume ratio		
Repeating Unit Cell:	1x8x14 ur m Forward Eu	hit cell with 28 square fibers (input anually) Ier	
Constituent Material Model:	Fiber: Matrix:	Elastic, transversely isotropic GVIPS - isotropic form	
Constituents:	Fiber: Matrix:	SCS-6 (temp. dep. properties input manually) TIMETAL21S	
Fiber Breakage:	Second implementation of debond model		
Fiber Breakage Parameters:	taken from room temperature SCS-6 fiber strength histogram with strengths decreased by		

Sample Input File for Longitudinal Discrete Fiber Breakage

effect.

5.8 % to account for elevated temperature

 $\Lambda = 1.0 x 10^{-5} \frac{i n^3}{k i p}, \qquad B = 10.\frac{1}{s}$

4

Note: In the case of longitudinal discrete fiber breakage, each fiber is modeled with an internal weak interface oriented normal to the fiber direction encompassing the entire fiber cross-section. This interface is then given a strength, σ_{DB} , which corresponds to the fiber ultimate strength. Thus, during simulated loading, when the longitudinal stress in the fiber reaches σ_{DB} , the fiber's internal interface debonds, and the longitudinal stress in the fiber begins to unload. This simulates a local fiber failure in a real composite.

In order to model longitudinal discrete fiber breakage as realistically as possible, a subcell containing 28 fibers has been used. Square-shaped fibers have been employed because longitudinal behavior is insensitive to fiber shape, and square fibers require the smallest number of subcells. Strength data for the SCS-6 fiber was taken from a vendor-supplied histogram. The room-temperature simulated distribution, as well as the actual strength distribution are shown in the subsequent figure. Note that the strengths employed in the example (in which the tensile simulation is performed at 650°C) were reduced by 5.8 % from the room-temperature values to account for the effect of the elevated temperature. In addition, the fiber strengths have been distributed over the strength ranges indicated by the histogram, rather than bunched up. That is, rather than providing 5 fibers with a strength of 600 ksi, the strengths of those 5 fibers were distributed between 575 ksi and 625 ksi (prior to the 5.8 % reduction).

The results of this example shown in the subsequent figure have been plotted only to an applied strain level of 0.0096 because at this point the slope of the predicted global stress-strain curve becomes negative. We are treating this as a criterion for simulated global failure of the composite.

```
test of longitudinal discrete fiber breakage
*PRINT
 NPL=1 %
*LOAD
 LCON=3 LOP=1 LSS=1 %
*MECH
 NPTW=5 TI=0.,24000.,57600.,64800.,64920. LO=0.0,0.0,0.,0.,0.012 %
*THERM
 NPTT=5 TI=0.,24000.,57600.,64800.,64920. TE=900.,534.583,23.,650.,650. %
*MODEL
 MOD=2 %
*SOLVER
 NTF=1 NPTS=5 TIM=0.,24000.,57600.,64800.,64920. STP=500.,40.,40.,0.4 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=2 MAT=U IFM=1
 NTP=6
 TEM=21.1,204.44,315.56,426.67.537.78,871.11
 EA=57.0E3,55.98E3,55.4E3,54.82E3,54.24E3,53.36E3
 ET=57.0E3.55.98E3.55.4E3.54.82E3.54.24E3.53.36E3
 NUA=0.25.0.25.0.25.0.25.0.25.0.25
 NUT=0.25,0.25,0.25,0.25,0.25,0.25
 GA=22.8E3,22.392E3,22.16E3,21.928E3,21.696E3,21.344E3
 ALPA=3.564E-6,3.618E-6,3.726E-6,3.906E-6,4.068E-6,4.572E-6
 ALPT=3.564E-6.3.618E-6.3.726E-6.3.906E-6.4.068E-6.4.572E-6
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=2 MAT=A %
*MRVE
 IDP=99
 NA=1 NB=8 NG=14
 D=1.0
#NOTE: VF = 35%
 H=0.5916,0.4084,0.5916,0.4084,0.5916,0.4084,0.5916,0.4084
 L=0.5916,0.4084,0.5916,0.4084,0.5916,0.4084,0.5916,0.4084 &
  0.5916,0.4084,0.5916,0.4084,0.5916,0.4084
 CM=F1.M1.F1.M1.F1.M1.F1.M1
 CM=M1,M1,M1,M1,M1,M1,M1,M1
 CM=F1,M1,F1,M1,F1,M1,F1,M1
 CM=M1,M1,M1,M1,M1,M1,M1,M1
 CM=F1,M1,F1,M1,F1,M1,F1,M1
 CM=M1,M1,M1,M1,M1,M1,M1,M1
 CM=F1,M1,F1,M1,F1,M1,F1,M1
 CM=M1,M1,M1,M1,M1,M1,M1,M1
 CM=F1,M1,F1,M1,F1,M1,F1,M1
 CM=M1,M1,M1,M1,M1,M1,M1,M1
```

CM=F1,M1,F1,M1,F1,M1,F1,M1 CM=M1,M1,M1,M1,M1,M1,M1,M1 CM=F1,M1,F1,M1,F1,M1,F1,M1 CM=M1,M1,M1,M1,M1,M1,M1,M1 *DEBOND NII=28 DBCH=2 NAI=1 NBI=1 NGI=1 FACE=1 BDN=357 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=1 NGI=3 FACE=1 BDN=403 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=1 NGI=5 FACE=1 BDN=469 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=1 NGI=7 FACE=1 BDN=565 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=1 NGI=9 FACE=1 BDN=496 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=1 NGI=11 FACE=1 BDN=503 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=1 NGI=13 FACE=1 BDN=612 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=1 FACE=1 BDN=450 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=3 FACE=1 BDN=655 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=5 FACE=1 BDN=603 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=7 FACE=1 BDN=512 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=9 FACE=1 BDN=407 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=11 FACE=1 BDN=517 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=3 NGI=13 FACE=1 BDN=556 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=1 FACE=1 BDN=716 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=3 FACE=1 BDN=593 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=5 FACE=1 BDN=631 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=7 FACE=1 BDN=640 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=9 FACE=1 BDN=678 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=11 FACE=1 BDN=584 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=5 NGI=13 FACE=1 BDN=488 GCN=0.00001 BCN=10. TOLN=1 & BDS=100 GCS=0.1 BCS=100 TI=0 DBCH=2 NAI=1 NBI=7 NGI=1 FACE=1 BDN=575 GCN=0.00001 BCN=10. TOLN=1 &

```
BDS=100 GCS=0.1 BCS=100 TI=0
DBCH=2 NAI=1 NBI=7 NGI=3 FACE=1 BDN=697 GCN=0.00001 BCN=10. TOLN=1 &
BDS=100 GCS=0.1 BCS=100 TI=0
DBCH=2 NAI=1 NBI=7 NGI=5 FACE=1 BDN=754 GCN=0.00001 BCN=10. TOLN=1 &
BDS=100 GCS=0.1 BCS=100 TI=0
DBCH=2 NAI=1 NBI=7 NGI=7 FACE=1 BDN=546 GCN=0.00001 BCN=10. TOLN=1 &
BDS=100 GCS=0.1 BCS=100 TI=0
DBCH=2 NAI=1 NBI=7 NGI=9 FACE=1 BDN=537 GCN=0.00001 BCN=10. TOLN=1 &
BDS=100 GCS=0.1 BCS=100 TI=0
DBCH=2 NAI=1 NBI=7 NGI=11 FACE=1 BDN=664 GCN=0.00001 BCN=10. TOLN=1 &
BDS=100 GCS=0.1 BCS=100 TI=0
DBCH=2 NAI=1 NBI=7 NGI=13 FACE=1 BDN=622 GCN=0.00001 BCN=10. TOLN=1 &
BDS=100 GCS=0.1 BCS=100 TI=0
*CURVE
 NP=3 %
*MACRO
 NT=2
 NC=1 X=1 Y=7 NAM=LONG-M
 NC=2 X=39 Y=1 NAM=LONG-T
*MICRO
 NT=3
 NC=1 CELL=1 X=1 Y=7 NAM=LONG-1
 NC=2 CELL=17 X=1 Y=7 NAM=LONG-2
 NC=3 CELL=33 X=1 Y=7 NAM=LONG-3 %
*END
```



Room Temperature SCS-6 Fiber Strength Histogram



Longitudinal Prediction of SCS-6/Timetal 21S with 35% fiber volume content at 650°C

6.14 Example N: Plain Weave Composite

Sample Input File for Plain Weave Composite: Step 1 - Determine Yarn Properties

Problem Summary

Load Type:	Mechanic	Mechanical		
Load Component:	33-direction	33-direction (transverse to fiber)		
Load History:	Monotoni	Monotonic		
Load Control:	Strain	Strain		
Load History Data:	$\dot{\varepsilon} = 1.0 \times 10$ $\Delta t_{mech} = -$	$\dot{\varepsilon} = 1.0 \times 10^{-4}$ /sec, $\varepsilon_{max} = 0.018$, $\varepsilon_{min} = 0$. $\Delta t_{mech} = 4$. sec.		
Micromechanics Model:	Double Pe	Double Periodicity		
Fiber Packing Arrangement:	Square Pa	Square Pack, R = 1., 65% fiber volume ratio		
Repeating Unit Cell:	26x26 circ	26x26 circular fiber cross-section approx.		
Integration Algorithm:	Forward E	Forward Euler		
Constituent Material Model:	Fiber: Matrix:	Elastic, transversely isotropic Elastic, isotropic		
Constituents:	Fiber: Matrix:	AS-4 Graphite Fiber (properties input manually) PMR-15 Epoxy (properties input manually)		

Note: Step 1 of analyzing a woven composite with MAC/GMC involves determining the effective properties of the fiber/matrix yarns which reinforce the woven composites. For the elastic example shown, this is quite simple and involves determining only the five transversely isotropic elastic properties and the two transversely isotropic CTEs for the unidirectional fiber/matrix yarn. These effective properties will then be employed in step 2 (using ncmd = 9, transversely isotropic model) to analyze the actual woven composite unit cell. Inelastic analysis of woven composites is possible with MAC/GMC, but the inelastic characterization of the unidirectional fiber/matrix yarns (for implementation with one of the transversely isotropic inelastic constitutive models) becomes more complex. Future versions of MAC/GMC may include automated routines to aid in the analysis of thermo-inelastic woven composites.

```
analysis of a graphite/epoxy yarn
*PRINT
NPL=3 %
*LOAD
LCON=2 LOP=3 LSS=1 %
*MECH
NPTW=2 TI=0.,108. LO=0.,0.018 %
*MODEL
 MOD=1 %
*SOLVER
NTF=1 NPTS=2 TIM=0.,108. STP=4. %
*FIBER
NFIBS=1
 NF=1 MF=6 NDPT=1 MAT=U IFM=1&
 EL=31.E6,2.E6,0.2,0.25,2.E6,-0.55E-6,5.6E-6 %
*MATRIX
 NMATX=1
 NM=1 MM=6 NDPT=1 MAT=U IFM=1 &
 EL=0.5E6,0.5E6,0.41,0.41,0.1773E6,57.E-6,57.E-6 %
*MRVE
 IDP=13 VF=0.65 R=1.0
*CURVE
NP=1 %
*MACRO
 NT=1
 NC=1 X=3 Y=9 NAM=YARN
*END
```

Results for fiber/matrix yarn:

EFFECTIVE ENGINEERING MODULI

E11S= .203E+08 N12S= .272E+00 E22S= .134E+07 N23S= .432E+00 E33S= .134E+07 G23S= .359E+06 G13S= .596E+06 G12S= .596E+06

EFFECTIVE THERMAL EXPANSION COEFFICIENTS .493E-07 .310E-04 .310E-04

Sample Input File for Plain Weave Composite: Step 2 - Composite Behavior

Problem Summary			
Load Type:	Mechanical		
Load Component:	33-direction (in the forcement)	plane of the woven rein-	
Load History:	Monotonic		
Load Control:	Strain		
Load History Data:	$\dot{\varepsilon} = 1.0 \times 10^{-4}$ /sec, $\varepsilon_{max} = 0.018$, $\varepsilon_{min} = 0$. $\Delta t_{mech} = 4$. sec.		
Micromechanics Model:	Triple Periodicity		
Repeating Unit Cell:	Input manually - approximates a plain weave reinforced composite		
Integration Algorithm:	Forward Euler		
Constituent Material Model:	Fiber/Matrix Yarns:	Elastic, locally trans-	
	Pure Matrix:	Elastic, isotropic	
Constituents:	Fibers 1 - 4:	V _f = 65% AS-4/PMR-15 yarns with different fiber orientations (properties	

Note: MAC/GMC currently limits the number of fiber and matrix constituents to 4 each.

Matrix 1:

Matrix 2, 3:

input manually)

input manually)

input manually)

PMR-15 Epoxy (properties

 $V_{f} = 65\% \text{ AS-4/PMR-15}$

yarns with different fiber orientations (properties

Note: Step 2 of analyzing a woven composite with MAC/GMC involves assembling a repeating unit cell that represents the woven composite using the effective fiber/matrix yarn properties determined in step 1. The repeating unit cell employed for a plain weave composite in this example is shown in the figure below. The heterogeneous subcells in this figure are represented by effective fiber/matrix yarn properties, while the local fiber direction of these subcells is accounted for by the D vector in the locally transversely isotropic constitutive model.



User defined IDP for woven architecture (not drawn to scale)

```
plain weave reinforced composite
*PRINT
 NPL= 3%
*LOAD
 LCON=2 LOP=3 LSS=1 %
*MECH
 NPTW=2 TI=0.,108. LO=0.,0.018 %
*THERM
 NPTT=2 TI=0..108. TE=23..23.
*MODEL
 MOD=2 %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,108. STP=4. %
*FIBER
 NFIBS=4
 NF=1 MF=9 NDPT=1 MAT=U IFM=1 &
 EL=20.3E6,1.34E6,0.272,0.432,0.596E6,0.0493E-6,31.E-6 D=0.,0.,1. %
 NF=2 MF=9 NDPT=1 MAT=U IFM=1&
 EL=20.3E6,1.34E6,0.272,0.432,0.596E6,0.0493E-6,31.E-6 D=0.,1.,0. %
 NF=3 MF=9 NDPT=1 MAT=U IFM=1&
 EL=20.3E6,1.34E6,0.272,0.432,0.596E6,0.0493E-6,31.E-6 D=0.25,1.,0. %
 NF=4 MF=9 NDPT=1 MAT=U IFM=1&
 EL=20.3E6,1.34E6,0.272,0.432,0.596E6,0.0493E-6,31.E-6 D=-0.25,1.,0. %
*MATRIX
 NMATX=3
 NM=1 MM=6 NDPT=1 MAT=U IFM=1 &
 EL=0.5E6,0.5E6,0.41,0.41,0.1773E6,57.E-6,57.E-6 %
 NM=2 MM=9 NDPT=1 MAT=U IFM=1 &
 EL=20.3E6,1.34E6,0.272,0.432,0.596E6,0.0493E-6,31.E-6 D=0.25,0.,1. %
 NM=3 MM=9 NDPT=1 MAT=U IFM=1 &
 EL=20.3E6,1.34E6,0.272,0.432,0.596E6,0.0493E-6,31.E-6 D=-0.25,0.,1. %
*MRVE
 IDP=99
 NA=4 NB=4 NG=4
 D=0.25,0.25,0.25,0.25
 H=1.,1.,1.,1.
 L=1.,1.,1.,1.
 CM=F1,M1,F2,M1
 CM=F1,F3,F2,F4
 CM=F2,F3,F1,F4
 CM=F2,M1,F1,M1
 CM=M1,M1,M1,M1
 CM=M3,M1,M2,M1
 CM=M3,M1,M2,M1
 CM=M1,M1,M1,M1
 CM=F2,M1,F1,M1
```
CM=F2,F4,F1,F3 CM=F1,F4,F2,F3 CM=F1,M1,F2,M1 CM=M1,M1,M1,M1 CM=M2,M1,M3,M1 CM=M2,M1,M3,M1 CM=M1,M1,M1,M1 *CURVE NP=1 % *MACRO NT=1 NC=1 X=3 Y=9 NAM=WEAVE *END Results for woven composite:

EFFECTIVE ENGINEERING MODULI

E11S= .920E+06 N12S= .288E+00 E22S= .158E+07 N23S= .151E+00 E33S= .158E+07 G23S= .357E+06 G13S= .351E+06 G12S= .351E+06

EFFECTIVE THERMAL EXPANSION COEFFICIENTS

.548E-04 .261E-04 .261E-04 .000E+00 .318E-22 -.567E-23





6.15 Example O: FEAMAC Analysis

Problem Summary: Thermomechanical; Load Control (ABAQUS) Load Type: From ABAQUS: Load History Data: STEP 1: Cool down Temperature: 860.0 to 427.0°C Strain: 0.0 - 0.0 STEP 2: Transverse loading Temperature held fixed at 427.0°C Strain: 0.0 to 0.015 **Double Periodic** Micromechanics model: Square Fiber (idp=2) Fiber Packing Arrangement: Triangular Pack at 33% fiber volume ratio Circular Fiber (idp=6) Rectangular Pack at 33% fiber volume ratio Forward Euler (ABAQUS) Integration Algorithm: Elastic (ncmd = 7) **Constituent Material Model:** Fiber: Matrix: TGVIPS (ncmd = 7) Fiber: SCS-6 **Constituents:** Matrix: Ti-6-4 (isotropic) First implementation of debond model Debonding: Consists of 2 (8-noded brick) elements each Finite Element Model: with a different material. In this way the example demonstrates the use of multiple material groups.

Sample Input Files For FEAMAC Analysis

FEAMAC Note: All data that is struck-out in the listing is not required by **FEAMAC** and is ignored if present. Warnings are generated informing the user that the data provided is being ignored. **FEAMAC** can be employed with any size of a finite element mesh. However, **FEAMAC** requires that an appropriate amount of state variable space be allocated for each user material found in the finite element model. The amount of state variable space to be allocated for a given user material is based on the size of the RVE requested in the corresponding **MAC**/ **GMC** input deck. The number of state variables must be less than 1000. The ***DEPVAR** option under each of the ***USER MATERIAL** sections should be used to define the size of this space. The number of <u>CONSTANTS</u> should be set to <u>one</u> on the ***USER MATERIAL** line and on the following line the <u>value should be set to</u> <u>one</u>. If the material model is linear elastic and no micro curve data files have been requested then no state variable space is required for that material.

To determine the amount of space to request use the following equation:

NG = Number of cells in the gamma direction

NB = Number of cells in the beta direction

NDEPVAR = Number of state variables to be to be provided to the *DEPVAR option.

NDEPVAR = 46 + (42 * NG * NB) < 1000

Therefore, based on the predefined RVE's in section 4.2.13:

IDP	NDEPVAR	
1	214	
2	718	
3	718	· · · · · · · · · · · · · · · · · · ·
4	718	
6	2104	Too Large: Must use alternate method described below
9	2104	Too Large: Must use alternate method described below
11	214	
13	28,438	Too Large: Must use alternate method described below

EXAMPLE:

*MATERIAL,NAME=MACDECK1 *USER MATERIAL, CONSTANTS=1 1 *DEPVAR 214 ... Where MACDECK1 references an IDP of size 2x2.

ALTERNATE METHOD:

If a chosen IDP requires more than 1000 state variables then the following alternate procedure must be followed:

On the *USER MATERIAL line set the number of <u>CONSTANTS</u> equal to 1 and on the following line set the value of this constant to be <u>two</u>. Also set the number of state variables on the *DEPVAR option to zero.

EXAMPLE: ... *MATERIAL,NAME=MACDECK2 *USER MATERIAL, CONSTANTS=1 2 *DEPVAR 0 ...

Where MACDECK2 references an IDP that would require more than 1000 state variables (e.g. IDP = 9).

FEAMAC Note: This alternate procedure will cause execution times to increase greatly.

FEAMAC Note: It is highly recommended that linear elastic materials with no requests for micro curve data be used in all areas of the model except in areas of suspected material nonlinearity, irrespective of memory approach.

MAC/GMC Input Listings: MACMAN1:

```
FEAMAC DEMO - Material group No. 1, identified in ABAQUS deck as MACMAN1
*PRINT
 NPL=1 %
*LOAD
LCON=3 LOP=3 LSS=1-%
*MECH
_NPTW=3 TI=0.0, 58000.0, 58150.0 LO=0.0, 0.0, 0.015 %
*THERM
--NPTT=3 TI=0.0, 58000.0, 58150.0 TE=860.0, 427.0, 427.0 %
*MODEL
 MOD=1 %
*SOLVER
-NTF=1&
-NPTS=3 TIM=0.0, 58000.0, 58150.0 STP=2000.0, 1.0 %
*FIBER
 NFIBS=1
 NF=1 MF=4 NDPT=1 MAT=U IFM=1 &
 EL=58.E3, 0.32, 3.5E-06 &
 VI=0.8E-9,1.E20,0.1E-5,0.,0.85E-4,0.05,1.,1.,1.,3.3,1.8,1.35,1.,0.01 %
*MATRIX
 NMATX=1
 NM=1 MM=7 NDPT=2 MAT=A D=1.0, 1.0, 1.0 %
*MRVE
 IDP=2 VF=0.33 %
*DEBOND
NII=2
DBCH=1 NBI=1 NGI=2 FACE=2 RN=0.1 BDN=35.0 RS=0.0 BDS=0.0 TI=58000.0
DBCH=1 NBI=1 NGI=3 FACE=2 RN=0.1 BDN=35.0 RS=0.0 BDS=0.0 TI=58000.0
%
*CURVE
 NP=1 %
*MACRO
 NT=1
 NC=1 X=3 Y=9 NAM=macman1_s3_e3n %
*END
```

MAC/GMC Input Listings: MACMAN2:

FEAMAC DEMO - Material group No. 2, identified in ABAQUS deck as MACMAN2 *PRINT **NPL=1 %** *LOAD LCON=3 LOP=3 LSS=1-% *MECH -- NPTW=3 TI=0.0, 58000.0, 58150.0 LO=0.0, 0.0, 0.015 % *THERM -NPTT=3 TI=0.0, 58000.0, 58150.0 TE=860.0, 427.0, 427.0 % *MODEL MOD=1 % ***SOLVER** -NTF=1-& -NPTS=3 TIM=0.0, 58000.0, 58150.0 STP=2000.0, 1.0 % *FIBER NFIBS=1 NF=1 MF=9 NDPT=2 MAT=A D=1.0, 0.0, 0.0 % *MATRIX NMATX=1 NM=1 MM=7 NDPT=2 MAT=A D=1.0, 1.0, 1.0 % *MRVE IDP=6 VF=0.33 R=0.75% *DEBOND Nil=10DBCH=1 NBI=4 NGI=1 FACE=2 RN=0.1 BDN=14.0 RS=0.0 BDS=0.0 TI=58000.0 DBCH=1 NBI=4 NGI=6 FACE=2 RN=0.1 BDN=14.0 RS=0.0 BDS=0.0 TI=58000.0 # DBCH=1 NBI=2 NGI=3 FACE=2 RN=0.1 BDN=22.0 RS=0.0 BDS=0.0 TI=58000.0 DBCH=1 NBI=2 NGI=4 FACE=2 RN=0.1 BDN=22.0 RS=0.0 BDS=0.0 TI=58000.0 # DBCH=1 NBI=3 NGI=2 FACE=2 RN=0.1 BDN=18.0 RS=0.0 BDS=0.0 TI=58000.0 DBCH=1 NBI=3 NGI=5 FACE=2 RN=0.1 BDN=18.0 RS=0.0 BDS=0.0 TI=58000.0 # DBCH=1 NBI=6 NGI=3 FACE=2 RN=0.1 BDN=22.0 RS=0.0 BDS=0.0 TI=58000.0 DBCH=1 NBI=6 NGI=4 FACE=2 RN=0.1 BDN=22.0 RS=0.0 BDS=0.0 TI=58000.0 # DBCH=1 NBI=5 NGI=2 FACE=2 RN=0.1 BDN=18.0 RS=0.0 BDS=0.0 TI=58000.0 DBCH=1 NBI=5 NGI=5 FACE=2 RN=0.1 BDN=18.0 RS=0.0 BDS=0.0 TI=58000.0 # *CURVE NP=1 % *MACRO NT=1 NC=1 X=3 Y=9 NAM=macman2 s3 e3n % *END

ABAQUS 5.8 Input Listing:

*HEADING VERIFICATION OF FEAMAC *NODE,NSET=ALL 1, 0.0, 0.0, 0.0 2, 0.0, 1.0, 0.0 3, 0.0, 1.0, 1.0 4, 0.0, 0.0, 1.0 5, 1.0, 0.0, 0.0 6, 1.0, 1.0, 0.0 7, 1.0, 1.0, 1.0 8, 1.0, 0.0, 1.0 9, 2.0, 0.0, 0.0 10, 2.0, 1.0, 0.0 11, 2.0, 1.0, 1.0 12, 2.0, 0.0, 1.0 ***ORIENTATION, NAME=TRANS** 0.0, 0.0, 1.0, 0.0, 1.0, 0.0 *ELEMENT, TYPE=C3D8, ELSET=EL1 1, 1, 2, 3, 4, 5, 6, 7, 8 *ELEMENT, TYPE=C3D8, ELSET=EL2 2, 5, 6, 7, 8, 9, 10, 11, 12 *SOLID SECTION, MATERIAL=MACMAN1, ELSET=EL1, ORIENTATION=TRANS 1.0 *SOLID SECTION, MATERIAL=MACMAN2, ELSET=EL2, ORIENTATION=TRANS 1.0 *MATERIAL,NAME=MACMAN1 *USER MATERIAL, CONSTANTS=1 1 *DEPVAR 718 ***USER DEFINED FIELD** *EXPANSION, TYPE=ANISO, USER *MATERIAL,NAME=MACMAN2 *USER MATERIAL, CONSTANTS=1 2 ***USER DEFINED FIELD** *EXPANSION, TYPE=ANISO, USER ***USER SUBROUTINE** *INCLUDE, INPUT=/home/smdant/macdev/feabench/man_demo/umacman.f ***INITIAL CONDITIONS, TYPE=TEMPERATURE** ALL, 860.0 *INITIAL CONDITIONS, TYPE=FIELD, VARIABLE=1 ALL, 1.0 **_<u>_____</u>

```
*STEP, AMPLITUDE=RAMP, INC=2000000
```

```
STEP ONE - COOL DOWN
*STATIC, DIRECT
2000.0, 58000.0
*TEMPERATURE
ALL, 427.0
*BOUNDARY
1, 1, 3
2, 1
2,3
3, 1
4, 1, 2
5, 2, 3
6,3
8,2
9, 2, 3
10, 3,
12, 2
*NSET, NSET=ALL
1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12
*EL PRINT, FREQ=1000
S
*EL PRINT, FREQ=1000
Ε
*EL PRINT, FREQ=1000
THE, TEMP
*NODE PRINT, FREQ=1000, NSET=ALL
U
*NODE PRINT, FREQ=1000, NSET=ALL
CF
*NODE PRINT, FREQ=1000, NSET=ALL
RF
*END STEP
**_____
*STEP, AMPLITUDE=RAMP, INC=2000000
STEP TWO - LOAD UP
*STATIC, DIRECT
1.0, 150.0
*BOUNDARY
1, 1, 3
2, 1
2,3
3, 1
4, 1, 2
5, 2, 3
6, 3
8, 2
```

9, 2, 3
10, 3,
12, 2
9, 1, 1, 0.015
10, 1, 1, 0.015
11, 1, 1, 0.015
12, 1, 1, 0.015
*TEMPERATURE
ALL, 427.0
*END STEP
**

The following figure was obtained from the macro x-y plot data files produced by the present example.



.

ABAQUS UMAT Subroutine Listing:

C COMMON TO SOLIDS AND SHELLS
C 111111111222222222233333333344444444445555555555
C ////_//_/////////////////////////////
C
C ORDER OF ABAQUS FUNCTION CALLS C
C – UMAT (SOLID) C (if needed) /
C USDFLD -=> UEXPAN -=< or
C - UGENS (SHELL) <-= TBD
C
SUBROUTINE USDFLD(FIELD, STATEV, PNEWDT, DIRECT, T, CELENT, & TIME, DTIME, CMNAME, ORNAME, NFIELD, NSTATV, NOEL, NPT, & LAYER, KSPT, KSTEP, KINC, NDI, NSHR)
INCLUDE `ABA_PARAM.INC'
PARAMETER (ZERO = 0.0D0, HALF = 0.5D0) PARAMETER (ONE = 1.0D0, TWO = 2.0D0, THREE = 3.0D0) PARAMETER (NFROM_UTHERM = 1, NFROM_UMECH = 2)
CHARACTER*8 CMNAME, ORNAME DIMENSION FIELD(NFIELD), STATEV(NSTATV), DIRECT(3,3) DIMENSION T(3,3), TIME(2)
CHARACTER*80 FNAME CHARACTER*80 EXTSION
CHARACTER*80 DFNAME
CHARACTER*80 TSTRING
CHARACTER*80 TNAME1 CHARACTER*80 TNAME2
INTEGER AB_PRINT
INTEGER AL_STORE
LOGICAL SIGAL_STATV
CHARACTER*8 FLGRAY(15) DIMENSION ARRAY(15), JARRAY(15)
COMMON / KINCINFO / AB_STEP, INC, NCYCLE

COMMON / KTIME / STEP_TIME, TOT_TIME, GDT COMMON / KMACFLAGS / NFIRSTTIME, NFIRSTELEM, NFIRSTIP, AL_STORE, N_PEN(100), N_PIN(100), NPEL, & N_PATPEN(100), N_PATPIN(100), NPATPEL, & AB_PRINT, GAB_PRINT, SIGAL_STATV, NCALL, NFEA_FROM ŵ COMMON / KABNAMES / PATH, FNAME, DFNAME, TNAME1, TNAME2, EXTSION COMMON / KPREMAC / NDU, NNOEL, NNPT, NNDI, NNSHR COMMON / KTHERMONLY / DTHERMSTRESS(6), DTHERMSTRAIN(6), DDTHERMSTRAIN_DT(6), THERMSTRESS(6), THERMSTRAIN(6) & DATA NFIRSTELEM / 0 / DATA NFIRSTIP / 0 / DATA NFIRSTTIME / 1 / $NFEA_FROM = 0$ C INITIALIZE NNOEL = NOEL NNPT = NPTNNDI = NDI NNSHR = NSHR NCALL = 0IF (NFIRSTTIME .EQ. 1) THEN CALL FEAMAC_INIT (NOEL, NPT) ENDIF AB PRINT = GAB_PRINT C DEFINE MAC INPUT DECK NAME CALL FILL_STRING(TSTRING, ``) CALL FILL_STRING(FNAME, ``) CALL STRING_COPY (FNAME, CMNAME) CALL STRING APPEND (FNAME, EXTSION) TSTRING = PATH CALL STRING_APPEND(TSTRING, FNAME) FNAME = TSTRING C DEFINE STEP INFORMATION IF ((NOEL .EQ. NFIRSTELEM) .AND. (NPT .EQ. 1) £ .AND. ((KINC - 1) .NE. INC)) THEN Sc. NCYCLE = -1ENDIF IF ((NOEL .EQ. NFIRSTELEM) A .AND. (NPT .EQ. 1)) THEN NCYCLE = NCYCLE + 1

.

ENDIF

```
AB_STEP = KSTEP
GDT = DTIME
STEP_TIME = TIME(1)
TOT_TIME = TIME(2)
INC = KINC - 1
```

C WE CAN MAKE CALLS TO GETVRM FROM HERE TO GET ADDTIONAL INFO IF NEEDED CALL GETVRM('THE', ARRAY, JARRAY, FLGRAY, JRDC)

```
DO J = 1, 6
THERMSTRAIN(J) = ARRAY(J)
THERMSTRESS(J) = 0.0
DTHERMSTRESS(J) = 0.0
DTHERMSTRAIN(J) = 0.0
DDTHERMSTRAIN_DT(J) = 0.0
ENDDO
```

IF	(AB_PRINT	GE. 1) THEN	
	WRITE(NDU,	·) · · ·	
	WRITE (NDU,	*)	
&	`=====	*======================================	
	WRITE (NDU,	') '-=> START OF USDFLD <=-'	
	WRITE (NDU,	') '-=> VALUE OF STEP TIME =', TIME(1), STEP_TIM	Ε
	WRITE (NDU,	<pre>', TIME(2), TOT_TIME</pre>	
	WRITE (NDU,	() '=> DTIME = ', DTIME, GDT	
	WRITE (NDU,	') '-=> KSTEP =', KSTEP, AB_STEP	
	WRITE (NDU,	(-=) KINC =', KINC, INC+1	
	WRITE (NDU,	') '-=> NOEL =', NOEL, NNOEL	
	WRITE (NDU,	(-=> NPT = (, NPT, NNPT)	
	WRITE (NDU,	*) '-=> NDI =', NDI	
	WRITE (NDU,	*) $'=>$ NSHR = ', NSHR	
	WRITE (NDU.	*) '-=> NCYCLE =', NCYCLE	
	WRITE (NDU,	*) '-=> NFIRSTELEM =', NFIRSTELEM	
	WRITE (NDU,	*) '-=> NFIRSTIP =', NFIRSTIP	
	WRITE (NDU,	*) '-=> NCALL =', NCALL	
	WRITE (NDU,	*) '-=> FNAME = ', FNAME, ' '	

ENDIF

RETURN END



```
С
С
С
   11
               \/
         1
С
              1
                1
С
С
С
                       1
С
              1
                11
С
           /_/
               11
                  1
                     1
С
                     1_/
С
          ******
С
  *** BEGIN USER EDITS ***
С
C *** USER NEEDS TO CHANGE THE NEXT LINES AS NEEDED...
C PATH TO WORKING DIRECTORY
      PATH = '/home/smdant/macdev/feabench/man_demo'
C NOTE: ABAQUS WILL CONVERT THE MATERIAL NAME TO UPPER CASE.
C MAC INPUT DECK NAME EXTENSION
      EXTSION = '.MAC'
C THE FOLLOWING IS ONLY USED TO RESOLVE VERSION DIFFERENCES BETWEEN
  SUN AND SGI
C NAME OF DIAGNOSTIC FILE
      DFNAME = 'MACDIAG'
      MAC PLOT FILES:
С
С
      - # OF POINTS TO PLOT, I.E # OF PAIRS TO FOLLOW
      NPEL = 2
C
      - ELEMENT # AND INTEGRATION POINT # PAIRS TO PLOT
            N PEN - ELEMENT NUMBERS
С
С
            N_PEN - INTEGRATION POINT NUMBERS
      J = 1
      N_PEN(J) = 1
      N_PIN(J) = 1
      J = J + 1
      N_PEN(J) = 2
      N_PIN(J) = 1
      PATRAN PLOT FILES:
C
С
       - # OF POINTS TO PLOT PATRAN DATA, I.E # OF PAIRS TO FOLLOW
      NPATPEL = 0
C
      - ELEMENT # AND INTEGRATION POINT # PAIRS TO PLOT PATRAN DATA
            N_PATPEN - ELEMENT NUMBERS
С
С
            N_PATPEN - INTEGRATION POINT NUMBERS
С
       J = 1
       N_PATPEN(J) = 1
С
С
       N_PATPIN(J) = 1
С
       \mathbf{J} = \mathbf{J} + \mathbf{1}
       N_PATPEN(J) = 2
С
С
       N_PATPIN(J) = 1
C THE FOLLOWING IS DEPRECIATED OPTION AND MAY NOT BE SUPPORTED
```

C IN THE NEXT RELEASE OF FEAMAC. IT'S USE IS NOT RECOMMENDED.

.



```
TSTRING = PATH
     CALL STRING_APPEND(TSTRING, '/ ')
     PATH = TSTRING
     TSTRING = PATH
     CALL STRING_APPEND(TSTRING, DFNAME)
     DFNAME = TSTRING
     DO 100 J = 1, NPEL
        IF ( (N_PIN(J) .LT. 1) .OR. (N_PIN(J) .GT. 8) ) THEN
          N_PIN(J) = 1
        ENDIF
100 CONTINUE
     NDU = 7
C UN-COMMENT NEXT 2 LINES IF YOUR ON A SGI
        NDU = 18
С
         OPEN(UNIT = NDU, FILE = DFNAME, STATUS = 'UNKNOWN')
С
     WRITE(NDU, *) 'BEGIN'
     NFIRSTELEM = NOEL
     NFIRSTIP = NPT
     NCYCLE = -1
     GAB_PRINT = AB_PRINT
     RETURN
     END
C
  C23456789012345678901234567890123456789012345678901234567890123456789
  012
С
Ç
   1111
           11
С
  /////////////
                    1
  11_1111
                   1/1
С
С
      _/_/ /_/_/
C
C
  C23456789012345678901234567890123456789012345678901234567890123456789
  012
     SUBROUTINE UMAT (STRESS, STATEV, DDSDDE, SSE, SPD, SCD, RPL,
DDSDDT, DRPLDE, DRPLDT, STRAN, DSTRAN, TIME, DTIME, TEMP,
     æ
          DTEMP, PREDEF, DPRED, CMNAME, NDI, NSHR, NTENS, NSTATV,
    &
          PROPS, NPROPS, COORDS, DROT, PNEWDT, CELENT, DFGRD0,
    æ
          DFGRD1, NOEL, NPT, LAYER, KSPT, KSTEP, KINC)
    &
C
  C23456789012345678901234567890123456789012345678901234567890123456789
  012
С
     STRESS(NTENS) - PASSED IN AS THE STRESSES TENSOR AT THE BEGINNING
С
          OF THE INCREMENT, SIGMA_I_J. (THIS SHOULD BE UPDATED IN
С
С
          THIS ROUTINE.)
          -IN/OUT
С
```

C C

Ċ С

С

00000000

С

C C C

С

С

С

C

С

С

С

С

С

С

```
С
      STATEV(NSTATV) - SOLUTION DEPENDANT STATE VARIABLES.
С
          -IN/OUT
С
            Format of state variable space ...
С
C
С
С
                   total strain
                                   (1-6)
"MACRO" quantities when
                   cauchy stress
                                   (7 - 12)
                                                micromechanics model
                                                or continuum model
                   inelastic strain (13-18)
                                                is used.
                   state variable(s) (19-30)
                   (space for 2 (6x1) vectors)
                   thermal strain
                                    (31 - 36)
                   temperature
                                     (37)
                   future use
                                    (38 - 40)
                   future use
                                     (41 - 43)
                   future use
                                     (44 - 46)
                      total strain
                                    (47 - 52)
                                               "MICRO" quantities:
                   cauchy stress
                                   (53-58)
                                                this block of NSVARS = 42
                 inelastic strain (59-64)
                L
                                            is repeated for each subcell
                   internal state variable(s)
                                    (65-76)
                   (space for 2 [6x1] vectors)
                   thermal strain
                                   (77-82)
                   rn & rd debond
                                    (83-88)
                   parameters
                   :
                                     (89 - ...)
                             etc.
     DDSDDE(NTENS, NTENS) - JACOBIAN MATRIX OF THE CONSTITUTIVE
          MODEL.
          -OUT
     SSE - SPECIFIC ELASTIC STRAIN ENERGY.
          -OUT
     SPD - PLASTIC DISSIPATION.
          -OUT
     SCD - CREEP DISSIPATION.
          -OUT
     RPL - VOLUMETRIC HEAT GENERATION PER UNIT TIME AT THE END
          OF THE INC.
          -OUT (COUPLED)
     DDSDDT(NTENS) - VARIATION OF STRESS INCREMENTS WITH
          RESPECT TO TEMPERATURE.
С
          -OUT (COUPLED)
С
     DRPLDE(NTENS) - VARIATION OF RPL WRT STRAIN INCREMENTS.
```

```
-OUT (COUPLED)
С
      DRPLDT - VARIATION OF RPL WRT THE TEMPERATURE INCREMENT.
С
           -OUT (COUPLED)
С
      STRAN(NTENS) - TOTAL STRAINS AT BEGINNING OF THE INCREMENT.
С
С
           -IN
      DSTRAN(NTENS) - STRAIN INCREMENTS
С
С
           -IN
      TIME(1) - VALUE OF STEP TIME AT BEGINNING OF CURRENT INCREMENT
С
С
           -IN
      TIME(2) - TOTAL TIME AT BEGINNING OF CURRENT INCREMENT
С
С
           -IN
      DTIME - TIME INCREMENT
С
С
           -IN
      TEMP - TEMPERATURE AT START OF INCREMENT
C
C
           -IN
      DTEMP - INCREMENT OF TEMPERATURE
С
С
           -IN
      PREDEF - ARRAY OF INTERPOLATED VALUES OF PREDEFINED FIELD
С
С
           VARIABLES.
С
           -TN
      DPRED - ARRAY OF INCREMENTS OF PREDEFINED FIELD VARIABLES.
С
С
           -IN
      CMNAME - NAME GIVEN ON *MATERIAL OPTION, LEFT JUSTIFIED.
С
С
           -IN
      NDI - NUMBER OF DIRECT STRESS COMPONENTS AT THIS POINT.
С
С
           -IN
      NSHR - NUMBER OF ENGINEERING SHEAR STRESS COMPONENTS AT
С
С
           THIS POINT.
С
           -IN
      NTENS - SIZE OF THE STRESS OR STRAIN COMPONENT ARRAY
С
С
            (NDI + NSHR)
С
            -IN
      NSTATV - NUMBER OF SOLUTION DEPENDANT STATE VARIABLES.
С
С
            -TN
      PROPS(NPROPS) - ARRAY OF MATERIAL CONSTANTS.
С
С
            -IN
      NPROPS - NUMBER OF MATERIAL CONSTANTS.
С
С
            -IN
      COORDS - ARRAY CONTAINING THE COORDINATES OF THIS POINT.
С
С
            -IN
       DROT(3,3) - ROTAION INCREMENT MATRIX.
С
С
            -TN
       PNEWDT - RATIO OF SUGGESTED NEW TIME INCREMENT TO THE
С
            TIME INCREMENT BEING USED.
С
            -UPDATE
С
       CELENT - CHARACTERISTIC ELEMENT LENGTH.
С
С
            -TN
       DFGRD0(3,3) - ARRAY CONTAINING DEFORMATION GRADIENT
С
            AT THE BEGINNING OF THE INCREMENT
С
С
            -TN
       DFGRD1(3,3) - ARRAY CONTAINING DEFORMATION GRADIENT
С
            AT THE END OF THE INCREMENT
С
С
            -IN
       NOEL - ELEMENT NUMBER
С
С
            -IN
       NPT - INTEGRATION POINT NUMBER
С
С
            -IN
       LAYER - LAYER NUMBER (FOR COMPOSITE SHELLS
С
            AND LAYERED SOLIDS).
 С
С
            -IN
       KSPT - SECTION POINT NUMBER WITHIN THE CURRENT LAYER
 С
 С
            -IN
 С
       KSTEP - STEP NUMBER
            -IN
 C
```

С KINC - INCREMENT NUMBER С -TNС С C23456789012 012 INCLUDE 'ABA PARAM. INC' PARAMETER (ZERO = 0.0D0, HALF = 0.5D0) PARAMETER (ONE = 1.0D0, TWO = 2.0D0, THREE = 3.0D0) PARAMETER (NFROM_UTHERM = 1, NFROM_UMECH = 2) CHARACTER*8 CMNAME CHARACTER*80 FNAME CHARACTER*80 EXTSION CHARACTER*80 DFNAME CHARACTER*80 PATH CHARACTER*80 TNAME1 CHARACTER*80 TNAME2 DIMENSION STRESS (NTENS) DIMENSION STATEV (NSTATV) DIMENSION DDSDDE (NTENS, NTENS) DIMENSION DDSDDT (NTENS) DIMENSION DRPLDE (NTENS) DIMENSION STRAN(NTENS) DIMENSION DSTRAN(NTENS) DIMENSION TIME(2) DIMENSION PREDEF(1) DIMENSION DPRED(1) DIMENSION PROPS(NPROPS) DIMENSION COORDS(3) DIMENSION DROT(3, 3) DIMENSION DFGRD0(3, 3) DIMENSION DFGRD1(3, 3) DIMENSION PLSTRAN(6) DIMENSION OSTRESS(6) DIMENSION OSTRAIN(6) DIMENSION ODSTRAIN(6) INTEGER N(2) INTEGER AB_PRINT INTEGER GAB_PRINT INTEGER AL_STORE INTEGER AB STEP COMMON / KINCINFO / AB_STEP, INC, NCYCLE COMMON / KTIME / STEP_TIME, TOT_TIME, GDT COMMON / KMACFLAGS / NFIRSTTIME, NFIRSTELEM, NFIRSTIP, AL_STORE, N_PEN(100), N_PIN(100), NPEL, & N_PATPEN(100), N_PATPIN(100), NPATPEL, Se . æ AB_PRINT, GAB_PRINT, SIGAL_STATV, NCALL, NFEA_FROM COMMON / KABNAMES / PATH, FNAME, DFNAME, TNAME1, TNAME2, EXTSION COMMON / KPREMAC / NDU, NNOEL, NNPT, NNDI, NNSHR

-

COMMON / KTHERMONLY / DTHERMSTRESS(6), DTHERMSTRAIN(6) DDTHERMSTRAIN DT(6), THERMSTRESS(6), THERMSTRAIN(6) &

```
LOGICAL SIGAL_STATV
```

```
NFEA_FROM = NFROM_UMECH
C*********
C KEEP TRACK OF THE NUMBER OF CALLS TO FEAMAC
C IF NCALL == 1->2 : WE HAVE ALREADY BEEN TO THERMAL
C IF NCALL == 0->1 : NO THERMAL WAS REQUESTED OR DIEMP IS ZERO
      NCALL = NCALL + 1
C FLAG TO DETERMINE HOW DATA WILL BE STORED
  IF SIGAL_STATV == TRUE - DATA STORED IN STATE VARIABLE
С
                                    SPACE PROVIDED BY ABAQUS
C
  IF SIGAL STATV == FALSE - DATA STORED IN FILES
С
       C**
      SIGAL_STATV = .TRUE.
      AB_PRINT = GAB_PRINT
      IF( NPROPS .GE. 1 ) THEN
          IF( PROPS(1) .EQ. 2 ) SIGAL_STATV = .FALSE.
          IF( NPROPS .GE. 2 ) AB_PRINT = INT(PROPS(2))
      ELSE
         SIGAL_STATV = .FALSE.
      ENDIF
      N(1) = NOEL
      N(2) = NOEL
      IF ( AB_PRINT .GE. 1 ) THEN
          æ
          WRITE(NDU, *) '-=> START OF UMAT <=-'
          WRITE(NDU, *) '-=> STORE IN STATEV ? =', SIGAL_STATV
          WRITE(NDU, *) `-=> VALUE OF STEP TIME =', TIME(1), STEP_TIME
WRITE(NDU, *) `-=> TOTAL TIME =', TIME(2), TOT_TIME
                                                 =', DTIME, GDT
=', KSTEP, AB_STEP
=', KINC, INC+1
=', NOEL, NNOEL
          WRITE(NDU, *) '-=> DTIME
          WRITE(NDU, *) '-=> KSTEP
         WRITE(NDU, *) '-=> KINC
WRITE(NDU, *) '-=> NOEL
WRITE(NDU, *) '-=> NPT
                                                 =', NPT, NNPT
=', NCYCLE
          WRITE(NDU, *) '-=> NCYCLE
          WRITE(NDU, *) '-=> NFIRSTELEM
                                                 =', NFIRSTELEM
          WRITE(NDU, *) '-=> NFIRSTIP
                                                 =', NFIRSTIP
          WRITE(NDU, *) '-=> NCALL
                                                  =', NCALL
          WRITE(NDU, *) '-=> NCALL - , NCALL
WRITE(NDU, *) '-=> FNAME =|', FNAME, `|'
WRITE(NDU, *) '-=> TEMP =', TEMP
WRITE(NDU, *) '-=> DTEMP =', DTEMP
WRITE(NDU, *) 'STRAIN IN:'
WRITE(NDU, *) `|', STRAN(1), STRAN(2), STRAN(3)
```

```
WRITE(NDU, *) `|', STRAN(4), STRAN(5), STRAN(6)
WRITE(NDU, *) `DSTRAIN IN:'
      WRITE(NDU, *) 'DSTRAIN IN:
WRITE(NDU, *) '|', DSTRAN(1), DSTRAN(2), DSTRAN(3)
WRITE(NDU, *) '|', DSTRAN(4), DSTRAN(5), DSTRAN(6)
WRITE(NDU, *) 'STRESS IN:'
WRITE(NDU, *) '|', STRESS(1), STRESS(2), STRESS(3)
WRITE(NDU, *) '|', STRESS(4), STRESS(5), STRESS(6)
      WRITE(NDU, *) ' CMNAME = | ', CMNAME, ' | '
WRITE(NDU, *) ' NDI = ', NDI
WRITE(NDU, *) ' NSHR = ', NSHR
WRITE(NDU, *) ' NTENS = ', NTENS
WRITE(NDU, *) ' NSTATV = ', NSTATV
WRITE(NDU, *) ' NPROPS = ', NPROPS
WRITE(NDU, *) ' CELENT = ', CELENT
WRITE(NDU, *) ' NOEL = ', NOEL
WRITE(NDU, *) ' NPT = ', NPT
WRITE(NDU, *) ' LAYER = ', LAYER
WRITE(NDU, *) ' KSTEP = ', KSTEP
WRITE(NDU, *) ' KINC = ', KINC
       WRITE(NDU, *) `-=> N=', N, `|'
       WRITE(NDU, *) '-=> INC
                                                   =', INC
       WRITE(NDU, *) '-=> NCYCLE =', NCYCLE
       WRITE(NDU, *) '-=> TEMP =', TEMP
       WRITE(NDU, *) '-=> DTEMP =', DTEMP
      WRITE(NDU, *) '-=> BEFORE FEAMAC PRE <=-'
 ENDIF
 DO J = 1, 6
      OSTRESS(J) = STRESS(J)
       STRAN(J) = STRAN(J) + THERMSTRAIN(J)
      DSTRAN(J) = DSTRAN(J) + DTHERMSTRAIN(J)
      OSTRAIN(J) = STRAN(J)
      ODSTRAIN(J) = DSTRAN(J)
 ENDDO
 CALL FEAMAC_PRE(DDSDDE, PLSTRAN, STRAN, DSTRAN, STRESS, STATEV,
&
          NSTATV, NTENS, N, NPT, LAYER, NDI, NSHR,
£
          TEMP, DTEMP, THERMSTRAIN)
 IF( (NCALL .EQ. 2) .AND. (NCYCLE .NE. 0) ) THEN
       DO I = 1, 6
            STRESS(I) = OSTRESS(I)
            DO J = 1, 6
                 STRESS(I) = STRESS(I)
&
                         + DDSDDE(I, J) * (ODSTRAIN(J) - DTHERMSTRAIN(J))
            ENDDO
       ENDDO
      DO J = 1, 6
```

-

```
IF ( DTEMP .NE. 0.0 ) THEN
             DDSDDT(J) = DTHERMSTRESS(J) / DTEMP
          ELSE
             DDSDDT(J) = 0.0
          ENDIF
          DRPLDE(J) = 0.0
        ENDDO
        RPL = 0.0
       DRPLDT = 0.0
     ENDIF
     IF ( AB_PRINT .GE. 1 ) THEN
WRITE(NDU, *) `-=> AFTER FEAMAC_PRE <=-'</pre>
       WRITE(NDU, *) 'DTHERMAL STRESS:'
        WRITE(NDU, *) '|', DTHERMSTRESS(1), DTHERMSTRESS(2),
            DTHERMSTRESS(3)
    æ
        WRITE(NDU, *) `|', DTHERMSTRESS(4), DTHERMSTRESS(5),
            DTHERMSTRESS(6)
    &
        WRITE(NDU, *) 'DTHERMAL STRAIN:'
        WRITE(NDU, *) '|', DTHERMSTRAIN(1), DTHERMSTRAIN(2),
            DTHERMSTRAIN(3)
    &
        WRITE(NDU, *) `|', DTHERMSTRAIN(4), DTHERMSTRAIN(5),
            DTHERMSTRAIN(6)
    æ
       WRITE(NDU, *) `DDSDDT:'
WRITE(NDU, *) `|', DDSDDT(1), DDSDDT(2),
            DDSDDT(3)
    æ
        WRITE(NDU, *) `|', DDSDDT(4), DDSDDT(5),
            DDSDDT(6)
    &
        WRITE(NDU, *) '-=> END OF UMAT <=-'
     ENDIF
C WHATEVER IS IN STRAN/DSTRAN WILL SHOW UP AS MECHANICAL STRAIN
C MECHANICAL STRAIN + THERMAL STRAIN WILL GIVE GLOBAL STRAIN
C WHATEVER IS IN STRESS WILL BE GIVEN AS GLOBAL STRESS
     DO I = 1, 6
        STRAN(I) = OSTRAIN(I) - THERMSTRAIN(I)
        DSTRAN(I) = ODSTRAIN(I) - DTHERMSTRAIN(I)
     ENDDO
     NFIRSTTIME = 0
     RETURN
     END
С
  7778
01234
C
   1111.
                1/ //
С
С
  1111
                 11 1_1 1
              С
  //_//__
             1
                 17
С
           .//_//_/_/
С
С
  7778
```

c 2 3 4 5 6 7 8 9 0 1 2

SUBROUTINE UEXPAN (EXPAN, DEXPANDT, TEMP, TIME, DTIME, PREDEF, DPRED, STATEV, CMNAME, NSTATV) æ INCLUDE 'ABA PARAM. INC' PARAMETER (ZERO = 0.0D0, HALF = 0.5D0) PARAMETER ($\dot{ONE} = 1.0D0$, TWO = 2.0D0, THREE = 3.0D0) PARAMETER (NFROM_UTHERM = 1, NFROM_UMECH = 2) CHARACTER*8 CMNAME DIMENSION EXPAN(*), DEXPANDT(*), TEMP(2), TIME(2), PREDEF(*) DIMENSION DPRED(*), STATEV(NSTATV) CHARACTER*80 FNAME CHARACTER*80 EXTSION CHARACTER*80 DFNAME CHARACTER*80 PATH CHARACTER*80 TNAME1 CHARACTER*80 TNAME2 DIMENSION STRAN(6), DSTRAN(6) DIMENSION STRESS(6) DIMENSION PLSTRAN(6) DIMENSION DDEDDT(6, 6) INTEGER N(2) INTEGER AB_PRINT INTEGER GAB PRINT INTEGER AL_STORE INTEGER AB_STEP COMMON / KINCINFO / AB_STEP, INC, NCYCLE COMMON / KTIME / STEP_TIME, TOT_TIME, GDT COMMON / KMACFLAGS / NFIRSTTIME, NFIRSTELEM, NFIRSTIP, & AL_STORE, N_PEN(100), N_PIN(100), NPEL, N_PATPEN(100), N_PATPIN(100), NPATPEL, & AB_PRINT, GAB_PRINT, SIGAL_STATV, NCALL, NFEA_FROM Sc. COMMON / KABNAMES / PATH, FNAME, DFNAME, TNAME1, TNAME2, EXTSION COMMON / KPREMAC / NDU, NNOEL, NNPT, NNDI, NNSHR COMMON / KTHERMONLY / DTHERMSTRESS(6), DTHERMSTRAIN(6), DDTHERMSTRAIN_DT(6), THERMSTRESS(6), THERMSTRAIN(6) & LOGICAL SIGAL_STATV NFEA FROM = NFROM_UTHERM AB_PRINT = GAB_PRINT DO J = 1, 6PLSTRAN(J) = 0.0STRAN(J) = 0.0DSTRAN(J) = 0.0

202

STRESS(J) = 0.0PLSTRAN(J) = 0.0 ς.

```
DTHERMSTRESS(J) = 0.0
          DTHERMSTRAIN(J) = 0.0
          DDTHERMSTRAIN_DT(J) = 0.0
          DO I = 1, 6
             DDEDDT(I, J) = 0.0
          ENDDO
          EXPAN(J) = 0.0
          DEXPANDT(J) = 0.0
      ENDDO
C NOTE THE "TIME" ARRAY FOR THIS ROUTINE IS PROVIDED AT THE -E N D-
  OF THE INCREMENT
       TIME(1) = TIME(1) - DTIME
      TIME(2) = TIME(2) - DTIME
C NOTE THE "TEMP" ARRAY FOR THIS ROUTINE IS PROVIDED AT THE -E N D-
  OF THE INCREMENT
      DTEMP = TEMP(2)
      TEMP(1) = TEMP(1) - DTEMP
      TEMPERATURE = TEMP(1)
       IF ( DTEMP .EQ. 0.0 ) THEN
           WRITE(NDU, *) 'UEXPAN EARLY EXIT - DTEMP = ', DTEMP,
С
            ' TEMP = ', TEMPERATURE
С
          RETURN
       ENDIF
      NCALL = NCALL + 1
      SIGAL_STATV = .FALSE.
      IF ( AB_PRINT .GE. 1 ) THEN
          WRITE(NDU, *) ``
          `_____
     æ
          WRITE(NDU, *) '-=> START OF UEXPAN <=-'
         WRITE (NDU, *)'-=> START OF GEAFAN <--</td>WRITE (NDU, *)'-=> VALUE OF STEP TIME =', TIME(1), STEP_TIMEWRITE (NDU, *)'-=> TOTAL TIMEWRITE (NDU, *)'-=> DTIMEWRITE (NDU, *)'-=> KSTEPWRITE (NDU, *)'-=> KSTEP=', AB_STEPWRITE (NDU, *)'-=> KINC=', INC+1
                                                    =', INC+1
          WRITE(NDU, *) '-=> NOEL
                                                     =', NNOEL
          WRITE(NDU, *) '-=> NPT
                                                     =', NNPT
          WRITE(NDU, *) '-=> NNDI
                                                    =', NNDI
          WRITE(NDU, *) '-=> NMSHR
WRITE(NDU, *) '-=> NCYCLE
WRITE(NDU, *) '-=> NFIRSTELEM
WRITE(NDU, *) '-=> NFIRSTIP
                                                    =', NNSHR
                                                    =', NCYCLE
                                                     =', NFIRSTELEM
=', NFIRSTIP
=', NCALL
          WRITE(NDU, *) '-=> NCALL
          WRITE(NDU, *) '-=> FNAME = |', FNAME, '|'
          WRITE (NDU, *) '-=> TEMPERATURE =', TEMPERATURE
          WRITE (NDU, *) '-=> DTEMP =', DTEMP
          WRITE(NDU, *) '-=> BEFORE FEAMAC_PRE <=-'
      ENDIF
      N(1) = NNOEL
      N(2) = NNOEL
      LAYER = 0
      NTENS = 6
```

.

C NOTE: HERE DSTRAN IS NOT A RATE BUT THE ALPHAS INSTEAD - I.E. THE VARIATION OF THERMAL STRAINS WITH RESPECT TO TEMPERATURE (NOT TIME) С CALL FEAMAC_PRE(DDEDDT, PLSTRAN, STRAN, DSTRAN, STRESS, STATEV, NSTATV, NTENS, N, NNPT, LAYER, NNDI, NNSHR, & TEMPERATURE, DTEMP, THERMSTRAIN) & C NOTE: THESE ARE INCREMENTS IN THERMAL STRAIN/STRESS C NEED TO FIGURE OUT A GOOD WAY TO GET THE SIZES OF THESE DO J = 1, 6EXPAN(J) = STRAN(J)DTHERMSTRAIN(J) = STRAN(J) $DDTHERMSTRAIN_DT(J) = DSTRAN(J)$ DEXPANDT(J) = DSTRAN(J)ENDDO DO I = 1, 6 DTHERMSTRESS(I) = 0.0DO J = 1, 6 DTHERMSTRESS(I) = DTHERMSTRESS(I) + DDEDDT(I, J) * DTHERMSTRAIN(J) & ENDDO STRESS(I) = DTHERMSTRESS(I) ENDDO RETURN END

.

С

6.16 Example P: Non-Symmetric Laminate

Sample Input File For Non-Symmetric Laminate Problem

The following example is used to explain the use of monolithic layer in more detail.

Problem Summary:

Load Type:	Thermomechanical
Load Component:	11-direction (applied mid-plane strain)
Load History:	Monotonic
Load Control:	Strain
Load History Data:	Cool-down from 371.1°C to 21.1 °C then hold temperature constant during mechanical load-ing
	$\dot{T} = 35.$ °C/sec
	$\dot{\epsilon} = 1.x 10^{-3} / \text{sec}, \ \epsilon_{max} = 0.01$,
	$\varepsilon_{min} = 0.$

 $\Delta t_{thermal} = 0.05 \text{ sec}, \Delta t_{mech} = 0.01 \text{ sec}$

Micromechanics model:

Laminate Option

Laminate Details:

Layer	Thickness (mm)	Fiber	Matrix	Fiber Volume Fraction	Unit Cell	Packing Arrangement	Angle
1	1	mono- lithic	Aluminum (2024-T4)	-	-	-	-
2	1	Graphite T-50	Aluminum (2024-T4)	30%	2x2 sub- cells	square	90
3	1	Graphite T-50	Aluminum (2024-T4)	30%	2x2 sub- cells	square	0

Integration Algorithm:

Forward Euler

Constituent Material Model:

Bodner-Partom

Note: The laminate analyzed in this example is shown in following figure. The macro output for the laminate option consists of stress resultants and mid-plane strains. The micro output is always in the local coordinates of the layer in question. Micro output is given for both integration points within the layer in question.



example of a non-symmetric laminate with monolithic and heterogeneous layers *PRINT NPL=0 % *LOAD LCON=3 LOP=1 LSS=1 *MECH NPTW=3 TI=0.0, 10.0, 20.0 LO=0.0, 0.0, 0.01 *THERM NPTT=3 TI=0.0, 10.0, 20.0 & TE=371.1, 21.1, 21.1 % *MODEL MOD=3 MATSYS=2 NLY=3 THK=0.001,0.001,0.001 CON=1,2,2 & SYS=1,2,2 ANG=0.,90.,0. % *SOLVER NTF=1 NPTS=3 TIM=0.0, 10.0, 20.0 STP=0.05, 0.01 % *FIBER NFIBS=1 NF=1 MS=2 MF=9 NDPT=2 MAT=A D=1.,0.,0.% *MATRIX NMATX=1 NM=1 MS=2 MM=1 NDPT=2 MAT=A % *MONOL NMON=1 NMO=1 MS=1 MMO=1 NDPT=2 MAT=A % *MRVE IDP=0.1.1 L=1 L=2 VF=0.3 L=3 VF=0.3 % ***CURVE** NP=20 % *MACRO NT=1 NC=1 X=1 Y=7 NAM=lam-x % *MICRO NT=3 NC=1 LYR=1 X=37 Y=1 NAM=lam-x1 NC=2 LYR=2 CELL=2 X=37 Y=2 NAM=lam-x2 NC=3 LYR=3 CELL=2 X=37 Y=1 NAM=lam-x3 % *END

The following figures were obtained from the macro and micro x-y plot data files produced by the present example.



Local Level Results

Note: The increasing magnitude of the strain towards the lower surface of the laminate is due to the increased bending resulting from the nonsymmetric lay-up.

6.17 Example Q: Yield Surface Plot

Sample Input File for a Yield Surface Plot

The following example is used to explain the yield surface capabilities in more detail.

Problem Summary:	
Load Type:	Themomechanical
Load Component:	Combined 11-component and 22-component
Load History:	Yield surface probing
Load Control:	Strain
Load History Data:	Constant temp. = 23 ° C $\dot{\epsilon}$ =1.0x10 ⁻⁵ /sec , ϵ_{max} =0.05, ϵ_{min} =0. (along each probing angle) Δt = 5. sec.
Micromechanics model:	Double Periodicity
Fiber Packing Arrangement:	Square pack at 35% fiber volume ratio
Integration Algorithm:	Forward Euler
Constituent Material Model:	Fiber: Elastic Matrix: GVIPS – isotropic form
Constituents:	Fiber: SCS-6 (properties input manually) Matrix: TIMETAL21S

```
test of yield surface option
*PRINT
 NPL=0 %
*LOAD
 LCON=3 LOP=1 LSS=1 %
*MECH
 NPTW=2 TI=0.,5000. LO=0.,0.05 %
*THERM
 NPTT=2 TI=0.,5000. TE=23.,23. %
*SURF
 NPRE=0 ISP=1 IAN=10
 C1=10.E-6 C2=1.45E-7 C3=0.0 C4=0.0
*MODEL
 MOD=1 %
*SOLVER
 NTF=1 NPTS=2 TIM=0.,5000. STP=5. %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=1 MAT=U IFM=1&
 EL=58.E3,58.E3,0.20,0.20,24.167E3,4.5E-6,4.5E-6 %
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=2 MAT=A %
*MRVE
 IDP=1 VF=0.35 %
*CURVE
 NP=1 %
*MACRO
 NT=1
 NC=1 X=2 Y=8 NAM=surf %
*END
```

Note: A zero value for a given criteria means that specific criteria will be immediately satisfied, consequently the output associated with that criteria is meaningless. The output associated with the nonzero criteria are correct.



Resulting yield surface

6.18 Example R: Ellipsoidal Inclusions

Sample Input File for Ellipsoidal Inclusions

Prob	lem	Sum	marv

Load Type:	Thermomechanical
Load Direction:	33
Load History:	Monotonic
Load Control:	Strain
Load History Data:	Hold temperature during mechanical loading $\dot{\epsilon}$ =6.667x10 ⁻⁵ /sec, ϵ_{max} =0.01, ϵ_{min} =0. $T = 427 \text{ °C}; \Delta t_{initial} =0.01 \text{ sec.}$
Micromechanics Model:	Triple Periodicity
Repeating Unit Cell:	Ellipsoidal Inclusion RVE (IDP=4) a = 1, b = 1, c = 1.3, d/h = 1, l/h = 1
Integration Algorithm:	Predictor/Corrector
Constituent Material Model:	Fiber: Elastic Matrix:TGVIPS
Constituents:	Fiber: SCS-6 Matrix:Ti-6-4

•

```
*PRINT
NPL=1 %
*LOAD
LCON=3 LOP=3 LSS=1 %
*MECH
NPTW=2 TI=0.0, 150.0 LO=0.0, 0.01 %
*THERM
NPTT=2 TI=0.0, 150.0 TE=427.0, 427.0 %
*MODEL
 MOD=2 %
*SOLVER
 NTF=2 ISTM=0.01 ISTT=5. ERR=0.01 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=2 MAT=E %
*MATRIX
 NMATX=1
 NM=1 MM=7 NDPT=2 MAT=A D=1.0, 1.0, 1.0 %
*MRVE
IDP=4
 OPT=2 VF=0.3 A=1 B=1 C=1.3 RD=1 RL=1 %
# OPT=2 VF=0.3 A=1 B=1 C=1.2 RD=1 RL=1 %
# OPT=2 VF=0.3 A=1 B=1 C=1. RD=1 RL=1 %
# OPT=2 VF=0.3 A=1 B=1 C=0.83333 RD=1 RL=1 %
*CURVE
 NP=1 %
*MACRO
NT=1
 NC=1 X=3 Y=9 NAM=ellipsoid %
*END
```

Note: To generate the four curves shown in the plot, change which line does not start with "#" under *MRVE



Results of Discontinuously reinforced composite with 30% volume fraction of ellipsoidal inclusions

Note: As the ellipsoidal inclusion becomes longer and thinner (i.e., increasing c) and thus more fiber-like, the response becomes stiffer.
6.19 Example S: PATRAN/MSC MACPOST Output

Sample Input File for PATRAN/MSC MACPOST Output

Problem Summary

Load Type:	Thermomechanical
Load Direction:	33
Load History:	Monotonic
Load Control:	Strain
Load History Data:	Cool from 900°C to 23°C, then hold temperature during mechanical loading \dot{T} =0.0152 °C/sec $\dot{\varepsilon}$ =1.667x10 ⁻⁴ /sec, ε_{max} =0.015, ε_{min} =0. $\Delta t_{thermal}$ = 100 sec.; Δt_{mech} =0.1 sec.
Micromechanics Model:	Double Periodicity
Fiber Packing Arrangement:	Square Pack, 40% fiber volume ratio
Repeating Unit Cell:	26x26 circular fiber cross-section approximation
Integration Algorithm:	Forward Euler
Constituent Material Model:	Fiber: Elastic Matrix: GVIPS – isotropic form
Constituents:	Fiber: SCS-6 Matrix: TIMETAL21S
PATRAN Output:	Start output after cooldown: <i>tpre</i> = 57600 sec. Write to output files every 300 time steps

```
test of PATRAN output
*PRINT
 NPL=1 %
*LOAD
LCON=3 LOP=3 LSS=1 %
*MECH
 NPTW=3 TI=0.,57600.,57690. LO=0.,0.,0.015 %
*THERM
 NPTT=3 TI=0.,57600.,57690. TE=900.,23.,23. %
*MODEL
 MOD=1 %
*PATRAN
 FN=apdxs TPRE=57600 STP=300 %
*SOLVER
 NTF=1 NPTS=3 TIM=0.,57600.,57690. STP=100,0.1 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=2 MAT=E %
*MATRIX
 NMATX=1
 NM=1 MM=4 NDPT=2 MAT=A %
*MRVE
 IDP=13 VF=0.4 R=1
*CURVE
 NP=2 %
*MACRO
 NT=1
 NC=1 X=3 Y=9 NAM=apdxs
*END
```



Macro stress-strain behavior results



Contour of the microlevel J_2 invariant at 1.5% macrostrain (@ pt. A)



Contour of the microlevel I_1 invariant at 1.5% macrostrain (@ pt. A)

6.20 Example T: Effective Thermal Conductivity Calculation

Sample Input File For Effective Thermal Conductivity Calculation

The following example is used to explain the calculation of effective thermal conductivities.

Problem Summary:

Loading:	Not Used (properties c	nplvl = -1 used to determine effective only)
Micromechanics Model:	Double Per	iodicity
Fiber Packing Arrangement:	Square Pac	ck, R = 1., 50% fiber volume ratio
Repeating Unit Cell:	7x7 circulaı	fiber cross-section approximation
Integration Algorithm:	Not Used	
Constituent Material Model:	Fiber:	Elastic, isotropic
	Matrix:	Bodner-Partom (viscoplastic properties not used)
Constituents:	Fiber:	Fictitious temperature dependent mate- rial (based loosely on tungsten)
	Matrix:	Fictitious temperature dependent mate- rial (based loosely on aluminum)

```
Test of user input properties with thermal conductivity
*PRINT
 NPL=-1 %
*LOAD
 LCON=3 LOP=2 LSS=1 %
*MECH
 NPTW=3 TI=0.,200.,218. LO=0.,0.,0.015 %
*THERM
 NPTT=3 TI=0.,200.,218. TE=600.,21.,21. %
*MODEL
 MOD=1 %
*COND
*SOLVER
NTF=1 NPTS=3 TIM=0.,200.,218 STP=0.2,0.01 %
*FIBER
 NFIBS=1
 NF=1 MF=6 NDPT=2 MAT=U IFM=1
 NTP=4
 TEM=18.,200.,400.,600.
 EA=314.1E9,300.E9,280.E9,200.E9
 ET=314.1E9,300.E9,280.E9,200.E9
 NUA=0.41,0.41,0.41,0.41
 NUT=0.41,0.41,0.41,0.41
 GA=111.38E9,106.38E9,99.29E9,70.92E9
 ALPA=4.5E-6.4.8E-6.5.1E-6.5.5E-6
 ALPT=4.5E-6,4.8E-6,5.1E-6,5.5E-6
 KA=0.2,0.25,0.31,0.44
 KT=0.2,0.25,0.31,0.44
*MATRIX
 NMATX=1
 NM=1 MM=1 NDPT=2 MAT=U IFM=1
 NTP=4
 TEM=18.,200.,400.,600.
 EA=72.E9,67.E9,55.E9,33.E9
 ET=72.E9,67.E9,55.E9,33.E9
 NUA=0.33,0.33,0.33,0.33
 NUT=0.33,0.33,0.33,0.33
 GA=27.07E9,25.19E9,20.68E9,12.41E9
 ALPA=22.E-6,24.E-6,28.E-6,33.E-6
 ALPT=22.E-6,24.E-6,28.E-6,33.E-6
 V1=1.E4, 1.E4, 1.E4, 1.E4
 V2=340.E6, 340.E6, 340.E6, 340.E6
 V3=435.E6, 435.E6, 435.E6, 435.E6
 V4=300.0, 300.0, 300.0, 300.0
 V5=10.0, 4.0, 1.6, 0.55
 V6=1.0, 1.0, 1.0, 1.0
```

KA=40.,38.,35.,29. KT=40.,38.,35.,29. *MRVE IDP=6 VF=0.5 R=1. % *CURVE NP=10 % *MACRO NT=1 NC=1 X=2 Y=8 NAM=apdxt % *END

Thermal conductivity results from outfile:

Effective Thermal Conductivities At Temperature = 18.0K Axial = 0.201E+02K Transverse = 0.889E+01At Temperature = 200.0K Axial = 0.191E+02K Transverse = 0.852E+01At Temperature = 400.0K Axial = 0.177E+02K Transverse = 0.795E+01At Temperature = 600.0K Axial = 0.147E+02K Transverse = 0.683E+01



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101 CELL INDEX 87 CM 63 CON **Keywords** 67 conductivity 67, 71 *COND 68 constituents 99 ***CURVE** 16 constitutive models 60, 69 *DAMAGE 77 CPER 95 *DEBOND 126 **Cross-Ply Laminate Problem** 68, 73, 74 ***FIBER *INTERFACE** 94 D 54.55 ***LOAD** 60,68 D 100 ***MACRO** Database 72 68,74 ***MATRIX** 95 DBCH 57 *MECH 42, 43 debonding 101 ***MICRO** Discrete Fiber Breakage Problem 168 63 *MODEL 60 DMAX 68 ***MONOL** 63 double periodicity 77 ***MRVE** 64 *PATRAN Ε ***PRINT** 53 Effective Thermal Conductivities 219 66 ***SOLVER** 86 Ellipsoidal 62 ***SURF** 212 **Ellipsoidal Inclusion Problem** 58 ***THERM** 36 engineering strains 66 ERR Symbols 69 ETU 52 % symbol 105 Examples 52 "#" symbol execution of MAC/GMC 46 - large finite element problems 49 Α 181 ABAQUS 14 Adams Bashforth F 15 Adams-Moulton 95 FACE ANG 63, 69 60 Fatigue 17 Anisotropic Elastic fatigue damage 30 156 Fatique Damage Analysis B 181 FEAMAC 95 **BCN** 181 **FEAMAC Analysis Example** 95 BCS 49 FEAMAC Input Deck 95 **BDN** 50 **FEAMAC Output Files** 95 BDS 60 FG **Biaxial Load Problem** 135 fiber and matrix constituents 68 69 BN finite element implementation 47 **Bodner-Partom** 18 60 FL BP 69 64 FN Forward Euler 14 С C1 62 G 62 C2 gauss quadrature points 28 C3 62 95 GCN 62 C4

GCS	95	Ν	
General Loading Problem	120	NA	87
generalized loading	54	NAI	95
generalized method of cells	s 3, 7	NAM	100, 101
GVIPS	22	NB	87
		NBI	95
н		NC	100, 101
н	87	NCY	60
Header Line	53	NDPT	68.94
	00	NF	68
1		NEIBS	68
	62	NG	87
	60	NGI	95
	77	NI	94
input and output files	// /7	NII	95
Input and output mes		NINT	94
Interface Modeling	42	NIY	63
internace would ing	76	NM	68
	62	NMATX	68
	66	NMO	68
	66	NMON	68
1511	00	nonisothermal	73
		Non-Symmetric Laminate	205
	70 07	NP	99
	70,07 26 05	NPI	53
Laminate	20, 95,	NPRE	62
IUI, IU2	62 101	NPTS	66
liaminate option	100, 101	NPTT	58
Laminate Problem	123	NPTW	57
	54 57	NT	101
	57	NTE	66
	54 54	numbering scheme	103
	04 101	Hambering Scheme	
LYR	101	0	
		OMEL	60
	045		60
	215		69
MAI	68, 69, 94	OMO	03
Material Constant Databas	se /2	Р	
matrix	68		77
MAISYS	63	Plain Weeve Composites	174
Mechanical Load Problem	105	Plain weave Composites	174
MF	68	predictor/corrector	14
MI	94	8	
micromechanics	/	R	
MM	68	R	//
MMO	68	H = X/Y	/8 77
MOD	63, 85	HAD	//
monolithic material	68	HAD1	//
MS	68	HAD2	- 17

÷

.

reformulation of GMC	11
residual stresses	111
RN	95
Robinson Viscoplastic	20
RS	95
Runge-Kutta	14

S	
SFL	69
SK	69
state variables	183
STO	64
STP	64, 66
SU	69
Subcell	103
SYS	63

_	_		
I			

Т	60
TE	58
temperature	58
TGVIPS	24
thermal conductivities	74
thermal conductivity	67
Thermal Load Problem	108
Thermomechanical Load	111
ТНК	63
ТІ	57, 58, 95
ТІМ	66
TOLN	95
Transverse Debonding	114
Triple Periodic GMC	129
triple periodicity	63

U

user defined architecture 132 User Defined Example 138 User Defined RVE Problem 132 User Supplied Isothermal 73 User Supplied NonIsothermal 73, 74 USRELS 37 USRMAT 34, 42

V	
V	60
VF	77
VF1	77
VF2	77

Х	
Х	100, 101
XA	77
XML	69
V	
Y	100 101
Y	100, 101
Yield Surface Plot	209

ĩ .

,

7

reformulation of GMC residual stresses RN Robinson Viscoplastic RS Runge-Kutta	11 111 95 20 95 14
S SFL SK state variables STO STP SU Subcell SYS	69 69 183 64 64, 66 69 103 63
T T TE temperature TGVIPS thermal conductivities thermal conductivity Thermal Load Problem Thermomechanical Load THK TI TIM TOLN Transverse Debonding Triple Periodic GMC	60 58 58 24 74 67 108 111 63 57, 58, 95 66 95 114 129

U

triple periodicity

user defined architecture 132 User Defined Example 138 User Defined RVE Problem 132 User Supplied Isothermal 73 User Supplied NonIsothermal 73, 74 USRELS 37 **USRMAT** 34, 42

63

60

77

77

77

۷ V VF VF1 VF2

^	
Х	100, 101
ХА	77
XML	69
Y	
Υ	100, 101
Yield Surface Plot	209

v

REPORT DOCUMENTATION PAGE

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